

CANADIAN BAT WHITE-NOSE SYNDROME NECROPSY PROTOCOL
Canadian Cooperative Wildlife Health Centre
November 1, 2013

PREAMBLE

The purpose of this necropsy protocol is to provide guidance to researchers, laboratories and diagnosticians participating in surveillance for the disease bat white-nose syndrome (WNS). To contribute to Canada's national surveillance program for WNS, **occurrence data must be reported using the diagnostic categories given at the end of this protocol and the criteria for each category of diagnosis must be strictly applied.** This is to ensure uniformity, consistency, accuracy and comparability of results. Also, Canada's national survey for WNS designates each annual surveillance period as the interval from **November 1 of a given year to May 31 of the following year.** During the defined bat WNS surveillance period this detailed necropsy protocol should be followed. This specifically means that all bats submitted will receive a complete necropsy combined with both a histological examination of wing membranes and RT-PCR for the fungus *Pseudogymnoascus* (formerly *Geomyces destructans*). **Best laboratory practices should be followed** to ensure quality assurance and control. As a **biosecurity guideline** for regions where bat WNS is not endemic, it is recommended that bat necropsies and other diagnostic tests involving materials potentially contaminated with *P. destructans* are done in a **biosafety cabinet.** Outside of the surveillance period, the diagnostic tests performed for bat WNS would be optional, and the biosecurity measures to prevent cross contamination of specimens and laboratory contamination could be relaxed.

GROSS EXAMINATION AND SWABBING FOR RT-PCR:

1. Weigh and sex the specimen. (**Note: If examining bats for bat WNS, only handle one bat at a time to prevent cross contamination. DO NOT weigh the bats until the time of necropsy and swab the specimen for RT-PCR for *Pseudogymnosascus destructans* (Pd) at the time of weighing. Weigh individual bats in a disposable weigh boat [discarded after a single use]. After completion of necropsy, change gloves and decontaminate instruments and working surfaces [see below] between each bat to prevent cross contamination of specimens. If multiple bats are submitted together in one bag, pool the sample by swabbing all of the bats with a single swab and the bats can be necropsied as a group without changing gloves or decontaminating instruments).**)
2. RT-PCR for Pd. With a single **polyester** swab, swab the muzzle, ears and dorsal and ventral surfaces of all wing membranes. Place individual swabs in separate labeled sterile whirlpak bags and keep chilled or frozen. **Change gloves between each bat.** Samples for RT-PCR can be shipped as diagnostic specimens to Dr. Hugh Cai, Animal Health Laboratory, University of Guelph (hcai@uoguelph.ca; 519-824-4120 ext 54316) or to the Animal Health Centre, Abbotsford, BC (<http://www.agf.gov.bc.ca/ahc/>; Toll Free 1-800-661-9903 [BC Only]; 604-556-3003). Please contact these laboratories prior to sending specimens. 2013-14 Bat WNS Surveillance Season costs per Pd RT-

PCR are \$34.00 per sample at the Animal Health Laboratory, University of Guelph and \$25.00 + GST per sample (BC **in province** submissions) or \$37.50 + GST per sample (BC **out of province** submissions) at the Animal Health Centre, Abbotsford, BC.

If the preference is to do your own RT-PCR, the current RT-PCR protocol is based on:

Muller, LK, JM Lorch, DL Lindner, M O'Connor, A Gargas and DS Blehert. 2013. Bat white-nose syndrome: a real-time TaqMan polymerase chain reaction test targeting the intergenic spacer region of *Geomyces destructans*. *Mycologia* 105:253-259.

3. Identify bat specimens to species. The indigenous bat species vary depending on the geographic location in Canada. Therefore, diagnosticians must become familiar with the bat species in their particular area. Important traits for differentiating among bat species are external morphological measurements including total body, tail, forearm, foot, tragus and ear lengths (**Note: ear and tragus lengths are difficult to measure consistently, resulting in high measuring error**); fur color; and presence or absence of a keel on the calcar. Most characteristics used in bat identification keys are those of adult bats and do not necessarily apply to nursing or immature individuals. Photographing a specimen and consulting a bat biologist or museum curator (eg. Dr. Hugh Broders, Saint Mary's University, Halifax, Nova Scotia; Dr. Don McAlpine, New Brunswick Museum, Saint John, New Brunswick; Dr. Brock Fenton; University of Western Ontario, London, Ontario; Dr. Craig Willis, University of Winnipeg, Winnipeg, Manitoba; and Dr. Cori Lausen, Bats R Us, Birchdale Ecological, British Columbia) may be necessary to obtain an accurate species identification. Nagorsen (2002) gives the following examples of species that are difficult to separate:
 - a. Keen's Long-eared Myotis (*Myotis keenii*) and Western Long-eared Myotis (*M. evotis*) cannot be reliably identified from external traits; their identification requires a cleaned skull.
 - b. Separation of other species of bats such as the Yuma Myotis (*M. yumanensis*) vs. Little Brown Bat (*M. lucifugus*) is very difficult to accomplish with a 100 percent certainty from their external features alone. Therefore, positive identification requires examination of a voucher specimen by a competent bat taxonomist.

The three species of bats currently most affected by bat white-nose syndrome (WNS) in Canada are the Little Brown Bat (LBB), Northern Long-eared Bat (*M. septentrionalis* or NLE) and Tricolored Bat (*Perimyotis subflavus* or TCB and previously known as the Eastern Pipistrelle). In general, the TCB is smaller than the LBB and NLE, but the overlapping weight and forearm length ranges for these three species make accurate species identification challenging.

The morphology of the tragus is a distinguishing feature for these species (see images below). However, it can be necessary to use a dissecting

scope to carefully and closely examine the tragus in post mortem specimens because dehydration of the individual and/or desiccation of the specimen can affect the size and shape of the tragus.

The LBB tragus is wide with a straight medial surface and bump on lateral edge (note: the bump is often curled inward on partially desiccated specimens so the tragus might have to be manipulated to be accurately visualized); the NLE tragus is conical or triangular in shape with straight lateral surface and slightly curved medial surface; and the TCB tragus is short, blunt and rounded. Placing a piece of white paper behind the tragus helps with visualization of this structure. In addition, the TCB can be recognized by its orange-red forearm and tricolored fur (black base, light brown center and dark tips).



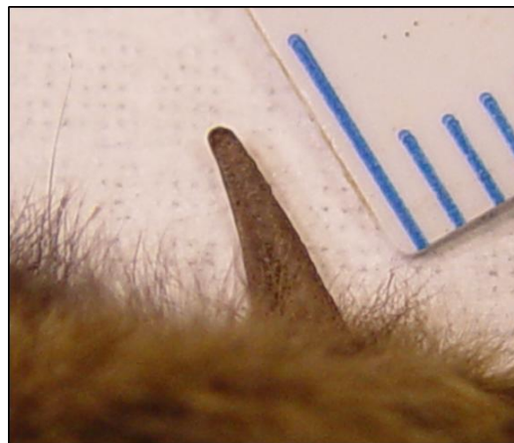
Little Brown Bat
Broders, Saint Mary's University



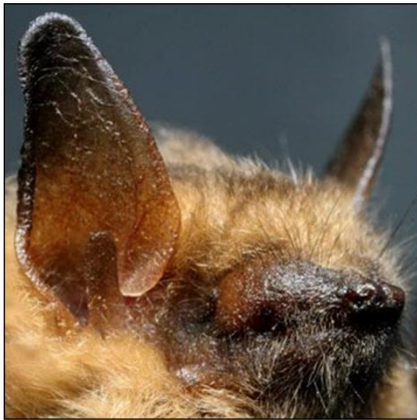
Little Brown Bat Tragus
McBurney and Needham, CCWHC, Atlantic



Northern Long-eared Bat
Broders, Saint Mary's University



Northern Long-eared Bat Tragus
McBurney, CCWHC, Atlantic



Tricolored Bat
© Hilton Pond Center



Tricolored Bat Tragus
McBurney, CCWHC, Atlantic



Tricolored bat (red-orange forearm)
© K. Miles, Tennessee Wildlife Resources Agency



Tricolored bat (black-brown-dark fur)
McBurney, CCWHC, Atlantic

There are many references online to help with identification of bat species, and the following are a few examples:

- a. Nagorsen, D.W. 2002. An identification manual to the small mammals of British Columbia. Ministry of Sustainable Resource Management, Ministry of Water, Land and Air Protection, Biodiversity Branch, and Royal British Columbia Museum. 165pp. Found at:
http://www.env.gov.bc.ca/wld/documents/techpub/id_keys_s.pdf
- b. http://www.ontarionature.org/discover/resources/PDFs/atlas/mammal_atlas_bats.pdf
- c. http://www.dnr.state.md.us/wildlife/Plants_Wildlife/bats/nhpbatfield.asp
- d. http://www.dnr.state.md.us/wildlife/Plants_Wildlife/bats/bat_key.asp - Dichotomous Key

Lastly, “van Zyll de Jong, C.G. 1985. Handbook of Canadian mammals. Volume 2 (Bats). National Museum of Canada and National Museum of Natural History, Ottawa, Ontario. 210pp.” is a good reference to have if you are working with bats in Canada.

5. Examination of Wings. In fresh specimens, examining the spread wings on a light table, backlit by a bright light or using UV light may reveal erosions in the skin of the wing membrane. The lesions may also be photographed at this time. Looking at the wings under a dissecting microscope is also helpful because mites and small skin lesions can be detected using this equipment.
6. Examination of Other Organ Systems. Grossly examine other body organ systems and collect representative samples of organs and any lesions observed for histological examination and/or other ancillary diagnostic tests (eg. bacteriology, virology and parasitology).
7. DNA Sampling. If DNA is needed for genetic studies, we typically collect the majority of the left uropatagium (see wing membrane diagram below to identify this structure) and the entire left ear, including the tragus. These are placed in a DMSO Sodium Solution. However, it is best to use a protocol specified by the individuals who are doing the genetic study.

NOTE: To prevent cross contamination of samples, dissection instruments and working surfaces should be cleaned and decontaminated after each necropsy and before proceeding with the next specimen's necropsy. However, flaming stainless steel dissection instruments or soaking them in 10% bleach solutions can damage them excessively. Therefore, this alternative protocol is provided to avoid flaming and minimize contact with 10% bleach solution:

Cleaning Tools for Processing Bat Tissues for PCR and Culture Analyses

(modified from J. Lorch, E. Bohuski, and D. Blehert, USGS – National Wildlife Health Center, June 2013)

1. Remove dissection tools from storage container, and before use, briefly dip tools in 10% bleach and rinse almost immediately by transferring to a tray containing deionized or distilled water.
2. Transfer rinse tray containing tools to the working surface, remove tools from tray, and place on a clean paper towel.
3. Spray tools on paper towel with a DNA decontamination solution (e.g., DNA-OFF™, Clontech Laboratories, Fisher Scientific Catalogue Number TAK9036; Decon™ ELIMINase™, Decon Laboratories, Fisher Scientific Catalogue Number 04-355-32; or D/RNase Free™ Decontaminant, Argos Technologies, VWR Catalogue Number 47751-044) and allow tools to sit for approximately 10 minutes or until dry.
4. Rinse tools in a tray containing deionized or distilled water.
5. Remove tools from rinse tray, dry with fresh paper towels and proceed with necropsy.

6. Following necropsy, clean tools with hot soapy water and dry them with a clean paper towel. Spray tools with DNA-Off solution and allow tools to sit for approximately 10 minutes or until dry. Lastly, dip them in 10% bleach and rinse immediately by transferring to a tray containing deionized or distilled water before returning tools to the storage container.
7. Douse working surface with 95% EtOH and allow the solution to evaporate. Then wet the working surface with 10% bleach and let stand 2 minutes. Wipe up the bleach solution and rinse working surfaces with deionized or distilled water to remove any residual bleach.

TISSUE SAMPLING:

1. Histology. (**Note: Place all skin samples in one cassette and immediately request a PAS stain at time of submission for histology if bat WNS is the suspect diagnosis.**)
 - a. Wing membranes: The wing membrane is thin and elastic. The NWHC's proposed method for collecting histology samples from this tissue is as follows: *Collect a large rectangle of wing membrane for histology which includes lesions if present. Roll the skin sample like a cinnamon roll or roll it onto a dowel and place in cassette with a piece of paper towel on it so the tissue doesn't unroll. Immerse the cassette in 10% neutral buffered formalin for fixation (24 hours). After fixation, cut multiple cross sections of the roll for histological processing.*

In practice, the technique above is difficult to do (especially in partially desiccated or decomposed carcasses), but a sample of wing membrane rolls into a ball quite readily. If you fix and cross section the ball of tissue, a very large surface area is available to examine microscopically. Therefore, collect the entire right and left dactylopatagia major, right and left plagiopatagia and right uropatagium, individually compress the membranes into small balls and place them into the cassette for processing (see diagram below for wing membrane nomenclature). Propatagia and dactylopatagia medius, minus and brevis can also be collected if lesions are observed in them. This methodology is best for simple diagnostic evaluation. However, if histological scoring/grading of wing damage is the goal, a more rigorous methodology is required and is described in Reeder *et al.*, 2012, *PLoS ONE* 7:e38920, Appendix S2.

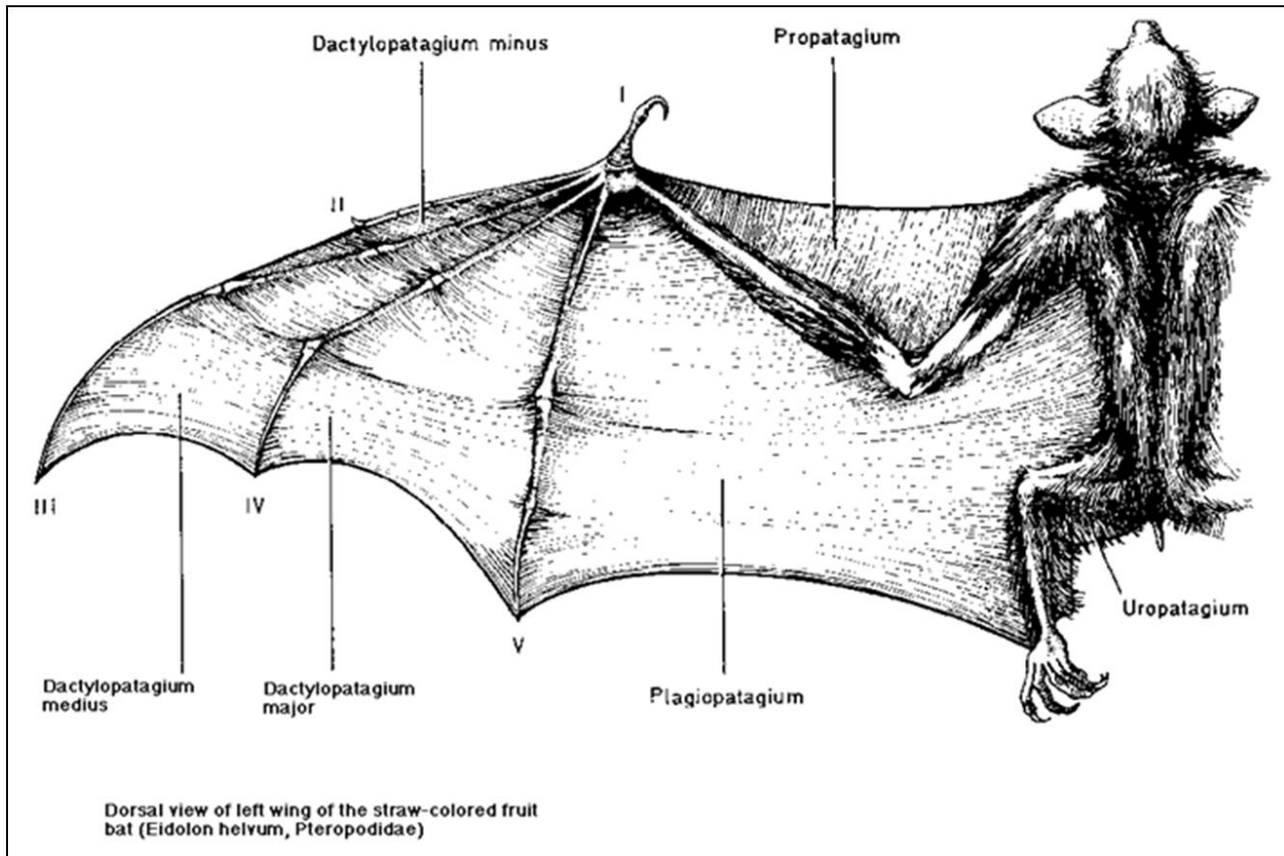


Diagram from: Halstead, L.B., and A.O. Segun, 1975. Dissection guides of common tropical animals, 3 Fruit bat (*Eidolon helvum*). Publication No. 6. Ethiopie Publishing House, Benin City. (**Note: *Uropatagium of Myotis and Perimyotis spp. forms a wide membrane that connects the leg to the tail***)

- b. Skin of the head and face: Take two transverse sections across the muzzle using a scalpel. Histology technicians can decalcify the surface of the block at the time of sectioning. Another option is to remove the skin from the nose and submandibular region for sectioning, but this is more time consuming. Include these sections in the same cassette containing wing membrane sections.
 - c. Other tissue samples for histology: One half of the brain, one entire lung, heart (entire), liver, spleen and stomach (together), intestine, kidney (often with adrenal) and other tissues with lesions as identified. Place in a second cassette and immerse it in 10% neutral buffered formalin for fixation (24 hours) and standard histological processing.
2. Screening for Rabies at CCWHC, Atlantic Region: All bats are screened for rabies using the Direct Rapid Immunohistochemical Test (dRIT). Half of the fresh brain is given to a wildlife technician for dRIT. After making the dRIT smears, the wildlife technician archives the brain tissue @ -80C for CFIA confirmation by FAT if dRIT positive.

3. Frozen Samples. (**-80 C is best**) The remainder of the bat carcass and tissues not utilized for diagnostic purposes can be placed in a labeled whirlpak bag and frozen for further diagnostic work or other research initiatives as deemed necessary.

DIAGNOSTIC CATEGORIES FOR CLASSIFYING AND REPORTING BAT WHITE-NOSE SYNDROME CASES

1. **Confirmed Positive** – Gross and histological lesions of Bat White-nose Syndrome (BWNS) are present **AND** RT-PCR for *Pseudogymnoascus destructans* (Pd) is positive.
2. **Presumptive Positive** – Gross lesions of BWNS are present. Histology and RT-PCR for Pd testing are not done, but the specimen is submitted from a site with confirmed positive cases in the same surveillance season. This category is for instances in which > 5 bats are submitted from a site experiencing ongoing bat WNS mortality, and it is not affordable to test all of the submitted specimens.
3. **Suspect Positive** – Gross and histological lesions of BWNS are present but RT-PCR for Pd is negative **OR** RT-PCR for Pd is positive but gross and histological lesions are not present.
4. **Negative** – Gross and histological lesions are not present **AND** RT-PCR for Pd is negative.

Protocol Revised November 1, 2013

Questions regarding this protocol should be directed to:

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