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SHORT COMMUNICATION



Divergent selection in trailing- versus leading-edge populations of *Biscutella laevigata*

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• *Background and Aims* Knowledge on how climate-induced range shifts might affect natural selection is crucial to understand the evolution of species ranges.

• *Methods* Using historical demographic perspectives gathered from regional-scale phylogeography on the alpine herb *Biscutella laevigata*, indirect inferences on gene flow and signature of selection based on AFLP genotyping were compared between local populations persisting at the trailing edge and expanding at the leading edge.

• *Key Results* Spatial autocorrelation revealed that gene flow was two times more restricted at the trailing edge and genome scans indicated divergent selection in this persisting population. In contrast, no pattern of selection emerged in the expanding population at the leading edge.

• *Conclusions* Historical effects may determine different architecture of genetic variation and selective patterns within local populations, what is arguably important to understand evolutionary processes acting across the species ranges.

Key words: Amplified fragment length polymorphism, *Biscutella laevigata* (Brassicaceae), gene flow, genome scan, landscape genetics, range margins, selection, species range shift.

INTRODUCTION

Phylogeographic studies over the past decade have uncovered how past climate oscillations have repeatedly impacted on species ranges and influenced the genetic composition of taxa (Hewitt, 1996; Hampe and Petit, 2005; Schönswetter et al., 2005). Across a distribution range, climate changes trigger the regression of populations where conditions become suboptimal and they render new territories suitable for colonization (Ackerly, 2003). Accordingly, two types of margins can be distinguished: the trailing edge, where populations persist despite climate changes, and the leading edge, where populations expand following climate changes. The Alps went through drastic climatic oscillations during the Pleistocene and represent an excellent landscape study system to explore the consequences of climate-induced rangeshifts. There is now accumulating evidence demonstrating long-term persistence of several plant taxa, often at the border of the Alps, and postglacial recolonization from the peripheral to the central Alps and towards higher elevation (Schönswetter et al., 2005; Parisod, 2008). Consequently, the Alps harbour populations with a varying, contrasting history over a restricted geographical scale (i.e. similar climatic conditions, landscapes, biogeographical areas), which may greatly help to understand the evolutionary mechanisms underlying patterns of genetic variation under changing climate.

The focus here is on the widespread alpine herb *Biscutella laevigata* (Brassicaceae). At the regional scale, the influence of past climate changes on the species range was evidenced as a gradient of genetic diversity along recolonization

pathways (Parisod and Besnard, 2007). Accordingly, the peripheral Alps represent the historical core of lineages as attested by ample haplotypic variation among localities and presently correspond to trailing-edge conditions. In contrast, recently glaciated areas at high altitude in the central Alps are typically formed of expanding populations fixed for a single haplotype and correspond to the leading edge. Taking advantage of such historical demographic perspective offered by phylogeography, selected marginal populations of B. laevigata within a particular haplotype lineage (E; according to Parisod and Besnard, 2007) were examined at a local scale with amplified fragment length polymorphism (AFLP): one population corresponding to the trailing edge in the peripheral Alps (Parisod and Christin, 2008) and another population corresponding to the leading edge in the central Alps (Parisod and Bonvin, 2008).

Both populations showed similar total genetic diversity and presented mosaic spatial patterns of genetic differentiation, but distinct patterns of within-population genetic structure. The trailing-edge population presented a significantly lower within-plot genetic diversity than the leading edge (Shannon index: 0.093 ± 0.033 vs. 0.270 ± 0.057 , respectively; Wilcoxon test, P < 0.001) and a stronger internal genetic structure (between-group eigenanalysis: $\beta_{ST} = 0.49$ vs. $\beta_{ST} = 0.32$, respectively). Furthermore, individuals with similar AFLP profiles were consistently associated with ecological factors in the trailing-edge population, suggesting local adaptation with respect to total solar radiation (Parisod and Christin, 2008). In contrast, the leading-edge population manifested signatures

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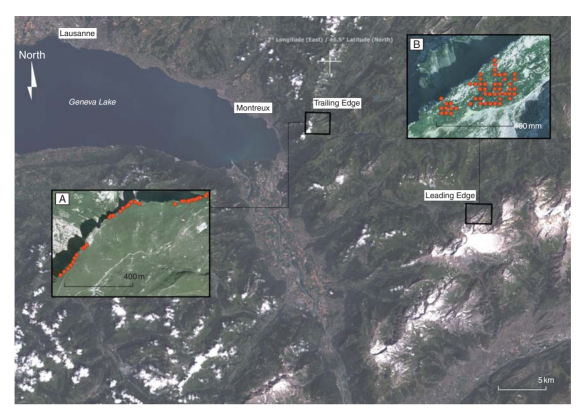


FIG. 1. Populations of *Biscutella laevigata* investigated at the trailing versus the leading edge of the species range. The basemap shows the western Swiss Alps (Landsat 7 image, July 1999). (A) The trailing-edge population sampled in the peripheral Alps; (B) the leading-edge population sampled in a glacier foreland in the central Alps. Dots represent the sampling plots.

of recent expansion since the distribution of both individual abundance and genetic diversity formed a cline along the local recolonization path (Parisod and Bonvin, 2008). Moreover, sampling plots from in between genetically homogeneous patches consistently showed higher genetic diversity, indicating ongoing admixture after recolonization.

The aim of this study is to explore the impact of divergent natural selection in different types of range margins by comparing indirect estimates of gene flow and genome scans for signature of selection between these trailing-edge and leading-edge populations. According to the hypothesis of Jansson and Dynesius (2002) postulating selection for local adaptation in areas with little climate-enforced range shifts, a clear signature of selection in the trailing-edge population and none or only a low one in the leading-edge population is expected.

MATERIALS AND METHODS

Investigated populations

Two populations of *Biscutella laevigata* L., one lying at the trailing and the other at the leading edge, were chosen for fine-scale surveys because they presented a large number of individuals arranged continuously across comparable altitudinal gradients (Fig. 1). Both populations showed great ecological similarity, sharing climatic conditions and presenting similar environmental heterogeneity as regards to the ecological factors assessed here on a local scale (Table 1; Supplementary data, available online). Individuals are strictly

outcrossing, characterized by a sporophytic incompatibility system (Olowokudejo and Heywood, 1984; C. Parisod, pers. obs.). Sampling was achieved across the whole populated area and four individuals were collected in evenly spaced 2×2 m plots according to a grid design. The trailing-edge population (JAM in Parisod and Besnard, 2007) was restricted to a linear ridge in the peripheral Alps (Parisod and Christin, 2008) and the leading-edge population (REU in Parisod and Besnard, 2007) was located at the altitudinal limit of the species in a glacier foreland of the central Alps (Parisod and Bonvin, 2008; Table 1). All individuals were genotyped with AFLPs, using the same combinations of fluorescent-labelled primer (M-CAG/E-ACA and M-CTC/E-AGG; Parisod and Bonvin, 2008; Parisod and Christin, 2008). Fragments were visualized after electrophoresis on 6 % Long Ranger denaturing gel using an ABI-PRISM 377 and manually scored (Table 1).

Comparative gene flow

Sp statistics are based on the spatial autocorrelation of genetic relatedness coefficients among individuals and represent convenient measures of gene flow within population, even under various sampling scheme such as linear vs. twodimensional populations (Vekemans and Hardy, 2004). Autocorrelation was assessed as the average relationship coefficient among individuals at increasing distance intervals with balanced number of pairwise comparisons, using SPAGeDi 1·2 TABLE 1. Description of the populations of Biscutella laevigata investigated at the trailing versus the leading edge of the species range

	Trailing edge	Leading edge	
Population's details			
Coordinates	6°58'/46°26'	7°12′/46°20′	
Historical features	Refugial populations in the peripheral Alps (Parisod and Christin, 2008)	Expanding population in the central Alps (Parisod and Bonvin, 2008)	
Spatial features	Linear $(1000 \times 25 \text{ m})$	Two-dimensional $(350 \times 200 \text{ m})$	
Grid sampling	31 plots of 4 individuals	51 plots of 4 individuals	
Environmental heterogene	eity*		
DEM: altitude (m)	(1851–1990); significant SA up to 140 m	(2154–2298); significant SA up to 140 m	
DDEG: degree-days	(1124-1260);	(839-921); significant	
during growing season (°d)	significant SA up to 140 m	SA up to 140 m	
ETPT: daily average evapotranspiration $(mm d^{-1})$	(2·5–6·6); low or no SA	$(2 \cdot 1 - 5 \cdot 6)$; low or no SA	
PDAY: number of precipitation days (d)	(62·9–63·75); low or no SA	(43-43); low or no SA	
SRAD: daily average radiation (kJ d^{-1})	(18057–75418); low or no SA	(17763–63928); low or no SA	
Slope (°) Genetic dataset	(2-69); low or no SA	(7–56); low or no SA	
Number of polymorphic AFLP bands	102	113	
Error rate (number of replicates)	1.8 % (20)	2.4 % (10)	

* Eco-climatic parameters discussed here are shown with their range of variation (in parenthesis) and a description of spatial autocorrelation (SA) over the sampling area (for details, see Table S1 in Supplementary data).

(Hardy and Vekemans, 2002). Confidence intervals (95%) were calculated after 999 permutations of individual locations among all individuals. In order to obtain a robust measure of relative gene flow in the trailing (A) and the leading-edge (B) populations, comparative Sp statistics were computed as the ratio

$$\frac{\sigma(A)}{\sigma(B)} = \sqrt{\frac{1 - F_1(A)}{1 - F_1(B)} \cdot \frac{D(B) \cdot b(B)}{D(A) \cdot b(A)}}$$

where F_1 is the relationship coefficient in the first distance class, D the effective density as estimated from field density (i.e. 3 and 2.5 individuals m⁻² in A and B, respectively), and b is the slope of the regression between the relationship coefficient and the logarithm of spatial distance. The surveyed populations are autotetraploid (i.e. with four segregating alleles per locus; Parisod *et al.*, 2010) and the relationship coefficient is thus increased by an unknown constant. This comparative approach has the advantage of circumventing such uncertainties. A 95 % confidence interval for the comparative Sp statistics was assessed by calculating the ratio again using opposite interval boundaries of the parameters (F_1 and b) for the two populations. In addition, autocorrelation and comparative *Sp* statistics were computed with all AFLP markers to deduce the realized gene flow and with neutral markers only (i.e. excluding those AFLP markers showing signature of selection as indicated by BAYESCAN, see below) to infer gene dispersal.

Detection of loci under selection

To detect signature of natural selection and to find ecoclimatic parameters putatively responsible for selection within the trailing- and the leading-edge populations of B. laevigata, two complementary approaches were used: (1) a Bayesian outlier locus approach estimating the posterior probability of each locus to be under selection using BAYESCAN software (Foll and Gaggiotti, 2008); and (2) an approach estimating the association between genetic data and environmental variables through logistic regressions as implemented in SAM (Joost et al., 2007, 2008). BAYESCAN is relying on estimates of genetic structure among sampling plots with the rationale that loci under positive selection present a significantly higher genetic differentiation than the multitude of neutral loci over the whole genome, while loci under purifying selection show lower differentiation. Accordingly, BAYESCAN was used on the subdivided populations (31 and 51 plots, according to the sampling scheme), running 150,000 iterations. Simulation studies demonstrated the robustness of BAYESCAN to a wide range of demographic scenarios violating underlying assumptions (Foll and Gaggiotti, 2008). In contrast, SAM does not rely on genetic models and works at the individual level, comprehensively inspecting correlations between loci and environmental variables by univariate logistic regressions (Joost et al., 2007). The eco-climatic parameters taken from the swisstopo 25-m-resolution digital elevation model and from Zimmermann and Kienast (1999) were used (see Table 1 for details). Although a formal comparison of the methods is beyond the scope of this paper, BAYESCAN and SAM profitably complement each other since the former authenticates signature of selection on theoretical grounds, while SAM identifies potentially effective selection pressure from the the environment.

RESULTS

The relationship coefficient among nearby individuals (F_1), as well as the slope of the regression between the relationship coefficient and the logarithm of spatial distance (b), was different in the trailing- and the leading-edge populations of *B. laevigata* (Table 2). In addition, F_1 and b remained similar after having removed putatively non-neutral loci from the dataset (see below), although the slope of the regression was lower in the trailing-edge population. Comparative Sp statistics computed from the whole AFLP dataset resulted in a ratio of 0.463 (95% confidence interval: 0.251; 0.908), indicating that realized gene flow was significantly more restricted in the trailing-edge population. Similarly, comparative Sp statistics computed from the neutral AFLP markers only resulted in a ratio of 0.468 (95% confidence interval: 0.281; 0.994).

In the trailing-edge population, BAYESCAN identified seven AFLP markers possibly under selection (Fig. 2A): A1 (posterior

	Trailing edge – all loci	Trailing edge – neutral loci	Leading edge – all loci	Leading edge – neutral loci
	(95 % CI*)	(95 % CI*)	(95 % CI*)	(95 % CI*)
$egin{array}{c} F_1^\dagger\ b^\ddagger \end{array}$	0.2280 (0.2109; 0.2510)	0·2094 (0·1925; 0·2307)	0.1526 (0.0765; 0.2145)	0·1531 (0·0769; 0·2153)
	-0.0397 (-0.0429; -0.0368)	-0·0337 (-0·0369; -0·0311)	-0.0069 (-0.0111; -0.0033)	-0·0071 (-0·0123; -0·0039)

TABLE 2. Spatial autocorrelation of AFLP genotypes in the trailing-edge and the leading-edge populations of Biscutella laevigata

* 95 % confidence interval (CI) as computed by 999 permutations.

[†] Relationship coefficient in the first distance class.

^{*} Slope of the regression between the relationship coefficient and the logarithm of spatial distance.

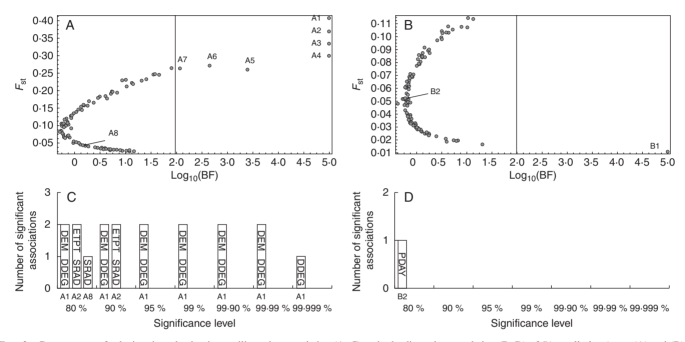


FIG. 2. Genome scans for loci under selection in a trailing-edge population (A, C) and a leading-edge population (B, D) of *Biscutella laevigata*. (A) and (B): BAYESCAN analysis, with the posterior probability for a locus to be under natural selection on the *x*-axis (log scale). The vertical line indicates the threshold beyond which there is a decisive evidence for selection. (C) and (D) SAM analysis, with significant associations between particular AFLP loci and eco-climatic parameters indicated on bars (DEM, altitude; DDEG, degree-days during growing season; ETPT, daily average evapotranspiration; PDAY, number of precipitation days during growing season; SRAD, daily average radiation). Sections on the *x*-axis refer to different significance levels. Loci detected by BAYESCAN and/or SAM are called A1 to A8 and B1 and B2.

probability = 1; \log_{10} of Bayes Factor = 5); A2 (pb = 1; \log_{10} BF = 5); A3 (pb = 1; \log_{10} BF = 5); A4 (pb = 1; \log_{10} BF = 5); A5 (pb = 1; \log_{10} BF = 3.4); A6 (pb = 0.998; \log_{10} BF = 2.66); A7 (pb = 0.992; \log_{10} BF = 2.07). SAM identified three loci putatively under selection (A1, A2 and A8) by total solar radiation, degree-days during growing season, altitude (poss-ibly correlated with other ecological determinants) (Fig. 2C). Two of the strongest outlier loci (A1 and A2) were clearly detected by both BAYESCAN and SAM. In the leading edge, BAYESCAN identified only one AFLP marker with strong evidence of purifying selection (Fig. 2B): B1 (pb = 1; \log_{10} BF = 5). SAM identified a different AFLP locus (B2) with a maximum confidence level of 80 % (Fig. 2D).

DISCUSSION

Genetic structure, as assessed here from the spatial distribution of AFLP genotypes through comparative Sp statistics, indicated that gene flow was above two times more restricted in the trailing-edge than in the leading-edge population of *B. laevigata* (ratio: 0.463). Despite differences according to uncontrolled factors (e.g. linear vs. two-dimensional habitats), the populations here surveyed presented great ecological similarity and were mostly distinguished by historical features (Table 1), suggesting an association between spatial genetic structure and the location at either the trailing edge or the leading edge.

Results from genome scans (BAYESCAN and SAM) pointed to a differential efficiency of selective processes in the two types of marginal populations (Fig. 2). Although such inferences in real populations have to be interpreted with caution (see Nielsen, 2005), the methods adopted here are reputed to provide reliable inferences under complex biological scenarios. The robustness of the BAYESCAN procedure against non-equilibrium situation was demonstrated, especially when assessing a reasonably large number of groups (Foll and Gaggiotti, 2008). Furthermore, SAM works at the individual level and is independent of any theoretical genetic model (Joost *et al.*, 2007). Comparative frameworks inspecting different populations with exactly the same approach, such as used here, should further alleviate from potential drawbacks since the main limitations are expected to apply equally to the contrasted situations. Finally, we argue that congruent results from totally independent approaches (BAYESCAN and SAM) bring decisive evidence favouring a reliable detection of the loci putatively under selection. Accordingly, the present results from genome scans indicate that, all else being equal, divergent selection was more efficient at the trailing edge than at the leading edge.

In the trailing-edge population, SAM highlighted loci (A1 and A2) referred as under directional selection by BAYESCAN to be significantly correlated to degree-days during growing season and total solar radiation. These ecological factors related to habitat thermal conditions were already associated with the genetic structure in this population (Parisod and Christin, 2008). The present results thus further indicate that selection by climatic factors may participate in maintaining pervasive genome-wide divergence at a local scale. Accordingly, since both BAYESCAN and SAM detected putative adaptive loci (Fig. 2), the stronger genetic structure observed in the trailing-edge population is linked to selection against migrants to a certain extent (Nosil et al., 2008). Since realized gene flow assessed with all AFLP markers was slightly lower than gene dispersal inferred with neutral markers only (Table 2), the loci detected here as under divergent selection might contribute to incipient reproductive isolation and, together with other barriers, sustain genome-wide association to environmental heterogeneity (Nosil et al., 2005; Parisod and Christin, 2008). In marked contrast, few loci putatively under selection were detected in the leading-edge population, and BAYESCAN and SAM provided incongruent results. No signature of divergent selection emerged in the leading-edge population of B. laevigata, although it was investigated with a stronger sampling than the trailing-edge one. This evidence supports the hypothesis that selective processes were less effective across the leading-edge population. Since expansion going on with high levels of admixture was detected in this population (Parisod and Bonvin, 2008), the lack of apparent selective pattern might be due to either comparatively intense recombination, limiting detection of selective patterns with genome scans, or to relatively high gene dispersal, hampering selection.

In this study, restricted gene flow at the trailing edge was observed with patterns of selection, while relatively high gene flow at the leading edge was associated with a lack of selection signature. Considering that species ranges experience strong historical heterogeneity due to past climate changes (Hampe and Petit, 2005), time passed since the establishment of populations may be an expected driver of spatial patterns of genetic variation. Indeed, gene pools persisting at the trailing edge obviously had long time periods to diversify and reach equilibrium, which suggests a strong influence of selective processes in sorting genotypes. In contrast, leading-edge populations are younger and characterized by non-equilibrium conditions, suggesting that dispersal processes may be prevalent in shaping genetic variation. Accordingly, the dynamic processes acting during expansion into empty habitats could promote either a thinning of genetic diversity, with an accordingly low potential of adaptation (Pujol and Pannell, 2008), and/or an increased gene flow, possibly preventing local adaptation (Parisod and Bonvin, 2008).

Replication at the population level is required to ascertain that historical fluctuation in population sizes and/or differential selection regimes shaped distinctive spatial genetic structure in trailing vs. leading-edge populations. However, equilibrium should not be expected across the entire range of species (Vucetich and Waite, 2003) and the present work stresses the need to rely on dynamic perspectives when investigating processes acting on populations at range margins. We advise using demographic insights from phylogeography to distinguish between trailing- and leading-edge populations in future studies, promising a deeper understanding of evolution and adaptation across the range of various species.

SUPPLEMENTARY DATA

Supplementary data are available online at www.aob.oxfordjournals.org and give the spatial autocorrelation of ecoclimatic parameters in the trailing-edge and the leading-edge populations of *Biscutella laevigata*.

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