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Whole genome sequencing revealed genetic diversity and selection of Guangxi indigenous chickens --Manuscript Draft--

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Whole genome sequencing revealed genetic diversity

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and selection of Guangxi indigenous chickens

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14 Abstract

Guangxi chickens play an important role in promoting the high-quality 15 development of the broiler industry in China, but their value and potential are yet to be 16 17 discovered. To determine the genetic diversity and population structure of Guangxi 18 indigenous chicken, we analyzed the whole genomes of 185 chicken from 8 phenotypically and geographically representative Guangxi chicken breeds, together 19 20 with 12 RJFt, 12 BRA and 12 WL genomes available from previous studies. Calculation 21 of heterozygosity (Hp), nucleotide diversity (π), and LD level indicated that Guangxi 22 populations were characterized by higher genetic diversity and lower differentiation 23 than RJFt and commercial breeds except HGFC. Population structure analysis also confirmed the introgression from commercial broiler breeds. Each population clustered 24 together while the overall differentiation was small, MA has the richest genetic diversity 25 26 among all varieties. Selective sweep analysis revealed BCO2, EDN3 and other 27 candidate genes have received strong selection in local breeds, these also provided novel breeding visual and data basis for future breeding. 28

29 Introduction

Chickens are the most widely distributed livestock species globally, more than half the total (53%) is found in Asia, one of the largest producers is China ^[1]. In China, poultry meat consumption accounts for the second largest proportion after pork. People in different regions have different preferences for the appearance, flavor, and cooking methods of chickens. Indigenous chickens with delicious meat quality and unique flavors are more in line with the habits and preferences of domestic consumers.

Guangxi Zhuang Autonomous Region is in mountainous terrain in the far south of 36 37 China, and its unique climate has created unique and rich chicken germplasm resources. 38 Six breeds are listed in Chinese national catalogue of livestock and poultry genetic 39 resources includes Guangxi Three-yellow chicken (SHC), Guangxima chicken (MA), Nandanyao chicken (NDYC), Xiayan chicken (XYC), Guangxiwu chicken (WC), 40 Longshengfeng chicken (LSFC), and two characteristic populations had a long history 41 of local breeding are Dongzhongai chicken (DZAC) and Cenxi Gudian chicken (GDC). 42 These germplasm resources provide good materials for the cultivation of high-quality 43 chicken breeds and promote the development of poultry industry in Guangxi. However, 44 only sporadic studies were reported on genomic information about them ^[2, 3]. 45

46 A comprehensive and deep understanding of the genome diversity of the

indigenous breeds could reveal the genetic diversity and population structure of these
breeds. This study therefore investigated genetic diversity, population structure, linkage
disequilibrium (LD), and signature selection within Guangxi indigenous chickens using
genome-wide single nucleotide polymorphisms (SNPs) generated from the whole
genome sequencing.

52 **Ethics statement**

This study was carried out in accordance with the guidelines of the Animal
Experimental Ethical Inspection Form of Guangxi Research Institute (20190318).

55 Materials and methods

56 Sampling and genotyping

A total of 185 blood samples from six breeds and two characteristic populations
were investigated from conservation center or breeding farms (S1 Table and S1 Fig)
were collected for genomic DNA extracting. Whole-genome sequencing libraries were
constructed by MGIEasy Universal DNA Library Prep Set and then sequenced using
MGISEQ-2000 with PE100 developed by BGI Genomics Co., Ltd.

62 Variant calling and annotation

After quality control, Pair-end reads were mapped onto the Gallus gallus GRC 63 using BWA version 0.7.12-r1039^[4]. The bam files were sorted using SortSam and 64 duplicated reads were marked using MarkDuplicates from Picard tools version 1.105. 65 SNPs were detected and filtered using HaplotypeCaller and VariantFiltration command 66 in GATK version 4.1.1.0. We applied hard filter command 'VariantFiltration' to exclude 67 potential false-positive variant calls with the parameter "QD < 2.0 || ReadPosRankSum 68 69 < -8.0 || FS > 60.0 || MQ < 40.0 || SOR > 3.0 || MQRankSum < -12.5 || QUAL < 30". To annotate the SNPs and InDels identified here, Vep (v95.3) was employed 70

71 Genomic diversity analysis

Genome-wide nucleotide diversity (π) and genetic differentiation (Fst) was performed using VCFtools (v0.1.13)^[5] with parameters 40kb sliding window and 20kb step size. Individual heterozygosity (Hp) was calculated by following the formula given by Rubin et al.^[6]:

76

$$Hp = \frac{2\sum n_{MAJ}\sum n_{MIN}}{(\sum n_{MAJ} + \sum n_{MIN})^2}$$

PopLDdecay version $3.40^{[7]}$ was used to assess patterns in the extent of linkage disequilibrium. ROH analysis was performed using plink version 1.9, the parameters were as follows, the minimum length of ROH > 10kb, the number of SNP per window > 20, and only one heterozygote was allowed, ROH had at least one variant per 1000 kb on average. The results of ROH analysis were visualized with the R package pheatmap, the total ROHs length of each chromosome was centered and scaled in breed's level.

83 Population structure analysis

To investigate the genetic background of the chickens, principal component 84 analysis (PCA) and structure analysis were conducted. SNPs in high linkage 85 disequilibrium were removed by PLINK version 1.9^[8], based on the pruned SNP data, 86 the individual ancestries were estimated using a maximum likelihood method 87 implemented in ADMIXTURE version $1.23^{[9]}$. Using the default parameter (folds = 5) 88 for cross-validation and the lowest cross- validation error was taken as the most 89 probable K value. VCF2Dis (v1.09) software was used to calculate the P distance 90 matrix, then use PHYLIPNEW (v3.69) to constructed a NJ-tree. 91

92 Sweep analysis, GO enrichment and KEGG pathway analysis

The selective sweep analysis was performed using vcftools (v0.1.13) in Guangxi 93 indigenous chickens. Scanning the whole genome selection signal with 40kb as the 94 95 sliding window and 20kb as the step size, and windows with less than 10 SNPs are excluded ^[10]. Fst values were Z-transformed: $ZFst = \frac{(Fst - uFst)}{\sigma Fst}$, where μ is the mean 96 of Fst and σ is the standard deviation of Fst. The sliding window with the top 1% of the 97 ZFst value were defined as significant selected region^[11]. GO enrichment analysis and 98 KEGG pathway analysis were performed using the packages clusterProfiler, KEGG.db 99 and org.Gg.eg.db in R. We select Benjamini-Hochberg method correction for multiple 100 comparisons, and GO terms with a p-value less than 0.05 were considered significantly 101 102 enriched.

103 **Results**

104 Variant calling and annotation

105 A total of 9.48 billion clean reads were obtained after quality filtering, corresponding to average depth and coverage is 9.39x and 96.97% (S3 Table). The 106 overall mapping rate is greater than 98.4%. SNPs with MAF < 0.05, call rate < 0.8 and 107 108 individual call rate < 0.9 were excluded, 13,245,769 high-quality SNPs and 3,790,305 109 indels were utilized for downstream analysis (Fig 1A). The Guangxi indigenous chickens harbored higher number of SNPs and indels than that of RJFt except for 110 DZAC, while WL exhibited the lowest ones. Vep was used to annotate SNPs, 55.66% 111 of these SNPs were aligned to the intron region, 28.43% were aligned to the intergenic 112 region, and only 2.27% located in the exon region (Fig 1B). 113

The genome is divided into isochrones with a sliding window of 100kb, and divided into five categories (L1, L2, H1, H2 and H3) according to different GC level to explore the potential impact between GC content and genetic variations (Fig 1C and 1D)^[12]. Our results shown that the L2 category has the largest number of isochrones, covering 37% of the genomic region, and the SNPs and Indels counts peak in this category. H1 category with higher GC level also contains a lot of genetic variations (S5 Table). In general, genomic regions with moderate GC content contain more variation.

121

Fig1. The number, distribution, and GC content of SNP and INDEL. (A) The SNP
and indel number in different breeds. (B) The distribution of SNPs. (C) Scatter plot of
SNP number and GC content in isochrones per 100kb window. (D) Scatter plot of Indel
number and GC content in isochrones per 100kb window.

126 Genetic diversity, LD and ROH analysis

To provide a more comprehensive understanding and profound insight into the 127 genome diversity of Guangxi indigenous chickens, we incorporated the sequencing data 128 129 of 12 Red jungle fowl population from Thailand (RJFt) and commercial breeds including 12 white layer (WL) and 12 Broiler A(BRA), which has been previously 130 published ^[13]. The nucleotide diversity (π) and heterozygosity were calculated to 131 132 evaluate the genetic diversity of all the chicken breeds. We observed Guangxi indigenous chickens harbored the higher genome-wide π than RJFt (π =0.00334) except 133 for DZAC ($\pi = 0.00332$), the lowest genome-wide π in WL ($\pi = 0.00152$), followed by 134 135 BRA ($\pi = 0.0031$) (Fig 2A). Unlike the results in nucleotide diversity, BRA harbored 136 the highest heterozygous SNP rate (He), followed by DZAC, while MA harbored the lowest (He = 0.1684) one (Fig 2B). The He in Z chromosome are lower than any 137 138 autosome among all populations (S2 Fig), probably due to sex chromosome had

undergone higher selective pressure than autosome ^[14]. Linkage disequilibrium (LD) 139 analysis showed that WL population had the slowest LD decay rate, significantly slower 140 than the followed BRA. MA had a faster LD decay rate than other chicken breeds, 141 142 DZAC and WC have similar LD level with RJFt in second group (Fig 2C).

The level of **ROH** reflects the recent inbreeding history of a population ^[15]. As 143 shown in Fig 2D, the average and total length of ROH of indigenous chicken are short, 144 145 RJFt had the lowest number of ROH. The difference of mROH among Guangxi breeds is relatively small (ranging from 37.74 kb to 66.78kb), which is very different from the 146 largest value of 167.3 kb in WL (S6 Table). HGFC (347.039Mb) had the same tROH 147 148 level with BRA (338.084 Mb), and the shortest tROH was observed in LYWC (185.483 149 Mb). 150

Fig2. Genome diversity and LD decay of 11 chicken populations. (A) Genome 151

152 nucleotide diversity was calculated with a window size of 40 kb and a step size of 20 kb. 153 (B) Genome heterozygosity within 500kb sliding window across the genome. (C) Linkage disequilibrium (LD) decay, denoted with one line for each population. (D) The 154 155 ROH of each chromosome in different breeds. The redder color represents longer ROH, 156 the bluer the shorter.

Population genetics analysis 157

158 As expected, the chickens from the same breeds clustered together according to 159 the PCA. The PC1 (26.79% variances explained totally) could separate the commercial 160 layer breed WL from other populations and PC2 (8.86% variances explained totally) displayed the genetic differentiation between commercial broiler breed BRA and other 161 populations except HGFC (Fig 3A). The 12 RJFt gathered with Guangxi fowls and 162 163 away from commercial chickens. When WL, BRA, RJFt and HGFC were removed from the dataset, DZAC and GDC could be identified as separate clusters, the two 164 populations of LSFC, MC and LSFC, are distributed separately. DZAC and LSFC have 165 a large variation within breed. MA, SHC and XYC are geographically close and tend 166 to get closer to each other. 167

PCA results could not completely reproduce the phylogenetic relationships, and 168 the neighbor-joining tree corroborates the findings of the PCA (Fig 3B). Individuals 169 from the same breed gathered were consistent with the breeding history and 170 geographical distribution. Cluster 1 consists of breeds of outgroup background (WL, 171

BRA and RJFt), and the hybrid HGFC also was grouped together. In cluster 2, WC, 172

NDYC and LSFC are grouped together. Following this group, the local populations 173 174 DZAC, GDC and several MA were arranged in the middle of the tree, but were not forming a visually distinct cluster. Cluster 4 consists exclusively of chickens sampled 175 in MA. Cluster 5 branched into two sub-clusters with SHC and XYC, an individual 176 from the WC was grouped with XYC, possibly because of a sampling error. 177 178

179 Fig3. Population structure analyses. (A) Principal component analysis (PCA), with 180 26.79% and 8.86% variance explained in PC1 and PC2, respectively. (B) Neighborjoining tree of 221 chickens, constructed with PHYLIPNEW version 3.69.650 (C) 181 Admixture analysis with K values running from 3 to 6. 182

183 To assess historical admixture patterns of the chickens, we conducted the ADMIXTURE analysis with K values running from 3 to 18. At K=3, genetic 184 divergency first occurred between commercial breeds and non-commercial ones. 185 HGFC shared the same ancestral lineage with BRA , Guangxi indigenous breeds shared 186

187 the same ancestral lineage with RJFt, WL consistent with the above PCA and phylogenetic tree result (Fig 3C). When K=4, the Guangxi indigenous breeds were 188 separated from others (except HGFC). Indigenous chickens gradually separated from 189 190 each other when K ranged from 5 to 14. MA experienced introgression from SHC, and the ancestral components of SHC, GDC and HGFC are pure. There is differentiation in 191 the breeding programs of LYWC and DLWC, as well as in LSFC and XYC breeds (S3 192 193 Fig). According to the calculated cross-validation value, the best fit was k=5, Guangxi indigenous breeds showed two ancestral components that are different from others. 194

195 Selective Sweep Analysis

The skin color influences consumers' preferences, and the yellow skin chickens 196 are more popular than white ones in south of China. We observed the highest ZFst 197 region occurring at chr24: 6.14-6.18 Mb (Fst = 0.63, ZFst = 35.39) between 121 vellow 198 199 skin and 64 non-vellow skin chickens. The top ten selected window annotated eight 200 genes, among which BCO2 gene is a classical yellow color gene in chicken. Then we collected the genotypes on this gene and found that the non-yellow skin clusters showed 201 202 a different genotypic pattern from LSFC and NDYC; as for WC, it has both two patterns 203 might due to the concealment of its black skin that the yellow skin has not been 204 deliberately selected (S4 Fig). We found a missense mutation at chr24: 6155481T>C (rs313409504) was consisted with the previous report^[16]. The strongest selective sweep 205 206 on chromosome 11 we detected was located at 19.12-19.16 Mb (Fst = 0.17, ZFst = 9.41) 207 near the gene MC1R, which plays a key role in controlling the deposition of melanin.

208 209

Fig4. The result of ZFst and Log2(pi) of yellow skin.

210 In oriental countries, nutritional and medicinal benefits have been attributed to the consumption of black-boned chickens. WC is distinguished compared with other breeds 211 212 because of its black beak, crown, skin, and shank. We compared WC with other Guangxi chickens and scanned the whole genome for selected regions. The strong 213 214 selected region contained four annotated functional genes which were associated with 215 dermal hyperpigmentation in chickens, beta-1 tubulin (TUBB1) and PRELI domain containing 3B (PRELID3B, also known as SLMO2), GNAS complex locus (GNAS), 216 217 encoding endothelin 3 (EDN3). GO enrichment analysis shows that gene TYRP1 on 218 chromosome Z and KITLG on chromosome 1 were significantly enriched in melanocyte differentiation (GO:0030318) and developmental pigmentation (GO:0048066). SYK 219 gene plays a role in regulation of bone resorption (GO:0045124). 220

221 XYC and GDC have a similar appearance with SHC, XYC is fat-deposited and GDC is smaller. We compared them with the SHC and scanned the whole genome for 222 223 selected regions related to fat deposition and body size. We found that the selected 224 genes of fat deposits were located on chromosome 12 and chromosome Z. HMGCS1 225 and OXCT1 are significantly enriched in the pathway of ketone body synthesis and 226 degradation (gga00072), ketone bodies are produced in the liver, mainly from the oxidation of fatty acids, and are exported to peripheral tissues for use as an energy 227 228 source. ATG7 (Autophagy Related 7) is a protein coding gene (gga:04140), it has been 229 associated with multiple functions, including axon membrane trafficking, axonal homeostasis, mitophagy, adipose differentiation, and hematopoietic stem cell 230 231 maintenance. A significant signal peak was detected at 49.32-49.36Mb on chromosome 5 (ZFst=8.5, Fst = 0.51) located in the *DLK1-DIO3* genomic region, which was 232 recognized to be an imprinted domain in placental mammals associated with 233 developmental programming^[17]. 234

235 **Discussion**

We assessed the genetic diversity of indigenous chickens from Guangxi provinces
using the re-sequencing, meanwhile, we performed selective sweep analysis of
phenotypes related to economic traits.

As shown in our study, most Guangxi chicken breed have various genetic diversity on the genome indicated by higher SNPs abundance, slower LD decay rate, lower ROH values and higher heterozygosity. HGFC, as a population of LSFC breed, is an exception, gene introgression from commercial broilers was detected. After further investigation, in order to improve the economic benefit of this group, broilers genes were introduced artificially in the process of breeding. The results of genetic structure are consistent with the origin of breeds, indicating the effectiveness of Guangxi local

chicken population in breed protection , even the HGFC also formed its own characteristics because the ancestral composition of it is different from that of BRA when k=6. Compared with commercial breeds, the genetic difference among indigenous chickens is relatively small, and Guangxi indigenous fowls do have a closer relationship with RJFt ^[18].

The difference of deposition location and amount of carotenoid and melanin in 251 252 chicken skin led to the diversity of chicken skin color. BCO2 gene encodes beta-253 carotene dioxygenase 2 could cleave colorful carotenoids to colorless apocarotenoids 254 by an asymmetric cleavage reaction^[19], is established as the causal gene for the yellow skin. According to the results of selection scanning, BCO2 gene is extremely strongly 255 256 selected in the population, and the SNP shows different patterns in the yellow and nonyellow skin population. Eriksson et al. demonstrate that regulatory mutations that 257 inhibit expression of BCDO2 in skin caused yellow skin, but not in other tissues ^[16]. 258 259 Fallahshahroudi's study showed the down-regulation of BCO2 in skin, muscle, and adipose tissue was associated with the derived haplotype ^[20]. Also, BCO2 has variety 260 variants in different breeds. Wang found a G>A mutation in exon 6 to be associated 261 with the concentration of carotenoids in Guangxi-huang and Qingjiao-ma chicken^[21]. 262 A GAG haplotype was fixed in commercial breeds of yellow skin^[16]. We also found 263 the missense mutant at chr24:6155481 led to the mutation of threonine to alanine. 264

265 The strongest selective sweep region with dermal hyperpigmentation on 266 chromosome 20 was located at 10.64-10.94 Mb consists of a set of 7 genes. EDN3 is a 267 gene with a known role in promoting melanoblast proliferation by encoding a potent mitogen for melanoblasts/melanocytes. Shinomiya et al. reported that the 268 overexpression of genes in a 130kb duplication region gives rise to the 269 hyperpigmentation in silk chickens ^[22], and then Dorshorst et al. extended this 270 discovery and strongly suggested that the increase of EDN3 expression caused by 271 272 duplication is the cause of FM in all breeds of chickens. The expression of two other 273 genes, SLMO2, and TUBB1 were also significantly increased in expression in both skin and muscle tissue from adult fibromelanosis chickens might contribute to the dermal 274 hyperpigmentation phenotype^[23]. Analysis of RNA-seq suggested that *SLMO2*, *ATP5e*, 275 and EDN3 were differentially expressed between the black and yellow skin groups, 276 277 combined analysis of genomic data found that EDN3 might interact with the upstream ncRNA LOC101747896 to generate black skin color during melanogenesis ^[24]. Wang 278 et al. study indicated that a T2270C mutation in GNAS gene promoter in chicken is 279 correlated strongly with the skin color traits^[25]. 280

DLK1 has been reported to be associated with the development and differentiation
 of adipose and muscle in chicken and *DLK1* promotes muscle development inhibitory

adipogenesis in mammals^[26]. Zhang et al.'s study provides strong in vivo evidence that 283 atg7, and by inference autophagy, is critical for normal adipogenesis ^[27]. AMACR 284 coding protein alpha-methylacyl-CoA racemase, this protein is involved in the pathway 285 286 bile acid biosynthesis, which is part of Lipid metabolism (gga00120). Bile acid is the main component of bile and its main function is to promote the digestion and absorption 287 of fat. HSD17B4 codes a bifunctional enzyme mediating dehydrogenation and 288 289 anhydration during β-oxidation of long-chain fatty acids, and a non-synonymous SNP has been reported to be related to meat-quality traits in pig ^[28]. PRKAA1 is associated 290 with skeletal muscle lipid accumulation^[29]. 291

292 **Conclusions**

In conclusion, a comprehensive analysis of genetic diversity, population genetic structure, LD, and the selection signatures of 8 indigenous chicken breeds distributed in Guangxi. The results suggested that indigenous chickens have abundant genetic diversity and potential, candidate genes related to economic traits can also provide a theoretical basis for breeding. Our analyses provide data for further research and local breeding of Guangxi indigenous chicken.

299 Data availability statement

The raw reads data have been submitted to NCBI Sequence Read Archive database
with the accession number PRJNA659069. The data reported in this study are also

302 available in the CNGB Nucleotide Sequence Archive (CNSA: https://db.cngb.org/cnsa ;

accession number CNP0001716). Furthermore, 36 individuals of commercial breeds
and red jungle fowls were downloaded from NCBI at ERP112703 (S2 Table).

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389 **Supporting information**

- 390 S1 Fig. Geographic distribution and appearances of typical female chickens.
- 391 S2 Fig. Boxplot showing heterozygous SNP rate of autosomes (left) and Z

- 392 chromosome (right) between each chicken population.
- 393 S3 Fig. Admixture analysis with K values running from 7 to 18. Each population
 394 separated by white dotted line.
- 395 S4 Fig. The genotype of fixed SNPs in chr24: 6.14Mb~6.18Mb of individuals. The
- 396 row represents the SNP position and the column represents the individual. Light blue
- denotes reference alleles while red indicates alternative homozygous alleles, yellowmeans heterozygous and dark blue means missing.
- 399 S5 Fig. ZFst values and Log 2 (pi). (A) WC and other indigenous population. (B)
- 400 XYC and SHC. (C) GDC and SHC
- 401 S1 Table. The character of chickens in this study.
- 402 S2 Table. The public data information.
- 403 S3 Table. The sequencing information of samples.
- 404 S4 Table. The distribution of variant, nucleotide diversity and HE.
- 405 **S5 Table. The distribution of isochrones.**
- 406 **S6 Table. ROH of chicken breeds.**
- 407 S7 Table. Selective sweep of dermal hyperpigmentation.
- 408 S8 Table. Selective sweep of yellow skin.
- 409 S9 Table. Selective sweep of body size.
- 410 S10 Table. Selective sweep of fat deposition.
- 411 S11 Table. Functional gene categories enriched for genes under selection.
- 412 S12 Table. KEGG pathway analysis of genes.









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