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The First Genome of the Balearic Shearwater (Puffinus mauretanicus) Provides a Valuable Resource for Conservation Genomics and Sheds Light on Adaptation to a Pelagic lifestyle

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Abstract

The Balearic shearwater (Puffinus mauretanicus) is the most threatened seabird in Europe and a member of the most speciose group of pelagic seabirds, the order Procellariiformes, which exhibit extreme adaptations to a pelagic lifestyle. The fossil record suggests that human colonisation of the Balearic Islands resulted in a sharp decrease of the Balearic shearwater population size. Currently, populations of the species continue to be decimated mainly due to predation by introduced mammals and bycatch in longline fisheries, with some studies predicting its extinction by 2070. Here, using a combination of short and long reads, we generate the first high-quality reference genome for the Balearic shearwater, with a completeness amongst the highest across available avian species. We used this reference genome to study critical aspects relevant to the conservation status of the species and to gain insights into the adaptation to a pelagic lifestyle of the order Procellariiformes. We detected relatively high levels of genome-wide heterozygosity in the Balearic shearwater despite its reduced population size. However, the reconstruction of its historical demography uncovered an abrupt population decline potentially linked to a reduction of the neritic zone during the Penultimate Glacial Period (~194–135 ka). Comparative genomics analyses uncover a set of candidate genes that may have played an important role into the adaptation to a pelagic lifestyle of Procellariiformes, including those for the enhancement of fishing capabilities, night vision, and the development of natriuresis. The reference genome obtained will be the crucial in the future development of genetic tools in conservation efforts for this Critically Endangered species.

Key words: Balearic shearwater, conservation genomics, comparative genomics, Procellariiformes marine adaptation.
The genomic sequence of a species accumulates valuable information on the evolutionary history, including demographic and selective events, and on the evolution of genes and traits (Jarvis et al. 2014; Foote et al. 2015; Nadachowska-Brzyska et al. 2015; Feng et al. 2020), information that is also crucial for the emerging field of conservation genomics (Allendorf 2017). The genetic diversity within a species represents a reservoir of adaptive variation that can help populations to cope with environmental variability (Dussex et al. 2021). Understanding the processes that shape genetic diversity and its distribution pattern within species is paramount to assess the conservation status or the factors responsible for a species decline (Brüniche-Olsen et al. 2021). This knowledge can inform the proposal of effective conservation and management plans, as for instance the definition of management units (Funk et al. 2012). In this context, next-generation sequencing techniques allow the analysis of an increased density of markers across the genome, providing unprecedented accuracy in the estimations of population genetic parameters relevant for scientific-based conservation recommendations (Supple and Shapiro 2018).

The Balearic shearwater is a medium-sized pelagic seabird endemic to the Balearic Islands that is listed as Critically Endangered by the IUCN Red List (IUCN 2021). Despite being the most endangered seabird in Europe, its genomic sequence has not been sequenced to date. At present, its population size is undergoing a fast annual decline of 7.4–14% (Genovart et al. 2016) mostly due to bycatch in longline fisheries and predation by invasive mammals in the colonies (Louzao et al. 2004; Arcos et al. 2008). Currently, it has a reduced number of breeding pairs (estimated as ca., 3,200, Arcos, 2011), with a total population size up to 30,000 individuals due to the vast contingent of floaters (Arcos et al. 2012; Arroyo et al. 2016). Genetic studies based on mtDNA and microsatellites found that this species has low levels of genetic diversity and high inbreeding coefficients (Genovart et al. 2012). Although local inbreeding and natal philopatry represent risk factors which may have negative effects on the viability of the species, the most significant threats come from human activities, and a population viability study based on demographic modeling predicted that the species would become extinct by 2070 (Genovart et al. 2016). Moreover, studies based on mitochondrial markers (Genovart et al. 2005) and also on morphology and migratory behavior (Austin et al. 2019), suggested a possible ongoing hybridization and introgression process between Balearic and Mediterranean (P. yelkouan) shearwaters, which may represent an additional threat for the species.

The Balearic shearwater belongs to the most diverse order of seabirds, the Procellariiformes. This order has a worldwide distribution and comprises more than 140 species (IUCN 2021) in four families: petrels and shearwaters (Procellariidae); northern storm petrels (Hydrobatidae); southern storm petrels (Oceanitidae); and albatrosses (Diomedeidae). There is a large variation in body mass and lifestyles within the Procellariiformes, ranging from 20 g to 15 kg; from bodies shaped for diving (e.g., short strong wings used for wing-propelled diving) to those fitted for an extremely agile lifestyle (thin elongated wings), and from a continuous flapping to dynamic soaring flight modes. Despite this diversity, all Procellariiformes show many morphological, physiological, and life-history traits associated with adaptation to a pelagic lifestyle. Such adaptations include prominent salt glands to facilitate the secretion of salt, adaptations to enhance fishing capabilities such as underwater vision and a particularly acute sense of smell, among other traits (Brooke 2004). Even so, there is no compiled study on the genomic signatures of adaptation to a pelagic lifestyle (but see Silva et al. 2020).

Here, we make available a high-quality reference genome for the Balearic shearwater along its structural and functional annotations. This resource allows us to provide important information for the conservation of the species, such as the estimation of genome-wide heterozygosity and historical demography of the species. Benefiting from this new genome assembly and seven additional Procellariiformes genomes (Feng et al. 2020), we also performed a comparative genomics analysis to uncover the putative adaptations to a pelagic lifestyle. The high-quality genome of the most endangered seabird in Europe presented here will be the basis for further population-based conservation genomics studies.

**Significance**

The Balearic shearwater (Puffinus mauretanicus) is the most threatened seabird in Europe and some studies predict their extinction by 2070. Here we provide a high-quality genome assembly of this species to study critical aspects relevant to its conservation status. We found that despite its low population size, the species harbors relatively high levels of heterozygosity, whereas its historical demography uncovered an abrupt population decline during the Penultimate Glacial Period. In addition, we identified some candidate genes that may have played an important role into the adaptation to a pelagic lifestyle in Procellariiformes. Our data will be useful for future conservation and management plans.
Results

Sequencing Data and Genome Assembly and Annotation

Illumina paired-end (2 × 150 bp) sequencing of the male-Mll yielded a throughput of 147.7 Gbp (table 1), representing a mean coverage of 118 ×. The five runs of ONT sequencing of the unsexed-Ei resulted in a 10 × coverage with a read N50 of 9,431 bp. RNA sequencing (RNA-seq) of the chick-Mll (2 × 100 bp) yielded 15 Gbp of data.

We obtained a hybrid assembly with MaSuRCA formed by 4,169 scaffolds, with an N50 of 2.1 Mbp, and an assembly length of 1.21 Gbp (table 2 and fig. 1b). The completeness analysis using BUSCO yields a value of 95.9%, and only 0.3% of the complete genes were duplicated and 1.1% were fragmented (table 2). Our de novo repeat annotation analysis shows that 9.95% of the genome consists of repetitive regions (supplementary table S1, Supplementary Material online), which is within the range of previously sequenced avian genomes (Zhang et al. 2014). Among repeat elements, long interspersed nuclear elements were the most abundant (4.45% of the genome). The genome annotation process resulted in a total of 21,959 protein-coding genes, of which 18,769 (85.5%) have at least one gene ontology (GO)-associated term, and 19,218 (87.5%) have at least one hit across the surveyed curated databases (supplementary table S2, Supplementary Material online).

Blood transcriptome assembly from the chick-Mll resulted in 224,904 transcripts (supplementary table S3, Supplementary Material online). However, BUSCO completeness was only 62.4%, which was far below genome completeness, probably due to the RNA coming from a single and not very transcriptionally active tissue.

The assembly of the mitogenome of Puffinus mauretanicus resulted in a single contig of 19,885 bp long, with a coverage (Illumina reads) of 371 ×, which is around three times higher than the coverage of the nuclear genome. This mitogenome has the same gene order as other published Procellariiformes’ mitogenomes (supplementary fig. S1, Supplementary Material online). The mitogenome has two copies of the nad6 gene, as predicted in P. iherminieri (Torres et al. 2019); the later feature was also confirmed analyzing the mean coverage (Illumina reads) across these genes (supplementary table S4, Supplementary Material online).

Table 1
Sequencing Data, Library Information, and Samples Used in this Study

| Library | Total Number of Base Pairs | Number of Reads | Coverage | Individual Code | Location |
|---------|---------------------------|----------------|----------|-----------------|----------|
| HiSeq X Ten – TruSeq DNA PCR Free. 2 × 150 bp | 143,765,593,200 | 958,437,288 | 118 × | Male-Mll | Sa Cella (Mallorca) |
| ONT – Ligation kit SQK-LSK109 1D | 12,142,789,693 | 2,576,486 | 10 × | Unsexed-Ei | Sa Conillera (Eivissa) |
| NovaSeq 6000 – TruSeq RNA Sample Prep Kit v2. 2 × 100 bp | 14,997,592,000 | 149,975,920 | – | Chick-Mll | Conills islet (Mallorca) |

Table 2
Balearic Shearwater Genome Assembly Metrics

| Assembly length (bp) | 1,218,519,395 |
|----------------------|---------------|
| Number of scaffolds  | 4,169         |
| Longest scaffold (Mbp) | 11.06       |
| N50 (Mbp)           | 2.13          |
| L50                 | 164           |
| GC content (%)      | 42.52         |
| Repetitive content (%) | 9.95       |
| Mitogenome (bp)     | 19,855        |
| No. protein-coding genes | 21,959    |
| BUSCO %             | 95.9          |
| Complete            | 95.9          |
| Single copy         | 95.6          |
| Duplicated          | 0.3           |
| Fragmented          | 1.1           |
| Missing             | 3.0           |

Historical Demography of the Balearic Shearwater

Multiple Sequentially Markovian Coalescent (MSMC2) analysis showed support for a steady growth in $N_e$ from an originally low $N_e$ followed by a sudden increase ~200 ka (fig. 1c). High effective population size did not last long, and the species suffered a sudden decrease to nearly one tenth of the population coinciding with the end of the glacial period before the last interglacial period (119–128 ka) and a prolonged period of low sea level (fig. 1c). Hereafter, $N_e$ remained stable until ~10 ka ago, as more recent MSMC2 time segments are regarded as being unreliable (Schiffels and Wang 2020).

Genome-wide heterozygosity in P. mauretanicus was 0.0024, which is within the range of genome-wide heterozygosities estimated for other Procellariiformes (ranging from 0.0014 in Thalassarche chlororhynchos to 0.0037 in Pelecanoides urinatrix) (fig. 2a). Among Procellariiformes, small-bodied species (~<200 g) tended to have higher mean heterozygosities but also higher variance than large-bodied species (~>450 g) (fig. 2b).

Phylogenetic Relationships

OrthoFinder analysis estimated 6,172 single-copy (1:1) ortholog genes across the 12 genomes surveyed. With this data we generated three supermatrices: (1) coding sequence (CDS) supermatrix of 10,534,506 bp long to extract
Comparative Genomics and Positive Selection Analyses

To identify genes associated with adaptation to a pelagic lifestyle in the Procellariiformes, we performed a positive selection analysis across 12 species including Procellariiformes species applying the HyPhy aBSREL model. We identified the hallmark of positive selection in 20 out of the 6,172 single-copy orthologs genes, after correcting for multiple testing (supplementary table S5, Supplementary Material online). We found among these genes enriched GO functions related with striated muscle cell differentiation, nutrient reservoir activity, response to starvation, olfactory receptor activity, or natriuresis (supplementary table S7, Supplementary Material online). We also performed an HyPhy RELAX analysis to assess the intensification/relaxation of natural selection in Procellariiformes, which uncovered a total of 310 genes (supplementary table S7, Supplementary Material online). The GO terms enriched in these genes include wound healing, response to wounding, inflammatory response, sensory perception of sound, smell and chemical stimulus, neurological system process, defense response, response to stress, camera-type eye development, renal system, and chloride transport among others (supplementary table S8, Supplementary Material online).

Using OrthoFinder 2.3.8, we identified 182,487 N:N orthogroups across all genes identified in the 12 analyzed genomes. These data, together with the estimated ultrametric tree, were used to estimate gene gains, losses, and number of genes in the ancestral nodes using BadiRate; for the analysis we selected the Free Rates (FR) model, because it was the best-fitted branch model. The analysis was conducted including all orthogroups, and the minimum number of gains and losses per branch is represented in the ultrametric tree. The analysis performed to explicitly account for incomplete lineage sorting (ILS) with ASTRAL using either the individual gene sequences (CDS gene trees) or the individual amino acid sequences (amino acid gene trees), also resulted in the same topology (supplementary figs. S4 and S5, Supplementary Material online). The normalized quartet score (proportion of input gene tree quartet trees in agreement with the species tree) was 0.78 for CDS gene trees and 0.64 for amino acid gene trees.
Our results showed a tendency to gain genes in Procellariiformes (+442/−34), whereas the branch leading to albatrosses (Diomedeidae) showed an opposite effect, with a noticeable loss of genes (+464/−3258); the branch leading to the rest of the Procellariiformes (+379/−15) is in the line of the general behavior of the tubenoses (supplementary table S9, Supplementary Material online).

Within the order, families Oceanitidae and Hydrobatidae present the same trend, with the branch leading to Hydrobates tethys presenting a stronger gene loss balance (+325/−966) than the branch leading to the ancestor of Oceanitidae (Oceanites oceanicus and Fregetta grallaria [+182/−414]).

We identified three gene families significantly expanded in the branch leading to the Procellariiformes (supplementary table S9, Supplementary Material online). These families encode zinc finger proteins (OG0000000), olfactory receptors (OG0000084), and avian histones (OG0000224).

**Discussion**

**A High-Quality Genome Assembly for the Most Endangered Seabird in Europe**

The assembly length and the GC content of the Balearic shearwater hybrid assembly presented here are similar to...
those reported in the seven Procellariiformes genomes released by the Bird 10,000 Genomes Project (B10K) (Feng et al. 2020). Repetitive content is remarkably higher (+33.4%) in the Balearic shearwater in comparison to the other genomes of the order, but within the range of avian genomes (Zhang et al. 2014). This up to a third increase in repetitive sequence can be due to the fact that we included a Procellariiform (Calonectris borealis) repeat library before running RepeatMasker, achieving a more precise library that encloses clade related repeats that are present in the genome but not found by the de novo RepeatModeler library. The genome assembly completeness (BUSCO 95.9%) is slightly higher than the obtained for other recently published bird genomes (Prost et al. 2019; Feng et al. 2020), and even higher than genome assemblies including optical mapping (Peñalba et al. 2020). Despite not being a chromosome-scale assembly, contiguity is also quite high (N50 2.1 Mbp), and higher than recent avian MaSuRCA hybrid assemblies (Gan et al. 2019; Leroy et al. 2019).

The retrieved proteome (21,959 protein-coding genes) is similar to previous genomes (Liu et al. 2021; Recuerda et al. 2021), but higher than the B10K 2020 genomes used in the comparative studies in this work (mean of 16K). This is probably due to the B10K annotation pipeline being fully based on homology, while we also used de novo prediction. The functional annotation quality in terms of genes having at least a GO term (85.9%) is comparable to recent chromosome-scale genomes (Recuerda et al. 2021).

The mitogenome of P. mauretanicus spans 19,885 bp, exhibiting the same order and the nad6 gene duplication observed in P. lherminieri (Torres et al. 2019). We did not find any cob duplication as it occurs in the Diomedeidae family (Abbott et al. 2005). Our result supports the hypothesis that nad6 duplication could be widespread in Procellariiformes (Torres et al. 2019), and, like cob, could have undergone various events of deletion or addition during the diversification of the order. Nevertheless, because some of the reported duplications could be artificial (Urantówka et al. 2020; Formenti et al. 2021), to fully identify the true number of gene duplications/deletions will require additional and specific experimental analyses.

Balearic Shearwater Heterozygosity Levels and Historical Demography

Current levels of intraspecific heterozygosity is a relevant parameter to determine the adaptive capacity of a population (or species) (Ørsted et al. 2019). Because the Balearic...
shearwater is categorized as Critically Endangered by the IUCN, we could naively expect low heterozygosity levels in the species when compared with other Procellariiformes. However, the fossil record suggests that the Balearic shearwater had a very large population (>30,000 pairs) until the arrival of human settlers in the Balearic Islands (Alcover et al. 1991), which hunted shearwaters (Ramis 2018) and introduced invasive mammals that also predated on them (Pinya and Carretero 2011).

In line with Genovart et al. (2007) results using mtDNA markers, we observed relatively high genome-wide heterozygosity levels, suggesting that the very recent demographic decline in the species is not yet visible in its genetic diversity.

Regarding the historical demography, our MSMC2 analysis shows an increase in \( N_e \) in the Balearic shearwater from around 1 Ma to later expand to reach high population sizes, until around 150,000 ya, when it suddenly suffered a sharp decline, resulting in lower \( N_e \) values maintained until 10,000 years ago. Because current MSMC2 analysis is based on the analysis of a single genome, we could not reliably infer more recent events (Schiffels and Wang 2020).

The Plio-Pleistocene eustatic variations resulted in a loss of neritic zones as sea level regressed (Pimiento et al. 2017), this may represent a loss of coastal habitat availability, which added to other oceanographic alterations (changes in ocean circulation or productivity) may have been the drivers of great population losses in marine megafauna, including seabirds. In the case of the Balearic shearwater, the MSMC2 analysis shows an abrupt decay of \( N_e \) associated with a long period of low sea level during the Penultimate Glacial Period (~194–135 ka) which may have resulted in an important loss of neritic zones. However, the particular timing of the sharp \( N_e \) decline should be interpreted with caution as it depends on both the mutation rate and generation time estimates; therefore, underestimations or overestimations of these parameters would result in biased timings (Nadachowska-Brzyska et al. 2015).

In view of the current critical population declines affecting the Balearic shearwater populations, understanding its impacts on current genetic diversity of the species and among colonies will be crucial to assess the conservation status of the Balearic shearwater. Future ongoing research, using a more powerful population genomics approach, will allow the reconstruction of more recent demographic histories of the species and to test the fossil-based hypotheses of a recent loss of population due to human colonization of the island, as well as why heterozygosity values have not decayed.

Association Between Body Size and Heterozygosity in Procellariiformes

We observed a strong association between body size and heterozygosity in the Procellariiformes. Small-bodied species (\( O. \) oceanicus, \( F. \) grallaria, \( H. \) tethys, and \( P. \) urinatrix) have higher heterozygosities and higher variance in heterozygosity levels across the genome than large-bodied species (\( F. \) glacialis, \( C. \) borealis, \( P. \) mauretanicus and \( T. \) chlororhynchos). This pattern is in line with the negative correlations reported between heterozygosity and body size across animals from many taxa (Romiguier et al. 2014; Brüniche-Olsen et al. 2018, 2021). Such well-supported correlations are likely due to the interplay between \( N_e \) and \( \mu \), both of which are correlated with body size (Romiguier et al. 2014; Brüniche-Olsen et al. 2018). Smaller animals generally live at higher population densities, which tend to be associated with higher \( N_e \). Indeed, the small-bodied species included here have a higher number of breeding pairs (200,000–7,000,000) than large-bodied species (3,193–400,000) (Billerman et al. 2020). In addition, the small Procellariiformes lineages included here, also show higher \( \mu \) (Nunn and Stanley 1998; Estandia et al. 2021), showing the likely interplay of \( N_e \) and \( \mu \) at driving the association between body mass and heterozygosity. Island species, such as most Procellariiformes, tend to show lower levels of heterozygosity and increased numbers of deleterious mutations, probably due to a lower ability of natural selection to efficiently remove weakly deleterious mutations in small population (Leroy et al. 2021). Because standing genetic variation is the main source for rapid adaptation (Barrett and Schluter 2008; Jamie and Meier 2020), species with low levels of heterozygosity might not be able to rapidly adapt to environmental changes. Due to their low levels of heterozygosity, large-bodied Procellariiformes may be particularly vulnerable to environmental change. This could represent an additional threat for these species that are already suffering from anthropogenic threats such as predation by invasive alien species and fisheries bycatch (Dias et al. 2019). Future studies should investigate the accumulation of deleterious mutations in Procellariiformes to understand the potential responses of procellariiform species to environmental changes.

Adaptation to a Pelagic Lifestyle in Procellariiformes

Our selection inference uncovered 20 genes evolving under positive selection in Procellariiformes, and are therefore potential candidates involved in the adaptation of the order to a pelagic lifestyle. Indeed, the function of these genes reveals biological processes related to striated muscle cell differentiation, response to starvation, and nutrient reservoir activity, that may be related to the high energy expenditure during the vast distances they cover in the open ocean, whereas visual related genes could be related with underwater vision to fish and night vision (Hayes et al. 1991; Martin and De 1991; Mitkus et al. 2016). Positive selection of genes related to natriuresis also makes sense for
Procellariiformes because this biological process plays a key role to maintain the osmotic equilibrium in a sodium-rich environment like the ocean (Goldstein 2001; Gutiérrez 2014), which Procellariiformes perform thanks to the development of salt glands (modified nasal glands engaged in secretion of salts). Olfactory receptors, also found here among enriched GOs of positively selected genes, showed signature of adaptive evolution in shearwaters (Silva et al. 2020), and are crucial to Procellariiformes for navigation (Gagliardo et al. 2013; Pollonara et al. 2015; Padget et al. 2017), partner recognition and mating (Bonadonna and Nevitt 2004; Strandh et al. 2012; Hoover et al. 2018), finding their own burrows (Bonadonna and Bretagnolle 2002), or foraging (Nevitt et al. 1995; Nevitt 2008; Bastos et al. 2020; Sin et al. 2022).

We also identified genes with intensified natural selection in Procellariiformes, for which the GOs annotations (supplementary table S8, Supplementary Material online) are similar and coherent to those in the candidate set of genes with positive selection in all tubenoses, or, in other words, related to the adaptation of the order to a pelagic lifestyle. For example, molecular functions such as sensory perception of sound, smell and chemical stimulus, neurological system process, camera-type eye development are related with oceanic navigation. On the other hand, functions such as homeostatic process, renal system, renal response, chloride transport, and regulation of ion transport point to the need of maintaining osmotic equilibrium. We also found intensified natural selection in genes participating in functions related to immune response (like inflammatory response, defense response, response to wounding, wound healing, positive regulation of phagocytesis, etc.), accompanied by a relaxation of natural selection in regulators of blood constituents, induction of bacterial agglutination, regulation of antigen processing and presentation, viral budding via host ESCRT complex, or macrophage antigen processing and presentation. As exposure to parasites in Procellariiformes is high (Khan et al. 2019) and their life-history traits favor parasite maintenance within populations (McCoy et al. 2016), we hypothesize that the tuning between the intensification and relaxation of natural selection in multiple biological processes and molecular functions related with immune response could have emerged following an arms race-like model. For example, as many parasites of tubenoses are blood-feeding, the intensified natural selection on the thrombin-activated receptor signaling pathway (GO:0070493), could be an evolutionary response to counter the anticoagulant activity that most blood-feeding parasites present (Bensaoud et al. 2018).

Among the gene families expanded in the branch of Procellariiformes, the one encoding olfactory receptors is remarkable as it is coherent not only with the finding of a gene with positive selection in all Procellariiformes with the same functional annotation (g16276.t1 in the P. mauretanicus reference annotation, supplementary table S4, Supplementary Material online) but also with three olfactory receptors genes with intensification selection in the same branch (g14377.t1, g16276.t1, and g17936.t1 in P. mauretanicus reference annotation, supplementary table S8, Supplementary Material online). This triple evidence highlights the importance of how the adaptation to a pelagic life resulted in the enhancement of the olfactory function in Procellariiformes, as discussed above. Moreover, similar results of positive selection in olfactory genes were obtained in C. borealis (Silva et al. 2020). Physiologically, tubenoses have one of the largest olfactory bulb to brain size ratio of all birds (Cobb 1968).

Conclusions
Our study highlights the utility of the hybrid assembly strategy using Illumina and ONT at recovering high-quality genome assemblies, especially regarding contiguity and completeness. Comparative genomics analyses identified candidate genes under selection to have played a major role in the adaptation of the Procellariiformes to a pelagic lifestyle such as changes in sensory perception, navigation, natriuresis, and physiological adaptations. The high-quality genome presented in this work will be a great tool for future population genomic analyses, that will reveal with more precision the genetic variability of the species, its recent demographic history and the potential introgression with its sister species, the Mediterranean shearwater (P. yelkouan). The data obtained will be of great help in future conservation and management plans for the species.

Materials and Methods
Sampling, DNA and RNA Extraction, and Sequencing
We sampled two Balearic shearwater adults and one chick. Adults were sampled on Sa Cella colony, Mallorca (male) and on Sa Conillera, Eivissa (unsexed) in 2004, whereas the chick was sampled on Conills islet (Mallorca) in July 2019. From here on the animals will be referred to as male-MII, unsexed-EI, and chick-MII, respectively. Special permits to obtain the samples were issued by Conselleria de Medi Ambient, Agricultura i Pesca (Govern de les Illes Balears, Spain).

We extracted DNA from blood samples preserved in absolute ethanol for both adults. The DNA extraction for the male-MII was performed with DNeasy Blood & Tissue Kit (Qiagen) following the manufacturer’s instructions, and with Blood & Cell Culture DNA Mini Kit (Qiagen) for the unsexed-EI. RNA was extracted from the chick-MII’s blood cells preserved in RNAlater 1:5 using the RNeasy Mini Kit (Qiagen) according to the manufacturer’s protocols. We performed the quality control with gel electrophoresis.

We performed the quality control with gel electrophoresis...
and NanoDrop One (Thermo Fisher Scientific, Waltham, MA, USA), and the quantification with an Invitrogen Qubit Fluorometer 2.0 (Broad Range kit).

We obtained the reference genome combining short-read and long-read sequencing libraries, and using RNA-seq data to assist with the annotation. First, an Illumina TruSeq DNA PCR Free library (insert size = 350 bp) was prepared by Macrogen (South Korea) using DNA from male-Mll, and sequenced using some HiSeq X Ten runs (2 × 150 bp). Second, long-read libraries were prepared, from the DNA of unsexed-Ei, using the Ligation kit SQK-LSK109 1D from ONT (Oxford Nanopore Technologies) (N50 of 9431 bp) at Centro Nacional de Análisis Genómico, Spain and sequenced through five runs of MinION on FLO-MIN106 flow cells. Third, we prepared RNA-seq libraries from the chick-Mll's RNA using the TruSeq RNA Sample Prep Kit v2 with Ribo-Zero, and we sequenced the libraries on a NovaSeq 6000 (2 × 100 bp) (Macrogen, South Korea).

**Genome Assembly**

We performed a de novo hybrid genome assembly with MaSuRCA 3.3.1 (Zimin et al. 2017), using short (Illumina) and long (ONT) reads. Before the assembly step, we filtered the ONT reads with a Phred quality score (Q ≥ 5) using the NanoFilt software included in NanoPack (De Coster et al. 2018). Paired-end Illumina reads were parsed into MaSuRCA without any preprocessing, as adapters and errors are handled by the QuORUM error corrector (Marçais et al. 2015), which is part of the MaSuRCA pipeline. MaSuRCA was run applying the following parameters: fragment mean (422), fragment stdev (312), and estimated genome size (1.2 Gbp). The resulting assembly was screened for contaminants with BlobTools v1.0 (Laetsch and Blaxter 2017) -x bestsumorder. Assembly completeness was assessed with BUSCO 4.0.2 (Sepey et al. 2019) using the 8,338 single-copy conserved genes in aves_odb10 database (Kriventseva et al. 2019).

**Transcriptome Assembly**

We trimmed RNA-seq raw reads for adapters with BBDuk (https://sourceforge.net/projects/bbmap/) (k = 23, tpe option), and used STAR 2.7.3a (Dobin et al. 2013) to map the filtered reads to the newly assembled reference genome. We obtained the transcriptome assembly with Trinity 2.8.6 (Grabherr et al. 2011) using the genome-guided bam mode (--genome_guided max_intron 82945). Transcripts were clustered with CD-HIT (Li and Godzik 2006; Fu et al. 2012) 4.8.1 (< 0.98) and CDS were predicted with TransDecoder 5.5.0 (https://github.com/TransDecoder).

**Mitogenome Assembly**

We trimmed adapters from Illumina raw reads with BBDuk (k = 23, tpe option), before using them as input to NOVOPlasty 2.7.2 (Dierckxsens et al. 2017). The P. thermienieri mitogenome (MH206163.1) was used as seed using the following parameters: Genome Range (16,000–24,000), Insert size (422), Insert range (1.74) and Insert range strict (1.3). The annotation was performed using the MITOS WebServer (Bert et al. 2013).

**Repeat Annotation**

We generated a de novo repeat library of the genome with RepeatModeler—1.0.11 (Smit et al. 2021) on scaffolds >100 kbp. This library was combined with all avian and ancestral consensus repeats from Dfam_Consensus-20181026 (Storer et al. 2021), RepBase-20181026 (Jurka et al. 2005), and the repeat annotation of the Cory’s shearwater (C. borealis) (Feng et al. 2020), which represents the most closely related sequenced genome. Redundancies among libraries were removed with the script ReannTE_MergeFasta.pl (https://github.com/4ureliek/ReannTE). We then ran RepeatMasker 4.0.7 (Smit et al. 2021) using the combined library as a reference, with the following parameters: -xsmall -e ncbi -s -gccalc -no_is -gff.

**Structural and Functional Annotation**

We performed the structural annotation with BRAKER 2.1.2 (https://github.com/Gaius-Augustus/BRAKER) (–etp-mode) using data from both the Cory’s shearwater proteome (Feng et al. 2020), and the RNA-Seq data generated in this work. Because the inclusion of RNA-Seq data appeared detrimental, we excluded this piece of information to perform the final annotation using the soft-masked genome with BRAKER 2.1.2 (–prg = gth –trainFromGth).

We made the functional annotation of the predicted genes using a similarity-based approach. We determined the protein domains with InterProScan 5.31–70.0 (Jones et al. 2014), used BLASTP (Altschul et al. 1990; Camacho et al. 2009) (--evalue 1e-5; --max_target_seqs 10) against the Swiss-Prot database (Boutet et al. 2016) and the Cory’s shearwater and the Zebra finch reference proteome (UP000007754) proteomes. Transcripts were annotated in the same manner. We also annotated the ncRNAs using cmscan from INFERNAL 1.1.2 (Nawrocki et al. 2009) with the covariance models (CMs) from the Rfam 14.1 database, and tRNA genes using tRNAscan-SE 2.0.5 (Chan and Lowe 2019).

**Demographic History**

We used the Multiple Sequentially Markovian Coalescent 2 (MSMC2) (Schiffels and Wang 2020) to infer the historical demography of the Balearic shearwater. MSMC2
implements a MSMC model, which allows the estimation of the effective population size ($N_e$) over time. To generate input files for MSMC2, first we mapped cleaned Illumina short reads from male-Mll to the reference genome using BWA-MEM 0.7.17 (Li and Durbin 2009). We then subsampled the bam file to only include scaffolds larger than 1 Mbp (343 scaffolds spanning 71.8% of the assembled genome), as recommended in Gower et al. (2018). Second, we called SNPs using samtools mpileup 1.9 (options: `-q 20 -Q 20 -C 50`) and then bcftools call 1.9 (options `-c -V indels`) (Danecek et al. 2021) and, for each scaffold, we generated a VCF file and a mask file in bed format, containing the regions on the scaffold that were sufficiently covered using the bamCaller.py script provided in the msmc-tools package (https://github.com/stschiff/msmc-tools). Third, we generated a mappability mask using GenMap (Pockrandt et al. 2020) with a k-mer size of 150 bp and allowing for up to two mismatches. We masked sites with a mappability score of <0.5 and we also masked sites within annotations of repetitive regions. In total, we masked 9.37% of the sites in the 343 scaffolds larger than 1 Mbp. Fourth, we generated input files for MSMC2 with the script generate_multihetsep.py, also provided in the msmc-tools package, by using the VCF and mask files as input files. Fifth, multiple sequentially markovian coalescent (MSCM) for two haplotypes was run with MSMC2 with time patterning specified as `-p 1*4 + 30 * 2 + 1*4 + 1*6 + 1*10`. Additional time patterning settings were used to assess the effect of the number of segments on the shape of the MSMC2 curves, but this had little effect on the shape.

We obtained the confidence intervals for $N_e$ estimates by running 100 bootstraps using multihetsep_bootstrap.py. We scaled time and $N_e$ using a generation time for the Balearic shearwater of 12.8 years (Genovart et al. 2016) and the Northern fulmar ($F. glacialis$) mutation rate ($\mu = 2.89 \times 10^{-9}$ substitutions per nucleotide per generation, Nadachowska-Brzyska et al. 2015).

**Genome-Wide Heterozygosity**

We estimated genome-wide heterozygosities using information from a single individual from all eight Procellariiformes species studied. We applied the Robinson et al. (2019) method, with minor modifications to take genome fragmentation into consideration, because we included genome assemblies with varying amounts of contiguity. The DNA sequence data (genome assemblies and whole-genome sequencing data) were downloaded from NCBI (PRJNA261828, PRJNAs545868, Jarvis et al. 2014; Feng et al. 2020). For each species, adapter-trimmed reads were aligned to its genome assembly using BWA-MEM (Li 2013), bam files were merged using Picard-Tools (http://broadinstitute.github.io/picard/) and variants were called using the GATK 4.1.9 HaplotypeCaller and GenotypeGVCFs (McKenna et al. 2010). Sites with a coverage $<1/3 \times$ or $>2 \times$ of the average coverage depth (of the particular genome) were filtered out using VCFtools 0.1.15 (Danecek et al. 2011). We computed per-site heterozygosity as the proportion of heterozygous sites per total number of called genotypes within a single individual in nonoverlapping 25 Kb windows across each scaffold. Windows with $<50\%$ of net sites (those excluding missing or filtered sites), were excluded from the analysis.

**Orthology Inference**

We performed the phylogenomics and comparative genomics analyses including information from 12 species with an available genome assembly: eight Procellariiformes ($P. mauretanicus$, $T. chlororhynchos$, $H. tethys$, $O. aethereus$, $F. gracilis$, $E. garzetta$, $P. urinatrix$, $C. borealis$), and four outgroups, Aptenodytes forsteri, Pygoscelis adeliae (Sphenisciformes); $Egretta garzetta$, Phalacrocorax carbo (Pelecaniformes). We inferred orthologous genes across the proteomes of these 12 species using OrthoFinder 2.3.8 (Emms and Kelly 2019) with default parameters.

**Inference of Phylogenetic Relationships**

In order to perform selection and gene family evolution analyses, we performed phylogenetic analyses using the inferred orthologous genes. Despite the Procellariiformes phylogeny has been recently resolved using UCE data (Estandia et al. 2021), we performed phylogenetic analyses to evaluate the robustness of this phylogeny when using different phylogenomic datasets. We built a multiple sequence alignment (MSA) for each 1:1 orthologs with the script generate_multihetsep.py, also provided in the msmc-tools package, by using the VCF and mask files as input files. Fifth, multiple sequentially markovian coalescent (MSCM) for two haplotypes was run with MSMC2 with time patterning specified as `-p 1*4 + 30 * 2 + 1*4 + 1*6 + 1*10`. Additional time patterning settings were used to assess the effect of the number of segments on the shape of the MSMC2 curves, but this had little effect on the shape.

We obtained the confidence intervals for $N_e$ estimates by running 100 bootstraps using multihetsep_bootstrap.py. We scaled time and $N_e$ using a generation time for the Balearic shearwater of 12.8 years (Genovart et al. 2016) and the Northern fulmar ($F. glacialis$) mutation rate ($\mu = 2.89 \times 10^{-9}$ substitutions per nucleotide per generation, Nadachowska-Brzyska et al. 2015).
orthologs MSAs were Before the analysis, nonreliable positions across all 1:1 (Wertheim et al. 2015) to test for relaxed/intensi
2015) to test for positive selection, and the RELAX method
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be associated with pelagic lifestyle. For this purpose, we
estimated gene turnover rates, the number of gene
turnover rates, several biological relevant hypotheses with
We tested, under the Birth-Death-Innovation model for
used the calibrated ultrametric tree estimated with r8s.
We generated an ultrametric tree with r8s v.1.81
(Sanderson 2003) using the 4D supermatrix ML tree. We
used four calibration points (in myr): root (max_age =
84 min_age = 73, Braun et al. 2011), most recent common
ancestor (MRCA) of Spheniscidae (min_age = 12.6,
Subramanian et al. 2013), MRCA Procellariiformes (min_
age = 49, Claramunt and Cracraft 2015), MRCA Procellariidae (min_age = 14, Prum et al. 2015), retrieved
from TimeTree (Kumar et al. 2017). We used the penalized
likelihood method and the Truncated Newton algorithm,
and the smoothing parameter was set to 100.

Positive Selection Analysis
We evaluated the selective constraints of genes that could
be associated with pelagic lifestyle. For this purpose, we
performed the analysis with HyPhy 2.5 (Kosakovsky Pond
et al. 2005), using Procellariiformes data (1:1 MSAs).
Before the analysis, nonreliable positions across all 1:1
orthologs MSAs were filtered with ZORRO (Wu et al.
2012) (default options; MSA with average quality <5
were filtered). We used the aBSREL method (Smith et al.
2015) to test for positive selection, and the RELAX method
(Wertheim et al. 2015) to test for relaxed/intensified selec
tion. We also performed a GO enrichment analysis of the
candidate genes using the GOstats (Falcon and
Gentleman 2007) R package against the background GOs
of 1:1 orthologs.

Gene Family Evolution
We estimated gene turnover rates, the number of gene
gains and losses across the phylogeny lineages, and in
ferred gene family contractions and expansions using
BadiRate 1.7 (Librado et al. 2012). For the analysis we first
inferred the orthogroups with OrthoFinder 2.3.8, and we
used the calibrated ultrametric tree estimated with r8s.
We tested, under the Birth-Death-Innovation model for
turnover rates, several biological relevant hypotheses with
three different branch models: FR, global rates, and branch
speciﬁc rates, and chose the best model based on the low
est Akaike information criterion value. To ensure an ap
propriate convergence we ran multiple times each model.

Supplementary Material
Supplementary data are available at Genome Biology and
Evolution online.
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