Sex-Biased Gene Expression on the Avian Z Chromosome: Highly Expressed Genes Show Higher Male-Biased Expression

Sara Naurin, Dennis Hasselquist, Staffan Bensch, Bengt Hansson*

Department of Biology, Lund University, Lund, Sweden

Abstract
Dosage compensation, the process whereby expression of sex-linked genes remains similar between sexes (despite heterogamety) and balanced with autosomal expression, was long believed to be essential. However, recent research has shown that several lineages, including birds, butterflies, monotremes and sticklebacks, lack chromosome-wide dosage compensation mechanisms and do not completely balance the expression of sex-linked and autosomal genes. To obtain further understanding of avian sex-biased gene expression, we studied Z-linked gene expression in the brain of two songbirds of different genera (zebra finch, Taeniopygia guttata, and common whitethroat, Sylvia communis) using microarray technology. In both species, the male-bias in gene expression was significantly higher for Z than for autosomes, although the ratio of Z-linked to autosomal expression (Z:A) was relatively close to one in both sexes (range: 0.89–1.01). Interestingly, the Z-linked male-bias in gene expression increased with expression level, and genes with low expression showed the lowest degree of sex-bias. These results support the view that the heterogametic females have up-regulated their single Z-linked homologues to a high extent when the W-chromosome degraded and thereby managed to largely balance their Z:A expression with the exception of highly expressed genes. The male-bias in highly expressed genes points towards male-driven selection on Z-linked loci, and this and other possible hypotheses are discussed.

Introduction
Many mammals and birds, and several important genetic model organisms, e.g. Drosophila spp. and Caenorhabditis elegans, have sex chromosomes that differ in gene content. In male heterogametic species (where females are XX and males either XY or X0), the X chromosome holds a large number of functional genes whereas in many species the Y chromosome is degraded due to cessation of recombination, leading to accumulation of deleterious mutations and loss of gene functionality [1–4]. Despite unequal sex chromosome copy numbers in males and females, the expression of X-linked genes is not much lower in the heterogametic males than in homogametic females in some species, including in Drosophila melanogaster, C. elegans, Anopheles gambiae, Tribolium castaneum, and possibly also in mammals [5–13] [see [10,14] for a contrasting view in mammals]. Moreover, in at least some tissues of Drosophila, Anopheles and mammals, mean expression on the single X chromosome in males is similar to that on autosomes [5–7,10,12,13,15]. The mechanisms resulting in this balance in expression between the sexes, and between sex-linked and autosomal expression, are referred to as dosage compensation [16–19]. It is believed that dosage compensation occurs because changes in gene copy number might otherwise have detrimental effects on gene expression in crucial genetic networks [20–22]. Dosage compensation was long believed to be essential although recent research in e.g. birds, butterflies, sticklebacks and platypus, clearly show that the sex chromosome to autosomal expression ratios (X:A or Z:A) of less than one can frequently be tolerated in the heterogametic sex [23–30] [reviewed in 19].

Sophisticated dosage compensation mechanisms, such as sex chromosome inactivation and hyper-transcription, have obviously evolved to an effective and chromosome-wide state in some male heterogametic systems, such as Drosophila, C. elegans, Tribolium and potentially mammals [5–9,31–33] [but see 34], and similar although transient mechanisms have been suggested in both platypus (XY) and chicken (ZW) [27,35,36]. This indicates the importance of retaining original expression levels of sex-linked genes in the heterogametic sex as the non-recombining sex chromosome degrades. However, recent genome-wide gene expression studies based on microarray and RNA sequencing data in both male and female heterogametic systems clearly demonstrate an overrepresentation of sex-biased genes on the sex chromosomes, and X:A or Z:A expression ratios of less than one in the heterogametic sex [8,23–30,37–41] [reviewed in 19]. This shows that complete dosage compensation is not an obligatory by-product of sex chromosome degradation and suggests that in several organisms the heterogametic sex either is not strongly negatively affected by inefficient dosage compensation or achieves sufficient dosage compensation without up-regulating the expression of their X- or Z-linked genes to the same levels as in the
homogametic sex [18,19,42,43]. To understand the complexity and evolution of sex-biased gene expression and dosage compensation mechanisms, additional gene expression data analysed from novel perspectives are clearly needed [10,11,18,19,42–44]. This is important since sex-biased gene expression has wide-ranging effect on the phenotype and is expected to be tightly linked to reproduction through natural and sexual selection [19,43–55].

We have previously studied the extent of sex-biased gene expression in two passerine birds, the zebra finch (Taeniopygia guttata) and the common whitethroat (Sylvia communis) [29]. These two species share a common ancestor approximately 24–51 million years ago [36,37]. We found that genes with male-biased gene expression were overrepresented on the Z chromosome when analysing brain tissue of the zebra finch and the common whitethroat [29]. A high proportion, 39%, of the 509 ESTs that were significantly sex-biased in the zebra finch, and 92% of the 345 ESTs that were sex biased in the common whitethroat, were Z-linked and male-biased [29]. Similar patterns have been described also in other zebra finch expression studies [25,26] as well as in chicken [24,39] and European crow Corvus corone [30].

In the present study, we address the potential association between overall gene expression and degree of sex-biased gene expression. The evolution of sex-biased gene expression is expected as a result of accumulation of sexually antagonistic mutations, and the outcome depends on the degree of dominance of the alleles and which of the two sexes benefits [58–60]. Sex-biased expression of sex-linked genes is expected to evolve when the heterogametic sex suppresses its expression as a response to the accumulation of dominant homogametic-advantageous/heterogametic-harmful mutations on Z (or X). Recessive mutations beneficial to the heterogametic sex may also accumulate, but will be expressed in the homogametic sex only when they have reached higher frequencies and so selection for down-regulation should be weaker compared to the case of dominant homogametic-advantageous/heterogametic-harmful mutations. From this follows the expectation that sex-biased genes should have lower levels of average expression than unbiased ones [58]. Connallon and Knowles [59] tested this hypothesis by comparing microarray expression data for male-biased, female-biased, and unbiased genes in Drosophila. Surprisingly, they found that sex-biased genes were on average transcribed at higher rates than unbiased genes [59] [see also 61], and similar patterns were recently detected in zebra finch [26] and European crow [30]. We quantify expression on Z-linked and autosomal genes in the brain of zebra finches and common whitethroats using microarrays spotted with genome-wide zebra finch ESTs [29,62]. Our main finding is a pronounced positive association between the degree of male-bias and overall expression of Z-linked genes in both species. We discuss this in the light of hypotheses of the evolution of sex-biased gene expression [19,43,58,63].

Materials and Methods

We have analyzed microarray gene expression data from hybridizations of brain tissue from 12 adult zebra finches (6 males and 6 females) and 22 adult common whitethroats (11 males and 11 females). The correct ethical approvals for conducting the experiments were obtained from Malmö/Lund’s ethical committee (”Malmö/Lund djurförsöksst ska nämnd”); www.sjv.se/amnesomraden/djur/forsoks djur/etskprövning). Birds were in good condition and were sacrificed through decapitation to ameliorate suffering. For detailed information regarding sample handling, quality control results, hybridization and initial analysis of data, including normalization and filtering, see Naurin et al. [29]. Background correction in these expression data sets is the same as in Naurin et al. [29] in all aspects but one; in the present analyses we have also removed the 2% of ESTs with the lowest signals. This is in line with Affymetrix’s standard approach where the 2% lowest probes are removed (which for standard arrays is performed before any background correction according to hybridization on mis-match probes). This was done in order to avoid including ESTs with signals close to the background level in the analyses.

The Lund-zfa Array

The Lund-zfa array is an Affymetrix custom array that consists of 22360 zebra finch ESTs, representing approximately 15000 genes [62,64]. All ESTs have been BLASTed against the 3.2.4 build of the zebra finch genome [29,41] and hits in that BLAST was considered significant if the E-value was ≤10−20. In total, 1104 of the ESTs with intensities higher than the background cutoff in our data set have significant hits against the Z chromosome and 18520 ESTs have significant autosomal hits. Out of these significantly annotated ESTs, 22 were identified as Z-linked and female-biased in a previous study of sexual dimorphism in gene expression [29]. These ESTs had lower identities with the Z chromosome than male-biased ESTs (90% for female biased ESTs and 99% for male-biased ESTs) and three of them had significant hits against the W chromosome in chicken (Gallus gallus) [29]. In order to avoid analyzing potential W gene expression, we excluded these ESTs from the present analyses (including these ESTs did not qualitatively affect our results).

The Common Whitethroat Hybridization Efficiency

The somewhat lower degree of hybridization on microarray probes caused by sequence divergence between whitethroat RNA and the zebra finch-based microarray can be efficiently controlled via filtering [see Naurin et al. [29,62] for description of the comparative genome hybridization, CGH, of the common whitethroat and how data generated from those studies can be used for filtering].

Analyses of Dosage Compensation

Log2 gene expression intensity data for all ESTs with a significant BLAST hit against either an autosome or the Z chromosome were imported into SPSS Statistics 17.0. For each species, we conducted one-way ANOVAs using expression intensities to compare Z-linked versus autosomal ESTs for females and males, respectively, as well as the intensity of Z-linked ESTs in females versus Z-linked intensity in males. Fold Change (FC) was calculated as the mean unlogged male/mean unlogged female expression intensity. Mean FC for Z-linked ESTs versus autosomal ESTs was tested with a one-way ANOVA. Kolmogorov-Smirnov two-sample tests were used in order to test whether or not the distribution of FC was significantly different for Z-linked and autosomal ESTs. All results as listed below remained unchanged when non-parametric tests were used instead of ANOVAs.

General Linear Models (GLM) were used in order to test whether or not male-biased gene expression increased with increasing expression of Z-linked genes. Residuals for all GLMs were normally distributed.

Results

Analyses of Dosage Compensation

In the zebra finch, Z-linked gene expression was on average significantly lower in females (mean ± SE: 7.26±0.056) than in males (7.57±0.060; \( F_{1, 2208} = 14.2 \), \( p<0.0001 \) (Figure 1A).
Moreover, gene expression in females was significantly lower for Z-linked genes (7.26±0.056) than autosomal genes (7.66±0.015; F1, 19623 = 42.9, p<0.0001) (Figure 1A). Male Z-linked expression (7.57±0.060) was not significantly different from autosomal expression (7.65±0.014; F1, 19623 = 1.72, p = 0.192). Fold Change (FC; i.e. male/female intensity) was significantly higher for Z (1.31±0.01) than for autosomes (0.997±0.001; F1, 19623 = 6686, p<0.0001; Figure 2A), and the distribution of FC-values was significantly different for Z-linked and autosomal ESTs (p<0.0001; Kolmogorov-Smirnov two-sample test; Figure 3A).

For the common whitethroat, Z-linked gene expression tended to be lower in females (6.14±0.048) than in males (6.28±0.050; F1, 2908 = 3.73, p = 0.053) (Figure 1B). As in the zebra finch, female intensity in the common whitethroat was significantly lower for Z-linked (6.14±0.048) than autosomal genes (6.90±0.013; F1, 19623 = 181.4, p<0.0001) (Figure 1B). Male Z-linked intensity in the common whitethroat (6.28±0.050) was also significantly lower than autosomal expression (6.89±0.013; F1, 19623 = 120, p<0.0001; Figure 1B); a pattern that is different from that found in the zebra finch. Fold Change (FC) was significantly higher for Z (1.12±0.005) than for autosomes (0.998±0.001; F1, 19623 = 2809, p<0.0001; Figure 2B). The distribution of FC for Z-linked genes was significantly different from the autosomal distribution (p<0.0001; Kolmogorov-Smirnov two-sample test; Figure 3B).

**Association between Male-biased Gene Expression and Overall Expression Level**

We tested how the male-bias in gene expression on the Z chromosome was related to the general gene expression level. The degree of male-bias in expression of Z-linked genes increased with increasing mean gene expression levels and this pattern was significant in both species (Figures 1 and 4). In GLMs with male intensity as dependent variable, and female intensity, chromosome type (Z or autosomal) and the interaction term as independent variables, the interaction term was highly significant (zebra finch: p<0.0001, Table 1; common whitethroat: p<0.0001, Table 2). In other words, male Z-linked expression increased more steeply with
female gene expression than did male autosomal expression (Figure 4). All results from GLMs remained significant even if the 25% of the dataset with the lowest expression was removed, and the results are therefore not affected by detection rates of the microarray. Linear regressions between male and female expression levels showed that Z-linked expression had steeper regression slopes (higher β coefficient) than autosomal expression in both species (zebra finch: $\beta_{\text{Z-linked}} = 1.068 \pm 0.005$, $\beta_{\text{autosomal}} = 0.992 \pm 0.001$; common whitethroat: $\beta_{\text{Z-linked}} = 1.053 \pm 0.004$, $\beta_{\text{autosomal}} = 0.998 \pm <0.0001$) (Figure 4).

Z-linked in Relation to Autosomal Gene Expression
In order to evaluate how the pattern of gene expression on the Z chromosome relates to autosomal gene expression, we divided the
Z-linked and the autosomal dataset into four equal parts (quartiles) by sorting all ESTs by mean log2 expression (calculated for all individuals regardless of sex). Each quartile was then analyzed separately (Figure 1).

In the zebra finch, female Z-linked expression was significantly lower than male Z-linked and autosomal expression in all four categories (p < 0.0001; Table 3). Male Z-linked expression was slightly (but significantly) lower than mean autosomal expression in the quartile with the second lowest expression (p = 0.013), while slightly higher in the quartile with the second highest expression (p = 0.008; Table 3). Overall, male gene expression closely followed that of the autosomes, while female expression was close to autosomal expression only in the quartile with lowest gene expression and then became progressively lower with higher mean expression in comparison to both male Z-linked and autosomal genes (Figure 1A). The mean Z-linked over mean autosomal expression ratios (Z:A ratios) for the four quartiles of gene expression ranged between 0.98 and 0.94 (with the lowest ratio in the high quartile) in females and between 0.99 and 1.01 in males.

In the common whitethroat, Z-linked expression was significantly lower than autosomal expression in both sexes in all four quartiles (p < 0.0001; Table 4; Figure 1B). Female Z-linked expression was significantly lower than male Z-linked expression in the three groups with highest expression intensities (p = 0.0001 for the highest and second highest and p = 0.002 for the second lowest group) but not significantly different in the lowest quartile (p = 0.223; Table 4; Figure 1B). The Z:A ratios for the four quartiles of gene expression ranged between 0.93 and 0.89 (with the lowest ratio in the high quartile) in females and between 0.93 and 0.92 in males.

**Sequence Divergence in Common Whitethroat**

Mean number of significantly hybridizing probes (out of the 11 possible) for Z-linked genes (9.62 ± 0.040) in our cross-species comparative genome hybridization (CGH) study [62] was...
significantly lower than mean number of hybridizing probes on autosomes (10.23 ± 0.009; F_1, 19615, p < 0.0001). This result suggests a high rate of sequence evolution on Z. Thus, our common whitethroat expression data might be affected by the somewhat higher degree of sequence divergence on the Z chromosome than on the autosomes. We therefore re-analyzed the data set for the

Figure 4. Male gene expression intensity plotted against female expression intensity in (A) the zebra finch and (B) the common whitethroat.
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common whitethroat, this time using only ESTs that had significant hybridization on all of the 11 probes for each EST on the array in the previous CGH study [62]. This data set included 10370 autosomal and 443 Z-linked ESTs. For this restricted data set the results remained quantitatively unchanged. However, as expected if Z evolves faster than the autosomes, Z-linked expression was more affected by the removal of ESTs with some degree of sequence divergence (female Z-linked intensity was on average 8.4% higher and male Z-linked intensity 8.6% higher, while the autosomal expression intensity was only 5.0% higher in both sexes).

Discussion

Our present results on common whitethroat and zebra finch, together with previous results on chicken, zebra finch and European crow [23–25,29,30,39,65], clearly demonstrate that male-biased gene expression on the avian Z chromosome cannot be explained by a simple effect of the double dose of Z chromosome in males compared to females. Ratios of Z-linked to autosomal gene expression (Z:A) were relatively close to one in both sexes of both species (range: 0.89–1.01) with the lowest ratios for females in the subset (quartile) of highly expressed genes (0.94 in zebra finches and 0.89 in common whitethroat). Previous studies report similar levels for chicken and zebra finch males [23–25], but somewhat lower values for zebra finch and European crow females (c. 0.80) [26,30]. These results indicate that in birds the heterogametic females have up-regulated their single Z-linked homologues to a high extent when the W chromosome degraded and have thereby largely kept the balance between Z-linked and autosomal expression. This suggests that female birds achieve dosage compensation to a very high degree, although without chromosome-wide mechanisms [25,26,36,66,67]. A recent study which analysed level of expression on the current Z and compared it to that of a putative proto-Z came to the same conclusion of partial, but not complete, dosage compensation in female chicken [68]. However, even though females achieve a high degree of compensation there is still extensive and significant male-biased Z-linked gene expression in both our study species. When examining this in more detail, we found the interesting pattern that in both species the male-biased expression of Z-linked genes increased with increasing gene expression level from a more or less sex-balanced expression among Z-linked genes with low expression. This mainly resulted from a more pronounced drop in female Z-linked expression with higher mean expression than was the case for both male Z-linked and autosomal gene expression (see Figure 1). Together with the indications of increasing male-bias

| Source                  | Type III Sum of Squares | df | Mean Square | F     | Sig.  | Observed Power |
|-------------------------|-------------------------|----|-------------|-------|-------|----------------|
| Corrected Model         | 46413.6                   | 3  | 15471.2     | 62631.2 | <0.001 | 1.00           |
| Intercept               | 0.3                      | 1  | 0.3         | 11.2  | 0.001 | 0.92           |
| Female log2 intensity   | 14568.5                  | 1  | 14568.5     | 589815.6 | <0.001 | 1.00           |
| Chromosome type         | 1.8                      | 1  | 1.8         | 73.7  | <0.001 | 1.00           |
| Chrom. type * Female log2 int. | 19.9                 | 1  | 19.9         | 807.6 | <0.001 | 1.00           |
| Error                   | 284.4                    | 11515 | 0.0         |       |       |                |
| Total                   | 712863.3                 | 11519 |            |       |       |                |
| Corrected Total         | 46698.0                  | 11518 |            |       |       |                |

| Source                  | Type III Sum of Squares | df | Mean Square | F     | Sig.  | Observed Power |
|-------------------------|-------------------------|----|-------------|-------|-------|----------------|
| Corrected Model         | 39293.4                 | 3  | 13097.8     | 1310155.4 | <0.001 | 1.00           |
| Intercept               | 1.7                     | 1  | 1.7         | 170.4  | <0.001 | 1.00           |
| Female log2 intensity   | 10846.4                 | 1  | 10846.4     | 1084952.2 | <0.001 | 1.00           |
| Chromosome type         | 2.4                     | 1  | 2.4         | 240.7  | <0.001 | 1.00           |
| Chrom. type * Female log2 int. | 8.0                 | 1  | 8.0         | 797.8  | <0.001 | 1.00           |
| Error                   | 115.2                   | 11528 | 0.0         |       |       |                |
| Total                   | 563563.7                | 11532 |            |       |       |                |
| Corrected Total         | 39408.6                 | 11531 |            |       |       |                |

Table 1. Results from a General Linear Model with male expression intensity as dependent variable, and female expression intensity, chromosome type (Z or autosomal) and their interaction term as independent variables in the zebra finch.

Table 2. Results from a General Linear Model with male expression intensity as dependent variable, and female expression intensity, chromosome type (Z or autosomal) and their interaction term as independent variables in the common whitethroat.
with increased gene expression found by Melamed and Arnold [23] in the chicken, and the finding of higher expression of male-biased than unbiased genes recently described in zebra finches and European crow [26,30], our results from yet another passerine, the common whitethroat, as well as for the zebra finch, indicate that the pattern of lower female expression for highly expressed genes on Z is a general phenomenon among birds.

In addition to those patterns concerning sex-biased gene expression, we found that in the common whitethroat both males and females had lower expression of genes on the sex chromosomes than of those on the autosomes, and this was particularly true for highly expressed genes (Figure 1). This is interesting in relation to the result from our previous CGH study (comparing the number of significantly hybridizing probes on the microarray) that showed high sequence divergence on the Z chromosome between the common whitethroat and the zebra finch [62]; and such patterns are also found in other species [48,51,69]. The pronounced drop in Z-linked gene expression for highly expressed genes in both male and female common whitethroats observed here suggests that divergence is particularly pronounced for Z-linked genes with medium to high expression levels. Similar patterns based on data of sequence divergence (dN/dS analyses) between chicken and zebra finch were reported by Itoh et al. [26].

The evolution of Z-linked genes is affected by the lower effective population size compared to autosomal genes [70–72]. Moreover, Z is implicated in adaptive evolution, perhaps to a larger extent than X in e.g. mammals [71,73]. The potential role of Z-linked genes in the evolution of male-specific traits and sexual selection makes the high rate of molecular evolution on Z highly interesting [19,43,50,63]. In Ficedula flycatchers genes involved in speciation are predominantly Z-linked and coding for sex specific traits, like male plumage characteristics and female species recognition [74]. However, also in some male heterogametic systems such pattern have been found, for example, in Drosophila, the X chromosome harbours genes involved in female mate-choice underlying sexual isolation [75] (see also [34,54,61,76]).

Male-biased Expression of Highly Expressed Genes on Z

Even if it may seem unlikely that the somewhat lower Z-linked than autosomal expression in females should disrupt critical networks, there must be a compelling reason for why females do not balance their Z-linked expression to that of the autosomes for genes with high mean expression while achieving full balance for genes with low expression. We suggest three possible explanations for this:

First, if there was a chromosome-wide mechanism of dosage compensation also in birds a general up-regulation of Z-linked genes along the chromosome may have occurred in females to a degree equal to the level of low to medium expressed genes (but not to the level of highly expressed genes). However, this explanation seems unlikely as the level of compensation is a highly variable process in birds and varies between tissues and age groups in a way that indicates that selection for up-regulation has taken

### Table 3. Gene expression intensities in all four quartiles in the zebra finch.

|            | female Z | male Z | female autosomes | male autosomes |
|------------|----------|--------|-----------------|----------------|
| lowest 25% |          |        |                 |                |
| mean log2 expression | 5.07     | 5.18   | 5.19            | 5.2            |
| standard error | 0.02     | 0.022  | 0.008           | 0.008          |
| female Z vs. male Z: F = 14.3; P < 0.0001 |          |        |                 |                |
| female Z vs. autosomes: F = 24.6; P < 0.0001 |          |        |                 |                |
| male Z vs. autosomes: F = 1.06; P = 0.304 |          |        |                 |                |
| 2nd lowest 25% |          |        |                 |                |
| mean log2 expression | 6.28     | 6.54   | 6.62            | 6.62           |
| standard error | 0.029    | 0.033  | 0.009           | 0.009          |
| female Z vs. male Z: F = 33.3; P < 0.0001 |          |        |                 |                |
| female Z vs. autosomes: F = 127.9; P < 0.0001 |          |        |                 |                |
| male Z vs. autosomes: F = 6.24; P = 0.013 |          |        |                 |                |
| 2nd highest 25% |          |        |                 |                |
| mean log2 expression | 7.93     | 8.33   | 8.27            | 8.24           |
| standard error | 0.036    | 0.041  | 0.009           | 0.009          |
| female Z vs. male Z: F = 56.5; P < 0.0001 |          |        |                 |                |
| female Z vs. autosomes: F = 114.2; P < 0.0001 |          |        |                 |                |
| male Z vs. autosomes: F = 7.03; P = 0.008 |          |        |                 |                |
| highest 25% |          |        |                 |                |
| mean log2 expression | 9.76     | 10.3   | 10.4            | 10.3           |
| standard error | 0.054    | 0.054  | 0.019           | 0.019          |
| female Z vs. male Z: F = 40.2; P < 0.0001 |          |        |                 |                |
| female Z vs. autosomes: F = 94.4; P < 0.0001 |          |        |                 |                |
| male Z vs. autosomes: F = 2.41; P = 0.121 |          |        |                 |                |

Also shown are results for one-way ANOVAs of female and male Z-linked and autosomal expression. doi:10.1371/journal.pone.0046854.t003
place on a gene to gene basis rather than on a chromosome-wide level [25,26,36,66,67].

Second, selection for higher expression in females (to compensate for loss of expression as W-linked gene homologue degrades) might cause a correlated response in males, thereby increasing their gene expression as well. If increased expression is detrimental in males, then Z-linked genes might face further selection for down-regulation (causing an opposing spillover effect in females). Such opposing selection pressures could explain why female have not fully balanced all their Z-linked genes. This scenario has been suggested to explain the complex sex-biased gene expression with female overcompensation (X:A\textsuperscript{-1}) in Tribolium beetles [8]. However, it is not clear why this should preferentially affect highly expressed genes. Moreover, if males are more sensitive to increased expression in high-expressed than in low-expressed genes, it might be expected that selection in females will affect the male Z:A ratio for lowly expressed genes; a pattern not seen in our data.

Third, some property of many of the highly expressed genes might make them less suitable for high expression in females. It is likely that females have faced selection for up-regulation of Z homologues as W degraded [16–21]. However, the two chromosomes would have evolved separately for some time before selection for dosage compensation took place. Male-biased mutation rate [77–79] and the potential for high divergence in sex-biased Z-linked genes [43,70,80,81] give rise to obvious questions: Have some of the Z-linked genes evolved to increase male function due to high degree of sexual selection in males; and when females cannot rely on the degrading W any longer, would they even benefit from lower levels of compensation in order to avoid malfunction of these male-adapted (female-detrimental) genes?

Similar results to these now found in birds [23,26,30, this study] were previously found by Connallon and Knowles [59] in Drosophila; sex-biased genes were on average expressed at higher rates than unbiased genes [see also 61]. Interestingly, the pattern of male-bias is reported to be reversed in the silkworm (Bombyx mori) where the male-bias decreases with increasing gene expression level, creating a pattern where male-biased genes are expressed at lower levels than un-biased genes [40] (however, see [82] for a re-analysis of Zha et al.’s data). Zha et al.’s result in the silkworm – but not the results in birds – is in line with Rice’s [58] original hypothesis suggesting that genes become sex-biased when their expression in the harmed sex is decreased or abolished, and therefore, on average, sex-biased genes should have lower levels of expression than unbiased ones.

It has been suggested that sex-biased genes may be more dispensable [83] — a hypothesis which is intuitively easy to grasp — genes with low fitness effects should be more prone to respond to any kind of selection (sexual or otherwise) and sex-biased genes do show high degrees of sequence evolution in general. However, if this is true, one would expect a pattern like the one in silkworm (where genes with low expression show a high degree of sex-bias [40]) to be the most commonly reported scenario. The fact that the

| Table 4. Gene expression intensities in all four quartiles in the common whitethroat. |
|--------------------------------|----------------|----------------|----------------|----------------|
|                                | female Z | male Z | female autosomes | male autosomes |
| lowest 25%                     |          |       |                 |                |
| mean log\textsubscript{2} expression | 4.28    | 4.32  | 4.62            | 4.63           |
| standard error                 | 0.023    | 0.022 | 0.023           | 0.01           |
| female Z vs. male Z: F = 1.48; P = 0.223 |          |       |                 |                |
| female Z vs. autosomes: F = 103.2; P < 0.0001 |          |       |                 |                |
| male Z vs. autosomes: F = 80.3; P < 0.0001 |          |       |                 |                |
| 2nd lowest 25%                 |          |       |                 |                |
| mean log\textsubscript{2} expression | 5.47    | 5.56  | 6.01            | 6.01           |
| standard error                 | 0.019    | 0.02  | 0.006           | 0.006          |
| female Z vs. male Z: F = 9.91; P = 0.002 |          |       |                 |                |
| female Z vs. autosomes: F = 697.5; P < 0.0001 |          |       |                 |                |
| male Z vs. autosomes: F = 495.7; P < 0.0001 |          |       |                 |                |
| 2nd highest 25%                |          |       |                 |                |
| mean log\textsubscript{2} expression | 6.52    | 6.65  | 7.19            | 7.19           |
| standard error                 | 0.043    | 0.044 | 0.013           | 0.013          |
| female Z vs. male Z: F = 16.6; P < 0.0001 |          |       |                 |                |
| female Z vs. autosomes: F = 755.5; P < 0.0001 |          |       |                 |                |
| male Z vs. autosomes: F = 482.6; P < 0.0001 |          |       |                 |                |
| highest 25%                    |          |       |                 |                |
| mean log\textsubscript{2} expression | 8.32    | 8.6   | 9.32            | 9.32           |
| standard error                 | 0.063    | 0.064 | 0.023           | 0.023          |
| female Z vs. male Z: F = 9.88; P < 0.0001 |          |       |                 |                |
| female Z vs. autosomes: F = 179.3; P < 0.0001 |          |       |                 |                |
| male Z vs. autosomes: F = 90.3; P < 0.0001 |          |       |                 |                |

Also shown are results for one-way ANOVAs of female and male Z-linked and autosomal expression.
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opposite seems to be true is highly interesting given the fact that highly expressed genes have a low degree of sequence evolution and are often assumed to be more essential [44,61,70,76,84–91]. Hence, the patterns of sex-bias in relation to gene expression levels reported to date do not support the idea that simple dispensability is what makes a gene prone to accumulate sex-bias or respond to sexual selection; other mechanisms must also be invoked if we are to understand which parts of the genome are more likely to obtain sex-specific functions.

To explain that highly expressed male-biased genes in Drosophila are less frequently located on the X chromosome than are lowly expressed male-biased genes [e.g. 76], Vicoso and Charlesworth [92] proposed the following hypothesis: The increase in expression required to make a gene male-biased is less likely when X-linked because there is an upper limit to the rate of expression that can be achieved [93,94] and because the X chromosome is often hyper-activated in Drosophila males as a result of the evolution of dosage compensation [5–8,31–33] [but see 34]. This type of hypothesis is however unlikely to explain the observation of increasing male-biased gene expression with increasing expression in female heterogametic birds, because there seem to be no chromosome-wide dosage compensation [24–26,36,66,67]. However, the idea that there may be an upper level to the rate of expression that can be achieved with a single gene copy, in this case on the Z in the hemizygous sex, due to space limitations on the DNA strand for binding transcriptional molecules, could explain why females may have difficulties in fully balancing the most highly expressed Z-linked genes. However, we find this suggestion somewhat less likely because the sex-biased Z-linked genes are not among the highest expressed in our data set, thus do not seem saturated (Figure 4), and because it remains unclear why such an expression ceiling would apply only to some species (e.g. this study) and not others [5,6,10,12,13,15,40].

The result that male-biased and unbiased genes in chicken are not only associated with certain expression levels but also have functional differences [23], and that female detrimental antagonism is overrepresented on the Z chromosome [95], would imply that selection on Z-linked genes in males have adapted them to male function in a way that makes them unsuitable for up-regulation in females [43,53]. However, observations of such functional differences and accumulation of female detrimental antagonisms do not provide an explicit hypothesis to explain the positive association between male-biased Z-linked expression and overall gene expression now found in birds. Moreover, the data available for autosomal genes from several taxa suggest that sex-biased genes evolve fast, whereas highly expressed genes – as mentioned above – in general have a low degree of sequence, protein and expression profile evolution [44,61,70,76,84–91]. Hence, it is surprising to find a proportionally large amount of male-bias in highly expressed Z-linked genes in both the chicken, crow, zebra finch and the common whitethroat, a finding that suggests that there has been a high degree of sex-specific selection at Z-linked loci in birds and therefore that highly expressed Z-linked genes are hotspots for evolution of sexual dimorphism in birds [19,34,43,58,63,73,74,91,96]. Further studies of gene expression and sexual antagonism in both female and male heterogametic systems are clearly needed to elucidate patterns and processes of sex-biased gene expression on the sex chromosomes [cf. e.g. 19,44].

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Author Contributions

Conceived and designed the experiments: SN DH SB BH. Performed the experiments: SN. Analyzed the data: SN. Wrote the paper: SN BH.
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