1	TITLE: Distribution and Environmental Persistence of the Causative Agent of White-
2	Nose Syndrome, Geomyces destructans, in Bat Hibernacula of the Eastern United States
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4	RUNNING TITLE: Distribution and Persistence of <i>Geomyces destructans</i>
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24 ABSTRACT

25

26 White-nose syndrome (WNS) is an emerging disease of hibernating bats caused by the 27 recently described fungus Geomyces destructans. First isolated in 2008, the origins of 28 this fungus in North America and its ability to persist in the environment remain 29 undefined. To investigate the correlation between manifestation of WNS and distribution 30 of G. destructans in the U.S., we analyzed sediment samples collected from 55 bat 31 hibernacula (caves and mines) both within and outside the known range of WNS using a 32 newly developed real-time PCR assay. Geomyces destructans was detected in 17 of 21 33 sites within the known range of WNS at the time the samples were collected; the fungus 34 was not found in 28 sites beyond the known range of the disease at the time that 35 environmental samples were collected. These data indicate that distribution of G. 36 *destructans* is correlated with disease in hibernating bats and support the hypothesis that the fungus is likely an exotic species in North America. Additionally, we examined 37 38 whether G. destructans persists in infested bat hibernacula when bats are absent. 39 Sediment samples were collected from 14 WNS-positive hibernacula, and the samples 40 were screened for viable fungus using a culture technique. Viable G. destructans was 41 cultivated from 7 of the 14 sites sampled during late summer when bats were no longer in 42 hibernation, suggesting the fungus can persist in the environment in the absence of bat 43 hosts for long periods of time. 44

45

46 INTRODUCTION

48	White-nose syndrome (WNS) is an emerging wildlife disease that by one estimate (see
49	http://www.whitenosesyndrome.org/news/north-american-bat-death-toll-exceeds-55-
50	million-white-nose-syndrome) has killed approximately 5.5 million hibernating bats in
51	North America since its discovery in 2007. The disease results from cutaneous infection
52	by the recently described fungus, Geomyces destructans (1,2), and has been implicated in
53	population declines of 72-88% for hibernating bat species inhabiting the northeastern
54	U.S. (3-5). Since 2007, the disease has spread across the eastern U.S. and Canada,
55	threatening the future of North American bat populations (5,6). The sudden emergence
56	and rapid spread of WNS has led to questions regarding the origin of G. destructans in
57	North America and how the lifecycle of the fungus allows it to exert such significant
58	impacts on hibernating bat populations.

60 Laboratory experiments have demonstrated that G. destructans is the causative agent of 61 WNS in a North American bat species, eliciting the disease in apparently healthy animals 62 (7,8). In addition, G. destructans has been found to occur on hibernating bats throughout 63 most of Europe, but it has not been associated with unusual bat mortality on the European 64 continent (9-12). Together, these findings prompt two hypotheses regarding the origin of 65 G. destructans in North America that are consistent with the emergence of a novel 66 infectious disease (13): 1) Geomyces destructans is endemic to North America, but a 67 pathogenic strain spontaneously emerged and is spreading across the landscape (8); or 2) 68 Geomyces destructans was recently introduced to North America where it is behaving as 69 an exotic pathogen among naïve populations of bats (8,9,11).

71	Recent research supports the exotic species hypothesis. For example, a European isolate
72	of G. destructans has been found to induce lesions diagnostic for WNS and mortality in
73	an experimentally infected North American bat species (8). This demonstrates that a
74	European isolate of the fungus is highly pathogenic to North American bats despite
75	having no apparent effects on wild bat populations of Europe and implicates Europe as
76	the possible source for introduction of G. destructans to North America (12). In addition,
77	isolates of G. destructans from the eastern U.S. appear to be genetically identical (14),
78	suggesting that G. destructans in North America is derived from a single isolate that may
79	have been introduced to this continent.

81 While evidence is mounting to support the hypothesis that G. destructans was introduced 82 to North America, it remains unclear why WNS-related mortality varies between bats of 83 North America and Europe (12) and why some North American species appear to be 84 more vulnerable than others (5). Environmental effects, genetic composition, and 85 behaviors differ among bat species and likely play a role in facilitating infection, disease 86 progression, and mortality, but such factors are difficult to tease apart without a basic 87 understanding of whether presence of G. destructans in a hibernaculum correlates with 88 manifestation of WNS in bats. Furthermore, the current assumption that G. destructans is 89 limited to areas where WNS has been observed may be biased by the primary means of 90 detecting the fungus through diagnostic analysis of samples derived from sick or dead 91 bats. A previous study addressed these issues by screening sediment samples from bat 92 hibernacula to determine whether G. destructans was indeed restricted to areas where

93	WNS has been observed in bats (15). While nucleic acid from <i>G. destructans</i> was found
94	to occur at three sites within the known range of WNS and no sites outside the range of
95	WNS, the number of sites sampled was too small given the low detection rate to conclude
96	the distribution of the fungus was correlated to that of the disease. In addition, the
97	method utilized in that study lacked specificity because similar species of Geomyces
98	cross-reacted with primers of the conventional PCR-based method (16) and may have
99	masked the presence of G. destructans if it was at a low abundance. Screening a greater
100	number of sediment samples from hibernacula using a more specific and sensitive
101	technique, such as a recently-described real-time PCR assay (17), may serve to better
102	determine whether distribution of G. destructans is limited to areas where WNS occurs or
103	whether the fungus is more widespread in North America than currently thought.
104	
105	The ability to detect G. destructans in environmental samples using PCR-based methods
106	could also reveal important information about WNS disease dynamics. Current
107	determination of whether G. destructans is present in a hibernaculum usually relies upon
108	first detecting the fungus on sick bats, which makes it difficult to address questions such
109	as: when did the fungus arrive at a given site; how long does it takes for disease to
110	manifest after the arrival of G. destructans at a new site; and is disease an inevitable
111	outcome of the fungus' presence in a hibernaculum? Understanding these aspects of the
112	pathogen's interaction with the environment and its host will facilitate disease
113	surveillance of bat hibernacula and potentially enable earlier deployment of
114	interventional strategies to more effectively limit the spread of and reduce mortality
115	caused by WNS.

117	Most fungi pathogenic to mammals can persist in the environment in the absence of a
118	host (18,19). Given the temperature requirements for growth of G. destructans (i.e. it
119	does not grow at or above approximately 20°C [20]), caves and mines have
120	environmental characteristics consistent with potential long-term reservoirs for the
121	fungus as they remain cool throughout the year, even when bats are absent during
122	summer months. While follow-up culture analyses of sediment samples that contained
123	DNA from G. destructans from a previous study (15) proved that viable G. destructans
124	was present (21), these samples were collected during the hibernation season and may
125	have represented only short-term survival of the fungus after it detached from a bat host.
126	Similarly, G. destructans was cultured from the wall of a cave in Estonia where an
127	infected bat had been observed nine days prior (12), again demonstrating only temporary
128	persistence. Thus, the ability of G. destructans to survive long-term in the environment
129	in the absence of its bat hosts remains uncertain.

130

131 The objectives of this study were to: 1) determine the distribution of G. destructans in 132 underground bat hibernacula of eastern North America and examine whether presence of 133 the fungus strictly correlates with occurrence of WNS; and 2) establish whether 134 hibernacula can serve as reservoirs for G. destructans during the summer months when 135 bats are largely absent. To address the first objective, we screened sediment samples 136 collected from bat hibernacula across the eastern U.S. for the presence of G. destructans 137 using a real-time PCR test (17). To address the second objective, sediment samples 138 collected from WNS-affected hibernacula were screened for viable G. destructans using a

- 139 previously-described culture technique (21) during seasons of both bat hibernation (when 140 bats are present in hibernacula) and activity (when bats are largely absent from 141 hibernacula). By demonstrating the utility of environmental sampling as a non-invasive 142 tool for detecting G. destructans, results from these investigations offer the potential to 143 refine WNS surveillance and management. 144 145 146 MATERIALS AND METHODS 147 148 Sample Collection. Sediment samples used for fungal distribution analysis were 149 collected from the eastern U.S. by volunteers during the winter of 2008-2009. For each 150 sample, clean latex gloves were worn and sterile wooden splints were used to transfer 151 sediment into sterile, labeled sampling bags. A minimum of five samples were collected 152 from the floor of each cave or mine (hereafter referred to as a "site") and immediately 153 shipped on ice to the U.S. Geological Survey – National Wildlife Health Center 154 (Madison, WI) where they were stored at -80°C. The samples included in this study represented a total of 56 sites from 22 states east of the 95th meridian west (95°W 155 156 longitude), including 8 states within and 14 states outside the known range of WNS at the 157 time the samples were collected (Fig. 1; Table 1). Exact locations of the sampled sites 158 are not provided due to the sensitive nature of bat hibernacula. 159 160 Sediment samples for environmental persistence analysis were collected from 14 bat
- 161 hibernacula in which bats with WNS had been previously identified. These consisted of

162	4 sites in New Hampshire, 2 sites in Vermont, 3 sites in Virginia, and 5 sites in West
163	Virginia (see Table 2). Five locations within each site were marked, and samples were
164	serially collected within 30 cm of the markers on three separate occasions: once in
165	February-March 2011 (during the bat hibernation period; hereafter referred to as winter
166	2010-2011), once in late July-late August 2011 (near the end of the active season and just
167	prior to large congregations of bats returning to the hibernacula; hereafter referred to as
168	summer 2011), and again in October 2011-March 2012 (during the next consecutive
169	hibernation period; hereafter referred to as winter 2011-2012). Exceptions were: samples
170	were not obtained from sites C6, C7, and C8 during the third sampling period (i.e. winter
171	2011-2012); several sampling markers could not be relocated within sites C7 and C9
172	during the second visit (i.e. summer 2011), and samples were collected from approximate
173	locations as determined by collectors. Additionally, site C4 flooded in September 2011;
174	the markers were relocated during the winter 2011-2012 visit although 2.5-20 cm of
175	sediment had been deposited on top of the previously sampled sediment.
176	
177	DNA Extraction and PCR Analysis. DNA was extracted from the sediment samples
178	for the distribution study using the PowerSoil [™] DNA Isolation Kit (MoBio Laboratories,
179	Inc., Carlsbad, CA) according to the manufacturer's instructions. If more than five
180	samples were collected from a given site, five were chosen at random for inclusion in the
181	study. All extracted DNA samples were stored at -20°C.
182	
183	Real-time PCR targeting the intergenic spacer (IGS) region of the rRNA gene complex of

184 G. destructans was performed on an Applied Biosystems 7500 Fast Real-Time PCR

186	microliters of each DNA extraction from sediment (diluted 1:1 and 1:10) were added to
187	each 25 µl PCR reaction. All plates included at least two positive (3.3 pg G. destructans
188	genomic DNA [gDNA]) and one negative (water added in place of template) control
189	samples. Individual samples that crossed the cycle threshold (set at 10% of the maximum
190	fluorescence of the positive control sample for each plate [17,22]) within 40 cycles were
191	considered positive for presence of G. destructans. Further, a sample was identified as
192	positive if either or both template dilutions (1:1 or 1:10) crossed the cycle threshold as
193	described above; a site was considered positive when at least one sample from that site
194	was PCR-positive.
195	
196	Sediment often contains humic acid and other substances that can inhibit DNA
197	amplification. Thus, prior to conducting the real-time PCR assay, all samples were
198	screened for PCR inhibition to reduce the chance of false negative results. Conventional
199	PCR targeting the internal transcribed spacer (ITS) region of the rRNA gene was
200	performed with primers ITS4 and ITS5 (23) using $GoTaq^{\mathbb{R}}$ Flexi DNA Polymerase
201	(Promega Corporation, Madison, WI) according to the manufacturer's instructions. Five
202	microliters of the 1:10 diluted sediment DNA extraction was used as template. Cycling
203	conditions were as follows: 98°C for 2 min, then 30 cycles of 98°C for 10 sec, 50°C for
204	30 sec, and 72°C for 1 min, followed by a final extension for 7 min at 72°C.
205	Amplification products were analyzed using an agarose gel. Control reactions containing
206	33 fg gDNA isolated from pure cultures of G. destructans (positive control) or without

System (Applied Biosystems, Foster City, CA) as described previously (17). Five

207 template added (negative control) were also included. Samples failing to yield bands by

208	the ITS PCR were subsequently spiked with gDNA from G. destructans and used as
209	template in a modified version of the real-time PCR assay (17) adapted for use by
210	conventional PCR. Reagents used were as described for the ITS PCR. Five microliters
211	of the 1:10 diluted sediment DNA extraction, 5 μL (containing 33 fg) gDNA, and 1.25
212	μL of each primer used in the real-time PCR were included in the 25 μL total reaction
213	volume. No probe was added. Control reactions as described for the ITS PCR were also
214	included. Cycling conditions were identical to those used for the real-time PCR assay
215	(17). Spiked samples that did not yield amplification products were considered
216	inhibitory. If one or more samples showed evidence of inhibition, the entire site was
217	excluded from the dataset. To ensure the qualitative results of the inhibition screen using
218	conventional PCR were consistent with the more quantitative real-time PCR, a subset of
219	samples was also screened for inhibition on the real-time PCR platform. Single samples
220	from 45 individual sites that tested negative for the presence of G. destructans by real-
221	time PCR were randomly selected, spiked with 33 fg of G. destructans gDNA, and used
222	as template in the real-time PCR assay. Samples within one C_t value of the positive
223	control well containing 33 fg G. destructans gDNA were considered non-inhibitory.
224	
225	PCR amplicons from each PCR-positive site were subjected to cloning and sequencing to
226	confirm an exact sequence match to G. destructans. PCR products were cloned as
227	described previously (24) and prepared for sequencing according to pre-established
228	methods (15) using primers SP6 (TATTTAGGTGACACTATA) and T7
229	(TAATACGACTCACTATAG), which target pGEM-T (Promega, Madison, Wisconsin)
230	up- and down-stream of the insert. Because of short size of the amplicons (103 bps),

blue/white screening of bacterial colonies was not possible, so approximately 8-16
random colonies were chosen for screening by PCR amplification, and those yielding
amplification products were further characterized by DNA sequencing.

234

235 **Culture Analysis.** For the environmental persistence analysis, approximately 200 mg of 236 each thawed sediment sample was placed into a sterile microcentrifuge tube, suspended 237 in 0.5 mL sterile, deionized water, and serially diluted as previously described (21). 238 Sabouraud dextrose agar plates containing chloramphenicol and gentamycin (BD 239 Diagnostic Systems, Sparks, MD) were inoculated by spreading 150 μ L of the 10⁻¹, 10⁻², and 10^{-3} dilutions onto the medium. Each dilution was plated in duplicate. The plates 240 241 were incubated at 7°C and checked at 30 days and once weekly thereafter for a total of 60 242 days. Colonies of G. destructans were initially identified by examining tape lifts of 243 suspect colonies using a 40X objective to identify characteristic crescent-shaped conidia 244 borne at the end of verticillately branching conidiophores (2). At least one colony of G. 245 *destructans* from each site at each time-point, when present, was isolated in pure culture. 246 The ITS region of the rRNA gene of each of these isolates was then sequenced to confirm 247 identification of G. destructans using primers ITS1-F and ITS4 (25) and PCR conditions 248 as described previously (21). 249

Data Analysis. Hibernacula for the fungal distribution analysis were categorized as
 occurring within one of three zones based on WNS distribution at the time of sample

collection: 1) the WNS zone (general geographic area within which the disease had been

253 documented); 2) the buffer zone (general geographic area within which WNS was

254	documented the year following sample collection); and 3) the outside zone (general
255	geographic area within which WNS was not documented until at least two years after
256	sample collection) (Table 1; Fig. 1). Not all hibernacula falling within the WNS zone
257	had been confirmed to contain bats exhibiting signs of the disease; similarly, not all
258	hibernacula within the buffer zone were confirmed to contain bats with WNS by the
259	following year. For this reason, individual sites were designated as being either diseased
260	sites (WNS documented at the site prior to or at the time the samples were collected),
261	buffer sites (WNS documented one year after the samples were collected), or clean sites
262	(WNS not documented to date or documented more than one year after the samples were
263	collected) based on interviews with individuals from state and federal wildlife agencies
264	(Table 1; Fig. 1).
265	
266	To test the null hypothesis that distribution of G. destructans is not associated with WNS
267	in North America, PCR results (i.e. the number of PCR-positive and PCR-negative bat
268	hibernacula) for sites that occurred within and outside the known range of WNS (WNS
269	and outside zones) at the time samples were collected were compared using Fisher's
270	exact test in SigmaPlot 11.2 (Systat Software, Inc., San Jose, CA). Sites that occurred

271 within the geographic buffer zone were excluded from this analysis because it was

272 equally plausible that *G. destructans* could be present or absent from those sites.

273

Estimated Probability of Detection. To determine the probability of PCR and culture analysis to detect *G. destructans*, the results of each test were formulated as a binomial variable (1 = G. *destructans* detected; 0 = G. *destructans* not detected). A detection 277 history for each sampled site was then created as a series of zeros and ones. For example, 278 a detection history of 101 for the environmental persistence analysis indicated that G. 279 destructans was detected on the first and last surveys but not the second survey. The 280 probability of detecting the fungus, if present, was then estimated in the following 281 manner. The observed values at site i and replicate (spatial or temporal) t $(y_{i,t})$, were the 282 detection histories, and represent the imperfect observation process uncorrected for the 283 ability of the diagnostic test to detect the fungus. These observations were modeled as 284 Bernoulli trials, where the probability of success (the probability that $y_{i,t}=1$, p.eff_{i,t}), was 285 the observed detection at site i and replicate t.

286

287 Mathematically, y_{i,t}~Bern(p.eff_{i,t}), and the observed detections of the fungus were a 288 function of the true infection status of the site (infected/clean) and the probability of the test correctly detecting the fungus if present. Therefore, $p.eff_{i,t} {=} z_i \, \text{X} \; p_{i,t};$ where z289 290 represented the true state of the site (infected or clean) and p was the true detection 291 probability. The observations, therefore, were imperfect reflections of the true state due 292 to imperfect detection probabilities. Further, when z was unknown, z was formulated as 293 Bernoulli trials, where probability of success = ψ , the true proportion of sites that were 294 infected (as opposed to the observed). The analysis was conducted in R (R Foundation 295 for Statistical Computing, Vienna, Austria) (26) using the R2WinBUGS library (27) 296 following pre-established methods (28). To aid in convergence, $logit(p_{i,t}) < -\alpha_1 + \beta_1 X$ was 297 formulated; where X was a matrix of covariate values (similarly, logit(ψ)<- α_2 + β_2 X). 298 The models for both analyses were checked for convergence using the Gelman-Rubin 299 diagnostics function (gelman.diag) in the CODA package in R (29).

301	For the PCR analysis, it was assumed the true infection status of buffer sites was
302	unknown and that the true infection status of diseased and clean sites was known. Thus,
303	z was set to 1 for diseased sites and 0 for clean sites. Priors for the estimates of αs and βs
304	were uniform (-5, 5). Ideally, detection and occupancy would have been estimated
305	separately for buffer and infected sites, but there were not enough buffer sites (n=4) for
306	this analysis to converge. Therefore, a pooled occupancy and detection probability was
307	estimated separately for buffer and infected sites.
308	
309	For the culture analysis it was assumed the true infection status for each site was known
310	as infected, and z was set to 1. Since multiple samples were collected within each cave
311	on 3 different sampling occasions, two models were run, one with an effect of time and
312	one including a random effect for cave. Random effects were formulated as (μ, τ) ,
313	where $\tau=1/(\sigma^*\sigma)$. Priors on μ were uniform (-5,5); priors on σ were also uniform (0,10).
314	
315	
316	RESULTS
317	
318	PCR Analysis. One of the 56 sites tested for the distribution portion of the study
319	exhibited PCR inhibition and was excluded from further analysis. The subset of samples
320	tested for inhibition using real-time PCR yielded identical results to the conventional
321	PCR, indicating the conventional PCR method used to screen samples for inhibition was
322	accurate. Sequences of PCR amplicons from all sediment samples that were PCR-

323 positive for *G. destructans* were 100% identical to the 103 nucleotide IGS region of the

324 type isolate of *G. destructans* (GenBank accession no. JX415267 [17]).

325

326 **Distribution and Environmental Detection of** *G. destructans.* Nucleic acid from *G.* 327 destructans was detected by real-time PCR in 47 samples collected during the winter of 328 2008-2009, representing 13 different sites (Table 1). Seven sites that were initially 329 sampled in winter 2008-2009 were re-sampled in winter 2010-2011 as part of the 330 environmental persistence study (see above). Four of these sites (C2, C10, C13, and 331 C14) that were PCR negative for G. destructans in winter 2008-2009 were subsequently 332 reanalyzed by real-time PCR using DNA extractions from samples collected during 333 winter 2010-2011 (site designations 23*, 22*, 20*, and 21*, respectively, in Table 1). 334 Bats from one of the four sites (site 20) had been diagnosed with WNS in winter 2008-335 2009, but bats from the remaining three sites (sites 21, 22, and 23) did not show signs of 336 the disease until the winters of 2009-2010 or 2010-2011. PCR analysis of these samples 337 (collected during winter 2010-2011) showed that DNA from G. destructans was present 338 in 9 of 20 samples, representing all four sites. 339 340 The occurrence of G. destructans as detected in the environmental samples was 341 synonymous with the known range of WNS at the time samples were collected (Fig. 1), 342 with hibernacula within the known range of WNS having significantly higher detection

rates for the fungus than hibernacula outside the range of the disease (p < 0.0001).

344 Sixteen of the 17 hibernacula in which the fungus was detected were diseased sites; the

remaining hibernaculum was a buffer site in which bats with WNS were observed the

346 following year. Of the 12 WNS-positive hibernacula from which G. destructans was 347 detected in winter 2008-2009, five had been designated as WNS-positive in winter 2007-348 2008, and seven were identified as WNS-positive in winter 2008-2009. Site 6 had not 349 been officially monitored since 1985, but the cave was considered WNS-positive based 350 upon observation of clinical signs in bats suggestive of the disease at the time the samples 351 were collected. The four hibernacula from which G. destructans was detected by PCR in 352 winter 2010-2011 included one site designated as harboring WNS-positive bats in winter 353 2008-2009, two in winter 2009-2010, and one in winter 2010-2011.

354

Environmental Persistence of *G. destructans*. *Geomyces destructans* was cultured from 27 of the 195 sediment samples collected from bat hibernacula in 2011-2012 with viable fungus detected in 11 of the 14 sites during at least one sampling interval (see Table 2). Seven of the 14 sites were found to harbor viable *G. destructans* in late summer when bats were either absent from the hibernacula or present in only low numbers. Sequences of the rRNA gene ITS regions of isolates from each site were 100% identical to the ITS region of the type isolate of *G. destructans* (GenBank accession no. EU884921 [2]).

362

Estimated Probability of Detection. All samples from the 36 caves designated as
"clean" (i.e. WNS-free) tested negative for *G. destructans*. The remaining 23 samples
(representing diseased and buffer sites) were combined into one dataset, and probability
of detection for the PCR assay was estimated. Estimated probabilities of detection for a
single sample were 0.56 [95% C.I. 0.47-0.67] for diseased sites and 0.11 [95% C.I. 0.020.29] for buffer sites. This indicated that with five samples from a given diseased site,

369	the probability of detecting the fungus was 0.98, and that 4 samples are sufficient to
370	obtain a mean estimated probability of detection >0.95. For buffer sites, the probability
371	of detecting G. destructans with 5 samples was 0.44, and at least 26 samples would be
372	needed from each site for the mean estimated probability of detection to be >0.95 .
373	
374	The overall probability of detection using the culture technique was 0.14 [95% C.I. 0.10-
375	0.19]. There was no difference between detection probabilities by time period (0.16
376	[95% C.I. 0.080-0.25] in winter 2010-2011; 0.13 [95% C.I. 0.062-0.22] in summer 2011;
377	and 0.13 [95% C.I. 0.054-0.23] in winter 2011-2012). However, the random effects
378	model indicated unexplained variation due to the effect of site (i.e. cave/mine). Greater
379	than 20 samples would be needed to have a mean estimated probability of detection
380	≥0.95.
381	
382	
383	DISCUSSION
384	
385	The sudden emergence and spread of WNS in North America has led to speculation that
386	G. destructans is an exotic species and may have been recently introduced from Europe
387	(8,9,11). If this hypothesis is valid, the distribution of <i>G. destructans</i> would be expected
388	to mirror that of the disease. We screened a total of 295 sediment samples collected from
389	55 caves and mines in the eastern U.S. using a real-time PCR assay specific for G .
390	destructans (17) and detected the fungus in 17 bat hibernacula. All 17 of these sites were
391	situated within the known range of WNS at the time the samples were collected, and G.

392	destructans was not found to occur outside that area (Fig. 1). Furthermore, the real-time
393	PCR findings paralleled WNS manifestation on a temporal scale. G. destructans was not
394	detected in three sites that were unaffected by WNS in the winter of 2008-2009, but the
395	fungus was later detected in sediment samples collected from those same hibernacula
396	subsequent to the appearance of the disease in bats at those sites. These findings suggest
397	that an endemic, less virulent strain of G. destructans likely did not occur in eastern
398	North America prior to arrival of WNS and offers further support for the exotic species
399	hypothesis to explain the emergence of G. destructans as a novel pathogen in North
400	America (8,13).

402 Sixteen of the 17 sites in which G. destructans was detected by real-time PCR in this 403 study contained bats showing signs of WNS prior to, or at the time of, sample collection. 404 In the remaining site, WNS was observed the following winter. While these results 405 would seem to suggest that WNS may be an inevitable outcome once G. destructans is 406 introduced into a hibernaculum, it is important to interpret these results cautiously 407 because a relatively small number of bat hibernacula were sampled, and all positive sites 408 were located within the same geographic area. Thus, it is unknown how clinical signs of 409 WNS and disease severity may vary as G. destructans spreads to new regions of North 410 America with different environmental conditions and host species. The potential 411 importance of site-specific factors in their relation to WNS may be highlighted by the 412 detection of G. destructans in only one of the four buffer sites, which could suggest that 413 different hibernacula have different latency periods between the arrival of G. destructans

415

416 417 This study represents the first application of a high-throughput PCR technique for 418 directly detecting G. destructans in the environment. A previously described PCR assay 419 (16) utilized in a prior study to detect G. destructans in the environment lacked 420 specificity and required cloning and sequencing procedures to differentiate DNA of G. 421 *destructans* from that of other closely related *Geomyces* spp. common in cave sediment 422 (15,21). Additionally, the real-time PCR assay that targets the Alpha-L-Rhamnosidase 423 gene of G. destructans (30) was not tested against environmental samples, but may lack 424 the sensitivity necessary to detect the fungus in sediment given that the Alpha-L-425 Rhamnosidase gene likely exists at a low copy number within the genome of G. 426 destructans (17). The work described herein confirms the specificity and sensitivity of a 427 previously developed PCR method that targets the IGS region of G. destructans (17) and 428 supports its application for use with environmental samples. Furthermore, the detection 429 of the fungus in a buffer site suggests that PCR screening of sediment samples within 430 caves may allow for early detection of G. destructans prior to manifestation of visible 431 signs of disease in bats inhabiting a hibernaculum. The relatively low estimated detection 432 probability for samples collected from buffer sites relative to diseased sites may have 433 been an effect of our inability to estimate infection status and detection probability 434 separately for buffer and infected sites. Thus, we may have underestimated detection 435 probability by overestimating the proportion of buffer sites that were infected. 436 Additionally, the probability of detecting the fungus at buffer sites prior to disease onset

and the manifestation of the disease in bats, or that fungal abundance thresholds that

result in the appearance of WNS (i.e. infective doses of the fungus) vary between sites.

438 months before a hibernaculum might become diseased as opposed to a full year before 439 manifestation of WNS). Also, future work to determine whether certain types of 440 environmental samples or specific locations within caves and mines are more likely to 441 harbor G. destructans may further enhance sensitivity of detection. 442 443 While using PCR to detect G. destructans can provide important information about 444 certain aspects of WNS disease ecology, the method is limited in that it cannot 445 discriminate between viable and non-viable fungus. This is of particular importance in 446 determining what role the environment plays in maintaining infectious populations of G. 447 destructans. Detection of live G. destructans in 7 of the 14 caves and mines in late 448 summer provides the first evidence that G. destructans is capable of surviving in bat 449 hibernacula when bats are either absent or at low densities and that caves and mines serve 450 as likely infection sources when bats return for hibernation in early autumn. However, 451 our ability to culture viable G. destructans from sediment samples collected in 2011 from 452 sites C5 and C9 suggests the fungus can survive much longer than a few months in the 453 environment in the absence of a bat host. Specifically, bats had not been observed in one 454 of the sites (C9) for approximately one year prior to sample collection, and site C5 had 455 been sealed such that bats were excluded from the hibernaculum for approximately two 456 years prior to sample collection. Demonstration that sediments from these two mines 457 contained live G. destructans one to two years after bats had been extirpated/excluded,

might be enhanced by collecting sediment samples in early fall instead of mid-winter (i.e.

458 indicates the fungus can persist long-term in caves and mines.

459

460	The culture technique used for this experiment lacked the sensitivity of the molecular
461	detection technique for sites known to be infested with G. destructans. Specifically, there
462	was a lack of correlation in detecting viable G. destructans across replicate, serially-
463	diluted, and spatially- and temporally-separated samples collected within the same sites.
464	The mean probability of detecting G. destructans from contaminated sediment was 0.14
465	with at least 20 samples required from an average site to have a 95% chance of detecting
466	the fungus using the described culture technique. However, detection probabilities varied
467	greatly by site with some sites still not reaching a 50% detection probability with 15
468	samples. Clumping or aggregation of G. destructans within sediment, competition or
469	inhibition by other fungi on the artificial culture medium, low abundance of G .
470	destructans in environmental samples relative to other fungi, differences in abundance of
471	G. destructans between sites, and/or differences in abundance between locations within
472	the same site may account for these discrepancies. Whatever the reason, the described
473	culture-based technique is valuable to demonstrate that viable G. destructans is present in
474	a tested sample. However, the technique is currently neither suitable for quantifying
475	abundance of G. destructans nor for proving the absence of the fungus in environmental
476	samples. Future work focusing on developing a medium that is more selective for G .
477	destructans may serve to improve the utility of culture-based methods for addressing
478	research questions such as how long the fungus remains viable in different environments,
479	what portions of hibernacula are most conducive to supporting G. destructans (including
480	cave ceilings where bats roost and therefore may be most likely to come into contact with
481	the fungus), whether G. destructans can propagate (as opposed to simply persist) in

482 hibernacula without bats, and how abundance of the fungus changes spatially or

483 temporally within sites.

484

485 Disease ecology is often represented by a triad that involves interactions between a host, 486 a pathogen, and an environment. To date, research on WNS has focused primarily on 487 bats, G. destructans, and interactions between the two. Relatively little information is 488 available regarding the interplay between the pathogen and the environment. This work 489 demonstrates the utility of environmental sampling for enhancing WNS surveillance and 490 furthering research on WNS epidemiology. Specifically, the results of this study show 491 that presence of G. destructans in environments where bats hibernate is strongly 492 correlated with disease manifestation; the fungus may be detectable in the environment 493 prior to disease manifestation; and the fungus can persist in the sediment of bat 494 hibernacula for long periods of time in the absence of bat hosts. Additional studies to more fully elucidate the role the environment plays in supporting proliferation of G. 495 496 destructans and facilitating the development and progression of WNS will reveal 497 important factors related to the epidemiology of WNS and may provide information 498 useful in WNS disease management. 499 500 501 REFERENCES 502

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657

658 FIGURES AND TABLES

659

664

Fig. 1. Map of the eastern U.S. showing the sampling locations described in Table 1.

The left halves of the circles depict the disease status of individual bat hibernacula at the time the samples were collected (red = diseased, blue = buffer, yellow = clean). The right halves of the circles represent the PCR results (red = G. destructans detected, yellow = G.

destructans not detected). The solid white line marks the general geographic range of

665	WNS (i.e. WNS zone) at the time of sample collection; the dotted white line marks the
666	general geographic range of WNS during the following winter (i.e. buffer zone) (see
667	http://www.whitenosesyndrome.org/resources/map). Geomyces destructans was only
668	detected in bat hibernacula that were situated within the known range of WNS as of the
669	winter of 2008-2009. The PCR results for sites 20, 21, 22, and 23 only represent those
670	from samples collected in the winter of 2008-2009 and not those collected in the winter
671	of 2010-2011.
672	

Table 1. Bat hibernacula in the eastern U.S. from which sediment samples were collected

674 in the winter of 2008-2009 to test for the presence of *G. destructans* by real-time PCR.

Site Designation	State	Geographic Zone	Cave/Mine Status	PCR Result
1	NH	WNS	diseased	+
2	NH	WNS	diseased	1
$\frac{2}{3}$	VT	WNS	diseased	-
4	VT	WNS	buffer	+
5	VT	WNS	diseased	+
6	MA	WNS	diseased	+
0 7	MA	WNS	diseased	+
8	MA	WNS	diseased	+
9	CT	WNS	diseased	+
10	NY	WNS	diseased	+
11	NY	WNS	diseased	+
12	NY	WNS	diseased	+
13	NY	WNS	diseased	+
14	PA	WNS	diseased	+
15	PA	buffer	clean	_
16	PA	buffer	clean	_
17	PA	buffer	clean	-
18	WV	WNS	clean	-
19	WV	WNS	diseased	+
20	WV	WNS	diseased	-
20*	WV	WNS	diseased	+
21	WV	WNS	clean	-
21*	WV	WNS	diseased	+
22	WV	WNS	buffer	-
22*	WV	WNS	diseased	+
23	VA	WNS	buffer	-
23*	VA	WNS	diseased	+
24	VA	WNS	buffer	-
25	VA	buffer	clean	-
26	NC	outside	clean	-
27	NC	outside	clean	-
28	GA	outside	clean	-
29	FL	outside	clean	-
30	FL	outside	clean	-
31	AL	outside	clean	-
32	AL	outside	clean	-
33	AL	outside	clean	-

34	TN	outside	clean	-
35	TN	buffer	clean	-
36	TN	buffer	clean	-
37	KY	outside	clean	-
38	KY	outside	clean	-
39	KY	outside	clean	-
40	KY	outside	clean	-
41	OH	outside	clean	-
42	OH	outside	clean	-
43	OH	outside	clean	-
44	IN	outside	clean	-
45	IN	outside	clean	-
46	IL	outside	clean	-
47	IL	outside	clean	-
48	IL	outside	clean	-
49	AR	outside	clean	-
50	MO	outside	clean	-
51	WI	outside	clean	-
52	WI	outside	clean	-
53	WI	outside	clean	-
54	MI	outside	clean	-
55	MN	outside	clean	-

677 * Sites re-sampled in winter 2010-2011; "geographic zone," "cave/mine status," and

678 "PCR result" depict results from winter 2010-2011. Sites with the same number, but

679 lacking this symbol represent results from winter 2008-2009 for these same sites.

680 Table 2. Bat hibernacula within the WNS-affected area of the U.S. in which sediment

samples were tested for the presence of viable *G. destructans* using a culture technique.

			Culture Result			
Site Designation	State	Sampling Location	Winter 2010-11	Summer 2011	Winter 2011-12	
C1	VA		-	+	+	
		1	-	-	-	
		2	-	+	-	
		3	-	-	-	
		4	-	-	+	
		5	-	-	-	
C2	VA		-	-	-	
		1	-	-	-	
		2	-	-	-	
		3	-	-	-	
		4	-	-	-	
		5	-	-	-	
C3	VA		-	-	-	
		1	-	-	-	
		2	-	-	-	
		3	-	-	-	
		4	-	-	-	
		5	_	-	-	
C4	VT		+	+	+	
		1	-	+	-	
		2	+	-	-	
		3	+	-	-	
		4	-	-	+	
		5	+	+	-	
C5	VT		+	-	+	
		1	+	-	+	
		2	-	-	-	
		3	-	-	+	
		4	-	-	+	
		5	-	-	-	
C6	NH		+	+	n/a	
		1	+	-	n/a	
		2	-	+	n/a	
		3	-	-	n/a	
		4	-	-	n/a	

		5	-	-	n/a
C7	NH		+	-	n/a
		1	-	-	n/a
		2	+	-	n/a
		3	-	-	n/a
		4	-	-	n/a
		5	_	-	n/a
C8	NH		+	+	n/a
		1	-	+	n/a
		2	+	+	n/a
		3	+	-	n/a
		4	-	-	n/a
		5	-	-	n/a
C9	NH		-	+	+
		1	-	-	-
		2	-	-	-
		3	-	-	+
		4	-	+	+
		5	-	-	-
C10	WV		-	-	-
		1	-	-	-
		2	-	-	-
		3	-	-	-
		4	-	-	-
		5	-	-	-
C11	WV		-	+	-
		1	-	+	-
		2	-	-	-
		3	-	-	-
		4	-	-	-
		5	-	-	-
C12	WV		+	-	-
		1	-	-	-
		2	+	-	-
		3	-	-	-
		4	-	-	-
		5	-	-	-
C13	WV		+	-	-
		1	-	-	-
		2	-	-	-
		3	-	-	-
		4	+	-	-

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			5	-	-	-
	C14	WV		+	+	-
			1	-	-	-
			2	+	-	-
			3	-	-	-
			4	-	+	-
			5	-	-	-
-						

683 n/a No sample collected

