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Genomics, Population Divergence, and Historical Demography of the World's Largest and Endangered Butterfly, The Queen Alexandra's Birdwing

Eliette L. Reboud (¹,*, Benoit Nabholz (¹,², Emmanuelle Chevalier , Marie-ka Tilak , Darren Bito , and Fabien L. Condamine (¹,*

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Abstract

The world's largest butterfly is the microendemic Papua New Guinean *Ornithoptera alexandrae*. Despite years of conservation efforts to protect its habitat and breed this up-to-28-cm butterfly, this species still figures as endangered in the IUCN Red List and is only known from two allopatric populations occupying a total of only ~140 km². Here we aim at assembling reference genomes for this species to investigate its genomic diversity, historical demography and determine whether the population is structured, which could provide guidance for conservation programs attempting to (inter)breed the two populations. Using a combination of long and short DNA reads and RNA sequencing, we assembled six reference genomes of the tribe Troidini, with four annotated genomes of *O. alexandrae* and two genomes of related species *Ornithoptera priamus* and *Troides oblongomaculatus*. We estimated the genomic diversity of the three species, and we proposed scenarios for the historical population demography using two polymorphism-based methods taking into account the characteristics of low-polymorphic invertebrates. Indeed, chromosome-scale assemblies reveal very low levels of nuclear heterozygosity across Troidini, which appears to be exceptionally low for *O. alexandrae* (lower than 0.01%). Demographic analyses demonstrate low and steadily declining *Ne* throughout *O. alexandrae* history, with a divergence into two distinct populations about 10,000 years ago. These results suggest that *O. alexandrae* distribution has been microendemic for a long time. It should also make local conservation programs aware of the genomic divergence of the two populations, which should not be ignored if any attempt is made to cross the two populations.

Key words: Conservation genomics, heterozygosity, low genetic diversity, *Ornithoptera alexandrae*, reference genome.

Significance

Despite its charisma, little is known about the demographic trends and taxonomic status of the two populations from the giant endangered birdwing butterfly *Ornithoptera alexandrae*. By sampling and sequencing individuals of this species and two closely related species, we study whether and how the population is structured, and we investigate the genomic diversity of the species and the "health" of their genomes and populations (e.g., demographic trend, evidence of inbreeding). Overall, the very low genomic diversity and steadily declining trend inferred by this study suggest that efforts need to be reinforced to conserve this amazing Papua New Guinean insect.

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Introduction

When in January 1906, Alfred S. Meek saw an enormous butterfly flying high above him in the canopy of this forest of the Northern Province of Papua New Guinea, two days walk from the coast, he took his rifle and shot down the beast. This is what one can read in the letter he sent to his correspondent Karl Jordan at the Natural History Museum of Tring (England) (letter no 155 of Meek's communications, Meek 1906; Ackery 1997; Tennent 2021). Meek let his funder, Lord Walter Rothschild describe in 1907 for the first time Ornithoptera alexandrae (Papilionidae: Troidini), known as the Queen Alexandra's birdwing butterfly, based on this female whose wing still bears the stigma of this extraordinary hunt. This "trophy" and the even larger congeners that followed have become representatives of the world's largest known butterfly species to date and contribute to the continuing amazement of scientists at the incredible diversity, size, and beauty of Papua New Guinea's insects (Parsons 1992; Mitchell et al. 2016). Indeed, many naturalists have been studying O. alexandrae, culminating with the comprehensive review of this butterfly by Mitchell et al. (2016) that serves as a basis for this work.

As the world's largest butterfly, *O. alexandrae* can measure up to 28–30 cm in wingspan (Mitchell et al. 2016 and references therein). *Ornithoptera alexandrae* is endemic to the Northern Province of Papua New Guinea, in a narrow range around Popondetta (Northern Province; fig. 1). Long-term field observations in the last decades have

shown there are two recognized allopatric populations: a lowland population in Popondetta plains (≤300 m above sea level), and a highland population occurring on the relatively inaccessible Managalas Plateau about 800 m above sea level (Collins and Morris 1985; Parsons 1999; Böhm 2018). A mountain range separates the two populations, bounded in the West by Mount Lamington volcano (1,700 m), and eastward to a mountain 2,140 m high. According to available data, there have been no sightings in between. The volcanic activity of Mount Lamington (last eruption in 1951 with an activity until 1956; Global Volcanism Program 2022), the flooding, drought, and fires occurring in the region as well as recent logging and agricultural activities might explain today's fragmented distribution of O. alexandrae (Parsons 1992; Mitchell et al. 2016). The relatively small distribution range composed of two patches has been interpreted by Haugum and Low (1979) as a relict occurrence, potentially due to an evolutionary bottleneck or demographic decline. However, genetic studies are crucially lacking to assess this hypothesis.

Ornithoptera alexandrae is considered a threatened species in the IUCN Red List (Böhm 2018). While this species is very rare over an area of occurrence of 8,710 km², its actual area of occupancy is only 128–140 km² fragmented into two populations, placing it in the endangered category (Böhm 2018). There are doubts about the previously thought monophagy of its caterpillar due to the lack of comprehensive morphological or genetic studies on the

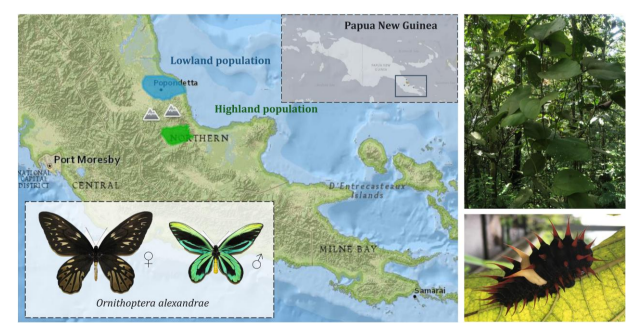


Fig. 1.—Left, the distribution range of *Ornithoptera alexandrae*, lowland population in blue and highland population in green. Topright, *Aristolochia cf. meridionaliana* plant in its environment. Bottomright, *O. alexandrae* larva. Photos: Fabien L. Condamine. Papua New Guinea gray map is from the IUCN Red List https://www.iucnredlist.org/ (Böhm 2018), and *O. alexandrae* distribution map has been designed on MapMaker https://mapmakerclassic.nationalgeographic.org based on Mitchell et al. (2016).

Southeast Asian genus Aristolochia (Parsons 1996; Buchwalder et al. 2014; Mitchell et al. 2016). It has long been thought that there was no particular restrictions due to host plant distribution to explain the local occurrence of O. alexandrae as there are many areas where the main larval food plants (previously thought to be Aristolochia dielsiana, possibly being A. meridionaliana in the highlands and A. alexandriana in the lowlands) grow prolifically. This pattern has been described in some monophagous lepidopteran species (Quinn et al. 1997) and may highlight the fact that the distribution of O. alexandrae species is also driven by other factors of a microclimatic, pedologic, or geological nature that might limit its distribution. On the other hand, some factors represent a threat or vulnerability for the species: it is a large species with a relatively specialized ecology (larvae on a single or very few host-plants and adults in a single habitat) (Koh et al. 2004, Palash et al. 2022), it is sympatric to other Ornithoptera species (Koh et al. 2004), and its habitat is or has been fragmented by fires, droughts, and volcanic eruptions, and is severely affected by agriculture (Parsons 1992; Mitchell et al. 2016) and other human activities, with habitat conversion leading to a local decline in larval host vines. Accordingly, O. alexandrae is placed at the top of the CITES list (Appendix I). However, although its international trade is prohibited, this species is highly prized and is subject to an illegal trade that is dangerous for the demography of the species, and its survival (Mitchell et al. 2016). Thus, better monitoring of this species is recommended by the IUCN to better track population trajectories (Böhm 2018), particularly because its numbers and the current trend in population dynamics are unknown.

The two allopatric populations of O. alexandrae are externally morphologically similar but express important biological differences, such as slight differences in size (on average 14% larger in highlands) and development time (on average 34.5% longer in highlands) (Straatman 1971; Mitchell et al. 2016). The soil and host plant eaten by the larvae might also differ in the two populations (Haugum and Low 1979; Mitchell et al. 2016), so it is unclear how divergent these two populations are. In fact, genomic research on other butterfly groups has revealed that superficial similarity in adults can hide a previously unrecognized cryptic lineage (e.g., Hebert et al. 2004; Burns et al. 2008). Knowledge of evolutionary and genetic history of the species and populations could help conservation and breeding programs to save the species. Genomics is considered a powerful tool for studying the past and present structure and diversity of populations and brings an invaluable source of information, especially for species that are naturally rare and difficult to study (e.g., Westbury et al. 2018; Van der Valk et al. 2020; Morin et al. 2021). Genome sequencing is increasingly recognized as an important contribution to identifying, characterizing, and conserving biodiversity (Formenti et al. 2022). Reference genomes provide primary data for understanding historical demography (Morin et al. 2021), gene and trait evolution (Warren et al. 2021), or even susceptibility to inbreeding depression and accumulation of deleterious mutations (Chattopadhyay et al. 2019; Van der Valk et al. 2020; Robinson et al. 2022). Genomic resources are also useful for broader studies of population structure, relatedness, and recovery potential (e.g., Garner et al. 2016; Morin et al. 2018; Tunstall et al. 2018), or for assessing correlations between current IUCN status and past demography (Nadachowska-Brzyska et al. 2015). These types of estimates (e.g., sequentially Markovian coalescent [SMC] methods, Li and Durbin 2011) have been widely used for conservation purposes for vertebrates, such as mammals (Morin et al. 2021) or birds (Nadachowska-Brzyska et al. 2015) and insect pests (Hazzouri et al. 2020; You et al. 2020). Despite the continuous increase of threatened insects (Sánchez-Bayo and Wyckhuys 2019), it has been much less used for insect conservation (but see Mikheyev et al. 2017; Podsiadlowski et al. 2021). It is indeed challenging to study the demography of invertebrates using polymorphism-based methods because the risk of violating the assumptions of SMC-type models is high. For instance, Sellinger et al. (2021) revealed that these methods of inference perform poorly when the ratio between the recombination and mutation rates is important, therefore highlighting that the consideration of this ratio is crucial and still much too little considered in this type of analyses in the literature.

Here we perform a first genomic study of O. alexandrae to understand the past and present demography of this species and to bring insights into its endangered status, which may have implications for conservation strategies. Since a local conservation program has been set up and is ongoing to rear the two populations, the taxonomic status of these two populations (i.e., populations or species) may inform conservation management of this threatened species (Mitchell et al. 2016). If the two populations are too divergent, it could be complicated to breed specimens from the Managalas Plateau with specimens from the Popondetta lowlands. Given the above-mentioned biological features of this butterfly species, we aim to 1) assemble high-quality and annotated whole genomes for the two populations; 2) assess the level of nuclear heterozygosity, 3) estimate the demographic history of the species and the two populations in relation to past environmental change and to test whether the current range of the species is relictual; and 4) provide information for the policy makers to improve their conservation strategy.

Results and Discussion

High-Quality Assemblies for Birdwing Butterflies

We collected live specimens from the two populations of *O. alexandrae* with one adult and one caterpillar per population and sequenced the DNA combining a mean of

72.5 x of long reads (LR) (Oxford Nanopore) for draft assembly, 82 x of short reads (SR) (Illumina) for polishing, as well as 51.7 million cleaned RNAseq reads (8.4 Gb) for genome annotation (see Materials and Methods). Using Flye assembler (Kolmogorov et al. 2019) and Pilon polisher (Walker et al. 2014), we assembled the four genomes of O. alexandrae that range from 321 Mb to 327 Mb. They are very contiguous with a mean of 582 contigs (ranging from 305 to 1,222 contigs) and a mean N50 of 9.9 Mb (table 1). Over a total of 5,286 core genes of the Lepidoptera database (odb10, Manni et al. 2021), BUSCO recovered on average $98.83\% \pm 0.05$ complete genes, $0.23\% \pm 0.05$ fragmented genes and $0.97\% \pm 0.06$ missing genes (table 1). The genome size and gene completeness of our O. alexandrae assemblies are comparable to the genome of a related species: Troides helena (330 Mb, BUSCO score = 96.6%), which was assembled with similar data and methods (He et al. 2022). Furthermore, the genome size stands among the smallest within the family Papilionidae but still is 30% to 40% larger than some Papilio (the sister tribe of Troidini; Liu et al. 2020; He et al. 2022) illustrating the dynamic genome size evolution of the family.

After removing potential exogenous contigs of the assemblies (see Materials and Methods), we selected FC563 as the reference genome for further analyses, as it had the best assembly statistics and we found no evidence of contamination (i.e., bacteria, plants). We assessed the quality of this O. alexandrae FC563 assembly by looking at its correspondence with the reference genome of Papilio bianor (chromosome-level assembly, Lu et al. 2019). We found 24 contigs that match with more than 70% of the length of P. bianor chromosomes out of 30 chromosomes. This represents a cumulative length of 81% of the total assembly. Among those, 18 contigs have a single reciprocal match with one *P. bianor* chromosome (representing a cumulative length of 48% of the assembly). One of the most fragmented chromosomes is the Z chromosome in which 11 contigs of O. alexandrae FC563 assembly match chromosome 30 (Z) of *P. bianor*. This is not surprising as FC563 is a female and, therefore, has half the coverage on the Z compared to the autosome. However, a similar analysis performed on the male FC560 led to eight contigs matching chromosome 30 (Z) of *P. bianor*, suggesting that chromosome Z is difficult to assemble. These eight contigs linked to the Z chromosome were independently identified using coverage and heterozygosity information in FC563, representing a cumulative length of 14.1 Mb (supplementary table S1, Supplementary Material online). The FC563 assembly is therefore composed of 24 full-length or nearly full-length chromosomes, including 18 full-length chromosomes (supplementary fig. S1, Supplementary Material online). Analysis of the chromosome-level synteny between O. alexandrae and P. bianor shows a high level of genomic synteny

Assembly Statistics for the Genomes of *Ornithoptera alexandrae*, *Ornithoptera priamus*, and *Troides oblongomaculatus*

Table 1

		Ornithopter	Ornithoptera alexandrae		Ornithoptera	Troides
	High	Highland	Low	Lowland	priamus	oblongomaculatus
	FC560 (♂)	FC561 (♀)	FC562 (♀)	FC563 (♀)	FC565 (♂)	FC569 (♀)
Raw data sequenced (Gb) (LR + SR)	30.98 + 33.2	31.82 + 31.0	41.18+33.8	29.44+37.6	38.99 + 36.9	38.34 + 57.4
Final mean coverage	66×+84×	59×+ 79×	95×+70×	70×+ 95×	×96 +×98	87×+ 137×
(LR + SR)						
Assembly size (bp)	326,746,405	325,591,695	320,565,121	321,134,305	316,500,205	348,219,047
Number of contigs	333	1,222	465	305	862	465
N50 (bp)	10,703,829	7,761,365	9,990,636	11,239,331	4,951,043	5,909,187
Max length (bp)	20,033,826	12,853,001	14,559,367	13,869,660	10,532,357	13,649,974
Nucleotide assembly BUSCO score (%)	S:98.6; D:0.2;	S:98.6; D:0.2;	S:98.7; D:0.2;	S:98.6; D:0.2;	S:98.4; D:0.3;	S:98.5; D:0.2;
	F:0.3; M:0.9	F:0.2; M:1.0	F:0.2; M:0.9	F:0.2; M:1.0	F:0.3; M:1.0	F:0.3; M:1.0
Protein-coding genes (number and mean size)	17,617	17,449	16,159	16,508	I	ı
	6,095bp	6,082bp	6,473bp	6,325bp		

long reads; SR, short reads; bp, base pairs. For BUSCO scores, S, single-copy genes; D, duplicated genes; F, fragmented genes; and M, missing genes out of 5,286 genes in odb10 lepidopteran database. Annotation was only done for O. alexandrae individuals 뭈

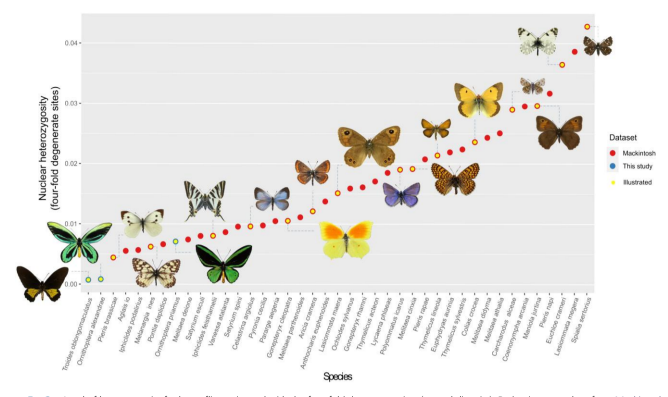


Fig. 2.—Level of heterozygosity for butterflies estimated with the four-fold degenerate sites (neutral diversity). Red points are values from Mackintosh et al. (2019) and blue points are from this study. The yellow point indicates illustrated species. Photos (not at scale): *T. oblongomaculatus* (CC BY 4.0, Peter Wing); *O. alexandrae* (Fabien L. Condamine); *P. brassicae* and *G. cleopatra* (CC BY-SA 3.0, Sarefo); *M. ines* (CC BY-SA 4.0, Atylotus); *O. priamus* (Eliette L. Reboud); *I. feisthamelii* and *M. cinxia* (CC BY-SA 4.0, Didier Descouens); *C. argiolus* (CC BY 3.0, Alan Cassidy); *A. cramera* and *P. icarus* (Robin Noel); *L. maera* and *C. crocea* (CC BY-SA 3.0, Vítězslav Maňák); *T. lineola* and *S. sertorius* (CC BY-NC-SA, Peter Huemer); *C. alceae* (CC BY-SA 3.0, Didier Descouens); *M. jurtina* (Public domain, Pekka Malinen); *E. crameri* (Alexander Slutsky).

(supplementary fig. S1, Supplementary Material online). Our results suggest that the combination of Nanopore LR and Illumina SR perform notably well to recover chromosome-scale assemblies as we reconstructed genome assemblies comparable to the best assemblies of Papilionidae available so far (comparison with *P. bianor*, Lu et al. 2019: table 1) without relying on Hi-C techniques.

Using transcriptomic data, protein homology, and de novo genes prediction, we annotated the genomes of the two populations (FC560 and FC653) using the MAKER pipeline (Holt and Yandell 2011). Protein predictions retrieved 17,617 genes for FC560 and 16,508 genes for FC563. We carried out BUSCO analyses with these two proteoms and estimated 97.7% complete genes, 0.9% fragmented genes and 1.4% missing genes for FC560, and 97.2% complete genes, 1.0% fragmented genes, and 1.8% missing genes for FC563. Because FC560 annotation contained more genes, it was transferred to the other two genomes of *O. alexandrae*. Overall, 33–35% of the genome was annotated as repeat sequences with mostly unclassified categories of interspersed repeats (supplementary table S2, Supplementary Material online). This proportion of repeats

is relatively high compared to other Papilionidae genomes already available (22% in Papilio glaucus, 22.4% in Papilio xuthus; Cong et al. 2015; Lu et al. 2019), except for the species with larger genomes such as P. bianor (55%; Lu et al. 2019) and Parnassius apollo (65%; Podsiadlowski et al. 2021). This is consistent with a positive correlation between assembly size and repeats content in insects (Petersen et al. 2019; Heckenhauer et al. 2022; Sproul et al. 2022). Within O. alexandrae, all individuals have similar cumulative repeat size, and no difference was detected between lowland and highland individuals despite the fact that highland individuals had a slightly larger genome assembly than lowland individuals (~5 Mb, supplementary table S2, Supplementary Material online). These annotated genomes are available in GenBank (BioProject PRJNA938052).

For comparison purposes, we also sequenced and assembled two other birdwing butterfly species (Troidini): *Ornithoptera priamus poseidon* and *T. oblongomaculatus papuensis*, both ranked as Least Concern in the IUCN Red List (see *Materials and Methods*). Both genomes were very similar to the *O. alexandrae*'s ones in quality and assembly statistics (table 1).



Area of occupancy: <140 km² Neutral diversity: 0.08155 Forewing length: 98 ~ 126 mm

Larval diet: monophagous? (Aristolochia alexandriana,

A. merdionaliana)

Area of occupancy: $>1,208 \text{ km}^2$ Neutral diversity: 0.7081Forewing length: $74 \sim 118 \text{ mm}$

Larval diet: oligophagous (*A. acuminata*, *A. alexandriana*, *A. meridionaliana*, *A. praevenosa*, *A. deltantha*, *A. indica*,...)

Area of occupancy: >156 km², likely underestimated

Neutral diversity: 0.07035Forewing length : $74 \sim 92$ mm

Larval diet: oligophagous (A. acuminata, A. momandul,

A. gaudichaudi)

Fig. 3.—Comparison of *O. alexandrae*, *O. priamus*, and *T. oblongomaculatus*. Distribution maps, IUCN status, and areas of occupancy are from IUCN Red List https://www.iucnredlist.org/(Böhm 2018). Neutral diversity of *O. alexandrae* is the mean neutral diversity values of the four individuals. Forewing length is from Nakae (2021) and its larval diet is from Mitchell et al. (2016) and Böhm (2018). Larval diets should be taken with caution due to uncertainties in the *Aristolochia* taxonomy. Host plant species in bold font are known to be the main diet of the species. Photos (male specimens, relatively scaled to the mean species forewing value): *O. alexandrae* (Fabien L. Condamine), *O. priamus* (Eliette L. Reboud), and *T. oblongomaculatus* (CC BY 4.0, Peter Wing NHM specimen).

The World's Largest Butterfly Has Low Levels of Genomic Diversity

Using SR data and annotation with MitoFinder (Allio et al. 2020) (see Materials and Methods), each mitogenome was reconstructed in a single contig and we retrieved 36-37 genes (including 13 protein-coding genes and 2 rRNA), with evidence of circularization. The mitogenomic diversity (π -diversity) including coding and noncoding region (such as the D-loop) was calculated at ~0.0704% (supplementary table S3, Supplementary Material online). This is comparable to the low level of mitochondrial diversity for mammalian species such as the Tasmanian devil or Island fox (Westbury et al. 2018). Up to our knowledge, there is no estimation of mitochondrial diversity based on whole mitogenomes of butterflies, but there are studies estimating mitochondrial diversity relying on the cytochrome c oxidase subunit I (COI) DNA barcode marker (e.g., π -diversity: Mackintosh et al. 2019; haplotype diversity: Dincă et al. 2021). The mitochondrial diversity of the DNA barcode for O. alexandrae is ~0.076% (supplementary table S3, Supplementary Material online), which ranks fifth lowest of 38 European butterflies (Mackintosh et al. 2019). However, low mitochondrial diversity does not equate to low autosomal diversity (Allio et al. 2017; Mackintosh et al. 2019).

Using mapping data on the best genome assembly of O. alexandrae FC563, we calculated the autosomal

heterozygosity of the two populations. All four individuals of O. alexandrae show a heterozygosity of around 0.08% (supplementary table S4, Supplementary Material online). Using a similar metric relying on four-fold degenerate sites from coding sequences, this heterozygosity is the lowest value observed within the butterfly studied by Mackintosh et al. (2019) (fig. 2, supplementary table S4, Supplementary Material online). We also performed sensitivity analyses to corroborate these results using different data types (SR vs. LR) and window sizes (100 kb vs. 1 Mb) and the proportion of missing data (10% vs. 20%). Overall, the heterozygosity of O. alexandrae ranges from 0.0745% to 0.0838% (supplementary table S4, Supplementary Material online), suggesting the low estimated heterozygosity is not an artifact due to methods and/or data. Determinants of heterozygosity are still a longterm debate (Romiguier et al. 2014; Ellegren and Galtier 2016; Mackintosh et al. 2019; Buffalo 2021) and intrinsic features of O. alexandrae like its large size, its tropicality, or its microendemism might explain such low heterozygosity. However, for two closely related species having similar lifestyle but larger distributional range and abundance than O. alexandrae (fig. 3), we found that O. priamus has a higher heterozygosity rate (autosomal 0.433%, neutral diversity 0.708%, ca. six and ten times higher) and that T. oblongomaculatus has an even lower heterozygosity than

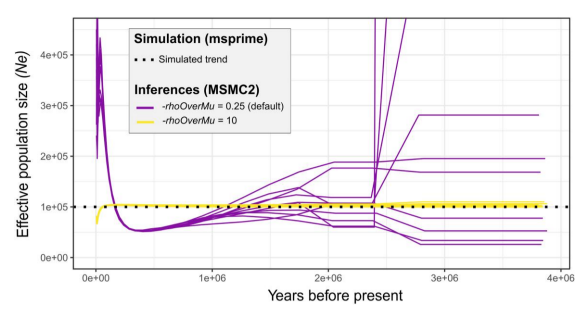


Fig. 4.—Performance of MSMC2 inferences on simulated data. Dotted line represents the simulated scenario produced by msprime (Stable Ne = 1e5, r = 1e-8, $\mu = 1.319e-9$). Colored lines represent the demography inferred by MSMC2 on this data with different *-rhoOverMu* settings (ten repetitions each).

O. alexandrae (0.0737% autosomal, 0.0704% neutral). Interestingly, these levels of heterozygosity are much lower than the level estimated in the genus *Papilio* ranging from 1.0% to 2.3% although estimated with a different method (k-mer distribution; Lu et al. 2019). Accordingly, the low level of heterozygosity cannot be explained by the species' range or body size, as suggested by Mackintosh et al. (2019) (fig. 3).

Historical Demography

Although the causes of the low genetic diversity of these birdwing butterflies are unclear, we can wonder how this translates into demographic history of the species given the threats they are experiencing. Predictions can be formulated such as a prolonged decline of effective population sizes such as inferred for the brown hyena and the Californian condor (Westbury et al. 2018; Robinson et al. 2021) or a low but stable effective population size such as in the vaquita porpoise (Morin et al. 2021; Robinson et al. 2022). To our knowledge, there are still few examples of demographic history in insects, other than pest insects (e.g., red-palm weevil, Hazzouri et al. 2020; diamondback moth, You et al. 2020; but see Walton et al. 2021; Manthey et al. 2022; García-Berro et al. 2023). Within swallowtail butterflies, the recent study of the Apollo butterfly (Parnassius apollo), ranked as a threatened species, showed using SNP data synchronous population declines throughout different mountain massifs despite high heterozygosity levels (Kebaïli et al. 2022).

Sequential Markovian coalescent models (e.g., PSMC, multiple sequentially Markovian coalescent [MSMC]) are

widely used to study the trajectory of the ancestral effective population size (Ne) over time. We applied the MSMC2 model to the best genome assembly of O. alexandrae as well as O. priamus and T. oblongomaculatus before focusing on the two populations of O. alexandrae. SMC models generally infer a Ne under the panmictic assumption. This strong assumption is often false in reality, and theoretical work and simulations have shown that SMC dynamics might also be caused by population structure and connectivity changes (Teixeira et al. 2021), so this should be borne in mind when interpreting these population size analyses (Teixeira et al. 2021; Bentley and Armstrong 2022). Furthermore, it has been shown that SMC methods do not perform well when the mutation rate μ is rarer than the recombination rate r (Sellinger et al. 2021). Unfortunately, this is likely to be very common in many groups such as fungi, sea cucumbers, nematodes, corals, insects, and plants (see Wilfert et al. 2007; Stapley et al. 2017 for recombination rates and Lynch et al. 2016; Wang and Obbard 2023 for mutation rates). In practice, there is concern that at least many small nonvertebrate genomes have a much lower μ than r (Sellinger et al. 2020; Sellinger et al. 2021) whose demographic inferences would be affected and poorly addressed in the literature (e.g., coral: Fuller et al. 2020; insects: Waldvogel et al. 2018; Hazzouri et al. 2020; You et al. 2020; Wang et al. 2022; Garcia-Berro et al. 2023).

Demographic inferences on simulated data with the range of recombination and mutation rate parameters of O. alexandrae (i.e., r about ten times higher than μ , see Materials and Methods) confirmed the methodological issue, which was almost completely reduced when the model

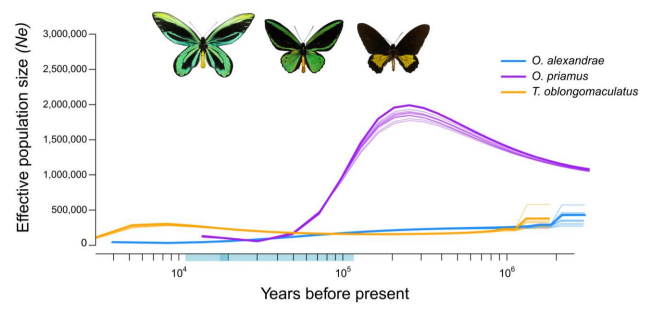


Fig. 5.—Estimated historical demography of three birdwing butterfly species. MSMC2 estimates of the effective population size (Ne) with -rhoOverMu = 10, for O. alexandrae (blue), O. priamus (purple), and T. oblongomaculatus (orange). Bootstraps are represented in clear lines. The pale blue rectangle along the time bar indicates the limits of the last glaciation period (11,700–115,000 years ago) with the last glacial maximum in darker blue (19–20,000 years ago). The recent present (last 4,000 years) is not represented.

was set with an appropriate initial value of the ratio (fig. 4). We do not know exactly how the model behaves with an appropriate initial value on real data, but our results changed significantly when the initial value was adjusted (supplementary fig. S2, Supplementary Material online), highlighting that this is probably a very important parameter to consider in SMC analyses and further emphasizing the importance of taking the results of any SMC analyses with caution.

Finally, MSMC2 analyses traced relatively different demographic histories for the three butterflies (fig. 5). The effective population size of O. alexandrae seems to have been at a low but continuously declining number from ~250,000 to 50,000 individuals during the last million years. However, we inferred a recent demographic change in the last 10,000 years (see below). This demography dynamic is similar to that of the vaquita porpoise, whose effective population size has never been elevated but remained stable for much of its known history and now has one of the lowest rates of heterozygosity known among marine mammals (Morin et al. 2021; Robinson et al. 2022). Altogether, these results suggest that the ancestral effective population of O. alexandrae has never been large, thus suggesting that it has been microendemic for a very long time. Interestingly, T. oblongomaculatus experienced a similar fairly low and regular Ne during most of its history with a Ne that culminated at ~0.3 million individuals around 10,000 years ago (fig. 5). It was followed by a decreasing trend toward the present, which seems consistent with the low rate of heterozygosity that we estimated today. In contrast, *O. priamus* shows an early peak of Ne (~2.0 million individuals ~250,000 years ago) followed by a severe decline until ~40,000 years ago.

In the literature, temporal variations of Ne are usually compared with past climatic fluctuations such as temperature and/or sea level, in particular in line with Quaternary glaciations (e.g., Nadachowska-Brzyska et al. 2015; Westbury et al. 2018; Hazzouri et al. 2020; Morin et al. 2021; Teixeira et al. 2021). Although it is tempting to associate the inferred Ne variations of the studied birdwing butterflies with the glaciation-interglaciation cycles, it remains difficult to extract a correlation because of climatic heterogeneity at the global scale and estimates of demographic parameters (Bentley and Armstrong 2022). The last million years was mostly a glacial period that has also been documented in New Guinea (e.g., Chappell and Polach 1991; Barrows et al. 2011), punctuated by intermittent warming events (e.g., Eemian event ~115,000-130,000), with the last glacial period (Würm glaciation event) starting ~115,000 and ending ~11,700 years ago (last glacial maximum at 19,000–20,000 years). During the last glacial maximum, temperatures were 5 °C colder (Barrows et al. 2011), and sea level was 70 m lower (Chappell and Polach 1991) than today. The last glaciation period coincides with the decrease of Ne for O. priamus. On the contrary, the scale induced by O. priamus' inference does not allow to discern any congruence between climatic changes and the Ne of T. oblongomaculatus and O.

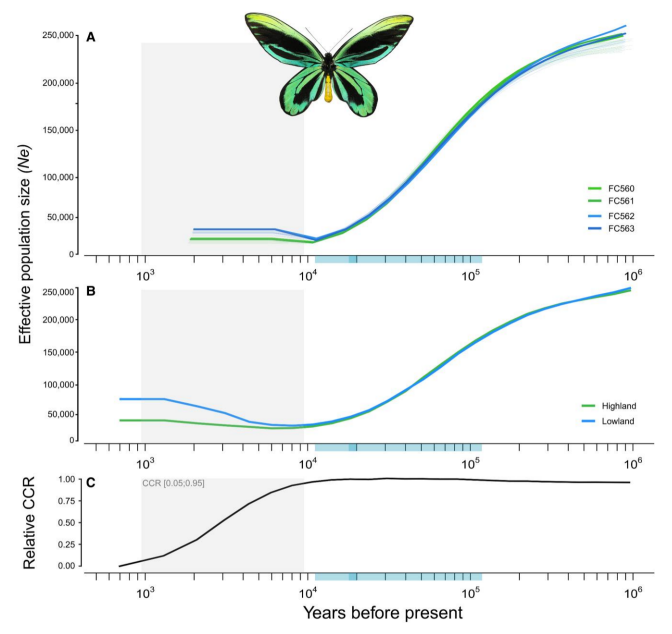


Fig. 6.—Demographic inferences of *O. alexandrae* populations. *A.* MSMC estimates for each individual of *O. alexandrae*, with bootstraps represented in clear lines. *B.* MSMC estimates for both populations of *O. alexandrae*. *C.* Relative cross coalescent rate (CCR) is estimated with the MSMC analysis, representing the time interval of population divergence. The gray zone represents the 5–95% CCR ([947; 9460] years ago), and has been reported on each plot. The pale blue rectangle along the time bar indicates the limits of the last glaciation period (11,700–115,000 years ago) with the last glacial maximum in darker blue (19–20,000 years ago). For every graph, the recent present (last 700 years) is not represented.

alexandrae. In addition, this result relies only on a single genome for *O. alexandrae* whereas the two populations of *O. alexandrae* might have had different demographic histories.

Within *O. alexandrae*, every individual showed a similar demographic pattern (low and steadily declining *Ne* during most of the history, fig. 6a). Nonetheless, the two populations of *O. alexandrae* seem to separate initially ~10,000 years ago, and have likely been followed by a relative

increase of the Ne in the last period (after the split). These results were validated when using multiple genomes per population (see Materials and Methods) (fig. 6b) and computed the cross coalescence rate (CCR) between populations (fig. 6c). The CCR gives the probability that a coalescence happens between rather than within the populations and, therefore, quantifies isolation (Wang et al. 2020). Using a generation time of 0.75 years, the CCR started to decrease \sim 10,000 years ago (CCR = 0.95) and

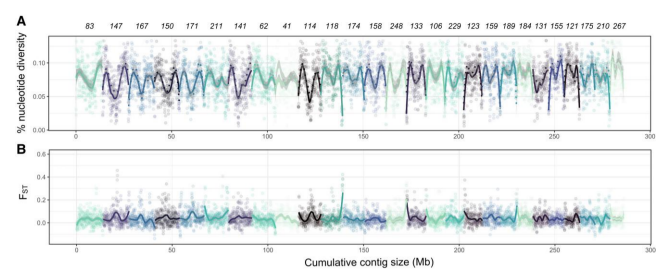


Fig. 7.—Genome-wide genetic diversity and differentiation within *Omithoptera alexandrae*. A. Percentage of nucleotide diversity of the genome of O. alexandrae for all individuals. B. Genetic differentiation (F_{ST}) between the highland and lowland populations. Nucleotide diversity and F_{ST} were computed on nonsex-related contigs >500 kb in length, over sliding windows of 100 kb with an overlap of 50 kb. Alternance of colors indicates the different chromosome-level contigs. Contig names are indicated at the top in italics.

was below 5% at \sim 1,000 years ago (CCR = 0.05) (fig. 6c). These results suggest that two populations have recently diverged, potentially just after the last glaciation, raising the question of the extent to which the two populations are fully isolated or whether some gene flow (migration) is still ongoing.

Population Structure

To explore the recent divergence between the two populations, we computed the median F_{ST} on sliding windows of 100 kb across the whole genome (see Materials and Methods). We estimated a median F_{ST} of 0.03 but the F_{ST} was highly variable among genomic positions with top 5% windows having F_{ST} above 0.14 (fig. 7). The F_{ST} and nucleotide diversity clearly decrease in the center of the large contigs in regions (fig. 7) as expected along chromosomes (e.g., Tine et al. 2014; Delmore et al. 2018). This large-scale variation in F_{ST} and genetic diversity, repeated among scaffolds, suggests the action of endogenous effects such as recombination rate variation, rather than the effect of exogenous local adaptation (Burri 2017). This is consistent with reduced recombination rate in centers and extremities of the chromosomes as recently reported in the painted lady (Vanessa cardui; Shipilina et al. 2022). In addition, the cumulative size of 100-kb windows with less than two heterozygous sites represents less than 1% of the genome in all individuals. Moreover, not a single window of 1 Mb with one heterozygous site or less was detected in any individuals. It suggests that the genome of *O. alexandrae* shows no sign of recent inbreeding.

Using the approximate Bayesian computation (ABC) framework implemented in DILS (Demographic Inferences

with Linked Selection) (Fraïsse et al. 2021), we fitted the four models (secondary contact [SC], strict isolation [SI], isolation with migration [IM], ancient migration [AM]) across randomly selected genomic regions to estimate population sizes, migration rates and time of population separation according to each model (i.e., the date when gene flow stopped) between lowland and highland populations (see Materials and Methods). On the four runs, DILS only set aside the SC model, but could not identify a best-fitting model between the three others: (two runs supported migration (IM model), and two runs found isolation (one run for AM and one run for SI). Depending on the best-fitted model, the time of split was either more recent than MSMC (isolation) or similar (migration) and varied from 5,100 generations (~3,825 years ago) to 16,850 generations (~12,637 years ago) for the median of the posterior distribution of the SI model and the IM model respectively (Supplementary table S5, Supplementary Material online).

Despite the uncertainty of the DILS scenarios, notably on the presence or intensity of migratory flows between the two populations, all the analyses seem overall to agree on the rather consistent isolation of the two populations for a few thousand years. This is interesting because field observations tend to say that the two populations are externally morphologically similar but express biological differences. The highland population is on average larger than the lowland population (264 vs. 218 mm wingspan on average; Mitchell et al. 2016). Furthermore, *O. alexandrae* populations may feed on two different *Aristolochia* species, which could be *A. meridionaliana* in the highlands and *A. alexandriana* in the lowlands (Haugum and Low 1979; Parsons 1996; Mitchell et al. 2016). However, it is difficult to reach a

conclusion on this point since little is known on the taxonomy of these two plant morphs and, as mentioned by Buchwalder et al. (2014) who started a taxonomic revision of the genus *Aristolochia*, there is still significant work to be done to update the nomenclature of this group. The two populations of *O. alexandrae* also differ slightly in their life cycle. From the egg, through larval and pupal stages to adults, the life cycle of *O. alexandrae* takes about 131 days (74 days as larva) in the lowland, while it takes 200 days (125 days as larva) in the highland of the Managalas Plateau (Straatman 1971; Mitchell et al. 2016). Such a difference may be explained by a temperature gradient between the low and highlands (up to 4 °C cooler than the plain).

Overall, our results indicate a lack of evidence of a strong genomic differentiation and a genomic barrier to gene flow between the two allopatric populations of *O. alexandrae* that could justify the description of two distinct species. Nevertheless, given the morpho-ecological differences and the fact that there is a modest population structure (F_{ST}) and divergence time probably initiated after the Last Glacial Maximum, it is possible that the populations deserve subspecies status. However, determining whether the observed morpho-ecological differences are fixed in the genome and to what extent the genitalia differ would shed more light on this proposal, although the latter is difficult to satisfy due to the lack of available individuals to dissect for this threatened and protected species.

Conservation Implications

Conservation today is often approached as a triage, leaving out some "unsalvageable" species in an attempt, at best, to save others (Hayward and Castley 2018). As discussed by Wiedenfeld et al. (2021), this approach to the sixth mass extinction should be primarily a matter of prioritizing the allocation of economic and material resources to the rescue of any species, and the current system is failing to allocate enough resources to save biodiversity. The choice to help certain species rather than others is risky since this choice is necessarily based on partial knowledge and limited data, such as their apparent rarity, sometimes interpreted as an abrupt decline, of certain species. However, it seems that comprehensive and meticulous works of some studies such as the one carried out on the kakapo (Dussex et al. 2021), Island fox (Robinson et al. 2018), or the vaguita porpoise (Morin et al. 2021, Robinson et al. 2022) in recent years are a further argument that species are not necessarily doomed to extinction due to their rarity and low numbers (e.g., Robinson et al. 2018; Wiedenfeld et al. 2021; Robinson et al. 2022). On the contrary, these rare species may have been rare for so long that this may have led them for instance to have purged more deleterious recessive alleles, they might not suffer from inbreeding depression, and their low number may be less of a problem in recovering their population than many other species that would appear to us less endangered at first sight. Protection measures are proposed and sometimes implemented as a result of scientific studies revealing the urgency of acting to save species. Most of these measures concern vertebrates (e.g., large mammals), often omitting threatened insect diversity from conservation policies (but see Mikheyev et al. 2017). As Harvey et al. (2020) advocate in their roadmap for insect conservation, there is a range of actions to be taken, from short-term actions such as drastically or immediately stopping threats, to medium- and long-term actions such as conducting new research to fill the knowledge gap regarding insect decline or launching sustainable funding initiatives with the aim of restoring, protecting and creating new vital habitats for insects.

Ornithoptera alexandrae is one of the most charismatic invertebrate species and is receiving special attention as a flagship species for invertebrate conservation (Mitchell et al. 2016). The biological characteristics of O. alexandrae correspond to a wide range of traits identified as vulnerability factors, notably its large size, its host, and environment specificity, the vulnerability of its host plant, or the presence of interspecific competition in its distribution range (Koh et al. 2004; Mattila et al. 2006). On the other hand, the relatively similar past evolutionary patterns and dynamics of O. alexandrae with the vaquita porpoise, showing a low heterozygosity and a low Ne for a long time, suggest that we can have hope about the capacity of O. alexandrae to avoid extinction. However, the species experienced declining trends in the past and these overall results should not be an argument to relax the protection measures. On the contrary, we take the opportunity to encourage the authorities and associations for increasing protection actions, especially since we know very little about the health of the butterfly populations for the last 4,000 years, while the destruction of its habitat has clearly not decreased since. Logging opens up previously inaccessible areas, resulting in changes in O. alexandrae habitat by increasing the number and extent of clearings of primary and secondary forests (Mitchell et al. 2016). Today, oil palm exploitation likely constitutes threats through planned or ongoing deforestation, as well as the increased human population on the Popondetta plain associated with this activity (Mitchell et al. 2016). Given the difficulty to access the Managalas plateau, the anthropic pressure is today higher in lowlands, suggesting that the lowland population is experiencing more threats. Therefore, we urge policy makers to 1) protect the rainforest environment of O. alexandrae, especially in the highlands where a specific protection area could be exclusively dedicated to it, and 2) strongly support the lowland population through the ongoing rearing program with breeding and release in the wild. Both proposals are discussed and analyzed in detail by Mitchell et al. (2016) and we fully agree with their statement on the urgency of action to protect O. alexandrae and

how to address the problem. Note that we advise avoiding interbreeding between the two populations as there is an ongoing process of natural evolution leading to genetic divergence between the two populations.

An interesting research perspective would be to perform a temporal sampling of genomic data from museum specimens. This can provide a more accurate approach to quantify genetic threats in endangered species and to estimate recent decreases in genome-wide diversity, increases in inbreeding levels, and accumulation of deleterious genetic variation (Díez-del-Molino et al. 2018) like it has been done on kākāpō (Dussex et al. 2021) and the vaquita porpoise (Robinson et al. 2022).

Materials and Methods

Sampling, DNA and RNA Extractions, and Sequencing

With permits, a total of four individuals of *O. alexandrae* were collected in Papua New Guinea (Oro Province) in November 2019 by F.L.C. and D.B. Two specimens per population were collected: one female and one caterpillar for the lowland population (near Popondetta), and one male and one caterpillar for the highland population in the Managalas Plateau (near Kawowoki village). For each specimen, head, thorax, and abdomen have been separated with scalpels, crushed with surgical scissors, and conserved separately in RNAlater, then stored in freezers at –20 °C after a few days at ambient temperature (and one day at 4 °C as a transition). One individual of *O. priamus* and one individual of *T. oblongomaculatus* have been collected under similar conditions during the same mission.

Tissues from the thorax or abdomen were used to extract high molecular weight DNA. As part of tests that ended up being similar in terms of quality of sequencing, we used two different extraction methods. The first was the phenolchloroform method, including a specific ratio of 0.8 x AMPure beads applied to retain the longest DNA fragments (Tilak et al. 2020). The second method was the use of the Qiagen genomic DNA kit. This second solution was ultimately applied to most samples due to a better 260/230 ratio in Nanodrop assays, as DNA purity is essential for long-read sequencing, especially for Oxford Nanopore Technology (ONT) sequencing. In addition, one of the four samples of FC561 was treated with the Short Read Eliminator Kit XS (Circulomics, PacBio, USA) to discard sequences below 10 kb long. Final DNA purity and concentrations were measured using Nanodrop (Thermo Fisher, USA) and Qubit (Thermo Fisher, USA). RNA was extracted for O. alexandrae only. Extraction and purification were conducted with the Qiagen RNeasy kit. We used part of the thorax of caterpillars (FC561/FC563) and part of the abdomen for adults (FC560/FC562) which were crushed in the lysis buffer.

Library Preparations and Sequencing

Whole-genome libraries were constructed using the resulting high molecular weight DNA as input for the Nanopore LSK-109 ligation kit (Oxford Nanopore Technologies, UK) following the manufacturer's protocol. Long-read sequencing was performed on a GridlON device with two to four R9.4.1 flow cells, depending on the individuals (supplementary table S6, Supplementary Material online).

Remaining DNA extractions of each individual were sent to Novogene Europe (Cambridge, UK) for library preparations. Libraries were generated using NEBNext DNA Library Prep Kit following manufacturer's recommendations and indices were added to each sample. Genomic DNA was randomly fragmented to a size of 350 bp by shearing, then DNA fragments were end-polished, A-tailed, and ligated with the NEBNext adapter for Illumina sequencing, and further PCR enriched by P5 and indexed P7 oligos. The PCR products were purified (AMPure XP system) and the resulting libraries were analyzed for size distribution by Agilent 2100 BioAnalyzer and quantified using real-time PCR. Since the genome sizes for the Troidini species were estimated to be about 320 Mb (Ornithoptera) and 340 Mb (Troides), Illumina 150 bp paired-end sequencing was run on a NovaSeq 6000 instrument to obtain about 32 and 34 Gb per sample corresponding to a genome depth-of-coverage of about 100x.

The quality and quantity of all RNAs were checked using Nanodrop, Qubit, and 1.0% agarose gel electrophoresis and sent to Novogene for library preparations. Messenger RNA was purified from total RNA using poly-T oligoattached magnetic beads. After fragmentation, the first strand cDNA was synthesized using random hexamer primers followed by the second strand cDNA synthesis. The library was ready after end repair, A-tailing, adapter ligation, size selection, amplification, and purification. The library was checked with Qubit and real-time PCR for quantification and BioAnalyzer for size distribution detection. Quantified libraries have been pooled and sequenced on Illumina platforms, according to effective library concentration to a data amount of about 8 Gb per sample.

Assembly of Reference Genomes

For GridION sequencing, all fast5 files were base called using Guppy 5.0.15 (developed by ONT) using super-high accuracy mode and quality control of 10 (min_score 10). Sequencing adapters were trimmed using Porechop 0.2.3 (https://github.com/rrwick/Porechop). Draft genome assemblies were performed with Flye 2.8.3 (Kolmogorov et al. 2019) with default options. Illumina reads were cleaned, filtered, and paired with fastp 20.0 (Chen et al. 2018) using default options. Paired-end sequences were mapped on the Flye assembly using BWA 0.7.17 (Li 2013). Resulting SAM files were converted to sorted

indexed BAM files with SAMtools (Li et al. 2009). Flye draft assemblies were polished with two rounds of Pilon 1.24 (Walker et al. 2014) using this mapping information. Assembly statistics were then assessed using the gVolante2 platform (Nishimura et al. 2017) to retrieve the number and size of contigs, the presence, completeness, and duplication of BUSCO genes of the Lepidoptera odb10 database (Manni et al. 2021). More information and statistics about quality of sequencing, assembly, and polishing are displayed in supplementary table S6, Supplementary Material online. Before submitting genomes assemblies to GenBank, we used BlobTools 1.1.1 (Laetsch and Blaxter 2017) set to the NCBI and diamond databases to check for possible contaminations. We found no evidence of artificial contamination coming from laboratory manipulation, but some contigs were clearly identified as belonging to exogenous organisms such as host plants and symbionts. We removed all contigs that were belonging to the plant (subsequently identified as Aristolochia by BLAST), Microsporidia (unicellular fungal insect parasites), or Pseudomonadota (Wolbachia, Enterobacter) phylum. The FC563 assembly showed no evidence of contamination, while 83% of the total contigs removed were from FC561 (supplementary table S7, Supplementary Material online). We used the scaffold function of RagTag 2.1.0 (https://github.com/malonge/RagTag; Alonge et al. 2019; Alonge et al. 2021) to find the correspondence between P. bianor chromosomes and O. alexandrae FC563 assembly.

Genome Annotation

We performed a full pipeline of annotations for the individuals FC560 and FC563. The pipeline was composed of the six following steps. First, we reconstructed the repeat sequences using RepeatModeler 2.0.1 (Flynn et al. 2020). The consensus sequences generated by RepeatModeler were blasted against the "reference transcriptome" database of UniProt (download in October 2021, https://www. uniprot.org/) using diamond blastx (Buchfink et al. 2015), and we excluded all the proteins that were not associated to repeat sequences from the consensus sequences. We then annotated the repeat sequences in the respective assemblies using RepeatMasker (Smit et al. 2013–2015) using both the Dfam libraries (Storer et al. 2021) setting the parameter "-species" on "Arthropoda" and the newly identified repeat sequences reconstructed using RepeatModeler. Second, we assembled the RNA-seg data by cleaning the reads with fastp 20.0 (Chen et al. 2018), mapped the read using HISAT2 (Kim et al. 2019) onto the reference genome, and we annotated the cDNA using StringTie (Pertea et al. 2015) producing a GTF file. The cDNA sequences were converted in fasta using the "gtf_genome_to_cdna_fasta.pl" script of TransDecoder (https://github.com/ TransDecoder/TransDecoder). The RNA-seq data of all the individuals were mapped against the reference genome of FC563. Third, we ran MAKER 2.31.11 (Holt and Yandell 2011) using the information of the annotated repeat sequences and the cDNA sequences provided as "rm_gff" and "est" options in the control file of MAKER, respectively. We also used the proteins sequences of Heliconius melpomene, Melitaea cinxia, Papilio machaon, P. xuthus, P. glaucus provided as "protein" option in the control file of MAKER to help identify genes using homology information. Fourth, SNAP (Korf et al. 2004) and AUGUSTUS (Stanke et al. 2006) were used to produce gene prediction models from the first round of MAKER. BUSCO 5 (Simão et al. 2015) with options "-long" and "-augustus" and the Endopterygota database was used to produce the gene prediction model of AUGUSTUS. Fifth, we ran again MAKER using the annotation from the first round and the gene models of SNAP and AUGUSTUS. Sixth, steps 3 and 4 were repeated using the second round of MAKER annotation to produce a third and final round of annotations. Finally, for individuals FC561 and FC562, we used Liftoff 1.6.3 (Shumate and Salzberg 2021) to map the annotation of FC560 on the two other assemblies.

Mitogenomic Diversity

Long reads of every individual were corrected with SR using LoRDEC 0.9 (Salmela and Rivals 2014). For each individual of O. alexandrae, the corrected LR were mapped with Minimap2 2.17 (Li 2018) on the reference mitogenome of O. richmondia from a previous study (NC_037869.1, Condamine et al. 2018). The reads that mapped with the references were filtered by quality via SAMtools (Li et al. 2009) ("view -q 30"). For each individual, a subset of reads from 3.6 Mb to 6.3 Mb was created so that mitogenomes would have an expected depth of coverage between $200 \times$ and $400 \times$. We used Flye 2.8.3 to assemble the mitogenomes and the resulting assemblies were given to MitoFinder 1.4 (Allio et al. 2020) to annotate (gene, tRNA, rRNA) and extract genes (gene, rRNA). The cleaned SR data were also directly given to MitoFinder to produce annotated mitogenomes based on the SR data only. As these mitogenomes were of slightly better quality (better annotation, presence of the complete or nearly complete D-loop), they were those submitted to GenBank (Accession numbers OQ59006-OQ59009). A nucleotide alignment was produced with MAFFT 7.310 (Katoh and Standley 2013) after having manually adjusted the sequences due to circularization. We used Seaview 4.7 (Gouy et al. 2010) to visualize the four whole mitochondrial genome alignment and to count pairwise differences using "Statistics" of Seaview, and the mean of this pairwise distance was calculated, and ultimately divided by the alignment length (supplementary table S3, Supplementary

Material online). The same steps were carried out with the COI mitochondrial gene only.

Nuclear Heterozygosity of Troidini

For O. alexandrae, we selected FC563 as the reference assembly, as it has the highest N50, mean coverage, and BUSCO score, the lowest number of contigs, and has no contamination (table 1). Genomes of O. priamus and T. oblongomaculatus were their own reference. The corrected reads (LoRDEC, see Mitogenomic diversity section) of every individual were mapped on their reference genome using "-a" option of Minimap2 2.17 (Li 2018). We used SAMtools to compress, sort, and index these mappings. SNP calling was performed with Longshot 0.4.1 (Edge and Bansal 2019), using a threshold of 15 x minimum and 150 x maximum for the depth of coverage (minimal quality of 20; default quality) and applying a transition/transversion rate for genotype prior estimation (ts_tv_ratio) of 2.0 (Edge and Bansal 2019). SNPs with a quality below 200 were excluded. All positions, SNPs and homozygous, must be contained within the coverage thresholds, otherwise, they were considered ambiguous. As the quality of phasing may be important for population genomics and demographic analyses, we checked the average size of phasing from Longshot. The average length of the phased blocks is 435 kb, and a haplotype N50 of 1.7 Mb (supplementary table S8, Supplementary Material online for phasing statistics per individual). In addition, to ensure that our heterozygosity estimates did not depend on the data and method (Bentley and Armstrong 2022), we also calculated the heterozygosity rate based on the SR data to evaluate the robustness of our results. Illumina cleaned reads were mapped on references using the speedseq pipeline (Chiang et al. 2015) that relies on BWA 0.7.17 (Li 2013). We excluded the socalled discordant and splitter reads and the reads with mapping quality below 30. Genotype calling was performed using FreeBayes 1.3.1 (Garrison and Marth 2012) set with the "-use-best-n-alleles 4" option, and the same coverage threshold as for LR data. SNPs with quality scores below 50 and out of these coverage thresholds were excluded. Homozygous positions were also selected based on the same coverage threshold and considered ambiguous otherwise. For ONT and Illumina data, heterozygosity was computed as the number of SNP divided by the total number of sites excluding ambiguous positions. Using the above criteria, heterozygosity levels were similar between ONT and Illumina data (supplementary table S4, Supplementary Material online). To compare our results with the values of Mackintosh et al. (2019), heterozygosity was also computed using only four-fold degenerate sites, as identified by a custom script using BIO++ library (Guéguen et al. 2013).

Estimation of the Demographic History and Effective Population Size

We used a sequential Markovian coalescent (SMC) model (McVean and Cardin 2005, e.g., PSMC, Li and Durbin 2011; MSMC2, Schiffels and Wang 2020) to estimate the ancestral effective population size (Ne) trajectory of the studied Troidini species.

The SMC model requires calibrations, in particular a value of mutation rate. We estimated this rate based on synonymous mutations by selecting the four-fold degenerate sites of the third codon positions of BUSCO genes from the odb10 lepidopteran database. We retrieved the set of fasta nucleotide sequences using BUSCOMP 0.13.0 (Edwards 2019) on local runs of **BUSCO** (odb10 Lepidoptera) of the six studied individuals and we considered only the genes that contained all individuals, which corresponded to 5,127 genes (~97% of the lepidopteran gene database). Assuming that these mutations are neutral, we applied the formula $D = 2 \times T \times \mu$ where D is the genetic divergence between two species, T is the divergence in millions of years and μ is the mutation rate per million years (Kimura 1983; Birky and Walsh 1988). Here, we chose O. priamus as the divergent species of O. alexandrae, and set T = 12.03166 Ma (median value of the divergence time between these two species, with 95% credibility interval = 7.9662-16.7068; sensu Allio et al. 2021). To estimate D, we split the 5,127 genes into six bins based on the GC-content and we estimated the branch lengths from the six corresponding trees inferred by IQ-TREE 1.6.12 (Nguyen et al. 2015) with a GTR+Γ6 substitution model. The divergence D ranged from 0.043 to 0.052 for the lowest GC-content bins to the highest, respectively. We then took an average D (= 0.0475) between the two Ornithoptera species to obtain a mean value of μ equal to 1.9740e-09 mutations per site per year. As O. alexandrae highland population produces one generation per year while the lowland population produces two (Mitchell et al. 2016), we set the generation time to one and a half generations per year, therefore, the mutation rate μ was estimated at 1.3160e-09 mutations per site per generation, which is at the lowest end of the range estimated for Heliconius ($\mu = 1.3-5.5e-09$; Keightley et al. 2015).

It has been shown that SMC models do not perform well when the ratio of recombination rate r over mutation rate μ becomes greater than one (Sellinger et al. 2021). Assuming O. alexandrae genome is 325 Mb long, distributed in 30 chromosomes, and there is a single crossover per tetrad per male meiosis, the recombination rate would be r=2.7e-8. An analysis of the nymphalid $Vanessa\ cardui$ (Shipilina et al. 2022) estimated an average r between 3.81e-8 and 4.05e-8 with substantial interchromosomal variation, meaning that the average recombination rate of O. alexandrae is an order of magnitude greater than its mutation rate at least.

To investigate if this parameter range was a problem in SMC analyses, we used msprime 1.2.0 (Kelleher et al. 2016) to produce ten simulated datasets of a stable demographic history scenario with Ne = 100,000 under a r = 1e-8 and a $\mu = 1.316e-9$ on 30 chromosomes of 10 Mb each. We ran the MSMC models implemented in the MSMC2 software (https://github.com/stschiff/msmc2; Schiffels and Wang 2020) with default options (i.e., -rhoOverMu = 0.25) to test whether the inferred demography was recovered stable. We ran the same data by setting an initial value -rhoOverMu = 10.

We applied the MSMC2 model on real data, first for each Troidini species (FC563 was selected for *O. alexandrae*). We used the VCF files generated using Longshot (as described in the *Nuclear heterozygosity of Troidini* section) and created the so-called "mask file" for each individual based on the depth of coverage thresholds of $>20 \times$ and $<150 \times$ using a custom python script. These files were then combined using the "generate_multihetsep.py" of MSMC2 to generate "multihetsep.txt" input files (https://github.com/stschiff/msmc-tools/blob/master/msmc-tutorial/guide.md). MSMC2 was run using default parameters (especially the initial value of the ratio $[r/\mu]$ -rhoOverMu = 0.25), and with -rhoOverMu = 10 (See supplementary fig. S2, Supplementary Material online for a comparison of the results with both options).

By applying the same methods, we ran MSMC2 on each individual of O. alexandrae and then applied the model to the two populations (both composed of two genomes). MSMC2 was run using default parameters, or with -rhoOverMu = 10. The "-I" option was used to consider relevant haplotypes depending on the analyses (i.e., single individuals, two populations, and CCR between populations). For each analysis on real data, we generated ten bootstraps per individual using the multihetsep_bootstrap.py script available at: https://github.com/stschiff/msmctools. We generated all graphs with the R package ggplot2 (Wickham 2016) by considering a generation time of 0.5 for O. priamus and T. oblongomaculatus (two generations per year), 0.75 for O. alexandrae (one and a half generations per year) and $\mu = 1.316e-9$ for every individual. Finally, we relied on the ABC framework implemented in DILS (Fraïsse et al. 2021) to test several scenarios of divergence between populations. Alternative methods like ∂a∂i (Gutenkunst et al. 2009) or FastSimCoal (Excoffier and Foll 2011) could not be implemented because of no modeling of Ne through time and sample size limitations of our dataset to compute site-frequency spectrum, respectively. DILS takes into account linkage information that is informative about past demography (Fraïsse et al. 2021). To fit DILS on our data, we randomly selected 5,000 windows of 4,000 bp (2 Mb in total) because the coalescence program is time-consuming to simulate large chunks of chromosomes with recombination. The analysis was replicated four times to evaluate variability and reproducibility of the ABC inferences. DILS implements a pipeline that selects the best-fitting demographic model by comparing models with variations in *Ne* and migration among loci allowing to consider linked selection and alleles that could be selected against during hybridization (Fraïsse et al. 2021). The four demographic scenarios tested include SI, IM, AM, and SC. DILS used a random forest method (Pudlo et al. 2016) to select the best model and estimate posterior parameters' distributions using rejection and neural network methods implemented in the R package abc (Csilléry et al. 2012).

Population Structure

The population differentiation due to genetic structure was estimated with the nucleotide diversity and fixation index (F_{ST}) that were computed using seq_stat_2pop (https:// github.com/benoitnabholz/seq_stat_2pop) using Bio++ library (Guéguen et al. 2013). The seq_stat_2pop program uses fasta sequences as input such that VCF files were converted into fasta sequences using a custom python program using coverage information for the homozygous sites as explained above (see Nuclear heterozygosity of *Troidini* section). Nucleotide diversity (π -diversity) was computed as the mean pairwise divergence between pairs of chromosomes. F_{ST} was computed using the nucleotide sequence as $F_{ST} = 1 - \pi_{intra}/\pi_{total}$; where π_{intra} is the mean nucleotide diversity of the two populations (π_{intra} = $(\pi_{highland} + \pi_{lowland})/2$) and π_{total} is the nucleotide diversity computed using all individuals (Nei 1982). F_{ST} and nucleotide diversity were computed on nonsex-related contigs >500 kb in length, over sliding windows of 100 kb with 50 kb overlapping regions (windows of 100 kb and overlapping regions of 100 kb were tested and led to similar results). It is considered that a F_{ST} value greater than 0.15 is significant in differentiating populations (Frankham et al. 2010).

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online (http://www.gbe.oxfordjournals.org/).

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Author Contributions

F.L.C. and D.B. conceived and supervised the project. F.L.C. and D.B. collected the samples. E.C. and M.-K.T. carried out molecular experiments. E.L.R. and B.N. performed the bioinformatic analyses. E.L.R., B.N., and F.L.C. discussed the results. E.L.R. and F.L.C. wrote the draft manuscript and B.N., M.-K.T. and D.B. made comments. All authors have read and approved the final manuscript.

Data Availability

The birdwing genomes, mitogenomes and sequencing data in the present study, including Nanopore, Illumina, and RNA data are available from the Genome database and Sequence Read Archive under the Bioproject accession PRJNA938052, the number with corresponding BioSamples accession numbers FC560: SAMN33424250; FC561: FC562: SAMN33424251; SAMN33424252; FC565: SAMN33424254: FC563: SAMN33424253: FC569: SAMN33424255.

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