Diagnosis of feline infectious peritonitis: update on evidence supporting available tests

Author Details
Séverine Tasker
Professor of Feline Medicine
BSc BVSc PhD DSAM DipECVIM-CA PGCertHE FHEA MRCVS
The Feline Centre, Langford Vets, Bristol Veterinary School, University of Bristol, Bristol, BS40 5DU, UK
s.tasker@bristol.ac.uk

Photo will be provided.

Structured Abstract

Practical relevance
Feline coronavirus (FCoV) infection is very common in cats, usually only causing mild intestinal signs such as diarrhoea. Up to 10% of feline coronavirus infections, however, result in feline infectious peritonitis (FIP), a fatal disease which is a common cause of death in young cats.

Clinical challenges
Obtaining a definitive diagnosis of FIP based on non-invasive approaches in cats is difficult. Confirmation of FIP relies on finding appropriate cytological or histopathological changes in cytological samples or biopsies in association with positive immunostaining for FCoV antigen. In FIP cases with effusions, cytology and immunostaining on effusion samples can be relatively easy to perform. In the absence of effusions, obtaining diagnostic samples more challenging. Often, and especially in FIP cases without effusions, collection of biopsies from tissues with gross lesions is necessary.

In the absence of a definitive diagnosis, a high index of suspicion of FIP may be obtained from a combination of the cat’s signalment (e.g. being 2 years or younger, originating from a multi-cat household), history (e.g. fluctuating pyrexia), clinical examination findings (e.g. pyrexia, jaundice, effusions, uveitis, neurological signs) and laboratory test results (e.g. lymphopenia, hyperglobulinaemia, reduced albumin to globulin ratio and/or elevated α1-acid glycoprotein concentrations in serum or effusion, positive reverse transcriptase-polymerase chain reaction (RT-PCR) for FCoV RNA in effusion, cerebrospinal fluid or biopsy samples, pyogranulomatous changes identified in cytological or biopsy specimens). These results, if largely consistent with FIP, can be used as a basis of discussion with the owner about whether additional, more invasive, diagnostic tests are warranted. In some cases it may be that euthanasia is discussed as an alternative to pursuing a definitive diagnosis ante-mortem, especially if financial limitations exist or when cats are very sick and concerns exist over a patient’s ability to tolerate invasive diagnostic procedures (e.g. surgical biopsy). Ideally confirmation of the diagnosis should be made in such patients, if euthanased, from samples taken at post-mortem examination.

Global importance
Feline infectious peritonitis occurs wherever FCoV infection is present in cats, and thus is found in most parts of the world. Only a few areas have been investigated (e.g. Galápagos Islands and Falkland Islands) which appear to be free from FCoV infection.

Evidence base
This review provides a comprehensive overview of how to approach the diagnosis of FIP, focusing on tests available to the veterinary practitioner and recent published evidence for the usefulness of these tests.
What are coronaviruses?
Coronaviruses are large, enveloped, positive-sense single-stranded RNA viruses with non-segmented genomes of around 30,000 nucleotides in length(Siddell, 1995 #371). Feline coronavirus (FCoV) is a sub-species of the alphacoronavirus I species, along with canine coronavirus (CCoV). Coronaviruses exhibit a high rate of mutation during RNA replication and therefore exist as clusters of genetically diverse populations, known as quasispecies(Denison, 2011 #372;Desmarests, 2016 #347). This genetic diversity, along with a readiness to recombine with other coronavirus strains, is associated with their pathogenicity and cross-species transmission. Interestingly, FCoVs appear to have emerged in the 1950s, possibly due to cross-species transmission(Addie, 2012 #373). Cats worldwide have been found to be infected with FCoV with the exception of those residing in a few areas, such as the Galápagos Islands(Levy, 2008 #387) and Falkland Islands(Addie, 2012 #259), likely due to quarantine of those cats by their isolated island habitat.

Two FCoV so-called serotypes are recognised: Type 1, which represents the vast majority of field strains found in naturally infected cats, although geographical variation exists(Benetka, 2004 #2;Hohdatsu, 1992 #52), and Type 2, which arose following recombination events between Type 1 FCoV and CCoV(Herrewegh, 1998 #377). The two FCoV serotypes are distinguished primarily by the genetic and serological differences in their transmembrane spike (S) gene and protein, respectively. The S protein (Figure 1) is important as it is the part of the FCoV that binds to the host (feline) receptor, mediating host cell entry(Jaimes, 2018 #409). The Type 2 FCoV S protein resembles that of CCoV, and, like CCoV, Type 2 FCoVs use aminopeptidase N as their feline host cell receptor. The receptor for Type 1 FCoVs is unknown(Dye, 2007 #323).

The complex relationship between feline coronavirus and feline infectious peritonitis (FIP)
Feline coronavirus infection is very common in cats. Around 40% of the domestic cat population has been infected with FCoV, and this figure increases to 90% in multi-cat households(Addie, 2000 #209;Addie, 1992 #111). Modern changes in feline husbandry (e.g. more cats kept indoors and in multi-cat households) are likely to have led to the increase in FCoV-related disease(Addie, 2012 #373).

Natural infections with FCoV are transient in ~70% of cats, whereas persistent infections occur in ~13% of cats(Addie, 2012 #373). These persistently infected cats are sometimes referred to as ‘carrier’ or ‘chronically shedding’ cats. In most cases, FCoV infection is asymptomatic or results in only mild gastrointestinal clinical signs (e.g. inappetence, diarrhea, vomiting), although occasionally more severe gastrointestinal disease is seen. Interestingly, around 5-10% of cats are believed to be resistant to FCoV infection(Addie, 2012 #373). However, in a small percentage of cases, FCoV infection results in the severe disease of FIP(Pedersen, 2009 #204;Addie, 1995 #40). Occasionally, and possibly with increasing frequency recently, outbreaks of FIP (when a larger percentage of cats are affected) in multi-cat households or shelters are reported(Wang, 2013 #269;Barker, 2013 #280). No curative treatments for FIP are currently available, although some novel treatments, including protease inhibitors, show promise(Kim, 2016 #360;Legendre, 2017 #374;Pedersen, 2017 #403).

Asymptomatic FCoV infection was previously believed to be confined to the intestinal tract, but we now know that healthy FCoV-infected cats can have systemic FCoV infection, albeit with lower FCoV viral loads than cats with FIP(Kipar, 2006 #130;Kipar, 2010 #227;Meli, 2004 #76;Desmarests, 2016 #347).

What factors contribute to the development of FIP following FCoV infection?
Viral factors are important in the pathogenesis of FIP. As outlined above, the S protein of FCoV mediates host cell entry. The S protein contains a putative fusion peptide that enables fusion of the FCoV envelope with the host cell membrane(Bosch, 2003 #351). Mutations in the S gene can result in amino acid substitutions in the transcribed S protein that influence the tropism of FCoV(Belouzard, 2012 #352). Studies have identified mutations in the fusion peptide sequence of the FCoV S gene that were thought to be markers of FIP(Chang, 2012 #255;Bank-Wolf, 2014 #395) as well as changes in the closely-related furin cleavage site that were also
thought to be correlated with FIP [Licitra, 2013 #277]. Recently it has been found that the fusion peptide mutations are likely to be markers of systemic FCoV infection, which can occur in both FIP and non-FIP cats, rather than FIP per se [Porter, 2014 #321]. However, these mutations are still important as it is probably via these, and/or other mutations, that the FCoV acquires its monocyte/macrophage tropism to allow it to spread systemically, outside of the intestinal tract, and contribute to the development of FIP. Other viral factors then mediating effective and sustained replication in monocytes, and activation of infected monocytes, are also likely to be important for the subsequent development of FIP following systemic FCoV infection [Kipar, 2014 #309].

**Host factors** are also very likely to play an important part in FIP development. These include the host immune response (e.g. T-lymphocyte depletion occurs in cats that develop FIP), the ability of monocytes to sustain FCoV replication, breed and genetics [de Groot-Mijnes, 2005 #314; Golovko, 2013 #294; Pedersen, 2016 #356; Dewerchin, 2005 #394].

**Environmental factors**, such as the level of stress and overcrowding in a household, may also play a role. These may feed into increasing the rate of viral replication in monocytes, and contribute to the development of FIP. These include factors such as the level of stress and overcrowding in a household, which contributing to the development of FIP.

**Approaching a diagnosis of FIP**

**Making a definitive diagnosis of FIP**

A definitive diagnosis of FIP traditionally relies on histopathological examination of tissues, usually with detection of the virus within lesions by immunohistochemistry for FCoV antigen. Immunostaining of FCoV antigen in effusion samples is also an option for definitive diagnosis of FIP in cases with effusions showing biochemical and cytological features consistent with FIP.

Histopathological examination, effusion analysis and FCoV antigen immunostaining are discussed in more detail later.

**Obtaining a high index of suspicion of a diagnosis of FIP**

In the absence of a definitive diagnosis, a high index of suspicion of FIP may be obtained from background information, clinical signs and routine clinicopathological results. With experience, this can be used as a basis to discuss with the owner whether additional, more invasive, diagnostic tests (e.g. biopsy of affected tissues) are warranted. In such cases it may be that euthanasia is discussed as an alternative to pursuing a definitive diagnosis ante-mortem, and this may be preferable in, for example, shelter cats, where there are financial limitations, or when cats are very sick and concerns exist over a patient’s ability to tolerate diagnostic procedures (e.g. surgical biopsy). If euthanasia is performed without a definitive diagnosis, post-mortem examination is strongly advised and relatively simple to perform [Tasker, 2017 #380]. This permits both assessment for gross changes consistent with FIP (Figure 2), and sampling for histopathological examination. If financial limitations preclude the latter, it is worth contacting researchers with an interest in FIP (e.g. the University of Bristol Feline Coronavirus Research Group & Bristol-Zurich FIP Consortium) to see if samples are being sought for research studies that could allow for analysis at a reduced cost or free of charge.

Many important differential diagnoses should be considered in cats suspected of having FIP such as toxoplasmosis, mycobacterial infection and lymphocytic cholangitis. These, and others, are described in Table 1, together with consideration of features distinguishing those diseases from FIP.

**Signalment and background evidence for FIP**

It should be remembered that FIP is most common in young cats (those less than three years of age, and especially those less than two years of age [Riemer, 2016 #358]) but a smaller peak of cases is seen in cats older than 10 years of age. Male cats are also at a slightly higher risk [Riemer, 2016 #358]. Some breeds in some countries may be predisposed to FIP [Pesteanau-Somogyi, 2006 #121; Worthing, 2012 #252], but this is likely due to the presence of unknown specific genetic risk factors in those breeds in those countries, and generalised
breed predispositions may not exist (Riemer, 2016 #358). A recent history of stress (e.g., adoption, being in a shelter, neutering, upper respiratory tract disease, vaccination etc.) may be apparent (Riemer, 2016 #358) and may play a part in triggering the development of FIP in a FCoV-infected cat. Although living in a multi-cat household increases the likelihood of being FCoV seropositive, a recent large study (Riemer, 2016 #358) found that the majority of cats presented to a university hospital with FIP were from households containing a small number of cats at time of diagnosis.

**Clinical signs of FIP**

Disease manifestations of FIP typically comprise a vasculopathy resulting in (‘wet’) effusions (up to 80% of FIP cases have effusions), granuloma formation only resulting in (‘dry’) mass lesions, or a combination of the two; indeed most FIP cases with effusions also have granulomatous lesions visible at post-mortem examination. Clinical signs (Figure 3) seen in both effusive and non-effusive FIP include lethargy, anorexia, weight loss (or failure to gain weight/stunted growth in younger cats), a fluctuating pyrexia that is usually non-responsive to drugs such as antibiotics or non-steroidal anti-inflammatories, and sometimes jaundice (more common in effusive FIP). A recent study evaluating referral cat cases with a history of pyrexia found that, at 20.8% (22 of 106 cats), FIP was the most common diagnosis made, showing the importance of FIP as a differential diagnosis in pyrexic cats in this population of referred cats (Spencer, 2017 #378). Another study (Riemer, 2016 #358) describing FIP cases only reported that body temperature exceeded 39.5°C in 81% of cats and 40°C in 39% of cats. Pyrexia was far more common in cats with effusive FIP than those with neurological non-effusive FIP (Riemer, 2016 #358). Lymphadenomegaly can also be present in both the effusive and non-effusive forms of FIP.

Effusive ‘wet’ FIP is associated with abdominal, pleural and/or pericardial effusions (occasionally in the scrotum too of entire male cats), and is often quite acute in nature, progressing within a few days or weeks and severely limiting survival (Ritz, 2007 #117). These cats can present with dyspnoea, tachypnoea and/or abdominal distension. Non-effusive ‘dry’ FIP is typically associated with neurological (can be focal, multifocal or diffuse in nature, often with central vestibular signs, occasionally as a T3-L3 myelopathy (Crawford, 2017 #397); Figure 3) and/or ocular (anterior and/or posterior uveitis; Figure 3) signs and is more chronic, progressing over a few weeks to months. Dermatological signs (manifested typically as small multiple non-pruritic papules or nodules) have also been reported in dry FIP. Renomegaly may occur in non-effusive FIP with renal involvement. Occasionally a diffuse pyogranulomatous pneumonia is reported.

It is important to remember that clinical signs of FIP can change over time so repeated clinical examinations are important to detect newly apparent signs (e.g. development of a small volume of effusion, ocular changes visible on retinal examination).

Focal non-effusive FIP occasionally occurs, presenting typically as a palpable abdominal mass, and can be particularly challenging to diagnose as the lesions can be hard to initially differentiate from neoplasia and mycobacterial infection. Focal FIP case reports have comprised cats presenting with mesenteric lymph node enlargement due to necrogranulomatous lymphadenitis (Kipar, 1999 #23), or solitary mural intestinal lesions of the colon or ileoceccolic junction with associated regional lymphadenopathy (Harvey, 1996 #36). The cats with focal intestinal FIP had a history of vomiting and diarrhoea.

**Possible routine laboratory test findings in FIP**

**Routine haematology**

Haematological changes in FIP are non-specific but there are a number of abnormalities that can be looked for to support a diagnosis. Lymphopenia is particularly common (55-77% of cases); although a recent study found only 49.5% of FIP cases to be lymphopenic (Riemer, 2016 #358), with neutrophilia (39-57%), a left shift, and mild-moderate normocytic, normochromic anaemia (37-54%) also reported (Tsai, 2011 #245; Sparks, 1991 #57; Norris, 2005 #246; Riemer, 2016 #358). An association between FIP and microcytosis (with or without
anaemia) was recently reported\cite{Riemer, 2016 #358}. Severe immune-mediated haemolytic anaemia (IMHA), with an associated regenerative anaemia, can occur with FIP\cite{Norris, 2005 #246}, but is uncommon.

**Serum biochemistry**

The changes in serum biochemistry in FIP cases are varied and often non-specific but there are a number of important abnormalities that should be looked for to support a diagnosis of FIP.

**Hyperglobulinaemia** is reported in 89% of cases, often with hypoalbuminaemia or low-normal serum albumin (seen in 64.5% of cases)\cite{Riemer, 2016 #358}. The presence of hypoalbuminaemia alongside hyperglobulinaemia means that hyperproteinaemia may not always occur; past reports documented hyperproteinaemia in up to 60% of cases, especially in dry FIP cases, but lower prevalences of 17.5% have been reported recently\cite{Riemer, 2016 #358}. The combination of hyperglobulinaemia and hypoalbuminaemia or low-normal albumin concentration also means that the albumin:globulin (A:G) ratio is low, and this parameter can be useful to evaluate how likely FIP is in an individual case. Reports of useful cut-off values for A:G ratios in the diagnosis of FIP vary, but it has been suggested that an A:G ratio of $<0.4$ makes FIP very likely, whilst an A:G ratio of $>0.8$ makes FIP very unlikely \cite{Tsai, 2011 #245; Sparkes, 1991 #57; Norris, 2005 #246}. Although these cut-off values are useful to consider, the author does not use a specific value but looks at the A:G ratio in conjunction with other diagnostic test results; the lower the value, the bigger the suspicion for FIP becomes, especially if other findings are consistent with a diagnosis of FIP. Interestingly, a study\cite{Jeffery, 2012 #264} in a population of cats with a low prevalence of FIP (akin to the situation that is usually encountered in veterinary practice) found that an A:G ratio of $>0.6$ was useful in ruling out FIP, but that lower ratios were not helpful in ruling in FIP. Additionally, the frequency and extent of hypoalbuminaemia, hyperglobulinaemia, low A:G ratio and serum protein electrophoresis (SPE) abnormalities reported in FIP cases have decreased recently\cite{Riemer, 2016 #358; Stranieri, 2017 #379}. With respect to SPE changes, in one study\cite{Stranieri, 2017 #379} cases diagnosed with FIP from 2013 to 2014 tended to have elevated α2-globulins rather than the elevated γ-globulins seen in cases from 2004 to 2009; this is possibly due to veterinarians diagnosing FIP earlier, meaning that cases have not progressed to show elevated γ-globulins. Polyclonal and monoclonal elevated γ-globulins have been reported with FIP\cite{Taylor, 2010 #388}, although polyclonal elevations are far more common.

**Hyperbilirubinaemia** occurs in 21-63% of FIP cases, and is especially seen with effusive FIP, often without marked elevations in alanine aminotransferase (ALT), alkaline phosphatase (ALP) or gamma-glutamyltransferase (GGT) enzyme activity (although these can be moderately elevated in FIP cases). Hyperbilirubinaemia due to IMHA is uncommonly reported with FIP\cite{Norris, 2005 #246}, and cats are often not severely anaemic. Thus the presence of hyperbilirubinaemia in the absence of elevated hepatic enzyme activities or severe anaemia should raise the index of suspicion of FIP (NB: sepsis and pancreatitis can also cause hyperbilirubinaemia in the absence of elevated hepatic enzyme activities [Table 1]). Hyperbilirubinaemia is more commonly identified in FIP cases as the FIP disease progresses; additionally any hyperbilirubinaemia present can worsen as the FIP disease progresses\cite{Tsai, 2011 #245}.

**Acute phase proteins** (APPs) are made in the liver in response to cytokines released from macrophages and monocytes (especially IL-1, IL-6 & TNF-α) in many inflammatory and non-inflammatory diseases. α1-acid glycoprotein (AGP) is an APP, and its measurement can be helpful in the diagnosis of FIP. Although AGP elevations ($>0.48$ mg/ml) per se are not specific for FIP, markedly elevated AGP levels ($>1.5$ mg/ml) are often seen in FIP cases, so the magnitude of the AGP increase may be helpful in aiding the diagnosis of FIP, with higher concentrations being more useful in raising the index of suspicion for FIP\cite{Giori, 2011 #242; Duthie, 1997 #244; Paltrinieri, 2007 #260; Hazuchova, 2016 #367}. Indeed, a study found that when the pretest probability of FIP was high (i.e. history and clinical findings being supportive of FIP), moderate serum AGP levels (1.5–2 mg/ml) could discriminate cats with FIP from cats without FIP, but only higher serum AGP levels ($>3$ mg/ml) could support a diagnosis of FIP in cats with a low pretest probability of disease (i.e. history and clinical findings not supportive of FIP)\cite{Paltrinieri, 2007 #260}. However, another, albeit very small, study of unusual cases of FIP...
actually found that modest AGP concentrations (>1.5 mg/ml) were still able to discriminate between FIP and non-FIP cases\cite{Giori, 2011 #242}.

**FCoV serology in FIP**

Serum FCoV antibody tests are usually enzyme-linked immunosorbent assays (ELISAs), indirect immunofluorescence antibody (IFA) tests or rapid immunomigration tests\cite{Addie, 2015 #331}. Most tests use CoV-infected swine or feline cells as a substrate and titres are read in distinct multiples of serum dilutions. A positive FCoV antibody test indicates that the cat has been infected with FCoV and has seroconverted (this takes 2-3 weeks from initial infection). Breed-related differences in median FCoV antibody titre have been detected, and may reflect differences in breed response to FCoV infection\cite{Bell, 2006 #125; Bell, 2006 #126}. Although FIP cats tend to have higher FCoV antibody titres than non-FIP cats, there is much overlap, with no difference between median FCoV antibody titres in healthy and suspected FIP cats, so the value in an individual cat to distinguish cats with FIP is very limited\cite{Bell, 2006 #126}. Many clinically healthy cats (especially those in multi-cat households) have positive, often very high, FCoV antibody titres, whilst ~10% of cats with FIP are seronegative (this could be due to the presence of virus in the sample binding antibody and rendering it unavailable to the serological test\cite{Meli, 2013 #271}), highlighting difficulties in interpretation. It may be that a negative FCoV antibody result in a suspected dry FIP case is more useful to rule out a diagnosis of FIP\cite{Addie, 2009 #206}; however, negative results have been reported in cases of neurological FIP\cite{Negrin, 2007 #392}. Clinicians vary as to whether they perform serology or not in suspected cases due to this issue, although a positive result certainly indicates exposure to FCoV.

**Analysis of effusion samples**

Analysis of any effusion sample in a suspected case of FIP is extremely helpful for diagnosis, so obtaining samples of effusions should always be prioritised in investigations of suspected cases. Ultrasonography is generally regarded as being more sensitive than radiography for the detection of small volumes of fluid in the thorax and abdomen, but this may depend on where pockets of fluid reside. Repeated ultrasonography to identify any small volume effusion is recommended and, similarly, ultrasonography can be used to guide sampling of small pockets of fluid.

FIP effusions (Figure 4) are usually clear, viscous/sticky, straw-yellow and protein-rich (thick eosinophilic proteinaceous backgrounds are often described on cytology), with a total protein concentration of >35 g/l (>50% globulins). Very occasionally chylous effusions are described. Typical FIP effusions have similar low A:G ratios (see above) and raised AGP concentrations to those in serum. A recent study found that effusion AGP concentrations (of >1.55 mg/ml) were most useful (sensitivity and specificity of 93%) in differentiating FIP and non-FIP cases compared with AGP levels in the serum or other APPs\cite{Hazuchova, 2016 #367}; however the ‘diagnosis’ of FIP in the cats in this study was not always confirmed by histopathology and immunostaining. FIP effusions are poorly cellular (usually <5 x10^9/l cells), and are typically pyogranulomatous in nature with macrophages, non-degenerate neutrophils and very few lymphocytes. The effusions are therefore often described as modified transudates based on cell counts (<5 x10^9/l cells) but as exudates based on protein concentrations (>35 g/l).

Rivalta’s test is a crude point-of-care assay that can be performed on an effusion sample to allow rapid differentiation of transudate from exudate. A positive result merely means that the effusion is an exudate and thus is not specific for FIP; positive results are reported in non-FIP cases e.g. bacterial/septic peritonitis and lymphoma\cite{Fischer, 2012 #263}. Although cytology can sometimes be successful in diagnosing bacterial/septic peritonitis and lymphoma, to help discriminate these from FIP, many vets may not be confident performing cytology in-house. To perform the Rivalta’s test, 8 ml of distilled water and one drop of 98% acetic acid (vinegar can be used) are mixed in a universal pot, and then one drop of effusion is carefully placed onto the surface of the liquid. A positive test is indicated by the drop staying attached to the surface of the liquid, retaining its shape with a connection to the surface (Figure 5), or floating slowly to the bottom. A negative test
is indicated by the drop disappearing and the solution remaining clear. Interpretation of results can be problematic due to subjectivity and difficulties in deciding whether a result is truly positive or negative (Fischer, 2013 #382).

One recent study measured FCoV antibody titres in effusion samples (Meli, 2013 #271) and found an inverse correlation between FCoV load and FCoV antibodies in some samples, possibly due to antibody being bound by FCoV and thus not available as a ligand in the serological test; thus making serology unreliable.

Serology for FCoV antibodies can also be performed on effusion samples, with very varied results (Meli, 2013 #271; Lorusso, 2017 #401), so the author does not perform this test in suspected cases of FIP.

Immunostaining for FCoV antigen and reverse-transcriptase polymerase chain reaction (RT-PCR) for FCoV RNA on effusion samples can also be performed (see later).

Miscellaneous diagnostic tests

In cases with neurological clinical signs, imaging of the brain by magnetic resonance imaging may be useful to demonstrate changes. For example, obstructive hydrocephalus, syringomyelia, foramen magnum herniation and marked contrast enhancement of the meninges, third ventricle, mesencephalic aqueduct and brainstem has been reported with FIP (Foley, 1998 #24; Penderis, 2009 #393; Crawford, 2017 #397). Cerebrospinal fluid (CSF) can be collected from neurological cases although care should be taken as the risk of brain herniation is significant. CSF may show elevated protein concentrations (>30 mg/dl [>0.3 g/l] cisternal samples, >46 mg/dl [>0.46 g/l] lumbar samples; occasionally FIP cats show marked elevations of >200 mg/dl [>2 g/l]) and an increased cell count (>8 cells/µl [>8 x 10⁶/l]) lumbar and cisternal samples; FIP cats can have cell counts of >1000 cells/µl [>1000 x 10⁶/l]), with the cell type being predominantly neutrophilic, mononuclear or mixed (Singh, 2005 #104; Crawford, 2017 #397). Some neurological cases of FIP have unremarkable CSF analysis results. Samples of CSF can also be submitted for RT-PCR for FCoV RNA and immunostaining for FCoV antigen (see below). Add notes on Ab measurement in CSF from {Soma, 2018 #408}{Foley, 1998 #24}

Reverse-transcriptase polymerase chain reaction for FCoV

Background information on FCoV RT-PCR

Reverse transcriptase-PCR assays are available to detect FCoV; however, they are not specific for FIP-associated FCoVs. FCoV RT-PCR assays amplify both cell-associated subgenomic mRNA (short lengths of transcriptional RNA produced when the FCoVs replicate), as well as cell-associated or virion-associated genomic RNA, with the relative abundance of each determined by the positioning of primers (i.e. where along the FCoV sequence the primers bind during PCR) (Barker, 2017 #383). As viral transcription starts at the 3’ end of the FCoV genome (Figure 6) there are more subgenomic mRNAs containing viral 3’ sequence than those containing viral 5’ sequence, hence quantitative assays (i.e. RT-qPCR) directed at the 5’ end of the genome (e.g. viral replicase complex genes) are less susceptible to viral load overestimation than those directed at the 3’ end of the genome (e.g. 7a/b non-structural protein genes).

Laboratories should be able to report the sensitivity and specificity of the RT-PCRs they are using to detect FCoV RNA, and the binding site of the primers they use can give some indication as to whether the assay will be prone to viral load overestimation (see above). As an RNA virus, FCoV shows a high rate of errors during replication and any viral mutations at the site of primer and/or probe binding can result in loss of PCR assay efficiency, and ultimately sensitivity. PCR conditions may be altered to tolerate such mutations, but this can result in a loss of specificity (Barker, 2017 #386).

Reporting of results for FCoV RNA RT-PCR can be rapid if the laboratory used has a fast turnaround time, although, once time taken to submit the sample to the laboratory is factored in, reporting of results can still take a few days. This is usually quicker than immunostaining on tissue samples and may be quicker than
immunostaining on effusion samples. However, it should be noted that immunostaining can provide a definitive diagnosis whereas RT-PCR does not. Recently a rapid molecular technique (loop mediated isothermal amplification) for detecting FCoV RNA in-house has been described (Stranieri, 2017 #370), although it suffered from poor sensitivity.

**Which samples can be tested for FCoV by RT-PCR?**

FCoV RT-PCR can be used to detect FCoV RNA in blood, effusion, tissue, CSF, or aqueous humour samples from suspected cases of FIP. Tissue samples should not be formalin fixed, as formalin degrades the target RNA and can decrease PCR sensitivity; indeed RNA is very sensitive to degradation and samples for research purposes are often collected into RNA preservation fluids. However, the need for special collection conditions of samples outlined above destined for routine diagnostic purposes by RT-PCR is unproven. The presence (particularly of high levels) of FCoV RNA in blood, effusion, tissue, CSF and/or aqueous humour samples can be highly supportive of a diagnosis of FIP, as outlined below, but cannot be regarded, in the author’s opinion, as delivering a definitive diagnosis.

FCoV RT-PCR can also be performed on faecal samples, but this is primarily used to identify cats that are shedding FCoV for the management of infection in a multi-cat household. Faecal FCoV RT-PCR is not used to aid in the diagnosis of FIP, but interestingly recent studies have found that cats with FIP are more likely to be shedding FCoV (Barker, 2017 #383), and have higher amounts of FCoV RNA as determined by RT-qPCR (Porter, 2014 #321), in their faeces than cats without FIP.

**FCoV RT-PCR on tissue samples**

Tissue samples from cats with FIP are significantly more likely to be FCoV RT-PCR positive (Barker, 2017 #383), and have significantly higher FCoV loads by RT-qPCR (Porter, 2014 #321), than tissues samples from non-FIP cats, although cats without FIP can still be positive for FCoV by RT-PCR. For example in a recent extensive study evaluating FCoV RT-PCR (Barker, 2017 #383), 90.4% of tissue samples from FIP cats were FCoV RT-qPCR positive compared to only 7.8% of tissue samples from non-FIP cats. Not surprisingly, in FIP cats, FCoV loads tend to correlate with histopathological findings suggestive of FIP (Pedersen, 2015 #326; Barker, 2017 #383). Thus, the presence of high (i.e. low threshold cycle values of around <30 using the PCR assay from our laboratory [http://www.langfordvets.co.uk/diagnostic-laboratories/diagnostic-laboratories/pcr-acarus/feline-coronavirus#overlay-context=dagnostic-laboratories/diagnostic-laboratories/pcr-acarus/feline-coronavirus]) levels of FCoV RNA in tissue samples appears to be highly supportive of a diagnosis of FIP. Selection of appropriate tissues to sample can be guided by clinical signs, imaging results, cytological findings (e.g. pyogranulomatous inflammation) etc., but non-invasive collection of such tissue can be difficult. Often laparotomy, or even laparoscopy, may be considered too invasive to perform in a sick cat in which the index of suspicion of FIP is very high. Alternatively, ultrasound-guided needle core (e.g. Tru-Cut) biopsies may be collectable relatively non-invasively, and a recent abstract (Freiche, 2016 #359) suggested that ultrasound-guided fine needle aspirates (FNAs) could be a good alternative to tissue samples for FCoV RT-PCR analysis.

**FCoV RT-PCR on effusion samples**

Effusion samples are usually obtainable using minimally invasive relatively easy techniques, especially if a moderate amount of abdominal or pleural effusion is present. Effusion samples in FIP cases often contain FCoV RNA (Pedersen, 2015 #326); recent published studies have amplified FCoV RNA in most (72-89%) FIP effusion samples but not in any non-FIP effusion samples (Felten, 2017 #350; Longstaff, 2015 #349; Doenges, 2017 #354), although our recent study (Barker, 2017 #383) did amplify FCoV RNA, albeit at a low level, in abdominal fluid from one cat (out of a total of 29 control cats) that did not have FIP. Overall, in tissue samples, the presence of (high) levels of FCoV RNA in an effusion sample that also has cytological and biochemical features consistent with FIP appears to be highly supportive of a diagnosis of FIP.

**FCoV RT-PCR on blood samples**
When FCoV RT-PCR was performed on plasma or serum samples from FIP and non-FIP cats (Doenges, 2017 #354; Felten, 2017 #350), none of the non-FIP cases and very few (9-15.4%) of the FIP cases gave positive results for FCoV RNA. A recent experimental study (Desmares, 2016 #347) also failed to show any FCoV RNA in the plasma of three FCoV-infected cats over the first 12 weeks of infection, and FCoV RNA was rarely detected in the blood of 20 FIP cases (Pedersen, 2015 #326). Thus, use of FCoV RT-PCR on blood, plasma or serum samples is not helpful in the diagnosis of FIP due to low sensitivity. Peripheral blood mononuclear cells (PBMCs) may be a better target for PCR than serum, as shown in one study (Doenges, 2017 #354), but sensitivity was still very poor at 28.6%. Similarly the experimental study (Desmares, 2016 #347) on three FCoV-infected cats only infrequently detected cell-associated FCoV in the blood over the first 12 weeks of infection; in this study the cells used to determine cell-associated viraemia were not stipulated as being PBMCs.

**FCoV RT-PCR on cerebrospinal fluid (CSF) samples**
A recent paper (Doenges, 2016 #345) described the use of FCoV RT-PCR on CSF samples and found it to have 100% specificity for FIP but a sensitivity of only 41.2%. Our study (Barker, 2017 #383) gave similar results. However, not all cats included in these studies had neurological signs as CSF was collected at post-mortem examination independent of presenting signs, such that the population tested may not reflect those that would have had CSF samples collected for diagnostic purposes. In one study, the sensitivity of RT-PCR rose to 85.7% (Doenges, 2016 #345) when only cats with neurological and ophthalmological signs were considered. Thus, FCoV RT-PCR on CSF appears to be a useful additional test in cats with neurological signs, as a positive result highly supports a diagnosis of FIP.

**FCoV RT-PCR on aqueous humour samples**
Robust studies have yet to be performed evaluating FCoV RT-PCR on aqueous humour samples, although positive results have been reported in cats with FIP (Barker, 2017 #383).

**Molecular techniques characterizing FCoV S mutations in FCoV-containing samples**

*How are mutations identified?*
Following the detection of FCoV RNA in a sample by RT-PCR, it may be possible to then characterise targeted sections of FCoV genomic sequences present in that sample using molecular techniques such as pyrosequencing, Sanger sequencing or PCR with sequence-specific hydrolysis probes. Such techniques are not always successful in samples positive for FCoV by RT-PCR if, for example, only low levels of FCoV are present (this can preclude sequence analysis) or if FCoV sequence variability means that targeted sequencing techniques cannot generate sequence results. Characterisation of FCoV genomic sequences would be most useful if FIP-specific mutations existed, as the detection of these mutations would be diagnostic for FIP.

*Is mutation analysis useful?*
Recent research has described amino acid differences in the fusion peptide encoded by the FCoV S gene as being markers of FCoVs associated with FIP (Chang, 2012 #255), raising the possibility that detection of the underlying S gene mutations could be used to definitively diagnose FIP. Similarly, amino acid differences in the furin cleavage motif, also encoded by the S gene, have been correlated with FIP disease (Licitra, 2013 #277). However, these S gene markers were identified by comparing the sequences of FCoVs found in the tissues of FIP cats with those found in the faeces of healthy non-FIP cats.

Researchers in our group hypothesized that the fusion peptide sequence mutations could reflect the cellular tropism of the FCoV (i.e. being systemic monocyte-/ macrophage-associated FCoV compared to intestinal epithelium associated FCoV) rather than being specific for FIP, knowing that non-FIP cats can have systemic FCoV infection. Thus we compared the S gene sequences, from the region of the previously described fusion peptide mutations, of FCoV detected in the tissues of FIP cats with those detected in the tissues of non-FIP cats (Porter, 2014 #321). This allowed us to evaluate the S gene sequences of FCoVs associated with systemic FCoV infection in both non-FIP and FIP cases. We found that the S gene mutations present in most of the FIP
tissues were also present in most of the tissues of non-FIP cats that had systemic FCoV infection. A recent more extensive study confirmed the same findings{Barker, 2017 #383}, and calculated that if the identification of S gene mutated FCoVs was included as an additional confirmatory step to the detection of FCoV alone by RT-PCR, this only slightly increased specificity for the diagnosis of FIP in tissue samples (from 92.6% to 94.6%) but moderately decreased sensitivity (from 89.8% to 80.9%), as non-mutated FCoVs were sometimes identified and mutation analysis was not possible in all tissue samples e.g. due to low FCoV copy numbers or the presence of Type 2 FCoVs). These results question the value of S gene mutation analysis over and above the detection of FCoV by RT-PCR, particularly in view of the extra financial expense and time required to perform this additional analysis.

Analysis for S gene mutations has also been performed on effusions in recently published studies{Felten, 2017 #350; Longstaff, 2015 #349}. The majority of FCoVs in the effusions of FIP cats do indeed have the mutations described{Chang, 2012 #255}. In one study{Longstaff, 2015 #349}, 12/17 FCoV-positive FIP effusion samples had S gene mutations, whilst one did not have a mutation and four could not be sequenced due to the low levels of FCoV present. In another study{Felten, 2017 #350}, 32/36 FCoV-positive FIP effusion samples had S gene mutations, whilst three did not have mutations and one could not be sequenced. Our recent extensive study{Barker, 2017 #383} calculated that the identification of S gene mutated FCoVs as an additional step to the detection of FCoV alone by RT-PCR did not increase specificity for the diagnosis of FIP in fluid (primarily effusions but also included CSF and aqueous humour) samples (specificity stayed at 97.9%) but markedly decreased sensitivity (from 78.4% to 60%, for the same reasons as described for the tissue samples above). In this study{Barker, 2017 #383} all FCoV-positive samples from cats with FIP had S gene mutations in CSF samples, whilst no non-FIP samples were positive for FCoV. Therefore, S gene mutation analysis in FCoVs does not substantially improve the ability to diagnose FIP in effusion or fluid samples as compared with detection of FCoV RNA alone by RT-PCR.

Histopathological examination of tissues
Samples of affected tissues, e.g. liver, kidney or mesenteric lymph nodes, can be collected ante-mortem by ultrasound-guided percutaneous needle-core biopsy, laparoscopy or laparotomy, although the invasive nature of collection may preclude carrying this out in sick cats. Often samples are collected following euthanasia due to a high index of suspicion of FIP at post-mortem examination. Samples are evaluated for characteristic histopathological changes of FIP, which when present are generally regarded as being reliable for diagnosis, although immunostaining for FCoV antigen (see below) is usually also recommended to confirm the diagnosis. However, a lack of histopathological lesions is more difficult to interpret, especially in cases with a high index of suspicion of FIP, as absence of gross lesions to guide biopsy could lead to sampling of non-affected organs or tissue{Giordano, 2005 #243}. A small study{Giori, 2011 #242} recently documented that 5/8 FIP cases did not have histopathology changes typically consistent with FIP, even though large representative biopsies were taken, and diagnosis in these cases was based on positive FCoV antigen immunostaining.

Immunostaining of FCoV antigen
Immunostaining is performed on formalin-fixed tissues using immunohistochemistry (IHC) or on cytological (typically effusion) samples using immunocytochemistry (ICC) or immunofluorescence (IF). These techniques exploit the binding of antibodies to host cell-associated FCoV antigens, which are subsequently visualised by enzymatic reactions producing a colour change (immunochemistry) or by fluorescence (immunofluorescence).

Positive FCoV antigen immunostaining of tissues is said to confirm a diagnosis of FIP (i.e. it is very specific) but a negative result does not exclude FIP as a diagnosis as FCoV antigens may be variably distributed within lesions{Giordano, 2005 #243} and thus are not detected in all histopathological sections prepared from lesions from FIP cases{Kipar, 2014 #309}. This somewhat contradicts the suggestion by some that immunostaining is mandatory to confirm/exclude FIP in doubtful cases{Giori, 2011 #242}, but may be overcome by taking multiple
and/or large samples with confirmed pathology, as well as possibly requesting additional sections of biopsies with pathology to be cut and stained.

Immunostaining of effusion samples has shown variable sensitivity (ranging from 57 to 100%)\cite{Felten, 2017 #355; Hartmann, 2003 #4; Hirscherber, 1995 #42; Litster, 2013 #274; Paltrinieri, 1999 #109; Parodi, 1993 #342}. Since this technique relies on staining FCoV within macrophages in the effusion, and the effusion is often cell-poor and/or the FCoV antigen is masked by FCoV antibodies in the effusion, a false negative result may be obtained. Immunostaining was thought to be very specific, although two (heart failure and cholangiocarcinoma cases) of seven non-FIP effusions were positive by IF in one study\cite{Litster, 2013 #274}, and eight (including two cats with heart failure and two cats with neoplasia) of 29 non-FIP effusions were positive by ICC in another\cite{Felten, 2017 #355}, questioning the specificity of ICC. However, the reported poorer specificity may be due to the methodology used in one study (i.e. double staining for both FCoV antigen and macrophages [via MHC II staining] was used), and the suboptimal storage of slides in the other, which could cause non-specific staining and false positive results. Some have suggested that using cell pellets prepared from centrifuged effusion samples to prepare formalin-fixed, paraffin embedded samples that can then be treated like a tissue specimen for IHC\cite{Kipar, 2014 #309}, can improve the reliability of detection of FCoV antigen\cite{Kipar, 2014 #309}, although the processing time required for this would be longer than for ICC.

FCoV antigen ICC staining has been reported as being successful in detecting FCoV in the CSF of a cat with neurological FIP\cite{Ives, 2013 #291}. A recent study evaluated ICC in the CSF of cats with and without FIP, that presented with and without neurological signs, collected at post-mortem examination\cite{Gruendl, 2016 #361}; this study found that 17 of 20 cats with FIP gave positive results but of 18 cats without FIP, three gave positive results, limiting the test's specificity, although methodology may again be an issue, as described above. These analyses excluded those cases in which cellularity was inadequate for ICC to be performed. Application of ICC to CSF samples collected ante-mortem from a larger number of cats with neurological signs due to FIP and non-FIP causes would be desirable to further evaluate the usefulness of this technique.

The use of FCoV antigen immunostaining has recently been described in aqueous humour samples collected at post-mortem examination from cats with and without FIP\cite{Felten, 2017 #384}. Being able to use aqueous humour for reliable diagnostic investigations in cases of FIP would be especially valuable as it would be possible to collect this in non-effusive cases, although the sample collection technique used in the study would need to be modified (e.g. smaller gauge needle) for use ante-mortem. The study evaluating FCoV ICC in aqueous humour samples from 25 cats with FIP (interestingly the majority were effusive FIP cases, and did not present with uveitis) and 11 non-FIP cats showed a sensitivity of 64% and specificity of 81.8%; positive results were obtained in two of the 11 control cats, one with lymphoma and one with pulmonary adenocarcinoma and both did not have aqueous humour cytological features consistent with FIP (pyogranulomatous inflammation). Further evaluation of ICC on aqueous humour samples collected ante-mortem from cats with uveitis due to FIP and non-FIP causes would be useful to further evaluate the usefulness of this technique.

It is possible that fine needle aspirates could also be used as samples FCoV antigen immunostaining but sensitivity may be poor due to difficulties in targeting lesions; further studies would be necessary to evaluate their utility in the diagnosis of FIP and no evidence to support this currently exists.

**Key Points**
- Look for features that could be suggestive of FIP in the history and on clinical examination: young cat, originally from a multi-cat household (including shelters or catteries), fluctuating non-responsive pyrexia, evidence of an effusion, ocular or neurological signs
- Look for a lymphopenia on haematology
- Look for hyperglobulinaemia, hyperbilirubinaemia (in the absence of moderate to severe increases in ALT and ALP enzyme activity, or anaemia), a reduced albumin:globulin ratio (A:G ratio) of <0.4 and elevated α1-acid glycoprotein (AGP) (> 1.5 mg/ml) on serum biochemistry
- Prioritise finding and sampling any effusion, whether pleural, peritoneal or pericardial in type. Effusions due to FIP are typically clear, viscous, straw-yellow and sticky with a total protein concentration of >35 g/l and a low A:G ratio of <0.4. AGP concentrations can also be raised as in the serum. FIP effusions are poorly cellular (usually <5 x10⁹/l cells) and typically pyogranulomatous in nature
- Demonstration of FCoV antigen in effusions or biopsies, in association with typical cytological or histopathological features of FIP, respectively, provides for a definitive diagnosis of FIP
- The detection of FCoV RNA by reverse transcriptase (RT)-PCR, especially if present at high levels, on diagnostic samples such as effusions, CSF and biopsies, is highly supportive of a diagnosis of FIP
- The detection of FCoV spike gene mutations following a positive FCoV RT-PCR result does not appear to offer additional information for the diagnosis of FIP
Table 1: Diseases to consider in the differential diagnoses of FIP.
Modified from Tasker S and Dowgray N (in production) with permission from BSAVA publications, Gloucester{Tasker, 2017 #380}.

<table>
<thead>
<tr>
<th>Disease/condition</th>
<th>Possible distinguishing features from FIP and course of action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toxoplasmosis</td>
<td><strong>Transmission/epidemiology:</strong> Acquired via vertical transmission (in young cats) or by hunting or eating raw meat. Clinical signs: Cats may have hepatic, pulmonary, neurological, muscle and/or pancreatic involvement. Signs can include lethargy, anorexia, dyspnoea (pneumonia and/or pleural effusions can occur), jaundice, abdominal effusions, uveitis (especially posterior) and/or neurological signs. Diagnostic testing: Clinical toxoplasmosis is less common than FIP and is not usually associated with the severe hyperglobulinaemia or reduced albumin:globulin ratio often seen with FIP. Hyperbilirubinaemia may occur. Serology (high IgM titre or rising IgG titre) may be helpful for diagnosis. Organisms may be found on sampling and microscopic examination of e.g. lung, lymph node. PCR can also be performed on such samples to demonstrate the presence of <em>Toxoplasma gondii</em> DNA. Cerebrospinal fluid (CSF) PCR can also be performed in cases with neurological signs. Treatment: If toxoplasmosis is suspected, trial treatment with clindamycin can be instigated to see whether there is a positive response.</td>
</tr>
<tr>
<td>Lymphocytic cholangitis (LC)</td>
<td><strong>Epidemiology:</strong> Can also occur in young cats. Persians may be predisposed. Clinical signs: Often associated with jaundice. Some cats with LC also have an abdominal (unicavitary) effusion. Diagnostic testing: The nature of the effusion is similar to that seen with FIP in terms of protein concentration (i.e. high), although cell counts in LC are usually higher than those seen with FIP. A marked hyperglobulinaemia can also be seen with LC. Both LC and FIP cases can be hyperbilirubinaemic. Unlike FIP, however, LC is also usually associated with marked increases in liver enzymes, especially cholestatic markers (i.e. ALP and GGT), compared to the more mild or modest increases that occur in FIP cats. Additionally, cats with LC are not usually as sick as those with FIP, e.g. they can be polyphagic rather than inappetent.</td>
</tr>
<tr>
<td>Neoplasia (e.g. lymphoma, abdominal carcinoma)</td>
<td><strong>Epidemiology:</strong> Lymphoma can affect young cats, but is seen in cats of all ages. Other neoplasias tend to be seen in older cats. Focal FIP lesions in the intestine or (especially mesenteric) lymph nodes can present very similar to cases with apparently solitary neoplasms of these organs. Clinical signs: Lymphoma can involve multiple body organs and, like FIP, it can result in lymphadenopathy and/or bicavity effusions. Cats are often systemically ill. Diagnostic testing: Sampling of affected tissues or effusions followed by cytology may yield a diagnosis of lymphoma rather than the mixed inflammatory cells typically seen on cytological sampling of FIP-affected tissues. Other neoplastic lesions, e.g. carcinomas, may be diagnosed on cytology of effusions.</td>
</tr>
<tr>
<td>Pancreatitis</td>
<td><strong>Clinical signs:</strong> Cats may present with anorexia, jaundice and weight loss. Marked pyrexia is not usually a feature, although pyrexia can occur in acute pancreatitis cases that are associated with severe pain and/or sepsis. Diagnostic testing: Hyperbilirubinaemia may occur. A small amount of abdominal fluid (typically with a high protein concentration and high cell count [non-degenerate neutrophils], in contrast to the high protein low cell count effusions with FIP) is sometimes present in acute cases. Pancreatitis can be diagnosed by ultrasonographic examination of the pancreas and measurement of feline pancreatic lipase immunoreactivity. Treatment: Trial treatment with antiemetics and analgesics may be warranted.</td>
</tr>
<tr>
<td>Condition</td>
<td>Epidemiology</td>
</tr>
<tr>
<td>-----------</td>
<td>--------------</td>
</tr>
<tr>
<td>Retroviral infection</td>
<td>Feline leukaemia virus (FeLV) and feline immunodeficiency (FIV) infections are both found more commonly in adults compared to juveniles, but age-related immunity plays a role in FeLV infection with younger cats being more prone to infection. Both viruses more likely to occur in outdoor cats. FeLV affects both males and females whereas males are at increased risk of FIV infection. Clinical signs: Both FeLV and FIV can be associated with pyrexia, lethargy, lymphadenopathy and/or uveitis. Diagnostic testing: FIV infection can be associated with a marked hyperglobulinaemia. NB: Retrovirus-positive status may act as a risk factor for the development of FIP.</td>
</tr>
<tr>
<td>Mycobacterial infection including tuberculosis (TB)</td>
<td>There is a geographical variation in prevalence and infection is usually associated with a history of outdoor access and hunting. Clinical signs: Lymphadenopathy, respiratory signs and/or uveitis may be seen, as well as draining non-healing wounds. Affected cats may be relatively well despite the disease, and pyrexia and inappetence are not common features although acute presentations of e.g. dyspnoea can occur. Mycobacterial infections that involve the lungs typically affect the lung parenchyma rather than presenting with pleural effusions as in FIP. Diagnostic testing: Mycobacterial infection is not usually associated with the severe hyperglobulinaemia or reduced albumin:globulin ratio seen with FIP. Hypercalcaemia may be present. Cytology of affected lymph nodes or organs shows inflammatory changes (macrophages prominent but inflammation can be similar to FIP with pyogranulomatous changes). Ziehl-Neelsen staining on cytology or biopsy samples may be positive, and samples can be submitted for culture (although positive culture results can take weeks to obtain with slower growing organisms, and may be impossible with some mycobacterial species). An interferon gamma test, performed on blood samples, is now available to assist in the diagnosis of suspected feline TB cases.</td>
</tr>
<tr>
<td>Pyothorax</td>
<td>Can be associated with pyrexia. A pleural effusion is seen. Diagnostic testing: Thoracic effusion analysis reveals very high cell counts due to marked neutrophilic inflammation, with degenerate changes and possibly intracellular bacteria, although any previous antibiotic treatment may mean that bacteria are not seen. Unicavity effusion.</td>
</tr>
<tr>
<td>Sepsis</td>
<td>Can be associated with many conditions e.g. septic peritonitis, pyothorax, pneumonia, pyelonephritis. Clinical signs: Can be associated with pyrexia (although cats can also present with low temperatures), tachycardia or bradycardia, tachypnoea, and other signs associated with source of sepsis. Diagnostic testing: Leucocytosis, band neutrophilia, hyperbilirubinaemia (in absence of raised hepatic enzyme activity) may be present.</td>
</tr>
<tr>
<td>Septic peritonitis</td>
<td>Can be associated with pyrexia. An abdominal effusion is seen. Diagnostic testing: Abdominal effusion analysis reveals very high cell counts due to marked neutrophilic inflammation, with degenerate changes and possibly intracellular bacteria, although any previous antibiotic treatment may mean that bacteria are not seen. Unicavity effusion. Glucose concentration in the abdominal effusion is lower than that in the blood (by &gt; 1.1 mmol/l).</td>
</tr>
<tr>
<td>Congestive heart failure (CHF)</td>
<td>Clinical signs: Bicavity effusions in the pleural and peritoneal spaces are possible, although pleural effusions are far more common with feline CHF than abdominal effusions, and abdominal effusions alone are very rarely seen with feline CHF. The presence of a gallop sound, arrhythmia, and possibly heart murmur, may increase...</td>
</tr>
<tr>
<td>Condition</td>
<td>Description</td>
</tr>
<tr>
<td>-----------</td>
<td>-------------</td>
</tr>
<tr>
<td>CHF</td>
<td>The index of suspicion for CHF. Jugular vein distension may be present with right-sided CHF. Pyrexia is not a feature. Diagnostic testing: The fluid is a modified transudate with little protein, in contrast to the fluid seen with FIP. Echocardiography will confirm cardiac disease and CHF.</td>
</tr>
<tr>
<td>Rabies</td>
<td>In countries where rabies is endemic, this must be considered as a differential diagnosis in unvaccinated cats presenting with neurological signs, especially acute behavioural changes and progressive paralysis.</td>
</tr>
</tbody>
</table>
Acknowledgements

The author would like to acknowledge the many contributions made by the University of Bristol Feline Coronavirus Research Group & Bristol-Zurich FIP Consortium to the viewpoints and discussions described in this review. Special thanks go to Emi Barker and Samantha Saunders for their helpful comments on this manuscript. Andrew Davidson, Anja Kipar and Stuart Siddell are also thanks for their valued contributions to past and current FCoV research. Additional thanks are made to the veterinary practices, cat breeders and rescue centres that helped in the acquisition of samples used in our research studies and we also thank our colleagues, current and past, at the Feline Centre and Veterinary Pathology Unit, Langford Vets, University of Bristol, who have assisted in obtaining samples.
**Figure 1:** Schematic diagram of a feline coronavirus particle.

*Modified with kind permission from Dr Emi Barker.*

The spike (S) protein binds to the feline ‘receptor’ mediating host cell entry. The feline receptor is known to be aminopeptidase N for Type 2 FCoVs, but is, as yet, unknown for Type 1 FCoVs.

**Figure 2:** Typical gross post-mortem examination findings in cases of FIP.

Granulomatous lesions in organs or fibrinous plaques on the serosa of organs may be visible in the abdominal or thoracic cavity; tissues that are good to examine for these are the mesenteric lymph nodes, liver, spleen, kidneys and intestinal surfaces, as well as the peritoneal lining of the abdominal wall and diaphragm. In effusive cases, yellow sticky fluid can be visible in the pleural and/or peritoneal cavities, but the pericardium can also be checked for fluid.

**Figure 3:** Examples of clinical signs seen in cases of FIP.

Clinical signs seen with FIP are typically assigned to wet (blue boxes) or dry (red boxes) FIP presentations, but much overlap is seen between the two presentations and many signs are seen in both forms (purple boxes).

**Figure 4:** Typical appearance of effusion seen in cases of FIP.

Effusions in FIP cases are typically clear, viscous and straw-yellow in colour.

**Figure 5:** Positive Rivalta’s test.

The Rivalta’s test is performed by adding a drop of effusion to the surface of a mixture of 8 mls of distilled water and 1 drop of 98% acetic acid (vinegar). A positive Rivalta’s test merely indicates that the effusion being tested is an exudate. It is cheap and quick to do in-house but is not specific for FIP. This is a positive result as the drop has retained its shape with a connection to the surface of the liquid.

**Figure 6:** Schematic diagram of the feline coronavirus genome.

*Modified with kind permission from Dr Emi Barker.*

FCoV RT-PCR assays detect FCoV RNA. The section of the genome amplified by different RT-PCRs varies depending on the position of the primers used in the assays. As viral transcription starts at the 3’ end of the FCoV genome, with the production of multiple subgenomic RNAs at this 3’ end, PCR assays with primers located at the 3’ end of the genome (e.g. in the M or N regions) will be susceptible to viral load overestimation as these will amplify these subgenomic RNAs, as well as the genomic RNA present in the FCoV. Conversely, PCR assays with primers located at the 5’ end of the genome (e.g. the RNA polymerase) will amplify primarily genomic RNA and will be less prone to viral load overestimation. Assays directed at the 3’ end of the FCoV genome will tend to be more sensitive in detecting the presence of FCoV, due to their ability to amplify both subgenomic and genomic RNA.

Coronaviruses, including FCoV, frequently undergo mutations and recombinations, meaning that PCRs designed to be specific for specific sequences may not amplify all FCoVs. PCRs can be designed to target conserved regions of the genome to minimise this, but elimination of FCoV sequence variability as a cause of non-amplification is impossible.
Fibrinous plaques visible on the surface of the spleen

Granulomas evident on surface of kidneys

Yellow sticky effusion visible in the abdomen

Granulomas visible on surface of intestines

Fibrinous plaques visible on the omentum together with enlargement of the mesenteric lymph nodes

Granulomas visible on surface of intestines

Granulomas visible on surface of the lungs
Wet (effusive) FIP

Abdominal effusion (abdominal distension)
Pleural effusion (dyspnoea, tachypnoea)
Pericardial effusion
Scrotal swelling

Dry (non-effusive) FIP

Lethargy
Pyrexia
Anorexia
Weight loss
Jaundice

Neurological signs (e.g. ataxia, head title, hyperaesthesia, nystagmus, seizures, behavioural changes)

Ocular signs (e.g. iritis [iris discolouration], corneal oedema, dyscoria/anisocoria, loss of vision, hyphaema, hypopyon, keratic precipitates, aqueous flare, perivascular cuffing, chorioretinitis, sub-retinal fluid accumulation causing partial retinal detachment)

Non-pruritic papules or nodules

Overlap between wet & dry forms
5' polyprotein including the RNA polymerase

1a

1b

S

spike protein

envelope protein

membrane protein

M

N

3' polyA

non-structural proteins

3 a/b/c

non-structural proteins

7 a/b

nucleocapsid protein
Hector, 3 year old MN Ragdoll

History: 4 weeks of progressive lethargy and reduction in appetite, with severe inappetance and a non-responsive pyrexia documented over the last week.

Clinical examination: Mild dehydration with slight pallor. Chest auscultation was unremarkable. Pyrexia at 40°C. On abdominal palpation a non-painful but large nodular mid-abdominal mass was palpable; it felt quite mobile. No other abdominal abnormalities were noted. Peripheral LNs were normal.

Haematology: A mild non-regenerative anaemia, monocytopenia and eosinopenia were present. The lymphocyte count was normal.

Serum biochemistry: Hypoalbuminaemia (20.4 g/l; RI 24-35) and hyperglobulinaemia (69.3 g/l; RI 21-51), with a markedly reduced albumin to globulin ratio of 0.29, were present. Liver parameters, including bilirubin, were normal. A mild hypokalaemia was present.

Imaging: Abdominal ultrasonography revealed multiple thickening of, and lesions in, an area of the small intestine and associated lymph nodes. Slight hepatomegaly was found, but with normal hepatic echogenicity and a left chronic renal infarct. Chest radiography revealed only mild sternal lymphadenopathy.

Ophthalmic and neurological examinations: These were both normal.

Initial management: Systolic blood pressure was low at 95 mmHg on admittance, and this, together with the presence of dehydration, necessitated administration of intravenous fluid therapy (IVFT), which corrected the dehydration and normalised blood pressure. Urine analysis, done on urine taken after the start of IVFT, was largely unremarkable.

Sampling of the abdominal lesions: Abdominal ultrasonography was repeated and attempts made to take fine needle aspirates of the abdominal masses – this was difficult as the masses were highly vascularised – but samples were taken of the lymph node, and cytology revealed the presence of lymphocytes, neutrophils, plasma cells and macrophages, although some of these cells could have been blood derived. Ziehl-Neelsen (ZN) staining of the cytological samples was negative.

Serum protein electrophoresis (SPE): This confirmed the hypoalbuminaemia and showed the presence of a polyclonal elevation in γ-globulins.

α1-acid glycoprotein (AGP): AGP was markedly elevated at 2.15 mg/ml (RI ≤ 0.48 mg/ml with FIP cases usually having values of >1.5 mg/ml).

Decision making: Hector’s signalment, history, serum biochemistry, SPE and AGP findings were all consistent with a diagnosis of FIP. The cytology of the lymph node was also supportive. On the day following admission, Hector became much duller and his pyrexia remained. Differential diagnoses were discussed with the owner and euthanasia was elected for in view of Hector’s condition and the likely diagnosis of FIP; mycobacterial infection was also considered as a differential diagnosis but Hector was an indoor cat, not a hunter and the ZN staining had been negative.

Post-mortem examination: This revealed a large mass involving the jejunal lymph nodes (centered around major blood vessels) and involving the small intestine. These findings were consistent with focal FIP or mycobacterial infection. Histopathology of sections of the mass showed severe necrosis and fibrosis with a mixed inflammatory cell infiltrate in the wall of the intestine with areas of mucosal ulceration and extension into the serosa; and similar areas of severe necrosis, fibrosis and inflammation in the lymph node mass. Immunostaining for FCoV antigen in the sections was strongly positive. A diagnosis of focal dry FIP was made.