Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Looking for Cryptosporidium: the application of advances in detection and diagnosis

Rachel M. Chalmers¹ and Frank Katzer²

¹Cryptosporidium Reference Unit, Public Health Wales Microbiology, Singleton Hospital, Swansea, UK
²Moredun Research Institute, Pentlands Science Park, Penicuik, Edinburgh, UK

The protozoan Cryptosporidium is a major public and animal health concern. Young children, immunocompromised people, and pre-weaning animals are especially vulnerable, but treatment options are limited and there is no vaccine. A laboratory diagnosis is required to confirm cases of cryptosporidiosis, and species and genotype determination is essential in distinguishing human from non-human sources, understanding transmission, and strengthening the epidemiological evidence for causative links in outbreaks. However, testing is not consistent, as demonstrated by investigation of a significant increase in cases in some European countries during 2012. Many methods employed are laborious and time-consuming; recent advances, translated into diagnostic assays, can improve testing and facilitate typing to support clinical and environmental investigations.

Why look for Cryptosporidium?
Cryptosporidium causes sporadic cases and outbreaks of gastroenteritis (cryptosporidiosis) through faeces-contaminated drinking and recreational water [1], food and beverages [2], person-to-person spread in households and institutions such as daycare settings [3], farmed animal contact [4], and in livestock, companion, and other animals [5] (Figure 1). Increased morbidity, mortality, and socioeconomic implications [6,7] are recognised by inclusion of Cryptosporidium in the Neglected Diseases Initiative of the World Health Organization (WHO) [8]. Cryptosporidiosis is a laboratory diagnosis because it is not pathogenic – in other words, the acute signs and symptoms can be similar to other infectious and non-infectious causes of gastroenteritis [9,10]. In cases where other parasites are also suspected, or in populations where they are prevalent, multiple faecal samples with and without chemical preservation may be required [11] (Figure 2). Differential diagnosis enables provision of appropriate advice on treatment, control, and management, and at the population level it is required for the identification of outbreaks and for monitoring trends, risk factors, and interventions.

The ability of Cryptosporidium to break through multiple water-treatment barriers and cause large-scale outbreaks has had huge impact on the water industry and its regulation [12], and it has been used as a reference pathogen for the faecal-oraly transmitted protozoa in the design and implementation of the WHO Guidelines for Drinking Water Quality [13,14]. Monitoring for oocysts is part of the surveillance to support water-safety plans [14].

During 2012 a significant but unexplained increase in the incidence of human cryptosporidiosis was reported by Fournet and colleagues in the UK, The Netherlands (NL), and Germany to the European Centre for Disease Control [15]. This report highlights difficulties in comparing data across countries, derived differently due to variations in laboratory testing and reporting. The lack of widespread, routine Cryptosporidium diagnosis and species identification, or a standardised subtyping scheme, impinges on investigation. These shortcomings also occur in water testing where only the genus is identified. Phylogenetic comparisons of isolates from cases and sources are not undertaken in most settings, impacting on source attribution and risk assessments for drinking water and other exposures [13].

Techniques for the detection, diagnosis, and characterisation of Cryptosporidium spp. from different sample types have been reviewed previously [16–18]. However, not all have been delivered from research to diagnostic settings. Here we examine the current workflows and review advances that have been translated into clinical and environmental investigations.

Cryptosporidium, an uncertain taxonomy
Taxonomic positions are important in understanding the evolution, biology, pathogenesis, sources, and transmission of pathogens as well as focussing efforts for identifying targets for diagnosis and drug therapies. The taxonomy of Cryptosporidium (phylum Apicomplexa, class Coccidea, order Eimeriidae) was disputed, because of life-cycle (Figure 1), phylogenetic, and biological differences observed in comparison with other Eimeriidae (e.g., Cyclospora, Cystoisospora, and Eimeria) [19]. Traditionally, Apicomplexan species were defined by host-specificity, location of endogenous stages, and morphology. For Cryptosporidium spp., however, host-specificity is variable, and oocyst morphology distinguishes only those with smaller,
near-spherical oocysts (4–6 μm in diameter) from those with larger, more oval oocysts (7 × 5 μm) (Table 1). Therefore, a combination of biological and genetic data is recommended for identifying and naming new species: (i) morphological features of the oocyst and demonstration of developmental stages; (ii) natural and experimental host range; (iii) detailed biological characterisation, including localisation and pathology associated with an individual host; (iv) differential diagnosis; (v) genetic characterisation at multiple loci, to include the small subunit ribosomal RNA (SSU rRNA) and other functional genes; (vi) deposition of definitive reference sequences and preserved material for further analyses; and (vii) compliance with International Committee for Zoonotic Nomenclature (ICZN) species-naming rules [20]. Where mainly genetic data are available, isolates are described as ‘genotypes’ [21]. The ICZN rules are open to interpretation, especially in the Principle of Priority, leading to conflicting opinions about renaming Cryptosporidium spp. (e.g., Cryptosporidium parvum as Cryptosporidium pestis) [22]. To avoid confusion and promote stability, here we use C. parvum for the species infective to calves and humans [21].

**Cryptosporidium species: infection and disease in relation to diagnosis**

At time of writing 26 Cryptosporidium species have been named (Table 1). There is good evidence for six as important causes of human cryptosporidiosis: most commonly Cryptosporidium hominis and C. parvum, followed by Cryptosporidium meleagridis and occasionally Cryptosporidium cuniculus, Cryptosporidium felis, and Cryptosporidium canis [18]. Infection is transmitted by oocysts and acquired mainly by the faecal–oral route, with very rare instances of respiratory infection following inhalation or aspiration [6]. Despite occasional reports in livestock, C. hominis appears to be transmitted anthroponotically.
Immunocompromised patients have provided sentinel alert to some new infections [23]. In animals, the clinically and economically important gastrointestinal species are *C. parvum* (pre-weaning ruminants), *C. meleagrisidis* (birds), *C. canis* (puppies), *C. felis* (kittens), *Cryptosporidium serpentis* (snakes), and *Cryptosporidium varani* (reptiles) [5]. *Cryptosporidium baileyi* is also associated with respiratory disease in birds [5]. The clinical significance of other species or genotypes in detected faeces is doubtful if disease is not described or tissue infection has not been demonstrated.

Human cryptosporidiosis (recently reviewed, [24]) is usually characterised by acute symptoms of watery diarrhoea accompanied by abdominal cramps, vomiting, low-grade fever, and loss of appetite. These indicate the pathogenesis with infection mainly of the small bowel (Figure 1), causing malabsorption and some elements of inflammation. The incubation period (normal range 3–12 days, usually 5–7 days) is dose-dependent. Cryptosporidiosis is self-limiting in immunocompetent people, and the symptoms vary in severity (mild to severe); duration can be for up to 1 month during which time relapse occurs in about one third of cases. Severity varies partially with the general health of the patient, and those experiencing mild, self-limiting diarrhoea will be unlikely to seek medical attention and obtain a diagnosis (Table 2).

In pre-weaning ruminants, *C. parvum* is usually associated with self-limiting gastroenteritis, characterised by profuse watery diarrhoea and signs of depression, dehydration, anorexia, and weakness, and in extreme cases it can be fatal [5]. The parasite is widely endemic and is one of the most frequently diagnosed causes of diarrhoea in pre-weaning calves [25]. Multiple pathogens can be involved in neonatal diarrhoea (rotavirus, coronavirus, pathogenic strains of *Escherichia coli* and *Salmonella* spp.), and cryptosporidiosis is confirmed by finding significant numbers of oocysts in diarrhoeic faeces in their absence [25]. However, coinfection may modulate the clinical presentation, and has been widely speculated to lead to more severe cryptosporidiosis [5], although experimental data to support this are still lacking. Infection with *C. felis* is often asymptomatic in cats but those also infected with the feline leukaemia virus are more likely to exhibit clinical signs [5]. A very different clinical spectrum may also be seen for example in birds where *C. baileyi* and *C. meleagrisidis* can also cause respiratory disease, requiring investigation of different sample types (Table 2).
### Table 1. Cryptosporidium species and their association with clinical disease

| Cryptosporidium species and genotypes* | Mean oocyst dimensions (μm)b | Major host(s) | Association with human cryptosporidiosis [18] | Association with animal cryptosporidiosis [5,21,23] | Additional key Refs |
|--------------------------------------|-------------------------------|---------------|-----------------------------------------------|--------------------------------------------------|-------------------|
| C. andersoni                         | 7.4 × 5.5                     | Cattle        | Rarely associated; individual reports from UK, Australia, and Malawi. | Usually seen in yearlings and adult cattle; not associated with diarrhoea but linked to reduced milk-yields and weight gain. | [63]              |
| C. baileyi                           | 6.2 × 4.6                     | Chickens, other birds | No association. | Many different bird species can be affected, mainly by infection in the upper respiratory tract which can be fatal; also linked to renal and ocular disease. Severe outbreaks and mortalities in poultry units. |                  |
| C. bovis (previously bovine B genotype) | 4.9 × 4.6                    | Cattle        | Rarely associated; individual reports from Australia and India of cases reporting cattle contact. | Shedding is seen mainly in weaned and sometimes pre-weaning calves but infection is considered non-pathogenic. | [63,64]           |
| C. canis (previously dog genotype)   | 5.0 × 4.7                     | Dog           | Epidemiologically linked to diarrhoea in children in a shanty town in Lima, Peru; occasional sporadic cases in various, especially non-industrialised, countries; some cases report contact with dogs. | Diarrhoea usually seen in puppies and younger dogs; asymptomatic shedding of oocysts has been reported in older and immune-competent dogs. |                  |
| C. cuniculus (previously rabbit genotype) | 5.6 × 5.4                    | Rabbit, humans | Caused a waterborne outbreak in UK; occasional, seasonal sporadic cases in UK, individual reports from France and children in Nigeria. | Asymptomatic shedding of oocysts in natural and experimental infections in rabbits. | [65–67]           |
| C. fayeri (previously marsupial genotype I) | 4.9 × 4.3                    | Marsupials    | Rarely associated; individual report from Australia of a person in contact with marsupials. | Oocyst shedding is not associated with diarrhoea and no clinical signs are described. | [63]              |
| C. felis                             | 4.6 × 4.0                     | Cat           | Epidemiologically linked to diarrhoea in children in a shanty town in Lima, Peru; occasional sporadic cases in various countries including developed countries, especially immunocompromised people and those with cat contact. | More common in kittens and younger cats and is then associated with diarrhoea. Often only low numbers of oocysts detectable. | [68]              |
| C. fragile                           | 6.2 × 5.5                     | Black spined toad | No association. | Infection appears to be host-specific because infections of four amphibian and one fish species failed. |                  |
| C. galli                             | 8.3 × 6.3                     | Chicken       | No association. | Infection of the proventriculus can lead to diarrhoea and fatalities; shedding can be prolonged and is not necessarily associated with diarrhoea. |                  |
| Cryptosporidium species and genotypes | Mean oocyst dimensions (μm) | Major host(s) | Association with human cryptosporidiosis [18] | Association with animal cryptosporidiosis [5,21,23] | Additional key Refs |
|-------------------------------------|-----------------------------|---------------|---------------------------------------------|-------------------------------------------------|-------------------|
| C. hominis (previously referred to as C. parvum human genotype, genotype I, and genotype H) | 4.9 × 5.2 | Humans | Common cause of diarrhoeal disease in sporadic cases and outbreaks. Infectivity data from experimental infections in adults extrapolated to dose–response indicate that ingestion of a single oocyst carries a discrete probability of infection. Transmitted 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. High parasite genetic heterogeneity in non-industrialised countries; more homogenous in industrialised countries. Children in Brazil shed more oocysts, had more frequent presence of faecal lactoferrin, and greater growth shortfalls than those infected with C. parvum, even in the absence of symptoms. HIV-positive persons in Lima, Peru, experienced more severe symptoms than those infected with other species. High virulence was evident within gp60 subtype IId but was absent in subtype families Ia and Ie. | There is no defined animal host for this species, although low density of the parasite has been detected occasionally in cattle, sheep and goat faeces. Experimental infection can be maintained in piglets and calves. Immunosuppressed Mongolian gerbils provide an asymptomatic small mammal model for infection. | [13] |
| C. macropodum (previously marsupial genotype II) | 5.4 × 4.9 | Eastern grey kangaroo | No association. | | [63] |
| C. meleagris | 5.2 × 4.6 | Homoeo-thermic birds; mammals | Infectivity data from experimental infections in adults indicate mild illness. Sporadic cases are reported more frequently in some populations, for example infections are as common as for C. parvum (10–20% of cryptosporidiosis cases) in Bangkok, Thailand and Lima, Peru; travel to endemic countries; possible outbreak on a farm in Sweden. | Associated with enteritis, diarrhoea and death in birds. Severe outbreaks and mortalities reported in poultry units and game birds. Asymptomatic infections and growth reductions have also been reported. Readily transmissible between birds and mammals. | [33,68,69] |
| C. molnari | 4.7 × 4.5 | Sea bream | No association. | | [70] |
### Table 1 (Continued)

| Cryptosporidium species and genotypes* | Mean oocyst dimensions (µm)ᵇ | Major host(s) | Association with human cryptosporidiosis [18] | Association with animal cryptosporidiosis [5,21,23] | Additional key Refs |
|--------------------------------------|-----------------------------|---------------|-----------------------------------------------|--------------------------------------------------|---------------------|
| C. muris                            | 7.0 × 5.0                   | Rodents       | Rarely associated; individual reports from various developing countries. | Clinical signs are not usually reported; infection and oocyst shedding has been reported in other animal species. | [13,71] |
| C. parvum (also sometimes previously called bovine genotype, genotype II, and genotype B) | 5.0 × 4.5                   | Humans, mammals | Common in sporadic cases and outbreaks; zoonotic transmission, either directly animal-to-person through recreational or occupational farm animal contact (especially young ruminants), or indirectly through contaminated drinking water, recreational water, environmental contact, food, or fomites. Person-to-person spread also occurs. Infectivity data from experimental infections in adults extrapolated to dose–response indicate that ingestion of a single oocyst carries a discrete probability of infection. The infectious dose with C. parvum in neonatal ruminants is similar. High heterogeneity; some subtypes (e.g., gp60 IIc) appear host-adapted to humans². | Common cause of diarrhoea in pre-weaning calves, lambs, and goats; infection can be fatal. Also reported in foals, alpaca, llama. Occasional respiratory symptoms. Disease is age-dependent, where older animals are usually asymptomatic. Some subtypes (e.g., gp60 IIc) have no defined animal host. Asymptomatic infectivity models include neonatal, immunosuppressed and immunocompromised mice. | |
| C. ryanae (previously deer-like genotype) | 3.7 × 3.2                   | Cattle        | No association. | Infection is host-specific and shedding is seen in weaned calves but is thought to be asymptomatic. | |
| C. scophthalmi†                     | 4.4 × 3.9                   | Turbot        | No association. | Infection occurs in the intestine and is seen seasonally in younger fish; has been linked to reduced growth rates. | [72] |
| C. scrofarum (previously pig genotype II) | 5.2 × 4.8                   | Pig           | Rarely associated; individual report from Czech Republic involving contact with pigs. | Infection is seen in pigs aged more than 6 weeks resulting in low levels of oocyst shedding and no association with diarrhoea. | [73] |
| C. serpensis                        | 6.2 × 5.3                   | Reptiles      | No association. | Infection is common. In snakes, infection manifests as anorexia, persistent postprandial regurgitation, lethargy, mid-body swelling, and chronic weight-loss, which is usually protracted and almost always fatal. In lizards infection is usually asymptomatic. | |
| C. suis (previously pig genotype I) | 4.6 × 4.2                   | Pig           | Rarely associated; individual reports from UK and Peru involving contact with pigs. | Infection is usually seen in pre-weaning pigs but is not associated with diarrhoea. | [73] |
The gut epithelium generally recovers after resolution of symptoms, but there are some indications that long-term sequelae may arise. For example, *C. hominis*, but not *C. parvum*, was associated with joint pain, eye pain, headaches, and fatigue during the 2 months following human infection [26]. Although a study of intestinal mucosal biopsies and serology from patients with inflammatory bowel disease (IBD) did not support a major role for *Cryptosporidium* in its pathogenesis [27], infection can trigger IBD flare, and inclusion of a test for *Cryptosporidium* is indicated in IBD patients seeking medical intervention for sudden exacerbation of digestive symptoms [28]. Experimental infection of rats with *C. parvum* triggers long-term pathological changes in the small intestine similar to those found in patients with irritable bowel syndrome (IBS) [29], supporting anecdotal reports of association. If a postinfectious (PI) association is found with *Cryptosporidium* for IBS (PI-IBS), reassurance could be provided to diagnosed patients that their altered bowel habit is consistent with PI-IBS and is likely to resolve without need for medical

| Cryptosporidium species and genotypes* | Mean oocyst dimensions (µm)† | Major host(s) | Association with human cryptosporidiosis [18] | Association with animal cryptosporidiosis (5,21,23) | Additional key Refs |
|---------------------------------------|-------------------------------|---------------|-----------------------------------------------|---------------------------------------------------|-------------------|
| *C. hyznderi* (previously mouse genotype l) | 4.6 × 4.2 | Mice | Rarely associated; individual report from Czech republic involving contact with wild mice. | Infection is usually most intense in the ileum but no clinical signs were recorded in experimental infections. | [74,75] |
| *C. ubiquitum* (previously cervine genotype) | 5.0 × 4.7 | Various mammals | Sporadic cases in various countries, especially developed countries, possibly involving untreated water supplies contaminated by animal hosts in the catchment. | Broad host-range and a common parasite of weaned lambs. No association of oocyst shedding and clinical symptoms in experimentally infected lambs. | |
| *C. viatorum* | 5.4 × 4.7 | Humans | Sporadic cases emerging in the UK and Sweden are linked to visits to the Indian subcontinent, South America, and Kenya. | No animal host is known for this species. | [76,77] |
| *C. varani* (syn. *C. saurophilum*) | 4.8 × 4.7 | Reptiles | No association. | Infection is seen in lizards and snakes. Clinical signs include anorexia, progressive weight-loss, abdominal swelling, and death, particularly in young animals. | |
| *C. wrairi* | 5.4 × 4.6 | Guinea pig | No association. | Disease not described. | [78] |
| *C. xiao* (previously *Cryptosporidium bovis*like genotype or *C. bovis* from sheep) | 3.9 × 3.4 | Sheep | No association. | In sheep, asymptomatic carriage; no association of oocyst shedding and clinical symptoms in experimentally infected lambs. Infection in goat kids can be associated with diarrhoea. | [79] |
| Chipmunk genotype I | Not reported | Chipmunk; possibly other Sciridae | Rarely associated; individual reports from the USA, France, Sweden. | No association of clinical signs with shedding oocysts. | |
| Horse genotype | 4.6 × 4.2 | Horses | Rarely associated; individual reports from the UK and USA | No association of oocyst shedding with diarrhoea. | [80] |
| *C. hominis* monkey genotype | Not reported | Monkey, human | Rarely associated; individual reports from UK and Malawi. | No association of oocyst shedding with diarrhoea recorded. | |
| Skunk genotype | Not reported | Skunk; possibly other mustelids | Rarely associated; an individual report from the UK where skunks are not found outside zoos. | No association of oocyst shedding with diarrhoea recorded. | |

*Only those genotypes found in humans have been included here.

†From the original papers describing the species or genotype.

*See [18] for an explanation of gp60 subtyping and nomenclature.

*Currently, *Cryptosporidium* is the only genus in the Family Cryptosporidiidae, but piscine species show considerable genetic distance, ultrastructural and developmental differences in comparison with other *Cryptosporidium* species, and a new genus, *Piscicryptosporidium*, has been proposed, pending study of additional piscine isolates [81].
### Table 2. Specimen types and applications of *Cryptosporidium* diagnostic tests

| Specimen type                                      | Relevant patient or animal group                                                                 | Sample preparation and standard test options                                                                 | Appropriate number of specimens | Test availability; key findings                                                                 |
|----------------------------------------------------|--------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------|---------------------------------|---------------------------------------------------------------------------------------------------|
| **Direct detection of parasite**                   | Patients/animals with diarrhea. Local policy may not include examination specifically for *Cryptosporidium* in all samples, even if a request for ‘ova, cysts, and parasites’ is made. Common selection criteria for examination include: *· Young age (for farmed animals, usually <1 month of age);* *· Recent travel to an endemic country;* *· Other risk factor; for example farm visits, local outbreak; immunocompromise.* Clinicians should familiarise themselves with local laboratory practice, and specify *Cryptosporidium* on the request form to ensure that appropriate testing is carried out. | Fresh, frozen or preserved faeces [e.g., in 10% formalin, merthiolate–iodine–formaldehyde; sodium acetate–acetic acid–formalin; other preservatives: e.g., polyvinyl alcohol (PVA) may optimise detection of other parasites, especially for the morphological examination of trophozoites, and preservation in different vials may be required]. Preservation pros: *· Preserves other parasite morphology or stages.* *· Allows transport at high ambient temperature.* *· Allows prolonged storage. Preservation cons: *· May impede PCR analysis, Strongyloides larvae concentration, and examination for trophozoites of other protozoans of interest.* *· Many immunosassays cannot be performed on PVA preserved samples.* *· Preserved faeces require concentration by a recognised method before microscopy. Some faeces may need additional processing, for example very liquid stools may be concentrated, high-fat faeces may require defatting, mucoid samples may need treatment with KOH or dithiothreitol, high-fibre faeces may need sieving to remove fibres. Test faeces or concentrate as shown in Figure 2. | For farm animals, ideally, collect faecal samples from a representative number of affected animals in the herd/flock. Repeat sampling may be required if samples are negative and *Cryptosporidium* is still suspected because shedding can be intermittent: *· Ideally, for humans, three stool specimens collected over no more than a 10 day period, usually collected every other day, before being deemed negative.* *· For animals, 2–3 consecutive negative faeces before confirmed to have cleared the infection. | Available locally. Prevalence among patients with gastrointestinal symptoms is 1–4% in Europe and North America and 3–20% in Africa, Asia, Australia, South and Central America [82]. Incidence rates may be underestimated because not all infections are symptomatic and persons with symptoms do not always seek medical attention, laboratory diagnosis may not be sought, laboratories may not include *Cryptosporidium* testing, and case reports or notifications may not be complete. In the UK, where most laboratories test all diarrhoeic stools for *Cryptosporidium*, the ratio of disease rates in the community and presenting to general practice relative to the rate of reported diagnoses to national surveillance) has been estimated to be 8.2 (95% CI 2.1–31.7) [42]. One of the most frequent causes of calf diarrhoea [25]. |
| Bowel content collected postmortem.               | Animals that died with signs of diarrhoea.                                                      | As above.                                                                                                      | As discussed with clinicians. | Available locally.                                                                                   |
| Small bowel scrapings or aspirates.              | Severely immunocompromised patients with persistent idiopathic gastrointestinal symptoms. Animals that died with clinical signs consistent with cryptosporidiosis. | Prepare a smear on a well slide and stain with immunofluorescent stains for oocysts and/or extract DNA and perform PCR. | As discussed with clinicians. | May be available locally; more likely refer for specialist testing.                                   |
| Small bowel biopsy or tissue sections collected postmortem. | As above.                                                                                         | Examine haematoxylin and eosin (H&E) histology sections for endogenous stages and pathology consistent with cryptosporidiosis and/or extract DNA and perform PCR. DNA quality from fixed tissue sections is reduced but can be used for PCR. | As discussed with clinicians. | As above.                                                                                           |
Table 2 (Continued)

| Specimen type                                      | Relevant patient or animal group                                                                 | Sample preparation and standard test options                                                                 | Appropriate number of specimens | Test availability; key findings |
|---------------------------------------------------|--------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------|--------------------------------|--------------------------------|
| Tracheal scrape collected postmortem.             | Birds that died with clinical signs consistent with respiratory cryptosporidiosis.              | As above.                                                                                                   | As above.                      | As above.                      |
| Bile from endoscopic retrograde cholangiopancreatography. | Severely immunocompromised patients with symptoms of cholangitis whose stool test is negative for Cryptosporidium. | Prepare a smear on a well slide and stain with immunofluorescent stains for oocysts and/or extract DNA and perform PCR. | As discussed with clinicians. | As above.                      |
| Liver biopsy, preferably in saline.               | Severely immunocompromised patients with symptoms of liver disease.                              | Examine H&E stained sections for endogenous stages and pathology consistent with infection and/or extract DNA and perform PCR. DNA quality from fixed tissue sections is reduced but can be used for PCR. | As discussed with clinicians. | As above.                      |
| Sputum or bronchoalveolar lavage.                 | Severely immunocompromised patients with unexplained respiratory symptoms.                      | May require pretreatment with dithiothreitol. Prepare a smear on a well slide and stain with immunofluorescent stains for oocysts and/or extract DNA and perform PCR. | As discussed with clinicians. | As above.                      |
| Antral washout.                                   | Severely immunocompromised patients with unexplained sinusitis.                                  | Prepare a smear on a well slide and stain with immunofluorescent stains for oocysts and/or extract DNA and perform PCR. | As above.                      | As above.                      |

Indirect detection of infection

Blood serum.  
Human populations for seroconversion, prevalence or seroepidemiological analysis.  
Large and mini-format Western blot, enzyme immunoassay (EIA), or microsphere assay incorporating recombinant proteins (e.g., 15/17 kD and 27 kD) for detection of Cryptosporidium-specific IgG and IgM and sometimes IgA.  
Depends on study design.  
Specialist test for research purposes. Strong IgG responses elicited to the 15/17 kD antigen occur within 6 months of infection; response to 27 kD antigen is usually much weaker but is detectable for longer [83].

Oral fluid.  
Populations for seroconversion, prevalence, or seroepidemiological analysis.  
Dissociation-enhanced lanthanide fluorescence immunoassay (DELFIA) or microsphere assays incorporating recombinant proteins for detection of Cryptosporidium-specific IgG and IgA.  
Depends on study design.  
A specialist test is under development for population monitoring [50]. Anti-Cryptosporidium IgG seroconversion reported in infected children [84]; detection of anti-Cryptosporidium IgG in oral fluid has been associated with recent diarrhoea [88].

Faeces/stool.  
Indication of active infection.  
Copro-antibody EIA for detection of Cryptosporidium-specific IgA.  
Depends on study design.  
Used for research purposes.

intervention. Potential impacts include reduction in use of health services (e.g., consultations and investigations).

In malnourished children, who are effectively immunocompromised, infection has been reported to cause substantial morbidity even for those with no acute symptoms [6]. Patients with severe immunodeficiency may suffer from chronic, severe and intractable cryptosporidiosis with significant mortality (reviewed in [30]). At most risk are those with primary or secondary T cell immunodeficiencies, notably untreated advanced HIV infection, lymphoma, and leukaemia (especially children). In this patient group, complications include pancreato-biliary infection, which may lead to sclerosing cholangitis and liver cirrhosis, and respiratory cryptosporidiosis (Table 2). Rapid detection of carriage in high-risk groups could help to limit clinical sequelae [31]. The findings of a recent study indicate that even low
doses of *C. parvum* infect dexamethasone-treated, severe combined immunodeficiency (SCID) mice, leading to digestive adenocarcinoma, implying that severely immunocompromised humans may also be susceptible [32], which highlights the need for vigilance in diagnosis.

In otherwise healthy adults, no significant difference in acute symptoms caused by *C. parvum, C. hominis*, and *C. meleagris* has been found [33]. However, species- and subtype-related differences in clinical manifestations have been reported in children and immunocompromised patients, indicating more severe disease with *C. hominis*, and some subtypes in particular, than with *C. parvum* (Table 1) [18]. A combination of parasite- and host-related factors influence the outcome of infection. Pathogenicity and virulence have been reviewed recently [34], and the reader is referred to reviews of host genetic susceptibility [35], host cell [36] and immune responses [37].

Asymptomatic carriage of *Cryptosporidium* occurs, and in humans has been mainly investigated in young children, including those in industrialized countries where carriage rates of up to 3.8% have been reported [38]. A study of children in daycare in the UK used highly-sensitive immunomagnetic separation (IMS) and direct immunofluorescence microscopy (IFM), and reported concentrations of 13367 *C. hominis* oocysts per gram (opg), 3471 *C. ubiquitum* opg, and 56 skunk genotype opg [38]. Only the *C. hominis*-positive sample contained sufficient oocysts for detection by some current methods (Figure 2), and low parasite density challenges most diagnostic assays (Box 1). This is also an issue for samples obtained late in an infection when oocyst numbers may be in decline or where partial immunity may result in few oocysts being shed [31].

Supportive therapy to prevent dehydration and restore electrolyte balance may be needed, but specific drug therapy is not usually required for the immunocompetent. Young children and immunocompromised patients may not resolve symptoms, and nitazoxanide is licensed by the US Food and Drug Administration (FDA) for treatment of cryptosporidiosis in immunocompetent individuals over 1 year of age; the latest Cochrane review confirmed the absence of evidence for effective agents for immunocompromised patients [39], which remains a key challenge. Immune reconstitution is most effective but not always possible. Halofuginone is licensed for the treatment and control of cryptosporidiosis in calves. However, this drug is toxic at twice the recommended dose and is contraindicated for use in diarrhoeic animals. A recent meta-analysis found that although it provided beneficial prophylactic use, there were insufficient data to establish a therapeutic effect [40]. However, a study in Ireland showed that halofuginone was effective in reducing clinical signs and environmental contamination with oocysts [41]. Screening for parasite clearance following treatment may require the more sensitive assays (Box 1, Figure 1).

**Recent advances incorporated in clinical diagnosis**

To diagnose enteric cryptosporidiosis a faecal sample must be tested, requiring a series of events to occur in the clinical consultation, specimen requesting, sampling, submission, testing and reporting process [42]. *Cryptosporidium* should be suspected in all patients who present with acute gastroenteritis, most especially young children or if the symptoms are prolonged (>3 days) [43], and in all neonatal ruminant diarrhoea; samples are also likely to include postmortem examinations of animals that died. Detection is very difficult in routine parasitological examinations (Figure 2) and, as with most pathogens, specific testing is often selective according to local policy [44].

The diagnostic trend during the last decade has been towards greater use of enzyme immunoassays (EIA) for detecting oocyst antigens [44], but there are no immunological tools that distinguish between oocysts of different species. One approach to overcoming this has been the commercial development of a PCR-EIA that identifies *Cryptosporidium* spp., *C. hominis* and *C. parvum* [45]. However, molecular diagnostic assays are being implemented in diagnostic laboratories, especially in virology and some aspects of bacteriology, and are emerging for enteric protozoa. Real-time PCR probe-based assays for the detection of multiple protozoa have been described, such as for *Cryptosporidium* spp., *Giardia lamblia*, *Entamoeba histolytica*, and *Dientamoeba fragilis*, with a testing algorithm for microscopy based on clinical information, which could be adapted to specific settings depending on the likelihood of involvement of other parasites [46]. The replacement of probes with carboxylated LumineX microspheres provides an alternative detection system, as demonstrated for several parasites [47]. An assay based on multiplex-tandem PCR (MT-PCR) describes the simultaneous detection of bacterial pathogens in *Cryptosporidium*-positive stools, and the authors suggest that the increased amplification efficiency would also be useful for water samples [48]. Commercial multiplex PCR-based assays for comprehensive panels of gastrointestinal pathogens are becoming available, utilising robotic platforms for DNA extraction and assay set-up. Although offering streamlined and high testing capacity, investigation of a wide range of pathogens in all samples increases the number of positive findings; this may not always be warranted clinically but may shed light on the importance of infection in patients previously not investigated, and of co-infections. Pathogen panels should ideally be tailored for the population under investigation. Downstream testing of extracted DNA for further species/genotype discrimination of *Cryptosporidium*-positive samples either in the same, or more likely a reference, laboratory could be readily implemented for epidemiological and outbreak investigations (Figure 2).

Thus, a variety of methods are used for *Cryptosporidium* diagnosis, as Fournet and colleagues discovered in their 2012 investigation: fluorescence microscopy (FM) using Auramine O (UK) or acid fast (AF) modified Ziehl–Neelsen staining (NL, UK, and Germany), EIA (NL, UK, and Germany), and multiplex PCR (NL) [15]. Species determination was performed routinely on a higher proportion of cases in the UK than the NL and not at all in Germany. Subtyping a proportion of the predominant *C. hominis* isolates at the gp60 locus [18] revealed mainly subtype IbA10G2, which predominates in these countries, and multilocus genotyping is underway. The investigation
Box 1. Attributes of current Cryptosporidium faecal diagnostic assays

- Selection criteria include resources (staff time, expertise, grade, finances, specialist equipment), desired turnaround time, the population being tested, desired sensitivity and specificity, and the need to identify other pathogens.
- Preservatives can inhibit PCR and must be removed by washing; this may not be possible if the preservative has penetrated the oocysts.
- Faeces can be concentrated by sedimentation using modified formol-ethyl acetate or solvent-free techniques, but significantly fewer parasites and larger deposits have been reported in a solvent-free system, and low-density parasites might be missed [86]. Conventional flotation (e.g., zinc sulphate, saturated sodium chloride or sucrose) results in cleaner preparations but low-density oocysts may be missed.
- Using fluorescent stains (e.g., Auramine O) oocyst observation is easier and less subjective, and slides can be scanned at lower magnification than acid-fast stained slides, but a fluorescence microscope is needed.
- IFM is less subjective than other microscopy stains and increases sensitivity and specificity, although some notable cross-reactions (e.g., algae) may occur in food, water, and environmental testing, necessitating observation of confirmatory features: internal contents and sporozoites by differential interference contrast and sporozoite nuclei by staining with a nuclear fluorogen.
- EIAs can be automated and are more sensitive than acid-fast microscopy. Immunochromatographic lateral flow (ICLF) is less sensitive, but requires little skill and can be performed anywhere as a rapid test, for example pen-side. ICLF and EIA kits may include other pathogenic protozoa in a single test (e.g., Giardia [87]), and additionally for humans, Entamoeba histolytica [88].

 Cryptosporidium species antigen variability may compromise immunodiagnostic tests based on antibodies raised against Cryptosporidium parvum; there is some evidence that Cryptosporidium felis may be less well detected [89].
- Conventional and real-time PCR assays have been described, but real-time PCR has been reported as the most sensitive and specific test for Cryptosporidium in diarrhoeal calf samples [90].
- Molecular assays are underpinned by efficient DNA extraction but oocysts may be present at low density in complex matrices (faeces, slurry, food, water) that may contain PCR inhibitors. Oocysts in faeces or semi-purified suspensions need first to be disrupted by procedures such as bead beating, freeze–thaw, cycles or enzymatic digestion [91]. Automated DNA extraction procedures can be used [48].
- PCR conditions must include mitigation of potentially inhibitory substances (e.g., heme, bilirubin, bile salts, or complex carbohydrates). PCR primers and conditions need to be selected carefully to amplify all Cryptosporidium species of interest and to avoid non-specific amplification. There is no gold-standard diagnostic test. For comparison of sensitivity and specificity, either one must be nominated or comparison tests such as McNemar’s or a Bayesian model must be applied [90]. The outcome depends not only on the test properties but also the nature of the samples (e.g., consistency, parasite density), the statistical power of the study, and skill in performing assays. Nevertheless, when performed under diagnostic conditions, the general consensus is as shown below (Figure 1).

Figure 1. Relative attributes of features of diagnostic assays, ranked low to high. Abbreviations: AF, acid fast; EIA, enzyme immunoassay; FM, fluorescence microscopy; ICLF, immunochromatographic lateral flow; IFM, immunofluorescence microscopy.

highlights the need for wider application of molecular epidemiology across the European Union.

Detection in food, water and environmental samples
Testing non-clinical samples for Cryptosporidium supports risk assessment and surveillance, and with the application of typing assists in the investigation of possible sources of contamination and routes of transmission, and provides microbiological evidence in outbreak investigations. In the USA, classification of waterborne outbreaks according to the strength of epidemiological, clinical, and environmental data now reflects the increasing use of molecular characterization of pathogens both in clinical and environmental samples [49]. However, outbreak investigations focus on specific incidents, and the contamination event will usually have passed some time before recognition of an outbreak. For understanding Cryptosporidium spp. transmission, baseline data need to be generated by well-structured sample surveys. Interventions such as those implemented to reduce waterborne disease need to be monitored for effectiveness, and one strategy is to compare population exposure. Various
serological methods have been investigated for *Cryptosporidium* antibodies (Table 2), which are moving in focus towards multiplex platforms to facilitate monitoring for a range of pathogens [50].

Standard methods for the detection and enumeration of *Cryptosporidium* in water are based on principles of large-volume sampling (10–1000 L), concentration by filtration and elution from the filter (or by flocculation for volumes <20 L), separation of oocysts from other particulates by IMS, and detection by IFM incorporating a nucleic acid stain as an identification aid ([51] for example). Alternative methods such as continuous-flow centrifugation have been shown to provide at least equivalent recovery efficiencies [51]. A similar sequence of oocyst recovery and detection from surface eluates of fresh food produce [2,17] forms the basis of a method currently being considered as an International Organization for Standardization (ISO) standard. However, the process is time-consuming and expensive. The production of automated, miniaturised systems has been sought as an alternative to laboratory testing of water samples, based on a variety of detection technologies including optical, mass, electrical, surface plasmon resonance, and nucleic acid detection. However, few have been performance-tested in the field, and a key step is sample preparation, still requiring IMS to deliver oocysts to the system [52].

Molecular detection of *Cryptosporidium* has not yet been adopted for regulatory purposes partly because only oocysts containing sporozoite DNA are detected, and no reliably quantitative method has been validated to replace oocyst counts. The small number of oocysts present in most water samples is challenging. One risk-assessment approach is to screen samples for a range of protozoa using universal primers, with fluorescent melt-curve analysis differentiating between genera [53]. Although a potentially valuable screening tool, the detection of less-abundant targets in field samples requires validation.

Standard methods provide no information about the species, virulence, or viability of oocysts; *Cryptosporidium* spp. need to be identified in water monitoring otherwise all oocysts detected must be assumed to present a public health risk. Sequencing DNA from nested PCR of the SSU rRNA gene, using a repetitive and limiting PCR template approach to improve analytical sensitivity and differentiation of multiple species or genotypes in a single sample, provides the benchmark [54]. However, this is technically demanding, and is outwith the scope of most water-testing laboratories. A simplified approach is being validated using single-round, conventional, or real-time PCR of the SSU rRNA and hsp70 genes to differentiate the main human pathogens *C. hominis*, *C. parvum*, and *C. meleagris* from other species commonly found in the environment (http://waterrr.org/Projects.aspx?PID=4099). A non-PCR-based method is loop-mediated isothermal amplification (LAMP), based on autocyclic strand-displacement DNA synthesis by Bst polymerase, and this may overcome the inhibitory potential of some samples and has been reported to detect organisms at relatively low concentrations in water samples [55], although it has not yet been subjected to independent evaluation.

**Impact of genomics on detection, diagnosis, and typing**

Molecular biology had a vast impact on the understanding of the species concept within the genus *Cryptosporidium*, resulting in a dramatic increase in species numbers and genotypes [21]. As a result, there is a need to understand the level of within-species genetic variation such that selected DNA sequences can be indentified as references. This is particularly important because a single locus (the SSU rRNA gene) is often used as species marker; confidence in species determination and subtyping is greatly increased by using multiple markers that are spread throughout the genome [56]. Phylogenetic analysis has proven to be an essential tool for determining species status; this approach provides more information than is achieved through individual sequence comparisons (such as BLAST searches) because more weight is given to informative mutations that may be species- or group-specific [20,54]. Use of reference sequences will help to reduce the confusion caused by sequences deposited in databases that may have been falsely attributed to a species due to limited understanding of species complexity at the time of submission. This is a considerable problem for a genus in which species numbers are still increasing.

Comparative analyses of the published *C. parvum* and *C. hominis* genomes (http://cryptodb.org) have revealed new possible diagnostic targets, for example a new family of telomERICally-encoded *Cryptosporidium* proteins, the *C. parvum*-specific Cops-1 and the *C. hominis*-specific Chos-1 [34]. The relatively small *Cryptosporidium* genome (9.1–9.2 Mb) combined with ever-reducing sequencing costs will allow more *Cryptosporidium* genomes to be sequenced, and from different species and from multiple isolates of the same species [57]. Publication of new genomes will allow detailed comparisons, resulting in identification of more species-specific markers, polymorphic mini- and micro-satellite markers, and single-nucleotide polymorphisms (SNPs) [58]. These markers can be exploited to develop improved PCR-based diagnostics utilising the more informative parts of the *Cryptosporidium* genome [16]. The SNPs may also be used in attempts to identify virulence genes or factors involved in host-specificity [58]. The increasing amount of genomic data will undoubtedly lead to further development of microarrays for detecting variation within parasite populations. However, this is a longer-term development and currently there is much focus on the use of mini- and micro-satellite markers. Generally, these are very variable because they are commonly in noncoding regions of the genome and are subject to fewer functional restraints that impede the accumulation of mutations. However, many *Cryptosporidium* satellite markers were identified before genome sequences became available, and are mostly within coding regions [59]. These satellite markers have been used successfully for studying *Cryptosporidium* population dynamics to evaluate transmission routes and zoonotic potential [59–62]. There is a need not only for the further development of subtyping methods for epidemiological purposes, but also for their standardisation. A recent review revealed that 55 markers have been used in various combinations on different platforms to investigate variation within *C. parvum* and *C. hominis* [59]. Although multi-locus typing is a potentially powerful
tool, global meta-analysis and comparisons are impossible if different methods are used, and there is a need for standardisation. Assessment of existing markers and suggestions for taking these forward in independent validation studies for harmonisation of typing schemes have been published recently by Robinson and Chalmers [59]. Typing tools developed based on Cryptosporidium genome analyses, if harmonised, will also have the potential to help with outbreak investigations, source attribution, investigations into the fate and transport of oocysts, persistence within hosts, studies into the role of strain-specific immunity, and the identification of virulence factors and genes involved in host specificity.

Concluding remarks
Cryptosporidium presents many challenges for detection and diagnosis, and the use of different diagnostic methods and the inconsistent application of typing techniques can make direct comparisons difficult or even impossible between clinical, veterinary, and environmental testing or between different regions or countries. Adoption of standardised techniques, where possible, for diagnosis and typing would allow better correlations between animal, human, and environmental data at national and international levels. This would aid source-tracking of outbreaks and enable better understanding of transmission of the parasite.

The absence of a gold-standard diagnostic test for comparison of sensitivity and specificity can be overcome either by nominating one or using appropriate statistical tests for comparison (Box 1). Small numbers of parasites may be clinically significant, and one of the challenges faced by scientists and clinicians working with Cryptosporidium is to increase the sensitivity of existing methods; this applies particularly to molecular detection and typing from water samples where the ultimate goal is to reliably speciate and multi-locus-type samples containing single oocysts.

Expected advances for clinical diagnosis of Cryptosporidium infections may come from the direct application of next-generation sequencing to faecal samples because this platform will allow the detection of multiple pathogens and their variants based directly on the presence of their DNA. More imminent is the implementation of molecular diagnostics. However, adopting these new technologies in a clinical laboratory involves significant changes to the current workflows, as depicted in Figure 2.

From the research community, the availability of genome sequences for different Cryptosporidium species and multiple genome sequences for the most clinically and economically important species is imminent. These genome datasets will facilitate the development of better diagnostic and typing tools at inter- and intra-species level, and efforts must be made to ensure that technological advances are also implemented in low-income countries where the burden of cryptosporidiosis is highest.

References
1 Baldursson, S. and Karanis, P. (2011) Waterborne transmission of protozoan parasites: review of worldwide outbreaks – an update 2004–2010. Water Res. 45, 6603–6614
2 Robertson, L.J. and Chalmers, R.M. (2013) Foodborne cryptosporidiosis: is there really more in Nordic countries? Trends Parasitol. 29, 3–9
3 Artieda, J. et al. (2012) Outbreak of cryptosporidiosis in a child day-care centre in Gipuzkoa, Spain, October to December 2011. Euro Surveill. 17, pii=20670
4 Gormley, F.J. et al. (2011) Zoonotic cryptosporidiosis from petting farms, England and Wales, 1992-2009. Emerg. Infec. Dis. 17, 151–152
5 Santín, M. (2013) Clinical and subclinical infections with Cryptosporidium in animals. N. Z. Vet. J. 61, 1–10
6 Shirley, D.A. et al. (2012) Burden of disease from cryptosporidiosis Curr. Opin. Infect. Dis. 25, 555–563
7 Featherstone, C.A. et al. (2010) Cryptosporidium species in calves submitted for postmortem examination in England and Wales. Vet. Rec. 167, 979–980
8 Savioli, L. et al. (2006) Giardia and Cryptosporidium join the ‘Neglected Diseases Initiative’. Trends Parasitol. 22, 203–208
9 Sreiner, T.S. and Mandell, G.L. (2009) Principles and syndromes of enteric infection. In Mandell, Douglas, and Bennett’s Principles and Practice of Infectious Diseases (7th edn) (Mandell, G.L. et al., eds), pp. 1335–1351, Churchill Livingstone
10 Ferreira, J. et al. (2011) Cryptosporidiosis masquerading as a Crohn’s flare. Inflamm. Bowel Dis. 17, E133–E134
11 Garcia, L.S. (2007) Diagnostic Medical Parasitology. ASM Press
12 Chalmers, R.M. (2012) Waterborne cryptosporidiosis outbreaks. Ann. Int. Super. Sanit. 48, 429–446
13 Medema, G. (2009) Risk Assessment of Cryptosporidium in Drinking-Water. World Health Organization
14 World Health Organization (2011) Guidelines for Drinking-Water Quality. (4th edn), WHO
15 Fournet, N. et al. (2013) Simultaneous increase of Cryptosporidium infections in the Netherlands, the United Kingdom and Germany in late summer season, 2012. Euro Surveill. 18, pii=20348
16 Jex, A. et al. (2008) Cryptosporidium – biotechnological advances in the detection, diagnosis and analysis of genetic variation. Adv. Parasitol. 77, 141–173
17 Smith, H. and Nichols, R.A. (2010) Cryptosporidium: detection in water and food. Exp. Parasitol. 124, 61–79
18 Xiao, L. (2010) Molecular epidemiology of cryptosporidiosis: An update. Exp. Parasitol. 124, 80–89
19 Bart, J.R. and Thompson, R.C.A. (2006) What is Cryptosporidium? Reappraising its biology and phylogenetic affinities. Trends Parasitol. 22, 463–468
20 Jirku et al. (2008) New species of Cryptosporidium Tytzer, 1907 (Apicomplexa) from amphibian host: morphology, biology and pylonogy. Folia Parasitol. 55, 81–94
21 Fayer, R. (2010) Taxonomy and species delimitation in Cryptosporidium. Exp. Parasitol. 124, 90–97
22 Xiao, L. et al. (2012) Cryptosporidium tyzzeri and Cryptosporidium pestis: which name is valid? Exp. Parasitol. 130, 308–309
23 Xiao, L. and Feng, Y. (2008) Zoonotic cryptosporidiosis. FEMS Immunol. Med. Microbiol. 52, 1–15
24 Chalmers, R.M. and Davies, A.P. (2010) Minireview: clinical cryptosporidiosis. Exp. Parasitol. 124, 138–146
25 Lorenz, I. et al. (2011) Calf health from birth to weaning. II. Management of diarrhoea in pre-weaned calves. Ir. Vet. J. 64, 9
26 Hunter, P.R. et al. (2004) Health sequelae of human cryptosporidiosis in immunocompetent patients. Clin. Infect. Dis. 39, 504–510
27 Chen, W. et al. (2001) Cryptosporidium parvum in intestinal mucosal biopsies from patients with inflammatory bowel disease. Am. J. Gastroenterol. 96, 3463–3464
28 Colussi, O. et al. (2010) Acute cryptosporidiosis as a cause of sudden recurrence of digestive symptoms in patients with Crohn’s disease. J. Crohns Collitis 4, 669–670
29 Khalil, S. et al. (2009) Cryptosporidium parvum isolate-dependent postinfectious jejunal hypersensitivity and mast cell accumulation in an immunocompetent rat model. Infect. Immun. 77, 5163–5169
30 Hunter, P.R. and Nichols, G. (2002) Epidemiology and clinical features of Cryptosporidium infection in immunocompromised patients. Clin. Microbiol. Rev. 15, 145–154
31 McLaughlin, J. et al. (2003) Polymerase chain reaction-based diagnosis of infection with Cryptosporidium in children with primary immunodeficiencies. Pediatr. Infect. Dis. J. 22, 329–334
32 Benamrouz, S. et al. (2012) Cryptosporidium parvum infection in SCID mice infected with only one oocyst: qPCR assessment of parasite...
replication in tissues and development of digestive cancer. PLoS ONE 7, e51292
33 Chappell, C.L. et al. (2011) Cryptosporidium meleagridis: infectivity in healthy adult volunteers. Am. J. Trop. Med. Hyg. 85, 238–242
34 Bouzid, M. et al. (2013) Cryptosporidium pathogenicity and molecular virulence factors. Clin. Microbiol. Rev. 26, 115–134
35 Flores, J. and Okhuysen, P.C. (2009) Host factors – genetics of susceptibility to infection with enteric pathogens. Curr. Opin. Infect. Dis. 22, 471–476
36 Yang, Y-L. et al. (2010) Cell sorting-assisted microarray profiling of host cell responses to Cryptosporidium parvum infection. Infect. Immun. 78, 1040–1048
37 Borad, A. and Ward, H. (2010) Human immune responses in cryptosporidiosis. Future Microbiol. 5, 507–519
38 Davies, A.P. et al. (2009) Asymptomatic carriage of protozoan parasites in children in day care centers in the United Kingdom. Paediatr. Infect. Dis. J. 28, 838–840
39 Abubakar, I. et al. (2007) Treatment of cryptosporidiosis in immunocompromised individuals: systematic review and meta-analysis. Br. J. Clin. Pharmacol. 63, 387–393
40 Silverlás, C. et al. (2009) Systematic review and meta-analyses of the effects of halofuginone against calf cryptosporidiosis. Prev. Vet. Med. 91, 73–84
41 De Waele, V. et al. (2010) Control of cryptosporidiosis in neonatal calves: use of halofuginone lactate in two different calf rearing systems. Prev. Vet. Med. 96, 143–151
42 Tam, C.C. et al. (2012) Longitudinal study of infectious intestinal disease in the UK (ID2 study): incidence in the community and presenting to general practice. Gut 61, 69–77
43 Guerrant, R.L. et al. (2001) Infectious Diseases Society of America. Practice guidelines for the management of infectious diarrhea. Clin. Infect. Dis. 32, 331–351
44 Chalmers, R.M. et al. (2010) Clinical laboratory practices for the detection and reporting of Cryptosporidium in community cases of diarrhoea in the United Kingdom, 2008. Euro Surveill. 15, pii=519731
45 Savin et al. (2008) Assessment of cryptodiagnosis for diagnosis of cryptosporidiosis and genotyping Cryptosporidium species. J. Clin. Microbiol. 46, 2590–2594
46 Brijnjesťijn van Coppenraet, L.E. et al. (2009) Parasitological diagnosis combining an internally controlled real-time PCR assay for the detection of four protozoa in stool samples with a testing algorithm for microscopy. Clin. Microbiol. Infect. 15, 869–874
47 Tanijchi et al. (2011) High throughput multiplex PCR and probe-based detection with Luminex beads for seven intestinal parasites. Am. J. Trop. Med. Hyg. 2, 322–327
48 Joux, A. et al. (2012) Detection of diarrhoeal pathogens in human faeces using an automated, robotic platform. Mol. Cell. Probes 26, 11–15
49 Centers for Disease Control (2011) Surveillance for waterborne disease outbreaks and other health events associated with recreational water – United States, 2007–2008 and surveillance for waterborne disease outbreaks associated with drinking water – United States, 2007–2008. MMWR Morb. Mortal. Wkly. Rep. 60, 1–75
50 Lamnie, P.J. et al. (2012) Development of a new platform for neglected tropical disease surveillance. Int. J. Parasitol. 42, 797–800
51 Environmental Protection Agency (2005) Method 1623: Cryptosporidium and Giardia in Water by Filtration/IMS/FA. US EPA
52 Bridle, H. et al. (2012) Detection of Cryptosporidium in miniaturised fluidic devices. Water Res. 46, 1641–1661
53 Lalonde, L.F. and Gajadhar, A.A. (2012) Detection and differentiation of coccidian oocysts by real-time PCR and melting curve analysis. J. Parasitol. 97, 725–730
54 Rueckert, N.J. et al. (2012) Molecular and phylogenetic approaches for assessing sources of Cryptosporidium contamination in water. Water Res. 46, 5135–5150
55 Bakheit, M. et al. (2008) Sensitive and specific detection of Cryptosporidium species in PCR-negative samples by loop-mediated isothermal DNA amplification and confirmation of generated LAMP products by sequencing. Vet. Parasitol. 25, 11–22
56 Widmer, G. and Lee, Y. (2010) Comparison of single- and multilocus genetic diversity in the protozoan parasites Cryptosporidium parvum and C. hominis. Appl. Environ. Microbiol. 76, 6639–6644
57 Widmer, G. and Sullivan, S. (2012) Genomics and population biology of Cryptosporidium species. Parasite Immunol. 34, 61–71
58 Widmer, G. et al. (2012) Comparative genome analysis of two Cryptosporidium parvum isolates with different host range. Infect. Genet. Evol. 12, 1213–1221
59 Robinson, G. and Chalmers, R.M. (2012) Assessment of polymorphic genetic markers for multi-locus typing of Cryptosporidium parvum and Cryptosporidium hominis. Exp. Parasitol. 132, 200–215
60 Morrison, L.J. et al. (2008) The population structure of the Cryptosporidium parvum population in Scotland: a complex picture. J. Parasitol. 94, 1217–1221
61 Tanriverdi, S. et al. (2008) Inferences about the global population structures of Cryptosporidium parvum and Cryptosporidium hominis. Appl. Environ. Microbiol. 74, 7227–7234
62 Drumbo, R. et al. (2012) Evidence of host-associated populations of Cryptosporidium parvum in Italy. Appl. Environ. Microbiol. 78, 3523–3529
63 Ryan, U. and Power, M. (2012) Cryptosporidium species in Australian wildlife and domestic animals. Parasitology 20, 1–16
64 Khan, S.M. et al. (2010) Molecular characterization and assessment of zoonotic transmission of Cryptosporidium from dairy cattle in West Bengal, India. Vet. Parasitol. 171, 41–47
65 The ANOVEL Cryptosporidium National Network (2010) Laboratory-based surveillance for Cryptosporidium in France, 2006–2009. Euro Surveill. 15, pii=19642
66 Molloy, S.F. (2010) Identification of a high diversity of Cryptosporidium species genotypes and subtypes in a pediatric population in Nigeria. Am. J. Trop. Med. Hyg. 82, 608–613
67 Robinson, G. et al. (2010) Re-description of Cryptosporidium caninum Inman and Takeuchi, 1979 (Apicomplexa: Cryptosporidiidae): morphology, biology and phylogeny. Int. J. Parasitol. 40, 1539–1548
68 Elwin, K. et al. (2011) The epidemiology of sporadic human infections with unusual cryptosporidia detected during routine typing in England and Wales, 2000 to 2008. Epidemiol. Infect. 140, 673–683
69 Silverlás, C. et al. (2012) Zoonotic transmission of Cryptosporidium meleagridis on an organic Swedish farm. Int. J. Parasitol. 42, 963–967
70 Sinjí-Bebadilla, A. et al. (2005) Epidemiology of Cryptosporidium molnari in Spanish gilthead sea bream (Sparus aurata L.) and European sea bass (Dicentrarchus labrax L.) cultures: from hatchery to market size. Appl. Environ. Microbiol. 71, 131–139
71 Blewett, D.A. et al. (1993) Infective dose size studies on Cryptosporidium parvum using gnotobiotic lambs. Water Sci. Technol. 27, 61
72 Alvarez-Pellitero, P. et al. (2009) Host and environmental risk factors associated with Cryptosporidium scophthalmi (Apicomplexa) infection in cultured turbot, Psetta maxima (L.) (Piscis, Teleostei). Vet. Parasitol. 165, 207–215
73 Némecj, K. et al. (2013) Occurrence of Cryptosporidium suis and Cryptosporidium scrofarum on commercial swine farms in the Czech Republic and its associations with age and husbandry practices. Parasitol. Res. 112, 1143–1154
74 Ren, X. et al. (2012) Cryptosporidium tyzzeri n. sp. (Apicomplexa: Cryptosporidiidae) in domestic mice (Mus musculus). Exp. Parasitol. 130, 274–281
75 Raskóvá, V. et al. (2013) Case of human cryptosporidiosis caused by Cryptosporidium tyzzeri and C. parvum presumably transmitted from wild mice. J. Clin. Microbiol. 51, 360–362
76 Elwin, K. et al. (2012) Cryptosporidium viatorum n. sp. (Apicomplexa: Cryptosporidiidae) among travellers returning to the United Kingdom from the Indian Subcontinent. Int. J. Parasitol. 42, 675–682
77 Insulander, M. et al. (2012) Molecular epidemiology and clinical manifestations of human cryptosporidiosis in Sweden. Epidemiol. Infect. 9, 1–12
78 Lv, C. et al. (2009) Cryptosporidium spp. in wild, laboratory, and pet rodents in China: prevalence and molecular characterization. Appl. Environ. Microbiol. 75, 7692–7699
79 Rieux, A. (2013) Molecular characterization of Cryptosporidium spp. in pre-weaned kids in a dairy goat farm in western France. Vet. Parasitol. 192, 268–272
80 Burton, A.J. et al. (2010) The prevalence of Cryptosporidium, and identification of the Cryptosporidium horse genotype in foals in New York State. Vet. Parasitol. 174, 139–144
Palenzuela, O. et al. (2010) Molecular characterization of Cryptosporidium molnari reveals a distinct piscine clade. Appl. Environ. Microbiol. 76, 7646–7649

Current, W.L. and Garcia, L.S. (1991) Cryptosporidiosis. Clin. Microbiol. Rev. 4, 325–358

Priest, J.W. et al. (2010) Multiplex assay detection of immunoglobulin G antibodies that recognize Giardia intestinalis and Cryptosporidium parvum antigens. Clin. Vaccine Immunol. 17, 1695–1707

Griffin, S. et al. (2011) Development of a multiplex microsphere immunoassay for the quantitation of salivary antibody responses to selected waterborne pathogens. J. Immunol. Methods 364, 83–93

Egorov, A. et al. (2010) Recent diarrhea is associated with elevated salivary IgG responses to Cryptosporidium in residents of an Eastern Massachusetts community. Infection 38, 117–123

Saez, A.C. et al. (2011) Comparison between the Midi Parasep and Midi Parasep Solvent Free (SF) faecal parasite concentrators. J. Clin. Pathol. 64, 901-904

Budu-Amoako, E. et al. (2012) Occurrence of Cryptosporidium and Giardia on beef farms and water sources within the vicinity of the farms on Prince Edward Island, Canada. Vet. Parasitol. 28, 1–9

Goti, P. et al. (2012) Evaluation of an immunochromatographic dip strip test for simultaneous detection of Cryptosporidium spp, Giardia duodenalis, and Entamoeba histolytica antigens in human faecal samples. Eur. J. Clin. Microbiol. Infect. Dis. 31, 2077–2082

Agnamey, P. et al. (2011) Evaluation of four commercial rapid immunochromatographic assays for detection of Cryptosporidium antigens in stool samples: a blind multicenter trial. J. Clin. Microbiol. 49, 1605–1607

de Waele, V. et al. (2011) Age-stratified Bayesian analysis to estimate sensitivity and specificity of four diagnostic tests for detection of Cryptosporidium oocysts in neonatal calves. J. Clin. Microbiol. 49, 76–84

Elwin, K. et al. (2012) A comparison of two approaches to extracting Cryptosporidium DNA from human stools as measured by a real-time PCR assay. J. Microbiol. Methods 89, 38–40