Supergene formation is associated with a major shift in genome-wide patterns of diversity in a butterfly.

Balancing selection at a wing pattern locus is associated with major shifts in genome-wide patterns of diversity and gene flow

Maria Ángeles Rodríguez de Cara1*, Paul Jay1*, Quentin Rougemont1*, Mathieu Chouteau1,2, Annabel Whibley3,4, Barbara Huber5, Florence Piron-Prunier3, Renato Rogner Ramos6, André V. L. Freitas6, Camilo Salazar7, Karina Lucas Silva-Brandão8, Tatiana Texeira Torres9, Mathieu Joron1$

* contributed equally

1 Centre d’Ecologie Fonctionnelle et Evolutive (CEFE), Univ Montpellier, CNRS, EPHE, IRD, Montpellier, France
2 Laboratoire Ecologie, Evolution, Interactions Des Systèmes Amazoniens (LEEISA), Université de Guyane, IFREMER, CNRS, Cayenne, Guyane Française
3 Institut de Systématique Evolution Biodiversité (ISYEB), Museum National d’Histoire Naturelle, CNRS, Sorbonne-Université, EPHE, Université des Antilles, Paris, France
4 School of Biological Sciences, University of Auckland, Auckland, New Zealand
5 Instituto de Ciencias Ecológicas y Ambientales (ICAE), Univ de los Andes, Mérida, Venezuela
6 Departamento de Biologia Animal, Instituto de Biologia, Unicamp, Campinas, São Paulo, Brazil
7 Department of Biology, Faculty of Natural Sciences, Universidad del Rosario, Carrera 24 No 63C-69, Bogotá 111221, Colombia.
8 Centro de Biologia Molecular e Engenharia Genética, Universidade Estadual de Campinas. Av. Candido Rondon 400. Campinas, São Paulo, Brazil
9 Department of Genetics and Evolutionary Biology, Institute of Biosciences, University of São Paulo (USP), São Paulo, Brazil

$ Corresponding authors: angeles.decara@gmail.com, paul.yann.jay@gmail.com, mathieu.joron@cefe.cnrs.fr, quentinrougemont@orange.fr
Abstract: Selection shapes genetic diversity around target mutations, yet little is known about how selection on specific loci affects the genetic trajectories of populations, including their genome-wide patterns of diversity and demographic responses. Adaptive introgression provides a way to assess how adaptive evolution at one locus impacts whole-genome biology. Here we study the patterns of genetic variation and geographic structure in a neotropical butterfly, Heliconius numata, and its closely related allies in the so-called melpomene-silvaniform subclade. H. numata is known to have evolved an inversion supergene via the introgression of an adaptive inversion about 2.2 million years ago, triggering a polymorphism maintained by balancing selection. This locus, which controls variation in wing patterns involved in mimicry associations with distinct groups of co-mimics, and butterflies show disassortative mate preferences and heterozygote advantage at this locus. We contrasted patterns of genetic diversity and structure 1) among extant polymorphic and monomorphic populations of H. numata, 2) between H. numata and its close relatives, and 3) between ancestral lineages in a phylogenetic framework. We show that H. numata populations which carry the introgressed inversions asin a balanced polymorphism show markedly distinct patterns of diversity compared to all other taxa. They show the highest genetic diversity and demographic effective population size estimates in the entire clade, as well as a remarkably low level of geographic structure and isolation by distance across the entire Amazon basin. By contrast, monomorphic populations of H. numata as well as its sister species and their ancestral lineages all show the lowest effective population sizes and genetic diversity in the clade, and higher levels of geographical structure across the continent. This suggests one hypothesis is that the large effective population size of polymorphic populations could be caused by the shift to a regime of balancing selection and disassortative preferences associated with inversions. Testing this hypothesis with forward simulations supported the observation of increased diversity in populations with the supergene. Our results are consistent with the hypothesis that the adaptive introgression formation of the inversion supergene triggered a shift from directional to balancing selection and a change in gene flow due to disassortative mating, causing a general increase in genetic diversity and the homogenisation of genomes at the continental scale.

Introduction: Genetic diversity is shaped by selective processes such as stabilizing or disruptive selection, and by demographic processes such as fluctuations in effective population size. Empirical studies on genetic diversity within and among populations abound, fuelled by an increasing availability of whole genome data, and spurred by our interest in understanding the underlying causes of variation in diversity (e.g. Beichmann 2018, Muers 2009; Murray 2017; Nielsen et al. 2009). At the locus scale, strong directional or disruptive selection tends to reduce diversity within populations (Mitchell-Olids et al. 2007), while balancing selection tends to enhance diversity (Charlesworth 2006). Genome-wide factors reducing diversity include low effective population sizes, generating drift, while high genetic diversity is enhanced by large population sizes and gene flow. Overall, it is well recognised that demographic changes should have a genome-wide effect on diversity, while positive selection is expected to play a role on the sites within and around the genes involved in trait variation (Glinka et al. 2003, Muers 2009, Nielsen et al. 2009).

Variation in behaviour and life-history traits, for instance involving changes in offspring viability or dispersal distance, may also affect species demography, and thus whole genome genetic diversity. However, whether and how genetic variability in a population may be driven by phenotypic evolution at certain traits is poorly understood, and confounding effects may affect patterns of
Genomic diversity, such as variation in census population size or colonization history. Dissecting how selection on a trait may affect genome-wide diversity can be tackled by comparing closely-related populations differing at this trait coupled with knowledge of when the differences evolved. Here, we took advantage of the dated introgressive origin of a chromosomal inversion associated with major life-history variation to study the demographics and whole genome consequences of changes in the selection regime at a major-effect locus.

*Heliconius* butterflies are aposematic, chemically-defended butterflies distributed over the American tropics from Southern Brazil to Southern USA (Emsley 1965; Brown 1979) (Fig 1A). *Heliconius* butterflies are well-known for visual resemblance among coexisting species, a relationship called Müllerian mimicry which confers increased protection from bird predators through the evolution of similar warning signals (Sheppard et al. 1985). Most species are locally monomorphic, but their mimicry associations vary among regions, and most species display a geographic mosaic of distinct mimetic “races” through their range. In contrast to most *Heliconius* species, the tiger-patterned *Heliconius numata* is well-known for maintaining both mimicry polymorphism within localities, with up to seven differentiated coexisting forms, and extensive geographic variation in the distribution of wing phenotypes (Brown & Benson 1974; Joron et al. 1999). Forms of *H. numata* combine multiple wing characters conveying resemblance to distinct sympatric species in the genus Melinaea and other local Ithomiini species (Nymphalidae: Danainae). Polymorphism in *H. numata* is controlled by a supergene, i.e. a group of multiple linked functional loci segregating together as a single Mendelian locus, coordinating the variation of distinct elements of phenotype (Brown & Benson 1974; Joron et al. 2006). Supergene alleles are characterized by rearrangements of the ancestral chromosomal structure, forming three distinct chromosomal forms with zero (ancestral type, Hn0), one (Hn1) or three chromosomal rearrangements (Hn123) (Fig 1B). The ancestral arrangement, Hn0, devoid of inversions, is fixed in most *Heliconius* species (although an inversion in the same region evolved independently in a distantly-related *Heliconius* lineage (Edelman et al. 2019)). Arrangement Hn1 contains a 400kb inversion called P1, originating from an introgression event about 2.2 My ago from *H. pardalinus*, in which P1 is fixed (Jay et al. 2018). This introgression is thought to be the founding event triggering the formation of the supergene and the maintenance of polymorphism in *H. numata* (Jay et al. 2018). Arrangement Hn123 displays two additional inversions, P2 and P3, in linkage with P1, and therefore originated after the introgression of P1 into the *H. numata* lineage (Jay et al. 2021).

*Heliconius numata* is widespread in the lowland and foothill tropical forests of the Amazon basin, the Guianas, and the Brazilian Atlantic Forest (Mata Atlântica), but the frequencies of the three chromosome arrangements vary across the range. Ancestral type Hn0 is fixed in the Atlantic Forest populations of Brazil (forms robigus or ethra), but segregates at intermediate frequencies in all other *H. numata* populations throughout the range (forms silvana and laura) (Fig 1C). Chromosome type Hn1 is associated with the Andean mimetic form bicoloratus and is found in the Eastern Andean foothills of Ecuador, Peru, and Bolivia. Chromosome type Hn123 is associated with a large diversity of wing-pattern forms of intermediate allelic dominance, including tarapotensis, arcuella and aurora, and is reported from Andean, lowland Amazonian and Guianese populations. Inversion polymorphism is therefore structured across the range, with populations being fixed for the ancestral chromosome (Atlantic Forest, see Text S1 & Table S1-2), or displaying a polymorphism with two (Amazon-Guiana) or three (Andes) chromosomal types in coexistence (Joron et al. 2011). Monomorphic populations of the Atlantic forest, devoid of rearrangements at the supergene locus,
might represent the ancestral state displayed by *H. numata* populations before the evolution of the supergene via introgression (Fig 1C).

The wing patterns of *H. numata* are subject to selection on their resemblance to local co-mimics (Chouteau et al. 2016), but the polymorphism is maintained by balancing selection on the chromosome types. Balancing selection is indeed mediated by disassortative mating favouring mixed-form mating (Chouteau et al. 2017) and is likely to have evolved in the response to the deleterious mutational load carried by inversions, which causes heterozygous advantage in *H. numata* (Jay et al. 2021, Faria et al. 2019, Maisonneuve et al. 2019). The introgression of P₁ and the formation of a supergene were associated with a major shift in the selection regime (Jay et al. 2018), and in the mating system was also changed during or after introgression. These events and may therefore have profoundly affected the population biology of the recipient species, *H. numata*. We investigate here whether the adaptive introgression of a balanced inversion is associated with a signature in the genetic diversity and geographic structure. In particular, we predict that genetic diversity should be higher in *H. numata* than in closely related taxa. Similarly nucleotide diversity should be higher in all polymorphic populations carrying either one segment (Hn1) or two (Hn1,Hn123) compared to the population that is monomorphic and only carries the non-inverted segment (Hn0) in the Brazilian Atlantic Forest. We analyse changes in the demographic history of the clade containing *H. numata* and closely related taxa, as well as their current patterns of diversity and demography, using three well separated populations of *H. numata* representing different states of inversion polymorphism. Our results are consistent with the selection regime and mating system associated with supergene formation having enhanced gene flow among populations and increased effective population size. Our results suggest that following supergene formation, a change in the selection regime and mating system may have facilitated gene flow among morphs and had key consequences in current patterns of genetic structure. Moreover, our findings highlight that balancing selection and a shift in mating systems associated with chromosomal polymorphism may reshape genomewide diversity, with crucial consequences on current patterns of genetic structure and population ecology.

**Material and Methods**

We used here whole genome resequencing from 137 specimens of *Heliconius*, including 68 *H. numata*. Sampling included specimens from populations in the Andean foothills (3 chromosome types), from the upper Amazon (2 chromosome types), from French Guiana (2 chromosome types) and from the Brazilian Atlantic Forests (1 chromosome type) (Fig 1C; Table S3). Related taxa were represented by the sister species *H. ismenius*, found west of the Andes (parapatric to *H. numata*), by Amazonian representatives of the lineage *H. pardalinus* (donor of the inversion), *H. elevatus*, *H. ethilla*, *H. besckei* as well as *H. hecale*, and by *H. melpomene* and *H. cydno* as outgroups. Only Andean, Amazonian and Guianese populations of *H. numata* display chromosomal polymorphism, all other taxa being fixed for the standard gene arrangement (Hn0), or for the inverted arrangement Hn1 (*H. pardalinus*) (Jay et al. 2018). Hereafter, *H. numata* populations from the Andes, Amazon and French Guiana will be collectively referred to as “Amazonian”, and populations from the Atlantic Forest as “Atlantic”. Butterfly bodies were preserved in NaCl saturated DMSO solution at 20°C and DNA was extracted using QIAGEN DNeasy blood and tissue kits according to the manufacturer’s instructions with RNase treatment. Illumina Truseq paired-end whole genome libraries were prepared and 2x100bp reads were sequenced on the Illumina HiSeq 2000 platform. Reads were mapped to the *H. melpomene* Hmel2 reference genome (Davey et al., 2016) using
Stampy (version 1.0.28; Lunter and Goodson, 2011) with default settings except for the substitution rate which was set to 0.05 to allow for the expected divergence from the reference of individuals in the so-called silvaniform clade (H. numata, H. pardalinus, H. elevatus, H. hecale, H. ismenius, H. besckei and H. ethilla). H. melpomene and H. cydno belonging to the so-called melpomene clade, their genomes were mapped with a substitution rate of 0.02. Alignment file manipulations were performed using SAMtools v0.1.3 (Li et al. 2009). After mapping, duplicate reads were excluded using the MarkDuplicates tool in Picard (v1.1125; http://broadinstitute.github.io/picard) and local indel realignment using IndelRealigner was performed with GATK (v3.5; DePristo et al. 2011). Invariant and polymorphic sites were called with GATK HaplotypeCaller, with options --min_base_quality_score 25 --min_mapping_quality_score 25 -stand_emit_conf 20 --heterozygosity 0.015.

FST, dxy and π, were calculated in overlapping windows of 25 kb based on linkage disequilibrium decay (Heliconius Genome Consortium 2012) using custom scripts provided by Simon H. Martin (https://github.com/simonhmartin), and the genome-wide average was calculated using our own scripts (available from https://github.com/angelesdecara). Distance in km between sampling sites was measured along a straight line, not taking into account potential physical barriers. Following Rousset (1997), in a 2-dimensional habitat, under a model of isolation by distance (IBD) differentiation, measured as FST/(1-FST), should increase as a function of the logarithm of the distance. Therefore, we tested for the existence and intensity of an IBD signal among species and between populations of H. numata using a linear model. If IBD is stronger in species not polymorphic for the inversion we should observed significantly steeper slopes in these species. To test this, we measured IBD (I) within populations of each species separately, (ii) for all H. numata within the Amazonian forest region but without (excluding the Atlantic forest populations) and (iii) for all H. numata including the Atlantic region. The slopes of FST/1-FST versus log(distance) were calculated using the R package lsmeans (Lenth 2016); the slope difference among species or between populations within species was estimated with an ANOVA and its significance evaluated with function pairs of this package (Text S1 and see example script on github.com/angelesdecara/).

Admixture (Alexander et al. 2009) analyses were run on a subset of the 68 H. numata genomes, keeping only 15 individuals from Peru to have a more balanced representation of individuals across the geographic distribution. Filters were applied to keep biallelic sites with minimum mean depth of 8, maximum mean depth of 200 and at most 50% genotypes missing. We only kept 1 SNP per kilobase to remove linked variants using the thinning function in vcftools, and we obtained the optimal number of clusters using cross-validation for values of K from 1 to 10 (Alexander et Lange, 2011a—2009). Principal component analyses (PCA) were performed with the same filters as for admixture, using the same H. numata genomes as for the admixture analyses, using smartpca (Patterson et al. 2006) plink2 (Chang et al. 2015).

In order to estimate demographic parameters independently of the effect of selection on diversity, we performed stringent filtering on the dataset. We removed all predicted genes and their 10,000 base-pair flanking regions, before performing G-PhoCS (Gronau et al. 2011) analyses as detailed below. Repetitive regions were masked using RepeatMasker and Tandem Repeat Finder (Benson 1999). GC islands detected with CpCluster.pl with parameters 50 and 1E-5 (Hackenberg et al., 2006) were also masked. Scaffolds carrying the supergene rearrangements (Hmel215006 to Hmel215028) were excluded, as were scaffolds from the sex chromosome (Z) and mtDNA, since
those are expected to show unusual patterns of diversity due to selection and different effective
population sizes.

We analysed the demographic history of *H. cydno, H. numata, H. ismenius, H. pardalinus* and *H. elevatus* with G-PhoCS, which allows for the joint inference of divergence time, effective
population sizes and gene flow. In order to detect differences in demography correlating with the
presence of the supergene in *H. numata*, we conducted analyses separating the Atlantic population
of *H. numata* from Amazonian populations. G-PhoCS is an inference method based on a full
coalescent isolation-with-migration model. Inferences are conditioned on a given population
phylogeny (based on Kozak et al. 2015) with migration bands (i.e. priors in the migration rates) that
describe allowed scenarios of post-divergence gene flow. The model assumes distinct migration rate
parameters associated with each pair of populations, and allows for asymmetric gene flow. Given
the computational burden of G-PhoCS, we selected two individuals per taxon or population,
retaining those with the highest sequencing depth (see Table S3). The input dataset consisted of
4092 genomic regions, each 1kb in length and spaced at approximately 30kb intervals (above the
value at which LD decay at more than half of its value) and with genotypes in at least one of the
two samples of each taxon We used as priors for coalescence times (τ) and genetic diversity (θ),
Gamma functions with α=1 and β=100, and for migration bands rates α=0.002 and β=0.00001.
These priors were chosen to allow good convergence while also ensuring non informativity. In
order to calculate the highest posterior density interval, we used the library HDInterval in R, and to
integrate such posterior densities we used the library sfsmisc in R. We rescaled the results using a
mutation rate of 1.9E-9 (Martin et al. 2016) and 4 generations per year (i.e., g=0.25). Migration
bands were considered significant following the criteria of Freedman et al. (2012): if the 95% HPD
interval did not include 0 or if the total migration rate (migration rate times the duration of the
migration band) was larger than 0.03 with posterior probability larger than 0.5.

**Demographic Reconstruction of population size changes, split and mixtures**

The G-phocs method provided useful information across all species but i) do not allow to quantify
the time scale of population size change, ii) is limited in the number of individuals it can handle and
iii) displayed limited accuracy to distinguish Ne and m in a simulation study (Gronau et al. 2011).
We thus constructed additional models to test the hypothesis that *H. numata* populations with
inversion polymorphism display an increased effective population size due to disassortative mating
To test this, we used ∂a∂i to reconstruct the demographic history of *H. numata* individuals from the
Amazonian forest, quantify their historical changes in effective population size and test their
divergence history from 1) *H. numata* from the Brazilian area, which do not carry the inversion; and
2) *H. pardalinus* individuals. We allowed for change in effective population size in both the
ancestral populations. Theoretically, the change in effective population size in *H. numata* associated
with the change in mating system should be more recent than the time of introgression of the
inversion into *H. numata*. To verify this hypothesis we allowed for change in Ne of the daughter
population at any time after the split. We tested different models of divergence with and without
(asymmetric) migration and included the effect of linked selection (e.g. Roux et al. 2016).
Since, the conditions of historical divergence are not known, we tested a model of divergence with
ongoing migration (IM) a model of divergence with ancient migration if gene-flow has stopped
recently (AM) and, in the case of divergence into multiple refugia, a model of secondary contact
(SC). We also included a model of Strict Isolation (SI) as a null model.
The models shared the following parameters: the ancestral populations of size Nanc1 can grow or
shrink to a size Nanc2 between Tanc and up until its splits at time Tsplit into two daughter populations of
size N1 and N2. Under the SI model, no gene flow occurs between the two populations. Under AM,
gene flow occurred between T_{split} and T_{am} and is followed by a period of strict isolation. Under IM, gene flow occurs at a constant rate at each generation between the two populations. Gene flow can be asymmetric, so that two independent migration rates $m_{12}$ (from population 2 to 1) and $m_{21}$ (from population 1 to 2) were modeled. Under the SC model, the population evolved in strict isolation between T_{split} and until T_{sc} where a secondary contact occurs continuously up to present time. Gene flow is modeled as $M = 2N_{REF}m$. In $\partial a_i / \partial t$, heterogeneity in effective population size was used to account for linked selection by defining two categories of loci with varying effective population sizes (proportion 1-Q of loci with a “neutral Ne” and a proportion Q of loci with a reduced effective population size due to either selection at linked site). To quantify how linked selection affects reduced $N_e$, we used a Hill-Robertson scaling factor (Hrf) to relate the effective population size of loci influenced by selection ($N_r = Hrf * N_e$) to that of neutral loci ($N_e$). A hierarchical approach was used to avoid over-fitting: first we compared models assuming constant effective population size. Second, the best identified models were modified to incorporate population expansion or decline, as expected given the observed distribution of genetic diversity. Population expansion was implemented using two additional parameters for population 1 and population 2, allowing each population to either grow or decline exponentially at any time after their split from the ancestral population (controlled by parameters $s_1$ and $s_2$ for population 1 and 2 respectively). Models were fitted using the diffusion theory implemented in $\partial a_i / \partial t$ (Gutenkunst et al. 2009) and includes the effect of linked selection. $\partial a_i$ uses the SFS as a summary of the data. For a given demographic model, the SFS is computed using diffusion approximation and compared to the empirical SFS using AIC. We used stringent filtering (GQ>30, 4 < mean depth < 80) and no missing data to keep high quality sites and remove potential paralogs or PCR duplicates exhibiting excessive read depth. To minimize linkage we subset our data to keep one SNP every 5kb. No MAF filter was used and singletons were kept to avoid ascertainment bias in estimates of demographic parameters. For each model, 32 independent replicate runs were performed and only models with the lowest AIC and ΔAIC were kept.

**Forward Simulations**

In order to better understand the nature of the processes that generate higher genetic diversity in *H. numata* compared to closely related taxa, we used simulations to test the hypothesis that disassortative mating generates an increase in levels of genetic diversity at a genome-wide scale. We hypothesized that such level of genetic diversity is higher than expected under i) random mating (a model similar to panmixia) or ii) assortative mating, as commonly observed in other *Heliconius* species. To test this hypothesis we run forward simulation under disassortative, assortative and random mating using slim v3.6 (Messer et al. 2013).

We simulated a stepping stone model with 10 demes, each composed of 1,000 diploid individuals and connected by a (symmetric) migration parameter (m). Each individual received neutral and deleterious (ratio 16:6) mutations at a rate $\mu = 1e-8 \mu/bp/generation$ (rescaled to $\mu = 1e-6$ for faster simulation of a larger population). We simulated an individual with a pair of 1Mb chromosome, including a single locus with 5 alleles with perfect dominance (allele 1 > allele 2 > allele 3 > allele 4 > allele 5) given 5 possible alternative phenotypes (referred hereafter as “morph”). Each allele was fully linked (no recombination) with a given deleterious recessive mutation, generating overdominance at this loci so that polymorphism is always maintained. Local adaptation was introduced in the model through a single parameter defining randomly which morphs were favored in each population. In each population, either 2 or 3 morphs benefited from a fitness advantage compared to the others. The fitness reduction varied between 0 (= fitness of zero for migrants in a demes) and 1 (no reduction of fitness). We tested 3 possible values for this parameter (0, 0.25 and 0.5).
Finally, disassortative mating was controlled by a mate choice parameter defining whether a morph would reproduce with another morph. The strength of the parameter varied between 0 (= complete disassortative mating) and 1 (= no mating weight). We tested 3 possible values for this parameter (0, 0.25 and 0.5).

We run the model for 80,000 generations to reach demographic equilibrium and assessed levels of synonymous diversity ($\pi$). We tested all combinations of the 3 values for levels of disassortative mating and local adaptation and ran 10 replicates per combination in order to estimate the variance around $\pi$.

Similarly, we run a model with strict assortative mating, controlled by a parameter defining whether similar morphs reproduced together. The strength of the parameter varied between 0 (complete assortative mating where a given individual mate only with an identical morph) and 1 (where individual mate randomly with regards to the morph). We tested 3 possible values for this parameter (0, 0.25 and 0.5). As for disassortative mating, all combinations of assortative mating and local adaptation values were tested. For each model we tested 3 values for the migration rate, $m = 1e^{-4}$, $1e^{-6}$ and $1e^{-8}$, resulting in a total of 54 comparisons.

For graphical display in Figure 4, the values of assortative/disassortative mating were rescaled on a scale between (0 and 1) with 0 indicating no disassortative mating but complete assortative mating and 1 complete disassortative mating (or no assoartive mating). A value of 0.5 was equivalent to random mating.

Results

Using cross validation error as a measure of the optimal number of clusters with Admixture, we found that $K=2$ was the optimal cluster number describing within-species genetic variation in $H. numata$ (Fig 2A). One cluster corresponds to the Atlantic population, forming a well-differentiated genetic entity compared to all other $H. numata$ populations. All Amazonian populations of $H. numata$ showed a remarkable uniformity, with the exception of a few individuals sharing some variation with SE Brazil. This pattern is consistent with the population structure inferred using microsatellite markers (Fig S1). Population structure revealed by PCA is in line with the admixture analysis (Fig 2B). Individuals from the Atlantic populations of $H. numata$ clustered together to one side of the first PCA axis, whereas all other individuals from all other populations clustered to the other side. The second axis of the PCA separates individuals from French Guiana from the other samples of the upper Amazon. This clustering separation was not found with Admixture (i.e. with $K=3$) from the complete dataset. To better investigate the existence of a hierarchical population structure, we excluded individuals from the Atlantic populations and compared individuals from French Guiana to a randomly sampled set of Peruvian individuals. In this case we found a clear separation in two groups corresponding to French Guiana and Peru (Fig S2A). The same pattern was observed when replacing Peru by Colombia or Ecuador (Fig S2B,C), suggesting that the divergence between Amazonian populations is very reduced. In accordance, pairwise genome-wide estimates of differentiation ($F_{ST}$) between $H. numata$ populations showed elevated values when comparing the Atlantic population to other populations, low values between French Guiana and other Amazonian population, and were the lower but very small values when comparing pairs of Amazonian populations outside of French Guiana, even at a large distance (Fig 2C, Table S4). For instance, the population from La Merced in Peru shows an $F_{ST}=0.032$ with the population from French Guiana at a distance of 3019km, but an $F_{ST}=0.311$ (an order of magnitude higher) with the Atlantic population at a similar distance. The comparison between La Merced and Ecuador was even lower ($F_{ST} = 0.0159$). Isolation by distance among Amazonian populations of $H. numata$,
estimated using the proxy $F_{ST}/1-F_{ST} \sim \log_{10}(\text{km})$ was significant ($R^2 = 0.41$, $p = 1.61e-06$, slope = 40.02). Comparison among other species did not reveal any significant IBD ($R^2 = 0.01$, $p = 0.29$, slope = 0.12), showing a very different pattern to other species, with a highly significantly shall. An analysis of the slope revealed a lower rate of increase in $F_{ST}$ with distance in H. numata compared to all other taxa (Fig 2C, Table S4, Supp Text S1). By contrast, differentiation as a function of distance(IBD) between Atlantic and Amazonian populations of H. numata is close to what is observed in other species, and not significantly different (see Supp. Text S1).

Analyses of genetic diversity show that all populations of H. numata, except those from the Atlantic Forest, have a similarly high genetic diversity (Fig 3A). By comparison, closely related Heliconius taxa show significantly lower genetic diversity (Fig 3A). These patterns are similar to those obtained using G-PhoCS to analyse the demographic histories in a phylogenetic context, where Amazonian populations of H. numata show higher population sizes compared to the Atlantic populations (Fig 3B, Table S5). G-PhoCS analyses also show a demographic history in which gene flow plays a crucial role (Table S6). For instance, our analyses show strongly significant gene flow right at the beginning of the divergence between H. ismenius and the other silvaniforms, as well as in the divergence between H. pardalinus and H. elevatus. The effective population sizes inferred from Atlantic genomes are one order of magnitude lower than those obtained using H. numata populations from other localities (Fig 3A and Table S5). In our cladogram, the increase in H. numata population size is restricted to the Amazonian branch, excluding Atlantic populations.

Demographic reconstruction from $\partial a/\partial i$

The model selection procedure based on AIC gave higher support for a model of secondary contact (SC) in the pairwise comparison between H. numata from Peru and H. numata from Brazil. The pairwise comparison between H. numata and H. pardalinus supported a model of divergence with continuous gene-flow (IM) (Table S7, Figure S3). All models supported an expansion occurring in the ancestral population, followed by further growth of the H. numata carrying the inversion supergene to reach a size of several millions, which was by far the largest effective size compared to all other species. This stands in stark contrast with the results observed in the samples from Brazil (which do not harbor the inversion) (Table 2). Accordingly, H. numata populations from the Atlantic forests of Brazil appear to have been subject to a bottleneck at the start of their divergence from Amazonian populations, followed by exponential growth, suggesting a strong (and recent) founding event, leading to a comparatively smaller population size than that observed in the rest of H. numata. It is worth noting however that effective population size was hard to estimate in pairwise comparisons between H. numata from Peru and SE Brazil. Indeed, parameter uncertainty was large, and model residuals (Figure S3) were also large. Our results indicated that H. pardalinus displayed an initially large population size followed by a comparatively smaller size expansion than H. numata (Table 2). Estimates of current effective population sizes are therefore qualitatively similar to those from G-phocs.

Forward simulations

Forward simulations under different levels of local adaptation (controlled by the strength of divergent selection), disassortative mating and migration are displayed in Figure 4B. The same results under a model of assortative mating involving different levels of selection and migration are displayed in Figure 4A. Overall, synonymous genome-wide nucleotide diversity ($\pi_S$) was higher in 73 % of the models including disassortative mating (average $\pi_S = 0.0145$) when compared to their
equivalent under assortative mating (average πS = 0.011), a weak but significant difference (p < 0.01, see Figure S5). In summary, modest differences were observed among models with different strengths of divergent selection or disassortative mating, the most influential variable being the rate of migration (Figure 4).

**Discussion**

Our results suggest that populations displaying inversion polymorphism in the P supergene in *H. numata* also display distinctive population demography and gene flow. Differences in demographic and differentiation regimes associated with structural variation at this locus are revealed when comparing polymorphic populations of *H. numata* to closely-related monomorphic taxa, such as (1) peripheral populations of *H. numata*, (2) sister taxa, and (3) inferred ancestral lineages. This suggests that the existence of a mimicry supergene controlling polymorphism in *H. numata* is may be associated, in time and in space, with major differences in population biology. We hypothesize this to be due to a change in the balancing selection regime due to heterozygote advantage (Jay et al. 2021) and in the associated evolution of disassortative mating (Chouteau et al. 2017) following the onset of inversion polymorphism, causing direct effects on ecological parameters such as gene flow, immigration success and effective population size. Testing this hypothesis through forward simulation yielded mixed evidence for a genome-wide effect of this disassortative mating, especially when compared to a simple model of random mating.

Our analyses show large-scale variation in genetic diversity among closely related taxa in this clade of *Heliconius* butterflies. Within *H. numata*, the genetic diversity of found in polymorphic Amazonian populations is ~4 times one to two orders of magnitude higher than the diversity found in populations from the Atlantic Forest. Generally, Amazonian populations of *H. numata* harbour the highest genetic diversity in the entire melpomene/silvaniform clade, which contrasts with the low diversity found in the most closely related taxa such as *H. ismenius* or *H. besckei*. Inferring historical demography during the diversification of the *H. numata* lineage reveals that the large effective population size in that species is only associated with the branch representing polymorphic, Amazonian *H. numata* populations, while internal branches all show very low diversity estimates. This suggests that ancestral and putatively monomorphic populations of *H. numata* were similar in their diversity parameters to current sister species *H. ismenius* populations, or to current peripheral Atlantic *H. numata* populations. Although low-diversity lineages could have lost diversity due to recent events such as strong bottlenecks, the distribution of parameters across as estimates of effective population size lineages rather from dadi indicated for the Atlantic population, suggests Nevertheless our dadi estimates do suggest that the Amazonian populations of *H. numata* underwent a dramatic increase in effective population size posterior to their split with Central American (*H. ismenius*), *H. pardalinus* and the Atlantic populations. Those findings are in agreement with G-Phocs analyses. The Amazonian branch of the *H. numata* radiation is characterized by the long-term maintenance of inversion polymorphism, triggered by the introgression of a chromosomal inversion about 2.2 Ma ago. Therefore, the major shift in demography between Amazonian and Atlantic populations indeed appears associated to coincide, at least in the broad sense, with the occurrence of inversion polymorphism, even though the lack of replication of this event impedes firmly establishing causality here.

Another striking result is the lack of genetic structure displayed by *H. numata* across the Amazon, with all Amazonian and Guianese populations forming a single genetic cluster. Only
Atlantic populations stand out and display high differentiation with other *H. numata* from the rest of the range. French Guiana and Peruvian populations, separated by over 3000 km across the Amazon, are remarkably genetically weakly genetically differentiated compared to pairs of populations at comparable distances in other species, and show only modestly stronger similar differentiation than pairs of *H. numata* populations taken at short distances. *H. numata* populations from the Amazon show significantly lower isolation by distance than all other taxa, as measured by the change in F\textsubscript{ST} across distance (F\textsubscript{ST}/km) (Fig. 2C), with a very distinctive, flat slope of isolation by distance. The only exception is found when comparing Amazonian populations with Atlantic populations of Brazil, displaying a level of differentiation in line with that of pairs of populations at similar distances within other taxa.

Effective population size is affected by census size, mating system, and the force and type of selection acting on traits (Charlesworth 2009). Selection is often viewed as a force only affecting the genetic variation around specific, functional loci in the genome, but it may also affect whole genome diversity, for instance when its action is sufficient to modify local demography or mating patterns. In *H. numata*, morphs and therefore inversion genotypes show disassortative mate preferences, i.e., they preferentially mate with individuals carrying different chromosome types (Chouteau et al. 2017). Disassortative mating enhances heterozygosity and the mating success of individuals expressing rare alleles (negative frequency dependence) (Knoppien 1985; Hedrick et al. 2018). Consequently, immigrants expressing rare, recessive alleles have a mating advantage in *H. numata*. Disassortative mating associated with the supergene should therefore bring an advantage to immigrant genomes in LD with recessive supergene alleles, possibly enhancing genome-wide gene flow. Supergenes are also characterised by monosingle-locus Mendelian inheritance, by which mimicry phenotypes are maintained in the face of recombination, even after immigration. This effective migration regime in populations harbouring a mimicry supergene is therefore likely to be quite different to that observed in other mimetic taxa such as *Heliconius melpomene* or *H. erato*, in which mimicry variation is controlled by multiple loci with diverse dominance patterns. In those taxa, hybrid offspring display recombinant patterns breaking down mimicry, even after multiple generations of backcrossing, and pure forms mate assortatively with respect to wing pattern (McMillan et al. 1997, Mallet et al. 1998, Jiggins et al. 2001); both processes select against mimetic variants migrating from adjacent areas with distinct warning patterns. The expectation is that immigrant genomes should be consistently associated with mimicry breakdown in the case of multilocus architectures, which should translate into an effect on effective migration genomewide, compared to situations with polymorphic mimicry supergenes. In *H. numata*, the evolution of a polymorphic mimicry supergene and disassortative mate preferences could therefore explain the relative lack, compared to other *Heliconius* taxa, of differentiation among polymorphic populations, even across large distances. Furthermore, enhanced gene flow could also cause an increase in effective population size estimates (Slatkin 1987), putatively explaining why polymorphic populations of *H. numata* harbour the highest genetic diversity, and display the highest *Ne* estimates in the entire *melpomene*-silvaniform clade of *Heliconius*. These hypotheses are also supported by our forward simulation which suggests that indeed, disassortative mating resulted in enhanced genetic diversity compared with assortative mating, although the effect was small. In addition, our results also suggest that a simple model of random mating may explain well the data, thus purely demographic expansions may also generate high genetic diversity and high effective population size, as observed from our \( \partial \alpha \partial \iota \) demographic modelling.
Alternative processes may of course contribute to the observed patterns. Amazonian and Atlantic populations may differ in other aspects that could also result in differences in genetic diversity. Habitat availability and structure may be different, possibly entailing differences in the maintenance of diversity. The Atlantic Forest is vast in area, but may represent a smaller biome compared to the Amazon. It is isolated from the bulk of the range of *H. numata*, which could result in a population's ecology—displaying characteristics of peripheral populations with smaller effective population sizes (Eckert et al. 2008). Reduced effective population size is supported by our data. One major caveat associated to our inference remains the small number of individuals (n = 12) from the Atlantic forest. Genetic diversity might be underestimated, notably if populations have a history of fragmentation in this area. The other *Heliconius* species in the clade have much in common with *H. numata* in terms of habitat and general ecology, yet their niche and life-history specificities and their phylogenetic histories may result in consistent differences with the polymorphic *H. numata* populations. All those specificities may contribute to the observed pattern in which polymorphic Amazonian populations of *H. numata* display high effective population size and a lack of weak geographic structure in genome-wide genetic variation. Yet this pattern of variation correlates parsimoniously with the evolution of a supergene causing disassortative mating and single-locus control of mimicry variation—in certain—Amazonian *H. numata* populations, which provides an elegant mechanism explaining their differences with extant and ancestral closely-related lineages. However, we cannot rule out a role for conjectural differences in ecology and geography with all other taxa.

In conclusion, our results show a remarkable contrast in the demography and differentiation of populations within the Amazonian range of *H. numata* compared to closely related taxa and ancestral lineages, as well as with other taxa in the *melpomene/silvaniform* clade. Although those populations may differ in many uncharacterized ways from all other taxa, one known and consistent difference is the maintenance of inversion polymorphism associated with a specific mating system and selection regime in Amazonian *H. numata*—Theis distinctiveness of the onlyis widely polymorphic populations species in the clade is consistent with the hypothesis that the evolution of a supergene maintained by balancing selection represents a major transition in this lineage, triggering changes in genome-wide patterns of diversity and population ecology over the last 2 million years since its formation. If this hypothesis is correct, the evolution of a locus under balancing selection may therefore feed-back on population ecology and diversification, and consequently on speciation.

Eco-evolutionary feedbacks between changes in genomic architecture and the ecological parameters of populations are still not well understood and few cases have been studied. The evolution of self-incompatibility loci in plants, affecting the rules of mating and feeding back on population ecology, connectivity, and demography, may be one example, but effects of the evolution of trait genetic architecture on population ecology may be more common than previously thought. In our study, more work on the determinants of variation in effective population sizes in the genus *Heliconius* is needed to determine the precise impact of the supergene on demography in *H. numata*. More work on the determinants of variation in effective population sizes in the *Heliconius* genus is needed to determine the precise impact of the supergene on demography of *H. numata*. We believe that our results emphasize a potential link between genomic architecture, selection and demography, and should inspire future theoretical and modelling studies. Finally, the eco-evolutionary feedbacks between changes in genomic architecture and the ecological parameters of populations are well
known when considering self-incompatibility loci in plants, but may be more common than previously thought. Indeed, Overall our result suggests that balancing selection maintaining structural polymorphisms affecting life-history traits may have a profound influence on species ecology.

Contributions:
MARdC, PJ, QR and MJ designed the study and wrote the manuscript. BH, AVLF, TTT, RRR, KLSB provided the Atlantic samples. CS provided the Colombian samples. MARdC and PJ and QR performed genomic analyses and simulations with input from AW. MARdC, PJ, MJ, FPP and MC collected the Peruvian and Ecuatorian samples. MC performed microsatellite analyses and organized fieldworks and butterfly rearing. All authors contributed to editing the manuscript.

Acknowledgements:
This work was funded by grants HYBEVOL-Hybevol (ANR-12-JSV7-0005) and Supergene (ANR-18-CE02-0019-01) from the Agence Nationale de la Recherche and European Research Council Grant MimEvol (StG-243179). We acknowledge the Genotoul and the Montpellier Bioinformatics Biodiversity (MBB) platforms for providing us with calculation time. We thank Dr. Vitor Becker, at the Serra Bonita Reserve (Bahia), Alexandre Soares, at the MN/UFRJ (Rio de Janeiro) and Dr. Marcelo Duarte at the MZ/USP (Sao Paulo) for their contribution to the collection of butterflies in Brazil. Field collections in Colombia were conducted under permit no. 530 issued by the Autoridad Nacional de Licencias Ambientales (ANLA). We are grateful to Marianne Elias and Violaine Llaurens for comments and discussions. AVLF acknowledges support from Fundação de Amparo à Pesquisa do Estado de São Paulo – (FAPESP) (Biotas-Fapesp grants 2011/50225-3, 2013/50297-0) and Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq) (421248/2017-3 and 304291/2020-0). KLSB acknowledges the financial support of FAPESP Process # 2012/16266-7. Brazilian specimens are registered under SISGEN (A701768).

Data availability:
The raw sequence data were deposited in NCBI SRA and accession numbers are indicated in Supplementary table 3.

References


Figure 1 | Genetic and population structure at the P supergene.

A. Schematic phylogeny of the sampled species. It includes all members of the silvaniform clade and two outgroups, *H. melpomene* and *H. cydno*.

B. Schematic description of the genetic structure of the P supergene. Three chromosomal arrangements coexist in *H. numata* and are associated with different morphs.

C. Origin of *H. numata* specimens used for analyses and distribution of chromosome arrangements across the neotropics. Numbers in brackets indicate sampled specimens in each locality (the Tarapoto population lumps several neighbouring subsamples on the map).
Figure 2 | *H. numata* is characterised by low population structure.

A. Admixture plot for *H. numata*. The optimal cluster number for *H. numata* is two, and it splits *H. numata* into two categories, whereas they come from Atlantic forest or the Amazon. BR=Brazil (Atlantic), PR=Peru, VE=Venezuela, CO=Colombia, EC=Ecuador, FG=French Guiana. B. Principal component analysis computed on whole genome SNP. Color code match those given in the asmixture label of panel A. C. Relationship between genetic differentiation (Fst/(1-Fst)) and logarithm of geographical distance. Fst is measured between morphs/populations of the same species. *H. numata* populations from the Amazon show low isolation by distance when compared to related species.
Figure 3 | Variation in present and past effective population size in Heliconius species

A. Variation in Pi in several Heliconius populations, showing higher genetic diversity in H. numata populations from the Amazon than other taxa. Population names indicate their origin as in Figure 2 (e.g. PR=Peru), with the addition of PA=Panama. The H. numata population with the lowest diversity is the one from the Atlantic forest (Brazil).

B. Schematic representation of Gphocs results showing that Amazonian populations of H. numata, which have the P supergene, show a dramatic increase in population size posterior to their split with the Atlantic populations of Brazil, which lack the supergene.
**Figure 2** | *H. numata* is characterised by low population structure.

A. Admixture plot for *H. numata*. The optimal cluster number for *H. numata* is two, and it splits *H. numata* into two categories, whereas they come from Atlantic forest or the Amazon. BR=Brazil (Atlantic), PR=Peru, VE=Venezuela, CO=Colombia, EC=Ecuador, FG=French Guiana. B. Principal component analysis computed on whole genome SNP. C. Relationship between genetic differentiation (Fst) and geographical distance. Fst is measured between morphs/populations of the same species. *H. numata* populations from the Amazon show low isolation by distance when compared to related species.
Figure 3 | Variation in present and past effective population size in Heliconius species

A. Variation in Pi in several Heliconius populations, showing higher genetic diversity in H. numata populations from the Amazon than other taxa. Population names indicates their origin as in Figure 2 (e.g. PR—Peru), with the addition of PA—Panama. The H. numata population with a lowest diversity is the one from the Atlantic forest (Brazil). B. Schematic representation of Gyphos results (presented in table S5-6). Gene flow was modelled but not represented graphically for clarity. Showing that Amazonian populations of H. numata, which have the P supergene, show a dramatic increase in population size posterior to their split with the Atlantic populations of Brazil, which lack the supergene.
Figure 4 | Weak but significant differences in synonymous nucleotide diversity ($\pi_S$) emerged at a genome-wide scale under divergent selection and mating region. Results from forward simulations of 10 populations undergoing local adaptation and different mating strategy are presented. Shown are levels of synonymous diversity obtained under assortative (A) versus disassortative mating (B) under different rates of migration and different local adaptation fitness. Each combination of parameters in brackets display the (dis)-assortative mating weight and the fitness value for local adaptation respectively. A left value of 0 in the bracket means complete assortative mating and 0.5 means no assortative mating or disassortative mating. Value of 1 means complete disassortative mating. A right value of 0 in the bracket mean fitness of 0 for non locally adapted individuals in a demes, A value of 0.5 means a reduced fitness of 0.5 relative to the maximum value.
List of Supplementary Materials:

Table S1-6
Fig S1-2
Text S1