RESEARCH ARTICLE



Phylogenetics of a Fungal Invasion: Origins and Widespread Dispersal of White-Nose Syndrome

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ABSTRACT Globalization has facilitated the worldwide movement and introduction of pathogens, but epizoological reconstructions of these invasions are often hindered by limited sampling and insufficient genetic resolution among isolates. Pseudogymnoascus destructans, a fungal pathogen causing the epizootic of white-nose syndrome in North American bats, has exhibited few genetic polymorphisms in previous studies, presenting challenges for both epizoological tracking of the spread of this fungus and for determining its evolutionary history. We used single nucleotide polymorphisms (SNPs) from whole-genome sequencing and microsatellites to construct high-resolution phylogenies of P. destructans. Shallow genetic diversity and the lack of geographic structuring among North American isolates support a recent introduction followed by expansion via clonal reproduction across the epizootic zone. Moreover, the genetic relationships of isolates within North America suggest widespread mixing and long-distance movement of the fungus. Genetic diversity among isolates of P. destructans from Europe was substantially higher than in those from North America. However, genetic distance between the North American isolates and any given European isolate was similar to the distance between the individual European isolates. In contrast, the isolates we examined from Asia were highly divergent from both European and North American isolates. Although the definitive source for introduction of the North American population has not been conclusively identified, our data support the origin of the North American invasion by P. destructans from Europe rather than Asia.

IMPORTANCE This phylogenetic study of the bat white-nose syndrome agent, *P. destructans*, uses genomics to elucidate evolutionary relationships among populations of the fungal pathogen to understand the epizoology of this biological invasion. We analyze hypervariable and abundant genetic characters (microsatellites and genomic SNPs, respectively) to reveal previously uncharacterized diversity among populations of the pathogen from North America and Eurasia. We present new evidence supporting recent introduction of the fungus to North America from a diverse Eurasian population, with limited increase in genetic variation in North America since that introduction.

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This article is a direct contribution from a Fellow of the American Academy of Microbiology. Solicited external reviewers: Matthew Fisher, Imperial College London; Jason Stajich, University of California, Riverside. **KEYWORDS** Chiroptera, *Pseudogymnoascus destructans*, emerging infectious disease, epizootic, microsatellite, whole-genome sequencing, wildlife

ungal diseases are emerging as major threats to ecosystem integrity at local, ľ regional, and global levels (1). For example, introduced fungal phytopathogens such as Cryphonectria parasitica (agent of chestnut blight) and Cronartium ribicola (agent of white pine blister rust) have caused dramatic declines in foundation tree species of American forests, with severe implications for regional forest structure and food webs (2, 3). Several fungal pathogens of vertebrates have also recently emerged and are of major conservation concern for wildlife, including Batrachochytrium dendrobatidis and Batrachochytrium salamandrivorans (agents of amphibian chytridiomycosis), as well as Ophidiomyces ophiodiicola (agent of snake fungal disease) (4–6). In particular, the ongoing chytridiomycosis panzootic ranks among the most devastating wildlife diseases due to its broad host range, high virulence, and rapid worldwide invasion (7). In 2006, another severe fungal epizootic, white-nose syndrome (WNS), emerged among hibernating bats in upstate New York in the United States. This disease was named for the white growth of the psychrophilic fungus Pseudogymnoascus destructans on the skin, especially muzzles, of infected bats (8). The pathogen has colonized nearly all surveyed bat hibernacula in the eastern United States and Canada over the last 10 years (9, 10). Millions of bats have died as a result of this disease, and several species are threatened with extinction, including Myotis septentrionalis (northern long-eared bat), the once plentiful Myotis lucifugus (little brown bat), and the endangered Myotis sodalis (Indiana bat) (11, 12). Bats provide widespread insect suppression services to natural and agricultural ecosystems and are primary suppliers of nutrients to unique cave ecosystems through deposition of quano (13). Thus, this catastrophic reduction in the bat populations of North America will likely have pervasive ecologic repercussions.

Rachowicz et al. (14) proposed the novel and endemic pathogen hypothesis for the origin of *B. dendrobatidis* and emphasized differences in management and research priorities for pathogens of different origins. A novel pathogen population would be expected to have limited diversity compared to its source population, reflecting a recent introduction and demographic bottleneck of the introduced organism (14, 15). Conversely, a pathogen with greater diversity implies a longer natural history in an area and supports the endemic pathogen hypothesis. Many bacterial, viral, parasitic, and fungal diseases of epizoological importance have expanded clonally (16), and genetic analyses of *P. destructans* have been consistent with this pattern.

Pseudogymnoascus destructans primarily reproduces asexually, producing abundant haploid conidia (17, 18). Although it is theoretically possible for *P. destructans* to reproduce sexually via a heterothallic mating system, this has not yet been observed, and only one of two required mating types has to date been detected in the genomes of North American fungal isolates (19). Previous studies of the origin and spread of WNS revealed little or no genetic diversity in North American isolates of *P. destructans* (20, 21). However, these previous studies were limited by low discriminatory power. Only eight loci were sequenced, and they did not exhibit sufficient polymorphisms to identify the exact source population or to distinguish among isolates of a recently emerged pathogen during an infectious disease outbreak (22).

In contrast, other genetic markers such as microsatellites (23) and whole-genome single nucleotide polymorphism (SNP) genotyping (24) use many loci from throughout the genome to increase the resolution of genotypes. The high variability of microsatellite loci leads to more heterogeneity than point mutations in genes, often allowing for discrimination between closely related or recently diverged isolates. Furthermore, the hallmark of whole-genome SNP genotyping is the large number of characters (ranging from hundreds to hundreds of thousands of SNP loci in a typical analysis) that can be used to construct a robust phylogeny of clonal organisms (22).

We report the results of a high-resolution phylogenetic analysis of *P. destructans* using both microsatellite and whole-genome SNP loci to complement the benefits and

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minimize the drawbacks of each approach. Our analysis includes isolates from throughout the WNS epizootic zone in North America that were collected from 2008 to 2014. We also included isolates from Europe, Mongolia, and China (where *P. destructans* was recently found to be widely distributed [25]) to more thoroughly investigate the genetic diversity of putative origin populations of the North American population.

RESULTS

SNP phylogeny. Illumina paired-end whole-genome shotgun sequences were obtained for 26 isolates of P. destructans from the North American WNS epizootic zone from the years 2008 to 2014, 5 isolates from Europe, and 3 isolates from Asia. These reads, as well as publicly available P. destructans and near-relative Pseudogymnoascus and Geomyces species assemblies, were aligned to a reference genome from P. destructans 20631-21 (NCBI accession no. GCA_001641265.1) (26). Oidiodendron maius Zn (Ascomycota, Leotiomycetes) was used as an outgroup based on our phylogenetic analyses of fungi closely related to Pseudogymnoascus (data not shown). Descriptive statistics of the alignments, as well as NCBI accession numbers of all samples analyzed, are displayed in Table 1. Duplicated/repeat regions accounted for 43.25% of the genome and were excluded from our SNP-based analyses. Only ~0.1% of the reference genome was orthologous in all samples, resulting in 4,757 SNP loci, of which 2,479 were synapomorphic (i.e., shared between one or more samples and useful to define clades). Maximum parsimony analysis resulted in four trees with a low rescaled consistency (RC) index (RC of 0.4704), indicating a high degree of homoplasy that one would expect from a data set representing a long evolutionary history (data not shown). Maximum likelihood analysis (Fig. 1) indicated strong bootstrap support for a single P. destructans clade that is sister to other all currently recognized members of the genus Pseudogymnoascus, including ancient samples from Russian permafrost. Moreover, in this analysis, isolate JH15CN0111a from China is basal to P. destructans isolates from Mongolia, Europe, and North America, allowing us to root the *P. destructans* phylogeny (Fig. 2).

As previously determined by multilocus sequence typing (MLST) (27), members of the genus Pseudogymnoascus are not all closely related to each other. We thus limited the sample set to just P. destructans and aligned only reads from this species to identify SNP loci shared specifically among P. destructans isolates. This resulted in the detection of 37,752 core SNP loci within the 20-Mb core genome alignment. Maximum parsimony produced one tree with an RC of 0.7729 (data not shown). Maximum likelihood analysis (Fig. 2) distinguishes three well-supported clades among the *P. destructans* isolates: China, Mongolia, and Europe. The North American isolates are nearly identical in this tree and are therefore represented as a group by the type isolate 20631-21, obtained in New York in 2008. This phylogeny clearly shows the North American clade of P. destructans as a member of a greater European clade and distinct from the eastern and central Asian isolates. The tree also shows that North American P. destructans isolate is most closely related to a Ukrainian isolate (Gd44) in our sample set and least related to an isolate from France (Gd41). This is in contrast to a previous MLST study (28), which showed the opposite trend using many of the same isolates. However, low sample sizes, long branch lengths, and poor bootstrap support for the branches between members of the European clade preclude stronger conclusions about the region of Europe that gave rise to North American P. destructans. With our current data set, the possibility that North American P. destructans was introduced from Asia, however, seems unlikely.

We reduced the sample set once again to include only isolates of *P. destructans* from North America and reanalyzed the alignments for SNPs. Only 51 core SNP loci were shared among the 26 North American isolates. Maximum parsimony produced one tree with an RC of 1.00 (data not shown). Thus, all homoplasy observed among *P. destructans* phylogenies was due to the Eurasian genomes. Only three SNPs were synapomorphic. Two SNPs show differences between the type isolate 20631-21 and all other samples. A single SNP shared by samples 23414-1W and 23434-1W, both obtained in Indiana in 2011, suggest the formation of a geographically isolated lineage. Less

TABLE 1 Whole-genome sequences aligned to the reference genome of <i>Pseudogymnoascus destructans</i> (NCBI accession no.
GCA_001641265.1)

					Read length	Avg read	% of core genome	
lsolate ID no.	Species ^a	Source ^b	Location ^c	Date	(bp)	depth	covered ^d	Accession no.
20631-21	P. destructans	M. lucifugus	NY	Feb 2008	NA ^e	NA	100.00	GCA_001641265.
GU999986	P. destructans	M. myotis	DEU	Mar 2009	2×10^{1}	155.83	98.73	SRR6011467
GU350433	P. destructans	M. myotis	CHE	Apr 2009	2×10^{1}	187.97	98.93	SRR6011468
GU350434	P. destructans	M. myotis	HUN	Mar 2009	2×10^{1}	169.29	99.00	SRR6011465
20693-1	P. destructans	M. lucifugus	MA	Mar 2008	2×10^{1}	206.39	99.79	SRR6011466
22429-8	P. destructans	M. septentrionalis	WV	Jan 2009	2×10^{1}	208.62	99.81	SRR6011471
22971-3	P. destructans	M. lucifugus	ON	Mar 2010	2×10^{1}	123.24	99.73	SRR6011472
22504-1	P. destructans	M. lucifugus	PA	Mar 2009	2×10^{1}	192.73	99.67	SRR6011469
22442-2	P. destructans	M. lucifugus	NJ	Feb 2009	2×10^{1}	200.79	99.71	SRR6011470
22948-1	P. destructans	M. septentrionalis	TN	Mar 2010	2×10^{1}	180.47	99.49	SRR6011473
20674-9	P. destructans	M. septentrionalis	VT	Mar 2008	2 imes 98	100.25	99.46	SRR6011474
20682-10	P. destructans	M. septentrionalis	MA	Mar 2008	2×10^{1}	156.00	99.11	SRR6011477
22004-1	P. destructans	M. lucifugus	CT	Apr 2008	2×10^{1}	213.78	99.20	SRR6011478
22426-2	P. destructans	M. lucifugus	CT	Jan 2009	2×10^{1}	170.49	99.42	SRR6011475
22469-1	P. destructans	P. subflavus	VA	Mar 2009	2×10^{1}	167.09	99.42	SRR6011476
22480-1	P. destructans	E. fuscus	NY	Mar 2009	2 imes 98	85.85	99.43	SRR6011481
22884-4W	P. destructans	M. lucifugus	VT	Jan 2010	$2 imes 10^{1}$	141.88	99.43	SRR6011482
22930-2	P. destructans	P. subflavus	TN	Feb 2010	$2 imes 10^{1}$	82.98	99.03	SRR6011479
22949-4	P. destructans	M. septentrionalis	MD	Mar 2010	2×10^{3}	211.59	99.54	SRR6011480
22972-2W	P. destructans	M. lucifugus	ON	Mar 2010	2×10^{1}	82.48	99.03	SRR6011483
22997-1	P. destructans	M. septentrionalis	TN	Apr 2010	2×10^{1}	271.20	99.97	SRR6011484
23414-1W	P. destructans	M. lucifugus	IN	Jan 2011	2 imes 98	120.00	99.59	SRR6011493
23434-1W	P. destructans	M. lucifugus	IN	Jan 2011	2×10^{1}	255.07	99.96	SRR6011492
23444-1	P. destructans	M. lucifugus	TN	Feb 2011	2 imes 98	96.67	99.49	SRR6011491
23455-1	P. destructans	M. lucifugus	VA	Feb 2011	2×10^{1}	163.17	99.70	SRR6011490
Gd41	P. destructans	M. myotis	FRA	Mar 2009	2 imes 251	89.04	99.31	SRR6011497
Gd44	P. destructans	M. myotis	UKR	Feb 2011	2 imes 301	92.68	98.71	SRR6011496
23874-1	P. destructans	M. lucifugus	ME	Dec 2011	2×10^{1}	200.76	99.97	SRR6011495
23877-1	P. destructans	M. septentrionalis	DE	Mar 2012	2×10^{1}	177.32	99.97	SRR6011494
23897-2	P. destructans	P. subflavus	MO	Mar 2012	2 imes 251	119.80	99.93	SRR6011489
W41203	P. destructans	Sub	NB	Apr 2012	2 imes 251	74.31	99.54	SRR6011488
JH15CN0111a	P. destructans	M. petax	CHN	Mar 2015	2 imes 150	74.31	96.78	SRR6011485
JH16MG088	P. destructans	P. ognevi	MNG	2016	2 imes 150	78.84	97.31	SRR6011486
JH16MG093	P. destructans	P. ognevi	MNG	2016	2 imes 150	15.28	53.10	SRR6011487
03VT05	Pseudogymnoascus sp.	Sed	VT	2008	NA	NA	76.63	GCA_001662645.1
UAMH10579	P. verrucosus	Peat	AB	2002	NA	NA	76.92	GCA_01662655.1
05NY08	Pseudogymnoascus sp.	Sed	NY	2008	NA	NA	76.68	GCA_001662605.1
23342-1-l1	Pseudogymnoascus sp.	P. subflavus	WI	2008	NA	NA	61.20	GCA_001662575.1
24MN13	Pseudogymnoascus sp.	Sed	MN	2008	NA	NA	69.28	GCA_001662595.1
WSF3629	Pseudogymnoascus sp.	Peat	WI	1960	NA	NA	77.02	GCA_001662585.1
F-103	Pseudogymnoascus sp.	Soil	NY	Contemporary	NA	NA	76.76	GCA_000750895.1
F-3557	Pseudogymnoascus sp.	Pine post	SWE	Contemporary	NA	NA	52.04	GCA_000750665.1
F-3775	Pseudogymnoascus sp.	Soil	DEU	Contemporary	NA	NA	50.60	GCA_000750715.1
F-3808	Pseudogymnoascus sp.	M. glareolus	RUS	Contemporary	NA	NA	52.51	GCA_000750675.1
F-4246	Pseudogymnoascus sp.	Sed	MNG	Contemporary	NA	NA	53.77	GCA_000750735.1
F-4281	Pseudogymnoascus sp.	Cryopeg	RUS	0.12–0.2 mya ^f	NA	NA	57.09	GCA_000750745.1
F-4513	Pseudogymnoascus sp.	Permafrost	RUS	1.8–3.0 mya	NA	NA	54.09	GCA_000750755.1
-4514	Pseudogymnoascus sp.	Permafrost	RUS	1.8–3.0 mya	NA	NA	51.80	GCA_000750795.1
-4515	Pseudogymnoascus sp.	Permafrost	RUS	1.8–3.0 mya	NA	NA	61.24	GCA_000750805.1
F-4516	Pseudogymnoascus sp.	Permafrost	RUS	1.8–3.0 mya	NA	NA	53.86	GCA_000750815.1
-4517	Pseudogymnoascus sp.	Permafrost	RUS	1.8–3.0 mya	NA	NA	60.86	GCA_000750875.1
-4518	Pseudogymnoascus sp.	Soil	RUS	Contemporary	NA	NA	67.47	GCA_000750925.7
-4519	Pseudogymnoascus sp.	Soil	RUS	Contemporary	NA	NA	77.04	GCA_000750935.7
-4520	Pseudogymnoascus sp.	Soil	RUS	Contemporary	NA	NA	67.01	GCA_000750935.
ATCC 16222	P. pannorum	Soil	DEU	NA	NA	NA	58.08	GCA_001630605.7
M1372	P. pannorum	Soil	CA	Jun 1961	NA	NA	58.55	GCA_000497305.
Zn	O. maius	NA	NA	NA	NA	NA	0.29	GCA_000827325.7

^aThe species shown are *Pseudogymnoascus destructans*, *Pseudogymnoascus verrucosus*, *Pseudogymnoascus sp.*, *Pseudogymnoascus pannorum*, and *Oidiodendron maius*. ^bThe species shown are *Myotis myotis*, *Myotis lucifugus*, *Myotis septentrionalis*, *Perimyotis subflavus*, *Eptesicus fuscus*, *Myotis petax*, *Plecotus ognevi*, and *Myodes glareolus* (bank vole). Sub, hibernaculum substrate (i.e., roosting surface); Sed, hibernaculum sediment.

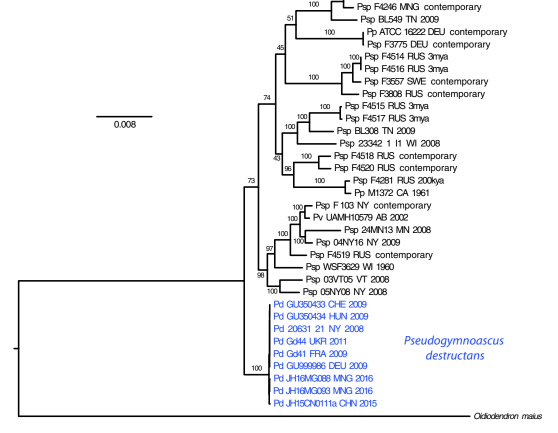
^cNY, New York; DEU, Germany; CHE, Chechnya; HUN, Hungary; MA, Massachusetts; WV, West Virginia; ON, Ontario, Canada; PA, Pennsylvania; NJ, New Jersey; TN, Tennessee; VT, Vermont; CT, Connecticut; VA, Virginia; MD, Maryland; IN, Indiana; FRA, France; UKR, Ukraine; ME, Maine; DE, Delaware; MO, Missouri; NB, New Brunswick; CHN, China; MNG, Montenegro; AB, Alberta; WI, Wisconsin; MN, Minnesota; SWE, Sweden; RUS, Russia; CA, California.

 d Coverage reported as percentage of core genome (i.e., portion of genome shared by all samples that is not in a duplicated region) that passed 10× minimum depth of coverage and 90% of read agreement filters.

^eNA, not available.

^fmya, million years ago.





Psp F4513 RUS 3mva

100 100

FIG 1 Maximum likelihood phylogenetic tree of the genus Pseudogymnoascus (previously Geomyces), based upon 4,757 SNPs (of which 2,479 are synapomorphic). Branches are labeled with bootstrap support values. Oidiodendron maius serves as outgroup. Pseudogymnoascus destructans is a sister clade to other members of the genus. Psp, Pseudogymnoascus sp.; Pd, P. destructans; Pp, P. pannorum; Pv, P. verrucosus. Additional data for each sample include isolate name, country/state of origin, and date of collection (estimated as preservation date for samples taken from permafrost). Samples without a precisely known collection date are indicated as "contemporary."

conservative filtering steps for SNP quality and read depth and removing the requirement for a locus to be present in all genomes failed to substantially increase the number of synapomorphies. Of the 51 SNP loci, none of the mutations were in coding sequences (Table 2), but rather were from intergenic sequences, introns, or untranslated regions (UTRs) of exons.

Isolate mating type was determined from whole-genome sequence alignments to the reference genome sequence of type isolate 20631-21, which was previously determined to mating type MAT1-1 (19). All isolates from North America in this study possessed the MAT1-1 mating type. Most Eurasian isolates did as well, although an isolate from Switzerland (GU350433) and an isolate from Mongolia (JH16MG088) were mating type MAT1-2.

Microsatellite phylogeny. Insufficient synapomorphic SNPs were discovered among the North American P. destructans whole-genome sequences to produce an informative phylogeny, such that analyses produced only a star-shaped phylogeny (inset in Fig. 3). Therefore, 96 North American isolates of P. destructans, including those analyzed by whole-genome sequencing, were genotyped with a 23-locus microsatellite panel (see Table S1 in the supplemental material). Thirty-one unique genotypes were identified, with one dominant genotype for 13 of the isolates. A phylogeny was constructed by the neighbor-joining (NJ) method from the microsatellite data (Fig. 3). These data, although based upon fewer genetic loci, contain many more samples and reveal far more genetic diversity than the SNP-based phylogeny of North American

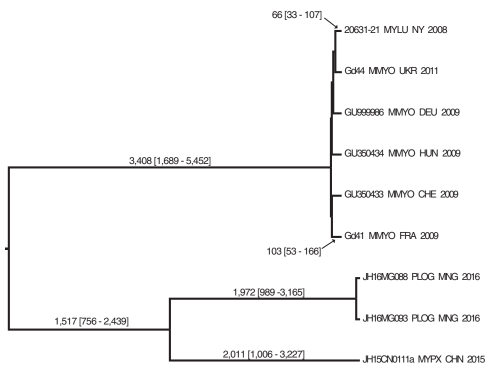


FIG 2 Maximum clade credibility tree for isolates of *Pseudogymnoascus destructans* from North America (n = 1), Europe (n = 5), and Asia (n = 3) based upon 37,752 core SNP sites (of which 22,594 are synapomorphic). All nodes had 100% posterior probabilities. Nodes are labeled with time (years before present), followed by the 95% high-probability density interval in brackets; dates on all shorter branches are not shown. North American isolates are nearly indistinguishable at these SNP loci, and so were represented in this tree by a single isolate (20631-21). The tree is rooted on the Chinese isolate based upon the Fig. 1 phylogeny that shows other *Pseudogymnoascus* spp. as a sister clade to *P. destructans*. Data for each sample include isolate name, bat host species code, country/state of origin, and date of collection.

P. destructans. This tree supports the single clade containing isolates 23431-1W and 23434-1W from Indiana identified in the whole-genome phylogeny, but includes a number of other isolates that did not share synapomorphic SNPs with the Indiana isolates. Other clades are evident as well, but they do not seem to correspond to geographic regions or sampling dates. A Mantel test comparing microsatellite-based genetic distance to geographic distance between isolates showed no correlation (r = 0.016, n = 99 replicates, P = 0.37). We note, however, that despite the appearance of defined groupings throughout the tree, there was limited bootstrap support for all of the branches within this phylogeny (all bootstrap values were <70).

Population genetics. Indices of genetic diversity and linkage were calculated for SNP genotypes. Asian samples were left out of this analysis due to the low sample size. The Simpson's index of genetic diversity was reduced in North America ($\lambda = 0.690$, standard error [SE] = 0. 130) compared to Europe ($\lambda = 0.800$), the putative source population for *P. destructans*. Average genomic linkage, represented by standardized index of association (29), was close to zero for both populations. Among North American isolates, $\bar{r}_D = 0.0585$, whereas for Europe, $\bar{r}_D = 0.00812$. Thus, recombination does not appear to be a significant factor in either population's genetic structure, and *P. destructans* isolates in both regions are reproducing clonally. The evenness of our sampling was not sufficient, nor did our data conform to various model assumptions to produce reliable population structure estimates (using models such as STRUCTURE and SNMF) from our SNP or microsatellite data (30).

DISCUSSION

Phylogeography. The two genetic markers used in this study, SNPs and microsatellites, provide complementary information regarding the phylogeography of *P. de*-

TABLE 2 Annotation of core SNP loci from isolates of P. destructans from North America

Accession no.	Position	Туре	Variant isolate(s)	Protein ID no.	Product
KV441386.1	1202839	UTR ^a	20693-1_MA_2008	OAF63413.1	NAD synthase
KV441386.1	2276958	Intergenic	22948-1_TN_2010	NA ^b	NA
KV441395.1	783753	UTR	22948-1_TN_2010	OAF58958.1	Hypothetical protein
KV441396.1	546602	Intergenic	23444-1_TN_2011	NA	NA
KV441398.1	551631	UTR	23877-1_DE_2012	OAF57976.1	Hypothetical protein
KV441399.1	235966	Intergenic	23414-1W_IN_2011,	NA	NA
		5	23434-1W_IN_2011		
KV441400.1	640730	UTR	22442-2_NJ_2009	OAF57273.1	Hypothetical protein
KV441401.1	139816	Intergenic	22948-1_TN_2010	NA	NÁ
KV441402.1	72139	Intergenic	23897-2_MO_2012	NA	NA
KV441402.1	226767	UTR	22948-1_TN_2010	OAF56852.1	Hypothetical protein
KV441402.1	510412	Intergenic	20693-1_MA_2008	NA	NÁ
KV441404.1	262446	UTR	22442-2_NJ_2009	OAF56419.1	Hypothetical protein
KV441387.1	430984	Intergenic	 23874-1_ME_2012	NA	NA
KV441387.1	820735	UTR	22426-2_CT_2009	OAF62808.1	Hypothetical protein
KV441387.1	1825781	Intergenic	22948-1_TN_2010	NA	NÁ
KV441387.1	2082665	UTR	22442-2_NJ_2009	OAF62574.1	ESCRT-I complex subunit VPS28
KV441387.1	685734	Intergenic	22442-2_NJ_2009	NA	NA
KV441387.1	785423	Intergenic	23897-2_MO_2012	NA	NA
KV441387.1	54855	Intergenic	22971-3_ON_2010	NA	NA
KV441387.1	193542	Intergenic	22948-1_TN_2010	NA	NA
KV441387.1	253155	Intergenic	22442-2_NJ_2009	NA	NA
KV441387.1	77282	Intergenic	23455-1_VA_2011	NA	NA
KV441387.1	120236	Intergenic	23897-2_MO_2012	NA	NA
KV441388.1	775461	Intragenic	22972-2W_ON_2010	OAF61744.1	Hypothetical protein
KV441388.1	950597	Intron	23897-2_MO_2012	OAF61915.1	Hypothetical protein
KV441388.1	120202	UTR	22469-1_VA_2009	OAF62044.1	Hypothetical protein
KV441388.1	57283	UTR	20631-21_NY_2008	OAF61813.1	Hypothetical protein
KV441389.1	343287	Intron	23455-1_VA_2011	OAF61688.1	AGC/AKT protein kinase
KV441389.1	419134	Intergenic	23414-1W_IN_2011	NA	NA
KV441389.1	867065	UTR	22004-1_CT_2008	OAF61368.1	Actin cytoskeleton-regulatory complex protein end
KV441389.1	1173430	Intergenic	22971-3 ON 2010	NA	NA
KV441389.1	1173432	Intergenic	22971-3_ON_2010	NA	NA
KV441389.1	50002	UTR	22948-1_TN_2010	OAF61394.1	Hypothetical protein
KV441390.1	643943	Intergenic	23897-2_MO_2012	NA	NA
KV441390.1	659862	UTR	22948-1_TN_2010	OAF61216.1	Hypothetical protein
KV441390.1	861423	UTR	22948-1_TN_2010	OAF60933.1	Hypothetical protein
KV441390.1	1243398	UTR	23434-1W_IN_2011	OAF60771.1	Hypothetical protein
KV441390.1	1524359	UTR	22948-1_TN_2010	OAF60872.1	Hypothetical protein
KV441391.1	151986	UTR	20693-1_MA_2008	OAF60366.1	Hypothetical protein
KV441391.1	1184294	Intergenic	22442-2_NJ_2009	NA	NA
KV441391.1	712894	Intergenic	23414-1W_IN_2011	NA	NA

^aUTR, untranslated region of exon.

^bNA, not available.

structans. Whole-genome sequencing revealed three distinct populations of P. destructans among the isolates we tested: one consisting of European isolates that included WNS epizootic isolates from North America, another of Mongolian isolates, and a third represented by an isolate from China. The phylogeny of European isolates determined in this study with 58,707 SNP loci was consistent with that described previously using 14 SNPs in an eight-member multilocus sequence typing analysis (MLST) (28): specifically, the genetic diversity of P. destructans was 14% higher in Europe than in North America. In this study, we identified only 51 core SNPs among North American isolates, which clearly demonstrates the homogeneity of *P. destructans* population structure across the epizootic zone and is consistent with previous reports of clonal spread (20, 21, 28, 31, 32). North American isolates formed a monophyletic group but appear to be representative of diversity from Europe (i.e., the branch lengths between the European samples were comparable between the European and North American isolates). This, as well as the 99.995% reduction in core SNP loci when non-American isolates are removed, represents a loss of genetic diversity consistent with previous conclusions of a recent introduction of a novel pathogen (33, 34). Moreover, star-shaped phylogenies containing few synapomorphies are typical in recent pathogen introductions (35-37),

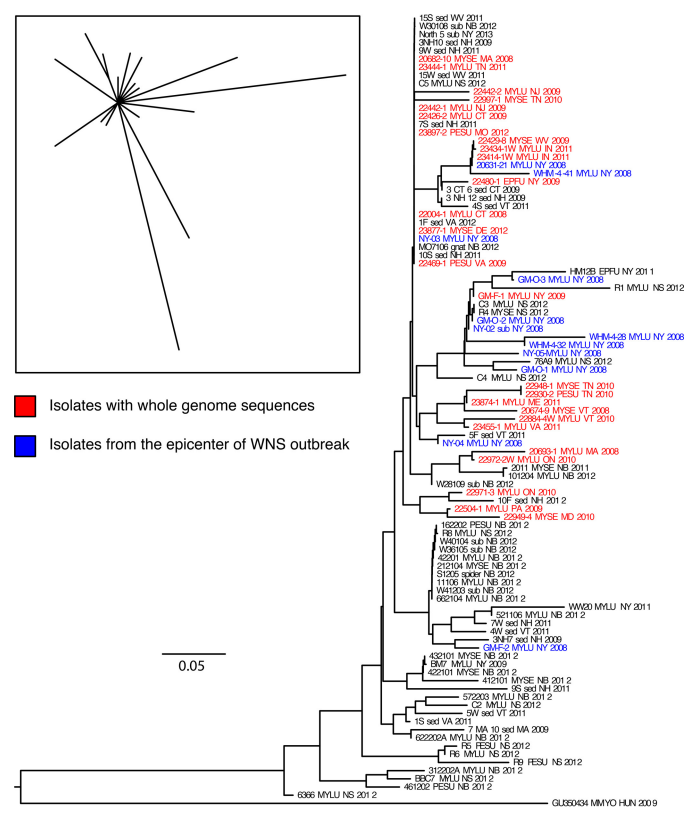


FIG 3 Phylogeny of 96 North American *Pseudogymnoascus destructans* isolates constructed from alleles of 23 microsatellite loci. Isolates that were included in whole-genome sequence phylogenies are colored red. The scale bar indicates genetic distance. Shown is a neighbor-joining tree of 96 North American isolates rooted with GU350434 (host, *M. myotis* [MMYO]; location, Hungary [HUN]; date, 2009). Samples from the epicenter of the WNS epizootic are colored blue. (Inset) Star-shaped maximum parsimony tree from comparisons of whole genomes, constructed from 51 SNPs, only 3 of which are synapomorphic.

with lack of rapid diversification into distinct lineages likely representing the norm rather than the rule. A recent report on the spread of WNS also identified few mutations of P. destructans upon introduction to North America (32). A strict interpretation of the dates in Fig. 2 would suggest that all P. destructans strains in North America and Europe diverged from each other within the past ~100 years and from the ancestor to Asian isolates roughly 3,400 years ago. While these estimates are plausible, we believe a more accurate estimation will require increased sampling of isolates from Eurasian populations. Perhaps analyses using P. destructans genomes from samples preserved long ago, as we did with Pseudogymnoascus sp. genomes from ancient Russian permafrost samples in Fig. 1, or detection of WNS-induced bat die-offs in Eurasia from the fossil record will enable more accurate dating. Given that P. destructans has likely been a bat pathogen for millions of years (73), the lack of mortality in European bats, considerable genetic diversity in microsatellite loci, and the broad distribution of the fungus across the palearctic—all changes that we envision taking millennia to evolve—it is likely that P. destructans is much older than a few thousand years old. Clearly more detailed molecular dating analyses are needed to resolve the question of the emergence and diversification of this species.

The microsatellite-based phylogeny containing more isolates illustrates some additional epizoological features with relevance to pathogen invasion and establishment. What is most striking about these data is the large amount of genetic diversity revealed among the North American isolates. Earlier studies indicated that 73 isolates of P. destructans from New York, Vermont, Pennsylvania, Ohio, West Virginia, North Carolina, Ontario, New Brunswick, Nova Scotia, and Prince Edward Island were genetically identical via MLST (20, 21, 31). Genotyping of these samples plus 78 others from eastern Canada and the United States resulted in a single genotype in 109 samples, plus two singleton genotypes from New Brunswick and one from Ontario. In contrast, microsatellite loci from our study defined 31 different genotypes among the isolates from North America. Another hallmark of these data is the lack of geographic or temporal correspondence between clades evident in the tree. For example, the Fig. 3 phylogeny shows that isolates collected from New York in 2008, at the beginning of the WNS outbreak, are distributed throughout the tree. Some isolates from the same sampling location and date (e.g., the Williams Hotel Mine) occur in different clades, indicating relatively high genetic diversity from this one region compared to other regions in North America. Unfortunately, these data do not indicate whether this diversity was present in the founding population or arose soon after introduction, a common problem with population genetic analyses of pathogens (38). New Brunswick and Nova Scotia isolates also occur in many clades associated with isolates from throughout the epizootic zone. A possible exception to the lack of geographic structuring we observed are several basal clades containing only northeastern Canadian isolates or isolates from New England and New York early in the outbreak. This region is relatively isolated and may represent an area that received P. destructans isolates geographically spreading from New England early in the outbreak and has had time since then to diversify genetically. Intense sampling in this region may also be contributing sampling bias to this observation, however.

Clonality. Questions regarding clonal expansion of *P. destructans* in North America can be answered more thoroughly by this study due to the large number of characters used in this data set. The strongest evidence for clonal reproduction of an ascomycete fungus is the lack of both mating types in a population, whereas an even distribution of mating types in a population would suggest the prevalence of sexual recombination. In a previous study (19), 5 out of 23 (22%) of isolates from Eastern Europe possessed the *MAT1-2* locus. With the addition of our samples from Eurasia, the proportion of MAT1-2 to MAT1-1 mating types is 7 out of 30 isolates (23%). In contrast, all 26 of the isolates from North America in this study possessed the MAT1-1 mating type. Additional analysis of *P. destructans* mating types in an eastern European population showed a more equal distribution of mating types (58.6% MAT1-1 and 41.3% MAT1-2; n = 41)

(39). The absence of the MAT1-2 mating type in North American *P. destructans* is evidence of both asexual reproduction and likely a recent genetic bottleneck.

The overrepresentation of widespread and persistent genotypes is also evidence of clonal reproduction as opposed to sexual reproduction (40), as was indicated by both SNP and microsatellite analyses in this study. Based upon SNP genotyping, 8 of the 26 isolates sequenced (31%) were found to be clonal, ranging from New England to Tennessee and sampled between 2008 and 2010. Similarly, two microsatellite genotypes account for 15 and 6 samples, respectively, out of a total of 55 North American isolates for which we obtained complete microsatellite genotypes (38%). Quantitatively, the Simpson's index of genetic diversity was slightly reduced in North America, although indexes of genetic diversity are highly dependent upon sample size, despite rarefaction, and the number of isolates from Europe compared to the number of isolates in North America analyzed in this study is very small. Average genomic linkage as measured by the standardized index of association indicates that both the European and North American populations of *P. destructans* are at linkage equilibrium, suggesting both populations reproduce primarily as clones.

Introduction of *P. destructans* **to North America.** Warnecke et al. (41) hypothesized that *P. destructans* was a novel pathogen introduced to North America from Europe based upon evidence that North American little brown bats developed WNS when inoculated with a European isolate of *P. destructans*. Their findings were further strengthened by Leopardi et al. (28), who demonstrated genetic similarity, based on MLST, between the North American and some European fungal populations, indicating the likely source population for this introduction to be from Europe. Subsequently, *P. destructans* has been isolated from bats throughout Europe (39, 42–46) and can also cause histopathological lesions consistent with WNS in European species of bats (47, 48). However, widespread mortality of bats from WNS has not been documented on the European continent (49). This suggests European bat species may have developed resistance or tolerance to infection by *P. destructans*, presumably due to coevolution between hosts and the fungal pathogen (41, 50). In contrast, the high mortality of North American bats infected with *P. destructans* is consistent with exposure of naive host species to a pathogen preadapted to similar hosts and environmental conditions.

Our work supports the hypothesis that *P. destructans* is a novel pathogen recently introduced to North America by demonstrating the relationship of North American isolates of the fungus to a population dominated by European isolates but genetically distant from isolates from Asia. Using high-resolution genetic markers, we document a loss of genetic diversity in epizootic *P. destructans* consistent with a recent introduction of this pathogen to North America.

Similarities to other fungal invasions. Pseudogymnoascus destructans follows a pattern of emergence and spread similar to other invasive fungal pathogens. In the case of Cryphonectria parasitica, the causative agent of chestnut blight (38), high genetic diversity in putative source populations coupled with a significant decrease in allelic richness in microsatellite and other multilocus genotypes among North American isolates suggested a founder effect associated with the introduction of the organism from Asia. North American isolates of C. parasitica were as genetically distant from Asian populations as the Asian populations were from each other. Analysis of population structure could narrow the source population down to a broad region (i.e., Honshu, Japan), but required the admixture of an unsampled population significantly different from the other Asian samples. A more detailed genomic analysis of an invading fungus was conducted with isolates of B. dendrobatidis originating from multiple continents and revealed a considerably more complex phylogeny of that fungus than was previously determined. A virulent panzootic strain apparently emerged and rapidly spread across the globe, yet the original source population of the lineage remains ambiguous despite a comprehensive sampling effort and thorough sequencing analyses (51). Consistent with a pattern also seen in our study, haplotypes from B. dendrobatidis did not correlate well with host or geography, suggesting other drivers of differentiation.

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Additionally, phylogenies constructed from different types of genetic markers yielded different, sometimes contradictory results for clones of the potato blight agent, *Phytophthora infestans* (52). A common thread in all of these case studies is that more genetic information lends complexity, but not necessarily clarity, to the natural history of an invading fungal pathogen. In terms of the natural history of *P. destructans*, it will require extensive sampling in Europe to find the population that was introduced to North America.

Using two high-resolution genetic characters—SNPs within whole-genome sequences and microsatellites—we have demonstrated that isolates of P. destructans in North America form a single clade much more closely related to isolates from Europe than to genetically distant populations sampled in Asia. The North American isolates represent a loss of genetic diversity in comparison to isolates from Europe, adding further support to the hypothesis that P. destructans was recently introduced to North America from Europe (28). Future sampling in Europe will be needed to more precisely define the origin of *P. destructans*, although the substantial diversity of *P. destructans* in Europe, coupled with results from other fungal invasion studies, suggests that identification of the exact source population may be challenging. Microsatellite loci were particularly suitable for distinguishing isolates from throughout the North American WNS epizootic zone, but the few synapomorphic SNPs found among these isolates could not resolve their phylogenetic relationships. However, sufficient SNPs were present to differentiate isolates of P. destructans from Europe, China, Mongolia, and North America. This work elevates our understanding of the origin and spread of P. destructans to a similar level to that achieved for B. dendrobatidis and C. parasitica, providing a framework for a global pathogen population structure that can be used for future epizoological investigations.

MATERIALS AND METHODS

Sampling and culturing *P. destructans.* Skin samples from bats or swab samples of bats and substrates were cultured on Sabouraud dextrose agar or potato dextrose agar and incubated at 7 to 10°C, as previously described (42, 53).

Whole-genome sequencing. Genomic DNA was extracted from P. destructans by a variety of methods, including the OmniPrep for Fungi kit (G-Biosciences, St. Louis, MO), the DNeasy blood and tissue kit (Qiagen, Hilden, Germany) using the supplementary protocol from the manufacturer "Purification of total DNA from yeast using the DNeasy Blood and Tissue kit (DIY13 Aug-06)" as described previously (54), or a phenol-chloroform extraction. Preparation of libraries for Illumina whole-genome sequencing was based upon a previously published method (55) but was modified as sequencing technology evolved during the course of the project. In summary, approximately 1 to 10 μ g of DNA sample in 200 μ l Tris-EDTA buffer was sheared with a SonicMan microplate sonicator (Brooks Automation, Chelmsford, MA) to produce fragments 200 to 1,000 bp in length, with the average fragment length being 600 bp. The shearing protocol was as follows: 75 s at 0°C prechill, followed by 20 cycles of sonication for 10.0 s at full power, 75 s at 0°C lid chill, 10 s at 0°C plate chill, and 75 s at 0°C postchill. End repair, dA-tailing, adapter ligation, and indexing followed the standard Illumina protocol "Preparing samples for multiplexed paired-end sequencing" (part no. 1005361) using reagents from the NEBNext DNA library prep master mix set for Illumina (New England Biolabs, Ipswich, MA) or the KAPA highthroughput library preparation kit "with bead" (Kapa Biosystems, Wilmington, MA). Libraries were indexed with the Illumina multiplexing sample preparation oligonucleotide kit 6-bp indices (Illumina, San Diego, CA) or custom 8-bp indices for higher index read discrimination (56). Size selection of 600-bp fragments was accomplished by excision from 2% agarose gels followed by purification with a QIAquick gel purification kit (Qiagen, Hilden, Germany), E-Gel SizeSelect gel (Life Technologies, Inc., Grand Island, NY), or Agencourt AMPure XP SPRI beads (Beckman Coulter, Inc., Indianapolis, IN) using the Kapa Biosystems "High-throughput NGS library preparation technical guide, Illumina platforms" (57). Products from each step of the library preparation were purified with either Agencourt AMPure XP SPRI beads using the same protocol or the QIAquick 96 PCR purification kit (Qiagen, Hilden, Germany). Libraries were quantified via quantitative PCR with a KAPA library quantification kit (Kapa Biosystems, Wilmington, MA) for the ABI Prism 9600 real-time PCR system (Life Technologies, Inc., Grand Island, NY). Prior to loading on the sequencer, the library fragment size distribution was qualitatively confirmed with an Agilent DNA high-sensitivity kit for the 2100 Bioanalyzer (Agilent, Santa Clara, CA). Individual libraries were run on an Illumina GAIIx sequencer (V2 chemistry) or an Illumina MiSeq (V3 chemistry) to produce 100-bp paired-end reads or 250-bp paired-end reads, respectively. Libraries sequenced on a HiSeq 2000 were multiplexed with a maximum of five libraries per lane to produce 150-bp paired-end reads with V3 chemistry.

Whole-genome sequence analysis. The sequence from resequencing and assembly of the genome of *P. destructans* 20631-21, collected in New York in 2008 (26) (NCBI GenBank assembly accession no. GCA_001641265.1), was used as a reference for SNP discovery using the Northern Arizona SNP Pipeline

(NASP) version 1.0.2 (58). With NASP, Illumina paired-end reads were aligned to the reference genome with the Burrows-Wheeler Aligner (59) version 0.7.5a mem algorithm. SNPs were detected in the alignments with the Unified Genotyper of the Genome Analysis Toolkit (60) build 2.5-2-gf57256b following recommended best practices (61, 62). Duplicated regions of the reference genome, including repeat regions and multiple gene copies, were determined by aligning the reference sequence to itself using the nucmer algorithm of MUMmer version 3.23 (63). SNPs that fell within these duplicate regions were excluded from further analysis to avoid false SNP calls due to ambiguous read alignment. SNP loci were also excluded if at least one sample lacked a base call at an SNP locus), had read coverage of less than $10\times$ in at least one sample, and less than 90% of the reads agreed with the SNP call. Default settings for each software package were used unless otherwise noted. Mating type was determined from the prevalence of reads mapping to the *MAT1-1* mating type locus in read alignments to the reference sequence (NCBI GenBank accession no. KV441390.1, bases 187500 to 188500). SNPs were annotated with SnpEff version 4.3q (64).

Microsatellite genotyping. DNA samples from *P. destructans* were tested with the 23-locus multilocus variable number tandem-repeat analysis (MLVA) panel described by Drees et al. (65). Briefly, we identified 2- to 6-bp repeats in the genome sequence of *P. destructans* type strain 20631-21 (GenBank accession no. GL573169 to GL575015). We then tested 127 primer pairs for microsatellites with length polymorphisms using a diverse set of DNA extracts from 39 isolates of *P. destructans* collected from North America and Europe. We selected 23 loci containing repeat number polymorphisms. Using these primer sets, we conducted fragment analysis using the Terminator v3.1 cycle sequencing kit on a 3130xl Genetic analyzer (Life Technologies, Inc., Grand Island, NY). Allele sizes were determined using the LIZ 1200 size standard in GeneMapper version 4.0.

Phylogenetics. PAUP* version 4.0 a150 (66) was used to conduct maximum parsimony analysis on SNP data, and neighbor-joining analysis (based on mean character difference) on microsatellite data. Extended majority rule maximum likelihood phylogenetic trees were created from concatenated SNP sequences with RAxML version 8.2.7 (67) using the ASC_GTRGAMMA substitution model with ascertainment bias correction for each base character to account for invariant loci and the autoMRE bootstopping criterion with a maximum of 1,000 bootstraps. Trees were plotted with FigTree version 1.4.0 (A. Rambaut, 2012; http://tree.bio.ed.ac.uk/software/figtree/).

BEAST2 version 2.4.5 (68) was used to create maximum clade credibility trees from SNP locus data. Briefly, the SNP data were first filtered to remove SNPs within 50 bp of each other to minimize the effects of linkage on the analyses. Optimum base substitution models for SNP data were determined with the R package phangorn version 2.1.1 (69), resulting in the generalized time-reversible (GTR) model for the full analysis and the JC69 model for the North American isolates alone. Both analyses used a strict molecular clock and a coalescent constant population tree model. Skyline demographic models, which may be more appropriate for the expanding populations in North America as well as a relaxed log normal molecular clock, were attempted for comparison but failed to coalesce. Because only core SNPs were being analyzed, the analyses were corrected for ascertainment bias with a count of invariant bases in the nonduplicated alignments to the reference genome. The Markov chain Monte Carlo (MCMC) chain length was set to 10 million, with 10% burn-in and logging every 1,000 steps. Five chains were run, producing a total of 50,000 trees, which were combined and subsampled to 10,000 trees with LogCombiner version 2.4.3 and used to create an annotated maximum clade credibility tree with TreeAnnotator version 2.4.3.

Population genetics. Simpson's index of genetic diversity and index of association were calculated with the R package poppr version 2.3.0 (70) as follows. SNP allele information was preprocessed by determining genetic distances between individuals with bitwise.dist, calculating the threshold distance for clustering isolates with cutoff_predictor, and assigning isolates to genotypes with mlg.filter. Simpson's index of genetic diversity was determined for both SNP and microsatellite genotypes with diversity_ci with 1,000,000 bootstraps. North American isolates were rarefied to equal the number of European isolates in the SNP analysis, which enabled the comparison of estimated Simpson's indices between the populations but prevented the calculation of standard error for the European population from SNP allele data was conducted with samp.ia, whereas the standard index of association and one-sided permutation test were calculated directly for microsatellite genotypes with ia.

Bruvo's genetic distance (which incorporates the number of repeats at a locus rather than just differences) for North American microsatellite genotypes was determined with poppr's bruvo.dist function. A pairwise geographic distance matrix between sampling locations was created with the R package geosphere v.1.5-5 (71). The Mantel test was conducted with the R package ade4 v.1.7-5 (72). Significance for this test was determined with 99 replicates of a Monte Carlo permutation test.

Accession number(s). Accession numbers for the sequences of the isolates examined in this study are listed in Tables 1 and 2.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio .01941-17.

TABLE S1, XLS file, 0.1 MB.

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