A Pedigree-Based Map of Crossovers and Noncrossovers in Aye-Ayes (*Daubentonia madagascariensis*)

Cyril J. Versoza^{1,†}, Audald Lloret-Villas^{1,†}, Jeffrey D. Jensen 1, Susanne P. Pfeifer 1,*

¹Center for Evolution and Medicine, School of Life Sciences, Arizona State University, Tempe, AZ, USA

*Corresponding author: E-mail: susanne@spfeiferlab.org.

[†]These authors contributed equally to this work.

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Abstract

Gaining a better understanding of the rates and patterns of meiotic recombination is crucial for improving evolutionary genomic modeling, with applications ranging from demographic to selective inference. Although previous research has provided important insights into the landscape of crossovers in humans and other haplorrhines, our understanding of both the considerably more common outcome of recombination (i.e. noncrossovers) as well as the landscapes in more distantly related primates (i.e. strepsirrhines) remains limited owing to difficulties associated with both the identification of noncrossover tracts as well as species sampling. Thus, in order to elucidate recombination patterns in this understudied branch of the primate clade, we here characterize crossover and noncrossover landscapes in aye-ayes utilizing whole-genome sequencing data from six three-generation pedigrees and three two-generation multi-sibling families, and in so doing provide novel insights into this important evolutionary process shaping genomic diversity in one of the world's most critically endangered primate species.

Key words: recombination, primate, strepsirrhine, Daubentoniidae, population genomics.

Significance

The rates of recombination—a fundamental process in mammalian meiosis—differ markedly between species. In this study, we provide the first recombination rate estimates for a strepsirrhine, the aye-aye (*Daubentonia madagascariensis*)—a highly endangered species and valuable comparative outgroup for future primate studies.

Introduction

Recombination is a fundamental biological process required for faithful gametogenesis in most sexually reproducing species (see the reviews of Baudat et al. (2013) and Johnston (2024)). Apart from being essential for the proper pairing of homologous chromosomes and their segregation into gametes during meiosis (Keeney 2001), recombination also plays an important evolutionary role in shuffling genetic variation, improving the efficacy of natural selection by breaking linkage interference between segregating beneficial and deleterious alleles (Hill and Robertson 1966; Felsenstein 1974; and see the review of Charlesworth and Jensen (2021)).

In primates, as in many other organisms, meiotic recombination predominantly occurs in 1 to 2 kb long regions of the genome—so-called recombination "hotspots"—the location of which is mainly determined by the zinc-finger protein PRDM9 (Baudat et al. 2010; Myers et al. 2010; Parvanov et al. 2010). By binding specific DNA sequence motifs and trimethylating histone H3 at lysines 4 and 36 (Powers et al. 2016), PRDM9 guides the meiotic machinery

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to initiate the formation of DNA double-strand breaks, the repair of which may result in either a reciprocal exchange between homologs (termed a crossover; CO) or a unidirectional replacement of a genomic region in one chromosome leaving the donor homolog unmodified (termed a noncrossover; NCO) (see the reviews of Wang et al. (2015), Lorenz and Mpaulo (2022), and Johnston (2024)).

While both COs and NCOs play important roles in shaping genetic diversity (Przeworski and Wall 2001), previous studies have suggested that NCOs tend to be considerably more common than COs in most organisms (Jeffreys and May 2004; Cole et al. 2010; Comeron et al. 2012; Li et al. 2019; Palsson et al. 2025). However, owing to the difficulty in detecting the often very small NCO events (with most tract lengths in humans being <1 kb; Jeffreys and May 2004; Williams et al. 2015; Halldorsson et al. 2016; Palsson et al. 2025), combined with the need for a segregating variant to be present in the donor homolog in order to allow identification (i.e. the tract remains undetectable if the donor and converted sequence are identical), the NCO landscape remains comparatively understudied. For these same reasons, combined with other underlying assumptions such as geometrically distributed tract lengths, computational approaches for NCO inference are generally characterized by poor accuracy (see discussion of Wall et al. (2022)). As such, time-consuming and costly direct pedigree-sequencing studies remain the most promising avenue for improving our understanding of this process (and see Peñalba and Wolf (2020) for a detailed overview of current methodologies).

With regards to primates specifically, although previous research has begun to elucidate the rates and patterns of recombination in haplorrhines—a suborder of primates that includes humans (Kong et al. 2002, 2010; Coop et al. 2008; Pratto et al. 2014; Williams et al. 2015; Halldorsson et al. 2016; Palsson et al. 2025), other great apes (Auton et al. 2012; Pfeifer and Jensen 2016; Stevison et al. 2016), and monkeys (Rogers et al. 2000, 2006; Cox et al. 2006; Jasinska et al. 2007; Xue et al. 2016, 2020; Pfeifer 2020; Wall et al. 2022; Versoza et al. 2024b)-little remains known about this process in strepsirrhines (though see Soni et al. 2024). As the most basal suborder of primates, gaining insights into the population genetic forces shaping the genomes of strepsirrhines is crucially important, not only to improve our understanding of primate evolution, but also for elucidating the scale at which recombination rates and spatial distributions of CO and NCO events may change between taxonomic groups, as this variation has been shown to be substantial (see the reviews of Paigen and Petkov (2010) and Stapley et al. (2017)). Given the vital role of recombination in maintaining genetic diversity, such inference is additionally important for the development of effective conservation strategies, particularly as many strepsirrhines are highly endangered (Gross 2017). For example, among the more than 100 species of strepsirrhines endemic to Madagascar (Mittermeier et al. 2010), aye-ayes (*Daubentonia madagascariensis*) are one of the most threatened by anthropogenic activities, such as slash-and-burn agriculture, logging, mining, and urbanization (Suzzi-Simmons 2023), with the resulting extensive habitat

2020). Thus, to elucidate the rates and patterns of recombination in this understudied branch of the primate tree—which represents an early split in the primate clade and thus a valuable comparative outgroup for future primate studies—we here investigate both the more commonly studied CO and the less-commonly studied NCO landscapes in aye-ayes. Using whole-genome sequencing data from six threegeneration pedigrees and three (partially overlapping) two-generation multi-sibling families, we present the first recombination rate estimates for a strepsirrhine and thereby provide novel insights into this important evolutionary process shaping genetic diversity in one of the world's most critically endangered primate species.

loss and fragmentation having already decimated their popu-

lations to an estimated 1,000 to 10,000 individuals (Louis et al.

Results and Discussion

To study the rates and patterns of recombination in aye-ayes, 14 individuals were selected from a multi-generation pedigree housed at the Duke Lemur Center, the genomes of which were sequenced to mean depths of 50x (supplementary table S1, Supplementary Material online; Versoza et al. 2025). After mapping reads to the speciesspecific genome assembly (Versoza and Pfeifer 2024), autosomal variants were called following the Genome Analysis Toolkit's Best Practices (van der Auwera and O'Connor 2020), and filtered using a set of coverage-, genotype-, and inheritance-based criteria, resulting in a high-confidence call set consisting of 1.8 million variants (supplementary table S2, Supplementary Material online). For the purpose of this study, this call set from a multi-generation pedigree was divided into partially overlapping six three-generation pedigrees and three two-generation nuclear families with multiple offspring (supplementary fig. S1, Supplementary Material online) for which gamete transmission could be tracked in order to identify recombination events based on "phase-informative" markers-that is, heterozygous variants for which the parent-of-origin could be determined (with an average of 0.5 million phase-informative markers per pedigree/family; supplementary table S3, Supplementary Material online, and see supplementary table S4 and figs. S2 to S15, Supplementary Material online for the marker density across each autosome per pedigree/family). Based on these phase-informative markers, recombination events were classified as either COs-that is, a single change of haplotype phase along a chromosome (from maternally inherited to paternally inherited haplotype blocks or vice versa)—or NCOs—that is, phase-informative markers that mismatched surrounding haplotype blocks (for additional details, see "Materials and Methods").

The Landscape of COs in Aye-Ayes

A total of 119 and 308 putative CO events were identified across the autosomes in 6 and 18 meioses through the pedigree-based approach and the family-based approach, respectively. Visual inspection of these initial datasets revealed a clustering (i.e. \geq 2 COs originating from the same meiosis located within <1 Mb) of 40 COs across 8 genomic regions, with the majority of these events (90%) identified through only one of the two approaches (supplementary table S5, Supplementary Material online). Although COs form at random during meiosis, CO interference tightly regulates their location within each chromosome (Muller 1916; Broman and Weber 2000), making such a pattern highly implausible in nature (see the discussions of Wall et al. (2022) and Versoza et al. (2024b)). More likely, these observations are driven by localized genotyping errors; thus, these CO events were excluded from further analyses, as is common practice. Subsequently, pedigree-based and family-based datasets were manually consolidated across the 20 meioses by removing duplicate recombination events observed in the four meioses that could be studied using both approaches (i.e. the meioses for the focal individual-offspring pairs 7-12, 7-13, 8-12, and 8-13; see supplementary fig. S1, Supplementary Material online), yielding a set of 324 distinct CO events. Thereby, the majority of consolidated events (54%) shared identical start and end coordinates at the break point; in cases of a partial overlap (i.e. a sharing of either start or end coordinates at the break point owing to differences in phase-informative markers in the two approaches), the CO events with the shortest resolution (i.e. the smallest region in which the CO could be located) were selected. Notably, CO events are supported by ~8,000 phase-informative markers on average and all CO events in the meioses that could be studied using pedigree-based and family-based approaches were also identified by both approaches, thus serving as a valuable validation of the employed methodology. After the datasets were consolidated, an additional 19 COs were removed either because the event was detected across multiple meioses (18 COs), possibly indicating a misplacement or inversion of a contig during the genome assembly, or because it occurred at the same position as an NCO in another individual (1 CO), thus likely resulting from a genotyping error (supplementary table S6, Supplementary Material online). The final dataset contained 305 COs: 163 and 142 COs in the 10 maternal and 10 paternal meioses, respectively (Fig. 1a; and see supplementary table S7, Supplementary Material online for the location of the observed events).

Aye-ayes exhibit one of the lowest levels of nucleotide diversity of any primate studied to date (Perry et al. 2013; Soni

et al. 2025; Terbot et al. 2025b)-yet, despite this low marker density across the pedigree, the median resolution of male and female CO events (12.9 and 20.0 kb; supplementary fig. S16, Supplementary Material online, and see supplementary table S8, Supplementary Material online for a summary) is on par with those previously obtained from similarly sized nonhuman primate pedigree studies (7.7 and 22.3 kb in olive baboons [Wall et al. 2022] and rhesus macaques [Versoza et al. 2024b], respectively). In accordance with CO assurance, ensuring an obligate CO between homologous chromosomes (or chromosome arms) during meiosis (Jones and Franklin 2006), an average of 1.1 COs were identified per chromosome and meiosis. In total, between 10 and 21 COs were observed per meiosis across the 14 autosomes (supplementary table S8, Supplementary Material online), with CO densities in males and females roughly inversely correlated with chromosome size (supplementary fig. S17, Supplementary Material online).

In order to gain a better understanding of the nature of hotspots, information from CO events with a resolution of less than 10 kb was leveraged to search for putative sequence motifs of PRDM9 binding sites in aye-ayes. Application of a de novo motif finding algorithm (Bailey and Elkan 1994) allowed for the identification of a degenerate sequence motif present in these CO regions (Fig. 1b)—a motif that exhibits considerable similarities with PRDM9 binding motifs previously identified in humans (see Fig. 1a in Altemose et al. (2017) for zinc-finger domains 7 to 12), despite a high turn-over rate of PRDM9 binding motifs between species. CO events were significantly enriched in regions harboring this degenerate sequence motif compared with the genomic background (Fisher's exact test, *P*-value < 2.2e-16). Interestingly, AlphaFold 3 (Abramson et al. 2024) predicted an interaction between PRDM9 and the putative DNA binding motif in aye-ayes with high confidence (ipTM = 0.91, with a score close to 1.0 suggesting that the protein–DNA interface is welldefined), confirming PRDM9–DNA binding in silico. Visualization of the PRDM9–DNA complex further supports this prediction (Fig. 1c), showing well-folded C2H2 zinc-finger domains (with a very high-confidence score of pLDDT > 90) binding DNA at the discovered motif.

Consistent with a preferential binding of PRDM9 in intronic and intergenic regions that are often more accessible in chromatin structure (Coop et al. 2008; Walker et al. 2015), CO events were enriched in these genomic regions at the broad scale (supplementary fig. S18, Supplementary Material online). At the fine scale, the number of COs was considerably lower near, and higher farther away from, genes (supplementary fig. S19, Supplementary Material online)—an observation in agreement with previous studies showing that PRDM9 directs COs away from transcription start sites in primates (Myers et al. 2005; Coop et al. 2008; Auton et al. 2012). Additionally, a clustering of COs was observed toward the telomeric ends in males, while females



Fig. 1. Characteristics of crossovers. a) The genomic distribution of CO events in females (shown as red circles) and males (blue circles) across the autosomes (note that scaffold 9, i.e. chromosome X, is not displayed). b) Motif of the predicted PRDM9 binding sequence in aye-ayes. c) PRDM9—DNA complex predicted by AlphaFold 3 (Abramson et al. 2024). Colors indicate confidence (dark blue: very high, predicted local distance difference test [pLDDT] > 90; turquoise: confident, 90 > pLDDT > 70; yellow: low, 70 > pLDDT > 50; orange: very low, pLDDT < 50). Predictions are provided for noncommercial use only, under and subject to AlphaFold Server Output Terms of Use found at alphafoldserver.com/output-terms. d) Relationship between female and male CO rates and relative proximity to telomeric regions. e) Relationship between maternal and paternal age at birth and the number of CO events.

displayed an overall larger number of COs that were more evenly spaced throughout the genome (Fig. 1d), as previously observed in other primates (e.g. Kong et al. 2002, 2010; Coop et al. 2008; Wall et al. 2022; and see Lenormand and Dutheil 2005; Sardell and Kirkpatrick 2020 for a discussion of this widespread phenomenon). Moreover, in agreement with recent studies in humans (Porubsky et al. 2024) and chimpanzees (Venn et al. 2014), the frequency of CO events decreased with both maternal and paternal age in aye-ayes (with 1.48 and 1.36 fewer COs per year,

Table 1	Number	of CO	events	and	genetic	distances	identified i	n
maternal/paternal meioses								

Scaffold	Length (Mb)	# CO events	сM	cM/Mb
1	316.17	20/18	200/180	0.63/0.57
2	290.59	14/9	140/90	0.48/0.31
3	261.42	18/14	180/140	0.69/0.54
4	219.69	13/14	130/140	0.59/0.64
5	215.45	14/8	140/80	0.65/0.37
6	204.02	16/13	160/130	0.78/0.64
7	199.60	9/16	90/160	0.45/0.80
8	162.77	11/9	110/90	0.68/0.55
10	114.90	12/10	120/100	1.04/0.87
11	102.08	9/10	90/100	0.88/0.98
12	67.30	4/5	40/50	0.59/0.74
13	62.73	11/8	110/80	1.75/1.28
14	34.25	6/5	60/50	1.75/1.46
15	28.26	6/3	60/30	2.12/1.06
Autosomal	2,279.23	163/142	1,630/1,420	0.94/0.77
(sex-averaged)		(152.5)	(1,525)	(0.85)

respectively; Fig. 1e); however, given the small sample size, this observation is not statistically significant (maternal: adjusted $R^2 = 0.1307$, P-value: 0.4583 and paternal: adjusted $R^2 = -0.2682$, *P*-value: 0.5864). Based on the number of COs per meiosis, the sex-averaged autosomal genetic map was estimated to be 1,525 cM in length (Table 1)approximately 25% to 35% shorter than those of catarrhines (Rogers et al. 2000, 2006; Cox et al. 2006; Jasinska et al. 2007; Wall et al. 2022; Versoza et al. 2024b) as may be anticipated from the lower karyotype (2n = 30 in aye-ayes)[Tattersall 1982] vs. 2n = 42 and 60 in rhesus macagues [Owen et al. 2016] and vervet monkeys [Finelli et al. 1999], respectively) (Pardo-Manuel de Villena and Sapienza 2001), which, in turn, exhibit shorter map lengths than hominoids (Kong et al. 2002, 2010; Venn et al. 2014). Similar to other primates, females exhibit an overall longer genetic map length than males (1,630 vs. 1,420 cM)—however, the ratio of the female to male autosomal map length (1.15) is lower than that observed in humans (1.36; Porubsky et al. 2024). The genome-wide average CO rates in males and females were thus estimated to be 0.77 and 0.94 cM/Mb (Table 1)—approximately 40% to 50% lower than the average rates of 1.3 and 2.0 cM/Mb reported in humans (Bhérer et al. 2017)-with the overall lower sex-averaged rate (0.85 cM/Mb) likely contributing to the low levels of genetic diversity observed in the species (Perry et al. 2013; Soni et al. 2025; Terbot et al. 2025b).

The Landscape of NCOs in Aye-Ayes

After excluding both complex events (involving multiple, noncontiguous NCO tracts within <5 kb) and NCOs with tracts >10 kb which were previously shown to frequently represent incorrect genotype calls and/or assembly errors (Smeds et al. 2016; Wall et al. 2022), 145 and 191 putative NCO events were identified through the pedigree-based approach and the family-based approach, respectively. Given the partial overlap between the two-generation nuclear families and the three-generation pedigrees (supplementary fig. S1, Supplementary Material online), NCOs identified in meioses that could be studied using both approaches were subsequently consolidated into 287 distinct NCO events. Careful manual curation using highly stringent guality metrics removed 87 out of these 287 NCOs, either because the event was detected across multiple meioses (85 NCOs) or because the phase-informative markers overlapped with a structural variant in the same individual (1 NCO; Versoza et al. 2024a) or a CO in another individual (1 NCO) (for details, see supplementary table S9, Supplementary Material online). The final dataset thus contained 200 NCOs, 95 and 105 of maternal and paternal origin, respectively (Fig. 2a; and see supplementary table \$10, Supplementary Material online for details). Notably, as expected from NCOs frequently localizing at the same recombination hotspots than COs (with ~70% of NCO events occurring within PRDM9-positioned hotspots in humans; Baudat et al. 2013; Williams et al. 2015; Halldorsson et al. 2016; Palsson et al. 2025), the degenerate hotspot motif (Fig. 1b) was observed in each 10 kb region centered around a NCO event, providing support that these events are likely genuine rather than sequencing artifacts.

On average, 0.71 NCOs were identified per chromosome and meiosis, with 1 to 20 NCOs per meiosis across the 14 autosomes (supplementary table S8, Supplementary Material online). Although guantitatively similar to recent estimates in catarrhines (e.g. 1.06 NCOs per chromosome and meiosis in rhesus macagues [Versoza et al. 2024b] as well as 2.39/2.90 NCOs per chromosome and paternal/maternal meiosis in humans [Palsson et al. 2025]), this represents a conservative estimate, both because NCO events occurring between phase-informative markers will inevitably be missed (a particular challenge for species exhibiting low heterozygosity—and thus low marker density—such as ave-aves) but also because the application of stringent quality metrics required to filter out false positives may have inadvertently removed genuine NCO events. Specifically, given an average of 517,832 phase-informative markers per pedigree/family (supplementary table S3, Supplementary Material online), markers are expected to occur roughly every 4,400 bp and thus, only a small subset of all genuine NCOs will be detectable in this dataset. To estimate the probability of a NCO event being detectable in the dataset on hand, 2,000 NCO events of varying length were simulated using the NCO length distribution recently reported for the best-studied primate, humans, as a reasonable proxy (Supplementary Table 4 in Palsson et al. 2025). As anticipated from the low marker density—and similar to a recent study of >5,000 human trios in which 2.3% (2.39 out of 105.0) and 3.6% (2.90 out of 81.6) of estimated NCO events per offspring were observed



Fig. 2. Characteristics of noncrossovers. a) The genomic distribution of NCO events in females (shown as pink circles) and males (blue circles) across the autosomes (note that scaffold 9, i.e. chromosome X, is not displayed). b) Distribution of sex-specific NCO tract lengths (reported in base pairs [bp]) based on phase-informative sites that mismatched the surrounding haplotype block.

for paternal and maternal meioses, respectively (Palsson et al. 2025)—the probability of identifying NCO events is low for short tracts (2.5% on average) and higher for extended tracts (58.3%), suggesting that the genuine number of NCO events per meiosis is much higher. This difficulty of detecting NCOs from pedigree data is well-appreciated in the limited number of vertebrate studies in which such events have been investigated in depth (see e.g. the discussions in Williams et al. (2015), Smeds et al. (2016), and Palsson et al. (2025)); however, as such, the ratio of NCO:CO cannot be meaningfully compared in this study.

NCOs are slightly less abundant in females than males (8.6 vs. 9.6) and exhibit shorter average minimal tract lengths (157 vs. 163 bp; Fig. 2b and see supplementary table S8, Supplementary Material online)—an observation that might, at least in part, be driven by the lower density of phase-informative markers in females than in males in the

two-generation nuclear families (though note that the difference in marker density is not statistically significant; twosample *t*-test: t = -2.4417; df = 2.9396; *P*-value = 0.09412; supplementary table S3, Supplementary Material online). The sex-averaged mean minimal tract length of 161 bp is similar to those observed in pedigree studies of other primates (with estimated mean tract lengths of 55 to 290 bp in humans [Jeffreys and May 2004; Palsson et al. 2025], 42 to 167 bp in baboons [Wall et al. 2022], and 155 bp in rhesus macaques [Versoza et al. 2024b])—however, it should be noted that differences in marker density will impact resolution, rendering such comparisons across species challenging. In fact, similar to a previous pedigree-based study in humans (Williams et al. 2015), the majority of events (140 out of 200, or 70%) include only a single phase-informative marker, exhibiting a minimal tract length of 1 bp (note that no significant difference between the mutation spectra of the phase-informative markers tagging NCO events, the phase-informative markers in the two-generation nuclear families, and the phaseinformative markers in the three-generation pedigrees was observed; Kruskal–Wallis $\chi^2 = 0.013995$, df = 2, *P*-value = 0.993; and see supplementary table S11, Supplementary Material online). Events involving multiple phase-informative markers generally exhibited short minimal tract lengths (median: 136 bp; mean: 534 bp); however, similar to humans (Williams et al. 2015), baboons (Wall et al. 2022), and rhesus macaques (Versoza et al. 2024b), several NCOs with minimal tract lengths longer than 1 kb were also observed (6 in males and 4 in females; supplementary table \$10, Supplementary Material online). Furthermore, in agreement with empirical patterns observed in other primates (Williams et al. 2015; Halldorsson et al. 2016; Wall et al. 2022; Versoza et al. 2024b; Palsson et al. 2025), the distribution of NCO minimal tract lengths (Fig. 2b) appears more consistent with a powerlaw (or heavy-tailed) distribution than the single geometric distribution frequently modeled (Frisse et al. 2001; Gay et al. 2007; Yin et al. 2009).

Based on the observed minimal tract lengths and an autosomal genome length of 2.28 Gb (supplementary tables S8 and S10, Supplementary Material online), aye-ayes exhibit a sex-averaged NCO rate of 6.8×10^{-7} per base pair per generation (95% CI: 2.9×10^{-7} to 1.1×10^{-6} /bp/gen), with similar rates between the sexes (6.4×10^{-7} /bp/gen in females vs. 7.4×10^{-7} /bp/gen in males; Mann–Whitney *U* test, *P*-value = 0.38) and an expected average of ~1.5 kb of autosomal sequence affected by NCOs in each generation. The actual sex-averaged NCO rate is likely much greater given the small sample size and low marker density of this study and, taking the simulation results at face value, is likely closer to 2.3×10^{-6} /bp/gen, affecting an estimated ~50 kb per generation.

In summary, the landscape of CO and NCO events in aye-ayes presented here provides novel insights into a parameter crucial for improving population genetic modeling in this highly endangered species, and will serve as a valuable resource for future comparative genomic studies seeking to understand the long-term evolution of recombination landscapes across the primate clade.

Materials and Methods

Sequencing, Read Mapping, and Variant Calling

A total of 14 aye-aye individuals (eight females and six males) were selected from six three-generation pedigrees and three (partially overlapping) two-generation multi-sibling families available at the Duke Lemur Center, the genomes of which were sequenced to an average coverage of 50× per individual (supplementary table S1, Supplementary Material online). Sequencing reads were mapped to the aye-aye genome assembly (DMad_hybrid; GenBank accession number:

GCA_044048945.1; Versoza and Pfeifer 2024) in order to call autosomal, biallelic single nucleotide polymorphisms (SNPs) genotyped in all individuals (see Versoza et al. (2025) for details regarding the autosomes and Terbot et al. (2025a) for a comparison with chromosome X). In the absence of a curated "gold standard" variant dataset that could serve as a basis for variant filtration in the species, initial calls were filtered using the Genome Analysis Toolkit's Best Practice "hard filter" criteria for germline variants (van der Auwera and O'Connor 2020). Following the methodology described by Versoza et al. (2024b), a set of additional coverage-, genotype-, and inheritance-based filters was applied to further increase the precision of the variant call set. Specifically, variants were excluded if (i) they were located within repetitive and low-complexity regions of the genome prone to base calling errors and mis-mappings (Pfeifer 2017), or if they exhibited (ii) a total depth of coverage (DP) of less than half or more than twice the average autosomal coverage for each sample, (iii) a genotype quality (GQ) less than 30 (corresponding to a probability of a genotyping error of more than 10^{-3}), (iv) an excess of heterozygosity (defined here as a P-value of 0.01), and/or (v) violated the patterns expected by Mendelian inheritance (as determined using the BCFtools + mendelian2 plugin [Danecek et al. 2021] together with the information of the pedigree). Additionally, to further reduce variant calling and/ or genotyping errors that could potentially lead to spurious recombination events, SNPs located within either (i) a cluster of variants (defined here as \geq 3 SNPs within a 10 bp window), (ii) 10 bp of an insertion/deletion, or (iii) 2 Mb from the chromosome ends, were also excluded from further analyses. The resulting high-confidence call set consisted of 1,830,431 SNPs with a transition-transversion ratio of 2.62 (supplementary table S2, Supplementary Material online).

Identification of CO and NCO Events

The high-confidence call set was divided into six threegeneration pedigrees and three (partially overlapping) two-generation nuclear families with multiple offspring (supplementary fig. S1, Supplementary Material online) for which gamete transmission could be tracked in order to identify recombination events based on "phase-informative" markers-that is, heterozygous SNPs for which the parent-of-origin could be determined (supplementary table S3, Supplementary Material online). In brief, in the threegeneration pedigrees, phase-informative markers were sites at which the P_0 individuals exhibited nonidentical genotypes, their F₁ offspring was heterozygous, and either the F₁'s partner or their joint F2 offspring was homozygous (for a schematic of the workflow, see Fig. 1b in Versoza et al. 2024b). In the two-generation nuclear families, maternally phase-informative markers exclusively included those for which the dam was heterozygous and the sire homozygous, whereas paternally phase-informative markers included

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sites at which the sire was heterozygous and the dam homozygous. Following the methodology outlined in Coop et al. (2008), recombination events were then identified by comparing these phase-informative markers across siblings using a randomly assigned offspring as "template" (note that although recombination events can be identified in a single sibling pair using this approach, it is not possible to unequivocally assign the meiosis in which the event occurred; see supplementary fig. S20, Supplementary Material online for a schematic of the workflow). In both cases, heterozygous genotype calls in phase-informative markers required the support of more than 25% but less than 75% of mapped reads to limit genotyping errors in the downstream analyses.

Based on these phase-informative markers, recombination events were classified as either COs—that is, a single change of phase along a chromosome (from maternally inherited to paternally inherited haplotype blocks or vice versa, ignoring single phase-informative markers within a region of consistent phase)-or NCOs-that is, phase-informative markers that mismatched surrounding haplotype blocks. CO interference prevents the occurrence of two CO events in close proximity to each other (Otto and Payseur 2019), thus following Venn et al. (2014), a minimum distance of 1 Mb was required between two COs that originated from the same meiosis (see supplementary table S5, Supplementary Material online for a summary of the removed CO events). Similarly, following Smeds et al. (2016), to reduce the number of spurious NCO events resulting from genotyping errors, a minimum distance of 5 kb was required between two consecutive NCOs; additionally, complex NCO events longer than 10 kb were excluded from further analyses as previous work demonstrated that these frequently represent assembly errors (Wall et al. 2022). For the four meioses that could be studied using both the three-generation pedigree approach and the twogeneration family approach (i.e. focal individual-offspring pairs 7-12, 7-13, 8-12, and 8-13; see supplementary fig. S1, Supplementary Material online), recombination events were manually consolidated. Notably, all CO events in these meioses were identified by both approaches, with the majority of events (54%) sharing identical start and end coordinates at the break point. In case of a partial overlap (i.e. a sharing of either start or end coordinates at the break point owing to different phase-informative markers in the two approaches), the CO events with the shortest resolution were selected. Shorter CO resolution tracts were generally observed using the pedigree-based approach (only 24% of events exhibited shorter resolutions in the family-based approach) due to a larger number of phase-informative markers (supplementary table S3, Supplementary Material online, and see supplementary table S4 and figs. S2 to S15, Supplementary Material online for the marker density across each autosome per pedigree/family). In contrast, due to the low density of markers and the short tract lengths, only 48 of the 120 NCO events (40%) were detected with both approaches. Of these,

83% exhibited identical break point coordinates; for the remaining, partially overlapping events, the NCOs with the longest length tract were selected. Lastly, to guard against assembly errors, CO and NCO events detected in more than one meiosis and those overlapping with structural variation (Versoza et al. 2024a) were removed (see supplementary tables S6 and S9, Supplementary Material online for COs and NCOs, respectively).

Annotation

High-confidence COs (defined here as COs with a resolution shorter than 5 kb) and NCOs (with a tract length shorter than 5 kb) were classified by genomic region (i.e. intergenic, upstream, exonic, exonic noncoding RNA [ncRNA], intronic, intronic ncRNA, 3' and 5' UTR, and downstream) and repeat content using ANNOVAR v.2020-06-08 (Wang et al. 2010) together with information from the aye-aye genome annotation (Versoza and Pfeifer 2024). CO and NCO events were checked for overlap with structural variants cataloged in the individuals of this study (Versoza et al. 2024a). Statistics were calculated and plotted using R v.4.2.2 (R Core Team 2022). Ideograms of the genome-wide maps were plotted with Rldeogram v.0.2.2 (Hao et al. 2020).

Motif Identification

CO events with a high resolution were leveraged to search for putative sequence motifs of PRDM9 binding sites in aye-ayes. Following Wijnker et al. (2013), the 110 CO regions with a resolution of less than 10 kb were extended to include the neighboring 500 bp in order to ensure that the complete region subject to double-strand break and repair was included, and the de novo motif finding algorithm implemented in MEME v.5.5.7 (Bailey and Elkan 1994) was run using the zero or one occurrence per sequence (ZOOPS) model. To account for potential nucleotide sequence bias, the search was corrected for genomic background using a model built from randomly selected nonbreakpoint regions with the same genomic characteristics (i.e. sequence length and GC content) as the breakpoint regions. Given that many human recombination hotspots harbor a degenerate 13-mer sequence motif (Myers et al. 2008), the search was limited to motifs with a minimum and maximum width of 5 and 15 bases, respectively. A single degenerate motif passed the discovery threshold of 1e-05. Similar to humans (Alternose et al. 2017), this degenerate motif is highly repetitive, and thus, a permutation test was performed to compare its distribution in breakpoint regions with that in nonbreakpoint regions. To this end, an additional 1,000 random nonbreakpoint regions were sampled and MOODS v.1.9.4.1 (Korhonen et al. 2009) was used to scan both breakpoint and nonbreakpoint regions for occurrences of the degenerate motif based on a positional weight matrix, considering matches with a P-value < 0.05. A Fisher's exact test was performed using R v.4.2.2 (R Core Team 2022) to test for statistical significance of the observed differences.

PRDM9–DNA Binding

In many primates, the location of recombination is mainly determined by the zinc-finger protein PRDM9 binding specific DNA sequence motifs (Baudat et al. 2010; Myers et al. 2010; Parvanov et al. 2010). In order to characterize PRDM9-DNA binding in silico, the nucleotide sequence of PRDM9 in aye-ayes was obtained from the recent genome annotation of the species (Versoza and Pfeifer 2024), visually checked for completeness and potential frameshift errors by alignment against the human PRDM9 protein sequence included in the Universal Protein Knowledgebase (UniProt Consortium 2025) using GeneWise v.2.4.1 (Birney et al. 2004), and translated into a protein sequence using the ExPASy web server (Appel et al. 1994; Gasteiger et al. 2003; Artimo et al. 2012; Duvaud et al. 2021). A BlastP (Altschul et al. 1990) search of the aye-aye PRDM9 protein sequence against the NCBI reference database demonstrated a high sequence similarity with PRDM9 in other primates (supplementary fig. S21, Supplementary Material online) and a multiple-sequence alignment with the PRDM9 protein sequence in humans provided evidence that the KRAB domain, the PR/SET domain, and the zinc-finger array are wellcharacterized (supplementary fig. S22, Supplementary Material online). The presence of these domains and the zincfinger array in ave-ayes was additionally supported by the protein sequence classification obtained from InterPro (Blum et al. 2025) (supplementary fig. S23, Supplementary Material online). With both the PRDM9 protein sequence and a putative degenerate binding motif (see "Motif identification") on hand, AlphaFold 3 (Abramson et al. 2024) was used to predict PRDM9–DNA binding.

Detectability of NCO Events

The detection of NCO events is complicated by both the short tract lengths and the need for a segregating variant to be present in the donor homolog in order to allow identification-a particular challenge for highly endangered species with low levels of heterozygosity and consequently low marker density. To estimate the probability of a NCO event being detectable in the aye-aye dataset on hand, 2,000 NCO events of varying length were simulated using the NCO length distribution recently reported for humans (Supplementary Table 4 in Palsson et al. 2025) as a reasonable proxy (as precise estimates of NCO tract lengths remain elusive in many species, including aye-ayes). Specifically, 500 maternal short NCO events were simulated based on a mean NCO length of 102 bp (95% CI: 71 to 125 bp), 500 paternal short NCO events were simulated based on a mean NCO length of 123 bp (95% CI: 94 to 135 bp), 500 maternal extended NCO events were simulated based on a mean NCO length of 9.1 kb (95% CI: 6.8 to 10.8 kb), and 500 paternal extended NCO events were simulated based on a mean NCO length of 7.2 kb (95% CI: 1.8 to 11.8 kb), and the number of NCO events containing at least one phase-informative marker was recorded. Power was calculated by dividing the number of detectable NCOs by the total number of simulated NCO events in each category, with sex-specific events combined as size differences between sexes were not significant in humans (Palsson et al. 2025).

Supplementary Material

Supplementary material is available at *Genome Biology and Evolution* online.

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Conflict of Interest

None declared.

Data Availability

All sequence data have been deposited under NCBI BioProjects PRJNA1179987 and PRJNA1181251. Computational workflows for the detection of crossover and noncrossover events were implemented using Snakemake v.7.32.4 (Mölder et al. 2021) and are available via GitHub: https://github.com/ PfeiferLab/aye-aye_CO_NCO.

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