INTRODUCTION

Approximately one-third of all known amphibian species are currently listed as threatened (Stuart et al. 2004). Declines of many amphibian species are attributed to the disease chytridiomycosis (Berger et al. 1998), which has been implicated in mass mortalities, population declines and extinctions of amphibians around the world. In almost all cases, amphibian species that have disappeared or declined due to chytridiomycosis coexist with non-declining species. One reason why some species decline from chytridiomycosis and others do not may be interspecific differences in behaviour. Host behaviour could either facilitate or hinder pathogen transmission, and transmission rates in the field are likely to vary among species according to the frequency of factors such as physical contact between frogs, contact with infected water and contact with environmental substrates containing Batrachochytrium dendrobatidis. We tracked 117 frogs (28 Litoria nannotis, 27 L. genimaculata and 62 L. lesueuri) at 5 sites where B. dendrobatidis is endemic in the rainforest of tropical northern Queensland and recorded the frequency of frog-to-frog contact and the frequency of contact with stream water and environmental substrates. Frequency of contact with other frogs and with water were highest in L. nannotis, intermediate in L. genimaculata and lowest in L. lesueuri. Environmental substrate use also differed among species. These species-specific opportunities for disease transmission were correlated with conservation status: L. nannotis is the species most susceptible to chytridiomycosis-related declines and L. lesueuri is the least susceptible. Interspecific variation in transmission probability may, therefore, play a large role in determining why chytridiomycosis drives some populations to extinction and not others.

KEY WORDS: Batrachochytrium dendrobatidis · Amphibian chytrid fungus · Chytridiomycosis · Disease transmission · Behaviour · Frogs
chytridiomycosis and others do not may be inter-
specific differences in behaviour. Transmission can be
the driving force in the dynamics of infectious diseases
(Begon et al. 2002), and the long-term spread and per-
sistence of many diseases depends largely on the con-
tact rate between susceptible hosts and infectious
pathogens (Swinton 1998). This suggests that differ-
ences among species in opportunities for the trans-
mission of Batrachochytrium dendrobatidis may affect
their susceptibility to chytridiomycosis. Despite the
importance of transmission in the epidemiological pro-
cess, very little is known about transmission of
Batrachochytrium dendrobatidis, particularly in the field. The only route of
transmission that has been established by controlled
experimentation is transmission via contact with water
that was previously in contact with infected tadpoles or
adults (Berger et al. 1998, Retallick 2002, Parris & Cor-
nelius 2004, Rachowicz & Vredenburg 2004). However,
infected zoospores are present on, and can be recov-
ered from, the skin surfaces of infected animals
(Berger et al. 1998, Pessier et al. 1999), and B. den-
drobatidis DNA has been detected on wet rocks at a site
during an epidemic (Lips et al. 2006). It therefore
seems likely that transmission also occurs via contact
with infected individuals or contaminated environ-
mental substrates.

Host behaviour could either facilitate or hinder
pathogen transmission, and transmission rates in the
field are likely to vary among species depending on the
frequency of behaviours such as physical contact
between frogs, contact with infected water and contact
with environmental substrates that may contain Batra-
chochytrium dendrobatidis. In this study, we tracked
3 species of rainforest stream frogs at 5 sites and
recorded their frequency of contact with other frogs,
contact with stream water and contact with environ-
mental substrates.

MATERIALS AND METHODS

The study was conducted at 5 tropical rainforest sites
in northern Queensland, Australia: Birthday Creek,
Paluma State Forest (18° 58’ 54” S, 146° 10’ 02” E, 800 m
above sea level [a.s.l.]), Python Creek (17°46’S, 145°35’E, 200 m a.s.l.), an unnamed creek (17°48’S, 145°41’E, 70 m a.s.l.) in Tully Falls Forest Reserve, an
unnamed creek in Kirrama State Forest (18°11’S, 145°52’E, 200 m a.s.l.) and Frenchman Creek, in Woo-
roonooran National Park (17°20’S, 145°55’E, 20 to
100 m a.s.l.). All sites were relatively undisturbed rain-
forest streams with rocky beds. All streams had pools
and riffles, and most sites had a number of waterfalls.

 Frogs of 3 species were tracked: the stony creek frog
Litoria lesueuri, which has not experienced population
declines (IUCN ‘least concern’; see below), the green-
eyed tree frog L. genimaculata, which declined and
then recovered (IUCN ‘least concern’; however, Aus-
tralian populations of this species are considered to be
‘near threatened’) and the waterfall frog L. nannotis,
which has experienced large and long-lasting popula-
tion declines (IUCN ‘endangered’; McDonald & Alford
1999, McDonald 2002, McDonald et al. 2005, IUCN et
al. 2006). Recently, the taxonomy of the L. lesueuri
group has been revised (Donnellan & Mahony 2004). Two species, L. jungguy and L. wilcoxii, occur in symp-
pathy at the study sites, hybridise, and are indistin-
guishable on the basis of morphology (Donnellan &
Mahony 2004). Population declines have not been ob-
erved in the region for either species (McDonald &
Alford 1999, McDonald et al. 2005). We therefore
continue to refer to the study population as L. lesueuri,
while recognising that the population contains 2 mor-
phologically indistinguishable species. All species of
frogs tracked were large- to medium-sized hylids
(males 5.4 to 12.5 g, females 6.5 to 41.3 g), and were
tracked using either radio-telemetry or harmonic di-
rection finding.

 Only frogs weighing >11 g were tracked via radio-
telemetry. Radio transmitters (Models BD-2N and
BD-2NT; Holohil Systems; weighing approximately
0.67 g, including a harness and with a battery life of
approximately 3 wk) were attached to a harness made
of silicone tubing, designed to minimise restrictions on
movement and avoid causing discomfort to the frog.
This harness/transmitter combination was placed
around the waists of frogs. Frogs that were too small to
be radio-tracked, and a number of larger individuals,
were tracked using harmonic direction finding (Lang-
kilde & Alford 2002). This required attachment of a
small diode with whip antenna to the same specially
designed harness (total weight approximately 0.23 g),
which was then placed around the waist of a frog.
Tracking devices were fitted in situ, and frogs were
released at the point of capture after <5 min of hand-
ing. Frogs wearing either tracking device always had
harnesses and associated equipment that weighed
<6% of their total body weight, just over half the rec-
ommended maximum relative weight for an attached
tag (10% of the body weight; Richards et al. 1994).

 Frogs fitted with radio-transmitters were tracked
using a Telonics TR-4 Tracking Receiver (Telonics;
2004 wet season only) and a HABIT Research HR2500
Osprey VHF Receiver (HABIT Research); we used a
3-element folding Yagi antennae with both receivers
(A.F. Antronics). Frogs fitted with diodes were tracked
using a portable RECCO R5 transmitter–receiver
(Recco Rescue Systems). The system consists of a
hand-held device that acts as both the transmitter and
receiver, and a battery and earphones. Tags were self-
built using commercial germanium diodes (see Langkilde & Alford 2002 for a description of methodology).

Surveys lasted 16 d and were conducted in July to September (the cool/dry season) and February to March (the warm/wet season) for each species at 2 sites, except for Paluma, which was only surveyed in the warm/wet season. *Litoria genimaculata* was tracked at Paluma in late 2003, *L. genimaculata* and *L. nannotis* were tracked simultaneously at Kirrama and Tully Gorge during 2004, and *L. lesueuri* was tracked at Lower Tully and Babinda during 2005. During surveys, the location of each frog was determined once during the day (09:00 to 18:00 h) and once at night (19:00 to 04:00 h).

We recorded instances of frog-to-frog contact when we observed direct skin-to-skin contact between frogs and when aggregations of >3 individuals occupied an area of less than 0.3 × 0.3 m. At each observation of contact between frogs, we recorded the species involved and the nature of the contact (i.e. amplexus, sharing a retreat site or a nocturnal perch site). We recorded instances of contact with water when we observed at least part of the frog in contact with stream water, or when the frog had crossed the stream since the last observation, and there was no way for the frog to cross the stream without entering the stream water (i.e. the frog could not have crossed the stream using dry rocks or overhead vegetation). This provides an estimate of the minimum rate of contact with water, since frogs may come into contact with water without crossing streams. During all observations, we recorded the substrate type with which the ventral surface of the frog was in contact. Substrate type was classified into 6 categories: bare ground, dry rock, leaf litter, vegetation, wood (dead wood such as logs or sticks) and wet rock. All these records represent possible contact between host and pathogen, or opportunities for transmission.

We excluded data from the night following tag replacement in July 2002. Any effects are unlikely to persist after the first night of tag attachment (Rowley & Alford 2007). To avoid pseudoreplication or biasing our results towards frogs that were located more frequently, we used individuals as replicates and compared summary statistics calculated for each animal. To analyse frog-to-frog contact and contact with water, we performed Kruskal-Wallis tests to detect differences among species and separate Mann-Whitney *U*-tests for each species to detect differences with respect to season and sex. We used Bonferroni adjustments to control Type I error rates. To analyse substrate use, we examined diurnal and nocturnal data separately, due to obvious differences between the diurnal and nocturnal behaviour of all species. We then performed multiresponse permutation procedures (MRPP) using Blossom statistical software (Cade & Richards 2005). We performed the analysis in a stepwise fashion, testing for differences between species, sexes, seasons and sites, in that order. If a difference was detected between groups, the next analysis was performed on each of those groups separately. We used Monte Carlo hypothesis testing, with 20000 iterations.

RESULTS

A total of 117 frogs (28 *Litoria nannotis*: 6 M, 22 F; 27 *L. genimaculata*: 15 M, 12 F; and 62 *L. lesueuri*: 32 M, 30 F) were tracked during the study period. Approximately equal numbers of frogs were tracked using both tracking techniques; 61 (52%) were tracked via radio-telemetry. Throughout the survey periods, a total of 2111 fixes or locations of individual frogs were obtained. On average, each frog was located 17 times (minimum 5, maximum 29).

The frequency of contact between frogs differed significantly among species (Table 1, Fig. 1). Individual *Litoria nannotis* were in contact with other frogs during an average 28% of observations, while *L. genimaculata* and *L. lesueuri* were only in contact with other frogs during an average of 3 and 2% of observations, respectively. Type of contact differed among species (Table 2); however, in all species, direct contact was always between conspecifics. The frequency of physical contact differed between seasons only in *L. nannotis*, which was more often in contact with conspecifics in the warm/wet season (Table 1, Fig. 1). There was no difference in the frequency of physical contact between sexes in *L. nannotis* and *L. genimaculata*, but

<table>
<thead>
<tr>
<th>Test statistic (χ² or Z)</th>
<th>p</th>
<th>Test statistic (χ² or Z)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>18.573</td>
<td>&lt;0.001*</td>
<td>60.492</td>
</tr>
<tr>
<td>Season</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>L. lesueuri</em></td>
<td>-1.008</td>
<td>0.313</td>
<td>-3.607</td>
</tr>
<tr>
<td><em>L. genimaculata</em></td>
<td>-0.789</td>
<td>0.648</td>
<td>-0.960</td>
</tr>
<tr>
<td><em>L. nannotis</em></td>
<td>-2.775</td>
<td>0.010*</td>
<td>-1.004</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>L. lesueuri</em></td>
<td>-2.636</td>
<td>0.008*</td>
<td>-0.993</td>
</tr>
<tr>
<td><em>L. genimaculata</em></td>
<td>-0.395</td>
<td>0.829</td>
<td>-0.541</td>
</tr>
<tr>
<td><em>L. nannotis</em></td>
<td>-0.852</td>
<td>0.460</td>
<td>-1.887</td>
</tr>
</tbody>
</table>

Table 1. *Litoria* spp. Differences in direct and indirect contact among species (Kruskal-Wallis tests), between seasons and between sexes (Mann-Whitney *U*-tests). *Significance at the Bonferroni-adjusted 0.05 level.
female *L. lesueuri* were more often in contact than male *L. lesueuri* (Table 1).

The frequency of contact with stream water also differed significantly among species, and was highest in *L. nannotis* (Table 2, Fig. 1). Overall, *L. nannotis* was observed in stream water or had crossed the stream since the last observation during 84% of observations. In contrast, *L. genimaculata* and *L. lesueuri* were observed in contact with water during an average of only 16 and 13% of observations, respectively. Most contact with water in *L. genimaculata* and *L. lesueuri* occurred at night, when the species were active and crossing the stream, in contrast to *L. nannotis*, which was often inactive and in contact with water for long periods of time (>12 h). We observed a number of these individuals crossing the stream, and in all cases these crossings were relatively brief (<2 h). Rates of contact with stream water differed between seasons only in *L. lesueuri*; they were higher in the wet season (Table 2, Fig. 1) and did not differ with sex for any species (Table 2).

Substrate use differed among species and between diurnal and nocturnal surveys (Tables 3 & 4). *Litoria lesueuri* were most often on bare ground or leaf litter during diurnal surveys, moving onto a wider range of substrate types at night. *L. genimaculata* were most often on vegetation, generally in the forest canopy, during both diurnal and nocturnal surveys. In instances when we could visually locate individual *L. genimaculata* in the canopy, frogs were sitting on leaves at the ends of branches. *L. nannotis* was most commonly on wet rock, with frogs typically sheltering under large boulders in the stream during the day and often moving on to terrestrial vegetation and dry rock at night.

Substrate use differed between sexes in *Litoria lesueuri* and *L. nannotis* (Tables 3 & 4). In *L. lesueuri*, substrate use differed subtly between sexes, but, in *L. nannotis*, males were almost always on wet rock, while females were found more often on other substrates, including vegetation and dry rock (Table 4). Substrate use differed with season for *L. lesueuri* males and for *L. genimaculata* (Tables 3 & 4). In the cool/dry season, *L. lesueuri* males spent more time on bare ground and less time on dry rocks and *L. genimaculata* spent more time on vegetation, generally in the forest canopy, during both diurnal and nocturnal surveys. In instances when we could visually locate individual *L. genimaculata* in the canopy, frogs were sitting on leaves at the ends of branches. *L. nannotis* was most commonly on wet rock, with frogs typically sheltering under large boulders in the stream during the day and often moving on to terrestrial vegetation and dry rock at night.

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Rowley & Alford: Frog behaviour may affect chytridiomycosis transmission

DISCUSSION

The frequencies of frog-to-frog contact and contact with stream water and different environmental substrates differed significantly among species. As these factors should affect rates of transmission of *Batrachochytrium dendrobatidis* via known or highly likely routes, transmission rates should also vary among species.

The frequency of contact between frogs was highest in *Litoria nannotis*, primarily because the species aggregates at retreat sites. The formation of aggregations by host populations promotes contact between individuals, and in many other host–pathogen systems it is positively correlated with both the prevalence and intensity of contact-transmitted parasites (Anderson & May 1979, Hoogland 1979, Brown & Brown 1986, Coté & Poulin 1995, Ezenwa 2004). Outbreaks of disease are also most commonly observed in aggregations of individuals (Vermeer 1969, Wobeser et al. 1979). In contrast, predominantly solitary or non-social species such as *L. lesueuri* and *L. genimaculata* will come into direct contact almost exclusively for reproduction, and it is likely that, for these species, most instances of direct transmission occur at this time (Loehle 1995). This may be especially true for the transmission of *Batrachochytrium dendrobatidis* in *L. lesueuri*, as *L. lesueuri* comes into contact with stream water infrequently. Therefore, while traditional models of direct transmission have assumed that contact rate is directly proportional to host population size or density (Anderson & May 1979, 1981, Watanabe 1987), the present study confirms that rates of contact between individuals may be almost entirely independent of population size due to host behaviour (Loehle 1995, McCallum et al. 2001, Ezenwa 2004).

In all 3 species, contact between frogs was always between conspecifics, providing almost no opportunities for cross-species pathogen transmission. Therefore, contact with stream water and possibly other environmental substrates that serve as reservoirs are likely to be the main source of transmission between species in this system. Contact with stream water was more frequent in *L. nannotis*, which was in contact with the stream during the majority of observations. Increased frequency of contact with stream water is likely to increase the rate of transmission of *Batrachochytrium dendrobatidis* in a species for a number of reasons. First, *B. dendrobatidis* is known to be transmitted via water in the laboratory (Berger et al. 1998, Parris & Cornelius 2004, Rachowicz & Vredenburg 2004) and in field enclosures (Retallick 2002). Second, *B. dendrobatidis* zoospores are aquatic, highly sensitive to desiccation (Johnson et al. 2003), and can sur-

Table 4. *Litoria* spp. Substrate use. Mean percent (and range) of observations on each substrate type, using individual frogs as replicates. Only categories that are significantly different from each other are shown

<table>
<thead>
<tr>
<th>Substrate Type</th>
<th>Female</th>
<th>Male</th>
<th>Female</th>
<th>Male</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare ground</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diurnal</td>
<td>19.34 (0–66.7)</td>
<td>39.01 (0–100)</td>
<td>9.1 (0–30.8)</td>
<td>0</td>
</tr>
<tr>
<td>Nocturnal</td>
<td>8.1 (0–33.3)</td>
<td>26.67 (0–66.7)</td>
<td>10.4 (0–28.6)</td>
<td>0</td>
</tr>
<tr>
<td>Dry rock</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diurnal</td>
<td>3.8 (0–60)</td>
<td>1.8 (0–25)</td>
<td>8.3 (0–33.3)</td>
<td>25.6 (0–100)</td>
</tr>
<tr>
<td>Nocturnal</td>
<td>8.2 (0–50)</td>
<td>2.3 (0–33)</td>
<td>20.2 (0–60)</td>
<td>21.4 (0–69.2)</td>
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<tr>
<td>Leaf litter</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diurnal</td>
<td>63.6 (0–100)</td>
<td>55.3 (0–100)</td>
<td>52.9 (0–100)</td>
<td>5.9 (0–33.3)</td>
</tr>
<tr>
<td>Nocturnal</td>
<td>39.9 (13.3–85.7)</td>
<td>16.4 (0–80)</td>
<td>30.8 (0–61.5)</td>
<td>2.8 (0–25)</td>
</tr>
<tr>
<td>Vegetation</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Diurnal</td>
<td>5.6 (0–41.7)</td>
<td>3.4 (0–22)</td>
<td>5.9 (0–40)</td>
<td>63.7 (0–100)</td>
</tr>
<tr>
<td>Nocturnal</td>
<td>30.5 (0–80)</td>
<td>48.8 (0–100)</td>
<td>25.2 (0–50)</td>
<td>72.7 (23.1–100)</td>
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<tr>
<td>Wood</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diurnal</td>
<td>2.7 (0–33.3)</td>
<td>0.5 (0–11.1)</td>
<td>14.7 (0–100)</td>
<td>0</td>
</tr>
<tr>
<td>Nocturnal</td>
<td>11.4 (0–16.7)</td>
<td>5.8 (0–44.4)</td>
<td>5.2 (0–20)</td>
<td>0</td>
</tr>
<tr>
<td>Wet rock</td>
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<td></td>
</tr>
<tr>
<td>Diurnal</td>
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<td>0</td>
<td>0</td>
<td>4.8 (0–22.2)</td>
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<tr>
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<td>0</td>
<td>8.3 (0–60)</td>
<td>3.1 (0–22.2)</td>
</tr>
</tbody>
</table>

Time on dry rock and less time on leaf litter, vegetation and bare ground (Table 4).
vive and remain infective in the laboratory for at least 7 wk in sterile lake water (Johnson & Speare 2003). Additionally, tadpoles often have high infection prevalence, may not be susceptible to the pathogen, and can persist in the stream environment for several years (Woodhams & Alford 2005, Rachowicz et al. 2006), thereby providing a likely continual source of zoospores to the stream. Lastly, amphibian population declines have been greatest in species with strong associations with streams (Williams & Hero 1998, McDonald & Alford 1999, Lips et al. 2003, Hero et al. 2005), and, while purely terrestrial species may be infected with *B. dendrobatidis* (Lips et al. 2006), they do not typically experience population declines or experience reduced rates of decline, even when in sympatry with rapidly declining species (Williams & Hero 1998, McDonald & Alford 1999, Hero et al. 2005, Lips et al. 2006).

Different species used different environmental substrates, and, in some cases, substrate use varied between sexes and seasons as well. Because the abundance and composition of other chytrids differ among macrohabitats (i.e. vegetation type; Letcher et al. 2004) and microhabitats (i.e. with distance from moss; Letcher & Powell 2002), it is highly probable that the abundance of *Batrachochytrium dendrobatidis* zoospores also differs among environmental substrates, and, hence, exposure to *B. dendrobatidis* probably differs among species and even between sexes and seasons for some species. However, there are a large number of uncertainties in quantifying the risks associated with contact with stream water or environmental substrates. There is currently no information on the relative abundance of zoospores in the environment, and we do not yet know the relative importance of different areas of the stream or different environmental substrates for the persistence and transmission of *B. dendrobatidis* in the field. While *B. dendrobatidis* DNA has been detected on environmental substrate samples during epidemics (Lips et al. 2006), it is not known whether these substrates contain viable zoospores. In addition, *B. dendrobatidis* DNA was not detected in environmental substrate samples taken at *Litoria lesueuri* retreat sites, where *B. dendrobatidis* is endemic (Rowley et al. 2007). More work to determine how important environmental substrates may be as reservoirs for infective zoospores is urgently required.

Frequency of contact with other frogs or with water differed between seasons only in *Litoria lesueuri*, which was more often in contact with water during the wet season. There was little evidence that the frequency of contact with other frogs or with water differed between sexes for any of the species. In other study species, sex may greatly influence opportunities for disease transmission. For example, host behaviour has been implicated in the differential survival of *Bufo boreas* during *Batrachochytrium dendrobatidis* outbreaks (Carey et al. 2006). In this species, adult males spend several weeks in frequent direct contact with other males and in continuous contact with water during the breeding season, while females are thought to spend <1 d at breeding sites (Carey et al. 2006). Perhaps because of this, adult females appear to live longer than adult males during outbreaks of *B. dendrobatidis* (Carey et al. 2006).

In addition to frequency of contact between frogs, with stream water or with environmental reservoirs, the duration of contact is likely to be important. In the laboratory, the duration of exposure to *Batrachochytrium dendrobatidis* can influence the probability of successful transmission and the speed of disease progression, with longer exposures to *B. dendrobatidis* resulting in shorter average survival times in *Bufo boreas* (Carey et al. 2006). In our study species, the majority of contact situations with water for *Litoria lesueuri* and *L. genimaculata* occurred briefly when crossing a stream, while *L. nannotis* spent all day and often all night in contact with stream water. In addition, *L. nannotis* often returned to the same retreat sites, which were often shared with a large number of conspecifics, after nocturnal excursions (author’s unpubl. data). Such behaviour is likely to increase the opportunities for transmission in *L. nannotis*, as parasites may accumulate in the hosts’ environment over time (Altizer et al. 2000). Species that rarely return to the same diurnal retreat sites, such as *L. lesueuri* (author’s unpubl. data), may reduce their chances of infection. Although *B. dendrobatidis* was not detected at retreat sites of *L. nannotis* and *L. lesueuri* at several of our sites (Rowley et al. 2007), when infection prevalence and intensity is high, the duration of time spent at retreat sites with a certain substrate type (i.e. wet rock) may be important.

The dose of *Batrachochytrium dendrobatidis* zoospores that a frog encounters may also influence the probability and outcome of infection in frogs. In *Bufo boreas*, high doses of *B. dendrobatidis* result in shorter average survival times (Carey et al. 2006). This pattern appears typical of other host–pathogen systems, with higher infective doses leading to higher mortality rates and decreased host survival times (van Beek et al. 2000, Arthurs & Thomas 2001, Brunner et al. 2005). Because the intensity of infection by *B. dendrobatidis* must reach a particular threshold of zoosporangia before individuals succumb to chytridiomycosis, larger inocula are likely to reach lethal levels sooner (Carey et al. 2006). It is currently not known which behaviours would expose frogs to the highest concentration of zoospores, although direct contact with highly infected frogs seems likely to expose individuals to the highest
zooospore concentrations. Contact with environmental substrates may also expose frogs to high zooospore concentrations if *B. dendrobatidis* is able to grow on these substrates or if long residence times of infected frogs at retreat sites lead to high concentrations of zooospores in the environment.

In this study, it is likely that we underestimated the frequency of contact in all frog species. As frogs were not observed continuously, individuals may have briefly come into contact with other individuals, or with the stream, without this being recorded, particularly at night, when frogs were active. While increasing the frequency of surveys would allow a more accurate estimate of opportunities for transmission, the relative differences between species are unlikely to change. Increasing survey frequency would also increase the probability of disturbing frogs and hence influencing their behaviour.

Despite its importance, contact between host and pathogen is only one part of the entire sequence of host–parasite interactions, beginning with pathogen survival in the environment and terminating in either host mortality or pathogen elimination. Even when a pathogen and its host come into contact with each other, successful invasion of the host by the pathogen is not guaranteed. For example, many fungal pathogens require very specific environmental conditions in order to be infective, and low humidity may completely prevent fungal pathogens from infecting potential hosts, regardless of how many spores are in contact with a potential host (Hajek & St Leger 1994). Other factors such as host microenvironment selection may also eliminate infection once it occurs, since exposure to temperatures of 37°C for <6 h can eliminate infection from captive *Litoria chloris* (Woodhams et al. 2003). Timing of transmission may therefore be extremely important. In particular, as host mortality due to chytridiomycosis is highest in winter, exposure to *Batrachochytrium dendrobatidis* during the winter months may have greater consequences for host survival.

In summary, due to behavioural differences, *Litoria lesueuri*, *L. genimaculata* and *L. nannotis* differ greatly in their rates of contact with other frogs, stream water and different environmental substrate types. Because these rates are likely to affect their levels of exposure to *Batrachochytrium dendrobatidis*, the transmission of *B. dendrobatidis* is likely to also differ between species. Frequency of frog-to-frog contact and contact with water are correlated with the conservation status of the frogs; the highest rates of contact between frogs and with water occurred in *L. nannotis*, the species most susceptible to chytridiomycosis-related declines, intermediate rates occurred in *L. genimaculata*, the species with intermediate susceptibility to decline, and the lowest rates occurred in *L. lesueuri*, the species least susceptible to chytridiomycosis-related declines. Interspecific variation in transmission may therefore play a role in determining why chytridiomycosis drives some populations to extinction and not others.

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