Programmed DNA elimination of germline development genes in songbirds

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Summary

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Genomes can vary within individual organisms. Programmed DNA elimination leads to dramatic changes in genome organisation during the germline–soma differentiation of ciliates¹, lampreys², nematodes^{3,4}, and various other eukaryotes⁵. A particularly remarkable example of tissue-specific genome differentiation is the germline-restricted chromosome (GRC) in the zebra finch which is consistently absent from somatic cells⁶. Although the zebra finch is an important animal model system⁷, molecular evidence from its large GRC (>150 megabases) is limited to a short intergenic region⁸ and a single mRNA⁹. Here, we combined cytogenetic, genomic, transcriptomic, and proteomic evidence to resolve the evolutionary origin and functional significance of the GRC. First, by generating tissue-specific de-novo linked-read genome assemblies and re-sequencing two additional germline and soma samples, we found that the GRC contains at least 115 genes which are paralogous to single-copy genes on 18 autosomes and the Z chromosome. We detected an amplification of >38 GRC-linked genes into high copy numbers (up to 308 copies) but, surprisingly, no enrichment of transposable elements on the GRC. Second, transcriptome and proteome data provided evidence for functional expression of GRC genes at the RNA and protein levels in testes and ovaries. Interestingly, the GRC is enriched for genes with highly expressed orthologs in chicken gonads and gene ontologies involved in female gonad development. Third, we detected evolutionary strata of GRC-linked genes. Developmental genes such as bicc1 and trim71 have resided on the GRC for tens of millions of years, whereas dozens have become GRC-linked very recently. The GRC is thus likely widespread in songbirds (half of all bird species) and its rapid evolution may have contributed to their diversification. Together, our results demonstrate a highly dynamic evolutionary history of the songbird GRC leading to dramatic germline-soma genome differences as a novel mechanism to minimise genetic conflict between germline and soma.

Text

Not all cells of an organism must contain the same genome. Some eukaryotes exhibit dramatic differences between their germline and somatic genomes, resulting from programmed DNA elimination of chromosomes or fragments thereof during germline–soma differentiation⁵. Here we present the first comprehensive analyses of a germline-restricted chromosome (GRC). The zebra finch (*Taeniopygia guttata*) GRC is the largest chromosome of this songbird⁶ and likely comprises >10% of the genome (>150 megabases)^{7,10}. Cytogenetic evidence suggests the GRC is inherited through the female germline, expelled late during spermatogenesis, and eliminated from the soma during early embryo development^{6,11}. Previous analyses of a 19-kb intergenic region suggested that the GRC contains sequences with high similarity to regular chromosomes ('A chromosomes')⁸.

In order to reliably identify sequences as GRC-linked, we used a single-molecule sequencing technology not applied previously in birds that permits reconstruction of long haplotypes through linked reads¹². We generated separate haplotype-resolved *de-novo* genome assemblies for the germline and soma of a male zebra finch (testis and liver; 'Seewiesen'; Supplementary Table 1). We further used the linked-read data to compare read coverage and haplotype barcode data in relation to the zebra finch somatic reference genome ('taeGut2')⁷, allowing us to identify sequences that are shared, amplified, or unique to the germline genome in a fashion similar to recent studies on cancer aneuploidies¹³. We also re-sequenced the germline and soma from two unrelated male zebra finches ('Spain'; testis and muscle; Extended Data Fig. 1) using short reads.

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We first established the presence of the GRC in the three germline samples. Cytogenetic analysis using fluorescence in-situ hybridisation (FISH) with a new GRC probe showed that the GRC is present exclusively in the germline and eliminated during spermatogenesis as hypothesised (Fig. 1a-b, Extended Data Fig. 2)6,11. We compared germline/soma sequencing coverage by mapping reads from all three sampled zebra finches onto the reference genome assembly (regular 'A chromosomes'), revealing consistently germline-increased coverage for single-copy regions, reminiscent of programmed DNA elimination of short genome fragments in lampreys² (Fig. 1c-d). A total of 92 regions (41 with >10 kb length) on 13 chromosomes exhibit >4-fold increased germline coverage in 'Seewiesen' relative to the soma (Fig. 1e, Supplementary Table 2). Such a conservative coverage cut-off provides high confidence in true GRC-amplified regions. We obtained nearly identical confirmatory results using another library preparation method for the 'Spain' birds (Fig. 1f). Notably, the largest block of testis-increased coverage spans nearly 1 Mb on chromosome 1 and overlaps with the previously⁸ FISH-verified intergenic region 27L4 (Fig. 1e-f).

Our linked-read and re-sequencing approach allowed us to determine the sequence content of the GRC. The GRC is effectively a non-recombining chromosome as it recombines with itself after duplication, probably to ensure stable inheritance during female meiosis⁸. We predicted that the GRC would be highly enriched in repetitive elements, similar to the female-specific avian W chromosome (repeat density >50%, compared to <10% genome-wide)¹⁴. Surprisingly, neither assembly-based nor read-based repeat quantifications detected a significant enrichment in transposable elements or satellite repeats in the germline samples relative to the soma samples (Extended Data Figure 3, Supplementary Table 3). Instead, most germline coverage peaks lie in

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single-copy regions of the reference genome overlapping 38 genes (Fig. 1e-f, Table 1, Supplementary Table 4), suggesting that these peaks stem from very similar GRC-amplified paralogs with high copy numbers (up to 308 copies per gene; Supplementary Table 5). GRC linkage of these regions is further supported by sharing of linked-read barcodes between different amplified chromosomal regions in germline but not soma (Fig. 1g-h), suggesting that these regions reside on the same haplotype (Extended Data Fig. 4). We additionally identified 245 GRC-linked genes through germline-specific single-nucleotide variants (SNVs) present in read mapping of all three germline samples onto zebra finch reference genes (up to 402 SNVs per gene; Supplementary Table 4). As a control, we used the same methodology to screen for somaspecific SNVs and found no such genes. We conservatively consider the 38 GRC-amplified genes and those with at least 5 germline-specific SNVs as our highest-confidence set (Table 1). We also identified GRC-linked genes using germline-soma assembly subtraction (Fig. 1i); however, all were already found via coverage or SNV evidence (Table 1). Together with the *napa* gene recently identified in transcriptomes (Fig. 1j)⁹, our complementary approaches yielded 115 high-confidence GRC-linked genes with paralogs located on 18 autosomes and the Z chromosome (Table 1; all 267 GRC genes in Supplementary Table 4).

We next tested whether the GRC is functional and thus probably physiologically important using transcriptomics and proteomics. We sequenced RNA from the same tissues of the two Spanish birds used for genome re-sequencing and combined these with published testis and ovary RNA-seq data from North American domesticated zebra finches^{9,15}. Among the 115 high-confidence genes, 6 and 32 were transcribed in testes and ovaries, respectively (Table 1). Note, these are only genes for which we could reliably separate GRC-linked and A-chromosomal paralogs using

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GRC-specific SNVs in the transcripts (Fig. 2a-b, Extended Data Fig. 5, Supplementary Table 6). We next verified translation of GRC-linked genes through protein mass spectrometry data for 7 testes and 2 ovaries from another population ('Sheffield'). From 83 genes with GRC-specific amino acid changes, we identified peptides from 5 GRC-linked genes in testes and ovaries (Fig. 2c-d, Extended Data Fig. 6, Table 1). We therefore established that many GRC-linked genes are transcribed and translated in adult male and female gonads, extending previous RNA evidence for a single gene⁹ and questioning the hypothesis from cytogenetic studies that the GRC is silenced in the male germline^{16,17}. Instead, we propose that the GRC has important functions during germline development, which is supported by a significant enrichment in gene ontology terms related to reproductive developmental processes among GRC-linked genes (Fig. 2e, Supplementary Table 7). We further found that the GRC is significantly enriched in genes that are also germline-expressed in GRC-lacking species with RNA expression data available from many tissues¹⁸ (Fig. 2f, Supplementary Table 8). Specifically, out of 65 chicken orthologs of high-confidence GRC-linked genes, 22 and 6 are most strongly expressed in chicken testis and ovary, respectively.

The observation that all identified GRC-linked genes have A-chromosomal paralogs allowed us to decipher the evolutionary origins of the GRC. We utilised phylogenies of GRC-linked genes and their A-chromosomal paralogs to infer when these genes copied to the GRC, similarly to the inference of evolutionary strata of sex chromosome differentiation¹⁹. First, the phylogeny of the intergenic 27L4 locus of our germline samples and a previous GRC sequence⁸ demonstrated stable inheritance among the sampled zebra finch populations (Fig. 3a). Second, 37 gene trees of GRC-linked genes with germline-specific SNVs and available somatic genome data from other

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birds identify at least five evolutionary strata (Fig. 3b-f, Extended Data Fig. 7, Table 1), with all but stratum 3 containing expressed genes (cf. Fig. 2a-d). Stratum 1 emerged during early songbird diversification, stratum 2 before the diversification of estrildid finches, and stratum 3 within estrildid finches (Fig. 3g). The presence of at least 7 genes in these three strata implies that the GRC is tens of millions of years old and likely present across songbirds (Extended Data Fig. 7), consistent with a recent cytogenetics preprint²⁰. Notably, stratum 4 is specific to the zebra finch species and stratum 5 to the Australian zebra finch subspecies (Fig. 3g), suggesting piecemeal addition of genes from 18 autosomes and the Z chromosome over millions of years of GRC evolution (Fig. 3h). The long-term residence of expressed genes on the GRC implies that they have been under selection, such as bicc1 and trim71 on GRC stratum 1 whose human orthologs are important for embryonic cell differentiation²¹. Using ratios of non-synonymous to synonymous substitutions (dN/dS) for GRC-linked genes with >50 GRC-specific SNVs, we found 17 genes evolving faster than their A-chromosomal paralogs (Supplementary Table 9). However, we also detected long-term purifying selection on 9 GRC-linked genes, including bicc1 and trim71, as well as evidence for positive selection on puf60, again implying that the GRC is an important chromosome with a long evolutionary history.

Here we provided the first evidence for the origin and functional significance of a GRC. Notably, our analyses suggest that the GRC emerged during early songbird evolution and we predict it to be present in half of all bird species. The species-specific addition of dozens of genes on stratum 5 implies that the rapidly evolving GRC likely contributed to reproductive isolation during the massive diversification of songbirds²². It was previously hypothesised that GRCs are formerly parasitic B chromosomes that became stably inherited^{23,24}. Our evidence for an enrichment of

germline-expressed genes on the zebra finch GRC is reminiscent of nematodes and lampreys where short genome fragments containing similar genes are eliminated during germline-soma differentiation²⁻⁴. All these cases constitute extreme mechanisms of gene regulation through germline-soma gene removal rather than transcriptional repression^{3,5,10}. Remarkably, the GRC harbours several genes involved in the control of cell division and germline determination, including *prdm1*, a key regulator of primordial germ cell differentiation in mice^{25,26}. Consequently, we hypothesise that the GRC became indispensable for its host by the acquisition of germline development genes and probably acts as a germline-determining chromosome. The aggregation of developmental genes on a single eliminated chromosome constitutes a novel mechanism to ensure germline-specific gene expression in multicellular organisms. This may allow adaptation to germline-specific functions free of detrimental effects on the soma which would otherwise arise from antagonistic pleiotropy.

References (max. 30 references)

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- 176 1 Chen, X. *et al.* The architecture of a scrambled genome reveals massive levels of genomic
- rearrangement during development. *Cell* **158**, 1187-1198 (2014).
- Smith, J. J. et al. The sea lamprey germline genome provides insights into programmed
- genome rearrangement and vertebrate evolution. *Nat. Genet.* **50**, 270-277 (2018).
- Wang, J. et al. Silencing of germline-expressed genes by DNA elimination in somatic
- 181 cells. Dev. Cell 23, 1072-1080 (2012).
- Wang, J. et al. Comparative genome analysis of programmed DNA elimination in
- nematodes. *Genome Res.* **27**, 2001-2014 (2017).
- Wang, J. & Davis, R. E. Programmed DNA elimination in multicellular organisms. *Curr*.
- 185 *Opin. Genet. Dev.* **27**, 26-34 (2014).

- 186 6 Pigozzi, M. I. & Solari, A. J. Germ cell restriction and regular transmission of an
- accessory chromosome that mimics a sex body in the zebra finch, *Taeniopygia guttata*.
- 188 *Chromosome Res.* **6**, 105-113 (1998).
- 189 7 Warren, W. C. *et al.* The genome of a songbird. *Nature* **464**, 757–762 (2010).
- 190 8 Itoh, Y., Kampf, K., Pigozzi, M. I. & Arnold, A. P. Molecular cloning and
- characterization of the germline-restricted chromosome sequence in the zebra finch.
- 192 *Chromosoma* **118**, 527-536 (2009).
- 193 9 Biederman, M. K. et al. Discovery of the first germline-restricted gene by subtractive
- transcriptomic analysis in the zebra finch, *Taeniopygia guttata*. Curr. Biol. 28, 1620-1627
- 195 (2018).
- 196 10 Smith, J. J. Programmed DNA elimination: keeping germline genes in their place. *Curr*.
- 197 *Biol.* **28**, R601-R603 (2018).
- 198 11 Pigozzi, M. I. & Solari, A. J. The germ-line-restricted chromosome in the zebra finch:
- recombination in females and elimination in males. *Chromosoma* **114**, 403-409 (2005).
- Weisenfeld, N. I., Kumar, V., Shah, P., Church, D. M. & Jaffe, D. B. Direct determination
- of diploid genome sequences. Genome Res. 27, 757-767 (2017).
- 202 13 Bell, J. M. et al. Chromosome-scale mega-haplotypes enable digital karyotyping of cancer
- aneuploidy. *Nucleic Acids Res.* **45**, e162-e162 (2017).
- Kapusta, A. & Suh, A. Evolution of bird genomes—a transposon's-eye view. Ann. N. Y.
- 205 *Acad. Sci.* **1389**, 164–185 (2017).
- Singhal, S. et al. Stable recombination hotspots in birds. Science **350**, 928-932 (2015).
- 207 16 del Priore, L. & Pigozzi, M. I. Histone modifications related to chromosome silencing and
- elimination during male meiosis in Bengalese finch. *Chromosoma* **123**, 293-302 (2014).
- 209 17 Goday, C. & Pigozzi, M. I. Heterochromatin and histone modifications in the germline-
- restricted chromosome of the zebra finch undergoing elimination during spermatogenesis.
- 211 *Chromosoma* **119**, 325-336 (2010).
- 212 18 Marin, R. et al. Convergent origination of a *Drosophila*-like dosage compensation
- 213 mechanism in a reptile lineage. *Genome Res.* **27**, 1974-1987 (2017).
- Lahn, B. T. & Page, D. C. Four evolutionary strata on the human X chromosome. *Science*
- **286**, 964-967 (1999).
- 216 20 Torgasheva, A. A. et al. Germline-restricted chromosome (GRC) is widespread among
- 217 songbirds. *bioRxiv* doi:10.1101/414276 (2018).

- 218 Uhlén, M. et al. Tissue-based map of the human proteome. Science 347 (2015).
- 219 Moyle, R. G. et al. Tectonic collision and uplift of Wallacea triggered the global songbird
- radiation. *Nat. Commun.* **7**, 12709 (2016).
- 221 23 Camacho, J. P. M. B chromosomes. in *The Evolution of the Genome* (ed T. Ryan
- Gregory) 223-286 (Elsevier Academic Press, 2005).
- 223 24 Camacho, J. P. M., Sharbel, T. F. & Beukeboom, L. W. B-chromosome evolution. *Philos*.
- 224 Trans. R. Soc. B **355**, 163-178 (2000).
- 25 Ohinata, Y. et al. Blimp1 is a critical determinant of the germ cell lineage in mice. Nature
- **436**, 207 (2005).
- 227 26 Vincent, S. D. et al. The zinc finger transcriptional repressor Blimp1/Prdm1 is
- dispensable for early axis formation but is required for specification of primordial germ
- cells in the mouse. *Development* **132**, 1315-1325 (2005).
- Hooper, D. M. & Price, T. D. Rates of karyotypic evolution in Estrildid finches differ
- between island and continental clades. *Evolution* **69**, 890-903 (2015).
- 232 28 Mossman, J. A., Birkhead, T. R. & Slate, J. O. N. The whole mitochondrial genome
- sequence of the zebra finch (*Taeniopygia guttata*). *Molecular Ecology Notes* **6**, 1222-
- 234 1227 (2006).

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Tables and Figures

Table 1 | The 115 high-confidence genes on the GRC with information on their A-chromosomal origin in the reference genome taeGut2, number of testis-specific SNVs, methods supporting their GRC linkage, testis/ovary RNA expression of the GRC paralog, testis/ovary protein expression of the GRC paralog, and evolutionary stratum on the GRC.

Gene symbol	Chr.	Start	End	SNV s	Method	RNA evidence	Protein evidence	GRC stratum
AAGAB	10	19608548	19634367	10	SNVs			S5
ADGRL2	8	14047115	14171612	10	SNVs			
ADGRL3	4	14919933	15404594	8	SNVs	ovary		
AKIRIN2	3	78683482	78688947	6	SNVs	ovary		S5
ALDH18A1	6	36280145	36301392	17	SNVs			S4
ALG13	4A	18474239	18501426	19	SNVs	ovary		
ARMC6	28	4942046	4946063	5	SNVs			
ATP2A2	15	2841010	2879975	8	SNVs			
BICC1	6	6355408	6434911	402	SNVs	ovary		S1
BMP15	4A	15596686	15598225	29	SNVs, coverage	ovary		S5
BMPR1B	4	18997710	19024248	47	SNVs, coverage			S5
CCND3	26_rando m			14	SNVs			
CD164	3	69169111	69174605	38	SNVs, coverage	ovary		
COPS2	10	10200701	10222248	1	SNVs, coverage	ovary		
CPEB1	10	3114181	3137661	114	SNVs	ovary		
CSNK1A1L	Un	13542220 1	13542579 2	NA	coverage			
CXCL14	13	9423543	9433139	12	SNVs			S5
DDX49	28	4913058	4918451	5	SNVs	ovary		
DIS3L	10	19097281	19112154	13	SNVs	ovary		S5
DNAAF5	14	13758049	13780402	NA	coverage			
DNAH5	2	81235805	81361091	7	SNVs			
DPH6	5	31543945	31606965	13	SNVs, coverage			
EFNB1	4A	5764021	5807953	86	SNVs	ovary		S5
ELAVL4	8	21034240	21098310	364	SNVs	ovary		

EPPK1	Un			52	SNVs		
FBXO16	3	11254186 5	11256894 8	6	SNVs		
FEM1B	10	19886491	19891616	9	SNVs	ovary	S5
FIG4	3	69023384	69073678	17	SNVs		S5
FRS3	26_rando m			42	SNVs, coverage		S5
GBE1	1	10582064 0	10593431 0	4	SNVs, coverage		
INTS9	3	11225995 1	11231351 2	NA	coverage		
LIAS	4	48132714	48139736	42	SNVs		S2
LIN54	4	13615974	13637371	17	SNVs		
LINC02027	1	10608659 6	10608703 3	NA	coverage		
LMBRD2	Z	41646446	41665840	NA	coverage		
LOC10022319 0	Z	69149414	69156994	41	SNVs		
LOC10022423 5	Un			5	SNVs		S5
LOC10022532 2	1A	47543094	47544622	6	SNVs	ovary	
LOC10022718 9	Un	15079714 2	15080199 7	NA	coverage		
LOC10022817 0	Un	55540047	55541360	NA	coverage		
LOC10123308 7	Z	47991391	47994344	7	SNVs		
LOC10123368 8	5	937818	939059	5	SNVs		S5
LOC10123376 7	18	8034939	8038005	11	SNVs		
LOC10123380 0	Un			16	SNVs		
LOC10123425	10	19184028	19186114	7	SNVs	ovary	S5
LOC10575846 4	23	46808	60360	14	SNVs		S5
LOC10575889 4	26_rando m			5	SNVs		

LOC10575897 6	2	34301994	34306899	16	SNVs		
LOC10575910 1	3	76396180	76401262	21	SNVs		
LOC10575916 7	4A	15573874	15574621	5	SNVs		
LOC10575919 5	4	14453003	14473747	18	SNVs		
LOC10575919 9	4	20714525	20720872	11	SNVs		
LOC10575926 0	5	1874731	1886007	32	SNVs		S5
LOC10575964 6	Un			7	SNVs		
LOC10575965 5	Un			8	SNVs		
LOC10575966 0	Un			18	SNVs		
LOC10575966 5	Un			5	SNVs		
LOC10575969 2	Un			12	SNVs		
LOC10575991 9	Un			8	SNVs		
LOC10576001	Un			7	SNVs		
LOC10576012 3	Un			18	SNVs		
LOC10576022 8	Un			14	SNVs		
LOC10576028 6	Un			18	SNVs		
LOC10576046	Un			10	SNVs		
LOC10576087 4	Z	60949696	60953194	19	SNVs	testis	
LOC10576093 6	16_rando m			12	SNVs		
LUC7L3	Un	35019850	35021569	NA	coverage		
MED20	26_rando m	110500	113183	28	SNVs, coverage		S5

MSH4	8	27964612	27983306	30	SNVs			S4
NAPA	NA			NA	Biederman et al. 2018		both	
NEUROG1	13	9450787	9451086	6	SNVs			
NFYA	26	4725655	4735626	7	SNVs			S5
NRBP2	2	15637934 5	15639822 5	48	SNVs			
PCSK4	28	4059367	4063775	21	SNVs			
PGC	26_rando m			24	SNVs			
PHKA1	4A	15562688	15593666	16	SNVs			
PIM1	26	603349	607242	50	SNVs	testis		
PIM3	1A	18426716	18430551	81	SNVs	ovary		
PMM1	1A	49038672	49047011	NA	coverage			
PRDM1	3	70624594	70644625	12	SNVs			
PRKAR1A	18	2200317	2211579	NA	coverage			
PRKAR1B	14	13784578	13872733	NA	coverage			
PRPSAP1	18	8008870	8033058	7	SNVs, coverage	ovary		S5
PSIP1	Z	59887174	59919902	57	SNVs, coverage	ovary		S3
PUF60	2	15635467 0	15637609 1	63	SNVs	ovary		
RFC1	4	48169638	48202709	77	SNVs	ovary		S2
RNF157	18	8048721	8062403	NA	coverage			
RNF17	1	45827734	45870640	69	SNVs	ovary	testis	S4
RNF20	Z_random			9	SNVs, subtraction	both		
ROBO1	1	10709452 1	10722850 9	19	SNVs, coverage	ovary		S5
ROBO2	1	10752936 5	10797930 2	25	SNVs			
RXRA	17	8320685	8355067	14	SNVs			S5
SCRIB	2	15623988 4	15632579 7	83	SNVs	ovary		S5
SECISBP2L	10	10159176	10193647	60	SNVs, coverage	ovary	both	S5

SHC4	10	10124441	10151124	11	SNVs, coverage			S4
SPHK1	18	7991834	7994408	2	SNVs, coverage	testis		
SRRT	Un			16	SNVs	both		
SUGP2	28	4930094	4937971	33	SNVs	ovary	both	S5
SURF4	17	7682661	7693000	50	SNVs	ovary		S3
TFEB	26_rando m	20475	21840	11	SNVs			S5
TIAM2	3	54800961	54890499	NA	coverage			
TRIM71	2	60893878	60907039	159	SNVs, subtraction			S1
UBE2O	18	7960889	7981633	NA	coverage			
UGDH	4	48113314	48126079	136	SNVs, coverage, subtraction	ovary	ovary	S2
UNC5C	4	19035187	19126466	13	SNVs, coverage			
Unnamed	Un	12457451 3	12457555 3	NA	coverage			
Unnamed	Un	12712981 9	12713050 3	NA	coverage			
Unnamed	16_rando m	26580	73126	NA	coverage			
Unnamed	Un	13010351 4	13010426 4	NA	coverage			
Unnamed	Un	50859565	50860210	NA	coverage			
Unnamed	Un	11535588 3	11535815 4	NA	coverage			
Unnamed	Un	12457859 5	12457932 6	NA	coverage			
VEGFA	3	31631385	31652650	34	SNVs, coverage	both		
WDR19	4	48204115	48240398	34	SNVs	ovary		S5
ZWILCH	10	19199771	19206407	8	SNVs	ovary		S5

Note: We were able to place only some genes on evolutionary strata due to our strict criteria for evaluating the maximum likelihood gene trees. The remaining genes lacked sequence information from several of the other sampled somatic genomes or had poorly resolved tree topologies.

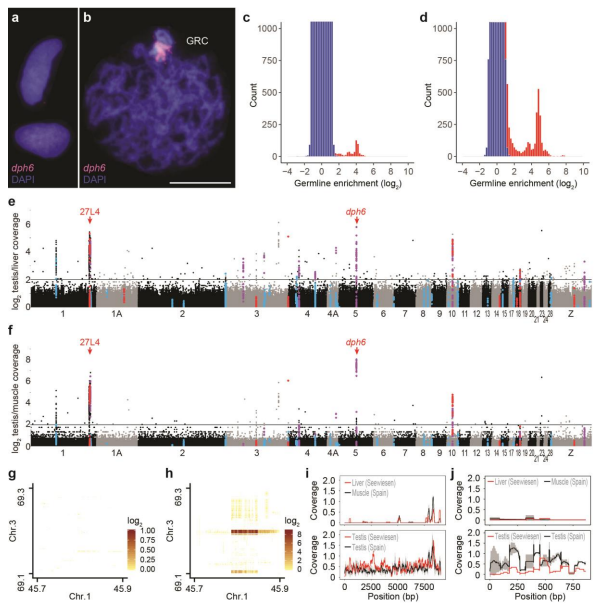


Figure 1 | The zebra finch germline-restricted chromosome contains genes copied from many A chromosomes. a-b, Cytogenetic evidence for GRC absence in muscle (a) and GRC presence in the testis (b) of the same bird (Spain 1) using fluorescence in-situ hybridisation (FISH) of our new GRC-ampliconic probe dph6 (selected due its high germline/soma coverage ratio; cf. panels e-f). The scale bar indicates 10 µm. c-d, Comparison of germline/soma coverage ratios (red) for 1 kb windows with an expected symmetrical distribution (blue) indicates enrichment of single-copy regions in the germline, similar to lamprey² both in Spain (c; average of Spain 1 and Spain 2 coverage; PCR-free short reads) and Seewiesen (d; linked reads) samples. Y-axis is truncated for visualisation. e-f, Manhattan plot of germline/soma coverage ratios in 1 kb windows across chromosomes of the somatic reference genome taeGut2. Colours indicate high-confidence GRC-linked genes and their identification (red: coverage, blue: SNVs, purple: both; Table 1). Note that the similarities between Seewiesen (e) and Spain_1/Spain_2 averages (f) constitute independent biological replicates for GRC-ampliconic regions, as the data are based on different domesticated populations and different library preparation methods. Red arrows denote two FISH-verified GRC-amplified regions (cf. panel b)8. Only chromosomes > 5 Mb are shown for clarity. g-h, Linked-read barcode interaction heatmaps of an inter-chromosomal rearrangement on the GRC absent in Seewiesen liver (g) but present in Seewiesen testis (h). i-j, Coverage plots of two examples of GRC-linked genes that are divergent from their A-chromosomal paralog, trim71 (i) and napa (j)9, and thus have very low coverage (normalised by total reads and genome size) in soma.

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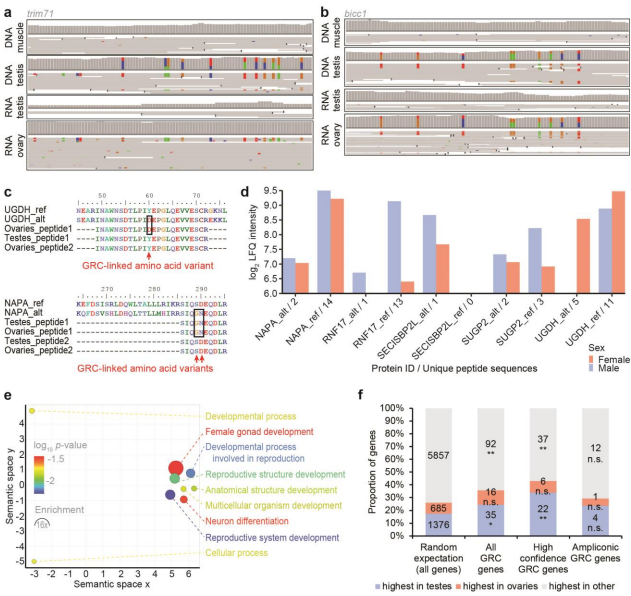


Figure 2 | The zebra finch germline-restricted chromosome is expressed in male and female gonads. a-b, Comparison of coverage and read pileups for DNA-seq data from Spain_1 and Spain_2 testis/muscle, RNA-seq data from Spain_1 and Spain_2 testis, and available ovary RNA-seq data9. Shown are 100-bp regions within *trim71* (a) and *bicc1* (b). Colours indicate SNVs deviating from the reference genome taeGut2. c, Example alignments of proteomics data showing a subset of peptide expression of the respective GRC-linked paralog of *ugdh* and *napa* (alternative or 'alt'; *cf.* reference or 'ref'). d, Proteomic evidence for GRC protein expression ('alt') in comparison to their A-chromosomal paralog ('ref') of 5 genes in 7 sampled testes and 2 sampled ovaries. For label-free quantification (LFQ), unique as well as razor (non-unique) peptides were used. Note that unique peptides may occur in several of the 9 samples. e, Gene ontology term enrichment analysis of the 115 high-confidence GRC-linked genes (77 mapped gene symbols). Colours indicate the log₁₀ of the false discovery rate-corrected *p*-value, circle sizes denote fold enrichment above expected values. f, Expression evidence for orthologs of three different sets of GRC genes in testes, ovaries, or other tissues in chicken¹⁸. Randomisation tests show a significant enrichment for germline-expressed genes among the 115 high-confidence GRC genes and all 267 GRC genes, but not the 38 ampliconic GRC genes.

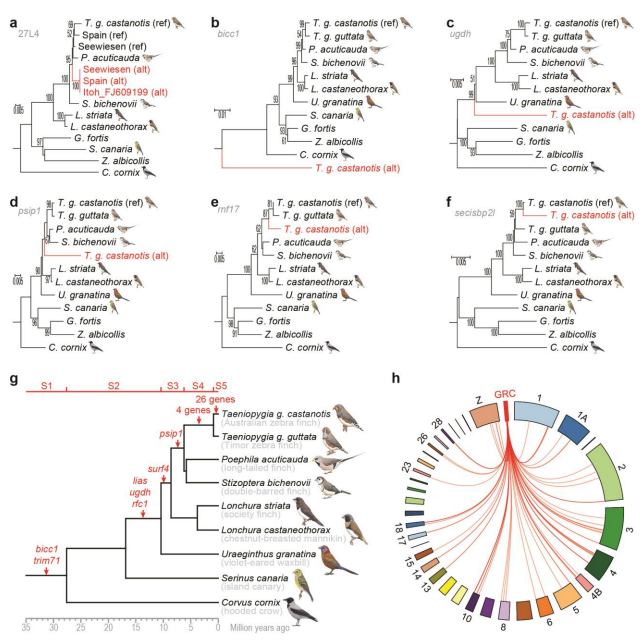
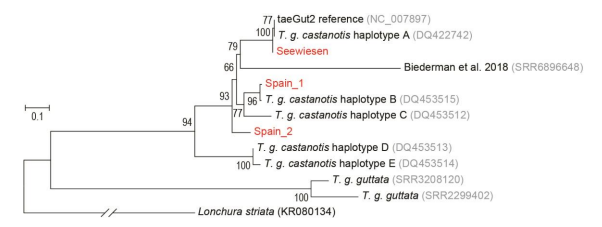
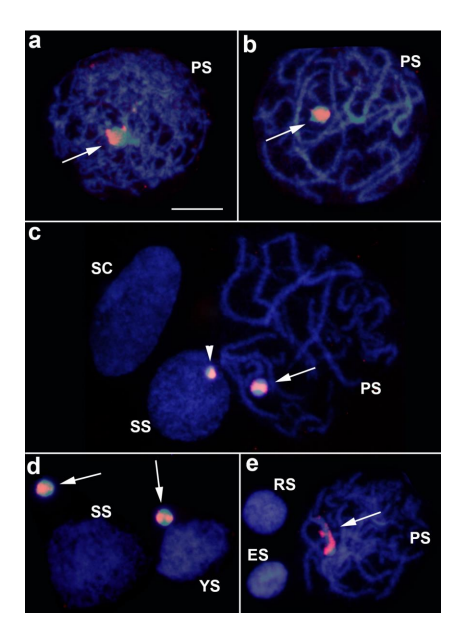


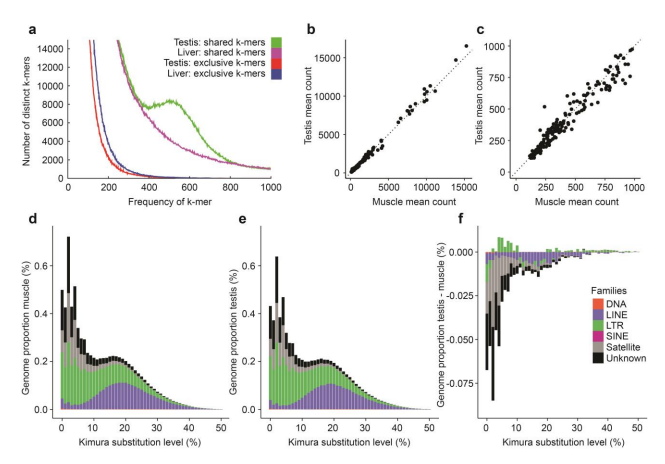
Figure 3 | The zebra finch germline-restricted chromosome is ancient and highly dynamic. a, Phylogeny of the intergenic 27L4 locus previously sequenced by Itoh et al.⁸ suggests stable inheritance of the GRC paralog (alternative or 'alt' in red; *cf.* reference or 'ref') among the sampled zebra finches. b-f, Phylogenies of GRC-linked genes ('alt', in red; most selected from expressed genes) diverging from their A-chromosomal paralogs ('ref') before/during early songbird evolution (b; *bicc1*, stratum 1; *cf.* Extended Data Fig. 7), during songbird evolution (c; *ugdh*, stratum 2), during estrildid finch evolution (d; *psip1*, stratum 3), in the ancestor of the zebra finch species (e; *rnf17*, stratum 4), and in the Australian zebra finch subspecies (f; *secisbp2l*; stratum 5). The maximum likelihood phylogenies in panels a-f (only bootstrap values ≥50% shown) include available somatic genome data from estrildid finches and other songbirds. g, Species tree of selected songbirds showing the emergence of evolutionary strata (S1–S5) on the GRC (red gene names). Molecular dates are based on previous phylogenies^{22,27}. Bird illustrations were used with permission from Lynx Edicions. h, Circos plot indicating A-chromosomal origin of high-confidence GRC-linked genes from 18 autosomes and the Z chromosome. Note that A-chromosomal paralogs of 37 genes remain unplaced on chromosomes in the current zebra finch reference genome taeGut2.



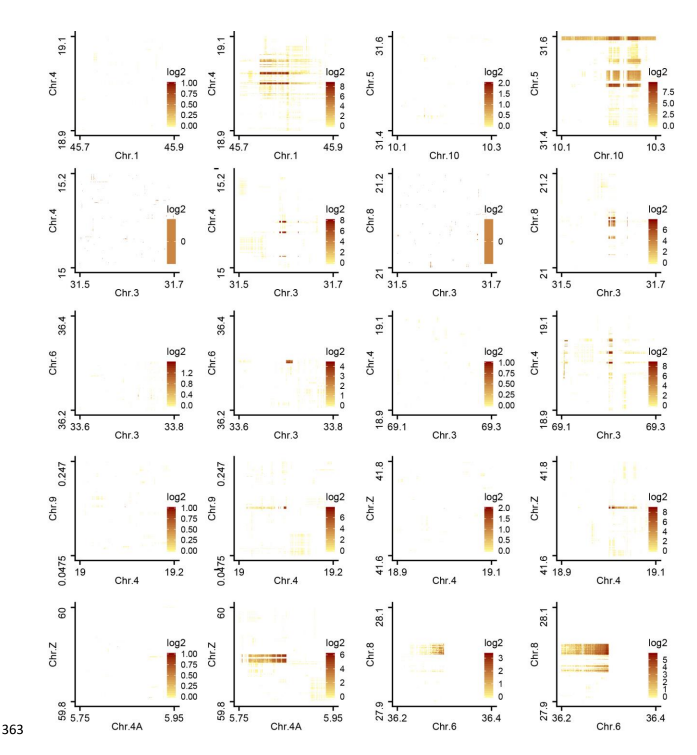
Extended Data Figure 1 | Maximum likelihood phylogeny of the five zebra finch mitochondrial haplotypes described by Mossman et al.²⁸ and mitogenomes assembled from all zebra finch Illumina libraries used in this work, comprising both the Australian zebra finch (*Taeniopygia guttata castanotis*) and the Timor zebra finch (*Taeniopygia guttata guttata*) subspecies. Note that the three individuals sequenced by us (red colour) and by Biederman et al.⁹ belong to different mitochondrial haplotypes.



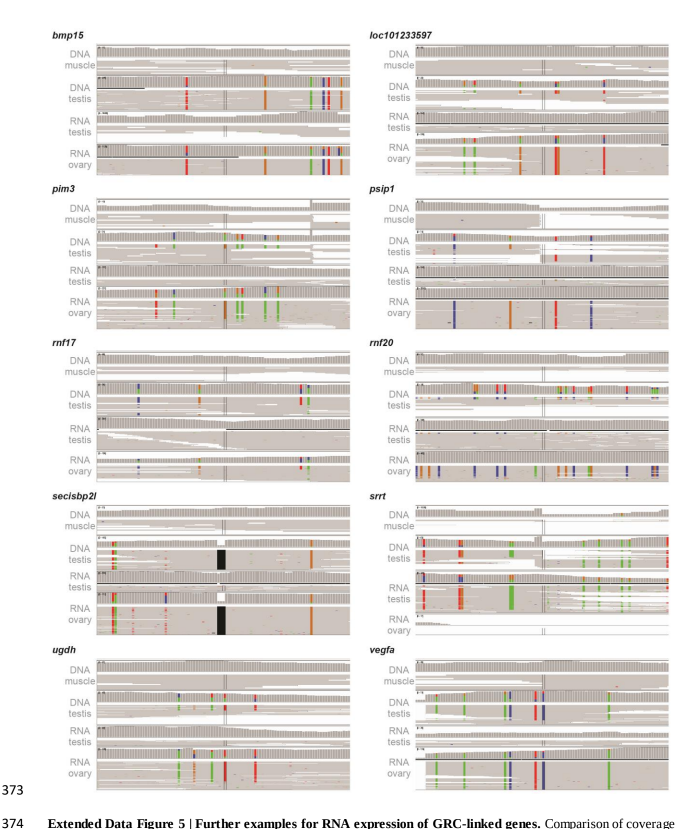
Extended Data Figure 2 | FISH analysis in testis cells of the Spain_1 zebra finch individual using the *dph6* probe (red) counterstained with DAPI (blue). Note the presence of primary (PS) and secondary (SS) spermatocytes, young spermatids (YS) and maturing spermatids at round (RS) and elongating (ES) stages. Also note that the *dph6* probe hybridises with only part of the GRC chromosome (arrow), and this is apparent in PS at leptotene-zygotene (a), pachytene (b-c, e) and in GRCs which failed to integrate into the main nucleus of SS or YS cells (d), with no FISH signal in somatic cells (SC) indicating GRC absence in somatic structural testis cells (c). The half size of GRC in the SS cell in panel c, compared with that in the PS next to it and that those lying outside nuclei in panel d, indicates that GRC sometimes divides equationally in the first meiotic division (resulting in the half sized GRC body in panel c) but, in most cases, it divides reductionally yielding the large sized GRCs in panel d. Note that RS and ES nuclei in panel e lack FISH signal, indicating GRC absence. All photographs were made at the same magnification, and the scale bar in panel a indicates 10 μm.



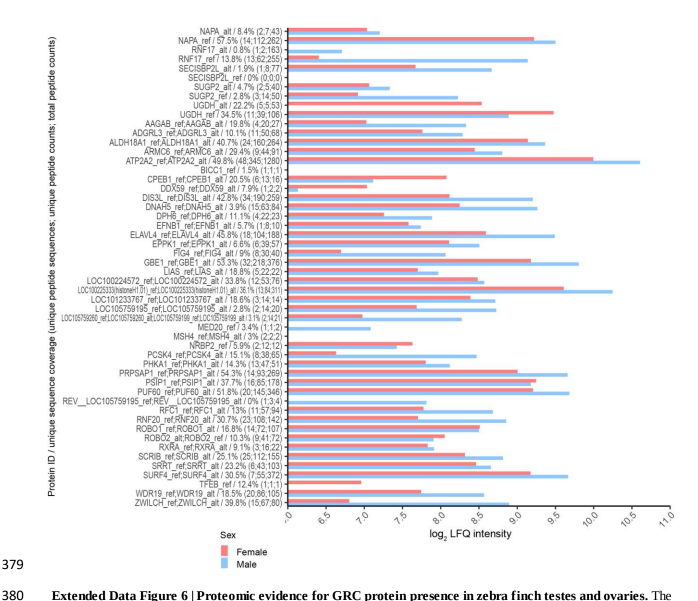
Extended Data Figure 3 | The zebra finch GRC is not enriched in satellites or specific transposable element families. **a**, Comparison of spectra for k-mers shared between or exclusive to genome sequencing data from testis and liver of the Seewiesen sample, showing that the germline is not enriched for exclusive high frequency k-mers, but is conspicuously enriched in high frequency k-mers shared with the soma. **b**, Comparison of simple repeat abundance as assessed by kSeek in the Spanish muscle samples relative to the testis samples. **c**, Same as in panel b, with a focus on low abundance simple repeats. **d-e**, Repeat landscapes based on RepeatMasker analyses showing the main repetitive element families for genome re-sequencing data from muscle (**d**) and testis (**e**) of the combined Spanish samples. **f**, Subtractive repeat landscape obtained by subtracting muscle from testis counts showing a general impoverishment of testis for most of the repetitive elements (negative values) due to the presence of the GRC.



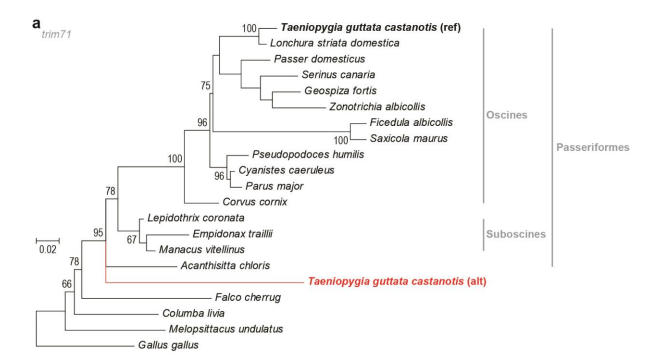
Extended Data Figure 4 | **Testis-specific linked-read barcode sharing between A chromosomes indicates GRC haplotypes.** Plots show side-by-side comparison of the inter-chromosomal barcode overlap for 200-kb regions for the liver and testis, respectively (chromosome position scale in Mb). With the exception of the interaction between chromosome 6 and chromosome 8 (bottom right) showing some background in the liver sample (potentially due to a shared A-chromosomal rearrangement), all inter-chromosomal structural variants were testis-specific and thus indicative of being on the same haplotype on the GRC. We exported barcode overlap matrices from the Loupe browser for testis-specific structural variants called by LongRanger and plotted them in R (v. 3.5.1). We reassigned 0 values to "NA" (shown in white on the plot) and log₂-transformed all values. Note that the scale varies across plots.

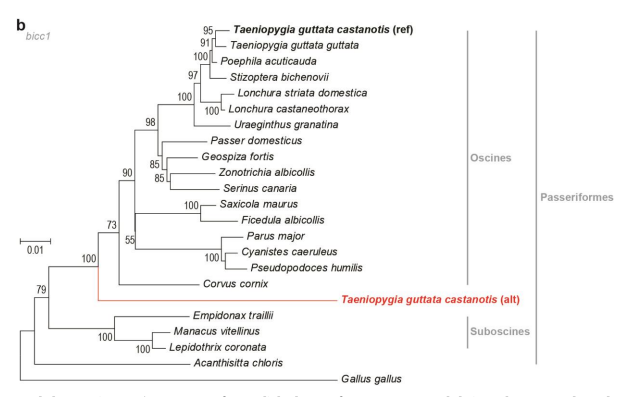


Extended Data Figure 5 | Further examples for RNA expression of GRC-linked genes. Comparison of coverage and read pileups for DNA-seq from Spain_1 and Spain_2 testis/muscle, RNA-seq data from Spain_1 and Spain_2 testis, and available ovary RNA-seq data⁹. Shown are 100-bp regions within 10 selected genes. Colours indicate SNVs deviating from the zebra finch reference genome taeGut2.



Extended Data Figure 6 | Proteomic evidence for GRC protein presence in zebra finch testes and ovaries. The five proteins listed at the top are also shown in Fig. 2d, i.e., those where we could differentiate between peptides from GRC vs. A chromosomes. GRC paralogs are denoted by the 'alt' suffix, whereas A-chromosomal paralogs are denoted by the 'ref' suffix. Unique sequence coverage corresponds to the peptide coverage percentage of the reference protein sequence. Note that unique peptides may occur in several samples (testes/ovaries). Entries of only one protein identification have sufficient evidence at the peptide level to differentiate between the GRC and A-chromosomal paralogs due to coverage of non-identical regions between the both reference sequences; entries of more than one protein identification contain evidence of presence based solely on identical regions, thus cannot be differentiated at the proteomic level. Entries of only one protein identification without the corresponding 'alt' or 'ref' variant contain evidence that span the non-identical region only, thus the alternate variant need not be called.





Extended Data Figure 7 | Gene trees of GRC-linked genes from stratum 1 and their A-chromosomal paralogs from broad taxon sampling imply GRC emergence in the ancestor of Passeriformes. a, Maximum likelihood gene tree of *trim71* (partitioned for codon positions) suggesting GRC linkage in the ancestor of Passeriformes. b, Maximum likelihood gene tree of *bicc1* (only 3' UTR) suggesting GRC linkage in the ancestor of oscine songbirds.

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Supplementary Information Methods and Supplementary Text Supplementary Table 1 | Assembly metrics of linked-read de-novo assemblies generated from liver and testis samples of the Seewiesen zebra finch individual. Supplementary Table 2 | Position, length, and library source of genomic blocks >10-kb with average germline/soma corrected coverage >4, with respect to the zebra finch reference genome (taeGut2). Supplementary Table 3 | Repeat annotation of the pseudohaploid testis and liver de-novo assemblies from the Seewiesen zebra finch individual. Supplementary Table 4 | All 267 genes on the GRC with information on their Achromosomal origin in taeGut2, number of testis-specific SNVs, methods supporting their GRC linkage, testis/ovary RNA expression of the GRC paralog, testis/ovary protein expression of the GRC paralog, and evolutionary stratum on the GRC. Supplementary Table 5 | Copy number estimates for 61 GRC-linked genes with at least 2 copies on the GRC as estimated from excess coverage in testis. Supplementary Table 6 | Transcriptome analyses of GRC-linked genes showing the number of 'alt' SNVs per transcript with a minimum of 100 reads and an 'alt'/'ref' SNV mapping ratio above 1% in testes and ovary RNA-seq data. Supplementary Table 7 | Enriched gene ontology terms for 167 mapped gene symbols from all 267 GRC-linked genes, and 77 mapped genes from 115 high confidence genes. Supplementary Table 8 | Enrichment analyses of GRC gene orthologs in chicken and human RNA-seq data for testes, ovaries, and other tissues. Supplementary Table 9 | Codon substitution rate analyses for 17 genes with at least 50 **GRC-specific SNVs. Supplementary Data**