

Amphibian ranaviruses in Europe: important directions for future research

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Abstract

Ranaviruses are an emerging group of pathogens capable of infecting all cold-blooded vertebrates. In Europe, ranaviruses pose a particularly potent threat to wild amphibian populations. Since the 1980s research on amphibian-infecting ranaviruses in Europe has been growing. The wide distribution of amphibian populations in Europe, the ease with which many are monitored, and the tractable nature of counterpart ex situ experimental systems have provided researchers with a unique opportunity to study many aspects of host–ranavirus interactions in the wild. These characteristics of European amphibian populations will also enable researchers to lead the way as the field of host–ranavirus interactions progresses. In this review, we provide a summary of the current key knowledge regarding amphibian infecting ranaviruses throughout Europe. We then outline important areas of further research and suggest practical ways each could be pursued. We address the study of potential interactions between the amphibian microbiome and ranaviruses, how pollution may exacerbate ranaviral disease either as direct stressors of amphibians or indirect modification of the amphibian microbiome. Finally, we discuss the need for continued surveillance of ranaviral emergence in the face of climate change.

Key words: microbiome, host–microbe interactions, wildlife disease, ecotoxicology, European amphibians

OPEN ACCESS

Citation: Campbell LJ, Pawlik AH, and Harrison XA. 2020. Amphibian ranaviruses in Europe: important directions for future research. FACETS 5: 598–614. doi:[10.1139/facets-2020-0007](https://doi.org/10.1139/facets-2020-0007)

Handling Editor: David Lesbarrères

Received: February 18, 2020

Accepted: May 11, 2020

Published: August 6, 2020

Note: This paper is part of a Collection titled “Ranavirus research: 10 years of global collaboration”.

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Published by: Canadian Science Publishing

Introduction

In 1979 a Scandinavian population of captive Atlantic cod (*Gadhus morhua*) began to exhibit elevated levels of morbidity and mortality. Affected fish exhibited a novel condition characterised by severe skin ulcerations, and investigation of the causes behind this disease outbreak implicated a virus belonging to the group *Iridoviridae* (Jensen et al. 1979). Since the advent of modern molecular techniques, researchers have revisited this case and genomic analysis of the isolated viruses has shown that this outbreak was, in fact, the first recorded incidence of ranavirosis (the often lethal disease caused by ranaviruses) in Europe (Ariel et al. 2010). Despite the importance of aquaculture and fisheries to many European economies (Crilly and Esteban 2013), the viral diversity and host range of ranaviruses infecting farmed and wild fishes in Europe has received only sporadic research effort over the last four decades (see recent reviews by Price et al. 2017b; Allain and Duffus 2019). However, fish are not the only group of vertebrates impacted by ranavirosis in Europe; ranaviruses are also known to establish infections in European herpetofauna (Cunningham et al. 1997; Balseiro et al. 2009). To date, reports of ranaviral infection and disease in reptiles are rare (Marschang 2011; Price et al. 2014, 2017b), but ranavirosis is considered to be an emerging infectious disease of

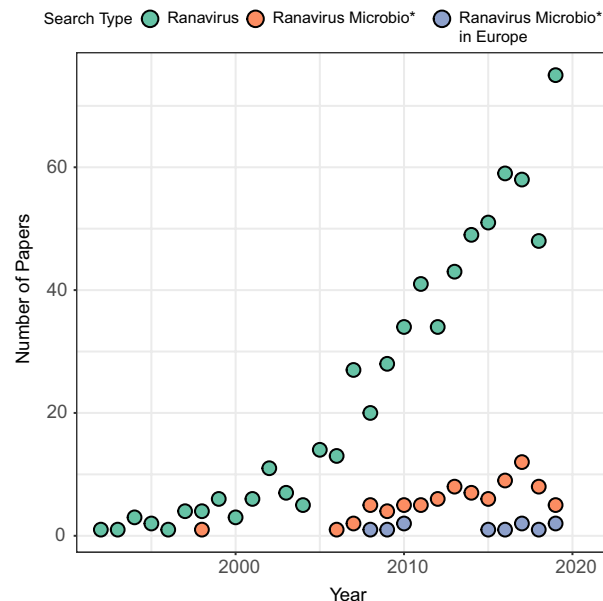


Fig. 1. Literature trends of research on ranavirus from a Scopus search, yielding number of papers per year when searching title, abstract, and keywords. Research on ranavirus has been increasing year on year since the 1990s (search term “ranavirus”, green circles). However, of those only a small subset represent distinct research on microbiomes (“ranavirus microbio*”, orange circles), though these will also include microbiology studies. Finally, of those, only a small fraction represent research on European hosts (blue circles).

European amphibians and is believed to pose a significant conservation threat (Daszak et al. 1999; Earl and Gray 2014; Campbell et al. 2018a).

The first incidences of ranavirosis in European amphibians were reported by members of the public in the United Kingdom during the late 1980s (Cunningham et al. 1997). Since that time, research effort devoted to these deadly pathogens has slowly grown both globally and in Europe (Fig. 1). It is currently accepted that there are two primary lineages of ranavirus circulating within European amphibians, the Frog Virus 3 (FV3)-like and the common midwife toad virus (CMTV)-like viruses (reviewed in Price et al. 2017a). Although there is overlap in the geographical range of these viruses, each is apparently impactful in a particular continental region, separating the study of amphibian ranaviruses in Europe into two distinct theatres: continental Europe and the islands that comprise the United Kingdom. The distribution and severity of ranavirus outbreaks in Europe have been given in-depth treatment in two recent reviews (Price et al. 2017a; Allain and Duffus 2019). Our goal in this article is not to supplant those works. Instead, here we highlight gaps in our knowledge concerning the forces shaping interactions between amphibian hosts and ranaviruses, and predictors of the severity of ranavirosis outbreaks, that should be prioritised for research. First, we present a brief summary of the key research findings from each of the main European theatres, before highlighting important questions that remain to be answered and potential avenues of future investigation.

Ranavirosis in the United Kingdom

The United Kingdom is densely populated, resulting in significant overlap between the human population and wildlife (Fuller and Gaston 2009). As a result, many United Kingdom amphibian populations reside in small water bodies, such as urban garden ponds, where they are easily observed by humans. In the late 1980s members of the British public began to report unusual mass mortality in

such populations of the European common frog (*Rana temporaria*; [Cunningham et al. 1997](#)). Deceased adult *R. temporaria* were found exhibiting severe skin ulceration and hemorrhaging of the internal organs ([Cunningham et al. 1997](#)). The Frog Mortality Project (FMP) was established in 1992 by scientists at the Zoological Society of London to collate and investigate these reports of disease incidence ([Cunningham et al. 1997](#)). The FMP solicited reports of frog mortality from the general public using an outreach campaign and facilitated the collection and post-mortem examination of carcasses collected at the site of reported outbreaks. Pathological and histological examination revealed that a ranavirus closely resembling FV3 was the likely cause ([Cunningham et al. 1997](#)). Comparative molecular analysis showed that this virus was highly similar to North American isolates of FV3 and was likely introduced into the United Kingdom via the international trade in amphibians and fish ([Hyatt et al. 2000](#)).

The FMP has generated a large archive of both amphibian tissue samples and associated viral isolates. Genomic analyses of ranavirus isolates collected as part of the FMP have shown that not only were there at least two distinct introduction events of FV3 into the United Kingdom ([Price et al. 2016, 2017a](#); [Duffus et al. 2017](#)) but also that an additional ranaviral species, CMTV, has been present in the United Kingdom since as early as 1995 (the earliest known detection of CMTV; [Price et al. 2017a](#)). Despite the presence of CMTV, it is believed that all records of ranavirosis in the United Kingdom are associated with FV3 ([Price et al. 2016, 2017a](#)). In stark contrast to ranavirosis outbreaks caused by FV3 elsewhere in the world, which often exhibit an extensive host range in terms of species and life stages impacted, mortality due to ranavirosis in the United Kingdom is almost entirely limited to *R. temporaria* ([Duffus et al. 2013](#); [Allain and Duffus 2019](#)) and is exclusively observed in post-metamorphic animals ([Duffus et al. 2013](#)).

Through the collection of disease incidence records the FMP has also spawned a database of *R. temporaria* populations that reside on private property and whose health is continually visually monitored by property owners ([Teacher et al. 2010](#); [Lawson et al. 2015](#)). With the help of these “citizen scientists” researchers in the United Kingdom have been able to establish a network of *R. temporaria* populations of known disease history, allowing for several comparative and (or) longitudinal studies that have advanced our understanding of interactions between ranaviruses and their hosts at the individual and the population level ([Lawson et al. 2015](#)). Research within this network of comparative populations has shown that the emergence of ranavirosis has resulted in a reduction of *R. temporaria* numbers by an average of over 80% in infected populations, though local extinction and recovery have also been documented ([Teacher et al. 2010](#)). Additionally, ranavirosis appears to have (i) altered the population genetics ([Teacher et al. 2009a](#)), (ii) reversed the mating system ([Teacher et al. 2009b](#)); and (iii) truncated the age structure of impacted *R. temporaria* populations ([Campbell et al. 2018a](#)). Applying predictive models to disease incidence reports and their associated environmental metadata has also revealed that the spread of FV3 within the United Kingdom is very likely facilitated by human translocation and pathogen pollution ([Price et al. 2016](#)) and that frequency and severity of ranavirosis outbreaks is tightly linked to the climactic conditions ([Price et al. 2019](#)). Data drawn from these incidence reports have also revealed that the occurrence and severity of ranavirosis outbreaks are correlated with host population density, the presence of secondary host species (hosts which can become infected with and shed virus but are themselves asymptomatic, including the common toad (*Bufo bufo*) and various species of fish), and the use of garden chemicals ([North et al. 2015](#)). Epidemiological models have demonstrated that the persistence of ranaviruses within *R. temporaria* populations can be maintained via exclusively adult to adult contact transmission ([Duffus et al. 2019](#)); however, the drivers of the distribution and intensity of ranavirus infection in the United Kingdom remain poorly understood. Though many monitored common frog populations in the United Kingdom appear to have never suffered outbreaks of ranavirosis, recent research detected the presence of FV3-like ranavirus in frogs originating from all

populations studied, including those with no history of observable ranavirosis (Campbell et al. 2018b). Critically, this suggests that ranaviruses may be ubiquitously distributed within *R. temporaria* populations in the United Kingdom, but that whether or not a population develops ranavirosis depends on some additional cryptic factors that promote infection of hosts and cause those infections to reach clinical or lethal thresholds.

For example, there is growing evidence suggesting a potential link between the occurrence of ranavirosis and composition of bacterial communities residing on the amphibian skin. Differing bacterial communities have been detected on the skin of frogs originating from populations with endemic ranavirosis versus frogs from populations with no history of disease, based on both transcriptomic (Campbell et al. 2018b) and genomic DNA (Campbell et al. 2019) sequences. Beyond these field-based studies, the outcome of acute ranaviral infection has been shown to be impacted by the diversity of the amphibian skin microbiome during an experimental study that showed that metamorphic *R. temporaria* with a depauperate bacterial microbiome were more likely to succumb to infection with FV3 than were metamorphic *R. temporaria* possessing a skin microbiome with higher species richness (Harrison et al. 2019).

Ranavirosis on continental Europe

Despite a long history of ranaviruses impacting fish on continental Europe (Jensen et al. 1979; Ariel et al. 2010), since mid-2000 ranavirosis has also been emerging in amphibians. The primary causes of ranavirus-induced mortality on continental Europe appear to be several strains of CMTV (reviewed by Price et al. 2017b). CMTV are considered a distinct species complex within the genus *Ranavirus* (Chinchar et al. 2017) and were first discovered during investigations of a large mortality event that occurred at the Picos de Europa National Park, Spain, in 2007 (Balseiro et al. 2009). This outbreak impacted the larval life-stage of the common midwife toad (*Alytes obstetricans*), lending the virus its name. Since 2007 there have been repeated outbreaks of ranavirosis attributed to CMTV-like ranaviruses throughout continental Europe, affecting a wide range of species and life stages. Mass mortality events have been observed in Danish edible frogs (*Pelophylax kl. Esculentus*; Ariel et al. 2009) and a French population of *R. temporaria* (Miaud et al. 2016). Worryingly, CMTV has also been documented to cause local outbreaks of ranavirosis that can span entire amphibian community assemblages and all life stages, causing pronounced and persistent population declines (Price et al. 2014; Rijks et al. 2016; Rosa et al. 2017).

Several strains of CMTV are known to be circulating in Europe, and experimental work has revealed these strains to vary markedly in pathogenicity (Saucedo et al. 2018, 2019). A lack of genetic diversity in CMTV isolates collected from various outbreaks and a high number of genetic loci that appear to be under strong selection has led to the hypothesis that CMTV is an invasive pathogen on continental Europe (Price 2015; Price et al. 2017b), though the route of invasion remains unknown. Although FV3-like ranaviruses are known to occur in Europe in both amphibians and reptiles (Price et al. 2014; Stöhr et al. 2015) and are known to occur in sympatry with CMTV-like viruses, such incidences are rare (Price et al. 2017b). Recent studies have shown that in regions where both viral species groups co-occur there is potential for high levels of recombination resulting in hybrid viruses, which could potentially possess elevated virulence compared to either CMTV- or FV3-like viruses (Price et al. 2014; Price 2015; Claytor et al. 2017; Rosa et al. 2017). Overlap also exists between CMTV and the amphibian chytrid fungi, and co-infections are known to establish (Rosa et al. 2017); however, little evidence currently exists that either pathogen exacerbates the impact of the other (Rosa et al. 2017). The gross pathology of ranavirosis caused by CMTV is identical to that which is caused by FV3 (e.g., Cunningham et al. 1997; Balseiro et al. 2009); however, CMTV is known to replicate initially in the oral cavity of infected amphibians, before invading the connective tissues and subsequently the organs, including the skin (Saucedo et al. 2019). Research that focused on an alpine population

of *R. temporaria* in France has also been key in proving the efficacy of environmental DNA or “e-DNA” at detecting the presence of ranaviruses in European amphibian populations (Miaud et al. 2019). Critically, the use of e-DNA not only allows for the detection of pathogens without the need for intensive or invasive sampling techniques of hosts, but also permits quantification of temporal change in pathogen loads in the environment (e.g., Hall et al. 2016; Miaud et al. 2019). Such data could prove invaluable for attempting to understand spatial variation in the severity of ranavirosis outbreaks.

Future directions

Although European research has been instrumental in advancing the study of amphibian infecting ranaviruses around the world, there remain significant gaps in our understanding that need to be addressed if effective management and mitigation strategies of these emerging pathogens are to be developed. Below, we outline these key directions of future research and how they may be pursued. Although we give special consideration to how these knowledge gaps may be addressed in European amphibian systems, these directions of research should be considered a priority of researchers globally.

Further investigation of the interaction between the amphibian microbiome and ranaviruses

There is substantial and growing evidence that the amphibian skin microbiome is vital to the ability of amphibians to resist or tolerate infection by pathogens (Harris et al. 2006, 2009; Becker et al. 2015). Although this evidence is primarily drawn from studies concerning amphibians infected with the chytrid fungi, including *Batrachochytrium dendrobatidis* (Bd) (Kueneman et al. 2016; Harrison et al. 2020) and *B. salamandrivorans* (Bates et al. 2019), recent research suggests that host-associated microbial communities in amphibians also represent a key component of their immune defense against ranaviruses. For example, skin microbiome structure has been found to correlate with population disease-status in the wild (Campbell et al. 2018b, 2019) and survival of acute ranaviral infection under laboratory conditions (Harrison et al. 2019). Furthermore, gut microbiomes of developing wood frogs (*Rana sylvatica*) have been shown to be critical modulators of later life resistance to ranavirus (Warne et al. 2019). These are potentially important findings as the ability of probiotic treatments to mitigate chytridiomycosis has been experimentally demonstrated (e.g., Kueneman et al. 2016) and similar treatments for ranavirosis would significantly advance amphibian conservation efforts. However, despite the clear importance of the host-associated microbial communities to emerging pathogens such as ranaviruses, several significant gaps in our knowledge remain that must be addressed before such probiotic treatments could be developed.

Generality of microbiome–pathogen interactions

To date, all evidence of a link between ranavirosis and the amphibian skin microbiome originates from within the United Kingdom *R. temporaria* field system outlined above, and studies on the interaction between gut microbiomes, host immunity, and ranavirus remain rare (e.g., Warne et al. 2019). Understanding the generality of these patterns in other host species and locations is a vital first step in quantifying the importance of amphibian microbial communities in shaping resistance or tolerance to ranaviruses. As such, further field and experimental studies of the amphibian microbiome in populations with varying ranaviral disease history, particularly in a diversity of vulnerable species outside of the United Kingdom, are a crucial first area of future research.

How should we measure host–microbe interactions?

Understanding the ecology of host-associated microbes, and how they influence host health is challenging in wild systems. Researchers face the choice of a variety of molecular methods for quantifying

the composition of microbial communities, including those that simply yield information on the presence and relative abundance of microbial taxonomic groups (e.g., 16S rRNA amplicon sequencing), and those that also directly estimate functional properties of the microbiome (e.g., metagenomics, metatranscriptomics, and metabolomics ([Harrison and Cameron 2020](#))). An outstanding issue is that the precision with which we can measure host–microbe interactions, and thus the inferences we make, depend on which of these methods we use. For example, [Campbell et al. \(2018b\)](#) profiled the *R. temporaria* microbiome using bacterial reads filtered from a transcriptomics study of the response of wild frog populations to endemic ranavirosis. They found that *Bacillus subtilis*, which is a bacterium commonly used as a probiotic in the aquaculture industry (e.g., [Aly et al. 2008](#); [Liu et al. 2012](#); [Ran et al. 2012](#)), was two orders of magnitude more abundant on frogs that were persisting at sites with endemic disease versus frogs from ranavirosis-free populations. However, the same pattern was not detected during a follow-up study that used 16S rRNA amplicon sequencing to categorise and compare the skin bacterial community structure of the same populations from samples collected the following year ([Campbell et al. 2019](#)). This discrepancy between results based on relative bacterial abundance versus bacterial transcription raises two interesting questions that warrant further study. Firstly, does transcriptional activity or relative biomass present the most informative indication of the importance of a microbial species in the amphibian skin microbiome? Understanding which potential measure is a truer reflection of genuine interaction between bacterial taxa and a pathogen will allow for more accurate detection of bacterial species which possess the potential to serve as probiotics in mitigation strategies. Targeted transcriptomic studies of the amphibian commensal skin bacteria (often termed the metatranscriptome) are rare ([Rebollar et al. 2016](#)). However, the incorporation of transcriptomic analysis in future research of the link between that amphibian skin microbiome and ranavirosis will be important in addressing this question. Moreover, “omics” technologies such as transcriptomics can also resolve the taxonomy of microbes to species and even strain level, yielding more precise information on which bacterial groups are responsible for observed differences in host response to disease. An experimental infection trial incorporating appraisal of shifts in the bacterial community structure using 16S rRNA amplicon techniques (relative abundance) versus changes to bacterial transcription in the same individuals following exposure to a ranavirus would be useful in determining whether bacterial biomass or transcription are more informative in predicting the outcome of infection by a pathogen. A targeted comparison of the metatranscriptomes of wild amphibian populations persisting at sites with endemic ranavirosis versus those from disease-free populations would also help to determine which species of bacteria are transcriptionally linked to ranavirosis under natural conditions.

Microbiome structure vs. stability

The differences between the two studies discussed above could also represent a temporal difference in the skin microbiomes of the animals at the populations sampled. Though a vast number of studies focus on drivers of microbiome structure in the wild, very few focus on microbiome stability and its consequences for the host. Recent experimental work has revealed that the stability of the amphibian skin microbiome is governed by the complexity of the environmental microbial reservoir ([Harrison et al. 2019](#)). This gives us strong reason to expect that there is marked variation in the stability of the skin microbiome in wild frogs, which may even vary by population in concert with local environmental conditions. Capture–mark–recapture techniques such as visible implant elastomer tagging or radio-frequency identification tags could be used to examine the temporal stability of the amphibian skin microbiome in the wild by identifying and repeat sampling individual wild amphibians in multi-year frameworks. This would allow for the quantification of the composition, stability, and functional activity of the amphibian skin microbiome and for these traits to be linked to ranaviral disease dynamics throughout longitudinal studies.

Understanding the mechanism underlying microbiome–disease relationships

Whilst present evidence represents an intriguing first demonstration of the potentially important role of the commensal microbiome for determining host response to ranavirosis, the mechanisms and directionality of this relationship are not understood (Campbell et al. 2019; Harrison et al. 2019). Particularly, it is unknown whether the composition of bacteria on amphibian skin is the result of originating from a population with endemic ranavirosis or whether certain bacterial community compositions can predispose a population to suffer outbreaks of ranavirosis. Both ranavirosis (Harrison et al. 2019) and chytridiomycosis (Jani and Briggs 2014) have been shown to perturb the amphibian skin microbiome, which could be expected to generate correlations between microbiome structure and disease status. Understanding the direction of this relationship is critical to appreciate the true potential of using probiotic treatments or bioaugmentation strategies to combat ranaviruses in wild populations. Development of techniques that enable the manipulation of bacterial communities provide the potential to perform experimental infection trials using amphibians with altered cutaneous microbiomes to examine how the presence or absence of key microbial species modulates response to pathogen exposure and subsequent survival. This type of experiment will prove invaluable in disentangling cause and effect in apparent microbiome–disease relationships. This is particularly relevant to when we think about how an individual bacterial species might be interacting with the pathogen. For example, many bacterial species have been shown to demonstrate inhibition of growth of *B. dendrobatidis* (Antwis and Harrison 2018; Harrison et al. 2020), suggesting a direct interaction. However, microbes can also effect changes to a host's response to a pathogen indirectly by stimulating host gene expression (Rebollar et al. 2018). To date, no study has discerned whether the microbiome's role in governing variance in host resistance to ranavirus is driven by direct interaction with the pathogen, or indirect effects on the host. Lack of knowledge regarding this mechanistic pathway remains a clear gap in our understanding that should be addressed as a priority.

Quantifying extrinsic drivers of ranavirosis outbreaks

It is widely appreciated that host–pathogen interactions in many systems are complex and often modulated by external factors, including the environment in which those interactions occur (e.g., James et al. 2015). Recent research conducted within the longitudinally studied *R. temporaria* populations in the United Kingdom clearly demonstrates that the same is true in the case of ranaviruses and amphibians. The severity of ranavirosis outbreaks within a population has been linked to the use of horticultural chemicals, particularly molluscicides, within or near the water body in which that population resides (North et al. 2015). Likewise, ambient temperature can alter ranavirosis dynamics and infection trajectories (Price et al. 2019), which will have marked implications for variation in disease risk across the geographical distribution of hosts. Here we briefly discuss key knowledge gaps regarding how these abiotic stressors, temperature, and pollutants, influence amphibian disease dynamics, and how they should be addressed.

Host–pollutant–pathogen interactions

The mechanisms by which pollutants may modulate host response to disease are not fully resolved but can broadly be separated into direct and indirect components (Fig. 2). For example, environmental pollutants can act as direct physiological stressors of the amphibian immune system (Robert et al. 2018; Thambirajah et al. 2019), heightening their susceptibility to disease (Fig. 2A). Interestingly, this relationship appears to be reciprocal and the presence of a ranavirus has also been shown to exacerbate the lethality of pesticides (Pochini and Hoverman 2017). Previous studies have shown that herbicides and pesticides can reduce resistance of amphibians to fungal pathogens (e.g., Krynak et al. 2017), perhaps by causing a reduction in anti-microbial peptide (AMP) production (Schadich 2009) and subsequently reduced skin peptide defense against Bd (Davidson et al. 2007; Fig. 2). Both

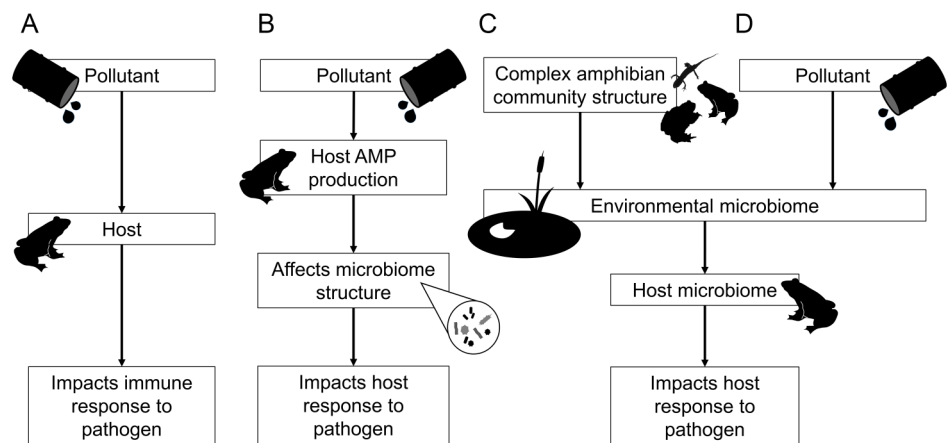


Fig. 2. Schematic diagram of the pathways via which pollutants may change host response to disease. (A) Pollutants may exert direct effects on host immunity. (B) Pollutants may effect changes in the microbiome indirectly by first driving differences in host AMP production that then select for differential microbial proliferation. Altered microbiomes then compromise host response to the pathogen. Finally, both (C) the wider amphibian community and (D) pollutants may alter the environmental reservoir of microbes capable of colonising a focal amphibian host, thus indirectly altering microbiome-mediated immunity to pathogens.

pathways involve a direct effect of the pollutant on the host immune response, which determines subsequent response to a pathogen (Fig. 2A).

However, despite the important role of the microbiome in governing host response to disease, the potential for pollutants to effect changes in amphibian pathogen resistance indirectly via their influence on the microbiome has received relatively little attention (but see McCoy and Peralta 2018). Both heavy metals from agricultural and industrial run-off, and antimicrobials from pharmaceutical pollutants (Casado et al. 2019), can reduce amphibian gut microbiome diversity (Zhang et al. 2016) and skin microbiome composition (Hernández-Gómez et al. 2020). Pollutant-microbiome relationships may arise either because of a direct effect of the pollutant on the microbiome, or indirectly because pollutants first alter host skin AMP secretion that in turn perturb skin microbial communities (Fig. 2B). Both scenarios represent important pathways for future investigation. Finally, there is strong evidence that the amphibian skin microbiome is populated from environmentally available bacterial species (Kueneman et al. 2014; Longo et al. 2015; Campbell et al. 2019; Harrison et al. 2019). Thus, anything that alters the pool of environmental microbes capable of colonizing the host may also drive differences in subsequent host microbiome structure. Local amphibian community complexity may similarly alter the dynamics and persistence of environmental microbes, for example by introducing novel microbes through immigration (Fig. 2C). Likewise, environmental pollutants can change the diversity of microbial assemblages within both soil and aquatic reservoirs (e.g., Bissett et al. 2013; Muturi et al. 2017; McCoy and Peralta 2018), and indirectly modulate the composition of the amphibian skin microbiome via these environmental effects (Fig. 2D).

Despite growing interest in amphibian microbiome research, studies of host-microbiome-environment interactions remain rare particularly in wild systems (e.g., Varga et al. 2019). Future work should seek to quantify the mechanism by which pollutants alter host susceptibility to disease by simultaneously studying host gene expression and skin microbiome structure in the presence of such contaminants. Surveying at sites with and without historical ranavirus prevalence has already uncovered systematic differences in the structure of the amphibian skin microbiome and the

expression of host genes related to the secretion of anti-microbial peptides and inflammation based on disease status (Campbell et al. 2018b, 2019). As amphibians source their skin microbiota from environmental reservoirs, we could expect different geographic locations to influence present amphibian population's microbiome structure (Campbell et al. 2019; Ross et al. 2019). Therefore, a priority for future studies would be to survey amphibian populations across a wide spatial scale encompassing multiple habitat types, but also varying in ecological gradients of both ranaviral pathogen loads and environmental pollutants. Focal pollutants could include heavy metals near (historically) industrial areas (e.g., Costa et al. 2016), pest-control agents such as molluscicides (North et al. 2015), and fertilizer chemicals (e.g., nitrates) known to affect amphibian physiology and susceptibility to pathogens (e.g., McCoy and Peralta 2018). Developing our understanding of how the amphibian skin microbiome, environmental contaminants, and ranavirosis are inter-linked is important for amphibian conservation efforts and for potential mitigation strategies, such as probiotic treatments; to work in wild systems we must have as full an appreciation as possible about how they will influence and be influenced by the complex interactions between host, pathogen, and environment.

Continued surveillance in the face of environmental change

Ranavirosis outbreaks due to FV3 appear to happen seasonally in the United Kingdom and most incidences of ranavirosis occur when ambient temperatures are 16 °C or higher (Cunningham et al. 1997; Teacher et al. 2010; Price et al. 2019). It has also been experimentally shown that FV3 grows faster at higher temperatures, resulting in a positive correlation between temperature and infection severity in *R. temporaria* (Price et al. 2019). If current trends in climate warming continue, then the annual time frame during which average monthly temperatures exceed 16 °C will increase and the geographical area over which these conditions are met will spread throughout the United Kingdom (Price et al. 2019). This spread will result in a higher number of populations experiencing ranavirosis outbreaks. It is also likely that climate change will drive distribution shifts of many amphibian species in the United Kingdom (Dunford and Berry 2013), potentially forcing yet more amphibian populations into regions where ranavirosis outbreaks are increasingly likely.

Not only has it been shown that a warming climate will increase the incidence of ranavirosis in the United Kingdom but also that associated climactic instability may be associated with reduced population viability of impacted populations, acting in tandem with age truncation of *R. temporaria* populations caused by endemic ranavirosis (Campbell et al. 2018a). Age-truncated *R. temporaria* populations exhibit an over-abundance of younger, smaller breeding individuals (Campbell et al. 2018a). As fecundity of *R. temporaria* is tightly correlated with body size, per capita recruitment rate of entire populations existing with endemic ranavirosis is lower than that of populations that are disease-free (Campbell et al. 2018a). Population matrix models suggest that this lower recruitment rate potentially heightens the impact of stochastic events that further reduce recruitment such as late frosts, or drought resulting in an increased likelihood of population collapse (Campbell et al. 2018a). As climactic instability also increases due to climate change, the impact of ranavirosis outbreaks on the population level may also be more severe across the ever-expanding range. However, additional population models of *R. temporaria* have shown that population dynamics are primarily influenced by the survival of adults and not larval or metamorphic individuals (Miaud et al. 1999; Biek et al. 2002). This means the potential for ranavirosis induced population instability may well be unique to the United Kingdom, where ranavirosis impacts adult *R. temporaria* rather than larval amphibians, which is the case elsewhere in Europe and around the world. Nevertheless, in the United Kingdom, it is important to continue to monitor *R. temporaria* populations and track the spread of ranavirosis throughout the country as closely as possible.

Little is yet known about the influence of climate on the virulence and spreading potential of CMTV. Understanding the influence of temperature on CMTV through experimental growth assays and in vivo challenge experiments will allow for predictions of how climate change will impact the spread of CMTV throughout continental Europe and the expansion of annual periods of disease outbreaks. The use of e-DNA monitoring tools to quantify environmental loads of ranavirus and subsequent infection risk will be crucial for refining predictive models of outbreak incidence and intensity across environmental gradients, such as intra- and interannual variation in temperature (e.g., [Hall et al. 2016](#); [Miaud et al. 2019](#)).

Studies looking into the indirect impacts of CMTV outbreaks on European amphibians are also rare. Field-based studies aimed at understanding how outbreaks of ranavirosis due to CMTV impact the structure of continental European amphibian populations through mechanisms such as age truncation or impinged recruitment will also allow for more accurate predictions of the impact of CMTV on amphibian populations across the continent. Additionally, understanding which regions of continental Europe stand to be more heavily impacted by the spread of ranavirosis requires that we understand which regions harbor the highest number of susceptible species. In vivo challenge experiments could help provide this information. Coupled with an effective surveillance project, likely partially implemented using e-DNA techniques, conducted at the leading edge of disease emergence, precious conservation resources can be targeted towards those regions which are most in peril or those regions where intervention stands the greatest chance of success.

Fill-in ecological blanks for better epidemiological models

Recent events have demonstrated the critical importance of well-parameterised epidemiological models in predicting the spread of emerging diseases and designing and implementing successful mitigation strategies (or not). A small number of studies have attempted to apply various types of epidemiological models to the emergence of ranaviruses in Europe ([Campbell et al. 2018a](#); [Duffus et al. 2019](#)), and have provided useful insight into several aspects of host-ranavirus interactions in a European context. Despite this, these studies have also demonstrated that a dearth of knowledge regarding the basic ecology of European amphibians and the ranaviruses that infect them can result in tenuous parameterization of these models from the literature, often drawn from studies focused on distantly related host species or systems with which there is minimal apparent ecological overlap ([Campbell et al. 2018a](#); [Duffus et al. 2019](#)). Refining our understanding of host-ranavirus interactions requires that we quantify and incorporate fundamental ecological variation into our models. Within populations, individuals will vary markedly in key life history traits such as growth rate, reproductive output, migration propensity, and (immune) genotype. Likewise, the ranaviral genomic variants hosts are exposed to are expected to vary in replication rates and subsequent transmission potential. Collectively these traits will interact to influence parameters such as the likelihood, frequency, and intensity of ranavirus infection within, and transmission among, amphibian populations. Tackling these knowledge gaps will undoubtedly require long term data sets of marked individuals paired with frequent monitoring, quantification, and genotyping of ranaviral genomic variants to understand the factors governing long term probability of persistence of both pathogen and hosts in natural populations. Using these long-term data sets to quantify rates of disease transmission within and among populations will undoubtedly lead to strong epidemiological models, increasing the accuracy with which researchers can make predictions about the spread of these deadly emerging pathogens.

Conclusion

Ranaviruses are an emerging disease threat to European amphibian populations. Both directly, through elevated mortality, and indirectly, through the effects of endemic disease on host populations,

ranaviruses have been found to cause catastrophic and sustained declines in amphibians in the United Kingdom and on continental Europe. As climate change drives global temperature increases and climatic instability, the pace of ranaviral emergence and the severity of ranavirosis outbreaks in Europe are likely to follow suit. This will result in the exposure of more populations and additional species over a wider annual time period and a reduction in the long-term viability of impacted populations. In light of this, it is imperative that we continue to monitor the spread of ranaviruses throughout Europe and to assess which currently naïve European amphibian species are likely to be most vulnerable to infection, allowing for the targeting of conservation efforts in regions where the impact of ranaviral emergence is likely to be particularly severe.

Happily, recent research suggests that the amphibian skin microbiome may offer a source of potential mitigation strategies in the form of probiotic treatment and environmental bioaugmentation, as it has done in response to the chytrid fungi (Kueneman et al. 2016; Antwis and Harrison 2018; Harrison et al. 2020). For the potential efficacy of such strategies to be appraised, a number of fundamental research questions need to first be addressed. Primarily, we must aim to assess the generality of relationships between the amphibian skin microbiome and ranavirosis in a wide array of systems. Additional questions regarding the measure by which we judge the importance of a microbial species in a microbiome and the subsequent impact of a complex environment on host/microbiome–pathogen interactions will need to be addressed. European research has been critical in generating knowledge regarding ranaviruses and their impact on amphibians. The proximity within which many European amphibian populations live to humans, coupled with the proven potential for significant public engagement in the study of ranavirosis in Europe, presents unique opportunities to address these important research questions that could lead to advances in amphibian conservation not just in Europe but around the world.

Author contributions

LJC and XAH conceived and designed the study. LJC, AHP, and XAH drafted or revised the manuscript.

Competing interests

The authors have declared that no competing interests exist.

Data availability statement

All relevant data are within the paper.

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