Origin of Chinese Goldfish and Sequential Loss of Genetic Diversity Accompanies New Breeds

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Abstract

**Background:** Goldfish, *Carassius auratus*, have experienced strong anthropogenic selection during their evolutionary history, generating a tremendous extent of morphological variation relative to that in native *Carassius*. To locate the geographic origin of goldfish, we analyzed nucleotide sequences from part of the control region (CR) and the entire cytochrome b (Cytb) mitochondrial DNA genes for 234 goldfish and a large series of native specimens. Four important morphological characteristics used in goldfish taxonomy–body shape, dorsal fin, eye shape, and tailfin–were selected for hypothesis-testing to identify those that better correspond to evolutionary history.

**Principal Finding:** Haplotypes of goldfish rooted in two sublineages (C5 and C6), which contained the haplotypes of native *C. a. auratus* from southern China. Values of *F*<sub>ST</sub> and *N*<sub>mt</sub> revealed a close relationship between goldfish and native *C. a. auratus* from the lower Yangtze River. An extraordinary, stepwise loss of genetic diversity was detected from native fish to goldfish and from Grass-goldfish relative to other breeds. Significantly negative results for the tests of Tajima’s *D* and Fu and Li’s *D*<sup>*</sup> and *F*<sup>*</sup> were identified in goldfish, including the Grass breed. The results identified eye-shape as being the least informative character for grouping goldfish with respect to their evolutionary history. Fisher’s exact test identified matrilineal constraints on domestication.

**Conclusions:** Chinese goldfish have a matrilineal origin from native southern Chinese *C. a. auratus*, especially the lineages from the lower Yangtze River. Anthropogenic selection of the native *Carassius* eliminated aesthetically unappealing goldfish and this action appeared to be responsible for the stepwise decrease in genetic diversity of domesticated goldfish, a process similar to that reported for the domestication of pigs, rice, and maize. The three-breed taxonomy–Grass-goldfish, Egg-goldfish, and Wen-goldfish–better reflected the history of domestication.

Citation: Wang S-Y, Luo J, Murphy RW, Wu S-F, Zhu C-L, et al. (2013) Origin of Chinese Goldfish and Sequential Loss of Genetic Diversity Accompanies New Breeds. PLoS ONE 8(3): e59571. doi:10.1371/journal.pone.0059571

Editor: Axel Janke, BiK-F Biodiversity and Climate Research Center, Germany

Received August 3, 2012; Accepted February 19, 2013; Published March 19, 2013

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Funding: This work was supported by grants from the State Key Basic Research and Development Plan (2007CB411600), Bureau of Science and Technology of Yunnan Province (2008GA001), and ‘Western Light programme’ of the Chinese Academy of Sciences to Y.G. and J.L. was funded by the State Key Laboratory of Genetic Resources and Evolution, and Natural Sciences and Engineering Research Council (Canada) Discovery Grant 3148 to R.W.M. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction

Goldfish, one of the first animals domesticated for ornamental purposes, has experienced extreme anthropogenic selection during its evolutionary history to create aesthetically appealing forms [1,2]. Widely distributed across Eurasia [3], native feral *Carassius* (crucian carp) can naturally change their body color from gray to red [1,2]. Feral red goldfish are thought to be the ancestral forms of Chinese goldfish [1,2].

The ability to change color has led to aquiculture of the fish for use in religion [4]. The earliest record of anthropogenic usage dates to the Ts’ìn Dynasty (265–419 A.D.) of China as noted in the Compendium of Materia Medica [4]. Strong anthropogenic selection during cultivation is likely responsible for much of the phenotypic variation seen today [1,2,4]. Some goldfish possess features such as egg-shaped bodies, celestial or telescopic eyes, fancy tailfins, lionhead morphotypes, a raspberry-like hood encasing the head (oranda), no dorsal fin, and other variants [2,5]. Around 1502 A.D. goldfish were exported to Japan [1] and around 1700 A.D. to Europe [1,2,5].

The tremendous extent of morphological variation in goldfish owing to anthropogenic selection causes difficulties in evolutionary taxonomy. Linnaeus (1758) originally named the goldfish as *Cyprinus auratus* because morphologically it is similar to the common carp, *Cyprinus carpio* [6]. Subsequently, *Cyprinus auratus* was transferred to the genus *Carassius* as *Carassius auratus* [6]. Several taxonomic schemes (Table 1) exist in China for goldfish, each of which focuses on different morphological features [4,7–9]. Most frequently, three terms are used to designate breeds: Grass-goldfish, Wen-goldfish, and Egg-goldfish. Assignment of an individual fish to one breed or another depends on body-shape (slender or egg-shaped) and condition of the dorsal fin (retained or
better mirrors evolutionary history, and not merely the extent of written history details the sequential development of the breeds. Thus, the Dragon-dorsal-goldfish has dragon-eyes and no dorsal fin [9]. Although the numbers of tailfins (single or double) is not a standard for grouping goldfish, it is an important morphological characteristic used to describe the breeds [4,7–9]. The detailed descriptions of the previous four morphological characteristics for the three taxonomic schemes are listed in Table 1. Taxonomy is more informative when based on evolutionary history, but the various taxonomies (Table 1) focus only on human-selected morphological characteristics and likely obscure history. Unfortunately, no written history details the sequential development of the breeds. This necessitates a reassessment of goldfish-taxonomy to assure it better mirrors evolutionary history, and not merely the extent of morphological divergence.

Chen [10] reported reproductive viability in hybrids between breeds of goldfish and native Carassius. The muscle proteins of native Carassius are similar to those of goldfish [11]. Analyses of nucleotide sequence data from partial fragments of mitochondrial DNA (mtDNA) control region (CR; 471 bp), also known as the D-loop, obtain the same conclusion [12]. Komiyama et al. [13] analyzed a portion of the mitochondrial genome (740 bp) from 67 specimens of Carassius including 44 specimens of goldfish, and further 11180 bp from seven specimens of goldfish. Although they evaluated most of the mitochondrial genome, their sampling on mainland China was limited. Their matrilineal history hypothesizes that the ancestral breed is the Gibelio group of Chinese Carassius. Our prior research on the biogeography of the East Asian C. auratus complex used 1876 partial CR (426 bp) and 187 complete cytochrome b (Cyb; 1140 bp) gene sequences from 67 localities representing most of the species’ range and identified three distinct, mostly geographically constrained matrilines [14]. These analyses provide an opportunity to investigate the origin and domesticated history of goldfish from the perspective of the large population of native Carassius.

Herein, we reconstruct the matrilineal relationships using de novo sequences of goldfish as well as mtDNA data of wild Carassius from prior studies [14]. Analyses are used to infer the geographic origin of goldfish and to investigate the genetic consequences of extreme anthropogenic selection. We use CR because of its high mutation rate, which facilitates the resolution of intraspecific matrilineal relationships [15–18]. We also employ nucleotide sequences of Cyb because this gene is less subject to substitution-saturation, which makes it more reliable than CR for evaluating interspecific relationships [15,17–20]. Furthermore, we select several important morphological characteristics of goldfish taxonomy in China for hypothesis-testing to identify those that better correspond to evolutionary history.

### Materials and Methods

#### Ethics Statement

All samples of fish from China used in this study were obtained and handled following the guidelines of the by-laws on experimentation on animals, and was approved by the Ethics and Experimental Animal Committee of Kunming Institute of Zoology, Chinese Academy of Science, China (KIZ_YP2010002).

#### Sampling and Molecular Methods

One hundred and ninety specimens of goldfish were collected from Hangzhou (37 specimens), Kunming (56 specimens), Changchun (20 specimens), Lanzhou (23 specimens) and Beijing (22 specimens), China, Seoul (11 specimens), South Korea, and Toronto (21 specimens), Canada. In addition, 5 sequences for Chinese goldfish [13], 39 sequences for Japanese goldfish [13] as well as 1876 sequences for native Carassius with detailed sampling localities and haplotypes [14] were obtained from GenBank (Table 2). Samples used for morphological analyses were photographed and then stored as voucher specimens. Either tailfin clips or muscle tissue samples were collected from individuals and stored at −20°C until processing. Genomic DNA from freshly frozen or ethanol-fixed tissues was extracted by the standard phenol/chloroform method.

PCR amplifications were performed in a total volume of 50 μl containing 1x buffer, 0.15 mM MgCl₂ (Sina-American, Beijing, China), 0.25 mM dNTPs (Amersco, Solon, OH, USA), 1 U Taq DNA polymerase (Sina-American) and 25–50 ng total DNA. Primers for amplification were identical to those of Gao et al. [14]. Amplifications were performed on a Gene Amp PCR system 9700 (Applied Biosystems, Foster City, CA, USA) following the conditions: pre-denaturation at 96°C for 2 min followed by 30 cycles of denaturation-annealing-elongation (96°C, 1 min; CR: 58°C and Cyb: 50°C, 1 min; 72°C, 1 min) and a final extension at 72°C for 10 min. The corresponding PCR products were purified on agarose gels and extracted (Watson Biomedical Inc., Shanghai). Sequencing reactions were run on a 3730 (ABI) with ABI PRISM BigDye Terminator Cycle Sequencing Ready Reaction

| Morphology | Three-breed system | Four-breed system | Five-breed system |
|------------|--------------------|------------------|------------------|
| Body-shape (slender vs. egg-shaped) | Grass- (slender); Wen- (Egg-shaped); Egg- (Egg-shaped); Dragon- (Egg-shaped) | Grass- (slender); Wen- (Egg-shaped); Egg- (Egg-shaped) | Grass- (slender); Wen- (Egg-shaped); Egg- (Egg-shaped) |
| Condition of dorsal fin (retained vs. loss) | Grass- (retained); Wen- (lost); Egg- (lost) | Grass- (retained); Wen- (retained); Egg- (retained or loss) | Grass- (retained); Wen- (retained); Egg- (loss); Dragon- (retained or loss) |
| Eye shape (normal vs. extended) | Grass- (normal); Wen- (normal or extended); Egg- (normal or extended) | Grass- (normal); Wen- (normal or extended); Egg- (normal or extended); Dragon- (normal or extended) | Grass- (normal); Wen- (normal or extended); Egg- (normal or extended); Dragon- (normal or extended) |
| Numbers of tailfins (single vs. double) | Grass- (single); Wen- (single or double); Egg- (double) | Grass- (single); Wen- (single or double); Egg- (double); Dragon- (single or double) | Grass- (single); Wen- (single or double); Egg- (double); Dragon- (single or double) |

Table 1. Features of different breed-systems of goldfish based on morphology.
Analyses of Sequence Data

Sequences were assembled using DNASTAR v.5.0 (DNASTAR Inc., Madison, WI, USA) and manually verified. Sequence alignments and information on nucleotide variation were obtained using MEGA v.4.0 [21,22]. DAMBE v.4.1.19 [23] was used to identify shared haplotypes. The new haplotypes identified from goldfish, plus the 180 haplotypes of combined CR and Cytb from a prior study [14], were used for the new reconstruction of the matrilineal genealogy.

Phylogenetic analyses were conducted using maximum likelihood (ML) and maximum parsimony (MP) in PAUP* 4.0b10 [24], and Bayesian inference (BI) in MrBayes v.3.0b4 [25]. All analyses were based on the concatenated Cytb and CR data. Likelihood ratios tests [26–28] implemented in MODELTEST v.3.7 [29] were employed to select the best-fitting models for the ML and BI analyses. The GTR+I+G model was selected for the combined dataset by the Akaike Information Criterion [30]. In the ML and MP analyses, a heuristic search with 100 random additions replicates was involved. BI used four simultaneous Markov-coupled Monte Carlo Markov chains running for 5,000,000 generations. Convergence to stationarity was evaluated by TRACER v.1.5 [31] using log-likelihood values. The first 50% of the trees were discarded as burn-in and the remaining tree samples were used to generate a consensus tree. Nodal support for the ML and MP tree building methods was assessed using nonparametric bootstrapping (BS) [32] calculated in PAUP* for the MP analysis (MPBS) and RAxML [33] for ML (MLBS) using 1000 pseudoreplicates each. Bayesian posterior probability (BPP) values, the frequency of nodal resolution in the majority rule consensus tree, were calculated through the BI analysis, and the BS for each node in MP as well as ML reconstructions were plotted on the tree.

According to the historical distribution of East Asian *Carassius* [3,34–38], we classified the 67 sampling localities for native *Carassius* from a prior study [14] into four geographic regions: northern China (NC); southern China (SC); Japan (JA); and Europe (Russia and Czech Republic; EU). Sampling localities for

### Table 2. Summary statistics for the CR and Cytb markers used in the study.

| Lineages | N  | V  | P  | NH | Hd  | D  | DP | F*  |
|----------|----|----|----|----|-----|----|----|-----|
| **CR (426 bp)** |    |    |    |    |     |    |    |     |
| *Carassius carassius* | 9  | 2  | 0  | 3  | 0.4167±0.1907 | 0.0030±0.0024 | −1.362 | −1.505 | −1.626 |
| Native *Carassius auratus* complex | 1867 | 98 | 86 | 208 | 0.9673±0.0017 | 0.1564±0.0746 | 0.169 | −1.218 | −0.628 |
| Lineage A: *C. a. cuvieri* | 5  | 1  | 0  | 3  | 0.7000±0.2184 | 0.0019±0.0018 | −0.817 | −0.816 | −0.772 |
| Lineage B: *C. a. langsfordii* | 421 | 42 | 37 | 46 | 0.9531±0.0032 | 0.0560±0.0272 | 0.665 | 0.471 | 0.674 |
| Sublineage C1 | 62  | 16 | 15 | 8 | 0.8636±0.0266 | 0.0129±0.0070 | −0.153 | 0.679 | 0.461 |
| Sublineage C2: *C. a. gibelio* | 243 | 26 | 19 | 30 | 0.5425±0.0387 | 0.0152±0.0080 | −1.678 | −2.008 | −2.262 |
| Sublineage C3 | 60  | 16 | 6 | 9  | 0.7994±0.0299 | 0.0132±0.0071 | −1.308 | −2.999 | −2.860 |
| Sublineage C4 | 277 | 17 | 16 | 18 | 0.5012±0.0358 | 0.0513±0.0251 | −0.475 | 0.073 | −0.163 |
| Sublineage C5 and C6: *C. a. auratus* | 799 | 57 | 47 | 87 | 0.9397±0.0040 | 0.0929±0.0447 | −1.336 | −0.879 | −1.323 |
| **Goldfish** |    |    |    |    |     |    |    |     |
| Egg-goldfish | 60  | 2  | 2  | 4  | 0.6471±0.0458 | <0.0001 | 0.964 | 0.734 | 0.933 |
| Wen-goldfish | 92  | 2  | 1  | 3  | 0.1949±0.0667 | <0.0001 | −0.993 | −1.059 | −1.214 |
| **Cytb (1140 bp)** |    |    |    |    |     |    |    |     |
| *Carassius carassius* | 4  | 0  | 0  | 1  |     |    |    |     |
| Native *Carassius auratus* complex | 187 | 224 | 184 | 103 | 0.9806±0.0044 | 0.0257±0.0125 | −0.765 | 0.314 | −0.634 |
| Lineage A: *C. a. cuvieri* | 2  | 1  | 0  | 2  |     |    |    |     |
| Lineage B: *C. a. langsfordii* | 24 | 80 | 44 | 20 | 0.9855±0.0159 | 0.0154±0.0079 | −0.704 | −1.109 | −1.152 |
| Sublineage C1 | 15 | 37 | 27 | 12 | 0.9714±0.0327 | 0.0092±0.0050 | −0.345 | −2.060 | −2.180 |
| Sublineage C2: *C. a. gibelio* | 19 | 14 | 8 | 10 | 0.9664±0.0446 | 0.0030±0.0018 | −0.493 | −0.628 | −0.683 |
| Sublineage C3 | 9  | 10 | 5 | 4  | 0.6944±0.1470 | 0.0028±0.0018 | −0.649 | −0.306 | −0.434 |
| Sublineage C4 | 32 | 29 | 10 | 10 | 0.7258±0.0749 | 0.0050±0.0027 | −0.757 | −2.700 | −2.434 |
| Sublineage C5 and C6: *C. a. auratus* | 84 | 64 | 40 | 45 | 0.9541±0.0149 | 0.0062±0.0033 | −1.573 | −2.736 | −2.726 |
| **Goldfish** |    |    |    |    |     |    |    |     |
| Grass-goldfish | 65 | 25 | 6 | 7  | 0.4509±0.0684 | 0.0017±0.0011 | −1.943 | −4.737 | −4.423 |
| Egg-goldfish | 34 | 0  | 0  | 1  |     |    |    |     |
| Wen-goldfish | 81 | 1  | 0  | 2  | 0.0230±0.0022 | <0.0001 | −1.043 | −1.995 | −1.992 |

N, number of sequences; V, variable sites of sequences; P, potentially parsimony-informative sites of sequences; NH, number of haplotypes; Hd, haplotype diversity; D, nucleotide diversity; D*, Tajima's D-statistic; DP, Fu and Li's D*-statistic; F*, Fu and Li's F*-statistic; *P<0.05. **P<0.01.

doi:10.1371/journal.pone.0059571.t002

Kit (ABI) following the manufacturer’s recommendations. All PCR products were sequenced in both directions.
native *Carassius* in China [14] were further classified into eight regions according to the distribution of Chinese drainages: Yunnan-Guizhou Plateau, Yellow River, Pearl River, middle Yangtze River, lower Yangtze River, Minjiang River, Amur River, and inland rivers in Xinjiang. We used ARLEQUIN v.3.1 [39] to calculate pairwise *F*<sub>ST</sub> values and the number of migrants in each generation (*N*<sub>m</sub>) based on the CR datasets. The analyses identified the extent of divergence and inferred gene flow between the native *Carassius* from these regions and goldfish. An analysis of molecular variance (AMOVA), implemented in ARLEQUIN, was used to evaluate the genetic divergence in goldfish from southern China (Hangzhou and Kunming), northern China (Changchun, Lanzhou and Beijing), Canada, South Korea and Japan. Haplotype diversity (*H*<sub>h</sub>) and nucleotide diversity (π) values for the different lineages of native *Carassius* and breeds of Chinese goldfish were calculated based on CR and *Cytb* datasets respectively using ARLEQUIN. The possible effects of demographic events during goldfish domestication were examined by the tests of Tajima's [40] and Fu and Li's [41] to calculate pairwise *F*<sub>ST</sub> values and the number of migrants in each generation (*N*<sub>m</sub>) based on the CR datasets. The analyses identified the extent of divergence and inferred gene flow between the native *Carassius* from these regions and goldfish. An analysis of molecular variance (AMOVA), implemented in ARLEQUIN, was used to evaluate the genetic divergence in goldfish from southern China (Hangzhou and Kunming), northern China (Changchun, Lanzhou and Beijing), Canada, South Korea and Japan.

Haplotype diversity (*H*<sub>h</sub>) and nucleotide diversity (π) values for the different lineages of native *Carassius* and breeds of Chinese goldfish were calculated based on CR and *Cytb* datasets respectively using ARLEQUIN.

Hypothesis-tests of Morphology and Genealogy

We tested for the correspondence between the three methods of morphologically grouping Chinese goldfish and genealogical history based on the concatenated dataset. The morphological data consisted of the four characters traditionally used for Chinese goldfish taxonomy (Table 1): body-shape (slender or egg-shaped), presence or absence of the dorsal fin, the eye shape (normal or derived), and single versus double tailfins. We identified specific haplotypes that were constrained for one morphological condition, and *C. carassius* was used as the outgroup taxon to determine character state polarities. For example, we evaluated whether the specific haplotypes for slender body-shape was constrained to one matriline, or not. The morphology-based trees were compared to the best unconstrained molecular tree. A MP molecular tree that represented a particular morphological topology was estimated using constrained tree searches in PAUP*, and a heuristic search with 100 random additions replicates was involved for each analysis. Each of the constrained trees was compared to the unconstrained MP topology using a non-parametric Templeton test [43] in PAUP*. Constrained and unconstrained topologies were similarly calculated under the ML criterion in a heuristic search with 100 random additions replicates and compared using the Shimodaira and Hasegawa [44] test (SH) implemented in PAUP*.

A Fisher's exact test was used to examine the matrilineal distribution of goldfish based on the concatenated *Cytb* and CR datasets. DNASP [42] calculated the polymorphic sites (variable and potentially parsimony-informative sites) for native *Carassius* and goldfish, respectively, and the Fisher's exact test was performed using SPSS version 13.0.

Results

Haplotype Nomenclature

To avoid confusion, we employed a nomenclature to distinguish haplotypes obtained from the two genes. Haplotypes starting with ‘h’ were used to denote CR data, those with the prefix ‘Jap’ were CR sequences unique to Japanese goldfish, and those starting with ‘B’ indicated *Cytb* data only. The designations were combined to indicate total mtDNA variation. Accordingly, a haplotype consisting of CR haplotype h13 and *Cytb* haplotype B10 was termed h13B10.

Sequence Variation

Sequence variation in CR and *Cytb* markers of goldfish was summarized in Table 2. The CR sequences (426 bp) of goldfish contained only 11 variable sites of which eight were potentially parsimony-informative. Analyses identified nine haplotypes from 234 specimens. Two haplotypes (Jap1, Jap2) were unique to goldfish, and seven were shared with native *Carassius*. Haplotype h20 was most common in goldfish, being shared by 181 specimens. Among the 1140 bp of *Cytb* data, 25 sites exhibited variation and among these only six sites were potentially parsimony-informative. For *Cytb*, eight haplotypes were identified from the 180 sequences of goldfish, of which three (B105–B107) were not shared with native *Carassius*. Shared by 160 specimens, haplotype B13 was the most common one. Combined, the CR and *Cytb* data identified 12 haplotypes. Seven of these haplotypes—h1_2B13, h20B13, h55B105, h55B106, h56B107, Jap1B13, and Jap2B13—were unique to goldfish. One hundred and forty seven specimens of goldfish shared the most common haplotype, h20B13; this haplotype was not shared with native *Carassius*. Only five haplotypes—h1_2B22, h13B10, h19B13, h32B25, and h36B9—were shared with native *Carassius*. More specimens of native *Carassius* were sequenced for CR (1876) than *Cytb* (187) [14] and this may have resulted in the resolution of a greater number of haplotypes unique to goldfish for *Cytb*.

Matrilineal History

Bayesian inference (BI), maximum likelihood (ML), and maximum parsimony (MP) analyses of the concatenated CR and *Cytb* sequences yielded the same topology (Fig. 1) and this was the same as that of Gao et al. [14]. Haplotypes identified in *C. a. cuvieri* (Lineage A) and *C. a. longidorsus* (Lineage B) were not found in goldfish. All haplotypes identified in the gibel carp (a.k.a. Gibelio) clustered together forming sublineage C2; none of these haplotypes was detected in goldfish. Haplotypes identified from goldfish occurred in sublineages C5 and C6 (Fig. 1) only. In our genealogy, sublineages C5 (*MLBS = 93%, MBPS = 99%, BPP = 100%) and C6 (*MLBS = 60%, MBPS = 56%, BPP = 100%) both containing the haplotypes identified in native *C. a. auratus* that generally occurred in southern China [14]. Sublineage C5 contained native *C. a. auratus*, fish especially associated with the Yangtze River [14]. Eleven of 12 haplotypes identified from goldfish clustered in sublineage C6, and only one haplotype (h32B13) from Grass-goldfish located in sublineage C5.

Genetic Divergence between Native *Carassius* and Goldfish

The *F*<sub>ST</sub> value between goldfish and native *Carassius* from southern China (SC) was substantially lower (0.2157) than that between goldfish and native *Carassius* from northern China (NC; 0.9958), Europe (EU; 0.8942), or Japan (JA; 0.7461). Values of *N*<sub>m</sub> also indicated more gene flow between the goldfish and native *Carassius* from SC (2.9847) than between goldfish and the native *Carassius* from NC (0.0234), EU (0.1337), and JA (0.2922). Values of *F*<sub>ST</sub> and *N*<sub>m</sub> between the native *Carassius* from eight Chinese drainages and goldfish were also calculated (Table 3). Native *Carassius* from the lower Yangtze River and goldfish had the lowest *F*<sub>ST</sub> value (0.2829). Values of *N*<sub>m</sub> indicated strong gene flow between goldfish and native *Carassius* from the lower Yangtze River (*N*<sub>m</sub> = 1.2674) and Yunnan-Guizhou Plateau (*N*<sub>m</sub> = 1.2580).

We classified the goldfish by the three-breed taxonomy and subsequently compared the divergence between these breeds and native *C. a. auratus*. For the CR dataset, the *F*<sub>ST</sub> value between
southern Chinese *C. a. auratus* (sublineages C5 and C6) and Grass-goldfish (0.0591) was lower than that between either Egg-goldfish (0.1355) or Wen-goldfish and southern Chinese *C. a. auratus* (0.1164). The Nm value between southern Chinese *C. a. auratus* and Grass-goldfish (6.5712) was higher than between Egg-goldfish (3.0126) or Wen-goldfish (3.4842) and southern Chinese *C. a. auratus*. Values of $F_{ST}$ among the goldfish from Hangzhou, Kunming, Changchun, Lanzhou, Beijing, Seoul, Toronto and Japan were low (0.0000–0.1805), and Nm values were always high (2.2698–infinity). The results of AMOVA also revealed no significant genetic divergence ($F_{CT} = 0.236$, $P > 0.5$) among southern China (Hangzhou and Kunming), northern China (Changchun, Lanzhou and Beijing), Canada, South Korea and Japan.

**Genetic Diversity**

Haplotype (Hd) and nucleotide ($\pi$) diversity based on CR and Cytb sequences separately (Table 2) of sublineages C5 and C6 (CR: $H_d = 0.9397 \pm 0.0040$, $\pi = 0.0929 \pm 0.0447$; Cytb: $H_d = 0.9541 \pm 0.0149$, $\pi = 0.0062 \pm 0.0033$) were always higher than in goldfish (CR: $H_d = 0.3782 \pm 0.0401$, $\pi = 0.0023 \pm 0.0018$; Cytb: $H_d = 0.1971 \pm 0.0384$, $\pi = 0.0007 \pm 0.0006$). These results revealed a reduction of genetic diversity from native goldfish to aquaculture goldfish.

We also classified the goldfish by the three-breed taxonomy and subsequently compared their levels of genetic diversity. Haplotype h19 was shared by the Egg-goldfish and Wen-goldfish. Haplotypes Jap1 and Jap2 only occurred in Egg-goldfish, which have double tail fins and no dorsal fins. Six Cytb haplotypes were unique to Grass-goldfish and only haplotype B13 occurred in all three breeds. Haplotype B9 was detected in Wen-goldfish only. Grass-goldfish had a higher level of genetic diversity (Table 2) for Cytb (CR: $H_d = 0.4436 \pm 0.0635$, $\pi = 0.0045 \pm 0.0029$; Cytb: $H_d = 0.4509 \pm 0.0684$, $\pi = 0.0017 \pm 0.0011$) than either Egg-goldfish (CR: $H_d = 0.6471 \pm 0.0458$, $\pi < 0.0001$; Cytb: all the specimens of Egg-goldfish shared the haplotype B13) or Wen-goldfish (CR: $H_d = 0.1949 \pm 0.0667$, $\pi < 0.0001$; Cytb: $H_d = 0.0230 \pm 0.0022$, $\pi < 0.0001$). These results revealed the loss of genetic diversity also occurred during the anthropogenic transition from Grass-goldfish to other breeds.

**Neutrality Tests**

Values for Tajima’s $D$ (CR: $-1.083$; Cytb: $-2.266$) and Fu and Li’s $D^*$ (CR: $-0.871$; Cytb: $-6.423$) and $P^*$ (CR: $-1.139$; Cytb: $-5.703$) were all negative in the goldfish and statistically significant ($P < 0.01$) in our analysis based on the Cytb dataset (Table 2); Tajima’s $D$ (CR: $-0.162$; Cytb: $-1.943$) and Fu and Li’s $D^*$ (CR: $-0.636$; Cytb: $-4.737$) and $P^*$ (CR: $-0.560$; Cytb: $-2.668$) were also negative in the goldfish.
Hypothesis-testing for Grouping Breeds

Four morphological constraint trees and the unconstrained best tree were represented in Figure S1. Results of the Templeton and SH tests were summarized in Table 4. The Templeton test rejected body-shape and eye-appearance \((P < 0.05)\) as being correlated with matrilineal history. P-values for the SH test showed that the best unconstrained ML topology differed significantly from the morphological constraint tree for eye-appearance \((P < 0.05)\). Differences in ln L values also revealed that the condition of the goldfish’s eye (normal or derived) was the least informative character for grouping by history \((23.66)\), followed by body-shape (egg-shaped or slender, 14.22), and dorsal fin (retained or loss, 14.22). The number of tailfins (single or double, 4.61) was most indicative of genealogical history. Therefore, the three-breed scheme (Grass-goldfish, Wen-goldfish, and Egg-goldfish) better reflected history than either the four-breed or the five-breed systems that emphasized eye condition.

Fisher’s exact test based on the concatenated \(Cytb\) and \(CR\) datasets obtained a highly significant \((P < 0.001)\) relationship between genetic variation sites and lineage. This indicated that the domestication of goldfish was constrained to particular matrilines.

Discussion

Origin of Goldfish

Our analyses suggest that Chinese goldfish have a matrilineal origin from native southern Chinese \(C. a. auratus\), especially lineages from the lower Yangtze River. The genealogical analyses resolve the origin of goldfish from native Chinese \(Carassius\), a finding consistent with that of Komiyama et al. [13]. The matrilineal genealogy (Figure 1) and \(F_{ST}\) values (Table 3) further indicate a much closer relationship between the goldfish and sublineages C5 and C6 of \(C. a. auratus\) from southern China rather than the gibel carp (sublineage C2) from northern China. This discovery differs from the suggestion of the origin of the goldfish being from the gibel carp [13]. Values of \(N_m\) (Table 3) also suggest strong gene flow occurs between goldfish and native \(Carassius\) from the lower Yangtze River. All analyses are consistent with the historical record, which suggests that Hangzhou and Jiaxin, Zhejiang, China might be the area of domestication [1]. Our analyses do not detect significant genetic divergence among the different regions where goldfish live; strong gene flow appears to occur among these regions. These results are not surprising considering the long history of commercialization, artificial selection, and hybridization among different breeds and regions of goldfish [1,2,4].

Other evidence excludes the gibel carp from being the ancestor of goldfish. Gibel carp are usually hexaploids with more than 150 chromosomes [45]. In contrast, goldfish are always tetraploids and have around 100 chromosomes [4,46]. Further, the historical distribution of the gibel carp (\(C. a. gibelio\)) appears to be restricted to the northern Amur River system and Europe [47–50]. Historically, the distributions for goldfish and the gibel carp did not overlap. Therefore, our resolution of a southern origin for goldfish is valid not only because of the strength of the historical geography and ploidy levels of these fishes, but also because of the incontrovertible exclusion of the matrilines of the gibel carp.

~4.423) values were also significantly negative in Grass-goldfish for the \(Cytb\) dataset (Table 2).

Table 3. Pairwise \(F_{ST}\) values and the numbers of migration in every generation \((N_m)\) between the wild \(Carassius\) from eight Chinese drainages and goldfish; pairwise \(F_{ST}\) values were below the diagonal; the numbers of migration in every generation \((N_m)\) were above the diagonal.

| Pairwise Carassius | Lower Yangtze River | Yunnan-Guizhou Plateau | Yellow River | Pearl River | Middle Yangtze River | Minjiang River | Inland rivers in Xinjiang | Amur River | Goldfish |
|-------------------|---------------------|------------------------|-------------|------------|----------------------|---------------|--------------------------|-----------|---------|
| Lower Yangtze River | 0.0137 | 0.0599 | 0.0395 | 0.0307 | 0.0293 | 0.0165 | 0.0186 | 0.0292 | 0.2829 |
| Yunnan-Guizhou Plateau | 0.0599 | 0.0478 | 0.0436 | 0.0307 | 0.0293 | 0.0165 | 0.0186 | 0.0292 | 0.2829 |
| Yellow River | 0.0395 | 0.0599 | 0.0448 | 0.0436 | 0.0307 | 0.0293 | 0.0165 | 0.0186 | 0.2829 |
| Pearl River | 0.0307 | 0.0395 | 0.0478 | 0.0448 | 0.0436 | 0.0307 | 0.0293 | 0.0165 | 0.2829 |
| Middle Yangtze River | 0.0293 | 0.0307 | 0.0395 | 0.0599 | 0.0478 | 0.0436 | 0.0307 | 0.0293 | 0.2829 |
| Minjiang River | 0.0165 | 0.0293 | 0.0307 | 0.0395 | 0.0478 | 0.0448 | 0.0436 | 0.0307 | 0.2829 |
| Inland rivers in Xinjiang | 0.0186 | 0.0165 | 0.0293 | 0.0307 | 0.0478 | 0.0448 | 0.0436 | 0.0307 | 0.2829 |
| Amur River | 0.0292 | 0.0292 | 0.0292 | 0.0292 | 0.0292 | 0.0292 | 0.0292 | 0.0292 | 0.2829 |
| Goldfish | 0.2829 | 0.2829 | 0.2829 | 0.2829 | 0.2829 | 0.2829 | 0.2829 | 0.2829 | 0.2829 |
Domestication History of Goldfish

Anthropogenic selection of native Carassius eliminated aesthetically unappealing goldfish and this action appears to be responsible for the stepwise decrease in genetic diversity of domesticated goldfish, i.e., the loss of genetic variation from native goldfish to Grass-goldfish in aquiculture followed by further loss within remaining breeds of goldfish. A strong reduction of genetic diversity should accompany the domestication and this is seen as a recent bottleneck event or founder effects, which occurs in domesticated pigs [51,52], maize [53], and rice [54]. Both the extraordinarily lower genetic diversity and the significantly negative results for the tests of Tajima's D and Fu and Li's D* and F* (Table 2) indicate founder effects and bottlenecking during the domestication of goldfish. Based on recorded history, native red Carassius were initially herding in ‘free life ponds’ at many temples in Hangzhou and Jiaxin, Zhejiang, China, and without anthropogenic breeding selection [1,4]. The morphology of the Grass-goldfish is less derived and more similar to the native Carassius than other breeds [1,7,8]. Our analyses reveal that the Grass-goldfish has higher level of genetic diversity than either Egg-goldfish or Wen-goldfish. The FST values also indicate that Grass-goldfish and southern Chinese C. a. auratus differ less from each other than do either Egg-goldfish or Wen-goldfish from southern Chinese C. a. auratus. These findings indicate that the Grass-goldfish is likely the first domesticated breed of Chinese goldfish.

Strong anthropogenic selection for aesthetics is likely responsible for the further loss of genetic diversity among different breeds of goldfish. Our analyses for the three-breed system detect a further decrease in genetic diversity from Grass-goldfish to Egg-goldfish (Table 2). Tajima's D and Fu and Li's D* and F* in Grass-goldfish are also significantly negative (Table 2). Wen-goldfish and Egg-goldfish both have many derived morphological features relative to the Grass-goldfish, such as the egg-shaped body, the possession of double tailfins, the absence of dorsal fins, and/or dragon-eyes [4,6–8]. These findings correspond to anthropogenic selection to eliminate aesthetically unappealing goldfish and the consequential dramatic reduction in genetic diversity, which occurs in Wen- and Egg-goldfish in the three-breed system.

Three-breed Taxonomy and the History of Domestication

Given the absence of a recorded history of the domestication of goldfish, we employed phylogenetic methods to reconstruct the past. Our analyses reveal that the three-breed taxonomy—Grass-goldfish, Egg-goldfish, and Wen-goldfish—better indicates the history of domestication than either the four-breed or the five-breed systems that emphasize eye-condition. The results of hypothesis-tests indicate that the condition of the fins is informative for grouping goldfish with respect to their evolutionary history. In the three-breed taxonomy, the Grass-goldfish and native Carassius differ only in the color of their scales [4,6–8], and the condition of dorsal fin (loss or retained) distinguishes the Egg-breed from the Wen-breed [1]. Biomechanically, the dorsal fin functions to maintain balance when swimming. Without the dorsal fin, most fishes cannot stay upright. Dorsal finlessness appears after the attainment of double tailfins, which compensate for the loss of dorsal fins [55]. Therefore, Egg-goldfish (no dorsal fin) likely have a more recent origin than goldfish with double tailfins. The further examination of other genes closely related to the morphological characteristics of goldfish can test this prediction.

Supporting Information

Figure S1 Four morphological constraint-trees and the best unconstrained matrilineal genealogy for goldfish. Unique haplotypes identified for each morphological characteristic were constrained to being monophyletic based on the concatenated Cytb and CR data, and using Carassius carassius as the outgroup taxon. The best unconstrained tree was shown at the bottom of the figure. Photographs of the goldfish for each morphological characteristic were mapped to the genealogy. (TIF)

Acknowledgments

We are very grateful to Mu-yong Lee and Yun-Fa Ge for their help in sample collection for this study. We thank Rui Du for his help in the laboratory. We also thank anonymous reviewers for their helpful suggestions.

Author Contributions

Conceived and designed the experiments: YPZ YG SYW. Performed the experiments: SYW. Analyzed the data: SYW YG RWM. Contributed reagents/materials/analysis tools: SYW YG JL RWM SFW CLZ. Wrote the paper: SYW RWM.

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