

Alpine endemic spiders shed light on the origin and evolution of subterranean species

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ABSTRACT

We designed a comparative study to unravel the phylogeography of two Alpine endemic spiders characterized by a different degree of adaptation to subterranean life: *Troglohyphantes vignai* (Araneae, Linyphiidae) and *Pimoa rupicola* (Araneae, Pimoidae), the latter showing minor adaptation to hypogean life. We sampled populations of the model species in caves and other subterranean habitats across their known geographical range in the Western Alps. By combining phylogeographic inferences and Ecological Niche Modeling techniques, we inferred the biogeographic scenario that led to the present day population structure of the two species. According to our divergent time estimates and relative uncertainties, the isolation of *T. vignai* and *P. rupicola* from their northern sister groups was tracked back to Middle–Late Miocene. Furthermore, the fingerprint left by Pleistocene glaciations on the population structure revealed by the genetic data, led to the hypothesis that a progressive adaptation to subterranean habitats occurred in *T. vignai*, followed by strong population isolation. On the other hand, *P. rupicola* underwent a remarkable genetic bottleneck during the Pleistocene glaciations, that shaped its present population structure. It seems likely that such shallow population structure is both the result of the minor degree of specialization to hypogean life and the higher dispersal ability characterizing this species. The simultaneous study of overlapping spider species showing different levels of adaptation to hypogean life, disclosed a new way to clarify patterns of biological diversification and to understand the effects of past climatic shift on the subterranean biodiversity.

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INTRODUCTION

Long term climatic changes are often invoked among the most important factors that drove surface-dwelling invertebrate populations to colonize subterranean habitats, causing their isolation and shaping their present day distribution patterns (*Jeannel, 1943; Peck, 1980; Holsinger, 1988; Botosaneanu & Holsinger, 1991; Culver & Pipan, 2010*). In this regard, the Pleistocene glaciations (reviewed in *Culver & Pipan, 2010*) and the

Messinian Salinity Crisis ([Faille et al., 2010](#)) have been pointed out among the main drivers of the present distribution patterns of the European subterranean biodiversity. The climate-driven isolation caused the allopatric divergence of subterranean populations, resulting in narrow patterns of distribution and high levels of endemism ([Christman et al., 2005](#); [Culver & Pipan, 2009](#); [Borges et al., 2012](#); [Cardoso, 2012](#)). Accordingly, population studies conducted so far have uncovered an extreme genetic structuring in terrestrial invertebrates in subterranean habitats ([Kane, Barr & Badaracca, 1992](#); [Allegrucci, Minasi & Sbordoni, 1997](#); [Hedin, 1997](#); [Gentile & Sbordoni, 1998](#); [Hedin & Thomas, 2010](#); [Ribera et al., 2010](#); [Snowman, Zigler & Hedin, 2010](#); [Dixon & Zigler, 2011](#); [Zhang & Li, 2013](#); see also [Bohonak, 1999](#) for an historical perspective on this topic). This general trend was mainly interpreted in light of the past climatic transition, as the result of the limited dispersal ability of subterranean organisms. Because of the adaptation to the hypogean medium, subterranean species develop narrower physiological tolerance (i.e., troglomorphism *sensu* [Christiansen, 1962](#)), which hamper their dispersal ability through non-subterranean habitats. For example, some cave-dwelling spiders with minor adaptations to hypogean life are known to disperse outside caves in different stages of their life cycle (e.g., *Meta* spiders; [Smithers, 2005](#); [Mammola & Isaia, 2014](#)). On the other hand, subterranean habitats, especially caves, are generally connected through a networks of small cracks and voids which may facilitate dispersal of the invertebrate fauna ([Juberthie, Delay & Bouillon, 1980](#); [Juberthie, Delay & Bouillon, 1981](#); [Uéno, 1987](#); [Romero, 2012](#); [Culver & Pipan, 2009](#); [Culver & Pipan, 2014](#); [Giachino & Vailati, 2010](#) among others).

Here we designed a comparative study aimed at providing insights on the origin and the evolution of the hypogean biodiversity. Specifically, we focused on two Alpine endemic species co-occurring in caves and other subterranean habitats across most of their known distribution range and exhibiting different levels of adaptation to subterranean life.

The first model species, *Troglohyphantes vignai* [Brignoli, 1971](#) (Araneae, Linyphiidae), is endemic to the Western Italian Alps (NW Italy), being discontinuously distributed from the Cottian (Province of Torino) to the Maritime Alps (Province of Cuneo) ([Isaia et al., 2011](#)). Since the description, all available records of this species refer to cave habitats ([Brignoli, 1971](#); [Brignoli, 1985](#); [Pesarini, 2001](#); [Arnó & Lana, 2005](#); [Isaia & Pantini, 2010](#); [Isaia et al., 2011](#)). *T. vignai* shows adaptations to the hypogean life, namely loss of pigmentation, reduction of the eye apparatus, thinning of integuments and heavy spination. *T. vignai* was described from the cave *Buco di Valenza* (Speleological cadastral number: 1009 Pi/CN; Po Valley) by [Brignoli \(1971\)](#). In the same publication, Brignoli also described *T. rupicapra*, which was distinguished from *T. vignai* by small morphological details of the epigynum. *T. rupicapra* was described on material from *Grotta superiore delle Camoscere* (Speleological cadastral number: 250 Pi/CN; Pesio Valley). According to the species description, and as later observed by [Isaia & Pantini \(2010, Figs. 15–18\)](#), *T. rupicapra* shows a higher degree of troglomorphism compared to *T. vignai*, namely higher depigmentation, reduction of eye diameter and loss of functional eyes, and lowering of the profile of the cephalothorax. The species validity of *T. rupicapra* was questioned by [Pesarini \(2001\)](#), who proposed the synonymy *T. rupicapra* = *T. vignai*, currently accepted in the [World Spider Catalog \(2015\)](#).

Our second model organism, *Pimoa rupicola* (Simon, 1884) (Araneae, Pimoidae), is an Alpine-Apenninic endemic element, recorded almost continuously from the Graian Alps to the Tuscan Apennines (Thaler, 1976; Hormiga, 1994; Isaia et al., 2011) and French Maritime Alps. Several authors (Brignoli, 1971; Brignoli, 1972; Brignoli, 1985; Thaler, 1976; Arnó & Lana, 2005; Isaia et al., 2011) referred to *P. rupicola* as a troglophile species (*sensu* Sket, 2008), being abundant in subterranean habitats and occasionally recorded from surface habitats, such as leaf litter, humid rocks covered by mosses and mountain screes (Bertkau, 1890; Jackson, 1926; Thaler, 1976; Hormiga, 1994; Isaia et al., 2015; Isaia, Paschetta & Chiarle, 2015). Given the sporadic collection of individuals outside cave (mainly pitfall trap data reported in Isaia et al. (2015); Isaia, Paschetta & Chiarle (2015), and additional unpublished data collected by two of us (SM and MI)), it seems likely that males and immatures of *P. rupicola* disperse through the epigeal environment. Morphologically, the species does not show any remarkable troglomorphic features: it has eight functional eyes, it is entirely pigmented and it has a well defined abdominal pattern.

To present knowledge (Arnó & Lana, 2005; Isaia et al., 2011), the Alpine range of *P. rupicola* encompasses the entire range of *T. vignai* and the two species often co-occur in the same cave.

In this contribution we investigated the biogeographic events that shaped present day population structure of the two species. Since the study was set at the species/population interface, we employed two fast-evolving DNA markers, namely the mitochondrial cytochrome oxidase I (*cox1*) and the nuclear second internal transcribed spacer region (*ITS-2*). The popularity of these markers stems from a generally high level of variation, which permits to reconstruct relationships among and within spider species, making them particularly suitable for population and biogeographic studies (Agnarsson, 2010; Videgar, Toplak & Kuntner, 2014).

Moreover, in accordance with Peterson (2009) we coupled genetic inferences with ecological niche modeling techniques, thus obtaining multiple supports to our research hypothesis. In particular, we hypothesized that past climatic changes played a key role in shaping the genetic structure of the populations of the two species. Given the contrasting degree of subterranean specialization exhibited by the two spiders, we further hypothesized that populations of *P. rupicola* show minor genetic structure than *T. vignai*.

Additionally, this study offered the opportunity to reveal the existence of cryptic species within the two lineages. In fact, it was observed during the course of the study that individuals belonging to the northern populations of *P. rupicola* presented subtle but consistent differences in their genital morphology. Hereinafter, we will restrict the use of the epithet '*rupicola*' to indicate the southern populations, and we will refer tentatively to the northern populations as 'n. sp.'

MATERIAL & METHODS

Sampling

Populations were collected in caves, abandoned mines and other hypogean habitats across the known distribution range of *Troglohyphantes vignai* and *Pimoa rupicola* in

the Western Alps. The distribution range of *T. vignai* was entirely covered (including type locality and former localities of *T. rupicapra*, indicated hereinafter as *T. vignai sensu rupicapra*), while for *Pimoa* we only sampled Alpine populations, thus excluding French and Apenninic populations. The complete list of localities is reported in Table 1 and Fig. 1. The toponomastics and classification of the different sectors and sub-sectors of the Alps follows the partition of the Alpine chain (SOIUSA: Marazzi, 2005). Specimens were hand-collected, preserved in 95% ethanol and stored in freezer. Given that the sampled environments were highly oligotrophic, in certain localities we were able to detect and collect only few individuals of the two investigated species. The number of individuals collected for each locality ranged from 2 to 8 in *P. rupicola* and *P. n. sp.* and from 1 to 7 in *T. vignai* (Table 2). Overall, 119 *Pimoa* specimens from 25 localities and 37 *Troglohyphantes* specimens from 8 localities were used in this study.

DNA extraction, amplification and sequencing

One leg was removed from each specimen for DNA extraction. Whole genomic DNA was extracted from the samples using the SpeedTools Tissue Extraction Kit (Biotools) following the manufacturer's protocol. A 676 bp region of the mitochondrial cytochrome oxidase subunit I (*cox1*) gene and a 400 bp region of the nuclear second internal transcribed spacer region (*ITS-2*) gene were amplified using polymerase chain reaction (PCR). We utilized the primers C1-J-1490 (5'-GGTCAACAAATCATAAAGATATTGG-3'; Folmer et al., 1994) and C1-N-2191 (5'-CCCGGTAAAATTTAAATATAAACTTC-3'; Simon et al., 1994) for the *cox1* and the ITS-5.8s (5'-GGGACGATGAAGAACGGAGC-3') and the ITS-28s (5'-TCCTCCGCTTATTGATATGC-3') for the *ITS-2* (White et al., 1990).

PCR amplifications were carried out in 25 μ L reaction volume in a final concentration of 0.1 μ L *Taq* polymerase (Promega), 5 μ L buffer (Promega), 2.25 μ L MgCl₂ (Promega), 0.2 mm of each dNTP, 0.5 μ L of each primer and 1.5 μ L of DNA sample. PCR conditions for amplification were as follows: initial denaturing step at 95 °C for 5 min, 35 amplification cycles (94 °C for 30 s, 45 °C for 35 s, 72 °C for 45 s *cox1* fragment and 94 °C for 45 s, 48 °C for 1 min, 72 °C for 60 s for *ITS-2* fragment) followed by a final extension at 72 °C for 5 min. For certain populations of *Pimoa rupicola* (localities #23 and #26), a slightly different annealing protocol for the *cox1* was utilized (94 °C for 30 s, 42 °C for 35 s, 72 °C for 45 s). PCR products were visualized on agarose gels.

PCR product were cycle-sequenced at Macrogen, Inc. (Seoul, Korea; <http://www.macrogen.com>). The DNA sequences obtained were preliminary assembled and edited using Geneious 7.1 (Kearse et al., 2012; <http://www.geneious.com>).

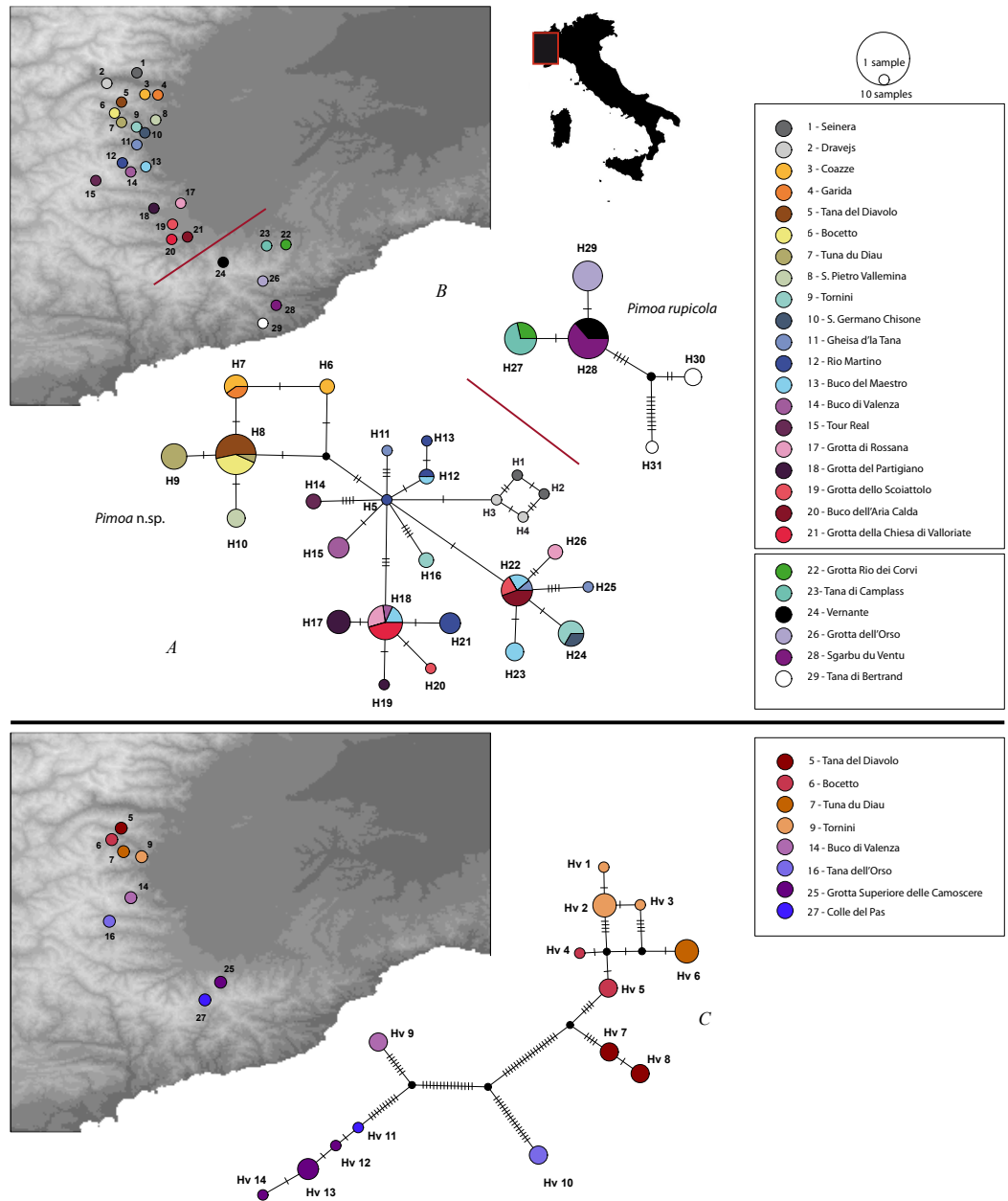
The alignment of the *cox1* sequences was trivial, as they showed no evidence of indel mutations. The *ITS-2* fragments were aligned with the online version of MAFFT (Katoh & Toh, 2008; <http://mafft.cbrc.jp>), using the Q-INS-I strategy with default options. We explored best partitioning schemes and substitution models simultaneously using PartitionFinder v.1.0.1 (Lanfear et al., 2012) under a Bayesian information criterion (BIC).

Table 1 Summary of the sampled localities. Sampled localities of *Pimoa rupicola*, *P. n. sp.* and *Troglohyphantes vignai* ordered by latitude (from North to South).

Cod	Valley	Locality	Habitat type	x	y	P. n.sp	P. rupicola	T. vignai	Date	Collector/s
1	Susa	(!) Seinerà	Abandoned mine	7,201	45,163	*			20.II.2011	Mammola S., Piano E., Giuliano D.
2	Susa	(!) Dravejs	Scree	7,039	45,118	*			13.VI.2014	Mammola S., Piano E.
3	Sangonetto	Coazze	Ruined building	7,241	45,067	*			20.II.2011	Isaia M.
4	Sangonetto	Garida	Abandoned mine	7,304	45,055	*			20.II.2011	Isaia M.
5	Chisone	[1591 Pi/TO] Tana del Diavolo	Wild cave	7,123	45,028	*		*	12.IX.2014	Isaia M., Mammola S.
6	Chisone	Bocetto	Abandoned mine	7,086	44,959	*			12.IX.2014	Isaia M., Mammola S.
7	Germanasca	[n.c. Pi/CN] Tuna du Diau	Wild cave	7,104	44,949	*		*	12.IX.2014	Isaia M., Mammola S.
8	Lemina	S. Pietro Val Lemina	Abandoned mine	7,297	44,937	*			12.IX.2014	Isaia M., Mammola S.
9	Germanasca	(!) Tornini	Abandoned mine	7,199	44,908	*		*	12.IX.2014	Isaia M., Mammola S.
10	Germanasca	S. Germano Chisone	Abandoned mine	7,225	44,901	*			13.XI.2014	Isaia M.
11	Pellice	[1538 Pi/TO] Gheisa d'la Tana	Wild cave	7,224	44,851	*			28.IX.2014	Isaia M., Mammola S., Paschetta M.
12	Po	Balma di Rio Martino (Opera 372)	Military bunker	7,140	44,702	*			13.XI.2014	Isaia M., Mammola S., Paschetta M.
13	Po	[1148 Pi/CN] Buco del Maestro	Wild cave	7,238	44,686	*			3.X.2014	Isaia M., Mammola S., Paschetta M.
14	Po	[1009 Pi/CN] Buco di Valenza	Wild cave	7,172	44,683	*		*	13.XI.2014	Isaia M., Mammola S., Paschetta M.
15	Varaita	(!) Tour Real	Blockhouse	6,982	44,645	*			29.VII.2014	Mammola S.
16	Varaita	[1019 Pi/CN] Tana dell'Orso di Casteldelfino	Wild cave	7,099	44,561	*		*	21.VII.2013	Mammola S.
17	Varaita	[1010 Pi/CN] Grotta di Rossana	Wild cave	7,431	44,534	*			20.VII.2013	Giresi A., Mammola S.
18	Maira	[n.c. Pi/CN] Grotta del Partigiano di Roccabruna	Wild cave	7,294	44,509	*			14.VII.2014	Isaia M.
19	Stura	[1122 Pi/CN] Grotta dello Scoiattolo	Wild cave	7,389	44,412	*			13.I.2015	Isaia M., Mammola S., Paschetta M.
20	Stura	[1102 Pi/CN] Buco dell' Aria Calda	Wild cave	7,462	44,349	*			03.X.2014	Isaia M., Mammola S., Paschetta M.
21	Stura	[1056 Pi/CN] Grotta della Chiesa di Valloriate	Wild cave	7,382	44,339	*			13.I.2015	Isaia M., Mammola S., Paschetta M.
22	Lisio	[884 Pi/CN] Grotta Rio dei Corvi	Wild cave	7,994	44,303	*			26.XII.2014	Isaia M., Mammola S.
23	Corsaglia	[113 Pi/CN] Tana di Camplass	Wild cave	7,887	44,297		*		26.XII.2014	Isaia M., Mammola S.
24	Vermenagna	Fort (B) of Vernante (Opera 14)	Military bunker	7,529	44,257		*		13.I.2015	Isaia M., Mammola S., Paschetta M.
25	Pesio	[250 Pi/CN] Grotta superiore delle Camoscere	Wild cave	- Protected data -	44,21719			*	26.XII.2014	Isaia M., Mammola S.
26	Tanaro	[118 Pi/CN] Grotta dell'Orso di Ponte di Nava	Wild cave	7,866	44,119		*		10.X.2014	Isaia M., Mammola S.
27	Pesio	(!) Unknown cave near Colle del Pas	Wild cave	7,774	44,166			*	20.VIII.2014	Badino G.
28	Argentina	[619 Li/IM] Sgarbu du ventu	Wild cave	7,937	44,002		*		27.XII.2014	Isaia M., Mammola S.
29	Argentina	[104 Li/IM] Tana di Bertrand	Wild cave	7,867	43,916		*		27.XII.2014	Isaia M., Mammola S.

Notes.

Cod, locality numeric code used in the analysis and figures. For each record we report the name of the locality, the name of the Alpine valley, the habitat type, the geographical coordinates (longitude and latitude in decimal degrees, WGS 84 reference system), the date and the collectors. For hypogean localities, we report the Speleological cadastral number in square brackets (e.g., 1591 Pi/TO), when available. An exclamation mark in parenthesis (!) before the name of the locality indicates new unpublished records found during this study.



Genetic analyses

Population structure

Standard genetic diversity indices (nucleotide and haplotype diversity, A-T bias, transition/transversion rate) were estimated with the PopGenome package (Pfeifer et al., 2014) in R environment (R Development Core Team, 2013). We tested for population

Table 2 Standard genetic diversity indices. Diversity measures for the *cox1* and *ITS-2* genes for the localities of *Pimoa* n. sp., *P. rupicola* and *Troglohyphantes vignai* sampled in this study.

Locality code	Species	<i>Cox1</i>				<i>ITS-2</i>			
		<i>N</i>	<i>H</i>	π	<i>h</i>	<i>N</i>	<i>H</i>	π	<i>h</i>
1	<i>P. n. sp.</i>	2	2	1,000	1,000	2	1	0,000	0,000
2	<i>P. n. sp.</i>	2	2	1,000	1,000	2	1	0,000	0,000
3	<i>P. n. sp.</i>	5	2	0,600	0,600	5	2	0,400	0,400
4	<i>P. n. sp.</i>	4	1	0,000	0,000	2	1	0,000	0,000
5	<i>P. n. sp.</i>	8	1	0,000	0,000	6	2	0,476	0,476
6	<i>P. n. sp.</i>	6	1	0,000	0,000	6	1	0,000	0,000
7	<i>P. n. sp.</i>	7	2	0,285	0,285	7	1	0,000	0,000
8	<i>P. n. sp.</i>	3	3	0,000	0,000	3	2	0,666	0,666
9	<i>P. n. sp.</i>	6	3	2,133	0,533	6	3	1,400	0,600
10	<i>P. n. sp.</i>	2	1	0,000	0,000	2	1	0,000	0,000
11	<i>P. n. sp.</i>	4	4	5,666	1,000	5	3	1,333	0,666
12	<i>P. n. sp.</i>	7	4	2,476	0,714	8	3	0,678	0,464
13	<i>P. n. sp.</i>	8	5	2,429	0,857	8	3	1,047	0,523
14	<i>P. n. sp.</i>	5	2	1,500	0,500	5	2	0,333	0,333
15	<i>P. n. sp.</i>	2	1	0,000	0,000	2	1	0,000	0,000
17	<i>P. n. sp.</i>	5	2	2,500	0,500	4	1	0,000	0,000
18	<i>P. n. sp.</i>	5	2	0,666	0,333	5	1	0,000	0,000
19	<i>P. n. sp.</i>	3	3	2,666	0,666	3	2	0,000	0,000
20	<i>P. n. sp.</i>	4	1	0,000	0,000	4	2	0,500	0,500
21	<i>P. n. sp.</i>	5	1	0,000	0,000	5	1	0,000	0,000
22	<i>P. rupicola</i>	2	1	0,000	0,000	2	1	0,000	0,000
23	<i>P. rupicola</i>	5	1	0,000	0,000	5	1	0,000	0,000
24	<i>P. rupicola</i>	4	1	0,000	0,000	5	1	0,000	0,000
26	<i>P. rupicola</i>	6	1	0,000	0,000	6	1	0,000	0,000
28	<i>P. rupicola</i>	7	1	0,000	0,000	7	5	2,285	0,857
29	<i>P. rupicola</i>	3	2	7,333	0,667	3	1	0,000	0,000
5	<i>T. vignai</i>	6	2	0,612	0,600	6	1	0,000	0,000
6	<i>T. vignai</i>	6	2	1,000	0,500	6	3	1,166	0,833
7	<i>T. vignai</i>	5	1	0,000	0,000	5	1	0,000	0,000
9	<i>T. vignai</i>	7	3	0,571	0,523	7	1	0,000	0,000
14	<i>T. vignai</i>	3	1	0,000	0,000	3	1	0,000	0,000
16	<i>T. vignai</i>	3	1	0,000	0,000	3	1	0,000	0,000
25	<i>T. vignai</i> *	6	3	0,666	0,600	6	1	0,000	0,000
27	<i>T. vignai</i> *	1	1	0,000	0,000	1	1	0,000	0,000

Notes.

N, number of individuals; *H*, number of haplotypes; π , nucleotide diversity; *h*, haplotype diversity.

* Indicates populations of *T. vignai* sensu *rupicapra*.

structure among localities in the *cox1* dataset using F_{ST} as implemented in ARLEQUIN 3.01 (Excoffier, Laval & Schneider, 2005). Significance was assessed by performing 10,000 permutations. We excluded from this analysis six localities (#1, #2, #10, #15, #22 and #27) where the sampling representativeness was questionable (i.e., less than 3 individuals).

Haplotype networks were constructed using the statistical parsimony method ([Templeton, Crandall & Sing, 1992](#); [Clement et al., 2002](#)) with a confidence limit of 95% as implemented in PopArt (online at: <http://popart.otago.ac.nz>).

Phylogenetic inference

Maximum likelihood (ML) and Bayesian inference (BI) were used to infer the gene trees and the concatenated tree for each genus. For this analysis we only included unique haplotypes. We concatenated *Cox1* and *ITS-2* gene fragments in Geneious and excluded taxa with partial sequences. Gaps in the *ITS-2* were recoded as absence/presence characters using the simple method proposed by [Simmons & Ochoterena \(2000\)](#) with the help of the computer program SeqState 1.4.1 ([Müller, 2005](#)). We used *Troglohyphantes nigraerosae* and *Pimoa edenticulata* as outgroups to root the respective trees based on the results of ongoing analyses on the two genera (MA Arnedo, 2015, unpublished data; G Hormiga, 2015, unpublished data).

ML analyses were performed in RAxML v.7.4.2 ([Stamatakis, 2006](#)) with the aid of the graphical interface RAXML-GUI v.1.3 ([Silvestro & Michalak, 2011](#)), by conducting 10 runs per 500 bootstrap replicates.

BI analyses were conducted in MrBayes v.3.2 ([Ronquist et al., 2012](#)) with two independent runs of 20 million generations with four Markov chains (one cold, three heated), sampling every 1,000 generations. The convergence of chains was checked in Tracer v.1.6 ([Rambaut et al., 2014](#)) until effective sample sizes (EES) was above 200, and the average standard deviation of split frequencies (ASDSF) of the two runs was below 0.02. The first 20% of trees in each run were discarded as burn-in. The majority-rule consensus tree was generated from remaining trees.

Divergence time estimation

Divergence time was estimated for the two lineages using a multispecies coalescent approach ([Heled & Drummond, 2010](#)), as implemented in BEAST ([Drummond et al., 2012](#)). Coalescent groups within each species were first identified by using the General Mixed Yule Coalescence (GMYC; [Fujisawa & Barraclough, 2013](#)) method and used as a proxy of species in the multicoalescent analyses. GMYC is a clustering method that provides an objective way to delimit putative independent evolutionary lineages (i.e., coalescent groups). For each *cox1* alignment, we generated a ML tree (see analytical protocol above) and we converted it to an ultrametric tree with the help of PATHd8 ([Britton et al., 2007](#)). The GMYC analysis was conducted *via* the package splits ([Ezard, Fujisawa & Barraclough, 2014](#)) in R, after removing zero-length branches and make the tree fully dichotomous.

For estimating the divergence time of *Troglohyphantes* and *Pimoa* lineages, we utilized the best gene partition schemes estimated with PartitionFinder. Because of the lack of reliable calibration points (e.g., fossils, relevant geological or biogeographical events) for any of the two lineages, we relied on informed priors of the substitution rates of the *cox1*, based on available information for spiders ([Bidegaray-Batista & Arnedo, 2011](#)). Preliminary analyses using a lognormal relaxed clock for the *cox1* gene showed that the posterior

distribution of the *ucl.d.mean* parameter accreted to zero and hence a strict clock was preferred. We set the prior rate parameter of the *cox1* strict clock to a normal distribution with mean $\pm sd = 0.02 \pm 0.006$. Similarly, we assigned a strict clock prior to the *ITS-2* partition. To speed up calculation, we defined a flat prior to the *ITS-2* mean rate parameter consisting in a uniform distribution with upper and lower bounds of 0.2 and 0.0001, respectively. We selected a Yule model for the tree prior.

For each species we ran three independent MCMC chains for 50 million generations, sampling every 10,000 generations. Convergence of the three chains and correct mixing was assessed in Tracer v.1.6 (Rambaut et al., 2014).

Ecological niche modeling

We relied on ecological niche modeling (ENM; see, e.g., Elith et al., 2006) to model the ancestral distribution of the target species. Detailed methodological protocol is provided in Supplemental Information 1. In a first step, we collected all records of the target species available in the literature. We managed to track down 22 localities for *Troglohyphantes vignai*, most of which clustered together. On the other hand, for *Pimoa* we recovered 110 localities (61 for *Pimoa* n. sp. and 49 for *P. rupicola*), including new unpublished records discovered during the present study. Given the low number of localities for *T. vignai*, we only inferred the ENM model for the *Pimoa* lineages. The dataset was corrected for potential spatial autocorrelation and haphazard sampling (Oliveira et al., 2014 and references therein). We obtained present day climatic data (19 'Bioclim variables', see Table 1 of Supplemental Information 1) and altitude a.s.l. from the WorldClim website (www.worldclim.org). We obtained downscaled and calibrated Paleoclimatic data for the Last Glacial Maximum (~22,000 years ago; hereinafter LGM) from three different simulations available from Global Climate Models (GCMs; Coupled Model Intercomparison Project phase 5; <http://cmip-pcmdi.llnl.gov/cmip5>). The climatic preferences of the two species were investigated via Principal Component Analysis (PCA) in the Vegan R package (Oksanen et al., 2013). We investigated collinearity among covariates and obtained a final set of uncorrelated variables (Annual mean temperature (Bio1), Temperature annual range (Bio7) and Mean temperature of the driest quarter (Bio9)).

We generated presence-only models with the Maximum Entropy Distribution Models available in MaxEnt (Phillips, Anderson & Schapire, 2006), as implemented in the dismo R package (Hijmans et al., 2014). Firstly, we computed the models on the present climate and on the occurrence points within the M area (*sensu* Barve et al., 2011; details in Supplemental Information 1). To generate the prediction, we ran each niche model 20 times using a loop script in R, keeping in all cases a random partition of 20% of the occurrence points, which was used to evaluate model performance via the Area Under the Curve (AUC) of the Receiver Operating Characteristic (ROC) plot (Fielding & Bell, 1997). We projected the MaxEnt models in the LGM climate, under each of the three GCMs climatic scenarios. We used a conservative approach to identify potential Pleistocene refugia: we first applied a threshold of 0.6 to the continuous probability surface of presence estimated after the projection. We then combined the three projections, by sub-sampling only those pixel classified as potentially occupied ($p > 0.6$) in each LGM forecast.

RESULTS

Population structure

The new sequences obtained in the present study are available in GenBank ([Supplemental Information 2](#)). A fragment of the mitochondrial *cox1* gene of 676 bp was obtained for 37 specimens of *Troglohyphantes vignai* in 8 localities, corresponding to 14 unique haplotypes. The *cox1* data set had 79 segregating sites and 9 parsimony informative sites. The overall mean distance (*p*-distance among haplotypes) was 0.0495 ± 0.0059 . Sequences of the nuclear intron *ITS-2* were obtained from the same individuals. The alignment was 400 positions long, 10 additional absence/presence gap characters were scored, corresponding to 10 *ITS-2* sequence types. The *ITS-2* had 16 segregating sites and 9 parsimony informative sites. The overall *p*-distance among caves was 0.0389 ± 0.0020 . We obtained 676 bp *cox1* sequence fragments of 119 *Pimoa* individuals from 25 localities. The 93 individuals from 19 localities of *Pimoa* n. sp. yielded 43 haplotypes (35 segregating sites and 7 parsimony informative sites) and the 27 individuals from 6 localities of *P. rupicola* yielded 7 haplotypes. The average *p*-distance within populations in *Pimoa* n. sp. and *P. rupicola* was 0.0076 ± 0.0017 and 0.0052 ± 0.0017 , respectively, and the maximum *p*-distance between the two lineages was 0.1164 ± 0.0111 . The nuclear *ITS-2* sequences were obtained from 118 *Pimoa* specimens. The final alignment included 411 positions and 4 additional gap characters. Individuals of *Pimoa* n. sp. (90 individuals) and *P. rupicola* (28 individuals) yielded 34 and 10 sequence types, respectively. The average *p*-distance within populations in *Pimoa* n. sp. and *P. rupicola* was 0.0102 ± 0.0027 and 0.0035 ± 0.0017 , respectively, and the maximum *p*-distance between the two lineages was 0.0701 ± 0.0121 .

The standard genetic diversity indices calculated for the *cox1* and the *ITS-2* for each locality are summarized in [Table 2](#). *Pimoa* n. sp. showed high levels of nucleotide diversity and low levels of haplotype diversity in most of the populations, while both *P. rupicola* and *T. vignai* showed low levels of haplotype and nucleotide diversity. This pattern was especially obvious in *P. rupicola*, since most individuals within populations were identical.

Pairwise F_{ST} values calculated for the localities of the three species are reported in [Table 3](#). Pairwise F_{ST} values between localities revealed contrasting patterns of gene flow. In *T. vignai*, pairwise F_{ST} values between localities were always higher than 0.8, and significant ($p < 0.05$), except for localities #14 and #16. A relatively strong population structure was also found in *P. rupicola*. Pairwise F_{ST} values between localities were always higher than 0.6, although significant comparisons involved exclusively southernmost localities (#28 and #29). *Pimoa* n. sp. showed instead a more shallow population structure, with several F_{ST} values below 0.5, generally corresponding to nearby localities.

Pimoa haplotypes were resolved as two independent networks, corresponding to *Pimoa* n. sp. (1) and *P. rupicola* (2), respectively, separated by 42 steps. *P. rupicola* haplotypes were limited to single populations, except H26 and H27, which were shared across populations. Generally, populations had low haplotype diversity, and in one case (#26) there was one single haplotype. Two divergent haplotypes (11 steps), however, were found in locality #29.

In *Pimoa* n. sp. haplotypes from nearby localities clustered together or were separated by only few steps (1–4). Several haplotypes (H7, H8, H12, H14) were shared among closely

Table 3 Population structure among localities. F_{ST} values for mtDNA *cox1* of *Pimoa* n. sp., *P. rupicola* and *Troglohyphantes vignai* based on the Tamura and Nei model. Locality codes are explained in Table 1. Localities #1, #2, #10, #15, #22 and #27 were excluded from the analysis, being represented by less than three individuals.

F_{ST} <i>Pimoa</i> n. sp.																
	3	4	5	6	7	8	9	11	12	13	14	17	18	19	20	21
3	0,000	–	–	–	–	–	–	–	–	–	–	–	–	–	–	–
4	0,250	0,000	–	–	–	–	–	–	–	–	–	–	–	–	–	–
5	0,723	0,018	0,000	–	–	–	–	–	–	–	–	–	–	–	–	–
6	0,787	0,123	0,332	0,000	–	–	–	–	–	–	–	–	–	–	–	–
7	0,557	0,322	0,412	0,341	0,000	–	–	–	–	–	–	–	–	–	–	–
8	0,700	0,764	0,597	0,422	0,857	0,000	–	–	–	–	–	–	–	–	–	–
9	0,433	1,000	0,733	0,733	0,590	0,733	0,000	–	–	–	–	–	–	–	–	–
11	0,200	1,000	0,500	0,500	0,357	0,500	0,230	0,000	–	–	–	–	–	–	–	–
12	0,340	0,561	0,642	0,642	0,500	0,642	0,376	0,142	0,000	–	–	–	–	–	–	–
13	0,289	0,642	0,589	0,578	0,446	0,589	0,322	0,006	0,218	0,000	–	–	–	–	–	–
14	0,500	0,589	0,800	0,800	0,657	0,800	0,533	0,300	0,442	0,357	0,000	–	–	–	–	–
17	0,450	1,000	0,750	0,750	0,607	0,750	0,483	0,250	0,392	0,186	0,470	0,000	–	–	–	–
18	0,533	0,750	0,833	0,830	0,690	0,833	0,566	0,333	0,476	0,422	0,633	0,583	0,000	–	–	–
19	0,366	0,860	0,660	0,666	0,523	0,666	0,400	–0,071	0,309	0,107	0,466	0,416	0,500	0,000	–	–
20	0,700	1,000	1,000	1,000	0,857	1,000	0,733	0,250	0,642	0,452	0,800	0,750	0,833	0,000	0,000	–
21	0,800	1,000	1,000	1,000	0,857	1,000	0,733	0,500	0,642	0,452	0,750	0,000	0,833	0,666	1,000	0,000
F_{ST} <i>Pimoa rupicola</i>					F_{ST} <i>Troglohyphantes vignai</i>											
	23	24	26	28	29		5	6	7	9	14	16	25			
23	0,000	–	–	–	–		5	0,000	–	–	–	–	–			
24	1,000	0,000	–	–	–		6	0,916	0,000	–	–	–	–			
26	1,000	1,000	0,000	–	–		7	0,964	0,855	0,000	–	–	–			
28	1,000	0,667	1,000	0,000	–		9	0,957	0,873	0,943	0,000	–	–			
29	0,876	0,876	0,876	0,667	0,000		14	0,991	0,988	1,000	0,991	0,000	–			
							16	0,989	0,984	1,000	0,989	1,000	0,000			
							25	0,987	0,984	0,992	0,988	0,975	0,988	0,000		

Notes.

Values in bold represent significant comparisons ($p < 0.05$).

located localities (e.g., occurring in the same Alpine valley or in adjacent valleys). In some instances, however, haplotypes from distant localities were found to be very similar (few steps), e.g., locality #1 and #2 with #12. Moreover, two haplotypes (H18 and H22) were found in individuals occurring in distant populations.

In *T. vignai*, haplotypes were not shared between localities. Haplotypes from closely located populations were generally separated by few mutations. Haplotype diversity was low within populations, and localities #7, #14 and #16 showed single haplotypes. Two localities (#9 and #25), on the other hand, had more than two haplotypes. The haplotypes of *T. vignai sensu rupicapra* were separated by 17–20 steps from the nearest locality (#14), which is actually the *locus typicus* of *T. vignai* (*Buco di Valenza* cave). A higher number of steps (38–56) separated this cave from the remaining localities.

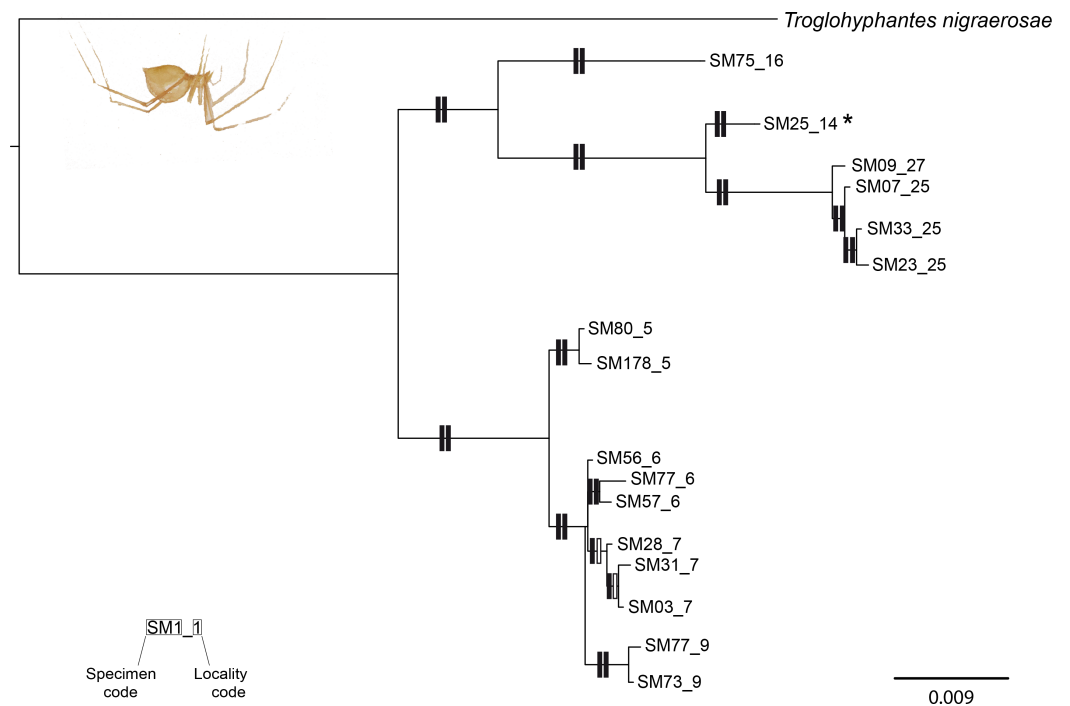


Figure 2 Phylogenetic tree of *Troglodyphantes*. Topology obtained in the concatenated Bayesian analysis for *Troglodyphantes vignai*. Only one individual per haplotype is shown. Vertical rectangles denote support as follows: Bayesian posterior probabilities (PP; left rectangles) and maximum likelihood bootstraps (ML; right rectangles); black: PP > 0.95, ML bootstrap support > 70%, white: support values lower than threshold values. The asterisk (*) indicate the *locus typicus* of *T. vignai*. Localities #25 and #27 refer to *T. vignai sensu rupicapra*.

Phylogenetic tree and estimation of the divergent time

Partition Finder selected the full codon as the best partition scheme for the alignments of both species. The models for each partition are reported in Table 2 of [Supplemental Information 1](#). Both the Bayesian and ML analyses of the concatenated data matrix of *T. vignai* yielded in similar tree topologies, and most branches were highly supported (i.e., posterior probabilities (PP) > 0.95, bootstrap support (BS) < 75%; [Fig. 2](#)). *T. vignai* was split in two main clades: one including the southern populations (#14, #16, #25 and #27), and the second one including the remaining northern populations (#5, #6, #7 and #9). The Bayesian and the ML analyses also recovered similar tree topologies in *Pimoa* ([Fig. 3](#)). Two well-supported clades were detected, corresponding to *Pimoa* n. sp. and *P. rupicola*, respectively. Individuals from geographical adjacent localities were closely related, although basal branches within *Pimoa* n. sp. were poorly supported.

The GMYC algorithm identified 2 coalescent clusters within *Pimoa cox1* sequences (ML = 100.8932; LR = 28.59767; $p < 0.000$), one including all sequences/localities of *P. rupicola* and the other including all sequences/localities of *Pimoa* n. sp. *Troglodyphantes vignai cox1* sequences were resolved as 7 coalescent clusters (ML = 424.2824; LR = 207.4615; $p < 0.000$; [Fig. 4](#)). Except for cluster A2, which included individuals from two caves, each cluster corresponded to individuals sampled from single caves.

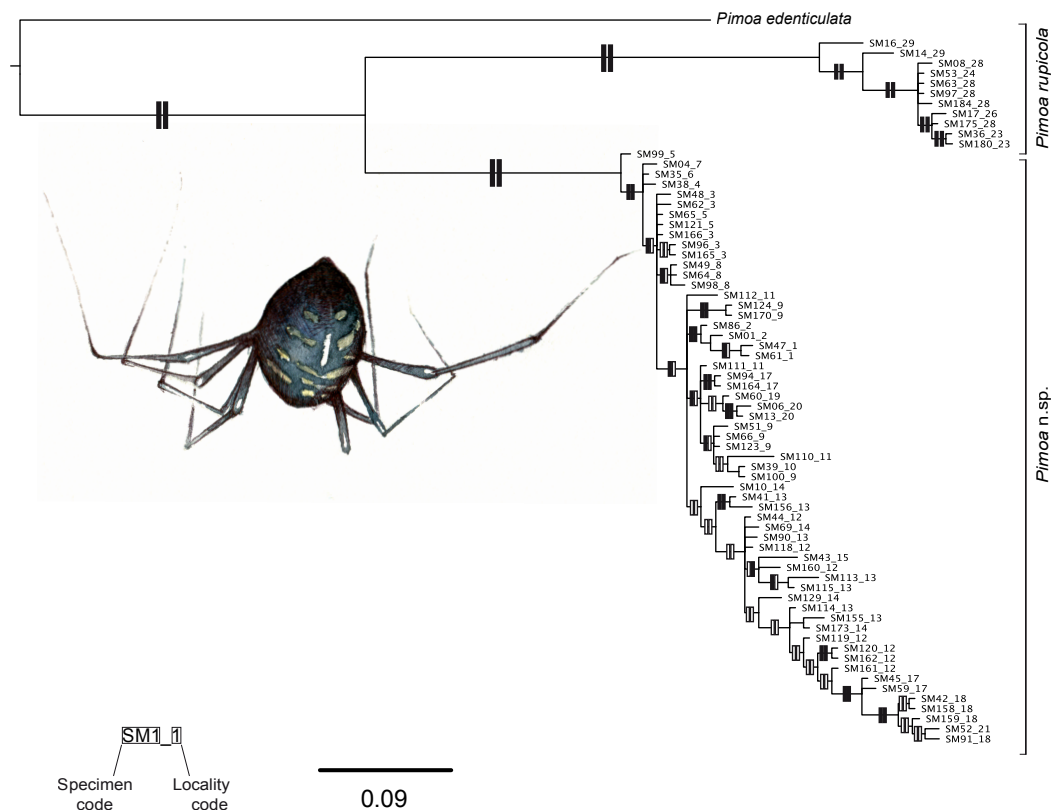


Figure 3 Phylogenetic tree of *Pimoa*. Topology obtained in the concatenated Bayesian analysis for *Pimoa*. Only one individual *per* haplotype is shown. Vertical rectangles denote support as follows: Bayesian posterior probabilities (PP; left rectangles) and maximum likelihood bootstraps (ML; right rectangles); black: PP > 0.95, ML bootstrap support > 70%, white: support values lower than threshold values.

The species trees and the embedded *cox1* gene tree recovered for each spider genus are shown in Fig. 4. The substitution rate estimated for the *Troglohyphantes cox1* was 0.0218 substitutions per lineage/million years (95% HPD = 0.010–0.033), and for the *ITS-2* was 0.0024 (95% HPD = 0.0005–0.0031). The split between *T. vignai* and *T. nigraerosae* was traced back to approximately 7.2 million years ago (Ma, 95% Highest Posterior Density, HPD = 13.7–3.5 Ma). Diversification of the extant *T. vignai* lineages occurred 2.9 million years ago (95% HPD = 5.4–1.5 Ma), while the diversification of the extant northern populations (D2, E2, F2, G2 clusters) occurred approximately 0.5 Ma (95% HPD = 0.9–0.2 Ma). The substitution rate estimated for *Pimoa cox1* was 0.0217 substitutions per lineage/million years (95% HPD = 0.011–0.033), and for the *ITS-2* was 0.006 (95% HPD = 0.002–0.013). The basal split between *Pimoa n. sp.* and *P. rupicola* was estimated to have occurred around 5.7 Ma (HPD = 12–2 Ma). The origin of the extant diversity of each lineage was estimated approximately at 0.4 Ma for both lineages (HPD = 1–0.15 and HPD = 0.85–0.15 Ma, for *P. rupicola* and *Pimoa n. sp.* respectively).

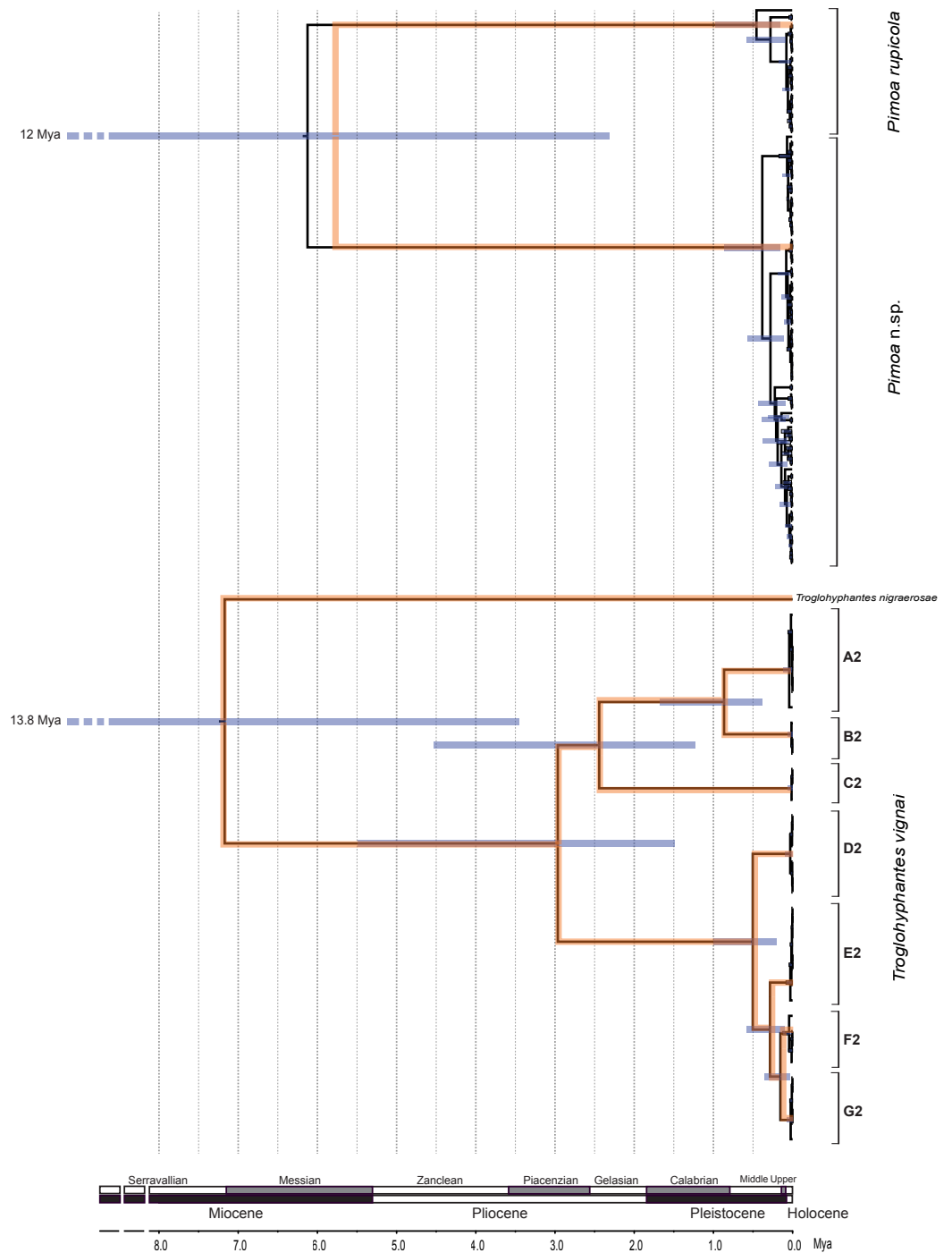


Figure 4 Timeframe of diversification. Chronograms obtained with the multispecies coalescent approach for the *cox1* and *ITS2* genes combined (orange topologies) and the *cox1* gene alone (black topologies). Grey node bars indicate the 95% HPD confidence intervals of the divergence time (for sake of clarity, only those HPD referring to the *cox1* gene are shown). The common x-axis is time in million years (Mya).

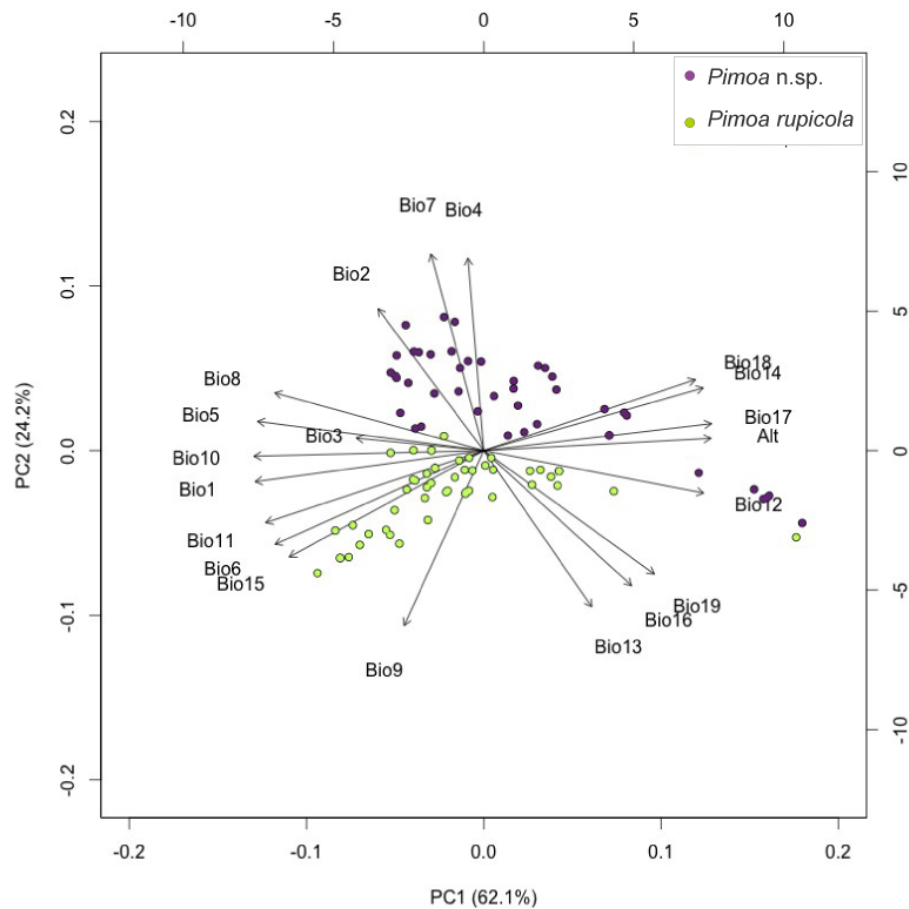


Figure 5 Climatic segregation of the *Pimoa* lineages. Bi-plot of Principal Component Analysis (PCA) scores for the first two axes based on 19 bioclimatic variables and altitude a.s.l. extracted for the localities of *Pimoa n. sp.* (purple dots) and *Pimoa rupicola* (green dots). For the explanation of the bioclimatic variables see Table 1 in [Supplemental Information 1](#).

Climatic segregation

We studied the climatic preference for the two *Pimoa* lineages with PCA, using bioclimatic variables (see Table 1 of [Supplemental Information 1](#)). The bi-plot of scores for the first two axes of the PCA is shown in [Fig. 5](#). The first two axes explained 86.2% of the variation in the data. The variance explained by other axis was negligible (<1% each). In respect to the first two axes, the two species segregate into two distinct clusters. The first axis (eigenvalues = 12.28; variance explained = 62.0%) mostly reflect a gradient of temperature. The second axis (eigenvalues = 4.79; variance explained = 24.2%) was positively correlated with variables reflecting diurnal and annual thermic excursion (Bio2, Bio4, Bio7) and anticorrelated with variables reflecting seasonal precipitation (Bio13, Bio16, Bio19). Although the first axis (PC1) explains most of the variance in the data, the localities of the two *Pimoa* species cluster in two groups according to the second axis (PC2), which combines bioclimatic variables referable to continentality. According to the original definition (see [Currey, 1974](#) for more details), continentality is intended as a measure

of how the climate is affected by its remoteness from the sea. Specifically, the distance from water masses influences the climate in terms of higher seasonality (Bio4), increasing diurnal (Bio2) and annual (Bio7) temperature ranges, as well as decreasing precipitation in the coldest (Bio19) and wettest (Bio13 and 16) periods. In light of our results, *Pimoa* n. sp. occurs in areas characterized by higher continentality in respect to *P. rupicola*.

Current and past distribution of Alpine *Pimoa*

The predictive performance of our bioclimatic models was fairly high both in *Pimoa* n. sp. (mean \pm SD AUC of the 20 runs = 0.845 ± 0.053) and in *P. rupicola* (0.908 ± 0.089). Overall, the suitable areas predicted by the model were congruent with the known distribution of the two species, at least in the Western Alps and Apennines. Current predictions identified suitable areas for *P. n. sp.* around the medium mountain belt (500–1500 m a.s.l.) of the Central and Western Alps, from the Camonica valley (province of Bergamo) down to the margin between Cottian and Maritime Alps (Stura valley, province of Cuneo). In respect to the known distribution, the predicted range of *P. n. sp.* extended northwards over the known limit of the species (see dotted line in Fig. 6A). More suitable areas were also detected in the northern edge of the Tuscan Apennines (Fig. 6A). The suitable areas of *P. rupicola* corresponded to the southern border of the Western Alps, in the coastal belt that spreads from Côte d'Azur (SW France) to the Ligurian eastern coast and Tuscan Apennines (Italy). Additional suitable areas were also found in Tuscany, Lazio and Corsica. The projection of the distribution model to the environmental condition of the Last Glacial Maximum (LGM) revealed that most of the current suitable areas were unsuitable in the LGM (Fig. 6B). Our threshold approach identified one main refugia for *P. n. sp.* (RF1) and two for *P. rupicola* (RF2 and RF3). All refugia corresponded to areas that were devoid from glaciers (in accordance with Ehlers, Gibbard & Hughes, 2011). The RF1 extended outside the southern edge of the actual distribution of *Pimoa* n. sp., in the hills and plains surrounding roughly the Northern border of Maritime and Ligurian Alps. RF2 corresponded to small areas along the French and Italian Riviera. RF3 extended over a wider geographic area in the Apennine, and in the northern part of the Corsica.

DISCUSSION

The history of two cave-dwelling spiders

The confounding effects of adaptation, biogeography and dispersal ability on the origin and the distribution patterns of cave organisms pose a stimulating challenge to biogeographers (Porter, 2007; Juan et al., 2010). According to Culver & Pipan (2010), long term climatic changes can be claimed as the main factors that prompted invertebrate species to colonize the subterranean habitats. In this regard, the Miocene climatic transitions and the Pleistocene glaciations are considered among the most important events. Here, by reconciling phylogeographic patterns and predictive ENMs, we provide support to this view, pointing out the Cenozoic climatic transitions has the most important factors shaping the present day genetic diversity and the distribution range of our model species.

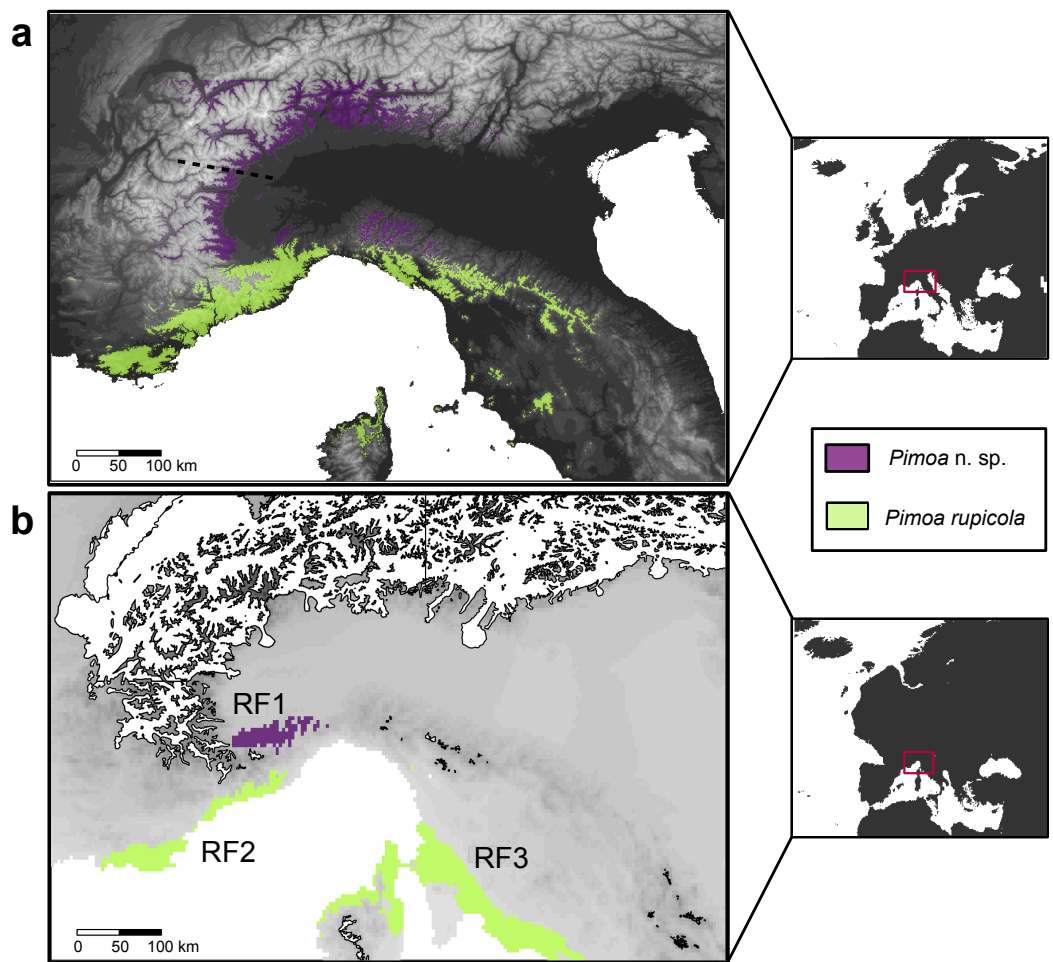


Figure 6 Current and past distribution of *Pimoa* lineages. Maps of the predicted environmental suitability according to the ENMs fitted to the occurrence points for *Pimoa* n. sp. (purple surface) and *P. rupicola* (green surface), at the present climate (A) and during the Last Glacial Maximum (B). Potential Pleistocene refugia (RF1, RF2, RF3) were identified by combining the three GCMs climatic reconstructions and applying a threshold of 0.6. The northern limit of the known distribution of *P. n. sp.*, corresponding to the Graian Alps (Isaia et al., 2011), is highlighted in the upper map with a dotted line. Limits of the ice cover in the Last Glacial Maximum (Ehlers, Gibbard & Hughes, 2011) are reported for Pleistocene projections (white shapes in the lower map).

Although special caution should be exercised when considering time estimates based on molecular data, especially in the absence of fossil record (Hipsley, 2014), our results fits well with some of the major climatic event undergone in the Western Alps during the Cenozoic. Accordingly, the two *Pimoa* lineages and *Troglodyphantes vignai* originated from the Middle (Serravallian) to the Late Miocene (Messinian), namely from 13 to 3.5 mya. More precisely, the isolation of *T. vignai* and *P. rupicola* from their northern sister groups (*T. nigraerosae* and *P. n. sp.*) dates back 7.2 and 5.7 mya, respectively. This time period approximately corresponds to the closure of the Gibraltar Strait and the onset of the so-called Messinian Salinity Crisis (MSC; after Ruggieri, Adams & Ager, 1967). However, given the large confidence intervals around our time estimates (~10 million years), it is

difficult to draw precise conclusions about the event that exactly determined the split of the different lineages.

It is worth noticing that the onset of a climatic transition in the Middle Miocene, marked the decline of the last global climate optimum conditions, leading to a progressive deterioration of the dominant subtropical climatic conditions (Suc, 1984; Shevenell, Kennett & Lea, 2004; Jiménez-Moreno, Fauquette & Suc, 2010). It is arguable that in parallel to the slow climate deterioration and the increase of seasonality, isolation of *Pimoides* and *Troglohyphantes* occurred.

Being possibly pre-adapted to shallow moist humid habitats (Deeleman-Reinhold, 1978; Hormiga, 1994; Zhang & Li, 2013; Wang et al., 2008), both species progressively colonized the subterranean habitat, most likely during the Pleistocene. Given their contrasting level of troglomorphy, it is likely that the process began earlier in *Troglohyphantes*.

The known distribution range of *T. vignai* stretches discontinuously along the Cottian (Chisone, Po and Varaita valleys) down to Ligurian (Valle Pesio) Alps. Conversely, the distribution of the northern sister group *T. nigraerosae* is adjacent southern Graian Alps (Isaia et al., 2011). The lack of shared haplotypes and the F_{ST} values close to 1 between the sampled localities of *T. vignai* indicates a strong isolation of the populations (Holsinger & Weir, 2009). The same idea is further corroborated by the identification of each population as an independent coalescent lineage (i.e., GYMC cluster). These results are in agreement with other studies on subterranean arachnids (Hedin, 1997; Hedin & Thomas, 2010; Dixon & Zigler, 2011) that support the “caves as islands” scenario (*sensu* Snowman, Zigler & Hedin, 2010), in which dispersal is virtually absent and the different populations diverge in allopatry. Under such conditions, it seems likely the subterranean habitat acted as an evolutionary *cul-de-sac* (see Fišer, Blejcek & Trontelj, 2012) for *T. vignai*. According to our time estimates the diversification of extant *T. vignai* lineages (especially northern population) occurred approximately during the Pleistocene glaciations. In this respect, it is worth noting that subterranean localities inhabited by *T. vignai* lie at the periphery of the Pleistocene glaciers (Ehlers, Gibbard & Hughes, 2011; see also local glacial limits reconstructed in Motta, 2014). Because subterranean populations most likely cannot survive under the ice cover (Culver & Papan, 2010), we suggest that the present day distribution range of *T. vignai* is the shadow of a wider ancestral distribution. Populations inhabiting the northern valleys in the Cottian Alps (Germanasca and Chisone valleys; see Fig. 4), where the ice shield was more compact, provide further evidences of the effect of Quaternary ice sheets.

In this area the distribution of *T. vignai* overlaps with the range of several hypogean species of *Doderotrechus* beetles (Carabidae, Trechini) (Giachino, 1993; Giachino & Vailati, 1997; Casale & Giachino, 2008). The similarity of distribution of both groups is most likely the result of a similar response to glacial dynamics. Gaining further knowledge on the biogeography of other subterranean species may provide further confirmation for the patterns here recovered.

In contrast with *T. vignai*, for which we did not detect any evidence of current population expansion, the topology of the chronogram obtained for *Pimoides* lineages

hints at a recent expansion following a bottleneck (see Fig. 4). The putative population expansion in both lineages of *Pimoa* would fall within the Quaternary Glacial Cycles, between 2.8 and 2.5 Ma (Gibbard, Head & Walker, 2010). The movement of glaciers as well as the continuous formation and melting of new ice sheets may have deeply affected the different populations, altering profoundly the local habitat suitability. Such transformations prompted either the migration of populations to more suitable areas at lower altitudes or latitudes or the local extinction of resident populations. Such scenario is congruent with the genetic fingerprint found in both *Pimoa* lineages. Accordingly, we hypothesize a glacial cycle-driven extinction of ancestral populations during cooler periods, followed by the expansion of populations which survived in climatic refugia during warmer periods. A similar pattern was observed in the Asian species *P. clavata* (Wang et al., 2008). The ENMs projected into the paleoclimatic reconstruction pointed out three putative areas devoid from glaciers that may have acted as glacial refugia for the surviving populations of each lineage (RF1-3; Fig. 6B). In the case of *P. n. sp.*, we detected one main macrorefugia (RF1) associated to the southernmost offshoots of the Alpine glacial masses. Notably, similar areas have also been classified as peripheral refugia for several Alpine plants, such as *Phyteuma globulariifolium* (Schönswetter et al., 2002) and *Ranunculus glacialis* (Schönswetter et al., 2003). For *P. rupicola*, potential macrorefugia were located along the SW French coast (RF2) and the Tuscan coast (RF3), including the northern part of Corsica. Even though the presence of *P. rupicola* in Tuscany is confirmed by literature records (Brignoli, 1971; Hormiga, 1994), recent investigation conducted by the authors did not confirm the present occurrence of the species in Corsica. Because of the larger spatial resolution of the LGM stacked rasters compared to the present day data (2.5 min versus 30 arc-seconds (~1 km)), it should be borne in mind that we may have not detected small point-like microrefugia (*sensu* Rull, 2009) within the interior of the Pleistocene ice shield covering the Alps.

As suggested by the lower F_{ST} values compared to *Troglohyphantes*, the relatively fast recolonization of ice-free areas is probably the result of the more effective dispersal ability of *Pimoa*. In particular, the population expansion followed a south-north direction, leading to the present distribution ranges of both lineages. Concerning *P. n. sp.*, we additionally predict suitable areas up to the Central Alps. Given the continuity of the suitable habitats predicted by the ENM and the supposed high dispersal ability of *Pimoa*, we hypothesize an ongoing expansion of the populations northwards. Indeed, the occurrence of *Pimoa* in outer shaded and humid habitats such as beech forests and other broadleaved forests (Bertkau, 1890; Jackson, 1926; Thaler, 1976; Isaia et al., 2015; Isaia, Paschetta & Chiarle, 2015) provides empirical evidence of the existence of epigeal dispersal. Because of the sex bias among the specimens collected in superficial habitats (Isaia et al., 2015; Isaia, Paschetta & Chiarle, 2015) and the general trend observed for spiders (Foelix, 1996), gene flow appears mostly mediated by males.

It should be borne in mind, however, that in light of the relatively low sample-size and the potential bias linked to the possible male-mediated dispersal, our *cox1* data (and the associated F_{ST} results) may not fully reflect patterns of gene flow (see, e.g., Willing,

Dreyer & Van Oosterhout, 2012; Davalos & Russell, 2014). Since low sample size is known to impact this kind of estimations, caution should be exercised when interpreting the values of nucleotide and haplotype diversity calculated for each locality (e.g., Goodall-Copestake, Tarling & Murphy, 2012). To minimize these potential bias, we left out from the calculation of the F_{ST} populations consisting of less than three individuals. Nevertheless, it is worth noticing that during data exploration we obtained very similar patterns of gene flow when comparing *cox1* and *ITS-2* results.

At present, the two lineages of *Pimoa* identified in this study show allopatric distributions (Fig. 6). *P. n. sp.* populations occur preferentially in areas characterized by higher continentality, and seem to tolerate cooler temperatures at higher altitudes and latitudes, as suggested by the results of the PCA (Fig. 5). On the other hand, *Pimoa rupicola* occurs in less continental areas, characterized by relatively small seasonal variations and high mean annual temperatures (i.e., Mediterranean climate). Similar complete niche partitioning between congeneric subterranean spiders has been reported elsewhere (Ribera, 1978; Gasparo & Thaler, 2000; Mammola & Isaia, 2014).

The application of ENM techniques has become a widespread practice to answer biogeographical and evolutionary questions (Franklin, 2009). In particular, ENM have been extensively used to identify Pleistocene refugia (e.g., Waltari et al., 2007; Rodriguez-Sanchez & Arroyo, 2008; Peterson, 2009; Planas et al., 2014). In constructing our scenarios, we have adopted a conservative approach, as our goal was to generate predictions under different levels of uncertainty. Although we relied on this approach, we are aware that ENMs have been rarely—and only recently—applied to study the hypogean ecosystems (see, e.g., Bryson et al., 2014; Camp et al., 2014; Naranjo, Moreno & Martiín, 2014). This is probably because, in first approximation, the link between the climatic variables (i.e., the external climate) and the subterranean habitat is not so straightforward. However, temperature of the underground compartment generally reflects the climatic regimen on the surface (Smithson, 1991; Badino, 2010). Although less intuitive, the regimen of rainfall plays an equally crucial role—if not more important—in determining such conditions (see details in Badino (2004) and Badino (2010)).

Overlooked diversity

In light of our results, some consideration regarding the overlooked diversity of our model species can be drawn. Concerning *T. vignai*, in this study we have included specimens from eight different localities, including topotypical material and material of *T. vignai* sensu *rupicapra*. The low levels of genetic variability observed between the latter and the topotypic material of *T. vignai* (Fig. 2, p -distance = 0.0022), provide further support for the synonymy between the two species proposed by Pesarini (2001). Therefore specimens of *T. vignai* sensu *rupicapra* have to be regarded as a population of *T. vignai* isolated in an area characterized by different climatic conditions (major Mediterranean influence). Such isolation could tentatively be related to the higher development of troglomorphism in *T. vignai* sensu *rupicapra*, as already observed by Brignoli (1971) and Isaia & Pantini (2010). Moreover, climatic factors provide a further line of interpretation for the presence of two

main lineages within *T. vignai*, corresponding to the northern and the southern clade (Figs. 1 and 2). Slight differences in the shape of the *lamella significativa* of the male palp could lead to consider the two lineages as candidate species (Vieites *et al.*, 2009), however the genetic distance between the two lineages, is about half of the value observed in the two nominal species *T. vignai* and *T. nigraerosae*.

The genus *Pimoa* is represented worldwide by 28 species (World Spider Catalog, 2015), many of which have only been described recently (Xu & Li, 2007; Xu & Li, 2009; Trotta, 2009; Hormiga & Lew, 2014). The application of molecular tools to investigate fine scale phylogeographic patterns in this group may uncover additional hidden diversity (Wang *et al.*, 2008). It is generally accepted that species delimitation and eventually species description should be based on the integration of multiple lines of evidence (Padial *et al.*, 2010). Here, we uncovered two deeply divergent genetic lineages (Fig. 3, GYMC clusters with *p*-distance above 9%) within *Pimoa rupicola*, which are further delimited both by genitalic morphology (S Mammola, G Hormiga, MA Arnedo, M Isaia, 2015, unpublished data) and different ecological requirements (Fig. 5).

CONCLUSIONS

Here, we have described the origin and the subsequent diversification of two species of spiders with contrasting levels of troglomorphy. We suggest that a different level of adaptation to subterranean life is an important factor to consider in the study of phylogeographic patterns. In particular, the major climatic events that occurred in the Alps during the Cenozoic determined from one side the complete isolation of pre-adapted subterranean species causing present day high population structuring, and from the other, the obliteration of surface-dwelling populations, causing their extinction or the lack of genetic structure in present day populations.

The parallel study of populations of subterranean species, especially when showing different levels of adaptation and overlapping ranges of distribution, may disclose new ways to understand patterns of biological diversification. Future research may include new highly variable nuclear markers and analytical tools, and also consider other taxa showing similar distributions (e.g., *Doderotrechus* beetles), to shed further light on the processes that shaped the present diversity of Alpine subterranean fauna.

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- Stefano Mammola conceived and designed the experiments, performed the experiments, analyzed the data, wrote the paper, prepared figures and/or tables, sampled hypogean habitats.
- Marco Isaia conceived and designed the experiments, reviewed drafts of the paper, sampled hypogean habitats.
- Miquel A. Arnedo conceived and designed the experiments, analyzed the data, contributed reagents/materials/analysis tools, reviewed drafts of the paper.

DNA Deposition

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