Genetic characterization of chytrids isolated from larval amphibians collected in central and east Texas

Thomas L. Marshall a, b, Carlos R. Baca a, Decio T. Correa b, Michael RJ. Forstner a, Dittmar Hahn a, David Rodriguez a, *

a Department of Biology, Texas State University, 601 University Dr., San Marcos, TX, 78666, USA
b Department of Integrative Biology, University of Texas at Austin, 2415 Speedway, Austin, TX, 78712, USA

Abstract

Chytridiomycosis, an emerging infectious disease caused by the fungal pathogen Batrachochytrium dendrobatidis (Bd), has caused amphibian population declines worldwide. Bd was first described in the 1990s and there are still geographic gaps in the genetic analysis of this globally distributed pathogen. Relatively few genetic studies have focused on regions where Bd exhibits low virulence, potentially creating a bias in our current knowledge of the pathogen's genetic diversity. Disease-associated declines have not been recorded in Texas (USA), yet Bd has been detected on amphibians in the state. These strains have not been isolated and characterized genetically; therefore, we isolated, cultured, and genotyped Bd from central Texas and compared isolates to a panel of previously genotyped strains distributed across the Western Hemisphere. We also isolated other chytrids from east Texas not known to infect amphibians. To identify larval amphibian hosts, we sequenced part of the COI gene. Among 37 Bd isolates from Texas, we detected 19 unique multi-locus genotypes, but found no genetic structure associated with host species, Texas localities, or across North America. Isolates from central Texas exhibit high diversity and genetically cluster with BdGPL isolates from the western U.S. that have caused amphibian population declines. This study genetically characterizes isolates of Bd from the south central U.S. and adds to the global knowledge of Bd genotypes.

1. Introduction

Emerging infectious diseases (EIDs) — those that have recently increased in incidence, impact, virulence, geographic or host range, or have recently evolved — are a growing threat to human health and global biodiversity (Daszak et al., 2000, 2003). Over the past two decades, an increasing proportion of EIDs have been attributed to pathogenic fungi. These include powdery mildew on important crop plants, white-nosed syndrome in bats, snake fungal disease, colony collapse disorder in bees, and sea-fan aspergillosis in corals (Fisher et al., 2012, 2016). Despite their immense economic and ecological impact, the genetic diversity of many of these fungi is still poorly understood (Fisher et al., 2016). None of these pathogens have been as devastating in their impact as Batrachochytrium dendrobatidis (Bd), the aquatic chytrid that induces chytridiomycosis, an epidermal infection in amphibians (Lips, 2016).

Bd, a member of the Rhizophydiales, causes infections when its flagellated zoospores encyst in a host’s skin and develop into zoosporangia (Longcore et al., 1999; Letcher et al., 2006). It was first described and attributed to disease outbreaks in the late 1990s (Longcore et al., 1999) and is now believed to be responsible for declines in amphibian populations dating back at least to the 1970s (Lips et al., 2004). Bd is found on every continent except Antarctica, is known to have infected over 700 species of amphibians, and has caused mass mortality events including species extinctions in Central and northern South America, the western U.S., Australia, and Spain (Berger et al., 1998; Bosch et al., 2001; Lips et al., 2006; Vredenburg et al., 2010). However, certain amphibian species and geographic regions have seemingly been unaffected by chytridiomycosis, even where Bd is present (Lips, 2016).

Phylogenomic analyses of Bd have revealed multiple lineages that include the widespread global panzootic lineage (BdGPL) (Farrer et al., 2011; Rosenblum et al., 2013), which is implicated in
most disease outbreaks and may have substructure representing a temperate clade (BdGPL1) and a tropical clade (BdGPL2) (Schloegel et al., 2012; Jenkinson et al., 2016). Other lineages include the putatively endemic and less virulent BdCape (found in South Africa and Spain), BdCH from Switzerland (Farrer et al., 2011), and BdKorea (Bataille et al., 2013). Isolation efforts have also led to the discovery of a strain in Japan that may be host specific (Goka et al., 2009) and BdBrazil, which was thought to be endemic in the Atlantic Forest of Brazil (Schloegel et al., 2012; Rodriguez et al., 2014; Jenkinson et al., 2016). A more recent analysis of global Bd genomes found that BdCH groups with a hyperdiverse Korean lineage (BdAsia-1), and BdBrazil groups with another Asian lineage (BdAsia-2) and has been subsequently renamed BdAsia-2/BdBrazil (O’Hanlon et al., 2018).

Recently revealed diversity within the Bd clade, coupled with the fact that there are still several geographic gaps in our current knowledge of Bd genotypes (James et al., 2015), suggests additional diversity is yet to be discovered. However, relatively few studies have examined Bd genotypes from regions lacking population declines, potentially creating a bias in our knowledge of the pathogen’s genetic diversity (Bataille et al., 2013). It is likely that Bd’s origin can be traced to a region in which amphibians have co-evolved with the fungus and where severe population declines and extinctions are not occurring (James et al., 2015). While recent work has begun to explore such regions of potential origin (O’Hanlon et al., 2018), genotyping in additional understudied areas is imperative and is facilitated by isolation and culturing of the fungus (Bataille et al., 2013; Jenkinson et al., 2016).

In the U.S., BdGPL1 has shown patterns of introduction and spread (Padgett-Flohr and Hopkins, 2009) and has caused host declines in the Sierra Nevada of California (Wake and Vredenburg, 2008; Briggs et al., 2010). On the other hand, long-term persistence and a lack of chytridiomycosis associated amphibian declines or extinctions have been detected. Studies have confirmed the presence of Bd in the eastern (Saenz et al., 2010) and central (Gaertner et al., 2009a, 2009b, 2010, 2012) parts of the state, yet no strains from these regions have been genotyped or isolated thus far (James et al., 2015). While a lack of known Bd related population declines or extinctions in the area could be an indication that the pathogen has a long local history, other factors, such as climate or host resistance, may be responsible for attenuating disease outbreaks in the region. Studying the genetics of the fungus in Texas and its amphibian hosts is an important first step in understanding potential disease dynamics between amphibians and Bd in the region. Additionally, by genetically characterizing isolates from Texas and placing them in the context of a more broadly distributed panel of strains, we can begin to gain a more complete evolutionary picture of the Bd group, and perhaps a better understanding of the origins and distribution of its most virulent lineages. Therefore, the goal of this study was to isolate Bd from larval anurans in central and east Texas, genetically identify anuran hosts, and compare genetic variation among chytrid strains present in this region to those distributed across the Western Hemisphere.

2. Methods and materials

2.1. Host collection

We collected anuran larvae from ponds in Travis, Hays, Bastrop, Kaufman, and Houston counties in central and eastern Texas from December 2015 to April 2016 (collection permit: SPR-0102-191) and stored them in 50 ml Falcon tubes or new plastic containers with water from the collection site (Table 1). We transported all collected tadpoles, in Styrofoam containers kept cool with synthetic ice packs, to Texas State University where they were euthanized following animal care and use protocols IACUC2014116499 and IACUC2015121220. Each tadpole was given a unique identifier, and a tail clipping from each tadpole was preserved in 95% ethanol for DNA extraction.

2.2. Fungal isolation, culture, and cryopreservation

We excised the keratinized jaw sheaths and tooth rows from each tadpole and transferred them to 1% tryptone agar Petri dishes. We cleaned the mouthparts by dragging them through fresh agar and transferred them to 1% tryptone agar Petri dishes with 200 mg/ml penicillin-G and 200 mg/ml streptomycin sulfate for isolation of chytrids (Longcore, 2000). We monitored the plates for growth of zoosporangia and active zoospores for up to two weeks. Once enough active zoospores were observed, we removed a section of the agar containing active zoospores to tryptone broth (16 g tryptone in 1000 ml deionized water; autoclaved) to cultivate the isolate. Once there was a high density of active zoospores, we transferred a 1-ml aliquot of broth from each isolate to a 1.5-ml centrifuge tube. We used a cryoprotectant solution consisting of 80 ml tryptone broth, 10 ml dimethyl sulfoxide, and 10 ml fetal calf serum to preserve all chytrid isolates (Boyle et al., 2003). For each isolate, we briefly centrifuged (376×g) tryptone broth cultures in 1.5 ml cryotubes, discarded the supernatant, and added 600 µl of cryoprotectant. All isolates were given a serially numbered unique identifier (TXST – Texas State University) and were stored at –80 °C (Supplementary appendix 1).

2.3. Genetic methods

To extract DNA, we centrifuged 1 ml aliquots of live broth cultures, discarded the supernatant, and extracted DNA from the pellet using the Mammalian Tissue and Rodent Tail Genomic DNA Purification Protocol in the Thermo Scientific GeneJET Genomic DNA Purification Kit #K0722 (Thermo Fisher Scientific, Inc.). We determined extraction success with gel electrophoresis using a 1% agarose gel stained with GelRed (Biotium) in 1X TBE buffer. We performed conventional polymerase chain reaction (PCR) on DNA from each successfully extracted isolate using primers for previously described multilocus sequence typing (MLST) markers (8009X2, BdC5, BdSc2.0, BdSc4.16, BdSc7.6, R6046, BdSC6.15) and the 18S rRNA gene (Morehouse et al., 2003; Morgan et al., 2007; James et al., 2009; Schloegel et al., 2012; Jenkinson et al., 2016) (Table S1). Amplifications were performed in 25 µl volumes consisting of 12.5 µl DreamTaq PCR Master Mix (2X) (Thermo Fisher Scientific, Inc.), 11.5 µl nuclease free water, 0.25 µl forward primer (10 µM), 0.25 µl reverse primer (10 µM), and 0.5 µl template DNA. Thermocycling conditions consisted of an initial denaturing step of 2 min at 95 °C, then 32 cycles of 30 s at 95 °C, 30 s at 52 °C–60 °C depending on the primer pair (Table S1), and 45 s at 72 °C, with a final extension of 10 min at 72 °C. We treated 5 µl aliquots of PCR product with 2 µl ExoSAP-IT (Affymetrix Inc.) and incubated the total volume at 37 °C for 15 min and then 80 °C for 15 min. Using each of the MLST primers, we performed cycle sequencing reactions using Big Dye v3.1 dye terminator (Applied Biosystems, Inc.). We incubated hydrated G-50 Sephadex (2.6 g/45 ml H2O) at room temperature for 30 min, pipetted 400-µl aliquots into individual wells in a filter plate, and centrifuged the plate at 1,258×g for 2 min to create a matrix through which cycle sequenced products were passed for purification. We dehydrated the purified cycle sequenced products, added 12 µl of formamide, and incubated them for 3 min at 94 °C and immediately cooled them to 4 °C. We electrophoresed the cycle sequenced products on an ABI 3100-
Avant genetic analyzer (Applied Biosystems, Inc.) and trimmed and edited the resulting chromatograms using Geneious v9.1.5 (Kearse et al., 2012).

2.4. Data analysis

We genotyped strains using reference sequences from Jenkinson et al. (2016). We then used a set of genotyped strains from across North and South America to perform comparative analyses and test for genetic structure across the Western Hemisphere. We limited our analyses to the Western Hemisphere owing to the relatively few number of strains sequenced at these same markers in the Eastern Hemisphere. These strains were selected from two previous studies that used at least six of the same genetic markers used in our study (Schloegel et al., 2012; Jenkinson et al., 2016). We excluded any strains that had been isolated from amphibians not found in the wild (this included animals from zoos, farms, and markets) for our analyses. The resulting dataset consisted of 177 strains, including the 37 Texas strains isolated in our study. Before calculating measures of genetic diversity, we divided our dataset into populations and subpopulations. These populations (and subpopulations) included temperate America BdGPL (subpopulations: eastern North America, Texas, and western North America), tropical America BdGPL (subpopulations: Panama and Brazil), BdAsia2/BdBrazil, and BdAsia2/BdBrazil-BdGPL hybrids. For each population and subpopulation, we calculated genotypic diversity and allelic richness using the R package PopGenReport (Adamack and Gruber, 2014). Next, we clone-corrected our datasets by removing identical genotypes from the same region. This resulted in a dataset that included the genotypes of 68 isolates, including 15 unique Texas isolates, at six MLST markers (Supplementary appendix 2). We then calculated gene diversity, or expected heterozygosity ($H_e$), and observed heterozygosity ($H_o$) using the R package adegenet (Jombart, 2008) and calculated pairwise $F_{ST}$ values using GENEPOP (Rousset, 2008).

To test for genetic structure among Bd isolates, we used STRUCTURE (Pritchard et al., 2000) with values of $K$ (number of genetic demes defined a priori) ranging from 1 to 12 and 5 iterations per $K$, with 500,000 Markov Chain Monte Carlo repetitions with a burn-in of 100,000. We determined the best number of populations by calculating $\Delta K$ (Evanno et al., 2005) using STRUCTURE HARVESTER (Earl and Vonholdt, 2012) and obtained average membership probabilities across all iterations per $K$ using the program CLUMPP (Jakobsson and Rosenberg, 2007). We performed an initial analysis with only our Texas isolates and their genotypes at all seven loci (8009x2, BdC5, BdSC4.16, BdSC7.6, R6046, BdSC6.15, BdSC2.0) to test for genetic structure within the state, followed by a second analysis combining the Texas isolates with the set of isolates from North and South America that had been sequenced and genotyped using at least six of the same markers (8009x2, BdC5, BdSC4.16, BdSC7.6, R6046, BdSC6.15). Both STRUCTURE analyses were performed using clone-corrected data.

2.5. Host sequencing and identification

We extracted DNA from tail muscle of collected tadpoles and toes of seven adult Rana berlandieri using the same extraction protocol described for Bd. Conventional PCR was performed to amplify the cytochrome c oxidase I (COX1) gene (Table S1), and PCR products were purified and sequenced using the same methods described for Bd sequencing. Chromatograms were trimmed, edited, and aligned in Genious v 9.1.5 (Kearse et al., 2012). Using CLUSTALW, implemented through Genious, we carried out a multiple sequence alignment of host sequence data (cost matrix = clustalw, gap open cost = 15, gap extend cost = 6.66). We used sequence similarity to assign anuran larvae to species and visualized the distance matrix (HKY) using a neighbor-joining topology. We included reference sequences from Pseudacris streckeri (GenBank: KJ536156), Rana muscosa (GenBank: KU985709), Rana. clamitans (GenBank: KYS87195), and Rana. sphenophela (GenBank: KT388406) and newly generated reference sequences for adult R. berlandieri (GenBank: MG969220; GenBank: MG969221; GenBank: MG969222; GenBank: MG969223; GenBank: MG969224; GenBank: MG969225; GenBank: MG969226). We suspected that some of the larvae belonged to these species based on geographic distribution and general morphology. R. muscosa was chosen as an outgroup to Texas ranids.

3. Results

We attempted to isolate Bd from a total of 140 tadpoles collected from sites in central, east, and north-central Texas (Fig. 1). From this total, we successfully cultured 41 chytrid isolates from 40 individual tadpoles comprising three host species. Of these 41 isolates, 37 were identified as Bd (Supplementary appendix 1). A total of 28 isolates have been characterized using all seven MLST markers, seven have been characterized at six loci, and two were characterized at five loci (Supplementary data). We were unable to amplify DNA of four isolates from two R. clamitans larvae using these markers but did amplify and sequence part of the 18S rRNA gene (Supplementary appendix 1). Using the NCBI BLAST search tool (Clark et al., 2016), we matched isolates TXST038 and TXST039 to Rhizophylicta harderi (GenBank: AF164272; match = 98%, coverage = 100%, E value = 0.0), which has recently been re-named Uebelmesseromyces harderi (Powell et al. 2015), TXST041 to Clytomyces sp. (GenBank: DS536486; match = 98%, coverage = 100%, E value = 0.0), and isolate TXST038 to Halyoraphidium curvatum (GenBank: NG_017172; match = 99%, coverage = 100%, E value = 0.0).
value = 0.0), all of which are species of chytrids not known to infect vertebrate hosts. Among the 37 Bd isolates, there were 19 unique multi-locus genotypes (MLGs), and no MLGs were found at more than one collection site (Supplementary data).

A neighbor-joining topology (Fig. S1) revealed four distinct clades among potential hosts. Three clades were associated with R. clamitans, R. sphenocephala, and Pseudacris streckeri reference sequences. We suspected that some of our tadpole specimens were Rana berlandieri, although a COI reference sequence for this species was unavailable in GenBank. Thus, we sequenced seven R. berlandieri metamorphs from central, south, western parts of Texas (Supplementary appendix 1) to verify species assignments. All newly generated sequences for both pathogen and host were accessioned into GenBank (chytrid 18S, MG979804–MG979842; amphibian COI, MG969220–MG969347).

### 3.1. Measures of genetic diversity

Gene diversity ($H_E$) among the clone corrected Western Hemisphere dataset was 0.506 and observed heterozygosity ($H_O$) was 0.370. When all BdGPL strains were divided into temperate and tropical groups, the temperate group had both higher $H_E$ and $H_O$ (0.402 and 0.394, respectively) than the tropical group (0.358 and 0.340, respectively). Both groups had higher $H_E$ and $H_O$ than BdAsia-2/BdBrazil (0.189 and 0.214, respectively). While larger pooled populations had higher expected than observed heterozygosity, the subpopulations in our dataset tended to show the reverse trend (except for the BdGPL in western North America and southeastern Brazil). While Texas strains had lower $H_E$ (0.348) than both eastern and western North American subpopulations, $H_O$ (0.387) was intermediate between the two other North American regions. All North American subpopulations, including Texas, had higher $H_O$ than Panama and southeastern Brazil, while Texas was the only North American subpopulation to have lower $H_E$ than the BdGPL in southeastern Brazil. Panama had the lowest $H_E$ and $H_O$ of all BdGPL subpopulations. The temperate BdGPL had higher genotypic diversity (0.491) and allelic richness (1.81) compared to the tropical BdGPL (0.250 and 1.68, respectively) and BdAsia-2/BdBrazil (0.280 and 1.34, respectively). Texas strains had lower genotypic diversity (0.405) and allelic richness (1.72) than eastern and western North American strains but displayed greater diversity in these measures than both tropical groups. Not surprisingly, the BdAsia-2/BdBrazil-BdGPL hybrid strains had higher measures of genetic diversity than all other groups (Table 2).

### 3.2. Cluster analyses

We performed a STRUCTURE analysis using seven loci on the 19 unique MLGs from Texas and did not detect population structure among these strains. We then conducted a STRUCTURE analysis on the combined dataset that included our Texas isolates, as well as additional Western Hemisphere genotypes from two previous
studies (Schloegel et al., 2012; Jenkinson et al., 2016). For these analyses, we chose six MLST markers (r6046, BdSc6.15, 8009x2, BdC5, BdSc4.16, BdSc7.6) that had been sequenced in all three studies, thus we used 15 unique MLGs from Texas to compare against other New World isolates (Supplementary appendix 2).

Our STRUCTURE analysis indicated two to three genetically distinct demes in our dataset, with the highest ∆K value occurring at K = 2 and second highest value at K = 3 (Fig. 2A). By using the already genetically identified isolates from the two previous studies as references, we determined our clusters corresponded to BdGPL and BdAsia-2/BdBrazil at K = 2. At K = 3 there was some support for weak substructure within the BdGPL, which corresponded to two genetic demes representing BdGPL1 and BdGPL2 (Fig. 2A). We did not detect the presence of any BdAsia-2/BdBrazil or hybrid strains in Texas. We set an arbitrary threshold of q = 0.80 for assigning strains to a particular cluster. Ambiguous strains that did not show high assignment to one cluster included the putative BdAsia-2/BdBrazil-BdGPL hybrids, two strains from Texas, and three from Brazil that showed some support for admixture between the two BdGPL demes. Pairwise FST values indicate much greater differentiation between BdAsia-2/BdBrazil and all other populations, as expected, while Texas isolates showed no difference compared to eastern North American isolates and relatively low differentiation (0.040) compared to isolates in western North America (Fig. 2B, Table S2).

4. Discussion

Bd has shown contrasting pathogen dynamics in North America (Wake and Vredenburg, 2008; Padgett-Rowl and Hopkins, 2009; Briggs et al., 2010; Talley et al., 2015), yet genetic information from cultured isolates is mainly restricted to the western and eastern U.S., leaving a large swath of the central U.S. without data on Bd genotypes, and thus information on strain types. We successfully isolated 37 chytrids identified as Bd and four other chytrids (three species) from three different amphibian host species native to central and east Texas. As a population, Bd isolates in this region do not show significant genetic differentiation from other North American isolates. Further, all Bd isolates collected in this study belong to the globally distributed BdGPL, and none show genetic affinities to other major lineages (e.g. BdAsia-2/BdBrazil, BdCH, BdCAPE, or BdAsia-1).

Our measures of genetic diversity across geographic regions in the Western Hemisphere are congruent with previous studies. James et al. (2009) found lower Hc and H0 among tropical BdGPL strains compared to temperate BdGPL strains using MLST markers. The latitudinal gradient in genetic diversity observed by Velo-Anton et al. (2012) from California to Panama is also congruent with our estimates. While North American strains have the highest measures of genetic diversity, Texas strains, which represent the southernmost group of North American strains in our dataset, tend to have slightly lower measures of diversity compared to the rest of the continent. Meanwhile, strains from Panama, the region closest to the equator in our dataset, have the lowest measures of diversity among the GPL. These trends could be reflective of a longer history of Bd in North America compared to the tropics, but we stress the need for further isolations from North America, especially in Mexico, and genomic sequencing of these strains to draw stronger conclusions about the demographic history of Bd in the Western Hemisphere.

A recent global genomic survey of Bd did not recover a strong distinction between BdGPL1 and BdGPL2 (O’Hanlon et al., 2018). This is congruent with our ∆K results showing stronger support for one BdGPL deme and less support for two genetic demes within New World BdGPL isolates. Although, at K = 3 these genetic demes correspond to the previously defined groups BdGPL1 and BdGPL2 (Schloegel et al., 2012; Jenkinson et al., 2016) and may reflect weak regional substructuring. Of the 37 cultured Texas isolates, 32 appear to cluster with BdGPL1 isolates. Three Texas isolates (TXST021, TXST026, and TXST035) may cluster with BdGPL2, and two isolates (TXST012 and TXST033) may exhibit some degree of assignment uncertainty between both BdGPL groups (Fig. 2A). Evidence for hybridization between Bd lineages has been detected (Schloegel et al., 2012; Jenkinson et al., 2016), thus it is possible that these ambiguous strains could be the result of admixture between two BdGPL demes. We should point out, however, that there is some ambiguity in assigning membership to these two sub-lineages within BdGPL. Genomic data may resolve these uncertainties and better inform tests for admixture between intraregional isolates.

At the continental scale, Bd isolates in Texas genetically cluster with isolates responsible for severe outbreaks of chytridiomycosis in the western United States (Fig. 2B). This finding leaves us with a still unanswered question—why do these strains, which are genetically similar, cause disease and mortalities in some regions but not others? If the pathogen is genetically similar in these different regions, the other two points of the epidemiological triangle—host and environment—should be explored further. One intuitive explanation for the lack of known chytridiomycosis outbreaks in Texas is the influence of climate. Piotrowski et al. (2004) reported Bd’s optimal temperature range to be 17–25 °C with a maximum threshold of 28 °C. In Texas, amphibians may be able to clear or reduce Bd infections before they become lethal coincident to hot summers, when temperatures regularly exceed 32 °C.

Table 2

<table>
<thead>
<tr>
<th>Region/Population</th>
<th>N</th>
<th>MLG</th>
<th>Genotypic Diversity</th>
<th>Mean Allelic Richness</th>
<th>Clone Corrected Data</th>
<th>Expected Heterozygosity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperate America (BdGPL)</td>
<td>57</td>
<td>28</td>
<td>0.491</td>
<td>1.81</td>
<td>0.394</td>
<td>0.402</td>
</tr>
<tr>
<td>Eastern N.A.</td>
<td>7</td>
<td>7</td>
<td>1.000</td>
<td>1.83</td>
<td>0.429</td>
<td>0.378</td>
</tr>
<tr>
<td>Western N.A.</td>
<td>13</td>
<td>13</td>
<td>1.000</td>
<td>1.91</td>
<td>0.385</td>
<td>0.435</td>
</tr>
<tr>
<td>Texas</td>
<td>37</td>
<td>37</td>
<td>0.405</td>
<td>1.72</td>
<td>0.387</td>
<td>0.348</td>
</tr>
<tr>
<td>Tropical America (BdGPL)</td>
<td>92</td>
<td>23</td>
<td>0.250</td>
<td>1.68</td>
<td>0.34</td>
<td>0.358</td>
</tr>
<tr>
<td>Panama</td>
<td>3</td>
<td>3</td>
<td>1.000</td>
<td>1.59</td>
<td>0.333</td>
<td>0.259</td>
</tr>
<tr>
<td>Southeastern Brazil</td>
<td>89</td>
<td>21</td>
<td>0.236</td>
<td>1.67</td>
<td>0.341</td>
<td>0.354</td>
</tr>
<tr>
<td>All BdGPL</td>
<td>149</td>
<td>47</td>
<td>0.315</td>
<td>1.96</td>
<td>0.373</td>
<td>0.419</td>
</tr>
<tr>
<td>Hybrids (BdAsia-2/BdBrazil-BdGPL)</td>
<td>3</td>
<td>2</td>
<td>0.667</td>
<td>1.34</td>
<td>0.214</td>
<td>0.438</td>
</tr>
<tr>
<td>BdAsia-2/BdBrazil</td>
<td>25</td>
<td>7</td>
<td>0.280</td>
<td>0.370</td>
<td>0.370</td>
<td>0.506</td>
</tr>
</tbody>
</table>

* Sample size.

b Number of multi-locus genotypes.
Indeed, *Bd* infections in *Acris crepitans* populations in central Texas have revealed a seasonal pattern in infection intensity, with peaks occurring in early spring followed by a decline throughout the summer months (Gaertner et al. 2009b, 2012). Additionally, temperatures in most parts of the state regularly exceed *Bd*’s thermal optimum for over six months of the year (U.S. climate data 2017). However, weather conditions cannot explain the lack of *Bd* associated morbidity in central Texas salamanders of the genus *Eurycea*, which dwell in cool, thermally stable aquifers and springs. While *Bd* infections in these salamanders have been documented (Gaertner et al., 2009a), symptoms of chytridiomycosis have not been detected in these populations despite several years of investigation (Bowles et al., 2006; Pierce et al., 2010; Bendik, 2017). Differences in host resistance among and

---

Fig. 2. (A) STRUCTURE and STRUCTURE HARVESTER results for *K* = 2 and *K* = 3 averaged from five iterations using CLUMPP. Assignment probabilities to the clusters are represented on the y-axis, and each bar represents one of the 68 isolates from a six MLST dataset, including 15 isolates collected for this study and 53 additional strains from Schloegel et al. (2012) and Jenkinson et al. (2016). (B) Heat map of pairwise *F*_{ST} values by origin calculated from the dataset of 68 *Batrachochytrium dendrobatidis* isolates, including 15 isolates from central Texas and 53 additional strains from Schloegel et al. (2012) and Jenkinson et al. (2016).
even within species have been documented (Woodhams et al., 2007; Savage and Zamudio, 2011), so it is possible that host species immunity plays a role in limiting the severity of disease outcomes in central Texas. The Bd strains isolated for this study will facilitate future studies necessary to dissect the effects of both climate and host resistance on the dynamics of Bd infection in this region.

Non-lethal Bd isolation and culturing efforts from amphibians collected in Madagascar, Portugal, and Taiwan have isolated other non-Bd chytrid species, Rappamyces sp. and Rhizophydom sp. (Fisher et al., 2018). Likewise, our isolation and culturing efforts recovered chytrids not known to infect vertebrate hosts, namely U. harderi (Rhizophydomia), Chytriales sp. (Chytridiales), and H. curvatum (Monoblepharidales) (Fig. 1, Supplementary appendix 1). Zoosporic fungi of the phylum Chytridomycota are generally saprobes or parasites of algae, plants, and invertebrates (Barr, 2001). It is possible that these chytrids were present in the pond water from which these tadpoles were collected, and that they somehow persisted on the tadpole mouthparts despite our following established procedures to clean them by dragging through sterile agar several times (Longcore et al., 1996). Although studies of the morphology and systematics of U. harderi, Chytriales, and H. curvatum can be found in the literature, little appears to have been written about the life history of these species (Ustunova et al., 2000; Foresti et al., 2002; Letcher and Powell, 2002; Powell et al., 2015). Many chytrids have similar, uninformative morphological features (James et al., 2000), and what appear to be successful isolations may not represent Batrachochytrium species. Yet, these isolations and cultures are still useful in diversity discovery efforts, especially among the Chytridomycota, a relatively understudied group of fungi (James et al., 2000). By sequencing isolates at the 18S rRNA gene, we were able to phylogenetically place these isolates in North America. There are also important conservation implications for the region as well. Texas is home to 16 endemic amphibian species, which are found in either the Edwards Plateau or the coastal prairies of Texas (Tipton et al., 2012). Knowledge of the Bd history of these species (Gaertner et al., 2010; Degler and Milillo, 2018) and habitat alteration.

Our investigation of genetic variation among Bd isolates from parts of Texas is an important first step in understanding disease dynamics between amphibians and Bd in this understudied region. We have gained a more complete picture of the distribution of Bd strains in this region and have begun to fill a large sampling gap for Bd isolates in North America. There are also important conservation implications to our understanding of chytridiomycosis dynamics in the region as well. Texas is home to 16 endemic amphibian species, many of which are federally listed as species of concern, and all of which are found in either the Edwards Plateau or the coastal prairies of Texas (Tipton et al., 2012). Knowledge of the Bd strains present in these regions can inform assessments of potential disease risks to host species in the face of continued climate change and habitat alteration.

Acknowledgements

We thank Abigail Romo, Charles and Rebecca Raleigh, and the Brackenridge Field Laboratory for access to collection sites. We dedicate this work to the memory of Jonathan Raleigh. We thank Dan Saenz, Trina Guerra, Stephen Roussos, Jeremy Weaver, Brandon Cranup, William Keitt, Jacob Owen, Madison Torres, Andrea Villamizar, Stephen Harding, Carlos Baca, Daniel Puckett, Edith Perez, Devin Jackson, Anna Gates, and Stephanie Toussaint for assistance in the lab and field. We thank Ana V. Longo for reviewing an earlier version of this manuscript. This work was funded by a National Science Foundation Graduate Research Fellowship to TLM (DGE-1144466), a Texas State University Research Enhancement Program grant to DR, Start-up funds from Texas State University to DR, and a CAPES fellowship (BEX 1176/13-7) and Texas Ecolab Program grant to DTC.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.funeco.2018.12.001.

References


Ustinova, I., Krienitz, L., Huss, V.A.R., 2000. Hyaloraphidium curvatum is not a green alga, but a lower fungus: Anoomoebidium parasiticum is not a fungus, but a member of the DRIPs. Protist 151, 253–262.


Wake, D.B., Vredenburg, V.T., 2008. Are we in the midst of the sixth mass extinc-