HIV medicine

Cryptosporidiosis in persons with HIV infection

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*Cryptosporidium parvum* is a coccidian protozoan parasite that belongs to the same family as *Isospora* and *Toxoplasma*. Since its discovery by Tyzzer in 1907 and until the advent of autoimmune deficiency syndrome (AIDS), *Cryptosporidium* was primarily regarded as a cause of diarrhoea among animals. Indeed, before 1982, only seven cases of human cryptosporidiosis had been reported in the medical literature. The last 15 years has witnessed a dramatic increase in the number of cases, owing to better diagnostic methods and a growing population of individuals infected with human immunodeficiency virus (HIV).

**Aetiology**

Symptoms of cryptosporidial infection begin 5–22 days after ingestion of as few as 30 oocysts of human, animal or environmental origin. After ingestion, oocysts excyst and release sporozoites, the infective form of the parasite. The sporozoites implant on the mucosal surface of host epithelial cells and undergo sexual and asexual reproduction with generation of oocysts and merozoites, respectively. Merozoites continue the process of mucosal auto-infection while oocysts are immediately infectious and are excreted in stools. Note that the entire life-cycle takes place within a single host. Though the most common site of infection is the gut, extra-intestinal infection has been reported rarely, most commonly in the respiratory tract.

The precise mechanism by which *Cryptosporidium* causes diarrhoea is unknown. Ultrastructural studies reveal that, while the parasite attaches to the mucosal epithelial lining, it does not invade the submucosa.

Mechanisms that promote clearance of *C parvum* remain unclear. On the basis of clinical and experimental data, it is generally accepted that both the humoral and cellular immune systems are required for control of infection. Studies using several animal models suggest that loss of T-lymphocyte function may lead to persistent cryptosporidiosis. The importance of B-cell-mediated immunity is suggested by the fact that immunocompromised persons with hypogammaglobulinaemia or isolated IgA deficiency may develop chronic cryptosporidiosis.

**Epidemiology**

*Cryptosporidium* is an ubiquitous parasite and infections have been traced to household or sick contacts, agricultural exposure, and contaminated water and food. The ease with which humans become infected is illustrated by attack rates of up to 70% in adult contacts of infected children in day-care centres. Widespread previous exposure or infection has been documented by several sero-epidemiological surveys revealing high prevalence of specific anti-cryptosporidial IgG in all paediatric and adult populations studied thus far.

Attempts to control environmental sources of infection have been thwarted by the resilient oocysts which can withstand extremes of temperature as well as standard disinfectants and chlorine. Particularly worrisome is the inability of water purification systems to eliminate *Cryptosporidium*. The oocyst is only 4 μm in diameter, making it difficult to clear through filtration. Community-wide outbreaks, such as the one in Milwaukee, US, which affected more than 400 000 people, have been traced to the public water supply.

**Clinical manifestations**

Acute infection is characterised by watery diarrhoea with 6–12 bowel movements/day, crampy epigastric pain, weight loss, anorexia, malaise and flatulence. Such symptoms are typical of diseases of the small bowel, the primary site of cryptosporidial infection. Nausea, vomiting and fever may also be present. A small subset of patients may present with colitis but primary...
Epidemiology

- ubiquitous parasite found worldwide
- common cause of diarrhoea
- prevalence in AIDS: 5% (US and Europe), 30–50% (tropical regions)
- infection 5–22 days after ingestion of as few as 30 oocytes
- sources of oocytes include sick contacts, animals, soil, contaminated food and water

Box 2

Diagnosis

- demonstration of oocytes in fecal or intestinal biopsy specimens by special stains or immunological assays
- commonly used stains include acid fast (for fecal matter) and haematoxylin and eosin (for intestinal biopsy material)
- if clinical suspicion is high and initial stool exam is negative, repeat stool studies given variation in oocyte shedding in serial stool specimens

Box 3

Clinical manifestations

- in normal hosts and HIV-infected persons with CD4 > 200, infection usually leads to diarrhoeal syndrome with spontaneous resolution
- in HIV-infected persons with CD4 < 100, infection most commonly becomes chronic and leads to an enteritis and, less commonly, cholangitis
- voluminous, non-bloody diarrhoea (6–20 bowel movements/day)
- other signs/symptoms may include: nausea, vomiting, crampy abdominal pain, right upper quadrant pain, fever, jaundice, wasting/weight loss

Box 4

Treatment

- oral or intravenous fluid repletion or maintenance
- nutritional supplements: oral, enteral, parenteral
- antimotility agents

In persistent infection:
- optimize antiretroviral therapy
- trial of paromomycin (symptomatic improvement in 50% of patients but rarely eradicates parasite)

Box 5

involvement of the large bowel is rare. Though the severity of symptoms may vary from individual to individual, a consistent finding is the spontaneous resolution of diarrhoea in the immunocompetent and well-nourished individual, usually within two weeks.

Two clinical patterns exist for HIV-related cryptosporidiosis: those persons with relatively preserved immunological function are able to clear infection, whereas those with depressed CD4 counts (ie, ≤180 cells/mm³) are usually chronically infected. Among persons with CD4 counts < 100 cells/mm³, a few individuals (< 10%) may have spontaneous resolution but the remainder have life-long diarrhoea of a waxing and waning nature, malabsorption and wasting. The chronicity and severity of the diarrhoea are usually what prompt HIV-infected individuals to seek medical treatment. In two clinical studies involving patients in the US and Mexico, clinical presentations were remarkably similar, with duration of symptoms averaging 9–12 weeks and weight loss at presentation averaging 14 kg.17,18

A particularly worrisome complication seen in over 10–15% of HIV-infected persons with cryptosporidiosis is infestation of the biliary tree leading to sclerosing or acalculous cholangitis.2 Clinical manifestations may include fever, right upper quadrant pain, jaundice, nausea and vomiting, with or without diarrhoea. Diagnostic work-up should include liver enzymes (alkaline phosphatase may be elevated up to 25 times the upper limit), noninvasive (ultrasound) and occasionally invasive (cholangiography) imaging.19,20 Persistence of the organism in the biliary tract may lead to seeding of the intestine and prolongation of diarrhoea. Though rarer, other clinical manifestations include pancreatitis, probably due to infection of the pancreatic duct and hepatitis.

Diagnosis

Cryptosporidiosis is diagnosed by identifying oocytes in stool in normal hosts and HIV-infected persons with cryptosporidiosis. Though many stool identification techniques exist, the most widely used method is the acid-fast stain (cold Kinyoun modified, hot Kinyoun, Ziehl-Neelsen). Oocytes stain red with varying intensity allowing differentiation from similarly sized and shaped yeast which are not acid fast. Other studies, such as the direct immunofluorescent assay which utilises a monoclonal antibody, may increase the sensitivity of stool identification. If initial studies are negative, the optimum number of stool specimens for diagnosis is unknown. Serial stool specimens in HIV-infected persons reveal 10-fold variation in cryptosporidial oocyte shedding.21 The importance of obtaining multiple specimens (eg, at least three specimens on three separate days) is supported by studies on other protozoa that demonstrate increased yield with additional stool samples.22

If stool studies are negative, endoscopy should be considered to establish a definitive diagnosis when diarrhoea is refractory to antimotility agents or when the disease process changes (eg, development of colitic symptoms, bloody diarrhoea, right upper quadrant pain). Endoscopic biopsy specimens may reveal Cryptosporidium when stool studies have been negative in < 10% of patients. Other pathogens such as cytomegalovirus or Microsporidia may also be identified.18 When examining intestinal biopsy material, Cryptosporidia can be identified on the brush border of the mucosal epithelial surface and appear basophilic with haematoxylin and eosin staining. Failure to identify Cryptosporidium in intestinal biopsy material does not rule out diagnosis, as cryptosporidial distribution may be patchy.23 Other identification methods include concentration techniques such as Sheather’s sugar flotation.

Treatment

NUTRITIONAL AND FLUID SUPPORT

Cryptosporidial diarrhoea is voluminous and individuals may lose up to 17 l/day in stool. Fluid loss is especially prominent in debilitated individuals who are unable to drink. Utmost attention needs to be paid to correction of electrolyte abnormalities and prevent dehydration with intravenous or oral rehydration solutions (ORS). A simple home-made ORS is 200 ml of water with an eighth of a teaspoon of salt and two teaspoons of sugar. Individuals should be encouraged to start ORS early to prevent even initial dehydration.

Optimum nutritional support should include a trial of oral nutritional supplements before considering total parenteral or intravenous hyperalimentation. Nutritional supplements containing medium chain fatty acids may be better absorbed in patients with small intestinal injury and malabsorption. Milk and dairy products should be avoided since lactose intolerance is common.
SPECIFIC ANTICYTOSPORIDIAL DRUGS
Though more than 100 drugs have been tried, none has proven uniformly effective in treating cryptosporidiosis.25 Paromomycin, a nonabsorbable aminoglycoside, has been shown to improve symptoms in approximately 50% of patients but complete resolution of diarrhea with eradication of parasite occurs in only 10% of individuals.17 Other agents that have been tried with unencouraging results include the macrolide antibiotics spiramycin and azithromycin.25

Another therapeutic approach has been the use of immune-based therapies such as hyperimmune bovine colostrum and bovine transfer factor. These agents are produced by inoculating cows with cryptosporidial oocysts and isolating specific anticytosporian antibody or transfer factor from the cows’ milk or lymph nodes, respectively. Trials have been limited by small numbers of patients, cumbersome production techniques and difficulty in standardising different batches of product.26,27

ANTIRETROVIRAL AGENTS
Although resolution of cytosporian infection with zidovudine has been documented in case report format, no studies have shown a definitive benefit of therapy in cryptosporidiosis.28 At present, combination antiretroviral therapy including protease inhibitors is recommended by the authors in the hope that the ensuing boost in immune function will help combat cytosporian infection.

ANTIMOTILITY AGENTS
Antimotility agents should be used for symptomatic relief and include first-line agents such as loperamide and diphenoxylate/atropine (lomotil). A stronger agent, tincture of opium, can be used for refractory diarrhea with the warning that over-use may lead to hypomotility, constipation and nausea. Antimotility agents should be stopped immediately if bloody diarrhea or fever develops and should not be restarted until the presence of invasive organisms has been ruled out. Trials of other antimotility agents such as octreotide, a somatostatin analogue, have shown them to be more effective than standard agents in controlling diarrhoeal symptoms.29

Conclusions
Cryptosporidium is one of multiple emerging pathogens that poses considerable challenges to medical science. As populations of immunocompromised persons grow, it will cause increasing morbidity due to chronic severe enteritis. Innovative methods to treat this infection are urgently needed. Immunotherapy trials with orally administered Cryptosporidium polyclonal or monoclonal antibodies are ongoing. Novel pharmacologic approaches based on a better understanding of the biology of this protozoa may be effective. A successful vaccine to induce mucosal immunity would be a great advance.
Medical Anniversary

JOHN COAKLEY LETTSON, 22 November 1744

John Coakley Lettsom (1744–1815) was born in Tortola, British Virgin Islands, less than 160 km east of Puerto Rico. He was one of the last of seven pairs of twins, and he and his brother Edward were the only pair to survive infancy. Their father was a plantation owner so he was able to send his son to a Quaker school in Lancashire. A year as a dresser at St Thomas' Hospital was eventually followed by medical education at Edinburgh and Leiden and a London practice with Dr John Fothergill, London's most celebrated Quaker physician. He and his wife Ann Miers had eight children and lived in his country residence The Grove, Camberwell. He became FRCP (Ed) in 1771 and FRS in 1773. His immortality is linked with the Medical Society of London, which he founded in 1773. He also founded the Sea-bathing Infirmary, Margate. After a full and satisfying life helping others, Lettsom died on 1 November 1815 and is buried at the Quaker Burial Ground, Bunhill Row, London.

He was blessed with a sense of humour as revealed in this epigram against himself:

When any sick to me apply
I physics, bleeds and sweets 'em
If after that they choose to die,
Why verily!

— DG James

Photo courtesy of The Medical Society of London
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