

High population differentiation in the rock-dwelling land snail (*Trochulus caelatus*) endemic to the Swiss Jura Mountains

Sylvain Ursenbacher · Caren Alvarez ·
Georg F. J. Armbruster · Bruno Baur

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Abstract Understanding patterns of genetic structure is fundamental for developing successful management programmes for isolated populations of threatened species. *Trochulus caelatus* is a small terrestrial snail endemic to calcareous rock cliffs in the Northwestern Swiss Jura Mountains. Eight microsatellite loci were used to assess the effect of habitat isolation on genetic population structure and gene flow among nine populations occurring on distinct cliffs. We found a high genetic differentiation among populations (mean $F_{ST} = 0.254$) indicating that the populations are strongly isolated. Both allelic richness and effective population size were positively correlated with the size of the cliffs. Our findings support the hypothesis that *T. caelatus* survived on ice-free cliffs during the Pleistocene glacier advancements from the Alps. Due to the establishment of beech and pine forest under recent, temperate climate conditions, dispersal between cliffs is no longer possible for rock-dwelling snails such as *T. caelatus*. Our results provide basic data for developing a conservation action plan for this endangered gastropod species.

Keywords Helicoidea · Land snails · Microsatellites · Population genetic structure · *Trochulus caelatus*

Introduction

The fragmentation of natural habitat is generally considered to be a major threat to many species. Population genetic theory predicts that the isolation of small populations lead to a reduction of genetic diversity. Human activities are often the main causes of habitat fragmentation, but geographical processes and/or specific habitat requirements may also contribute to natural segregation of populations. Species with limited dispersal ability particularly suffer from isolation, which may lead to a marked genetic divergence among populations (e.g. Conner and Hartl 2004). Furthermore, in isolated populations with decreasing population size, the risk of genetic drift increases which may lead to a reduction of genetic diversity (e.g. Frankham 2003) and an enhanced risk of inbreeding (Saccheri et al. 1998; Reed and Frankham 2003). Consequently, the evaluation of the extent of isolation of the existing populations and their genetic diversity are of major concern to assess the risk of disappearance of any threatened taxa.

Land snails are ideal model organisms to study population genetics of species with restricted dispersal ability. Several studies have focused on the micro-scaled structure of land snail populations (Schilthuizen and Lombaerts 1994; Anderson 2007; Armbruster et al. 2007a). *Trochulus caelatus* (formerly *Trichia caelata*) is a small terrestrial gastropod (shell height: 4 mm, width: 8 mm; Kerney et al. 1983) endemic to the northwestern Swiss Jura Mountains. This species is a highly specialised rock-dwelling snail restricted to limestone cliffs between 400 and 1,280 m above sea level with exposed rock faces resulting in an extremely patchy distribution in a small area (Turner et al. 1998). Due to its particular habitat requirements and the very limited distribution area, *T. caelatus* is considered as

S. Ursenbacher (✉) · C. Alvarez · G. F. J. Armbruster · B. Baur
Department of Environmental Sciences, Section of Conservation
Biology, University of Basel, St. Johannis-Vorstadt 10,
4056 Basel, Switzerland
e-mail: s.ursenbacher@unibas.ch

G. F. J. Armbruster
Botanisches Institut der Universität Basel, Schönbeinstrasse 6,
4056 Basel, Switzerland

potentially threatened on the Red List of Switzerland (Turner et al. 1994). During the last glacier advancement, this gastropod survived in ice-free tundra regions (nunataks) in the Swiss Jura Mountains. The temperature increase during the Holocene led to the establishment of beech forest and this species had to retract to the last open areas such as rock cliffs. The present distribution of *T. caelatus* coincides largely with areas that were ice-free during the last glacier advancement from the Alps (Turner et al. 1998), suggesting that this snail has not spread since the last glaciation. The surrounding forest may strongly reduce or even prevent dispersal among distinct cliffs. Consequently, we can presume that gene flow (if occurring) is restricted among populations.

We first examined 132 cliffs in the distribution area of the species in order to determine where *T. caelatus* is still occurring. When sufficient numbers of individuals were observed, samples were collected and levels of genetic variation and differentiation were investigated using microsatellite markers. We also conducted landscape genetic analyses to determine whether population groups

may occur and whether some evolutionary significant units (ESUs) or managements units (MUs; see Moritz 1994) can be determined. Our investigation will help to develop a conservation action plan for this rare gastropod.

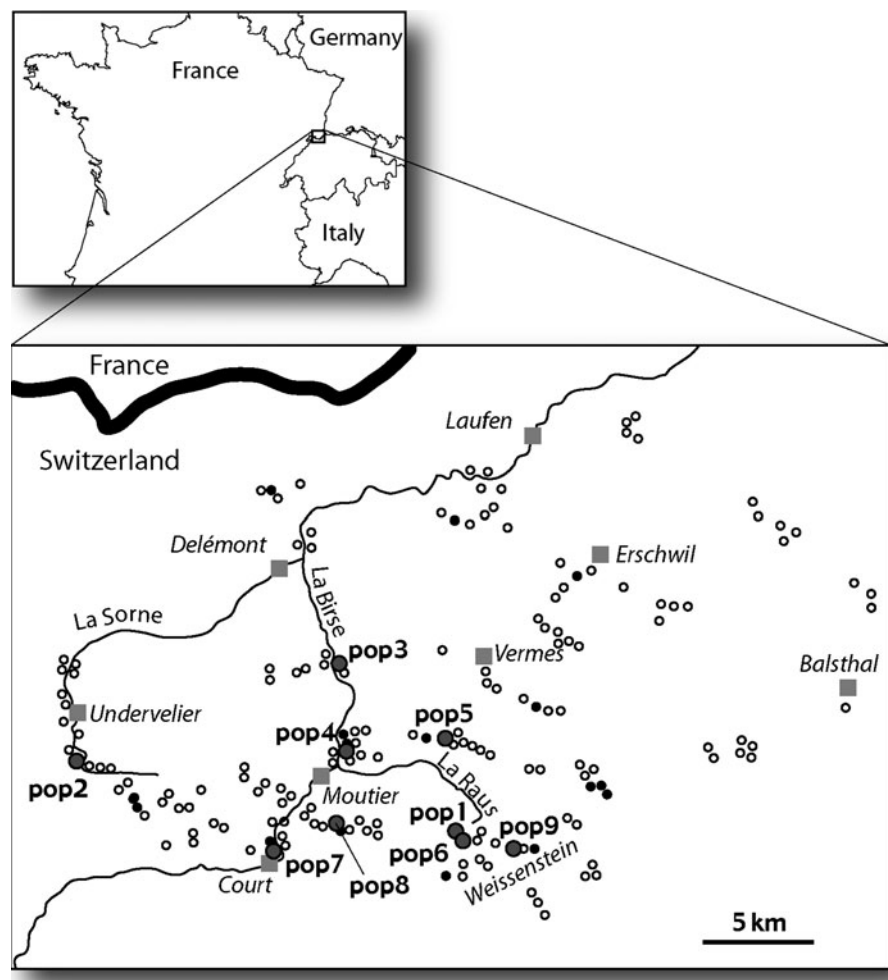
Material and methods

Field survey and sampling

One hundred and twenty-two cliffs, spread over the entire distribution area of *T. caelatus* were examined for the presence of the species between April and August 2006 (Fig. 1). The cliffs examined were located at altitudes ranging from 380 to 1,360 m above sea level. They consist of Jurassic coral chalks. The cliff bases are covered by different stands of deciduous forests belonging to *Fagetum* and *Tilietum* associations (Burnand and Hasspacher 1999).

One sampling plot (2 m high and 10 m long) was randomly placed along the foot of a cliff. A cliff was

Fig. 1 Locations of the cliffs examined for *Trochulus caelatus*. *Solid dots* indicate cliffs where the species was found, *open dots* indicate cliffs where the species was not found and *grey dots* to populations analysed in this study



considered as an isolated unit when it was separated from neighbouring cliffs by a strip of soil (>2 m) covered with vegetation. In each plot, the vertical rock surface, fissures and pockets were carefully searched for *T. caelatus* for 10 min., and the number of individuals was recorded. The presence of other snail species (*Trochulus montanus*, *Pyramidula rupestris*, *Cochlostoma septemspirale*, *Chondrina avenacea* and *Abida secale*) was also noted. Possible associations between species were tested using Chi-square tests. For each cliff the following variables were recorded: altitude (in metres above sea level), the horizontal length of the cliff (with five categories: ≤ 10 , 11–20, 21–50, 51–100 and >100 m), the height of the cliff (three categories: ≤ 5 , 5–10 and >10 m), the size of a cliff (calculated by cliff length \times cliff height), exposure of the cliff face, number of trees occurring at distances of 0–5 and 5–10 m from the cliff's foot and the cliff area (%) covered by grass and moss. Relations between occurrence of *T. caelatus* and cliffs parameters were tested using Chi-square tests and logistic regressions. For conservation reasons, samples of ten snails were only taken in the nine largest populations. Snail sampling was conducted on another day than the density estimate.

DNA extraction and microsatellite amplification

Collected snails were stored at -80°C . Genomic DNA was extracted from the whole soft body of the snail using the DNeasy Blood & Tissue Kit (Qiagen). DNA concentration was measured with a NanoDrop[®] ND-1000 UV-Vis Spectrophotometer (Thermo Fisher Scientific) and the DNA samples were diluted to a standard concentration of 10 ng/ μl .

Eight microsatellite loci (primers 08, 09, 10, 13, 16, 18, 22 and 26) developed for *T. caelatus* were used (Armbruster et al. 2007b). PCR was performed in 25 μl volume following the protocol described by Armbruster et al. (2007b). Spreadex EL-400 or EL-600 gels (Elchrom Scientific AG, Switzerland) were used to resolve allelic patterns by horizontal electrophoresis performed on SEA 2000 advanced submerged gel electrophoresis equipment using M3 size marker (Elchrom Scientific AG, Switzerland; see Armbruster et al. 2005).

Microsatellite analyses

Microsatellite results of each population were checked for null alleles and mis-scoring using MICRO-CHEKER version 2.2.3 (Van Oosterhout et al. 2004). GENEPOP version 4.0 was applied to assess the null allele frequency for each locus in all populations (Raymond and Rousset 1995; Rousset 2008). We estimated allele frequency, allelic richness (A_R),

observed and expected heterozygosity (H_O , H_E) using FSTAT version 2.9.3 (Goudet 1995). Heterozygote deficit within populations (F_{IS}) was estimated and divergence from Hardy–Weinberg (HW) equilibrium was tested using FSTAT with 1,000 permutations. Effective population size (N_e) was estimated using the linkage disequilibrium method proposed by Bartley et al. (1992) implemented in NEESTIMATOR version 1.1.1 (Peel et al. 2004). Spearman's rank correlations were calculated to examine relationships between genetic diversity (i.e. allelic richness; A_R), effective population size and environmental parameters (cliff size and snail abundance).

Genetic relationships within and among populations were investigated using FSTAT to estimate F -statistics (Weir and Cockerham 1984). Isolation by distance was tested using Mantel's test (Mantel 1967) by comparing pairwise ($F_{ST}/(1-F_{ST})$) values with the corresponding geographic distances (log-transformed) using the program FSTAT. This transformation has been showed to be more accurate in detecting isolation by distance than F_{ST} versus untransformed geographic distance (Rousset 1997).

In addition, Bayesian individual assignment approach as implemented in STRUCTURE version 2.2.3 (Pritchard et al. 2000) was tested using a model of admixture with a number of populations (K) ranging from 1 to 12. Likelihood values for ten replicates of each K value were estimated after 150,000 iterations (with the first 50,000 iterations discarded as burn-in). The best K value was chosen by applying the method of Evanno et al. (2005) which considers a second order rate of change to determine the most likely value of K .

Results

Habitat characteristics

Individuals of *T. caelatus* were found in 24 of the 122 (19.7%) plots examined (Fig. 1). The 24 populations were situated at an altitudinal range of 460–1,360 m in an area of 25 km \times 20 km. *T. caelatus* occurred more frequently on large cliffs (cliff length >50 m) than on small cliffs ($\chi^2 = 5.21$, $df = 1$, $P = 0.023$). Furthermore, the occurrence of *T. caelatus* was positively associated with a presence of *C. avenacea* or *A. secale* ($\chi^2 = 4.26$, $df = 1$, $P = 0.039$) and negatively associated with the number of trees at a distance of 5–10 m from the cliff's foot (logistic regression: odd ratio = 0.461 (95% CI: 0.243–0.875), $P = 0.016$). The presence of *T. caelatus* was, however, neither affected by cliff height, altitude, exposure of the cliff face, and the percentages of the rock surface covered by grass and moss (in each case $P > 0.1$). Interestingly, *T. caelatus* was found on cliffs exposed to all directions.

On most cliffs the abundance of *T. caelatus* was low. At 13 out of 24 sites (54.2%) only 1–5 individuals were found during 10 min. of searching. The largest numbers of snails per plot was 24 during a 10 min search period.

Population genetic diversity

Between 6 (P13) and 23 (P08) alleles were observed in the 90 individuals analysed in nine populations. Using MICROCHECKER, possible occurrence of null alleles was detected for locus P18 in one population, for P08, P10, P13 and P26 in two populations and for locus P16 in four populations. Allelic richness within populations varied between 2.91 (Pop 5) and 4.64 (Pop 8) (Table 1). The frequency of null alleles per population estimated with GENEPOP ranged from 0 to 0.450. Considering single primers, mean null allele frequency varied from 0 (locus P09) to 0.145 (locus P13). Four of eight markers showed significant deviations from HW equilibrium (P10, P13, P16 and P26). Consequently, H_O , H_E and F -statistics were calculated on the basis of the remaining four microsatellite markers (P08, P09, P18 and P22; Table 1). Estimates of the effective population size varied between 5.7 (Pop 5) and 25.9 (Pop 2). However, no estimates were available for Pop 3 and Pop 9 with the linkage disequilibrium method (Table 1).

Significant positive relationships between cliff size and allelic richness (Spearman's test: $r_s = 0.73$, $n = 9$, $P = 0.024$) and cliff size and effective population size

Table 1 Genetic diversity in nine populations of the snail *Trochulus caelatus* endemic to the Swiss Jura Mountains

Population	N_f	N_A	A_R	H_O	H_E	F_{IS}	N_e	95% CI
Pop 1	15	10	4.11	0.725	0.674	-0.076	25.5	12.4–240
Pop 2	15	10	4.13	0.575	0.537	-0.070	25.9	12.7–220
Pop 3	15	10	4.33	0.544	0.711	0.235	NA	–
Pop 4	20	10	3.79	0.625	0.718	0.130	7.1	4.9–11.4
Pop 5	24	10	2.91	0.500	0.526	0.050	5.7	3.6–10.4
Pop 6	16	10	3.28	0.575	0.628	0.084	8	5.0–15.4
Pop 7	5	10	4.37	0.700	0.736	0.049	15.9	9.0–46.3
Pop 8	16	10	4.64	0.625	0.7375	0.153	14.4	9.2–27.9
Pop 9	11	10	4.53	0.800	0.693	-0.15	NA	–

N_f corresponds to the number of snails found in an area of 20 m² within 10 min of search, indicating the population density. N_A corresponds to the number of snails genetically analysed. Allelic richness (A_R) was estimated for six diploid individuals for all eight loci. Observed and expected heterozygosity (H_O and values H_E), and heterozygote deficit within populations (F_{IS}) were estimated on the basis of the four loci in Hardy–Weinberg equilibrium. Effective population size estimates (N_e) with the 95% confidence interval (CI) were calculated using the linkage disequilibrium method implemented in NEESTIMATOR (Peel et al. 2004)

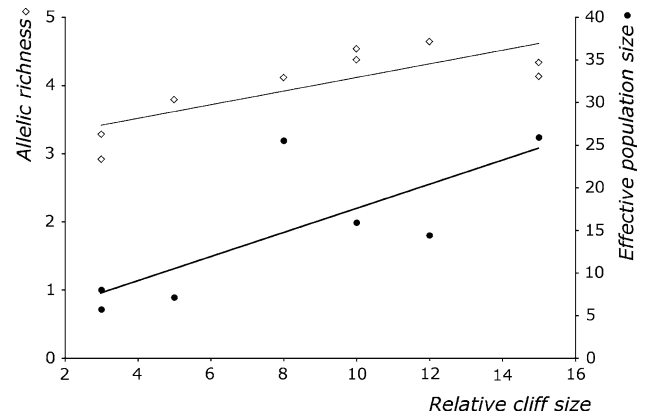


Fig. 2 Relationships between cliff size (height \times length) and allelic richness based on eight microsatellite markers in nine populations of *Trochulus caelatus* (left scale and open diamonds; Spearman's rank correlation: $r_s = 0.73$, $n = 9$, $P = 0.024$) and cliff size and effective population size calculated using genetic markers with a linkage disequilibrium method implemented in NEESTIMATOR (Peel et al. 2004) (right scale and full dots; $r_s = 0.88$, $n = 7$, $P = 0.008$)

($r_s = 0.88$, $n = 7$, $P = 0.008$) were found (Fig. 2). Furthermore, a marginally significant negative relationship between snail abundance and allelic richness was found ($r_s = -0.64$, $n = 9$, $P = 0.064$).

Population differentiation

High genetic differentiation was observed among the populations examined (mean $F_{ST} = 0.254$). All pairwise F_{ST} -values were significant (Table 2). The lowest value (0.099) was found between Pop 1 and Pop 6, which are isolated by only 100 m of unfavourable habitat. Isolation by distance was found (Mantel test: $r^2 = 0.213$, $n = 36$, $P = 0.011$), even if the fraction of genetic divergence explained by distance is limited.

The number of population groups was estimated to $K = 2$ using the method of Evanno et al. (2005). The populations 1, 5, 6 and 9 belong to one cluster, the populations 2, 3, 4, 7 and 8 to another. A sub-structure was detected in the first group. The program STRUCTURE suggested three sub-populations (one consisting of Pop 5, Pop 6 and Pop 9; one of four individuals of Pop 1 being regrouped with Pop 9 and one of six individuals of pop 1 together with Pop 6) and five sub-populations in the second group, each consisting of a distinct population (Pop 2, Pop 3, Pop 4, Pop 7, Pop 8).

Discussion

The field survey showed that the distribution of the endemic *T. caelatus* is restricted to a moderate number of small

Table 2 Pairwise F_{ST} estimates based on the analysis of four microsatellite loci in nine populations of the snail *Trochulus caelatus* endemic to the Swiss Jura Mountains

	Pop 2	Pop 3	Pop 4	Pop 5	Pop 6	Pop 7	Pop 8	Pop 9
Pop 1	0.275**	0.263	0.231	0.339	0.099**	0.197	0.202**	0.120**
Pop 2		0.321**	0.361	0.458**	0.337**	0.187**	0.299	0.302
Pop 3			0.230**	0.307	0.216**	0.147**	0.197	0.263
Pop 4				0.335	0.221**	0.243**	0.178	0.216**
Pop 5					0.383**	0.305**	0.311**	0.307
Pop 6						0.266**	0.182	0.127**
Pop 7							0.169	0.229**
Pop 8								0.165**

Significance was calculated with FSTAT (Goudet 1995) using 36'000 permutations. ** $P < 0.01$; for all other pairwise estimates, $P < 0.005$

populations in an area of 25×20 km. Relating the number of cliffs examined (122) to the number of cliffs existing in this region of the Jura Mountains, we assume that we have found 30–50% of the extant populations.

The analysis of genetic markers revealed a high degree of genetic differentiation between the examined *T. caelatus* populations, even in populations separated by less than 100 m. The results of individual oriented analysis (STRUCTURE) suggest a very restricted amount of gene flow between most populations. *Trochulus caelatus* is a rock-dwelling species living on vertical rock faces. The snail does not occur in the leaf litter of the forests surrounding the cliffs (Alvarez 2005). Considering the limited dispersal ability as indicated by the genetic analyses and by direct observations in other rock-dwelling snails (e.g. a maximum dispersal of 2.64 m per year was observed in *Chondrina clienta*; Baur and Baur 1995), we can conclude that most current populations are completely isolated. Between a few populations (Pop 1, Pop 6 and Pop 9), however, a low level of genetic differentiation was detected. The STRUCTURE analysis was not able to completely separate these populations which are situated about 100 m apart. Hence, we presume that a few gene exchanges might have occurred between these populations in the last decades.

Microsatellite markers have rarely been used to examine the population genetic structure of land snail species. In the closely related species *Trochulus villosus* a mean F_{ST} of 0.159 between 20 sites (maximum distance between sites: 300 km) has been reported (Dépraz et al. 2008). In *Helix aspersa* and *Cepaea nemoralis*, two snail species with larger shells, F_{ST} values of respectively 0.093 and 0.076 were found (Arnaud and Laval 2004; Schweiger et al. 2004). *Trochulus villosus* lives in the ground vegetation of deciduous forests, a habitat that frequently extends over hundreds of metres. Nevertheless, a continuous distribution of *T. villosus* seems to reduce the genetic divergence between sites probably due to gene flow. For isolated populations, high F_{ST} values were observed among nearby

populations in *H. aspersa*, *C. nemoralis* and *T. caelatus*. The F_{ST} values were slightly higher in *T. caelatus* than in the other species. However, this does not indicate a significant higher genetic differentiation compared to other land snail species.

Numerous other land snail species have small distribution areas with isolated populations. For example, *Trochulus biconicus* is endemic to a small mountain area in the Swiss Alps (M. Baggenstross, personal communication). *Arianta aethiops* occurs at a few localities in the southern Carpathian Mountains, Romania (Baur et al. 2000), and various land snails can be found on single carstic towers in Malaysia (Clements et al. 2008). Knowledge of genetic differentiation of the extant and mostly isolated populations of snail species with small distribution areas is extremely important for the success of future conservation and restoration actions.

Results of both population genetic and capture-mark-recapture studies (e.g. *Arianta arbustorum*, Baur 1986; *Albinaria corrugata*, Schilthuizen and Lombaerts 1994) confirm dispersal distances of only a few metres per year (not considering passive dispersal) in terrestrial snails. Long distance dispersal events due to passive transportation by wind, water, avalanches, mammals and birds are rare in land snails (Bengtsson and Baur 1993; Baur et al. 1997; Akçakaya and Baur 1996) and most probably insufficient to maintain gene flow between otherwise isolated populations. Furthermore, isolation by distance as found in this study suggests that the probability of passive dispersal is related to the distance between populations.

During the last glacial advancement, *T. caelatus* survived in ice-free sites in the Jura Mountains and did not spread afterwards (Turner et al. 1998). The lack of dispersal, combined with a limited number of ice-free sites in the Jura Mountains (Stehlik et al. 2001, 2002; Schönschwetter et al. 2005; Dépraz 2008) may explain the current distribution. We also found a positive association between cliff size and both genetic diversity and effective population size (in

contrast to *Discus macclintocki*, also a relict snail; Ross 1999). This finding implies the occurrence of numerous small refugia for *T. caelatus* during the last glaciation. Finally, the restricted dispersal may have limited the post-glacial colonisation of cliffs released from the ice sheet during the Holocene.

Conservation suggestions

The significant genetic differentiation of the populations examined and the snails' restricted dispersal ability indicate that current populations are completely isolated when they are separated by more than 200 m. Consequently, all these populations should be treated and protected as separated units (Management Units). The severe fragmentation of this endemic snail species, its small distribution area and the small population sizes strongly increase the risk of local and global extinction.

Changes in forestry practice (e.g. clear-cutting), which alter the lichen, moss and plant communities on the cliff walls, as well as rock climbing are the major threats to *T. caelatus*. Various specialised land snail species inhabiting rock cliffs in the Jura Mountains are negatively affected by climbing activities (B. Baur, unpublished data). The prohibition of sport climbing on cliffs with endangered land snails, lichens and plants is the most effective and adequate measure to preserve these species.

Small populations are also exposed to a high risk of extinction due to environmental or demographic stochasticity. Moreover, the relationship between cliff size and allelic richness indicates an increased risk of inbreeding and the occurrence of negative effects of inbreeding depression in small sites, and consequently a threat of disappearance of some populations. Because (re)colonisation of favourable habitats is restricted to extremely rare events of passive dispersal, the present population structure cannot be considered as a metapopulation complex. Therefore, any loss of a population would have a considerable impact on the survival of the endemic *T. caelatus* in the long-term.

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