

## Chromosome-length genome assembly and structural variations of the primal Basenji dog (*Canis lupus familiaris*) genome

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## Abstract

**Background:** Basenjis are considered an ancient dog breed of central African origins that still live and hunt with tribesmen in the African Congo. Nicknamed the barkless dog, Basenjis possess unique phylogeny, geographical origins and traits make understanding their genome structure relative to more modern dog breeds of great interest. Here, we report the *de novo* assemblies of two Basenji: a female, China, and a male, Wags. We conduct pairwise comparisons and report structural variations between assembled genomes of three dog breeds: Basenji (CanFam\_Bas), Boxer (CanFam3.1) and German Shepherd Dog (GSD) (CanFam\_GSD). We then align representative whole genome sequences from 58 dog breeds and show the importance of genome reference when assessing variation among dog breeds.

**Results:** Here we present two high quality Basenji genome assemblies, CanFam\_Bas (China) and Wags. CanFam\_Bas is superior to CanFam v3.1 in terms of genome contiguity and comparable overall to the high quality CanFam\_GSD assembly. The increasing number of available canid reference genomes allows us to examine the impact the choice of reference genome makes with regard to reference genome quality and breed relatedness. By aligning short read data from 58 representative dog breeds to three reference genomes, we demonstrate how the choice of reference genome significantly impacts both read mapping and variant detection. Further, we generate a conservative list of structural variant calls using a consensus of both Pacific Bioscience and Oxford Nanopore long reads to identify large structural breed differences. Collectively this work highlights the importance the choice of reference genome makes in canid variation studies.

**Conclusions:** The growing number of high-quality canid reference genomes means the choice of reference genome is an increasingly critical decision in subsequent canid variant analyses. The basal position of the Basenji makes it suitable for variant analysis for targeted applications of specific dog breeds. However, as is increasingly being employed in other

model organisms, we believe more comprehensive analyses across the entire family of canids is more suited to a pangenome approach.

**Keywords:** canine genome, domestication, comparative genomics, artificial selection



## Background

85 Dogs were the first animals to be domesticated by humans some 30,000 years ago [1] and exhibit exceptional levels of breed variation as a result of extensive artificial trait selection [2]. It is not clear whether dogs were domesticated once or several times, though the weight of accumulating evidence suggests multiple events [3-9]. By establishing genome resources for more ancient breeds of dog, we can explore genetic adaptations perhaps unique to the  
90 modern dog breeds. Basenjis are an ancient breed that sits at the base of the currently accepted dog phylogeny [10]. Basenji-like dogs are depicted in drawings and models dating back to the Twelfth Dynasty of Egypt [11] and they share many unique traits with pariah dog types. Like dingoes and New Guinea Singing dogs (NGSD), Basenji's come oestrus annually—as compared to most other dog breeds, which have two or more breeding seasons  
95 every year. Basenji's, dingoes and NGSDs are prone to howls, yodels, and other vocalizations over the characteristic bark of modern dog breeds. One explanation for the unusual vocalisation of the Basenji is that the larynx is flattened [12]. The shape of the dingo and NGSD larynx is not reported.

Basenjis were originally indigenous to central Africa, wherever there was tropical  
100 forest. Primarily, what is now the DRC Congo, Southern Sudan, Central African Republic and the small countries on the central Atlantic coast. Today their territory has shrunk to the more remote parts of central Africa. The Basenji probably made its debut in the western world in around 1843. In a painting of three dogs belonging to Queen Victoria and Prince Albert entitled "Esquimaux, Niger and Neptune". Niger is clearly a Basenji. In total, 71  
105 Basenji's have been exported from Africa and, to date, ~ 56 have been incorporated into the registered Basenji breeding population.

The first dog genome to be sequenced was of Tasha the Boxer [13], which was a tremendous advance and continues to be the resource guiding canine genomics research

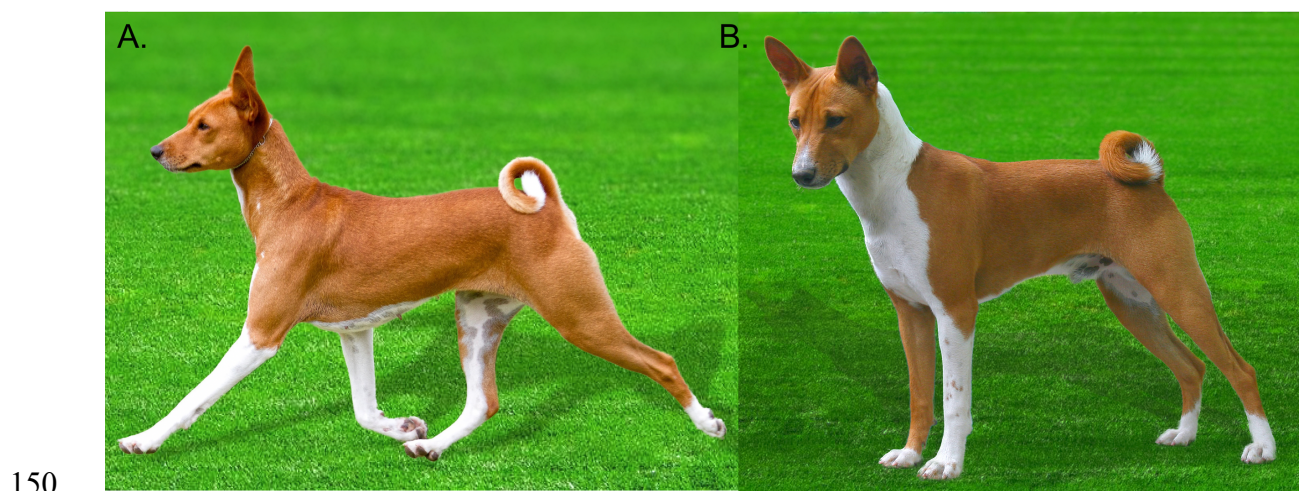
today. The Boxer is a highly derived brachycephalic breed that has been subject to  
110 generations of artificial selection. Further, due to its discontinuous sequence representation it  
has been difficult to accurately detect structural variations (SV's) in other domestic dog  
breeds. Now, a new generation of breed-specific chromosome-level genome reference  
assemblies are becoming available (5 breeds in October 2020 according to the NCBI  
assembly archive). For example, we previously published a chromosome-level German  
115 Shepherd dog (GSD) genome assembly (CanFam\_GSD) that is comprised of only 410  
scaffolds and 716 contigs [14].

Here, we first report the sequence and *de novo* assembly of two Basenji genomes,  
female and male. We then compare these assemblies with the Boxer (CanFam3.1) and GSD  
(CanFam\_GSD) [13, 14]. We conduct pairwise comparisons and report single-nucleotide  
120 variants (SNVs) and structural variants (SV) between Basenji, Boxer [13] and GSD [14]. We  
distinguish an SNV as a variation in a single nucleotide without any limitations on its  
frequency. SV comprises a major portion of genetic diversity and its biological impact is  
highly variable. Chen et al. [15] used high-resolution array comparative genome  
hybridization to create the first map of DNA copy number variation (CNV) in dogs. Many  
125 canine CNVs were shown to effect protein coding genes, including disease and candidate  
disease genes, and are thus likely to be functional. In this study, we find all types of genetic  
variation are impacted by the choice of reference genome. The basal position of the Basenji  
makes it useful as a general reference for variant analysis, but the generation of clade-specific  
genomes is likely to be important for canine nutrition and disease studies. We recommend a  
130 pan-genome approach for comprehensive analyses of canid variation.

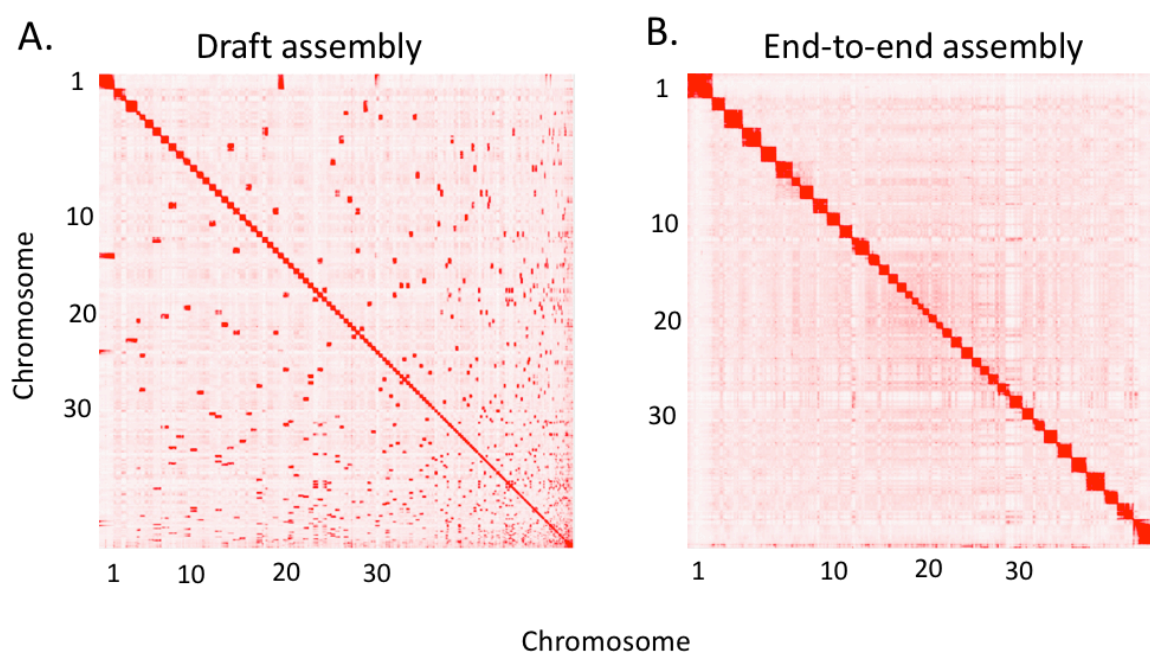
## Results

### Basenji female assembly, CanFam\_Bas

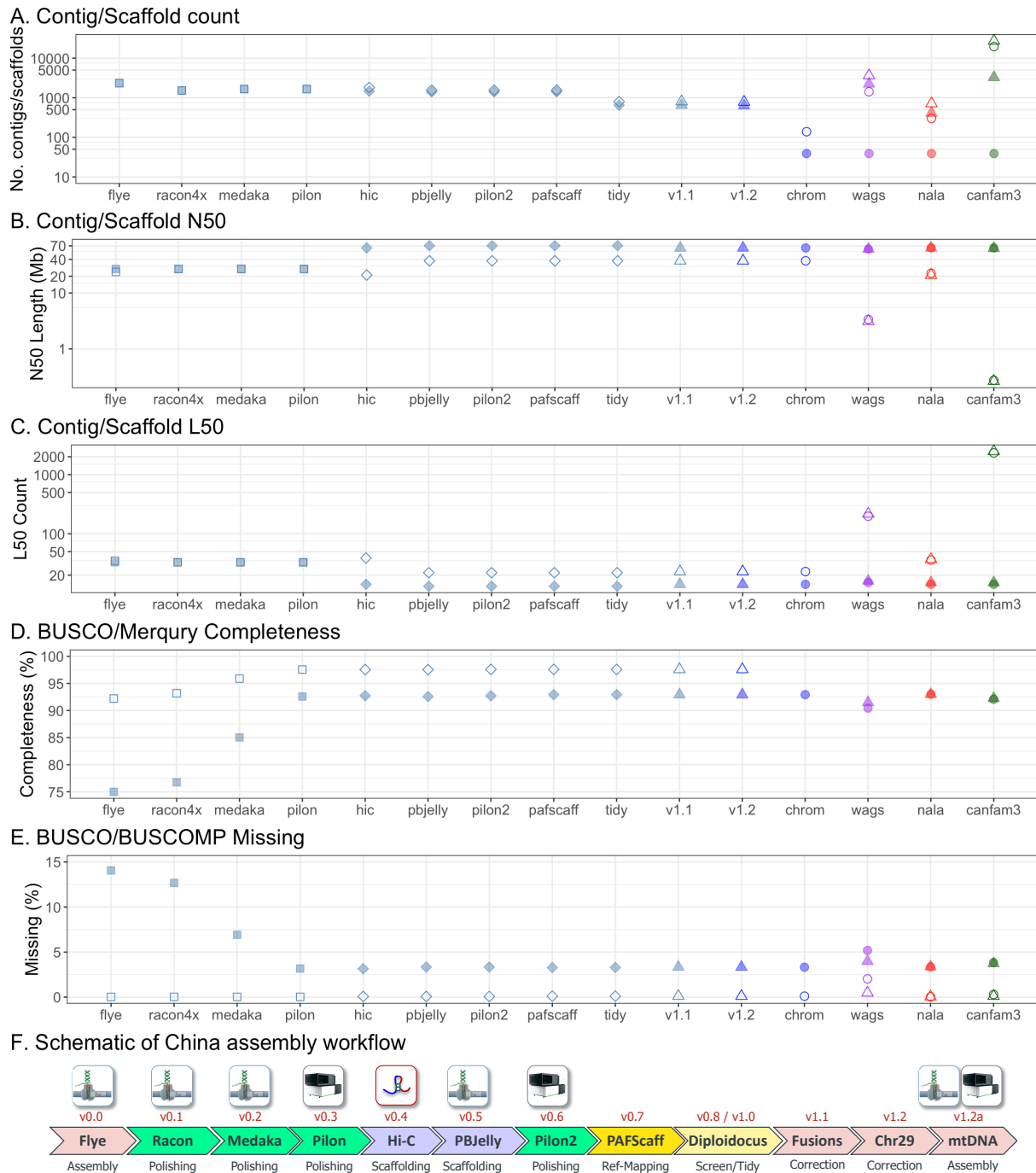
The female Basenji, China (Fig. 1A), was initially assembled from 84.5 Gb Oxford Nanopore Technologies (ONT) PromethION reads (approx. 35X depth based on a 2.41 Gb genome size) using Flye (v2.6) [16, 17] and subjected to long read polishing with Racon v1.3.3 [18] and Medaka 1.03 [19] (Supplementary Fig 1A). Additional short read Pilon [20] error-correction was performed with 115.1 Gb (approx. 47.7X) BGISEQ data. Hi-C proximity ligation was used with the DNA zoo pipeline [21-23] to scaffold 1,657 contigs into 1,456 scaffolds, increasing the N50 from 26.3 Mb to 63.1 Mb and decreasing the L50 from 33 to 14 (Figs 2 and 3, Supplementary Table 1). Scaffolds were gap-filled by PBJelly (pbsuite v.15.8.24) [24] using the ONT data, reducing the number of gaps from 348 to 148 and the number of scaffolds to 1,407. Following a final round of Pilon [20] BGISEQ-polishing, scaffolds were mapped onto the CanFam3.1 [13] using PAFScaff v0.40 [25]. Diploidocus vector filtering removed one 5.7kb contig and masked a 3.3kb region of Chromosome X as lambda phage (J02459.1) contamination. Seven rounds of iterative Diploidocus tidying of the remaining sequences removed 277 (832 kb) as low coverage/quality and 481 (1.58 Mb) as probable haplotigs, retaining 483 core scaffolds and 165 probable repeat-heavy sequences [14] as China v1.0 (Supplementary Fig 2, Supplementary Table 2).



155 **Figure 1. The Basenji dogs included in the study.** **A.** China. Is registered as Australian  
Kennel Club Supreme Champion Zanzipow Bowies China Girl. Her registration is  
#2100018945. She was born in 2016 and she is free of all known genetic diseases. Her sire  
and dam are Australian bred and her most recent ancestor from Africa was 18 generations  
160 ago. **B.** Wags. is registered as American Kennel Club Champion Kibushi the Oracle, born in  
in 2008. His registration number is HP345321/01. His sire is an American bred dog while his  
dam was imported from the Haut-Ule district of the DRC congo, 3°24'04.0"N 27°19'04.6"E,  
in 2006 .



165 **Figure 2. Contact matrices generated by aligning the CanFam\_Bas (China) Hi-C data  
set to the genome assembly A. before the Hi-C upgrade (draft assembly). B. After Hi-C  
scaffolding (End-to-end assembly).**



**Figure 3. Key contiguity, quality and completeness metrics for different assembly stages and comparison dog genomes.** Square, pre-scaffolded China; Diamond, scaffolded China; Triangle, complete assembly; Circle, main chromosome scaffolds only; Blue, China; Purple, Wags; Red, CanFam\_GSD; Green, CanFam3.1. **A.** Contig (open) and scaffold (filled) numbers. **B.** Contig (open) and scaffold (filled) N50. **C.** Contig (open) and scaffold (filled) L50. **D.** Genome completeness estimated by BUSCO v3 (filled) and Merquy (open). **E.** The percentage of missing BUSCO genes (filled) and BUSCOMP genes (those found to be Complete in any assembly) (open). **F.** Schematic of China assembly workflow. CanFam\_Bas is China v1.2.



**Genome assembly correction.** Two pairs of fused chromosomes in China v1.0 were incorrectly joined by PBJelly. Pre-gap-filled HiC scaffolds were mapped onto the assembly using Minimap2 v2.17 [26] and parsed with GABLAM v2.30.5 [27] to identify the gap  
180 corresponding to the fusion regions. These were manually separated into four individual chromosomes, gap lengths standardised, and scaffolds re-mapped onto CanFam v3.1 using PAFScaff v0.4.0. D-GENIES [28]. Analysis against CanFam\_GSD chromosomes confirmed that PBJelly had incorrectly fused two pairs of chromosomes: chromosomes 8 with 13, and  
185 chromosome 18 with 30. These were manually separated and the assembly re-mapped onto CanFam 3.1 as China v1.1. PAFScaff assigned 112 scaffolds to chromosomes, including 39 nuclear chromosome-length scaffolds.

It was observed that the mitochondrial chromosome was missing and China v1.1 Chromosome 29 contained a 33.2 kb region consisting of almost two complete copies of the  
190 mitochondrial genome that were not found in other dog genome assemblies. The 26 ONT reads that mapped onto both flanking regions were reassembled with Flye v.2.7.1 [16, 17] into a 77.2 kb chromosome fragment, which was polished with Racon v1.3.3 [18] and Medaka 1.03 [19]. This was mapped back on to the Chromosome 29 scaffold with GABLAM v2.30.5 [27] (blast+ v2.9.0 megablast [29]) and the mitochondrial artefact replaced with the  
195 corrected nuclear mitochondrial DNA (NUMT) sequence. Finally, scaffolds under 1kb were removed to produce the China v1.2 nuclear genome.

**Mitochondrial genome assembly.** In total, 4,740 ONT reads (52.1 Mb) mapping on to mtDNA were extracted. To avoid NUMT contaminants, a subset of 80 reads (1.32 Mb) with 99%+ mtDNA assignment and 99%+ mtDNA coverage, ranging in size from 16,197 kb to  
200 17,567 kb, were assembled with Flye 2.7b-b1526 [16, 17] (genome size 16.7 kb) into a 33,349 bp circular contig consisting of two mtDNA copies. This contig was polished with Racon [18] and Medaka [19], before being re-circularised to a single-copy matching the

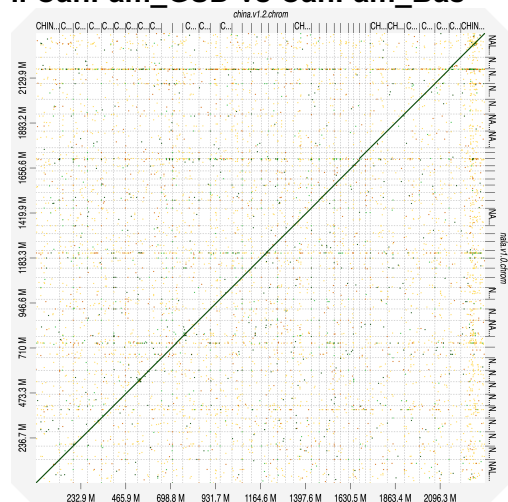
CanFam3.1 mtDNA start position. After final Pilon [20] correction of SNPs and indels, the 16,761 bp mitochondrial genome was added to the China v1.2 nuclear assembly.

205 ***CanFam\_Bas (China) reference genome.*** The resulting chromosome-length CanFam\_Bas reference genome is 2,345,002,994 bp on 632 scaffolds with 149 gaps (76,431 bp gaps) (Table 1). The 39 nuclear plus mitochondrial chromosome scaffolds account for 99.3% of the assembly and show a high level of synteny with CanFam3.1 and CanFam\_GSD (Fig 4). CanFam\_Bas represents the most contiguous dog chromosomes to date, with a contig N50 of 210 37.8 Mb and contig L50 of 23, which is slight improvement over CanFam\_GSD and considerably more contiguous than the standard dog reference genome, CanFam3.1 (Fig 3, Table 1). The completeness and accuracy of the genome as measured by BUSCO v3 [30] (laurasiatherian,  $n=6253$ ) is also superior to CanFam3.1 and approaches that of CanFam\_GSD (92.9% Complete, 3.75% Fragmented, 3.34% Missing).

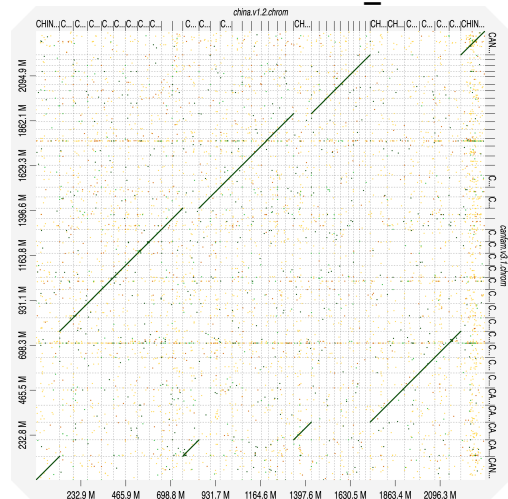
#### 215 **Methylomic identification of putative regulatory elements**

Additionally, we profiled whole genome methylation of Basenji's blood DNA using MethylC-seq [31]. The number of unmethylated (un) highly methylated (high) CpG sites in Basenji's genome were similar to that of GSD (Supplementary Fig. 3A). Importantly, high resolution DNA methylation data can be utilised to identify the putative regulatory elements 220 in a given tissue type. That is, CpG-rich unmethylated regions (UMRs) mostly correspond to gene promoters, while CpG-poor low-methylated regions (LMRs) correspond to distal regulatory elements, both characterised by DNase I hypersensitivity [32] Using MethylSeekR algorithm [33], we performed the segmentation of Basenji DNA methylome and discovered 20,660 UMRs and 54,807 LMRs (Supplementary Fig. 3B,C), in line with 225 previously reported numbers in mammalian genomes [14, 33, 34].

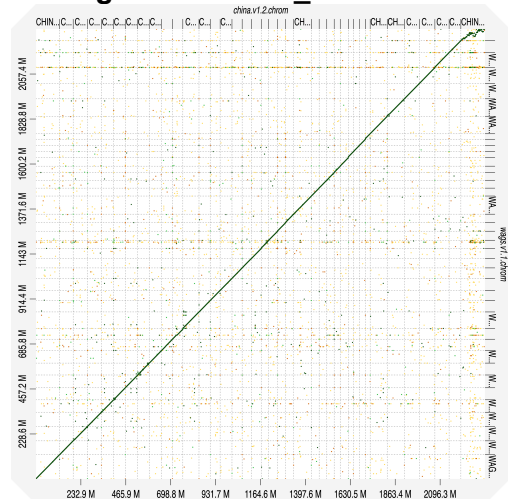
**A. Main nuclear chromosomes scaffolds**  
**i. CanFam\_GSD vs CanFam\_Bas**



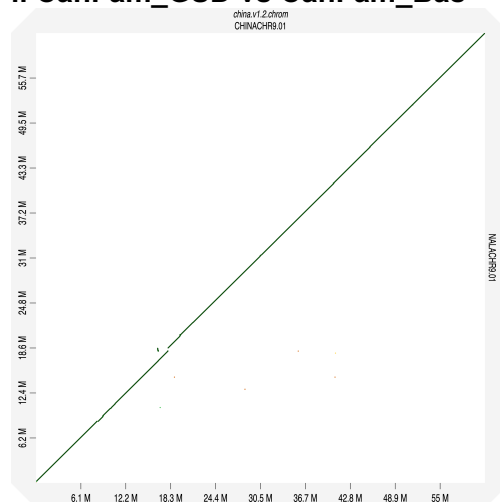
**ii. CanFam3.1 vs CanFam\_Bas**



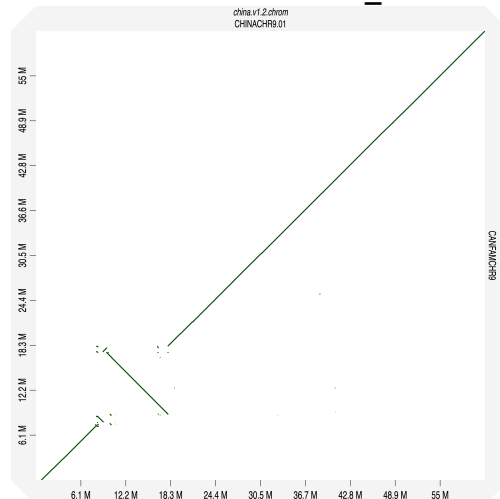
**iii. Wags vs CanFam\_Bas**



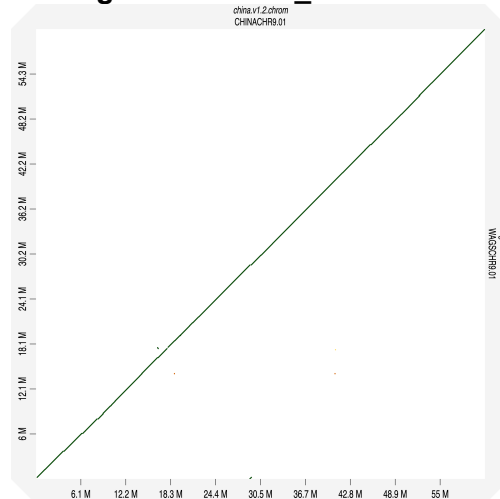
**B. Main chromosome 9 scaffold**  
**i. CanFam\_GSD vs CanFam\_Bas**



**ii. CanFam3.1 vs CanFam\_Bas**



**iii. Wags vs CanFam\_Bas**



**Figure 4. D-GENIES synteny plots of main chromosome scaffolds for three dog genome assemblies against CanFam\_Bas.** In each case, CanFam\_Bas (China v1.2) is on the x-axis and the comparison assembly on the y-axis. Gridlines demarcate scaffolds. Thick black lines indicate regions of genomic alignment. **A.** All-by-all main chromosome scaffold alignments with **(i)** CanFam\_GSD, **(ii)** CanFam\_3.1, and **(iii)** Wags. **B.** Main chromosome 9 scaffold alignment with **(i)** CanFam\_GSD, **(ii)** CanFam\_3.1, and **(iii)** Wags.



**Table 1. Genome assembly and annotation statistics for Basenji assemblies vs CanFam3.1 and CanFam\_GSD.**

	CanFam_Bas (China)	Wags	CanFam3.1	CanFam_GSD
Total sequence length	2,345,002,994	2,410,429,933	2,410,976,875	2,407,308,279
Total ungapped length	2,344,926,563	2,410,291,233	2,392,715,236	2,401,163,822
Number of scaffolds	632	2,243	3,310	430
Scaffold N50	64,291,023	61,087,166	45,876,610	64,346,267
Scaffold L50	14	16	20	15
Number of contigs	780	3,630	27,106	736
Contig N50	37,759,230	3,131,423	267,478	20,914,347
Contig L50	23	217	2,436	37
No. chromosomes	40	40	40	40
Percentage genome in main chromosomes	99.3%	94.8%	98.3%	96.5%
BUSCO complete (genome)	92.9% (1.14% Duplicated)	91.5% (1.31% Duplicated)	92.2% (1.17% Duplicated)	93.0% (1.38% Duplicated)
BUSCO fragmented (genome)	3.74%	4.53%	4.03%	3.73%
BUSCO missing (genome)	3.34%	3.98%	3.73%	3.37%
BUSCO complete (proteome)	98.5% (1.9% Duplicated)	97.8% (2.4% Duplicated)	95.1% (1.0% Duplicated)	98.9% (2.4% Duplicated)
BUSCO fragmented (proteome)	1.2%	1.5%	1.9%	1.0%
BUSCO missing (proteome)	0.3%	0.7%	3.0%	0.1%

### 235 Male Basenji assembly, Wags

For the male Basenji, Wags, (Fig 1B) we generated Pacific Bioscience Single Molecule Real Time (SMRT) sequences to approximately 45X genome coverage and assembled the genome to ungapped size of 2.41 Gb (Supplementary Fig 1B). Assembly contiguity metrics of 3,630 total contigs show N50 contig and scaffold lengths of 3.1 and 61 Mb length, respectively (Table 1). Wags alignment to China revealed a high level of synteny. However, the Wags assembly of the X chromosome is smaller in size (59 Mb vs 125 Mb) and shows multiple rearrangements as a result of lower sequence coverage on the sex chromosomes (~21x).

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Therefore, all comparative analyses reported herein were done with China, the female Basenji, CanFam\_Bas. In addition, the Wags assembly includes 3.6 Mb of the Basenji dog Y  
245 for future comparative studies of this unique chromosome.

### Genome annotation

The CanFam\_Bas and Wags assemblies were annotated using the homology-based gene prediction program GeMoMa v1.6.2beta [35] and nine reference species [14]. In total, CanFam\_Bas (China) and Wags had similar numbers of predicted protein-coding genes at  
250 27,129 (68,251 transcripts) and 27,783 (65,511) transcripts, respectively (Supplementary Table 3). Analysing the longest protein isoform per gene with BUSCO v3 [30] (laurasiatherian,  $n=6253$ , proteins mode), CanFam\_Bas was measured to be 98.5 % complete (1.9% duplicated) and Wags was measured as 97.8% complete (2.4% duplicated). Both proteomes compare favourably with CanFam3.1 in terms of completeness (Table 1). To  
255 correct for differences introduced by the annotation method, CanFam3.1 was annotated with the same GeMoMa pipeline. Approximately 90% of the Quest For Orthologues (QFO) reference dog proteome [36] is covered by each GeMoMa proteome, confirming comparable levels of completeness (Supplementary Table 3).

When the CanFam\_Bas GeMoMa proteome was compared to Wags, CanFam3.1  
260 and CanFam\_GSD, over 91% genes had reciprocal best hits for at least one protein isoform (Supplementary Table 3). To investigate this further, the Wags, CanFam3.1 and CanFam\_GSD genomes were mapped onto CanFam\_Bas and the coverage for each gene calculated with Diploidocus v0.10.0. Of the 27,129 predicted genes, 26,354 (97.1%) are found at least 50% covered in all four dogs, whilst only 30 (0.11%) are completely unique to  
265 CanFam\_Bas. In total, Wags is missing 302 predicted genes, CanFam\_GSD is completely missing 95 predicted genes, and CanFam3.1 is missing 211 predicted genes (Table 2). A considerably greater proportion of the missing genes in Wags (64.2% versus 11.4% in

CanFam3.1 and 15.8% in CanFam\_GSD) were on the X chromosome. To test for artefacts due to assembly errors we mapped the long read data for Wags and CanFam\_GSD onto  
 270 CanFam\_Bas. Only 7 of the 302 missing Wags genes (2.3%) had no long read coverage, whilst 21/95 (22.1%) of genes missing in CanFam\_GSD were confirmed by an absence of mapped long reads.

275 **Table 2. Predicted copy numbers for CanFam\_Bas GeMoMa genes based on A. assembly mapping, and B. long read mapping.**

<b>A. Dog</b>	<b>Missing</b>	<b>Partial (&lt;50%)</b>	<b>Single (1n)</b>	<b>Duplicate (2n)</b>	<b>3n+</b>
CanFam_Bas	0 (0)	0 (0)	27129 (25788)	0 (0)	0 (0)
CanFam_Wags	302 (108)	120 (58)	26103 (25035)	486 (472)	118 (115)
CanFam3.1	211 (187)	167 (161)	26404 (25125)	251 (223)	96 (92)
CanFam_GSD	95 (80)	48 (42)	26586 (25304)	299 (266)	101 (96)

<b>B. Data</b>	<b>0n</b>	<b>0.5n</b>	<b>1n</b>	<b>1.5n</b>	<b>2n</b>	<b>2.5n+</b>
CanFam_Bas (ONT)	2 (2)	1257 (1201)	24116 (22954)	1508 (1403)	80 (68)	166 (160)
CanFam_Wags (PacBio)	7 (2)	4717 (3476)	21140 (21049)	1028 (1024)	79 (79)	158 (158)
CanFam_GSD (ONT)	21 (18)	1893 (1795)	22412 (21350)	2503 (2349)	109 (98)	191 (178)

Figures in brackets exclude predicted genes on X chromosome.

### Amylase copy number

Two copies of the *Amy2B* gene were identified in a tandem repeat on Chromosome 6 of the  
 280 CanFam\_Bas assembly. The single-copy read depth for CanFam\_Bas, calculated as the modal read depth across single copy complete genes identified by BUSCO v3 [30], was estimated to be 34X. This was verified by BUSCO complete genes, which gave mean predicted copy numbers of  $1.008 \pm 0.005$  (95% C.I.) (Supplementary Fig 4). The two complete *Amy2B* coding sequence copies had a mean depth of 97.5X, equating to 2.87N, or a total

285 copy number estimate of 5.78N (2 x 97.5 / 34). The full CanFam\_GSD *Amy2B* repeat region was also found in two complete copies with a mean depth of 98.1X, estimating 5.77 copies (2 x 98.1 / 34). Similar results were obtained restricting analysis to reads at least 5kb (6.01 gene copies; 5.98 region copies) or 10kb (6.18 gene copies; 6.04 region copies) to minimise repeat-based fluctuations in read depth. In contrast, droplet digital PCR (ddPCR) estimated  
290 that China had 4.5 copies per individual chromosome. This slight difference suggests that the raw sequence data slightly overestimated copies or the ddPCR primers did not capture all the genetic variation.

Wags assembly has a single copy of the *Amy2B* region, which includes 90% of the *Amy2B* coding sequence (data not shown). Single-copy depth analysis of Wags estimates  
295 4.97 (90% at 253.8X) and 5.51 (100% at 253.5X) copies of the *AMY2B* coding sequence and tandem repeat unit, respectively.

### **Nuclear mitochondrial DNA fragments**

During the assembly of China, the mitochondrial genome was erroneously assembled into a nuclear mitochondrial DNA (NUMT) fragment on chromosome 29. A blastn search ( $e < 10^{-4}$ ) identified 291 putative NUMTs, ranging in size from 34 bp to 6,580 bp (Table 3). These  
300 total 190.5 kb (approx. 11.4 mtDNA copies) and span the entire mtDNA with reasonably even coverage and no obvious bias (Supplementary Fig. 5A). All 291 NUMTs are well-supported with at least 3 reads that span the entire NUMT plus at least 5 kb each side (Supplementary Table 4). Only 1 NUMT was not found to be full-length in CanFam\_GSD  
305 (Supplementary Fig. 5B). An additional 26 NUMTs are partially covered in CanFam3.1 and 9 are entirely absent. Whilst this could represent a breed difference, 19 of the 35 additional incomplete NUMTs in CanFam3.1 (65.5%) are also incomplete in Wags, whilst Wags has a further ten incomplete NUMTs that are present in CanFam3.1 (Supplementary Table 4). This

is consistent with these regions being generally harder to assemble and/or align. Further  
310 analyses of these regions may provide insight into domestic dog genealogies.

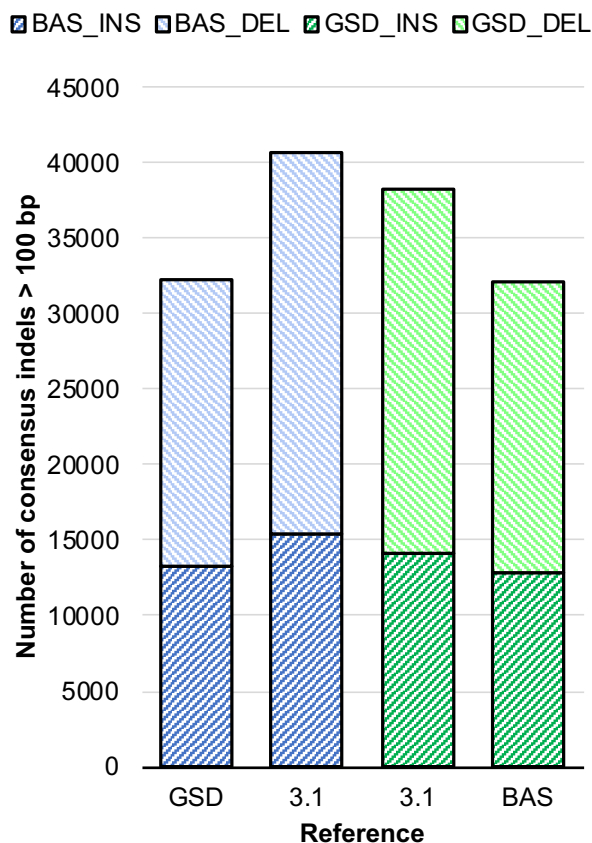
### **Whole genome assembly comparisons**

To discover unique large-scale structural differences in assembled genomes of the three  
breeds – Basenji, German Shepard and Boxer – we performed pairwise alignments of  
CanFam\_Bas, CanFam3.1 and CanFam\_GSD. Overall, genome synteny was maintained and  
315 there were limited large scale genomic rearrangements observed (Fig 4A). There was,  
however a large inversion in CanFam3.1 that was not present in CanFam\_Bas or  
CanFam\_GSD (Fig 4B). To investigate this further, we aligned CanFam\_Bas against Wags  
(Fig 4A iii). As expected there was no inversion on Chromosome 9 (Fig 4B iii), suggesting  
the inversion or perhaps an assembly error, occurs in CanFam3.1.

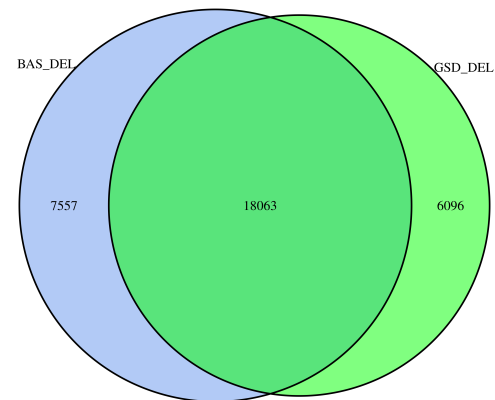
### **320 Long read structural variant detection**

To generate a conservative set of high-quality structural variants, the consensus of ONT and  
SMRT long read sequences data from both Basenji (China ONT and Wags SMRT) and GSD  
(Nala ONT and SMRT) was mapped onto Basenji (CanFam\_BAS), Boxer (CanFam3.1) and  
GSD (CanFam\_GSD) reference genomes (Supplementary Table 5). One difference in the  
325 data sets is that both ONT and SMRT reads from GSD were sourced from the same  
individual dog whereas for Basenji the ONT and SMRT reads are from different individual  
Basenji dogs. These high-quality SVs were additionally annotated for their overlap to both  
genes and exons. Of these high-quality ONT/SMRT consensus SVs, the Basenji long reads  
overlap 814 CanFam3.1 exons and 568 CanFam\_GSD exons while the GSD long reads  
330 overlap 825 CanFam3.1 exons and 495 CanFam\_Bas exons. In total, these SVs represent  
92.19 Mb, 97.21 Mb, and 78.99 Mb of deleted sequence and 4.11 Mb, 4.09 Mb, and 7.69 Mb  
of inserted sequence for the CanFam\_Bas, CanFam3.1, and CanFam\_GSD assemblies  
respectively.

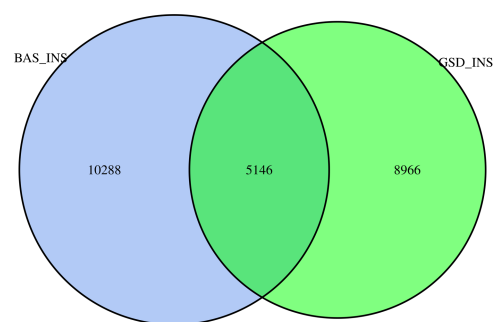
### A. Consensus SVs >100 bp



### B. Consensus deletions >100 bp vs CanFam3.1



### C. Consensus insertions >100 bp vs CanFam3.1



**Figure 5. Consensus structural variant calls for combined ONT and PacBio data.** High-quality set of consensus structural variant (SV) calls generated from the intersection of the ONT and PacBio SV calls for each breed versus reference comparison, limited to SVs >100 bp long. **A.** Total numbers of SVs called from Basenji CanFam\_Bas reads (red) versus CanFam3.1 and CanFam\_GSD, and German Shepherd long reads (green) versus CanFam3.1 and CanFam\_Bas. **B.** Numbers and overlap of consensus deletion calls for Basenji reads (blue) and GSD reads (green) versus CanFam3.1. **C.** Numbers and overlap of consensus insertion calls for Basenji reads (blue) and GSD reads (green) versus CanFam3.1.

To reduce the impact of small indels arising from local mis-assembly errors, the high-quality consensus SVs were further reduced to those over 100 bp in length (Fig 5). Overall, we observe similar number of total SV calls from Basenji long reads relative to CanFam\_GSD and GSD long reads relative to CanFam\_BAS. Both breeds had a substantially larger number of consensus SV calls against the CanFam3.1 reference overall, with Basenji long reads generating more SVs calls than GSD long reads. We next overlapped the CanFam SV calls relative to CanFam v3.1 and found 18,063 long read deletion calls overlapped between Basenji and GSD. For Basenji this represented 70.00% of the total 25,260 Basenji

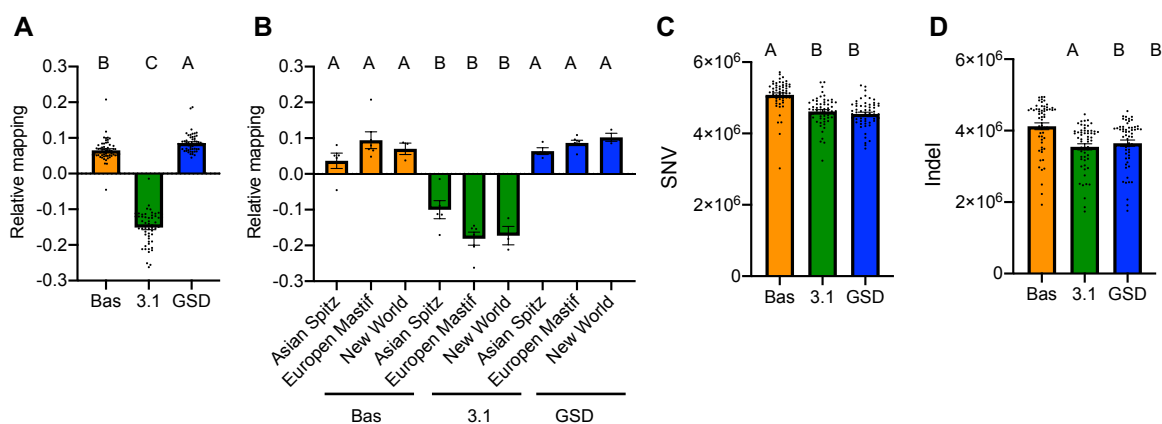
deletions while for GSD this represented 73.25% of the total 24,138 GSD deletions (Fig. 5B). Insertions were fewer in number and degree of overlap, however we still found 5,146 overlapping insertions between Basenji and GSD long reads representing 33.33% of the total 15,434 Basenji insertions and 36.46% of the total 14,111 GSD insertions (Fig. 5C). The high degree of overlap in SVs from GSD and Basenji relative to CanFam v3.1 represent Boxer-specific SVs or potential issues with the current canid reference assembly.

### **Short read mapping, SNV / small indel detection**

Mapping multiple individuals to a reference genome for variant calling is standard practice in population genomics and is known to be prone to biases in the reference genome. To investigate whether differences identified in variant analyses were due to the Basenji being a basal breed or due to assembly quality difference, short read data from 58 different dog breeds belonging to sixteen different well-supported clades from Parker et al. [10] (Supplementary Table 6) were mapped on to three reference genomes Basenji (CanFam\_BAS), Boxer (CanFam3.1) and GSD (CanFam\_GSD). Large-scale structural differences between breeds would be expected to significantly affect read mapping efficiencies for closely-related versus distantly-related breeds, whilst missing assembly data would be expected to result in a systematic reduction in mapping across all breeds. In our analysis, we observe such systematic and breed specific changes in both the number of mapped reads and variants detected.

For the systematic changes, overall trends are exhibited in the total percentage of reads mapped across the three references, with the highest percent of reads mapping to CanFam\_GSD, followed closely by CanFam\_BAS and the lowest percent of reads mapping to CanFam3.1 (Fig 6A). ANOVA shows these differences are significant ( $F_{2, 171}=819.53$ ,  $P<0.0001$ ). To investigate this result further and test for interactions we focused upon breeds within each of the monophyletic clades close to or associated with the three reference

genomes. Specifically, we included the short-read sequences from the four breeds in the Asian Spitz clade as this close to the basal Basenji lineage. We also included the six breeds European Mastif clade containing the Boxer and three breeds within the New World clade containing the GSD. Overall, the CanFam\_Bas and CanFam\_GSD performed equally well while the relative mapping was lowest for CanFam3.1 (Fig 6B). Once again ANOVA shows this result is significant ( $F_{8, 30}=32.01$ ,  $P<0.0001$ ). Next, we considered the capacity of each reference genome to detect SNV's and indels. In this case CanFam\_BAS detected higher number of changes than did either CanFam3.1 or CanFam\_GSD (Fig 6C and D; ANOVA  $F_{2, 171}=30.71$ , and  $F_{2, 171}= 12.08$ , respectively with  $P<0.0001$  for each). In combination these data attest to the quality of the CanFam\_Bas assembly. There is a noticeable depletion of variant calls for the reference breed (Supplementary Fig 6), but there were no significant interactions between clades and reference genome (data not shown).



**Figure 6. Comparative short read mapping and single nucleotide variant calling for 58 dog breeds versus three reference genomes: CanFam\_Bas (Bas), CanFam3.1 (3.1) and CanFam\_GSD (GSD).** **A.** Relative percentage of reads mapped to each reference calculated by subtracting the mean percentage of reads for the three reference genomes from the number of reads for each reference. **B.** Relative percentage of reads mapped to each reference for three closely related clades determined from Parker et al. [13]. The Basenji is closely related to the Asian Spitz clade. The Boxer is a member of the European Mastiff clade and the GSD is a member of the New world clade. **C.** The number of SNVs. **D.** The number of small indels.



## Discussion

In this manuscript we present a reference-quality assembly of a female Basenji (China),  
400 which we designate CanFam\_Bas. We also present a second Basenji genome, Wags. The  
Wags build is of high-quality but not as good as CanFam\_Bas, partly because Wags was a  
male. In total, 64.2% of the missing genes are on the X chromosome, compared to under 16%  
in the other two individuals. Analysis of long read data mapped onto the CanFam\_Bas  
genome annotation further supports assembly issues as a contributor, with only two  
405 autosomal genes lacking any coverage in Wags SMRT data. Equivalently, two other  
autosomal genes also lacked any coverage in China ONT data, suggesting that even high  
quality assemblies can have missing genes in an assembly that may not be biologically lost.

CanFam\_Bas is considerably more contiguous and complete than CanFam3.1  
(Boxer breed) and slightly more contiguous (Contig N50 37.8 kb vs 20.9 kb) but has slightly  
410 lower completeness (BUSCO completeness 92.9% vs 93.0%) than CanFam\_GSD [14] (Table  
1, Supplementary Table 1). The ungapped nuclear chromosomes lengths of CanFam\_Bas,  
CanFam3.1, and CanFam\_GSD are 2.33 Gb, 2.32 Gb, 2.36 Gb and, respectively. Likely,  
both assembly quality and breed differences contribute to the 211 missing genes in  
CanFam3.1. CanFam\_GSD had fewer predicted genes missing (95) but a higher percentage  
415 (22.0%) also lacking long read coverage. This uncertainty notwithstanding, our analysis  
highlights the need to consider multiple reference genomes in a pan-genomic approach when  
a comprehensive analysis is required. These results also highlight the need for considering  
missing genes carefully on a case-by-case basis as, even with high quality genome assemblies  
such as those compared in this study, an apparent absence may reflect missing data rather  
420 than missing biology.

We noted that the mitochondrial chromosome was missing from the China 1.0 Basenji  
assembly. Exhaustive search of the genome detected a 33.2kb region consisting of almost two

complete copies of the mitochondrial genome on Chromosome 29 that was not present in other dog assemblies. The real NUMT fragment was 4,302 bp in size and a total of 291  
425 (190.5 kb) were detected across the genome. NUMTs are troublesome insertions that are universal in nuclear DNA, but they are not ubiquitous in nuclear genome assemblies [37]. Very often NUMTs are edited out during the genome annotation process [38], but it is becoming increasingly clear that that such organelle insertions must be properly accounted for in nuclear genome assemblies as they have the potential to provide unique insight into  
430 population histories and animal well-being [39]. In domestic cats, the nuclear cat genome includes 7.9 kb fragment that is duplicated in tandem 38–76 times [40]. In humans NUMTs are polymorphic at 141 loci. NUMT polymorphisms have been used to estimate the age of of insertions in human and other primate lineages [39] while five insertions have been implicated in human disease [37, 41].

435         Sequence analyses detected six copies of *Amy2B* in Chia and five in Wags. Previous studies have shown that Basenji's may have 4-18 copies [6]. This high variation in *Amy2B* copy number suggests at least three evolutionary histories of the gene in Basenji's. First, *Amy2B* copies may have differentially accumulated in specific lineages since the divergence from other dog breeds. Second, the ancestral founding population of the modern Basenji may  
440 have been polymorphic for *Amy2B*. Third, it remains possible that the *Amy2B* region from other breeds have differentially introgressed into the modern breed dog. Basenji-like dogs are depicted in In Egyptian drawings and models dating back to 1550-1900 BC [11]. Possibly, the Basenji is derived from the Abuwtiyuw, which was a lightly built hunting dog with erect ears and a curly tail that died before 2280BC [42]. Likely, sequencing the region around  
445 *Amy2B* in Basenji's with a higher copy numbers will aid resolving the alternate hypotheses. This large genetic diversity in Basenji *Amy2B* copy has important dietary and veterinary implications. Veterinarians frequently recommend a rice based diet following complex

intestinal surgeries and following diarrhoea because it is thought to be bland and easily digestible. However, dogs with low *Amy2B* copy numbers may not be able to digest rice readily as their serum amylase levels are lower [43].

CanFam\_Bas and CanFam\_GSD represent two high-quality reference genomes from different breeds. The availability of corresponding Basenji and GSD long read data, provided an excellent opportunity to further investigate the contributions of breed and assembly differences, as real SVs will be represented in the raw read data even if assembly errors have introduced false positives or negatives. Possessing ONT and SMRT data for both GSD and Basenji allows us to overlap the disparate SV call sets to generate a conservative list of SV calls relative to CanFam3.1. This analysis identified over 70,000 SVs in CanFam\_Bas relative to CanFam3.1 and over 64,000 SVs in GSD relative to CanFam3.1 (Supplementary Table 5). There is a high degree of overlap in SVs from GSD and Basenji relative to CanFam v3.1 (68% of total basenji SV calls and 77% of GSD SV calls), which highlights potential issues with the current canid reference assembly. Further, each consensus set contains several hundred SVs overlapping annotated exons, highlighting the importance of the selection of appropriate reference genome for analysis of specific genomic regions.

Next, we examined the overlap of the consensus calls for SVs over 100bp of GSD and Basenji relative to CanFam v3.1. We find a high degree of agreement for deletions, with 70.1% of GSD calls and 74.75% of Basenji calls overlapping (Fig 5B), compared to insertions where 33.3% of basenji calls and 36.4% of calls overlap (Fig 5C). Basenji and GSD long reads have 25.9% and 19.1% more SVs called when CanFam3.1 is used as the reference, respectively, and the high degree of overlap is consistent with Boxer-specific SVs and/or assembly issues. GSD-specific and Basenji-specific SV calls are also evident.

To get additional insight into breed differences and the influence of reference genome selection, we mapped short read data from 58 breeds onto CanFam\_Bas, CanFam3.1, and CanFam\_GSD. If assembly differences are dominated by real differences between breeds then we might expect different breeds to map with different efficiencies onto the three  
475 genomes. Differences to due quality, on the other hand, should be reflected across all breeds. With some minor exceptions, short read data from the different breeds consistently mapped better onto CanFam\_GSD than CanFam\_Bas, which was in turn better than CanFam3.1 (Fig 6A). This is consistent with assembly quality being a dominating factor.

480 After adjusting for read mapping difference, different reference genomes also produce different SNV and small indel densities for the 58 mapped breeds (Supplementary Fig 6). Using a Basenji reference genome consistently identifies more variants than either GSD or Boxer (Fig 6, Supplementary Fig 6). This probably reflects the basal position of Basenji in the breed phylogeny [10]. The basal position of the Basenji makes it useful as a  
485 reference for variant analysis as there are clear biases affecting related breeds seen for both the GSD and Boxer reference genomes. On the other hand, CanFam\_GSD has slightly higher read mapping levels across breeds, which may provide better total coverage.

Together, these data suggest that a single high-quality reference should be sufficient for most general analyses, but the generation of clade specific genomes is likely to be  
490 important for canine nutrition and disease studies. Fanconi Syndrome has been reported in several dog breeds, but the vast majority (75%) have occurred in Basenjjs. Approximately, 10-30% of Basenji's in North America are affected. The most severe and common ailment in Basenjjs is Fancioni Syndrome, in which the renal tubes fail to reabsorb electrolytes and nutrients [44]. In 2011, it was shown that Basenji Fanconi Syndrome is caused by a 370bp  
495 deletion on canine chromosome 3 [45]. To date no other breeds have been recoded with this same deletion, although other mutations can cause the disease. Likely complex diseases in

dogs belonging to different clades may have different underlying causes. For example, it remains unclear whether the same suite of mutations causes hip dysplasia in the GSD (New World clade) and the Labrador (Retriever/ Spaniel clade).

## 500 **Conclusions**

Here, we present two high quality *de novo* Basenji genome assemblies: CanFam\_Bas (China, female) and Wags (male). CanFam\_Bas offers improved genome contiguity relative to CanFam v3.1 and can serve as a representative basal breed in future canid studies. We generate core genomic information for the Basenji that has the potential to inform future studies of population history and aid disease management. We generate high-quality variants (SV's, SNVs, small indels, and SVs) relative to CanFam\_Bas, CanFam v3.1, and CanFam\_GSD. We demonstrate the impact that the reference genome makes on both read mapping and variant detection, illustrating the importance of either selecting the appropriate reference genome or employing a pan-genome approach in future canid studies.

## 510 **Methods**

### **Sequencing and genome assembly of female Basenji, China**

**Sampling.** China or Zanzipow Bowies China Girl is an Australian Supreme Champion Kennel Club show dog. Her Australian National Kennel Council registration number is 2100018945. She is bred primarily from Australian lines, with her most recent common ancestor coming from Africa 18 generations previously. She was born on 14 Jan 2016 and is free from all known currently reported diseases.

**Sequencing.** High molecular weight DNA was extracted from 100 µl of blood using the DNeasy Blood and Tissue kit (Qiagen). For long read (Oxford Nanopore) sequencing, 1 µg of DNA was prepared via the Genomic DNA by Ligation kit (SQK-LSK109) as per the manufacturer's protocol. Prepared DNA (180 ng) was loaded onto a PromethION (FLO-

PRO002) flowcell and sequenced with standard parameters. After 48 hours, a nuclease flush was performed and an additional 115 ng was loaded onto the run. Base-calling was performed after sequencing with the GPU-enabled guppy basecaller (v3.0.3) using the PromethION high accuracy flip-flop model with config 'dna\_r9.4.1\_450bps\_hac.cfg'.

525 For short read sequencing whole blood was shipped to BGI, Hong Kong. High molecular weight DNA was extracted, a paired-end library prepared, and the sample run on the BGISEQ-500 sequencing system to generate high quality PE100 data. A total of 767,111,208 clean reads (115.1Gb) were produced with a lower base call accuracy (Q20) of 95.92%.

530 **Assembly.** An overview of the China assembly workflow is given in Supplementary Fig 1A. The ONT reads were assembled with the Flye (v2.6-release) assembler [16, 17]. The resulting contigs were polished with ONT reads using four rounds of Racon (v1.4.3) [18] followed by Medaka (v0.10.0) [19] to minimise error propagation. BGI-seq reads were aligned to the polished assembly with BWA-mem (v 0.7.17) [46] and Pilon (v1.23) (diploid) [20] was used for further error correction. A second assembly was performed using Canu assembler v1.8.0 [47] and correcting the sequencing errors using 2 rounds of Arrow polishing [48]. The Flye assembly was considered more contiguous and therefore selected as the core assembly.

540 **Scaffolding.** The ONT assembly was scaffolded to chromosome-length by the DNA Zoo following standard DNA zoo methodology ([www.dnazoo.org/methods](http://www.dnazoo.org/methods)). Briefly, an *in situ* Hi-C library was prepared [23] from a blood sample from the same individual and sequenced to ~30x coverage (assuming 2.4 Gb genome size). The Hi-C data was processed using Juicer [49], and used as input into the 3D-DNA pipeline [50] to produce a candidate chromosome-length genome assembly. We performed additional finishing on the scaffolds using Juicebox  
545 Assembly Tools [21]. The matrices are visualized in Juicebox.js, a cloud-based visualization

system for Hi-C data [22] and are available for browsing at multiple resolutions. After scaffolding, all ONT reads were aligned to the assembly with Minimap2 (v2.16) (-ax map-ont) [26] and used by PBJelly (pbsuite v.15.8.24) [24] to fill gaps. BGI reads were re-mapped with BWA-mem (v 0.7.17) and another round of polishing was performed with Pilon (v1.23) (diploid, SNP and indel correction) [20].

**Final clean-up.** The Pilon-polished genome underwent a final scaffold clean-up to generate a high-quality core assembly following Field and colleagues' processing of Canfam\_GSD [14]. Scaffolds were reoriented and assigned to CanFam3.1 chromosomes using PAFScaff v0.3.0 [14] (Minimap2 v2.16 mapping) [26]. Diploidocus v0.9.0 [51] was used to screen contamination, remove low-coverage artefacts and haplotig sequences, and annotate remaining scaffolds with potential issues as described in [14]. ONT reads were mapped onto the assembly using Minimap2 v2.17 (-ax map-ont --secondary=no) [26] and read depth summaries calculated with BBMap v38.51 pileup.sh [52]. Any scaffolds with median coverage less than three (e.g., less than 50% of the scaffold covered by at least three reads) were filtered out as low-coverage scaffolds. Single-copy read depth was estimated using the modal read depth of 35X across the 5,5578 single copy complete genes identified by BUSCO v3.0.2b [30]. This was used to set low-, mid- and high-depth thresholds for PurgeHaplotigs v20190612 [53] (implementing Perl v5.28.0, BEDTools v2.27.1 [54, 55], R v3.5.3, and SAMTools v1.9 [56] at 8X, 25X and 68X. PurgeHaplotig coverage was adjusted to exclude gap regions and any scaffolds with 80%+ bases in the low/haploid coverage bins and 95%+ of their length mapped by PurgeHaplotigs onto another scaffold were filtered as haplotigs or assembly artefacts. Any other scaffolds with 80%+ low coverage bases were filtered as Low Coverage. Remaining scaffolds were further classified based on read depth profiles. Scaffolds with <20% diploid coverage and 50%+ high coverage were marked as probable collapsed repeats. Scaffolds with "Diploid" depth as the dominant PurgeHaplotigs

coverage bin and >50% match to another scaffold were marked as a possible repeat sequences [14].

**Genome assembly correction.** Main chromosome scaffold integrity was checked using D-GENIES [28] comparisons of different assembly stages with CanFam\_GSD chromosomes [14]. Two pairs of fused chromosomes were identified following incorrect joins made by PBJelly (pbsuite v.15.8.24) [24]. Pre-gap-filled HiC scaffolds were mapped onto the assembly using Minimap2 v2.17 [26] and parsed with GABLAM v2.30.5 [27] to identify the gap corresponding to the fusion regions. These were manually separated into four individual chromosomes, gap lengths standardised, and scaffolds re-mapped onto CanFam v3.1 using PAFScaff v0.4.0 [25]. Finally, scaffolds under 1kb were removed following correction of Chromosome 29 (below).

**Correction of mitochondrial insertion into Chromosome 29.** NUMT analysis identified a 33.2 kb region consisting of almost two complete copies of the mitochondrial genome, not present in other dog genome assemblies. GABLAM v2.30.5 [27] was used to confirm that this region was also absent from the Canu assembly of *China*. ONT reads that mapped onto both flanking regions of the 33.2 kb putative NUMT were extracted and reassembled with Flye v2.7.1 [16, 17]. The new NUMT was approx. 2.8 kb long. To avoid repeated problems with mitochondrial reads mis-polishing the sequence, a subset of 4.82M ONT reads (72.7 Gb, ~30X) was extracted and mapped onto the assembled region with Minimap 2.17 [26]. Reads mapping to at least 5 kb of the assembled region including some immediate flanking sequence were extracted (66 reads, 1.50 Mb) and polished with one round of Racon v1.4.5 [18] (-m 8 -x -6 -g -8 -w 500) and Medaka v0.7.1 [19] (model r941\_prom\_high). The polished NUMT region was mapped on to the Chromosome 29 scaffold with GABLAM v2.30.5 [27] (blast+ v2.9.0 [57] megablast) and stretches of 100% sequence identity



595 identified each side of the NUMT. The mtDNA sequence between these regions of identity was replaced with the re-assembled NUMT sequence.

**Mitochondrial genome assembly.** To assemble the mitochondrion, ONT reads were mapped onto a construct of three tandem copies of the CanFam3.1 mtDNA with minimap2 v2.17 [26] (-ax map-ont --secondary=no). Reads with hits were extracted using SAMTools v1.9 [56] 600 fasta and mapped onto a double-copy CanFam3.1 mtDNA with GABLAM v2.30.5 [27] (blast+ v2.9.0 [57] megablast). “Pure complete” mtDNA ONT reads were identified as those with 99%+ of their length mapping to mtDNA and 99%+ coverage of the mtDNA. These reads were assembled with Flye 2.7b-b1526 [16, 17] (genome size 16.7 kb) and polished with Racon v1.4.5 [18] (-m 8 -x -6 -g -8 -w 500) followed by Medaka v0.7.1 [19] (model 605 r941\_prom\_high). The polished mtDNA assembly was mapped onto CanFam3.1 mtDNA with GABLAM v2.30.5 [27] (blast+ v2.9.0 [57] megablast) and circularised by extracting a region from the centre of the assembly corresponding to a single full-length copy with the same start and end positions. Final correction of SNPs and indels was performed by adding the mtDNA to the nuclear assembly, mapping BGI reads with BWA v0.7.17 and polishing 610 the mtDNA with Pilon v1.23 [20]. The polished mtDNA was then added back to the nuclear genome for the final *China* assembly.

**Genome assembly quality assessment.** At each stage of the assembly, summary statistics were calculated with SLiMSuite SeqList v1.45.0 [58], quality was assessed with Merquy v20200318 (Meryl v20200313, bedtools v2.27.1 [54, 55], SAMTools v1.9 [56], java v8u45, 615 igv v2.8.0) and completeness assessed with BUSCO v3.0.2b [30] (BLAST+ v2.2.31 [57], HMMer v3.2.1 [59], Augustus v3.3.2, EMBOSS v6.6.0, laurasiatherian lineage (n=6253)). To account for fluctuations in BUSCO ratings, presence of complete BUSCO genes across assembly stages was also assessed with BUSCOMP v0.9.4 [60]. Final assembly scaffold statistics and quality assessment was performed with Diploidocus v0.10.2 (KAT v2.4.2, perl

620 v5.28.0, BEDtools v2.27.1 [54, 55], SAMTools v1.9 [56], purge\_haplotigs v20190612, java  
v8u231-jre, bbmap v38.51, minimap2 v2.17 [26], BLAST+ v2.9.0 [57]).

***DNA methylation calling.*** China's blood DNA methylome libraries were sequenced on the  
Illumina HiSeq X platform (150 bp, PE), generating 336 million read pairs and yielding 14x  
625 sequencing coverage. Sequenced reads were trimmed using Trimmomatic [61] and mapped  
to the China v1.0 genome reference using WALT [62] with the following parameters: -m 10 -  
t 24 -N 10000000 -L 2000. The mappability of the MethylC-seq library was 86%. Duplicate  
reads were removed using Picard Tools v2.3.0. Genotype and methylation bias correction  
were performed using MethylDackel with additional parameters: minOppositeDepth 5 --  
maxVariantFrac 0.5 --OT 10,140,10,140 --OB 10,140,10,140. The numbers of methylated  
630 and unmethylated calls at each CpG site were determined using MethylDackel  
(<https://github.com/dpryan79/MethylDackel>). Bisulphite conversion efficiency was 99.71%,  
estimated using unmethylated lambda phage spike-in control.

***UMR and LMR calling.*** Segmentation of basenji's blood DNA methylome into CpG-rich  
unmethylated regions (UMRs) and CpG-poor low-methylated regions (LMRs) was  
635 performed using MethylSeekR [33] (segmentUMRsLMRs(m=meth, meth.cutoff=0.5,  
nCpG.cutoff=5, PMDs = NA, num.cores=num.cores, myGenomeSeq=build,  
seqLengths=seqlengths(build), nCpG.smoothing = 3, minCover = 5).

### **Sequencing and genome assembly of male Basenji, Wags**

Wags DNA was derived from blood of a single male. He is registered as American Kennel  
640 Club Champion Kibushi the Oracle, born on December 3, 2008. His registration number is  
HP345321/01. Sire is AM Ch C-Quests Soul Driver, HM827502/02, and his dam is  
Avongara Luka, HP345312/01, a native bitch imported from the Haut-Ule district of the DRC  
Congo, 3°24'04.0"N 27°19'04.6"E, in 2006. SMRT sequences for Wags (Fig. 1B) were  
generated on the Pacific Biosciences Sequel instrument (V2 chemistry) to approximately 45x

645 genome coverage based on a genome size estimate of 2.5 Gb. An overview of the China  
assembly workflow is given in Supplementary Fig 1B. All SMRT sequences were assembled  
with the HGAP4 algorithm, a Falcon based assembly pipeline available through the SMRT  
Link interphase (SMRT Link v5.0.1.9585) [63]. The assembly was then error corrected with  
the original SMRT sequences using the Arrow error-correction module [63]. Additional  
650 polishing of the assembly for residual indels was done by aligning 32x coverage of Illumina  
data and the Pilon algorithm [20]. Chromosomal level scaffolds were generated with the  
same DNA source using the Proximo<sup>TM</sup> Hi-C genome scaffolding software (Phase Genomics  
Inc) and finalized by alignment to the CanFam3.1 reference.

### **Locus copy number estimation**

655 Copy numbers for specific assembly regions were calculated using Diploidocus v0.10.0  
(runmode=regcnv) [51]. For each animal, long reads were mapped onto the assembly with  
Minimap2 v2.17 (no secondary mapping) [26] and the modal read depth across single-copy  
Complete genes identified by BUSCO v3 [30] (laurasiatherian\_db) calculated using  
SAMTools v1.9 [56] mpileup. This set the expected single-copy read depth,  $X_{SC}$ . Copy  
660 number for a region,  $N_{reg}$  was estimated by dividing the mean read depth across that region,  
 $X_{reg}$ , by  $X_{SC}$ . The variance and standard deviation of the estimate was calculated using  $X_{reg}$  for  
all single copy BUSCO genes. For Wags (a male), genes on the X chromosome were  
excluded from this analysis.

***Amylase copy number.*** The copy number of the beta amylase gene *Amy2B* was calculated  
665 using Diploidocus v0.10.0 (runmode=regcnv) [51] using a modification of the single locus  
copy number estimation (above) to account for multiple copies of the gene in the assembly.  
First, the AMY2B protein sequence from CanFam3.1 (UniprotKB: J9PAL7) was used as a  
query and searched against the genome with Exonerate v2.2.0 [64] to identify assembled  
copies of the *Amy2B* gene. A full-length (14.8 kb) *Amy2B* gene repeat was also extracted

670 from the CanFam\_GSD assembly and mapped onto the assembly with Minimap2 v2.17 (-x  
asm5) [26]. Estimated  $N_{reg}$  values were converted into a number of copies by multiplying by  
the proportion of the query found covered by that region. The total genome *Amy2B* copy  
number was then calculated as the summed copy number over each hit. To further investigate  
the robustness of the method and improve the *Amy2B* copy number estimate in China,  
675 analysis was repeated with ONT reads at least 5 kb in length and at least 10 kb in length.  
These reads should be less susceptible to poor mapping at repeat sequences, but at a cost of  
reduced coverage.

We also used droplet digital PCR (ddPCR) to directly quantify the *Amy2B* copy  
number of China [65]. ddPCR was performed using a QX100 ddPCR system (Bio-rad). Each  
680 reaction was performed in a 20 $\mu$ l reaction volume containing 10  $\mu$ l of 2x ddPCR Supermix  
(Bio-rad), 1 $\mu$ l of each 20x primer/probe, 1 $\mu$ l of DraI restriction enzyme (New England  
BioLabs #R0129S), 5 $\mu$ l of DNA template (4ng/ $\mu$ l) and 2 $\mu$ l ddH<sub>2</sub>O. Primer sequence for  
*Amy2B*: forward 5'-CCAAACCTGGACGGACATCT-3' and reverse 5'-  
TATCGTTCGCATTCAAGAGCAA-3' with FAM probe: 6FAM-  
685 TTTGAGTGGCGCTGGG-MGBNFQ. Primer sequence for *C7orf28b-3*: 5'-  
GGGAAACTCCACAAGCAATCA-3' and reverse 5'-GAGCCCATGGAGGAAATCATC-3'  
with HEX probe HEX-CACCTGCTAACAGC-MGBNFQ.

### **Nuclear mitochondrial DNA (NUMT) fragment analysis**

To make sure that any nuclear mitochondrial DNA fragments (NUMTs) were identified as a  
690 single region, a double-copy CanFam3.1 mtDNA sequence was constructed and then  
searched against the CanFam\_Bas nuclear genome and compressed to unique hits using  
GABLAM v2.30.5 [27] (blast+ v2.9.0 [57] blastn, localunique) with a blastn e-value cut-off  
of 1e-4 [29]. Predicted copy number was calculated for each NUMT in *China* using the  
method described above. Diploidocus v0.10.0 was also used to calculate number of reads

695 spanning each entire NUMT plus flanking regions of 0 bp, 100 bp, 1 kb and 5 kb. In addition,  
assembly coverage for each NUMT was calculated for Wags, CanFam3.1 and CanFam\_GSD.  
Each genome was split into 1 Mb tiled fragments and mapped onto CanFam\_Bas with  
Minimap2 v2.17 [26]. Each BAM file was used for Diploidocus v0.10.0 regenv [51] analysis  
with a single-copy read depth of 1X. Mitochondrial genome coverage was analysed by  
700 extracting all 291 NUMT regions with SeqSuite v1.23.3 [66] and mapping them onto the  
CanFam3.1 mtDNA chromosome using GABLAM v2.30.5 [27] (blast+ v2.10. [57] tblastn).

### Genome annotation

Following Field et al. [14] each genome was annotated using the homology-based gene  
prediction program GeMoMa [35] (v1.6.2beta) and nine reference organisms. For each  
705 reference organism, coding exons of full-length transcript were extracted and translated to  
peptides using the GeMoMa module Extractor. These peptides were searched in each Basenji  
genome using mmseqs2 [67] (v5877873cbcd50a6d954607fc2df1210f8c2c3a4b). Based on  
the results of mmseqs2 and Extractor, transcripts were predicted for each Basenji genome  
from each reference organism independently. These nine gene annotation sets were then  
710 combined into a final gene annotation using the GeMoMa module GAF. To make a fair  
comparison of the influence of genome quality and completeness on annotation, CanFam3.1  
was annotated with the same pipeline. Annotation of CanFam\_GSD using the same pipeline  
was obtained from Field et al. [14].

***Annotation summary and quality assessment.*** Annotation summary statistics and the longest  
715 protein isoform per gene were generated with SAAGA v0.4.0 [68]. Annotation completeness  
was estimated using BUSCO v3 [30] (laurasiatherian,  $n=6253$ , proteins mode), run on a  
reduced annotation consisting of the longest protein per gene. To check for truncated or  
fragmented protein predictions, predicted proteins were mapped onto the Quest For  
Orthologues reference dog proteome [36] with mmseqs2 v [67]. The best protein hit for each

720 gene was used to calculate a protein length ratio (length of predicted protein / length of  
reference protein). Percentage coverage of query and hit proteins was calculated with  
mmseqs2 v [67]. A reciprocal search identified the closest predicted protein for each  
reference protein. Any reciprocal best hits were marked as predicted orthologues.

***Annotation copy number and coverage analysis.*** Predicted copy number was calculated for  
725 every protein-coding gene in CanFam\_Bas using the method described above. In addition,  
assembly coverage for each CanFam\_Bas gene was calculated for Wags, CanFam3.1 and  
CanFam\_GSD. Each genome was split into 1 Mb tiled fragments and mapped onto  
CanFam\_Bas with Minimap2 v2.17 (-ax asm5 -L) [26]. Each BAM file was used for  
Diploidocus v0.10.0 regcnv analysis with a single-copy read depth of 1X. In addition, SMRT  
730 reads from Wags and ONT reads from the GSD were mapped onto CanFam\_Bas with  
Minimap2 v2.17 (--secondary=no -L -ax map-pb or -ax map-ont) [26] and the standard  
predicted copy number calculation applied. Genes with zero coverage were marked 0n. Other  
genes were binned according to coverage: for mapped assemblies, coverage was rounded to  
the nearest integer; for long read mapping, coverage was rounded to the nearest 0.5n. Genes  
735 with greater than zero but less than 50% coverage were assigned to 0.5n. Any genes  
exceeding a rounded coverage of 2n were grouped as “3+”.

***Ribosomal RNA prediction.*** For each genome, genes for rRNA were predicted with Barrnap  
v0.9 [69] in the eukaryotic mode, implementing Perl v5.28.0, HMMer v3.2.1 [59] and  
BEDTools v2.27.1 [54, 55].

#### 740 **Whole genome assembly comparisons**

Whole genome synteny analysis were performed for the main chromosome scaffolds using  
the D-GENIES [28] web portal.

#### **Long read structural variant detection**

Structural variant calls were generated using a combination of minimap2 v2.17-r943-dirty

745 [26], SAMTools v1.9 [56], and sniffles v1.0.11 [70]. In total, four sets of long reads from  
three samples were analyzed consisting of China the Basenji (Oxford Nanopore), Wags the  
Basenji (SMRT) and Nala the German Shepherd (Oxford Nanopore and SMRT). Variants for  
the Basenji and Nala were annotated with gene model predictions generated using GeMoMa  
[35] (v1.6.2beta) while CanFam3.1 variants were annotated with Ensembl gene annotations  
750 v100 for CanFam v3.1 [71]

### **Short read mapping, SNV / small indel detection**

Representative Illumina data was identified from [72]. All SRA IDs associated with the  
DBVDC bioproject on NCBI (SRP144493) were downloaded along with their metadata and  
reduced to 126 samples representing the biggest sequencing run (no. bases) per annotated  
755 breed. These 126 samples were used for initial read mapping and variant calling  
(Supplementary Table 7). Following removal of unknown/mixed/village dog samples, canids  
other than domestic dogs, and duplicates the remaining breeds were mapped onto those used  
by Parker et al. [10]. In total, 58 breeds in the significantly monophyletic clades (greater than  
70% bootstrap) designated by Parker et al. [10] were considered for analyses (Supplementary  
760 Table 6).

SNVs and small indels were called from the Illumina reads of the 58 representative  
breeds against three reference genomes (Basenji China v1.0, CanFam3.1, and  
CanFam\_GSD). All Illumina reads were downloaded from the short read archive using the  
SRA toolkit v2.10.4 [73]. All samples were analysed using a modified version of an existing  
765 variant detection pipeline [74]. Briefly, the pipeline employs the Burrows-Wheeler Aligner  
v0.7.17-r1188 (BWA) for read alignment [75], SAMTools v1.9 [56] and picard v2.4.1  
(<http://broadinstitute.github.io/picard>) for binary alignment map (BAM) file preprocessing,  
and Genome Analysis ToolKit v3.6 (GATK) for calling SNVs and small indels [76]. The  
workflow follows GATK best-practices using default parameters throughout. Additionally,

770 SRA reads from each reference genome were aligned using BWA and SAMTools and a  
consensus variant list generated using an approach described previously [77]. The consensus  
variant lists for each reference genome were utilized in GATK's BaseRecalibrator step as the  
'-knownSites' argument to serve as the required variant truthset. Variant alignment statistics  
were generated using SAMTools flagstat. Joint variant calls were generated for the larger  
775 dataset of 126 samples relative to the three reference genomes and the total number of SNVs  
and small indels for individual samples tabulated. Variants for the Basenji and CanFam\_GSD  
were annotated with gene model predictions generated using GeMoMa (see above) while  
CanFam variants were annotated with ENSEMBL gene annotations v100 for CanFam v3.1.  
Read mapping statistics for each sample were calculated using SAMTools to remove  
780 secondary mapping and then summarized with v38.51 pileup.sh [52]. Numbers were then  
converted into relative values for each reference by averaging the score for each breed over  
the three reference genomes and then calculating the difference from the mean. Breeds were  
considered individually, but mean values for each clade were also calculated. Three clades  
are of particular importance as they are closely related, or include, the three reference  
785 genome assemblies. The Asian Spitz clade is considered closely related to the Basenji. This  
clade contained the Alaskan Malamute, Shar-Pei, Shiba Inu and Tibetan Mastiff. The  
European Mastiff Clade contained the Boxer, Bull Terrier, Cane Corso, Great Dane, Mastiff  
and Rhodesian Ridgeback. The New World clade contained the Berger Picard, Chinook and  
German Shepherd. One-way ANOVA's were employed to detect significant differences  
790 between groups. As the same short read samples were examined relative to the three  
reference genomes statistical significance was set to be  $P < 0.01$ .

## Acknowledgements

We would like to thank Jennifer Power for introducing us to China and Barbara Reisinger,  
Jon Curby and Jared Reisinger for access to Wags. For China, Brooke Morgan-Burke at the



795 Vineyard Veterinary Hospital provided constant encouragement. ONT data were collected at the Garvan Institute and the nuclease flush by Jillian Hammond, the BGI -seq data was provided by BGI and the Hi-C data collected at Baylor College of Medicine. For Wags, DNA sequencing was completed at the The Hudson Alpha Institute and assembly at the The McDonnell Genome Institute, Washington University School of Medicine.

## 800 **Funding**

This work was supported by the the University of New South Wales/ School of Biotechnology and Biomolecular Sciences Genomics Initiative and the Basenji Health Endowment Inc, Poynette, WI and. The DNA Zoo initiative of E.A.L. funded the Hi-C data collection and analyses. M.F. is funded by NHMRC APP5121190. E.L.A. was supported by 805 an NSF Physics Frontiers Center Award (PHY1427654), the Welch Foundation (Q-1866), a USDA Agriculture and Food Research Initiative Grant (2017-05741), an NIH 4D Nucleome Grant (U01HL130010), and an NIH Encyclopaedia of DNA Elements Mapping Center Award (UM1HG009375).

## **Author contributions**

810 J.W.O.B. coordinated the project. R.J.E., M.A.F., and J.W.O.B. designed the study. J.W.O.B. and W.W. funded the project. For China, R.A.Z. provided the sample. J.F. and B.D.R. performed the initial assembly and polishing, O.D., A.O. and Z.C. performed the Hi-C experiment, and O.D. and E.L.A. conducted the Hi-C analyses. R.J.E performed the final assembly polishing, curation, correction, analyses of genome completeness, mitochondrial 815 genome assembly, NUMT analysis, and read-depth gene copy number analyses. S.G.T. completed the digital droplet PCR. K.S. and O.B. conducted the DNA methylation analyses. M.A.F. performed the structural comparisons, dog breed short read mapping and variant calling. M.A.F., R.J.E. and J.W.O.B. performed the dog breed variant analysis. E.R., L.H., G.S.J., and W.W. performed the Wags assembly. J.K. performed the genome annotation.

820 R.J.E. M.F., M.A.F, B.D.R., O.D., O.B. and J.W.O.B. wrote the manuscript. All authors  
edited and approved the final manuscript.

## **Ethics declarations**

### **Ethics approval and consent to participate**

For China, all experimentation was performed under the approval of the University of New  
825 South Wales Ethics Committee (ACEC ID: 18/18B).

### **Competing interests**

JF has received travel and accommodation expenses to speak at Oxford Nanopore  
Technologies conferences. Otherwise, the authors declare that the submitted work was  
carried out in the absence of any professional or financial relationships that could potentially  
830 be construed as a conflict of interest.

### **Supplementary information**

Supplementary information accompanies this paper at:  
<http://www.slimsuite.unsw.edu.au/research/basenji/>

### **Availability of data and materials**

835 Chromosome-length genome assemblies for both Basenji are available at NCBI  
(CanFam\_Bas (*China*), GCA\_013276365.1; *Wags*, GCA\_004886185.2). The CanFam\_Bas  
mitochondrial genome is available at NCBI Genbank: MW051511. To review GEO  
accession GSE159396: Go to  
<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE159396>

840 Enter token ylc1qswyvh1tclr into the box

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