

7th edition

SWINE DISEASE DIAGNOSTIC MANUAL

A COMPREHENSIVE GUIDE TO
SWINE DISEASE DIAGNOSIS

Respiratory
Enteric
Multi-systemic
Neurologic
Reproductive



va  inova



Vaxxinova is a highly focused, technology-based company dedicated to providing timely, science-based solutions to food-animal disease problems. Our products and services are delivered and supported by a dedicated and experienced sales staff and Veterinary Service team.

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Swine Diagnostic Submission Guide

Disease Suspected	Specimen	Sample Preparation	Laboratory Procedure
<i>Actinobacillus pleuropneumoniae</i>	Lung	Refrigerate	Culture, Sensitivity, Whole Genome Sequencing
		10% Formalin	Histopathology
Arthritis	Joint Fluid, Joint Swab, Synovium	Refrigerate	Culture, Sensitivity
<i>Clostridium perfringens</i>	Duodenum, Jejunum, Ileum, Colon	Refrigerate	Anaerobic Culture, Sensitivity, Whole Genome Sequencing
		10% Formalin	Histopathology
<i>Clostridioides (Clostridium) difficile</i>	Duodenum, Jejunum, Ileum, Colon, Fecal Swabs, Colon Content	Refrigerate	Culture
		10% Formalin	Histopathology
<i>Escherichia coli (E. coli)</i> Colibacillosis Edema Disease	Duodenum, Jejunum, Ileum, Colon, Brain with Brainstem, Fecal Swabs, Colon Content	Refrigerate	Culture, Sensitivity, Whole Genome Sequencing
		10% Formalin	Histopathology
Enteritis (non-specific)	Duodenum, Jejunum, Ileum, Colon, Fecal Swabs, Colon Content	Refrigerate	Culture, Sensitivity, Anaerobic Culture, TGEV qPCR, Whole Genome Sequencing, Rotavirus qPCR
		10% Formalin	Histopathology
Erysipelas	Heart, Lymph Node, Liver, Spleen, Synovium, Joint Fluid	Refrigerate	Culture, Sensitivity, Whole Genome Sequencing
		10% Formalin	Histopathology
<i>Glaesserella (Haemophilus) parasuis</i> (Glässer's disease – polyserositis)	Lung, Pleural Fluid, Heart, Pericardial Fluid, Pericardium, Synovium, Joint Fluid	Refrigerate	Culture, Sensitivity, qPCR, Whole Genome Sequencing
		10% Formalin	Histopathology
Hemorrhagic Bowel Syndrome (HBS)	Duodenum, Jejunum, Ileum, Colon	Refrigerate	Gross lesions and culture to rule out mimic infections
		10% Formalin	Histopathology
Ileitis (<i>Lawsonia intracellularis</i>)	Ileum, Spiral Colon, Cecum, Feces	Refrigerate	qPCR
		10% Formalin	Histopathology
Mulberry Heart Disease	Lung, Pericardial Fluid, Heart, Left and Right Ventricles	Refrigerate	Culture to rule out septicemia
		10% Formalin	Histopathology
<i>Mycoplasma hyorhinis</i> (Polyarthritis and Polyserositis)	Lung with Pleura, Pericardium, Pericardial Fluid, Synovium, Joint Fluid	Refrigerate	Mycoplasma Multiplex qPCR, Culture
		10% Formalin	Histopathology
<i>Mycoplasma hyosynoviae</i> (Polyarthritis)	Synovium, Joint Fluid	Refrigerate	Mycoplasma Multiplex qPCR, Culture
		10% Formalin	Histopathology
<i>Mycoplasma pneumonia (M. hyopneumoniae)</i>	Lung	Refrigerate	Mycoplasma Multiplex qPCR, Culture
		10% Formalin	Histopathology
	Serum	Refrigerate	Serology

Swine Diagnostic Submission Guide

Disease Suspected	Specimen	Sample Preparation	Laboratory Procedure
Pasteurella Pneumonia and Rhinitis (<i>P. multocida</i>)	Lung, Turbinates if Investigating Rhinitis	Refrigerate	Culture, Sensitivity, Serogrouping PCR
		10% Formalin	Histopathology
Porcine Circovirus Disease (PCV2, PMWS, PDNS)	Lung, Spleen, Liver, Lymph Nodes, Kidney, Ileum, Pancreas, Serum	Refrigerate	qPCR, Virus Isolation
		10% Formalin	Histopathology
Porcine Epidemic Diarrhea (PEDV)	Duodenum, Jejunum, Ileum, Colon, Colon Content or Feces, Fecal Swabs, Serum	Refrigerate	qPCR, VI
		10% Formalin	IHC, Histopathology
Porcine Reproductive & Respiratory Syndrome (PRRS)	Lung	Refrigerate/Frozen	qPCR, Virus Isolation, Full Genome Sequencing
		10% Formalin	Histopathology, IHC
	Serum	Refrigerate	Serology, qPCR, Virus Isolation, Full Genome Sequencing
	Blood Swab, Semen	Refrigerate	qPCR, Virus Isolation
Reproductive Diseases/ Abortions (Parvovirus, Lepto, PRRS, etc)	Fetus, Mummies, Stillborns, Weakborns, Serum	Refrigerate	Culture, Sensitivity, IgG, PCR
Rotavirus Enteritis	Duodenum, Jejunum, Ileum, Colon, Colon Content or Feces, Fecal Swabs	Refrigerate	qPCR, Virus Isolation, Whole Genome Sequencing
		10% Formalin	Histopathology
Salmonella	Duodenum, Jejunum, Ileum, Colon, Liver, Lung, Spleen, Mesenteric Lymph Node, Colon Content or Feces, Fecal Swabs	Refrigerate	Culture, Sensitivity, Whole Genome Sequencing
	Duodenum, Jejunum, Ileum, Colon, Liver, Lung	10% Formalin	Histopathology
<i>Streptococcus suis</i>	Brain Including Cerebellum and Brain Stem, Lung, Liver, Spleen, Synovium	Refrigerate	Culture, Sensitivity, Whole Genome Sequencing
		10% Formalin	Histopathology
Swine Dysentery (<i>Brachyspira hyodysenteriae</i>)	Feces, Cecum, Spiral Colon	Refrigerate	Culture, Sensitivity
		10% Formalin	Histopathology
Influenza A Virus – Swine (IAV-S)	Trachea, Lung, Nasal Swabs (in Viral Transport Media)	Refrigerate	qPCR, Virus Isolation, Full Genome Sequencing, HTSN
	Lung	10% Formalin	Histopathology, IHC
	Serum	Refrigerate	Serology (HI/ELISA)
Transmissible Gastroenteritis (TGEV)	Duodenum, Jejunum, Ileum, Colon, Colon Content or Feces, Fecal Swabs, Serum	Refrigerate	qPCR, VI
		10% Formalin	Histopathology, IHC
Gastric Ulcers	Stomach	Refrigerate	Gross Lesions

For bacterial culture, we recommend swabs with transport media to prevent desiccation.

For virus isolation, swabs should be placed into viral transport media; see Sample Submission Guidelines on next page or call the lab for information.

FREE Diagnostic Submission Kits available.

Sample Submission Guidelines

Laboratory test results are directly affected by animal selection, necropsy technique, specimen selection and handling, how well-preserved samples are, and speed of shipment to the laboratory.

The following sections provide an overview and best practices to consider. Vaxxinova provides diagnostic services in support of the development of custom-made vaccines. Initial, exploratory diagnostic investigations involving tissues and necropsy should be conducted at university laboratories. Isolates can then be forwarded to Vaxxinova, or specific follow-up diagnostic samples in support of an ongoing investigation can be submitted directly to Vaxxinova.

Contact Vaxxinova at +1 800 220 2522 or email us at info.us@vaxxinova.com if you have any questions regarding sample collection or the diagnostic process.

PRE-COLLECTION CONSIDERATIONS



Timing

Populations of pathogens change over time; therefore, diagnostic monitoring should be performed on an ongoing basis. When selecting animals for laboratory analysis, they should be free from antibiotic therapy and in an early or acute disease stage. Pigs that are displaying typical clinical signs and immediately necropsied will yield the most reliable diagnostic data.



Histories

A meaningful history of the disease outbreak and a tentative diagnosis, based upon clinical evaluation and necropsy findings, should be included. This information can help laboratories determine the most appropriate diagnostic tests for your case. It can also help guide follow-up testing that may be required, based on preliminary results.

For any materials submitted to Vaxxinova for analysis, Vaxxinova solely owns the work developed or derived from the materials submitted as unique work product and an invention by Vaxxinova. All written materials and other works which may be subject to copyright, and all patentable and unpatentable inventions, ideas, improvements or discoveries conceived or made by Vaxxinova arising out of the developments, shall be the sole and entire property of Vaxxinova. Any and all intellectual property rights related to the vaccine and the development of the vaccine belong solely to Vaxxinova.



Quantity and Volume

Tissue samples should be collected from at least two or three humanely euthanized pigs, ideally in the early stages of disease. Oral fluids, processing fluids, nasal swabs and serum should be collected from appropriate numbers of animals or litters based on the diagnostic test(s) being requested. For more information about specific sample types and appropriate sampling protocols, please contact the Vaxxinova Customer Service team or continue reading.

Sample Submission Guidelines



Method and System of Collection

Selected specimens should be collected using aseptic technique whenever possible to reduce the risk of contamination. Tissue specimens should also be kept separate to avoid cross-contamination. Diagnostic kits are available from Vaxxinova. Contact the Vaxxinova Customer Service team for assistance.



Specimen Identification

Clearly identify any sample you wish to submit. All samples should include the following information:

- **Farm ID number**, including site and building, when applicable
- **Animal ID number**
- **Source of material** (i.e., oral fluid, nasal swab, joint fluid, tracheal wash, tissue type, etc.)



Fixative

The selected tissues should be fixed in 10 percent neutral buffered formalin. Use 10 times the volume of the tissues being fixed to ensure good perfusion of the sample and to maintain the tissue architecture. After 24 hours of fixation, excess formalin can be safely poured off, and a smaller formalin volume can be then be used for shipping.

NEUTRAL BUFFERED FORMALIN FORMULAS

37–40% formaldehyde.....	100 mL
Distilled water.....	900 mL
Sodium phosphate, monobasic monohydrate.....	4.0 g
Sodium phosphate, dibasic anhydrous.....	6.5 g



Use the appropriate collection method and transport system for the desired specimen type. Details by type are provided in the Collection Considerations section on page 8.



Temperature

It is important that the samples arrive at the lab before they degrade. To help extend viability, as soon as a sample is collected, keep it refrigerated or on ice until and during shipment. Be sure to include enough cold packs for delivery time. For assistance determining this, contact the Vaxxinova Customer Service team.

Sample Submission Guidelines

COLLECTION CONSIDERATIONS



Fresh Tissues

TARGET:
Collect sample from visible lesions with adjacent normal tissue.

SIZE:
Approximately 2 by 4-inch samples.

TRANSPORT SYSTEM:
Double-bag in Whirl-Pak® bags.

CONSIDERATIONS:
Do not mix swabs, intestines or brains with other tissues in a single bag. Remember to clearly label each bag.



Swabs

Aerobic Culture

TRANSPORT SYSTEM:
Commercial swab.

MEDIA TYPE:
Stuart or Amies recommended to prevent desiccation.

MEDIA VOLUME:
Minimum 1 mL.

CONSIDERATIONS:
Remember to clearly label each swab.

Anaerobic Culture

TRANSPORT SYSTEM:
Port-A-Cul™ (BBL™) or other anaerobic transport system. (The Port-A-Cul tube can be used for anaerobic, facultative and aerobic bacteria.)

CONSIDERATIONS:
For abscesses or exudates, use a capped syringe with the needle removed or a tube with a snug cap.



Nasal Swabs

Bacterial Suspect

TARGET:
Nasal cavity.

TRANSPORT SYSTEM:
Stuart or Amies.

CONSIDERATIONS:
Clean the external nares and internal nostrils with a moist towel to remove common contaminants. Insert the swab into the nasal cavity and rotate. Once sample is collected, return the swab to the accompanying sterile plastic sheath. Crush the ampule at the base of the sheath to release the transport medium.

For bacterial isolation, use bacterial media and avoid using Mycoplasma or viral media, which contain antimicrobials that may inhibit growth of the desired pathogen.

Viral Suspect

TARGET:
Nasal cavity.

TRANSPORT SYSTEM:
Universal Viral Transport Kit (Becton Dickinson #220528) or equivalent.

CONSIDERATIONS:
Prepare the nostrils and sample as described above in the Bacterial Suspect section.

For viral isolation, use Mycoplasma or viral media and avoid using bacterial media. Use of incorrect swab and media may jeopardize the ability to detect or culture the pathogen of interest.



Oral Fluids

Oral fluids can be collected from pens of pigs by hanging a length of clean cotton rope in a way that allows pigs to chew on it, but without contacting areas of excessive feces or other debris. After pigs have sufficiently saturated the rope, the fluid can be squeezed from the rope into a plastic bag and transferred to a sterile screw-/snap-top tube.



Processing Fluids

Processing fluids are collected during castration and tail docking. Tissue should be placed into a clean container lined with a plastic bag and allowed to sit overnight in a refrigerator. The fluid that has collected in the bottom of the bag should then be transferred to a sterile screw-/snap-top tube.



Histopathology

TARGET:
Samples should be taken from multiple sites or types of lesions and should include both normal and diseased tissue, as well as the line of demarcation.

SIZE:
No more than 1 inch thick. Smaller sizes of tissues result in a more rapid and complete penetration of the fixative.

TRANSPORT SYSTEM:
Doubled Whirl-Pak® and 10 percent formalin solution.

Sample Submission Guidelines

CONSIDERATIONS:
Selected tissues should be cut with a sharp knife or scalpel, since the squeezing action of scissors crushes and tears tissues.

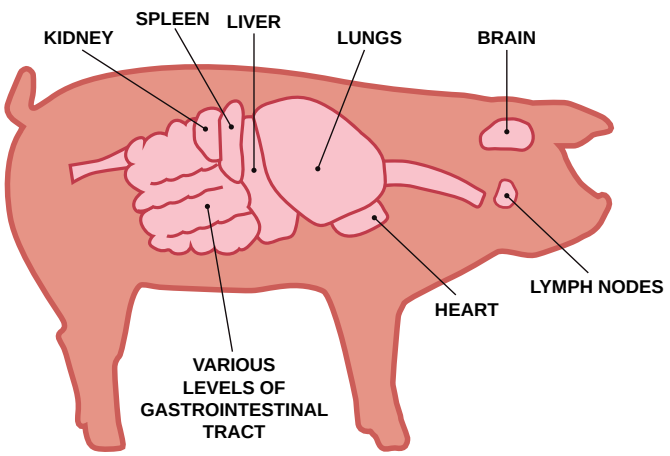
Autolysis and freezing will make samples unsuitable for histopathological evaluation.

For information about how to fix samples and prepare the fixative solution for histopathological analysis, see the Neutral Buffered Formalin Formulas on page 7.

If hollow organs (gut or uterus) retain significant amounts of content, they should be gently flushed with 10 percent formalin — without disturbing the mucosal lining — before being placed in the formalin bag.

CAUTION: Be sure to take proper precautions when handling and disposing of formalin.

Check the recommended samples in the Swine Diagnostic Submission Guide on pages 6-7. If the cause of death is unknown or the clinical syndrome is vague, then submit samples exhibiting gross lesions and sections from all of the following:



Sample Submission Guidelines



Blood Samples

SIZE:
1–3 mL non-hemolyzed blood, depending on the number of tests.

TRANSPORT SYSTEM:
Sterile tubes (serum separator).

CONSIDERATIONS:
Fill vacutainer tubes three-fourths of the way full, and allow to stand at room temperature for an hour to permit a solid clot to form and retract. Pipette the serum into sterile tubes with snap caps. (3-mL plastic tubes with snap caps, Falcon #2054, are recommended.)

POST-COLLECTION AND SHIPPING CONSIDERATIONS



Diagnostic Shipping Kits and Submission

Vaxxinova provides free diagnostic shipping kits for sample submission, which include all necessary forms and shipping materials.

When filling out diagnostic submission forms, remember these key considerations:

- Clearly specify the test(s) being requested.
- Use one form per client site, and identify the sample tubes by different barns or age groups, depending on the test(s) being requested.
- When sending paired sera, distinguish the acute samples from the convalescent samples on the tube and on the form.



- Make sure caps are securely closed to avoid leaking during transit.
- Use permanent markers and underline the ID numbers to indicate orientation (e.g., 16 vs. 91).
- Do not freeze whole blood or samples with clots.
- Contaminated or toxic samples cannot be used in virus isolation tests.



To request shipping kits or submission forms email us at info.us@vaxxinova.com or call us at +1 800 220 2522.

Sample Submission Guidelines



Packing Specimens

To avoid leaking in transit, double-bag all samples. Whirl-Pak® bags or equivalent are recommended.

Wrap sample bags and two to four ice packs in absorbent paper (e.g., newspaper).

Place the package into a styrofoam container. Completed submission forms should be inserted into a separate bag in case of leakage and clearly attached to the matching specimens.

This is especially important if your package contains specimens from multiple clients and/or sites.

Avoid mixing intestinal samples with other tissues.

If you need more information about shipping samples to Vaxxinova, please call us at +1 800 220 2522.



Mailing

Diagnostic samples should be submitted by the fastest means possible to avoid deterioration of specimens. Next-day or overnight delivery is preferred.

We recommend these reliable services:

- UPS
- FedEx
- Spee-Dee
- U.S. Postal Service as last resort (above services preferred over USPS)

Hours and Contact Information

Vaxxinova is open for service from 8:00 a.m. to 5:00 p.m. (CST) Monday through Friday, with the exception of holidays.

Please email us at info.us@vaxxinova.com or call +1 800 220 2522 to request a diagnostic shipping kit or submission forms. Visit us online at www.vaxxinova.us.com for additional information.

Our mailing address for diagnostic sample submission and other general inquiries is listed below.

Diagnostic Shipping Address

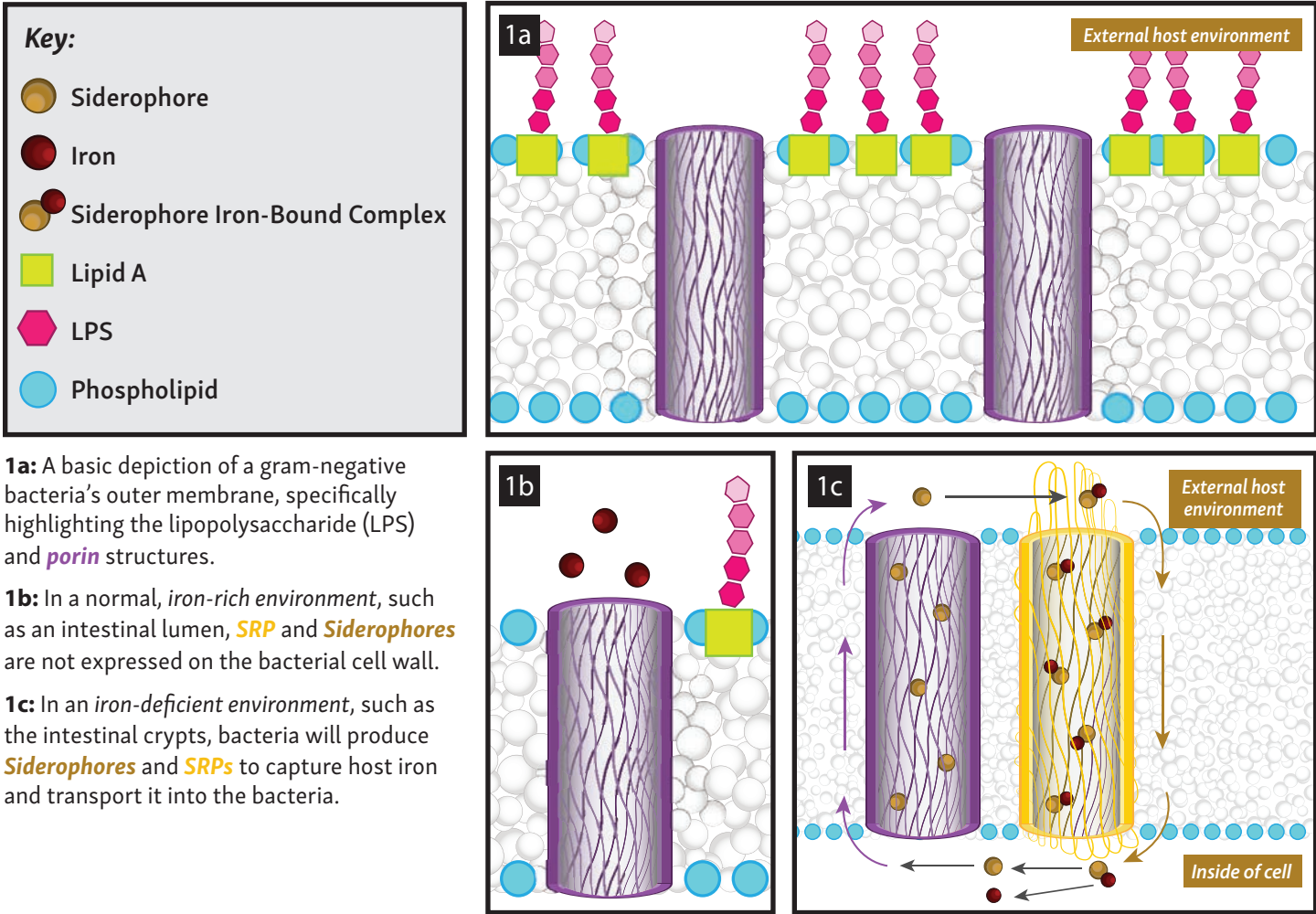
va^{xx}inova

Vaxxinova US
1524 Prairie Drive
Worthington, MN 56187

General Adjuvant and Manufacturing Considerations

SRP Technology

SRP technology is a highly purified, antigenic vaccine that helps induce a broad immune response to gram-negative bacteria. These bacteria, like *E. coli* and *Salmonella*, become problematic when they are unable to absorb iron, turning on protein receptors called Siderophore Receptor Proteins (SRP) to collect host-bound iron. With these SRPs activated, the damage begins, destroying the host’s ability to absorb nutrients, while expressing toxins. To prevent this damage from occurring, Vaxxinova scientists have found a way to express, collect, and purify SRPs; thus, manufacturing a vaccine with the SRP antigens, allowing the host to develop a highly specific immune response to those pathogenic gram-negative bacteria.



Amplivac™ (formerly TS6)

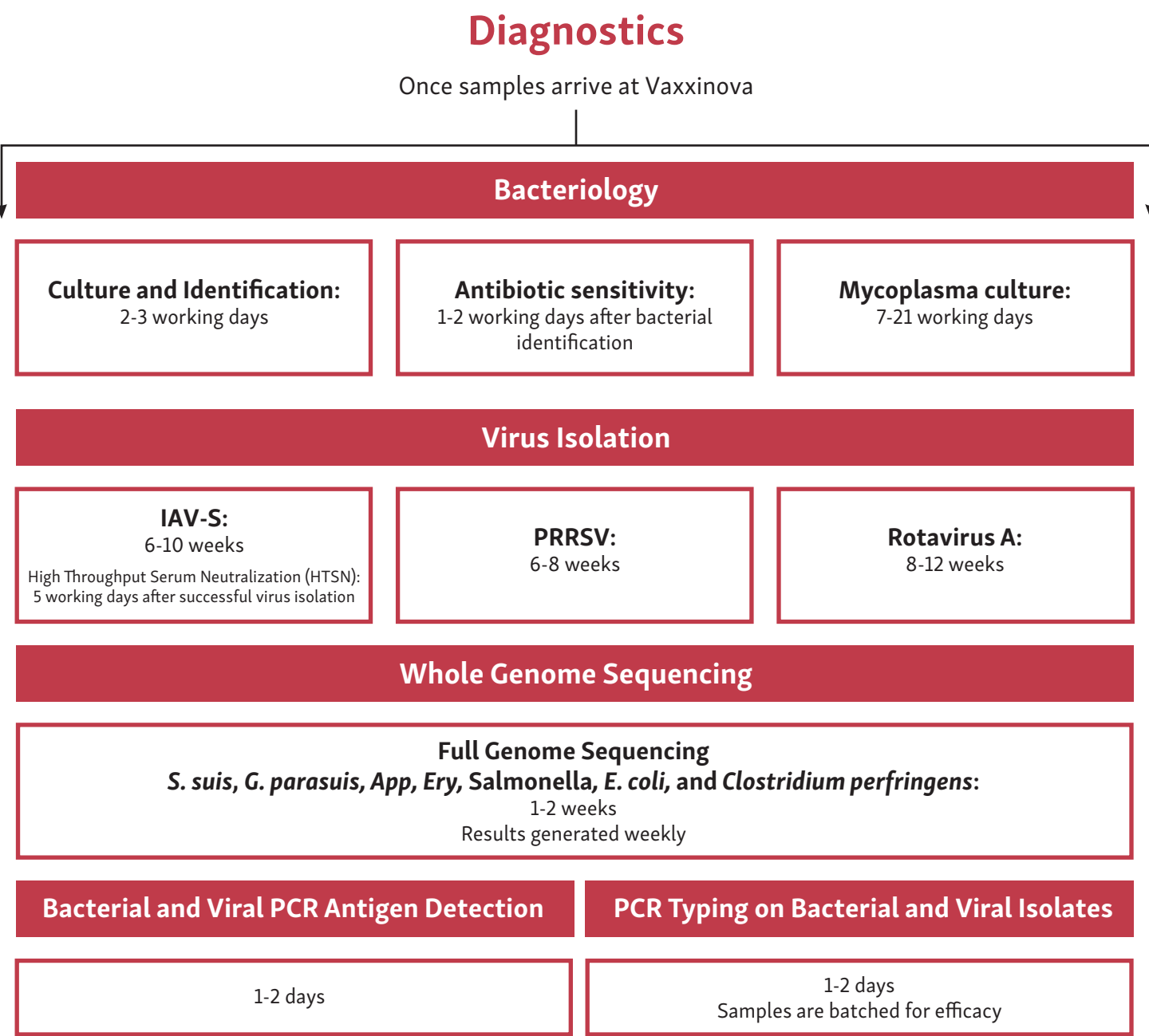
This is the best all-around adjuvant, particularly for viruses, *Strep. suis* and *Mycoplasma* custom-made vaccines. Previous Vaxxinova challenge studies support this. Amplivac can cause mild-to-moderate local injection-site reactions. To minimize the risk of not achieving minimum antigen concentrations and possibly reactions within animals and the bottle of vaccine, a maximum of four isolates should be included in a custom-made vaccine with Amplivac. If more isolates are needed, consult with Vaxxinova customer service.

Trigen™

Trigen is another effective Vaxxinova adjuvant that is used in specific situations when certain Gram-negative bacteria (such as *Actinobacillus suis*) need to be included in a custom-made vaccine.

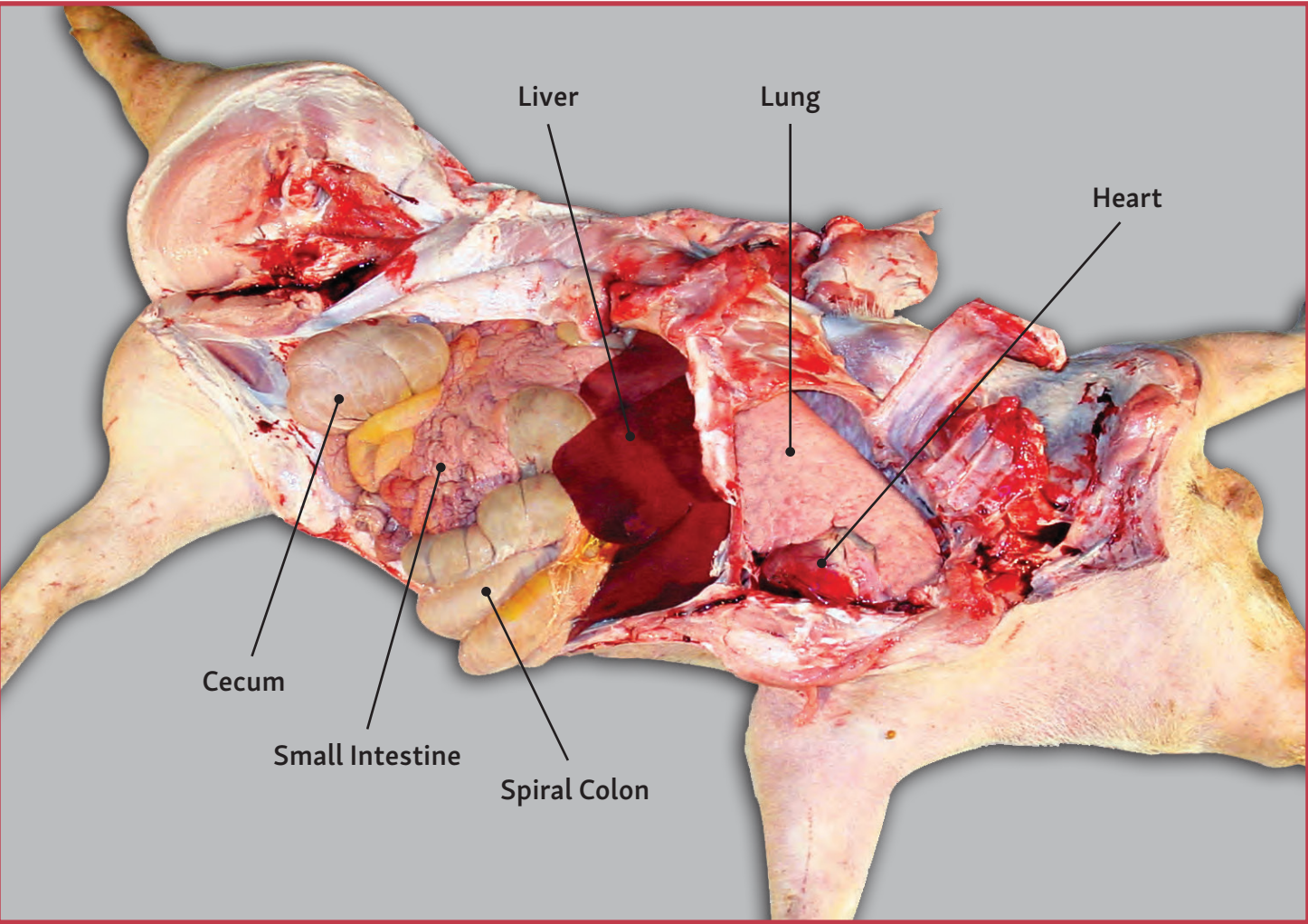
Diagnostics

Overall, timelines vary by the diagnostic tests that are requested. In most circumstances, diagnostic work is required prior to beginning vaccine production, which is critical for helping understand which isolates are most appropriate for vaccine development. If this is the first time an isolate is being considered for vaccine production, these diagnostic timeline estimates are in addition to production timelines. Furthermore, consider any other diagnostic timelines from University Veterinary Diagnostic Laboratories that might be part of an initial disease workup. In general, Vaxxinova’s diagnostic capabilities are primarily aimed at providing specific support of vaccine manufacturing, rather than complete diagnostic evaluations.

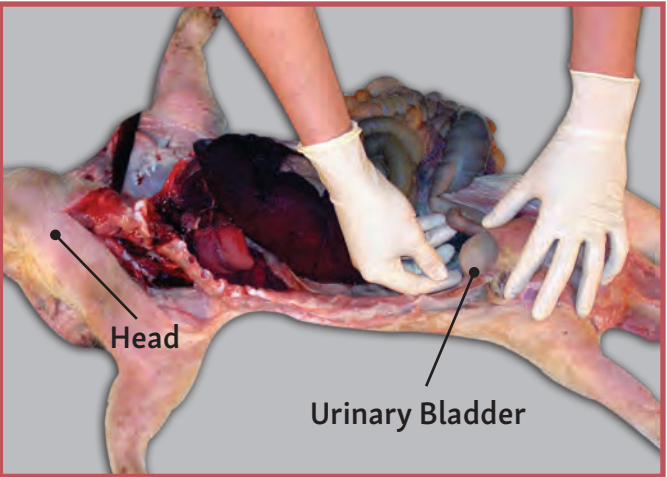
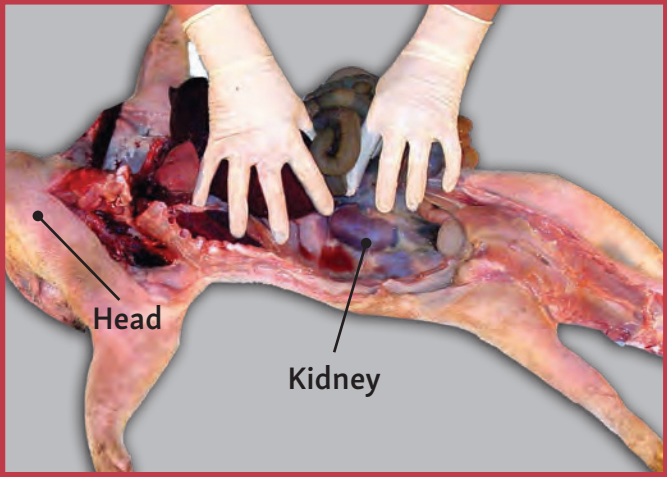
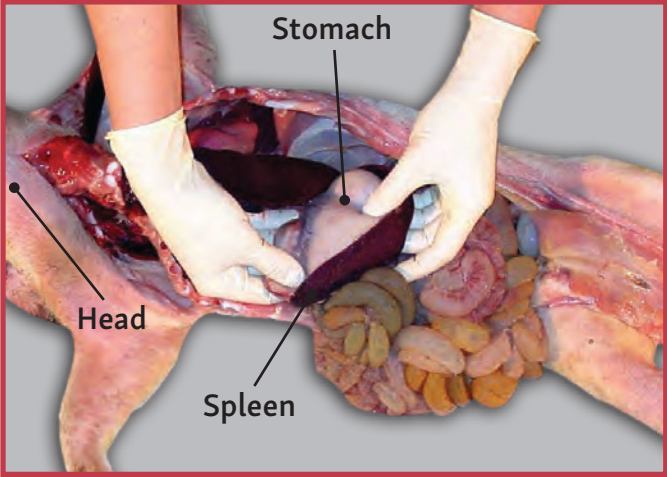


Clostridium perfringens, *Pasteurella multocida*, *E. coli* fimbriae
Actinobacillus pleuropneumoniae, *Glaesserella parasuis*
PRRS, IAV-S, subtype
**Anticipated these diagnostics will move into whole genome sequencing in the future.*

Major Pig Organs



Major Pig Organs



Nursery Pig Necropsy Instructions



Important: Start with a sharp knife.



Cut through the axilla to partially separate the front limb from the rib cage. Repeat for other side.



After cutting through the axillae, the pig will lie upright on its back.



Hook the knife under the cranial sternum. Cut through the cartilage of all the ribs on both sides.

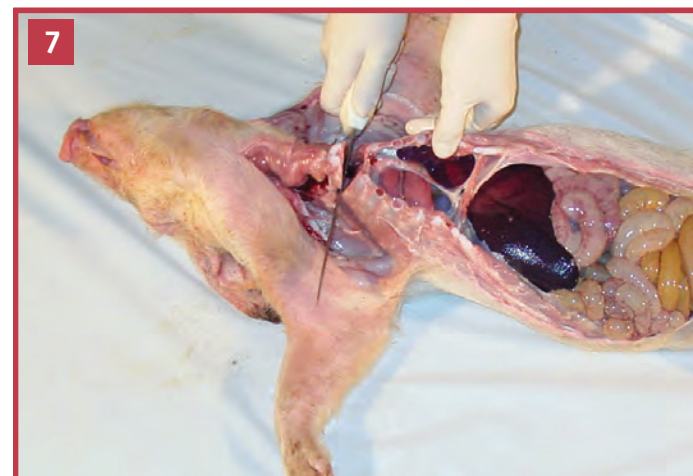


Continue this cut to remove the skin with sternum, and the ventral abdominal wall (belly) of the pig.

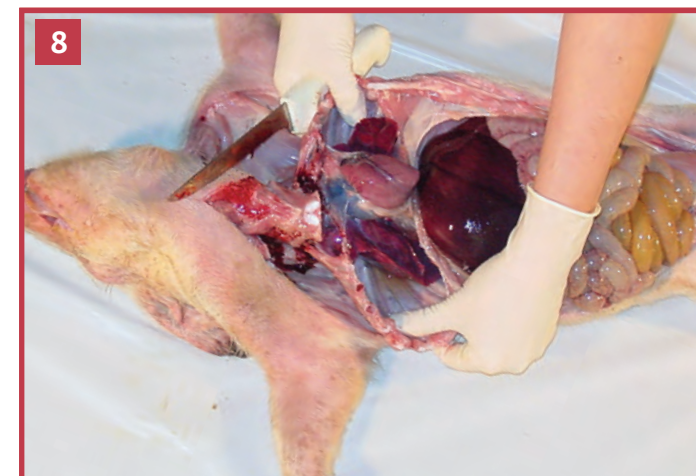


Most organs are now visible.

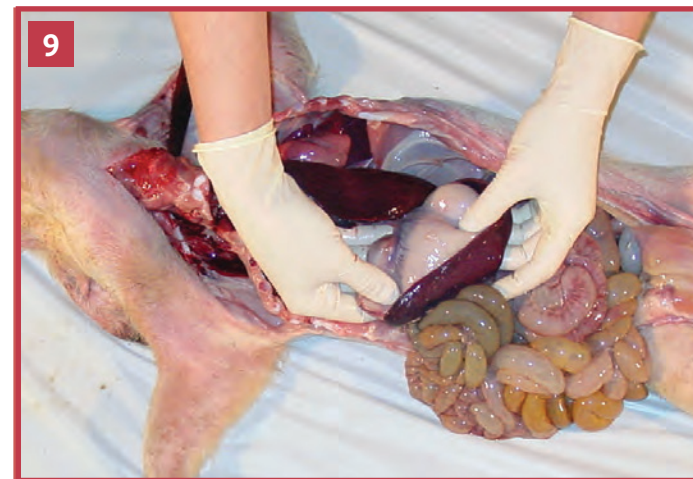
Nursery Pig Necropsy Instructions



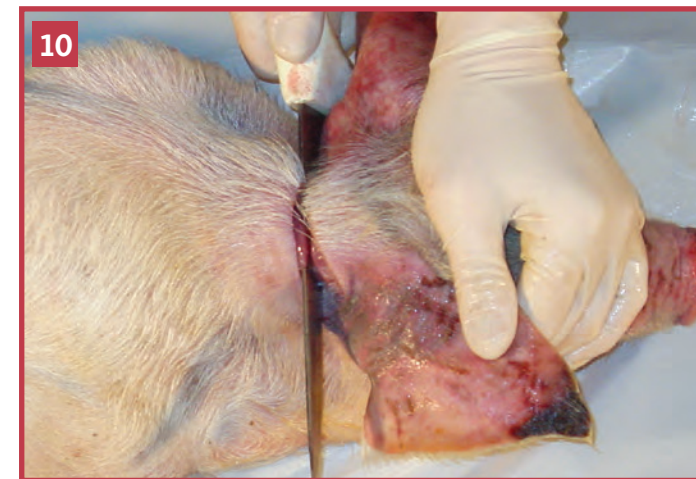
Cut between the ribs below the collarbone.



Spread and crack the ribs open.



Organs are easily examined.



Brain Swabs: Cut through the skin and muscle behind the ears at the base of the skull.



Flex the nose toward the floor and the ears down to open the space between the top vertebra and the skull. This will allow room to cut between them.



Place a swab on the exposed spinal cord toward the brain. This is an excellent way to test for strep.

Grower-Finisher Necropsy Instructions



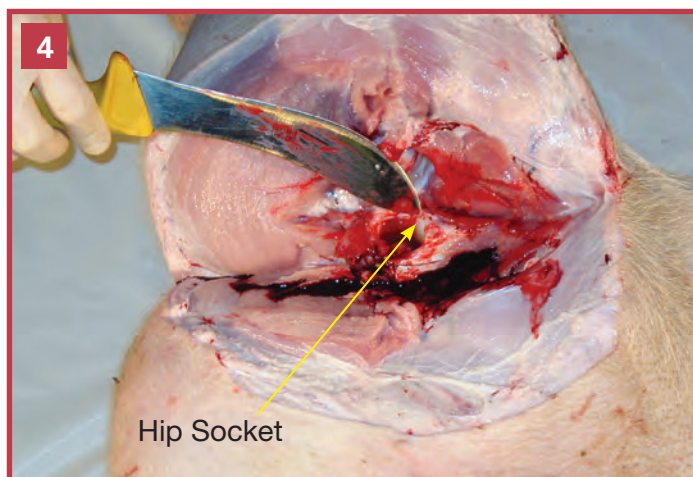
With the pig on its side, hold the lower front limb down with your foot while pulling up on the upper front limb.



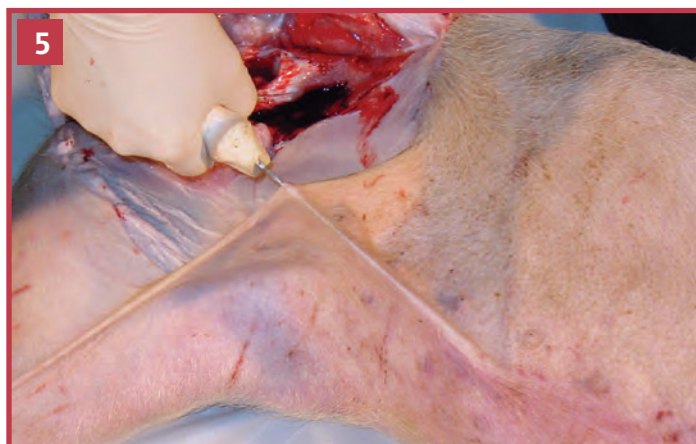
Use the knife to cut through the axilla (armpit) to separate the leg from the rib cage.



The upper hind limb is cut and laid back likewise.



As you push the hind limb back, up and over the hip by cutting muscle in the area, the hip socket will become exposed. Cut through the socket and continue pushing the limb straight back over the hip.



Cut between the skin and body wall, beginning at the pelvis, along the midline all the way to the neck.



Continue the cut along the ventral midline toward the neck.

Grower-Finisher Necropsy Instructions



Following the cut just made, dissect the skin away from the body wall, reflecting it over the back.



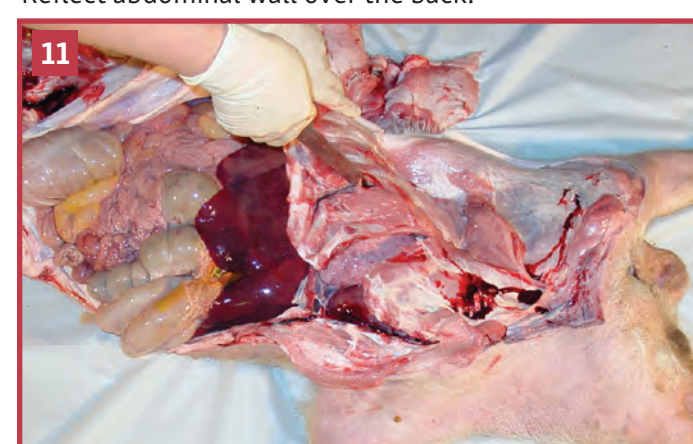
Continuation of previous step. Note that abdominal wall and back muscles are being exposed, but abdomen is not open.



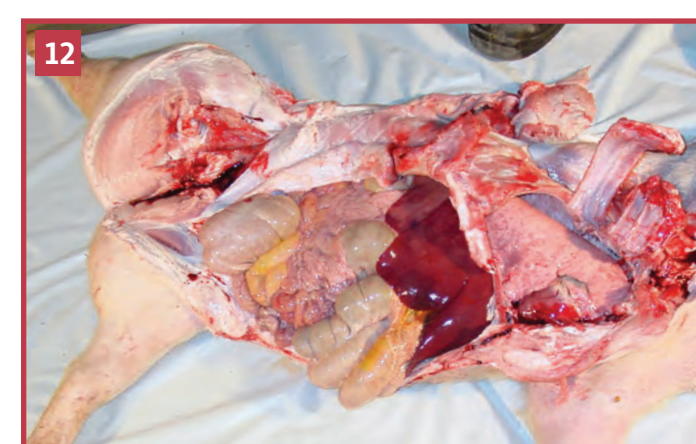
Carefully open the abdomen wall without cutting into intestines or urinary bladder beginning near the pelvic floor and working toward the head along the midline. Reflect abdominal wall over the back.



Puncture the diaphragm near caudal sternum. Cut through cartilage of the sternum all the way to neck.



Cut muscles between ribs in pairs; break ribs by pushing one or two at once over the back.



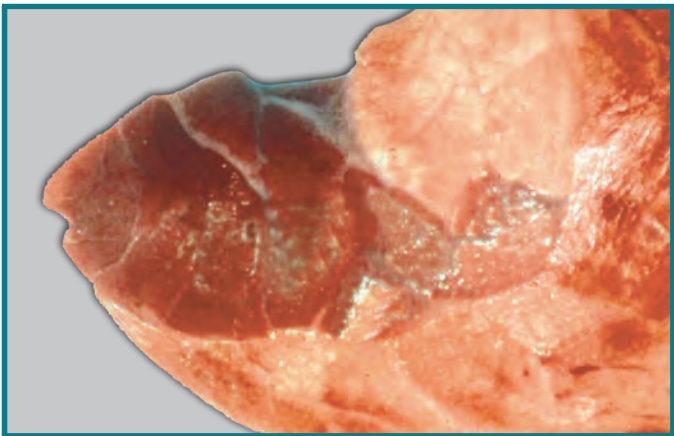
Organs are now exposed for examination.

Mycoplasma Pneumonia

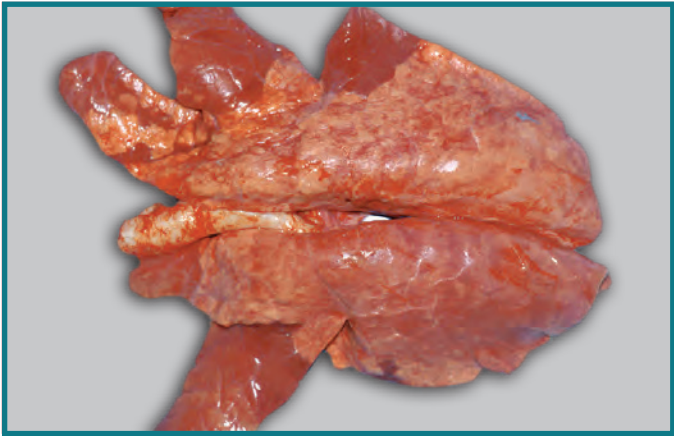
Mycoplasma hyopneumoniae



Normal lung



Purplish-tan consolidated lung without pleuritis.



Purplish-tan consolidated lung without pleuritis. Note cranioventral involvement.



Purplish-tan consolidated lung without pleuritis. Note cranioventral involvement.

MYCO

Tissues to Submit

- Lung
- Serum

Diagnostic Tests

- Mycoplasma Multiplex PCR
- Mycoplasma Culture
- Histopathology
- Serology

Clinical Signs and History

- Coughing is the most common sign and is most obvious when pigs are roused.
- Sporadically, individual pigs or groups develop severe pneumonia.
- Often accompanied with secondary *Pasteurella multocida* or other bacterial infections.

Stages of Production

- Nursery
- Grow-Finish

Diagnosis

- Affected lung tissue is gray or purple, most commonly in the apical and cardiac lobes (cranioventral).
- Lesions are clearly demarcated from normal lung.
- The associated lymph nodes may be enlarged.

Pneumonic Pasteurellosis

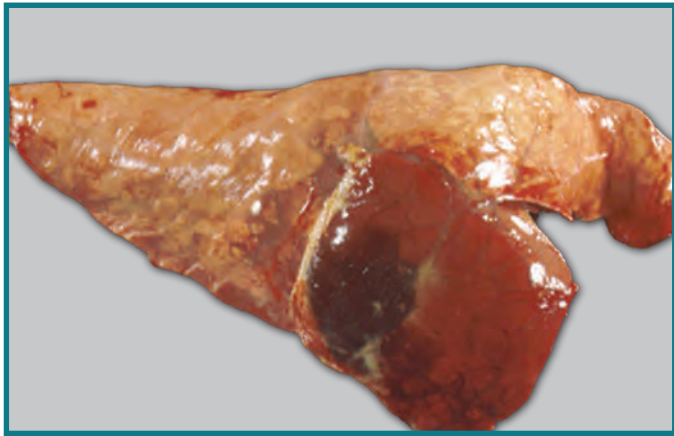
Pasteurella multocida



Normal Lung



Mycoplasma Lung



Mycoplasma and Pasteurella Lung

PASTEURELLA

Tissues to Submit

- Lung

Diagnostic Tests

- Culture
- Sensitivity
- Serogrouping PCR

Clinical Signs and History

- Respiratory: Cough, rapid breathing (thumping).
- A temperature of 104° – 107° F develops, and there is anorexia, depression and occasionally mild rhinitis and dyspnea with coughing.
- Some pigs become lame with painful, warm, swollen joints.
- Chronic arthritis and occasionally meningitis and convulsions may develop.

Stages of Production

- Nursery
- Grow-Finish

Diagnosis

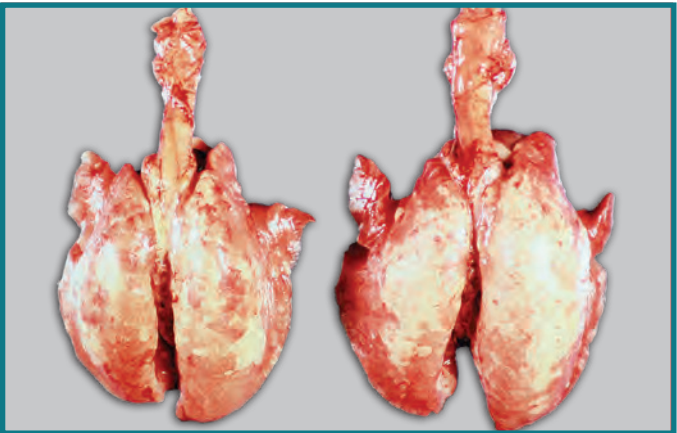
- Diagnosis is based on necropsy findings and culture of *P. multocida* from the lesions.
- Exudative bronchopneumonia, sometimes with pericarditis and pleuritis.

PRRSV

Porcine Reproductive and Respiratory Syndrome



Normal Lung



Lungs stay upright due to diffuse interstitial pneumonia; slightly rubbery but not consolidated.



Lungs stay upright due to diffuse interstitial pneumonia.



Ventral view of PRRSV-infected lung.

PRRSV

Tissues to Submit

- Lung
- Serum

Diagnostic Tests

- PCR
- Virus Isolation
- Sequencing
- Serology
- Histopathology
- IHC

Clinical Signs and History

- Respiratory: Cough, rapid breathing (thumping), unthrifty pigs.
- Reproductive: late-term abortions after 90 day's gestation with fresh and autolyzed piglets, stillborns, weak live piglets.

Stages of Production

- Gestation
- Farrowing
- Nursery
- Grow-Finish

Diagnosis

- Characteristic lesions and organism identification.
- Diagnosis is based on herd history and virus isolation (VI) or viral antigen testing (PCR or IHC).

Influenza A Virus – Swine (IAV-S) / Pneumonia



Normal Lung: pink, collapses uniformly, soft upon palpation



Dorsal view of IAV-S pneumonia: Lungs stay upright due to diffuse interstitial pneumonia with patchy red lobules of consolidation; palpates rubbery compared to normal.



IAV-S pneumonia: Diffuse interstitial pneumonia as described above with cranioventral consolidation related to secondary bacterial bronchopneumonia (arrow).

IAV-S

Tissues to Submit

- Lung
- Nasal Swabs
- Serum
- Trachea

Diagnostic Tests

- PCR
- Virus Isolation
- Full Genome Sequencing
- HTSN
- Serology (HI, ELISA)
- Histopathology
- IHC

Clinical Signs and History

- Respiratory: Rapid spread of severe cough throughout the barn, rapid breathing (thumping), depression, fever to 108° F, anorexia, dyspnea, weakness, prostration and a mucous discharge from the eyes and nose.
- Outbreak is characterized by sudden onset and rapid spread through the entire herd, often within one to three days.

Stages of Production

- Farrowing
- Nursery
- Grow-Finish

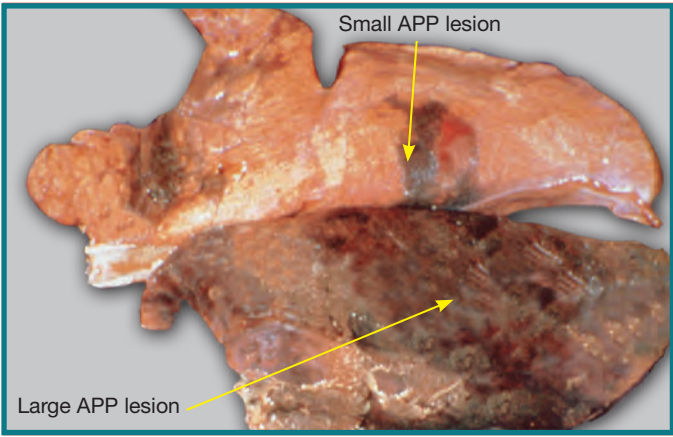
Diagnosis

- In uncomplicated infections, lesions are usually confined to the lungs.
- Necrotizing bronchiolitis becomes proliferative in chronic cases; IAV-S is confirmed by IHC or PCR or VI.
- The airways contain a copious mucopurulent exudate, and the bronchial and mediastinal lymph nodes are edematous and enlarged.

Swine Pleuropneumonia *Actinobacillus pleuropneumoniae* (APP)



Normal Lung



Hemorrhagic necrotic lung lesions, with pleuritis involving dorsal lung at small lesion. Extensive lobar consolidation in larger lesion.



Cross section of APP lung showing distinct demarcated areas of multi-lobular necrotizing and hemorrhagic pneumonia with pleuritis.

APP

Tissues to Submit

- Lung
- Serum

Diagnostic Tests

- Culture
- Sensitivity
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- Respiratory distress is severe; there is thumping and occasionally open-mouth breathing with a blood-stained frothy nasal and oral discharge, fever, anorexia and reluctance to move.

Stages of Production

- Grow-Finish

Diagnosis

- An explosive disease onset is suggestive.
- The pneumonia is usually bilateral, but often unevenly distributed with unique dorsal and caudal lung lobe involvement.
- The characteristic lesion is a severe fibrinonecrotic and hemorrhagic pneumonia with accompanying fibrinous pleuritis.
- In acute cases, the lesions are sharply delineated, dark consolidated regions that ooze bloody fluid from the cut surface. The involved pleura and inter-lobular septa are thick with exudate.
- The trachea may contain blood-stained froth. Bloody nasal/oral discharge is common.
- In chronic cases, the lesions are more organized and adhesions between the lung and rib cage become fibrous.

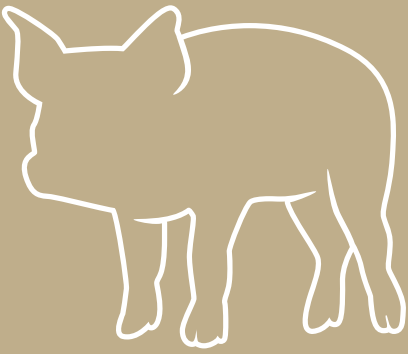
Diarrheal Diseases at Various Ages

Infection with and/or agent	Suckling Piglets	Nursery Pigs	Grow/Finish Pigs	Adults
Porcine Sapovirus	++++	+++		
PRRSV	++++	+++		
Seneca Valley Virus	++++	++		
Non-Hemolytic <i>E. coli</i>	++++	++		
Hemolytic <i>E. coli</i>	+++	++++	+ (early grower)	
Rotavirus	++++	+++	+ (early grower)	
Transmissible Gastroenteritis Virus (TGE)	++++	+++	+++	+++
<i>Clostridoides (Clostridium) difficile</i>	++++			
<i>Clostridium perfringens</i> Type A	++++			
<i>Clostridium perfringens</i> Type C	++++	+ (rare)		
Coccidiosis <i>Isopara suis</i>	++++	++	+	
Salmonellosis	+	++	++++	+
Swine dysentery <i>Brachyspira hyodysenteriae</i>	+	++	++++	++
Proliferative enteropathies <i>Lawsonia intracellularis</i>		++	++++	++
Porcine Epidemic Diarrhea Virus (exotic)	++++	++++	++++	++++
Whipworm infection <i>Trichuris suis</i>		++	++++	+++
Hemorrhagic Bowl Syndrome			+++	+++

Key: Clear cells indicate groups that seldom, if ever, are infected; 1+, 2+, and 3+ indicate increasing frequencies of infections with 4+ indicating the age groups that most often are infected.

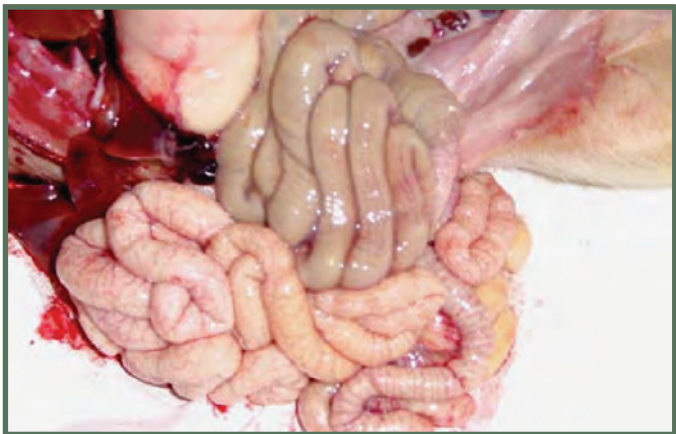
Did you know diarrhea in young pigs is often a multifactorial etiology? Here are a few examples of what can increase the chances of scours:

- Diet change
- Pig movement
- Temperature change
- Enteric, and some respiratory pathogens
- Mineral content in drinking water



Clostridial Enterocolitis

C. perfringens & *C. difficile*



Intestinal distension – *C. perfringens* Type A.



Dark-red intestine, gas bubbles visible beneath the serosa. Lumen will be filled with bloody necrotic content. *C. perfringens* Type C.

Tissues to Submit

- Small Intestine
- Large Intestine
- Colon
- Colon Content
- Fecal Swabs

Stages of Production

- Farrowing
- Nursery

Diagnostic Tests

- Culture
- Sensitivity
- Whole Genome Sequencing
- Histopathology



Mesocolonic edema – *C. difficile*

CLOSTRIDIUM

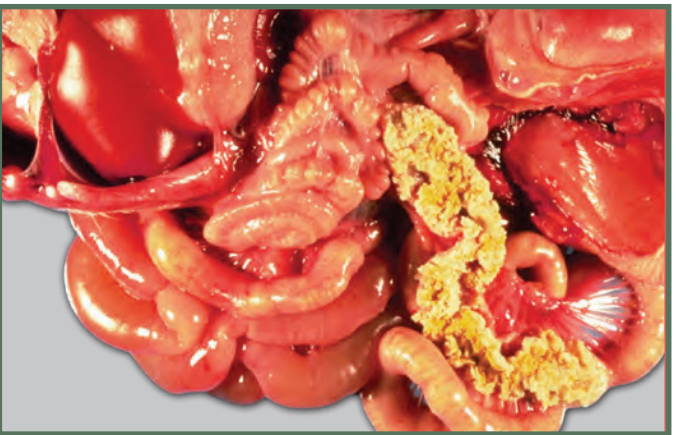
Clinical Signs and History

- Diarrhea is the most common sign in enteric clostridial infections.
- Sudden onset of hemorrhagic diarrhea followed by collapse and death is characteristic in piglets 1–3 days old as a result of *Clostridium perfringens* Type C.
- *Clostridium perfringens* Type A and *Clostridioides (Clostridium) difficile* most frequently cause diarrhea without hemorrhage in pigs 3–15 days of age.

Diagnosis

- Necropsy is usually sufficient to establish the diagnosis of *C. perfringens* Type C in the peracute hemorrhagic form and in the acute form with jejunal emphysema. Histologic observation of villous necrosis with mucosal colonization by numerous large Gram-positive rods is adequate for confirmation.
- Isolation and identification of the organism is necessary to diagnose *C. perfringens* Type A and *C. difficile*.
- *C. perfringens* Type C – In acute cases, gas bubbles (gut emphysema) will be visible through the serosa and within the mucosa. The disease is often segmental, and normal areas can be adjacent to severely diseased areas. Important to find and submit specimens from diseased areas.
- *C. perfringens* Type A – Lesions are much milder than seen with *C. perfringens* Type C and are similar to those seen with *E. coli*.
- Mesocolonic edema can be seen in *C. difficile* and *C. perfringens* Type A cases.

Coccidiosis



Coccidiosis: note diffuse dull necrotic yellow-tan membrane covering the normally shiny mucosa.

COCCIDIOSIS

Tissues to Submit

- Small Intestine
- Large Intestine
- Fecal Swab

Diagnostic Tests

- Smear
- Histopathology

Clinical Signs and History

- Diarrhea in farrowing house caused by *Isospora suis*, usually after 5 days of age. The disease is most intense from 7–14 days of age. Less common in nursery pigs, where it can be associated with *I. suis* or other types of coccidia.
- Clinical signs of coccidiosis are due to destruction of the intestinal epithelium and, frequently, the underlying connective tissue of the mucosa.
- Infection is characterized by a watery or greasy diarrhea, usually yellowish to white and foul smelling. Piglets may appear weak, dehydrated and undersized; weight gains are depressed and sometimes piglets die.

Stages of Production

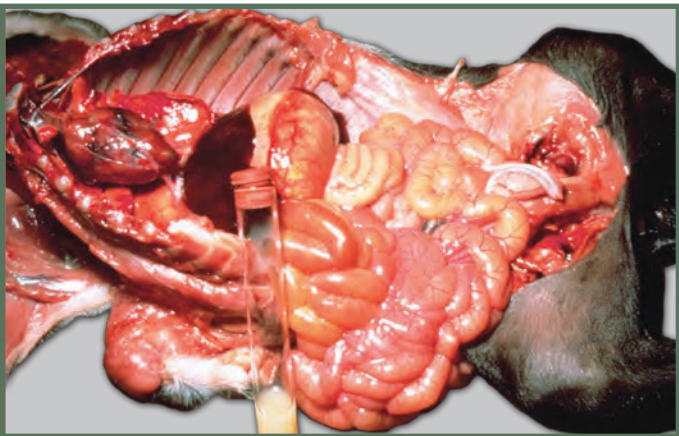
- Farrowing house disease after 5 days of age; intense between 7-14 days of age
- Nursery (less common)

Diagnosis

- Diagnosis is by histopathological observation of sporozoites in the diseased mucosa; or by finding sporozoites in mucosal smears via direct microscopic examination.

Intestinal Colibacillosis and Edema Disease

E. coli



Distension of the intestine with yellowish fluid.



Litter with diarrhea.



Pig down on side, paddling.

E. COLI

Tissues to Submit

- Small Intestine
- Large Intestine
- Fecal Swab
- Brain

Diagnostic Tests

- Culture
- Sensitivity
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- Diarrhea
- Sudden Death
- CNS (Edema Disease)

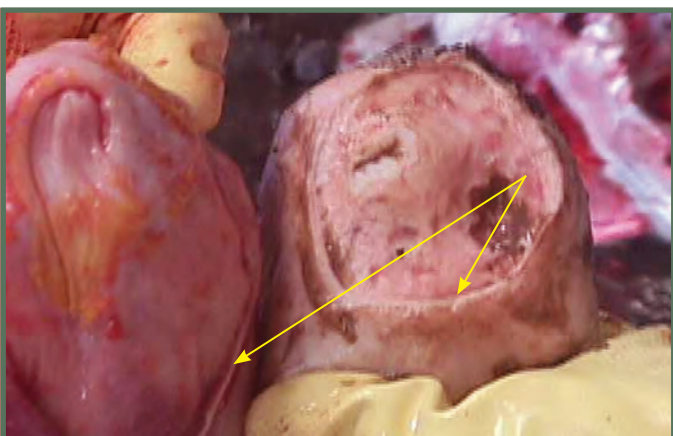
Stages of Production

- Farrowing
- Nursery

Diagnosis

- Confirmation is based on histologic observation of villous colonization and isolation of pathogenic *E. coli*.
- Dehydration and distension of the small intestine and colon with yellowish, watery to cream-like fluid. Mesenteric lacteals are still white with milk fat, indicating absorption is still normal, but hypersecretion is producing diarrhea.

Gastric Ulcers



Gastric ulcers in two stomachs. Arrows point to ulcer edges.



Gut content in distal small intestine and colon is dark brown (arrow) due to digested blood coming from the gastric ulcers.

ULCERS

Tissues to Submit

- Stomach

Diagnostic Tests

- Post-mortem Exam

Clinical Signs and History

- Sudden death related to gastric bleeding; hematoma (large blood clot) found in the stomach.
- In the “chronic” form, hemorrhage results in anorexia, weakness, anemia, and black, tarry feces.

Stages of Production

- Grow-Finish

Diagnosis

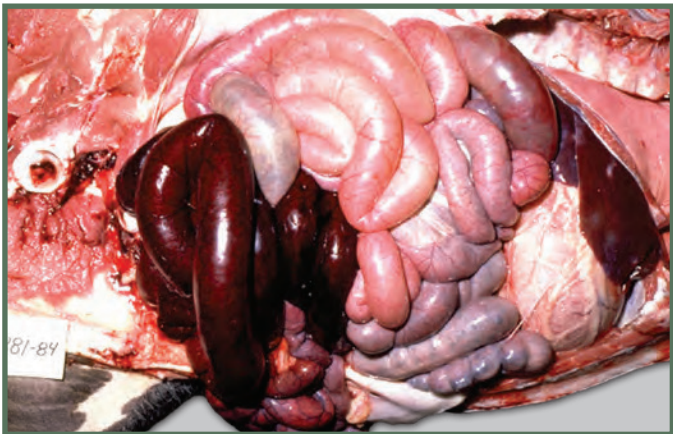
- Appearance in a pen of one or two listless, anorexic pigs that show weight loss, anemia and dark feces.
- Sometimes, dyspnea is suggestive of gastric ulceration, as is the sudden death of an apparently healthy pig.
- The typical terminal ulcer lesion is found in the gastric mucosa near the esophageal opening (cardia) in the rectangular area of white, glistening, non-glandular, squamous epithelia.
- In cases of sudden death, the stomach will contain a large hematoma (blood clot) that originates from a chronic bleeding ulcer.

Hemorrhagic Bowel Syndrome – HBS

Mesenteric Torsion of the Small Intestine

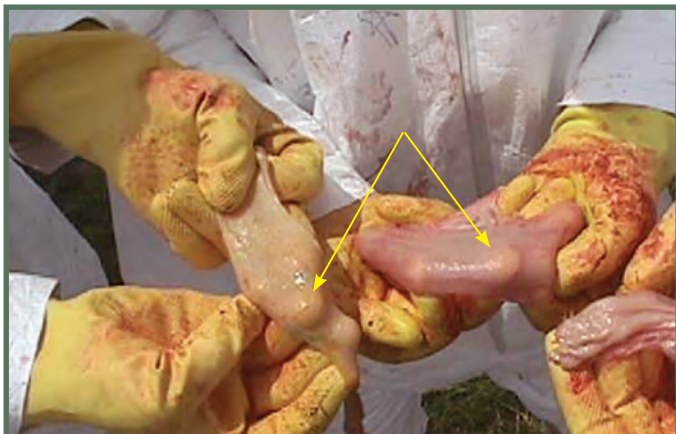


Bright-red gas-distended loops of small intestine with bloody content.

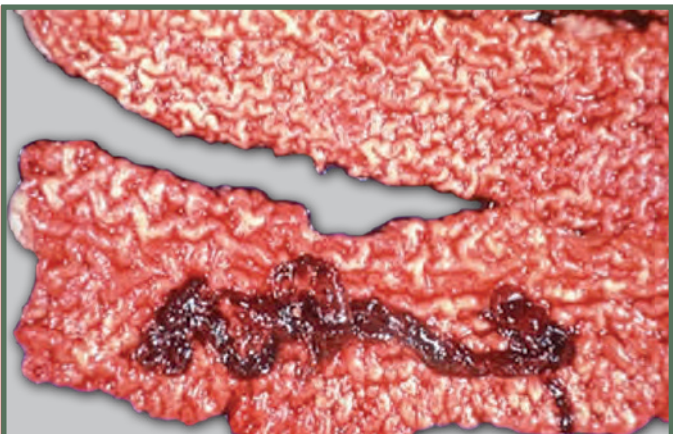


Ileitis

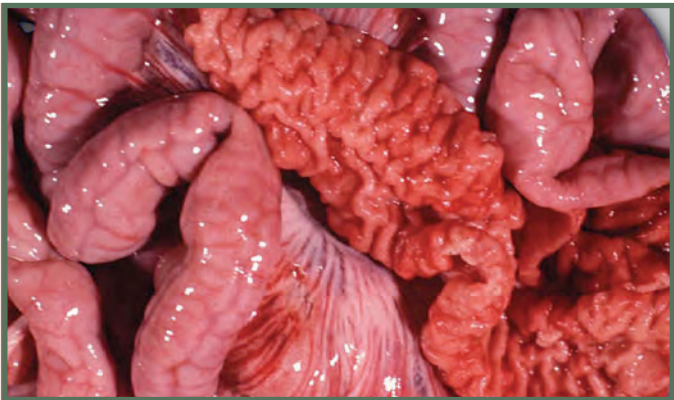
Lawsonia intracellularis



Normal intestine – can see fingers through lining.



Thick hyperplastic ileal mucosa with blood clot in lumen.



Thick hyper-plastic ileal mucosa.

HBS

Tissues to Submit

- Small Intestine
- Colon

Diagnostic Tests

- Post-mortem Exam
- Tests to rule out Salmonellosis, Ileitis, and Swine Dystentery

Clinical Signs and History

- Sudden death of 4-6 month-old grow-finish and young breeding pigs.
- Only involves a few animals; not a large outbreak.

Stages of Production

- Grow-Finish

Diagnosis

- Sudden death of previously healthy grow-finish pigs and characteristic post-mortem findings.
- Before manipulating the intestines, palpate the mesenteric root (tissue coming down from the lumbar back and supporting the gut mass) for a twist or torsion. When present, this is diagnostic for mesenteric torsion. Smaller lesions may only involve a torsion within the mesentery of a portion of the small intestine.
- The involved gut loops are thin-walled, gas-filled, and red due to congestion, and contain bloody fluid.

ILEITIS

Tissues to Submit

- Ileum
- Feces

Diagnostic Tests

- PCR
- Histopathology

Clinical Signs and History

- Diarrhea
- Ileitis can be either a chronic disease in growing pigs, or an acute hemorrhagic form in market-weight and adult pigs.

Stages of Production

- Grow-Finish

Diagnosis

- Confirmation is based on histologic observation of characteristic proliferation and inflammation of mucosal crypts.
- Lesions may occur anywhere in the lower half of the small intestine, cecum or colon, but are most frequent and obvious in the ileum.
- The wall of the intestine is thickened, and the mesentery may be edematous.

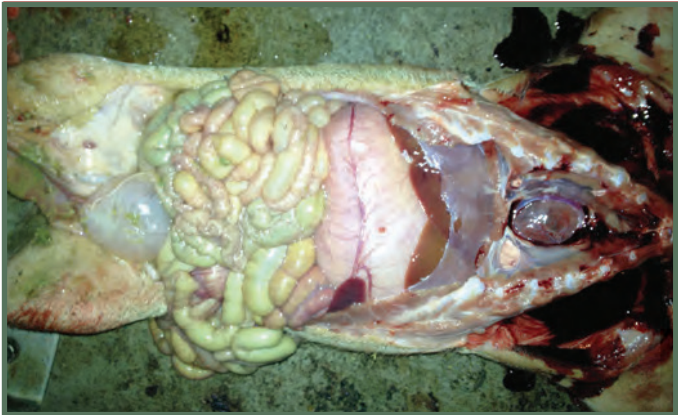
Porcine Epidemic Diarrhea PEDv



Gaunt, dehydrated piglets.



Vomiting sow.



Voluminous, fluid-filled intestinal loops.

PEDv

Tissues to Submit

- Small Intestine
- Colon

Diagnostic Tests

- PCR
- Histopathology

Clinical Signs and History

- The primary clinical sign in outbreaks that occur in previously naïve herds is severe diarrhea in all ages.
- Clinical signs will be essentially identical to those expected with acute TGEv infections.
- Virus is shed in the feces and transmission is via the fecal-oral route.
- The incubation period is 12–24 hours after exposure, with clinically ill pigs shedding virus for 7–10 days.
- Mortality rate in suckling pigs in a naïve herd can be 30–100 percent.

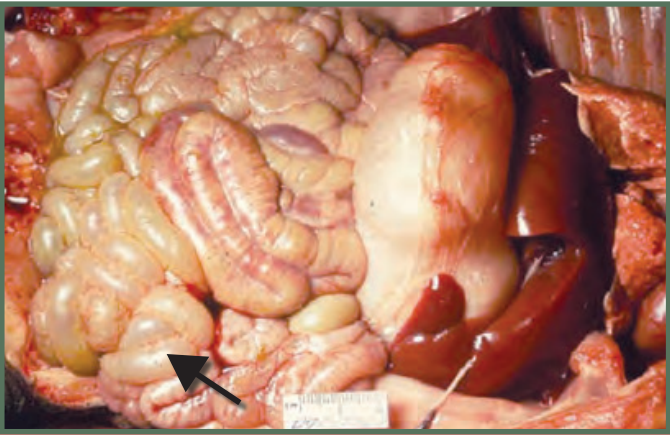
Stages of Production

- Farrowing
- Nursery
- Grow-Finish

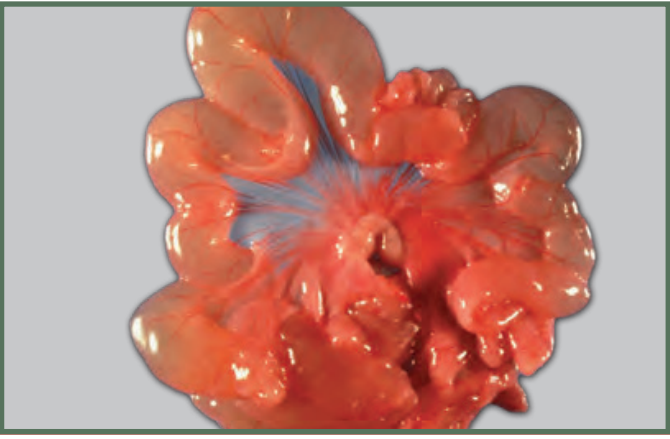
Diagnosis

- Clinical signs with severe diarrhea begin explosively in naïve herds, leading to a presumptive diagnosis of TGEv or PEDv.
- PEDv in naïve herds affects animals of all ages.
- The most common sources of infected feces are pigs, trucks, boots, clothing or other fomites.
- Preferred samples for diagnostic testing are live pigs in acute stages of disease, fresh and formalin-fixed small intestine and colon.

Transmissible Gastroenteritis TGE Virus



Thin-walled Intestine



Thin-walled, fluid-filled intestines. Mesenteric lacteals do not contain milk fat.



Thin-walled Intestine



Weak, gaunt piglet covered with malodorous diarrheal fluids.

TGE

Tissues to Submit

- Small Intestine
- Large Intestine
- Fecal Swab
- Serum

Diagnostic Tests

- PCR
- Histopathology

Clinical Signs and History

- In susceptible herds, vomiting often is the initial sign, followed by profuse watery diarrhea, dehydration and excessive thirst.
- Feces of nursing pigs often contain curds of undigested milk.
- Mortality is nearly 100 percent in piglets < 1 week old, whereas pigs > 1 month old seldom die.
- Gestating sows occasionally abort, and lactating sows often exhibit vomiting, diarrhea and agalactia.
- Diarrhea in surviving nursing piglets continues for five days, but older pigs may be diarrheic for a shorter period.
- Clinically and pathologically mimics PEDv.

Stages of Production

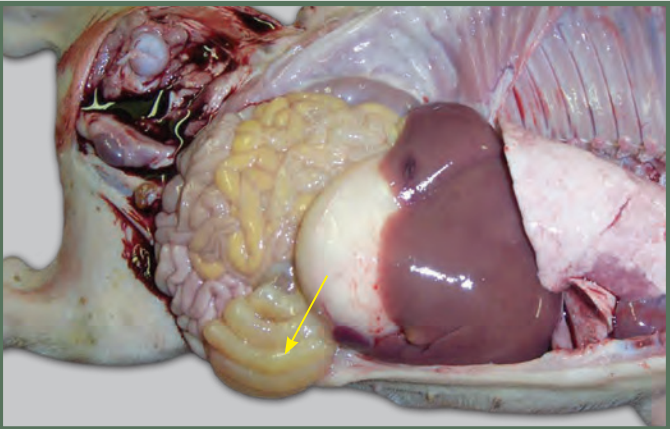
- Farrowing
- Nursery
- Grow-Finish

Diagnosis

- Clinical signs in the epidemic form of TGE usually provide a presumptive diagnosis.
- In the mild endemic form, laboratory procedures are required. Histologic and immunofluorescent examination of the small intestine to demonstrate typical lesions and the presence of TGE viral antigen provide confirmatory evidence.
- Piglets are severely dehydrated and the skin is soiled with liquid feces.
- The stomach usually contains milk curd, but may be empty.
- The small intestine is thin-walled, and the entire intestine contains greenish or yellow watery fluid and clumps of undigested milk.

Rotavirus A

Rotavirus Enteritis



Thin-walled fluid-filled small intestine and spiral colon (arrow).



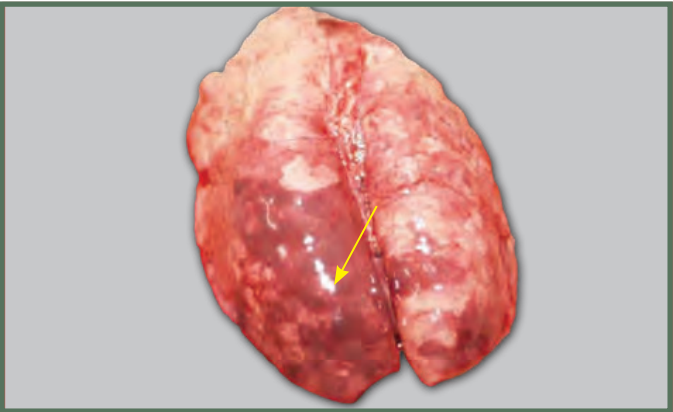
Evidence of diarrhea around anus of a nursery pig.

Salmonellosis

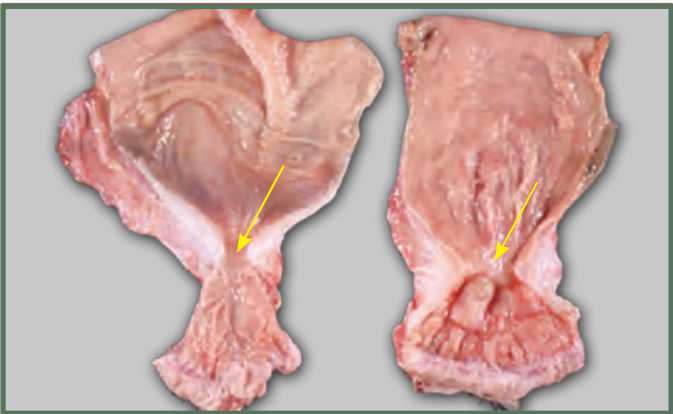
Enteritis and Septicemia



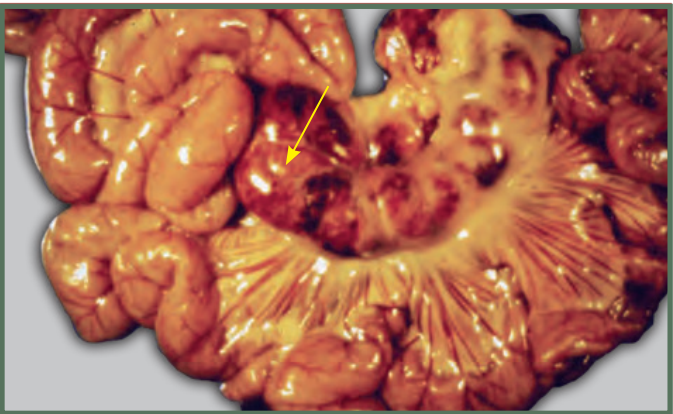
Intestine distended; edematous and thickened wall.



Diffuse interstitial pneumonia with congestion, edema and patchy consolidation (arrow).



Proctitis with rectal stricture.



Swollen lymph nodes.

ROTA

Tissues to Submit

- Small Intestine
- Large Intestine
- Fecal Swabs

Diagnostic Tests

- PCR
- Virus Isolation
- Full Genome Sequencing
- Histopathology

Clinical Signs and History

- Diarrhea
- Commonly, the infection is endemic in a herd. Sows have varying levels of antibody in the colostrum and milk, which provides varying degrees of passive protection to nursery pigs.
- Diarrhea often begins in pigs 5 days to 3 weeks old, and is very common immediately after weaning.

Stages of Production

- Farrowing house and nursery piglets

Diagnosis

- Laboratory procedures are required for accurate diagnosis.
- The small intestine appears thin-walled, and the cecum and colon contain abundant liquid and usually yellow feces.

SALMONELLA

Tissues to Submit

- Small Intestine
- Large Intestine
- Liver
- Lung
- Spleen
- Mesenteric Lymph Nodes

Clinical Signs and History

- Septicemia is the usual syndrome in pigs up to 6 months of age. Illness is acute, depression is marked, fever (105° – 107° F) is common, and death occurs in 24–48 hours. Nervous signs may occur in pigs; these animals may also suffer from pneumonia. Mortality may reach 100 percent.
- Nursing pigs may develop diarrhea, but usually succumb to generalized septicemia.
- Weaning or grow-finish pigs are febrile, and have liquid feces that may be yellow and contain shreds of necrotic debris.

Diagnostic Tests

- Culture
- Sensitivity
- Whole Genome Sequencing

Stages of Production

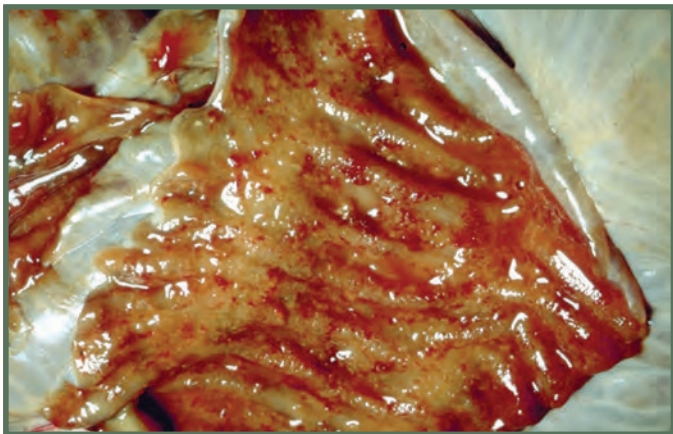
- Farrowing
- Nursery
- Grow-Finish

Diagnosis

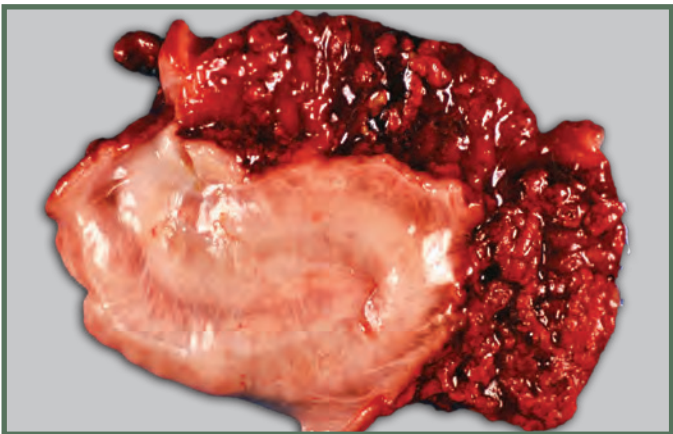
- Depends on the clinical signs and on the laboratory examination (culture) of feces, tissues from affected animals, feed (including all mineral supplements used), water supplies and feces from wild rodents and birds that may inhabit the premises.
- A dark-red to purple discoloration of the skin is common, especially at the ears and ventral abdomen.
- Also, a swollen spleen, liver and lymph nodes can be seen, as well as rubbery congested hemorrhagic lungs and roughened necrotic intestinal mucosa with ulceration and accumulation of debris.

Swine Dysentery

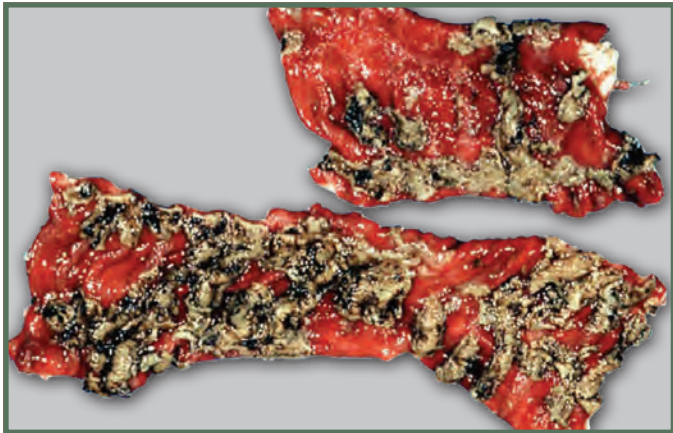
Brachyspira hyodysenteriae



Edematous and hemorrhagic large intestinal mucosa (inner lining).



Thick-walled, hemorrhagic and edematous large intestine.



Fibrinonecrotic debris with dark blood clots on the colon mucosa.



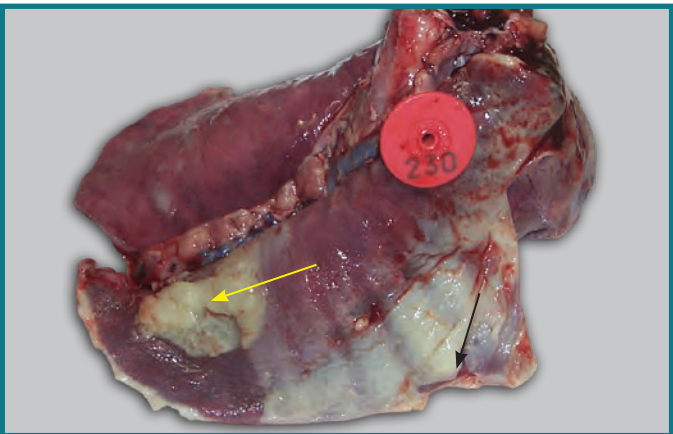
Chronic Swine Dysentery: Less blood, but thickened mucosa covered with adherent yellow-tan necrotic membrane.

Glässer's Disease

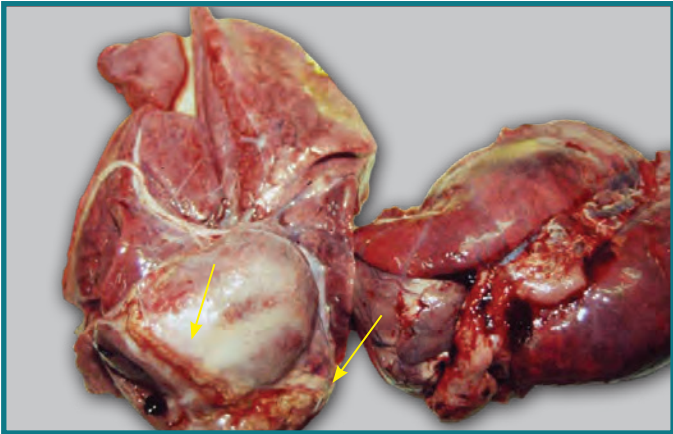
Glaesserella (Haemophilus) parasuis



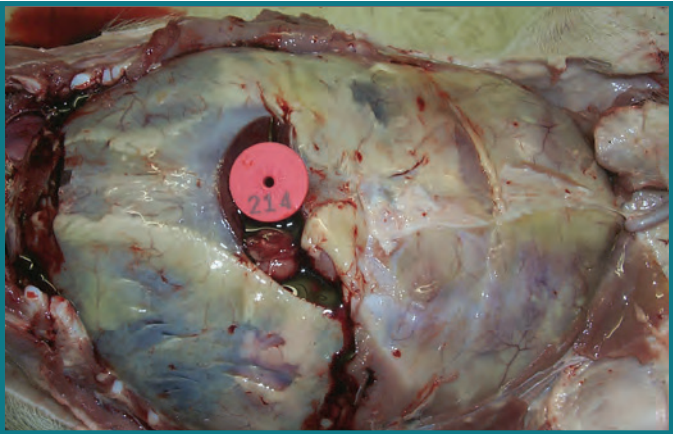
Lame pig



Polyserositis — fibrin-covered lungs



Fibrin on heart



Polyserositis — fibrin-covered abdominal organs

SCOURS

Tissues to Submit

- Feces
- Cecum
- Spiral Colon

Diagnostic Tests

- PCR
- Culture
- Histopathology

Clinical Signs and History

- Bloody diarrhea, affects large intestine, partial anorexia, soft feces, dehydration, profoundly weak, gaunt and emaciated.

Stages of Production

- Grow-Finish

Diagnosis

- Presumptive diagnosis can be based on necropsy and direct examination of smears prepared on slides from fresh colonic mucosa or feces.

Lesions

- Diffuse superficial lesions, confined to cecum, spiral colon and rectum.

GPS

Tissues to Submit

- Synovium
- Joint Fluid
- Meninges
- Lung with Pleura
- Exudate
- Heart with Pericardium

Diagnostic Tests

- PCR
- Culture
- Sensitivity
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- Sudden death.
- A temperature of 104° – 107° F develops, and there is anorexia, depression and occasionally mild rhinitis and dyspnea with coughing.
- Some pigs become lame with painful, warm, swollen joints.
- Chronic arthritis and occasionally meningitis and convulsions may develop.

Stages of Production

- Nursery
- Grow-Finish

Diagnosis

- Based on history, clinical signs and necropsy. Confirmed by culture of the organism from joint fluids, involved tissues, or CSF.
- Polyserositis, polyarthritis, and meningitis.

Erysipelas

Erysipelothrix rhusiopathiae



Diamond, square or rhomboid skin lesions (infarcts) associated with Erysipelas.

ERY

Tissues to Submit

- Heart
- Lymph Nodes
- Liver
- Spleen
- Joints
- Joint Fluid

Diagnostic Tests

- Culture
- Sensitivity
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- Acute septicemia (the skin [subacute] form), chronic arthritis and vegetative endocarditis may occur together or separately.
- Pigs with acute septicemia may die suddenly without previous signs. This occurs most frequently in finishing pigs weighing 100–200 pounds.
- Acutely infected pigs are febrile (104° – 108° F), walk stiffly, and lie on their sternums separately rather than piling in groups. They squeal when handled, and may shift weight from foot to foot when standing.
- Skin discoloration may vary from widespread erythema and purplish discoloration of the ears, snout and abdomen to diamond-square-or rhomboid-shaped skin lesions (infarcts) almost anywhere on the body, particularly the lateral and dorsal areas.

Stages of Production

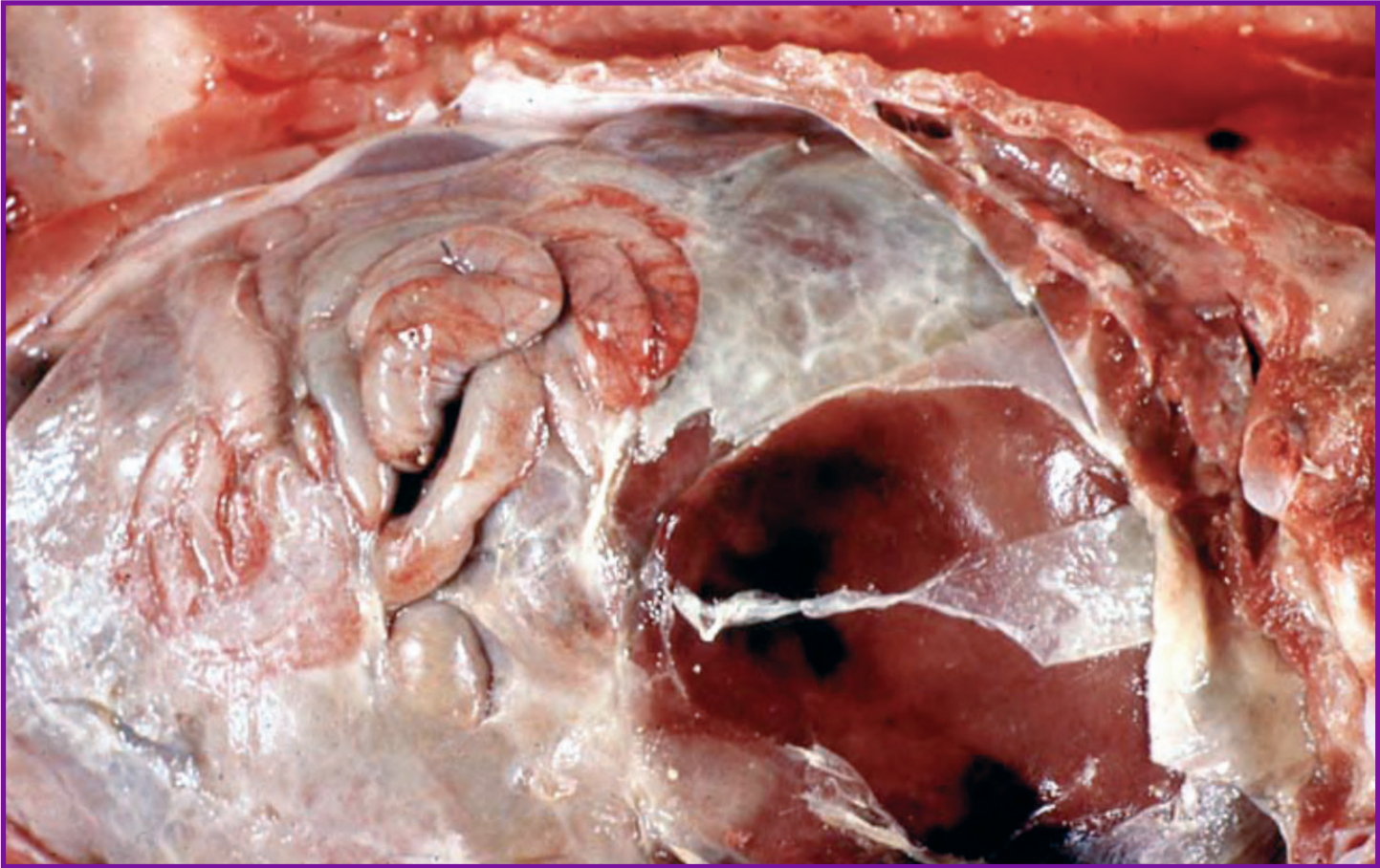
- Grow-Finish

Diagnosis

- Acute Erysipelas is difficult to diagnose in pigs showing only fever, poor appetite and listlessness.
- The typical diamond-shaped skin lesions are highly characteristic when found, but are not always present, and can sometimes be seen with other bacterial septicemias.
- Arthritis and endocarditis are not diagnostic in the live animal because other agents can cause similar syndromes.
- In acute infection, in addition to skin lesions, lymph nodes are usually enlarged and congested, the spleen is noticeably enlarged, and the lungs are edematous and congested.
- Petechiae may be found in the kidneys, heart and occasionally elsewhere.

Polyserositis and Polyarthritis

Mycoplasma hyorhinis



Fibrinous peritonitis and polyserositis over intestinal serosa, peritoneum and liver capsule.

HYORHINIS

Tissues to Submit

- Consolidated Lung
- Heart with Pericardium
- Joint Fluid
- Synovium
- Peritoneal Fluid

Diagnostic Tests

- Myco Multiplex PCR
- Culture
- Serology

Clinical Signs and History

- *M. hyorhinis*, *Strep suis* *H. parasuis* signs are similar because these organisms all can cause polyarthritis and polyserositis.
- *M. hyorhinis* generally occurs in 3- to10-week-old pigs becoming unthrifty, with roughened coat, slight fever, difficult movement, swollen joints and lameness with duration up to 14 days.

Stages of Production

- Nursery

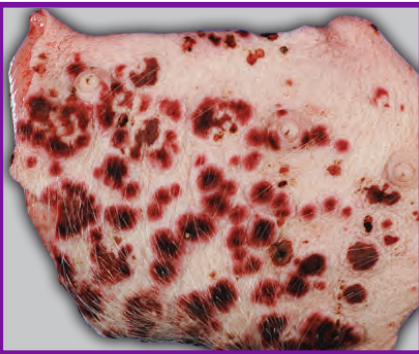
Diagnosis

- Isolation of organism from acute and subacute cases depends on freshly necropsied pigs.
- Polyserositis-affected lungs, pleura, pericardium, epicardium, and peritoneum.

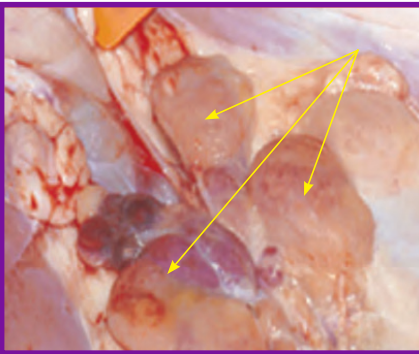
Porcine Circovirus PCV2 / PMWS / PDNS



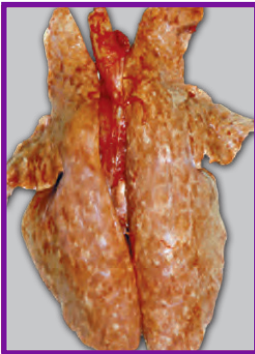
Thin, wasting pig.



Porcine Dermatitis and Nephropathy Syndrome (PDNS) is one manifestation of PCV2 infection. Here the skin shows striking multi-focal hemorrhagic dermatitis from the ventral abdomen.



Enlarged mediastinal lymph nodes.



Lung non-collapsed heavy and firm, or rubbery compatible with diffuse interstitial pneumonia.

PCV2

Tissues to Submit

- Lung
- Spleen
- Lymph Nodes
- Kidney
- Intestine with Peyer's Patches
- Pancreas

Diagnostic Tests

- PCR
- Virus Isolation
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- The most frequent clinical sign is wasting or failure to thrive. In decreasing order of frequency, other signs include dyspnea, enlarged lymph nodes, diarrhea, pallor and jaundice.
- All of the fundamental clinical signs are often not observed in a single pig, but most affected farms will present the majority – if not all – of the signs over a period of time.
- Less common clinical signs include coughing, fever, gastric ulceration, multi-focal hemorrhagic dermatitis and central nervous disorders.

Stages of Production

- Nursery
- Grow-Finish

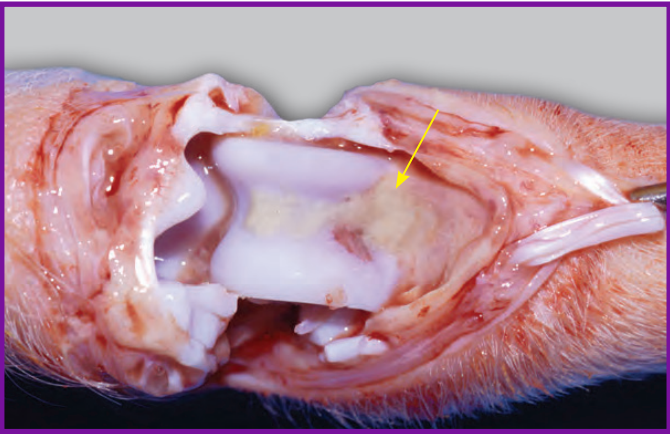
Diagnosis

- Diagnosis of PCV2 requires that a pig or group of pigs has a specific set of clinical signs and microscopic lesions.

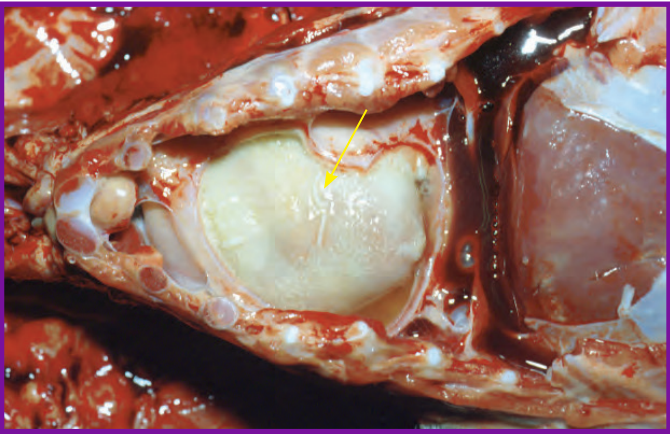
PCV2 Diagnostic Criteria

- Microscopic Lesions: depletion of lymphoid tissues and/or lymphohistiocytic to granulomatous inflammation in any organ (predominantly lung, lymphoid tissue, liver, kidney, intestine, pancreas), or interstitial pneumonia with bronchiolitis.
- PCV2 antigen or genetic material within characteristic lesions;
- Clinical signs alone are not diagnostic. Gross lesions alone are not diagnostic.
- Role of Co-infections: Field observations and scientific literature suggest that PCV2, although essential for development of PCVAD, may require other factors or agents to induce the full spectrum of clinical signs and lesions associated with advanced PCVAD in conventional pigs:
- PRRS + PCV2
- Mycoplasma + IAV-S + PCV2

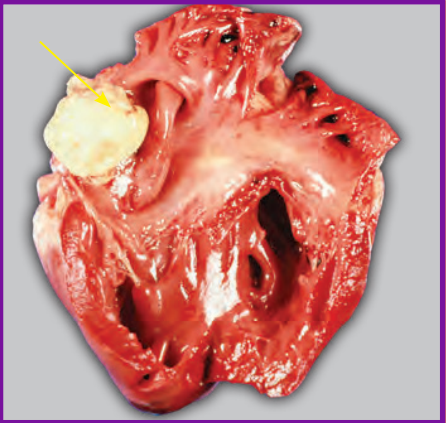
Strep Streptococcus suis



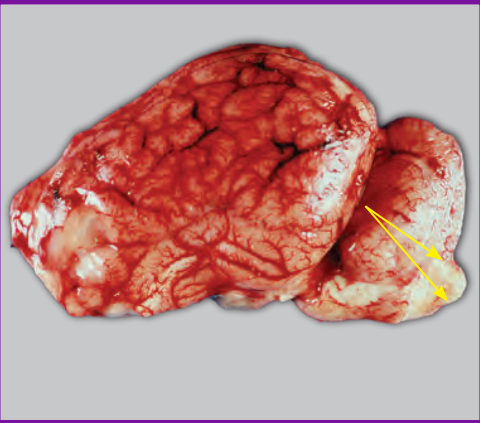
Joint exudate.



Fibrinous lesions on the epicardium (heart surface).



Prominent valvular endocarditis lesion; typical of pigs with bacterial septicemia.



Pus covering cerebellum and brain stem (arrow). Diffuse engorgement of meningeal blood vessels due to hyperemia of septicemia.



Pig down paddling, CNS signs with head tilted back.

STREP

Tissues to Submit

- Brain
- Joint
- Lung
- Liver
- Spleen

Diagnostic Tests

- Culture
- Sensitivity
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- CNS/brain disease; lateral recumbency, paddling, ataxia, head tilt, convulsions, sudden death, arthritis with warm swollen joints, endocarditis (heart).

Stages of Production

- Farrowing
- Nursery

Diagnosis

- Definitive diagnosis depends on gross and microscopic lesions and isolation and identification of the organism. The disease can be confused with other streptococcal infections, other bacterial infections (such as Erysipelas, Salmonellosis or acute Glässer's disease), water deprivation or pseudorabies.
- The skin may be reddened in patches. Lymph nodes are often enlarged and congested, and fibrinopurulent polyserositis is common.
- Joint capsules may be thickened and joints may contain excessive clear or cloudy fluid.
- Affected lungs may show varying degrees of diffuse rubbery interstitial change or patchy consolidation due to bronchopneumonia.

Lepto/Parvovirus/PRRSV



Parvovirus-infected sow litter following abortion. Note mummified fetuses, uneven sizes and post-mortem change indicative of *in utero* death.



One litter from a PRRS virus-associated abortion. Note litter has late-term piglets, at varying stages of *in utero* decomposition, typical of the disease infecting one piglet at a time *in utero*.

REPRODUCTIVE

Tissues to Submit

- Aborted Fetus – two of the freshest
- Mummies – two typical
- Stillborns
- Weakborns
- Sow Serum

Diagnostic Tests

- PCR
- Virus Isolation
- Whole Genome Sequencing
- Histopathology

Clinical Signs and History

- Abortions, mummies, stillborns, weakborns.
- PPV is the most commonly identified cause of reproductive failure, with associated mummification.
- Lepto can cause abortions occurring two to four weeks before farrowing, and is the most common manifestation of leptospirosis in pigs.

Stages of Production

- Gestation
- Farrowing

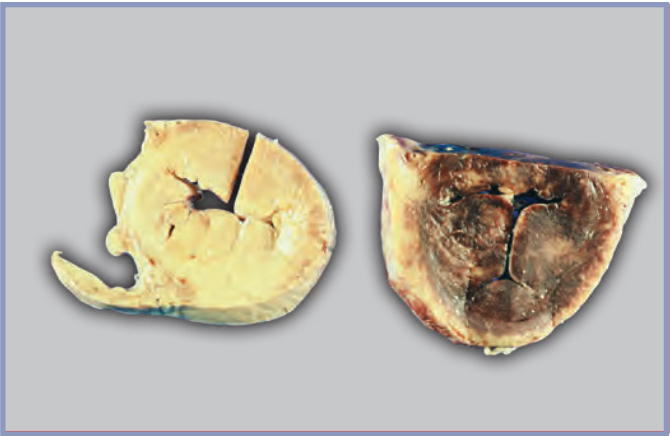
Diagnosis

- Porcine parvovirus (PPV) is usually asymptomatic in adults.
- Sows infected with PPV before 70 days of gestation may abort mummified or near-term autolyzed fetuses.
- PRRS causes late-term abortions including fresh and autolyzed pigs, or weakborn piglets.

Mulberry Heart Disease
Nutritional Cardiomyopathy of Pigs



Fresh heart with multiple prominent hemorrhages on the epicardial surface.



Cross sections of formalin-preserved pig heart. Heart on the left is normal. Heart on the right shows severe diffuse hemorrhage and necrosis of entire left ventricular wall.

MULBERRY HEART

Tissues to Submit

- Fresh Lung and Pericardial Fluid
- Formalin-Fixed Left and Right Ventricles

Diagnostic Tests

- Culture of Lung and Pericardial Fluid to Rule Out Septicemia
- Histopathology Reveals Diagnostic Lesions

Clinical Signs and History

- Sudden death in healthy, rapidly growing piglets and young pigs.
- One or a few pigs in a barn.
- No premonitory signs, but collapse may be precipitated by exercise.

Stages of Production

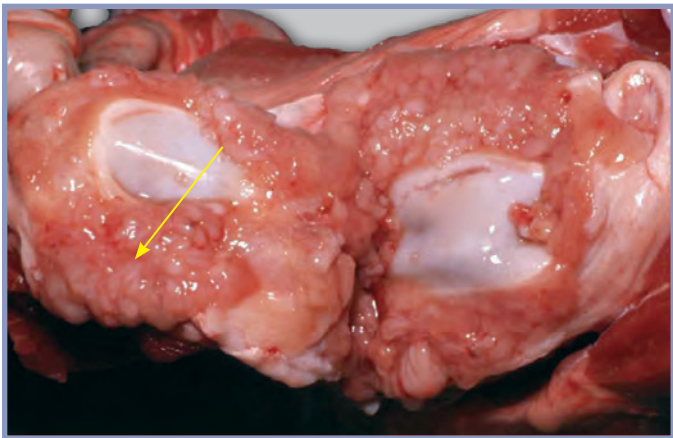
- Farrowing House or Nursery
- 2-16 weeks old

Diagnosis

- Necropsy reveals pericardial effusion and marked epicardial hemorrhages.
- Cross sections of the ventricles show hemorrhages extend throughout the wall.
- Hemorrhages are not superficial on the epicardium, as seen with bacterial septicemias.
- Histopathological heart lesions are pathognomonic. Send formalin-fixed cross section of ventricles for definitive diagnosis.
- A Vitamin E-/selenium-responsive disease.
- Diets may be low in active form of Vitamin E or selenium (Se).
- Factors that may increase Se demand include low concentrations of dietary protein (especially sulfur-containing amino acids), diets with an excess of selenium-antagonistic compounds, and possibly genetic influences on selenium metabolism.
- Vitamin E demand may increase with diets high in polyunsaturated fatty acids, Vitamin A, mycotoxins or rancid fats.

M. hyo Polyarthritits

Mycoplasma hyosynoviae



Proliferative synovitis



Collecting joint fluid.



Dog-sitting pig.

Please reach out to your Vaxxinova vet or representative to help get the best control over endemic diseases.

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ARTHRITIS

Tissues to Submit

- Joints
- Synovium

Diagnostic Tests

- Myco Multiplex PCR
- Culture

Clinical Signs and History

- Lameness typically occurs at 3–5 months of age, appearing acutely and may occur in more than one leg.
- Slight reduction in appetite, resulting in weight loss.

Stages of Production

- Grow-Finish

Diagnosis

- Infected joints are swollen with edema and hyperemia of synovial membranes.
- On necropsy, lesions are restricted to the joints, especially the stifles.
- Joints contain excess of clear, yellow synovial fluid while surrounding tissues are unaffected.
- Definitive diagnosis is made based on isolation of organism.

Notes

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WORTHINGTON OFFICE

1520 Prarie Drive
Worthington, MN 56187
+1 800 220 2522

WILLMAR OFFICE

1801 Biotech Ave NE
Willmar, MN 56201
+1 844 777 8299

INFO.US@VAXXINOVA.COM

WWW.VAXXINOVA.US.COM

