

## Electronic Supplementary Materials

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## 3 Repeated sex chromosome evolution in vertebrates supported 4 by expanded avian sex chromosomes

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39 **Supplementary Methods S1**

40 **S1a. DNA extraction and sequencing**

41 DNA from the blood samples was extracted from each sample ( $n = 8$ ) using a phenol–chloroform protocol  
42 [1]. The extracted DNA was sequenced with Illumina HiSeqX (150 bp, paired-end) by SciLifeLab Sweden.

43 **S1b. *De novo* assembly and mapping**

44 To obtain reference genomes, we created *de novo* genome assemblies from the male sequence data for each  
45 of the study species. The resequencing data was trimmed using nesoni clip (<https://github.com/Victorian->  
46 Bioinformatics-Consortium/nesoni) with a minimum read quality of 20 and minimum read length of 20 bp.  
47 The trimmed reads were then assembled with Spades v3.5.0 [2] using 5 different kmer lengths (21, 33, 55,  
48 77 and 127) and the setting “careful”. Scaffolds shorter than 1 kbp were discarded. Quality statistics from  
49 the assemblies were calculated using Quast v4.5.4 [3]. Samples from all species were aligned to their  
50 corresponding reference genome. However, based on results from downstream analyses (S1d; figure S2),  
51 two of the reference genomes (the horned lark and the bearded reedling) were discarded from the final  
52 analyses, and the samples from these species were instead analysed using the reference genome of their  
53 closest relative, the Raso lark. This decision was made as the level of heterozygosity in these two species  
54 influenced the relative mapping success between the female and male sample on a genome-wide scale, as  
55 the reference genome was based on the same male sample, thus not revealing the sex-linked genomic regions  
56 as clearly as when the samples were aligned to the reference genome of their relative, the Raso lark (see  
57 figure S2). This strategy meant keeping the assemblies of the Raso lark ( $N_{50}=103$  kbp, 28304 scaffolds) and  
58 the bearded reedling ( $N_{50}=68$  kbp, 36455 scaffolds), while the assemblies for the Eurasian skylark ( $N_{50}=8$

59 kbp, 256822 scaffolds) and horned lark (N50=22 kbp, 109088 scaffolds) were discarded. Genome assembly  
60 statistics are given in electronic supplementary material, table S1.

61           Reads from all of the samples (n = 8) were cleaned for adaptor sequences with Trimmomatic  
62 v.0.3.6 [4] using the adaptor file TruSeq3-PE and options seedMismatches: 2, palindromeClipThreshold: 30  
63 and simpleClipThreshold: 10. Trimming of low-quality bases was done using a quality threshold of 15 from  
64 the leading end and 30 from the trailing end. The reads were further trimmed for a minimum quality of 20  
65 over sliding windows of 4 bp. Lastly, any reads shorter than 90 bp were excluded from further analyses. The  
66 number of remaining reads in our samples ranged from 176 to 404 million. The samples were quality  
67 checked using fastqc v0.11.7 (<https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>).

68           The male and female samples of Raso lark and bearded reedling were aligned to their  
69 respective genome assemblies, while the Eurasian skylark and horned lark individuals were aligned to the  
70 genome assembly of their closest relative, the Raso lark. The alignment was done with bwa mem v0.7.17  
71 [5], marking shorter split hits as secondary (option -M) for downstream compatibility. The aligned reads  
72 were sorted with samtools v1.7 [6], and duplicated reads were removed using picardtools v2.18.0  
73 (<http://broadinstitute.github.io/picard>).

74           Assembly and alignment statistics are provided in electronic supplementary material, table  
75 S1 and table S2.

### 76 **S1c. Chromosome anchoring**

77 The scaffolds in the genome assemblies were grouped into different chromosomes and ordered into  
78 chromosome-level using the genome assembly of the zebra finch (*Taeniopygia guttata*). The zebra finch  
79 genome assembly (taeGut.3.2.4 [7]) was downloaded from Ensembl [8] and transformed into a database  
80 using the last v876 [9] program lastdb. The two genome assemblies, of the Raso lark and bearded reedling,  
81 were aligned to the zebra finch genome using the program lastal and converted to psl format using the script  
82 maf-convert, both from the same software suite last v876. From there, we extracted chromosome anchoring  
83 coordinates based on the longest match to the zebra finch genome for each 5 kbp window in the Raso lark  
84 and bearded reedling assemblies, with a minimum requirement of 500 matching base pairs per 5 kbp

85 window. The assembly positions from the output files of the coverage and single nucleotide variant (SNV)  
86 analyses were then translated to the starting positions of the match to the zebra finch genome assembly (see  
87 section below). With this method, between 81 and 95% of the scaffolds larger than 5 kbp were anchored to  
88 the zebra finch reference genome, with the largest proportion aligning to the Raso lark (95%) and bearded  
89 reedling (94%). See statistics in table S3.

90 **S1d. Identification of sex-linked genomic regions**

91 We identified sex-linked regions using two different kinds of genomic signatures: (i) differential mapping  
92 success in males and females (i.e. sex-specific genome coverage), and (ii) an excess or deficit of female-  
93 specific genetic variation. Sex chromosomes almost invariably evolve recombination suppression in the  
94 heterogametic sex so that regions that have been sex-linked for a long time (such as the sex chromosomes  
95 that formed in the ancestor of all birds) will show pronounced sequence divergence and degeneration in the  
96 non-recombining chromosome (the W in birds [10]). The regions belonging to the ancestral sex  
97 chromosomes can thus be identified by lower female coverage, and fewer female-specific genetic variants  
98 compared to males. This is because reads from the female-specific W-chromosome will either not map to  
99 the male reference genome due to substantial differentiation between the Z and W or because of deletions  
100 on the W chromosome. More recently formed sex-linked regions may be identified by lower mapping  
101 success in females, although we expect a subtler difference as the W-linked genomic region may not have  
102 yet developed substantial differentiation from the Z-linked homologous region. Here, we also expect a  
103 higher amount of female-specific mutations compared to males, due to differentiation between the Z and  
104 W sex chromosome copies.

105 To uncover differential mapping success in males and females for each species, we calculated  
106 genome-wide coverage for 5 kbp windows with bedtools v2.71.1 [11] from alignment files with increasingly  
107 strict settings for maximum allowed mismatches between the samples and the reference genome, allowing  
108 a maximum of (i) 4, (ii) 3, (iii) 2, (iv) 1 and lastly (v) 0 mismatches. All genome coverage values were  
109 normalised between the female and the male sample by dividing the median female-to-male coverage ratio  
110 for each 5 kbp window by the genome-wide median female-to-male coverage ratio. We also excluded

111 windows with a read count >1500 in either sample or a coverage ratio >2. Both of these thresholds were  
112 set to be well outside the normal distribution for all species and was done in order to filter out repeat regions  
113 which may have a large effect on the differences between the two samples. To identify sex-linked regions,  
114 we binned the female-to-male coverage ratio values for every 1 Mbp genomic region of chromosomes larger  
115 than 1 Mbp and extracted the mean value from each bin. Results from all maximum mismatches settings  
116 were visually inspected and we decided on a cut-off of maximum 2 mismatches, which best reveals the sex-  
117 linked regions in the data. We then binned the data into 0.1 Mbp windows and calculated the mean female-  
118 to-male coverage ratio within each bin.

119 To analyse female-specific variation, we called variants in the alignment files (with all  
120 mismatches allowed) with freebayes v1.1.0 for each species separately ( $n = 2$  in each analysis) using  
121 freebayes-parallel [12] and parallel v20180322 [13]. The output was then parsed for any SNV that had been  
122 marked with a flag other than PASS (--remove-filtered-all), a minimum quality of 20 and minimum depth  
123 of 3x using vcftools v0.1.15 [14]. Private alleles (minor alleles occurring only in one sample in a heterozygous  
124 state) were extracted with vcftools using option --singletons. We calculated the difference between the  
125 number of female-specific private alleles and male-specific private alleles for each 5 kbp window and  
126 extracted the average difference across 1 Mbp and 0.1 Mbp windows.

127 Supplementary figure S2 shows the results from the 1 Mbp window analysis from the  
128 Eurasian skylark and horned lark aligned to the reference genome of the Raso lark, as well as to the reference  
129 genome of their respective species.

### 130 **S1e. Extraction of Z–W sequences of gametologous genes**

131 We used the whole-genome synteny aligner program SatsumaSynteny v. 2.0 [15] to align the Raso lark  
132 assembly to the zebra finch assembly (taeGut.3.2.4), and then used kraken [16] to make a lift-over of the  
133 zebra finch annotations to the Raso lark assembly. Of the 18204 transcripts and 17488 genes in the zebra  
134 finch annotation, 14466 transcripts (79%) from 13764 genes (79%) were annotated in the Raso lark. We  
135 used Freebayes v.1.1.0 [12] (--report-monomorphic) to call variants for every base pair within all exons in  
136 the Raso lark based on the genome coordinates from the lift-over.

137 We *in silico* extracted gametologous gene sequences from sex-linked regions using in-house  
138 scripts (code provided as Code S1; general methodology described in [17] based on the genotypes of the  
139 female and male samples). The scripts use sex-specific genotype information to phase the data into a Z and  
140 W sequence. We replaced a site by “N” if it had a quality score or sequence depth below 20 or if the genotype  
141 in either sample was not called. Any site without variants in either sample was extracted as such, and  
142 remaining variants between the male and female were phased based on sex-specific allele compositions  
143 provided in electronic supplementary material, table S4.

144 To confirm that both Z and W gametologs were present in the data, we calculated genome  
145 coverage values for every exon using bedtools v.2.27.1 multicov [11] and normalised the values between the  
146 two samples of each species. Exons with female coverage less than 75% of the male sample were masked  
147 with N:s, as this suggests absence of a W gametolog.

148 We used TransDecoder v3.0.1 (<https://github.com/TransDecoder/TransDecoder/>) to  
149 find the longest open reading frames for each of the sequences (--retain\_pfam\_hits, --retain\_blastp\_hits, --  
150 single-best-orf). Orthologous genes from the zebra finch, collared flycatcher (*Ficedula albicollis*) and chicken  
151 (*Gallus gallus*) were downloaded through BioMart (database: Ensembl Genes 93), and the longest transcript  
152 from the flycatcher and chicken corresponding to the zebra finch transcripts in the annotation was added  
153 as additional sequences. The gene sequences of the zebra finch were later used to analyse the relative  
154 evolutionary rate between Z and W gametologs. The sequences were codon-aware aligned using *Prank*  
155 v.150803 [18]. Each N in the sequences was transformed into “-”, and all sites including this character in  
156 any sequence were then removed using gblocks v0.91b [19]. We calculated pairwise substitution rates  
157 between Z–W gametologs using codeml (estimated kappa and codon frequency F3X4) from the PAML  
158 v4.9 package [20]. Gene sequences longer than 500 bp, and where the Z and W sequences within a species  
159 had a synonymous substitution rate (dS) value above 0.01, were kept for further analysis. No comparisons  
160 between gene sequences had synonymous substitution values (dS) > 1, which may have been biasing the  
161 divergence estimates due to mutation saturation. The maximum dS value in our dataset was 0.4697 with >  
162 90% having dS values < 0.1.

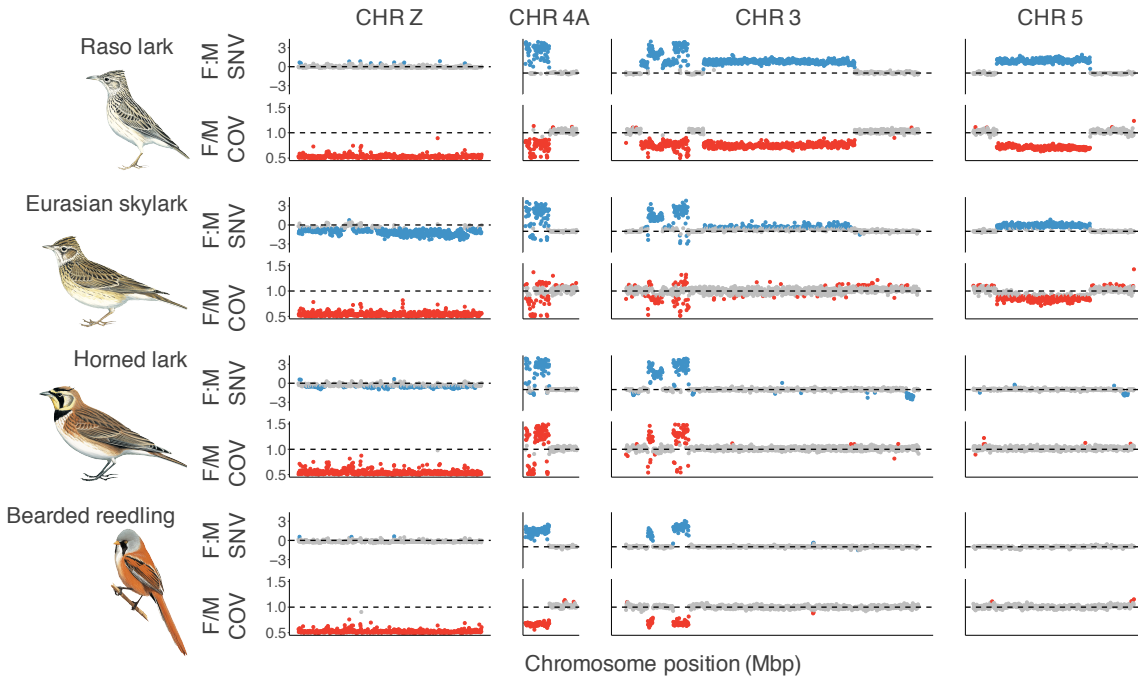
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164 S1f. References for Supplementary Methods S1

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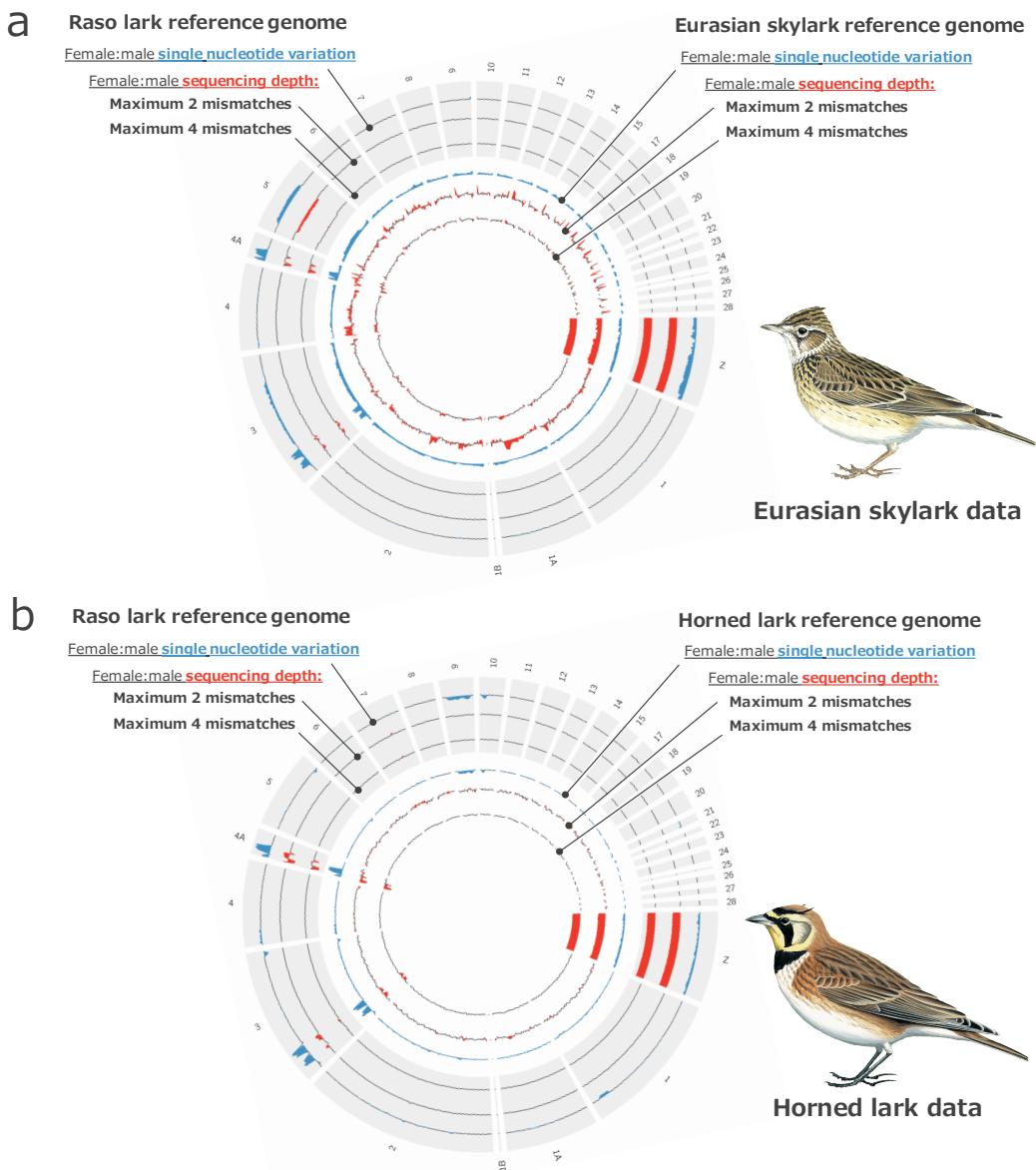
220 Supplementary Figures



221

222 **Figure S1.** Distribution of female-to-male difference in number of private single nucleotide  
223 variants (SNVs), and female-to-male coverage ratio, in the four study species across chromosomes  
224 Z, 4A, 3 and 5. Chromosome names and positions follow the zebra finch genome structure (see  
225 Methods section). The upper panel for each chromosome shows sex-specific SNV differences (tick  
226 marks represent 3000, 0 and -3000, with values > 500 or < -500 in blue) and the lower panel shows  
227 female-to-male coverage ratios (with values > 0.9 or < 1.1 in red). Both measurements are means  
228 across 0.1 Mbp windows. Bird drawings ordered from HBW.

229



230

231 **Figure S2.** For each species, Eurasian skylark (a) and horned lark (b), the grey ring shows the  
 232 same data that is presented for this species in Figure 1. The inner rings with white background  
 233 show the results when aligned to its own reference genome, starting with single nucleotide  
 234 variation data (in blue), followed by genome coverage data (in red) when parsed for 2 and 4  
 235 allowed mismatches. It is evident from the single nucleotide data for both species that the sex-  
 236 linked regions are the same whether the reference genome is the Raso lark or their respective  
 237 species. The genome coverage is however less clear, especially in the Eurasian skylark, which  
 238 is why the Raso lark was chosen as a reference genome for these two species.

239

240 Supplementary Tables

241

242 **Table S1. Genome assembly statistics (related to Supplementary Method S1b)**

243 Statistics was computed using the software Quast version 4.5.4 [3].

Assembly	Raso lark	Eurasian skylark	Horned lark	Bearded reedling
<b>N contigs (<math>\geq 0</math> bp)</b>	28304	256822	109088	36455
<b>N contigs (<math>\geq 1</math> kbp)</b>	28304	256822	109088	36455
<b>N contigs (<math>\geq 5</math> kbp)</b>	17977	79725	51289	24408
<b>N contigs (<math>\geq 10</math> kbp)</b>	14382	31198	32587	19461
<b>N contigs (<math>\geq 25</math> kbp)</b>	9767	3546	10545	12025
<b>N contigs (<math>\geq 50</math> kbp)</b>	6145	609	2805	6436
<b>Total length (<math>\geq 0</math> bp)</b>	1003896371	1276061662	1086584534	1020354129
<b>Total length (<math>\geq 1</math> kbp)</b>	1003896371	1276061662	1086584534	1020354129
<b>Total length (<math>\geq 5</math> kbp)</b>	979649475	887743951	962416262	991752640
<b>Total length (<math>\geq 10</math> kbp)</b>	953912007	543408884	827105499	955927387
<b>Total length (<math>\geq 25</math> kbp)</b>	878299503	145311035	479556955	833768162
<b>Total length (<math>\geq 50</math> kbp)</b>	748288932	49632478	216102095	634399538
<b>N contigs</b>	28304	256822	109088	36455
<b>Largest contig (bp)</b>	851956	346983	447328	534950
<b>Total length (bp)</b>	1003896371	1276061662	1086584534	1020354129
<b>GC (%)</b>	42.31	42.59	42.17	42.14
<b>N50</b>	103210	8477	21578	68188
<b>N75</b>	49107	4130	10418	32980
<b>L50</b>	2726	41480	13291	4314
<b>L75</b>	6239	94969	31395	9642
<b># N's per 100 kbp</b>	0.87	23.63	16.02	0.85

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246

247 **Table S2. Alignment statistics (related to Supplementary Method S1b)**

248 Alignment statistics for the female and male sample from the four studied species. No. read pairs refer to the  
 249 number of read pairs remaining after quality trimming. The number of properly aligning reads was calculated with  
 250 samtools v1.7 flagstat [6]. Sample ID correspond to the raw data files uploaded to NCBI short read archive.

Sample ID	Species	Reference genome	Sex	No. read pairs	Properly paired mapped reads (after deduplication)	
					Before mismatch filtering	<=2 mismatches
QL-1681-95694_S52_L008	Raso lark	Raso lark	female	101453455	174881744	161376774
QL-1681-246_S51_L008	Raso lark	Raso lark	male	87822864	152776522	147354471
QL-1681-19_S46_L006	Eurasian skylark	Raso lark	female	151270768	224364208	111786956
QL-1681-21_S47_L006	Eurasian skylark	Raso lark	male	147067636	227283590	114762381
QF-1504-H-19_S8_L003	Horned lark	Raso lark	female	130242267	180857630	69507055
QF-1504-H-88_S7_L003	Horned lark	Raso lark	male	202099679	275199656	104740879
QF-1504-2KR32024_S2_L001	Bearded reedling	Bearded reedling	female	98173644	138219068	125403224
QF-1504-1ET92164_S3_L001	Bearded reedling	Bearded reedling	male	163911536	230747108	218069639

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253

254 **Table S3. Proportion of reference genomes aligned to the zebra finch genome (related to Supplementary**  
255 **Method S1c)**

256 Statistics showing the length of each reference genome and the proportion that aligns to the zebra finch genome in  
257 the synteny analysis. The table show the total length of each genome in addition to the total length of each genome  
258 only counting scaffolds larger than 5 kbp. As only scaffolds larger than 5 kbp were kept in the synteny analysis, the  
259 proportion of the genome aligning to the zebra finch is based on this value.

<b>Assembly</b>	<b>Raso lark</b>	<b>Eurasian skylark</b>	<b>Horned lark</b>	<b>Bearded reedling</b>
Total length (>= 1000 bp)	1003896371	1276061662	1086584534	1020354129
Total length (>= 5000 bp)	979649475	887743951	962416262	991752640
Matching ZF genome	934820000	717725000	842675000	931690000
Prop. of >5kbp scaffolds aligned	0.95	0.81	0.88	0.94

261 **Table S4. Gametolog extraction criteria (related to Supplementary Method S1e)**

262 Z and W gametolog sequences were extracted based on the genotype distributions between the male and female  
 263 sample.

264

<b>Variant type</b>	<b>Nr of mutations</b>	<b>Male</b>	<b>Female</b>	<b>Extracted Z allele</b>	<b>Extracted W alle</b>
biallelic	0	0/0	0/0	0	0
	1	0/1	0/0	N	N
	1	0/0	0/1	0	1
	2	0/1	0/1	N	N
	3	1/1	0/1	1	0
	3	0/1	1/1	N	N
	4	1/1	1/1	1	1
triallelic	1	1/2	1/1	N	1
	1	1/1	1/2	1	2
	2	1/2	1/2	N	N
	3	2/2	1/2	2	1
	3	1/2	2/2	N	N
	4	2/2	2/2	2	2

0/0 = no variation in relation to reference genome

0/1 = heterozygote genotype in relation to reference genome

1/1 = homozygote genotype with one allele not found in reference genome

1/2 = heterozygote genotype with two different variants not found in reference genome

2/2 = homozygote genotype not found in reference genome

265

266 **Table S5. Signatures of sex-linkage across 1 Mbp windows (related to Figure 1)**

267 The mean female-to-male coverage ratio (cov) and difference in number of female-specific mutations and male-  
 268 specific mutations (snv) of all 1 Mbp windows per chromosome.

Chromosome	Raso lark		Eurasian skylark		Horned lark		Bearded reedling		
	coverage	snv	coverage	snv	coverage	snv	coverage	snv	
1	1.03	-55.76	1.03	59.39	1.01	-428.35	1.00	-90.65	
1A	1.03	-453.08	1.04	174.25	1.02	265.73	1.01	-193.37	
1B	0.98	130.50	1.06	278.00	1.03	-83.00	1.06	37.00	
2	1.03	129.38	1.03	91.55	1.02	46.71	1.00	-30.68	
3	0.84	13459.84	1.00	4958.29	1.03	2775.37	0.98	1898.89	
4	1.02	218.61	1.03	-121.41	1.01	503.89	1.00	-121.34	
4A	0.88	13033.77	0.97	9762.91	1.05	10978.41	0.85	10497.82	
5	0.83	11011.22	0.93	5102.22	1.02	-90.25	1.01	-122.97	
6	1.03	-64.81	1.03	52.16	1.02	43.51	1.01	78.08	
7	1.03	119.59	1.03	-65.32	1.02	-11.24	1.02	-64.37	
8	1.02	-309.62	1.03	3.86	1.01	-6.00	1.01	-62.31	
9	1.03	-727.96	1.03	409.54	1.01	-5353.43	1.02	21.68	
10	1.02	950.95	1.03	55.23	1.02	-1323.82	1.02	-171.36	
11	1.02	637.95	1.03	47.00	1.01	-18.09	1.02	-275.05	
12	1.03	130.96	1.03	-15.91	1.02	74.39	1.02	-9.70	
13	1.02	191.17	1.03	-6.50	1.02	-120.17	1.03	-367.50	
14	1.02	107.88	1.04	132.82	1.02	99.76	1.03	99.94	
15	1.02	-902.73	1.03	9.80	1.02	-8.47	1.03	-188.53	
17	1.02	-112.08	1.03	80.69	1.02	140.23	1.04	-448.00	
18	1.02	-666.17	1.03	-43.50	1.02	-71.42	1.03	467.58	
19	1.02	102.77	1.03	97.54	1.02	64.31	1.04	38.54	
20	1.02	-137.18	1.03	18.59	1.02	-19.94	1.04	-65.41	
21	1.02	-880.43	1.03	52.14	1.02	-1986.71	1.03	-563.29	
22	1.03	246.75	1.03	27.75	1.02	328.00	1.03	716.75	
23	1.01	-780.00	1.03	76.00	1.03	-99.57	1.04	-20.29	
24	1.03	-355.78	1.03	57.00	1.03	-78.22	1.04	285.00	
25	1.03	-99.50	1.07	-681.50	1.04	231.00	1.06	90.50	
26	1.02	-198.17	1.05	-1245.50	1.02	-95.50	1.05	103.00	
27	1.01	336.33	1.04	93.33	1.03	-228.83	1.04	160.33	
28	1.02	-52.50	1.04	187.83	1.03	134.33	1.05	-35.83	
269	Z	0.53	-174.12	0.54	-9719.91	0.54	-3682.34	0.52	-1570.14

270

271

272 **Table S6. Sex-linked regions in each of the four species (related to Figure 2).**

273 Mean values for the female-to-male coverage ratio (coverage) and female-to-male difference in number of private  
 274 SNVs across 0.1 Mbp windows for each stratum. The strata are numbered according to the most parsimonious order  
 275 of emergence based on phylogenetic analyses (see Main text).

Stratum	Chromosome	Genomic region (Mb)	Stratum size (Mb)				
				Raso lark	Eurasian skylark	Horned lark	Bearded reedling
1	Z	0-72.9	72.9	Coverage	0.53	0.54	0.54
				SNV	-17.77	-992.10	-375.85
2	4A	0-9.6	9.6	Coverage	0.74	0.91	1.07
				SNV	2957.35	2227.36	2518.46
3	3	8.4-10.4, 18.1-24.1	8	Coverage	0.75	0.96	1.22
				SNV	3386.68	2435.65	3404.79
4	3	10.4-14.0	3.6	Coverage	0.71	0.86	0.98
				SNV	3019.06	2161.91	2512.71
5a	3	5.8-8.4, 14.0-18.1, 29.8-88.0	64.9	Coverage	0.75	1.00	NA (1.01)##
				SNV	1785.39	451.48	NA (1.73)##
5b	5	9.1-45.4	36.3	Coverage	0.71	0.87	NA (1.01)##
				SNV	1952.04	879.14	NA (20.19)##

276

277 #Bearded reedling and horned lark have no sex-linkage for stratum 5a and 5b, and bearded reedling not for stratum 4, and are  
 278 therefore marked with NA in addition to the corresponding values.

279

280 **Table S7. Nucleotide substitution values for gametologous (Z-W) gene pairs (related to Figure 3)**

281 Nucleotide substitution values for gametologous (Z-W) gene pairs from Raso lark and Eurasian skylark for each sex  
 282 chromosome strata.

Stratum	Chromosome		No. genes	Median dS	Median dN	Median dN/dS
1	Z	Raso lark	9	0.224	0.030	0.097
		Eurasian skylark	10	0.213	0.030	0.082
2	4A	Raso lark	33	0.080	0.010	0.124
		Eurasian skylark	32	0.085	0.013	0.123
3	3	Raso lark	23	0.073	0.009	0.123
		Eurasian skylark	23	0.072	0.010	0.137
4	3	Raso lark	18	0.032	0.008	0.294
		Eurasian skylark	20	0.036	0.006	0.196
5a	3	Raso lark	168	0.018	0.002	0.116
		Eurasian skylark	186	0.018	0.002	0.115
5b	5	Raso lark	164	0.019	0.004	0.187
		Eurasian skylark	169	0.020	0.004	0.168

283

284 **Table S8. Nucleotide differentiation between evolutionary strata (related to Figure 3)**  
 285 P values from Kruskal-Wallis tests of nucleotide differences between pairs of evolutionary strata (for details, see  
 286 methods).

		Stratum 1	Stratum 2	Stratum 3	Stratum 4	Stratum 5a
Raso lark	Stratum 2	< 0.001				
Eurasian skylark		0.002				
Raso lark	Stratum 3	< 0.001	0.167			
Eurasian skylark		0.001	0.029			
Raso lark	Stratum 4	< 0.001	< 0.001	< 0.001	< 0.001	
Eurasian skylark		0.001	< 0.001	< 0.001	< 0.001	
Raso lark	Stratum 5a	< 0.001	< 0.001	< 0.001	< 0.001	
Eurasian skylark		< 0.001	< 0.001	< 0.001	< 0.001	
Raso lark	Stratum 5b	< 0.001	< 0.001	< 0.001	< 0.001	0.185
Eurasian skylark		< 0.001	< 0.001	< 0.001	< 0.001	0.087
<hr/>						
		Analysis of non-synonymous substitutions (dN)				
Raso lark	Stratum 2	0.273				
Eurasian skylark		0.66				
Raso lark	Stratum 3	0.273	0.887			
Eurasian skylark		0.660	0.986			
Raso lark	Stratum 4	0.142	0.746	0.772		
Eurasian skylark		0.208	0.66	0.66		
Raso lark	Stratum 5a	< 0.001	< 0.001	< 0.001	< 0.001	
Eurasian skylark		0.001	< 0.001	< 0.001	< 0.001	
Raso lark	Stratum 5b	0.001	< 0.001	0.001	0.003	< 0.001
Eurasian skylark		0.007	< 0.001	0.002	0.01	0.002
<hr/>						
		Analysis of rate of evolution (dN/dS)				
Raso lark	Stratum 2	0.861				
Eurasian skylark		0.874				
Raso lark	Stratum 3	0.861	0.861			
Eurasian skylark		0.874	0.874			
Raso lark	Stratum 4	0.105	0.073	0.861		
Eurasian skylark		0.089	0.133	0.874		
Raso lark	Stratum 5a	0.861	0.861	0.861	0.005	
Eurasian skylark		0.874	0.874	0.874	0.125	
Raso lark	Stratum 5b	0.861	0.861	0.861	0.861	0.002
Eurasian skylark		0.535	0.758	0.874	0.874	0.032

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291 **Table S9. Rates of evolution between Z gametologs and zebra finch, and W gametologs and zebra finch**  
 292 **(related to Results & Discussion 2b)**

293 The table show median substitution rates between zebra finch and Z-linked gametologs (Zlinked) and between zebra  
 294 finch and W-linked gametologs (Wlinked) for all evolutionary strata that have formed after the split with the zebra  
 295 finch (i.e. strata 2-5b). P values from paired samples Wilcoxon test, and after Benjamini & Hochberg correction for  
 296 five tests (one per stratum).

Stratum	Chromosome	Species	Substitution type	No. Genes	Median substitution rate to zebra finch			
					Zlinked	Wlinked	paired samples	Wilcoxon test
					p value	adjusted p value		
2	4A	Raso lark	dN	33	0.005	0.015	0.000	< 0.001
		Raso lark	dS	33	0.125	0.115	0.437	0.501
		Eurasian skylark	dN	31	0.008	0.015	0.000	< 0.001
		Eurasian skylark	dS	31	0.113	0.119	0.184	0.307
3	3	Raso lark	dN	23	0.007	0.017	0.000	< 0.001
		Raso lark	dS	23	0.106	0.095	0.501	0.501
		Eurasian skylark	dN	23	0.007	0.016	0.000	< 0.001
		Eurasian skylark	dS	23	0.103	0.101	0.482	0.603
4	3	Raso lark	dN	19	0.012	0.014	0.005	0.005
		Raso lark	dS	19	0.108	0.087	0.169	0.281
		Eurasian skylark	dN	19	0.010	0.018	0.003	0.003
		Eurasian skylark	dS	19	0.089	0.094	0.768	0.768
5a	3	Raso lark	dN	209	0.014	0.015	0.000	< 0.001
		Raso lark	dS	209	0.117	0.117	0.009	0.024
		Eurasian skylark	dN	209	0.013	0.014	0.000	< 0.001
		Eurasian skylark	dS	209	0.112	0.118	0.000	< 0.001
5b	5	Raso lark	dN	184	0.011	0.013	0.000	0.019
		Raso lark	dS	184	0.134	0.129	0.004	0.019
		Eurasian skylark	dN	184	0.011	0.013	0.000	< 0.001
		Eurasian skylark	dS	184	0.128	0.132	0.000	< 0.001

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298

299 **Table S10. Tests for enrichment of sex-related genes (related to Figure 4)**

300 Results from binomial tests for significant enrichment of genes involved in a range of sex-related functions (see  
 301 Methods section 4.6 for details). “Sex-related genes within region” corresponds to the observed genes in Figure 4.  
 302 Expected genes in Figure 4 was calculated as the total number of sex-related genes within the genome (n = 323)  
 303 divided by “Proportion of genome”. Genes mentioned in the Discussion section are marked in red.  
 304

Chromosome	Region	Proportion of genome		Sex-related genes	P value	Adjusted p value
<i>Strata level analysis</i>						
Z	1	Observed	Estimated (CI)	Genome-wide	Within region	
4A	2	0.058	0.05 (0.029-0.079)	323	16 <sup>#</sup>	0.633
3	3	0.008	0.015 (0.005-0.036)	323	5 <sup>\$</sup>	0.105
3	4	0.006	0.015 (0.005-0.036)	323	5 <sup>&amp;</sup>	0.058
3	5a	0.003	0 (0.000-0.011)	323	0	1
5	5b	0.052	0.087 (0.058-0.123)	323	28 <sup>€</sup>	0.008
						<b>0.047</b>
		<i>Total sex-linked region per chromosome analysis</i>				
Z	1	0.029	0.015 (0.005-0.036)	323	16	0.633
4A	2	0.029	0.015 (0.005-0.036)	323	5	0.105
3	3 + 4 + 5a	0.029	0.1 (0.071-0.140)	323	33	0.005
5	5b	0.029	0.015 (0.005-0.036)	323	5	0.365

# *B4GALT1, CRHBP, DDX4, DMRT1, DMRT3, DNAJ41, FANCC, FANCG, HEXB, KIF2A, LHFPL2, RAD23B, RPS6, SKP2, SPIN1, ZNF366*

\$ *AR, DACH2, DLAPH2, MSN, SEPT6*

& *FSHR, LBH, MEA1, MSH2, LHCGR (NA in zebra finch)*

€ *CAMD1, ARID4B, CCR6, CGA, CITED2, CYP1B1, ESR1, FOXO3, HSF2, LAT51, MAP3K4, MCM9, MEI4, NA (Uncharacterized protein), P4CRG, PGM3, QKI, ROS1, RWDD1, SLC22A16, SRD5A2, STRN, TBPL1, TCF21, TCTE1, UBE2J1, UBR2, UFL1*

! *CELF1, EIF2B2, SLIRP, TTL5, TYRO3*

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