The Microbiology Section of Veterinary Diagnostic Services encompasses diagnostic bacteriology, mycology and parasitology. Culture, microscopy and immunologic methods are used for the detection of pathogenic bacteria, fungi, protozoa, helminths and arthropods. PCR assays for certain bacteria and protozoa are conducted by the Virology and Molecular Diagnostics Section.

Turnaround Times

Estimated turnaround times for all tests are outlined in the VDS Fee Schedule. The time for completion of bacterial culture is the most variable, as it depends on the nature of the specimen and the type of bacteria involved. When specimens are collected from a normally sterile site and contain no bacteria or a single rapidly-growing species, final reports will usually be sent within two working days. When specimens contain slow-growing bacteria (e.g. anaerobes, actinomycetes) or multiple bacterial species (e.g., chronic otitis cases, contaminated samples), it will take longer to report the final result. Preliminary reports will be sent as appropriate.

Specimen Selection and Collection

Directly sample the site of suspected infection as indicated by clinical signs, gross lesions or medical imaging.

When collecting from normally sterile sites, avoid contaminating the specimen with environmental microorganisms or the commensal bacteria of skin (the animal’s or the collector’s), feces or mucosal surfaces.

Antemortem specimens should be placed in appropriate sterile containers. Do not use anticoagulant tubes or serum separator tubes. Swabs should have dacron, rayon or nylon tips and plastic handles. Cotton and wood can inhibit the growth of fastidious bacteria. Swabs with transport media (Amies, Stuart’s) will be composed of appropriate materials, and in most situations, submitting the swab in transport media is preferable. If the swab is intended for culture, do not use transport media intended for virus isolation or PCR testing. “Dry” swabs are only acceptable if they will be processed at VDS within a few hours of collection.

Specimens should be refrigerated and submitted to VDS as soon as possible after collection. If more than two days will elapse between specimen collection and processing at VDS, samples of tissue or exudate should be frozen.

Exceptions: Tritrichomonas pouches and blood culture bottles should remain at room temperature – never refrigerate or freeze these specimens.
Specimen containers must be labeled with the specimen type and at least one unique identifier (animal identification, owner name).
Whenever possible, collect specimens for bacterial culture before starting antimicrobial therapy. Drugs that fail to control an infection may still inhibit in vitro growth. For follow-up testing, it is ideal to cease treatment for 48 to 72 hours before specimen collection. Specimens collected during antimicrobial therapy are still acceptable and may yield significant results; however, negative cultures obtained from these specimens will be questionable.

**Superficial Skin and Ear Canal**

*Otitis Externa and Pyoderma:* Gram staining will be done on swabs submitted for bacterial culture. Yeast such as *Malassezia* and *Candida* will be seen on the Gram stain and may grow on bacterial media if concurrent bacterial infection is absent. Requesting fungal culture is generally not warranted in these cases. If fungal culture is desired, submit a separate swab. Ears should be swabbed separately. Using the same swab for both ears is acceptable but not recommended. The Microbiology lab will not pool multiple swabs for a single culture.

*Dermatophytosis:* Collect at least 2 cubic centimetres of hair and scab material from the periphery of lesions and submit for fungal culture. If enough sample is received, it will be examined by the KOH-calcofluor white method which has the potential to detect fungal elements rapidly. Dermatophyte growth usually takes at least 5 days; negative culture will be reported after 3 weeks.

*Acariasis:* If mange is suspected, submit skin scrapings for a parasitology direct examination. Mites such as *Sarcoptes* and *Demodex* are often difficult to detect, so multiple skin scrapings are recommended.

*Macroparasites:* Submit lice, fleas and ticks for parasite identification. Ticks and unusual arthropods will be referred to the University of Manitoba Department of Entomology for identification. Black-legged ticks (*Ixodes scapularis*) will be forwarded to the National Microbiology Laboratory for PCR assays that detect *Borrelia burgdorferi*, *Anaplasma phagocytophilum* and *Babesia microti*.

**Wounds, Cellulitis, Draining Tracts and Abscesses**

Gram stain, aerobic culture and anaerobic culture are done on specimens from lesions of this type. Swabs are acceptable, but samples of exudate (1 mL or more) or fresh tissue biopsies are better, especially for anaerobic culture. Sample from the periphery of abscesses, as the central exudate may not contain viable bacteria.

*Blastomycosis:* Smears from appropriate dog and cat specimens will be examined for the presence of *Blastomyces dermatitidis* (no fee is applied for this test). If blastomycosis is strongly suspected by the clinician, smears should be sent to the Clinical Pathology Section for cytology.
Granulomas: If lesions are suspected to be granulomas submit fresh specimens for bacterial culture, fungal culture and acid-fast staining; samples may be sent to a referral laboratory for mycobacterial culture. Fixed samples for histopathology are also recommended.

Histotoxic Clostridial Infections

In cases of suspected blackleg or malignant edema/gas gangrene in ruminants, submit samples of subcutaneous tissue and/or skeletal muscle for the clostridial fluorescent antibody test (FAT) panel. The panel includes Clostridium chauvoei, Clostridium septicum, Clostridium novyi and Clostridium sordellii. A positive result for C. chauvoei almost always indicates blackleg. The other species rapidly colonize tissues after death; if autolysis is advanced, consider testing for C. chauvoei only.

Heart and other internal organs can be appropriate specimens for the Clostridium chauvoei FAT if necropsy reveals myocardial necrosis or serofibrinous exudate in the pericardial sac, thoracic cavity or abdominal cavity. The full clostridial panel is not appropriate for internal organs.

If bacillary hemoglobinuria or black disease (infectious necrotic hepatitis) are suspected, submit a portion of necrotic liver for the Clostridium novyi FAT. This test will detect both C. novyi type B (the agent of black disease) and C. haemolyticum (the agent of bacillary hemoglobinuria, formerly C. novyi type D).

Conjunctivitis

Moisten the swab with sterile saline and avoid contact with the eyelid skin. If corneal ulceration is present, particularly in horses, collect a separate swab for fungal culture. In cattle, submit a separate swab in viral transport media for Bovine herpesvirus 1 PCR testing.

Urine

Urine collected by cystocentesis is the ideal specimen for quantitative bacterial culture. Free catch (midstream voided) samples and catheterization samples are acceptable for cats and male dogs, as well as large animals.

During cystotomy, urine or visibly inflamed bladder wall can be collected for culture. While urolithiasis in dogs is commonly associated with urease-producing bacteria, bladder stones are not useful for culture.

When requesting both urine culture and urinalysis, submit urine in two separate containers for the Microbiology and Clinical Pathology Sections.
Milk

**Bovine Mastitis:** Individual quarter samples are ideal and should always be collected in clinical mastitis cases. Composite samples (milk from 2-4 quarters) can be used to screen herds for major mastitis pathogens.

The California Mastitis Test (CMT) will be done on quarter samples that have not been frozen. If the CMT indicates mastitis, but no pathogens are detected by routine culture, mycoplasma culture will be set up (the PCR test for *Mycoplasma bovis* cannot be done on milk samples).

Udder abscesses or other lesions should be noted in the history, as should any signs of systemic illness.

Milk samples are often collected by dairy producers. Veterinarians are responsible for instructing their clients in proper collection techniques that will avoid contamination. Many of the minor mastitis pathogens (e.g., coagulase-negative staphylococci) are normal residents on skin and therefore are common contaminants. Environmental pathogens and *Staphylococcus aureus* can be present in feces and reside transiently on skin, so severely contaminated milk samples can give misleading culture results. When contamination is severe (3 or more species isolated), only major mastitis pathogens will be reported.

**Other Species:** Milk specimens should be collected from individual mammary glands, not pooled. Mammary gland abscesses should be noted in the history. Consider aspirating material directly from abscesses.

**Respiratory Tract**

**Rhinitis:** Nasal swabs or nasal flushes can be submitted for bacterial culture. Submit separate swabs for fungal culture.

**Tracheobronchitis and Pneumonia:** Transtracheal aspirates and bronchoalveolar lavages are the ideal samples. Nasal swabs are acceptable but will often be contaminated by nasopharyngeal flora. Note that many of the bacteria involved in causing pneumonia can also be nonpathogenic residents of the upper respiratory tract. Lung samples collected at necropsy should be taken from areas with visible pneumonia or pleuritis. If sampling lung that lacks definite lesions, collect from the cranial and middle lung lobes.

**Blood**

When bacteremia is suspected, blood collected in a red-top or anticoagulant tube is not a suitable specimen for culture. Special blood culture bottles are required. These bottles can be obtained from VDS at no charge.
Joint, Pleural and Peritoneal Fluids

A fluid sample (at least 0.5 mL) is preferable to a swab. Fluid submitted in an EDTA tube for fluid analysis (see Clinical Pathology Section) cannot be used for bacterial culture – submit a separate sample in a red top tube.

Feces

Submit fecal samples in rigid, sealable plastic containers. Tied-off disposable gloves are not a suitable specimen container.

Fecal Bacteriology

Culture on feces and fecal swabs will routinely include Salmonella culture and, in susceptible species, Campylobacter culture.

Escherichia coli is almost universally present in feces – it will only be reported if there is reason to suspect E. coli diarrhea. E. coli isolated from calves with onset of diarrhea during the first 3 days of life will be reported and tested for the F5 (K99) fimbrial antigen – F5 positive isolates should be considered significant. Hemolytic E. coli isolates from swine in susceptible age groups will be reported and tested for the F4 (K88) antigen. E. coli may be reported in other scenarios if there is histologic evidence of infection. PCR testing for E. coli virulence factor genes is also available (see the Virology and Molecular Diagnostics Section).

Clostridium perfringens is part of the large intestinal flora, so isolating it from a fecal sample is a poor indicator of infection. In dogs and cats, the ELISA test for the C. perfringens enterotoxin is used to test for clostridial diarrhea. For diagnosis of C. perfringens type A diarrhea in neonatal piglets, anaerobic culture should be done on small intestinal samples collected shortly after euthanasia. Isolates will be sent for PCR typing and to detect the beta-2 toxin gene that is associated with piglet diarrhea. When C. perfringens type D enterotoxemia (lambs) or C. perfringens type C necrotizing enteritis (multiple species) is suspected, anaerobic culture should be done on small intestine as soon as possible after death, and any C. perfringens isolates should be sent for PCR typing. For necrotic enteritis of poultry, affected segments of intestine should be submitted for aerobic and anaerobic culture.

A Clostridium difficile toxin ELISA is available for piglet fecal or colon samples and canine fecal samples.

Fecal Parasitology

A direct smear examination can be done when only a small volume is available. This method may be more sensitive for certain trematodes and protozoa. Fecal flotation is
generally preferred for detecting cestodes, nematodes and coccidia. Flotation requires about 10 mL of liquid feces or a “walnut-sized” portion of solid feces.

A fluorescent antibody test panel is done to detect Cryptosporidium and Giardia in fecal samples. This test requires about 3 grams of feces. If this amount is not available, PCR testing for Cryptosporidium can be done in the Virology and Molecular Diagnostics Section.

Macroscopic worms recently passed by the animal or found during necropsy can be submitted for parasite identification.

Postmortem Specimens

After death, body sites that normally contain few or no microorganisms are rapidly colonized by endogenous bacteria originating mainly from the gastrointestinal tract and upper respiratory tract. Even mild autolysis can result in bacterial culture results that are unrewarding or difficult to interpret. Specimens should be collected as soon as possible after death.

During necropsy, avoid touching the oral cavity and gastrointestinal tract until culture samples have been collected. Alternatively, change gloves before collecting bacteriology specimens. Organ samples should be placed in separate labeled bags. Different organs from the same animal should not be pooled – mixing thoracic and abdominal organs is especially detrimental. It is also not acceptable to pool the same organ from different animals. When submitting multiple organ samples from multiple animals, the label on each bag should include the animal identification as well as the specimen identification (e.g., Calf A lung, Calf B lung or Turkey 1 liver, Turkey 2 liver).

Antimicrobial Sensitivity Testing (AST)

VDS uses the Kirby-Bauer disk diffusion method for AST. Zones of bacterial growth inhibition around antimicrobial drug-impregnated disks are measured and interpreted according to criteria set by the Clinical and Laboratory Standards Institute (CLSI). This method works well for fast-growing aerobic bacteria but cannot be done on anaerobes or certain fastidious bacteria. When disk diffusion is not appropriate, minimum inhibitory concentration determination by microbroth dilution may be available at a referral laboratory.