Genetic Dissection of Growth Traits in a Unique Chicken Advanced Intercross Line

Yuzhe Wang1,2*, Lina Bu2†, Xuemin Cao2, Hao Qu3, Chunyuan Zhang2, Jianguo Ren2, Zhuolin Huang2, Yiqiang Zhao2, Chenglong Luo3*, Xiaoxiang Hu2*, Dingming Shu3 and Ning Li2

1 College of Animal Science and Technology, China Agricultural University, Beijing, China, 2 State Key Laboratory of Agrobiotechnology, College of Biological Sciences, China Agricultural University, Beijing, China, 3 State Key Laboratory of Livestock and Poultry Breeding, Guangdong Key Laboratory of Animal Breeding and Nutrition, Institute of Animal Science, Guangdong Academy of Agricultural Sciences, Guangzhou, China

The advanced intercross line (AIL) that is created by successive generations of pseudo-random mating after the F2 generation is a valuable resource, especially in agricultural livestock and poultry species, because it improves the precision of quantitative trait loci (QTL) mapping compared with traditional association populations by introducing more recombination events. The growth traits of broilers have significant economic value in the chicken industry, and many QTLs affecting growth traits have been identified, especially on chromosomes 1, 4, and 27, albeit with large confidence intervals that potentially contain dozens of genes. To promote a better understanding of the underlying genetic architecture of growth trait differences, specifically body weight and bone development, in this study, we report a nine-generation AIL derived from two divergent outbred lines: High Quality chicken Line A (HQLA) and Huiyang Bearded (HB) chicken. We evaluate the genetic architecture of the F0, F2, F8, and F9 generations of AIL and demonstrate that the population of the F9 generation sufficiently randomized the founder genomes and has the characteristics of rapid linkage disequilibrium decay, limited allele frequency decline, and abundant nucleotide diversity. This AIL yielded a much narrower QTL than the F2 generations, especially the QTL on chromosome 27, which was reduced to 120 Kb. An ancestral haplotype association analysis showed that most of the dominant haplotypes are inherited from HQLA but with fluctuation of the effects between them. We highlight the important role of four candidate genes (PHOSPHO1, IGF2BP1, ZNF652, and GIP) in bone growth. We also retrieved a missing QTL from AIL on chromosome 4 by identifying the founder selection signatures, which are explained by the loss of association power that results from rare alleles. Our study provides a reasonable resource for detecting quantitative trait genes and tracking ancestor history and will facilitate our understanding of the genetic mechanisms underlying chicken bone growth.

Keywords: chicken, advanced intercross line, bone growth, ancestral inference, QTL fine-mapping, genome-wide association study, selective sweep, haplotype association study
INTRODUCTION

Identifying key polymorphisms and dissecting the genetic architecture of complex growth traits is of considerable interest in fields like agriculture breeding and evolution. F\textsubscript{2} crosses between divergent outbred lines are traditionally used to map quantitative trait loci (QTL) in domestic animal and plant populations (Andersson et al., 1994; Perez-Enciso et al., 2001). However, it is not enough to rely on a single generation of meiotic recombination to break up and randomize the parental genomes to finely map causal variants of complex quantitative traits (Flint et al., 2005). Improved strategies, such as the use of larger sample cohorts, the construction of advanced intercross lines (AIL) (Besnier et al., 2011; Parker et al., 2012), nested association mapping population (NAM), and multi-parent advanced generation inter-cross (MAGIC) in animals and plants (Poland et al., 2011; Gatti et al., 2014; Pascual et al., 2015) can increase the precision of quantitative trait loci (QTL) mapping by introducing more recombination events and together provide a series of alternatives to the traditional association mapping of populations.

Advanced intercross lines (AILs) were first introduced by Darvasi and Soller (1995). An AIL is created by successive generations of pseudo-random mating after the F\textsubscript{2} generation, and recombinations are accumulated continuously between generations and are easier to construct in species with short generation intervals and a high tolerance of inbreeding decline. To date, AIL has been used as a common strategy to improve the mapping resolution for the genome wide association studies (GWASs) of model animals, such as fruit flies (Mackay et al., 2012), mice (Gonzales et al., 2018), chicken (Zan et al., 2017), and C. elegans (Doitsidou et al., 2016). The significant advantages of AILs include reducing the QTL confidence interval by 3- to 27-fold and finely splitting the original QTL into two linked QTLs (Besnier et al., 2011; Parker et al., 2014; Arends et al., 2016). However, we should always pay attention to the tradeoff between mapping resolution and statistical power, as the causal allele may become rare with a continuous increase of the inbreeding coefficient in the AIL (Yalcin et al., 2010; Parker et al., 2016).

Bone growth is crucial to poultry production, as skeletal problems are associated with economic benefits and animal welfare issues (Tszudzuki et al., 2007; Kapell et al., 2012). Too long legs give rise to leg problems in high body weight chickens (Deeb and Lamont, 2002). In healthy chicken, shank length (SL) and shank circumference (SC) are the two most commonly used parameters for evaluating bone growth in chickens (Tszudzuki et al., 2007) and are highly correlated with body weight (BW) (Gao et al., 2010). Moreover, shank traits can be measured without slaughter and we can track bone growth of different periods. We previously reported two major QTLs for growth traits located on chicken chromosome 1 (GGA1) and GGA27 via a linkage analysis in the F\textsubscript{2} generation (Sheng et al., 2013). To promote a better understanding of the underlying genetic architecture for growth trait differences, specifically body weight and bone development, here, we report a nine-generation AIL derived from High Quality chicken Line A (HQLA) × Huiyang Bearded (HB) chicken. Detailed information on HQLA, HB, and AIL is presented in the materials and methods and Figure 1. We employed genotyping-by-sequencing (GBS) SNPs from F\textsubscript{0}, F\textsubscript{8}, and F\textsubscript{9} (Wang et al., 2017) and Beadchip SNPs from the F\textsubscript{2} of AIL (Sheng et al., 2013). Based on these data, we characterized the gradient of the population structure over these generations and the potential functional genes of growth traits by a genome wide association study (GWAS), selective sweep analysis, haplotype association, and ancestral inference. The integrated analysis of selection in F\textsubscript{0} and GWAS for AIL provides both power and precision and demonstrates the transmission of important genetic information between generations.

MATERIALS AND METHODS

The AIL Population

The High Quality chicken Line A (HQLA) is a closed population founded by the commercial Anak broiler breed and a Chinese indigenous chicken line, followed by strong artificial selection over more than 10 generations, according to a weight-based selection index, while maintaining the meat quality and plumage color. The Huiyang Bearded chicken (HB) is a Chinese meat-type breed with a long history (1,600 years), which is characterized by its slow growth, high meat quality, and muff and beard phenotype (Guo et al., 2016); currently, HB is in the stage of conservation and breeding. At 7 weeks of age, the HQLA was 3.2-times the body weight of HB (Figure 1). The F\textsubscript{2} cross was generated by the reciprocal crossing of the founder lines [4 HQLA♂ × 12 HB♀ and 4 HB♂ × 12 HQLA♀, details presented in Sheng et al. (2013)]. Later AIL generations (F\textsubscript{3} to F\textsubscript{9}) were founded by birds from the F\textsubscript{2} population and bred using random mating (Figure 1). The population size of each generation was maintained at more than 1,000 individuals. The body weight at 7 weeks of age (BW\textsubscript{7}) was around 900 g.

Phenotype

For the F\textsubscript{9} generation, body weight was measured at hatching and every other week until 12 weeks of age. During weeks 4–12, the shank length and shank circumference were also measured every 2 weeks. Boxplots for each phenotype were generated to scan for outliers. Individuals that were further than 1.5 times IQR away from the upper or lower quartile of the boxplots were removed. Descriptive statistics of the phenotypes are provided in Supplementary Table S1.

Genotype

We employed GBS SNPs of F\textsubscript{0}, F\textsubscript{8}, and F\textsubscript{9} (Wang et al., 2017) and Illumina Chicken 60K Beadchip SNPs of F\textsubscript{2} (Sheng et al., 2013) for further filtering and analysis. In brief, for F\textsubscript{9} generation, double-enzyme GBS (EcoRI/MseI) libraries were prepared and sequencing was performed on an Illumina Nextseq500 sequencer. The TASSEL-4.0 GBS analysis pipeline was used to discover SNPs. Using VCFtools (0.1.17), the raw GBS SNP filter criteria was set to: -maf 0.05 – max-alleles 2 – min-alleles 2 – minDP 5 – minGQ 98 – max-missing 0.2 (Danecek et al., 2011). Genotype phasing of the clean SNPs was done using Beagle 5.0 (Browning and Browning, 2007) with gt model and impute = true parameters,
other parameters were left as default. The GBS SNPs were evenly distributed across chromosomes (Supplementary Figure S1). In F2 generation, SNPs (autosome 1–28) that failed to meet the following criteria were removed: individual call rate (>0.9), individual SNP call frequency (>0.9), and minor allele frequency (MAF >0.05). All the genomic coordinates of the SNPs were uniformly converted to the chicken reference genome Gallus gallus, version 5.0 (Ensembl release 94). After that, we kept 161,376 GBS SNPs for 16 HQLA, 14 HB, 185 F8, 602 F9 individuals, and 43,472 Chip SNPs for 492 F2 individuals.

Genetic Parameter Estimation

We evaluated the changes in the population genetic parameters as a component of generation transmission. LD decay statistics were analyzed by PopLDDecay 3.31 (Zhang et al., 2019) with a max distance of 2 Mb. The inbreeding coefficient (F), nucleotide diversity (π), nucleotide divergent, and minor allele frequency (MAF) were evaluated by VCFtools (0.1.17) (Danecek et al., 2011). The heritability and genetic correlations of all traits were estimated using GCTA package (v1.92) (Yang et al., 2011).

Genome Wide Association Study

The mixed linear model (MLM) approach was used for the GWAS of the F9 generation, as implemented in the GCTA package (v1.92) (Yang et al., 2011). The basic model was: $y = a + bx + g + e$, where $y$ is the phenotype, $a$ is the mean term, $b$ is the additive effect (fixed effect) of the candidate SNP to be tested for association, $x$ is the SNP genotype indicator variable, $g$ is the polygenic effect (as captured by the GRM calculated using all SNPs), and $e$ is the residual. The GWAS statistical model of body weight included the sex and batch as discrete covariates and hatch weight as a quantitative covariate. For shank traits, body weight at the same age were also included as a covariate, because we focused on QTL scans which are associated with bone growth. A quantile-quantile (Q-Q) plot generated in CMplot1 was used to assess the potential impact of population stratification (Supplementary Figure S2). Bonferroni correction was applied to correct the number of estimated independent markers. A subset of SNPs that were in approximate linkage equilibrium was obtained by removing one in each pair of SNPs if the LD was greater than 0.2 using PLINK v1.07 (Purcell et al., 2007). QTL intervals were established after the remaining top SNPs and their neighboring SNPs with $r^2 >0.3$.

Selective Sweep

To investigate the signatures of selection between HQLA and HB, four statistical tests were used, including XP-EHH and iHH (linkage disequilibrium-based), Tajima’s D (frequency spectrum-based), and $F_{st}$ (population differentiation-based), to investigate the signatures of selection between HQLA and HB. The XP-EHH and iHH value at each locus were estimated by The selfscan program (v1.2.0a) (Szpiech and Hernandez, 2014), and the genetic map for our population was 3 cM/Mb. The statistics for $F_{st}$ and Tajima’s D were calculated using VCFtools (0.1.17) (Danecek et al., 2011) with a window size of 200 Kb and step size of 100 Kb.

Haplotype-Based Association Analyses

A haplotype-based association analysis was performed in the ~120 Kb fine-mapped QTL region on GGA27 using the following model: $Y = X\beta + Zu + e$, where $Y$ is a column vector containing the SL10 of the F9 individuals. $X$ is the design matrix including the coding for the sex of the birds. For each specific interval, there are $n$ haplotypes for $m$ individuals constructed by several SNPs. $Z$ is the design

---

1https://github.com/YinLiLin/R-CMplot
matrix (m \times n) containing each haplotype count (coded as 0, 1, 2) of each individual. β is a vector that estimates the fixed effect of sex, u is a column vector that estimates the allele substitution effects for each haplotype, and e is the normally distributed residual.

**Ancestral Inference**

The RFmix software (v2.03) (Maples et al., 2013) is based on a discrimination analysis model that can be used to estimate the genetic ancestry composition of each individual and each chromosome. Using the F0 population as the ancestor population, RFmix was used to evaluate the local ancestral source of individuals in the F9 generation. To determine the haplotype window size, we set conditional random field spacing (# of SNPs) (-c) to 9 based on the results of LD with window size, we set conditional random field spacing (# of SNPs) of 2 Kb in F9, 8 Kb in F10, 16 Kb in F11, and fully mixed in the F12 generation. The recombination and continued to accumulate as the distance increased. The r2 decayed rapidly in F8 and F9 individuals in comparison to F2 populations (r20.1 = 27 Kb in F9 and r20.1 = 570 Kb in F2), supporting the suitability of the F9 population for high-resolution mapping.

We used GBS SNPs to estimate the nucleotide polymorphisms (π) in each population (except F2) (Figure 2D) and the inbreeding coefficient (F) (Figure 2E). HQLA showed higher nucleotide polymorphisms but lower heterozygosity levels (higher F value) than HB. This profile is consistent with the strong artificial selection history of the HQLA population. Fortunately, the F9 generation maintained a high nucleotide polymorphism, and only 552 SNPs (0.34%) were lost compared to the F0 GBS data. Considering the distribution of minor allele frequencies (MAFs) (Figure 2F), a high proportion SNPs (30.43%) in F9 had lower allele frequencies (MAF < 0.1) than F0 (22.46%). This pattern shows that the AIL population still experiences a slight genetic drift and bottleneck between F0 and F9.

**GWAS Identified Two Major QTLs Affecting Growth Traits in the F9 Generation**

The growth traits of this study population have high heritability (0.48–0.82, Supplementary Table S2). Using a mixed linear model, we performed GWAS between the 161,376 GBS SNPs and 17 growth traits, including BW2-BV14, SC4-SC12, and SL4-SL12, in 599 F9 individuals. All traits have a high phenotypic and genetic correlation, especially between the same traits at different periods. The correlation between SC and BW at different periods is higher than the correlation between SL and BW (Supplementary Table S3). At a Bonferroni of 5% (1.01 × 10−6, 0.05/49,318), we identified a large major QTL mainly affecting body weight at GGA1: 168.6–171.7 Mb (Q1) and a small major QTL mainly affecting shank development at GGA27: 3.60–3.72 Mb (Q2) (Figures 3A,B and Supplementary Figure S3). These QTL peaks were narrower than those of the F2 linkage analysis (Sheng et al., 2013). The most significant associations were for BW8 at GGA1: 169,241,142 bp (p = 3.8 × 10−16) and SL10 at GGA27: 3,608,297 bp (p = 6.1 × 10−8). Q1’s confidence interval was 25-fold that of Q2, partly because the recombination rate of GGA1 (12.05 cM/Mb) was 4.9-fold that of GGA2 (2.44 cM/Mb) (Sheng et al., 2013) and the LD in GGA1 is more extensive than that in GGA2 (Supplementary Figure S4). The broad loci in Q1 make it difficult to infer which genes are responsible for the association. We speculate that there is more complex genetic architecture concealed in Q1, such as multiple linked minor QTLs. However, clarifying this architecture further is very difficult. The following fine-mapping work mainly focuses on the Q2 interval. It should be noted that the Z chromosome was excluded in this study due to the pre-GWAS of 297 cocks (ZZ) showing no significant signal on the Z chromosome.

**Selective Sweep Analysis on the F0 Generation Retrieved a Missing QTL on GGA4**

The genes or variants underlying the large phenotypic differences between HQLA and HB likely evolved rapidly after artificial selection. Based on this principle, we employed different statistical tests to investigate the signatures of selection, including frequency spectrum-based Tajima’s D, the linkage disequilibrium-based XP-EHH method, and the population differentiation-based Fst method. However, one must carefully evaluate the results of selection signals since small sample sizes may introduce large drift effects. Hence, we combined our GWAS results with the Animal Quantitative Trait Loci Database2 to conduct a further screening of each selection signal interval. By comparing the growth traits associated QTLs with the candidate selection interval obtained by at least one method, we identified a total of 10 clear selection signal intervals (S1–S10), four of which occurred mainly in HQLA and six of which occurred in HB (Figures 3C–E).

Among these, we highlight the narrowed S6 interval on GGA4 (Figures 3C, 4A,B) matched the QTL database’s lists of

2https://www.animalgenome.org/cgi-bin/QTLdb/index
growth traits. This signal spans GGA4: 75.28–75.67 Mb, harbors some candidate genes (PACRGL, SLIT2, KCNIP4, and mir-218-1), and has been reported to be significantly associated with chicken body weight in different populations, such as White Leghorn × Rhode Island Red cross (Sasaki et al., 2004), Silky Fowl × White Plymouth Rock cross (Gu et al., 2011), Beijing-You chickens (Liu et al., 2013), New Hampshire × White Leghorn cross (Nassar et al., 2015), and Dongxiang Blue-shelled chickens × White Leghorn cross (Guo et al., 2020). However, our association results were negative at this location because HQLA and HB were selected in the same direction (nearly fixed in HB, Figure 4C), resulting in an extremely low allele frequency difference (ΔAF) between them, which led to a further loss of statistical power in F9-GWAS.

**Fine-Mapping and Local Ancestral Inference for the Mosaic QTL on GGA27**

A 120 Kb QTL region (GGA27: 3.60–3.72 Mb) was identified by 34 GBS SNPs and aggregated using $r^2 > 0.3$ with the top five SNPs of SL10. The GWAS significant SNPs were not continuously distributed across the region but were instead located in two peaks separated by regions with no genetic hitchhiking (Figure 4D). It is difficult to identify the selected interval by the window-based selection method under large allele frequency fluctuations (Figures 4E,F) (the mosaic association model).

We further analyzed the genetic architecture of this QTL using haplotype association analysis. To identify the haplotypes contributing to the association signal, a multilocus backward-elimination analysis was performed across the 34 SNPs in the 1.2 Mb region, and the top SNP on GGA1:169,241,142 bp was selected to control for Q1 effects. Four SNPs (GGA27: 3,608,297 bp, 3,620,306 bp, 3,644,245 bp, and 3,686,628 bp) were identified to have statistically independent associations with SL10 at a 5% False Discovery Rate (FDR) threshold. The haplotypes tagged by these 4 SNPs were estimated; in total, 12 haplotypes were detected (MAF > 0.01 in F9). Tracing back to the F0 generation, we confirmed three HQLA-origin haplotypes (A, red), four HB-origin haplotypes (B, blue), two shared haplotypes (gray), and three recombination haplotypes (orange) (Figure 5). We first focused on the dominance-recessiveness relationship and computed the phenotype scale for AA, AB, and BB as $1.37 \pm 0.65$ cm ($n = 95$), $0.08 \pm 1.22$ cm ($n = 201$), $-1.53 \pm 0.64$ cm ($n = 151$), respectively. The results showed that HQLA carries the main length increasing alleles, and heterosis does not commonly exist in crosses of AB. Next, the additive haplotype substitution effects on SL10 were estimated. There was a gradient distribution of haplotype allele effects between decreasing SL10 by 1.57 cm and increasing it by 2.38 cm (Figure 5). Although most of the length increasing haplotypes are inherited from HQLA, there is still fluctuation among them, which is the same in HB. This effect distribution does not seem to be caused by only one causal mutation. Some well-known candidate genes related to body size and bone growth are located within this interval and are worthy of follow-up research, including PHOSPHO1, IGF2BP1, ZNF652, and GIP (Figure 4D). However, the proportion of our
FIGURE 3 | Joint analysis of GWAS in F9 and selection signature identification in F0. The Manhattan plots for BW8 (A) and SL10 (B). The genome-wide 5% significance threshold $-\log_{10}P$ was 5.99. (C) XP-EHH and iHH in HQLA and HB using a ±2 cutoff (top 4.4% genomic region). (D) Tajima’s D in HQLA and HB and (E) the $F_{ST}$ value with a 200 Kb window using the 99th percentile cutoff. The orange vertical dashed (marked by the letter Q) represents the QTL interval, and the red and blue vertical dashed (marked by the letter S) represent the selection signature intervals dominated by HQLA and HB, respectively.
recombination haplotypes is too small to give further genetic evidence in the current population. Further recombinations (more offspring in AIL $F_9$) and higher density markers (Davies et al., 2016; Yang et al., 2019) will clarify this issue.

**DISCUSSION**

Crosses among well-characterized strains are a mainstay of modeling organism genetics. We reported a running chicken AIL that was generated by crossing HQLA × HB, which was differentially bred for fast growth and slow growth prior to subsequent intercrossing. Systematic characterization of the genetic architecture of AIL makes it possible to evaluate the suitability of different genomic situations for GWAS. Overall, the $F_9$ of AIL has low linkage disequilibrium between markers to obtain accurate mapping resolution, an absence of population structure to prevent false positive associations, and relatively stable allele frequency to ensure a high enough power to detect the majority of quantitative trait loci (QTLs). We highlighted the fine-mapped QTL on the GGA27 derived from GWAS, haplotype association, and local ancestry inference, which implicated four candidate genes corroborated by extant human, mouse, and chicken genetic data.

Although the basic strategy to build the AIL was similar to that in other studies, certain practical considerations, along with the AIL’s complex genetic background, affected the design of this study and its outcomes in important ways. For example, we observed rich diversity and intense LD decay in the $F_0$ generation, even though the four males were full siblings, and the 12 females were either half or full siblings, in the HQLA and HB founders, which is very different from the inbred AIL line of mice (Gonzales et al., 2018) and other model organisms (Mackay et al., 2012; Nicod et al., 2016). Compared with other chicken AILs, HQLA-HB-AIL presents lower levels of SNP diversity loss (11% SNPs with MAF < 0.05 in $F_9$) than broiler × Fayoumi AIL (60% SNPs with MAF < 0.05 in $F_{18}$ and $F_{19}$) (Van Goor et al., 2015). Moreover, our previous study (Guo et al., 2016) reported that the single-nucleotide genome-wide polymorphisms of $F_0$ were 1.38-fold those of the founders of the Virginia chicken AIL population (Zan et al., 2019), which illustrates the high polymorphisms in our population. Thus, this AIL is more human-like or similar to laboratory outbred mice (Yalcin et al., 2010) than the inbred AIL mice model, namely that this AIL has lower levels of LD, lower MAFs, and is more abundant haplotype diversity; the resulting mosaic association model also supports this conclusion. This is a double-edged sword that improves fine-mapping accuracy but affects power.
Haplotype association analysis for the shank length at 10 weeks of age in the 120 Kb candidate region on GGA27. Four SNPs were associated with SL10 in a multilocus backward-elimination analysis across the segment. Haplotypes were estimated in the founder populations (HB and HQLA) and the F9 AIL generation across these markers. Twelve haplotypes were inferred in F9 at haplotype frequencies (HF) > 0.01, including five unrecombined HQLA haplotypes (red), six unrecombined HB haplotypes (blue), two shared haplotypes (gray), and three recombination haplotypes (orange). Overall, the haplotype substitution effects exhibited a gradual distribution of effects on SL10 in F9.

by increasing the multiple testing burden (Parker et al., 2016). In addition, the diversity of F0 may be due to the breeding process because HQLA is a commercial strain formed first by crossing and then by directed selection. This factor gives this population greater similarities to the three-ancestor MASIC population from the perspective of ancestors, which can be monitored by estimating individual ancestry (Supplementary Figure S5) using the unsupervised ADMIXTURE method (Alexander et al., 2009).

We also presented a joint analysis of GWAS, and selective sweep of this AIL was able to comprehensively extract more genomic features. Firstly, although we cannot rule out the effect of genetic drift on the selection results, the diversity of F0 still reduces the false positive rate of the selection signal to some extent, which allows all candidate intervals to be further studied based on their association with other phenotypes. Secondly, we showed a typical example of failing to replicate prior results on GGA4, which we explained by the loss of GWAS power that results from rare alleles. This result demonstrated that local diversity may be lost, even if two founder strains generally maintain large phenotypic/genotypic differences, which is also a major performance difference between AIL and MAGIC. Besides, this study used ~160 K SNPs, which means that a very large sample size which not available currently is required to meet the multiple testing correction of detecting epistatic QTLs. Therefore, a comprehensive analysis of interaction between directional epistasis and mutation effects will also be a very interesting issue to be explored in the near future.

Another core issue of this study is the dissection of key growth-related (especially for bone development) genes. We focused on the narrow QTL on GGA27 that contains fewer genes, and we highlighted some genes that are noted by the existing literature for their role in the corresponding traits. This result finely replicated the F2 finding (Sheng et al., 2013) and is consistent with the QTL-mapping in Japanese cockfighting (Tsuzuki et al., 2007) and Pekin ducks (Zhou et al., 2018). The lead SNP at chr27:3,608,297 is associated with the shank length in F9 AILs, which lies in the intron of the ZNF652 gene. Although the function of this gene has not been reported in detail, discoveries from human GWAS have replicated the significant correlation between ZNF652 and body height in two independent cohorts [rs35587648, $p = 7 \times 10^{-42}$ in Lango Allen et al. (2010) and rs2072153, $p = 4 \times 10^{-8}$ in Kichaev et al. (2019)]. Interestingly, three other genes at this locus, PHOSPHO1, IGF2BP1, and GIP, have been reported to be related to skeletal development. PHOSPHO1 is a phosphoethanolamine/phosphocholine phosphatase that has been implicated in the generation of Pi for matrix mineralization, a process central to skeletal development. Phospho1−/− mice display growth plate abnormalities, spontaneous fractures, bowed long bones, osteomalacia, and scoliosis in early life. Insulin-like growth factor II mRNA-binding protein 1 (IGF2BP1) belongs to a family of RNA-binding proteins implicated in mRNA localization, turnover, and translational control (Bell et al., 2013). The IGF2BP1−/− mice were, on average, 40% smaller than their wild-type and heterozygous littermates; growth retardation was apparent from E17.5 and remained permanent into adult life (Hansen et al., 2004). Moreover, a GWAS study revealed that a putative regulatory mutation causes the continuous expression of the IGF2BP1 gene after birth, which increases body size of Pekin ducks by 15% (Zhou et al., 2018). Glucose-dependent insulinotropic polypeptide (GIP) also has been recognized in the last decade as an important contributor to bone remodeling and is necessary for optimal bone quality (Guo et al., 2020). GIP
stimulates osteoblasts and increases bone formation. A decline in GIP leads to a decline in bone metabolism, which could be one of the mechanisms that induces osteopenia in diabetics (Zofkova, 2015). It is possible that all four genes are associated with shank length and further affect body weight as they are all involved in growth traits, which is also suggested by the progressive haplotype accumulation effect (Figure 5). In short, the above genes provide a starting point to further study the shank traits. The next analysis requires multi-omics methods, i.e., combined with a map-based approach, gene expression analysis, metabolic regulation analysis, causality analysis, and other optional methods to investigate the molecular mechanism and causal mutations in this region.

In summary, the HQLA-HB-AIL chicken, which balanced the avoidance of rare alleles with the requirement for rapid linkage disequilibrium (LD) decay, is a reasonable resource for detecting quantitative trait genes. This AIL yielded a much narrower QTL than the F3 generations, especially the QTL on chromosome 27. Further, we highlighted the important role of four candidate genes (PHOSPHO1, IGF2BP1, ZNF652, and GIP) for bone development. We also identified a missing QTL on chromosome 4 via the joint analysis of GWAS and a selection signature analysis, which demonstrated the local limitations of this population but can be remedied by a multidimensional analysis. Overall, our study provides a promising resource for this field of study and will facilitate our understanding of the genetic mechanisms underlying chicken bone growth.

DATA AVAILABILITY STATEMENT

All data are published and can be found in Sheng et al. (2013) and Wang et al. (2017). The genotyping data can be found in doi: 10.6084/m9.figshare.12581063.v1. The phenotype data can be found in Supplementary Table 4.

ETHICS STATEMENT

The animal welfare committee of the State Key Laboratory approved all animal care and experimental procedures for Agrobiotechnology of China Agricultural University with approval number SKLAB-2014-06-07.

AUTHOR CONTRIBUTIONS

XH, CL, HQ, NL, and YW conceived and designed the experiments. YW and XC performed the experiments. YW, LB, and CZ analyzed the data. YW, LB, XC, JR, and ZH contributed reagents, materials, and analysis tools. YW wrote the manuscript. DS, HQ, CL, XC, and XH provided comments on the manuscript. All authors contributed to the article and approved the submitted version.

FUNDING

This work was financially supported by the National Natural Science Foundation of China (31961133003, 31672411, and 31902143) and the Open Research Program of State Key Laboratory for Agro-Biotechnology (2020SKLAB6-25).

ACKNOWLEDGMENTS

We gratefully acknowledge Jingyi He and Xinchun Xu for their help in collecting samples.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2020.00894/full#supplementary-material

REFERENCES

Alexander, D. H., Novembre, J., and Lange, K. (2009). Fast model-based estimation of ancestry in unrelated individuals. Genome Res. 19, 1655–1664. doi: 10.1101/gr.094052.109

Andersson, L., Haley, C. S., Ellegren, H., Knott, S. A., Johansson, M., Andersson, K., et al. (1994). Genetic mapping of quantitative trait loci for growth and fatness in pigs. Science 263, 1771–1774. doi: 10.1126/science.8134840

Arends, D., Heise, S., Karst, S., Trost, J., and Brockmann, G. A. (2016). Fine mapping a major obesity locus (Obes1) using a Berlin Fat Mouse x B6N advanced intercross population. Int. J. Obes. 40, 1784–1788. doi: 10.1038/ijo.2016.150

Bell, J. L., Wachter, K., Muhleck, B., Pazaitis, N., Kohn, M., Lederer, M., et al. (2013). Insulin-like growth factor 2 mRNA-binding proteins (IGF2BPs): post-transcriptional drivers of cancer progression? Cell Mol. Life Sci. 70, 2657–2675. doi: 10.1007/s00018-012-1186-z

Besnier, F., Wahlberg, P., Ronnegard, L., Ek, W., Andersson, L., Siegel, P. B., et al. (2011). Fine mapping and replication of QTL in outbred chicken advanced intercross lines. Genet. Sel. Evol. 43:3. doi: 10.1186/1297-9686-43-3
Deeb, N., and Lamont, S. J. (2002). Genetic architecture of growth and body composition in unique chicken populations. J. Hered. 93, 107–118. doi:10.1093/jhered/93.2.107

Doitsidou, M., Jarriault, S., and Poole, R. J. (2016). Next-Generation Sequencing-Based Approaches for Mutation Mapping and Identification in Caenorhabditis elegans. Genetics 204, 451–474. doi:10.1534/genetics.115.186197

Gao, Y., Du, Z. Q., Feng, C. G., Deng, X. M., Li, N., Da, Y., et al. (2010). Identification of quantitative trait loci for shank length and growth at different development stages in chicken. Anim. Genet. 41, 101–104. doi:10.1111/j.1365-2052.2009.01018.x

Gatti, D. M., Svensson, K. L., Shabalina, A., Wu, L. Y., Valdar, W., Simecek, P., et al. (2014). Quantitative trait locus mapping methods for diversity outbred mice. G3: 1623–1633. doi:10.1534/g3.113.013748

Gu, X., Feng, C., Ma, L., Song, C., Wang, Y., Da, Y., et al. (2011). Genome-wide association analysis in a mouse advanced intercross line. Nat. Commun. 5:9562. doi:10.1038/ncomms9562

Hansen, T. V., Hammer, N. A., Nielsen, J., Madsen, M., Dalbaeck, C., Wewer, U. M., et al. (2014). Quantitative trait locus analysis in crosses between New Hampshire and White Leghorn chicken lines: IV. Growth performance. Anim. Genet. 46, 441–446. doi:10.1111/age.12298

Kichaev, G., Bhatia, G., Loh, P. R., Gazal, S., Freund, M. K., et al. (2019). Leveraging Polygenic Functional Enrichment to Improve GWAS Power. Am. J. Hum. Genet. 104, 65–75. doi:10.1016/j.ajhg.2018.11.008

Lango Allen, H., Estrada, K., Lettre, G., Berndt, S. I, Weedon, M. N., Rivadeneira, F., et al. (2010). Hundreds of variants clustered in genomic loci and biological pathways affect human height. Nature 467, 832–838. doi:10.1038/nature09410

Liu, R., Sun, Y., Zhao, G., Wang, F., Wu, D., Zheng, M., et al. (2013). Genome-wide association study identifies Loci and candidate genes for body composition and meat quality traits in Beijing-You chickens. PLoS One 8:e61172. doi:10.1371/journal.pone.0061172

Maples, B. K., Gravel, S., Kenny, E. E., and Bustamante, C. D. (2013). RFMix: a discriminative modeling approach for rapid and robust local-ancestry inference. Am J Hum Genet. 93, 278–288. doi:10.1016/j.ajhg.2013.06.020

Nassar, M. K., Gorogara, Z. S., and Brockmann, G. A. (2015). Quantitative trait loci segregating in crosses between New Hampshire and White Leghorn chicken lines: IV. Growth performance. Anim. Genet. 46, 441–446. doi:10.1111/age.12298

Parker, C. C., Carbonetto, P., Sokoloff, G., Park, Y. J., Abney, M., and Palmer, A. A. (2014). High-resolution genetic mapping of complex traits from a combined analysis of F2 and advanced intercross mice. Genetics 198, 103–116. doi:10.1534/genetics.114.167056

Perez-Enciso, M., Fernando, R. L., Bidanel, J. P., and Le Roy, P. (2001). Quantitative trait locus analysis in crosses between outbred lines with dominance and inbreeding. Genetics 159, 413–422.

Poland, J. A., Bradbury, P. J., Buckler, E. S., and Nelson, R. J. (2011). Genome-wide nested association mapping of quantitative resistance to northern leaf blight in maize. Proc. Natl. Acad. Sci. U. S. A. 108, 6893–6898. doi:10.1073/pnas.1008491048

Sasaki, O., Odawara, S., Takahashi, H., Nirasawa, K., Oyamada, Y., Yamamoto, R., et al. (2004). Genetic mapping of quantitative trait loci affecting body weight, egg character and egg production in F2 intercross chickens. Anim. Genet. 35, 188–194. doi:10.1111/j.1365-2052.2004.01133.x

Sheng, Z., Pettersson, M. E., Hu, X., Luo, C., Qu, H., Shu, D., et al. (2013). Genetic dissection of growth traits in Chinese indigenous x commercial broiler chicken cross. BMC Genomics 14:151. doi:10.1186/1471-2164-14-151

Szeicz, Z. A., and Hernandez, R. D. (2014). selscan: an efficient multithreaded program to perform EHH-based scans for positive selection. Mol. Biol. Evol. 31, 2824–2827. doi:10.1093/molbev/msu211

Tsdzuki, M., Onitsuka, S., Akiyama, R., Iwamizu, M., Goto, N., Nishibori, M., et al. (2007). Identification of quantitative trait loci affecting shank length, body weight and carcass weight from the Japanese cockfighting chicken breed, Oh-Shamo (Japanese Large Game). Cytogeten Genome Res. 117, 288–295. doi:10.1159/000103190

Van Goor, A., Boele, K. J., Ashwell, C. M., Persia, M. E., Rothschild, M. F., Schmidt, C. J., et al. (2015). Identification of quantitative trait loci for body temperature, body weight, breast yield, and digestibility in an advanced intercross line of chickens under heat stress. Genet. Sel. Evol. 47:96. doi:10.1186/s12711-015-0176-7

Wang, Y., Cao, X., Zhao, Y., Fei, J., Hu, X., and Li, N. (2017). Optimized double-digest genotyping by sequencing (ddGBS) method with high-density SNP markers and high genotyping accuracy for chickens. PLoS One 12:e0179073. doi:10.1371/journal.pone.0179073

Yalcin, B., Nicod, J., Bomhra, A., Davidson, S., Cleak, J., Farinelli, L., et al. (2010). Commercially available outbred mice for genome-wide association studies. PLoS Genet 6:e1001085. doi:10.1371/journal.pgen.1001085

Yang, J., Lee, S. H., Goddard, M. E., and Visser, P. M. (2011). GCTA: a tool for genome-wide complex trait analysis. Am. J. Hum. Genet. 88, 76–82. doi:10.1016/j.ajhg.2010.11.011

Yang, L., Guo, X., Zhu, D., Bian, C., Zhao, Y., Tan, C., et al. (2019). Genome-wide association analyses of multiple traits in Duroc pigs using low-coverage whole-genome sequencing strategy. bioXiv

Zan, Y., Payen, T., Lillie, M., Honaker, C. F., Siegel, P. B., and Carlborg, O. (2019). Genotyping by low-coverage whole-genome sequencing in intercross pedigrees
from outbred founders: a cost-efficient approach. *Genet. Sel. Evol.* 51:44. doi: 10.1186/s12711-019-0487-1

Zan, Y., Sheng, Z., Lillie, M., Ronnegard, L., Honaker, C. F., Siegel, P. B., et al. (2017). Artificial Selection Response due to Polygenic Adaptation from a Multilocus, Multiallelic Genetic Architecture. *Mol. Biol. Evol.* 34, 2678–2689. doi: 10.1093/molbev/msx194

Zhang, C., Dong, S. S., Xu, J. Y., He, W. M., and Yang, T. L. (2019). PopLDdecay: a fast and effective tool for linkage disequilibrium decay analysis based on variant call format files. *Bioinformatics* 35, 1786–1788. doi: 10.1093/bioinformatics/bty873

Zhou, Z., Li, M., Cheng, H., Fan, W., Yuan, Z., Gao, Q., et al. (2018). An intercross population study reveals genes associated with body size and plumage color in ducks. *Nat. Commun.* 9:2648. doi: 10.1038/s41467-018-04868-4

Zofkova, I. (2015). Bone tissue as a systemic endocrine regulator. *Physiol Res.* 64, 439–445. doi: 10.33549/physiolres.932900

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2020 Wang, Bu, Cao, Qu, Zhang, Ren, Huang, Zhao, Luo, Hu, Shu and Li. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.