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Distribution and environmental limitations of an amphibian pathogen in the Rocky Mountains, USA

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ABSTRACT

Amphibian populations continue to be imperiled by the chytrid fungus (*Batrachochytrium dendrobatidis*). Understanding where *B. dendrobatidis* (*Bd*) occurs and how it may be limited by environmental factors is critical to our ability to effectively conserve the amphibians affected by *Bd*. We sampled 1247 amphibians (boreal toads and surrogates) at 261 boreal toad (*Bufo boreas*) breeding sites (97 clusters) along an 11° latitudinal gradient in the Rocky Mountains to determine the distribution of *B. dendrobatidis* and examine environmental factors, such as temperature and elevation, that might affect its distribution. The fungus was detected at 64% of all clusters and occurred across a range of elevations (1030–3550 m) and latitudes (37.6–48.6°) but we detected it in only 42% of clusters in the south (site elevations higher), compared to 84% of clusters in the north (site elevations lower). Maximum ambient temperature (daily high) explained much of the variation in *Bd* occurrence in boreal toad populations and thus perhaps limits the occurrence of the pathogen in the Rocky Mountains to areas where climatic conditions facilitate optimal growth of the fungus. This information has implications in global climate change scenarios where warming temperatures may facilitate the spread of disease into previously un- or little-affected areas (i.e., higher elevations). This study provides the first regional-level, field-based effort to examine the relationship of environmental and geographic factors to the distribution of *B. dendrobatidis* in North America and will assist managers to focus on at-risk populations as determined by the local temperature regimes, latitude and elevation.

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1. Introduction

Chytridiomycosis is a potentially lethal disease of amphibians caused by the amphibian chytrid fungus (*Batrachochytrium dendrobatidis* [*Bd*]) that has been associated with population declines in several amphibian species throughout the world (Daszak et al., 1999; Carey, 2000; Green et al., 2002; Lips et al., 2006). Much of our understanding of the host–pathogen relationship of amphibians and *Bd* in the wild comes from the

tropics, particularly Central and South America and Australia. Research in the tropics suggests that mid- to high-elevations provide ideal temperatures for *Bd* and amphibian communities at such elevations are more susceptible to infection (Daszak et al., 2003; Berger et al., 2004; Woodhams and Alford, 2005; Drew et al., 2006). Other studies from the tropics suggest that global climate change may be enabling *Bd* to expand its range due to increasingly warm temperatures and moisture at higher elevations (Bosch et al., 2006; Pounds et al., 2006;

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Seimon et al., 2006). Although there is evidence to support this paradigm, it has not been well tested in the northern hemisphere and information on environmental factors limiting the distribution of *Bd* in temperate regions is lacking.

The boreal toad (*Bufo boreas*) is a terrestrial anuran that is distributed widely across western North America. Toads in the Southern Rocky Mountains tend to be found at higher elevations (2615–3385 m, Colorado, Hammerson, 1999) than those in the northern parts of the region (1609–2865 m, in Yellowstone and Grand Teton National Parks (Koch and Peterson, 1995); 2810 m maximum in Montana (Maxell et al., 2003)). *Bufo boreas* appears to have suffered increasingly widespread declines along a gradient in the Rocky Mountains from north to south, a pattern possibly associated with the distribution of *Bd*. For example, *Bufo boreas* has disappeared from many historic localities, especially in Colorado, over the last 20–30 years (Carey, 1993; Corn, 2000; Corn et al., 2005). Some of these declines have been linked to *Bd* infection (Muths et al., 2003; Scherer et al., 2005). In contrast, *Bufo boreas* populations in the Northern Rocky Mountains are perceived to be fairly stable (Corn et al., 2005) and *Bd* was not found in museum specimens from Montana or Canada prior to 1977 (B. Maxell, Montana Heritage Program, pers. comm.; Ouellet et al., 2005).

Recent studies have provided information on the distribution of *Bd* in parts of North America (Ouellet et al., 2005, North America; Pearl et al., 2007, Pacific Northwest; Longcore et al., 2007, Northeast), but information on the distribution of *Bd* in the Rocky Mountains is incomplete. These data are needed given that *Bd* is a known pathogen of boreal toads in the laboratory (Blaustein et al., 2005; Carey et al., 2006) and has been isolated from sick and dying toads in the field (Raverty and Reynolds, 2001; Muths et al., 2003). The earliest known occurrences of *Bd* for locations in this study are in Colorado in the mid 1970s (*Rana pipiens*, Carey, 2000) with later occurrences in Wyoming (*Bufo baxteri*, Muths et al., 2003), Montana and Idaho (*B. boreas* and *Dicamptodon aterrimus*, respectively, D.E. Green, USGS, National Wildlife Health Laboratory, personal communication). Distributional information can provide insights on what species or populations may be resistant to chytridiomycosis and, importantly, what amphibian populations are at risk and should be targeted for conservation actions.

Information on the environmental limitations of *Bd* in the wild is critical to the conservation of amphibians affected by this disease. Variation in environmental factors clearly influences the distribution of *Bd*, although there is little empirical data to fully understand the fundamental or realized niche of this pathogen (e.g., Ron, 2005). Laboratory studies have demonstrated that this fungus produces zoospores in vitro between 4 and 25 °C (Piotrowski et al., 2004), pathogenicity decreases > 23 °C (Lamirande and Nichols, 2002; Woodhams et al., 2003; Berger et al., 2004) and *Bd* persists for up to 3 months in moist, sterile river sand (Johnson and Speare, 2003), but little else is known. Although other aspects of climate, such as rainfall, likely play a role in this host–pathogen interaction, we focused on temperature for this study because previous work indicates that temperature is one of the most important environmental parameters determining the persistence and perhaps pathogenicity of the fungus (Piotrowski et al., 2004; Drew et al., 2006; Kriger and Hero, 2007; Kriger et al., 2007). Given global climate change scenarios, and

mounting evidence that rising temperatures, diminished snowpack, earlier run-offs and drought are affecting the western United States now (Barnett et al., 2008), it is increasingly important to understand how rising temperatures, particularly at northern latitudes and high altitudes, will affect host–pathogen relationships.

The goals of this study were to (1) determine the current distribution of *Bd* at boreal toad breeding localities along a latitudinal gradient, roughly paralleling the continental divide, (2) explore how this distribution may be affected by local climate, and (3) discuss how environment–pathogen relationships may influence declining amphibian species in the Rocky Mountains.

2. Materials and methods

2.1. Sample site selection

We sampled all known boreal toad breeding sites in Colorado (Jackson, 2005) and 30 boreal toad breeding sites each in Montana and Wyoming randomly from all known breeding localities (C. Peterson and D. Patla, Idaho State University, personal communication, Maxell et al., 2003, Wyoming Game and Fish Department and Wyoming Natural Diversity Database [<http://uwadmweb.uwyo.edu/wyndd/>]). Additional samples were collected opportunistically in Montana, Idaho, Wyoming, and Colorado. The northernmost site was at 48.6° N. latitude and the southernmost at 37.6° N. latitude. Sites ranged in elevation from 1030 m to 3800 m.

2.2. Site characteristics

We used a global positioning system (± 10 m accuracy) to determine geographic coordinates of each collection site. Elevation data were derived from the national elevation dataset (Gesch et al., 2002). We used program PRISM (2.5 min digital elevation model, Daly et al., 2002, <http://www.ocs.oregonstate.edu/prism/index.phtml>) to estimate mean maximum and minimum daily temperatures averaged from 2002 to 2004 (study period weather) and from 1970 to 2005 (long-term climate) during the active season (May–September) for each sample location. We then projected mean maximum and minimum daily temperatures during the active season at each sample location to the year 2100 (Govindasamy et al., 2003).

2.3. Sample collection and analysis

We sampled 1151 adult boreal toads and 96 individuals from 5 other species when toads were not abundant or not found at our selected sites. Samples were collected between April and September 2000–2004; most samples (84%) were collected between May and July. At randomly selected sites, surveyors attempted to capture 10 adult boreal toads to collect ventral skin swabs or scrapings. If toads were unavailable, other adult amphibians were captured and sampled as surrogates. In Colorado, sites were surveyed for boreal toads during the day or night and toe clips or ventral swabs were obtained from up to 20 individual toads per site. Fewer swabs were collected at sites that were sampled opportunistically due primarily to time constraints.

The ventral surface and the hind foot webbing of each animal was swabbed or scraped ≥ 25 strokes and the sample was placed in 70% ethanol in a 0.5 ml o-ring vial and labeled. A few individuals were sampled by clipping a single digit. Differences in detectability between the sample methods are not significant (swab versus scrape, Livo, 2004; swab versus toe clip, Hyatt et al., 2007). To avoid potential contamination of the collected tissue, we adhered scrupulously to clean procedures in the field for capturing, handling and sample collec-

tion, including single-use (=single animal) gloves, and single-use containers. Each vial was sealed in an individual plastic bag. Samples were sent to Pisces Molecular (Boulder, CO) where polymerase chain reaction (PCR) techniques (Annis et al., 2004 modified by J. Wood, Pisces Molecular, Boulder, CO, personal communication) were used to detect *Bd*. This technique does not differentiate between the simple presence of the fungus on the skin of the amphibian and a clinical infection that is affecting the animal's health (Smith, 2007).

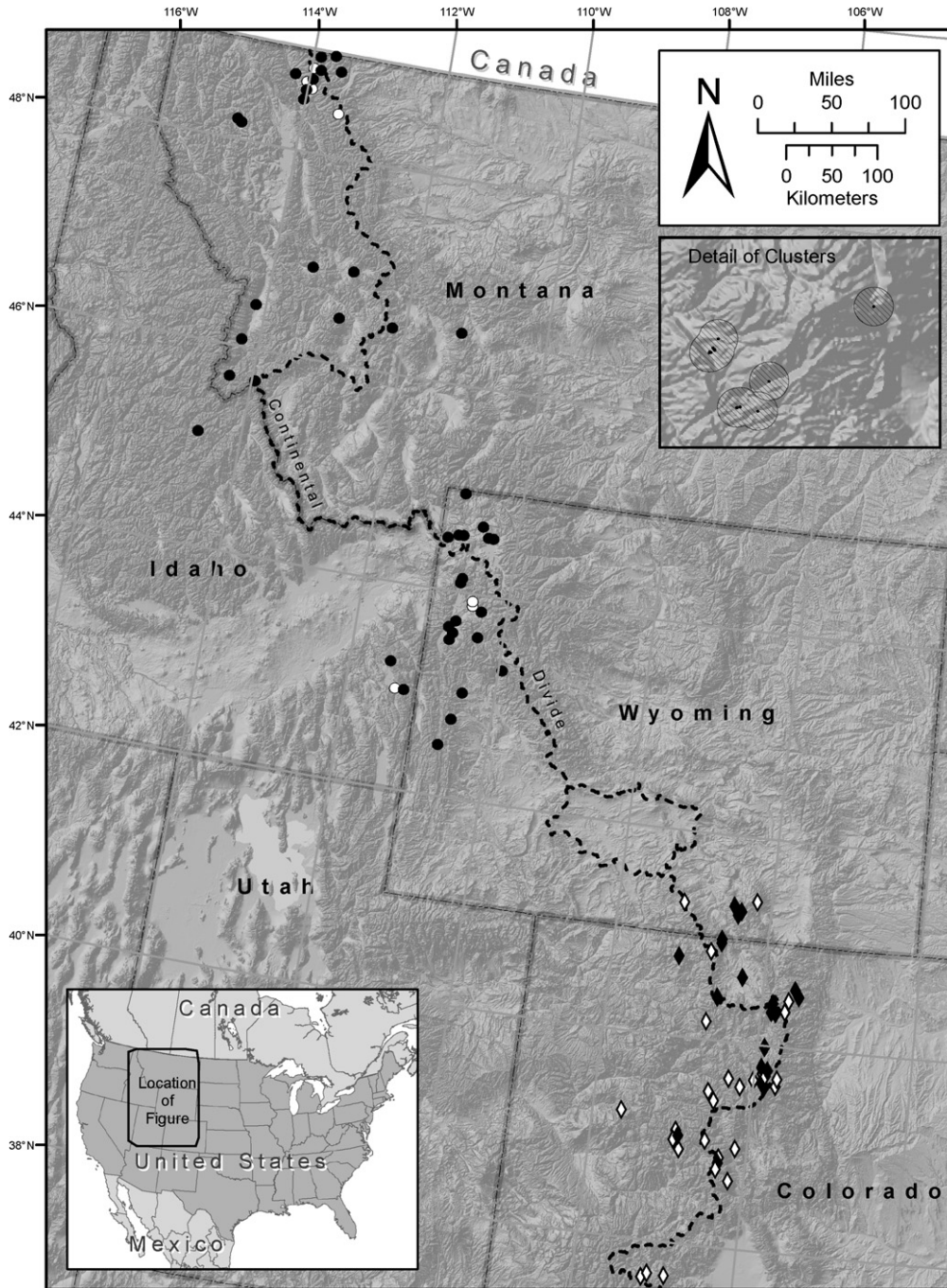


Fig. 1 – Map of clusters sampled along the Continental Divide (- - -). Clusters ($n = 97$) are composed of boreal toad breeding sites (=sampling sites). Sites were grouped such that sites closer than 3 km formed a cluster. However, buffer zones less than 6 km apart may overlap (e.g., clusters in detail). Diamonds indicate southern clusters, squares indicate northern clusters; filled symbols are *Bd* positive clusters, open symbols are *Bd* negative clusters.

2.4. Data analysis

Sample locations were considered individual sites only if they were separated from other sample locations by ≥ 3 km. If sample locations were < 3 km from one another, they were merged into a cluster formed by the overlap of each 3-km buffer zone and the center of the cluster determined using GIS (ARCGIS 9.0, ESRI Corp, Redlands, CA) (Fig. 1). By pooling individual toads (=results) from sites within clusters, we increased our sample size at each cluster (Table 1), thus reducing potential false negative results (i.e., reporting *Bd* was absent when it was actually present) and increasing the independence between our sample units (Petranka et al., 2004). Although we have observed boreal toads traveling > 6 km between breeding sites (E.M. personal observation), we chose 3 km as a buffer distance because this represents approximately twice the mean maximum one-way migration distance of boreal toads published in other studies (Muths, 2003; Bartelt et al., 2004).

We analyzed the occupancy of *Bd* in 97 clusters, including 89 clusters with boreal toads and 8 clusters where other species were swabbed as surrogates because toads were not found. Eight clusters contained samples from 2 or more species. We did not examine prevalence (the number of cases of a disease in a defined population at a particular point in time; sensu Hunter, 1997), because we were not confident that the presence of *Bd* isolated from the skin of a small number of animals in a cluster is an accurate measure of prevalence of *Bd* in a population. Furthermore, detection of *Bd* via PCR in a population is strongly influenced by season and tempera-

ture (Kriger and Hero, 2007) and we did not randomize the timing of sampling.

We plotted mean temperatures (minimum and maximum) during the active season for all clusters and placed our data within the context of what is known about temperature tolerances of *Bd* in the laboratory (Piotrowski et al., 2004). T-tests were used to compare mean temperature values between the northern and southern portions of the continental divide transect.

We used logistic regression to assess the effect of selected environmental variables (date, elevation, and temperature) on occurrence. The number of samples collected at each cluster was modeled as a measure of effort. Potential over-dispersion in the data was assessed using the deviance/d.f. value for the most parameterized model. We used Akaike's information criterion corrected for small sample sizes (AIC_c) and Akaike weights to determine which model best reflected our data (Burnham and Anderson, 2003). We also assessed the probability of observing a false negative result by first computing $p (=n_i/n)$ for each cluster where *Bd* was detected; where p = the probability of the fungus occurring, n_i = the number of positive samples and n = the total number of samples. Then we calculated a mean value for $p (=p^*)$ for all clusters and used this value to calculate the probability of a false negative at clusters where we did not detect the fungus $(1 - p^*)^n$. Finally, we examined the sensitivity of our results to this potential error (false negative) by re-assigning a random selection of potential false negative clusters (based on the probability described above) to positive and re-assessing the top model using a standard bootstrapping approach (1000 iterations). All analyses were conducted in SAS (SAS Institute, 2003) except the bootstrapping, which was programmed in R (www.r-project.org/).

Table 1 – Sampling effort: Number of clusters and sites; number of samples tested for *Batrachochytrium dendrobatidis*

Group	No. of clusters	No. of sites	No. samples tested	Median no. of samples/cluster (range)
<i>Northern</i>				
Montana	24	107	225	6 (1–47)
Idaho	4	10	18	4 (2–8)
Wyoming	21	45	225	7 (1–86)
<i>Southern</i>				
Colorado	43	83	752	13 (1–134)
Wyoming	5	16	29	4 (1–12)
Total	97	257	1248	7 (1–134)

3. Results

DNA from *Bd* was isolated from all 6 species sampled (Table 2). Although the sample size for some species was very small, boreal toads had the lowest overall proportion of individuals that tested positive for *Bd*. This pattern was influenced strongly by samples from the southern region of our study area, where only 10% of boreal toads tested positive for *Bd* compared with 46% in northern region (Table 3).

We detected *Bd* in 64% of the 97 clusters sampled (Table 3). *Batrachochytrium dendrobatidis* was detected from the lowest (37.6 N) to the highest (48.6 N) latitudes and across a range of elevations (1030–3550 m) (Fig. 3). Although *Bd* was detected

Table 2 – Summary of *Batrachochytrium dendrobatidis* (*Bd*) results from 6 species tested

Species	States	No. of clusters	No. of individuals sampled	No. of individuals <i>Bd</i> positive	No. of individuals <i>Bd</i> negative	Percent of individuals <i>Bd</i> positive
<i>Ambystoma tigrinum</i>	WY	2	2	1	1	50
<i>Bufo boreas</i>	ID, CO, MT, WY	89	1151	265	886	23
<i>Pseudacris maculata</i>	CO, WY	7	14	6	8	43
<i>Rana luteiventris</i>	ID, MT, WY	12	50	20	30	40
<i>Rana pipiens</i>	CO	1	4	3	1	75
<i>Rana sylvatica</i>	CO, WY	6	26	9	17	35
Total			1247	304	943	24

Table 3 – Summary of the detection of *Batrachochytrium dendrobatidis* (Bd) across the region relative to average temperature (1) during the study; (2) the 35 year average temperature (PRISM, Daly et al., 2007); and (3) predicted temperature for the year 2100 (Govindasamy et al., 2003)

Clusters	Average temperature during study (2002–2004)				Average temperature 1970–2005 (35 years)				Average temperature 2100 (predicted)			
	TempMin		TempMax		TempMin		TempMax		TempMin		TempMax	
	Mean	Range	Mean	Range	Mean	Range	Mean	Range	Mean	Range	Mean	Range
<i>Northern</i> (1030–2661 m)												
Bd+ (n = 41)	3.8	1.0–7.4	19.7	15.5–25.1	2.6	–0.2–6.5	19.5	15.1–25.0	4.3	–0.8–8.0	23.1	18.7–27.8
Bd– (n = 8)	4.4	2.2–6.4	19.2	15.3–21.1	3.2	1.7–6.0	18.8	14.7–20.9	5.0	2.7–7.0	22.7	18.2–24.6
All sites North	3.9	1.0–7.4	19.6	15.3–25.1	2.7	–0.2–6.5	19.4	14.7–25.0	4.4	–0.8–8.0	23.0	18.2–27.8
<i>Southern</i> (2458–3805 m)												
Bd+ (n = 21)	1.9	–0.6–4.4	18.6	15.2–21.5	1.4	–2.3–3.8	17.3	13.3–20.9	3.9	2.3–5.6	21.1	15.7–24.6
Bd– (n = 27)	1.5	1.1–6.2	16.8	15.3–19.9	0.2	–3.1–5.9	15.3	13.5–19.7	3.2	1.3–6.2	19.5	14.8–24.7
All sites South	1.6	–0.6–6.2	17.8	15.2–21.5	1.1	–2.3–5.9	16.2	13.3–20.9	3.5	1.3–6.2	20.2	14.8–24.7

Northern sites n = 49, Bd detected at 84% of sites; Southern sites, n = 48, Bd detected at 65% of sites.

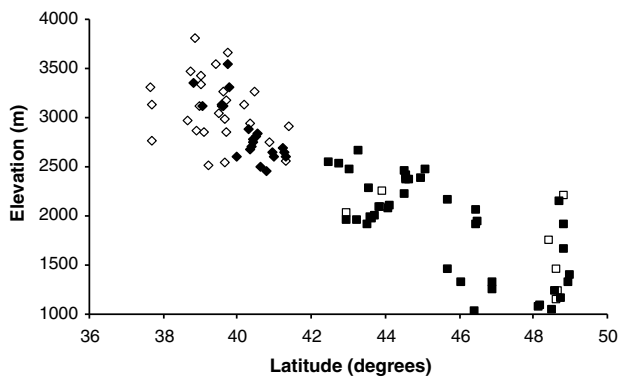


Fig. 2 – Clusters plotted relative to elevation and latitude illustrating that boreal toad sites (=clusters) at lower latitudes tend to be at higher elevations. Diamonds indicate southern clusters, squares indicate northern clusters; filled symbols are Bd positive clusters, open symbols are Bd negative clusters.

from low to high latitudes, geographical differences between the northern and southern regions are clear (Fig. 2), with sites in southern clusters (lower latitudes) occurring at higher elevations than sites in northern clusters. This pattern may reflect putative species differences predicted for *Bufo boreas* in the Rocky Mountains (e.g., Goebel, 2005). As expected, maximum and minimum temperatures at high elevation clusters were lower than at low elevation clusters (Table 3). Mean maximum temperatures at southern clusters where Bd was detected were at the lower limit of the optimal or “faster” growth zone (17–25 °C) as defined by Piotrowski et al. (2004). Mean maximum temperatures at southern clusters where the fungus was not detected were, on average, 2° colder.

Plots of the temperature data indicate that all clusters have daily minimum temperatures within the reduced growth zone (<17 °C) and 84% of these experience minimums <4 °C. Forty-two percent of clusters have daily maximum temperatures in the reduced growth zone. Mean maximum temperature at southern clusters during the active season was lower than the mean maximum temperature at northern

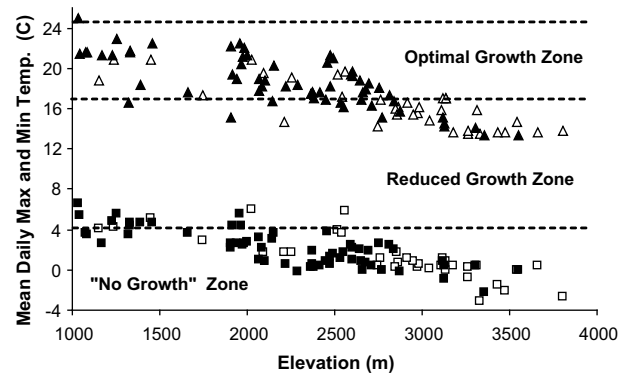


Fig. 3 – Mean daily maximum (triangles) and minimum (squares) temperatures at clusters. 10–25 °C is suitable for growth of *Batrachochytrium dendrobatidis*; growth is optimal between 17–25 °C and slower between 4–10 °C. (Piotrowski et al., 2004). Berger et al. (2004) reported that pathogenicity of Bd decreased at 23 °C.

clusters ($T = -7.238$, d.f. = 95, $P < 0.0001$) and the mean minimum temperature at southern clusters during the active season was lower than the mean minimum temperature at clusters in the north ($T = -5.783$, d.f. = 95, $P < 0.0001$) (Table 3). Of the clusters where Bd was detected ($n = 62$), 44% experienced daily mean maximum temperatures greater than 17 °C. In contrast, clusters where Bd was not detected ($n = 35$), only 11% experienced daily mean maximum temperatures greater than 17 °C (Fig. 3).

The model best supported by the data, as indicated by Akaike weights and ΔAIC_c (Table 4) was: Bd occurrence $\sim 4.083 + 0.960$ (Region) -0.006 (Elev) -0.051 (Julian Day) $+ 0.479$ (TempMax) -0.574 (TempMin) $+ 0.000$ (Elev \times Julian Day). Confidence intervals (95%) for estimates for TempMax and TempMin did not overlap zero (Table 5). The over-dispersion factor for the most parameterized model was 1.07.

Our cluster-level detection probability for Bd was >65% and the probability of a false negative (assigning a cluster as negative when it was really positive) was 11% (4 of 35). TempMax,

Table 4 – Models and AIC_c results

Model	Number of Parameters (K)	AIC _c	ΔAIC _c	Akaike weight
BdOcc ~ Elev + JulianDay + Elev × JulianDay + Region + TempMax + TempMin	8	112.494	0.000	0.652
BdOcc ~ Elev + JulianDay + Elev × JulianDay + Region + TempMax	7	115.179	2.685	0.170
BdOcc ~ Elev + JulianDay + Elev × JulianDay + Region	6	116.215	3.721	0.101
BdOcc ~ Elev + JulianDay + Elev × JulianDay + Region + TempMin	7	116.785	4.291	0.076

N = 97 for all models. Bd Occ = Bd Occurrence; Elev = Elevation in meters; Region = North or South; TempMax = mean maximum temperature; TempMin = mean minimum temperature. All models included Effort = number of samples collected for each cluster.

Table 5 – Estimates, standard error (S.E.) and 95% confidence intervals (C.I.) for variables in the model with the most support from the data (BdOcc ~ Elev + JulianDay + Elev × JulianDay + Region + Effort + TempMax + TempMin + Effort)

	Estimate	S.E.	Lower 95% C.I.	Upper 95% C.I.
Intercept	4.0829	12.0871	-19.3031	28.4256
Region	0.9599	0.9308	-0.8451	2.8498
Elevation	-0.0056	0.0039	-0.0140	0.0016
Julian Day	-0.0508	0.0519	-0.1569	0.0482
TempMax	0.4785	0.1981	0.1106	0.8955
TempMin	-0.5742	0.2633	-1.1201	-0.731
Elevation × Julian Day	0.0000	0.0000	-0.0000	0.0001
Effort	0.0191	0.0149	-0.0081	0.0541

TempMin, Elev and Effort were the variables least affected by a potential 11% misclassification of negative clusters. For these variables, the magnitude of the parameter estimate and the sign did not change. The mean value produced by the simulations was similar to the modeled parameter estimate. Estimates for other model parameters (Elev × Julian Day, Region, Julian Day, Intercept,) switched signs (<1%, <1%, 2%, and 22%) of the 1000 iterations, respectively). Therefore, the model is relatively robust to misclassification of negative clusters especially for the two variables of interest, TempMax and TempMin.

4. Discussion and conclusions

We found that *Bd* is widespread in the Rocky Mountains from northern Montana to southern Colorado. *Batrachochytrium dendrobatidis* was documented at a higher proportion of low versus high elevation boreal toad breeding sites. Our results suggest that, in the higher elevations of the temperate zone, the occurrence or persistence of *Bd* is limited by temperature which is constrained by elevation. The positive relationship between the occurrence of *Bd* and maximum daily temperatures indicates that, in the Rocky Mountains (1) maximum ambient temperatures rarely get warm enough for *Bd* to be fully successful and (2) there may be threshold temperatures (minima and maxima) that reduce the probability of *Bd* occurrence.

One-way to explain the patterns we observed is to examine fundamental niche models that predict the occurrence of *Bd* worldwide and suggest that temperature and precipitation are important in determining where the fungus may occur (Ron, 2005). Most studies (which are almost entirely from

tropical regions) have focused on maximum temperatures exceeding the tolerance levels of the fungus (e.g., Pounds et al., 2006; Drew et al., 2006). In the Rocky Mountains, ambient temperatures are rarely high enough to be lethal to the disease (i.e., 30 °C, Piotrowski et al., 2004) such that low maximum temperatures in the Rocky Mountains, particularly at higher elevations, may actually limit the success of *Bd*. Conversely, in Australia, frogs experiencing cooler temperatures may be less resilient to stressors and those farther from the equator (higher latitudes) carry more zoospores of *Bd* than frogs closer to the equator (Kriger et al., 2007). Kriger et al. (2007) also conclude that while optimal temperatures may occur at high-elevations in the tropics, high-elevations in temperate areas are too cold to support significant populations of *Bd*. Our data support this hypothesis.

Alternatively, ambient air temperature may not be a good predictor of *Bd* occurrence because it may not reflect water temperatures. For example, Seimon et al. (2006) argue that subfreezing temperatures and large diurnal temperature fluctuations do not limit the pathogenicity of *Bd*, positing instead that solar heating of small ponds may provide the thermal opportunities for *Bd* in an otherwise cool environment. Furthermore, ambient air temperatures are not necessarily correlated with operative body temperature of individual toads. Amphibians are capable of thermal regulation by microclimate selection and behavior and adult anurans are capable of clearing themselves of *Bd* when they can maintain elevated body temperatures for several hours (Woodhams et al., 2003). Boreal toads are able to raise body temperatures above the lethal limit of *Bd* (25 °C) by basking (Bartelt and Peterson, 2005) and they have been observed basking at high-elevations (Muths and Corn, 1997). However, we emphasize that the air temperature data used in our analysis are 35 year averages providing a view of climate rather than a short term weather snapshot and illustrating what the fungus would encounter over generations, likely affecting its persistence. Our study focuses on the patterns at a regional scale and does not attempt to predict situations at specific sites.

Another possible explanation for the low number of *Bd* observations at high-elevations is the relative topographic position of sites in a particular drainage. For example, in the Rocky Mountains, the fundamental niche model for *Bd* (Ron, 2005) predicts a mosaic pattern of occurrence. We suggest that this result is caused by drainage networks and steep topography in the region and that this feature may be key in predicting *Bd* occurrence. Whereas the absolute elevation affects environmental temperatures, the relative elevation and position within a particular drainage may also be a predictor of *Bd* infection in toad populations. Because *Bd* is an

aquatic pathogen it may have difficulty moving upstream in mountain systems and therefore be less likely to occur in headwater wetlands. For example, in Gunnison County, Colorado, boreal toads disappeared (coincident with *Bd* observations (Jackson, 2005)) from much of the lower East River drainage in the 1970s (Carey, 1993), while at the headwaters of the same drainage (Copper Creek tributary, 3446 m elevation), an extant toad population has tested negative for *Bd* since 2001 (Jackson, 2005). This site is >400 m higher than nearby sites where toads disappeared or are currently *Bd* positive; suggesting that relative elevation within a drainage plays a role in the potential for invasion by this fungus.

The effects of global climate change on species ranging from mammals to grasses are well documented and appear to be coherent in pattern (Parmesan and Yohe, 2003; Root et al., 2003) and regional manifestations of recent climate change are evident (e.g., Parmesan, 2006; Logan et al., 2003; Barber et al., 2000). Concerns are especially relevant to high latitude and high elevation locations because that is where effects on organisms are likely to be most dramatic (e.g., Schneider et al., 2007; Wagner, 2003). Pounds et al. (2006) discussed the importance of elevation with respect to changing environmental conditions ascribed to global warming and notes that “low temperatures as well as high ones may limit the impact of *Bd*.” More broadly, Kriger and Hero (2007) pointed out that temperatures vary across seasons, elevations and latitudes and that such variation, in concert with the sensitivity of amphibian immune systems and the thermal restrictions of *Bd*, are likely important in the ecology of this host–pathogen interaction. In North America, Longcore et al. (2007) report that ambient temperatures in the Northeastern United States vary between high enough and low enough to limit growth of *Bd*, further emphasizing the importance of local conditions. Our data provide evidence that the distribution or persistence of this disease in the Rocky Mountains is limited by temperature.

As global warming progresses, *Bd* may increase in occurrence and perhaps in the severity of its impact on Rocky Mountain amphibians. Predicted ambient temperatures at our study sites by 2100 are expected to fall within the optimal growth zone for *Bd*, reducing the temperature limitations to *Bd* growth and allowing possible range expansion of the pathogen to higher elevations. Specifically, predictions for the Rocky Mountains include warmer night-time temperatures, higher temperatures during the winter and reduced snow-packs (Wagner, 2003) resulting in a thermal regime at higher elevations that might be more conducive to *Bd* growth and persistence. At our study sites, average daily maximum temperatures at *Bd* positive clusters are predicted to increase at least 3.6 °C, while average daily minimum temperatures are predicted to increase at least 1.7 °C (Table 3). The average temperatures during the years of the study (2002–2004) are greater than the 35 year average temperatures (Table 3) suggesting that warming is already occurring.

The distribution of *Bd* in this region highlights a paradox. We found that the occurrence of *Bd* was lower in the southern (and higher elevation) portion of the study area, yet boreal toad populations in the Southern Rocky Mountains are declining (Corn et al., 1989; Carey, 1993; Stuart and Painter, 1994; Muths et al., 2003). In contrast, toad populations in

the northern portion of the study area appear to be stable. Possible explanations for this paradox include previous outbreaks of *Bd* resulting in resistant populations, or higher susceptibility to *Bd* with consequent local extirpations in the southern portion of the study area. For example, Briggs et al. (2005) point out that long-term persistence of *Bd* at a breeding pond may depend on some individuals surviving and carrying the fungus past metamorphosis (i.e., resistance). Susceptibility to chytridiomycosis may also vary with genetic differences among boreal toad populations. Recent genetic evidence, supported by habitat characteristics (e.g., elevational range), suggests that populations of boreal toads in the Southern Rocky Mountains represent a distinct species or subspecies (Goebel, 2005). In addition to resistance and susceptibility issues in the host, there may be differences in the pathogenicity of *Bd* strains within a region (Berger, 2005; Morgan et al., 2007). Alternatively, aspects of local environments may affect host–pathogen interactions differently. For example, toads in the Colorado Front Range inhabit environments with elevated atmospheric nitrogen deposition (Fenn et al., 2003).

The regional-level field data presented here demonstrate that the occurrence of *Bd* varies geographically. These results are particularly important given that this study focused on breeding sites for a single species and still revealed patterns. For example, temperature considerations are likely to be critical to understanding the distribution and prevalence of *Bd* and can play a role in the selection of sites for future salvage operations (sensu Mendelson et al., 2006) and reintroductions of threatened and endangered amphibians. Increased temperatures due to global warming are likely to reduce the environmental constraints acting on the fungus, allowing it to expand its distribution into high elevation areas. In conclusion, we suggest that it is low maximum temperatures and relative geography that may actually restrict the occurrence or persistence of this pathogen in the Rocky Mountains.

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