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What is This?
TOXOPLASMA GONDII INFECTION IN CATS
ABCD guidelines on prevention and management

Katrin Hartmann, Diane Addie, Sándor Belák, Corine Boucrat-Baralon, Herman Egberink, Tadeusz Frymus, Tim Gruffydd-Jones, Margaret J Hosie, Albert Lloret, Hans Lutz, Fulvio Marsilio, Karin Mósti, Maria Grazia Pennisi, Alan D Radford, Etienne Thiry, Uwe Truyen and Marian C Horzinek

Agent properties

Toxoplasma gondii is an obligate intracellular coccidian parasite that can infect virtually all species of warm-blooded animals, including people. Domestic cats and other felids are the natural hosts – non-feline species serve only as intermediates.1,2

Three infectious structures can be distinguished: sporozoites in oocysts, tachyzoites (the actively multiplying stage), and bradyzoites (the slowly multiplying stage) enclosed in tissue cysts. Oocysts are excreted in faeces, whereas tachyzoites and bradyzoites are found in tissues and milk.1,2

Pathogenesis

Enterop epithelial life cycle

This cycle is found only in the feline host. Most cats are infected by ingesting intermediate hosts – typically rodents – infected with tissue cysts. Bradyzoites are released in the stomach and intestine from the tissue cysts when digestive enzymes dissolve the cyst wall. They enter epithelial cells of the small intestine and give rise to schizonts, initiate five types of predetermined asexual stages, and merozoites released from the schizonts eventually form male and female gamonts. After fertilisation, a wall is formed around the fertilised macrogamont to form an oocyst. Oocysts are round to oval, 10 x 12 μm in size, and are still unsporulated (not infectious) when passed in faeces. After exposure to air and moisture for 1–5 days they sporulate to contain two sporocysts, each with four sporozoites.1,2

Overview: Toxoplasma gondii infection is common in cats, but the clinical disease is rare. Up to 50% of cats, especially free-roaming ones, have antibodies indicating infection and the presence of cystic stages.

Disease signs: Clinical signs only appear in few cats when they become immunosuppressed – in these situations cystic stages can be reactivated. Commonly affected are the central nervous system (CNS), muscles, lungs and eyes.

Human infection: Cats can pose a risk for humans when they shed oocysts. However, this happens only once in their lifetime, usually only for 3–10 days after ingestion of tissue cysts. Thus, cats that have antibodies to T gondii no longer shed oocysts, and do not pose a risk to humans.
The cycle is usually completed within 3–10 days of ingestion of tissue cysts, which is the route of infection in up to 97% of naive cats. In the rare event that cats ingest oocysts or tachyzoites, formation of new oocysts is delayed and shedding can occur for up to 18 days (occasionally longer). However, only 20% of cats fed oocysts will shed.1,2

Extraintestinal life cycle
The extraintestinal development of *T. gondii* is the same for all hosts, including cats, dogs, and people, irrespective of whether tissue cysts or oocysts have been ingested. After the ingestion of oocysts, sporozoites hatch in the lumen of the small intestine and enter intestinal cells, including those in the lamina propria. Sporozoites divide into two by an asexual process known as endodyogeny, thereby becoming tachyzoites. These are lunate (falciform) in shape, approximately 6 x 2 μm, and multiply in almost any cell of the body. When the cell ruptures, releasing the tachyzoites, these infect new cells. Otherwise, tachyzoites multiply intracellularly for an undetermined period, and eventually encyst. Tissue cysts vary in size from 15–60 μm and usually conform to the shape of the parasitised cell. Tissue cysts are formed mainly in the CNS, muscles and visceral organs, and probably persist for the life of the host. They can be reactivated after immunosuppression, which may then lead to clinical signs.1,2

Parasitaemia during pregnancy of the host can cause placentitis and spread of tachyzoites to the fetus. Many kittens born to queens infected with *T. gondii* during gestation become infected transplacentally or when suckling. Clinical signs are common in these kittens, varying with the stage of gestation at the time of infection; some of these newborn kittens shed oocysts.1,2

Epidemiology
Antibody prevalence to *T. gondii* varies geographically: in Portugal, 24% of cats had antibodies in one study; in the USA, 16–40% were antibody-positive, depending on the state.2 However, only 3/326 faecal samples from cats in California and 1/252 in Switzerland contained *T. gondii* oocysts.4 The annual burden in the environment is about 90–5000 oocysts per square metre.5 The age of the cat does not play a role in the frequency of *T. gondii* shedding, but the season does: shedding is more common between July and December in the northern hemisphere.6

The three major modes of transmission of *T. gondii* in all host species are congenital infection, ingestion of infected tissue, and ingestion of oocyst-contaminated food or water.2

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Clinical signs
Clinical signs develop very rarely in infected cats and are caused by inflammation and tissue necrosis resulting from intracellular growth of tachyzoites.2 Congenital infection tends to be more serious than infection of the adult cat.2 Clinical toxoplasmosis develops during dissemination and intracellular replication of tachyzoites. It usually originates from reactivation of a latent infection rather than after a newly acquired infection. If a carrier cat is immunosuppressed, bradyzoites in tissue cysts replicate rapidly and disseminate again as tachyzoites. Clinical toxoplasmosis has been documented in some cats infected with feline immunodeficiency virus (FIV) or feline leukaemia virus (FeLV) [EBM grade III].16 Commonly used doses of glucocorticoids do...
not predispose to reactivation of \( T\) \textit{gondii} cysts.\textsuperscript{17} However, administration of ciclosporin to cats with renal transplants or dermatological disease has been associated with clinical manifestations [EBM grade IV].\textsuperscript{18-20}

The most commonly affected tissues are the CNS, muscles (Figure 1), lungs (Figure 2) and eyes. Hepatic and pancreatic involvement is less likely. Cats with toxoplasmosis show neurological signs (eg, seizures, ataxia), muscle hyperaesthesia, dyspnoea, uveitis, icterus, diarrhoea, fever, depression, anorexia and weight loss.\textsuperscript{2} Transplacentally or lactogenically infected kittens develop more severe signs and frequently die of pulmonary or hepatic disease [EBM grade III].\textsuperscript{21} Immune complex formation\textsuperscript{22} and deposition in tissues, as well as delayed hypersensitivity reactions, can be involved in chronic forms of toxoplasmosis. Since \( T\) \textit{gondii} is not cleared from the body, either naturally or through treatment, toxoplasmosis can recur.

**Immunity**

Immunity to \( T\) \textit{gondii} in the cat is poorly understood. In the mouse and in humans, it is highly dependent on cell-mediated effector responses.\textsuperscript{23} All infected cats develop IgG and IgA antibodies; about 80\% also have IgM antibodies. IgG can take 4–6 weeks to appear, and maximal antibody titres are achieved within 2–3 weeks of first appearance.\textsuperscript{2}

**Diagnosis**

Oocyst shedding is diagnosed by microscopy of faecal samples. Diagnosis of the disease is only confirmed when the organism is found in body fluids or tissue. If suitable samples cannot be taken, a tentative diagnosis is sometimes based on rising IgM titres, exclusion of other causes for the clinical signs, and a favourable clinical response to anti-\( T\) \textit{gondii} drugs [EBM grade II].\textsuperscript{1,2}
Detection of oocysts in faeces

*T. gondii* oocysts are 10 μm in size and best demonstrated by centrifugation using Sheather’s sugar solution (saccharose solution with a specific gravity of 1.27 g/ml) during the shedding period. *T. gondii* oocysts are morphologically indistinguishable from those of *Hammondia hammondi*, *Besnoitia oryctofelisi* and *Besnoitia darlingi*. A cesium chloride method for easy purification of *T. gondii* oocysts from faeces of infected cats has been described.\(^2\)

Detection of tachyzoites

Ante-mortem diagnosis of clinical toxoplasmosis ideally is based on the detection of the organism by cytology or polymerase chain reaction (PCR) [EBM grade III]. Tachyzoites can be detected in various tissues and body fluids during acute illness (Figure 3). They are rarely found in blood, but occasionally in cerebrospinal fluid (CSF) or aqueous humour, fine-needle aspirates of organs (eg, lymph nodes), and transtracheal or bronchoalveolar lavage fluid. Detection of tachyzoites results in a definitive diagnosis. Alternatively, a PCR can be performed using CSF, aqueous humour or bronchoalveolar lavage fluid.

Detection of antibodies

Using the immunofluorescence assay (IFA), antibodies of the IgM, IgG and IgA isotypes can be detected. For assessing human health risks, antibody test results from healthy cats are useful. An antibody-negative cat can be shedding oocysts (early during infection, before antibodies have had time to develop) and will likely shed oocysts if exposed for the first time. An antibody-positive cat does not shed oocysts: antibodies need 2–3 weeks to develop, by which time cats usually no longer shed; and a cat sheds only once in its lifetime. It is also unlikely to shed oocysts if re-exposed or immunosuppressed [EBM grade II].\(^1\)

Antibodies are commonly found in both healthy and sick cats. Thus, their presence does not prove clinical toxoplasmosis. Antibodies of the IgM class are also commonly detected in healthy cats and do not correlate with clinical signs.

### Treatment

Clindamycin is the treatment of choice\(^24\) and should be administered at 10–12 mg/kg orally q12h for 4 weeks (Table 1) [EBM grade III]. Cats with systemic disease and uveitis should be treated with clindamycin in combination with topical, oral or parenteral glucocorticoids, to avoid secondary glaucoma and lens luxation [EBM grade III].\(^25\) Prednisolone acetate (1% solution) applied topically to the eye three to four times daily is generally sufficient.

Clinical signs not involving the eyes or the CNS usually begin to resolve within the first 2–3 days of clindamycin administration. CNS and ocular toxoplasmosis tend to respond more slowly. In cases of pulmonary toxoplasmosis, radiographic abnormalities might not resolve for several weeks. The prognosis is usually poor in pulmonary or hepatic disease, particularly in immunocompromised animals.\(^26\)

### Prevention of infection

Preventing toxoplasmosis in cats involves measures intended to reduce the incidence of infections and the shedding of oocysts into the environment. Cats should preferably be fed commercially available, processed food. Prevalence of feline *T. gondii* infection is higher in countries where raw meat is fed. Freezing or irradiation can kill tissue cysts without affecting meat quality. Pets should be prevented from hunting and eating intermediate hosts (rodents) or mechanical vectors, such as cockroaches and earthworms. If meat is fed, it should be thoroughly cooked, even if frozen. Cats should be prevented from entering buildings where food-producing animals are housed or where feed storage areas are located.\(^1\)

**Table 1** Treatment of toxoplasmosis

<table>
<thead>
<tr>
<th>Drug</th>
<th>Comment</th>
<th>ABCD recommendation</th>
<th>EBM grade</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antiparasitic therapy</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clindamycin</td>
<td>10–12 mg/kg PO q12h for 4 weeks (treatment of choice)</td>
<td>III</td>
<td></td>
</tr>
<tr>
<td><strong>Symptomatic therapy</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prednisolone acetate (1%)</td>
<td>For cats with Toxoplasma-induced uveitis (to avoid secondary glaucoma and lens luxation)</td>
<td>Use in addition to systemic antibiotic treatment. Apply topically to the eye q6–8h</td>
<td>IV</td>
</tr>
</tbody>
</table>

Antibodies are commonly found in both healthy and sick cats. Thus, their presence does not prove clinical toxoplasmosis.
The risk of infection from cats is low, except for young children and people with HIV infection. Although toxoplasmosis is more common in HIV-infected persons, the disease results from reactivation of a previous infection rather than from acquiring a new infection.

Most people are infected with *T. gondii* through ingestion of undercooked meat, especially goat, mutton and pork. The risk of infection from cats is low, except for young children playing in soil contaminated with sporulated oocysts.

Bites or scratches from an infected cat do not transmit the infection.

Infected cats under treatment with immunosuppressive drugs at standard doses do not start shedding oocysts in their faeces.

Infected cats also do not re-shed oocysts in their faeces when they become immunosuppressed due to infection with FIV or FeLV. Cats infected with FIV or FeLV that are subsequently infected with *T. gondii* do not shed oocysts for any longer or in any greater numbers than other cats.

Newly identified strains of *T. gondii* are highly infectious for species other than cats; thus, cats might actually become less important in the spread of this infection.
**KEY POINTS**

- *T. gondii* infection is common in cats. The clinical picture of toxoplasmosis, however, is rare.
- Up to 50% of cats have antibodies, which show they are infected and harbour *T. gondii* cysts.
- Clinical signs usually occur when cats become immunosuppressed, due to the reactivation of cystic stages.
- Commonly affected sites are the CNS, muscles, lungs and eyes.
- Cats shedding oocysts can pose a risk to humans. However, they shed only once in their lifetime, and usually for only 3–10 days after ingestion of tissue cysts.
- Cats with *T. gondii* antibodies no longer shed oocysts; they neither are nor will become a zoonotic risk.
- A definitive diagnosis of toxoplasmosis relies on the detection of the organism in body fluids or tissues.
- The treatment of choice for cats with the disease toxoplasmosis is clindamycin.

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**Conflict of interest**

The authors do not have any potential conflicts of interest to declare.

**References**


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