Reference Genes for Quantitative Gene Expression Studies in Multiple Avian Species

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Abstract

Quantitative real-time PCR (qPCR) rapidly and reliably quantifies gene expression levels across different experimental conditions. Selection of suitable reference genes is essential for meaningful normalization and thus correct interpretation of data. In recent years, an increasing number of avian species other than the chicken has been investigated molecularly, highlighting the need for an experimentally validated pan-avian primer set for reference genes. Here we report testing a set for 14 candidate reference genes (18S, ABL, GAPDH, GUSB, HMBS, HPRT, PGK1, RPL13, RPL19, RPS7, SDHA, TFR2, VIM, YWHAZ) on different tissues of the mallard (Anas platyrhynchos), domestic chicken (Gallus gallus domesticus), common crane (Grus grus), white-tailed eagle (Haliaeetus albicilla), domestic turkey (Meleagris gallopavo f. domestica), cockatiel (Nymphicus hollandicus), Humboldt penguin (Spheniscus humboldti), ostrich (Struthio camelus) and zebra finch (Taeniopygia guttata), spanning a broad range of the phylogenetic tree of birds. Primer pairs for six to 11 genes were successfully established for each of the nine species. As a proof of principle, we analyzed expression levels of 10 candidate reference genes as well as FOXP2 and the immediate early genes, EGR1 and CFOS, known to be rapidly induced by singing in the avian basal ganglia. We extracted RNA from microbiopsies of the striatal song nucleus Area X of adult male zebra finches after they had sang or remained silent. Using three different statistical algorithms, we identified five genes (18S, PGK1, RPS7, TFR2, YWHAZ) that were stably expressed within each group and also between the singing and silent conditions, establishing them as suitable reference genes. In conclusion, the newly developed pan-avian primer set allows accurate normalization and quantification of gene expression levels in multiple avian species.

Introduction

The zebra finch (Taeniopygia guttata) and the domestic chicken (Gallus gallus f. dom.) have become widely used model organisms for biologists studying neurobiology and behavior, ecology as well as diseases and their transmission [1–5]. With both genomes sequenced and technologies adapted from traditional genetic model systems, the finch and the chicken are becoming increasingly amenable to genetic investigation [6–9]. The more than 10,000 avian species are known for their extraordinary differences in behavior and physiology. This offers opportunities to find a suitable species to address particular questions about the genetic background of migration, mating systems, parental care, flight, niche partition, vocal behavior or cognition, to name just a few.

Quantitative real-time polymerase chain reaction (qPCR) is currently the most economic, efficient and reliable method to measure gene expression levels in low to medium throughput approaches. It is sensitive, specific and reproducible even with limited mRNA copy numbers [10]. As in any quantitative study, it is necessary to correct for sample to sample variations in order to obtain reliable results. The most commonly used method is normalization to the expression of an internal control gene, also called reference gene. Hypothetically, the ideal reference gene is expressed at stable levels irrespective of tissue type, species, treatment, metabolism or sampling conditions. To date, no such ideal gene has been found and most likely does not exist, thus it is becoming clear that reference genes need to be established for each new experimental design, controlling for the numerous variables which might bias the results, i.e., number of cells and their transcriptional activity, RNA quality and reverse transcriptase efficiency [11]. In recent avian gene expression experiments, genes such as 18S (18S ribosomal RNA), ACTB (beta actin) and GAPDH (glyceraldehyde 3-phosphate dehydrogenase) have been used most frequently [12–17] but several studies have shown that using single or inappropriate reference genes for normalization may dramatically bias the results of mRNA copy number quantification [10,18–20].

For mammals, multiple sets of reference genes have been published [21–25] but are of limited use for non-mammalian vertebrates including birds due to the phylogenetic distance. Certainly, the advent of Next Generation Sequencing and the now fully sequenced genomes of the domestic chicken, domestic turkey (Meleagris gallopavo f. domestica), zebra finch, duck (Anas platyrhynchos) and the collared flycatcher (Ficedula albicollis) has made it easier and faster to conduct gene expression studies in these species [7,9,26–28]. However, currently more than 10,000 avian species are known, more and more of which are also investigated at the
molecular level [29–32]. The tools available to compare those less genetically amenable bird species are highly limited, one problem being the lack of sequence information to design PCR-primers. To date, avian reference genes have been published only for the domestic chicken [33,34], great tit (Parus major) [32], Japanese quail (Coturnix c. japonica) [35] and domestic pigeon (Columba livia f. domestica) [36].

This prompted us to develop a set of pan-avian PCR primers for the amplification of reference genes that can be tested for their suitability to normalize gene expression levels in avian studies involving qPCR analyses. We designed primer pairs for the amplification of 14 candidate reference genes classically used in mammalian research, including 18S, ABL (Abelson murine leukemia viral oncogene homolog), GAPDH, GUSB (beta glucuronidase), HMBS (hydroxymethylbilane synthase), HPRT (hypoxanthine-guanine phosphoribosyltransferase 1), PGK1 (phosphoglycerate kinase 1), RPL13 (60S ribosomal protein L13), RPL19 (60S ribosomal protein L19), RPS7 (40S ribosomal protein S7), SDHA (succinate dehydrogenase complex, subunit A), TFRC (transferrin receptor protein 1), VIM (vimentin) and YWHAZ (tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, zeta polypeptide). For each of the nine avian species investigated in this study we successfully established between six and 11 primer pairs.

To demonstrate the suitability of these genes for normalization purposes, we conducted an experiment investigating singing-induced gene expression changes in a striatal song control region, Area X of zebra finches. We measured the expression of three genes of interest, among them two immediate early genes known to be upregulated by singing [37–39], as well as 10 potential reference genes, on samples of birds that sang undirected song (not for a female conspecific) or remained silent for the same amount of time. Using the statistical algorithms geNorm [11], NormFinder [40] and BestKeeper [41] we determined the most stably expressed genes and employed them to normalize the expression of our genes of interest.

### Material and Methods

Animal experiments were carried out in accordance with the guidelines provided and approved by the governmental institutions (LAGeSo, Berlin, Permit Number: T 0298/01). All zebra finch samples originated from a captive breeding colony approved by local authorities (permits: ZH147). The brain of the

### Table 1. Gene names and primer pairs tested on all nine avian species.

| Gene symbol | GenBank accession number G. gallus | Primer sequences 5’-3’ (forward/reverse) | Amplicon length (bp) | Size on genomic level chicken (bp) |
|-------------|----------------------------------|------------------------------------------|----------------------|-----------------------------------|
| 18S | AF173612 | CGAAGGATTTGGCAGAGAAT/98–99 GGTCACCTGGGACG | 98–99 | 98 |
| ABL | XM_001233811 | GCTGTCGCGGAGAACC/218 TGATGATGTAATTGCTGGGGACC | 218 | 940 |
| GAPDH | NM_204305 | GTTTGCTGCTTGGTTGATCACT/269–270 GGCAGGCACCTGCAACTC | 269–270 | 534 |
| GUSB | NM_001039316 | GCGAAACCTCTCCGGAC/222–224 TCAAGGTTTGGGACG | 222–224 | 858 |
| HMBS | XM_417846 | CTTCTTTGTCGCTGGACG | 181 | 1575 |
| HPRT | NM_204848 | GATGAAACCTCCAGAACTGGAGA | 181 | 1575 |
| PGK1 | NM_2049958 | AAAATTTCGATCAATCAGCTG/167 GCACTCAGGCTGCAATC | 167 | 450 |
| RPL13 | NM_204999 | CCAACGGAGACTGCTGAGG/135 TATAGCCACCCCTGCCAGAC | 135 | 434 |
| RPL19 | NM_001030929 | CCAACGGACCCACGCAGACATC/152–153 TATAGCCACCCCTGCCAGAC | 152–153 | 629 |
| RPS7 | XM_419936 | TATGAGTGGGACGGAACG/156 TGGTCGCTGCCGACGAT | 156 | 1773 |
| SDHA | XM_419054 | CAGAAACAGCTTTGCCCAATG/238 TATAGCCACCCCTGCCAGAC | 238 | 1821 |
| TFRC | NM_205256 | GAAATTCCGGCTGATCTG/111–113 CGACTCAGCAGCTGCCG | 111–113 | 723 |
| VIM | NM_001048076 | GGAAGATGATGACGCTGACG/145 GCAAAATCTCCTGACATGC | 145 | 761 |
| YWHAZ | NM_001031343 | GTGAGAGCTTGGTGATGCTG/222–224 GCGGACCTTGGTGATGCTG | 222–224 | 326 |
Summary of primer test and efficiency for each primer pair and species.

| Species                  | No. of established primer pairs | Species                  | No. of established primer pairs |
|--------------------------|---------------------------------|--------------------------|---------------------------------|
| Zebra finch             | nd                              | White tailed eagle       | nd                              |
| Cockatiel                | nd                              | Humboldt penguin         | nd                              |
| White-tailed eagle       | nd                              | Common crane             | nd                              |
| Hummingbird              | nd                              | Chicken                  | nd                              |
| Common crane             | nd                              | Turkey                   | nd                              |
| Turkey                   | nd                              | Mallard                  | nd                              |
| Ostrich                  | nd                              | Jellyfish                | nd                              |

Numbers refer to the efficiency determined (bold: efficiency ok); nt, primer not tested; nw, primer tested but not working; nd: efficiency not tested.

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Sample collection

Three sets of tissues were collected. (1) We collected samples from nine phylogenetically distant avian species: The brains from a mallard (Anas platyrhynchos), a domestic chicken (Gallus gallus domesticus), a white-tailed eagle (Haliaeetus albicilla), a domestic turkey (Meleagris gallopavo f. domestica), a cockatiel (Nymphicus hollandicus) and an ostrich (Struthio camelus); from a common crane (Grus grus) we took blood and from a Humboldt penguin (Spheniscus humboldti) the lung with fungal pneumonia. (2) From adult zebra finches we collected brains and gonads (10 females, 10 males, below called 'tissue dataset'). (3) We collected microbiopsies of Area X from another 12 adult zebra finches (below called 'song data set'). All samples were immediately snap frozen after dissection and stored at ~80°C until further use.

Song data set. Adult male zebra finches were housed in free flight aviaries in our breeding colony and kept alone in sound attenuated chambers overnight. After lights were switched on in the morning at 8 a.m., song was monitored and recorded. Birds that did not sing during the first 30 min were sacrificed ('silent group'). Birds that started singing (while being alone, called ‘undirected song’) within the first two hours were sacrificed 30 min after the onset of song (‘song group’). Birds were sacrificed by an isoflurane overdose and brains were immediately dissected, cut into hemispheres, embedded in Tissue-Tek O.C.T. compound (Sakura Finetek) and stored at ~80°C. To obtain microbiopsies of Area X, hemispheres were mounted on a cryostat (Cryo-Star HM 560 Cryostat, MICROM) and 20 µm sagittal sections were cut from medial to lateral until Area X became visible. Then a 1 mm diameter coring tool (Harris Unicore) was used to mark Area X from another 12 adult zebra finches (below called 'song data set'). All samples were immediately snap frozen after dissection and stored at ~80°C until further use.

Selection of candidate reference genes and primer design

Candidate reference genes were selected from previous reports for mammalian species [21–25] (Table 1). All primers except RPL13 [36] were designed de novo on homologous gene segments of the chicken and zebra finch derived from GenBank (http://www.ncbi.nlm.nih.gov) and Ensembl (http://www.ensembl.org/index.html) databases using NetPrimer (http://www.premiersoft.com/netprimer/netprimer.html). Primers were designed to reside in the open reading frame (ORF) of genes to grant maximal conservation across all birds. ORFs were identified by Biowire Jellyfish 1.5. Criteria for primer design were a predicted melting temperature of 56°C, primer length of 15-25 nucleotides, a guanine-cytosine content of 40–70% and amplicon lengths smaller than 250 base pairs [10,42]. All primer pairs except those for 18S were designed to span exon-exon boundaries as identified by MEGA4 [43].
Table 3. Sequence similarity of all amplified products to the chicken (ICGSC Gallus gallus 4.0) and zebra finch (WUGSC 3.2.4/taeGut1) genomes.

| Gene symbol | Sequence identity (%) to G. gallus to T. guttata |
|-------------|-----------------------------------------------|
|              | A. platyrhynchos | G. gallus | G. grus | H. albicilla | M. gallopavo | N. hollandicus | S. humboldti | S. camelus | T. guttata |
| 18S          | 100/100          | 100/100   | 100/100 | 98/98        | 100/100      | 98/98         | 97/97       | 100/100    | 100/100    |
| ABL          | nw               | 100/89    | nw      | 95/92        | 98/88        | 97/93         | 97/93       | 94/89      | 89/100     |
| GAPDH        | 95/94            | 100/92    | 93/95   | 94/95        | 98/93        | 93/96         | 93/94       | 94/92      | 92/100     |
| GUSB         | 91/89            | 97/86     | 88/89   | 91/89        | 96/87        | 90/89         | 90/89       | 87/90      | 87/100     |
| HMBS         | 93/89            | 100/89    | 88/93   | 92/96        | 97/98        | 92/90         | 88/92       | 88/92      | 89/100     |
| HPRT         | 95/98            | 100/94    | 96/98   | 95/99        | 97/95        | 95/98         | 95/99       | 95/99      | 94/100     |
| PGK1         | 92/95            | 100/92    | 93/96   | 93/97        | 98/92        | 93/97         | 93/95       | 93/96      | 92/100     |
| RPL13        | nd               | 100/85    | nd      | nd           | nd           | nd            | nd          | 86/-       | 90/95      |
| RPL19        | nd               | 100/86    | nd      | nd           | nd           | nd            | nd          | 89/84      | 85/97      |
| RPS7         | 98/93            | 100/93    | 96/93   | 97/95        | 100/93       | 97/93         | 97/92       | 97/92      | 93/100     |
| SDHA         | 93/89            | 100/90    | 94/93   | 95/93        | 98/91        | 92/91         | 93/91       | 90/90      | 90/100     |
| TFRC         | 99/87            | 100/86    | 86/95   | 87/90        | 99/87        | 89/86         | 90/93       | 94/89      | 86/100     |
| VIM          | 97/90            | 100/88    | 93/90   | 96/91        | 98/90        | 95/92         | 96/90       | 93/90      | 88/100     |
| YWHAZ        | 97/95            | 100/96    | 96/95   | 96/97        | 99/96        | 97/97         | 97/97       | 97/96      | 96/99      |

*nd, not done; nw, not working.

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RNA extraction, cDNA synthesis and qPCR

All tissues except for Area X tissue samples (‘song data set’). RNA was extracted and purified from approximately 100 mg of each sample using Trizol (Invitrogen, Carlsbad, CA) or the NucleoSpin RNA II kit (Macherey-Nagel, Düren, Germany) following the manufacturer’s recommendations. Concentration and purity of total RNA were determined at 260/280 nm absorbance ratio to be above 1.8 in all cases (NanoDrop 1000 Spectrophotometer, Thermo Fisher Scientific, Wilmington, DE). RNA integrity numbers (RIN) of all tissue samples were measured using the RNA Nano kit on an Agilent 2100 Bioanalyzer (RNA 6000 Nano Kit, Agilent, Santa Clara, CA) and used only if above 8.0. For cDNA synthesis, 100 ng RNA was reverse transcribed at 25°C for 5 min followed by 30 min at 42°C containing 200 U iScript (Bio-Rad, Hercules, CA), 100 mmol MgCl2, 50 ng random hexamers, 0.2 mmol DTT, 40 U RNase Out (Promega, Madison, USA) and 10 nmol of each dNTP. The initial tests for successful product amplification were run on the MX 3000P or the CFX96 qPCR detection system using the MX Pro 3.0 (Agilent) or CFX Manager 2.0 (Bio-Rad) software packages, respectively. All subsequent PCR experiments were performed on the CFX96 system. PCR reactions were carried out in 96-well polystyrene plates (Sarstedt, Numbrecht, Germany) in a volume of 20 µL. 5 µL cDNA was added to 15 µL reaction mix containing 10 µL Brilliant SYBR Green QPCR Master Mix (Applied Biosystems, Carlsbad, CA) and 4.5 pmol of each primer. Cycling conditions were as follows: 10 min at 95°C followed by 40 cycles of 30 sec at 95°C, 1 min at 58°C, and 30 sec at 72°C and a subsequent melting curve analysis. All reactions were run in triplicate and the mean Ct-values were used for further analysis. If one of the three Ct-values deviated from the other two by more than 0.5 it was excluded from further analysis or the experiment was repeated. Nuclease-free water was used for the no-template controls. PCR efficiencies (E) were calculated using the equation $E = 10^{-1/\text{slope}}$ by measuring a ten-fold dilution series over five orders of magnitude for each primer pair and for each avian species as indicated in Table 2. Specificity of the primers was confirmed by sequencing and sequence identity was evaluated with BLASTN [44] against coding sequences of the chicken and zebra finch derived from GenBank (Table 3).

Song data set. After microscopic verification of proper targeting, we extracted the RNA from the Area X biopsies of each animal individually, resulting in 12 separate RNA extractions. RNA was extracted using the NucleoSpin RNA XS kit (Macherey-Nagel) and quantified using the Qubit RNA assay (Life Technologies). For cDNA synthesis 70 ng RNA was reverse transcribed in 20 µL using 200 ng random hexamer primers, 10 nmol of each dNTP, 50 mM DTT, 200 U SuperScript III (Invitrogen) and 40 U RNasinPlus (Promega). The temperature program was chosen as follows: 5 min at 65°C, cool down on ice, 5 min 25°C, 45 min at 50°C, 15 min at 72°C. We included reverse transcriptase-free reactions to control for DNA contaminations. All cDNAs were diluted 10-fold with nuclease free water and 5 µL were used in each PCR reaction.

QPCR reactions were carried out in 96-well polystyrene plates (Sarstedt, Numbrecht, Germany) in a volume of 20 µL and in triplicates. 5 µL cDNA was added to 15 µL reaction mix containing 10 µL KAPA SYBR FAST Universal QPCR mix (Peqlab) and 10 pmol (18 pmol in the case of FoxP2) of each primer. PCRs were run on a MX3005P system (Agilent) with the following cycling conditions: 10 min at 95°C followed by 40 cycles of 30 sec at 95°C, 30 sec at 65°C (64°C for FoxP2) and a
Figure 2. Specificity of 10 reference genes of the zebra finch with single peaks in melting curves. All amplifications were run in triplicate on 32 tissue samples. doi:10.1371/journal.pone.0099678.g002
subsequent melting curve analysis. Primer pairs for FOXP2 (CCTGGCTGTGAAAGCGTTTG/ATTTGCACCCGACACTGAGC [8]), EGR1 (ACTTCATCATCGCCATCCTC/TGGAATTGGGAAATGTTGGT) and CFOS (AGCTGGAGGAGGAAGTCC/CTCCTCGGAGAAGCACAACT) were designed for the zebra finch only.

Analysis of expression stability

The stability of expression was evaluated for 10 zebra finch candidate reference genes (18S, GUSB, HMBS, HPRT, PGK1, RPS7, SDHA, TFRC, VIM and YWHAZ) on samples from 10 female and 10 male brains and six gonads of each sex, and samples from Area X of male zebra finches that sang before sacrifice (n = 5) or were silent (n = 7). Mean Ct values from qPCR runs were exported to Excel (Microsoft Excel 2010) and analyzed by the three commonly used statistical algorithms geNorm (version 3.5), NormFinder (version 0.953) and BestKeeper (version 1.0) [11,40,41]. For the geNorm analysis, we employed the comparative Ct method, taking into account the efficiency of each assay as outlined in the manual. The resulting values were used as input into the geNorm analysis. For the NormFinder analysis we followed the input procedure outlined in [45]. For the BestKeeper analysis the raw Ct values were entered into the Excel mask.

The song data set was analyzed for differences in gene expression using the Mann-Whitney-U-test.

Results

Normalization to reference genes is a crucial step to obtain meaningful and reliable results, when gene expression levels are quantified using qPCR. In this study we aimed to establish and test a set of primer pairs that could ideally be used on the entire avian class. We selected 14 candidate genes (18S, ABL, GAPDH, GUSB, HMBS, HPRT, PGK1, RPS7, SDHA, TFRC, VIM and YWHAZ) and designed non-degenerate primer pairs.

Establishing and testing primer pairs on the avian class

To test our primers, we collected samples from different branches of the avian tree in an attempt to represent phylogenetic diversity. We chose the zebra finch, cockatiel and white tailed eagle to represent land birds, the Humboldt penguin for water birds, the common crane for the gruiformes, the chicken, turkey and mallard for galliformes and the ostrich for paleognaths. We tested all primer pairs except RPL13 and RPL19 on all species. The performance of each primer pair was evaluated for each species separately and was considered successful if the following criteria were met:

1. Amplification of a single product indicated by a single peak in the melting curve analysis
2. Sequence of the PCR product confirming amplification from the proper gene
3. Efficiency of amplification between 90 and 110%.

Concerning the first criterion, a single product was amplified by most primer pairs in most species (Figure 1, 2; Table 2), except for
Expression stability analysis of candidate reference genes
for normalization in the zebra finch

To test if the established primers could be used for normalization in a proof of principle experiment, we conducted two independent screens with 10 candidate genes (18S, GUSB, HMBS, HPRT, PGK1, RPS7, SDHA, TFRC, VIM, and YWHAZ) on zebra finch tissues (Figure 3).

1) Microbiopsies of the striatal song nucleus Area X, either taken 30 min after the onset of undirected singing or of a non-singing control (henceforth called “song data set”).

2) Zebra finch entire brains (female and male separately), testes and ovaries.

We determined the expression stability of the 10 genes in both screens using three different algorithms: geNorm [11], NormFinder [40] and BestKeeper [41]. Table 4 and Table 5 display the rankings derived from each of the algorithms for the two data sets.

All raw data were prepared for input into the three software applications as described in the methods section. We excluded GUSB from the analysis in the song data set, as it was expressed below the detection limit of our assay.

GeNorm. The geNorm program is a Visual Basic application (VBA) tool for Microsoft Excel [11]. The algorithm assumes that the ratio of two reference genes is unaffected by any experimental treatment or condition. It calculates the pairwise variation of each gene with all other control genes. The stability value M for a gene is the average of all pairwise variations of this particular gene. By stepwise exclusion of the gene with the highest M values, the two most stable reference genes are determined. One critical assumption of the algorithm is that the tested genes are not co-regulated [46]. If they are, they will falsely be considered as suitable reference genes, because of their low co-variation.

In the case of the song dataset, geNorm analysis recommended 18S and PGK1 as the most stably expressed genes (Table 4). In contrast, the 18S gene was one of the least stably expressed genes in all four tissues tested (Figure 3, Table 5).

Besides determining which genes are most suited as reference genes, the geNorm algorithm also gives a recommendation on how many genes should be used for normalization. A Vn/n+1 pairwise variation value below 0.15 indicates that n genes are sufficient for normalization and the gene n+1 should not be included. For the song dataset the V2/3 value (0.079) was already below 0.15 indicating that two genes are sufficient for normalization. For the other tissues the same was true if the tissues were analyzed separately. When all four tissues were analyzed together, three genes were recommended.

NormFinder. The NormFinder program, also a VBA based Microsoft Excel tool, is regarded as the most reliable of the three algorithms we employed [40]. In contrast to the other two approaches, NormFinder takes the experimental groups from which samples are drawn into account [46]. The algorithm distinguishes between variation across samples irrespective of the experimental condition, and variation across experimental groups due to experimental condition. To determine the most suited reference genes, NormFinder analyzes the inter- and intra-group variability and calculates a so called “stability value” which takes both measures into account. Consequently, the genes which are most suited as reference genes have the lowest stability values.

For the song dataset YWHAZ and 18S were recommended as reference genes, though 18S alone did not have the second lowest stability value, but the two genes together had a very low stability.
Table 5. Results of testing the stability of expression analysis for the four different zebra finch tissues and three different statistical methods.

| Gene     | All (Brain, Gonads) | Brain male | Brain female | Testes | Ovaries |
|----------|----------------------|------------|--------------|--------|---------|
|          | gN  | Nf | BK | gN  | Nf | BK | gN  | Nf | BK | gN  | Nf | BK | gN  | Nf | BK |
| 18S      | 7  | 8 | 9 | 7  | 8 | 9 | 7  | 8 | 9 | 7  | 8 | 9 | 7  | 8 | 9 |
|          | 10 | 9 | 10 | 10 | 9 | 10 | 10 | 9 | 10 | 10 | 9 | 10 | 10 | 9 | 10 |
| GUSB     | 678 | 567 | 677 | 666 | 557 | 43 | 66 | 3 | 5 | 29 | 43 | 3 | 56 | 2 | 4 |
|          | 11 | 5 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 | 12 |
| HPRT     | 232 | 113 | 334 | 111 | 224 | 3 | 5 | 21 | 12 | 5 | 21 | 12 | 5 | 21 |
|          | 5 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 | 12 | 4 |
| PGK1     | 113 | 443 | 44 | 68 | 8 | 68 | 8 | 68 | 8 | 68 | 8 | 68 | 8 | 68 | 8 |
|          | 2 | 11 | 3 | 2 | 11 | 3 | 2 | 11 | 3 | 2 | 11 | 3 | 2 | 11 |
| RPS7a    | 32 | 67 | 7 | 2 | 55 | 24 | 3 | 58 | 8 | 2 | 55 | 24 | 3 | 58 | 8 |
|          | 67 | 7 | 67 | 7 | 67 | 7 | 67 | 7 | 67 | 7 | 67 | 7 | 67 | 7 | 67 |
| SDHA     | 5 | 12 | 3 | 5 | 12 | 3 | 5 | 12 | 3 | 5 | 12 | 3 | 5 | 12 | 3 |
|          | 3 | 5 | 12 | 3 | 5 | 12 | 3 | 5 | 12 | 3 | 5 | 12 | 3 | 5 | 12 |
| TFRC     | 865 | 887 | 865 | 887 | 865 | 887 | 865 | 887 | 865 | 887 | 865 | 887 | 865 | 887 | 865 |
|          | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 | 21 |
| VIM      | 331 | 43 | 441 | 89 | 23 | 5 | 5 | 23 | 5 | 5 | 23 | 5 | 5 | 23 |
|          | 3 | 5 | 23 | 5 | 23 | 5 | 23 | 5 | 23 | 5 | 23 | 5 | 23 | 5 | 23 |
| YWHAZ    | 2 | 65 | 76 | 22 | 2 | 65 | 76 | 22 | 2 | 65 | 76 | 22 | 2 | 65 | 76 |
|          | 55 | 75 | 65 | 75 | 65 | 75 | 65 | 75 | 65 | 75 | 65 | 75 | 65 | 75 | 65 |

Samples were either analyzed together (all), by tissue type (brains, gonads) or separately. The first four ranked genes are shown in bold. gN, geNorm; Nf, Normfinder; BK, BestKeeper.

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value. For the different tissues the results again differed (Table 5). One interesting aspect here was that 1WH1A was the second most stable gene in the male brain but only ranked as the seventh most stable gene in the female brain, hinting towards a sexually dimorphic expression in the brain.

**BestKeeper.** The BestKeeper Excel template offers several measures to detect the most stable genes [41]:

1. The standard deviation (SD) of the Cts of all samples for one gene.
2. The correlation coefficient with the BestKeeper Index. The BestKeeper Index for each sample is the geometric mean of the Cts of all reference genes.
3. The coefficient of variation of a potential reference gene.

The BestKeeper does not indicate how to weigh the importance of these three measures, leaving it up to the experimenter to choose which values to consider most relevant. As a guideline, the authors recommend to exclude all genes with an SD > 1.5. However, in the song dataset all candidate genes displayed SDs higher than 1.5, consistent with a previous report finding that the BestKeeper algorithm may be better suited for analysis of gene expression in homogeneous cell populations rather than complex tissues [46]. Given that previous studies used various selection criteria, for example SD only [46], R² only [47], an unspecified combination of SD+CV [48] and in one case a self-designed decision criterion [49], we decided to rank all potential reference genes based on each of the values separately and then calculate the mean of the rankings to determine the final rank of a gene. Employing this method, RPS7, TFRC, VIM and 1WH1A were the four most stably expressed genes. We decided to use RPS7 and TFRC for normalization as TRFC, unlike VIM and 1WH1A, was equally stable in all three values contributing to the final ranking (Table 4). For the other tissues the BestKeeper ranking largely agreed with the ranking of the other two programs (Table 5).

**Reference gene validation**

To evaluate the usefulness of the reference genes suggested by our data analysis, we measured the expression of EGR1, CFOS and FOXP2 genes in the song dataset. It is known that EGR1 and CFOS mRNA is upregulated in a sexually dimorphic subregion of the avian striatum, Area X, after the bird sings undirected song for at least 30 minutes [38,39]. FOXP2 is downregulated by undirected singing that lasts for two hours. Whether FOXP2 is also regulated as early as 30 minutes after the onset of song is currently unknown [50].

We normalized the expression of each of our genes of interest to the geometric mean of the two best genes recommended by each algorithm. Regardless of which genes we chose for normalization, we detected a significant upregulation of CFOS and EGR1 by more than 7-fold in the singing animals compared to the silent controls (Figure 4). We failed to detect such a significant result for FOXP2. The upregulation of CFOS was most significant if normalized to 18S and 1WH1A as suggested by NormFinder. The results for EGR1 and FOXP2 did not differ much between the different normalizations. These results support the suitability of our pan-avian primer set to identify appropriate reference genes for gene expression studies on avian cells and tissues.

**Discussion**

Gene expression studies in diverse bird species are used to explore the molecular mechanisms underlying physiology and behavior. Due to the richness of species, the comparative
approach to understand the evolution of particular traits is especially promising in the avian lineage. As more genomes are being sequenced, this endeavor becomes increasingly feasible. Frequently quantitative gene expression data derived from qPCR experiments are used as a first step to probe functional consequences of gene activity [8,51,52]. For these data to be meaningful it is essential to choose an appropriate method to normalize the expression level of the gene of interest.

Because the genomes of only very few avian species have been fully explored to date [7,9,26,27], it can be difficult and time consuming to establish primers for qPCR which work on all species under study. Systematic evaluations of reference genes for avian species other than the chicken [33,53,54] are scarce. The present study aimed to establish primers for a set of 14 candidate reference genes and to test whether they can be used to normalize gene expression in qPCR experiments on any given avian species. First we tested the adequacy of the primer pairs for each of our candidate genes on nine different avian species spanning the phylogenetic tree from the ostrich to the zebra finch. Subsequently, we evaluated the suitability of the tested genes as reference genes on samples from zebra finch brain microbiopsies. Moreover, we examined the expression levels of the candidate reference genes in gonads and brains from male and female zebra finches.

Of the 14 primer pairs tested, six to 11 turned out to match all our quality criteria on each species. Even though the GAPDH primers amplified properly on all species tested, we excluded the primers because of the low amplification efficiency for the zebra finch and chicken. Although we have not yet tested the primers RPL13 and RPL19 on all nine avian species of this study, our initial results from the chicken, ostrich and zebra finch indicate that both could serve as additional sets to the established ones. Additionally, we previously used RPL13 and RPL19 for normalization in the domestic pigeon (Columba livia f. dom.) [36] and the cockatiel (A. Meyer, personal communication).

The overall evaluation of all our analyses revealed that PGK1 was consistently among the most stably expressed genes. Notably, the results calculated by the three algorithms employed (BestKeeper, geNorm, NormFinder) varied considerably between different tissues and sexes. No single gene ranked as the most stable one in all tissues tested. This emphasizes the importance of determining the most reliable reference genes for each new experimental design [11]. This point gains even more emphasis when comparing the results of the stability analysis in the song

Figure 4. Singing induced gene regulation in Area X of adult male zebra finches. Relative gene expression levels for EGR1, CFOS and FOXP2 were calculated by normalizing to the geometric mean of the two genes recommended by each of the three different algorithms. Relative expression values of the singing group are compared to the silent group. Statistical differences were calculated using the Mann Whitney U test and are indicated in the diagram (p<0.05 *, p<0.01 **, no significant difference ns). doi:10.1371/journal.pone.0099678.g004
needs to be cautious to only use it if the experimental conditions
RNA Polymerase II [20]. RNA Polymerase I whereas synthesis of mRNA is performed by
the 18S gene for all but the most abundantly expressed genes. Additionally, PGK1 is one of the most stably expressed genes in the female brain, while it is among the least stable genes in the male brain. The sexually dimorphic variation of PGK1 and YWHAZ in the zebra finch brain might point towards hormonal regulation of gene expression or even a functional role in, e.g. the song system, which is fully developed only in male zebra finches.

As expected, 18S was the most abundant mRNA and GUSB the least abundant mRNA in all samples tested (Figure 3). In fact, in the song data set GUSB expression was below the detection limit of the qPCR-assay. The use of 18S to normalize gene expression in a qPCR experiment has been heavily debated in the past [20,55,56]. A control gene should be expressed at roughly the same level as the gene of interest to minimize the influence of the technical error [57]. It thus might be advisable to avoid using 18S as reference gene for all but the most abundantly expressed genes. Additionally, 18S might not be a suitable reference gene as it is produced by RNA Polymerase I whereas synthesis of mRNA is performed by RNA Polymerase II [20]. 18S is not polyadenylated and one thus needs to be cautious to only use it if the experimental conditions allow it. For example, if the RNA for the experiment was extracted via oligo(dT) purification or reverse transcribed using oligo(dT)’s, 18S should clearly not be used as reference gene. In spite of this criticism, 18S has repeatedly been used to normalize gene expression in qPCR experiments and we thus decided not to exclude it from our results [56,58–60]. In conclusion, the practical and easy-to-use pan-avian reference gene primer panel will greatly facilitate molecular research in multiple avian species.

Supporting Information

Document S1 FASTA file of all sequencing results yielded from the PCR products during establishing the primer pairs on nine different species. (FASTA)

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Author Contributions

Conceived and designed the experiments: PO IA AM ADG. Performed the experiments: PO AM IA. Analyzed the data: PO AM IA. Contributed reagents/materials/analysis tools: PO IA CS ADG. Wrote the paper: PO IA CS ADG.

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42. Pan Avian Primer Set to Select Reference Genes