Molecular Detection of Pseudogymnoascus destructans (Ascomycota: Pseudeurotiaceae) and Unidentified Fungal Dermatitides on Big Brown Bats (Eptesicus fuscus) Overwintering inside Buildings in Canada

Authors: Donald F. McAlpine, Scott McBurney, Mary Sabine, Karen J. Vanderwolf, Allysia Park, et. al.

Source: Journal of Wildlife Diseases, 52(4) : 902-906

Published By: Wildlife Disease Association

URL: https://doi.org/10.7589/2015-03-076
Molecular Detection of *Pseudogymnoascus destructans* (Ascomycota: Pseudeurotiaceae) and Unidentified Fungal Dermatitides on Big Brown Bats (*Eptesicus fuscus*) Overwintering inside Buildings in Canada

Donald F. McAlpine,1,6 Scott McBurney,2 Mary Sabine,3 Karen J. Vanderwolf,1,4 Allysia Park,2 and Hugh Y. Cai1
1New Brunswick Museum, 277 Douglas Avenue, Saint John, New Brunswick E2K 1E5, Canada; 2Canadian Wildlife Health Cooperative, Atlantic Region, Atlantic Veterinary College, University of Prince Edward Island, 550 University Avenue, Charlottetown, Prince Edward Island C1A 4P3, Canada; 3New Brunswick Department of Natural Resources, PO Box 6000, Fredericton, New Brunswick E3B 5H1, Canada; 4Canadian Wildlife Federation, 350 Promenade Michael Cowpland Drive, Kanata, Ontario K2M 2W1, Canada; 5Animal Health Laboratory, University of Guelph, 419 Gordon Street, Building 89, Guelph, Ontario N1G 2W1, Canada; 6Corresponding author (email: donald.mcalpine@nbm-mnb.ca)

**ABSTRACT:** Big brown bats (*Eptesicus fuscus*) overwintering outside the underground environment are not believed to play a role in the epidemiology of the disease white-nose syndrome (WNS), caused by the fungus *Pseudogymnoascus destructans* (*Pd*). Using quantitative real-time PCR (qPCR), we provide molecular evidence for *Pd* on four big brown bats overwintering in heated buildings in New Brunswick, Canada. Two of the affected individuals also had very mild, focal, pustular, fungal dermatitis identified microscopically. A third bat, which was qPCR *Pd*-negative, had similar fungal lesions. Despite determining that these fungal lesions were caused by a suspected ascomycete, the intralesional fungi were not confirmed to be *Pd*. These findings demonstrate that bats overwintering in heated buildings and other above-ground sites may have subclinical or preclinical WNS, or be contaminated with *Pd*, and could play a role in local dispersal of *Pd*. Our inability to determine if the ascomycetes causing pustular lesions were *Pd* highlights the need for ancillary diagnostic tests, such as in situ hybridization or immunohistochemistry, so that *Pd* can be detected directly within a lesion. As the host–pathogen relationship for *Pd* evolves, and where bat species are exposed to the fungus under varying temperature regimes, lesions may become less stereotypic and such tests could help define these changes.

**Key words:** Big brown bat, *Eptesicus fuscus*, fungal infection, *Pseudogymnoascus destructans*, white-nose syndrome.

White-nose syndrome (WNS), caused by the fungus *Pseudogymnoascus destructans* (*Pd*), is decimating populations of eastern North American bats. First detected in New York in 2006–07, it is believed *Pd* was introduced from Europe (Leopardi et al. 2015). Estimates suggest >6.7 million bats have died from WNS (US Fish and Wildlife Service 2012), and several species may be at risk of regional extinction (Frick et al. 2010). White-nose syndrome is associated with bats in underground hibernacula during periods of extended torpor (hibernation), when they fail to show a cellular immune response to *Pd* invasion (Meteyer et al. 2011, 2012) and where environmental conditions provide high humidity and the range of temperatures at which *Pd* grows (Verant et al. 2012).

More than 50% of the 45 North American bat species hibernate underground (Foley et al. 2011), most above 40°N. Other bats may enter torpor for short or extended periods while roosting in buildings, hollow trees, and under bark. As winter approaches some migratory bats leave the northern portion of their range, possibly avoiding the need for torpor. Although WNS is reported in cave-roosting big brown bats (*Eptesicus fuscus*) and *Pd* prevalence on this species can be high (Langwig et al. 2014), we are not aware that *Pd* or WNS has been documented in hibernating bats overwintering outside underground hibernacula. We present incidents where *E. fuscus* overwintering in buildings had Suspect (b) category diagnoses of WNS (Canadian Wildlife Health Cooperative [CWHC] 2014), based on premature egression from hibernacula (a clinical or field sign associated with WNS) and cryptic *Pd* infections as determined by quantitative real-time PCR (qPCR) for *Pd*, but with no external...
Signs of the fungus (see Janicki et al. 2015). The affected individuals also lacked grossly visible or microscopic lesions consistent with WNS (Meteyer et al. 2009), providing further evidence of the cryptic nature of these preclinical or subclinical infections. Although \( Pd \) on the surface of bat skin or fur could yield positive qPCR without infection, molecular detection of \( Pd \) on multiple bats during hibernation, 195 km from \( Pd \)-positive underground hibernacula strongly suggests infection. In addition, some affected bats had a fungal dermatitis caused by unidentified ascomycetes. We discuss epidemiologic implications and diagnostic significance of these results.

Surveillance for WNS and \( Pd \) in bats in Atlantic Canada follows a standardized necropsy protocol (CWHC 2014) and has been underway since winter 2008–09. The disease was first recorded in New Brunswick in 2011. Most postmortem submissions of bats include little brown bats (\( Myotis lucifugus \)) and northern long-eared bats (\( Myotis septentrionalis \)) which overwinter in underground hibernacula in the region. However, 10 big brown bats were submitted from New Brunswick during mid-November to late May 2013–14 (Table 1). Six of these bats were discovered active and alive inside the living spaces of human-occupied, heated buildings in Fredericton (45°57′N–66°38′W). The remaining four were found dead or alive outside, near buildings, where they may have been overwintering, in Fredericton (\( n=3 \)) and near Royalton (46°28′N–67°45′W), 85–190 km north of the closest known underground hibernaculum. Prior to necropsy, swabs for \( Pd \) were taken with one sterile, polyester swab from the dorsal and ventral wing membranes, ears, and muzzle of each bat (CWHC 2014). Swabs were frozen at \(-80^\circ C\) and sent to the Animal Health Laboratory, University of Guelph (Ontario) to test for \( Pd \) using qPCR for WNS surveillance (Muller et al. 2013). Skin from all wing membranes, ears, and two cross-sections of the muzzle were fixed in 10% neutral-buffered formalin, dehydrated in alcohol and xylene, and embedded in paraffin; 5-μm sections were stained with H&E and periodic acid–Schiff to detect fungi.

### Table 1. Findings in big brown bats (\( Eptesicus fuscus \)) collected in or adjacent to buildings, during winter in New Brunswick, Canada, and submitted to Canadian Wildlife Health Cooperative (CWHC) November 2013–May 2014. Included are quantitative real-time PCR (qPCR) results for \( Pseudogymnoascus destructans \) (\( Pd \)), white-nose syndrome status based on histologic evaluation, whether fungal elements were seen histologically, and minimum outdoor temperature on day of collection.

<table>
<thead>
<tr>
<th>Date collected</th>
<th>CWHC</th>
<th>Location</th>
<th>Status</th>
<th>Sex</th>
<th>Age</th>
<th>Weight (g)</th>
<th>( Pd ) qPCR</th>
<th>WNS</th>
<th>Fungal lesions</th>
<th>Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>16 November</td>
<td>107455</td>
<td>O</td>
<td>Alive</td>
<td>M</td>
<td>Imm</td>
<td>11.6</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>-5.6 C</td>
</tr>
<tr>
<td>2 January</td>
<td>107775</td>
<td>B</td>
<td>Alive</td>
<td>M</td>
<td>?</td>
<td>12.8</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>-29.5 C</td>
</tr>
<tr>
<td>2 January</td>
<td>107768</td>
<td>O</td>
<td>Dead</td>
<td>M</td>
<td>Imm</td>
<td>14.5</td>
<td>P (Cq=29.46)</td>
<td>N</td>
<td>N</td>
<td>-31.0 C</td>
</tr>
<tr>
<td>8 January</td>
<td>107769</td>
<td>B</td>
<td>Alive</td>
<td>F</td>
<td>Adult</td>
<td>19.4</td>
<td>P (Cq=29.07)</td>
<td>N</td>
<td>P</td>
<td>-16.1 C</td>
</tr>
<tr>
<td>9 January</td>
<td>107774</td>
<td>B</td>
<td>Alive</td>
<td>F</td>
<td>Imm</td>
<td>11.7</td>
<td>P (Cq=30.51)</td>
<td>N</td>
<td>P</td>
<td>-19.5 C</td>
</tr>
<tr>
<td>26 January</td>
<td>110398</td>
<td>B</td>
<td>Alive</td>
<td>F</td>
<td>Imm</td>
<td>10.2</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>-15.6 C</td>
</tr>
<tr>
<td>7 February</td>
<td>108006</td>
<td>B</td>
<td>Alive</td>
<td>F</td>
<td>Adult</td>
<td>13.0</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>-18.7 C</td>
</tr>
<tr>
<td>25 March</td>
<td>109752</td>
<td>B</td>
<td>Alive</td>
<td>M</td>
<td>Adult</td>
<td>11.9</td>
<td>P (Cq=35.00)</td>
<td>N</td>
<td>N</td>
<td>-21.6 C</td>
</tr>
<tr>
<td>16 April</td>
<td>110401</td>
<td>B</td>
<td>Alive</td>
<td>F</td>
<td>Imm</td>
<td>12.0</td>
<td>N</td>
<td>N</td>
<td>[10.3 C]f</td>
<td></td>
</tr>
<tr>
<td>23 May</td>
<td>110014</td>
<td>O</td>
<td>Alive</td>
<td>F</td>
<td>Imm</td>
<td>11.7</td>
<td>N (Cq=44.19)</td>
<td>N</td>
<td>P</td>
<td>8.0 C</td>
</tr>
</tbody>
</table>

*O* = outside; *B* = inside building.

* M = male; *F* = female.

* Imm = immature.

* N = negative; *P* = positive; *Cq* = quantification cycle.

* Environment Canada.

* Data not available for 15 April; 16 April data presented.
Cause of death for the 10 bats included rabies (n=1), possible hypothermia (n=1), and euthanasia of emaciated (n=7) and apparently healthy individuals (n=1). Four of the 10 bats had positive qPCR results for Pd (Table 1) and two of these also had a pustular, focal dermatitis with intralesional, unidentified ascomycetes. Fungi were identified as ascomycetes based on the presence of septa without clamp connections, the typical morphologic feature distinguishing the group (Alexopoulos et al. 1996). Although the monokaryon hyphae of basidiomycetes share this feature, most filamentous fungal infections are caused by ascomycetes and most pathogenic infections associated with basidiomycetes present solely in the single-celled yeast state (de Hoog et al. 2000). A third bat with similar, multifocal lesions was qPCR Pd-negative based on a quantification cycle >40, the cut-off for WNS surveillance protocol (Muller et al. 2013). Although Pd qPCR was positive in four of 10 E. fuscus, lesions consistent with WNS (Meteyer et al. 2009) were not present. Additionally, fungal hyphae or conidia morphologically consistent with the typical growth described for intralesional Pd in bats with WNS (Meteyer et al. 2009) were not identified microscopically. Skin lesions caused by the unidentified ascomycetes were only found on the muzzle and were minor and not associated with significant disease.

In the first case (CWHC 107769; Table 1), the fungus was on the epidermal surface in a solitary, intraepidermal, microscopic pustule on the muzzle of an adult female in good body condition that had been found alive inside a building. The fungal hyphae within the pustule were narrow (2.5–5.0 µm diameter) and septate with random branching. In contrast, the hyphae within the epidermis were often bulbous and almost torulose, with diameters <5.0 µm (Fig. 1A). In the second case (CWHC 107774), the fungus was in a solitary, microscopic, intraepidermal pustule in the muzzle skin of an immature female that was emaciated and found alive outside, adjacent to a building. The hyphae of one fungus were branched with infrequent septation and were variable in diameter (2.5–7.5 µm; Fig. 1C); these hyphae were not confined to the intraepidermal pustule but extended into the surrounding dermis. The morphology of the second fungus varied depending on its location in the lesion. On the epidermal surface, and within the intraepidermal pustule, there were slender (1.8–2.5 µm diameter), unbranched, septate hyphae. However, where they extended into the surrounding dermis, the diameters of the hyphae were 2.5–5.0 µm (Fig. 1D). Conidio genesis was not observed from any of the fungi.

Outside New Brunswick, big brown bats share caves with Myotis spp. and tricolored bats (Perimyotis subflavus). However, big brown bats are known to hibernate only in heated buildings in New Brunswick (McAlpine et al. 2002). Such behavior is common in the northern US (Whitaker and Gummer 2000). It is unknown how the Pd-positive big brown bats reported here might have acquired Pd. Although big brown bats may participate in swarming behavior at New Brunswick hibernacula, there is no evidence to support this.

Verant et al. (2012) found the optimal Pd growth temperature to be 12.5–15.8 °C, with atypical fungal morphologies observed above 12 °C; fungal morphologies typical for Pd were displayed between 0–7 °C. Vanderwolf et al. (2012) report average winter dark zone temperatures of 5.1±1.1 °C for New Brunswick underground hibernacula. While the descriptions of Verant et al. (2012) for atypical Pd morphologies share some similarities with the unidentified fungi we describe, and big brown bats were hibernating in buildings that likely provided the higher temperatures associated with atypical growth of Pd, there is no in situ or immunohistochemical diagnostic test
available to confirm that these unidentified fungi are \( Pd \). Nonetheless, our data demonstrate that \( Pd \) may occur on \( E. fuscus \) overwintering in buildings, although risk of the subsequent development of WNS is unclear. In spite of temperature conditions in buildings that are more conducive for \( Pd \) growth than are underground hibernacula, we could not confirm WNS in any of the \( Pd \)-infected big brown bats, consistent with Frank et al. (2014) findings that big brown bats have some resistance to WNS. Langwig et al. (2014) suggested that even occasional movements among underground hibernacula during winter could be important in \( Pd \) spread. Although big brown bats do not appear to use underground hibernacula in New Brunswick, these bats can regularly move in and out of winter roosts in buildings and caves (Beer 1955; Whittaker and Gummer 2000).

We can neither conclude nor discount that the fungi in the big brown bat lesions were \( Pd \). Ancillary diagnostic tests, such as in situ hybridization or immunohistochemistry, are needed so that \( Pd \) can be detected directly within a microscopic lesion, particularly in the absence of identifying features such as conidia. As the host–pathogen relationship for \( Pd \) evolves, and where bats are exposed to the fungus under varying temperature regimes, lesions may become less stereotypic, and such tests could help define these changes.

We are grateful to the following members of the public for submitting bat specimens for
testing: John Boudreau, Cara Hazelton, Wendell Price (Canadian Border Services Agency), Joe Robichaud, and Larry Tannahill. We are especially grateful to Melissa Behr, University of Wisconsin, School of Veterinary Medicine; Carol Meteyer, US Geological Survey, National Wildlife Health Center; and David Overy, Department of Pathology and Microbiology, Atlantic Veterinary College, University of Prince Edward Island, who graciously provided constructive comments on the pathology as it related to the fungal infections reported here, as well as David Groman with Aquatic Diagnostic Services, Atlantic Veterinary College, for preparation of photomicrographs and Hamid Haghighi and Patricia Bell-Rogers with Animal Health Laboratory, University of Guelph, for qPCR testing of our samples.

LITERATURE CITED


Submitted for publication 26 March 2015. Accepted 7 January 2016.
ERRATUM

“Molecular Detection of *Pseudogymnoascus destructans* (Ascomycota: Pseudeurotiaceae) and Unidentified Fungal Dermatitides on Big Brown Bats (*Eptesicus fuscus*) Overwintering in Buildings” by Donald F. McAlpine et al. [Journal of Wildlife Diseases, 2016, 52(4):902–906; DOI: 10.7589/2015-03-076]. The table (p. 903) contains errors. Those errors have been corrected in the online version.