CHAPTER 3.1

African horse sickness

Fast facts
African horse sickness (AHS) is an exotic arthropod-borne viral disease of Equidae transmitted by certain species of Culicoides.

Organism
AHS virus (AHSV):
- has nine different serotypes
- multiplies in regional lymph nodes and is followed by primary viraemia, with subsequent dissemination to target organs.

Susceptible species
AHS affects:
- all species of Equidae (although not all show clinical signs)
- dogs and camels (dead-end hosts).

Economic impact
There is potential for significant economic loss associated with high mortality in horses, as well as restrictions on animal movements and international trade.

Epidemiology
AHS:
- distribution is determined by the presence of competent vectors
- morbidity depends on species, previous immunity and the form of disease
- can cause mortalities, with horses particularly susceptible (up to 95 per cent).

Transmission
Transmission occurs primarily through Culicoides vectors.

Clinical signs
The disease may present as a peracute (pulmonary) form, subacute oedematous (cardiac) form, and acute mixed form or as horse sickness fever. Clinical signs may include:
- pyrexia
- laboured breathing and coughing
- pulmonary oedema
- oedema of the head, conjunctiva, neck, brisket, ventral thorax and abdomen
- death (which is common and can occur suddenly).

Post-mortem
Gross lesions vary depending on the form of the disease, and lesions are not pathognomonic. These may be found:
- subpleural and interlobular oedema
- subcutaneous and intermuscular connective tissue oedema.

Samples
Collect:
- blood samples (EDTA and plain tubes)
- lymph nodes (especially mesenteric and bronchial), lungs, spleen from dead horses.

Actions to take
If you suspect a case of AHS:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including movement controls and protection of animals from vectors) until advised by government veterinary authorities.
Introduction

African horse sickness (AHS) is an arthropod-borne infectious (but non-contagious) viral disease affecting all species of Equidae.

Disease agent and susceptible species

African horse sickness virus (AHSV):
- is a double-stranded RNA virus belonging to the genus Orbivirus and family Reoviridae
- has nine different serotypes and some serotypes are cross protective
- mainly affects equids, with horses, mules, donkeys and zebras the primary hosts
- causes the most serious infections in horses (mortality up to 95 per cent)
- is also known to affect dogs, usually (but not exclusively) following ingestion of virus-infected meat. However, dogs are considered dead-end hosts
- is thought to be maintained in the environment in most regions of Africa by zebras which are often subclinically infected natural reservoir hosts
- antibodies have been reported in camels and African elephants but these species are not considered significant in the epidemiology.

Distribution

AHS:
- geographical distribution and seasonal occurrence are dependent on competent vectors, and studying the dynamics and behaviour of Culicoides spp. is therefore essential to understanding the disease
- appears to be seasonal in endemic areas, where it is preceded by seasons of heavy rain alternating with hot and dry climatic conditions
- occurs in eastern and southern Africa (all serotypes of AHSV)
- virus serotypes 2, 4 and 9 have been found in North and West Africa, from where they occasionally spread into countries surrounding the Mediterranean.

For the latest information on the distribution of AHS, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

None reported.

Epidemiology

Modes of transmission

AHS is not contagious and does not spread by direct contact between horses.

Transmission occurs via:
- vectors, during blood-feeding by Culicoides species—Culicoides imicola is the principal vector in Africa and is present in Australia but other Australian Culicoides species could be involved. The virus has also been isolated from Rhipicephalus sanguineus (a dog tick) and Hyalomma dromedarii (a camel tick) and has been experimentally transmitted by mosquitoes, including Aedes aegypti and Culex pyriens, both of which are present in Australia
• wind, which was implicated in the dispersal of infected Culicoides in some epidemics
• mechanical transmission by other biting flies, but these are unlikely to play a significant role
• parenteral injection of infectious blood.

Disease dynamics
Following ingestion by a competent female Culicoides vector, the virus replicates in the insect gut then translocates and replicates in the salivary glands before infection of the next mammalian host. Following infection of a mammalian host:
• multiplication of AHSV occurs in regional lymph node and is followed by primary viraemia, with subsequent dissemination to endothelial cells of target organs
• the incubation period lasts 2–10 days, depending on viral load, viral virulence and host factors
• the viraemic phase typically lasts only 2–8 days (for horses) but reservoir mammalian host species (such as zebra) have a longer infectious period
• lifelong immunity following recovery is rare as different serotypes are not necessarily cross-protective
• animals that survive infection do not become carriers of the virus
• morbidity and mortality can be as low as 30 per cent and 10 per cent respectively in endemic areas
• high morbidity and high case mortality (of up to 100 per cent) may occur in naïve horses.

Persistence of the agent
AHSV:
• is inactivated in the laboratory with formalin, Virkon® S or radiation
• is destroyed at a pH less than 6, or pH 12 or greater. Acidic disinfectants such as acetic or citric acid have been recommended for decontamination when warranted. Alkaline disinfectants such as sodium hypochlorite are also recommended
• can survive in frozen meat, but is inactivated at temperatures greater than 60°C
• is rapidly destroyed in carcasses that have undergone rigor mortis due to pH fluctuations.

Diagnosis and pathology
Clinical signs
The disease may present as a peracute (pulmonary) form, subacute oedematous (cardiac) form, an acute mixed form, or as horse sickness fever.

Signs of the peracute pulmonary form (‘Dunkop’) include:
• acute pyrexia, followed within a day or two by the sudden onset of severe respiratory distress
• short incubation period, usually 3–4 days
• tachypnoea, forced expiration, profuse sweating, spasmodic coughing and a frothy serofibrinous nasal exudate
• forelegs spread apart stance, extended head and dilated nostrils
dyspnoea, which usually progresses rapidly and causes death within a few hours after respiratory signs appear

pyrexia which subsides gradually while the breathing remains laboured for several days (in recovering animals).

Signs of the **subacute oedematous cardiac form** (‘Dikkop’) include:

- a more protracted and milder incubation than the pulmonary form
- pyrexia lasting less than a week
- oedematous swellings appear in the supraorbital fossae and eyelids, spreading to involve the face, tongue, intermandibular space, laryngeal region, and sometimes the neck, shoulders and chest
- absence of oedema of the ventral side and lower legs
- severe depression, colic, petechiae or ecchymoses on the ventral surface of the tongue, and petechiae in the conjunctivae (all are usually seen in the terminal stages of the disease)
- death from cardiac failure (which often occurs within 1 week of the onset of pyrexic reaction).

The **acute mixed form** is most commonly seen in outbreak cases, where the disease may cause mortality of up to 80 per cent. Signs include:

- a mix of clinical signs from the pulmonary and cardiac forms of the disease
- pyrexia
- mild pulmonary or subclinical cardiac disease followed by oedema, cardiac failure or respiratory failure
- death, usually occurring 3–6 days after the onset of pyrexic reaction.

A combination of pathologies (often not clinically apparent) are usually found at post-mortem. **Horse sickness fever** is:

- the mildest form of the disease
- frequently not clinically diagnosed as mild pyrexia in animals may be subclinical
- usually observed in donkeys, reservoir species (zebras) and partially immune horses infected with heterologous serotypes of the virus.

**Pathology**

Gross lesions are quite variable and depend on the form of the disease. In the respiratory forms of the disease, this may be found:

- interlobular oedema of the lungs
- hydropericardium, pleural effusion
- oedema of the thoracic lymph nodes
- petechial haemorrhages of the pericardium.

In the cardiac forms of the disease, this may be found:

- subcutaneous and intramuscular gelatinous oedema
- epicardial and endocardial ecchymosis, myocarditis
- haemorrhagic gastritis.
Differential diagnosis

Consider in the differential diagnosis:

- **Exotic diseases**
  - dourine
  - equine encephalosis
  - equine piroplasmosis
  - equine viral arteritis (avirulent strains of the virus circulate in Australia)
  - Nipah virus infection
  - surra

- **Endemic diseases**
  - anthrax
  - equine infectious anaemia
  - Hendra virus infection
  - purpura haemorrhagica (a complication of strangles)

- **Non-infectious causes**
  - acute poisoning (plant or chemical)
  - heat stress.

Samples required

Sample collection

Collect these samples from clinically affected animals:

- **Serum**, preferably paired sera
- **EDTA blood** from live animals (at least 10 if possible), preferably at early pyrexic stages (7–10 ml per animal)
- **Fresh tissue** from lymph nodes (especially the mesenteric and bronchial nodes), lungs, spleen.

Transport of samples

For transport:

- chill blood samples and unpreserved tissue samples either at 4°C, or with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.
Diagnostic tests

For AHS diagnosis:

- serological tests available include complement fixation test (CFT), competitive blocking ELISA (which is serogroup specific based on VP7) and viral neutralisation
- nucleic acid detection includes real-time reverse transcription polymerase chain reaction (real-time RT-PCR)
- viral isolates can be detected in blood during the early pyrexic stage.

Reporting requirements

AHS is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect AHS, report it immediately by phoning the **Emergency Animal Disease Watch Hotline on 1800 675 888**, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for AHS, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases (and protecting from vectors, if possible) and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). You need to thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


FIGURE 3.1.1 Abundant froth draining from the nostrils reflects severe pulmonary oedema

[Image credit: PIADC and CFSPH]

FIGURE 3.1.2 Severe interlobular oedema of the lungs with petechiae on the pulmonary pleura and the splenic capsule

[Image credit: PIADC and CFSPH]
### African swine fever

**Fast facts**

African swine fever (ASF) is an exotic highly contagious viral disease of swine clinically indistinguishable from classical swine fever.

**Organism**

ASF virus:
- replicates in *Ornithodoros* ticks
- is found in all tissues and body fluids
- can survive >30 days in pig pens and >140 days in some pork products
- strains vary greatly in virulence.

**Susceptible species**

Only pigs and other suid species are susceptible.

**Economic impact**

There is potential for significant economic loss associated with high mortality and morbidity in pigs, as well as restrictions on animal movements and international trade.

**Epidemiology**

ASF:
- may cause morbidity approaching 100 per cent in naïve herds
- mortality varies with strain virulence and ranges from 0–100 per cent
- may be asymptomatic in wild pigs
- incubation usually lasts 4–20 days.

**Transmission**

Transmission occurs via:
- direct contact, usually oronasal
- ingestion of contaminated swill
- contact with infected carcasses
- indirectly by breeding and fomites
- insect vectors including *Ornithodoros* ticks (biological) and stable flies (mechanical).

**Clinical signs**

ASF clinical signs:
- vary according to the disease form (peracute, acute, subacute or chronic) as influenced by the virus strain, route and dose of infection
- include marked pyrexia, reduced food intake, erythema, cyanosis, recumbency, diarrhoea, abortion and death
- in the chronic form, include transient recurring pyrexia, stunting, emaciation, multi-focal erythema, raised or necrotic areas, coughing and painless joint swelling.

**Post-mortem**

The most common lesions are:
- haemorrhagic spleen (enlarged, friable, dark red or black)
- haemorrhagic lymph nodes, kidney and heart
- focal skin necrosis
- fibrinous pericarditis
- generalised lymphadenopathy
- swollen joints
- consolidated lobules in lung.

**Samples**

Also refer to the detailed section on sampling for sudden death in pigs.

Collect:
- blood (EDTA and plain tubes) from living animals
- fresh tissue from spleen, lymph nodes, kidney, lung and ileum on post mortem.

**Actions to take**

If you suspect a case of ASF:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

African swine fever (ASF) is a highly contagious disease of pigs. It is clinically indistinguishable from classical swine fever (CSF), and similar lesions seen at post-mortem examination. ASF and CSF can only be distinguished by laboratory testing.

Disease agent and susceptible species

ASF virus:
- is an enveloped DNA virus, genus *Asfivirus*, family *Asfarviridae*
- replicates in *Ornithodoros* ticks
- is found in all tissues and body fluids
- is highly resistant to inactivation
- strains vary greatly in their virulence. Domestic and feral pigs (*Sus scrofa*) are the only susceptible mammalian species in Australia outside zoos. European wild boar are also susceptible to ASF and present similar clinical signs and disease dynamics to domestic pigs. In Africa, the warthog, giant forest hog, and bushpigs serve as wild suid reservoirs, and have an inapparent infection.

Distribution

ASF:
- is present in most of sub-Saharan Africa
- was first detected outside Africa in 1957 in Portugal, and then in Belgium, Brazil, Cuba, the Dominican Republic, France, Haiti, Italy, Malta, the Netherlands, Portugal and Spain from the 60s to 90s. The disease has since been eradicated from most of these countries, but is endemic in feral pigs in Sardinia, Italy
- outbreaks have been reported since June 2007 in many countries—most notably in a number of countries in central Europe where the disease had not previously been reported, including Georgia and Russia
- a major outbreak occurred in China in 2018 and subsequently spread throughout South-East Asia.

For the latest information on the distribution of ASF, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

None reported.

Epidemiology

Modes of transmission

Transmission occurs via:
- **contact with infected carcasses**—the carcasses of pigs that die during acute ASF contain more virus than carcasses of chronic carriers of ASF and are therefore more infective to other pigs. Carcasses of chronic carriers have also been linked to the spread of ASF
- **direct contact with infected pigs**—the virus is shed in high concentrations in all secretions and excretions that contain blood. Oral and respiratory infection can also occur between pigs in close contact. Moving infected pigs is the most important way to spread disease between piggeries. Feral pigs could become an important reservoir of the virus in Australia
• **Vectors**—soft ticks belonging to the *Ornithodoros* sp. are important vectors of ASFV in parts of Africa. ASFV replicates in *Ornithodoros* ticks and infection can persist over long periods, allowing the virus to be transmitted during blood meals. Other haematophagous insects such as stable flies (*Stomoxys calcitrans*) feeding on viraemic pigs have been experimentally implicated in the mechanical spread of ASF. Two native species of *Ornithodoros* ticks are found in Australia—*Ornithodoros capensis* (penguin tick) and *Ornithodoros gurneyi* (inornate kangaroo tick). Although *O. capensis* only parasitises sea birds, *O. gurneyi* may feed on dogs, cattle, humans, horses, mice, rats, rabbits and kangaroos (its natural host).

• **Contact with infected faeces and urine**

• **Semen and embryos**—the virus is present in semen and artificial insemination can transmit the virus. The International Embryo Transfer Society has indicated that there is not enough information to reach a conclusion regarding the transmission risk of ASF virus via embryos, but you should assume transmission via embryo transfer until proven otherwise.

• **Ingesting contaminated swill**—ASF virus has the ability to survive for many months in raw, unprocessed, frozen meat as well as meat stored at fridge temperatures and in some processed hams. Ingesting pig meat or products infected with the virus is an important means of spread, and is commonly the way it is introduced into a country. Australia has strict import conditions to address the risk of an ASF incursion and swill feeding is illegal.

• **Fomites**—because of the stability of ASF virus in the environment, indirect contact by means of contaminated pig pens, vehicles and other fomites (such as hypodermic needles) is a very important means of spread of the disease.

• **Airborne spread**—ASF virus is not transmitted in the air from one piggery to another, but it does spread via aerosols within a piggery. Respiratory infection can occur between pigs in close contact.

### Disease dynamics

• Following infection via the upper respiratory tract, the virus replicates in the tonsils and lymph nodes draining the head and neck, followed by generalised infection via the blood stream.

• Incubation period is usually between 4–20 days.

• Clinical disease may be peracute, acute, subacute or chronic.

• In the acute form, has a duration of 1–19 days and case mortality rate of up to 100 per cent.

• In the subacute form, persists for longer (3–4 weeks) and the case mortality rate is lower, with deaths more likely in younger pigs.

• Pigs that survive the acute disease or are infected by mild strains of the virus usually become chronically infected for at least several months. However, such pigs do not readily excrete the virus for more than about 5 or 6 weeks after their initial infection and their role in the spread of the disease after that is unclear.

• It is reported that most recovered pigs are virus carriers for long periods, perhaps for life.

• Transmission to other pigs through direct contact can occur for up to 1 month after recovery from infection.
Persistence of the agent

ASFV:
- can survive at least 30 days in pens and up to 300 days in some pork products
- is highly resistant to inactivation but can be killed by high temperatures and some disinfectants—potassium peroxymonosulfate, hypochlorites, and phenols and related compounds (cresols).

Diagnosis and pathology

Clinical signs
Signs vary according to whether disease is peracute, acute, subacute or chronic.
In the peracute form, pigs are usually found dead with no prior clinical signs and if seen before death have marked pyrexia and are moribund.
In the acute form signs are:
- pyrexia up to 42°C
- hyperaemia or cyanosis of extremities, particularly ears and snout
- loss of appetite or irregular appetite
- inability or unwillingness to stand up
- convulsions
- incoordination or stiff gait
- huddling together or piling one on top of another
- laboured breathing or coughing
- dysentery or diarrhoea (often bloody)
- conjunctivitis
- mucopurulent nasal discharge
- vomiting
- abortion.
In the subacute form clinical signs are:
- as for the acute form but generally milder and persisting longer (3–4 weeks)
- pyrexia, which may fluctuate irregularly and may exceed 40.5°C
- occasionally, a purple colour over the pig’s body
- bleeding from injection sites
- abortion.
In the chronic form (generally seen in pigs surviving the subacute form) signs are:
- recurrent transient pyrexia
- ill-thrift, stunting and emaciation
- pneumonia
- arthritis
- cutaneous ulcers
- poor body condition.
### Pathology

In the peracute form pigs may show no gross lesions. In the acute form, findings include:
- enlarged and haemorrhagic lymph nodes, often resembling blood clots
- the gastrohepatic, renal, mesenteric and submandibular lymph nodes are most often affected
- enlarged spleen (2–3 times its normal size), which may be necrotic, dark, friable or pulpy
- haemorrhages in almost any organ which are most commonly seen on serosal membranes and in kidneys (as subcapsular petechiae), heart, urinary bladder, lung and gall bladder
- septal oedema of lungs, resulting in prominent interlobular septa
- fluid in body cavities.

In the subacute form findings are more variable than for the acute form and include:
- lymph node and renal haemorrhage
- enlarged but not congested spleen
- lobular consolidation of cranial lung lobes
- haemorrhage of the intestinal lining, lymph nodes and kidney.

In the chronic form findings include:
- enlarged lymph nodes
- fibrinous pericarditis and pleurisy
- lobular consolidation of lungs, which may progress to lobular necrosis
- small, hard, nodular white masses in lungs
- arthritis
- cutaneous ulcers
- poor body condition
- less commonly seen are haemorrhages in additional organs (petechiae, ecchymoses) and oedema in the lungs and gall bladder.

### Differential diagnosis

Many other viral and bacterial pathogens of pigs that are often confused with ASF may cause concurrent infections. It is important to take into consideration that isolation of other pathogens does not exclude an underlying ASF virus infection. Consider in the differential diagnosis:
- exotic diseases
  - *Aujeszky’s disease* (respiratory signs)
  - *classical swine fever* (do not attempt clinical differentiation of classical swine fever and ASF)
  - porcine deltacoronavirus
  - porcine respiratory disease complex
  - *porcine reproductive and respiratory syndrome*
  - post-weaning multi-systemic wasting syndrome
  - swine enteric coronavirus diseases such as those caused by *porcine epidemic diarrhoea* virus
  - *transmissible gastroenteritis*
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• endemic diseases
  – acute salmonellosis (diarrhoea)
  – acute septicaemia due to *Streptococcus suis* or *Haemophilus parasuis* (Glässer’s disease)
  – infection with other pestiviruses such as border disease virus and bovine viral diarrhoea virus (mucosal disease)
  – pasteurellosis (pneumonia)
  – porcine circovirus 2-associated diseases such as those seen in porcine dermatitis and nephropathy syndrome
  – erysipelas (haemorrhagic skin lesions)
• non-infectious causes
  – heavy metal poisoning
  – salt poisoning with water deprivation
  – thrombocytopaenia purpura (haemorrhages, especially in pigs 2–3 weeks old)
  – warfarin poisoning (causing death with haemorrhages, generally sporadic)
  – any cause of chronic ill-thrift in pigs.

Samples required

Submit several pigs (at least five if possible) for post-mortem examination, as individual animals may present with great variability in lesions. Record a composite picture of all lesions seen.

Sample collection

Collect samples from affected, especially pyrexic, pigs killed immediately before a post-mortem examination and from pigs that have recently died (including stillborn piglets and aborted foetuses). Collect:

• serum, 30 samples from suspected chronically infected animals
• EDTA blood (7–10 ml/animal) from live, clinically affected animals
• fresh tissue from the spleen, lymph nodes (gastro-hepatic, mesenteric and sub-mandibular) tonsils, lung, kidney and ileum (2 g of each tissue); bone marrow may also be useful from dead wild animals that have been dead for some time, as it is likely to be relatively well preserved
• fixed tissue, a full range of tissues (including the brain) in neutral-buffered formalin.

Transport of samples

For transport:

• chill blood samples and unpreserved tissue samples either at 4°C, or with frozen gel packs
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days.
• formalin fixed tissue can be sent at room temperature.
Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
You must confirm the diagnosis by laboratory testing because there is considerable overlap in the clinical and pathological signs seen in ASF with those of a number of other diseases. For ASF diagnosis:
- the initial approach is screening by real-time PCR (qPCR). An antigen ELISA is also available if required and virus isolation will be attempted. Further characterisation and genotyping by sequence analysis can be carried out on primary samples or on isolates
- serological tests are available but are generally more important in defining the nature and extent of any outbreak, and in the proof-of-freedom phase
- histopathology and immunohistochemical staining are also useful diagnostic tests.

Reporting requirements
ASF is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect ASF, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
There are no public health implications for ASF, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). You need to thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information
**FIGURE 3.2.1** Multiple sharply demarcated foci of cutaneous haemorrhage and/or necrosis; haemorrhagic lesions may contain dark red (necrotic) centres

Image credit: PIADC and CFSPH

**FIGURE 3.2.2** Haemorrhagic submandibular lymph nodes

Image credit: AAHL
FIGURE 3.2.3 Petecchial haemorrhages in the intestines

Image credit: AAHL

FIGURE 3.2.4 Haemorrhagic fluid in the abdominal cavity

Image credit: AAHL
CHAPTER 3.3
Aujeszky’s disease

Fast facts
Aujeszky’s disease, also called pseudorabies, is an important exotic viral disease of pigs which affects the nervous, respiratory and reproductive systems.

Organism
Aujeszky’s disease is caused by *Suid alphaherpesvirus 1*. The virus:
- is inactivated by drying, exposure to sunlight and high temperatures
- strain virulence varies from mild to severe.

Susceptible species
Pigs are the only reservoir host but the virus can infect cattle, sheep, cats, dogs, goats and wildlife.

Economic impact
There is potential for significant economic loss associated with production losses, as well as restrictions on animal movement and international trade.

Epidemiology
Aujeszky’s disease is characterised by:
- morbidity varying from 0–100 per cent
- mortality which may approach 100 per cent in newborn pigs, 5–10 per cent in weaners and 1–2 per cent in growers and finishers
- slow spread between farms but rapid within-farm spread
- often mild or inapparent infection in adult pigs
- high mortality in non-porcine species.

Clinical signs
Clinical signs are not pathognomonic but Aujeszky’s disease should be suspected in:
- pig herds with high mortality
- young piglets with central nervous system (CNS) signs
- older animals with lower mortality and respiratory signs
- non-porcine species which present with intense pruritis (‘mad itch’) and CNS signs.

Post-mortem
At post-mortem, gross lesions are often minimal or absent.

Samples
Also refer to the [detailed section](#) on sampling for neurological diseases in pigs. Collect:
- unpreserved spleen, tonsils, lymph nodes, distal ileum and brain
- sera from animals (30) suspected of having chronic disease (or of giving birth to piglets with chronic disease)
- a full range of tissues (including brain and spinal cord) in neutral buffered formalin.

Actions to take
If you suspect a case of Aujeszky’s disease:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Aujeszky’s disease, also called pseudorabies, is an acute, potentially fatal porcine disease which causes significant production losses.

Disease agent and susceptible species

Aujeszky’s disease is caused by

- an enveloped, double stranded DNA virus called suid alphaherpesvirus 1 (SuHV-1) in the family *Herpesviridae*
- there is only one serotype but strains vary in virulence (from mild to severe), minimum infective dose and tissue tropism.

Pigs are the only natural host for Aujeszky’s disease virus and the only latent carrier; however:

- the virus can infect nearly all domesticated and wild mammals (including cattle, sheep, cats, dogs and goats) but infections in horses are rare
- other species may be infected when they come in contact with infected pigs, resulting in a fatal central nervous system (CNS) disease
- infection of incidental hosts is often characterised by intense pruritus
- sporadic infection of incidental hosts results in acute neurological disease and death.

Distribution

- occurs in most countries of Europe and Asia
- in parts of the United States (feral pigs only)
- occurs in Central and South America
- was eradicated from the United Kingdom during the 1980s
- New Zealand declared itself free from Aujeszky’s disease in 2000, following an eradication campaign in the 1990s.

For the latest information on the distribution of Aujeszky’s disease, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int](http://www.oie.int).

Occurrences in Australia

None reported.

Epidemiology

Modes of transmission

Transmission occurs via:

- **direct contact** with oronasal secretions (between pigs); the virus is spread principally by nose-to-nose contact
- **ingestion of contaminated pig meat** (including head or offal tissue) by other susceptible carnivores
- **transplacental infection**, or infective semen or vaginal secretions
- **ingestion of infectious colostrum or milk**
• **ingestion of infectious animal products**—Aujeszky’s disease virus can survive in offal (head and neck tissues, and thoracic and abdominal viscera). However, though you can isolate the virus from meat products, it is easily inactivated. Movements of live animals or infected genetic material are generally the way it is introduced to disease-free countries, rather than importing carcass meat (excluding offal and heads)

• **fomites**—in addition to people, vehicles and equipment, contaminated pig pens, bedding and water can also spread the disease

• **airborne spread**—under certain favourable conditions in densely populated pig-farming areas, windborne spread of the virus between farms can occur over distances of more than 2 km.

**Disease dynamics**

The virus is maintained in wild and domestic pigs, in which the virus can establish latent infections like other herpes viruses. In addition:

- in general, pigs excrete virus oronasally during the 2–4 week period following the primary infection
- animals other than pigs are generally regarded as dead-end hosts, as infection is usually short and self-limiting (most die after an illness of short duration, usually 2–3 days after clinical signs appear)
- the incubation period can be as short as 2–4 days in sucking pigs and 3–6 days in finishers
- virus excretion begins 2–5 days after infection and can continue for at least 14 days (this may precede the onset of clinical signs)
- a very high percentage of recovered pigs become latent carriers for 1 year or longer, with intermittent virus excretion at times when the animal is stressed, such as at parturition.

**Persistence of the agent**

Aujeszky's disease virus is a large virus with a lipid envelope which is sensitive to many disinfectants, including detergents. The virus:

- is rapidly inactivated at 37°C in sunlight and in dry conditions
- has fairly stable infectivity at pH 5–9, but extreme acidity and alkalinity have a rapid inactivating effect
- survives for extended periods under cold conditions (below 4°C)
- survives in contaminated straw and feeding troughs for 10–30 days at 24°C or for up to 46 days at 20°C, and on other fomites for 2–7 days at 25°C
- survives in faeces and urine for up to 3 days
- survives in well water and green pasture for up to 7 days.

**Diagnosis and pathology**

**Clinical signs**

The most striking feature of infection in species other than pigs is intense pruritus of a localised area or areas of skin, often leading to self-mutilation. Neurological signs vary and include excitement, salivation, spasms and paralysis. Death usually occurs about 2 days after the onset of signs. Some animals may die suddenly before other clinical signs develop.
In pigs:
- disease affects the nervous, respiratory and reproductive systems
- clinical signs can vary considerably according to the strain, dose of virus and the age of the pig
- neurological signs are more commonly seen in younger animals, whereas respiratory signs are usually observed in older animals
- clinical signs in pigs are highly variable and largely age-dependent
  - pigs less than 2 weeks old—prostration is often the only clinical sign but pyrexia, altered appetite, vomiting, depression, central nervous system and respiratory involvement may also be seen. Death occurs within hours of the onset of clinical signs
  - weaners—respiratory signs are usually more prominent than neurological signs. These include coughing, sneezing, laboured breathing and conjunctivitis. The case mortality rate is generally 5–10 per cent
  - growers/finishers—respiratory disease is most common and may be mild. Morbidity may reach 100 per cent, with mortality in uncomplicated cases low (1–2 per cent)
  - adults—infection is usually mild or inapparent, with respiratory signs most common. However, severe outbreaks have been reported with severe respiratory signs, and sporadic cases of neurologic disease.
  - pregnant sows—may reabsorb infected foetuses, abort or give birth to weak, trembling neonates. Often retrospective analysis of reproductive parameters may give an indication of the time of infection.

Pathology
In species other than pigs, gross lesions are often minimal and may consist of spinal cord oedema, congestion and haemorrhage. Gross lesions are often minimal or absent in pigs and, when present, are typically seen in young animals. This may be seen:
- multifocal, pinpoint areas of acute necrosis and haemorrhage in the tonsils, liver, spleen, lung, gastrointestinal tract and adrenal glands
- oedematous and congested lungs
- foetal resorption, mummification, abortion, stillbirths and placental necrosis.

Differential diagnosis
Consider in the differential diagnosis:
- exotic diseases
  - African swine fever
  - classical swine fever
  - haemagglutinating encephalomyelitis virus
  - influenza A viruses in pigs (exotic strains)
  - porcine reproductive and respiratory syndrome
  - post-weaning multi-systemic wasting syndrome
  - porcine enterovirus
  - rabies
• endemic diseases
  – bacterial meningitis (*Streptococcus suis, Haemophilus parasuis*)
  – bacterial pneumonia including *Mycoplasma hyopneumoniae*
  – porcine circovirus 2-associated diseases such as those seen in porcine dermatitis and nephropathy syndrome
  – influenza A viruses in pigs (human-origin strains, subtypes H1N1, H1N2 and H3N2)
• non-infectious causes
  – congenital defects
  – iron toxicity
  – middle ear infection
  – nutritional deficiencies (e.g. pantothenic acid)
  – poisons such as arsenic, mercury, monensin and organophosphorus compounds
  – porcine stress syndrome
  – splay leg
  – water deprivation or salt poisoning.

Samples required

Sample collection

For pigs, collect:
• a full range of tissues (including the brain) in neutral-buffered formalin
• 30 serum samples from suspected chronically infected animals (serum antibodies are detected 7–10 days after infection)
• EDTA blood (7–10 ml/animal) from at least 10 live, clinically affected animals
• sera from sows suspected to have had piglets with chronic disease
• nasal swabs from live clinically affected animals, and place in viral transport medium
• fresh tissue (collected aseptically at post-mortem) including spleen, tonsils, pharyngeal mucosa, lymph nodes and distal ileum (2 g of each tissue). Lung, kidney and liver may be included principally for differential diagnostic workup. One half of the brain should be collected aseptically (the trigeminal nerve ganglia should be included if latent infection is suspected).

For other species, also collect the skin and subcutaneous tissue at the site of the itching.

Transport of samples

For transport:
• chill blood samples and unpreserved tissue samples either at 4°C, or with frozen gel packs
• place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days
• formalin fixed tissue can be sent at room temperature.
Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Since clinical signs are relatively non-specific, a presumptive diagnosis of Aujeszky’s disease may be based initially on histopathology and confirmed by positive serology, or virus or antigen detection in tissue or serum. In addition:

- Aujeszky’s disease can be diagnosed by virus isolation, detection of viral DNA, antigens and serology. ELISAs and virus neutralisation tests are the prescribed tests for international trade.
- If vaccination is to be used in any eradication program, certain vaccines permit the serologic discrimination of natural versus vaccine-derived antibody (DIVA).
- Serology may not be helpful in species other than pigs as these animals often die before mounting an antibody response.

Reporting requirements
Aujeszky’s disease is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect Aujeszky’s disease, report it by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
There are no public health implications for Aujeszky’s disease, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information
FIGURE 3.3.1 Crusting of the mucosal membranes around the eye and nares, and the eye has periorbital serous exudate

Image credit: AFIP and CFSPH
CHAPTER 3.4
Avian influenza

Fast facts
Avian influenza (AI) is a zoonotic, highly contagious viral disease of birds causing two types of disease—highly pathogenic avian influenza (HPAI) and low pathogenic avian influenza (LPAI).

Organism
AI virus:
- varies in virulence between subtypes and species
- replicates in respiratory and intestinal epithelial cells (LPAI) or replicates in many cell types, including heart and brain (HPAI)
- inactivated by most disinfectants.

Susceptible species
AI affects all bird species, with a spillover infection in mammals, including humans.

Economic impact
There is potential for significant economic loss associated with control and eradication, and restrictions on international trade.

Epidemiology
In infected flocks:
- infection with LPAI viruses is usually subclinical
- mortality can reach 100 per cent in the acute form of HPAI
- the incubation period varies depending on the strain, and can range from hours up to 16 days.

Transmission
Transmission occurs:
- via the fecal oral route from wild waterfowl into commercial poultry
- via aerosol and direct and indirect contact within flocks
- between flocks primarily via contaminated fomites.

Clinical signs
Signs vary with virus virulence, and species, and can include:
- mortality
- subcutaneous oedema and cyanosis of the comb and wattles
- mild respiratory disease
- huddling, decreased feed and water consumption and diarrhoea
- reduced egg production.
- Waterfowl (including ducks) typically resistant to disease but some HPAI viruses may cause significant mortality.

Post-mortem
This may be found:
- gross lesions of internal organs with multifocal pinpoint haemorrhages, particularly in the trachea, coronary fat, and surfaces of the gastrointestinal tract
- catarrhal air sacculitis and peritonitis, usually associated with secondary infection.
Samples
Collect:
• tracheal and cloacal swabs in virus transport media
• blood samples (EDTA and plain tubes)
• proventriculus, pancreas, intestine, liver, caecal tonsil, trachea, lung and cloaca tissue.

Actions to take
If you suspect a case of AI:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

Public health considerations
• AI is a zoonotic disease
• people may become infected after close contact with infected birds, excreta or litter. There is also a risk of infection from aerosol droplet and air contamination, though this is more rare
• consult with the relevant state or territory health authorities as soon as AI is strongly suspected or confirmed.
Introduction

Avian influenza (AI) is a highly contagious viral disease of birds which may take the form of highly pathogenic avian influenza (HPAI) or low pathogenic avian influenza (LPAI).

Disease agent and susceptible species

AI:
- is caused by the Influenza A virus, which is an enveloped, single stranded negative sense RNA virus in the family Orthomyxoviridae, genus Influenza A
- viruses have two surface antigens haemagglutinin and neuraminidase. Avian viruses have 16 types of haemagglutinin antigen (H1-16) and 9 types of neuraminidase antigen (N1-9)
- wild waterfowl and shorebirds (which rarely show disease) are the reservoir hosts for all influenza A virus subtypes
- influenza A affects all bird species but the domestic poultry species most susceptible to disease are chickens, turkey, guinea fowl, quail and pheasants. Waterfowl (including ducks) are typically resistant to disease, but HPAI viruses may still cause mortalities
- HPAI and LPAI are distinguishable by lethality in chickens as well as a molecular sequence in the haemagglutin protein of the virus. Only H5 and H7 viruses are known to cause HPAI, and H5 and H7 LPAI viruses may mutate spontaneously into HPAI especially following passage in poultry flocks
- LPAI viruses typically replicate in respiratory and intestinal epithelial cells while HPAI viruses replicate in many cell types, including heart and brain
- Both LPAI and HPAI virus can infect humans via close contact with infected birds. Most human infections have occurred in China, South-East Asia, Egypt and Indonesia. Mortality is especially reported in young children, older people, and immunosuppressed people, though HPAI can cause death in healthy adults. The main subtypes that infect humans include H5N1, H5N6, H7N9 and H9N2; however, subtypes H6N1, H7N4, H7N7 and H10N8 have also been reported to infect people.

Public health considerations

- people may become infected after close contact with infected birds, excreta or litter
- people in close contact with infected poultry and poultry facilities should use personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots), safety glasses and powered air purifying respirators
- it is important that personnel seek medical advice if they develop influenza-like symptoms or otherwise become unwell. Usually, symptoms of AI infection appear 2–8 days after infection (see http://www.health.gov.au/avian_influenza)
- consult the relevant state or territory health authorities in the case of an AI outbreak.
Distribution
LPAI and HPAI viruses of all H and N subtypes are endemic in wild waterfowl and shorebirds globally. LPAI outbreaks (clinical and subclinical) and HPAI outbreaks occur in poultry flocks globally, linked to the circulation of viruses in wild migratory birds and live poultry markets.

For the latest information on the distribution of AI, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int] or the FAO EMPRESi Global Animal Disease Information System [http://empres-i.fao.org/eipws3g/].

Occurrences in Australia
In Australia, all seven outbreaks of HPAI were eradicated and involved H7 subtype viruses causing clinical disease in commercial poultry in:
- Victoria—1976, 1985 and 1992
- Queensland—1994

In Australian poultry several LPAI viruses, including H5, H7 and H9 subtypes, have been detected. There was evidence of contact with waterfowl in many cases, and indications that LPAI viruses were initially involved in the outbreaks. These then evolved to HPAI with multiple passages in the flocks.

Epidemiology
Modes of transmission
Transmission occurs via:
- **fomites** such as water, food supplies, clothing, bird cages, pallets, egg filler flats, manure, feedstuff, people, equipment and vehicles that are contaminated with infected faeces or respiratory secretions. Water or food supplies contaminated by wildfowl faeces can introduce AI to caged poultry, and the movement of infected birds and fomites is the primary way to spread HPAI virus between flocks to
- **eggs** may be a source of AI infection through shell contamination. While LPAI virus shedding in eggs has not been detected to date, HPAI viruses have been found in the yolk and albumen of eggs from chickens, turkeys and quail, suggesting the potential for vertical transmission exists
- **direct contact between birds**—LPAI and HPAI are likely to be introduced through direct contact with waterfowl (in the case of free range farms). Contact with faeces and respiratory secretions is important for AI transmission between birds, and aerosol transmission is possible within the poultry flock due to the close proximity of birds.
Disease dynamics

Wild aquatic birds (which rarely show disease) are the reservoir host for all influenza A virus subtypes, particularly members of the orders Anseriformes (ducks and geese) and Charadriiformes (gulls and shorebirds). Migratory birds are considered to be one of the ways the disease travels across and between continents.

In domestic poultry:

- the incubation period of AI can vary from hours to up to 2–3 days in individual birds depending on the strain
- LPAI virus infections in the flock may not be evident for 2 weeks
- infected birds can spread virus via respiratory and faecal secretions from as early as 24 hours after infection (spread is therefore possible during the incubation period and before the onset of clinical signs)
- most chickens excrete LPAI viruses for 1–2 weeks in contrast to water birds which may shed some LPAI or HPAI viruses for 3–4 weeks
- uncontrolled H5 or H7 LPAI infections have been known to mutate into virulent HPAI infections.

Persistence of the agent

AI viruses:

- can survive in aerosols, faeces and litter and survival is prolonged by high relative humidity and low temperature (the virus can survive within the poultry house environment for up to 5 weeks)
- remain infective in lake water for up to 4 days at 22°C and for more than 30 days at 0°C
- can be inactivated by heat of 60°C for 60 minutes, or extremes of pH (pH<3 or >10)
- are highly susceptible to disinfectants, including detergents, but only if organic matter is removed from surfaces before disinfection.

Diagnosis and pathology

Clinical signs

LPAI viruses usually cause:

- subclinical infections or mild respiratory disease in poultry
- high morbidity (>50 per cent) and low mortality (<5 per cent), but mortality can be high in young birds or when AI is accompanied by secondary pathogens
- respiratory signs of varying severity, including coughing, sneezing, rales, rattles, nasal discharge and excessive lacrimation
- general clinical signs such as huddling, ruffled feathers, listlessness, decreased feed and water consumption and diarrhoea
- reduced egg production (in layers and breeders).
While there are no clinical signs pathognomonic for HPAI, signs in chickens may include:

- sudden onset of disease and rapid death (in very severe, peracute forms)
- profuse watery diarrhoea, oedema and cyanosis of the unfeathered skin and combs, ecchymoses on the shanks and feet, swelling of the face, coughing and neurological signs
- depressed birds with reduced egg production and egg quality that is more pronounced than with LPAI
- Clinical signs in other gallinaceous birds (such as turkeys and quail) are similar to chickens. However, there are cases where H5 or H7 AI viruses have caused only mild illness in chickens and turkeys despite being classified as HPAI virus by molecular pathotyping.
- Ducks and other waterfowl are typically resistant to disease caused by HPAI viruses, but some virus strains, including Asian-lineage H5 HPAI viruses, may cause acute mortality in these species.

**Pathology**

With LPAI:

- gross lesions are variable depending on the virus strain, host species and presence of secondary infection
- the infraorbital sinuses may be swollen, especially in turkeys, with accompanying mucoid nasal discharge
- the most frequent lesions are rhinitis and sinusitis of varying severity
- tracheal mucosa may be oedematous and congested, with occasional haemorrhages or exudate
- catarrhal air sacculitis and peritonitis may also be present, usually associated with secondary infection rather than LPAI
- swollen kidneys may occur in laying hens, as well as regression or haemorrhage of ovaries and ruptured ova ('egg yolk peritonitis')
- enteritis may be observed, especially in turkeys, and turkeys may have a pale and mottled pancreas.

With HPAI:

- gross lesions vary with the stage of infection
- in the peracute phase (when death occurs 1–2 days post-infection) there may be no gross lesions evident
- from 2–5 days post-infection, necrosis and cyanosis of the comb and wattles are common (subcutaneous oedema is also evident around the head, legs and feet, along with ecchymotic subcutaneous haemorrhages)
- internal organs most consistently have focal haemorrhages, particularly in the trachea, coronary fat, epicardium, proventriculus, Peyer’s patches and caecal tonsils
- severe oedema and haemorrhage in the lungs can occur.
Differential diagnosis

Note that many other viral and bacterial pathogens of chickens can present with similar clinical signs, or cause concomitant infections with AI. It is important for you to take into consideration that isolation of other pathogens may mask an underlying AI virus infection, and AI should be tested for if the clinical signs and pathology are present. This is especially important to remember considering the spectrum of signs that can be seen with LPAI.

Consider in the differential diagnosis:

- **exotic diseases**
  - duck plague (duck viral enteritis)
  - velogenic Newcastle disease
- **endemic diseases**
  - avian chlamydiosis
  - avirulent Newcastle disease
  - bacterial septicaemia (enterococcosis, staphylococcosis, streptococcosis, colibacillosis, pasteurellosis, erysipelas)
  - cellulitis of the comb and wattles (*Escherichia coli* and other bacteria)
  - fowl cholera (*Pasteurella multocida*)
  - infectious bronchitis
  - infectious coryza
  - infectious laryngotracheitis
  - mycoplasmosis
- **non-infectious causes**
  - acute poisoning
  - dehydration
  - heat stress
  - smothering.

Samples required

Sample collection

Where AI is suspected, collect:

- **tracheal, cloacal and fecal swabs** from live, clinically affected birds
- **serum** from birds suspected to have been clinically affected for a number of days (at least seven birds) or recovered birds
- **fresh tissue samples and whole blood** (EDTA tube, 2 ml/bird) from clinically affected birds (killed immediately before a post-mortem examination) and from any recently dead birds. Fresh tissue should include samples from the brain, proventriculus, pancreas, intestine, liver, caecal tonsil, trachea, lungs and cloaca
- **impression smears** of the bursa of Fabricius, pancreas, brain, and spleen
- **a full range of tissues** (including the brain, proventriculus, pancreas, intestine, liver, caecal tonsil, trachea, lung, bursa, kidney and cloaca) in neutral-buffered formalin
- **several whole birds** (at least 5 birds) for post-mortem examination, as there may be great variability in lesions presented in individual animals. A composite picture of all lesions seen should be recorded.
Transport of samples

For transport:
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- chill blood samples and unpreserved tissue samples (including whole carcasses) at 4°C, or with frozen gel packs
- DO NOT freeze samples at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be shipped at room temperature.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests

For AI:
- the most rapid, sensitive and specific diagnostic procedure is the detection of viral nucleic acid in whole blood and/or tissues by PCR (a positive result can be obtained within 1 day of pyrexia first developing in an animal)
- in order to quickly determine whether the virus is highly pathogenic, molecular pathotyping is carried out by sequencing the gene encoding the haemagglutination protein; this relies on adequate PCR detection
- commercial ELISA kits are available for detection of AI virus antigen and antibody but are comparatively less sensitive than molecular techniques, and antibodies against the AI virus will not be present in acutely affected birds
- viral antigen can also be detected in frozen sections of bursa of Fabricius, pancreas, brain, and spleen using direct immunofluorescence, and may yield a result in a matter of hours
- isolation of the virus is attempted by inoculation of tissue suspensions into embryonated chicken eggs. This takes 1–10 days and may be needed to improve genetic detection.

Reporting requirements

Cases of AI in commercial poultry flocks are nationally notifiable, and infection with HPAI and H5/H7 LPAI viruses are notifiable to the OIE. Australia has an international obligation to report cases. If you suspect AI, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.
**Biocontainment and personal protective equipment**

AI is a public health risk. Human infections are often due to close contact with infected birds; however, influenza A viruses are generally transmissible by fomites and aerosol droplets. You must take this zoonotic risk into consideration when investigating an outbreak of sudden deaths, gastrointestinal or respiratory disease in poultry. Therefore, it is important to seek advice from government veterinary authorities, noting:

- **biocontainment** requires training in the use and appropriate disposal of personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots) and powered air purifying respirators
- **personal decontamination** is required when leaving infected premises. Thoroughly clean and then disinfect vehicles and equipment before leaving the property
- you need to avoid the spread of infection from premises by contaminated fomites such as personnel clothing, poultry litter, equipment and manure
- any people participating in the biocontainment of a HPAI outbreak must be vaccinated against seasonal influenza.

**Further information**


FIGURE 3.4.1 Congested and markedly oedematous wattles and comb of a chicken

Image credit: Dr D Swayne, USDA and CFSPH

FIGURE 3.4.2 Swollen (oedematus) and extensively reddened (haemorrhagic) chicken legs

Image credit: Dr D Swayne USDA and CFSPH
CHAPTER 3.5
Bluetongue disease

Fast facts
Bluetongue is an arthropod-borne viral disease of ruminants transmitted by certain species of *Culicoides* midges.

Organism
Bluetongue virus (BTV):
- virulence may vary between serotypes and (in some cases) within serotypes
- is present in Australia (several serotypes) but clinical disease has not been reported in commercial Australian livestock.

Susceptible species
BTV affects all ruminant species, including goats, sheep, cattle, buffaloes, antelopes and deer (not all will develop clinical signs).

Economic impact
There is potential for significant economic loss associated with sheep production losses, as well as potential restrictions on international trade.

Epidemiology
- BTV distribution is determined by the presence of competent *Culicoides* vectors
- causes up to 100 per cent morbidity and 30–70 per cent mortality in sheep
- causes very low morbidity and mortality in cattle
- commonly causes subclinical infection in cattle and other species (for certain strains)
- has an incubation period of generally 2–8 days (but which may last as long as 20 days)
- viraemia is considered prolonged but not persistent
- has no recorded carrier state.

Transmission
Transmission occurs primarily through *Culicoides* vectors.

Clinical signs
Clinical signs of bluetongue disease vary in severity and are characterised by inflammation of mucous membranes, widespread haemorrhages and oedema. They include respiratory disease, lameness, oral lesions and nasal/ocular/oral discharges.

Post-mortem
Haemorrhages in the tunica media at the base of the pulmonary artery are regarded as being very characteristic of bluetongue.

Diagnosis and samples
Collect:
- serum and EDTA blood from affected and in contact live animals
- fresh spleen, heart and mesenteric lymph nodes from deceased animals.

Actions to take
If you suspect a case of bluetongue disease:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including movement controls and protection of animals from vectors) until advised by government veterinary authorities.
Introduction

Bluetongue disease is an arthropod-borne viral disease (arbovirus) of ruminants, which is characterised by inflammation of mucous membranes, widespread haemorrhages and oedema.

Disease agent and susceptible species

Bluetongue virus (BTV):

- is a non-enveloped double stranded RNA virus belonging to the *Orbivirus* genus of the *Reoviridae* family
- targets the epithelial cells of blood vessels and mucous membranes
- as of 2018 at least 27 serotypes have been identified worldwide of which 12 have been isolated in Australia
- affects all ruminant species, including sheep, goats, cattle, buffaloes, antelopes and deer
- affects sheep most severely whereas infection in cattle is generally subclinical, although this varies with the virus strain with which they are infected. However, cattle are of epidemiological importance as an amplifier of virus that can spill over into sheep.

Distribution

Bluetongue virus (BTV):

- occurs on all continents except Antarctica (New Zealand, Iceland and the Hawaiian Islands do not have *Culicoides* midges and are therefore free of BTV)
- caused outbreaks of disease across Europe and the United Kingdom (primarily associated with BTV-8) involving cattle, which reached as far north as Scandinavia in 2007.

For the latest information on the distribution of bluetongue, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int](http://www.oie.int).

Occurrences in Australia

As of 2018, 12 serotypes of the virus (BTV-1, 2, 3, 5, 7, 9, 12, 15, 16, 20, 21 and 23) have been detected in Australia. Experimental infections of a number of these serotypes show variable virulence in sheep, but clinical disease has not been reported in commercial livestock.

Only two serotypes, BTV-1 and BTV-21, are routinely detected along the east coast, with most serotypes geographically limited to the northern parts of the Northern Territory and Western Australia. This limitation in geographical distribution is thought to be due to vector-virus interactions.

Changes to the epidemiology of bluetongue in Australia can occur with the introduction of new bluetongue viruses or insect vectors into northern Australia, and/or spread of viruses capable of causing disease into sheep-raising districts in eastern/southern Australia (which would have significant implications for trade of livestock and livestock products).
Bluetongue virus (BTV):
- is present in northern and eastern parts of Australia, but is associated with subclinical infection due to the limited geographical distribution of virulent strains capable of causing disease in sheep
- the virus has not been detected in Victoria, South Australia or Tasmania
- distribution and prevalence varies from year to year according to climate and distribution and density of vector species
- distribution of viral transmission is monitored through the National Arbovirus Monitoring Program (NAMP) [https://www.animalhealthaustralia.com.au/what-we-do/disease-surveillance/national-arbovirus-monitoring-program/].

**Epidemiology**

**Modes of transmission**

BTV is not contagious and does not spread by direct contact between animals.

Transmission occurs via:
- **vectors**—certain species of *Culicoides* midges (of which there are over 1,400 species worldwide). Less than 1 per cent of species are capable of transmitting BTV. The primary vector species varies between regions.
- **Wind**, implicated in the dispersal of infected *Culicoides* in some outbreaks
- **transplacental transmission and transmission via ingesting placentas** from infected cows are possible, but not considered important in the epidemiology of the virus (BTV is not contagious)
- can be found in **semen**, but sexual transmission does not appear to be a major route of infection
- transmission of BTV-26 (and infection) through **direct contact** under experimental conditions is also documented.

**Disease dynamics**

BTV:
- incubation is generally 2–8 days (but incubation may last as long as 20 days)
- viraemia is prolonged but not persistent
- incubation and viraemia are both dependent on the strain of infecting virus
- immunity is prolonged but normally serotype specific, with animals exposed to one serotype generally remaining susceptible to others (limited cross-protection between specific serotypes can occur)
- vaccines are available but are serotype specific
- serotype is not itself a determinant for virulence, which varies not only between serotypes but also between strains within a given serotype, particularly those from different geographical locations.
Persistence of the agent

BTV does not persist in the environment, nor in animal carcasses or products. However, infected *Culicoides* vectors remain infected for life and can be carried over long distances by wind, enabling dispersal of virus to susceptible hosts. A carrier state is not observed in ruminants.

Diagnosis and pathology

Clinical signs

Disease results from extensive virus-mediated vascular injury that produces oedema, hyperaemia, haemorrhage and tissue infarction.

Morbidity and mortality are highly variable and dependent on the infecting virus strain and breed of sheep. In sheep, subacute and mild disease may be seen with varying combinations of clinical signs, including:

- acute reddening and petechial haemorrhages on the coronary band (animals may stand with arched backs and be reluctant to move due to pain associated with the coronary band)
- pyrexia
- general weakness and lethargy
- laboured breathing
- lameness due to coronitis
- hyperaemic mucous membranes
- nasal discharge (initially clear, later mucopurulent)
- oedema and cyanosis of the tongue (in severe cases) which give rise to the name 'bluetongue'
- oral erosions and ulcers
- petechial haemorrhages on the oral, nasal and conjunctival mucosae
- profuse haemorrhagic diarrhoea (in severe cases).

In the European outbreak of BTV-8 (2006–2007) clinical signs in cattle were marked and similar to those in sheep, although generally milder and with very low mortality. However, in cattle:

- most infections are subclinical
- dermatitis, oral and nasal lesions predominate
- a drop in milk production may be noted.

Infection in goats is usually subclinical. Disease presentation amongst wild ungulates varies between species but white-tailed deer appear to be the most susceptible of this group, with peracute, acute and chronic clinical disease described.

Pathology

Haemorrhages of 2–15 mm in diameter located in the tunica media at the base of the pulmonary artery are highly characteristic of bluetongue and:

- hyperaemia, oedema and haemorrhages may be widespread
- the most prominent gross lesions are found in and around the mouth
gastrointestinal mucosa may be oedematous, hyperaemic and possibly cyanotic, and petechial or ecchymotic haemorrhages may be present

• excoriations, which may be covered by grey necrotic material, may be found on the lips, dental pad, tongue and cheeks

• hyperaemia of the ruminal pillars and reticular folds is commonly found

• congestion, catarrhal inflammation and petechial haemorrhages maybe present in the small and large intestines

• lymph nodes and spleen may be moderately enlarged and haemorrhagic

• petechiae and pale areas of necrosis may be scattered through the skeletal musculature and myocardium

• catarrhal inflammation of the upper respiratory tract and oedema of the lungs may be present

• large quantities of plasma-like fluid may be seen in the thoracic cavity and pericardial sac.

**Differential diagnosis**

Consider in the differential diagnosis, depending on the species and predominant clinical signs:

• exotic diseases
  - [epizootic haemorrhagic disease](#) (virus is present in Australia but no clinical disease has been reported)
  - [foot-and-mouth disease](#)
  - [peste des petits ruminants](#)
  - [sheep and goat pox](#)
  - [vesicular stomatitis](#)

• endemic diseases
  - acute haemonchosis (with depression and submandibular oedema)
  - bovine parainfluenza 3 infection
  - bovine viral diarrhoea (BVDV1)
  - contagious pustular dermatitis (ecthyma, orf, scabby mouth)
  - footrot
  - infectious bovine rhinotracheitis
  - malignant catarrhal fever (bovine herpesvirus 2 is present in sheep Australia but clinical disease in cattle is rare)
  - pneumonia
  - **Oestrus ovis** infestation

• non-infectious diseases
  - photosensitisation
  - plant poisoning
  - other causes of lameness.
Samples required

Sample collection

Collect:

- **serum** from at least 10 live, clinically affected animals, and from exposed animals (particularly those that are convalescent)
- **EDTA blood** (7–10 ml/animal) from live, clinically affected animals
- **fresh tissue**, spleen, heart, mesenteric lymph node (2 g of each tissue) from recently dead affected animals.

Transport of samples

For transport:

- chill blood samples and unpreserved tissue samples to 4°C, or with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests

Confirm the diagnosis of bluetongue disease by laboratory testing, noting that:

- tests are available for detection of bluetongue include PCR assays that exploit the nucleic acid specificity of the viral genome and virus isolation
- the sequencing of segments of the genome can also be used to confirm serotype and regional genotypes
- serological assays include a group specific ELISA and virus neutralisation tests.

Reporting requirements

Infection with bluetongue is notifiable to the OIE and Australia has an international obligation to report cases. If you suspect bluetongue disease, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for bluetongue disease, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases (and protecting from vectors, if possible) and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.
Further information


FIGURE 3.5.1 Bilateral nasal exudate, erosion of the nasal planum, and excessive salivation in a sheep

Image credit: PIADC and CFSPH
FIGURE 3.5.2 Adherent crusting over a bovine muzzle, with underlying hyperaemic and eroded tissue

Image credit: PIADC and CFSPH

FIGURE 3.5.3 Erosion of most of the dental pad and necrosis of the remaining pale mucosa in an ovine mouth

Image credit: AFIP and CFSPH
FIGURE 3.5.4 Extensive coalescing ulceration of the teat skin of a bovine mammary gland

Image credit: PIADC and CFSPH

FIGURE 3.5.5 Multiple petechiae in the hoof wall, and marked hyperaemia of the coronary band of an ovine foot

Image credit: AFIP and CFSPH
CHAPTER 3.6

Bovine spongiform encephalopathy

Fast facts

Bovine spongiform encephalopathy (BSE) is due to a prion that results in a fatal neurodegenerative disease.

Clinical signs

Clinical signs in ruminants include:
- changes in mentation, behaviour and posture or gait
- neurological disease accompanied by weight loss and emaciation.

Agent

- BSE is caused by the accumulation of structurally-altered, stable and protease-resistant forms of prion protein (known as PrP\textsuperscript{\textsc{res}}).
- atypical BSE is caused by prions with either higher (H-type) or lower (L-type) molecular mass fragments than classical BSE
- prions are highly resistant to physical and chemical inactivation procedures.

Susceptible species

BSE affects:
- cattle
- humans (variant Creutzfeldt-Jacob disease, also known as vCJD).

Economic impact

There is potential for significant economic loss associated with restrictions on animal movements and international trade.

Epidemiology

- the incubation period is protracted, typically 2–8 years
- the disease therefore manifests in adult animals
- disease is always fatal.

Transmission

- natural transmission has not been observed with atypical forms of BSE
- consumption of BSE-infected central nervous system tissue has been implicated in vCJD in people, so a case of BSE has food safety implications.

Post-mortem

No gross lesions are detectable on post-mortem examination.

Samples

If safe to do so, and while wearing personal protective equipment, conduct a post-mortem and collect:
- whole brain fixed in 10 per cent neutral buffered formalin (obex, caudal cerebellar peduncles and midbrain samples)
- a 1–2 cm segment of fresh cervical spinal cord or caudal medulla

Actions to take

If you suspect a case of BSE:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

Public health considerations

- BSE is a zoonotic disease
- people may become infected after close contact with infected tissues
- consult the relevant state or territory health authorities as soon as BSE is strongly suspected or confirmed.
Introduction

Bovine spongiform encephalopathy (BSE) is a progressive neurodegenerative disease of adult cattle. The three strains identified in cattle include classical BSE and two strains of atypical BSE (H-type and L-type).

Disease agent and susceptible species

BSE is caused by the accumulation of structurally-altered, stable and protease-resistant forms (known as PrP$\text{\textsuperscript{TSE}}$) of prion protein in the CNS. In addition:

- classical BSE is caused by transmission of the BSE prion, whereas atypical BSE is thought to be a spontaneously occurring, non-contagious disease
- the two atypical strains—H-type BSE and L-type BSE—are caused by prions with either higher (H-type) or lower (L-type) molecular mass fragments than classical BSE
- BSE is principally a disease of domestic cattle, but other bovids including buffalo may be affected
- experimental infection has been induced in other ruminants (sheep and goats), mice, mink and non-human primates
- spongiform encephalopathies of various captive exotic species (wild ungulates and felids) are associated with ingestion of bovine carcasses or dietary supplementation of prion-contaminated meat-and-bone meal
- a spike in the incidence of acquired or variant Creutzfeldt-Jakob disease (vCJD) in humans occurred following the BSE epidemic in the United Kingdom. The consumption of food containing CNS tissue from BSE-affected cattle causes vCJD, a fatal neurodegenerative disease.

Public health considerations

- people may become infected after close contact with infected tissues
- if undertaking post-mortem examinations, wear personal protective equipment such as gloves and coveralls, rubber boots (or disposable boots), safety glasses and mask
- consult the relevant state or territory health authorities in the case of a BSE outbreak.

Distribution

BSE has been reported in the United Kingdom, Europe, Canada, Japan and Israel. For the latest information on the distribution of BSE, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

Classical and atypical BSE have not been detected in cattle in Australia.
Epidemiology

Modes of transmission
Infection requires exposure to CNS tissue from affected cattle. Transmission occurs via:

- **feeding of contaminated material**—the primary risk factor for BSE transmission is ingesting prion-contaminated feed. These products include: tonsils, distal ileum, brain, eyes, spinal cord, skull and vertebral column (which may be found in ruminant-derived meat-and-bone meal products, gelatin and tallow). For this reason, the feeding of ruminant-derived meat-and-bone meal to cattle is banned.

Disease dynamics
In countries other than the United Kingdom, BSE cases have originated from imported cattle or feeding young animals contaminated meat-and-bone meal. In addition:

- due to the protracted incubation of prion diseases, most cases of BSE in cattle occur in 3–8 year old animals
- the BSE in the United Kingdom was fundamentally a food-borne epidemic with BSE transmitted to young animals via feed contaminated with infectious prion. During the epidemic, most cattle were infected within the first 6 months of life and developed disease at 5–7 years of age.

Persistence of the agent
Prion strains are typically highly resistant to inactivation. In addition:

- PrP\textsuperscript{TSE} is resistant to physical and chemical inactivation procedures, including freezing, desiccation, ultraviolet radiation, degradation by various proteolytic enzymes, burial, and common methods of chemical and heat disinfection that destroy bacteria and viruses
- some organic acids can inactivate infectious prion proteins
- prions can adhere very strongly to stainless steel which may require special chemical disinfection.

Diagnosis and pathology

Clinical signs
The clinical course of BSE is protracted and slowly progressive over a period of weeks to months. Clinical signs of atypical and classical BSE are similar and:

- animals may have changes in mentation (fear, apprehension, aggression), sensation (hyperaesthesia), and posture and gait (ataxia, tremor, recumbency)
- non-specific signs such as weight loss and reduced milk yield may be present.

Pathology
No gross CNS lesions are present in cases of BSE.
Differential diagnosis

Consider BSE in cattle over 30 months of age presenting with neurological signs. Differential diagnoses include disorders that affect the neurological function and locomotion, which includes:

- **exotic diseases**
  - rabies

- **endemic diseases**
  - botulism
  - brain or spinal cord abscess
  - bovine herpesvirus
  - *Clostridium perfringens* type D (focal symmetrical encephalomalacia)
  - *Histophilus somni* encephalitis
  - *Listeria monocytogenes*
  - malignant catarrhal fever (sheep-associated malignant catarrhal fever is endemic)

- **non-infectious causes**
  - brain or spinal cord trauma
  - lead or plant toxicosis
  - metabolic disease (hypomagnesemia, hypocalcemia, polioencephalomalacia)
  - neoplasia
  - nutritional myopathy (white muscle disease).

Samples required

Sample collection

There is currently no validated diagnostic test available to detect BSE in live animals. Diagnostic tests require brain and spinal cord obtained at post-mortem examination. Note that you should:

- take stringent precautions during sampling, including wearing full personal protective equipment; where BSE is strongly suspected, direct efforts at collecting the brain and cervical spinal cord samples with minimal damage
- take care when disposing of or decontaminating equipment used for the post-mortem removal of brain tissue from suspected BSE cases
- be able to recover diagnostic material following mechanical injury to the brain (e.g. euthanasia by captive bolt).

For suspected cases of BSE, collect whole brain fixed in 10% neutral buffered formalin, and a 1–2 cm segment of unfixed (fresh) cervical spinal cord or caudal medulla.
Transport of samples

For transport:
- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests

Histologic examination is required to identify characteristic lesions within significant areas of the CNS, and to exclude differential diagnoses. In addition:
- immunohistochemistry (IHC) detects PrP\textsuperscript{TSE} aggregates in formalin-fixed brain sections and can be evaluated in conjunction with histological changes
- detection by Western blotting of PrP\textsuperscript{TSE} in unfixed tissue homogenates has similar diagnostic sensitivity to IHC
- rapid ELISAs are typically used as a screening strategy in surveillance programs.

Reporting requirements

BSE is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect BSE, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

BSE is a public health risk. Live cattle do not pose a risk, but human infections may result from contact with tissues of infected animals (as well as aerosols generated during slaughter or post-mortem examination). Post-mortem examinations of animals that have died from BSE pose a risk for animal-to-human transmission. Therefore, keep suspected cases of BSE isolated and on the property until advised by government veterinary authorities, noting you should:
- wear appropriate personal protective equipment (coveralls and boots, gloves, safety glasses or full face visor and protective respirator) when examining animals, performing post-mortem examinations, or handling tissue
- minimise the use of sharps and where possible use single-use disposable items (e.g. scalpels)
- use suitable hand protection such as armoured or cut resistant gloves, especially during collection of the brain or spinal cord
- extract central nervous system tissue using hacksaws and bone forceps to reduce the risk of exposure to bone dust contaminated with neural tissue (as would occur with oscillating saws)
- undertake personal decontamination when leaving infected premises. Thoroughly clean and then disinfect vehicles and equipment before leaving the property.
Further information


**FIGURE 3.6.1** The red box indicates the obex, which is the region of the brain that must be obtained for the diagnosis of BSE and other spongiform encephalopathies such as scrapie and chronic wasting disease.
Fast facts

Classical swine fever (CSF), also called hog cholera or swine fever, is a highly contagious, generalised exotic viral disease of pigs clinically indistinguishable from African swine fever.

Organism

CSF virus:
- replicates in blood and lymphatic tissues but is found in all tissues and body fluids
- has only one serotype
- is highly resistant to inactivation.

Susceptible species

Only pigs are susceptible to CSF.

Economic impact

There is potential for significant economic loss associated with high mortality and morbidity in pigs, as well as restrictions on animal movements and international trade.

Epidemiology

CSF:
- causes very high morbidity and up to 90 per cent mortality in the acute form
- has an incubation period of 2–6 days
- chronic or mild forms play a key role in disease spread through sub-clinically infected carriers
- piglets born to infected sows may also remain chronically infected, continuing to excrete the virus for several months before showing overt disease.

Transmission

Transmission occurs via:
- direct contact, usually oronasal
- contact with carcasses
- indirect transmission from ingestion, breeding (including artificial insemination) and fomites
- mechanical transmission by vectors.

Clinical signs

Affected animals may suffer from acute, chronic or subclinical disease and clinical signs vary according to the disease form (as influenced by the virus strain, route and dose of infection). Signs include:
- marked pyrexia, drowsiness, depression, anorexia and death (acute form)
- depression, anorexia, ill-thrift, fluctuating pyrexia and death after day 30 post-infection (chronic form).

Post-mortem

Pigs that die of acute disease may have few gross lesions, otherwise:
- lymph nodes are swollen, congested and haemorrhagic
- petechial or ecchymotic haemorrhages are found in many locations, including the renal cortex, larynx, trachea and small intestine.

Samples required

Also refer to the detailed section on sampling for sudden death in pigs.

Collect:
- blood samples (EDTA and plain tubes) from living animals
- fresh tissue from tonsils, spleen, lymph nodes, kidney, lung, liver, ileum and brain.

Actions to take

If you suspect a case of CSF:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Classical swine fever (CSF), also known as hog cholera or swine fever, is a highly contagious viral disease of pigs. CSF is clinically similar to African swine fever (ASF) but is caused by a different virus. The variation in clinical signs makes clinical identification difficult and definitive diagnosis requires laboratory testing.

Disease agent and susceptible species

CSF virus:
- is an enveloped RNA virus in the family Flaviviridae, genus Pestivirus
- has only one serotype but strains vary considerably in virulence and antigenicity
- affects domestic and feral pigs (Sus scrofa), which are the only susceptible species in Australia outside zoos.

Distribution

CSF:
- occurs in much of Asia, and in Central and South America, and parts of Europe and Africa
- was identified in the southern part of West Papua, Indonesia, in 2004
- has been eradicated from most of western Europe but a foci of infection persists in eastern Europe.

For the latest information on the distribution of CSF, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

Outbreaks occurred in Australia in 1903, 1927–28, 1942–43 and in 1960–61. In each case the disease was eradicated and Australia has been free of CSF since 1961.

Epidemiology

Modes of transmission

Transmission occurs via:
- **direct contact with infected pigs**—pigs with acute disease shed virus in high concentrations in all secretions and excretions. Respiratory infection can also occur between pigs in close contact. Moving infected pigs is the most important means of spread between piggeries. Heavily infected pigs excrete virus until they die and recovered animals excrete virus for 30 days. Feral pigs could become an important reservoir of the virus
- **ingesting contaminated swill**—CSF virus can survive for many months in raw, unprocessed, frozen meat as well as meat stored at fridge temperatures and in some processed hams. Ingesting pig meat or products containing the virus is an important means of spread. In Australia, we have strict import conditions to address the risk of a CSF incursion and swill feeding is illegal
- **fomites**—because of the stability of CSF virus in the environment, indirect contact by means of contaminated pig pens, vehicles and other fomites (such as hypodermic needles) is a very important means of spread of the disease
• contact with infected carcasses of pigs
• contact with infected faeces
• vectors—less commonly, bloodsucking insects feeding on viraemic pigs have been implicated in the mechanical spread of CSF
• semen and embryos—the virus is present in semen and can be transferred through breeding. Sows infected with low-virulence strains may continue to carry the virus in their foetuses because the virus can cross the placenta.

**Disease dynamics**

CSF virus:
• spreads in the nasal, lachrymal and saliva secretions of pigs as soon as 1 day after infection
• spread is possible during the incubation period (usually 2–6 days) and before the onset of clinical signs. Onset of shedding in faeces and urine occurs a few days later
• replication is detectable in the blood after 15–24 hours post-infection, which is concurrent with the occurrence of leukopenia and thrombocytopenia
• if infection occurs late in gestation, live piglets are occasionally born that remain chronically infected and continue to excrete the virus for several months before showing clinical disease.

Pigs that survive the acute disease or are infected by mild strains of the virus usually become chronically infected for at least 3–4 weeks post-infection. However, such pigs do not readily excrete the virus for more than about 5–6 weeks after initial infection and their role in the spread of the disease after that is unclear.

The chronic and subclinical forms usually last three months and consist of three distinguished phases:
• the early phase (2–3 weeks post-infection) is similar to the acute form of CSF
• the second phase (3–6 weeks post-infection) is characterised by convalescence with an increased neutralising antibody response and decreased levels of virus in the blood. Virus shedding is not affected by these changes
• the third phase is characterised by the reappearance of clinical signs with increased viral replication and death.

**Persistence of the agent**

CSF virus:
• can persist in contaminated pig pens and on other fomites for up to 2 weeks, depending on environmental conditions and the amount of infectious virus present
• is susceptible to ether and other lipid solvents, and inactivated by chlorine-based disinfectant. Sodium hydroxide (2%) is considered suitable for disinfecting premises.

**Diagnosis and pathology**

**Clinical signs**

CSF is an extremely variable disease and you cannot diagnose it based on clinical signs and gross pathology alone. Disease may be acute, subclinical or chronic.
In the acute form most pigs die between 10–20 days post-infection. Clinical signs for the acute form include:

- pyrexia (above 39.5°C, and as high as 42°C)
- hyperaemia or cyanosis of extremities, particularly ears and snout, which can present as red or purple skin blotching on the ears, snout limbs and abdomen
- loss of appetite or irregular appetite
- inability or unwillingness to stand up, convulsions
- incoordination, stiff gait
- huddling together, piling one on top of another
- laboured breathing, coughing
- dysentery or diarrhoea
- conjunctivitis
- nasal discharge
- vomiting
- abortion, mummifications, stillbirth and foetal abnormalities
- severe leucopenia
- death (case fatality rate up to 100 per cent).

In the subclinical disease form, pigs may become chronic carriers without showing any of the clinical signs listed. This is most likely to occur when strains of low virulence infect older breeding animals or when piglets are infected in utero.

In the chronic disease form, clinical signs are as for the acute form but generally milder and persisting longer (3–4 weeks). They include:

- pyrexia which may fluctuate irregularly
- ill-thrift
- pneumonia
- loss of appetite
- diarrhoea
- alopecia and dermatitis
- death—often due to secondary bacterial infections
- lower case fatality rate than the acute form.

**Pathology**

With the acute disease form, findings are:

- enlarged and haemorrhagic lymph nodes, often resembling blood clots (the gastrohepatic, renal, mesenteric and submandibular lymph nodes are most often affected)
- pinpoint haemorrhages on the tonsils (tonsils are frequently enlarged, with necrotic foci and pustules)
- pyramidal splenic infarcts along the margin
- haemorrhages in almost any organ (most commonly on serosal membranes and in kidneys as subcapsular petechiae or ‘turkey egg’ appearance, heart, urinary bladder, epiglottis, lung and gall bladder)
- septal oedema of lungs
- fluid in body cavities.
Findings for the chronic form are more variable than for the acute form, as they are often complicated by secondary bacterial infections. They include:

- lymph node and renal haemorrhage
- mucosal intestinal haemorrhage
- enlarged lymph nodes
- thymic atrophy
- fibrinous pericarditis and pleurisy
- lobular consolidation of lungs which may progress to lobular necrosis and bronchopneumonia
- poor body condition
- ulceration of the large intestine
- button ulcers, particularly near the ileocaecal valve.

**Differential diagnosis**

Many other viral and bacterial pathogens of pigs often confused with CSF may cause concomitant infections. It is important that you take into consideration that detection of other pathogens may mask an underlying CSF virus infection.

Consider in the differential diagnosis:

- **exotic diseases**
  - *African swine fever* (do not attempt clinical differentiation of CSF and ASF)
  - *Aujeszky’s disease* (respiratory signs)
  - *porcine reproductive and respiratory syndrome*
  - *porcine deltacoronavirus*
  - *porcine respiratory disease complex*
  - *post-weaning multi-systemic wasting syndrome*
  - swine enteric coronavirus diseases such as those caused by *porcine epidemic diarrhoea* virus
  - *transmissible gastroenteritis*

- **endemic diseases**
  - acute salmonellosis (diarrhoea)
  - acute septicaemia due to *Streptococcus suis* or *Haemophilus parasuis* (Glässer’s disease)
  - border disease virus infection
  - bovine viral diarrhoea virus infection
  - *erysipelas* (haemorrhagic skin lesions)
  - *pasteurellosis* (pneumonia)
  - *porcine circovirus-associated* diseases (such as those seen in *porcine dermatitis and nephropathy syndrome*)

- **non-infectious causes**
  - heavy metal poisoning
  - salt poisoning with water deprivation
  - warfarin poisoning (causing death with haemorrhages, generally sporadic)
  - thrombocytopenia purpura (haemorrhages, especially in pigs 2–3 weeks old)
  - any cause of chronic ill-thrift in pigs.
Samples required
Submit at least five pigs for post-mortem examination, as individual animals may present with great variability in lesions. Record a composite picture of all lesions seen.

Sample collection
Collect samples from affected, especially pyrexic, pigs killed immediately before a post-mortem examination and from pigs that have recently died (including stillborn piglets and aborted foetuses). Collect:
- **serum**, 30 samples from suspected chronically infected animals
- **EDTA blood** (7–10 ml/animal) from live, clinically affected animals
- **fresh tissue** from the spleen, gastro-hepatic lymph node, mesenteric lymph node, tonsils, lung, kidney and ileum (2 g of each tissue)
- **fixed tissue**, a full range of tissues (including the brain) in neutral-buffered formalin.

Transport of samples
- chill blood samples and unpreserved tissue samples either at 4°C or with frozen gel packs
- DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
The most rapid, sensitive and specific diagnostic procedure is the detection of viral nucleic acid in blood or in lymphoid tissue by qRT-PCR. A positive result can be obtained within one day of pyrexia first developing in an animal. In addition:
- commercial ELISA kits are available for detection of CSF virus antigen and antibody but are comparatively less sensitive than molecular techniques
- specific antibodies against CSF virus may be slow to appear due to the immunosuppressive nature of the disease
- viral antigen can be detected in frozen sections of tonsil, spleen or lymph node using direct immunofluorescence, and may yield a result in a matter of hours
- isolation of the virus is attempted by inoculation of blood or tissue suspensions into porcine kidney cell cultures
- because most strains of CSF virus do not produce recognisable cytopathic effects, viral growth is identified by immunolabelling methods such as fluorescent antibody or immunoperoxidase tests, or by qRT-PCR
- antibodies to border disease virus or bovine viral diarrhoea virus in pigs may confuse serological surveys for CSF. Cross-neutralisation tests for these and other relevant circulating pestiviruses need to be carried out to confirm CSF antibody.
Reporting requirements

CSF is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect CSF, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for CSF, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


FIGURE 3.7.1 Multiple coalescing, swollen, dark red infarcts along the margins of a spleen

Image credit: PIADC and CFSPH
FIGURE 3.7.2 Multiple petechiae and pale infarcts surrounded by haemorrhage in the cortex of a kidney

![Image of a kidney with petechiae and infarcts](image1)

Image credit: PIADC and CFSPH

FIGURE 3.7.3 Reddened mucosa that contains multiple discrete (button) ulcers surrounded by zones of haemorrhage in the colon

![Image of colon with ulcers](image2)

Image credit: PIADC and CFSPH
Chapter 3.8
Epizootic haemorrhagic disease

Fast facts

Epizootic haemorrhagic disease (EHD) is a vector-borne viral disease. Historically a disease of wild ruminants, EHD has become an emerging disease in cattle.

Organism

EHD virus (EHDV):
- has 7 recognised serotypes (1–2 and 4–8) of which 6 occur in Australia (but no clinical disease has been reported)
- has a strain called Ibaraki virus, which causes the Ibaraki disease of cattle in eastern Asia.

Susceptible species

EHD affects:
- most cervids but clinical disease is not reported in deer species introduced to Australia
- various wild and domestic ruminant species, which are susceptible to sub-clinical infection.

Economic impact

There is potential for significant economic loss associated with production losses and mortality in cattle or farmed deer, and restrictions on animal movement and international trade.

Epidemiology

Infection is most common during peak activity periods of Culicoides vectors. Persistent ruminant infection does not occur.

Transmission

The non-contagious virus is transmitted between ruminant hosts by haematophagous species of Culicoides biting midges.

Clinical signs

Most infections in cattle are subclinical. Peracute to chronic illness occurs in deer and clinical signs may include: pyrexia, severe and rapid oedema of the head and neck, hypersalivation, erosive and ulcerative lesions of the oral and oesophageal mucosa, weakness and respiratory distress.

Post-mortem

Post-mortem lesions are characteristic of systemic haemorrhagic disease, with petechiae, ecchymoses and haemorrhages in various mucosal and serosal surfaces, organs and tissues.

Samples

Collect:
- whole blood in EDTA tubes
- fresh spleen, lung and lymph nodes samples
- paired serum samples.

Actions to take

If you suspect a case of EHD:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including movement controls and protection of animals from vectors) until advised by government veterinary authorities.
Introduction

Epizootic haemorrhagic disease (EHD) is a vector-borne, non-contagious viral disease transmitted by *Culicoides* spp (haematophagous *Culicoides* midges). EHD causes a severe haemorrhagic disease of wild cervids in North America but is also recognised as an emerging disease in cattle overseas.

Disease agent and susceptible species

The EHD virus (EHDV):
- is a non-encapsulated, double-stranded RNA virus and a member of the *Orbivirus* genus of the family *Reoviridae*
- serogroup contains 7 recognised serotypes (numbered 1–2 and 4–8), 6 of which occur in Australia (serotypes 1, 2, 5, 6, 7, 8) but no clinical disease has been reported
- shares morphologic and structural similarities to other members of the *Orbivirus* genus (there are also epidemiological parallels with bluetongue virus through shared patterns of *Culicoides* spp. transmission) and shows immunological cross reactivity with bluetongue virus
- various wild and domestic ruminant species (including water buffalo) are susceptible to sub-clinical infection. In addition:
  - most cervids are susceptible to infection but clinical disease is not known to occur in deer species in Australia
  - sheep can be infected experimentally but rarely develop clinical signs, whereas goats may be susceptible to EHDV infection but their role as a host is uncertain
  - the Ibaraki strain of EHDV serotype 2 causes Ibaraki disease in cattle in eastern Asia. This is not present in Australia.

Distribution

EHDV is found in Australia, Africa, North and South America, Asia and the Middle East.

Outbreaks of EHD in cattle have been reported in Japan, South Korea, Taiwan (Ibaraki disease syndrome), the United States, North Africa and countries bordering the European Union. In addition:
- there were few descriptions of EHD in cattle outside Asia until 2004, when outbreaks were reported on Reunion Island (located near Madagascar), followed by outbreaks in the Middle East in 2006, the Caribbean in 2011, and the United States in 2012
- cattle were the only domesticated animals affected during the outbreaks in the Caribbean and the Middle East
- bison, yaks, elk and sheep developed clinical signs during the 2012 outbreaks in the United States. The case fatality rates in most species are unknown, but deaths were reported in bison and yaks.

For the latest information on the distribution of EHD, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].
Occurrences in Australia

In Australia:

- naturally occurring disease has not been observed in cattle, deer or other susceptible species
- though there have been no reports of clinical disease, the virus is regularly isolated in the Northern Territory and Queensland under the National Arbovirus Monitoring Program (NAMP)
- the distribution of EHDV is presumed to mirror bluetongue virus (BTV), so regions in Western Australia, Northern Territory, Queensland and New South Wales defined as BTV transmission zones by NAMP are regarded as zones of potential transmission for EHDV. It is assumed that incursions from overseas may occur in northern Australia due to long distance movement of Culicoides vectors on the wind, as is demonstrated for BTV. This may explain the greater diversity of EHDV serotypes (1, 2, 5, 6, 7 and 8) detected in the Northern Territory, while only serotypes 2, 5, 6 and 7 have been detected in Queensland.

Epidemiology

Modes of transmission

EHD is not contagious and is not transmitted by direct contact between animals. Transmission occurs via:

- **vectors**—haematophagous Culicoides spp midges transmit the virus between ruminant hosts and disease transmission patterns are typically seasonal, corresponding with vector distribution and abundance (cattle may serve as temporary reservoir hosts)
- **wind**—implicated in the dispersal of infected Culicoides in some outbreaks
- **reproductive material**—being closely related to bluetongue virus, it is likely that the semen of viraemic bulls may sometimes contain EHDV.

Disease dynamics

EHD:

- virus replicates in the endothelial cells of lymphatic vessels
- in deer the disease is characterised by an acute haemorrhagic disorder with a progressive thrombocytopenia
- viral incubation period lasts 5–10 days in deer and 2–10 days in cattle
- clinical signs develop in white-tailed deer (Odocoileus virginianus) about 7 days after exposure
- viraemia duration differs for each Australian EHDV serotype but typically ranges from 2–4 weeks.

Persistence of the agent

The virus:

- is stable in blood and tissue samples but sensitive to pH <6.0 and >8.0
- can be readily inactivated by acids and alkalis, sodium hypochlorite, 2% weight per volume glutaraldehyde, acids, iodofores and phenolic compounds (but like other non-enveloped RNA virus it is resistant to lipid solvents such as ether and chloroform and resistant to ultraviolet and gamma irradiation)
- can also be inactivated at temperatures of 50°C for 3 hours, 60°C for 15 minutes and 121°C for 15 minutes.
Diagnosis and pathology

Clinical signs
In deer, peracute to chronic illness occurs and clinical signs may include:

- pyrexia, severe and rapid oedema of the head and neck, hypersalivation, erosive and ulcerative lesions of the oral and oesophageal mucosa, weakness and respiratory distress
- anorexia, muscle weakness, lameness, tachypnea and tachycardia, and haemorrhagic nasal discharge.

Experimental infection of cattle with Australian EHDV strains has not resulted in clinical signs; however, these signs have been recorded overseas:

- pyrexia
- anorexia and hypersalivation
- dehydration and emaciation
- lameness associated with coronitis
- erosive and ulcerative lesions of the oral and oesophageal mucosa
- dysphagia is considered pathognomonic for Ibaraki disease (associated with striated muscle damage to the pharynx, larynx, tongue and oesophagus)
- abortions and stillbirths (foetal hydranencephaly).

Pathology
Post-mortem lesions are characteristic of systemic haemorrhagic disease, with petechiae, ecchymoses and haemorrhages in various mucosal and serosal surfaces, organs and tissues. In deer, you may see:

- extensive haemorrhages on the serosal surfaces of the intestine and rumen
- erosions and ulcerations in the mouth, rumen, and omasum
- dry, gray-white necrotic lesions of the hard palate, tongue, dental pads, esophagus, larynx, rumen, and abomasum
- sloughing of the tips or walls of the hooves
- lungs may be oedematous and cyanotic.

In cattle, you may see:

- degeneration of the striated muscles in the oesophagus, larynx, pharynx, tongue, and skeletal muscles (pathognomonic lesions for Ibaraki disease)
- marked oedema and haemorrhages in the mouth and lips, abomasum, and around the coronets
- erosions or ulcerations in the oral cavity, rumen and omasum
- emaciated and dehydrated carcass.
Differential diagnosis

Depending on the species and predominant clinical signs, Consider in the differential diagnosis:

- **exotic diseases**
  - bluetongue disease (infection present in Australia but without clinical disease)
  - foot-and-mouth disease
  - vesicular stomatitis
- **endemic diseases**
  - bovine viral diarrhoea (BVDV-1)
  - bovine ephemeral fever
  - infectious bovine rhinotracheitis
  - malignant catarrhal fever (sheep-associated viruses are present in Australia but clinical disease in cattle is rare)
- **non-infectious diseases**
  - photosensitisation
  - plant poisoning
  - other causes of lameness.

Samples required

Sample collection

Collect:

- **serum** from at least 10 live, clinically affected animals, and from exposed animals (particularly those that are convalescent)
- **paired serum samples** if possible
- **EDTA blood** (10 ml/animal) from live, clinically affected animals
- **other samples**, fresh spleen, lung, and lymph nodes (2 g of each tissue) from recently dead affected animals.

Transport of samples

For transport:

- chill blood samples at 4°C with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.
Diagnostic tests

EHDV can readily be detected by direct detection of genome with molecular assays and isolation of virus or indirectly through the detection of group or type-specific antibodies. The diagnostic tests available in Australia include virus isolation, virus neutralisation, agarose gel immunodiffusion and ELISA.

Reporting requirements

EHD is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect EHD, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for EHD, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases (and protecting from vectors, if possible) and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


FIGURE 3.8.1 Lungs of a deer with moderate to marked swelling of the interlobular septa due to oedema and a few small pleural haemorrhages. There is diffuse congestion of the tracheal mucosa.

Image credit: Dr E Howerth, University of Georgia, Department of Veterinary Pathology, and CFSPH
Equine influenza

Fast facts
Equine influenza (EI) is an acute, highly contagious exotic viral disease which can cause respiratory disease in horses and other equids.

Organism
EI virus:
• has two subtypes that infect horses—H3N8 and H7N7
• inactivated by ultraviolet light and most disinfectants.

Susceptible species
EI affects:
• horses and other equids
• dogs (this chapter will only address infection in horses).

Economic impact
There is potential for significant economic loss associated with high morbidity if the disease became established, as well as potential restrictions on international trade. The 2007 Australian outbreak cost an estimated AUD$1 billion.

Epidemiology
EI:
• has an incubation period of 18 hours to 5 days
• infected animals can shed virus for 7–10 days.
• causes very high morbidity but low mortality

Transmission
Transmission occurs via:
• direct contact with infected animals or fomites
• droplets (droplets >10 µm)
• aerosol (droplets <5 µm).

Clinical signs
Clinical signs include:
• pyrexia (up to 42°C)
• serous nasal discharge
• dry cough.

Post-mortem
Since the disease causes very low mortality post-mortem is unlikely.

Samples
Collect:
• blood samples (EDTA and plain tubes)
• swabs (nasopharyngeal or nasal) or nasal wash samples in adults
• nasal swabs from foals.

Actions to take
If you suspect a case of EI:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Equine influenza (EI) is a highly contagious disease equids causing upper respiratory tract infection.

Disease agent and susceptible species

*Equine influenza virus* (EIV):
- is a member of the *Influenza virus type A* genus, in the *Orthomyxoviridae* family
- is not known to infect humans
- infects horses and other equids and transmission to dogs has been reported (this chapter will only address infection in horses)
- is more stable than human influenza viruses; however, point mutations in the haemagglutinin and neuraminidase genes can cause significant changes in the antigenicity of the virus
- has two subtypes that infect horses—H3N8 and H7N7—and both are believed to have evolved from avian influenza virus ancestors. The H3N8 subtype circulates widely whereas the H7N7 subtype was last detected in 1979.

Distribution

EI:
- outbreaks have been reported all over the world with the exception of a small number of island nations including New Zealand and Iceland
- is endemic in Europe (except Iceland), North and South America, North Africa and Asia.

For the latest information on the distribution of EI, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int](http://www.oie.int).

Occurrences in Australia

Australia had its first outbreak in 2007, but the disease was eradicated and Australia was declared free of EI in 2008.

Epidemiology

Modes of transmission

Transmission occurs via:
- **direct contact with infected horses**—respiratory infection can occur between horses in close contact, and outbreaks of disease occur most often when you congregate susceptible animals and house them in close contact with each other
- **droplet transmission**—droplets greater than 10 µm are projected over moderate distances by coughing and sneezing
- **airborne transmission**—droplet nuclei are greater than 5 µm and capable of wide dissemination, and reaching the lower respiratory tract. Windborne spread of several kilometers has been reported anecdotally; however, evidence supporting this is inconclusive and alternative transmission routes (such as by fomites) play a larger role in the epidemiology
• **fomites**—such as clothing, horse equipment, people, buildings that have recently housed sick horses, vehicles, floats, grooming and veterinary instruments. Depending on environmental conditions (humidity, temperature, exposure to sun light), the virus can remain infectious for days on contaminated surfaces and objects.

**Disease dynamics**

**EI:**
- may occur at any time of the year
- infects horses of all ages, though disease is generally mild in young foals
- incubation period can range from 18 hours to 5 days, dependent on level of exposure
- live virus shedding is variable and can persist for 7–10 days, and viral RNA is detectable by PCR for up to 15 days.

**Persistence of the agent**

**EI:**
- virus survival depends on environmental conditions and the amount of infectious virus present
- virus can persist for up to 48 hours on hard non-porous surfaces (such as stainless steel and plastic) and 12 hours on porous surfaces (such as clothing and paper)
- virus can survive in tap and canal water for up to 2 weeks
- virus is inactivated by exposure to ultraviolet light for 30 minutes, by heating at 50°C for 30 minutes, by acid (pH 3) treatment and exposure to sunlight for 15 minutes at 15°C also inactivates the virus
- virus is inactivated by most disinfectants (including citric acid, and Virkon®) suitable for disinfecting surfaces and equipment (providing organic material has been first removed)
- in endemic countries, vaccinated horses can represent an important source of infection because of the occurrence of subclinical disease with viral shedding.

**Diagnosis and pathology**

**Clinical signs**

Signs include:
- rapid onset pyrexia (up to 42°C)
- serous nasal discharge
- a persistent, harsh, dry cough
- mild swelling of the submandibular lymph nodes
- dull attitude
- loss of appetite
- fatigue
- myalgia
- limb oedema
- vaccination can reduce the incidence, severity and duration of clinical signs.
Pathology
Gross and microscopic lesions are not highly specific. There may be hyperaemia or inflammation of the mucosa of the upper respiratory tract.

Differential diagnosis
Consider in the differential diagnosis:
• endemic diseases
  – equine adenovirus
  – equine rhinitis
  – equine rhinopneumonitis (equine herpesvirus 1 and 4 infection)
  – equine viral arteritis (infection present with no clinical disease)
  – Hendra virus infection
  – infection with *Streptococcus zooepidemicus, Streptococcus pneumoniae*, *Streptococcus zooepidemicus* subspecies *equi* (strangles)
  – parasitic infections, including ascarids and lungworms
  – pasteurellosis
• non-infectious causes
  – inflammatory airway disease due to exposure to environmental irritants and allergens.

Samples required
Sample collection
Collect:
• EDTA blood, from live, clinically affected animals (7–10 ml/animal)
• swab samples (nasopharyngeal or nasal swabs in adults; nasal swabs in foals under 6 months of age) or nasal washes containing both mucosal cells and nasopharyngeal secretions, as soon as possible after onset of clinical disease (preferably within 24–48 hours).

Transport of samples
For transport:
• swab samples should be placed in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
• chill blood samples and unpreserved tissue samples at 4°C, or with frozen gel packs
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.
Diagnostic tests

For EI:

- the most rapid, sensitive and specific diagnostic procedure is the detection of viral nucleic acid in blood or nasal secretions by qRT-PCR
- serology is available for the determination of acute and convalescent antibody titers using an ELISA and/or haemagglutination inhibition tests
- isolation of the virus is attempted by inoculation of nasal or nasopharyngeal swab samples into cell cultures (typically Madin-Darby canine kidney cells and embryo-nated chicken eggs).

Reporting requirements

EI is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect EI, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for EI, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


CHAPTER 3.10
Foot-and-mouth disease

Fast facts
Foot-and-mouth disease (FMD) is a highly contagious, exotic viral vesicular disease of cloven-hoofed animals.

Organism
FMD virus (FMDV):
• primarily replicates in epithelial cells in the pharynx and dorsal soft palate and then spreads via the blood to secondary sites
• can remain infective in the environment for several weeks and possibly longer in the presence of organic matter
• is inactivated by high and low pH and many disinfectants.

Susceptible species
All cloven-hoofed animals are susceptible, including cattle, sheep, pigs, goats and deer. Of domestic livestock, the disease is most severe in cattle and pigs.

Economic impact
There is potential for significant economic loss associated with eradication, production losses and restrictions on international trade.

Epidemiology
FMD:
• has an incubation period of 2–14 days, depending on the dose and route of exposure
• causes morbidity of up to 100 per cent in domestic livestock
• causes mortality in <5 per cent adult animals, higher in young animals
• virus can persistently infect a number of animals for varying periods.

Transmission
Transmission occurs via:
• aerosol transmission
• direct contact, usually oronasal

• swill feeding
• fomites
• artificial insemination.

Clinical signs
Clinical signs include:
• vesicles and ulcers in the mouth, feet and teats
• lameness
• pyrexia
• unwillingness to eat
• excessive salivation
• abortion
• drop in milk production
• sudden death in young animals.

Post-mortem
At post-mortem, vesicles may be present in ruminal pillars and ‘tiger heart’ striping of cardiac muscle in young animals.

Samples
Also refer to the detailed section on sampling for animals with vesicular disease.
• blood (EDTA and plain tubes)
• vesicular fluid, oropharyngeal fluid, oral, nasal and tonsillar swabs, epithelium, and epithelial tags (in buffer or virus transport medium).

Actions to take
If you suspect a case of FMD:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Foot-and-mouth disease (FMD) is a highly contagious viral vesicular disease of cloven-hoofed animals. The most significant risk for entry of FMD into Australia is through illegal importation of meat and dairy products from infected animals fed to pigs as swill (swill feeding).

Disease agent and susceptible species

FMD virus (FMDV):

- is a member of the genus *Aphthovirus* within the family *Picornaviridae*
- is a single-stranded RNA virus with no envelope
- is differentiated into seven serotypes (A, O, C, Asia 1, SAT 1, SAT 2 and SAT 3) by various serological tests. There is no cross-protection between serotypes. Serotypes have been further classified into genotypes and topotypes based on genetic similarity and their geographical origin and occurrence
- has numerous antigenic variants and studies are required to assist with the selection of the most appropriate vaccine strains
- can infect all wild and domestic cloven-hoofed animals (of the domestic livestock species, cattle, buffalo, pigs, sheep, goats and deer are susceptible to FMD, with the disease most severe in cattle and pigs)
- has some strains which may affect Bactrian camels and camelids, and the Asian elephant
- infection of humans has been reported but is extremely rare, and infections were minor and self-limiting. FMD is not a public health concern.

Distribution

FMD:

- is endemic in the Middle East, South Asia, South-East Asia, China, central Asian countries and most parts of Africa (but the serotype distribution differs in these regions)
- was controlled or eradicated with the use of mass vaccination programs in South America, and most South American countries are free of the disease
- incursions into previously free countries have had severe economic impacts, for example, in the United Kingdom, Japan and the South Korea.

For the latest information on the distribution of FMD, please refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

FMD has not occurred in Australia since the late 1800s when minor outbreaks are thought to have occurred in 1801, 1804, 1871 and 1872. The 1872 incident occurred in Victoria where the disease was introduced with an imported bull, and involved two farms before being eradicated.
Epidemiology

FMD is one of the most contagious viral animal diseases as virus is excreted in high quantities in expired air, in all secretions and excretions (including saliva, milk and semen), and from ruptured vesicles. Note that:

- pigs are regarded as important amplifying hosts for the disease, because of their capacity to excrete large quantities of virus in their exhaled breath
- cattle are regarded as indicator hosts because of their susceptibility to infection
- sheep and goats are considered silent or maintenance hosts, because the infection can spread through flocks with limited clinical disease.

Modes of transmission

Transmission occurs via:

- **direct contact between infected and susceptible animals**—the main route of infection is via the oronasal route; virus can also enter through breaks in the skin or mucosae.
- Stocking density is a determinant of the rate of spread of disease within an infected premises, with higher stocking densities seeing faster rate of spread. Movement of infected animals enables rapid dissemination of the virus over wide geographical areas
- **mechanical spread** by people, rodents, insects and birds
- **contact with the carcasses** of infected animals
- **contact with secretions**
- **semen and embryos** containing virus
- **ingestion of infected swill**—pigs have a higher relative susceptibility to infection through ingestion of contaminated feed, and swill feeding is illegal in Australia. Transmission occurs through the feeding of infected animal products including meat scraps and bones, or untreated raw milk and milk products from infected animals
- **fomites**—indirect contact by means of contaminated animal feed, straw, water, vehicles (including milk tankers) and other fomites (such as hypodermic needles) can spread the disease
- **airborne spread**—cattle and sheep are most susceptible to infection by inhalation of contaminated aerosols and extremely small doses of virus can initiate infection. Airborne spread can occur over long distances. The pattern of airborne spread has generally been from pigs as source to cattle downwind, and is likely to occur only when there are high concentrations of the appropriate livestock species at these locations. Long distance spread only occurs under very specific climatic conditions.

The transmission of FMD tends to be different in tropical and semitropical regions from that in temperate regions. In tropical areas, reasonably close contact is required between infected and susceptible animals for successful transmission, probably because of faster inactivation of the virus in hotter environments. The epidemiology of the disease in northern Australia could thus be expected to be different from that in southern Australia.

Healthy people can harbour FMDV subclinically in the nasal passages and throat for up to 28 hours, which means that during FMD outbreaks responders should avoid moving between premises for a directed amount of time (to avoid the possibility of transmitting FMDV to susceptible, uninfected animals).
Disease dynamics

FMDV is considered highly contagious, but mortality is low. In addition:

- the incubation period is variable depending on the virus strain, exposure dose, route of entry and species infected
- the incubation period for clinical signs is 1–14 days, but most commonly is 2–5 days
- excretion of FMDV can begin up to 4 days before clinical signs become apparent
- excretion of the virus decreases about 4–6 days after the appearance of vesicles, when circulating antibodies appear
- excretion of virus from foot lesions tends to last a day or two longer than from mouth lesions, so foot lesions may be a better source of virus for diagnostic purposes in older cases
- FMDV has been detected in the milk and semen of experimentally infected cattle for 23 and 56 days post-infection, respectively
- following infection with FMDV, it is possible for ruminants (but not pigs) to become persistently infected when virus persists in the pharynx in the presence of circulating antibody. Despite a number of anecdotal reports, as yet there is no evidence from the field that persistently infected animals (other than African buffalo) have been responsible for initiating new infections in susceptible animals
- vaccinated animals may become infected even when they are fully protected against clinical disease. Such animals are capable of excreting virus for about 1 week, though at lower levels. A proportion of vaccinated infected ruminants may also become persistently infected.

Persistence of the agent

FMDV:

- is most stable at pH 7.2–7.6 but will survive at pH 6.7–9.5 if the temperature is reduced to 4°C or lower
- has a half-life of approximately 12 hours at pH 6.5, 1 minute at pH 6, and 1 second at pH 5 (although inactivation times depend on many factors)
- can be inactivated with acidic solutions (e.g. acetic acid, straight vinegar or citric acid solutions), alkaline solutions (sodium hydroxide or sodium carbonate) or the disinfectant Virkon®
- survival time is reduced by higher temperatures and most strains of FMDV are inactivated when placed at 56 °C for 60 minutes. The effect of temperature on viral infectivity is influenced by the presence of organic material
- remains stable for prolonged periods at temperatures below freezing
- is largely unaffected by sunlight
- can remain infective in the environment for several weeks, and possibly longer in the presence of organic matter (such as soil, manure and dried animal secretions) or on chemically inert materials such as straw, hair and leather.

Diagnosis and pathology

The classical signs and lesions of FMD are described below. Note that a wide range of clinical syndromes can occur, ranging from inapparent disease with minimal lesions to severe clinical disease.
Clinical signs

Morbidity is close to 100 per cent in fully susceptible cloven-hoofed domestic animals but variable in wildlife species. Mortality due to FMD virus infection is very low in adult animals (<5 per cent) but is higher in young calves, lambs and piglets (20 per cent or higher).

The course of FMD in cattle may last for 2–3 weeks, or longer if there are serious secondary bacterial infections. Long-term sequelae may include hoof deformities and permanent damage to the udder. Clinical signs in cattle include:

- pyrexia accompanied by severe depression, inappetence and (in milking animals) a sudden drop or cessation of milk production
- followed (within 1–2 days) by the appearance of vesicles, the predilection sites for which are the tongue, lips, gums, dental pad, interdigital skin of the feet, coronary bands, bulbs of the heels and teats. Note that vesicles can burst readily, within 24 hours, so it is more common to see ulcers
- occasionally, vesicles and ulcers appear inside the nostrils or on the muzzle or vulva (see Table 3.10.1 for description of lesions)
- in the mouth, vesicles and ulcers are particularly prominent on the tongue, dental pad and cheeks. In severe cases, most of the dorsal surface of the tongue may slough. In uncomplicated cases, mouth lesions heal fairly rapidly over a 7–10 day period and eating may resume within a few days of rupture of vesicles
- painful stomatitis (associated with unruptured and freshly ruptured vesicles), which causes excess salivation, lip smacking and cessation of eating, leading to rapid loss of body condition
- foot lesions, which are accompanied by acute lameness and reluctance to move (secondary infections may lead to severe involvement of the deeper structures of the foot)
- teat lesions, which may lead to secondary mastitis
- abortion
- infection of very young calves may cause sudden death due to cardiac involvement, without vesicular lesions (mortality can be as high as 50 per cent).

Clinical signs in pigs include:

- initially, pyrexia (>40°C), inappetence and lameness or reluctance to move
- vesicles and ulcers (within 1–2 days of disease onset)
- vesicles and ulcers are most pronounced on the feet and result in acute lameness, pain and recumbency, particularly if the pigs are housed on a hard floor (the disease may be difficult to detect when affected pigs are housed on soft bedding)
- vesicles and ulcers may occur on the coronets, interdigital skin, or bulbs of the heel, or other areas of friction
- vesicles and ulcers that encircle the coronet may lead to separation of the keratinised layers of the hoof from the corium—in severe cases there may be sloughing of the hoof; otherwise a line of separation between old and new horn moves steadily down the hoof at a rate of about 1 mm per week, starting a week after rupture of coronary band vesicles (the age of FMD lesions in pigs can often be estimated in this way)
- vesicles occur on the snout, but rupture quickly
- vesicles and ulcers on the tongue are relatively uncommon in pigs, and when they occur are small and heal rapidly
• sows often develop vesicles and ulcers on their teats
• pregnant sows may abort (in some herds this is the first overt sign of the disease)
• there may be high mortality in suckling pigs, with sudden deaths but no vesicular lesions (in some herds this is the first overt sign of the disease).

Clinical signs in sheep and goats include:
• lameness (often the only overt sign of disease in a flock)
• foot lesions are most pronounced on the coronary bands and interdigital skin
• foot lesions are particularly prone to secondary bacterial infections, including footrot
• mouth lesions are not prominent (vesicles and ulcers are most likely to occur on the dental pad and the posterior portion of the dorsal surface of the tongue, and they tend to be small and heal rapidly)
• pregnant animals may abort
• sudden deaths may occur in young lambs as a result of cardiac lesions (the mortality rate may be as high as 90 per cent, but is more usually about 50 per cent).

FMD is generally much milder in small ruminants than in other species, and may escape detection. Therefore, careful individual examination of all individuals in a flock may be required to detect the disease.

Lesions usually progress in this manner:
• a small, blanched, whitish area develops in the epithelium
• fluid fills the area and a vesicle is formed
• vesicle enlarges and may coalesce with adjacent ones
• vesicle ruptures
• vesicular covering sloughs leaving an eroded red area
• grey fibrinous coating forms over the eroded area
• coating becomes yellow, brown or green
• epithelium is restored, but the line of demarcation remains (this gradually fades with time).

The following table can be used as a guide for ageing FMD lesions in cattle and pigs. Lesions in sheep are too transient to be used for gauging the time of infection.

<table>
<thead>
<tr>
<th>Day of clinical disease</th>
<th>Appearance of lesion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1</td>
<td>blanching of epithelium, followed by formation of fluid-filled vesicles</td>
</tr>
<tr>
<td>Day 2</td>
<td>freshly ruptured vesicles, characterised by raw epithelium, a clear edge to the lesion and no deposition of fibrin</td>
</tr>
<tr>
<td>Day 3</td>
<td>lesions start to lose their sharp demarcation and bright red colour; deposition of fibrin starts to occur</td>
</tr>
<tr>
<td>Day 4</td>
<td>considerable fibrin deposition has occurred, and regrowth of epithelium is evident at the periphery of the lesion</td>
</tr>
<tr>
<td>Day 7</td>
<td>extensive scar tissue formation and healing have occurred; some fibrin deposition is usually still present</td>
</tr>
</tbody>
</table>
Histological lesions are not specific. However, in addition to the external gross lesions already described, the following may be seen at post-mortem:

- vesicular lesions and ulcers on the ruminal pillars (ruminants)
- in young animals, focal necrosis of cardiac muscle. The lesions appear as small grey foci of irregular size and may give the myocardium a stripped appearance (‘tiger heart’) and similar lesions may also occur on skeletal muscles (all species).

**Pathology**

Gross lesions at post-mortem are generally restricted to vesicles described in the diagnosis and pathology section of Foot-and-mouth disease. Erosions may also be present on the ruminal pillars. There can also be ‘tiger heart’ striping of cardiac muscle in young animals.

**Differential diagnosis**

FMD is clinically undistinguishable from other exotic vesicular viral diseases of livestock, namely:

- Senecavirus A (Seneca Valley virus) infection
- swine vesicular disease
- vesicular exanthema of swine (not seen globally since 1956)
- vesicular stomatitis.

Regard any cattle, pigs, sheep or goats exhibiting vesicular lesions and ulcers on their feet or teats as being infected with FMD until proven otherwise by laboratory testing.

A number of other diseases cause similar clinical signs to the viral vesicular diseases, particularly during their more advanced clinical stages. Although mouth and muzzle lesions in these diseases are not vesicular, they could be confused with FMD lesions. These include:

- exotic diseases
  - bluetongue disease (infection present in Australia but without clinical disease)
  - peste des petits ruminants
  - rinderpest (now recognised by OIE as eradicated from all countries with susceptible populations)
- endemic diseases
  - bovine papular stomatitis
  - dermatophilus and other types of mycotic stomatitis
  - footrot
  - infectious bovine rhinotracheitis
  - mucosal disease (BVDV-1 only)
- non-infectious diseases that can also be confused with the viral vesicular diseases are:
  - phototoxic dermatitis with vesicle formations from contact with the leaves of plants of the family Umbelliferae (parsley, parsnips and celery)
  - chemical irritants and scalding
  - traumatic lesions of the mouth and feet.
Samples required

Sample collection

FMD virus is very sensitive to both acid and alkaline conditions. Inappropriate buffer conditions can inactivate the virus making virus isolation difficult or impossible. To maximise chances of virus isolation:

- use phosphate buffered saline or virus transport media with a pH of 7.6.
- if a sample is to be submitted after 24 hours or more, add glycerol to the phosphate buffered saline
- dilute oropharyngeal fluid collected with a probang in an equal volume of phosphate buffered saline pH 7.6, and mixed vigorously for 1 minute.

Note that collection of samples in these buffers is optimal (for the growth of FMD virus) but not essential for RNA or antigen detection assays.

Take samples from at least 10 live, clinically affected animals. Collect:

- serum, 7–10 ml/animal in plain tubes
- vesicular fluid, carefully use a syringe and needle to aspirate the vesicular fluid from unruptured vesicles, and place in a sterile container. Alternatively, collect fluid from small vesicles onto a swab and place the swab in 500 µl of buffer, such as phosphate buffered saline or virus transport medium
- fresh tissue, epithelium, epithelial tags, oral, nasal and tonsillar swabs and oropharyngeal fluid and submit in phosphate buffered saline or virus transport medium, if available.

Transport of samples

- chill blood samples and unpreserved tissue samples at 4°C, or with frozen gel packs
- DO NOT FREEZE SAMPLES at –20°C, it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests

Laboratory tests currently available for primary FMD testing include PCR, antigen ELISA, virus isolation, and serological assays for the detection of antibodies. Testing strategies are based on samples submitted and clinical and epidemiological information provided.

Reporting requirements

FMD is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect FMD, report it immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.
**Biocontainment and personal protective equipment**

There are no public health implications for FMD, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls, rubber boots (or disposable boots) and a mask.

Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property. Remember that people can harbour FMDV subclinically in the nasal passages and throat for up to 28 hours, which means that during FMD outbreaks responders should avoid moving between premises for a directed amount of time.

**Further information**


FIGURE 3.10.1 Cow with hypersalivation

FIGURE 3.10.2 Vesicles on the lower lip of a cow, suggestive of one-day old lesions

Image credit: EuFMD
FIGURE 3.10.3 Fresh lesions (2–3 days old) with epithelial tags on the upper gum of a cow

Image credit: EuFMD

FIGURE 3.10.4 Lesions with fibrin deposition on the upper gums of a cow. This is suggestive of 4-5 day old lesions

Image credit: EuFMD
FIGURE 3.10.5 Lesions on the upper gum of a cow. The re-epithelisation is suggestive of an 8-10 day old lesion

Image credit: EuFMD

FIGURE 3.10.6 Fresh lesions (~2–3 days old) on the teat of a cow

Image credit: EuFMD
FIGURE 3.10.7 Fresh vesicles on the snout of a pig (note the blanched epithelium at the base of the snout)

Image credit: EuFMD

FIGURE 3.10.8 Coronary band lesions on the feet of a pig

Image credit: EuFMD
FIGURE 3.10.9 Coronary band lesion on the foot of a sheep

Image credit: EuFMD
Hendra virus infection

Fast facts
Hendra virus (HeV) is a highly fatal zoonotic paramyxovirus primarily infecting horses which develop respiratory and neurological signs.

Organism
HeV:
- is susceptible to inactivation by desiccation, acidic conditions, heat and common disinfectants
- has an estimated maximum survival time in the environment of 5 days.

Susceptible species
- HeV affects dogs, flying foxes, horses and humans.
- HeV has infected pigs, cats, guinea pigs, hamsters, ferrets, and mice under experimental conditions.

Economic impact
There is potential for minor economic loss associated with disease control and fatalities.

Epidemiology
HeV causes high case fatality rates in horses and humans, and:
- flying foxes are the reservoir hosts
- outbreaks in horses (which may occur any time of year) have occurred annually since 2006
- incubation in horses lasts 4–16 days
- infection may occur anywhere horse and flying fox distributions overlap.

Transmission
Transmission occurs via:
- contact with pasture contaminated with infectious flying fox urine, saliva, faeces, and birthing fluids (flying fox to horse)
- direct or indirect transmission via contact with secretions and bodily fluids of affected horses (horse to horse, human, dog)
- fomite transmission possible (horse to horse, human, dog).

In addition:
- horses may transmit the virus before the onset of clinical signs
- endotracheal intubation, nasal lavage, endoscopy, and necropsy are high risk activities for vets
- transmission via respiratory droplets has not been evaluated and a 5-metre perimeter around infected horses is recommended.

Clinical signs
Clinical signs in horses include:
- acute-onset illness with rapid deterioration
- increased body temperature and heart rate
- pulmonary oedema and congestion, dyspnea, nasal discharge
- ataxia, altered consciousness, head tilt, circling, muscle spasms, seizures, recumbency
- colic-like signs
- facial oedema.
Post-mortem

Only perform post-mortem if strict biosecurity measures can be achieved. Field post mortems should not be attempted by untrained personnel.

Samples

Collect:

- blood samples (EDTA and plain tubes)
- nasal, oral and rectal swabs (in virus transport media)
- a blood clot from the jugular (if safe to do so) and a tissue sample from the mandibular lymph node from dead animals
- samples from in-contact horses and other susceptible animals.

Actions to take

If you suspect a case of HeV:

- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities
- minimise contact with suspected cases and wear personal protective equipment.

Public health considerations

- HeV infection is a zoonotic disease
- people may become infected after close contact with infected horses
- consult the relevant state or territory health authorities as soon as HeV infection is strongly suspected or confirmed.
Introduction
Hendra virus (HeV), previously known as equine morbillivirus, is a lethal zoonotic disease in Australia which primarily affects horses and people. To date, the disease has only been seen in the eastern states of Queensland and New South Wales.

Disease agent and susceptible species
HeV:
• is an enveloped RNA virus belonging to the genus Henipavirus, family Paramyxoviridae
• affects horses, humans, dogs and flying foxes
• is known to have naturally infected two dogs without causing clinical signs of illness. This has been confirmed experimentally. In one case, HeV genome was detected in bodily fluids and tissues by qRT-PCR, and in the other case the dog had seroconverted to HeV
• has infected pigs, cats, guinea pigs, hamsters, ferrets, and mice (rats seroconvert without the development of clinical signs) under experimental conditions
• a highly effective vaccine is available for use in horses.

Public health considerations
• people may become infected after close contact with infected horses
• use personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots), goggles or safety glasses, and P2 respirators or N95 masks when you are in close contact with infected horses
• consult the relevant state or territory health authorities in the case of a HeV outbreak.

Distribution
HeV has been found in flying fox populations in Australia and Papua New Guinea, and the closely related Nipah virus is found in flying fox populations in South-East Asia (please refer to the chapter on Nipah virus for more information).

Occurrences in Australia
HeV is endemic in most flying fox populations throughout Australia. To date we have seen more spillover events in northern New South Wales and South East Queensland. An observation is that spillover events have only occurred in regions where the spectacled flying fox and/or the black flying fox is present.
Epidemiology

Modes of transmission

No human-to-human transmission has been reported. However, human-to-human transmission has occurred with the closely related Nipah virus. Transmission occurs via:

- **spillover from flying fox populations**—infection in horses most likely occurs after close contact with bat urine, faeces and birthing fluids. This is thought to be due to contact with contaminated pasture, and transmission risk may be increased in horses fed under trees that bats roost in. It is also plausible that horses may become infected by inhaling droplets via the nasal route. Serological studies of bat carers suggest that bat-to-human direct transmission has not occurred.

- **direct contact** with the nasal secretions and excretions (urine, blood) of affected horses may cause horse-to-horse and horse-to-human spread of HeV. Horse-to-human transmission has only occurred with very close, prolonged contact during invasive procedures. HeV shedding, and therefore transmission, can occur before clinical signs are evident.

- **airborne spread**—inhaling infected aerosols may be a method of transmission to horses and humans. No definitive studies have been performed to describe the distance over which respiratory droplets can spread from horses. For biosecurity measures, a distance of 5 metres is used to define an at-risk area.

- **fomites**—HeV may survive on fomites for hours under mild climatic conditions and horse-to-horse transfer via fomites may occur.

Disease dynamics

There are no reports of relapsing disease in animals, but there has been one case of relapsing disease in humans. HeV infection in horses:

- is thought to occur through the naso-oral route, where a local replication in the nasal cavity or nasopharynx is followed by systemic spread of the virus

- has an incubation period of between 4–16 days. Horses may shed virus in nasal secretions before the onset of clinical signs, and are therefore a transmission risk

- causes damage to vascular endothelium, and subsequent vasculitis is thought to precipitate disease in the various organ systems (e.g. respiratory, neurologic, gastrointestinal)

- pyrexic and clinically affected horses shed virus from the nasal secretions and excretions such as urine and blood

- the case fatality rate is difficult to assess as it is likely that mild cases or cases of sudden death are not diagnosed

- humans have only been infected after exposure to HeV infected horses.

Persistence of the agent

The virus:

- has a lipid-envelope and outside the host is susceptible to desiccation and changes in temperature

- may survive from several hours to several days in the environment, depending on the environmental conditions (for disease control purposes, 5 days is presumed to be the maximum survival time under optimal environmental conditions)
is inactivated by disinfectants, including soap and detergents, Virkon®, hypochlorites, iodophors/iodine, biguanidines (e.g. chlorhexidine), and quaternary ammonium compounds.

a subunit vaccine is available for use in horses and is highly effective.

**Diagnosis and pathology**

**Clinical signs**

In humans, HeV infection causes an influenza-like illness and/or encephalitis.

In horses, the disease presents with a wide range of clinical signs and a definitive diagnosis requires laboratory testing. In fatally infected horses, illness typically lasts just over 48 hours from first clinical signs to death. An elevated body temperature or increased heart rate should be an early warning for possible HeV infection.

Common clinical signs in horses include:

- acute onset of illness with rapid deterioration, usually with respiratory and/or neurological signs
- pyrexia
- tachycardia
- discomfort/weight shifting between legs
- depression.

Respiratory signs include:

- pulmonary oedema and congestion
- respiratory distress (tachypnea)
- terminal nasal discharge which may be a stable white or blood-stained froth
- pulmonary disease leading to terminal weakness, ataxia and collapse.

Neurological disease signs include:

- ataxia
- abnormal behavior (for example, loss of vision in one or both eyes, aimless walking)
- head tilting, circling
- muscle twitching
- urinary incontinence
- recumbency with inability to rise
- facial paralysis and/or locked jaw
- spasms of the jaw or involuntary chomping
- opisthotonus
- seizures.

Other clinical signs that may be seen include:

- previous unexplained horse deaths, or a high case fatality rate where there are multiple cases
- colic-like signs (rolling, thrashing, quiet abdominal sounds on auscultation of the abdomen, teeth grinding, straining to pass manure)
- facial oedema
- wide-based stance
• anorexia
• congestion of oral mucous membranes
• stranguria
• protruding penis.

Pathology
This may be found:
• enlarged and oedematous submandibular, bronchial and/or sternal lymph nodes
• dilated pulmonary lymphatics
• pulmonary oedema and congestion with gelatinous distension of the subpleural lymphatics
• petechial haemorrhages on the pleural surfaces
• (less frequently) oedema of the mesentery, increased pleural and pericardial fluid and thick stable foam in the airways.

Differential diagnosis
There are no pathognomonic signs for horses with HeV infection. Affected horses display a large number of often vague clinical signs, resulting in a large number of differential diagnoses. Only a brief list is included here for reference. Consider in the differential diagnosis:
• exotic diseases
  - African horse sickness
  - equine influenza
  - equine protozoal myeloencephalitis
  - Japanese encephalitis
  - rabies
• endemic diseases
  - anthrax
  - bacterial infection
  - botulism
  - equine herpesvirus EHV-1 (while the neuropathogenic strain of EHV-1 is considered exotic to Australia, the abortigenic strain is endemic in certain areas)
  - inflammatory airway disease
  - Australian bat lyssavirus infection
  - tetanus
  - West Nile virus infection (Kunjin virus)
• non-infectious causes
  - snake bite
  - toxicity
  - trauma.
Samples required
Take stringent precautions during sampling. Wear full personal protective equipment with a P2 respirator for anyone in contact with the horse. Only perform post-mortem and tissue collection if you have appropriate workplace health and safety controls in place. Do not perform routinely.

Sample collection
Collect:
- serum, both plain and clotted whole blood
- EDTA blood, 7–10ml per animal
- swabs, nasal, oral, rectal and urogenital or urine
- fresh tissue in the form of a blood clot from a dead horse (collected by jugular cut-down, if safe to do so) and sample of mandibular lymph node.

Transport of samples
For transport:
- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
For HeV:
- the most rapid, sensitive and specific diagnostic procedure is the detection of viral nucleic acid in blood or swabs by qRT-PCR
- virus isolation is also conducted, but takes several days or more to complete
- serological tests, such as ELISA and virus neutralisation test are available for the detection of antibodies which have been produced through natural infection or via vaccination.

Reporting requirements
HeV infection is a notifiable disease in Australia and we have a legal obligation to report cases. If you suspect HeV infection, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.
Biocontainment and personal protective equipment

HeV infection is a serious public health risk. Human infections are due to close contact with infected horses. Post-mortem examinations of horses that have died from acute HeV infection are high-risk for horse-to-human transmission. Therefore it is important for you to seek advice from government veterinary authorities, noting:

• stringent biosecurity procedures must be enforced to protect human health
• sick/dead horses should be isolated from people as soon as possible. Limit human and other susceptible animal (e.g., dog/cat) contact with the sick/dead horse and other horses on the property until HeV diagnostic testing has been performed
• establish ‘clean’ and ‘dirty’ areas to reduce transmission of disease throughout the property
• people in the dirty area must wear personal protective equipment, including impervious rubber boots, splash-proof overalls, disposable impermeable gloves, face shield or safety eyewear and a particulate respirator
• on leaving the dirty area, remove the personal protective equipment and double bag it in clinical waste bags with Virkon® disinfectant. You must disinfect all equipment thoroughly, and wash your hands and other exposed skin thoroughly. Ensure you shower and change clothes before coming into contact with other horses
• people who have had unplanned or accidental close contact with a suspect HeV case should wash with soap and water and seek medical advice as soon as possible

Further information


FIGURE 3.11.1 Severe interlobular oedema of the lungs

Image credit: Dr M Williamson, CSIRO, and CFSPH
CHAPTER 3.12

Influenza A in pigs

Fast facts
Influenza A viruses in pigs are highly contagious and typically present as acute to chronic respiratory disease with low mortality and high morbidity.

Organism
Influenza A virus (IAV):
• replicates primarily in epithelial cells of the respiratory tract
• transfers readily between humans and pigs
• constantly evolves genetically and virulence varies with the strain of the virus
• three subtypes usually associated with disease in pigs are H1N1, H1N2 and H3N2.

Susceptible species
• Pigs, humans, ferrets, turkeys are susceptible to IAVs.

Economic impact
There is potential for severe economic loss due to high morbidity, associated production losses and the potential for secondary bacterial infections, which may result in death.

Epidemiology
• morbidity can reach 100 per cent in herds naïve to the infecting strain
• mortality is usually low but can reach up to 30 per cent due to secondary infection
• incubation period is usually 1–3 days
• peak shedding from the nasal cavity occurs at 48–72 hours with a shedding duration of 7 days.

Transmission
• is by direct contact, usually nasal discharges
• aerosol droplets from sneezing and coughing
• indirect by ingestion of infected slurry and fomites.

Clinical signs
IAV infection typically presents as an acute to chronic respiratory disease with:
• marked pyrexia
• lethargy
• inappetence
• harsh, barking cough
• sneezing
• nasal discharge.

Post mortem
• cranioventral lung consolidation
• concurrent bacterial bronchopneumonia, if present, will produce more extensive lesions.

Samples
Also refer to the detailed section on sampling for pigs with respiratory disease.
Collect:
• nasal, oral or respiratory swabs, particularly from clinically affected pigs, piglets and pigs aged 10–22 weeks
• blood in EDTA and plain tubes, particularly from sows and older pigs
• trachea and lung tissue (chilled and fixed).
Actions to take
If you suspect a case of IAV:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

Public health considerations
• IAV infection is a zoonotic disease
• people may become infected after close contact with infected pigs. There is also a risk of infection from aerosol droplet and air contamination
• use personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots), and appropriate masks if you are in close contact with infected pigs
• consult the relevant state or territory health authorities if IAV infection is strongly suspected or confirmed.
Introduction

Influenza A viruses (IAVs) are highly transmissible respiratory pathogens that circulate in poultry, wild birds and mammals worldwide. IAVs are constantly evolving, leading to new subtypes and changes in characteristics such as virulence, infectivity and host range. The genetic diversity and extent of IAV infection in Australian piggeries is largely unknown, as is the potential contribution of IAV to porcine respiratory disease outbreaks.

Disease agent and susceptible species

IAVs in pigs:
- are single stranded negative sense RNA viruses from the Orthomyxoviridae family
- H1N1, H1N2 and H3N2 are the three subtypes usually associated with disease, with multiple strains recognised within these subtypes
- when established in pig populations, can evolve further by reassortment and genetic mutation
- other subtypes including H2N3, H3N1, H4N6, H6N6, H4N8, H7N6 and H9N2, have occasionally been linked to outbreaks, but have not become established in pig populations
- can result from reverse zoonotic transmission from piggery workers to pigs, resulting in the exposure of pigs to new viruses
- can infect humans in close contact with pigs, with people usually showing symptoms similar to seasonal influenza.

Public health considerations

- IAV infection is a zoonosis
- people may become infected after close contact with infected pigs
- use personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots) and masks when you are in close contact with infected pigs
- seek medical advice if you/others develop influenza-like symptoms or otherwise become unwell
- consult the relevant state or territory health authorities if IAV infection is strongly suspected or confirmed.

Distribution

- Influenza has been clinically recognised in pigs since 1918 and IAVs have been isolated from the domestic pig populations of Europe, Asia and the United States since the 1930s.
- H3N2, H1N1 and H1N2 viruses circulate in pig herds worldwide.
- There are five global lineages of IAVs circulating in swine: classical swine H1, human seasonal H1, human seasonal H3, Eurasian avian-like H1, and triple reassortant internal gene.
- In 2009 a novel H1N1 virus (H1N1pdm09) that was made up of genes from classical swine H1, Eurasian avian-like H1 and triple reassortant lineages emerged in humans in North America quickly spreading to become a worldwide pandemic. This virus is now considered endemic in many pig populations around the world due to reverse zoonotic transmission.
- IAVs cause production losses in intensively farmed poultry and pigs and in many countries where vaccine strategies are implemented in order to control outbreaks in naïve animals.
Occurrences in Australia

- Before 2009, Australian domestic piggery herds were considered free of influenza A infection, based on serosurveys conducted in 1977 and 1997-1999.
- Following the outbreak of H1N1pdm09 in people in Australia, the virus was identified as a cause of respiratory disease outbreaks in pigs in New South Wales, Queensland, Victoria and Western Australia as a result of reverse zoonotic transmission.
- In 2012, novel reassortant H3N2 and H1N2 viruses were isolated from respiratory disease outbreaks in piggeries in Western Australia and Queensland. The genomes of these viruses were a mixture of human seasonal H1 and human seasonal H3 lineages from the 1970s to the 1990s, as well as H1N1pdm2009. Further detections of these viruses occurred in Western Australia in 2016.
- It is possible that IAV with genes from the human seasonal H1, human seasonal H3 and H1N1pdm2009/classical swine lineages are circulating in Australian pigs due to historical and ongoing reverse zoonotic transmission. Viruses from the Eurasian avian-like and triple reassortant lineages are less likely to be present in Australian herds due to restrictions on the importation of live pigs.

Epidemiology

Modes of transmission

Transmission occurs via:

- **pig movements**—between herds, the main method of spread of swine-origin IAV is through the movement of infected pigs, for example via introduced stock
- **direct contact**—within herds, IAV are primarily transmitted among pigs in close contact through nasal discharges and aerosols from sneezing and coughing
- **airborne spread**—between pigs via aerosolised respiratory droplets readily occurs
- **fomites**, between piggery sheds may occur via fomites such as contaminated equipment, footwear and vehicles in enterprises with poor biosecurity
- **other species**—humans and birds may be a source of infection and spread of IAVs to pigs. Transmission of IAVs from pigs to humans also occurs. Feral pigs may be sources of infection.

Disease dynamics

- the incubation period for IAV in pigs is usually 1–3 days
- although a distinct seasonality of IAV incidence in pig herds has not been consistently documented, as with seasonal human IAV circulation, outbreaks are more likely to occur during the colder seasons
- pigs begin excreting the virus from the upper respiratory tract within 24 hours of infection, with peak shedding at 48–72 hours and a duration of shedding of up to 7 days
- morbidity can reach 100 per cent in herds naïve to the infecting strain but mortality is usually low
- in endemic herds, pigs aged 10–22 weeks are most likely to be affected by IAV and should be targeted for detection of active infection.
Persistence of the agent

- Due to the fragile viral lipid envelope, IAVs are highly susceptible to detergents and most commonly used disinfectants including sodium hypochlorite, 70% ethanol, oxidizing agents, quaternary ammonium compounds, aldehydes (formalin, glutaraldehyde, formaldehyde), phenols, acids, povidone-iodine and lipid solvents.
- IAVs can be inactivated by heating at 56°C for a minimum of 60 minutes (or higher temperatures for shorter periods) and by low pH (pH <2).
- Cool, moist conditions contribute to environmental survival of IAVs.
- Studies have shown that IAVs survive for 24–48 hours on hard, nonporous surfaces such as stainless steel and plastic and for up to 8–12 hours on cloth, paper and tissues.
- IAVs survived on hands for up to 5 minutes after transfer from the environmental surfaces.
- Studies have also reported sustained detection of IAV RNA in air during outbreaks in piggeries.

Diagnosis and pathology

Clinical signs

- A typical outbreak of IAV in a pig herd presents as an acute, high morbidity, low mortality infection that spreads rapidly through groups of pigs.
- During the first 1–2 days of infection, there may be a rapid onset of pyrexia up to 41°C, lethargy and inappetence. Respiratory signs such as serous nasal discharge, slight coughing and laboured abdominal breathing may also be present in this early phase. When pigs are forced to move, the heaves and tachypnea will become more pronounced.
- By days 3 and 4, pigs demonstrate a harsh, deep, barking cough which is the hallmark clinical sign of this disease.
- In the absence of secondary bacterial infection, pigs usually recover from IAV infection within 6–7 days.
- Due to pyrexia or transplacental infection, pregnant animals may abort.
- In combination with other viruses or bacteria, IAVs may play a role in a severe, multifactorial respiratory disease syndrome known as the porcine respiratory disease complex.
- Not all pigs will be infected simultaneously, so clinical signs in the herd may last 2 or more weeks before abating, followed by gradual weight gain and return to normal body condition.
- Many pig herds can become infected with one or more IAV subtypes without showing clinical signs.
- Mortality rates vary depending on the presence of concurrent bacterial infection.

Pathology

- IAVs are a primary respiratory pathogen whereby infection with the virus alone can induce pathology in respiratory epithelial cells.
- Pathology relating to IAV infection in pigs is strain dependent but the most common macroscopic manifestation of IAV respiratory infection is a cranioventral consolidation that can affect as much as 40 per cent of the lung.
- Field cases will often present with a concurrent bacterial bronchopneumonia which results in more extensive lesions.
- Although the virus infects the epithelial lining of the nasal turbinates and trachea these organs are usually unremarkable.
- Bronchitis, bronchiolitis and interstitial pneumonia are usually evident to some degree on histopathology.
Differential diagnosis

Consider in the differential diagnosis:

- **Exotic diseases**
  - Aujeszky’s disease
  - Nipah virus infection
  - Porcine reproductive and respiratory syndrome

- **Endemic diseases**
  - Actinobacillus pleuropneumoniae
  - Atrophic rhinitis (Bordetella bronchiseptica)
  - Inclusion body rhinitis (porcine cytomegalovirus)
  - Miscellaneous opportunistic bacterial infections (Haemophilus parasuis, Pasteurella multocida, Streptococcus suis)
  - Mycoplasmal pneumonia (Mycoplasma hyopneumoniae, Mycoplasma hyorhinis)
  - Porcine circovirus type 2 associated diseases
  - Influenza A viruses (human-origin strains, subtypes H1N1, H1N2 and H3N2).

Samples required

Sample collection

Submit several pigs (at least five) for post mortem examination, as there may be significant variability in lesions seen. Record a composite picture of all lesions. Older pigs such as sows, boars and finisher pigs are most likely to have serum antibodies against IAVs while piglets and weaned pigs aged 10–22 weeks are more likely to be shedding infectious virus (maternal antibodies wane from 6–12 weeks of age).

Collect:

- **Nasal swabs** from untreated, acutely ill pigs, within 24–72 hours of development of clinical signs
- **EDTA blood** (7–10 ml per animal) from live, clinically affected animals.
- **Oral fluids**, using cotton ropes hung in pig pens, preferably from weaners and growers
- **Blood serum** samples from 30 suspected chronically infected animals (preferably sows, finishers, boars) to screen for seroconversion
- **Fresh tissue**, from lung, tonsils and trachea and any organ showing gross pathology, chilled and immersed in virus transport media or phosphate buffered saline (PBS).
- **Fixed tissue**, bronchiole and lung tissue and any organ showing gross pathology in neutral-buffered formalin.

Transport of samples

For transport:

- Chill blood samples and unpreserved tissue specimens to 4°C, or with frozen gel packs
- Place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- **Do not freeze samples** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days
• formalin fixed tissue can be shipped at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
• the most rapid, sensitive and specific diagnostic procedure is the detection of viral nucleic acid in whole blood and/or tissues by PCR
• whole genome sequencing is attempted on samples in order to characterise the virus, which may differ from pig to pig due to propensity for reassortment (even within a single outbreak)
• serology is most useful as a surveillance tool however, due to antigenic diversity in IAVs, detection would require further investigation before an active infection is confirmed
• IAV can also be identified using an ELISA or haemagglutination inhibition (HI) test
• virus isolation can be achieved using embryonated chicken eggs and continuous cell lines
• specialised staining for IAV nucleoprotein can be done on fixed tissues to identify IAV antigen and any associated histopathology.

Reporting requirements
Influenza A infection in pigs is a notifiable disease in Australia and there is a legal obligation to report cases. If you suspect IAV infection in pigs, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
IAV infection in pigs is a public health risk. Human infections can occur following close contact with infected pigs; however, it is important to remember that reverse zoonosis can occur, when humans infect pigs. This transmission is probably by fomites and aerosol droplets. It is important that you seek advice from government veterinary authorities, noting:
• biocontainment requires the appropriate use and disposal of personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots) and masks
• personal decontamination is required when leaving infected premises. Thoroughly clean and then disinfect vehicles and equipment before leaving the property
• avoid the spread of infection from premises by contaminated fomites such as personnel clothing, poultry litter, equipment and manure
• discourage staff or visitors to enter the piggery if feeling unwell
• any people participating in the biocontainment of an IAV outbreak must be vaccinated against seasonal influenza.
Further information


CHAPTER 3.13

Japanese encephalitis

Fast facts
Japanese encephalitis is a mosquito-borne viral disease of humans and animals.

Organism
Japanese encephalitis virus (JEV):
• strains can vary in virulence
• is readily inactivated by heat, photochemical treatment, pasteurisation, treatment with detergents and low pH.

Susceptible species
Humans, horses, donkeys and pigs are susceptible.

Economic impact
There is potential for significant economic loss due to high morbidity and reproductive losses in pigs, as well as restrictions on animal movements and international trade.

Epidemiology
Japanese encephalitis:
• causes morbidity (50 per cent) and mortality (30 per cent) in humans
• causes low morbidity in horses
• has a case fatality rate in horses of between 5–15 per cent in endemic areas and 30–40 per cent in seasonal epidemics
• can cause high reproductive losses in pig herds (up to 50–70 per cent) and 100 per cent mortality in pig litters
• subclinical infection is most common
• incubation period in horses is 4–14 days and in pigs is 1–3 days.

Transmission
Japanese encephalitis is mosquito-borne and Culex species are the main vectors.

Clinical signs
Clinical signs are:
• mild pyrexia or no overt signs in adult pigs
• reproductive failure in sows or gilts
• encephalitis in horses and humans.

Post-mortem
Typical findings include:
• lesions in the testes of boars
• microscopic lesions typical of non-suppurative meningoencephalitis in horses and adult pigs
• hydrocephalus, subcutaneous oedema, cerebellar hypoplasia and spinal hypomyelination in aborted foetuses
• gross lesions in horses (including severe congestion of the meninges, hypophysis and cerebral cortex).

Samples
Collect:
• a range of tissues in formalin
• whole blood and serum samples (paired serum samples should be collected 2–4 weeks apart)
• placental tissues from aborted foetuses
• cerebrospinal fluid and a range of brain tissue from animals with neurological signs.
**Actions to take**

If you suspect a case of Japanese encephalitis:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

**Public health considerations**
- Japanese encephalitis is a vector-transmitted zoonosis
- use insect repellents or netting to protect against mosquitoes
- consult the relevant state or territory health authorities as soon as Japanese encephalitis is strongly suspected or confirmed.
**Introduction**

Japanese encephalitis is a mosquito-borne viral disease of humans and animals that occurs throughout much of Asia. Japanese encephalitis virus (JEV) infections are mainly subclinical, but can be associated with abortion in pigs and encephalitis in humans and horses. Although an effective vaccine is available, approximately 67,000 human cases occur annually.

**Disease agent and susceptible species**

The Japanese encephalitis serogroup comprises antigenically related viruses including West Nile (and the Australian subtype Kunjin), St Louis encephalitis, Murray Valley encephalitis, Alfuy and Usutu.

**JEV:**
- is enveloped and has a single-stranded RNA genome
- is a member of the *Flavivirus* genus of the family *Flaviviridae*
- replicates in various tissues with tropism for neurological and lymphoid tissues
- comprises strains that vary in virulence.

The species affected include:
- humans, horses and donkeys, which are dead-end hosts but can develop severe encephalitis following infection
- pigs and infection of pregnant sows or gilts can lead to reproductive failure with abortions and stillbirths, while infection of boars has been associated with aspermia. Encephalitis has also been observed in piglets following experimental infection.
- Other species that can be infected but do not exhibit disease include:
  - cattle, dogs, sheep and goats (also dead-end hosts)
  - waterbirds (herons and egrets), which are the main reservoir for spreading JEV
  - pigs and waterbirds, which are important amplifying hosts
  - several bat species, and fruit bats of the order Microchiroptera, which can develop prolonged viraemia capable of infecting mosquitoes under experimental conditions
  - lizards, which have been shown experimentally to become persistently infected and transmit virus to mosquitoes.
  - marsupial species, such as wallabies and possums, have also been shown experimentally to become infected and develop viraemia that may be sufficient to infect vector mosquitoes, although this has not been tested
  - the susceptibility of other Australian native fauna is not known.

**Public health considerations**

- Japanese encephalitis is a vector-transmitted zoonosis
- use insect repellents or netting to protect against mosquitoes
- consult the relevant state or territory health authorities in the case of a Japanese encephalitis outbreak.
Distribution

Japanese encephalitis is widely dispersed throughout Asia as follows:

• its geographic range extends to maritime areas of the north-east Russia, China and Philippines in the east, India and Pakistan at its western limits, and throughout South-East Asia.

• outbreaks of Japanese encephalitis have also been reported in Papua New Guinea (PNG) and northern Australia, as well as in the Western Pacific islands of Guam and Saipan.

• in some Asian countries, pigs are vaccinated as part of control efforts, and horses in endemic areas are vaccinated to protect against disease and for international travel.

Occurrences in Australia

Occurrences in Australia are described:

• Japanese encephalitis emerged in the Torres Strait, northern Australia in 1995 and subsequently spread to northern Queensland where infections of pigs and humans occurred. In 1995, three human cases occurred on Badu Island (in the Torres Strait) and two of these were fatal. A serological survey found further evidence of infection in people (subclinical), pigs, horses and dogs on Badu and other outer islands. Virus was isolated from two subclinical human cases and from Culex annulirostris mosquitoes, which were implicated as the major vector involved in the outbreak.

• The first isolate of JEV from mosquitoes in the NPA was made from collections undertaken in 2004 following seroconversions in sentinel pigs. Despite these detections on mainland Australia, JEV does not appear to have become established and no further cases have been detected.

• The source of the periodic incursions that have occurred are thought to have been infected windblown mosquitoes or migrating water birds from PNG, where the virus is endemic.

Epidemiology

Modes of transmission

Transmission occurs via:

• **vectors**—mosquitoes primarily transmit JEV. JEV has been isolated from over 30 species of mosquito. In South-East Asia, the major vector is the rice field-breeding *Cx. tritaeniorhynchus*, whereas in Australia, the principle vector mosquito is *Cx. annulirostris*. JEV vectors are opportunistic feeders and are zoophilic, preferring animals to humans. While host availability is an important factor influencing feeding, bovine species are a preferred source of blood meals for *Cx. tritaeniorhynchus*. Since cattle are dead-end hosts, they may dampen JEV transmission. In Australia, *Cx. annulirostris* preferentially feed on cattle and marsupials.

• **vertical transmission** can occur in mosquitoes (transovarial) and pigs (transplacental).

• **direct transmission** has been demonstrated experimentally in pigs via the oronasal route.
Disease dynamics

Pigs are the main amplifying hosts of JEV and can act as maintenance hosts in endemic areas. Water birds, particularly herons and egrets, are important reservoir hosts, but may also act as amplifiers in areas where pigs are absent or in low numbers. Following natural infection, pigs and waterbirds develop a prolonged and high level viraemia capable of infecting mosquito vectors. The high birth rate and rapid turnover of pigs in commercial production systems provide a continual source of susceptible hosts and large epidemics of Japanese encephalitis have occurred when JEV spreads to new areas where susceptible hosts and suitable vectors co-exist.

A basic cyclical pattern of JEV transmission between pigs, mosquitoes and humans is recognised which involves two amplification cycles in pigs as follows:

- the first cycle infects a proportion of the herd, from which vector mosquitoes become infected (virus may be detected in mosquitoes prior to amplification in pigs in some instances)
- after a 1–2 week incubation period, mosquitoes transmit the virus to remaining susceptible pigs
- after a second incubation period, human and equine cases commence.

Japanese encephalitis predominantly affects rural areas where pig-rearing, irrigated agriculture and human habitation are in close proximity. Typically, two epidemiological patterns exist: endemic activity in tropical regions and epidemic activity in temperate and subtropical regions. In tropical areas, the virus is endemic and sporadic cases occur throughout the year, with small outbreaks occurring during monsoonal seasons. In temperate areas, the disease is epidemic with outbreaks occurring in summer or early autumn. Exceptions to these patterns can occur, such as in tropical northern Australia where peak JEV activity has followed a temperate pattern.

Several overwintering mechanisms have been proposed for JEV in temperate areas. These include survival in dormant mosquito eggs following vertical transmission, survival in hibernating mosquitoes or animals, such as bats or reptiles, or transmission from persistently infected pigs. JEV may also be re-introduced by viraemic migratory birds or from wind-blown infected mosquitoes. One or more of these methods may contribute to Japanese encephalitis epidemiology in different areas.

In addition:
- morbidity and mortality in humans is approximately 50 per cent and 30 per cent, respectively
- Japanese encephalitis in livestock has not been accurately quantified, but mortality rates in horses are estimated at 5–15 per cent in endemic areas and 30–40 per cent in seasonal epidemics
- the incubation period in horses is 4–14 days and in pigs is 1–3 days
- the morbidity rate in horses is low (<1 per cent) and up to 1.4 per cent during epidemics
- reproductive losses in pig herds can be up to 50–70 per cent during epidemics
- mortality in litters of pigs can reach 100 per cent but is rare in adult pigs.
Persistence of the agent

JEV is relatively unstable and can be inactivated by a number of means, including heat inactivation in serum at 37°C for 30 minutes and a wide range of chemicals and disinfectants (e.g. 1% sodium hypochlorite, 70% ethanol, iodophore, quaternary ammonium disinfectants).

Diagnosis and pathology

Clinical signs

In pigs, the most common form of disease is reproductive failure in pregnant sows or gilts infected before 60–70 days gestation, which leads to abortion and litters containing stillborn or mummified foetuses or live, weak piglets. In addition:

- adult pigs do not generally show overt signs of infection
- under experimental conditions, pigs can become pyrexic (40–41°C) after 24 hours of infection, lasting up to 5 days, with inappetance and depression
- viraemia can be detected experimentally from 1–3 days post-inoculation
- central nervous signs indicative of encephalitis, such as hind limb tremor, have been occasionally reported in piglets
- infected boars may have oedematous, congested testicles resulting in lowered motile sperm counts and abnormal spermatozoa.

In horses, subclinical infection is most common. Note that neurological sequelae can occur in surviving horses, and has manifested as incoordination, paraplegia, ataxia and incontinence. Three clinical syndromes have been described as:

- **transient type**—pyrexia of up to 40°C for 2–4 days, with anorexia, sluggish movement, congested or jaundiced mucous membranes, and rapid and uneventful recovery
- **lethargic type**—fluctuating pyrexia up to 41°C, with lethargy, anorexia, stupor, grinding of teeth and chewing motions, difficulty in swallowing, jaundice, petechial haemorrhages in mucous membranes, incoordination, staggering and falling, transient neck rigidity, radial paralysis, impaired vision and recovery within 1 week
- **hyperexcitable type**—marked pyrexia (>41°C), with aimless wandering, violent and demented behaviour, blindness, profuse sweating, muscle trembling, bruxism, collapse, coma and death.

Pathology

There are no characteristic gross lesions in infected sows. Otherwise, the following may be seen:

- accumulation of mucous fluid in the cavity of the tunica vaginalis, fibrous thickening of the epididymis and visceral lamina of the connective tissue surrounding the testes (tunica vaginalis)
- hydrocephalus, subcutaneous edema, cerebellar hypoplasia and spinal hypomyelinogenesis in stillborn foetuses and live, weak piglets
- severe congestion of the meninges, hypophysis and cerebral cortex with foci of malacia and haemorrhage in horses
- lesions throughout the cerebral hemispheres in horses.
Differential diagnosis

In pigs, clinical disease characterised by abortion, foetal mummification or stillbirth, and encephalitis in animals up to 6 months old should be considered in the differential diagnosis. Japanese encephalitis in horses must be differentiated from other causes of neurological diseases. In Australia this includes infections with the closely-related subtype Kunjin virus and Murray Valley encephalitis viruses, which have caused outbreaks of equine encephalitis and have very similar epidemiology to Japanese encephalitis.

Consider in the differential diagnosis:

• exotic diseases:
  - Aujeszky’s disease
  - blue eye disease (rubulavirus infection)
  - Borna disease
  - enteroviral encephalomyelitis
  - equine arboviral encephalitides (e.g. Elsey viruses, eastern equine encephalitis, western equine encephalitis, Venezuelan equine encephalitis viruses)
  - Nipah virus infection
  - porcine polioencephalomyelitis (Teschen disease)
  - porcine reproductive and respiratory syndrome
  - rabies
  - swine vesicular disease

• endemic diseases:
  - bacterial meningoencephalitis
  - botulism
  - haemagglutinating encephalomyelitis
  - Hendra virus infection
  - herpes myeloencephalopathy (while the neuropathogenic strain of EHV-1 is considered exotic to Australia, the abortigenic strain is endemic in certain areas)
  - Menangle virus infection (not reported in Australia since 1997)
  - porcine brucellosis (infection in feral pigs, with rare occurrence in domestic pigs)
  - porcine parvovirus infection
  - tetanus
  - verminous encephalitis

• non-infectious diseases:
  - hepatic encephalopathy (e.g. Crotalaria plant poisoning or post-vaccinal hepatitis)
  - indigofera plant poisoning (Birdsville disease)
  - mycotoxin poisoning and nigropallidal encephalomalacia (the latter is rare in Australia while mycotoxin poisoning is more common)
  - poisons such as arsenic, mercury, monensin and organophosphorus compounds
  - pantothenic acid deficiency (vitamin B)
  - water deprivation or salt poisoning.
Samples required

Sample collection
Collect samples from animals euthanised in the acute stage of the disease or from animals that have been dead for less than 12 hours.

Collect:

- **whole blood** (EDTA) and serum samples for virus detection and serology (for serum, collect at least 7–10 mL of blood from animals in the acute phase and convalescent stage of the disease). Collect paired serum samples 2–4 weeks apart
- **cerebrospinal fluid** (CSF) in animals presenting with neurological signs. Place into a sterile container
- **fresh brain, spinal cord, tonsils** from pigs, **placenta** from fetuses and neonates. Since viremia only lasts a few days and virus is rarely cultured from blood or CSF, isolation is best achieved from infected brain and post-mortem tissues. Remove the brain aseptically and collect a range of brain tissue samples into sterile containers
- **formalin fixed tissues** (full range of tissues including brain).

Transport of samples
For transport:

- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Laboratory diagnosis for JEV is achieved by virus isolation, molecular detection and serological tests. In addition:

- molecular methods (reverse transcriptase-PCR) are used to detect and identify viral RNA genome in a range of samples, including infected tissues, CSF, sera and whole blood
- infection with JEV can be detected in fixed tissues using immunohistochemistry
- diagnosis of JEV infections is often reliant on detection of antibodies
- as there is a high degree of serological cross-reactivity between flaviviruses, take care in interpreting results in areas where related flaviviruses co-circulate, especially those of the JE serogroup (these viruses should be included in parallel testing)
- a reliable diagnosis may be made where IgM to JEV is detected in the CSF in the absence of IgM to other flaviviruses
- IgG is even more broadly cross-reactive, and identifying virus-specific antibody is generally confirmed using the plaque-reduction neutralisation test; a diagnosis of JEV infection is based on the demonstration seroconversion between paired serum samples (≥ 4 fold rise in antibody titre to JEV compared to related circulating flaviviruses).
Reporting requirements

Japanese encephalitis is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect Japanese encephalitis, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

JEV is a public health risk. Human infections can potentially occur via needlestick and mucosal exposure or inhalation of infected samples. Contact with aborted foetuses, bodily fluids and tissues of infected animals (as well as aerosols generated during post-mortem examination) are also potential sources of human infection.

Therefore, keep suspected cases of Japanese encephalitis isolated and on the property (and protected from vectors, if possible) until you seek advice from government veterinary authorities, noting:

- wear appropriate personal protective equipment (coveralls and boots, gloves, mask) when examining animals, performing post-mortem examinations, or handling tissue, carcasses or abortion materials
- isolate sick/dead animals and limit contact with the sick/dead animals
- personal decontamination is required when leaving infected premises. Thoroughly clean and then disinfect vehicles and equipment before leaving the property
- vaccination is recommended for personnel working in Japanese encephalitis endemic areas, such as the Torres Strait.

Further information


CHAPTER 3.14
Lumpy skin disease

Fast facts
Lumpy skin disease (LSD) is a poxvirus disease of cattle, characterised by pyrexia, and nodules on the skin, mucous membranes and internal organs.

Organism
Lumpy skin disease virus (LSDV):
• closely related antigenically to sheeppox virus and goatpox virus
• may remain viable in environment for prolonged periods.

Susceptible species
Cattle and water buffalo are the only livestock species affected by LSD.

Economic impact
There is potential for significant economic loss due to high morbidity and drop in milk yield in cattle, as well as restrictions on animal movements and international trade.

Epidemiology
• morbidity usually varies between 10 and 20 per cent
• mortality is usually negligible
• incubation period is usually 12 days, but may range from 4–28 days.

Transmission
Transmission occurs:
• mainly by arthropod vector (biting insects)
• less commonly by direct contact with skin lesions, saliva, nasal discharge, milk or semen of infected animals.

Clinical signs
Clinical signs include:
• sudden onset of pyrexia, reluctance to move, nasal and ocular discharges, hypersalivation and lymphadenopathy
• cutaneous nodules of 2–5 cm in diameter, particularly on the head, neck, limbs, udder, genitalia and perineum
• fibrinous plaques eventually form over the top of the larger nodules.

Post-mortem
Typical findings include:
• extensive greyish-pink skin nodules with caseous necrotic centres
• similar nodules may be found in the nasopharynx, trachea, bronchi, lungs, rumen, abomasum, renal cortex, testicles and uterus
• swollen, congested lymph nodes with petechial haemorrhages.

Samples
Collect:
• fresh tissue samples—skin lesion biopsies, scabs, vesicular fluid (if present) and lesions in the respiratory and gastrointestinal tracts
• skin scrapings
• whole blood in EDTA.

Actions to take
If you suspect a case of LSD:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Lumpy skin disease is a generalised viral disease of cattle that is spread by biting insects. The virus, which is closely related to the pox viruses of sheep and goats, causes nodular skin lesions on the animal’s body. It can cause considerable production losses.

Disease agent and susceptible species

Lumpy skin disease is caused by lumpy skin disease virus (LSDV) which is a pox virus. LSDV:

- is an enveloped, double stranded DNA virus in the family Poxviridae, subfamily Chordopoxvirinae, genus Capripoxvirus
- is very closely related to sheep and goat pox
- infects all cattle (Bos taurus are more susceptible than Bos indicus) and water buffalo (Bubalis bubalis)
- can infect a number of wildlife species, including giraffe and impala; however, a wildlife reservoir has not been identified
- does not infect humans.

Distribution

Globally, LSDV:

- is endemic in Africa, being virtually throughout the entire continent, including Madagascar
- spread into the Middle East in the 1990s
- by 2013 spread to Turkey, and from there westwards into Europe including Greece, Russia, Serbia and Georgia

For the latest information on the distribution of LSD, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

LSD has never been reported in Australia.

Epidemiology

Modes of transmission

Transmission occurs via:

- vectors—most commonly, insects (such as mosquitoes and biting flies) act as mechanical vectors of the virus
- secretions and excretions—less commonly, transmission can be through direct contact with secretions and excretions
- semen—shedding in the semen of clinically affected bulls may be prolonged; viral DNA has been found in the semen of some bulls for at least 5 months after infection, and live virus for up to 42 days; subclinically affected bulls may shed virus for at least 12 days.
Disease dynamics

LSDV:
- may be present in all secretions and excretions (including milk and semen) of infected animals. However, the level of virus shedding from mucosal surfaces is generally low
- is present at high concentrations in skin lesions
- may cause disease ranging from inapparent or mild to severe, depending on age, breed, immunity, and other factors
- the incubation period usually lasts 12 days, but may vary from 4–28 days
- would likely cause severe disease in Australian cattle populations (which are fully susceptible)
- skin nodules up to 5 cm or more in diameter develop within 1–2 days of the onset of pyrexia, especially around the head, neck, genitalia, limbs, udder and perineum
- although few adult cattle die from the disease, many become debilitated and can remain in extremely poor body condition for prolonged periods (up to 6 months or longer)
- has no carrier state and recovered animals develop lifelong immunity.

Persistence of the agent

Capripoxviruses are very stable in the environment and can remain viable for long periods, on or off the animal host. LSDV:
- can survive in necrotic skin nodules for up to 33 days or longer, in desiccated crusts for up to 35 days, and at least 18 days in air-dried hides
- in dark environmental conditions, such as contaminated animal sheds, can persist for many months
- is susceptible to sunlight and detergents containing lipid solvents and acids. Therefore, acids combined with detergents (such as Virkon®) are the disinfectants of choice, particularly for areas where organic matter is prevalent. Hypochlorites and aldehydes are useful for disinfecting clean surfaces. Citric acid, alcohols and iodophors are suitable for personal disinfection
- is inactivated after heating for 2 hours at 55°C.

Diagnosis and pathology

Clinical signs

Clinical signs include:
- may vary widely between individuals within the same herd
- nodular and nasal discharge and hypersalivation (usually observed first)
- a sudden onset of pyrexia, which may exceed 41°C
- sharp drop in milk yield
- highly characteristic, nodular skin lesions of 2–5 cm in diameter, particularly on the head, neck, limbs, udder, genitalia and perineum within 48 hours of onset of pyrexia. These nodules are circumscribed, firm, round and raised, and involve the skin, subcutaneous tissue and sometimes even the underlying muscles
- centres of large nodules may become necrotic and eventually fibrotic and persist for several months
- anorexia
- conjunctivitis
• enlarged superficial lymph nodes
• vesicles, erosions and ulcers may develop in the mucous membranes of the mouth and alimentary tract, and in the trachea and lungs
• acute respiratory distress, and in more severe cases, commonly secondary pneumonia, which can be fatal
• oedema of the limbs, brisket and genitals may occur
• abortion, intrauterine infections and temporary sterility of bulls and cows may also occur.

Pathology
In addition to the lesions described:
• nodular lesions may be scattered through the nasopharynx, trachea, bronchi, lungs and the gastrointestinal tract, including the rumen and abomasum, the renal cortex, testicles and uterus
• there may also be enlarged and haemorrhagic lymph nodes.

Differential diagnosis
Acute cases in naïve populations should be readily diagnosed based on clinical signs. However, clinical signs may vary widely between animals within the same herd some strains of low virulence may produce only mild clinical signs or subclinical disease. Consider in the differential diagnosis:
• exotic diseases
  – cutaneous tuberculosis
  – screw-worm fly myiasis
• endemic diseases
  – bovine herpes virus 2 (pseudo-lumpy skin disease)
  – bovine papular stomatitis
  – dermatophilosis
  – ectoparasites
  – pseudocowpox
• non-infectious causes
  – insect and tick bites
  – photosensitisation
  – skin allergies.

Samples required
Sample collection
Virus isolation will be possible within the first week of clinical signs developing, before neutralising antibodies develop. Collect:
• serum from at least 10 live, clinically affected animals, and from exposed animals (particularly those that are convalescent)
• EDTA blood from live, clinically affected animals (7–10 ml/animal)
• fresh tissue, characteristic skin lesions from skin as well as internal lesions if present, and regional lymph nodes (2 g of each tissue)
• fixed tissue, characteristic skin lesions from skin as well as internal lesions if present, and regional lymph nodes (in neutral-buffered formalin).
Transport of samples
For transport:
• chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
• send samples with dry ice if the journey is expected to take several days
• formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
For LSD:
• a rapid, tentative laboratory diagnosis can be made by electron microscopy of tissue samples. Electron microscopic examination of tissue samples reveals typical capripoxvirus particles
• confirmation of the diagnosis is obtained by specifically identifying the virus in tissues from early lesions or in cell culture using virus-specific tests
• the most rapid, sensitive and specific diagnostic procedure is the detection of viral DNA in characteristic pox virus lesions or nasal swabs by real-time or conventional PCR. A positive result can be obtained within 1 day of samples being received at the laboratory.

Reporting requirements
LSD is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect LSD, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.
Biocontainment and personal protective equipment

There are no public health implications for LSD, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


SECTION 3 EMERGENCY ANIMAL DISEASES

FIGURE 3.14.1 Disseminated cutaneous papules

Image credit: PIADC and CFSPH

FIGURE 3.14.2 Disseminated cutaneous papules with necrotic centres

Image credit: PIADC and CFSPH
FIGURE 3.14.3 Sloughing of the necrotic centres of two cutaneous papules

Image credit: PIADC and CFSPH

FIGURE 3.14.4 Haemorrhagic exudate associated with the necrotic centre of a cutaneous papule

Image credit: PIADC and CFSPH
FIGURE 3.14.5 Multiple sharply-demarcated slightly raised papules, with eroded surfaces, that extend into the nares

FIGURE 3.14.6 Marked generalised interlobular oedema of the lungs with a small cluster of red nodules on the left side of the specimen
CHAPTER 3.15
Newcastle disease

Fast facts
Newcastle disease (ND) is a highly pathogenic disease of poultry which can cause significant mortalities. Newcastle disease only refers to infection in poultry (not wild birds) with virulent strains of avian avulavirus 1.

Organism
Avian avulavirus 1:
- varies in virulence with strain and host species
- can survive at room temperature for extended periods of time.

Susceptible species
ND replicates in all bird species, including poultry. ND can cause mild infection in humans.

Economic impact
There is potential for significant economic loss associated with poultry morbidity and mortalities, control and eradication costs, as well as restrictions on animal movements international trade.

Epidemiology
Varies with the pathotype of the virus and the species infected. In addition:
- velogenic viruses cause high morbidity and mortality and spread rapidly
- higher pathogenicity viruses usually cause shorter incubation periods
- Incubation periods are usually less than 1 week but (2–6 days) can be as long as 15 days.

Transmission
Transmission occurs via:
- the faecal-oral route and inhalation
- vertical transmission
- fomites
- mechanical vectors (rodents, insects and humans).

Clinical signs
Clinical signs vary with virus strain and host species and include:
- mortality
- conjunctivitis
- diarrhoea
- dyspnoea
- ataxia, paresis, paralysis, torticollis
- decreased egg production.

Post-mortem
In young birds, or in infection with a strain causing rapid death, no gross lesions may be present. Otherwise, the following may be found:
- haemorrhage/oedema of conjunctiva, ovaries and diphtheritic membranes overlying tracheal mucosa
- haemorrhage/oedema/ulceration of lymphoid accumulations (in the Peyer’s patches, caecal tonsils, proventriculus and bursa)
- splenomegaly
- swelling of the head.
Samples
Collect:
- serum
- oropharyngeal and cloacal swabs
- fresh tissues (lung, kidney, intestine, spleen, brain, heart)
- formalin fixed tissues (a full range).

Actions to take
If you suspect a case of ND:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

Public health considerations
- ND is a zoonotic disease
- people (predominantly poultry workers and laboratory staff) may become infected after close contact with infected birds or materials.
Introduction

Newcastle disease (ND) is a zoonotic, highly pathogenic disease of birds which can cause significant mortalities, particularly in poultry. The disease varies in pathogenicity and is associated with gastrointestinal, respiratory and/or neurological signs.

Disease agent and susceptible species

ND is caused by virulent forms of avian avulavirus (AAvV-1). Also known as Newcastle disease virus (NDV), and formerly known as avian paramyxovirus 1, AAvV-1:

- is a virus within the genus *Avulavirus*, family *Paramyxoviridae* and order *Mononegavirales*
- can be classified into five pathotypes: 1) viscerotropic velogenic, 2) neurotropic velogenic, 3) mesogenic, 4) lentogenic, 5) avirulent (and only velogenic or mesogenic pathotypes fulfill the OIE definition of ND)
- is an RNA virus which can be separated into class I and class II viruses. Class I viruses are predominantly found in wild waterfowl and are avirulent. However, class II viruses are found in both wild birds and poultry and contain both avirulent and virulent viruses
- is a zoonotic agent and infections are occasionally seen in people, predominantly in poultry workers and laboratory staff. Clinical signs observed include mild to moderate conjunctivitis and, rarely, flu-like symptoms
- replicates in all bird species but species susceptibility to disease varies
- chickens are highly susceptible to virulent forms of NDV but the virus can also be pathogenic for turkeys
- NDV rarely causes clinical disease in ducks and geese but game birds (including pheasants, partridges, and guinea fowl) are susceptible to disease with the virus being particularly pathogenic in quail
- disease in wild bird species is not common but significant outbreaks of disease have been seen in young double-crested cormorants (*Phalacrocorax auritus*) in the United States and Canada.

Public health considerations

- people may become infected after close contact with infected birds or material
- when in close contact with infected poultry use personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots), safety glasses and mask.

Distribution

ND is endemic in many countries (particularly throughout southern and Central America, Asia, the Middle East and Africa) and sporadic outbreaks of disease regularly occur worldwide. The pigeon-adapted strain of AAvV-1, pigeon paramyxovirus (PPMV-1), is also present worldwide.

For the latest information on the distribution of ND, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

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1 Highly virulent
2 Intermediate virulence
3 Low virulence
Occurrences in Australia

While avirulent strains of NDV are widespread and have been detected in Australia since the 1960s, Australia is free of virulent ND. In addition:

- there have been no detections of virulent ND since 2002
- Victoria experienced two outbreaks of ND in 1930 and 1932 (both outbreaks were quickly eradicated) and a single outbreak in 2002
- NSW experienced Australia’s largest outbreaks of ND between 1998 and 2000 in the Mangrove Mountain region. In 2002 another outbreak occurred in Greater Western Sydney
- the 1998–2000 and 2002 outbreaks are thought to be a result of mutations in one or more avirulent strains circulating in wild birds
- PPMV-1 was first detected in pigeons in 2011 but the virus has not yet been detected in any poultry species
- vaccination against NDV is available and comprises both live and inactivated vaccines.

Epidemiology

Modes of transmission

Transmission occurs via:

- **inhalation** of secretions or **ingestion via the faecal-oral route** (these are the main mode of transmission between birds)
- **vertical transmission** via embryonated eggs, although virulent strains will typically kill embryos before hatching occurs
- **fomites**, (clothing, equipment, vehicles, feed and water) as the virus survives in faecal material
- **mechanical vectors** in the form of rodents, insects and humans.

Disease dynamics

Disease presentation varies with the strain of virus and the host species and immune status. Typically:

- post-exposure, the incubation period of the virus is 2–15 days (but incubation periods of up to 21 days are possible)
- virus is shed during the incubation period, while clinical signs are present, and until early in the convalescent period
- virus is maintained in the environment by wild birds, gulls, waterfowl and shorebirds, some of which may be reservoir hosts for lentogenic pathotypes, which can become virulent following mutation in domestic poultry.
Persistence of the agent

NDV:
• is able to persist for extended periods of time in faecal material at room temperature and in water
• is inactivated by temperatures of 56°C for 3 hours or 60°C for 30 minutes, acid pH 2, and disinfectants such as formalin, chlorhexidine, sodium hypochlorite and oxidizing agents (removal of organic material prior to disinfection is important)
• can remain in eggs laid by infected chickens for months at room temperature, for more than a year at 4°C, and similar virus survival times have been observed on contaminated feathers and in contaminated premises.

Diagnosis and pathology

Clinical signs
Clinical signs of Newcastle disease are variable, depending on the pathotype. However, there is considerable overlap between pathotypes. Previous vaccination may greatly diminish the severity of clinical signs seen and may mask clinical disease. Poultry affected by the avirulent pathotype may display no clinical signs. Otherwise, clinical signs may include:
• anorexia, lethargy, ruffled feathers, respiratory distress, diarrhoea and conjunctivitis along with high mortality (velogenic viscerotropic pathotype)
• depression, ruffled feathers, increased respiratory effort, torticollis, ataxia and head tremors along with high mortality (velogenic neurotropic pathotype)
• respiratory and neurological disease, particularly in young birds, along with low mortality (mesogenic pathotype)
• mild respiratory disease, often in conjunction with other diseases such as mycoplasmosis (lentogenic pathotype).

Pathology
The most significant gross pathological lesions are seen with velogenic pathotypes. Lesions may include:
• swelling of the periorbital area or head
• haemorrhagic and oedematous conjunctiva
• haemorrhages of the tracheal mucosa and upper respiratory tract with diphtheritic membranes
• haemorrhage, oedema and/or ulceration throughout the gastrointestinal tract, particularly in the proventriculus and caecal tonsils at sites of lymphoid tissue accumulation
• haemorrhage of the ovaries
• increased opacity of air sacs
• splenomegaly with pale mottling.
Differential diagnosis

Clinical signs of ND are not pathognomonic and therefore other poultry diseases should be considered as part of the diagnostic process. Consider in the differential diagnosis:

- **exotic diseases**
  - duck plague (duck viral enteritis)
  - highly pathogenic avian influenza
- **endemic diseases**
  - aspergillosis
  - avian chlamydiosis
  - bacterial septicaemia (enterococcosis, staphylococcosis, streptococcosis, colibacillosis, pasteurellosis, erysipelas)
  - cellulitis of the comb and wattles (Escherichia coli and other bacteria)
  - fowl cholera (Pasteurella multocida)
  - infectious bronchitis
  - infectious coryza
  - infectious laryngotracheitis
  - mycoplasmosis
- **non-infectious causes**
  - acute poisoning
  - dehydration
  - heat stress
  - poor ventilation
  - smothering.

Samples required

Ideally, submit whole birds for post-mortem examination. In particular, examining whole birds may provide information on other differential diagnoses. Documenting gross lesions is also useful in determining the most appropriate samples for laboratory testing.

Sample collection

Collect:

- **sera** from live birds (to assess exposure to NDV or to determine vaccination titres)
- **oral and cloacal swabs** from sick and recently dead birds (for large numbers of samples, pools of three swabs may be submitted in viral transport media)
- **fresh tissue** samples from the trachea, lungs, spleen, caecal tonsils, bursa of multiple dead birds (keep intestinal tract samples separate from other tissue samples to avoid contamination with bacteria during virus isolation)
- **fixed tissue**, a full range of tissues (including the brain), in particular tissue showing gross pathological changes, fixed in 10% neutral buffered formalin.
Transport of samples
For transport:
- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Laboratory testing is required for the diagnosis of ND and is accomplished by one or more of the these procedures: haemagglutination inhibition (more common), ELISA, virus isolation, qPCR, pathotyping PCR, histopathology, immunohistochemistry.

Initial testing for AAvV-1 relies on qRT-PCR due the speed of testing and the high sensitivity and specificity of the test. Results are typically available within 24 hours of samples arriving at the laboratory.

Reporting requirements
Virulent ND is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect virulent ND, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
ND is a public health risk. It can cause conjunctivitis in humans, but the condition is usually self limiting. It is important to seek advice from government veterinary authorities, noting:
- biocontainment requires the use and appropriate disposal of personal protective equipment such as gloves, coveralls, rubber boots (or disposable boots), safety glasses and masks
- isolate sick birds
- exclude wild birds from the premises
- thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.
Further information


FIGURE 3.15.1 Marked haemorrhage evident in the comb, wattle and adjacent skin

Image credit: AFIP and CFSPH
FIGURE 3.15.2 Conjunctival haemorrhage that is most severe in the nictitans

Image credit: California Animal Health and Food Safety Laboratory System and CFSPH

FIGURE 3.15.3 Diphtheritic laryngo-tracheitis

Image credit: California Animal Health and Food Safety Laboratory System and CFSPH
CHAPTER 3.16

Nipah virus infection

Fast facts
Nipah virus (NiV) is a highly lethal zoonotic paramyxovirus primarily infecting pigs, horses and people.

Organism
NiV:
- replicates in the nasal cavity or nasopharynx
- is susceptible to drying and desiccation and most disinfectants.

Susceptible species
NiV affects:
- flying foxes
- pigs
- humans
- dogs, cats and horses.

Economic impact
There is potential for minor to moderate economic loss associated with disease control and human health care costs, as well as restrictions on animal movements and international trade.

Epidemiology
NiV:
- has an incubation period of 6–14 days in pigs
- has high rates of pig-to-pig transmission, causing high prevalence of infection on pig farms
- causes high mortality in humans.

Transmission
While data is limited on the role domestic species play in transmission to people, other transmission occurs via:
- the oronasal route (in pigs)
- contact with the secretions and bodily fluids of infected pigs (in pig-to-pig and pig-to-human transmission)
- spillover from flying foxes to pigs and humans (flying foxes can shed the virus in urine, saliva, and uterine fluids).

Clinical signs
Nipah virus can present with a wide range of clinical signs and a definitive diagnosis requires laboratory testing. Clinical signs in pigs include:
- increased body temperature
- respiratory signs (barking cough, laboured respiration, open-mouth breathing)
- neurological signs (behavioral changes, head-pressing, nystagmus, champing, uncoordinated gait, spasms, myoclonus and hindlimb paresis)
- early abortion.

Post-mortem
Post-mortem poses a high health risk for humans and should only be performed if strict biosecurity can be achieved. Field post mortems should not be attempted. There are no pathognomonic signs on gross pathology (but on histopathology, endothelial syncytia are highly indicative).
**Samples**

Collect:
- blood samples (EDTA and plain tubes)
- swabs—nasal, oral, rectal and urogenital swab or urine
- a blood clot from the jugular (if safe to do so).

**Actions to take**

If you suspect a case of NiV infection:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- minimise contact with suspect animals and wear adequate personal protective equipment
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

**Public health considerations**

- NiV infection is a zoonotic disease
- people may become infected after close contact with infected pigs
- you must consult the relevant state or territory health authorities as soon as NiV infection is strongly suspected or confirmed.
Introduction

Nipah virus (NiV) is an exotic highly lethal zoonotic paramyxovirus. In the largest outbreak to date, it primarily infected pigs but had low mortality, causing either respiratory or neurological signs. It has a broad mammalian host range and can cause severe disease, particularly in humans.

Disease agent and susceptible species

NiV:
- a nonsegmented negative strand RNA virus, NiV belongs to the genus *Henipavirus* of the family *Paramyxoviridae*
- to date there have been two identified strains—Nipah virus-Malaysia (NiV-MY) and Nipah virus-Bangladesh (NiV-BD)
- infection does not clinically affect flying foxes (*Pteropus* spp.), the natural reservoir of NiV
- infection (natural cases) has been reported in pigs, cats, dogs, horses and humans.

Public health considerations

- people may become infected after close contact with infected pigs
- when in close contact with infected pigs use personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots), goggles or safety glasses, and P3 respirators or N95 masks
- consult the relevant state or territory health authorities in the case of a NiV infection outbreak.

Distribution

NiV:
- has been recorded in pigs in Malaysia, and in people in Malaysia, Singapore, Bangladesh and India (a number of states including West Bengal, and Kerala)
- has been found in flying fox (*Pteropus* spp.) populations in multiple locations in South-East and southern Asia
- may have caused an outbreak of disease in horses and people in the Philippines in 2014
- has been isolated from flying foxes in Cambodia but no outbreaks have been reported from that country to date.

For the latest information on the distribution of NiV infection, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int](http://www.oie.int).

Occurrences in Australia

None recorded.
Epidemiology

Modes of transmission

Transmission occurs through:

- **spillover from flying fox populations**—infection in pigs most likely occurs after close contact with flying fox urine, faeces, uterine fluids and fruit spats (chewed up fruit). Consuming contaminated date palm sap or contaminated fruit has been linked to recurrent spillovers of NiV into the human population in Bangladesh (*flying fox to pig, flying fox to human*).
- **direct contact between pigs (pig to pig spread)**
- **direct human contact** with excretions (urine, blood) of infected animals. Pig farm workers and abattoir workers have the highest risk of pig-to-human spread. Preparation and consumption of infected horse meat implicated in Philippines outbreak (*horse-to-human spread*)
- **fomites**—as NiV may survive on fomites for hours under mild climatic conditions (*pig to human, pig to pig spread*)
- close contact associated with sick or dead humans (*human-to-human spread*).

Disease dynamics

NiV is thought to infect susceptible species through the naso-oral route, with local replication in the nasal cavity or nasopharynx followed by systemic spread of the virus. In addition:

- infection of vascular endothelial cells, damage to vascular endothelium and subsequent vasculitis is thought to precipitate disease in the various organ systems (for example, respiratory, neurologic, gastrointestinal). As the disease progresses, you may see predominantly respiratory and/or neurological signs
- pyrexic and clinically-affected pigs shed virus from the nasal secretions and excretions such as urine and blood. Therefore, post-mortem examinations of pigs that have died from acute NiV infection are high-risk for pig-to-human transmission
- viral shedding in respiratory secretions of subclinical pigs has been demonstrated experimentally
- available data suggests incubation period in pigs is between 6–14 days
- while morbidity is high in pigs, mortality is low
- in other species, such as horses and cats, mortality is high.

Persistence of the agent

NiV:

- has a lipid envelope and outside the host is susceptible to desiccation and changes in temperature
- survival in the environment varies from several hours to several days, depending on the environmental conditions (for disease control purposes, 5 days is presumed to be the maximum survival time under optimal environmental conditions)
- disinfectants include soap and detergents, Virkon®, hypochlorites, iodophors/iodine, biguanidines (e.g. chlorhexidine) and quaternary ammonium compounds.
Diagnosis and pathology

Clinical signs
A high incidence of subclinical infection in pigs was reported during the outbreak in Malaysia. The main presenting sign was respiratory disease, with some developing encephalitis. Clinical signs in pigs are:
- increased body temperature
- early abortion
- respiratory signs including barking cough, rapid or labored respiration, open-mouth breathing, pulmonary congestion and oedema on post-mortem examination
- terminal nasal discharge which may be a stable white or blood-stained froth
- neurological signs (including behavioral changes, head-pressing, nystagmus, champing, uncoordinated gait, spasms, myoclonus, and hindlimb paresis.

In humans, NiV causes an influenza-like illness, encephalitis and/or respiratory syndrome.

Pathology
The following are seen:
- generalised vasculitis
- dilated pulmonary lymphatic vessels
- pulmonary oedema
- exudate in airways
- (less frequently) oedema of the mesentery, increased pleural and pericardial fluid and thick stable foam in the airways.

Differential diagnosis
Consider in the differential diagnosis of NiV infection in pigs:
- exotic diseases
  - Aujeszky’s disease (in adult animals)
  - classical swine fever (chronic form)
  - Japanese encephalitis
  - porcine reproductive and respiratory syndrome
  - rabies lyssavirus infection
  - post-weaning multisystemic wasting syndrome (in weaned pigs)
- endemic diseases
  - anthrax
  - Australian bat lyssavirus infection
  - bacterial infection
  - botulism
  - Hendra virus infection
  - West Nile virus infection (Kunjin virus)
non-infectious causes
- snake bite
- toxicity
- trauma.

Samples required
Take stringent precautions during sampling. Personnel must wear full personal protective equipment with P3 respirator when they are in contact with animals strongly suspected to be infected with Nipah virus. Sample susceptible animals that have had close contact with a known infected animal three times—as soon as possible, 12 days post-contact and 20 days post-contact.

Sample collection
Only perform post-mortem and tissue collection if you have appropriate workplace health and safety controls in place. Do not perform post-mortem and tissue collection in the field.

Collect:
- EDTA blood (7–10ml per animal)
- swabs, nasal, oral, rectal and urogenital swabs or urine
- serum in the form of plain/clotted whole blood.

Transport of samples
For transport:
- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Laboratory testing is required for the diagnosis of NiV and is accomplished by one or more of the following:
- ELISA
- virus neutralisation test
- PCR.

The most rapid, sensitive and specific diagnostic procedure is the detection of viral nucleic acid in blood or swabs by qRT-PCR.
Reporting requirements

NiV infection is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect NiV infection, report the disease by phoning the **Emergency Animal Disease Watch Hotline on 1800 675 888**, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

NiV infection is a serious public health risk. Post-mortem examinations of animals that have died from acute NiV infection are a high-risk for animal-to-human transmission. Therefore it is important to seek advice from government veterinary authorities, noting:

- stringent biosecurity procedures must be enforced to protect human health
- sick/dead animals should be isolated from people as soon as possible. Limit human contact with the sick/dead animals and other potentially exposed animals on the property until NiV diagnostic testing has been performed. Stop or limit animal and human movements onto and off the property (if possible)
- establish ‘clean’ and ‘dirty’ areas to reduce transmission of disease throughout the property
- wear personal protective equipment in the dirty area, including impervious rubber boots, splash-proof overalls, disposable impermeable gloves, face shield or safety eyewear and a particulate respirator
- on leaving the dirty area, you must remove your personal protective equipment and double bag it in clinical waste bags with Virkon® disinfectant. You must disinfect all equipment thoroughly, and wash your hands and other exposed skin thoroughly. Shower and change your clothes before coming into contact with other animals
- anyone who has had unplanned or accidental close contact with a suspect NiV case must wash with soap and water and seek medical advice as soon as possible

Further information


CHAPTER 3.17
Peste des petits ruminants

Fast facts

Peste des petits ruminants (PPR) is a highly contagious exotic acute viral disease of small ruminants which primarily affects sheep and goats.

Organism

Peste des petits ruminants virus (PPRV):
- enters through the upper respiratory tract epithelium
- does not survive long outside the host
- is inactivated by a range of disinfectants.

Susceptible species

PPR affects:
- small ruminants, with goats more severely affected than sheep
- captive and wild ungulates and camels.

Economic impact

There is potential for significant economic loss associated with high morbidity and mortality in small ruminants, as well as restrictions on animal movements and international trade.

Epidemiology

PPR:
- incubation period is usually 4–6 days but this may range from 3–10 days
- animals are infectious during the incubation period and for up to 12 days after clinical signs first appear.

Transmission

Transmission is:
- mainly via close-contact airborne transmission (e.g. respiratory aerosols)
- via all secretions and excretions
- via fomites.

Clinical signs

Affected animals may suffer from peracute, acute or subacute disease.

Signs include:
- pyrexia, depression and sudden death with high mortality (peracute form)
- serous naso-ocular secretions, inappetence, diarrhoea, dysentery and ulcerative and necrotic erosive lesions on mucous membranes, pneumonia (acute form)
- milder signs in the subacute form.

Post-mortem

This may be found:
- dehydrated carcasses
- inflammatory, necrotic and erosive lesions of the oral and nasal cavities, gastrointestinal tract and less commonly the genital mucous membranes
- ileocecal valve congestion
- splenomegaly and engorgement of mesenteric lymph nodes
- primary and secondary bronchopneumonia.

Samples

Collect:
- swabs of conjunctival discharge, nasal, buccal and rectal mucosa and lymph nodes (especially the mesenteric and bronchial nodes)
- whole blood in EDTA and plain tubes
- full range of tissue including lungs, intestine, spleen, liver and kidneys in 10% neutral-buffered formalin.

Actions to take

If you suspect a case of PPR:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Peste des petits ruminants (PPR) is a highly contagious acute exotic disease of small ruminants caused by the peste des petits ruminants virus (PPRV). The disease is characterised by pyrexia, necrotic stomatitis, enteritis, bronchopneumonia and high morbidity and mortality.

Disease agent and susceptible species

PPRV:
- is a non-segmented, enveloped, negative-sense RNA virus belonging to the genus Morbillivirus in the family Paramyxoviridae
- has four genetic lineages (1–4) and multiple strains
- is antigenically similar to rinderpest virus (which was eradicated globally in 2011)
- hosts are principally sheep and goats, with goats more severely clinically affected
- causes sub-clinical infection and seroconversion in cattle, pigs and buffalo (but these animals are not known to play a role in transmission)
- infects other ungulates (including water buffalo and camels) and cause outbreaks of clinical disease in captive and wild ruminants
- infects deer under experimental conditions
- causes sub-clinical infection in camels or a mild form of the disease characterised by respiratory signs, diarrhoea, weight loss and decreased milk production
- is transmitted from camels to other camels and goats (but not to sheep)
- causes sub-clinical and clinical infection in captive and wild ungulates (including from the families Gazellinae, Tragelaphus, Antilopinae and Caprinae)
- does not affect humans.

Distribution

PPR:
- occurs in most of sub-Saharan Africa, North Africa, the Arabian Peninsula, the Middle East and South Asia
- has expanded to east Asia (including Vietnam, China and Mongolia) and the European continent, with reports of PPR disease events in the Republic of Georgia in 2016.

For the latest information on the distribution of PPR, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

None reported.
Epidemiology

Modes of transmission
Transmission occurs via:
- **direct contact** between small ruminants and inhalation of fine droplets released during coughing and sneezing is considered an important mode of transmission
- **indirect contact** with infected urine/faeces
- **semen and embryos** (potentially)—while the ability of PPRV to transmit via artificial breeding is unknown, rinderpest virus (which is a similar virus) can be transmitted in reproductive material. The most likely risk entry pathway to Australia is the illegal importation of infected semen or embryos for artificial breeding purposes
- **fomite** transmission is possible.

Disease dynamics
- the incubation period lasts from 3–10 days, usually lasting 4–6 days
- PPRV can be shed during the incubation period, and animals often remain infectious for 10–12 days after clinical signs first appear
- often produces prolonged convalescence and surviving animals develop lifelong immunity
- may cause peracute, acute or subacute clinical disease and disease severity depends on factors including species, breed, age and immune status
- can cause morbidity and mortality of 100 per cent in immunologically naïve flocks.

Persistence of the agent
PPRV:
- does not typically survive for long periods in carcasses or the environment
- is rapidly deactivated by UV light and desiccation but can survive for longer periods in chilled and frozen tissues
- is a large, lipid-based, enveloped virus susceptible to a range of disinfectants including alkalis, halogens, phenolic compounds, citric acids, alcohol and ionophores.

Diagnosis and pathology

Clinical signs
Disease may be peracute, acute or subacute. There is no chronic form of this disease. Infections are most commonly seen in animals of all ages in immunologically naïve herds, and in animals older than 4–6 months in endemic regions. This is associated with waning of maternal immunity.

**Peracute**
Clinical signs include pyrexia, serous nasoocular secretions, mucous membrane congestion, diarrhoea, depression and sudden death.
Acute
Disease duration ranges up to 14 days and signs include:
• pyrexia, depression, inappetence, serous to mucopurulent nasoocular discharge, mucous membrane congestion, particularly in the buccal cavity
• mucopurulent discharge can cause eyelid matting and obstruct nostrils which may result in dyspnoea
• ulcerative and necrotic erosive lesions on the hard palate and other mucous membranes (appearing approximately 5 days after initial clinical signs), with gums becoming hyperaemic
• caseous fibrin deposits found on the tongue with associated hypersalivation and halitosis
• primary or secondary respiratory infections, including pneumonia
• profuse watery or haemorrhagic diarrhoea which may contain flecks of intestinal epithelium
• dehydration and emaciation
• abortion
• erosion and crusting of the lips, eyes, nose and vagina or vulva
• death, which may occur within 5–10 days of the onset of clinical signs.

Sub-acute
Sub-acute cases are more likely to occur in sheep in endemic regions and sub-clinical animals can act as viral shedders (but there is no long-term carrier state). Severe immunosuppression in sub-clinical animals may result in secondary infections such as pasteurellosis and *Mannheimia haemolytica* pneumonia.

Pathology
Findings may include:
• emaciated and dehydrated carcass
• well-demarcated, inflammatory, necrotic and erosive lesions of the oral and nasal epithelium, gastrointestinal tract and (less commonly) the genital mucous membranes
• linear engorgement and blackening of folds in the abomasum, caecum, proximal colon and rectum
• congestion of the ileocecal valve
• congestion and villi sloughing of the intestinal tract with mucosal haemorrhage, well-demarcated areas of ulceration or necrosis (erosive lesions are common in the abomasum)
• gastrointestinal necrosis of the mucosa and necrosis and collapse of Peyer's patches
• hypertrophy, necrosis and haemorrhage of mesenteric lymph nodes and splenic atrophy and congestion
• enlarged spleen with petechial haemorrhages, red hepatisation, oedema of lymph nodes, especially the mesenteric lymph nodes
• pulmonary congestion, consolidation, and raised patches of emphysema on the lung
• erosions and petechiation of the nasal mucosa, turbinates, larynx, trachea and bronchopneumonia.
**Differential diagnosis**

Consider in the differential diagnosis of PPR in small ruminants:

- **exotic diseases**
  - bluetongue disease (infection present in Australia but without clinical disease)
  - foot-and-mouth disease
  - contagious caprine pleuropneumonia
  - heartwater (infection with *Ehrlichia ruminantium*)
  - rinderpest (now recognised as globally eradicated by the OIE)
  - sheep and goat pox

- **endemic diseases**
  - contagious ecthyma (orf)
  - malignant catarrhal fever (bovine herpesvirus 2 is present in sheep Australia)
  - coccidiosis
  - pasteurellosis
  - *Mannheimia haemolytica*

- **non-infectious causes**
  - mineral poisoning.

**Samples required**

**Sample collection**

Collect samples from clinically affected animals, ideally during the pyrexic stage, before the onset of diarrhoea. Alternatively, collect samples from animals that have recently died, or immediately after euthanised for post-mortem.

Samples to collect for diagnostic testing are:

- **serum**, from at least 10 animals (if possible) towards the end of the acute phase or from recovered animals
- **EDTA blood**, from live, clinically affected animals (7–10 ml/animal) during the acute phase (preferably from pyrexic animals)
- **fresh tissue**, including lymph node (especially the mesenteric and bronchial nodes), lungs, spleen, tonsils, tongue, affected areas of the alimentary tract (2 g of each tissue)
- **fixed tissue**, comprising a full range of tissues including lungs, intestine, spleen, mesenteric lymph nodes, liver and kidneys (in 10% neutral-buffered formalin)
- **swabs**, of conjunctival discharge as well as nasal, buccal and rectal mucosa. Collect from live, clinically affected animals (early during the acute phase) and preferably from pyrexic animals.
Transport of samples

For transport:

- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- **DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests**
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests

For PPR:

- available serological tests include competitive ELISA, indirect ELISA and viral neutralisation
- antigen detection tests include agar gel immunodiffusion test and an ELISA
- nucleic acid detection includes real-time reverse transcription polymerase chain reaction (real-time RT-PCR). These tests can differentiate the lineages of PPRV from the vaccination lineage
- viral isolates can only be detected during acute infection which is usually within days from the onset of clinical signs. Tests include viral isolation, antigen capture ELISA and lateral flow immunoassay.

Reporting requirements

PPR is an OIE-listed disease and Australia has an international obligation to report confirmed cases. If you suspect PPR, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for PPR, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.
Further information


CHAPTER 3.18

Porcine epidemic diarrhoea

Fast facts
Porcine epidemic diarrhoea (PED) is an exotic viral disease of pigs characterised by diarrhoea and weight loss, and high piglet mortality.

Organism
PED virus (PEDV):
• replicates primarily in the small intestine, in villous enterocytes
• virulence varies with the strain of virus
• survives for 28 days at 4°C and 7 days at 25°C, and will not survive for longer than 7 days at room temperature
• is inactivated by most disinfectants.

Susceptible species
Only pigs are affected.

Economic impact
There is potential for significant economic loss due to the high number of mortalities in young animals, as well as restrictions on animal movements and international trade.

Epidemiology
PED:
• has an incubation period ranging from 1–4 days
• causes morbidity of up to 100 per cent, (particularly in a naïve herd)
• causes mortality of up to 100 per cent in neonatal piglets, and 5 per cent in adult pigs.

Transmission
Transmission occurs:
• via the faecal-oral route
• via fomites
• via contaminated pig blood products.

Clinical signs
Signs include:
• diarrhoea, often watery
• vomiting
• anorexia
• dehydration
• depression.

Post-mortem
You may see thinning of the intestines and fluid intestinal contents.

Samples
Also refer to the detailed section on sampling for pigs with diarrhoea.
• whole blood in EDTA and plain tubes
• faeces in sterile tubes
• full range of tissue including lungs, intestine, spleen, liver and kidneys fresh and in 10% neutral-buffered formalin.

Actions to take
If you suspect a case of PED:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Porcine epidemic diarrhoea (PED) is an exotic viral disease of pigs which predominantly affects neonatal pigs. The disease is of particular concern in naïve herds (such as those in Australia) in which the disease may cause large outbreaks with near 100 per cent mortality in younger animals. Significant outbreaks have recently been seen in previously unaffected countries (such as the United States) resulting in 80 per cent mortality in suckling pigs and declines in pork production of 6–7 per cent.

Disease agent and susceptible species

PED virus (PEDV)

- is an enveloped, single-stranded RNA virus in the *Alphacoronavirus* genus within the family *Coronaviridae*
- can affect pigs already exposed to other porcine coronaviruses (transmissible gastroenteritis coronavirus and hemagglutinating encephalomyelitis virus) as exposure to these viruses does not provide cross protection to PED
- only affect pigs (*Sus scrofa*) and disease has only been detected in domestic pigs (the susceptibility of wild pigs is unknown).

Distribution

PED:

- was first detected in 1971 in the United Kingdom and subsequently caused epidemics of disease throughout Europe and Asia in the 1970s and 1980s
- causes occasional epidemics in Europe (where the disease is endemic), with disease usually manifesting as diarrhoea in weaned and grower pigs
- continues to cause significant disease in piglets in Asia
- was reported for the first time in 2013 in the United States, and then spread throughout the United States and Canada causing large numbers of mortalities in young pigs.

Occurrences in Australia

None recorded.

Epidemiology

Modes of transmission

Transmission occurs via:

- **faeco-oral contamination** is the most important mode of transmission
- **fomites**, as a result of faecal contamination of inappropriately decontaminated vehicles and farm equipment, has been an important transmission route between farms in the North American epidemics.
Disease dynamics

After ingestion of viral particles in faeces:
- PEDV replicates in the intestinal epithelial cells within the small intestine and (to a lesser extent) in the large intestine. This leads to exfoliation of the enterocytes and blunting of intestinal villi and an osmotic and malabsorptive diarrhoea
- the incubation period of the virus is estimated at between 1–4 days
- PEDV may be excreted for up to 35 days after clinical signs are noted.

Persistence of the agent

PEDV persistence in the environment is dependent on the temperature, humidity and substrate, with cooler temperatures and moist conditions allowing the virus to persist for longer periods of time. PEDV:
- has been shown to survive for 7 days at 25°C in dry feed, and up to 14 days at 25°C in wet feed
- is inactivated at temperatures above 60°C
- is susceptible to inactivation by a range of disinfectants—formalin (1%), anhydrous sodium carbonate (4%), lipid solvents, iodophores in phosphoric acid (1%) and sodium hydroxide (2%).

Diagnosis and pathology

Clinical signs

Clinical signs vary with age and immune status, with younger pigs exhibiting more severe clinical signs. Signs include:
- watery diarrhoea
- vomiting
- anorexia
- depression.

Laboratory testing is required to exclude other causes of diarrhoea in pigs.

Pathology

Gross pathological lesions are usually confined to the gastrointestinal tract and may include:
- thinning of the intestine
- watery intestinal contents
- dilated intestinal loops
- undigested milk in the stomach and intestines of piglets
- faecal staining of the perineum.
Differential diagnosis

Differential diagnoses for PED are those which include diarrhoea as part of the clinical syndrome. Consider these diseases:

- **exotic diseases**
  - [African swine fever](#)
  - [classical swine fever](#) (hog cholera)
  - porcine deltacoronavirus
  - post-weaning multisystemic wasting syndrome
  - [transmissible gastroenteritis](#)

- **endemic diseases**
  - [Clostridium spp.](#)
  - coccidiosis
  - colibacillosis
  - internal parasites
  - porcine circovirus 2 infection
  - proliferative enteropathy (ileitis)
  - rotavirus infection
  - salmonellosis
  - swine dysentery
  - yersiniosis

- **non-infectious causes**
  - copper poisoning
  - fluoride toxicity
  - fungal/toxic mould poisoning
  - iron toxicity in piglets
  - lead poisoning
  - mercury toxicity
  - organochlorine toxicity
  - organophosphate toxicity
  - overfeeding
  - pantothenic acid deficiency
  - paraquat poisoning
  - phosphorus toxicity
  - salt poisoning
  - zinc deficiency.
Samples required

Sample collection
Collect samples to investigate suspected endemic and exotic causes of gastroenteritis. Collect:
- **formalin fixed tissues**—a full range of tissues including the small and large intestine
- **fresh faeces and tissues**—small and large intestine
- **serum**, in the form of whole blood.

Transport of samples
For transport:
- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Laboratory testing is required to differentiate PEDV from other causes of diarrhoea in pigs. Real-time PCR is the preferred method for detecting viral nucleic acid, with a range of serological tests available for antibody detection. Other tests available include ELISA, virus isolation, immunohistochemistry, immunofluorescence and virus neutralisation.

Reporting requirements
PED is not an OIE-listed disease but is nationally notifiable within Australia. If you suspect PED, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
There are no public health implications for PED, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.
Further information


Porcine reproductive and respiratory syndrome

Fast facts
Porcine reproductive and respiratory syndrome (PRRS) is a highly contagious exotic viral disease of pigs.

Organism
PRRS virus (PRRSV):
- is present in most pig-producing countries
- is found in all tissues and body fluids
- is readily inactivated by heat and chemical disinfection and its environmental stability is poor.

Susceptible species
Only pigs are susceptible.

Economic impact
There is potential for significant economic loss associated with production losses, as well as restrictions on animal movements and international trade.

Epidemiology
- mortality varies with virulence of the virus isolate but is typically low
- the incubation period is 4–8 days but it may take longer to see signs in a herd
- prolonged (persistent) infection occurs.

Transmission
Transmission occurs via:
- direct contact, usually oronasal contact with infected fluid (e.g. saliva, oropharyngeal mucus, urine)
- contaminated fomites
- artificial insemination
- aerosol transmission.

Clinical signs
There are two syndromes: reproductive failure in sows and respiratory disease in young pigs. In boars and unbred replacement gilts and sows, transient pyrexia and anorexia may be observed.

Post-mortem
- the most common lesions are interstitial pneumonia and generalised lymphadenopathy
- lesions are more pronounced in younger animals and may be exacerbated by concurrent bacterial infection
- gross lesions are typically absent or non-specific in aborted foetuses.

Samples
Also refer to the detailed section on sampling for pigs with reproductive problems.

Samples from younger pigs are preferred. Collect:
- blood (EDTA and plain tubes)
- semen, oral swabs
- tissue from tonsils, heart, spleen, lymph nodes, kidney, lung.

Actions to take
If you suspect a case of PRRS:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Porcine reproductive and respiratory syndrome (PRRS) is a highly contagious exotic viral disease (arterivirus) of pigs characterised by reproductive failure (abortions), death in piglets (stillborn and weak animals) and respiratory disease in all age groups. Other names applied to the syndrome include blue ear, swine infertility and respiratory syndrome, and pig high-fever disease.

Disease agent and susceptible species

PRRS virus (PRRSV) has only one known host species—domestic and feral pigs (Sus scrofa). PRRSV:

- is an enveloped RNA virus (genus Arterivirus, family Arteriviridae)
- has two related but antigenically distinguishable genotypes: type 1 (European genotype) and type 2 (American genotype). The latter predominates in North America and Asia
- within the two genotypes are multiple distinct variants that differ in infectivity, pathogenicity and epidemiology, and rapid genetic change and emergence of further novel strains is expected in the future.

Distribution

PRRS is reportedly the most prevalent disease of pigs in the world and:

- is endemic in most major pig-producing countries, with the highest economic impact in North America
- has been successfully eradicated a number of countries but has since recurred in some of these
- severe outbreaks (attributed to highly pathogenic PRRSV) have had a dramatic effect on pig health in China and Vietnam.

For the latest information on the distribution of PRRS, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

None reported.

Epidemiology

Modes of transmission

Transmission occurs via:

- **direct contact with infected pigs** (oral or intranasal) and their saliva, nasal secretions, urine, milk, colostrum and faeces
- **vectors**—insects such as biting flies and mosquitoes can play a role in the transmission of the virus
- **reproduction** involving infective semen and vaginal secretions
- **vertical transmission** from viraemic dams to foetuses, resulting in foetal death or birth of infected pigs that are weak or appear to be normal. The virus crosses the placenta efficiently only in the last trimester of pregnancy
- **airborne spread**—aerosol transmission has been reported.
Disease dynamics

The maintenance of infection within herds is facilitated by persistent infection and the virus:

- infects macrophages at the site of infection (mucosal surfaces), followed by extension to regional lymphoid tissue and rapid viremia. Within 12 hours of experimental intranasal inoculation, virus is detectable within macrophages within the nasal mucosa, tonsils and lungs, as well as respiratory epithelial cells
- impairs host immunity, resulting in increased susceptibility to opportunistic pathogens
- incubation period is 4–8 days but clinical signs may take longer to emerge in a herd
- has been detected in serum and oropharyngeal mucus for 3 and 22 weeks, respectively
- is shed for an extended period in saliva (42 days), urine (14 days) and semen (43 days)
- has been recovered from the oropharyngeal area 157 days after experimental infection
- pigs have transmitted disease to commingled susceptible sentinel pigs 22 weeks after originally being infected and viral RNA has been demonstrated as late as 210 days after birth in the serum of pigs infected in utero.

Persistence of the agent

PRRSV:

- cannot withstand drying, so does not persist on equipment or other fomites beyond 1 day
- is most stable between pH 5.5 and pH 6.5; infectivity is reduced by over 90 per cent at a pH lower than 5 or higher than 7
- is unstable in low concentrations of detergents and is rapidly inactivated by lipid solvents and chlorine based disinfectants
- can survive in water for 11 days
- has been isolated from muscle and lymphoid tissue
- survives freezing in cell culture for a prolonged period and has been isolated after 1 month from muscle frozen at -20°C, but levels of virus decrease with cooling, hardening and freezing
- can be recovered from muscle tissues 0–24 hours after slaughter but not from muscle held at 4°C for 48 hours
- will survive in bone marrow for several weeks when stored at 4°C
- is inactivated by heat.
Diagnosis and pathology

Clinical signs
PRRS is a severe systemic disease characterised by profound viraemia (often prolonged), and virus dissemination and replication in multiple organs. The clinical presentation of PRRS is highly variable with subclinical, mild or severe disease which may be associated with secondary infections. Diagnosis is strongly reliant on recovery and identification of the infective agent. In addition:

- in the field, the incubation period is variable, ranging from 3–37 days but commonly is between 4–8 days
- clinical manifestations vary greatly and depend on the strain of virus, the age and immune status of the pigs affected, the presence of co-infection, and husbandry factors
- clinical syndromes associated with PRRSV infection can be broadly divided into late-term reproductive failure and respiratory disease
- in endemically infected herds, PRRS presentation varies from asymptomatic infection to endemic respiratory disease, ongoing foetal loss or failure to thrive in weaners and grower-finished pigs
- highly pathogenic PRRSV causes more severe disease characterised by a persistent marked pyrexia and anorexia with red cutaneous discolouration and blue ears
- infection with PRRSV is a predisposing factor for bacterial pneumonia and septicaemia, and the clinical scope of disease may be complicated by co-infection with one or more concurrent pathogens.

Clinical signs in naïve herds include:
- anorexia and lethargy in all age groups
- reproductive losses including sporadic abortions or abortion storms, stillbirth, mummified foetuses and weak neonatal piglets
- high pre-weaning mortality in neonatal piglets, which are listless, weak and dyspnoeic
- pneumonia characterised by dyspnoea and respiratory distress in nursery and grower/finisher pigs
- ‘sow abortion and mortality syndrome’ resulting in up to 10 per cent sow mortality.

Pathology
As with clinical presentation, gross and histologic changes are poorly defined, and tend to vary with the strain of the virus and age of the affected animal. In addition:

- post-natal PRRSV infection is characterised by interstitial pneumonia and generalised lymphadenopathy
- post mortem changes consistent with interstitial pneumonia include failure of the lungs to collapse, and a firm and rubbery texture with rib impressions
- lymph nodes are variably enlarged and white or tan coloured, consistent with lymphoid hyperplasia
Differential diagnosis

The clinical picture of PRRS is not specific and varies with PRRSV strain. In the field, suspicion of PRRS is based on clinical signs of reproductive failure and high levels of neonatal mortality. Analysis of farm records will provide helpful information.

Consider in the differential diagnosis:

- **exotic diseases**
  - African swine fever
  - Aujeszky’s disease
  - classical swine fever
  - foot-and-mouth disease
  - Nipah virus infection
  - porcine enterovirus infection
  - postweaning multisystemic wasting syndrome
  - influenza A viruses in pigs (exotic strains)

- **endemic diseases**
  - enzootic pneumonia
  - haemagglutinating encephalomyelitis virus infection
  - haemophilus parasuis infection (Glässers disease)
  - leptospirosis
  - porcine parvovirus infection
  - porcine respiratory coronavirus infection
  - proliferative and necrotising pneumonia
  - syncitial pneumonia and myocarditis
  - influenza A viruses in pigs (human-origin strains, subtypes H1N1, H1N2 and H3N2)

- **non-infectious causes**
  - carbon monoxide poisoning
  - ingested toxins (e.g., pyrrolizidine alkaloids, paraquat)
  - poor air quality such as high levels of dust and ammonia
  - poor management
  - poor nutrition
  - zearalenone poisoning.
Samples required
Samples from younger rather than older animals are preferred as the virus is generally detected in higher amounts and for a longer time in younger pigs.

Sample collection

Collect:
- **fixed tissues** in neutral-buffered formalin taken from recently euthanased affected pigs and from pigs that have recently died. Include cranioventral lung, tonsil, lymph nodes, heart, brain, thymus, spleen and kidney tissue and any lesions detected on post-mortem examination. Process tissues within 48 hours of fixation to avoid degradation of PRRSV antigens and loss of immunohistochemical positive cells
- **fresh tissue** from spleen, gastro-hepatic lymph node, mesenteric lymph node, tonsils, lung, kidney, ileum (2 g of each tissue)
- **EDTA blood**—7–10 ml/animal from live, clinically affected animals
- **serum** from at least 20 exposed animals in the herd. You can also detect antibody in oral fluid from ropes, though this method is not as sensitive
- **other samples**—oral swab, semen, umbilical cords and umbilical cord blood, tonsils, lung, thymus, bronchial lymph nodes, heart, kidney, spleen, muscle transudate. Samples from live born pigs are preferred, as samples from mummified or aborted litters are unlikely to yield virus for agent detection.

Transport of samples
For transport:
- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Confirm the diagnosis by laboratory testing as there is considerable overlap in the clinical and pathological signs seen in PRRS with those of a number of other diseases (see differential diagnosis). For PRRS diagnosis the tests available are:
- screening by real-time PCR (qPCR) using a multiplexed real-time PCR that detects both type 1 and type 2 viruses
- an indirect enzyme-linked immunosorbent assay (ELISA) is used for serological screening
- immunoperoxidase detection on virus-infected cells is used as a confirmatory test
- histopathology and immunohistochemical staining.
Reporting requirements

PRRS is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect PRRS, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for PRRS, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


Fast facts

Rabies is a zoonotic neurological disease almost invariably fatal to its hosts. Non-bat variants of lyssavirus (including rabies lyssavirus) are exotic to Australia and their introduction would have major public health implications.

Organism

Rabies lyssavirus belongs to the genus Lyssavirus, family Rhabdoviridae. In addition:
- the Lyssavirus genus includes 14 other recognised species and other proposed variants, including Australian bat lyssavirus (ABLV)
- lyssaviruses are readily inactivated by all main disinfectants and by ultraviolet light and heat treatment.

Susceptible species

Lyssaviruses infect all mammals, including humans (infections are almost invariably fatal).

Economic impact

An incursion of rabies lyssavirus would result in moderate to severe economic costs associated with intensive vaccination campaigns, as well as restrictions on animal movements.

Epidemiology

Rabies lyssavirus:
- is transmitted through saliva by the bite of an infected animal showing clinical signs
- is shed from saliva during clinical stages and can be shed for a short period before clinical signs are seen
- has a case fatality close to 100 per cent in humans and other mammals
- has an incubation period of normally around 2–6 weeks.

Transmission

Rabies lyssavirus is transmitted by the bite of an infected animal during clinical disease. Other lyssaviruses may be transmitted in a similar manner.

Clinical signs

Clinical signs of rabies lyssavirus and other lyssavirus infections are highly variable but all are attributable to the neurological effects of infection. Signs in animals fall into six main categories:
- excitation (aggression, hyperaesthesia, wandering, restlessness, self-inflicted trauma)
- paralysis (ataxia, paresis, dysphagia, facial asymmetry)
- loss of normal social and behavioural responses (personality changes and unusual tameness of wild animals)
- unusual vocalisation
- pica
- coma.
Post-mortem
There are no specific gross post-mortem changes.

Samples
- brain tissue (or entire head)
- (if brain is not available) spinal cord, ganglia, peripheral nerves and salivary gland.

Keep samples cold for storage and transport. If there is no cold chain, place samples in formalin.

Actions to take
If you suspect a case of rabies due to rabies lyssavirus or other lyssaviruses such as ABLV:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- confine and isolate suspected animal cases where it is safe to do so. Alternatively, destroy the suspected cases and submit the brain material (or entire head) for laboratory confirmation. Implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

Public health considerations
- rabies caused by rabies lyssavirus and other lyssaviruses is a zoonotic disease
- people may become infected after being bitten or scratched by an infected animal or through close contact with saliva from an infected animal
- if a person is bitten, scratched or has close exposure to saliva from an infected animal, the bite or scratch wound should be washed thoroughly with soap and running water for 15 minutes and contact human health authorities immediately to ensure post-exposure prophylaxis is provided
- consult the relevant state or territory health authorities as soon as rabies is strongly suspected or confirmed.
Introduction

Rabies is a zoonotic neurological disease, which is almost invariably fatal to its hosts. Non-bat variants of lyssavirus (including rabies lyssavirus) do not occur in Australia and their introduction would have major public health implications.

Disease agent and susceptible species

Rabies is caused by infection with rabies lyssavirus and related lyssaviruses. Rabies lyssavirus:
• belongs to the genus Lyssavirus, family Rhabdoviridae
• affects mammals, including humans (infections are almost invariably fatal)
• is maintained by various carnivore species (particularly domestic dogs) and bats.

Bats are the natural hosts of most lyssavirus species. As little is understood of the virus transmission cycles, all bats should be considered capable of supporting lyssavirus replication and shedding. Lyssaviruses:
• cause similar acute, fatal, neurological disease in mammals
• can infect all mammals, however; different viruses appear adapted to a particular host species (hence, dog rabies lyssaviruses are optimally adapted to dogs, and although other mammals can become infected, the transmission of infection in these species is less efficient)
• many species of lyssavirus are maintained in insectivorous bat species as well as in fruit bats
• in 2018, there were 14 virus species in the lyssavirus genus.

Public health considerations

• people may become infected after being bitten or scratched by an infected animal, or through close contact with saliva from an infected animal
• people in close contact with infected animals should be vaccinated against rabies
• when in close contact with infected animals use personal protective equipment such as bite resistant gloves and coveralls, rubber boots (or disposable boots), safety glasses and mask
• if personnel are bitten or scratched by a potentially infected animal that they thoroughly wash the wound with soap and running water for 15 minutes and immediately seek medical advice
• consult the relevant state or territory health authorities in the case of a rabies outbreak.
Distribution
Rabies lyssavirus:
- occurs throughout most of the world except in Antarctica, Australia, New Zealand, Papua New Guinea, Japan, much of western Europe and many of the small islands
- has persisted in wildlife species in many areas of the world where dog rabies has been controlled
- in wildlife has been eradicated over large areas of Europe and North America by distributing baits with oral vaccines
- has been eradicated (in dog-cycle rabies) in Europe, North America and Japan by using injectable vaccines, which have also reduced its frequency in many other regions.

Other lyssaviruses have been reported in Europe and Asia, Africa and Australia. Antibodies to ABLV have been reported from bats from the Philippines. It is possible that bat lyssaviruses occur wherever bats are present.

For the latest information on the distribution of rabies lyssavirus and other lyssaviruses, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia
In Australia:
- Rabies lyssavirus was suspected in several related cases in Tasmania in 1867. There have been a number of confirmed human cases in returning travellers from rabies endemic countries
- ABLV is present in flying-fox (Pteropus species) populations (Pteropus variant virus) and has been reported in a small number of insectivorous yellow-bellied sheathtailed bats (Saccolaimus flaviventris), (insectivorous variant virus)
- ABLV infection has been reported in three humans and two horses. The insectivorous variant virus was involved in one of the human cases and in the two horse cases. The Pteropus variant virus was involved in two of the human cases.

Epidemiology

Modes of transmission
Transmission occurs through:
- inoculation of saliva into bites, scratches and wounds—rabies lyssavirus is shed in saliva so bites (the most frequent means of transmission) and contact with saliva from infectious animals pose a risk of transmission. Two cases of human infection (leading to death from a disease indistinguishable from rabies) with ABLV have occurred following bites and scratches from infected bats
- ingesting carcasses of animals affected by lyssavirus. Animal products, and byproducts from an animal affected by lyssavirus should be regarded as potentially infectious and not permitted to enter the food chain
- inhalation of aerosol is also a potential route of transmission.
Disease dynamics

Current knowledge indicates that rabies lyssavirus and other lyssaviruses have essentially the same pathogenesis. After the inoculation of virus into a wound:

- the virus replicates in local tissues and within hours to days after a bite and soon afterwards invades the peripheral nerves. During the incubation period, the virus is transmitted by axons to the central nervous system (CNS). The CNS is invaded by virus, clinical signs become apparent, and the disease course becomes irreversible
- the incubation period (from bite to initial neurological signs) in all animals is rarely less than 10 days and is normally around 2–6 weeks (and in rare cases may exceed 6 months)
- the duration of the incubation period varies according to virus strain and dose, the distance of the bite site from the CNS, and the richness of the sensory innervation at the site of virus entry into the body
- no clinical, pathological or immunological signs are apparent during the incubation period
- the virus is shed in saliva from about the time of onset of clinical signs
- viral shedding coincides with behavioural changes that lead to biting of other animals
- disease, once it appears, is acute and progressive, invariably causing death
- there is a typical progression of excitatory to paralytic disease.

The following is known about lyssavirus hosts.

- Ecological factors determine epidemiology in different host species.
- Dog rabies lyssavirus (urban rabies) is dominant in areas (urban and rural) where there is a high proportion of unvaccinated and semi-owned or stray dogs.
- For rabies lyssavirus in wildlife species (sylvatic rabies), it is usually one wildlife species (e.g. foxes, raccoons, skunks or bats) in the ecosystem that is responsible for perpetuation of the disease.
- In dogs and other reservoir hosts, outbreaks of rabies can occur in local epidemics that last for several months.
- Cattle, sheep and other herbivores (spillover hosts) are usually infected with rabies lyssavirus following attacks by a reservoir host animal showing clinical signs of rabies. These spillover host animals do not play a role in perpetuation of virus transmission cycles.
- Vaccination markedly reduces the frequency of rabies lyssavirus in reservoir and spillover hosts. In reservoir hosts, an immunisation coverage of over 50 per cent of the population can result in a significant decline and if coverage exceeds 70 per cent this can lead to eradication of the disease.
- The transmission of ABLV to individual animals of other species is considered unlikely to result in the establishment of persistent transmission cycles in these species, as this would require adaptation of the virus strain to a new host species other than bats.

Persistence of the agent

Lyssaviruses:

- are comparatively fragile and do not retain infectivity for long outside the host
- are readily inactivated by all the main disinfectants (detergents, ethanol and iodine preparations) and by ultraviolet light and heat treatment.
Diagnosis and pathology

Clinical signs

CNS infection patterns, and therefore clinical signs, are varied but often include behaviour changes that lead to biting of other animals. The clinical signs of rabies caused by rabies lyssavirus can often be subtle and change rapidly. ABLV infection causes similar clinical signs in affected people and animals.

Clinical signs of disease in infected animals fall into six main categories:

- **excitation**—includes unprovoked aggression, over-reactive to external or perceived stimuli, aimless wandering, restlessness, self-inflicted trauma (scratching, biting)
- **paralysis**—can affect any of the motor systems, causing ataxia, knuckling of distal limbs, paresis, oral dysphagia and facial paralysis
- **loss of normal social and behavioural responses**—as wild animals ignore territoriality and encroach on urban areas, and nocturnal species become active by day. In companion animals, owners usually report personality change
- **unusual vocalisation**—whereby animals vocalise relentlessly. In dogs, muscular incoordination of the larynx causes a characteristic howling
- **pica**—can cause animals, particularly carnivores, to chew and swallow anything in their environment (such as soil, plants and bedding)
- **coma**—at the terminal stage of disease.

In addition, signs include: dehydration, red eyes, salivation, poor condition, unkempt pelage and signs of trauma. Cases of sudden death with few, if any, preceding signs have been reported.

Pathology

No remarkable gross pathological signs are usually evident, and when they are present are secondary to neurological effects. The following may be found in cases of rabies lyssavirus:

- dehydrated carcasses in poor physical condition, with signs of trauma (for example, broken teeth)
- signs of pica in carnivores (such as soil and plant material in the mouth and stomach).

Differential diagnosis

You should consider any other causes of neurological dysfunction as differential diagnoses for rabies lyssavirus or other lyssavirus infection. In all mammals the disease is acute, progressive and fatal. Where this is not the case, rabies generally can be excluded.

Consider in the differential diagnosis:

- **exotic diseases**
  - Aujeszky’s disease
  - Borna disease
  - bovine spongiform encephalopathy
  - eastern, western and Venezuelan equine encephalomyelitis viruses
  - Japanese encephalitis virus and other flaviviruses
  - scrapie
• endemic diseases
  – bacterial and mycotic diseases of the central nervous system, including listeriosis and cryptococcosis
  – canine distemper and infectious canine hepatitis
  – protozoal infections (such as babesiosis and toxoplasmosis)
  – West Nile-Kunjin virus infection
• non-infectious causes
  – foreign bodies in the oropharynx or oesophagus and other traumatic injuries
  – poisons, including ‘1080’ (sodium fluoroacetate), heavy metals (such as lead), chlorinated hydrocarbon and organophosphate pesticides, urea and nitrogen trichloride.

Samples required
Take stringent precautions when collecting samples (including wearing full personal protective equipment) due to the zoonotic risk. A current rabies vaccination is necessary when working with bats. Likewise, submit whole bats rather than attempting to remove the head or brain. Unvaccinated persons must not remove animal heads or brains.

For the laboratory confirmation of lyssavirus:
• the brain is the most important sample
• the brain stem is the most consistently reliable sample for detection of infectious virus or viral antigen (other regions of the brain, including the hippocampus, are negative in up to 5 per cent of rabid animals)
• if the brain is not present, other suitable tissues include spinal cord, ganglia, peripheral nerves (taken from points close to the CNS) and salivary gland.

Sample collection
Collect:
• fresh composite brain samples—including several different parts (the preferred samples) of the brain for diagnosis. Do not store in formalin as a fluorescent antibody test cannot be performed reliably on formalin-fixed samples
• fixed composite brain samples—particularly if there will be a delay in the samples reaching the laboratory
• serum—serological tests are useful for confirming vaccine responses in animals and humans and for epidemiological studies in wildlife species, including bats.

Transport of samples
For transport:
• chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days
• formalin fixed tissue can be sent at room temperature.
Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Laboratory testing is required for the diagnosis of lyssavirus infection and is accomplished by one or more of the following procedures: PCR, immunohistochemistry and fluorescent antibody test.

Reporting requirements
Rabies lyssavirus infection is an OIE-listed disease and Australia has an international obligation to report cases. ABLV is a nationally notifiable disease in both animals and humans. If you suspect rabies due to infection with rabies lyssavirus or ABLV, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
Rabies caused by rabies lyssavirus and other lyssaviruses including ABLV, is a public health risk. You must implement stringent biosecurity procedures to protect human health:

- isolate and confine suspect animal cases (where it is safe to do so) or destroy animals and submit the brain material (or entire head or whole bat) for laboratory confirmation of lyssavirus infection
- wear protective (bite resistant) gloves, glasses, mask and coveralls, and avoid animal bites, aerosols and scratches when handling potentially infected animals
- if working with potentially infected animals, personnel must be vaccinated against rabies virus
- immediately seek medical advice if personnel are bitten or scratched by a potentially infected animal
- post-exposure prophylaxis must be rapidly initiated before the virus can replicate in the central nervous system. This consists of three components—washing and disinfection of the exposure site for at least 15 minutes, immunisation with vaccine, and immunoglobulin administration
Further information


CHAPTER 3.21

Rift Valley fever

Fast facts

Rift Valley fever (RVF) is an exotic acute, arthropod-borne zoonotic viral disease principally affecting domestic ruminants and people.

Organism

RVF virus (RVFV):
- some isolates differ in virulence
- is killed by strong sunlight and a range of disinfectants.

Susceptible species

RVF affects sheep and goats (highly susceptible), cattle, camels and humans.

Economic impact

There is potential for significant economic loss due to high morbidity and mortality in ruminants, as well as restrictions on animal movements and international trade.

Epidemiology

RVF:
- outbreaks occur over large areas following heavy rains and flooding
- the incubation period ranges from 1–30 days
- morbidity depends on which species are affected.

Transmission

Transmission occurs:
- mainly from mosquitoes to vertebrate hosts
- via direct or indirect contact (by humans) with the blood or organs of infected animals, or from mosquito bites.

Clinical signs

In mature sheep, RVF may manifest as acute, subacute, chronic or convalescent disease. Disease is less severe in cattle but can cause abortion, and calf mortality may be high. Signs include:
- high mortality and abortion storms (in sheep)
- rapid onset pyrexia, anorexia and haemorrhagic diathesis leading to death (in neonatal lambs)
- pyrexia, vomiting, anorexia, haemorrhagic diarrhoea and mucopurulent nasal discharge (mature sheep with acute disease)
- infertility in sheep and cattle.

Post-mortem

Post-mortem poses a health risk for humans and should only be performed if good biosecurity can be achieved. The most characteristic post-mortem lesions involve the liver and aborted foetuses have swollen, yellow livers.

Samples

Collect:
- blood samples (EDTA and plain tubes)
- liver, lymph nodes, spleen, foetal membranes (viral titres are high in aborted foetuses and foetal membranes)
- fresh tissues.
**Actions to take**

If you suspect a case of RVF:

- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

**Public health considerations**

- RVF is a zoonotic disease
- people may become infected after close contact with blood, body fluids or tissues of RVFV-infected animals
- less commonly, people can be infected with RVFV from bites of infected mosquitoes
- consult the relevant state or territory health authorities as soon as RVF is strongly suspected or confirmed.
Introduction

Rift Valley fever (RVF) is an acute, arthropod-borne, zoonotic viral disease mainly affecting domestic ruminants. In ruminants the disease is characterised by abortion, high mortality in young animals and hepatopathy. People are also susceptible to the disease.

Disease agent and susceptible species

RVF:

- is caused by Rift Valley fever virus (RVFV), an enveloped RNA virus belonging to the family Bunyaviridae, genus Phlebovirus
- can cause severe disease in ruminants. Ruminants indigenous to regions in Africa where RVFV circulates endemically are relatively resistant to RVF, particularly Bos indicus cattle, but the susceptibility of Australian Bos indicus cattle is unknown
- causes significant disease in goats, and in sheep the virus is highly pathogenic
- affects camels and buffalo, which may become infected and develop mild disease
- causes influenza-like illness in humans, with severe complications in a minority of cases
- causes subclinical infection in cats and dogs (kittens and puppies are highly susceptible), horses, pigs and various wild species (for example, monkeys and rodents) but these species do not represent a major source of virus for vectors.

Public health considerations

- people may become infected after close contact with blood, body fluids, or tissues of RVFV-infected animals or from infected mosquitoes
- wear personal protective equipment such as gloves, disposable coveralls, rubber boots (or disposable boots) and masks when working with potentially infected animals to avoid any exposure to blood or tissues of RVFV-infected animals
- use insect repellents or netting to protect against mosquitoes and other bloodsucking insects
- consult the relevant public health authorities in the case of a RVF outbreak.

Distribution

RVF occurs throughout the African continent, with significant outbreaks in the last 30 years occurring in Egypt, southern and eastern African countries and Madagascar. RVF has been reported sporadically in Saudi Arabia and Yemen since 2000. A large outbreak occurred in Somalia and Kenya in 2006.

For the latest information on the distribution of RVF, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

None recorded.
Epidemiology

Large epidemics of RVF have occurred across a variety of ecological zones throughout Africa and the Arabian Peninsula at irregular intervals of 5–40 years, often in association with periods of above-average rainfall increasing mosquito vector breeding. Other risk factors include the presence of wetlands or waterlogged areas, large populations of arthropod vectors and susceptible livestock, large-scale irrigation projects and the transboundary movement of infectious livestock and their products.

Modes of transmission

Transmitting the disease via byproducts such as wool and hides is not well understood, and although semen collected from infectious animals is likely to contain the virus, the main mode of transmission is via vectors (from mosquito to animal). What is known is that:

- RVF is transmitted between animals by biting arthropods
- once RVFV is circulating and amplified in ruminants, it has the potential to infect a large number of arthropod vectors, including ticks and a variety of biting flies
- infected adult mosquitoes are likely to remain infectious for life and may transmit RVFV for the duration of their lifespan (up to 4 weeks)
- vertical (trans-ovarial) transmission occurs within Aedine spp. vectors, enabling agent survival and persistence between seasons between inter-epidemic periods.

To date, no human-to-human transmission of RVF has been documented. Human infections:

- mostly result from exposure to blood and fresh tissues (organs, carcasses, aborted materials) from infected animals, either via aerosol transmission or skin abrasions, or by biting mosquitoes (or, less commonly, flies)
- occur among farmers, herders, abattoir workers, veterinarians and laboratory personnel, who are at higher risk of infection through direct or indirect contact with the blood or organs of infected animals
- may result from drinking unpasteurised milk from infected animals (especially while animals are viraemic).

Disease dynamics

RVF:

- incubation period ranges from 1–30 days
- may be mild or sub-clinical in endemic areas
- epidemics cause abortion storms, devastating neonatal mortality and variably severe disease in mature animals with mortality rates up to 30 per cent
- virus is highly pathogenic in sheep and goats, but disease is relatively less severe in cattle
- is characterised (particularly following periods of above average rainfall) by the simultaneous occurrence of numerous cases of abortion and disease in ruminants, together with human illness.
Persistence of the agent

RVF:

- virus is stable in aerosol form and may persist in dried blood for up to 3 months (but survival times of up to 4 months and 21 days have been observed at 25°C and 27°C respectively)
- viral particles are susceptible to a range of disinfectants including chlorine-based disinfectant and detergents
- viral particles are destroyed by strong sunlight
- virus in milk is inactivated by pasteurisation
- viral titres in meat decline rapidly following slaughter and chilled meat is not likely to represent a hazard to human health.

Diagnosis and pathology

Clinical signs

RVF:

- disease in mature ruminants manifests as peracute, acute, subacute, chronic or convalescent forms
- disease is most severe in young sheep, goats and cattle
- commonly causes abortion in both sheep and cattle, and infertility may follow retention of foetal membranes and subsequent endometritis. Abortion may occur at any gestational age. Foetal death occurs rapidly in utero and the foetus may be aborted in an autolyzed state
- common haematology and clinical chemistry findings include leukopenia, thrombocytopenia and elevated liver enzymes.

Sheep are particularly susceptible to RVF and mortalities may exceed 90 per cent in neonatal lambs, 40–60 per cent in weaner lambs and up to 30 per cent in adults. In addition:

- adults commonly suffer subacute disease, with animals pyrexic for 24–96 hours
- lamb death may be sudden
- lambs may be viraemic within 8–12 hours following infection with RVFV, followed by a pyrexia at 24–36 hours preceding a rapidly progressive clinical course including anorexia, diarrhoea, jaundice and death within 24 hours.

Cattle mortality is usually less than 10 per cent (in adults) and may be accompanied by subacute diarrhoea and jaundice. Although RVF is less pathogenic in cattle, calf mortality may reach 70 per cent and is accompanied by pyrexia, anorexia and weakness.

Human infection with RVFV may result in mild or (less often) severe disease. In addition:

- disease is most commonly subclinical or mild, and associated with non-specific flu-like symptoms (including fever, muscle and joint pain and appetite loss lasting 4–7 days)
- a small proportion of people develop severe complications which fall into three syndromes: viral haemorrhagic fever, ocular disease or meningoencephalitis
- human mortalities are rare and tend to occur in patients who develop haemorrhagic fever.
Pathology

In aborted foetuses:
- foetal lesions are most consistently localised to the liver which is swollen and discoloured yellow to dark red
- gross lesions reflect viral tropism for hepatocytes.

In young animals, lesions are typically more acute and:
- the liver is swollen, friable and discoloured a yellow-tan, and generalised icterus may be present
- widespread petechiae and ecchymoses may be present on the serosal surfaces of abdominal viscera, including the liver, kidneys, gastrointestinal tract and lymph nodes.

In older animals:
- lesions may reflect an acute to subacute time course
- the liver is darker with scattered 1–2 mm areas of pallor throughout the liver indicative of necrosis.

Differential diagnosis

RVF should be highly suspected in ruminant disease outbreaks characterised by acute abortion storms and peracute neonatal death accompanied by hepatic necrosis and haemorrhagic diathesis. This is particularly the case when acute febrile illness is also reported in people handling sick animals.

Consider in ruminants presenting with the clinical signs and gross lesions:
- exotic diseases
  - bluetongue disease
  - ovine enzootic abortion (*Chlamyphila abortus*)
  - *Peste des petits ruminants*
  - Wesselsbron virus
  - *Brucella abortus* and *B. melitensis*
- endemic diseases
  - Akabane virus
  - any cause of septicemia
  - *Brucella suis*
  - *Campylobacter fetus*
  - enterotoxemia (*Clostridium perfringens* type C or D)
  - leptospirosis
  - *Listeria monocytogenes*
  - malignant catarrhal fever (only sheep-associated malignant catarrhal fever is endemic)
  - *Neospora caninum*
  - Q fever (*Coxiella burnetti*)
  - *Toxoplasma gondii*
• non-infectious causes
  – hepatotoxins (such as aflatoxin and phytotoxins)
  – copper toxicosis
  – anticoagulant toxicosis (for example from a pesticide such as Brodifacoum).

**Samples required**

**Sample collection**

Collect:

• **fresh tissue samples** of liver, spleen, lymph nodes and foetal membranes aseptically and place in sterile containers. Where possible, collect tissue for virus isolation from recently deceased animals or aborted foetuses

• **whole blood** from pyrexic animals using EDTA tubes

• **fixed tissue** samples for histopathology from the liver, spleen, lymph nodes, foetal membranes and brain (fixed in 10% neutral buffered formalin).

**Transport of samples**

For transport:

• chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs

• **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests

• send samples with dry ice if the journey is expected to take several days

• formalin fixed tissue can be sent at room temperature.

**Sample submission**

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

**Diagnostic tests**

When RVF is suspected, undertake diagnostic tests at a biocontainment laboratory where the risk of exposure to highly infectious material can be appropriately managed. Characteristic histopathology findings (such as acute hepatocellular necrosis with eosinophilic intranuclear inclusion bodies) may increase suspicion of RVF. However, definitive diagnosis requires identification of the virus through one or a combination of the following tests: PCR, virus isolation, immunofluorescence, ELISA or virus neutralisation.

**Reporting requirements**

RVF is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect RVF, report the disease immediately by phoning the **Emergency Animal Disease Watch Hotline on 1800 675 888**, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.
Biocontainment and personal protective equipment

RVF infection is a public health risk. Human infections may result from contact with aborted foetuses, bodily fluids and tissues of infected animals (as well as aerosols generated during slaughter or post-mortem examination). Post-mortem examinations of animals that have died from RVFV infection pose a risk for animal-to-human transmission. Therefore, isolate and keep suspected cases of RVF on the property (and protected from vectors, if possible) until advised by government veterinary authorities, noting:

- wear appropriate personal protective equipment (coveralls and boots, gloves, mask) when examining animals, performing post-mortem examinations, or handling tissue, carcasses or abortion materials
- isolate sick/dead animals and limit contact with the sick/dead animals
- personal decontamination is required when leaving infected premises. You must thoroughly clean and then disinfect vehicles and equipment before leaving the property.

Further information


FIGURE 3.21.1 An emphysematous bovine foetus with skin stained with meconium

Image credit: PIADC and CFSPH
FIGURE 3.21.2 The cut surface of a swollen ovine liver, which is pale and contains many petechiae

Image credit: PIADC and CFSPH

FIGURE 3.21.3 Severe haemorrhagic colitis in a sheep

Image credit: PIADC and CFSPH
CHAPTER 3.22

Scrapie

Fast facts
Scrapie is due to a prion that results in a fatal neurodegenerative disease.

Agent
- Scrapie is due to a structurally-altered, protease-resistant form of prion protein (known as PrPSc)
- atypical scrapie (Nor98) is caused by spontaneous folding of prion protein
- prions are resistant to physical and chemical inactivation procedures.

Susceptible species
- Sheep and goats, but with sheep, the host genotype influences susceptibility to both forms of scrapie.

Economic impact
There is potential for significant economic loss associated with restrictions on animal movements and international trade.

Epidemiology
- incubation period is protracted, typically 2–5 years (disease therefore manifests in adult animals)
- disease is always fatal.

Transmission
- horizontal transmission between animals likely occurs via ingestion
- vertical transmission may occur with classical scrapie in sheep
- natural transmission has not been observed with atypical forms of scrapie.

Clinical signs
Clinical signs include:
- changes in mentation, behaviour and posture or gait
- pruritis and self-trauma
- neurological disease accompanied by weight loss and emaciation.

Post-mortem
No gross lesions are detectable on post-mortem examination.

Samples
If safe to do so, and while wearing personal protective equipment, conduct a post mortem and collect:
- whole brain fixed in 10% neutral buffered formalin
- 1–2 cm segment of fresh cervical spinal cord and/or caudal medulla (caudal to the obex).

Actions to take
If you suspect a case of scrapie:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Scrapie is a progressive, fatal neurodegenerative disease of adult sheep, goats and mouflons. It may occur due to sporadic or heritable polymorphisms in the prion protein gene, or due to transmission of infectious prion between animals.

Disease agent and susceptible species

A structurally-altered, protease-resistant form (known as PrPSc) of the prion protein is the infectious agent transmitted between animals and is central to the pathogenesis of classical scrapie. Sheep and goats are susceptible and:

- susceptibility to disease varies between sheep breeds. Sheep with the ARR variant of the PrP gene are most resistant to disease whereas sheep homozygous for the variant VRQ/VRQ are most susceptible to scrapie
- in 1998, a variant form (‘atypical scrapie’ or ‘Nor98’) was found in a sheep in Norway
- the atypical form causes spontaneous disease in older animals and is distinguished from classical scrapie by distinct clinicopathological characteristics.

Distribution

Classical scrapie occurs in the United States, Canada, the United Kingdom, several European countries, Israel, India, Japan and Brazil. For the latest information on the distribution of scrapie, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

Following importation of affected animals from the United Kingdom, an isolated case of classical scrapie occurred on a single property in Victoria in 1952 and was successfully eradicated. Australia is free from classical scrapie.

Isolated cases of atypical scrapie have been detected globally, including in Australia.

Epidemiology

Modes of transmission

The precise mechanisms of scrapie transmission are not well understood, but what is known is that:

- animals are usually infected early in life, likely via oral exposure although vertical transmission may occur
- the prospect of maternal transmission is significant as affected ewes are likely to have several affected lambs before developing clinical signs
- scrapie spreads laterally among sheep grazing the same pasture and persists in affected flocks for prolonged periods
- foetal membranes, milk and contaminated materials, equipment, structures and vaccines are known sources of infection.
Disease dynamics

Disease presentation and progression is highly variable and may occur over a period of weeks to months. In addition:

- the agent initially proliferates in lymphoid tissue and the intestinal tract, and may take as long as 2 years to enter the CNS (potentially via splanchnic nerves)
- an additional 2 years may lapse before the onset of clinical signs
- due to the protracted incubation period, affected animals typically develop clinical disease at 2–5 years of age. With atypical scrapie, clinical disease usually appears in animals over 5 years of age and affects only single animals in a herd.

Persistence of the agent

Prion strains vary greatly in resistance to inactivation, and some are over a thousand times more resistant than others. In addition:

- PrP\textsuperscript{Sc} is resistant to physical and chemical inactivation procedures, including freezing, desiccation, ultraviolet radiation, degradation by various proteolytic enzymes, burial, and common methods of chemical and heat disinfection that destroy bacteria and viruses
- some organic acids can inactivate infectious prion proteins
- prions can adhere very strongly to stainless steel which may require special chemical disinfection.

Diagnosis and pathology

Clinical signs

The clinical course of scrapie is variable and protracted, and neurological signs may be slowly progressive over a period of weeks to months. In addition:

- the onset of clinical disease is typically characterised by mild changes in behavior and gait
- affected sheep present with paresthesia which manifests as pruritus, giving rise to the name ‘scrapie’
- animals may rub against fences or compulsively nibble at their legs and flanks resulting in self trauma and wool damage
- neurological signs are accompanied by weight loss, and eventually emaciation and death.

Pathology

No gross abnormalities are present in the brain.

Differential diagnosis

Differential diagnoses include disorders that affect the neurological function and locomotion, which includes:

- exotic diseases
  - louping ill
  - maedi-visna
  - rabies
Emergency animal diseases

• endemic diseases
  – botulism
  – brain or spinal cord abscess
  – *Clostridium perfringens* type D (focal symmetrical encephalomalacia)
  – ectoparasites
  – *Listeria monocytogenes*

• non-infectious causes
  – brain or spinal cord trauma
  – lead or plant toxicosis
  – metabolic disease (hypomagnesemia, hypocalcemia, polioencephalomalacia)
  – neoplasia
  – nutritional myopathy (white muscle disease).

**Samples required**

**Sample collection**

There is currently no validated diagnostic test available in Australia for detecting scrapie in live animals. Diagnostic tests require brain and spinal cord obtained at post-mortem examination. Although scrapie is not considered to be a zoonosis, note that you should:

• take precautions during sampling, including wearing full personal protective equipment
• take care when disposing or decontaminating equipment used for the postmortem removal of brain tissue from suspected scrapie cases.

In addition:

• where scrapie is strongly suspected, direct efforts at collecting the brain and cervical spinal cord samples with minimal damage
• it may be possible to recover diagnostic material following mechanical injury to the brain (e.g. euthanasia by captive bolt)
• for suspected cases of scrapie, collect whole brain fixed in 10% neutral buffered formalin, and a 1–2 cm segment of unfixed (fresh) cervical spinal cord and/or caudal medulla (caudal to the obex).

**Transport of samples**

For transport:

• chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days
• formalin fixed tissue can be sent at room temperature.

**Sample submission**

Sample packaging and consignment for delivery to CSIRO-AAHL should be coordinated by the relevant state or territory laboratory.
Diagnostic tests

Histologic examination is required to identify characteristic lesions within relevant areas of the CNS (which differ from atypical scrapie), and to exclude differential diagnoses. In addition:

- immunohistochemistry (IHC) detects PrPSc aggregates in formalin-fixed brain sections and can be evaluated in conjunction with histological changes
- detection by western blotting of PrPSc in unfixed tissue homogenates has similar diagnostic sensitivity to IHC
- rapid ELISAs on unfixed tissue are typically used as a screening strategy in surveillance programs.

Reporting requirements

Scrapie is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect scrapie, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no known public health implications for scrapie, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment.

During a post-mortem examination:

- minimise environmental contamination during post-mortem examination
- protect eyes and mucous membranes with safety spectacles and masks or a full face visor
- use a protective respirator
- minimise the use of sharps and where possible use single-use disposable items (e.g. scalpels)
- use suitable hand protection such as armoured or cut resistant gloves, especially during collection of the brain or spinal cord
- contaminated bone dust will be minimised if hacksaws or bone forceps are used (rather than reciprocating or oscillating saws) for removal of the brain or spinal cord.

Further information


CHAPTER 3.23

Sheep pox and goat pox

Fast facts

Sheep pox and goat pox are clinically indistinguishable, contagious, generalised exotic viral diseases of small ruminants.

Organism

Sheep pox virus (SPV) and goat pox virus (GPV):
- strains may vary considerably in host specificity and virulence
- are very stable viruses in the environment.

Susceptible species

SPV and GPV affect sheep and goats and strains are mostly host-specific, but some have equal virulence in both species.

Economic impact

There is potential for significant economic loss associated with high morbidity and mortality in small ruminants, as well as restrictions on animal movements and international trade.

Epidemiology

SPV and GPV:
- peak mortality usually occurs about 2 weeks after the development of lesions
- may cause morbidity approaching 100 per cent and mortality reaching 50 per cent in adults
- may cause morbidity and mortality approaching 100 per cent in young animals
- usually has an incubation period of 12 days but this may range from 2–14 days.

Transmission

Transmission occurs:
- mainly by direct contact via respiratory aerosols
- via skin abrasions
- via all secretions and excretions
- via insects (over short distances)
- via fomites.

Clinical signs

Signs include:
- sudden onset of pyrexia, nasal and ocular discharges, hypersalivation and lymphadenopathy
- laboured breathing (secondary pneumonia is common)
- classical pox lesions all over the skin
- lesions on the mucous membranes of the mouth, nostrils and vulva.

Post-mortem

You may see:
- lung lesions consisting of grey-white or red nodules
- swollen, congested lymph nodes with petechial haemorrhages
- skin lesions and haemorrhagic ulcerations in the linings of the trachea and gastrointestinal tract.

Samples

Collect:
- fresh tissue samples—skin lesion biopsies, scabs, vesicular fluid (if present) and from lesions in the respiratory and gastrointestinal tracts
- skin scrapings
- whole blood in EDTA.

Actions to take

If you suspect a case of sheep or goat pox:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction
Sheep and goat pox are contagious, generalised viral diseases of small ruminants. Clinically indistinguishable, the two diseases are the most serious poxvirus diseases of livestock and may cause considerable production losses and mortalities.

Disease agent and susceptible species
Sheep and goat pox are caused by sheep pox virus (SPV) and goat pox virus (GPV). SPV and GPV:
- are enveloped, double stranded DNA viruses in the family Poxviridae, subfamily Chordopoxvirinae, genus Capripoxvirus
- have strains which may vary considerably in host specificity and virulence
- share the same serotype and are morphologically and serologically indistinguishable from each other
- are closely related to lumpy skin disease virus (which affects cattle)
- can recombine and produce a spectrum of strains with intermediate host specificity and virulence across species
- usually demonstrate host specificity and cause clinical disease in either sheep or goats; however, some strains have equal virulence in both species
- infect sheep (merino and European breeds of sheep are more susceptible to SPV than other breeds)
- infect goats, although breeds vary in susceptibility to GPV and breeds exotic to the source area are more severely affected
- have no known wildlife reservoir
- do not infect humans.

Distribution
Globally, SPV and GPV:
- are endemic in Africa (north of the equator), the Middle East, Central Asia, including southern Russia and western China, and the Indian subcontinent
- spread into south-eastern Europe causing outbreaks in Turkey, Greece and Bulgaria in 2013–2015.

For the latest information on the distribution of sheep and goat pox, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia
SPV and GPV have never been reported in Australia.
Epidemiology

Modes of transmission

Transmission occurs via:

- **secretions and excretions**—sheep and goats with acute disease shed virus in all secretions and excretions. Infection is mainly via respiratory aerosols but may also occur through cuts and abrasions in the skin or mucosal membranes.

- **fomites**—the viruses are readily transported on fomites, including clothing and equipment. Due to the stability of the viruses in the environment, indirect contact by means of contaminated yards, vehicles and other fomites (such as hypodermic needles) is an important means of spread of the diseases.

- **vectors**—less commonly, insects (such as biting flies) can act as mechanical vectors of the viruses over short distances.

- **contact with the carcasses** of infected sheep and goats.

- **semen and embryos**—while no information is available on the transmission of SPV or GPV in semen and embryos, you should assume that semen and embryos would contain the viruses during the viraemic period. The closely related lumpy skin disease virus of cattle is shed in the semen of clinically affected bulls for up to 22 days, and for at least 12 days in subclinically affected bulls.

Disease dynamics

Sheep and goat pox:

- viruses are present in all secretions and excretions (including milk and scabs from skin lesions) of infected animals at every stage of the disease, but scabs are not considered to be important sources of transmission.

- viruses may cause disease ranging from inapparent or mild to severe, depending on age, breed, immunity, virus strain and other factors.

- have a peracute form (usually seen in initial outbreaks in an area) characterised by pyrexia, death and (in some European goat breeds) flat haemorrhagic papules that coalesce to form widespread cutaneous ulceration.

- viral incubation usually lasts 12 days but may vary from 4–14 days.

- would likely cause severe disease in Australian sheep and goat populations (which are fully susceptible).

- lesions follow the classical pox cycle (over about 2 weeks) of macule (skin erythema), papule (0.5–1.5 cm diameter), vesicle (rare), pustule with exudation, encrustation and scab formation (exudate from ruptured pustules can cause the fleece to matt).

- skin lesions heal slowly, taking 5–6 weeks with secondary skin infections and fly strike being common complications.

- may cause death at any stage of the disease, with peak mortality occurring about 2 weeks after lesions appear.

- have no carrier state and recovered animals develop lifelong immunity.
In addition:

- in experimentally infected sheep and goats, a cell-associated viremia occurs from days 4–6 post-inoculation, and clears within 2–3 weeks
- shedding of virus in nasal, conjunctival and oral secretions occurs from as early as day 6–8 post-inoculation, peaks between days 10–14, and may persist at low levels for up to an additional 3–6 weeks
- infectious virus has also been detected between days 8–15 post-inoculation (at low levels) in the faeces of experimentally infected sheep and goats, and the urine of sheep
- movement of infected sheep and goats is the most important means of spreading the diseases to new areas
- feral goats could potentially become an important reservoir of the viruses in Australia.

**Persistence of the agent**

Capripoxviruses are very stable in the environment and can remain viable for long periods, on or off the animal host. SPV and GPV:

- can persist for up to 3 months in wool and hair from infected animals, for up to 6 months in a cool, dark environment (such as in shaded animal pens) and for many years in dried scabs at ambient temperatures
- are susceptible to lipid solvents and acids. Therefore, acids combined with detergents (such as Virkon®) are the disinfectants of choice, particularly for areas where organic matter is prevalent. Hypochlorites and aldehydes are useful for disinfecting clean surfaces. Citric acid, alcohols and iodophors are suitable for personal disinfection
- are inactivated after heating for 1 hour at 55°C.

**Diagnosis and pathology**

**Clinical signs**

For both sheep and goat pox:

- disease may vary from inapparent or mild to severe, with severe disease typically seen in fully susceptible flocks
- mild disease (typically seen in partially immune animals or in resistant breeds in endemic regions) is often missed clinically because skin lesions are mild, few in number and often restricted to certain areas such as the ears and around the tail.

**Peracute disease**

Peracute cases may die before lesions become very obvious.

**Acute disease**

Clinical signs for both sheep and goat pox include:

- anorexia
- conjunctivitis
- lesions within the mouth, nostrils and vulva
- reluctance to move, standing with arched backs
- a sudden onset of pyrexia, which peaks at 40–42°C, with ocular and nasal discharge and hypersalivation
lymphadenopathy which may be particularly prominent in prescapular nodes
- typical pox lesions in the gastrointestinal tract, trachea and lungs
- acute respiratory distress and commonly secondary pneumonia
- mortality, which may reach 50 per cent in adults and approach 100 per cent in young animals
- flat haemorrhagic papules which in some European goat breeds which coalesce to form widespread cutaneous ulceration
- death, which may occur at any stage of the disease, with peak mortality occurring about 2 weeks after lesions appear
- formation of erythematous macules in the skin within 1–2 days of onset of pyrexia. Macules progress to papules within 1–2 days and are most readily detected by palpation, particularly in animals with hairy fleeces. They may extend all over the skin, but are most obvious where wool or hair is shortest, such as on the face, ears, axillae, groin, perineum and under the tail. Encrustation of papules and scab formation occurs within the following 1–2 weeks.

**Nodular disease**

A nodular form of sheep and goat pox—known as stonepox—is also described. Stonepox resembles lumpy skin disease (of cattle), with skin lesions of 0.5–3 cm in diameter hyperaemic, thickened and raised above the surrounding skin.

**Pathology**

In addition to the lesions described above, internal lesions are usually present in the acute disease. Findings include:
- ulcerations or nodules in the mucous membranes of the oral cavity, trachea and the gastrointestinal tract, including the rumen, abomasum and reticulum
- pulmonary lesions in about 40 per cent of cases. These consist of grey-white or red nodules that are generally sub-pleural but may be readily palpated as hard nodules throughout the lungs
- nodules in the omentum, kidneys and liver
- enlarged and haemorrhagic lymph nodes.

**Differential diagnosis**

Acute cases in naïve populations should be readily diagnosed based on clinical signs. However, some strains of low virulence may produce only mild clinical signs. Consider in the differential diagnosis:
- exotic diseases
  - [bluetongue disease](#) (infection present in Australia but without clinical disease)
- endemic diseases
  - contagious pustular dermatitis (scabby mouth or orf)
  - ectoparasites
  - mycotic dermatitis
- non-infectious causes
  - insect bites
  - photosensitisation.
Samples required

Sample collection
You will be able to isolate the virus within the first week of clinical signs developing, before neutralising antibodies develop. Collect:

- **serum** from at least 10 live, clinically affected animals, and from exposed animals (particularly those that are convalescent)
- **EDTA blood** from live, clinically affected animals (7–10 ml/animal)
- **fresh tissue**—characteristic pox lesions from skin as well as respiratory and gastrointestinal tracts (2 g of each tissue)
- **nasal swabs** from live, clinically affected animals
- **fixed tissue**—characteristic pox lesions from skin as well as respiratory and gastrointestinal tracts (in neutral-buffered formalin).

Transport of samples
For transport:

- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- place swab samples in virus transport media containing antibiotics and antifungals (or saline if other media is unavailable)
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
For both sheep and goat pox:

- a rapid, tentative laboratory diagnosis can be made by electron microscopy of tissue samples. Electron microscopic examination of tissue samples reveals typical capripoxvirus particles, which are readily differentiated morphologically from those of contagious pustular dermatitis (orf) virus
- confirmation of the diagnosis is obtained by specifically identifying the virus in tissues from early lesions or in cell culture using virus-specific tests
- the most rapid, sensitive and specific diagnostic procedure is the detection of viral DNA in characteristic poxvirus lesions or nasal swabs by real-time or conventional PCR. A positive result can be obtained within 1 day of the laboratory receiving samples.
Reporting requirements

Sheep and goat pox are OIE-listed diseases and Australia has an international obligation to report cases. If sheep or goat pox are suspected, report the diseases immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for sheep and goat pox, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


FIGURE 3.23.1 Multiple red-brown papules on the scrotum and inguinal skin of a sheep

Image credit: PIADC and CFSPH

FIGURE 3.23.2 A caprine muzzle with several papules and partially covered by a haemorrhagic nasal exudate

Image credit: PIADC and CFSPH
FIGURE 3.23.3 Several coalescing cutaneous pox with pale tan (necrotic) centres on ovine skin

Image credit: PIADC and CFSPH

FIGURE 3.23.4 Raised pale nodules in the lung of a small ruminant

Image credit: PIADC and CFSPH
FIGURE 3.23.5 Endometrium with several tan papules (pox) amongst the caruncles in an ovine uterus

Image credit: PIADC and CFSPH
CHAPTER 3.24
Swine vesicular disease

Fast facts
Swine vesicular disease (SVD) is a highly contagious, exotic viral disease of pigs clinically indistinguishable from foot-and-mouth disease (FMD).

Organism
SVD virus (SVDV):
- primarily replicates in the endothelial cells of the gastrointestinal tract
- can survive in the environment for long periods and is highly resistant to heat and a wide pH range. The most effective disinfectants are strong alkalis.

Susceptible species
Only pigs are susceptible.

Economic impact
There is potential for significant economic loss resulting from difficulty of eradication, as well as restrictions on animal movements and international trade.

Epidemiology
SVD:
- causes high morbidity which can reach 100 per cent but very low (zero) mortality
- has an incubation period of 2–7 days; longer if dose is small
- infection can be subclinical.

Transmission
Swill feeding has been responsible for most primary outbreaks. In addition:
- virus is shed in nasal secretions and excretions of infected pigs
- pigs can excrete the virus in faeces up to 48 hours before clinical signs are seen, and for up to 3 months following infection.

Clinical signs
SVD can be a subclinical, mild or severe vesicular condition depending on the virus strain, route and dose of infection. Housing on damp concrete may exacerbate clinical signs, with younger animals more severely affected. The main clinical signs are:
- vesicles and ulcers on the snout, mammary glands, coronary band, interdigital areas and on the skin of the lower legs, particularly at pressure points such as the knees
- pyrexia
- lameness
- salivation.

Post-mortem
Post-mortem lesions of SVD are restricted to vesicle formation and resolution.

Samples required
Also refer to the detailed section on sampling for animals with vesicular disease.
- blood (EDTA and plain tubes)
- vesicular fluid, oropharyngeal fluid, oral, nasal and tonsillar swabs, epithelium, and epithelial tags (in buffer or virus transport medium).

Actions to take
If you suspect a case of SVD:
- call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
- isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Swine vesicular disease (SVD) is an acute, highly contagious but rarely fatal, exotic vesicular viral disease of pigs. SVD is of international concern because it is clinically indistinguishable from foot-and-mouth disease (FMD). Rapid differentiation from FMD is essential.

Disease agent and susceptible species

Pigs are the only species susceptible to SVD virus (SVDV). SVDV:
- is a member of the Enterovirus genus of the family Picornaviridae
- has two serotypes (SVDV-1 and SVDV-2) within the species Enterovirus B
- is a non-enveloped virus with a single-stranded RNA genome (non-enveloped viruses are in general more stable than enveloped viruses)
- strains vary in virulence, with subclinical infection the most frequent condition observed during recent years.

Distribution

Since the 1970s, the disease appears to have been eradicated from most countries. However, as the disease is no longer OIE-listed, countries are not obligated to report cases, which makes it difficult to determine distribution. SVD is endemic in southern Italy and possibly in parts of Asia.

Occurrences in Australia

None reported.

Epidemiology

Modes of transmission

Transmission occurs via:
- direct contact with infected pigs, as the virus is shed in secretions and excretions (including faeces) of infected pigs. Movement of infected pigs is the most important means of spread
- contact with faeces is an important means of spread as the virus can be shed in the faeces for up to 3 months following infection
- ingestion of pig meat or pig meat products infected with the virus is an important means of spread. In Australia, strict import conditions are in place to address the risk of a SVD incursion and swill feeding is illegal
- fomites, because of the stability of SVDV in the environment, contaminated pig pens, water, vehicles and other fomites (such as hypodermic needles) can spread disease
- contact with the carcasses of pigs affected by SVD
- mechanical spread by people, rodents, insects and birds (but this plays a minor role in transmission).
Disease dynamics

SVDV is considered to be highly contagious, but mortality is low. Dynamics are as follows:

- virus enters host pigs through damaged epithelia and multiplies in epithelial cells
- pigs can secrete the virus from the nose or mouth, and excrete the virus in faeces up to 48 hours before clinical signs are seen
- the incubation period in natural outbreaks is 2–7 days. The incubation period for SVD varies by route of transmission, with 2–5 days incubation more common if the virus is ingested in contaminated meat, and 2–7 days more common if acquired through contact with infected animals or faecal material
- subclinical infection may occur if pigs are exposed to small amounts of virus, particularly by inhalation or ingestion
- no persistent infections have been reported, and infected pigs develop protective antibody against SVDV upon recovery
- there is no evidence of a carrier state and no reservoir hosts are known
- on some properties, the disease apparently runs its course and, once all animals are infected, dies out. On other properties, two waves of disease approximately 3 months apart have been observed.

Persistence of the agent

SVDV is highly stable in the environment, which complicates eradication and makes prompt recognition essential for control. SVDV:

- can persist for many months in a range of pork products and frozen carcasses but has not been detected in cooked, canned ham
- is resistant to heat up to 69°C, pH ranging from 2–12 and many common detergents and disinfectants
- is protected by manure, fats and other organic matter, which must be completely removed during cleaning
- can survive in pig faeces for at least 138 days
- is most effectively inactivated with strong alkalis (for example, sodium hydroxide), although hypochlorites or acid-containing iodophors can be used when organic material is not present.

Diagnosis and pathology

Suspect SVD and other vesicular diseases when vesicles or erosions are found on the mouth and/or feet of pigs. Other vesicular diseases must be ruled out with laboratory tests.
Clinical signs
SVD can be a subclinical, mild or severe vesicular condition depending on the strain of virus involved, the route and dose of infection, and the husbandry conditions under which the pigs are kept. Clinical signs can be more severe in animals housed on damp concrete, with younger animals more severely affected. Clinical signs are often mild and overlooked (particularly in muddy yards or where the animals are infrequently observed), and affected pigs recover rapidly (usually occur within 2–3 weeks) with little permanent damage. A dark, horizontal line on the hoof may appear where growth has been temporarily interrupted. Clinical signs are:

- very similar to FMD (including pyrexia, salivation and lameness) but with lower morbidity and less severe lesions
- a transient pyrexia (following incubation) of up to 41°C, after which vesicles develop on the coronary band, typically at the junction with the heel (these may affect the whole coronary band resulting in loss of the hoof)
- vesicles and ulcers on the snout, mammary glands, coronary band, interdigital areas and on the skin of the lower legs, particularly at pressure points such as the knees, but vesicles in the oral cavity are rarely seen
- anorexia
- shivering, unsteady gait, and chorea (rhythmic jerking) of the legs.

Pathology
Post-mortem lesions of SVD are restricted to vesicle formation and resolution.

Differential diagnosis
SVD is clinically undistinguishable from the other exotic viral vesicular diseases, namely:

- foot-and-mouth disease
- Senecavirus A (Seneca Valley virus) infection.
- vesicular exanthema of swine (not seen globally since 1956)
- vesicular stomatitis.

A number of other diseases cause similar clinical signs to the viral vesicular diseases, particularly during their more advanced clinical stages. Although mouth and muzzle lesions in these diseases are not vesicular, they could be confused with FMD lesions. These include:

- non-infectious causes
  - chemical or thermal burns
  - phototoxic dermatitis with vesicle formations from contact with the leaves of plants of the family Umbelliferae (parsley, parsnips and celery)
  - chemical irritants and scalding.

Differential diagnoses of lameness include bad/hard floors and Glässers disease (*Haemophilus parasuis*).
Samples required

Take samples from at least five affected pigs in the group and from lesions in the mouth or the feet, or at other sites with suitable lesions. Because SVD may circulate unnoticed, you will need to look for seroconversion to SVDV in apparently healthy animals to ascertain how long infection has been present.

Sample collection

FMD virus (an important differential diagnosis for SVD) is extremely sensitive to both acid and alkaline conditions. Inappropriate buffer conditions can inactivate the virus making virus isolation difficult or impossible. To maximise chances of virus isolation:

- use phosphate buffered saline or virus transport media with a pH of 7.6
- if a sample is to be submitted after 24 hours or more, add glycerol to the phosphate buffered saline
- dilute oropharyngeal fluid collected with a probang in an equal volume of phosphate buffered saline pH 7.6, and mix vigorously for 1 minute.

Note that samples collection in these buffers is optimal (for the growth of FMD virus), but not essential for RNA or antigen detection assays.

Take samples from at least 10 live, clinically affected animals. Collect:

- serum, 7–10 ml/animal in plain tubes
- vesicular fluid—carefully use a syringe and needle to aspirate the vesicular fluid from unruptured vesicles, and place in a sterile container. Alternatively, collect fluid from small vesicles onto a swab and place the swab in 500 µl of buffer, such as phosphate buffered saline or virus transport medium
- fresh tissue, epithelium, epithelial tags, oral, nasal and tonsillar swabs and oropharyngeal fluid and submit in phosphate buffered saline or virus transport medium, if available.

Transport of samples

For transport:

- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests

SVDV is identified using an antigen enzyme-linked immunosorbent assay (ELISA), the direct complement fixation test, virus isolation in pig-derived cell cultures and PCR. Virus neutralisation is the definitive serological test. Serological ELISAs are used for surveillance.
Reporting requirements

SVD is a nationally notifiable disease and there is an obligation to report cases. If you suspect SVD, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

There are no public health implications for SVD, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


FIGURE 3.24.1 A deep ulcer on the dorsum of the snout

Image credit: Iowa State University, College of Veterinary Medicine and CFSPH

FIGURE 3.24.2 Ulcers on coronary bands

Image credit: PIADC and CFSPH
CHAPTER 3.25
Transmissible gastroenteritis

Fast facts
Transmissible gastroenteritis (TGE) is a highly contagious exotic disease of pigs characterised by diarrhoea and transient vomiting with associated high mortality in piglets under 5 weeks of age.

Organism
TGE virus (TGEV):
• replicates primarily in the small intestine causing cell lysis and damage to villus epithelium
• is rapidly inactivated at temperatures above 37°C and considered highly sensitive to sunlight
• is inactivated by the use of formalin (0.03%), sodium hypochlorite, NaOH, iodine, ether and quaternary ammonium compounds.

Susceptible species
TGE primarily affects pigs but cats, dogs and foxes are all are considered carriers.

Economic impact
There is potential for significant economic loss associated with high mortality in young pigs in a naïve herd, as well as restrictions on animal movements and international trade.

Epidemiology
TGE:
• has a higher transmission in herds during winter months
• may cause mortality of 100 per cent in piglets less than 2–3 weeks of age
• incubation lasts 18 hours to 3 days.

Transmission
Transmission occurs:
• typically via faecal-oral route
• via nasal secretions and milk during acute infections
• via fomites (during colder months).

Clinical signs
Signs include:
• watery diarrhoea
• vomiting
• rapid weight loss/wasting
• anorexia
• dehydration.

Post-mortem
This may be found:
• small intestine distention with foamy, yellow fluid and undigested/curdled milk
• +/- gastric petechial haemorrhage.

Samples
Refer to the detailed section on sampling for pigs with diarrhoea.
• whole blood in EDTA and plain tubes
• faeces in sterile tubes
• full range of tissue including lungs, intestine, spleen, liver and kidneys fresh and in 10% neutral-buffered formalin.

Actions to take
If you suspect a case of TGE:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.
Introduction

Transmissible gastroenteritis (TGE) is an acute, highly contagious exotic disease of pigs with a high mortality case rate in piglets aged under 5 weeks.

Disease agent and susceptible species

TGE:
- is caused by the TGE virus (TGEV), an enteropathogenic coronavirus belonging to the Alphacoronavirus genus within the family Coronaviridae
- is related to other porcine coronaviruses, including porcine epidemic diarrhoea virus (PEDV), porcine deltacoronavirus (PDCoV), porcine respiratory coronavirus (PRCV) and porcine hemaglutinating encephalomyelitis virus (HEV)
- virus has only one serotype; however, the virus is closely related to PRCV which will cross-react serologically
- causes clinical disease in pigs (Sus scrofa)
- can be carried by cats, dogs, foxes and possibly mink (which do not display clinical signs).

Distribution

TGEV is present in most parts of Europe, the Americas (North, South and Central), South-East Asia, China, Japan and South Korea and West Africa. For the latest information on the distribution of TGE, refer to the WAHIS information database website of the World Organisation for Animal Health (OIE) [http://www.oie.int].

Occurrences in Australia

TGE has not been reported in Australia.

Epidemiology

Modes of transmission

While the exact mechanism is unclear, faeco-oral contamination is considered the most likely mode of transmission. Transmission via fomites is possible but the virus is rapidly deactivated in warm environmental conditions (in sunlight and at room temperature or above). Otherwise, transmission occurs via:
- direct contact with the faeces or nasal secretions of infected pigs
- ingestion of litter, feed or water contaminated by the faeces or nasal secretions of infected pigs
- virus shed in the milk of sows during the acute stage of the disease (piglets may become infected from milk in this way)
- contact with reservoirs/carriers, which includes wild carnivores such as dogs, cats and foxes
- swill feeding of contaminated swill
- mechanical vectors in the form of wild birds and flies.
**Disease dynamics**

TGEV infection is associated with high case morbidity and high case mortality (up to 100 per cent) in piglets less than 2–3 weeks of age, and:

- the incubation period is extremely short, estimated at between 18 hours and 3 days
- infection usually spreads very rapidly through a susceptible population (but spread may be slower in warmer environments)
- faecal shedding may persist for up to 2 weeks after recovery from infection
- pigs that recover from enteric infections develop immunity, but the longer-term carrier status of recovered animals is difficult to assess.

**Persistence of the agent**

Freezing and post-slaughter tissue acidification to not reduce the infectivity of TGE virus in pig products. TEGV:

- can persist in the environment for up to 72 hours (but is rapidly inactivated at temperatures above 37°C)
- can survive for up to 2 weeks in manure slurry at 20°C
- is susceptible to sunlight and most chemical disinfectants
- can be inactivated with 1% phenol, 0.01% betapropiolactone, sodium hypochlorite, 1 mM binary ethylenamine, sodium hydroxide, iodines, quaternary ammonium compounds, ether and chloroform.

**Diagnosis and pathology**

**Clinical signs**

The incubation period varies from between 18 hours and 3 days. Typical clinical signs in young pigs include:

- watery foul smelling yellow diarrhoea
- transient vomiting
- rapid weight loss
- anorexia
- depression
- rapid dehydration
- death within 2–7 days
- high mortality of pigs under 2–3 weeks of age (can be as high as 100 per cent).

The disease in grower and breeding herds can be clinically indistinguishable from porcine epidemic diarrhoea (PED), with typically watery diarrhoea, inappetence, ill-thrift and agalactia in lactating sows.

Laboratory testing is required to exclude other causes of diarrhoea in pigs.
Pathology

Typically, gross pathological lesions indicative of enteritis are found (usually confined to the gastrointestinal tract). Other gross lesions found may include:

- stomach distention with undigested/curdled milk
- +/- gastric petechial haemorrhage
- stomach congestion with regional haemorrhage on diaphragmatic surface (in 50 per cent of cases)
- small intestine distention with foam yellow fluid and milk
- thinning of intestinal wall (which may appear almost transparent due to villus atrophy).

Differential diagnosis

Differential diagnoses for TGE are those which involve atrophic enteritis, which includes:

- exotic diseases
  - porcine epidemic diarrhoea (PED)
  - porcine deltacoronavirus
  - Senecavirus A (Seneca Valley virus) infection (this would be a differential for neonates <2–3 weeks)
- endemic diseases
  - colibacillosis
  - enterotoxemia (Clostridium perfringens type C)
  - haemagglutinating encephalomyelitis virus (HEV)
  - porcine circovirus type 2-associated diarrhoea
  - porcine rotavirus infection
  - proliferative enteritis
  - salmonellosis
  - swine dysentery
- non-infectious causes
  - arsenic poisoning.

Samples required

Viral antigen is best detected in piglets sacrificed at a very early stage of disease. Collect enough samples (from at least five animals) to investigate suspected endemic causes of gastroenteritis, as you can frequently observe clinical signs of TGEV in other enteric infections.

Sample collection

Collect:

- **serum** from acute and convalescent animals (at least 10 animals)
- **fresh tissue** in the form of loops of affected ileum from acutely ill cases and preferably collected within 24 hours of the onset of clinical signs (tie off ileum and store in sterile containers on ice, and if transit is likely to take more than 24 hours, add glycerol buffer—pH 7.4—to the samples)
- **fixed tissue** in the form of sections of small intestine (both unpreserved and in neutral buffered formalin) from different parts of the small intestine.
Transport of samples
For transport:
• chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
• DO NOT FREEZE SAMPLES at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
• send samples with dry ice if the journey is expected to take several days
• formalin fixed tissue can be sent at room temperature.

Sample submission
The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.

Diagnostic tests
Laboratory testing is required for the diagnosis of TGE and is accomplished by virus neutralisation and isolation, electron microscopy and immuno-electron microscopy, and immunofluorescence. A PCR test is also available and can be used to differentiate TGE virus from other porcine coronaviruses.

Reporting requirements
TGE is an OIE-listed disease and Australia has an international obligation to report cases. If you suspect TGE, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment
There are no public health implications for TGE, but you should implement biocontainment protocols until advised by government veterinary authorities. This includes isolating suspected cases and using and appropriately disposing of personal protective equipment such as gloves, coveralls and rubber boots (or disposable boots). Thoroughly disinfect and decontaminate clothing, vehicle and equipment before leaving the property.

Further information


CHAPTER 3.26

Vesicular stomatitis

Fast facts

Vesicular stomatitis (VS) is an exotic zoonotic, vesicular viral disease of livestock, clinically indistinguishable from other important livestock vesicular diseases, such as foot-and-mouth disease (FMD).

Organism

VS is caused by a number of vesiculoviruses:
• viral antigens localise in keratinocytes in the coronary band
• virus can retain infectivity in the environment for several days but is sensitive to a wide range of disinfectants (e.g. 1% chlorine bleach, quaternary ammonium).

Susceptible species

Susceptible species include:
• cattle, horses and pigs
• South American camelids, sheep and goats occasionally have clinical signs
• people.

Economic impact

There is potential for significant economic loss resulting from restrictions on animal movements and international trade.

Epidemiology

VS:
• incubation period is usually 2–8 days
• causes morbidity ranging from 5–90 per cent but many infections in livestock are subclinical
• mortality is rare in cattle and horses but higher mortality rates have been seen in some pigs.

Transmission

Transmission of VS is incompletely understood. Insect vectors, mechanical transmission and movement of animals may transmit infection.

Clinical signs

Signs include:
• pyrexia
• hypersalivation (often the first sign)
• vesicles that progress to erosions in the mouth, nares, muzzle, teats and feet.

Post-mortem

Gross lesions at post-mortem are restricted to erosive ulcerative lesions.

Samples required

Also refer to the detailed section on sampling for animals with vesicular disease; collect:
• blood (EDTA and plain tubes)
• vesicular fluid, oropharyngeal fluid, oral, nasal and tonsillar swabs, epithelium, and epithelial tags (in buffer or virus transport medium).

Actions to take

If you suspect a case of VS:
• call the Emergency Animal Disease Watch Hotline (1800 675 888) immediately or contact a government veterinarian in your state or territory. Isolate animals until a definitive diagnosis is determined
• isolate suspected cases and implement biocontainment protocols (including farm movement controls) until advised by government veterinary authorities.

Public health considerations

• VS is a zoonotic disease
• humans can become infected with VSV when having direct contact with infected animals
• VS causes flu-like symptoms and rarely causes oral blisters in people.
Introduction

Vesicular stomatitis (VS) is an important exotic vesicular viral disease which produces significant economic losses in livestock, and poses a zoonotic risk to people. As with all vesicular diseases, rapid differentiation from FMD is essential.

Disease agent and susceptible species

VS is caused by a number of viruses belonging to the *Vesiculovirus* genus in the family *Rhabdoviridae* (vesicular stomatitis New Jersey (VSNJV) and vesicular stomatitis Indiana subtypes 1, 2 and 3). In addition:

- VS primarily affects equids (horses, donkeys, mules) and cattle and, occasionally, pigs, sheep, goats, llamas and alpacas
- most cases occur in adults and young cattle; horses under a year of age are rarely affected
- humans can also become infected, producing influenza-like symptoms
- South American camelids, sheep and goats occasionally display clinical signs
- serological evidence of infection is found in many other animals including deer, bats, raccoons, skunk, monkeys, sloths, rodents and wild pigs, but there is a noticeable absence of a viremic mammalian host, which is an unusual feature for an arbovirus.

Public health considerations

- VS is a zoonosis
- use appropriate personal protective equipment as humans (in particular veterinarians and animal handlers) become infected after direct contact with infected animals
- symptoms include flu-like signs and oral blisters (rarely).

Distribution

VS:

- is limited to the Americas
- is known to be an endemic disease in the warmer regions of North, Central and South America, but outbreaks of the disease in other temperate geographic parts of the Americas occur sporadically
- outbreaks have occurred in a number of states in the United States throughout the 21st century.

Occurrences in Australia

None reported.
Epidemiology

Modes of transmission

The mechanism of transmission of VSV is unclear, but it is thought to involve:

- **direct contact** between animals, as infected animals shed VSV in vesicular material, saliva and to a lesser extent, in nasal secretions and broken skin or mucous membranes may facilitate entry of the virus. Faecal shedding has been reported occasionally in pigs. VSV does not appear to be shed in milk.

- **insect vectors**, which introduce VSV into populations of domesticated animals. Vesiculoviruses are transmitted to cattle, horses and pigs by various blood-sucking insects found to be infected during epidemics, including sandflies (Lutzomyia spp), blackflies (family Simuliidae) and biting midges (Culicoides).

- **fomites**—animals can be infected by contaminated food, water, milking machines and tack.

- **mechanical** spread by human hands.

- **semen and embryos**—in bovines transmission via semen is thought to be possible. Transmission is considered unlikely via embryos, but has not been ruled out definitively. In pigs and horses, the virus is present in and can be transmitted via semen contaminated with vesicular fluid. Virus is known to be present in ova.

While vector-borne transmission to humans is possible, it is most likely to occur through inhaling infective aerosols or contact with infective fluids when handling infected livestock.

Disease dynamics

Much remains unknown about disease dynamics as outbreaks in other countries have proven to be erratic and unpredictable. It is thought that domestic animals are probably not the primary hosts of VS virus. What is known is that:

- the incubation period ranges from 2–8 days.
- most cases occur in adults, and young cattle and horses under a year of age are uncommonly affected.
- typically, 5–20 per cent of the animals in a herd show clinical signs, but up to 100 per cent seroconvert.
- infectivity diminishes rapidly and may be lost within a week after the vesicles rupture (thus, virus is shed in vesicular fluids and saliva for a few days only).
- mortality is low and morbidity is very variable, ranging from 5 per cent to more than 90 per cent.
- deaths are very rare in cattle and horses, but higher mortality is seen in some pigs.

Persistence of the agent

VSV:

- is relatively unstable and readily inactivated by sunlight, and therefore does not survive for long periods in the environment except in cool, dark places.
- can remain infectious for 3–4 days on fomites.
- can be inactivated with a wide range of disinfectants (e.g. 1% sodium hypochlorite, 70% ethanol, iodophore, quaternary ammonium disinfectants).
Diagnosis and pathology

Clinical signs

VS causes clinical signs resembling foot-and-mouth disease (FMD) in cattle and pigs, and less often in sheep and goats. However, the disease occurs in horses, which is not the case with other vesicular diseases.

VS can significantly affect production in dairy cattle and performance in horses.

In addition:

- many infections with VS are subclinical
- excessive salivation is often the first sign of the disease
- VS is characterised by vesicles, papules, erosions and ulcers and these lesions are found particularly around the mouth but also may be present on the nares, muzzle, feet, udder and prepuce. Vesicles vary widely in size, with some as small as a pea while others can cover the entire surface of the tongue. In addition, vesicles rupture to become erosions or ulcers. **Frequently this happens before any vesicles are seen.** In some animals, the vesicles may go unnoticed and the disease may appear as crusting scabs on the muzzle and lips
- some animals can have a catarrhal nasal discharge, bleeding from ulcers, or a foetid mouth odour
- transient pyrexia usually develops when lesions appear.

Horses appear to be most contagious for the first 6 days after infection and:

- vesicles occur most often on the upper surface of the tongue, the gums and lips, and around the nostrils and corners of the mouth
- lesions may develop on the teats or prepuce, and on the coronary band and interdigital space of the hooves
- the turbinates, nasopharynx and larynx may be affected, resulting in nose bleeding and difficulty in eating and breathing.

In cattle:

- vesicles are usually found on the hard palate, tongue, lips and gums, sometimes extending to the nostrils and muzzle
- some papules do not become vesiculated
- lesions may develop on the teats or prepuce, and on the coronary band and interdigital space of the hooves
- lesions on the teats can cause mastitis from secondary bacterial infections (particularly in dairy cattle) and weight loss may be severe, reducing milk production
- reduction in milk production can be permanent
- some animals may appear to be normal, but only eat approximately half their normal feed ration.
In pigs, vesicles usually appear first on the feet and the first sign may be lameness. The muzzle and lips are also frequently affected (though the predominant sites of lesions can vary by outbreak).

In humans:
- symptoms resemble influenza
- the incubation period is usually 3–4 days, but may be as short as 24 hours or as long as 6 days
- most people recover without complications in 4–7 days.

Pathology
Gross lesions at post mortem are restricted to the vesicles described. There are no disease-specific histopathological changes.

Differential diagnosis
Although VS may be suspected when horses are involved as well as pigs and cattle, VS is clinically undistinguishable from the other exotic viral vesicular diseases, namely:
- foot-and-mouth disease
- swine vesicular disease
- vesicular exanthema of swine (not seen since 1956)
- Senecavirus A (Seneca Valley virus) infection
- other exotic diseases
  - bluetongue disease (infection present in Australia but without clinical disease)
  - rinderpest (now recognised by OIE as eradicated from all countries with susceptible populations)
- endemic diseases
  - bovine papular stomatitis
  - dermatophilus and other types of mycotic stomatitis
  - footrot
  - infectious bovine rhinotracheitis
  - mucosal disease (BVDV-1 only)
- non-infectious causes
  - phototoxic dermatitis with vesicle formations from contact with the leaves of plants of the family Umbelliferae (parsley, parsnips and celery)
  - chemical irritants and scalding
  - traumatic lesions of the mouth and feet.
Samples required

Because VS may be mild or subclinical, you will need to look for seroconversion to VS in apparently healthy animals to ascertain the length of infection.

Sample collection

Take samples from several affected animals in the herd, flock or group (take these from lesions in the mouth or the feet, or at other sites with suitable lesions).

FMD virus (an important differential diagnosis for VS) is very sensitive to both acid and alkaline conditions. Inappropriate buffer conditions can inactivate the virus making virus isolation difficult or impossible. To maximise chances of virus isolation:

- use phosphate buffered saline or virus transport media with a pH of 7.6
- if submitting a sample after 24 hours or more, add glycerol to the phosphate buffered saline
- dilute oropharyngeal fluid collected with a probang in an equal volume of phosphate buffered saline pH 7.6, and mix vigorously for 1 minute.

Note that sample collection in these buffers is optimal (for the growth of FMD virus) but not essential for RNA or antigen detection assays.

Take samples from at least 10 live, clinically affected animals. Collect:

- **serum**—7–10 ml/animal in plain tubes
- **vesicular fluid**—carefully use a syringe and needle to aspirate the vesicular fluid from unruptured vesicles, and place in a sterile container. Alternatively, collect fluid from small vesicles onto a swab and place the swab in 500 µl of buffer, such as phosphate buffered saline or virus transport medium
- **fresh tissue**—epithelium, epithelial tags, oral, nasal and tonsillar swabs and oropharyngeal fluid and submit in phosphate buffered saline or virus transport medium, if available.

Collect fresh and formalin fixed samples from several tissues (lymph nodes, thyroid and adrenal glands, kidney, spleen and heart) from dead animals. If neurological signs are present, submit the brain.

Transport of samples

For transport:

- chill blood samples and unpreserved tissue samples at either 4°C, or with frozen gel packs
- **DO NOT FREEZE SAMPLES** at –20°C; it reduces the sensitivity when used for virus isolation and molecular diagnostic tests
- send samples with dry ice if the journey is expected to take several days
- formalin fixed tissue can be sent at room temperature.

Sample submission

The relevant state or territory laboratory should coordinate sample packaging and consignment for delivery to CSIRO-AAHL.
Diagnostic tests

VS or other vesicular diseases should be suspected when vesicles or erosions are found on the mouth and/or feet of livestock. Other vesicular diseases must be ruled out with laboratory tests.

VS can be identified using an antigen enzyme-linked immunosorbent assay (ELISA), polymerase chain reaction (PCR), virus isolation, and serological assays for the detection of antibodies. Testing strategies are based on samples submitted and clinical and epidemiological information provided.

Reporting requirements

VS is no longer an OIE-listed disease; however, it is a nationally notifiable disease and there is an obligation to report cases of VS. If you suspect VS, report the disease immediately by phoning the Emergency Animal Disease Watch Hotline on 1800 675 888, wherever you are in Australia. Alternatively, contact a government veterinarian in your state or territory.

Biocontainment and personal protective equipment

VSV infection is a public health risk. Human infections may result from direct contact with infected animals. Isolate suspected cases of VS and keep them on the property until advised by government veterinary authorities, noting:

• wear appropriate personal protective equipment (coveralls and boots, gloves, mask) when examining animals, performing post-mortem examinations, or handling tissue, carcasses or abortion materials
• personal decontamination is required when leaving infected premises. Thoroughly clean and then disinfect vehicles and equipment before leaving the property.

Further information


FIGURE 3.26.1 Extensive erosion at the mucocutaneous junction of an equine lip

Image credit: PIADC and CFSPH

FIGURE 3.26.2 Extensive ulceration of a bovine dental pad, with hypersalivation

Image credit: PIADC and CFSPH
FIGURE 3.26.3 A large vesicle (bulla) on the dorsal snout of a pig

Image credit: Iowa State University, College of Veterinary Medicine, and CFSPH

FIGURE 3.26.4 Severe erosion on the distal teat of a cow

Image credit: Iowa State University, College of Veterinary Medicine, and CFSPH