Epistatic Effects on Abdominal Fat Content in Chickens: Results from a Genome-Wide SNP-SNP Interaction Analysis

Fangge Li¹,²,³, Guo Hu⁴, Hui Zhang¹,²,³, Shouzhi Wang¹,²,³, Zhipeng Wang¹,²,³, Hui Li¹,²,³

¹ Key Laboratory of Chicken Genetics and Breeding, Ministry of Agriculture, Harbin, People’s Republic of China, ² Key Laboratory of Animal Genetics, Breeding and Reproduction, Education Department of Heilongjiang Province, Harbin, People’s Republic of China, ³ College of Animal Science and Technology, Northeast Agricultural University, Harbin, People’s Republic of China, ⁴ Heilongjiang River Fishery Research Institute, Chinese Academy of Fishery Sciences, Harbin, People’s Republic of China

Abstract

We performed a pairwise epistatic interaction test using the chicken 60 K single nucleotide polymorphism (SNP) chip for the 11th generation of the Northeast Agricultural University broiler lines divergently selected for abdominal fat content. A linear mixed model was used to test two dimensions of SNP interactions affecting abdominal fat weight. With a threshold of P<1.2×10⁻⁶ by a Bonferroni 5% correction, 52 pairs of SNPs were detected, comprising 45 pairs showing an Additive x Additive and seven pairs showing an Additive x Dominance epistatic effect. The contribution rates of significant epistatic interactive SNPs ranged from 0.62% to 1.54%, with 47 pairs contributing more than 1%. The SNP-SNP network affecting abdominal fat weight constructed using the significant SNP pairs was analyzed, estimated and annotated. On the basis of the network’s features, SNPs Gga_rs14303341 and Gga_rs14988623 at the center of the subnet should be important nodes, and an interaction between GGAZ and GGA8 was suggested. Twenty-two quantitative trait loci, 97 genes (including nine non-coding genes), and 50 pathways were annotated on the epistatic interactive SNP-SNP network. The results of the present study provide insights into the genetic architecture underlying broiler chicken abdominal fat weight.

Introduction

Epistasis – the interactions between polymorphic loci, such as SNPs, genes or quantitative trait loci (QTLs) – is a hot topic in quantitative genetics [1]. Epistasis is a major factor that determines variation in quantitative phenotypes [2]. Mapping the epistatic loci affecting quantitative traits will advance our understanding of the genetic architecture of these complex phenotypes for humans and model organisms [2,3]. Recently, the study of epistasis among genes, SNPs or QTLs has progressed rapidly [4–8], as has the study of epistasis in biological experiments [9].

Much progress has been made using genome-wide association studies (GWASs). Performing genome-wide SNPs interaction analysis represents the next step for detecting the variations of quantitative traits, because single-locus tests cannot identify the interactions among SNPs, genes or other genetic or environmental factors [10]. The markers detected by GWAS only explained a fraction of the heritable variance [11–13]. Identifying markers that show interactions would help to explain a higher proportion of the heritable variance. From sequence to phenotype, the SNP-SNP and gene-gene interactions can provide new insights into the genetic basis of complex traits.

In chickens (Gallus gallus), it has been suggested that epistatic interactions between genes (or QTLs) are important for quantitative traits such as growth and fatness traits [14–16]. Abdominal fat in meat-type chicken is a quantitative trait, and is an adverse economic factor; therefore, many QTLs and SNPs affecting fatness traits have been identified [17–21]. A previous study on chicken abdominal fat traits identified epistatic interactions among 10 candidate genes, and constructed networks of the interacting genes [15]. However, a large proportion of the effects of the epistatic interactions between polymorphic loci on fatness traits remain undetermined, and the whole genetic network is unclear. Hence, it is essential to study the effects of interactions between SNPs, genes and QTLs on fatness traits.

In this study, we aimed to determine the interactions between polymorphic loci affecting chicken abdominal fat weight (AFW). Pairwise epistatic interaction effects among genome-wide SNPs were detected. A network of significant SNP pairs was analyzed and annotated. The results provide further insight into the genetic network controlling abdominal fat deposition in chickens.

Materials and Methods

Ethics Statement

All animal work was conducted according to the guidelines for the care and use of experimental animals established by the Ministry of Science and Technology of the People’s Republic of China (Approval number: 2006–398) and was approved by the...
Epistatic Effects on Abdominal Fat in Chickens

Experimental Populations
The experimental chickens were from the Northeast Agricultural University. The experimental population was constructed by the Laboratory Animal Management Committee of Northeast Agricultural University.

SNP Genotyping
Genotyping was carried out using the Illumina Inc. (San Diego, CA, USA) chicken 60 K SNP chip, which contains 57,636 SNPs. Markers with minor allele frequencies less than 5% or monomorphic loci were filtered out. Individuals were removed who had 5% or more missing SNP genotypes. Finally, 45,611 SNPs in 475 individuals were used for the pairwise interaction analyses. SNP genotypes were provided by Zhang et al. [23].

Genome-wide Pairwise Interaction Analysis
The EPISNP3 module in epiSNP_v4.2_Windows software package (Y. Da, Department of Animal Science, University of Minnesota, USA, http://animalgene.umn.edu/episnp/download.html) was used to test for epistatic effects [24].

The EPISNP's statistical model for testing epistatic effects for AFW is

\[ y = \mu + SNP_1 + SNP_2 + SNP_1 \times SNP_2 + F + e \]

where \( y \) is the dependent variable (AFW), \( \mu \) is the population mean, \( SNP_1 \) and \( SNP_2 \) are the two single-locus genotypic effects, \( SNP_1 \times SNP_2 \) is the two-locus interaction effect, \( F \) (family effect) is a random effect, and \( e \) is the random error.

The two-locus interaction effect was partitioned into four individual epistatic effects using the extended Kempthorne model, which allows Hardy-Weinberg disequilibrium and linkage disequilibrium (LD) to be considered. Additive \( \times \) Additive (AA), Additive \( \times \) Dominant (AD), Dominant \( \times \) Additive (DA), and Dominant \( \times \) Dominant (DD) effects [24]. An F-test was used to test the significance of the two-locus interaction effect.

The \( P \)-value of model (1) was adjusted by a Bonferroni correction \((4.16 \times 10^{-6})\). Independent tests using the same data set, with a significance threshold of \( P < 0.05 \), and the significance threshold of the test was determined as \( P < 0.2 \times 10^{-2} \).

Contribution Rate Calculation
The contribution rate of every significant epistatic interactive SNP pair to phenotypic variation was calculated by the formula,

\[ c = \frac{SS_{SNP_1 \times SNP_2}}{Var_F} \times 100\% \]

where \( SS_{SNP_1 \times SNP_2} \) is the variance of the significant \( SNP_1 \times SNP_2 \) interactive effect (AA, AD, DA or DD), and \( Var_F \) is the phenotype variance.

SNP-SNP Network
The figure showing the SNP-SNP network with significant epistatic effects was drawn using the Cytoscape 2.8 software package [25]. The interactive SNPs affecting AFW with a pairwise interaction \( P \)-value of \( P < 1.20 \times 10^{-11} \) were loaded into Cytoscape to visualize the network. Based on entropy theory [26], the importance of the different subnets was evaluated by the formula,

\[ w = \sum_{i=1}^{n} (- \log p_i) \times c_i \]

where \( w \) is the importance of the subnet, \( n \) is the edge number (i.e., the number of SNP pairs), \( p_i \) is the \( P \)-value of the interactive effect being tested for the \( i \)-th SNP pair, and \( c_i \) is the contribution rate of the \( i \)-th SNP pair.

Annotation of SNP-SNP Network
The purpose of this step was to annotate significant SNPs and mine QTLs or gene networks affecting abdominal fat traits. First, the significant interactive SNPs were put into the Chicken QTL database (http://www.animalgenome.org/cgi-bin/QTLDb/GG/index), and mapped in terms of QTLs. Second, the network was annotated with genes and pathways. A mean value of \( r^2 \geq 0.8 \) [27] was observed in pairwise distances of approximately 0.2 Mb, as calculated by HAPLOVIEW v4.1 [28], and genes whose physical distance to the significant SNP was smaller than 0.2 Mb were selected. The positions of genes were acquired from UCSC Genome Bioinformatic site (http://genome.ucsc.edu/). Genes in the same network were put into the KEGG program (http://www.kegg.jp/kegg/) to identify pathways that might affect the traits.

Results
SNP Makers
A total of 45,005 SNPs on 28 autosomes, the Z chromosome, linkage groups and 606 SNPs not assigned to any chromosomes in chickens were included in this study (Table 1). These markers covered 1026.23 Mb of the genome, with an average of 16.63 kb between adjacent markers.

Pairwise Interaction Effects Analysis
EPISNP3 [24] was used to analyze the pairwise interaction effects among genome-wide SNPs for AFW across both lines. Fifty-two pairs of significant SNPs (Table 2), containing 45 pairs of AA and seven pairs of AD epistatic effects, were detected. No DA or DD epistatic effects were detected. The \( P \)-value of the epistatic effect between SNP Gga_rs13569377 on GGA18 (Gallus gallus, GGA) and Gga_rs14988623 on GGA13 reached \( 2.54 \times 10^{-11} \), which was the most significant effect detected. There were six pairs of SNPs on the same chromosome (one pair on GGA2, four pairs on GGA3 and one pair on GGA10). The other 46 pairs of SNPs were distributed on different chromosomes. The contribution rate of the significant epistatic interactive SNP pairs ranged from 0.62% to 1.54%, with 47 pairs having a contribution rate of more than 1%. The 52 pairs of significant SNPs comprised 68 single SNPs covering 18 chromosomes. The position of each SNP is shown in Table S1. Many SNPs were detected to interact with the same locus; for example, seven SNPs on GGA3 all had epistatic interactions with Gga_rs14303341 on GGA27, and four SNPs on GGA18 interacted with Gga_rs14988623 on GGA13. The SNP pairs having the same SNPs are shown in Figure 1.

SNP-SNP Network Analysis
To investigate the complex mechanism of epistatic effects on AFW, a network of SNPs having epistatic interaction affecting AFW was constructed. SNP epistatic interaction subnets containing more than three nodes are shown in Figure 1. The degree of nodes refers to the number of edges connecting it, and an edge stands for a two-way interaction. For example,
Table 1. Summary of genome-wide markers.

| GGA | SNPs' number | GGA length (Mb) | Mean distance (kb) | GGA | SNPs' number | GGA length (Mb) | Mean distance (kb) |
|-----|--------------|----------------|-------------------|-----|--------------|----------------|-------------------|
| 1   | 7,143        | 200.95         | 28.13             | 17  | 844          | 10.61          | 12.57             |
| 2   | 5,299        | 154.46         | 29.15             | 18  | 845          | 10.89          | 12.89             |
| 3   | 4,081        | 113.65         | 27.85             | 19  | 804          | 9.89           | 12.30             |
| 4   | 3,314        | 94.16          | 28.41             | 20  | 1,460        | 13.92          | 9.53              |
| 5   | 2,172        | 62.23          | 28.65             | 21  | 726          | 8.88           | 9.48              |
| 6   | 1,714        | 35.84          | 20.91             | 22  | 295          | 3.89           | 13.19             |
| 7   | 1,770        | 38.17          | 21.56             | 23  | 577          | 6.02           | 10.43             |
| 8   | 1,394        | 30.62          | 21.97             | 24  | 676          | 6.23           | 9.22              |
| 9   | 1,168        | 24.02          | 20.57             | 25  | 170          | 2.02           | 11.88             |
| 10  | 1,297        | 22.42          | 17.29             | 26  | 617          | 5.03           | 8.15              |
| 11  | 1,196        | 21.87          | 18.29             | 27  | 472          | 4.84           | 10.25             |
| 12  | 1,324        | 20.45          | 15.45             | 28  | 563          | 4.46           | 7.92              |
| 13  | 1,128        | 18.32          | 16.24             | LEG22 | 103     | 0.89         | 8.64              |
| 14  | 984          | 15.76          | 16.02             | LEG64 | 2       | 0.005        | 2.27              |
| 15  | 1,010        | 12.93          | 12.80             | Z         | 1,844   | 74.63        | 40.47             |
| 16  | 13           | 0.17           | 13.08             | UN*      | 606     | /            | /                 |

*These SNPs were not assigned to any chromosomes.

doi:10.1371/journal.pone.0081520.t001

Discussion

Abdominal fatness traits in chickens are important quantitative traits, and are under complex genetic control. Understanding the molecular mechanisms underlying them will help efficient growth selection in broiler chickens. In this study, analyses using genome-wide SNPs were performed to examine the genetic contribution of epistatic effects to the phenotypic variation of chicken abdominal fatness traits.

GWASs have identified many significant association markers. However, many studies showed that the identified markers explain only a fraction of the heritable variance [11–13]. This discrepancy has been ascribed to the insufficient power of GWAS to identify gene–gene interactions [13]. Zink et al. [29] showed that the “missing” heritability of Crohn’s disease is 62.8%, and that genetic interactions could account for 80% of this missing heritability. Recently, most genome-wide association analyses in chickens have focused on individual SNP marker association analysis [20,30]. However, quantitative traits often arise from the combined effects...
Table 2. Genome-wide significant pairwise epistatic interactive SNP pairs for abdominal fat weight, P<1.20×10^{-11}.

| GGAF\(^1\) | SNP\(_1\) name | GGAS\(^1\) | SNP\(_2\) name | Epistatic effect type | P-value | c (%) |
|---|---|---|---|---|---|---|
| 0 | GGaluGA194739 | 1 | GGaluGA012915 | AA | 1.18×10^{-11} | 1.08 |
| 1 | Gga_rs13866305 | 3 | Gga_rs13717259 | AD | 4.90×10^{-12} | 0.68 |
| 1 | Gga_rs13749637 | 13 | Gga_rs16002106 | AA | 7.47×10^{-12} | 1.40 |
| 1 | Gga_rs15227054 | 13 | Gga_rs16002106 | AA | 8.52×10^{-12} | 1.38 |
| 1 | Gga_rs13749637 | 13 | GGaluGA097211 | AA | 1.02×10^{-11} | 1.44 |
| 1 | Gga_rs13749637 | 13 | GGaluGA097233 | AA | 7.50×10^{-13} | 1.54 |
| 1 | Gga_rs15227054 | 13 | GGaluGA097233 | AA | 8.57×10^{-13} | 1.51 |
| 1 | GGaluGA060937 | 14 | GGaluGA101229 | AA | 3.17×10^{-12} | 1.26 |
| 2 | Gga_rs16026770 | 2 | GGaluGA146662 | AA | 6.60×10^{-12} | 1.05 |
| 2 | Gga_rs14214495 | 13 | Gga_rs15683090 | AA | 9.31×10^{-12} | 1.14 |
| 3 | Gga_rs14319575 | 3 | Gga_rs14402423 | AA | 5.96×10^{-12} | 0.91 |
| 3 | Gga_rs14380677 | 3 | Gga_rs16306728 | AA | 8.25×10^{-12} | 1.10 |
| 3 | Gga_rs16306728 | 3 | GGaluGA213041 | AA | 1.09×10^{-11} | 1.07 |
| 3 | Gga_rs14319575 | 3 | GGaluGA361222 | AA | 5.96×10^{-12} | 0.91 |
| 3 | Gga_rs14388313 | 5 | Gga_rs14521876 | AA | 8.46×10^{-13} | 1.14 |
| 3 | Gga_rs14368109 | 10 | GGaluGA071224 | AD | 6.80×10^{-12} | 0.62 |
| 3 | Gga_rs14368127 | 10 | GGaluGA071224 | AD | 7.23×10^{-12} | 0.63 |
| 3 | Gga_rs16228738 | 14 | Gga_rs14075705 | AA | 8.12×10^{-13} | 1.42 |
| 3 | Gga_rs16222762 | 20 | Gga_rs14272866 | AA | 1.30×10^{-12} | 1.13 |
| 3 | Gga_rs16228738 | 23 | Gga_rs13622160 | AA | 7.99×10^{-12} | 1.20 |
| 3 | Gga_rs16228738 | 23 | GGaluGA188871 | AA | 9.80×10^{-12} | 1.15 |
| 3 | Gga_rs14340790 | 27 | Gga_rs14303341 | AA | 1.16×10^{-11} | 1.25 |
| 3 | Gga_rs14341204 | 27 | Gga_rs14303341 | AA | 1.42×10^{-12} | 1.29 |
| 3 | Gga_rs14341224 | 27 | Gga_rs14303341 | AA | 6.31×10^{-13} | 1.36 |
| 3 | Gga_rs14341242 | 27 | Gga_rs14303341 | AA | 5.68×10^{-13} | 1.38 |
| 3 | Gga_rs14341255 | 27 | Gga_rs14303341 | AA | 5.68×10^{-13} | 1.38 |
| 3 | Gga_rs16254447 | 27 | Gga_rs14303341 | AA | 6.31×10^{-13} | 1.36 |
| 3 | GGaluGA216762 | 27 | Gga_rs14303341 | AA | 1.36×10^{-12} | 1.31 |
| 4 | Gga_rs15480969 | 20 | Gga_rs14276105 | AA | 1.05×10^{-11} | 1.54 |
| 7 | GGaluGA317680 | 14 | Gga_rs15718248 | AA | 1.05×10^{-11} | 1.23 |
| 9 | Gga_rs16674724 | 3 | Gga_rs14319575 | AA | 1.05×10^{-12} | 1.01 |
| 10 | Gga_rs15583507 | 6 | Gga_rs13561344 | AA | 2.80×10^{-12} | 1.32 |
| 10 | Gga_rs14009265 | 6 | Gga_rs14560750 | AA | 5.35×10^{-13} | 1.18 |
| 10 | Gga_rs15583507 | 6 | Gga_rs14560750 | AA | 2.15×10^{-12} | 1.30 |
| 10 | GGaluGA066690 | 10 | GGaluGA066877 | AA | 4.66×10^{-13} | 1.27 |
| 10 | GGaluGA069801 | 23 | Gga_rs13622160 | AA | 5.56×10^{-12} | 1.15 |
| 10 | GGaluGA069801 | 23 | Gga_rs14290610 | AA | 4.64×10^{-12} | 1.26 |
| 14 | Gga_rs14068999 | 23 | GGaluGA188871 | AA | 3.29×10^{-12} | 1.12 |
| 14 | Gga_rs15717370 | 23 | GGaluGA188871 | AA | 3.29×10^{-12} | 1.12 |
| 18 | Gga_rs10729280 | 13 | Gga_rs14988623 | AD | 6.56×10^{-14} | 1.10 |
| 18 | Gga_rs13569377 | 13 | Gga_rs14988623 | AD | 2.54×10^{-14} | 1.14 |
| 18 | Gga_rs14416916 | 13 | Gga_rs14988623 | AD | 9.00×10^{-14} | 1.07 |
| 18 | Gga_rs15469971 | 13 | Gga_rs14988623 | AD | 2.92×10^{-14} | 1.10 |
| Z | Gga_rs14748835 | 8 | Gga_rs14658668 | AA | 1.14×10^{-11} | 1.20 |
| Z | Gga_rs16094710 | 8 | Gga_rs14658668 | AA | 1.14×10^{-11} | 1.20 |
| Z | Gga_rs16758057 | 8 | Gga_rs14658668 | AA | 1.14×10^{-11} | 1.20 |
of multiple loci [14–16]. In the present study, the contribution rates of the significant epistatic interactive SNP pairs to AFW ranged from 0.62% to 1.54%. Two SNP pairs showed the largest contribution rate of 1.54%, and the contribution rates of 47 pairs were more than 1%. In addition, we calculated the contribution rates of single SNPs in the GWAS (Table S5). The single SNP Gga_rs14276105, which was significant in GWAS (Table S5), showed the largest contribution rate 1.46%. Comparing the results showed that the contribution rates of significant SNP pairs was larger than that of the single SNPs contained in the pairs. Although the contribution rates of the significant epistatic interactive SNP pairs might be overestimated for model (1), there were no high dimension interactions; therefore, we concluded that epistasis has an important effect on the phenotypic variance and showed the largest contribution rate 1.46%. Comparing the results showed that the contribution rates of significant SNP pairs was larger than that of the single SNPs contained in the pairs. Although the contribution rates of the significant epistatic interactive SNP pairs might be overestimated for model (1), there were no high dimension interactions; therefore, we concluded that epistasis has an important effect on the phenotypic variance and

Table 2. Cont.

| GGA F   | SNP name  | GGA S   | SNP name  | Epistatic effect type | P-value | c (%) |
|---------|-----------|---------|-----------|-----------------------|---------|-------|
| Z       | Gga_rs14748835 8 GgaluGA333545 | AA       | 1.14 x 10^-11 | 1.20 |
| Z       | Gga_rs16094710 8 GgaluGA333545 | AA       | 1.14 x 10^-11 | 2.05 |
| Z       | Gga_rs15991936 10 Gga_rs15589655 | AA       | 8.33 x 10^-12 | 1.28 |

GGA F = The first chromosome in the pairwise epistasis analysis; GGA S = The second chromosome in the pairwise epistasis analysis; AA = Additive x Additive effect, AD = Additive x Dominance effect, DA = Dominance x Additive effect, DD = Dominance x Dominance effect; P-value = P-value of the effect being tested; c = The contribution rate (%) of every significant epistatic interactive SNP pair.

doi:10.1371/journal.pone.0081520.t002

Figure 1. Epistatic network among SNPs affecting abdominal fat weight (AFW) in NEAUHLF. A node represents a SNP. The chromosome in which the SNP is located is shown in the circle. A pair of SNPs connected by an edge has a significant effect. The colors of the nodes represent P-values of the interaction (P<1 x 10^-13 = red; P<1 x 10^-12 = blue; P<1 x 10^-11 = green; P<1 x 10^-10 = white). The color of the edge indicates the epistatic effect type (AA = red; AD = purple).

doi:10.1371/journal.pone.0081520.g001
cannot be ignored. Our results also showed that Gga_rs14276105 was the only SNP that was significant in an association study among the significantly interacting SNPs (Table S3). Wu et al. [31] also demonstrated that the majority of significantly interacting SNPs did not show marginal association in their data. This suggested that testing interactions of SNPs with only one significant locus association would lose the majority of interactions.

The features of scale-free networks are several nodes with large degrees and many nodes with few connections. Further, the nodes with large degrees should be the hub sites of the network. Our SNP-SNP network displayed the features of a scale-free network. The degree of one node was 7, and of another was 4; and 47 nodes had a degree of 1 in our SNP–SNP network. Thus, the nodes (Gga_rs14989823 and Gga_rs14303341) with larger degrees should be the important ones. Scale-free behavior has been demonstrated in many biological interaction networks, such as protein or gene interaction networks [32].

The formula for evaluating the importance of subnets proposed in the present study was based on the entropy theory [26]. The evaluation indices comprised the edge number, P-value and the contribution rate of every SNP pair in the subnets. The importance evaluation score stands for the information quantity of a subnet. A subnet containing more information would be more important. We suggested that subnets A and C are the most important; as they received the top two importance evaluation scores (Table 3).

Subnet A was radial, with seven SNPs connecting the center. A previous study concluded that many significant epistatic effects involving one locus were less likely to be random than was a single epistatic effect [39]; i.e., the network is less likely to be random. Combined with the results of the scale-free network analysis, we identified Gga_rs14303341 as the important SNP. The annotated information indicated that Gga_rs14303341 maps to QTLs (11809, 11817) [34], and the genes GOSR2, GFCL, CDDC43, WNT3, ASF and EFTUD2 are within the defined 0.4 Mb region. These QTLs and genes should receive attention in a future study.

Subnet C was a loop graph, and contained six SNPs: three on GGA3 and three on GGA8. Between the two chromosomes, nine (3 × 3) pairs of SNP interactions were obtained, and eight of these interactions were significant. In addition, the SNPs were adjacent to one another on GGA3 and GGA8. The result indicated that the interaction occurs between the two regions on GGA3 (59,561,642–60,090,313) and GGA8 (28,415,409–28,925,822).

The epistatic interactions of SNPs might provide the basis for identifying interactions among genes, QTLs, or even pathways; therefore, we annotated the network. The result showed that 35% (24/68) of the SNP markers are located in regions containing 22 QTLs associated with chicken AFW (Table S2). The radial network based on subnet A suggested interactions between QTLs. A QTLs network with a similar topological structure was found in a study of chicken growth traits by Carlborg et al. [16]. They identified five significant epistatic QTL pairs that also formed a radial network and that accounted for a significant amount of the phenotypic variance.

Gene-by-gene interaction plays a major role in genetic studies of quantitative traits; however, the detection of gene–gene interaction has been traditionally assessed by SNP interactions. Single SNPs cannot capture the total variation of a gene; therefore, SNP interactions cannot represent gene–gene interactions. Thus, Cui et al. [35] and Li et al. [36] extended the idea of single SNP interactions to haplotype interactions, and proposed a novel statistical approach for capturing variations in genes and potential interactions between two genes. In the present study, we found an interesting phenomenon in some subnets, in which SNPs on the circumference were very near in physical distance. Therefore, we believe that the definition of block interaction may provide additional biological insights, compared with the analysis of SNP interactions.

In this study, we detected 26 regions on 12 chromosomes. Ninety-seven genes (including nine non-coding genes), including BMPRIA, GIPR, GRB2, LITAF, SOCS3, WNT3 and PDPK1, were identified in these regions. Based on the literature, some genes are associated with obesity; for example, BMPRIA is associated with human obesity [37]. GIPR on GGA18 is associated with fat droplet formation [38]. LITAF plays an important role in the regulation of tumor necrosis factor alpha (TNF-α), which is important in adipose tissue [39,40]. WNT3 is part of the Wnt-signaling pathway, which is one of the pathways operating in fat cells [41]. PDPK1 affects the fat-pad mass in mice [42]. We identified 50 pathways related to AFW, according to gene annotation analysis (Table S4). Some of the pathways are related to obesity, such as the Jak-STAT-signaling pathway and the insulin-signaling pathway [43,44].

### Table 3. Description of SNPs epistatic interaction network.

| Subnet | Importance | Node number | Edge number | The greatest degree | Structure | GGA$^3$ |
|--------|------------|-------------|-------------|--------------------|----------|--------|
| Subnet A | 1.115 | 8 | 7 | 7 | Tree graph | 3, 27 |
| Subnet B | 0.961 | 8 | 7 | 3 | Tree graph | 3, 10, 14, 23 |
| Subnet C | 1.051 | 6 | 8 | 3 | Loop graph | 8, Z |
| Subnet D | 0.837 | 5 | 5 | 3 | Loop graph | 1, 13 |
| Subnet E | 0.589 | 5 | 4 | 4 | Tree graph | 13, 18 |
| Subnet F | 0.325 | 4 | 3 | 3 | Tree graph | 3, 9 |
| Subnet G | 0.450 | 4 | 3 | 2 | Tree graph | 6, 10 |
| Subnet H | 0.139 | 3 | 2 | 2 | Tree graph | 3, 10 |
| Subnet I | 0.239 | 3 | 2 | 2 | Tree graph | 3 |

$^1$The greatest degree = The greatest number of nodes in the subnet;
$^2$Structure = Topological structure of the subnet;
$^3$Chromosomes = Chromosomes contained in the subnet.

doi:10.1371/journal.pone.0081520.t003

3Chromosomes = Chromosomes contained in the subnet.
2Structure = Topological structure of the subnet;
1The greatest degree = The greatest number of nodes in the subnet;
Conclusions

Fifty-two pairs of significant epistatic interactive SNPs and their interaction regions were proposed based on the network of significant epistatic interactive pairs. Relevant genes, pathways and QTLs were annotated. The results advanced our understanding of the genetic architecture of abdominal fatness traits, hinted at
their molecular mechanisms, and will contribute to faster progress in the artificial genetic selection of broiler chickens.

Supporting Information

Table S1 SNPs contained in the significant SNP pairs. (DOC)

Table S2 SNPs mapped to QTLs. The QTL information was obtained from http://www.animalgenome.org/cg-bin/ QTLdb/GG/i-index. (DOC)

Table S3 Information on protein coding genes. The information was obtained from http://www.genatlas.org/. (DOC)

Table S4 Genes in pathways. Pathway information was obtained from http://www.genome.org/kegg/. (DOC)

Table S5 Results of the genome-wide association study of significantly interacting SNPs. The threshold of GWAS is $P<2.7 \times 10^{-7}$ by a Bonferroni 5% correction. (DOC)

Acknowledgments

We thank all the members of the Poultry Breeding Group of the College of Animal Science and Technology, Northeast Agricultural University, for managing the chickens and collecting the data. We appreciate the help of Prof. Y. Da from the University of Minnesota in providing the epiSNP software package [24].

Author Contributions

Conceived and designed the experiments: HL, ZW. Performed the experiments: FL, GH, HZ, SW, ZW, HL. Contributed reagents/materials/analysis tools: FL, GH, HZ, SW, ZW, HL. Performed the bioinformatics analyses: HL, ZW. Performed the statistical analyses: FL, GH, HZ, SW, ZW, HL. Performed the experiments and managed the chickens and collecting the data: We appreciate the help of Prof. Y. Da from the University of Minnesota in providing the epiSNP software package [24].

References

1. Cordell HJ (2002) Epistasis: what it means, what it doesn’t mean, and statistical methods to detect it in humans. Hum Mol Genet 11: 2463–2468.
2. Huang W, Richards S, Carbone MA, Zhu D, Andhur RR, et al. (2012) Epistasis dominates the genetic architecture of Drosophila quantitative traits. Proc Natl Acad Sci U S A 109: 15553–15559.
3. Rajon E, Ploix JB (2012) The evolution of genetic architectures underlying quantitative traits. arXiv preprint arXiv:12100843.
4. Ritchie MD, White BC, Parker JS, Hahn LW, Moore JJ (2003) Optimization of neural network architecture using genetic programming improves detection and modeling of gene–gene interactions in studies of human diseases. BMC Bioinformatics 4: 28.
5. Lou XY, Chen GB, Yan L, Ma JZ, Zhu J, et al. (2007) A generalized combinatorial approach for detecting gene-by-gene and gene-by-environment interactions with application to nicotine dependence. Am J Hum Genet 80: 1125–1137.
6. Zhang Y, Liu JS (2007) Bayesian inference of epistatic interactions in case-control studies. Nat Genet 39: 1167–1173.
7. Alvarez-Castro JM, Carlborg O (2007) A unified model for functional and statistical epistasis and its application in quantitative trait Loci analysis. Genetics 176: 1151–1167.
8. Alvarez-Castro JM, Le Rouziec A, Andersson L, Siegel PB, Carlborg O (2012) Modelling of genetic interactions improves prediction of hybrid patterns: a case study in domestic fowl. Genet Res (Camb) 94: 255–266.
9. Fierst JL, Phillips PC (2012) Variance in epistasis links gene regulation and evolutionary rate in the yeast genetic interaction network. Genome Biol Evol 4: 1089–1097.
10. Cordell HJ (2009) Detecting gene-gene interactions that underlie human diseases. Nat Rev Genet 10: 392–404.
11. Eichler EE, Flint J, Gibson G, Kong A, Leal SM, et al. (2010) Missing heritability and strategies for finding the underlying causes of complex disease. Nat Rev Genet 11: 446–450.
12. Yang J, Benyamin B, McEvoy BP, Gordon S, Henders AK, et al. (2010) Common SNPs explain a large proportion of the heritability for human height. Nat Genet 42: 565–569.
13. Manolio TA, Collins FS, Cox NJ, Goldberg DB, Hindorff LA, et al. (2009) Finding the missing heritability of complex diseases. Nature 461: 747–753.
14. Kwok M, Marklund S, Ragan-Kelley A, Siegel P, Muir W, et al. (2012) Generation of a multi-locus chicken introgression line to study the effects of genetic interactions on metabolic phenotypes in chickens. Front Genet 3: 29.
15. Hu G, Wang SZ, Wang ZP, Li YM, Li H (2010) Genetic epistasis analysis of 10 peroxisome proliferator-activated receptor gamma-correlated genes in broiler lines divergently selected for abdominal fat content. Poult Sci 89: 2541–2550.
16. Carlborg O, Jacobsson L, Ahgren P, Siegel P, Andersson L (2006) Epistasis and the release of genetic variation during long-term selection. Nat Genet 38: 418–420.
17. Wang R, Hu XX, Wang LP, Li XC, Wang QG, et al. (2012) Quantitative trait loci associated with body weight and abdominal fat traits on chicken chromosomes 3, 5 and 7. Genet Mol Res 11: 956–965.
18. Le Magnien G, Piel F, Gilbert H, Le Bihan-Duval E, Viguoles F, et al. (2009) A comprehensive analysis of QTL for abdominal fat and breast muscle weights on chicken chromosome 5 using a multivariate approach. Anim Genet 40: 157–164.
19. Ahabat B, Dekkers JC, Lamont SJ (2006) Review of quantitative trait loci identified in the chicken. Poult Sci 85: 2079–2096.
20. Ahabat B, Lamont SJ (2007) Genome-wide association analysis reveals cryptic alleles as an important factor in heterosis for fatness in chicken F2 population. Anim Genet 38: 491–498.
21. Liu X, Li H, Wang S, Hu X, Gao Y, et al. (2007) Mapping quantitative trait loci affecting body weight and abdominal fat weight on chicken chromosome one. Poult Sci 86: 1084–1089.
22. Guo L, Sun B, Shang Z, Leng L, Wang Y, et al. (2011) Comparison of adipose tissue cellularity in chicken lines divergently selected for fatness. Poult Sci 90: 2024–2034.
23. Zhang H, Wang SZ, Wang ZP, Da Y, Wang N, et al. (2012) A genome-wide scan of selective sweeps in two broiler chicken lines divergently selected for abdominal fat content. BMC Genomics 13: 794.
24. Ma L, Runestad HB, Dvorak D, Garbe JR, Da Y (2008) Parallel and serial computing tools for testing single-locus and epistatic SNP effects of quantitative traits in genome-wide association studies. BMC Bioinformatics 9: 315.
25. Smoot ME, Ono K, Ruscheinski J, Wang P-L, Ideker T (2011) Cytoscape 2.8: new features for data integration and network visualization. Bioinformatics 27: 431–432.
26. Liviasey AK, Skilling J (1965) Maximum entropy theory. Acta Crystallographica Section A: Foundations of Crystallography 41: 113–122.
27. Verbeek EC, Bakker IM, Beleva MR, Rohnhorst Z, Rizzi P, et al. (2012) A fine-mapping study of 7 top scoring genes from a GWAS for major depressive disorder. PLoS One 7: e37384.
28. Barrett JC, Fry B, Maller J, Daly MJ (2005) Haploview: analysis and visualization of LD and haplotype maps. Bioinformatics 21: 263–265.
29. Zaat O, Hechtner E, Sunyaev SR, Lander ES (2012) The mystery of missing heritability: Genetic interactions create phantom heritability. Nat Rev Genet 13: 199–206.
30. Guo X, Feng C, Ma L, Song G, Wang Y, et al. (2011) Genome-wide association study of body weight in chicken F2 resource population. PLoS One 6: e21872.
31. Wu X, Dong H, Liao L, Zhu Y, Peng G, et al. (2010) A novel statistic for genome-wide interaction analysis. PLoS Genet 6.
32. Khairin R, Wu E (2008) How scale-free are biological networks. Journal of computational biology 15: 810–818.
33. Ma L, Dvorak D, Garbe JR, Da Y (2007) Genome-wide analysis of single-locus and epistasis single-nucleotide polymorphism effects on anti-cyclic citrullinated peptide as a measure of rheumatoid arthritis. BMC Proc 1 Suppl 1: S127.
34. Campos RL, Nones K, Ledur MC, Moura AS, Pinto LF, et al. (2009) Quantitative trait loci associated with fatness in a broiler-layer cross. Anim Genet 40: 729–736.
35. Cui Y, Fu W, Sun K, Romero R, Wu R (2007) Mapping Nucleotide Sequences that Encode Complex Binary Disease Traits with HapMap. Curr Genomics 8: 307–322.
36. Li M, Romero R, Fu WJ, Cui Y (2010) Mapping haplotype-haplotype interactions with adaptive LASSO. BMC Genet 11: 79.
37. Böttcher Y, Unbehauen H, Klotzing N, Ruschke K, Körner A, et al. (2009) Adipose tissue expression and genetic variants of the bone morphogenetic protein receptor IA gene (BMPRIA) are associated with human obesity. Khunet 56: 2119–2120.
38. Weaver RE, Donnelly D, Wabitsch M, Grant PJ, Balabanoff AJ (2008) Functional expression of glucose-dependent insulinotropic polypeptide receptors is coupled to differentiation in a human adipocyte model. Int J Obes (Lond) 32: 1705–1711.
39. Tang X, Metzger D, Leeman S, Amar S (2006) LPS-induced TNF-α factor (LITAF)-deficient mice express reduced LPS-induced cytokine: evidence for LITAF-dependent LPS signaling pathways. Proceedings of the National Academy of Sciences 103: 13777–13782.
40. Winkler G, Koo S, Kezethly I, Saji Z, Ory L, et al. (2005) Expression of tumor necrosis factor (TNF)-alpha protein in the subcutaneous and visceral adipose tissue in correlation with adipocyte cell volume, serum TNF-alpha, soluble...
serum TNF-receptor-2 concentrations and C-peptide level. Eur J Endocrinol 149: 129–135.

41. Schinner S, Willenberg H, Krause D, Schott M, Lamounier-Zepter V, et al. (2007) Adipocyte-derived products induce the transcription of the StAR promoter and stimulate aldosterone and cortisol secretion from adrenocortical cells through the Wnt-signaling pathway. International journal of obesity 31: 864–870.

42. Belgardt BF, Hasch A, Rother E, Ernst MB, Wunderlich FT, et al. (2008) PDK1 deficiency in POMC-expressing cells reveals FOXO1-dependent and-independent pathways in control of energy homeostasis and stress response. Cell metabolism 7: 291–301.

43. Saltiel AR, Pozin JE (2002) Insulin signaling pathways in time and space. Trends in cell biology 12: 65–71.

44. Wunderlich CM, Hovelmeyer N, Wunderlich FT (2013) Mechanisms of chronic JAK-STAT3-SOCS3 signaling in obesity. JAK-STAT 2: 0–1.