# Secondary contact rather than co-existence – Erebia butterflies in the Alps

Short title: Secondary contact in Erebia butterflies

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# Abstract

Secondary contact zones are ideal systems to study the processes that govern the evolution of reproductive barriers, especially at advanced stages of the speciation process. An increase in reproductive isolation resulting from selection against maladaptive hybrids is thought to contribute to reproductive barrier buildup in secondary contact zones. While such processes have been invoked for many systems, it remains unclear to which extent they influence contact zone dynamics in nature. Here, we study a very narrow contact zone between the butterfly species *Erebia cassioides* and *E. tyndarus* in the Swiss Alps. We quantified phenotypic traits related to wing shape and reproduction as well as ecology in order to compare the degree of intra- and interspecific differentiation. Even though only very few first-generation hybrids occur, we find no strong indications for current reinforcing selection, suggesting that if reinforcement occurred in our system, it likely operated in the past. Additionally, we show that both species differ less in their ecological niche at the contact zone than elsewhere, which could explain why co-existence between these butterflies may currently not be possible.

Key words: Secondary contact, reinforcement, P matrix, Lepidoptera

### Introduction

When closely related lineages become geographically isolated, they may accumulate genetic incompatibilities through drift and divergent selection over time (Turelli et al. 2001; Coyne & Orr 2004). The evolutionary consequence of secondary contact between such lineages depends on the presence and strength of reproductive barriers that evolved during allopatry (Butlin et al. 2012; Canestrelli et al. 2016). Outcomes can range from substantial admixture when barriers are weak, to the formation of hybrid zones whose widths depend on the strengths of the barriers involved, and eventually to co-existence without gene flow (Harrison & Larson 2014; Gompert et al. 2017). Secondary contact zones provide excellent opportunities to investigate the evolution and interaction of reproductive barriers, often at an advanced stage of the speciation process (Gompert et al. 2017; Kulmuni et al. 2020). The reason is that selection against hybrids upon secondary contact could trigger the evolution of additional barriers to gene flow through reinforcement, *i.e.* the evolutionary process by which reproductive isolation increases in response to costly hybridisation (Dobzhansky 1940; Servedio & Noor 2003; Butlin & Smadja 2018). Although reinforcement has been invoked for some systems (e.g. Hoskin et al. 2005; Kronforst et al. 2007; Hopkings & Raucher 2012; Turelli et al. 2014), the relative frequency and importance of this process in nature remain debated (Matute & Cooper 2021). Expanding on findings from a former study (Lucek et al. 2020), we investigated the outcome of secondary contact between two sibling butterfly species of the genus Erebia from the Swiss Alps that form a very narrow contact zone (< 500 m) and assessed the potential for phenotypic signatures of reinforcement.

Reinforcement is predicted to be associated with the evolution of pre-zygotic barriers and may result in increased interspecific phenotypic differentiation in a zone of secondary contact compared to allopatric sites (Coyne & Orr 2004; Servedio 2009, but see Butlin & Ritchie 2013). Such character displacement often results in phenotypic and genetic clines across the contact zone, and in cases of resource competition or niche segregation, ecological clines may similarly occur (Barton & Hewitt 1985; Goldberg & Lande 2006; Gompert et al. 2017). The drivers of pre-zygotic isolation and the associated traits that experience selection often differ between taxa (Ravinet et al. 2017; Merot et al. 2017). If pre-zygotic isolation involves mate choice, divergence in mating relevant traits may be reinforced upon secondary contact, leading to reproductive character displacement (RCD; Gröning & Hochkirch 2008; Pfennig & Pfennig 2009). RCD has been shown for advertisement calls in chorus frogs (Pseudacris sp.; Lemmon 2010) or colour patterning of butterfly wings (Hinojosa et al. 2020). Divergence in genital morphology has similarly been invoked to result in reproductive character displacement (Hollander et al. 2018). The latter may be especially true for organisms with internal fertilisation, such as insects, where lock-and-key mechanisms have been suggested to be a powerful agent of selection against hybrids (Sota & Kubota 1998). For example, increased difference in the lengths of the male copulatory organ upon secondary contact has been shown to lead to failure of heterospecific matings in carabid beetles (Carabus sp.), resulting in the evolution of increased premating isolation (Usami et al. 2006; Nishimura et al. 2022). Importantly, character displacement resulting from reinforcement may increase trait divergence in either one of the two species involved, or in both (Cooley 2007; Wheatcroft & Qvarnstrom 2017; Dyer et al. 2018).

Secondary contact and reinforcement have been suggested to affect the multivariate phenotypic covariance structure (Blows & Higgie 2003; Dochtermann & Matocq 2016). Multivariate phenotypic evolution is thought to be constrained along so-called "lines of least resistance", *i.e.* the leading eigenvector of the **G** matrix ( $g_{max}$ ), which summarises the additive genetic variances and covariances (Lande 1979; Lande & Arnold 1983; Schluter 1996; Steppan *et al.* 2002). Biologically, this

axis captures the largest fraction of the genetic variance and is predicted to be shaped by selection and drift (Lande & Arnold 1983; Steppan *et al.* 2002; Marroig & Cheverud 2005; Arnold *et al.* 2008). In the absence of quantitative genetic data, the **G** matrix may be surrogated by the **P** matrix and  $p_{max}$  based on phenotypic data from wild populations (Cheverud 1988). This method is valid when phenotypic traits are heritable (Lande 1979), as has been found for many taxa (Cheverud 1988; Leinonen *et al.* 2011), including butterfly wing patterns (*e.g.* Palmer & Kronforst 2020; Nadeau *et al.* 2016) or insect genital morphology (*e.g.* Higgins *et al.* 2009; Andrade *et al.* 2009). Different **P** matrices can be compared by calculating the angle  $\theta$  between different **p**<sub>max</sub> (Schluter 1996). However, while the effects of gene flow and hybridisation on the **G/P** matrices have been studied both from a theoretical and empirical perspective (*e.g.* Guillaume & Whitlock 2007; Seehausen *et al.* 2014; Lucek *et al.* 2017), few empirical studies have looked at the outcome of secondary contact on the **G/P** matrices (Blows & Higgie 2003; Dochtermann & Matocq 2016).

*Erebia* is a genus of cold-adapted butterflies (Sonderegger 2005; Peña *et al.* 2005). The diversification of *Erebia* has been associated with differentiation in distinct glacial refugia due to the Quaternary glacial cycles (Sonderegger 2005; Schmitt *et al.* 2006; Schmitt & Haubrich 2008; Albre *et al.* 2008; Schmitt *et al.* 2014). Following postglacial range expansions, distantly related *Erebia* species often co-exist and exploit different microhabitats (Kleckova *et al.* 2014). However, closely related species or lineages exclude each other in several cases by forming very narrow secondary contact zones (Schmitt & Müller 2007; Descimon & Mallet 2009; Cupedo 2014; Lucek *et al.* 2020). Given the abundance of contact zones between different *Erebia* species or lineages, they provide an excellent system to study the outcome of secondary contact. For example, *E. cassioides* and *E. tyndarus*, two evolutionarily young species that split about two Myr ago (Peña *et al.* 2015), recolonized the Alps from different refugia (Schmitt *et al.* 2016; Gratton *et al.* 2016, Lucek *et al.* 2020), and form a very narrow contact zone in the central Alps, which has been stable since at least the 1950's (Warren 1954; Sonderegger 2005). A preliminary study with few individuals, found that the two closely-

related species form a very narrow genomic cline and found only a few F<sub>1</sub> hybrid individuals, suggesting selection against interspecific gene flow in this system (Lucek *et al.* 2020). The genomic cline overlapped with the presence/absence of the endosymbiotic bacterium *Wolbachia*, where 90% of *E. cassioides* were infected, as were the F<sub>1</sub> hybrids, while none of the studied *E. tyndarus* carried the symbiont (Lucek *et al.* 2020). While the potential role of *Wolbachia* in *Erebia* is still unknown (Lucek *et al.* 2021), it may act as an intrinsic postzygotic barrier to gene flow, potentially causing sterility of hybrids as in other butterflies (Nice *et al.* 2009). Indeed, *E. tyndarus* can be crossed with moderate success with *E. cassioides* when for the latter a distinct lineage from the Eastern Alps is used (Lorkovic 1958) that shares a *Wolbachia* strain with nearby *E. tyndarus* populations (Lucek *et al.* 2021). The genomic cline also overlapped with a phenotypic cline on wing patterns (Lucek *et al.* 2021). Wing-pattern recognition is often related to mate choice in butterflies (*e.g.* Kemp & Rutowski 2011; Hinojosa *et al.* 2020). Reproductive character displacement could thus have evolved to avoid costly hybridisation. However, Lucek *et al.* (2020) could not test this, as allopatric populations needed for comparison were unavailable.

Here we expand on the study of Lucek *et al.* (2020) and assess the potential footprint of reinforcement upon secondary contact, *i.e.*, evidence for reproductive character displacement on male genital morphology and wing shape. In a first step, we quantify the degree of intra- and interspecific phenotypic differentiation between individuals from geographically distant allopatric sites and expand on the formerly described contact zone in terms of sampling and geographical extent. Under reproductive character displacement, we predicted increased phenotypic differentiation in the secondary contact zone compared to the degree of allopatric differentiation in one or both species. We further compared the degree of multivariate phenotypic differentiation using **P** matrices, similarly predicting a shift in the multivariate phenotype between individuals from the contact zone and the allopatric sites or along the secondary contact zone. Finally, we also test for ecological character displacement along the contact zone of *E. cassioides* and *E. tyndarus*.

#### **Material & Methods**

## Sampling and data collection

We collected 841 male specimens of E. cassioides and E. tyndarus from 13 sites across Switzerland between June-September 2017-2020, with the vast majority caught in August (Fig. 1A, Fig. S1, Table S1). All individuals were caught by hand-netting and immediately euthanised with an overdose of ethyl acetate. For each specimen, we recorded its place-of-catch (GPS). We clipped the wings of each specimen, kept them in paper bags for further morphological analyses, and stored the body at -20°C. We used the coordinates to retrieve abiotic environmental parameters from a 25mresolution climatic dataset for Switzerland (Broennimann 2018). Monthly data on mean, minimum and maximum temperature (°C), precipitation (mm), growing degree days (i.e., the accumulation of temperature units during days where the temperature is above the 0°C threshold for alpine plant growth) and potential evapotranspiration (mm/day) were extracted for July-September, the months of active flight for *E. cassioides* and *E. tyndarus* (Sonderegger 2005). We further extracted the 19 bioclim variables as designated in the Worldclim database for the same time period (Fick & Hijmans 2017). Finally, because the geological substrate is often associated with broad-scale species distributions of mountain butterflies, since it may be related to the presence of food plants (Sonderegger 2005; Illán et al. 2010), we extracted substrate information for each specimen from the EuroGeoSurvey European Geological Data Infrastructure (Tulstrup et al. 2016).

Intra- and interspecific phenotypic differences in wing morphology, especially in the shape and extent of the orange spot on the dorsal surface of the forewing, have been observed for various *Erebia* species in the Swiss Alps (Sonderegger 2005). We assessed the phenotypic variation of our samples by digitising the right dorsal surface of the forewings of 833 out of 841 specimens with a flatbed scanner. Eight samples were discarded due to insufficient wing quality. When damaged, we scanned the left dorsal surface of the forewing instead, and the image was flipped horizontally (N=127). We captured phenotypic variation among and within *Erebia* populations by placing 23 landmarks on scanned wing images, focusing on defining wing shape based on venation and the shape and extent of the orange spot (Fig. S2A). Landmarks were placed using TPSDIG2 2.31 (Rohlf 2015), and Procrustes coordinates were calculated in MORPHOJ 1.07a (Klingenberg 2011). Additionally, we measured wing length, defined as the distance between landmarks 1 and 4 (Fig. S2A), using ImageJ 1.53a (Abràmoff *et al.* 2004). We applied a size correction by taking the regression residuals of the untransformed traits against measured wing length for each individual.

Like wing morphology, male genital morphology is a common characteristic to distinguish between *Erebia* species and subspecies (Sonderegger 2005; Cupedo 2014). Therefore, we clipped the genital apparatus of all 833 specimens used for wing morphology before macerating them in a 13% sodium hydroxide solution at room temperature for 24 h. Forceps were used to remove additional tissue. We stained the remaining sclerotised genitalia with a 3% Eosin Y solution in 70% ethanol for 5 min. Subsequently, we washed the genitalia in 70% ethanol, once for 5 minutes and then for 20 minutes. Stained genitalia were stored in 90% ethanol at -20°C. We photographed the valve structures using a stereomicroscope (Leica M205 C, Leica Microsystems, Wetzlar, Germany). We placed 11 landmarks on each resulting image with TPSDIG2, covering valve length and shape and shape and positioning of the first three-valve teeth (Fig. S2B), reflecting traits known to differ between *E. cassioides* and *E. tyndarus* (Sonderegger 2005).

#### Genotyping and species assignment

In order to assess the extent of interspecific gene flow, we used the restriction site associated DNA (RAD) sequence data from Lucek *et al.* (2020; N=84; NCBI BioProject Accession: PRJNA640280) and combined it with information from 235 newly genotyped individuals. Of the

latter, 152 were collected along the contact zone and the remainder across ten allopatric sites (Fig. 1A). DNA extraction followed Lucek *et al.* (2020). We used whole-genome resequencing (WGS) on a single Illumina NovaSeq 6000 S4 flow cell. Library preparation and sequencing were outsourced to the Genomics Facility Basel (D-BSSE of ETH Zurich). We aligned all data against the *Erebia ligea* genome (NCBI GCA\_917051295.2) using BWA mem v. 0.7.17 (Li 2013), followed by genotyping with BCFtools v. 1.15 (Li 2011). Only sites covered by both the RAD and the WGS datasets were retained. We then used VCFtools v. 0.1.16 (Danecek 2011) to apply a filter for minor allele frequencies (MAF)  $\geq$ 0.04, remove indels and to remove SNPs that were non-biallelic, had a genotype quality score  $\leq$ 20, had >60% missing data across all samples or a depth <5 or >30. Our filtering resulted in 2'387 SNPs for a total of 319 individuals. To perform a principal component (PC) analysis, we used Plink v. 2 (Chang *et al.* 2015). We further ran ADMIXTURE v. 1.3.0 (Alexander *et al.* 2009) assuming two genetic clusters (K = 2) to test for interspecific gene flow.

*Erebia cassioides* and *tyndarus* can be challenging to distinguish in the field, particularly upon secondary contact, where potential hybrids may show intermediate phenotypes (Sonderegger 2005; Lucek *et al.* 2020). We consequently employed a linear discriminant analysis (LDA) including all morphological traits for both wing and genital shapes to assign the individuals from the contact zone that were not genotyped to either species. To estimate the reliability of this approach, we first conducted an LDA using all allopatric individuals (N=347) and all genotyped males from the contact zone from Lucek *et al.* 2020 & 2021 (N=42). We then assigned all remaining individuals from the contact zone (N=444) to either species with the *predict* function in R version 3.6.3 (R Core Team 2020). This initial prediction was validated based on the 152 newly genotyped individuals from the contact zone. Of these, 97% were correctly assigned by the initial LDA to their respective species. Finally, we repeated our LDA assignment using all genotyped individuals, and to reduce potential false-positive assignments, we included only individuals within the 97% confidence interval for the respective phenotypic traits studied in all subsequent analyses. Given the low frequency of hybrids

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(N=3, Fig. 1), and their phenotypically intermediate characters (Fig. S3), we excluded these from subsequent phenotypic analyses.

#### Phenotypic differentiation between allopatric sites

To test if individuals from the contact zones would differ phenotypically from allopatric sites, we first summarised the phenotypic variation across all individuals with principal component (PC) analyses for wing shape, orange spot, and genital shape, respectively. We then tested for intra- and interspecific morphological differences between allopatric individuals and individuals from the contact zone by fitting linear mixed-effects models with Ime4 (Bates *et al.* 2015) in R on the PC1 scores for wing shape, orange spot, and genital shape, with *species* and *population type (i.e.,* allopatric or contact zone) as explanatory variables. We included location as a random effect and estimated the significance of each model with a type II Wald  $\chi^2$  test.

To capture intra- and interspecific changes in the phenotypic variance-covariance structure, we also compared the phenotypic covariance matrices (**P** matrices) among populations, by firstly calculating the angles ( $\theta$ ) between their leading eigenvectors  $p_{max}$  and secondly, the pairwise Mahalanobis distances between **P** matrices. For  $\theta$ , we calculated the dot product's inversed cosine between two  $p_{max}$  divided by the summed length of both  $p_{max}$  (Schluter 1996). To establish the statistical significance of each comparison, we used 10'000 bootstrap replicates as implemented in Lucek *et al.* (2014a & b). We estimated both  $\theta$  and the Mahalanobis distances for wing shape, the orange spot and genital shape, within species (using allopatric populations )and among species (using the individuals of either species that were collected from the eastern- or westernmost part of the contact zone). Given the much denser sampling, we excluded individuals closer to the secondary contact zone. We subsequently compared  $\theta$  and the Mahalanobis distances within and among species using an ANOVA with a Tukey HSD *post-hoc* test.

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We then tested whether the phenotypic traits of *E. cassioides* and *E. tyndarus* become more dissimilar the closer they are in proximity, which could indicate character displacement. For this, we repeated the PC analysis, including only individuals from across the contact zone to constrain the morphospace to phenotypic differentiation upon secondary contact, as PCs based on the full dataset could be driven by interspecific differentiation between allopatric populations. We then analysed the scores of PC1 for wing shape, orange spot, and genital shape, as well as for all individual traits, by fitting linear models. Explanatory variables were the *distance* of each specimen from the westernmost individual (in km), the *species (E. cassioides* or *E. tyndarus*) and their interaction. We subsequently repeated the analysis by fitting the same linear models for each morphological trait separately, also applying a False Discovery Rate (FDR; Benjamini & Hochberg 1995) correction to account for multiple testing.

Next, we fitted simple sigmoid clines across all individuals from the contact zones separately for PC scores of wing shape, orange spot and genital shape for PC1 – PC4. Cline estimations, using maximum likelihood approximation (BBMLE package in R, Bolker 2017), were based on the equations of Derryberry *et al.* (2014) adapted from Westram *et al.* (2018) to allow for an individual-based analysis. We fitted the clines using the individual geographic distances (km) from the westernmost individual. The best model was selected using Akaike's Information Criterion (AIC). We also performed a cline analysis for the genomic PC1 axis, for which hybrid individuals were included.

Finally, to test if the intraspecific **P** matrix may change along the contact zone, we employed an overlapping sliding-window approach, where for each species, we took 30 individuals starting from the point of contact and estimated  $\theta$  and the pairwise Mahalanobis distance between these samples and individuals from the utmost western- (for *E. cassioides*) or easternmost (for *E. tyndarus*)

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part of the contact zone. Window step size was by ten individuals, *i.e.*, removing the ten individuals closer to the point of contact and including the following ten individuals closer to the respective western- or easternmost parts of the range. We estimated the significance of  $\theta$  and the Mahalanobis distances with 10'000 bootstrap replicates. Because we included individuals that were not genotyped but assigned to a species by the LDA, we performed an additional local PCA for each species and phenotypic dataset and removed individuals outside the 95% CI from the subsequent **P** matrix analyses. We tested for changes in the **P** matrix along the contact zone for each phenotypic dataset using linear mixed effect models. The response variable was  $\theta$  or the Mahalanobis distance, with the fixed effect being the interaction between *distance* (the average position along the contact zone in km from west to east) and *species*. The random effect was the state of statistical significance estimated by our bootstrap approach.

# Differentiation in the abiotic environment

To assess the potential for niche differentiation in the abiotic environment between *E*. *cassioides* and *E*. *tyndarus*, we first tested whether the multivariate habitat would differ between the species among allopatric populations and between allopatric populations and the contact zone. As such, we summarised the environmental parameters in a PC analysis, and used the PC scores of the leading axis to fit a linear mixed-effects model with *species* and *population type (i.e.,* allopatric or contact zone) as explanatory variables and location as a random effect. We estimated the significance of each model with a type II Wald  $\chi^2$  test. We further selected the seven least-correlated, ecologically meaningful variables based on Pearson's correlation coefficient and variance inflation factor: potential evapotranspiration (mm/day) in July (EvapoJul), isothermality (Isothermal), precipitation seasonality (PrecSeason), precipitation in September in mm (PrecSept), mean temperature of the wettest quarter (TMeanWetQ), the minimum temperature in August (TMinAug),

and temperature seasonality (TSeason). Based on these variables, we ran niche similarity tests with 1'000 replications using the *ecospat* package (Di Cola *et al.* 2017) in R to quantify niche overlap (Broennimann *et al.* 2012) based on Warren's *I* (Warren *et al.* 2008) and Schoener's *D* (Schoener 1968) between *E. cassioides* and *E. tyndarus* in the allopatric and the contact zone and to determine if the species may undergo niche divergence.

We then focused on differences across the contact zone by fitting a linear model based on PC1 scores for contact zone individuals only, with *distance* from the westernmost individual and *species* as explanatory variables. We extracted the variance components of this model to disentangle each abiotic variable's contribution and then fitted individual sigmoid clines for the seven focus variables across the contact zone as for the phenotypic clines. We similarly compared substrate classes among species.

## RESULTS

### Genomic structure

Consistent with former studies (Gratton *et al.* 2016; Lucek *et al.* 2020), the genomic PC1 accounted for 70.2% of the total variation and clearly separated the two focal species, both between allopatric populations and along the contact zone (Fig. 1). We further identified three putative F1 hybrids that were genetically and phenotypically intermediate between *E. cassioides* and *E. tyndarus*, and found no apparent backcrossing (Fig. 1, Fig. S3). ADMIXTURE similarly separated the two species (Fig. 1C).

#### Phenotypic differentiation between allopatric sites

Of 830 specimens, 811 were retained as being within the 97% confidence interval, with 464 samples coming from the contact zone. Across the contact zone, we counted 186 *E. cassioides* (N<sub>Genotyped</sub>=71; N<sub>Assigned</sub>=115 and 278 *E. tyndarus* (N<sub>Genotyped</sub>=115; N<sub>Assigned</sub>=163), respectively.

Across all samples, we found wing shape to be significantly different between species along PC1, accounting for the majority of phenotypic variation (52.2%;  $\chi^2_{1,811}$  = 861.64, *p* < 0.001, Fig. 2A). Here, *E. cassioides* and *E. tyndarus* differ by the shape and extent of the orange spot on the forewing, which in *E. cassioides* is generally smaller, does not extend as far downwards towards the anal margin of the wing and often does not reach the cell of the wing (Table S2, Fig. 2A). Wing shape did not differ within species between individuals from the contact zone and allopatric sites ( $\chi^2_{1,811}$  = 2.39, *p* = 0.122). Focusing on the orange spot only, we observed the same pattern (Fig. 2B), *i.e.*, a marked difference between species along PC1, explaining a majority of phenotypic variation (59.3%;  $\chi^2_{1,811}$  = 874.97, *p* < 0.001), but no difference between individuals from the contact zone and allopatric sites ( $\chi^2_{1,811}$  = 2.46, *p* = 0.117). The shape of the male genitalia, a key character in distinguishing between the species ( $\chi^2_{1,811}$  = 1405.29, *p* < 0.001, Fig. 2C) along PC1 (32.5%). Overall, *E. cassioides* had shorter genital valves, and their first tooth on the valve was larger and wider than for *E. tyndarus* (Fig. 2C, Table S3). Like wing shape and the orange spot, genital shape did not overall differ between individuals from the contact zone and allopatric sites ( $\chi^2_{1,811}$  = 0.14, *p* = 0.714).

We also assessed differentiation within and between *E. cassioides* and *E. tyndarus* in their multivariate phenotypic covariance matrices. First by estimating the angle  $\theta$ , which captures the pairwise difference of the leading eigenvectors (PC1) between populations. Secondly, by calculating the Mahalanobis distance between population pairs, to quantify the overall differentiation between

two matrices. For  $\theta$  we found a significant differentiation for all phenotypic trait categories (ANOVA wing shape:  $F_{2,87} = 5.23$ , p = 0.007; orange spot:  $F_{2,87} = 5.91$ , p = 0.004; genital shape:  $F_{2,87} = 56.69$ , p < 1000.001; Fig. 3). Post hoc Tukey HSD indicate that intraspecific phenotypic differentiation was significantly higher among allopatric *E. tyndarus* than allopatric *E. cassioides* (wing shape: p = 0.009; orange spot: p = 0.005; genital shape: p < 0.001, Fig. 3). Similarly,  $\theta$  for interspecific comparisons was significantly higher than for intraspecific comparisons of *E. cassioides* (wing shape: p = 0.047; orange spot: p = 0.029; genital shape: p < 0.001) but not of *E. tyndarus* (wing shape: p = 0.387; orange spot: p = 0.365; genital shape: p = 0.085). The Mahalanobis distances differed similarly for all phenotypic trait categories (ANOVA wing shape:  $F_{2,87}$  = 23.05, p < 0.001; orange spot:  $F_{2,87}$  = 16.93, p < 0.001; genital shape:  $F_{2,87}$  = 108.00, p < 0.001; Fig. 3). However, post hoc Tukey HSD suggest no difference in intraspecific differentiation between allopatric *E. tyndarus* and *E. cassioides* (wing shape: p = 0.993; orange spot: p = 0.108; genital shape: p = 0.366, Fig. 3), yet all interspecific comparisons were significantly higher than the intraspecific comparisons for both E. tyndarus and E. cassioides (all p<0.001). Together these results suggest that the level of intraspecific differentiation is smaller than interspecific differentiation but that the leading eigenvectors differ even among populations within a species.

#### Phenotypic differentiation along the contact zone

We tested if individuals sampled closer to the point of contact were phenotypically more distinct than individuals caught further away by assessing whether our phenotypic traits changed with distance across our 14.58 km wide transect (Fig. 4A-C). For wing shape, although there was a marked difference between species (linear model:  $F_{1,464} = 249.91$ , p < 0.001, Fig. 4G) along PC1 (54.3%), there was no differentiation across the transect (*distance*:  $F_{1,464} = 1.21$ , p = 0.271) for neither species (*species*×*distance*:  $F_{1,464} = 2.22$ , p = 0.137). Using the same model for each individual

landmark, we similarly found that most wing traits (40 out of 46) differ between species (Table S4) but do not vary with distance across the transect (all p>0.05, Table S4).

For the orange spot, results were similar to those of wing shape. The phenotypic variation in the orange spot along PC1 (59.2%), differs between *species* ( $F_{1,464} = 260.01$ , p < 0.001, Fig. 4H) but individuals closer to the point of contact did not show increased phenotypic differentiation (*distance*:  $F_{1,464} = 1.08$ , p = 0.299; *species×distance*:  $F_{1,464} = 1.84$ , p = 0.175). For genital shape the same interspecific differentiation was found as for the overall dataset along PC1 (32.7%;  $F_{1, 464} = 447.21$ , p < 0.001, Fig. 4I). While *distance* was not significant ( $F_{1,464} = 0.14$ , p = 0.710), we found a significant *species×distance* interaction ( $F_{1, 464} = 10.82$ , p = 0.001), driven by *E. tyndarus*, where genital shape shifts towards the point of contact as individuals seem to become phenotypically more similar to *E. cassioides*. The latter is reflected at the level of individual landmarks, where *distance* played no role, but interspecific differentiation occurred in 13 out of 22 landmarks and an intraspecific shift across *distance* was found for five landmarks (Table S5).

The clines for wing shape, the orange spot and genital shape all overlapped and centred around the transition between *E. cassioides* and *E. tyndarus* (Fig. 4D-F, Table S6; wing shape: 9.08 km from the westernmost specimen [95% CI: 8.95-9.21]; orange spot: 9.10 km [95% CI: 8.99-9.21]; genital shape 8.97 km [95% CI: 8.95-8.99]). The clines were narrow compared to transect distance, *i.e.* ranging from 35 m (genital shape), over 216 m (orange spot) to 259 m (wing shape). Clines on subsequent PC axes could only be fitted for the second PC axes for all trait categories (Fig. S4). For these, the cline centres again overlapped at around 9 km, and all were narrow (wing shape: 175 m, orange spot: 199 m, genital shape: 227 m). Likewise, the genomic cline overlapped with the phenotypic clines, as its cline centre lies at 9.02 km from the westernmost individual [95% CI: 8.96-9.08]. Notably, the cline is only 125 m wide (Fig. 1D).

Intraspecific changes in the **P** matrix also occurred along the contact zone for both species (Fig. 5). For  $\theta$ , these changes often involved several phenotypic changes, as indicated by differences in the trait loadings of the local leading PC axes compared to individuals from the eastern or westernmost part of the contact zone (Tables S7-S9). For *E. tyndarus*, differences in  $\theta$  occurred primarily close to the contact zone, but an association with the distance gradient only occurred for genital shape as indicated by an overall *species*×*distance* interaction ( $\chi^2_1$ =11.72, *p* < 0.001; Fig. 5C). Here, intraspecific phenotypic changes primarily occurred along the horizontal axes, where *E. tyndarus* showed shorter valves near the contact zone than further away (Table S9). For *E. cassioides*,  $\theta$  differed only in some cases for wing shape and the orange spot, more closely to the western part of the contact zone (Fig. 5). While the overall **P** matrix similarly varied along the contact zone to the contact zone (Fig. 5).

# Abiotic environment

Allopatric *E. cassioides* and *E. tyndarus* differed significantly in their abiotic environment along PC1 ( $\chi^2_{1,811}$  = 9.64, *p* = 0.002, Fig. 6A), which accounts for 58.1% of the total variation and is mainly driven by temperature-related variables, growing degree days and some precipitation-related variables (Table S10). While individuals from the contact zone and the allopatric sites did overall not differ in their abiotic environment ( $\chi^2_{1,811}$ = 0.43, *p* = 0.512) there is some differentiation between species ( $\chi^2_{1,811}$  = 4.30, *p* = 0.038). Niche similarity based on the seven least correlated variables showed limited overlap between allopatric *E. cassioides* and *E. tyndarus* (Ca *vs.* Ta: Warren's *I* = 0.10, Schoener's *D* = 0.07) while the overlap was higher at the contact zone (Cc *vs.* Tc: *I* = 0.50, *D* = 0.30). The niche of allopatric individuals and individuals at the contact zone was more similar for *E. cassioides* (Ca *vs.* Cc: *I* = 0.38, *D* = 0.23) than *E. tyndarus* (Ta *vs.* Tc: *I* = 0.02, *D* = 0.08). Randomised replications suggest that the compared niches were not more diverged than expected by chance (all p > 0.05). In addition, for allopatric individuals, the two species occurred on different substrates, where *E. cassioides* is found primarily on limestone and *E. tyndarus* on gneiss substrates (Fig. S5). In contrast, individuals along the contact zone were collected exclusively on limestone, independent of species (Figs. 6B, S5).

In the contact zone, the abiotic environment changed with *distance* across the transect  $(F_{1,464} = 25.52, p < 0.001)$ , and differed between and within *species*  $(F_{1,464} = 4.44, p = 0.036; species×distance: <math>F_{1,464} = 339.40, p < 0.001$ ). At the level of individual ecological variables, differences most commonly include changes across the transect within one species (*E. cassioides*), while interspecific differentiation was found for only nine variables related to temperate and precipitation (Table S11). This shift within *E. cassioides* was reflected in the clines of the environmental variables, which were shifted westward compared to the phenotypic and genomic clines (Fig. 6C), indicating that the abiotic environmental variables are not correlated with the position of the contact zone. For PC1 the cline centre lies 2.82 km east of the westernmost individual [95% CI: 2.54-3.11 km] and the cline centres for each of the seven least-correlated variables overlap at about 3.5 km (Table S12). These results suggest a transition in the abiotic environment within the *E. cassioides* habitat from a slightly warmer to a colder environment with higher isothermality and potential evapotranspiration, but lower precipitation.

# DISCUSSION

Secondary contact may trigger reinforcement of existing or additional barriers to gene flow within a contact zone, but to which degree reinforcement occurs in nature or contributes to speciation remains unclear (Kulmuni *et al.* 2020; Matute & Cooper 2021). A final stage of speciation is co-existence and widespread sympatry, which may not always be achieved *e.g.*, when species or lineages fail to evolve enough ecological differentiation (Tobias *et al.* 2020; but see M'Gonigle *et al.*  2012) or when hybridization occurs even under high assortative mating (Irwin & Schluter 2022). As such, secondary contact zones may differ from one another in where they could be placed within the "grey zone" of late-stage speciation (Roux *et al.* 2016; Burbrink *et al.* 2020). Here, we studied the inter-and intraspecific phenotypic and environmental variation of two closely-related *Erebia* butterfly species that form a very narrow contact zone in the Swiss Alps and rarely hybridise (Descimon & Mallet 2009; Gratton *et al.* 2016; Lucek 2020; Fig. 1).

### The geographic extent of secondary contact

Contact zones in butterflies may extend over tens to hundreds of kilometres, often with substantial gene flow between lineages as found in *Lycaeides* butterflies from North America (Gompert *et al.* 2010) or tropical *Heliconius* butterflies (van Belleghem *et al.* 2021). In contrast, the closely related species *E. cassioides* and *E. tyndarus* are a case of exceptionally limited geographical contact, with their contact zone being less than a kilometre wide (Lucek *et al.* 2020; Fig. 1 & 4). However, an important gap in the aforementioned study was that it used only few individuals and focused on the narrow geographic region where the two species meet, precluding any inference about the full geographic scale of secondary contact and the potential outcome of reinforcing selection. When we expanded the sampled range for both species to several allopatric sites across the Swiss Alps, we found significant interspecific genomic differentiation with few intermediates (Fig. 1, S3), as well as differentiation for all phenotypic trait categories both for the respective leading axes of phenotypic variation (Fig. 2) and for the population-based multivariate covariance (**P**) matrices (Fig. 3). The **P** matrices further indicate that intraspecific differentiation is more predominant across allopatric *E. tyndarus* populations for the leading eigenvectors, especially for wing shape-related traits (Fig. 3A – B).

Likewise, genital shape showed little intraspecific differentiation for *E. cassioides*, compared to *E. tyndarus*, especially along the leading axis (Fig. 3C), suggesting stronger genetic constraints for these traits in the former species. Notably, at the broad scale, we did not find a statistical difference between allopatric and contact zone individuals, suggesting that putative ongoing reinforcement may only be at play across our studied contact zone. Indeed, our studied transect comprising the contact zone is relatively wide (14.58 km) compared to the actual point of contact (Fig. 4). As dispersal capabilities of *Erebia* butterflies are often limited, *i.e.*, a few hundred metres (Polic *et al.* 2014), individuals at the extreme end of this transect may experience only little to no interspecific contact, limiting the spread of potential reinforcing selection.

Given the consistent interspecific phenotypic differentiation among allopatric sites, we subsequently focused on the contact zone to assess the potential for reinforcement by fitting clines. Theory implies that the cline width and slope for a given trait depend on the extent of gene flow and the strength of selection against hybrids in a contact zone (Bímová et al. 2011; Bewick & Dyer 2014). A steep cline may result from strong selection against genetically or phenotypically intermediate individuals, suggesting that reinforcement could be at play (May et al. 1975; Bewick & Dyer 2014). If clines of different traits overlap, it may indicate that they are coupled, either because the different reproductive barriers may be genetically linked or because the strength and type of selection acting on them are the same (Slatkin 1975; Kruuk et al. 1999; Bierne et al. 2011; Bewick & Dyer 2014). The genomic cline and the clines for wing shape, orange spot and genital shape across the contact zone were extremely steep and narrow, with estimated widths for the cline centre ranging between 35 and 259 m (Fig. 4, S4). Therefore, direct or indirect selection against intermediate phenotypes for these traits seems likely, especially for genital shape, given the abrupt phenotypic transition (Fig. 4F). Notably, the phenotypic clines spatially overlap within this very narrow zone despite a putative lack of apparent geographic or environmental barriers at that location (Fig. 6B, 6C), indicating that they may experience a common selection regime. The phenotypic and genomic clines could otherwise be

a by-product of interspecific differentiation in the absence of heterospecific mating, yet hybridization occurs at this narrow point of contact. In contrast, most of the clines for the abiotic environment are much smoother and wider, and none overlap with the phenotypic clines (Fig. 6C), suggesting that ecological differences do not primarily drive the phenotypic clines.

#### Reinforcement - or the lack thereof

Given the narrow phenotypic clines, we tested for the "classic" signal of reinforcement, *i.e.*, whether the phenotypic traits become more dissimilar close to the point of contact when compared to allopatry (Coyne & Orr 2004; Servedio 2009). Importantly, this is only an indirect estimation, as character displacement across a contact zone may be consistent with reinforcement, but does not provide a direct measure of the presence, strength or impact of this process. Reinforcement has been suggested to act on wing shape and colour patterns in butterflies when involved in mate choice, as in the genus *Agrodiaetus (e.g.* Lukthanov *et al.* 2005), or mimetic *Heliconius* butterflies (Jiggins *et al.* 2001; Kronforst *et al.* 2007). For our *Erebia* contact zone, we found limited intra- but substantial interspecific variation in wing shape with no statistical support for potential ongoing reinforcement (Fig. 4G and 4H).

The phylogenetic splits between many butterfly species that currently form secondary contact zones across Europe are relatively old, as they predate several glaciation cycles (Ebdon *et al.* 2021). Such species may consequently have already been in contact during the Pleistocene and potentially experienced reinforcing selection in the past, as could be the case for *E. cassioides* and *E. tyndarus*, whose split has similarly proposed to predate the last glaciation (Peña *et al.* 2015). Reinforcing selection could therefore have occurred during a past interglacial period and phenotypes subsequently fixed in different glacial refugia. This could have caused the limited level of intraspecific differentiation, especially within *E. cassioides*. Reinforcement could also have occurred more recently, as the footprint of reinforcement is expected to diminish when intermediates

become rarer, especially if the trait on which reinforcement acts has a weak or intermediate effect on isolation (Bank *et al.* 2012). In such cases, the completion of speciation requires additional factors, which in our case may include differential infection by *Wolbachia* (Telschow *et al.* 2007; Lucek *et al.* 2020). If the traits under reinforcement caused strong isolation, then reinforcement alone can suffice to complete speciation (Bank *et al.* 2012), which in the case of *E. cassioides* and *E. tyndarus* seems not to have happened.

Unlike wing shape, the genital shape did show evidence for intraspecific phenotypic differentiation for PC1 along the contact zone for *E. tyndarus*, whose individuals close to the point of contact differed significantly in their genital shape when compared to individuals from the eastern part of the transect (Fig. 4I). Surprisingly, individuals at the point of contact tend to become more similar to *E. cassioides* (Fig. 4). Phenotypic convergence in sympatry can result from interspecific competition for essential resources, territoriality, or behavioural signals (Cody 1973; Leary 2001; Reifová et al. 2011). While increased convergence is contrary to classic predictions of reinforcement, different scenarios may account for the observed patterns: First, genital morphology may have resulted from introgressive hybridisation (Reifová et al. 2011). However, this seems unlikely given the apparent low hybridisation rates (Lucek et al. 2020; Fig. 1) and the narrow genomic (Fig. 1D) and phenotypic clines (Fig. 4). Secondly, intrinsic genetic incompatibilities may have evolved in allopatry in one or both species, preventing interspecific gene flow upon secondary contact. We could then still expect selection towards increased differentiation in mate-choice related traits in order to avoid interbreeding, unless assortative mating also arose as a by-product (Kulmuni & Westram, 2017). In such a scenario, reproductive interference could be at play, *i.e.* interspecific mating interactions leading to negative fitness effects on either one or both species, often resulting from incomplete species recognition. Much like reinforcement, reproductive interference can lead to a pattern of reproductive character displacement (RCD; Gröning & Hochkirch 2008). The key difference with reinforcement is that under reproductive interference, selection may act directly on phenotypes

related to mating behaviour in order to enhance preference for heterospecific signals (Shaw & Mendelson 2013). Reproductive interference may operate even when there is no interspecific gene flow upon secondary contact, except perhaps for a few F1 hybrids that do not backcross (Hollander et al. 2018). While the presence of only F1 hybrids with no further gene flow in our system (Fig. 1D) is consistent with reproductive interference, a clear distinction between reproductive interference (direct selection due to wasteful mating interactions) from reinforcement (indirect natural selection to avoid unfit hybrids and a gradual reduction in gene flow) would require to test for past gene flow (Hollander et al. 2018). Furthermore, both reinforcement and reproductive interference could result in RCD, and would require increased differentiation in traits associated with mate choice, which does not seem to involve the traits we assessed for wing shape (Fig. 2 & 4). However, pre-zygotic species recognition may, in this case, involve additional characters, such as olfactory cues through chemical signalling, which are often involved in mate choice in butterflies (Andersson et al. 2007; Constanzo & Monteiro 2007; Li et al. 2017). Although we could not assess the possibility of reinforcement of chemical signals between E. cassioides and E. tyndarus, this could be a promising avenue for future research in this system. Similarly, 3D micro-imaging of genital morphology could provide further insights in this system.

Even if closely related species are phenotypically strongly differentiated, they may evolve along shared evolutionary trajectories, whereby the leading eigenvectors of the species-specific **P** matrices ( $p_{max}$ ) would align (Dochtermann & Matocq 2016). Conversely, our intra- and interspecific comparisons between allopatric populations suggest that  $p_{max}$  differ between *E. cassiodes* and *E. tyndarus*, especially for genital shape (Fig. 3). The difference between wing shape and genital shape could result from a higher genetic integration of these traits (Arnold 2008). *E. tyndarus* shows a high level of intraspecific phenotypic differentiation for wing shape, which could indicate a higher standing genetic variation for these traits or additional intraspecific differentiation (Eroukhmanoff & Svensson 2011). Similarly to individual traits, the intraspecific **P** matrix may change along a contact zone (Dochtermann & Matocq 2016). Based on our sliding window approach, intraspecific changes occasionally occur for both species along the contact zone for  $p_{max}$  but not necessarily for the overall **P** matrix (Fig. 5). The latter was especially true for *E. tyndarus* individuals spatially close to the contact zone; however, to which degree these patterns could reflect local responses to selection requires further investigation. While it has been suggested that the **G** and **P** matrix estimation may require large sample sizes (Melo *et al.* 2015), Eroukhmanoff & Svensson (2011) suggest that small sample sizes are more likely to result in increased similarity. As such, our estimates are probably on the more conservative side. Overall, our analyses implicate that *E. tyndarus* is likely less phenotypically constrained than *E. cassioides*, as indicated by its increased level of intraspecific differentiation both across allopatric populations and along the contact zone (Figs. 3 & 5).

Despite significant phenotypic differentiation in traits linked to mate choice in other butterfly systems (e.g. Kemp & Rutowski 2011; Hinojosa *et al.* 2020), our focal species fail to co-exist. In addition, the presence (*E. cassioides*) or absence (*E. tyndarus*) of *Wolbachia* (Lucek *et al.* 2020, 2021), which may act as an intrinsic postzygotic barrier between the two species, nonetheless does not seem to prevent interspecific gene flow. Temporal isolation also seems unlikely to cause strong isolation, given that both species fly together at the contact zone (Figs. S1, S3). A common requirement for spatial co-existence is the utilisation of different ecological niches (Leibold & McPeek 2006), though other factors such as sexual selection can similarly promote co-existence on their own (M'Gonigle *et al.* 2012). Conversely, even a very small amount of hybridisation, or just interbreeding itself between lineages with strong assortative mating could suffice to prevent coexistence (Irwin & Schluter 2022). However, if neither ecology, sexual selection, nor their interaction suffices to complete reproductive isolation, competing species may stay in stable parapatry at contact zones (Tobias *et al.* 2020). Co-occurring *Erebia* species have been shown to differ in their microhabitat use (Kleckova *et al.* 2014), but to which degree this may be the case for our studied species remains unknown. Both *E. cassioides* and *E. tyndarus* use *Festuca sp.* grasses as

their larval host plants (Sonderegger 2005), and these occur in abundance in the contact zone (*personal observation*). However, it is not known whether they share the same host plant species. Similarly, there is limited evidence that nectar plants are shared by adults (Sonderegger 2005).

Focusing on more broad-scale ecological data, we found that the niches of the two species differ between their allopatric populations, but become more similar at the contact zone, however there is no indication of niche divergence among species. Interestingly, *E. tyndarus* from the contact zone seem to occupy a different niche than their allopatric counterparts (Fig. 6A). The latter is especially true for the geological substrate, where both species occur on limestone along the contact zone, whereas allopatric *E. tyndarus* primarily occur on gneiss (Fig. S5). The geological substrate is a commonly used proxy to describe species distributions of Alpine butterflies (Illán *et al.* 2010), including *Erebia* (Sonderegger 2005). Different substrates may have different effects on caterpillars in terms of temperature, humidity and presence of fungal endophytes, even when host plants are otherwise the same (Johnson *et al.* 1968). However, which aspects of the environment may be causal in shaping the actual distributions of our *Erebia* is unknown. Our result suggests that *E. tyndarus* may be able to expand its niche to different geological substrates, while this seems less likely for *E. cassioides*.

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Focusing on the zone of secondary contact, we did not detect a signal of reinforcement linked to ecology, as the occupied habitat appears not to be significantly differentiated between the two focal species at the contact zone. Indeed, niche overlap between the species was highest there, and the cline in the abiotic environment does not overlap with the phenotypic clines (Fig. 6C). Therefore, it may be possible that both species are more generalistic in their habitat use, as has previously been found in butterflies (*e.g.* Vodă *et al.* 2015), or that they are genetically constrained and cannot occupy different microhabitats. To which degree this could have contributed to the lack of co-existence requires further experimental investigation, and assessing the potential difference in microhabitats may be especially promising.

### Conclusion

Given their extremely narrow contact zone together with the limited level of interspecific gene flow, *E. cassioides* and *E. tyndarus* fall within the grey zone of advanced or late-stage-speciation (Roux *et al.* 2016; Kulmuni *et al.* 2020). However, speciation is not complete as the species fail to co-exist. Given the scarcity of hybrids, other pre-zygotic barriers are likely at play, but to which degree they, and the apparent lack of ecological niche divergence, could have contributed to the formation of a secondary contact zone that has been stable for decades (Warren 1954; Sonderegger 2005), requires further investigation. Interestingly, we did not find strong evidence for current reinforcement of our studied traits. Given that the split between the two species could be old, we may observe the outcome of repeated secondary contact following past reinforcement. The above suggests that the *cassioides-tyndarus* system provides an intriguing case of nearly-complete speciation, allowing to study the interplay between selection and ecology on the formation of barriers to gene flow and species co-existence. Similar processes may be more commonly at play among alpine species, where closely related species either form zones of secondary contact or exclude each other (*e.g.* Descimon & Mallet 2009; Capblancq *et al.* 2020).

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Figure 1: Sampling locations and genomic ancestry of Erebia cassioides and E. tyndarus. A – Map depicting sampling locations across Switzerland (Source: Google Maps, 2021, see Table S1). B,C – Genetic assignment of Erebia cassioides and E. tyndarus. B.) Barplot of the first principal component (PC) axis based on 2387 SNPs. Red = allopatric E. cassioides (Ca), orange = contact zone E. cassioides (Cc), green = hybrids, dark blue = allopatric E. tyndarus (Ta), light blue = contact zone E. tyndarus (Tc). C.) ADMIXTURE result for K = 2. D.) Cline fit for scaled scores of the genomic PC1 along a west-east transect in the contact zone. The black line represents the fitted cline, the vertical black bar the cline centre with its 95% CI in grey.



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Figure 2: Boxplots representing the phenotypic variation among populations along the leading principal component axes (PC1) for A - wing shape, B - the shape of the orange spot and C - genital shape. Wireframes depict phenotypes at a score of 0.1 and -0.1, respectively. Populations for each species are arranged from west to east. The contact zone is separated by species. Colours indicate species and location: red = allopatric Erebia cassioides, orange = contact zone Erebia cassioides, light blue = contact zone Erebia tyndarus, dark blue = allopatric Erebia tyndarus. Pictograms depict representative morphologies of each species (following Sonderegger 2005).



Figure 3: Inter- and intraspecific changes of populationbased **P** matrices. Boxplots summarise the angles  $\theta$  between the population-specific leading eigenvectors (**p**<sub>max</sub>) (A-C) or the overall Mahalanobis distances between **P** matrices (D-F) for: wing shape (A & D), the shape of the orange spot (B & E) and genital shape (C & F). Filled circles indicate significant and open circles non-significant estimates, respectively, based on 10'000

bootstrap replicates. Population comparisons are as follows: Ca – allopatric E. cassioides, Cc– contact zone E. cassioides, Ta –

allopatric E. tyndarus, Tc– contact zone E. tyndarus.  $\theta$  for each trait

category was compared, grouping intra- and interspecific comparisons. Significance levels based on a post hoc TukeyHSD test: \*\*\* p < 0.001; \*\* p < 0.01; \* p < 0.05; ns p > 0.05.



Figure 4: Summary of the phenotypic outcome of secondary contact. A-C – Elevation maps of the contact zone showing the locations of all studied Erebia cassioides (dots) and E. tyndarus (rhombi) individuals coloured by their respective PC1 scores based on A – wing shape, B – shape of the orange spot, C – genital shape. D-F – Cline fits between E. cassioides (C in orange) and E. tyndarus (T in light blue) along a west-east 42

transect for the same traits as A-C. The black line represents the fitted cline, the vertical black bar indicates the cline centre, with the grey area depicting its 95% confidence interval. Each cline was fitted based on the distance (km) from the westernmost individual. PC scores were rescaled. G-I – Model fits based on linear models testing for an interaction between the distance from the westernmost individual and species with the respective 95% CI for each model in grey. Colours depict species (orange: E. cassioides; light blue: E. tyndarus) and symbols indicate if species were determined based on genotyping (circles) or statistically assigned to a species (crosses).



Figure 5: A.-C.) Intraspecific changes in the P matrix along the contact zone for angles  $\vartheta$  between the subset specific leading eigenvectors ( $p_{max}$ ) (A-C) or the overall Mahalanobis distances between P matrices (D-F) for wing shape (A & B), the shape of the orange spot (B & E) and genital shape (C & F). The differences between the westernmost or easternmost individuals for E. cassioides (orange) and E. tyndarus (light blue),

respectively and subsets of individuals along the contact zones based on a sliding window approach are shown. Comparisons were only made within a species, where full circles depict significant, and the open circles non-significant values based on 10'000 bootstrap replicates. The line in C depicts a significant increase in  $\vartheta$  based on a linear mixed-effect model.



Figure 6: Summary of the abiotic environment. A - biplot of the two leading principal component axes based on all environmental variables (red = allopatric E. cassioides (Ca), orange = contact zone E. cassioides (Cc), light blue = contact zone E. tyndarus (Cc), dark blue = allopatric E. tyndarus (Ta). B - bar plots summarising the geological substrate classes where allopatric individuals and individuals from the contact zone were collected. C - Cline fits between E. cassioides and E. tyndarus along a west-east transect for the seven least correlated environmental variables: EvapoJul = potential evapotranspiration (mm/day) in July, Isothermal = isothermality, PrecSeason = precipitation seasonality, PrecSept = precipitation in September in mm, TMeanWetQ = mean temperature of the wettest quarter, TMinAug = minimum temperature in August, TSeason = temperature seasonality, and for PC1 based on all environmental variables = PC1 Environment (84%). The full vertical black bar indicates the cline centre for PC1 environment, with the grey area depicting its 95% confidence interval. The dotted vertical line indicates the cline centre for PC1 of genital morphology (see Fig. 4F). All cline fits were performed on scaled values (see methods).

