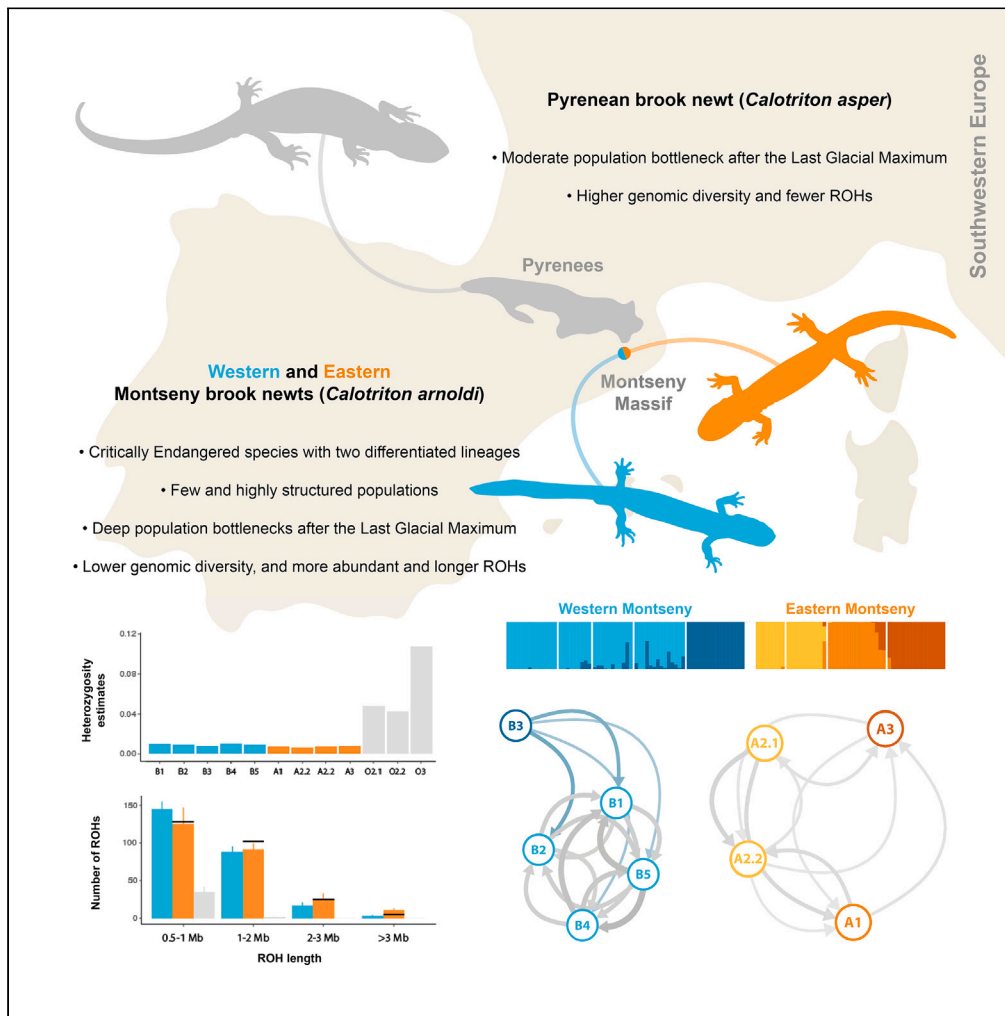


Article

Genomic insights into the Montseny brook newt (*Calotriton arnoldi*), a Critically Endangered glacial relict



Adrián Talavera, Marc Palmada-Flores, Bernat Burriel-Carranza, ..., Elena Obon, Tomàs Marquès-Bonet, Salvador Carranza

adrian.talavera@csic.es

Highlights

Third genome assembly of a urodele and first genomic study of the Montseny brook newt

Inferred demographic oscillations are accurately predicted by past climate changes

Bottlenecks have led to low genomic diversity and runs of homozygosity

Montseny brook newt populations are highly structured due to the lack of land dispersal

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Article

Genomic insights into the Montseny brook newt (*Calotriton arnoldi*), a Critically Endangered glacial relict

Adrián Talavera,^{1,13,14,*} Marc Palmada-Flores,^{1,13} Bernat Burriel-Carranza,^{1,2} Emilio Valbuena-Ureña,³ Gabriel Mochales-Riaño,¹ Dean C. Adams,⁴ Héctor Tejero-Cicuéndez,^{1,5} Anna Soler-Membrives,⁶ Fèlix Amat,⁷ Daniel Guinart,³ Francesc Carbonell,⁸ Elena Obon,⁸ Tomàs Marquès-Bonet,^{1,9,10,11,12} and Salvador Carranza^{1,12}

SUMMARY

The Montseny brook newt (*Calotriton arnoldi*), considered the most endangered amphibian in Europe, is a relict salamandrid species endemic to a small massif located in northeastern Spain. Although conservation efforts should always be guided by genomic studies, those are yet scarce among urodeles, hampered by the extreme sizes of their genomes. Here, we present the third available genome assembly for the order Caudata, and the first genomic study of the species and its sister taxon, the Pyrenean brook newt (*Calotriton asper*), combining whole-genome and ddRADseq data. Our results reveal significant demographic oscillations which accurately mirrored Europe's climatic history. Although severe bottlenecks have led to depauperate genomic diversity and long runs of homozygosity along a gigantic genome, inbreeding might have been avoided by assortative mating strategies. Other life history traits, however, seem to have been less advantageous, and the lack of land dispersal has driven to exceptional levels of population fragmentation.

INTRODUCTION

During the Quaternary, the shifting climate of Europe and its complex geography, resulted in an intricate dance of expanding and retracting species ranges.¹ While warm periods fostered the recolonization of the continent by some thermophilic taxa expanding from diverse glacial refugia, alpine species ranges shattered into patches, and conversely during glaciations. Such fragmentation into the so-called alpine archipelago, resulted in copious examples of disjunct distributions and, ultimately, in vicariant sibling species.² One example are the newts in the genus *Calotriton*, comprising the relatively widespread Pyrenean brook newt *C. asper* and the Montseny brook newt *C. arnoldi*. Whereas the former inhabits the Pyrenees and neighboring lesser mountain ranges, the latter is a Critically Endangered microendemism exclusive to the Montseny Massif (Catalonia, NE Spain; Figure 1).³

At this Eurosiberian exclave, the Montseny brook newt is thought to have been evolving in isolation for the last 1.76 million years (my).⁴ Currently, natural populations of this species are scattered across only eight streams, barely covering an occupancy area of 10 km².⁵ Although this minute range might not seem prone to fragmentation, the deep Tordera River valley divides the Montseny into two submassifs from West to East, breaking apart two sets of populations. The connectivity between them is disrupted not only because of geographical barriers, but also due to inherent biological features of the species: whereas *Calotriton asper* undergoes a 2-year-long terrestrial stage following metamorphosis,⁶ dispersal in the Montseny brook newt is greatly hampered by the lack of functional lungs.³ This adaptation to avoid undesired buoyancy in strong-current brooks makes them fully dependent on groundwater during dry periods, and directly restricts dispersal to waterway routes. In turn, the 5 km width of the Tordera River valley represents a largely unsuitable >30 km water pathway, crossing unfavorable

¹Institute of Evolutionary Biology (CSIC-Universitat Pompeu Fabra), Barcelona, Spain

²Museu de Ciències Naturals de Barcelona, P^o Picasso s/n, Parc Ciutadella, 08003 Barcelona, Spain

³Servei de Gestió de Parcs Naturals, Diputació de Barcelona, Spain

⁴Department of Ecology, Evolution, and Organismal Biology, Iowa State University, Ames, IA 50010, USA

⁵Department of Biodiversity, Ecology and Evolution, Faculty of Biology, Universidad Complutense de Madrid, 28040 Madrid, Spain

⁶Departament de Biologia Animal, de Biologia Vegetal i d'Ecologia, Universitat Autònoma de Barcelona, Barcelona, Spain

⁷Àrea d'Herpetologia, BiBIO, Museu de Granollers – Ciències Naturals. Palauàries 102, Granollers, Barcelona, Spain

⁸Centre de fauna salvatge de Torreferrussa (Forestal Catalana, SA), Santa Perpètua de Mogoda, Spain

⁹CNAG-CRG, Centre for Genomic Regulation (CRG), Barcelona Institute of Science and Technology, Barcelona, Spain

¹⁰Institut Català de Paleontologia Miquel Crusafont, Universitat Autònoma de Barcelona, Cerdanyola del Vallès, Spain

¹¹Catalan Institution of Research and Advanced Studies (ICREA), Barcelona, Spain

¹²Senior authors

¹³These authors contributed equally

¹⁴Lead contact

*Correspondence: adrian.talavera@csic.es

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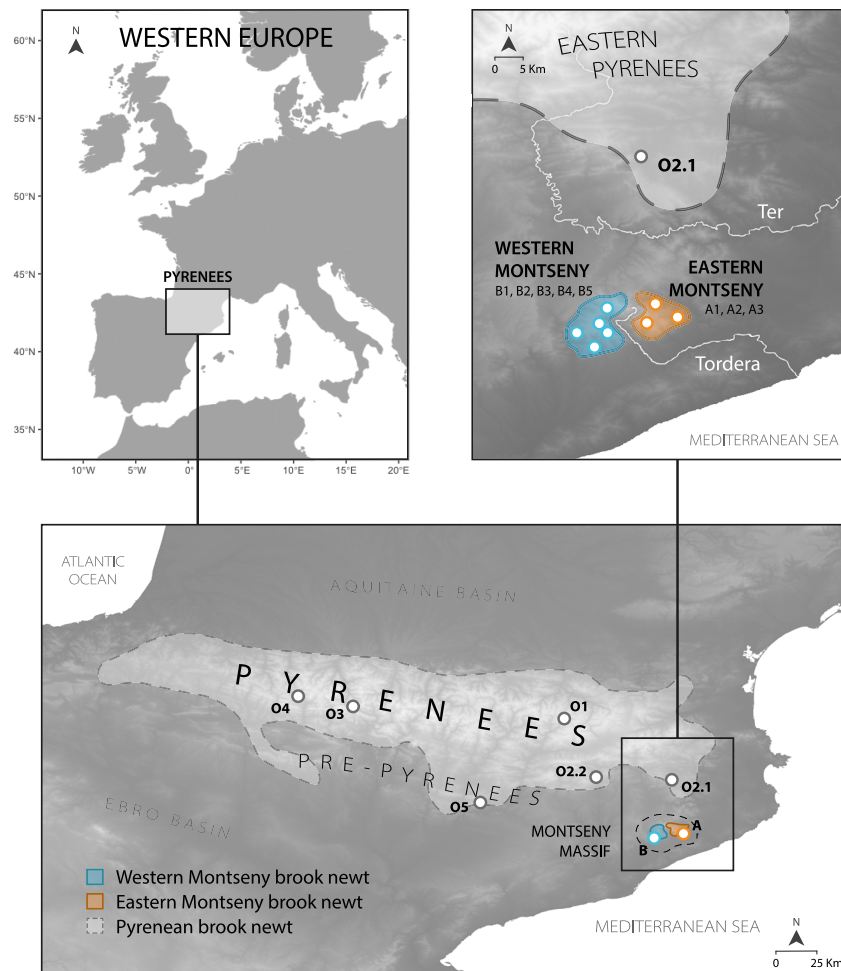


Figure 1. Distribution of brook newts *Calotriton* spp. in Western Europe

First close-up (below) shows the Pyrenees and adjacent mountain ranges, including the Montseny Massif (dashed). Second close-up (top-right) shows the area from this massif to the neighboring Eastern Pyrenees, and the rivers Ter and Tordera (in white), which divide the ranges of the Pyrenean-Montseny brook newts, and Western-Eastern Montseny lineages, respectively. Montseny brook newt (*Calotriton arnoldi*) range in blue (Western lineage) and orange (Eastern lineage). Pyrenean brook newt (*Calotriton asper*) distribution in light gray, within dashed line. White circles identify the sampling localities used in this study. Sampling points at Montseny do not correspond to exact geographic localities for conservation reasons.

environments for this strictly aquatic amphibian. Consequently, genetic studies showed that the Western and Eastern *C. arnoldi* sets of populations constitute two differentiated lineages, isolated since the late Middle Pleistocene.⁴ The combination of long-term isolation and small population sizes has resulted in incipient morphological differentiation, including the fixation of distinct color patterns. However, contrary to expectations, genetic studies did not find reduced diversity compared to *C. asper*.^{4,7}

Loss of genetic variability has been theoretically and empirically linked to decreases in the mean fitness and evolutionary potential of populations.^{8–11} In amphibians, genetic erosion has been found to be an important factor diminishing resistance to the parasitic fungus *Batrachochytrium dendrobatidis* in anurans.¹² *Batrachochytrium* infections are one of the main global drivers of declines in amphibian biodiversity.^{13,14} While *Batrachochytrium salamandrivorans* wreaks havoc on urodele populations across Europe¹⁵ and already caused an outbreak in a Natural Park 15 km south of Montseny in 2018,¹⁶ its lethal effect in *Calotriton arnoldi* has been confirmed by lab trials.¹⁷ On the other hand, groundwater-dependent amphibians are one of the most vulnerable components of global diversity due to rapid groundwater depletion at global and regional scales.¹⁸ Indeed, nearly one third of the total bottled water consumed in Spain is abstracted in the Montseny Massif and its surroundings.¹⁹ This groundwater overdraft has resulted in the desiccation of the brook headwaters, leading to the disappearance of the species in some of these areas.^{3,5} On top of that, despite being aquatic, the Montseny brook newt requires specific types of forests covering the streams. Unfortunately, some of these forests are being replaced by non-native timber plantations, while local species, such as beech trees, are shifting toward higher elevations due to the effects of global warming,²⁰ limiting the potential habitat available for the Montseny brook newt.

Altogether, the menace of emergent diseases and habitat loss due to global warming and human activities impose significant pressure on these small, fragmented, and declining populations, whose size has been estimated at approximately 1,000–1,500 adult individuals in the wild.⁵ This situation led to the declaration of the Montseny brook newt as the only ‘Critically Endangered’ urodele species in Europe by the IUCN, and one of the most threatened amphibian species in Europe⁵ with the Karpathos frog (*Pelophylax cerigensis*) (but see Toli et al.²¹ for the current need of conservation reassessment in the later). As a result, two years after its description as a new species, a provisional conservation program by the Catalan government began with the purpose of guaranteeing its future,²² followed later by a LIFE project (LIFE15 NATO/SE/000757) in 2016. Thanks to the genetic studies corroborating the existence of two well differentiated lineages,^{3,4,7} representatives from both submassifs were chosen to be founders of an international *ex situ* breeding program. Since then, these lineages have been treated as different management units, allowing no admixture to preserve their distinctiveness, despite the lack of any genomic insight. Ideally, any conservation effort should be guided by genomic studies, which provide valuable information such as effective population size, inbreeding, demographic history, and population structure.²³ However, those are yet scarce in urodeles, whose gigantic genomes²⁴ render the sequencing of enough data for reference genomes challenging.

Here, we present the reference genome of the Montseny brook newt, the third for the order Caudata,^{25,26} as well as exhaustive population genomic analyses based on 22 low-coverage whole genomes (lcWGS) and ddRADseq data for 132 individuals from all Montseny populations and several populations of the closely related Pyrenean brook newt *C. asper*. Our results reveal a series of population bottlenecks during the Late Pleistocene and Holocene, coinciding with warm periods in Europe, as well as their ultimate signatures: long Runs of Homozygosity and depauperate diversity values in all extant populations. Overall, this study highlights both the urge to take conservation actions under the current global climate crisis and the deep-rooted vulnerability of Europe’s most threatened amphibian species.

RESULTS

We generated the third *de novo* genome assembly for the order Caudata using two captive-bred siblings from one Eastern populations (A1) of the Montseny brook newt. This resulted in a genome assembly size of 22.89 Gb, N50 = 1.65 Mb and 82.4% of vertebrata ortholog completeness (Figures S1–S3; Tables S1 and S2) which was used to map both lcWGS and ddRADseq data from 21 to 132 wild animals, respectively. The sampling comprises all *C. arnoldi* natural populations and representatives of the five mitochondrial lineages of *C. asper*.

Brook newts’ evolutionary history was shaped by climatic oscillations

We found the effective population sizes of both brook newt species to be significantly dependent on climate oscillations along the Late Pleistocene and Holocene, showing negative correlations with waning ice sheets and, mainly, rising temperatures (Table S3), which explains up to 85% of *C. arnoldi*’s demographic trends. While large effective population sizes for both *C. arnoldi* and *C. asper* along the Late Pleistocene were estimated, population declines toward the present were also inferred (Figure 2). The Pyrenean brook newt shows a slow but constant decrease in effective population size, with the largest drop at the end of the Last Glacial Maximum (LGM, c. 26.5–19 kya). On the other hand, the Montseny brook newt suffered a more recent and severe bottleneck, coinciding with two warming periods: the Late Glacial Interstadial (LGI, c. 14.67–12.89 kya), and especially the sudden end of the Last Glacial Period (LGP, 110–11.65 kya), in which near-modern temperatures were reached. In between both events, the colder Young Dryas (YD, c. 12.89–11.65 kya) seems to have partially buffered their collapse. Nevertheless, during a brief warm period, the Roman Climatic Optimum (RCO, c. 2.25–1.55 kya), the effective population size reached its minimum with <500 effective individuals. Afterward, the populations recovered to some extent coinciding with the beginning of a locally colder and wetter period in modern Europe: the Little Ice Age (LIA, 14th–18th centuries).

Comparisons of simulations with different migration patterns between lineages support both recent migration between Montseny submassifs ($m_{arnoldi}$ in Table S4) and, at higher rates, ancestral migration from the Pyrenees into the Montseny (m_{asper} , Figure S4; Tables S4 and S5). Effective population sizes were lower for the Eastern Montseny *C. arnoldi* than for its Western counterpart, and both were extremely reduced compared to *C. asper*’s current N_e (N_A for current Eastern Montseny, N_B for Western Montseny, and N_O for the Pyrenean brook newt in Table S4). Large ancestral populations (N_{AB} for ancestral *C. arnoldi* population before lineage divergence and N_{ABO} for ancestral *Calotriton* population before species divergence in Table S4) and bottlenecks are in accordance with the demographic inference results from Stairway-plot analyses and lead to the idea of a progressive rarefaction of the genus.

At a larger scale, we found divergence events within the genus *Calotriton* to match certain cold or warm periods (Figure 3). Montseny and Pyrenean newts shared their most recent common ancestor 1.69 mya (95% highest probability density [HPD] intervals: 1.05–2.35 mya), partially coinciding with the Donau glaciation (1.8–1 mya). Divergence between *Calotriton asper* lineages, around 160 kya (101–258 kya), overlaps with another cold period: the Riss glaciation (300–130 kya). In contrast, divergence between Eastern and Western Montseny lineages seems to have taken place later, around 111 kya (68–155 kya), roughly matching with a warmer period: the Riss-Würm Interglacial (130–115 kya).

Genetic erosion and unexpected levels of fragmentation

Observed heterozygosity (H_O) for the Montseny brook newt was estimated to be between 10.18 and 21.14 (from lcWGS; Table S6) and 4.45–17.23 (from ddRADseq; Table S7) times lower for the Montseny brook newt than for its sister species, the Pyrenean brook newt. Moreover, H_O was slightly higher in the Western Montseny populations than in the Eastern ones according to both sequencing techniques, and at a finer scale, sparsely populated creeks (e.g., B3, A2.1) yielded lower values than populous well-connected demes (i.e., B1–2, B4–5). However, for all *C. arnoldi* populations (and two out of three *C. asper* demes), H_O was always higher than expected (H_E), suggesting absence of inbreeding

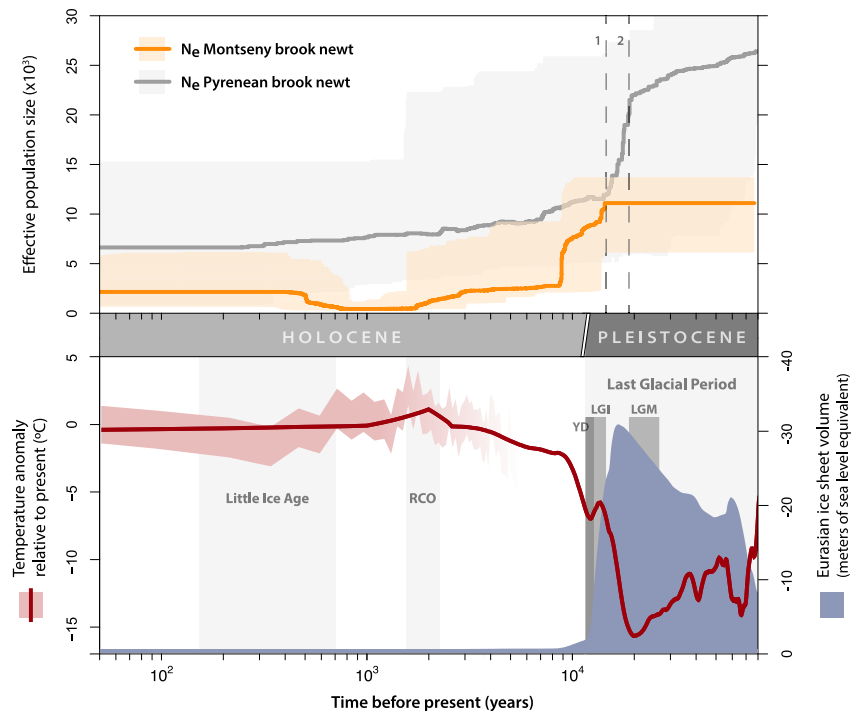


Figure 2. Demographic inference (above) coupled with climatic oscillations (below) along the Holocene and Late Pleistocene

Median effective population sizes ($N_e \times 1,000$) calculated with Stairwayplot2 for *Calotriton arnoldi* (orange line, below) and *C. asper* (gray line, above) are shown with corresponding 95% CI (areas shaded in orange or gray, respectively). Climatic oscillations are represented by the mean temperatures in the Northern Hemisphere (de Boer et al., 2013; red line) and West Mediterranean temperatures within 95% CI (red shaded area; Margaritelli et al., 2020). Eurasian ice sheet volume (blue area) is depicted as negative meters of the sea level in the corresponding time compared to the present. Time before present shown in logarithmic scale (x axis). The Pleistocene-Holocene transition and some other important climatic events are highlighted: the Little Ice Age, Roman Climatic Optimum (RCO), Last Glacial Period, Younger Dryas (YD), Late Glacial Interstadial (LGI) and Last Glacial Maximum (LGM). The two events associated with the main population bottlenecks in the brook newts are highlighted by dashed lines: 1) the beginning of the LGI for *C. arnoldi* and 2) the end of the LGM for *C. asper*.

(G_{15}) based on ddRADseq data (Figure 4A; Table S7). Low coverage WGS showed a higher number of ROHs in *C. arnoldi* compared to *C. asper*, including the presence of medium- and long-sized ROHs in the former, especially in the Eastern Montseny lineage (Figure 4B; Table S8).

Despite low levels of heterozygosity in the Montseny brook newt, the fixation of such diversity was unexpectedly high. The mean fixation index (F'_{ST} ; i.e., the percentage of shared diversity lost across a barrier in between demes) between Western and Eastern Montseny submassifs was higher ($F'_{ST} = 0.801$; $p < 0.001$) than between Western and Eastern Pyrenees *C. asper* ($F'_{ST} = 0.554$; $p < 0.001$), in spite of involving a much smaller geographical scale. Within Montseny lineages, population-pairwise fixation indices were relatively lower but still significant (Figure 4C; Table S9). Genomic PCAs showed agreeing results with both types of genomic data. First components of lcWGS and ddRADseq PCAs including *C. asper* explained a vast majority of the variance, sorting out the two sister species (Figures S5 and S6). Second and third components split lineages within *C. asper* Western (lineages O2–3) and Eastern Pyrenees (O1, O4–5) and *C. arnoldi* respectively. Within *C. arnoldi*, PCAs explained mostly the split between Western and Eastern submassifs, with A3 and B3 as the most divergent populations for each lineage.

Genus-level Admixture analysis with ddRADseq data recovered three clusters without admixture among them: *Calotriton asper*, East Montseny and West Montseny (reaching cross validation errors (CV) plateau at K3, Table S10; Figure 5A). Migration networks similarly supported the complete isolation between Montseny lineages, which represent two independent metapopulations without current migration between them (Figure 5B). Isolation-by-distance (IBD) alone did not fully explain genomic differences between Montseny submassifs ($p < 0.001$), despite considering water-restricted dispersal. However, structure within both Montseny lineages is concordant with the same IBD pattern (Figure S8). Although Admixture analysis within Montseny yielded the lowest CV errors at $K = 2$, separating again Eastern and Western Montseny lineages, $K = 5$ holds clear biological significance with a close CV mean value (Table S10), separating A1–A2 (which remain lumped at $K = 4$) and distinguishing the three Eastern inhabited brooks as different clusters (Figure 5A). Matching the high levels of isolation in the East, migration was found to be low within the metapopulation, even between A2.1–A2.2 (Figure 5B). These two subpopulations, despite belonging to the same creek and genomic cluster, show migration rates similar to other populations within the Eastern submassif, likely because a road, with the concomitant desiccation of a creek stretch, has artificially disconnected them. On the other hand, at the Western submassif, two putative clusters were identified: one population, B3, remains free from admixture with any other population, whereas the

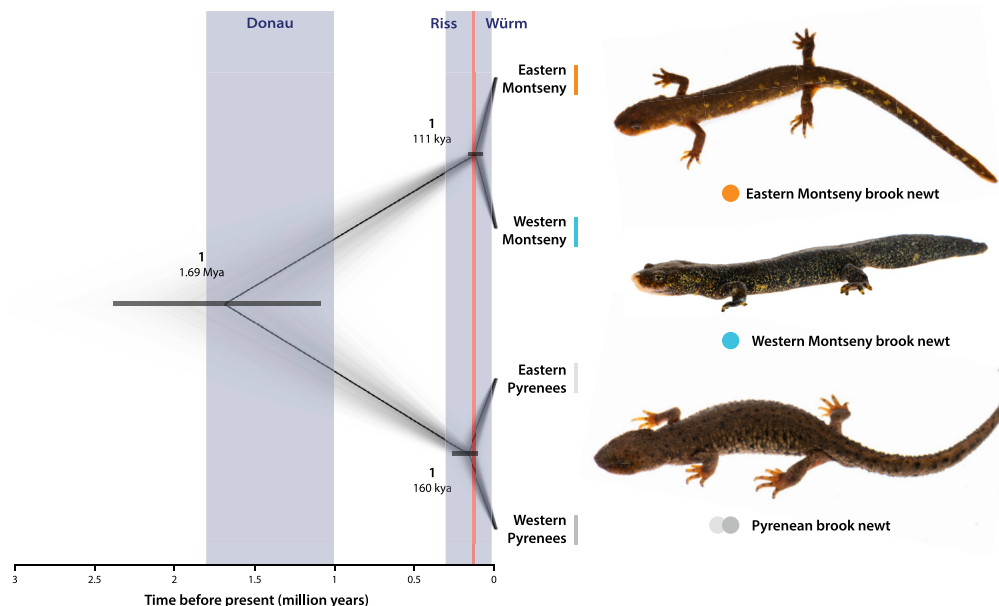


Figure 3. Genomic phylogeny of the genus *Calotriton*

Left, time-calibrated Bayesian phylogeny from ddRADseq data (cloudogram+consensus tree). The nodes are shown with associated posterior probabilities (in bold) and estimated mean times of divergence with their 95% CIs (horizontal gray bars). Three glacial periods (Donau, Riss and Würm/LGP) are highlighted in light blue, whereas the warmer Riss-Würm Interglacial is displayed as the red vertical line. Right, pictures of representative individuals of the three different brook newt lineages in the wild: from top to bottom, Eastern and Western Montseny brook newt *Calotriton arnoldi* and Pyrenean brook newt *C. asper*.

rest of the Western populations are merged into the same cluster (i.e., B1–2, B4–5), exhibiting high levels of connectivity (Figure 5B) and varying levels of admixture with B3 (Figure 5A). Migration from B3 to other Western populations appears to be significantly asymmetrical, being a source but not receiving migrant from other populations. This source-sink dynamic can be explained by the watershed architecture: B3 is physically isolated from the other populations by a large waterfall, rendering upstream dispersal impossible.

DISCUSSION

Evolutionary history

The tighter dependence of *C. arnoldi* population size on climatic oscillations than *C. asper* agrees with a narrower range of preferred body temperatures in the former,^{27,28} despite inhabiting more meridional and temperate places (Table S3; Figure 2). Such intimate relation between demographic trends and climatic oscillations seems to have shaped the evolutionary history of *Calotriton* spp. On the one hand, glacial periods (i.e., Donau and Riss) may have caused divergences in the Pyrenean–Montseny and the Western Pyrenees–Eastern Pyrenees species and lineages, respectively (Figure 3). Indeed, glaciations have been considered as drivers of transverse alpine speciation in temperate mountain ranges all around the globe, including the Pyrenees (see Wallis et al.²⁹). Species survival during glacial periods at high-altitude refugia,³⁰ is consistent with higher genomic diversity values in high altitude populations (Table S6). In addition, cold-adapted species are expected to thrive during glaciations and expand their ranges across ice-free suitable areas. Accordingly, our results infer larger population sizes for both species during the LGP (Figure 2). This is also supported by species distribution modeling (SDM) studies, which show, in turn, continuous potential distribution ranging from the Pyrenees to the Montseny Massif at the LGM.⁷ Moreover, a much wider range could vindicate a possible *Calotriton* fossil record quite beyond their current range (Cueva de las Hienas, Asturias³¹; see discussion in Lobo et al,³²). Putative expanding distributions and populations in the past would provide opportunities for ancestral inter- and intra-specific migration events that are in agreement with our results (Tables S4 and S5). However, evidence of periglacial events at the summit of the Montseny Massif at LGM³³ led to the hypothesis that *C. asper* never reached the Tordera River basin after divergence from *C. arnoldi* due to the harsh alpine conditions, suggesting that both species never came into secondary contact again.⁷ Regardless, migrants between the two Montseny lineages were probably never common due to their limited dispersal capacity, which could explain why they remained differentiated after divergence from *C. asper*. In contrast, *C. asper* expanded and occupied a larger, mostly continuous range after divergence from *C. arnoldi*, resulting in reduced isolation among its main lineages despite their older divergence compared to the Montseny brook newt lineages.

On the other hand, warm periods seem to have historically compromised the survival of the brook newts. First, the divergence of Western and Eastern Montseny lineages is associated with the Riss-Würm Interglacial (Figure 3), in which temperatures were similar or higher than the present. In this suboptimal scenario, *C. arnoldi*'s range probably split up into at least two marginal refugia at the headwaters of the different watersheds, and distribution has remained disjoint to this day. Secondly, increasing temperatures triggered severe population bottlenecks in

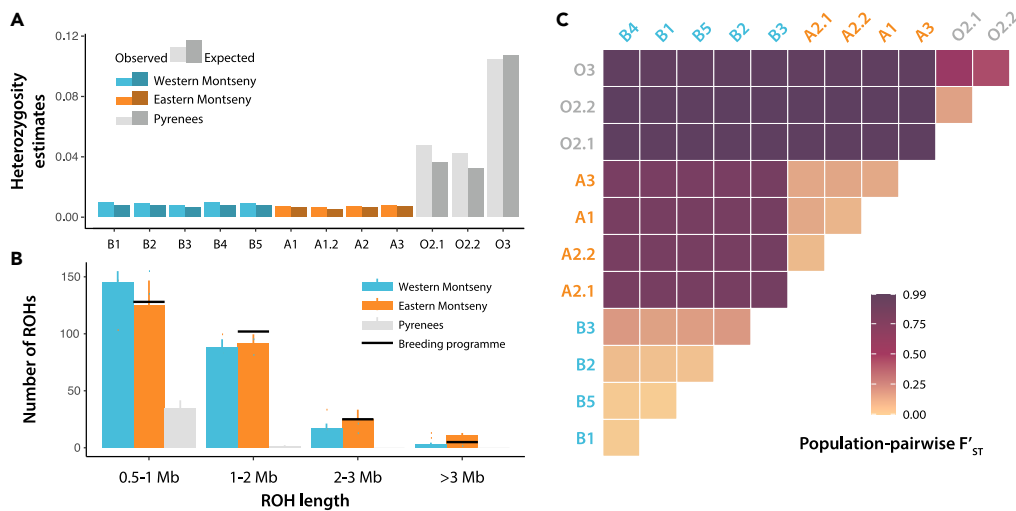


Figure 4. Genomic diversity, runs of homozygosity (ROHs) and fixation indices in genus *Calotriton*

(A) Population-level observed and expected heterozygosity estimates from 132 ddRADseq samples, including all Montseny and three Pyrenean populations. Observed heterozygosity is lower in Montseny brook newts than in Pyrenean brook newts, but is always higher than expected. (B) Bar plot of the mean + standard deviation counts of ROHs found in the Western Montseny (blue, $n = 10$), Eastern Montseny (orange, $n = 6$), and Pyrenean (gray, $n = 5$; *C. asper*) brook newts from lcWGS sorted by four increasing length categories. Black bars show the respective counts obtained from one F1 captive-bred Eastern individual used in the reference genome assembly. Montseny brook newts show a higher number of short ROHs than Pyrenean brook newts and, moreover, exhibit medium- and long-sized ROHs in their genomes, as expected in bottlenecked and isolated taxa. Within Montseny, the Eastern lineage shows more long-sized ROHs than the Western lineage. (C) Population-pairwise fixation indices (F_{ST}) among the 12 populations studied with ddRADseq. F_{ST} and significance values are shown in Table S9. Fixation of diversity between Western Montseny populations (in blue) and Eastern Montseny ones (orange) is more similar to fixation between *C. arnoldi* and *C. asper* (in gray) than between populations belonging to the same lineages.

brook newts, especially in *C. arnoldi* (Figure 2). This is reflected in a higher number of ROHs in the Montseny species compared with its sister taxon *C. asper* (Figure 4B; Table S8), as expected in isolated and bottlenecked populations.³⁴ Post-glacial population expansions after LGM and LGP have been inferred for many vertebrate groups,^{35–42} including Eurasian salamandrids from similar latitudes.^{43–45} However, well documented post-glacial bottlenecks are scarcer in the genomic literature and typically limited to either relict or extinct megafauna.^{37,46–49} Although paleodemographic modeling has statistically shown tight links between demography and climate oscillations for humans at the end of LGP from archaeological radiocarbon dates,^{39,50} to our knowledge this is the first time the dependence of a genomically inferred past demography on climate variables is proven, moreover, on a non-model species. Regarding future scenarios, a major decrease in the potential distribution of the Pyrenean brook newt and, concomitantly, of its genetic diversity and evolutionary history has been foreseen.⁵¹ In fact, the situation could be even more menacing for *C. arnoldi*, as climate change is expected to cause more grievous effects in species with low genetic diversity, small ranges, low effective population sizes, specific habitat requirements and limited dispersal capabilities,³⁰ all characteristics exhibited by the Montseny brook newt.

Population genomics

Previous studies based on microsatellites concluded that Pyrenean and Montseny brook newts exhibited similar genetic diversity,⁴ in contrast with our genomic estimates, which show that diversity in all Montseny populations is much lower than in Pyrenean populations, supported both by lcWGS (Table S6) and ddRADseq data (Figure 4A; Table S7). Although traditional genetics do not always disagree with genomics, this illustrates how conservation can benefit from the advent of Next-Generation Sequencing and its superior resolution power.^{52–55} Within Montseny, results show higher genetic diversity and a lower number of long ROHs in the Western Montseny lineage relative to the Eastern (Figure 4B; Table S8), as expected in larger and admixed populations.³⁴ Interestingly, the highest ddRADseq H_O value at the Eastern Montseny lineage was found in population A3 (Table S7), which is a small population that could be virtually extinct at present. Despite exhaustive searches in different seasons, no newts have been found at this locality (previously disturbed by intense human activities) since the storm “Gloria” lashed Montseny in January 2020 and changed the geomorphology of the brook. Such dire event clearly evidences how the survival of small-ranged endemisms can be compromised by stochastic phenomena.⁵⁶

Despite long ROHs and low genomic diversity in all *C. arnoldi*, we found no evidence of inbreeding. In fact, observed heterozygosity values higher than expected were already reported based on microsatellite data,⁴ leading to the hypothesis of putative assortative mating strategies to avoid the negative effects of breeding among close relatives. Assortative mating has been reported in a number of salamandrid species, either involving particular preferences by males⁵⁷ or, more usually, by females.^{58–60} However, those strategies do not take place exclusively

become detectable, i.e., several generations for spatial genetic structure to arise,^{65,66} we can conclude that the isolation of Montseny Massif populations into five clusters is long enough to be attributed to natural reasons. In contrast, and regarding the artificial fragmentation of A2 creek, recent human impacts on the environment (i.e., most roads) are relatively young in terms of animal generations. Therefore, only weak genetic differences are expected, and only methods inferring contemporary migration are able to detect a barrier effect.⁶⁷ The threatening impact of roads in amphibians is not new, and it has been linked to an increased mortality, reduced genetic diversity, habitat fragmentation and, ultimately, an increased risk of extinction by reducing demographic and genetic input from immigrants and decreasing the probability of recolonization after extinction,^{68–71} while road verges promote the expansion of invasive species⁶⁷ and, thus, of potential pathogen vectors.

Differences across populations at the Montseny Massif are driven not only by natural- and human-related barriers, but also by simple distance effects. In contrast with previous studies,⁴ probably because of the higher resolution of genomic approaches, our results point to the existence of IBD within Montseny submassifs, but not in between: i.e., geographic distance is not enough to explain differences between the two lineages (Figure S7). Rivers are one of the main geographical barriers shaping amphibian distributions,⁷⁰ in our case (Tordera River, Figure 1) due to unsuitable conditions in terms of temperature, pollution, and the presence of predatory fishes, but also because of the meagre dispersal abilities of *C. arnoldi* compared to *C. asper* (mean displacements of barely 7 m in a 2-year period⁷²). Low dispersal rates and small population sizes lead to an accelerated differentiation,⁷³ reflected in high fixation indices (Figure 4C; Table S9). F_{ST} from genetic data between Pyrenean brook newt populations had already shown values comparable to reptiles and amphibians inhabiting different islands within archipelagos.³⁰ Furthermore, our genomic data shows even higher values of isolation between Montseny lineages at a much smaller scale, comparable to values found in troglobiont amphibians across different cave systems,⁷⁴ in desert-dwelling frogs from fragmented populations >300 km apart,⁷⁵ or even in between *Calotriton* species (Figure 4C). Consequently, in view of their genomic and phenotypic differentiation, the Montseny brook newt lineages might need further taxonomic reassessment.

Limitations of the study

This study presents not only the third genome assembly in the order Caudata, but also the first population genomic study in the group based on whole-genome resequencing. Due to the large genome sizes in urodeles, the sequencing of these amounts of data is always challenging and, therefore, yields ultra-low coverages that compel to analyze genotype likelihoods instead of true genotypes. The lack of confidence in the called genotypes, added to mapping difficulties of short reads to a highly repetitive genome, could hamper the correct identification of ROHs. Furthermore, the fragmentary nature of the reference genome limits the amount of ROHs that can be detected, especially the longest ones. We prioritized the number of samples rather than their coverage for lcWGS, so that we could compare results not only between lineages but within groups and populations as well. In the same way, despite the lack of other similar whole-genome studies in urodeles to compare with, we ensured the soundness of the ultra-low coverage data by comparing it to ddRADseq results.

Regarding *C. asper*, the sampling includes fewer individuals, while its range is much wider. Although we have chosen representatives of the different mitochondrial lineages, other genomic clusters may remain undiscovered. Future studies on this species should improve the sampling by covering new areas and exploring the contact zone between the Western and Eastern Pyrenees to assess the extent of lineage isolation.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.isci.2023.108665>.

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AUTHOR CONTRIBUTIONS

A.T., T.M.-B., and S.C. conceived and designed the study. F.A., E.V.-U., S.C., A.T., B.B.-C., G.R., F.C., A.S.-M., and D.G. performed sample collection. M.P.-F. performed genome assembly and analyzed data. A.T. carried out laboratory work, analyzed the data, designed figures and wrote the manuscript with input from all other authors, who approved the paper in its final form.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
<i>Critical commercial assays</i>		
MagAttract HMW DNA kit	Qiagen	N/A
<i>Deposited data</i>		
Calotriton <i>arnoldi</i> Reference genome assembly (aCalArn1)	This study	PRJEB57772
ORG.one Nanopore data used for the assembly	This study	PRJEB57582
Illumina data for lcWGS population genomic analyses and Nanopore and Illumina data generated outside ORG.one used for the assembly	This study	PRJEB57773
ddRADseq data from 154 Calotriton <i>arnoldi</i> and <i>C. asper</i> individuals	This study	PRJNA1023478
<i>Software and algorithms</i>		
Cutadapt v3.2	Martin ⁷⁶ , 2011	https://github.com/marcelm/cutadapt
Meryl v1.7	Rhie et al. ⁷⁷ , 2020	https://github.com/marbl/meryl
Smudgeplot v.0.2.5	Ranallo-Benavidez et al. ⁷⁸ , 2020	https://github.com/KamilSJaron/smudgeplot
KMC v.3.1.1	Kokot et al. ⁷⁹ , 2017	https://github.com/refresh-bio/KMC
GenomeScope v2.0	Ranallo-Benavidez et al. ⁷⁸ , 2020	https://github.com/schatzlab/genomescope
Filtlong v0.2.0	N/A	https://github.com/rwrick/Filtlong
Flye v2.8.3	Kolmogorov et al. ⁸⁰ , 2019	https://github.com/fenderglass/Flye
BWA v.0.7.15	Li ⁸¹ , 2013	https://github.com/lh3/bwa
HyPo v1.0.3	Kundu et al. ⁸² , 2019	https://github.com/kensung-lab/hypo
Minimap2 v2.17-r941	Li ⁸³ , 2018	https://github.com/lh3/minimap2
purge_dups v1.2.5	Guan et al. ⁸⁴ , 2020	https://github.com/dfguan/purge_dups
BlobToolKit	Challis et al. ⁸⁵ , 2020	https://github.com/blobtoolkit/blobtoolkit
RepeatMasker v4.1.2	Smit et al. ⁸⁶ , 2013-2015	https://www.repeatmasker.org/
RepeatModeler v1.0.11	Smit & Hubley ⁸⁷ , 2008	https://www.repeatmasker.org/RepeatModeler/
FastQC v0.11.8	Andrews ⁸⁸ , 2010	https://www.bioinformatics.babraham.ac.uk/projects/fastqc/
fastp v0.23.1	Chen et al. ⁸⁹ , 2018	https://github.com/OpenGene/fastp
PicardTools v2.8.2	BroadInstitute, 2021	https://github.com/broadinstitute/picard
SAMtools v1.9	Danecek et al. ⁹⁰ , 2021	https://github.com/samtools/samtools
ipyrad v0.9.78	Eaton & Overcast ⁹¹ , 2020	https://github.com/dereneaton/ipyrad
PLINK v2.0	Purcell et al. ⁹² , 2007	https://www.cog-genomics.org/plink/2.0/
easySFS	N/A	https://github.com/isaacovercast/easySFS
Stairway Plot v2.1.1	Liu & Fu ⁹³ , 2020	https://github.com/xiaoming-liu/stairway-plot-v2
R v.3.6.3	R Core Team ⁹⁴ , 2021	https://www.r-project.org/
RRPP v. 1.3.1	Collyer & Adams ⁹⁵ , 2018	https://github.com/mlcollyer/RRPP
nlme v3.1-140	Pinheiro et al. ⁹⁶ , 2016	https://github.com/cran/nlme/blob/master/R/lme.R
fastsimcoal2	Excoffier et al. ^{97,98} , 2021	http://cmpg.unibe.ch/software/fastsimcoal2/

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
RAxML-NG v1.0.2	Kozlov et al. ⁹⁹ , 2019	https://github.com/amkozlov/raxml-ng
BEAST v2.6.4	Bouckaert et al. ¹⁰⁰ , 2019	http://www.beast2.org/
ANGSD v0.935	Korneliussen et al. ¹⁰¹ , 2014	http://www.popgen.dk/angsd/index.php/ANGSD
ngsF-HMM	Vieira et al. ¹⁰² , 2016	https://github.com/fgvieira/ngsF-HMM
ggplot2 v3.4.0	Wickham ¹⁰³ , 2016	https://ggplot2.tidyverse.org/
GenoDive v3.06		http://www.patrickmeirmans.com/software/Home.html
Admixture v1.3	Alexander et al. ¹⁰⁴ , 2009	https://dalexander.github.io/admixture/
diveRsity	Keenan et al. ¹⁰⁵ , 2013	https://github.com/kkeenan02/diveRsity
prabclus	Hennig & Hausdorf ¹⁰⁶ , 2019	https://rdr.io/cran/prabclus/

RESOURCE AVAILABILITY

Lead contact

Further information can be requested via the lead contact, Adrián Talavera (adrian.talavera@csic.es).

Materials availability

This study did not generate any new reagents.

Data and code availability

All data is publicly available as of the date of publication in the following NCBI BioProjects: aCalArn1 genome assembly (PRJEB57772), ORG.one Nanopore raw data used for the assembly (PRJEB57582), Illumina data for lcWGS population genomic analyses and Nanopore and Illumina data generated outside ORG.one used for the assembly (PRJEB57773), and demultiplexed ddRADseq data (PRJNA1023478).

Any additional information required to reanalyze the data reported in this paper is available from the lead contact (adrian.talavera@csic.es) upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Material and methods

Sampling and ethics statement

The Montseny brook newt (*Calotriton arnoldi*) reference genome was sequenced from heart tissue samples of two F1 captive-bred siblings (OR192 & OR193) from the “Centre de Fauna de Torreferrussa-Generalitat de Catalunya” (Santa Perpètua de Mogoda, Spain), which were sacrificed due to their unsuitability to breed. These animals were direct descendants of founder individuals of the breeding programme (TOR16 & TOR18), from the Eastern Montseny lineage (type locality of the species). For population genomic analyses, a total of 154 tail samples were collected, including 134 *C. arnoldi* from all the known natural populations at the Montseny Massif (namely the Eastern Montseny lineage from populations A1, A2 and A3 and the Western Montseny lineage from populations B1, B2, B3, B4, B5), and 23 *C. asper* from six populations belonging to the five described mitochondrial lineages¹⁰⁷ (namely lineages O1–5; Figure 1). Tissue samples (i.e., tail tips) were stored in 99% ethanol at -20°C. Due to the species’ susceptibility to chytridiomycosis, strict biosecurity measures were taken while sampling. New nitrile gloves were worn for each animal and all tools and equipment were disinfected with Virkon™ (Zotal).

Genomic data production

High molecular weight DNA extractions were performed using MagAttract HMW DNA kit (Qiagen) to sequence the reference genome. By using 10 different flowcells, we sequenced 562.08 Gbp of Oxford Nanopore Technologies (ONT) data (63.88 M reads, read N50 > 34 kbp and 98.2% of the reads with QV>7) from OR192 and OR193 individuals. Data was basecalled with Guppy versions v3.2.10, v4.3.4 or v5.0.11, using the most recent version available at the time data was generated. On the other hand, we built an Illumina library following the BEST protocol¹⁰⁸ with minor changes and 588.3 Gbp of Illumina data (1.96x2 M 150-pb reads) was sequenced with HiSeq4000 from OR193 individual.

For lcWGS and ddRADseq libraries, DNA was extracted from tissue samples following standard high-salt protocols¹⁰⁹ and quality-checked with 1% agarose gel electrophoresis. DNA concentrations were quantified with BroadRange Qubit 2.0 fluorometer (Invitrogen). Out of the 154 DNA samples, we selected one male and female of each known inhabited brook at the Montseny, plus one representative of each *C. asper* lineage, to build 21 libraries for lcWGS, following the BEST protocol and sequencing at low coverage on HiSeq4000 with 150 pb PE reads. On the other hand, we followed the ddRADseq protocol by Peterson et al.¹¹⁰ with minor changes and double-digested 154 samples of all *C. arnoldi* and three *C. asper* populations -belonging to two lineages- with the enzymes SbfI and MspI. Digested fragments were ligated

with barcoded Illumina adapters and size-selected for 415–515 bp. Finally, ddRAD libraries were quality-checked with Bioanalyzer (Agilent) and sequenced with 75 bp SE reads using Illumina NextSeq 500.

METHOD DETAILS

De novo reference genome assembly

Illumina raw data was trimmed afterwards with Cutadapt v3.2.⁷⁶ Then, we generated a k-mer database using Meryl v1.7. A 22-mer database (in complex mode) was built using the `best_k.sh` script from Merqury v.1.1,⁷⁷ with the k-mer length decided based on the length of the axolotl genome assembly ASM291563v2²⁶ as an estimate of the genome size. Diploidy of *C. arnoldi* was assessed and confirmed with Smudgeplot v.0.2.5⁷⁸ based on another 22-mer database obtained with KMC v.3.1.1,⁷⁹ adding the option `-middle` (Figure S1). GenomeScope v2.0⁷⁸ was employed to estimate the genome assembly haploid length (22.75–28.38 Gb), the proportion of repeats (15.46–18.04 Gb) and heterozygosity with the 22-mer Meryl database (98.75–100% of homozygosity; Figure S2).

We used Filtlong v0.2.0 (available at <https://github.com/rwick/Filtlong>) to aim for the 340 Gb of the best ONT data available (aiming for 12–15X coverage), setting a minimum size of 4 kbp and looking for high-quality reads (`-minlen 4000 -min_mean_q 80`). The amount of ONT data obtained was ~12x of the maximum length calculated by GenomeScope (=28.3 Gb). The genome was assembled with Flye v2.8.3⁸⁰ adding two internal rounds of polishing, with the ONT raw reads option and scaffolding enabled by default. The assembly process required almost 80 computing days to obtain a draft assembly in a node with 48 CPUs, reaching >1Tb of maximum memory consumption. Trimmed Illumina data was mapped to the draft assembly with BWA v.0.7.15⁸¹ and used to polish it with HyPo v1.0.3.⁸² We used Minimap2 v2.17-r941⁸³ with the option `"-x map-ont"` to map the reads to the polished assembly and then run `purge_dups v1.2.5`⁸⁴ to remove haplotigs. We manually changed cutoffs based on the k-mer histogram from "5 40 120" to "5 21 150". Finally, we used the Meryl k-mer database (k=22) to run Merqury in order to evaluate the k-mer completeness and base accuracy of the final assembly, yielding 35.2 QV of base accuracy and 97.54% of k-mer completeness. Functional completeness was assessed using BUSCO v5.2.2 (82.3% BUSCO(S) vertebrata odb10, n=3,354) and contiguity (scaffold N50=1.65 Mb; 47,257 scaffolds) metrics were summarized as a snail plot (Figure S3) using BlobToolKit⁸⁵ as well as compared with other previously published assemblies in the order Caudata (Table S1) and Vertebrate Genome Project's (VGP) standards (Table S2).

Finally, we used RepeatMasker v4.1.2⁸⁶ to mask the resulting genome assembly following Nowoshilow et al.²⁶ First, we used the available Vertebrata, Tetrapoda and Amphibia repeat databases followed by the use of the repeat custom library used for the axolotl genome in Nowoshilow et al. (2018) provided by the authors. Afterwards, we ran RepeatModeler v1.0.11⁸⁷ to create a Montseny brook newt specific repeat database and ran again RepeatMasker with the resulting database.

Processing of lcWGS data

Low-coverage WGS raw data, along with the high-coverage WGS raw data from the reference genome animal (OR193), was quality-checked using FastQC v0.11.8⁸⁸ and visualized with MultiQC.¹¹¹ Data was trimmed using fastp v0.23.1,⁸⁹ allowing deduplication, requiring a minimum read length of 30 bp, allowing for the detection of adapters in PE data for adapter removal and providing the adapter sequences. We used BWA⁸¹ with BWA-MEM algorithm to map the trimmed data to the *de novo* assembled reference genome. After mapping, we marked duplicates of the re-sequenced genomes with Picard v2.8.2.¹¹² Finally, we filtered with SAMtools v1.9,⁹⁰ removing unmapped reads, secondary alignments and previously marked PCR duplicates (`-F 3332`). We also removed any alignment with mapping quality <25.

Processing of ddRADseq data

A total of 3.34×10^8 reads were obtained from the ddRAD sequencing, with a mean of 2.17×10^6 reads per sample. ddRAD sequencing output was quality-checked using FastQC and MultiQC. Sample demultiplexing, alignment, SNP-calling and some basic filtering steps were carried out with *ipyrad* v0.9.78.⁹¹ We tested clustering thresholds of 0.85, 0.89, 0.92 and 0.95 for *de novo* assembly and reference-assisted assembly with the *C. arnoldi* reference genome scaffolds, retrieving a different number of loci (Table S11). For downstream analyses, only the reference-assisted method was chosen. PhredQ score offset was kept by default at 33 and minimum samples per locus at 4. The output from *ipyrad* was later filtered in different ways to produce a number of datasets. The final number of unlinked SNPs achieved for each dataset, the performed filtering steps and the analyses in which they were used are reported on Table S12. A main dataset (*general*, Table S12) was filtered by common markers, followed by an iterative filter which discards alternatively individuals and genotypes with higher missingness than a 2%-step-decreasing threshold, from 100% missing data to 75% (script from Burriel-Carranza et al.¹¹³, following recommendations by O'Leary et al.¹¹⁴). Later, we applied a hard filter of 15% genotype missingness, removed singletons (MAC = 1), monomorphic sites and heterozygote excess (command `"-hwe 1e-50 midp keep-fewhet"` of PLINK v2.0⁹²) and kept only biallelic SNPs. Finally, we used PLINK v1.9 to prune the dataset by linkage disequilibrium, allowing a maximum correlation between SNPs of 0.5 within 50-SNP sliding windows to achieve a dataset comprising 132 individuals and 9,470 unlinked SNPs. As the sampling was carried out along several years and given the tissue regeneration skills of urodeles, a last filtering step using PLINK v2.0 was applied to detect putative pseudo-replicates. For demographic inference analyses, we removed all 1st-degree or closer relatives by using PLINK v2.0 function `-king-cutoff` and applied the same above-mentioned filtering steps to create the dataset *no_relatives*, comprising 88 individuals and 8,841 unlinked SNPs.

Another dataset (*no_outgroup*) was created directly from the *ipyrad* output selecting just the *C. arnoldi* samples which passed the post-processing filtering of the *general* dataset, and applying the same filters but allowing 30% genotype missingness; resulting in a dataset of 115 animals and 1,339 uSNPs. Also using *ipyrad*, we created a branch at step 3 to generate the dataset *loci*, allowing for 15% genotype

missingness. Finally, the seven individuals with the least missingness, representing different pure genomic clusters from Admixture (see [results](#)) were used to create the dataset *no_missingness*, which was filtered allowing no missingness and, alike the dataset *general*, keeping just unlinked biallelic SNPs.

QUANTIFICATION AND STATISTICAL ANALYSES

Demographic inference

The ddRADseq dataset *no_relatives* was downprojected by means of the *easySFS* script (<https://github.com/isaacovercast/easySFS>) into two sets of Site Frequency Spectrum (SFS) files. For the first one (*2_spp_SFS*), we split the individuals by species and chose the optimum number of retained haploid sequences without missing data based on the number of segregation points (*C. arnoldi* 124 sequences, 525 segregation points; *C. asper* 10 seq., 1,357 seg. points). The second set of SFS *-3_lineages_SFS-* files was created assuming there are 3 lineages: *C. asper*, Western *C. arnoldi* and Eastern *C. arnoldi*. 10, 70 and 60 sequences and 1,367, 245 and 208 segregation points were retrieved as the best option to downproject.

From *2_spp_SFS*, both intraspecific and unidimensional folded SFS files were used as input for Stairway Plot v2.1.1.⁹³ The number of total sites was estimated from the percentage of monomorphic and non-biallelic sites in each species. We assumed 4-year generation times and a mutational rate of 3.08×10^{-8} substitutions per site per generation.¹¹⁵ Singletons were considered only for *C. asper* analyses, as some populations retained just one individual after the downprojection. The medians of N_e estimates based on 200 runs per species were plotted for both species, along the respective 95% confidence intervals for the Montseny brook newt. Demographic inferences were coupled with reconstructions of the continental mean surface air temperature anomalies of the Northern Hemisphere (40°–80° N) and the total ice volume for the Eurasian ice sheet (in meters of sea level equivalent) for the last 200 ky (dataset from de Boer et al.,¹¹⁶ see [STAR methods](#) in de Boer et al.¹¹⁷). To account for local events, we included a Western Mediterranean Sea surface temperature reconstruction for the last 5 ky.¹¹⁸ Ultimately, some major and minor periods of climatic relevance were also highlighted. Afterward, we statistically tested for an association between *Calotriton* spp. median effective population sizes and either surface-air temperature or ice sheet volumes in Eurasia. We fitted four Generalized Least Squares (GLS) models over 10,000 permutations using a randomized residual procedure with 'RRPP' package v. 1.3.1.⁹⁵ To account for temporal autocorrelation (i.e. the non-independence of the residuals), we previously built matrices of covariance for each variable to explain the decay of autocorrelation with time by means of a corAR1 model using 'nlme' package v3.1-140⁹⁶ in R.⁹⁴ The resulting matrices were incorporated in the structure of the respective error terms of each model, but, globally, little temporal autocorrelation was found ($\phi=0$).

On the other hand, the multidimensional SFS from *3_lineages_SFS* was used for demographic simulations and parameter inference under the composite-likelihood approach of *fastsimcoal2*.⁹⁷ Four different models were designed, differing in the allowed migration matrices ([Figure S4](#)). The first model lacked migration at all, the second allowed for recent migration between *C. arnoldi* lineages, the third for ancient asymmetrical migration from *C. asper* to *C. arnoldi*; and the fourth model considered both migrations patterns. We chose not to test bidirectional migration between the two species since *C. asper* dispersal capacity is considerably higher than *C. arnoldi*'s. All models had the same topology, the resize option was permitted after whichever lineage split, and an extra instantaneous bottleneck was allowed along *C. arnoldi* history (T_{BOT} ; [Table S4](#)), so that the estimated divergence time between Montseny lineages (T_{DIV} ; [Table S4](#)) was less biased due to extreme changes in effective population changes. Eastern and Western *C. arnoldi* effective population sizes (N_A and N_B in [Table S4](#), respectively) were bounded up to 12,000 haploid sequences (i.e., 6,000 diploid individuals) - the 95%-confidence maximum for the species at present according to Stairwayplot2 results. Generation time was set to 4 years and the *C. asper*-*C. arnoldi* split was fixed 440,000 generations ago (1.76 my, as the median of the mitochondrial estimate by Valbuena-Ureña et al.;⁴ 95% HPD 1.24–2.44 my). For each scenario, 100 independent runs with different initial values were carried out and the best-fitting model was chosen based on Akaike's Information Criterion ($AIC_i = 2k_i - 2 \ln \text{MaxLhood}_i$),¹¹⁹ and the model normalized relative likelihood ($w_i = \exp(-0.5\Delta AIC_i) / \sum_r \exp(-0.5\Delta AIC_r)$) based on Excoffier et al.⁹⁸ For each run of each of the 4 models, we set 500,000 coalescent simulations, up to 50 ECM cycles in Brent optimization of parameters, with monomorphic sites masked.

Phylogenomic reconstructions

First, we constructed a Maximum Likelihood (ML) phylogeny based on the ddRADseq *loci* dataset of all individuals ([Table S12](#)) to assess the genomic monophyly of *Calotriton* spp. and their intra-specific lineages, namely Western Montseny, Eastern Montseny, Western Pyrenees and Eastern Pyrenees. To do so, we used RAXML-NG v1.0.2⁹⁹ with a GTRGAMMA model, starting from 50 random and 50 parsimony-based trees and with 1,000 bootstrap replicates ([Figure S8](#)). Then, we inferred a time-calibrated Bayesian Inference (BI) phylogeny based on SNPs under the coalescent model, which directly models incomplete lineage sorting, a common issue when dealing with divergence among young taxa.¹²⁰ We ran SNAPP (implemented on BEAST v2.6.4¹⁰⁰) on the ddRADseq dataset *no_missingness* of seven individuals with the least missingness, which were treated as belonging to one of the four monophyletic lineages (2 Western Montseny; 3 Eastern Montseny, 1 Eastern Pyrenees and 1 Western Pyrenees). The vcf file comprising 8,139 unlinked SNPs was converted to xml format following Stange et al.¹²¹ by using the script *snapp_prep.rb* (available at: https://github.com/mmatschiner/snapp_prep). *Calotriton arnoldi* and *C. asper* were constrained to be monophyletic and the divergence between them was set with a lognormal distribution in 1.76 ± 0.2 mya following a previously published calibration.⁴ Analyses consisted of three independent runs with 6 million generations each. Tracer v1.7.2¹²² was used to check for stationarity (estimated sample sizes of all parameters ≥ 200) and convergence between runs. Converging log and tree files were combined using LogCombiner v1.10 with 10% burn-in. The cloudogram was visualized with DensiTree v. 2.6.4,¹²³ and TreeAnnotator v2.6.3 was used to construct the 'Maximum clade credibility' tree and annotate nodes with the corresponding posterior probabilities. We plotted together the cloudogram, plus the consensus tree and the posterior probabilities of each node along with the mean divergence times and their

95% confidence intervals. Furthermore, important climatic events of each time frame were highlighted, specifically the glaciations of Donau (c. 1.8-1.0 mya), Riss (c. 300-130 kya) and Würm (c. 115-11.7 kya), and the interglacial period separating the latter two (130-115 kya).

Population genomic analyses and ROHs with lcWGS

We estimated the heterozygosity of the 22 *Calotriton* spp. individuals with WGS data (see Table S6 for details) using ANGSD v0.935¹⁰¹ on the downsampled (1.3x coverage) filtered bam files as follows: “*angsd -i \$file.bam -anc aCalArn_1.scaffolds.fa -dosaf 1 -gl 1*”. Furthermore we applied additional filters (*-remove_bads 1 -baq 1 -C 50 -minQ 20 -minMapQ 30*) and calculated heterozygosity values for the largest scaffold and genome-wide, including and excluding repetitive regions, for each individual following the ANGSD manual.

ANGSD was used as well to call genotype likelihoods to perform Principal Component Analyses (PCAs) in order to explore population structure, and to estimate runs of homozygosity (ROHs, or identity-by-descent tracks) from all WGS samples, including the reference sample from the breeding center (an F1 Eastern-lineage individual). We calculated genotype likelihoods for five different datasets: *all_wild&captivity* (22 samples), *all_wild* (21 samples), *only_arnoldi* (16 samples), *only_west* (10 samples), *only_east* (6 samples); varying the number of samples and the *minInd* option (*minInd* = 18 for *all_wild&captivity* and *all_wild*; *minInd* = 15 for *only_arnoldi*, and *minInd* = (number of individuals – 1) for *only_west* and *only_east*). We inferred the minor and major alleles from likelihoods (*-doMajorMinor 1*), avoiding sites where the reference does not contain either the major or the minor alleles inferred (*-skipTriallelic 1*). Genotype likelihoods were reported as a beagle file (*-doGlf2*) and the allele frequency estimations were performed assuming the major allele inferred is known (*-doMaf2*). We removed sites with minor allele frequency lower than 0.04 (*-minMaf 0.04*) and sites with a p-value greater than $1e-6$ (*-SNP_pval 1e-6*). Furthermore, different quality filters were applied to the imputed BAM files: we excluded non-primary alignments, and unmapped and duplicate reads (*-remove_bads 1*), discarded reads with multiple mappings (*-uniqueOnly 1*), kept only pairs of reads for which both were properly mapped (*-only_proper_pairs 1*), performed Base Alignment Quality calculation (*-baq 1*), avoided trimming (*-trim 0*), adjusted base quality for excessive mismatches (*-C 50*) and set a minimum mapping quality (*-minMapQ 25*). PCAs were performed using the output beagle files with PCAngsd v0.8. A total of 23.50M SNPs for *all_wild*, 2.08M SNPs for *only_arnoldi*, 1.55M for *only_west* and 0.75M for *only_east* datasets were used.

Finally, we used ngsF-HMM¹⁰² to estimate runs of homozygosity (ROHs, or identity-by-descent tracks) from the resulting beagle file for *all_wild&captivity*. ngsF-HMM uses an iterative approach that stops when a dual condition threshold is reached.¹⁰² We followed the authors' recommendation of lowering this threshold to “*-min_epsilon 1e-9*”. Only the 50 largest scaffolds from our assembly (with lengths 11.6-7.1 Mb) were selected for this analysis. We compared the ROH counts between lineages (i.e. Western Montseny, Eastern Montseny and Pyrenees) at four different ranges in terms of ROH length (0.5-1, 1-2, 2-3 and >3 Mb) by one-way ANOVA tests and corresponding Bonferroni-adjusted post-hoc tests. Means and SD of ROH counts along those 50 largest scaffolds by lineage and by categories of ROH length were plotted in barplots with ggplot2 v3.4.0.¹⁰³

Population genomic analyses with ddRADseq

GenoDive v3.06¹²⁴ was used to estimate observed (H_o) and expected heterozygosity (H_s) and inbreeding coefficients (G_{IS}) per population from the *general* dataset. The same software was also used for calculating the fixation indices, both lineage- and population-pairwise. Observed and expected heterozygosities per population from ddRAD data were plotted as a barplot and all the population-pairwise indices were plotted on a heatmap with ggplot2. To explore population structure, we performed two PCAs by means of PLINK v1.9, one at genus level (dataset *general*) and another at *C. arnoldi* species level (dataset *no_outgroup*), and then plotted results with ggplot2. Furthermore, to assess structure and admixture between populations, we run Admixture v1.3,¹⁰⁴ again hierarchically, at genus- and species-level, for K values from 1 to the number of sampling localities (i.e., twelve and nine at genus and species level, respectively) and calculated 20 cross-validation error (CV) values with 20 replicates. Optimal K values were chosen based on CV scores (see Table S10), the plateau profile of those values and the likelihood of the results in biological terms. Additionally, geneflow among *C. arnoldi* populations was estimated with the function *divMigrate* in the *diveRsim* package¹⁰⁵ in R to obtain a relative migration network graph from the dataset *no_outgroup*. Geneflow, scaled to the largest magnitude estimated, was measured in N_m (i.e. effective number of migrants^{125,126}) and the significance of asymmetric gene flow was tested with 1,000 bootstrap iterations and $\alpha = 0.05$. Finally, the resulting matrix was plotted with the R package *qgraph*,¹²⁷ and edges below 0.35 were disregarded (Figure 5B). Geneflow arrows which are significantly asymmetric are colored in these graphs, whereas the rest remain in gray scale depending on their intensity. Since the Montseny brook newt strictly disperses by water-way routes, migration networks were depicted with a scheme of brook connectivity within the Tordera river basin.

To explore the differences in population structure between Western and Eastern Montseny lineages, we tested for isolation-by-distance (IBD) in *C. arnoldi* populations following Hausdorf & Hennig¹²⁸ These IBD tests check if genetic distances between individuals or, in our case, populations, belonging to two different lineages (Eastern and Western Montseny as candidate species *sensu* Hausdorf & Hennig¹²⁸ or primary species hypotheses *sensu* Bamberger et al.¹²⁹) are not larger than expected based on their geographical distances and within-group IBD patterns (null hypothesis *sensu* Hausdorf & Hennig¹²⁸ or secondary species hypothesis *sensu* Bamberger et al.¹²⁹). Genetic distances were measured as $F'_{ST}/(1-F'_{ST})$ ¹³⁰ based on previous results. We measured shortest water-way linear distances between populations in meters and then log-transformed values. We used R package *prabclus*¹⁰⁶ to perform a Jackknife-based test for equality of two independent within-lineage regressions (i.e. within the Western and the Eastern Montseny lineages) to confirm that IBD evolves alike within both metapopulations, and thus that they can be merged into a single joint within-lineages regression (intercept $p=0.802$; slope $p=0.745$). Second, another Jackknife-based test was run to check for the equality of the latter and another regression over all comparisons, both within- and between Montseny lineages. The non-equality of both regressions would imply that the observed genetic differences cannot be explained by IBD alone.