Antimicrobial peptide and sequence variation along a latitudinal gradient in two anurans

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Abstract

Background: While there is evidence of both purifying and balancing selection in immune defense genes, large-scale genetic diversity in antimicrobial peptides (AMPs), an important part of the innate immune system released from dermal glands in the skin, has remained uninvestigated. Here we describe genetic diversity at three AMP loci (Temporin, Brevinin and Palustrin) in two ranid frogs (Rana arvalis and R. temporaria) along a 2000 km latitudinal gradient. We amplified and sequenced part of the Acidic Propiece domain and the hypervariable Mature Peptide domain (~150-200 bp) in the three genes using Illumina Miseq and expected to find decreased AMP genetic variation towards the northern distribution limit of the species similarly to studies on MHC genetic patterns.

Results: We found multiple loci for each AMP and relatively high gene diversity, but no clear pattern of geographic genetic structure along the latitudinal gradient. We found evidence of trans-specific polymorphism in the two species, indicating a common evolutionary origin of the alleles. Temporin and Brevinin did not form monophyletic clades suggesting that they belong to the same gene family. By implementing codon evolution models we found evidence of strong positive selection acting on the Mature Peptide. We also found evidence of diversifying selection as indicated by divergent allele frequencies among populations and high Theta k values.

Conclusion: Our results suggest that AMPs are an important source of adaptive diversity, minimizing the chance of microorganisms developing resistance to individual peptides.

Keywords: Rana, MHC, AMPs, Trans-specific polymorphism, Multi-locus system

Background

Vertebrates fight pathogens using both the adaptive and the innate immune systems. The adaptive immune system is composed of highly specialized tissues and systemic cells, which synthetize antibodies and recognize an infinite diversity of antigens [1]. The adaptive immune system usually clears the infection and protects the host against reinfection with the same pathogen [1, 2]. The innate immune system provides a non-specific response to pathogens, and when a pathogen is detected it acts immediately with an early induced inflammatory response recognized by non-specific effectors [2]. The vertebrate innate immune system includes macrophages and neutrophils [3], natural killer cells [4] and antimicrobial peptides (AMPs [5, 6]), the latter often being secreted from the granular glands on the dermal layer of the skin. AMPs are generally short (15–45 amino acid residues) [7], cationic [8], amphipathic [9] and α helical [10]. These molecules are found in a great variety of taxa such as mollusks [11], fish [12], amphibians [13], birds [14] and mammals, including humans [15].

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In amphibians, AMPs consist of a highly conserved Signal Peptide, an Acidic Propiece domain and a hypervariable C-terminus domain [16], Fig. 1). Amphibian AMPs exhibit potent activity against antibiotic resistant bacteria, protozoa, yeast, virus and fungi by killing cells via disrupting the membrane integrity [17, 18]. They have been implicated as a defense against the lethal fungus *Batrachochytrium dendrobatidis* (Bd) causing global amphibian population declines (e.g., [19–21]). It has been firmly established that AMPs may be encoded by duplicated genes [17, 22]. A high number of amphibian AMPs has been characterized with the aim to develop peptide-based therapeutic agents [23, 24]. However, most of the AMP studies have focused on the antimicrobial peptide variation between amphibian species, and the genetic variation at the population level has received less consideration. Indeed, while there is evidence of geographic variation in effectiveness of antimicrobial defenses in various organism groups [22, 25–29], population-level variation in peptide profiles [27, 29] and positive selection acting on the Mature Peptide domain [6, 7, 16], little attention has been directed towards understanding the evolutionary forces acting on the genes coding for AMPs [17, 22, 30, 31]. This is especially the case for the role of selection acting upon sequence variation in AMP genes, and how this variation is distributed within populations as well as at larger geographical scales.

Intraspecific genetic diversity is a fundamental dimension of biodiversity. Intraspecific genetic diversity is a reflection of both past and current evolutionary processes, as well as an indicator of a population’s potential for adaptation to future stressors [32]. In temperate and boreal regions, historical colonizations events such as postglacial colonizations have a strong effect on genetic variation at all spatial scales. Genetic diversity both within and between species generally decreases with increasing latitude [33]. Moreover, understanding the role of selection and gene flow is pivotal for understanding the genetic diversity displayed in contemporary populations. For instance, diversifying selection in different parts of the gradient, especially in cases in which a cline coincides with a demographic shift, may cause population divergences [34]. Thus, genetic diversity patterns could be attributed to diversifying selection, different demography and/or gene flow along a geographical gradient [35, 36]. To which extent demography, selection, gene flow and their interactions contribute to shape large-scale genetic variation remains a complex and unresolved issue.

In this study, we assessed variation in AMP genes in two ranid frogs, the moor frog *Rana arvalis* and the common frog *R. temporaria*, along a latitudinal gradient from northern Germany to the northern margin of the distribution of the two species. We investigated both the historical postglacial colonization patterns along the latitudinal gradient across northern Europe (Additional File 3: Figure 1) and current evolutionary bottlenecks (selective forces, drift/demography, migration) driving AMP intraspecific genetic variation. *R. temporaria* shows an almost unidirectional postglacial colonization of Scandinavia, colonizing most of Sweden from the north via Finland and meeting another lineage colonizing from the south via Denmark in southernmost Sweden [37, 38]. *R. arvalis* shows a different pattern of dual postglacial colonization from south and north to Scandinavia, the contact zone being situated in central northern Sweden [39, 40]. Populations at the northern edge are smaller, more fragmented and isolated relative to populations at the core of the distribution range [41–43]. As a consequence of decreasing effective population size at range margins, many species show lower genetic variation at the edge of their distribution range in concordance with the central-marginal hypothesis [44]. *R. arvalis* shows a
clear and structured allele distribution with a decreasing pattern of genetic variation towards northern latitudes at microsatellite loci as well as at the major histocompatibility complex (MHC) class II exon 2, an important component of the adaptive immune response [40]. These results are in concordance with the central-marginal hypothesis. Following previous results, we expected the lowest genetic diversity at the AMP loci at the northern edge of the distribution of both species.

To our knowledge, this is the first investigation of large-scale genetic variation in AMPs 1) within and between populations of a species and 2) comparing two different species. We compared AMP genetic variation among 14 R. arvalis and 17 R. temporaria populations from northern Germany to northern Scandinavia. We expected a decrease in AMP genetic variation towards the northern distribution limit of the species. We characterized skin antimicrobial peptides genetically using ultra-deep Illumina sequencing in order to elucidate the underlying evolutionary processes acting on AMP genes. Specifically, we asked: 1) are patterns of AMP genetic variation influenced by demographic history? 2) is there evidence of selection at AMP loci? and 3) to which extent is selection affecting AMP gene characteristics?

Results

AMP diversity: Miseq run summary
The total number of reads obtained by run varied from 2.597.554 to 5.124.580. We obtained an average of 4.048.930 reads ±987.541 (s.d.) with intact primer barcode information from four separate runs (Additional File 1: Table 1). The average number of reads per amplicon was 218.503.14 ± 35.720 (s.d.). Amplicons with < 300 reads were discarded. We included replicated individuals for each of the four Miseq runs (15–20%); (Additional File 1: Table 1). Replicates were randomly assigned across different pools to avoid false allele identification. All replicates produced the same alleles in each sample. We extracted DNA from a total of 320 individuals (150 R. arvalis and 170 R. temporaria). For all three genes (Temporin, Brevinin and Palustrin), we amplified and sequenced a total of 849 amphibian DNA samples, plus the duplicates (Raar = 376; Rate = 473; See Table 1; (Additional File 1: Table 1). For Temporin, we genotyped (275) (Raar = 109; Rate = 166), for Brevinin 296 (Raar = 131; Rate = 165) and for Palustrin 278 (Raar = 136; Rate = 142) individuals. Among the 849 sequenced samples, we assigned 58 valid AMP sequences with length variation within and among loci (from 111 to 204 bp) by using the DOC method [45].

Multi-locus family genes
We detected more than two alleles per individual in both species, indicating multiple copies in all sequenced AMP genes. Number of valid AMP alleles per individual varied between one and 11 in Temporin, one and seven in Brevinin, and one and six in Palustrin. The average number of gene copies per AMP ranged from 2 to 4 (Temporin: Raar = 4.17 ± 1.10 [s.d.], Rate = 3.68 ± 2.17 [s.d.]; Brevinin: Raar = 3.09 ± 1.38 [s.d.], Rate = 2.00 ± 1.09 [s.d.]; and Palustrin: Rate = 3.59 ± 1.13 [s.d.]).

AMP nucleotide gene diversity
Allele frequency pie-charts showed a non-structured genetic pattern from northern to southern populations at all loci for both species (Additional File 4: Figure 2). Pairwise nucleotide differences (Theta k) varied considerably among species and group of genes. However, Theta k values were very similar within regions, showing the same allelic pattern in the allele frequency plots throughout the gradient (Table 1; (Additional File 4: Figure 2; Additional File 5: Figure 3). In Temporin, three out of 21 alleles (Rate_Temp*21, Rate_Temp*20, Rate_Temp*17) were widespread at high frequencies in all R. temporaria populations (Additional File 4: Figure 2; Additional File 5: Figure 3). Similarly, the three most common alleles (Raar_Rate_Temp*02, Raar_Rate_Temp*05 and Raar_Rate_Temp*17) out of the 13 in R. arvalis were found in all populations at high frequency (Additional File 4: Figure 2; Additional File 5: Figure 3). In Palustrin, Rate_Palu*07 and Raar_Rate_Palu*04 were the most frequent alleles out of the 12 alleles in R. temporaria (Additional File 4: Figure 2; Additional File 5: Figure 3) and found in all populations. Palustrin was monomorphic in R. arvalis. In Brevinin the most predominant allele Raar_Rate_Brev*02 was found in all populations along the gradient in both species (Additional File 4: Figure 2; Additional File 5: Figure 3). In R. arvalis, four (Raar_Brev*10, 11, 12, 16) Brevinin alleles out of the total of 14 were present in only southern populations from Uppsala to Germany.

The number of alleles per region was significantly different among regions for the Brevinin group of genes in R. arvalis (F_{4,9.62} = 5.48, P = 0.014), but there was no difference in R. temporaria (F_{5,25.78} = 1.82, P = 0.14). The results were the opposite for the Temporin group of genes, where we found a higher number of alleles in the two southern regions in R. temporaria (F_{5,10.85} = 4.29, P = 0.021), but not in R. arvalis (F_{4,8.53} = 0.61, P = 0.41). For the Palustrin group of genes we found no significant differences in number of alleles among the regions in R. temporaria (F_{5,10.78} = 2.54, P = 0.092). The number of Brevinin alleles was significantly higher in R. arvalis than in R. temporaria (F_{1, 25.34} = 22.16, P < 0.001; Fig. 2), and almost so in Temporin (F_{1,25.53} = 5.36, P = 0.054).

Phylogenetic analyses
The Neighbor-joining (NJ) phylogenetic tree and the haplotype network generated for Temporin showed that R. arvalis and R. temporaria alleles were distinctly
| Loci | Population | Region | \( N^o \) of Ind Genotyped | Species | \( k \) | \( \hat{h}_{alleles} \) | \( \pi \) | \( \hat{r}_{alleles/Region} \) | \( N^o \) of alleles/Region | Average \( N^o \) alleles/regional | Total \( N^o \) alleles/ind | Total \( N^o \) Peptide | Average \( N^o \) Peptides/ind |
|------|------------|--------|---------------------------|---------|------|----------------|------|-----------------|---------------------|--------------------------|----------------|----------------------|----------------------|
| M    | L Germany  |         | 26.45                     | R. arvalis | 1.18 | 24.36         | 0.19 | 20.32           | 0.15                | 3.21±0.49               | 13              | 4.17±1.00           | 6                    | 2.69±0.28            |
| A    | L Germany  |         | 20.89                     | R. arvalis | 1.17 | 15.36         | 0.13 | 12.04           | 0.10                | 3.08±0.35               | 13              | 4.17±1.00           | 6                    | 2.69±0.28            |
| T    | S Skitee   |         | 20.10                     | R. arvalis | 1.13 | 15.36         | 0.13 | 12.04           | 0.10                | 3.08±0.35               | 13              | 4.17±1.00           | 6                    | 2.69±0.28            |
| AO   | O Uppsala  | V       | 38.27                     | R. arvalis | 2.66 | 35.96         | 0.39 | 28.60           | 0.26                | 3.81±1.29               | 13              | 4.17±1.00           | 6                    | 2.69±0.28            |
| NY   | H Umeå     |         | 26.61                     | R. arvalis | 1.18 | 24.36         | 0.19 | 20.32           | 0.15                | 3.21±0.49               | 13              | 4.17±1.00           | 6                    | 2.69±0.28            |
| B    | E Luleå    |         | 20.04                     | R. arvalis | 1.17 | 15.36         | 0.13 | 12.04           | 0.10                | 3.08±0.35               | 13              | 4.17±1.00           | 6                    | 2.69±0.28            |

**Table 1** Diversity statistics summary for Population/Region of Temporin, Brevinin and Palustrin group of genes in two different frog species (R. arvalis and R. temporaria). K: Theta K pairwise nucleotide comparison. K_Region: Theta K pairwise comparison within regions. \( \pi \): nucleotide diversity and \( \hat{r}_{alleles/Region} \): nucleotide diversity within regions. Number of alleles in brackets represent unique alleles within regions.
separated with the exception of two alleles which were present in both species (Raar_Rate_Temporin*02,05; Fig. 3; (Additional File 6: Figure 4). In contrast, the trees generated for Brevinin and Palustrin indicated that many alleles (species-specific and shared) were not separated between the two species. The alleles of both species were clustered into three, five and two clades for Temporin, Brevinin and Palustrin, respectively (Additional File 7: Figure 5).

We constructed a NJ phylogenetic tree where all alleles earlier assigned to Temporin, Brevinin and Palustrin were clustered in seven different groups/clades according to the connected branches and nodes of the tree (Fig. 3). Surprisingly, Temporin and Brevinin alleles were widespread and did not form monophyletic clades in the phylogenetic trees, occurring in the same groups/clades. Consequently, we consider Temporin and Brevinin as a part of the same gene family and they were grouped together for consecutive analyses.

AMP peptide diversity

When DNA sequences were translated to amino-acids, variant frequency showed a structured pattern from northern to southern populations in Temporin for R. temporaria and in Brevinin for R. arvalis (Temporin; $F_{A,10.85} = 4.29$, $p = 0.021$, Brevinin; $F_{A,8.51} = 3.59$, $p = 0.054$; Fig. 4; (Additional File 8: Figure 6 and Additional File 9: Figure 7), indicating lower peptide variation at higher latitudes. The Palustrin group did not show any pattern of peptide variation along the gradient in R. temporaria (Fig. 4; (Additional file 10: Figure S8, Additional file 11: Figure S9, Additional file 12: Figure 10, Additional file 13: Figure S11 and Additional file 14: Figure S12). Palu_Amino*05 was the only amino-acid variant found in R. arvalis for Palustrin (Fig. 4; From (Additional file 10: Figure S8, Additional file 11: Figure S9, Additional file 12: Figure 10, Additional file 13: Figure S11 and Additional file 14: Figure S12). The Temporin variant Temp_Amino*07, and two amino-acid variants Temp_Brev_Amino*01,02 present at both Brevinin and Temporin, were the only functional amino-acid variants shared between the two species (Fig. 4; (Additional File 8: Figure 6 and Additional File 9: Figure 7).

Adaptive evolution/positive selection in AMPs multi-locus gene family

We examined patterns of nucleotide diversity ($\pi$) and average pairwise nucleotide differences (Theta $\kappa$) within each of the 46 Mature Peptide and Acidic Propiece domains among four set of clades (Temporin-Brevinin) and 12 Mature Peptide and Acidic Propiece domains among two set of clades (Palustrin) (Table 2). The diversity at the Mature Peptide domain was higher ($\pi_{\text{Mature Peptide}} = 0.437$; $\pi_{\text{Mature Peptide}} = 0.35$, $k_{\text{Mature Peptide}} = 14.439$; $k_{\text{Mature Peptide}} = 23.136$) and more divergent than at the Acidic Propiece domains ($\pi_{\text{Acidic Propiece}} = 0.115$; $\pi_{\text{Acidic Propiece}} = 0.116$; $k_{\text{Acidic Propiece}} = 11.76$; $k_{\text{Acidic Propiece}} = 2.78$) in both Temporin-Brevinin and...
Fig. 3 (See legend on next page.)
Palustrin group of genes, respectively. The net charges of the Mature Peptide and the Acidic Propiece showed a negative relationship among Temporin-Brevinin and Palustrin sequences. The corresponding “Im” slopes were $-2.75 \pm 0.45$ for Temporin-Brevinin and $-1.02 \pm 0.41$ for Palustrin group of genes, indicating coordinated changes between the Mature Peptide domain and the Acidic Propiece domain. Net charges of the Mature Peptide and the Acidic Propiece domains for Temporin-Brevinin and Palustrin ranged from $-6$ to $3$ and from $-1$ to $4$, respectively (Additional File 15: Figure 13).

We found evidence of positive selection on specific codon sites in the Mature Peptide and the Acidic Propiece domains using the maximum likelihood models implemented in CODEML [46]. The model allowing for selection fitted the data significantly better than neutral models only for the Temporin-Brevinin group (Additional File 2; Table 2, Additional File 16; Figure 14 and Additional File 16), while the evidence of selection on specific codon sites in the Palustrin group was not significantly better for the selection model compared to the neutral model. The models allowing for selection (PAML, FEL and REL) identified a total number of 14 sites under selection in Temporin-Brevinin and five in Palustrin. The Mature Peptide domain displayed 12 out of 14 and five out of five, for Temporin-Brevinin and Palustrin respectively, of the positively selected sites within the hypervariable region. Ten sites (3, 21, 40, 44, 47, 49, 50, 51, 56) out of 14 in Temporin-Brevinin group were identified by at least two methods.
different approaches with posterior probabilities > 0.95 (See Aditional File 2).

**Discussion**

We studied variation in AMP genes in two amphibians along a latitudinal gradient from northern Germany to the northern margin of distribution of the two species (~2000 km). We investigated the relative contribution of selective forces, drift/demography and migration driving AMP genetic variation. We expected that genetic variation at the AMP loci would be lower towards the species distribution margin in the north [40]. Contrary to our expectations, we found genetic variation in all AMPs was largely equally distributed among the regions. The \( \Theta_k \) values did not differ systematically among populations and regions, suggesting similar effects of selection and demography shaping the genetic variation at AMP loci. However, we found evidence for reduced latitudinal variation at the peptide level in Brevinin for \( R. arvalis \), and in Temporin for \( R. temporaria \). Moreover, we found trans-specific polymorphism between \( R. arvalis \) and \( R. temporaria \), alleles in the Temporin and Brevinin group of genes clustering to a large extent by allele similarity. We also detected signatures of historical positive selection on AMP genes, as well as multiple lines of evidence suggesting strong diversifying selection maintaining adaptive variation within the Mature Peptide region.

Neutral vs. functional diversity?

As a consequence of decreasing effective population size \( (N_e) \) at range margins, many widely distributed species show lower genetic variation in populations at the edge of their distribution range [41, 42]. While \( R. arvalis \) and \( R. temporaria \) have slightly different postglacial recolonization history in Scandinavia, both species have colonized Scandinavian Peninsula both from the south (via Denmark) and the north (via Finland) and lost neutral genetic variability during this process [37, 40, 42]. Nevertheless, a study of Scandinavian \( R. arvalis \) found a clear and structured allelic distribution with a decreasing pattern of genetic variation towards northern latitudes at microsatellite loci as well as at the major histocompatibility complex (MHC) class II exon 2, an important component of the adaptive immune response [40]. These results were mainly explained by a complex pattern of varying levels of drift and selection along the latitudinal gradient which is influenced by colonization history of the species [40, 47]. Both in \( R. temporaria \) and \( R. arvalis \) the highest genetic variation both in neutral and adaptive allele frequencies is found in the southernmost regions of the latitudinal gradient [43, 48, 49].

We found that AMP alleles are largely equally distributed within species in \( R. arvalis \) and \( R. temporaria \) populations. Opposed to the findings on MHC, this study presents no clear loss of alleles towards the north of the gradient. However, we found a decrease in functional amino-acid diversity towards the northern latitudes in Brevinin in \( R. arvalis \) and in Temporin in \( R. temporaria \). This may suggest 1) independent effects of ecological and/or demographic conditions on genetic and amino-acid variability of AMPs, 2) differential importance of pathogens along the gradient 3) and/or drift. Furthermore, the differences in AMP diversity and allele/peptide geographic distribution along the gradient between the species might indicate different selective pressures acting on AMP genes in the two species.

In general, the intensity of species interactions decreases towards higher latitudes [50, 51]. The lower AMP amino-acid variability might be associated with the weaker biotic interactions at higher latitudes, which are strongly dependent on environmental conditions. Gurnier and co-workers [52] suggested that stochastic changes in environmental temperature, humidity and precipitation may affect parasitic and infectious microorganism diversity disproportionately, weakening the effect of pathogens at northern latitudes. Infectious diseases are also likely to be fewer in the north [50] and hence select for fewer and functionally similar AMP alleles at northern latitudes, while southern populations may have to cope with a more

| Loci                | Region      | A  | N  | S  | k    | \( \pi \) |
|---------------------|-------------|----|----|----|------|--------|
| Temporin-Brevinin    | Complete seq| 46 | 183| 63 | 26.20| 0.194  |
| Palustrin            |             | 12 | 141| 56 | 25.92| 0.288  |
| Temporin-Brevinin    | Mature Peptide | -  | 102| 31 | 14.439| 0.437  |
| Temporin-Brevinin    | Acidic propiece | -  | 117| 32 | 11.76| 0.115  |
| Palustrin            | Mature Peptide | -  | 87 | 50 | 23.136| 0.35   |
| Palustrin            | Acidic propiece | -  | 39 | 6  | 2.78 | 0.116  |
diverse flora of pathogens. Also anthropogenic alteration of natural environments and its effect on the intensity of species interactions can be weaker in the north [53]. We hypothesize that the latitudinal variation in biotic and abiotic interactions may contribute to the observed functional pattern found at amino-acid level. The lower genetic variation commonly observed at high latitudes [40, 54, 55] might suggest lower adaptive potential of AMPs (Temporin in *R. temporaria* and Brevinin in *R. arvalis*) against climate mediated changes in parasite and disease regimes in northern populations.

Our study highlights the need for additional work on the genetic basis for disease resistance in amphibians facing population declines in many parts of the world due to emerging infectious diseases [21, 56–58]. *Bd* infections are widespread in southern and central Sweden [59, 60], but data from northern Scandinavia are largely lacking [60, 61]. Several studies have investigated the complex relationship between skin microbiota composition and AMPs as a direct measure to fight *Bd* infections (e.g. [62, 63]), while other studies have shown that purified AMPs inhibit *Bd* growth under laboratory conditions [64, 65]. Therefore, understanding how genetic variation in AMPs is distributed across large spatial scales is important for future studies and may help to explain direct effects of AMPs on disease resistance and *Bd*-related declines.

**Copy number and trans-species polymorphism variation**

Other vertebrates have evolved multiple copies per individual in AMPs [66] and specifically in some amphibian species [31]. However, this is the first study to show that AMP group of genes in *R. arvalis* and *R. temporaria* is a clear multi-locus system by using NGS techniques with multiple alleles per individual as a result of gene duplication. A minimum of four loci were found in Temporin and three in Brevinin and Palustrin. The importance of duplicated genes as a reservoir of raw genetic material and its contribution in evolution has been recognized for long [67], but very little is known about AMPs copy number variation and their location within amphibian genomes. Earlier studies have shown the presence of more than one MHC class II loci in several species [68–70], but before our study this was not described for AMPs along a latitudinal gradient in ranids.

One of our main results is that the alleles within each AMP clade/paralog were clustered by allele and not by species, and similar alleles occurred in both species. This pattern is a hallmark of trans-specific polymorphism (TSP), which has been found especially in immune genes in a variety of taxa [71–73]. In our study, topologies for phylogenetic trees based on nucleotide sequences constructed for Brevinin, Temporin and Palustrin group of genes show that alleles share more similarity between, rather than within, species. This suggests that these AMPs have originated before the two species diverged at least 3.0 mya [74] or more than 10 mya [75]. Therefore, incomplete lineage sorting or hybridization is an unlikely explanation for the observed TSP. Our results are in concordance with Duda et al. [6] who showed that all alleles in Temporin, Ranalexin and Gaegurin loci clustered together within the ranid phylogeny, strengthening the idea of AMPs originating via concerted evolution before the divergence of the species.

Temporin alleles in *R. arvalis* (Raar_Temp*23,24,11, 25,27,22,19,18,16,17,30) were grouped in two distinct clades, supporting a recent genetic duplication of Raar_Temp*23,24,11,25 in *R. arvalis* after the two species diverged. Rest of the alleles (Raar_Temp*27,22,19,18,16,17,30) were more similar to Brevinin than Temporin alleles. The similarity between Temporin and Brevinin group of genes can be explained by several duplication events followed by subsequent episodes of evolutionary divergence. To demonstrate TSP conclusively, further research is needed to elucidate the causes of similarity between the groups of loci and the locations of these genes within the genomes.

**Selection on AMPs**

Our results indicate that the AMPs in *R. temporaria* and *R. arvalis* exhibit the same characteristics as observed in other amphibians [6, 16, 17], including high levels of polymorphism, differences in sequence length and strong signatures of selection within the Mature Peptide region (C-terminal antimicrobial-coding region). We also found that the three antimicrobial peptides have evolved in a coordinated manner as a result of a maintained charge balance between the Acidic Propiece and the Mature Peptide over evolutionary time, which is in concordance with studies on mammalian defensins [76]. Interestingly, this was not the case in an earlier study [6], which suggested AMPs in hylids, but not in ranids, evolved in a coordinated manner. We hypothesize that under the coordinated evolution scenario, evolutionary processes might be driven by selective forces derived from the composition of the pathogen communities to which the species and populations are exposed. However, no data exist to test this hypothesis. Therefore, further investigations are needed regarding pathogen community composition in specific environments and how the bacterial communities drive coevolution in AMPs.

We found footprints of historical positive selection on specific amino-acid positions within both the Acidic Propiece and the Mature Peptide. The majority of the codons under positive selection are situated within the Mature Peptide, suggesting a strong selective pressure on the hypervariable Mature Peptide region, and more relaxed selection within the Acidic Propiece. Pathogen-driven selection is a well-known mechanism in MHC
genes (e.g. [77–80]) but how these mechanisms act on AMP genes is not clear. Following the same principle as for MHC genes, a potential explanation for a strong selective pressure within the Mature Peptide domain in AMP genes may be due to differences in pathogen species composition along the latitudinal gradient. Moreover, temporal variation in pathogen-driven selection could also play a role on the Mature Peptide domain. In general, our results confirm the hypothesis that the hypervariable Mature Peptide domain is a result of adaptation to challenges by pathogens leading to increasing genetic diversity [6, 16]. Signatures of selection found this study could be contrasted with other markers such as neutral markers. However, given the complexity of multi-locus systems, such analyses of selection should be interpreted with caution.

**Conclusions**

We investigated genetic diversity as well as selective forces shaping AMP diversity in two species of amphibians over a latitudinal gradient. We described novel AMP sequences and high AMP genetic variation in both species. We found strong evidence for trans-specific polymorphism and the estimated number of loci is high for each AMP. Our data support positive selection within the Mature Peptide domain. The evolution and diversification of this extensive family of hypervariable genes can be due to the contribution of different processes such as speciation events, gene duplication and targeted hyper-mutations, and we suggest the gene complex has ultimately evolved under diversifying selection. From a conservation perspective, identifying and characterizing AMPs is important for ultimately understanding the adaptive processes and the ability of populations to combat novel or altered pathogens. Hence, our results will facilitate further studies on the evolutionary and conservation ecology in amphibians.

**Methods**

Sample collection and DNA extraction

We sampled *R. arvalis* eggs in five regions, from northern Germany (Hanover) to northern Sweden (Luleå, Table 3). The eggs were collected at three sites in each region, with the exception of one region (Umeå), where we collected at two sites. For *R. temporaria*, we took samples in seven regions from northern Germany (Hanover) to northernmost Finland (Kilpisjärvi). We collected the eggs at three sites in each region except the regions in northern central Sweden (Umeå), northernmost Sweden (Kiruna) and northern Finland (Kilpisjärvi), where we collected eggs at two, two and one site, respectively. The eggs were collected in spring 2014 and 2015 except in Kilpisjärvi where the eggs were collected in 2009. The average distance between collection sites within a region was 20 km (range 8 to 50 km) for both species (Table 3). At each site we collected ca. ten eggs from ten freshly laid clutches. The species coexist in many of our sampling locations and the species were identified in the field by differences in the color of the jelly surrounding the eggs [81]. The eggs were transported to the laboratory in Uppsala and kept at 16 °C. After hatching, the tadpoles (stage 25, [82]) were euthanized with an overdose of MS222, preserved in 96% ethanol and stored at 4 °C until DNA extraction.

Genomic DNA was extracted from ten individuals per site (one individual/clutch). We extracted genomic DNA from a total of 320 individuals (150 *R. arvalis* and 170 *R. temporaria*), using the DNeasy Blood and Tissue kit (Qiagen® Sollentuna, Sweden). Purity and concentration of DNA were determined with a NanoDrop® 2000 spectrophotometer and Qubit®3.0 fluorometer Quantitation Kit (Invitrogen™). Species verification was carried out by mtDNA cytochrome b amplification followed by the addition of HaeIII restriction enzyme (Palo and Merilä 2003). Digestion by HaeIII produces different, easily distinguishable banding patterns in *R. arvalis* and *R. temporaria*.

Primer design, antimicrobial peptide amplification and preparation for sequencing

We used the primers designed by [17] for the amplification of Brevinin loci (see Table 4). PCR products were visualized and isolated from agarose gels (1.5%). The targets bands were excised from the gel and extracted using the MinElute Gel Extraction Kit (Qiagen® Sollentuna, Sweden) as Brevinin primers produced multiple PCR bands in *R. arvalis* and *R. temporaria*. We sequenced the resultant PCR product from the 3’end and 5’end and used it to design the primer pair BrevMF1 (5’-TTCAGTTTTGTGGCATCCCG-3’) and BrevR6 (5’-CAAGTTTCCAAAGTTCAACAT-3’). We designed primers for Temporin loci (summarized in Table 4) using several available *R. temporaria*, *R. japonica* and *R. versabilis* temporin cdNA sequences (Accession number: YO93934.1, YO93935.1, YO93935.1 [83]; Accession: AB593684.1 [84]; Accession: AM113510.1 [85]; PCR products were visualized and isolated from agarose gels (1.5%). The target bands were excised from the gel and extracted using the MinElute Gel Extraction Kit (Qiagen® Sollentuna, Sweden). The resulting PCR product was partially sequenced from the 3’ and used to design the primer pair TempoF6 (5’-GTCAGGATTGAAATACAGTTTGTGTA-3’) and TemporinR3 (Table 4). Finally, we designed primers for Palustrin loci: PaluF1, PaluR2 (summarized in Table 4) using available Palustrin cdNA *R. versabilis* sequences (Accession: AM113507.1, AM745092.1 [86]; PCR products were visualized and isolated from agarose gels (1.5%). The resulting product was sequenced for DNA-sequence and fragment length verification. All PCR products were standard Sanger-sequenced at Macrogen Europe (Netherlands).
The forward primers were positioned within the "Signal Peptide" (Brevinin, Temporin) or in the "Acidic Pro-piece" (Palustrin) region, the most conserved regions of the antimicrobial peptides, and all the reverse primers were positioned in the 3′ UTR-region (Fig. 1). The length of the fragments varied from 100 bp to 204 bp among loci and alleles within each locus. For Illumina Miseq sequencing, both forward and reverse primers for Temporin, Brevinin and Palustrin were modified with an individual 8 bp barcode and a sequence of three N (to facilitate cluster identification). Each amplicon was marked with an individual combination of a forward and a reverse barcode for identification. Reactions were conducted in a total volume of 20 μl containing 1 μl of genomic DNA, 2 μl of 10X Dream taq buffer (Thermo scientific lab), 0.4 μl of 2 mM of each dNTP, 0.5 μl of each 10 μM primer (BrevMF1-BrevR6; TempMF6-TempoR3; PaluF1-PaluR2, respectively), 1.5 μl of Bovine serum albumine (BSA; 5 mg/ml) and 0.25 μl of Dream taq DNA polymerase (5 U/μl, Thermo scientific lab) in deionized water. Thermocycling was performed on an ABI 2720 (Applied Biosystems®). All reactions were carried out using filter tips in separate (pre- and post-PCR) rooms, and negative controls were included in all amplifications to avoid contaminations. PCR products were run and visualized on a 1.5% agarose gel using gel green (BIOTIUM). To reduce the number of samples for subsequent purification, 3–9 PCR products with similar concentrations were pooled based on visual estimations from the gel image. These sample pools were run on a 1.5% agarose gel, the target band was excised from the gel and extracted using the MinElute Gel Extraction Kit.
The concentration of each sample pool was measured with Qubit 4 Fluorometer dsDNA assay kit (Invitrogen Life Technologies, Stockholm, Sweden). The final amplicon pooling was conducted according to the measured concentrations and consisted of equimolecular amounts of each sample. A total of eight final amplicon pools were generated per run, and libraries were prepared using the Illumina Tru-seq DNA PCR-Free Sample preparation kit (Illumina Inc., San Diego, CA). Eight pools each were combined into a Miseq run, and sequencing of four Miseq runs was carried out at the National Genomic Infrastructure (NGI), the SNP&SEQ Technology Platform hosted at SciLifeLab in Uppsala (Sweden).

Miseq data analyses

Sequencing data were extracted from the raw data and paired-end reads were combined into single forward reads using FLASH [87], each of the eight amplicon pools was analyzed independently. In total, eight fastq files were generated per group of genes and transformed into fasta (multifasta) files using Avalanche NextGen package (DNA Baser Sequence Assembler v4 (2013), Heracle BioSoft, www.DnaBaser.com). The jMHC software [88] was used to remove primer sequences and unique tags, and to generate alignments of all alleles per amplicon. Generally, in multilocus system studies using NGS techniques, rigorous quality control and filtering procedures have to be applied to distinguish PCR and sequencing artefacts from true alleles. In this particular case amplicon coverage and replication rate were remarkably high (ca. 20%). We assigned the most frequent alleles within each amplicon as valid AMP alleles that occurred in at least 3% of the reads [89, 90]. We discarded amplicons with <300 reads from the analysis for quality reasons. In addition, we used the DOC method [91], not assuming any specific number of loci to identify and estimate the number of alleles (Ai) per individual. This procedure is based on the break point in sequencing coverage between alleles within each individual and avoids choosing a subjective threshold to separate true alleles from artefacts. Alleles are sorted top-down by coverage, followed by the calculation of the coverage break point (DOC statistic) around each allele. The allele with the highest DOC value is assumed to be the last true allele (see [91]). All valid alleles were imported and aligned by ClustalW in MEGA v7.0 [92]. Alleles were extensively compared to other sequences from the same putative locus in order to define the “Mature Peptide” and the “Acidic Propiece” region boundary within our alleles. Valid AMP alleles were named following the nomenclature suggested by [93] for MHC loci: a four-digit abbreviation of the species name followed by gene species*numeration, e.g. Raar_Brev*01.

Data analyses

Relative allele frequencies were estimated for each AMP (Brevinin, Temporin and Palustrin) with ARLEQUIN v. 3.5 [94]. Allele frequency plots were created in R using the “ggplot2” package (Wickham 2016). We calculated the number of pairwise nucleotide differences by population (Theta λ) and within region (Theta λ Region) with DNAsp v.5 (Librado and Rozas 2009). We tested for differences in number of loci between the populations, regions and species by running a Generalized Mixed Model (GLMM) in R using the package ‘lme4’ [95]. Allele frequency was...
considered as the response variable, region and species as fixed factors and (population: region) as random factors of the model.

We constructed two phylogenetic trees to illustrate the phylogenetic relationship among AMPs sequences from different gene families (Temporin, Brevinin and Palustrin group of genes): 1) separately by genes, 2) all genes together, present in both species. We used the Neighbor method with bootstrapping (1000 replicates) implemented in MEGA v7.0 [92]. We also constructed a haplotype network per gene and per gene and species by using Minimum Spanning network inference [96] in the software PopART.

Relative amino-acid frequencies were estimated for each AMP with ARLEQUIN v.3.5 (Excoffier, et al. 2010). Allele frequency plots were created in R using the “ggplot2” package [97]. AMP nucleotide sequences were grouped according to the phylogenetic tree based on amino-acid sequences. Some nucleotide sequences differed only by synonymous substitutions and translated into the same amino acid sequence (e.g Rate_Temp*06, Rate_Temp*22 and Rate_Temp*08 were named as Temp_Amino*01; see the phylogenetic tree based on Amino-acid sequences; (Additional File 8: Figure 6).

Nucleotide diversity (n), number of segregating sites (S), average number of pairwise nucleotide differences (Theta k) and Tajima’s D (D) were calculated for the entire AMP region, for the Acidic Propiece and for the Mature Peptide for every locus in DNAsp v.5 [98]. Net charges were calculated at Peptide property calculator platform. Signatures of natural selection at specific codon sites were also detected by using the common method on order to simplify the haplotype, network every individual in the plot is reconstructed with neighbor join methods. Name of sequences was reconstructed with neighborjoining method for: a) Temporin, b) Brevinin and c) Palustrin. Bootstrap values from 1000 replicates greater than 50% are indicated on branches. Alleles that belong to the same group are included in the same colored square. Alleles shared by both R. arvalis and R. temporaria are shown in bold. Valid alleles were named following the nomenclature suggested by Klein (1975) for MHC loci: a four-digit abbreviation of the species name followed by species_gene*numeration, e.g. Raar_Brev*01.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s12863-020-00839-1.
single individual. Populations within regions along the gradient are represented with the same colour.

Additional file 11: Figure 9. Minimum Spanning Network representing Temporin nucleotide variants in R. temporaria. Every circle represents one single individual. Populations within regions along the gradient are represented with the same colour.

Additional file 12: Figure 10. Minimum Spanning Network representing Brevinin nucleotide variants in R. temporaria. Every circle represents one single individual. Populations within regions along the gradient are represented with the same colour.

Additional file 13: Figure 11. Minimum Spanning Network representing Palustrin nucleotide variants in R. temporaria. Every circle represents one single individual. Populations within regions along the gradient are represented with the same colour.

Additional file 14: Figure 12. Minimum Spanning Network representing Palustrin nucleotide variants in R. temporaria. Every circle represents one single individual. Populations within regions along the gradient are represented with the same colour.

Additional file 15: Figure 13. Net charge of the Acidic Propiece domain is relatively related to the net charge of the Mature Peptide domain for a) Temporin and Brevinin sequences and b) Palustrin sequences. The regression line is plotted in green and brown, respectively in A) and B).

Additional file 16: Figure 14. Antimicrobial peptide alignment. The Signal Peptide is re-presented in blue, the Acidic Propiece in black and the Mature Peptide in red. Codons under selection were marked with an asterisk (*). Codons under selection detected at least by two different methods are colored in yellow. A) represent Temporin-Brevinin group of genes and B) palustrin group of genes, respectively.

Abbreviations

AMPs: Antimicrobial peptides; DOC: Degree of change; MHC: Major histocompatibility complex

Acknowledgements

We thank Emilien Luquet, Frank Johansson, Alberto Corral, Miriam Rubin and Cátia Chaves for invaluable help in the field. Many thanks to Gunilla Engström for support in the lab and Alba Cortáz-Chinarro (A.Cortazar) for helping with the illustrations. We also thank Anders Hallengren (Länstyrelsen Skåne), Anders Forsgren and Stefan Andersson (Piteå Kommun), Thomas Brandt (Ökologische Schutzstation Steinhuder Meer e.V), and Bernd Rittberg (Authority for Nature Protection of Lower Saxony and the Hanover region) for helping with the sampling permissions.

Authors’ contributions

MC collected the data, did the practical work in the laboratory, analyzed the data, participated actively in the conception of the study and wrote the manuscript. ARB helped in the field work and made contribution to the conception of the study. TVW helped with the haplotype networks. YML, AL and JH made substantial contribution at the conception, design, analyses and interpretation of data. All the authors read and approved the final version of the manuscript.

Funding

Funding was provided by the Swedish Research Council Formas (grant 46400178 to JH), Stiftelsen Oscar och Lili Lamms Minne (grant D0213–0013 to JH), Swedish Research Council (grant 621–2013-4503 to AL) and Stiftelsen för Zoologisk Forskning. The funding bodies played no role in the design of the study and collection, analyses and interpretation of data and in writing the manuscript.

Availability of data and materials

The datasets used and/or analysed during the current study are available from the corresponding author on request.

Ethics approval and consent to participate

Sample collection was conducted in Germany and Sweden according to the relevant sample collection guidelines. Permission were approved from the administrative institutions (The County board [Sweden] and The Authority for Nature Protection of Lower Saxony and the Hanover region in [Germany]) in the respective countries.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interest

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Received: 14 August 2019 Accepted: 6 March 2020

Published online: 30 March 2020

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