

Single-haplotype comparative genomics provides insights into lineage-specific structural variation during cat evolution

Received: 23 June 2023

Accepted: 20 September 2023

Published online: 02 November 2023

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The role of structurally dynamic genomic regions in speciation is poorly understood due to challenges inherent in diploid genome assembly. Here we reconstructed the evolutionary dynamics of structural variation in five cat species by phasing the genomes of three interspecies F1 hybrids to generate near-gapless single-haplotype assemblies. We discerned that cat genomes have a paucity of segmental duplications relative to great apes, explaining their remarkable karyotypic stability. X chromosomes were hotspots of structural variation, including enrichment with inversions in a large recombination desert with characteristics of a supergene. The X-linked macrosatellite *DXZ4* evolves more rapidly than 99.5% of the genome clarifying its role in felid hybrid incompatibility. Resolved sensory gene repertoires revealed functional copy number changes associated with ecomorphological adaptations, sociality and domestication. This study highlights the value of gapless genomes to reveal structural mechanisms underpinning karyotypic evolution, reproductive isolation and ecological niche adaptation.

Comparative genomics is a powerful approach for inferring the genetic basis of adaptation and speciation. Its success depends on accurate and representative whole-genome alignments that precisely quantify genetic similarities and differences between evolutionary lineages to make predictions regarding the impact of genomic divergence on phenotypic evolution and diversification. The application of long-read sequencing has enabled increasingly precise comparisons between taxa, facilitating the assembly of 92–96% of a diploid genome

sequence into chromosomes^{1,2}. However, tracing the evolutionary history of regions of high structural complexity and allelic divergence has remained challenging. Until the completion of the human telomere-to-telomere (T2T) project^{3–5}, genomic ‘dark matter’ (refs. 6,7) that encompasses satellite arrays, centromeres, segmental duplications (SDs) and complex gene families had been missing from nearly all comparative genomic studies. Consequently, for most species, we still have a limited understanding of the evolutionary dynamics of the

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most repetitive genomic sequences and how their divergence manifests in reproductive isolation and phenotypic innovation.

The cat family Felidae represents a speciose and successful apex predator radiation that occupies diverse biomes across the globe. Previous comparative genomic studies have illuminated their rapid diversification in the Miocene^{8,9}, frequent postspeciation gene flow^{9,10}, the impacts of demographic changes on genetic diversity and fitness^{11–13}, and the genetic consequences of domestication¹⁴. Here we applied the trio-binning approach¹⁵ to three divergent interspecific crosses amenable to high-resolution haplotype phasing (Fig. 1a) to generate near-gapless genome assemblies from multiple species pairs along the felid phylogeny. Comparisons of these assemblies provided an unprecedented glimpse into the properties of large and complex gene families and functional repetitive elements that were previously inaccessible^{14,16,17}. We describe insights into the cauldron of repetitive genetic variation with potentially large effects on chromosome function and speciation.

Results

Phased genome assembly reveals remarkable collinearity

We used long-read PacBio sequencing to phase and assemble six single-haplotype genomes from five cat species (domestic cat, leopard cat, Geoffroy's cat, tiger and lion) through the application of trio-binning to three F1 interspecies hybrids¹⁵. The parent species of the crosses diverged ≥ 4 million years ago (MYA; Fig. 1a), enabling $>99.5\%$ of the long sequence reads to be accurately phased into the parental haplotypes¹⁸ (Fig. 1a, Supplementary Figs. 1–4 and Supplementary Table 1). De novo assembly produced ultracontiguous assemblies with contig N50 = 77–104 Mb (Table 1 and Fig. 1b). At least 99.6% of the euchromatic sequence was assembled into chromosome-length scaffolds using high-throughput chromosome conformation capture (Hi-C; Supplementary Fig. 5), with an average of just 53 gaps per genome assembly, 15 gapless chromosomes across all species and 62% of the assembled autosomes containing two or fewer gaps (Fig. 1c and Supplementary Fig. 6), exceeding comparable parameters from all other domestic species reference assemblies (Fig. 1b). The canonical telomeric sequence shared by vertebrates is TTAGGG¹⁹; however, different blocks of microsatellites are found in telomeres of other species of the generalized pattern (TxAYGz)²⁰. To determine which chromosome assemblies extended into one or both telomeres, we searched for telomere-like repeat sequences by requiring 80% of the terminal 100 bases of the chromosome to be labeled as a repeat family or a tandem repeat. Then, we extended the search window progressively. About 61% of the chromosomes in the six assemblies likely extend into both telomeres, 32% extend into one telomere and the remaining 7% lack terminal repeats and are likely incomplete. Only 32% of the assembled chromosomes possess the canonical TTAGGG tandem array at the telomere, while 21 chromosomes terminated with the FA-satellite^{21,22} (Supplementary Table 2).

Pairwise whole-genome alignments between the five species' assemblies revealed near-complete karyotypic stasis after they diverged from a common ancestor ~ 11 to 15 MYA^{9,10} (Fig. 1d). The only change in chromosome number is a single Robertsonian translocation of two small acrocentrics (chrF1 and chrF2), producing a medium-size metacentric (chrC3) shared by all species of the neotropical cat genus *Leopardus* (Fig. 1e,f)²³. Close inspection of alignments between *Leopardus geoffroyi* and *Felis catus* showed that chromosome C3 was the product of a centric fusion, followed by a near chromosome arm-length inversion that reoriented $>99\%$ of C3q relative to the ancestral chrF2 homolog (Fig. 1g). All other chromosomal rearrangements between species were inversions several orders of magnitude smaller in size (<2 Mb; Fig. 2a and Supplementary Table 3). We identified 172 fixed inversions >50 bp (Fig. 2a) across the five species phylogeny that samples >50 million years of independent branch length. By comparison, great ape genomes contain the products of 1,326 fixed

inversions >50 bp (ref. 1) (Fig. 2a). Felids and great apes diverged on a very similar evolutionary timescale, matching nearly 1:1 for divergence events (Fig. 2a). Given the similarity in sampled evolutionary history, great ape genomes possess 7.7-fold more rearrangements than felids suggesting that great ape genomes are more structurally prone to chromosome rearrangement than felids.

SDs have been hypothesized to be major drivers of chromosome evolution and disease susceptibility in the great ape lineage by promoting nonallelic homologous recombination^{24,25}, particularly because of their uniquely interspersed distribution²⁶. In support of this hypothesis, SDs flank 82–86% of known primate inversions²⁷. To determine whether SDs might be a primary driver of felid inversions, we used SEDEF²⁸ to identify SDs in each cat haplotype. The total bases in felid SDs range from 25 to 35 Mb, or 1% to 1.5% of each genome (Supplementary Fig. 7). By comparison, the SD frequency (7%) estimated in the human T2T genome²⁹ is five- to seven-fold higher than in felid genomes. Compared to great apes, the similar-fold reduction in chromosomal rearrangements and SD frequency in felid genomes supports the hypothesis that the overall frequency of SDs is the primary driver of chromosome evolution in these two lineages. Future analysis of near-gapless genomes in other mammalian lineages with highly variable rates of karyotypic evolution will enable the testing of this hypothesis.

Structural variation is enriched on chromosome X

The hemizygous nature of the X chromosome (chrX) in male heterogametic taxa promotes faster rates of evolution relative to the autosomes and the accumulation of loci associated with reproductive isolation and speciation^{30,31}. Previous studies revealed a higher fixation rate of inversions on chrX relative to autosomes^{32,33}. In cats, chrX was an outlier in terms of the number of inversions relative to chromosome length (Fig. 2b). For each branch in the phylogeny, the mean inversion was significantly larger on chrX than the autosomes (Fig. 2c). Inversions accumulated disproportionately in an ~ 45 -Mb recombination cold spot on chrX that is enriched for barriers to gene flow across multiple felid lineages¹⁰ (Fig. 2d). Two thirds (24/36) of the X-linked inversions were fixed versus polymorphic (Supplementary Table 3). About 70% of fixed inversions harbored at least one protein-coding gene (mean 1.3 genes/fixed inversion). In contrast, only 33% of polymorphic inversions spanned or overlapped with a single protein-coding gene. In half of these cases, the inversion was located within a long intron (Supplementary Table 4). These results support previous observations in insects³³ and suggest that the fixed X-linked inversions within the 45-Mb recombination cold spot may harbor beneficial alleles given their longer length and enrichment with protein-coding genes. Previous studies of small and big cats identified signatures of natural selection within the large recombination cold spot^{14,34}. We hypothesize that this gene-rich, inversion-rich region is a major X-linked supergene locus underpinning felid reproductive isolation that warrants future comparative genomic analyses.

Satellite elements have been implicated in speciation but are poorly represented in diploid genome assemblies^{35,36}. Cat chrX harbors the only X-linked speciation gene identified in mammals; the macrosatellite repeat *DXZ4* (ref. 37). *DXZ4* has been well studied regarding its putative role in mammalian chrX inactivation (XCI). Human *DXZ4* consists of a single 3-kb tandem repeat array containing 56 monomers, where each repeat contains a single CTCF-binding site⁴ (Fig. 3a). Long noncoding RNAs (*DANT1* and *DANT2*) expressed from *DXZ4* on the inactive chrX (Xi) promote superlooping with other macrosatellites on the Xi³⁸ and facilitate the localization of the Barr body in female placental mammals to the nucleolar membrane³⁹ (Fig. 3a). The human T2T genome assembly first completely resolved the *DXZ4* array structure, but a complete assembly of *DXZ4* sequences in other mammalian taxa is largely lacking, clouding our understanding of its evolution and function. *DXZ4* was resolved in all six cat assemblies, revealing a unique

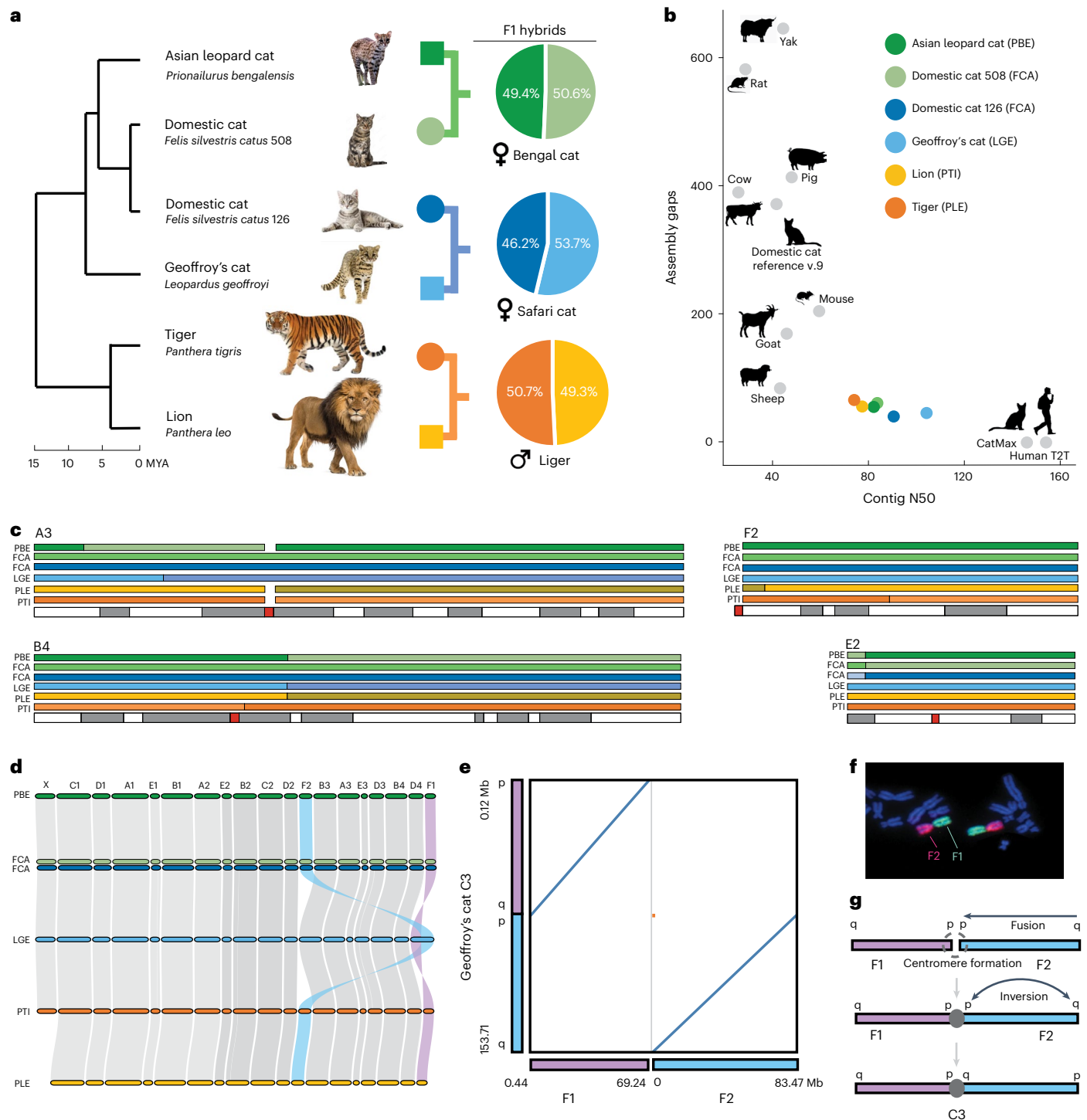


Fig. 1 | Assembly and synteny comparisons among the genomes of five cat species. a, Phylogeny and timescale of the parent species of the three hybrid trios used for assembly and comparative analysis. Pie charts illustrate the phasing results (% of total reads) for the F1 PacBio CLR long reads. **b**, Comparison of contig N50 statistics and number of assembly gaps against other highly contiguous mammalian reference genomes from domestic species. CatMax refers to the theoretical N50 maximum based on domestic cat chromosome sizes. PBE, *Prionailurus bengalensis*; FCA, *Felis catus*; LGE, *Leopardus geoffroyi*; PTI, *Panthera tigris*; PLE, *Panthera leo*. **c**, Contig alignments for the six felid single-haplotype assemblies from chrsA3, B4, E2 and F2/C3 to the felCat9 diploid domestic cat long-read genome assembly, depicted on the bottom as a G-banded ideogram. Inferred centromere locations are indicated by red bars.

The bars above each ideogram are colored by species and represent assembly contigs > 1 Mb. Breaks between contigs are indicated by a black line and a shift in color contrast. The full set of chromosome alignments is found in Supplementary Fig. 6. **d**, Synteny plot⁷⁰ illustrating extensive collinearity of the five species assemblies. Blue and purple alignment tracks highlight the only chromosome number change in Felidae, the Robertsonian fusion of chrF1 and chrF2 present in all felid genera, and the derived C3 chromosome observed in Geoffroy's cat and all species of the genus *Leopardus*. **e, f**, Dot plot alignment (left) of Geoffroy's cat chrC3 and domestic cat chrF1 and chrF2 (e) (illustrated with multicolor FISH in f). **g**, Note the orange alignment fragment (in e) indicating a small centromeric fragment of chrF2 that defines the inversion breakpoint on the ancestral chrF2.

Table 1 | Felid single-haplotype genome assembly statistics

Species/hybrid sequenced	Domestic cat-508 (Bengal cat F1)	Domestic cat-126 (Safari cat F1)	Asian leopard cat (Bengal cat F1)	Geoffroy's cat (Safari cat F1)	Lion (liger F1)	Tiger (liger F1)
Sex of parent haplotype	♀	♀	♀	♀	♂	♀
Chromosomes	18, X	18, X	18, X	17, X	18, Y	18, X
Contigs	123	103	132	88	103	135
Largest contig	205,171,639	172,124,406	240,846,738	239,106,607	166,870,000	166,130,000
Ungapped assembly length (Mb)	2,422,283,418	2,425,722,929	2,435,689,660	2,426,362,316	2,297,542,863	2,408,668,598
Contig N50 (Mb)	84,507,663	92,686,623	83,696,501	104,474,415	77,781,637	74,360,613
Scaffolds	71	70	83	46	53	74
Total assembly length (Mb)	2,422,299,418	2,425,747,038	2,435,718,761	2,426,370,816	2,297,568,983	2,408,695,688
Scaffold N50 (Mb)	147,603,332	148,491,486	148,587,958	152,606,360	147,402,474	146,942,463
Chromosome gaps	60	39	56	45	55	65
Complete BUSCO genes (mammalia_odb10)	8,621	8,619	8,621	8,612	8,417	8,630
Percent complete	93.4	93.4	93.4	93.3	91.2	93.5
Single copy	8,599	8,596	8,599	8,592	8,383	8,601
Duplicated	154	160	154	152	143	147
Missing	451	447	451	462	666	449
Complete+partial (%)	95.1	95.2	95.1	95.0	92.8	95.1

compound tandem repeat composed of two highly divergent (mean P distance = 0.67) repeat arrays, RA and RB (Fig. 3b). Both monomer types contain CTCF-binding sites, but notably differ in the number and orientation of the sites that are important for CTCF-binding affinity and loop extrusion directionality⁴⁰, suggesting divergent superlooping functions between the arrays. The human and mouse genomes notably lack the RB array.

Studies using interspecific backcross hybrids of the domestic cat and Jungle cat (*Felis chaus*) identified *DXZ4* as a major-effect hybrid male sterility locus, with a likely role in reproductive isolation within the *Felis* genus³⁷. The testicular germ cells of sterile male hybrid cats possess RA-specific methylation defects and *DANTI* misregulation, culminating in the failure of meiotic sex chromosome inactivation (MSCI) and meiotic arrest, hallmark phenotypes in mammalian interspecies hybrids³¹. Evidence that *DXZ4* functions in male meiotic silencing was intriguing, given the parallels between the heterochromatic Barr body formed during female XCI and the condensed X-Y body in male MSCI. Although the hybrid sterility phenotype was attributed to *DXZ4* interspecific divergence, the precise mechanism is not well understood. Here our expanded sampling of felid genomes demonstrates that the compound RA and RB repeat structure is copy number variable across all species (Fig. 3b), suggesting copy number-mediated expression effects may have an important role in speciation in other felids. In addition, StainedGlass⁴¹ plots illustrate the rapidity of *DXZ4* repeat array sequence divergence (Fig. 3c). RA and RB arrays evolve two- to three-fold faster than the flanking and intervening noncoding spacer sequences. Notably, a genome-wide analysis of pairwise interspecific genetic divergence calculated across 28,312 5-kb alignment windows (94.1% of the multispecies alignment) placed *DXZ4* RA in the top 0.5% of the most rapidly evolving genomic loci (Fig. 3d), supporting its role as a speciation gene³⁷.

To determine whether the compound *DXZ4* array structure in cats is the exception or the rule in placental mammals, we searched for *DXZ4* arrays in long-read genome assemblies from species representing divergent superorders (Fig. 3e and Supplementary Figs. 8–11). Most assemblies possessed a gap within or adjacent to the predicted

position of *DXZ4* (Supplementary Figs. 12 and 13). We were able to recover sufficient repeat array resolution at the edge of some assembly gaps to characterize the CTCF array. Although the *DXZ4* monomer sequence diverges rapidly to the point of phylogenetic saturation and lack of phylogenetic patterning (Supplementary Fig. 14), we observed the conservation of the CTCF-binding motif patterns across species from different ordinal lineages. Euarchontoglires (for example, primates, rodents and rabbits) possessed only RA or RB, while members of Laurasiatheria possess RA, RB or both types (Fig. 3e). RA and RB were, therefore, present in the most recent common ancestor of boreoeutherian mammals. Moreover, the repeat unit length is relatively constrained (between 3 and 4.9 kb) across species despite rapid sequence divergence and little conservation outside the CTCF motif⁴². Given this unusual combination of spatial and structural evolutionary conservation and an extremely fast rate of sequence evolution, we predict that *DXZ4* satellite divergence may have a more widespread role in establishing and maintaining species boundaries in other mammalian clades.

Intriguingly, all sampled species from the family Bovidae lack *DXZ4* in their assembly, suggesting they may have evolved compensatory mechanisms for its putative loss. Multiple studies have shown that ablation of *DXZ4* has no significant impact on the silenced state of the inactive chrX in mouse and human cells^{40,43}. Nonetheless, the high degree of syntenic, CTCF⁴² and spatial conservation of the *DXZ4* repeat array over the past 104 million years of the placental mammal radiation suggest that *DXZ4* expression and long-range chromatin interactions are functionally important for some heretofore unidentified cellular role during XCI and MSCI⁴⁴. Pan-autosomal gene downregulation is one noteworthy cellular phenotype shared by in vivo *DXZ4*-knock-out mice⁴⁵ and sterile feline interspecific hybrid testes³⁷. These observations raise the possibility that *DXZ4*, acting alone or in concert with other X-linked macrosatellites, may function in RNA-dependent, chrX-autosomal crosstalk associated with the chrX 'counting' process in XCI⁴⁵ and proper sequestration of the DNA damage response factors from the autosomes to the X-Y body during MSCI^{46,47}. Gapless chrX assemblies from a diverse sampling of mammalian genomes will be

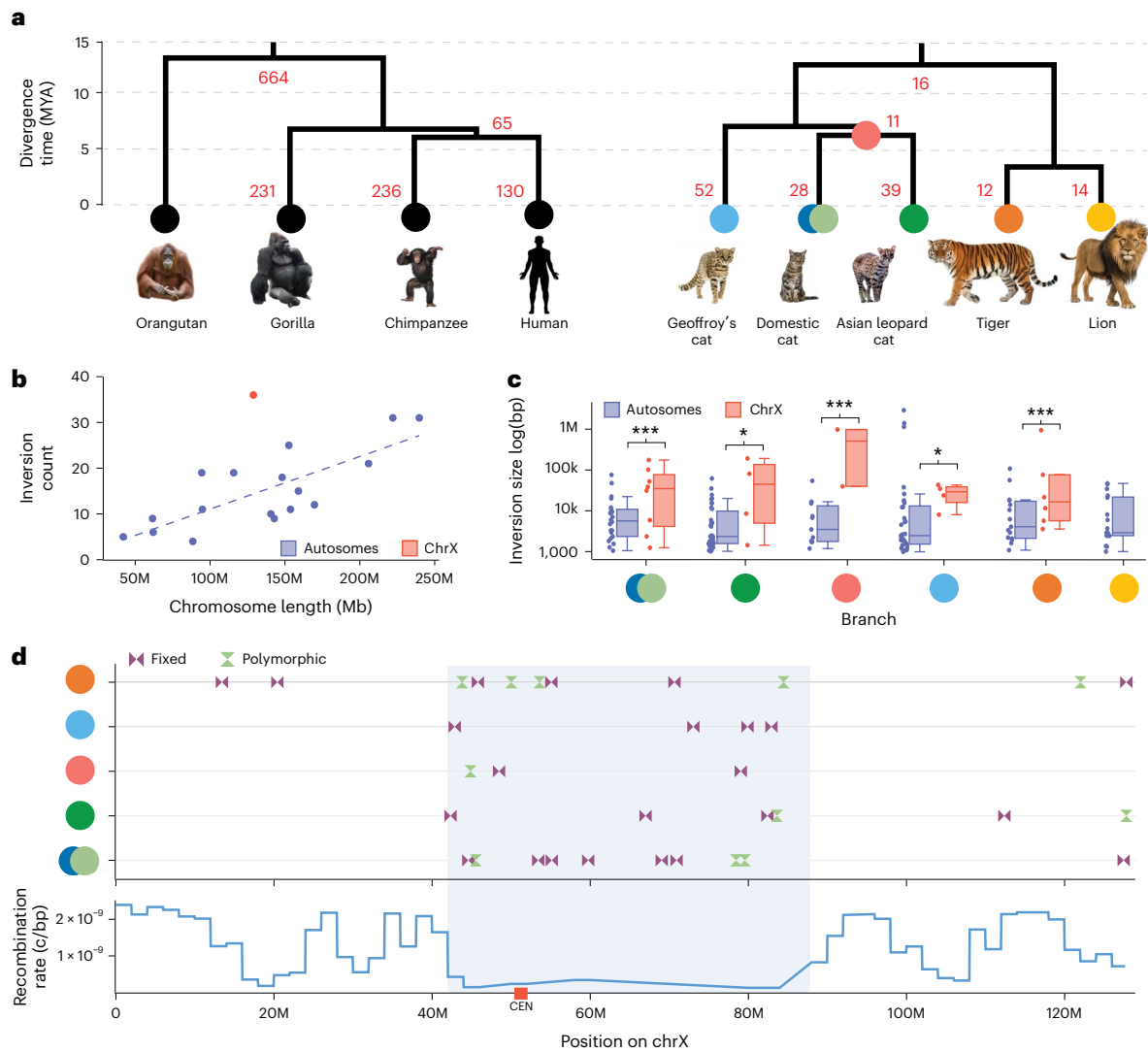


Fig. 2 | Felid structural variation. **a**, Comparison of fixed inversions (red numbers) plotted on branches of the phylogeny of felids (right) and great apes (left) (5). Note the similar divergence times between ape and felid species sampled. **b**, Per chromosome inversion counts plotted against chromosome length. Autosomes are indicated with blue dots and chrX in red. **c**, Comparison of inversion size between the autosomes and chrX for each branch of the phylogeny (colored dots) shown in **a** (except for the lion genome, which is derived from the paternal haplotype of the male F1 liger; Supplementary Table 3). A one-sided Wilcoxon rank sum test determined significance ($*P < 0.05$, $***P < 0.01$). Domestic cat ($n = 40$ autosomal inversions, $n = 11$ chrX inversions, $U = 2.52$, $P = 5.9 \times 10^{-3}$), Geoffroy's cat ($n = 33$ autosomal inversions, $n = 4$ chrX inversions, $U = 2.15$, $P = 1.6 \times 10^{-2}$), Asian leopard cat ($n = 40$ autosomal inversions, $n = 6$ chrX inversions, $U = 1.92$, $P = 2.7 \times 10^{-2}$), domestic cat + Asian leopard cat

($n = 17$ autosomal inversions, $n = 3$ chrX inversions, $U = 2.59$, $P = 4.8 \times 10^{-3}$), tiger ($n = 34$ autosomal inversions, $n = 11$ chrX inversions, $U = 2.54$, $P = 5.6 \times 10^{-3}$), lion ($n = 34$ autosomal inversions). Box plots show the interquartile range with the center line representing the median. Whiskers indicate the highest and lowest value within the upper and lower fences (upper fence = 75% quantile + 1.5 × interquartile range, lower fence = 50% quantile - 1.5 × interquartile range). **d**, The physical distribution of fixed and polymorphic inversions (Supplementary Table 4) on chrX for each branch of the phylogeny relative to the tiger genome. The chrX genome sequences are otherwise collinear across species. A tiger recombination map estimated from population genomic data (Supplementary Fig. 30) is depicted at the bottom (Methods) and is highly conserved with the recombination rate profile of the domestic cat chrX^{9,71}. The shaded area refers to a large recombination cold spot shared with domestic cats, humans and pigs^{9,10}.

critical to understanding the functional relevance of *DXZ4* in the chrX biology of placental mammals.

Variation in centromere structure and size

Current human and great ape centromere sequence models portray large tandem repeat arrays of α satellites flanked by other satellite repeat types, SDs, transposable elements and even some genes⁴⁸. Whether centromere structure is conserved across mammalian lineages is poorly understood because they are not sequence-resolved in most genome assemblies. Therefore, we sought to determine whether our assemblies possessed genomic signatures characteristic of centromeric satellites⁵. Given the absence of previously annotated centromeric

sequences, we first characterized the overall landscape of feline repetitive elements to enable de novo prediction of the most probable centromeric satellites (Supplementary Fig. 15). Interspersed repeats comprise 38% of each genome with a marked distinction between Felinae (*Felis*, *Prionailurus* and *Leopardus*) and *Panthera*, with Felinae showing an average SINE insertion rate of $\sim 2.7\times$ higher than *Panthera*, while conversely, the LINE insertion rate in *Panthera* is $\sim 1.6\times$ higher than Felinae (Supplementary Fig. 16).

Next, we searched for novel repeat enrichment within narrowly defined chromosomal regions for which we had a strong priori evidence classifying that region as centromere-containing based on integrative analysis of comparative mapping approaches^{9,14,17}

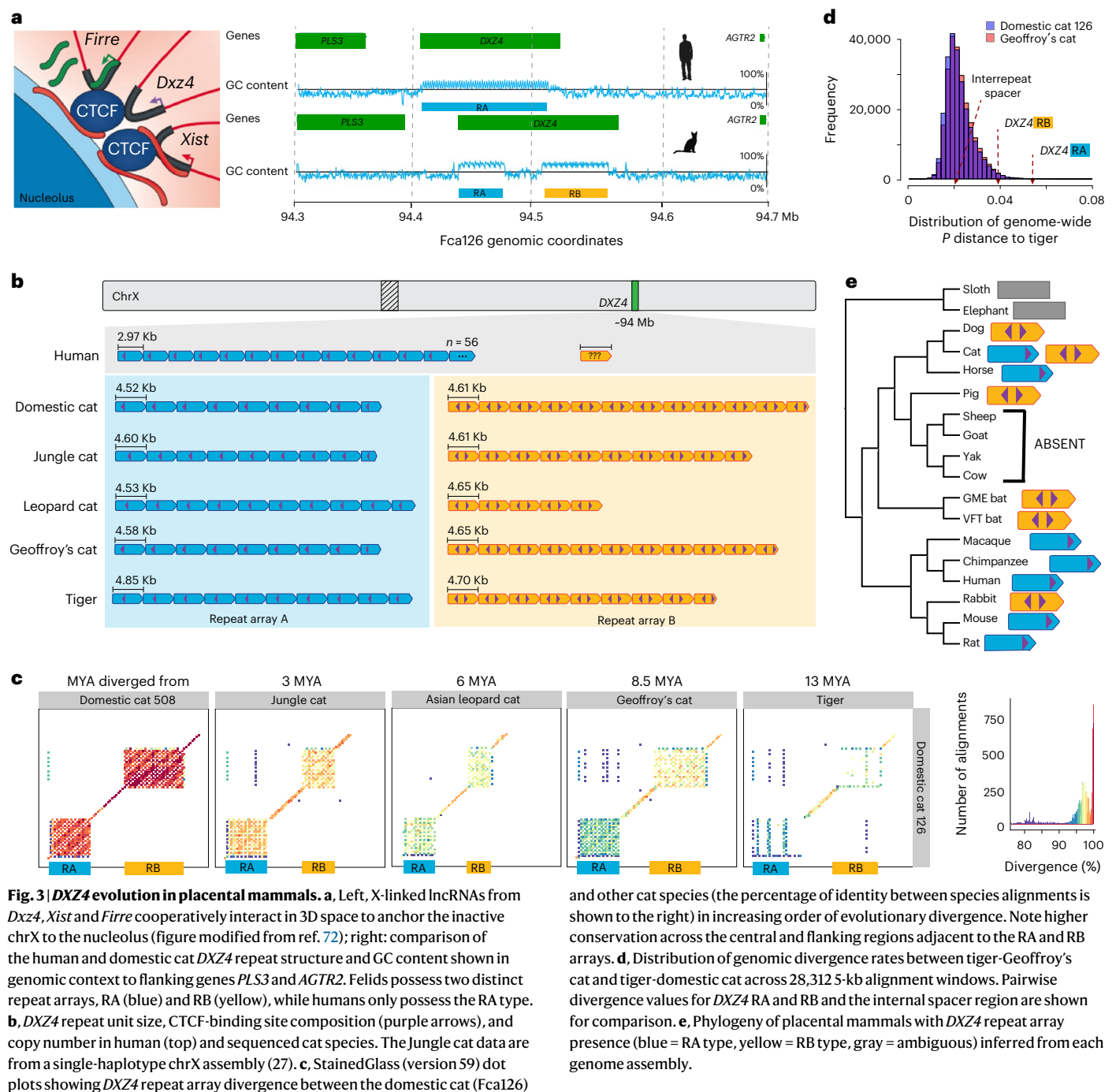


Fig. 3 | *DXZ4* evolution in placental mammals. **a**, Left, X-linked lncRNAs from *Dxz4*, *Xist* and *Firre* cooperatively interact in 3D space to anchor the inactive chrX to the nucleolus (figure modified from ref. 72); right: comparison of the human and domestic cat *DXZ4* repeat structure and GC content shown in genomic context to flanking genes *PLS3* and *AGTR2*. Felids possess two distinct repeat arrays, RA (blue) and RB (yellow), while humans only possess the RA type. **b**, *DXZ4* repeat unit size, CTCF-binding site composition (purple arrows), and copy number in human (top) and sequenced cat species. The Jungle cat data are from a single-haplotype chrX assembly (27). **c**, StainedGlass (version 59) dot plots showing *DXZ4* repeat array divergence between the domestic cat (Fca126)

and other cat species (the percentage of identity between species alignments is shown to the right) in increasing order of evolutionary divergence. Note higher conservation across the central and flanking regions adjacent to the RA and RB arrays. **d**, Distribution of genomic divergence rates between tiger-Geoffroy's cat and tiger-domestic cat across 28,312.5-kb alignment windows. Pairwise divergence values for *DXZ4* RA and RB and the internal spacer region are shown for comparison. **e**, Phylogeny of placental mammals with *DXZ4* repeat array presence (blue = RA type, yellow = RB type, gray = ambiguous) inferred from each genome assembly.

(Supplementary Fig. 17). This strategy identified a single, most probable centromere-containing interval for each chromosome enriched >1,000-fold with a small class of tandem repeats (Supplementary Fig. 18). The location of these intervals was highly conserved across species and consistent with stability of the felid karyotype. Like human and ape centromeres, several better-resolved cat centromeres (for example, chrE3; Fig. 4a) consisted of a central satellite array of higher-order repeats. The predominant satellite repeat was 113 bp in length, ~25% smaller than the 151-bp α satellite typical of great ape centromeres^{5,48} (Supplementary Fig. 19). StainedGlass analysis of these candidate satellite arrays revealed patterns of monomer divergence similar to great ape centromere arrays, with more divergent monomers flanking higher identity monomers within the central satellite array (Fig. 4a). The Geoffroy's cat possessed the largest centromeric repeat arrays

on most chromosomes (Supplementary Fig. 20). This species' karyotype also has the distinct C3 metacentric chromosome, a product of a Robertsonian chromosome fusion between chrF1 and chrF2, which occurred in the ancestor of the *Leopardus* lineage ≥ 3 MYA (refs. 9,10). StainedGlass and syntenic alignment plots (Fig. 4b and Supplementary Fig. 21) reveal that Geoffroy's cat chrC3 centromeric region retains the highest pattern and sequence similarity to the ancestral chrF1 centromeric satellite array.

Centromere sizes and repeat composition varied markedly between chromosomes and across felid species. Although we cannot exclude incomplete/collapsed sequences for some of this variation (Supplementary Figs. 22–25), the centromeric regions of three autosomes were gapless in all six felid genomes (chrs. B4, D4 and E2), likely due to reduced satellite array repeat complexity. For example, *Felis*

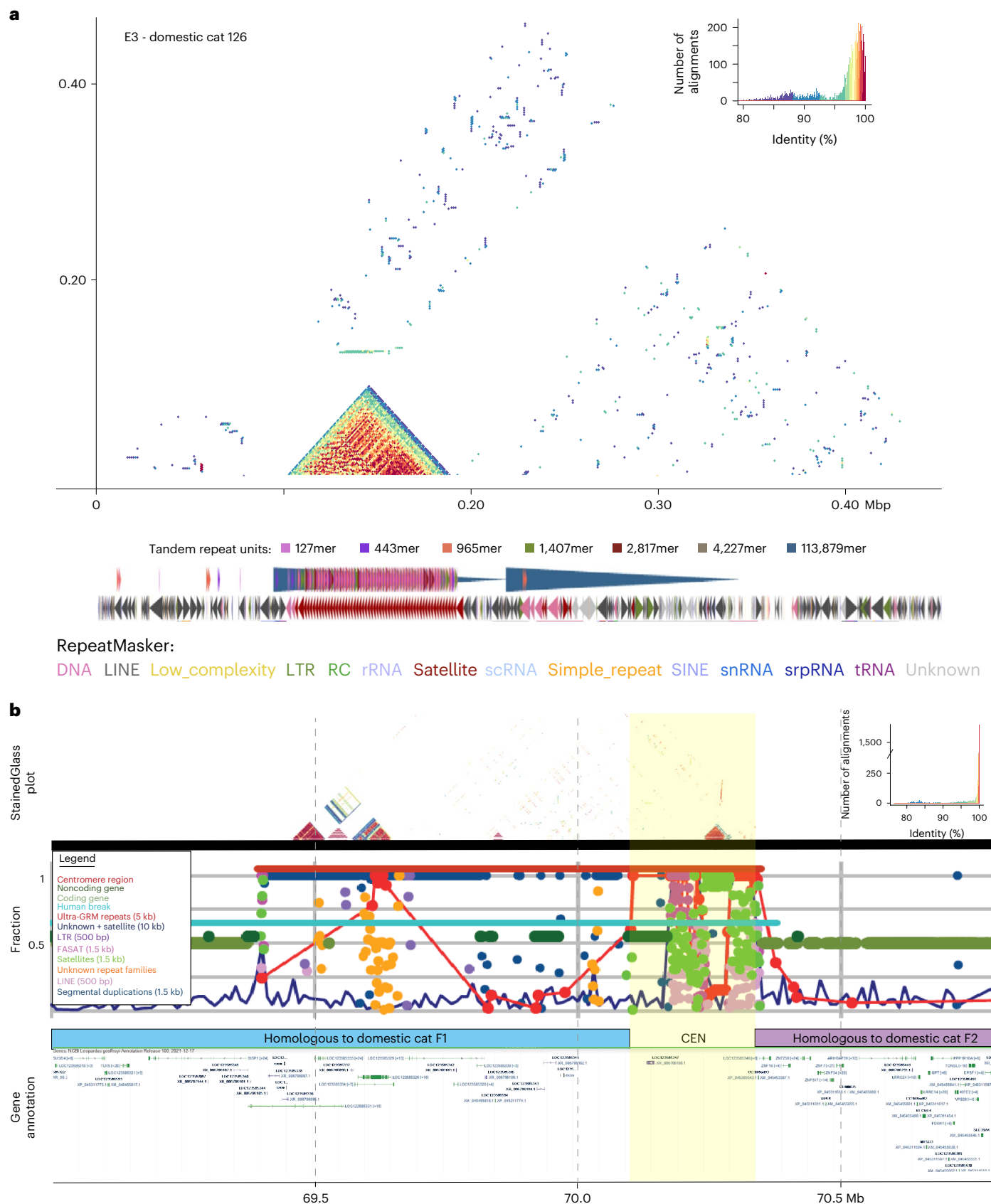


Fig. 4 | Centromere annotation and evolution. a, StainedGlass⁴¹ dot plot of domestic cat-126 chrE3 centromere region showing percent identity of self-alignments within the satellite repeat array (colored triangle, with % identity scale and distribution shown in the upper right). Below the chromosome are tracks for tandem repeat annotations (colors indicate different GRM-defined repeat units) and RepeatMasker annotations (key at bottom). **b**, Geoffroy's cat chrC3

centromere region. The lower two panels display NCBI CpG and gene annotations and inferred homology to the domestic cat F1 and F2 centromeric regions. The top tracks show StainedGlass plots and repeat annotations (and fractions observed on y axis). The most probable centromeric repeat array is highlighted in yellow and supported by alignments in Supplementary Fig. 21. CEN, centromere.

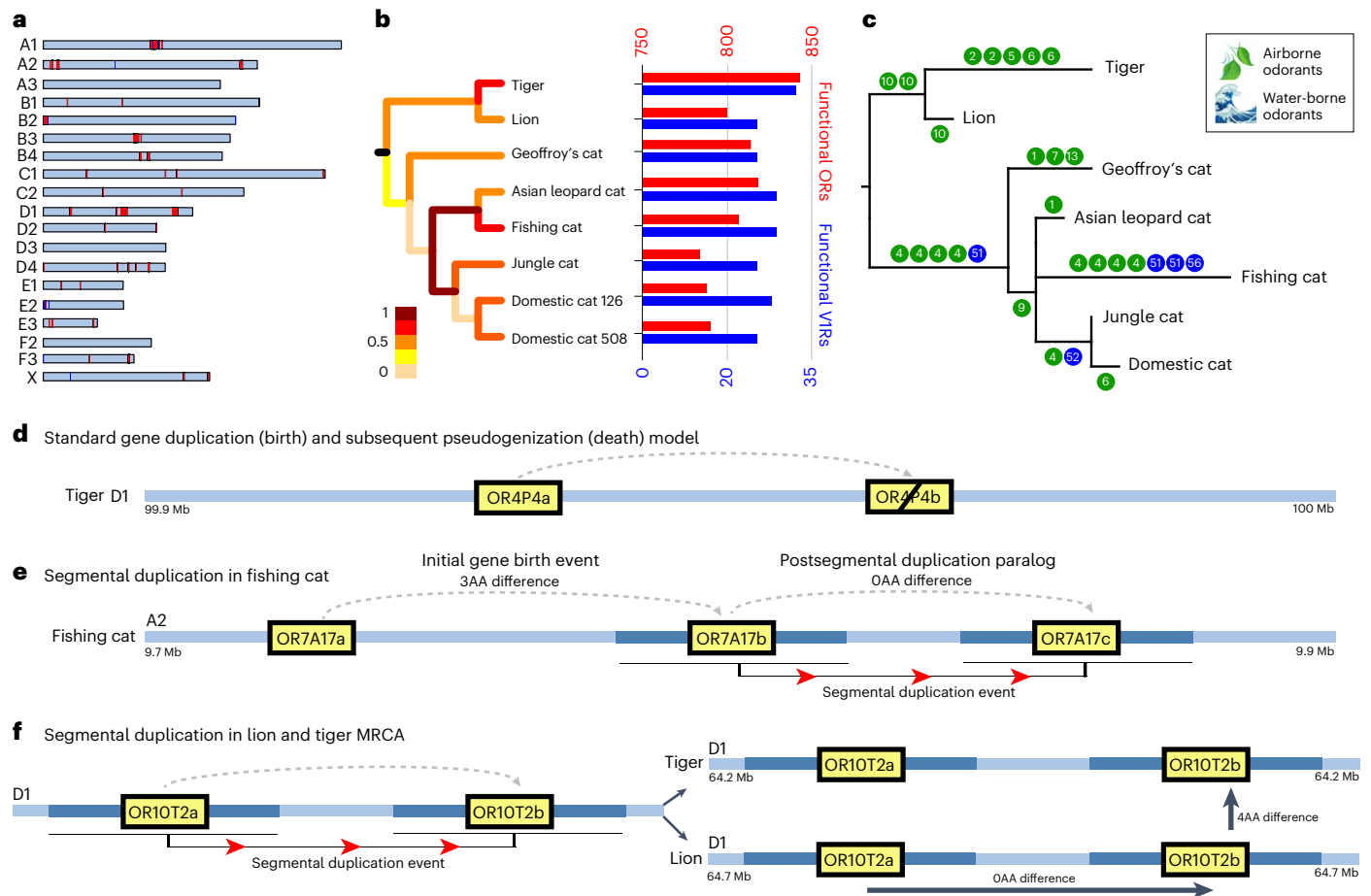


Fig. 5 | ORG and V1R gene evolution in cats. **a**, Chromosomal distribution of ORG (red) and V1R gene (blue) within the domestic cat genome. **b**, Phylogeny and rate of ORG family duplications (scale to lower left). Barplots to the right illustrate per species ORG (red) and V1R (blue) functional gene copy number. **c**, Number of per branch unique ORG retention, classified into class I (blue = ‘water-borne’) and class II (green = ‘airborne’) receptor types. Each circle represents a

uniquely retained gene, with its subfamily classification depicted by the number. **d–f**, Models of ORG birth and death with specific examples—the standard birth and death (pseudogenization) model illustrated by tiger chrD1 (OR4P4a and OR4P4b) (**d**); a gene birth followed by paralog birth via SD in the fishing cat (**e**) and a gene birth via SDs in the *Panthera* ancestor preceding speciation of the lion and tiger lineages (**f**).

chrB4 possesses a narrower centromeric interval and lacks the large satellite arrays observed on other chromosomes (Supplementary Fig. 26). Some mammalian families, such as equids (donkeys, onagers and zebras), also exhibit considerable variability in the presence/absence of satellite repeats at their centromeres^{49,50}. By contrast, the chrD4 centromere possesses a mostly conserved satellite array and illustrates the rapidity with which the central satellite monomer array sequences diverge relative to the flanking sequence (Supplementary Figs. 27 and 28), similar to great apes⁵. These new assemblies pave the way to exploring the potential role of interspecific centromeric satellite variation in felid speciation⁵¹.

Evolutionary innovations in sensory supergene families

Olfactory receptor genes (ORGs) encode receptors that detect odorants and represent the largest gene superfamily, dispersed across the majority of mammalian chromosomes⁵² (Fig. 5a). Variation in repertoire size and functional content has been linked to shifts in ecology, diet and life history traits, which are likely crucial components of adaptation to new environments^{53,54}. Most comparative studies of ORG variation were based on short-read assemblies, which confound allelic discrimination and gene copy number differences. Indeed, the previous enumeration of differences in OR repertoire sizes between cats and tigers produced opposing results^{14,54}. We quantified the functional ORG and vomeronasal receptor (V1R) gene profiles within each

genome assembly and added published repertoire reconstructions from the jungle cat (*Felis chaus*)³⁷ and a fishing cat (*Prionailurus viverrinus*) based on Hi-Fi reads⁵⁵. These assemblies showed gapless ORG and V1R gene cluster inclusion with contiguity metrics approaching the single-haplotype assemblies (mean cN50 = 80 versus 91 Mb).

We observed large ORG copy differences (>10% of the maximum repertoire size) between species (Fig. 5b and Supplementary Table 5). Felids retain >70% functional ORGs (Supplementary Table 6), larger than most mammals⁵⁴. This elevated functional repertoire may reflect their predatory behaviors, with an acute sense of smell to track and locate prey across great physical distances⁵⁶. The tiger is solitary, with among the largest home range sizes and habitat diversity of any living felid⁵⁷. It possesses the most extensive functional ORG repertoire and the highest number of gene duplications of any sampled species for airborne Class II ORGs (Fig. 5b,c and Supplementary Tables 6 and 7). Several ORGs that are known to bind volatile compounds in the blood (*ORIG1*: nonanal, *OR2W1* and *ORS1V1*: hexanal)^{58,59}, and the pheromone androstenone (*OR7D4*)⁵⁸ had relatively high copy numbers (Supplementary Fig. 29). The tiger and Geoffroy’s cat lineages both possessed specific duplications in ORGs associated with blood-associated odorants. By contrast, the ancestor of the domestic cat lineage had the fewest ORG duplication events, potentially reflecting relaxed evolutionary pressure on olfaction before or during domestication.

Class I ORG families (OR51, OR52, OR55 and OR56) are generally considered the ‘water-borne’ odorant-binding class, and selection for functional copies is usually rare in terrestrial mammals. The fishing cat (*Prionailurus viverrinus*) is one of two felids with pronounced aquatic adaptations such as foot webbing and other otter-like morphological adaptations to the head and tail⁵⁶. The fishing cat possesses one of the largest relative percentages of functional water-borne ORGs (75%), similar to the two domestic cats (74% and 76%) and higher than the other wild felids (lion: 67%, tiger: 71%, Geoffroy’s cat: 72% and leopard cat and jungle cat: 73%; Supplementary Table 8). Notably, the adaptive importance of water-borne OR receptors to the fishing cat is reflected in the lack of any class I-specific pseudogenization events within its lineage and the retention of three functional class I ORGs that have subsequently been pseudogenized in all other felid species (Fig. 5c and Supplementary Table 9).

ORG sequences evolve through an evolutionary pattern known as the birth-and-death model⁶⁰ (Fig. 5d). This model assumes new ORGs are ‘born’ through tandem gene duplication and retained via subfunctionalization or neofunctionalization⁶¹. Gene death occurs from nonsense mutations or larger-scale genic deletions. Analysis of the chromosomal regions flanking ORG clusters revealed that while many of the inferred duplication events consisted of the ORG sequence alone, 18 of 198 detected lineage-specific gene duplications (9.1%) were the product of larger SDs spanning $\geq 2,000$ bp (Fig. 5e,f), similar to the frequency (10%) of SD-driven ORG duplications in humans⁶². A mean rate of 2.73 amino acid mutations was observed between functional segmentally duplicated ORGs compared to 2.3 amino acids in gene-specific duplicates, suggesting differences in the rate of natural selection acting on ORG evolution may be dependent on the duplication mechanism. This distinction is important because all genes duplicated as part of a larger block may not be targets of selection. SD likely explains some of the more extensive ORG repertoires observed in mammals, as in the African elephant, which is estimated to possess over 2,000 functional genes but more than 1,000 pseudogenes⁶³. Future analyses of sensory genes in T2T genomes will allow further exploration of this model of ORG evolution in a range of vertebrate taxa.

VIR detects pheromones and other sociochemicals. We recovered complete VIR gene repertoires for each species, ranging from 67 genes in the jungle cat to 85 genes in the tiger (Fig. 5b), with ~36% of VIR genes retaining function across species (Supplementary Tables 10 and 12). The Tiger genome possessed the most functional VIR loci. Like their large functional ORG repertoire, this is potentially attributable to the large physical distances necessary for tigers to detect scent marks and discriminate potential conspecific and reproductively receptive mates⁶⁴. Most of the estimated gene duplication events occurred in tiger and lion genomes. They may reflect divergent adaptations to the use of social/sexual cues in both solitary and social life histories. Interestingly, we observed the highest frequency of nonfunctional (68%) VIR genes within the lion genome. Because lions live in highly cooperative groups in physical proximity, we hypothesize that the increased pseudogenization rate may be the product of relaxed selection on the use of chemical cues for determining sexual status and identifying mates relative to solitary species. Furthermore, while there were no unique lineage or species-specific retention of functional VIR genes like in the ORG family, the only unique VIR gene loss event occurred in the ancestor of the domestic cats, evidence of relaxed selective pressures during domestication¹⁴.

Discussion

Here we applied feline hybrid models to produce multiple well-annotated and near-gapless sequence assemblies spanning the felid radiation. Despite their similar evolutionary ages, great ape and felid lineages possess distinct differences in SD densities that provide a genomic explanation for the striking karyotypic stability observed across the cat radiation. Resolving recalcitrant sequence structures

also clarifies how natural selection continues to shape different axes of genomic diversity. The chemosensory system is particularly relevant in this sense, as gene family variation has large fitness effects, and here we showed that precisely resolved gene repertoires allow for discriminating the ecological relevance of gene birth and death. Notably, large differences in ORG and VIR gene repertoires between the closely related lion and tiger likely mirror the outcome of natural selection on evolved differences in social versus solitary life histories. The private retention of aquatic-borne odorant receptors in the fishing cat also helps to clarify the role of natural selection in ecological niche adaptation. Future studies of sensory gene repertoire variation within species occupying broad geographic ranges and habitats (for example, tiger, puma and bobcat) using phased assembly approaches will provide critical insights into the genetic basis of local sensory adaptation.

Speciation studies typically focus on the landscape of divergence, seeking outlier loci or ‘islands of speciation’ to uncover the genetic barriers that maintain species boundaries in the face of gene flow⁶⁵. Our study illustrates the rapidity with which functional satellite elements evolve relative to background rates of gene sequence variation and provides additional evidence as to the role of *DXZ4*’s exceptional divergence in felid speciation. Yet satellites are often invisible to divergence scans as these highly repetitive regions are typically missing^{4,37} or misassembled in most diploid genome assemblies. Future genomic prospecting from T2T genomes^{3,66} promises to lend new insights into the landscape of genomic and structural divergence in adaptive phenotypic variation. We anticipate exciting breakthroughs inferring the genetic mechanisms of speciation and enabling genomically informed biodiversity conservation^{67–69}.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information, details of author contributions and competing interests and statements of data and code availability are available at <https://doi.org/10.1038/s41588-023-01548-y>.

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Methods

Biological materials and genome sequencing

Fibroblast cell lines were established at the National Cancer Institute under protocols approved under contract N01-CO-12400. The parent-offspring trio of the Safari cat was composed of a random-bred domestic cat (*Felis silvestris catus*) dam, a Geoffroy's cat (*Leopardus geoffroyi*) sire and a female F1 offspring. Cell lines were karyotyped to confirm species identity and F1 status (Supplementary Fig. 31). The details of the Bengal cat F1 trio were previously reported^{18,73,74}. The parent-offspring trio of the liger was composed of a tiger dam, a lion sire and a male F1 offspring (LxT-3). A karyotype of the F1 male liger was generated (Supplementary Fig. 32).

High molecular weight genomic DNA was extracted from cells using a modified salting-out protocol⁷⁵. PacBio SMRT libraries were size selected (>20-kb) and sequenced on the Sequel IIe instrument to yield approximately 158× and 153× coverage for the Safari and Liger F1, respectively. The Bengal F1 reads¹⁸ were sequenced on the Sequel I platform to 90× coverage.

Illumina fragment libraries (~300-bp average insert size) were prepared for the parent samples of trios using the NEBNext Ultra II FS DNA Library Kit (New England Biolabs). Samples were sequenced to ~30× coverage with 2 × 150-bp reads on the NovaSeq 6000 platform.

Hi-C library preparation and sequencing

Fibroblasts were fixed as a monolayer using 1% formaldehyde, divided into 4.2×10^6 cell aliquots, snap-frozen in liquid nitrogen and stored at -80 °C (ref. 76). Cells were lysed, resuspended in 200 µl of 0.5× DNase I digestion buffer and chromatin digested with 1.5 units of DNase I for 4 min. Downstream library preparation was performed as described⁷⁶ and sequenced on the Illumina NovaSeq 6000 to ~78× coverage.

Genome assembly and annotation

Haplotype binning. All Illumina data were processed with FastQC v0.11.8 (ref. 77) and adapter trimming using Trim Galore! v0.6.4. Illumina sequences were unavailable for the parents of the F1 Safari cat. Therefore, we used the domestic cat parent (Fca-508) of the Bengal F1 hybrid and published Geoffroy's cat Illumina data (Oge-3: SRR6071645)¹⁰ for phasing. Long reads were phased into haplotype bins using the trio-binning feature of Canu v1.8 (TrioCanu)^{15,78}.

De novo assembly. Haplotyped long reads for each species were assembled using NextDenovo v2.2-beta.0 (github:Nextomics/Nextdenovo) with the configuration file (.cfg) altered for inputs: minimap2_options_raw = -x ava-pb, minimap2_options_cns = -x ava-pb. The seed_cutoff= option was adjusted to 32k for all assemblies. Lion Y chromosome contigs were identified using published procedures³⁷.

Contig polishing and QC. NextPolish v1.3.0 (ref. 79) and NextDenovo corrected long reads were used to polish the raw contigs. Changes to the NextPolish configuration file included: genome_size=auto, and task=best, which instructs the program to perform two iterations of polishing using the corrected long reads. The sgs option was removed as polishing with the parental diploid short reads could lead to the conversion of consensus sequence to reflect the alternate haplotypes not present in the F1. The lgs options within the configuration file were left at default settings except for modification for PacBio long reads by adjusting minimap2_options= -x map-pb. Basic assembly stats were generated using QUAST v5.0.2 (ref. 80) with the --fast run option selected. BUSCO v4.0.6 (ref. 81) was used to assess genome completeness, with the -m genome setting with -l mammalia_odb10 database selected (9,226 single copy genes). Visual assessment of the assemblies was performed through alignment to the domestic cat assembly Fcat_Pben_1.0_maternal_alt (Fca-508: GCA_016509815.1)¹⁸ using nucmer (mummer3.23 package)⁸² with default settings.

Delta files were used to generate dot plots using Dot: interactive dot plot viewer for genome-genome alignments (DNAnexus).

We also assessed assembly quality based on *k*-mer accuracy and completeness. Illumina data from each respective F1 hybrid were used to generate Meryl (v1.3) *k*-mer databases for the two parents and a child. Resulting Meryl databases were then used to generate hapmer databases using Merqury's (v1.3) hapmer script (\$sh \$MERQURY/trio/hapmers.sh). The parental hapmer databases and child database were then passed to Merqury to evaluate assembly quality. We also assessed assembly quality using Inspector (<https://github.com/Maggi-Chen/Inspector>; v1.0.2).

Scaffolding. Polished contigs from the domestic and Geoffroy's cat were scaffolded using Hi-C data generated from the F1 Safari cat hybrid fibroblasts. Hi-C reads were binned into parental haplotypes prior to scaffolding by aligning the offspring reads to both polished parental assemblies using bwa mem v0.7.17 (ref. 83) and the classify_by_alignment (https://github.com/esrice/trio_binning/; v0.2.0) program as described in ref. 84. Haplotyped reads were mapped to polished contigs using the pipeline and scripts described in ref. 84 (<https://github.com/esrice/slurm-hic/>) using SALSA v2.2 (refs. 85,86) with parameters -e none -m yes. We removed all Y chr contigs prior to scaffolding to prevent incorporation of repetitive Y chromosome contigs into paralogous autosomal regions. Previously published Hi-C data for tiger (SRR8616865) and lion (SRR10075807/SRR10075808) (DNA Zoo⁸⁷) were used to scaffold their respective assemblies with SALSA parameters -e GATC -m yes. The resulting scaffolds were inspected using QUAST, nucmer and Hi-C contact maps. RagTag v1.0.1 (ref. 88) was used to align scaffolds relative to Fcat_Pben_1.0_maternal_alt (Fca-508: GCA_016509815.1). Selected RagTag parameters included -remove-small, -f10000 and -j unplaced.txt. RagTag scaffolds were manually inspected with Hi-C maps generated using Juicer v1.5.7 (ref. 89) with option -s for compatibility with DNase Hi-C libraries. Maps were visualized using Juicebox v1.11.08 (ref. 90) and Juicebox Assembly Tools with scripts from 3d-dna v.180922.

Genome annotation. The NCBI annotation pipeline provided the final assembly annotations used in our analyses. Identification and annotation of *DXZ4* repeat units were performed manually using GC content traces, CTCF motif annotations and self-self dot plots for the region using Geneious Prime v2021.0.3 and FlexiDot v1.06 (ref. 91). CTCF motifs were annotated using the Geneious Annotate & Predict tool with a sequence motif of GAGTTTCGCTTGATGGCAGTGTGCACCACGAAT, based on the conserved CTCF motif logo⁹², with the most prevalent nucleotide representative of each position. A max mismatch of 13 was selected to allow for interspecific ambiguity within the motif. CTCF sites annotated using this method corresponded to the approximate location within human *DXZ4* repeat units originally described by Chadwick⁹³. Independent repeat units were aligned using the Mafft Multiple Aligner v1.4.0, and maximum likelihood (ML) trees were generated with RAXML v8.2.11 (ref. 94) under a GTR+I+G model of sequence evolution. Trees were pruned using Mesquite v3.61 (ref. 95) and visualized using FigTree v1.4.4. Mean within- and between-group *P* distances for masked (10% gaps masked) *DXZ4* repeat unit alignments were calculated using Mega-X v10.0.5 (ref. 96). To compare the rate of *DXZ4* repeat evolution to the remainder of the genome, we created a multiple-sequence alignment with the domestic cat genome (Fca126) and Geoffroy's cat aligned to the tiger SHA reference. The alignment was passed to Tree House Explorer (v1.0.2)⁹⁷ where the THeXBuilder function was used to calculate *P* distances in 10 kb windows with a strict missing data threshold of 0.0.

Comparative genomic analyses of *DXZ4* were assessed with contiguous long-read genome assemblies from all placental mammal superordinal clades⁹⁸ downloaded from NCBI. We chose male assemblies,

where available, due to their single chrX haplotype. Reference gene annotations for *PLS3* and *AGTR2* were used with Liftoff to identify the location of *DXZ4* (ref. 92). Centromere positions were identified using a combination of NCBI annotations, interspecific alignments and the Atlas of Mammalian Chromosomes, Second Edition⁹⁹. Dot plots were generated using FlexiDot. We determined the presence/absence of *DXZ4* based on the presence of repeat structure, CTCF-binding motifs and location relative to *PLS3* and *AGTR2*. Human, cat and pig *DXZ4* repeat monomers were also queried against the chrX using the discontinuous megablast BLAST algorithm.

Repetitive landscape, centromere annotation and analysis

Repeats. Repeats in each of the genomes were masked using RepeatMasker (RepeatMasker-4.1.2-p1; Smit, AFA, Hubley, R & Green, P. RepeatMasker Open-4.0. 2013–2015 <<http://www.repeatmasker.org>>) with the Dfam3.5+RepBase (rbrm-20181026) libraries where RepeatMasker was configured to use trf4.09.1 for identifying tandem repeats, rmbblastn (2.10.0+) to generate alignments and called with the -species cat option to mask using the cat-specific libraries. All repeats identified with that RepeatMasker run using the standard cat libraries were then masked as Ns, and RepeatModeler2 (ref. 100) (RepeatModeler 2 v2.0.2a; rmbblast 2.10.0+; TRF 4.10, RECON, RepeatScout 1.0.6, RepeatMasker 4.1.2 = -p1; LTR Structural Analysis: Enabled (GenomeTools 1.6.2, LTR_Retrieve v2.9.0, Ninja 1.10.2, MAFFT 7.453, CD-HIT 4.8.1)) was used to model additional repetitive elements with the LTRstruct option enabled (LTR_retriever v2.9.0 configured to use rmbblast2.10.+; RepeatMasker; hmmer3.3.2; cdhit4.8.1). All identified repeats were masked, RepeatModeler2 was run again and the genomes were N-masked. Finally, to be certain the centromeres had been fully sampled, centromere regions from the final N-masked genomes were used as the input to RepeatModeler to create a final set of repeat models that were added to the Dfam3.5 + RepBase + the two previous rounds of RepeatModeler. The RepeatModeler consensus sequences were extended when the full repeat was not modeled and trimmed when the repeat model ran into a neighboring exon, concatenated, and redundancy was removed.

SDs. Before identifying SDs, repetitive elements identified using the RepeatMasker/RepeatModeler approach described above, as well tandem repeats identified by GRM¹⁰¹ and ULTRA¹⁰² (version 0.99.17ultra; using period=10, period=100 and period=4000), were masked. SDs were defined using SEDEF²⁸ with default parameters.

Centromeres. Initial outer bounds for the centromere region of each chromosome were defined by aligning known bounding markers¹⁰³ against each cat genome using blat¹⁰⁴. The location was further refined by identifying human/cat synteny breakpoints by aligning each cat genome to the human genome (GCA_000001405.27_GRCh38.p12_genomic.fna) using nucmer⁸² with default parameters, and then filtered using a 70% identity filter (delta-filter -i 70). Many felid chromosome arms are painted by separate human chromosomes using Zoo-FISH data, hence synteny breaks should define centromeric regions¹⁰⁵. Reciprocal best alignments were extracted (show-coords -cT) and human/cat breakpoints were identified. To identify the centromere boundary, beginning at the human/cat alignment breakpoint, we move into the centromere analyzing the repeat density of Unknown+Satellite repeats in 25 kb windows in 1 kb steps. When that repeat density exceeded 0.3, we stepped 'back' to the base of the repeat density peak. To identify the position at which there was a significant change in the Unknown+Satellite repeat density, we identified the change point with a probability of at least 0.75 (ref. 106). From that point, we again walked 'away' from the centromere using a window size of 1.5 kb on the density of all repetitive elements that were enriched >500× within the centromere to incorporate any missed elements (density > 0.25) within 30 kb and to incorporate missed

tandem repeats (repeat unit sizes 100 to 4,000, window size = 5k; density > 0.20). Finally, we checked that any boundary was between and not within a predicted gene.

Sensory receptor annotation and analysis. To identify both OR and V1R genes, we combined both the BLAST^{107,108} and the Olfactory Receptor Assigner⁵³ algorithms into a single workflow. Initially, genomic regions containing putative sequences were identified by mapping a set of mammal-annotated ORG and V1R sequences, available via RefSeq, to each genome using blastn. A minimum of 85% sequence identity and 200 bp covered per hit were used to highlight potential sensory gene sequences and exclude nonspecific GPCR-like regions. Genomic regions for each hit were extracted with an additional 500 bp up and downstream to ensure start and stop codons were included. ORA uses a set of reference profile hidden Markov models (HMMs) to annotate ORG/V1R genes for each region extracted. Profile HMMs specific to V1Rs were generated using HMMER3 (ref. 109). ORG/V1R genes were classified as nonfunctional if they contained an in-frame stop codon or if they were less than 650 bp in length (that is, not long enough to complete the seven-transmembrane domain). Identified ORG/V1R sequences were mapped to the original RefSeq data to confirm they were definitive sensory genes. All ORG/V1R genes were mapped (blastn) between felid genomes to ensure no sequences were missing between species. ORA was used to classify all ORG and V1R genes into 13 subfamilies (OR1/OR3/OR7, OR2/OR13, OR4, OR5/OR8/OR9, OR6, OR10, OR11, OR12, OR14, OR51, OR52, OR55 and OR56) and eight subfamilies (V1R1, V1R2, V1R3, V1R4, V1R5, V1R48, V1R90 and V1R100), respectively.

ML gene trees per gene family per chromosome were inferred using IQTREE v.1.6.12(GTR+I+G)¹¹⁰ based on multiple-sequence alignments generated with Clustal Omega¹¹¹. The number of lineage-specific gene duplication events per species was estimated using Notung¹¹². Additionally, by splitting gene trees into all possible subtrees via the 'ape' package in R¹¹³, gene presence/absence per subtree was used to characterize putative one-to-one orthologs across species. Ambiguous orthologous relationships were further resolved using both genomic coordinates and blast hits. To determine if lineage-specific ORG/V1R gene duplications consisted of only the specific receptor gene or represented the duplication of a larger chromosomal region, 1,000 bp both up and downstream of each sequence was extracted and analyzed for SDs as described above.

Tiger recombination map. Publicly available short-read data for four individual *Panthera tigris jacksoni* (SRR7152390, SRR7152389, SRR7152391 and SRR715294) were trimmed, filtered and mapped to the *Panthera tigris* (P.tigris_Pti1_mat1.1) reference genome. Mapping results were evaluated and summarized using the Qualimap function bamqc¹¹⁴. Samtools¹¹⁵ was used to remove duplicate reads. Base quality score recalibration was performed using GATK^{116,117} by generating an initial reference set of SNPs from the dataset itself. Variants were then called, and all samples were jointly genotyped. Variants were filtered to remove variants in repeatmasked regions using GATK. Variants were further filtered, removing variants within 5 bp of an indel and those which did not meet the following quality criteria: -e%QUAL<30|INFO/DP<16|INFO/DP>62|QD<2|FS>60|SOR>10|ReadPosRankSum<-8|MQRankSum<-12.5|MQ<40' in bcftools (<https://github.com/samtools/bcftools>). VCFtools (https://vcftools.github.io/man_latest.html) was used to remove indels, leaving 3,067,994 biallelic SNPs for further analysis. ReLERNN v.1.0.0, a deep learning approach that uses recurrent neural networks, was used to model the genome-wide recombination rate¹¹⁸. A mutation rate of 2.2×10^{-9} (ref. 119), was used. ReLERNN was run using the simulate, train, predict and bscorrect modules with default settings. Inferred recombination rates were averaged in 2 Mb blocks in 50 kb sliding windows.

Structural variant/inversion identification and analysis

Initial inversion call set detection with PAV. An initial variant call set was generated using PAV¹²⁰ (GitHub commit: 24efbea) with minimap2 (v2.24)¹²¹ parameters ‘-x asm20 --secondary=no -a -t {params.cpu} --eqx -Y -B 2 -z 10000,50 --end-bonus=100’ and PAV configuration settings ‘inv_region_limit: 3000000’, henceforth referred to as the PAV-mm2 call set. The ‘sv_inv.bed.gz’ bed files containing inversion calls for each sample were then used for downstream filtration and validation. As an additional line of validation, we also ran PAV using Long-Read Aligner (LRA) (v1.3.2)¹²² with parameters ‘-CONTIG -p s -t’. The resulting ‘sv_inv.bed.gz’ inversion bed file was used for validation of the PAV-mm2 initial call set. Inversions overlapping regions identified as collapsed SDs identified by SDA¹²³ were removed from the analysis.

PBSV. CLR reads were mapped to Geoffroy’s cat reference assembly (O.geoffroyi_Oge1_pat1.0) using pbmm2 (v1.9.0) using the parameters ‘--sort --median-filter’. The variant call set was generated using PBSV (v2.8.0) by first identifying signatures of structural variants using the discover command ‘pbsv discover --tandem-repeats tandem_repeats.bed <input.bam> <output.svsig.gz>’, where tandem repeats were identified by GRM and ULTRA. Then, variants are called using the call command ‘pbsv call <reference.fasta> <output.svsig.gz> <output.vcf>’.

Sniffles. CLR reads were mapped to Geoffroy’s cat reference assembly (O.geoffroyi_Oge1_pat1.0) using pbmm2 (v1.9.0) using the parameters ‘--sort --median-filter’. Variants were then called using Sniffles (v2.0.7)^{124,125} with parameters ‘-t <cpu_count> -i <input.bam> -v <output.sniffles.vcf> --tandem-repeats <reference-repeats.bed>’.

Long-read mapping-based call set filtration. Call sets from PAV-LRA, PBSV and Sniffles were used to filter the initial PAV-mm2 call set by removing variants that were not supported by at least one of the three additional variant call sets. We utilized BEDTools (v2.30.0)¹²⁶ to call inversion variants with a 50% reciprocal overlap. Inversions identified on unplaced scaffolds were excluded. We identified large inversions (>500 kbp) not called by PAV with Saffire (<https://github.com/mrvollger/Saffire>). Input paf files were generated by mapping each assembly to the Geoffroy’s cat reference assembly (O.geoffroyi_Oge1_pat1.0) with minimap2 (v2.24) with parameters ‘-x asm20 -t <cpu_count> -c --eqx’ and then rustybam (<https://github.com/mrvollger/rustybam> - bioconda v0.1.31) parameters ‘rb trim-paf sample.paf | rb break-paf --max-size 5000 | rb orient | rb filter --paired-len 100000 | rb stats --paf > sample.Saffire.bed’. Inversions greater than 500 kbp were called if supported by both Saffire- and Nucmer-based⁸² dot plots.

Short-read genotyping and inversion classification. Pangenie (v1.0.1)¹²⁷ classified inversions as species/lineage-specific, paraphyletic with breakpoint use or polymorphic. Paired-end Illumina datasets for the lion ($n = 14$), tiger ($n = 14$), domestic cat ($n = 10$) and Asian leopard cat ($n = 10$) were downloaded from NCBI’s SRA database and interleaved utilizing Seqkit’s (v0.16.0)¹²⁸ concat function. The interleaved FASTQ files and fully-phased VCF files were then passed to Pangenie using the parameters ‘-u -s <sample_name> -o <sample_name> -i <sample_interleaved_fastq> -r <reference_assembly> -v <fully_phased_PAV_inversions.vcf>’. We could not genotype Geoffroy’s cat-specific inversions using Illumina data. They were called if supported by inverted alignments to all query species. An initial phylogenetic matrix was constructed by merging inversions across all samples based on 50% reciprocal overlap (calculated by pybedtools v0.9.0)^{126,129}.

Annotation of SV-overlapping/containing SDs, gaps, genes and repetitive elements. Further, pybedtools (v0.9.0) intersected the breakpoint positions of the inversions with the coordinates of SDs,

gaps, genes and repetitive elements. SciPy’s (v1.7.3)¹³⁰ rank sum function (one-sided, greater) determined if inversions flanked by SDs were significantly larger than inversions not flanked by SDs. Inversions flanked by repetitive elements sharing more than 90% identity were identified using pandas (v1.4.0). Repetitive elements within 100 kb of the inversion breakpoints were aligned using biopython’s (v1.79)¹³¹ pairwise2.align.globalmx (upstream_seq, downstream_seq, 1, 0, score_only=True).

Statistics and reproducibility. The one-sided Wilcoxon rank sum test was used to determine differences in inversion sizes between the autosomes and chrXs. In this study, no statistical method was used to predetermine sample size, no data were excluded from the analyses and the experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

Assemblies are available in NCBI under accession numbers [GCA_016509475.2](https://ncbi.nlm.nih.gov/assembly/GCA_016509475.2), [GCA_016509815.2](https://ncbi.nlm.nih.gov/assembly/GCA_016509815.2), [GCA_018350155.1](https://ncbi.nlm.nih.gov/assembly/GCA_018350155.1), [GCA_018350175.1](https://ncbi.nlm.nih.gov/assembly/GCA_018350175.1), [GCA_018350195.2](https://ncbi.nlm.nih.gov/assembly/GCA_018350195.2) and [GCA_018350215.1](https://ncbi.nlm.nih.gov/assembly/GCA_018350215.1). OR gene sequences and *DXZ4* alignments are found at: <https://figshare.com/s/68266360874d5078bdf5>.

Code availability

Publicly available software and packages were used in this study. No custom code was used. All software and packages used in this study are described within Methods section.

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Acknowledgements

We thank the High-Performance Research Computing Center at Texas A&M University for support. DNA library preparation and long-read sequencing were performed at the University of Maryland Institute for Genome Sciences (L. Tallon and L. Sadzewicz). Illumina sequencing was performed through the Texas A&M Institute for Genome Sciences & Society research core (A. Hillhouse). We thank R. Stanyon for the flow-sorted domestic cat chromosomes for FISH experiments. This research was supported by grants from the Morris Animal Foundation (grant D19FE-04 to W.J.M. and W.C.W.), the National Science Foundation (grants DEB-1753760 and DEB-2150664 to W.J.M.) and the National Institutes of Health (NIH; grant R01 GM59290 to M.A.B.). A.J.H. was funded, in part, by a training grant from the National Institute of General Medical Sciences, NIH (T32 GM135115). The funders had no role in study design, data collection and analysis, decision to publish or preparation of the manuscript.

Author contributions

W.J.M. and W.C.W. were responsible for conceptualizing the project. K.R.B., L.H., A.J.H., G.H., N.M.F., C.L., R.C., J.M.S., E.R., B.W.D., T.R., L.A.L. and S.J.O. developed the methodology. K.R.B., L.H., A.J.H., N.M.F., G.H.

and T.R. were involved in data visualization. Funding for the project was acquired by W.J.M. and W.C.W. W.J.M. and W.C.W. were responsible for project administration. Supervision of the project was provided by W.J.M., W.C.W. and M.A.B. The original draft of the manuscript was prepared by W.J.M., K.R.B., L.H., A.J.H., G.H., N.M.F. and W.C.W. All authors contributed to the investigation phase of the study and participated in the review and editing of the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41588-023-01548-y>.

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Peer review information *Nature Genetics* thanks Michael Hiller and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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Data collection Genome sequence data was generated on Pacific Biosciences Sequel IIE and Illumina NovaSeq 6000 platforms using standard commercial software.

Data analysis We used publicly available software/code for data analysis. We used the following programs and versions: FastQC v0.11.8, Trim Galore! v0.6.4, Canu v1.8, NextDenovo v2.2-beta.0, NextPolish v1.3.0, QUAST v5.0.2, BUSCO v4.0.6, mummer3.23, Merqury v1.3, Inspector v1.0.2, bwa mem v0.7.17, Samtools v1.9, BEDTools suite v2.30.0, R v.3.5.1, BLAST v2.9.0, RepeatMasker v4.0.9, seqTK subseq v1.3, SALSA v2.2, RagTag v1.0.1, Juicer v1.5.7, Juicebox v1.11.08, 3d-dna v.180922, Liftoff v1.4.2, Geneious Prime v2021.0.3, FlexiDot v1.06, Mafft Multiple Aligner v1.4.0, RAXML v8.2.11, Mesquite v3.61, FigTree v1.4.4, Mega-X v10.0.5, Tree House Explorer v1.0.2, RepeatMasker-4.1.2-p1, RepeatModeler 2 v2.0.2a, GenomeTools 1.6.2, LTR_Retrieve v2.9.0, Ninja 1.10.2, MAFFT 7.453, CD-HIT 4.8.1, hmmer3.3.2; cdhit4.8.1, ULTRA version 0.99.17, GRM, SEDEF, IQ-TREE v1.6.12, ReLERNNv1.0.0, PAV, minimap2 v2.24, Long-Read Aligner v1.3.2, PBSV v2.8.0, pbmm2 v1.9.0, Sniffles v2.0.7, SDA, SafFire, Pangenie v1.0.1, pybedtools v0.9.0, SciPy v1.7.3, pandas v1.4.0, biopython v1.79

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

All genome assemblies are available in NCBI, under the following accession numbers: GCA_016509475.2, GCA_016509815.2, GCA_018350155.1, GCA_018350175.1, GCA_018350195.2, GCA_018350215.1. OR gene sequences are found at: <https://figshare.com/s/68266360874d5078bdf5>

Research involving human participants, their data, or biological material

Policy information about studies with [human participants or human data](#). See also policy information about [sex, gender \(identity/presentation\), and sexual orientation](#) and [race, ethnicity and racism](#).

Reporting on sex and gender	N/A
Reporting on race, ethnicity, or other socially relevant groupings	N/A
Population characteristics	N/A
Recruitment	N/A
Ethics oversight	N/A

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	No statistical method was used to predetermine sample size
Data exclusions	No data was excluded in our analyses.
Replication	All attempts to replicate results were successful
Randomization	The experiments were not randomized
Blinding	The Investigators were not blinded to allocation during experiments and outcome assessment.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

Methods

- n/a | Involved in the study
- Antibodies
- Eukaryotic cell lines
- Palaeontology and archaeology
- Animals and other organisms
- Clinical data
- Dual use research of concern
- Plants

- n/a | Involved in the study
- ChIP-seq
- Flow cytometry
- MRI-based neuroimaging

Eukaryotic cell lines

Policy information about [cell lines and Sex and Gender in Research](#)

Cell line source(s)	The single haplotype genome assemblies were derived from DNA sequencing of primary skin fibroblast cell lines.
Authentication	All cell lines were karyotyped using standard methodology from early passages (p1-p3) and verified by comparison to known karyotypes and analyzed with PacBio long read technology to confirm species id.
Mycoplasma contamination	The cells were not tested for mycoplasma contamination
Commonly misidentified lines (See ICLAC register)	N/A