Avian W and mammalian Y chromosomes convergently retained dosage-sensitive regulators

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After birds diverged from mammals, different ancestral autosomes evolved into sex chromosomes in each lineage. In birds, females are ZW and males are ZZ, but in mammals females are XX and males are XY. We sequenced the chicken W chromosome, compared its gene content with our reconstruction of the ancestral autosomes, and followed the evolutionary trajectory of ancestral W-linked genes across birds. Avian W chromosomes evolved in parallel with mammalian Y chromosomes, preserving ancestral genes through selection to maintain the dosage of broadly expressed regulators of key cellular processes. We propose that, like the human Y chromosome, the chicken W chromosome is essential for embryonic viability of the heterogametic sex. Unlike other sequenced sex chromosomes, the chicken W chromosome did not acquire and amplify genes specifically expressed in reproductive tissues. We speculate that the pressures that drive the acquisition of reproduction-related genes on sex chromosomes may be specific to the male germ line.

In birds and mammals, one pair of chromosomes differs between males and females. In birds, females are ZW and males are ZZ; in mammals, females are XX and males are XY. The sex chromosomes of birds and mammals are not orthologous: genes that are sex linked in birds are autosomal in mammals, and vice versa1–5. The orthologs of sex-linked genes from emu and ostrich—where two-thirds of the Z chromosome still crosses over with the W chromosome in lengthy pseudoautosomal regions—to the chicken, where the Z and W chromosomes are almost completely differentiated.

The current understanding of the biology and evolution of sex-specific chromosomes is largely based upon the reference sequences of several male-specific Y chromosomes10–15. Vertebrate sex chromosomes commonly contain ampliconic sequences, long stretches of duplicated sequences that have high nucleotide identity4,8,10–15. Resolving these sequences requires a methodology with an extraordinary level of accuracy and precision—specifically, the sequencing of large-insert clones derived from a single haplotype. High-quality, clone-based Y-chromosome reference sequences have identified two major phenomena in the evolution of male-specific Y chromosomes: the acquisition and amplification of testis-expressed gene families that preserve or enhance male fertility11,13,15 and the preservation of widely expressed, dosage-sensitive ancestral genes that may have crucial roles in Turner syndrome and in sexual dimorphism in health and disease10.

Analogous evolutionary pressures are expected to act on female-specific W chromosomes. Genes on the chicken W chromosome was among the most conservative, retaining 4–5% of ancestral genes, while decay was more severe on the mouse Y chromosome, where only 1% of ancestral genes remain10,11. The extent of divergence between the Z and W chromosomes varies widely across birds, from emu and ostrich—where two-thirds of the Z chromosome still crosses over with the W chromosome in lengthy pseudoautosomal regions—to the chicken, where the Z and W chromosomes are almost completely differentiated17.

Although the avian ZW sex chromosomes are the mirror image of the mammalian XY pair with respect to sex, these two chromosome pairs followed parallel evolutionary trajectories. In each lineage, a series of events, most likely inversions on the sex-specific (W or Y) chromosome, suppressed crossing-over between the sex chromosomes, leading to the formation of evolutionary strata5–7. In the absence of crossing-over, the sex-specific W and Y chromosomes diverged from their counterparts, the Z and X chromosomes. The Z and X chromosomes retained 98% of the genes that existed on the ancestral autosomes4,8. In contrast, the sex-specific W and Y chromosomes became subject to genetic decay9. Few ancestral genes remain on mammalian Y chromosomes; the opossum Y chromosome
respond to female-specific selection\(^\text{16}\) and are expected to accumulate genes expressed solely in female-specific tissues\(^\text{17}\). W-linked genes in chicken, turkey, and duck are evolving with significant contributions from purifying selection\(^\text{16}\). In chicken\(^\text{19}\) and flycatcher\(^\text{20}\), the combined expression of Z–W gene pairs in females is comparable to the expression from both Z genes in males, leading some investigators to hypothesize that the surviving ancestral W-linked genes in birds also should be enriched for broadly expressed, dosage-sensitive regulators\(^\text{20}\).

Here we reconstruct the evolutionary trajectory of the genes ancestral to the avian sex chromosomes and examine whether evolutionary pressures analogous to those faced by the mammalian Y chromosomes generated biases in the gene content of female-specific avian W chromosomes. To enable a systematic and comprehensive analysis of gene acquisition and preservation, we produced the first high-quality, clone-based reference sequence from the female-specific chicken W chromosome, supported by physical, linkage, and cytological maps. These sequences, made immediately available in GenBank, have already enabled the design and interpretation of recent studies of avian sex chromosomes\(^\text{18–20}\). We took advantage of our previous reconstruction of the ancestral gene content of the avian sex chromosomes\(^\text{4}\), as well as the candidate W-linked genes reported in draft genome assemblies from several avian lineages\(^\text{18,20,21}\), to extend our parallel analysis across the surviving ancestral genes on the W chromosomes of 14 species of birds.

Genetic decay devastated the chicken W chromosome: only 28 of 685 ancestral genes remain. All of the genes on the chicken W chromosome derive from the ancestral autosomes and are expressed across a broad array of adult and embryonic tissues. Relative to other ancestral genes, surviving Z–W gene pairs on the W chromosomes of chicken and other birds are enriched for dosage-sensitive, broadly expressed genes, under strong purifying selection. We conclude that selection to maintain the ancestral dosage of homologous sex-chromosome gene pairs was the driving force behind the survival of ancestral W-chromosome genes in the chicken and across the avian lineage. Further, we speculate that differences in selective pressures operating on chromosomes in male and female germ lines may explain why no W-linked genes are expressed exclusively in female-specific tissues in the chicken.

## RESULTS

### Sequencing and analysis of the chicken W chromosome

We sequenced the euchromatic portion of the chicken W chromosome (Fig. 1), using the super-resolution methodology that we previously employed on mammalian Y, human X, and chicken Z chromosomes (Online Methods)\(^\text{4,8,10–13,15}\). We obtained a tiling path of 7 Mb in 13 contigs (Fig. 1a–c, Supplementary Table 1, and Supplementary Data 1), containing 28 genes (Fig. 1b). The resulting sequence is accurate to about 1 nt per 36 kb (Online Methods and Supplementary Table 1). We employed two methods to order and orient these contigs (Online Methods). First, we assigned each sequence contig to one of three distinct linkage groups on the radiation hybrid map (Fig. 1a,d and Supplementary Data 2), and, second, we ordered the three radiation hybrid linkage groups along the W chromosome using lampbrush FISH (Fig. 1e,f and Supplementary Fig. 1).

We validated each putative chicken W-chromosome gene by verifying transcriptional activity (Supplementary Data 3) and comparing its ORF to its human ortholog (Supplementary Data 4). All 28 genes on the W chromosome are broadly expressed across adult tissues (Fig. 2). Of the 28 genes, 27 are each present in a single copy on the chicken W chromosome; only HINTW has been amplified into a multicopy family (Supplementary Fig. 2) (refs. 18,22). Ampliconic sequences, which are long stretches of duplicated sequence that have high nucleotide identity, are a common feature of mammalian Y chromosomes\(^\text{11–13}\). The HINTW array is the only ampliconic sequence on the chicken W chromosome, with approximately 40 copies of a 5-kb repeat unit, ranging from 95–99.9% nucleotide identity (Fig. 1a,e and Supplementary Fig. 2).

### Reconstructing the ancestral autosomes

Our previous comparisons of the chicken Z chromosome with the orthologous human autosomes identified a set of 671 ancestral genes that were present on the ancestral amniote autosomes that became the chicken Z and W sex chromosomes (Table 1 and Supplementary Data 1)
Species as lower-confidence predictions. In these species, candidate W-chromosome evolution across all birds (including galloanserae (chicken)—allowing broad conclusions about paleognathae (emu), neoaves (crested ibis and collared flycatcher), and ostrich that could allow us to determine whether gene gains or losses occurred before the common ancestor of extant birds (Fig. 3a and Online Methods). With this revised reconstruction, we identified 685 genes as present on the ancestral autosomes that became the avian Z and W sex chromosomes (Table 1 and Supplementary Table 2).

Table 2) (ref. 4). We revisited our reconstruction of the gene content of the ancestral autosomes in light of recent improvements to the annotations of the chicken and human genomes, as well as newly published genome sequences from anole lizard23, American alligator24,25, and ostrich26 that could allow us to determine whether gene gains or losses occurred before the common ancestor of extant birds (Fig. 3a and Online Methods). With this revised reconstruction, we identified 685 genes present on the ancestral sex chromosomes that became the avian Z and W sex chromosomes (Table 1 and Supplementary Table 2).

Ancestral Z–W gene pairs in other avian species

We also searched for surviving ancestral Z–W gene pairs among the published W-linked genes from the 13 other avian species with published female genomes but without clone-based assemblies of the W chromosome (Fig. 3a)7,18,21. We stratified these candidate W-linked genes into two groups, on the basis of the amount of information used to identify W-linked genes for each species.

In three species—emu27, crested ibis28, and collared flycatcher20—candidate W-linked genes had been ascertained by comparing male and female genome assemblies to identify female-specific sequences7,20, but W linkage was not confirmed by PCR or other additional mapping information. Fifty additional candidate W-linked genes were members of our set of 685 ancestral genes in one or more of these three species (Fig. 3b and Supplementary Table 3). Combining these 50 genes with the 28 genes from the chicken W chromosome gives a total of 78 genes of intermediate or high confidence in one or more of these three species (Fig. 3a).

We regarded candidate W-chromosome genes in the ten remaining species7 as lower-confidence predictions. In these species, candidate W-linked genes had been predicted directly from a female genome assembly. Without the control of a male genome assembly, two factors could potentially confound these gene predictions and diminish our ability to detect enrichments among surviving Z–W gene pairs. First, sequencing biases cause local variations in genome coverage that make it difficult to accurately identify the twofold changes in read depth that distinguish autosomal sequences from sex-linked sequences. Second, autosomal paralogs of Z-linked genes may appear similar to genuine Z–W gene pairs. Including predictions from these 10 species yielded another 79 ancestral genes, for a total of 157 putative ancestral genes in one or more of all 14 species (Fig. 3b and Supplementary Table 3).

Table 1 Reconstruction of the gene content of the autosomal ancestors of the chicken sex chromosomes

| Loss | Ancestral | Gain |
|------|-----------|------|
| Bellott et al.4 | 49 | 671 | 493 |
| Not unique | 0 | 0 | –427 |
| Updated annotations, new | +5 | +5 | +1 |
| Updated annotations, withdrawn | –7 | –49 | –18 |
| Gained before birds diverged | 47 | 627 | 49 |
| Lost after birds diverged | –19 | +19 | 0 |
| Total | 28 | 685 | 10 |

See also Supplementary Table 2.

Strategies for gene survival on sex-specific chromosomes

On the male-specific Y chromosomes of mammals, two evolutionary strategies contributed to gene survival despite widespread genetic decay: the retention and amplification of testis-expressed gene families and the conservation of ancestral X–Y gene pairs to maintain comparable expression between males and females10,29. Although analogous strategies should act on W chromosomes, and W chromosomes are expected to accumulate genes expressed solely in female-specific

Figure 3 Ancestral Z–W gene pairs from 14 avian species. (a) Phylogenetic tree of the species in this study, with branches colored to highlight relationships among species. Humans diverged from birds 325 million years ago. Green anole lizard and American alligator diverged from birds 275 and 219 million years ago, respectively, and were used to resolve gene gains and losses between birds and mammals. Birds diverged from each other starting around 120 million years ago (yellow). The branches of major avian lineages are colored: green, galloanserae; purple, neovaves; red, paleognathae. (b) Euler diagram showing overlapping sets of ancestral Z–W gene pairs identified in chicken (dark pink); 4 species (chicken, collared flycatcher, crested ibis, and emu) (medium pink); and all 14 published female avian genomes (light pink), as subsets of all 685 ancestral Z genes (light yellow). See also Supplementary Table 3.
We found that the female-specific chicken W chromosome has no genes that are exclusively expressed in sex-specific tissues (Fig. 2). In contrast to ampliconic genes on mammalian X and Y chromosomes and even the chicken Z chromosome, the sole ampliconic gene on the W chromosome, HINTW, is broadly expressed (Fig. 2). Therefore, the first strategy cannot explain the survival of ancestral Z–W gene pairs in the chicken.

Despite widespread genetic decay on the sex-specific chromosome, dosage-sensitive genes functioning across many tissues and cell types may survive because their loss would have too great an impact on reproductive fitness and even viability. We looked for evidence that selection to maintain the correct dosage of ancestral genes might spare W-linked genes from genetic decay.

Z–W gene pairs are more dosage sensitive

First, we examined whether surviving Z–W gene pairs show signs of dosage sensitivity. We compared each of these three lists of surviving ancestral Z–W gene pairs (from chicken alone, or 4 species, or all 14 species) to the other ancestral genes, reanalyzing published data sets for evidence that Z–W pair genes systematically differ from the ancestral genes on the Z chromosome that lack W homologs with regard to dosage sensitivity, breadth of expression, and intensity of purifying selection (Fig. 4).
probability of haploinsufficiency for the human genome\textsuperscript{32}, mapped on
to their orthologs in the chicken genome. We found that the human orthologs of surviving Z–W gene pairs were more likely to be haploinsufficient than the human orthologs of ancestral Z-chromosome genes that lack W-chromosome homologs (one-tailed Mann–Whitney U test: chicken, $P < 5.8 \times 10^{-5}$; 4 species, $P < 1.6 \times 10^{-3}$; 14 species, $P < 8.34 \times 10^{-4}$) (Fig. 4a and Supplementary Table 2).

Additional evidence for the dosage sensitivity of specific Z–W gene pairs comes from human congenital disorders. Of the 28 chicken Z–W pairs, 3 have human orthologs implicated in congenital disorders caused by heterozygous loss-of-function mutations. Haploinsufficiency for TCF4 is responsible for Pitt–Hopkins syndrome.\textsuperscript{36} Parke–Weber syndrome is caused by heterozygous inactivating mutations in RASA1. Cornelia de Lange syndrome results from haploinsufficiency for NIPBL.\textsuperscript{37} To assess the statistical likelihood of finding 3 demonstrably haploinsufficient human orthologs among these 28 Z–W pairs, we examined all 4,562 human phenotypes with a known molecular basis catalogued in Online Mendelian Inheritance in Man (OMIM). Specifically, we searched for entries containing the word “haploinsufficient” or “haploinsufficiency” and found 238 disorders attributed to haploinsufficiency for a human gene. Three of 11 phenotypes mapping to human orthologs of Z–W pair genes were due to haploinsufficiency, whereas only 23 of 4,551 phenotypes in the rest of the human genome were due to haploinsufficiency (one-tailed Fisher’s exact test, $P < 0.017$). Taken together, the elevated haploinsufficiency probabilities and enrichment for human disorders caused by haploinsufficiency suggest that Z–W pairs are enriched for haploinsufficient genes.

Z–W gene pairs are more broadly expressed

Z–W gene pairs functioning across many tissues and cell types face additional selective constraints, which could prevent the loss of the W-linked copy, leading to an enrichment for broadly expressed genes among surviving Z–W pairs. Across adult chicken tissues, we observed that the Z homologs of Z–W gene pairs are more broadly expressed than ancestral Z genes that lack W homologs in chicken and in four species (one-tailed Mann–Whitney U test: chicken, $P < 2.1 \times 10^{-3}$; 4 species, $P < 3.8 \times 10^{-3}$; 14 species, $P < 0.059$) (Fig. 4b and Supplementary Table 2). This increased breadth of expression extends to the human orthologs of Z–W gene pairs; the human orthologs of Z–W gene pairs are more broadly expressed than the human orthologs of ancestral Z genes that lack W homologs in chicken and in four species (one-tailed Mann–Whitney U test: chicken, $P < 1.6 \times 10^{-3}$; 4 species, $P < 0.047$; 14 species, $P < 0.13$) (Supplementary Fig. 3 and Supplementary Table 2). We conclude that the autosomal precursors of Z–W gene pairs were broadly expressed across adult tissues in the common ancestor of birds and mammals.

This breadth of expression also extends to the earliest stages of development. Ancestral Z–W pairs are more highly expressed in chicken blastocysts than are ancestral Z genes that lack W orthologs (one-tailed Mann–Whitney U test: chicken, $P < 7.7 \times 10^{-7}$; 4 species, $P < 1.1 \times 10^{-3}$; 14 species, $P < 2.8 \times 10^{-3}$) (Fig. 4c and Supplementary Table 2). We also examined the human orthologs of ancestral Z genes in published human embryonic transcriptome data.\textsuperscript{34} We found that the human orthologs of ancestral Z–W pairs are more highly expressed in human blastocysts than are human orthologs of ancestral Z genes that lack W homologs in chicken and in 14 species (one-tailed Mann–Whitney U test: chicken, $P < 5.4 \times 10^{-3}$; 4 species, $P < 0.087$; 14 species, $P < 0.011$). We conclude that the autosomal precursors of the Z–W pairs were more broadly expressed across developmental time as well as across tissues in the amniote ancestor.

Z–W gene pairs are subject to stronger purifying selection

Previous comparisons among Z–W pairs in chicken, turkey, and duck showed that purifying selection has contributed significantly to the evolution of W-linked genes.\textsuperscript{18} We reasoned that if Z–W gene pairs are haploinsufficient, alleles that impair the function of Z-linked homologs should be detrimental in both males and females, so that the Z homologs of Z–W gene pairs should also show signs of strong purifying selection. We examined Ensembl chicken ortholog alignment data for evidence that the Z-linked homologs of Z–W gene pairs were subject to stronger purifying selection than other ancestral Z-linked genes. In comparison to ancestral genes on the Z chromosome that lack W homologs, the Z-linked homologs of Z–W gene pairs have a reduced ratio of nonsynonymous to synonymous substitution (dN/dS) rates when chicken genes are compared to orthologs in duck for chicken and 14 species (one-tailed Mann–Whitney U test: chicken, $P < 0.022$; 4 species, $P < 0.052$; 14 species, $P < 3.6 \times 10^{-3}$) (Fig. 4d and Supplementary Table 2), and for all three groups both in collared flycatcher (one-tailed Mann–Whitney U test: chicken, $P < 8.6 \times 10^{-5}$; 4 species, $P < 7.7 \times 10^{-5}$; 14 species, $P < 2.9 \times 10^{-5}$) (Fig. 4e and Supplementary Table 2), and zebra finch (one-tailed Mann–Whitney U test: chicken, $P < 9.5 \times 10^{-5}$; 4 species, $P < 1.3 \times 10^{-5}$; 14 species, $P < 1.6 \times 10^{-3}$) (Fig. 4f and Supplementary Table 2).

We conclude that, on avian W chromosomes, strong purifying selection has preserved a subset of ancestral genes that are more widely expressed and more dosage sensitive, just as it has on mammalian Y chromosomes.

Functional coherence of Z–W gene pairs

We recently characterized human X–Y gene pairs as performing an array of functions in gene expression and regulation, suggesting that X–Y pair genes could govern the expression of targets throughout the genome.\textsuperscript{10} We asked whether our high-confidence set of ancestral genes that survived on the chicken W chromosome could be characterized as carrying out regulatory functions similar to the survivors on mammalian Y chromosomes. In comparison to ancestral genes on the Z chromosome that lack W homologs, Z–W pair genes are enriched for Gene Ontology (GO) annotations such as nucleic acid binding, nucleus, and transcription (Supplementary Table 4)
suggest regulatory functions. We therefore looked more closely at the molecular functions of the 28 chicken Z–W pairs.

We observe that, in addition to the functions in transcription, translation, and protein stability attributed to mammalian X–Y pairs, chicken Z–W pairs are also predicted to act in protein secretion and signal transduction pathways (Fig. 5) (refs. 38,39). Specifically, several Z–W pairs share annotations that suggest roles in transducing TGF-β signaling (SMAD2, SMAD7, and NEDD4L) and modulating Wnt signaling (UBE2R2, HINT1, and SPIN1). Interactions between the TGF-β and Wnt pathways are critical for axis and pattern formation in early development40, and, as morphogens, each can induce different cellular responses as a function of concentration, or dosage.

**DISCUSSION**

The preservation of broadly expressed, dosage-sensitive genes by purifying selection on avian W chromosomes offers a striking example of convergent evolution of the ZW and XY sex-chromosome systems. This survival strategy has been documented across diverse sets of ancestral Y-linked genes 10,30,31. In Drosophila miranda, surviving gene pairs on the Neo-X and Neo-Y chromosomes are expressed at higher levels and across more tissues than those genes that were lost to decay30. Likewise, in mammals, the surviving ancestral X–Y gene pairs are more broadly expressed across developmental time and adult tissues10. This strongly suggests that genes whose expression is required across a broad array of tissues are subject to greater constraints on gene dosage, making even the loss of a single copy costly. Like the Z–W gene pairs in birds we report here, the surviving X–Y gene pairs in mammals had higher predicted probabilities of haplinsufficiency, as well as ties to human syndromes caused by changes in gene dosage10. Similarly, the surviving X–Y gene pairs of both D. miranda and threespine stickleback were enriched for genes encoding proteins with many partners in protein–protein interaction networks30,31. Macromolecular complexes are sensitive to imbalances in the stoichiometry of their components, and an abundance of interactions is correlated with dosage sensitivity41. The repeated finding, across both female-specific (W) and male-specific (Y) chromosomes, that surviving ancestral genes are enriched for dosage-sensitive genes functioning across many tissues and cell types, contradicts the dire predictions of the imminent demise of sex-specific chromosomes due to genetic decay42. Purifying selection has been effective at preserving the ancestral dosage of critical genes on the sex-specific chromosome, even in the absence of crossing-over, through hundreds of millions of years of evolution.

Despite the evolutionary similarities, we note that the chicken W chromosome is remarkably divergent from all sequenced Y chromosomes, in that it lacks any genes expressed specifically in sex-specific organs or tissues. The gene content of mammalian Y chromosomes is frequently dominated by massively amplified testis-specific gene families that did not originate on the autosomal ancestors, even though they may have X-linked homologs11,13–15. In mammalian Y chromosomes, ancestral genes that narrowed their expression to male-specific tissues and became amplified into multicopy gene families were preserved across a greater number of species10.

The relative simplicity of the W chromosome, with only broadly expressed ancestral genes and only one multicopy gene family, may be because its transmission is restricted to the female germ line. X, Y, and Z chromosomes pass through the male germ line, and all have acquired and amplified testis-expressed gene families4,8,11–13,15,28. This marked absence of acquired genes that are specifically expressed in the ovary or other female-specific tissues, even on a female-specific chromosome, suggests that, in amniotes, there is greater pressure to preserve or enhance male reproductive functions. Meiotic drive, which pits each chromosome against its homolog in a competition for transmission to the next generation, is one source of pressure on reproduction in males and females. However, there are more opportunities for meiotic drive to exert pressure during spermatogenesis than during oogenesis. Developing sperm are connected by cytoplasmic bridges, forming a syncytium that provides a venue for competition both during and after meiosis. For example, the ampicolous gene families on the X and Y chromosomes of the mouse are implicated in meiotic drive11,43–45, even though they are expressed predominantly in post-meiotic germ cells46. During oogenesis, the arena for competition is narrower; any competition between homologous chromosomes must be resolved by the first meiotic division, when homologs separate and one is ejected into a polar body. Thus, W chromosomes, which only pass through the female germ line, may be subject to less disruptive selective pressures than those experienced by X, Y, and Z chromosomes in the male germ line. The complete sequences of W chromosomes from other birds, or the independently evolved Z and W chromosomes of snakes, could show whether the absence of acquired gene families that we observe in the chicken is a general feature of female-specific chromosomes.

We previously proposed that the dozen broadly expressed, dosage-sensitive genes on the human Y chromosome, along with their X-linked homologs that escape X-chromosome inactivation, are essential for the viability of 46,XY fetuses10. Two key observations support this hypothesis: first, that X–Y gene pairs are enriched for genes expressed in early development10 and, second, that 99% of human 45,X conceptuses are not viable, while the remainder are often mosaic for all or part of a second sex chromosome47–49.

Parallel lines of evidence in the chicken lead us to propose that the single-copy chicken Z–W pairs function to ensure female survival by providing the correct dosage of genes, especially those functioning in critical signaling pathways during early embryonic development. All 27 single-copy Z–W pairs are expressed in the developing chicken blastoderm, and the combined expression of Z–W gene pairs in females is comparable to the expression of the two Z homologs in males19. Both 2A:ZZW and 2A:Z0 aneuploid embryos have been observed in chicken at the blastocyst stage50, but these embryos do not survive past 4 to 5 d of incubation50,51, and sex chromosome aneuploidy is widely regarded as embryonic lethal in the chicken. Considering the severity of the three congenital developmental disorders linked to human orthologs of Z–W gene pairs, we conclude that hemizygosity for all Z–W gene pairs would likely result in early lethality.

In addition to their critical roles in maintaining embryonic viability, chicken Z–W and human X–Y gene pairs may have broader roles in sex determination and sexual dimorphism. Evidence of cell-autonomous sex determination in chickens has emerged from the study of lateral gynandromorphs52, along with sexually dimorphic gene expression that precedes gonadal differentiation19,52. This leads us to speculate that one or more of the broadly expressed regulators found on the chicken W chromosome may have evolved to direct aspects of female fate in cell types across the body. In mammals, we are just beginning to understand the consequences of a fundamental sexual dimorphism, at the cellular level, arising from genetic differences between developmental regulators encoded by the X and Y chromosomes. In humans, for example, somatic mutations in the X-linked members of X–Y gene pairs were recently linked to the increased incidence of cancer in human males53. It will surely be of interest to compare and contrast...
birds and mammals, taking advantage of the parallel evolutionary trajectories of avian ZW and mammalian XY chromosomes, to uncover new paradigms for understanding the regulation and development of sexual dimorphism in both health and disease.

METHODS

Methods, including statements of data availability and any associated accession codes and references, are available in the online version of the paper.

AUTHOR CONTRIBUTIONS

D.W.B., H.S., W.C.W., A.G.C., E.G., R.K.W., and D.C.P. planned the project. D.W.B. and T.-J.C. performed RH mapping. D.W.B. and H.S. performed clone mapping. D.W.B., T.-J.C., D.W.B., H.S., W.C.W., A.G.C., E.G., R.K.W., and D.C.P. performed the project (grant 16-04-01823a).

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COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Single-haplotype iterative mapping and sequencing. We employed the single-haplotype iterative mapping and sequencing (SHIMS) strategy to assemble the chicken W chromosome sequence from 41 BAC and 123 fosmid clones (Supplementary Table 1). These clones were obtained from four BAC libraries (CH261, TAM31, TAM32, and TAM33) (refs. 54,55) and two fosmid libraries (J_AD and J_AE)56, which provide combined ~16-fold coverage of the W chromosome. The single female red jungle fowl of the UCD001 line (RFJ 256) (ref. 56). Thirty-seven BACs and 5 fosmids were sequenced on ABI 3730 machines at the McDonnell Genome Institute, and 4 BACs and 118 fosmids were sequenced on an Illumina MiSeq instrument at the Whitehead Institute; see individual GenBank records for assembly details. We previously used the SHIMS strategy to produce finished sequence from mammalian Y, human X and chicken Z chromosomes4,8,10–13,15. The major steps in the SHIMS strategy are outlined below.

Marker generation. We identified female-specific sequence contigs in the draft assembly of the chicken genome using existing genetic linkage data 56,57, direct sequencing of flow-sorted W-chromosome DNA, electronic searches for close homologs of Z-linked gene sequences4, and electronic subtraction using short-read genomic data from a male White Leghorn58. We used these sequences to develop sequence-tagged sites (STS) and verified that they were female specific by PCR on DNA from a male and female red jungle fowl.

Initial BAC selection and sequencing. We identified large-insert BAC and fosmid clones and organized them into contigs of overlapping clones on the basis of (i) high-density filter hybridization using pools of overgo probes, (ii) electronic mapping of clone end sequences to female genomic sequences, and (iii) BAC fingerprint contig analysis. We confirmed the resulting contigs by PCR using female-specific STS markers and selected tiling paths for sequencing.

Distinguishing repeat copies and finding true tiling paths. We scrutinized overlaps between clones within repetitive regions for sequence differences or sequence family variants (SFVs). The presence of SFVs indicates that the clones belong to distinct copies of the same repeat family, and we used SFV patterns to identify clones that truly overlap. This produced new tiling paths. We reiterated this process until all overlaps were consistent.

Extension and joining of large-insert clone contigs. We identified clones that extend outward from or link existing contigs using high-density filter hybridization and electronic mapping of clone end sequences.

Gap closure. Regions composed of repeats with units less than 10 kb and greater than 99% identity frustrate the assembly of individual clones and are not well represented in our assemblies. These regions include both gene-poor regions, like centromeres, telomeres, and heterochromatin, and gene-rich regions, such as the HINTW array. No current technology is able to access these regions. Wherever possible, we attempted to find the boundaries of these arrays and obtain a representative repeat unit.

Calculation of sequence accuracy. The initial error rate estimated for clone sequencing and assembly was 1 in 28 kb. However, as 23% of our sequence was covered redundantly by two BACs, we were able to identify and resolve all discrepancies in redundantly covered regions so that the error rate for these regions was zero. Therefore, the final error rate was estimated to be 0.77 ± 0.23 × 0 = 1/36,000, or 1 in 36 kb.

Ordering and orienting sequence contigs. The structure of the chicken W chromosome presents a unique challenge to traditional techniques for chromosome mapping and assembly. Islands of euchromatin are separated by massive heterochromatic tandem arrays, each composed of one of three families of genome-typical interspersed repeats59–61. We employed two independent and complementary methods to order and orient the 13 contigs of our sequence map along the W chromosome.

Radiation hybrid mapping. We tested 119 STS markers (Fig. 1d and Supplementary Data 2) on the ChickRH6 panel62, a 6,000-rad panel consisting of 90 hybrid clones, and constructed an RH map using RHMAPPER63. We thereby assigned each of the 13 sequence contigs to one of three distinct linkage groups on the RH map (Fig. 1a-d).

Lampbrush FISH. We ordered the three RH linkage groups along the W chromosome using lampbrush FISH (Fig. 1e-h and Supplementary Fig. 1). The lampbrush W chromosome features a series of seven condensed heterochromatic chromomeres along its axis; these are numbered in ascending order from the tip of the long arm (Fig. 1e). The three major repetitive sequence families of the W chromosome were previously mapped to chromomeres 1, 3, 5, and 6 by lampbrush FISH59,60 (Fig. 1e). We found that the three remaining chromomeres—2, 4, and 7—correspond to the three radiation hybrid linkage groups (Fig. 1a,c,e,f and Supplementary Fig. 1). The first linkage group contains 3 contigs, spans 2 Mb, contains 8 genes, and corresponds to chromomere 2, on the long arm of the W chromosome (Fig. 1e,f and Supplementary Fig. 1). This linkage group terminated in sequences from the XhoI repeat family (Fig. 1e and Supplementary Table 1). The XhoI repeat family was previously mapped to the adjacent chromomere 3 (ref. 60), suggesting that we captured the border of this heterochromatic array (Fig. 1e).

The second linkage group contains 6 contigs, spans 3 Mb, contains 12 genes, and corresponds to chromomere 4 (Fig. 1e,g, Supplementary Fig. 1, and Supplementary Table 1), near the centromere of the W chromosome64. Consistent with proximity to the centromere, single-copy markers assigned to this linkage group were retained at higher frequency in the radiation hybrid panel than were markers in the other two linkage groups (Fig. 1d).

The third linkage group consists of 4 contigs, spans 2 Mb, contains 8 genes, and corresponds to chromomere 7, on the short arm of the W chromosome near the pseudoautosomal region where the Z and W chromosomes cross over during female meiosis (Fig. 1e,h, Supplementary Fig. 1, and Supplementary Table 1). Consistent with the results of lampbrush FISH, the proximal end of this linkage group contained the only ampliconic sequence on the W chromosome, a tandem array of HINTW genes (Fig. 1c and Supplementary Fig. 1). The HINTW array was previously mapped to the short arm of the W chromosome by metaphase FISH65. The third linkage group terminated in the pseudoautosomal region, which contained no genes but instead consisted entirely of telomeric and subtelomeric repeats shared with the Z chromosome (Fig. 1e and Supplementary Fig. 1).

Chromosomal FISH analyses. We performed FISH assays on Gallus gallus domesticus lampbrush chromosomes as previously described64. Briefly, lampbrush chromosomes were manually isolated from oocyte nuclei, dehydrated in 96% ethanol, air dried, and treated with RNase A. BAC probes were labeled with digoxigenin and denatured together with unlabeled competitor DNA and the lampbrush chromosomes before hybridization. Probes were detected with antibodies against digoxigenin conjugated to Cy3 (JacksonImmunoResearch Laboratories, 200-162-156) at a 1:400 dilution. Chromosomes were stained with DAPI and imaged on a fluorescence microscope. Experimental procedures involving chicken oocytes were approved by the Saint Petersburg State University Ethics Committee (statement 131-03-2).

Interspersed repeats. Interspersed repeats were electronically identified with RepeatMasker.

Identification of genes and transcription units. We identified genes and transcripts as previously described4,10. Briefly, we used TWINSCAN55,56 with human as the informant genome and EST sequences from the BBSRC ChickEST database67, supplemented by our own ESTs from adult ovary (SRP000997). We specifically searched for homologs of all genes found in the finished sequence of the chicken Z chromosome to detect ancestral genes as well as any genes co-acquired by the Z and W chromosomes. We validated transcription of predicted genes by RT–PCR and capillary sequencing, as well as 454 sequencing of ovary cDNA and Illumina-based RNA–seq (PRNA204941).

Dot plots. Triangular dot plots (representing intrachromosomal sequence similarity) and square dot plots (representing interchromosomal sequence similarity) were generated with a custom Perl script (see URLs).
Reconstructing ancestral autosomes. Our previous comparisons of the chicken Z chromosome with orthologous regions of human autosomes identified 720 genes that were present on the ancestral amniote chromosomethat became the chicken Z and W sex chromosomes (Table 1 and Supplementary Table 2) (ref. 4). Of these 720 genes, 671 had syntentic orthologs in both human and chicken. The other 49 genes had syntentic orthologs in human and an outgroup species (amphibians or fish), but not in chicken, indicating that these genes were lost along the lineage leading to chicken (Table 1 and Supplementary Table 2). We also identified 66 distinct families of genes (493 genes in all) that had been added to the chicken Z chromosome but were not present on the ancestral amniote autosomes (Table 1 and Supplementary Table 2) (ref. 4).

Previously, we relied on assignments of chicken and human orthologs in the Ensembl database (version 52). We reexamined all 786 distinct genes or families (720 ancestral plus 66 acquired) in light of recent improvements to annotations of the chicken and human genomes (Ensembl version 80) (ref. 35). This allowed us to eliminate genes that represented errors or redundancies in previous annotations of the chicken and human genomes and to add genes that had been overlooked by previous annotation efforts (Table 1 and Supplementary Table 2). This reduced the number of ancestral genes maintained on the chicken Z chromosome from 671 to 627 (Table 1 and Supplementary Table 2). This also reduced the number of ancestral genes evidently lost from the chicken Z chromosome from 69 to 58 (Table 1 and Supplementary Table 2). Of the 497 genes that had been lost from the chicken Z chromosome, 19 maintained a syntenic ortholog in the ostrich, indicating that these genes were lost along the lineage leading to birds (Table 1 and Supplementary Table 2). Combining the 627 ancestral genes maintained on the chicken Z chromosome with the 19 genes lost from the chicken Z chromosome after the radiation of birds, plus the 39 genes gained by the ancestral autosomes before birds diverged, yields a total of 685 genes present on the ancestral autosomes that became the avian Z and W sex chromosomes (Table 1 and Supplementary Table 2).

W-linked gene expression. We quantified the abundances of chicken transcripts from the Chickpress RNA–seq data set (PRJNA204941) using kallisto version 0.42.3 (ref. 69) and edgeR (ref. 70). We normalized the transcript abundances for each gene to the abundance in the highest expressing tissue for that gene.

OMIM. We downloaded the full text of OMIM (ref. 54) and searched entries for “haploinsufficiency” or “haploinsufficiency,” limiting our search to phenotypes with a known molecular basis. We examined each of the resulting entries to verify that there was evidence that the phenotype was caused by haploinsufficiency.

Functional annotation. We mapped published functional annotation data onto our set of 685 ancestral genes and their human orthologs. For expression breadth, we normalized the expression of each gene to the highest RPKM in any tissue and took the average expression across all tissues. We used UniProt annotations to identify chicken Z–W pair genes involved in regulatory processes.

Statistics. We tested whether the human orthologs of Z–W gene pairs were enriched for phenotypes caused by haploinsufficiency, relative to the rest of the human genome, using a one-tailed Fisher’s exact test, because of the small number of Z–W gene pairs whose orthologs are annotated in OMIM. We tested for enrichments in the annotations of ancestral Z–W gene pairs identified in chicken, 4 species (chicken, collared flycatcher, crested ibis, and emu), and all 14 published female avian genomes versus the remainder of ancestral Z genes using one-tailed Mann–Whitney U tests. We report all of our comparisons, and, in every case, all three classes of Z–W pairs differed from other ancestral genes in the expected direction, making correction for multiple comparisons unnecessary. We attribute the reduced significance for comparisons involving the sets of 4 and 14 species to noise from low-confidence gene predictions in these species.

The exact numbers used to calculate the P values for Figure 4, along with the associated test statistic, U, are as follows. The human orthologs of ancestral Z–W pairs had a higher probability of haploinsufficiency than other ancestral Z genes. Chicken Z–W pairs n = 17, other ancestral Z genes n = 443, P = 5.8 × 10−5, U = 5,840.5; 4–species Z–W pairs n = 49, other ancestral Z genes n = 411, P = 1.6 × 10−4, U = 12,666; 14-species Z–W pairs n = 103, other ancestral Z genes n = 537, P < 8.3 × 10−4, U = 22,122.5. The chicken Z orthologs of ancestral Z–W pairs were more broadly expressed in adult chicken tissues than other ancestral Z genes. Chicken Z–W pairs n = 25, other ancestral Z genes n = 511, P = 2.1 × 10−3, U = 8,561; 4–species Z–W pairs n = 70, other ancestral Z genes n = 466, P = 3.8 × 10−3, U = 19,546; 14-species Z–W pairs n = 138, other ancestral Z genes n = 398, P = 0.069, U = 29,283. The chicken Z orthologs of ancestral Z–W pairs were more highly expressed in chicken blastocysts than other ancestral Z genes. Chicken Z–W pairs n = 28, other ancestral Z genes n = 613, P = 7.7 × 10−7, U = 13,188; 4–species Z–W pairs n = 78, other ancestral Z genes n = 563, P < 1.1 × 10−3, U = 26,684; 14–species Z–W pairs n = 156, other ancestral Z genes n = 485, P = 2.8 × 10−3, U = 43,410.5. The chicken Z orthologs of ancestral Z–W pairs had a reduced dN/dS ratio in alignments with their orthologs in duck. Chicken Z–W pairs n = 26, other ancestral Z genes n = 560, P = 0.022, U = 5,580.5; 4–species Z–W pairs n = 74, other ancestral Z genes n = 512, P = 0.052, U = 16,728.5; 14–species Z–W pairs n = 149, other ancestral Z genes n = 437, P < 3.6 × 10−3, U = 27,753. The chicken Z orthologs of ancestral Z–W pairs had a reduced dN/dS ratio in alignments with their orthologs in collared flycatcher. Chicken Z–W pairs n = 25, other ancestral Z genes n = 582, P < 8.6 × 10−5, U = 4,048; 4–species Z–W pairs n = 72, other ancestral Z genes n = 535, P < 7.7 × 10−7, U = 13,971; 14–species Z–W pairs n = 149, other ancestral Z genes n = 458, P < 2.9 × 10−5, U = 26,636. The chicken Z orthologs of ancestral Z–W pairs had a reduced dN/dS ratio in alignments with their orthologs in zebra finch. Chicken Z–W pairs n = 24, other ancestral Z genes n = 568, P < 9.5 × 10−7, U = 3,750.5; 4–species Z–W pairs n = 72, other ancestral Z genes n = 520, P < 1.3 × 10−4, U = 13,741.5; 14–species Z–W pairs n = 149, other ancestral Z genes n = 443, P < 1.6 × 10−4, U = 26,476.5.

Data availability. Genbank accession numbers for BAC and fosmid sequences are listed in Supplementary Table 1. Z and W transcript sequences assembled from PRJNA204941 have been deposited at DDBJ, EMBL, and GenBank under accession GENL00000000. The version described in this paper is the first version, GENL01000000.
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