Evolution of immune genes in island birds: reduction in population sizes can explain island syndrome

Mathilde BARTHE^{1*}, Claire DOUTRELANT², Rita COVAS³⁻⁵, Martim MELO³⁻⁵, Juan Carlos ILLERA⁶, Marie-Ka TILAK¹, Constance COLOMBIER¹, Thibault LEROY^{1,7}, Claire LOISEAU^{2,3a}, Benoit NABHOLZ^{1,8a}

¹ ISEM, Univ Montpellier, CNRS, IRD, Montpellier, France

² Centre d'Ecologie Fonctionnelle et Evolutive, CNRS, Univ Montpellier, EPHE, IRD, Montpellier, France

³ CIBIO, Research Centre in Biodiversity and Genetic Resources, InBIO Associated Laboratory, Campus Agrário de Vairão, Vairão, Portugal

⁴ MHNC-UP, Natural History and Science Museum of the University of Porto, Porto, Portugal

⁵ DST/NRF Centre of Excellence, FitzPatrick Institute, University of Cape Town, Rondebosch, South Africa

⁶ Biodiversity Research Institute (CSIC-Oviedo University-Principality of Asturias), Oviedo University, Mieres, Spain

⁷IRHS-UMR1345, Université d'Angers, INRAE, Institut Agro, SFR 4207 QuaSaV, 49071, Beaucouzé, France

⁸Institut universitaire de France, Paris

* Corresponding author : mathilde.barthe.pro@gmail.com

^a Last co-authors

1 Abstract

Shared ecological conditions encountered by species that colonize islands often lead to the 2 3 evolution of convergent phenotypes, commonly referred to as the "island syndrome". Reduced immune functions have been previously proposed to be part of this syndrome, as a consequence 4 of the reduced diversity of pathogens on island ecosystems. According to this hypothesis, 5 immune genes are expected to exhibit genomic signatures of relaxed selection pressure in 6 7 island species. In this study, we used comparative genomic methods to study immune genes in 8 island species (N = 20) and their mainland relatives (N = 14). We gathered public data as well 9 as generated new data on innate (TLR: Toll-Like Receptors, BD: Beta Defensins) and acquired 10 immune genes (MHC: Major Histocompatibility Complex classes I and II), but also on hundreds of genes with various immune fonctions. As a control, we used a set of 97 genes, not 11 12 known as involved in immune functions based on the literature, to account the increased drift effects for the lower effective population sizes in island species. We used synonymous and 13 14 non-synonymous variations to estimate the selection pressure acting on immune genes. BDs and TLRs have higher ratios of non-synonymous over synonymous polymorphisms (Pn/Ps) 15 16 than randomly selected control genes suggesting that they evolve under a different selection regime than non-immune related genes. However, simulations analyses show that this is 17 unlikely to be explained by ongoing positive selection or balancing selection. For the MHC 18 evolving under balancing selection, we used simulation to estimate the impact of population 19 20 size variation. We found a significant effect of drift on immune genes of island species leading 21 to a reduction in genetic diversity and efficacy of selection. However, the intensity of relaxed 22 selection was not significantly different from control genes, except for MHC class II genes. 23 These genes exhibit a significantly higher level of non-synonymous loss of polymorphism than expected assuming only drift and an evolution under frequency dependent selection, possibly 24 due to a reduction of extracellular parasite communities on islands. Overall, our results showed 25 26 that demographic effects lead to a decrease in the immune functions of island species, but the 27 relaxed selection caused by a reduced parasite pressure may only occur in some immune genes 28 categories.

Keywords: genetic drift, island evolution, immunity, Toll-Like Receptors, Beta-Defensins,
 major histocompatibility complex, molecular evolution, population genomics

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32 Introduction

Island colonizers face new communities of competitors, predators and parasites in a small area 33 34 with limited resources, which generally result in high extinction rates of colonizers (Losos and Ricklefs, 2009). Oceanic island faunas are characterized by a low species richness, coupled 35 with high population densities for each species (MacArthur and Wilson, 1967; Warren et al., 36 2015) - which translates in communities with, on average, low levels of inter-specific 37 interaction and high levels of intra-specific competition (but see Rando et al., 2010 for an 38 39 example of character displacement due to competition among island finch species). These 40 shared island characteristics are thought to underlie the evolution of convergent phenotypes, in 41 what is called the 'island syndrome' (Baeckens and Van Damme, 2020). Convergence has been 42 documented in multiple traits, such as size modification (dwarfism or gigantism; Lomolino, 43 2005), reduction of dispersal (Baeckens and Van Damme, 2020) shift towards K life history strategies (Boyce, 1984; Covas, 2012; MacArthur and Wilson, 1967), evolution of generalist 44 45 traits (Blondel, 2000; Warren et al., 2015), or changes in colour and acoustic signals 46 (Doutrelant et al., 2016; Grant, 1965).

47 Reduced immune function has also been hypothesized as an island syndrome trait, directly 48 linked to reduced parasite pressure on islands (Lobato et al., 2017; Matson and Beadell, 2010; 49 Wikelski et al., 2004). Island parasite communities are i) less diverse (Beadell et al., 2006; 50 Illera et al., 2015; Loiseau et al., 2017; Maria et al., 2009; Pérez-Rodríguez et al., 2013), and 51 ii) could be less virulent due to the expansion of the ecological niche expected by the theory of 52 island biogeography. In fact, island parasites are generally more generalists than their mainland 53 counterparts, which could lead to a reduced virulence due to the trade-off between replication 54 capacity and resistance against host immune defenses (Garamszegi, 2006; Hochberg and Møller, 2001; Pérez-Rodríguez et al., 2013). Overall, a reduction of parasitic pressure should 55 lead to a weakening of the immune system due to the costs of maintaining efficient immune 56 57 functions (Lindström et al., 2004; Matson and Beadell, 2010; Wikelski et al., 2004). Such 58 reduction may have important implications for the ability of these populations to resist or tolerate novel pathogens. The introduction of avian malaria in the Hawaiian archipelago, and 59 the subsequent extinctions and population declines of many endemic species is the most 60 emblematic example (Van Riper III et al., 1986; Wikelski et al., 2004). 61

Immunological parameters, such as blood leukocyte concentration, antibodies or other immune
 proteins (e.g. haptoglobin), hemolysis, and hemagglutination (Lee et al., 2006; Matson and

64 Beadell, 2010) may serve as proxies to determine population immune functions. To date, the majority of studies that focused on island avifauna have found ambiguous results, with either 65 no support for a reduced immune response on island species (Beadell et al., 2007; Matson, 66 2006), or contrasted results, such as a lower humoral component (total immunoglobulins) on 67 68 islands, but a similar innate component (haptoglobin levels) between island and mainland 69 species (Lobato et al., 2017). The use of immune parameters as proxies of immune function is 70 fraught with difficulties (Lobato et al., 2017). The study of molecular evolution of immune 71 genes therefore represents an alternative strategy to tackle this question. However, it is 72 necessary to distinguish neutral effects, the demographic effects resulting from island colonization, from selective ones, the potential relaxation of selection pressures due to the 73 74 changes in the pathogen community.

75 First, the bottleneck experienced by species during island colonization leads to a decrease in genetic variability (Frankham, 1997). A reduced genetic diversity at loci involved in immunity 76 77 should have a direct implication on immune functions (Hale and Briskie, 2007 but see; Hawley et al., 2005; Spurgin et al., 2011). Second, small population sizes increase genetic drift, which 78 79 may counteract the effect of natural selection on weakly deleterious mutations (Ohta, 1992). 80 Several recent studies found a greater load of deleterious mutations in island species (Kutschera 81 et al., 2020; Leroy et al., 2021b; Loire et al., 2013; Robinson et al., 2016; Rogers and Slatkin, 82 2017). Finally, it is necessary to differentiate genes involved in the innate versus the acquired immune response. The innate immune response is the first line of defense and is composed of 83 phagocytes, macrophages and dendritic cells. These cells allow non-specific recognition of 84 85 pathogens (Akira, 2003; Alberts et al., 2002). For example, Toll-Like Receptors (TLR; transmembrane proteins) trigger a chain reaction leading to the production of various 86 87 substances, including antimicrobial peptides such as beta-defensins (BD) that have active 88 properties in pathogen cell lysis (Velová et al., 2018). On the other hand, the acquired immune 89 system allows a specific response, characterized by immune memory. Major Histocompatibility Complex (MHC) genes code for surface glycoproteins that bind to antigenic 90 peptides, and present them to the cells of the immune system; class I and II genes ensure the 91 92 presentation of a broad spectrum of intra- and extracellular-derived peptides, respectively (Klein, 1986). Although all these genes are directly involved in the identification and 93 neutralization of pathogens, previous studies found that they evolve under different selection 94 regimes: TLRs and BDs are under purifying selection which usually results in the selective 95

96 removal of deleterious alleles and stabilizing selection (Grueber et al., 2014; van Dijk et al.,

97 2008), whereas MHC genes are under balancing selection (Bernatchez and Landry, 2003).

98 Recent studies on birds (Gonzalez-Quevedo et al., 2015a, 2015b), amphibians (Belasen et al., 2019), and lizards (Santonastaso et al., 2017) found that the demographic history of island 99 populations led to the loss of genetic variation at immune genes involved in pathogen 100 recognition, such as TLRs and MHC. For example, (Santonastaso et al., 2017) demonstrated 101 102 that the polymorphism pattern in MHC genes and microsatellites covary positively with island 103 area in *Podarcis* lizards, suggesting a dominant role for genetic drift in driving the evolution 104 of the MHC. Gonzalez-Quevedo, et al. (2015a) found a similar pattern comparing TLR and 105 microsatellite polymorphism in the Berthelot pipit, Anthus berthelotii, an endemic species from 106 Macaronesia, supporting a predominant role of genetic drift in TLR evolution. However, these 107 studies did not explicitly test the hypothesis of a relaxed selection pressure on islands imposed 108 by an impoverished parasite community. All other things being equal, it is expected that the 109 polymorphism pattern of a coding sequence decreases with population size (Buffalo, 2021; 110 Leroy et al., 2021b). Therefore, a decrease in polymorphism with population size could not be 111 taken as a proof of a relaxation in the selection pressure.

Here, we study a dataset of 34 bird species (20 insular and 14 mainland species; Figure 1) combining the 24 species of Leroy et al. (2021b) and 10 newly generated by targeted-capture sequencing (Table 1). To be able to demonstrate a change in natural selection, a traditional approach is to contrast polymorphism of synonymous sites (Ps) with polymorphism of nonsynonymous sites (Pn). Synonymous mutations refer to mutations that do not alter amino acid sequences, whereas non-synonymous mutations do.

Following population genetic theory, in a diploid population, $Ps = 4 Ne \mu$ and $Pn = 4 Ne \mu f$, 118 119 where Ne is the effective population size, μ is the mutation rate and f is a function that integrates 120 the probability of an allele to segregate at a given frequency. f depends on the distribution of 121 the fitness effect (DFE) of mutations (Eyre-Walker and Keightley, 2007). This distribution 122 scales with Ne as the fitness effect is dependent on Ne multiplied by the coefficient of selection s (Kimura, 1962). The nearly-neutral theory predicts that the DFE includes a large proportion 123 124 of mutations with a Ne*s close to 0 (Ohta, 1992). As a consequence, an increase of Ne will lead to an increase of the fitness effect of weakly deleterious mutations, in such a way that these 125 126 mutations will be more easily removed from the population by natural selection, therefore 127 reducing Pn relative to Ps leading to a negative correlation between Pn/Ps and Ps (through Ne;

Welch et al., 2008). The presence of linked positively selected mutations does not change
qualitatively this relationship (Castellano et al., 2018; Chen et al., 2020 and our simulations
below).

131 Shifts in the parasitic community on islands are expected to have an impact on the ratio Pn/Ps 132 of immune genes. However, the fixation probability depends on the product Ne s, and a variation in Ne is also expected to impact the efficacy of selection and thus the ratio Pn/Ps 133 134 across the entire transcriptome, particularly in the presence of slightly deleterious mutations 135 (Charlesworth and Eyre-Walker, 2008; Leroy et al., 2021b; Loire et al., 2013; Ohta, 1992). In addition, due to their lower population sizes, island birds compared to continental species 136 137 exhibit a genome-wide reduction in genetic diversity and efficacy of selection (Kutschera et 138 al., 2020; Leroy et al., 2021b). Therefore, we expect a similar reduction in immune genes 139 diversity even without any change in the parasite pressure.

140 To disentangle the effect of population size from a change in parasite pressure and estimate the impact of demography on the efficacy of selection, we randomly selected protein-coding genes 141 142 (i.e., control genes) involved in various biological functions (Fijarczyk et al., 2016; Leroy et al., 2021b). The selection pressure acting on the randomly selected control genes is expected 143 144 to be similar between island and mainland bird species. Therefore, the variation of Pn/Ps of the 145 control genes is only dependent on the variation of Ne. In contrast, if a reduced parasite pressure on islands directly impacts the evolution of immune genes, the Pn/Ps of immune genes is 146 147 expected to show a larger variation between island and continental species than the control 148 genes. More specifically, for genes under purifying selection, non-synonymous weakly 149 deleterious mutations, normally eliminated under strong selection, would be maintained, 150 leading to an increase of Pn/Ps. By contrast, for genes under balancing selection, non-151 synonymous advantageous mutations, normally maintained in the polymorphism under strong 152 selection, would be fixed or eliminated leading to a decrease of Pn/Ps (Figure 2).

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159 Figure 1: Phylogeny based on mitochondrial genes of species from the dataset reconstructed by 160 maximum likelihood method (IQTREE model GTR+Gamma). Species names in yellow indicate island 161 species, and in green, mainland species. Ultrafast bootstrap values are provided in the supplementary methods. Some relationships are poorly supported. Bird representations are not to scale. Photos from 162 163 top to bottom : P. major, C. caeruleus, P. trochilus, Z. borbonicus, T. pelios, F. albicollis, C. olivacea, 164 P. acuticauda, P. grandis, F. coelebs, C. fusca, G. conirostris. Photo credits: A. Chudý, F. Desmoulins, 165 E. Giacone, G. Lasley, Lianaj, Y. Lyubchenko, B. Nabholz, J.D. Reynolds, K. Samodurov, A. 166 Sarkisyan, Wimvz, Birdpics, T. Aronson, G. Lasley, P. Vos (iNaturalist.org); M. Gabrielli (Zosterops 167 borbonicus).

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173 Figure 2: Conceptual diagram showing the expected results under the hypothesis of a relaxation in the 174 selection pressure of the immune genes in island species due to a change in the parasitic community. 175 A) Genes evolving under purifying selection where control genes are randomly selected protein-coding 176 genes. B) Genes evolving under balancing selection where controls are obtained from SLiM simulations 177 of genes evolving under the same balancing selection but different population size. Under the hypothesis 178 of a relaxed selection as a consequence of the reduced diversity of pathogens on island ecosystems, the 179 difference in Pn/Ps between categories (Δ Pn/Ps) is expected to be different between species' origin, 180 leading to a statistical interaction between gene categories and origin.

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182 Methods

183 Dataset

Alignments of Coding DNA Sequences (CDS) of individuals from 24 species were obtained from Leroy et al. (2021b). In addition, data for ten other species (six and four from islands and mainland, respectively) were newly generated for this study by targeted-capture sequencing. Blood samples and subsequent DNA extractions were performed by different research teams. The complete dataset consisted of 34 bird species (20 and 14 insular and mainland species respectively; Table 1; Figure 1). We filtered alignments in order to retain only files containing a minimum of five diploid individuals per site (Table 1).

Sequence enrichment was performed using MYBaits Custom Target Capture Kit targeting 21 immune genes: 10 Toll-Like receptors (TLR), 9 Beta Defensins (BD), 2 Major Histocompatibility Complex (MHC) and 97 control genes (see below). We followed the manufacturer's protocol (Rohland and Reich, 2012). Illumina high-throughput sequencing using a paired-end 150 bp strategy was performed by Novogene (Cambridge, UK).

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197	Table 1: List of species and sampling localities, along with the type of data obtained and the
198	number of individuals (N).

Species	Origin	Island/Country	N	Reference genome	Reference for population genomics data	Type of data	
Cyanistes teneriffae palmae	Island	La Palma	15				
Cyanistes teneriffae teneriffae	Island	Tenerife	14	(This study)	(Mueller et al., 2016)	Capture	
Cyanistes caeruleus	Mainland	France	15	(This study)			
Parus major	Mainland	Europe	10	Parus major (Laine et al., 2016)	Parus major (Laine et al., 2016) (Corcoran et al., 2017)		
Phylloscopus trochilus	Mainland	Europe	9	<i>Phylloscopus trochilus</i> (Lundberg et al., 2017)	(Lundberg et al., 2017)	Whole genome	
Zosterops virens	Mainland	South Africa	7				
Zosterops olivaceus	Island	Réunion	15	Zosterops borbonicus	(Laroy et al. 2021b)	Whole conomo	
Zosterops mauritianus	Island	Mauritius	9	(Leroy et al., 2021a)	(Lefoy et al., 2021b)	whole genome	
Zosterops borbonicus	Island	Réunion	25				
Ficedula semitorquata	Mainland	Europe	20				
Ficedula albicollis	Mainland	Europe	20	Ficedula albicollis	(Ellermon et al. 2012)	XX /1 1	
Ficedula speculigera	Mainland	Nord Africa	20	(Ellegren et al., 2012)	(Ellegren et al., 2012)	whole genome	
Ficedula hypoleuca	Mainland	Europe	20				
Turdus olivaceofuscus	Island	São Tomé	15	Turdus pelios	This starder	Capture	
Turdus pelios	Mainland	Gabon	15	(This study)	This study		
Cyanomitra olivacea	Island	Príncipe	15	Cyanomitra olivacea	This starder	Contract	
Cyanomitra olivacea	Mainland	Gabon	15	(This study)	This study	Capture	
Ploceus grandis	Island	São Tomé	13			1	
Ploceus princeps	Island	Príncipe	13	(This study)	This study	Capture	
Ploceus nigerrimus	Mainland	Cameroon Gabon	14	(This study)			
Poephila acuticauda acuticauda	Mainland	Australia	10	Taeniopygia guttata	(Singhal at al. 2015)	Whole conome	
Taeniopygia guttata castanotis	Mainland	Australia	19	(Warren et al., 2010)	(Siliginal et al., 2013)	whole genome	
Fringilla teydea	Island	Tenerife	10	Eminailla agalaha			
Fringilla canariensis palmae	Island	La Palma	15	(Requerda et al. 2021)	(Leroy et al., 2021b)	Whole genome	
Fringilla coelebs	Mainland	Spain	9	(Recuciua et al., 2021)		-	
Certhidea olivacea	Island	Santiago (Galápagos)	5				
Certhidea fusca	Island	San Cristobal (Galápagos)	10			l	
Certhidea fusca	Island	Española (Galápagos)	10				
Geospiza difficilis	Island	Pinta(Galápagos)	10	~			
Platyspiza crassirostris	Island	Santa Cruz (Galápagos)	5	(Zhang et al., 2012)	(Lamichhaney et al., 2015)	Whole genome	
Pinaroloxias inornata	Island	Coco (Galápagos)	8				
Camarhynchus pallidus	Island	Santa Cruz (Galápagos)	5				
Geospiza difficilis	Island	Wolf (Galápagos)	8				
Geospiza conirostris	Island	Española (Galápagos)	10				

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200 Newly generated draft genome sequence

201 We generated whole genome sequences at moderate coverage (~40X) for Turdus pelios,

202 Ploceus cucultatus and Cyanomitra olivacea (from Gabon). Library preparation from blood

DNA samples and Illumina high-throughput sequencing using a paired-end 150 bp strategy were performed at Novogene (Cambridge, UK). Raw reads were cleaned using FastP (vers. 0.20.0; Chen et al., 2018). Genomes assemblies were performed using SOAPdenovo (vers. 2.04) and Gapcloser (v1.10) (Luo et al., 2012) with parameters "-d 1 -D 2" and a kmers size of 33. Protein annotation was performed by homology detection using genBlastG (She et al., 2011; http://genome.sfu.ca/genblast/download.html) and the transcriptome of the collared flycatcher (*Ficedula albicollis*; assembly FicAlb1.5; Ellegren et al., 2012) as reference.

210 *Capture data processing*

Reads from targeted-capture sequencing were cleaned with FastP (vers. 0.20.0; Chen et al., 211 212 2018). Reads of each individual were mapped respectively to the nearest available reference genomes using bwa mem (vers. 0.7.17; Li, 2013; Table 1), with default parameters. Samtools 213 214 (vers. 1.3.1; Li et al., 2009) and Picard (vers. 1.4.2; Picard Toolkit 2019) were used to convert the mapping files, order and index reads according to their position on the chromosomes (or 215 216 scaffolds) of the reference genomes or on the draft genomes generated in this study for *Ploceus*, Cyanomitra and Turdus. Duplicate reads were marked using MarkDuplicates (vers. 1.140; 217 Picard Toolkit 2019). SNP calling was performed with Freebayes (vers. 1.3.1; Garrison and 218 219 Marth, 2012). Freebayes output file (VCF file) was converted to a fasta file by filtering out 220 sites with a minimum quality of 40 and a sequencing depth between 10 and 1000X (sites outside 221 these thresholds were treated as missing data, i.e., 'N'). CDS were then extracted from the 222 alignments using the coordinates of the annotations (gff files). CDS were aligned using 223 MACSE (vers. 2.03; Ranwez et al., 2011) to prevent frameshift mutation errors and GNU-224 parallel (Tange, 2018) was used to parallelise the computation.

225 Selection and identification of immune and control genes

We defined several groups of immune genes to compare with the control genes. The control group consisted of 97 protein-coding genes randomly selected in the genome of *Zosterops borbonicus* (Leroy et al., 2021a). These control genes allowed the estimation of the average selection pressure that a gene, not involved in the immune response, undergoes in the genome. These genes are single copy (absence of paralogue) and have a variable GC content representative of the whole transcriptome.

For the immune genes, we selected three sets of genes from i) a limited set of genes (Core Group) where functions are unambiguously related to immunity, and ii) two larger sets of genes (Database-group & Sma3s-group), obtained through an automatic annotation pipeline.

235 The Core Group included MHC class I and class II genes, 10 Toll-Like Receptors (TLRs; 236 Velová et al., 2018) and 9 Beta Defensins (BD; Chapman et al., 2016). The Database group included genes identified by Immunome Knowledge Base (Ortutay and Vihinen, 2009, 237 238 http://structure.bmc.lu.se/idbase/IKB/; last access 04/02/2020) and InnateDB (Breuer et al., 239 2013, http://www.innatedb.com; last access 04/02/2020). We also added a set of genes for which the genetic ontology indicated a role in immune functions. To do so, we used the chicken 240 241 (Gallus gallus) annotation (assembly GRCg6a downloaded from Ensembl database in March 242 2020; https://www.ensembl.org/). We identified genes with the terms "immun*" or 243 "pathogen*" in their Gene Ontology identifiers description (directory obtained from 244 http://geneontology.org/). This set included 2605 genes considered to be involved in immunity, 245 although some may be only indirectly involved in immunity or have a small impact on immune functions. Finally, the third set of genes (Sma3s-group) has been built up through the Sma3s-246 247 group program (vers. 2; Munoz-Mérida et al., 2014). This program annotated sequences in 248 order to be associated with biological functions through gene ontology identifiers. The 249 annotation of the genome of F. albicollis allowed us to identify 3136 genes associated with the 250 genetic ontology "immune system processes". Like for the Database group, this set may include 251 genes with various functions in the immune response. It should be noted that Sma3s-group and Database-group are not mutually exclusive, and some genes are present in both groups. An 252 253 analysis was performed to identify and exclude genes under balancing selection from Database-254 group and Sma3s-group sets using BetaScan (vers. 2; Siewert and Voight, 2020), due to the 255 potentially antagonistic responses of these genes. Very few genes (only 2 and 3 genes from Database-group and Sma3s-group sets) were identified and removed from the analysis (see 256 257 Detection of genes under balancing selection in Supplementary Methods).

258 Test for contamination and population structure

We use the program CroCo (vers. 1.1; Simion et al., 2018) to identify candidates for crossspecies contamination (see supplementary materials for details). Overall, we did not detect a clear case of cross-species contamination in our dataset (Figure S1). Contigs identified as potential contamination always involve a pair of species belonging to the same genus. In this

263 case, contamination could be difficult to identify due to the low genetic divergence between264 species.

For the newly sequence becies, we also performed PCA analyses on using allele frequencies of control genes. We use the function dudi.pca of adegenet R packages (Jombart and Ahmed, 2011). This analysis aims to check for population structure and to detect potentially problematic individuals (i.e., contaminated individuals). This analysis led to the exclusion of 4 individuals (*Ploceus princeps* P6-174; *P. grandis* ST10_094; *P. nigerrimus* G3_016; *C. teneriffae* TF57) for which we suspected contamination. Otherwise, no extra population structure was detected (Figure S2-S4).

272 Hidden paralogy

273 We compute the statistic Fis = 1-HO/He where H0 is the average number of heterozygous 274 individual observed (H0 = #heterozygous / n ; where n is the sample size) and He is the 275 expected number of heterozygous individuals at Hardy-Weinberg (HW) equilibrium (He = 276 (n/(n-1) 2 * p * (1-p))*n where n is the sample size and p the allele frequency of a randomly 277 chosen allele). Fis varies between -1 and 1 with positive value representing excess of 278 homozygous individuals and negative value representing excess of heterozygous individuals 279 compared to the HW proportions. Gene with high value of nucleotide diversity (Pi) and 280 negative value of Fis could represent potential cases where hidden paralogous sequences have 281 not been separated and where all the individuals present heterozygous sites in the positions 282 where a substitution occurred between the paralogous copies. Five TLR21 genes appear 283 problematic (Pi > 0.01 and Fis < -0.5; Figure S5) and were excluded from further analyses.

284 The MHC genes are more difficult to analyse. Indeed, heterozygosity could be comparable to 285 divergence under balancing selection. This makes the identification of orthologs very difficult. 286 We identify a variable number of genes among species (from 1 to 10 genes for MHC class I 287 and MHC class II). We checked the sequence similarity for the 10 copies of the MHC class II 288 in F. albicollis and the 7 copies of the MHC classI genes in C. caeruleus using cd-hit (Fu et 289 al., 2012). For MHC class II, sequence divergences are always higher than 15% indicating that 290 reads will likely be correctly assigned to their corresponding gene copy. For MHC class I, 291 sequence identities could be as high as 95%. In this case, we rely on the fact that the reads from 292 very similar paralogous copies will not be confidently assigned to a gene copy sequence by the 293 mapping software. This will lead to a low mapping score quality and are likely to be discarded

during the genotype calling procedure. For example, 3 out of 7 genes of the *Cyanistes* MHC
 class I genes could not have been correctly genotyped and are missing from our final dataset.

296 Data Analysis

297 SLiM simulations

We use SLiM (vers. 3.3.2; Haller and Messer, 2017) to estimate the impact of demographic 298 299 changes on polymorphism patterns under various selection regimes. The following parameters were used in all simulations. Sequences of 30kb with a mutation = f 300 $4.6e^{-9}$ 301 substitutions/site/generation were simulated (Smeds et al., 2016). Recombination was set to be 302 equal to mutation rate. Introns/exons pattern was reproduced by simulating fragments of 3kb 303 separated by one bp with a very high recombination rate of 0.1 rec./site/generation. Five types 304 of mutations were possible: i) neutral synonymous mutations, ii) non-synonymous mutations with a Distribution of Fitness Effect (DFE) following a gamma law of mean = -0.025 and shape 305 306 = 0.3, which corresponds to the DFE estimated in Passerines by Rousselle et al. (2020), iii) 307 codominant non-synonymous mutations positively selected with s = 0.1, iv) non-synonymous 308 mutations under balancing selection with an effect on fitness initially set at 0.01 but re-309 estimated by the program at each generation according to the mutation frequency in the 310 population, thus including a frequency-dependent effect and v) non-synonymous mutations 311 under overdominance using with a dominance coefficient of 1.2.

We simulate a coding sequence organization where positions one and two of the codons were considered as non-degenerated sites, with the non-synonymous types of mutations previously described are possible in various proportions. The third position was considered as completely neutral where only synonymous mutations could appear.

In the absence of control genes evolving under balancing selection, we use SLiM to generate a set of control genes for this category. We simulate two populations of 270,000 and 110,000 individuals, representing mainland and island effective population size respectively.

We also explore the effect of positive and balancing selection on the pattern of Ps and Pn/Ps in a population of size 50,000, 110,000, 270,000 and 500,000. In order to speed up the computational time, we reduce the population size by a factor 100 and rescale mutation rate, recombination rate and selection coefficient accordingly.

All the details of the simulation parameters, calculations of non-synonymous polymorphism rate (Pn) and synonymous polymorphism rate (Ps) of simulated sequences, as well as SLiM command lines are provided in Supplementary Methods and Supplementary Materials.

326 Polymorphism analyses

Synonymous (Ps) and non-synonymous (Pn) nucleotide diversity were estimated from 327 328 seq stat coding written from the Bio++ library (Available as Supplementary data; Guéguen et 329 al., 2013). The mean Pn/Ps was computed as the sum of Pn over the sum of Ps (Wolf et al., 330 2009). Ps of concatenated sequences of control genes were estimated for each species of our 331 dataset. For the whole-genome sequence species, we compared the Pn/Ps and Ps estimated 332 obtained using the 97 control genes with the values from Leroy et al., (2021b; ~5000 genes 333 used in their study). Pn/Ps and Ps correlations showed a R² of 0.6 and 0.95 respectively (Figure 334 S6). Thus, the 97 control genes used in our study were representative of a larger set. This 335 allowed us to identify *Phylloscopus trochilus* as an outlier. Unlike for all other species (e.g. 336 Fringilla coelebs, Figure S7), synonymous polymorphism level was correlated to the amount 337 of missing data in *P. trochilus* alignments (Figure S7). As such, we excluded *P. trochilus* from 338 further analysis.

The mean Pn/Ps, calculated from the concatenated sequences of genes from the same gene class (control genes; BD; TLR; MHC I; MHC II; Database-group; Sma3s-group), was estimated for each bird species. Alternative transcripts were identified based on the genomic position in the GFF file. If several transcripts were available, one transcript was randomly selected. Pn/Ps estimates based on less than four polymorphic sites were excluded from the analysis, as were those with no polymorphic non-synonymous sites.

345 Statistical analyses

346 To estimate the impact of demographic history on genome-wide polymorphism of island 347 species and the potentially reduced constraints on their immune genes, we computed the ratio 348 of non-synonymous nucleotide diversity over synonymous nucleotide diversity (Pn/Ps). A 349 linear mixed model was performed, using the Pn/Ps ratio as dependent variable and, as 350 explanatory variables, the mainland or insular origin of species as well as the category of genes 351 (packages lme4 and lmerTest (Bates et al., 2012; Kuznetsova et al., 2017)). In order to take the 352 phylogenetic effect into account, the taxonomic rank "family" was included as a random effect 353 in the model. We also used a generalized linear mixed model (using the function glmer of the

354 package lme4) with the family "Gamma(link="log")" which leads to the same results (Figure 355 S15 to S24). Five linear mixed models were defined i) null model, ii) model with only the 356 origin parameter, iii) model with only the gene category parameter, iv) model using both origin 357 and gene category parameters, and finally v) model including those two parameters and the 358 interaction effect. In some cases, the phylogenetic effect was difficult to estimate because the number of species per family was reduced to one. In that case, we chose to reduce the number 359 360 of families by grouping Turdidae with Muscicapidae, Nectariniidae, and Estrildidae with 361 Ploceidae and Fringillidae within Thraupidae. The results obtained with these family groupings 362 were similar to the original model (Table S1), except when stated. The categories Databasegroup and Sma3s-group were tested separately from the Core group because they contained 363 364 hundreds of genes annotated using the automatic pipeline that were only available for species with genome wide data. Database-group and Sma3s-group were not analysed simultaneously 365 366 because they contained a partially overlapping set of genes. Finally, genes evolving under purifying selection and genes evolving under balancing selection were also analysed 367 368 separately. Model selection was based on two methods. First, we use the difference in corrected 369 Akaike Information Criterion ($\Delta AICc$) calculated using the qpcR package (Spiess and Spiess, 370 2018). Second, a model simplification using an ANOVA between models was also performed.

371 We also tested an alternative model using the difference between Pn/Ps of immune genes and 372 control genes (Δ Pn/Ps) as dependent variable, and species origin as explanatory variable. Under 373 the hypothesis of a relaxation in selection pressure on islands due to a change in the parasite 374 community, we expect the $\Delta Pn/Ps$ to be higher on island species compared to the mainland ones and, therefore, the species origin (i.e., mainland or island) to be significant. In this model, 375 we used the Phylogenetic Generalized Least Squares model (PGLS; implemented in the "nlme" 376 377 packages; Pinheiro et al., 2017). This model assumes that the covariance between species follows a Brownian motion evolution process along the phylogeny (implemented using the 378 379 "corBrownian" function from the "ape" package; Paradis and Schliep, 2019). The species phylogeny was estimated using mitochondrial genes and a maximum likelihood inference 380 381 implemented in IQTREE (model GTR+Gamma and ultrafast bootstrap; Nguyen et al., 2014; 382 median of 11,134 bp analysed per species). The phylogeny with the bootstrap support is 383 provided as supplementary material.

All the statistical analyses were performed using R (R Core Team, 2018), and dplyr package (Wickham, 2016). Graphical representations were done using ggplot2, ggrepel, ggpubr and ggpmisc (Aphalo, 2020; Kassambara, 2018; Slowikowski et al., 2018; Wickham, 2016).

387

388 Results

For the 150 individuals (10 species with 15 individuals each) for which we generated new data by targeted capture sequencing, an average of 3.3 million paired-ends reads per individual was generated (Table S1). After mapping, genotyping and cleaning, we analysed 86 control and 16 immune genes on average per species, out of the 141 targeted genes (120 control and 21 immune related genes; Table S4). For the species with whole-genome sequences, we analysed 106 control and 20 immune genes on average per species, out of the 141 targeted genes, and 875 and 688 genes on average in the Database-group and Sma3s-group respectively (Table S4).

For the species for which full genome sequences were available, the Ps and Pn/Ps estimatedusing the control genes reflect the Ps and Pn/Ps of the whole transcriptome (Figure S6).

398 Population genetics of BD and TLR Immune genes



In order to characterize the selection regimes shaping the BD and TLR polymorphisms, we first analyze the variation of Pn/Ps ratios among gene categories using a linear mixed model.

401

402 <u>Figure 3:</u> Pn/Ps according to species origin (mainland in green and insular in orange) for
 403 different gene categories under purifying selection. The number of species (N), and the mean
 404 Pn/Ps are shown for each modality.

405 Model selection based on AICc as well as model selection approach based on simplification with ANOVA identified the model including the origin (i.e., mainland or island) and gene 406 407 category without interaction (Table 2). In this model, island origin of species is associated with 408 a greater Pn/Ps (0.14 vs. 0.10; Table 3; p < 0.01). Gene categories corresponding to TLRs and 409 BDs showed a significantly higher Pn/Ps than control genes (Table 3; p < 0.001). Our statistical 410 analysis confirms that island birds have a higher Pn/Ps ratio than mainland relatives, in 411 agreement with the nearly-neutral theory of evolution. It also reveals that immune genes have 412 a higher Pn/Ps than randomly selected control genes suggesting that BD and TLR evolve under 413 a different selection regime than non-immune related genes.

414 Next, we investigate the cause of the higher Pn/Ps of immune genes by testing three hypotheses. 415 First, we exclude a bias due to a lower number of genes in immune genes, and therefore higher 416 variance in the estimation of Pn/Ps, in immune genes. Immune genes still have significantly 417 higher Pn/Ps compared to a random subsample of control genes of comparable size (Figure S8 418 & S9). Second, the Pn/Ps of immune genes could be inflated by positive selection. It is well 419 known that immune genes are subject to frequent adaptation due to harm race evolution with 420 pathogens (Enard et al., 2016; Shultz and Sackton, 2019; Velová et al., 2018). We evaluate the 421 effect of positively selected genes on the Pn/Ps using SLiM simulations with both positively 422 and negatively selected mutations. The presence of recurrent positive selection could increase 423 the Pn/Ps leading to a higher Pn/Ps in immune genes if this category is more prone to adaptive 424 evolution (Figure 4A). However, positive selection always leads to a drastic decrease in Ps due to genetic sweep effect at linked sites (Figure 4B). BDs and TLRs have a slightly higher or 425 426 similar Ps than control genes (Figure S9, mean Ps = 0.007, 0.004 and 0.003 for BDs, TLRs and control genes respectively, effect of gene category p < 0.1) and, as a consequence, even if 427 428 positive selection is likely to have impacted the evolution of immune genes, it is not the cause of the higher Pn/Ps observed here. Third, balancing selection could be present, at least 429 430 temporarily, in the evolution of BDs and TLRs genes (Kloch et al., 2018; Levy et al., 2020). Simulation analyses confirm that balancing selection causes an increase of Ps and Pn/Ps (Figure 431 432 4C & 4D). However, a change in effective population size has an opposite effect on the Pn/Ps 433 according to the type of selection. In the presence of slightly deleterious mutations, Pn/Ps 434 decreases with *Ne* whereas it increases in the presence of balancing selection. Island birds have 435 higher Pn/Ps ratios than mainland birds for BDs and TLRs. Therefore, we can rule out balancing selection as the main factor explaining the high Pn/Ps of immune genes because, in 436 437 this case, Pn/Ps of island birds should be lower. The last possible explanation we can think of is a relaxed selection of immune genes. It is likely that immune genes are overall less
constrained than the control genes. It has been shown that evolutionary constraints are more
related to gene expression than to function (Drummond et al., 2005; Drummond and Wilke,
2008) and therefore, functionally important genes could still have a high Pn/Ps.

442 Overall, our analyses do not support a strong impact of ongoing adaptive mutation or balancing 443 selection on BDs and TLRs. However, these immune genes do not evolve as the random genes 444 not involved in immune functions and present a significantly higher Pn/Ps of 0.20 (p < 0.001; 445 Table 3).

446 No evidence of a reduced impact of the parasite communities on the polymorphism pattern = 447 immunes genes in island birds

For BDs and TLRs, the best model selected includes the origin (i.e., mainland or island) and gene category without interaction (see above and Table 2). This model has no interaction between origin and gene categories invalidating the hypothesis of a reduced parasite communities on island (Figure 2).



453 <u>Figure 4:</u> Neutral polymorphism (Ps) and ratio of selected over neutral polymorphism (Pn/Ps)
454 estimated from SLiM simulations. A) Pn/Ps as a function of population size, N and B) Ps as a

function of N. In both A and B, colour indicates the frequency of positively selected mutation compare to deleterious mutation. C) Pn/Ps as a function of N and D) Ps as a function of N. In both C and D, yellow indicates simulations with overdominance mutation (h = 1.2) and negatively selected mutations and green indicates simulations with only negatively selected mutations.

460

<u>Table 2:</u> Statistical model explaining Pn/Ps variation of Toll-Like Receptors, Beta-Defensins
 genes, and control genes. The p-values of ANOVA test between simpler models are not
 reported if a more complex model explains a larger proportion of the variance.

Model		Mod	ANOVA test					
n°	Details	AICc	ΔAICc	Likelihood	n° 1	2	3	4
1	Pn/Ps~ 1+ category +origin+ category *origin	-5.39	8.83	0.01		0.63		
2	Pn/Ps~ 1+ category +origin	-14.22	0	1			0.002	3.71E-05
3	Pn/Ps~1+ category	-11.8	2.42	0.3				
4	Pn/Ps~1+ origin	-6.83	7.39	0.02				
5	Pn/Ps~1	-6.44	7.78	0.02				

464

<u>Table 3</u>: Summary of the best statistical model selected using AICc explaining variation in
Pn/Ps in control genes, Toll-Like receptors and Beta-Defensins genes under purifying selection
with origin, gene category parameters. * indicates significances : * < 0.05; ** < 0.01; *** <
0.001.

Model	Pare	ameters			
WIUGEI	Origin	Category	Estimate	P.value	
Origin	Intercept mainland	Control genes	0.10	2.65E-02	*
and Gene	island		0.14	4.56E-03	**
category		Toll-Like Receptors	0.20	7.43E-05	***
(n°2)		Beta-Defensins genes	0.20	3.16E-04	***

469

For larger sets of genes, identified using an automatic pipeline and gene annotation, model selection based on AICc and simplification with ANOVA (Table S5, S8) identified models that included origine parameters which associated a higher Pn/Ps of at least 0.07 for island species (p < 0.001; Table S6, S7, S9, S10, Figure 5). Selection model by simplification with

ANOVA identified models with interaction effect between origin and gene category associated
with a reduced Pn/Ps for immune genes of island species that invalidate our hypothesis (Table
S7, S10).

477

The alternative statistical approach using the difference between Pn/Ps of immune genes and control genes (Δ Pn/Ps) as dependent variable, and species origin as explanatory variable under a PGLS framework lead to similar results. Island was never associated to a statistically higher Δ Pn/Ps (table S2) providing no support for an increased relaxed selection of immune genes in island species.



483



487 Genes under balancing selection

First, we estimated the effect of population size variation on the Pn/Ps of the genes evolving under balancing selection by simulating sequences under frequency dependent or overdominance selection using SLiM (see Methods and Supplementary Methods). The simulation under frequency dependence selection revealed an average Pn/Ps equal to 0.8 for island species and 1.2 for mainland species (Figure 6). Under overdominance, simulated sequences from island and mainland populations respectively have an average Pn/Ps equal to 0.54 and 1.03 (Figure 6).



496 <u>Figure 6:</u> Boxplot of Pn/Ps according to species origin (mainland in green and insular in orange)
 497 for different gene categories under balancing selection. The number of species (N), and the
 498 mean Pn/Ps are shown for each modality. The control groups correspond to the results obtained
 499 from simulated sequence via SLiM (see Methods and Supplementary Methods Simulation of
 500 control genes under balancing selection).

Using simulations under frequency dependence selection as well as simulations under the 501 502 overdominance, model selection by AIC identifies the model with origin as the best, contrary 503 to the method by simplification with ANOVA which identified the full model therefore including significant interaction between origin and genes category (Table 4). This interaction 504 effect is significant for the MHC II (p < 0.05, Table S12) but not for MHC I. As expected, 505 island species have a significantly lower Pn/Ps in MHC genes compared to mainland species 506 (p < 0.01; except for the full model based on control genes evolving under overdominance 507 Table S12). 508

<u>Table 4</u>: Statistical model explaining Pn/Ps variation of genes under balancing selection (i.e
 MHC class I and II), and simulated sequences under i) frequency dependence or ii)
 overdominance. The p-values of ANOVA test between simpler models are not reported if a
 more complex model explains a larger proportion of the variance.

]	Model selection by AIC				AN	ANOVA test			
Type of balancing selection	n°	Details	AICc	ΔAICc	Likelihoo d	n°1	2	3	4
Frequency	1	Pn/Ps~1+ category +origin+ category	157.17	5.62	0.06		0.019		

dependence		*origin				
	2	Pn/Ps~1+ category +origin	157.85	6.31	0.04	
	3	Pn/Ps~1+ category	187.58	36.04	0.00	
	4	Pn/Ps~1+ origin	151.54	0.00	1.00	
	5	Pn/Ps~1	180.52	28.97	0.00	
	1	Pn/Ps~1+ category +origin+ category *origin	140,56	8,50	0,01	0.024
Overdominance	2	Pn/Ps~1+ category +origin	140,56	8,50	0,01	
	3	Pn/Ps~1+ category	185,91	53,85	0,00	
	4	Pn/Ps~1+ origin	132,05	0,00	1,00	
	5	Pn/Ps~1	177,54	45,49	0,00	

513

514 Discussion

On oceanic islands, the depauperate parasite community is expected to lead to a relaxation of 515 516 selection on the immune system. In this study, we found support for such an effect, but only on 517 MHC class II genes and using simulated sequences under balancing selection as control. No effect was detected for MHC class I genes nor for innate immune genes (TLRs and BDs), 518 evolving under purifying selection. On these gene sets, increased drift effects on island 519 520 populations limit the efficacy of selection in accordance with the nearly-neutral theory (Ohta, 521 1992). The ability to distinguish between the selective and nearly-neutral processes (relaxed 522 selection due to environmental change vs. drift) could only be achieved by our approach of using random genes (i.e., "control genes") to estimate the genome-wide effect of potential 523 variation in effective population size between populations. 524

525 *Effects of effective population size variation*

526 Our results support the nearly-neutral theory of evolution for those genes under purifying 527 selection, whereby strong genetic drift acting on small island populations reduces the efficacy 528 of natural selection, leading to an increase in non-synonymous nucleotide diversity compared 529 to the mostly neutral, synonymous nucleotide diversity (i.e., Pn/Ps; Ohta, 1992). This is 530 materialized by a genome-wide increase in frequency of weakly deleterious mutations 531 (Kutschera et al., 2020; Leroy et al., 2021b; Loire et al., 2013; Robinson et al., 2016; Rogers 532 and Slatkin, 2017).

533 For genes evolving under balancing selection, we performed simulations under the hypotheses of overdominance (heterozygote advantage) or frequency dependence (rare-allele advantage). 534 535 Our results showed reduced Pn/Ps for smaller population sizes (Figure 6, S10, S11). This 536 simulation confirmed our expectations (Figure 2) that a reduction in the efficacy of selection 537 results in a decrease in the frequency of non-synonymous polymorphism, as, under normal 538 circumstances, selection maintains those mutations at intermediate frequencies. It also matches 539 what we obtained in the empirical results, where both MHC classes I and II had a reduced Pn/Ps 540 in island birds. This result supports that the fitness effect of having non-synonymous 541 polymorphisms segregating at high frequencies is not strong enough to counteract entirely the 542 effect of genetic drift on islands, therefore extending the nearly-neutral theory to the 543 overdominance type of selection.

544

545 *Effects of selection on immune genes*

For immune genes, we try to characterize the nature of the selection acting on BDs and TLRs 546 genes. Comparing those genes with control genes and using simulations, we were able to rule 547 548 out that directional positive selection and balancing selection had a major impact shaping the polymorphism of these immune genes. In contrast, the pattern of Pn/Ps between island and 549 550 mainland populations is in line with the effect of purifying selection in the presence of slightly deleterious mutation. However, no effect was detected on insular species, beyond what could 551 552 be attributed to genetic drift. This is in line with the result of Gonzalez-Quevedo et al. (2015b) 553 and Grueber et al. (2013) who found that TLR genetic diversity was mostly influenced by 554 genetic drift. At first sight, this result seems not in line with the fact that island parasite 555 communities are less diverse (Beadell et al., 2006; Loiseau et al., 2017; Maria et al., 2009; 556 Pérez-Rodríguez et al., 2013; but see Illera et al., 2015). However, a reduced pathogens number 557 has also been found to be associated with a higher prevalence in birds and reptiles from the 558 Macaronesian archipelago (Illera and Perera, 2020). Therefore, these two patterns, i.e. a less 559 diverse pathogen's community on islands with a higher prevalence, could still imply a strong 560 selection pressure on immune genes.

In contrast, for MHC genes that unambiguously evolve under balancing selection, MHC class II genes presented a reduction in non-synonymous polymorphism larger than the effects of drift alone, when simulated sequences are used as control. This was the only case where a role for relaxed selection pressures in the molecular evolution of immune genes could be invoked.

565 Our results are in accordance with the hypothesis of Lee (2006), which proposes that innate 566 and acquired immunity may exhibit distinct responses to changes in pressures due to different 567 costs and benefits. However, our result contrasts with the study of Santonastaso et al. (2017) 568 that identified no change in selection pressures on MHC II genes in a lizard species, concluding 569 that their evolution was mostly governed by drift. Similarly, Agudo et al. (2011) also found a 570 prominent role for genetic drift over selection in the evolution of MHC II genes in the Egyption 571 vulture (*Neophron percnopterus*).

572 Our results rely on simulations that may be affected by the choice of the parameter values. 573 First, we performed simulations using a fixed effective population size (Ne) estimated from the 574 polymorphism data. Using others values of *Ne* had a weak impact on the relative difference 575 between island and mainland species for the overdominance type of selection (Figure S10, 576 S11). Secondly, we simulated two types of selection, namely overdominance (Doherty and 577 Zinkernagel, 1975) and frequency dependence (Slade and McCallum, 1992), but it has been 578 argued that the maintenance of MHC polymorphism could be the result of fluctuating selection 579 (Hill, 1991). Additionally, recombination and gene conversion has also been put forward as a 580 mechanism responsible for generating diversity (Spurgin et al., 2011). Therefore, our results 581 for the MHC II, which is based on the relative difference between Pn/Ps of island and mainland species comparing empirical and simulated data, should be taken cautiously as their 582 583 significance can be dependent on the specific parameters that we used, although we did our 584 best to select a realistic range of parameters.

585 The observed difference between MHC class I and II could be explained by their different 586 pathogen targets: MHC class I genes are primarily involved in the recognition of intracellular 587 pathogens (Kappes and Strominger, 1988), while MHC class II genes are directly involved in 588 the recognition of extracellular pathogens (Bjorkman and Parham, 1990). These differences could lead to variable selection pressures depending on the extracellular versus intracellular
parasite communities present on islands. In addition, the relaxed selection pressures on MHC
II genes from adaptive immunity is in line with a reduction in acquired immunity parameters
observed by Lobato et al. (2017) that used partly the same sets of species.

593 As a perspective of our work, we should mention that there is an extensive variation in the number of MHC gene copies across the avian phylogeny (Minias et al., 2019; O'Connor et al., 594 595 2020). Particularly, it was recently discovered that Passerines have a very dynamic evolution 596 of duplication/loss events compared to other birds (Minias et al., 2019). Here, we used the two copies of MHC gene I and II currently annotated in the collared flycatcher genome as target 597 598 sequences for our targeted-capture sequencing. The future improvement of genome assembly, 599 thanks to the development of long-reads technology (Peona et al., 2021, 2018), will certainly 600 help to precisely annotate all MHC copies and to study the whole repertoire of MHC genes.

601 *Consequences of drift effect-and selection on immunity*

602 The potential relaxation of the natural selection acting on immune genes in island species is 603 expected to reduce immune functions and increase susceptibility of island populations to 604 pathogens. This is true even if this relaxation is only the consequence of a reduction in the 605 effective population size and not caused by a reduction of the pressure exerted by the parasitic community. This is in line with the results of Hawley et al. (2005) and Belasen et al. (2019) 606 607 who showed that a decrease in diversity of immune loci (MHC II or through immune proxy) 608 was associated with a reduction in immune functions. It should be noted that even if migration 609 rate is reduced on islands, sedentary and endemic island species are not completely free from 610 the exposure of exogen pathogens through migratory birds (Levin et al., 2013).

611 As a final remark, we would like to stress that more research is still needed (i) to ascertain both selection pressures on innate and adaptive immune responses and the load of deleterious 612 613 mutations due to drift, also identified by an increasing body of work (Loire et al., 2013; Robinson et al., 2016; Rogers and Slatkin, 2017; Kutschera et al., 2020; Leroy et al., 2021b), 614 615 and (ii) to describe island parasite communities. To date, most of the studies investigated 616 intracellular parasite communities on islands, and more specifically haemosporidian parasites, 617 avian pox and coccidian parasites (Cornuault et al., 2012; Illera et al., 2015, 2008; Ishtiaq et al., 2010; Loiseau et al., 2017; Martinez et al., 2015; Padilla et al., 2017; Pérez-Rodríguez et 618 al., 2013; Silva-Iturriza et al., 2012), whereas very few evaluated the extracellular parasite 619

diversity, such as helminths (Nieberding et al., 2006) but see the review of Illera and Perera (2020) for reptiles. Metabarcoding of parasites is a new technique to evaluate at the same time both communities of intracellular and extracellular parasites (Bourret et al., 2021) and might therefore be a promising approach to evaluate their communities in island and mainland populations.

625 *Conclusion*

Our comparative population genomics study has investigated the combined effects of drift and 626 627 selection on immune genes from island and mainland passerines. The study of synonymous and non-synonymous polymorphism of these genes confirmed that island species, with smaller 628 629 population sizes than their mainland counterparts, were more impacted by drift, which induces 630 a load of weakly deleterious mutations in their genome. Indeed most of the genes studied here 631 involved in the immune response do not show a statistically different pattern from control 632 genes. Only MHC II genes, involved in the recognition of extracellular pathogens, showed a 633 reduction in their non-synonymous polymorphism in island species. This response, which may be attributed to reduced selection pressures on these genes, could be associated with the 634 635 suspected reduced parasitic communities on islands. The increased load of deleterious 636 mutations as well as the potential relaxed selection pressures on MHC II support the reduced immune functions of island species, which could be added to the list of other convergent 637 638 responses of the island syndrome.

639 Data availability

Datasets, scripts, supplementary figures and texts are available on figshare :
 <u>https://figshare.com/s/ab7004cc2f4415b4058f</u>. The reads newly generated for this study have
 been deposited in the NCBI Sequence Read Archive under the bioproject PRJNA724656.

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