

A whole-genome scan for evidence of positive and balancing selection in aye-ayes (*Daubentonia madagascariensis*) utilizing a well-fit evolutionary baseline model

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The aye-aye (*Daubentonia madagascariensis*) is one of the 25 most endangered primate species in the world, maintaining amongst the lowest genetic diversity of any primate measured to date. Characterizing patterns of genetic variation within aye-aye populations, and the relative influences of neutral and selective processes in shaping that variation, is thus important for future conservation efforts. In this study, we performed the first whole-genome scans for positive and balancing selection in the species, utilizing high-coverage population genomic data from newly sequenced individuals. We generated null thresholds for our genomic scans by creating an evolutionarily appropriate baseline model that incorporates the demographic history of this aye-aye population, and identified a small number of candidate genes. Most notably, a suite of genes involved in olfaction—a key trait in these nocturnal primates—were identified as experiencing long-term balancing selection. We also conducted analyses to quantify the expected statistical power to detect positive and balancing selection in this population using site frequency spectrum–based inference methods, once accounting for the potentially confounding contributions of population history, mutation and recombination rate variation, as well as purifying and background selection. This work, presenting the first high-quality, genome-wide polymorphism data across the functional regions of the aye-aye genome, thus provides important insights into the landscape of episodic selective forces in this highly endangered species.

Keywords: primate; strepsirrhine; selective sweep; balancing selection; genome scan; demography

Introduction

A strepsirrhine endemic to Madagascar and the world's largest nocturnal primate, the aye-aye (Daubentonia madagascariensis) is the only extant member of the Daubentoniidae family, exhibiting a geographical range wider than any other member of the Lemuroidea superfamily (Sterling 1993). However, rapid habitat destruction is thought to have contributed to a severe population decline over the past few decades (Louis et al. 2020; Suzzi-Simmons 2023), along with direct human predation at least partially owing to the regional Malagasy cultural belief that aye-ayes are a harbinger of illness and death (Andriamasimanana 1994). These ongoing trends-coupled with harboring amongst the lowest genetic diversity of any primate measured to date (Perry et al. 2012; Kuderna et al. 2023)-have placed the ave-ave on the list of the 25 most endangered primate species in the world, according to the International Union for Conservation of Nature and Natural Resources Species Survival Commission Primate Specialist Group (Schwitzer et al. 2013; Louis et al. 2020; and see the discussion in Gross 2017). As such, characterizing patterns of genetic variation within aye-aye populations, and the relative influences of neutral and selective processes in shaping that variation, remains of significant importance.

Signatures of episodic selection

Although it is well understood that the demographic history of a population together with the recurrent action of natural selection acts to shape patterns of polymorphism at the DNA sequence level, disentangling the effects of these processes remains an ongoing concern (e.g. Ewing and Jensen 2016; Charlesworth and Jensen 2022; Jensen 2023). Nevertheless, distinguishing these processes is fundamental for gaining an improved understanding of general evolutionary dynamics, inasmuch as it would improve our understanding of the relative importance of adaptive and nonadaptive factors in shaping levels of genetic variation in natural aye-aye populations as well as facilitate the accurate identification of genomic regions that have experienced recent bouts of episodic selection. Existing methods for detecting recent beneficial fixations rely on the changes in patterns of variation at linked sites (i.e. by characterizing the resulting effects of the associated selective sweep), though the nature of these changes will naturally depend on the details of the selective pressure (see the review of Stephan 2019). The term "selective sweep" describes the process whereby a positively selected mutation rapidly increases in frequency and fixes within a population, with linked variation

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following the same trajectory to an extent determined by the level of linkage (Maynard Smith and Haigh 1974 and see the review of Charlesworth and Jensen 2021). The fixation of these linked variants is expected to temporally reduce local nucleotide diversity (Berry et al. 1991). Under a single selective sweep model with recombination, there is also an expectation of a skew in the site frequency spectrum (SFS) toward both high- and low-frequencyderived alleles within the vicinity of the beneficial mutation (Braverman et al. 1995; Simonsen et al. 1995; Fay and Wu 2000). The theoretical expectations under this model of a single, recent selective sweep have been well-described (e.g. Kim and Stephan 2002; Kim and Nielsen 2004), and a composite likelihood ratio (CLR) test was developed by Kim and Stephan (2002) based on these expectations that detects such local reductions in nucleotide diversity and skew in the SFS along a chromosome. This signature is used to identify candidate loci that have experienced the recent action of positive selection by comparing the probability of the observed SFS under the standard neutral model with that under the model of a selective sweep. Subsequent work demonstrated that certain demographic histories may be problematic however, with, for example, severe population bottlenecks often replicating patterns of positive selection (Jensen et al. 2005). Thus, to help reduce this issue, Nielsen et al. (2005) adapted the CLR method for genomewide data utilizing a null model instead derived from the empirically observed SFS, which they termed SweepFinder (along with a more recent implementation, SweepFinder2; DeGiorgio et al. 2016).

Unlike signatures of selective sweeps, those of balancing selection (see the reviews of Fijarczyk and Babik 2015; Bitarello et al. 2023)—a term that encapsulates a variety of selective processes that maintain genetic variability in populations—can potentially last for considerable timescales (Lewontin 1987). Indeed, the temporal history of a balanced allele has been split into multiple phases, with detectable genomic signatures varying in each phase. For example, Fijarczyk and Babik (2015) characterized these phases as recent ($<0.4N_e$ generations), intermediate (0.4–4 N_e generations), and ancient (>4 N_e generations), where N_e is the effective population size. The initial trajectory of a newly introduced mutation under balancing selection is indistinguishable from that of a partial selective sweep (Soni and Jensen 2024), whereby the newly arisen mutation rapidly increases to its balanced frequency, conditional on escaping stochastic loss. The signatures of these partial sweeps include a potential excess of intermediate frequency alleles, extended linkage disequilibrium (LD) owing to the associated genetic hitchhiking effects (see the reviews of Crisci et al. 2013; Charlesworth and Jensen 2021), and weaker genetic structure at genes experiencing balancing selection (Schierup et al. 2000). Once the balanced frequency is reached, the allele under balancing selection fluctuates about this frequency, with recombination breaking up the aforementioned LD patterns (Wiuf et al. 2004; Charlesworth 2006; Pavlidis et al. 2012). If balancing selection persists in species with a divergence time predating the expected coalescent time, the allele under selection may continue to segregate as a trans-species polymorphism (Klein et al. 1998; Leffler et al. 2013). To facilitate detection of long-term balancing selection, Cheng and DeGiorgio (2020) developed a class of CLR-based methods for detecting balancing selection that utilizes a mixture model, combining the expectation of the SFS under neutrality with the expectation under balancing selection, to infer the expected SFS at both a putatively selected site and at increasing distances away from that site, released under the BalLeRMix software package (Cheng and DeGiorgio 2020). As with SweepFinder2, this class of methods utilizes a null model directly derived from the empirical SFS in an attempt to account for

deviations from the standard neutral expectation in a model-free manner. Such approaches have been shown to be well powered in detecting long-term balancing selection (> $25N_e$ generations in age; Soni and Jensen 2024), depending, as they do, on new mutations accruing on the balanced haplotype and thereby generating the expected skew in the SFS toward intermediate frequency alleles.

Inferring positive and balancing selection in nonhuman primates

As one might expect, the majority of scans for positive selection in primates have focused upon human population genomic data. However, a number of studies have found signals of putative positive selection in nonhuman primates, chiefly in the great apes (The Chimpanzee Sequencing and Analysis Consortium 2005; Enard et al. 2010; Locke et al. 2011; Prüfer et al. 2012; Scally et al. 2012; Bataillon et al. 2015; McManus et al. 2015; Cagan et al. 2016; Munch et al. 2016; Nam et al. 2017; Schmidt et al. 2019), as well as in biomedically relevant species such as rhesus macaques (The Rhesus Macaque Genome Sequencing and Analysis Consortium et al. 2007) and vervet monkeys (Pfeifer 2017b), often with contradictory results, likely owing to differing methodological approaches as well as to high false-positive rates related to a neglect of demographic effects. For example, Enard et al. (2010) proposed a variant of the Hudson-Kreitman-Aguadé test (Hudson et al. 1987) and applied it to chimpanzee, orangutan, and macaque population genomic data, finding a high number of orthologous genes exhibiting simultaneous signatures of selective sweeps; conversely, Cagan et al. (2016) utilized a variety of test statistics for detecting positive selection across differing timescales, finding relatively little overlap between species.

Although the first whole-genome, short-read assembly for the aye-aye was published over a decade ago (Perry et al. 2012), and a more recent long-read assembly was made available in 2023 (Shao et al. 2023), the lack of protein-coding gene annotations has greatly limited the ability to scan for recent, episodic selective events in the species. However, the new release of a fully annotated, chromosome-level hybrid de novo assembly (based on a combination of Oxford Nanopore Technologies long reads and Illumina short reads, and scaffolded using genome-wide chromatin interaction data; Versoza and Pfeifer 2024) provides a unique opportunity to identify patterns of selection in this endangered species. In this study, we have performed scans for selective sweeps and balancing selection using SweepFinder2 (DeGiorgio et al. 2016) and BalLeRMix (Cheng and DeGiorgio 2020), respectively, utilizing unique, high-quality, whole-genome, population-level data. Importantly, to account for the confounding effects of demography (Barton 1998; Ewing and Jensen 2014; Poh et al. 2014; Harris and Jensen 2020; and Charlesworth and Jensen 2024), we used the demographic model of Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025), which was generated using nonfunctional regions in the aye-aye genomes that are free from the effects of background selection. This demographic history fits neutral population data exceedingly well and suggests a history in which the aye-aye population size was greatly reduced with first human contact in Madagascar 3,000-5,000 years ago, with an additional decline owing to recent habitat loss over the past few decades. This well-fitting null model is thus utilized here to determine thresholds for positive and balancing selection scans across functional regions in order to avoid extreme false-positive rates (Thornton and Jensen 2007; Poh et al. 2014; Johri, Aquadro et al. 2022; Johri, Eyre-Walker et al. 2022; Johri et al. 2023; Soni et al. 2023)—a particularly important feature in this application given the severe bottleneck history of the species (Terbot, Soni, Versoza, Pfeifer *et al.* 2025). Through this process, we have identified a number of candidate loci with evidence of positive and balancing selection effects and discuss these results in light of the recently available gene annotations.

Materials and methods Animal subjects

This study was approved by the Duke Lemur Center's Research Committee (protocol BS-3-22-6) and Duke University's Institutional Animal Care and Use Committee (protocol A216-20-11). The study was performed in compliance with all regulations regarding the care and use of captive primates, including the US National Research Council's Guide for the Care and Use of Laboratory Animals and the US Public Health Service's Policy on Human Care and Use of Laboratory Animals.

Samples and whole-genome sequencing

We sequenced 4 wild-born aye-aye (*D. madagascariensis*) individuals as well as 1 colony-born aye-aye of wild-born parents using DNA extracted from whole blood samples previously collected at the Duke Lemur Center (Durham, NC, USA). A 150-bp paired-end library was prepared for each sample using the NEBNext Ultra II DNA PCR-free Library Prep Kit (New England Biolabs, Ipswich, MA, USA) and sequenced at high-coverage (>50-fold) on the Illumina NovaSeq platform (Illumina, San Diego, CA, USA). Information regarding samples and their sequencing coverage is provided in Supplementary Table S1.

Calling variant and invariant sites

Calling of variant and invariant sites followed the best practices for nonmodel organisms as described in Pfeifer (2017a) and van der Auwera and O'Connor (2020). In brief, we preprocessed raw reads by removing adapters and low-quality bases from the read-ends using the Genome Analysis Toolkit (GATK) MarkIlluminaAdapters v.4.2.6.1 (van der Auwera and O'Connor 2020) and TrimGalore v.0.6.10 (https://github.com/FelixKrueger/TrimGalore), respectively. We mapped the prepared reads to the aye-aye reference assembly (DMad_hybrid; GenBank accession number: GCA_044048945.1; Versoza and Pfeifer 2024) using the Burrows Wheeler Aligner (BWA-MEM) v.0.7.17 (Li and Durbin 2009) and marked duplicates using GATK's MarkDuplicates v.4.2.6.1. We further refined the read mappings by performing multiple sequence realignments using GATK's RealignerTargetCreator and IndelRealigner v.3.8, recalibrating base quality scores using GATK's BaseRecalibrator and ApplyBQSR v.4.2.6.1, and conducting another round of duplicate removal using GATK's MarkDuplicates v.4.2.6.1. As population genomic resources for aye-ayes are limited, no "gold standard" dataset exists for this recalibration step; instead, we utilized a high-confidence training dataset obtained from pedigreed individuals (see Versoza et al. 2025 for details). We then called variant and invariant sites from high-quality read alignments ("-minimum-mapping-quality 40") using GATK's HaplotypeCaller v.4.2.6.1 with the "pcr_indel_model" parameter set to NONE as a PCR-free protocol was used during library preparation, the "-heterozygosity" parameter set to 0.0005 to reflect speciesspecific levels of heterozygosity (Perry et al. 2013), and the "-ERC" parameter set to BP_RESOLUTION to output both variant and invariant sites. Next, we jointly assigned genotype likelihoods across all 5 individuals at all sites ("-all-sites") using GATK's GenotypeGVCFs v.4.2.6.1, accounting for species-specific levels of heterozygosity as detailed above. Following the GATK Best Practices, we applied a set of site-level "hard filter" criteria (i.e. QD < 2.0, QUAL < 30.0, SOR>3.0, FS>60.0, MQ<40.0, MQRankSum<-12.5, and

ReadPosRankSum < -8.0) to the sites genotyped in all individuals (AN = 10) and applied upper and lower cutoffs on the individual depth of coverage ($0.5 \times DP_{ind}$ and $2 \times DP_{ind}$) to remove regions with an unusual read depth indicative of erroneous calls. The resulting dataset was limited to the autosomes and divided into variant [i.e. single nucleotide polymorphisms (SNPs)] and invariant sites for downstream analyses (Supplementary Table S2).

Generating genome-wide null thresholds for selection inference

We simulated the demographic model of Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025) to generate genome-wide null thresholds for the inference of selection using the coalescent simulator msprime v.1.3.2 (Baumdicker et al. 2022). Although the Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025) model involves 2 populations—one population from northern Madagascar, consisting of 4 individuals previously sequenced at low coverage (Perry et al. 2013), and one population from the rest of the island, consisting of 8 individuals previously sequenced at low coverage (Perry et al. 2013) as well as 5 individuals newly sequenced at high coverage (Terbot, Soni, Versoza, Pfeifer et al. 2025)—we here only consider the single population of newly sequenced individuals (n = 5), given the much higher data quality of this sample. As such, while the Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025) work includes only the noncoding regions of these newly sequenced individuals, we here present the full genome data including functional regions as well. The demographic history of this population involves a relatively ancient size reduction (likely associated with human colonization of Madagascar), followed by a period of recent decline up to the current day (likely related to recent habitat loss). Based on this demographic history, we simulated 100 replicates for each of the 14 autosomal chromosomes included in the most recent aye-aye genome assembly (Versoza and Pfeifer 2024). In the absence of fine-scale mutation and recombination rate maps for this species, we modeled a mutation rate of 1.52e-8 per base pair per generation (i.e. the average mutation rate previously reported in another lemur species; Campbell et al. 2021) and a broad-scale recombination rate of 1 cM/Mb (as recently inferred from pedigree data; Versoza, Lloret-Villas et al. 2025).

To generate genome-wide null thresholds, we ran both SweepFinder2 v.1.0. (DeGiorgio et al. 2016) and the B_{OMAF} method of Cheng and DeGiorgio (2020) on the allele frequency files generated from our simulated demographic data. In brief, we performed inference at each SNP with SweepFinder2 using the following command: SweepFinder2 -lu GridFile FreqFile SpectFile OutFile. Additionally, we utilized 2 inference schemes in B_{OMAF}: (1) windows containing 10 SNPs with a 5 SNP step size and (2) windows containing 100 SNPs with a 50 SNP step size, using the following command: python3 BalleRMix+_v1.py -I FreqFile -spect SfsFile -o OutFile -w W -s S usePhysPos -noSub -MAF -rec 1e-8, where W is the window size and S is the step size, both based on the number of SNPs. Because we lacked information on the polarization of SNPs, allele frequencies were folded, and only polymorphic sites were included in the analyses. Notably, the highest CLR value across all null model simulations was set as the null threshold for inference, under the assumption that this is the highest value that can be generated in the absence of positive or balancing selection, thereby providing a conservative scan to reduce false-positive rates.

Inferring positive and balancing selection in the aye-aye genome

We ran SweepFinder2 and B_{OMAF} on the 14 aye-aye autosomes using the same inference schema discussed above. Only those inference values greater than the null threshold values were considered as putatively experiencing positive or balancing selection. The genes in which these selected sites were located were identified using the genome annotations of Versoza and Pfeifer (2024), whilst overlaps with structural variants were also identified, based on those described by Versoza, Jensen et al. (2024). Because the number of identified genes was relatively small (<200 for both the sweep and balancing selection scans), we manually curated our candidates. In brief, for sweep candidates, we first identified genes under the significant likelihood surface. These candidate genes were then run through the NCBI database (Savers et al. 2022) and Expression Atlas (Madeira et al. 2022) in order to identify function and expression patterns in other primate species. Additionally, we performed a Gene Ontology (GO) analysis (The Gene Ontology Consortium 2023) using the Database for Annotation, Visualization, and Integrated Discovery (DAVID; Ma et al. 2023) on our candidate genes (conducted for all candidates together and also separately for selective sweep and balancing selection candidates). As several of our candidate genes were related to olfaction, and as primate olfactory genes are organized in large tandem repeats that may be challenging for the accurate mapping of short-read sequencing data, we computed the confidence score of the read mappings to measure the confidence that the observed variants were indeed assigned to the correct genomic location.

Power analyses

To assess how much statistical power exists to detect episodic selection in this aye-aye population given the details of both the demographic history and of the dataset itself, we simulated the Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025) demographic model forward-in-time in SLiM v.4.0.1 (Haller and Messer 2023). Thereby, the simulated region was comprised of 3 functional regions, separated by intergenic regions of length 16,489 bp. Each functional region contained 9 exons of length 130 bp, separated by introns of length 1,591 bp, for a total region length of 91,161 bp. These details were estimated from the Versoza and Pfeifer (2024) genome annotations to represent common aye-aye genomic architecture. Mutations in intronic and intergenic regions were modeled as effectively neutral, while exonic mutations were drawn from a distribution of fitness effects (DFEs) comprised of 4 fixed classes (Johri et al. 2020), whose frequencies are denoted by fi:f0 with $0 \le 2N_e s < 1$ (i.e. effectively neutral mutations), f1 with $1 \le 2N_e s < 10$ (i.e. weakly deleterious mutations), f2 with $10 \le 2N_e s < 100$ (i.e. moderately deleterious mutations), and f3 with $100 \le 2N_e s < 2N_e$ (i.e. strongly deleterious mutations), where N_e is the effective population size and s is the reduction in fitness of the mutant homozygote relative to wild type. Within each bin, s was drawn from a uniform distribution. We utilized the general DFE shape recently inferred in both humans and aye-ayes (Johri et al. 2023; Soni et al. 2025). This modeling of a realistic DFE in functional regions enabled us to account for the effects of purifying and background selection, in addition to population history, when assessing a baseline model of commonly acting evolutionary processes in the species.

To model uncertainty and heterogeneity in the underlying mutation and recombination rates, each 1-kb region of the simulated chromosome was assigned a different rate. Rates were drawn from a uniform distribution such that the chromosome-wide average was approximately the fixed rate previously observed in pedigree data (i.e. 1 cM/Mb; Versoza, Lloret-Villas et al. 2025). For variable recombination rates, the minimum and maximum parameters of the uniform distribution were 0.01 and 10 cM/Mb. respectively (i.e. a 100-fold decrease and a 10-fold increase on the fixed rate), mimicking both cold spots and hotspots. For variable mutation rates, the minimum and maximum parameters of the uniform distribution were set at 0.61e-8 and 3.8e-8 per base pair per generation (i.e. 0.5x and 2.5x the fixed rate previously reported in lemurs, respectively; Campbell et al. 2021). Notably, as much remains unknown about the fine-scale distribution of mutation and recombination rates (including the recombinational hotspot landscape) in this highly endangered species, the underlying heterogeneity is thus being represented by empirically informed variance estimates. However, as both higher quality recombination and mutation rate maps across the aye-aye genome become available, it would become natural to utilize those maps directly when performing such inference.

Simulations had a 10N generation burn-in time, where N is the ancestral population size of 23,706. A further 10N generations were then simulated for the sweep analysis, whilst a further 85N generations were simulated for the balancing selection analysis. In each simulation replicate, a single positively selected mutation was introduced. For the selective sweep analysis, the beneficial mutation was introduced at $\tau = [0.1, 0.2, 0.5, 1, 2]$, where τ is the time before sampling in N generations. Three different beneficial selection coefficients were simulated: $2N_e s = [100, 1,000, 10,000]$, where N_e is equal to the ancestral population size of 23,706. For the balancing selection analysis, the balanced mutation was introduced at $\tau = [10N, 50N, 75N]$. The balanced mutation experienced negative frequency-dependent selection, which was modeled such that the selection coefficient of the balanced mutation was dependent on its frequency in the population: $S_{bp} = F_{eq} - F_{bp}$, where S_{bp} is the selection coefficient of the balanced mutation, $\ensuremath{F_{\text{eq}}}$ is the equilibrium frequency of the balanced mutation (here set to 0.5), and $F_{\rm bp}$ is the frequency of the balanced mutation in the population. Simulations were structured such that if the selective sweep failed to fix, or the balanced mutation was either fixed or lost from the population, the simulation would restart at the point of introduction of the selected mutation.

Scans for selective sweeps and balancing selection were then performed on the simulated data, as per the procedure discussed above. Receiver operating characteristic (ROC) plots were generated in order to summarize expected performance. Because selective sweep inference was performed on each SNP, 100 bp, 1 kb and 10 kb windows were generated for creating ROC plots for SweepFinder2 results (note that this was not necessary for B_{OMAF} as SNP-based windows were used for inference).

Results and discussion

We sequenced the genomes of 5 unrelated aye-aye (*D. madagascar*iensis) individuals (4 wild-born and 1 colony-born of wild-born parents) housed at the Duke Lemur Center to an average coverage of >50×. After mapping reads to the recently published chromosome-level genome assembly for the species, we called variant and invariant sites following the best practices for nonmodel organisms (Pfeifer 2017a; van der Auwera and O'Connor 2020). We ran genomic scans across this newly generated populationlevel dataset, using the 14 autosomal scaffolds from the aye-aye genome assembly of Versoza and Pfeifer (2024) (excluding the



Fig. 1. Genome scans for selective sweeps using SweepFinder2. Blue data points are CLR values inferred at each SNP. The dashed line is the threshold for sweep detection, determined by the highest CLR value across 100 simulated replicates of each of the 14 autosomal scaffolds (see Materials and Methods section for further details). The x-axis represents the position along the scaffold, and the y-axis represents the CLR value at each SNP.

X-chromosome, i.e. scaffold 9; see Terbot, Soni, Versoza, Mihaven et al. 2025). The CLR methods implemented in SweepFinder2 and the B_{0MAF} statistic of the BalLeRMix software package were used to infer selective sweeps and balancing selection, respectively. Selective sweep inference was performed at each SNP, whilst balancing selection inference was performed in windows of size 10 and 100 SNPs. Figures 1 and 2 provide the results of the genomewide scans for SweepFinder2 and B_{0MAF} based on 100 SNP windows, respectively (and see Supplementary Fig. S1 for the genome-wide scan results with B_{0MAF} on 10 SNP windows), highlighting a number of peaks along the likelihood surface for each analysis.

Although it is common to use outlier approaches to identify candidate regions experiencing positive selection, it has previously been shown that such approaches are often associated with extreme false-positive rates (Teshima et al. 2006; Thornton and Jensen 2007; Jensen et al. 2008; Jensen 2023; Soni et al. 2023). Furthermore, these approaches are problematic as any evolutionary model (including standard neutrality) will naturally have a 1 and 5% tail, and thus assuming that genes in these tails of an observed empirical distribution are likely sweep candidates is inherently flawed (see Harris et al. 2018). Moreover, it is not a given that recent sweeps, if they exist, will necessarily even appear in the tails of the empirical distribution under any given demographic model. Thus, we instead followed the recent recommendations of Johri, Eyre-Walker et al. (2022) in carefully constructing an evolutionarily appropriate baseline model accounting for commonly acting evolutionary processes (as summarized in the diagrams shown in Figures 5 and 6 of Johri, Aquadro et al. 2022). Furthermore, we characterized the expected true- and falsepositive rates of the utilized statistical approaches for our given

genomic dataset, as positive and balancing selection will not necessarily be detectable within the context of any given baseline model (Barton 1998; Thornton and Jensen 2007; Poh *et al.* 2014; Harris and Jensen 2020); importantly, even if these events are not detectable, this approach remains necessary for managing false-positive rates.

We therefore simulated 100 replicates for each of the 14 autosomes using msprime (Baumdicker et al. 2022), under the aye-aye demographic model inferred by Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025), utilizing the genome assembly of Versoza and Pfeifer (2024), and modeling the specific data details of our newly presented whole-genome dataset. We performed SweepFinder2 and B_{OMAF} analyses on these baseline simulations, using the maximum CLR values across all simulation replicates as the conservative null thresholds for positive and balancing selection inference, reasoning that these are the maximum values that can be generated in the absence of episodic selective processes under the baseline model considered, and any CLR values that exceeded the thresholds in our empirical analyses were considered to represent meaningful candidate regions. The identified threshold values under this model were 211.747 for SweepFinder2 inference at each SNP, 50.817 for BOMAF inference on 10 SNP windows, and 244.382 for BOMAF inference on 100 SNP windows.

Signatures of episodic selection in the aye-aye genome

A total of 3,462 loci met our null threshold for selective sweep inference using SweepFinder2, which mapped to 71 genes within the aye-aye genome. Scaffolds 1, 7, 10, and 11 contained at least one candidate region, although numerous regions on other scaffolds



Fig. 2. Genome scans for balancing selection using the B_{OMAF} method. Blue data points are CLR values inferred over windows of length 100 SNPs. The dashed line is the threshold for detection, determined by the highest CLR value across 100 simulated replicates of each of the 14 autosomal scaffolds (see *Materials and Methods* section for further details). The x-axis represents the position along the scaffold, and the y-axis represents the CLR value at each window.

show peaks that were below our null threshold. For balancing selection inference with B_{OMAF} , no windows met our null threshold for windows of size 10 SNPs, and 163 windows met our null threshold for windows of size 100 SNPs. The latter windows mapped to 60 candidate genes, covering all autosomal scaffolds apart from scaffolds 11, 14, and 15. Supplementary File S1 provides tables of candidate regions overlapping genes exhibiting CLR values greater than the null thresholds for these analyses.

We manually curated the 71 sweep candidate genes, particularly noting those associated with the peak of each significant likelihood surface (e.g. see Fig. 3 for scaffolds 1 and 7 for zoomed inset plots and Supplementary Figs. S2 and S3 for additional scaffolds containing candidate regions). Because balancing selection candidate genes were based on 100 SNP windows, these peaks were already quite localized by comparison, though we similarly manually curated this associated set of 60 candidate genes (see Fig. 4 for the results in scaffold 1 with mapped genes marked on the plot and Supplementary Figs. S4-S12 for additional scaffolds containing candidate regions). The gene that exhibited the strongest signal of positive selection (i.e. the highest CLR value) was SMPD4 on scaffold 10, whilst the gene exhibiting the strongest signal of balancing selection was LRP1B on scaffold 6 for the 100 SNP window analysis. Biallelic loss-of-function variants in SMPD4 have been found to cause microcephaly, a rare and severe neurodevelopmental disorder with progressive congenital microcephaly and early death in humans (Magini et al. 2019; Smits et al. 2023). LRP1B is a putative tumor suppressor (Brown et al. 2021), and one of the most altered genes in human cancer (Principe et al. 2021). Mutations in LRP1B have been associated with an increased tumor mutation burden (Yu et al. 2022). In one of the earliest large-scale

primate genome scans, Nielsen *et al.* (2005) found that a number of genes involved in tumor suppression were identified as positive selection candidates in humans, as well as genes involved in spermatogenesis, which our analysis in aye-ayes also identified (see below).

A single candidate region (scaffold 1:315,975,047–315,981,365) in the 100 SNP window balancing selection analysis was found to overlap with a 89,250-bp inversion (scaffold 1:315,972,759– 316,062,009). This region may represent a false positive, owing to the reduced recombination related to the inversion potentially generating long haplotype structure (Stevison *et al.* 2011). However, inversions may themselves be selectively maintained, particularly if they contain a beneficial combination of alleles (e.g. Hager *et al.* 2022; and see Villoutreix *et al.* 2021); thus, this candidate region will require future dissection.

Gene functional analysis

A gene function analysis using the Database for Annotation, Visualization, and Integrated Discovery (DAVID; Ma *et al.* 2023) predicted an enrichment for 19 GO terms in aye-ayes at a $P \leq 0.05$ for recent selective sweep candidate genes—however, no terms passed a false discovery rate (FDR) threshold of 0.05 or 0.1 for selective sweep candidate genes. Conversely, 14 of the 43 GO terms identified for the 100 SNP window analysis passed both the $P \leq 0.05$ and the FDR ≤ 0.05 thresholds for balancing selection. Table 1 provides the top 12 functional categories for balancing selection candidate genes for the 100 SNP analysis (and see Supplementary File S1 for all enriched categories). These categories show considerable enrichment, and the majority are involved in olfaction.



Fig. 3. SweepFinder2 selective sweep scan results for a) scaffold 1 and b) scaffold 7. Inset plots zoom in on likelihood surface peaks, with genes in these regions highlighted. The x-axis represents the position along the scaffold, and the y-axis represents the CLR value at each SNP.



Fig. 4. B_{OMAF} balancing selection scan results for 100 SNP window analysis on a) scaffold 1 and b) scaffold 7. Red vertical lines map to candidate genes. Instances where CLR values meet the null threshold, but no gene is denoted, indicate that no gene overlap was found. The x-axis represents the position along the scaffold, and the y-axis represents the CLR value of each 100 SNP window.

Table 1. Top 12 hits from gene functional analysis with DAVID on balancing selection candidate genes from 100 SNP window analysis.

Category	Term	P-value	Fold enrichment	FDR
KEGG_PATHWAY	hsa04740:olfactory transduction	3.22E-06	7.23	2.96E-04
GOTERM_BP_DIRECT	GO:0050911~detection of chemical stimulus involved in sensory perception of smell	3.72E-06	9.32	8.16E-04
GOTERM_MF_DIRECT	GO:0004984~olfactory receptor activity	5.06E-06	8.92	4.71E-04
INTERPRO	IPR000725:Olfact_rcpt	5.07E-06	8.98	8.17E-04
UP_KW_BIOLOGICAL_PROCESS	KW-0552~Olfaction	6.24E-06	8.16	1.44E-04
GOTERM_MF_DIRECT	GO:0004930~G protein-coupled receptor activity	2.00E-05	6.22	9.32E-04
INTERPRO	IPR000276:GPCR_Rhodpsn	2.63E-05	6.05	1.70E-03
INTERPRO	IPR017452:GPCR_Rhodpsn_7TM	3.17E-05	5.91	1.70E-03
UP_SEQ_FEATURE	DOMAIN:G protein-coupled receptors family 1 profile	6.06E-05	6.36	2.12E-02
UP_KW_BIOLOGICAL_PROCESS	KW-0716~sensory transduction	7.17E-05	5.82	8.25E-04
GOTERM_BP_DIRECT	GO:0007186~G protein-coupled receptor signaling pathway	1.62E-04	4.76	1.78E-02
GOTERM_MF_DIRECT	GO:0005549~odorant binding	2.20E-04	16.38	6.83E-03

Olfaction

Seven enriched gene functions for balancing selection (enrichment score >1) were related to olfaction, all with high-fold enrichment (ranging from 5.82 to 16.38). A number of olfactory receptor (OR) genes were also found to meet our null threshold for selective sweeps. OR genes provide the basis for the sense of smell (e.g. Buck and Axel 1991) and comprise the largest gene superfamily in mammalian genomes (Glusman et al. 2001; Zozulya et al. 2001). Although olfaction plays a role in locating food, mating, and avoiding danger, the importance and sensitivity of smell varies significantly even amongst closely related species, and it has been suggested that this gene family is subject to a birth-and-death model of evolution, whereby new genes are formed by gene duplication and some of the duplicate genes differentiate in function, whilst others become inactive or are removed from the genome (Niimura and Nei 2003, 2005a, 2005b, 2007). Indeed, OR genes within primates show relaxed selective constraints in apes relative to Old and New World monkeys (Dong et al. 2009), and whilst patterns of variability in chimpanzees are consistent with purifying selection acting on intact OR genes, patterns in humans have been suggested to indicate the action of positive selection (Gilad, Bustamante et al. 2003). These findings were partially corroborated by Williamson et al. (2007), who found evidence of recent selective sweeps in numerous OR genes in human populations. Gilad, Bustamante et al. (2003) argued that the differing selective processes acting on OR genes in humans and chimpanzees are likely reflections of differences in lifestyles between humans and other great apes, resulting in distinct sensory needs. Further studies have suggested that both positive selection (Gilad, Man et al. 2003) and balancing selection (Alonso et al. 2008) are acting on OR genes in humans. Importantly, it has been proposed that heterozygosity in ORs can increase the number of different odorant-binding sites in the genome (Lancet 1994), and thus heterozygote advantage may be an important process. Given that aye-ayes have been shown to discriminate based on scent (Price and Feistner 1994) and use scent-marking behaviors to attract mates (Winn 1994), OR genes represent interesting candidate loci for having experienced ongoing positive and balancing selection.

The organization of primate OR genes in tandem arrays poses challenges for both the accurate assembly of gene organization and copy number as well as the accurate mapping of short-read sequencing data to these regions. However, several lines of evidence suggest that our candidate loci are likely genuine rather than artifacts. First, the aye-aye genome assembly was built from high-coverage, ultra-long Oxford Nanopore reads with average length of 38.7 kb (Versoza and Pfeifer 2024)—a read length well above that required to fully resolve the majority of OR genes in primate genomes (e.g. the median length of OR genes in the "gold standard" telomere-to-telomere human assembly is 1.9 kb; Nurk et al. 2022), in particular those meeting our null thresholds (with 10 OR genes being <1.7 kb and 1 OR gene being ~10 kb). Second, using Sanger sequencing for validation, a recent study in humans (Trimmer et al. 2019) demonstrated that short-read Illumina data sequenced at a minimum of 15-fold coverage can accurately identify genetic variation in OR genes when sequencing reads map with a high confidence (>99.9%). Ten out of the 11 OR genes meeting our null thresholds for selective sweeps and balancing selection surpass this benchmarked highconfidence threshold in our high-coverage (>50-fold) data (Supplementary Table S3). The single candidate OR gene below this high-confidence threshold (OR4A16) exhibits considerable sequence similarity with other OR genes in the aye-aye genome, hindering reliable mapping with short-read data. Future long-read sequencing may provide an avenue to obtain additional insights into this candidate (though it should be noted that long-read sequencing remains challenging in highly endangered species such as aye-ayes due to the inherently low sample availability).

Rhodopsin

Six enriched gene functions for balancing selection (enrichment score >1) were related to G protein-coupled receptors (GPCRs), with fold enrichment scores ranging from 3.75 to 6.22. GPCRs are cell surface receptors responsible for detecting extracellular molecules and activating cellular responses; consequently, they are involved in numerous physiological processes. Two of the 6 enriched GPCR functions were specifically related to rhodopsin, which is the opsin responsible for mediating dim light vision (Litman and Mitchell 1996). It has previously been shown that, despite being nocturnal, aye-ayes maintain dichromacy-potentially supporting previous work that found that the red/green opsin gene survived the long nocturnal phase of mammalian evolution, which has been hypothesized to relate to a role in setting biorhythms (Nei et al. 1997). These identified candidate regions may also lend credence to the speculation that dichromatic nocturnal primates may be able to perceive color while foraging under moonlight conditions (Perry et al. 2007).

PATE gene family

The PATE gene family has been shown to express in the testis, encoding a sperm-related protein (Bera *et al.* 2002). Of the 4 genes that make up the PATE gene family, 3 were found in candidate selective sweep regions in our analysis (Fig. 3), whilst PATE1 was also found in our balancing selection scans based on 100 SNP windows (Fig. 4). Soler-García *et al.* (2005) found that PATE is highly expressed in the male genital tract and that proteins are secreted into the semen, suggesting a potential in mammalian sperm maturation, whilst Margalit *et al.* (2012) found that PATE proteins are involved in sperm-oolemma fusion and penetration.

Although primate sperm displays a general uniformity, previous studies have found variations in sperm morphology (Cummins and Woodall 1985; Gage 1998), often predicated on the absence or presence of sperm competition. Indeed, the use of coagulated ejaculate that forms sperm plugs to avoid sperm competition and increase male fertilization success has been described in multiple species of primates, particularly those exhibiting polygynandrous mating systems (i.e. multimale, multifemale mating systems; Dixson *et al.* 2005; Martinez and Garcia 2020). As aye-ayes are polygynandrous (Quinn and Wilson 2004), these candidate loci may similarly be hypothesized to relate to mate competition/sexual selection shaping the PATE family of genes.

Zinc-finger genes

Multiple zinc-finger (ZNF) genes were identified as having undergone recent selective sweeps or balancing selection in our scans of the aye-aye genome. ZNF genes are the largest family of transcription factors in mammalian genomes and play an important role in gene regulation. Previous studies have identified a number of KRAB-ZNF genes-a subfamily of the deeply conserved Kruppel-type zinc-finger (KZNF) genes (Bellefroid et al. 1993; Looman et al. 2002; Huntley et al. 2006)—with evidence of positive selection in humans (Nielsen et al. 2005; Nowick et al. 2010, 2011; Jovanovic et al. 2021); additional research has suggested that KRAB-ZNF genes follow a species-specific—as opposed to tissuespecific—pattern of expression (Kapopoulou et al. 2016), suggesting that these genes have different tissue preferences in different species and thus have functionally diversified across the primate lineage (Liu et al. 2014). In addition, numerous gene regulatory factors were identified as putative candidate regions, likely related to the well-described roles of these genes in modifying expression patterns (Berrio et al. 2020; Liu and Robinson-Rechavi 2020; Jovanovic et al. 2021; and see the review of McDonald and Reed 2023).

Quantifying power to detect positive and balancing selection in aye-ayes

It has previously been demonstrated that the statistical power to detect positive selection in any given population will depend on a variety of factors, ranging from the details of the population history to the amount and configuration of the analyzed data itself (e.g. Johri et al. 2021; Johri, Eyre-Walker et al. 2022; Soni et al. 2023, 2025; Soni and Jensen 2025). In order to quantify the specific power of this analysis to detect candidate regions in aye-ayes, we ran forward-in-time simulations in SLiM (Haller and Messer 2023), utilizing the well-fitting demographic history presented in Terbot, Soni, Versoza, Pfeifer et al. (2025) and Terbot, Soni, Versoza, Milhaven et al. (2025). Recent work has inferred the neutral and deleterious DFE in aye-ayes (Soni et al. 2025), finding a similar distribution to that previously observed to characterize human genes (Johri et al. 2023), which was here utilized to model and thus account for purifying and background selection effects. In order to model the impact of mutation and recombination rate heterogeneity (Johri, Aquadro et al. 2022; Soni et al. 2024), we drew rates from a uniform distribution for each 1-kb window, such that the mean across each simulation replicate was equal to the mean genomic rate (see the Materials and Methods section for more details).

In each simulation replicate, a single beneficial mutation was introduced. For selective sweep models, 3 different selection regimes were considered, with population-scaled strengths of selection, $2N_es$, of 100, 1,000, and 10,000, where N_e is the ancestral population size and s is the strength of selection acting on the beneficial mutation. Five different introduction times of the beneficial mutation were considered, $\tau = 0.1, 0.2, 0.5, 1, and 2$, where τ is the time before sampling in N generations. These values were chosen as there is not expected to be power to detect selective sweeps beyond 4N generations, though power generally decays much more rapidly (Kim and Stephan 2002; Przeworski 2002, 2003; Ormond et al. 2016). Only simulation replicates in which the beneficial mutation fixed were retained. For balancing selection, the beneficial mutation was modeled as experiencing negative frequency-dependent selection, and introduced at $\tau = 10N$, 50N, and 75N generations, as it has been shown that SFS-based methods have little power to detect recent balancing selection (Soni and Jensen 2024). Only simulation replicates in which the balanced mutation was still segregating at the time of sampling were retained.

Figures 5 and 6 provide ROC plots for selective sweep inference with SweepFinder2 and B_{OMAF}, respectively. As shown, the power to detect positive selection in aye-ayes is expected to be reasonably poor in all cases, though results at 100-bp windows are most encouraging in suggesting sufficient power. This general result is likely consistent with the observation that aye-ayes have undergone a recent bottleneck, an event which may replicate patterns of variation consistent with a selective sweep (e.g. Barton 1998; Poh et al. 2014; Harris and Jensen 2020), followed by a period of further decline; relatedly, balancing selection inference is expected to be partially confounded by the resulting skew in the SFS toward intermediate frequency alleles (Soni and Jensen 2024). Additionally, identifying the episodic and locus-specific patterns of positive selection is more challenging in populations that have experienced bottlenecks owing to the large genealogical variance generated by this event, leading to widely dispersed test statistics across the genome which can give the illusion of a locusspecific pattern (Thornton and Jensen 2007). While power to detect selective sweeps generally related with strength as expected, this relationship was partially off-set by the fact that at higher strengths of selection the beneficial mutation fixed more rapidly in the population, and thus the time since fixation was longer thereby reducing inference power, as has been previously described analytically (Kim and Stephan 2000). By contrast, power to detect balancing selection increased with time since the introduction of the balanced mutation, as has been previously shown (e.g. Soni and Jensen 2024). In summary, these power analyses provide a key for interpreting our empirical analysis, in demonstrating that any statistically detectable selective sweep would need be both strong and recent, while any statistically detectable loci experiencing balancing selection would need to be relatively ancient.

Concluding thoughts

We ran the first large-scale scans for loci having experienced selective sweeps and balancing selection in the aye-aye genome, using newly generated, high-quality, whole-genome, populationlevel data, and utilizing the recent fully annotated genome assembly of Versoza and Pfeifer (2024) together with the well-fitting demographic model inferred from nonfunctional genomic regions of Terbot, Soni, Versoza, Pfeifer *et al.* (2025) and Terbot, Soni,



Fig. 5. ROC plots for SweepFinder2 showing the change in true-positive rate (TPR) as the false-positive rate (FPR) increases, for sweep inference in aye-ayes across 100 simulation replicates under the Terbot, Soni, Versoza, Pfeifer *et al.* (2025) and Terbot, Soni, Versoza, Milhaven *et al.* (2025) demographic model, with mutation and recombination rates drawn from a uniform distribution such that the mean rate per simulation is equal to the fixed rate (see Materials and Methods section). Power analysis was conducted across 3 selection regimes (population-scaled strengths of selection of $2N_es = 100, 1,000, and 10,000)$, 5 different times of introduction of the beneficial mutation (r = 0.1, 0.2, 0.5, 1, and 2, in N generations), and 3 window sizes (100 bp, 1 kb, and 10 kb). If no ROC is plotted, this is a case in which the beneficial mutation was unable to fix prior to the sampling time in any of the simulation replicates (e.g. at $2N_es = 100$ and r = 0.1).

Versoza, Milhaven et al. (2025). By simulating an evolutionarily appropriate baseline model employing these details (Johri, Aquadro et al. 2022; Johri, Eyre-Walker et al. 2022), we were able to generate conservative null thresholds for these scans, thereby greatly reducing the types of false-positive rates associated with outlier approaches (Teshima et al. 2006; Thornton and Jensen 2007; Jensen et al. 2008; Soni et al. 2023). The differences between our conservative baseline model approach and a traditional genomic outlier approach are considerable (and see the discussions in Howell et al. 2023; Jensen 2023; Johri et al. 2023; Terbot et al. 2023). For example, selection inference with B_{OMAF} on 10 SNP windows yielded no candidate windows, and inference on 100 SNP windows yielded 163 candidate windows. An outlier approach interpreting the (commonly used) 5% tail of the empirical distribution of CLR values as candidate loci would instead give 22,971 candidate windows at 10 SNP windows, and 2,280 for 100 SNP windows. More to the point however, as demonstrated in our power analyses, the great majority of density in these tail distributions may be readily generated by the population history alone, and thus need not invoke positive or balancing selection as an explanation.

Through this more thorough and conservative approach, we have identified a number of promising candidate genes with evidence of having been episodically impacted by positive and balancing selection during the recent evolutionary history of the species, with particularly notable examples being those involved in olfaction and spermatogenesis. Given the ongoing destruction of aye-aye habitats, it may well be hypothesized that these functions are experiencing changing selection pressures, particularly in light of their solitary lifestyle and polygynandrous mating system.

Data availability

Sequence data are available under NCBI BioProjects PRJNA1085541 and PRJNA1179987. All scripts to generate and analyze simulated data, as well as results from selection scans, are available at the



Fig. 6. ROC plots for B_{0MAF} showing the change in true-positive rate (TPR) as the false-positive rate (FPR) increases, for balancing selection inference in aye-ayes across 100 simulation replicates under the Terbot, Soni, Versoza, Pfeifer *et al.* (2025) and Terbot, Soni, Versoza, Milhaven *et al.* (2025) demographic model, with mutation and recombination rates drawn from a uniform distribution such that the mean rate per simulation rate is equal to the fixed rate (see *Materials and Methods* section). Power analyses were conducted across 3 different times of introduction of the balanced mutation (r = 10N, 50N, and 75N generations).

GitHub repository: https://github.com/vivaksoni/aye_aye_recent_positive_selection.

Supplemental material available at G3 online.

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Conflicts of interest

The author(s) declare no conflicts of interest.

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