Simone M. Cacciò Giovanni Widmer *Editors*

Cryptosporidium: parasite and disease



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Preface

It is more than a 100 years since Cryptosporidium parasites were first described by Edward Tyzzer.¹ In addition to the initial discovery, two events stand out for having significantly impacted our awareness of these parasites and the diseases they cause. The HIV epidemic and the emergence of cryptosporidiosis as a potentially severe opportunistic infection in people living with AIDS was significant, not only because of the clinical implications but because it motivated a substantial research effort and a desire to better understand these parasites. A second notable event was the waterborne cryptosporidiosis outbreak which occurred in Milwaukee in 1993. The importance of this outbreak is illustrated by the fact that the article² reporting on the epidemiological investigation of the outbreak was cited over 600 times, more than any other paper with the term *Cryptosporidium* in the title listed in PubMed. Subsequently, the application of genotyping methods based on the polymerase chain reaction revealed the common occurrence of cryptosporidiosis in animals, particularly in young livestock, and the importance of zoonotic transmission of this parasite. Whereas waterborne outbreaks have provided incentives for improving drinking water quality, progress in treatment and prevention of cryptosporidiosis has been disappointing. As described in this book, basic research has generated a wealth of information on many aspects of Cryptosporidium biology, but this knowledge has not had a decisive impact on the progress towards the production of effective treatments or vaccines. Cryptosporidiosis thus remains a serious infection, not only for immunocompromised individuals but also for children living in underdeveloped countries. In recognition of the disease burden, and to underline its link with poverty, the World Health Organization (WHO) has included cryptosporidiosis in the Neglected Diseases Initiative since 2004.

¹Tyzzer, E.E. (1907) A sporozoon found in the peptic glands of the common mouse. Proc. Soc. Exp. Biol. Med. 5,12–13.

² Mac Kenzie WR, Hoxie NJ, Proctor ME, Gradus MS, Blair KA, Peterson DE,Kazmierczak JJ, Addiss DG, Fox KR, Rose JB, et al. (1994) A massive outbreak in Milwaukee of *Cryptosporidium* infection transmitted through the public water supply. N Engl J Med. 331, 161–167.

The content of this book reflects the extent to which our knowledge of *Cryptosporidium* parasites has expanded in recent years. The 13 chapters are written by scientists, clinicians and veterinarians having many years of experience with these parasites and who together have published hundreds of research papers. This practical experience and scholarly activity is reflected in the quality of the contributions. The book covers a wide range of subjects, ranging from clinical cryptosporidiosis to the epidemiology, taxonomy, host-parasite interaction and molecular biology. Recent progress in the field of *Cryptosporidium* "omics" is also covered.

We have grouped the chapters in four parts. Part I covers taxonomy and epidemiology. It includes four chapters on the molecular taxonomy, epidemiology, evolution and ecology of *Cryptosporidium* species infecting humans, livestock and other vertebrates. Part II covers "omics"-related subjects (genomics, transcriptomics and proteomics), as well as an overview of *Cryptosporidium* metabolism and its many unique features. Host-parasite interaction is the focus of Part III. Clinical cryptosporidiosis, immunology and the current state of drug development are covered. The last part is devoted to waterborne cryptosporidiosis. Two chapters review the implications of *Cryptosporidium* oocysts in drinking and recreational water and give an overview of treatments to remove and inactivate oocysts in drinking water.

This book is evidence of the devotion of 34 authors residing in seven countries who together have invested a significant effort in reviewing and interpreting recent progress in their respective fields. The editors would like to express their gratitude to the authors for taking on this task.

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Part I Taxonomy and Epidemiology of Cryptosporidium

Chapter 1 Taxonomy and Molecular Taxonomy

Una Ryan and Lihua Xiao

Abstract Cryptosporidium parasites belong to the phylum Apicomplexa and possess features of both the coccidia and gregarines. Currently, 25 species of Cryptosporidium are recognized in fish, amphibians, reptiles, birds and mammals. All 25 species have been confirmed by morphological, biological, and molecular data. Cryptosporidium duismarci and C. scophthalmi lack sufficient biological and/or molecular data to be considered valid species. In addition to the named species, more than 40 genotypes from various vertebrate hosts have been described. For these genotypes to receive taxonomic status, sufficient morphological, biological, and molecular data are required and names must comply with the rules of the International Code for Zoological Nomenclature (ICZN). A different interpretation of the ICZN led to the proposal that Cryptosporidium parvum be renamed as Cryptosporidium pestis and that C. parvum be retained for C. tyzzeri. However, this proposal violates the guiding ICZN principle of maintaining taxonomic stability and avoiding confusion. In addition, C. pestis lacks a full taxonomic description and therefore is not a valid species. The taxonomic status of *Cryptosporidium* spp. is rapidly evolving and many genotypes are likely to be formally described as species in the future.

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1.1 Introduction

The genus *Cryptosporidium* is composed of protozoan parasites that infect epithelial cells in the microvillus border of the gastrointestinal tract of all classes of vertebrates. The parasite causes self-limiting diarrhoea in immunocompetent individuals but the infection may be chronic and life-threatening to those that are immunocompromised (Hunter et al. 2007). The incubation period for illness symptoms is approximately 7 days (range 1–14 days) and illness is usually selflimiting, with a mean duration of 6–9 days (Hunter et al. 2007), although longer times (mean duration 19–22 days, maximum up to 100–120 days) were reported in Australian cases (Robertson et al. 2002). Relapses are common; reports indicate 1–5 additional episodes in 40–70 % of patients (Hunter et al. 2004). The predominant symptom is watery diarrhoea, sometimes profuse (1–2 L/day in a small minority of cases, usually very young or old), sometimes mucous but rarely bloody.

The oocyst, the environmental stage of *Cryptosporidium*, is ubiquitous and therefore cryptosporidiosis can be acquired through a variety of transmission routes including direct contact with infected persons (person-to-person transmission), contact with animals (zoonotic transmission) and ingestion of contaminated food (foodborne transmission) or water (waterborne transmission) (Xiao 2010) (Chap. 2).

The advent of molecular tools for characterization and phylogenetic analysis of *Cryptosporidium* has revolutionized our understanding of *Cryptosporidium* taxonomy, which in turn underlies our ability to understand the biology, epidemiology and health related importance of various *Cryptosporidium* species (Xiao et al. 1999a, 2004a; Fayer 2010).

1.2 Cryptosporidium's Taxonomic Relationship to Gregarine Parasites

The genus *Cryptosporidium* belongs to the phylum Apicomplexa, which comprises parasitic eukaryotes possessing an apical complex at some stage in their life cycle. Within the Apicomplexa, *Cryptosporidium* species have been traditionally considered to be intestinal coccidian parasites, based on the possession of life cycle features and morphological characteristics similar to other enteric coccidian parasites (Levine 1988). Despite similarities, *Cryptosporidium* demonstrates several peculiarities that separate it from any other coccidian. These include (1) the location of *Cryptosporidium* within the host cell, where the endogenous developmental stages are confined to the apical surfaces of the host cell, where a multimembranous attachment or feeder organelle is formed at the base of the parasitiphorous vacuole to facilitate the uptake of nutrients from the host cell; (3) the presence of two morpho-functional types of oocysts, thick-walled and

thin-walled, with the latter responsible for the initiation of the auto-infective cycle in the infected host; (4) the small size of the oocyst ($4.9 \times 4.4 \,\mu\text{m}$ for *C. parvum*) which lacks morphological structures such as sporocyst, micropyle and polar granules (Tzipori and Widmer 2000; Petry 2004); (5) the insensitivity to all anticoccidial agents tested so far (Blagburn and Soave 1997; Cabada and White 2010); and (6) the presence of a novel gamont-like extracellular stage similar to those found in gregarine life cycles (Hijjawi et al. 2002; Rosales et al. 2005).

Gregarines are a diverse group of apicomplexan parasites that consist of large single-celled parasites that inhabit the intestines and other extracellular spaces of invertebrates and lower vertebrates, which are abundant in natural water sources (Barta and Thompson 2006; Levine 1988). Molecular phylogenies suggest that Cryptosporidium is evolutionarily divergent from other coccidia but related to gregarines and is an early branch at the base of the apicomplexan phylum (Zhu et al. 2000; Barta and Thompson 2006). Genomic and biochemical data also indicates that Cryptosporidium differs from other apicomplexans in that it has lost the apicoplast organelle (Zhu et al. 2000; Abrahamsen et al. 2004). In addition, *Cryptosporidium* appears to have lost many *de novo* biosynthetic pathways, such as the capacity to synthesize amino acids and nucleotides. A recent whole-genomesequence survey for the gregarine, Ascogregarina taiwanensis supports the phylogenetic affinity of Ascogregarina with Cryptosporidium at the base of the apicomplexan clade (Templeton et al. 2010). More recently, genome sequencing of C. muris has identified the presence of mitochondrial structure and proteins that are absent from C. parvum and C. hominis but present in gregarines (Widmer and Sullivan 2012). Ascogregarina and Cryptosporidium however also possess features that unite them with the Coccidia, including an environmental oocyst stage, metabolic pathways such as the Type I fatty acid and polyketide synthetic enzymes, and a number of conserved extracellular-protein-domain architectures (Templeton et al. 2010). Future genome studies of other gregarine parasites will hopefully provide a clearer understanding of the correct taxonomic placement of the genus Cryptosporidium.

1.3 Standards for Taxonomic Status in the Genus *Cryptosporidium*

Traditionally, species delimitation within the Apicomplexa has been based on combinations of morphological features detected by light microscopy or electron microscopy (EM), unique life cycles and host specificity. Because oocysts of *Cryptosporidium* are among the smallest exogenous stages of any apicomplexan and lack distinctive morphological features to clearly differentiate *Cryptosporidium* species, morphology alone cannot be used to differentiate *Cryptosporidium* species (Fall et al. 2003).

To provide clarity, the following minimum requirements have been proposed for the naming of new species of *Cryptosporidium*: (i) morphometric studies of oocysts; (ii) genetic characterization with sequence information deposited in GenBank; (iii) demonstration of natural and, whenever feasible, at least some experimental host specificity; and (iv) compliance with International Code of Zoological Nomenclature (ICZN) (Egyed et al. 2003; Xiao et al. 2004a; Jirků et al. 2008).

1.4 Valid Species Within the Genus Cryptosporidium

To date a total of 25 species of *Cryptosporidium* have been formally described and are considered valid, including the recently described *C. viatorum* in humans and *C. scrofarum* (previously pig genotype II) in pigs (Tables 1.1, 1.2, 1.3, and 1.4 and Fig. 1.1). There are also over 40 genotypes, with a high probability that many of these will eventually be given species status with increased biological and molecular characterisation. Understanding the transmission dynamics of *Cryptosporidium* has traditionally been difficult because most species of *Cryptosporidium* are morphologically identical. Therefore, alternative molecular characterization tools such as PCR and DNA sequence analysis have been required to reliably differentiate/identify species and genotypes of *Cryptosporidium*. The 18S ribosomal RNA (rRNA) gene and the hypervariable 60-kDa glycoprotein (gp60) gene have been widely used as targets to identify species and track transmission (Xiao 2010; Plutzer and Karanis 2009; Ng et al. 2011) (Chap. 3).

1.5 Species in Marine Mammals and Fish

Very little is known about the prevalence and genetic diversity of species of *Cryptosporidium* in marine environments and the role that marine animals play in transmission of these parasites to humans. Molecular research on marine mammals has identified *C. muris* and seal genotypes 1 and 2, in ringed seals in Canada (Santín et al. 2005). Seal genotype 3 was identified in a harp seal (*Pagophilus groenlandicus*) from the Gulf of Maine (Bass et al. 2012) and a genotype similar to the *Cryptosporidium* sp. skunk genotype was identified in an Antarctic Southern elephant seal (*Mirounga leonina*) (Rengifo-Herrera et al. 2011). More recently, seal genotype 4 was identified in Weddell seals (*Leptonychotes weddellii*) (Rengifo-Herrera et al. 2013) (Table 1.1).

Cryptosporidium has been described in both fresh and marine water piscine species with parasitic stages located either on the stomach or intestinal surface, or at both sites (Table 1.1). The first account of *Cryptosporidium* in a piscine host was *Cryptosporidium nasorum*, identified in a Naso tang, a tropical fish species (Hoover et al. 1981). Hoover and colleagues also noted a similar infection in an unnamed

					-
					Cenbank
					accession
		Site of			number
Species	Host	Infection	Size $(\mu m) L \times W$	Reference	(18SrRNA)
Cryptosporidium sp.seal	Ringedseals (Phoca			Santín	AY731234
genotype 1				et al. 2005;	(AY731236-
and 2, C. muris	seals (Phoca			Bass et al. 2012	hsp70),
	vituline),				AY731235
	Hooded seal				(AY731238-
	(Cystophora				hsp70),
Cumtoconoui dirum on cool	п.,			Dans of al 2012	TNIOSOUDD
Cryptosportatum sp.scar	Па	I	I	Dass et al. 2012	6060CONT
c addining	(r agopnuus oroenlandicus)				-+00000111)
Cumtornoui dimm on	Southam alarhant			Donatfo Homoro	
Cryptosportatum sp.	Southern elephant	I	I	Kenglio-Herrera	00421420 2771422
(similar	seal (Mirounga			et al. 2011,	(GQ421420-
to skunk genotype)	leonina)			2013	hsp70)
Cryptosporidium	Weddell seals			Rengifo-Herrera	JQ740103
sp. seal	(Leptonychotes			et al. 2013	(JQ740105-
genotype 4	weddellii)				hsp70)
C. nasorum	Naso Tang(Naso	Intestine	3.6 imes 3.6	Hoover	I
	literatus)			et al. 1981	
Cryptosporidium sp.	Carp (Cyprinus carpio)	Intestine	1	Pavlásek 1983	I
Cryptosporidium sp.	Cichlid	Stomach	I	Landsberg and	I
	(Oreochromis			Paperna 1986	
	sp.)				
Cryptosporidium sp.	Brown trout (Salmo trutta)	Intestine	I	Rush et al. 1987	I
Crvntosnoridium sn.	Barramundi (Lates	Intestine	I	Glazebrook and Campbell 1987	I
. J	calcarifer)				

Table 1.1 Cryptosporidium sp. reported in marine mammals and fish

7

					GenBank
					accession
		Site of			number
Species	Host	Infection	Size $(\mu m) L \times W$	Reference	(18SrRNA)
Cryptosporidium sp.	Rainbow trout (Oncorhynchus mykiss)	Stomach	5-7	Freire-Santos et al. 1998	1
Cryptosporidium sp.	Red drum(Sciaenops ocellatus)	Stomach	7×4	Camus and Lopez 1996	I
Cryptosporidium sp.	Pleco (<i>Plecostomus</i> sp.) Intestine and Stomach	Intestine and Stomach	I	Muench and White 1997	I
Piscicryptosporidium reinchenbachklinkei	Gourami (<i>Trichogaster</i> leeri)	Stomach	$2.4 - 3.18 \times 2.4 - 3.0$	Paperna and Vilenkin 1996	I
Piscicryptosporidium- cichlidis (previously Cryptocystidium villithecum)	Cichlid (<i>Oreochromis</i> sp.)	Stomach	$4.0 - 4.7 \times 2.5 - 3.5$	Paperna and Vilenkin 1996	1
Piscicryptosporidium sp. (previously Chloromyxum-like)	Gilthead sea bream (Sparus aurata)	Stomach	I	Paperna and Vilenkin 1996	1
C. molnari	Gilthead sea bream (Sparus aurata), European sea bass (Dicentrarchus labrax), Murray cod (Maccullochella peelii peelii)	Stomach (and intestine)	4.72 (3.23–5.45) × 4.47 (3.02–5.04)	Alvarez-Pellitero and Sitj- à-Bobadilla 2002; Palenzuela et al. 2010; Barugahare et al. 2011	HM243548, HM243550, HQ585890
Cryptosporidium sp. (C. scophthalmi)	Turbot (Scophthalmus maximus)	Intestine	$\begin{array}{c} 4.44 \; (3.7{-}5.03) \times \\ 3.91 \; (3.03{-}4.69) \end{array}$	Alvarez-Pellitero et al. 2004	I

Table 1.1 (continued)

8

AY 524773 -	FJ769050 HM989833	HM989834	HM991857, JQ995776 (continued)
Ryan et al. 2004a Ziegler et al. 2007 Murphy et al. 2009	orine et al. 2012	Zanguee et al. 2010	Zanguee et al. 2010; Morine et al. 2012
4.6×4.4 - 3.4 $\times 4.1$	1 1	1	I
Stomach - Stomach	Intestine Intestine	I	1
Guppy (Poecilia Stomach reticulata) Alewife (Alosa – pseudoharengus) Angelfish (Pterophyllum Stomach	scalare) Mullet (Mugil cephalus) Intestine Golden algae eater Intestine (<i>Crossocheilus</i> <i>aymonieri</i>), Kupang damsel (<i>Chrysiptera</i> hemicyanes) Oscar fish (<i>Astronatus</i> <i>ocellatis</i>), Neon tetra (<i>Paracheirodon</i>	Angelfish (Prerophyllum scalare), Butter bream (Monodactylidae), Golden algae eater (Crossocheilus armonirer)	Guppy (<i>Poecilia</i> <i>reticulata</i>), Gourami (<i>Trichogaster</i> <i>trichopterus</i>)
Piscine genotype 1 <i>Cryptosporidium</i> sp. Piscine genotype 2	Piscine genotype 3 Piscine genotype 4	Piscine genotype 5	Piscine genotype 6, piscine genotype 6-like

1 Taxonomy and Molecular Taxonomy

Species	Host	Site of Infection	Size (μm) L \times W	Reference	GenBank accession number (18SrRNA)
C. parvum, C. parvum-like C. xiaoi and C. scrofarum	Whiting (Sillago vittata), Barramundi (Lates calcarifer)			Reid et al. 2010; Gibson-Kueh et al. 2011	1
Cryptosporidium sp.	Barramundi (<i>Late</i> scalcarifer)	Distal stomach and proximal small intestine		Gabor et al. 2011	I
Piscine genotype 7	Red eye tetra (Moenkhausia sanctaefilomenae)	I	I	Morine et al. 2012	JQ995773
Piscine genotype 8	Oblong silver biddy (Gerres oblongus)	I	I	Koinari et al. 2013	KC807985

Table 1.1 (continued)

Species/ genotype	Avian host species	Site of infection	Size (μ m) L \times W	References	GenBank accession no. 18S
C. meleagridis	Turkey (Meleagris gallopavo) (type host), Indian ring-necked parrot (Psittacula krameri), Red-legged partidge (Alectoris rufa), Cockatiels (Nymphicus hollandicus); Bohemian waxwing (Bombycilla garrulus), Rufousturle dove (Streptopelia orientalis), and fan-tailed pigeon (Columba livia), Chicken (Gallus gallus); Quails (Coturnis gallus); pigeonica), Pekin ducks (Anas pidarvhynchos)	Intestine	4.5-6.0 imes 4.2-5.3	Slavin 1955; Lindsay et al. 1989; Morgan et al. 2000a, 2001; Glaberman et al. 2001; Abe and Iseki 2004; Pagès-Manté et al. 2007; Abe and Makino 2010; Wang et al. 2010; Qi et al. 2011; Berrilli et al. 2012; Wang et al. 2012	AF112574
C. baileyi	Chicken (Gallus gallus)(type host): Brown quail (Synoicus australis) Cockatiels (Nymphicus hollandicus), Whooping crane (Grus vipio), Gray-bellied bulbul (Pycnontus spp.), Black vul- ture (Coragyps atratus), Saf- fron finch (Sicalis flaveola), mixed-bred falcons (Falco rusticolus x Falco cherrug) Ruddy Shelduck (Tadornaferruginea), Red-billed leiothrixes (Leiothrix lutea). Pekin ducks	Cloaca, bursa, trachea	Cloaca, bursa, $4.8 (4.8-5.7) \times 6.4 (5.6-7.5)$ Current et al. 1986; Morgan trachea trachea Ng et al. 2001; Abe and Iseki Ng et al. 2007; Kimura et al. van Zeeland et al. 2009; Al Makino 2010; Amer et al. 2009; Al Makino 2010; Amer et al. 2011; Sevá et al. 2011; Sevá et al. 2011; Sevá et al. 2011; Sevá et al. 2012; Warr et al. 2012; W	Current et al. 1986; Morgan et al. 2001; Abe and Iseki 2004; Ng et al. 2006; Huber et al. 2007; Kimura et al. 2004; van Zeeland et al. 2008; Nakamura et al. 2009; Abe and Makino 2010; Wang et al. 2010; Qi et al. 2011; Sevá et al. 2011; Sevá et al. 2012; Wang et al. 2011; Schulze et al. 2012; Wang et al. 2012	L19068

Species/	A	Site of	111 ····· 1 ······ 10		GenBank accession
	(Anas platyrhynchos), Buffy- fronted seedeater (Sporophila frontalis), Java sparrows (Padda oryzivora), Mynas (Acridotheres tristis), Zebra finches (Taeniopygia guttata), Crested Lark (Galerida cristata), Gouldian finch (Chloebia gouldiae), Black- billed magpie (Pica pica), Ostriches (Struthio camelus), Red-breasted Merganser (Mergus serrator), Quails				
C. galli	Chicken (Gallus gallus)(type host), Finches (Spermestidae and Fringillidae), Capercaille (Tetrao urogallus), Pine gros- beak (Pinicola enucleator), Turqoise parrots (Neophema pulchella), Cuban flamingo (Phoenicopterus ruber, Rhinocerous hornbill (Buceros rhinoceros), Red-cowled car- dinal (Paroaria dominicana), Zebra finches (Taeniopygia guttata), Chocolate parson finches (Peophila cincta), Chestnut finches (Lonchura castaneothorax), Painted		Preventriculus 8.25 (8.0–8.5) \times 6.3 (6.2–6.4) Pavlásek 1999, 2001; Ryan et al. 2003a; Ng et al. 2008; Nal Antunes et al. 2008; Nal et al. 2009; da Silva et al Qi et al. 2011; Sevá et al Qi et al. 2011; Sevá et al	Pavlásek 1999, 2001; Ryan et al. 2003a; Ng et al. 2006; Antunes et al. 2008; Nakamura et al. 2009; da Silva et al. 2010; Qi et al. 2011; Sevá et al. 2011a	AF316624, AY168847

Table 1.2 (continued)

	DQ650339	DQ650340	(continued)
	Ng et al. 2006; Nakamura et al. 2009	Meireles et al. 2006; Ng et al. 2006; Nakamura et al. 2009; Sevá et al. 2011a; Nguyen et al. 2013	
		$6.0 - 6.5 \times 4.8 - 6.6$	
firetail finches (Emblema picta), Canaries (Serinus sp.), Glosters (Serinus canaria), Hazel hen (Tetrastes bonasia rupestris), Lesser seed-finches (Oryzoborus angolensis), Rufousbellied thrush (Turdus ruffventris), Green-winged saltators (Saltator similis), Slate-coloured seedeater (Sporophila schistacea), Gold- finch (Carduelis carduelis), Great-billed seed-finch (Oryzoborus maximiliani), Ultramarine grosbeak (Cyanocompsa brissonii), Rusty-collared seedeater (Sporophila collaris), Bohe- mian waxwings (Bombycilla garrulus), silver-eared Mesia (Leiothrix argentauris)	Red Factor Canary (<i>Serinus</i> – <i>canaria</i>), Canary (<i>S. canaria</i>). Indian peafowl (<i>Pavo cristatus</i>)	 Eclectus (Eclectus roratus), Galah (Eolophus roseicapilla), Cock- atiel (Nymphicus hollandicus), Major Mitchell Cockatoo (Cacatua leadheateri, Ostriches (Struthio camelus), White-eyed parakeet (Aratinga leucophthalma) 	
	Avian Genotype I	Avian Genotype II	

					,
Species/		Site of			GenBank accession
genotype	Avian host species	infection	Size (μm) L \times W	References	no. 18S
Avian Genotype III	Galah (Eolophus roseicapilla), Cockatiel (Nymphicus hollandicus), Sun Conure	Proventriculus 7.0×6.0	7.0 imes 6.0	Ng et al. 2006; Nakamura et al. 2009; Makino et al. 2010; Gomes et al. 2012	DQ650342
	(Aratinga solstitialis), Peach- Faced Lovebirds (Agapornis roseicollis), Java sparrow (Padda oryzivora), cockatiel (Nymphicus hollandicus)		7.4×5.8 8.25×6.3	Ng et al. 2006	DQ650344
Avian Genotype IV	Japaneese whiteeye (Zosterops japonica)				
Avian Genotype V	Cockatiels (Nymphicus hollandicus)	Intestine	5.8 imes 4.5	Abe and Makino 2010; Qi et al. 2011	HM116381
Eurasian wood- cock Genotype	Eurasian woodcock (Scolopax rusticola)		8.5×6.4	Ryan et al. 2003b; Ng et al. 2006 AY273769 Morgan et al. 2001, Zhou AF316630 et al. 2004; Jellison et al. 2004	AY273769 AF316630
Duck Genotype	Black duck (Anas rubripes), Canada Geese (Branta canadensis)				
Goose Genotype I	Goose Genotype Canada Geese (Branta canadensis) I			Xiao et al. 2002a; Zhou et al. 2004; AY120912 Jellison et al. 2004	AY120912
Goose Genotype II	Goose Genotype Canada Geese (Branta canadensis) II			Zhou et al. 2004, Jellison et al. 2004	AY504512
Goose Genotype III	Goose Genotype Canada Geese (Branta canadensis) III			Jellison et al. 2004	AY324638
Goose Genotype IV	Goose Genotype Canada Geese (Branta canadensis) IV - isolate KLJ-3b				
Goose Genotype V	Goose Genotype Canada Geese (<i>Branta</i> V <i>canadensis</i>)- isolate KLJ-7			Jellison et al. 2004	AY324041

Table 1.2 (continued)

Species/ genotype	Amphibian/reptile host species	Site of infection	Size (μm) L \times W	Reference	GenBank accession number (18SrRNA)
C. fragile	Black-spined toads (Duttaphrynus melanostictus)	Stomach	$6.2 (5.5-7.0) \times 5.5 (5.0-6.5)$ Jirků et al. 2008) Jirků et al. 2008	EU162751-EU162754
C. serpentis	Amazon tree boa (Corallus Stomach hortulanus), Black rat snake (Elaphe obsoleta obsolete), Bormueller's viper (Vipera bornmueller's viper (Vipera bornmuelleri), Bull snake (Pituophis melanoleucus), California kingsnake (Lampropeltis getulus californiae), Cornsnake (Lampropeltis getulus californiae), Connsnake (Elaphe guttata guttata), Common death adder (Acamhophis antarticus), Desert monitor (Varauus griseus), Eastem/Main- land Tiger snake (Natechis scutatus), Frilled lizard (Chlamydosaurus kingui), Giant madagascar or Marchar's chonchar's cho	Stomach	$6.2 (5.6-6.6) \times 5.3 (4.8-5.6)$ Levine 1980; Kimbell et al. 1999; Morga et al. 1999a; Xiao et al. 2004b; Hajdt et al. 2004; Pedraz Díaz et al. 2001; Richter et al. 2011b; Rinaldi et al. 2012) Levine 1980; Kimbell et al. 1999; Morgan et al. 1999a; Xiao et al. 2004; Hajdusek et al. 2004; Pedraza- Díaz et al. 2011; Sevá et al. 2011; Rinaldi et al. 2012	AF151376

Species/ genotype	Amphibian/reptile host species	Site of infection	Size (μm) L \times W	Reference	GenBank accession number (18SrRNA)
	(<i>Chamaeleo oustaleti</i>), Leopard gecko (<i>Eublepharis</i> <i>macularius</i>), Mexican black kingsnake (<i>Lampropeltis getulus</i> <i>nigritus</i>), Milk snake (<i>Lampropeltis</i> <i>triangulum</i>), Mountain viper (Vipera wagneri), Python (Python molurus), Savamah monitor (Varanus exanthematicus), Skink (Mabuya perroteti), Tainan (Oxvuranus				
C. varanii	scutellatus) African fat-tailed gecko (Hemitheconyx caudicinctus) Leopard gecko (Eublepharis macularius), Boa con- strictor, Cornsnake (Elaphe guttata guttata), Leopard gecko (Eublepharis macularius), Desert monitor (Varanus griseus), Gecko	Intestine and Cloaca	4.7 (4.2–5.2) × 5 (4.4–5.6)	Pavlásek et al. 1995; Koudela and Modry 1998; Morgan et al. 1999a; Xiao et al. 2004b; Hajdusek et al. 2004; Plutzer and Karanis 2007; Pedraza- Díaz et al. 2009; Richter et al. 2011	AF112573

	AY120915	r120914	GQ504270	(continued)
	Xiao et al. 2004b; Richter AV et al. 2011	Xiao et al. 2002a, b; Alves AY120914 et al. 2005; Pedraza- Díaz et al. 2009; Griffin et al. 2010; Richter et al. 2012	Traversa et al. 2008; GG Pedraza-Díaz et al. 2009; Griffin et al. 2010; Traversa 2010; Richter et al. 2012	
		Stomach	Intestine	
(Gekkoninae sp.), Green iguana (Jguana iguana), Lampropeltis sp.; Louisiana pine snake (Pituophis ruthveni), Plated lizard (Gerrhosaurus sp.), Schneider's Skink (Eumeces schneideri), Taipan (Oxyuranus scutellatus)	Leopard gecko (Eublepharis macularius), Corn snake (Pantherophis guttatus)	ortoises one elegans), s tortoise (Tes- nanii), Ball Python regius), ortoise mys [Testudo] ii)	toise (Tes- nata), Ball thon regius) meleon), Pancake	
	Lizard genotype/ C. <i>serpentis</i> - like	Tortoise geno- type I	Tortoise geno- type II (C. duismarci)	

Species/	Amphibian/reptile host	Site of			GenBank accession
genotype	species	infection	Size (μ m) L × W	Reference	number (18SrRNA)
	tornieri), Russian tor-				
	toise (Agrionemys [Tes-				
	tudo] horsfieldii)				
Snake	New Guinea Viper boa			Xiao et al. 2002a; Kuroki	AB222185
genotype I	(Candoia asper), Japa-			et al. 2008	
	nese grass snakes (Rhabdophis tigris)				
Snake genotype	Snake genotype Boa constrictor (Boa con-			Xiao et al. 2004b	AY268584
Π	strictor ortoni)				

 Table 1.3 (continued)

	:	•			GenBank accession
Species	Major host(s)	Size of oocysts (in mm)	Site of infection	References	number (18SrRNA)
C. muris	Rodents	$6.1 (5.6-6.4) \times 8.4 (8.0-9.0),$	Stomach	Tyzzer 1907	AB089284
		8.1~(7.0-9.0) imes 5.9~(5.0-6.5)			
C. parvum	Cattle/sheep/	$4.9~(4.5-5.4)~\times~4.4~(4.2-5.2)$	Small intestine	Tyzzer 1912	AF308600
	humans				
C. wrairi	Guinea pigs	$4.6~(4.0{-}5.0) \times 5.4~(4.8{-}5.6)$	Small intestine	Vetterling et al. 1971	AF115378
C. felis	Cats	$4.5~(5.0{-}4.5) imes~5.0~(6.0{-}5.0)$	Small intestine	Iseki 1979	AF108862
C. andersoni	Cattle	$7.4~(6.0{-}8.1) imes 5.5~(5.0{-}6.5)$	Abomasum	Lindsay et al. 2000	AF093496
C. canis	Dogs	4.95 imes 4.75	Small intestine	Fayer et al. 2001	AF112576
C. hominis	Humans	$4.8~(64.4{-}5.4)~\times~5.2~(4.4{-}5.9)$	Small intestine	Morgan-Ryan et al. 2002	AF108865
C. suis	Pigs	$4.6~(4.9-4.4) \times 4.2~(4.0-4.3)$	Small and large	Ryan et al. 2004b	AF115377
			intestine		
C. bovis	Cattle	$4.89(4.76-5.35) \times 4.63(4.17-4.76)$	Small intestine	Fayer et al. 2005	AY741305
C. fayeri	Marsupials	$4.9(4.5-5.1) \times 4.3(3.8-5.0)$	Small intestine	Ryan et al. 2008	AF159112
C. macropodum Marsupials	Marsupials	$4.9~(4.5{-}6.0) imes~5.4~(5.0{-}6.0)$	Small intestine	Power and Ryan 2008	AF513227
C. ryanae	Cattle	$3.16(2.94 - 4.41) \times 3.73(2.94 - 3.68)$	Small Intestine	Fayer et al. 2008	AY587166
C. xiaoi	Sheep	$3.94 (2.94 - 4.41) \times 3.44 (2.94 - 4.41) \mu m$	Small intestine	Fayer and Santín 2009	EU408314
C. ubiquitum	Sheep/wildlife	$5.19(4.92 - 5.61) \times 4.87(4.45 - 5.44)$	Small intestine	Fayer et al. 2010	AF262328
C. cuniculus	Rabbits	$5.98(5.55-6.40) \times 5.38(5.02-5.92)$	Small intestine	Robinson et al. 2010; (Inman and Takeuchi 1979)	FJ262725
C. tyzzeri	Mice	$4.64\pm0.05 \ \mu m imes 4.19\pm0.06$	Small intestine	Ren et al. 2012	AF112571
C. viatorum	Humans	$5.35(4.87 - 5.87) \times 4.72(4.15 - 5.20)$	Small intestine	Elwin et al. 2012b	HM485434
C. scrofarum	Pigs	$5.16(4.81 - 5.96) \times 4.83(4.23 - 5.29)$	Small Intestine	Kváč et al. 2013	EU331243

Table 1.4 Mammalian Cryptosporidium species

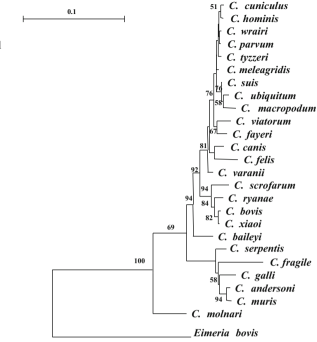


Fig. 1.1 Phylogenetic relationships of currently accepted *Cryptosporidium* species at the 18S ribosomal RNA locus inferred by neighbour-joining analysis based on genetic distances calculated using Kimura-2 parameters. Bootstrap values (>60 %) from 1,000 pseudoreplicates are shown

species of marine fish (Hoover et al. 1981). Levine (1984) suggested that *C. nasorum* should be given the status of species. However, lack of viable oocyst measurements and taxonomically valid diagnostic features and the fact that only developmental stages on the intestinal microvillous surface were described, has resulted in *C. nasorum* being considered a nomen nudem (i.e., a name that is invalid because an insufficient description was published) (Ryan et al. 2004a; Xiao et al. 2004a).

Cryptosporidium molnari was described in gilthead sea bream (*Sparus aurata*) and European sea bass (*Dicentrarchus labrax*) in 2002 (Alvarez-Pellitero and Sitjà-Bobadilla 2002) (Table 1.1). *Cryptosporidium molnari* primarily infects the epithelium of the stomach and seldom the intestine (Alvarez-Pellitero and Sitjà-Bobadilla 2002). In 2004 *Cryptosporidium scophthalmi* was described in turbot (*Psetta maxima, syn. Scophthalmus maximus*) and mainly infects the epithelium of the intestine and very seldom the stomach (Alvarez-Pellitero et al. 2004). An unusual feature of both parasites is that while merogonial and gamogonial stages appeared in the typical extracytoplasmic position, oogonial and sporogonial stages were located deeply within the epithelium (Alvarez-Pellitero and Sitjà-Bobadilla 2002; Alvarez-Pellitero et al. 2004). *Cryptosporidium molnari* has recently been characterised genetically (Palenzuela et al. 2010), but no DNA sequences are available for *C. scophthalmi*. Until genetic sequences are provided for *C. scophthalmi*, it cannot be considered a valid species because of the high genetic

heterogeneity and morphological similarity among *Cryptosporidium* species and genotypes in fish.

A total of 13 additional species/genotypes have been identified in fish using molecular tools; piscine genotype 1 from a guppy (Poecilia reticulate) (Ryan et al. 2004a); piscine genotype 2 from a freshwater angelfish (*Pterophyllum scalare*) (Murphy et al. 2009), piscine genotype 3 from a sea mullet (*Mugil cephalus*) (Reid et al. 2010); piscine genotype 4 from a golden algae eater (Crossocheilus aymonieri), a kupang damsel (Chrysiptera hemicyanes) and an oscar fish (Astronatus ocellatis); piscine genotype 5 from an angelfish (Pterophyllum scalare), a butter bream (Monodactylidae) and a golden algae eater (Crossocheilus aymonieri); piscine genotype 6 from a guppy (Poecilia reticulate) (Zanguee et al. 2010); piscine genotype 7 from red eye tetra (Moenkhausia sanctaefilomenae) (Morine et al. 2012), piscine genotype 8 from Oblong silver biddy (Gerres oblongus) (Koinari et al. 2013); C. parvum, C. xiaoi and C. scrofarum in whiting (Sillago vittata) (Reid et al. 2010); C. parvum in Nile tilapias (Oreochromis niloticus) and a Silver barb (Puntius gonionotus) and C. parvum and C. hominis in mackerel scad (Decapterus macarellus) from Papua New Guinea (Koinari et al. 2013) and rat genotype III in a goldfish (Carassius auratus) (Morine et al. 2012).

The prevalence of Cryptosporidium in fish ranges from 0.8 % to up to 100 % mostly among juvenile fish (Landsberg and Paperna 1986; Sitjà-Bobadilla et al. 2005; Alvarez-Pellitero et al. 2004, 2009; Murphy et al. 2009; Reid et al. 2010; Zanguee et al. 2010; Barugahare et al. 2011). The pathogenesis of Cryptosporidium in fish has not been extensively studied; however C. molnari has been associated with clinical signs in gilthead sea bream, consisting of whitish feces, abdominal swelling and ascites which resulted in mortalities in fingerlings (Alvarez-Pellitero and Sitjà-Bobadilla 2002). Another study, which examined interactions between bacteria and C. molnari in gilthead sea bream under farm and laboratory conditions reported that the ability of gilthead sea bream to survive an additional bacterial infection may be compromised where a previous mixed infection of bacteria and C. molnari existed (Sitjà-Bobadilla et al. 2006). Piscine genotype 1 was associated with high mortalities amongst guppies and was detected in the stomach, with oogonial and sporogonial stages observed deep within the epithelium, similar to C. molnari (Ryan et al. 2004a). Previous studies on piscine genotype 2, which was identified in a hatchery, revealed that infected fish exhibited variable levels of emaciation, poor growth rates, swollen coelomic cavities, anorexia, listlessness and increased mortality (Murphy et al. 2009). In affected fish, large numbers of protozoa were identified both histologically and ultrastructurally associated with the gastric mucosa. A high prevalence of Cryptosporidium was associated with high mortalities in immature hatchery barramundi (Lates calcarifer) in Australia (Gabor et al. 2011).

In 1996, Paperna and Vilenkin proposed a new genus, designated *Piscicryptosporidium*, for *Cryptosporidium*-like species infecting a number of piscine hosts. The genus included two species, *P. reichenbachklinkei* and *P. cichlidaris* previously described as *Cryptosporidium* sp. from cichlid fishes of the genus

Oreochromis (Paperna and Vilenkin 1996). Several unique features were cited to support the genus including the covering of the surface of the parasitophorous sac by rudimentary microvilli and the localisation of the oocysts deep within the gastric mucosa (Paperna and Vilenkin 1996). However, these apparently differential features have also been described in some mammalian *Cryptosporidium* spp. For example, *C. parvum* has been occasionally found within some cells (Beyer et al. 2000; Marcial and Madara 1986) and microvilli are usually retained in different mammalian species (Alvarez-Pellitero and Sitjà-Bobadilla 2002). Recently, *P. reichenbachklinkei* and *P. cichlidaris* were published as *Cryptosporidium reichenbachklinkei* and *C. cichlidaris* (Jirků et al. 2008), with the comment that the genus was considered a synonym of *Cryptosporidium*.

No molecular data was provided by Paperna and Vilenkin, (1996) to support the genus or species, however more recent characterization of *C. molnari* and piscine genotypes 1–7 indicate that piscine-derived species and genotypes of *Cryptosporidium* are genetically very distinct and primitive to all other species (Ryan et al. 2004a; Palenzuela et al. 2010; Reid et al. 2010; Zanguee et al. 2010; Barugahare et al. 2011; Morine et al. 2012). However, further studies at additional loci are required to validate whether or not *Piscicryptosporidium* is a valid genus for piscine species.

1.6 Species in Birds

Currently only three avian *Cryptosporidium* spp. are recognised; *Cryptosporidium* meleagridis, *Cryptosporidium baileyi* and *Cryptosporidium galli*. Two other species of *Cryptosporidium* have been named from birds: *Cryptosporidium anserinum* from a domestic goose (*Anser domesticus*) (Proctor and Kemp 1974) and *Cryptosporidium tyzzeri* from chickens (*Gallus gallus domesticus*) (Levine 1961). Neither of these reports gave adequate description of oocysts or provided other useful information and are therefore not considered valid species (Lindsay and Blagburn 1990). Naturally occurring cryptosporidiosis in birds manifests itself in three clinical forms; respiratory disease, enteritis and renal disease. Usually only one form of the disease is present in an outbreak (Lindsay and Blagburn 1990).

Tyzzer first described avian cryptosporidiosis in 1929 (Tyzzer 1929) but it was not until 1955 that Slavin found a structurally similar parasite in the ileum of turkey poults and named the parasite *C. meleagridis* (Slavin 1955) (Table 1.2). Since then *C. meleagridis* has been characterized genetically and has been detected in many avian hosts (Table 1.2). It is the third most prevalent species infecting humans (Leoni et al. 2006; Elwin et al. 2012a). In some studies, *C. meleagridis* prevalence is similar to that of *C. parvum* (Gatei et al. 2002a; Cama et al. 2007). The ability of *C. meleagridis* to infect humans and other mammals, and its close relationship to *C. parvum* and *C. hominis* at the 18S rRNA, actin and 70 kDa heat shock protein (HSP70) loci, has led to the suggestion that mammals actually were the original hosts, and that the species has later adapted to birds (Xiao et al. 2002a, 2004a). Recently, sequence analysis of the 18S gene and HSP70 loci, has been used to provide evidence of zoonotic transmission of *C. meleagridis* from chickens to a human on a Swedish farm (Silverlås et al. 2012).

A second species of avian *Cryptosporidium*, originally isolated from commercial broiler chickens, was named *C. baileyi* based on its life cycle and morphologic features (Current et al. 1986). Natural *C. baileyi* infections have been reported in many anatomic sites in avian hosts including the conjunctiva, nasopharynx, trachea, bronchi, air sacs, small intestine, large intestine, ceca, cloaca, bursa of fabricius, kidneys and urinary tract, but is more frequently associated with respiratory cryptosporidiosis (Lindsay and Blagburn 1990). *Cryptosporidium baileyi* is probably the most common avian *Cryptosporidium* species and has been reported in a wide range of avian hosts (Table 1.2). The relationship between *C. baileyi* and the rest of the intestinal and gastric parasites is unclear as phylogenetic analysis at the SSU rRNA, actin and COWP loci group *C. baileyi* with the intestinal parasites but analysis at the HSP70 locus placed *C. baileyi* in a cluster that contained all gastric *Cryptosporidium* instead of the intestinal parasite cluster (Xiao et al. 2002a).

A third species of avian *Cryptosporidium* was first described by Pavlásek (Pavlásek 1999, 2001) in hens on the basis of biological differences. More recently, the parasite was re-described on the basis of both molecular and biological differences (Ryan et al. 2003a). Unlike other avian species, life cycle stages of *C. galli* developed in epithelial cells of the proventriculus and not the respiratory tract or small and large intestines (Pavlásek 1999, 2001). *Cryptosporidium galli* is the most prevalent *Cryptosporidium* species in Passeriformes where it frequently causes chronic infection (da Silva et al. 2010). Natural *C. galli* infections have been reported in a wide range of species (Table 1.2).

Blagburn et al. (1990) may also have detected *C. galli* in birds when they used light and electron microscopy to characterize *Cryptosporidium* sp. in the proventriculus of an Australian diamond firetail finch that died of acute diarrhea. A subsequent publication also identified a species of *Cryptosporidium* infecting the proventriculus in finches and inadvertently proposed the name *Cryptosporidium* blagburni in Table 1.1 of the paper (Morgan et al. 2001). However, Pavlásek (Pavlásek 1999, 2001) had provided a detailed description of what appeared to be the same parasite and named it *C. galli* and molecular analyses have revealed *C. galli* and *C. blagburni* to be the same species (Ryan et al. 2003a). Distance, parsimony and maximum likelihood analysis of 3 loci (18S rRNA, HSP70 and actin) identified *C. galli* as a distinct species (Ryan et al. 2003a). The gastric location of *C. galli* in the host and its large size suggest that it is most closely related to the other gastric *Cryptosporidium* species and this has been supported by molecular analysis.

In addition to the three recognized species of *Cryptosporidium*, 11 genotypes: the avian genotypes I–V, the black duck genotype, the Eurasian woodcock genotype and goose genotypes I–IV have been reported (Table 1.2). These genotypes may be renamed as distinct species in the future once more biological and molecular data become available. In addition, *C. hominis, C. parvum, C. serpentis, C. muris, C. andersoni* and muskrat genotype I have also been identified in a small

number of birds, most of which were probably the results of accidental ingestion of oocysts of these organisms (Ryan et al. 2003a; Zhou et al. 2004; Ng et al. 2006; Chatterjee 2007; Zylan et al. 2008; Jellison et al. 2007, 2009; McEvoy and Giddings 2009; Quah et al. 2011; Gomes et al. 2012).

1.7 Species in Amphibians

The first report of *Cryptosporidium* in amphibians was the description of oocysts in the feces of a captive mice-fed ornate horned frog, *Ceratophrys ornata* (Crawshaw and Mehren 1987). The identity of the species is unknown and, as mice can be infected with several species and genotypes of *Cryptosporidium*, it remains unknown if the frog was naturally infected or passing oocysts from infected mice. Currently the only accepted species is *Cryptosporidium fragile*, which was described from the stomach of naturally infected black-spined toads (*Duttaphrynus melanostictus*) from the Malay peninsula (Jirků et al. 2008). Developmental stages were confined to the surface of gastric epithelial cells. In transmission experiments, *C. fragile* was not infective for one fish species (*Poecilia reticulate*), four amphibian species (*Bufo bufo, Rana temporaria, Litoria caerulea, Xenopus laevis*), one species of reptile (*Pantherophis guttatus*) and SCID mice. Phylogenetic analysis of the full length 18S rRNA revealed *C. fragile* to be genetically distinct and grouped with the gastric species.

1.8 Species in Reptiles

Cryptosporidium infections are common in reptiles and have been reported in more than 57 reptilian species (O'Donoghue 1995; Xiao et al. 2004b). Unlike in other animals in which *Cryptosporidium* infection is usually self-limiting in immuno-competent individuals, cryptosporidiosis in reptiles is frequently chronic and sometimes lethal in snakes. *Cryptosporidium serpentis* was named by Levine (1980) based on a report of hypertrophic gastritis in four species of snakes by Brownstein et al. (1977), but was a nomen nudum until Tilley et al. (1990) provided supporting morphologic and biological data. Since then it has been reported in lizards and other reptiles (Table 1.3). *Cryptosporidium serpentis* has also been identified in cattle (Azami et al. 2007; Chen and Qiu 2012).

Cryptosporidium varanii was described by Pavlásek et al. (1995) from an Emerald Monitor (*Varanus prasinus*) in the Prague Zoo, based on oocyst morphology, histology of endogenous stages in the intestine, and failure of oocysts to infect mice. It was subsequently genetically characterized and identified in other lizards and in snakes (Pavlásek and Ryan 2008) (Table 1.3). *Cryptosporidium saurophilum* was named by Koudela and Modry (1998) from a Schneider's skink (*Eumeces schneideri*). Cross-transmission experiments showed that the species was

transmissible to lizards but not to snakes and mice. Molecular characterization revealed *C. saurophilum* and *C. varanii* were the same species (Xiao et al. 1999b, 2004b; Pavlásek and Ryan 2008) and therefore *C. varanii* took precedence over *C. saurophilum* based on its earlier publication date (Pavlásek and Ryan 2008).

Tortoise genotype I has been identified in the feces of Indian star tortoises (*Geochelone elegans*), Hermann's tortoises (*Testudo hermanii*) and a ball python (*Python regius*) (Xiao et al. 2004b; Alves et al. 2005; Pedraza-Díaz et al. 2009; Richter et al. 2012). In 2008, Traversa et al. characterized a second tortoise genotype from the feces of a marginated tortoise (*Testudo marginata*) (*Cryptosporidium* sp. ex *Testudo marginata* CrIT-20). The same genotype was subsequently identified in a chameleon (*Chamaeleo calyptratus*), a python (*Python regius*), a Pancake tortoise (*Malacochersus tornieri*), a Russian tortoise (*Agrionemys [Testudo] horsfieldii*) and Hermann's tortoises (Pedraza-Díaz et al. 2009; Griffin et al. 2010; Richter et al. 2012).

In 2010, Traversa proposed the name *Cryptosporidium ducismarci* for this second tortoise genotype (Traversa 2010), however the manuscript lacks the traditional formal descriptions associated with naming a new parasite species including a description of the oocysts. While genetic evidence suggests that is likely to be a separate species, *C. ducismarci* cannot be considered a valid species until it is formally described and should therefore be referred to as tortoise genotype II. Tortoise genotype II has been identified in gastric mucosa in a Russian tortoise, while tortoise genotype II has been identified in intestinal lesions in a Russian tortoise and a Pancake tortoise (Griffin et al. 2010). Phylogenetically, tortoise genotype II is closer to *C. serpentis* (Griffin et al. 2010).

Other reptile-associated genotypes include snake genotype 1 which was identified in a New Guinea Viper boa (*Candoia asper*) and Japanese grass snakes (*Rhabdophis tigris*) (Xiao et al. 2002b; Kuroki et al. 2008). Snake genotype II has been identified in a Boa constrictor (*Boa constrictor ortoni*) (Xiao et al. 2004b). *Cryptosporidium parvum, C. tyzzeri* (mouse genotype I) and *C. muris* have also been identified in snake and lizard feces. It is unclear whether these species are infecting the animals or passing through from ingested prey (Upton et al. 1989; Xiao et al. 2004b; Morgan et al. 1998, 1999a and Traversa et al. 2008). Avian genotype V has also been detected in two green iguanas (*Iguana iguana*) (Kik et al. 2011). In that study, *Cryptosporidium* developmental stages were identified on the intestinal epithelium and the infection was associated with cloacal prolapse and cystitis (Kik et al. 2011).

1.9 Species in Mammals

Over 150 species belonging to 12 mammalian Orders have been reported as hosts for *Cryptosporidium* species (Fayer 2008, 2010). However, the majority of these reports were based on microscopic observations of oocysts and lack data necessary

for species designation. Currently, 18 species of *Cryptosporidium* are recognized in mammals, *C. muris, C. parvum, C. wrairi, C. felis, C. andersoni, C. canis, C. hominis, C. suis, C. bovis, C. fayeri, C. macropodum, C. ryanae, C. xiaoi, C. ubiquitum, C. cuniculus, C. tyzzeri, C. viatorum and C. scrofarum.* All of these species have been identified using combinations of morphologic, biological, and molecular data. Many of these species will be discussed in greater depth in subsequent chapters and therefore they are briefly discussed here in chronological order of their dates of publication (Table 1.4). In addition to named species, more than 28 *Cryptosporidium* genotypes have been described in mammals including horse, hamster, ferret, skunk, squirrel, bear, deer, fox, mongoose, hedgehog, wildebeest, muskrat I and II, opossum I and II, chipmunk I to III, rat I to V and deer mouse I to IV (Fayer 2010; Xiao et al. 2004a). For these genotypes to receive taxonomic status, sufficient morphological, biological, and molecular data are required and names must comply with the rules of the International Code for Zoological Nomenclature (ICZN).

Cryptosporidium muris, the type species of Cryptosporidium, was described as a protozoan parasite in the gastric glands of the stomach of the common mouse (Tyzzer 1907, 1910). Transmission was successful to other mice but not to a rat. Nearly 80 years later (Iseki et al. 1989), reported that oocysts from rats (C. muris strain RN66) were infectious to SPF laboratory rats, mice, guinea pigs, rabbits, dogs, and cats. Stages developed in the stomach and all hosts excreted oocysts (Iseki et al. 1989). Molecular data at multiple loci have confirmed the genetic distinctness of C. muris, which groups phylogenetically with the other gastric parasites (C. andersoni, C. galli, C. serpentis) (Xiao et al. 1999a; Morgan et al. 2000b). Cryptosporidium muris is well known to have broad host specificity. In addition to various rodent species, natural C. muris infections have been documented for pigs, Bactrian camels, giraffes, dogs, cats, nonhuman primates, seals, bilbies, and tawny frogmouth (Fayer 2010; Feng et al. 2011). It has also been reported in humans (Katsumata et al. 2000; Guyot et al. 2001; Gatei et al. 2002b, 2003, 2006; Tiangtip and Jongwutiwes 2002; Palmer et al. 2003; Muthusamy et al. 2006; Al-Brikan et al. 2008).

Cryptosporidium parvum was described as a coccidium in the small intestine of the common mouse (Tyzzer 1912). Subsequently, over 150 mammals were reported hosts of *Cryptosporidium* based on finding oocysts resembling *C. parvum* in their feces (Fayer 2010). The advent of molecular typing, however, has revealed that many *Cryptosporidium* species and genotypes have oocysts that are indistinguishable from *C. parvum* and therefore molecular tools are necessary to confirm the identity of a species. The terms *C. parvum* bovine genotype, genotype II, and genotype B have been used in the past to distinguish this species from *C. hominis*. The *C. parvum* genome project (Abrahamsen et al. 2004) and numerous phylogenetic analyses at >15 loci have confirmed the genetic uniqueness of *C. parvum* (Fayer 2010). *Cryptosporidium parvum* primarily infects cattle, sheep and humans but has been reported in a wide range of hosts (Xiao and Ryan 2008; Fayer 2010).

Cryptosporidium wrairi, first described as a coccidian in guinea pigs (*Cavia porcellus*) (Jervis et al. 1966), was named by Vetterling et al. (1971) as an acronym

for the Walter Reed Army Institute of Research. Endogenous stages, but not oocysts, were described from guinea pigs with enteritis. Molecular studies at numerous loci have confirmed that it is genetically distinct (Xiao et al. 2004b). Infections in nature, found only in guinea pigs, strongly suggest that *C. wrairi* is host specific.

Cryptosporidium felis oocysts were first described from the feces of cats (Iseki 1979). Oocysts failed to infect mice, rats, guinea pigs and dogs (Asahi et al. 1991). *Cryptosporidium felis* has also been detected in cattle and in humans (Bornay-Llinares et al. 1999; Raccurt 2007; Lucio-Forster et al. 2010; Cieloszyk et al. 2012; Elwin et al. 2012a; Gherasim et al. 2012; Insulander et al. 2012). Molecular analysis confirmed the genetic distinctness *C. felis* (Xiao et al. 2004b).

Cryptosporidium andersoni infects the abomasum of cattle (*Bos taurus*) and produces oocysts that are morphologically similar to, but slightly smaller than those of *C. muris* and was originally mistakenly identified in cattle as *C. muris* based on its oocyst size. In 2000, it was described as a new species based on location of endogenous stages in the abomasum, its host range, and genetic distinctness at multiple loci (Lindsay et al. 2000). Oocysts were not infectious to outbred, inbred immunocompetent, or immunodeficient mice, nor to chickens or goats (Lindsay et al. 2000). More recently, a novel *C. andersoni* genotype from Japan has been reported to infect SCID mice (Satoh et al. 2003; Koyama et al. 2005; Matsubayashi et al. 2005). In contrast to *C. muris, C. andersoni* is mostly a parasite of cattle, having been found only occasionally in other animals such as Bactrian camels, sheep, and goats (Fayer 2010; Feng et al. 2007; Waldron et al. 2011).

Cryptosporidium canis (previously dog genotype 1) was first identified as the dog genotype by Xiao et al. (1999b) and described as a species in 2001 (Fayer et al. 2001) on the basis that *C. canis* oocysts were infectious for calves but not mice and were genetically distinct from all other species. *Cryptosporidium canis* and its sub-genotypes (*C. canis* fox genotype and *C. canis* coyote genotype) have been reported in dogs, foxes and coyotes (Fayer 2010). *Cryptosporidium canis* has also been reported worldwide in humans (Lucio-Forster et al. 2010; Fayer 2010; Elwin et al. 2012a).

Cryptosporidium hominis, a pathogenic species found primarily in humans and originally referred to as the *C. parvum* human genotype, genotype 1, and genotype H was named a new species based on molecular (16 genetic loci) and biological differences between isolates from human and bovine sources (Morgan-Ryan et al. 2002). Under experimental conditions *C. hominis* was non-infective for mice, rats, cats, dogs, and cattle but infective for calves, lambs, and piglets (Morgan-Ryan et al. 2002; Xiao et al. 2002b). Natural infections have been reported in a dugong, cattle, goats and marsupials (Morgan et al. 2000c; Park et al. 2006; Fayer 2010; Abeywardena et al. 2012; Ryan and Power 2012).

Cryptosporidium suis, previously recognized as pig genotype I (Morgan et al. 1998), was named a new species based on analysis of the 18S rDNA, HSP70, and actin loci (Ryan et al. 2004b). It is not infectious for mice (Morgan et al. 1999b) and poorly infectious for cattle (Enemark et al. 2003). It has been

reported in naturally infected pigs worldwide, but causes only mild or no clinical signs in pigs (Enemark et al. 2003; Zintl et al. 2007). It has been reported in four humans (Xiao et al. 2002a; Leoni et al. 2006; Cama et al. 2007; Wang et al. 2013) and a *C. suis*-like genotype has been reported in cattle (Geurden et al. 2006; Fayer et al. 2006; Khan et al. 2010; Ng et al. 2012) and rodents (Ng-Hublin et al. 2013).

Cryptosporidium bovis from calves, identified as *Cryptosporidium* genotype bovine B (Xiao et al. 2002b) was named a new species based on the fact that it was not transmissible to neonatal BALB/c mice or lambs in transmission experiments and based on its genetic distinctness at the 18S rRNA, HSP70, and actin loci (Fayer et al. 2005). This species primarily infects young cattle and has a wide geographic distribution (Feng et al. 2007). It is closely related to *C. ryanae* (Fayer et al. 2005).

Cryptosporidium fayeri (previously marsupial genotype I) was first described in a red kangaroo (*Macropus rufus*) in 1988 (Morgan et al. 1998) and was formally described as a species in 2008 (Ryan et al. 2008). *Cryptosporidium fayeri* oocysts are indistinguishable from *C. parvum* but were found to be infectious for neonatal ARC Swiss mice (Ryan et al. 2008). Its host range is largely confined to marsupials with the exception of one report in sheep (Ryan et al. 2005) and a recent report in a 29-year-old immunocompetent woman who suffered prolonged gastrointestinal illness in Sydney (Waldron et al. 2010). Identical subtypes were found in marsupials in the area (Waldron et al. 2010). *Cryptosporidium fayeri* is genetically very closely related to the opossum genotype I identified in North American opossums (*Didelphis virginiana*), which is considered a sub-genotype of *C. fayeri* (Ryan et al. 2008).

Cryptosporidium macropodum (previously marsupial genotype II or EGK3) was first identified in eastern grey kangaroos (*Macropus giganteus*) in 2003 (Power et al. 2004) and was named a new species in 2008 (Power and Ryan 2008). It has only been reported in marsupial hosts. It is genetically distinct and not closely related to *C. fayeri* but rather more closely related to *C. suis* (Power and Ryan 2008). Both *C. fayeri* and *C. macropodum* appear to be host adapted to marsupial species as demonstrated by absence of clinical signs in marsupial hosts despite excretion of high oocyst numbers (Power and Ryan 2008).

Cryptosporidium ryanae (previously deer-like genotype) was named a new species based on transmission and genetic data in 2008 (Fayer et al. 2008). The deer-like genotype has never been found in deer but was called "deer-like" because its 18S rRNA sequence was very similar to the deer genotype (Xiao et al. 2002a). Oocysts of *C. ryanae* are not infectious for BALB/c mice or lambs (Fayer et al. 2008). Together with *C. bovis, C. ryanae* is responsible for the majority of *Cryptosporidium* infections in post-weaned calves. It was the only species identified in zebu cattle and water buffaloes in a recent study in Nepal (Feng et al. 2012).

Cryptosporidium xiaoi (previously *C. bovis*-like or *C. bovis*) from sheep was initially identified by Chalmers et al. in 2002 and was formally described as a species in 2009 (Fayer and Santín 2009). *Cryptosporidium xiaoi* mainly infects sheep and appears to be asymptomatic but has been reported in yaks, goats, fish and

kangaroos (Fayer and Santín 2009; Reid et al. 2010; Yang et al. 2011; Rieux et al. 2013). *Cryptosporidium xiaoi* is genetically distinct but very closely related to *C. bovis* (Fayer and Santín 2009).

Cryptosporidium ubiquitum (previously cervine genotype, cervid, W4 or genotype 3) was first identified by Xiao et al. (2000) in storm water samples in lower New York State (storm water isolate W4, GenBank accession no. AF262328). Subsequently, Perez and Le Blancq (2001) identified this genotype in white tailed deer-derived isolates from lower New York State and referred to it as genotype 3. Since then it has been described in a wide variety of hosts worldwide including humans and was formally described as a species in 2010 (Fayer et al. 2010). *Cryptosporidium ubiquitum* infects the greatest number of host species of any species of *Cryptosporidium* that has been substantiated by molecular testing and is one of the main *Cryptosporidium* species infecting sheep (along with *C. xiaoi*). It is also commonly found in humans (Fayer et al. 2010; Cieloszyk et al. 2012; Elwin et al. 2012a). It is genetically very distinct and is most closely related to *C. suis* (Fayer et al. 2010).

Cryptosporidium cuniculus (previously rabbit genotype) was first described in rabbits by Inman and Takeuchi (1979), who described the microscopic detection and ultra-structure of endogenous Cryptosporidium parasites in the ileum of an asymptomatic female rabbit. The rabbit genotype was first identified in rabbits from the Czech Republic (Ryan et al. 2003a) and C. cuniculus was formally re-described as a species in 2010 (Robinson et al. 2010). Cryptosporidium cuniculus oocysts were infectious for weanling rabbits, immunosuppressed Mongolian gerbils and immunosuppressed Porton mice but not neonatal mice (Robinson et al. 2010). Cryptosporidium cuniculus has a close genetic relationship with C. hominis with limited differences at the 18S rRNA, HSP70 and actin genes (0.51 %, 0.25 % and 0.12 %, difference, respectively) and is known to infect humans. In 2008, it was responsible for a human cryptosporidiosis outbreak in the UK (Chalmers et al. 2009), which has raised considerable awareness about the importance of investigating rabbits in drinking water catchments as a source of *Cryptosporidium* transmissible to humans. A recent study in the UK reported that C. cuniculus was the third most commonly identified Cryptosporidium species in patients with diarrhea (Chalmers et al. 2011). Cryptosporidium cuniculus has also been identified in a human patient in France and in children in Nigeria (Anon 2010; Molloy et al. 2010).

Cryptosporidium tyzzeri (previously mouse genotype I) was first reported in mice in Australia (Morgan et al. 1998) and was named a new species in 2012 (Ren et al. 2012). It has a wide geographic distribution and has been found in various countries, including the United States, Australia, the United Kingdom, Spain, Portugal, Poland, China, and Japan (cf. Morgan et al. 1999b; Ren et al. 2012). *Cryptosporidium tyzzeri* has smaller oocysts than *Cryptosporidium parvum* (see Table 1.4) and is not infectious for calves and lambs (Ren et al. 2012). It is also not infectious for pigs (Kváč et al. 2012). Unlike *C. parvum* in mice, *C. tyzzeri* does not appear to infect the colon (Ren et al. 2012). Although phylogenetically related to *C. parvum* (which infects mainly ruminants and humans), *C. tyzzeri* is genetically

distinct from *C. parvum* and mostly infects domestic mice and small rodents but has been recently been reported in humans (Rasková et al. 2013).

Cryptosporidium viatorum was recently described from human travelers returning to Great Britain from India, Nepal, Bangladesh and Pakistan (Elwin et al. 2012b). Natural infection results in diarrhoea, abdominal pain, fever and occasionally nausea and/or vomiting. As with most intestinal *Cryptosporidium* species, *C. viatorum* oocysts are indistinguishable from *C. parvum* and measure $5.35 \times 4.72 \,\mu$ m with a length to width ratio of 1.14 (Elwin et al. 2012b). *Cryptosporidium viatorum* is phylogenetically most similar to *Cryptosporidium fayeri* (<98 % similarity) at the 18S rRNA, HSP70 and actin genes (Elwin et al. 2012b). It has been also identified in two human patients in Sweden (Insulander et al. 2012). No other natural hosts of *C. viatorum* have been reported and potential reservoirs remain unknown.

Cryptosporidium scrofarum (previously pig genotype II) was first reported in pigs in Australia (Ryan et al. 2003c) and was formally described as a species in pigs in 2013 (Kváč et al. 2013). *Cryptosporidium scrofarum* oocysts were not infectious for adult SCID mice, adult BALB/c mice, Mongolian gerbils (*Meriones unguiculatus*), Southern multimammate mice (*Mastomys coucha*), yellow-necked mice (*Apodemus flavicollis*), or guinea pigs (*Cavia porcellus*). Phylogenetic analysis indicates that it is most closely related to *C. ryanae*, *C. xiaoi*, and *C. bovis* (Kváč et al. 2013). It has been reported in domestic pigs in eight countries and is prevalent in adult pigs, generally asymptomatic and with a generally low infection intensity (Kváč et al. 2013). In addition, wild boars, calves, marine fish, rodents and humans have been reported as natural hosts of this species (Kváč et al. 2009a, b, c; Němejc et al. 2012; Ng et al. 2011; Reid et al. 2010; Ng-Hublin et al. 2013). Evidence suggests that *C. suis* infects pigs of all ages, whereas *C. scrofarum* only infects pigs >6 week of age (Jeníková et al. 2011).

1.9.1 The C. tyzzeri Versus C. pestis Debate

In 2006, Slapeta proposed in an opinion article to rename *C. parvum* as *Cryptosporidium pestis* and retain *C. parvum* for mouse genotype I (now *C. tyzzeri*) (Slapeta 2006). This was largely based on a rigid interpretation of the Principle of Priority of the ICZN as Slapeta maintained that when Tyzzer described *C. parvum* in 1912 in mice (Tyzzer 1912), it is probable that he was in fact describing *C. tyzzeri*. This led to a debate in the literature as to the validity of *C. pestis* (Xiao et al. 2007). The debate was re-ignited in 2012, when Ren et al. re-named mouse genotype I as *C. tyzzeri*, the validity of which was disputed by Slapeta (Ren et al. 2012; Slapeta 2011; Xiao et al. 2012).

The dispute on the validity of *C. pestis* and *C. tyzzeri* originates from the uncertainty of the identity of *C. parvum* described by Tyzzer in 1912 and on the interpretation of the Principle of Priority of the ICZN. Because most animals are infected with multiple species and genotypes of *Cryptosporidium* spp., we will never know what Tyzzer originally described. It may have been *C. tyzzeri*, but may

also have been any of the other *Cryptosporidium* species and genotypes that have been found in naturally infected domestic mice (Ren et al. 2012), including mouse genotype II and *C. parvum*. In addition, the ICZN clearly states that the Principle of Priority is to be used to promote stability and is not intended to upset a longaccepted name (Ride et al. 1999). Importantly, in 1985, Upton et al. re-described *C. parvum* specifically for bovine and human isolates (Upton and Current 1985). Because mice are known to be naturally infected with *C. parvum* and have been used widely as a laboratory model for *C. parvum*, the 1985 taxonomic re-description of *C. parvum* was done following ICZN rules. Prior to this re-description, the name was seldom used but since then the name *C. parvum* has been widely accepted by almost all researchers in the community (a recent PubMed search of *C. parvum* found no publications before 1985 and over 3,240 publications since 1985).

The designation of *C. tyzzeri* for the mouse genotype I, brings further clarity to the taxonomy of *Cryptosporidium* spp. in humans, cattle, and domestic mice (Ren et al. 2012). Slapeta objected to *C. tyzzeri* (Slapeta 2011), because the name was used by Levine (1961) for a *Cryptosporidium* sp. in chickens originally described by Tyzzer (1929), while the *C. tyzzeri* described by Levine (1961) is considered a synonym of *C. meleagridis* Slavin 1955. On the basis of this, Slapeta suggested that the name *C. tyzzeri* by Ren et al. (2012), was both a primary homonym and a junior synonym. However, homonymy and synonymy refer to the application of the same name to different taxa and different names to the same taxon, respectively. Because the taxon *C. tyzzeri* Levine 1961 is not an established one, the homonymy and synonymy suggested by Slapeta do not exist. This is because the nomen nudum nature of *C. tyzzeri* Levine 1961 was previously pointed out in several reviews of *Cryptosporidium* taxonomy (Upton 2003; Fayer 2008), and was stated clearly in the Etymology section of *C. tyzzeri* (Ren et al. 2012).

Cryptosporidium pestis has never been formally described and therefore is not a valid species as it lacks a full taxonomic description, and the name has only been used in the literature in four publications since 2006 (three of which were by Slapeta). To re-name *C. parvum* as *C. pestis* would be confusing to not only *Cryptosporidium* researchers but also the wider veterinary and medical community and water industry, who struggle to keep up with the taxonomy as it is. It would also be in violation of the underlying principles of the ICZN. Therefore, the prevailing name *C. parvum* for the species infective to calves and humans must be retained to avoid confusion.

1.10 Conclusions

To date 25 *Cryptosporidium* species are recognized as valid. Undoubtedly many *Cryptosporidium* genotypes will be formally described as species in the future. As new species of *Cryptosporidium* are named, recommendations from the "Code" in achieving stability and ensuring the uniqueness of each species should be borne in mind.

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Chapter 2 Epidemiology of Human Cryptosporidiosis

Simone M. Cacciò and Lorenza Putignani

Abstract Cryptosporidium species are protozoan parasites that infect the epithelial cells of the gastrointestinal tract of vertebrates. In humans, cryptosporidiosis is usually a self-limiting infection in immunocompetent individuals, but severe diarrhoea and dissemination to extra-intestinal sites can occur in high-risk individuals, such as the very young, the elderly and immunosuppressed individuals, particularly those with HIV infection. The oocyst, the infectious stage of Cryptosporidium, is immediately infectious upon excretion with the host faeces, which favours direct transmission. Oocysts have the capacity to persist in the environment and to withstand standard water treatment and some species of Cryptosporidium, particularly C. parvum, have a wide host range and can be transmitted to humans by direct contact with animals or through ingestion of water and food contaminated with oocysts. Due to the presence of multiple transmission routes, the epidemiology of cryptosporidiosis is complex. The investigation of sporadic cases and outbreaks of cryptosporidiosis has contributed to a better understanding of risk factors and infection sources. Genotyping techniques have enabled a better understanding of the epidemiology of cryptosporidiosis in different geographical, seasonal and socioeconomic context.

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2.1 Introduction

Cryptosporidia are obligate intracellular parasites of many species from all vertebrate classes. First described in laboratory mice by Tyzzer in 1912, *Cryptosporidium* was recognized as a cause of diarrheal disease in animals and then in humans during the 1970s. In the following decade, cryptosporidiosis emerged worldwide as a common cause of severe or life-threatening infection in immunocompromised patients, especially those with AIDS, and of acute, self-limiting gastroenteritis in otherwise healthy subjects, especially children (see Chap. 9).

In humans, infection is caused mainly by the zoonotic species *Cryptosporidium parvum*, which is highly prevalent in young livestock, and *Cryptosporidium hominis*, which is essentially a human parasite. With the development of improved genotyping methods, an increasing number of species and genotypes have been recognized as human pathogens, albeit with low prevalence (Putignani and Menichella 2010). The epidemiology of the infection involves both direct transmission from animals to humans or from person to person, as well as indirect transmission through ingestion of water and food contaminated with infectious oocysts (Cacciò et al. 2005; Smith et al. 2006b).

This chapter focuses on the epidemiology of human infections. The molecular epidemiology of human cryptosporidiosis is presented in Chap. 3.

2.2 Life Cycle of Cryptosporidium

The *Cryptosporidium* life cycle is represented in Fig. 2.1, and the major phases are briefly described here. The life cycle of the parasite is completed within a single host (i.e., a monoxenous cycle), and involves both asexual and sexual replication. For a more detailed account, the readers are referred to previously published articles (Chen et al. 2002; Smith et al. 2005; Fayer 2008).

Excystation: after ingestion of oocysts by a susceptible host, the first step towards infection is excystation, the process by which the oocyst wall opens along a suture to allow the release of four infectious sporozoites. This process has been studied under in vitro conditions and several factors that mimic the transit through the acidic stomach to the alkaline small intestine have been shown to enhance excystation. In particular, temperature and pH appear to be the most important triggers. *Cryptosporidium* species that infect the stomach of the host (like *Cryptosporidium muris*) respond more rapidly to those triggers compared to species that infect the intestine of the host, indicating the need of gastric species to rapidly excyst and release the sporozoites upon ingestion (Widmer et al. 2007). Therefore, the role of host derived triggers during the excystation process varies for different *Cryptosporidium* species (Smith et al. 2005).

Attachment and invasion: like other apicomplexans, *Cryptosporidium* possesses an apical complex formed by the apical ring, the conoid and secretory

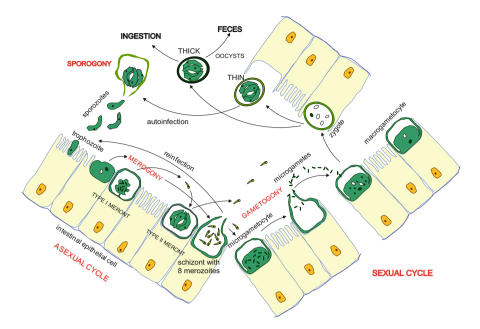


Fig. 2.1 Life cycle of parasite. The infection is acquired through the ingestion of sporulated oocysts. Motile sporozoites, from opened oocyst, attach to intestinal epithelial cells. The trophozoite undergoes an asexual replication (merogony), resulting in the production of eight merozoites (type I meronts). Merozoites, released into the intestinal lumen, infect new intestinal epithelial cells, and originate type II meronts, characterized by four merozoites. The merozoites can undergo a sexual cycle (gametogony) and develop into macrogametocytes. The microgametocyte produces numerous microgametes which are released into the intestinal lumen. A microgamete will fuse with a macrogamete and the resulting zygote undergoes sporogony. Fully sporulated thick and thin oocysts are shed into the intestinal lumen at the completion of sporogony. The infectious thick oocysts are excreted in the feces, thus completing the life cycle. An autoinfection in which excystation takes place within the same host may also be possible and is mediated by 'thin-walled' oocysts

organelles (a single rhoptry and micronemes). The apical complex is intimately involved in the process of attachment and invasion of host cells (Borowski et al. 2008). When the sporozoite contacts the host cell membrane, the rhoptry extends to the attachment site, while micronemes and dense granules move to the apical region. The content of secretory organelles is released, which triggers the process of recognition and attachment to the host cell, invasion, and formation of the parasitophorous vacuole in which the parasite replicates. The sporozoites within the parasitophorous vacuole are not directly in contact with the host cell, and occupy a unique intracellular but extracytoplasmic niche, typical of *Cryptosporidium* parasites (also named parasitophorous sac, Valigurová et al. 2008). A complex process at the host-parasite interface leads to the formation of the so-called feeder organelle, which is thought to be essential in salvaging nutrients and directly separates the parasite cytoplasm from the affected host cell (Umemiya et al. 2005). As the internalization process progresses, the sporozoite differentiates into a spherical trophozoite.

Asexual reproduction: merogony is characterized by the division of the trophozoite nucleus and cytoplasm to generate meronts. In *C. parvum*, there are two types of meronts, type I and type II. Type I meronts have six or eight nuclei. Each gives rise to a merozoite, which is structurally similar to the sporozoite. Mature merozoites leave the meront and invade other host cells, where they can develop into another Type I or into a Type II meront. Type II meronts have four nuclei and generate four merozoites.

Sexual reproduction: upon infection of new host cells, merozoites from Type II meronts can initiate the sexual phase by differentiating into either a microgamont or a macrogamont. Nuclear division in the microgamont leads to the production of numerous microgametes (equivalent to sperm cells) that are released from the parasitophorous vacuole. Each macrogamont (equivalent to an ovum) may be fertilized by a microgamete. The product of fertilization, the zygote, develops into an oocyst.

Sporogony: the zygote differentiates into four sporozoites (sporogony) within the oocyst. It is thought that fully sporulated oocysts are released into the lumen of the intestine and pass out of the body with the faeces, where they are immediately infectious for other susceptible hosts. Oocysts that sporulate in the respiratory tract are found in nasal secretions and sputum (Mor et al. 2010). *Cryptosporidium* oocysts are spherical, measuring only $3-6 \mu m$ in diameter. Each contains four haploid sporozoites and possesses a tick wall.

It is believed that some of the oocysts possess a thin wall and can cause autoinfection in the same parasitized host by liberating their sporozoites in the gut lumen. The process of autoinfection is believed to occur only in species of *Cryptosporidium* and *Caryospora*. Upon their release, sporozoites undergo the developmental processes of schizogony, gametogony and sporogony in enterocytes of the same infected host. Therefore, the life cycle of *Cryptosporidium* ensures the production of very large numbers of infective oocysts, due to the recycling of merozoites to produce further type I generations of schizonts, and the endogenous re-infection from thin-walled oocysts.

2.3 Cryptosporidium Species Infecting Humans

The first human cases of cryptosporidiosis were reported in 1976 (Nime et al. 1976; Meisel et al. 1976). The patients were a 3-year-old child and a 39-year old individual who had severe bullous pemphigoid (a skin disease characterized by blisters) and received treatment with cyclophosphamide and prednisolone. Both patients lived on a farm with cattle and had a dog. They presented with severe watery diarrhea. Diagnosis was based on microscopic examination (including electron microscopy) of rectum and jejunal biopsy specimens. At that time, identification at the level of species was not possible, and infections were attributed to *Cryptosporidium* spp. In 1978, it was shown that oocysts are shed with the feces of infected hosts (Pohlenz et al. 1978) and since then, the diagnosis of cryptosporidiosis has been based on the demonstration of oocysts in feces.

A series of experiments performed using parasite isolates from different hosts (a calf, a lamb, a human and a deer) showed that *Cryptosporidium* could be transmitted to newborn animals (mice, rats, guinea pigs, piglets, calves and lambs), thus demonstrating a lack of host specificity and underlining a zoonotic potential (Tzipori et al. 1980). It was therefore argued that a single species, *C. parvum*, was the causative agent of cryptosporidiosis in mammals, including man.

The role of *Cryptosporidium* as a serious human pathogen was firmly demonstrated in the 1980s in individuals with HIV/AIDS, who experienced a persistent and life threatening infection, often involving parasite dissemination to the hepatobiliary and the respiratory tracts in addition to the entire gastrointestinal (GI) tract (Ma and Soave 1983; Navin and Juranek 1984). The next major event that attracted worldwide interest towards the parasite was the 1993 massive waterborne outbreak in Milwaukee, Wisconsin, that involved an estimated 403,000 person (MacKenzie et al. 1995). This event demonstrated the ability of *Cryptosporidium* to resist water treatment and be transmitted through drinking water. Following these major events, molecular methods for the detection and identification of *Cryptosporidium* has dramatically changed our understanding of the taxonomy and epidemiology of *Cryptosporidium*.

In the early 1990s, the application of Southern blotting (Ortega et al. 1991), Western blotting (Nina et al. 1992) and isoenzymes profiles (Ogunkolade et al. 1993) provided the first evidence of genetic heterogeneity among *C. parvum* isolated from humans and livestock. These studies demonstrated for the first time that humans were infected with two types of *Cryptosporidium* parasites, one being apparently the same as found in cattle and the other exclusively found in humans. However, due to the large amount of biological material needed, these techniques have not been used to characterize field isolates, and have been largely replaced by DNA amplification techniques (PCR). Different assays, including PCR followed by restriction length fragment polymorphism (PCR-RFLP) (Ortega et al. 1991), PCR followed by sequencing (Morgan et al. 1997), Random Amplification of Polymorphic DNA (RAPD) (Morgan et al. 1995), and length polymorphisms of simple DNA repeats (Cacciò et al. 2000), were developed to investigate genetic heterogeneity among isolates.

A number of studies, based on the analysis of single or multiple genetic markers (e.g., Peng et al. 1997; Spano et al. 1998), confirmed the presence of two genetically distinct subgroups within *C. parvum*, which were referred in the literature to as 'human' and 'cattle,' H and C, or Type 1 and Type 2. This result was subsequently confirmed in many laboratories and demonstrated the existence of two distinct transmission cycles of human cryptosporidiosis, one comprising ruminants and humans (potentially zoonotic cycle) and the other exclusively comprising humans (solely anthroponotic cycle).

In 2002, on the basis of accumulating observations of genotypic and biological differences, a new species, *C. hominis*, was finally proposed for *C. parvum* parasites exclusively infecting humans (Morgan-Ryan et al. 2002; see Chap. 1 for the taxonomy of *Cryptosporidium*).

The increasing application of genotyping techniques in epidemiological surveys has demonstrated that the large majority (>90 %) of human cases of cryptosporidiosis are due to *C. hominis* and *C. parvum* (Xiao 2010). However, it is now clear that other species have the potential to cause infection in humans, including *Cryptosporidium meleagridis* and, more occasionally, *C. canis*, *C. felis*, *C. ubiquitum* and *C. viatorum*, and other *Cryptosporidium* genotypes of unknown taxonomic status. The prevalence of these less common species and genotypes varied geographically: for example, *C. meleagridis* was found to be as prevalent as *C. parvum* in children from Peru and in Thailand (Cama et al. 2008; see Chap. 3).

2.4 Incubation Period

Based on evidence from experimental infections, it has been estimated that the incubation period is between 5 and 7 days. Data from experimental infection of healthy volunteers (Okhuysen et al. 1996; Chappel et al. 2006, 2011), and investigations of waterborne and foodborne outbreaks are the main source of information.

Experimental infections of healthy volunteers have been performed using the three main human pathogens, namely *C. parvum*, *C. hominis* and *C. meleagridis*. For *C. parvum*, volunteers were challenged with different inocula (10 to >10,000 oocysts) of three strains of animal origin (Iowa, TAMU and UCP, MD, Okhuysen et al. 1996). Results showed that 12 (86 %) of 14 volunteers who received the TAMU isolate developed diarrhoea, as did 15 (52 %) of 29 who received the Iowa isolate and ten (59 %) of 17 who received the UCP isolate (Okhuysen et al. 1996). The mean incubation period was 7.7 days (for IOWA), 7 days (for UCP) and 4 days (for TAMU).

In the study with *C. hominis*, 21 healthy adults were challenged with 10–500 oocysts of the TU502 (Xu et al. 2004) isolate, and 13 developed diarrhoea, while nine had oocysts detected in faecal samples (Chappel et al. 2006). The mean incubation period was 5.4 days (range, 2–10 days). Finally, in the case of *C. meleagridis*, the study involved five volunteers, who were challenged with 10^5 oocysts of the isolate TU1867 (Akiyoshi et al. 2003). The incubation period was of 5.3 days (range, 4–7 days) (Chappel et al. 2011).

During the large waterborne outbreak of cryptosporidiosis in Milwaukee, the mean incubation period was estimated to range from 3 to 7 days, but it appeared to be shorter in the elderly (5–6 days) compared to either children (7 days) or adults (8 days) (Naumova et al. 2003).

In 1996, another large waterborne outbreak occurred in Japan, and was caused by contamination of the town's potable water (Yamamoto et al. 2000). An estimated number of 9,140 individuals were affected. The median incubation period for the 14 persons for whom this calculation was possible was 6.4 days (range, 5–8 days).

In 2002, an outbreak of cryptosporidiosis occurred among visitors to a public swimming pool in Sweden and affected an estimated with about 800–1,000 individuals (Insulander et al. 2005). The median incubation period was estimated at 5 days (range, 2–13 days).

Three investigations of foodborne cryptosporidiosis have also provided data on incubation period. In 1993, an outbreak linked to consumption of contaminated apple cider occurred in central Maine (Millard et al. 1994). The median incubation period was 6 days (range, 10 h to 13 days).

In 1997, an outbreak of acute gastroenteritis occurred among members of a group attending a dinner banquet catered by a restaurant in Spokane (Anon 1998). Foodborne transmission was implicated through a contaminated ingredient in multiple menu items. The incubation period was estimated between 3 and 9 days. In 2006, acute gastroenteritis caused by *C. parvum* was reported by four members from the same company who had eaten a raw meat dish called "Yukke: Korean-style beef tartar" and raw liver at a rotisserie in Sakai City, Japan (Yoshida et al. 2007). Based on information from interviews, the median incubation period was 5.5 (range, 5–7 days).

Further data comes from other investigations. A study of an outbreak of GI illness among a class of 96 undergraduate veterinary students in New Zealand (Grinberg et al. 2011) indicated a median incubation period of 5 days (range, 0-11 days).

Variability in the estimated duration of the latent period can be attributed to inaccuracies in the estimation of the time from exposure to the onset of symptoms, or to difference in inoculum size (Chappel et al. 2006), but may also reflect alterations in the incubation period due to partial immunity derived from prior exposure to the parasite. Phenotypic differences among parasite isolates may also explain the observed range.

2.5 Asymptomatic Infection

Little information is available on asymptomatic carriage of *Cryptosporidium* in humans. In studies of experimental infection of healthy volunteers with *C. parvum*, it was noted that some individuals passed oocysts in their stools in the lack of overt symptoms (diarhoea), indicating that asymptomatic shedding of the parasite occurs (Okhuysen et al. 1999). In contrast, asymptomatic shedding was not seen in any of the volunteers experimentally challenged with *C. hominis* oocysts (Chappel et al. 2006).

A prospective study in the US found asymptomatic cryptosporidiosis in 12 of 78 (6.4 %) immunocompetent and 11 of 50 (22 %) immunodeficient children (Pettoello-Mantovani et al. 1995). By comparison, *Cryptosporidium* was found in

4.4 % of immunocompetent and 4.8 % of immunodeficient children of a control symptomatic population. A recent study in the UK screened 230 asymptomatic children in preschool day care centres, using a highly sensitive detection protocol based on immunomagnetic separation and DNA typing (Davies et al. 2009). The observed prevalence (1.3 %) was lower than that from the study of Pettoello-Mantovani et al. (1995), that was based on the less sensitive microscopic technique. However, the detection of *C. ubiquitum* and of the skunk genotype in those asymptomatic children raises the possibility that some species/genotypes may have low pathogenicity and be more common than previously thought.

In Scandinavian countries (Denmark, Sweden, Norway and Finland) a metaanalysis of asymptomatic and symptomatic cryptosporidiosis in adults showed a lower prevalence (0.99 %) in the former compared to the latter (2.99 %). In a community-based study in Melbourne, Australia, 1,091 faecal specimens from asymptomatic individuals were screened for the presence of bacteria, viruses and parasites (Hellard et al. 2000), and *Cryptosporidium* was found in four samples (0.4 %). The role of carriers in the transmission of *Cryptosporidium* was also suggested by a large-scale case–control study of sporadic cases in the United Kingdom (Hunter et al. 2004), that identified changing diapers as an independent risk factor for infection with *C. hominis*, even if the child did not have diarrhoea.

The prevalence of asymptomatic infection in other countries is only partially known. A study of 377 Aymara school students (5–19 years of age) from villages of the northern Bolivian Altiplano (Esteban et al. 1998) found a very high prevalence of *Cryptosporidium* (31.6 %) based on microscopy. The authors argued that the mild or asymptomatic infections observed in children were due to some level of immunity that develops after continuous exposure to the parasite.

In Jeddah, Saudi Arabia, a study of asymptomatic children from nurseries found that 9 of 190 (4.7 %) were positive for *Cryptosporidium*, as compared to 20 of 63 (32 %) children with diarrhoea from paediatric clinics of the same city (Al-Braiken et al. 2003). In Makkah, Saudi Arabia, *Cryptosporidium* was detected in 4 % of 589 stool samples from asymptomatic school children aged 7–12 years from 13 primary schools (Al-Harthi 2004). Finally, in a study of 276 asymptomatic aboriginal (Orang Asli) children (141 boys and 135 girls, aged 2–15 years) living in villages in the Malaysia (Al-Mekhlafi et al. 2011), *Cryptosporidium* was detected in 20 (7.2 %) children.

An interesting study aimed at determining the contribution of asymptomatic immigrants in the spreading of the disease in the Kashmir region (India), where cryptosporidiosis is not considered to be endemic. Analysis of stool samples from 45 non-diarrheic and nine diarrheic HIV-infected individuals revealed that all were carriers of *Cryptosporidium* spp. (Masarat et al. 2012). Remarkably, epidemiological traits revealed that the asymptomatic individuals were non-Kashmiri army personals and travellers (immigrants), while local emigrant merchants represented symptomatic cases. This suggests that the non-diarrheic HIV positive population may be a potential source of endemic spread of cryptosporidiosis, and that obligatory laboratory testing in HIV positive immigrant population, like merchants and

travellers, regardless of symptoms, should be mandatory to understand patterns of transmission.

2.6 Symptomatic Infection

The symptomatic period of infection is characterized by diarrhoea, abdominal pain, nausea or vomiting, mild fever, anorexia, malaise, fatigue and weight loss (Fayer and Ungar 1986; Casemore 1990). Diarrhoea can be of sudden onset and is generally watery and voluminous; between three and six stools (but sometimes many more) may be passed each day, which are sometimes offensive and may contain mucus. Symptoms usually last up to 3 weeks, but some patients experience chronic diarrhoea of a month or longer. Oocysts may continue to be shed for a mean period of 7 days (range 1–15 days) after symptoms have ceased, although exceptionally for up to 2 months (Jokipii and Jokipii 1986) (see Chap. 9).

2.7 Risk Factors

Our knowledge of risk factors for acquiring cryptosporidiosis is mainly derived from investigations of outbreaks, albeit the majority of human infection is sporadic. Outbreak investigations have demonstrated the existence of multiple transmission routes, including contact with infected animals, person-to-person transmission in households and care settings, consumption of contaminated foods and drinks, consumption of water from private and public supplies, exposure to recreational water in swimming pools or water parks, and travel to endemic countries (Nichols et al. 2009) (Fig. 2.2). It is important to stress that risk factors are likely to differ for *C. parvum* and *C. hominis* and in different geographical settings, due to the difference in host range, transmission cycles and prevalence. This underscores the importance of identifying species or genotype in epidemiologic studies, both from developed (Hunter et al. 2004; Lake et al. 2007) and developing countries (Cama et al. 2008; Molloy et al. 2011).

In industrialised countries, case–control studies have been conducted in the US (Roy et al. 2004), the UK (Hunter et al. 2004; Lake et al. 2007), and Australia (Robertson et al. 2002). In all these studies, statistically significant risk factors, identified by multivariate analysis, included contact with persons with diarrhoea, particularly young children, and contact with cattle, especially calves. Further, travel abroad was significantly associated with an increased risk in the US and UK studies (Roy et al. 2004; Hunter et al. 2004). History of travel abroad was excluded in the Australian study to allow focus on endemic disease (Robertson et al. 2002). In addition, the US study identified swimming in fresh water as a risk factor and the Australian study found swimming in a chlorinated swimming pool as

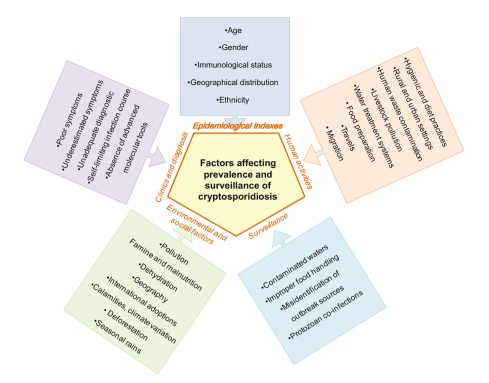


Fig. 2.2 Factors affecting prevalence and adequate surveillance of cryptosporidiosis

a risk factor. Both the UK and Australian studies identified a dose dependent risk associated with drinking unboiled water, which was also reported in a regional study in the UK (Goh et al. 2004). Negative associations with eating ice cream and raw vegetables was found in the UK and US studies.

A more recent study from the UK (Lake et al. 2007) investigated the role of wider environmental and socioeconomic factors (e.g. water supply, socioeconomic status, land use, livestock densities and healthcare accessibility) upon human cryptosporidiosis. By comparing 3,368 laboratory-confirmed cases to an equal number of controls, the authors concluded that risk factors for *C. hominis* and *C. parvum* must be considered separately. Indeed, for *C. hominis* cases the strongest risks factors were living in areas with many higher socioeconomic status individuals, living in areas with a high proportion of young children and living in urban areas. In contrast, agricultural land use surrounding the place of residence and the water supply were significant risk factors for *C. parvum* illness (Lake et al. 2007).

Despite the higher prevalence of cryptosporidiosis in developing and tropical countries, few studies have been conducted to identify risk factors in those areas. In Peru, a total of 156 cases of cryptosporidiosis were found in 109 of 553 children during a 4-year longitudinal birth cohort study (Cama et al. 2008). Investigation

into risk factors did not identify statistically significant associations between *Cryptosporidium* spp. and any of the variables considered (basic aspects of sanitation and zoonotic, foodborne, and waterborne transmission), possibly because children are exposed to these parasites through different transmission routes, which makes single exposure variables difficult to demonstrate (Cama et al. 2008) (Fig. 2.2).

In Indonesia, a study of 917 patients with acute diarrhea (715 in-patients and 202 out-patients from the Hospital of University of Airlanaga in Surabaya) found *C. parvum* oocysts in 26 (2.8 %) of the patients and in 15 (1.4 %) of 1,043 control patients. Investigation of risk factors by multiple logistic regression indicated that contact with pets (cats), rainy season, occurrence of a flood, and crowded living conditions were significant risk factors for cryptosporidiosis (Katsumata et al. 1998).

A 2-year study in the Nile River Delta in Egypt examined risk factors for cryptosporidiosis in children with diarrhea (Abdel-Messih et al. 2005). A prevalence of 17 % (241 of 1,275) was observed, and clinical findings included vomiting, persistent diarrhoea and the need for hospitalization. Children <12 months of age were 2.4 times more likely to be infected with *Cryptosporidium* (p<0.01) and children 12–23 months were 1.9 (p<0.05) times more likely to be infected with the organism as compared to older children. Breastfeeding had a trend towards protection against *Cryptosporidium*-associated diarrhoea (p = 0.07).

Finally, a study in Nigeria found that 134 of 692 children (19.4 %) were infected with *Cryptosporidium* spp. The study also provided information on the species for 49 isolates (Molloy et al. 2011). Using generalized linear mixed-effects models, risk factors were identified for all *Cryptosporidium* infections, as well as for *C. hominis* and *C. parvum* both together and separately. Malaria and absence of *Ascaris* infection were risk factors for all *Cryptosporidium* infections, whereas stunting and younger age were highlighted as risk factors for *C. hominis* infections. Stunting and malaria were identified as risk factors for *C. parvum* infection (Molloy et al. 2011).

Together, studies in developing countries indentified several risk factors: age <2 years, absence of breastfeeding, contact with pets, living in overcrowded conditions, low birth weight, male gender, malnourishment and co-infections as significant risk factors for cryptosporidiosis (Putignani and Menichella 2010) (Fig. 2.2).

The role of host genetics in susceptibility to infection was studied in a cohort of 226 Bangladeshi children aged 2–5 years, who were prospectively followed for >3 years (Kirkpatrick et al. 2008). Ninety-six children (42.5 %) were diagnosed with *Cryptosporidium* infection. A total of 51 (22.6 %) had asymptomatic infection, whereas 58 (25.7 %) had symptomatic cryptosporidiosis, of whom 17 (29.3 %) had recurrent disease. Infected children, both asymptomatic and symptomatic, were more likely to carry the human leukocyte antigen (HLA) class II DQB1*0301 allele (P = 0.009), and a strong association was found between the DQB1*0301/DRB1*1101 haplotype and the development of both asymptomatic and symptomatic infection (P = 0.009). Infected children were also more likely to carry the

B*15 HLA class I allele. This was the first description of a genetic component of the immune response to *Cryptosporidium* infection, which included HLA class I and II alleles (Kirkpatrick et al. 2008).

2.8 Post-Infectious Sequelae

Little is known on the long-term consequences of *Cryptosporidium* infection in humans. The gut epithelium generally recovers after resolution of symptoms, but there are some indications that long-term sequelae may arise (Cacciò et al. 2009).

In immunocompetent individuals, the medium-term health effects of cryptosporidiosis is characterized by the recurrence of loss of appetite, vomiting, abdominal pain, and diarrhoea, independently if the patients have been infected by *C. parvum* or by *C. hominis* (see Chap. 9).

Interestingly, significant differences in the occurrence of extra-intestinal symptoms as sequelae of cryptosporidiosis following infections with *C. hominis* and *C. parvum* have been reported in immunocompetent persons (Hunter et al. 2004). Indeed, eye pain and recurrent headache are associated with *C. hominis* infection but not with *C. parvum* infection, whereas other symptoms, such as fatigue and joint pains, are present after infections with both species, but are significantly more common after *C. hominis* infection (Hunter et al. 2004). The relatively small number of case patients who reported joint pains (13 control subjects versus 36 case patients) means that the establishment of firm conclusions about the nature and distribution of joint symptoms will require further investigations (Hunter et al. 2004).

The potential association of *Cryptosporidium* with the inflammatory bowel syndrome (IBS), a common GI disorder characterized by abdominal pain and alterations in bowel habits, is still uncertain. Indeed, a study of intestinal mucosal biopsies and serology from patients with IBS did not support a major role for the parasite in its pathogenesis (Chen et al. 2001). However, experimental *C. parvum* infection in a rat model resulted in jejunal hypersensitivity to distension, which was also associated with activated mast cell accumulation at 50 days post-infection (Khaldi et al. 2009). These findings are consistent with the observations that IBS patients have a marked increase in mast cell numbers and higher tryptase concentrations in jejunal fluid. Thus, further studies are needed to understand the role that *Cryptosporidium* infection may have in the establishment of IBS.

A seronegative reactive arthritis secondary to cryptosporidiosis has been reported in adults (Hay et al. 1987; Ozgül et al. 1999; Collins and Highton 2004) and children (Shepherd et al. 1989; Cron and Sherry 1995), including one report of Reiter's syndrome (arthritis, conjunctivitis and urethritis) (Cron and Sherry 1995).

2.9 Burden of Disease

2.9.1 Overall Prevalence of Infection

In the developed countries, diarrhoea is the most common reason for missing work, while in the developing world, it is a leading cause of death. Internationally, the mortality rate is 5-10 million deaths each year (Nemes 2009). In this scenario, Cryptosporidium is a major cause of diarrheal disease, globally (Shirley et al. 2012). Unlike many common causes of infectious enteritis, control and treatment of this infection are still problematic. Indeed, control is focused mainly on prevention and no widely effective vaccine or drug-based intervention strategies are available (see Chap. 11). Furthermore, control strategies are particularly deficient for infections of severely immunocompromised individuals, the elderly, children or malnourished people, especially in developing countries. Cryptosporidiosis also presents a significant burden on immunocompetent individuals, and can have permanent effects on physical and mental development of children infected at an early age (Jex et al. 2011). Therefore, a tight monitoring of global infection is nowadays even more essential than in the past, because of the need to include control strategies for different categories of individuals, also in the absence of symptomatic evidence.

Moreover, *Cryptosporidium* is considered an emergent pathogen, often underappreciated for limitations or absence of appropriate diagnostics tools (e.g., microscopy often fails to detect low parasite load) and/or for only indirect association to severe medical growth faltering, malnutrition, and diarrheal mortality.

Recently, significant advances in molecular typing and subtyping analyses have yielded new insights into the epidemiology of cryptosporidiosis; however, in developing countries, point-of-care sites remain crucial for control of enteritis, especially in large areas such as the Indian subcontinent, where they are still lacking in many territories, while highly active antiretroviral therapy (HAART) is not equally distributed. In industrialized countries, outbreaks due to food-borne and water-borne transmission routes, growing transplantation surgery in routine medical practice and related long-term steroid treatments given in combination, are strongly contributing to the increase in disease incidence (Putignani and Menichella 2010; Shirley et al. 2012).

2.9.2 Seasonality

The incidence of cryptosporidiosis exhibits strong seasonality, with low endemic levels followed by pronounced seasonal outbursts (McLauchlin et al. 2000). In a recent meta-analysis on the seasonality of cryptosporidiosis, which was based on 61 published studies, increases in temperature and precipitation were associated with an increase in the incidence of cryptosporidiosis (Jagai et al. 2009).

Precipitation was found to be a strong seasonal driver for cryptosporidiosis in moist tropical climates. On the other hand, in temperate climates, the incidence of cryptosporidiosis peaked with an increase in temperature. The study further shows that while climatic conditions typically define a pathogen habitat area, meteorological factors affect timing and intensity of seasonal outbreaks (Jagai et al. 2009).

Clearly, the seasonal patterns tend to vary with location. For example, in India, the incidence of cryptosporidial diarrhea among children residing in the more temperate northern part of India correlated positively with temperature and negatively with humidity, but this correlation is not observed for children residing in the more tropical southern region. In another study from North-Eastern India, the highest prevalence of cryptosporidiosis was observed during the rainy months, and symptomatic as well as asymptomatic cryptosporidiosis in children was found to increase with increasing rainfall in Kolkata (Desai et al. 2012). In Kuwait, peak incidence occurred during the months of March and April, with no cases during the hottest months of July and August (Daoud et al. 1990).

Due to difference in transmission routes, it is likely that seasonal patterns may vary for different *Cryptosporidium* species. Indeed, genotyping studies conducted in the United Kingdom and New Zealand have found that human cases due to *C. parvum* peak in the late spring whereas those caused by *C. hominis* peak in the fall (McLauchlin et al. 2000; Learmonth et al. 2004). This was interpreted as increased exposure to animal oocysts following the calving and lambing season for *C. parvum*, and to increased travel, exposure to water and attendance to day care centres for *C. hominis*. The dramatic decline in human cases due to *C. parvum* observed in the United Kingdom after the large food-and-mouth outbreak of 2001, was initially explained by a reduced exposure of people to animals and, therefore, reduced zoonotic transmission (Smerdon et al. 2003). However, as the decline has continued, intervention measures, such as the introduction of water regulations and major structural changes in public water supply, are now believed to have played a major role (Sopwith et al. 2005).

2.9.3 Geographic Distribution

Cryptosporidiosis has a worldwide distribution, but the prevalence of infection is assumed to be higher in developing countries (Putignani and Menichella 2010). It should be noted, however, that in many developed and developing countries, surveillance systems for routine detection of cryptosporidiosis are not in place, and few studies have been conducted to estimate how prevalence can vary over time (Nichols 2008).

The distribution of the major *Cryptosporidium* species infecting humans varies geographically. Previous studies have shown that *C. parvum* and *C. hominis* are responsible for >90 % of human cases of cryptosporidiosis in most areas (Xiao and Ryan 2008). In the United Kingdom, in other European countries and in New

Zealand, *C. parvum* is responsible for slightly more infections than *C. hominis* (Xiao 2010). In the Middle East, *C. parvum* is the dominant species in humans. In contrast, *C. hominis* is responsible for more infections than *C. parvum* in the United States, Australia, China and Japan, as well as in most developing countries. Notably, the prevalence of *C. meleagridis* can be as high as that of *C. parvum* in certain areas of the world (Cama et al. 2008). Major differences in transmission routes may account for the observed differences in the distribution of *Cryptosporidium* species (Xiao 2010). The distribution of *C. parvum* and *C. hominis* can also vary within a single country; for example, *C. parvum* is more common than *C. hominis* in rural states in the United States, Ireland and New Zealand (Feltus et al. 2006; Zintl et al. 2009; Snel et al. 2009). Further geographic differences can be observed in the distribution of *C. parvum* and *C. hominis* subtypes (see Chap. 3).

2.9.4 Infection in Children

Cryptosporidiosis occurs more frequently in infants and children than in adults, both in developed and developing countries (Snelling et al. 2007; Xiao 2010; Putignani and Menichella 2010). This is likely to reflect both exposure and immunity. In the United States, endemic parasitic infections are more frequent than commonly perceived. Cryptosporidiosis occurs mainly in children aged 1–9 years, with the onset of infection peaking in the summer in association with communal swimming venues and recreational water use (Barry et al. 2013).

Needless to say, however, the impact of cryptosporidiosis is much higher in the poorest regions of the world. Globally, one in ten child deaths result from diarrhoeal disease during the first 5 years of life, and most occur in sub-Saharan Africa and south Asia (Liu et al. 2012). The role of various pathogens have been very recently studied in 9,439 children with moderate-to-severe diarrhoea and 13,129 control children without diarrhoea (Kotloff et al. 2013). Remarkably, *Cryptosporidium* was identified as a significant pathogen regardless of HIV prevalence and site of collection, and was the second most common pathogen in infants, and was associated with an increased risk of death in toddlers aged 12–23 months (Kotloff et al. 2013).

Children in developing countries are uniquely vulnerable to persistent infection because of the independent and synergistic effects of immune naiveté, malnutrition, and HIV infection (Mor and Tzipori 2008). In these areas, cryptosporidiosis is most prevalent during early childhood, with as many as 45 % of children experiencing the disease before the age of 2 years (Valentiner-Branth et al. 2003). *Cryptosporidium* also plays a causal role in childhood malnutrition and has been linked to impaired physical fitness in late childhood. Studies in Sub-Saharan Africa (reviewed by Mor and Tzipori 2008) have documented significantly higher cryptosporidiosis prevalence among malnourished children. It is difficult to ascertain the direction of this association, i.e., if malnutrition predisposes to infection or if infection actually impairs nutrient absorption and therefore causes weight loss

and growth stunting. Similar findings have been reported from longitudinal studies in Peru (Checkley et al. 1998) and Brazil (Bushen et al. 2007).

The prevalence and predictors of *Cryptosporidium* infection, and its effect on nutritional status, have recently been explored among 276 children (aged 2–15 years) in aboriginal villages in the Malaysian state of Selangor (Al-Mekhlafi et al. 2011). Faecal smears were examined by microscopy while socio-economic data were collected using a standardized questionnaire. Nutritional status was assessed by anthropometric measurements. *Cryptosporidium* infection was detected in 7.2 % of the children, and was found to be significantly associated with low birth weight (≤ 2.5 kg), being part of a large household, and prolonged breast feeding (>2 years).

The impact of *Cryptosporidium* on children has been demonstrated also in Arab countries, such as Egypt, Jordan, Kuwait, Libya, Palestine, Saudi Arabia and Tunisia. Prevalence rates of 1-43 % (mean 8.7 %) in diarrheic immunocompetent children and of 1-82 % (mean 41 %) in immunocompromised children and adults were reported (Ghenghesh et al. 2012). Higher infection rates were found in children living in rural and semi-urban areas than in those residing in urban areas. *Cryptosporidium*-associated diarrhoea occurred mainly in children aged 1 year or less and was inversely correlated with age (Ghenghesh et al. 2012).

Access to quality drinking water in poor regions of the world is thought to be important in limiting gastro-intestinal infections. A study was conducted to determine whether or not bottled drinking water, intended such as "protected" water supply, could prevent or delay cryptosporidiosis among children in an endemic semi-urban community in Southern India (Sarkar et al. 2013). A total of 176 children were enrolled and received either bottled (n = 90) or municipal (n = 86)drinking water. Weekly surveillance visits were conducted until children reached their second birthday. Stools were collected every month and during diarrheal episodes, and tested for the presence of Cryptosporidium spp. by PCR. Cryptosporidiosis, mostly in an asymptomatic form, was observed in 118 of 176 (67 %) children during the follow-up period at a rate of 0.59 episodes/child-year. Diarrhea associated with Cryptosporidium spp. tended to be longer in duration and more severe. Stunting at 6 months and a higher disease burden were associated with a higher risk of cryptosporidiosis, but interestingly drinking bottled water was not associated with a reduced risk of cryptosporidiosis. The lack of association between drinking bottled water and cryptosporidiosis suggests possible spread from asymptomatically infected individuals involving multiple transmission pathways.

2.9.5 Infection in Immunocompromised Individuals

Cryptosporidiosis is a leading cause of severe diarrhoea and extraintestinal infection in immunocompromised individuals (see Chap. 9). Besides HIV-infected patients, individuals at high risk include those with X-linked hyperimmunoglobulin M syndrome (XHIM), CD40 ligand or gamma-interferon deficiency, children with leukemia, and organ transplant recipients (Cacciò et al. 2009).

The importance of cryptosporidiosis in HIV-infected people is well established and the subject has been extensively reviewed from the point of view of the epidemiology and clinical features (Hunter and Nichols 2002; Del Chierico et al. 2011), as well as from the perspective of treatment and control (Abubakar et al. 2007). In short, the risk of faecal carriage, the severity of illness and the development of unusual complications of cryptosporidiosis are related to the CD4 cell count (Pozio et al. 1997). Indeed, patients with CD4 counts of less than 50 are at greatest risk for both the severity of the disease and prolonged carriage. In severely immunocompromised persons, the parasite can also colonize extra-intestinal sites, particularly the gall bladder, biliary tract, pancreas, and lungs (Hunter and Nichols 2002).

The introduction of the highly active antiretroviral therapy (HAART) has had a remarkable impact on many opportunistic viral, bacterial and parasitic infections, resulting in a marked reduction in their occurrence and clinical course, at least in developed countries (Pozio and Gomez Morales 2005). The HAART therapy is based on a combination of nucleoside and non-nucleoside reverse transcriptase inhibitors and HIV protease inhibitors, and results in immune restoration, characterized by an increase in memory and naïve CD4⁺C T cells and the recovery of CD4⁺C lymphocyte reactivity against opportunistic pathogens. Cryptosporidiosis, however, remains a major problem for patients failing HAART, for most individuals living with AIDS in developing countries without access to HAART, and for severely malnourished children.

An extended study on intestinal parasitic infections, including cryptosporidiosis, was carried out in Congo by enrolling hospitalized AIDS patients (Wumba et al. 2010). Stool samples were collected from 175 patients older than 15 years. Parasites were detected by microscopy, immunofluorescence antibody tests and PCR (for diagnosis of microsporidia). At baseline, 19 patients (10.8 %) were under HAART and 156 (89.2 %) were eligible for it. Hospitalization was essentially due to intestinal infection associated with diarrhoea (49.7 % of the patients). A parasite was found in 47 of 175 (26.9 %) patients, and 27 out of 175 (15.4 %) were infected with at least one opportunistic parasite. The overall prevalence rate for *Cryptospo*ridium sp. was 9.7 %, and increased to 12.6 % when only patients with diarrhoea were considered. A number of other protozoan and helminths were observed, but no significant relationship was established between any individual parasite and diarrhoea. These results underline the importance of opportunistic infections in symptomatic AIDS patients regardless of diarrhoea at the time of the hospitalisation, and showed that routine microscopic examination for Cryptosporidium spp. should be considered due to the absence of clinical markers.

In HIV patients, hepatic parenchymal and biliary tract diseases are common. A recent study focused on clinical aspects of AIDS-related cholangiopathy (De Angelis et al. 2009). Although the etiology is unclear, several opportunistic infections (including *Cryptosporidium* and cytomegalovirus) are suspected to cause it. The most common finding after endoscopic retrograde cholangiopancreatography is

diffuse sclerosing cholangitis in combination with papillary stenosis. Clinically, the most common manifestations are right upper quadrant pain and fever accompanied by an elevated serum alkaline phosphatase level. In vitro experiments have shown that concurrent active HIV replication and *C. parvum* infection synergistically increase cholangiocyte apoptosis and thus jointly contribute to AIDS-related cholangiopathies (De Angelis et al. 2009).

Primary immunodeficiencies are rare inherited disorders of the innate, cellular, and/or humoral immune system. A particular susceptibility to infection with *Cryptosporidium* is observed in children with X-linked hyper-immunoglobulin M syndrome (XHIM), resulting from CD40 ligand deficiency (CD40L), hyper-IgM syndrome type 3 caused by CD40 deficiency, primary CD4 lymphopenia, severe combined immunodeficiency syndrome, and gamma interferon deficiency (Cacciò et al. 2009). These patients are unable to clear the infection and extraintestinal infection, particularly of the bile tract, may result in chronic liver inflammation or even lead to liver cirrhosis. Colonization of the biliary system may also predispose to the development of sclerosing cholangitis (SC) and cholangiocarcinoma.

In a study of the association between XHIM and tumors of the pancreas, liver, and biliary tree, 14 of 20 boys (70 %) were found to be infected with *Cryptosporid-ium* (Hayward et al. 1997). In these patients, cholangiopathy and/or cirrhosis preceded the development of the tumors, suggesting that infection or inflammation of bile ducts caused by *Cryptosporidium* may play an important role in the development of five patients with XHIM and in a single patient with primary CD4 lymphopenia, and reported SC in these patients (Wolska-Kusnierz et al. 2007). Further support for the association between *Cryptosporidium* infection and SC was found in a study in the United Kingdom that enrolled 35 children with clinical evidence of liver disease and found 12 of 27 children (44 %) infected with the parasite, among whom nine had SC (Rodrigues et al. 2004).

In a recent analysis of 126 patients with XHIM syndrome reported to the European Society for Immunodeficiency registry, approximately one-sixth developed liver disease, and in more than 50 % of cases this was associated with *Cryptosporidium* species infection (Toniati et al. 2002). These figures may even be an underestimate, because more-sensitive molecular techniques reveal that a number of patients are colonized by *Cryptosporidium* without evidence of its presence on conventional microbiology screening (Rodrigues et al. 2004; Wolska-Kusnierz et al. 2007). Thus, unrecognized cases of cryptosporidiosis in children with primary immunodeficiencies may lead to serious consequences, with development of sclerosing cholangitis, liver cirrhosis, and cholangiocarcinoma.

Data on cryptosporidiosis in solid-organ transplant recipients are limited, but those available illustrate the need of a high index of suspicion in any transplant patient who presents with severe diarrhea (Cacciò et al. 2009; Krause et al. 2012). Intestinal infections with *Cryptosporidium* species have been reported in renal transplant patients. Some cases resulted in either mild disease or asymptomatic carriage, but severe cryptosporidiosis can occur in these patients, including biliary involvement treated with reduction of immunosuppression and a short course of

antiparasitic agents (Abdo et al. 2003; Hong et al. 2007). In the recent study of Krause et al. (2012), *Cryptosporidium* was detected as the cause of gastroenteritis in six children (four kidney recipients, one liver and kidney recipient, and one heart transplant recipient). All patients were hospitalized due to prolonged diarrhoea, fever, abdominal pain and weight loss, and most presented deterioration of kidney functions and abnormal values of liver enzymes.

About ten cases of cryptosporidosis has also been reported in liver transplant recipients. In a study from Belgium on 461 children following liver transplantation, three (0.65 %) developed diffuse cholangitis associated with intestinal Cryptosporidium species carriage (Campos et al. 2000). All three recipients required reoperation on the bile duct anastomosis, but biliary cirrhosis developed in one patient, requiring retransplantation. In a retrospective study from Pittsburgh, four (0.34 %) pediatric cases of cryptosporidiosis were identified among 1,160 nonrenal, abdominal organ transplant recipients (Gerber et al. 2000). Three of these four cases occurred in patients receiving liver transplants, and one occurred following a small bowel transplantation. All four patients spontaneously resolved their infections. Manz and Steuerwald (2007) reported a case of cryptosporidiosis in an adult patient treated with interferon and ribavirin for recurrent hepatitis C after liver transplantation. The patient did not have HIV infection or immunoglobulin deficiency and recovered after the treatment was stopped and the dosage of immunosuppressant was lowered. A case of multiple infections with distinct Cryptosporidium species has been described for a transplanted ileum (Pozio et al. 2004). In Iran, among 44 liver transplant children in Shiraz Nemazee hospital, C. parvum and C. meleagridis were detected in 11.36 % of the children (Agholi et al. 2013).

The study by Sulżyc-Bielicka et al. (2012) evaluated the prevalence of *Crypto-sporidium* spp. in 87 patients with diagnosed colorectal cancer, by using a ProSpecT *Cryptosporidium* Microplate Assay. *Cryptosporidium* sp. was found in 12.6 % of the patients, with a prevalence comparable to patients with immune deficiency; however, no specific correlation was found between *Cryptosporidium* spp. infection and gender, age, neoplasm differentiation grade, or neoplastic tumour localisation.

2.9.6 Infection in Travellers

In 2011, approximately 980 million people travelled internationally. More than 522 million people from developed countries travelled overseas; an estimated 50–100 million people travelled to developing countries (Ross et al. 2013). Approximately 8 % of travellers to the developing world require medical care during or after travel and more than a quarter of those who seek medical assistance present with GI symptoms. Traveller diarrhoea (TD) occurs in 20–60 % of European or North American travellers in inter-tropical areas (Cavallo and Garrabé 2007). Bacteria, viruses and parasites all may contribute to TD, but their relative importance is still uncertain; however, protozoan infections with *Giardia* and *Cryptosporidium* are frequently identified as the cause of GI complaints in returning travellers

(Freedman et al. 2006; Thielman and Guerrant 1998; Okhuysen 2001). Travelling represents an important risk factor for acquiring infection also with spore-forming protozoa such as *Cyclospora*, Microsporidia, and *Isospora* (Goodgame 2003). Several studies have shown that a large proportion of travellers and immigrants from tropical and subtropical countries are affected by GI disorders and harbour intestinal pathogens in the absence of evident GI problems (Saiman et al. 2001; Freedman et al. 2006; Ansart et al. 2005; Caruana et al. 2006; Whitty et al. 2000; Fotedar et al. 2007).

One of the first studies of travellers returning from developing countries was performed in Germany in 1997 (Jelinek et al. 1997). To estimate the prevalence of *C. parvum* and *Cyclospora cayetanensis*, 978 stool samples were taken from 795 patients (469 suffering from diarrhoea) returning from developing countries. Of the 795 patients, infection with *C. cayetanensis* was detected in five subjects (1.1 %), while 13 patients (2.8 %) were infected with *C. parvum*. All patients with either *C. parvum* or *C. cayetanensis* infection suffered from watery diarrhoea, suggesting that in cases of persistent watery diarrhoea these pathogens should be always considered in the differential diagnosis (Jelinek et al. 1997).

In the Netherlands, a new diagnostic strategy was recently implemented for the routine diagnosis of intestinal parasites in returning travellers and in immigrants (ten Hove et al. 2009). Over a period of 13 months, unpreserved stool samples, patient characteristics and clinical data were collected from those attending a travel clinic. Stool samples were analysed on a daily basis by microscopic examination and antigen detection, and compared with a weekly performed multiplex real-time PCR analysis for *Entamoeba histolytica*, *Giardia*, *Cryptosporidium* and *Strongyloides stercoralis*. Microscopy and antigen detection screening of 2,591 stool samples showed *E. histolytica*, *Giardia*, *Cryptosporidium* and *S. stercoralis* in 0.3 %, 4.7 %, 0.5 %, and 0.1 % of the cases. These detection rates were lower than those obtained with real-time PCR. PCR positivity was 0.5 %, 6.0 %, 1.3 %, and 0.8 % (ten Hove et al. 2009), showing that high-throughput molecular screening could provide more accurate estimates of the prevalence of these pathogens.

It has been hypothesized that travellers may be exposed to parasite species/ genotypes that do not circulate in their home countries, and that symptomatic infection with these parasites occurs because of the lack of or insufficient crossprotection resulting from previous exposures. In agreement with this hypothesis, Tanriverdi et al. (2008) used microsatellite typing to show that the *C. hominis* isolates infecting patients that had travelled to the UK from Pakistan less than 2 weeks prior to the isolation of the parasites were significantly different from the large cluster of autochthonous cases from the UK. Furthermore, a novel *Cryptosporidium* species was recently identified among travellers with gastro-intestinal symptoms returning to the UK from the Indian subcontinent (Elwin et al. 2012). Based on morphological and molecular data, this parasite was designated a new species, *Cryptosporidium viatorum*. The name was chosen to underscore its link to foreign travel (Elwin et al. 2012). The epidemiology of *C. viatorum* cases was found to be different from that of individuals infected with *C. parvum* and *C. hominis*: most *C. viatorum* cases occurred in the first months of the year, vomiting was reported less often, but the duration of symptoms was longer. The cases of *C. viatorum* were all travellers to the Indian subcontinent, whereas cases of *C. hominis* and *C. parvum* were more likely associated to travel elsewhere (Elwin et al. 2012).

Another study on *Cryptosporidium* infections contracted whilst travelling abroad used a PCR-coupled single-strand conformation polymorphism analysis, followed by targeted sequencing of the gp60 gene (Jex and Gasser 2008). The study investigated *C. hominis* and *C. parvum* isolates (n = 115) from UK citizens inferred to have been infected while travelling abroad (to 25 countries) or in the UK. Isolates were classified to the genotype and sub-genotype levels, leading to the identification of five *C. hominis* and four *C. parvum* gp60 genotypes. A particular *C. hominis* subgenotype (IbA10G2R2) was found in the majority (71 %) of the isolates, collected from individuals who have travelled to 14 different countries, while other subgenotypes appeared to be quite rare (Jex and Gasser 2008).

2.10 Transmission Routes

There are multiple transmission routes through which humans can acquire *Cryptosporidium* infections. The main routes of transmissions are shared by the two major human pathogens, *C. hominis* (Fig. 2.3) and *C. parvum* (Fig. 2.4); however animal pollution and environmental contamination particularly affect zoonotic transmission of *C. parvum* (Fig. 2.4).

2.10.1 Person-to-Person Transmission

Cryptosporidium is easily transmitted among children and staff members in nurseries (Hannah and Riordan 1988), day care centres (Heijbel et al. 1987), and schools (Lee and Greig 2010).

Nosocomial infection is also well documented, and both direct (Baxby et al. 1983; Koch et al. 1985) and indirect (Martino et al. 1988; Navarrete et al. 1991) person-to-person transmission (via contaminated hands) has been incriminated as the likely route. Nosocomial infection can cause secondary cases among roommates (Bruce et al. 2000) and family members (Pandak et al. 2006), further supporting the highly infectious capacity of the parasite.

Direct transmission among HIV-positive men who have sex with men occurs more frequently than in HIV-positive drug users (Pedersen et al. 1996), and a case–control study in Australia (Hellard et al. 2003) has shown that men having more than one sexual partner are more likely to have *Cryptosporidium* diarrhoea, therefore indicating that sexual contacts represent a risk factor for faecal-oral parasite transmission.

In developing countries, the high prevalence of C. hominis and of anthroponotic subtypes of C. parvum, particularly in children, has also been

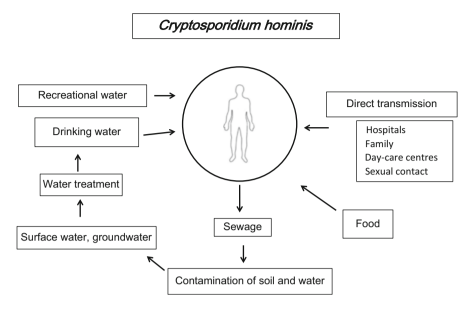


Fig. 2.3 Main transmission routes of Cryptosporidium hominis

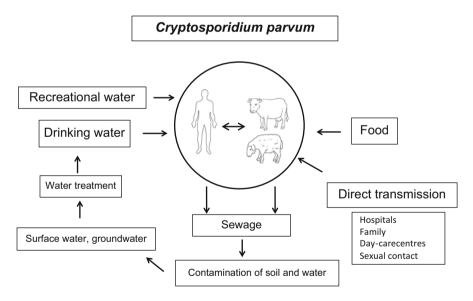


Fig. 2.4 Main transmission routes of Cryptosporidium parvum

taken as an indication of the importance of person-to-person transmission (Xiao 2009) (Figs. 2.3 and 2.4).

2.10.2 Zoonotic Transmission

Cryptosporidium parvum is the most important zoonotic agent of cryptosporidiosis, with a large range and abundance of animal reservoirs, mainly in young farmed animals (see Chap. 4). Therefore, individuals who come in contact with those animals, either for occupational or recreational reasons, may be at risk. Outbreaks of cryptosporidiosis have been reported among veterinarians and veterinary students (Pohjola et al. 1986; Preiser et al. 2003; Gait et al. 2008), other people exposed to agricultural animals (Stantic-Pavlinic et al. 2003) and children visiting farms (Shield et al. 1990; Stefanogiannis et al. 2001; Hoek et al. 2008). Contact with farmed animals was identified as a significant risk factor for sporadic cases of human cryptosporidiosis in the UK (Hunter et al. 2004; Goh et al. 2004).

Due to the high prevalence of *C. parvum* infection and the high numbers of oocysts shed in faeces (up to $>5 \times 10^6$ oocysts per gram), calves are considered to pose the most significant threat to environmental contamination and transmission to humans (Current et al. 1983). Initially, it was believed that human infection with *C. parvum* were all of zoonotic origin and calves have been implicated as the main source of infectious oocysts, but further studies based on highly polymorphic markers have shown that certain *C. parvum* subtypes are found in humans but not in animals, and are likely to be transmitted through an anthroponotic cycle (Mallon et al. 2003). Thus, a significant fraction of human *C. parvum* infections may not originate from livestock reservoirs (Grinberg et al. 2008). Nevertheless, calves are frequently infected with a *C. parvum* subtype that is commonly found in humans in the same geographic areas (Xiao 2009) and epidemiologic studies have supported the occurrence of zoonotic transmission (Hunter et al. 2007).

Epidemiologic investigations have demonstrated the role of sheep in human cryptosporidiosis more than 20 years ago (Casemore 1989), and this has been further supported by molecular studies, particularly in the UK, where five human outbreaks have been linked to contacts with lambs (Chalmers and Giles 2010). Recently, a case of zoonotic transmission of a rare *C. parvum* subtype from infected lambs to a children has been reported in Italy (Cacciò et al. 2013).

On the contrary, little is known about the role of goats in zoonotic cryptosporidiosis. Goats can be infected with *C. xiaoi*, which is a non-zoonotic species, but also with *C. parvum* (Rieux et al. 2013).

The zoonotic potential of canine and feline cryptosporidiosis has been a major concern to both veterinarians and physicians, but the actual risk of transmission from pets to human appears to be minimal, at least in develop countries. To date, there have been only 26 *C. canis* and 97 *C. felis* cases reported in people, and the majority was from immunocompromised individuals (Lucio-Forster et al. 2010). Epidemiological investigations in the UK and USA (Goh et al. 2004; Glaser et al. 1998) have failed to establish a significant relationship between owning a dog and infection with *Cryptosporidium*, whereas owning a dog or a cat was found to be a risk factor for cryptosporidiosis in Guinea Bissau and Indonesia (Miron et al. 1991; Katsumata et al. 1998). There is only one report of possible transmission

of *C. canis* between a dog and two siblings living in a household in Peru (Xiao 2009). It is likely that only immunocompromised individuals are at risk of acquiring cryptosporidiosis from pets.

Rabbits are the natural host of *Cryptosporidium cuniculus* (Robinson et al. 2010), a species that is now known to infect humans (Chalmers et al. 2009). First identified as a human pathogen during a waterborne outbreak, *C. cuniculus* appears to be the third most commonly identified species in patients with diarrhea in the UK, after *C. parvum* and *C. hominis* (Chalmers et al. 2011). The existence of identical *C. cuniculus* subtypes in humans and pets or wild rabbits suggest zoonotic potential (Zhang et al. 2012).

Zoonotic transmission of *C. meleagridis*, an avian parasite that can infect humans, has been recently reported in Sweden (Silverlås et al. 2012). Interestingly, results of molecular characterization suggest laying hens or broiler chickens as the source of infectious oocysts.

Fish, reptiles and amphibians appear not to pose a risk for human cryptosporidiosis (see Chap. 5).

2.10.3 Waterborne Cryptosporidiosis

Waterborne infectious diseases are a globally emerging public health issue. Various community outbreaks due to contamination of water have highlighted the importance of intestinal protozoa in public health. Among these important pathogens are *Giardia duodenalis*, *E. histolytica*, *C. cayetanensis*, *Isospora belli*, Microsporidia and, of greater relevance, *Cryptosporidium* (Karanis et al. 2007).

The ubiquitous presence of *Cryptosporidium* spp. in the aquatic environment is explained by the large number of hosts, the extremely high number of oocysts shed by these hosts, and the remarkable stability of oocysts (Smith et al. 2006b). Thus, water represents a very important vehicle of infection for the population, and waterborne cryptosporidiosis is a serious public health concern, particularly for populations at risk of severe infection (pregnant women, children, HIV-positive and transplanted patients) (Chap. 12).

Indeed, out of the 71 *Cryptosporidium*-linked outbreaks described in the last decade, 40 (56.3 %) appear to be correlated to waterborne transmission. Geographically, the outbreaks seem to be concentrated in the USA, Canada, Australia and Europe, especially in the UK and Ireland, and affect both adults and children (Putignani and Menichella 2010; Chalmers 2012). Surveillance data has revealed the presence of *Cryptosporidium* spp. in the entire water treatment system (see Chap. 12), which represents an unacceptable health risk, particularly for at risk populations (pregnant women, children, HIV-positive and transplanted patients). Such evidence suggests that focus ought to be placed on prevention of human and animal waste contamination especially in authorized recreational waters. Remarkably, cryptosporidiosis is the most frequently reported gastrointestinal illness in

outbreaks associated with treated (disinfected) recreational water venues in USA (Yoder and Beach 2007; Chap. 12).

Following several large outbreaks linked to drinking water in UK and USA, most notably the 1993 Milwaukee outbreak that involved an estimated 403,000 cases of cryptosporidiosis (MacKenzie et al. 1995), emphasis on monitoring and intervention of water supplies, and greater awareness and investigation of water as a transmission vehicle, has led to a decrease in the number of outbreaks due to drinking water (Chap. 12).

Remarkably, however, cryptosporidiosis remains the most frequently reported GI illness in outbreaks associated with treated (disinfected) recreational water venues in the UK and the USA (Hlavsa et al. 2011; Smith et al. 2006a; Yoder and Beach 2007; Yoder et al. 2010). Outbreaks have occurred in swimming pools, paddling or wading pools, water parks ad fountains, and were caused by C. hominis and C. parvum (see Chap. 12). During the summer of 2007, Utah experienced a statewide outbreak of GI illness caused by Cryptosporidium (CDC, MMWR Report 2012). Of 1,506 interviewed patients with laboratory-confirmed cryptosporidiosis, 1,209 (80 %) reported swimming in at least one of approximately 450 recreational water venues during their potential 14-day incubation period. Because swimmers were the primary source of Cryptosporidium contamination, healthy swimming campaigns are needed to increase awareness and practice of healthy swimming behaviours, especially not swimming while ill with diarrhoea. The healthy swimming campaign, as part of a multipronged prevention effort, might have helped prevent recreational water-associated outbreaks of cryptosporidiosis in Utah (CDC, MMWR Report 2012). Local and state health departments can use cryptosporidiosis surveillance data to better understand the epidemiologic characteristics and the disease burden of cryptosporidiosis in the USA, to design prevention strategies that reduce disease spread and to establish research priorities. The role of water in the transmission of *Cryptosporidium* in developing countries is less known. Clearly, the potential for transmission is enhanced by the absence of sanitary and parasitological drinking water monitoring, and the burden of the infection is surely underestimated due to the scarcity of appropriate surveillance programs and the relative inadequacy of laboratory diagnosis (Mak 2004). A Quantitative Microbiological Risk Assessment (QMRA) conducted in Africa (Hunter et al. 2009) demonstrated that interruptions in water supplies that forced people to revert to drinking raw water caused a greater risk of infection, particularly in young children. Thus, poor reliability of drinking water supplies has an impact on the achievement of health improvement targets (Hunter et al. 2009).

The importance of stability of the treatment process and the importance of watershed protection has been stressed by a comprehensive QMRA performed on a source water survey from 66 waterworks in 33 major cities across China (Xiao et al. 2012). The annual diarrhoea morbidity caused by *Cryptosporidium* in drinking water was estimated to be 2,701 cases per 10⁵ immunocompromised persons and 148 cases per 10⁵ immunocompetent persons, giving an overall rate of 149 cases per 100,000 population. The burden of cryptosporidiosis associated with drinking water treated with the conventional process was higher than the reference risk level

suggested by the WHO, but lower than that suggested by the United States Environmental Protection Agency (Xiao et al. 2012).

A survey of *Giardia* and *Cryptosporidium* conducted in 206 samples of surface waters used as drinking water sources by public water systems in four densely urbanized regions of Sao Paulo State, Brazil and a QMRA, showed 102 samples positive for *Giardia* and 19 for *Cryptosporidium*, with maximum concentrations of 97.0 cysts/L and 6.0 oocysts/L, respectively. The probability of *Giardia* infection was close to the rates of acute diarrheic disease for adults (1-3 %) but lower for children (2-7 %). The daily consumption of drinking water was an important contributing factor for these differences (Sato et al. 2013).

2.10.4 Foodborne Cryptosporidiosis

Contamination of different types of food with *Cryptosporidium* oocysts has been demonstrated in studies from different regions of the world. Those studies have mainly focused on fruits and vegetables, because these foods are prone to contamination and are often consumed raw or after minimal thermal treatment, therefore increasing the possibility of transmission (Robertson and Chalmers 2013). Due to the highly variable, and usually low, rates of recovery of oocysts from food matrices, improved methods have been recently developed and validated in the UK (Cook et al. 2006).

Studies in Costa Rica and Peru (Monge and Arias 1996; Ortega et al. 1997) have shown contamination of numerous raw vegetables, including basil, cabbage, celery, cilantro, green onions, leeks, lettuce, parsley, and yerba buena. A more recent study in Costa Rica (Calvo et al. 2004) investigated the presence of *Cryptosporidium* spp., *Cyclospora* spp., and Microsporidia on lettuce, parsley, cilantro, strawberries and blackberries collected from five local markets. Fifty different samples of each product, 25 taken in the dry season and 25 in the rainy season, were evaluated. All products were found contaminated with *Cryptosporidium* spp., *Cyclospora* spp., and/or Microsporidia. *Cryptosporidium* was not detected in strawberries, microsporidia were absent on blackberries and *Cyclospora* was only isolated from lettuce during the dry season. These results show the importance of introducing good agricultural practices, especially due to the resistance of *Cryptosporidium* and *Cyclospora* to disinfecting agents (Putignani and Menichella 2010).

Food contamination with *Cryptosporidium* oocysts, however, is not limited to developing countries. In Norway, a search for parasites in fruits and vegetables was undertaken in the period from 1999 to 2001 (Robertson and Gjerde 2001). Of the 475 samples, 29 were found to be positive for *Cryptosporidium* oocysts and *Giardia* cysts, of which 19 only for *Cryptosporidium* (lettuce and mung bean sprouts). Mung bean sprouts were significantly more likely to be contaminated with *Cryptosporidium* oocysts or *Giardia* cysts than the other fruits and vegetables, even if concentrations were generally low (approximately 3 (oo)cysts per 100 g product). There was no association between imported produce and detection of parasites.

Crvptosporidium oocysts and *Giardia* cysts were also detected in water samples concerned with field irrigation and production of bean sprouts (Robertson and Gjerde 2001).

In Poland, 163 samples comprising Peking cabbages, leeks, white cabbages, red cabbages, lettuces, spring onions, celery, cauliflowers, broccoli, spinach, Brussels sprouts, raspberries, strawberries, all from local markets, were tested (Rzezutka et al. 2010). *Cryptosporidium* oocysts were detected in one leek sample, one celery sample, four cabbage samples, including all cabbage types tested. Of note, *Cryptosporidium*-positive samples came from districts with the highest number of cattle herds. In Spain, 19 fresh produce samples from local markets were tested, and oocysts were found in 33 % of Chinese cabbage, 75 % of Lollo rosso lettuce, and 78 % of Romaine lettuce (Amorós et al. 2010).

These studies demonstrated that contamination of fruits and vegetables occurs also in developed countries.

A few outbreaks have been linked to the consumption of contaminated vegetables. In 2008, a *C. parvum* outbreak in Sweden was linked to chanterelle sauce with fresh parsley added after the preparation of the sauce (Insulander et al. 2008), while in Finland a salad mixture was the suspected vehicle for a *C. parvum* outbreak (Ponka et al. 2009).

Contamination of dairy products and fruit juices has also been linked to outbreaks of cryptosporidiosis. The consumption of unpasteurized cow milk has been suggested as the cause of outbreaks in the UK and Australia (Gelletlie et al. 1997; Harper et al. 2002). Outbreaks associated with consumption of fruit juice is becoming an emergent public health problem since the early 1990s, when the first outbreak associated with apple cider was described (Millard et al. 1994). However, in the period from September to November 2003, 12 local residents in Northern Ohio were diagnosed with cryptosporidiosis for having drunk ozonated apple cider (Blackburn et al. 2006). In response to epidemiologic investigations of outbreaks in which juice is implicated, the USA Food and Drug Administration has implemented process control measures to regulate the production of fruit juice, according with the Hazard Analysis Critical Control Point (HACCP) plan. However juice operations that are exempt from processing requirements or do not comply with the regulation, continue to be implicated in outbreaks of illness. The CDC receives reports of food-associated outbreaks of illness (FoodNet, http://www.cdc. gov/FoodNet/) and its Foodborne Outbreak Reporting System has reviewed, from 1995 through 2005, ten implicating apple juice or cider, eight linked to orange juice, and three involving other types of fruit juice-associated outbreaks (Putignani and Menichella 2010). Among the 13 outbreaks of known aetiology, two were caused by Cryptosporidium and one by Shiga toxin-producing E. coli O111 and Cryptosporidium (Vojdani et al. 2008). The incidence of foodborne disease outbreaks caused by contaminated low pH fruit juices is increasing (Lynch et al. 2006). The association of *Cryptosporidium* with fruit juice is a raising safety concern in food industries. In 1998, CDC implemented enhanced surveillance for foodborne-diseases outbreaks by increasing communication with state, local, and territorial health departments and revising the outbreak report form. Since 2001,

reports are submitted through a web internet application called electronic Foodborne Outbreak Reporting System (Putignani and Menichella 2010).

Outbreak investigations have also put into focus the role of food handlers as a source of food contamination and subsequent transmission of cryptosporidiosis (Robertson and Chalmers 2013). In 1998, a large outbreak of gastroenteritis occurred in >100 persons at a University campus in Washington DC. A case–control study of 88 case patients and 67 control subjects showed that eating in one of two cafeterias was associated with diarrheal illness. Further epidemiologic and molecular evidence indicate that an ill foodhandler was the likely outbreak source, and *C. hominis* as identified as the etiological cause of gastroenteritis (Quiroz et al. 2000).

In 2005 an outbreak of diarrhoea, affecting a group of 99 company employees, was described near Copenhagen (Ethelberg et al. 2009). All people were ill and 13 tested positive for *C. hominis*. Disease was associated with eating from the canteen salad bar on one, possibly two, specific weekdays. Three separate salad bar ingredients were found to be likely sources: peeled whole carrots served in a bowl of water, grated carrots, and red peppers. The likely source of infection was an infected food-handler, who may have contaminated food served at the buffet (Ethelberg et al. 2009).

Recently, the role of the food handlers has been investigated in Venezuela, where cryptosporidiosis is an important public health problem (Freites-Martinez et al. 2009). Despite a basic investigation approach, 14 out of 119 fecal samples from food workers were found positive for *Cryptosporidium* spp. in association with other protozoa, the most frequently detected being *Endolimax nana*, *Blastocystis hominis*, *Entamoeba coli*, *Giardia*, and *E. histolytica/Entamoeba dispar*.

Finally, because of their capacity to filter large volumes of water potentially contaminated with oocysts from human and animal faecal waste, and because they are usually consumed raw, molluscs have been postulated as a route of transmission (Robertson 2007). Indeed many species, including mussels, oysters and clams, have been found to harbour *Cryptosporidium* oocysts in their digestive tract. However, there are no evidence of infections or outbreaks linked to consumption of molluscs.

2.11 Conclusions

Cryptosporidium is an important cause of diarrhoea, worldwide. In developed countries, large waterborne outbreaks continue to occur, emphasizing the need for better regulation and for improvements of drinking water treatment processes. Also, the increasing number of outbreaks linked to the use of recreational water (swimming pools, water parks) indicate the need for better control measures and guidelines. Immunocompromised individuals are particularly susceptible to *Cryptosporidium*, and may develop severe infections and extra-intestinal dissemination, yet an effective therapy to eradicate the parasite is not available. In developing

countries, the parasite is endemic and significantly associated with moderate-tosevere diarrhoea in infants, a finding that highlights the need to develop resources to diagnose, treat, and prevent cryptosporidiosis in resource-poor settings. Under this situation, routine diagnosis and effective reporting of *Cryptosporidium* to local and national surveillance organizations remain of key importance in understanding the epidemiology of this important, but often underestimated, pathogen.

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Chapter 3 Molecular Epidemiology of Human Cryptosporidiosis

Gordon L. Nichols, Rachel M. Chalmers, and Stephen J. Hadfield

Abstract Most human cryptosporidiosis is caused by the species Cryptosporidium hominis and C. parvum, with less common species globally including C. meleagridis, C. cuniculus, C. ubiquitum, C. viatorum, C. canis and C. felis, although the distribution and prevalence may vary. Minor species that are only very rarely found in symptomatic people include C. andersoni, C. suis, C. scrofarum, C. bovis, and C. muris. The use of molecular methods to characterise isolates has contributed to our understanding of the overall epidemiology of cryptosporidiosis and its transmission, the biology and host range of each species and the evolution from a common ancestor. The development of more discriminatory typing methods has shown that outbreaks may be caused by subtypes identified at particular loci. The 60 kDa glycoprotein (gp60) gene has been the most widely used locus due to its high sequence variation. A gp60 naming system has been proposed and widely adopted which has greatly assisted with global epidemiological analysis. However, limitations of a single locus scheme have been identified including low gp60 subtype diversity in certain countries. A plethora of multilocus genotyping (MLG) methods have been developed to provide increased discrimination and are beginning to deepen our understanding of parasite population structures and transmission dynamics. However, there is a need for a standard MLG scheme that, when applied to routine surveillance datasets, will distinguish between outbreak and sporadic cases. This will allow investigation of geographically and temporally dispersed outbreaks which is not currently possible in areas of low gp60 sequence variation.

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3.1 Introduction to Molecular Epidemiology

Cryptosporidiosis is a disease that can be controlled by interventions; these are currently directed at limiting exposure to the parasite. Although vaccination has been attempted (Roche et al. 2013) there are no commercial vaccines and specific treatment options are limited (McDonald 2011). In order to intervene and prevent disease, epidemiological investigations should focus on identifying those at most risk, determining sources, transmission pathways, hosts and vectors and risk factors that are linked with infection. The evidence to date suggests clear human and animal sources and transmission routes that commonly involve water as a vehicle. However, there remain unanswered epidemiological questions and the role of molecular typing, both to the species level and further, in addressing these questions needs to be well documented. A review of the molecular epidemiology of Cryptosporidium infections undertaken in 2007 (Nichols 2008) informed us of a number of things. First, that not all cryptosporidiosis is caused by the same organism (Nichols et al. 1991), a concept proposed in 1980 (Tzipori et al. 1980). Second, it showed that the host range for Cryptosporidium species and genotypes varies (Xiao and Ryan 2004); third, that there is clear evidence of distinct human transmission routes and sources (Peng et al. 1997), with more than one species responsible for human disease; and fourth, that C. parvum and C. hominis are generally the most common species causing human cryptosporidiosis (McLauchlin et al. 2000; Morgan-Ryan et al. 2002; Peng et al. 1997). There was evidence that the distribution of these and other species differs geographically, with C. meleagridis being more common in some settings (for example in a shanty town in Peru) (Cama et al. 2003; Xiao et al. 2001b). It was shown that outbreaks can sometimes involve more than one Cryptosporidium species and individuals can be co-infected (McLauchlin et al. 2000; Peng et al. 1997) and that contaminated water can contain multiple species and genotypes (Xiao et al. 2006). Differences in the descriptive epidemiology for sporadic disease between C. hominis and C. parvum (Hunter et al. 2004; Nichols 2008) were demonstrated and though unrelated, the degree of genetic variation in different species (panmictic vs. clonal) can vary (Mallon et al. 2003b). The term genotype is used to refer to a taxonomic rank below the species level using a defined set of genetic markers, which for *Cryptosporidium* includes isolates with sequences sufficiently different at the SSU rRNA gene (similarity usually close to 99.5 %) but that don't have enough other data for species status. The term subtype is used when defined loci are investigated to detect variation within species. The term isolate is used for a specific population of organisms from a human or animal, and the term strain should be restricted to isolates that are maintained in culture, although these may not necessarily be genetically homogeneous. In one multi-locus study, some subtypes of C. parvum from human patients were found to be uncommon in agricultural animals (Mallon et al. 2003b), while natural infection with C. hominis in sheep and cattle can occur occasionally (Smith et al. 2005). The potential for significant disease reduction was demonstrated with interventions to reduce contamination of drinking water (Sopwith et al. 2005). Molecular epidemiology has also demonstrated the potential for identifying new transmission routes (McLauchlin et al. 2000). These themes will be explored below.

3.1.1 Phenotyping

Early in the development of *Cryptosporidium* epidemiology, phenotypic methods were used to try to delineate species and provide investigation tools (McLauchlin et al. 1998; Nichols 1992; Nichols et al. 1991). Approaches used included monoclonal antibody and polyacrylamide gel electrophoresis or isoenzyme analysis to examine differences between *Cryptosporidium* species (Nina et al. 1992a, b; Ogunkolade et al. 1993). However, such approaches did not have the sensitivity of genetic approaches and have not proved to be very useful in the practice of molecular epidemiology, other than highlighting the presence of several *Cryptosporidium* species present in different hosts.

3.1.2 Genotyping to Species Level

Application of molecular methods for typing (genotyping) *Cryptosporidium* isolates has provided improved sensitivity and specificity over phenotyping and enabled more widespread investigation of variation in clinical and subclinical infections and in food, water and environmental samples. One of the important achievements of the application of molecular epidemiology has been the identification of infection sources (that is, the hosts providing a source of human or animal infection), by delineating the species of *Cryptosporidium*, their natural hosts and the evidence for infectivity and pathogenicity of these species for humans. The rationale for species designation was reviewed in 2010 by Fayer (2010) and is discussed, along with the current taxonomy of the genus, in Chap. 1. The most current nomenclature is used here, assuming in places that the prior name correctly represents the organism. *Cryptosporidium* species and their host distribution with some molecular epidemiological information are summarised in Table 3.1.

To date human cryptosporidiosis has been strongly linked to *C. hominis*, *C. parvum*, *C. meleagridis* and *C. cuniculus*. Experimental infections conducted in human volunteers and outbreak investigations has provided further evidence (Table 3.1). In some settings, *C. canis and C. felis* are also associated with human cryptosporidiosis, and *C. ubiquitum* and *C. viatorum* are recognised as emerging human pathogens. In the case of many of the minor species (those only rarely found in humans, see Table 3.1) there is merely the occasional finding of oocysts in faeces of symptomatic patients, including *C. andersoni*, *C. suis*, *C. scrofarum*, *C. bovis*, and *C. muris*. Some species have not been described in humans. This may reflect the greater adaptation of these species to their hosts, especially those found in fish, reptiles and amphibians, that is reflected in the taxonomy (Palenzuela et al. 2010).

Table 3.1 Cryptospo	Table 3.1 Cryptosporidium species and their involvement in human infections	an infections	
Cryptosporidium			
species	Involvement in human cryptosporidiosis	Animal reservoirs of human infection	Molecular epidemiology
Species strongly linke <i>C. hominis</i> (Morgan- Ryan et al. 2002)	 Species strongly linked to human infections <i>C. hominis</i> (Morgan- The species most commonly linked to human- derived infection, either directly person-to-person (especially in daycare centres, household contacts, toileting or nappy changing), or indirectly via contaminated drinking water, recreational water, food or fomites A common cause of sporadic cases and outbreaks of cryptosporidiosis Infectivity data from experimental infections in adults indicates that ingestion of a single occyst carries a discrete probability of infection 	There is no defined animal host for this species, although low density shedding of oocysts has been reported occasion- ally from cattle, sheep and goats	High parasite genetic heterogeneity has been reported in non-industrialised countries, compared with apparently less variation in some industrialised countries (Tanriverdi et al. 2008). A genome sequence for isolate TU502, has been published (Xu et al. 2004)
C. parvum (Xiao et al. 2012)	The species commonly transmitted from young ruminants to humans, and sometimes between humans, either directly through farm animal contact or person-to-person, or indirectly through contaminated drinking water, recrea- tional water, environmental contact, food or fomites A common cause of sporadic cases and outbreaks of cryptosporidiosis Infectivity data from experimental infections in adults indicates variation between isolates, although virulence factors are as yet undefined. Ingestion of a single oocyst of some strains car- ries a discrete probability of infection	A common cause of diarrhoea in pre-weaned calves, lambs and goat kids, and also reported in foals, alpaca, llama (see Chaps. 4 and 5). Infection may be asymptomatic especially in older animals and sometimes in lambs. High oocyst output from pre-weaned animals increases risk of direct and indirect transmission to humans	 High heterogeneity in isolates from humans has been reported from many countries; some subtypes (e.g. <i>gp60</i> IIc) appear host-adapted to humans with no defined animal host A full genome sequence for the zoonotic IOWA isolate (gp60 subtype IIaA15G2R1a) has been published (Abrahamsen et al. 2004). A three-way comparison with anthroponotic <i>C. parvum</i> TU114 (gp60 IIc) and <i>C. hominis</i> identified loci where the anthroponotic <i>C. parvum</i> sequence is more similar to <i>C. hominis</i> than to the zoonotic <i>C. parvum</i>, suggesting that proteins encoded by these genes may influence the host range (Widmer et al. 2012)

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Subtyping <i>C. meleagridis</i> in Japan has identified several genotypes infecting birds (Abe 2010; Abe and Makino 2010)	May be overlooked by commonly used typing tools due to genetic similarity to <i>C. hominis</i> (Chalmers et al. 2011a). <i>C. cuniculus</i> gp60 subtype Va is found in humans and only occasionally in rabbits (including the rabbit that caused the outbreak in the UK) and Vb is found in both hosts (Zhang et al. 2012). Sea- sonal and sex distribution of <i>gp60</i> subtypes has been reported in humans in UK (Chalmers et al. 2011a)		(continued)
Associated with enteritis, diarrhoea and death in birds. Readily transmissible between birds and mammals	Caused a drinking waterborne outbreak in the UK when a wild rabbitAsymptomatic shedding of oocysts in nat- ural and experimentally infected rabbitsMay be overlooked by commonly used typing tools due to genetic similarit.Caused a drinking waterborne outbreak in the UK when a wild rabbitAsymptomatic shedding of oocysts in nat- ural and experimentally infected rabbitsMay be overlooked by commonly used typing tools due to genetic similarit.Contaminated treated water (Chalmers et al. 2011a). Has also been demonstrated to occur in sporadic dis- ease in the UK (Chalmers et al. 2011a; Elwin et al. 2013). Individual reports from France (Network 2010) and chil- dren in Nigeria (Molloy et al. 2010)Nay be overlooked by commonly used typing tools due to genetic similarit. C. <i>hominis</i> (Chalmers et al. 2011a). C. <i>cuniculus</i> gp60 subtype Va is fo in humans and only occasionally in rabbits (including the rabbit that cau the outbreak in the UK) and Vb is fo in both hosts (Zhang et al. 2012). S sonal and sex distribution of <i>gp60</i> subtypes has been reported in huma in UK (Chalmers et al. 2011a)	Diarrhoea usually seen in puppies and younger dogs; asymptomatic shedding of oocysts has been reported in older and immune- competent dogs	
One of the more common non- <i>C. parvum</i> , non- <i>C. hominis Cryptosporidium</i> spe- cies infecting humans Mainly sporadic cases, more frequent in some populations, for example in Bangkok, Thailand and Lima, Peru. Cases in UK have been linked to travel to endemic countries A possible farm-related outbreak has been reported in Sweden linked to poultry (Silverlas et al. 2012) Infectivity data from experimental infections in adults indicates mild illness	Caused a drinking waterborne outbreak in the UK when a wild rabbit contaminated treated water (Chalmers et al. 2011a). Has also been demonstrated to occur in sporadic dis- ease in the UK (Chalmers et al. 2011a; Elwin et al. 2012a). Individual reports from France (Network 2010) and chil- dren in Nigeria (Molloy et al. 2010)	The species found in dogs that has been isolated from some infected humans (Elwin et al. 2012a), and epidemiolog- ically linked to diarrhoea in children in a shanty town in Lima, Peru (Priest et al. 2006)	
C. meleagridis (Slavin 1955)	<i>C. cumculus</i> (Robinson et al. 2010)	C. camis (Fayer et al. 2001)	

Table 3.1 (continued)	(]		
Cryptosporidium species	Involvement in human cryptosporidiosis	Animal reservoirs of human infection	Molecular epidemiology
C. felis (Iseki 1979) Species occasionally f	<i>C. felis</i> (Iseki 1979) The species found in cats and has been detected in infected humans in the UK (Chalmers et al. 2011b), France (Network 2010) and Spain (Cieloszyk et al. 2012). Having a compromised immune system and being in contact with a cat were risk factors for human infection (Elwin et al. 2012a). <i>C. felis</i> has been associated with diarrhoea in children in a shanty town in Lima, Peru (Priest et al. 2006)	More common in kittens and younger cats and has been associated with diarrhoea. <i>Cryptosporidium</i> was not detected in 57 healthy kittens in the UK (Gow et al. 2009) and uncommon (4 %) in 149 kittens in New York (Spain et al. 2001), although transmission from cat to human has been reported (Egger et al. 1990)	
C. andersoni (Lindsay et al. 2000)	C. andersoni (Lindsay This species has been reported in a few et al. 2000) human infections in the UK, Australia, and Malawi (Leoni et al. 2006; Morse et al. 2007; Waldron et al. 2011a), including in HIV positive people (Agholi et al. 2013)	Occurs in the stomach of cattle (Lindsay et al. 2000), and has been reported in one study in bird facces (Ng et al. 2006). This species has also been detected in wastewaters (Ben Ayed et al. 2012; Liu et al. 2011; Nichols et al. 2005; Ruecker et al. 2013; Xiao et al. 2001a), molluscs (Miller et al. 2005; Nichols et al. 2006a) and tapwater (Castro-Hermida et al. 2008; Feng et al. 2011c)	C. andersoni subtyping of cattle strains has identified at least five MLST types (Zhao et al. 2013)
C. baileyi (Current et al. 1986)	There is a single unsubstantiated report of human infection (Ditrich et al. 1991) with this bird species, made without molecular analysis	Many bird species can be affected, partic- ularly chickens (Abbassi et al. 2000)	

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	A genome sequence has been published (Widmer and Sullivan 2012)		The frequent detection of this organism in drinking water in China without human illness being commonly identified (Feng et al. 2011c) suggests the organ- ism is poorly infectious or lacks patho- genicity for most people (continued)
Mainly shed by weaned calves with no or few clinical signs Found in kangaroos but no clinical signs are described in animals (Ryan et al. 2008; Yang et al. 2011)	Infection is of the stomach of mice; clinical A genome sequence has been published signs are not usually reported; infection (Widmer and Sullivan 2012) and oocyst shedding has been reported in other animal species	Infection is seen in pigs older than 6 weeks resulting in low levels of oocyst shed- ding and no association with diarrhoea	Infection is usually seen in pre-weaned pigs but is not associated with diarrhoea
This species is usually found in cattle (Fayer et al. 2005); small numbers of human infections have been reported of people stating cattle contact in Australia and India (Helmy et al. 2013; Ng et al. 2012) This kangaroo species been detected in a individual infected person in Australia who had contact with marsuipials (Waldron et al. 2010)	The first <i>Cryptosporidium</i> species to be described (Tyzzer 1910). It occurs, as its name suggests, in mice. <i>C. muris</i> has larger oocysts than most species and is reported to infect humans only occasionally (Tosini et al. 2010).	This species is found in farmed pigs worldwide (Kvac et al. 2013), in wild pigs (Nemejc et al. 2012) and has also been identified in fish (Reid et al. 2010) and a calf (Ng et al. 2011). To date this species has been identified in a single human immunocompetent case in Czech Republic who had contact wild pigs (Kvac et al. 2000)	This species is found in domestic pigs (Ryan et al. 2004a) and has been detected in a small number of immu- nocompetent and in HIV positive patients with diarrhoea (Cama et al. 2007; Leoni et al. 2006; Wang
C. bovis (Fayer et al. 2005) C. fayeri (Ryan et al. 2008)	C. muris (Tyzzer 1910)	C. scrofarum (Kvac et al. 2013)	C. suis (Ryan et al. 2004a)

Cryptosporidium			
species	Involvement in human cryptosporidiosis	Animal reservoirs of human infection	Molecular epidemiology
	et al. 2013; Xiao et al. 2002) in UK and Peru who had contact with pigs		
C. tyzzeri (Ren	This species is considered to be the species Infection is usually most intense in the	Infection is usually most intense in the	
et al. 2012)	first found in mice by Tyzzer in 1912	mouse ileum but no clinical signs were	
	(1 ang et al. 2011; 1 yzzer 1910). 1 here is one individual report from Czech	recorded in experimental infections	
	Republic involving contact with wild mice (Raskova et al. 2013)		
C. ubiquitum (Fayer	Has been demonstrated in infected humans A common parasite of weaned lambs	A common parasite of weaned lambs	Newly developed gp60 primers have been
et al. 2010)	in the UK (Chalmers et al. 2011b;	(Fiuza et al. 2011; Shen et al. 2011;	used to compare human and animal
	Elwin et al. 2012a), Canada (Wong and	Sweeny et al. 2011) and has been found	isolates in different geographical
	Ong 2006), Spain (Cieloszyk	in many other mammals indicating a	locations revealing differences in
	et al. 2012) and Nigeria (Molloy	broad host range encompassing	transmission (Lihua paper submitted -
	et al. 2010)	ruminants, rodents and primates. Con-	check we can state this)
		tamination of water sources,	
		stormwaters and wastewaters in many	
		geographical locations	
C. viatorum (Elwin et al. 2012c)	A species identified from infected people returning to the UK from the Indian	It is not clear whether this species is found only in humans, as an animal host has	
	subcontinent (Jellison et al. 2009),	not been identified yet	
	sporadic cases have also been reported		
	in Sweden, following visits to South		
	et al. 2013)		
Species not reported in humans	n humans		
C. fragile (Jirku	This species has been identified in toads	Infection seems host specific as infections	
et al. 2008)	(Wang et al. 2013)	of four amphibian and one fish species	
		Ialleu	

Table 3.1 (continued)

Infection of the proventriculus can lead to diarrhoea and fatalities, shedding can be prolonged and is not necessarily associated with diarrhoea Infection is limited to marsupials, without any clinical signs	Infection is seasonal, mostly in younger fish in the spring	Infection is host specific and shedding is seen in weaned calves but is thought to be asymptomatic	Infection occurs in the intestine and is seen seasonally in younger fish; has been linked to reduced growth rates	Infection is common. In snakes, infection manifests as anorexia, persistent post- prandial regurgitation, lethargy, mid-body swelling and chronic weight loss, which is usually protracted and almost always fatal. In lizards infection	is usually asymptomatic Infection is seen in lizards and snakes. Clinical signs include anorexia, pro- gressive weight loss, abdominal swelling and death, particularly in young animals
 <i>C. galli</i> has been isolated from wild and pet Infection of the proventriculus can lead to birds (Antunes et al. 2008; da Silva diarrhoea and fatalities, shedding can et al. 2010; Nakamura et al. 2009; Qi be prolonged and is not necessarily et al. 2011; Seva Ada et al. 2011) associated with diarrhoea This species is found in marsupials includ- Infection is limited to marsupials, without ing kangaroos (Yang et al. 2011) any clinical signs 	A fish species (Alvarez-Pellitero and Sitja- Infection is seasonal, mostly in younger Bobadilla 2002), first found in farmed fish in the spring bream and bass and a <i>C. molnari</i> -like species has been detected in guppies (Rvan et al. 2004b) (Ne et al. 2011)	This species is found in buffalos (Amer et al. 2013; Feng et al. 2012) and cattle (Fayer et al. 2008; Helmy et al. 2013; Liu et al. 2009; Santin and Zarlenga 2009)	C. scophthalmi (Alvarez-Pellitero et al. 2004) are fish species found in turbot	This species is common in snakes and lizards (Pavlasek and Ryan 2008; Richter et al. 2011)	This species is found in lizards and snakes Infection is seen in lizards and snakes. (Pavlasek and Ryan 2008) Clinical signs include anorexia, progressive weight loss, abdominal swelling and death, particularly in young animals
C. galli (Ryan et al. 2003) C. macropodum (Power and Ryan	<i>C. molnari</i> (Alvarez- Pellitero and Sitja- Bobadilla 2002)	C. ryanae (Fayer et al. 2008)	<i>C. scophthalmi</i> (Alvarez-Pellitero et al. 2004)	C. serpentis (Levine 1980)	<i>C. varanii</i> (Pavlasek and Ryan 2008)

(continued)

Animal reservoirs of human infection Disease not described In sheep, asymptomatic carriage; no asso- ciation of oocyst shedding and clinical symptoms in experimentally infected lambs, while infection in goat kids can be associated with diarrhoea	I able 3.1 (continued)			
-	Cryptosporidium			
-	species	Involvement in human cryptosporidiosis	Animal reservoirs of human infection	Molecular epidemiology
	C. wrairi (Vetterling et al. 1971)	C. wrairi (Vetterling et al. 1971) was identified from guinea pigs	Disease not described	
ct at. zutt) and tisit (Network at at. zutu)	<i>C. xiaoi</i> (Fayer and Santin 2009)	<i>This species</i>) is found in lambs and sheep (Fayer and Santin 2009) in many countries (Imre et al. 2013; Sweeny et al. 2011), in goats (Diaz et al. 2010), as well as in kangaroos (Yang et al. 2011) and fish (Reid et al. 2010)	In sheep, asymptomatic carriage; no asso- ciation of oocyst shedding and clinical symptoms in experimentally infected lambs, while infection in goat kids can be associated with diarrhoea	Closely related to C. bovis

Table 3.1 (continued)

However, some zoonotic species have been detected, for example *C. parvum*, *C. scrofarum* and *C. xiaoi*, in fish (Reid et al. 2010).

The genotypes of *Cryptosporidium* that have been reported comprise those isolates that have yet to be given species status, those that are sub-species of one of the recognised species and those that have been shown to be different from recognised species but are not fully characterised. Most are named after the host in which they are first reported. It is likely, based on previous experience, that several of these will, in the fullness of time, be described as separate species. Some of these genotypes have also been found in humans, including the C. hominis monkey genotype (Xiao et al. 1998) detected in humans in UK and Malawi (Elwin et al. 2012a; Mallon et al. 2003b), a chipmunk genotype I identified in chipmunks (Lv et al. 2009), squirrels, geese (Jellison et al. 2009) and reported in humans in the USA, France and Sweden (Insulander et al. 2013; Network 2010), a skunk genotype found in skunks, racoons (Chavez et al. 2012), geese (Jellison et al. 2009), an elephant seal (Rengifo-Herrera et al. 2011) and reported in human infections in the UK (Chalmers et al. 2009; Davies et al. 2009; Elwin et al. 2012a) (interestingly there are no skunks outside of zoos in the UK) and a horse genotype identified in foals (Burton et al. 2010) and reported in humans in the UK and USA (Elwin et al. 2012a).

Many other genotypes (over 50) have not been detected in humans. For example, a seal genotype (Bass et al. 2012); two muskrat genotypes (Perz and Le Blancq 2001; Zhou et al. 2004) that have also been found in samples from geese (Jellison et al. 2009); a fox genotype (Zhou et al. 2004) and three different rat genotypes (Paparini et al. 2012). This list of genotypes not found in humans is not definitive, and does not mean these types cannot cause human infection. The species and genotype host range data suggest that over time there has been selection and speciation of *Cryptosporidium* lineages not only in individual hosts, but also in host phyla. However the host range of some species appears to be wider than for others. In addition it remains likely that some "hosts" are merely transport vectors of oocysts from their environment rather than being infected themselves (e.g. *C. parvum* found in molluscs, fish, or geese).

3.1.3 Subtyping

Efforts to investigate genetic diversity within and between *Cryptosporidium* species, aimed at identifying infection sources and investigating transmission dynamics, have mainly been focussed on *C. parvum* and *C. hominis* as the most important human pathogens. *C. cuniculus, C. meleagridis, C. andersoni* and *C. muris* have also been investigated (Feng et al. 2011a; Glaberman et al. 2001; Wang et al. 2012; Zhang et al. 2012). Caution must be expressed for "subtypes" based on sequence analysis of the ssu rRNA gene alone, arising from the polymorphic nature of different copies of the gene (Xiao et al. 1999). Because of a lack of identified genetic markers, other species have been less investigated. The 60 kDa glycoprotein (*gp60*, also called *gp40/15*) gene (Strong et al. 2000) is the most common marker that has been used, either alone or as part of a multi-locus panel. However, as a protozoan with a sexual

cycle, *Cryptosporidium* species diversity is best investigated at multiple loci, although there is, as yet, no standardised multilocus subtyping scheme. There is experimental evidence that gp60 subtypes of *C. parvum* and *C. hominis* do not segregate with a multi-locus subtyping method (Widmer and Lee 2010), confirming the need for a standardised multi-locus approach to isolate identification.

3.2 Factors Influencing the Molecular Epidemiology of Human Infections

The same factors that impact the prevalence of *Cryptosporidium* in humans (Chap. 2) also influence the molecular epidemiology, i.e.: host-related factors of age, sex, immunological status, geographical location and ethnic group; exposure including hygiene, diet, water consumption, rural and urban settings, contact with human waste and livestock, food and water protection, travel and immigration; and environmental and socio-economic factors including animal pollution, malnutrition and dehydration, geography, calamities (e.g. floods), weather and climate (Nichols 2008; Nichols et al. 2006; Putignani et al. 2010). Exposure factors will also be influenced by the distribution of Cryptosporidium species in various hosts and in vehicles of transmission such as water (see Chaps. 4, 5, and 12). In addition, parasite-related infectivity and pathogenicity factors also influence the molecular epidemiology of human infections (Bouzid et al. 2013; Nichols 2008; Xiao 2008). Although 25 putative virulence factors have been proposed, none of them has yet been confirmed (Bouzid et al. 2013), yet various studies have shown differences in outcome of infection between C. parvum and C. hominis. For example, a study of sporadic cryptosporidiosis in the UK showed significantly longer duration of diarrhoea in C. hominis patients than in those with C. parvum infection (Hunter et al. 2004). Comparison of three C. parvum strains in adult volunteer studies found differences in ID50, attack rate and duration of diarrhoea (Okhuysen et al. 1999). The association between gp60 subtypes and outcome of infection is discussed later. When host and parasite factors are coupled, they help explain the molecular epidemiology and potential risks posed by different exposures and Cryptosporidium species to human health. For example, the microbiological evidence from the rare occurrence of C. felis and C. canis in humans in most settings is that dog and cat faeces are probably not a major source of human infection (Bowman and Lucio-Forster 2010). This is supported by many analytical epidemiological studies, none of which identify contact with companion animals as a risk factor for human cryptosporidiosis (Bowman and Lucio-Forster 2010). Nevertheless, some individuals are susceptible and it is sensible to advise hand washing after contact with any animal, although there does not seem to be any particular susceptibility to infection with these species in immunocompromised patients.

The common occurrence and low infectious dose of *C. parvum* and *C. hominis* in human infections (Chap. 2), and the rare occurrence of many of the other *Cryptosporidium* species, suggests that there is some species barrier that prevents many of these rarer species infecting people. Because *C. parvum* and *C. hominis* are

common in the environment as a result of animal and human faecal contamination from animal waste and sewage inputs, there may also be a relationship between infection and exposure to oocysts that are common in the environment. It has been shown that in some countries a broader range of species can be detected in all patients (Morgan-Ryan et al. 2002). This contrasts to the view that some *Cryptosporidium* species are more commonly found in immunocompromised patients than in immunocompetent ones. However, with improved protocols for detecting different species it should be recognised that the ability to detect some species has improved. A recent example of this is *C. cuniculus* that was identified as *C. hominis* using routine PCR screening methods but has been detected as a small but significant component of the UK *Cryptosporidium* cases following more detailed typing.

3.3 Genotyping and Subtyping Methods

Molecular testing or screening stools for *Cryptosporidium* using PCR protocols are much more sensitive than microscopic methods (Hadfield et al. 2011), allowing larger numbers of cases of cryptosporidiosis to be diagnosed, and improving estimates of carriage. However, assays for the detection of presence/absence of the genus may not be designed to identify species, for which conventional (single round and nested) and real-time PCRs have been described. PCR primers and conditions need to be selected carefully to amplify all *Cryptosporidium* species of interest and to avoid non-specific amplification. Any PCR-based method will be dependent, among other things, on efficient DNA extraction. This is also discussed in this section.

3.3.1 DNA Extraction from Clinical Samples

There is no recommended method, but efficient DNA extraction underpins the application of molecular assays (Elwin et al. 2012b). The most commonly investigated human samples for epidemiological purposes are stools, and extraction can be undertaken readily either directly from fresh or frozen stools or from semipurified oocyst suspensions or sediments, including immuno-magnetically separated oocysts, providing the oocysts are disrupted so that sporozoite DNA can be made available (Elwin et al. 2012b).

Options for oocyst disruption processes include boiling (Elwin et al. 2001), freeze-thaw cycles (Bialek et al. 2002; Nichols et al. 2006b) and chemical, enzymatic or mechanical treatments (Balatbat et al. 1996; Lindergard et al. 2003; McLauchlin et al. 1999; Pieniazek et al. 1999) or a combination such as bead-beating, followed by guanidine thiocyanate extraction and silica purification (Amar et al. 2007; Boom et al. 1990; McLauchlin et al. 1999; Patel et al. 1998; Zintl et al. 2009).

Options for DNA extraction include commercial spin columns, glassmilk, and chelex resin. One study reported that direct extraction by bead-beating, guanidine thiocyanate and silica purification provided slightly lower PCR positivity and higher threshold cycle (Ct) values in real-time PCR, indicating lower DNA yield, than spin-column extraction from boiled, semi-purified oocyst suspensions, although direct extraction is cheaper, and amenable to automation (Elwin et al. 2012b). *Cryptosporidium* DNA extraction has been demonstrated successfully in an automated, universal DNA extraction method for enteric bacteria, viruses and protozoan based on bead-beating (MagNa Lyser instrument, Roche Diagnostics Ltd) and lysis under denaturing conditions in the presence of proteinase-K followed by nucleic acid binding to the silica surface of magnetic particles (QIAsymphony® SP, Qiagen) (Halstead et al. 2013). Contaminants are removed by washing, and pure nucleic acid is eluted in either modified TE buffer or water.

If the original stool sample is not available, the stained microscopy slides left over from the original diagnosis can be used; the smeared material can be scraped off into lysis buffer and extracted by a suitable method (Amar et al. 2001).

Some preservatives and fixatives applied to stool samples, such as 10 % formalin, sodium acetate–acetic acid–formalin (SAF) and 2.5 % potassium dichromate, may inhibit downstream PCR. If they have not penetrated the oocysts, they can be removed by washing. In one study, *Cryptosporidium* DNA was successfully amplified from "aged" stool samples that had been collected and processed in 10 % formalin from HIV-AIDS patients by including washing steps in PBS before multiple freeze-thaw cycles and preceding stool extraction kit by 95 °C incubation in lysis buffer (QIAamp DNA stool mini kit; Qiagen (Del Chierico et al. 2011). The repeated PBS washings were deemed to remove excess formalin. Further, the selection of a relatively small size DNA target (358 bp of the gp60 gene) minimized the effect of formalin-induced DNA damage, including fragmentation and cross-linking.

Stools can contain many PCR inhibitors, including heme, bilirubin, bile salts, and complex polysaccharides. Boiling in 10 % polyvinylpolypyrrolidone before oocyst disruption and DNA extraction can reduce inhibition, alternatively spin columns appear to be effective. The use of reagents such as 400 ng/ μ l of non-acetylated bovine serum albumin in the PCR is recommended.

The choice of disruption and extraction processes will depend on the types and consistency of samples being tested, the normal procedures in the laboratory, the number of parasites present, whether other pathogens are also being sought, the laboratory resources (expertise, time, consumables and equipment available), and the requirement for any long-term archiving of material.

3.3.2 Species Identification by Conventional PCR

Conventional PCRs which either produce amplicon sizes characteristic of individual species or requiring downstream analysis usually by restriction enzyme digestion identifying restriction fragment length polymorphisms (RFLP), or by sequencing, have been described in detail previously (Chalmers et al. 2011c). There is no single recommended method, although sequencing provides the definitive identification, as some species/genotypes have RFLP patterns that are indistinguishable.

Conventional PCR assays have targeted various genes including small subunit (SSU) rRNA, *Cryptosporidium* oocyst wall protein (COWP), thrombospondin-related adhesive proteins, 70-kDa heat shock protein (HSP70), and actin genes (Chalmers et al. 2011c) and were described in detail (Xiao 2008). Sequencing the SSU rRNA gene is regarded as the benchmark and has been widely used in many studies, because its analysis offers improved sensitivity due to the multiple copies of the gene and because it has both conserved and polymorphic regions. Use of other gene targets may lead to under-estimation of the presence of some species, as the designed primers don't amplify DNA from all species. However, conventional PCR is time-consuming and resource-heavy.

An alternative method for species identification is capillary electrophoresis single strand conformation polymorphism (CE-SSCP). One study found that species could be readily differentiated based on the SSCP mobility of amplified ssu rRNA gene molecules, and clones that differed by single-nucleotide polymorphisms could be distinguished in this way (Power et al. 2011).

3.3.3 Species Identification by Real-Time PCR

Real-time PCR methods for *Cryptosporidium* were first developed over 12 years ago (Amar et al. 2003; Fontaine and Guillot 2002; Higgins et al. 2001; MacDonald et al. 2002) and can provide a more rapid result than conventional PCR. When coupled with modern filtration sampling methods and immunomagnetic separation, real-time PCR can also be used for examining *Cryptosporidium* isolates from water (Fontaine and Guillot 2003). Real-time PCR offers advantages of improved work flow and efficiency for large-scale molecular surveillance, as well as reduced handling avoiding cross contamination and carryover of PCR products, amplification performance monitoring, and lower consumables costs and staff time (Hadfield et al. 2011).

There is no recommended method and various real-time PCR assays have been described for the identification of *Cryptosporidium* species in human clinical samples, especially where many years of base-line work has established the usual prevalence of each species (Table 3.2). In the UK, where the vast majority of clinical cases are caused by C. parvum or C. hominis (Chalmers et al. 2009, 2011c), typing has evolved to streamline the process: Hadfield et al. (2011) initially developed and applied a TaqMan minor groove binding (MGB) probe-based assay that provides specific identification of C. hominis and C. parvum by targeting a locus of unknown function, LIB13 (Tanriverdi et al. 2003), while simultaneously detecting all known Cryptosporidium species and genotypes by amplification of a region of the SSU rRNA gene (Morgan et al. 1997) with identification by direct sequence analysis. Designed for investigation of human samples, semi-purification of oocysts prior to DNA extraction (Elwin et al. 2001) effectively screens out many potential cross-reacting genera which can be problematic when targeting this gene which is ubiquitous and conserved among eukaryotes. This assay improved typability of microscopy-positive stools to 99.5 % compared with 96.9 % by

Table 3.2 Some real-time	-time PCR approaches used in	investigation of hun	PCR approaches used in investigation of human samples for Cryptosporidium	
Cryptosporidium amplification target	Detection platform and chemistry	Purpose of application	Most suitable application	Reference
SSU rRNA gene	Light cycler (Roche) Hybridisation probes	Species identification	Differentiation of intestinal species	Limor et al. (2002)
β-tubulin gene	Light cycler (Roche) SYBR Green/Hybridisation probes	Species identification	Differentiation of C. parvum and C. hominis	Tanriverdi et al. (2002)
Lib13 locus	Light cycler (Roche) and SYBR Green	Species identification	Differentiation of C. parvum and C. hominis	Tanriverdi et al. (2002)
COWP gene	Light cycler (Roche) SYBR Green	Species identification	Real-time, then RFLP as melt curves couldn't differentiate	Amar et al. (2004)
DNA-J like protein gene	iCycler (BioRad) MGB TaqMan	Genus detection in diagnostic narastrology	Used in a multiplex assay for simultaneous detection of <i>Entamoeba histolytica</i> , <i>Giardia duodenalis</i> , and <i>C. parvum</i>	Verweij et al. (2004)
SSU rRNA gene	iCycler (BioRad) Scorpion probes	Species identification	Detection of <i>Cryptosporidium</i> spp. Specific detection of <i>C. parvum</i> and <i>C. meleagridis</i>	Stroup et al. (2006)
SSU rRNA gene	iCycler (BioRad) SybrGreen	Genus detection	Detection of <i>Cryptosporidium</i> spp. in cell culture and human faecal samples	Parr et al. (2007)
COWP gene	iCycler (BioRad) TaqMan	Genus detection in diagnostic parasitology	Used in a multiplex assay for simultaneous detection of Haque et al. (2007) <i>Cryptosporidium</i> spp. <i>E. histolytica</i> and <i>G. duodenalis</i>	Haque et al. (2007)
SSU rRNA gene and iCycler (BioRad) + Lib13 locus Mx3000p (Stratage	iCycler (BioRad) + Mx3000p (Stratagene)	Genus detection and species identification	Detection of <i>Cryptosporidium</i> spp. Differentiation of <i>C. parvum</i> and <i>C. hominis</i>	Jothikumar et al. (2008)
SSU rRNA gene	Rotor-Gene 6000 (Qiagen) Melt curve (dye not stated)	Genus detection	Multiplexed detection of Cryptosporidium spp., Dientamoeba fragilis, Entamoeba histolytica and G. duodenalis	Stark et al. (2011)

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SSU rRNA gene, and Lib13 locus.	SSU rRNA gene, and Rotor-Gene 6000 (Corbett Research) Lib13 locus, MGB TaqMan	Species identification	Differentiation of <i>C. parvum</i> and <i>C. hominis</i> . Detection and identification by sequencing of other <i>Cryptosporidium</i> spp.	Hadfield et al. (2011)
DNA-J like protein gene	ABI PRISM 7300 (Life Technologies) Dual-labelled probe	Genus detection	Detection of Cryptosporidium spp.	Calderaro et al. (2011)
gp60 Va and Vb alleles	Rotor-Gene 6000 (Corbett Research) Melt curve analysis (Sybr Green)	Species detection and subtype family identification	Species detection Detection of <i>C. cuniculus</i> . Identification of Va and Vb and subtype families subtype family identification	Hadfield and Chalmers (2012)
SSU rRNA gene	ViiA7 (Life Technologies) TaqMan singleplex array card	Genus detection	 Detection of <i>Cryptosporidium</i> spp., with parallel detection of enterotoxigenic <i>E. coli</i>, enteropathogenic <i>E. coli</i>, enteroaggregative <i>E. coli</i>, and Shiga-toxigenic <i>E. coli</i>, <i>Shigellal</i> enteroinvasive <i>E. coli</i>, <i>Salmonella</i>, <i>Campylobacter jejuni/Campylobacter coli</i>, <i>Vibrio cholerae</i>, and <i>Clostridium difficile</i>, <i>G. duodendis</i>, <i>E. histolytica</i>, <i>Ascaris lumbricoides</i>, <i>Trichuris trichiura</i>, rotavirus, norovirus group 2, adenovirus, astrovirus, and sapovirus 	Liu et al. (2013)
DNA-J like protein gene	ABI 7500 (Life Technologies) Probe not stated	Genus detection	Detection of <i>Cryptosporidium</i> spp. multiplexed <i>E. histolytica</i> and <i>G. duodenalis</i>	McAuliffe et al. 2013 J Infect 67:122–129
SSU rRNA gene	Platform not stated TaqMan and hybridisation probes	Genus detection and species identification	Single-tube detection of <i>Cryptosporidium</i> spp. and identification of <i>C. hominis/C. cuniculus</i> and <i>C. parvum</i>	Mary et al. (2013)
DNA J-like protein	ABI 7900HT Fast/ABI 7500 (Applied Biosystems) TaqMan MGB	Genus detection in diagnostic parasitology	Detection of <i>Cryptosporidium</i> spp., with parallel detec- Mejia et al. (2013) tion of <i>A lumbricoides, Necator americanus, Ancylostoma duodenale, G. duodenalis, E. histolytica, T. trichiura</i> , and <i>Strongyloides stercoralis</i>	Mejia et al. (2013)

PCR-RFLP COWP gene and increased the detection of unusual species (Chalmers et al. 2013). The assay originally incorporated an exogenous internal control (IC) for the identification of inhibited reactions. The IC was considered superfluous for reference genotyping of clinical samples as no inhibition was identified in analysis of over 2,000 samples, largely due to the semi-purification step in sample preparation (Elwin et al. 2012b), but should be considered when using PCR for primary testing. Because the vast majority of samples are *C. parvum* or *C. hominis*, and mixed infections were detected in <1 % of samples, the assay has since been re-formatted (Hadfield et al., in preparation) as a single tube, multiplex PCR using re-designed *C. hominis* primers and probe based on the CpA135 gene (Tosini et al. 2010) that avoid amplification of *C. cuniculus* which is co-detected in most assays (Chalmers et al. 2009). The reference test algorithm based on this single multiplex real-time PCR includes re-testing the small number of samples that do not amplify in the *C. parvum/C. hominis* assay by amplification of a region of the SSU rRNA gene with identification by direct sequence analysis.

Alternative molecular algorithms need to be applied for primary testing incorporating genotyping, where it is not known whether *Cryptosporidium* are present or absent. For example, one paper describes a real-time PCR assay for detection and quantification of all *Cryptosporidium* species with a conserved sequence TaqMan MGB probe that is then followed by specific identification of *C. hominis* and *C. parvum* in positive samples using a duplex assay with specific TaqMan MGB probes, performed on the same amplicon (Mary et al. 2013). However, one limitation of the assay pointed out by the authors is that other *Cryptosporidium* species cannot be characterized. Other papers describe multiplex assays for primary testing of stools for a variety of parasites (Table 3.2).

A TaqMan array 384 well real time PCR technique has been used for screening for a range of 19 enteric pathogens including *Cryptosporidium* (Liu et al. 2013). Similar approaches have been used by others. If applied uniformly this sort of approach could give a much better uniformity of surveillance than current targeted approaches. The singleplex real-time PCR format has advantages over multiplex approaches (Haque et al. 2007; Jothikumar et al. 2009; Nazeer et al. 2013; Stark et al. 2011; Taniuchi et al. 2011), which have predominantly been focussed on parasites, in covering a broader spectrum of pathogens and possibly being amenable to use as a single screening method for all the common enteric pathogens. Re-formatting the assay targets would permit identification of *Cryptosporidium* species/genotypes.

Further developments in PCR-based detection and quantitation include digital PCR which has the potential to have utility in food and water microbiology, where more precise quantitation is needed (Morisset et al. 2013).

3.3.4 Characterisation of Cryptosporidium in Stools by Alternative Methods

An alternative to PCR-based methods is loop-mediated isothermal amplification (LAMP), based on autocycling strand-displacement DNA synthesis by *Bst* polymerase which may overcome the inhibitory potential of some samples; one study reported a high prevalence of *Cryptosporidium* in animal faeces that were negative by PCR (Bakheit et al. 2008), and another demonstrated improved detection over microscopy (Nago et al. 2010) although improved specificity is required for species characterisation and independent evaluation is required.

The presence of two extra-chromosomal linear virus-like double stranded RNA elements in some *Cryptosporidium* species has been utilised in a heteroduplex mobility assay (HMA) where heterogeneity between two segments of DNA, one being a reference sequence and the other a test sample, is identified; heating to denature the DNA into single strands and slow cooling encourages annealing of both the original homoduplex and the formation of a heteroduplex comprising the two different start strands. These can be differentiated by the slower PAGE migration by the heteroduplexes. One group in the UK developed a HMA for epidemiological investigations (Leoni et al. 2003). Although unusual strains identified by HMA were confirmed by sequencing, and epidemiological utility identified, this method has not been widely adopted.

3.3.5 GP60 Subtyping Methods

Sequencing the hypervariable gp60 gene has been used to further classify some *Cryptosporidium* species and genotypes, most commonly *C. parvum* and *C. hominis*, demonstrating extensive intra-species diversity. This analysis has been used to increase our understanding of the epidemiology of human cryptosporidiosis and to assist in outbreak investigations and source tracking of contamination of food and water.

The gp60 gene encodes two surface glycoproteins, gp15 and gp40, and contains hyper-variable regions and a microsatellite sequence of tandem tri-nucleotide repeats coding for the amino acid serine at the 5' (gp40) end of the gene (Strong et al. 2000). GP60 subtype nomenclature is based on that proposed for *C. parvum* by Sulaiman et al. (2005) and *C. hominis* by Cama et al. (2007). A hyper-variable region towards the 3' end of the gene is used to classify subtype families and provide the first part of the nomenclature: for *C. hominis* termed Ia, Ib, Id, etc. and for *C. parvum* IIa, IIb, IIc etc. Note there is now no Ic as the isolate to which it was ascribed was actually *C. parvum* and is re-classified as IIc; this family is regarded as human-restricted *C. parvum* as there is no defined animal host, although IIc has been reported in a hedgehog (Dyachenko et al. 2010), and a IIc isolate has been shown to be infectious to immunosuppressed mice (Tanriverdi et al. 2007). The second part of the nomenclature, used to designate the "subtype", is based on the

number and combination of various serine repeats, whether TCA (represented by the letter A), TCG (represented by the letter G), or TCT (represented by the letter T). A third expression may be used for some isolates which have one or more characteristic sequences repeated immediately after the serine microsatellite region, and is represented by the letter R and a number. Finally, where isolates diverge at the 3' end of the gene but are insufficiently different to form a new subtype family, lower case letters are used to indicate the difference. Readers are referred to standard nomenclature as shown in Table 3.3. Typical examples are *C*. *hominis* IbA10G2 (family Ib with ten TCA and two TCG serine repeats) and *C*. *parvum* IIaA15G2R1 (family IIa with 15 TCA and two TCG serine repeats and one characteristic repeat sequence).

The most commonly used gp60 assay is a nested PCR using external primers AL3531 and AL3535 which amplify ~850 to ~900 bp region of the gene and the internal primers AL3532 and AL3534 which amplify a ~810 to ~870 bp region (Alves et al. 2003). The sensitivity of the assay is not as good as that described for nested SSU rRNA (Chalmers et al. 2005), and an alternative, amplifying a shorter product of ~358 to ~400 bp, can be helpful (Sulaiman et al. 2005). Commonly used gp60 primers do not amplify species other than *C. parvum* and *C. hominis*, and different primers have been developed, for example for *C. meleagridis* (Glaberman et al. 2001) and *C. fayeri* (Power et al. 2009).

Sequencing the gp60 gene has identified a large number of distinct "subtypes" within *C. hominis* and *C. parvum* from humans, which suggested genetic 'richness' (i.e. high number of distinct types) but low diversity (i.e. the relationship between the number and relative prevalence of each those distinct types) (Jex and Gasser 2010). However, at a more local level, this is dependent on the population studied (see Sect. 3.4 and Table 3.4). As indicated earlier there is the need for a standardised multi-locus approach for subtyping *C. parvum* and *C. hominis*.

Alternatives to sequencing the *gp60* gene have been investigated for comparing isolates. PCR-based restriction endonuclease fingerprinting was used by one group to measure mutations in the gene, allowing single-nucleotide substitutions for amplicons of approximately 1 kb in size to be detected (Pangasa et al. 2010). Another group used a dual fluorescent terminal-restriction fragment length polymorphism (T-RFLP) analysis of the gp60 gene to investigate the genetic diversity of *Cryptosporidium* subtype populations in a single host infection. Variation within one human *Cryptosporidium* sample and mouse samples from seven consecutive passages with *C. parvum* was revealed, showing that differences in the ratio of subtype populations occur between infections (Waldron and Power 2011). Single strand conformational polymorphism analysis has also been used (Jex et al. 2007b; Wu et al. 2003) and can provide a rapid, cost-effective alternative for differentiation in population-based studies but sequencing would be required to confirm subtypes. Fragment sizing of gp60 amplicons has also been used in multilocus subtyping, which is discussed below.

As a subtyping tool, investigation of the gp60 gene offers both advantages and disadvantages. It is the most variable gene identified so far, and there is a large amount of sequence data available. The data are readily generated, although there is no single reference repository for comparison against reference subtypes and

Species and family	Dominant trinucleotide repeat	Other repeat (R)	GenBank accession no.
C. hominis			
Ia	TCA	AA/GGACGGTGGTAAGG	AF164502 (IaA23R4)
Ib	TCA, TCG, TCT	I	AY262031 (IbA10G2), DQ665688 (IbA9G3)
Id	TCA, TCG	I	DQ665692 (IdA16)
Ie	TCA, TCG, TCT	1	AY738184 (leA11G3T3)
If	TCA, TCG	1	AF440638 (IfA19G1)
Ig	TCA	1	EF208067 (IgA24)
Ih	TCA, TCG	1	FJ971716 (IhA14G1)
Ii (C. hominis monkey genotype)	TCA	1	HM234173 (IiA17)
Ij ^a	TCA	I	JF681174 (IjA14)
C. parvum			
Па	TCA, TCG	ACATCA	AY262034 (IIaA15G2R1), DQ192501 (IIaA15G2R2)
IIb	TCA	I	AF402285 (IIbA14)
Пс	TCA, TCG	1	AF164491 (IIcA5G3a)
	TCA, TCG	1	AF164501 (IIcA5G3b)
	TCA, TCG	I	EU095267 (IIcA5G3c)
	TCA, TCG	I	AF440636 (IIcA5G3d)
	TCA, TCG	I	HM234172 (IIcA5G3e)
	TCA, TCG	1	HM234171 (IIcA5G3f)
	TCA, TCG	1	Accession no. not available; IIcA5G3g (REF)
	TCA, TCG	I	Accession no. not available; IIcA5G3h (REF)
	TCA, TCG	I	AM947935 (IIcA5G3i)
	TCA, TCG	1	HQ005749 (IIcA5G3j)
	TCA, TCG	I	JN867336 (IIcA5G3k)
Пd	TCA, TCG	1	AY738194 (IIdA18G1)
IIe	TCA, TCG	1	AY382675 (IIeA12G1)
Πf	TCA	1	AY738188 (IIfA6)

Table 3.3 (continued)			
Species and family	Dominant trinucleotide repeat	Other repeat (R)	GenBank accession no.
IIg	TCA	1	AY873780 (IIgA9)
IIh	TCA, TCG	1	AY873781 (IIhA7G4)
IIi	TCA	1	AY873782 (IIiA10)
IIk	TCA	I	AB237137 (IIkA14)
ΠΙ	TCA	I	AM937006 (IIIA18)
IIm	TCA, TCG	1	AY700401 (IImA7G1)
IIn	TCA	1	F1897787 (IInA8)
По	TCA, TCG	1	JN867335 (IIoA16G1)
C. meleagridis			
IIIa	TCA, TCG	1	AF401499 (IIIaA24G3)
IIIb	TCA, TCG	1	AF401501 (IIIbA13G1), AB539720 (IIIbA20G1)
IIIc	TCA	1	AF401497 (IIIcA6)
IIId	TCA	I	DQ067570 (IIIdA6)
IIIe	TCA, TCG	1	AB539721 (IIIeA20G1)
C. fayeri			
IVa	TCA, TCG, TCT	1	FJ490060 (IVaA11G3T1)
IVb	TCA, TCG, TCT	1	FJ490087 (IVbA9G1T1)
IVc	TCA, TCG, TCT	1	FJ490069 (IVcA8G1T1)
IVd	TCA, TCG, TCT	1	FJ490058 (IVdA7G1T1)
IVe	TCA, TCG, TCT	1	FJ490071(IVeA7G1T1)
IVf	TCA, TCG, TCT	1	FJ490076 (IVfA12G1T1)
C. cuniculus			
Va	TCA	1	FJ262730 (VaA18)
Vb	TCA	1	FJ262734 (VbA29)
Horse genotype			
VIa ^b	TCA, TCG	I	FJ435960 (VlaA11G3), DQ648547 (IIjA15G4)
VIb	TCA	1	FJ435961 (VIbA13)

C. wrairi			
VIIa	TCA, TCT	I	GQ121020 (VIIaA17T1)
Ferret genotype			
VIIIa	TCA, TCG	I	GQ121029 (VIIIaA5G2)
C. tyzzeri			
IXa	TCA	A/GTTCTGGTACTGAAGATA	A/GTTCTGGTACTGAAGATA GQ121030 (IXaA6R3), AY378188 (IIfA6R2)
IXb	TCA	I	HM234177 (IXbA6R2)
Mink genotype			
Xa	TCA, TCG	1	HM234174 (XaA5G1)
Opossum genotype			
XIa	TCA, TCG, TCT	I	HM234181 (XIaA4G1T1)
C. ubiquitum			
XII ^c	1	1	Not available
Hedgehog genotype			
XIII	TCA	ACATCA	KC305643 (XIIIaA22R9)
^a Baboon isolate, originally referred to as Ii (ref) ^b Originally referred to as IIj ^c Lihua Xiao, personal communication	ed to as Ii (ref) ation		

3 Molecular Epidemiology of Human Cryptosporidiosis

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Geographical region		C. p	arvu	n cat	C. parvum cases subtyped	ubtyp	bed		C. I	iomi	C. hominis subtyped	ubtyp	peq		I		
UN) and country	Patient group	z	Па	IIb) IIc	: IId	I IIe	Other	Z	Ia	Ib	Id	Ie	If	Ig	Ih	
Europe																	
Belgium	Sporadic cases	9	4	0	1	1	0	0	13	0	13	0	0	0	0	0	Geurden et al. (2009)
England, North	Sporadic cases	Not	inve	Not investigated	ted				8	0	×	0	0	0	0		Glaberman et al., (2002)
France	Waterborne outbreak	1	1	0	0	0	0	0	22	0	21	-	0	0	0		Cohen et al., (2006)
Ireland	Sporadic cases	79	78	0	0	1	0	0	25	0	25	0	0	0	0	0	Zintl et al. (2009)
Ireland	Sporadic cases	249	170	0	0	0	0	0	Not	inve	Not investigated	ted					Zintl et al. (2011)
Italy	Unspecified patients	Non	e ide	None identified	ģ				5	0	2	0	0	0	0		Wu et al. (2003)
Italy	AIDS patients	×	4	0	4	0	0	0	1	-	0	0	0	0	0	0	Del Chierico et al. (2011)
Netherlands	Sporadic cases ^a	13	6	0	1	б	0	0	64	0	63	-	0	0	0	0	Wielinga et al. (2008)
Northern Ireland	Waterborne outbreak, spo-	52	52	0	0	0	0	0	68	1	67	0	0	0	0		Glaberman et al. (2002)
-				¢		¢	¢	c	,	0	¢	¢	¢	Ŧ	¢		
Portugal	Unspecified	4		0	m	0	0	0	m	0	0	0	0	-	0		Peng et al. (2001)
Portugal	HIV + adults ^a	25	6	-	2	×	0	0	15	-	10	1	0	-	0	0	Alves et al. (2006)
Slovak Republic	Family members	Not i	inve	investigated	ted				1	0	-	0	0	0	0	0	Ondriska et al. (2013)
Slovenia	Children and adults	31	29	0	1	0	0	1	0	-	1	0	0	0	0	0	Soba and Logar (008)
Slovenia	Patients with gastro- intestinal disorders	21	21	0	0	0	0	0	71	0	53	12	0	1	ŝ	0	Koehler et al. (2013)
Slovenia	Unspecified	Э	ŝ	0	0	0	0	0	Noi	le id	None identified	ed					Peng et al. (2001)
Sweden	2 Foodborne outbreaks	27	4	0	0	23	0	0	0	0	1	-	0	0	0	0	Gherasim et al. (2010)
Sweden	Sporadic cases	107	69	0	11	24	-	2xIIo	63	S	44	10	1	С	0	0	Insulander et al. (2013)
Switzerland	HIV+	0	-	0	0	1	0	0	С	0	0	-	0	0	0	0	O'Brien et al. (2008)
UK	Unspecified	Non	e ide	None identified	ģ				б	0	б	0	0	0	0	0	O'Brien et al. (2008)
UK	Sporadic cases	69	56	0	1	6	0	б	Not	inve	Not investigated	ted					Chalmers et al. (2011b)
UK	Patients from farms ^a	11	11	0	0	0	0	0	Not	inve	Not investigated	ted					Smith et al. (2010b)
UK	Sporadic cases	Not	inve	Not investigated	ted				101	S	5 93	0	0	-	0	0	Chalmers et al. (2008)
1117			;	4													

	Chalmers et al. (2005)	Jex et al. (2008)	Jex et al. (2007a)	O'Brien et al. (2008)	Waldron et al. (2009)	Ng et al. (2012)	Waldron et al. (2011a)	Ng et al. (2010b)	Mayne et al. (2011)	Jex et al. (2008)	Ng et al. (2010b)	Ng et al. (2008)	Grinberg et al. (2011)	Grinberg et al. (2008a)	Ng et al. (2010b)		Ong et al. (2008)	Budu-Amoako et al. (2012)	Trotz-Williams et al. (2006)		Xiao et al. (2009)	Sulaiman et al. (2001)	Xiao and Ryan (2004)	Herges et al. (2012); Gatei et al. (2006)	Feltus et al. (2006)	Hira et al. (2011)	Ajjampur et al. (2007)	(continued)
	0	0	0	0	0		0	0	0			0			0		0		0		0				0	0	0	
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	0	ŝ	0	1	0	None identified	4	0	0	Э	0	0		Not investigated	0		0	Not investigated	0		43	11		Not investigated	0	×	35	
	12	38	74	41	37	None	164	48	×	38	195	3	0	Not	21		4	Not	4	0	51	55		Not	-	48	47	
	6 6 0 0 0 0 0	24 23 0 1 0 0 0	23 18 0 5 0 0 0	9 8 0 1 0 0 0	$32 \ 30 \ 0 \ 1 \ 1 \ 0 \ 0$	7 5 0 0 1 0 0	80 79 0 0 1 0 0	None identified	Not investigated	24 23 0 1 0 0 0	49 48 0 0 1 0 0	4 4 0 0 0 0 0 0 0 0	4 4 0 0 0 0 0 0 0 0	41 41 0 0 0 0 0	3 3 0 0 0 0 0		None identified	5 5 0 0 0 0 0	6 4 0 0 0 0 2	7 7 0 0 0 0 0	5 5 0 0 0 0 0	Not investigated	6 1 0 5 0 0 0	166	30 30 0 0 0 0 0	4 0 0 0 0 0 4 x Hm	7 0 0 7 0 0 0	
	Sporadic cases	Unspecified	Unspecified	Children ^a	Unspecified	Farm dwellers ^a	Sporadic cases ^a	Outbreak	Outbreak	Sporadic cases	n Sporadic cases	ו Unspecified ^a	Outbreak ^a	Sporadic cases ^a	Outbreak		Outbreak	Sporadic cases ^a	Adults ^a	Outbreak	Cases	Various	IS HIV + adults	vest Cases	Unspecified	Children	Children	
Oceania	Australia	Australia	Australia	Australia	Australia	Australia	Australia	Australia	Australia	Australia	Australia, Western	Australia, Western	New Zealand	New Zealand	Australia	Northern America	Canada	Canada	Canada	USA	USA, 4 states	USA	USA, New Orleans	USA, Upper Midwest	USA, Wisconsin Southern Asia	Bangladesh	India	

lable 3.4 (continued)																	
Geographical region		$C. p_0$	arvur	n cas	es su	C. parvum cases subtyped	pa		C.h	omi	C. hominis subtyped	ıbtyp	bed				
(UN) and country	Patient group	z	IIa	IIb		IIc IId IIe	IIe	Other	z	Ia	Ib	Id	Ie	If	Ig	Ih	
India	HIV + adults	6	0	5	з	0	0	1	28	9	×	5	0	×	0	0	Muthusamy et al. (2006)
India		14	0	0	9	5	ю	0	39	12	1	×	13	Ś	0	0	
India	Children	9	0	0	1	1	0	2xIIm, 2xIIn	59	16	11	7	16	5	0	0	Ajjampur et al. (2010)
India	Unspecified	0	0	0	0	0	0	0	٢	S	-	1	0	0	0		Peng et al. (2001)
India, Kolkata	Children with diarrhoea	Not	inves	Not investigated	pe				43	5	15	16	ŝ	0	0		Gatei et al. (2007)
Iran	Children ^a	22	٢	0	0	15	0	0	б	0	0	-	0	7	0		Nazemalhosseini-Mojarad et al. (2011)
Nepal	Unspecified patients	None	e idei	None identified	q				9	9	0	0	0	0	0		Wu et al. (2003)
Tehran, Iran	Children	17	9	0	0	11	0	0	7	0	0	0	0	0	0		Taghipour et al. (2011)
Sub-Saharan Africa																	
Ethiopia	Patients with diarrhoea	13	13	0	0	0	0	0	-	0	1	0	0	0	0	0	Adamu et al. (2010)
Kenya	Paediatric population	23	0		17			2xⅢi,	28	10	10	4	Э	0	0	-	Molloy et al. (2010)
								2xIIm									
Kenya	Unspecified	4	0	4	0	0	0	0	0	0	0	0	0	0	0		Peng et al. (2001)
Madagascar	Children	1	0	0	-	0	0	0	10	Э	0	4	-	0	0		Areeshi et al. (2008)
Malawi	Children	0	0	0	1	0	1	0	26	4	S	Ξ	9	0	0	0	Peng et al. (2003)
Nigeria	Patients	1	0	0	0	0	1	0	0	0	0	0	0	0	0	0	Maikai et al. (2012)
Nigeria	Children	0	0	0	0	0	0	0	б	0	1	0	0	0	0	0	Ayinmode et al. (2012)
South Africa	HIV + children	5	0	0	S	0	0	0	15	-	4	S	S	0	0	0	Leav et al. (2002)
Uganda	Children	15	0	0	10	0	0	5	13	4	1	-	Г	0	0	0	Akiyoshi et al. (2006)
Northern Africa																	
Egypt Ismailia province Western Asia	Children ^a	14	5	0	0	Г	0	0	Did	not	Did not amplify	ify					Helmy et al. (2013)
Iordan	Children	13	6	0	ç	ø	0	0	15	\sim	ø	٢	0	0	0		Hiiianni at al (2010)
			5		10			- c	5 -		, c					¢	
Kuwait	Children	61 61	50 F0		12	20 20		- 0	52	⊃ ∞	7 C	12	- 0				Idbal et al. (2011)
Yemen	Patients attending clinic	32	٢	0	0	0	0	0	1	0	0	0	1	0	0		Alyousefi et al. (2013)

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Southeast Asia									,								
aysia	HIV/AIDS patients	9	2	0	0	0		-	m	0	m	0	0	0	0	Lim et al. (2011)	
aysia	HIV + patients	13	12	0	0	-	0	0	S	ŝ	0	0	0	0	0	Iqbal et al. (2012)	
Eastern Asia																	
na		Non	None identified	ntifie	q				6	-	9	0	0	0	0	Wang et al. (2011)	
China	Unspecified	1	0 1 0 0	-	0	0	0	0	7	0	1	-	0	0	0	Peng et al. (2001)	
an	Unspecified ^a	0	1	0	1	0	0	0	б	-	1	0		0	0 0	0 Abe et al. (2010)	
an	Unspecified patients	1	1	0	0	0	0	0	7	1	0	0	-	0	0	Wu et al. (2003)	
Latin America and Caribbean	oean																
Cuba	Children	Non	None identified	ntifie	q				9	-	æ	0	0	0	0	Pelayo et al. (2008)	
Guatemala	Unspecified	1	1 0 0 1 0	0	-	0	0	0	б	0	0	0	ŝ	0	0	Peng et al. (2001)	
Jamaica	HIV + adults	7	0	0	٢	0	0	0	25	0	22	0	З	0	0	Gatei et al. (2008)	
Peru	HIV + adults	22	0	0	22 (0	0	0	141	35	39	40	13	13 0 0	0	Cama et al. (2007)	
Peru	Children	15	0	0	15	0	15 0 0 0	0	56	15	56 15 16 7 15 3 Mixed	٢	15	3 M	ixed	Cama et al. (2008)	
Studies included testing of animals	g of animals																i i

non-standardised nomenclature has been used by some researchers. Investigation of samples from animals and humans has provided information about the distribution of families and subtypes useful in supporting (or otherwise) zoonotic transmission (see below). However, the sequence similarity between isolates and the relationship to similarity at other loci is not extensively known. The proteins encoded by the gene have a functional role in invasion of host cells and initiation of disease. As such, gene products are exposed to the host immune response, and the gene is likely to be under considerable selective pressure, reflected by its diversity. Interestingly, when Widmer (2009) compared the distribution of serine repeat length in *C. parvum* isolates from humans with those from animals, the former showed a bi-modal distribution with more shorter repeats than occur in animals, indicative of host-driven selective pressure.

Recombinant progeny of *C. parvum* strains have been used to examine co-infections in immunosuppressed mice (Tanriverdi et al. 2007). The recombinants were obtained by crossing two genetically distinct isolates co-infected in immunosuppressed mice and then by targeted propagation in mice of the progeny oocysts from populations lacking one parental allele at one or more loci. Sixteen progeny clones were genotyped at 40 loci located on each of the eight chromosomes and a range for the meiotic crossover frequency was determined. Like other Apicomplexan parasites, *C. parvum* exhibited a high rate of recombination. Evolution is therefore likely to be rapid as recombination events between different strains can occur; new subtypes are constantly being placed on GenBank. More compatible with the biology of *Cryptosporidium*, and the presence of a sexual phase in the life cycle, is a multilocus subtyping scheme, for both epidemiological and population biology investigations.

3.3.6 Multilocus Typing Methods

Multilocus genotyping (MLG) methods have become more widely used to improve on the discrimination offered by gp60 single locus analysis. Most often this involves investigation of amplified PCR products from multiple loci to produce either a multilocus fragment type (MLFT) or multilocus sequence type (MLST). MLFT identifies variations in the length of microsatellite (repeating unit lengths up to 4 bp) or minisatellite (>4 bp) markers, driven by the number of tandem repeats and is measured cost-effectively by gel or capillary electrophoresis. MLST, as the name suggests, is based on the heterogeneity identified by DNA sequencing providing not only the number but also variation in the markers, which may also include those with only single nucleotide polymorphisms (SNPs), providing improved discrimination between *Cryptosporidium* isolates (Gatei et al. 2006).

One of the most striking features of multi-locus investigations of *C. parvum* and *C. hominis* is the diversity of loci that have been investigated. There is no standardised MLFT or MLST scheme, and a variety of loci have been used in global, regional and local parasite population studies and in epidemiological investigations (Table 3.5). Robinson and Chalmers (2012) attempted to use

		Number of MLGs identified	ILGs			
Location	Samples subtyped	Loci investigated C. hominis C. parvum	C. hominis	C. parvun	Key findings	Reference
Global comparison studies Investigation of population structure in <i>C. hominis</i> from US, Kenya, Australia, Guatamala	62 <i>C. hominis</i> samples	gp60 hsp70 Poly-T SSR-1 SSU rDNA TRAP-C2	6		The same number of MLGs and gp60 subtypes were identified. Did not iden- tify recombination, supporting clonal popu- lation but acknowledged the need for subsequent studies involving the use of more polymorphic genes	Sulaiman (2001)
Investigation of geographi- cally distinct isolates from Kolkata, India, Nairobi, Kenya, Lima, Peru and New Orleans, USA	69 <i>C. hominis</i> sporadic human cases	cp47 cp56 dzhrgp gp60 hsp70 msc6-7 Mucin1	34	1	Geographical clustering was observed when gp60 excluded	Gatei et al. (2006)
Investigation of genetic variability in isolates from various countries (USA, Scotland, Israel, Uganda, Australia)	Reference isolates of human and animal origin: 10 <i>C. hominis</i> , 16 <i>C. parvum</i>	rpgr 1887 5B12 Cb492	10	16	Differential rate of repeat lengths inferred for <i>C. hominis</i> and <i>C. parvum</i>	Tanriverdi (2006)

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Location Samı						
		Number of MLGs identified	1LGs			
	Samples subtyped	Loci investigated C. hominis C. parvum	C. hominis	C. parvum	Key findings	Reference
		MS5— Mallon				
		MS9—				
		Mallon				
		MSA				
		MSB				
		MSC				
		MSE				
		MSF				
		MSG				
		ISM				
		MSK				
		TP14				
Investigation of population C. he	C. hominis: 51 human cases	ML1 ML2	8	17 in humans	Most C. hominis isolates (Tanriverdi et al. 2008)	Ngouanesavanh
Port-au-Prince, Haiti		ZTINI			were of one MLG	
<i>C. p.</i> a	<i>C. parvum</i> : 42 human cases MS5— and 14 animal samples Ma	MS5— Mallon		10 in calves	From analysis of human isolates in France a basic	
	(7 calf; 7 goat)			5 in goats	clonal population struc-	
		Mallon)	ture was observed and in <i>C. hominis</i> basic or epi- demic clonal population structure. In Haiti, the low degree of polymor- phism suggested epi- demic clonality	

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Tanriverdi et al. (2008)	Caccio (2001) Leoni et al. (2007)	et al. (2007)	(continued)
For both parasite species, a quasi-complete phyloge- netic segregation was observed among the countries. Statistical tests, linkage disequilib- rium (LD), and eBURST analysis indicated both clonal and panmictic population structures. Gene flow was not suffi- cient to erase genetic divergence among geo- graphically separated populations, which basi- cally remained allopatric	Some alleles were found in different hosts while some were host or geo- graphically restricted Isolates of both <i>C. hominis</i> and <i>C. norrum</i> from sen-	and <i>C. parvum</i> from sep- arate waterborne outbreaks were geneti- cally homogeneous, suggesting preferential or point source transmission	
74 (not differentiated by host in the analysis)	6 25 human 5 animal	5 animal	
81	9		
1887 MS9— Mallon MSA MSB MSC MSC MSC MSC MSC MSC TP14	ML1 ML2 gp60	ML1 MS5— Mallon	
 C. <i>hominis</i>: 290 sporadic and outbreak human cases C. <i>parvum</i>: 227 human cases and animal isolates (calves, horse, goat) 	 59 C. parvum (9 HIV + ve humans, 29 calves, 11 kids, 10 lambs) C. hominis: 104 human cases 08 shoradic 	 cases (28 sporadic, cases (28 sporadic, 60 linked to 3 waterborne outbreaks) 16 from 8 familial outbreaks) <i>C. parvum:</i> 126 human (73 sporadic, 40 linked to 3 waterborne outbreaks, 13 from 6 familial outbreaks), 17 sporadic livestock samples (lambs or calves) 	
Investigation of global partitioning in <i>C</i> . <i>parvum</i> and <i>C</i> . <i>hominis</i> populations and com- parison of the popula- tion structures among isolates from Uganda, USA, England and Wales (including for- eign travellers), Israel, New Zealand, Serbia and Turkey	Area-specinc studies Investigation of genetic variability in northern, central and southern Italy Investigation of genetic variability in Fnoland	variability in England	

Table 3.5 (continued)						
		Number of MLGs identified	LGs			
Location	Samples subtyped	Loci investigated C. hominis C. parvum	C. hominis	C. parvum	Key findings	Reference
Investigation of genetic variability and risk factors in North West England and Wales	<i>C. hominis</i> : 118 sporadic human cases <i>C. parvum</i> : 63 sporadic human cases	gp60 ML1 ML2	6	31	Analysis with epidemiologic data found an association between reported contact with farm animals and individual <i>C. parvum</i> microsatellite alleles	Hunter et al. (2007)
Investigation of population structure in Aberdeenshire, Scotland	<i>C. hominis</i> : 71 sporadic human cases <i>C. parvum</i> : 64 sporadic human cases; 44 bovine cases	gp60 hsp70 ML1 MS5 Mallon MS9 Mallon MS12 TP14	٢	26 human 15 bovine	Overall cluster analysis identified three <i>C. parvum</i> and one <i>C. hominis</i> cluster. <i>C. hominis</i> isolates were essentially clonal. Human <i>C. parvum</i> isolates were in LD suggesting clonality, whereas bovine isolates were not, suggesting pannixia	Mallon et al. (2003a)
Investigation of population structure in Orkney and Thurso, Dumfriesshire, Scotland	<i>C. parvum:</i> 40 human cases, 190 bovine cases, 11 ovine cases	gp60 ML1 hsp70 MS12 MS5- Mallon MS9- Mallon TP14	7	36 human 24 bovine 6 ovine	Found no evidence to support Mallon (2003b) geographic or temporal geographic or temporal sub-structuring of the <i>C. parvum</i> populations, but host-substructuring was identified by cluster analysis, with human <i>C. parvum</i> isolates falling into two human-only groups and a human/animal group	Mallon (2003b)

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Morrison et al. (2008) C. <i>parvum</i> data from Mallon (2003b)	Gatei (2007)	Gatei (2008)	Peng (2003)	(continued)
Limited sub-structuring found within a small geographical area, but substantial sub-structuring over larger geographical distances	Four major groups identified. Gatei (2007) Significant intra- and inter-genic LD observed with minimum recombi- nation or expansion of limited subtypes, indica- tive of a mostly clonal population structure	Presence of homogeneous <i>C. hominis</i> and <i>C. parvum</i> populations which were distinct from isolates from other countries	LD analysis showed possible Peng (2003) recombination. Crypto- sporidiosis in the study area was largely caused by anthroponotic transmission	
95 (not differentiated by host in the analysis)	25	4	10	
I	I	7	1	
gp60 hsp70 ML1 MM5 MM19 MM19 MS5- Mallon MS9- Mallon MS12 TP14	Chrom3T CP47 CP56 GP60 HSP70 MSC6-7 Mucin1	CP47 DZ-HRGP HSP70 MSC6-7 RPGR		
297 <i>C. parvum</i> samples (human, cattle, sheep)	C. hominis: 47 paediatric patients	HIV + ve patients: 25 C. hominis 7 C. parvum	41 HIV + ve and –ve <i>C</i> . <i>hominis</i> patients (MLGs obtained for 18 only)	
Re-analysis of <i>C. parvum</i> cases from cases from Aberdeenshire, Orkney and Dumfriesshire, Scotland (Mallon et al. 2003a, b)	Investigation of population structure in Kolkata, India	Study of genetic diversity in Kingston, Jamaica	Investigation of population structure in Malawi	

Table 3.5 (continued)						
		Number of MLGs identified	S			
Location	Samples subtyped	Loci investigated C. hominis C. parvum	hominis C. pa	um	Key findings	Reference
Investigation of population structure in northern, central, and southern Italy	<i>C. parvum</i> : 9 human, 122 bovine, 21 ovine, 21 caprine <i>C. hominis</i> : 9 human	MS1 2 MS9 TP14 MM5 MM18 MM19 GP60	102 (9 human)	nan)	Predominantly clonal popu- lation structure, but also evidence that part of the diversity could be explained by genetic exchange	Drumo et al. (2012)
Investigation of population structure in Minnesota and Wisconsin, USA	212 <i>C. parvum</i> : 166 human, 46 cattle	TP14 - TP14 - CP47 MS5 MS5 MS9 MS06-7 MS06-7 DZ-HRGP GRH gp60	94 (90 hi	94 (90 human, 13 bovine)	Among the 94 MLGs identified, 60 were represented by a single isolate. Approximately 20 % of isolates belonged to MLT 2, a group that included both human and cattle isolates. Population analyses revealed a pre- dominantly panmictic population with no apparent geographic or host substructuring	Herges et al. (2012)
Outbreak investigations Various locations in Wales and England	<i>C. parvum:</i> 25 human cases 4 lambs 2 calves 1 effluent sample (also contained <i>C. ubiquitum</i>)	hsp70 – ITS-2 (SSCP) gp60 ML1 ML2	4		Single MLGs identified in each outbreak. Outbreak cases and suspected sources were linked	Chalmers et al. (2005)

Mattsson (2008)	Hunter et al. (2008)
MLGs were clustered according to pool use: one MLG was found among outdoor pool users and secondary cases, and another MLG from users of indoor pool	Clustering of one <i>C. parvum</i> Hunter MLG was observed (sta- et al tistically significant) in patients living in the area most dependent on one water supply
7	5
I	-
gp60 hsp70 TP14	gp60 ML1 ML2
Outbreak cases associated with a swimming pool	88 human samples from drinking waterborne outbreak:10 <i>C. hominis</i>78 <i>C. parvum</i>
Stockholm, Sweden	North West England

published data to identify the most informative markers as candidates for the development of a standardised multi-locus fragment size-based typing (MLFT) scheme for epidemiological purposes. They identified that in 31 MLFT studies, 55 markers had been used, of which 45 were applied to both *C. parvum* and *C. hominis*. Further analysis of 11 selected studies that provided sufficient raw data from three or more markers of >2 bp repeats, and a sampling frame containing at least 50 samples, revealed that some markers were present in all schemes at an average redundancy (i.e. markers which did not significantly enhance discrimination) of 40 % for *C. hominis* and 27 % for *C. parvum*. They ranked markers based on the most discriminatory combinations, which revealed two different sets of potentially most informative candidate markers, one for each species. It is likely, however, that wider application of whole genome sequencing will enable the identification of more informative markers.

Accurate analysis of microsatellite and minisatellite markers can be problematic due to the difficulties encountered by DNA polymerases during PCR amplification resulting in multiple fragment sizes evidenced by multiple "stutter" bands on electrophoresis gels or mixed sequence chromatograms (Diaz et al. 2012). Shorter microsatellites appear to be more problematic in this respect as well as being more technically challenging for size assignment by MLFT methods. Careful selection of marker and analysis method with validation with sequenced isolates is therefore essential. It should also be noted that sizes given by different MLFT methods are not necessarily equivalent.

Although sequence analysis is perhaps most desirable as it can potentially give the most accurate fragment size measurement and greater discrimination, if MLG is to be used during outbreak investigations fragment sizing provides more rapid assessment of relationship between isolates. Either way, there is a need for technical evaluation including typability (percentage of samples generating a complete multilocus type) and discriminatory power by direct fragment size analysis and analysed for correlation with epidemiological data in suitable sampling frames.

An alternative sequence based approach to analysis of micro- and minisatellites and the gp60 gene was reported by Bouzid et al. (2010a) who reported SNPs in ten slower mutating loci identified in published genomes. As more whole genome data becomes available it is likely that more such SNPs will be discovered, allowing rapid and more extensive analysis using array-based techniques making genomewide association studies possible for investigation of aspects of the parasite's biology e.g. virulence.

3.4 Strain Differences in Epidemiology and Outcome

More than 80 studies have been published investigating gp60 sequences in humanderived *C. parvum* and *C. hominis* isolates (Table 3.4). Broadly, these show some differences in distribution of gp60 families between studies, and it is clear that some sampled populations have greater diversity than others (Table 3.4). For example, *C. hominis* family Ib appears to predominate in most studies in Europe, North America and high income countries in Oceania. There are two common Ib subtypes: IbA10G2 which is especially prevalent in northern Europe, and IbA9G3 which is common in Australia, India and some countries in Africa, but more rare in northern Europe (Chalmers et al. 2008; Wielinga et al. 2008). However, one study in Western Australia found Id and Ig to be more common than Ib, especially in young children (Ng et al. 2010a). All *C. hominis* outbreaks in Europe, and nearly half of those in the USA have been caused by IbA10G2 (Xiao 2010). In Peru and China studies have revealed greater diversity within Ib than elsewhere (Cama et al. 2007; Feng et al. 2009; Peng et al. 2001).

In most high-income countries, *C. hominis* Id and Ia occur only occasionally and other gp60 families more rarely. There are some notable exceptions, in addition to the study in Western Australia mentioned above, another study in Australia reported C. *hominis* Id as the second most common after Ib (O'Brien et al. 2008). During the investigation of increased numbers of reported cases in 2007 in four states in the USA, Xiao et al. (2009) found that *C. hominis* Ia was most prevalent, and identified as rare subtype IaA28R4. This subtype had been found in cases linked to recreational water outbreaks in that year and in previous years, and the authors speculate that the cases may have been part of a larger, multistate outbreak for which typing was not widely undertaken (Xiao et al. 2009). In Sweden and Slovenia, Id is reported second to Ib, and is also common in reports from the Indian sub-continent, Peru, Kenya, Malawi and in one study in the US (Table 3.4). In a study of sporadic cases in the UK, having a non-IbA10G2 subtype was significantly linked to recent travel outside Europe (Chalmers et al. 2008).

There is greater diversity of *C. hominis* subtypes in less industrialised nations, and in studies in Peru these have found linkage between subtype and clinical outcomes that appear to vary with the particular host population under investigation, producing different spectra in children and HIV + adults. In HIV + patients in Lima, having Id was significantly associated with diarrhea in general and chronic diarrhea in particular, while infections with Ia and Ie were more likely asymptomatic (Cama et al. 2007). In a longitudinal cohort study of children in the same area of Lima, all *C. hominis* infections were associated with diarrhea, although Ib was significantly associated with diarrhea, nausea, vomiting and general malaise during both first and subsequent infections and thus appeared more virulent (Cama et al. 2008). In first episodes of infection, Ia was also very pathogenic being significantly associated with diarrhea, nausea and vomiting (Cama et al. 2008).

Among *C. parvum*, IIc is considered to be anthroponotic, having no defined animal host. It is widespread, especially where *C. hominis* is prevalent, like in Africa, Western Asia, Latin American and the Caribbean (Table 3.4), and the lack of a defined animal host for IIc supports its anthropomorphic transmission. Although *C. hominis* IbA10G2 has been found on rare occasions in animals, there is as yet no evidence for animals as a reservoir of human infection.

The other major gp60 families common in humans, IIa and IId, have been found in animals, especially ruminants, and this supports the zoonotic potential of *C. parvum* IIa and IId. One study in Spain found that family IIa is particularly associated with bovines, and IId with ovines and caprines (Quilez et al. 2008), although human outbreaks associated with sheep (linked to bottle feeding lambs) have occurred with IIa in the UK (Chalmers and Giles 2010). In most high income countries the anthroponotic *C. parvum* family IIc is generally rare, and *C. parvum* isolates are mainly IIa, although in Sweden IId is more common. In contrast, with a single exception, there have been few or no reports of IIa in sub-Saharan Africa, the Indian sub-continent, South and Central America and the Caribbean, where IIc is commonly found; coupled with the relatively high proportion of *C. hominis* cases in these countries, the importance of anthroponotic sources and transmission is highlighted here. The exception is one study in Ethiopia where IIa predominated and IIc was not found; zoonotic transmission here is supported by the predominance of *C. parvum* over *C. hominis* (Adamu et al. 2010), possibly reflecting the importance of cattle in that country. In North Africa, Western and South Eastern Asia, zoonotic transmission is also indicated by gp60 family distribution.

Any distribution analysis also needs to take into account the population investigated e.g. diarrhoea status, age, sex, HIV status, urban versus rural location, season, etc. Although inference about zoonotic potential can be made from prior knowledge of the host distribution of subtypes, further evidence for interpretation of the distribution of gp60 families and subtypes has been provided in some studies where humans were sampled alongside animals in the same area. For example, studies in Canada, Australia, the Netherlands and Egypt found a high proportion of human cases with the same gp60 subtypes as those found in livestock animals (Budu-Amoako et al. 2012; Helmy et al. 2013; Ng et al. 2012; Wielinga et al. 2008) indicating they may be a reservoir for infection. In one study in New Zealand, all foal, human, and bovine C. parvum isolates investigated had the gp60 IIaA18G3R1 subtype (Grinberg et al. 2008b). A high prevalence of C. parvum IIaA15G2R1 was found in both humans and cattle in the Netherlands (Wielinga et al. 2008), in 9/15 HIV + patients and all bovine, one of two sheep and all zoo ruminants tested in Portugal (Alves et al. 2006), but in only 2/9 Australian patients, 2/7 Australian cattle samples and all Canadian, European and US cattle sampled (O'Brien et al. 2008). Other Australian patients in that study had IIaA18G3R1, IIaA19G2R1, IIaA19G3R1, IIaA19G4R1, IIcA5G3a1. In fact, IIaA18G3R1 was found in farm dwellers and calves on the same farms in New South Wales (Ng et al. 2012). A greater diversity of subtypes was detected previously in human and cattle samples across the state (Ng et al. 2008). In Canada, a study on Prince Edward Island revealed that of three subtypes found in beef cattle,IIaA16G2RI, IIaA16G3RI and IIaA15G2RI, two, IIaA16G2RI and IIaA15G2RI, were also identified in the human isolates (Budu-Amoako et al. 2012). In Ontario, greater diversity was identified in both humans and dairy claves in the province of Ontario where three isolates from humans were the same subtypes as isolates from the calves (Trotz-Williams et al. 2006). In Iran, cattle were deemed to be a possible source of IIa subtypes (IIaA15G2R1, IIaA16G3R1), and one IId subtype (IId A15G1) in children, but there were more IId subtypes in children than cattle, the source of which was not known (Nazemalhosseini-Mojarad et al. 2011). Another study in Iran, also found that IId was common in children in Tehran, this time IIdA20G1a (Taghipour et al. 2011). In a study of bovines and humans in Egypt, both the bovine and the human infections were with IIaA15G1R1 and IIdA20G1, although the humans also had IIaA15G2R1 (Helmy et al. 2013). In England and Wales IIaA17G1R1 and IIaA15G2R1 predominate in cattle and lambs (Brook et al. 2009; Mueller-Doblies et al. 2008; Smith et al. 2010b) and in sporadic human *C. parvum* cases (Chalmers et al. 2011c). However, in Ireland, as in New South Wales Australia, IIaA18G3R1 predominates in both humans (Zintl et al. 2011) and cattle (Thompson et al. 2007), but is rare in England and Wales.

Data arising from an on-farm study in England and Wales, showing the proportion of human cases that had the same subtype as those found in suspected source animal contacts (Smith et al. 2010b), were used to estimate, from national surveillance data, the proportion of all laboratory confirmed cases that might have been acquired from direct contact with farm animals (Chalmers et al. 2011a). The proportion was 25 % for *C. parvum* cases, and 10 % of all reported *Cryptosporidium* cases.

Some studies have combined analytical epidemiology with analysis of gp60 families and subtypes. For example, in a study of *C. parvum* cases in the UK, cryptosporidiosis cases with IIa were more likely than IId to have visited a farm, or had contact with farm animals or with their faeces in the 2 weeks prior to illness (Chalmers et al. 2011c). Within gp60 family IIa, genotypes IIaA15G2R1 and IIaA17G1R1 predominated (22 cases each); nine other IIa genotypes accounted for 12 cases. The IId genotypes were mainly IIdA17G1R1 and IIdA18G1 (three each). Cases with IIaA17G1R1 were particularly linked to zoonotic exposures: visiting a farm or having farm animal contact in the 2 weeks prior to illness.

The analysis of a single locus is not sufficient to infer geographical diversity of subtypes. Global gp60 subtype distribution revealed limited microbiological evidence for host specificity (Jex and Gasser 2008). Although many gp60 subtypes have only been found in humans, many are rare findings and we do not know whether they have an animal host or not.

Some studies have used MLG to make inferences about the global population structures of *C. hominis* and *C. parvum* (Table 3.5). For example, Gatei et al. (2006) compared fragment sizing and sequence-based analysis of *C. hominis* subtypes at multiple loci, both including and excluding gp60. When gp60 was excluded from the analysis, all but three of 27 MLGs identified were found in samples from single countries. They concluded that geographical isolation has led to the emergence of distinct *C. hominis* subtypes. Targets other than gp60 were identified as suitable to discriminate geographically distinct subtypes, including the MSC6–7, DZHRGP, CP47 and HSP70 loci which showed genetic differentiation by region, and their inclusion in a multi-locus scheme was deemed to provide a superior subtyping tool for *C. hominis*.

A fragment sizing approach used by Tanriverdi et al. (2008), indicated that gene exchanges do not seem sufficient to erase genetic divergence among geographically separated populations, and that multilocus genotyping enables the tracking of *Cryptosporidium* isolates to their country of origin. They suggested that, rather than conforming to a strict paradigm of either clonal (deriving from a common ancestor without interbreeding with other types within the species) or panmictic (interbreeding freely within the species) species, their data are consistent with the

co-occurrence of both modes of propagation, with the relative contribution of each mode appearing to vary according to the prevailing ecological determinants of transmission. This seems to be important as apparently conflicting conclusions have been drawn from studies in different countries and different regions (Table 3.5). For example, in another study in a smaller geographical area, Scotland, lack of partitioning may be explained by the high frequency of cattle movements, typical in the UK, preventing the establishment of settled and epidemiologically isolated herds (Morrison et al. 2008).

It should also be recognised that population studies based on clinical isolates tend to be biased towards analysis of the most virulent strains and may not be representative of overall population structure (Gupta and Maiden 2001). Another limitation of these studies is that the presence of mixed MLGs is often not accounted for in analyses and the contribution of mixed infections to population diversity is not well understood.

Interestingly, in the interim since Gatei's work in 2006, although DZHRGP and CP47 featured in the preferred marker list generated by the analysis of Robinson and Chalmers (2012), MSC6-7 and hsp70 did not appear in the top half of the most informative rankings, other markers being deemed more useful.

Only a small number of publications have reported on epidemiological investigations applying MLG to both sporadic and outbreak cases. In outbreaks, MLG has been demonstrated to be useful in identifying related cases. In the UK, Hunter et al. (2008) demonstrated that although 17 different MLGs were detected among cases, clustering of one MLG was statistically significant for patients resident in the area most dependent on one water supply. In a swimming pool-related outbreak in Sweden, MLGs were clustered according to pool use: one MLG was found among outdoor pool users and secondary cases, and another MLG from users of an indoor pool (Mattsson et al. 2008).

Overall there is a need for the application of a standard multi-locus subtyping scheme applied to more host species and most especially in non-industrialised countries where the burden of illness and long-term sequelae from *Cryptosporidium* infection are greatest.

3.5 Application of Molecular Methods in Sporadic and Outbreak Related Cryptosporidiosis

3.5.1 Outbreak Detection, Investigation, Management, Control and Surveillance

Molecular typing and subtyping have proved valuable in the investigation of outbreaks of cryptosporidiosis, with subtyping of *C. parvum* and *C. hominis* allowing a degree of confidence that isolates can be recognised as an "outbreak strain". Early investigations indicated that outbreaks could be caused by *C. hominis* or *C. parvum*, and that drinking water and recreational water outbreaks could be

caused by either species (McLauchlin et al. 2000). However, the development of more specific and sensitive methods of species identification and subtyping has resulted in an ability to more sensitively examine outbreaks that are not from these sources.

3.5.2 Foodborne Outbreaks

Many outbreaks linked to food have been reported (Centers for Disease C, Prevention 1995, 1996, 1997; Djuretic et al. 1997; Millard et al. 1994; Robertson and Chalmers 2013), but the investigation of such sources has been limited because of the lack of systematic subtyping and reporting of strains to national surveillance (Chap. 2). Outbreaks in Sweden in Stockholm and Umea caused by C. parvum subtype IIdA24G1 and another one in Orebro by C. parvum IIdA20G1e illustrate the value in applying subtyping to the investigation of Cryptosporidium cases. C. parvum seems to predominate in foodborne outbreaks suggesting animal sources of contamination. However, some foodborne outbreaks have been linked to contamination by food handlers (Ethelberg et al. 2009; Harper et al. 2002; Lee and Greig 2010; Ponka et al. 2009; Ouiroz et al. 2000), where the local nature of the outbreak makes subtyping less essential as the route of transmission is clear and interventions exist to prevent future outbreaks (improvement of hygiene). It remains possible that foodborne cryptosporidiosis is more common than we currently recognise, particularly through salad items and soft fruits as has been common for Cyclospora infections (Chalmers et al. 2000b). A large Cryptosporidium outbreak across England and Scotland in 2012 was attributed epidemiologically to a bagged salad product, partly as a result of the identification of an outbreak strain based on gp60 subtyping of C. parvum (unpublished data). However, there is a need for routine subtyping of all clinical Cryptosporidium isolates using even more specific methods, along with these results feeding into national surveillance, so that other clusters of cases can be detected and links to sources of contamination investigated. This was particularly the case in the late summer of 2012 when there was a simultaneous increase in C. hominis infections in three EU countries the cause of which is not known (Fournet et al. 2012).

One of the difficulties in investigating foodborne cryptosporidiosis is that the implicated foods are generally short-shelf life products which will have been discarded before the cases of illness have been identified, leaving no relevant food samples for testing. Further, issues of sampling from large batches of products are raised where the distribution of the contamination may be uneven and missed by sampling procedures. There are as yet no standard methods for the detection of *Cryptosporidium* in food, although an ISO method for soft fruit and leafy green vegetables is in preparation. This does not include molecular tests but they could be incorporated in the process in a similar way to that applied to water, by either testing part of the pellet by immunomagnetic separation (IMS) or by scraping material from *Cryptosporidium*-positive microscopy slides (Ruecker et al. 2013). Although this procedure has been informative in investigations (see below) and gp60 subtyping has been applied, multi-locus methods have not, and careful

preparation will be needed to ensure that sufficient DNA is extracted to permit the sequencing of multiple markers.

3.5.3 Waterborne Outbreaks

Typing has been useful in the analysis and follow-up of waterborne outbreaks, some of which are mentioned above (see also Chap. 12). Microsatellite typing was used in the investigation of cases that may have been linked to a UK drinking water related outbreak and was useful in identifying cases most likely to have been involved (Hunter et al. 2008). A unique drinking water related outbreak of cryptosporidiosis in a city in England was linked to water contamination from a rabbit, the *Cryptosporidium* species, *C. cuniculus*, only identified by typing (Robinson and Chalmers 2010; Smith et al. 2010a) (Pulleston et al., submitted for publication). A state-wide outbreak of cryptosporidiosis throughout New South Wales, Australia used genetic characterization to identify *C. hominis* IbA10G2 subtype as the causative parasite acquired from swimming pools (Waldron et al. 2011b). A drinking water related outbreak in Wales was attributed to *C. hominis* gp60 subtype IbA10G2, which was also found in source waters serving the drinking water supply (Chalmers et al. 2010).

3.5.4 Source Tracking (Isolates from Water) and Source Attribution (Animal Sources of Human Infections)

Comparing isolates from water and host animals with those from humans can be used to track the levels of exposure to humans through source waters used for drinking water and to attribute human infection to particular source animals (Ruecker et al. 2012). For example, a rare gp60 subtype IIaA20G2R1 that caused an outbreak of cryptosporidiosis among lambs but not calves in a mixed sheep/ cattle farm in central Italy, was also found in the faeces of an 18 month old boy who lived on the farm and had been hospitalised due to acute gastrointestinal symptoms demonstrating transmission from lambs to humans (Caccio et al. 2013). This route of transmission is a common cause of outbreaks in the spring in the UK (Gormley et al. 2011), and typing has been applied in the investigation of farm-related outbreaks (Chalmers and Giles 2010).

Cryptosporidium has been one of the most successfully studied organisms with respect to source attribution. The common occurrence of *C. parvum* in agricultural animals and the almost complete absence of *C. hominis* has been key in focussing the attention of water authorities on the differential contributions of human and animal sources of contamination of drinking water (Nichols 2008). The identification of a variety of *Cryptosporidium* species that have a narrow host specificity has also been useful in ruling out some potential transmission routes, for example dogs and cats, as being common (Bowman and Lucio-Forster 2010). Comparing isolates from water

and host animals with those from humans can be used to track the levels of exposure to humans through source waters used for drinking water (Nichols et al. 2003, 2006a, 2010) and to attribute human infection to particular source animals (Ruecker et al. 2012). Some studies have identified most of the *Cryptosporidium* isolates to be not strains that commonly cause human disease (Yang et al. 2008).

The value of this work, in addition to identifying transmission pathways and sources, has been to extend our understanding of the host range of *Cryptosporidium* spp. present in the natural environment, and to reassure us that most of these isolates do not seem to be linked with human disease.

3.5.5 Outbreak Detection, Investigation, Management and Control

There have been numerous reviews and analyses of outbreaks of cryptosporidiosis, some general enteric ones and others exclusively due to *Cryptosporidium* (Baldursson and Karanis 2011; Barwick et al. 2000; Beaudeau et al. 2008; Chalmers 2012; Chalmers and Giles 2010; Craun et al. 2005; Curriero et al. 2001; Djuretic et al. 1997; Dziuban et al. 2006; Furtado et al. 1998; Glaberman et al. 2002; Hlavsa et al. 2011; Karanis et al. 2007; Kramer et al. 1996; Lee and Greig 2010; Lee et al. 2002; LeJeune and Davis 2004; Levine et al. 1990; Levy et al. 1998; McLauchlin et al. 2000; Nichols et al. 2006, 2009; Risebro et al. 2007; Rooney et al. 2004; Schuster et al. 2005; Semenza and Nichols 2007; Smith et al. 2006; Vojdani et al. 2008; Xiao 2008; Yoder et al. 2004, 2008). In 2008, Xiao and Morgan reviewed outbreaks where gp60 genotypes were identified (Xiao 2008). In reviewing the molecular epidemiology it is always useful to assess what using modern molecular methods has brought to the investigation and control of such outbreaks.

The approaches to outbreak investigation of protozoan infections have been reviewed recently (Pollock and Nichols 2012). The detection and investigation of outbreaks is crucially dependent on having good microbiological diagnostic regimens in place in primary diagnostic laboratories. In the absence of this only the largest of outbreaks will be detected. It is not generally possible for primary laboratories to type strains to species or sub-species level, and therefore having a reliable reference service, or a research centre of excellence is fundamental to providing the detail derived from subtyping. These reference services also interact with each other to provide a common nomenclature of species and types so that strain comparison can be understood internationally. Recent reviews have addressed taxonomy and nomenclature (Xiao et al. 2004). The typing is useful in real time to focus investigations on linked cases and eliminate background cases that are not part of the outbreak. There needs to be communication between public health teams and local and national governments, drinking water suppliers and regulators as well as swimming pool operators, in order that these groups understand the specific problems resulting from the chlorine resistance of *Cryptosporidium* oocysts,

so that when an outbreak is detected there is some agreement about the science. In particular, what testing is required, what results from water testing mean, what levels of oocyst contamination are acceptable, if any, what are the best approaches to intervention to limit the outbreak, who is in charge of the investigation, what approach should be used for the investigation and also so that the people involved know each other and therefore can have some prior experience of trusting each other.

Detection of a cluster of cases locally should result in a problem assessment group being established, upgraded to an outbreak control team should the evidence warrant this. Because cryptosporidiosis is commonly waterborne, cases usually present as a defined local cluster. However, outbreaks linked to overseas travel or to widely distributed food items, mean that outbreaks can arise that are not local. It is here, particularly, where speciation and subtyping are important in identifying an outbreak strain. For outbreaks where faecally contaminated water is the likely source there is a greater chance of a mixed outbreak caused by more than one strain, because the contamination may have come from several infected people or animals, compared to an outbreak caused by a water contaminated from a single individual. In the case of swimming pools there is good evidence for mixed outbreaks caused by C. parvum and C. hominis. The same is true for drinking water outbreaks. Outbreaks of foodborne illness where the infection derives from a single human case are thought to be predominantly derived from a single outbreak strain. The consequences of these difficulties is that in the absence of typing and subtyping some outbreaks can be difficult to investigate. In many of these cases older outbreaks reported there was a delay between when cases occured and when the results were available.

The control and prevention of outbreaks of cryptosporidiosis relies on the use of outbreak data to identify the sorts of combinations of events that can contribute to outbreaks. Some of these were outlined in a previous review (Nichols et al. 2006). For drinking water there are specific issues related to backwash water re-cycling, catchment management, water safety plans, source and treated water monitoring, animal management around treatment facilities, and emergency planning (Chaps. 12 and 13). For swimming pools the management approaches are established, but while reliance on the optimal operation of coagulation and filtration is essential, there is an increasing recognition that secondary disinfection using UV is necessary to deal with viable oocysts that break through conventional filtration at times.

3.6 Cryptosporidiosis in Immunocompromised People

Examinations of *Cryptosporidium* infections in people with some specific defects in immunity (Hunter and Nichols 2002) has been greatly assisted by molecular epidemiology (Wang et al. 2013). The occurrence of different *Cryptosporidium* species in immunocompetent and immunocompromised people was reviewed by Xiao and Morgan (2008). Both groups had infections with a range of species, and there do not appear to be *Cryptosporidium* species that are infectious only for any of

the immunocompromised groups, although infected patients with compromised immunity will excrete the parasite for extended periods.

3.7 Serological Markers

The detection of serum antibodies in response to *Cryptosporidium* infection has been useful in the examination of pathogenicity of individual *Cryptosporidium* species for humans (Chappell et al. 2006, 2011; DuPont et al. 1995; Moss et al. 1998). It has also been of use in comparing the incidence of cryptosporidiosis in different populations, particularly those served by different water supplies (Egorov et al. 2004; Frost et al. 2000a, b, 2001, 2003, 2005; Kozisek et al. 2008). For detailed molecular epidemiology however, the current tools for examining the response to infection will not clearly differentiate between different *Cryptosporidium* species, and are unlikely to be able in the future to differentiate between different genetic subtypes without specific cloned proteins screened in microarrays.

3.8 Conclusions and Future Developments

This chapter started by reviewing what past molecular epidemiology studies have shown us. The question that remains is what issues are still left to address. For us, one critical missing piece of the story at present is what outbreaks and sources of contamination and infection we are missing using current methods and surveillance algorithms. In particular, there seems to be the possibility of outbreaks occurring that are not being detected using current surveillance and/or typing and subtyping methods. When an outbreak strain is identified, it is commonly one that is also found in sporadic cases, and therefore being definitive about its role in the outbreak is difficult. This problem in common bacterial causes of diarrhoea like Salmonella is being addressed through whole genome sequencing of isolates to allow greater differentiation of strains (Singh et al. 2013; Zankari et al. 2013). For *Cryptosporidium* the problem is a bit harder because strains cannot readily be cultured to yield high numbers of parasite for genome sequencing. However, it is likely that an approach to whole genome sequencing larger numbers of strains using oocyst purification and whole genome amplification may yield approaches that are applicable to use in disease surveillance. The advantages of genome sequencing are that the dramatic and continuing reduction in the costs make this a feasible tool for use in molecular epidemiology and the finer differentiation of strain differences will lead to a much more proactive approach to the surveillance and control of cryptosporidiosis.

Using single sporozoite sequencing of the SSU rRNA gene from *C. andersoni* shows individual haploid sporozoites can contain two different copies of the gene (Ikarashi et al. 2013) (Le Blancq et al. 1997). More systematic whole genome sequencing will undoubtedly shed light on the recombination events that give rise to this and to distinctive features of *Cryptosporidium* genetics.

All diagnostic laboratories that test faecal samples for enteric and foodborne infections should be screening for *Cryptosporidium* using the recognised standard methods. While the infection is more common in young children, we also find that testing people of all ages is useful. Experience suggests that systematic subtyping of positive samples obtained through this approach can be enormously helpful in both understanding the causes of both cases and outbreaks, but also in response to outbreaks when they occur. Despite the widespread occurrence of large outbreaks across the World, it remains likely that there are more cases occurring in small outbreaks, most of which are not being detected. This is particularly the case for swimming pool related outbreaks where the exposed population may not be as large as occurs when a water catchment is contaminated. Food related outbreaks resulting from salad or fruit with a wide distribution will probably not be picked up by traditional epidemiological methods and we think that there is still a need for more discriminatory genetic methods to separate out strains associated with outbreaks from background sporadic cases.

The mapping (Piper et al. 1998) and sequencing of the complete genomes (Abrahamsen 2001; Abrahamsen et al. 2004; Liu et al. 1999; Widmer and Sullivan 2012; Xu et al. 2004) of three species of Cryptosporidium and an anthroponotic strain of C. parvum (Widmer et al. 2012) has lead to an increased understanding of the biochemistry and genetics of this genus (Chap. 6). Approaches to amplifying Cryptosporidium whole genome fragments have been examined and the commercial kits available have been compared (Bouzid et al. 2010b). Cryptosporidium genes can be examined through a Library of Apicomplexan Metabolic Pathways at http://www. llamp.net (Shanmugasundram et al. 2013), the Cryptosporidium Genomics Resource http://cryptodb.org/cryptodb/ (Heiges et al. 2006), the EPICDB relational database that organizes and displays experimental, high throughput proteomics data for T. gondii and C. parvum http://toro.aecom.yu.edu/cgi-bin/biodefense/main.cgi (Madrid-Aliste et al. 2009) and an extended database of protein domains for several eukaryotic pathogens http://www.atgc-montpellier.fr/EuPathDomains/ (Ghouila et al. 2011). However, there are still a considerable number of genes that are not annotated and assigned to a function, and there remains a strong need for additional work on individual proteins (Rider and Zhu 2010). Comparison of zoonotic and anthroponotic strains of C. parvum and C. hominis suggest these genes may influence the host range (Widmer et al. 2012). However, more gene sequences are needed for variants within these species and of other species, both human pathogenic and non pathogenic for comparison and further investigation.

Rather than an approach that passively collects *Cryptosporidium* surveillance data and then conducts questionnaires on cases that appear to be part of a cluster, the use of enhanced surveillance, where a questionnaire is administered to all patients, allows epidemiological, demographic and risk information to be collected systematically. Such an approach, conducted over an extended time period, can be useful in conducting ad-hoc case-case comparisons across years. Such an approach proved useful in the investigation of a multi-region outbreak in 2012 (unpublished) that early case-case examination suggested the outbreak might be linked to salad items

from a national retailer and which was subsequently shown to be from this source by a national case–control study.

Other issues that remain to be definitively answered follow. Does the rare occurrence of some species in humans reflect greater susceptibility in some human phenotypes, some immunocompromised people or some age groups? Is the detection of oocysts in the faeces always correlated with human infection and disease, and can some species infect but not produce oocysts in humans? Are any oocyst detections in humans a result of the consumption of heavily contaminated food with species that are not infectious to humans?

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Chapter 4 Cryptosporidiosis in Farmed Animals

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Abstract Cryptosporidiosis was first identified as a disease of veterinary, rather than human medical, importance, and infection of farmed animals with different species of *Cryptosporidium* continues to be of veterinary clinical concern. This chapter provides insights into *Cryptosporidium* infection in a range of farmed animals – cattle, sheep, goats, pigs, cervids, camelids, rabbits, water buffalo and poultry – presenting not only an updated overview of the infection in these animals, but also information on clinical disease, infection dynamics and zoonotic potential. Although extensive data have been accrued on, for example, *Cryptosporidium parvum* infection in calves, and calf cryptosporidiosis continues to be a major veterinary concern especially in temperate regions, there remains a paucity of data for other farmed animals, despite *Cryptosporidium* infection causing

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significant clinical disease and also, for some species, with the potential for transmission of infection to people, either directly or indirectly.

4.1 Introduction

4.1.1 Species of Cryptosporidium Relevant to Different Farmed Animals: Overview

Farmed animals, also commonly referred to as livestock or domesticated animals, are those animals that are reared in an agricultural setting in order to produce various commodities - usually food (meat, organs, eggs, dairy products), and/or hair or wool. In some settings farmed animals are also used to supply labour, and the manure of domesticated animals is often used as fertilizer. Animals were probably first farmed, that is their breeding and living conditions controlled by their human owners, around 7000-8000 BC during the first transitions from huntergatherer lifestyles to more settled agricultural living. The physiologies, behaviours, lifecycles of farmed animals generally differ quite substantially from those characteristics of the equivalent wild animals, and this difference impacts the interactions of these farmed animals with their parasites. Farmed animals are exposed to different stresses than wild animals, are kept at different densities, and their lifecycles regulated to such an extent that a parasite-host interaction in a farmed animal may differ significantly from that in a wild animal. Additionally, for infections that are of significant clinical importance, farmers may implement control measures (including treatment or prophylaxis) that alter the infection dynamics. With respect to *Cryptosporidium* infection, for which a satisfactory chemotherapeutic cure or prophylaxis is not yet available, different species infect different species of farmed animal, and may or may not be of clinical relevance.

Table 4.1 provides an overview of the farmed animals included in this chapter, the species of *Cryptosporidium* to which they are susceptible and brief notes on the clinical relevance. Greater details are provided in the appropriate chapter sections. Various categories of animals that are 'farmed', including mink, foxes, guinea pigs etc. are not included, largely because of a lack of information on *Cryptosporidium* in these animals in the domesticated setting. Additionally, farmed fish are not included in this chapter.

4.1.2 Relevance of Cryptosporidiosis in Farmed Animals to Human Infections

While cryptosporidiosis in farmed animals is of veterinary relevance, resulting in clinical morbidity, mortality, and associated production losses, the zoonotic nature

	Species of	
Farmed animal	Cryptosporidium	
Bovines, including cattle (<i>Bos taurus</i> and <i>B. indicus</i>),	C. parvum ^a	Common in pre-weaned calves – acute onset diarrhoea. Intestinal location
banteng (Bos javanicus), gayal (Bos frontalis),	C. bovis	Common in post-weaned calves – less pathogenic than <i>C. parvum</i>
water buffalo (<i>Bubalus bubalis</i>), and yaks (<i>Bos grunniens</i>)	C. andersoni	Older post-weaned calves, yearlings and adults- some failure to thrive. Infects the gastric glands of the abomasum
	C. ryanae	Common in post-weaned calves
	bovines. These	species has been reported from cattle and other e seem to be unusual and are apparently cal significance
Small ruminants, including	<i>C. parvum</i> ^a	Relatively common in pre-weaned lambs,
sheep (Ovis aries)	C. par van	associated with diarrhoea
and goats (Capra aegagrus hircus)	C. xiaoi	Common in older lambs and sheep, often apparently asymptomatic
	C. ubiquitum ^a	Common in older lambs and sheep, often apparently asymptomatic
Pigs (Sus scrofa domesticus)	C. parvum ^a	Less common than in bovines and small ruminants; diarrhoea and vomiting
	C. suis	Relatively common, mild symptoms
	C. scrofarum	Relatively common, mild symptoms
Deer (cervids), including red deer (<i>Cervus elaphus</i>) fallow deer (<i>Dama dama</i>), elk/wapiti (<i>Cervus</i> <i>canadensis</i>), white-tailed deer (<i>Odocoileus</i> <i>virginianus</i>), and reindeer (<i>Rangifer tarandus</i>)	C. parvum ^a C. ubiquitum ^a	Information on species detected amongst farmed deer is lacking; diarrhoea in young calves, possibly severe, but can also be asymptomatic
Camelids, including dromedaries (<i>Camelus</i> <i>dromedarius</i>), llama (<i>Lama glama</i>), and alpaca (<i>Lama pacos</i>)	C. parvum ^a	Relatively little information on species that are infectious to camelids; diarrhoeal disease, particularly in young alpaca (crias)
Rabbits (Oryctolagus cuniculus)	C. cuniculus ^a	Clinical symptoms in rabbits are apparently mild or lacking
Poultry, including chickens (Gallus gallus domesticus), ducks (Anas platyrhynchos), turkeys (Meleagris gallopavo),	C. meleagridis ^a	Appears to have a wide host range, including farmed poultry (and mammals). Mostly infects the intestines and has been associated with generally mild clinical symptoms
geese (Anser anser domesticus), ostriches (Struthio camelus), pigeons (Columba livia domestica) etc.	C. baileyi ^a	A wide avian host range reported, including various farmed poultry species. Detected in many different anatomical sites including digestive tract, respiratory tract, and urinary tract. Has been associated with high morbidity and mortality

Table 4.1 Overview of farmed animals and major relevant species of Cryptosporidium

(continued)

Farmed animal	Species of Cryptosporidium Clinical notes
	C. galli May affect a wide range of avian species, including farmed poultry. Infects the proventriculus, and has been associated with acute diarrhoeal disease
	Various host species-associated genotypes have been described from different poultry, including ducks and geese. Their clinical importance and host specificity is generally not known

^aNoted for zoonotic potential

of various *Cryptosporidium* species (see Table 4.1) means that public health may also be affected by infections in farmed animals. Infection may be direct, from animal to human, or indirect, via a transmission vehicle. A large number of small outbreaks associated with C. parvum in calves and in veterinarians or veterinary students that have been exposed to calf faeces are documented in the literature (e.g. Grinberg et al. 2011; Gait et al. 2008; Robertson et al. 2006). In addition, a number of outbreaks have been documented associated with children visiting 'petting farms' or similar venues, where interaction with young animals such as lambs or calves is encouraged. Less commonly, transmission between animals such as camels or alpacas and their carers has also been reported. Drinking water, and less often food, has been associated with transmission of Cryptosporidium infection from animals to human populations, with C. parvum from grazing cattle contaminating water supplies particularly implicated. The high densities of farmed animals in water catchment areas mean that implementation of catchment control measures, including preventing defecation into water courses, may have a significant effect on minimising the risk from this potential transmission pathway.

4.2 Cryptosporidium Infection in Bovines

There are various species of farmed bovines, with cattle and zebu (*Bos taurus* and *Bos indicus*, respectively) amongst the most important livestock worldwide. Both provide meat, milk and other dairy products, and are also used as draught animals, with an estimated 1.5 billion head globally (3 cattle for every 14 people). A comprehensive overview of *Cryptosporidium* infection in cattle has been published by Santín and Trout (2008) and the information presented here is largely an update built on this solid basis. Domesticated water buffalo (*Bubalus bubalis*) is also important domestic livestock in the bovinae sub-family. Domesticated buffalo consist of swamp buffalo and river buffalo. In 2011, the world population of buffalo was estimated to approximately 195 million animals, of which 97 % were in Asia (FAOSTAT 2012).

4.2.1 Occurrence (Prevalence)

4.2.1.1 Cattle

The first report on bovine cryptosporidiosis was published in 1971, when parasites were identified in an 8-month-old heifer with chronic diarrhoea (Panciera et al. 1971). Since then, Cryptosporidium infection in cattle has been documented in most countries worldwide. Four major Cryptosporidium species infect cattle: C. parvum, C. bovis, C. ryanae and C. andersoni (Table 4.2; Fayer et al. 2007; Feng et al. 2007; Santín et al. 2004; Langkjaer et al. 2007). Cryptosporidium parvum has a broad host range and apparently has the ability to infect most mammals, including humans and cattle. In contrast, the other three species have almost exclusively been found in cattle. In addition to these four common species, sporadic natural infections with C. felis, C. hominis, C. scrofarum, C. serpentis, C. suis and C. suis-like genotype have been detected in cattle (Bornay-Llinares et al. 1999; Geurden et al. 2006; Langkjaer et al. 2007; Santín et al. 2004; Smith et al. 2005; Chen and Huang 2012). The extent to which these findings reflect true infections or accidental carriage, i.e. ingested oocysts that pass intact through the gastrointestinal tract, remains to be clarified. Cattle have also been experimentally infected with C. canis, but natural infection has not been reported (Fayer et al. 2001).

Many reports on *Cryptosporidium* in cattle in different countries and settings have been published over the years, showing that *Cryptosporidium* spp. infections are common worldwide. Both dairy and beef cattle are infected and the prevalence estimates vary considerably among studies. Reported herd level prevalences range from 0 to 100 % (Olson et al. 1997; Chang'a et al. 2011; Maddox-Hyttel et al. 2006; Santín et al. 2004). Infected animals have been reported from all age groups but infection is most common in preweaned calves. When calves up to 2 months of age have been investigated in point prevalence surveys, 5–93 % of the calves shed oocysts (Table 4.2; Maddox-Hyttel et al. 2006; Santín et al. 2004; Uga et al. 2000). Longitudinal studies performed in infected dairy herds showed that all calves in such herds shed oocysts at some time during their first months of life (O'Handley et al. 1999; Santín et al. 2008).

The overall picture is that there is an age-related pattern in the species distribution. *C. parvum* is mostly found in preweaned, monogastric calves up to 2 months of age where it is often the most prevalent species, responsible for more than 80 % of *Cryptosporidium* infections (Brook et al. 2009; Fayer et al. 2007; Plutzer and Karanis 2007; Santín et al. 2004, 2008; Trotz-Williams et al. 2006). In some areas, however, *C. bovis* is the dominating species found in preweaned calves (Budu-Amoako et al. 2012a; Wang et al. 2011a; Silverlås et al. 2010b). The prevalence of *C. parvum* is considerably lower in older calves and young stock, and there are few reports of *C. parvum* infection in adult cows (Fayer et al. 2007; Langkjaer et al. 2007; Silverlås et al. 2010b; Castro-Hermida et al. 2011a; Khan et al. 2010; Ondrácková et al. 2009; Muhid et al. 2011; Budu-Amoako et al. 2012a, b).

Table 4.2 Locat	ion, age, and prevale	Table 4.2 Location, age, and prevalence of <i>Cryptosporidium</i> spp. in cattle in some recent studies ^a	p. in cattle in	some recent studies ^a		
Location	Age	No. of animals/farms or locations	Prevalence	Microscopy (M), ELISA (E), Molecular (Mo) or Other (O)	Molecular identification/ species or genotype (% investigated)	Reference
Australia, NSW	Calves	196/20 herds	74 %	Mo	18S rRNA and GP60/	Ng et al. 2012
					C. parvum 59 % C. bovis 20 %	
					C. ryanae 10 % Mixed infections 10 %	
					Not identified 1 %	
Brazil	≤30 days	196/dairy herds	$11 \ \%$	Mo	18S rRNA and GP60/	Meireles et al. 2011
					C. parvum 33 %	
					C. bovis 5 %	
					C. ryanae 10 %	
					C. andersoni 10 %	
					Not identified 42 %	
Canada	<2 months	752/20 dairy herds	$17 \ \%$	Μ	18S rRNA/	Budu-Amoako
	2–6 months		14 %		C. parvum 5 %	et al. 2012a
	>6 months		15 ~%		C. bovis 51 %	
					C. ryanae 17 %	
					C. andersoni 27 %	
Canada	≤ 6 months	739/20 beef herds	$18 \ \%$	М	18S rRNA and HSP70/	Budu-Amoako
	>6 months		15 %		C. parvum 24 %	et al. 2012b
					C. bovis 20 %	
					C. ryanae 7 %	
					C. andersoni 49 %	
Czech Republic	20–60 days	750/24 dairy herds	21 %	М	18S rRNA RFLP/	Kváč et al. 2011
					C. parvum 86 %	
					C. bovis 2 %	
					C. andersoni 13 %	

China	0->48 months	2,056/14 dairy herds	19 %	W	18s rRNA/ C. parvum 48 % C. bovis 16 % C. andersoni 29 % C. hominix 6 %	Chen and Huang 2012
China	0-8 weeks	801/8 herds	21 %	Μ	C. serpentis 1 % C. serpentis 1 % Blas rRNA and GP60/ C. parvum 31 % C. bovis 38 % C. ryanae 11 % C. andersoni 7 %	Wang et al. 2011b
England and Wales	\leq 3 months	229 dairy or beef calves/diagnostic lab	45 %	W	MIXED INFECTIONS 12 % 18s rRNA/ C. parvum 91 % C. bovis 2 % MA: idanifad 7 %	Featherstone et al. 2010a
England and Wales	Preweaned Immature Adult	116/11 herds connected with human cryptosporidiosis	81 % 58 % 19 %	M	C. parvum 77 % C. parvum 77 % C. bovis 5 % C. andersoni 16 % Not identified 7 %	Smith et al. 2010
Egypt	<6 weeks	96/2 dairy herds	30 %	М	C anderconi 7 %	Amer et al. 2010
Egypt	1 day–3 months >3 months–1 year >1–2 years >2 years	593	30 % 13 % 13 % 5 %	0	C. name som P. 18 18S rRNA and GP60/ C. parvum 65 % C. hovis 4 % C. ryanae 14 % Mixed infections17%	Helmy et al. 2013 (continued)

Location	Age	No. of animals/farms or locations	Prevalence	Microscopy (M), ELISA (E), Molecular (Mo) or Other (O)	Molecular identification/ species or genotype (% investigated)	Reference
Hungary	Preweaned calves	79 diarrhoeic/52 herds	49 %	М	18S rRNA and GP60/ C. parvum 95 % C. rvanae 5 %	Plutzer and Karanis 2007
India	<3 months	461/various	16 %	Μ	18s sRNA/ C. parvun 100 %	Maurya et al. 2013
India	0–2 months 3–12 months >12 months	180/2 dairy herds	20 % 14 % 4 %	M	18S rRNA/ C. parvum 29 % C. bovis 38 %	Khan et al. 2010
					C. ryanae 14 % C. andersoni 14 % C. suis-like 5 %	
İran	1-20 weeks	272/15 dairy herds	19 %	Μ	18S rRNA/ C. parvum 73 % C. bovis 8 % C. andersoni 18 % Atypical isolates 2 %	Keshavarz et al. 2009
Italy	0 day-<12 months	0 day-<12 months 2,024/248 dairy and beef herds	8 %	ELISA, M	COWP and GP60/ C. parvum 100 %	Duranti et al. 2009
Japan	3-48 days	80 diarrhoeic/different herds	75 %	Мо	18S rRNA/ C. parvum 53 % C. bovis 2 % Not identified 45 %	Karanis et al. 2010
Malaysia	1 day- \leq 4.5 months 250/16 herds >4.5-12 months	250/16 herds	31 % 23 %	Мо	18S rRNA and GP60/ C. parvum 17 % C. bovis 25 % C. ryanae 15 %	Muhid et al. 2011

Table 4.2 (continued)

Maikai et al. 2011	Thompson et al. 2007	Imre et al. 2011	Castro-Hermida et al. 2011a	Díaz et al. 2010a	Silverlås et al. 2009b; Silverlås et al. 2010b	Santín et al. 2004 (continued)
C. andersoni 20 % Mixed infections 6 % Not identified: 17 % 188 rRNA/ C. bovis 45 % C. ryanae 26 % C. andersoni 16 % Mixed infections 13 %	18S rRNA/ C. parvum 95 % C. bovis 4 % C. ryanae 1 %	18S rRNA and GP60/ <i>C. parvum</i> 100 %	18S rRNA/ C. parvum 56 % C. andersoni 23 % Not identified 21 %	18S rRNA and GP60 <i>C. parvum</i> 100 %	18S rRNA and GP60/ C. parvum 14 % C. bovis 75 % C. ryanae 9 % C. andersoni 2 %	18S rRNA/ C. parvum 50 % C. bovis 28 % C. ryanae 16 % C. andersoni 6 %
No	W	М	W	Μ	X	W
16 %	37 %	25 %	$\begin{array}{c} 61 \ \% \\ 15 \ \% \\ 8 \ \% \end{array}$	49 %	52 % 29 % 6 %	50 % 20 %
194/20 herds	779 diarrhoeic/ diagnostic lab	258 diarrhoeic/9 dairy herds	649/not specified	61 diarrhoeic/27 herds	1,202/50 dairy herds	971/15 dairy herds
2–365 days	<1 month	1–30 days	Neonatal Heifers Cows	≤21 days	≤2 months 4-12 months Cows	5 days-2 months 3-11 months
Nigeria	Northern Ireland <1 month	Romania	Spain	Spain	Sweden	USA 7 states

		No. of animals/farms		Microscopy (M), ELISA Molecular identification/ (F) Molecular (Mo) or suboies or genetitine (%)	Molecular identification/	
Location	Age	or locations	Prevalence Other (O)	Other (O)	investigated)	Reference
USA 7 states	12-24 months	571/14 dairy herds	12 %	Mo	18S rRNA/	Fayer et al. 2006
					C. parvum 6 %	
					C. bovis 35 %	
					C. ryanae 15 %	
					C. andersoni 43 %	
					C. suis 1 %	
USA 20 states	6–18 months	819/49 beef herds	20 %	Mo	18S rRNA/	Fayer et al. 2010
					C. bovis 23 %	
					C. ryanae 9 %	
					C. andersoni 68 %	
^a See e.g. Santín and Trout		(2008) for a review of older studies				

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^a See e.g.

Table 4.2 (continued)

In older calves and young stock, *C. bovis* and *C. ryanae* are the most commonly found species (Santín et al. 2008; Fayer et al. 2007; Muhid et al. 2011; Langkjaer et al. 2007; Silverlås et al. 2010b). *C. andersoni* is mainly found in young stock and adult cattle (Enemark et al. 2002; Wade et al. 2000; Fayer et al. 2007; Ralston et al. 2003). Older studies, based on microscopy alone, overestimated the *C. parvum* prevalence in weaned animals because the similarity in oocyst size makes it impossible to differentiate between *C. parvum* (~ $5.0 \times 4.5 \mu$ m), *C. bovis* (~ $4.9 \times 4.6 \mu$ m) and *C. ryanae* (~ $3.7 \times 3.2 \mu$ m).

Molecular studies have revealed different genetic subtypes within the C. parvum and C. hominis species and DNA sequence analysis of the 60 kDa glycoprotein gene is commonly used to further characterize the isolates. A number of C. parvum GP60 subtype families, designated IIa-IIo, have been described. Of these, IIc and IIe are considered anthroponotic, whereas IIa and IId are commonly found in both humans and animals. The other subtype families are uncommon and their zoonotic potential has not been determined. In cattle, C. parvum of the IIa subtype family is especially common. In addition to IIa, IId and occasionally III (sometimes named IIj) subtypes are found (Table 4.3; Xiao 2010). Whether there is a difference in pathogenicity between subtypes is currently unknown, as most genotyping studies to date have focused on herds with a history of calf diarrhoea. It has been suggested that herd management strategies affect subtype distribution. Studies from areas with closed herd management (limited animal movements between herds) have shown a high number of subtypes in the calf population, but only one subtype in each herd (Brook et al. 2009; Mišic and Abe 2007; Soba and Logar 2008; Silverlås et al. 2013). It has also been shown that a unique GP60 subtype can persist over time in a closed dairy herd (Björkman and Mattsson 2006). In contrast, only a few subtypes have been identified in areas with more animal movements between herds, but several subtypes could be present in a herd (Brook et al. 2009; Peng et al. 2003; Trotz-Williams et al. 2006).

4.2.1.2 Water Buffalo

The information on the distribution of *Cryptosporidium* infection in water buffalo is rather fragmentary. Reported prevalences vary between 3 % and 38 % (Table 4.4). An association between prevalence of infection and age of the animals, with the highest prevalence in young calves, has been found (Helmy et al. 2013; Maurya et al. 2013; Nasir et al. 2009; Bhat et al. 2012; Díaz de Ramírez et al. 2012).

The first report on *Cryptosporidium* species identification in water buffalo was published in 2005, in which Gómez-Couso et al. (2005) used molecular tools to characterize *Cryptosporidium* oocysts from an asymptomatic neonatal calf in a dairy buffalo farm in Spain. Sequence analysis of a fragment of the oocyst wall protein (COWP) gene revealed that the isolate was closely related to the *Cryptosporidium* 'pig' genotype. A few years later, *C. parvum* was identified in water buffalo calves from Italy (Cacciò et al. 2007) and today *C. ryanae*, *C. bovis* and

Table 4.3 Cryptospor.	idium parvum	Table 4.3 Cryptosporidium parvum GP60 subtypes in cattle in some recent studies ^a		
Country	Number subtyped	IIa subtypes ^b	IId subtypes ^b	Reference
Australia	36	IIaA16G3R1 (1), IIaA17G4R1 (4), IIaA18G3R1 (29), IIaA20G3R1 (2)	77	Waldron et al. 2011
Australia	84	IIaA18G3R1 (57), IIaA19G3R1 (11), IIaA17G2R1 (7), IIaA19G2R1 (6), IIaA16G3R1 (2), IIaA20G3R1 (1)		Ng et al. 2012
Brazil	7	IIaA15G2R1 (7)		Meireles et al. 2011
China	67		IIdA19G1 (54)	Wang et al. 2011b
Czech Republic	1	IIaA16G1R1 (1)		Ondráčková et al. 2009
Czech Republic	131	IIaA16G1R1 (56), IIaA15G2R1 (48), IIaA22G2R1 (12), IIaA15G1R1 (12), IIA18G1R1 (3)		Kváč et al. 2011
England and Wales	13	IIaA17G1R1(10), IIaA15G2R1 (3)		Smith et al. 2010
Egypt	24	IIaA15G2R1 (1)	IIdA20G1 (23)	Amer et al. 2010
Egypt	71	IIaA15G1R1 (16)	IIdA20G1 (54), IIdA19G1 (1)	Helmy et al. 2013
Hungary	21	IIaA16G1R1 (15), IIaA17G1R1 (3), IIaA18G1R1 (1)	ПdA19G1(1), ПdA22G1 (1)	Plutzer and Karanis 2007
Iran	25	IIaA15G2R1 (22), IIaA16G3R1 (1)	IIdA15G1 (2)	Nazemalhosseini-Mojarad et al. 2011
Italy	62	IIaA15G2R1 (34), IIaA18G2R1 (10), IIaA17G2R1 (9), IIa A14 (5), IIaA13 (4)		Duranti et al. 2009
Malaysia	8	IIaA17G2R1 (2), IIaA18G3R1 (1)	IIdA15G1 (5)	Muhid et al. 2011
Romania Spain	13 27	IIaA15G2R1 (8), IIaA16G1R1 (5) IIaA15G2R1 (26), IIaA13G1R1 (1)		Imre et al. 2011 Díaz et al. 2010a
Sweden	13	IIaA15G1R1 (2), IIaA18G1R1 (2), IIaA21G1R1 (2), IIaA16G1R1 (1)	IIdA20G1 (2), IIdA23G1 (2) IIdA16G1 (1), IIdA22G1 (1)	Silverlås et al. 2010b

A17GIR1c (17), IIdA20G1e (14), IIdA16G1b (4), Silverlås et al. 2013 IIaA17GIR1 (6), IIdA22G1 (4), IIdA24G1c (4), Silverlås et al. 2013 IIaA22GIR1 (7), IIdA22G1c (3), IIdA24G1c (4), IIdA22G1 (3), IaA15G2R1 (4), IIdA22G1c (3), IIdA23G1 (3), IIdA17G1d (2), IIdA19G1 (2), IaA15G1R1 (3), IIdA26G1b (2), IIdA19G1 (2), IIAA13G1R1 (3), IIdA26G1b (2) IIdA16G1 (2), IIAA13G1R1 (1), IIAA17R1 (1), IIAA17R1 (1),	llaA23GIR1 (1) ies ype
IIAA16GIR1 (58), IIAA17GIR1c (17), IIAA21GIR1 (12), IIAA17GIR1 (6), IIAA18GIR1c (7), IIAA22GIR1 (7), IIAA20GIR1 (5), IIAA15G2R1 (4), IIAA13GIR1 (3), IIAA15G1R1 (3), IIAA14R1 (2), IIAA16GIR1b (2), IIAA14GIR1b (1), IIAA17R1 (1), IIAA14GIR1b (1), IIAA17R1 (1),	IIaA2IGIR1b (1), IIaA23GIR1 (1) v of results of older studies of samples with the subtype
en 171	See Xiao et al. (2010) for a summary of results of older studies Numbers in parentheses are number of samples with the subtype
Sweden	^a See

Table 4.4	Table 4.4 Location, age, and prev	valence of Cryptos	sporidium spp. an	and prevalence of Cryptosporidium spp. and C. parvum subtypes in water buffalo ^a	in water buffalo ^a		
Location	Symptoms	Host age	Study design	Prevalence % positive (no. positive/no. examined)	Diagnostic technique	Molecular analyses gene/species or subtype ^b	Reference
Egypt	Diarrhoeic and non-diarrhoeic	<3 months	Cross-sectional	Individual 14 % (65/458) Herd 55	Modified Ziehl- Neelsen (mZN)		El-Khodery and Osman 2008
Egypt	Diarrhoeic and non-diarrhoeic	Not stated	Cross-sectional	22 % (16/71)	mZN and Sheather's flotation followed by mZN		Shoukry et al. 2009
Egypt	Not stated	1 week -4 months and adults	Cross-sectional	Calves: 10 % (17/179) Adults: 0 Herds: 71 % (5/7)	Sheather's flotation followed by mZN	18S rRNA/ C. parvum (7) C. ryanae (10) GP60/ IIdA20G1 (5) IIAA15G1R1 (2)	Amer et al. 2013
Egypt	Not stated for calves, diarrhoeic adults	Calves and adults	Cross-sectional	Overall: 20 %. (43/211) 1 day–3 months: 40 % (34/85) <3 months–1 year: 11 % (6/56) >1 year: 4 % (3/70)	Antibody based copro-antigen test	18S rRNA/ C. parvum (30) C. bovis (2) C. ryanae (3) Mixed infections (10) GP60/ IIdA20G1 (28) IIAA15G1R1 (4)	Helmy et al. 2013
India	Not stated	Not stated	Cross-sectional 17 % (10/60)	17 % (10/60)	Kinyon acid fast stain		Dubey et al. 1992

Mohanty and Panda 2012	Bhat et al. 2012	Maurya et al. 2013	Feng et al. 2012	Nasir et al. 2009	(continued)
		18S rRNA/ C. parvum (16)	18S rRNA/ C. ryanae (16) Unique variant		
mZN and formol- ether sedimentation followed by mZN	NZM	mZN or Sheather's flotation followed by mZN	Water-ethyl sedimentation followed by mZN	Sheather's flotation followed by mZN	
25 % (76/305)	Overall prevalence: 38 % (62/162) 1-2 months: 49 % (18/37) 2-3 months: 34 % (13/38) 3-4 months: 20 % (76/305) 4-5 months: 6 %	24 % (64/264)	37 % (30/81)	Overall prevalence: 24 % (60/250) 1-30 days: 42 % (29/69) 1-3 months: 26 % (17/66) 4-8 months: 15 % (9/60) 9 months->1 year: 9 % (5/55)	
Cross-sectional 25 % (76/305)	Cross-sectional	Cross-sectional 24 % (64/264)	Cross-sectional	l day->1 year Cross sectional	
Not stated	<5 months	<3 months	2-7 months	1 day->1 year	
Diarrhoeic and non-diarrhoeic	Diarrhoeic and non-diarrhoeic	Diarrhoeic and non-diarrhoeic	Not stated	Diarrhoeic and non-diarrhoeic	
India	India	India	Nepal	Pakistan	

Table 4.4 (continued)	continued)						
Location	Symptoms	Host age	Study design	Prevalence % positive (no. positive/no. examined)	Diagnostic technique	Molecular analyses gene/species or subtype ^b	Reference
Philippines	Not stated	1–12 days	Longitudinal	3 % (1/38)	Formalin ether sedimentation followed by Kinyon acid fast stain		Villanueva et al. 2010
Venezuela	Not stated	0-12 weeks	Longitudinal	88 % (22/25)	Formalin-		Díaz de
					ethylacetate sedi- mentation and sodium chloride flotation followed by carbol-fuchsin staining		Ramírez et al. 2012
Italy	Asymptomatic	Calves	Cross-sectional	14 % (8/57)	ELISA, followed by formol-ether extraction and IFAT	18S rRNA/ C. parvum	Cacciò et al. 2007
Italy	Not stated	2–60 days/ 43 farms	Cross-sectional		ELISA		Rinaldi et al. 2007a
Italy	Not stated	1–9 weeks/ 90 farms	Cross-sectional	15 % (51/347) Herd 24 % (22/90)	ELISA		Rinaldi et al. 2007b
^a A few studi	^{a}A few studies are not included in the table due to inaccessibility of the publications	n the table due to	inaccessibility of th	he publications			

C. ubiquitum have also been reported (Table 4.4). Most molecular investigations have been done in calves and thus it is not known if the species distribution differs between animals of different age. One of the few studies that include faecal samples from both calves and older animals was done in Egyptian smallholder herds (Helmy et al. 2013). *C. parvum* was most common in calves younger than 3 months but was detected in animals up to 2 years of age. In another study, also from Egypt but from another part of the country, *C. parvum* was only found in calves, whereas none of the sampled cows shed any *Cryptosporidium* oocysts (Amer et al. 2013). When the *C. parvum* isolates were subtyped by sequence analysis of the GP60 gene, subtype families IId and IIa were found, with a majority of IId in both studies. Subtype IId is the dominating subtype family also in cattle in Egypt (Amer et al. 2010; Helmy et al. 2013).

4.2.2 Association of Infection with Clinical Disease

4.2.2.1 Cattle

Cryptosporidium parvum and C. andersoni are the two species that have been associated with clinical disease in cattle. C. parvum infection is considered a major cause of diarrhoea in young calves (Blanchard 2012; Radostits et al. 2007). Calves are often already infected during the first week of life (Uga et al. 2000) and clinical cryptosporidiosis is mostly seen in calves up to 6 weeks of age. The most prominent finding is pasty to watery diarrhoea, sometimes accompanied by lethargy, inappetence, fever, dehydration and/or poor condition. The calf most often recovers spontaneously within 1-2 weeks, but there is a large variation between individuals in how they respond to, and recover from, infection. In some cases the infection may be fatal (Tzipori et al. 1983; Fayer et al. 1998). A decrease in growth rate may be seen in the weeks after the calves have recovered from the acute phase of the disease (Klein et al. 2008), but no long-term effects on growth and performance have been reported. The pathogenesis of bovine cryptosporidiosis is not fully understood but the clinical signs are attributed to both malabsorption and an increase in fluid secretion in the ileum and proximal portions of the large intestine. For references and a brief overview see O'Handley and Olson (2006). Cryptosporidiosis may be seen in individual calves, but frequently it soon develops among the calves into a herd problem. Concomitant infection with other pathogens, e.g. rotavirus, coronavirus and enteropathogenic Escherichia coli (E. coli F5+) can worsen the clinical signs and prolong the duration of illness (Blanchard 2012).

A number of studies have reported an association between *C. parvum* infection and diarrhoea in young calves. Many of these were published at the time when all *Cryptosporidium* oocysts of around 3–5 μ m in diameter were considered to be *C. parvum*. More recent investigations, applying molecular methods to analyse faecal samples from diarrhoeic calves, corroborate these earlier findings. When samples from young calves with diarrhoea were analysed, *C. parvum* is found to be the dominant species (Quílez et al. 2008b; Imre et al. 2011; Karanis et al. 2010; Soba and Logar 2008; Plutzer and Karanis 2007). Interestingly, this dominance of *C. parvum* in diarrheic calves was also seen in a recent Swedish investigation of diarrheic calves (Silverlås et al. 2013), although *C. bovis* is the predominant species in randomly selected calves in Sweden.

Only one experimental trial has been performed with *C. bovis* (Fayer et al. 2005). Three calves under 1–8 weeks of age were orally inoculated with oocysts. This resulted in subclinical infection in 2 of 3 calves. Both animals had, however, previously been infected with *C. parvum* and cross-protective immunity could not be excluded. Calves with diarrhoea are significantly more likely to be infected with *C. parvum* than with *C. bovis* (Silverlås et al. 2013; Starkey et al. 2006; Kváč et al. 2011). Based on these findings, and based on the fact that *C. bovis* is not common in calves, but is a widespread subclinical infection in older animals in most countries, *C. bovis* is commonly considered to be apathogenic to cattle. However, the pathogenic potential deserves further attention as high numbers of *C. bovis* occysts in samples from diarrhoeic calves have been reported, even in the absence of *C. parvum* or other diarrhoeal agents (Silverlås et al. 2010a, b, 2013).

Cryptosporidium ryanae was first described as a separate species by Fayer et al. (2008), and until then it was known as *Cryptosporidium* deer-like genotype. An experimental trial was performed in two colostrum-deprived calves 17–18 days old. Both calves started excreting oocysts 11 days after inoculation, but neither of them showed any clinical signs (Fayer et al. 2008). There are several reports of the distribution of *Cryptosporidium* deer-like genotype and *C. ryanae*. Most studies found a predominance of the parasite in older calves and young stock. So far no association with clinical disease has been reported.

In contrast to the other species, *C. andersoni* infects the abomasum. It does not cause diarrhoea, but *C. andersoni* infections have been associated with maldigestion. The infection may cause moderate to severe weight gain impairment in young stock and reduced milk production in cows (Anderson 1998; Esteban and Anderson 1995; Lindsay et al. 2000).

A major obstacle from a disease control perspective is the lack of effective means to control *Cryptosporidium* infection and decrease the level of contamination of the environment with oocysts. Preventive hygiene measures and good management are currently the most important tools to control cryptosporidiosis. Reducing the number of oocysts ingested by neonatal calves may reduce the severity of infection and allow immunity to develop. A common recommendation is to ensure good hygiene in calf facilities and ascertain that all newborn calves ingest an adequate amount of colostrum during their first 24 h of life. Sick calves should be housed in a clean, warm, and dry environment and isolated to prevent spreading of the infection to other calves. Acutely infected animals may need supportive care with fluid and electrolytes, and milk should be given in small quantities several times daily to optimise digestion and minimise weight loss.

Over the years, several substances have been tested for potential anticryptosporidial effects with limited success (Santín and Trout 2008). Halofuginone lactate has shown some beneficial effects such as milder clinical signs and reduced oocyst output when used as prophylactic treatment (De Waele et al. 2010; Silverlås et al. 2009a). This drug is approved in Europe to treat calf cryptosporidiosis. However, the safety margin is narrow and the substance is toxic at only twice the therapeutic dose, so careful dosage is necessary. Halofuginone lactate treatment should only be considered in herds with severe diarrhoeal problems strongly associated with *C. parvum*. When treatment is used, it should always be in conjunction with applying measures to reduce environmental contamination and risk of infection.

A recent study investigated if an antibody-biocide fusion consisting of a monoclonal antibody "armed" with membrane-disruptive peptides (biocides) could be used for treatment of cryptosporidiosis in calves (Imboden et al. 2012). Calves 36–48 h of age were challenged once with *C. parvum* oocysts and were simultaneously administered the antibody-biocide fusion mixed with milk replacer. The antibody-biocide fusion treatment was repeated 5–8 times. Control calves were given milk replacer with placebo. Calves receiving the antibody-biocide fusion had a significantly higher health score and shed fewer oocysts than control calves. These results suggest that this concept might be effective in cattle, but further testing is necessary (Imboden et al. 2012).

Vaccination is successfully used to control many infectious diseases in livestock. However, it takes weeks for a protective immune response to develop after a vaccine has been administered, and as calves may be exposed to *Cryptosporidium* oocysts immediately after birth, vaccination of newborn calves is unlikely to be successful in preventing cryptosporidiosis. Thus it has been suggested that the most feasible approach is likely to involve passive immunisation (Innes et al. 2011). Dams are immunised in late gestation and their colostrum is fed to the calves. A recent study investigated antibody responses in calves fed colostrum from heifers vaccinated with a recombinant *C. parvum* oocyst surface protein (rCP15/60). The calves had measurable quantities of the specific antibody in their serum. However, as the calves were not subsequently challenged with oocysts it remains to be seen whether this immunisation scheme can also prevent symptomatic infection and eliminate oocyst shedding (Burton et al. 2011).

4.2.2.2 Water Buffalo

An association between *Cryptosporidium* oocyst shedding and diarrhoea in buffalo calves has been reported from investigations performed in Egypt, India and Venezuela (Mohanty and Panda 2012; Bhat et al. 2012; Maurya et al. 2013; El-Khodery and Osman 2008; Díaz de Ramírez et al. 2012) suggesting the *Cryptosporidium* infection is part of the calf diarrhoea syndrome in water buffalo, as it is in cattle. Species identification was only performed in one of these studies, and *C. parvum* was the only species that was found (Maurya et al. 2013).

4.2.3 Infection Dynamics: Oocyst Excretion and Transmission

4.2.3.1 Cattle

Calves begin shedding C. parvum oocysts 2-6 days after infection and shedding continues for 1-13 days (Fayer et al. 1998; Tzipori et al. 1983). During the first 2 weeks an infected calf can shed millions of oocvsts (Faver et al. 1998; Uga et al. 2000) resulting in heavy environmental contamination, and efficient dissemination of the parasite within the herd and to the environment. In faecal samples obtained from symptomatic calves naturally infected with C. parvum 10^6-10^8 occysts per gram faeces (OPG) are often seen (Silverlås et al. 2013). In herds with established C. parvum infection, most calves are excreting oocysts between 2 and 4 weeks of age (O'Handley et al. 1999; Santín et al. 2008; Uga et al. 2000). When Santín et al. (2008) repeatedly sampled the same 30 calves in a dairy herd from birth to 2 years of age they found C. parvum oocysts in faeces of all individuals before they were 3 weeks old, i.e. a cumulative prevalence of 100 %. C. parvum oocysts were also found in samples collected from a calf at 16 weeks of age and from another at 6 months of age, indicating that oocysts can be shed intermittently over a long period after the initial infection. Alternatively, these late-shedding individuals might not have developed a fully protective immunity after the first infection, and rather than this being a sign of prolonged infection, they had acquired new infections. In this study, molecular analyses indicated the same sub-genotype at the GP60 locus. However, this does not necessarily indicate prolonged infection, as re-infection with the same genotype in the environment may occur if the immunity is not protective. It has been suggested that an increase in C. parvum oocyst shedding may occur in adult cows around calving (so called periparturient rise), but to date there have been few reports to support this. In a recent study, however, dams in a suckler beef herd were found to shed low levels of C. parvum oocysts around the time of calving (De Waele et al. 2012).

Only one experimental infection for each of *C. bovis* and *C. ryanae* has been reported so far. Regarding *C. bovis*, one calf shed oocysts from 10 to 28 days after infection and the other only for 1 day (day 12) (Fayer et al. 2005). For *C. ryanae*, oocyst shedding started 11 days after inoculation. Both infected calves excreted oocysts during 15 and 17 consecutive days, respectively (Fayer et al. 2008). Shorter prepatent periods have been seen for both *C. bovis* and *C. ryanae* in natural infections (Silverlås et al. 2010b; Silverlås and Blanco-Penedo 2013). No oocyst excretion rate values were determined from the experimental infections, but in naturally infected calves 300 to 8×10^6 OPG and 100–835,000 OPG have been reported for *C. bovis* and *C. ryanae*, respectively (Silverlås and Blanco-Penedo 2013).

Young stock and adults may also be infected by the larger *C. andersoni* (oocyst size $\sim 7.4 \times 5.5 \,\mu$ m) and may shed oocysts intermittently for many years (Olson et al. 2004; Ralston et al. 2003). A periparturient rise in *C. andersoni* oocyst shedding, seen both as increase in prevalence and in number of oocysts in faeces, has been reported (Ralston et al. 2003).

Several studies have shown that age is associated with *Cryptosporidium* infection and that young calves have the highest risk of being infected (Maddox-Hyttel et al. 2006; Santín et al. 2004; Sturdee et al. 2003; Fayer et al. 2007). This is also the age group that is most often infected with C. parvum and suffers from clinical cryptosporidiosis. Thus, from clinical and zoonotic perspective, knowledge on the epidemiology of cryptosporidiosis in young calves is highly valuable. When potential risk factors for Cryptosporidium infection in pre-weaned calves have been explored, the results differ between studies. One factor that recurs in several studies is the type of flooring in the calf housing area. In Spain, Castro-Hermida et al. (2002) found that straw on the floor or earth floors in the calf pens increased the risk for infection compared with cement flooring, and in Malaysia calves kept in pens with slatted floors and sand floors had an increased risk compared with those in pens with cement floors (Muhid et al. 2011). A protective effect of cement floors was also reported from the USA (Trotz-Williams et al. 2008). It was suggested that the reason for this protective effect of cement floors is that they facilitate thorough cleaning. This assumption corroborates the finding that a low frequency of cleaning of the calf pens increased the risk for infection (Castro-Hermida et al. 2002). It is also consistent with the finding that the use of an empty period in the calf pen between introductions of calves was associated with a lower risk for infection in Danish dairy herds (Maddox-Hyttel et al. 2006). When cows as a cause of infection were investigated, a higher risk of infection was identified in calves that were housed separately from their dams (Duranti et al. 2009), and a lower risk of infection in dairy calves kept with the cow for more than 6 h after birth (Silverlås et al. 2009b).

In one of the few reports to investigate risk factors for infection with different *Cryptosporidium* species in pre-weaned dairy calves to date (Szonyi et al. 2012), risk of infection with *C. parvum* differed to some extent from that of *C. bovis*. Both *C. parvum* and *C. bovis* were more common in the younger calves, but herd size and hay bedding were associated with an increased risk for *C. parvum* infection, whereas Jersey breed was a risk factor for *C. bovis* infection.

4.2.3.2 Water Buffalo

Experimental *Cryptosporidium* infections in water buffalo have not been reported. However, oocyst shedding dynamics were investigated in naturally infected buffalo calves in a farm located in a tropical dry forest area in Venezuela. Twenty-five calves were sampled from birth to 12 weeks of age. Oocysts were detected from day 5 and 72 % of the calves shed oocysts before they were 30 days (Díaz de Ramírez et al. 2012).

Regarding risk factors for infection, there are some reports of seasonal variations in prevalence (Bhat et al. 2012; El-Khodery and Osman 2008; Maurya et al. 2013; Mohanty and Panda 2012), and (El-Khodery and Osman 2008) identified type of flooring, frequency of cleaning and water source as risk factors for infection in small-scale herds in Egypt.

4.2.4 Zoonotic Transmission

4.2.4.1 Cattle

There are numerous reports of cryptosporidiosis outbreaks in humans after contact with infected calves. These have often involved veterinary students and students at farm schools (see, e.g., Gait et al. 2008; Grinberg et al. 2011; Pohjola et al. 1986; Robertson et al. 2006; Kiang et al. 2006), but also young children have fallen ill after visiting petting zoos or open farms (Gormley et al. 2011; Smith et al. 2004). Contact with cattle has been identified as a risk factor for disease also in casecontrol studies of sporadic human cryptosporidiosis (Hunter et al. 2004; Robertson et al. 2002; Roy et al. 2004). Altogether, there is plenty of evidence to conclude that *Cryptosporidium* can be transmitted from calves to humans by direct contact or by contaminated equipment. The risk for zoonotic transmission is likely to be highest in herds with Cryptosporidium associated calf diarrhoeal problems, where oocyst contamination in the barn can reach high levels and where contact with naïve individuals is most likely to occur. Key measures to prevent visitors becoming infected are to ensure good hygiene in the visitor area, providing suitable handwashing facilities and ensure that they are used when workers and visitors leave the premises. C. bovis infections have recently been detected in a few persons living or working on cattle farms (Khan et al. 2010; Ng et al. 2012). It is not known if these were active infections and the implication of these findings is thus unclear.

As molecular typing methods become more accessible, epidemiological studies can investigate C. parvum GP60 subtype distribution in cattle and human populations in different regions. The reports so far indicate that in many areas the subtypes that are most common in cattle are those most often found in humans. For example, C. parvum IIaA15G2R1 was the predominant subtype in both bovine and human infections in Slovenia and Portugal (Soba and Logar 2008; Alves et al. 2006). In New South Wales, Australia C. parvum IIaA18G3R1 dominated in both calves and people living on cattle farms (Ng et al. 2012), whereas IIaA16G2R1 was the predominant genotype identified in beef cattle and humans in Prince Edward Island, Canada (Budu-Amoako et al. 2012c). Further information is provided in the review by Xiao (2010). That the same subtypes are found in cattle and humans might be taken as an indication of zoonotic transmission. However, it is important to note that even when zoonotic C. parvum subtypes are identified in humans, cattle are not necessarily the source of the infection. These zoonotic subtypes can circulate and propagate in the human population in addition to the anthroponotic subtypes. The occasional finding of C. hominis (Smith et al. 2005; Chen and Huang 2012) in cattle highlights the fact that cryptosporidiosis may be transmitted not only from cattle to humans, but also from humans to cattle.

Food-related cryptosporidiosis outbreaks have sometimes been associated with cattle. Foodborne transmission was implicated in cases of children who had drunk unpasteurized milk (Harper et al. 2002) or cider, made from apples collected in an orchard where calves from an infected herd had grazed (Millard et al. 1994).

Other outbreaks in which cattle were suspected as the source involved vegetables that had been sprayed with water that could have been contaminated with cattle faeces. Often there was only circumstantial evidence that cattle were the source of contamination, and it was not possible to exclude other potential sources (see e.g. CDC 2011; Robertson and Chalmers 2013).

Outbreaks of cryptosporidiosis associated with drinking water have often been attributed to contamination of water catchments by cattle manure. The evidence implicating cattle has sometimes been substantial (Bridgman et al. 1995; Smith et al. 1989), but for others the evidence was not conclusive. Grazing cattle or slaughterhouse effluent contaminating Lake Michigan were mentioned as two possible sources of *Cryptosporidium* oocysts in the large outbreak in Milwaukee, Wisconsin in 1993 (Mac Kenzie et al 1994), but retrospective analysis of clinical isolates revealed that it was caused by the anthroponotic species C. hominis (Sulaiman et al. 1998). This was also the case in the most recent outbreaks in UK (compiled by Chalmers 2012) and a large drinking waterborne outbreak in Sweden in 2010 (Anonymous 2011). Given that pre-weaned calves are the most likely age group to shed C. parvum oocysts, any measure to prevent waterborne zoonotic transmission should be directed towards this age group. Protective measures could be to prevent young ruminants from accessing water catchments, and compost or spread calf manure on fields where runoff cannot occur. The manure of older ruminants is generally not a zoonotic concern with respect to Cryptosporidium.

4.2.4.2 Water Buffalo

The seemingly common occurrence of *C. parvum* in buffalo calves highlights the potential role of water buffalo in zoonotic transmission. Thus the same precautions to prevent transmission of the parasite to humans, by direct contact or through food or water, are also applicable to water buffalo.

4.3 Cryptosporidium spp. in Small Ruminants

Sheep (*Ovis aries*) and goats (*Capra aegagrus hircus*) are important in the global agricultural economy – producing meat, milk and wool – both in developing countries such as India and Iran, and industrialised countries such as Australia and The United Kingdom (de Graaf et al. 1999; Noordeen et al. 2000; Robertson 2009). In 2010, the world stocks were approximately one billion sheep and 910 million goats (FAOSTAT 2013a). Asia has the largest populations of both species, with 42 % and 60 % of the total world populations, followed by Africa (FAOSTAT 2013a). According to a FAO report, over 90 % of the goat population can be found in developing countries (FAO 2012). Sheep and goats tend to be managed differently to cattle, with flocks grazing large enclosures rather than being kept indoors. There have been fewer studies on *Cryptosporidium* infection in sheep than in cattle, and

even fewer studies have been performed on goats. Nevertheless, it is known that these protozoans are economically important parasites in both ruminant species (Noordeen et al. 2000; Robertson 2009). Infection and disease was first described in 1974 for sheep (Barker and Carbonell 1974) and in 1981 for goats (Mason et al. 1981). Younger animals are more susceptible to infection than older ones, reflected in high shedding rates and diarrhoeal prevalences in lambs and kids up to 1 month of age, whereas infection in older animals is usually subclinical with lower shedding rates (Vieira et al. 1997).

4.3.1 Occurrence (Prevalence)

As for other animals, ovine and caprine Cryptosporidium infection can be found throughout the world. The prevalence varies widely between studies, from 0 % to 77 % in sheep and from 0 % to 100 % in goats. All age groups are susceptible, but infection is more common in lambs and kids than in older animals (Tables 4.5 and 4.6). Study design factors other than age of sampled animals, such as whether only diarrhoeal animals were sampled or not, if a point prevalence study or a longitudinal study was performed and the diagnostic method(s) used, also can affect prevalence data. The effect of using different diagnostic methods is evident in, for example, Giadinis et al. (2012; see also Tables 4.5 and 4.6) and Ryan et al. (2005), where microscopy resulted in lower prevalences than detected by ELISA and PCR, respectively. Prevalence and species distribution for studies conducted on sheep dating back to 2007 are summarised in Table 4.5. Specific data for studies on sheep published before 2007 can be found in "Cryptosporidium and Cryptosporidiosis" (Santín and Trout 2008; Tables 18.9 and 18.10). Prevalence rates and species distribution for all identified surveys of goats are summarised in Table 4.6.

Several species and genotypes have been identified in sheep, and the species distribution varies between studies and with age of the animals. Cryptosporidium parvum, C. ubiquitum (previously Cryptosporidium cervine/cervid genotype) and C. xiaoi (previously C. bovis-like genotype) are the most common species. Sporadic infection with C. hominis, C. suis, C. andersoni, C. fayeri (previously marsupial genotype I), C. scrofarum (previously pig genotype II), sheep genotype I, and unknown/novel genotypes have been identified (Chalmers et al. 2005; Giles et al. 2009; Karanis et al. 2007; Ryan et al. 2005, 2008; Sweeny et al. 2011b; Wang et al. 2010c). Species distribution differs between studies and between age groups within studies. For instance, C. parvum is commonly found in lambs in Italy, Romania, Spain and the UK (Díaz et al. 2010a; Imre et al. 2013; Mueller-Doblies et al. 2008; Paoletti et al. 2009). In other studies, C. ubiquitum or C. xiaoi is the most common species (Fiuza et al. 2011a; Geurden et al. 2008; Robertson et al. 2010; Sweeny et al. 2011b). For example, Wang et al. (2010c) identified C. ubiquitum in 90 % of all analysed samples, and the species dominated in all age groups, whereas Sweeny et al. (2011b) found C. xiaoi to be the most common

aals Age ≤8 w ≤8 w 2 w-8 m 2 w-8 m Ewes, 4 m prp Ewes, 2 w prp 2 -3 m 4 -5 m 2 -6 m >12 m >12 m preveaned postweaned		Positive	Cryptosporidium spp.	
a 235 2 w-8 m a 235 2 w-8 m b Ewes, 4 m prp b Ewes, 2 w prp a 96 Ewes, 4 m prp b Ewes, 2 w prp a 200 2-3 m 137 ≤10 w 90 2-6 m 35 >12 m 378 preveaned 585 postweaned	Animal prevalence $\%$	herds	(% of all determined)	Reference
a 235 2 w-8 m 96 Ewes, 4 m prp Ewes, 2 w prp 137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveated 585 postveated	9.3-56.3	5/5	C. bovis 36.5	(Yang et al. 2009)
a 235 2 w-8 m 96 Ewes, 4 m prp Ewes, 2 w prp 2-3 m 4-5 m 90 2-6 m 35 >12 m 10 >3 m preveated 585 postveated	Overall 24.5		C. parvum 46.1	
a 235 2 w-8 m a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m preveated 585 postveated			C. ubiquitum 8.7	
a 235 2 w-8 m a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 2-3 m 4-5 m 35 > 12 m 378 preveated 378 preveated 585 postveated			Mixed infections 8.7	
a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 2-3 m 4-5 m 4-5 m 35 >12 m 35 >12 m 10 >3 m 378 preveated 385 postveated	Age group point prevalences: 18.5–42.6	2/2	C. andersoni 1.0 ^d	(Sweeny et al. 2011b)
a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 2-3 m 4-5 m 4-5 m 35 >12 m 35 >12 m 10 >3 m 378 preveated 585 postveated	Cumulative prevalences:		C. parvum 5.9	
a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 2-3 m 4-5 m 137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveated 585 postweated	Herd A: 81.3		C. ubiquitum 9.8	
a 96 Ewes, 4 m prp Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 4-5 m 4-5 m 137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveaned 585 postweaned	Herd B: 71.4		C. xiaoi 73.7	
a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 2-3 m 4-5 m 137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveaned 585 postweaned			Sheep genotype I 1.6	
a 96 Ewes, 4 m prp Ewes, 2 w prp Ewes, 2 w prp 4-5 m 4-5 m 90 $2-3$ m 4-5 m 35 >10 w 35 >12 m 10 >3 m 378 preveated preveated			Mixed infections 8.0	
Eves, 2 w prp 4-5 m 4-5 m 4-5 m $137 \leq 10 \text{ w}$ 90 2-6 m 35 > 12 m 10 > 3 m 378 preveated 585 postweated	prp Herd A, B: 6.3	2/2	C. xiaoi 100 ^d	(Sweeny et al. 2011b)
a 200 2-3 m 4-5 m 137 $\leq 10 \text{ w}$ 90 2-6 m 35 >12 m 10 >3 m 378 preveaned 585 postveaned	prp Herd A, B: 8.3			
4-5 m 137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveaned 585 postveaned	Herd I: 33.6, II: 31.9	2/2	C. parvum 21.0 ^d	(Sweeny et al. 2012)
137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveaned 585 postweaned	Herd I: 28.1, II: 23.6		C. ubiquitum 44.5	
137 $\leq 10 \text{ w}$ 13 90 $2-6 \text{ m}$ 2. 35 >12 m 0 1 $<3 \text{ m}$ 0 378 preveaned 10 585 postweaned 4.	Cumulative prevalences:		C. xiaoi 31.9	
137 $\leq 10 \text{ w}$ 13 90 $2-6 \text{ m}$ 2. 35 >12 \text{ m} 0 1 $<3 \text{ m}$ 0 10 >3 \text{ mexeaned} 10 585 postweaned 4.	Herd I: 40.6		Mixed infections 2.5	
137 ≤10 w 90 2-6 m 35 >12 m 10 >3 m 378 preveated 585 postweated	Herd II: 31.9			
90 2–6 m 35 >12 m 1 <3 m 10 >3 m 378 preveaned 585 postweaned	13.1	4/10	C. parvum 10.0	(Geurden et al. 2008)
90 2-6 m 35 >12 m 1 <3 m 10 >3 m 378 preweaned 585 postweaned			C. ubiquitum 90.0	
35 >12 m 1 <3 m 10 >3 m 378 preveated 585 postweated	2.2		C. ubiquitum 100	(Fiuza et al. 2011a)
1 <3 m 10 >3 m 378 preweaned 585 postweaned	0			
1 <3 m 10 >3 m 378 preweaned 585 postweaned	Overall 1.6			
10 >3 m 378 preveaned 585 postweaned	0	6/0	I	(Sevá et al. 2010)
378 preweaned 585 postweaned	0			
postweaned		5/5	C. andersoni 4.9	(Wang et al. 2010c)
			C. ubiquitum 90.2	
ewe, prp	2.1		C. xiaoi 4.9	
158 ewe, pp 2.5	2.5			
China 213 6–9 m 14.6	14.6	3/8	C. ubiquitum 100 ^e	(Shen et al. 2011)

4 Cryptosporidiosis in Farmed Animals

Table 4.5 (continued)	ued)					
	Number of			Positive	Cryptosporidium spp.	
Country	animals	Age	Animal prevalence %	herds	(% of all determined)	Reference
Cyprus ^c	39	4–15 d	61.5 (mZN) 76.9 (E)	12/15	ND	(Giadinis et al. 2012)
India	55	<3 m	1.8	I	ND	(Maurya et al. 2013)
Italy ^c	21	I	100	I	C. parvum 100	(Drumo et al. 2012)
Italy	149	2 w-3 m	17.4	5/6	C. parvum 100	(Paoletti et al. 2009)
Norway ^a	567	5-6 w	14.6	6/6	C. ubiquitum 83.3	(Robertson et al. 2010)
	528	6–10 w	24.1		C. xiaoi 16.7	
Romania ^c	58	1-7 d	13.8	5/7	C. parvum 83.3	(Imre et al. 2013)
	68	8–14 d	16.2		C. ubiquitum 8.3	
	49	15–21 d	10.2		C. xiaoi 8.3	
Spain ^c	127	<21 d	30.7	17/28	C. parvum 63.6	(Díaz et al. 2010a)
					C. ubiquitum 36.4	
Spain	446	Adult	5.3	13/38	ND	(Castro-Hermida et al. 2007)
St Kilda ^{a, b}		4–≥40 m	9.0–28.6	I	ND	(Craig et al. 2007)
Tunisia	30	<3 m	16.7	3/3	C. xiaoi ^f	(Soltane et al. 2007)
	59	>1 y	8.5			
Turkey ^c	151	≤7 d	44.4	26/34	ND	(Sari et al. 2009)
	104	8–14 d	37.5			
	95	15–21 d	40.0			
	50	22 d ≤ 1 m	22.0			
			Overall 38.8			
United Kingdom	260	Mixed	39.6	11/17	C. parvum 88.9 C. xiaoi ^g 11.1	(Mueller-Doblies et al. 2008; Smith et al. 2010)
d days, w weeks, m month ^a Longitudinal study ^b Feral population	1 months, <i>ND</i> r y	not done, Prp prep	s, ND not done, Prp prepartus, pp post partum, mZN modified Ziehl-Neelsen, E ELISA	dified Ziehl-Ne	elsen, E ELISA	
^c Only diarrhoeal samples collected ^d Species distribution for all positive	amples collecte on for all positi	collected I positive samples independent of age	endent of age			
		· · · · · ·	, ,			

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^eOnly four isolates successfully amplified and sequenced ^fReported as similar to *C. bovis*, 99.5–99.7 % identical to *C. xiaoi* isolates in GenBank ^sReported as *C. bovis*, 100 % identical to *C. xiaoi* isolates in GenBank

					Crvntosnoridium snn	
Country	Number of animals	Age	Animal prevalence %	Positive herds	(% of all determined)	Reference
Belgium	148	$\leq \! 10 \text{ w}$	9.5	6/10	C. parvum (100)	(Geurden et al. 2008)
Brazil	49	<12 m	10.2	2/6	ND	(Bomfim et al. 2005)
	56	>12 m	0			
			Overall 4.8			
China	42	I	35.7	I	C. xiaoi ^c	(Karanis et al. 2007)
					Novel genotype ^d	
Cyprus ^b	75	4–15 d	64(mZN) 86.7 (E)	25/28	ND	(Giadinis et al. 2012)
France ^a	40	0-weaning	Age related point prevalences: 0-100	1/1	C. xiaoi (94.7)	(Rieux et al. 2013)
			Cumulative 77.5		C. parvum (5.3)	
France	200	5-12 m	2.5	I	QN	(Castro-Hermida et al. 2005)
France	879	5-30 d	16.2	32/60 (53.3)	ND	(Delafosse et al. 2006)
India	116	<3 m	3.5	I	C. parvum (100)	(Maurya et al. 2013)
Italy ^b	21	I	100	I	C. parvum (100)	(Drumo et al. 2012)
Spain ^b	5	<21 d	40	1/1	C. xiaoi	(Díaz et al. 2010b)
Spain	116	Adult	7.7	6/20 (30)	ND	(Castro-Hermida et al. 2007)
Spain	134	≤15 d	10.4	4/4	ND	(Sanz Ceballos et al. 2009)
	144	>15 d-2 m	13.4			
	304	>2 m	25.2			
Sri Lanka	558	≤6 m	33.0	23/24	ND	(Noordeen et al. 2000)
	133	6–12 m	30.8			
	329	>12 m	20.1			
			Overall 28.5			
Sri Lanka	72	1-12 m	$0-75^{\rm b}$	1/1	ND	(Noordeen et al. 2001)
Tunisia	184	1-7 y	0	0/5	Ι	(Soltane et al. 2007)
UK	15	Adult	20	1/4	Negative	(Smith et al. 2010)
<i>d</i> days, <i>w</i> weeks, <i>m</i> ^a Longitudinal study	<i>d</i> days, <i>w</i> weeks, <i>m</i> months, <i>ND</i> n ^a Longitudinal study	ot done, Prp pi	s, ND not done, P_{TP} prepartus, pp post partum, mZN modified Ziehl-Neelsen, E ELISA	odified Ziehl-Neel	lsen, E ELISA	
^b Only diar	^b Only diarrhoeal samples collected	ч 				
Only two	Only two samples successfully amplified	nplified		(-	
"Clustered	with C. bovis, C. ryam	<i>ie</i> (previously ("Clustered with C. bovis, C. ryanae (previously Cryptosporiatium deerlike genotype) and Cryptosporiatium deer genotype	pe) and <i>Cryptospo</i>	<i>ridium</i> deer genotype	

Table 4.6 Published studies on prevalence of Cryptosporidium infections in goats

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species in lambs (73.1 %) and the only species in ewes (Table 4.5). *Cryptosporidium bovis* has also been reported in sheep (Mueller-Doblies et al. 2008; Soltane et al. 2007; Wang et al. 2010c). Whether *C. bovis* has actually been identified in sheep, or if it is the closely related species *C. xiaoi* is uncertain. For instance, Soltane et al. (2007) reported isolates similar to *C. bovis*, and Mueller-Doblies et al. (2008) reported *C. bovis*, but a BLAST search of the GenBank accession numbers identified these isolates as *C. xiaoi*. For the isolates reported as *C. bovis* by Yang et al. (2009), no GenBank accession numbers are available, so the true identity of those isolates is uncertain. Similarly, no GenBank records are available from the study of Ryan et al. (2005) reporting the "New bovine B genotype" in sheep. Since *C. bovis* was first identified as the bovine B genotype, this could actually be the "*C. bovis*-like genotype", i.e. *C. xiaoi*.

Cryptosporidium andersoni has been identified in a few naturally infected adult sheep (Wang et al. 2010c), but experimental infection in 4-month-old lambs failed (Kváč et al. 2004).

A couple of apparently related surveys from Mexico have been published, but because of lack of clarity in the data, they will not be reviewed in this text. Two studies from Brazil (Sevá et al. 2010) and Mongolia (Burenbaatar et al. 2008) failed to identify *Cryptosporidium* in any of the collected samples, but the number of sampled sheep was small -11 and 5 animals, respectively.

Because of the small number of studies and isolates analysed, it is hard to draw any conclusion about the species distribution in goats. *Cryptosporidium parvum*, *C. xiaoi* and a novel genotype have been identified in naturally infected goats (Table 4.5). In addition, one report of natural infection with *C. hominis* is also available (Giles et al. 2009). The identification of *C. xiaoi* in a number of samples is in contrast with a failed attempt to infect 36-week-old goats to determine the host range of *C. xiaoi* (Fayer and Santín 2009). Because there is only scant information about cryptosporidiosis in goats, we do not know if an age-related resistance or immunity from a previous *Cryptosporidium* infection could have affected this experiment, as natural infections indicate that *C. xiaoi* is infectious to goats.

Two studies from Mongolia (Burenbaatar et al. 2008) and the United Kingdom (Smith et al. 2010) failed to identify *Cryptosporidium* in any of the collected samples, but the number of sampled goats was small -16 and 15 animals, respectively.

4.3.2 Association of Infection with Clinical Disease

Cryptosporidiosis has been associated with high morbidity and mortality rates in both lambs and goat kids (Cacciò et al. 2013; Chartier et al. 1995; de Graaf et al. 1999; Giadinis et al. 2007, 2012; Johnson et al. 1999; Munoz et al. 1996; Paraud et al. 2011; Vieira et al. 1997). High mortality has been described both from natural infection and from experimental studies, where infection doses are generally high (Chartier et al. 1995; Giadinis et al. 2007; Paraud et al. 2010). In fact, it has been stated to be one of the most important pathogens associated with diarrhoeal disease and mortality in neonatal lambs and kids (Quílez et al. 2008a).

Anorexia and apathy/depression are common symptoms, accompanied by abdominal pain and pasty to watery, yellow and foul-smelling diarrhoea (de Graaf et al. 1999; Snodgrass et al. 1984). Diarrhoea can last from a few days up to 2 weeks (de Graaf et al. 1999). Faecal consistency is correlated with oocyst excretion (de Graaf et al. 1999; Paraud et al. 2010, 2011), and a longer duration of diarrhoea is potentially associated with infection early in life (Paraud et al. 2010). Body condition score and growth are affected (de Graaf et al. 1999), probably due to both anorexia and the intestinal damage, that can reduce nutrient uptake for weeks (de Graaf et al. 1999; Klein et al. 2008).

Infection in animals older than 1 month is usually subclinical, and even younger animals can be subclinically infected. However, the infection can still affect production, with reduced body condition score (Sweeny et al. 2011a, 2012), reduced growth rate, and reduced carcass weight and dressing percentage at slaughter (Sweeny et al. 2011a).

As discussed above for cattle, before molecular methods were developed *C. parvum* was the only species considered to infect and cause disease in sheep and goats (Chartier et al. 1995; de Graaf et al. 1999; Munoz et al. 1996). *Cryptosporidium parvum* infection has since been associated with diarrhoea in studies using molecular methods (Caccio et al. 2013; Díaz et al. 2010a; Drumo et al. 2012; Imre et al. 2013; Mueller-Doblies et al. 2008). However, *C. xiaoi* has also been associated with mild to severe diarrhoea and mortality (Díaz et al. 2010b; Navarro-i-Martinez et al. 2007; Rieux et al. 2013), and *C. ubiquitum* too has been found in a few diarrhoeal samples from lambs (Díaz et al. 2010a), indicating that *C. parvum* is not the only pathogenic species in small ruminants.

4.3.3 Infection Dynamics: Oocyst Excretion and Transmission

The prepatent period is 3–4 days in goat kids (Paraud et al. 2010) and 2–7 days in lambs (de Graaf et al. 1999). The patent period can last for at least 13 days (Paraud et al. 2010). Shedding peaks a few days to a week into the patent period, and maximum shedding can be as high as 2×10^7 OPG (Rieux et al. 2013). The length of the patent period and shedding intensity are determined by age, immune status and infection dose (de Graaf et al. 1999). A natural age-related resistance to infection seems to be present. In one study, the prepatent period increased and intensity of shedding at 17 days of age and excretion peaked at a mean of 23×10^4 OPG 5–11 days later (Rieux et al. 2013), whereas another group of animals studied by the same authors started shedding at the age of 10 days, with a mean peak of 3×10^6

OPG 0–4 days later, indicating higher virulence, infection pressure or an age-related higher sensitivity in the latter group.

Factors such as hygienic conditions, milking practices, herd size (population density), season, climatic zone (within a country), and lambing/kidding season are factors that have been associated with prevalence of infection, prevalence of clinical cryptosporidiosis and intensity of oocyst shedding (Alonso-Fresan et al. 2009; Bomfim et al. 2005; Craig et al. 2007; Delafosse et al. 2006; Giadinis et al. 2012; Maurya et al. 2013; Noordeen et al. 2000, 2001). However, factors associated with infection and shedding intensity are also impacted by different management systems and climatic conditions; results from small farms in, for example, India cannot be extrapolated to large-scale farming in, for example, the United Kingdom.

4.3.4 Zoonotic Transmission

The significance of sheep and goats as reservoirs for zoonotic cryptosporidiosis is unclear (Robertson 2009). The first case of suspected zoonotic transmission from sheep was described in 1989 (Casemore 1989), but at that time diagnosis was based solely on microscopy and thus zoonotic transmission cannot be confirmed. However, since the introduction of molecular tools in diagnostics, a number of cases and outbreaks with suspected or confirmed zoonotic transfer from sheep have been described (Cacciò et al. 2013; Gormley et al. 2011). In the UK, where sheep farming is an important industry, a seasonal pattern with spring and autumn peaks of human cryptosporidiosis cases is observed, with the spring peak concurring with the lambing season (Anonymous 2002; Gormley et al. 2011; McLauchlin et al. 2000; Nichols et al. 2006). Lambs in petting zoos seem to be a common infection source (Chalmers et al. 2005; Elwin et al. 2001; Gormley et al. 2011; Pritchard et al. 2007). The incidence of human cryptosporidiosis, especially due to C. parvum, dropped significantly during the foot-and-mouth outbreak in the spring and summer 2001 (Hunter et al. 2003) when >6 million livestock animals (~4.9 million sheep; Anonymous 2001) were slaughtered and there were restrictions in animal movements and farm visits, providing further evidence of the importance of zoonotic transmission in this region.

In 2012, an outbreak occurred in Norwegian schoolchildren visiting a farm raising several animal species. *Cryptosporidium parvum* oocysts of an identical and unusual GP60 subtype were identified in faecal samples from six human patients, two lambs and one goat kid. Another human outbreak with the same *C. parvum* subtype had occurred at the same farm 3 years previously, but at that time very few oocysts were detected in animal faecal samples and molecular analyses were not conducted (Lange et al. submitted). Cacciò et al. (2013) described a case where a farmer's son fell ill with cryptosporidiosis, being infected with the same and unusual *C. parvum* subtype that caused high morbidity and mortality in the farm's lambs (Table 4.7).

Studies on *C. parvum* GP60 subtypes have been performed with sheep and goat isolates, and all isolates were found to belong to the IIa and IId families (Table 4.7).

Host			
Country	GP60 subtype	Number of samples	Reference
Sheep			
Australia ^a	IIa	1	(O'Brien et al. 2008)
Italy	IIaA20G2R1	3	(Cacciò et al. 2013)
	IIaA15G2R1	1	
Belgium	IIaA15G2R1	1	(Geurden et al. 2008)
Norway	IIaA19G1R1	2	(Lange et al. submitted)
Romania	IIaA17G1R1	2	(Imre et al. 2013)
	IIaA16G1R1	1	
	IIdA20G1	2	
	IIdA24G1	1	
	IIdA22G2R1	1	
Spain	IIaA15G2R1	3	(Díaz et al. 2010a)
	IIaA16G3R1	7	
Spain	IIaA15G2R1	2	(Quílez et al. 2008a)
	IIaA18G3R1	1	
	IIdA14G1	2	
	IIdA15G1	3	
	IIdA17G1a	44	
	IIdA17G1b	26	
	IIdA18G1	15	
	IIdA19G1	33	
	IIdA21G1	1	
	IIdA22G1	2	
	IIdA24G1	2	
United Kingdom	IIaA15G2R1	3	(Chalmers et al. 2005)
United Kingdom	IIaA15G2R1	1	(Smith et al. 2010)
	IIaA17G1R1	9	
	IIaA17G2R1	1	
Goats			
Belgium	IIaA15G2R1	3	(Geurden et al. 2008)
	IIdA22G1	8	
Norway	IIaA19G1R1	1	(Lange et al. submitted)
Spain	IIdA17G1a	8	(Quílez et al. 2008a)
	IIdA19G1	4	
	IIdA25G1	2	
	IIdA26G1	3	
Italy ^a	IIa	1	(O'Brien et al. 2008)

Table 4.7 Cryptosporidium parvum GP60 subtypes identified in sheep and goats

^aIIaA12G2R1 or IIaA15G2R1 or IIaA19G4R1 were identified in a sheep, a goat and an alpaca isolate, but it is not reported which subtype was identified in which sample

Several studies using multi-locus genotyping (MLG) have found evidence of specific host associated *C. parvum* populations (Drumo et al. 2012; Mallon et al. 2003a, b; Morrison et al. 2008). In the United Kingdom, sheep MLGs clustered with human and bovine isolates (Mallon et al. 2003a, b), indicating frequent zoonotic transmission, whereas only one of the 34 MLGs identified in

sheep/goats in Italy was also identified in human samples (Drumo et al. 2012), indicating a low rate of zoonotic transmission. However, the latter study included very few human isolates.

Zoonotic transmission is commonly observed with *C. parvum*, but *C. ubiquitum* has also been identified in a number of sporadic human cases (Cieloszyk et al. 2012; Elwin et al. 2012; Feltus et al. 2006; Leoni et al. 2006; Ong et al. 2002; Soba et al. 2006; Trotz-Williams et al. 2006). Oocysts of this species have been identified in storm water, wastewater, raw water and drinking water (Jiang et al. 2005; Liu et al. 2011; Nichols et al. 2010; Van Dyke et al. 2012). In Scotland, *C. ubiquitum* was the third most common species in raw water and the most common species identified in drinking water (Nichols et al. 2010). Thus, in areas where *C. ubiquitum* is common in sheep and goats, this species could be a more important cause of zoonotic infection than *C. parvum*. In addition, the relatively common presence of this species in water indicates a potential for waterborne outbreaks.

Natural infection with *C. hominis* has been reported in one goat and two sheep (Giles et al. 2009; Ryan et al. 2005) and in three lambs following experimental infection (Ebeid et al. 2003; Giles et al. 2001), but since animals are not natural hosts for this species, risk of zoonotic transmission with this species should be negligible compared with the risk of human-to-human transmission.

It is important to note that because *Cryptosporidium* infection can be subclinical, the zoonotic potential is not restricted to contact with diarrhoeic *Cryptosporidium*-infected animals (Pritchard et al. 2007).

4.4 Cryptosporidium spp. in Pigs

Since domestication around 4900 BC in China, the pig has been an important food source (Moeller and Crespo 2009). Pigs are farmed worldwide, with the global swine inventory estimated at over 800 million in 2002. Because Asian countries are major consumers but do not produce sufficient pigs for their needs, there is a significant international trade in live and slaughtered pigs. China has the world's largest pig population, mostly small herds consisting of only a few animals, and is a net importer of pigs. The United States, European Union, and Canada are major exporters with relatively few but very large production units (Moeller and Crespo 2009). The global trend is for fewer producers responsible for larger numbers of pigs, and more concentration within the swine industry.

4.4.1 Occurrence (Prevalence)

Pigs are the primary host for *C. suis* (Ryan et al. 2004) formerly identified as *Cryptosporidium* pig genotype I and for *C. scrofarum* (Kváč et al. 2013) formerly identified as *Cryptosporidium* pig genotype II. Farm pigs have also been found

infected with *C. parvum*, *C. muris*, on one occasion with *C. tyzzeri*, a species common to mice, and with the novel genotype isolate Eire 65.5 (Kváč et al. 2013). Cryptosporidiosis occurs in pigs of all ages in 21 countries on 6 continents (Table 4.8). Before molecular methods were developed *C. parvum* was thought to infect 152 species of mammals and to consist of several genotypes. Consequently some early studies erroneously reported *C. parvum* infection in pigs based on the identification of oocysts in faeces by microscopy. Subsequent use of molecular methods provided the necessary tools to identify and distinguish species.

Overall, prevalence data for locations, herds and age groups vary greatly and are not directly comparable because some data represent pooled samples (some from litters, others from fecal slurry), some data originate from single farms while other data come from multiple farms. Some surveys have studied individual pigs at various ages, or only those pigs with diarrhoea, or simply specimens submitted to diagnostic laboratories from unspecified locations (Table 4.8). Even in comparable populations, such as preweaned pigs in the same country or indifferent countries, data differences are too great to draw any conclusions on prevalence. For example: in Australia- reports of 32.7 % versus 6.0 % prevalence (Johnston et al. 2008 vs. Ryan et al. 2003a); in the Czech Republic- reports of 21.8 % versus 5.7 % prevalence (Kváč et al. 2009a vs. Vitovec et al. 2006); or between Serbia and Spain - reports of 32 versus 0 % prevalence (Mišić et al. 2003 vs. Quílez et al. 1996a). Some studies found significant association between the presence of a particular species and the pigs' age, with C. suis prominent in piglets and C. scrofarum prominent in weaners (Enemark et al. 2003). In contrast, others found no significant association between species and age or housing conditions (Featherstone et al. 2010b). These prevalence data reflect vast differences in management practices from location to location with too many unknown factors to draw valid conclusions on cause and effect or location within the 49 cited studies in Table 4.8 that reported a prevalence of infection between 0.1 % and 100 %. The only variable repeatedly associated with detection of *Cryptosporidium* is age. Most positive samples were from weaters and growers (Table 4.8). Generally, prevalence increased until pigs were 10 weeks of age, then gradually declined.

4.4.2 Association of Infection with Clinical Disease

The first reports of cryptosporidiosis in pigs found one piglet among 81 herds of nursing piglets with necrotic enteritis, but the significance of this finding was described as unknown (Bergeland 1977) and *Cryptosporidium* was found at necropsy in three pigs without clinical signs (Kennedy et al. 1977).

Although a higher prevalence of diarrhoea was found in *Cryptosporidium*-infected pigs than in uninfected pigs (Hamnes et al. 2007), others found no significant relationship between infection and diarrhoea (Quílez et al. 1996b; Guselle et al. 2003; Maddox-Hyttel et al. 2006; Vitovec et al. 2006; Suárez-Luengas et al. 2007).

I able 4.0 Location, age,	tion, age, and prevalence of <i>Cryptosportatum</i> in pigs identified by microscopy and molecular methods	sporiaium in pigs idenii	nea by microscopy and	a molecular met	SDOL	
					Molecular identification/species	
Location	Age	No. of animals/farms Prevalence or locations (% positive	Prevalence (% positive)	Microscopy (M)	or genotype (no. or % identified) ^a	Reference
Australia	Not specified	78/diagnostic lab	0	М	No/unknown	O'Donoghue et al. 1987
Australia	3–8 weeks	Samples from 3 herds with clinical disease	3 herds: 2, 5, and 4 positive samples	W	18S rRNA/C. suis (8) Morgan C. parvum (4) et al.	Morgan et al. 1999
Australia	1–8 weeks	648/22 indoor and outdoor herds	6.03 0.5 17.2	W	18S rRNA/C. suis C. scrofarum	Ryan et al. 2003a
Australia	pre-weaned weaned	156 123/4 piggeries	32.7 10.6	M	18S rRNA C. suis (6) C. scofarum (32)	Johnson et al. 2008
Brazil São Paulo	Not specified	25	0	М	No/unknown	Sevá et al. 2010
Brazil Rio de Janeiro	Not specified	91/10 piggeries	2.2	Not indicated 18S rRNA C. scrofaru	18S rRNA C. scrofarum	Fiuza et al. 2011b
Canada	5-15 days	1,453 diarrhoeic samples/diagnos- tic lab	0.5	Μ	No/unknown	Sanford 1983
Canada	1-30 weeks	3491samples/diag- nostic lab	5.3	М	No/unknown	Sanford 1987
Canada	Not specified	100/5 farms	11	М	No/unknown	Olson et al. 1997b
Canada Alberta Not specified	Not specified	25–50 samples/1 farm	0	М	No/unknown	Heitman et al. 2002
Canada	21 days-6 months	33/1 farm	100 longitudinal study, cumulative prevalence	М	HSP70/C. suis (10)	Guselle et al. 2003

Table 4.8 Location, age, and prevalence of Cryptosporidium in pigs identified by microscopy and molecular methods

<i>m</i> Farzan et al. 2011	Bu	Wang et al. 2010b	Lai et al. 2011	Chen et al. 2011	Yin et al. 2011	Vitovec et al. 2006	Kváč et al. 2009a	Kváč et al. 2009b
18S rRNA/C. parvum Farzan C. scrofarum et a	18S rDNA HSP70/C. suis, S. scofarum, C. parvum, C. tyzzeri	18S rRNA C. suis C. scrofarum	No/unknown	18S rRNA C. suis (82.6 %) C. scrofarum (8.7 %) mixed infection (8.7 %)	18S rRNA C. scrofarum	18S rRNA C. suis (394)	18S rRNA C. suis, C. muris C. scrofarum	18S rRNA GP60 C. suis, C. parvum C. scrofarum
Μ	Μ	W	M	Μ	Not indicated 18S rRNA C. scrofaru	W	W	M
/ 55.7	26	10.2 20.6 0 Avs 8.2		34.4	14.3–25.0	0 5.7 24.1	21.8 29.0 17.1 2.5 Ave 21.2	123/14 farms
122 pooled samples/ 55.7 10 farms	633/21 herds	1,350/14 farms in 10 prefectures	2,971/14 intensive 29 extensive farms in 13 counties	2,323/8 farms	94/6 farms	135 3,368 835/8 farms	119 131 123 40 Total 413/1 farm	Finishers sows
		<l< td=""><td>Boars and sows fatteners, growers, weaners</td><td>60–180 days of age</td><td>Not specified</td><td>Sows Pre-weaned weaned</td><td>Pre-weaners Starters Pre-growers Sows</td><td>Abattoir</td></l<>	Boars and sows fatteners, growers, weaners	60–180 days of age	Not specified	Sows Pre-weaned weaned	Pre-weaners Starters Pre-growers Sows	Abattoir
Canada Ontario	Canada Prince Edward Island	China Henan	China Chongqing	China Shanghai	China Yangtze River Delta	Czech Republic	Czech Republic	Czech Republic South Bohemia

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Table 4.8 (continued)	tinued)					
					Molecular identification/species	
Location	Age	No. of animals/farms Prevalence or locations (% positive	Prevalence (% positive)	Microscopy (M)	or genotype (no. or % identified) ^a	Reference
Croatia	Not specified	5 pigs each/ 38 production	65.8 of all units	W	No/unknown	Bilic and Bilkei 2006
		units				
Cuba Villa	Suckling weaned	45	6.7	Е	No/C. parvum based	de la Fe
Clara		45	13.3		on DAS/ELISA	et al. 2013
		All diarrhoeic/ 6 piggeries				
Denmark	<1 month	488	6	М	No/unknown	Maddox-Hyttel
	1–4 months	504	71			et al. 2006
	SOWS	245/50 herds	4			
Denmark	Same as above	1,237/50 herds	Same as above	Not indicated	Not indicated 18S rRNA, HSP70 C. suis (50)	Langkjaer et al. 2007
					C. scofarum (133)	
England East	Suckler	20	20.0	Μ	18S rRNA	Featherstone
Anglia	Weaner	36	38.9		C. scrofarum (64.1 %)	et al. 2010b
	Grower	93	48.4			
	Finisher	119	36.9		C parvum (20.5 %)	
	Adult	39	30.8		C suis (15.4 %)/39	
		Total 308/72 farms	Avg 38.6		samples	
		Diagnostic lab				
Germany	1-42 days	287 diarrhoeic samples/	1.4	M	No/unknown	Wieler et al. 2001
		24 farms				
Germany	Not specified	1,427 samples/ diagnostic lab	0.1	М	No/unknown	Epe et al. 2004

Ireland	Not applicable	56 slurry samples/ 33 farms	50 at 1 farm 40.6 at all other farms	М	No/unknown	Xiao et al. 2006
Ireland	Weaners Finishers Sows Gilts Boars	342/5 units in 4 counties	15.0 7.4 13.3 Rare Rare Mean: 11.4	Not indicated 18S rRNA C. suis, C. C. parvum,	18S rRNA C. suis, C. scofarum, C. parvum, C. muris	Zintl et al. 2007
Italy	Piglets	200 litters (pooled feces from each litter)	0.5	W	No/unknown	Canestri-Trotti et al. 1984
Japan Kanagawa	1–3 months 6 months	232 252/8 farms	33.2 NS	М	No/unknown	Izumiyama et al. 2001
Japan Hokkaido	6 months	108/abattoir	0	М	No/unknown	Koyama et al. 2005
Korea	6–8 months	500/location(s) not specified	19.6	M	No/unknown	Rhee et al. 1991
Korea	Not specified	589/4 slaughter locations	62	M	No/unknown	Yu and Seo 2004
Malawi	Not specified	92/2 regions over 3 seasons	17.7–60	M	No/unknown	Banda et al. 2009
Norway	433 days	684 litters (each pooled)/100 indoor farms	8.3	М	18S rRNA COWP, Actin/C. parvun C. suis C. scrofarun	Hannes et al. 2007
Serbia	1–30 days 1–3 months 3–6 months	50 40 38	32 62.5 44.7	М	No/unknown	Mišić et al. 2003

Table 4.8 (continued)	tinued)					
					Molecular identification/species	
Location	Age	No. of animals/farms Prevalence or locations (% positive	Prevalence (% positive)	Microscopy (M)	or genotype (no. or % identified) ^a	Reference
	6–9 months	45	31.1			
	9–12 months	42	23.8			
	>12 months	45	15.5			
		Total 260				
Spain	<2 months	329/farms and	3 (all in 1 litter of ten M	М	No/unknown	Villacorta
		abattoirs	OU-day-old pigs)			et al. 1991
Spain	1–2 months	36	36	М	No/unknown	Fleta et al. 1995
	2–6 months	11	6			
	mature	10 abattoirs	0			
Spain Aragon	<1 months	49	0	М	No/unknown	Quílez
	1–2 months	49	59.2			et al. 1996a
	2–6 months	312	34.3			
	mature	210	0			
		Total 620/27 farms				
Spain Aragon	<1 months	14	0	М	No/unknown	Quílez
	1–2 months	16	87.5			et al. 1996b
	2–6 months	30	56.7			
	Mature	30	0			
		Total 90				
Spain Zaragosa	Weaned (1–2 months)	75	30.7	М	18S rRNA	Suárez-Luengas
	Fattening (2–6 months)	42	11.9		C. suis	et al. 2007
	Sows	25	16.0		C. scrofarum	
		Total 142/24 farms	22.5			
Trinidad and Tobago	Not specified	275/locations not specified	19.6	М	No/unknown	Kaminjolo et al. 1993
0		1				

Trinidad	Not specified	52/3 locations	1.9	М	No/unknown	Adesiyun et al 2001
USA-California Not specified	Not specified	200/stockyard	S	М	No/unknown	Tacal et al. 1987
USA Ohio	7–25 days	44 1	15.9 of litters	Μ	No/unknown	Xiao et al. 1994
	5–8 weeks	176 200 î	~ ~			
	Sows	62/2 tarms	0			
USA-California ≤8 months	≤8 months	62	11	М	No/unknown	Atwill
	≥ 9 months	159/10 feral groups	3			et al. 1997
USA Georgia	Separate farrowing, nursing,	10 swine waste	100 (every sample	М	18S rRNA/C. suis and Jenkins	Jenkins
	finishing, and gestation	lagoons;	was positive)		C. scofarum	et al. 2010
	effluents	12 monthly			(95–100 %); also	
		samples from			C. muris and	
		each			C. parvum	
Vietnam	4-8 weeks	farms	23.5	М	No/unknown	Koudela
						et al. 1986
^a The species nam that appeared in	^a The species names C. suis and C. scrofarum are used throughout. In some places they replaced the former designations Cryptosporidium pig genotype I and II that appeared in the original references published before these species were named	l throughout. In some pl fore these species were	aces they replaced the f named	ormer designati	ons Cryptosporidium pig	genotype I and II

Cryptosporidium was detected histologically in the microvillus brush border of 5.3 % of 3,491 pigs from 133 farms examined for routine diagnostic evaluation (Sanford 1987). Most infected pigs were 6–12 weeks old. Organisms were detected in the jejunum, ileum, caecum, and colon, but primarily in microvilli of dome epithelium in the ileum. Twenty six percent of *Cryptosporidium*-infected pigs had diarrhoea but most of these also had other primary agents capable of causing diarrhoea. Similar observations have been made by others. Whereas most infections are asymptomatic or cause only mild, non-specific colitis (Higgins 1999), pigs known to be naturally infected with *C. suis* or *C. scrofarum* have not been found with clinical signs of infection while pigs infected with *C. parvum* or co-infected other enteropathogens such as rotavirus, *Salmonella*, or *Isospora* have had associated diarrhoea and some have died (Enemark et al. 2003; Núñez et al. 2007).

Experimental infections with different species have helped to clarify the relationship of species with clinical disease. Pigs experimentally infected with *C. suis* (Enemark et al. 2003) or *C. scrofarum* (Kváč et al. 2013) showed no clinical signs. The pathogenicity of *C. parvum* isolated from calves was demonstrated in early transmission studies to pigs (Moon and Bemrick 1981; Tzipori et al. 1981b, 1982; Argenzio et al. 1990; Vitovec and Koudela 1992; Pereira et al. 2002). Experimental infection with the avian species, *C. meleagridis*, obtained from a human infection, consistently resulted in oocyst excretion and diarrhoea in pigs, although mucosal changes were milder than those described for *C. parvum* (Akiyoshi et al. 2003). Piglets infected with *C. suis* had mild or no clinical signs despite excreting large numbers of oocysts, in contrast to those infected with *C. parvum* that had diarrhoea for a mean duration of 3.5 days and developed inappetence, depression and vomiting (Enemark et al. 2003).

Developmental stages of *Cryptosporidium* have been observed throughout the intestinal tract. Villous atrophy, villous fusion, crypt hyperplasia, and cellular infiltration of the lamina propria have been observed (Kennedy et al. 1977; Moon and Bemrick 1981; Tzipori et al. 1982, 1994; Sanford 1987; Vitovec and Koudela 1992; Argenzio et al. 1990; Pereira et al. 2002; Enemark et al. 2003; Núñez et al. 2003; Vitovec et al. 2006). Lesions caused by *C. parvum* were the most severe, as were clinical signs associated with that species. Changes in the location of stages have been noted. In the first days of infection more stages were found in the proximal intestine, but later more stages were found in distal locations (Tzipori et al. 1982; Vitovec and Koudela 1992). Extra-intestinal infections also have been reported in pigs. In two naturally infected piglets, the gall bladder was infected (Fleta et al. 1995). In experimentally immunosuppressed piglets, the gall bladder, bile ducts, and pancreatic ducts were found infected (Healey et al. 1997). Infections in the trachea and conjunctiva were detected in experimentally infected normal piglets (Heine et al. 1984).

4.4.3 Infection Dynamics: Oocyst Excretion and Transmission

A survey of faecal slurry from swine finishing operations in Ireland found C. suis, C. scrofarum and C. muris and concluded that Cryptosporidium oocysts can persist in treated slurry and potentially contaminate surface water through improper discharge or uncontrolled runoff (Xiao et al. 2006). Hamnes et al. (2007) reported C. suis and C. scrofarum in faces of suckling pigs in Norway and reasoned that farrowing operations were sources of these parasites. Additional data on oocyst concentrations, numbers of oocysts excreted, how long oocysts remain infectious under environmental condition, and modes of transmission of Cryptosporidium species and genotypes are rare or non-existent. A year-long investigation was conducted at four types of swine operations (finishing, farrowing, nursery and gestation) in Georgia, USA (Jenkins et al. 2010). Mean oocyst concentrations ranged from 11 to 354 oocysts per ml of lagoon effluent; the nursery had the highest concentration of oocysts and the greatest percentage of viable oocysts (24.2 %), C. suis and C. scrofarum were the dominant species with some C. muris and C. parvum. Experimental attempts to transmit C. scrofarum to adult SCID mice, adult BALB/c mice, Mongolian gerbils, southern multimammate mice, yellow-necked mice, and guinea pigs were unsuccessful, suggesting that rodents are an unlikely source of transmission of this species under natural conditions (Kváč et al. 2013).

4.4.4 Zoonotic Transmission

Cryptosporidium suis was detected by immunofluorescence microscopy and RFLP analysis of PCR products in stools from an HIV patient in Peru (Xiao et al. 2002a). The patient was not severely immunosuppressed and was asymptomatic. He had a dog but reported no contact with other animals or animal faeces, including pigs and pig faeces, so the source and method of transmission are unknown.

4.5 *Cryptosporidium* spp. in Other Farmed Mammals

4.5.1 Cryptosporidium in Farmed Deer

Many countries, including Australia, Canada, China, Korea, Norway, Russia, Sweden, UK, USA and Vietnam, have thriving deer farming industries. New Zealand, a country where deer are not native, has the world's largest and most advanced deer farming industry. Although it is difficult to find estimates on the numbers of deer farmed worldwide, more than one million deer were being farmed in New Zealand in 2011 (Sources: Statistics New Zealand and Deer Industry New Zealand http://www.deernz.org/about-deer-industry/nz-deer-industry) – compared with five million dairy cows – and there are over 2,800 deer farmers. More than 90 % of the New Zealand deer industry's products are exported, with approximately half of the export going to Germany and Benelux.

Species of deer which are commercially farmed varies regionally, but the following species are now being farmed in various parts of the world: red deer (*Cervus elaphus*), wapiti or elk (*Cervus canadensis*), fallow deer (*Dama dama*), sika (*Cervus nippon*), rusa deer (*Rusa timorensis*), and reindeer (*Rangifer tarandus*) (FAO http://www.fao.org/docrep/004/X6529E/X6529E02.htm).

4.5.1.1 Occurrence (Prevalence)

Although farmed deer are an important resource in many countries, much of the published information on Cryptosporidium in deer refers to studies on wild or free-ranging cervids (e.g. white-tailed deer in USA, roe deer in Spain, caribou in Canada, and moose, red deer, roe deer and reindeer in Norway; Rickard et al. 1999; Castro-Hermida et al. 2011b; Johnson et al 2010; Hamnes et al. 2002). While these studies on free-ranging cervids may give useful information regarding the species or genotype of Cryptosporidium that might infect farmed deer (C. ubiquitum, C. parvum), as farmed deer probably differ quite substantially from their wild counterparts regarding exposures and stresses, extrapolation of prevalence data from wild to farmed deer may give an incorrect picture. Indeed, a study in Poland found that the prevalence of Cryptosporidium was significantly higher in wild red deer than farmed red deer (27 % compared with 4.5 %), and mean oocyst concentration was also five times higher in faecal samples from wild red deer (Paziewska et al 2007). However as the sample size was relatively small (52 wild deer, 66 farmed deer) and from only single locations and as age and symptoms were not indicated, it is not possible to determine the reason for these differences.

The few studies on the prevalence of *Cryptosporidium* infection in different farmed or domesticated cervids are summarised in Table 4.9.

4.5.1.2 Association of Infection with Clinical Disease

The lack of surveys for *Cryptosporidium* infection in farmed deer is surprising, given the clear association of infection with clinical disease in farmed cervids. Some of the first published studies on *Cryptosporidium* infection in cervids are case reports of severe (high mortality) outbreaks among farmed red deer calves. In one outbreak in Scotland, UK among 82 artificially reared red deer calves, 56 developed cryptosporidiosis and 20 subsequently died; 80 % of the calves with diarrhoea and 50 % of apparently asymptomatic calves excreted oocysts and post mortem histopathological examination of the intestines demonstrated lesions similar to

le 4.5	Cryptospo	ridium in differe	ent species of d	Table 4.9 Cryptosporidium in different species of deer identified by microscopy	roscopy			
_	Location Host species Symptoms	Symptoms	Host age	Study design	Prevalence (% positive – no. positive/ no. examined)	Diagnostic technique	Molecular analyses	Reference
	Ireland Red deer	Asymptomatic Adult hinds and calve	Adult hinds and calves	Longitudinal – monthly samples	Hinds: 0–63 % according to month (overall prevalence: 39.3 % – 114/290) Calves: 40–100 % according to month (overall prevalence: 60 % – 21/35)	Water-ether sedimentation followed by sucrose flotation and IFAT	Not conducted	Skerret and Holland 2001
Poland	Red deer	Not stated	Not stated	Semi-longitudinal – seasonal sampling	Overall prevalence: 4.5 % – 3/66	Sheather's flotation followed by mZN and IFAT	Unsuccessful	Paziewska et al 2007
China	Red deer, Pere David's deer, Sika deer	Asymptomatic 4 months- 10 yeau r	4 months- 10 years	Cross-sectional study at 4 deer farms	Red deer: 0 (0/8) Pere David's deer: 0 (0/33) Sika deer: 2.4 (2/83)	Sheather's flotation followed by light microscopy	 1 sample C. ubiquitum – PCR and sequence analysis at 4 genes 	Wang et al. 2008a

those seen in other species (Tzipori et al. 1981a). Another outbreak among new-born red deer calves in the UK also resulted in high mortality, with calves dying at 24–72 h of age. However, this outbreak was not characterized by diarrhoea, and terminal uraemia was proposed as the symptom leading to death (Simpson 1992). Outbreaks of cryptosporidiosis in red deer calves have also been reported from New Zealand, again with relatively high mortality (10 out of 10 calves dying within a few days of illness onset in one outbreak, and 7 out of 10 calves dying within 3 days of illness in another outbreak) (Orr et al. 1985). Severe subactute enteritis in the small and large intestine were reported in both outbreaks.

Information from other species of farmed deer is more scanty, but a retrospective study of neonatal mortality in farmed elk (Pople et al 2001) identified *Cryptosporidium* infection as one of the most important causes of enteritis leading to death (7 cases out of 11 of infectious enteritis from a total of 111 cases in which 46 had no specific cause of death identified). Among these 7 cases, 4 were associated with an outbreak on a single farm (Pople et al 2001).

Unfortunately, information on the species of *Cryptosporidium* infection associated with clinical disease in deer and *Cryptosporidium* associated with asymptomatic infection is lacking. One outbreak on a Scottish farm occurred when deer were put to graze on a pasture that had previously been grazed by a *Cryptosporidium*-infected herd of cattle (Angus 1988), and therefore it seems probable that this might indicate infection with *C. parvum*; it might be speculated that infection of deer with deer-adapted *C. ubiquitum* is less likely to cause severe symptoms.

4.5.1.3 Infection Dynamics: Oocyst Excretion and Transmission

Published information about infection dynamics in farmed deer is minimal. However, a longitudinal study in asymptomatic farmed red deer in Ireland (Skerrett and Holland 2001) provides some interesting data. Asymptomatic low-level (<10 OPG) oocyst shedding from adult hinds appeared to continue throughout the year, except during the calving season (May - June), when there was a 2-log increase in oocyst excretion rate; the highest level recorded was 67,590 OPG. The authors speculate that this may be related to hormonal or immunological changes, or perhaps alterations in stress levels. The authors note that this preparturient rise in oocyst shedding results in contamination of the environment for the new born calves. However, in this study, although calves became infected, oocyst excretion was low (not exceeding 150 OPG, and usually less), and clinical disease was not observed. Again, the species of *Cryptosporidium* in these infections is unknown.

4.5.1.4 Zoonotic Transmission

Both *C. parvum* and *C. ubiquitum* have zoonotic potential, but there appear to be no documented cases of proven zoonotic transmission from/to deer. Those studies

(USA and Australia) that have investigated deer as sources of contamination in watersheds, have focused on wild deer only and provided contrasting results (Cinque et al 2008; Jellison et al 2009).

4.5.2 Cryptosporidium in Farmed Camelids

Camelids are members of the family *Camelidae*, and include the tribe *Camelini* (including dromedaries and Bactrian camels), and the tribe *Lamini* (llamas, alpacas, vicuñas, and guanacos).

There are two species of camel. Approximately 14 million domesticated onehumped dromedaries (*Camelus dromedarius*) are found in Middle Eastern countries including the Sahel and Horn of Africa, as well as parts of Southern Asia where they provide people with milk, food, and transportation. Nearly two million domesticated two-humped bactrians (*Camelus bactrianus*) are native to the steppes of central Asia the Gobi and Taklamakan Deserts in Mongolia and China.

Alpacas (*Vicugna pacos*) and llamas (*Lama glama*) exist only in the domesticated state and are found worldwide. However, both are native to South America and are raised primarily for fibre production although llamas were once used extensively as work animals. The young of both are called crias.

4.5.2.1 Occurrence (Prevalence)

Camels and dromedaries: In the relatively few studies of dromedaries in Northern Africa and the Middle East, the prevalence of cryptosporidiosis varied greatly (Table 4.10). None of 23 camels from Iraq (Mahdi and Ali 1992) and none of 110 camels on farms in Tunisia were found positive for *Cryptosporidium* (Soltane et al. 2007). *Cryptosporidium* was detected in one of four camel calves in Egypt (Abou-Eisha 1994). However, in an abattoir in Yazd Province in Iran, microscopic examination of 300 faecal specimens detected 61 (20.3 %) positive for *Cryptosporidium* and 12 (12 %) positive abomasal mucosa specimens. At an abattoir in Isfahan Province in central Iran, of 63 adult male and 40 adult female dromedary camels examined, 39 (37.9 %) were *Cryptosporidium*-positive (Razawi et al. 2009). In northwestern Iran, of 170 faecal samples from camels 17 (10 %) were positive for *Cryptosporidium*-like oocysts (Yakhchali and Moradi 2012). The prevalence was significantly higher (20 %) in calves less than a year old.

Oocysts have also been recovered from wild and zoo-housed camels. Faeces from a 3-year-old Bactrian camel in the Wild Animals Rescue Centre of Henan Province in China were found positive for *C. andersoni* (Wang et al. 2008b). Oocysts from a zoo-housed Bactrian camel (Fayer et al. 1991) were infectious for mice (Anderson 1991) and were identified as *C. muris* (Xiao et al. 1999; Morgan et al. 2000); those from camels in the Czech Republic were identified as *C. andersoni*. Other zoo-housed

Location	Host	Age	No. infected/ no. examined	Method	Reference
Basrah, Iraq	Camels	Adults	0/24	Microscopy	Mahdi and Ali 1992
Australia	Camel	Not stated	1	Molecular	Morgan et al. 2000
Tunisia	Camels	Not stated	0/110	Microscopy	Soltane et al. 2007
China	Bactrian camel	3-year-old	1	Molecular	Wang et al. 2008b
Isfahan Province, Iran abattoir	Camels	2-14 years	39/103	Microscopy	Razawi et al. 2009
NW Iran	Dromedary camels	Calves and adults	17/170	Microscopy	Yakhchali and Moradi 2012
Yazd Province, Iran abattoir	Camels	<5–>10 years	61/300 faeces 12/100 abomasums	Microscopy	Sazmand et al. 2012

 Table 4.10 Cryptosporidium detected in camels

camels have been found to be infected with *Cryptosporidium* (Abou-Eisha 1994; Gomez et al. 2000; Gracenea et al. 2002).

Alpacas and llamas Cryptosporidium oocysts have been detected in both these species. However, in California none of 354 llamas from 33 facilities were found positive (Rulofson et al. 2001) nor were 61 alpacas on two farms in Maryland (Trout et al. 2008). Elsewhere in North America, Europe, and Australia small numbers of alpacas, llamas and guanacos have been examined and a few have been found positive for Cryptosporidium (Table 4.11). Most examinations were conducted by microscopy, but those that utilized molecular methods identified only C. parvum (Morgan et al. 1998; Starkey et al. 2007; O'Brien et al. 2008; Twomey et al. 2008). The exception is a study in which a cria was found infected with C. ubiquitum (Gomez-Couso et al. 2012). A national survey of 5,163 1-15-day-old alpacas in 105 Andean herds in Peru, the natural habitat for nearly 80 % of the world's alpacas, found 2 % of the youngest alpacas increasing to 20 % of the oldest alpacas, infected with Cryptosporidium spp., with an overall prevalence of 13 % (Lopez-Urbina et al. 2009). More recently in Peru, 4.4 % of 274 alpacas from 12 herds were found positive for *Cryptosporidium* spp. (Gomez-Couso et al. 2012). Herd prevalence was 58.3 % (7/12 herds) for Cryptosporidium. The highest prevalence (20 %) was found in the 8-week-old group (Gomez-Couso et al. 2012).

4.5.2.2 Association of Infection with Clinical Disease

Camels and dromedaries Few data are available on the subject of clinical illness associated with cryptosporidiosis in camels. Of 170 faecal samples, 17 camels (10 %) were positive for *Cryptosporidium*-like organisms (Yakchali and Moradi 2012). The prevalence was significantly higher in camel calves (<1 years old) (20 %) than other age groups, in which the diarrhoeic calves had a prevalence of 16 %.

			No infected/	Detection	
Location	Host	Age	no. examined	method	Reference
Wisconsin, USA	Llama	Cria	1/1	Not stated	Hovda et al. 1990
Peru	Alpaca	Not given	1	Molecular	Spano et al. 1997
					Morgan et al. 1998
England	Alpaca	9–30 days old	3/3	Microscopy	Bidewell and Cattell 1998
California, USA	Llama	Crias and Adults	0/354	Microscopy	Rulofson et al 2001
Oregon, USA	Llama and Alpaca	31–210 days old	4/45	Microscopy	Cebra et al 2003
Czech Republic	Alpaca	Not given	1/1	Molecular	Ryan et al. 2003c
Canada	Alpaca	Crias	Multiple (number not given)	Microscopy	Shapiro et al. 2005
Scotland	Alpaca	Not given	Not given	Not stated	Stewart et al. 2005
New York, USA	Alpaca	Crias	8/total not given	Microscopy and Molecular	Starkey et al. 2007
Maryland, USA	Alpaca	10 weeks-10 years	0/61	Molecular	Trout et al. 2008
Ohio	Llama and Alpaca	7–100 days old	15/58	Not stated	Whitehead and Anderson 2008
SW England	Alpaca	Crias; Adults Crias	3/6; 0/24 5/14	Molecular	Twomey et al. 2008
Australia	Alpaca	Not given	1/1	Molecular	O'Brien et al.2008
Washington, USA	Alpaca	8-18 days old	20/20	Microscopy	Waitt et al. 2008
Peru	Alpaca	1–15 days old	24/241 666/5.163	Microscopy	Lopez-Urbina et al. 2009
Peru	Alpaca	≥5 weeks	12/274	Microscopy and Molecular	Gomez-Couso et al. 2012
New York and Pennsylvania, USA	Alpaca	Crias Adults	8/110 9/110	Microscopy	Burton et al. 2012

Table 4.11 Cryptosporidium in llamas and alpacas

Alpacas and llamas Not all alpacas and llamas infected with *Cryptosporidium* show clinical signs of infection. Of 110 healthy crias and their 110 dams 7 % and 8 %, respectively, were found excreting oocysts (Burton et al. 2012). Oocysts of *C. parvum* were detected in 4 of 14 faecal samples from healthy crias and in one sample from a cria with diarrhoea (Twomey et al. 2008).

Cryptosporidium was observed in a post-operative neonatal llama with diarrhoea, cachexia, dehydration and electrolyte abnormalities (Hovda et al. 1990). During 8 days that intravenous fluids and nutritional support were provided, these signs were not observed.

Of 20 *Cryptosporidium*-infected alpaca crias with diarrhoea, 15 exhibited weight loss and 5 had a poor appetite (Waitt et al. 2008). Most were 8–18 days old when examined. Additional potential gastrointestinal pathogens were found in 7 of these crias. Sixteen crias recovered after supportive therapy that included intravenous rehydration, with partial parenteral administration of nutrients, antimicrobials, oral nutrients, plasma, insulin and other palliative treatments.

Additional reports of diarrhoea associated with cryptosporidiosis have been reported in alpaca and llama crias (Cebra et al. 2003; Shapiro et al. 2005; Whitehead and Anderson 2006; Starkey et al. 2007). Three fatal cases (2 with diarrhoea) of cryptosporidiosis were reported in alpaca crias less than 30 days of age (Bidewell and Cantell 1998). At necropsy, intestinal congestion and distension were noted, oocysts were detected in Ziehl-Neelsen stained smears, and no other significant organisms or toxins were detected.

In South America, llama and alpaca husbandry is a vital economic activity and neonatal diarrhoea syndrome (NDS) is the most common and costly enteric disease in newborn llamas and alpacas (Lopez-Urbina et al. 2009). However, the role of cryptosporidiosis in NDS has not been clearly identified.

4.5.2.3 Zoonotic Transmission

Camels and dromedaries Only rare circumstantial data of zoonoses are available and the link is very tenuous. In Yazd Province in Iran, 24 of 100 people in long-term contact with camels were found infected with *Cryptosporidium* spp. (Sazmand et al. 2012). Infection was higher in winter than summer (16/50 compared with 8/50).

Alpacas and llamas In New York, *Cryptosporidium parvum* infection was identified in 5 crias, 3 of their caretakers were confirmed to have cryptosporidiosis, and three others were suspected to have cryptosporidiosis, suggesting zoonotic transmission (Starkey et al. 2007).

4.5.3 Cryptosporidium in Farmed Rabbits

Rabbit farming (cuniculture) for meat, wool, and fur production occurs in a variety of settings around the world, and mostly involves the European (or common) rabbit (*Oryctolagus cuniculus*). Small-scale backyard cuniculture is common in many

countries (especially in Africa and South America), but commercial operations on a larger scale are found in Europe (particularly Italy, Spain and France) and Asia (particularly China and Indonesia). In the EU, rabbit meat production was estimated to be around 520,000 tonnes carcass-weight equivalent in 2005 (EFSA-AHAW 2005). In addition, rabbits continue to be bred for biomedical purposes – but this type of rabbit breeding will not be considered further in this chapter. Production and consumption of rabbit meat is relatively low in North America. Different rabbit breeds are used for meat, wool, and fur – with the most commonly used meat breeds being New Zealand, Californian, Florida White and Altex, all having good growth rates and desirable reproductive characteristics.

Much of the information presented in this section is derived from a comprehensive review article from 2010 (Robinson and Chalmers 2010).

4.5.3.1 Occurrence (Prevalence)

The majority of published prevalence information on *Cryptosporidium* in rabbits refers to studies on wild rabbits. Nevertheless, there have been several studies on the occurrence of infection in farmed rabbits and also in laboratory rabbits. The majority of these studies (involving both wild and domestic rabbits) are summarised in Robinson and Chalmers (2010). In Table 4.12, selected prevalence studies (rather than case reports) from farmed rabbits only are summarized, including two recent studies from China. Additionally, a further three studies from China and referenced in Zhang et al. (2012) are not included in Table 4.12 due to inaccessibility of the original publications. Zhang et al. (2012) do not provide any details on these studies and it is not certain that they refer to farmed rabbits. Although some surveys refer to *Cryptosporidium parvum*, all those studies in which genotyping has been used (including from wild rabbits; e.g. Nolan et al. 2010) suggest that the majority of natural infections in rabbits, if not all, are caused by *C. cuniculus*. Nevertheless, experimental infections with other species of *Cryptosporidium* have been established in rabbits, as summarised by Robinson and Chalmers (2010).

4.5.3.2 Association of Infection with Clinical Disease

Although the majority of surveys do not report symptoms associated with cryptosporidiosis in rabbits, experimental infections in preweaned rabbits have been associated with diarrhoea and high mortality (e.g. as reported by Robinson and Chalmers 2010; Mosier et al 1997) and also as described by Pavlásek et al. (1996) in farmed rabbits. However, even asymptomatic infection may result in some pathology, as noted by Inman and Takeuchi (1979), who reported blunted villi, a decrease in villus-crypt ratio, and mild oedema in the lamina propria in an apparently asymptomatic adult rabbit. Thus, even asymptomatic infection may reduce stock productivity.

					Prevalence (% positive –			
Location	Breed	Symptoms	Host age	Study design	no. positive/no. examined)	Diagnostic technique Molecular analyses	Molecular analyses	Reference
Czech Republic	Broiler rabbits of 6 different breeds or crossbreeds	of Variable, including diarrhoea s and inappe- tence. Seven deaths recorded	23–33 days and 82–92 days (all post- weaning)	Longitudinal – with pooled samples	Variable throughout study, but at peak pooled samples from 12/28 cages	Giemsa staining in faeces; post mortem examination of digesta and intestinal scrapings. Histology	Not conducted	Pavlásek et al. 1996
Tunisia	Not stated	Not stated	Not stated	Cross-sec- tional at 1 farm	Overall preva- lence: 0 % – 0/178	Formol-ether sedi- mentation followed by mZN	Not applicable	Soltane et al 2007
China (Henan Province)	Various, includ- ing Standard Rex and New Zealand White	Asymptomatic	5 age groups: <1 month, 1–3 months,	Cross-sec- tional at 8 farms	Overall prevalen ce: 3.4 % – -37/1,081 <1 month: 4.1 %-3/73 1-3 months: 10.9 %-27/	Sheather's flotation followed by modified acid- fast stain	All (36/37successful) <i>C. cuniculus^a</i> – PCR and RFLP and sequence analysis at 18S rRNA gene. 8 samples further analysed at 3 other genes	Shi et al. 2010
			4–6 months, 7–12 months, >12 months		247 246 months: 1.3 %-6/474 7-12 months: 0.4 %-1/230 >12 months: 0 %-0/57		 GP60 subtyping – 30 of 37 successful: VbA29 (18 samples), VbA35 (4 samples), VbA36 (8 samples) 	
China (Heilongj- iang province	Not stated	Not stated	4-6 months	Cross-sec- tional at 8 farms	Overall preva- lence: 2.4 %–9/378 (positive samples from 4 farms only)	Sheather's flotation followed by bright-field microscopy	All (9/9successful) C. cuniculus – PCR and sequence analysis at 18S rRNA gene GP60 subtyping – 9 of 9 suc- cessful; VbA21 (6 samples), VbA32 (3 samples)	Zhang et al 2012

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 Table 4.12
 Cryptosporidium in farmed rabbits

^aDescribed as rabbit genotype in publication

Although no outbreaks of cryptosporidiosis in rabbit farms have been documented in the literature, acute outbreaks of diarrhoea with high mortality rates are frequently observed in rabbits (Banerjee et al 1987). Although bacterial agents are frequently considered to be the aetiological agent, it seems probable that some may be due to undiagnosed cryptosporidiosis. For example, the parasitological techniques (direct microscopy and flotation) used for investigating epizootic outbreaks of diarrhoea, characterized by a high morbidity and mortality, in different commercial rabbit farms in Mexico (Rodríguez-De Lara et al 2008) may have been insufficient for detecting *Cryptosporidium* infection, particularly if the operators had little experience in diagnosing this infection.

4.5.3.3 Infection Dynamics: Oocyst Excretion and Transmission

Information on the dynamics of *Cryptosporidium* infection in farmed rabbits is mostly lacking, although low oocyst excretion rates were reported in the majority of studies on rabbits in general (not just farmed rabbits). The studies from the Czech Republic provide some data, but, as the animals were not sampled individually, the data are difficult to interpret, and suggest that the source of infection for young rabbits may be low-level excretion of oocysts from mother rabbits at around parturition (Pavlásek et al. 1996).

4.5.3.4 Zoonotic Transmission

C. cuniculus is rarely, but sporadically, identified in human infections. In 3030 *Cryptosporidium*-positive faecal samples submitted for routine typing in UK between 2007 and 2008, 37 (1.2 %) were identified as *C. cuniculus*, with both GP60 Va and Vb subtype families detected (Chalmers et al 2011). However, the greatest evidence for *C. cuniculus* from rabbits having a significant zoonotic potential came from a waterborne outbreak of cryptosporidiosis in England in 2008 affecting 29 people; *C. cuniculus*, subtype VaA18 was identified in eight patients, a water sample from the implicated supply, and from the colon of a carcass of a rabbit (presumably wild) that was found in a tank at the water treatment works (Chalmers et al 2009). Nevertheless, transmission of *Cryptosporidium* to humans from farmed rabbits has not been recorded, and an investigation exploring associations between farm animals and human patients with cryptosporidiosis did not implicate rabbits as a source of infection (Smith et al 2010).

4.6 Cryptosporidium spp. in Poultry

The world stock of birds in production in 2011 was estimated to 22×10^9 animals (FAOSTAT 2013b). Approximately 56 % of the world stock was found in Asia, whereas Europe, North America and South America had approximately 10–11 %

each of the population. The largest group was chickens, with 90 % of the total stock. Ducks, turkeys and geese/guinea fowls constituted 6.1 %, 2.1 % and 1.7 % respectively, and other birds (ratites, pigeons etc.) only constituted 0.1 % of the world stock. The main chicken, duck and goose/guinea fowl production is in Asia (54 %, 90 % and 91 % within each group respectively), and most of the turkey production in North America (54 %), followed by Europe (23 %). For other birds, 50 % of the reported production was located in Asia, 41 % in Africa and 9 % in Europe.

Chickens (*Gallus gallus domesticus*) are descendants of the Red jungle fowl (*Gallus gallus*), with some hybridization with the Grey junglefowl (*G. sonneratii*). Broilers are usually kept in intense systems and reach slaughter size at about 6 weeks of age. Organically bred broilers and broilers kept on free range grow a bit more slowly. Laying hens can produce over 300 eggs in their first production year, but after that production declines rapidly.

Domesticated ducks (*Anas platyrhynchos domesticus*) are, except for the Moscovy duck (*Cairina moschata*), descendants of the Mallard (*Anas platyrhynchos*). The majority of domesticated geese (*Anser anser domesticus*) descend from the Greylag goose (*Anser anser*), but the breeds Chinese goose and African goose are derived from the Swan goose (*Anser cygnoides*). Ducks and geese are bred for meat, eggs and down, and ducks, to a lesser degree, also for the production of foie gras.

The domestic turkey (*Meleagris gallopavo*) is a progeny of the wild turkey, which is found in the wild in the United States http://www.turkeyfed.com.au/Turkey_Info. php. Turkeys are bred for meat production. The breed used is the white broad breasted turkey, introduced into commercial production in the 1950s http://bizfil.com/turkey-raising-primer. As with commercial chicken broiler farming, turkey farming is intense. The poults are extremely fast-growing, and reach approximately 6 kg at 10 weeks of age if given proper nutrition http://bizfil.com/turkey-raising-primer/. The United States has the highest consumption of turkey meat per person, and they are also the largest turkey producer, with 7.32 billion pounds of turkey meat produced in 2011 http://www.agmrc.org/commodities_products/livestock/poultry/turkey.

Among ratites, mainly ostriches (*Struthio camelus*) are farmed, but rheas (*Rhea americana*) and emus (*Dromaius novaehollandiae*) are also kept for production. Ratites are bred for meat, egg, and feather and leather production. Farming for feather production began already in the nineteenth century. Partridges, such as the Grey or English partridge (*Perdix perdix*) and red-legged partridge (*Alectoris rufa*), are gallinaceous birds used as game, and have been introduced in different parts of the world for this purpose. Another gallinaceous bird is the helmeted guinea fowl (*Numida meleagris*). They are used for pest control, eating ticks and other insects, and can be kept as an alarm system among other domesticated birds due to their loud and shrieking warning call. The meat is considered a delicacy. The Japanese quail (*Coturnix japonica*) is bred for meat and eggs. Domestic pigeons (*Columba livia domestica*) are the progeny of the world's oldest domesticated bird, the Rock pigeon. Pigeons are bred for meat, sporting competitions, homing, as exhibition birds or pets.

Cryptosporidium infection has been associated with large morbidity and mortality in different bird species (Bezuidenhout et al. 1993; Hoerr et al. 1986; Pages-Mante et al. 2007; Penrith et al. 1994; Ritter et al. 1986; Santos et al. 2005) and can thus be of great economic importance.

4.6.1 Prevalence

Avian cryptosporidiosis was first described in chickens (Tyzzer 1929). The infection was subclinical and situated in the caecum. Invasive stages looked identical to those of *C. parvum*, but no oocyst description was made, and no name was proposed. Today, three valid species have been identified in poultry. In addition, five genotypes have been identified in wild ducks and geese, and five additional genotypes have been described from other birds.

The *Cryptosporidium* oocysts identified by Slavin in 1955 were morphologically similar to *C. parvum*, described in mice in 1912 (Tyzzer 1912), and the infection site was the distal ileum. Slavin identified this bird *Cryptosporidium* as a unique species, *C. meleagridis*. When molecular methods were introduced as a means of species determination, it was verified that *C. parvum* and *C. meleagridis* were indeed different species (Sreter et al. 2000).

A species with a larger oocyst, first identified in chickens, and infecting the intestine, bursa and cloaca, was described and named *C. baileyi* (Current et al. 1986). This species is also involved in respiratory cryptosporidiosis, infecting the epithelium of sinuses, air sacs, nasopharynx, trachea and bronchi (Itakura et al. 1984; Lindsay et al. 1987). Infection of the conjunctiva (Chvala et al. 2006) and urinary tract, including the kidneys has also been shown (Abbassi et al. 1999; Trampel et al. 2000).

A third species, *C. galli*, infecting the proventriculus of chickens, was described by Pavlásek in 1999 and 2001, and re-described in 2003 (Pavlásek 1999, 2001; Ryan et al. 2003b). The species was probably described in finches already in 1990 (Blagburn et al. 1990) and later the name *C. blagburni* was proposed (Morgan et al. 2001). However, molecular analyses have shown that *C. blagburni* is the same species already described as *C. galli*, and thus the latter is considered to be the valid species name.

In addition, isolates referred to as goose genotypes I-IV have been identified in Canada geese and a duck genotype has been described in a Black duck and Canada geese. Of the other genotypes described in birds (avian genotypes I-IV and the Eurasian woodcock genotype), avian genotype II has been detected in ostriches.

Two proposed species are today considered as *nomen nudum* due to lack of sufficient data. *Cryptosporidium tyzzeri* in chickens was described in 1961 (Levine 1961) and later *C. anserinum*, found in the large intestine of geese was described (Proctor and Kemp 1974).

Based on 18S rDNA phylogeny, C. galli and the woodcock genotype belong to the clade of gastric cryptosporidia together with C. andersoni, C. muris and

C. serpentis, whereas *C. meleagridis*, *C. baileyi*, goose genotypes I and II and the duck genotype belong to the intestinal clade (Xiao et al. 2004). *Cryptosporidium meleagridis* is closely related to the group including *C. parvum* and *C. hominis*; *C. baileyi* is closely related to the snake genotype, goose genotypes I, II and the duck genotype cluster together and are closely related to *C. scrofarum*, *C. bovis*, *C. ryanae* and the deer genotype. Goose genotypes III-IV and avian genotypes I-IV were not included in the phylogenetic tree. In another publication, goose genotypes I (goose #1, 2, 3, 6 and 8), II (goose #9) and the duck genotype (goose #7) are more distant (Jellison et al. 2004). The avian genotypes are more scattered. Avian genotypes I and II belong to the intestinal clade and are closely related to *C. baileyi*. Genotypes III and IV belong to the gastric clade, where genotype III is closely related to the Eurasian woodcock genotype and *C. serpentis*, and genotype IV is closely related to *C. galli* (Ng et al. 2006).

Prevalence data based on fecal examination could be affected by the time from sampling to analysis. This has been observed when oocyst numbers in chicken faeces dropped to approximately one third in samples stored for a week from first to second analysis, and where first analysis was performed on the day after sampling (C. Axén, unpublished data). It is possible that oocysts die and are quickly degraded by detrimental effects (extreme pH) due to the high ammonium content of bird droppings.

4.6.1.1 Chickens

A flock prevalence of 41 % (23/56), with 10–60 % within-flock prevalence, was reported for *C. baileyi* respiratory infection in broilers in the USA (Goodwin et al. 1996). In Morocco, *Cryptosporidium* sp. were found in 14 (37 %) of 38 investigated flocks. Within-flock prevalence ranged from 14 % to 100 %, and the highest prevalence (52 %) was identified in broilers aged 36–45 days, with no infection prior to 25 days of age (Kichou et al. 1996). Diagnosis was based on histopathology.

An overall *Cryptosporidium* prevalence of 10.6 % for layer chickens and 3.4 % for broiler chickens was shown in a study of faecal samples from 2015 birds in China (Wang et al. 2010a). The highest prevalence (24.6 %) was found in 31–60-day old laying chickens, whereas prevalences in broiler chickens never exceeded 5 %. DNA analysis identified *C. baileyi* as the major species, with 92/95 investigated samples, and only 3 samples were positive for *C. meleagridis* (Wang et al. 2010a). In contrast, another recent study identified *C. meleagridis* as the major species in chickens (Baroudi et al. 2013). The overall *Cryptosporidium* prevalence was 34.4 % by histopathology, and the highest prevalence (46.2 %) was identified in 16–30-day-old chickens, which is in line with the results from Kichou et al. (1996) and Wang et al. (2010a). The majority of the birds were infected with *C. meleagridis* only (n = 25). *Cryptosporidium baileyi*

only was detected in four birds and a mixed *C. meleagridis/C. baileyi* infection was found in one bird. However, these chickens had died from diarrhoea, which could affect the outcome regarding *Cryptosporidium* sp.

4.6.1.2 Turkeys

A morbidity of 5–10 % due to sinusitis was reported for a flock where Cryptosporidium sp. could be isolated from diseased poults (Glisson et al. 1984). It was stated that macroscopic *post mortem* examination of the infraorbital sinuses of healthy birds was normal compared with those of diseased birds, but it was not clearly stated whether *Cryptosporidium* sp. was also identified in the healthy birds and thus the infection prevalence cannot be estimated. Goodwin et al (1988b) identified invasive Cryptosporidium stages in turkey poults form a farm where the poults suffered from self-limiting diarrheal of unknown aetiology, but no prevalence estimation was given. Prevalences of 80 % in 17-day-old poults, 38 % in 24-day-old and 0 % in \geq 60-day-old poults was found by Woodmansee et al. (1988). Oocysts were identified as C. meleagridis based on morphology and infection site. A 35.5 % (17/60) prevalence in diarrhoeic or just unthrifty poults was reported in Iran (Gharagozlou et al. 2006). Prevalence was based on histological examination of intestinal, bursal and cloacal tissues. Examination of faeces revealed that only 29 % of the infected birds shed oocysts. Infection was identified in 1–7-week-old poults, whereas the 43 uninfected poults all were older than 7 weeks. DNA analysis was not performed and oocyst size was not stated in the publication, but based on infection site, host species and symptoms the authors suggested that C. meleagridis was the species responsible. A 10.0 %, 10.5 % and 2.5 % pre-slaughter prevalence respectively (age 4-9 weeks) was detected upon faecal examination of three flocks from the same farm (McEvoy and Giddings 2009). One of 59 turkeys was positive at post-slaughter examination (age 14 weeks). Upon DNA analysis, all six positive samples were identified as C. parvum. In a recent study, a 43.9 % prevalence of C. meleagridis was shown in deceased turkeys, with the highest prevalence (57.9 %) in poults aged 16–30 days (Baroudi et al. 2013).

4.6.1.3 Ducks and Geese

In one study, 73 (57 %) of 128 ducklings and 44 (59 %) of goslings aged 8–35 days were infected with *Cryptosporidium* (Richter et al. 1994). Infection was present in both intestinal and respiratory tract, but oocyst morphology was not described.

In a study on experimental infection with Usutu virus in geese, *Cryptosporidium* developmental stages in tissue samples were an accidental finding. This was further investigated by *in situ*-hybridization, and *Cryptosporidium* infection was detected in 89 % of conjunctival tissue samples and 88 % of bursal tissue samples. DNA analysis revealed presence of *C. baileyi* (Chvala et al. 2006). *C. baileyi* was also identified in two ducks in Rio de Janeiro (Huber et al. 2007).

4.6.1.4 Other Birds

Ratites *Cryptosporidium* infection in ostriches was first described in the early 1990s (Allwright and Wessels 1993; Bezuidenhout et al. 1993; Gajadhar 1993, 1994; Penrith et al. 1994; Penrith and Burger 1993). Infection was first identified in faecal samples from 14 (8.5 %) of 165 ostriches imported from Africa to Canada (Gajadhar 1993). Penrith and Burger (1993) identified invasive stages in a section of the small intestine of a 4-week old chick that has suffered from rectal prolapse, and Allwright and Wessels (1993) identified *Cryptosporidium* in histology sections of the bursa, intestine and pancreatic ducts. In 1994, Gajadhar et al. characterized the isolated oocysts and investigated host specificity. The oocysts were morphologically similar to those of *C. meleagridis*, but attempts to infect suckling mice, chickens, turkeys and quail failed, indicating that this was probably another species. In addition, only faecal samples were investigated, so the infection site was not determined (Gajadhar 1994). As this study was conducted before molecular tools were commonly used for *Cryptosporidium* species determination, the true identity of this isolate will remain unknown.

A low prevalence, with only 2 (0.6 %) of 336 investigated samples from ostriches aged 2 months-5 years being Cryptosporidium positive, was found in Greece (Ponce Gordo et al. 2002). Oocysts were of two sizes, $3.8 \times 3.8 \ \mu m$ and $5.7 \times 4.8 \ \mu\text{m}$, indicating the presence of two different species. In contrast, in a Spanish study a 60 % Cryptosporidium prevalence in adult rheas and ostriches was found (Ponce Gordo et al. 2002). The authors reported an oocyst diameter of 3–5 µm, which is similar to the description provided by Gajadhar (1994). Molecular analysis of the isolates was not performed. Oliveira et al. (2008) found 44 % prevalence in 77 ostriches based on microscopy. Oocysts were generally morphologically similar to C. baileyi and Cryptosporidium avian genotype II (Ryan and Xiao 2008). However, the morphometric variation was so large that the authors suggested that more than one species had been identified (Oliveira et al. 2008), but this was not verified by molecular analysis. An isolate similar to C. baileyi in both oocyst morphology and PCR-RFLP banding pattern was described from Brazilian ostriches (Santos et al. 2005). The isolate was characterized as a sister taxon to C. baileyi by sequence analysis of the 18S rDNA, HSP70 and actin genes (Meireles et al. 2006), and was named *Cryptosporidium* avian genotype II by another research group (Ng et al. 2006). Experimental infection (oral or intratracheal) with the Brazilian isolate in chickens failed (Meireles et al. 2006). The avian genotype II has also been identified in Vietnam. On a single ostrich farm 110 (23.7 %) of 464 samples were positive for Cryptosporidium oocysts. The highest prevalence as well as the highest shedding intensity (35.2 %) was found in 2-3 month-old animals. Of 17 samples used for molecular characterization, all were found to be avian genotype II (Nguyen et al. 2013).

Wang et al (2011b) reported *Cryptosporidium* infection in 53 (11.7 %) of 452 investigated ostrich samples. Prevalence peaked at the age of 4–8 weeks with 16.2 %. No infection was detected in birds younger than 1 week or older than 12 months. Molecular analysis of positive samples identified only *C. baileyi*.

Quails and partridges Enteric cryptosporidiosis in quails, with oocysts similar to *C. meleagridis*, was first described in 1986 (Hoerr et al. 1986; Ritter et al. 1986). Early attempts at experimental infection of quail with *C. baileyi* isolated from chickens failed (Current et al. 1986; Lindsay et al. 1986), but were later successful (Cardozo et al. 2005). Natural infection was first documented in 2001 (Morgan et al. 2001). Since then, natural *C. baileyi* infection has been described in two reports (Murakami et al. 2002; Wang et al. 2012). One large survey of *Cryptosporidium* infection in quails was performed in China (Wang et al. 2012). Out of 1,818 faecal samples, 239 (13.1 %) from 29 (61.7 %) farms were positive. Infection was most common among 72–100-day old quails (23.6 %). DNA analysis revealed *C. baileyi* in 237 samples and *C. meleagridis* in two samples. One case of *Cryptosporidium* infection in partridges was described (Pages-Mante et al. 2007).

Pigeons There are a few reports of cryptosporidiosis in pigeons (Ozkul and Aydin 1994; Qi et al. 2011; Radfar et al. 2012; Rodriguez et al. 1997). Radfar et al. (2012) describe an overall prevalence of 2.9 % in 102 examined adult and nestling birds, with 3.4 % prevalence in adults and 2.3 % prevalence in nestlings. The other articles are case reports (Ozkul and Aydin 1994; Rodriguez et al. 1997) and a study on pet birds in general, where *C. meleagridis* was found in one pigeon (Qi et al. 2011).

4.6.2 Association of Infection with Clinical Disease

4.6.2.1 Chickens

Respiratory as well as intestinal and bursal *Cryptosporidium* infections cause disease in chickens, but infection without clinical symptoms has also been observed (Fletcher et al. 1975; Taylor et al. 1994).

In Spain, a 90 % morbidity due to respiratory infection in one flock was caused by *Cryptosporidium* sp. Weekly mortality rates were 0.9–1.5 % (Fernandez et al. 1990). Infection was detected in the trachea and oesophagus. In another flock investigated in the same study, weight loss was the primary symptom, and bursal cryptosporidiosis was diagnosed (Fernandez et al. 1990). Goodwin et al. (1996) found a correlation between *C. baileyi* infection of the trachea and severity of tracheitis symptoms, airsacculitis and condemnation of birds.

In respiratory cryptosporidiosis, co-infection with other pathogens has been identified in a number of studies. *Cryptosporidium* sp. and concurrent adenovirus infection was identified in a large broiler flock with respiratory disease (Dhillon et al. 1981). In a retrospective study on *post mortem* diagnoses of respiratory cryptosporidiosis, it was found that co-infection with virus or bacteria was common (Goodwin et al. 1988a). In another study, *Cryptosporidium* sp. and *Aspergillus* or bacteria were detected in the lungs of four layer chickens that died from pneumonia. *Cryptosporidium* were also found in the ureters and kidneys (Nakamura and Abe 1988). The effect of *Cryptosporidium* infection alone on development of clinical

symptoms in these cases cannot be estimated, but there is probably a synergistic effect of co-infections, increasing the severity. Such a synergistic effect of co-infection with infectious bronchitis virus or *Escherichia coli* has been reported (Blagburn et al. 1991).

Respiratory symptoms were reported from chickens that had been experimentally inoculated intra-tracheally with *C. baileyi*, whereas infection was successful but caused no symptoms in orally inoculated chickens (Lindsay et al. 1988).

C. meleagridis infection was associated with diarrhoea and mortality in one study of Algerian chickens (Baroudi et al. 2013). Experimental *C. meleagridis* infection of chickens has been observed to result in the chickens becoming indolent and having soiled feathers. Growth retardation was reported, but compensatory growth occurred after a few weeks (Tumova et al. 2002).

4.6.2.2 Turkeys

Turkey was the first animal species in which clinical cryptosporidiosis was described (Slavin 1955). Infection was associated with diarrhoea at 10–14 days of age, but other parasites (including *Histomonas*, *Trichomonas* and Strongylides) were also detected. Experimental infection (crop inoculation) with *C. meleagridis* produced infection of the ileum, caecum and bursa, but was not associated with clinical symptoms (Bermudez et al. 1988). The isolate used was from symptomatic poults; however, these were simultaneously infected with reovirus (causing enteritis and hepatitis). Co-infection with *Cryptosporidium* and reovirus in turkeys with enteritis and hepatitis, leading to increased mortality, has also been shown in another study (Wages and Ficken 1989). The presence of other pathogens in these studies could indicate a low to moderate primary pathogenicity of *C. meleagridis*. Self-limiting diarrhoea (moderate to severe in character), a slower growth rate and growth deformities were reported from one farm where diseased poults were diagnosed with *Cryptosporidium* infection (Goodwin et al. 1988b). Other pathogens were not excluded, as was also mentioned by the authors.

In dead poults that had suffered from depression and diarrhoea (faeces adhered on the hind part of the body), necropsy revealed lesions in the small intestine. Microscopic investigation identified *Cryptosporidium* sp. in the respiratory tract and kidneys, as well as in the gastrointestinal tract (Tacconi et al. 2001). Diarrhoea, emaciation, lethargy and reduced growth associated with natural *C. meleagridis* infection have been reported from Iran, but the presence of other pathogens was not excluded (Gharagozlou et al. 2006). Of 60 diarrhoeal and/or unthrifty birds, 17 (35.3 %) were identified as *Cryptosporidium* positive by histology, and *C. meleagridis* in 25 (44 %) of 57 examined turkeys that died from diarrhoea, but infection with other pathogens was not investigated.

Respiratory cryptosporidiosis has also been described in turkeys (Ranck and Hoerr 1987; Tarwid et al. 1985). Tarwid et al. (1985) identified *Cryptosporidium* sp. in necropsied birds from two outbreaks of colibacillosis. Colibacillosis is, according to

the authors, a secondary disease in turkeys, and *Cryptosporidium* sp. was identified as the primary pathogen. Symptoms were frothy conjunctivitis and increased mortality. Necropsy revealed pathological changes such as pericarditis, peritonitis and air-sacculitis in addition to the conjunctivitis that was observed in live birds.

Thirteen birds with respiratory disease were all positive for *Cryptosporidium* sp. by histology (Ranck and Hoerr 1987). Microscopy of sinus and/or tracheal exudates revealed oval oocysts in some samples, but oocyst size was not described, and both *C. baileyi* and *C. meleagridis* can appear oval (length/width ratios of 1.05–1.79 and 1.00–1.33 respectively (Ryan and Xiao 2008)). Symptoms such as coughing, rattling, sneezing, frothy eyes and swollen sinuses were reported. Other pathogens were present in all but two of the examined birds, and it is unclear whether the infection with *Cryptosporidium* played a primary role in the pathogenesis or not. Studies on cryptosporidiosis in turkeys, including clinical symptoms, are summarised in Table 4.13.

4.6.2.3 Ducks and Geese

Clinical cryptosporidiosis in ducks and geese seems to be less common and milder (see Table 4.14) than in other poultry. Only mild respiratory symptoms resulted from experimental *C. baileyi* infection (both oral and intratracheal inoculation) in ducks (Lindsay et al. 1989). Respiratory and intestinal infection occurred for both infection routes, but symptoms (sneezing, rales, mild dyspnea) were only present in animals infected by the intratracheal route. Mason (1986) described a case of conjunctival cryptosporidiosis. However, since only one of 97 affected ducks was *Cryptosporidium* positive, the author concluded that the parasite was not the cause of the disease. Similarly, no symptoms occurred in geese in which *Cryptosporidium* infection was detected in the conjunctivas and bursas (Chvala et al. 2006); and Richter et al. (1994) noted that enteritis and upper respiratory tract symptoms were equally present in infected and non-infected ducks and geese. Mortality was not increased in the positive flocks (Richter et al. 1994).

4.6.2.4 Other Birds

Ratites *Cryptosporidium* infection in ostrich chicks has been associated with cloacal and phallus prolapse, leading to high mortality (Bezuidenhout et al. 1993; Penrith et al. 1994; Santos et al. 2005). Bezuidenhout et al (1993) found that prolapsed cloacas were heavily infected, whereas Penrith et al. (1994) described heavy infection of both the bursa and cloaca in affected chicks, but healthy chicks were not infected. Santos et al (2005) also identified *Cryptosporidium* infection in the rectum, coprodeum, urodeum and bursa of two dead chicks with cloacal prolapse, both originating from a farm with high mortality rates in 7–30-day-old chicks. However, the authors did not associate the problems with the infection, since changed management practices decreased clinical symptoms and mortality,

l able 4.13	itudies on <i>Cryp</i>	lable 4.1.3 Studies on <i>Cryptosportatum</i> infection in turkeys	ska				
			Location of		Cryptosporidium		
Country	Age	Symptoms	parasites	Oocyst size	sp.	Diagnostic method	Reference
United Kingdom	10–14 days	Diarrhea, mortality	Distal jejunum, ileum	$4.5 imes 4.0 \ \mu m$	C. meleagridis	Feces (smears) histopathology	(Slavin 1955)
United	7 weeks	Sinusitis, serous	Infraorbital	Not stated	Cryptosporidium	Histopathology	(Glisson
States	1	conjunctivitis	sinuses				et al. 1984)
Canada	5 weeks	Bronchopneumonia, con- junctivitis, mortality	Trachea	Not stated	Cryptosporidium spp.	Histopathology	(Tarwid et al. 1985)
United States	2.5-11 weeks	Coughing, gasping, sneez- ing, rattling, sinusitis	Turbinates, sinuses, tra- chea, bronchi	Not stated	Cryptosporidium spp.	Exudate (smears) histopathology	(Ranck and Hoerr 1987)
United States ^a	5–26 days	None	Ileum, caecum, bursa	4.9 mu Ø	C. meleagridis	Feces (smears + auramine O) histopathology	(Bermudez et al. 1988)
United States	Not stated	Diarrhea, depression, growth retardation, abnormal feathers, mis- shapen bones	Mid - to distal small intestine	Not stated, developmental stages 2-4 µm Ø	C. meleagridis	Pathology, histopathology	(Goodwin et al. 1988b)
United States	25 days	Enteritis, depression, stunted growth, mortality	Ileum, ileo- caecal junction	5 µm	C. meleagridis	Feces (smears + auramine O) histopathology	(Wages and Ficken 1989)
Hungary ^a	1 weeks	None stated	Mainly small intestine ^b	$4.8 \times 4.2 \ \mu m$	C. meleagridis	Histopathology, 18S rDNA PCR +	(Sreter et al. 2000)
Italy	30 days	Diarrhea, depression, huddling	Ileum, caecal tonsil, caecum, rectum, bursa, duodenum, proventricu- lus, kidney, trachea, lung	4.5–5.0 µm Ø	C. meleagridis	mucosal scrapings (flotation + mZN) histopathology	(Tacconi et al. 2001)

Table 4.13 Studies on Crvotosnoridium infection in turkeys

Iran	1–7 weeks	diarrhea, emaciation, leth- Duodenum, argv. growth retardation ieiunum.	Duodenum, ieiunum.	Not stated, developmental	C meleagridis	Feces (flotation + (Gharagozlou mZN) pathol- et al. 2006)	(Gharagozlou et al. 2006)
		0				ogy, histopathology	
			colon, cloaca, bursa	•			
United States	4, 9, 18 weeks	None stated	unknown/ caecum ^c	Not stated	C. parvum	18S rDNA analysis (McEvoy and of fecal Giddings	(McEvoy and Giddings
						droppings and post-slaughter caecal content	2009)

^aExperimental infection ^bDetermined in chickens and mice, oocysts first passaged through turkey poults ^cOne caccum-positive

Table 4.14	Studies on Cr	Table 4.14 Studies on Cryptosporidium spp. in ducks and geese	pp. in ducks and ε	geese				
Country	Age	Species	Symptoms	Location of parasites	Oocyst size	Cryptosporidium sp.	Diagnostic method	Reference
Ducks United States ^b	4 days	Domestic ducks	None or mild respiratory disease	Trachea, Bursa	Not stated	C. baileyi	Histology	(Lindsay et al. 1987)
Germany	9–35 days	Peking ducks	Not stated	Bursa, cloaca, intestine, respiratory tract, conjunctiva	Not stated	Cryptosporidium spp.	Tissue scrapings + mZN, histology + IFA	(Richter et al. 1994)
Australia	Not stated	Black duck (wild)	Not stated	Not stated	Not stated	Cryptosporidium duck genotype	18S rDNA PCR + sequencing	(Morgan et al. (2001)
United States	Unknown	Wild ducks	Unknown	Intestine	Not stated	Cryptosporidium spp.	Faecal flotation + IFA, 18S rDNA PCR	(Kuhn et al. 2002)
Brazil	Not stated	Domestic ducks	Not stated	Not done	Not stated	C. baileyi	18S rDNA PCR-RFLP and sequencing	(Huber et al. 2007)
Geese United States	25 days	Domestic geese	Not stated	Large intestine	Not stated Schizont 3.6 µm Macrogamete 4.5 µm	Cryptosporidium anserium, nomen nudum	Histology	(Proctor and Kemp 1974)
Germany	8–35 days	Domestic geese (Dan- ish breed)	Not stated	Bursa, gastro- intestinal and respira- tory tract	Not stated	Cryptosporidium spp.	Tissue scrapings + mZN, his- tology + IFA	(Richter et al. 1994)

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8	Q	4)	(4)	
(Graczyk et al. 1998)	(Chvala et al. 2006)	(Jellison et al. 2004)	(Zhou et al. 2004)	
TRAP C2 and β-tubulin PCR + genotyping	Histology with in-situ hybridization, 18S rDNA PCR + sequencing	18S rDNA PCR and sequencing	18S rDNA PCR-RFLP and sequencing	
C. parvum ^c	C. baileyi	<i>Cryptosporidium</i> goose genotypes I, II, III. IV, V ^d	Cryptosporidium goose genotypes I, II, Cryptosporidium duck genotype, C. parvum ^c , C. hominis ^c	
Not done	Not done	Not done	Not done	ading
Not done	Bursa, conjunctiva	Not done	Not done	^a Experimental infection study for Usutu virus, <i>Cryptosporidium</i> accidental finding b Experimental infection ^b Experimental infection ^a Not named in this publication
Unknown	Not stated	Unknown	Unknown	us, <i>Cryptospor</i> t manifest infe
Canada geese	Domestic geese	Canada geese	Canada geese	Experimental infection study for Usutu virus, <i>Cryptosporidium</i> Experimental infection Finding reported as passage of oocysts, not manifest infection Not named in this publication
Unknown	16-36 days	Unknown	Unknown	Experimental infection study 1 DExperimental infection Frinding reported as passage of Not named in this publication
United States	Austria ^a	United States	United States	^a Experimental infection ^b Experimental infection ^c Finding reported as pas ^d Not named in this publ

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although *Cryptosporidium* infection was still present on the farm. Enteritis was indicated by the presence of intestinal invasive stages and rectal prolapse in one chick examined by Penrith and Burger (1993). Because diarrhoea was not reported it is unknown whether the prolapse was caused by intense bowel movements or something else. *Cryptosporidium* infection has also been associated with pancreatic necrosis (Allwright and Wessels 1993).

Quails and partridges *Cryptosporidium* infection has been shown in both diarrhoea and respiratory disease in quails (Guy et al. 1987; Hoerr et al. 1986; Murakami et al. 2002; Ritter et al. 1986). Hoerr et al. (1986) reported high mortality rates from 5 days of age in quails infected with *Cryptosporidium* sp., and with no bacterial or viral pathogens detected. Acute fatal diarrhoea with mortality rates of up to 45 % in 0–17-day-old birds was described by Ritter et al (1986). Reovirus was also detected in necropsied birds, but another study reported that experimental infection with reovirus alone did not produce diarrhoea, whereas infection with *Cryptosporidium* sp., either alone or simultaneously with reovirus, resulted in severe diarrhoea and mortality (Guy et al. 1987). A synergistic effect of co-infection was, however, shown, since oocyst shedding was higher and reovirus infection became systemic and liver necrosis occurred (Guy et al. 1987).

Muramaki et al. (2002) reported a daily mortality rate of 5.7 % in one farm, where birds suffered from upper respiratory tract disease and decreased egg production. Respiratory symptoms were head swelling, nasal discharge and increased lacrimation, and necropsy revealed sinusitis, airsacculitis and egg peritonitis. Co-infection of *Cryptosporidium* sp., *Mycoplasma gallisepticum* and other bacteria was shown. The authors concluded that *M. gallisepticum* was the primary pathogen, but that the mixed infections in conjunction with high ammonia concentrations in the air worsened the symptoms. The role of *Cryptosporidium* infection in respiratory disease in quails thus remains unclear. Wang et al. (2012) reported that no clinical symptoms were seen in 1,818 sampled quails, of which 239 were *Cryptosporidium* positive.

C. meleagridis was the only pathogen identified in an outbreak of diarrhoea and cough in red-legged partridge chicks (Pages-Mante et al. 2007). Morbidity rates were 60–70 % and mortality more than 50 %, indicating high pathogenicity. Invasive stages were identified in both the respiratory and intestinal tract, suggesting that not only *C. bailey* might be associated with respiratory avian cryptosporidiosis.

Pigeons Diarrhoea associated with cryptosporidiosis in pigeons has been described in four birds (Ozkul and Aydin 1994; Rodriguez et al. 1997). Rodriguez et al. (1997) described a 40 % morbidity of yellow watery diarrhoea, weight loss, dehydration and weakness in a farm with 280 pigeons. Mortality was 5 % and necropsy of three birds revealed invasive stages of *Cryptosporidium* in the small intestine, caecum, colon, cloaca, and bursa. No viruses or bacteria could be isolated. Ozkul and Aydin (1994) identified invasive stages in the small intestine of a pigeon that had been depressed and had evidence of diarrhoea in the form of faeces in its hind feathers.

4.6.3 Infection Dynamics: Oocyst Excretion and Transmission

Isolates of both *C. baileyi* and *C. meleagridis* derived from one domestic bird species have been successfully transmitted to other domestic birds (Current et al. 1986; Lindsay et al. 1987). *C. galli* has not been experimentally transmitted between different domestic birds, but has been shown in finches as well as chickens (Blagburn et al. 1990; Pavlásek 1999, 2001; Ryan et al. 2003b), and thus has the potential to infect different bird species.

4.6.3.1 Chickens

The prepatent period of *C. baileyi* is approximately 4–8 days (Hornok et al. 1998; Lindsay et al. 1988; Rhee et al. 1991; Tumova et al. 2002). However, in the first report on *C. baileyi* infection in chickens, a prepatent period of up to 24 days was described (Current et al. 1986). Older chicks have a slightly longer prepatent periods than younger ones (Lindsay et al. 1988; Rhee et al. 1991; Taylor et al. 1994; Tumova et al. 2002).

The patent period varies more. At oral inoculation of 2-day-old chicks, a patent period of 26 days was seen, whereas it was 11–15 days in chicks inoculated at 14 days of age, 11–12 days in chicks inoculated at 28 days of age and <7 days in chicks inoculated at 42 days of age (Lindsay et al. 1988). With intratracheal inoculation, the same authors described patent periods of 27, 11–19, 10–11 and <7 days in these age groups (Lindsay et al. 1988). Rhee et al. (1991) and Tumova et al. (2002) observed a mean patent period of approximately 14 days. Oocyst excretion peaked on day 12 and days 11–17 post inoculation, respectively (Rhee et al. 1991; Tumova et al. 2002). Taylor et al. (1994) showed shorter patent periods and lower total oocyst output in older than younger chickens. There was also an effect of infection dose, in that oocyst output was higher and declined more slowly with lower infection doses (Taylor et al. 1994). Similar observations were made for 1- and 9-week old chicks (Sreter et al. 1995). In that study, the mean patent period for 1-week old chicks was 32 days, but one chicken shed oocysts for 151 days.

C. meleagridis was shed in the facees on days 4-7 post infection in two chickens experimentally infected at 6 weeks of age (Woodmansee et al. 1988). Tumova et al. (2002) infected 7-day-old chicks. Oocysts first appeared 3 days later and the patent period lasted for 16–17 days. Shedding rates were significantly lower than in chicks inoculated with the same number of *C. baileyi* oocysts.

The prepatent and patent period of *C. galli* has been described to be 25 and 6 days respectively (Pavlásek 2001), but was later reported as unknown when *C. galli* was redescribed (Ryan et al. 2003b).

4.6.3.2 Turkeys

The prepatent period of *C. meleagridis* in turkeys inoculated at 7–11 days of age was 2–4 days (Bermudez et al. 1988; Sreter et al. 2000; Woodmansee et al. 1988).

Woodmansee et al. (1988) reported that oocysts were shed for only 4 days; Sreter et al. (2000) found the patent period to be 8–10 days, whereas Bermudez et al. (1988) reported oocyst shedding and invasive stages still being present at day 21 post inoculation. Oocyst shedding rates were moderate (Sreter et al. 2000), and low to moderate (Bermudez et al. 1988).

Experimental infection with *C. baileyi* induced mild infection of the bursa (Current et al. 1986). Lindsay et al. (1987) inoculated turkey poults via the intratracheal, oral and intracloacal route. All three experiments caused infection, but only poults inoculated via the trachea developed symptoms (Lindsay et al. 1987).

4.6.3.3 Ducks and Geese

Lindsay et al (1986) described a prepatent period of 5 days and a possible patent period of 9–10 days in experimentally infected Muscovy ducks, based on investigation of pooled faecal samples (Lindsay et al. 1986). Oocyst morphology was not described. The intestine, bursa and cloaca were positive for invasive stages, but these tissues can be infected by both *C. baileyi* and *C. meleagridis* (Table 4.14).

4.6.3.4 Other Birds

For quails, one study describes a prepatent period of 7 days and a patent period of 21 days for *C. baileyi* (Cardozo et al. 2005). Otherwise, no data are available.

4.6.4 Zoonotic Transmission

Only one of the species and genotypes commonly infecting birds – *Cryptosporidium meleagridis* – has, so far, proved to be important in human cryptosporidiosis. This species is the third most common species in human cryptosporidiosis worldwide. In the industrialised world, *C. meleagridis* infection is usually associated with cryptosporidiosis cases in travellers to Asia or Africa (Elwin et al. 2012; Insulander et al. 2013; Leoni et al. 2006), but autochthonous cases have also been described (Elwin et al. 2012; Leoni et al. 2006; Silverlås et al. 2012). Studies on *Cryptosporidium* prevalence and species distribution in humans in South America have identified *C. meleagridis* infection at about the same prevalence as *C. parvum* (Cama et al. 2003, 2007, 2008). Although this is a true zoonotic species, there is only one report in which the bird source has been identified, and in that case,

chickens and not turkeys were involved (Silverlås et al. 2012). It is not known whether anthroponotic transmission occurs with this species, but it has been indicated by the fact that not all *C. meleagridis*-infected patients in an epidemiological investigation had had bird or animal contact (Elwin et al. 2012). *C. meleagridis* has the potential to infect other mammalian species as well, and experimental infection of mice, rats, rabbits, pigs and calves has been reported (Akiyoshi et al. 2003; Darabus and Olariu 2003). Due to the wide host range and the close relationship of *C. meleagridis* to *C. parvum* and *C. hominis*, it has been proposed that this species originated as a mammalian *Cryptosporidium* species, and later adapted to birds (Xiao et al. 2002b, 2004).

One study has identified *C. parvum* in turkeys (McEvoy and Giddings 2009), indicating that this species could play a role in zoonotic transmission. However, only one of 59 birds post-slaughter was positive compared to 2.5-10.5 % of the 5-10-week younger poults, which means risk of transmission via contaminated meat should be very small. The higher prevalence in poults should not pose a risk as long as the flocks are closed to the public. The shedding intensity was not reported, but prevalence indicates that infection rather than just intestinal passage was present. Some studies have identified *C. parvum*, *C. hominis* and *C. hominis*-like isolates in Canada geese (Jellison et al. 2004, 2009; Zhou et al. 2004). The authors conclude that these findings are probably not associated with infection and parasite proliferation, but rather transient carriage. Nevertheless, this indicates that domesticated ducks and geese can potentially act as transmission vehicles for these species.

Infection with *C. baileyi* has been identified in one immunodeficient patient. Diagnosis was based on oocyst morphology and biology – experimental infection of mice failed whereas inoculated chickens developed infection of the intestine, bursa and trachea (Ditrich et al. 1991). Since this patient was immunodeficient and no other reports exist, this species should not be considered as a true zoonotic agent.

4.7 Conclusion

Ever since animals were first domesticated, and humans became dependent upon them for the commodities that they supply, particularly food and fibre, the infections that affect the health and productivity of livestock have been a concern. Cryptosporidiosis was first identified as a disease of veterinary significance in the 1950s (in turkeys) and then in the early 1970s in calves, but major interest in cryptosporidiosis only developed with the first report of a human cases later that decade, and the recognition that *Cryptosporidium* infection was also of medical importance. Since then our knowledge on the veterinary significance of *Cryptosporidium* infection has expanded enormously – particularly in the livestock sector most impacted by cryptosporidiosis – young calves. However, as demonstrated in this chapter, it should not be forgotten almost all farmed animals may be pathologically affected by at least one species of *Cryptosporidium*, often causing clinical disease that in some instances may be fatal. For some *Cryptosporidium* species in some farmed animal species, transmission may be anthropozoonotic.

Cryptosporidium is a hugely successful parasite, as demonstrated by its host range and wide geographic distribution, and its control has proved challenging. As long as humans raise and depend on animals, there will be a need to control the transmission of cryptosporidiosis amongst livestock species.

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Chapter 5 Cryptosporidiosis in Other Vertebrates

Martin Kváč, John McEvoy, Brianna Stenger, and Mark Clark

Abstract *Cryptosporidium* has adapted to a broad range of hosts in all major vertebrate classes, and the species associated with humans and livestock represent a small fraction of the diversity in the genus. This review focuses on *Cryptosporidium* and cryptosporidiosis in terrestrial vertebrates other than humans and livestock. As the known host range of *Cryptosporidium* continues to expand, major orders of amphibians (Anura), reptiles (Squamata and Testudines), avians (17 out of 26 orders), and mammals (18 out of 29 orders) are now represented. The greatest *Cryptosporidium* diversity appears to be in mammals, which may be an Artifact of undersampling in other classes, but more likely reflects a different mechanism of *Cryptosporidium* diversification in mammals relative to other classes.

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5.1 Introduction

The emergence of *Cryptosporidium* as a serious human pathogen in the 1980s spurred research efforts to understand the biology and ecology of this enigmatic parasite. While early studies were hindered by a lack of tools to type isolates, the widespread use of genotyping during the last 15 years has begun to uncover the enormous diversity that exists in the genus *Cryptosporidium*.

The major focus of *Cryptosporidium* research over the past 30 years has been the control of cryptosporidiosis in humans and livestock, and the primary motivation to study other vertebrates has been to understand the ecology of human cryptosporidiosis. Wildlife-associated cryptosporidia account for a relatively small but significant proportion of human cryptosporidiosis cases (Feltus et al. 2006; Robinson et al. 2008; Elwin et al. 2011), the recent emergence of C. cuniculus as a human pathogen illustrates the connection between wildlife and public health. Rabbitadapted C. cuniculus was not on the public health radar until the misadventures of a rabbit at a UK water treatment facility in 2008 resulted in a waterborne outbreak of cryptosporidiosis (Chalmers et al. 2009). Genetically, C. cuniculus is remarkably similar to the major human pathogen C. hominis, and the two cannot be differentiated by some routine molecular diagnostics (Robinson et al. 2010). However, prompted by the waterborne outbreak, samples from 3,030 sporadic cryptosporidiosis cases in the UK were reexamined using enhanced diagnostics, and C. cuniculus was detected in 1.2 % of samples (Elwin et al. 2012a). Apart from the obvious benefit of knowing the species, and hence the likely source of the Cryptosporidium causing human disease, these studies raise interesting and fundamental questions about the biology of Cryptosporidium; such as, how does Cryptosporidium diversify and how does diversification affect host specificity and disease potential? These questions can be addressed by examining Cryptosporidium diversity across the range of vertebrate hosts, not just humans and livestock.

This chapter addresses the occurrence of *Cryptosporidium* in vertebrate hosts other than fish (Chap. 1), humans (Chaps. 2 and 3), and livestock (Chap. 4). Information is organized by vertebrate class (Amphibia, Reptilia, Aves, and Mammalia), and by order within each class. Background information is provided on each order, including orders that do not currently contain identified *Cryptosporidium* hosts. It is hoped that this approach will facilitate the identification of knowledge gaps and will stimulate ideas for future research.

5.2 Cryptosporidium and Cryptosporidiosis of Amphibians

To date, *Cryptosporidium* has been detected in only one of the three orders of amphibians: the Anura. It has not been detected in the Caudata, an order that contains more than 550 species in ten families and approximately 30 genera of salamanders, or the Gymnophonia, which are snake-like amphibians with about

180 species in ten families and 35 genera. The Caudata require freshwater for reproduction and they consume invertebrates. Gymnophonia are tropical in distribution, have larvae that may only be partially aquatic, and typically inhabit loose soil layers.

5.2.1 Anura

Anura is the largest order of Amphibians, with over 6,000 species of frogs and toads in 33 families and over 65 genera. Frogs and toads require at least some water for reproduction, but they occupy a wide variety of terrestrial and freshwater habitats. There are no marine species, and only a few inhabit brackish waters. All Anurans consume invertebrate prey, lacking teeth or grinding organs in their digestive tract.

Cryptosporidium fragile, the only amphibian-adapted species described to date, was isolated from the doubtful toad (*Duttaphrynus melanostictus*) originating in Malaysia (Jirků et al. 2008). Oocysts of *C. fragile* measure 6.2 µm (5.5–7.0 µm) × 5.5 µm (5.0–6.5 µm) and are readily lysed in hypertonic solutions. *Cryptosporidium fragile* infects the gastric epithelium and clusters with gastric cryptosporidia in molecular phylogenies (Table 5.1 and Fig. 5.1). Other reports of *Cryptosporidium* and cryptosporidiosis in reptiles have been rare. Cryptosporidiosis was diagnosed in a captive South African clawed frog (*Xenopus laevis*) that was euthanized after becoming grossly emaciated from illness (Green et al. 2003). Endogenous *Cryptosporidium* stages were identified in the gastric mucosa, and oocysts were isolated from water in the infected frog's enclosure. *Cryptosporidium* oocysts also have been reported in a captive Bell's horned frog (*Ceratophrys ornata*) (Crawshaw and Mehren 1987).

5.3 Cryptosporidium and Cryptosporidiosis of Reptiles

Cryptosporidium has been detected in two of the four orders of reptiles: Squamata and Testudines. It has not yet been reported in Crocodylia, a small order of 25 species in three families and eight genera, or the Rhynchocephalia, which is represented by two living species: the Brother's Island tuatara (*Sphenodon guntheri*) and the northern tuatara (*Sphenodon punctatus*). Crocodylians, which include crocodiles and alligators, have a two-chambered stomach that is unique among reptiles. The first chamber contains stones and functions much like the avian gizzard, and the second chamber is extremely acidic to facilitate almost complete digestion of prey. Tuataras, which are found only in New Zealand, are similar to the Squamates, but have different dentition and skull characteristics. They feed primarily on invertebrates, but occasionally eat small vertebrates or eggs.

Class	Order	Common names/groups	Host species (scientific name) and GenBank accession numbers
Amphibia		Toad	Doubtful toad (Duttaphrynus melanostictus) [EU162751]
Reptilia	Squamata	Lizards	Central bearded dragon (<i>Pogona vitticeps</i>) [AY382169, AY382170]
			Desert monitor (Varanus griseus) [AF112573]
			Leopoard gecko (Eublepharis macularius) [AY120915, AY282714, EU553556]
			Savannah monitor (Varanus exanthematicus) [AF093500, AY282713]
			Schneiders skink (Eumeces schneideri) [AY282715]
			Veiled chameleon (Chamaeleo calyptratus) [EU553587]
		Snakes	Australian taipan (Oxyuranus scutellatus) [AF108866]
			Ball phyton (<i>Python regius</i>) [EU553589, EU553590]
			Boa constrictor (Boa constrictor) [AY268584]
			Corn snake (<i>Pantherophis guttatus guttatus</i>) [EF50204, AF151376]
			Japanese grass snake (Rhabdophis tigrinus) [AB222185]
			New Guinea boa snake (Candoia aspera) [AY120913]
	Testudines	Tortoises and	Marginated tortoise (Testudo marginata) [EF547155]
		turtles	Pancake tortoise (Malacochersus tornieri) [GQ504270]
			Russian tortoise (Agrionemys horsfieldii) [GQ504268]
_			Indian star tortoise (Geochelone elegans) [AY120914]

Table 5.1 GenBank accession numbers of representative *Cryptosporidium* small subunit rRNA gene sequences from amphibians and reptiles. A phylogeny constructed from sequences in this table is presented in Fig. 5.1

5.3.1 Squamata

The order Squamata – snakes and lizards – is the largest and most diverse reptilian order, with more than 9,000 species in 50 families. They occupy diverse habitats, though most species inhabit drier terrestrial areas. All are predators that consume a wide variety of prey. Many produce venom or toxins to assist in immobilization of prey, which are swallowed without mastication.

Two *Cryptosporidium* species have been described in the Squamata: *Cryptosporidium serpentis*, which infects the gastric epithelium, and *C. varanii* (syn. *C. saurophilum*), which is an intestinal species (Table 5.1 and Fig. 5.1).

Cryptosporidium serpentis, which was described by Brownstein et al. (1977) and later named by Levine (1980), can cause chronic, insidious, and often fatal hypertrophic gastritis in both immature and mature ophidian hosts (Brownstein et al. 1977; Cranfield and Graczyk 1994; Kimbell et al. 1999). Clinical signs include

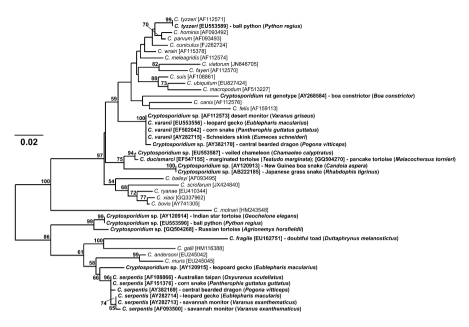


Fig. 5.1 A SSU rDNA-based maximum likelihood (GTRCAT model) tree of *Cryptosporidium* spp. sequences showing clades of cryptosporidia infecting amphibians and reptiles. *Cryptosporidium* sp. detected in amphibians and reptiles are in bold. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) is shown at nodes. Bootstrap values <50 are not shown. Interrupted branches have been shortened fivefold

mid-abdominal swelling and postprandial regurgitation, but not all cases are symptomatic (Brownstein et al. 1977; Cranfield and Graczyk 1994).

Although it is primarily a parasite of snakes (Table 5.2), *C. serpentis* also causes asymptomatic infections in a number of lizard species (Table 5.3). Unsuccessful attempts to experimentally infect Balb/c mice (Fayer et al. 1995), Pekin ducklings (*Anas platyrhynchos*) (Graczyk et al. 1998b), African clawed frogs (*Xenopus laevis*), and wood frogs (*Rana sylvatica*) (Graczyk et al. 1998c) suggest that *C. serpentis* is restricted to reptiles. However, strict reptile-specificity is brought into question by a recent report that *C. serpentis* was isolated from cattle in China, and was infectious for immunosuppressed and non-immunosuppressed Balb/c mice under experimental conditions (Chen and Qiu 2012).

Snakes are not susceptible to experimental infection with cryptosporidia from homothermic animals, including *C. andersoni* (calves), *C. baileyi* (chickens), *C. meleagridis* (turkeys), *C. muris* (mice), *C. muris*-like (Bactrian camels), and *C. wrairi* (guinea pigs), but they are susceptible to other reptilian cryptosporidia (Graczyk and Cranfield 1998). Reports of *C. tyzzeri* (previously mouse genotype I) in a black rat snake (*Elaphe obsoleta obsoleta*), boa constrictor (*Boa constrictor ortoni*), California king snake (*Lampropeltis getulus californiae*), corn snake (*Pantherophis guttatus*), emerald tree boa (*Corallus caninus*), fox snake

-	Host species		
Family	(scientific name)	Cryptosporidium taxa	Reference
Boidae	Amazon tree boa (Corallus hortulanus)	C. serpentis	(Morgan et al. 1999c; Xiao et al. 2004)
	Boa constrictor (Boa constrictor)	C. muris, C. serpentis, C. tyzzeri, C. varanii, rat genotype I	(Xiao et al. 2004; Pedraza-Diaz et al. 2009; Rinaldi et al. 2012)
	Brazilian rainbow boa (Epicrates cenchria)	C. serpentis	(Graczyk and Cranfield 2000; Sevá Ada et al. 2011b)
	Colombian rainbow (Epicrates cenchria maurus)	C. serpentis	(Ryan et al. 2003a)
	Emerald tree boa (Corallus caninus)	C. serpentis, C. tyzzeri	(Xiao et al. 2004)
	Green anaconda (Eunectes murinus)	C. serpentis	(Sevá Ada et al. 2011b)
	Kenyan or East African sand boa (Eryx colubrinus)	Cryptosporidium sp.	(Pedraza-Diaz et al. 2009)
	Madagascar tree boa (Sanzinia madagascariensis)	C. serpentis	(Levine 1980)
	New Guinean viper boa (Candoia aspera)	C. serpentis, C. varanii, Cryptosporidium sp.	(Graczyk and Cranfield 2000; Xiao et al. 2004)
	Rosy boa (Lichanura trivirgata)	C. serpentis	(Ryan et al. 2003a)
Colubridae	Black rat snake (Pantherophis obsoleta obsoleta)	C. muris, C. serpentis, C. tyzzeri, C. varanii, Crypto- sporidium sp.	(Cranfield and Graczyk 1994; Morgan et al. 1999c; Xiao et al. 2004)
	Brazos water snake (Nerodia harteri harteri)	Cryptosporidium sp.	(Upton et al. 1989)
	Bull snake (Pituophis melanoleucus melanoleucus)	C. serpentis, C. varanii	(Morgan et al. 1999c; Graczyk and Cranfield 2000; Xiao et al. 2004)
	California king snake (Lampropeltis getulus californiae)	C. serpentis, C. tyzzeri	(Xiao et al. 2004; Pedraza-Diaz et al. 2009)
	Common garter snake (Thamnophis sirtalis)	Cryptosporidium sp.	(Brower and Cranfield 2001)
	Corn snake (Pantherophis guttatus guttatus)	C. muris, C. serpentis, C. tyzzeri, C. varanii	(Levine 1980; Cranfield and Graczyk 1994; Kimbell et al. 1999; Morgan et al. 1999c; Graczyk and (continued)

 Table 5.2 Cryptosporidium spp. identified in snakes

(continued)

Table 5.2 (continued)

Family	Host species (scientific name)	Cryptosporidium taxa	Reference
			Cranfield 2000; Xiao et al. 2004; Plutzer and Karanis 2007; Pedraza-Diaz et al. 2009; Richter et al. 2011; Sevá Ada et al. 2011b; Rinaldi et al. 2012)
	Diamondback water snake (Nerodia rhombifer rhombifera)	Cryptosporidium sp.	(Upton et al. 1989)
	Eastern indigo snake (Drymarchon corais couperi)	Cryptosporidium sp.	(Cerveny et al. 2012)
	Fox snake (Pantherophis vulpina gloydi)	C. serpentis, C. tyzzeri	(Xiao et al. 2004)
	Jalisco milk snake (Lampropeltis triangulum arcifera)	Cryptosporidium sp.	(Upton et al. 1989)
	Louisiana pine snake (Pituophis ruthveni)	C. varanii	(Xiao et al. 2004)
	Milk snake (Lampropeltis triangulum)	C. serpentis, C. tyzzeri, C. varanii, Crypto- sporidium sp.	(Graczyk and Cranfield 2000; Xiao et al. 2004; Pedraza- Diaz et al. 2009; Sevá Ada et al. 2011b)
	Mexican black kingsnake (Lampropeltis getulus nigritus)	C. serpentis	(Pedraza-Diaz et al. 2009)
	Mexican kingsnake (Lampropeltis mexicana)	C. serpentis	(Sevá Ada et al. 2011b)
	Pine snake (Pituophis melanoleucus)	C. serpentis, C. varanii, Cryptosporidium sp.	(Xiao et al. 2004)
	Prairie king snake (Lampropeltis calligaster)	C. tyzzeri	(Xiao et al. 2004)
	Rough green snake (Opheodrys aestivus)	Cryptosporidium sp.	(Brower and Cranfield 2001)
	Ruthven's kingsnake (Lampropeltis ruthveni)	C. serpentis	(Graczyk and Cranfield 2000)
	Spotted leaf-nosed snake (Phyllorhynchus decurtatus)	C. serpentis	(Graczyk and Cranfield 2000)

(continued)

Family	Host species (scientific name)	Cryptosporidium taxa	Reference
Tanniy	Texas rat snake	<i>Cryptosporidium</i> taxa	(Upton et al. 1989)
	(Elaphe obsolete lindheimeri)	Cryptosportatium sp.	
	Trans-Pecos rat snake (Bogertophis subocularis)	C. serpentis	(Levine 1980)
	Tricolor hognose snake (Lystrophis semicinctus)	Cryptosporidium sp.	(Cerveny et al. 2012)
	Western fox snake (Pantherophis vulpina vulpina)	Cryptosporidium sp.	(Upton et al. 1989)
	Yellow rat snake (Elaphe obsoleta quadrivittata)	C. serpentis, C. muris	(Cranfield and Graczyk 1994; Ryan et al. 2003a)
Elapidae	Cape coral snake (Aspidelaps lubricus lubricus)	Cryptosporidium sp.	(Cerveny et al. 2012)
	Common death adder (Acanthophis antarticus)	C. serpentis	(Morgan et al. 1999c)
	Eastern/mainland tiger snake (Notechis scutatus)	C. serpentis	(Morgan et al. 1999c)
	Mulga/king brown snake (Pseudechis australis)	Cryptosporidium sp.	(Morgan et al. 1999c)
	Red-bellied black snake (Pseudechis porphyriacus)	C. tyzzeri	(Morgan et al. 1999c)
	Taipan (Oxyuranus scutellatus)	C. serpentis	(Morgan et al. 1999c; Xiao et al. 2004)
Pythonidae	Ball python (Python regius)	C. ducismarci, C. muris, C. serpentis, C. tyzzeri, tortoise genotype, Crypto- sporidium sp.	(Xiao et al. 2004; Pedraza-Diaz et al. 2009; Sevá Ada et al. 2011b)
	Burmese python (Python mollurus)	C. serpentis	(Xiao et al. 2004)
	Boelen's python (Morelia boeleni)	C. serpentis	(Xiao et al. 2004)
	Green python (Chondropython viridis)	C. serpentis, C. varanii, Crypto- sporidium sp.	(Xiao et al. 2004)

Table 5.2 (continued)

(continued)

	Host species		
Family	(scientific name)	Cryptosporidium taxa	Reference
	Indian rock python (Python molurus)	C. serpentis	(Rinaldi et al. 2012)
	Woma python (Aspidites ramsayi)	C. tyzzeri, Cryptospo- ridium sp.	(Morgan et al. 1999c; Cerveny et al. 2012)
Viperidae	Bornmueller's viper (Vipera bornmuelleri)	C. serpentis	(Xiao et al. 2004)
	Eastern diamondback rattlesnake (Crotalus adamanteus)	C. serpentis	(Graczyk and Cranfield 2000)
	Jararaca (Bothropoides jararaca)	C. serpentis	(Sevá Ada et al. 2011b)
	Jararacussu (Bothrops jararacussu)	C. serpentis	(Sevá Ada et al. 2011b)
	Mountain viper (Vipera wagneri)	C. serpentis	(Xiao et al. 2004)
	Nikolski viper (Vipera nikolski)	C. serpentis	(Ryan et al. 2003a)
	Northwestern tropical rattlesnake (Crotalus durissus culminatus)	Cryptosporidium sp.	(Upton et al. 1989)
	Pit viper (Crotalus viridis viridis)	Cryptosporidium sp.	(Cerveny et al. 2012)
	Timber rattlesnake (Crotalus horridus)	C. serpentis	(Levine 1980; Heuschele et al. 1986)
	Tropical rattlesnake (Caudisona durissa)	C. serpentis	(Sevá Ada et al. 2011b)

Table 5.2	(continued)
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(*Elaphe vulpina gloydi*), milk snake (*Lampropeltis triangulum*), mangrove monitor (*Varanus indicus*), and prairie king snake (*Lamproletis calligaster*); *C. muris* in a corn snake (*Pantherophis guttatus guttatus*); and rat genotype I in a boa constrictor probably represent passive transmission as a consequence of feeding on infected rodents (Xiao et al. 2004).

Koudela and Modrý (1998) identified and named the species *C. saurophilum* from five lizard species: the desert monitor (*Varanus griseus*), emerald monitor (*Varanus prasinus*), leopard gecko (*Eublepharis macularius*), Schneider's skink (*Eumeces schneideri*), and skink (*Mabuya perrotetii*). Pavlásek and Ryan (2008) subsequently showed that *C. saurophilum* was indistinguishable from *C. varanii*, a species described 4 years earlier in an emerald monitor (Pavlásek et al. 1995), and proposed that *C. varanii* should take precedence as the species name. *Cryptosporidium varanii* (syn. *saurophilum*) infects the intestine and cloaca of a number of lizard species (Table 5.3), causing weight loss and abdominal swelling (Koudela and Modrý 1998). In contrast to *C. serpentis* in snakes, the disease is observed in

Family	Host species (scientific	Cryptosporidium	Deference
Family	name)	taxa	Reference
Agamidae	Bearded dragon (Pogona vitticeps)	C. serpentis, C. varanii	(Xiao et al. 2004)
	Damara rock agama (Agama planiceps)	Cryptosporidium sp.	(Upton et al. 1989)
	Frilled lizard (Chlamydosaurus kingui)	C. serpentis, Cryp- tosporidium sp.	(Pedraza-Diaz et al. 2009; Rinaldi et al. 2012)
	Ground agama (Agama aculeata)	Cryptosporidium sp.	(Upton et al. 1989)
Chamaeleonidae	Giant Madagascar chameleon (Chamaeleo oustaleti)	C. serpentis	(Pedraza-Diaz et al. 2009)
	Mountain chameleon (Chamaeleo montium)	C. serpentis, C. varanii	(Xiao et al. 2004)
	Veiled chameleon (Chamaeleo calyptratus)	C. ducismarci, C. varanii, Cryptosporid- ium sp.	(Koudela and Modrý 1998; Pedraza-Diaz et al. 2009; Rinaldi et al. 2012)
Gekkonidae	African fat-tailed gecko (Hemiteconyx caudicintus)	C. varanii	(Pedraza-Diaz et al. 2009)
	Gargoyle gecko (Rhodocodactylus auriculatus)	C. serpentis, C. varanii	(Xiao et al. 2004)
	Gecko (Gekkoninae sp.)	C. parvum, C. varanii	(Xiao et al. 2004)
	Giant ground gecko (Chondrodactylus angulifer)	Cryptosporidium sp.	(Upton et al. 1989)
	Leopard gecko (Eublepharis macularius)	C. parvum, C. serpentis, C. varanii, Cryptosporid- ium sp.	(Koudela and Modrý 1998; Xiao et al. 2004; Pedraza-Diaz et al. 2009; Richter et al. 2011; Rinaldi et al. 2012)
	Madagascar giant day gecko (Phelsuma madagascariensis grandis)	Cryptosporidium sp.	(Upton et al. 1989)
	Mediterranean house gecko (Hemidactylus turcicus turcicus)	Cryptosporidium sp.	(Upton et al. 1989)
Gerrhosauridae	Plated lizard (Gerrhosaurus sp.)	C. varanii	(Xiao et al. 2004)

 Table 5.3 Cryptosporidium spp. identified in lizards

(continued)

	Host species (scientific	Cryptosporidium	
Family	name)	taxa	Reference
Iguanidae	Green iguana (Iguana iguana)	C. varanii, C. parvum, avian genotype V	(Xiao et al. 2004: Kik et al. 2011)
Lacertidae	European green liz- ard (Lacerta viridis)	C. varanii	(Koudela and Modrý 1998)
Scincidae	Ocellated skink (Chalcides ocellatus)	C. varanii	(Koudela and Modrý 1998)
	Schneider's skink (Eumeces schneideri)	C. varanii	(Koudela and Modrý 1998)
	Skink (Mabuya perrotetii)	C. serpentis, C. varanii	(Koudela and Modrý 1998; Xiao et al. 2004)
Varanidae	Crocodile monitor (Varanus salvadori)	C. muris	(Ryan et al. 2003a)
	Desert monitor	C. serpentis,	(Koudela and Modrý 1998)
	(Varanus griseus)	C. varanii	(Ryan et al. 2003a; Xiao et al. 2004)
	Emerald monitor (Varanus prasinus)	C. varanii, C. serpentis	(Pavlásek et al. 1995; Koudela and Modrý 1998)
	Mangrove monitor (Varanus indicus)	C. tyzzeri	(Xiao et al. 2004)
	Monitor sp. (Varanus sp.)	C. varanii, C. parvum	(Xiao et al. 2004)
	Nile monitor (Varanus niloticus)	C. serpentis	(Xiao et al. 2004)
	Savannah monitor (Varanus exanthematicus)	C. serpentis	(Morgan et al. 1999c; Xiao et al. 2004)

Table 5.3 (continued)

juveniles but not adults (Koudela and Modrý 1998). Oocysts are smaller than *C. serpentis* at 5.0 μ m (4.4–5.6 μ m) × 4.7 μ m (4.2–5.2 μ m) with a shape index of 1.09 (1.04–1.12) (Koudela and Modrý 1998). Under experimental conditions, *C. varanii* was not infectious for snakes, birds, or mice (Koudela and Modrý 1998); however, natural *C. varanii* infections have been detected in at least nine snake species from three families (Table 5.2).

Other *Cryptosporidium* taxa identified in Squamata include a genotype sharing 99 % identity with *C. serpentis* from a leopard gecko (Richter et al. 2011), avian genotype V from a green iguana (*Iguana iguana*) (Kik et al. 2011), *C. ducismarci* and the tortoise genotype from a ball python (Pedraza-Diaz et al. 2009), *C. ducismarci* from a veiled chameleon (Pedraza-Diaz et al. 2009), and a genotype from a New Guinea boa (*Candoia aspera*) and Japanese grass snake (*Rhabdophis tigrinus*) sharing 98 % similarity with *C. ducismarci* and avian genotype II (Xiao et al. 2002; Kuroki et al. 2008).

5.3.2 Testudines

The order Testudines is comprised of 250 species of turtles in 14 families and more than 20 genera. All species possess a bony and cartilaginous carapace. Turtles are found in a great variety of habitats, including dry upland regions as well as marine and fresh waters. Diet is similarly varied, with mostly omnivorous species, but also some herbivorous and carnivorous species.

Cryptosporidium ducismarci (previously known as *Cryptosporidium* sp. ex *Testudo marginata* CrIT-20) has been proposed as a new *Cryptosporidium* species (Traversa et al. 2008; Traversa 2010). This species, which infects the intestinal epithelium of *Testudo marginata*, also has been detected in an asymptomatic veiled chameleon and ball phyton in Spain (Pedraza-Diaz et al. 2009) and in the pancake tortoise (*Malacochersus tornieri*) and Russian tortoise (*Testudo horsfieldi*) in the US (Griffin et al. 2010). Biological data, including descriptions of pathology, and oocyst morphometry data are lacking for this species description.

The *Cryptosporidium* tortoise genotype has been identified in an asymptomatic Indian star tortoise (*Geochelone elegans*) at the Lisbon Zoo (Alves et al. 2005). The SSU rRNA sequence of the isolate identified by Alves et al. (2005) is identical to a sequence previously reported by Xiao et al. (2002) in an Indian star tortoise at the St. Louis Zoo (GenBank accession no. AY120914), and a sequence from an environmental isolate in Maryland, USA (Yang et al. 2008). In a study of pet reptiles, Pedraza-Diaz et al. (2009) detected *Cryptosporidium* sp. in 17.6 % of tortoise samples and identified the *Cryptosporidium* tortoise genotype in a Hermann's tortoise (*Testudo hermanni*) and a ball python (*Python regius*).

5.4 Cryptosporidium and Cryptosporidiosis of Birds

GenBank accession numbers and phylogenetic relationships among representative *Cryptosporidium* small subunit rRNA sequences from birds are presented in Table 5.4 and Fig. 5.2. Three avian-associated *Cryptosporidium* species are recognized: *Cryptosporidium meleagridis*, *Cryptosporidium baileyi*, and *Cryptosporidium galli*. These species infect a broad range of birds; however, they differ in host range and site of infection. *Cryptosporidium meleagridis* also causes disease in humans (McLauchlin et al. 2000; Alves et al. 2003; Cama et al. 2003).

Cryptosporidium meleagridis is primarily a parasite of intestinal epithelial cells of birds, particularly turkeys (Slavin 1955; Sreter and Varga 2000). *Cryptosporidium baileyi* inhabits the respiratory tract, bursa of Fabricius, and cloaca of the domestic chicken and a broad range of other birds (Current et al. 1986; Sreter and Varga 2000). Endogenous stages of *C. galli* are localized on glandular epithelial cells of the proventriculus (Ryan et al. 2003b).

The three avian species can be distinguished based on oocyst morphology. Oocysts of *C. meleagridis* are the smallest of the three, measuring

Table 5.4 GenBank accession numbers of representative *Cryptosporidium* small subunit rRNAgene sequences from avians. A phylogeny constructed from sequences in this table is presented inFig. 5.2

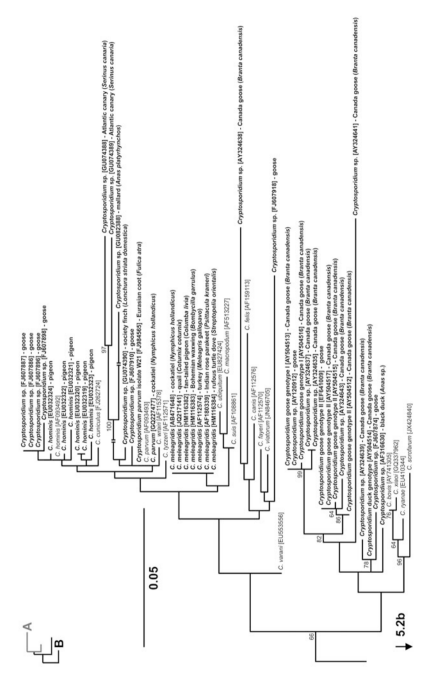
Order	Common names/groups	Host species (scientific name) and GenBank accession numbers
Anseriformes	Waterfowl: geese and ducks	Black duck (species unspecified) [AF316630]
		Goose (species unspecified) [AY120912, EF641009, FJ607874, FJ607886, FJ607887, FJ607896, FJ607898, FJ607910, FJ607918]
		Canada goose (Branta canadensis) [AY324635, AY324637–AY324639, AY324641, AY324643, AY504512– AY504517]
		Goose (Anser anser f. domestica) [FJ984564]
		Mallard (Anas platyrhynchos) [GU082388]
Charadriiformes	Gulls and woodcock	Eurasian woodcock (Scolopax rusticola) [AY273769]
		Kelp gull (Larus dominicanus) [GQ355891]
Columbiformes	Pigeons and doves	Fan-tailed pigeon (Columba livia) [HM116382]
		Pigeon (species unspecified) [EU032319– EU032324]
		Rufous turtle dove (<i>Streptopelia orientalis</i>) [HM116384]
Falconiformes	Black vulture	Black vulture (Coragyps atratus) [GQ227474]
Galliformes	Fowl: chickens, turkey, quail, Indian peafowl	Chicken (Gallus gallus domesticus) [AF093495, JX548291, JX548292, JX548299, GQ227476, AY168847, AY168848]
		Cockatiel (Nymphicus hollandicus) [GQ227477]
		Indian peafowl (Pavo cristatus) [GQ227478]
		Quail (Coturnix coturnix) [JQ217141, JQ217142]
		Quail (species unspecified) [AF316631]
		Turkey (Meleagris gallopavo) [AF112574]
Gruiformes	Coot	Eurasian coot (Fulica atra) [FJ984565]
Passeriformes	Perching birds: finches, canaries, sparrows, nightingale,	Atlantic canary (Serinus canaria) [GU074388, GU074389]
	waxwings, corvids	Aurora finch (<i>Pytilia phoencoptera</i>) [AF316627–AF316629]
		Blackbilled magpie (<i>Pica pica</i>) [HM116380]

(continued)

0.1		Host species (scientific name) and
Order	Common names/groups	GenBank accession numbers
		Bohemian waxwing (Bombycilla garrulus) [HM116383, HM116388]
		Canary (Serinus canaria) [GQ227479, EU543269]
		Common myna (Acridotheres tristis) [HM116374]
		Crested lark (Galerida cristata) [HM116379]
		Gold finch (<i>Carduelis carduelis</i>) [AY168846]
		Gouldian finch (Erythrura gouldiae) [AF316623–AF316625, HM116377]
		Java sparrow (Padda oryzivora) [GU074384]
		Lesser seed finch (Oryzoborus angolensis) [EU543270]
		Plum-headed finch (<i>Neochmia modesta</i>) [AF316626]
		Red-billed Leiothrix (<i>Leiothrix lutea</i>) [HM116375]
		Red-billed blue magpie (Urocissa erythrorhyncha) [HM116386]
		Silver eared leiothrix (Leiothrix argentauris) [HM116387]
		Society finch (<i>Lonchura striata domestica</i>) [GU074390]
		White Java sparrow (Padda oryzivora) [HM116376]
		Zebra finch (<i>Taeniopygia guttata</i>) [HM116378]
Psittaciformes	Parrots, cockatiels, lovebirds	Cockatiel (Nymphicus hollandicus) [AB471645–AB471648, EU543268, GQ227477, GQ227481, GU074385–GU074387, HM116381, HM116385]
		Indian rose parakeet (Psittacula krameri) [AF180339]
		Peach-faced lovebird (Agapornis roseicollis) [GQ227480]
Struthioniformes	Ostrich	Ostrich (Struthio camelus) [DQ002931]

Table 5.4 (continued)

4.5–6.0 μ m × 4.2–5.3 μ m with a shape index (length/width) of 1.0–1.3. *Cryptosporidium baileyi* oocysts measure 6.0–7.5 μ m × 4.8–5.7 μ m with a shape index of 1.1–1.8 (Lindsay et al. 1989), and oocysts of *C. galli* measure 8.2 × 6.3 μ m with a shape index of 1.3 (Ryan et al. 2003b). Relatively little is known about the biology of other cryptosporidia infecting birds.





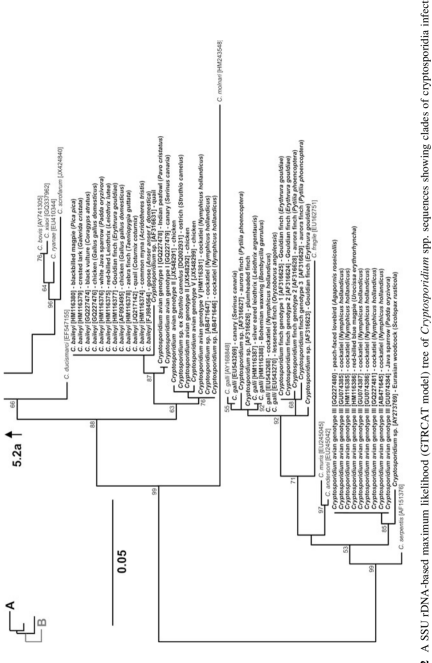


Fig. 5.2 A SSU rDNA-based maximum likelihood (GTRCAT model) tree of Cryptosporidium spp. sequences showing clades of cryptosporidia infecting avians. Cryptosporidium sp. detected in avians are in bold. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) is shown at nodes. Bootstrap values <50 are not shown. Interrupted branches have been shortened fivefold *Cryptosporidium* has been reported in 17 out of 26 avian orders. Members of the following orders have not yet been identified as hosts:

- **Tinamiformes:** This is a very old avian lineage comprising one family, nine genera, and 47 species. Members are ominvorous, feeding on a variety of seeds and invertebrates. Many species occur in forests or open grasslands, but they generally prefer dryland habitats.
- **Gaviiformes:** This order comprises one family, one genus, and five species of aquatic birds that primarily inhabit North America and Eurasia. They feed on fish, amphibians, or aquatic invertebrates and inhabit both freshwater and marine environments during their annual cycle.
- **Podicipediformes:** This order comprises 22 species in one family and six genera of aquatic birds. Molecular phylogenies indicate that they are closely related to loons and flamingoes. They feed on fish, amphibians, and aquatic invertebrates.
- **Procellariiformes**: This order comprises four families, 26 genera, and about 112 species of seabirds. All inhabit offshore marine environments and most return to land only for nesting. Although, most feed exclusively on marine invertebrates or fish, a couple of species scavenge carrion on islands.
- **Opisthocomiformes:** This order comprises one species, genus, and family of tropical birds that inhabit swamps and mangroves of South America. They feed on vegetable matter (seeds, nuts, leaves, and fruits), and have well-developed ceca that facilitates fermentation in the digestive tract.
- **Cuculiformes:** This is a medium sized order comprising six families, 30 genera, and 143 species. Members inhabit Africa, Asia, Australasia, Eurasia, and the Americas (South, Central, and North).
- **Apodiformes:** This order comprises two families containing approximately 24 genera and 100 species of small birds. They mostly inhabit the northern hemisphere and can be found in a variety of habitats. They feed aerially on insects.
- **Trochiliformes:** These small, nectar-feeding birds include hermits and hummingbirds. There are approximately 330 species in one family with about 100 genera. The majority of species occur in tropical and subtropical Central and South America.
- **Trogoniformes**: Trogons comprise approximately 39 species in one family with six to eight genera. They inhabit tropical forests, where they feed mainly on insects or fruit.

5.4.1 Struthioniformes

The order Struithioniformes (including cassowaries, emus, kiwis, ostriches, and rheas) is comprised of some of the oldest avian species. There are about 12 species in five families and six genera. All are flightless, forage on insects or vegetation, and lay large eggs with thick shells. Ceca are often well developed in these species.

Cryptosporidium infections in ostriches can be subclinical (Gajadhar 1993), or characterized by a prolapsed phallus/cloaca (Bezuidenhout et al. 1993; Santos et al. 2005), and edema/necrosis of the pancreas (Jardine and Verwoerd 1997). In Brazil, ostriches with a prolapsed cloaca shed oocysts of *Cryptosporidium* avian genotype II measuring 6.0 μ m (5.0–6.5 μ m) \times 4.8 μ m (4.2–5.3 μ m) with a shape index of 1.31 (Santos et al. 2005; Meireles et al. 2006). Cryptosporidium avian genotype II is not infectious for 1-day-old chicks under experimental conditions (Meireles et al. 2006), although it has been detected in a naturaly infected chicken (GenBank accession no. JX548291; unpublished). Cryptosporidium avian genotype II was detected in 23 % of farmed ostriches (Struthio camelus) in Vietnam (Nguyen et al. 2013). The highest prevalence and oocyst shedding intensity was observed in birds aged 61–90 days; younger and older birds had a lower prevalence and shed fewer oocysts (Nguyen et al. 2013). Oocysts from subclinical Cryptosporidium infections in ostriches being imported into Canada measured 4.6 um $(3.9-6.1 \text{ um}) \times 4.0 \text{ um}$ $(3.3-5.0 \,\mu\text{m})$ with a shape index of 1.15 (range 1.0-1.4) (Gajadhar 1993), which is similar to oocysts of C. meleagridis. Oocysts were not infectious for suckling mice, chickens, turkeys, or quail (Gajadhar 1993, 1994). Cryptosporidium baileyi oocysts were identified in an ostrich from the Czech Republic (Ryan et al. 2003a). Cryptosporidium sp. was detected in 60 % of farmed ostriches and rheas originating from Belgium, France, Netherlands Portugal, and Spain (Gordo et al. 2002).

5.4.2 Anseriformes

Waterfowl, including ducks, geese and swans, form the order Anseriformes, with about 162 species in three families and 52 genera. These birds are omnivorous, feeding on combinations of vegetation and invertebrates, with a few groups feeding on small aquatic vertebrates.

Three species and five named genotypes of *Cryptosporidium* have been identified in ducks, geese, mergansers, and swans (Table 5.5). In addition, five unnamed genotypes have been reported in Canada geese (*Branta canadensis*) (Jellison et al. 2004), however, sequences have not been deposited in GenBank, and it is not clear if they differ from subsequently named genotypes. Canada geese have also been shown to host the human pathogens *C. hominis* and *C. parvum*, and the rodentspecific muskrat genotype I (Graczyk et al. 1998d; Zhou et al. 2004b; Jellison et al. 2009). It is hypothesized that birds serve as mechanical vectors for mammalian cryptosporidia, and this is supported by data showing that *C. parvum* does not infect Canada geese or Pekin ducks (*Anas platyrhynchos*) under experimental conditions but oocysts remain viable following passage (Graczyk et al. 1996b, 1997).

Cryptosporidium has been reported to cause varying degrees of respiratory distress in ducks and geese (O'Donoghue et al. 1987). *Cryptosporidium baileyi* has been identified in the bursa of Fabricius and cloaca of ducks with mild symptoms (Lindsay et al. 1989), and the bursa of Fabricius and conjunctiva of domestic geese with no clinical symptoms (Chvala et al. 2006).

Host species		
(scientific name)	Cryptosporidium taxa	References
American widgeon (Anas americana)	Cryptosporidium sp.	(Kuhn et al. 2002)
Black swan (Cygnus atratus)	Cryptosporidium sp.	(Rohela et al. 2005)
Blue-winged teal (Anas discors)	Cryptosporidium sp.	(Kuhn et al. 2002)
Canada goose (Branta canadensis)	<i>C. hominis, C. hominis</i> -like, <i>C. parvum</i> , muskrat genotype I, duck geno- type, goose genotype I, goose genotype II, 5 unnamed genotypes	(Graczyk et al. 1998d; Jellison et al. 2004, 2009; Zhou et al. 2004b)
Common merganser (Mergus merganser)	Cryptosporidium sp.	(Kuhn et al. 2002)
Domestic goose (Anser anser f. domestica)	C. baileyi	(Chvala et al. 2006)
Green-winged teal (Anas cercca carolinensis)	Cryptosporidium sp.	(Kuhn et al. 2002)
Hooded merganser (Lophodytes cucullatus)	Cryptosporidium sp.	(Kuhn et al. 2002)
Mallard (Anas platyrhynchos)	Cryptosporidium sp.	(O'Donoghue et al. 1987; Kuhn et al. 2002)
Swan goose (Anser cygnoides)	Cryptosporidium sp.	(Rohela et al. 2005)

Table 5.5 Cryptosporidium spp. identified in Anseriformes species

5.4.3 Galliformes

There are approximately 290 species in five families and 80 genera of gallinaceous birds. These terrestrial species feed mostly on seeds and small insects. This group has been extensively domesticated for agricultural purposes, and many species have been introduced to areas well outside of their historic range.

A number of gallinaceous birds including the domestic chicken (*Gallus gallus domesticus*), grouse (*Tetrastes bonasia rupestris*), and capercaillie (*Tetrao urogallus*), are natural hosts of *C. galli* (Ryan et al. 2003b; Ng et al. 2006).

Non-genotyped cryptosporidia have been detected in the common quail (*Coturnix coturnix*), common peafowl (*Pavo cristatus*), great argus (*Argusianus argus*), great currasow (*Crax rubra*), red-legged partridge (*Alectoris rufa*), and ring-necked pheasant (*Phasianus colchicus*) (O'Donoghue et al. 1987; Rohela et al. 2005; Lim et al. 2007). Infections in the common quail and ring-necked pheasant have been associated with clinical signs of respiratory disease, and oocysts recovered from these hosts were infectious for chickens (O'Donoghue et al. 1987).

Cryptosporidium meleagridis caused an outbreak of respiratory and diarrheal cryptosporidiosis with greater than 50 % mortality in red-legged partridges (*Alectoris rufa*) on a game farm in Spain (Pages-Mante et al. 2007).

5.4.4 Phoenicopteriformes

Phoenicopteriformes is a small order of five species in one family and three genera of flamingoes. These long-legged wading birds use a specialized bill to filter-feed invertebrates or algae from shallow waters. The intestine is relatively long compared to other avian species.

In the only report from this order, *C. galli* was identified in a Cuban flamingo (*Phoenicopterus ruber*) from the Czech Republic (Ng et al. 2006).

5.4.5 Sphenisciformes

Sphenisciformes is comprised of 17 penguin species in one family and six genera. Penguins are flightless birds of the southern oceans, feeding on fish and krill in offshore areas. Very little is known about cryptosporidiosis in this group of birds.

Cryptosporidium oocysts have been detected in 6.6 % (11/167) of stools from the Adelie penguin (*Pygoscelis adeliae*) from the Antarctic territory (Fredes et al. 2007), and in 32.8 % (21/64) of gentoo penguins (*Pygoscelis papua*) from Ardley Island, King George Island, South Shetland Islands, and the Antarctic Specially Protected Area no. 150 (Fredes et al. 2008).

5.4.6 Pelecaniformes

Pelecaniformes includes waterbirds comprising approximately 65 species in eight families and ten genera. These birds feed mostly on aquatic vertebrates. A gular pouch is usually present, and prey is often swallowed whole. Almost all inhabit marine waters for part of their annual cycle.

There have been just two reports of *Cryptosporidium* from the order Pelecaniformes. *Cryptosporidium baileyi* and a non-genotyped *Cryptosporidium* sp. were reported in a cormorant and great cormorant (*Phalacrocorax carbo*), respectively (Jellison et al. 2004; Plutzer and Tomor 2009).

5.4.7 Ciconiiformes

There are approximately 116 ciconiform species in three families and 39 genera. All are long-legged wading birds, feeding in aquatic environments on a variety of invertebrates and vertebrates.

Cryptosporidium has been reported in a marabou stork (*Leptoptilos crumeniferus*) from a zoo in Malaysia (Rohela et al. 2005) and 12.5 % (3/24) of white storks (*Ciconia ciconia*) from Poland (Majewska et al. 2009). The species infecting white storks was identified as *C. parvum* using fluorescence in situ hybridization.

5.4.8 Falconiformes

There are approximately 304 species in three to five families and 83 genera of falconiform birds. These birds feed on vertebrate prey or carrion. Most nest in trees or cliffs and are terrestrial in their habits.

Cryptosporidium baileyi and C. parvum have been identified in Falconiforms, and both are associated with disease. Cryptosporidium parvum caused anorexia, mild respiratory difficulties, and mild bilateral ocular discharge in a 3-month old gyrfalcon (*Falco rusticolus*), and C. baileyi caused mild bilateral conjunctivities and sinusitis in a 13-year-old gyrfalcon × Saker falcon hybrid (*Falco rusticolus* × *Falco cherrug*) (Barbon and Forbes 2007). Both raptors recovered following treatment with paromomycin. Cryptosporidium baileyi similarly caused an upper respiratory tract infection in three mixed-bred falcons (*Falco rusticolus* × *Falco cherrug*) (van Zeeland et al. 2008). One of the birds had epiglottal swelling and laryngeal stridor, and two had nasal discharge and sneezing. Oocyst shedding was not detected in any of the birds. Cryptosporidium baileyi also was identified in a Saker falcon (*Falco cherrug*) with inflammation of the middle ears, conjunctivae, third eyelids, choanae, larynx, trachea, salivary glands of the tongue, syrinx, and turbinates (Bougiouklis et al. 2012), and in a captive black vulture (*Coragyps atratus*) (Nakamura et al. 2009).

5.4.9 Charadriiformes

Charadriiformes is a diverse order of shorebirds, with about 367 species distributed among 17 families and 88 genera. Most species feed on invertebrate prey, and utilize marine habitats during part of their annual cycle.

Cryptosporidium sp. has been reported in black-headed gulls (*Chroicocephalus ridibundus*) (Pavlásek 1993; Smith et al. 1993; Ryan et al. 2003a), herring gulls (*Larus argentatus*) (Smith et al. 1993; Bogomolni et al. 2008), and kelp gulls

(*Larus dominicanus*) (GenBank accession no. GQ355891; unpublished). Smith et al. (1993) detected *Cryptosporidium* oocysts measuring 4.7 μ m (4.4–5.2 μ m) × 5.2 μ m (4.7–5.8 μ m) in feces from gulls trapped at two refuse sites and other locations in Scotland, and found no difference in *Cryptosporidium* prevalence between the black-headed gull and the herring gull. *Cryptosporidium baileyi* was identified as a cause of significant morbidity and mortality in 28–100 % of black-headed gulls in the Czech Republic (Pavlásek 1993). *Cryptosporidium baileyi* from black-headed gulls infected 4-day-old chickens (*Gallus gallus f. domestica*), causing 40 % mortality (Pavlásek 1993). The intestinal locations of developmental stages are similar to those in the black-headed gull, and a novel *Cryptosporidium baileyi* was reported in a black-headed gull, and a novel *Cryptosporidium genotype* (GenBank accession no. AY273769) was identified in a wild-caught Eurasian woodcock (*Scolopax rusticola*) from the Czech Republic (Ryan et al. 2003a).

5.4.10 Gruiformes

Gruiformes contains 212 species distributed among 11 families and 61 genera. Most species inhabit marshland areas, and feed on invertebrates, seeds, or other vegetation. Ceca and hindgut fermentation are common in this group.

Cryptosporidium baileyi has been identified in a crane from the Czech Republic (Ng et al. 2006). In Korea, a 4-month old white naped crane (*Grus vipio*) that died from a disseminated *Eimeria* infection also had an incidental *Cryptosporidium* infection of the cloaca (Kim et al. 2005). In Hungary, Plutzer and Tomor (2009) identified *C. parvum* in a wild Eurasian coot (*Fulica atra*) (GenBank accession no. FJ984565).

5.4.11 Columbiformes

Pigeons and doves make up the order Columbiformes. Although there are 308 species worldwide, only one or two families containing 42 genera are recognized. These birds occupy a diverse range of terrestrial habitats, feeding almost exclusively on seeds and grains.

There have been relatively few reports of *Cryptosporidium* in pigeons (Ozkul and Aydin 1994; Rodriguez et al. 1997; Abreu-Acosta et al. 2009; Qi et al. 2011; Radfar et al. 2012). *Cryptosporidium hominis* has been reported in rock pigeons (*Columba livia*) on Tenerife, one of the Canary Islands, suggesting that pigeons can play a role in the transmission of human-pathogenic cryptosporidia (Abreu-Acosta et al. 2009). In a study of *Cryptosporidium* in birds at pet stores in China, Qi et al. (2011) detected *C. meleagridis* in a fan-tailed pigeon and a rufous turtle dove (*Streptopelia orientalis*).

5.4.12 Psittaciformes

Birds in the order Psittaciformes order, which contains 364 species from 85 genera in the family Psittacidae, inhabit trees in dense, tropical or subtropical forests, and feed almost exclusively on fruits and seeds. Although there have been few studies of psittaciforms in their natural environment, captive psittaciforms host a number of *Cryptosporidium* species and genotypes (Table 5.6). The greatest diversity of taxa has been detected in the cockatiel (*Nymphicus hollandicus*), which hosts four species (*C. baileyi*, *C. galli*, *C. meleagridis*, and *C. parvum*) and three genotypes (avian genotypes II, III, and V) (Abe and Iseki 2004; Ng et al. 2006; Antunes et al. 2008; Nakamura et al. 2009; Abe and Makino 2010; Qi et al. 2011; Gomes et al. 2012). The galah (*Eolophus roseicapilla*), which is in the same subfamily as the cockatiel, also hosts avian genotype II (Ng et al. 2006).

Although most cases of *Cryptosporidium* infection in psittaciform birds have been asymptomatic, there have been reports of clinical disease. Latimer et al. (1992) diagnosed cryptosporidiosis in four cockatoos with psittacine beakand-feather disease. *Cryptosporidium* infection was confined to the bursa of Fabricius in three of the birds, and was more widespread in the intestine of the fourth. All birds had intermittent diarrhea. Makino et al. (2010) detected avian genotype II in 35 % (13/37) of infected peach-faced lovebirds (*Agapomis roseicollis*). All birds had symptoms of infection including weight loss and chronic vomiting.

5.4.13 Strigiformes

The owls make up a small order of about 180 species in two families and 29 genera. All species are predators of small vertebrates, principally rodents. Most species are nocturnal and inhabit forested areas, though a few nest underground in burrows.

In a study of 12 adult owls held in captivity in south Brazil, da Silva et al. (2009) isolated *Cryptosporidium* oocysts measuring 5–6 μ m × 4–5 μ m from a barn owl (*Tyto alba*), great horned owl (*Bubo virginianus*), and striped owl (*Phinoptynx clamator*). Owls showed no clincal signs of disease.

Molina-Lopez et al. (2010) diagnosed ocular and respiratory cryptosporidiosis in 16 wild fledgling scops owls (*Otus scops*) up to 2 months after they were admitted to a wildlife rehabilitation center in Catalonia, northern Spain; the owls were born in the wild and were healthy when they arrived at the center. Blepharoedema, conjunctival hyperaemia, and mucopurulent ocular discharge were diagnosed unilaterally in 75 % (12/16) of the birds and bilaterally in 25 % (4/16). Five owls (31 %) developed diffuse epithelial corneal edema, one exhibited mild anterior exudative uveitis, and another developed rhinitis. *Cryptosporidium baileyi*, measuring 6.5–7.0 μ m × 5.0–5.5 μ m, was identified in samples from two birds that were

Host species (scientific name)	Cryptosporidium taxa	References
Alexandrine para- keet (Psittacula eupatria)	Avian genotype II, Cryptosporid- ium sp.	(Ng et al. 2006; Papini et al. 2012)
Blue-fronted Ama- zon (Amazona aestiva)	Cryptosporidium sp.	(Papini et al. 2012)
Budgerigar (Melopsittacus undulatus)	Cryptosporidium sp.	(Nakamura et al. 2009)
Cockatiel (Nymphicus hollandicus)	C. baileyi, C. galli, C. meleagridis, C. parvum, avian genotype II, avian genotype III, avian geno- type V	(Abe and Iseki 2004; Ng et al. 2006; Antunes et al. 2008; Nakamura et al. 2009; Abe and Makino 2010; Qi et al. 2011; Gomes et al. 2012)
Eastern rosella (Piatycercus eximius)	Cryptosporidium sp.	(Papini et al. 2012)
Eclectus parrot (Eclectus roratus)	Avian genotype II	(Ng et al. 2006)
Galah (Eolophus roseicapilla)	Avian genotype II	(Ng et al. 2006)
Goffin's cockatoo (Cacatua goffini)	Cryptosporidium sp.	(Nakamura et al. 2009)
Indian ring-necked parrot (Psittacula krameri)	C. meleagridis	(Morgan et al. 2000b)
Major Mitchell's cockatoo (Lophochroa leadbeateri)	Avian genotype II	(Ng et al. 2006)
Peach faced lovebirds (Agapomis roseicollis)	Avian genotype III	(Makino et al. 2010)
Princess parrot (Polytelis alexandrae)	Avian genotype II	(Ng et al. 2006)
Red-bellied macaw (Orthopsittaca manilata)	Cryptosporidium sp.	(Nakamura et al. 2009)
Red-crowned Ama- zon (Amazona viridigenalis)	C. baileyi	(Ryan et al. 2003a)
Rose-ringed para- keet (Psittacula krameri)	C. meleagridis	(Ryan et al. 2003a)

 Table 5.6 Cryptosporidium spp. identified in Psittaciformes species

(continued)

Host species (scientific name)	Cryptosporidium taxa	References
Salmon-crested cockatoo (Cacatua moluccensis)	Cryptosporidium sp.	(Rohela et al. 2005)
Sun conure (Aratinga solstitialis)	Avian genotype II	(Ng et al. 2006)
Turquoise parrot (Neophema pulchella)	C. galli	(Ng et al. 2006)
White-eyed para- keet (Aratinga leucophthalma)	Avian genotype II	(Sevá Ada et al. 2011a)

Table 5.6 (continued)

euthanized due to the severity of their disease. The remaining owls recovered following a 15-day treatment with azithromycin.

5.4.14 Caprimulgiformes

This small order of insectivorous birds has about 118 species in five families and 22 genera. Most feed nocturnally, on the wing. These species generally occur near upland forests, and almost all species nest on the ground.

Cryptosporidium muris has been detected in the tawny frogmouth (*Podargus strigoides*); although, the bird may have been a mechanical vector (Ryan et al. 2003a; Ng et al. 2006).

5.4.15 Piciformes

The piciform birds include woodpeckers and toucans, with approximately 398 species in five to nine families and eight genera. Most species occur in forests of the Americas, and almost all nest in cavities. The woodpeckers feed mostly on insects, but toucans are fruigivorous and the honeyguides have the unique ability to eat beeswax.

Cryptosporidium infection has been reported in a channel-billed toucan (*Rhamphastus vitellinus*), chestnut-eared aracari (*Pteroglossus castanotis*), and Toco toucan (*Ramphastos toco*) (Ryan et al. 2003a; Nakamura et al. 2009). The channel-billed toucan was infected with *C. baileyi* (Ryan et al. 2003a).

5.4.16 Coraciiformes

Hornbills, kingfishers, and rollers make up the order Coraciiformes, which has about 209 species in 11 families and 51 genera that are found mostly in Eurasia. These birds are generally omnivorous, feeding on a variety of invertebrates, small vertebrates, or fruits and seeds. The coraciiform birds typically inhabit forests, but ground dwelling species occur in sub-Saharan Africa. Most nest in cavities or burrows.

To date, only a few species from the order Coraciiforme have been identified as hosts for *Cryptosporidium*. *Cryptosporidium galli* was identified in a captive rhinocerous hornbill (*Buceros rhinoceros*) in the Czech Republic (Ng et al. 2006). In Malaysia, *Cryptosporidium* sp. was detected in a wrinkled hornbill (*Aceros corrugatus*) and a wreathed hornbill (*Aceros undulates*) at the Kuala Lumpur National Zoo and Zoo Negara, respectively (Rohela et al. 2005; Lim et al. 2007).

5.4.17 Passeriformes

Passeriformes is the largest avian order, with over 5,700 species in approximately 96 families and more than 1,200 genera. All species have feet for perching, and they occupy a wide range of habitats from dense forests to open grasslands. Fruits, seeds and invertebrates are the dominant foods in their diet.

Passerines are relatively frequent hosts of *Cryptosporidium* sp., some of which are human pathogens (Table 5.7). Yet, despite having more than 50 % of the avian diversity, passerines host only four species (*C. baileyi*, *C. galli*, *C. meleagridis*, and *C. parvum*) and three genotypes (avian genotypes I, III, and IV) of *Cryptosporidium* (Table 5.6).

Australian passerines infected with avian genotypes I, III, and IV showed no clinical signs; whereas, birds from the Czech Republic, which were primarily infected with *C. baileyi* and *C. galli*, had diarrhea and anorexia (Ng et al. 2006).

5.5 Cryptosporidium and Cryptosporidiosis of Mammals

Oocyst size and gastrointestinal localization of *Cryptosporidium* species reported in wild mammals are shown in Table 5.8. In contrast to cryptosporidia of domestic animals, oocyst morphometry, infection site, and course of infection are unknown for many *Cryptosporidium* genotypes infecting wild mammals, including bear, beaver, brushtail possum II, chipmunk II and III, *C. bovis*-like, *C. hominis*-monkey, *C. muris*-like, *C. ryanae*-variant, *C. suis*-like, deer, deer mouse I–IV, elephant seal, fox, giant panda, guinea pig, hamster, hedgehog, horse, kangaroo I, mink, mouse II,

5 Cryptosporidiosis in Other Vertebrates

Family	Host species (scientific name)	Cryptosporidium taxa	Reference
Alaudidae	Crested lark (Galerida cristata)	C. baileyi	(Qi et al. 2011)
Bombycillidae	Bohemian waxwing (Bombycilla garrulous)	C. galli, C. meleagridis	(Qi et al. 2011; Sevá Ada et al. 2011a)
Cardinalidae	Green-winged saltator (Saltator similis)	C. galli	(Sevá Ada et al. 2011a)
Corvidae	Black-billed magpie (<i>Pica pica</i>)	C. baileyi	(Qi et al. 2011)
	Hooded crow (Corvus cornix)	Cryptosporidium sp.	(Plutzer and Tomor 2009)
	Red-billed blue magpie (Urocissa erythrorhyncha)	Avian genotype III	(Qi et al. 2011)
Estrildidae	Aurora finch (Pytilia hypogrammica)	Cryptosporidium sp.	(Morgan et al. 2001)
	Australian diamond firetail finch (Stagonoplura bella)	Cryptosporidium sp. ^a	(Blagburn et al. 1990)
	Bengalese finch (Lonchura striata domestica)	C. parvum ^b	(Gomes et al. 2012)
	Black-throated finch (Poephila cincta)	Cryptosporidium sp.	(Gardiner and Imes 1984)
	Bronze mannikin finch (Lonchura cucullata)	Cryptosporidium sp.	(Lindsay et al. 1991)
	Chestnut finch (Lonchura castaneothorax)	C. galli	(Ng et al. 2006)
	Diamond firetail finch (Stagonopleura guttata)	Cryptosporidium sp.	(Lindsay et al. 1991)
	Gouldian finch (<i>Erythrura gouldiae</i>)	C. baileyi, C. galli	(Morgan et al. 2001; Qi et al. 2011)
	Java sparrow (Padda oryzivora)	C. baileyi, avian genotype III	(Qi et al. 2011; Gomes et al. 2012)
	Painted firetail finch (Emblema pictum)	C. galli	(Ng et al. 2006)
	Parson's finch (<i>Poephila cincta</i>)	C. galli	(Ng et al. 2006)
	Plum-headed finch (Neochmia modesta)	Cryptosporidium sp.	(Morgan et al. 2001)
	Red-face aurora finch (<i>Pytilia hypogrammica</i>)	Cryptosporidium sp.	(Morgan et al. 2001)
	Zebra finch (Taeniopygia guttata)	C. galli, C. baileyi	(Ng et al. 2006; Qi et al. 2011)
Fringillidae	Canary (Serinus canaria)	C. galli, avian geno- type I	(Ng et al. 2006; Nakamura et al. 2009)
	Goldfinch (Carduelis tristis) Pine grosbeak (Pinicola	C. galli C. galli	(Sevá Ada et al. 2011a) (Ryan et al. 2003b)
	enucleator) White lored euphonia (Euphonia chrysopasta)	Cryptosporidium sp.	(Lindsay et al. 1991)

 Table 5.7 Cryptosporidium spp. identified in Passeriformes species

(continued)

Family	Host species (scientific name)	Cryptosporidium taxa	Reference
Hirundinidae	Cliff swallow (<i>Petrochelidon pyrrhonota</i>)	Cryptosporidium sp.	(Ley et al. 2012)
Icteridae	Chopi blackbird (Gnorimopsar chopi)	Cryptosporidium sp.	(Nakamura et al. 2009)
	Crested oropendola (<i>Psarocolius decumanus</i>)	C. baileyi	(Ryan et al. 2003b)
	Red-rumped cocique (<i>Cacicus haemorrhous</i>)	C. baileyi	(Ryan et al. 2003b)
Leiothrichidae	Red-billed leiothrix (Leiothrix lutea)	C. baileyi	(Qi et al. 2011)
	Silver-eared mesia (Leiothrix argentauris)	C. galli	(Qi et al. 2011)
Ploceidae	Eastern golden-backed weaver (Ploceus jacksoni)	C. baileyi	(Ng et al. 2006)
Pycnonotidae	Gray-bellied bulbul (<i>Pycnonotus cyaniventris</i>)	C. baileyi	(Ng et al. 2006)
Sturnidae	Common myna (Acridotheres tristis)	C. baileyi	(Qi et al. 2011)
Thraupidae	Double-collared seedeater (Sporophila caerulescens)	Cryptosporidium sp.	(Nakamura et al. 2009)
	Lesser seed finch (Oryzoborus sp.)	C. galli	(Antunes et al. 2008; Nakamura et al. 2009)
	Red-cowled cardinal (Paroaria dominicana)	C. galli	(Ng et al. 2006)
	Saffron finch (Sicalis flaveola)	C. baileyi, C. galli	(Nakamura et al. 2009; Sevá Ada et al. 2011a)
	Slate-coloured seedeater (Sporophila schistacea)	C. galli	(Sevá Ada et al. 2011a)
Turdidae	Rufous-bellied thrush (<i>Turdus rufiventris</i>)	C. galli, Cryptospo- ridium sp.	(Sevá Ada et al. 2011a)
Zosteropidae	Japaneese white eye (Zosterops japonicus)	Avian genotype IV	(Ng et al. 2006)

 Table 5.7 (continued)

^aAssociated with clinical disease

^bC. parvum was identified using molecular tools; however, oocysts from the sample measured 7.2 μ m \times 5.8 μ m, which suggests that another species was also present

muskrat I and II, opposuum II, raccon, rat I–IV, Sbey/Sbld A, Sbey B, Sbey/Sbld/ Sltl C, Sbld D, seal I and II, shrew, skunk, vole, W12, W18, and many other unnamed genotypes (Fig. 5.3).

Table 5.9 shows GenBank accession numbers for representative SSU rRNA gene sequences from mammal-associated cryptosporidia. *Cryptosporidium* has been identified in the three major groups of mammals: the egg-laying mammals (Protheria), marsupials (Metatheria), and placental mammals (Eutheria).

Cryptosporidium			
taxa	Oocysts size (µm)	Infection site	Reference
C. andersoni	6.0-8.1 × 5.0-6.5	Abomasum	(Lindsay et al. 2000)
C. baileyi	6.0–7.5 × 4.8–5.7	Brusa of Fabricius, cloaca	(Lindsay et al. 1989)
C. canis	$3.68-5.88 \times 3.68-5.88$	Small intestine	(Fayer et al. 2001)
C. cuniculus	$5.55-6.40 \times 5.02-5.92$	Small intestine	(Robinson et al. 2010)
C. fayeri	$4.5-5.1 \times 3.8-5.0$	ND	(Ryan et al. 2008)
C. felis	4.6 × 4.0 (3.2–5.1 × 3.0–4.0)	Small intestine	(Iseki 1979)
C. macropodum	$4.5-6.0 \times 5.0-6.0$	ND	(Power and Ryan 2008)
C. muris	7.5–9.8 × 4.6–6.3	Stomach	(Upton and Current 1985)
C. parvum	5.2–5.7 × 4.7–5.3	Small intestine	(Vítovec et al. 2006)
C. scrofarum	$4.81 - 5.96 \times 4.23 - 5.29$	Small intestine	(Kváč et al. 2013a)
C. suis	$6.0-6.8 \times 5.3-5.7$	Large intestine	(Vítovec et al. 2006)
C. tyzzeri	$4.64 \pm 0.05 \times 4.19 \pm 0.06$	Small intestine	(Ren et al. 2012)
C. ubiquitum	$4.71-5.32 \times 4.33-4.98$	Small intestine	(Fayer et al. 2010)
C. wrairi	5.4 × 4.6 (4.8–5.6 × 4.0–5.0)	Small intestine	(Tilley et al. 1991)
Brushtail genotype I	$3.92 \pm 0.25 \ imes 4.12 \pm 0.34$	ND	(Hill et al. 2008)
Chipmunk genotype I	5.3-6.6 × 4.7-5.9	ND	(Kváč et al. 2008a)
Ferret genotype	$4.9-6.0 \times 4.7-5.6$	ND	(Kváč et al. 2008a)

Table 5.8 Oocyst morphology and infection site of *Cryptosporidium* taxa detected in wild mammals

ND not determined

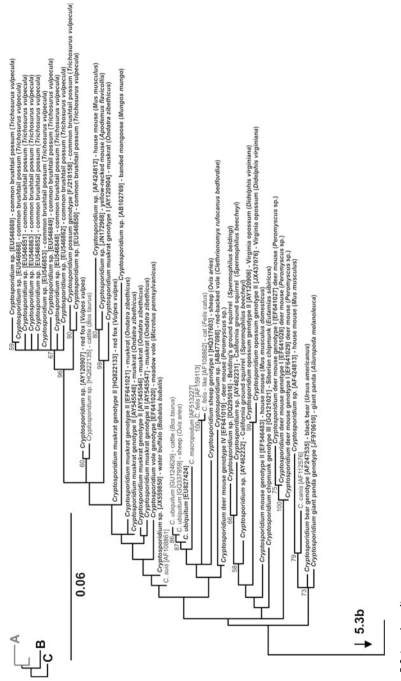
5.6 Egg-Laying Mammals

Monotremata, which is the only order of egg-laying mammals, contains two families, three genera, and five species. All lack teeth as adults, forage on insects or other invertebrates, and are native to Australia and New Guinea.

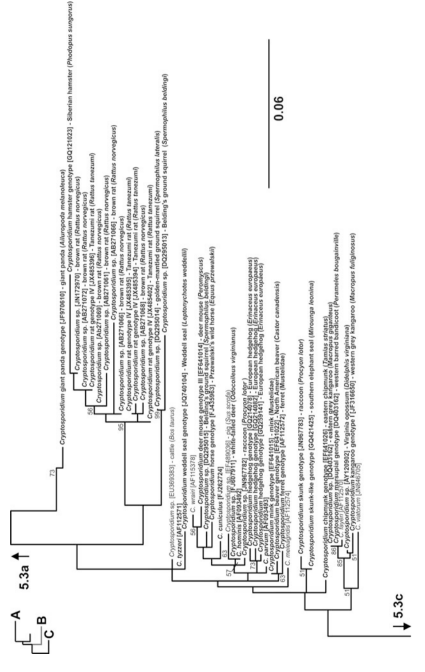
The only report of *Cryptosporidium* from this order has been in a short-beaked echidna (*Tachyglossus aculeatus*), a species that inhabits Australia and New Guinea (O'Donoghue 1995).

5.7 Marsupials

Marsupials comprise seven orders of about 334 species primarily inhabiting Australia and surrounding islands, and South America. *Cryptosporidium* has thus far been detected in species from the orders Dasyuromorphia, Peramelemorphia, and Diprotodontia in Australia, and Didelphimorphia in South America. Members









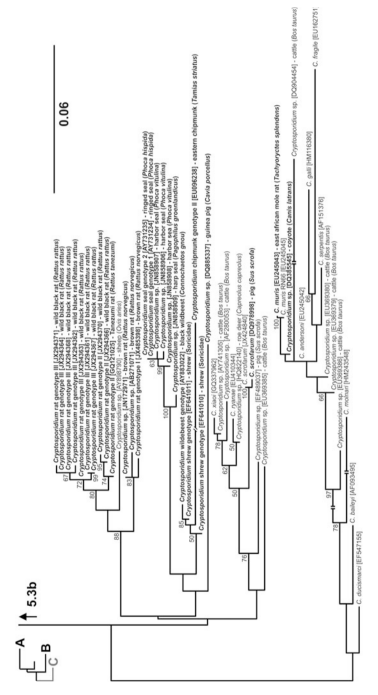


Fig. 5.3 A SSU rDNA-based maximum likelihood (GTRCAT model) tree of Cryptosporidium spp. sequences showing clades of cryptosporidia infecting mammals. Cryptosporidium sp. detected in mammals are in bold. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) is shown at nodes. Bootstrap values <50 are not shown. Interrupted branches have been shortened fivefold

-		1 5	e 1 e
			Host species (scientific name) and GenBank accession
Infraclass	Order	Common names/groups	numbers
Placental mammals	Artiodactyla	Ungulates: sheep, cows, pigs, deer, alpine ibex,	Alpine ibex (Capra ibex) [EF613340]
		water buffalo	Black wildbeest (Connochaetos gnou) [AY883022]
			Deer [AY120910, FJ607911, FJ607928]
			Roe deer (<i>Capreolus</i> <i>capreolus</i>) [HQ822140]
			Sika deer (Cervus nippon) [DQ898159]
	Carnivora		Water buffalo (Bubalus bubalis) [JX559850]
		Deces	Wild boar (Sus scrofa) [U96770]
		Bears	Black bear (Ursus americanus) [AF247535]
			Giant panda (Ailuropoda melanoleuca) [JF970610, JN790957]
		Canines: dog, foxes, coyote	Coyote (Canis latrans) [DQ385545]
			Dog (Canis lupus) [AF112576]
			Red fox (Vulpes vulpes) [AY120907, HQ822133]
			Raccoon dog (Nyctereutes procyonoides viverrinus) [AB104730]
		Cats	Cat (Felis catus) [AF108862, AF112575, AF159113]
		Mongoose	Banded mongoose (Mungos mungo) [AB102769]
		Skunk, raccoons, otter, mink, ferret	Ferret (Mustela) [AF112572] Mink (Mustela vison) [EF428186, EF428187, EF428189, EF641015]
			Raccoon (Procyon lotor) [AY120903,FJ607943, GQ426097, JN967782– JN967784]
			River otter (Lontra canadensis) [DQ288166]
			(continued

 Table 5.9
 Select GenBank accession numbers for *Cryptosporidium* small subunit rRNA gene sequences from mammals. Hosts are separated by taxonomic order and common name groupings

Infraclass	Order	Common names/groups	Host species (scientific name) and GenBank accession numbers
		Seals	Harbor seal (Phoca vitulina) [JN858906–JN858908]
			Harp seal (Pagophilus groenlandicus) [JN858909]
			Hooded seal (Cystophora cristata) [JN858905]
			Southern elephant seal (Mirounga leonina) [GQ421425, JQ740100– JQ740102]
			Ringed seal (Phoca hispida) [AY731234, AY731235]
			Weddell seal (Leptonychotes weddellii) [JQ740103, JQ740104]
	Hyracoidea	Нугах	Rock hyrax (<i>Procavia capensis</i>) [AF161579]
	Erinaceomorpha	Hedgehogs	European hedgehog (Erinaceus europaeus) [GQ214078, GQ214082, [GQ259141]
	Soricomorpha	Shrews and moles	Shrew (Soricidae) [EF641010, EF641011]
	Lagomorpha	Rabbits	European rabbit (Oryctolagus cuniculus) [AY120901, AY273771, FJ262725, GQ865536, HQ397716]
	Perissodactyla	Horse	Prezewalski's wild horse (Equus przewalskii) [FJ435963]
	Non-human Primates	Lemur	Black-and-white colobus (Colobus guereza) [F342450]
			Coquerel's sifaka (Propithecus verreauxi coquereli) [AF442484]
			Gray lagur (Semnopithecus entellus thersites) [EF446673 EF446678]
			Mountain gorilla (Gorilla gorilla gorilla) [JQ837801]
			Olive baboon (Papio anubis [JF681172–JF681174]

Table 5.9 (continued)

(continued)

Infraclass	Order	Common names/groups	Host species (scientific name) and GenBank accession numbers
			Purple-faced lagur (Trachypithecus vetulus philbricki) [EF446679]
			Red colobus (Procolobus rufomitratus) [JF342488– JF342495]
			Rhesus macaque (<i>Macaca</i> <i>mulatta</i>) [HM234173, JX000568–JX000570]
			Toque macaque (<i>Macaca</i> sinica sinica) [EF446672; EF446674–EF446677]
	Rodentia	Mice and rats	Brown rat (<i>Rattus</i> norvegicus) [AB271061, AB271062, AB271064, AB271066, AB271068, AB271069, AB271071–AB271073, JX485398, EU245045, JN172970, JN172971]
			Deer mouse (<i>Peromyscus</i> sp.) [AY120905, EF641014, EF641019, EF641027– EF641030]
			Mouse (species unspecified) [AF108863, AF112571]
			House mouse (Mus musculus) [AF424812, AF424813]
			House mouse (Mus domesticus) [EF546483]
			Tanezumi rat (<i>Rattus</i> <i>tanezumi</i>) [GQ121025, JX485394–JX485396, JX485402]
			Wild black rat (<i>Rattus rattus</i>) [JX29435–JX294371]
			Yellow-necked mouse (Apodemus flavicolis) [JN172968]
		Squirrels: chipmunks, tree and ground squirrels	Belding's ground squirrel (Spermophilus beldingi) [DQ295013, DQ295015, DQ295016]
			California ground squirrel (Spermophilus beecheyi) [AY462231–AY462233, DQ295012]

			Host species (scientific name) and GenBank accession
Infraclass	Order	Common names/groups	numbers
			Golden-mantled ground squirrel (Spermophilus lateralis) [DQ295014]
			Eastern gray squirrel (Sciurus carolinensis) [EU096237]
			Eastern chipmunk (Tamias striatus) [EU096238, EF641026]
			Red squirrel (<i>Sciurus</i> <i>vulgaris</i>) [EU250844, EU250845]
			Siberian chipmunk
			(Eutamias sibiricus) [GQ121021]
		Muskrats, voles, hamster	Boreal red-backed vole (Myodes gapperi) [EF641012, EF641013, or EF641016] ^a
			Meadow vole (Microtus pennsylvanicus) [EF641020]
			Muskrat (Ondatra zibethicus) [AY120904, AY545546– AY545548, EF641021] ^b
			Red-backed vole (<i>Clethrionomys rufocanus</i> <i>bedfordiae</i>) [AB477098]
			Siberian hamster (Phodopus sungorus) [GQ121023]
			Brazilian porcupine
			(Coendou prehensiles) [HM209375]
		Guinea pigs, porcupine	Brazilian porcupine (Coendou prehensiles) [HM209375]
			Guinea pig (Cavia porcellus) [AF115378, DQ885337]
		Beaver	North American beaver (<i>Castor canadensis</i>) [EF641022]
Marsupials	Didelphimorphia	American opossums	Opossum (<i>Didelphis</i> virginiana) [AY120902, AY120906,
			JX437075–JX437079]
			(continued)

Table 5.9 (continued)

Infraclass	Order	Common names/groups	Host species (scientific name) and GenBank accession numbers
	Diprotodontia	Kangaroos, possums, koala	Eastern grey kangaroo (Macropus giganteus) [AF513227, AY237630, DQ403162]
			Common brushtail possum (Trichosurus vulpecula) [EU546848–EU546853, EU546862, EU546863, EU546868 EU546869, FJ218158]
			Red Kangaroo (<i>Macropus</i> <i>rufus</i>) [AF108860, AF112570]
			Western grey kangaroo (Macropus fuliginosus) [JF316650, JF316651]
	Peramelemorphia	Bandicoot	Western barred bandicoot (Perameles bougainville) [DQ403162]

Table 5.9 (continued)

^aIt is not clear from the GenBank entries which of these three sequences was isolated from the boreal red-backed vole

 $^{\mathrm{b}}\mathrm{It}$ is not clear from the GenBank entries which of these three sequences was isolated from the muskrat

of the Notoryctemporphia, which contains two species of marsupial moles (*Notoryctes typhlops* and *Notoryctes caurinus*) that are native to Australia; the Microbiotheria, which contains a single species (*Dromiciops gliroides*) that is native to the southwestern part of South America; and the Paucituberculata, which contains six species of shrew opossum (*Rhyncholestes raphanurus, Lestoros inca, Caenolestes caniventer, Caenolestes condorensis, Caenolestes convelatus,* and *Caenolestes fuliginosus*) that are native to South America, have yet to be identified as hosts for *Cryptosporidium*.

5.7.1 Dasyuromorphia

This order of carnivorous species includes the Tasmanian Devil (*Sarcophilus harrisii*), numbat (*Myrmecobius fasciatus*), and the shrew-like antechinus species.

To date, only the brown antechinus (*Antechinus staurtii*) has been identified as a host for *Cryptosporidium* (Barker et al. 1978).

5.7.2 Peramelemorphia

This order of omnivorous marsupials, which includes bandicoots and bilbies, is found in Australia and New Guinea.

Cryptosporidium sp. and *C. fayeri* have been reported in a southern brown bandicoot (*Isoodon obesulus*) and western-barred bandicoot (*Peremeles bougainville*), respectively (O'Donoghue 1995; Power 2010). *Cryptosporidium muris* was detected in greater bilbies (*Macrotis lagotis*) at a captive breeding colony in Australia. Although some animals cleared the infection within 2 months, others remained infected for 6 months (Warren et al. 2003). Mice trapped in the pens of infected bilbies were positive for *C. muris* and were considered to be the likely source of the infection.

5.7.3 Diprotodontia

Diprotodontia, a large and diverse order of herbivorous marsupials that are native to Australia and surrounding islands, hosts a number of *Cryptosporidium* species and genotypes. Cryptosporidium fayeri, which was first isolated from a koala (Phascolarctos cincereus) (Morgan et al. 1997), has been reported in the red kangaroo (Macropus rufus), eastern grey kangaroo (Macropus giganteus), and yellow-footed rock wallaby (Petrogale xanthopus) (Ryan et al. 2008). Cryptosporidium macropodum (previously marsupial genotype II) has been reported in the red kangaroo, eastern grey kangaroo, western grey kangaroo, and swamp wallaby (Wallabia bicolor). Neither C. fayeri nor C. macropodum are known to cause clinical disease in diprodonts or any other marsupial. The Cryptosporidium kangaroo genotype I, which was identified in the western grey kangaroo (Yang et al. 2011), clusters with the opossum genotype in a neighbour-joining phylogeny of SSU rRNA sequences (Fig. 5.3b). Brushtail possum genotypes I and II have been identified in the brushtail possum (Trichasuris vulpecula) (Power et al. 2003; Hill et al. 2008). Non-genotyped Cryptosporidium sp. have been reported in the red-necked wallaby (Macropus rufogriseus), pademelon (Thylogale billardierii), and koala (Jakob 1992; O'Donoghue 1995).

5.7.4 Didelphimorphia

The didelphimorphs (opossums) are native to the American continent.

The *C. fayeri*-opossum genotype (previously opossum genotype I; this genotype is 99 % similar to *C. fayeri* at the SSU rRNA locus), *C. fayeri* (identified by RFLP analysis), and opossum genotype II have been reported in Virginia opossums (*Didelphis virginiana*) from California and New York, USA (Xiao et al. 2002;

Ziegler et al. 2007b). *Cryptosporidium* also has been detected in the white-eared opossum (*Didelphis albiventris*) (Zanette et al. 2008).

5.8 Placental Mammals

Of the 21 orders of placental mammals, 13 contain species that have been identified as hosts for *Cryptosporidium*. The following eight orders have yet to be identified as hosts:

- **Cingulata and Pilosa**: These two orders totalling 19 species in five families and 18 genera constitute the superorder Xenarthra. Representatives of these orders include armadillos (Cingulata), anteaters (Cingulata), and sloths (Pilosa). Many species are now extinct. The habitats and diets within the group are quite varied. Most are omnivores, but many are insectivores or herbivores.
- **Macroscelididae**, **Afrosordida**, **and Tublidentata**: These are three of the six orders in the superorder Afrotheria. They include tenrecs (Afrosoricida), elephant shrews (Macroscelidia), and aardvarks (Tubulidentata).
- **Pholidota**: This order compromises about eight species in one family and one genus. They are insectivores, inhabiting Africa and southern areas of Asia.
- **Dermoptera**: These comprise a small family of two genera and two arboreal species of gliding mammals, colugos, in Southeast Asia.
- **Scandentia**: These are native of Indonesia and comprise two families, five genera, and 20 species. These small omnivores, commonly referred to as treeshrews, are found in densely forested habitats.

5.8.1 Hyracoidea

Hyracoidea (hyraxes) are in the superorder Afrotheria. They have a digestive system that is similar in function to that of ruminants.

A single species, *C. muris*, has been identified in the rock hyrax (*Procavia capensis*) (Graczyk et al. 1996a; Xiao et al. 1999a).

5.8.2 Sirenia

The order Sirenia, which contains aquatic herbivores such as the dugong and manatee, is most closely related to elephants in the order Proboscidea.

Reports of *Cryptosporidium* in the sirenians have been rare. Hill et al. (1997) described a case of intestinal cryptosporidiosis in a dugnong (*Dugong dugon*) from Queensland, Australia. Three dugongs died as a consequence of the disease and a

fourth, the subject of the study, was humanely euthanized (Hill et al. 1997). Sequence analysis of SSU rRNA and acetyl CoA synthethase genes amplified from preserved tissue specimens revealed that the dugong was infected with C. hominis (Morgan et al. 2000a).

5.8.3 Proboscidea

There have been few studies of *Cryptosporidium* in the herbivorous elephants that comprise the order Proboscidea.

Cryptosporidium sp. has been detected in African elephants (*Loxodonta africana*) at Kruger National Park, South Africa (6/144 positive) (Abu Samra et al. 2011) and the Barcelona Zoo (Gracenea et al. 2002).

5.8.4 Erinaceomorpha

Hedgehogs (Erinceomorpha) contain 24 species in one family (Erinceidae) and ten genera. Some recent phylogenies include this group in the order Soricomorpha (moles and shrews); hedgehogs are similar to shrews in many ways, including an omnivorous diet that may include carrion.

Cryptosporidium can cause clinical and sometimes fatal infections in hedgehogs. Graczyk et al. (1998a) reported fatal cryptosporidiosis in juvenile African hedgehogs (Ateletrix albiventris) housed at the Baltinore Zoo. Cryptosporidium developmental stages were detected in the ileum, jejunum, and colon, and moderate to severe villous atrophy was detected in the ileum and jejunum. Meredith and Milne (2009) reported on a case of cryptosporidiosis in an adult European hedgehog (Erinaceus europaenus) with hemorrhagic diarrhoea. Similar to the juvenile African hedgehogs, moderate to severe villous atrophy was detected in the ileum and jejunum. Sturdee et al. (1999) detected Cryptosporidium oocysts in feces of a free-living European hedgehog from the UK with an estimated infection intensity of 3,000 oocysts per gram. Cryptosporidium spp. was detected in 30.0 % (56/188) of European hedgehogs in Germany (Dyachenko et al. 2010). Cryptosporidium parvum from gp60 subtype families IIa and IIc, and Cryptosporidium hedgehog genotype from gp60 subtype family XIIa (previously VIIa) were identified in positive samples. Although Cryptosporidium positive hedgehogs had diarrhea, any association between a Cryptosporidium sp. and clinical signs was not determined.

5.8.5 Soricomorpha

The Soricomorpha (moles and shrews) represent about 420 species in four families and 44 genera. These are small omnivorous mammals, though invertebrates (especially insects) form a large component of the diet. Many are nocturnal or fossorial. Shrews and moles occupy a variety of habitats, and often live in close association with water.

Cryptosporidium has been reported in the greater white-toothed shrew (*Crocidura russula*), common shrew (*Sorex araneus*), masked shrew (*Sorex cinereus*), northern short-tailed shrew (*Blarina brevicauda*), pygmy shrew (*Sorex minutus*), and Brewer's mole (*Parascalops brewer*) (Siński 1993; Sturdee et al. 1999; Torres et al. 2000; Ziegler et al. 2007a). The shrew genotype (also known as W5) has been identified in 2/5 northern short-tailed shrews in New York (Feng et al. 2007). It is not known to what extent *Cryptosporidium* infection of shrews is associated with clinical disease.

5.8.6 Cetacea

Cetaceans include approximately 90 species in 11 families and 40 genera of dolphins and whales. There are two main groups. The Mysticeti (baleen whales) have baleen rather than teeth for filtering invertebrates from water or bottom sediments. The Odontoceti (toothed whales) have teeth and generally are piscivorous. Most species inhabit marine waters, but a few occur in coastal rivers. Cetaceans are most closely related to the hippopotamus.

Cryptosporidium has been detected in fecal samples from 5.1 % (2/39) of bowhead whales (*Balaena mysticetus*) and 24.5 % (12/49) of North Atlantic right whales (*Eubalaena glacialis*) (Hughes-Hanks et al. 2005). Isolates were not genotyped.

5.8.7 Artiodactyla

The Artiodactyl mammals include about 240 species in 10 families and 89 genera. These even-toed ungulates include three main groups that differ in their digestive system: Suiformes (pigs) have non-ruminating stomachs, Tylopoda (camels) have three-chambered ruminating stomachs, and Ruminantia (deer, antelope, cows, and hippos) have four-chambered ruminating stomachs. They occupy a wide variety of habitats, but many utilize grasslands or savannahs.

5.8.7.1 Pigs

Most research has focused on the domestic pig, and there is relatively limited data regarding *Cryptosporidium* and cryptosporidiosis in wild pigs (described variously as wild boars and feral pigs). Although domestic pigs can host several *Cryptosporidium* taxa, including *C. felis*, *C. hominis*, *C. meleagridis*, *C. muris*, *C. parvum*, *C. scrofarum*, *C. suis*, *C. tyzzeri*, *Cryptosporidium* sp. Eire w65.5, *Cryptosporidium* rat genotype, and *C. suis*-like (Morgan et al. 1999a; Ebeid et al. 2003; Chen and Huang 2007; Kváč et al. 2009a, c), only *C. parvum*, *C. suis*, and *C. scrofarum* have been detected in wild pigs (Table 5.10) (Atwill et al. 1997; Němejc et al. 2012; Garcia-Presedo et al. 2013). The occurrence of non-adapted cryptosporidia in domestic pigs, and their absence from wild pigs, may be a consequence of farming operations that place pigs in close proximity to humans and cattle (e.g. *C. parvum*), cats (*C. felis*), and rodents (*C. tyzzeri*, *C. muris*, and rat genotype).

Similar to domestic pigs, it appears that shedding of *Cryptosporidium* oocysts by wild pigs is associated with age and population density. Atwill et al. (1997) reported a 4.2-fold greater likelihood of oocyst shedding in animals younger than 8 months compared to older animals. Also, wild pigs from sites with ≥ 2.0 wild pigs/km² had approximately a tenfold greater likelihood of shedding *Cryptosporidium* oocysts compared to animals from sites with ≤ 1.9 feral pigs/km².

There is no association between diarrhea in wild pigs and the presence of cryptosporidia, and infection intensity is generally less than 2,000 oocyts per gram of feces (Castro-Hermida et al. 2011a; Němejc et al. 2012).

5.8.7.2 Camels and Llamas

Camels are susceptible to infection with two closely related gastric cryptosporidia, *C. muris* and *C. andersoni*, which prior to 2001 were considered a single species, *C. muris* (Lindsay et al. 2000). A key difference between these species is that only *C. muris* is infectious for neonatal mice under experimental conditions (Lindsay et al. 2000). Therefore, when Anderson (1991) isolated *C. muris*-like oocysts from a camel and demonstrated their infectivity for 2–20-day-old mice, it is probable that the isolate was *C. muris*, and not *C. andersoni*. Molecular studies have subsequently shown that *C. muris* from a Bactrian camel clusters with *C. muris* from mice, a hamster, and a rock hyrax in phylogenies constructed from SSU rRNA, Internal transcribed spacer region 1, and HSP-70 sequences (Xiao et al. 1999a; Morgan et al. 2000c).

A C. and ersoni isolate with oocysts measuring 7.0–7.2 μ m × 5.1–5.2 μ m (shape index: 1.37–1.39) from a 3-year-old Bactrian camel did not infect immunosuppressed or immunocompetent calves, immunosuppressed or immuno-competent Kun-ming mice, or severe combined immunodeficiency mice (Wang et al. 2007).

Cryptosporidium			
taxa	Country	Prevalence	Reference
Cryptosporidium sp.	Spain	11.5 % (20/175)	(Castro-Hermida et al. 2011a)
	Spain	ND	(Gómez et al. 2000)
	USA	ND	(Pereira et al. 1998)
C. parvum ^a	Spain	11.1 % (3/27)	(Garcia-Presedo et al. 2013)
	USA	5.4 % (12/221)	(Atwill et al. 1997)
C. scrofarum	Austria	2.3 % (1/44) ^b	(Němejc et al. 2013)
		13.6 % (6/44) ^c	
	Czech Republic	0 % (0/193) ^b	(Němejc et al. 2012)
		13.0 % (25/193) ^c	
	Czech Republic	1.7 % (4/231) ^b	(Němejc et al. 2013)
		11.3 % (26/231) ^c	
	Poland	2.3 % (3/129) ^b	(Němejc et al. 2013)
		7.8 % (10/129) ^c	
	Slovak Republic	0 % (0/56) ^b	(Němejc et al. 2013)
		1.8 % (1/56) ^c	
	Spain	70.4 % (19/27) ^c	(Garcia-Presedo et al. 2013)
C. suis	Austria	2.3 % (1/44) ^b	(Němejc et al. 2013)
		11.4 % (5/44) ^c	
	Czech Republic	0 % (0/193) ^b	(Němejc et al. 2013)
		13.0 % (25/193) ^c	
	Czech Republic	0.9 % (2/231) ^b	(Němejc et al. 2013)
		10.8 % (25/231) ^c	
	Poland	0 % (0/129) ^b	(Němejc et al. 2013)
		0.8 % (1/129) ^c	-
	Slovak Republic	0 % (0/56) ^b	(Němejc et al. 2013)
	-	3.6 % (2/56) ^c	-
	Spain	18.5 % (5/27) ^c	(Garcia-Presedo et al. 2013)

Table 5.10 Cryptosporidium spp. identified in wild pigs

^aAlthough, the sequence obtained from the feral pig isolate was confirmed as *C. parvum* by the appropriate location of and 100 % sequence homology with two internal probes and three internal primers, in retrospect is impossible to determinate the species and genotypes

^bDetermined using microscopy and

^cDetermined using PCR

ND not determined

Kváč et al. (2008b) infected two out of three lambs and three out of three goat kids with *C. muris* isolate CB03, which originated from a naturally infected Bactrian camel. *Cryptosporidium muris* CB03 was also infectious for Balb/c mice and southern multimammate mice (*Mastomys coucha*). Developmental stages were detected in the *plicae spirales curvature major* in the abomasum of infected lambs and kids. Gastric glands were slightly dilated, and covered with cuboidal and metaplased cells. Infiltration or congestion in the *lamina propria* was not detected, and there was no diarrhea.

Oocysts isolated from rectal samples of camels at a slaughterhouse in Iran had a diameter of 4.56 \pm 0.65 μm (range: 4.20–5.70 μm), which is consistent with the

morphology of intestinal cryptosporidia (Razawi et al. 2009). However, the isolates were not genotyped so this could not be confirmed. Ryan et al. (2003a) identified *C. parvum* in an alpaca (*Lama pacos*), and Gómez et al. (2000) detected *Cryptosporidium* sp. in a guanaco (*Lama guanicoe*).

A study of dromedary camels (*Camelus dromedaries*) in Iran showed a high *Cryptosporidium* prevalence in animals younger than 1 year (20 %) relative to adult camels (6.5 %) (Yakhchali and Moradi 2012). Another study in Iran showed a relatively high prevalence of *Cryptosporidium* sp. (20.3 %; 61/300) in asymptomatic camels (Sazmand et al. 2012).

5.8.7.3 Giraffe and Okapi

Cryptosporidium muris and *Cryptosporidium* sp. have been detected in giraffes (*Giraffa camelopardalis*) from zoos in the Czech Republic and Spain, respectively (Gómez et al. 1996; Kodádková et al. 2010). The *C. muris* isolate was identical to isolates from the rock hyrax and Bactrian camel at the SSU rRNA locus, and was not infectious for Balb/c mice under experimental conditions (Kodádková et al. 2010).

5.8.7.4 Deer and Moose

Cryptosporidium parvum, C. ubiquitum, and the Cryptosporidium deer genotype have been detected at varying rates in deer and moose worldwide. Cryptosporidium *parvum* and C. *ubiquitum* have been reported in 12.5 % (4/32) of swamp deer (Cervus duvauceli) from Nepal (Feng et al. 2012), in a roe deer from England (Robinson et al. 2011), in a red deer (Cervus elaphus) from the Czech Republic (Hajdušek et al. 2004), and in a sika deer (Cervus nippon) from China (Wang et al. 2008a). The Cryptosporidium deer genotype, which is closely related to C. ryanae, has been reported in white-tailed deer (Odoileus virginianus) from the US (Xiao et al. 2002). Cryptosporidium sp. has been detected in 1.3 % (2/149) and 6 % (3/49) of caribou (*Rangifer tarandus*) from Canada and Alaska, respectively; 8.3 % (2/24), 8.8 % (4/35), and 5.0 % (72/360) of white-tailed deer from the US; 7.9 % (3/38) of black-tailed deer (Odocoileus hemionus columbianus) from the US; 25 % (10/40) of tule elk (Cervus canadensis ssp. nannodes) from the US; 1.3 % (3/224), 9.1 % (2/22), and 6.2 % (18/291) of roe deer (*Capreolus capreolus*) from Spain, Poland, and Norway, respectively; 14.4 % (17/118), 0.3 % (1/289), 41.5 % (135/325), and 100 % (2/2) of red deer from Poland, Norway, Ireland, and the US, respectively; and 3.3 % (15/455) of moose (Alces alces) from Norway (Simpson 1992; Fayer et al. 1996; Deng and Cliver 1999; Rickard et al. 1999; Skerrett and Holland 2001; Siefker et al. 2002; Hamnes et al. 2006; Ziegler et al. 2007b; Johnson et al. 2010; Castro-Hermida et al. 2011b). Other deer that have been identified as hosts of Cryptosporidium sp., include the fallow deer (Dama dama), axis deer (Axis axis), barasingha deer (Cervus duvauceli), Eld's deer, muntjac deer, sambar (Rusa

unicolor), Thorold's deer (*Cervus albirostris*), and Père David's deer (*Elaphurus davidianus*) (Heuschele et al. 1986; Sturdee et al. 1999).

5.8.7.5 Bovids

Cryptosporidium has thus far been identified in wild bovid species representing eight of the ten bovid subfamilies (Table 5.11). It has not yet been identified in the Tibetan antelope (*Pantholops hodgsonii*) or grey rhebok (*Pelea capreolus*), which are the sole representatives of the Pantholopinae and Peleinae subfamilies, respectively. Most studies of wild bovid species have been limited to captive animals outside their native habitat.

In a study of cryptosporidiosis in wild bovids, *Cryptosporidium* developmental stages were identified in the small intestine, cecum, spiral colon, and colon (Van Winkle 1985). Four mountain gazelles (*Gazella cuvieri*) at the Munich Zoo with cryptosporidiosis had anorexia and weight loss but no diarrhea (Pospischil et al. 1987). *Cryptosporidium* developmental stages were detected in the abomasum only. The mucosa was hyperplastic and diffusely infiltrated with small lymphocytes, and the mucosal glands were elongated and hypercellular.

Hippopotamus

Cryptosporidium sp. has been detected in a pygmy hippopotamus (*Choeropsis liberiensis*) at the Barcelona Zoo (Gómez et al. 2000).

5.9 Chiroptera

Bats represent a diverse order of flying mammals with over 1,100 species in 18 families and 202 genera. The group is split into the Megachiropterans, which are large, fruit or nectar-eating bats that lack certain ear structures, and the Microchiropterans, which are small, omnivorous or insectivorous species that utilize echolocation and possess associated ear structures. Most (but not all) bats hibernate, and are generally associated with forest habitats.

Despite representing about 20 % of the mammalian fauna, few bat species have been identified as hosts for *Cryptosporidium*. Cryptosporidiosis was diagnosed in a big brown bat (*Eptesicus fuscus*) from Oregon, USA (Dubey et al. 1998). In New York, *Cryptosporidium* sp. and *C. parvum* were identified in a small brown bat (*Myotis lucifugus*) and big brown bat, respectively (Ziegler et al. 2007b). *Cryptosporidium tyzzeri* was detected in a large-footed bat (*Myotus adversus*) from New South Wales, Australia (Morgan et al. 1999b).

Subfamily	Host species (scientific name)	Cryptosporidium taxa	Reference
Aepycerotinae	Impala (Aepyceros melampus)	C. ubiquitum, Crypto- sporidium sp.	(Abu Samra et al. 2011; 2013) (Heuschele et al. 1986)
Alcelaphinae	Black wildebeest (Connochaetes gnou)	Similar to shrew genotype (W5), <i>Cryptosporidium</i> sp.	(Mtambo et al. 1997; Alves et al. 2005)
	Blesbok (Damaliscus dorcas philipsi)	C. ubiquitum	(Ryan et al. 2003a)
	Blue wildebeest (Connochaetes taurinus) ^a	C. parvum	(Gómez et al. 1996; Morgan et al. 1999b)
Antilopinae	Addra gazelle (Gazella dama ruficollis) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
	Blackbuck (Antilope cervicapra)	Cryptosporidium sp.	(Van Winkle 1985; Heuschele et al. 1986)
	Dorcas gazella (Gazella dorcas neglecta) ^a	Cryptosporidium sp.	(Gómez et al. 1996)
	Mountain gazelle (Gazella cuvieri) ^a	Cryptosporidium sp.	(Pospischil et al. 1987)
	Persian gazelle (Gazella subgutturosa) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
	Slender-horned gazelle (Gazella leptoceros) ^a	C. parvum, Crypto- sporidium sp.	(Heuschele et al. 1986; Geurden et al. 2009)
	Springbok (Antidorcas marsupialis) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
	Thomson's gazelle (Eudorcus thomsonii)	Cryptosporidium sp.	(Canestri-Trotti 1989)
Bovinae	African buffalo (Syncerus caffer)	C. ubiquuitum, C. bovis, Crypto- sporidium sp.	(Gómez et al. 1996; Mtambo et al. 1997; Abu Samra et al. 2011; 2013)
	Bison, American (Bison bison) ^a	C. tyzzeri, Cryptospo- ridium sp.	(Alves et al. 2005; Geurden et al. 2009)
	Bison, European (Bison bonasus) ^a Bongo antelope	C. andersoni, Crypto- sporidium sp. C. parvum, Crypto-	(Paziewska et al. 2007) (Ryan et al. 2003a) (Geurden et al. 2009)
	(Tragelaphus eurycerus) ^a	<i>sporidium</i> sp.	(Géurden et al. 2009) (Gómez et al. 2000)

 Table 5.11 Cryptosporidium in wild bovids grouped by subfammily

(continued)

Table 5.11	(continued)
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Subfamily	Host species (scientific name)	Cryptosporidium taxa	Reference
	Eland (Taurotragus oryx) ^a	C. parvum, Crypto- sporidium sp.	(Heuschele et al. 1986; Geurden et al. 2009)
	Lowland anoa (Bubalus depressicornis) ^a	Cryptosporidium sp.	(Gómez et al. 2000)
	Nilgai (Boselaphus tragocamelus) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
	Nyala (Tragelaphus angasi) ^a	C. ubiquitum	(Ryan et al. 2003a)
	Water buffalo (Bubalus bubalis)	C. ryanae variant	(Feng et al. 2012)
	Yak (Bos mutus) ^a	C. parvum, Crypto- sporidium sp.	(Geurden et al. 2009) (Karanis et al. 2007)
	Zebu (Bos primigenius indicus)	<i>C. ryanae</i> variant	(Feng et al. 2012)
Caprinae	Alpine ibex (<i>Capra ibex</i>) ^a	C. ubiquitum	(Karanis et al. 2007)
	Angora goat (Capra hircus)	Cryptosporidium sp.	(Mason et al. 1981)
	Armenian mouflon (Ovis orientalis gmelini) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
	Barbary sheep (Ammotragus lervia) ^a	C. tyzzeri	(Karanis et al. 2007)
	Mouflon sheep (Ovis musimon) ^a	C. ubiquitum, Crypto- sporidium sp.	(Gómez et al. 2000; Ryan et al. 2003a)
	Takin (Budorcas taxicolor) ^a	C. tyzzeri	(Karanis et al. 2007)
	Turkomen markhor (<i>Capra falconeri</i>) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
Hippotraginae	Urial (Ovis orientalis) Addax (Addax nasomaculatus)	Cryptosporidium sp. Cryptosporidium sp.	(Ducatelle et al. 1983) (Van Winkle 1985) (Heuschele et al. 1986)
	Fringe-eared oryx (Oryx beisa callotis)	Cryptosporidium sp.	(Van Winkle 1985)
	Sable antelope (Hippotragus niger) ^a	C. parvum, Crypto- sporidium sp.	(Hajdušek et al. 2004) (Van Winkle 1985; Heuschele et al. 1986)
	Scimitar-horned oryx (Oryx dammah)	Cryptosporidium sp.	(Van Winkle 1985) (Heuschele et al. 1986)

(continued)

Subfamily	Host species (scientific name)	Cryptosporidium taxa	Reference
	White antelope (Addax nasomaculatus)	Cryptosporidium sp.	(Van Winkle 1985)
Reduncinae	Nile lechwe (Kobus megaceros) ^a	Cryptosporidium sp.	(Heuschele et al. 1986)
_	Waterbuck (Kobus ellipsiprymnus)	Cryptosporidium sp.	(Gómez et al. 1996)

Table 5.11 (continued)

^aWild bovids under captive condition

5.10 Perissodactyla

The Perissodactyl mammals include about 16 species in three families and six genera. These are odd-toed ungulates that include horses, tapirs and rhinoceroses. The digestive system is much simpler than that of the ruminants, with cellulose broken down by fermentation in the latter portion of the gut. They occupy mainly grassland or savannah habitat.

5.10.1 Equids

Equids include the zebra, ass, kiang, and wild horse. *Cryptosporidium* was detected in 28 % (7/25) of zebras (*Equus zebra*) at Mikumi National Park in Tanzania (Mtambo et al. 1997). *Cryptosporidium* horse genotype was identified in a Przewalski's wild horse foal at the Prague Zoo (Ryan et al. 2003a). Although, no other cryptosporidia have been reported in wild Equids, domestic horses have been identified as hosts for *C. parvum*, *Cryptosporidium* horse genotype, and *Cryptosporidium* hedgehog genotype (Laatamna et al. 2013).

5.10.2 Rhinoceros and Tapirs

Cryptosporidium sp. has been reported in a southern white rhinoceros (*Ceratotherium simum simum*), rhinoceros (*Rhinoceros unicornis*), and a South American tapir (*Tapirus terrestris*) at the Barcelona Zoo (Wang and Liew 1990; Gómez et al. 1996, 2000).

5.11 Carnivora

5.11.1 Fin-Footed Mammals (Pinnipedia)

Cryptosporidium muris, which was isolated from a ringed seal (*Phoca hispida*) in Northern Quebec, Canada (Santín et al. 2005), remains the only species identified in Pinnipeds to date. Cryptosporidium sp. has been reported in the ringed seal (Phoca hispida), harbor seal (Phoca vitulina), harp seal (Pagophilus groenlandicus), seal (Leptonychotes weddellii), California sea lion (Zalophus Weddell californianus), grey seal (Halichoerus grypus), and southern elephant seal (Mirounga leonina) (Deng et al. 2000; Hughes-Hanks et al. 2005; Santín et al. 2005; Bogomolni et al. 2008; Rengifo-Herrera et al. 2011, 2013; Bass et al. 2012). In a neighbor-joining phylogeny, SSU rRNA gene sequences from the ringed seal (seal genotype I, GenBank accession no. AY731234 and seal genotype II, GenBank accession no. AY731235), harbor seal (GenBank accession nos. JN858906–JN858908), harp seal (GenBank accession no. JN858909), and hooded seal (GenBank accession no. JN858905) form a separate clade that is most closely related to a group containing sequences from the shrew (Soricidae), wildebeest (Connochaetes gnou), and eastern chipmunk (Tamias striatus) (Fig. 5.3c). Isolates from southern elephant seals (*Mirounga leonina*) on the west coast of the Antarctic Peninsula shared 99 % identity with a skunk genotype and C. tyzzeri at the SSU locus and are named southern elephant seal genotype (Rengifo-Herrera et al. 2011, 2013). Isolates from the Weddell seal are most closely related to the ferret genotype and C. parvum (99 % identity at the SSU locus). Cryptosporidium sp. detected in California sea lions (Zalophus californianus) from the northern California coastal area share 98 % identity with C. parvum at the COWP locus (Deng et al. 2000).

5.11.2 Domestic Dogs

There have been few molecular studies of *Cryptosporidium* in dogs (Table 5.12). *Cryptosporidium canis* is the most frequently identified species in dogs (Abe et al. 2002) and it appears to be relatively host adapted. Other species found in dogs include *C. parvum* (Hajdušek et al. 2004), *C. muris* (Ellis et al. 2010), and *C. meleagridis* (Hajdušek et al. 2004). *Cryptosporidium muris* caused chronic gastritis in an 18-month male, mixed-breed dog (Ellis et al. 2010). It appeared that the dog had a concurrent infection with *Helicobacter*.

In a study of parasites infecting sled dogs in Poland, Bajer et al. (2011) found a higher prevalence of *Cryptosporidium* in dogs infected concurrently with *Giardia* (35.5 %) than in dogs without *Giardia* (2.7 %). In contrast, *Cryptosporidium* prevalence was three times lower in dogs infected with nematodes. Other studies have shown a higher prevalence of *Cryptosporidium* in dogs with diarrhea (Mirzaei

Table 5.12 Prevalence (able 5.12 Prevalence of Cryptosporidium in dogs from different countries		
Country	Prevalence	Cryptosporidium taxa	Reference
Australia	10.9 % (54/493)	Cryptosporidium sp.	(Bugg et al. 1999)
Argentina	2.2 % (48/2,193)	Cryptosporidium sp.	(Fontanarrosa et al. 2006)
	1.0%(1/100)	Cryptosporidium sp.	(Soriano et al. 2010)
Brazil	1	C. canis	(Thomaz et al. 2007)
	9.6 % (43/450)	Cryptosporidium sp.	(Lallo and Bondan 2006)
	4.9 % (4/81)	Cryptosporidium sp.	(Mandarino-Pereira et al. 2010)
Canada	74.3 % (52/70)	Cryptosporidium sp.	(Shukla et al. 2006)
	3.2 % (5/155)	Cryptosporidium sp.	(Himsworth et al. 2010)
Costa-rica	1.7 % (1/58)	Cryptosporidium sp.	(Scorza et al. 2011)
Czech Republic	ND	C. meleagridis, C. parvum	(Hajdušek et al. 2004)
Iran	2.0 % (11/548)	Cryptosporidium sp.	(Mirzaei 2012)
Iran	5.2 % (4/77)	Cryptosporidium sp.	(Beiromvand et al. 2013)
Japan	3.9 % (3/77)	C. canis	(Yoshiuchi et al. 2010)
Korea	9.7 % (25/257)	Cryptosporidium sp.	(Kim et al. 1998)
Netherlands	8.6 % (13/152)	Cryptosporidium sp.	(Overgaauw et al. 2009)
Poland	13 % (14/108)	Cryptosporidium sp.	(Bajer et al. 2011)
Spain	5.1 % (4/79)	Cryptosporidium sp.	(Dado et al. 2012)
	6.3 % (32/505)	Cryptosporidium sp.	(Gracenea et al. 2009)
USA	10.2 % (5/49)	Cryptosporidium sp.	(Jafri et al. 1993)
	17 % (17/100)	Cryptosporidium sp.	(Juett et al. 1996)
	3.8 % (5/130)	Cryptosporidium sp.	(Hackett and Lappin 2003)
	2.0 % (4/200)	Cryptosporidium sp.	(el-Ahraf et al. 1991)
	2.3 % (3/129)	C. canis	(Wang et al. 2012)
	ND	C. muris	(Ellis et al. 2010)
	12.0 % (6/50) with diarrhea;	Cryptosporidium sp.	(Tupler et al. 2012)
	2.0 % (1/50) without diarrhea		
	2.5 % (3/120)	Cryptosporidium sp.	(McKenzie et al. 2010)

 Table 5.12
 Prevalence of Cryptosporidium in dogs from different countries

ND not determined

2012; Tupler et al. 2012), and dogs less than 1-year-old (Mirzaei 2012). It is not known if the association with diarrhea is dependent on the *Cryptosporidium* species/genotype causing the infection.

Cryptosporidium canis does occasionally cause human disease (Leoni et al. 2006; Hijjawi et al. 2010), and a UK study found that dogs are the most likely companion animals to shed *Cryptosporidium* (46/139) (Smith et al. 2009).

5.11.3 Raccoon Dogs

Cryptosporidium parvum has been identified in a raccoon dog (*Nyctereutes procyonoides viverrinus*) at a zoo in Osaka, Japan (Matsubayashi et al. 2005).

5.11.4 Foxes

Cryptosporidium has been reported in the two genera of true foxes: *Vulpes* and *Urocyon*. Current (1989) identified *Cryptosporidium* sp. in the gray fox (*Urocyon cinereoargenteus*), which is native to southern parts of North America and northern parts of South America. *Cryptosporidium canis* fox genotype (n = 4), *C. canis* (n = 1), and muskrat genotype I (n = 1) were identified in 6/76 *Cryptosporidium* positive samples from unidentified fox species in Maryland, USA (Zhou et al. 2004a). *Cryptosporidium* was detected in 38.7 % (24/62) and 8.1 % (10/124) of red foxes (*Vulpes vulpes*) from the Slovak Republic and Ireland, respectively (Nagano et al. 2007; Ravaszová et al. 2012). Two of the isolates from the study in Ireland were identified as *C. parvum* by sequence analysis of the SSU rRNA and gp60 loci. In contrast to the relatively high prevalence reported in the Slovak Republic, *Cryptosporidium* sp. was detected in only 2.2 % (6/269) and 8.7 % (2/23) of red foxes from Norway and England, respectively (Sturdee et al. 1999; Hamnes et al. 2007).

5.11.5 Wolves

Two large Canadian studies were consistent in showing a relatively low prevalence of *Cryptosporidium* in gray wolves. The prevalence was 1.7 % in a study of 1,558 fecal samples from wolves in British Columbia (Bryan et al. 2012), and 1.2 % in a study of 601 fecal samples from wolves in Manitoba (Stronen et al. 2011). In contrast, 54.9 % of fecal samples (28/51) from wolves in northeast Poland were positive for *Cryptosporidium* (Kloch et al. 2005). A subsequent study in the same region of Poland reported a prevalence of 35.7 %, and *C. parvum* was

identified by sequence analysis of the COWP gene (GenBank accession number: AF266273) (Paziewska et al. 2007).

5.11.6 Coyotes

Cryptosporidium has been detected in 27 % (6/22) and 26.3 % (5/19) of fecal samples from coyotes (*Canis latrans*) in northeastern Pennsylvania (Trout et al. 2006) and New York (Ziegler et al. 2007a), respectively. One isolate from the Pennsylvania study shared 99.7 % sequence identity with *C. muris* at the SSU rRNA locus, and the remaining five isolates were identical to the *C. canis* coyote genotype (Trout et al. 2006). In a study of coyotes in southern Alberta and Saskatchewan, Canada, *Cryptosporidium* was not found during winter, and the prevalence was 17.4 % during summer. SSU rRNA sequences from two of the isolates were identified as the *C. canis* coyote genotype (Thompson et al. 2009).

5.11.7 Bears

Cryptosporidium has been detected in three of the five genera in the family Ursidae (*Ailuropoda, Helarctus*, and *Ursus*). There have been no reports in the sloth bear (*Melursus ursinus*) or spectacled bear (*Tremarctos ornatus*), which are the only extant representatives of their genera.

Cryptosporidium has been reported in two captive Malayan sun bears (*Helarctos malayanus*) at a zoological park in Taiwan (Wang and Liew 1990).

Cryptosporidium has been detected in black bears (*Ursus americanus*), brown bears (*Ursus arctos*), and polar bears (*Ursus maritimus*) in the genus *Ursus* (Siam et al. 1994; Duncan et al. 1999; Xiao et al. 2000; Ravaszová et al. 2012). Xiao et al. (2000) identified the *Cryptosporidium* bear genotype in a black bear and showed that it is most closely related to *C. canis* at the SSU rRNA and HSP-70 loci.

Karanis et al. (2007) detected *C. tyzzeri* (previously mouse genotype I) in a lesser panda (*Ailurus fulgens*). An isolate from a giant panda (*Ailuropoda melanoleuca*) was named the *Cryptosporidium* giant panda genotype based on sequences of the SSU rRNA, actin, COWP, and HSP-70 genes. Although the study has not been published, the sequences are available in GenBank under the accession numbers JF970610, JN969985, JN588570, and JN588571.

5.11.8 Raccoons

A number of studies have identified *Cryptosporidium* in raccoons (*Procyon lotor*) (Snyder 1988; Perz and Le Blancq 2001; Zhou et al. 2004a; Ziegler et al. 2007a;

Chavez et al. 2012). Zhou et al. (2004a) identified the skunk genotype in 3.9 % (2/51) of raccoon samples collected in the Chesapeake Bay area of Maryland, USA. *Cryptosporidium* was detected in 11 out of 44 fecal samples from raccoons in Colorado, USA. Five out of six of the isolates that were genotyped were identified as the skunk genotype, and one was identified as *C. parvum* (Chavez et al. 2012).

Cryptosporidium also causes clinical disease in raccoons. A juvenile raccoon, estimated to be between 6 and 12-months-old was found in a moribund condition in Fort Collins, Colorado. The animal was emaciated and dehydrated, and had diarrhea and a mucoid oculonasal discharge. *Cryptosporidium* sp. was identified on intestinal villi (Martin and Zeidner 1992).

5.11.9 Ferrets

Rehg et al. (1988) reported cases of subclinical cryptosporidiosis in 40 % of ferrets at an animal research facility in Tennessee. Subsequent studies identified the *Cryptosporidium* ferret genotype in the black-footed ferret (*Mustela nigripes*) and domestic ferret (*Mustela putorius furo*) in the USA and Japan (Xiao et al. 1999b; Sulaiman et al. 2000; Abe and Iseki 2003). The ferret genotype is genetically closely related to *C. parvum*, sharing 99 %, 98 %, 98 %, and 97 % identity at the SSU rRNA, actin, COWP, and HSP-70 genes, respectively. Ferret genotype isolates from the USA (Sulaiman et al. 2000) and Japan (Abe and Iseki 2003) have identical sequences, suggesting that this genotype is geographically conserved.

Cryptosporidium also causes clinical disease in ferrets. Gómez-Villamandos et al. (1995) reported an outbreak of fatal cryptosporidiosis in captive 11–12 month-old non-pregnant female ferrets (*Mustela putorius furo*) kept on a goat farm. Animals died 48–72 h after the onset of signs, which included anorexia, depression, and diarrhea. *Cryptosporidium* developmental stages were identified in feces and tissue, but the species/genotype causing the infection was not identified.

5.11.10 Otters

Cryptosporidium has been detected in the European or wild otter (*Lutra lutra*), river otter (*Lontra canadensis*), and sea otter (*Enhydra lutris nereis*) (Feng et al. 2007; Gaydos et al. 2007; Méndez-Hermida et al. 2007; Oates et al. 2012). In Spain, *Cryptosporidium* oocysts were detected in 17 out of 437 fecal samples from wild otters (Méndez-Hermida et al. 2007). In the USA, nine river otters from the Puget Sound Georgia Basin were positive for the *Cryptosporidium* mink genotype (Gaydos et al. 2007), and a river otter in New York was positive for the *Cryptosporidium* sp. was detected in a single sea otter in California (Oates et al. 2012).

5.11.11 Mink and Martens

Rademacher et al. (1999) isolated Cryptosporidium oocysts measuring 3-5 µm from captive beech martens (Martes foina) with episodes of diarrhea. Cryptosporidium oocysts (4.5–5.5 µm) were also detected in 24.2 % (8/33) of fecal samples from captive American minks (Mustela vison) in Spain. The animals did not show clinical signs associated with cryptosporidiosis. Three isolates had COWP sequences that differed from the ferret genotype by a single nucleotide polymorphism (Gómez-Couso et al. 2007). In China, eight Cryptosporidium isolates were obtained from 469 fecal samples of American minks (Mustela vison) originated from a farm in Hebei Province in China. Six of the eight Cryptosporidium-positive samples contained a novel genotype, the Cryptosporidium mink genotype, that was most closely related to the Cryptosporidium ferret genotype (Wang et al. 2008b). In New York, one of four mink (Mustela vison) in a watershed study was positive for Cryptoporidium mink genotype, and an ermine (Mustela erminea) had a mixed infection with Cryptosporidium shrew genotype (also known as W5 genotype) and W18 genotype (Feng et al. 2007). Also in New York, Ziegler et al. (2007a) detected Cryptosporidium sp. in one of three ermine (Mustela erminea) and in one of 58 American mink (Musela vison).

5.11.12 Skunk

Cryptosporidium has been detected in the striped skunk (*Mephitis mephitis*) in New York, USA (Perz and Le Blancq 2001; Ziegler et al. 2007a), and the *Cryptosporidium* skunk genotype has been described from this host (Xiao et al. 2002; Feng et al. 2011). *Cryptosporidium* prevalence in the striped skunk in New York was 14 % (12/86) (Ziegler et al. 2007a).

5.11.13 Badger

Cryptosporidium has been reported in 15.4 % (4/26) of European badgers (*Meles meles*) in England (Sturdee et al. 1999). The *Cryptosporidium* species/genotype infecting badgers is not yet known.

Country	Prevalence	Cryptosporidium taxa	Reference
Argentina	2.2 % (48/2,193)	Cryptosporidium sp.	(Fontanarrosa et al. 2006)
Canada	7.1 % (5/70)	Cryptosporidium sp.	(Shukla et al. 2006)
Germany	5.3 % (1/19)	Cryptosporidium sp.	(Sotiriadou et al. 2013)
Japan	12.7 % (7/55)	C. felis	(Yoshiuchi et al. 2010)
Korea	9.7 % (25/257)	Cryptosporidium sp.	(Kim et al. 1998)
Netherlands	8.6 %	Cryptosporidium sp.	(Overgaauw et al. 2009)
Spain	6.3 % (32/505)	Cryptosporidium sp.	(Gracenea et al. 2009)
UK	5.9 % (3/51)	Cryptosporidium sp.	(Smith et al. 2009)
USA	5.3 % (11/206)	Cryptosporidium sp.	(Hill et al. 2000)
	4.7 % (16/344)	Cryptosporidium sp.	(Mekaru et al. 2007)
	3.8 % (10/263)	Cryptosporidium sp.	(Spain et al. 2001)
	6.4 % (11/173)	Cryptosporidium sp.	(Nutter et al. 2004)
	12.0 % (30/250) – IFA	Cryptosporidium sp.	(Ballweber et al. 2009)
	4.8 % (12/250) – PCR	C. felis	

Table 5.13 Prevalence of Cryptosporidium in cats from different countries

5.11.14 Banded Mongoose

The banded mongoose (*Mungos mungo*) has been identified as a host of a *Cryptosporidium* genotype that is most closely related to the bear genotype at the SSU rRNA and HSP-70 loci (Abe et al. 2004).

5.11.15 Domestic Cat

Two *Cryptosporidium* species, *C. felis* and *C. muris*, have been reported in domestic cats (*Felis catus*). *Cryptosporidium felis* was first identified in cats in Japan (Iseki 1979) and has since been reported in cats worldwide. *Cryptosporidium muris* has been reported in cats less frequently (Santín et al. 2006; Pavlásek and Ryan 2007; FitzGerald et al. 2011). FitzGerald et al. (2011) reported a *C. felis/C. muris* coinfection in a cat with chronic diarrhea. *Cryptosporidium muris* was identified within the glands of the gastric mucosa. Interestingly, the cat appeared to be persistently infected with *C. muris*, but not *C. felis. Cryptosporidium felis* has been identified as an infrequent cause of human cryptosporidiosis (Pedraza-Diaz et al. 2001; Caccio et al. 2002; Caccio 2005; Leoni et al. 2006). The prevalence of *Cryptosporidium* sp. in cats from different countries is shown in Table 5.13.

5.11.16 Wild Cats

Cryptosporidium tyzzeri has been identified in black leopards (*Pantera pardus*) (Karanis et al. 2007). *Cryptosporidium* sp. also has been reported in bobcats (*Lynx rufus*) in California (Carver et al. 2012) and New York, USA (Ziegler et al. 2007a).

5.12 Lagomorpha

Lagomorpha (rabbits, hares and pikas) includes about 87 species in two families and 12 genera. The phylogeny of the lagomorphs is not well understood. They have four upper incisors, and all are herbivorous. All produce both a hard fecal pellet and a soft, grease-like pellet derived from the cecum. The soft pellet is reingested, and contains a much higher concentration of vitamins and minerals than the uningested hard pellet. Lagomorphs occupy both forested and grassland areas.

Rabbits are the major host of *C. cuniculus* (previously *Cryptosporidium* rabbit genotype). *Cryptosporidium cuniculus* is closely related to *C. hominis*, sharing 99.5, 100, 99.9, and 99.5 % nucleotide sequence identity at the SSU rRNA, COWP, actin, and HSP-70 loci, respectively (Robinson et al. 2010). It emerged as a human pathogen in 2009 when it was identified as the cause of a waterborne cryptosporidiosis outbreak in Northhamptonshire, England (Robinson et al. 2010). A subsequent study identified *C. cuniculus* in 1.2 % of 3,030 samples from human cryptosporidiosis cases in the UK during the period 2007–2008 (Chalmers et al. 2011). In a comprehensive review of the literature on *Cryptosporidium* in rabbits, which will not be duplicated here, Robinson and Chalmers (2010) noted that in studies employing genotyping tools, *C. cuniculus* was the only species/genotype detected. More recently, *C. cuniculus* was the only species detected in rabbits in Australia (Nolan et al. 2010, 2013) and China (Shi et al. 2010; Zhang et al. 2012), and prevalence was highest in younger (1–3-month-old) rabbits (Shi et al. 2010).

5.13 Rodentia

Rodents comprise about 40 % of the mammalian diversity, with over 2,200 species in 31 families and 481 genera. Five suborders are typically recognized: Myomorpha (mice, rats, gerbils, and relatives), Sciuromorpha (tree squirrles and relatives), Castorimorpha (beavers, gophers, and relatives), Anumaluromorpha (scaly-tailed squirrels, and springhares), and Hystricomorpha (gundis, capybaras, and relatives). With such diversity, rodents occupy a wide range of habitats, and generalizations regarding their natural history are difficult. All rodents have two upper and two lower incisors that grow continuously (and are self-sharpening) but lack canines. Most species are herbivorous or omnivorous.

5.13.1 Muridae

Old world mice and rats comprise the subfamily Murinae in the largest mammal family, the Muridae. Gerbils, which also are members of the family Muridae, have yet to be identified as a natural host for *Cryptosporidium*, although they are susceptible to experimental infections (Baishanbo et al. 2005; Kváč et al. 2009d).

Four Cryptosporidium species (C. tyzzeri, C. muris, C. parvum, and C. scrofarum) and six genotypes (C. suis-like genotype, rat genotypes I-IV and mouse genotype II) have been reported as natural infections in several species of Mus, Apodemus, and Rattus in Australia, China, the Czech Republic, Germany, Kenya, New Zealand, the Philippines, Poland, Portugal, Spain, the UK, and the US (Table 5.13). The C. suis-like genotype and C. scrofarum were reported in rats in the Philippines; C. suis-like genotype was found exclusively in Asian house rats, and C. scrofarum was additionally identified in brown rats (Ng-Hublin et al. 2013). The C. suis-like genotype, which is 99.7 % similar to C. suis at the SSU rRNA locus, has additionally been reported in cattle and humans (Ong et al. 2002; Langkjær et al. 2007; Robinson et al. 2011). Cryptosporidium scrofarum is a pig-adapted species that also has been reported in cattle and humans (Kváč et al. 2009b; Ng et al. 2011). Cryptosporidium muris, a rodent adapted species, has been detected the domestic mouse (Mus musculus domesticus) in the UK (Chalmers et al. 1997), the Algerian mouse (*Mus spretus*) in Spain (Torres et al. 2000), and the East African mole rat (Tachyoryctes splendens) in Kenya (Kváč et al. 2008b). Cryptosporidium mouse genotype II has been identified in 11 domestic mice in Australia (Foo et al. 2007). Rat genotypes I-IV have been reported in various rat species in Australia, China, and the Philippines (Table 5.14). Although C. tyzzeri (previously mouse genotype I) has been reported in the yellownecked mouse, voles, snakes, and rats (Morgan et al. 1998, 1999b; Bajer et al. 2003; Xiao and Ryan 2004; Karanis et al. 2007), the house mouse is considered the major host for this species (Ren et al. 2012).

The fate of the house mouse (*Mus musculus*) has been intimately connected to humans since the establishment of a commensal relationship at the dawn of civilization. Two house mouse subspecies, *Mus musculus musculus* and *Mus musculus domesticus*, diverged approximately 0.5 million years ago in the Middle East (Geraldes et al. 2008; Duvaux et al. 2011; Auffray and Britton-Davidian 2012; Bonhomme and Searle 2012). *Mus m. musculus* spread to northern Eurasia and migrated westward through Europe. *Mus m. domesticus* expanded westward through Asia Minor to southern and western Europe, northern Africa, and the New World (Boursot et al. 1993; Guénet and Bonhomme 2003; Rajabi-Maham et al. 2008; Duvaux et al. 2011; Auffray and Britton-Davidian 2012; Bonhomme and Searle 2012). About 6,000 years ago, westward migrating

Host (scientific		
name)	Cryptosporidium taxa	Reference
Algerian mouse (Mus spretus)	C. muris, C. parvum	(Torres et al. 2000)
Asian house rat (Rattus tanezumi)	C. scrofarum, C. tyzzeri, C. suis-like, rat genotypes II–IV	(Lv et al. 2009; Ng-Hublin et al. 2013)
Black rat (Rattus rattus)	C. parvum, rat genotypes II–III, Cryptosporidium sp.	(Miyaji et al. 1989; Yamura et al. 1990; Webster and Macdonald 1995; Chilvers et al. 1998; Torres et al. 2000; Paparini et al. 2012)
Brown rat (Rattus norvegicus)	C. tyzzeri, C. scrofarum, C. muris, rat genotypes I–IV, Cryptosporidium sp.	(Iseki 1986; Miyaji et al. 1989; Yamura et al. 1990; Quy et al. 1999; Kimura et al. 2007; Lv et al. 2009; Ng-Hublin et al. 2013)
House mouse (Mus musculus)	C. tyzzeri, C. muris, mouse genotype II, Cryptosporidium sp.	(Klesius et al. 1986; Chalmers et al. 1997; Chilvers et al. 1998; Xiao et al. 1999b; Foo et al. 2007; Ziegler et al. 2007a; 2007b; Lv et al. 2009; Kváč et al. 2013b)
Japanese field mouse (Apodemus speciosus)	Cryptosporidium sp.	(Nakai et al. 2004; Hikosaka and Nakai 2005)
Striped field mouse (Apodemus agrarius)	Cryptosporidium sp.	(Siński et al. 1998)
East African mole rat (Tachyoryctes splendens)	C. muris	Kváč et al. 2008b
Wood mouse (Apodemus sylvaticus)	C. parvum, C. muris, Cryptosporidium sp.	(Chalmers et al. 1997; Torres et al. 2000; Hajdušek et al. 2004)
Yellow-necked mouse (Apodemus flavicollis)	C. tyzzeri, C. parvum, Cryptosporidium sp.	(Siński 1993; Torres et al. 2000; Bajer et al. 2002; 2003; Bednarska et al. 2003)

Table 5.14 Cryptosporidium spp. identified in species of old world mice and rats

M. m. musculus and eastward-migrating *M. m. domesticus* reestablished contact in central Europe, and formed a stable, narrow (approximately 20 km wide) hybrid zone that stretches 2,500 km from Norway to the Black Sea (Macholán et al. 2003; Jones et al. 2011; Ďureje et al. 2012). This hybrid zone affords a rare opportunity to study speciation through the interactions between two subspecies that have been separated for about 500,000 years.

The hybrid zone was recently used to test the hypothesis that the house mouseadapted species *C. tyzzeri* is coevolving with subspecies of *M. musculus*. *Cryptosporidium tyzzeri* isolates from naturally infected *M. m. musculus* and *M. m. domesticus* in the hybrid zone differed genetically, morphometrically, and biologically (Kváč et al. 2013b), which supports the coevolution hypothesis.

5.13.2 Cricetidae

Cricetidae, which includes muskrats, deer mice, voles, cotton rats, and hamsters, are the second largest family of mammals with greater than 681 species in 130 genera. To date, members of this family have been reported as hosts for four *Cryptosporidium* species (*C. muris, C. parvum, C. andersoni*, and *C. ubiquitum*) and ten genotypes (vole genotype, muskrat genotypes I and II, W12, hamster genotype, chipmunk genotype I, and deer mouse genotypes I–IV) (Table 5.15).

Within the Cricetidae, *C. muris* and *C. andersoni* have been detected only in hamsters (Ryan et al. 2003a; Lv et al. 2009). *Cryptosporidium andersoni*, a ruminant adapted species, was detected in 5.9 % (8/136) of pet hamsters from three different species in China (Lv et al. 2009), suggesting that hamsters may be a significant host for *C. andersoni*, at least in that country. *Cryptosporidium muris* and *C. parvum* are the only *Cryptosporidium* species or genotypes known to infect members of both the Muridae and Cricetidae family. These species also are found in the family Sciuridae (see below).

5.13.3 Sciuridae

Five species (*C. andersoni*, *C. baileyi*, *C. muris*, *C. parvum*, and *C. ubiquitum*) and ten genotypes have been detected in the family Sciuridae, which includes squirrels and chipmunks (Table 5.16). In addition to *C. parvum* and *C. muris*, which are found in Muridae, Cricetidae, and Sciuridae, two species (*C. andersoni* and *C. ubiquitum*) and two genotypes (chipmunk genotype I and deer mouse genotype III) are common to the Cricetidae and Sciuridae (Fig. 5.4).

5.13.4 Hystricomorpha

Hystricomorphs include the guinea pig, capybara, and chinchilla, which were the first rodents to colonize South America about 40 million years ago, probably from Africa, and they remained the only rodents in South America until the arrival of murids about five million years ago (Antoine et al. 2012).

Table 5.17 shows the *Cryptosporidium* sp. that have been reported in Hystricomorphs. *Cryptosporidium wrairi* is found only in guinea pigs (Vetterling et al. 1971; Lv et al. 2009), and the prevalence of infection can be as high as 85 % in pet guinea pigs (Lv et al. 2009). Guinea pigs also host a guinea pig genotype

Host (scientific		Deferre
name)	Cryptosporidium taxa	Reference
Bank vole (Myodes glareolus)	Cryptosporidium sp.	(Siński 1993; Laakkonen et al. 1994; Chalmers et al. 1997; Bull et al. 1998; Siński et al. 1998; Torres et al. 2000; Bajer et al. 2002; Bednarska et al. 2007)
Campbell hamster (Phodopus campbelli)	C. andersoni, C. muris, C. parvum	(Lv et al. 2009)
Common vole (Microtus arvalis)	Cryptosporidium sp.	(Siński et al. 1998; Bednarska et al. 2007)
Cotton rat (Sigmodon hispidus)	Cryptosporidium sp.	(Elangbam et al. 1993)
Deer mouse or white-footed mice (Peromyscus spp.)	C. parvum, C. ubiquitum, muskrat genotype II (W16), chipmunk genotype I (W17), deer mouse genotypes I–IV	(Perz and Le Blancq 2001; (Xiao et al. 2002; Feng et al. 2007; Ziegler et al. 2007a, b)
Golden hamster (Mesocricetus auratus)	C. andersoni, C. muris, C. parvum	(Ryan et al. 2003a; Lv et al. 2009)
Meadow vole (Microtus pennsylvanicus)	Vole genotype (W15), muskrat genotype II (W16)	(Feng et al. 2007; Ziegler et al. 2007a; 2007b)
Muskrat (Ondatra zibethicus)	Muskrat genotype I (W7), muskrat genotype II (W16)	(Siński et al. 1998; Perz and Le Blancq 2001; Xiao et al. 2002; Zhou et al. 2004a; Feng et al. 2007; Ziegler et al. 2007b)
Siberian hamster (Phodopus sungorus)	C. andersoni, C. muris, C. parvum, hamster genotype	(Lv et al. 2009)
Southern red-backed vole (Myodes gapperi)	<i>C. parvum</i>, vole cluster, muskrat genotype I (W7),Muskrat genotype II (W16), W12	(Feng et al. 2007; Ziegler et al. 2007a, b)

 Table 5.15
 Cryptosporidium spp. identified in the Cricetidae

(GenBank Accession nos. DQ885337 and DQ885338) that is only 93 % similar to *C. wrairi* at the SSU rRNA locus. *Cryptosporidium parvum* has been identified in capybara (*Hydrochoerus hydrochaeris*) in Brazil (Meireles et al. 2007); however, it is likely that the infections resulted from exposure of capybara to water polluted by anthroponotic activities.

Host (scientific name)	Cryptosporidium taxa	Reference
American red squirrel (Tamiasciurus hudsonicus)	C. ubiquitum, chipmunk genotype I	(Ziegler et al. 2007a, b)
Bobak marmot (Marmota bobac)	C. andersoni	(Ryan et al. 2003a)
Belding's ground squirrel (Spermophilus beldingi)	Sbey/Sbld A, Sbey/Sbld/Sltl C, Sbld D	(Pereira et al. 2010)
California ground squirrels (Spermophilus beecheyi)	Sbey/Sbld A, Sbey B, Sbey/Sbld/Sltl C	(Atwill et al. 2001; 2004)
Eastern chipmunk (Tamias striatus)	C. andersoni, C. baileyi, C. ubiquitum, chipmunk genotypes I–II	(Perz and Le Blancq 2001; Feng et al. 2007; Ziegler et al. 2007a, 2007b)
Eastern grey squirrel (Sciurus carolinensis)	<i>C. baileyi, C. muris, C. parvum,</i> <i>C. ubiquitum,</i> chipmunk genotype I, deer mouse genotype III, skunk genotype	(Sundberg et al. 1982; Feng et al. 2007; Ziegler et al. 2007a, 2007b)
Eurasian red squirrel (Sciurus vulgaris)	C. ubiquitum, ferret genotype, chip- munk genotype I (W17)	(Bertolino et al. 2003; Feng et al. 2007; Kváč et al. 2008a; Lv et al. 2009)
Fox squirrel (Sciurus niger)	C. ubiquitum, Cryptosporidium sp.	(Current 1989) (Stenger et al. unpublished)
Golden-mantled ground squirrel (Spermophilus lateralis)	Sbey/Sbld/Sltl C	(Pereira et al. 2010)
Siberian chipmunk	C. muris, C. parvum, chipmunk	(Matsui et al. 2000; Lv
(Tamias sibiricus)	genotype III, Ferret genotype	et al. 2009)
Southern flying squirrel (Glaucomys volans)	Cryptosporidium sp.	(Current 1989)
Thirteen-lined ground squirrel (Ictidomys tridecemlineatus)	Cryptosporidium sp.	(Current 1989)
Woodchuck or Groundhog (Marmota monax)	C. ubiquitum	(Feng et al. 2007; Ziegler et al. 2007b)

Table 5.16 Cryptosporidium spp. identified in the Sciuridae – chipmunks and squirrels

5.13.5 Beaver and Gophers

Cryptosporidium has been detected in the American and European beavers and the plains pocket gopher (Table 5.18). Combining the data from three studies in the eastern US (Fayer et al. 2006; Feng et al. 2007; Ziegler et al. 2007b), the prevalence of *Cryptosporidium* in American beavers (*Castor Canadensis*) was 4.0 % (7/176).

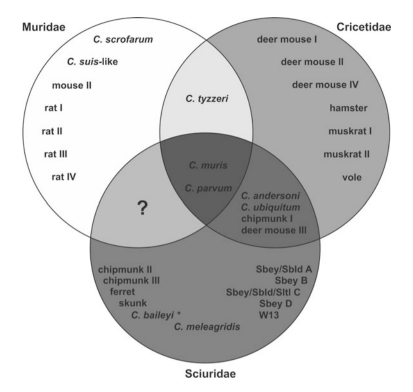


Fig. 5.4 Venn diagram showing group specific and overlapping *Cryptosporidium* taxa from the muridae, cricetidae, and scuridae. **C. baileyi* is generally associated with avian hosts

In Europe, *Cryptosporidium* was detected in 19.2 % (10/52) of European beavers (*Castor fiber*) in northeastern Poland (Paziewska et al. 2007). *Cryptosporidium ubiquitum* and a novel beaver genotype were identified in beavers in New York. The beaver genotype clusters with mink and ferret genotypes in a neighbor-joining phylogeny of SSU rRNA sequences, (Fig. 5.3b) and is most similar to a ferret-like genotype, sharing 99.6 % identity at the SSU rRNA locus.

5.14 Non-Human Primates

There are about 375 species of non-human primates in 15 families and 68 genera. All are omnivorous, and occupy both forest and open grassland habitats. There is potential for *Cryptosporidium* transmission between humans and endangered non-human primates in areas where the two live in close proximity (Nizeyi et al. 1999, 2002; Graczyk et al. 2001). Yet, knowledge of *Cryptosporidium* and cryptosporidiosis in our closest relatives remains relatively poor, and few studies

Host (scientific name)	<i>Cryptosporidium</i> taxa	Reference
Capybara (Hydrochoerus hydrochaeris)	C. parvum	(Meireles et al. 2007)
Chinchilla (Chinchilla laniger)	Cryptosporidium sp.	(Yamini and Raju 1986)
Guinea pig (Cavia porcellus)	C. wrairi, guinea pig genotype	(Vetterling et al. 1971; Xiao et al. 1999b; Huber et al. 2007; Lv et al. 2009)
Indian crested porcupine (Hystrix indica)	Cryptosporidium sp.	(Fayer et al. 2000)
North American porcupine (Erethizon dorsatum)	Cryptosporidium sp.	(Ziegler et al. 2007a, 2007b)
Nutria/coypu (Myocastor coypus)	Cryptosporidium sp.	(Ryan et al. 2003a)

Table 5.17 Cryptosporidium spp. identified in hystricomorphs

Table 5.18 Cryptosporidium spp. identified in the beaver and gophers

Host (scientific name)	Cryptosporidium taxa	Reference
American beaver (Castor canadensis)	C. ubiquitum, beaver genotype	(Fayer et al. 2006; Feng et al. 2007; Ziegler et al. 2007b)
European beaver (<i>Castor fiber</i>)	Cryptosporidium sp.	(Siński et al. 1998; Paziewska et al. 2007)
Plains pocket gopher (Geomys bursarius)	Cryptosporidium sp.	(Current 1989)

have genotyped isolates (Table 5.19). Among those that have, the potential for zoonotic transmission is clearly evident in the species detected.

Cryptosporidium parvum has been identified in the mountain gorilla (*Gorilla gorilla beringei*) from Uganda; toque macaque (*Macaca sinica sinica*), gray lagur (*Semnopithecus entellus thersites*), and purple-faced lagur (*Trachypithecus vetulus philbricki*) from Sri Lanka; and rhesus macaque (*Macaca mulatta*) from China (Graczyk et al. 2001; Ekanayake et al. 2007; Ye et al. 2012). *Cryptosporidium hominis* has been reported in rhesus macaques from the US and China and olive baboons (*Papio anubis*) from Kenya (Feng et al. 2011; Li et al. 2011; Ye et al. 2012). Isolates from the red-tailed guenon, red colobus, and black-and-white colobus were indistinguishable from *C. hominis*, *C. parvum* and *C. cuniculus* at the COWP locus (Salyer et al. 2012). Other *Cryptosporidium* taxa detected in primates include *C. felis* in rhesus macaques at a park in China (Ye et al. 2012) and a *Cryptosporidium* sp. closely related to *C. suis* in a Coquerel's sifaka (*Propithecus coquereli*) in the US (da Silva et al. 2003).

The clinical signs of cryptosporidiosis in infant non-human primates resemble those seen in human infants. Cryptosporidiosis was diagnosed in 81/157 infant primates, predominantly pigtailed macaques (*Macaca nemestrina*), housed in a nursery unit at the Washington Regional Primate Research Center. The mean age at onset of oocyst shedding was 38 ± 25 days, and animals shed oocysts for

 Table 5.19 Cryptosporidium spp. identified in non-human primates

Host (scientific name)	Country	Cryptosporidium taxa	Reference
Captive – zoo			
Black-and-white ruffed lemur (<i>Lemur variegatus</i>)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Black-capped squirrel monkey (Saimiri sciureus boliviensis)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Black lemur (Eulemur macaco)	Spain	Cryptosporidium sp.	(Gómez et al. 2000; Gracenea et al. 2002)
Bonnet macaque (Macaca radiata)	Malaysia	Cryptosporidium sp.	(Lim et al. 2008)
Brown spider monkey (Ateles belzebuth hybridus)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Brown lemur (Lemur macacomayottensis)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Campbell's mona (Cercopithecus campbelli)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Collared mangabey (<i>Cercocebus</i> <i>torquatus lunulatus</i>)	Spain	Cryptosporidium sp.	(Gracenea et al. 2002)
Common marmoset (<i>Calithrix jacchus</i>)	USA	Cryptosporidium sp.	(Kalishman et al. 1996)
Cotton-top tamarin (Saguinus oedipus)	USA	Cryptosporidium sp.	(Heuschele et al. 1986)
Drill (Mandrillus leucophaeus)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Gray langur (Semnopithecus)	Malaysia	Cryptosporidium sp.	(Lim et al. 2008)
Lesser slow loris (Nycticebus pygmaeus)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Mangabey (Cercocebus albigena)	Spain	Cryptosporidium sp.	(Gómez et al. 1992, 2000; Gracenea et al. 2002)
Marimonda spider monkey (Ateles belzebuth)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Olive baboon (Papio anubis)	Italy	Cryptosporidium sp.	(Fagiolini et al. 2010)
Patas monkey (Erythrocebus patas)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Pig-tailed macaque (Macaca leonina)	Malaysia	Cryptosporidium sp.	(Lim et al. 2008)
Red ruffed lemur (Vaecia varie- gate rubra)	USA	Cryptosporidium sp.	(Heuschele et al. 1986)
Ring-tailed lemur (Lemur catta)	Spain Italy	Cryptosporidium sp.	(Gómez et al. 2000) (Fagiolini et al. 2010)
Siamang (Hylobates syndactylus)	Spain	Cryptosporidium sp.	(Gómez et al. 2000; Gracenea et al. 2002)
Southern talapoin monkey (Miopithecus talapoin)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Stump-tailed macaque (Macaca arctoides)	Malaysia	Cryptosporidium sp.	(Lim et al. 2008)

(continued)

Table 5.19	(continued)
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Host (scientific name)	Country	Cryptosporidium taxa	Reference
Talapoin monkey (<i>Cercopithecus talapoin</i>)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Tibetan Macaque (<i>Macaca</i> <i>thibetana</i>)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Velvet monkey (<i>Cercopithecus aethiops</i>)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
Western gorilla (Gorilla gorilla gorilla)	Spain	Cryptosporidium sp.	(Gómez et al. 2000; van Zijll Langhout et al. 2010)
White-collared monkey (Cercocebus torquatus)	Spain	Cryptosporidium sp.	(Gómez et al. 1992)
White-crowned mangabeys (Cercocebus torquatus lunulatus)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
White-faced saki (Pithecia pithecia)	Spain	Cryptosporidium sp.	(Gómez et al. 2000)
Captive – non-zoo		<i>a</i>	
Crab-eating macaque (Macaca fascicularis)	USA	C. muris-like	(Dubey et al. 2002)
Coquerel's sifaka (Propithecus coquereli)	USA	Cryptosporidium sp.	(Charles-Smith et al. 2010) (da Silva et al. 2003)
Rhesus macaque (Macaca	USA	Cryptosporidium sp.	(Yanai et al. 2000)
mulatta)	USA	Cryptosporidium sp.	(Osborn et al. 1984; Feng et al. 2011; Ye et al. 2012)
	Not reported	C. hominis monkey genotype	Feng et al. 2011;
Southern pig-tailed macaque (Macaca nemestrina)	USA	Cryptosporidium sp.	(Miller et al. 1990a)
Free living Black-and-white colobus (Colobus guereza)	Uganda	Cryptosporidium sp.	(Salzer et al. 2007; Salyer et al. 2012)
Gray lagur (Semnopithecus entellus thersites)	Sri Lanka	C. parvum, Cryptospo- ridium sp.	(Ekanayake et al. 2006)
Green monkey (<i>Cercopithecus aethiops</i>)	Ethiopia	<i>Cryptosporidium</i> sp.	(Legesse and Erko 2004)
Mountain gorilla (Gorilla beringei beringei)	Uganda	Cryptosporidium sp.	(Nizeyi et al. 1999; Sleeman et al. 2000; Graczyk et al. 2001)
	Rwanda	C. parvum	Graczyk et al. 2001)
		Cryptosporidium sp.	Sleeman et al. 2000,
			(continued)

(continued)

Host (scientific name)	Country	Cryptosporidium taxa	Reference
Olive baboon (Papio anubis)	Kenya	Cryptosporidium sp.	(Muriuki et al. 1997, 1998; Hope et al. 2004; Legesse and Erko 2004; Li et al. 2011)
		C. hominis	Li et al. 2011)
	Uganda	Cryptosporidium sp.	Hope et al. 2004
	Ethiopia	Cryptosporidium sp.	Legesse and Erko 2004
Purple-faced lagur (Trachypithecus vetulus)	Sri Lanka	Cryptosporidium sp.	(Ekanayake et al. 2006)
		C. parvum	(Ekanayake et al. 2007)
Rhesus macaque (Macaca mulatta)	China	C. felis, C. hominis, C. parvum	Ye et al. 2012
Red colobus (Procolobus tephrosceles)	Uganda	Cryptosporidium sp.	(Salzer et al. 2007, Salyer et al. 2012)
Red-tailed guenon (<i>Cercopithecus ascanius</i>)	Uganda	Cryptosporidium sp.	(Salzer et al. 2007) (Salyer et al. 2012)
Savanna chimpamzee (Pan troglodytes schweinfurthii)	Tanzania	Cryptosporidium sp.	(Gonzalez-Moreno et al. 2013) (Ekanayake et al. 2006)
Toque macaque (Macaca sinica)	Sri Lanka	Cryptosporidium sp.	Ekanayake et al. 2007)
Vervet monkey (Chlorocebus pygerythrus)	Kenya	Cryptosporidium sp.	(Muriuki et al. 1997, 1998)
Western gorilla (Gorilla gorilla gorilla)	Gabon	Cryptosporidium sp.	(van Zijll Langhout et al. 2010)

 Table 5.19 (continued)

7–78 days. All but one of the animals had clinical symptoms of enteric infection, which included severe diarrhea and dehydration. The outbreak was confined to the nursery and no cases were detected among juvenile or adult animals (Miller et al. 1990a). A follow-up study (Miller et al. 1990b) characterized the course of experimental infections in pigtailed macaques infected with 10 or 2×10^5 oocysts from naturally infected macaques (the species/genotype of the isolate was not reported). Infected animals suffered clinical enteritis, watery stools, lethargy, and loss of appetite. Animals infected with 10 and 2×10^5 oocysts began shedding oocysts on day 8 and day 7, respectively, and there was no difference in the duration of intense oocyst shedding or clinical symptoms between the treatments.

Cryptosporidiosis in immunocompromised non-human primates resembles cryptosporidiosis in immunocompromised humans with respect to extraintestinal involvement. Kaup et al. (1994) diagnosed biliary and pancreatic cryptosporidiosis in several rhesus monkeys following an experimental simian immunodeficiency virus (SIV) infection. Yanai et al. (2000) reported moderate to severe bronchopneumonia with cryptosporidiosis in macaques experimentally infected with SIV. Endogenous stages were detected in the trachea, lungs, bile ducts, pancreas, and intestine. Conjunctival infections were detected in six SIV-infected, immunode-ficient rhesus monkeys (Baskin 1996). Although conjunctival cryptosporidiosis has been reported in birds, there have been no reported cases in humans. Singh et al. (2011) examined the time during an SIV infection when a self-limiting *Cryptosporidium parvum* infection becomes persistent due to mucosal immune defects. One group of SIV-infected macaques was challenged with *C. parvum* during acute SIV infection and the second group was challenged during the chronic infection phase. Interestingly, persistent cryptosporidiosis developed during acute SIV infection.

5.15 Diversity of *Cryptosporidium* in Different Vertebrate Classes

The diversity of mammalian cryptosporidia appears to reflect mammal diversity. We compared *Cryptosporidium* diversity in the order Rodentia – the most diverse order of mammals - to diversity in other mammalian orders. Figure 5.5 presents 68 mammalian Cryptosporidium taxa (species and genotypes) in a Venn diagram with two sets: 'Rodentia' and 'other mammals' Among the taxa that are members of both sets (the intersection), C. andersoni, C. scrofarum, and the ferret genotype primarily infect non-rodent hosts and are therefore included with 'other mammals'. Similarly, C. tyzzeri and C. muris are primarily parasites of rodents and are included with the Rodentia set. Cryptosporidium baileyi was excluded from the analysis because it primarily infects avian hosts. Forty percent (27/67) of mammalian Cryptosporidium taxa are associated with the order Rodentia, which reflects the diversity of Rodentia within mammals (Rodentia contains 40 % of mammalian species). Three percent (2/67) of Cryptosporidium taxa (C. parvum and C. ubiquitum) can be considered generalists, and 57 % (38/67) of taxa are associated with non-rodent mammals. These data suggest that cryptosporidia diverged in close association with mammal species.

In contrast to mammals, it appears that the diversity of avian cryptosporidia does not reflect avian diversity. The order Passeriformes comprises about 50 % of avian species; species diversity in the Passeriforme order is comparable to that of the entire mammal class. Figure 5.6 presents 17 *Cryptosporidium* taxa from avians in a Venn diagram with two sets: 'Passeriformes' and 'other avians'. Muskrat genotype I, *C. hominis, C. hominis*-like, *C. muris*, and *C. parvum* were excluded from the analysis because they are not considered avian-associated taxa. Seventeen percent (2/12) of avian *Cryptosporidium* taxa are exclusively found in passerines (avian genotypes I and IV), 50 %

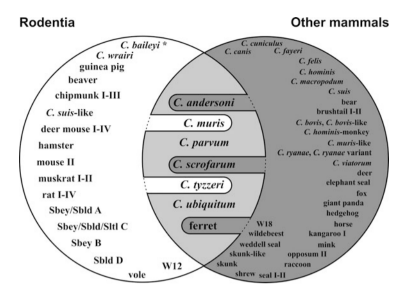


Fig. 5.5 Venn diagram showing group-specific and overlapping *Cryptosporidium* taxa in rodent and non-rodent mammal groups. **C. baileyi* is generally associated with avian hosts

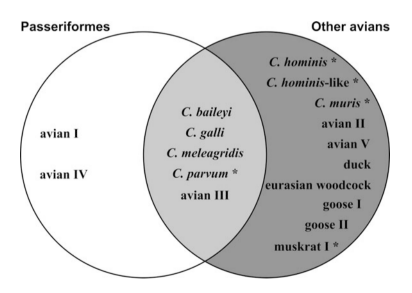


Fig. 5.6 Venn diagram showing group-specific and overlapping *Cryptosporidium* taxa from passerine and non-passerine bird groups

(6/12) are found only in non-passerine orders, and 33 % (4/12; *C. baileyi*, *C. meleagridis*, *C. galli*, and avian genotype III) infect multiple avian orders and can be considered generalists. From these data, it can be hypothesized that the mechanisms of *Cryptosporidium* diversification in avians and mammals are fundamentally different.

The diversity of *Cryptosporidium* in reptiles and amphibians appears to be low. Despite a comparable faunal diversity with birds, and considerably greater faunal diversity than mammals, relatively few *Cryptosporidium* taxa have been reported in reptiles or amphibians.

5.16 Gaps in Knowledge and Suggestions for Future Work

- *Cryptosporidium* remains poorly characterized in many vertebrate orders. For example, Chiroptera (bats), an order that comprises 20 % of mammal species, has not been well studied. If *Cryptosporidium* diversity reflects bat diversity, then we should expect to find many bat-adapted *Cryptosporidium* taxa.
- *Cryptosporidium viatorum* is a recently described species that has thus far been identified only in cases of human cryptosporidiosis in the UK (n = 10) and Sweden (n = 2); all cases were associated with travel to countries in the Indian subcontinent, Africa, and South America (Elwin et al. 2012b; Insulander et al. 2013). Studies are needed to determine if there is a major animal host for this species. Greater sampling in general, particularly of animals in their natural habitat, will help to help to identify emerging sources of human disease.
- Relatively little is known about the general biology of many of the *Cryptosporidium* genotypes identified. Experimental evidence of infection characteristics and host specificity will enhance understanding of the biological diversity and will clarify taxonomy.

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Part II Molecular Biology

Chapter 6 Cryptosporidium: Current State of Genomics and Systems Biological Research

Aaron R. Jex and Robin B. Gasser

Abstract Recent years have seen an unprecedented expansion in our knowledge of Cryptosporidium and cryptosporidiosis through the emergence of the genomics and systems biological age. High-quality draft genome sequences are now published for C. parvum and C. hominis, and the draft assembly of C. muris has been made publicly available. These genome sequences reveal a highly stream-lined parasite with limited metabolic and biosynthetic pathways and a heavy reliance on the hostcell. Bottlenecks in these pathways may be exploited for new drugs, which remain stubbornly illusive for these parasites. As more genomic information becomes available, fundamental research into gene regulation, genomic evolution and genome-wide variation becomes possible. This research will provide new insights into the transmission dynamics of these parasites and markers associated with host-specificity, virulence and pathogenicity, and will allow the identification of novel loci for use as molecular-diagnostic markers and genes under heavy immunoselection, potentially providing a basis for vaccine development. With the accelerating reduction in costs associated with 'omic' research, improved accessibility to analytical tools and in vitro culture of *Cryptospordium*, this field has tremendous potential to shape our understanding of their biology in the coming years.

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6.1 Genomics of Cryptosporidium

Research of Cryptosporidium in the genomics age is landmarked by the publication of the complete genome sequence of C. parvum Iowa strain (Abrahamsen et al. 2004). This genome was sequenced to ~13-fold coverage using a shotgun Sanger sequencing approach (Table 6.1). Briefly, this process involved isolating total genomic DNA from purified C. parvum oocysts, randomly shearing the DNA (2-5 kb) and constructing plasmid libraries, from which 120,000 clones were sequenced. In addition, to facilitate scaffolding and the resolution of repeat regions, the authors sequenced the genome to ~ 0.5 -fold depth using data generated from large-insert (~15 kb) λ -phage libraries constructed from genomic DNA of the same Iowa isolate. The assembly of the sequence data from both libraries was guided by a physical map, which had been generated previously for this C. parvum isolate using ~200 oligonucleotide primers to screen haploid amounts (i.e., a "HAPPY" map) of C. parvum DNA by the polymerase chain reaction (PCR) for marker regions spaced ~50 kp apart (Piper et al. 1998; Abrahamsen et al. 2004) and a draft sequence for chromosome 6 (Bankier et al. 2003). The combination of the long sequence reads, large-insert libraries and the HAPPY map, combined with selective PCR-based sequencing for the purpose of gap closure, allowed the assembly of the C. parvum genome into its eight chromosomes, leaving five physical gaps. The final assembly revealed an AT-rich (70 %) and highly compact genome of ~9.1 Mb, which is relatively small in comparison with related apicomplexans, such as Eimeria tenella (~60 Mb) (Shirley 1994, 2000) and *Plasmodium falciparum* (~23 Mb) (Gardner et al. 2002a) and, unlike many apicomplexans, includes no mitochondrial or apicoplast genomes.

The annotation of the C. parvum genome was achieved primarily through the prediction of open reading frames (ORFs) of at least 67 amino acids in length from the assembled genome sequence. Although gene prediction methods based on the use of hidden Markov and machine learning models for the analysis of expressed sequence tag (EST) data, and, more recently, deep RNA-Sequencing data (Wang et al. 2009) are considered standard approaches for current genomic sequencing projects (Jex et al. 2011a; Young et al. 2012), extensive EST data are not yet available for *Cryptosporidium* species and are challenging to generate from the endogenous developmental stages. Noting this, the C. parvum genome appears to have few intronic sequences and, thus, the straight-forward prediction of ORFs was sufficient to allow the identification of 3,952 (3,807 protein-coding) genes in C. parvum. These genes were fewer in number than predicted to be encoded by the genomes of related apicomplexans, such species of *Plasmodium* (~5,300 genes) (Gardner et al. 2002b), Toxoplasma (~7,700) (Radke et al. 2005) and Theileria (~4,000) (Gardner et al. 2005), which, coupled to a lack of introns and limited non-coding sequence (~25 % of the entire genome), explains the smaller size of the C. parvum genome (see also Bankier et al. 2003).

Feature	<i>C. parvum</i> Iowa II	C. hominis TU502	C. parvum TU114	C. muris RN66
Project accession Assembly size (Mb)	AAEE01000000 9.1	AAEL000000 8.7	not available not available	PRJNA19553 9.2
Assembly metrics (Contigs: N50)	9.1 8: 1.10 Mb	8.7 1,422: 14.5 kb	not available	9.2 45: 0.72 Mb
Number of genes	3,952	3,994	~3,952	3,986
GC richness	30.3	31.7	30.3	27.8
% identity (to C. parvum Iowa II)	-	~97	~99.8	not available
Human infective	Yes	Yes	Yes	No
Human specific	No	Yes	Yes	No

 Table 6.1
 Comparison of characteristics and metrics of each Cryptosporidium genome sequenced and available in CryptoDB (www.cryptodb.org)

The completion of the *C. parvum* genome was rapidly followed by the publication of the genome of *C. hominis* TU502 strain (Xu et al. 2004). As for *C. parvum*, the *C. hominis* genome was sequenced using a shotgun Sanger method to a depth of ~12-fold from a plasmid insert library. Unlike for *C. parvum*, a physical map was not available for *C. hominis*, although the authors constructed large (~7–8-fold coverage), bacterial artificial chromosome (BAC) libraries for this species and used the *C. parvum* HAPPY map to guide assembly. Using this approach, the *C. hominis* genome is currently represented by ~1,400 contigs and ~300 scaffolds, with <250 gaps. The gene content for the *C. hominis* genome appears to be similar to that of *C. parvum*, with the genome predicted to encode 3,994 genes. Xu et al. (2004) estimated that 5–20 % of the *C. hominis* genes represent introns, with the higher end of this range differing markedly from that predicted for *C. parvum*, emphasizing the need for increased transcriptomic research of these *Cryptosporidium* species to validate their genome annotation.

An alignment of the *C. parvum* and *C. hominis* genomes showed the extent of similarity between the two species (Xu et al. 2004). Upon pairwise comparison, Xu et al. (2004) found that the genomes diverged in sequence by no more than 3-5 %, with no evidence of large insertion, deletion or recombination events. Their respective gene sets appear to be almost identical, with genes of *C. parvum* found to have homologues in *C. hominis*, and the <50 genes predicted for *C. hominis* and apparently absent from *C. parvum* relating to known gaps in the genome of *C. hominis*, suggesting that both parasites have the same or similar biological pathways, with the differences in their phenotypic behaviour (e.g., host specificity, infectivity and symptomology; see Chaps. 4 and 9) possibly relating to transcriptomic differences and/or mutations (Xu et al. 2004).

6.2 The Inferred Cryptosporidium Proteome

The compact genomes of C. parvum and C. hominis seem to reflect the biology of these parasites and have a reduced complement of genes associated with both anabolic and catabolic phases of metabolism. Energy generation (i.e., ATP) appears to be dependent exclusively upon the degradation of simple sugars via anaerobic glycolysis, with no evidence of a mitochondrial genome or many of the nuclear genes associated with the Krebs cycle or electron transport chain (Abrahamsen et al. 2004; Xu et al. 2004). In addition, there is no evidence for the presence of genes associated with energy production via the digestion of fatty acids/proteins or the urea and nitrogen cycles (i.e., for amino acid synthesis) or the shikimate pathway (Xu et al. 2004). The 'minimalistic' metabolome of these organisms substantially reduces their ability to synthesize essential building blocks (e.g., some nucleotides and amino acids), likely leading to a major dependency on the host cell. Supporting this hypothesis is the identification of a variety of amino acid transporter genes, which are proposed to be involved in amino acid salvaging, and a variety of enzymes necessary for the conversion of amino acids and nucleotides (e.g., pyrimidines to purines and purines to pyrimidines) (Abrahamsen et al. 2004; Striepen and Kissinger 2004; Xu et al. 2004).

In addition to the core metabolome predicted from the genomes of *C. parvum* and *C. hominis*, these parasites encode a variety of proteins involved in a wide array of biological processes. Not surprisingly, many of the fundamental biological processes of the cell (e.g., DNA replication, transcription and translation) appear to be linked to standard eukaryotic pathways. A notable exception to this, as highlighted by Abrahamsen et al. (2004) for *C. parvum*, but also consistent with *C. hominis* (see Xu et al. 2004), is a diminished spliceosomal pathway with a reduced U6 spliceosomal complex, and a lack of identifiable homologues to DICER and argonaute, suggesting an altered role for small interferring RNAs in gene regulation in these microorganisms.

Of major interest among the structural proteins encoded by the *Cryptosporidium* genomes are those involved in the structure and function of the apical complex and/or adorning the cell-surface. It appears that most of the known proteins associated with the apical complex in species of *Plasmodium* have homologues in *Cryptosporidium*, suggesting that the overall molecular biology of this crucial organelle is conserved across many, if not all apicomplexans. Among the most studied groups of cell-surface proteins encoded by an apicomplexan genome are the variable surface proteins, which provide *Plasmodium* with its infamous capacity to evade host immune responses (Reeder and Brown 1996). Abrahamsen et al. (2004) explored the *C. parvum* genome in-depth for these sequences but did not find them. These authors did note several groups of molecules with strong SER- and THR-rich glycosylation signals and transmembrane domains, consistent with a diverse panel of cell-surface glycoproteins, which appear to have undergone an expansion via duplication since the evolutionary emergence of *Cryptosporidium*. Many of these molecules were actively transcribed during in vitro development (Abrahamsen

et al. 2004), and it would seem likely that at least some of them play important roles in cell-cell adhesion and other mechanisms associated with host invasion and infection, as has been shown to be the case for the 15 and 40 kilodalton (kDa) glycoprotein (G15 and GP40), both of which are encoded by the 60 kDa glycoprotein (*gp60*) gene (Strong and Nelson 2000).

6.3 The Druggable Genome of Cryptosporidium

Despite substantial research into the molecular biology of *Cryptosporidium* in the past decades, there are still no highly effective drugs or vaccines for intervention, and few examples of compounds that show or appear to show efficacy against members of the genus (Jex et al. 2010). A major goal of genomic research of these parasites is to identify novel drug targets to address the serious lack of chemotherapeutic treatment options. In sequencing the C. parvum genome, Abrahamsen et al. (2004) highlighted that the 'minimalist' metabolic/biosynthetic pathways of this microorganism present numerous, novel avenues for drug target and drug discovery. Most notable among these pathways, given the limited biosynthetic capabilites of Cryptosporidium species, is the observation that the parasites have a single enzyme, 5' inosine monophosphate dehydrogenase (IMPDH), for the conversion of adenosine monophosphate to guanosine monophosphate. Mycophenolic acid is a known inhibitor of IMPDH (Senda et al. 1995), but attempts to inhibit Cryptosporidium-encoded IMPDH using this compound class have not been successful. The bacterial origin of the enzyme in these parasites was highlighted as a possible confounding issue (Umejiego et al. 2004). However, more recently, using in silico docking experiments and high-throughput screening, several compounds (Compounds "A", "F", "G" and "H": Umejiego et al. 2008) with specific efficacy against Cryptosporidium in cell-culture superior to the performance of paramomycin were identified. Abrahamsen et al. (2004) also predicted the potential of a wide variety of transporter proteins as targets, such as those used by the parasite to salvage nucleotides and essential amino acids from the host, as well as a mitochondrial-associated alternative oxidase, which catalyzes the reduction of oxygen to water without the production of free radicals and/or oxidizing reagents (e.g., superoxide/H₂O₂). These authors (Abrahamsen et al. 2004) highlighted the potential of salicylhydroxamic acid to inhibit this latter enzyme.

In a recent review (Jex et al. 2011b), we conducted an in-depth exploration of the *Cryptosporidium* genome for potential drug targets. We found, for example, that 17 of 31 high priority 'druggable' molecules proposed for *P. falciparum* (see Crowther et al. 2010) had close homologues in *Cryptosporidium* species. Given the recent focus on repurposing anti-malarial drugs via the Medicines for Malaria Venture (www.mmv.org), there are clear prospects for screening these "malaria-box" compounds for efficacy against *Cryptosporidium*. In addition to assessing these homologues, we conducted a global prediction of druggability on the complete gene set of *C. parvum and C. hominis* (see Jex et al. 2011b) by assessing their

homology to essential genes in Saccharomyces cerevisiae (yeast). Approximately 33 % of the annotated Cryptosporidium genes have an homologue in yeast, and >500 of them are associated with a lethal phenotype; 313 of these 'essential' genes have homologous sequences in the BRENDA/CHEMBL databases (Schomburg et al. 2002; Gaulton et al. 2012). Most notable among these are 60 proteinsynthesizing GTPases, for which >50 inhibitors (primarily stemming from anticancer research) are contained within these database, including a variety of aminoglycosides, such as dihydrostreptomycin, hygromycin, neomycin and viomycin (Jex et al. 2011b). Interestingly, one of the few available compounds with some efficacy against *Cryptosporidium* is an aminoglycoside (i.e., paromomycin) (Rossignol 2010), although it is not known to have an inhibitory effect on GTPases. Aminoglycides (often named with the suffix *-mycin*) have been widely used in humans as topical antibiotics, but are known to be toxic to mammalian cells (Martinez-Salgado et al. 2007; Rizzi and Hirose 2007; Guthrie 2008). Recent research shows that this toxicity can be reduced through careful administration (Murakami et al. 2008; Pannu and Nadim 2008) and/or modified formulations (Tugcu et al. 2006; Jeyanthi and Subramanian 2009; Nagai and Takano 2010), suggesting that the aminoglycosides might be re-considered as anti-cryptosporidial compounds.

6.4 Additional Cryptosporidium Genomes

An interesting aspect of the biology of Cryptosporidium, and of critical importance in the control of cryptosporidiosis, is the variable host-specificity of the species of this genus. For example, epidemiological studies of these parasites in the past decades, coupled to molecular systematics in more recent times, have provide a wealth of data to indicate that human infections are largely caused by C. parvum and C. hominis, with most other species of Cryptosporidium rarely being detected in otherwise healthy humans and displaying varying levels of adaptation to other hosts (Xiao et al. 2004). Of the Cryptosporidium species known to infect humans, C. hominis is thought to be human-specific, whereas C. parvum is transmitted by either anthroponotic or zoonotic routes (Xiao et al. 2004). However, C. hominis has been reported in naturally occurring infections in animals, such as cattle in Scotland (Smith et al. 2005) and, has been detected (although infections were not confirmed) in the faeces of cattle in New Zealand (Abeywardena et al. 2012) and a dugong in Australia (Morgan et al. 2000). In addition, although cattle, in particular, have been identified as a major source of C. parvum for transmission to humans, population genetic and molecular epidemiological data suggest that human-specific lineages might yet exist within this well-studied species (Mallon et al. 2003; Jex and Gasser 2010).

Several aspects might explain the variable host-specificity or adaption of these species: (a) the differing gastrointestinal conditions (e.g., pH, and body temperature) between and among vertebrate groups known to be infected by at least one species

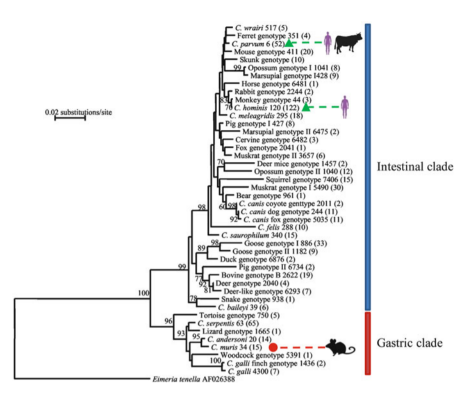


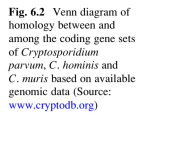
Fig. 6.1 Current molecular phylogeny (based on 18S ribosomal DNA sequence) of *Cryptosporidium*, highlighting the species for which complete genomic data are presently available (phylogram modified from Xiao et al. 2004). Genomes that are both sequenced and published are represented by *green triangles*. Genomes that have been sequenced but not yet published in the peer-reviewed literature are represented by a *red circle*. Human/animal images represent that common hosts for each species for which complete genomic data are presently available

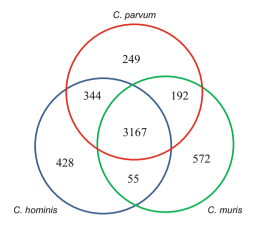
of *Cryptosporidium*; (b) differences between and among the *Cryptosporidium* species or their hosts in the cell-surface molecules involved in host-cell binding and/or invasion by the activated sporozoite; and (c) the differing innate or acquired immunological responses of the host against *Cryptosporidium*. Understanding whether these factors influence the host range of different species will be of critical importance in the effective control of cryptosporidiosis.

In pursuit of this understanding, the recent sequencing of the genomes of *C. parvum* TU114 (Widmer et al. 2012) and *C. muris* RN66 (NCBI bioproject: PRJNA19553) genomes represents a major advance, with the former appearing to be a humanadapted/specific lineage of *C. parvum* (classified subgenotype IIcA5G3R2 at gp60 (Jex and Gasser 2010)) and the latter being representative of a gastric-colonizing species not known to infect humans (Fig. 6.1: Xiao et al. 2004).

To faciltate comparisons among C. parvum TU114 (IIcA5G3R2), C. parvum IOWA (IIaA15G2R1) and C. hominis TU502 (IaA21), two C. parvum TU114 samples were selected and their genomes sequenced using Illumina technology to a mean depth of \sim 20–40-fold, revealing, upon alignment to the *C*. parvum IOWA sequence, 12,748 single nucleotide polymorphisms (SNPs) (i.e. ~1.4 per kilobase) common to both TU114 isolates. The authors (Widmer et al. 2012) hypothesized that the human-infective phenotype of both C. parvum TU114 and C. hominis was a likely consequence of the same positive selective force, and explored their data for genomic regions in the TU114 samples that had diverged from C. parvum IOWA and converged with C. hominis. These analyses identified several genomic regions of C. parvum TU114 which were significantly more similar to C. hominis than to C. paryum IOWA, despite the two C. paryum genome sequences being tenfold more similar to each other than either was to C. hominis overall. Three exonic regions in the genome, corresponding to the genes cgd1 650, cgd3 3370 and cgd6 5260. contained regions that were significantly more similar to orthologues in C. hominis than those in the IOWA strain of C. parvum (Widmer et al. 2012). This pattern was shown to be consistent for five other C. parvum IIc isolates in conventional PCR-based sequencing study of these gene regions (Widmer et al. 2012). Although their specific function is not known, gene regions cgd1 650, cgd3 3370 are predicted to encode signal peptide domains but no transmembrane domains, suggesting that their products are secreted by the parasite. At present, no other information on the function of these genes is available in the literature. Clearly, this aspect warrants detailed exploration. gene cgd6_5260 appears to encode an ABC transporter (ATpase with 2 AAA domains). AAA + ATPases have a wide range of cellular functions, including membrane fusion, proteolysis and DNA replication (IPR003593). In the current context, membrane fusion is noteworthy, as it might suggest cell-cell interactions between the parasite and the host; however, cg6 5260 also appears to encode a nucleotide-binding domain, suggesting that its role more likely relates to DNA replication, which is obviously more difficult to reconcile with host-specificity.

The sequencing of the *C. muris* genome will also provide new insights into aspects of *Cryptosporidium* biology and host-parasite relationships. *C. muris* is infective primarily to rodents and is located in the stomach of the host, whereas the other sequenced *Cryptosporidium* species are pathogens of the small intestine. Currently, although the *C. muris* genome has not been published in the peerreviewed literature, the complete, annotated genome assembly is available through the GenBank database (accessible via www.ncbi.nlm.nih.gov) and CryptoDB (www.cryptodb.org). The current assembly is ~9.2 Mb and consists of 84 scaffolded sequences, with 50 % of the genome (i.e., N50) represented by contigs of at least 720 kb. The *C. muris* genome is predicted to encode ~3,900 protein-coding genes, with 3,511 and 3,359 of them predicted to have identifiable orthologues in *C. parvum* IOWA and *C. hominis*, respectively, and 3,167 genes being shared among all three species (Fig. 6.2: www.CryptoDB.org). Characterizing the functional similarities and differences associated with these overlapping gene-sets as well as further comparative analyses of the *C. muris*, *C. parvum* and *C. hominis*.





genomes both at the structural level (e.g., overall gene composition and organisation, synteny and the identification of large insertion/deletion or translocation events) and the characterization of genetic variation (i.e., SNPs) across the genome should be considered a high priority and will likely yield significant insights into the genetics and evolution of members of the genus. For example, an alignment of the genomes of C. muris, C. parvum IOWA, C. parvum TU114 and C. hominis provides scope to assess regions of the genome under neutral versus purifying selective pressure at a much deeper level than has been possible previously for this genus. On a genome-by-genome comparison, this information might be useful for assessing phenotypic differences, such as host-specificity and siteselection, among the species. Considering all four genomes, these analyses should provide insight into the genomic 'dark matter' of each species. For example, at present ~75 % of the Cryptosporidium genome is annotated as protein-coding. However, as no RNA-Sequencing data are available for *Cryptosporidium* species, there is no direct support for many of these predicted genes. Given that nearly two thirds of the currently annotated Cryptosporidium genes represent 'hypothetical proteins', for which no functional data are available, more support for the gene-sets is needed. In addition, very little is known about functional, non-coding regions in Cryptosporidium genomes sequenced to date. Studies of the accumulation of mutations in a variety of eukayrotic species (e.g., Homo sapiens, Caenorhabditis elegans and Saccharomyces cerevisiae) have shown that at least 4-8 % of the non-exonic regions of the genomes of these organisms are under purifying selection (Siepel et al. 2005; Gerstein et al. 2010; Ponting and Hardison 2011). In these eukaryotes, many of the non-coding, but evolutionarily constrained regions have important regulatory functions, including non-coding RNAs, transcriptional signal sites, and promotor and/or enhancer regions. Defining regions in Cryptosporidium genomes undergoing purifying selection will assist in identifying such regulatory elements are present in these parasites. Indeed, in sequencing the C. parvum TU114 genome, some authors (Widmer et al. 2012) found evidence to suggest that at least some non-coding regions of the genome appears to be under purifying selection. Including the genome of the distantly related C. muris in these analyses might improve comparisons and provide greater confidence in identifying regions likely to be enriched for regulatory elements (Siepel et al. 2005). In a recent study, Mullapudi et al. (2007) used pattern recognition software to identify conserved motifs in the regions up-stream of the annotated *C. parvum* genes. They proposed these motifs as putative *cis*-regulatory elements (CREs) and found evidence, based on real-time PCR, for the co-transcription of numerous genes sharing the same 'CRE' conserved motif. This research represents one of the first explorations of gene regulation in *Cryptosporidium*. A subsequent study using a similar approach has also predicted conserved CREs associated with the co-transcription of heat shock proteins (HSPs) during *C. parvum* excystation (Cohn et al. 2010). Given the compact nature of *Cryptosporidium* genomes, their 'regulome' may represent a close-approximation of the minimum regulatory network required for eukaryotic cells to function, and might have far reaching biological implications for understanding the eukaryotic cell.

6.5 Systems Biology Research of Cryptosporidium

Referred to initially as 'next-generation sequencing' and, more recently, as 'highthroughput sequencing' or 'massively parallel sequencing', advanced technologies, such as the 454 Roche (Margulies et al. 2005) and Illumina (Bentley et al. 2008), have vastly reduced the cost and time associated with conducting genomic research. New developments in large-scale and benchtop sequencing are expected to further accelerate the reduction in costs of such research and drive innovation across all disciplines of biology and biomedicine (Niedringhaus et al. 2011).

6.5.1 Deep Exploration of the Systems Biology of C. parvum and C. hominis

The innovation associated with the expansion of genomic, transcriptomic and bioinformatic technologies will have profound implications for *Cryptosporidium* research. Initial applications of these technologies will likely include refinements of existing genome sequences for *C. parvum* and *C. hominis*. For example, although the *C. parvum* genome has been assembled, the *C. hominis* genome is presently represented by several hundred scaffolds of >1,400 individual sequence contigs. Finishing these two genomes will allow the gene sets to be confirmed and genetic and genomic differences between these two important, human-infective parasites to be re-assessed, and will also provide a basis to further explore regulatory pathways associated with non-coding RNAs and other regions of intergenic DNA (Siepel et al. 2005; Gerstein et al. 2010; Ponting and Hardison 2011). Furthermore, although the alignment of genome sequences currently provides little indication of significant

structural differences between *C. parvum* and *C. hominis*, the absence of a physcial map for the latter species prevents a thorough comparison. Rearrangements may have relevance, not only in comprehensively determining the differences between the species and their coding gene sets, but also in establishing rates of inter- and intraspecific recombination, which might have implications for the emergence and spread of drug resistance, following the future introduction and administration of effective and safe anti-cryptosporidial compounds on a large scale. Although the in vitro cultivation of *Cryptosporidium* species is presently challenging, recent advances in culture methodologies (reviewed by Jex et al. 2010) might pave the way for the the construction of genetic crossing experiments, as a means of assessing genotypic causes of phenotypic traits, such as virulence, host-specificity and infectivity.

In addition to characterizing and finishing the genome sequence of these species, there is a clear need to support gene annotations. Specifically, most of the proteincoding genes currently predicted for Cryptosporidium species are not supported by transcriptomic or expression data. Previously, this lack of information has related to the challenges associated with isolating enough parasite material for molecular study. However, the advent of high-throughput sequencing technologies provides opportunities to enhance our understanding of the genomics and genetics of Cryptosporidium. Traditional obstacles to the transcriptomic research of species of Cryptosporidium seem to be related primarily to limitations associated with the specific isolation of RNA from the endogenous stages of the life cycle and the relative transcriptional 'dormancy' of the oocyst stage. In vitro culturing of Cryptosporidium has, until recent times, been a major obstacle, and, despite advances in these technologies (Hijjawi et al. 2010), the transcriptome might be quite different from that be the host. Laser capture and microdissection technologies, in the hands of skilled microscopists, might provide a means to overcome the difficulty in exploring the transcriptome of specific endogenous stages when coupled with single-cell RNA-Sequencing technologies (Tang et al. 2010). Such information could pave the way for testing of current gene sets for Cryptosporidium, the assessment of intron-exon boundaries, definition of promoter and transcription start sites, and the characterization of stage-specific changes in gene transcription, identification of alternative splicing of transcripts associated with multi-exon genes or multi-gene operons (Wang et al. 2009). Clearly, such new developments will have major implications for understanding Cryptosporidium molecular biology by allowing the exploraton of key genes and their roles in major life-cycle phases (e.g., infection, reproduction or cellular destruction). In addition, characterizing constitutively transcribed and/or stage-specific genes should also assist significantly toward prioritizing novel drug candidates.

Further expansions of transcriptomic research of *Cryptosporidium* may seek to explore regulatory (i.e., non-coding) RNAs. Although there is little evidence that the genomes of *Cryptosporidium* species have genes encoding argonaute and DICER (Abrahamsen et al. 2004; Xu et al. 2004), commonly involved in RNA silencing pathways, other non-coding RNAs, such as long non-coding, small nuclear and/or *piwi*-RNAs, might play key regulatory roles that remain unexplored in these species to date.

In addition to characterizing genes and differential (including developmentally regulated) transcription, increased insights into the proteome of *Cryptosporidium* is highly desirable, particularly considering that approximately two-thirds of all known genes of *Cryptosporidium* are considered to encode 'hypothetical proteins' (Abrahamsen et al. 2004; Xu et al. 2004). To date, requirements for sufficient amounts of parasite material and the costs associated with proteomic analyses have limited, detailed expression profiling of *Cryptosporidium* species. However, although in vitro cultivation might assist, there have been massive technological enhancements in proteomic technologies, such that low cost and highly sensitive analytical analyses are now feasible (Cox and Mann 2011).

Culturing methods also have major implications for studying other areas of systems biology, particularly in reconstructing the *Cryptosporidium* metabolome. Genomic sequence data indicate that metabolomes of different species of Cryptosporidium are 'minimalistic', with a heavy reliance on simple sugars, anaerobic glycolysis and the harvesting of essential building blocks from the host through a range of small molecule salvage pathways. However, as indicated, ~66 % of the protein-coding genes currently annotated for C. parvum and C. hominis encode hypothetical proteins. Given the large evolutionary distances between Cryptosporidium and eukaryotic model organisms for which functional genomic data are available, it is probable that some of these genes may relate to known metabolic pathways, but are too divergent in sequence to be identified using currently available bioinformatic tools. Considering the extent with which the genome of these parasites appears to have been shaped through the horizontal acquistion of genes from bacteria (Abrahamsen et al. 2004; Xu et al. 2004), one cannot discount the possiblity that these Cryptosporidium species have novel, highly divergent (i.e., relative to other eukaryotes) biosynthetic pathways to account for their metabolic demands that cannot presently be identified among the 'orphan' proteins encoded in the genome. By coupling advanced in vitro cultivation methods for Cryptosporidium to new advances in metabolomic mapping (i.e., highly sensitive gas chromatography - mass spectrophotometry, nuclear magnetic resonance and matrix-assisted laser desorption/ionisation-time of flight (MALDI-TOF) mass spectrophotometric technologies), it will soon be possible to directly explore metabolism in these parasites, as has been done in recent years for species of Trypanosoma and Leishmania (Creek et al. 2012). Such applications have significant potential to improve our knowledge of Cryptosporidium species and assist in finding ways of disrupting metabolic pathways as a new means of intervention. Even without the use of culturing methods, recent advances in molecular and biochemical technologies (Fritzsch et al. 2012) should allow detailed investigations of gene function in Cryptosporidium and are likely to have substantial impact in the coming years.

6.5.2 Exploring the Cryptosporidium variome

In addition to harnessing new molecular technologies to improve the annotation and our understanding of the biology of *Cryptosporidium*, such technologies, particularly DNA sequencing technologies, may be applied to assess genomic variation within and among species. For example, at present, a major molecular marker for the exploration of genetic variation within, between and among Cryptosporidium populations is the gp60 gene (Jex and Gasser 2010). This highly polymorphic glycoprotein allows the clustering of *Cryptosporidium* populations into genotypes, and microsatellite variability in the form of a serine/threonine rich glycosylation region, allowing a further subdivision of Cryptosporidium genotypes into subgenotypes (or subtypes) (Strong et al. 2000). Recently, we reviewed the global richness (= the number of sequence types) and diversity (= the relative proportional distribution and/or domination of these sequence types) of all gp60 sequence types reported for C. parvum and C. hominis and the frequency with which they occur among the hosts for which such data are available (Jex and Gasser 2010). This review showed that, of the ~150 subgenotypes (representing 11 C. parvum and 6 C. *hominis* genotypes) for which data were available at that time (i.e., 2010), the majority (85%) of them had been reported from humans. Notably, despite this high richness of subgenotypes, six specific subgenotypes (namely C. hominis IbA10G2, C. homins IbA9G3, C. hominis IeA11G3T3, C. parvum IIcA5G3R2, C. parvum IIaA15G2R1 and C. parvum IIaA18G3R1) were associated with the ~70 % of infections in humans from >40 countries (for which information was available) (Jex and Gasser 2010). Indeed, evidence demonstrates clearly C. hominis subgenotype IbA10G2 to be the cause of the most infamous outbreak of cryptosporidiosis in Milwaukee in 1992 (Zhou et al. 2003).

This finding might have profound implications for the treatment and control of cryptosporidiosis, but the extent to which gp60 sequence is representative of population-level variation in Cryptosporidium is unclear. A recent publication by Tanriverdi et al. (2008) explored population structuring in C. parvum and C. hominis using numerous isolates from a broad geographical range, assessed at several genetic loci, and found evidence of some population structuring but also substantial 'genetic mixing'. This information would appear to suggest that the dominance of a few gp60subgenotypes infecting humans might relate to functional constraints on the genes themselves. Given the role of protein GP15/40 in binding to the host cell membrane, some forms of this molecule (expressed by subgenotypes) are likely to bind effectively to host-derived surface proteins, enabling subsequent cellular invasion. Understanding whether human cryptospordiosis is largely dominated by six subgenotypes of C. parvum and C. hominis and the molecular mechanisms linked to infectivity may be a key to the control of human cryptosporidiosis, with implications for the development of new intervention approaches. Antibody responses against proteins GP15 and G40 have been detected (by immunoblot) in the sera from humans following cryptosporidiosis (Strong et al. 2000). Interestingly, a recent study (Preidis et al. 2007) showed that recombinant C. hominis (but not C. parvum)

GP15 stimulated the production of interferon-gamma in whole blood samples taken from donors that were seropositive for *Cryptosporidium*. As noted by the authors of that study (Preidis et al. 2007), interferon-gamma is known to play a key role in the host immune response to *Cryptosporidium* infection (Jex et al. 2010). Although this research hints to recombinant *C. hominis* GP15 as a potential vaccine, the absence of a similar response against *C. parvum* GP15 (Preidis et al. 2007) suggests speciesor genotype-specific immunogenic responses. Understanding, on a genomic level, the richness and diversity of genetic variants of *Cryptosporidium* infecting humans might have implications for vaccine development against various species of *Cryptosporidium*.

6.6 Concluding Remarks

The year 2007 marked the centenary of the discovery of *Cryptosporidium*. The past century has seen immense changes in our understanding of these organisms, with most advances emerging in the last three decades. That this time-scale corresponds to the advent of PCR and the dawn of the molecular biological age is no co-incidence. The application of PCR technology has revolutionized our understanding of *Cryptosporidium*, from systematics to transmission dynamics, population genetics, diagnostics and molecular/cellular biology. Massively parallel genomic sequencing and high-throughput technologies are just emerging. These platforms bring an analytical and exploratory power beyond anything that we have seen in biology to date, and mark a new era in the biological and biomedical sciences. In the field of *Cryptosporidium* research, the application of these emerging technologies has only just begun and, yet, the promise of this research captures the imagination.

Clearly, whole-genome sequencing of Cryptosporidium from humans and other animals should be considered a priority as is the case for an ongoing effort to sequence additional Cryptosporidium species. In previous years, such research would have been considered unaffordable and far too time consuming. However, given the capacity of modern sequencing devices, this is no longer the case. Benchtop sequencing platforms, such as the Illumina MiSeq, coupled to sample multiplexing technologies, now provide sufficient data to sequence complete *Cryptosporidium* genomes to a depth of \geq 100-fold for less than ~\$100 per sample. New developments in DNA library construction further facilitate this research, with the sequencing of whole genomes from nanograms (or less) of genomic DNA being now possible (Ling et al. 2009; Grindberg et al. 2011; Bankevich et al. 2012). Such research can be further enabled through the use of whole genome amplification (WGA) technologies, which allow the synthesis of micrograms of DNA from nanogram or even pictogram amounts of starting template. These methods, also coupled to other 'omic' technologies, allow us to explore the biology of cryptosporidia from humans and other animals to depths never before achieveable.

Nonetheless, significant challenges remain, not least the accessibility of phenetically and genetically well-defined *Cryptosporidium* isolates ("strains") to underpin research efforts. Limitations associated with in vitro cultivation as well as the lack of a known function for most proteins encoded in *Cryptosporidium* genomes represent significant challenges. Despite the complexity of the technologies and the enormity of 'systems biology', as always, addressing these challenges will require persistance, significant technical skill, and the imagination and innovation to solve many unforseen obstacles. After 100 years of research and advances in this field, perhaps Edmund Tyzzer would be pleased to see that some things never change.

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Chapter 7 From Genome to Proteome: Transcriptional and Proteomic Analysis of *Cryptosporidium* **Parasites**

Jonathan M. Wastling and Nadine P. Randle

Abstract The ability to generate large-scale transcriptome and proteome datasets has changed the landscape in parasite biology. These data enable an integrated, whole organism approach to understanding how parasites function and interact with their hosts. The difficulty in propagating *Cryptosporidium* means that work has focused on the oocyst and sporozoite stages of the parasite. Transcriptional studies using expressed sequence tags (ESTs), microarrays and quantitative real-time PCR (qRT-PCR) have given a valuable insight into how both the parasite and host cells respond to infection. In addition, global proteomics analyses have characterised the expressed proteomes of oocysts and sporozoites of *C. parvum*. Currently, *Cryptosporidium* research is lagging behind some other pathogens in terms of global 'omics' analyses. However, there have been significant technological and bioinformatics advances in transcriptome and proteome analyses in recent years, which are set to continue. Exploiting these technologies and capitalising on the resulting "systems-biology" data mean that exciting times are ahead in the field of *Cryptosporidium* biology.

7.1 Introduction

The availability of the genome sequences for *Cryptosporidium parvum* (Abrahamsen et al. 2004) and *C. hominis* (Xu et al. 2004) has greatly advanced our knowledge of the biology of these parasites, but is only part of the story of trying to understand the evolutionary success of *Cryptosporidium*. Although genomes provide information concerning the potential repertoire of genes in *Cryptosporidium*, they provide no information on the actual expression of these genes, or on the translation of genes into functional proteins. In order to investigate the most

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interesting biological problems, such as how *Cryptosporidium* adapts to its unique biological niche, how it controls development through its life-cycle and how it interacts with its host environment, it is necessary to determine the expression of both genes and proteins. Analysis of the mRNA transcriptome and protein expression by proteomics has the potential to yield a wealth of "systems-biology" expression data to help answer some of these essential questions.

The difficulty in propagating in *Cryptosporidium* spp. in vitro has hampered gene and protein expression studies. While it appears that it might be possible to maintain *C. parvum* in cell culture for a period of time (Arrowood 2002; Current and Haynes 1984; Hijjawi et al. 2001, 2002; Upton et al. 1994), insufficient material can be produced for the later life-cycle stages to perform comprehensive transcriptomic and proteomic analyses. The oocyst and sporozoites are currently the only life-cycle stages of *C. parvum* that are readily accessible for transcript and proteome analyses, thus, such studies have been restricted to these stages.

This chapter describes the recent advances in the transcriptomic and proteomic analyses of *Cryptosporidium* and how these data can be used to further the understanding of its biology.

7.2 Global mRNA Transcriptional Analysis

The transcriptome describes all the RNA transcripts present in a cell and is highly dynamic, reflecting all the genes that are being actively expressed at a given point in time. Examining the expression profiles of mRNA transcripts in a population of cells gives a good understanding of the functional processes occurring. Gene expression profiles vary depending on the cell type, developmental stage of the organism and environmental conditions, thus unlike the relatively static genome, the transcriptome is highly dynamic and responsive to biological events. Two main approaches can be taken in order to characterise and quantify the transcriptome of a cell or organism: hybridisation- or sequence-based methods. A brief overview of the main techniques is given here, but there are numerous previously published general reviews of these to which reference should be made (Holt and Jones 2008; Mutz et al. 2013; Wang et al. 2009).

Microarrays are the most widely used hybridisation-based technique for simultaneously measuring gene expression levels for thousands of genes, identifying SNPs and for comparative genome analysis (Gobert et al. 2005). They consist of glass or silicon slides with thousands of DNA probes attached in an ordered array; these probes correspond to mRNA sequences from the target species and can be oligonucleotides, cDNA or PCR products. Hybridisation to complementary DNA (cDNA) arrays is more specific than to oligonucleotide arrays, but the latter are considerably cheaper. When performing a microarray experiment, cDNA is usually labelled with a fluorescent dye, such as Cy3 or Cy5, prior to being hybridised to the microarray. Samples can be labelled with different dyes and hybridised to the same array for comparative studies. When the sample binds to the probe, it emits a signal that can be detected at a specific wavelength and is proportional to the amount of target sequence that has bound.

Reverse-transcription PCR (RT-PCR) is used to detect mRNA expression, but is relatively non-quantitative. However, this technique can be readily adapted to obtain relative or absolute gene expression levels using quantitative real-time PCR (qRT-PCR). By incorporating fluorescent dyes or probes, such as SYBR Green or TaqMan, into the PCR products as they are amplified, the amount of DNA present can be measured in real-time. This technique is widely used for quantifying gene expression levels.

There have been significant technological advances in recent years, with numerous high-throughput "next-generation" sequencing platforms now available. This technology has revolutionised the extent and complexity of transcriptome sequence data that can be obtained for an organism. RNA-Seq is highly sensitive, with very low background noise and a large dynamic range that can quantify transcripts of very low to very high abundance (Mortazavi et al. 2008). Briefly, a complementary DNA (cDNA) library is generated from RNA by amplification from adaptors attached to either end of the RNA strands. Single-end or paired-end sequences are generate by sequencing from either one or both adaptors, respectively. The cDNA fragments are sequenced in a high-throughput manner to produce reads of 30–1,000 nucleotides (nt), depending on the sequencing platform used. These reads can either be aligned (mapped) to an existing genome sequence or assembled de *novo*; the resulting transcriptional map yields data about both the structure and expression level of each gene (inferred from the number of reads for a given transcript). The SOLiD sequencing platform is based on ligation and generates short reads of up to 50 nt. The Illumina and 454 platforms both take a "sequencing by synthesis" approach; the Illumina sequencers produce reads of 50-300 nt while the 454 produces much longer reads of up to 1,000 nt. The short read lengths produced by SOLiD and Illumina are ideally suited to analysing transcriptomes of good quality, well annotated genomes and can be useful in identifying intron-exon boundaries and sequence variations (Wang et al. 2009). When there is either an incomplete, or no reference genome available the longer reads produced by 454 sequencing are more appropriate as they maximise the possibility of reads overlapping.

A limiting factor with microarrays and PCR product-based transcriptome analyses is the need for good quality genome sequences. Many parasite genomes are incomplete which can present major challenges when designing probes and primers for these types of studies (Chen et al. 2004). The RNA-Seq approach to transcriptome analysis offers considerable advantages over microarrays: it requires less mRNA (a considerable advantage when working with microorganisms like *Cryptosporidium* which are difficult to propagate in vitro), has a much larger dynamic range for accurately detecting and quantifying transcript expression levels, has very low background noise and is highly reproducible. Correctly mapping the mRNA sequence reads to the appropriate genome and/or assembling sequences *de novo* can still present a challenge, although the use of paired end sequencing has made this task substantially easier (Holt and Jones 2008; Roach et al. 1995).

7.3 Proteomics Technologies

Proteins are arguably the most relevant functional component in a biological system; however, they are in a constant state of flux and activation, presenting a significant analytical challenge in comparison to mRNA. The development of mass spectrometry-based proteomics as a routine technology has transformed the way in which protein expression can be analysed. In common with high-throughput transcriptome sequencing, characteristic signatures (peptide fragmentation patterns in the case of proteins) can be matched by bioinformatics to specific genes. This means that the physical data from a mass spectrometer can be de-convoluted to provide high-throughput detection and analysis platforms for proteins, providing a corresponding genome sequence is available. Mass spectrometry (MS) analysis of proteins involves the ionization, separation and detection of molecular ions based on their mass-to-charge ratio (m/z). The resulting spectra are then searched against a corresponding virtual database of peptide mass data. Briefly, the virtual database uses previously established gene models, or translates the genome sequence in all six frames and the protein sequences are then digested in silico to generate every possible peptide sequence. The m/z for each ion detected by the mass spectrometer is then queried against the virtual database to find a corresponding peptide from an annotated protein with the same m/z. A typical proteomics workflow is shown in Fig. 7.1.

In order to reduce the complexity of a sample for MS analysis, proteins can be separated by gel electrophoresis or chromatography before digesting with an enzyme, such as trypsin, to produce peptides. Alternatively, peptides can be separated after enzymatic digestion using methods such as liquid chromatography. For the former, one- and two-dimensional gel electrophoresis (1-DE and 2-DE) have been widely employed to separate proteins but these techniques are most suited to analysing abundant hydrophilic proteins. In 1-DE proteins are separated by their mass so bands frequently contain multiple proteins. Using 2-DE, proteins are first separated according to their isoelectric point and then by mass, producing a highly resolved protein map in which each spot only contains one protein. The 2-DE approach reveals something of the complexity of the proteome, as proteins may appear in multiple spots due to post-translational modifications and alternative splicing.

Multidimensional protein identification technology (MudPIT) uses liquid chromatography (LC) to separate peptides for identification by MS. This can be done by excising bands from a 1-DE gel, digesting them and then performing the separation step or by digesting the proteins in solution and performing two rounds of LC followed by MS analysis. Advances in mass spectrometry instrumentation and protein separation technology have increased the resolution and accuracy of m/z data acquisition. This, when coupled with the use of more sophisticated bioinformatics tools, has led to more reliable and statistically sound protein identifications with much reduced false discovery rates (FDR).

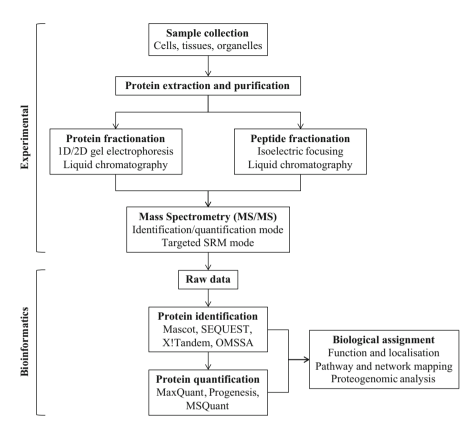


Fig. 7.1 Schematic representing a typical proteomics workflow. The identification and quantification of proteins involves an experimental and bioinformatics stage. In the experimental stage samples are collected, protein is extracted and purified, samples are separated and digested (in either protein or peptide space) and mass spectrometry analysis is performed. The bioinformatics analysis uses the raw spectra data to interrogate protein identification databases and specialised quantification programmes calculate the relative and absolute protein abundances. Biological relevance is then assigned using pathway mapping tools and predictions of the protein function and localisation

7.4 The Transcriptome of *C. parvum* Oocysts and Sporozoites

Large-scale transcriptional analysis from *Cryptosporidium* parasites began with sequenced cDNA libraries of oocysts/sporozoites which generated expressed sequence tag (EST) data (Strong and Nelson 2000). There are currently 98,058 EST transcripts from *C. parvum* and *C. muris* on CryptoDB (version 5.0) (http://cryptodb.org/cryptodb/; Heiges et al. 2006), which provide evidence for 2,386 genes. One of the early studies to profile the genes expressed by *C. parvum* sporozoites sequenced cDNA libraries and generated 567 ESTs (Strong and Nelson

2000). To complement the EST data, and to expand the study to other life-cycle stages, the authors also analysed 1,507 genome survey sequences (GSS) that were generated from randomly sheared genomic DNA. This was performed prior to the release of the genome, but BLAST analyses found that 32 % of the ESTs showed similarity to existing sequences in publicly available databases, leaving a high frequency of sequences with no known homology or function. More recently over 1,000 ESTs were sequenced from oocysts undergoing excystation, this study has shown that transcripts encoding for structural proteins are highly up-regulated during the early stages of the excystation process (Jenkins et al. 2011). In addition, a high percentage of hypothetical proteins were identified as well as a large number of ESTs involved in cell signalling, proliferation and metabolism. In concordance with proteomic evidence, heat shock proteins (HSP) and ribosomal proteins were also abundant (Snelling et al. 2007). In a study to assess the quality of the current gene models, 5'-EST data generated for C. parvum, Toxoplasma gondii and a variety of *Plasmodium* species were compared with existing genome sequences. In contrast to other Apicomplexa, gene annotations for C. parvum were found to be highly reliable, undoubtedly due to the small size of the genome and low frequency of introns (Wakaguri et al. 2009).

Recently, a *C. parvum* microarray (CpArray15K) containing oligonucleotide probes for 3,805 genes has been developed (Zhang et al. 2012). The array, coupled with qRT-PCR, was used to compare gene expression profiles between untreated and UV irradiated oocysts to identify genes involved in the response to environmental stress. In the untreated oocysts, genes associated with protein synthesis (transcription, translation and ribosome biogenesis) were highly expressed. From the high expression levels of ubiquitin and proteasome, the authors suggest that the oocyst may degrade and reuse amino acids. Genes involved in DNA repair, intracellular trafficking and cytoskeleton rearrangement were up-regulated in response to UV irradiation, as were members of the T-complex protein 1 (TCP-1) family, and some thioredoxin-related genes which are known to be associated with responses to stress.

Remarkably, at the time of press, no RNASeq data for *Cryptosporidium* has been published, despite the significant advances in sequencing technologies. However, the first comprehensive in vitro analysis of the *C. parvum* transcriptome has recently been completed (Mauzy et al. 2012). The transcript abundances of 3,302 genes (representing 87 % of the genome) were determined by qRT-PCR. This monumental experiment was designed as a temporal study, in which samples of infected human epithelial cells were collected at seven time points over a 72 h infection period. For each of the genes analysed, transcripts were detected for at least one time point. Although expression levels varied throughout over time, two thirds of the transcripts were detected at every time point. Gene expression patterns were analysed by looking at relative fold changes that were normalised to 18S rRNA expression levels. Distinct expression profiles were associated with different developmental stages, as evidenced by the lack of congruence between metabolic, ribosomal and proteasome protein expression, and 18 s rRNA levels. The number of distinct transcripts detected increased from 6 h post infection, with a third of the

genes being most highly expressed at either 48 or 72 h. When looking at the complete dataset, cluster analysis revealed nine distinct clusters; genes within each cluster did not map to the same chromosomal region indicating that gene expression is independent of chromosome location. From the cluster analysis, expression of transcription related genes peaked at 2 h, proteins involved in translation dominated at 6 h while transcripts related to structural genes, such as myosin and tubulin, were most prevalent at 12 h post infection. An increase in transcript abundance, coinciding with the sexual reproduction of the parasites, was observed from 36 to 72 h. These data suggest that *C. parvum* has a more complex system of transcription regulation than previously thought (Mauzy et al. 2012; Templeton et al. 2004).

7.5 Modulation of the Host Cell Transcriptome by *Cryptosporidium*

Transcriptional studies have been used to determine host-response, as well as parasite gene expression patterns in cell monolayers, including several microarray studies that characterised the host cell (human) responses to infection with C. parvum (Castellanos-Gonzalez et al. 2008; Deng et al. 2004; Liu et al. 2009; Yang et al. 2009, 2010). These studies have shown that significant changes occur in the host cell in response to infection with C. parvum, particularly in genes associated with the inflammatory response, apoptosis, cell proliferation and the cytoskeleton. The composition of the host cell membrane is affected by the presence of the parasite (Yang et al. 2009). One study showed that heat shock proteins and pro-inflammatory cytokines were up-regulated in response to C. *parvum* infection (Deng et al. 2004). Interestingly, the differential expression of actin related genes was also observed with those for actin binding proteins being down-regulated while genes for host actin and tubulin were up-regulated. This corresponds well with previous findings of host cell actin polymerisation and cytoskeleton remodelling in response to C. parvum infection (Chap. 10) (Chen et al. 2001; Elliott and Clark 2000; Elliott et al. 2001). Apoptosis of host cells was observed to be differentially regulated throughout the infection process with antiapoptotic genes being more highly expressed during the first 12 h of infection with C. parvum and no apoptosis being observed in culture; from 24 h post infection, pro-apoptotic genes were up-regulated (Liu et al. 2009). Genes related to apoptosis have been found to be differentially regulated in the host in other studies (Deng et al. 2004; Yang et al. 2009), consistent with previous evidence of host cell apoptosis in C. parvum infected cell monolayers (Griffiths et al. 1994; Widmer et al. 2000). Coupling biopsy infection with microarray analysis revealed that the osteoprotegerin (OPG) gene was up-regulated in response to infection with C. parvum and C. hominis and that OPG expression was greater in response to C. hominis. These findings were confirmed in HCT-8 cells and OPG was also shown to be expressed in response to an in vivo infection of a volunteer with *C. meleagridis*. Thus, a role for OPG in modulating the host cell response to *Cryptosporidium* infection has been proposed (Castellanos-Gonzalez et al. 2008).

A difficulty faced by studies to analyze the in vitro expression of both parasite and host genes during *Cryptosporidium* cell invasion is the relatively low multiplicity of infection that can be achieved using current cell culture techniques. Since the majority of cells remain uninfected in culture, signals from infected cells may appear relatively weak as compared to the majority of uninfected cells. In order to overcome this problem, Yang et al. (2010) used fluorescence-activated cell sorting (FACS) to separate infected and uninfected cells from a single monolayer to compare host cell responses using a human genome microarray. At 24 h post infection, 417 genes were up-regulated and 361 were found to be down-regulated. In sorted cells, glycoproteins predominated in the up-regulated genes of infected cells (Yang et al. 2010); lectin-binding studies confirmed that the sugars on these glycoproteins were mainly N-linked oligosaccharides. This differs from studies using unsorted cells where the majority of the up-regulated genes were related to apoptosis and cell proliferation (Deng et al. 2004; Liu et al. 2009; Yang et al. 2009).

7.6 The Proteome of *C. parvum*

In contrast to other apicomplexan parasites, relatively few proteomic analyses have been performed for *Cryptosporidium* to date. Early studies of *Cryptosporidium* by mass spectrometry were limited to identifying characteristic spectral patterns that could be used to distinguish between species. Prior to the release of the *C. parvum* genome, MALDI-TOF MS was used to identify spectral peaks that could identify *Cryptosporidium* at the genus level and others that could be used to distinguish between *C. parvum* and *C. muris* oocysts (Magnuson et al. 2000). Further work also revealed spectral peaks that were specific to the oocyst and sporozoites (Glassmeyer et al. 2007).

Subsequently, facilitated by the availability of the genome sequence, large-scale global analyses of the expressed proteome of *C. parvum* oocysts and sporozoites have been performed (Sanderson et al. 2008; Snelling et al. 2007). In the first proteomic analysis of *C. parvum*, a combination of LC-MS/MS and iTRAQ isobaric labelling were used to identify around 200 proteins, representing about 6 % of the predicted proteome, from excysted and non-excysted oocysts (Snelling et al. 2007). For both stages of the parasite, the soluble fraction was labelled with iTRAQ reagents for quantitative analysis on a QSTAR QqTOF mass spectrometer and the insoluble fraction was subject to qualitative MudPIT analysis. By using iTRAQ to label the samples, Snelling and co-workers were able to perform a relative quantification of the proteins identified between the excysted and non-excysted parasites. In the soluble fractions, a total of 26 proteins were found to have significantly higher expression levels post-excystation, relative to unexcysted oocysts. These proteins include a variety of heat shock proteins,

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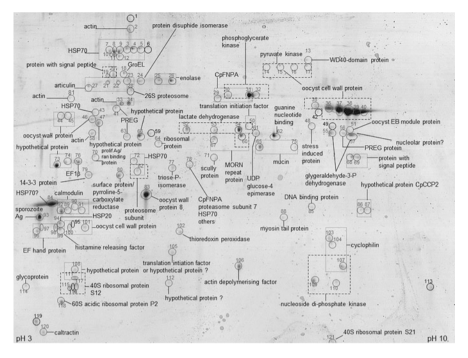


Fig. 7.2 2-DE analysis of *Cryptosporidium parvum* sporozoites. Proteins were resolved using a broad non linear pH3-10 gradient followed by separation on a 12.5 % denaturing poly acrylamide gel before staining with Coomassie Colloidal (Sanderson et al. 2008)

ribosomal proteins and metabolic enzymes as well as three apicomplexan-specific and five *Cryptosporidium*-specific proteins. Several of these proteins contained signal peptides, were predicted to be secreted and were hypothesised to play a role in host cell invasion by the parasite.

In the second, and most comprehensive of these studies, 1,237 non-redundant proteins were identified from excysted oocysts and sporozoites, equivalent to 30 % of the predicted proteome (Sanderson et al. 2008). To maximise the coverage of the proteome, and increase the confidence in the proteins identified, three complementary approaches were used: 1-DE LC-MS/MS, 2-DE LC-MS/MS and MudPIT. Proteins excised from 1-DE and 2-DE gels were digested with trypsin and analysed by either MALDI-ToF or LTQ ion trap mass spectrometry; the resulting spectra were used to interrogate a *C. parvum* genome database in Mascot® (Perkins et al. 1999) (Fig. 7.2). In the MudPIT experiment, spectra from lysed soluble and insoluble fractions were searched using SEQUEST® (Eng et al. 1994). A total of 642, 282 and 1,154 non-redundant proteins were identified from the 1-DE, 2-DE and MudPIT analyses, respectively. Approximately 20 % of the proteins identified were done so based on a single peptide hit from one platform and thus would not have been identified if a single approach had been taken.

Analysis of protein function revealed structural information for 1,165 of the non-redundant proteins identified, with 213 (18 %) predicted to contain transmembrane domains (about 50 % of these proteins have no known function). The MudPIT dataset contained a greater proportion of hydrophobic proteins than the other two platforms. Putative functional categories were assigned to proteins using a combination of gene ontology (GO) classifications and homology and literature searches. However, no information was available for approximately 40 % of the identified proteins, which were designated as unclassified. Key findings include peptide evidence for 18 of the mitochondrial and iron-sulphur metabolism genes predicted in the *C. parvum* and *C. hominis* genomes, consistent with the theory that a functional relic mitochondrion is present in *Cryptosporidium* (Putignani et al. 2004). Peptide evidence was also detected for 14 of the predicted micronemal proteins, 12 putative rhoptry (ROP) proteins, a few dense granule proteins and a large number of surface proteins.

7.7 Sub-proteome Analysis of C. parvum Sporozoites

Despite huge advances in the sensitivity of mass-spectrometry as a protein discovery tool, global proteomics experiments are still not able to cover the complete depth of the expressed proteome in a single experiment. One way to increase the depth of coverage is to simplify the proteome by fractionation prior to analysis, creating "sub-proteomes" that contain fewer proteins. For example, sub-proteomes for apical invasion related organelles have been characterised for a range of apicomplexan parasites including the rhoptries of *Toxoplasma gondii*, *Eimeria tenella* and several *Plasmodium* species (Bradley et al. 2005; Bromley et al. 2003; Sam-Yellowe et al. 2004); the micronemes of *E. tenella* (Bromley et al. 2003) and excreted/secreted antigens of *T. gondii* (Zhou et al. 2005). In *Cryptosporidium*, there have been relatively few equivalent studies, largely due to the difficulties of obtaining sufficient amounts of parasite material with which to perform organelle fractionation by techniques such as density gradient centrifugation.

In one study to investigate how sporozoites are tethered to the oocyst wall, mass spectrometry analysis was employed to identify the key components of the oocyst wall and affinity purified glycoproteins from the sporozoites (Chatterjee et al. 2010). Excysted and sonicated oocysts were centrifuged through sucrose to isolate the walls, while the glycoprotein fractions were prepared using Sepharose columns. The authors found that the *Cryptosporidium* oocyst wall proteins (COWPs) constitute about 75 % of the proteins identified by mass spectrometry in the oocyst walls. The dominant oocyst wall protein in *C. parvum* is COWP1, with more than 50 % of peptides identified by mass spectrometry belonging to this protein. COWP8 and COWP6 are also relatively abundant while COWP2, COWP3 and COWP4 were only detected at very low levels and COWP5, COWP7 and COWP9 were not detected. These results confirmed the previous

findings of Sanderson et al. (2008). Another protein detected at low levels associated with the oocyst wall was gp40/15, a mucin-like glycoprotein that localises to the sporozoites surface and inner surface of the oocyst wall (Chatterjee et al. 2010; O'Connor et al. 2007). The authors also identified five possible oocyst wall proteins (POWPs) that were detected at low levels and are unique to *C. parvum*. When analysing the affinity purified glycoprotein components of the oocyst, gp40/15, gp900 and six other glycoproteins (CpMPA1 to CpMPA6) were identified, all are unique to *C. parvum*.

7.8 Publicly Accessible Transcriptomic and Proteomic Databases for *Cryptosporidium*

CryptoDB (http://cryptodb.org/cryptodb/) is an integrated database of genomic and functional genomic data from representative strains of *C. parvum*, *C. hominis* and *C. muris* (Heiges et al. 2006). It contains the most recently annotated genome sequences for each species alongside experimental data from the research community. CryptoDB is a part of EuPathDB (Aurrecoechea et al. 2007), a freely available and accessible resource providing easy access to a range of "-omics" data for *Cryptosporidium* and other protozoan parasites. Transcriptomic and proteomics data from a variety of studies, primarily on *C. parvum*, have been fully integrated into CryptoDB. By mapping the transcript and peptide data generated from these global expression studies to the genome, proteins that were not predicted by the original gene models have been verified. These expression data have provided empirical evidence that has been instrumental in improving the annotation of the genome (Sanderson et al. 2008).

Currently, CryptoDB (version 5.0) contains 98,058 ESTs, providing transcript evidence for 1,806 *C. parvum* (70 560 ESTs) and 580 *C. muris* RN66 (27,498 ESTs) genes along with RT-PCR temporal expression data for *C. parvum* (Mauzy et al. 2012). In addition, proteomics data from five independent studies have been integrated into CryptoDB (Sanderson et al. 2008; Snelling et al. 2007); at the moment this data is limited to the IOWA II strain of *C. parvum*. Expression data for 1,320 proteins, representing approximately a third of the predicted proteome, has been uploaded to the database for the oocyst and sporozoite stages of the parasite. This is a reasonable coverage (for both transcript and protein evidence) given that these life-cycle stages are unlikely to express stage specific proteins from the sexual stages and merozoites, although lower than for some other Coccidia, such as *Toxoplasma gondii* (Xia et al. 2008).

Genes of interest can be searched for by a variety of methods such as, by gene identifier (e.g. cgd6_2090) or by text search (e.g. oocyst wall protein), as illustrated in Fig. 7.3. Batch searches, such as "identify genes based on EST evidence", can also be performed to search the database for experimental data. An overview of the

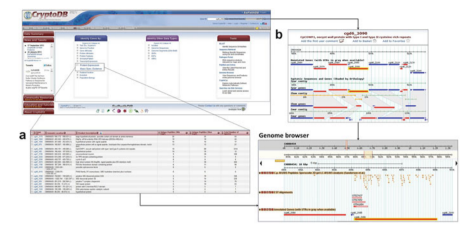


Fig. 7.3 Visualisation and mining of proteomic and transcriptomic data on CryptoDB (http:// cryptodb.org/cryptodb/). In this example data for COWP1 (cgd6_2090) has been mined by (a) using tools such as 'Identify genes based on Mass Spec Evidence' and (b) gene ID or text searching. The resulting peptide, EST, sequence and gene model data can then be viewed in Genome Browser

data available for individual genes is displayed, including the location of the gene in the genome, any orthologs to it in other *Cryptosporidium* species along with any transcript and/or peptide evidence for the gene product and functional information. Data can also be visualised in GBrowse (Stein et al. 2002) for more in-depth data mining.

An alternative proteomics database is EPICDB, which contains protein predictions and mass spectrometry data for *T. gondii* and *C. parvum* (Madrid-Aliste et al. 2009). EPICDB also includes functional annotations, ESTs, open reading frames (ORFs) and experimental data. The curated Library of Apicomplexan Metabolic Pathways (LAMP) provides core metabolic pathways for *C. parvum*, *C. hominis* and *C. muris*, along with other apicomplexan parasites (Shanmugasundram et al. 2013). This database combines proteomic, genomic and transcriptomic data from EuPathDB with evidence from the literature to reconstruct metabolic pathways and networks.

7.9 Conclusions

The analysis of gene and protein expression in pathogens and their hosts has been transformed by the advent of high-throughput transcriptional analysis and proteomics. Both techniques have been applied to the study of *Cryptosporidium*, although there remains much to do. The depth of proteomics coverage of *Cryptosporidium* is still relatively poor compared to other *Apicomplexa* such as *Toxoplasma* and

Plasmodium, and almost nothing is known about gene expression in life-stages other than the oocyst and sporozoite. Nonetheless, the application of both transcriptional and proteomics analysis techniques to Cryptosporidium research has done much to help build a platform on which more targeted biological experiments can now be performed. However, both types of data, valuable as they are, need to be seen in context. The increasing abundance of simultaneously collected transcriptomics and proteomics data from both pathogens and their hosts reveals that the relationship between protein and transcript measurements is not always as faithful as might be supposed (Wastling et al. 2009). Since proteins are the functional molecules in a biological system, this adds complexity to making functional interpretations from steady state mRNA expression data alone. Moreover, the abundance of a protein is not always the most important feature, with alternative splicing and posttranslational modifications (PTMs), such as phosphorvlation, also influencing its activity. Finally, proteins do not act in isolation and protein-protein interactions play an important role in the functioning of a cell/ organism. All of this means that we are some way off from developing a truly comprehensive "systems" view of *Cryptosporidium* biology until we are able to provide greater coverage and complexity in transcriptional and proteomic studies. We anticipate these datasets will be acquired and made available to the research community in the near future, providing significant advances in our knowledge and understanding of this fascinating parasite.

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Chapter 8 Cryptosporidium Metabolism

Guan Zhu and Fengguang Guo

Abstract Rather than the presence of unique metabolic pathways, it is the absence of many pathways that characterizes the metabolism of *Cryptosporidium*. In fact, this genus of parasites has lost its ability of synthesizing de novo virtually all nutrients such as amino acids, nucleotides and fatty acids, thus relying on a large number of transporters to scavenge nutrients from the host. Members of this genus lack an apicoplast and associated pathways that are present in other apicomplexans. They lack cytochrome-based respiration, and rely mainly on glycolysis for energy production. Core metabolic pathways are highly streamlined, and redundancy is rare. These features make *Cryptosporidium* different from other apicomplexans. This chapter summarizes these features based on the analysis of genome sequences and published biochemical data in the context of drug targets and drug development.

8.1 Introduction

The genus *Cryptosporidium* comprises many species with different host specificities. *C. parvum* and *C. hominis* are the major human pathogens. The genomes of *C. parvum* and *C. hominis* have been reported (Abrahamsen et al. 2004; Xu et al. 2004), that of *C. muris* has been sequenced but remains to be published, and many more are being sequenced by an NIH-funded "Comparative Genomics of *Cryptosporidium* Species" project (http://www.genome.gov/26525388). The three sequenced genomes are accessible at GenBank and EuPathDB (http://www.EuPathDB.org). The availability of these genome sequences has greatly contributed to our understanding of parasite biology, and provided new opportunities to study *Cryptosporidium* metabolism. Currently, our

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knowledge on *Cryptosporidium* metabolism is mainly derived from genome annotations and analyses. A growing but limited number of enzymes have been investigated for their functions, biochemical features and potentials as therapeutic targets. However, systematic studies at the pathway level are still limited. Two reviews on *Cryptosporidium* biochemistry and metabolism were published as a book chapter in 2008 and a journal article in 2010 (Rider and Zhu 2010; Zhu 2008), These publications were mainly based on the *C. parvum* and *C. hominis* genomes and biochemical data available then. In this chapter, we will highlight the major metabolic features in *Cryptosporidium* with the addition of the newly sequenced *C. muris* genome and up-to-date biochemical data.

8.2 General Features of Cryptosporidium Metabolism

Phylogenetic and phylogenomic analyses have consistently placed Cryptosporid*ium* at the base of the Apicomplexa with a closer relationship to gregarines than coccidia and haematozoa (Barta and Thompson 2006; Templeton et al. 2010; Zhu et al. 2000a). This view differs from the conventional taxonomy that considers Cryptosporidium as a sister group to the intestinal coccidia. The divergence of cryptosporidia from coccidia is also supported at the genomic and metabolic level. *Cryptosporidium* species have highly compact genomes (i.e., ~ 10 Mb), which are 3-5 times smaller than those of other apicomplexans and feature short intergenic sequences and very few introns (Abrahamsen et al. 2004; Xu et al. 2004). More importantly, this genus of parasites has lost the ability to synthesize virtually any nutrients de novo, such as amino acids, nucleotides and fatty acids. Cryptosporidium also lacks an apicoplast and mitochondrial genomes and associated metabolic pathways that are present in coccidia and haematozoa (Abrahamsen et al. 2004; Xu et al. 2004; Zhu et al. 2000b). Members of the genus possess a mitochondrial remnant that lacks the cytochrome-based respiratory chain, and retains only limited functions such as the assembly of ion-sulfur clusters (Lei et al. 2010; Kang et al. 2008; Keithly et al. 2005; Slapeta and Keithly 2004; Roberts et al. 2004; Riordan et al. 2003). Therefore, Cryptosporidium metabolism is extremely simplified. This feature differentiates these species from the evolutionarily more closely related gregarines that are able to synthesize many nutrients (e.g., Templeton et al. 2010).

Because *Cryptosporidium* cannot synthesize most nutrients de novo, it relies on a large family of transporters to scavenge nutrients such as amino acids (~11 transporters), sugars (~9), and nucleotides (at least one) (Abrahamsen et al. 2004; Xu et al. 2004). About 24 putative ATP-binding cassette (ABC) transporters can be identified in the *C. parvum* genome, but their substrates remain to be determined (Benitez et al. 2007; Li and Mun 2005; Bonafonte et al. 2004; Zapata et al. 2002; Perkins et al. 1999). ABC transporters are a large family of proteins with various substrate preferences including ions, sugars, amino acids, peptides, lipids, sterols and drugs. Therefore, many of them could be utilized by the parasite to scavenge

nutrients. The parasite also possesses at least seven P-type ATPases (P-ATPases) that are mostly involved in cation transport. Among them, a putative Ca^{2+} -ATPase and a heavy metal ATPase with binding specificity for reduced copper [Cu(I)] have been reported (LaGier et al. 2002, 2001; Zhu and Keithly 1997). One of the CpATPases belongs to the phospholipid transporter family and may be involved in lipid transport or membrane remodeling.

8.3 Carbohydrate and Energy Metabolisms

Carbohydrates are a source of energy and serve as building blocks for various biomolecules. Cryptosporidium is able to synthesize amylopectin as an energy storage polysaccharide. This biosynthetic capability is supported by earlier biochemical analyses and the presence of two glycogen branching enzymes (Harris et al. 2004; Zhang et al. 2012). The parasite can use polysaccharides, disaccharides or hexoses (e.g., glucose) to produce pyruvate and acetyl-CoA via the glycolytic pathway (Fig. 8.1). However, it employs two pyrophosphate-dependent phosphofructokinase (PPi-PFK) isoforms, rather than an ATP-PFK, to minimize the consumption of ATP. It also uses a unique pyruvate: NADP⁺ oxidoreductase comprised of pyruvate:ferredoxin oxidoreductase (PFO) and P450 reductase domains, rather than a pyruvate dehydrogenase complex to convert pyruvate into acetyl-CoA (Abrahamsen et al. 2004; Ctrnacta et al. 2006; Rotte et al. 2001). It has been speculated that NADP⁺ oxidoreductase is associated with the anti-cryptosporidial action of nitazoxanide (NTZ) that is currently the only drug approved by the FDA to treat cryptosporidial infections in immune-competent patients (Coombs and Muller 2002). Unlike classic inhibitor-enzyme interactions, NTZ may not act on NADP⁺ oxidoreductase. Rather, it is converted to a biotoxic free radical molecule by the PFO domain, similar to the reductive activation of 5-nitroimidazole metronidazole by PFO in the anaerobic protists *Trichomonas* and *Giardia* (Leitsch et al. 2011; Crossnoe et al. 2002; Yarlett et al. 1986). To better understand the enzymes in the carbohydrate pathway, protein crystals have been obtained for C. parvum LDH, glyceraldehyde 3-phosphate dehydrogenase (G3PDH), and pyruvate kinase (Nguyen et al. 2011; Cook et al. 2009; Senkovich et al. 2005). The crystal structures of pyruvate kinase and triosephosphate isomerase are resolved, and the structure of the active site of pyruvate kinase displays no obvious difference compared to the human homologue (Nguyen et al. 2011; Cook et al. 2009).

There are at least two types of cryptosporidial mitochondrial remnants: the "intestinal-type" *C. parvum* and *C. hominis* lack both the tricarboxylic acid (TCA) cycle and the cytochrome-based respiratory chain (Abrahamsen et al. 2004; Xu et al. 2004); whereas the "gastric-type" *C. muris* similarly lacks the respiratory chain but, based on the *C. muris* genome annotation, retains a complete set of enzymes for the TCA cycle and a type II NADH dehydrogenase (unpublished). The absence of an electron transport chain in all three species indicates that their mitochondria are unlikely to be a major source of energy,

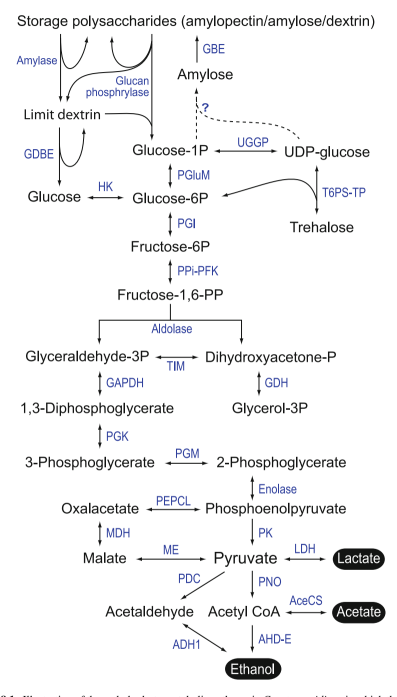


Fig. 8.1 Illustration of the carbohydrate metabolic pathway in *Cryptosporidium*, in which the core components are glycolytic and fermentative enzymes. Abbreviations: *AceCS* acetyl-CoA synthetase (also known as acetate-CoA ligase), *ADH1* alcohol dehydrogenase 1 (monofunctional), *ADH-E* type E alcohol dehydrogenase (bifunctional), *GDH* glycerol phosphate dehydrogenase, *GAPDH* glyceraldehyde phosphate dehydrogenase, *GBE* glycogen branching enzyme, *GDBE* glycogen

although it is possible that a certain level of electron potential may be generated via the type II NADH dehydrogenase in *C. muris*. Without respiration, the TCA cycle is possibly used by *C. muris* to supply intermediate metabolites. Additionally, all three genomes encode a plant-type alternative oxidase, which may be involved in detoxification of oxygen and/or generating certain electron potential.

Cryptosporidium does not carry a mitochondrial genome and the machinery for the replication, transcription and translation of organellar genomes. In addition to the enzymes discussed above, their nuclear genomes also encode a number of other proteins with mitochondrial-targeting signals, which include several translocases of outer and inner membranes, heat-shock proteins (HSPs), solute carriers, nucleotide anti-porters, ferredoxin and ferredoxin reductase and a small set of enzymes involved in ion-sulfur [Fe-S] cluster assembly (Abrahamsen et al. 2004; Lei et al. 2010; Kang et al. 2008; Mogi and Kita 2010; LaGier et al. 2003). It is believed that the [Fe-S] cluster assembly is one of the core functions retained in the mitochondrial remnant.

Collectively, we may conclude that Cryptosporidium relies mainly on glycolysis to produce energy. This notion is also supported by the presence of fermentative enzymes for producing three organic end products to avoid the accumulation of pyruvate and acetyl-CoA at the end of glycolysis (Fig. 8.1). These include lactate produced by lactate dehydrogenase (LDH), acetic acid by acetyl-CoA synthetase (AceCS; also known as acetate-CoA ligase, AceCL) and alcohol by a type-E bifunctional alcohol dehydrogenase (ADH-E) from acetyl-CoA or monofunctional ADH coupled with pyruvate decarboxylase from pyruvate (Zhang et al. 2012). Among them, LDH and ADH are bacterial-type enzymes. Cryptosporidium LDH originated from malate dehydrogenase (MDH) by a relatively recent gene duplication event which occurred after this genus separated from other apicomplexans. In fact, all apicomplexan MDH and LDH are bacterial-type, derived from an α-proteobacterial MDH (Zhu and Keithly 2002; Madern et al. 2004). Interestingly, a recent microarray-based transcriptome analysis also revealed that LDH has the highest level of expression among all genes in C. parvum oocysts, suggesting that the parasite mainly depends on LDH to keep the glycolytic pathway unobstructed in the external environment (Zhang et al. 2012).

Cryptosporidium possesses a plant-type pathway for synthesizing trehalose, which is accomplished by UDP-glucose/galactose pyrophosphorylase (UGGP) and a bifunctional enzyme fusion containing trehalose-6P synthase and trehalose phosphatase (T6PS-TP) (Yu et al. 2010). The presence of trehalose in *C. parvum* oocysts has been confirmed biochemically (Yu et al. 2010). However, the genome

Fig. 8.1 (continued) debranching enzyme, *HK* hexokinase, *LDH* lactate dehydrogenase, *MDH* malate dehydrogenase, *ME* malic-enzyme, *PDC* pyruvate decarboxylase, *PEPCL* phosphoenol-pyruvate carboxylase, *PGI* phosphoglucose isomerase, *PGluM* phosphoglucose mutase, *PGK* phosphoglycerate kinase, *PGM* phosphoglycerate mutase, *PK* pyruvate kinase, *PNO* pyruvate: NADP⁺ oxidoreductase, *PPi-PFK* pyrophosphate-dependent phosphofructokinase, *TGPS-TP* tre-halose-6-phosphate synthase-trehalose phosphatase, *TIM* triosephosphate isomerase, *UGGP* UDP-galactose/glucose pyrophosphorylase

lacks trehalase, suggesting that the parasite is unlikely to reuse trehalose as a carbon source, unless using reversed reactions by UGGP and T6PS-TP. This differs from the intestinal coccidian *Eimeria* that has a trehalase, but lacks UGGP and T6PS-TP. Instead, *Eimeria* possesses a mannitol cycle that is absent in most other apicomplexans, including *Cryptosporidium* (Coombs and Muller 2002; Schmatz 1989). Trehalose and mannitol are known to function as anti-desiccants, antioxidants or protein-stabilizing agents in microorganisms, plants and some invertebrates, thus likely playing an important role in protecting the parasite against environmental stress.

The glycolytic pathway also provides GDP-mannose derived from fructose-6P or mannose-6P for N-glycan biosynthesis. Cryptosporidium appears to have a complete set of enzymes for synthesizing N-glycans in the lumen of the endoplasmic reticulum (ER) (e.g., various asparagine-linked glycosylation [ALG] transferases and an oligosaccharidyl-lipid flippase RFT1). N-glycan synthesis is also connected to the GPI (glycosyl-phosphatidyl-inositol) anchor synthesis. Like other apicomplexans and protists, these parasites lack enzymes to make more complex N-glycans in the Golgi apparatus that are common in fungi and plants. The C. parvum genome encodes ~30 mucin-like proteins, many of which are (or are predicted to be) membrane or secretory proteins based on the presence of signal peptides (e.g., Cevallos et al. 2000a, b; Barnes et al. 1998; Chatterjee et al. 2010). Most of the mucins contain both N- and O-glycosylation sites, and at least four enzymes involved in mucin-type O-glycosylation have been identified in the C. parvum genome, including UDP-N-acetyl-D-galactosamine-polypetide N-acetylgalactosaminyl transferases (Wanyiri and Ward 2006). The involvement of mucins in parasite attachment to, and invasion of, host cells is being actively investigated. Binding of some of the mucin-like proteins by antibodies can block or reduce infection in vitro and/or in vivo, suggesting that mucins may be targets for developing immunotherapeutics (Wanyiri and Ward 2006).

8.4 Amino Acid Metabolism

Cryptosporidium cannot synthesize any amino acids de novo, which differs from other apicomplexans that possess complete pathways to make at least some amino acids. Instead, it retains only enzymes to interconvert certain amino acids coupled with other metabolic pathways. These include: glutamine synthetase for recycling glutamate produced by GMP synthetase back to glutamine; serine hydroxymethyl transferase for converting glycine to serine within the folate cycle; asparagine synthetase for producing asparagine from aspartate (which may be required to recycle ammonia released by AMP-deaminase); S-adenosylmethionine (SAM or AdoMet) synthetase to catalyze the formation of SAM to serve as an important methyl donor for transmethylation; and S-adenosylhomocysteine (SAH) synthase (SAHS; also known as SAH hydrolase, SAHH) for converting SAH derived from SAM after transmethylation to homocysteine and adenosine.

Among these enzymes, the general molecular and biochemical features of *C*. *parvum* SAHH (CpSAHH) has been characterized. Its inhibitors D-eritadenine and 9-(S)-(2, 3-dihydroxypropyl)adenine [(S)-DHPA] display efficacy at low micromolar levels against growth of *C. parvum* in vitro (Ctrnacta et al. 2007, 2010).

8.5 Nucleotide Metabolism

Most apicomplexans scavenge purines, but are capable of synthesizing pyrimidines de novo. However, *Cryptosporidium* lacks synthetic pathways for both purines and pyrimidines. The highly simplified purine salvage pathway starts with the uptake of adenosine by a nucleoside transporter. Adenosine is converted to AMP by adenosine kinase, IMP by AMP deaminase, XMP by IMP dehydrogenase (IMPDH), and GMP by GMP synthase (Fig. 8.2) (Striepen and Kissinger 2004). IMPDH genes in *C. parvum* and *C. hominis* were acquired from ε -proteobacteria, and differ from the eukaryotic type IMPDHs found in humans, animals and other apicomplexans (Striepen et al. 2002). However, an IMPDH gene has yet to be identified in the current version of the *C. muris* genome. Based on the essential role of this enzyme, *C. muris* needs IMPDH unless it directly scavenges GMP from the host. The *C. parvum* and *C. hominis* IMPDH genes are located at the end of chromosome 6. As this region is not represented in the sequenced *C. muris* genome, it is likely that a *C. muris* IMPDH gene is located in an unsequenced region.

Because *Cryptosporidium* lacks hypoxanthine-xanthine-guanine phosphoribosyl transferase to serve as an alternative purine salvaging pathway, blocking this AMP-GMP pathway can effectively kill the parasite. The bacterial-type IMPDH that is highly divergent from humans and animals has been considered an attractive drug target. Its protein structure has been determined, and a number of potent inhibitors have been identified and are being evaluated for drug development (Johnson et al. 2013; Gorla et al. 2012; Sharling et al. 2010; Umejiego et al. 2004, 2008).

For pyrimidine salvaging, Cryptosporidium may utilize uracil, uridine and cystidine by converting them to UMP and CMP by uridine kinase (UK) and a bifunctional enzyme with UK fused to uracil phosphoribosyltransferase (UK-UPRT) (Fig. 8.2). Thymidine can also be used and converted to dTMP by a bacterial-type thymidine kinase (TK), which is subsequently converted to dUMP by thymidylate synthase (TS) coupled with the folate cycle, and to dCMP by dCMP deaminase. dCMP, CMP and UMP can be further converted to dCDP/dCTP, CDP/CTP and UDP/UTP by a multi-functional UMP kinase and UDP kinase. The pyrimidine nucleotide pathways are interconnected by three а ribonucleoside-diphosphate reductase enzyme. Therefore, inhibition of a single pathway may be insufficient to block the pyrimidine synthesis, unless interconvertion is restricted by the presence of rate-limiting enzymes and/or if the supply of any single source of precursors is limited. Although the pyrimidine salvaging pathways appear to be redundant, a recent study has shown that

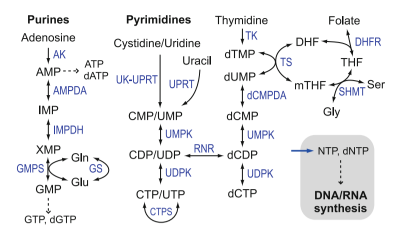


Fig. 8.2 Streamlined purine and pyrimidine salvaging pathways and folate cycle in *Cryptosporidium*. Abbreviations: *AK* adenosine kinase, *AMPDA* AMP deaminase, *CTPS* CTP synthase, *dCMPDA* dCMP deaminase, *DHFR* dihydrofolate reductase, *GMPS* GMP synthase (glutaminehydrolyzing), *IMPDH* IMP dehydrogenase, *RNR* ribonucleoside-diphosphate reductase, *SHMT* serine hydroxymethyl transferase, *TK* thymidine kinase, *TS* thymidylate synthase, *UDPK* UDP/CDP kinase, *UK* uridine kinase, *UMPK* UMP/CMP kinase, *UPRT* uracil phosphoribosyltransferase

TK-mediated pro-drug activation may be utilized as an effective strategy for treating cryptosporidiosis (Sun et al. 2010).

In the folate cycle, dihydrofolate reductase and thymidylate synthase (DHFR-TS) are fused into a bifunctional enzyme in apicomplexans and some other protists (Vasquez et al. 1996). The linker between the DHFR and TS domains in *Cryptosporidium* is unique as it contains an 11-residue α -helix with extensive interactions with the opposite DHFR-TS monomer of the homodimeric enzyme (O'Neil et al. 2003). The active site of *C. parvum* DHFR contains unique residues that are analogous to the point mutations associated with antifolate resistance in other DHFRs, suggesting CpDHFR may be intrinsically resistant to some antifolate inhibitors (Vasquez et al. 1996). However, several novel CpDHFR inhibitors have been identified using a yeast complementation system and structure-based virtual screens, but their anti-cryptosporidial activity in vitro or in vivo remains to be determined (Senkovich et al. 2009; Martucci et al. 2009; Bolstad et al. 2008; Popov et al. 2006; Anderson 2005; Lau et al. 2001; Brophy et al. 2000).

8.6 Lipid Metabolism

Fatty acids are a source of energy in many organisms and major components of all biomembranes. However, *Cryptosporidium* is unable to use fatty acids as an energy source due to the absence of the β -oxidation pathway. It also lacks an apicoplast and

its associated pathways such as isoprenoid synthesis and the Type II fatty acid synthase (FAS) system. Therefore, the parasite cannot synthesize fatty acids (Zhu 2004). However, Cryptosporidium possesses a 25-kb intronless Type I FAS gene that resembles bacterial polyketide synthase (PKS) and predicts a ~900 kDa megasynthase comprised of at least 21 enzymatic domains (Zhu et al. 2000c). The basic biochemical features have been studied using recombinant proteins. Its N-terminal loading unit containing an acyl-ligase (AL) and an acyl-carrier protein (ACP) has a substrate preference towards long chain fatty acids (LCFAs), indicating that CpFAS1 functions as a fatty acid "elongase" rather than synthesizing fatty acids (Zhu et al. 2004). This notion is further supported by functional analysis of its C-terminal reductase domain that is only active with very long chain (VLC) fatty acyl-CoAs (i.e., >C20:0) (Zhu et al. 2010). The reductase domain-catalyzed reductive reaction may release final products as fatty acvl aldehvdes or fatty acvl alcohol, which differs from classic Type I FAS (in humans and animals) and Type II FAS (in prokaryotes, plants and plastidcontaining apicomplexans) that use thioesterase to release acyl chains as fatty acids by hydrolysis (Zhu 2004). Between the loading unit and the reductase domain are three internal acyl elongation modules, each consisting of a complete set of 6 enzymes: (1) ACP for carrying acyl chains; (2) acyl-transferase (AT) for loading malonyl-CoA and transferring an acyl-chain from the previous module to ACP; (3) ketoacyl-ACP synthase (KS) for condensing a two-carbon (C2) unit from malonyl-CoA into the acyl-chain by a carboxylation reaction; (4) ketoacyl-ACP reductase for the reduction of a keto group; (5) hydroxyacyl-ACP dehydrase for the dehydration of a hydroxyl group; and (6) enoyl-ACP reductase for the reduction of double bonds (Zhu 2004; Zhu et al. 2000c, 2004, 2010). Therefore, at least three C2 units can be added into the acyl precursors (e.g., C16:0 palmitic acid) to form very long fatty acyl chains (e.g., C22:0) that are released as fatty acyl aldehydes or alcohol. Because aldehydes are biotoxic, fatty acyl alcohols are likely the final products that can be produced by two series of reductive reactions.

In parallel to Type I FAS, the *Cryptosporidium* genome also encodes a giant 45-kb intronless PKS, which represents the first PKS discovered in a protist (Zhu et al. 2002). Molecular and biochemical analysis reveals that CpPKS1 is similarly structured as CpFAS1, but contains seven internal acyl elongation modules that lack one or more of the five enzymatic domains. Therefore, elongated acyl chains will contain keto groups, hydroxyl groups and/or double bonds, which are characteristic of polyketides. The CpPKS1 loading unit also displays substrate preference towards LCFAs, suggesting the final product(s) may contain 30 or more carbons (Fritzler and Zhu 2007).

The ACP domains in all types of FAS and PKS systems require a posttranslational modification by phosphopantetheinyl transferase (PPT) to add a prosthetic phosphopantetheine to a serine residue to become a functional holo-ACP. There are two types of PPT with different substrate preferences: SFP-type for activating Type I ACPs and ACPS-type for Type II ACPs. In fact, the types of PPT present in various apicomplexans match well with the types of FAS systems. For example, *Cryptosporidium* possesses only Type I FAS and SFP-PPT, *Plasmodium* has only Type II FAS and ACPS-PPT, while *Toxoplasma* contains both types I and II FAS and both SFP-PPT and ACPS-PPT. The activation of CpFAS1-ACP domains by SFP-PPT has been biochemically characterized and demonstrated (Cai et al. 2005).

In addition to type I FAS and PKS, *Cryptosporidium* possesses another set of enzymes capable of elongating fatty acyl chains, in which the substrates are fatty acyl-CoA thioesters, rather than acyl-ACP. The hallmark enzyme is a long chain acyl elongase (LCE). The biochemical features of CpLCE1 have been characterized using recombinant protein expressed in HEK-293 T cells as its expression in bacterial systems was found to be difficult (Fritzler et al. 2007). CpLCE1 is a membrane protein localized on the surface of sporozoites and the parasitophorous vacuole membrane (PVM). Localization on the surface contrasts with CpFAS1 and CpPKS1 that are mainly cytosolic. CpLCE1 is able to add a single C2 unit to LCFAs with substrate preference towards C14:0 myristoyl-CoA and C16:0 palmitoyl-CoA (Fritzler et al. 2007).

Long chain fatty acyl-CoA synthetase (ACS; aka fatty acid-CoA ligase, ACL) is another family of important enzymes in lipid metabolism. It catalyzes the first reaction in all fatty acid metabolisms by activating free fatty acids to form fatty acyl-CoA thioesters except for the Type I and II FAS/PKS systems. In fact, AL domains in Type I FAS/PKS systems share similar molecular and biochemical properties with ACS, and can also catalyze the formation of fatty acyl-CoA (Fritzler and Zhu 2007). *Cryptosporidium* possesses three ACS genes, which are under investigation in our laboratory. Our data have shown that CpACS enzymes prefer LCFAs as substrates and their inhibitors could inhibit the growth of *C. parvum*, suggesting that ACS can be explored as a novel therapeutic target in the parasite (unpublished observation).

Cryptosporidium also possesses a long-type fatty acyl-CoA binding protein (ACBP) that is responsible for restraining the movement of fatty acyl-CoA and/or forming an acyl-CoA pool in cells. This function is important, as free fatty acyl-CoA may be harmful to cellular membranes due to its "detergent effect" if it is not restrained or immediately routed into other metabolic pathways. CpACBP1 is also a membrane protein localized to PVM. It prefers binding to LC and VLC fatty acyl-CoA thioesters (Zeng et al. 2006). More recently, a fluorescence assay was developed for CpACBP1 and used to screen 1,040 known drugs, from which 28 drugs displayed inhibitory effects on CpACBP1 at sub-micromolar concentrations. Among them, four drugs (i.e., broxyquinoline, cloxyquin, cloxacillin sodium and sodium dehydrocholate) displayed efficacies against the growth of *C. parvum* with ID_{50} values at low micromolar levels. This observation raises hopes for potential repurposing of known drugs to treat cryptosporidiosis (Fritzler and Zhu 2012).

Two distinct oxysterol binding protein (OSBP)-related proteins (ORPs) have also been identified in *C. parvum*, designated as CpORP1 and CpORP2 (Zeng and Zhu 2006). The short-type CpOPR1 contains only a ligand binding domain, while the long-type CpORP2 contains Pleckstrin homology and ligand-binding domains. Lipid–protein overlay assays have revealed that CpORP1 and CpORP2 could specifically bind to phosphatidic acid, various phosphatidylinositol phosphates (PIPs), and sulfatide, but not to other types of lipids with simple heads. However, cholesterol was not a ligand for these two proteins. Like CpLCE1 and ACBP, CpORP1 also localized to the PVM, while CpORP2 localized only in intracellular merozoites (Zeng and Zhu 2006).

Due to the incapability of synthesizing fatty acids de novo, *Cryptosporidium* needs to scavenge fatty acids/lipids from the host. However, it is unclear how the parasite scavenges lipids as no specific fatty acid or lipid transporters have been identified or experimentally validated. A recent study has provided strong evidence that *C. parvum* is able to scavenge cholesterol from host cells and from the intestinal lumen (Ehrenman et al. 2013). Among lipoproteins, LDL is an important source of cholesterol, and *C. parvum* can obtain cholesterol that is incorporated into micelles and internalized into enterocytes by the NPC1L1 transporter. Pharmacological blockage of NPC1L1 function by ezetimibe or moderate down-regulation of NPC1L1 expression decreases parasite infectivity (Ehrenman et al. 2013).

The presence of acyl-CoA binding protein, long-chain acyl elongase, acyl-CoA synthetase and OSBP-related protein in PVM indicates that this unique membrane structure is involved in lipid metabolism including transport, activation and/or elongation of fatty acids in Cryptosporidium. Some ACS proteins in bacteria and yeast are also known to function as fatty acid transporters (Black and DiRusso 2007: DiRusso and Black 1999). Based on the most recent data, we have formulated a working hypothesis that fatty acids may be directly transported into the parasite via an undefined pathway(s) as free fatty acids, or by a PVM-specific ACS coupled with the formation of fatty acyl-CoA (Fig. 8.3). Free fatty acids may be elongated by the Type I FAS or PKS, or activated by ACS within the parasite to form fatty acyl-CoA. All fatty acyl-CoA thioesters may be immediately routed into subsequent metabolic pathways (such synthesis of complex lipids and biomembranes, or protein palmitoylation), undergo chain elongation by LCE, or bound to ACBP to form an acyl-CoA pool before entering subsequent pathways. Cryptosporidium may also scavenge phospholipids as implied by the presence of a type IV P-ATPase (cgd7_1760) with predicted substrate affinity towards phospholipids. However, it remains to be determined if this transporter is truly involved in lipid scavenging or is simply responsible for intracellular trafficking of phospholipids in parasite cells.

Cryptosporidium has a small set of enzymes involved in synthesis of complex lipids. In glycerolipid synthesis, it only retains enzymes for conversions between 1, 2-diacyl-sn-glycerol-3-phosphate, 1,2-diacyl-sn-glycerol and triacylglycerol by phosphatidate phosphatase, diacylglycerol kinase, and diacylglycerol acyltransferase 1, indicating that the parasite may store fatty acids in the form of triacylglycerol. Phosphatidyl-ethanolamine may be synthesized from diacyl-snglycerol and CDP-ethanolamine (produced from phospho-ethanolamine) by ethanolamine-phosphotransferase (ETHPT), or via the diacyl-sn-glycerol-3-phosphate to CDP-diacyl-glycerol to phosphatidyl-L-serine to phosphatidylentholamine pathway by CDP-diacylglycerol synthase (CDS), phosphatidylserine synthase and phosphatidylserine decarboxylase (PSDC). However, a phosphatidylserine synthase gene has not be identified in the *Cryptosporidium* genomes.

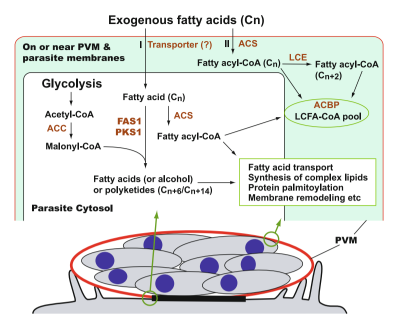


Fig. 8.3 Working hypothesis on the fatty acid metabolism associated with the parasitophorous vacuole membrane (PVM) in *Cryptosporidium*. Abbreviations: *ACBP* fatty acyl-CoA binding protein, *ACC* acetyl-CoA carboxylase, *ACS* Fatty acyl-CoA synthetase, *FAS1* Type I FAS, *LCE* long chain fatty acyl-elongase, *LCFA-CoA* long chain fatty acyl-CoA, *PKS1* Type I PKS

Additionally, enzymes involved in synthesizing lipoproteins and glycolipids are present in the *Cryptosporidium* genomes, which include up to nine DHHC family palmitoyl transferases for post-translational S-palmitoylation of proteins, and PIG-A, PIG-C, PIG-H, PIG-P, PIG-M and PIG-U involved in GPI anchor synthesis.

8.7 Stress-Related Pathways

Cryptosporidium must face various external and internal environmental stresses, including UV irradiation, temperature changes and dehydration in the natural environment, as well as hazardous chemicals, free radical molecules, drugs and host immune responses at various life cycle stages. The parasite genome encodes several classes of proteins that are important in handling these stresses, including heat shock proteins (HSPs) and various anti-oxidant molecules. HSPs are well known stress proteins that are generally up-regulated as part of the stress responses by participating in protein folding, maintaining proper protein conformation, and monitoring cellular proteins. The expression of the *CpHSP70* gene has been found to be highly up-regulated in response to chlorine-based oxidants and heat treatment (Bajszar and Dekonenko 2010).

Cryptosporidium possesses a number of anti-oxidant molecules, including superoxide dismutase (SOD), glutathione S-transferase, glutathione peroxidase, thioredoxin reductase, and a number of thioredoxin related proteins (e.g., Kang et al. 2008; Zhang et al. 2012; Yoon et al. 2012). In *C. parvum* oocysts, all three glutaredoxin-associated genes and five out of 13 thioredoxin-associated genes are expressed at various levels. Upon UV treatment, three putative t-complex protein 1 (TCP-1) subunits and several thioredoxin-associated genes are up-regulated in oocysts, whereas various HSP/DNAj family members, SOD and glutaredoxin-related genes are not up-regulated or are even down-regulated, suggesting that different stress proteins play different roles in responses to different stresses (Zhang et al. 2012).

DNA damage may occur more frequently in the natural environment than during DNA replication in the host cell due to the exposure of oocysts to UV irradiation. *Cryptosporidium* possesses a machinery for DNA excision repair. Several genes encoding excision repair enzymes were found to be up-regulated upon UV treatment, including the replication protein large subunit 1B (CpRPA1B) (Zhang et al. 2012; Rider and Zhu 2008; Rochelle et al. 2005). There are two types of RPA1 proteins in *Cryptosporidium* (i.e., RPA1A and RPA1B) which are involved in DNA replication, repair and recombination (Rider and Zhu 2008; Zhu et al. 1999; Millership and Zhu 2002). Several studies have indicated that RPA1A is mainly responsible for general DNA replication in the parasite, whereas RPA1B may play a role in DNA recombination and repair (Rider and Zhu 2008; Rider et al. 2005).

Trehalose synthesis is another important anti-stress pathway as described in Sect. 8.3 (Yu et al. 2010). Despite *Cryptosporidium* lacking amino acid synthetic pathways, a single standalone bacterial-type tryptophan synthase β -subunit gene is present in the *C. parvum* and *C. hominis* genomes. It is known that tryptophan starvation is one of the innate immunity's strategies in humans and animals to kill cells infected with certain pathogens including *Toxoplasma* and probably *Cryptosporidium* by activating the tryptophan degradation pathway (MacKenzie et al. 2007; Habara-Ohkubo et al. 1993). It is possible that *Cryptosporidium* may use tryptophan synthase β -subunit to synthesize tryptophan from indole present in the gut to evade tryptophan depletion.

8.8 Conclusions

Cryptosporidium is extremely well adapted to a parasitic life style. These parasites rely on the host to supply virtually all nutrients for their highly streamlined metabolic pathways. The insensitivity of *Cryptosporidium* to many anti-apicomplexan drugs is explained by the absence of the drug targets which are common in other apicomplexans. Examples of such pathways are de novo biosynthetic pathways and the cytochrome-based respiratory chain. Evolutionary divergence from other apicomplexans (e.g., bacterial type IMPDH and DHFR-TS with unusual amino acids at the active site) was also observed and may be relevant to

drug design. The availability of whole genome sequences provides opportunities not only to better understand the unique metabolic features of these parasites, but also to identify key enzymes and study their biochemical features of interest to drug development. Indeed, a number of potential drug targets have been proposed and/or are currently pursued by various laboratories, including protein kinases and enzymes in the carbohydrate, energy, nucleotide, and fatty acid metabolic pathways.

Currently, research on *Cryptosporidium* is still hampered by the lack of genetic tools and by technical difficulties in manipulating of the parasite in the laboratory. Therefore, recombinant protein-based biochemical analysis is still the best approach to study the metabolism and characterize potential drug targets. On the other hand, although gene knockout or knockdown tools are not available to validate drug targets, we can still effectively predict potential drug targets in the streamlined core metabolic pathways that are essential to the parasite. Drug development against cryptosporidial infection has been progressing slowly, but promising new data are being reported (see also Chap. 11). Research will be accelerated by new knowledge generated by biochemical analysis, target-based high-throughput screening of drugs, and structure-based analysis of protein-inhibitor interactions.

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Part III Host-parasite Interaction

Chapter 9 Human Cryptosporidiosis: A Clinical Perspective

Henry Shikani and Louis M. Weiss

Abstract Cryptosporidiosis is a diarrheal illness that in humans is most commonly due to infection by *Cryptosporidium parvum* or *C. hominis*. It has a world-wide distribution and is associated with both human to human and zoonotic infections, depending on the *Cryptosporidium* species causing the infection. Cryptosporidiosis can occur in both immune competent individuals, where it causes a self limiting diarrheal illness, or in patients with immune deficiency where the diarrhea can be extensive and result in death. The severity of illness is dependent upon factors such as age, environment, co-existent diseases, and host immune status. Infection typically involves the gastrointestinal cells that line the epithelial surface of the small and large intestines, but, depending on the species of *Cryptosporidium*, can also involve other epithelial surfaces such as the respiratory tract, particularly in immune compromised hosts. This chapter reviews the available data on the pathogenesis of cryptosporidiosis in humans, the host response to this infection, the clinical complications associated with illness, and the therapeutic and preventative strategies for management of this infection.

9.1 Introduction

Cryptosporidiosis is a common cause of diarrheal illness due to infection with species of the Apicomplexan protozoan parasite *Cryptosporidium*. Although *Cryptosporidium* was described in mice in 1907, it was not until 1976 that it was first

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reported to have an association with diarrhea in humans, in a healthy child and an adult with immune suppression. To date, about 20 Cryptosporidium species have been identified (White 2010). While Cryptosporidium parvum and Cryptosporid*ium hominis* are the most common causative agents of human disease, a number of other species can also infect humans including Cryptosporidium canis, C. meleagridis, C. felis. C. suis, C. baylei, C. muris, and C. andersoni (Cama et al. 2007, 2008; Gatei et al. 2006; Leoni et al. 2006; Muthusamy et al. 2006; Nichols 2008; Pedraza-Diaz et al. 2001; Fayer 2008; Tumwine et al. 2003) Infection characteristics and clinical features associated with cryptosporidiosis are shown in Tables 9.1 and 9.2 (Chalmers and Davies 2010; Guerrant 2008). These organisms usually cause diarrhea. Spontaneous recovery is the rule, and there is no universally effective specific therapeutic agent. Cryptosporidiosis is commonly identified as an opportunistic infection in immune compromised patients, such as those with acquired immunodeficiency syndrome (AIDS) (Bushen and Guerrant 2006; Chen et al. 2002; Kosek et al. 2001). It is also, however, a cause of diarrhea in immune competent individuals in both developed and developing countries, with a significant effect on public health (Bushen and Guerrant 2006; Chen et al. 2002; Kosek et al. 2001). Cryptosporidiosis-associated diarrheal syndromes can be stratified into three types, which are dependent on the specific population affected: (1) a selflimited diarrhea most often seen in immune competent individuals; (2) a persistent diarrhea which most commonly afflicts children in developing countries and can be associated with nutritional and growth effects in afflicted children; and (3) a chronic diarrhea syndrome most often seen in immune compromised individuals (Bushen and Guerrant 2006; Chen et al. 2002; Kosek et al. 2001). Many of the studies in the literature were done prior to the recognition of C. hominis as a distinct species from C. parvum and, therefore, whenever C. parvum is mentioned, in older literature, it is likely that these reports include cases due to both organisms. Cryptosporidium hominis is found primarily in humans and is responsible for most water borne outbreaks of this parasite, including the outbreak in 1993 in Milwaukee that involved 403,000 people (Chalmers and Davies 2010; MacKenzie et al. 1995).

9.2 Clinical Disease

Following *Cryptosporidium spp*. infection, humans are asymptomatic for typically 1 week during which the parasite multiplies in its target tissue, the epithelial lining of the intestine (Jokipii and Jokipii 1986; Mac Kenzie et al. 1994; Okhuysen et al. 1999, 2002). The incubation period between infection and presentation of symptoms can range, however, from 1 day to 1 month, depending on strain/species of parasite and the immune status of the host (Okhuysen et al. 1999, 2002; Pereira et al. 2002) (Table 9.1). Cryptosporidiosis typically presents as diarrhea, however, as mentioned above, the severity of disease and exact diarrheal syndrome can vary depending on the immune status and age of affected individuals (White 2010). Three distinct susceptible populations have been described: (1) immune competent

Table 9.1 Infection characteristics and clinical features associated with various Cryptosporidium isolates. Adapted from Chalmers and Davies (2010)	al features associated with various	Cryptosporidii	um isolates. Ad	apted from Cl	halmers and]	Davies (2010)
Study isolate	C. parvum IOWA ^a	C. parvum IOWA ^b	C. parvum TAMU ^b	C. parvum UCP ^b	C. parvum ^c C. hominis TU502	C. hominis TU502 ^d
Source of isolate	Calf ^a	Calf ^b	Foal ^b		Moredun Deer ^c	Human ^d
Infectious dose ID50	132ª (clinical definition with microbiological confirmation)	87 ^{b.e} ; 74.5 ^{b.f}	9 ^{b,e} ; 125 ^{b,f}	1,042 ^{b.e} ; 2788 ^{b,f}	300 ^{c,e} ; 375 ^{c,f}	10 ^{d.e.} ; 83 ^{d.f.}
Clinical attack rate (percent of subjects developing symptoms)	100 $\%^{a.e.}_{a.h.}$ 20 $\%^{a.i.}_{a.i.}$	52 % ^{b,g}	86 % ^{b.g}	59 % ^{b.g}	80 % cs	$\begin{array}{c} 62 \ \%^{\rm d.g.} \ 75 \ \%^{\rm d.j.} \\ 71 \ \%^{\rm d.k.} \ 60 \ \%^{\rm d.i.} \\ 40 \ \%^{\rm d.l} \end{array}$
Asymptomatic shedding of oocysts Prenatent neriod (mean davs)	Yes ^a 6 ^{a.g.} 9 ^{a.m.} 10 ^{a.n}	Yes ^b 7.7 ^{b,g}	Yes ^b 4 ^{b,g}	${ m Yes}^{ m b}$	No ^c 5 ^{c,g}	No ^d
Incubation period: mean; median (days)	$9^{a}, 6.5^{a}$	$9^{\rm b}; 7^{\rm b}$	$5^{\rm b}; 5^{\rm b}$	$11^{\rm b}; 6^{\rm b}$	Median: 3 ^c	5.4 ^d ; 4 ^d
Patent period: mean duration of shedding (days)	$12^{a,g}$; $10^{a,m}$; $2^{a,n}$	$8.4^{b,g}$	$3.4^{\mathrm{b,g}}$	$3.3^{b,g}$	5 ^{c,g}	
Duration of diarrhea: mean (hours)		64.2 ^b	94.5 ^b	81.6^{b}	Median:	1,373 ^d
Severity (mean number of unformed stools)	12.7 ^a	\mathcal{T}^{b}	9 ⁶	8 ^b	19°	
Relapse rate (%)		18 ^b	58 ^b	46^{b}		
^a DuPont et al. (1995) ^b Okhuysen et al. (1999) ^c Okhuysen et al. (2002) ^d Chappell et al. (2006) ^d Chappell et al. (2006) ^d Criadia = symptoms reported ^f Microbiological = oocysts detected in stools ^g Creater than 1,000 oocysts ^h Creater than 300 oocysts ¹ 30 oocysts ¹ 30 oocysts ¹ 10 oocysts ¹ 10 oocysts ¹ 10 oocysts ¹ 30 oocysts ¹ 30 oocysts ¹ 10 oocysts ¹ 30 oocysts						

Clinical features	<i>C. parvum</i> : 30–1,000,000 oocysts/dose (29 adults)	<i>C. hominis</i> : 10–500 oocysts/ dose (21 adults)
Infection rate ^a	18 (62 %)	9 (43 %)
Enteric symptoms rate ^b	11 (38 %)	14 (67 %)
Diarrhea attack rate ^c	7 (24 %) ^d	13 (62 %) ^e
Incubation period (mean days)	9	5.4
Duration of illness (mean hours)	74	137.3
Maximum number of unformed stools per day (mean)	6.4	3.2
Number of unformed stools per illness (mean)	12.7	8.9
Total stool weight per episode of diarrhea (mean, kg)	1.23	1.08

Table 9.2 Clinical features of healthy adults infected with *C. parvum* or *C. hominis* Oocysts. Adapted from Warren and Guerrant (2008)

^aInfection = excretion of oocysts in stools more than 36 h after ingestion of oocysts

^bRate of development of fever, nausea, vomiting, abdominal pain/cramps, tenesmus, gas-related intestinal symptoms, fecal urgency or fecal incontinence

^cDiarrhea in *C. parvum* study: passage of three unformed stools in 8 h or passage of more than three unformed stools in 24 h in addition to the presence of at least one enteric symptom; Diarrhea in *C. hominis* study: passage of ≥ 200 g of unformed stools per day or ≥ 3 unformed stools in 8 h or \geq unformed stools in 24 h

^dAll volunteers excreted oocysts

e6/13 volunteers excreted detectable oocysts

individuals in developed countries including tourists travelling in developing countries, (2) young children in developing countries whose immune status is not yet fully functional, and (3) immune compromised individuals (typically those with AIDS) (White 2010). Tables 9.3 and 9.4 describe the clinical characteristics of individuals in these patient populations (Chalmers and Davies 2010; Guerrant 2008).

9.2.1 Immune Competent Individuals

Infection of immune competent hosts in developed countries usually occurs because of water-related outbreaks, travel or animal/human contact (Hunter et al. 2004b; Roy et al. 2004). Children and the elderly are most frequently affected, and disease normally presents as a watery diarrhea that lasts for 1–2 weeks, but occasionally can persist for a month (Gambhir et al. 2003, Guerrant 2008; Mac Kenzie et al. 1994; Naumova et al. 2003; Neill et al. 1996). Oocysts are shed for a mean period of 7 days (range 1-15 days) after symptoms have ceased, although exceptionally for up to 2 months (Chalmers and Davies 2010; Jokipii and Jokipii 1986). Associated symptoms include abdominal cramps, nausea, vomiting and fever (Mac Kenzie et al. 1994). Recurrence of gastrointestinal symptoms can occur, e.g. in the Milwaukee outbreak this was reported by 30 % of patients

	Immune-competent:	Immune- compromised:	Immune- competent:
	Laboratory-diagnosed patients seeking medical care in United Kingdom	AIDS, haematological malignancy, pri- mary T-cell deficiency	Children in prospec- tive studies in developing countries
Length/ severity of illness	Self-limiting (mean: 13 days, median: 11 days) ^a	Severe/chronic/intrac- table; increased morbidity/mortality	Acute or chronic dis ease/carriage
Clinical	Gastrointestinal:	Gastrointestinal ^{c,d} :	Gastrointestinal ^e :
features	Diarrhea (98 %) ^b : watery (81 %) ^b ; loose (17 %) ^b ; bloody (11 %) ^a ; relapsing (40 %) ^a	Diarrhea: transient; relapsing; chronic; cholera-like	Watery diarrhea (96 %)
	Abdominal pain (60–96 %) ^{b,a}	Abdominal pain	Vomiting (57 %)
	Vomiting (49–65 %) ^{b,a}	Vomiting	Fever (37 %)
	Fever (36–59 %) ^{b,a}	Fever	Malnutrition ^{f,g,h,i}
	Nausea (35 %) ^b	Nausea	Failure to thrive ^{f,k}
		Severe weight loss	Weight loss ^f
		Biliary tract involvement ^d :	Impaired cognitive function ^j
		Cholangitis; pancreati- tis; sclerosing cholangitis;	
		Liver cirrhosis	
		Respiratory involvement ^d :	
		cough; sinusitis	

 Table 9.3
 Clinical characteristics of cryptosporidiosis in different patient groups. Adapted from Chalmers and Davies (2010)

^aHunter et al. (2004a, b) ^bPalmer and Biffin (1990) ^cManabe et al. (1998) ^dHunter and Nichols (2002) ^eKhan et al. (2004) ^fMolbak et al. (1997) ^gSallon et al. (1988) ^hSarabia-Arce et al. (1990) ^ILima et al. (1992) ^jGuerrant et al. (1997)

(Chalmers and Davies 2010; MacKenzie et al. 1995) and in an English study of sporadic cases by 40 % of patients (Chalmers and Davies 2010; Hunter et al. 2004a). The clinical presentation of infection may vary due to the infecting *Cryptosporidium* species. Oocysts are shed for longer and the number of oocysts detected in stools is higher in *C. hominis* infections than *C. parvum* (Bushen et al. 2007; Cama et al. 2008; Chalmers and Davies 2010; McLauchlin et al. 1999; Xiao et al. 2001). Cryptosporidiosis-associated disease in immune

Characteristics	Immune competent individuals	Immune compromised individuals	Other populations
Susceptible population	Healthy adults/ volunteers	AIDS patients, solid-organ transplant recipients, malignant individuals, individuals suffering from malnutrition, individuals undergoing hemodialysis, individuals with primary immunodeficiency diseases	Children (primarily those younger than 2 years of age), elderly adults
Site of infection Enteric presentation	Intestine Asymptomatic or acute	Intestine or extraintestine Asymptomatic, transient, chronic or fulminant	Intestine (primarily) Asymptomatic, acute or persistent
Signs and symptoms	Diarrhea, abdominal discomfort, fatigue	Diarrhea, weight loss, abdominal pain, nausea, vomiting, fever	Diarrhea, dehydration, mal- nutrition, weight loss
Duration of illness	4–14 days	14 days to several months	Outbreak: 5–17 days; Greater than 14 days to sev- eral weeks in children
Clinical outcome	Self-limiting	Increased morbidity and mortality	Long-term developmental impact on malnourished children; increased trans- mission and hospitaliza- tion in elderly

 Table 9.4
 Clinical features of human Cryptosporidiosis. Adapted from Warren and Guerrant (2008)

competent individuals is generally mild and can often be asymptomatic (White 2010), reflecting the importance of a functional immune system in controlling infection. This is reflected in reports of outbreaks and studies that have demonstrated that *C. parvum* infected immune competent hosts can have no symptoms (Chappell et al. 2006; Cicirello et al. 1997; DuPont et al. 1995; Mac Kenzie et al. 1994; Okhuysen et al. 1998, 1999, 2002; Pozio et al. 1997), but shed oocysts. Thus, as with many infections, the prevalence of individuals with symptomatic infection underestimates the extent of an infection in the community.

9.2.2 Young Children in Developing Countries

In developing countries, cryptosporidiosis-associated diarrhea most frequently occurs in children younger than 5 years (White 2010). In developing countries in Asia, Africa and Latin America, 5-10 % of diarrheal disease in children is due to infection with *Cryptosporidium spp*. (White 2010). Symptoms include watery diarrhea, abdominal cramps and, less frequently, fever and cough (White 2010).

Almost half of infected children develop a persistent diarrhea that lasts over 2 weeks (Newman et al. 1994; Sodemann et al. 1999), which increases the likelihood of additional episodes of diarrhea and death in these children (Agnew et al. 1998; Amadi et al. 2001; Lima et al. 2000; Molbak et al. 1993). Infection with Cryptosporidium is one of the major causes of persistent diarrhea in developing countries and can also manifest as chronic diarrhea and lead to malnutrition (Amadi et al. 2002; Behera et al. 2008). Clinical presentation varies dependent on Cryptosporidium spp., and perhaps the specific strain, with C. hominis being associated with the most severe disease syndromes in children and C. meleagridis with milder disease (Ajjampur et al. 2007; Bushen et al. 2007; Cama et al. 2008; Hunter et al. 2004a). Children in a birth cohort in Brazil with C. hominis infection had increased fecal lactoferrin and delayed growth compared with those with C. parvum (Bushen et al. 2007; Chalmers and Davies 2010). Children in Lima, Peru with C, hominis had diarrhea, nausea, vomiting and general malaise while only diarrhea was reported in those with C. parvum, C. meleagridis, C. canis or C. felis infection (Cama et al. 2008; Chalmers and Davies 2010). At a subtype level (GP60 gene subtyping) C. hominis subtypes Ia, Ib, Id and Ie were associated with diarrhea and Ib was also associated with nausea, vomiting and general malaise (Cama et al. 2008; Chalmers and Davies 2010).

Cryptosporidiosis is strongly associated with malnutrition in children in developing countries (Hunter and Nichols 2002). Infection causes malnutrition, and disease is significantly more severe, and can be fatal, in malnourished individuals (Amadi et al. 2001; Behera et al. 2008; Hunter and Nichols 2002; Javier Enriquez et al. 1997; Macfarlane and Horner-Bryce 1987; Newman et al. 1999; Sallon et al. 1988, 1994; Sarabia-Arce et al. 1990; Tumwine et al. 2003). A study conducted in the West Indies found that 15 out of 77 fecal specimens from malnourished children were positive for Cryptosporidium (Hunter and Nichols 2002; Macfarlane and Horner-Bryce 1987). Of the 15 cases, all experienced fever and diarrhea, most demonstrated dehydration and vomiting and two died due to the infection (Hunter and Nichols 2002; Macfarlane and Horner-Bryce 1987). A study in Israel reported that 30 out of 221 malnourished children with diarrhea also had Cryptosporidium infection (Hunter and Nichols 2002; Sallon et al. 1988). In a study performed in India, 50 children with diarrhea were examined (Hunter and Nichols 2002; Jaggi et al. 1994). Seven of those children were infected with Cryptosporidium, six of which were also malnourished (Hunter and Nichols 2002; Jaggi et al. 1994). A report from Gabon found the Cryptosporidium carriage rate in malnourished children (31.8 %) to be twice that of children with proper nourishment (16.8 %) (Duong et al. 1995; Hunter and Nichols 2002). In a Tanzanian study, 7 out of 55 children with acute diarrhea were malnourished and infected with Cryptosporidium (Cegielski et al. 1999; Hunter and Nichols 2002).

Malnourished children infected with *Cryptosporidium* before the age of one commonly demonstrate significant and unrecoverable weight loss and frequently experience stunted growth (Agnew et al. 1998; Checkley et al. 1998; Molbak et al. 1997). A study of infected infants in Guinea-Bissau found that, on average, boys with cryptosporidiosis lost 392 g and girls lost 294 g of body weight (Hunter

and Nichols 2002; Molbak et al. 1997). A study conducted in Bangkok, Thailand found that diarrhea was more common in children infected with *Cryptosporidium* compared to another intestinal parasite, *Giardia lamblia* (Hunter and Nichols 2002; Janoff et al. 1990). *Cryptosporidium*-infected children also lost more weight relative to those with *G. lamblia* (Hunter and Nichols 2002; Janoff et al. 1990). These reports are consistent with the idea that cryptosporidiosis is a precursor to malnutrition, however this cause-effect notion remains controversial (Hunter and Nichols 2002). The available studies also demonstrate that malnourished children infected with *Cryptosporidium* experience a more serious and potentially fatal disease characterized by a significant loss in weight (Hunter and Nichols 2002).

9.2.3 Immune Compromised Individuals

Cryptosporidiosis in immune compromised hosts often presents as a complication of HIV-1 infection, usually when the patient has clinically defined AIDS with a CD4+ T cell count under 200 cells/mm³ (White 2010). The disease variability seen in the presentation of cryptosporidiosis in various immune compromised patients is remarkable and currently unexplainable. Oddly, a large portion of HIV/Cryptosporidium co-infected individuals, even with CD4+ T cells under 200 cells/mm³, demonstrate mild or no symptoms, while others experience chronic watery diarrhea, significant weight loss and severe malabsorption (White 2010). This variability might occur because of co-infections with other pathogens that are either exacerbating or preventing symptoms of cryptosporidiosis (Hashmey et al. 1997; Lumadue et al. 1998). With the emergence of effective combination antiretroviral therapy (cART), the prevalence of cryptosporidiosis in the HIV-1 infected population in the areas with widespread use of cART has dramatically decreased (Kim et al. 1998; Le Moing et al. 1998). Therapy with cART helps maintain adequate levels of CD4⁺ T cells necessary to control the infection (Blanshard et al. 1992; Hashmey et al. 1997; Lumadue et al. 1998; Manabe et al. 1998).

Blanshard et al. examined 128 HIV positive patients with cryptosporidiosis and divided them into four clinical presentations: (1) asymptomatic infection, (2) transient infection, (3) chronic infection, and (4) fulminant infection (Blanshard et al. 1992; Farthing 2000). Infected, asymptomatic cases demonstrated normal bowel patterns, with less than three stools per day (Blanshard et al. 1992; Farthing 2000). This was the least common presentation, in this study, occurring in 3.9 % of patients (Blanshard et al. 1992; Farthing 2000). This study, however, did not specifically sample all HIV positive patients, but was based on samples submitted to a diagnostic laboratory, and thus likely underestimated the asymptomatic carriers of this organism. Transiently infected individuals represented 28.7 % of the patients and were defined as those who experienced diarrhea for under 2 months (Blanshard et al. 1992; Farthing 2000). At the 2-month time point, all symptoms ceased and the parasite could no longer be found in fecal samples (Blanshard et al. 1992; Farthing 2000). Chronic infection was the most common disease outcome (Blanshard

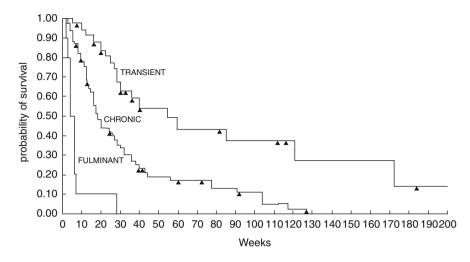


Fig. 9.1 Kaplan-Meier plots of the differential survivals of patients with transient, chronic and fulminant cryptosporidiosis (Reprinted with permission from Blanshard et al. (1992))

et al. 1992; Farthing 2000) comprising 59.6 % of these patients. In these chronically infected cases, diarrhea and Cryptosporidium spp. shedding in stools lasted over 2 months (Blanshard et al. 1992; Farthing 2000). Lastly, 7.8 % of patients experienced fulminant infection (Blanshard et al. 1992; Farthing 2000). These individuals experienced the most aggressive clinical outcome, passing at least two liters of watery diarrhea per day, i.e., a cholera-like diarrheal syndrome (Blanshard et al. 1992; Farthing 2000). Asymptomatic or transiently infected cases typically exhibited higher CD4⁺ T cell counts than the other two groups (Blanshard et al. 1992; Farthing 2000). All patients with fulminant infection had CD4⁺ T cell counts lower than 50 cells/mm³ (Blanshard et al. 1992; Farthing 2000), demonstrating the importance of T cells and the cellular immune response in controlling this disease. Survival (Fig. 9.1) and weight were also strongly associated with clinical presentation, as asymptomatic and transiently infected individuals survived much longer (median: 36 weeks) and exhibited a higher weight (median: 64.7 kg) than chronic (median: 20 weeks; 59.3 kg) and fulminant (median: 5 weeks; 55.2 kg) cases (Blanshard et al. 1992; Farthing 2000).

Chronic and fulminant patterns of disease are frequent outcomes in individuals in developing areas of the world (Farthing 2000). Sub-Saharan African AIDS patients co-infected with *Cryptosporidium* commonly demonstrate the chronic diarrhea wasting syndrome known as 'slim disease' (Farthing 2000). This illness has a major effect on morbidity and mortality, particular in areas with limited use of cART.

9.2.4 Additional Gastrointestinal Infections Due to Cryptosporidiosis

The severity of cryptosporidiosis is associated with the ability of the parasite to spread within the human gastrointestinal tract. The ability of the organism to infect various organs of the gastrointestinal tract varies by host and likely is highly dependent on the host immune status. When infection is controlled and asymptomatic, Cryptosporidium spp, infection is usually restricted to the colon (Clayton et al. 1994; Farthing 2000). In circumstances where disease becomes more severe, Cryptosporidium colonization occurs throughout the gastrointestinal tract, including such areas as the pharynx, esophagus, stomach, small bowel, small intestine and rectum (Berk et al. 1984; Farthing 2000; Godwin 1991; Kazlow et al. 1986; Kelly et al. 1998). Due to a defective immune response, it is well known that Cryptosporidium spp. infection in severely immune compromised individuals, such as AIDS patients, can involve the biliary system epithelium (Clavel et al. 1996; Farthing 2000; Hashmey et al. 1997; Lopez-Velez et al. 1995; Meynard et al. 1996; Vakil et al. 1996). Cryptosporidium spp. have also been reported in the duodenal papilla and the pancreatic duct, and infection has been associated with the production of overt pancreatitis (Farthing 2000; Gross et al. 1986). Additionally, ampullar infection can lead to severe constriction and obstruction of the bile duct, increasing susceptibility to cholangitis (Farthing 2000; Hasan et al. 1991). Gallbladder and biliary system infection is common in HIV-positive patients with cryptosporidiosis, occurring in 10-26 % of cases (Farthing 2000; Hasan et al. 1991; McGowan et al. 1993: Vakil et al. 1996).

AIDS-related sclerosing cholangitis is a serious clinical complication that manifests in patients with AIDS and is strongly associated with C. parvum intestinal infection (Farthing 2000). Symptoms include abdominal pain, fever and jaundice. Patients with this syndrome demonstrate increased serum alkaline phosphatase and significant anatomical damage to the biliary system on various imaging studies (Farthing 2000). Cryptosporidiosis patients with biliary disease have lower CD4⁺ T cell counts and a significantly higher rate of mortality than individuals without biliary symptoms (Farthing 2000; Vakil et al. 1996). Infection of the biliary tract with C. parvum has been associated with rapid onset of death (Hashmey et al. 1997; Vakil et al. 1996). Among 24 patients with AIDS and biliary involvement and symptoms from the large waterborne outbreak of cryptosporidiosis in Milwaukee, it was reported that biliary symptoms was a strong indicator of the prognosis since 83 % of patients with symptoms died within the following year compared to only 48 % of those without these symptoms (Hunter and Nichols 2002; Vakil et al. 1996). This may have reflected the advanced state AIDS in patients with biliary tract involvement. Lastly, widespread squamous metaplasia of the epithelial lining of the bile duct has been reported in the context of chronic Cryptosporidium infection (Farthing 2000; Kline et al. 1993).

9.2.5 Extraintestinal Infection

Cryptosporidiosis is known to spread as far as the respiratory tract (Farthing 2000). In addition, this may be more common with other species of *Cryptosporidium*, e.g. Cryptosporidium meleagridis is known to be associated with respiratory disease in birds, Cryptosporidium spp. have been found in humans in sputum, in bronchial aspirates, and in the bronchial epithelium (Blanshard et al. 1992; Clavel et al. 1996; Farthing 2000; Kocoshis et al. 1984; Travis et al. 1990). Symptoms of respiratory infection include cough, dyspnea, fever and thoracic pain with pneumonia being described as the most severe clinical outcome (Farthing 2000). Respiratory disease frequently results from co-infection with Cryptosporidium spp. and other pathogens, namely Mycobacterium tuberculosis or Pneumocystis carinii, however single infections with this parasite have also been observed (Clavel et al. 1996; Farthing 2000). Parasitic sinusitis has also been found in individuals co-infected with HIV and Cryptosporidium spp. (Dunand et al. 1997; Hunter and Nichols 2002). These cases typically presented fever, chills, and local tenderness and discharge (Dunand et al. 1997; Hunter and Nichols 2002). Diseases such as reactive arthritis and Reiter's syndrome have also been seen in patients with cryptosporidiosis.

9.3 Additional Complications of Cryptosporidiosis

9.3.1 Pneumatosis Cystoides Intestinales

A number of case reports on AIDS patients with cryptosporidiosis have described additional consequences of infection and in particular, pneumatosis cystoides intestinales has been described (Collins et al. 1992; Hunter and Nichols 2002; Samson and Brown 1996; Sidhu et al. 1994). Patients with pneumatosis cystoides intestinales are defined by the presence of rupture-prone, gas-containing cysts in their intestinal wall that are associated with the production of pneumoretroperitonea (air in the retroperitoneum) and pneumomediastina (air in the mediastinum) (Hunter and Nichols 2002).

9.3.2 Esophageal Damage and Appendicitis

A case was reported of a 2-year-old child with cryptosporidiosis who experienced damage to the esophagus, causing vomiting and dysphagia (Hunter and Nichols 2002; Kazlow et al. 1986). Cryptosporidiosis has also been associated with appendicitis (Hunter and Nichols 2002; Oberhuber et al. 1991). In a review of the

literature, 16 cases of gastric cryptosporidiosis were identified (Hunter and Nichols 2002; Ventura et al. 1997). In an endoscopic study of 24 patients with AIDS and chronic diarrhea due to cryptosporidiosis, 16 (67 %) had parasites in the gastric epithelium (Hunter and Nichols 2002; Rossi et al. 1998). The majority of these patients had no specific symptoms related to this gastric colonization. A significant complication of gastric involvement by *Cryptosporidium spp*. is antral narrowing and gastric outlet obstruction (Cersosimo et al. 1992; Garone et al. 1986; Hunter and Nichols 2002; Iribarren et al. 1997; Moon et al. 1999). This obstruction causes nausea, vomiting and a reduction in nutrient intake.

9.4 Risk Factors for Cryptosporidiosis

Susceptibility to cryptosporidiosis depends on several factors, including environmental conditions, host immune status, age, geographic location and contact with infected humans/animals (Table 9.5). The prevalence of cryptosporidiosis ranges from as low as 1 % in more westernized regions of the world, such as North America and Europe, to as high as 30 % in tropical areas, especially in developing countries (Farthing 2000). A tropical climate is ideal for survival of the *Cryptosporidium* oocyst. The average prevalence of diarrheal cases due to infection with the parasite is almost threefold higher in developing (6.1 %) versus developed (2.1 %) countries (Farthing 2000; Adal and Guerrant 1995).

Cryptosporidiosis is a very common cause of waterborne disease and has been associated with drinking water and swimming pool contact. Cryptosporidiosis is responsible for one of the largest waterborne outbreaks ever described (Hunter and Nichols 2002; Mac Kenzie et al. 1994). The infectious dose of Cryptosporidium in human hosts is thought to be low (Blagburn and Current 1983; Guerrant 2008). In one study involving infection of immune competent volunteers with the Iowa isolate of C. parvum (Abrahamsen et al. 2004), a 50 % infectious dose (ID₅₀) of 132 oocysts was estimated (DuPont et al. 1995; Guerrant 2008). There is significant unpredictability in infectivity, however, among different strains of *C. parvum*, as demonstrated by distinct ID_{50S} of individual isolates in various host species (Guerrant 2008; Okhuysen et al. 1999). In a study where immune competent volunteers were infected with C. hominis, the ID_{50} ranged from 10 to 83 oocysts (Chappell et al. 2006; Guerrant 2008). Susceptibility to re-infection and illness following primary infection was investigated by re-challenging volunteers (DuPont et al. 1995) with the homologous isolate 1 year later (Okhuysen et al. 1999). This study demonstrated that infection still occurred and that the onset and duration was similar to the primary infection, although the degree of diarrhea and number of oocysts shed was decreased.

	Immune competent hosts	Immune compromised hosts
Developed countries	Age (young most susceptible)	Recreational water (i.e. swimming pools, lakes, rivers, beaches)
	Recreational water (i.e. swimming pools, lakes, rivers, beaches)	Poorly sanitized sources of water
	Poorly sanitized sources of water	Primary immunodeficiency diseases (e g. common variable immunodeficiency)
	Travel	Agents that cause immunodeficiency (HIV/AIDS)
	Hospitals	Transplantation
	Animal reservoirs	Immune suppressant medications
Developing countries	Age (young most susceptible)	Agents that cause immunodeficiency
	Lack of proper breast-feeding	(HIV/AIDS)
	Poorly sanitized sources of water	Poorly sanitized sources of water
	Rain	Rain
	Animal reservoirs	Animal reservoirs

Table 9.5 Risk factors for human Cryptosporidiosis

9.4.1 Immunodeficiency Diseases

As noted previously, patients with immune deficiency, particularly those with AIDS, are prone to developing severe cryptosporidiosis. Several other immune defects have been associated with cryptosporidiosis. Another immunodeficiency complication associated with cryptosporidiosis is severe combined immunodeficiency syndrome (Hunter and Nichols 2002). One case report described how, despite receiving both extensive therapy as well as a thymus transplantation, an infected individual with this syndrome died 5 months after being diagnosed with cryptosporidiosis (Hunter and Nichols 2002; Kocoshis et al. 1984). Furthermore, this patient exhibited a widespread infection with extraintestinal spread as Cryptosporidium parasites were identified in the small intestine, pancreatic duct and bronchioles (Hunter and Nichols 2002; Kocoshis et al. 1984). In another case of cryptosporidiosis, deficiencies in selective immunoglobulin A and Saccharomyces opsonin were discovered (Hunter and Nichols 2002; Jacyna et al. 1990). Several patients with X-linked hyper-immunoglobulin M syndrome have simultaneously demonstrated chronic diarrhea and liver damage due to cryptosporidiosis (Hayward et al. 1997; Hunter and Nichols 2002; Levy et al. 1997). This is a highly fatal immunodeficiency syndrome, characterized by a mutation in the gene for CD40 ligand and significant impairment of the T lymphocytes (Hunter and Nichols 2002). Mouse models have suggested that the CD40/CD40 ligand system is critical in the immune response to this pathogen. Lastly, Gomez Morales et al. described a child with chronic cryptosporidosis and diarrhea with a marked loss in weight who demonstrated deficiencies in interferon- γ production (IFN- γ) (Gomez Morales et al. 1996; Hunter and Nichols 2002).

9.4.2 Cancer

Several studies have looked at associations of cancer and cryptosporidiosis, however, a consistent association has not been demonstrated. A study performed in Turkey on 106 fecal specimens from patients with diarrhea and cancer found that 17.0 % were positive for parasite oocysts (Hunter and Nichols 2002; Tanyuksel et al. 1995). In a similar study conducted in India, only 1.3 % of diarrhea/cancer cases had *Cryptosporidium* in their stools (Hunter and Nichols 2002; Sreedharan et al. 1996), suggesting a possible country-specific susceptibility to infection in this patient population. In two separate studies performed on children with cancer and diarrhea (one in New South Wales and the other in Malaysia), 0 % and 2 % of patients, respectively, were positive for *Cryptosporidium* (Burgner et al. 1999; Hunter and Nichols 2002; Menon et al. 1999), further suggesting a possible country-specific susceptibility.

9.4.3 Leukemia/Lymphoma

The relationship of cryptosporidiosis and leukemia has not been definitely determined. In one study, a child with acute lymphocytic leukemia and severe diarrhea recovered (Hunter and Nichols 2002; Stine et al. 1985), while, in a different study, an infected child with acute lymphoblastic leukemia had a relapse (Hunter and Nichols 2002; Lewis et al. 1985). A study on six children with leukemia or lymphoma found that four cases survived with chemotherapy while persistently infected patients died (Foot et al. 1990; Hunter and Nichols 2002).

9.4.4 Bone Marrow and Solid-Organ Transplantation

Severe cryptosporidiosis with diarrhea can occur in patients undergoing bone marrow transplantation (Gentile et al. 1991; Hunter and Nichols 2002; Manivel et al. 1985). Pulmonary cryptosporidiosis, sometimes fatal, has also been described in bone marrow transplant cases (Hunter and Nichols 2002; Kibbler et al. 1987; Nachbaur et al. 1997). Additionally, one bone marrow transplantation facility experienced an outbreak of cryptosporidiosis (Hunter and Nichols 2002; Martino et al. 1988). Cryptosporidiosis has been observed in children undergoing liver transplantations (Campos et al. 2000; Gerber et al. 2000; Hunter and Nichols 2002; Vajro et al. 1991). A Belgian study found that three out of 461 children experienced a diffuse *Cryptosporidium*-associated cholangitis post-transplant (Campos et al. 2000; Hunter and Nichols 2002). In a Pittsburgh study, it was discovered that, of 1,160 patients receiving abdominal organ transplants, four children had cryptosporidiosis and resolved their disease (Gerber et al. 2000;

Hunter and Nichols 2002). Three of the four children were undergoing liver transplants while the fourth was a small bowel transplant patient (Gerber et al. 2000; Hunter and Nichols 2002). The rate of cryptosporidiosis in patients undergoing renal transplantation has also been explored in multiple studies (Hunter and Nichols 2002). In two separate studies conducted in Brazil and Turkey, renal transplantation was associated with an increase in infection with *Cryptosporidium spp*. (Chieffi et al. 1998; Hunter and Nichols 2002; Ok et al. 1997).

9.4.5 Diabetes

An association between cryptosporidiosis and diabetes has been proposed. In separate reports, three diabetes patients also experienced chronic diarrhea due to *Cryptosporidium* infection (Chan et al. 1989; Hunter and Nichols 2002; Trevino-Perez et al. 1995). The significance of this observation is not clear as the majority of individuals with diabetes do not demonstrate simultaneous evidence of cryptosporidiosis and in large studies of cryptosporidiosis, diabetes has not emerged as a risk factor (Hunter and Nichols 2002).

9.5 Host Immune Response

9.5.1 CD4⁺ and CD8⁺ T Cells

As with all diseases, an effective host response to infection with *Cryptosporidium* is critical to controlling the pathology and clinical outcome associated with the parasite. Arguably, the most important cell type in this response are $CD4^+$ T cells, as evidenced by the fact that Cryptosporidium presents as one of the main opportunistic infections in patients with low cell counts due to HIV (Blanshard et al. 1992; Flanigan et al. 1992; Hashmey et al. 1997; Pantenburg et al. 2008). The importance of this T cell subtype has also been seen in in vivo mouse studies. Mice that lack functional CD4⁺ T cells demonstrate chronic Cryptosporidium infection. However, mice given an infusion of these cells demonstrate significant improvement (Aguirre et al. 1994; Chen et al. 1993; Culshaw et al. 1997; McDonald et al. 1992, 1994; Perryman et al. 1994; Ungar et al. 1991). The role of the effector CD8⁺ T cell subtype in controlling *Cryptosporidium* infection is not as clear (White 2010). While these cells localize to sites of infection (Abrahamsen 1998; Wyatt et al. 1997) and are known to produce IFN- γ , a crucial cytokine for combating infection, it is not known whether a lack of CD8⁺ T cells exacerbates disease (Kirkpatrick et al. 2008; Pantenburg et al. 2008; Preidis et al. 2007). Furthermore, CD8⁺ T cells do not appear to be important for preventing cryptosporidiosis in mice

(Aguirre et al. 1994; McDonald et al. 1994), making it more challenging to characterize their role during infection.

9.5.2 IFN-γ

The importance of IFN- γ in controlling cryptosporidiosis is well established. Mice that lack this pro-inflammatory cytokine develop a persistent infection described as worse than in mice without CD4⁺ T cells (Mead and You 1998; Theodos et al. 1997; Tzipori et al. 1995). Moreover, studies in mice also demonstrate that interleukin (IL)-12, a cytokine which contributes to the production of IFN- γ , is also critical for regulating Cryptosporidium infection (Campbell et al. 2002; Urban et al. 1996). As has been shown in vitro with C. parvum, IFN- γ may help clear the parasite by activating intestinal epithelial cells (Pollok et al. 2001). While evidence points strongly towards IFN-y playing a key role in controlling Cryptosporidium infection, this cytokine does not always help regulate the disease. Infected individuals with low levels of IFN-y have been shown to control this infection (Gomez Morales et al. 1999; Kirkpatrick et al. 2002). Additionally, murine studies demonstrate that functional IFN- γ is critical to the survival of C57BL/6, but not BALB/c mice, suggesting that other immune factors must also be working to reverse or prevent disease (Gomez Morales et al. 1999; Kirkpatrick et al. 2002; Mead and You 1998). A number of other cytokines have been proposed to function in limiting infection, including the pro-inflammatory cytokine, tumor necrosis factor-alpha (TNF- α), which has been shown to clear the parasite in vivo and in vitro (Lacroix et al. 2001; Pollok et al. 2001; Wyatt et al. 1997).

9.5.3 Antibody

While the role of cellular immunity, specifically $CD4^+$ T cells, in responding to *Cryptosporidium* infection is clear, the function of the humoral arm of the immune response remains unknown (Pantenburg et al. 2008). Human studies have demonstrated that individuals infected with *C. parvum* exhibit high antibody levels (Benhamou et al. 1995; Cozon et al. 1994; Dann et al. 2000), but the reasons for this are not clear. Murine studies have shown that the disease is not more severe in infected mice with non-functional B cells, however, treatment of infected mice with antibody to *Cryptosporidium* has been shown to facilitate parasite clearance (Arrowood et al. 1989; Chen et al. 2003; Jenkins et al. 1999; Perryman et al. 1999; Riggs 2002; Sagodira et al. 1999; Taghi-Kilani et al. 1990).

9.6 Diagnosis

Cryptosporidium infection is typically diagnosed by stool examination (White 2010). A number of approaches have been attempted to increase diagnostic sensitivity, i.e. concentration methods involving formalin buffer, flotation methods and the use of immunomagnetic beads aimed at isolating the organism (Pereira et al. 2002; Smith 2008; Webster et al. 1996). Various staining methods have also been tested for diagnosis, including acid-fast staining, fluorescence and immunofluorescence with antibodies that target the oocyst (White 2010). Because of its high sensitivity compared to other assays, immunofluorescence is now considered the gold standard for detecting *Cryptosporidium* in stool samples (Alles et al. 1995; Balatbat et al. 1996; Dagan et al. 1995; Garcia and Shimizu 1997; Ignatius et al. 1997; Johnston et al. 2003; Kehl et al. 1995). Other approaches that focus on detecting Cryptosporidium-specific antigens, such as enzyme-linked immunosorbent assay (ELISA) and immunochromatography, have been generally described as similar in sensitivity to other techniques, but more specific (Cryptosporidiosis 2009; Dagan et al. 1995; Garcia and Shimizu 1997, 2000; Garcia et al. 2003; Johnston et al. 2003; Smith 2008). Lastly, PCR tests for detection of Cryptosporidium DNA have been used and are considered more sensitive than stool examination (Ajjampur et al. 2008; Amar et al. 2007; Kaushik et al. 2008; Nair et al. 2008; ten Hove et al. 2007). Identification of *Cryptosporidium* species, in clinical specimens, is most commonly done using molecular techniques, such as PCR.

9.7 Pathology

Cryptosporidium invasion of host cells is restricted to the luminal border of the epithelial lining of the gastrointestinal tract (Fig. 9.2). Severe infection leads to displacement of the microvillous border and loss of the surface epithelium, causing changes in the villous architecture, with villous atrophy, blunting and crypt cell hyperplasia, and mononuclear cell infiltration in the lamina propria. The cause of diarrhea has yet to be elucidated but osmotic, inflammatory and secretory processes have been suggested. It is likely, that pathogenesis is due to several factors including the effect of the parasite and the substances it produces on the epithelial layer and the immunological and inflammatory responses of the host, leading to impaired intestinal absorption and enhanced secretion (Farthing 2000).

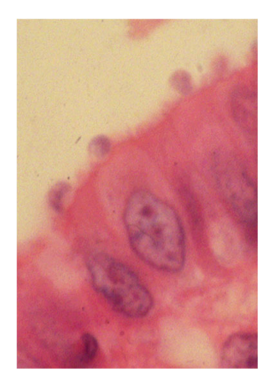


Fig. 9.2 Intestinal biopsy demonstrating *Cryptosporidium parvum* at the surface of the intestinal epithelial cells. Hematoxylin and eosin stain, 100X oil objective

9.8 Pathogenesis

The extent of pathology associated with *Cryptosporidium* infection largely depends on the individual's immune status. These parasites typically infect gastrointestinal epithelial cells. In immune competent cases, organisms normally concentrate in the distal small intestines and proximal colon, however immunodeficiency can cause the parasite to spread to other areas in the body, such as the gut and respiratory tract (Godwin 1991; Greenberg et al. 1996; Kelly et al. 1998). Additionally, the infectious dose can also contribute to pathogenesis as a large dose of oocysts has the potential to lead to a disseminated infection, causing a dysregulated involvement of such immune cells as lymphocytes and neutrophils (Clayton et al. 1994; Genta et al. 1993; Goodgame et al. 1995; Greenberg et al. 1996; Lumadue et al. 1998).

A balanced immune response is critical to controlling any pathogenic infection, including that of *Cryptosporidium*. While the importance of pro-inflammatory cytokines such as IFN- γ in regulating cryptosporidiosis is well documented, their dysregulated activity may also contribute to some of the characteristic features of disease, including increased permeability in the intestines (Roche et al. 2000). Antiinflammatory cytokines such as transforming growth factor beta (TGF- β) have been shown in vitro to prevent *C. parvum*-mediated epithelial barrier breakdown (Roche et al. 2000), demonstrating the necessity of a regulated immune response.

9 Human Cryptosporidiosis: A Clinical Perspective

The major clinical feature associated with cryptosporidiosis is diarrhea (White 2010). Physiologically, diarrhea is believed to manifest because of several factors, including the malabsorption of sodium, the secretion of chloride and the increased permeability of the intestinal epithelial layer (White 2010). These consequences likely result, in part, from the cascade of nuclear factor kappa-light-chain-enhancer of activated B cells (NF- κ B)-mediated events that occur intracellularly following *Cryptosporidium* infection of epithelial cells in the intestine. Infection induces activation of the multi-functional transcription factor NF- κ B (Chen et al. 2001; McCole et al. 2000). NF- κ B subsequently activates a variety of anti-apoptotic factors as well as pro-inflammatory cytokines and chemokines, including TNF- α , IL-1 β and IL-8 (Alcantara et al. 2003; Kandil et al. 1994; Kirkpatrick et al. 2002, 2006; Lacroix et al. 2001; Pantenburg et al. 2008; Robinson et al. 2001; Seydel et al. 1998; Wyatt et al. 1997). Inflammation causes a dysregulated influx of immune and inflammatory cells to infection sites, which probably contributes to enhanced epithelial permeability as well as other aspects of disease (White 2010).

Other factors which have been implicated in the pathogenesis of cryptosporidiosis include the TNF- α -mediated increased production of prostaglandins (Kandil et al. 1994). This increased production is believed to contribute to sodium malabsorption and chloride secretion in porcine and bovine cryposporidiosis (Argenzio et al. 1996; Cole et al. 2003; Gookin et al. 2004; Kandil et al. 1994; Laurent et al. 1998). Evidence thus far suggests that prostaglandins do not play a role, however, in human cryptosporidiosis (Okhuysen et al. 2001; Robinson et al. 2001; Sharpstone et al. 1996; Snijders et al. 1995).

The specific neuropeptide substance P has also been associated with cryptosporidiosis (Garza et al. 2008; Hernandez et al. 2007; Robinson et al. 2003, 2008). It has been shown that levels of this neuropeptide correlate with severity of disease both in infected immune competent and immune compromised individuals (Robinson et al. 2003). Both in non-human primate and murine models of infection, it was demonstrated that therapeutically targeting substance P itself or its receptor significantly ameliorates disease (Garza et al. 2008; Hernandez et al. 2007; Robinson et al. 2008; Sonea et al. 2002). Substance P is thought to contribute to both chloride secretion and enhanced intestinal permeability (Garza et al. 2008; Hernandez et al. 2007).

Cell death in the intestine, both by apoptosis and necrosis, is a predominant feature of cryptosporidiosis. While infected epithelial cells are programmed initially to survive, uninfected adjacent cells undergo apoptosis (Chen et al. 1999, 2001; McCole et al. 2000; Motta et al. 2002). Interactions between FAS ligand and its receptor are thought to facilitate cell death (Chen et al. 1999, 2001; McCole et al. 2000; Motta et al. 2002). Moreover, once *Cryptosporidium* organisms complete their intracellular cycle, they induce necrosis of infected cells (Chen et al. 1998; Elliott and Clark 2003; Griffiths et al. 1994). All of these events together likely contribute to the decreased intestinal surface area, increased permeability and malabsorption that occur during disease (White 2010).

9.8.1 Malabsorption

Disease caused by *Cryptosporidium* typically increases in severity when the organism spreads to the proximal small intestine (Clayton et al. 1994; Farthing 2000). Infection of both humans and animals results in significant anatomical disturbances to the crypt and villous architecture of the small intestine, with complete atrophy and hyperplasia occurring in certain circumstances (Argenzio et al. 1990; Clayton et al. 1994; Farthing et al. 1996, Farthing 2000; Genta et al. 1993; Phillips et al. 1992). These alterations are associated with an uncontrolled influx of inflammatory cells to the lamina propria and epithelium, leading to both a decrease in surface area and a decline in the enzymatic activity of disaccharidases and aminopeptidases in the microvillous membrane (Farthing 2000). The reduced activity of the above enzymes is likely related, in part, to the damage to the microvilli which takes place during infection (Bird and Smith 1980; Farthing 2000). This microvillous damage is most severe at points where Cryptosporidium organisms are attaching to epithelial cells (Bird and Smith 1980; Farthing 2000). Furthermore, it has been demonstrated that intracellular organelles undergo significant structural change during infection, e.g. swelling of mitochondria. These alterations in structure correlate with an impairment in microvillous function, including a reduction in both the absorption of monosaccharides and the co-transport of sodium and glucose (Argenzio et al. 1990; Farthing 2000; Modigliani et al. 1985). In cases of more severe disease, i.e. AIDS patients with cryptosporidiosis, poor absorption of vitamin B_{12} and fat has also been described (Farthing 2000; Goodgame et al. 1995; Modigliani et al. 1985).

9.8.2 Dysregulated Intestinal Secretion

Characterized by aggressive bouts of watery diarrhea, the clinical presentation of fulminant cryptosporidiosis has been compared to that caused by enterotoxinproducing organisms (i.e., cholera) (Farthing 2000). This comparison evokes the possibility that cryptosporidiosis is a diarrheal disease which results from both defective absorption and increased secretion of important electrolytes (Farthing 2000). Studies on human stool samples attempting to confirm this "secretory" characterization have thus far been controversial, however, enterotoxin-like activity has been found in fecal specimens from *C. parvum*-infected calves (Farthing 2000; Guarino et al. 1994, 1995, 1997; Kelly et al. 1996). It was also discovered that individuals with cryptosporidiosis have increased intestinal levels of the potent secretagogue, hydroxytryptamine (5-HT) (Farthing 2000; Sears and Guerrant 1994). The secretory diarrhea which results from infection with *Cryptosporidium* may therefore result from a combination of enterotoxin-like activity and 5-HT (Farthing 2000).

9.9 Clinical Management of Cryptosporidiosis

Effective management of *Cryptosporidium* infection can minimize the severity of this diarrheal disease (White 2010). Diarrhea can result in extreme dehydration, therefore management begins with replacing lost fluids and electrolytes, whether through oral rehydration or intravenous approaches. Rehydration and repletion of electrolyte losses by either the oral or intravenous (IV) route are important. Severe diarrhea can exceed 10 L/day among patients with AIDS, often requiring intensive support. Oral rehydration should be pursued aggressively with oral rehydration solutions. Patients should be monitored closely for signs and symptoms of volume depletion, electrolyte and weight loss, and malnutrition. Total parenteral nutrition might be indicated in certain patients. Unfortunately, cryptosporidiosis causes a loss in enzymes such as lactase, therefore rehydration regimens should be lactosefree (Phillips et al. 1992). In addition, in the recovery stage following infection, lactase deficiency can persist, contributing to diarrhea. Therefore, after treatment it may be necessary to restrict milk products and/or use lactase until the brush border is restored. To date, there are limited options for the direct treatment of infection, as a large number of antiparasitic agents have no effect especially in immune compromised hosts. The cornerstone of treatment for cryptosporidiosis is, therefore, the restoration of host immune function, e.g. cART in patients with AIDS. Case reports had suggested that oral therapy with anti-Cryptosporidium immunoglobulin preparations (bovine colostrum) was effective; however, a clinical trial of 20 g/day for treatment did not demonstrate any benefit (Fries et al. 1994). Patients with biliary tract involvement may require endoscopic retrograde choledocoduodenoscopy for diagnosis and may benefit from sphincterotomy and/or stenting (Hashmey et al. 1997).

9.9.1 Restoration of the Immune System

Cryptosporidiosis is a disease which frequently presents in immune compromised individuals, such as patients with AIDS. Any therapeutic agent, therefore, that either restores a functional immune system or targets the pathogen responsible for damaging the immune response could ameliorate clinical outcome (White 2010). In AIDS patients with cryptosporidiosis, antiretroviral therapy (cART) has been shown to significantly improve diarrheal disease (Carr et al. 1998; Foudraine et al. 1998; Grube et al. 1997; Maggi et al. 2000; Miao et al. 2000; Okhuysen et al. 2001). Interestingly, the antiretroviral medications that function by inhibiting HIV proteases have demonstrated specific activity against *Cryptosporidium* species both in vivo and in vitro (Hommer et al. 2003; Mele et al. 2003). Immune reconstitution inflammatory syndrome (IRIS) has not been described in association with treatment of cryptosporidiosis.

9.9.2 Antimotility Agents

The intestinal epithelial cell layer breakdown which occurs during cryptosporidiosis results in an increase in intestinal transit (Brantley et al. 2003; Sharpstone et al. 1999) that contributes to malabsorption (Brantley et al. 2003; Sharpstone et al. 1999). Therapeutics that regulate motility in the intestine are, therefore, critical for regulating disease severity (White 2010). Opiates are typically the primary antimotility drugs used for treatment (White 2010). Tincture of opium may be more effective than loperamide. Octreotide in case reports was also effective, however, a clinical trial did not demonstrate benefit for severe diarrhea due to cryptosporidiosis (Cello et al. 1991).

9.9.3 Antiparasitic Agents

There have been a large number of studies aimed at developing a satisfactory therapy for cryptosporidiosis, particularly in patients with AIDS. In vitro screening of over 200 agents failed to identify a highly active compound (White 2010). Recently halofuginone lactate (HalocurTM) was identified and developed as a commercial veterinary drug for the prevention and treatment of this disease in cattle. To date, there have been no antiparasitic agents found which have clinical activity for the treatment of cryptosporidiosis in immune compromised patients. Multiple agents have been investigated in small randomized controlled clinical trials of HIV-infected adults, including nitazoxanide, paromomycin, spiramycin, bovine hyperimmune colostrum, and bovine dialyzable leukocyte extract. No pharmacologic or immunologic therapy directed specifically against Cryptosporidium has been shown to be consistently effective when used without cART (Cabada and White 2010). Agents which have variable activity, but are used clinically, include macrolides, e.g. clarithromycin and azithromycin, and the aminoglycoside paromomycin (Griffiths 1998; Hashmey et al. 1997; Hunter and Nichols 2002; Tzipori 1998). Nitazoxanide has been approved by the Food and Drug Administration (FDA) for treatment of C. parvum in immune competent children and adults. Because of the clinical significance of cryptosporidiosis, a trial of nitazoxanide or other anti-parasitic drugs in conjunction with cART, but never instead of cART, may be considered in patients with AIDS. Combination therapy with multiple antiparasitic agents has not been studied for the treatment of cryptosporidiosis and merits further investigation.

9.9.4 Nitazoxanide

To date, the only licensed treatment for cryptosporidiosis is the thiazole-containing therapeutic, nitazoxanide with in vivo activity against a broad range of helminths, bacteria, and protozoa (Shirley et al. 2012). It has been approved by the Food and Drug Administration (FDA) for treatment of C. parvum in children and adults (Chap. 11). Nitazoxanide has demonstrated efficacy in reducing the severity and duration of disease in immune competent individuals infected with the parasite (Shirley et al. 2012). Regardless of dosage and length of treatment, nitazoxanide has not proven to be highly effective in immune compromised patients, specifically those with AIDS (Amadi et al. 2009; Shirley et al. 2012). In patients with AIDS and cryptosporidiosis it has been administered at 500 to 1,000 mg twice daily in adults with some evidence of response when used on a compassionate expanded use program in patients with advanced AIDS (Rossignol 2006, Rossignol et al. 1998, 2001). Response depended on immune status as 70 % of patients with CD4+ counts > 50 cells/mm³ responded compared to 20 % of those with lower CD4+ cell counts. HIV-infected adults with cryptosporidiosis with CD4⁺ >50 cells/µL treated with 500 to 1,000 mg twice daily of nitazoxanide for 14 days experienced substantially higher rates of parasitological cure and resolution of diarrhea than persons receiving placebo treatment (Rossignol et al. 1998); however, this was not confirmed in two randomized trials in children (Amadi et al. 2002, 2009). This is probably due to key deficiencies in the host immune system, which are necessary for the thiazole to be effective. Ideal treatment for AIDS cases with cryptosporidiosis would therefore consist of cART to target the virus and antiparasitic agents to target the parasite (Shirley et al. 2012).

Adverse events associated with nitazoxanide are limited and typically mild, and no important drug-drug interactions have been reported. Nitazoxanide is not teratogenic in animals but human data on use in pregnancy are not available.

9.9.5 Paromomycin

Paromomycin is a nonabsorbable aminoglycoside that is used for the treatment of giradiasis and intestinal amebiasis. In animal models, it has shown efficacy in the treatment of cryptosporidiosis (Tzipori et al. 1994) A meta-analysis of 11 published human studies of paromomycin noted an overall response rate of 67 %; however, relapses were common, with a long-term success rate of only 33 % (Hashmey et al. 1997) The dose in adults is 25 to 35 mg/kg/day in 2 to 4 divided doses. Two randomized trials and a meta-analysis of these trials comparing paromomycin with placebo among patients with AIDS and cryptosporidiosis demonstrated a limited effectiveness of this drug; however, this analysis is limited by the small sample size and methodological issues (Cabada and White 2010; Hewitt et al. 2000; White et al. 1994). There appeared to be an improved response rate if paromomycin

was combined with cART in AIDS patients (Maggi et al. 2000, 2001). Some clinicians believed that paromomycin response correlated with the infection being primarily present in the colon rather than small bowel. In AIDS patients, paromomycin could be used instead of nitazoxanide, but needs to be combined with cART.

9.9.6 Macrolides

Macrolides including spiramycin, azithromycin, roxithromycin and clarithromycin have shown some activity for cryptosporidiosis, but this has been variable. Spiramycin in children at a dose of 100 mg/kg/day was associated with a more rapid resolution of cryptosporidiosis in an initial trial; however a second trial demonstrated no efficacy (Wittenberg et al. 1989). Intravenous spiramycin was associated with therapeutic benefit; however, this was associated with significant drug induced side effects (Stockdale et al. 2008). Azithromycin has efficacy in vitro and in animal models (Stockdale et al. 2008) and in many case reports in humans (Hicks et al. 1996; Kadappu et al. 2002; Nachbaur et al. 1997). In placebo controlled trials of azithromycin (900 mg/day) there was no effect seen (Blanshard et al. 1997; Stockdale et al. 2008; White 2010). Clarithromycin was active in animal and in vitro studies (Faver and Ellis 1993) as well as in some case reports from children in Egypt with cryptosporidiosis. Data from patients on this drug for MAI prophylaxis also suggested a protective effect. Roxithromycin demonstrated activity in two uncontrolled trials (Sprinz et al. 1998; Uip et al. 1998). Overall, in immune compromised patients with AIDS, macrolides could be used instead of nitazoxanide, but would need to be combined with cART.

9.9.7 Preventing Infection

Immune suppressed patients should be counseled regarding the routes of transmission of *Cryptosporidium spp.*, e.g. direct contact with infected adults, diaper-aged children and infected animals; coming into contact with contaminated water during recreational activities; drinking contaminated water; and eating contaminated food. Handwashing and careful hygiene can reduce the risk for diarrhea (Huang and Zhou 2007). Patients with HIV-infection should be advised to wash their hands after any potential contacts that could result in transmission of cryptosporidiosis. Immune suppressed individuals should avoid unprotected sex practices that could lead to direct (e.g., oral-anal) or indirect (e.g., penile-anal) contact with feces. HIV-infected patients (especially if their CD4⁺ is < 200 cells/µL), should avoid direct contact with diarrhea or stool. Immune suppressed patients should not, in general, consume raw shellfish as oocysts can survive in oysters for >2 months and have been found in oysters taken from commercial oyster beds (Fayer et al. 1999; Freire-Santos et al. 2000; Gomez-Couso et al. 2004; Graczyk and Schwab 2000). It should be appreciated, however, that there is no direct evidence, in humans, of transmission by this route.

Outbreaks of cryptosporidiosis have been linked to drinking water from municipal water supplies (Berkelman 1994; Bouzid et al. 2008; Leclerc et al. 2002). During outbreaks or in other situations that warrant a community advisory to boil water, boiling water for at least 3 min will eliminate the risk for cryptosporidiosis. Freezing does not kill this organism, so ice made from contaminated water is not safe for consumption. Immune suppressed patients can consider the routine use of submicron point of use water filters or bottled water as this can also reduce the risk for infection from municipal or well water sources.

As chronic cryptosporidiosis occurs primarily in persons with advanced immune deficiency, the initiation of cART in AIDS patients will prevent the majority of these infections. Rifabutin and clarithromycin when taken for *Mycobacterium avium-intracellulare* prophylaxis appear to provide protection against cryptosporidiosis; however, data are insufficient to recommend this as routine chemoprophylaxis for cryptosporidiosis (Fichtenbaum et al. 2000; Holmberg et al. 1998).

9.10 Conclusion

Cryptosporidiosis is a worldwide public health concern which can affect a variety of individuals irrespective of their immune status. Employing means to prevent both contamination of food and water is crucial for limiting the spread of this easily transmissible disease. Research needs to continue to focus on the development of effective therapeutic agents and preventative strategies, including vaccination. Significant work remains to identify optimal treatment regimens. In the meantime, public health measure such as consistent hand hygiene and safe food and water practices are critical for preventing infection, particularly in patients with a compromised immune system.

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Chapter 10 Immunology of Cryptosporidiosis

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Abstract Cryptosporidium spp. infect the gastrointestinal epithelium of vertebrate hosts. Intestinal species typically cause self-limiting diarrhea in immunocompetent individuals, suggesting an efficient host immune defense to eliminate the infection. Both innate and adaptive immunity are involved in host anti-parasite defense. Because of the "minimally invasive" nature of *Cryptosporidium* infection, mucosal epithelial cells are critical to the host's anti-Cryptosporidium immunity. Epithelial cells not only provide the first and rapid defense against *Cryptosporidium* infection, but also mobilize immune effector cells to the infection site to activate adaptive immunity. Attachment to the apical cell surface by Cryptosporidium, as well as molecules inserted into host cells after attachment, can activate host cell signal pathways and thereby alter cell function. Pathogen recognition receptors (e.g., Tolllike receptors) in epithelial cells recognize Cryptosporidium and initiate downstream signaling pathways (e.g., NF-kappaB) which trigger a series of antimicrobial responses and activate adaptive immunity. Non-coding RNAs are critical regulators of mucosal immunity to infection, while release of exosomes from epithelial cells may be a relatively unexplored, important component of mucosal anti-parasite defense. Conversely, it appears that *Cryptosporidium* has also developed strategies of immune evasion to escape host immunity, at least at the early stage of infection.

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Immune responses contribute to the pathophysiologic features of cryptosporidiosis. A better understanding the immunology of cryptosporidiosis will provide a framework for the potential development of novel therapeutic strategies.

10.1 Introduction

Host immune defense is critical to eliminate *Cryptosporidium* spp. infection. The self-limiting nature of Cryptosporidium infection in immunocompetent subjects suggests that the host activates an efficient immune response to eliminate the infection. After entry into host epithelial cells, the parasite resides within a unique intracellular but extracytoplasmic niche, separating the parasite from a direct interaction with other cell types. Therefore, Cryptosporidium spp. are classified as "minimally invasive" mucosal pathogens (Chen et al. 2002). Evidence from in vitro and in vivo studies indicates that both innate and adaptive immunity are involved in the resolution of cryptosporidiosis and resistance to infection (Chen et al. 2005b; Pantenburg et al. 2008). However, it appears that *Cryptosporidium* can survive the host innate immune attack during early stages of infection (Pantenburg et al. 2008; Zhou et al. 2009). Complete elimination of infection requires adaptive immune responses, particularly those mediated by CD4⁺ T cells (Pantenburg et al. 2010). Interferon (IFN)- γ , mainly released by activated CD4⁺ T cells (Boehm et al. 1997), is also critical in the control of cryptosporidiosis (Pollok et al. 2001).

Because of the "minimally invasive" nature of *Cryptosporidium* parasites, innate immune responses by epithelial cells are critical to the host's defense against the infection. Upon Cryptosporidium infection, epithelial cells quickly initiate a series of innate immune reactions including production of antimicrobial peptides (e.g., β -defensions) and release of inflammatory chemokines/cytokines, such as interleukin-8 (IL-8) (Hu et al. 2010; Zhou et al. 2012). Production and secretion of antimicrobial peptides (e.g., β -defensin 2) and nitric oxide (NO) can kill Cryptosporidium or inhibit parasite growth (Zhou et al. 2012). These chemokines/ cytokines of epithelial cell origin can mobilize and activate immune effector cells (e.g., lymphocytes, macrophages and neutrophils) to the infection sites (Blikslager et al. 2007). Recent evidence implicates that the release of epithelial exosomes may additional element of epithelial anti-Cryptosporidium be an defense (Hu et al. 2013). In addition, the molecular mechanisms of parasite-epithelial interactions in triggering epithelial defense and how immune responses contribute to the pathology of Cryptosporidium infection have been vigorously investigated (Pantenburg et al. 2008). Nevertheless, how parasite antigens are presented through the mucosal epithelium and how adaptive immunity is subsequently activated remain unclear. The potential strategies of Cryptosporidium immune evasion have recently been recognized and experimentally tested (Choudhry et al. 2009). Both transcriptional and post-transcriptional mechanisms are involved in the regulation of mucosal anti-Cryptosporidium defense. This chapter will highlight recent

advances in the immunology of cryptosporidiosis, particularly those pertaining to the human pathogenic *Cryptosporidium* species, *C. parvum*, and to the role of host epithelial anti-*C. parvum* defense. It will only briefly describe the immune responses from T cells and B cells and readers are referred to recent detailed reviews on this topic (Borad and Ward 2010; Petry et al. 2010).

10.2 Parasite-Epithelial Cell Interactions

The initial interaction between intestinal epithelia and the parasite is essential for parasitism, and hence for immune and pathological sequelea. C. parvum occupies a unique intracellular, yet extracytoplasmic, niche in the host cell, which may help protect the parasite against host immune responses as well as antimicrobial drugs. All developmental stages of C. parvum (asexual and sexual) occur in a single host. Upon ingestion of an oocyst, four infective sporozoites are released into the intestinal lumen. Immediately following this process (excystation), the sporozoites contact and attach to host derived mucins and exhibit gliding motility, a method of motility conserved in all apicomplexans (Chen et al. 2002; Bhat et al. 2007). Gliding requires the parasite cytoskeleton and the release of adhesive glycoproteins from micronemes, which are apical secretory organelles (Tomley and Soldati 2001). Ultimately, the apical pole of the parasite adheres to an epithelial cell, representing the first step of internalization (Lumb et al. 1988; Aji et al. 1991). A few characterized C. parvum secretory proteins have been implicated in the invasion process. These include GP900 (Petersen et al. 1992), the circumsporozoite-like (CSL) glycoprotein p23 (Perryman et al. 1996; Riggs et al. 1997), the Cpgp40/15 gene (Cevallos et al. 2000b), thrombospondin related proteins TRAP-C1 and TSP2-TSP12 (Spano et al. 1998; Deng et al. 2002; Putignani et al. 2008), and mucin-like antigens CpMuc4 and CpMuc5 (O'Connor et al. 2009). In addition, Gal/GalNAcspecific lectins may be essential for the initial attachment to host epithelial cells, including the Gal/GalNAc-specific lectin p30, that has been identified in both C. *parvum* and *C. hominis*, and mediates attachment to and invasion of epithelial cells in vitro (Bhat et al. 2007).

The invasion process, and consequently, the morphological and functional alterations of host epithelia, requires intimate contact between host plasma membrane and the apical pole of the parasite (Chen et al. 2004, 2005b). Early investigations demonstrated that *C. parvum* infection induced actin polymerization at sites of infection utilizing the actin branching and nucleation machinery of the Arp2/3 complex of proteins (Elliott et al. 2001). Multiple signaling pathways have been identified that modulate *C. parvum*-induced actin reorganization and parasite internalization, suggesting redundancy in these processes (Chen et al. 2003b, 2004). The identified signaling axes include: PI3-kinase-dependent activation of the small GTPase, CDC42; c-Src-dependent activation of cortactin; and Ca⁺⁺- dependent activation of the serine/threonine kinases PKC α and/or PKC β (Hashim et al. 2006; Perez-Cordon et al. 2011). Hence, actin dynamics and actin-dependent

membrane protrusion events are likely central to the formation of the intracellular/ extracytoplasmic niche occupied by *C. parvum*. Moreover, in *C. parvum*-infected biliary epithelial cells, the Na+/Glucose cotransporter, SGLT1, and aquaporin 1, a channel protein selective for the movement of water, accumulate at invasion sites, and participate in membrane protrusion events induced by the parasite (Chen et al. 2005a). The localization of these proteins is dependent on the localized contractility of myosin IIB (O'Hara et al. 2010a). Interestingly, SGLT1 may also protect host epithelial cells against pathogen-induced apoptosis (Yu et al. 2005, 2006a, 2008). Further research is needed to clarify whether these epithelial alterations reflect a protective innate host response, and/or a process that ensures efficient parasite entry into a viable cell.

The host-cell signaling cascades culminate in the rearrangement of the host actin cytoskeleton, and parasite internalization within a parasitophorous vacuole membrane. The parasitophorous vacuole membrane envelops the parasite on the luminal surface of the cell in a parasite-modified host membrane (Robert et al. 1994; Bonnin et al. 1995; McDonald et al. 1995; O'Hara et al. 2004). As the invasion process ensues, a unique electron dense structure forms within the host, adjacent to a network of microfilaments. The adjacent parasite membrane becomes highly invaginated. Collectively, these structures are referred to as the feeder or attachment organelle. Hence, the parasite occupies a niche that is selectively isolated from the intestinal lumen and host cytoplasm.

10.3 Activation of Pattern Recognition Receptors (PRRs) in Epithelial Cells

The gastrointestinal epithelium, in addition to providing a natural barrier that limits infection, also plays a critical role in the initial recognition of parasites and the triggering of adaptive immunity. Pathogen-associated molecular pattern molecules (PAMPs) derived from microorganisms can be recognized by PRR-bearing cells (Medzhitov 2007). When ligated by the PAMPs, these PRRs trigger cytokine and costimulatory signals that initiate both innate and adaptive cellular responses. Activation of downstream signaling cascades of PRRs regulate many biological processes that are critical to immune response to infection. Epithelial cells along the gastrointestinal and biliary tracts express a variety of PRRs, such as the Toll-like receptors (TLRs) and nucleotide binding NOD-like receptors, RIG-I-like receptors, and oligomerization domain-like receptors, which recognize pathogens or PAMPs (Nasu and Narahara 2010).

Studies on the activation of PRRs and associated downstream signaling cascades in epithelial cells following *C. parvum* infection have focused on the TLR4/NF- κ B pathway. The NF- κ B pathway controls many genes involved in immune responses, inflammation and cell differentiation (Zhou et al. 2009; Oeckinghaus et al. 2011). Activation of the NF- κ B pathway has been demonstrated in epithelial cells following *C. parvum* infection in a variety of studies using both in vivo and in vitro models. Activation of NF- κ B signaling in epithelia cells following *C. parvum* infection appears to be a "late" but persistent response, compared with the early and transient NF- κ B activity in epithelial cells following stimulation of lipopoly-saccharide (LPS) and tumor necrosis factor- α (TNF- α) (Zhou et al. 2009). It is also of interest that *C. parvum*-induced NF- κ B activation requires direct parasite-epithelial cell interaction. Indeed, it appears that parasite-induced NF- κ B activation is limited to infected epithelial cells, at least in the early stage of infection. Hence, *C. parvum*-induced TLR4/NF- κ B activation, likely requires the process of parasitic invasion (Chen et al. 2001, 2005b).

While the precise molecular mechanisms underlying C. parvum-induced activation of TLR4/NF-kB signaling are still unclear, it is possible that parasite-derived molecules act as PAMPs and are recognized by host cells as a TLR4 ligand during the invasion process of the parasite. However, the potential PAMPs from C. parvum required for TLR4 activation are not known. Indeed, the C. parvum genome lacks enzymes associated with endotoxin synthesis (Abrahamsen et al. 2004), hence, C. parvum does not express LPS or LPS-like molecules. Moreover, C. parvum sporozoites do not initiate limulus amebocyte lysate activity (unpublished data). Recruitment of TLR4 and TLR2 to the attachment/invasion site was reported in H69 human biliary epithelial cells after exposure to C. parvum in vitro (Chen et al. 2005b). Whether this recruitment is required for the activation of TLR4 signaling is unclear. Furthermore, whether infection can trigger activation of other PRRs and downstream signaling pathways, in particular, the inflammasome activation pathways (Newton and Dixit 2012), is yet to be explored. However, knockdown of TLR4 and its adaptor protein MyD88 can block C. parvum-induced NF- κ B activation (Chen et al. 2005b), suggesting an integral role of TLR4 initiated signaling. Importantly, as discussed in detail below, activation of TLR/NF-KB signaling may account for the majority of epithelial anti-parasite defense.

10.4 Epithelial Defense Response

10.4.1 Epithelial Cell-Derived Effector Molecules

Human gastrointestinal epithelial cell lines increase the expression and production of the potent neutrophil chemoattractants interleukin-8 (IL-8) and chemokine (C-X-C motif) ligand 1 (CXCL1) following *C. parvum* infection. IL-8 and CXCL1 were shown to be predominantly released from the basolateral surface of infected cultures using polarized epithelial monolayers cultured in Transwell chambers (Ojcius et al. 1999). Cultured human colon epithelial cells infected with *C. parvum* increased prostaglandin E2 (PGE2) production as much as 50-fold and prostaglandin F2alpha (PGF2 α) production by up to tenfold compared with uninfected cells. Increases were observed by 12 h after infection, were maximal by 36 h and were sustained over a 72 h culture period (Kagnoff and Eckmann 1997). In addition, CXCL10 is highly upregulated in intestinal epithelial cells of AIDS patients with active cryptosporidiosis (Wang et al. 2007). Induction of CX3CL1 by *C. parvum* has been reported in cultured human intestinal and biliary epithelial cells (Harada and Nakanuma 2012). *C. parvum* infected epithelial cells respond by upregulating certain C, C–C, and C–X–C class chemokines as shown in neonate mice (Lacroix-Lamande et al. 2002). Epithelial cells are also a source of IL-18 (IFN- γ -inducing factor) (Okazawa et al. 2004). The IL-18 gene is upregulated in ilea of mice in response to infection (Tessema et al. 2009), and IL-18 mRNA and protein are upregulated in vitro upon *C. parvum* infection of intestinal epithelial cells (McDonald et al. 2006).

Release of these cytokines and chemokines from epithelial cells following *C*. *parvum* infection will attract inflammatory infiltration, and facilitate mucosal antimicrobial defense (Fig. 10.1). Besides cytokine/chemokine release, *C. parvum* infection also induces expression of adhesion molecules in epithelial cells. Intercellular adhesion molecule (ICAM-1) is long known for its importance in stabilizing cell-cell interactions and is critical for the firm arrest and transmigration of leukocytes out of blood vessels into tissues (Muller 2009). Upregulation of ICAM-1 transcription and protein expression was detected in human gastrointestinal epithelial cells following *C. parvum* infection in vitro (Gong et al. 2011). Expression of adhesion molecules, such as ICAM-1, on infected epithelial cell surfaces may facilitate recruitment of lymphocytes to the site of infection (Gong et al. 2011). Indeed, expression of ICAM-1 on infected epithelial cells facilitated epithelial adherence of co-cultured Jurkat T cells (Gong et al. 2011).

Several molecules synthesized in epithelial cells following *Cryptosporidium* infection reveal anti-*C. parvum* capacity, and thus play a direct role in antimicrobial defense (Fig. 10.1). Nitric oxide (NO) is produced from arginine by the enzyme nitric oxide synthase (NOS), which exists in three isoforms. One isoform, the inducible NOS (iNOS), is known to be inducibly expressed in human intestinal epithelial cells (Ricciardolo et al. 2004), and NO has been shown to have a broad range of antimicrobial properties. NO synthesis by the epithelium is significantly increased following *C. parvum* infection; and, in the absence or inhibition of iNOS, infection and oocyst shedding are significantly exacerbated (Leitch and He 1999; Gookin et al. 2005, 2006). L-arginine administered enterically significantly reduced fecal oocyst shedding in athymic nude mice chronically infected with *C. parvum* infection induced NO production in host epithelial cells in a NF- κ B-dependent manner, with the involvement of the stabilization of iNOS mRNA (Zhou et al. 2012).

In addition, α - and β -defensins, as well as other low-molecular mass antimicrobial peptides, comprise part of the arsenal of innate defenses present in the gastrointestinal tract. Some of these peptides have been shown to have anticryptosporidial activity in vitro (Dommett et al. 2005). *C. parvum* induces expression of β -defensin-2 in human biliary epithelial cells via TLR-mediated NF- κ B activation, whereas β -defensin-1 appears to be constitutively expressed. Both

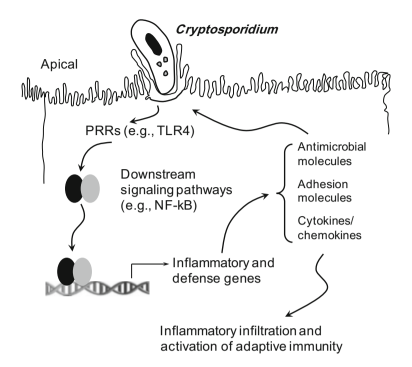


Fig. 10.1 Production and release of epithelial cell-derived effector molecules is essential for the activation of mucosal anti-*Cryptosporidium* immunity. *C. parvum* infection activates pathogen recognition receptors (e.g., TLR4) and their downstream signaling pathways (e.g., NF-κB). This activation induces the expression of cytokine and chemokine inflammatory genes and antimicrobial defense genes. Release of these cytokines and chemokiens from epithelial cells following infection will attract inflammatory infiltration, resulting in activation of adaptive immunity. Several molecules synthesized in epithelial cells following infection, including LL-37, β-defensins and NO, reveal anti-*C. parvum* capacity and play a direct role in anti-microbial defense

β-defensin-1 and β-defensin-2 have anti-*C. parvum* activity as evidenced by a decreased sporozoite viability following incubation with recombinant β-defensin-1 or 2 (Zaalouk et al. 2004). One β-defensin, enteric β-defensin, was shown to be highly expressed in epithelial cells in bovine distal small intestinal crypts and colon (Tarver et al. 1998). Moreover, expression of enteric β-defensin mRNA was upregulated by five to tenfold following *C. parvum* infection of calves, further supporting a possible role of β-defensins in the innate immune response to *C. parvum* infection (Tarver et al. 1998).

Secretion of mucins in the gastrointestinal tract is an important element of mucosal defense. Nevertheless, their role in host defense to *Cryptosporidium* infection is still unclear. Mucins are a family of high molecular weight, heavily glycosylated proteins (glycoconjugates) produced by epithelial tissues in most metazoans (Marin et al. 2008). Although some mucins are membrane-bound due to the presence of a hydrophobic membrane-spanning domain that favors retention

in the plasma membrane, most mucins are secreted onto mucosal surfaces. It has been speculated that epithelial mucin expression could protect the host against further infection by extracellular stages of *C. parvum*. Notably, several mucin-like molecules have been identified in *C. parvum* which may mediate the parasite attachment to or invasion of the host cells (Cevallos et al. 2000a).

Moreover, recent studies indicate that mucosal epithelial cells along the gastrointestinal and urogenital tracts can secrete several specific cytokines, so called epithelial-derived cytokines, such as IL-25, IL-33, IL-34, and thymic stromal lymphoietin (TSLP), upon extracellular stimuli (Maynard et al. 2012). Release of these epithelial-derived cytokines plays a critical role in mucosal immunity against mucosal infection (Taylor et al. 2009). Additional epithelial cell-derived molecules such as serum amyloid A 3 (Saa3) and regenerating islet-derived 3 gamma (Reg3g) have also recently been demonstrated to display anti-microbial activity (Reigstad et al. 2009; Choi et al. 2013). Their potential involvement in *Cryptosporidium* infection of gastrointestinal epithelium has yet to be investigated.

10.4.2 MicroRNAs (miRNAs)

miRNAs are short strands of non-coding RNAs of 18-24 nucleotides in length (Bartel 2009). These RNA molecules are emerging as key mediators of many biological processes and impact gene expression at the post-transcriptional level (Bartel 2009). Similar to other RNA molecules, most miRNAs are initially transcribed as primary transcripts (termed pri-miRNAs) by RNA polymerase II and processed by the RNase III Drosha (in the nucleus) and a second RNase III Dicer (in the cytoplasm) to generate mature miRNA molecules (Lee et al. 2003). Mature miRNAs are incorporated as single-stranded RNAs into a ribonucleoprotein complex, known as the RNA-induced silencing complex (RISC). The RISC identifies target mRNAs based on perfect or nearly perfect complementarity between the miRNA and the target mRNA 3'-untranslated region (3'UTR). The interaction between the RISC and mRNA causes either mRNA cleavage and/or translational suppression, resulting in gene suppression at the post-transcriptional level (Bartel 2009). More importantly, it has become clear that miRNAs have diverse expression patterns and play essential roles in biological processes that include development, maintenance of genome stability and viral adaptive defense mechanisms (Lewis et al. 2005). It has been predicted that miRNAs control 20-30 % of human genes (Lewis et al. 2005; Bartel 2009).

miRNAs are involved in regulation of epithelial responses to *C. parvum* infection. Differential mature miRNA expression profiles in biliary epithelial cells following *C. parvum* infection were reported using an in vitro model of human cryptosporidiosis. Transcription of a subset of miRNA genes was found to be regulated through NF- κ B activation in human biliary epithelial cells following *C. parvum* infection (Zhou et al. 2009). Specifically, inhibition of NF- κ B activation by SC-514, an IKK2 inhibitor, blocked LPS- and *C. parvum*-induced upregulation of a

subset of pri-miRNAs, including pri-miR-125b-1, pri-miR-21, pri-miR-23b-27b-24-1 and pri-miR-30b. Moreover, direct binding of NF- κ B p65 subunit to the promoter elements of *mir-125b-1*, *mir-21*, *mir-23b-27b-24-1* and *mir-30b* genes was identified by chromatin immunoprecipitation analysis and confirmed by a luciferase reporter assay using constructs covering the potential promoter elements of these miRNA genes (Zhou et al. 2009, 2010). In addition to the upregulation of some miRNAs, NF- κ B signaling may be also involved in the downregulation of miRNA genes. Transcription of the *let-7i* gene in biliary epithelial cells in response to LPS stimulation and *C. parvum* infection has been reported to be suppressed through promoter binding by NF- κ B subunit p50 along with CCAAT/enhancerbinding protein beta (C/EBP β) (O'Hara et al. 2010b). Importantly, functional inhibition of selected NF- κ B-responsive miRNAs in epithelial cells increased *C. parvum* burden (Zhou et al. 2009). Thus, upregulation of miRNAs through NF- κ B signaling may be relevant to the regulation of epithelial anti-microbial defense.

NF-kB-responsive miRNAs may modulate epithelial anti-parasite responses at every step of the innate immune network (Fig. 10.2). Elements of the TLR/NF-κB pathway are potential targets for the NF- κ B-responsive miRNAs, thus, providing feedback regulation of TLR/NF- κ B signaling. For example, TLR4 is targeted by *let-7* miRNAs and suppression of *let-7* facilitates the synthesis of TLR4 protein in epithelial cells following Cryptosporidium infection, which promotes epithelial innate defense (Chen et al. 2007). Targeting the suppressors of cytokine signaling family of proteins (CIS/SOCS) by miR-98/let-7 may also exert a positive feedback regulation of the innate immune response in infected cells. C. parvum infection induces CIS/SOCS expression in human biliary epithelial cells through TLR/NF-kB-suppressed expression of miR-98 and let-7 (Hu et al. 2009, 2010). Induction of CIS expression enhances IκBα degradation promoting NF-κB activation (Hu et al. 2009). On the other hand, upregulation of miR-21 in infected cells may result in a negative feedback regulation of the TLR4/NF-κB signaling through targeting of PDCD4 (Sheedy et al. 2010). It was reported that LPS decreases expression of PDCD4 through induction of miR-21, resulting in subsequent inhibition of NF-κB signaling activity and promotion of IL-10 expression in human peripheral blood mononuclear cells (Sheedy et al. 2010). Similarly, miR-21 targeting of PDCD4 was shown to influence TNF-induced activation of NF-kB.

NF-κB-responsive miRNAs may play a role in the expression of many NF-κBregulated immune or inflammatory genes. Based on bioinformatics analysis, approximately 29 % of cytokine/chemokine mRNAs have potential target sites for miRNAs (Asirvatham et al. 2008). Many of these cytokine/chemokine mRNAs are transcripts of NF-κB-regulated genes. Expression of IFN- β , the main type I IFN cytokine important to the initiation of innate responses in response to infection including *Cryptosporidium*, is finely controlled by various miRNAs. It has been experimentally confirmed that miR-26a, -34a, -145, and *let-7b* directly regulate IFN- β production by targeting IFN- β 3'UTR (Witwer et al. 2010).

Intriguingly, miRNAs may target RNA-binding proteins to regulate mRNA stability. Several RNA-binding proteins, including the KH-type splicing regulatory protein (KSRP, also known as KHSRP), tristetraprolin (TTP) and Hu antigen R

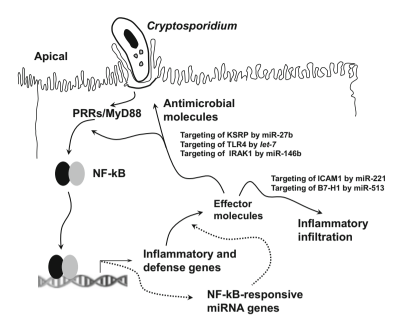


Fig. 10.2 Post-transcriptional gene suppression mediated by miRNAs is involved in the finetuning of epithelial anti-*Cryptosporidium* defense. *C. parvum* infection alters the expression profile of miRNA genes in infected epithelial cells. A panel of miRNA genes is trans-activated or trans-suppressed through the activation of the NF-κB pathway. These NF-κB-responsive miRNAs may modulate epithelial anti-parasite responses at every step of the innate immune network, including feedback regulation to the TLR/NF-κB signaling, post-transcriptional regulation of many of NF-κB-regulated immune or inflammatory genes, and production of anti-microbial effector molecules

(HuR), recognize AU-rich elements (AREs) within the 3'UTRs of mRNAs and diminish their half-life in the cytoplasm (Chen and Shyu 1995; Dean et al. 2004; Anderson 2008). For example, many cytokine mRNAs have AREs in their 3'UTRs and usually exhibit a very short half-life (in the range of 10-30 min) in resting cells, effectively preventing cytokine production in inactive cells. In activated cells, cytokine mRNA half-lives are in the range of several hours, accounting for a significant increase in protein production (Stoecklin and Anderson 2006; Anderson 2008). In this regard, KSRP interacts with these mRNAs that have the AREs within their 3'UTRs, including mRNAs for iNOS, IL-8 and cyclooxygenase-2 (COX-2), and is a key mediator of mRNA decay (Winzen et al. 2007; Subramaniam et al. 2008; Pautz et al. 2010). Thus, the level of KSRP can potentially coordinate mRNA stability in response to extracellular stimuli (Winzen et al. 2007). Indeed, induced expression of IL-8, NO (synthesized by iNOs) and PGE₂ (catalyzed by COX-2) has been implicated in the response to a variety of pathogens, including C. parvum (Stoecklin and Anderson 2006; Anderson 2008). We recently demonstrated that C. parvum infection of epithelial cells increases transcription of the mir-23b-27b-24-1 gene in an NF- κ B-dependent manner. Moreover, KSRP was

downregulated in epithelial cells following *C. parvum* infection. Importantly, miR-27b targeted the 3'UTR of KSRP mRNA and suppressed its translation. Hence, *C. parvum* infection resulted in decreased KSRP expression through upregulation of miR-27b. Functional manipulation of KSRP or miR-27b caused reciprocal alterations in iNOS mRNA stability in infected cells. Additionally, forced expression of KSRP and inhibition of miR-27b resulted in an increased *C. parvum* burden (Zhou et al. 2012). Therefore, downregulation of KSRP and upregulation of miR-27b stabilize iNOs/IL-8/COX-2 mRNAs, contributing to epithelial anti-*C. parvum* defense (Fig. 10.2). Notably, no complementarity to IFN- γ or anti-microbial peptide mRNAs (such as LL-37 and β -defensins) has been identified for these NF- κ B-responsive miRNAs.

The involvement of NF- κ B-responsive miRNAs in the expression of co-stimulatory molecules, in the epigenetic impact and in the shuttling of miRNAs through release of exosomes in *Cryptosporidium* infection will be discussed in the following sections. In addition, miRNAs have been identified in both mammalian cells and non-mammalian cells including virus and parasites (Bartel 2004; Scaria et al. 2006; Winter et al. 2009). Expression of miRNAs in *C. parvum* has not yet been examined and whether *C. parvum*-derived miRNAs, if there are any, can be localized in infected host cells is unknown. The concept that a pathogen encodes mRNAs targeted by host miRNAs has recently emerged as an important mechanism of anti-viral defense (Pedersen et al. 2007). Likewise, it is of interest to test the possibility that host cell miRNAs target the internalized parasite mRNAs and silence genes of the pathogen. The direct *C. parvum*-host cell cytoplasmic tunnel-connection (Huang et al. 2004) could mediate exchange of molecules, including miRNAs, between the host cells and the internalized parasite.

10.4.3 Exosomes

Exosomes represent a specific subtype of secreted membrane vesicles that are about 30–100 nm in size and are formed inside secreting cells in endosomal compartments called multi-vesicular bodies (MVBs) (Thery 2011). Exosomes are produced by a variety of cells (e.g., reticulocytes, epithelial cells, neurons, tumor cells) and have been found in bronchoalveolar lavage, urine, serum, bile, and breast milk (Thery 2011; Hu et al. 2012a). The composition of exosomes is heterogenous, depending on their cellular origin. Exosomes do not contain a random array of intracellular proteins, but a specific set of protein families arising from the plasma membrane, the endocytic pathway, and the cytosol, especially those of endosomal origin, such as CD63, ICAM-1, and MHC molecules (Bobrie et al. 2011). These vesicles also mediate the secretion of a wide variety of other molecules, such as lipids, mRNAs, and miRNAs, and thereby traffic molecules from the cytoplasm and membranes of one cell to other cells or extracellular spaces (Smalheiser 2007; Valadi et al. 2007). There is increasing evidence that secreted vesicles play an important role in normal physiological processes, development, viral infection, and

other human diseases (Liegeois et al. 2006; Yu et al. 2006b; Kolotuev et al. 2009; Thery et al. 2009).

We recently demonstrated that luminal release of exosomes from the biliary and intestinal epithelium is increased following infection by *C. parvum* (Hu et al. 2013). Secretion of exosomes is regulated by various stimuli, including the activation of P2X receptor by ATP on monocytes and neutrophils, thrombin receptor on platelets, and TLR4 by LPS on dendritic cells (Bhatnagar and Schorey 2007). Formation of exosomes within MVBs and targeting of transmembrane proteins involve a complex intracellular sorting network, including the endosomal sorting complex required for transport (ESCRT) machinery (van Niel et al. 2006). Fusion of MVBs with plasma membrane is an exocytic process that requires the association of v-SNAREs (from the vesicles) and t-SNAREs (at the membrane) to form a ternary SNARE (SNAP receptor) complex. The SNARE complex brings the two membranes in apposition, a necessary step in overcoming the energy barrier required for membrane fusion (Sudhof and Rothman 2009; Kennedy and Ehlers 2011). Several Rab family proteins, including Rab11 and Rab27b, are key regulators of the exosome secretion pathway and are involved in MVB docking at the plasma membrane (Ostrowski et al. 2010). We found that C. parvum-stimulated release of exosomes involves TLR4/IKK2 activation and the SNAP23-associated vesicular exocytic process (Hu et al. 2013). Whereas a basal level of exosomal luminal release exists in cultured biliary epithelial monolayers and in the murine biliary tract, a TLR4-dependent increase in luminal release of epithelial exosomes was detected following C. parvum infection. We demonstrated a significant co-localization of Rab11 and Rab27b with MVBs in the cytoplasm of epithelial cells following C. parvum infection. C. parvum infection also increases expression and enhances phosphorylation of SNAP23 in infected cells. Activation of TLR4 may contribute to both events: TLR4 signaling increases SNAP23 protein expression through diminished let-7- mediated SNAP23 translational repression, and stimulates SNAP23 phosphorylation through activation of IKK2 (Suzuki and Verma 2008).

Intriguingly, released exosomes contain antimicrobial peptides with anti-*C*. *parvum* activity, thus contributing to mucosal anti-*C. parvum* defense (Fig. 10.3). The anti-*C. parvum* activity of β -defesin-2 and LL-37 was previously reported in intestinal and biliary cryptosporidiosis (Zaalouk et al. 2004; Chen et al. 2005b). Both β -defesin-2 and LL-37 were identified as components of apical exosomes released from human biliary epithelial cell H69 monolayers (Hu et al. 2013). Moreover, apical exosomes from infected cells demonstrated an increase in β -defesin-2 and LL-37 content compared to exosomes from uninfected cells, while inhibition of TLR4 signaling diminished their presence in exosomes (Hu et al. 2013). Because biliary epithelial cell derived-exosomes display a protein profile different from the whole cell lysate, targeting of cytoplasmic proteins to the MVBs within epithelial cells, such as β -defesin-2 and LL-37, must be controlled through highly selective and regulated mechanisms. Moreover, exposure of *C. parvum* sporozoites to released exosomes decreases their viability and infectivity both in vitro and ex vivo (Hu et al. 2013). Anti-*C. parvum* activity of apical exosomes released

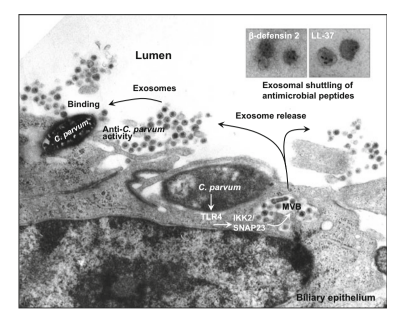


Fig. 10.3 Luminal release of exosomes from epithelium contributes to TLR4-mediated mucosal anti-*Cryptosporidium* defense. Apical exosomal vesicles are secreted by biliary epithelial cells through activation of the TLR4 signaling pathway following *C. parvum* infection. Released exosomal vesicles shuttle antimicrobial peptides, such as cathelicidin-37 and β -defensin 2, and activation of TLR4 signaling increases exosomal shuttling of antimicrobial peptides. Functionally, released exosomal vesicles can directly bind to the *C. parvum* sporozoite surface and display antimicrobial activity

from the epithelium may involve direct binding to the *C. parvum* sporozoite surface. Direct binding of exosomes to the *C. parvum* sporozoite surface was evident by scanning and transmission EM. Confocal microscopy analysis with exosomal labeling confirmed the binding of exosomes to *C. parvum* sporozoites after incubation. Interestingly, treatment of exosomes with Gal/GalNAc-specific lectin arachis hypogaea (PNA, peanut agglutinin) diminished their anti-*C. parvum* activity, suggesting that these molecules may be involved, at least partially, in exosomal binding to the *C. parvum* sporozoite surface. The lifecycle of *C. parvum*, both in vitro and in vivo, has extracellular stages (i.e., sporozoites, merozoites, and microgametocytes), and they are likely vulnerable to exosomal binding/targeting.

Release of epithelial cell-derived exosomes to the basolateral domain has also been demonstrated following *Cryptosporidium* infection (Hu et al. 2013). Although these basolateral exosomes don't exhibit anti-*C. parvum* activity as the apical exosomes, they do shuttle MHC proteins (Hu et al. 2013) and therefore, may contribute to antigen presentation (as described below). In addition, the basolateral exosomes have been shown to modulate lymphocyte immune responses during other mucosal infections (Mallegol et al. 2007). Intestinal epithelial cell-derived exosomes containing $\alpha\nu\beta6$ integrin and food antigen induced the generation of tolerogenic dendritic cells in a model of tolerance induction (Chen et al. 2011). The presence of these intestinal epithelial cell-derived exosomes impacted the development of antigen-specific T regulatory cells (Chen et al. 2011). Intriguingly, exosome-shuttled miRNA molecules can be delivered to other cell types through exosomal uptake (Hu et al. 2012b). Therefore, exosome-mediated transport of miRNAs may provide a novel mechanism of gene regulation between cells. Ohshima et al. showed that there is the enrichment of *let-7* miRNA family in the exosomes from AZ-P7a cells (Ohshima et al. 2010). Given the importance of miRNAs in epithelial innate immune responses, it would be interesting to determine if exosomes from epithelial cells also carry miRNAs and thus modulate epithelial-immune cell interactions and epithelial antimicrobial defense, via exosomal delivery of miRNAs.

10.5 Antigen Presentation

C. parvum resides at the apical surface of intestinal epithelial cells and does not invade deeper layers of the human gastrointestinal mucosa. The mechanisms by which infection with a pathogen, like C. parvum, that is confined to epithelial cells, result in activation of adaptive immunity is a subject of considerable interest. Microscopic analyses identified the presence of C. parvum in M cells in AIDSrelated intestinal cryptosporidiosis (McCole et al. 2000). Infiltration of various inflammatory cells to the lumen of the biliary tract has been observed in liver tissues from AIDS patients with biliary cryptosporidiosis (Lumadue et al. 1998; Chen and LaRusso 1999). Therefore, it is possible that direct interaction between parasite and immune cells, including macrophages or dendritic cells (DCs), may present parasite antigens and activate the adaptive immunity. Until now, there is relatively little work directly addressing the role of DCs in the immune response to C. parvum (McDonald et al. 2013). Bone-marrow derived mouse dendritic cells exposed to C. parvum sporozoites or parasite antigens expressed IFN- α and IFN- β within a few hours (Barakat et al. 2009b). Soluble sporozoite antigens or recombinant parasite antigens induced maturation of these dendritic cells and stimulated production of IL-12, IL-1β and IL-6 (Barakat et al. 2009b). Human monocytic dendritic cells also increased production of IL-12 after exposure to soluble sporozoite antigens or live sporozoites (Barakat et al. 2009b; Bedi and Mead 2012).

Several pieces of evidence suggest that epithelial cells may play a role in antigen presentation during infection. The machinery required for antigen processing and presentation include major histocompatibility complex (MHC) class I and class II molecules and co-stimulatory molecules (Hershberg and Mayer 2000). As in other tissues, macrophage phagocytosis of apoptotic cells containing the pathogens may contribute to antigen presentation. Apoptotic cell death of gastrointestinal epithelial cells during *C. parvum* infection has been reported both in vivo and in vitro (Lacroix et al. 2001; Liu et al. 2008). Additionally, induction of MHC-I, MHC-II and the co-stimulatory molecule B7-H1 has been reported in biliary and intestinal

epithelial cells following *Cryptosporidium* infection (Gong et al. 2010). Moreover, as described above, release of epithelial-derived exosomes has recently been implicated in antigen presentation during mucosal infection. Our recent studies demonstrate basolateral release of epithelial-derived exosomes using in vitro models of intestinal and biliary cryptosporidiosis. Of importance, those released exosomes carry both MHC-I and MHC-II (Hu et al. 2013). Coupled with the fact that epithelial cells release an array of cytokines/chemokines and adhesion molecules to attract immune cell infiltration, epithelial cells may play an active role in the antigen presentation during *Cryptosporidium* infection.

10.6 IFNs, NK Cells, Macrophages, and the Complement System

A critical role for NK cells and IFN-y in anti-Cryptosporidium immunity has been documented in T and B cell-deficient mice (Barakat et al. 2009a; McDonald et al. 2013). In a recent study using mice lacking T-cells and NK-cells (SCIDbg mice) and SCIDbg mice additionally lacking macrophages and neutrophils (SCIDbgMN mice), C. parvum developed an acute infection with high mortality in SCIDbgMN mice but not in SCIDbg mice (Takeuchi et al. 2008). NK cells may carry out their anti-Cryptosporidium function through release of IFN- γ , because the resistance of SCID mice deficient in T- and B- lymphocytes to C. parvum infection is IFN-y-dependent (Ungar et al. 1991; Chen et al. 1993; McDonald and Bancroft 1994). Splenic NK-cells from SCID mice which had been activated in vitro by C. *parvum* antigen produce IFN- γ (Chen et al. 1993). IFN- γ may also be released by cells other than NK-cells, including macrophages (McDonald et al. 2013). In support of this, a diligent study by Barakat et al. (2009a), demonstrated that in mice deficient in B-, T-, and NK-cells (Rag $2^{-/-}\gamma c^{-/-}$), oocyst excretion is higher than in $Rag2^{-/-}$ mice that are deficient in B- and T-cells only. Although lacking Tand NK-cells, the level of infection can be exacerbated in Rag $2^{-/-}$ yc $^{-/-}$ mice by antibody neutralization of IFN- γ , suggesting that an additional intestinal cell type may be an important source of IFN- γ .

The complement system, a part of the humoral innate immune system, can be activated and can participate in host anti-*Cryptosporidium* defense. Petry et al (2008) analyzed the in vitro binding and activation of the human and mouse complement systems and tested the susceptibility to infection in complement-deficient mouse strains (Petry et al. 2008). Their results show that *C. parvum* can activate both the classical and lectin pathways, leading to the deposition of C3b on the parasite. Using real-time PCR, parasite development could be demonstrated in adult mice lacking mannan-binding lectin (MBL-A/C^{-/-}) but not in mice lacking complement factor C1q (C1qA^{-/-}) or in wild type C57BL/6 mice. MBL belongs to the C-type lectin superfamily and can recognize carbohydrate patterns of microorganisms, resulting in activation of the lectin pathway of the complement system

(Worthley et al. 2005). The contribution of the complement system and the lectin pathway in particular to the host defense against cryptosporidiosis may become apparent in situations of immunodeficiency such as HIV infection or in early childhood.

10.7 T Cells and B Cells

The role of acquired immunity in resistance to and clearance of *C. parvum* infection has been studied most extensively in murine models in which B- and T-cell populations, or both, can be selectively depleted. Both humoral and cellular immunity play a role in the control of this infection, but the latter plays the major role, mainly in the intestinal mucosa.

Acquired resistance to cryptosporidial infection in humans is dependent upon T-cells with the $\alpha\beta$ -form T-cell receptor; in addition, the CD4⁺ T-cell subset has a protective role whereas $\gamma\delta^+$ or CD8⁺ subpopulations of T-cells appear irrelevant or subordinate (Mombaerts et al. 1993; Ladel et al. 1995). The ability to clear acute and chronic *C. parvum* infection in humans correlates closely with host CD4 T-cell levels (Takeuchi et al. 2008). Similarly, studies in congenitally athymic (nude) mice and reconstitution studies in immunodeficient SCID mice demonstrate a critical role for T cells in resistance to, and clearing of, infection (Ruittenberg and van Noorle Jansen 1975). Both $\alpha\beta$ T cells and $\gamma\delta$ T cells are important for host defense to *C. parvum* in mice (Eichelberger et al. 2000). As in humans, CD4, but not CD8 T cells, were shown to be most important for clearing infection in mice (Wong and Pamer 2003).

Cytokines of Th1 (e.g., IFN- γ , IL-2, and IL-12), Th2 (e.g., IL-4, IL-5, IL-6, and IL-10), and Th17 (e.g., IL-23) may all play a role in host anti-*Cryptosporidium* immunity (Ehigiator et al. 2005; Weaver et al. 2006). It has been suggested that the capacity to produce all Th1, Th2 and Th17 cytokines, rather than the presence of Th2 cytokines alone, determines the effective immune response against *C. parvum* infection. The susceptibility or resistance to *C. parvum* infection depends on a delicate balance between the production of Th1 cytokines, needed to control parasite growth, and Th2 cytokines, to limit pathology. IL-18 appears to be involved in the regulation of the Th1/Th2 responses in *C. parvum*, because neutralization of IL18 resulted in a cytokine imbalance with upregulation of systemic (spleen) Th2 cytokine genes, notably IL-4 and IL-13 (Tessema et al. 2009).

C. parvum infection is accompanied by an antibody-mediated Th2 response in infected individuals with production of parasite-specific immunoglobulin (Ig) of all major classes (Ungar et al. 1986; Williams 1987; Hill et al. 1990; Peeters et al. 1992). IgG, IgM and IgA have been detected in serum and mucosa of humans and animals following infection. High levels of these immunoglobulins have been detected in AIDS patients with chronic cryptosporidiosis (Ungar et al. 1986; Cozon et al. 1994; Reperant et al. 1994; Jakobi and Petry 2008). Mucosal IgA may assist the cellular immune response by coating surfaces of the infective stages and

preventing attachment to the epithelial cells. However, studies employing B-celldeficient $IMT^{-/-}$ mice indicate that B-cells are not essential for resistance to and recovery from *C. parvum* infections in mice (Chen et al. 2003a). Therefore, the antibody response observed during *C. parvum* infection may be of minor importance as it is raised mainly to the exposed antigens of the extracellular stages of the parasite. This remains the major obstacle to the use of classical vaccines for prevention of cryptosporidiosis.

10.8 Evasion of Host Immunity and Defense

Host cell death following invasion by intracellular pathogens is an ancient defense mechanism (Velmurugan et al. 2007), which involves infiltration and activation of various immune effector cells such as macrophages, neutrophils and lymphocytes. Many intracellular pathogens have developed strategies to inhibit the induction of host cell death (Luder et al. 2001; Ashida et al. 2011). The intricate interaction between host and *C. parvum* is exemplified by the modulation of epithelial cell apoptosis during the course of cryptosporidiosis (Fig. 10.4). *C. parvum* has the ability to either induce or inhibit enterocyte apoptosis. Completion of the *C. parvum* life cycle requires viable host cells and detachment or apoptosis of infected cells in vitro leads to decreased parasite numbers (Widmer et al. 2000). Therefore, it is proposed that apoptosis may play a role in limiting and/or clearing the infection.

Early studies demonstrated that C. parvum infection of intestinal epithelial cells correlates with the activation of NF-κB target genes (Laurent et al. 1997, 1998; Seydel et al. 1998). NF- κ B activation has been shown to prevent apoptosis in several experimental models (Beg and Baltimore 1996; Orange et al. 2005) and *Cryptosporidium*-infected cells activate anti-apoptotic pathways (Chen et al. 1998; McCole et al. 2000). During later stages of the infection cycle, pro-apoptotic mediators dominate and the host cell dies (McCole et al. 2000). Indeed, studies found that C. andersoni (a Cryptosporidium species predominantly in post-weaned and adult cattle) increased enterocyte apoptosis in vitro (Buret et al. 2003). Additionally, microarray analysis of gene expression of C. hominis and C. parvum infected ileal tissue explants revealed that the NF-kB-regulated osteoprotegerin (OPG) gene is upregulated (Castellanos-Gonzalez et al. 2008). OPG, a member of the tumor necrosis receptor superfamily, is released from intestinal epithelial cells and functions as a soluble decoy receptor, preventing the pro-apototic effects of tumor necrosis factor-related apoptosis inducing ligand (TRAIL) (O'Hara and Chen 2011). Treatment of a Cryptosporidium-infected cultured intestinal cell line with TRAIL induced apoptosis and decreased parasite numbers. However, the addition of recombinant OPG antagonized TRAIL-induced parasite reduction, suggesting both TRAIL-mediated regulation of parasite viability and a novel role for OPG in modulating apoptosis during the early phase of infection (Castellanos-Gonzalez et al. 2008). As described above, Cryptosporidium activation of TLR4 and TLR2 on the surface of biliary epithelial are required for NF- κ B activation, which induces

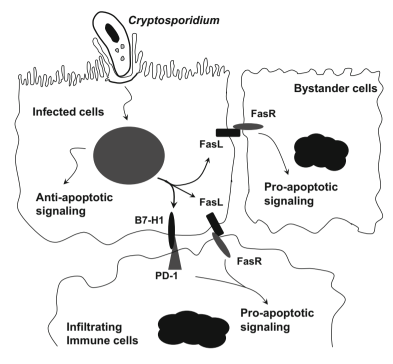


Fig. 10.4 Modulation of pro- and anti-apoptotic signals in various cell types at infection sites during *Cryptosporidium* infection. *C. parvum* reveals the ability to either induce or inhibit enterocyte apoptosis. The activation of NF- κ B and the antagonism between anti- and pro-apoptotic mechanisms both promote early parasite propagation, but is a likely mechanism for parasite eradication. Induction of both FasL and B7-H1 may be involved in *C. parvum*-induced apoptotic cell death in bystander un-infected cells, including infiltrating inflammatory cells, and contributes to parasite evasion of host immune and defense

anti-apoptotic factors, but also induces the expression of proinflammatory mediators (Laurent et al. 1997; Chen et al. 2005b) and β -defensins (Tarver et al. 1998), an antimicrobial peptide that is likely to be cytopathic to zoites (Chen et al. 2005b). Therefore, evidence supports that the activation of NF-kB and the antagonism between anti- and pro-apototic mechanisms both promote early parasite propagation, but may function as a mechanism for parasite eradication.

Clearing the pathogen infected cells through apoptosis is usually facilitated by infiltrating inflammatory cells. *C. parvum* may induce apoptotic resistance in infected host epithelial cells through upregulation of B7-H1, thus preventing killing by infiltrating inflammatory cells (Fig. 10.4). B7-H1 is a key member of the B7 family of costimulatory molecules which provide signals for both stimulating and inhibiting T cell activation (Dong et al. 1999; Gong et al. 2010). Upregulation of B7-H1 has been reported in a variety of infectious diseases, including parasitic diseases (Dong and Chen 2006; Joshi et al. 2009). Recent studies indicate that B7-H1 itself acts as a receptor and can respond to PD-1 and B7-1 (Azuma

et al. 2008; Keir et al. 2008). B7-H1 has been demonstrated as a ubiquitous antiapoptotic receptor on cancer cells: B7-H1 on cancer cells receives a signal from PD-1 to rapidly induce resistance against T cell-mediated killing (Azuma et al. 2008). Interestingly, *C. parvum* induces B7-H1 expression in host epithelial cells both in vitro and in vivo (Gong et al. 2010). Whether infiltration of inflammatory cells contributes to apoptotic resistance in infected epithelial cells through interactions between B7-H1 and PD-1 is yet to be investigated.

It is important to note that C. parvum-induced apoptosis in vitro and in vivo may include uninfected bystander cells, in particular, during the early stage of infection (Liu et al. 2009). The Fas (APO-1 and CD95)/Fas ligand (FasL) system is an important cellular pathway that regulates the induction of apoptosis in a variety of tissues (Matter et al. 2006; Juric et al. 2009). Fas is a widely expressed, 45-kDa type I membrane protein of the TNF-nerve growth factor family of cell surface receptors. By activating a variety of downstream effector cellular proteases, including members of the caspase family, Fas induces apoptosis (Cryns and Yuan 1998). The biological importance of the Fas/FasL system has been extensively studied in T cells, where it plays a critical role in the clonal deletion of autoreactive T cells and in the activation-induced suicide of T cells (Cryns and Yuan 1998). It was reported, using an in vitro model of biliary cryptosporidiosis, that C. parvum stimulates FasL membrane surface translocation, increases FasL protein expression in infected biliary epithelia, and induces a marked increase of soluble FasL in supernatants from infected cells. Infected epithelial cells can induce apoptotic cell death in co-cultured Fas-sensitive Jurkat cells, which can be blocked by a neutralizing antibody to FasL and a metalloprotease inhibitor (Chen et al. 1999). Besides FasL, induction of B7-H1 in infected epithelial cells may also play a role in apoptosis in the bystander un-infected cells. PD-1, the receptor for B7-H1, is a CD28/CTLA-4 like molecule expressed on T cells, B cells and macrophages (Ishida 1992; Freeman 2000). As described above, C. parvum induces B7-H1 expression in host epithelial cells both in vitro and in vivo. Infected human biliary epithelial cells can induce apoptotic cell death in co-cultured activated human T cells, which can be partially blocked by a neutralizing antibody to B7-H1 (Gong et al. 2010). Therefore, induction of both FasL and B7-H1 may be involved in C. parvuminduced apoptotic cell death in bystander un-infected cells, including infiltrating inflammatory cells, and contributes to parasite evasion of host defense (Fig. 10.4).

Intestinal epithelial cells infected with *C. parvum* were shown to be less susceptible to activation by IFN- γ , at least in part due to parasite induced depletion of the signal transducer and activator of transcription 1 (STAT1) protein (but not STAT1 mRNA depletion) (Choudhry et al. 2009). Moreover, the depletion of STAT1 did not involve protein ubiquitination (Choudhry et al. 2009). Whether miRNA-mediated post-transcriptional suppression plays a role in STAT1 depletion is unclear. Of note, miR-146a targets the 3'UTR of STAT1 mRNA and has been demonstrated to regulate levels of STAT1 in various cell types in response to LPS stimulation (Tang et al. 2009). However, expression of miR-146a remains unchanged in biliary epithelial cells following *C. parvum* infection (Zhou et al. 2009). Given that the IFN- γ response is critical for host defense against *C*.

parvum, the parasite could benefit from the suppression of activation of epithelial cells in response to IFN- γ in infected cells.

10.9 Immunopathology of Cryptosporidiosis

10.9.1 Pathogenic Mechanisms

Intestinal epithelia serve as an absorptive/secretory surface required for nutrient acquisition, and electrolyte and water transport. Also, the epithelium functions as a protective barrier between the underlying tissues and the potentially pathogenic contents of the lumen. If either of these functions is altered, pathological consequences ensue; both are altered during cryptosporidiosis. The intestinal epithelium has a remarkable capacity for fluid and electrolyte absorption under the tight control of neurotransmitters, hormones, inflammatory mediators, and intraluminal factors (Berkes et al. 2003). Chloride secretion, involving the concerted effort of several transporters (Berkes et al. 2003), is the principal determinant of luminal hydration. With chloride secretion, paracellular movement of sodium follows providing an osmotic gradient for the diffusion of water. Chloride hypersecretion has been reported in cryptosporidiosis, and represents one pathophysiological event responsible for diarrhea (Argenzio et al. 1994; Troeger et al. 2007). Compounding this effect, cryptosporidia cause malabsorption of sodium, nutrients and water, and accelerate intestinal transit (Argenzio et al. 1990; Buret et al. 1992; Deselliers et al. 1997).

Epithelial cells maintain the intestinal barrier by forming apical junctional and adherent complexes between enterocytes. These junctions result from the association of various transmembrane and cytosolic plaque proteins, and several cytosolic regulatory proteins (Gonzalez-Mariscal et al. 2003; Turksen and Troy 2004). Intact epithelial junctions are critical to normal physiological function and prevention of diseases. The disruption of intestinal tight junctions results in a "leaky" tight junction barrier, allowing the paracellular translocation of toxic luminal substances, food antigens, and luminal microbes (Baker et al. 2003; O'Hara and Buret 2008). Indeed, impaired intestinal barrier function has been implicated in the pathophysiology of a variety of enteric infections, and chronic gastrointestinal disorders (Farquhar and Palade 1963; Ukabam et al. 1983; Hollander 1988; Furuse et al. 1996; Clayburgh et al. 2004; Dokladny et al. 2006; O'Hara and Buret 2008).

Cryptosporidiosis impairs epithelial barrier function. Epithelial tight junctions are transiently disrupted during cryptosporidial infection (Zhang et al. 2000; Buret et al. 2003; Huang and White 2006). Further research is needed to assess whether these alterations in vivo are initiated by parasitic products, the invasion process itself, or host inflammatory responses to *Cryptosporidium*. However, in the absence of other host cells, infection of bovine and human epithelial cell lines with *C. andersoni* altered the distribution of tight junctional zonula occludens 1 protein, and

promoted apoptosis, consistent with a direct pathogenic effect of the parasite (Buret et al. 2003). Treatment of intestinal cells with TGF- β 1 inhibits *C. parvum*-induced permeability (Roche et al. 2000).

At the structural level, cryptosporidiosis is associated with crypt hyperplasia, villous atrophy, and a diffuse shortening or loss of brush border microvilli (Argenzio et al. 1990; Farthing 2000; Huang and White 2006). Hence, the absorptive capacity of the villi surface is diminished. This imbalance of absorption and secretion is likely a major contributor to disease manifestation. While the precise mechanism of diffuse villus blunting remains elusive, it is possible, given the tissue alterations, that activated CD8⁺ T-cells play a significant role as seen with giardiasis (Faubert 2000; Eckmann 2003; Scott et al. 2004). Physiologically, cryptosporidiosis is associated with reduced glucose-NaCl absorption, increased chloride anion secretion (Cl⁻), and decreased epithelial barrier function, all of which contribute to dehydration, diarrhea, and malabsorption associated with cryptosporidiosis. Of significant interest, expression of substance P, a neuropeptide that localizes to areas of inflammation and stimulates active Cl⁻ secretion from intestinal epithelial cells, is elevated in experimentally infected human subjects, and in AIDS patients with uncontrolled cryptosporidiosis. Expression levels of this neuropeptide seem to correlate with severity of diarrhea (Robinson et al. 2003). Furthermore, a molecular antagonist of the substance P receptor (NK1) reduced both ion secretion and corrected the *Cryptosporidium*-induced glucose malabsorption in a macaque intestinal explant model of cryptosporidiosis (Hernandez et al. 2007).

10.9.2 Chromatin Dynamics and Cryptosporidium Infection

Epigenetic regulation refers to reversible modifications introduced by various chromatin modifying proteins on DNA and histories to regulate gene expression via modulating chromatin structures (Graeff et al. 2011). In general, DNA methylation is associated with transcriptional repression while histone acetylation is related to transcriptional activation. Depending on the lysine residue methylated, histone methylation can either activate or repress gene transcription (Cloos et al. 2008). Dynamic epigenetic regulation participates in modulating almost all important biological processes including cellular differentiation. Gene expression in eukaryotes is regulated by histone acetylation/deacetylation, an epigenetic process mediated by histone acetyltransferases (HATs) and histone deacetylases (HDACs) whose opposing activities are tightly regulated. The acetylation of histones by HATs increases DNA accessibility and promotes gene expression, whereas the removal of acetyl groups by HDACs has the opposite effect. Epigenetic modifications protect against the excessive response and autotoxicity of acute pro-inflammatory gene products (McCall et al. 2010). Evidence has accumulated showing that HDACs have immunomodulatory activity and are important to regulation of anti-microbial defense (Roger et al. 2011). Multiple reports have shown that HDAC inhibitors possess suppressive effects on the induction of immune response genes (Bode et al. 2007; Roger et al. 2011). Therefore, it is not surprising that distinct epigenetic mechanisms regulate the inflammatory responses and contribute to the pathogenetic features in many infectious diseases (Paschos and Allday 2010).

Epigenetic mechanisms may be involved in C. parvum-induced alteration of miRNA transcription in infected cells. Among those miRNAs suppressed in host epithelial cells following C. parvum infection, the let-7i gene is suppressed through histone-H3 deacetylation (O'Hara et al. 2010b). Analysis of the mechanism revealed that microbial infection promotes the formation of an NF-KB p50-C/ EBPβ silencer complex in the *let-7i* regulatory sequence. Chromatin immunoprecipitation assays demonstrated that the repressor complex binds to the let-7i promoter following C. parvum infection and promotes histone-H3 deacetvlation. Suppression of the *mir-424-503* gene following *C. parvum* infection also involves epigenetic mechanisms (Zhou et al. 2013). C. parvum-induced transcriptional suppression of the mir-424-503 gene is associated with an increased recruitment of NF-kB p50 to the promoter. Intriguingly, promoter recruitment of several HDACs, including HDAC1, HDAC2 and Sirt1, was also detected in infected cells. Such promoter recruitment of NF-KB p50 and HDAC complex appears to be associated with a decrease of H3 acetylation and account for the repression of the mir-424-503 gene in infected cells. Functionally, induction of CX3CL1 may be associated with downregulation of mature miR-424 and miR-503, both of which target the CX3CL1 3'UTR, suppress its translation and induce RNA degradation (Zhou et al. 2013).

Our understanding of epigenetic regulation of the host response to infection is advancing, but studies on the role of epigenetic modifications during *Cryptosporid-ium* infection are still very limited. Of particular interest, epigenetic processes may impact the metabolic functions in enterocytes and thus play a role in the pathogenicity of *Cryptosporidium* infection. Epigenetic changes may also impact mucosal immunity and determine the severity of infection.

10.10 Conclusion and Perspectives

While we have begun to recognize the critical role for mucosal epithelial cells in the immune defense against *Cryptosporidium* infection, how epithelial cells are activated to mount an efficient innate and adaptive anti-parasite immune response is still unclear. The parasite-epithelial cell interactions are very complex. PRRs expressed in epithelial cells, such as TLRs, and their downstream signaling pathways, appear to be the key effector regulators accounting for epithelial cell defense, including production and release of a panel of effector molecules of anti-microbial activity or attracting the infiltration of inflammatory immune cells to infection sites. Both transcriptional and post-transcriptional mechanisms are involved in the regulation of such epithelial responses, in particular, miRNA-mediated post-transcriptional gene suppression. *Cryptosporidium* parasites have

also developed strategies of immune evasion to escape host defense, at least at the early stage of infection. Despite significant advances in understanding mucosal anti-*Cryptosporidium* immunity, how parasite antigens are presented through the mucosal epithelium and how adaptive immunity is regulated remains unclear.

Identification of parasite molecules involved in the activation of signaling pathways in infected host cells would provide new insights into the molecular mechanisms of immune activation following Cryptosporidium infection. Evaluating the secretion and function of these newly-identified cytokines secreted by epithelial cells will provide additional insights into how epithelial cells mediate anti-Cryptosporidium defense. Cell-cell communications mediated by exosomes may have a much broader impact on the innate and adaptive immune network. In particular, these mechanisms may recruit infiltrating inflammatory cells to infection sites, and may represent a new target for therapeutic intervention. The development of epithelial cell-specific knockouts will greatly advance our understanding of epithelial-mediated immune responses in vivo. Unraveling the regulatory functions of ncRNAs in anti-parasite immunity is still in its infancy, but will likely yield new insights into our understanding of the immunobiology and immunopathology of cryptosporidiosis. Given the importance of adaptive immunity to clear infection and the intramembranous localization of the parasite in epithelial cells after invasion, modulation of epithelial innate immune defense through lympho-epithelial interactions may be critical for anti-Cryptosporidium immunity, an understudied area of Cryptosporidium research. Better understanding of the impact of epigenetic modifications on the immune response will help us to evaluate the potential impact of cryptosporidial infection on malnutrition and possible co-infections identified in young children.

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Chapter 11 Treatment of Cryptosporidiosis

Jan R. Mead and Michael J. Arrowood

Abstract *Cryptosporidium* species are protozoan parasites that infect the epithelial cells of the gastrointestinal tract. In humans, infections occur primarily in the small intestine resulting in diarrheal illness. *Cryptosporidium* has a worldwide distribution and is considered an emerging zoonosis. Despite intensive efforts to develop workable experimental models, and the evaluation of nearly 1,000 chemotherapeutic agents, adequate therapies to clear the host of these parasites are still lacking. The reasons for the lack of drug efficacy are probably manifold and may include the unusual location of the parasite in the host cell, distinct structural and biochemical composition of drug targets, or its ability to either block import or rapidly efflux drug molecules. Understanding the basic mechanisms by which drugs are transported to the parasite and identifying unique targets are important steps in developing effective therapeutic agents.

11.1 Introduction

Therapeutic agents and treatment strategies for cryptosporidiosis have been pursued for over 35 years since *Cryptosporidium* was first identified in humans (Nime et al. 1976). Even though promising targets and lead compounds have been identified, robust therapies that clear the host of these parasites are still lacking. Particular groups would benefit from an effective therapy against cryptosporidiosis:

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children, the elderly, and immunocompromised individuals including organ transplant recipients, patients undergoing cancer chemotherapy, patients with congenital or disease-induced immunodeficiencies, and especially HIV-infected individuals. Even if not fully curative, many patients would benefit if drug therapy only reduced the parasite burden. Chemotherapy would also be useful among immunocompetent populations in outbreak situations to curb the spread of disease. While long recognized as one of the most ubiquitous and frequent cause of protozoal diarrhea in humans (Huston and Petri 2001), the recent Global Enteric Multicenter Study (GEMS) of children under 5 years old in developing countries found *Cryptosporidium* to be among the top 4 causes of moderate-to-severe diarrhea and that such diarrhea is a "high risk factor for linear growth faltering and death" (Kotloff et al. 2013).

Among the more important groups in need of therapy are individuals infected with human immunodeficiency virus (HIV). While cryptosporidiosis can be serious in immunocompetent people, it can be devastating to those with AIDS. In these individuals, symptoms may include chronic or protracted diarrhea that can become life threatening. Infections among HIV-infected individuals may also become extra-intestinal, spreading to other sites including the gall bladder, biliary tract, pancreas and pulmonary system (Lopez-Velez et al. 1995). Drug development should take into consideration the location of the parasite within the host (intestinal vs. extraintestinal sites) and the unique relationship between the parasite and host including location in the host cell and development of asexual and sexual life cycle stages.

The introduction and widespread use of highly-active anti-retroviral therapy (HAART) in AIDS patients has resulted in a decrease in opportunistic infections, however many adults and children living with HIV/AIDS in sub-Saharan Africa are currently not being treated with ART. Prevalence of Cryptosporidium among HIV-positive children with diarrhea has been reported to range between 13 % and 74 % in sub-Saharan Africa (Mor and Tzipori 2008). An early study of the impact of HAART on AIDS-defining illnesses in HIV-infected patients noted a 60 % decrease in the incidence of cryptosporidiosis (Ives et al. 2001). It is not known, however, how long these effective treatments will persist since multi-drug resistance and severe side effects associated with protease inhibitors may result in rebounding viral loads and, ultimately, increases in opportunistic infections. Despite intensive efforts, workable experimental models and testing of hundreds of chemotherapeutic agents, adequate therapies to clear the host of these parasites are lacking. Unique characteristics of Cryptosporidium species and the cryptosporidial life cycle in the host may account for the failure of most drug treatments tested to date. These include unique biochemical pathways (drug targets), lack of potential drug targets, unique or selective transporters functioning in the parasitophorous vacuole, or other features unique to the location of the parasite in the host cells (Chap. 2, Fig. 2.1) that facilitates sequestration of the parasite from drugs or other therapies.

11.2 Clinically-Evaluated Therapeutics

Controlled treatment trials have provided useful information about the activity of potential anti-cryptosporidial agents, but have not resulted in the identification of a robustly effective therapeutic agent. While many compounds have demonstrated anti-cryptosporidial activity using in vitro assays, fewer therapeutic agents have demonstrated significant potency using animal models. Further, many case reports describe attempts to treat cryptosporidiosis, but provide limited statistical value in aggregate analyses. The therapies most widely tested in animal models, case reports, and clinical trials are summarized below.

11.2.1 Macrolide Antibiotics

Some of the first drugs to be tested against *Cryptosporidium* species were macrolide antibiotics. Spiramycin, evaluated in the mid to late 1980s, produced encouraging results in pilot studies but was found to have little efficacy in larger, controlled trials and exhibited potentially unacceptable toxicity (Portnoy et al. 1984; Saez-Llorens et al. 1989; Wittenberg et al. 1989). Indeed, escalating high dose i.v. spiramycin treatment of cryptosporidiosis in two AIDS patients led to worsening of intestinal symptoms without significantly impacting cryptosporidial infection. Upon discontinuation of treatment one patient continued to deteriorate, while the second showed worsening diarrhea although gross and histological improvement was observed in upper and lower intestinal biopsies (Weikel et al. 1991).

A newer macrolide, azithromycin, was evaluated with mixed results. Azithromycin treatment resulted in symptomatic relief in immunosuppressed children and in patients with chronic cryptosporidiosis (Russell et al. 1998; Vargas et al. 1993). When given at a dose of 500 mg daily to patients with AIDS it was reported to be both effective (Kadappu et al. 2002) or ineffective (Blanshard et al. 1997). Extraintestinal infections (respiratory) in immunocompromised (HIV) patients responded positively to azithromycin alone or in combination with paromomycin (Palmieri et al. 2005; Tali et al. 2011). Apparent parasitologic cure was achieved in a liver transplant patient with cryptosporidial cholangitis after combination treatment with azithromycin and paromomycin, despite ongoing immunosuppression with tacrolimus (Denkinger et al. 2008). Combination treatment using azithromycin and paromomycin of an ileum transplant patient infected with C. hominis resulted in transient disappearance of cryptosporidia in biopsy samples and a cessation of clinical signs. However, immunosuppressive therapy was maintained and the patient became infected with C. parvum within 3 months and, despite additional treatment regimens including pyrimethamine-sulphadiazine and paromomycin, intermittently shed oocysts over the next 3 years (Pozio et al. 2004). Repeated treatment with azithromycin and paromomycin failed to eradicate cryptosporidiosis in a patient with X-linked hyper-IgM syndrome

(an immunodeficiency disease resulting from a mutation in the CD40L gene) (Rahman et al. 2012).

Clarithromycin demonstrated modest efficacy as a prophylactic drug. The prevalence of cryptosporidiosis in HIV-infected patients was reduced when patients received clarithromycin daily, suggesting a protective effect (Holmberg et al. 1998; Jordan 1996). In a contrasting study, rifabutin, but not clarithromycin, decreased the risk of developing cryptosporidiosis in HIV-infected patients (Fichtenbaum et al. 2000). Therapeutic efficacy was reported with clarithromycin (Fujikawa et al. 2002) and with roxithromycin (Uip et al. 1998; Sprinz et al. 1998). Symptomatic improvement and resolution of oocyst shedding in an AIDS patient followed a 3 week course of clarithromycin (800 mg/day), although CD4⁺ counts concomitantly rose from 152 to 180/µL perhaps indicating immune responsiveness played an important role (Fujikawa et al. 2002). In an uncontrolled study, patients received oral roxithromycin (300 mg b.i.d.) for 4 weeks. Out of the 22 patients that completed the study, 15 patients (68 %) were considered to be cured, six patients (27 %) improved, and one patient failed treatment (5 %) (Uip et al. 1998). In another uncontrolled study of 24 AIDS patients with cryptosporidiosis, treatment with roxithromycin (300 mg b.i.d.) for 4 weeks produced symptomatic improvement of diarrhea in 79 % of cases, with 50 % of patients achieving complete response (Sprinz et al. 1998).

11.2.2 Aminoglycoside Antibiotics

Paromomycin has been one of the most widely used agents to treat cryptosporidial infections in AIDS patients. It is a poorly absorbed aminoglycoside that is related to neomycin and kanamycin and achieves high concentrations in the gut, in part due to poor bio-availability. Paromomycin has shown efficacy in animal models (Healey et al. 1995; Tzipori et al. 1995) and as a prophylactic treatment in neonatal calves (Fayer and Ellis 1993b), lambs (Viu et al. 2000) and goats (Johnson et al. 2000; Mancassola et al. 1995).

Patients have reported positive responses to oral paromomycin treatment with doses ranging from 1,500 to 2,000 mg/day (Flanigan and Soave 1993). The drug was considered partially effective in that decreases in symptoms (frequency of stools) were noted but the parasite was not eradicated. From 1990 to 1996, 11 studies reported an overall response rate of 67 % (Hewitt et al. 2000). In a placebo-controlled, double blind study, treatment was found to be partially effective (White et al. 1994). Ten patients with AIDS and cryptosporidiosis were randomized to paromomycin (500 mg t.i.d. or q.i.d.) or placebo in a double-blind trial. After 14 days, patients were switched to the other treatment for 14 additional days. During the paromomycin treatment phase, oocyst excretion decreased, suggesting clinical improvement in the treated group. However, the parasite load remained high in some patients with disseminated biliary tract infections. In another study in which AIDS patients were treated with paromomycin (500 mg q.i.d.), 19 out of

25 people had improvement in clinical symptoms but in only six cases was the parasite eliminated (Stefani et al. 1996). An NIAID multi-site study was initiated in 1995. Thirty-five adults with $CD4^+$ cell counts of $<150/\mu$ L were enrolled through the AIDS Clinical Trials Group. Initially, 17 patients received paromomycin (500 mg q.i.d.) and 18 received matching placebo for 21 days. Then all patients received paromomycin (500 mg q.i.d.) for an additional 21 days. The results of this study were disappointing, showing no significant difference between the treated and placebo groups (Hewitt et al. 2000). Three paromomycin recipients (17.6 %) versus two placebo recipients (14.3 %) responded completely. Rates of combined partial and complete responses in the paromomycin arm (8 out of 17, 47.1 %) and the placebo arm (5 out of 14, 35.7 %) of the study were similar (P = 0.72). Consequently, many patients initially respond to paromomycin treatment with a decrease in diarrhea but then relapse. A statistical meta-analysis of data from two studies (Hewitt et al. 2000; White et al. 1994) found no evidence that paromomycin was effective in reducing the frequency of diarrhea nor in achieving parasitological cure (Abubakar et al. 2007). Nevertheless, paromomycin as a therapeutic agent is routinely used as a positive control in animal studies and in vitro (Graczyk et al. 2011; Benitez et al. 2007; Umejiego et al. 2008; Klein et al. 2008).

11.2.3 Benzeneacetonitrile Antibiotics

The benzeneacetonitrile derivatives diclazuril and letrazuril have been evaluated in clinical trials. Diclazuril was evaluated in a double-blind, placebo-controlled study in which patients were given 50–800 mg/kg daily (Soave et al. 1990). No significant differences were observed between the diclazuril and placebo-treated groups. Letrazuril, a derivative of diclazuril, showed promise in a pilot study. At a dosage of 150–200 mg daily, letrazuril was associated with an improvement in symptoms in 40 % of patients treated and cessation of excretion of cryptosporidial oocysts in the stool in 70 % (Blanshard et al. 1997). However, biopsies from these patients remained positive. Letrazuril also demonstrated modest activity in clinical trials (Harris et al. 1994). Of 14 patients evaluated, five showed a major response (symptomatic improvement and eradication of cryptosporidial oocysts from the stool), two had a minor response (symptomatic improvement with persistence of oocysts in stool), and seven had no response to therapy with letrazuril. A 1997 review concluded diclazuril and letrazuril had modest anticryptosporidial activities and were not very promising since patients experienced little clinical improvement (Blagburn and Soave 1997).

11.2.4 Immunotherapeutic Treatment (Colostrum)

There has been a great deal of interest in the potential of immunotherapy for cryptosporidiosis over the years. The earliest reports of immunotherapy involved individual cases where treatment with immune or hyperimmune bovine colostrum was associated with both success as well as failure (Jenkins 2004; Kelly 2003; Crabb 1998). These included a child with hypogammaglobulinemia (Tzipori et al. 1986), and patients with either hypogammaglobulinemia (Saxon and Weinstein 1987) or AIDS (Nord et al. 1990; Saxon and Weinstein 1987; Ungar et al. 1990). Reduction in parasite load was observed in animals given immune colostrum (Fayer et al. 1989; Perryman and Bjorneby 1991). In an observational study in Nigeria, HIV-associated diarrhea was alleviated by a 4 week treatment regimen with a commercial bovine colostrum product (Floren et al. 2006). Among a 30 patient group with diarrhea, 20 had cryptosporidiosis. The study did not differentiate which patients, on average, benefited most from colostrum treatment, nor whether parasite shedding ceased, but average daily bowel movements decreased from 7.0 \pm 2.7 (s.d.) to 1.6 \pm 0.9 (s.d.) after 4 weeks of treatment (week 5 of the study). Interestingly, CD4⁺ cell counts increased from an average of 153 ± 62 (s.d.) cells/ μ L to 310 \pm 106 (s.d.) cells/ μ L at week 7 of the study which the authors attribute, in part, to the colostrum product since ART was not initiated until the end of week 7.

Several open label studies using hyperimmune bovine colostrum (HBC) to treat AIDS patients with cryptosporidiosis have been reported. Patients treated with 48 enteric-coated capsules (40 total grams) per day over a 21-day period showed decreases in mean stool weight and stool frequency. Unfortunately, parasite load was not evaluated in this study, so HBC's direct effect upon parasite growth could not be determined (Greenberg and Cello 1996).

HBC was also used in a placebo-controlled, double-blind, one-way cross-over study in which AIDS patients were treated for 1–2 weeks with 20 g/day followed by increasing doses to 80 g/day in some patients. No statistical differences were observed in clinical symptoms but a reduction in oocyst shedding was reported (Fries et al. 1994). The prophylactic effect of hyperimmune bovine anti-Cryptosporidium colostrum immunoglobulin (BACI) was evaluated in healthy adults challenged with C. parvum (Okhuysen et al. 1998). Sixteen volunteers were randomized to receive either a (1) BACI prior to C. parvum challenge and a nonfat milk placebo 30 min later, (2) BACI prior to and 30 min after challenge (reinforced BACI group), or (3) nonfat milk placebo prior to and 30 min after challenge. Subjects received BACI (10 g) or nonfat milk placebo three times a day for a total of 5 days and were monitored for clinical symptoms and oocyst excretion for 30 days. Subjects receiving BACI or nonfat milk placebo had a 100-fold reduction in oocyst excretion as compared with excretion in the baseline group, however no difference was observed between the BACI and nonfat milk placebo treatment groups. In terms of clinical symptoms, no significant differences were observed in the duration of disease, time to onset of diarrhea, and severity of the disease among groups.

11.2.5 Thiazolide Antibiotics

Nitazoxanide (NTZ), a thiazolide drug with reported broad antiparasitic activities, is currently the only FDA-approved drug for use against cryptosporidiosis in immunocompetent patients. Early studies demonstrated moderate efficacy in cell culture systems and in neonatal mice and gnotobiotic piglets (Blagburn et al. 1998; Gargala et al. 2000; Theodos et al. 1998). Duration of oocyst shedding and severity of diarrhea in experimentally-infected newborn calves was reduced following treatment with NTZ (1,500 mg b.i.d.) (Ollivett et al. 2009). Calves received colostrum on day 1, were infected between days 2 and 3 and NTZ treatment began between days 5 and 7 (to allow infections to become established) and continued for 5 days.

Although early clinical trials in the U.S. were not encouraging, one uncontrolled open study in Mali suggested that the drug was efficacious against the parasite in stage 4 AIDS patients with "light" cryptosporidial infections, but failed in "heavily infected" stage 4 AIDS patients (Doumbo et al. 1997). In a double-blind placebo-controlled study, patients were randomly treated orally with either 500 or 1,000 mg (b.i.d.) NTZ or placebo for 14 days (Rossignol et al. 1998). Patients on NTZ then crossed over to placebo while the placebo patients crossed over to NTZ therapy at either the high or low dose depending on their randomization. Patients were considered 'cured' if oocysts were not found in post-treatment fecal examinations. Both doses of NTZ produced parasitological cure rates superior to the placebo responses (12/19 [63 %, P = 0.016] for patients receiving 1 g/d and 10/15 [67 %, P = 0.013] for those receiving 2 g/d). It should be noted that 25 % (5/20) of placebo-treated patients were parasitologically "cured" and 50 % (10/20) had complete resolution of clinical diarrhea before crossing over to nitazoxanide therapy.

An HIV positive patient with persistent diarrhea was sequentially treated with paromomycin, azithromycin, paromomycin with octreotide, and finally nitazoxanide, all of which failed to adequately control the cryptosporidial infection 1999). Interestingly, seven samples of the patient's (Giacometti et al. cryptosporidial isolate(s) were collected separately over a nearly 1.5 year period and tested for susceptibility to paromomycin, azithromycin and nitazoxanide in an in vitro cultivation system. Although some activity was noted for each drug, none completely inhibited parasite growth. Maximum inhibition for azithromycin was 26.5 % at 8 mg/L (10.7 µM), for paromomycin 63.4 % at 1 mg/L (1.6 µM), and for nitazoxanide 67.2 % at 10 mg/L (33.0 μ M). It was also noted that nitazoxanide was toxic for the cell cultures at the maximum concentration tested (100 mg/L (330 µM)). In a subsequent study by the same researchers, nitazoxanide alone at 8 mg/L (26 μ M) yielded an inhibition of parasite growth in vitro of approximately 55 %, but inhibition increased to approximately 80–84 % when combined with rifabutin at 8 mg/L (9.5 μ M) or azithromycin at 8 mg/L (10.7 μ M), respectively (Giacometti et al. 2000).

In a prospective, randomized, double-blind, placebo-controlled study, 50 adults and 49 children from the Nile delta of Egypt, were treated with nitazoxanide or placebo for diarrhea caused by *C. parvum*. Nitazoxanide was administered in 500 mg doses b.i.d. for 3 days in adults and adolescents, in 200 mg doses b.i.d. for 3 days in children aged 4–11 years, and in 100-mg doses b.i.d. for 3 days in children aged 1–3 years (Rossignol et al. 2001). Seven days after initiation of therapy, diarrhea resolved in 39 (80 %) of the 49 patients in the nitazoxanide treatment group, compared with 20 (41 %) of 49 in the placebo group (P < 0.0001). Diarrhea resolved in most patients receiving nitazoxanide within 3 or 4 days of treatment initiation. Nitazoxanide treatment reduced the duration of diarrhea (P < 0.0001) and oocyst shedding (P < 0.0001). When children were excluded, 72 % (18/25) of nitazoxanide-treated adults and adolescents resolved diarrhea while 44 % (11/25) of placebo-treated adults and adolescents resolved diarrhea; differences were not statistically significant.

Additionally, a prospective, randomized, double-blind, placebo-controlled study was performed in children with diarrhea. The efficacy and safety of nitazoxanide compared with placebo was assessed in the treatment of diarrhea caused by *C*. *parvum* in HIV-infected and HIV-uninfected Zambian children, most of whom were younger than 3 years and malnourished. A 3-day course of nitazoxanide significantly improved the resolution of diarrhea, parasitological eradication, and mortality in HIV-seronegative children. However, HIV-seropositive children did not benefit from nitazoxanide treatment (Amadi et al. 2002).

In an in vitro study, NTZ and tizoxanide were reportedly more active against sporozoites while tizoxanide glucuronide appeared to be more active against intracellular stages (Gargala et al. 2000). To improve the efficacy of this class of drugs, second generation compounds were generated. Among a group of 39 thiazolide and thiadiazolide compounds evaluated for efficacy compared to NTZ, several were identified as more potent than NTZ (Gargala et al. 2010). One of these compounds, RM-4865, demonstrated efficacy in a gerbil model and was equally as potent as NTZ in reducing oocysts in this model while RM-5038 was significantly more active than NTZ (Gargala et al. 2013).

The mode of action for NTZ against *Cryptosporidium* has not been determined and the following in vitro data are worth noting when interpreting the anticryptosporidial efficacy. These observations should be kept in mind when testing any agent for anticryptosporidial activity. NTZ has been reported to cause dose-dependent toxicity in cell cultures (Theodos et al. 1998; Giacometti et al. 1999; Gargala et al. 2000). A clear association between mammalian cell cycle and thiazolide toxicity (via induction of apoptosis) has also been reported (Müller et al. 2008). In the latter study, growth inhibition was pronounced for proliferating Caco2 cells (transformed) [IC₅₀ 6.8 \pm 1.3 μ M] and human foreskin fibroblasts (primary cell culture) [IC₅₀ 18.9 \pm 1.2 μ M], but much less so for confluent, non-proliferating cells. The proposed mode of action (toxicity) against the cell lines is via inhibition of glutathione-S-transferase P1. Since the mode of action for NTZ against *Cryptosporidium* is unknown, the toxicity/apoptosis exhibited by NTZ against rapidly proliferating cells (e.g. intestinal epithelium) may partially explain the capacity to reduce parasite numbers independent of a direct mode of action against the parasite itself.

11.2.6 Combination Therapy

There have been a number of studies over the years that used combination chemotherapy (some mentioned above). These include in vitro studies (Giacometti et al. 1996, 1999, 2001; You et al. 1998), in vivo studies (Fayer and Ellis 1993a; Kimata et al. 1991) and treatment given to a patient with disseminated cryptosporidiosis (Giacometti et al. 1999).

In an open-label, combination study by Smith and colleagues (Smith et al. 1998), patients with AIDS, chronic cryptosporidiosis, and low CD4⁺ cell counts (<100 CD4⁺ cells/ μ L) were given paromomycin (1.0 g b.i.d.) plus azithromycin (600 mg once daily) for 4 weeks, followed by paromomycin alone for 8 weeks. In 11 patients, median stool frequency decreased from 6.5/day (baseline) to 4.9/day (week 4) and 3.0/day (week 12). Median reductions in 24 h oocyst excretion were 84 %, 95 %, and >99 % at 2, 4, and 12 weeks, respectively. Of five survivors at 12–30 months follow-up, three remained asymptomatic off medications, and two had chronic, mild diarrhea. Both partial clinical and parasitological responses were observed, as treatment of cryptosporidiosis with azithromycin and paromomycin was associated with significant reduction in oocyst excretion and some clinical improvement. Unfortunately, no placebo group was included in the study.

The use of potent highly active anti-retroviral therapy (HAART) in patients with advanced HIV infection can improve symptoms or lead to the clearance of *C. parvum* oocysts from stools. Patients treated with double anti-retroviral therapy or protease inhibitors have demonstrated excellent responses and sustained therapeutic effects after follow-up (Ives et al. 2001; Miao et al. 2000). In other studies it was found that combination antiretroviral therapy that included a protease inhibitor restored immunity to *C. parvum* in HIV-1 infected individuals, resulting in complete clinical, microbiological, and histological responses (Carr et al. 1998). HAART-associated resolution of diarrhea is linked to increased CD4⁺ cell counts rather than to modulation of the viral load. Four patients had recurrent diarrhea at 7–13 months (one with positive stool microscopy), associated with declining CD4⁺ counts.

Although HAART is not thought to have a direct effect on *C. parvum*, there is evidence that the protease inhibitor, indinavir (IND) can reduce *C. parvum* infection in both in vitro and in vivo models (Bobin et al. 1998; Hommer et al. 2003; Mele et al. 2003; Miao et al. 2000).

A retrospective cohort study spanning 1999 through 2009 of 1,243 HIV⁺ patients in Morocco receiving HAART identified 91 deaths (Sodqi et al. 2012). Average CD4⁺ counts were 95 cells/ μ L at enrollment and initiation of HAART. Cryptosporidiosis was the second most common cause of death (19 %) among these patients (following tuberculosis at 35 %). Of those patients that died, HAART duration averaged 9 months and 63 % died within the first year of treatment. At the time of death, ~53 % had CD4⁺ counts above 200 cells/ μ L. Although some patients succumbed to cryptosporidiosis, it appears that prompt initiation of HAART and effective immune reconstitution had the most efficacious impact on surviving opportunistic infections, including cryptosporidiosis.

Looking historically at the range of drugs tested clinically against *Cryptosporidium*, it appears several drugs may have had promising initial assessments, not because the drugs were efficacious against the parasite itself, but rather because of their impact on the gut microflora and/or the epithelial (host) cells of the gastrointestinal tract.

11.3 Innate Drug Resistance

Why are so many therapies ineffective against this parasite? It seems that *C. parvum* has an innate resistance to drug therapy. Several factors may contribute to this lack of efficacy. These include: (1) lack of specific targets or differences in targets either at the molecular or structural levels compared to the host, (2) differences in biochemical pathways, (3) the parasite's unique location in the host cell which may affect drug concentration (transported from the lumen or host cell cytoplasm across to the parasite), and (4) existence of transport proteins or efflux pumps that move drugs out of the parasite to the lumen or back into the host cell.

11.3.1 Genomic, Proteomic, Metabolic Characteristics

The establishment of large scale sporozoite-expressed sequence tag (EST) and genome sequencing projects has aided the identification of *C. parvum* genes and in understanding phylogenetic similarities and diversities within the genome. It appears that *C. parvum* is more divergent and less related to other coccidia. Comparisons of small-subunit ribosomal RNA gene sequences first demonstrated that the gregarine/*Cryptosporidium* clade is separated from the other major apicomplexan lineages (Carreno et al. 1999; Leander et al. 2003). Another study compared six different proteins along with the SSU rRNA and concluded that *Cryptosporidium* is an early emerging branch of the Apicomplexa (Zhu et al. 2000a). Full genome sequencing of *C. parvum* demonstrated significant differences between cryptosporidia and coccidia (Abrahamsen et al. 2004), and though it reinforced the relatedness to gregarines, there are significant differences between cryptosporidia and gregarines. In particular, the streamlined metabolism of cryptosporidia is represented by the absence of the shikimate pathway, synthesis of

select amino acids, and *de novo* pyrimidine synthesis, all of which are commonly found in apicomplexans (Templeton et al. 2010). Nevertheless, cryptosporidia contain a number of metabolic targets in common with coccidia (and gregarines), including Type I fatty acid and polyketide synthesis enzymes (see Chap. 8).

Other structural differences have been noted. Unlike most of the Apicomplexa, C. parvum appears to lack a plastid genome (Riordan et al. 1999; Zhu et al. 2000b; Abrahamsen et al. 2004). Primers based upon the highly conserved plastid small- or large-subunit rRNA (SSU/LSU rRNA) and the tufA-tRNAPhe genes of other members of the phylum Apicomplexa failed to amplify products from intracellular stages of C. parvum grown in human adenocarcinoma (HCT-8) cells (Zhu et al. 2000b). The plastid (apicoplast) in other apicomplexan parasites has been shown to code for ribosomal proteins and tRNAs (Wilson and Williamson 1997) as well as metabolic pathways involved in Type II fatty acid synthesis and isoprenoid metabolism (Templeton et al. 2010). The activities of clindamycin and other macrolide antibiotics may be achieved through inhibition of protein synthesis within the plastid (Beckers et al. 1995; Fichera and Roos 1997). Clindamycin, an effective anti-Toxoplasma drug, expressed some in vitro anticryptosporidial activity (Woods et al. 1996) but was not efficacious in vivo (Rehg 1991). In general, macrolides have not demonstrated consistent anticryptosporidial efficacy even after long-term administration. While this lack of efficacy may or may not be related to the "missing" plastid, drug development targeting a plastid genome or metabolic pathways associated with it, may not be useful.

Despite the "missing" plastid and, presumably, the associated isoprenoid biosynthetic pathway (Clastre et al. 2007; Templeton et al. 2010), a nonspecific polyprenyl pyrophosphate synthase (CpNPPPS) was identified in C. parvum that is distinctly different from similar isoprenoid synthesis enzymes in other organisms (Artz et al. 2008). At least three prenyl synthase enzymes are coded for in the cryptosporidial genome. While downstream enzymes are present in Cryptosporid*ium* that can utilize substrates in the isoprenoid pathway, no evidence exists that cryptosporidia have either the mevalonate (MVA, e.g. human) or methylerythritol phosphate (MEP, see *Plasmodium* and *Toxoplasma*) pathways to generate these medium-length isoprenoid substrates (Clastre et al. 2007). It is unknown whether Cryptosporidium relies on an uncharacterized enzyme pathway to produce these substrates or whether they are acquired by a salvage pathway from the host cells. Interestingly, CpNPPPS was inhibited by sub-nanomolar concentrations of nitrogen-containing bisphosphonates and significant inhibition of in vitro parasite growth in MDCK cells was observed (Artz et al. 2008). Of the bisphosphonates tested, ibandronate showed the highest anticryptosporidial activity (IC₅₀ = $3.0 \,\mu\text{M}$) in cell culture.

Evidence suggests that *C. parvum* has a putative mitochondrion and mitochondrion-associated enzymes markedly different in structure from those of its nearest relatives (Riordan et al. 1999) but lacks much of the electron transport chain (Templeton et al. 2010). This may, in part, explain why atovaquone (targets cytochrome C) is not effective against *Cryptosporidium* (Rohlman et al. 1993; Giacometti et al. 1996; Mather et al. 2007).

Some parasite targets (e.g. enzymes, structural proteins) are different from those of other related parasites. For example, the dihydrofolate reductase (DHFR) gene of C. parvum differs from the homologous Plasmodium gene. Sequencing of the DHFR gene has suggested that the enzyme may be intrinsically resistant to 2,4-diaminopyrimidine inhibitors (Vasquez et al. 1996). The C. parvum DHFR active site contains novel residues at several positions analogous to those at which point mutations have been shown to produce antifolate resistance in other DHFRs (Vasquez et al. 1996). This may, in part, explain why C. parvum is resistant to clinically used antibacterial and anti-protozoal antifolates. An analysis of the crystal structure of C. hominis DHFR led to the development of novel trimethoprim (TMP) analogs with significantly enhanced (368-fold) inhibition of DHFR (TMP $IC_{50} = 14 \,\mu\text{M}$ versus 38 nM for analog # 37) in a cell free assay system (Pelphrey et al. 2007). Recent drug design efforts utilizing "virtual screening" of a non-active site pocket in the cryptosporidial DHFR linker region identified a noncompetitive inhibitor, flavin mononucleotide (FMN) (Martucci et al. 2009). As a first-generation lead compound, FMN exhibited an IC₅₀ = 55 μ M versus 14 μ M for TMP.

11.3.2 Location and Physiology

A factor that may contribute to the ineffectiveness of many drugs is the unusual location of the parasite in the host cell. The parasite has a unique niche inside the cell that is considered "intracellular" but "extracytoplasmic." The organism resides in a parasitophorous vacuole (PV) surrounded, in part, by host cell membrane but is outside the cytoplasm. It has been postulated that the PV basal membranes, including the "feeder organelle", modulate the transport of certain drugs, so that drugs entering the cytoplasm of the host cell may not be transported across to the parasite. In cell culture studies, apical but not basolateral exposure of these drugs led to significant parasite inhibition. There is evidence to suggest that this is the case for geneticin and paromomycin, a clinically-relevant drug (Griffiths et al. 1998). Alternatively, the parasite membranes into the PV, perhaps bypassing the host cytoplasm.

Another example where physiology may affect drug uptake was shown when the cryptosporidial inosine 5'-monophosphate dehydrogenase (CpIMPDH) gene was introduced into *T. gondii* under the control of a *T. gondii* promoter (Gorla et al. 2012). Many drugs were effective in this *Toxoplasma* model and were due to inhibition of the CpIMPDH (since the native *T. gondii* IMPDH gene was rendered inactive). However, these same drugs were generally not as active in a *C. parvum* in vitro cultivation assay suggesting reduced efficacy may be due to differences in the biology of the two parasites (Sharling et al. 2010). A phthalazinone-based inhibitor of CpIMPDH with promising efficacy and selectivity was not effective in an in vivo mouse model of cryptosporidiosis (Johnson et al. 2013). Location, structure, and function of PV from these two parasites is

clearly different: e.g. the PV membrane of T. gondii is in direct contact with the host cell cytoplasm whereas C. parvum remains beneath the apical membrane of the host cell and separated from the host cell cytoplasm by a feeder organelle (Sharling et al. 2010). Differences in drug efflux transporters between these parasites may also play a role in drug susceptibility.

11.3.3 ABC and Other Parasite Transporters

Transporters may also be involved in the rapid efflux of drugs as well as nutrient uptake. The existence of multi-drug resistant (MDR) transporters could facilitate drug resistance. P-glycoproteins and MDR transporters are members of the ATP-binding cassette (ABC) superfamily that are responsible for drug resistance by extruding drugs against a concentration gradient. In cancer chemotherapy, chemosensitizing agents are used to reduce efflux of chemotherapeutic agents from tumor cells by MDR pumps. To date, a set of 21 ABC genes has been identified in the fully sequenced C. parvum genome (Sauvage et al. 2009). Originally, three ABC transporters were identified in C. parvum. These include CpABCC3 (CpABC1) and CpABCC4 (CpABC2) that are similar to the multidrug resistance-associated (MRP) subfamily and CpABCB1 (CpABC3) that groups with the MDR subfamily (Zapata et al. 2002). The CpABCC3 gene shows similarity to human cystic fibrosis transmembrane receptor (CFTR) and to rat conjugate export pump protein (Strong and Nelson 2000) and is localized in sporozoites and at the boundary between host cells and the extracytoplasmic meront (Perkins et al. 1999; Zapata et al. 2002). CpABCC4 shows similarity to human CFTR and MRP-like protein from Candida albicans and has been identified in the apical end of sporozoites (Zapata et al. 2002). Gene expression of CpABCB1, an ABC transporter with similarity to MDR1, was detected in intracellular stages of C. parvum but its localization is unknown (Bonafonte et al. 2004; Zapata et al. 2002). These transporters are constitutively expressed in infected cell cultures, with transcript levels of CpABCC3 4 logs higher than either CpABCB1 or CpABCC4. In addition, MDR or MRP modulators (cyclosporine A, verapamil and probenicid) significantly affected transcript levels of all three of these ABC transporters in C. parvum (Bonafonte et al. 2004) and several sesquiterpenes have been shown to bind to the recombinant nucleotide-binding site of an MDR-like efflux pump when assayed on the recombinant nucleotide-binding domain of CpABCB1 (Lawton et al. 2007). While classic drug resistance has not been demonstrated with any of these ABC proteins, treatment by paromomycin, one of the most widely used agents to treat cryptosporidial infections in AIDS patients, significantly upregulates transcription of CpABCC14 (cgd7_4510) and of the CpABCB4 (cgd1_1350 or CpABC4 or C_p ATM1) (Benitez et al. 2007), a Pgp homolog half transporter. In order to characterize the catalytic site of this half-transporter, an extended region of the nucleotide-binding domain of CpABCB4 (H6-1350NBD) was expressed and purified as an N-terminal hexahistidine-tagged protein in E. coli. A dose-dependent quenching of the domain's intrinsic fluorescence was observed with the fluorescent analogue substrate TNP-ATP as well as the flavonoids quercetin and silibinin, previously shown to inhibit parasite development in a cell-based assay.

11.4 New Areas of Drug Research

A number of drug targets have been proposed over the years but have not been pursued either because the presumed drug target could not be found, toxicity of the drug was too great, or because the drug lacked efficacy in animal models or in clinical trials. Some of the following therapies remain promising because the target is selective or unique and efficacy has been demonstrated either in vitro or in vivo.

11.4.1 Flavonoids

Flavonoids, polyphenolic compounds found in plants, have demonstrated activity against several parasites and can augment the efficacy of other drugs by either increasing the uptake or decreasing the efflux of these drugs. The flavonoids apigenin and genistein inhibited C. parvum growth in vitro (Forney et al. 1999; Mead and McNair 2006). When the dinitroaniline drug trifluralin was tested in combination with the isoflavone genistein in an in vitro cultivation model the effective concentration that inhibited growth by 50 % (EC₅₀) dropped by 1 log (from 0.7 to 0.09 μ M), suggesting that isoflavone compounds may be used alone or in combination with other moderately active drugs to increase efficacy (Mead and McNair 2006). Flavonoids such as genistein have been shown to inhibit epidermal growth factor receptor protein tyrosine kinases (EGFR PTK), which are located at the surface of target cells and may be involved in the activation of the intracellular tyrosine kinase (TK) domain, initiating downstream signaling pathways. Synthetic analogs retaining the 5,7-dihydroxyisoflavone core of genistein were found to have in vitro activity against C. parvum (Stachulski et al. 2006). Two of these agents with demonstrable activity in vitro decreased oocyst shedding and limited C. parvum intracellular development in the gut and in the biliary tract using an immunosuppressed-gerbil model of cryptosporidiosis.

11.4.2 Bisphosphonates

The isoprenoid metabolic pathway is another potential drug target in parasitic protozoa including those causing malaria, Chagas' disease, toxoplasmosis, and leishmaniasis (Ling et al. 2005; Martin et al. 2001; Moreno et al. 2001). Bisphosphonates (used in bone resorption therapy) are known to inhibit the

biosynthesis of isoprenoids. In a mouse xenograft model of cryptosporidiosis (Moreno et al. 2001), the pyrophosphate analog drug risedronate was shown to inhibit *C. parvum* growth. Nitrogen-containing bisphosphonates (N-BPs) are capable of inhibiting *C. parvum* in infected MDCK cells at low micromolar concentrations (Artz et al. 2008). Crystal structures of the enzyme (nonspecific polyprenyl pyrophosphate synthase) with risedronate and zoledronate show how it could accommodate larger substrates and products.

In the process of screening a library of drug repurposing candidates (NIH Clinical Collections), researchers employing a high-throughput in vitro assay system identified itavastatin, an inhibitor of human 3-hydroxy-3-methyl-glutaryl-coenzyme A (HMG-CoA) reductase with potent in vitro activity against *C. parvum* (IC₅₀ = 0.62 μ M) (Bessoff et al. 2013). The investigators predict that the anticryptosporidial activity is mediated by the inhibition of host cell HMG-CoA, thus limiting the availability of isoprenoid precursors. *Cryptosporidium* species appear to lack the enzymes necessary for synthesis of isoprenoid precursors and depend on the host cell for these precursors. It remains to be seen if the parasite can scavenge sufficient precursors necessary for propagation under itavastatin drug pressure in an animal model or in clinical trials.

11.4.3 Polyamine Biosynthesis

It has also been determined that *C. parvum* differs fundamentally from other eukaryotes in polyamine metabolism. Mammals and most parasitic protozoa, including the coccidian parasites *E. tenella* and *P. falciparum*, convert arginine into ornithine. Ornithine is subsequently converted to putrescein via ornithine decarboxylase, and finally into spermidine and spermine (Bacchi and Narlett 1995). However, polyamine biosynthesis in *C. parvum* occurs via a pathway used by plants and certain bacteria, in which arginine is converted to agmatine by the action of arginine decarboxylase (Keithly et al. 1997). Neither arginine decarboxylase nor agmatine is found in other parasitic protozoa. Agmatine serves as the precursor for other polyamines.

C. parvum also has a reverse polyamine biosynthetic pathway not found in other protozoa that enables the interconversion of spermine, spermidine, and putrescein. Treatment with a polyamine analogue (SL-11047) prevented C. parvum infection in suckling TCR-alpha-deficient mice and cleared an existing infection in older mice. Treatment with putrescein, while capable of preventing infection, did not clear C. parvum from previously infected mice. Inhibitors of arginine decarboxylase (ADC), including difluoromethylarginine (DFMA), significantly reduced intracellular growth of C. parvum (Keithly et al. 1997), whereas inhibitors of ornithine had no effect upon ADC activity or upon growth of the parasite. Back-conversion of spermine spermidine and putrescein via spermidine:spermine-N1to acetyltransferase (SSAT) was also detected. Although ADC activity was

consistently detected in *C. parvum* parasites, a gene encoding an ADC homolog has not been identified in the completed genome sequence.

11.4.4 Fatty Acid Synthase

C. parvum possesses a unique Type I fatty acid synthase (*Cp*FAS1) and a putative polyketide synthase (*Cp*PKS1) encoded by 25-kb and 40-kb intronless open reading frames (ORFs), respectively (Zhu et al. 2002, 2000c) (see Chap. 8). Since *Cp*FAS1 and *Cp*PKS1 are structurally and functionally different from human Type I FAS, these enzymes may serve as novel drug targets. The fatty acid synthase gene (*Cp*FAS1) also differs from the organellar type II fatty acid enzymes identified in *Toxoplasma gondii* and *Plasmodium falciparum* (Zhu et al. 2000c). The FAS inhibitor cerulenin inhibited the growth of *C. parvum* in vitro by 96 % at 10 µg/mL, reinforcing the potential efficacy of FAS as a drug target.

11.4.5 Anti-tubulin Agents

Microtubules of various parasites differ from mammalian microtubules in a number of ways, including being relatively stabile and resistant to low temperatures. Antitubulin agents such as the benzimidazoles and the dinitroaniline herbicides have demonstrated efficacy against a number of parasitic agents (Roos 1997; Stokkermans et al. 1996; Traub-Cseko et al. 2001). Although benzimidazoles are widely used against helminth infections and some protozoans, C. parvum did not have the predicted amino acids for benzimidazole drug sensitivity in the betatubulin-coding gene (Edlind et al. 1994; Katiyar et al. 1994). In a subsequent study, benzimidazole treatment was not effective against C. parvum when evaluated in mice (Fayer and Fetterer 1995). Conversely, dinitroaniline herbicides were effective against C. parvum in vitro (Arrowood et al. 1996). The efficacy of this class of compounds has been determined in neonatal mice. At doses of 100 mg/ kg body weight administered twice daily for 3 consecutive days, trifluralin had no statistically significant effect on the number of oocysts recovered from the excised gut of either rats or mice compared with controls, whereas at the same concentration, oryzalin caused 90 % and 79 % inhibition of oocysts recovered from mice and rats, respectively (Armson et al. 1999). Treatment with oryzalin doubled the villus/ crypt ratio in the duodenum, jejunum and ileum following doses of 5 mg, 50 mg and 200 mg/kg respectively (Armson et al. 2002).

Dinitroanilines, as a rule, are poorly soluble which complicates their development as therapeutic agents. Synthetic derivatives with increased solubility were compared to parent compounds (Benbow et al. 1998; Mead et al. 2003). Acetylated glycoconjugate forms of dinitroaniline compounds were more active in vitro than non-acetylated forms, suggesting acetylation may facilitate drug transport across cell membranes (Mead et al. 2003). Although enhanced solubility did not necessarily correlate with better activity, several analogs were produced with similar anticryptosporidial activities but with toxicities lower than those of the lead compounds (Benbow et al. 1998). It may be possible that these compounds could be used at lower concentrations when combined with other efficacious drugs (Mead et al. 1999).

11.4.6 Dihydrofolate Reductase (DHFR) Inhibitors

Dihydrofolate reductase (DHFR), occurring as a bifunctional protein with thymidylate synthase (DHFR–TS) in apicomplexan protozoa, has been an established drug target for treatment of protozoal infections for many years. DHFR utilizes the cofactor NADPH to catalyze the reduction of dihydrofolate to tetrahydrofolate. In initial studies with *C. parvum*, standard dihydrofolate reductase (DHFR) inhibitors were not very effective, which was explained by nucleic acid sequences encoding resistant dihydrofolate reductase-thymidylate synthase (Vasquez et al. 1996). The sequence of *C. hominis* DHFR contains 3 amino acid mutations known to confer antifolate resistance in *P. falciparum* DHFR (*Pf*DHFR).

Several lipophilic DHFR inhibitors were reported to be active in a complementation assay but were nonselective (Brophy et al. 2000). Ninety-three lipophilic diand tricyclic diaminopyrimidine derivatives were evaluated in a cell-free enzyme assay for the ability to inhibit recombinant DHFR cloned from human and bovine isolates of *C. parvum* (Nelson and Rosowsky 2001). The library of compounds was also tested for anticryptosporidial activity in an in vitro cultivation assay using Madin-Darby canine kidney (MDCK) cells cultured in folate-free culture medium supplemented with thymidine (10 μ M) and hypoxanthine (100 μ M). Cell culture assays identified 16 compounds with IC₅₀s <3 μ M, of which five had IC₅₀s <0.3 μ M. Anticryptosporidal activities of the latter were comparable to trimetrexate.

The crystal structure showed that the *C. hominis* DHFR domain has a unique 9stranded rather than the 8-stranded β -sheet (O'Neil et al. 2003). Through protein: ligand interactions it was suggested that TMP did not extend deep enough into the hydrophobic pocket occupied by the natural substrate, dihydrofolate (Popov et al. 2007). A series of C7-TMP derivatives, designed to exploit a unique pocket in *Ch*DHFR was synthesized and evaluated. Modification of the TMP structure generated analogs with four times greater activity against *Ch*DHFR compared to TMP (Popov et al. 2006).

11.4.7 Inosine Monophosphate Dehydrogenase Inhibitors

Another potential target is the enzyme inosine 5-monophosphate dehydrogenase (*Cp*IMPDH), used for the production of guanine nucleotide. Through phylogenetic analysis, it was suggested that *Cryptosporidium* sp. obtained their IMPDH gene from an ε -proteobacterium by lateral gene transfer (Striepen et al. 2002). The bacterial origin of *Cp*IMPDH and the different kinetic properties of these enzymes compared with their human counterparts suggested *Cp*IMPDH-selective inhibitors could be obtained and, in fact, several different inhibitors active in the μ M range were identified in initial high throughput screens (Umejiego et al. 2008). All of these compounds were shown to bind to the nicotinamide portion of the NAD site suggesting that they stack up against the purine ring of IMP. Subsequent modifications of these initial hits, through medicinal chemistry, resulted in increased drug potency when evaluated in a *Toxoplasma* model expressing *Cp*IMPDH (Sharling et al. 2010).

11.4.8 Pyrimidine Salvage Enzymes

Since *Cryptosporidium parvum*, unlike other protozoa, lacks de novo pyrimidine nucleotide biosynthesis, two other potential targets are the pyrimidine salvage enzymes thymidine kinase (*Cp*TK) and kinase-uracil phosphoribosyltransferase (*Cp*UK-UPRT). Both trifluoromethylthymidine (TFT) and 5-fluorodeoxyuridine (FUdR) inhibit in vitro growth of the parasite at concentrations much lower than those that inhibit host cell proliferation (20–400-fold, respectively). This cytotoxicity differential is comparable to the therapeutic index for ribavirin, and indeed, TFT treatment in an acute mouse model of cryptosporidial infection resulted in a significant decrease in oocyst shedding (Sun et al. 2010).

11.4.9 Protein Kinases

An analysis of the *C. parvum* genome identified over 70 protein kinases, which may include promising drug targets (Wernimont et al. 2010; Artz et al. 2011). Protein kinases are recognized as essential to cell cycle regulation and exhibit a high degree of conservation, especially in the ATP-binding site. While the latter may challenge drug design, the observation that almost a quarter of the *C. parvum* kinases have no known orthologues outside *Cryptosporidium* species affords opportunities for selective drug development. A comparison of kinase orthologues in *P. falciparum* and *T. gondii* revealed distinct features associated with the 'gatekeeper' amino acid that can be exploited for drug design (Wernimont et al. 2010; Ojo et al. 2010). Cell

free enzyme assays of expressed kinases identified 13 pyrazolopyrimidine-based inhibitors with IC_{50} values < 10 nM (Artz et al. 2011).

A calcium-dependant protein kinase-1 was identified in *C. parvum* as well as a homologue in *T. gondii* (Wernimont et al. 2010; Murphy et al. 2010). These enzymes were targeted for drug development, in part, due to their characteristics related more to plant kinases than animal kinases. Pyrazolopyrimidine-based inhibitors showed selectivity for the protozoan kinases and exhibited low mammalian cell toxicity and significant inhibition of *T. gondii* invasion and proliferation with $EC_{50}s < 1 \mu M$ and some compounds showing <100 nM (Johnson et al. 2012). Subsequent drug development based on modification of the base compound, mebendazole, identified compounds with $IC_{50}s <50$ nM in cell free enzyme inhibition assays and low toxicities for mammalian cells ($EC_{50}s > 30 \mu M$) (Zhang et al. 2012). However, none of the compounds showed activity against cryptosporidial or toxoplasma growth in vitro below 1 μM , although it should be noted that no details were provided for the *Cryptosporidium* in vitro growth assay (Zhang et al. 2012).

11.5 Summary Comments on Current Drug Therapy

Considering the overall pattern of drug treatment data, it is apparent that no drugs that are available against other coccidian parasites have shown a robust efficacy against cryptosporidial infections. The potent efficacy of trimethoprimsulfamethoxazole (TMP-SMX) against Toxoplasma, Isospora, and Cyclospora is in stark contrast to the drugs used against Cryptosporidium (Goldberg and Bishara 2012; Seddon and Bhagani 2011). Even HIV-infected AIDS patients with chronic and severe diarrhea caused by Cyclospora or Isospora responded quickly and completely to TMP-SMX (Verdier et al. 2000). In a 2003–2004 study of chronic diarrhea among HIV/AIDS patients in Haiti (Dillingham et al. 2009), the occurrence of Cyclospora and Isospora at enrollment was observed to be <10 % compared to approximately 25 % of diarrheic patients in an earlier 1997–1998 study at the same Port-au-Prince clinic (Verdier et al. 2000). The lower prevalence of Cyclospora and Isospora was attributed to the widespread use of TMP-SMX as a prophylaxis or therapeutic agent in HIV/AIDS patients. This contrast is especially evident when comparing the activity of anticryptosporidial agents in immunocompetent versus immunocompromised patients. The largely modest activity of anticryptosporidial agents in immunocompetent patients combined with the essential failure of these same drugs to control or cure infections in immunocompromised patients argues that even the best drugs currently available are successful only in the context of an effective immune response, rather than by a direct and potent activity against the parasite itself.

11.6 Immunotherapy

11.6.1 Hyperimmune Colostrum

As described above, hyperimmune colostrum used as passive immunotherapy for cryptosporidiosis has been pursued as a strategy in humans since the late 1980s. In addition to the use of colostrum generated against whole antigen (Ag), several studies have evaluated colostrum or monoclonal antibodies (MAb) produced against specific antigens, many of these antigens involved in parasite attachment or invasion of host cells. For example, passive protection against cryptosporidiosis was obtained by treating immunosuppressed mice with immune colostrum generated in cows injected with recombinant pCP15/60 plasmid DNA before and after C. parvum infection (Jenkins et al. 1999). Immune bovine colostrum induced by immunization with C. parvum recombinant protein rC7, which is the C terminus of the Cp23 protein, provided substantial protection against cryptosporidiosis in neonatal calves (Perryman et al. 1999). Likewise, treatment with different polyclonal or MAbs resulted in the reduction in oocyst shedding as well as easing of clinical symptoms, although colonization still occurred, but at a considerably reduced level (Arrowood et al. 1989; Doyle et al. 1993; Perryman et al. 1990). One MAb, designated 3E2, which recognized multiple 46-770 kDa sporozoite Ags and a 1,300-kDa Ag designated CSL, was able to neutralize infectivity in vitro and control murine infection in vivo (Riggs et al. 1997). The 3E2 MAb combined with other antibodies, including anti-GP25-200 and anti-Cp23 demonstrated significant additive protection over that of the individual MAbs, reducing infection levels by 86-93 %. In addition, infection was completely prevented in up to 40 % of mice administered 3E2 alone or in combination with 3H2 and 1E10 Mabs (Schaefer et al. 2000).

Hyperimmune colostrum may also be used to reduce severity of diarrheal disease in farm animals, such as neonatal calves. This could potentially decrease transmission of *C. parvum* to animals and humans in agricultural settings or in developing countries where families and livestock live in close proximity to one another. In a recent study, cows vaccinated with rCP15/60 produced a significantly greater antibody response compared to controls and this response was strongly associated with the subsequent level of colostral antibody. Calves fed rCP15/60-immune colostrum showed a dose-dependent absorption of antibody, also associated with colostral antibody levels (Burton et al. 2011). Induction of the antibody was clearly evident but treatment efficacy was not demonstrated.

Despite the variable performance of immune colostrum in clinical trials, immunotherapy may still be useful in conjunction with conventional drug therapy or as a mechanism to decrease the severity of infection in neonatal animals or moderately immunocompromised individuals.

11.6.2 Monoclonal Antibodies

An example of MAb-based immunotherapy includes the use of a human CD40 agonist MAb, CP-870,893 to treat two X-linked hyper IgM syndrome patients with biliary cryptosporidiosis (Fan et al. 2012). The MAb activated B cells and antigen presenting cells (APCs) in vitro, restoring class switch recombination in XHM B cells and inducing cytokine secretion by monocytes. Although specific antibody responses were lacking, frequent dosing in one subject primed T cells to secrete IFN-gamma and suppressed oocyst shedding in stools. Nevertheless, oocyst shedding relapse occurred after discontinuation of therapy.

Another antibody-based immunotherapy involved the generation of an antibodybiocide fusion protein. *Cryptosporidium*-specific antibodies were fused with the antimicrobial peptide LL-37 and administered orally to neonatal mice in a prophylactic model of cryptosporidiosis (Imboden et al. 2010). Infections in treated mice were reduced by as much as 81 % in the mucosal epithelium of the gut. When administered simultaneously with oocyst inocula, several versions of antibody fusion proteins that differed in antigen specificity and in the biocide conjugate inhibited parasite growth in mouse intestinal tissue (up to 82 %), although none completely prevented infection.

11.7 Vaccines

Because of the lack of efficacious drug treatments, vaccine development that prevents disease or reduces the severity of infection is a relevant option. This is particularly true for certain groups such as immunocompromised individuals and children in developing countries since cryptosporidiosis in early childhood has been reported to be associated with subsequent impairment in growth, physical fitness, and intellectual capacity (Dillingham et al. 2002). Numerous immunogenic antigens of the C. parvum invasive stages involved in attachment or penetration of host cells have been identified (reviewed in (Boulter-Bitzer et al. 2007)). Several cryptosporidial antigens are immunodominant; some are surface and/or apical complex proteins that may mediate attachment and invasion. Sera from infected animals and humans recognize a number of immunodominant sporozoite antigens, including polypeptides of approximately 11, 15, 23, 44, 100, 180 and >200 (Boulter-Bitzer et al. 2007). These include the surface antigens CSL, Cp900, Cp23/27, Cp40/45, Cp15/17, Muc4 and Muc5, some of which are partially or heavily glycosylated. Antibodies developed against some of these antigens demonstrated therapeutic efficacy in mouse and animal models. Much of this work has focused on the Cp15 and Cp23 antigens.

DNA immunization has been used to induce antigen-specific B and T cell responses in various infection model systems (Hong-Xuan et al. 2005; Jenkins et al. 1995; Sagodira et al. 1999a, b). DNA vaccines expressing the Cp15/60

gene, a sporozoite surface antigen (Tilley et al. 1991), induced primarily a type-1 immune response when injected either intranasally or intramuscularly into mice (He et al. 2004; Sagodira et al. 1999b). Additionally, efficacy has been demonstrated by the generation of Cp23-specific immune responses: mice immunized with Cp23-DNA developed partial protection against *C. parvum* infection as shown by the >60 % reduction in oocyst shedding after challenge (Ehigiator et al. 2007). Administration of a DNA vaccine encoding *C. parvum* Cp15 and Cp23 resulted in induction of Th1 immune response and increased resistance to infection (Wang et al. 2010). Evaluation of a DNA vaccine comprised of P2 (CpP2), an important immunodominant marker in *C. parvum* infection (Priest et al. 2010), showed that CpP2-DNA followed with P2 protein (prime-boost), significantly increase antibody production over immunization with just the protein or CpP2-DNA alone. When challenged, reduction in oocysts production was not statistically significant, although a trend in reduction of infection was observed in the CpP2-DNA-immunized mice (Benitez et al. 2011).

Both humoral and cellular responses were elicited using a Salmonella strain-andvector combination that delivered Cp23 and Cp40 fused to the C-terminal fragment of tetanus toxin (Benitez et al. 2009). In another study, three antigens, Cp15, profilin, and a Cryptosporidium apyrase, were delivered in a heterologous primeboost regimen as fusions with cytolysin A (ClyA) in a Salmonella live vaccine vector and as purified recombinant antigens, and were found to induce specific and potent humoral and cellular immune responses (Manque et al. 2011). Profilin is a potent inducer of immune responses in mice by both Eimeria and Toxoplasma and works through the toll receptor TRL11. However, an analogous receptor (TRL11) has not been found in humans. Recently, a prime-boost immunization regimen using an intranasal route followed by oral Salmonella live vaccine vector of the Cp15 antigen increased immune responses but did not result in decreased infection (Roche et al. 2013). It is possible that other vectors (e.g. Listeria, adenovirus) may increase vaccine efficacy. Identification of other vaccine targets, multi-antigen formulations or constructs, or use of an attenuated Cryptosporidium strain could result in better immunological responses and protection from infection.

11.8 Conclusion

Key to the development of selective, efficacious anticryptosporidial agents is understanding cryptosporidial biochemistry and cell biology including studies into the basic mechanisms by which chemicals and drugs are transported from the host cell to the parasite (and vice versa), discerning stage-specific developmental changes, and identification of nutrient requirements and unique enzymatic pathways. The continuing increase in genome sequence data should aid in the identification and characterization of drug targets and increase our understanding of host/parasite interactions. However, the inability to cryopreserve the parasite, propagate the parasite continuously in vitro, and the lack of a genetic model hinders many avenues of chemotherapeutic research. Cryopreservation of the parasite would facilitate the establishment of standard isolates that could be used for multiple studies with consistency and reduced variability. The ability to propagate the parasite continuously in vitro could lead to establishment of a genetic model that would be useful for mutant/transfection studies and the identification of molecules as viable drug targets. Standardized positive control drugs are likewise needed when evaluating and validating potential anticryptosporidial agents in both in vitro and in vivo models. Understanding host-parasite interactions and the essential elements of immunity to *Cryptosporidium* spp. may lead to the development of effective immunotherapies or vaccines. Despite the many impediments faced by researchers working on *Cryptosporidium* parasites, active research in this area will hopefully overcome the barriers to success and more efficacious drugs and therapies will be developed to treat this potentially severe disease.

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Part IV *Cryptosporidium* and Water

Chapter 12 *Cryptosporidium* Oocysts in Drinking Water and Recreational Water

Paul A. Rochelle and George D. Di Giovanni

Abstract Oocysts belonging to a wide variety of species and genotypes of Cryptosporidium are common in livestock, wild animals, and humans. Consequently, water is frequently contaminated through direct contact with infected animals and their waste, run-off from contaminated land, or structural and engineering failures in water conveyance, storage, or treatment facilities. Oocysts are resistant to chlorine disinfection at the concentrations typically applied during drinking water treatment but properly operated treatment plants that utilize filtration usually remove oocysts from source water with high efficiency. Nevertheless, waterborne Cryptosporidium continues to be a public health concern. Outbreaks have been linked to treated drinking water, but regulations enacted over the last decade, better watershed management, and operational improvements have led to a decline in drinking water related cryptosporidiosis in some countries. However, the same period has seen a marked increase in cryptosporidiosis outbreaks caused by contamination of recreational water, particularly swimming pools. This chapter reviews recent waterborne outbreaks of cryptosporidiosis, discusses oocyst prevalence in drinking water and recreational water, examines the risk of waterborne transmission, and describes the principal methods for detecting oocysts in water, including genotyping environmental oocysts.

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12.1 Cryptosporidium in Water

The story of the 1993 cryptosporidiosis outbreak in Milwaukee is well known within the drinking water industry (MacKenzie et al. 1994). It remains the largest waterborne disease outbreak in U.S. history with an estimated 400,000 illnesses and approximately 150 deaths linked to the contamination. However, 20 years after the Milwaukee event, outbreaks of cryptosporidiosis still occur, linked to both drinking water and recreational water, so *Cryptosporidium* in water continues to be a public health concern. The response to the Milwaukee outbreak by regulators, public health professionals, academic researchers, and water utility scientists led to a plethora of detection methods, development of viability and infectivity assays, the application of molecular tools for genotyping oocysts in water, and evaluation of various disinfection technologies. In addition, many oocyst occurrence surveys were conducted, culminating in the U.S. with a period of regulatory monitoring under the Long Term 2 Enhanced Surface Water Treatment Rule (LT2ESWTR; USEPA 2006) and an intensive 10-year finished water monitoring program in the U.K. (DWI 1999). The LT2ESWTR is designed to improve the control of Cryptosporidium (and other pathogens) in drinking water systems and implementation of the regulation is expected to reduce the annual incidence of cryptosporidiosis by an estimated 89,000-1,459,000 cases and prevent 20-314 related premature deaths (USEPA 2006).

Cryptosporidium is particularly problematic for water utilities because it is common in surface waters and the oocyst stage of the parasite, which is found in environmental waters and is immediately infective after being excreted from infected hosts, is resistant to chlorine disinfection at the concentrations typically applied during drinking water treatment (see Chap. 13). However, despite everything that has been learned since the Milwaukee outbreak, it is still difficult to assess the true risk to public health from waterborne *Cryptosporidium*.

In the immediate aftermath of the Milwaukee outbreak, the only concern for the drinking water industry was "Cryptosporidium". However, research using polymerase chain reaction (PCR)-based techniques greatly enhanced our knowledge of the species and genotypes of Cryptosporidium oocysts found in surface waters. We now know that human-infectious oocysts predominantly belong to just two species (C. parvum and C. hominis), with a few other species playing a minor role in human infection (see Chap. 3). Additionally, there are many other animal-associated species and genotypes that do not appear to be pathogenic to humans. At least eight species and seven genotypes have been linked to human disease but it is generally recognized that the vast majority of human cryptosporidiosis cases are caused by C. parvum and C. hominis, and it appears that only C. hominis, C. parvum, and C. meleagridis readily infect immune-competent humans in all age groups (McLauchlin et al. 2000; Leoni et al. 2006). A genotyping study of 2,414 human clinical specimens from England obtained between 1985 and 2000 found C. parvum and C. hominis in 99 % of samples (Leoni et al. 2006). The remaining samples were identified as containing C. meleagridis (0.9 %), C. felis (0.2 %), C. andersoni (0.1 %), C. canis (0.04 %), C. suis (0.04 %) and the Cryptosporidium cervine genotype (0.04 %; now recognized as C. ubiquitum). Similarly, Chalmers et al. (2009a) genotyped 7,758 human clinical specimens obtained in England and Wales between 2000 and 2003, and 96 % were identified as C. parvum and/or C. hominis. The remaining 1 % of typeable samples was identified as C. meleagridis (0.7%), C. felis (0.05%), C. ubiquitum (0.05%), C. canis (0.01%), horse genotype (0.01%), and skunk genotype (0.01%). There have also been two documented cases of cryptosporidiosis due to the W17 chipmunk genotype (Feltus et al. 2006). So, finding C. parvum or C. hominis in drinking water sources is certainly a cause for concern and finding any of the other species that have been recovered from human infections would need investigating. However, oocysts of species and genotypes that do not usually pose a serious risk of human disease frequently contaminate surface waters. While these wildlife- and livestock-associated oocysts may be genetically distinct from the primary human-pathogenic species, they are morphologically similar and cross-react with the antibodies that are used in the most widely adopted detection methods, thus complicating risk assessments. Consequently, important issues have been raised with regard to *Cryptosporidium* risk assessment and the development of watershed management strategies. From a watershed management perspective, different strategies are needed to control human, wildlife, and livestock sources of oocysts. It is therefore reasonable to expect that detection methodologies which differentiate animal-related genotypes from the three Cryptosporidium species causing the majority of cryptosporidiosis in immunecompetent humans (C. hominis, C. parvum and C. meleagridis) may meet the needs of regulatory agencies and the water industry in addressing *Cryptosporidium* risk and control.

Species of Cryptosporidium are common in many different types of animals and consequently oocysts belonging to various species and genotypes are common in the environment. Oocysts are readily mobilized by rainfall, and so even if they are not deposited directly into a body of water by animal defecation, they are easily transported across land into water (Miller et al. 2008). There have been few studies of oocyst survival in untreated water under realistic environmental conditions, although some studies have examined survival in livestock wastewater lagoons and soil (Jenkins et al. 1999). However, compared to most human pathogens, Cryptosporidium is relatively stable in environmental waters, remaining infectious for weeks or months at typical surface water temperatures. Factors that determine whether or not oocysts released into surface waters will cause outbreaks or sporadic cases of disease include: the source of contamination and hence the species; the concentration of contaminating oocysts; the length of time oocysts are in the water; environmental temperature; and the presence of effective filtration and/or effective disinfection at treatment plants. The fact that waterborne outbreaks occur demonstrates that human-pathogenic oocysts can survive in environmental waters and retain their infectivity, but the amount of attenuation due to combined environmental stressors is not known. Oocysts are very sensitive to inactivation by UV-C irradiation (Rochelle et al. 2004; Johnson et al. 2005) and it has been suggested that sunlight may be the most significant inactivating agent in environmental waters (see Chap. 13). Water temperature is also a critical factor in oocyst survival in the environment. In a study using inclusion/exclusion of fluorescent dyes to assess loss of viability, oocyst suspensions lost 70 % of their infectivity when incubated in natural mineral waters for 84 days at 20 °C (Nichols et al. 2004). Cell culture assays have also been used to measure reductions in infectivity of oocysts in natural waters. Infectivity of oocysts stored in river water at 21–23 °C decreased by 2.6–3.3 log₁₀ over 84 days and was undetectable after 98 days (Pokorny et al. 2002). Infectivity of *C. parvum* oocysts suspended in raw surface water, as measured by cell culture combined with reverse transcription polymerase chain reaction (RT-PCR), was reduced by 3-log₁₀ in about 120 days at 18 °C and 40 days at 26 °C (Johnson et al. 2008). These values translated to a daily inactivation rate of 0.072-log₁₀/day at 18 °C. Similarly, inactivation of oocysts suspended in groundwaters and surface waters ranged from 0.009 log₁₀/day at 5 °C to 0.2 log₁₀/day at 30 °C (Ives et al. 2007).

12.2 Waterborne Outbreaks of Cryptosporidiosis

Cryptosporidiosis outbreaks linked to both drinking water and recreational water continue to occur around the world (Tables 12.1 and 12.2). Of the 325 water-associated protozoan disease outbreaks reported worldwide for the hundred years ending in 2004, 51 % were caused by *Cryptosporidium* spp. (Karanis et al. 2007). During the next 6 years (2004–2010) *Cryptosporidium* caused 60 % of the 199 reported outbreaks of protozoan diseases linked to all sources (Baldursson and Karanis 2011). Almost 200 waterborne outbreaks of cryptosporidiosis had been reported globally by the end of 2010 and have occurred in the U.S., U.K., Australia, New Zealand, Germany, Sweden, Japan, Spain, Norway, Denmark, France, and Ireland. It is highly likely that outbreaks also occur in most other countries but they may lack suitable reporting mechanisms or detection methods, and comprehensive monitoring data are not available for most countries.

During the period 1991–2002, *Cryptosporidium* caused 7 % of the 207 waterborne disease outbreaks in the U.S. However, due to the Milwaukee outbreak, *Cryptosporidium* caused more illnesses than all other waterborne pathogens combined during the 11 year period of 1991–2002 (Craun et al. 2006). Although the overall number of documented waterborne cryptosporidiosis outbreaks has increased substantially in the U.S. over the last 20 years (4 outbreaks in 1992 vs. 39 in 2007; Yoder et al. 2010), there have been no outbreaks linked to community surface water supplies in the last two decades (Yoder et al. 2012).¹ In fact, the majority of outbreaks are associated

¹ The 20-year period without a cryptosporidiosis outbreak linked to a municipal surface water supply in the U.S. ended recently. An outbreak in Baker City, Oregon during the summer of 2013 was linked to an unfiltered surface water supply with up to 91 oocysts/L in one supply creek (www. bakercity.com; accessed Sept. 25, 2013).

Date	Location	Source/cause	Cases ^b
2000	North West England	Ground water under the influence of surface water, heavy rain- fall, inadequate treatment	58
2000	Scotland	Inadequate filtration	
2000	Northern Ireland	Human sewage from septic tank contaminated water supply, <i>C. parvum</i>	168
2001	Saskatchewan, Canada	Treatment plant failure	1,907
2001	Northern Ireland	Structural failures at treatment plant, wastewater contamination, <i>C. hominis</i>	306
2004	Bergen, Norway	Possible sewage contamination, coincident with large giardiasis outbreak, <i>C. parvum</i>	133
2005	South East England	Public water supply, C. hominis	140
2005	North Wales, UK	Contaminated source water reservoir, C. hominis	231
2007	Galway, Ireland	Contaminated source water lake and heavy rainfall	>240
2008	Northampton, England	Contamination of treated water by rabbit feces/carcass, <i>C. cuniculus</i>	>400
2010	Ostersund, Sweden	Sewage contamination of source water lake, C. hominis	12,700
2011	Skellefteå, Sweden	Sewage contamination of river, C. hominis	147

Table 12.1 Examples of cryptosporidiosis outbreaks linked to drinking water^a

^aCompiled from various sources: (Baldursson and Karanis 2011; Chalmers 2012; EPA 2011; Glaberman et al. 2002; Karanis et al. 2007)

^bCase numbers for some outbreaks are based on laboratory confirmed infections while others are based on epidemiological studies and the size of the potentially impacted population

Date	Location	Source/cause	Cases
2002	Sweden	Swimming pool, C. parvum	800
2003	Majorca, Spain	Hotel swimming pool, C. parvum	391
2004	California, USA	Treated recreational water in water park, C. parvum	336
2004	California, USA	Water slide in a water park	>250
2005	Nagano, Japan	Hotel swimming pool, C. hominis	41
2005	New York, USA	Interactive fountain	2,307
2005	Ohio, USA	Community swimming pool	523
2005	London, England	Swimming pools, C. hominis	>120
2005	NSW, Australia	Swimming pool	254
2006	Missouri, USA	Interactive fountain in water park	116
2007	Utah, USA	Various treated recreational water venues	1,302
2008	Texas, USA	Lake, pools, interactive fountain, C. hominis	2,050
2009	Australia	Swimming pools, C. hominis	1,141
20			2012

Table 12.2 Examples of cryptosporidiosis outbreaks linked to recreational water^a

^aCompiled from various sources: (Baldursson and Karanis 2011; Beach 2008; Chalmers 2012; Hlavsa et al. 2011)

with recreational use of water, particularly treated water such as swimming pools (Yoder et al. 2012).

There were 149 outbreaks of cryptosporidiosis in the U.K. for the period 1983–2005 (Nichols et al. 2006). Fifty five of these outbreaks were linked to municipal drinking water, six were associated with private water supplies, and swimming pools were the source of infections in 43 outbreaks. During the period 1983–1997, there were 25 waterborne outbreaks of cryptosporidiosis in the U.K. with 3,455 reported cases of illness (Nichols 2003). As recently as May 2013, an outbreak of cryptosporidiosis with at least 13 confirmed cases in Roscommon, Ireland, led to a boil water advisory affecting over 10,000 people (Roscommon County Council 2013). While acute contamination and failures at treatment plants have often been implicated as the cause of cryptosporidiosis outbreaks, they can occur in water produced by treatment plants meeting all regulatory standards (Neira-Munoz et al. 2007).

Although much of the early research and regulatory focus was on drinking water, Cryptosporidium has become the leading cause of recreational water outbreaks in the U.S. The spatial and temporal proximity of contaminating individuals and "recipient" individuals in recreational water, particularly swimming pools, means that outbreaks linked to recreational water probably result from recent contamination and so oocyst survival in recreational water is not a primary issue. In the 30 years spanning 1971-2000, Cryptosporidium caused 15 % of outbreaks associated with recreational water (Craun et al. 2005). More recently (2007-2008), 59 out of 82 disease outbreaks with an identified infectious agent linked to treated recreational water were caused by Cryptosporidium. Between 1988 and 2008, 136 documented outbreaks of cryptosporidiosis linked to recreational use of water in nine countries sickened a total of 19,271 people (Beach 2008). The average number of people infected in these outbreaks was 142 but there were over 5,000 infections in one outbreak. There were 5,697 cases of illness in a statewide outbreak linked to swimming pools in Utah in 2007 (Hlavsa et al. 2011) leading to restrictions on the use of public pools by young children. Of the U.K. outbreaks linked to recreational water in the period 1983-2005, the number of cases ranged from 3 to 152 and the sources of infection were public and private swimming pools, rivers, interactive water features (e.g., splash zones), and water fountains. During a 1 year monitoring period of seven swimming pools in the Netherlands, 4.6 % of samples contained Cryptosporidium oocysts (Schets et al. 2004). Finally, at the time of writing this chapter, another outbreak during the first quarter of 2013 in Victoria, Australia was being linked to swimming pools (Lester 2013). Detailed reviews of outbreaks associated with recreational water are provided by Beach (2008) and Chalmers (2012).

Some of the earliest applications of genotyping tools demonstrated that in the U.K. and the U.S., 67 % of waterborne cryptosporidiosis outbreaks were caused by *C. hominis* while *C. parvum* was the causative agent in the remaining 33 % (McLauchlin et al. 2000; Sulaiman et al. 1998). Genotyping of *Cryptosporidium* isolates associated with the 1993 Milwaukee outbreak identified *C. hominis* as the etiologic agent (Zhou et al. 2003). With the exception of one recent incident, only

C. parvum and *C. hominis* have been identified as causes of waterborne outbreaks of cryptosporidiosis (Xiao and Ryan 2008). A 2008 outbreak in Northamptonshire, England was caused by a rabbit genotype, now renamed *C. cuniculus* (Chalmers et al. 2009b; Robinson et al. 2010) (Chap. 3). It is noteworthy that multilocus genotyping of the rabbit genotype indicated that it is very closely related to *C. hominis*, as opposed to typical animal-associated genotypes (Robinson et al. 2010). Reanalysis of 3,030 sporadic human cryptosporidiosis samples previously identified as containing *C. hominis* revealed that *C. cuniculus* had a prevalence of 1.2 % among clinical isolates (Chalmers et al. 2011).

12.3 Risk Assessment

There are many risk factors for acquiring cryptosporidiosis, including recreational use of treated water (swimming pools and water parks), swimming in untreated water, consuming contaminated food, person to person transmission, contact with farm animals, exposure in day care facilities, drinking contaminated water, and foreign travel (see Chap. 2). However, the potential magnitude of outbreaks from drinking water is greater than for most other sources. Consequently, there have been numerous efforts to derive estimates of the risk to public health from *Cryptosporidium* oocysts in water but the contribution of drinking water to the overall cryptosporidiosis disease burden is still not clear. Many factors contribute to the uncertainty of calculating this risk, including:

- · Widely varying oocyst occurrence data in source and finished waters
- Relative prevalence of human-pathogenic and animal-associated species and genotypes
- · Variable recovery efficiencies of detection methods
- · Non-standardized viability and infectivity assessment methods
- · Differing risk assessment models and underlying assumptions
- Varying susceptibility to infection within different human populations
- · Possible protective immunity resulting from low-level endemic exposure
- Varying infectious doses for different species and strains of human-pathogenic *Cryptosporidium* spp. (Okhuysen et al. 1999; Chappell et al. 2006)
- · Relative contributions of other sources of infection

According to one of the first risk assessment studies, based on an individual's average water consumption of 1.5 L/day and an acceptable daily oocyst intake of 6.5×10^{-5} (calculated from human dose response data), the theoretical maximum acceptable concentration in drinking water was 4.4×10^{-2} oocysts/1,000 L (Haas et al. 1996). This value is considerably lower than the detection limit of currently used monitoring methods. In another study, based on a finished water concentration of one oocyst/1,000 L, it was estimated that tap water was responsible for 6,000 cases of illness annually in New York City (Perz et al. 1998). More recently, it was estimated that the daily infection risk for the general immune-competent population

in New York City was 3–10 cases per 100,000 people (Makri et al. 2004). However, it is well established that some sections of the community are more susceptible than others to infection by *Cryptosporidium*. For example, in a case–control study of AIDS patients in San Francisco up to 85 % of endemic cryptosporidiosis cases were attributed to drinking unboiled tap water (Aragón et al. 2003).

In contrast to the above risk assessments, some studies have found no association between drinking water and endemic cryptosporidiosis. The most significant risk factors for sporadic cryptosporidiosis among 282 immunocompetent individuals in seven U.S. states were recent international travel, contact with cattle, contact with young children with diarrhea, and swimming in untreated surface water (Roy et al. 2004). Drinking water was not associated with *Cryptosporidium* infection in six of these states but consumption of well water within Minnesota was a significant risk factor. Similarly, in a case–control study of immunocompetent individuals in the San Francisco Bay Area, there was no significant association between cryptosporidiosis and drinking water (Khalakdina et al. 2003). The major risk factor for cryptosporidiosis within this population was recent travel to another country. Similar results were obtained in the U.K. where epidemiological and case–control studies found no association between drinking water and sporadic cases of cryptosporidiosis in northwest England (Hunter et al. 2004; Hughes et al. 2004).

Nevertheless, regulatory improvements within the water industry, better watershed management, and changes in treatment practices have resulted in reduced disease incidence, confirming that drinking water is a contributing factor to cryptosporidiosis in the community. Following the introduction of new drinking water regulations in England and Wales in 1999, the annual incidence of cryptosporidiosis was reduced by 6,770 cases (Lake et al. 2007). There was a significant reduction in disease incidence during the first 6 months of each year but interestingly there was no change during the second half of the year. This disparity in seasonal disease incidence may have been due to improved land and water management practices reducing domestically acquired infections during the first part of the year. However, travel-associated illnesses starting in the summer and continuing through the second part of the year were not impacted by regulatory changes in the U.K. After installing membrane filtration at two treatment plants in the U.K., the annual incidence of 22 cryptosporidiosis cases per 100,000 people declined to less than 10 cases per 100,000 people (Goh et al. 2005). The authors concluded that drinking cold, unboiled municipal tap water was one of the greatest risk factors for sporadic cryptosporidiosis.

Regardless of the theoretical or actual risk of waterborne cryptosporidiosis, when oocysts are detected in drinking water, utilities and regulatory agencies take measures to minimize community-wide outbreaks. The usual action is to issue boilwater orders or advisories. However, a recent cost-benefit analysis determined that nine illnesses per 10,000 people was the point at which the expected monetized benefit of a boil water notice would exceed the expected costs (Ryan et al. 2013). This infection rate far exceeds the USEPA "acceptable" infection rate of 1 in 10,000, and it corresponded to a raw water concentration of 46 oocysts/L and a finished water concentration of 0.046 oocysts/L (assuming 99.9 % removal). But

the costs of disease are not just economic, particularly at the level of the individual. Utilities that are routinely detecting human-pathogenic oocysts in finished water should take mitigating action regardless of the economic break-even point.

Despite the emphasis on drinking water as a potential mode of transmission, it is well established that sources such as food, contact with animals, and international travel also commonly transmit *Cryptosporidium* oocysts. In fact, some studies have suggested that food and other modes of transmission may be at least as important as drinking water and may be more likely to transmit higher oocyst doses (Frost et al. 2005). There is some indication that low level endemic exposure to oocysts in drinking water may confer protective immunity during outbreaks (Frost et al. 2005; Chappell et al. 1999). This has led to speculation that cryptosporidiosis only became a serious epidemic disease in western countries after improvements in sanitation and drinking water treatment reduced levels of low-dose exposure that had formerly maintained protective immunity in the community (Frost et al. 2005).

12.4 Detection Methods

USEPA Method 1623 is the most commonly used oocyst detection method in the U.S. and is recognized as the standard method for *Cryptosporidium* and *Giardia* in surface waters (USEPA 2005). It was used in the most recent round of regulatory Cryptosporidium monitoring of untreated water in the U.S. under the LT2ESWTR (USEPA 2006). It involves sample concentration by filtration and centrifugation, oocyst purification by immunomagnetic separation (IMS), and detection by immunofluorescence assay microscopy. Method 1623 includes four options for initial sample concentration: Envirochek and Envirochek HV filter capsules, Filta-Max compressed foam filters, and a portable continuous flow centrifuge. Each of these methods requires different post-concentration elution and processing prior to sample clean up by IMS. Ultrafiltration through 50–80 kDa molecular weight cut-off hollow-fiber filters has also proved efficient for recovering Cryptosporidium oocysts from large volumes of environmental and finished water samples (Hill et al. 2007; Kuhn and Oshima 2002; Lindquist et al. 2007). However, ultrafiltration has not yet been validated for use with Method 1623. The Envirochek filter is a pleated membrane of polyethersulfone whereas the membrane in the Envirochek HV (high volume) capsule is polyester. The minimum volume of untreated surface water required for analysis by Method 1623 is 10 L, although depending on sample turbidity more than one filter may be required to filter 10 L, and special precautions are sometimes necessary to ensure sample integrity (Fig. 12.1). The Envirochek HV filter can typically be used for up to 50 L of surface water and up to 1,000 L of finished water. However, some authors have reported difficulty in passing 10 L of turbid source waters through Envirochek and Envirochek HV filters (DiGiorgio et al. 2002). Detected organisms are verified by staining with 4'6-diamidino-2phenylindole (DAPI) and differential interference contrast (DIC) microscopy. The method was recently updated to include pre-elution of the filter in sodium



Fig. 12.1 Special sampling precautions to ensure samples remain adequately cooled during prolonged filtration of large volumes of finished water using USEPA Method 1623 filtration equipment. In this example, finished drinking water samples were being filtered for the detection of infectious *Cryptosporidium* using a cell culture assay (Rochelle et al. 2012). The ambient temperature was 43 °C (110 °F) and samples needed to be kept cool during filtration to prevent inactivation of oocysts (Photos courtesy of J. Hernandez, City of Scottsdale Water Resources Dept. (Arizona, USA))

hexametaphosphate and adding an additional rinse to the IMS (Method 1623.1; USEPA 2012).

Until 2008, regulations in the U.K. required continuous *Cryptosporidium* monitoring of treated drinking water (DWI 1999). The method required continuous sampling at a flow rate of at least 40 L/h with the filter being changed every 24 h (\geq 960 L in 24 h). As with Method 1623, the U.K.-approved method involved sample collection by filtration through Envirochek HV capsules or FiltaMax compressed foam filters, secondary concentration and purification by IMS, staining with FITC-labeled anti-*Cryptosporidium* antibody, and examination by fluorescence and DIC microscopy (DWI 2005). The method used 0.5 % sodium polyphosphate for the pre-elution treatment. A slightly modified IMS procedure allowed solid-phase automated scanning cytometry to be used for slide examination.

Although numerous PCR-based methods and primers for the detection of *Cryp*tosporidium in water have been described and compared, PCR has not been developed as a routine tool for monitoring or regulatory compliance within the water industry. The various PCR assays have different nucleic acid targets and include modifications for viability determination by detecting mRNA (RT-PCR), different species specificities (e.g. all *Cryptosporidium* spp. or individual species) and reported sensitivities as low as one oocyst. Amplification targets include the 18S ribosomal RNA gene, β -tubulin, heat shock protein 70, and oocyst wall proteins, among others (Leetz et al. 2007; Rochelle 2001; Xiao et al. 2006). Most PCR-based assays were developed as presence/absence methods but assays using various quantitative-PCR formats applied to oocysts in water have also been developed (Guy et al. 2003; Kishida et al. 2012; Masago et al. 2006). However, despite reports of sensitive and reproducible PCR-based detection of oocysts in water, a recent study concluded that although quantitative-PCR methods are useful for the detection and species identification of *Cryptosporidium* in environmental samples, they cannot accurately and consistently measure the low concentrations of oocysts that are typically present in sources of drinking water (Staggs et al. 2013).

C. parvum, C. hominis, and other species of Cryptosporidium are morphologically similar but can be readily differentiated by DNA sequence variations in many genes. Consequently, the real power of applying PCR-based methods to Cryptosporidium in water comes from the ability to analyze amplification products by restriction fragment length polymorphisms (RFLP) or sequencing. The current USEPA Method 1623 and the U.K. DWI regulatory methods are microscopybased, as described above. While these methods are capable of enumerating low levels of oocysts, they do not determine the species or genotypes of the detected Cryptosporidium. Researchers have attempted to address this shortcoming by developing PCR methods for genotyping oocysts directly recovered from water samples or from microscope slides following analysis by fluorescence microscopy. While valuable information can be obtained by directly genotyping oocysts from water samples, accurate quantification of oocysts is not possible with current PCR technology. Therefore, methods focused on genotyping oocysts recovered from regulatory slides are the most useful since the overriding regulatory requirement of oocyst quantification (including empty oocysts) is satisfied. Molecular species identification or genotyping of oocysts detected by Method 1623 have not been approved by the USEPA, and it is unlikely that genotyping will be a regulatory requirement in the near future in the U.S. However, a simplified genotyping method for oocysts present on Method 1623 slides has been developed (Di Giovanni et al. 2010) and a genotyping method was developed for oocysts isolated during routine water monitoring in the U.K. (Nichols et al. 2010).

Nested polymerase chain reaction restriction fragment length polymorphism (PCR-RFLP) methods have been used for the sensitive detection and discrimination of Cryptosporidium species and genotypes in water (e.g., Xiao et al. 2000; Nichols et al. 2010; Ruecker et al. 2013). These methods involve two rounds of PCR amplification for increased sensitivity. The amplified DNA fragments are then subjected to restriction enzyme digestion which generates DNA fragment patterns unique to different Cryptosporidium species and genotypes. Although nested PCR-RFLP genotyping methods have contributed greatly to our understanding of waterborne Cryptosporidium, they are not practical for routine use by most water utility and water quality laboratories. Nested PCR is prone to contamination with PCR product, and the extensive manipulation of PCR products for RFLP analysis increases this risk. At present, the majority of utility and water quality laboratories have limited or no molecular experience. In response, a slide genotyping method that can readily be used by water quality laboratories to distinguish humanpathogenic (i.e. C. hominis, C. parvum, and C. meleagridis) from animal-associated Cryptosporidium oocysts was developed under Water Research Foundation Project 4099 (Di Giovanni et al. 2010). The developed method uses single-round multiplex PCR targeting the Cryptosporidium 18S rRNA and heat shock protein 70 (hsp70) genes and is compatible with conventional or real-time formats. The melting temperature (T_m) of PCR products (amplicons) is dictated by their DNA sequences. High resolution melt (HRM) analysis allows precise determination of amplicon T_m and further discrimination of human and animal-associated *Cryptosporidium* spp. The method is currently undergoing international multi-laboratory evaluation as part of Water Research Foundation Project 4284. Although slide genotyping will not be a regulatory requirement under the next round of LT2ESWTR monitoring (due to start in April, 2015), its use as a research tool will provide added value to the monitoring, aid development of effective watershed management plans, and provide critical data for refinement of *Cryptosporidium* human health risk assessments.

12.5 Occurrence in Source Waters

The actual occurrence of human-pathogenic Cryptosporidium oocysts in source and finished waters is still not clear due to natural variation in oocyst occurrence, variable method recovery efficiencies, and the effect of the water matrix on the sensitivity of the detection method. Various research surveys conducted both before the Milwaukee outbreak and in the immediate aftermath generally reported high oocyst prevalence in untreated and finished drinking waters. In the first survey of oocyst occurrence in surface waters, samples that were impacted by domestic and agricultural waste had Cryptosporidium concentrations as high as 5,800 oocysts/L (Madore et al. 1987). A large survey of North America spanning 1988-1993 reported that 60 % of samples were positive for Cryptosporidium oocysts (LeChevallier and Norton 1995) while a study in Canada demonstrated lower levels of contamination with oocysts detected in 4.5 % of raw water samples (Wallis et al. 1996). Other studies detected oocysts in 6 % of stream samples in Wisconsin (Archer et al. 1995), 63 % of river samples in Pennsylvania (States et al. 1997), and 13 % of surface waters in New Zealand (Ionas et al. 1998). More recently, a version of Method 1623 using Filta-Max filters was used to detect oocysts in water at Portuguese river beaches (Julio et al. 2012). Oocysts were detected in 82 % of samples with concentrations up to 5.3 oocysts/L. In addition, a study of raw water samples in Canada using USEPA Method 1623 (approximately 20 L grab samples), reported that >45 % of water samples contained Cryptosporidium oocysts (N = 1,296; Ruecker et al. 2013). In general, occurrence surveys using a variety of detection methods reported high oocyst prevalence in rivers and streams, with 100 % of samples being positive in some surveys and concentrations of less than one to hundreds of oocysts per liter (Clancy and Hargy 2008).

Regulatory monitoring programs typically demonstrate lower prevalence than most research studies. The first regulatory monitoring of *Cryptosporidium* in U.S. waters was conducted under the USEPA's Information Collection Rule (ICR). This survey of 5,838 untreated source waters throughout the U.S. reported an average occurrence of 6.8 % with a mean concentration of 0.067 oocysts/L (Messner and Wolpert 2003). In follow-up supplementary surveys of fewer facilities, 14 % of samples were positive with an average concentration of 0.053 oocysts/L. Starting in 2006, monitoring for Cryptosporidium was mandated in the U.S. by the LT2ESWTR (USEPA 2006). Drinking water facilities serving over 10,000 people were required to monitor for oocysts in at least 24 consecutive monthly samples of raw water using USEPA Method 1623. Out of almost 40,000 samples (mostly 10 L raw water grab samples) collected under the first round of monitoring at 1,670 sampling sites from 1,376 facilities across the U.S., 7 % contained at least one *Cryptosporidium* oocyst (Messner 2011a). Over half of the facilities (51 %) reported no positive samples at all throughout the entire monitoring period. In oocyst-positive samples, concentrations ranged from 0.1 oocysts/L (1,720 samples out of 2,895 positives) to a high of 16 oocysts/L (Ongerth 2013a). A total of 57 samples (0.14 %) had concentrations greater than one oocyst per liter. In the positive samples, a single oocyst was detected in more than half of the samples (59 %), 94 % had 1-4 oocysts, 4 % had 5-9 oocysts, and 2 % had ten or more oocysts. The locations with the highest prevalence of oocysts were on the Mississippi River downstream of the Ohio River and Missouri River confluence. Based on the LT2ESWTR monitoring data, at least 80 treatment plants in the U.S. serving over 10,000 people will be required to make improvements to increase oocyst removal or inactivation.

However, as with most microbe detection methods, recovery efficiencies for Method 1623 are rarely 100 % and can vary considerably between laboratories, between analysts, and between different sampling sites, depending on a number of factors, including the chemical and biological characteristics of water samples. Recovery efficiencies for sample sites monitored during the LT2ESWTR ranged from zero to >90 % with a mean of 39.7 % (N = 3,370; Messner 2011b). Similarly, Method 1623 recoveries ranged from 36 % in highly turbid water (99 NTU) to 75 % in lower turbidity samples (20 NTU; DiGiorgio et al. 2002) and were 3.1–90.3 % in samples from coastal streams in Australia (Ongerth 2013b). Consequently, since most oocyst occurrence surveys do not account for matrix effects and measurement of recovery efficiency, it is likely that occurrence is actually higher than most reported values. Analyzing the LT2ESWTR dataset by cumulative probability distributions and accounting for recovery efficiencies, a recent study concluded that one oocyst per liter will occur at 50 % of raw water sample sites at least 25 % of the time (Ongerth 2013a).

12.6 Species Identification and Genotyping of Oocysts in Water

Most genotyping studies of oocysts in environmental waters have demonstrated the presence of a wide variety of species and genotypes, including human infectious oocysts and species/genotypes that have not been linked to human disease (Jiang et al. 2005; Nichols et al. 2010; Ruecker et al. 2005; Wilkes et al. 2013; Xiao et al. 2000; Yang et al. 2008). PCR-RFLP targeting the 18S rRNA gene

demonstrated that 93 % of storm water samples were positive for *Cryptosporidium* spp. with 12 different genotypes represented. None of the detected genotypes matched those typically found in human, farm animals, or domestic animals. However, four were identical or closely related to *C. baileyi*, and *Cryptosporidium* genotypes from opossums and snakes, indicating that wildlife was the primary source of oocyst contamination of surface water during storms (Xiao et al. 2000). The same method was also used to analyze untreated surface water and wastewater samples. *Cryptosporidium* was detected in 45.5 % of surface water samples and 24.5 % of raw wastewater samples (Xiao et al. 2001). The predominant genotypes in surface water were *C. parvum* and *C. hominis*, while *C. andersoni* was most commonly detected in wastewater. In the South Nation River Basin in Canada, 21 % of raw water samples (N = 690) contained wildlife-specific genotypes and 13 % contained livestock genotypes, while avian and human-specific genotypes were identified in 3 % and <1 % of samples, respectively (Wilkes et al. 2013).

Several research groups have developed methods for genotyping *Cryptosporidium* oocysts recovered from regulatory slides. A nested PCR-RFLP assay for the 18S rRNA gene was used to analyze 1,042 *Cryptosporidium* IFA microscopy positive slides obtained from Scottish raw and finished drinking waters under the Scottish Water Routine *Cryptosporidium* Monitoring Programme (Nichols et al. 2010). Volumes of filtered water ranged from 300 L for turbid raw waters to a maximum of 4,000 L for groundwaters. The frequency distribution was a single oocyst on 34 % of slides, 75 % of slides had 1–4 oocysts, 12 % had 5–9 oocysts, and 11 % had ten or more oocysts. Genotyping success was 45 % for single oocyst slides and overall genotyping success was 62 % for all slides. Genotyping revealed that animal-associated *Cryptosporidium* were present in almost 90 % of the positive samples. The most frequently observed species and genotypes were *C. andersoni* (27 %), *C. ubiquitum* (20 %), human-pathogenic *C. parvum* and *C. hominis* (combined at 12 %), and muskrat genotypes (7 %).

In a similar study of 1,296 Canadian raw water samples (approximately 20 L grab samples) analyzed using USEPA Method 1623, more than 45 % of water samples tested positive for Cryptosporidium by IFA microscopy, yielding 601 slides for subsequent genotyping analysis (Ruecker et al. 2013). Distribution of oocysts on slides was very similar to that reported by Nichols et al. (2010), with 30 % of slides having a single oocyst, 71 % had 1-4 oocysts, 19 % had 5-9 oocysts, and 10 % had ten or more oocysts. Slides were analyzed by nested PCR-RFLP of the 18S rRNA gene for species/genotype identification (Ruecker et al. 2011). Genotyping success was 57 % for single oocyst slides and overall genotyping success was 70 % for all slides. Genotyping results were also similar to the Nichols et al. (2010) study in Scotland, with over 90 % of the samples found to contain animal-associated Cryptosporidium species. The most frequently observed species and genotypes were C. andersoni (44 %), muskrat genotypes (48 %), C. ubiquitum (5 %), and human-pathogenic C. parvum and C. hominis occurring in only 4 % (combined) of genotyped samples. Unfortunately, provisions for slide archival and genotyping for research purposes were not in place during the first round of the LT2ESWTR.

12.7 Oocysts in Treated Drinking Water

Oocysts are resistant to chlorine disinfection at the concentrations typically applied during drinking water treatment but correctly operating treatment plants that utilize filtration usually remove oocysts from source water. However, oocysts were reported in 3.8-40 % of treated drinking water samples at concentrations up to 0.48 oocysts/L (Rose et al. 1997). A survey of treatment plants in Wisconsin detected oocysts in 4.2 % of finished water samples (Archer et al. 1995) and another survey detected oocysts in 3.5 % of treated drinking waters (Wallis et al. 1996). In 1998, high oocyst concentrations detected in the treated water of Sydney, Australia and multiple detections in source water led to a period of on-again, off-again boil water advisories over 2 months. Although no infections were attributed to this contamination, the incident had long-ranging political, technical, operational, and managerial consequences in Australia (McClellan 1998). During the subsequent 6 years of monitoring Sydney's water supply that was precipitated by this incident, oocysts were detected in just 0.04 % of treated drinking water samples (N = 4,961; O'Keefe 2010). However, the monitoring period was characterized by an extreme multi-year drought and so low oocyst occurrence may not have been reflective of "normal" conditions. Finished water was also monitored at a treatment plant in Japan. No oocysts were detected in 365 consecutive daily samples of 180 L each (Masago et al. 2004).

Regulatory or systematic and widespread monitoring of Cryptosporidium in either raw or treated water has only been conducted in a few countries (Table 12.3). Until October 2008, the UK drinking water regulations included the most intensive Cryptosporidium monitoring program ever undertaken. The regulation required continuous monitoring of Cryptosporidium oocysts in finished drinking water for at least 23 h per day at a flow rate of at least 40 L per hour (DWI 1999). Although the majority of samples analyzed during this decade-long monitoring program were negative, Cryptosporidium oocysts were occasionally detected in finished drinking water. During the period 2000–2002, a total of 97,999 samples were analyzed (total volume = 115,303,050 L), 5.5 % were positive, and the average oocyst concentration was 0.0002 oocysts/L (Smeets et al. 2007). In the earlier years of the monitoring program, oocysts were detected at least once in the finished water from many plants. For example, in 2002, 1.9 % of samples were positive but oocysts were detected at least once from 68 % of sample sites (DWI 2002). Similarly, in 2003, 1.1 % of samples were positive with 54 % of sample locations reporting detection in at least one sample. So clearly, the public is being exposed to low levels of Cryptosporidium oocysts in finished drinking water. Nevertheless, the results of this extensive monitoring program allowed the DWI to conclude that treated drinking water is not a major source of exposure of the population to Cryptosporidium oocysts. In 2008, a total of 50,569 samples were analyzed (total volume = 46,523,480 L) from 204 plants but none of them exceeded the treatment standard of <1 oocyst/L (DWI 2008).

Country	Regulation/requirement	Monitoring dates	Type of water	Oocyst occurrence
Sydney, Australia	Outcome of Sydney Water Enquiry, 1998	1998–2004	Finished	0.04 %
England and Wales	UK Water Supply Regulation (Amended), 1999	1999–2008	Finished	2.8 %
Scotland	Cryptosporidium (Scottish Water)	2006-2007	Finished	7 %
	Directions, 2003		Raw	26 %
United States	Long Term 2 Enhanced Surface Water Treatment Rule, 2006	2006–2008	Raw	7 %

Table 12.3 Regulatory or required monitoring of Cryptosporidium in water

Despite significant efforts in watershed management and treatment upgrades, Cryptosporidium remains a challenge to drinking water quality in Scotland. Consequently, regulatory monthly monitoring is required at all treatment plants in Scotland (The Cryptosporidium [Scottish Water] Directions 2003). During 22 months of monitoring finished water (2006-2007), a total of 20,249 treated water samples from 304 drinking water supplies were analyzed. Oocysts were detected in 7 % of samples with a maximum concentration of 1.6 oocysts/L (Wilson et al. 2008). Oocysts were detected at least once in 54 % of supplies. From a public health perspective, the most risky source water catchments were those that were drier than average but had occasional high rainfall events, which flushed oocysts into the water. DNA could not be amplified from all of these monitoring slides, but of those that could be amplified, 19 % were identified as C. andersoni, 8 % were a cervine genotype, 5 % were C. parvum, 0.3 % were C. baileyi, 0.3 % were C. bovis, and 0.3 % were a muskrat genotype. Similarly, C. andersoni, C. parvum, and C. ubiquitum were identified on 4.1 %, 4.3 %, and 12.6 % of regulatory finished water monitoring slides in Scotland (Nichols et al. 2010). Mixed species or genotypes were detected on 19 % of slides. During 2011, a total of 8,919 finished water samples were analyzed under the regulatory directive in Scotland. Of these, 378 (4.2 %) samples tested positive for oocysts, the highest percentage of positive samples since 2008. Similarly, 91 of 264 (34 %) treatment plants reported at least one positive sample (Drinking Water Quality Regulator for Scotland 2012).

12.8 Infectivity of Waterborne Oocysts

Cryptosporidium infectivity assays have typically been used to assess the efficacy of disinfection processes but they can also be used to inform risk assessment models by determining the infectivity of oocysts in environmental water samples. The infectivity of oocysts recovered by Method 1623 can be tested in cell culture assays by modification of the oocyst-magnetic bead disassociation step at the end of the IMS procedure and omitting microscopic examination of oocysts. A variety of methods have been developed for detecting infection in cell culture and assays have

Type of water	Number of samples	Mean sample volume	Positive samples	Reference
Finished drinking water	370	943 L	0 %	Rochelle et al. (2012)
Finished drinking water	1,690	100 L	1.4 %	Aboytes et al. (2004)
Source water	560	10 L	3.9 %	LeChevallier et al. (2003)
Source water	122	10 L	4.9 %	Di Giovanni et al. (1999)
Filter backwash water	121	10 L	7.4 %	Di Giovanni et al. (1999)
Disinfected reclaimed effluent	15	400 L	40 %	Gennaccaro et al. (2003)
Raw wastewater	9	4 L	22 %	Lalancette et al. (2012)
Secondary undisinfected waste- water effluents	10	4 L	30 %	Lalancette et al. (2012)
Raw wastewater	18	1 L	33 %	Gennaccaro et al. (2003)

Table 12.4 Prevalence of infectious Cryptosporidium oocysts in water

been used to detect infectious *Cryptosporidium* oocysts in raw wastewater, disinfected reclaimed effluent, raw source water, treatment plant filter backwash water, and finished drinking water (Table 12.4). Although cell culture-based methods for assessing *Cryptosporidium* infectivity have been standardized within individual laboratories, and detailed protocols have been developed, they have not yet been validated in large-scale multi-laboratory trials and there are no procedures approved for regulatory use. There are a variety of approaches that include different cell lines and media formulations, oocyst treatments prior to inoculation, incubation periods, and infection detection assays.

Optimized cell culture assays are very sensitive to infections with low levels of oocysts (Fig. 12.2) and have been used to assess the prevalence of infectious oocysts in finished water. A study utilizing cell culture to assess oocyst infectivity reported that 26.8 % of surface water treatment plants (N = 82) were releasing infectious oocysts in their finished water (Aboytes et al. 2004). Overall, 1.4 % of treated drinking water samples (N = 1,690) contained infectious *Cryptosporidium* oocysts but in all cases the follow-up repeat samples were negative. This detection rate translated into a calculated annual risk of 52 infections in 10,000 people, far exceeding the USEPA's 1 in 10,000 risk goal. In a study with contrasting results, treated drinking water samples from 14 treatment plants across the U.S. were analyzed using a modified version of Method 1623 coupled with a cell culture infectivity assay and immunofluorescence detection of infection (Rochelle et al. 2012). Sample volumes were 83.5–2,282 L with an average of 943 L (N = 370). None of the 370 finished water samples produced infections. This lack of infectious oocysts in a total volume of 349,053 L translated to an annual

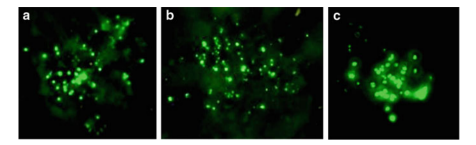


Fig. 12.2 Assay of oocyst infectivity in cultured cell monolayers. Clusters of intracellular life cycle stages (infectious foci) from infectious *C. parvum* oocysts recovered from large volume, spiked drinking water samples (**a** and **b**) and *C. hominis* from an unspiked treated wastewater sample (**c**). Human ileocecal adenocarcinoma (HCT-8) cell monolayers grown in multi-well chamber slides were inoculated with oocysts to allow the development of intracellular *Cryptosporidium* growth stages. Infected monolayers were fixed with methanol and stained with fluorescently-labeled antibodies that recognize sporozoites and other intracellular life cycle stages. Infectious foci were visualized using fluorescence microscopy

risk of less than one infection per 10,000 people. So, more research and more extensive sampling are needed to accurately determine the prevalence of humaninfectious oocysts in treated drinking water and thus improve the accuracy of risk assessments.

There have been several recent methodological advances in the detection of infectious Cryptosporidium in water. A method for enumerating both total and infectious oocysts in individual water samples was recently described (Lalancette et al. 2010). The method, referred to as 3D-CC-IFA, relies on dual direct detection using differential immunofluorescent staining to detect both oocysts and cell culture infection foci for each sample. This method helps address issues related to splitting of water samples, which typically contain low numbers of oocysts, between oocyst enumeration and cell culture infectivity assays (LeChevallier et al. 2003). The 3D-CC-IFA method was used to quantify the proportion of total Cryptosporidium oocysts in raw sewage and wastewater treatment plant effluents that were potentially human-infectious (Lalancette et al. 2012). Human-infectious Cryptosporidium were observed in 20 % of the samples. The maximum proportion of total oocvsts that were infectious was 22 % for raw sewage and 7 % for treated effluents (with one exception). Both of these values are well below commonly used quantitative microbial risk assessment values for the occurrence of infectious Cryptosporidium in water. Further application of the 3D-CC-IFA method will provide significant cost and labor savings and improved estimates of infectious oocyst fractions to aid risk assessments.

Numerous studies have examined the interplay between oocyst pretreatment, cell lines, media formulations, pH, infectivity periods, and other factors on the development of *Cryptosporidium* in cell culture. In particular, there are several, and sometimes conflicting reports in the literature examining the effects of bile salts and sodium taurocholate on oocyst excystation, sporozoite mobility, and susceptibility

of host cells to parasite invasion (e.g., Feng et al. 2006; Gold et al. 2001; Kato et al. 2001; King et al. 2012; Woodmansee et al. 1987). However, significant increases in the in-vitro development of *Cryptosporidium* have been reported using combination of cell monolayer growth medium with a high fetal bovine serum concentration (approximately 10 %) followed by centrifugal inoculation and infection development in medium with low (approximately 1 %) fetal bovine serum (King et al. 2012; B. King, personal communication). Additional research is needed in this area to fully understand these factors and develop standardized protocols that can then be used to gain a more accurate assessment of the prevalence of infectious oocysts in treated drinking water.

12.9 Conclusions

Waterborne cryptosporidiosis was recognized as a major problem for the drinking water industry after large outbreaks in the 1980s and 1990s. More recently, recreational water, particularly treated water venues such as swimming pools, has emerged as the primary waterborne route of transmission. Remarkably, there has been no drinking water outbreak linked to treated surface water for the last 2 decades in the U.S. (Yoder et al. 2012).² Nevertheless, routine detection in raw waters, occasional detection in finished water, resistance to chlorine disinfection, continued transmission in the community, and the large number of potential cases in the event of an outbreak, all continue to ensure that waterborne Cryptosporidium remains a matter of great concern to the water industry. While still not accurately quantified, the risks of endemic disease and outbreaks are clearly greater than zero as illustrated by recent outbreaks linked to drinking water in some countries and frequent swimming pool-related outbreaks. Some of the questions that still need to be answered regarding Cryptosporidium in water include: the relative contribution of drinking water to the overall burden of cryptosporidiosis compared to other sources of infection; the prevalence in source waters of human-infectious species or genotypes compared to species that do not infect humans; accurate measures of the frequency of oocyst breakthrough into treated water; the relative virulence of human-infectious species in water; and the contribution to population immunity of low-level endemic exposure to waterborne oocysts.

 $^{^{2}}$ The 20-year period without a cryptosporidiosis outbreak linked to a municipal surface water supply in the U.S. ended recently. An outbreak in Baker City, Oregon during the summer of 2013 was linked to an unfiltered surface water supply with up to 91 oocysts/L in one supply creek (www. bakercity.com; accessed Sept. 25, 2013).

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Chapter 13 Removal and Inactivation of *Cryptosporidium* **from Water**

Paul Monis, Brendon King, and Alexandra Keegan

Abstract Water is a major route of transmission for *Cryptosporidium* and oocysts commonly occur in surface and recreational waters as a consequence of fecal contamination from Wildlife or anthroponotic sources. There are many characteristics possessed by *Cryptosporidium* oocysts that allow them to persist in aquatic environments, including recreational waters, and to bypass water treatment processes. These types of events lead to outbreaks of cryptosporidiosis, caused by direct exposure to contaminated recreational water (such as swimming pools) or by drinking contaminated potable water. Previous chapters have discussed the epidemiology of *Cryptosporidium* in relation to waterborne transmission and also the sources and presence of oocysts in drinking and recreational waters. This chapter will review the processes contributing to the removal and inactivation of *Cryptosporidium* oocysts from surface waters and wastewaters, including natural processes that occur in surface waters and engineered processes used for the production of drinking water or for the treatment of wastewater.

13.1 Measurement of Inactivation

In order to review the removal and inactivation of *Cryptosporidium* in water, it is important to first define key terms, such as viability and infectivity, and to briefly review the methods used to measure inactivation. In general terms, "viability" is loosely taken to indicate that an organism is capable of living or developing under favorable conditions. Many techniques that measure viability at the level of a cell provide general indicators of cell health, such as membrane integrity or enzyme activity. Infectivity is more specifically taken to indicate that an organism is able to

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survive and multiply in a host. In the case of *Cryptosporidium*, viability measurements have historically been used as surrogates for infectivity, but as will be shown below, there are cases where this assumption has been proven to be invalid. In the context of protecting public health, it is critical to understand how treatment processes affect oocyst infectivity.

Animal bioassays have been considered the "gold standard" for assessing Cryptosporidium oocyst infectivity. The first description of experimental infection in mice using Cryptosporidium oocysts from a diarrheic calf found that neonatal mice were readily infected but that 21 day old mice were only mildly infected (Sherwood et al. 1982). The study of Sherwood et al. (1982) also provided the first evidence that oocysts were susceptible to freezing but retained infectivity for 4-6 months at 4 °C. The neonatal mouse model has since been used extensively in the assessment of Cryptosporidium parvum oocyst inactivation by a number of mechanisms (Anderson 1985, 1986; Peeters et al. 1989; Korich et al. 1990; Finch et al. 1993a, b; Fayer 1994, 1995; Black et al. 1996; Fayer et al. 1996; Harp et al. 1996; Xunde and Brasseur 2000; Araki et al. 2001; Craik et al. 2001; Biswas et al. 2003; Li et al. 2004, 2005; Garvey et al. 2010), which will be discussed elsewhere in this chapter. Mouse infectivity has been shown to be reproducible under standardized conditions using a logistic dose response model (Korich et al. 2000). Aside from practical, cost and ethical issues around the use of animals for measuring infectivity, the mouse model has limited applicability because the most common species that infects humans, Cryptosporidium hominis, cannot infect mice (Morgan-Ryan et al. 2002). Gnotobiotic piglets have been described for the propagation of C. hominis (Widmer et al. 2000; Akiyoshi et al. 2002) and for measuring drug efficacy against C. hominis (Theodos et al. 1998) but such a system is even less practical and more costly than neonatal mice. More recently, an animal model using immunosuppressed gerbils has been described for the propagation of C. parvum and C. hominis (Baishanbo et al. 2005) and also for C. muris and C. andersoni (Kvac et al. 2007, 2009). The gerbil model has been used to study drug efficacy against C. parvum (Baishanbo et al. 2006) but does not appear to have been further used to evaluate the effectiveness of disinfectants against C. hominis.

Viability assays have long been pursued as surrogates for measuring infectivity, being cheaper, simpler, faster and more amenable to high-throughput quantitative analysis than using animal models of infectivity. Vital dyes were shown in an early study to be poor predictors of viability (Korich et al. 1990), although excystation correlated with mouse infectivity for measuring inactivation by oxidant-based disinfectants (Korich et al. 1990). However, according to Korich et al. (1990), reliable use of excystation required sporozoite counts, rather than oocyst counts, since high doses of disinfectants resulted in empty oocysts that could be confused with mouse infectivity for assessing the effect of chemical disinfectants found that the in vitro viability assays consistently underestimated inactivation (Finch et al. 1993a; Black et al. 1996). Although vital dyes such as propidium iodide have been shown to poorly correlate with mouse infectivity, nucleic acid dye staining has been shown to correlate with mouse infectivity for chemical disinfectants

(Belosevic et al. 1997; Neumann et al. 2000). However, inter-laboratory comparison of in vitro viability assays, including dye exclusion, nucleic acid binding dyes and excystation, with mouse infectivity found that the in vitro techniques could not reliably predict inactivation following ozone treatment (Bukhari et al. 2000). While some agreement has been reported between viability techniques and animal infectivity for chemical disinfectants, the same has not been found for alternative methods of inactivation such as exposure to ultraviolet light. Excystation has been shown to greatly underestimate inactivation by UV compared with animal infectivity (Morita et al. 2002). When compared with cell culture infectivity, viability methods have also been shown to underestimate inactivation caused by oocyst ageing at 15 °C (Schets et al. 2005). Detection of ribosomal RNA using fluorescent in situ hybridization (FISH) has also been reported as a viability indicator to measure inactivation of oocysts in soil (Davies et al. 2005). The reported oocyst inactivation rate at 35 °C was only 2 log₁₀ after 25 days (Davies et al. 2005), compared with 4 \log_{10} within 4 days at 37 °C and 4 \log_{10} after 14.6 days at 30 °C measured using cell culture infectivity (King et al. 2005), suggesting that the FISH technique is a poor surrogate for infectivity when measuring metabolic exhaustion of oocysts. While cost effective and simple to use, in vitro viability assays should be used with caution because of the inconsistent agreement with animal infectivity measures, and such assays should only be used when validated against an infectivity assay for a particular mode of inactivation.

An in vitro alternative to viability assays is the use of cell culture systems to measure oocyst infectivity. Complete development of Cryptosporidium was first reported using cultured human fetal lung cells and primary cultures of chicken and pig kidney cells (Current and Haynes 1984). However, while this system supported the production of sporulated oocysts it did not appear to be adopted for further use, possibly because of the requirement for primary-derived cells, with mouse infectivity being used predominantly in the 1980s. In subsequent years, a number of cell lines were evaluated for their ability to support *Cryptosporidium* development, but not as quantitative assays for measuring infectivity. For example, a canine kidney cell line (MDCK) was reported to support high rates of infection but did not produce sporulated oocysts (Gut et al. 1991), while a human endometrial carcinoma cell line was reported to support the complete development of Cryptosporidium in vitro (Rasmussen et al. 1993). An extensive comparison of cell lines suggested that HCT-8 cells (a human ileocecal adenocarcinoma cell line) supported the production of the highest number of parasite developmental stages (Upton et al. 1994b). The timing of sporozoite release from oocysts was also shown to be critical for successful infection of cell monolayers, with highest infection achieved when oocysts excysted when in contact with the monolayer (Upton et al. 1994a). While various cell lines have continued to be evaluated (Yang et al. 1996; Lawton et al. 1997; Deng and Cliver 1998; Lacharme et al. 2004), HCT-8 cells are currently the most commonly used for measuring in vitro infectivity.

Different end-points have been used for detecting cell infectivity. PCR-based assays have been used for end-point measurement of infection of cell monolayers either qualitatively (Rochelle et al. 1997; Di Giovanni et al. 1999) or through the

use of quantitative PCR to measure the relative change in infectivity due to exposure to disinfectants or environmental exposures (Keegan et al. 2003; King et al. 2005). A chemiluminescence immunoassay has also been described for measuring Cryptosporidium proliferation during cell infection (You et al. 1996). Possibly the most useful end-point measurement has been the application of fluorescent antibodies to detect intracellular developmental stages of Cryptosporid*ium*, allowing the detection of infectious foci (Slifko et al. 1997). In this method, a single focus of infection is caused by the localized infection of nearby cells by sporozoites released from a single oocyst followed by the subsequent infection of neighboring cells by other life-cycle stages. The focus detection method (FDM) allows quantitation of infectious oocysts in a sample (assuming each focus arises from a single occyst). If the input number of occysts is know the FDM also allows measurement of the proportion of infectious oocysts in a sample. For FDM to be accurate it is critical to control oocvst excystation so that sporozoites are released when the oocyst is on the monolayer to prevent dispersal of sporozoites, which could result in multiple foci from a single oocyst (King et al. 2011, 2012). An alternative approach for the detection of intracellular stages has been the use of in situ hybridization with a Cryptosporidium-specific oligonucleotide probe (Rochelle et al. 2001), although this technique does not appear to be in widespread use. A recent comparison of infectivity methods for both C. parvum and C. hominis has shown that the FDM method is less prone to the detection of false positives compared with a PCR end-point for infectivity detection and that FDM was better at detecting infectious oocysts spiked into water samples (Johnson et al. 2012). The robustness of cell culture infectivity using HCT-8 cells and FDM has been demonstrated in an interlaboratory trial (Bukhari et al. 2007).

Multiple comparison studies of in vitro cell culture infectivity assays with mouse infectivity have been conducted and these methods shown to be equivalent (Rochelle et al. 2002; Slifko et al. 2002; Garvey et al. 2010; Alum et al. 2011), although not for assessing the efficacy of some anti-cryptosporidial drug treatments (Theodos et al. 1998). A comparison of in vitro infectivity for three cell lines measured using a reverse-transcription PCR assay with CD-1 mouse infectivity found that HCT-8 cells gave the best correlation with the "gold standard" for oocysts exposed to UV light and ozone (Rochelle et al. 2002). A similar study compared HCT-8 infectivity using the focus detection method with BALB/c mouse infectivity for dose response and response to UV and chlorine dioxide disinfection (Slifko et al. 2002). The results were not significantly different, although it was suggested that cultured cells might allow more sensitive detection of infection compared with measuring infection in mice (Slifko et al. 2002). A HCT-8 infectivity assay using qPCR also provided comparable results with a SCID mouse assay for UV inactivation (Garvey et al. 2010). Considering that these studies used different cell culture infection measurement end-points and different mouse strains, the correlation between cell culture and mouse infectivity can be considered to be robust.

13.2 Removal and Inactivation by Natural Processes

13.2.1 The Terrestrial Environment

13.2.1.1 Heat

Oocysts can enter surface waters when infected animals defecate directly into creeks or streams or as a result of discharge from sewage systems or sewage treatment plants. However, in many instances fecal deposition is directly onto the ground, so the terrestrial environment often poses the first barrier to the entry of oocysts into surface waters. Dispersal from animal feces relies on rainfall to release oocysts from the fecal matrix and wash them into surface waters (Davies et al. 2004), and this mobilization is one of the major factors influencing the output of process-based models for oocyst fate in catchments (Ferguson et al. 2007). While in the terrestrial environment, oocysts can be exposed to stresses such as temperature and desiccation, which are particularly effective at inactivating oocysts. Experiments using moist heat demonstrated complete oocyst (presumably C. parvum) inactivation, measured using mouse infectivity, for exposure times between 5 and 20 min at 45 °C, although details of oocyst doses and oocyst infectivity before heat treatment were not provided (Anderson 1985). Mouse infectivity data have also shown that C. parvum oocysts suspended in water were completely inactivated after 1 min at 72 °C and 2 min at 64 °C (Fayer 1994). In the study by Fayer (1994), the oocyst dose administered to each mouse was 1.5×10^5 and considering that the oocysts were at most 1 month old and isolated directly from experimentally infected calves, it would be reasonable to infer a reduction in infectivity of at least 4 log₁₀ units. Similar results were obtained using a pasteurization temperature of 71.7 $^{\circ}$ C for 5–15 s, resulting in complete inactivation using doses of 10^5 treated oocysts administered to mice (Harp et al. 1996). Sensitivity to high temperature appears to be common to Cryptosporidium species, with C. parvum, C. muris and a Cryptosporidium spp. isolated from a chicken all exhibiting complete inactivation after 15 s at 60 °C or 30 s at 55 °C using a dose of 10⁶ oocysts into mice for C. parvum and C. muris or 2-week-old chickens for the Cryptosporidium spp. (Fujino et al. 2002). These temperatures can occur in animal feces exposed to sunlight, with bovine fecal material shown to exhibit temperature peaks between 40 °C and 70 °C once air temperature exceeds 25 °C (Li et al. 2005). Using diurnal temperature cycles typical of spring – autumn conditions for California rangelands, an inactivation rate of $3.27 \log_{10}/day$ was observed using a mouse infectivity model (Li et al. 2005). In these experiments, Li et al. (2005) observed that the primary loss of infectivity appeared to be due to partial or complete excystation of oocysts in the fecal matrix. The rate of loss of infectivity was much slower for winter diurnal temperatures, with an internal fecal matrix temperature of 30 °C resulting in an inactivation rate of 0.2 \log_{10}/day and a rate of 0.03 \log_{10}/day at 20 °C (Li et al. 2010).

13.2.1.2 Freezing and Desiccation

While oocysts are sensitive to heat, they are able to withstand short periods of freezing, surviving days at -10 °C and hours at -15 °C (Fayer and Nerad 1996). However, oocysts have been shown to be susceptible to freeze/thawing cycles particularly in soil (Jenkins et al. 1999; Kato et al. 2002), where soil particles cause abrasion and fragmentation of oocysts (Jenkins et al. 1999). Oocysts have been shown to be particularly sensitive to desiccation, which is significant in a terrestrial environment. Desiccation can occur within the fecal matrix, as a consequence of fecal processing by insects such as dung beetles (Ryan et al. 2011) or once oocysts have been washed from the fecal matrix into soil. Loss of infectivity from desiccation can be relatively rapid, with oocysts in calf feces dried for as little as 1 day losing infectivity for neonatal mice (Anderson 1986). Within 2 h of air-drying on slides, 97 % of oocysts were dead (as measured by vital dye staining). Mortality increased to 100 % of oocysts after 4 h (Robertson et al. 1992). Similar results were obtained when assessing oocyst survival following desiccation on stainless steel surfaces (Deng and Cliver 1999). Aside from desiccation, the physical, chemical and biological properties of soil may affect oocyst survival (Ferguson et al. 2003). Viability assays have shown that oocyst survival is affected by soil type (Jenkins et al. 2002; Davies et al. 2005), in particular soil texture and possibly low soil pH (Jenkins et al. 2002). In these studies temperature was suggested as a major factor affecting oocyst inactivation, and soil moisture appeared to have no correlation with loss of viability (Jenkins et al. 2002; Davies et al. 2005). However, a study evaluating the effect of water potential (using osmolarity as a surrogate) suggests that soil moisture content will greatly influence the rate of oocyst degradation. The study of Walker et al. (2001) used water potentials representing soil moisture contents required to support normal crop growth, the wilting point (the minimum soil moisture content required to prevent plant wilting) and extremely dry conditions. Water potential stresses were found to combine with temperature effects and freeze-thawing to enhance oocyst degradation. Degradation was likely to be more rapid in dry soil compared with calf feces or in water at low temperatures (Walker et al. 2001). A more recent study using cell culture infectivity provided further support for the findings of Walker et al. (2001), suggesting that saturated loamy soil did not affect oocyst inactivation compared with oocysts suspended in distilled water, with both resulting in 0.93 log₁₀ inactivation after 10 days at 30 °C, whereas dry loamy soil had a greater effect, with 2.5 log₁₀ inactivation after 10 days at 32 °C (Nasser et al. 2007). Furthermore, the study of Nasser et al. (2007) compared viability and infectivity measures for the different test conditions and found that the viability method overestimated oocyst survival by 1-2 log₁₀, measuring no loss of infectivity at 30 °C and only 0.5 log₁₀ inactivation at 32 °C.

13.2.1.3 Ammonia

Ammonia can be present at high levels in decomposing feces, especially in manure storages (Muck and Steenhuis 1982; Muck and Richards 1983; Patni and Jui 1991), ranging from 0.05 M, rising to 0.2 M in cattle slurries depending on storage time (Whitehead and Raistrick 1993). Oocysts have been shown to be sensitive to ammonia, initially in a study investigating gaseous disinfection of oocysts using different compounds (Fayer et al. 1996). In that study, oocysts suspended in water were exposed to one atmosphere of pure ammonia at 21–23 °C for 24 h and were rendered non-infectious to mice. Another study used excystation and vital dye staining assays to assess the effect of a wide range of ammonia concentrations (0.007–0.148 M) on oocyst viability, finding that all concentrations reduced viability and calculating that 0.06 M ammonia would inactivate 99.999 % oocysts in 8.2 days at 24 °C (Jenkins et al. 1998). The inactivation rate was temperature dependent, taking much longer (55 days) at 4 °C. Considering the sensitivity of oocysts to ammonia, it is likely that some oocyst inactivation will occur from this source in animal feces and that practices such as long-term storage of animal waste will effectively reduce Cryptosporidium risk, even at low temperatures (Hutchison et al. 2005).

13.2.1.4 Transport into Receiving Waters

Oocysts need to be mobilized from soil surfaces/sub-surfaces into receiving waters. There are conflicting reports of the interaction between oocysts and soil particles. Oocysts have been reported not to attach to soil particles in water (Dai and Boll 2003) and to behave as single particles, rather than aggregates (Kaucner et al. 2005; Brookes et al. 2006). However, in the context of agricultural runoff, significant attachment of oocysts was shown for clay loam and for both clay loam and sandy loam in the presence of manure (Kuczynska et al. 2005). Irrespective of adsorption/ desorption, soils can greatly limit the mobility of pathogens by acting as a filter (Tufenkji et al. 2004) and for this reason the preferred method for disposal of human waste in wilderness areas is by digging small holes and burying feces (Cilimburg et al. 2000). Retention in soils has been shown to be a function of surface roughness and not to be affected by soil organic matter (Santamaria et al. 2012). A laboratory study using simulated rainfall and soil cores for three contrasting soil types (clay loam, silty loam, sandy loam) found that the distribution of Cryptosporidium parvum oocysts was similar for all three soils, with the majority of oocysts (73 %) in the top 2 cm of soil (Mawdsley et al. 1996). However, oocysts have been shown to rapidly pass through saturated macroporous soils, although oocyst losses through the soil columns were 90-99.9 % (Darnault et al. 2003), similar to that reported by Kuyczynska et al. (2005). The presence of the majority of oocysts near the soil surface suggests that they could be readily mobilized during a precipitation event. Removal of oocysts through soil appears to be variable and the

observed transport not consistent with colloid theory (Santamaria et al. 2011). The variability in removal, particularly as a function of travel distance, has been suggested to be due to variability within the population of oocysts (Santamaria et al. 2011). The degree of agricultural land use can also affect transport, particularly the use of tile drainage, which is used to remove excess water from the soil sub-surface to improve crop growth, minimize erosion and reduce the impact of livestock on land. In the case of land with tile drainage, pathogen transport (including *Cryptosporidium*) has been shown to be predominantly via the tile drainage system and not by surface transport, except on rare occasions (Dorner et al. 2006).

Soil is not the only factor affecting oocyst mobilization, with vegetation proving to be effective at reducing *Cryptosporidium* numbers in surface runoff, causing a reduction of up to 1.44 log_{10} per meter of vegetated buffer (Tate et al. 2004). The combination of soil type and vegetated buffer influences oocyst removal, with buffers on sandy loam being less effective at removing Cryptosporidium oocysts $(1-2 \log_{10} \text{ reduction/meter})$ compared with vegetation on silty clay or loam (2-3)log₁₀ reduction/meter) (Atwill et al. 2002). Rainfall simulation using artificial 1 kg cow fecal pats spiked with 10^7 Cryptosporidium oocysts on a loam soil (49 % sand, 27 % silt, 24 % clay) demonstrated that vegetation promoted infiltration (compared with a bare soil surface) and that there were fewer oocysts $(1-4 \log_{10} \text{ lower})$ in vegetated runoff compared with bare runoff. In these simulations, the number of oocysts detected in the vegetated runoff and infiltrate was much lower than that in the runoff and infiltrate from the bare soil (Davies et al. 2004). A more extensive study examining a wider range of slopes, vegetation level and rainfall conditions found that 2-5 log₁₀ oocysts were retained in soil without buffer, but that 3-8.8 \log_{10} oocysts were retained by soils with grassland buffers (Atwill et al. 2006). Considering that infected calves can excrete 6×10^7 oocysts per gram of feces (Uga et al. 2000) and up to 10^{10} occysts per day for up to 14 days during the infection (Meinhardt et al. 1996), significant numbers of oocysts could still be mobilized into the aquatic environment, despite attenuation through soils and vegetation.

13.3 Surface Waters

13.3.1 Hydrological Parameters

Surface waters can provide an excellent environment for supporting the survival of oocysts, providing a thermal buffer from temperature extremes (in contrast to the terrestrial environment) and also providing a medium for transmission to and ingestion by a susceptible host. However, there are biotic and abiotic factors that can cause removal or inactivation of oocysts in an aquatic environment and also

processes that control transport and distribution within a water body (Brookes et al. 2004). Key factors include particle interaction, temperature, sunlight (UV) exposure and grazing, all of which are affected by the hydrodynamics of a stream or reservoir (e.g. dispersion, dilution) (Brookes et al. 2004). Oocvst transport within lakes/reservoirs is predominantly driven by inflows, particularly those from rainfall events, and also by circulation patterns from wind-generated currents and internal waves (Brookes et al. 2004). Inflows are considered to be major sources of pathogens for reservoirs. The behavior of an inflow is controlled by the inflow density relative to that of the receiving water, with water temperature being a major factor affecting transport of oocysts. Inflow water that is warmer than the receiving water (and so less dense) will travel across the top of the receiving water surface, whereas inflow water that is colder will be more dense and flow along the bottom of the receiving water (Brookes et al. 2004). Entrainment of the receiving water by the inflow leads to dilution of oocyst concentrations. Inflow temperature, velocity, insertion depth and entrainment rate are critical factors determining the hydrodynamic distribution of oocysts in lakes and reservoirs (Brookes et al. 2004).

Sedimentation rate, along with flow velocity, is a key factor affecting the position of oocysts within a water column. Oocyst position can influence survival, oocysts at the surface can be exposed to stressors such as sunlight, whereas oocysts at depth or in sediments may be protected from sunlight but may be subjected to microbial interactions via predation or interaction with biofilms. Empirically determined oocyst sedimentation rates have been shown to closely match theoretical calculations based on oocyst size and density, being measured to be 0.27-0.35 µm/s (Medema et al. 1998; Dai and Boll 2006). Attachment to particles or entrapment within an organic matrix will greatly affect sedimentation rates (Searcy et al. 2005), but as discussed in the previous section, there have been conflicting reports on the interaction of oocysts with particles and characteristics of the water such as type and concentration of organic matter, as well as the nature of the sediment contributing particles, may influence oocyst attachment to particulates. In the context of wastewater, oocysts have been shown to readily attach to effluent particles (Medema et al. 1998), so oocysts in water resulting from sewage effluent discharge may behave differently compared with oocysts from land runoff. While the sedimentation velocity of single oocysts is too low to result in significant settling in a large water body, the rate can increase dramatically (28.9 μ m/s) for oocysts attached to particles (Medema et al. 1998). A study of the oocyst loss rate in Lake Burragorang (a water storage in Sydney, Australia) measured a high sedimentation rate, $57.9-115.7 \mu m/s$, much higher than the laboratory-measured values, and proposed that oocyst aggregates with other oocysts or particles was responsible for the accelerated removal (Hawkins et al. 2000). The results of Hawkins et al. (2000) suggest that transport under field conditions can vary dramatically from those observed under laboratory-simulated conditions. Factors that may not be accounted for are the effect of turbulence in inflows on oocyst interaction/aggregation with particles and the influence of processes within a water body, such as internal waves, which can cause rapid movement of oocysts through the water column (Brookes et al. 2004). Sediment re-suspension events, caused by turbulence from underflows or internal waves (Michallet and Ivey 1999), can remobilize oocysts, which will

still pose a water quality challenge in drinking water service reservoirs. Storm events have also been shown to cause significant re-suspension of pathogens in river sediments (Dorner et al. 2006).

13.3.2 Predation and Microbial Activity

Oocysts are exposed to a variety of aquatic environments where they may come into contact with other microorganisms that are antagonistic or will reduce oocyst persistence. Despite the potential importance of biological interactions for the removal of oocysts in natural and engineered systems, relatively little has been published in this field. One of the first indications of the potential role of biological interactions was the study of Chauret et al. (1998), who observed better survival of oocysts in 0.2 µm-filtered river water compared with unfiltered river water and concluded that this was due to biological antagonism (Chauret et al. 1998). It is unclear whether the antagonism was due to predation or other microbial interactions that resulted in decreased oocyst viability (measured by excystation). Predation of oocysts by a free-living nematode (*Caenorhabditis elegans*) has been demonstrated in soils (Huamanchay et al. 2004) and interaction with Pseudomonas aeruginosa extracts (modeling soil exposure) led to degradation of the oocyst wall, reducing oocyst infectivity (Nasser et al. 2007). Ingestion of oocysts by C. elegans did not reduce the infectivity of excreted oocysts (Huamanchay et al. 2004). A variety of aquatic organisms have been shown to be capable of ingesting oocysts, including rotifers (Fayer et al. 2000; Stott et al. 2003; King et al. 2007), ciliates (Stott et al. 2001, 2003; King et al. 2007), amoebae (Stott et al. 2003; King et al. 2007), gastrotrichs (King et al. 2007) and platyhelminths (King et al. 2007). Examples are shown in Figs. 13.1, 13.2, 13.3, 13.4, 13.5, and 13.6 (reproduced from (King et al. 2007), http://www.wqra.com.au/publications/document-search/? download=81).

The effect of ingestion on oocyst infectivity has not been assessed. However, Fayer et al. (2000) noted that rotifers excreted oocysts in boluses and King et al. (2007) also identified oocyst clumping following biotic activity. As discussed above, aggregated oocysts will settle faster and so may be less mobile. Excreted oocysts, which may have been exposed to proteolytic digestion and also may be coated in other material as a consequence of the ingestion process, may also interact with particles differently, which may further increase sedimentation and decrease opportunity for further transport through a water body. In feeding experiments conducted by King et al. (2007), it was observed that ingestion by some predators resulted in the degradation of oocysts in food vacuoles (e.g. Fig. 13.1). One limitation of feeding experiments is that in general high oocyst numbers have been used (e.g., greater than 10^4 oocysts/ml) whereas in environmental waters oocyst concentrations are very low, at most 1–10 oocysts L under base-flow conditions (Brookes et al. 2004). In the environment many of the predators are size selective feeders and *Cryptosporidium* oocysts would only be an incidental

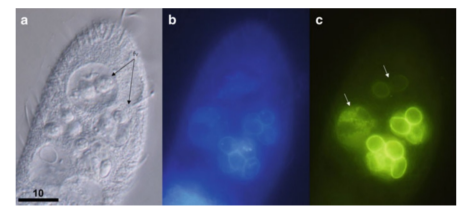


Fig. 13.1 *Paramecium* sp. enlarged to show oocysts within the food vacuoles. (a) DIC, (b) DAPI staining under UV, (c) oocysts labelled with a FITC antibody under *blue light*. Bar = $10 \mu m$

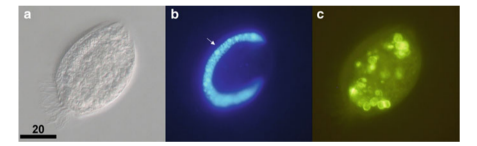


Fig. 13.2 *Euplotes* sp. containing ingested oocysts of *Cryptosporidium parvum*. The arrow indicates the *horseshoe shaped* macronucleus of the cell. (a) DIC, (b) DAPI staining under UV, (c) oocysts labelled with a FITC antibody under *blue light*. Bar = 20μ m

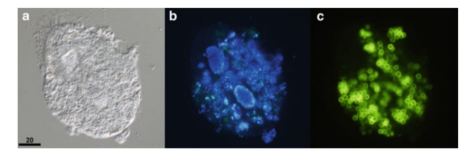


Fig. 13.3 Oxytricha sp. containing oocysts of *Cryptosporidium parvum*. (a) DIC, (b) DAPI staining under UV, (c) oocysts labelled with a FITC antibody under *blue light*. Bar = $20\mu m$

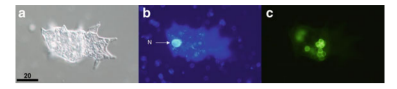


Fig. 13.4 *Mayorella* sp. containing oocysts of *Cryptosporidium parvum*. The amoeba displays conical pseudopodia (*black arrow*), which are a classifying feature of the species (**a**). The nucleus (N) of the amoeba is highlighted by the DAPI staining (**b**). (**a**) DIC, (**b**) DAPI staining under UV, (**c**) oocysts labelled with a FITC antibody under *blue light*. Bar = 20μ m

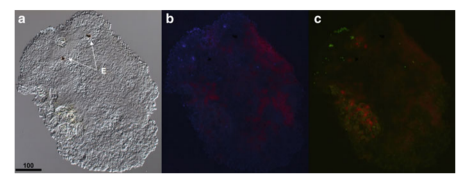


Fig. 13.5 Rhabdocoel platyhelminth containing ingested oocysts of *Cryptosporidium parvum*. Contraction and lysis of the organism occurred upon fixation, but it is still possible to determine whether the organism has ingested oocysts. (a) DIC, (b) DAPI staining under UV, (c) oocysts labelled with a FITC antibody under *blue light*. Bar = 100μ m

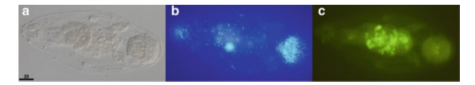


Fig. 13.6 Bdelloid rotifer (unidentified genus) containing ingested oocysts of *Cryptosporidium* parvum. (a) DIC, (b) DAPI staining under UV, (c) oocysts labelled with a FITC antibody under blue light. Bar = $20\mu m$

food source amongst the many other organisms (such as yeast, algae and bacteria) in that size range in water. Therefore, food density would be sufficient to sustain predator numbers and oocysts in aquatic environments rich in predators (such as sediment water interfaces or biofilm surfaces) may be effectively removed.

Another mechanism for removal and release of oocysts in water is attachment to biofilms. Flow chamber experiments, using biofilms established from environmental biofilm microorganisms collected during different seasons, have been used to compare attachment and detachment of oocysts (Wolyniak et al. 2010). Oocysts were found to rapidly attach and reach a steady state with between 46 % and 77 %of oocysts attached to the biofilm during oocyst challenge, with steady detachment of oocysts once the oocyst challenge had been removed to a lower steady state of 24–65 % of oocysts retained within the biofilm (Wolyniak et al. 2010). Biofilms collected at different seasons had different levels of oocyst retention, with the spring biofilm retaining the highest proportion of oocysts (65 %) and the autumn biofilm retaining the fewest (24 %). Oocyst retention appeared to be a specific function of the microbial community harvested and was not affected by temperature (5 °C or 20 °C), biofilm thickness or water characteristics, with water harvested during autumn not reducing retention by the spring biofilm and water harvested during spring not increasing retention by the autumn biofilm (Wolyniak et al. 2010). The steady-state attachment and persistence of oocysts in natural wastewater biofilms has also been demonstrated for (Skraber et al. 2007) and drinking water and wastewater biofilms established in annular reactors (Helmi et al. 2008). The attachment and release of oocysts by natural biofilms is in contrast with earlier reports using a biofilm established with P. aeruginosa, which strongly retained oocysts even under high flow conditions (Searcy et al. 2006). An extension of the study by Wolyniak et al. (2010) found that seasonal differences in oocyst retention were consistent across different sites but not across different years, and that oocyst retention correlated with biofilm roughness. Challenging biofilms with increasing numbers of oocysts demonstrated that saturation could be achieved, suggesting that the number of attachment sites for oocysts is limited (DiCesare et al. 2012b). Biofilms in shallow waters may play a role in protecting oocysts from inactivation by solar radiation, with oocysts embedded at the bottom of a biofilm shown to retain infectivity up to two times better compared with oocysts at the top of a biofilm or oocysts with no biofilm in solar radiation experiments (DiCesare et al. 2012a).

Another potential source of oocyst removal in freshwaters are bivalve molluscs, which have been proposed as biological indicators of Cryptosporidium in river water (Graczyk et al. 2001; Izumi et al. 2006). Zebra mussels collected from a river were shown to contain significant numbers of oocysts, up to 220/g tissue (Graczyk et al. 2001). Experimental exposure of the clam *Corbicula japonica* to oocysts found that oocysts were excreted in the clam feces at high rates, with 1-3 % of the input oocysts retained in tissue (based on the challenge doses, similar numbers in tissue compared with the report of Graczyk et al. 2001) (Izumi et al. 2006). Passage through the clams had no effect on oocyst infectivity, as measured using cell culture infectivity. Clams placed in a river were able to concentrate oocysts, which were isolated and identified by genotyping to be C. parvum (Izumi et al. 2006). The high clearance rate reported suggests that detection of low numbers of oocysts in clam tissue may represent previous contamination events, and high numbers of oocysts representative of recent events. Infectious oocysts have been recovered from marine molluscs harvested for food (Gomez-Bautista et al. 2000). While filterfeeding molluscs may play a role in the removal of oocysts in freshwater systems, there appears to have been little research activity in this area (aside from use as a sentinel organism), with research activity predominantly in the area of oocyst contamination of commercially harvested shellfish.

13.3.3 Inactivation by Temperature

The temperatures in aquatic environments are generally more moderate compared with the high temperatures that can be encountered in a terrestrial environment. While inactivation at high temperatures is rapid, particularly at temperatures above 40 °C, it can be relatively slow at temperatures more relevant in water bodies. Oocysts have been shown capable of retaining infectivity for at least 24 weeks at 4 °C and 16–24 weeks at 15 °C (Fayer et al. 1998; Keegan et al. 2008), with at most a 1 log₁₀ reduction in infectivity. Using mouse infectivity, Fayer et al. (1998) found that oocysts retained infectivity after 24 weeks stored at 20 °C and 12 weeks stored at 25 °C, but that the number of mice infected and the level of infection was greatly reduced compared with control oocysts. Quantitative measurement of inactivation using cell culture infectivity found more than $3 \log_{10}$ inactivation of oocysts after 12 weeks stored at 20 °C, 8 weeks stored at 25 °C and 3 weeks stored at 30 °C (King et al. 2005). In cooler climates, temperature is unlikely to play a role in oocyst inactivation unless the detention time in water is extremely long. However, depending on reservoir hydrodynamics and oocyst detention times, tropical and possibly sub-tropical climates may well influence oocyst survival in the aquatic environment.

The mechanism responsible for oocyst inactivation, at least at temperatures of 37 °C or less, has been shown to be metabolic exhaustion. *Cryptosporidium* oocysts contain finite carbohydrate energy reserves, stored as amylopectin, which are required to initiate infection and are consumed due to metabolic activity at ambient temperatures (Fayer et al. 1998). Oocyst energy levels (measured as ATP) have been shown to closely correlate with retention of infectivity in cell culture for temperatures between 4 °C and 37 °C (King et al. 2005). The finite energy reserve within sporozoites is a key factor in determining host cell infection success, meaning that the timing of sporozoite release is critical and must be controlled for optimal measurement of infectivity using cell culture models (Upton et al. 1994b; King et al. 2011).

13.3.4 Inactivation by Sunlight

Sunlight is possibly the most potent environmental stressor in the aquatic environment. The most damaging components of sunlight are short-wavelength ultraviolet (UV) light, specifically UV-B (280–320 nm) and UV-A (320–400 nm), with UV-B principally causing DNA damage and UV-A causing damage to a variety of cellular components including lipids and proteins (Caldwell 1971; Friedberg et al. 1995; Malloy et al. 1997; Ravanat et al. 2001). UV exposure can be detrimental to a variety of organisms ranging from bacteria to plants in both terrestrial and aquatic environments (King and Monis 2007).

In terms of Cryptosporidium in water, the bulk of research appears to have focused on measuring the effectiveness of solar disinfection (SODIS) for oocyst inactivation, which combines solar irradiation with temperature inactivation and may have variable wavelengths of UV involved, depending on the UV transmissivity of the containers used. However, several studies have assessed the effectiveness of solar radiation on oocyst infectivity under environmentally relevant conditions. An early study utilized excystation to measure a 90 % reduction in oocyst viability following a 3-day exposure to sunlight in marine water (Johnson et al. 1997). More recently, solar radiation and artificial UV-B have been shown to be effective at reducing the infectivity of oocysts suspended in phosphate-buffered saline (Connelly et al. 2007), although it was not clear how inactivation was calculated in that study. A dose of 66 kJ/m² (measured at 320 nm) resulted in a 97.5 % reduction in infectivity compared with the dark control, and a claimed >99.99 % loss of infectivity was observed following 10 h exposure to sunlight on a mid-summer day. However, some of the calculated reductions may not be accurate, based on the data presented (triplicate samples of 5,000 oocysts used in solar radiation experiments, each replicate analyzed in triplicate using 100 oocysts per analysis, measuring infectivity from a total of 900 oocysts) At most a 3 \log_{10} inactivation (99.9 %) could be measured if all of the oocysts were infectious at the start of the experiment (but at most 22 % were infectious in the controls). Additionally, there was also between two and almost tenfold difference in the infectivity measured for dark and laboratory control oocysts, suggesting either large variability in the measured infectivity, or some other effect on infectivity aside from solar radiation (temperature can be excluded because it did not exceed a maximum of 22.5 °C on any of the testing days). The authors noted the high variability in some samples and suggested it could be due to differences within the oocyst population, oocyst clumping (causing underestimation of infectious foci) or variability in the quality/density of the cell monolayer (Connelly et al. 2007). Irrespective of these possible shortcomings of the cell assay or inactivation calculations, the findings clearly demonstrated the impact of solar radiation on oocyst infectivity.

Another study assessed the effect of solar radiation on the survival of *C. parvum* oocysts suspended in tap water or environmental waters, using a cell culture infectivity assay (King et al. 2008). These experiments were conducted on days with different levels of solar insolation, measured using a pyranometer, and compared with the UV index. The UV index is a measure developed by the World Health Organization to raise public awareness of the risk of UV radiation and provides an indication of the daily danger of solar UV radiation intensity (http://www.who.int/uv/intersunprogramme/activities/uv_index/en/index.html). It is calculated at solar noon, when the sun's radiation is most intense, with each point on the index equivalent to 25 mW/m² of UV radiation 290 nm and 400 nm. The water quality ranged from 2.8 mg/L dissolved organic carbon (DOC), 0.23 nephelometric turbidity units (NTU) turbidity and a color of 2 Hazen units (HU) for the tap water, to maximums of 12.3 mg/L DOC, 7.42 NTU turbidity and 77 HU color for the reservoir waters. For days with UV indices of 4 or higher, reductions in

infectivity of between 2.2 and 3 \log_{10} were achieved within a single day of exposure (King et al. 2008). The inactivation rates on different days were compared using T_{90} , which is the time (hours) required to achieve 90 % inactivation. The T_{90} was larger for low UV index days (e.g. 6.4 for a day with a UV index of 1) and smaller for high UV index days (e.g. 0.4 for a day with a UV index of 12). Comparison of T_{90} for the different water types suggested that DOC had the largest influence on oocyst inactivation by solar UV, with waters with the T_{90} increasing as the DOC increased, and waters with similar DOC having similar T_{90} , despite differing by up to tenfold in color and turbidity (King et al. 2008). The penetration of UV in water has been shown to be highly dependent on the concentration and type of DOC (Morris et al. 1995; Jerome and Bukata 1998; Hutchison et al. 2005), providing protection from solar radiation for oocysts that are at sufficient depth to allow UV attenuation. However, particular events such as warm-water inflows into cold reservoirs will place oocysts within a zone where they can be inactivated by sunlight, which can cause >90 % inactivation after a few hours on a high UV index day.

The relative contribution of the different wavelengths of sunlight has been investigated using long-pass filters (Connelly et al. 2007; King et al. 2008). Connelly et al. (2007) used a <404 nm long pass filter to remove UV-B and UV-A. Compared with dark controls, oocysts exposed to sunlight with the UV component removed had between 42 % and 61 % reduction in infectivity, compared with reductions of 93 and 97 % for oocysts exposed to solar UV. King et al. (2008) used cut off filters to remove UV-B (<323 nm) and UV-A and UV-B (<400 nm). In these experiments, full sunlight achieved 2.5–2.7 log₁₀ oocyst inactivation, whereas the removal of UV-B reduced the amount of inactivation to $0.8-1.3 \log_{10}$ and the removal of both UV-A and UV-B abolished any reduction in infectivity compared with the dark control, which is in contrast with the finding of Connelly et al. (2007). While UV-B might be more effective in causing DNA damage and reducing infectivity (causing approximately two thirds of the observed inactivation), UV-A light has also been previously shown to cause cytotoxic and mutagenic effects, although to a smaller extent than UV-B (Ravanat et al. 2001). While UV-A causes less damage and a longer exposure time is required, it is likely to be of greater ecological significance than UV-B in the context of oocyst inactivation within a water column because longer wavelengths are better able to penetrate deeper into water.

The mechanism of solar UV inactivation has been further elucidated through the study of the effect of UV radiation on sporozoite DNA and cellular processes (King et al. 2010). One of the main mechanisms of inactivation caused by short wavelength UV irradiation is the production of DNA photoproducts, specifically cyclobutane pyrimidine dimers (CPDs) (Ravanat et al. 2001). Oocysts were exposed to varying doses of UV-C and an anti-CPD antibody was used to readily detect CPDs in oocyst DNA extracts (King et al. 2010). However, CPDs could not be detected in oocysts exposed to levels of solar UV causing 3 log₁₀ inactivation, whereas CPDs could be detected in ocysts exposed to a UV-C dose causing the same level of inactivation (approx. 10 mJ/cm²). It was possible to detect CPDs in the DNA from solar UV irradiated oocysts, but only by increasing the amount of

DNA assayed tenfold and also by increasing the exposure times for the luminescent assay used to detect antibody binding events (King et al. 2010). Other cellular targets were assessed to determine if they were affected by solar UV causing loss of infectivity. Solar radiation was found to induce membrane depolarization in sporozoites, leading to an increase in intracellular calcium and apical organelle discharge, which was apparent as a loss of internal granularity as measured by flow cytometry (King et al. 2010). The ultimate consequence of exposure to solar radiation was the accelerated depletion of ATP (compared with dark controls) and a decrease in the ability of sporozoites to attach to and invade host cells (King et al. 2010).

13.3.5 Inactivation by Other Environmental Stressors

While ammonia levels may become high enough in animal feces/manure storages to reduce oocyst viability, the impact of free ammonia on oocyst viability is likely to be negligible in drinking water reservoirs (Brookes et al. 2004). Brookes et al. (2004) estimated that levels of ammonia in lakes would typically be 1,000fold lower than the lowest concentration tested by Jenkins et al. (1998) and that ammonia levels would be unlikely to significantly affect oocyst viability even in lakes undergoing eutrophication. However, anthropogenic activity could introduce other stressors into the aquatic environment. For example, algicides are often used to control blooms of toxic or taste and odor-producing cyanobacteria, which commonly occur in water storage reservoirs (due to eutrophication, often as a consequence of nutrient inputs from agriculture and urban development) or waste stabilization ponds and cause problems with the production of potable or reuse water. These blooms were historically treated using compounds such as copper sulfate, but due to environmental concerns, alternatives have been developed, such as stabilized hydrogen peroxide. Hydrogen peroxide has been shown to be effective at inactivating oocysts (Barbee et al. 1999; Weir et al. 2002; Quilez et al. 2005; Castro-Hermida et al. 2006), but the available studies have focused on high doses of peroxide for rapid inactivation of *Cryptosporidium*. It is not known if oocysts are also inactivated by stabilized hydrogen peroxide or if the amounts dosed to control cyanobacteria will provide a high enough concentration to be effective against Cryptosporidium oocysts.

13.4 Removal by Treatment Processes

Water treatment provides a key barrier to the entry of *Cryptosporidium* into drinking water supplies. For the purpose of this chapter, we will consider water treatment processes to be those engineered processes responsible for the physical removal of oocysts from water and will consider disinfection processes separately

(although they are part of the treatment plants used to produce drinking water or reuse water). As discussed elsewhere in this book (see Chaps. 3 and 12), many of the largest waterborne outbreaks of cryptosporidiosis around the world have been caused by failures in water treatment systems (e.g., Hayes et al. 1989; Mac Kenzie et al. 1994; Yamamoto et al. 2000), so water treatment is a critical barrier for the protection of public health (Risebro et al. 2007).

13.4.1 Flocculation

Cryptosporidium poses a challenge to conventional water treatment plants. Most older plants utilize treatment processes designed to remove turbidity and to a certain degree natural organic matter (NOM), but have not been designed to target the removal of particles in the Cryptosporidium oocyst size range. The first step in conventional filtration plants is the addition of coagulants, such as alum or ferric chloride, as well as coagulant aids (cationic polyelectrolytes), to remove particulates by flocculation followed by sedimentation. Coagulation results in the aggregation of suspended particles in source water, forming a gel like precipitate (aluminium hydroxide in the case of alum). The removal of Cryptosporidium through flocculation can be quite effective and is the basis of a cost effective method for the concentration of oocysts in water samples for routine monitoring (Vesev et al. 1993). Poor coagulation conditions have been shown to result in poor filtration performance (Emelko 2003). A pilot plant study using alum as a coagulant removed approximately 1 log₁₀ oocysts after coagulation and sedimentation (Hsu and Yeh 2003). Jar testing of river water, using alum as the coagulant and an oocyst surrogate (with similar characteristics to an oocyst $-4-6 \mu m$ in size, 1.05-1.10 specific gravity, -25 mV (pH 6.0-6.5) Zeta potential), measured 1.2-1.5 log₁₀ surrogate removal by coagulation/sedimentation (Lee et al. 2007). The mechanism of oocyst removal during flocculation is dependent on the oocyst charge (Bustamante et al. 2001). A study of oocyst zeta potential in the presence of different coagulants suggests that oocysts maintain a negative charge in the presence of ferric chloride and so sweep flocculation (physical entrainment in the developing floc) is required for oocyst removal, whereas oocysts undergo charge neutralization in the presence of alum, meaning that oocysts can adsorb to alum flocs, as well as be removed by sweep flocculation (Bustamante et al. 2001). These findings have been further supported by studies of oocyst charge characteristics in the presence of different salt concentrations and coagulants (Butkus et al. 2003). The chemicals used for flocculation do not appear to reduce oocyst infectivity. Examination of filter backwash water from water treatment plants detected infectious oocysts (Di Giovanni et al. 1999) and jar testing using alum as the coagulant did not measure any reduction in oocyst infectivity. A small amount of infectivity reduction of 0.4 \log_{10} also occurred in the control jars without alum, indicating an effect due to mechanical processing (Keegan et al. 2008).

13.4.2 Media Filtration

Sand filtration was first reported to have potential for removal of oocysts based on a laboratory-scale experiment using a 1 m column of pool filter sand challenged with oocvsts purified from a human clinical sample (Chapman and Rush 1990). In this experiment, 10⁶ oocysts were used to challenge the filter and none were detected in the filtrate, while oocysts were detected only in the top 300 mm of the filter bed (Chapman and Rush 1990). The transport of oocysts through a filter medium has been measured for wastewater treated with sand filters. Grain size was a key factor, with fine-grained sand effectively removing oocysts while coarse-grained media had more oocyst break-through, depending on hydraulic loading rates (Logan et al. 2001). Filter removal efficiency has been shown to be affected by the presence of biofilm and NOM. The removal efficiency was measured for oocysts coagulated using calcium chloride and filtered through silicon beads, finding that biofilm reduced the filtration efficiency by 50 % and the presence of 5 ppm of NOM reduced efficiency by 75 % (Dai and Hozalski 2002). The reduction in filtration efficiency was likely due to charge interactions, with improved filtration performance observed using alum as the coagulant, even in the presence of NOM (Dai and Hozalski 2002). Filtration efficiency has also been shown to be affected by ionic strength and pH (Hsu et al. 2001). However, a more recent study suggests that oocyst removal by a packed quartz sand column was not overly affected by ionic strength and that physical straining is an important capture mechanism (Tufenkji et al. 2004). Atomic force microscopy has characterized the interaction between oocysts and sand particles, suggesting that proteins on the oocyst surface contribute to electrostatic repulsion against the surface of sand particles, and that dissolved organic carbon and dissolved calcium assist with overcoming this repulsion (Considine et al. 2002). A further study of oocyst wall macromolecules suggests that they are responsible for electrostatic repulsion from surfaces, even in the presence of ionic strengths predicted to overcome any electrostatic repulsion, and that treatment of the oocysts with proteinase K to remove the macromolecules resulted in improved oocyst attachment to surfaces (Kuznar and Elimelech 2006).

A monitoring program of source water and drinking water produced by a conventional treatment plant (coagulation-flocculation, sedimentation and rapid filtration) measured a combined oocyst removal of 2.5–2.7 \log_{10} , depending on the method used to determine oocyst numbers (geometric mean versus 90 % observation level) (Hashimoto et al. 2001). Pilot-scale experiments using similar treatment processes measured a combined oocyst removal of 5–6 \log_{10} (Gale et al. 2002). There appear to be fewer studies examining filtration performance separately from coagulation/sedimentation. Sand filter column testing, using a *Cryptosporidium* tracer (described in the *Flocculation* section) and water that had been coagulated with alum measured a 1.3–1.5 \log_{10} removal of the tracer (Lee et al. 2007). Poorer performance (0.5 removal) was reported for a pilot plant sand filter receiving secondary treated effluent from an activated sludge plant, although performance was greatly increased by dosage of alum (2 \log_{10} removal) (Suwa and

Suzuki 2003). Other studies have reported variable filter performance in wastewater treatment plants, ranging from 0.4 to 1.5 \log_{10} depending on the filter type (Levine et al. 2008; Fu et al. 2010). The differences in performance are likely due to differences in operational parameters such as hydraulic load, chemical pretreatments and backwashing regimes (Levine et al. 2008). In contrast, laboratory-scale slow sand filters have achieved $>5 \log_{10}$ oocyst removal, a similar performance as observed for pool sand filters by Chapman et al. (1990) (Hijnen et al. 2004). This removal rate was confirmed in a later study by loading a mature pilot sand filter with oocysts, dosing for 100 days and monitoring for 150 days, demonstrating oocyst removal of 4.7 \log_{10} (Hijnen et al. 2007). Oocysts did not accumulate in the filter and it was hypothesized that they were removed as a result of predation by zooplankton (Hijnen et al. 2007), Granular activated carbon (GAC), often used in filters to reduce NOM or remove organic contaminants, has also been shown to be effective at removing oocvsts $(1.3-2.7 \log_{10})$ (Hijnen et al. 2010). A study on the fate of oocysts on GAC also observed oocyst loss from the filter medium due to biotic activity (Bichai et al. 2010). Zooplankton were isolated, identifying rotifers as a dominant group (previously shown to be predators of Cryptosporidium) and oocysts were recovered from zooplankton following release by sonication (Bichai et al. 2010).

Conventional filters using sand, gravel, anthracite, etc., are not the only types of filters in use for the production of drinking water or the treatment of wastewater. Bank filtration, which passes water through a river bank, is in use in some parts of the world as a pretreatment for drinking water or for wastewater. The removal efficiency of sandy alluvial riverbank sediment in bank filtration, based on modeled distribution profiles, has been claimed to be $23-200 \log_{10}/m$ (Faulkner et al. 2010). In the case of bank filtration, straining is not the only removal mechanism and surface charge of the medium has been suggested to play a major role, such that gravel soil can have better performance than sandy soils for removal of microorganisms (Hijnen et al. 2005; Faulkner et al. 2010). The performance of bank filtration for oocyst removal has been found to correlate with the surface coverage of the sediment grains by metal oxides, with higher coverage providing better removal (Metge et al. 2010). A novel filtration medium was developed using Zeolite treated with a quaternary ammonium chloride. This modified Zeolite exhibited an oocyst removal efficiency of 0.54, which is comparable with sand filters but of most interest it also resulted in 1.19 log₁₀ inactivation of the oocysts (Abbaszadegan et al. 2006). Surface functionalization of the sand in sand filters may improve filter performance. Functionalizing silica with 3-(2-aminoethyl) aminopropyltrimethoxysilane improved the filtration performance from $1 \log_{10}$ for the uncoated silica to $1.3-1.4 \log_{10}$ for the functionalize silica, depending on the level of coating (Majewski and Keegan 2012).

13.4.3 Dissolved Air Flotation

Dissolved Air Flotation (DAF) treatment of surface waters and wastewaters uses coagulants to develop flocs, which are floated to the surface using aeration and then collected from the top of the reactor and disposed of as sludge (for a detailed review of this process see Edzwald 2010). The effectiveness of the DAF process, like conventional flocculation processes, is affected by turbidity and natural organic matter and has reached oocyst removals of $1.7-2.5 \log_{10}$, increasing to $> 5.4 \log_{10}$ when combined with filtration (DAFF) for drinking water (Edzwald et al. 2001). In the case of DAFF, the operation of the plant in terms of filter backwashing is important for determining if oocysts are concentrated in the floated sludge or in the filters, with the volume of backwash water used determining the level of oocyst build-up in the plant influent (Edzwald et al. 2001).

13.4.4 Membrane Filtration

Membrane filtration systems can provide excellent removal of oocysts, through size exclusion. Most research on membrane filters in terms of Cryptosporidium has been in relation to the concentration of oocysts from large volume water samples (e.g. Hill et al. 2007; Lindquist et al. 2007; Liu et al. 2012), rather than validating the technology for oocyst removal. Full-scale validation of membrane filters will often measure removal of bacteria (such as E. coli) and use calculated log₁₀ removal values (LRV) for smaller organisms to provide a conservative LRV for the much larger oocysts. In practice, the performance of filters is higher than the maximum LRV that will be credited by health regulators. As a guiding principle, health regulators (at least in Australia) will not provide a credit of more than 4 LRV for any individual treatment barrier. In the case of membranes this is partly driven by limitations in real-time detection of membrane failure, for example parameters such as turbidity may not detect failures that will allow breakthrough of small particles (such as viruses) while still removing larger particles and producing acceptable levels of turbidity in the filtrate. Ultrafiltration has been shown to be effective $(>1.84 \log_{10})$ at removal of oocysts in a full scale testing for tertiary treatment of secondary effluent (Fu et al. 2010). A pilot-scale evaluation of conventional and membrane filters has been reported, but the numbers of oocysts in the raw water were low (56 oocysts/10L) and so limited conclusions can be drawn from that work regarding oocyst removal efficiency (Hsu and Yeh 2003).

13.4.5 Activated Sludge and Other Secondary Wastewater Treatment Processes

As water becomes scarcer, increasing attention is being given to the reuse of wastewater. Most of the common conventional wastewater treatment processes, such as activated sludge process (ASP) or waste stabilization ponds, are designed to reduce nutrients to a level suitable for environmental discharge, with pathogen removal more often being achieved by disinfection prior to discharge, although some pathogen attenuation may occur during treatment. Wastewater reuse usually requires tertiary treatment to further improve the quality of the water and also to remove pathogens and protect human health. The activated sludge process is essentially a large bioreactor designed to achieve nitrification and denitrification of the wastewater to remove nitrogen from the wastewater. Being an active biological process, there is some opportunity for removal or inactivation of oocysts, either through entrapment in biological flocs or predation. An early study of lab-scale ASP using C. parvum oocysts demonstrated a 1 log_{10} reduction in the intensity of infection measured using mice (Villacorta-Martinez de Maturana et al. 1992). A pilot plant ASP showed an oocyst removal efficiency of 2, which was increased to $3 \log_{10}$ by dosing with alum (Suwa and Suzuki 2003). Monitoring of a full-scale wastewater treatment plant measured a $1.52 \log_{10}$ oocyst removal for conventional ASP and higher removals for an oxidation ditch process $(2.17 \log_{10})$ and an anaerobic-anoxic process $(1.79 \log_{10})$ (Fu et al. 2010). Another study of a full-scale ASP reported better removal of oocysts, with an LRV of almost 3 log₁₀ (Neto et al. 2006). There are many different configurations for ASP, and these may have a large effect on removal efficiency. A lab-scale ASP, using return activated sludge from a full-scale ASP, was able to achieve a 2.4 log₁₀ reduction in oocyst number, whereas the full-scale plant only removed 0.7 \log_{10} (Wen et al. 2009).

13.4.6 Lagoons/Waste Stabilization Ponds

Wastewater stabilization ponds and constructed wetlands provide attractive low technology and low energy solutions for treating contaminated waters such as wastewaters or storm waters, especially in areas where there is sufficient space. Provided that detention times are adequate and there are reduced opportunities for disturbance of sediments, lagoons and constructed wetlands have the potential to achieve excellent removal of oocysts from wastewater or storm water. Sedimentation is a major mechanism for oocyst reduction. Analysis of oocysts in a surface flow wetland showed that oocyst numbers in sediment were 1–3 orders of magnitude higher compared with oocyst numbers in the water column (Karim et al. 2004). Monitoring of lagoon systems used for treating wastewater has shown 1–2.5 log₁₀ oocyst removal (Ulrich et al. 2005), with constructed wetlands capable of 1–2 log₁₀ LRV (Quinonez-Diaz et al. 2001; Reinoso et al. 2008). As previously mentioned,

some of this removal is thought to be a result of protozoan predation (Stott et al. 2001). Some variability in the performance of constructed wetlands has been noted, with a system receiving sewage effluent only removing 64 % of oocysts (Thurston et al. 2001).

High rate algal ponds (HRAP) are a variation of waste stabilization ponds, actively using energy to provide a higher level of oxidation to support algal growth, which mediates nutrient removal. The HRAP design has been shown to be effective at oocyst inactivation, causing 2–3 log_{10} reduction in mouse infectivity (Araki et al. 2001). Further study of oocyst inactivation in a HRAP using semipermeable bags suggested that the inactivation was due to a combination of ammonia, pH and sunlight exposure (Araki et al. 2001). Sunlight has also been suggested to be a major cause of inactivation of oocysts in waste stabilization ponds, with 40 % inactivation observed after 4 days exposure to sunlight (Reinoso and Becares 2008). Considering that secondary treated effluent is high in organics, it is tempting to speculate that UV-A is a primary factor causing inactivation in this scenario. A comparison of anaerobic ponds, a facultative pond and a maturation pond suggested that sunlight and water chemistry were the main factors influencing oocyst removal in the anaerobic and maturation ponds, whereas predation may have been more important in the facultative pond (Reinoso et al. 2011).

13.5 Disinfection of *Cryptosporidium* in Water

This chapter has so far reviewed natural and engineered processes that remove or inactivate *Cryptosporidium* oocysts in environmental waters or in wastewaters. The final barrier, either for the production of safe drinking water or for the production of fit for purpose reuse water, is disinfection. Many processes have been evaluated, with some now in widespread use for the management of any *Cryptosporidium* oocyst that may break through water treatment processes. Disinfection methods, which will be discussed below, include chlorine, chloramine, chlorine dioxide, ozone, mixed oxidants and ultraviolet light (UV).

13.5.1 Oxidant-Based Disinfection

The resistance of *Cryptosporidium* to disinfectants such as chlorine is one of the primary reasons *Cryptosporidium* has been an issue for the water industry and the cause of so many drinking water-related outbreaks. The ineffectiveness of chlorine and chloramine was recognized in the early 1990s, with disinfection studies showing that a contact time (Ct) of 1,600 mg.min/L using chlorine was required to achieve a 1 log₁₀ reduction in oocyst infectivity, measured using mice (Korich et al. 1990). Contact time, which is the product of the disinfectant dose (mg/L) multiplied by time of exposure (minutes), provides a measure for comparing the

effectiveness of different chemical disinfectants and also provides a way of ensuring delivery of an adequate dose during water treatment. In the case of the study of Korich et al. (1990), the Ct of 1,600 mg.min/L was delivered using 80 ppm (ppm = mg/L) of free chlorine with an exposure time of 20 min. Chloramine was shown to be equally ineffective in the same study. While it is not possible to deliver the required Ct for treating drinking water or reuse water, chlorine can be used to effectively inactivate oocysts in swimming pools following a fecal accident. However, there is usually insufficient contact time immediately following a fecal accident, so any pool users in the vicinity will likely become infected if they have ingested contaminated water before the pool has been treated with elevated levels of chlorine. A more recent study using two different isolates of C. parvum and cell culture infectivity demonstrated that the Ct required for 3 log₁₀ inactivation was 10,400–15,300 mg.min/L, which is larger than the previous recommendation from the Centre for Disease Control and Prevention for the treatment of recreational waters following a fecal accident (Shields et al. 2008). Chlorine dioxide was shown to be more effective, while ozone treatment (1 ppm) was the best option for inactivation of oocysts, achieving 2 log₁₀ inactivation for a Ct estimated between 5 and 10 mg.min/L and >4 \log_{10} inactivation for a Ct of 10 mg.min/L (Korich et al. 1990). Similar results were observed in an independent study using 1.11 ppm ozone, achieving approximately 4 log₁₀ inactivation for a Ct of 6.6 mg.min/L (Peeters et al. 1989).

Electrochemically produced oxidants mixed (chlorine plus other uncharacterized species produced by a MIOX unit) have been shown to inactivate $>3 \log_{10}$ oocysts with a Ct of 1,200 mg.min/L (5 ppm mixed oxidant with 4 h contact), whereas no inactivation was observed after exposure to chlorine for Cts of 1,200–7,200 mg.min/L (5 ppm for 4–24 h contact) (Venczel et al. 1997), although it was not clear from the study if chlorine decay was measured to properly calculate a Ct. The lack of any observed inactivation for chlorine for the nominated Ct range is suspect considering the earlier findings of Korich et al. (1990). A study of the inactivation kinetics of chlorine dioxide for oocysts compared excystation and cell culture infectivity, showing that for a Ct of 1,000 mg.min/L, the excystation method only measure $0.5 \log_{10}$ inactivation, whereas the infectivity method measured a $2 \log_{10}$ inactivation (Chauret et al. 2001). This result suggests that the earlier finding of Venczel et al. (1997) may have in part been affected by the method used for estimating viability. The performance of MIOX units appears to be variable. The effectiveness of MIOX could not be reproduced in a subsequent study using a cell culture infectivity assay, where the MIOX mixed oxidants and chlorine treatment performed similarly (Keegan et al. 2003). However, a different study using mouse infectivity measured inactivation for oocysts exposed to 2-5 mg/L MIOX (measured as free chlorine) but not for 5 mg/L free chlorine (Sasahara et al. 2003). An on-site evaluation of an electrolytic system (ECO, using the same system as MIOX) found that the system was not effective at inactivating oocysts, producing equivalent results to chlorine (Venczel et al. 2004).

Conflicting results have been reported for the sensitivity of oocysts as a function of age. A study using oocysts aged in dialysis cassettes suspended in a river reported

no change in chlorine sensitivity, although this study used excystation to measure chlorine inactivation (Chauret et al. 1998). In contrast, oocysts stored at 4 °C for 24 weeks were shown to have approximately 1 \log_{10} more inactivation for Ct 300 mg.min/L compared with fresh oocysts using an infectivity assay (Keegan et al. 2008).

13.5.2 Ultraviolet Light

Discussion within the scientific literature regarding the use of UV-C for the disinfection of potable water first began in the late 1980s/early 1990s in response to proposed changes to the US Safe Drinking Water Act (Wolfe 1990). At that time, there was no information on UV inactivation of Cryptosporidium, but oocysts were presumed to be more resistant than bacteria on the basis of the observed resistance of oocysts to chlorine (Wolfe 1990). There are two types of UV-C lamps used in disinfection systems, medium pressure (MP)-UV lamps that deliver UV light in the full UV-C spectrum (200–300 nm) and low pressure (LP)-UV lamps that deliver germicidal UV-C (254 nm). The first report of the effectiveness of UV against Cryptosporidium used mouse infectivity to demonstrate complete inactivation of oocysts following a UV dose of 15,000 mW/s for 150 min, although the dose delivery was not measured to allow comparison with more recent studies (Lorenzo-Lorenzo et al. 1993). A subsequent study used viability assays (vital dyes and excystation) to demonstrate 2-3 log 10 reduction in viability using a total low pressure LP-UV dose of 8,748 mJ/cm² (Campbell et al. 1995) and on the basis of this and similar studies using viability techniques it was believed for many years that Cryptosporidium was relatively UV resistant.

A detailed study of the dose response of Cryptosporidium to LP- and MP-UV, using mouse infectivity and a collimated beam apparatus for precise UV dose delivery, demonstrated inactivation of 2 \log_{10} for 10 mJ/cm², 3 \log_{10} for 25 mJ/cm² and 3.4 – >4.9 \log_{10} for 119 mJ/cm² (Craik et al. 2001). Evaluation of a UV disinfection device that delivered 120 mJ/cm² was shown to reduce infectivity in mice by 5.4 \log_{10} (Drescher et al. 2001). The most effective wavelengths of UV-C were determined to be 250-275 nm (Linden et al. 2001). A direct comparison of animal infectivity and excystation methods demonstrated that high doses of UV were required to reduce excystation, but much lower doses were required to reduce infectivity (Morita et al. 2002). Dose-response studies at low UV doses measured >2.6 \log_{10} for 3 mJ/cm² and >3.9 \log_{10} for 10 mJ/cm² (Shin et al. 2001), with a similar bench-scale study using cell culture infectivity demonstrating $>3 \log_{10}$ inactivation for 5–10 mJ/cm² doses of LP-UV (Bukhari and LeChevallier 2003). Due to ease of production in calves or mice, the vast majority of UV studies have used C. parvum. Using oocysts isolated from a human patient, C. hominis has been shown to possess a similar UV sensitivity compared with C. parvum (Johnson et al. 2005). The requirement for lower UV doses has translated into suitable UV reactor design, with full-scale UV systems described for *Cryptosporidium* disinfection and validated using MS2 bacteriophage as a challenge organism (Bukhari and LeChevallier 2003).

A study of ex vivo and in vivo reactivation of oocysts found no evidence for reactivation, with oocysts exposed to 60 mJ/cm² of MP-UV remaining inactivated. as measured using mouse infectivity (Belosevic et al. 2001). Other studies also examined photoreactivation and dark repair, finding that although Cryptosporidium possesses UV repair genes and that cyclobutane pyrimidine dimers were shown to be repaired, infectivity was not restored (Oguma et al. 2001; Morita et al. 2002; Rochelle et al. 2004). Based on more recent knowledge regarding the importance of energy stores for infectivity and the impact of sunlight on infectivity (King et al. 2010), it is likely that sporozoite repair of UV-induced damage results in energy depletion and loss of infectivity. Transcriptome analysis supports this, showing an upregulation of genes for DNA repair and intracellular trafficking (Zhang et al. 2012), which is consistent with the finding of King et al. (2010) in terms of UV-B causing premature exocytosis. Some studies have recommended the use of MP-UV to prevent reactivation (Kalisvaart 2004), but the available evidence suggests that LP-UV is sufficient to cause a permanent reduction in infectivity for Cryptosporidium.

Conventional UV reactors used for treatment of bulk water use UV lamps enclosed in a quartz sleeve and installed either in the center of a stainless steel pipe, or suspended in an open channel. Water is passed through the pipe or channel and the UV dose delivered is a function of the lamp intensity, flow rate, mixing through the system and the UV transmissivity of the water. In conventional UV systems the lamps are always on while the system is operational. An alternative system that has been evaluated more recently uses pulsed UV light, which has been shown to achieve $6 \log_{10}$ inactivation with an equivalent UV dose of 278 mJ/cm² and $2 \log_{10}$ with a dose of 15 mJ/cm² (Lee et al. 2008). One issue with the use of pulsed UV technology is the difficulty in measuring the UV fluence (dose) delivered to oocysts (Garvey et al. 2010).

13.5.3 Combined Processes

Sequential disinfection has been shown to improve oocyst inactivation levels. The combination of ozone followed by free chlorine increased oocyst inactivation by four to sixfold compared with ozone alone (Li et al. 2001). Similar results were obtained using a combination of ozone and monochloramine, although the increase in inactivation was only 2.5-fold (Rennecker et al. 2001). The synergistic effective-ness of ozone followed by chlorine has been demonstrated using both animal infectivity (Li et al. 2001; Biswas et al. 2005) and excystation assays (Corona-Vasquez et al. 2002). Low ozone doses have been combined with UV disinfection to allow adequate disinfection of water while reducing the formation of disinfection by-products by ozone (Meunier et al. 2006). An added advantage of this combination is that the ozone will react with organics in the water, helping to increase the

UV transmissivity and so improve UV dose delivery. In terms of the treatment of wastewater to produce reuse water, combined treatment trains such as UV and chlorine disinfection have been recommended to allow control of both chlorine resistant pathogens such as *Cryptosporidium* and UV resistant pathogens such as adenovirus (Montemayor et al. 2008).

13.5.4 Heat

As has been previously discussed in this chapter, *Cryptosporidium* are sensitive to heat, making this an effective method for disinfection. Moist heat is highly effective (Anderson 1985) and the temperatures achieved in thermophilic aerobic digestion of sewage sludge (55 °C) are sufficient for ready inactivation of oocysts (Whitmore and Robertson 1995). Pasteurization has been shown to be effective for the inactivation of oocysts in water and milk (Harp et al. 1996), but while this treatment can be used to inactivate oocysts in food (e.g. Deng and Cliver 2001), it has not been used for large scale treatment of water or wastewater.

13.5.5 Other

Photocatalytic inactivation of Cryptosporidium oocysts has been assessed using titanium dioxide films and a black light lamp (essentially UV-A), achieving up to 78 % oocyst inactivation after 3 h exposure (Sunnotel et al. 2010). However, this study used viability surrogates rather than infectivity and so may have underestimated the effectiveness of the treatment. The measured inactivation was not due to UV-A exposure based on comparison of a control exposed to the light source in the absence of the titanium dioxide. A fibrous ceramic titanium oxide photocatalyst, used in combination with UV-C, has also been shown to be effective at reducing oocyst numbers, suggesting that the photo byproducts produced are active at disrupting oocysts (Navalon et al. 2009). The mechanism of photocatalytic titanium oxide or ferrioxolate inactivation is the generation of the hydroxide radical, which has been reported to be $10^4 - 10^7$ times more effective compared to ozone, chlorine dioxide or chlorine (Cho and Yoon 2008). However, another study examining LP-UV and titanium dioxide suggested that reactive oxygen species were only responsible for a relatively small amount of inactivation when used in isolation, but that when used in combination with UV produced a synergistic effect resulting in more inactivation than either exposure alone (Ryu et al. 2008).

Sonication has been shown to be effective in physically disrupting oocysts, causing oocysts to release contents, and also to reduce the infectivity of any remaining intact oocysts (Oyane et al. 2005). Using a horn-type sonicator, 60 s at 52 W resulted in 52 % inactivation, while 126 W for the same exposure time caused

94.9 % inactivation (Oyane et al. 2005). A flow through sonicator was also able to inactivate oocysts, but it was only effective at low flow rates (Oyane et al. 2005).

13.6 Conclusions

Cryptosporidium continues to be a public health concern around the world, through outbreaks caused by contaminated drinking water or recreational water. However, we have developed an increasingly better understanding of the fate and transport of oocysts through the environment and of the processes required to remove or inactivate them in drinking water or wastewater. There are many processes, starting from the deposition of oocysts into the terrestrial environment, through to overland transport into surface waters and hydraulic transport through to water treatment plants, that either cause the removal or inactivation of a significant number of oocysts. There are also processes that can be used to treat surface waters to remove or inactivate oocysts for the production of potable water. Despite these advances, outbreaks from drinking water will still occur, either due to treatment barrier failures, or because of insufficient risk characterization of catchments, resulting in inadequate treatment barriers to prevent entry of infectious oocysts into drinking water systems. Although there are advanced treatment options, outbreaks will still occur from recreational waters and swimming pools because it is not possible to prevent transmission when fecal events occur in close proximity to other water users. However, at least for drinking water, the technology is available to remove the threat of a Cryptosporidium outbreak, although at the cost of additional infrastructure and operating costs.

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