

Inoculation of bats with European *Geomyces destructans* supports the novel pathogen hypothesis for the origin of white-nose syndrome

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Edited by Jitender P. Dubey, US Department of Agriculture, Beltsville, MD, and approved March 9, 2012 (received for review January 9, 2012)

White-nose syndrome (WNS) is an emerging disease of hibernating bats associated with cutaneous infection by the fungus *Geomyces destructans* (*Gd*), and responsible for devastating declines of bat populations in eastern North America. Affected bats appear emaciated and one hypothesis is that they spend too much time out of torpor during hibernation, depleting vital fat reserves required to survive the winter. The fungus has also been found at low levels on bats throughout Europe but without mass mortality. This finding suggests that *Gd* is either native to both continents but has been rendered more pathogenic in North America by mutation or environmental change, or that it recently arrived in North America as an invader from Europe. Thus, a causal link between *Gd* and mortality has not been established and the reason for its high pathogenicity in North America is unknown. Here we show that experimental inoculation with either North American or European isolates of *Gd* causes WNS and mortality in the North American bat, *Myotis lucifugus*. In contrast to control bats, individuals inoculated with either isolate of *Gd* developed cutaneous infections diagnostic of WNS, exhibited a progressive increase in the frequency of arousals from torpor during hibernation, and were emaciated after 3–4 mo. Our results demonstrate that altered torpor-arousal cycles underlie mortality from WNS and provide direct evidence that *Gd* is a novel pathogen to North America from Europe.

fungal pathogen | infectious disease | invasive species | Chiroptera | wildlife conservation

White-nose syndrome (WNS) is a rapidly spreading wildlife disease caused by the cold-tolerant fungus *Geomyces destructans* (*Gd*) (1). WNS has killed millions of bats across 16 US states and four Canadian provinces since its emergence in New York State in 2006 (2). So far, nine bat species from three genera, all of which hibernate in caves or mines, have been found to carry *Gd*, and mortality of infected bats has been observed in six of these species (2). The population dynamics of most affected species are not well understood but the effects of current declines are likely to be drastic; for example, the little brown bat (*Myotis lucifugus*) was the most widespread and common bat species in North America before WNS, but is now predicted to face local extinction in WNS-affected areas within two decades (3). During hibernation, the skin of WNS-affected bats is colonized by *Gd*, which invades cutaneous tissues of the muzzle, ears, and wings (4, 5). Major inflammation is usually not observed in infected tissues (6), possibly because immune responses in hibernating animals are suppressed (7). Mortality occurs in the second half of the hibernation season and affected bats are typically emaciated. Recently Lorch et al. (1) showed that experimental inoculation of *M. lucifugus* with *Gd* caused the characteristic wing lesions associated with WNS, and confirmed that *Gd* can be spread by

direct contact between bats. However, no study has established a causal mechanism linking *Gd* with bat mortality.

One possible explanation for mortality from WNS is that *Gd* causes a disruption of energy balance during hibernation. Hibernating mammals spend the majority of their time in torpor, a state of controlled reduction in body temperature (T_b) and metabolic rate, which is interrupted by brief periodic arousals to normothermic T_b (8). Although these arousals last less than 24 h in most species, the high metabolic cost of thermoregulation during normothermia at a low ambient temperature (T_a) means they account for the vast majority of over-winter energy expenditure (8, 9). Food is unavailable for most temperate-zone bats during winter, so they must survive on stored fat (9). Therefore, one hypothesis to explain WNS-related mortality is that *Gd* causes bats to increase the duration and/or frequency of periodic arousals, resulting in premature depletion of fat and consequently starvation (10). Preliminary support for this hypothesis was found based on an energetic model (11) but, to date, there is no experimental evidence that bats infected with *Gd* spend more time out of torpor than uninfected controls.

In addition to the mechanism underlying mortality, the origin of WNS is still unknown. There are two competing explanations for the origin of any emerging infectious disease (12). Such a disease may result from a pathogen that has been present historically but is rendered more pathogenic by a genetic mutation or environmental change (i.e., the endemic pathogen hypothesis). Alternatively, a pathogen may arrive in a new geographic area and encounter a naive host population (the “novel” or invasive pathogen hypothesis) (12). It is now established that *Gd* occurs at low levels on bats throughout Europe, where it has been isolated from eight *Myotis* spp., but with no evidence of mass mortality (13, 14). Given that *Gd* went undiscovered in Europe until WNS was observed in North America, one possibility is that *Gd* has occurred historically at low levels on bats from both continents but went unnoticed until mass mortality of bats in North America led to intensive sampling for a potential pathogen. This theory is cause for concern because European bats could be at risk from the accidental introduction of North

Author contributions: P.M.C., D.S.B., and C.K.R.W. designed research; L.W., J.M.T., T.K.B., V.M., and C.K.R.W. performed research; J.M.L., G.W., and D.S.B. contributed new reagents/analytic tools; L.W., J.M.T., and C.K.R.W. analyzed data; and L.W., J.M.T., and C.K.R.W. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

See Commentary on page 6794.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1200374109/-DCSupplemental.

American *Gd* to bat hibernacula in Europe. Alternatively, *Gd* may have arrived in North America as a recent invader from Europe, perhaps introduced by tourists visiting caves. Wibbelt et al. (14) hypothesized that under this novel pathogen scenario European bats may have coevolved with *Gd* over many years, and differences in its apparent pathogenicity for North American versus European bats could reflect differences in the physiology or behavior of the bats or differences in their environments, rather than intercontinental differences in *Gd*. Confirming one or the other of these hypotheses is essential because different disease-management strategies are warranted for invasive versus endemic pathogens (12).

We conducted an inoculation experiment with *M. lucifugus* to evaluate three hypotheses important for our understanding of WNS. First, we tested a key prediction of the novel pathogen hypothesis, which predicts that *Gd* isolated from Europe should cause the same clinical signs in a North American bat species as *Gd* isolated from North America. Therefore, we inoculated individual *M. lucifugus* with either a North American isolate of *Gd* (*NAGd*) or a European isolate (*EUGd*) and assessed clinical signs following several months of infection. Second, we tested whether inoculation with *Gd*, alone, is sufficient to cause mortality, a fundamental question about WNS that has still not been addressed. Third, by monitoring skin temperatures of bats

following inoculation, we assessed the hypothesis that infection with *Gd* causes bats to increase the frequency and/or duration of periodic arousals during hibernation, leading to premature fat depletion (10, 11). Importantly, we kept animals in environmental conditions closely matched to those of *M. lucifugus* hibernacula (9), particularly in terms of high relative humidity (RH).

Results

All bats entered multiday torpor bouts (i.e., began hibernating) within the first week of the study (Fig. 1). Average torpor bout duration over the entire study period was 9.0 ± 1.0 d for *NAGd* bats (individual range 1.2–32.4 d), 6.1 ± 0.6 d for *EUGd* bats (1.0–21.8 d), and 16.0 ± 0.9 d for sham-inoculated control group (*CO*) bats (2.0–33.4 d). Both *NAGd* and *EUGd* caused a progressive increase in the frequency of periodic arousals over the course of the experiment. There was no significant difference among groups during Interval 1, but treatment groups aroused significantly more often throughout the rest of hibernation (Table 1). In fact, during Interval 3, arousal frequency of *NAGd* bats was three times—and *EUGd* bats four times—that of *CO* bats (Fig. 2 *A* and *B*). The effect of time on arousal frequency was significant for each group (Table 2) with a significant increase over time for both *NAGd* and *EUGd* bats, and a decrease for *CO* bats (Fig. 2*A*). In contrast, the duration of periodic arousals was not affected by inoculation (Fig. 2*C* and Table 1), nor by time for any group (Table 2).

Both isolates of *Gd* caused all known clinical signs of WNS (5, 6), including loss of elasticity, irregular pigmentation and stickiness of wing tissue, and white surface growth. Histopathology confirmed infection in all *NAGd* and *EUGd* bats as fungal hyphae penetrated the epidermis and damaged underlying tissues, consistent with previous studies (5, 6) (Fig. 3 *A* and *B*). *Gd* was also cultured from sections of wing tissue for bats from both treatment groups but no *CO* bats showed any evidence of infection by *Gd* (Fig. 3*C*).

There was a highly significant effect of inoculation on survival for both *NAGd* ($\chi^2 = 17.1$, $P < 0.001$) and *EUGd* bats ($\chi^2 = 26.4$, $P < 0.001$) compared with controls, and *NAGd* bats survived significantly longer than *EUGd* bats (Fig. 4) ($\chi^2 = 20.3$, $P < 0.001$). Mortality was first observed for *EUGd* bats on day 71, and surviving bats from the group were euthanized on day 91 after 16 bats reached moribund status. Mortality first occurred for *NAGd* bats on day 88 and surviving bats from this group were terminated on day 114 after 12 bats reached moribund status. Two *EUGd* and four *NAGd* bats were unable to arouse from torpor when removed from the chamber at the end of the experiment and were therefore also considered moribund. Based on necropsies, bats from both treatment groups had virtually no fat reserves remaining. On day 119, when we terminated the *CO*

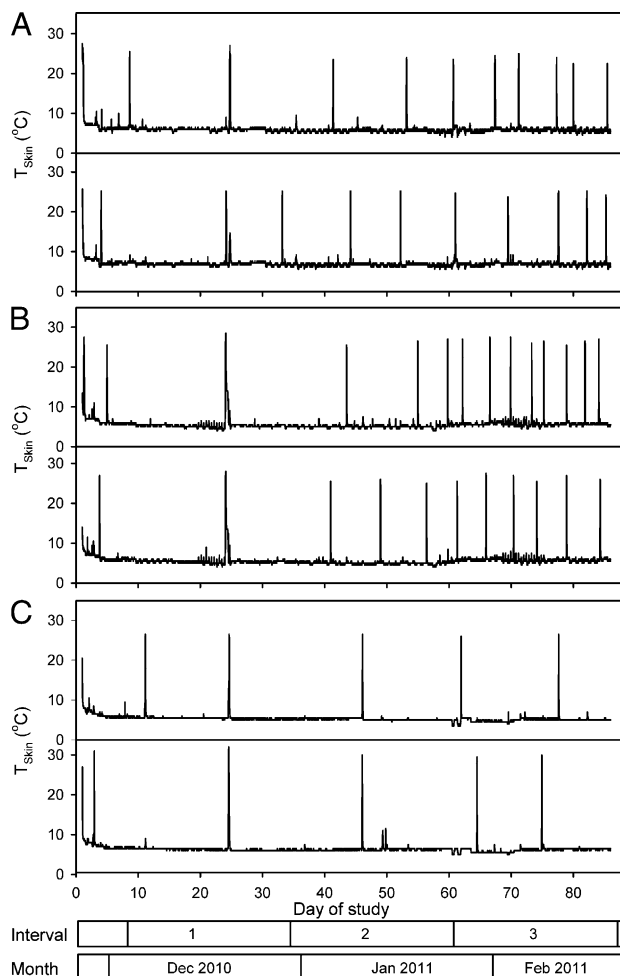


Fig. 1. Representative traces of skin temperature (T_{skin}) for six *M. lucifugus*, two each from the following groups: (*A*) inoculated with *NAGd*; (*B*) inoculated with *EUGd*; (*C*) sham-inoculated control. The x axis shows the day of study, where day 1 is November 27, 2010; the bars at the bottom indicate the division of the study period into 26-d intervals, and months.

Table 1. Sample sizes and ANOVA results for arousal frequency and arousal duration

	Interval	<i>NAGd</i> (n)	<i>EUGd</i> (n)	<i>CO</i> (n)	df_1	df_2	F	P
Arousal frequency	1	10	11	18	2	36	0.52	0.601
	2	10	11	18	2	36	20.79	<0.001
	3	14	11	18	2	40	52.82	<0.001
	4	4	—	5	1	7	10.79	0.013
Arousal duration	1	8	6	15	2	26	0.24	0.789
	2	10	13	18	2	38	1.57	0.221
	3	14	12	18	2	41	0.21	0.811
	4	4	—	5	1	7	0.76	0.411

Sample sizes and ANOVA results for arousal frequency (arousal bat⁻¹·d⁻¹) and arousal duration (length of time above skin temperature threshold) for the four time intervals for *M. lucifugus* inoculated with *NAGd* and *EUGd* compared with sham-inoculated controls (*CO*). The results of the pair-wise post hoc comparisons (SNK) are indicated in Fig. 2. Significant results are in bold.

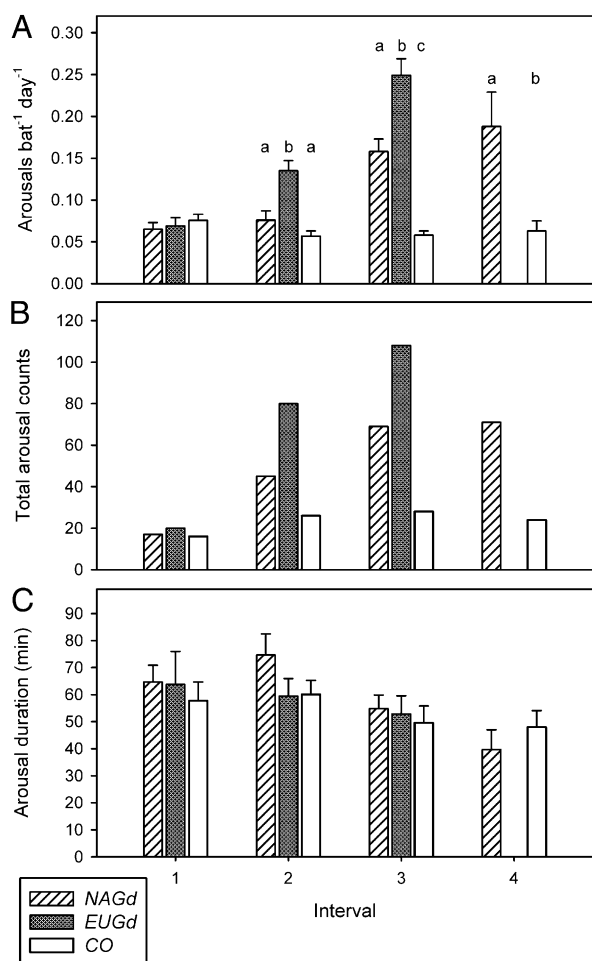


Fig. 2. Changes in torpor patterns in *M. lucifugus* following inoculation with *NAGd*, *EUGd*, or *CO*. Frequency of arousals based on skin temperature (A), total count of arousals based on video observations (B), and mean arousal duration (C). Within intervals, different letters above bars indicate significant differences between groups (SNK post hoc tests following significant ANOVA in Table 1).

group, all bats were alive, capable of endogenous arousal from torpor, and still had subcutaneous fat reserves.

Discussion

The susceptibility of a North American bat species to both *EUGd* and *NAGd* strongly supports the novel pathogen hypothesis that accidental introduction of *Gd* from Europe is responsible for the WNS-related mass mortality of bats in North America. Our data suggest that the absence of mortality observed among European bats infected with *Gd* reflects different physiological and behavioral responses of European versus North American bats rather than a heightened pathogenicity of *NAGd* (14). This finding also supports the hypothesis of Wibbelt et al. (14) that *Gd* may have impacted European bat populations in the past and that bats in Europe have coevolved resistance to (e.g., via immune system responses), or tolerance of (e.g., via behavioral adaptations), infection with *Gd*. These findings have significant implications for management and future research. Endemic pathogens are best addressed via management of factors that enhance virulence of the pathogen (e.g., environmental or biotic cofactors), and novel pathogens are best dealt with by managing the agents that spread the disease (12). Managing agents of spread for WNS will be impractical, if not impossible, because

Table 2. Repeated-measures ANOVA results for within-group effects of time on arousal frequency and arousal duration

	Group	n	df ₁	df ₂	F	P
Arousal frequency	<i>NAGd</i>	10	2	8	16.10	0.002
	<i>EUGd</i>	10	2	8	34.59	<0.001
	<i>CO</i>	18	2	16	8.28	0.003
Arousal duration	<i>NAGd</i>	8	2	6	1.43	0.310
	<i>EUGd</i>	6	2	4	1.59	0.311
	<i>CO</i>	15	2	13	1.61	0.238

Repeated-measures ANOVA results for within-group effects of time on arousal frequency (arousal bat⁻¹·d⁻¹) and arousal duration (length of time above skin temperature threshold) for *M. lucifugus* inoculated with *NAGd* and *EUGd* compared with sham inoculated controls (*CO*). Significant results are in bold.

the putative agents (i.e., the bats) are highly cryptic, widely dispersed for much of the year, and wide-ranging. However, our results support the high priority of research aimed at understanding temporal and spatial aspects of *Gd* transmission in the wild, as this work could aid in the development of management strategies focused on critical locations or times of year when *Gd* is likely to be transmitted. Encouragingly, our findings suggest that European bats face little risk from the possible reintroduction of *Gd* from North America to Europe, although it would be useful to repeat our experiment with a European bat species.

Interestingly, we found that *EUGd* affected *M. lucifugus* more quickly than *NAGd* (Figs. 1 and 4). Rapid evolution of the host-pathogen interaction between *Gd* and bats could help explain this pattern (12, 15). For example, if European bats exhibit resistance to infection, *Gd* in Europe may face intense selection pressure for increased production of potential virulence factors and more rapid growth to facilitate its propagation and transmission. However, if the production of virulence factors and rapid growth are costly for *Gd*, selective trade-offs could quickly favor a less pathogenic, slower growing variant of the fungus as it infected a naive host population in North America. Moreover, dramatic population declines of North American bats in the early years of the epizootic could have reduced the potential for transmission among bats, enhancing selection for reduced pathogenicity in North America. Despite this potentially encouraging finding, clearly the version of *Gd* now present in North America is highly pathogenic to a number of bat species. Thus, more laboratory and field experiments are necessary to better understand interactions between bats and *Gd*, particularly studies aimed at better understanding transmission of the fungus in the wild.

Our study also confirms that *Gd* causes mortality of hibernating bats and provides direct evidence for the hypothesis that an increase in arousal frequency during hibernation is the mechanism underlying mortality. The three- to fourfold increase in arousal frequency we observed for infected bats is similar to the pattern predicted by Boyles and Willis (11) based on an energetic model. The additional arousals would prematurely deplete the stored energy of a small hibernator like *M. lucifugus* which, in its northern distribution, must survive >190 d exclusively on fat reserves (9, 16). Periodic arousals account for only 1.2% of the hibernation time budget, yet the thermoregulatory cost of each arousal amounts to about 5% of the winter energy budget (9). Hence, each additional arousal shortens the time a bat is able to hibernate by about 9 d. WNS-affected bats are often observed flying outside hibernacula during the daytime in winter (4), possibly searching for food and, like the *Gd*-inoculated bats in our study, WNS-affected carcasses collected from hibernacula after mass mortality events were emaciated (4). Hence, we conclude that infection with *Gd* causes an increase in arousal frequency, leading to emaciation because fat reserves are used prematurely.

20631–21 (American Type Culture Collection, ATCC MYA-4855) (5) isolated from a *M. lucifugus* collected in New York on February 2, 2008. The *EUGd* isolate (MmyotGER2) was obtained from a greater mouse-eared bat (*Myotis myotis*) collected in Thuringia, Germany, on March 7, 2009 (14). CO bats were sham-inoculated with 20 μ L of PBS-Tween-20 solution lacking fungal conidia.

Skin Temperature. All bats were equipped with one of two types of device to record skin temperature (T_{skin}): either temperature-sensitive radio transmitters (LB-2NT; Holohil Systems) or data loggers (DS1922L-F5 Thermochron iButton, Maxim; and iBBat, Alpha Mach). T_{skin} was recorded every 15 min.

Behavior. Infrared cameras inside each environment chamber allowed us to monitor behavior and count the total number of arousals from torpor for each group within each interval (Fig. 2B). These data were clearly consistent with T_{skin} (compare Fig. 2 A and B).

Histopathology. We examined multiple sections from the left wing, as well as nose and ear, following Meteyer et al. (6). Tissues were fixed in formalin immediately after bats were euthanized and later stained for histopathological examination using the periodic-acid Schiff method. All *NAGd* and *EUGd* bats exhibited the epidermal lesions typical of WNS (5, 6).

Analyses. The study period was divided into four intervals of 26.3 d each. We tabulated the number of arousals from torpor for each individual to generate mean values for each treatment group. Full-factorial ANOVA was used to analyze differences in torpor bout duration and arousal duration among groups within each interval. Student-Newman-Keuls (SNK) post hoc tests were used for pair-wise comparisons following a significant ANOVA result. To examine the effect of time on torpor patterns (i.e., arousal frequency and arousal duration) we used repeated-measures ANOVA testing for differences among the first three intervals within each group. A Breslow-Gehan survival analysis was used to test for differences in the time to mortality/moribund status for the three groups with a Bonferroni correction to account for multiple comparisons between each pair of groups. All analyses were conducted using *statistiXL* v7.0 and *Systat* v11.0.

ACKNOWLEDGMENTS. We thank M. Burmester, P. Mason, and M. Weiss for animal care support; C. Rainbow, C. Wilson, and M. Zimmer for pathology assistance; P. Withers for assistance with statistical analyses; M. Kilpatrick and W. Frick for helpful discussions; and M. Brigham, B. Fenton, and an anonymous reviewer for excellent comments on drafts of the manuscript. Funding was provided by a US Fish and Wildlife Service grant (to C.K.R.W., D.S.B., and P.M.C.); grants from the Natural Sciences and Engineering Research Council, the Canada Foundation for Innovation and Manitoba Research, and Innovation Fund (to C.K.R.W.); and a Government of Canada Post-Doctoral Research Fellowship and a fellowship within the Postdoc Programme of the DAAD, German Academic Exchange Service (to L.W.).

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