# GigaScience

# Genomic evidence of neo-sex chromosomes in the Eastern Yellow Robin --Manuscript Draft--

Manuagint Number			
Manuscript Number:	GIGA-D-19-00237R1		
Full Title:	Genomic evidence of neo-sex chromosomes in the Eastern Yellow Robin		
Article Type:	Data Note		
Funding Information:	Australian Research Council (ARC) Discovery Project (DP180102359)	Prof Paul Sunnucks	
	ARC Linkage Grant (LP0776322)	Prof Paul Sunnucks	
Abstract:	Background: Understanding sex-biased natural selection can be greatly enhanced by access to well-annotated chromosomes including ones inherited in sex-specific fashions. The Eastern Yellow Robin (EYR) is an endemic Australian songbird inferred to have experienced climate-driven sex-biased selection and is a prominent model for studying mitochondrial-nuclear interactions in the wild. However, the lack of an EYR reference genome containing both sex chromosomes (in birds, a female bearing Z and W chromosomes) is limiting current efforts to understand the mechanisms of these processes. Here, we assemble the genome for a female EYR and use low depth (10 ×) genome resequencing data from 19 individuals of known sex to identify chromosome fragments with sex-specific inheritance.		
	categories of sex-linked genomic regions. First, 653 W-linked scatfolds (25.7 Mb) were anchored to the W sex chromosome and 215 Z-linked scatfolds (74.4 Mb) to the Z. Second, 1138 W-linked scatfolds (70.9 Mb), and 179 Z-linked scatfolds (51.0 Mb), were anchored to a large section (coordinates ~5 to ~60 Mb) of Zebra Finch chromosome 1A. The first ~5 Mb and last ~14 Mb of the reference chromosome 1A had only autosomally-behaving EYR scatfolds mapping to them.		
	Conclusions: We report a female (W-chrom provide genomic evidence for a neo-sex (ne EYR, involving most of a large chromosome autosomal in passerines.	osome containing) EYR genome and eo-W and neo-Z) chromosome system in e (1A) previously only reported to be	
Corresponding Author:	Han Ming Gan AUSTRALIA		
Corresponding Author Secondary Information:			
Corresponding Author's Institution:			
Corresponding Author's Secondary Institution:			
First Author:	Han Ming Gan		
First Author Secondary Information:	indary Information:		
Order of Authors:	Han Ming Gan		
	Stephanie Falk		
Hernán E. Morales			

Powered by Editorial Manager  ${\ensuremath{\mathbb R}}$  and ProduXion Manager  ${\ensuremath{\mathbb R}}$  from Aries Systems Corporation

	Christopher M. Austin
	Paul Sunnucks
	Alexandra Pavlova
Order of Authors Secondary Information:	
Response to Reviewers:	Editor: -In your revised manuscript, please include a citation to your upcoming GigaDB dataset (including the DOI link) to your reference list, and cite this in the data availability section and elsewhere in the manuscript, where appropriate. -In the data availability section, please write something along the lines of " Supporting data, including the genome assembly, annotations, and [ mention other data types] are available via the GigaScience database, GigaDB [xx]."
	Added GigaDB in the data availability citation as per recommended format. (DOI yet to be made available)
	Reviewer #1:
	-I did not understand the reasoning behind the "evolutionary strata" argument in Figure 5C. I only realized what the point of this analysis was when I read the suggestion by Reviewer 3. Please explain this better in the ms.
	We have adjusted the text accordingly: One expectation under sex chromosome evolution is the development of evolutionary strata – regions of suppressed recombination identified by spatial clusters of Z-W orthologs with similar divergence estimates [43].
	-Line 54: "sex-linked genome region". genomic or regions. Genome region -> genomic regions
	-line 174: Remove underlined ".". Removed
	-line 264: "Surprisingly,A substantial". Fix capital "A" Fixed
	Reviewer #2:
	Line 264 - "A" should be lowercase Fixed
	Line 326 - Fix "to decrease" Fixed
	Line 327 - More appropriate citations would be Charlesworth & Charlesworth 2000 Phil Trans Roy Soc B and Charlesworth 1994 Genet Res. Also, note that reduced purifying selection due to the reduced effective population sizes of the W/Y chromosomes is only one of the reasons why we expect accumulation of deleterious mutations, along with Mueller's ratchet and hitchhiking (see Bachtrog 2006 Current Opinion in Genetics & Development).

	Citations added. We also added 'along with Mueller's ratchet and hitchhiking'.	
	Line 345 - Remove comma (or insert comma between "approaches" and "followed" in previous line) Inserted comma in previous line.	
	Line 357 - Fix to "neoZ-1A differences" Fixed	
	Figure 2A - This figure is not currently cited in the manuscript	
	Did a search in Microsoft word and confirm that Figure 2A is cited in the manuscript. "genome profiling based on Jellyfish2-calculated k-mer frequency (k=25) that estimated a genome size of 993 Mb with 1.12% heterozygosity for EYR054 (Figure 2A) [22-24]. We used MaSuRCA v3.2.4"	
Additional Information:	Posponeo	
Are you submitting this manuscript to a	No	
Full details of the experimental design and statistical methods used should be given in the Methods section, as detailed in our Minimum Standards Reporting Checklist. Information essential to interpreting the data presented should be made available in the figure legends.		
Have you included all the information requested in your manuscript?		
Resources	Yes	
A description of all resources used, including antibodies, cell lines, animals and software tools, with enough information to allow them to be uniquely identified, should be included in the Methods section. Authors are strongly encouraged to cite <u>Research Resource</u> <u>Identifiers</u> (RRIDs) for antibodies, model organisms and tools, where possible.		

Have you included the information requested as detailed in our <u>Minimum</u> <u>Standards Reporting Checklist</u> ?	
Availability of data and materials	Yes
All datasets and code on which the	
conclusions of the paper rely must be	
either included in your submission or	
deposited in publicly available repositories	
(where available and ethically	
appropriate), referencing such data using	
a unique identifier in the references and in	
the "Availability of Data and Materials"	
section of your manuscript.	
Have you have met the above	
requirement as detailed in our Minimum	
Standards Reporting Checklist?	

## Click here to view linked References

1	Genomic evic	lence of neo-sex chromosomes in the Eastern Yellow Robin	
2			
3	Han Ming Gan <sup>1,2</sup> *, Stephanie Falk <sup>3</sup> , Hernán E. Morales <sup>4</sup> , Christopher M. Austin <sup>1,2</sup> , Paul		
4	Sunnucks <sup>3</sup> , Alexandra Pavlova <sup>3*</sup>		
5			
6	<sup>1</sup> Centre for Integrative Ecology, School of Life and Environmental Sciences, Deakin University,		
7	Geelong, Vict	oria, Australia	
8	<sup>2</sup> Deakin Gene	omics Centre, Deakin University, Geelong, Victoria, Australia	
9	<sup>3</sup> School of Biological Sciences, Monash University, Clayton Campus, Clayton, Victoria, Australia		
10	<sup>4</sup> Centre for M	larine Evolutionary Biology, Department of Marine Sciences, University of	
11	Gothenburg, Göteborg, Sweden		
12			
13	* Correspond	ling author:	
14	Name:	Han Ming Gan, PhD	
15	Address:	Building KA4, School of Life and Environmental Sciences, Deakin University,	
16		Waurn Ponds, Victoria 3220, Australia	
17	Phone:	0490786277	
18	Email:	han.gan@deakin.edu.au	
19			
20	Name:	Alexandra Pavlova	
21	Address:	School of Biological Sciences	
22		Monash University, Clayton Campus	
23		Clayton Victoria 3800	
24	Phone:	0399055902	
25	Email:	alexandra.pavlova@monash.edu	
26			
27			
28			
29			
30			
31			
32			
33			
34			

Ŧ

- 35 Abstract
- 36

**Background**: Understanding sex-biased natural selection can be greatly enhanced by access to 37 well-annotated chromosomes including ones inherited in sex-specific fashions. The Eastern 38 39 Yellow Robin (EYR) is an endemic Australian songbird inferred to have experienced climatedriven sex-biased selection and is a prominent model for studying mitochondrial-nuclear 40 interactions in the wild. However, the lack of an EYR reference genome containing both sex 41 chromosomes (in birds, a female bearing Z and W chromosomes) is limiting current efforts to 42 43 understand the mechanisms of these processes. Here, we assemble the genome for a female EYR and use low depth (10  $\times$ ) genome resequencing data from 19 individuals of known sex to identify 44 chromosome fragments with sex-specific inheritance. 45

46

Findings: MaSuRCA hybrid assembly using Nanopore and Illumina reads generated a 1.22 Gb 47 EYR genome in 20,702 scaffolds (94.2% BUSCO completeness). Scaffolds were tested for W-48 linked (female-only) inheritance using a k-mer approach, and for Z-linked inheritance using 49 median read-depth test in male and female reads (read-depths must indicate haploid female and 50 diploid male representation). This resulted in 2,372 W-linked scaffolds (total length: 97,872,282 51 52 bp, N<sub>50</sub>: 81,931 bp) and 586 Z-linked scaffolds (total length: 121,817,358 bp, N<sub>50</sub>: 551,641 bp). Anchoring of the sex-linked EYR scaffolds to the reference genome of a female Zebra Finch 53 54 revealed two categories of sex-linked genomic regions. First, 653 W-linked scaffolds (25.7 Mb) were anchored to the W sex chromosome and 215 Z-linked scaffolds (74.4 Mb) to the Z. Second, 55 56 1138 W-linked scaffolds (70.9 Mb), and 179 Z-linked scaffolds (51.0 Mb), were anchored to a large section (coordinates ~5 to ~60 Mb) of Zebra Finch chromosome 1A. The first ~5 Mb and last 57 58 ~14 Mb of the reference chromosome 1A had only autosomally-behaving EYR scaffolds mapping 59 to them. 60

Conclusions: We report a female (W-chromosome containing) EYR genome and provide genomic
evidence for a neo-sex (neo-W and neo-Z) chromosome system in EYR, involving most of a large
chromosome (1A) previously only reported to be autosomal in passerines.

- 64
- 65

66 Keywords: Eastern Yellow Robin, *Eopsaltria australis*, passerine, songbird, genome, sex

67 chromosome, W-chromosome, neo-W, neo-Z

#### 68 Data description

69

Wildlife species that have genomic variation distributed heterogeneously through environmental 70 and geographic space can be excellent models for studying evolutionary processes under natural 71 72 conditions. Eastern Yellow Robin (EYR, NCBI:txid44318), Eopsaltria australis, is a common 73 endemic eastern Australian songbird (Figure 1) that shows geographically discordant patterns of 74 mitochondrial and nuclear genome variation. Whereas nuclear DNA variation in EYR is structured 75 mainly north-to-south, its two mitochondrial lineages (mitolineages) occur in contrasting climates 76 in an east-west (coast-to-inland) direction, with a narrow contact zone between them, despite ongoing male-mediated gene flow [1]. This pattern is inferred to have arisen when EYR 77 experienced two instances of climate-driven mitochondrial introgression into different nuclear 78 backgrounds: from the northern population into the southern through the inland, and from the 79 southern into the northern population along the coast [2]. Because mitogenome divergence is 80 mirrored by a fraction of the EYR nuclear genome that maps to the chromosome 1A of Zebra 81 Finch and is enriched for genes with mitochondrial functions, each inferred mitochondrial 82 83 introgression is hypothesized to have been accompanied by co-introgression of a co-evolved nuclear region [3]. Accordingly, the species has been highlighted as an exceptional model in the 84 85 emerging field of 'mitonuclear ecology', which addresses evolutionary interactions between mitochondrial and nuclear genomes and their products [4]. 86

87 Whereas progress on understanding mitonuclear interactions in EYR has been made by mapping genomic reads to a male Zebra Finch Taeniopygia guttata reference genome [5], the ~40 88 89 million years of evolution between the two species limits the assumptions that can be made about the degree of synteny of their genome organization. Moreover, the male reference lacks the 90 female-specific W chromosome in birds. Nuclear genomic architecture (for example, 91 concentrations of genes with mitochondrial functions that are subject to suppressed 92 93 recombination), has considerable potential to be a driver of mitonuclear evolution [6]. Furthermore, female-specific selection has been inferred for EYR, based on fine-scale spatial 94 95 separation of mitolineage distributions and their correlation with climate, despite male-biased gene flow in a species with female-biased dispersal [1]. Accordingly, genomic architecture with the 96 97 potential to impact the sexes differently could be a key player in mitonuclear evolution in this species. Thus to test among alternative hypotheses concerning mechanisms of potential co-98 evolution between elements of the nuclear genome and maternally-transmitted mtDNA, reference 99 sequences of both sex chromosomes are required. For example, the female-specific W-100 chromosome is necessarily co-inherited with mitochondrial DNA, and a species could experience 101

evolution so that W-chromosome bore genes relevant to mitochondrial function [1]. Substantial
female-specific gene regions are known from birds, notably neo-sex chromosome systems that can
provide females with gene sequences unavailable to males [7, 8].

Using a combination of Illumina and Nanopore reads, which have been shown to produce contiguous genome assemblies [9-12], we assembled a female inland EYR reference genome and utilized population genomic data from populations harbouring only inland mitochondrial lineages [13] to identify and annotate W and Z sex chromosomes. This procedure could also detect sexlinked chromosomes other than the typical W and Z avian sex chromosomes such as neo-sex chromosomes (caused by fusions between autosomal and sex chromosome elements) that are uncommon but known in birds, notably throughout the Sylvioidea, and in a honeyeater [7, 8, 14-

112

16]

113

#### 114 Sample collection, library construction and sequencing

115

Two EYR females, EYR054 and EYR056, were captured at Stuart Mill, western Victoria in the 116 same net on 6<sup>th</sup> of April 2009, as part of another project [17, 18]. DNA was extracted from 40 µL 117 of blood using a Qiagen DNAeasy Blood and Tissue Kit. A standard paired-end Illumina library 118 119 was constructed from 100 ng of QSonica-fragmented (~ 350 bp fragment size) EYR054 DNA using the NEBUltra Illumina Library Preparation kit (New England Biolabs, Ipwich, MA). The 120 121 library was quantified with a Tapestation 4000 (Agilent) and sequenced on the Novaseq6000 ( $2 \times$ 150 bp run configuration) at the Deakin Genomics Centre. Two Oxford Nanopore sequencing 122 123 libraries were constructed from G-tube fragmented (~8 kb) EYR054 gDNA using the LSK108 library preparation kit. Sequencing was performed on two MinION R9.4.1 flowcells for 48 hours 124 125 followed by fast5 base-calling using Albacore v2.0.1. A total of 6.63 Gb Nanopore data in 916,218 reads ( $N_{50} = 10,224$  bp) were generated after adapter-trimming using Porechop v0.2.3 126 (https://github.com/rrwick/Porechop). Nanopore reads used for this study had 13% error rate, 127 estimated based on mean pairwise sequence similarity of 87% (median= 89%) between Nanopore 128 reads and the assembled EYR genome, aligned using Minimap2 [19]. The DNA of EYR056 was 129 used to construct a mate-pair library with an insert size of 1 kb and sequenced by BGI for earlier 130 studies [18]. EYR054 is similar genetically to EYR056 according to whole mitogenomes, 131 microsatellites, and being female contemporaries in an area of the species' range where only the 132 inland mitolineage occurs, in an isolated habitat patch characterized by high local genetic 133 relatedness [3, 18, 20]. 134

For low (~10  $\times$ ) depth whole genome resequencing, 10 female and 9 male EYR individuals 135 bearing inland mitogenomes (EYR-A) were selected from northern (n=9) and southern (n=10) 136 populations [2, 13] away from the contact zone between the inland and coastal mitolineages 137 (Supplemental Table 1). Prior to Illumina sequencing, EYR individuals were genetically sexed 138 based on the intron length-variation of homologous sections of CHD (chromo-helicase-DNA-139 binding) genes located on W (female-limited) and Z (occurs in both sexes) chromosomes [21]. 140 These fragments have been sequenced previously for EYR for both sexes [1]. DNA extraction 141 from 16 blood samples and five tissues (Supplemental Table 1) was performed using a Qiagen 142 143 DNAeasy extraction kit. Illumina library construction and whole genome sequencing were performed at the Deakin Genomics Centre using the methods described above, generating an 144 average of 17 Gb (min = 12 Gb; max = 31 Gb) sequencing output per sample (Supplemental Table 145 146 1).

147

#### 148 Genome size estimation, hybrid *de novo* assembly and annotation

149

Raw Illumina EYR054 reads were poly-G, adapter- and quality trimmed using fastp v0.18.0 [22]. 150 The trimmed reads were used for genome profiling based on Jellyfish2-calculated k-mer frequency 151 152 (k=25) that estimated a genome size of 993 Mb with 1.12% heterozygosity for EYR054 (Figure 2A) [22-24]. We used MaSuRCA v3.2.4 (MaSuRCA, RRID:SCR\_010691) [25] to perform a 153 154 hybrid assembly of the EYR054 Nanopore and poly-G trimmed Illumina reads followed by gapclosing with Sealer v2.0.2 [26]. For the MaSuRCA assembly, Illumina reads were first error-155 156 corrected and used to construct contigs using the de Bruijn graph approach. These contigs were then used to error-correct the Nanopore long reads generating "mega reads" contigs and used for 157 158 Overlap-Layout-Consensus assembly. Subsequently, the MaSuRCA hybrid assembly was gapclosed with Sealer v2.0.2 using Illumina paired-end reads from the same individual. Given that 159 160 EYR056 and EYR054 are from the same population away from the hybrid zone (Harrisson et al. 2012, Morales et al. 2018) and thus likely possess similar versions of chromosomes, the EYR054 161 assembly was further scaffolded with mate-pair data from EYR056 using BESST [27] to generate 162 the final assembly for subsequent analyses (Table 1). Using mate-pair data improved the assembly 163 164 N<sub>50</sub> from 585 kb to 987 kb. The Sealer-gap-closed EYR054-only assembly is also made available in the GigaDB [28], should the future work on this species require single-individual assembly. 165 BUSCO v3 (BUSCO, RRID:SCR\_015008) [29] assessment of the assembled genome 166 based on the avian protein database (aves\_odb9), indicates 94.2% genome completeness with a 167 low level of duplicated genes (Table 1). Prior to gene prediction, the genome was masked for 168

- repeats using RepeatModeler (RepeatModeler, RRID:SCR\_015027) v1.0.11 and RepeatMasker
- 170 (RepeatMasker, RRID:SCR\_012954) v4.0.7 [30, 31]. The soft-masked genome (15.77% masked,
- 171 Table 1) along with the reference proteome of a male Collared Flycatcher [32] were used as the
- input for BRAKER2 annotation [33], resulting in the prediction of 23,905 genes. The Collared
- 173 Flycatcher proteome was used here in preference to Zebra Finch because the former has greater
- 174 protein similarity to EYR.
- 175

#### 176 Identification of sex chromosome scaffolds

177

Scaffolds inherited in sex-specific fashions ('sex-linked', 'W-linked' or 'Z-linked') were identified 178 using two methods (explained below) applied to sequence data obtained from 10 female and 9 179 male EYR individuals as detailed above. Paired-end reads for each re-sequenced male and female 180 were poly-G, quality- and adapter-trimmed using fastp (default setting) [22]. The trimmed reads 181 were mapped to the EYR genome using Bowtie2 (Bowtie, RRID:SCR\_005476) v2.3.4 [34]. High 182 mapping rates ranging from 97.82 to 98.53% were observed across all 19 individuals, indicating 183 robust assembly of the female EYR genome. The read mapping quality reported by Bowtie2 is 184 relatively constant (MapQ >30) across the assembly albeit with lower quality in the repetitive 185 186 regions as short reads will not be able to uniquely map to these regions. Subsequently, 90 million mapped PE reads were subsampled from each individual (to equalize coverage across individuals) 187 188 and used to estimate for each individual the median read-depth for each scaffold, and the fraction of the length of each scaffold that was covered by reads, using BAMStat04 as implemented in the 189 190 jvarkit package [35, 36].

Genome-wide identification of sex-linked scaffolds based on pooled male and female reads 191 192 could be compromised if any individuals had their sexes mis-assigned. Accordingly, to confirm the sex of the individual to which each set of sequence data was ascribed, the read-depth profiles for 193 194 all 19 EYR were assessed for the CHD sexing region noted above. BLASTN was used to align the CHD-W and CHD-Z nucleotide sequences (GenBank accession KC466840 - KC466844 CHD-W 195 and KC466845 - KC466853 CHD-Z) to two separate, long scaffolds (W chromosome scaffold: 196 QKXG01001703.1 - 310,213 bp; Z chromosome scaffold: QKXG01001459.1 - 211,357 bp). For 197 198 comparison, an autosomal scaffold, QKXG01002030.1 (3,864,097 bp) was identified that contained a fragment of the single-copy autosomal GAPDH (glyceraldehyde-3-phosphate 199 200 dehydrogenase) gene, sequenced previously for EYR (Genbank accession KC466694- KC466739) [1]. For the Z chromosome scaffold, a median read-depth centered on  $\sim 5 \times$  (haploid depth) was 201 observed in females, and  $\sim 10 \times$  (diploid) in males, while for the W chromosome fragment it was 202

 $\sim 5\times$  (haploid) in females and  $\sim 0\times$  (absent) in males;  $\sim 10\times$  diploid depth was observed for the autosomal scaffold in both sexes (Figure 2B).

BAM files from individual EYR were merged by sex using samtools v1.9 [37] to generate 205 one pooled alignment BAM file per sex. A histogram of read-depth frequency for each sex was 206 then generated using 'samtools depth' to estimate the read-depth cut-off for the identification of 207 candidate W- and Z-linked scaffolds (Figure 2C). The expected diploid depth for each sex was 208 209 estimated based on the peak observed read-depth (male =  $77\times$ ; female =  $83\times$ , Figure 2C). A minor peak corresponding to haploid read-depth ( $\sim 40 \times$ ) was observed for females but not males, 210 211 consistent with females being hemizygous for sex-linked regions (Figure 2C). A strong peak of low read-depth sequences ( $< 5 \times$ ) was seen only for males, consistent with them lacking a W 212 chromosome (Figure 2C). 213

To identify candidate W-linked scaffolds, we applied two established approaches with 214 complementary strengths that take advantage of sequence data being available for each sex. First, 215 we used a differential mapping approach, based on the expectation that a W-linked scaffold should 216 217 exhibit zero median read-depth in males, with a more than 75% of the scaffold having female reads mapping to it [7, 8]. Second, we used the YGS ('Y chromosome Genome Scan') k-mer approach, 218 designed for detecting W- or Y-linked regions [38]. The k-mer approach removes identical 219 220 repetitive sequences that might lead to false-positive matches to W-linked regions while retaining useful information from unique variants of repetitive regions: this is an advantageous attribute in 221 222 the face of the elevated repetitiveness expected of W chromosome sequences [38]. The k-mer approach was implemented as follows. For the pooled male reads, pooled female reads and the 223 224 female EYR genome assembly dataset, separate lists were built of all overlapping 16-bp sequences ('16-mers'): k=16 was chosen on the basis of genome size, and empirical validation that it 225 226 produced bimodal frequency distributions of *k*-mer presences in larger (>1 Gb) genomes [38]. Then, scaffolds from the assembled female genome are assumed to be W-linked if >75% of their 227 228 single-copy k-mers are absent in the pooled male reads but present in both of the female genome and pooled female reads. 229

Together, the two approaches identified 2,372 candidate W-linked scaffolds (total length of 97.87 Mb) that were used for downstream analyses. A great majority (1,952, 82.3 %, amounting to 86.32 Mb) of the candidate W-linked scaffolds were identified by both approaches, with 174 (7.3 %, 2.64 Mb) being exclusive to the *k*-mer approach, and 246 (10.4 %, 8.91 Mb) found only by the differential mapping approach. Inspection of the repetitiveness in the candidate W-linked scaffolds identified only by the *k*-mer approach indicates that they are 80% repetitive (total repeat length/total sequence length × 100%), consistent with the high sensitivity of *k*-mer approach in identifying repetitive sex-linked scaffolds [38]. In contrast, the candidate W-linked scaffolds found
by the differential mapping approach alone were only 32.6% repetitive.

Since Z-linked scaffolds are present in males and females, it is not possible to utilize the 239 YGS k-mer approach to identify candidates. Thus, we identified putative Z-linked scaffolds on the 240 basis of differences in read-depth between males and females, similar to the differential mapping 241 method for W-linked scaffold discovery outlined above. To allow for variation in sequencing 242 depth, we conservatively defined a candidate Z-linked scaffold as one exhibiting more than  $58 \times$ 243 median read-depth in males (i.e. 0.75 times the observed male diploid read-depth of 77×) and less 244 245 than 62× median read-depth in females (i.e. 1.5 times the observed female haploid read-depth of  $41.5\times$ ). Scaffolds passing these thresholds were further filtered to retain only those having both 246 male and female reads mapping to > 75% of the scaffold length. This resulted in the identification 247 of 586 candidate Z-linked scaffolds with a total length of 121.8 Mb and N<sub>50</sub> of 551.6 kb. 248

The total lengths of W-linked scaffolds (97.87 Mb) and Z-linked scaffolds = 121.82 Mb are much greater than expected from the typical sizes of sex chromosomes in Passerida, of which EYR is a member (e.g. in Xu *et al.* 2019 [39], Passerida W chromosomes range from 3.37-4.75 Mb and Z chromosomes range from 68.8-74.7 Mb) [39]. These observations raised the possibility of the presence of a neo-sex chromosome system, and hence it was of great interest to compare the sex-

linked regions to a well-annotated reference genome, as follows.

255

#### 256 Genomic evidence of neo-sex chromosomes in Eastern Yellow Robin

257

258 To assess the accuracy of our approaches for detecting sex-linked regions known in related

reference genomes, and to test for possible neo-sex chromosomes, the candidate W- and Z-linked
scaffolds were separately anchored to the female Zebra Finch genome (bTaeGut2:

261 <u>https://vgp.github.io/genomeark/Taeniopygia\_guttata/</u>, accessed on 19<sup>th</sup> December 2018) using

RaGoo v1.0 (with default settings) [40]. A total of 215 Z-linked scaffolds (74.4 Mb, ) were

anchored to the Zebra Finch Z chromosome, and 653 W-linked scaffolds (25.7 Mb) to the Zebra

Finch W chromosome. Surprisingly, a substantial proportion of candidate W-linked (n=1138, 70.9

Mb) and Z-linked (n=179, 51.0 Mb) scaffolds were also anchored to the autosomal Zebra Finch

chromosome 1A. Thus, each sex-linked scaffold anchored to one of three female Zebra Finch

chromosomes: W, Z or chromosome 1A. Using the entire EYR draft genome assembly as the

- 268 RaGoo input led to the anchoring of several W- and Z-linked scaffolds with the best hits to the
- 269 Zebra finch chromosome 1A, resulting in a substantially larger pseudomolecule for chromosome
- 1A (143.6 Mb), a length that is nearly double that of the Zebra Finch chromosome 1A (71.64 Mb)

which suggests the presence of two separate sex-linked versions of chromosome 1A in EYR (Fig.

- 3). By re-anchoring the EYR scaffolds in the absence of first the candidate W-linked and then the
- 273 candidate Z-linked scaffolds, two distinct versions of a chromosome 1A pseudomolecule were
- recovered that we designated putative neoZ-1A and neoW-1A chromosomes and used for
- 275 subsequent analyses.

To assess the robustness of the sex-based scaffold assignment approach, and to check the 276 sex-specific read-depth and length coverage along the putative neo-sex chromosomes involving 277 chromosome 1A (which we refer to as "pseudomolecules neoW-1A (Fig. 3 Chr1A: pink bar) and 278 279 neoZ-1A (Fig. 3 Chr1A: light blue bar)"), pooled female and male reads were mapped to the constructed EYR Z, W, autosomal chromosome 5, and neoZ-1A and neoW-1A pseudomolecules. 280 The mean read-depth in 100 kb non-overlapping sliding windows was calculated using the 281 'coverage' command in bedtool v2.25.0 [41] and visualized with ggplot2 in R v3.5.2 [42]. The 282 mean read-depth across the pseudomolecules was largely consistent with the scaffold sex-283 assignment i.e. zero depth for males and haploid for females for the W chromosome (Figure 4C) 284 and neoW-1A (Figure 4A), diploid depth for males and haploid for females for the Z chromosome 285 (Figure 4D) and neoZ-1A (Figure 4B), and diploid depth for both sexes for autosomal 286 chromosome 5 (Fig. 4E). In contrast to the W and Z chromosomes, several distinct genomic 287 288 regions with read-coverage consistent with that of an autosomal chromosome (Fig 4E) were observed for neoW-1A (Figure 4A) and neoZ-1A (Figure 4B), mostly at the pseudomolecule 289 290 termini.

291

#### 292 Identification of chromosome 1A-anchored gametologous gene pairs

293 Using FastANI, we calculated the pairwise sequence identity between the neoW-1A or 294 neoZ-1A pseudomolecule and the Zebra Finch chromosome 1A [43] and found that both exhibited substantial sequence similarity (calculated mean nucleotide identity of 86%) across the whole of 295 296 Zebra Finch chromosome 1A (Figure 5A, B). NeoW-1A exhibited ~ 20 Mb greater assembled length (92.5 Mb) than did neoZ-1A (72.5 Mb) (Figure 5 A, B). Accumulation of repeats 297 contributed to this: 36.6% of the EYR neoW-1A sequence was characterized as repetitive by 298 RepeatMasker, while this value is only 10% for the EYR neoZ-1A sequence. One expectation 299 under sex chromosome evolution is the development of evolutionary strata - regions of suppressed 300 recombination identified by spatial clusters of Z-W orthologs with similar divergence estimates 301 [44]. Accordingly, using FastANI, we calculated the pairwise sequence identity between EYR 302 neoW-1A and neoZ-1A in a non-overlapping sliding window of 10 kb. By aligning the putative 303 neoW-1A to the neoZ-A, we observed high (mostly >90%) pairwise sequence identity throughout 304

the pseudomolecule (Figure 5C). However, there was considerable heterogeneity in absolute
sequence similarity, with zones of ~100 %, ~98 %, ~95 %, and ~92 % identity clumped along the
pseudomolecules, suggestive of evolutionary strata (Figure 5C) [44].

Orthologous genes shared between the EYR and Collared Flycatcher (higher protein 308 similarity to EYR compared to Zebra Finch) were inferred using OrthoFinder2 [45]. Of the 957 309 genes located on the Collared Flycatcher chromosome 1A, 725 formed a one-to-one (n=513) or 310 one-to-many (n=212) orthologous group with the EYR genes located on the neoZ-1A or neoW-1A 311 pseudomolecule. We restricted the ortholog analysis to only genes predicted from the sex-linked 312 313 scaffolds (identified based on EYR scaffold assignment) since the "autosomal-behaving" scaffolds on the neoZ-1A and neoW-1A pseudomolecules (Figure 4A and B) may consist of unassigned sex-314 linked, recombining sex-linked (collapsed into a single scaffold) or truly autosomal scaffolds that 315 will affect gametologous pairing. This resulted in the identification of 419 Z-linked genes on EYR 316 neoZ-1A pseudomolecule and 221 W-linked orthologs on neoW-1A, to a total of 488 different sex-317 linked genes. Among these were 148 putative gametologous gene pairs (i.e. homologs with 318 319 sufficiently low recombination for one version to be identifiably W- and one Z-linked) between EYR neoW- and neoZ-1A (Supplemental Table 2). The fewer number of W-linked EYR genes that 320 formed orthologous group with the Collared Flycatcher chromosome 1A genes compared to that of 321 322 Z-linked EYR genes may be due to the more fragmented assembly and higher repetitiveness of Wlinked EYR scaffolds that precludes the accurate annotation of genes in the W-linked scaffolds 323 324 when using the default BRAKER2 annotation settings [46]. It is also possible that W-linked EYR genes on chromosome 1A have diverged, been lost or degraded beyond detection, as expected 325 326 under sex chromosome evolution [44].

Neo-sex chromosomes have reduced effective population size relative to the autosomes that 327 328 contribute to them: this is expected to decrease the effectiveness of purifying selection, especially when compounded by reduced recombination [47-50]. These effects, along with Mueller's ratchet 329 330 and hitchhiking, should promote the accumulation of deleterious mutations, commonly revealed as elevated non-synonymous to synonymous (dN/dS) ratios, particularly for sex-limited 331 chromosomes such as the neo-W in birds [7, 8, 51]. To calculate the dN/dS ratios for EYR neo-sex 332 gametologous gene pairs, protein alignment was first performed for all 148 putative neo-sex 333 334 gametologous gene pairs with their respective Collared Flycatcher orthologs using Clustal Omega v1.2.1 [52] followed by codon-based alignment with pal2nal (-nogap option to remove gaps and 335 inframe stop codons) [53]. The pal2nal output for each orthologous group was used to calculate 336 dN/dS ratios via codeml in paml v4.9i package [54]. When the Collared Flycatcher orthologous 1A 337 genes were used as the reference for each comparison, 120 out of 148 the neoW-1A gametologs 338

exhibited higher dN/dS than their neoZ-1A gametologous partners (Wilcoxon paired samples
signed rank test, p = 3.9e-14; Fig. 5 D).

341

#### 342 Conclusion

343

We report a hybrid genome assembly using Nanopore and Illumina reads of a female Eastern 344 Yellow Robin (EYR), the first published genome for the family Petroicidae. The identification of 345 sex-linked scaffolds using a combination of read-depth and k-mer YGS approaches, followed by 346 347 chromosomal anchoring to the genome of a female Zebra Finch, provided strong evidence for the presence of a neo-sex chromosome system in EYR involving most of chromosome 1A. The 348 inferred neoW-1A pseudomolecule showed the characteristics expected of a sex-limited neo-sex 349 chromosome, including elevated dN/dS ratios, increased levels of repetitive sequences and signals 350 of strata of levels of sequence divergence [7, 8, 51]. Further work is required to understand the 351 formation of the neo-sex system we infer. One relatively simple model is that one copy of 352 353 chromosome 1A fused with the W chromosome, and the second copy of chromosome 1A became 354 inherited in a neo-Z fashion; but more complicated scenarios are possible [7, 8, 15]. Given that the divergence between inland and coastal EYR lineages is partly due to a genomic region enriched for 355 356 nuclear genes with mitochondrial functions that maps to the autosomal chromosome 1A in other songbirds [3] (Morales et al. 2018), the role of neo-sex chromosomes in maintaining lineage 357 358 divergence despite nuclear gene flow warrants further investigation involving a female genome of the coastal lineage. Future work should also test whether unlikely but possible neoZ-1A 359 360 differences between the EYR054 used for assembly here and EYR056 used for scaffolding affected the neo-Z assembly. Chromosome 1A is not one of the chromosomes implicated in 361 362 multiple known vertebrate neo-sex systems [51], but given its unusual concentration of genes with mitochondrial functions, it will not be surprising if subsequent equivalent cases are found [3]. Our 363 results show that assuming close synteny between a songbird of interest and a distantly-related 364 reference genome can lead to incomplete or incorrect evolutionary inferences. The present genome 365 assembly will be an important molecular resource for understanding and re-evaluating genome 366 evolution in EYR, a key model wildlife species in the emerging field of 'mitonuclear ecology' [4]. 367 The discovery of neo-sex chromosomes in this bird species adds another, independent model to the 368 limited number in which sex chromosome evolution can be studied through the lens of relatively 369 370 young sex chromosomes [15, 51].

#### 371 Availability of supporting data

372	The genome assembly has been deposited in the NCBI database with the accession number
373	QKXG010000000. Raw sequencing data have been deposited in the NCBI Sequence Read Archive
374	(SRA) database and linked to the Bioproject ID PRJNA476023. Additional supporting data,
375	including the initial EYR054 MaSuRCA genome assembly (prior to scaffolding with EYR056
376	mate-pair data), BUSCO calculations, genome annotations, candidate W- and Z-linked sequences,
377	RaGOO scaffolding output and dN/dS ratio calculations are available via the GigaScience
378	database, GigaDB [28].
379	
380	Competing interests
381	The authors declare that they have no competing interests.
382	
383	Funding
384	This study was funded by the Monash School of Biological Sciences, Australian Research Council
385	(ARC) Discovery Project (DP180102359), ARC Linkage Grant (LP0776322), the Victorian
386	Department of Sustainability and Environment (DSE), Museum of Victoria, Victorian Department
387	of Primary Industries, Parks Victoria, North Central Catchment Management Authority, Goulburn
388	Broken Catchment Management Authority, CSIRO Ecosystem Sciences, and the Australian
389	National Wildlife Collection Foundation.
390	
391	Authors' contributions
392	HMG, PS and AP conceived the study. HEM and SF collected the samples and extracted the
393	genomic DNA. CMA contributed sequencing reagents and computing resources. HMG performed
394	whole-genome sequencing, genome assembly, genome annotation and comparative genomics
395	analysis. HEM assessed the assembly quality and genomics analysis. HMG, AP and PS wrote the
396	manuscript. All authors read, edited and approved the final manuscript.
397	
398	Acknowledgments
399	We thank Hans Zauner, Jason Sardell, Leonardo Campagna and Qi Zhou for comments that helped
400	improve the manuscript. Thanks to Nevil Amos, Lana Austin and all other collectors of specimens
401	and all agencies who granted permission to collect specimens. Field samples were collected under
402	scientific research permits issued by the Victorian Department of Environment and Primary

- 403 Industries (numbers 10007165, 10005919 and 10005514), New South Wales Office of
- 404 Environment and Heritage (SL100886), in accordance with Animal Ethics approvals AM13-05,
- 405 BSCI\_2012\_20 and BSCI\_2007\_07, using bands issued by the Australian Bird and Bat Banding

406 Scheme. We also thank the Australian National Wildlife Collection, Leo Joseph, Robert Palmer

and Richard Major for providing genetic samples. We are grateful to Erich Jarvis and Olivier

408 Fedrigo for giving us access to the female Zebra Finch assembly. We thank Gabriel Low for

409 helpful discussion.

410

411 Figure legends

412

413 Figure 1. The Eastern Yellow Robin. Photo by Geoff Park

414

415 Figure 2. Genomic profiling and *in-silico* sexing of Eastern Yellow Robin. (A) Genomescope

416 profile calculated from trimmed Illumina data of EYR054 using a *k*-mer length of 25. (B) Median

417 coverage per individual for three sets of scaffolds with different inheritance, for the female sample

418 (n=10) and male sample (n=9), with individuals sequenced at ~  $10 \times$  coverage each. Autosomal =

419 GAPDH-containing chromosome fragment [1] scaffold QKXG0002030; W = W-chromosome

420 fragment: scaffold QKXG0001703; Z = Z chromosome fragment: scaffold QKXG0001459. (C)

421 Frequency distribution of base-by-base read-depth calculated from the mapping of pooled male

422 (blue line) and female (red line) reads to the female genome assembly. This is subsequently used to

423 estimate the read-depth of haploid and diploid scaffolds.

424

Figure 3. The assembled lengths of Eastern Yellow Robin (EYR) chromosome 1A, W and Z

426 pseudomolecules constructed by anchoring different scaffold inputs to the female Zebra Finch

427 reference genome (ZF; grey bars). Inputs included: EYR genome (EYR; yellow bars); EYR

428 candidate W-linked scaffolds (EYR W-linked; red bars); EYR candidate Z-linked scaffolds (EYR

429 Z-linked; dark blue bars); EYR genome without Z-linked scaffolds (EYR w/o Z-linked; pink bars);

430 EYR genome without W-linked scaffolds (EYR w/o W-linked; light blue bars). Neo-sex-

431 chromosome pseudomolecules were built using the latter two datasets (the length of neoW-1A is

432 shown by the light blue bar and that of neoZ-1A by the light pink bar for Ch1A).

433

Figure 4. Read-depth of pooled male (blue dots) and pooled female (red dots) reads across EYR

435 pseudomolecules neoW-1A, neoZ-1A, W, Z and autosomal chromosome 5. Read-depth (the

436 number of reads for each nucleotide in the genome) was estimated for each 100 kb sliding window.

437 The locations of candidate W-linked scaffolds on the neoW-1A and Z-linked scaffolds on neoZ-1A

438 pseudomolecules are indicated by the black lines below the read-depth plots. Coordinate (Mb)

439 refers to the position on the pseudomolecule.

441	Figure 5. Characterization of the inferred neo-sex chromosomes in Eastern Yellow Robin. Linear
442	genome comparison of the (A) neoW-1A and (B) neoZ-1A pseudomolecules (blue horizontal bars)
443	with the Zebra Finch chromosome 1A (green horizontal bars). The neoW-1A alignment is $\sim 20$ Mb
444	longer than that of neoZ-1A. The red lines denote regions of nucleotide similarity with more than
445	70% nucleotide identity calculated over 10 kb non-overlapping sliding window. (C) Pairwise
446	sequence identity per 10 kb sliding window (to obtain high resolution) between the neoW-1A and
447	neoZ-1A scaffolds mapped along the neoZ-1A pseudomolecule, with coordinates relating to the
448	neoZ-1A pseudomolecule. Zones of different levels of sequence similarity can be seen along the
449	pseudomolecule. The blue line denotes the smoothed conditional means for pairwise identity and
450	the grey zone around it indicates the 95% confidence interval. (D) Paired box plots showing the
451	dN/dS ratios of neoW-1A and neoZ-1A gene copies (gametologs) of the Eastern Yellow Robin
452	compared with Collared Flycatcher orthologs as references. Collared Flycatcher was used here in
453	preference to Zebra Finch because the former has greater protein similarity to EYR. Grey lines
454	connecting red and blue dots represent gametologs.

## **Table 1**

Parameter	Details
Organism	Eopsaltria australis (Eastern Yellow Robin)
Isolate	EYR054 (sex = female; data type: Illumina
	standard paired-end and Nanopore long read)
	EYR056 (sex = female; data type: Illumina mate-
	pair)
Bioproject	PRJNA476023
Biosample	SAMN09425179 (isolate EYR054)
	SAMN10581952 (isolate EYR056)
GenBank assembly accession	GCA_003426825.1 (QKXG01)
Assembled Length	1,228,344,903 bp
Scaffold N <sub>50</sub>	987,278 bp

457 Table 1. Genome assembly and annotation statistics of the Eastern Yellow Robin

Number of scaffolds	20,702
Number of predicted protein-coding	23,905
genes	

# **Repeat Annotation:**

LINEs	39,888,415 bp (3.25%)
LTR elements	85,519,635 (6.96%)
DNA elements	6,416,492 (0.52%)
Unclassified repeats	42,749,317 (3.48%)
Satellites	1,967,923 (0.16%)
Simple repeats	14,300,770 (1.16%)
Low complexity	3,128,912 (0.25%)

<b>BUSCO completeness (Ave odb9)</b>	Whole genome	Predicted proteome
Complete BUSCO	4,627 (94.2%)	3,795 (77.2%)
Complete and single-copy BUSCO	4,436 (90.3%)	3,302 (67.2%)
Complete and duplicated BUSCO	191 (3.9%)	493 (10.0%)
Fragmented BUSCO	163 (3.3%)	590 (12.0%)
Missing BUSCO	125 (2.5%)	530 (10.8%)
Total BUSCO groups search	4,915	4,915

#### **References**

461	1.	Pavlova A, Amos JN, Joseph L, Loynes K, Austin JJ, Keogh JS, et al. Perched at the mito-nuclear
462		crossroads: divergent mitochondrial lineages correlate with environment in the face of ongoing
463		nuclear gene flow in an Australian bird. Evolution. 2013;67 12:3412-28.
464	2.	Morales HE, Sunnucks P, Joseph L and Pavlova A. Perpendicular axes of differentiation generated
465		by mitochondrial introgression. Molecular ecology. 2017;26 12:3241-55.

466 3. Morales HE, Pavlova A, Amos N, Major R, Kilian A, Greening C, et al. Concordant divergence of 467 mitogenomes and a mitonuclear gene cluster in bird lineages inhabiting different climates. Nature 468 ecology & evolution. 2018;2 8:1258. 4. Hill GE. Mitonuclear ecology. Oxford, United Kingdom: Oxford University Press; 2019. 469 470 5. Warren WC, Clayton DF, Ellegren H, Arnold AP, Hillier LW, Künstner A, et al. The genome of a 471 songbird. Nature. 2010;464:757. doi:10.1038/nature08819 https://www.nature.com/articles/nature08819#supplementary-information. 472 473 Sunnucks P, Morales HE, Lamb AM, Pavlova A and Greening C. Integrative approaches for studying 6. 474 mitochondrial and nuclear genome co-evolution in oxidative phosphorylation. Frontiers in 475 genetics. 2017;8:25. 476 Dierickx E, Sin S, van Veelen P, Brooke MdL, Liu Y, Edwards S, et al. Neo-sex chromosomes and 7. 477 demography shape genetic diversity in the Critically Endangered Raso lark. bioRxiv. 2019:617563. 478 doi:10.1101/617563. 479 Leroy T, Anselmetti Y, Tilak M-K, Bérard S, Csukonyi L, Gabrielli M, et al. A bird's white-eye view on 8. 480 neosex chromosome evolution. bioRxiv. 2019:505610. doi:10.1101/505610. 481 9. Austin CM, Tan MH, Harrisson KA, Lee YP, Croft LJ, Sunnucks P, et al. De novo genome assembly 482 and annotation of Australia's largest freshwater fish, the Murray cod (Maccullochella peelii), from 483 Illumina and Nanopore sequencing read. Gigascience. 2017;6 8:1-6. 484 10. Tan MH, Austin CM, Hammer MP, Lee YP, Croft LJ and Gan HM. Finding Nemo: hybrid assembly 485 with Oxford Nanopore and Illumina reads greatly improves the clownfish (Amphiprion ocellaris) 486 genome assembly. GigaScience. 2018;7 3:gix137. 487 Zimin AV, Puiu D, Luo M-C, Zhu T, Koren S, Marçais G, et al. Hybrid assembly of the large and 11. 488 highly repetitive genome of Aegilops tauschii, a progenitor of bread wheat, with the MaSuRCA mega-reads algorithm. Genome research. 2017. 489 490 Zimin AV, Puiu D, Hall R, Kingan S, Clavijo BJ and Salzberg SL. The first near-complete assembly of 12. 491 the hexaploid bread wheat genome, Triticum aestivum. Gigascience. 2017;6 11:1-7. 492 doi:10.1093/gigascience/gix097. 493 13. Morales HE, Pavlova A, Joseph L and Sunnucks P. Positive and purifying selection in mitochondrial 494 genomes of a bird with mitonuclear discordance. Mol Ecol. 2015;24 11:2820-37. 495 doi:10.1111/mec.13203. 496 14. Sardell J. Evolutionary Consequences of Recent Secondary Contact Between Myzomela 497 Honeyeaters. 2016. 498 15. Pala I, Naurin S, Stervander M, Hasselquist D, Bensch S and Hansson B. Evidence of a neo-sex 499 chromosome in birds. Heredity (Edinb). 2012;108 3:264-72. doi:10.1038/hdy.2011.70. 500 16. Brooke Mde L, Welbergen JA, Mainwaring MC, van der Velde M, Harts AM, Komdeur J, et al. 501 Widespread translocation from autosomes to sex chromosomes preserves genetic variability in an 502 endangered lark. Journal of molecular evolution. 2010;70 3:242-6. doi:10.1007/s00239-010-9333-503 3. 504 17. Harrisson KA, Pavlova A, Amos JN, Takeuchi N, Lill A, Radford JQ, et al. Fine-scale effects of habitat 505 loss and fragmentation despite large-scale gene flow for some regionally declining woodland bird 506 species. Landscape Ecology. 2012;27 6:813-27. doi:10.1007/s10980-012-9743-2. 507 18. Morales HE, Pavlova A, Sunnucks P, Major R, Amos N, Joseph L, et al. Neutral and selective drivers 508 of colour evolution in a widespread Australian passerine. Journal of biogeography. 2017;44 3:522-509 36. 510 19. Li H. Minimap2: pairwise alignment for nucleotide sequences. Bioinformatics. 2018;34 18:3094-511 100. doi:10.1093/bioinformatics/bty191. 512 20. Amos JN, Harrisson KA, Radford JQ, White M, Newell G, Nally RM, et al. Species- and sex-specific 513 connectivity effects of habitat fragmentation in a suite of woodland birds. Ecology. 2014;95 514 6:1556-68. doi:10.1890/13-1328.1. 515 21. Griffiths R, Double MC, Orr K and Dawson RJ. A DNA test to sex most birds. Molecular ecology. 516 1998;7 8:1071-5.

517 22. Chen S, Zhou Y, Chen Y and Gu J. fastp: an ultra-fast all-in-one FASTQ preprocessor. 518 Bioinformatics. 2018;34 17:i884-i90. doi:10.1093/bioinformatics/bty560. Marçais G and Kingsford C. A fast, lock-free approach for efficient parallel counting of occurrences 519 23. 520 of k-mers. Bioinformatics. 2011;27 6:764-70. doi:10.1093/bioinformatics/btr011. 521 24. Vurture GW, Sedlazeck FJ, Nattestad M, Underwood CJ, Fang H, Gurtowski J, et al. GenomeScope: 522 fast reference-free genome profiling from short reads. Bioinformatics. 2017;33 14:2202-4. 523 25. Zimin AV, Marçais G, Puiu D, Roberts M, Salzberg SL and Yorke JA. The MaSuRCA genome 524 assembler. Bioinformatics. 2013;29 21:2669-77. 525 26. Paulino D, Warren RL, Vandervalk BP, Raymond A, Jackman SD and Birol I. Sealer: a scalable gap-526 closing application for finishing draft genomes. BMC bioinformatics. 2015;16 1:230. 527 27. Sahlin K, Chikhi R and Arvestad L. Assembly scaffolding with PE-contaminated mate-pair libraries. Bioinformatics. 2016;32 13:1925-32. 528 529 28. Gan HM, Falk S, Morales HE, Austin CM, Sunnucks P and Pavlova A. Supporting data for "Genomic 530 evidence of neo-sex chromosomes in the Eastern Yellow Robin". GigaScience Database. 2019. 531 http://dx.doi.org/10.5524/100639 532 29. Waterhouse RM, Seppey M, Simão FA, Manni M, Ioannidis P, Klioutchnikov G, et al. BUSCO 533 applications from quality assessments to gene prediction and phylogenomics. Molecular biology 534 and evolution. 2017;35 3:543-8. 535 30. Smit AF, Hubley R and Green P. RepeatMasker. 1996. 536 31. Smit A and Hubley R. RepeatModeler Open-1.0. Available fom http://wwwrepeatmaskerorg. 2008. 537 32. Ellegren H, Smeds L, Burri R, Olason PI, Backström N, Kawakami T, et al. The genomic landscape of 538 species divergence in Ficedula flycatchers. Nature. 2012;491 7426:756. 539 33. Hoff KJ, Lange S, Lomsadze A, Borodovsky M and Stanke M. BRAKER1: unsupervised RNA-Seq-540 based genome annotation with GeneMark-ET and AUGUSTUS. Bioinformatics. 2015;32 5:767-9. 541 34. Langmead B and Salzberg SL. Fast gapped-read alignment with Bowtie 2. Nature methods. 2012;9 542 4:357. 543 Lindenbaum P. JVarkit: java-based utilities for Bioinformatics. 2015. 35. 544 36. Lindenbaum P and Redon R. bioalcidae, samjs and vcffilterjs: object-oriented formatters and filters 545 for bioinformatics files. Bioinformatics. 2018;34 7:1224-5. doi:10.1093/bioinformatics/btx734. 546 37. Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, et al. The Sequence Alignment/Map 547 format and SAMtools. Bioinformatics. 2009;25 16:2078-9. doi:10.1093/bioinformatics/btp352. 548 38. Carvalho AB and Clark AG. Efficient identification of Y chromosome sequences in the human and 549 Drosophila genomes. Genome research. 2013;23 11:1894-907. doi:10.1101/gr.156034.113. 550 39. Xu L, Auer G, Peona V, Suh A, Deng Y, Feng S, et al. Dynamic evolutionary history and gene content 551 of sex chromosomes across diverse songbirds. Nat Ecol Evol. 2019;3 5:834-44. 552 doi:10.1038/s41559-019-0850-1. Alonge M, Soyk S, Ramakrishnan S, Wang X, Goodwin S, Sedlazeck FJ, et al. Fast and accurate 553 40. 554 reference-guided scaffolding of draft genomes. bioRxiv. 2019:519637. doi:10.1101/519637. 555 Quinlan AR and Hall IM. BEDTools: a flexible suite of utilities for comparing genomic features. 41. 556 Bioinformatics. 2010;26 6:841-2. doi:10.1093/bioinformatics/btq033. 557 42. Wickham H. ggplot2: Elegant Graphics for Data Analysis. Springer Publishing Company, 558 Incorporated; 2009. 559 43. Jain C, Rodriguez-R LM, Phillippy AM, Konstantinidis KT and Aluru S. High throughput ANI analysis 560 of 90K prokaryotic genomes reveals clear species boundaries. Nature communications. 2018;9 561 1:5114. 562 44. Wright AE, Dean R, Zimmer F and Mank JE. How to make a sex chromosome. Nature 563 communications. 2016;7:12087. Emms DM and Kelly S. OrthoFinder: solving fundamental biases in whole genome comparisons 564 45. 565 dramatically improves orthogroup inference accuracy. Genome biology. 2015;16 1:157. 566 46. Rutkowska J, Lagisz M and Nakagawa S. The long and the short of avian W chromosomes: no 567 evidence for gradual W shortening. Biology letters. 2012;8 4:636-8. doi:10.1098/rsbl.2012.0083.

- 56847.Smeds L, Warmuth V, Bolivar P, Uebbing S, Burri R, Suh A, et al. Evolutionary analysis of the569female-specific avian W chromosome. Nature communications. 2015;6:7330.
- 570 48. Charlesworth B. The effect of background selection against deleterious mutations on weakly
  571 selected, linked variants. Genetical research. 1994;63 3:213-27.
- 49. Charlesworth B and Charlesworth D. The degeneration of Y chromosomes. Philos Trans R Soc Lond
  573 B Biol Sci. 2000;355 1403:1563-72. doi:10.1098/rstb.2000.0717.
- 57450.Bachtrog D. A dynamic view of sex chromosome evolution. Current opinion in genetics &575development. 2006;16 6:578-85. doi:10.1016/j.gde.2006.10.007.
- 576 51. Sigeman H, Ponnikas S, Videvall E, Zhang H, Chauhan P, Naurin S, et al. Insights into Avian
  577 Incomplete Dosage Compensation: Sex-Biased Gene Expression Coevolves with Sex Chromosome
  578 Degeneration in the Common Whitethroat. Genes. 2018;9 8 doi:10.3390/genes9080373.
- 579 52. Sievers F and Higgins DG. Clustal Omega, accurate alignment of very large numbers of sequences. 580 Multiple sequence alignment methods. Springer; 2014. p. 105-16.
- 581 53. Suyama M, Torrents D and Bork P. PAL2NAL: robust conversion of protein sequence alignments 582 into the corresponding codon alignments. Nucleic acids research. 2006;34 suppl\_2:W609-W12.
- 583 54. Yang Z. PAML 4: phylogenetic analysis by maximum likelihood. Molecular biology and evolution.
- 584 2007;24 8:1586-91.

585













Supplementary Table 1

Click here to access/download Supplementary Material SupplementalTable1.xlsx Supplementary Table 2

Click here to access/download Supplementary Material SupplementalTable2\_revised.xlsx