A new chicken 55K SNP genotyping array

Ranran Liu1†, Siyuan Xing1,2†, Jie Wang1, Maiqing Zheng1, Huanxian Cui1, Richard P. M. A. Crooijmans2, Qinghe Li1, Guiping Zhao1,3,4* and Jie Wen1,3,4*

Abstract

Background: China has the richest local chicken breeding resources in the world and is the second largest producer of meat-type chickens. Development of a moderate-density SNP array for genetic analysis of chickens and breeding of meat-type chickens taking utility of those resources is urgently needed for conventional farms, breeding industry, and research areas.

Results: Eight representative local breeds or commercial broiler lines with 3 pools of 48 individuals within each breed/line were sequenced and supplied the major SNPs resource. There were 7.09 million - 9.41 million SNPs detected in each breed/line. After filtering using multiple criteria such as preferred incorporation of trait-related SNPs and uniformity of distribution across the genome, 52.18 K SNPs were selected in the final array. It consists of: (i) 19.22 K SNPs from the genomes of yellow-feathered, cyan-shank partridge and white-feathered chickens; (ii) 5.98 K SNPs related to economic traits from the Illumina 60 K SNP Bead Chip, which were found as significant associated SNPs with 15 traits in a Beijing-You crossed Cobb F2 resource population by genome-wide association study analysis; (iii) 7.63 K SNPs from 861 candidate genes of economic traits; (iv) the 0.94 K SNPs related to residual feed intake; and (v) 18.41 K from chicken SNPdb. The polymorphisms of 9 extra local breeds and 3 commercial lines were examined with this array, and 40 K - 47 K SNPs were polymorphic (with minor allele frequency > 0.05) in those breeds. The MDS result showed that those breeds can be clearly distinguished by this newly developed genotyping array.

Conclusions: We successfully developed a 55K genotyping array by using SNPs segregated from typical local breeds and commercial lines. Compared to the existing Affy 600 K and Illumina 60 K arrays, there were 21,41 K new SNPs included on our Affy 55K array. The results of the 55K genotyping data can therefore be imputed to high-density SNPs genotyping data. The array offers a wide range of potential applications such as genomic selection breeding, GWAS of interested traits, and investigation of diversity of different chicken breeds.

Keywords: Chicken, Commercial line, Genotyping array, SNP

Background

With a total of 107 chicken breeds, China has one of the richest local breed resources [1]. This diverse chicken genetic resource is a vital part of the diversity of biological genetic resources around the world and provides excellent material for breeding new varieties or to genetically improve breed.

China is the second-largest broiler producer and consumer all over the world, which accounts for approximately 11% of the chicken production across the globe (FAOSTAT, 2017). In China, chicken is the second largest meat product after pork, making up to 17% of the total meat production. Chicken meat is mainly obtained from the introduced white feather broilers and domestic yellow-feathered meat-type chickens (meat-type local chicken breed, meat-type bred variety and a relevant strain containing the consanguinity of Chinese indigenous chicken), each accounting for half of the consumption. However, the current challenge is how to effectively protect and maintain the existing local varieties. On the other hand, if breeding efficiency is promoted, new chicken lines breeding would be accelerated. The genome-wide SNP chip, also known as SNP array, arranges up to 25 million of DNA marker flanks on glass or special silicon chip to form the SNP probe array. It functions by means of the reaction of base pairing between the chip fixed DNA marker

* Correspondence: zhaoguiping@caas.cn; wenjie@caas.cn
†Ranran Liu and Siyuan Xing contributed equally to this work.
1Institute of Animal Sciences, Chinese Academy of Agricultural Sciences, No. 2 Yuanmingyuan West Road, Beijing 100193, People’s Republic of China
2Institute of Animal Sciences, Chinese Academy of Agricultural Sciences, No. 2 Yuanmingyuan West Road, Beijing 100193, People’s Republic of China
3Institute of Animal Sciences, Chinese Academy of Agricultural Sciences, No. 2 Yuanmingyuan West Road, Beijing 100193, People’s Republic of China
Full list of author information is available at the end of the article
flanks with the target genome, so as to accurately identify the genetic information.

The genotyping arrays have been developed for pig [2], cow [3], dairy cattle [4], sheep [5], salmon [6], and buffalo [7] et al. In chicken, the first 3 K genotyping array was developed in 2005 with 3072 SNPs [8]. After that, in 2008, Groenen et al. did develop a 60 K bead chip for chicken which evenly covered the whole genome [9]. To date, the only available commercial arrays for chicken is Chicken the Affy 600 K SNP Array (Axiom Genome-Wide Chicken Genotyping Array), which was developed by Kranis et al [10]. The other arrays are privately owned by commercial companies. The array supplied an important tool for the genetic diversity analysis, breeds relationship analysis, GWAS, quantitative character positioning analysis of QTL, selective evolution investigation, and Genomic Selection [11]. Up till now the most efficient ways for SNP genotyping, biodiversity measuring, QTL mapping and genomic selection is using SNP arrays. These applications provide improved technical support for the conservation of indigenous breeds and development of new genetic lines/breeds.

One pitfall of all current chicken SNP arrays is the bias towards western commercial lines. The current chicken arrays, however, lack the genomic variation information of Chinese indigenous breeds. Therefore, it is imperative to develop a new type of genome-wide SNP chip with moderate flux in the chicken breeding industry, and also contains the genetic variation information specific to Chinese indigenous breeds.Overlap with the current arrays of the different platforms (Axiom and Illumina) is essential to link the commercial SNP arrays.

Through whole genome re-sequencing of a variety of Chinese native breeds and commercial chicken lines, integrating SNPs associated with economic traits detected in a crossing breed (either indigenous and commercial), a new public available moderate density (55 K) chicken array (IASCHICK) has been developed.

**Results**

The SNPs selection was performed in four groups. The roadmap is shown in Fig. 1, and the establishment of the four groups are indicated in the following paragraphs.

**Genome re-sequencing of chickens supplying the first SNP group**

Eight Chinese local chicken breeds or inbred lines were selected for whole genome sequencing. Each breed/line holds 3 pools of 16 individuals per library without individual barcodes (Table 1). The data summary of each library is provided in the Additional file 1. The number of SNPs per breed/line varied from 7.09 million to 9.41 million SNPs. The average number of detected SNPs was 8.61 M in the local lines, and 7.73 M in the commercial broilers. The total number of SNPs detected overall 8 breeds/lines was 15.2 M. The SNPs with minor allele frequency (MAF) < 0.05 and with low $\Delta F$ were excluded for further steps. The 140 K SNPs, which allelic frequencies distinct to the control breeds, were subsequently used as the first group of candidate SNPs.

| Genome Re-sequencing detected 7.0M-9.4M SNPs from 8 breeds | GWAS of 15 traits | Differentially expressed genes of target traits from candidate genes |
|---------------------------------------------------------------|-------------------|---------------------------------------------------------------|
| 140K SNPs with differentiation in AF between breeds/lines     | 7.42 K SNPs with top 1% p-value | 15.18K SNPs within or near the 861 candidate genes and 0.80 K SNPs for IgY |

Non-recommended SNPs probes in silico validation were excluded

1 or 2 SNPs per 22kb window

- First, 14.39K SNPs in II, III, IV categories were selected
- Second, 19.22K from I category for different breeds were obtained
- Last, 18.41K SNPs from chicken SNPdb to fill the gaps

Fig. 1 The roadmap for the design of the new chicken 55K SNP array.
Selection of the second group of candidate SNPs based on the GWAS of 15 traits

The 7.42 K SNPs were demonstrated to have the top 1% genome-wide significance in 15 traits and were selected as the second group of SNPs. The details are shown in Additional file 2.

Selection of the third group of candidate SNPs based on the genes associated with economic traits

SNPs in the regions of 861 candidate genes related to economic traits were used according to previous studies of gene/protein expression profiles. A total of 66.37 K SNPs in 383 genes for breast muscle and intramuscular fat development in embryonic and post-hatching periods [12], 24.69 K SNPs in 286 genes for body fat metabolism [13], 32.59 K SNPs in 146 genes for disease resistance [14], and 7.24 K SNPs in 46 genes that exhibited possible influence on other chicken economic traits [15] were selected (Additional file 3). The SNPs located in the 5 Kb of either side of the genes’ up- and down-stream were also considered.

According to the SNPs detected by the genome resequencing of the previously mentioned 8 breeds, 15.18 K candidate SNPs were selected from 118.47 K SNPs on all those genes, which had priority with mutations in exons, splicing regions, promotors, and the 3’ and 5’ untranslated regions (UTRs).

In addition, a batch of 798 SNPs from an unpublished capture sequencing of chicken chromosomes 11, 16, and 19 were included in the third candidate group (Additional file 4). The SNPs were significantly related to high IgY levels in Beijing-You and White Leghorn chickens.

There were 15.98 K candidate SNPs that were selected for the design of the final genotyping array.

The fourth group of candidate SNPs are derived from whole genome sequences of low- and high-RFI chickens

Whole genome sequencing of low- and high-RFI chickens were performed to locate the genomic variants for RFI based on differences in allelic frequency between high- and low-RFI chickens as described in our previous study [16]. The selected 4.32 K SNPs (3.74 K RFI related SNP in Beijing-You chickens and 0.58 K RFI related SNPs in Cobb chickens) were used as the candidate SNPs for the design of the final genotyping array in the next step.

Designing the Affy 55K genotyping array

Based on the above four groups of candidate SNPs, a custom-made algorithm was used to fix the final array. Finally, 52,184 SNPs were selected for the final array. The mean physical distance of SNPs in each involved chromosome shows in Table 2. The priority 1 SNPs (the SNPs in group 2, 3 and 4) and 25 INDELs were first placed on the final SNP panel. The next step was addition of the priority 2 SNPs (the SNPs in group 1). The remaining 18.41 K SNPs was selected for the blank windows in the whole chicken genome which the SNPs in the four groups cannot be covered.

The SNPs positions of 55K array were given in Additional file 5. The selected SNPs were derived from the following five groups (Table 3): (i) 19.2 K SNPs from whole genome sequencing of the eight chicken breeds/lines; (ii) 7.42 K trait-related SNPs from the Illumina 60 K SNP Bead Chip, which were found as SNPs significantly associated with 15 economic traits; (iii) 15.98 K SNPs from 861 candidate genes of target traits and high IgY level related region; (iv) 4.32 K SNPs related to chicken RFI; and (v) 18.41 K from chicken SNPdb. In the final genotyping array, 99.85% of SNPs could be annotated (Table 4). The distribution of SNPs on the chromosomes is shown in Fig. 2.

### Table 1

| Type                    | Breeds                              | Individuals in each pool | No. of pools | Number of detected SNPs (with QC ≥ 20) |
|-------------------------|-------------------------------------|--------------------------|--------------|----------------------------------------|
| Local Yellow-feathered chicken | Beijing-You                         | 16                       | 3            | 8,505,214                              |
|                         | Jingxing-Huang                      | 16                       | 3            | 8,349,627                              |
|                         | Sanhuang                            | 16                       | 3            | 9,405,319                              |
| Cyan shank partridge    | Cyan shank partridge (fast growth rate) | 16                       | 3            | 8,954,795                              |
|                         | Cyan shank partridge (mediate growth rate) | 16                       | 3            | 8,884,232                              |
| Commercial white-feathered | Cobb maternal line                  | 16                       | 3            | 7,093,225                              |
|                         | Cobb paternal line                  | 16                       | 3            | 8,372,769                              |
|                         | Recessive White                     | 16                       | 3            | 7,556,464                              |
| **Total**               |                                     |                          |              | 15,312,402                             |

*Each pool contained 8 males and 8 females

Based on Gallus_gallus-4.0
The comparisons of the Affy 55K array with the existing chicken arrays (Affy 600 K array, and Illumina 60 K)
All the SNPs of this 55K array, Affy 600 K array [10], and Illumina 60 K array [9] were mapped to the latest chicken genome (GRCg6a). The overlap of the 3 arrays is shown in Fig. 3. There are 6740 SNPs (13%) which overlap between the Affy 55K array and the Illumina 60 K array. When comparing to the Affy 600 K array, there are 24,227 SNPs that overlap between the 55K array which accounts for 46%. There were 21,412 new SNPs included in 55K array compared to the existing arrays.

Validation of the 55K array in 13 chicken breeds/lines
All samples from 10 Chinese local breeds (Chahua, Dagu, Liyang, Luhua, Qingyuan, Silkie, Wenchang, Bai’er, Xianju, and Jingxing-Huang) and 3 commercial lines (Hubbard, Cobb, and White Leghorn) were genotyped.

The average call rate for each breed ranged from 97.0% (Qingyuan) to 98.7% (Cobb). Across all populations, 76.7 to 88.0% of the 52,184 SNPs were polymorphic, with MAF ≥ 0.05. The average MAF ranged from 0.22 (Bai’er chicken) to 0.27 (Wenchang chicken) (Table 5).

An MDS analysis was performed using the genotyped data to investigate the ability of the 55K panel to detect population stratification in the validated samples. Figure 4 shows the relative coordinates of individuals when plotted using the two largest principal components. Individuals originating from the commercial broilers, Hubbard and Cobb tightly clustered. The Chinese indigenous meat-type breeds clustered together. The two Chinese indigenous egg-type breeds, Xianju and Bai’er, clustered together. The remaining local breeds (mainly

| Table 2 | The number of SNPs of the 55K array on each chromosome and their distance* |
|---------|---------------------------------|
| Chromosome | Number of SNPs | Mean Distance (Kbp) |
| 1 | 10,228 | 19.21 |
| 2 | 7077 | 21.14 |
| 3 | 5196 | 21.44 |
| 4 | 4589 | 19.91 |
| 5 | 2705 | 22.10 |
| 6 | 1750 | 20.36 |
| 7 | 1684 | 21.63 |
| 8 | 1314 | 22.81 |
| 9 | 1236 | 18.98 |
| 10 | 1399 | 14.59 |
| 11 | 1373 | 14.67 |
| 12 | 1389 | 14.34 |
| 13 | 1041 | 17.67 |
| 14 | 1118 | 14.43 |
| 15 | 761 | 16.71 |
| 16 | 81 | 7.37 |
| 17 | 724 | 14.31 |
| 18 | 736 | 14.94 |
| 19 | 725 | 13.74 |
| 20 | 867 | 16.04 |
| 21 | 503 | 13.53 |
| 22 | 153 | 30.00 |
| 23 | 321 | 17.76 |
| 24 | 390 | 15.98 |
| 25 | 106 | 26.67 |
| 26 | 339 | 15.36 |
| 27 | 277 | 20.22 |
| 28 | 317 | 15.91 |
| Z | 3785 | 21.67 |
| Summary | 52,184 |

* The distance between SNPs based on Gallus_gallus-5.0

| Table 3 | The number of SNPs from five candidate groups in the final 55K array |
|---------|---------------------------------|
| Resource Category | Number of SNPs in 55K array |
| I. Genome Re-sequencing of eight breeds | 
| White-feathered | 12,555 |
| Yellow-feathered | 3040 |
| Cyan-shank Partridge Chicken | 2724 |
| II. SNPs based on GWAS of 15 traits | 5980 |
| III. SNPs on the candidate genes | 7630 |
| IV. SNPs related to RFI | 943 |
| V. SNPs from chicken SNPdb | 18,412 |
| Total | 52,184 |

| Table 4 | Summary of the SNPs effect prediction in 55K array |
|---------|---------------------------------|
| Item | Count | Percent (%) |
| Total number of SNPs in the panel | 52,184 |
| Annotation possible | 52,108 | 99.85 |
| intergenic variant | 16,106 | 30.86 |
| intron variant | 25,275 | 48.43 |
| intron variant & noncoding transcript variant | 3981 | 7.63 |
| missense variant | 590 | 1.13 |
| missense variant & splice region variant | 13 | 0.02 |
| synonymous variant | 1601 | 3.07 |
| Splicing | 187 | 0.36 |
| start/stop gained/lost/retained | 12 | 0.02 |
| 3 prime UTR variant | 1358 | 2.60 |
| 5 prime UTR variant | 229 | 0.44 |
| upstream gene variant (1 kb) | 871 | 1.67 |
| downstream gene variant (1 kb) | 1014 | 1.94 |
| noncoding transcript exon variant | 871 | 1.67 |

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characteristic of meat-types) were located relatively close to each other compared to egg-type breeds and commercial broilers. The commercial layer White Leghorn chickens were placed relative far away from the Chinese local breeds and commercial broilers in Fig. 4.

The linkage disequilibrium (LD) in Jingxing-Huang chicken and Cobb paternal line chicken were calculated, respectively. Figure 5 a and b shows the LD decay of the Jingxing-Huang and Cobb paternal lines for chromosome 1 and 2, respectively. The average levels of LD between adjacent SNPs in Jingxing-Huang breeds of chromosome 1 is 0.61 and for chromosome 2 is 0.58, whereas in the Cobb line these LD levels are 0.56 and 0.46, respectively. The mean LD level decay is around 0.22 in 40 Kb. The r² of LD in the Jingxing-Huang chickens are larger than that in Cobb paternal line. Additional file 6 and Additional file 7 showed the r² of LD decreasing with the increased SNPs distance in the two populations in whole genome level.

**Discussion**

The 52.18 K SNPs selection was performed in four groups and NCBI SNPdb using several criteria. The first column of Table 5 shows the number of polymorphic loci in local breeds and introduced lines.

**Table 5** Number of polymorphic loci in local breeds and introduced lines

| Breeds          | Average Call Rate (%) | Polymorphic loci | Mean MAF ³ |
|-----------------|-----------------------|------------------|------------|
| Chahua          | 97.56                 | 42.3 K           | 0.242      |
| Qingyuan        | 97.00                 | 45.2 K           | 0.267      |
| Wenchang        | 97.10                 | 46.5 K           | 0.277      |
| Luhua           | 97.21                 | 44.4 K           | 0.261      |
| Liyang          | 97.23                 | 40.4 K           | 0.229      |
| Dagu            | 97.46                 | 43.7 K           | 0.253      |
| Bai’er          | 97.36                 | 40.0 K           | 0.222      |
| Xianju          | 97.13                 | 40.1 K           | 0.235      |
| Silkie          | 97.03                 | 45.0 K           | 0.258      |
| Hubbard         | 97.88                 | 46.1 K           | 0.269      |
| Cobb            | 98.70                 | 45.2 K           | 0.237      |
| White Leghorn   | 97.99                 | 43.5 K           | 0.249      |

⁴MAF > 0.05.
⁵Across all 52.2 K loci.
Fig. 4 Results of multidimensional scaling analysis of 12 breeds/lines. The scatters show the individuals’ position in the MDS plot, different colors represent different breeds/lines.

Fig. 5 The LD decay plots. (a) from the Cobb and Jingxing-Huang (JXH) chickens in chromosome 1; (b) from Cobb and JXH chickens in chromosome 2.
The linkage disequilibrium (LD) in the Jingxing-Huang breeds and Cobb paternal line were calculated and compared. The mean LD level decay to around 0.22 in 40 Kb. This result is similar with the result of Fu et al. in 2015 [21]. The \( r^2 \) of LD in the Jingxing-Huang breed is larger than that in Cobb paternal line. The Jingxing-Huang is an inbred line with a
relatively small effective population size whereas the Cobb paternal line is three to four times larger.

In China, indigenous yellow-feathered chickens are highly diverse (more than 100 local breeds and 70 crosses). The major obstacle in applying genomic selection for improvement of local breeds is the cost of genotyping array. The 55K array has a medium SNPs density, cost-efficient, and optimal for Chinese local breeds compared with the existing 600 K commercial array. Furthermore, the 55K genotyping array incorporated known SNPs loci that possess a high potential for association with economic traits and traits that are expensive and difficult to measure, which will be interesting for both GWAS and genomic selection (GS) projects.

With the rapid development of next-generation sequencing technologies and reduction of the costs, genotyping with re-sequencing (IBS) will be the focus of future research. In the current phase, however, the IBS system is more complex and not as solid as the SNP array. The array genotyped data can be easily analyzed and standardized according to constant array SNP positions. The batch effect can be excluded by different laboratories and companies.

Conclusions
In conclusion, we developed Affy 55K genotyping array that was designed to use SNPs that are segregated in Chinese local chicken breeds and commercial lines/breeds, and where large number of SNPs are associated with economic traits. Compared to the existing Affy 600 K and Illumina 60 K arrays, 21,41 K new SNPs were included in the 55K SNP array. The results from the our Affy 55K genotyping array can be imputed to the high-density SNPs genotyping data. This array offers wide range of potential applications, such as the evaluation of germplasm resources of chicken breeds, investigation of diversity of different chicken breeds, implementation of genome-wide association studies and genomic selection.

Methods
Animals
For whole genome sequencing, the 384 chickens were sampled from eight local breeds or inbred lines (Table 1). Chickens were supplied by Institute of Animal Sciences in CAAS (local breed Beijing-You, inbred Jingxing-Huang line), Jiangsu Lihua Co. Ltd. (Cyan-shank Partridge lines with fast and mediate growth rates, respectively), Institute of Poultry Sciences of CAAS (Sanhuang chicken and Recessive White chicken), Xinguang Nongmu Co. Ltd. (paternal lines in parent generation from Cobb and Hubbard), the Institute of Animal Sciences of CAAS (White Leghorn). Two groups with 87 and 100 chickens from Jingxing-Huang and Cobb were also used for SNP array evaluation. The blood samples used in this study were all collected from chickens under the veterinary supervision and the Guidelines for Experimental Animals established by the Ministry of Science and Technology (Beijing, China), and with the approval of Animal Ethics Committee of the Institute of Animal Sciences. No anesthesia or euthanasia methods were used. There was no evidence at health examination that any of the involved chickens had clinical diseases caused by the sampling.

Whole genome re-sequencing
Genomic DNA was isolated from blood samples by the phenol-chloroform method. Samples DNA quality were validated by gel electrophoresis and Nanophotometer. The individual DNA samples (48 from each breed/line) were pooled to construct three libraries, with each library containing 8 males and 8 females. The libraries were constructed using the Nextera DNA Library Preparation Kit (Illumina Inc., San Diego, CA) according to the manufacturer's standard protocol. All libraries were sequenced on the Illumina Hiseq2500 (2 × 125 bp).

Genome sequence alignment and detection of the first group of candidate SNPs
Reads were filtered for low quality (> 10 consecutive nucleotides with Phred scores < 10), adaptor sequences, and sequences without a quality control-passed paired read using NGSSCQ toolkit (v2.3.3) [22]. Each trimmed pool sequencing coverage are shown in Table S5. Filtered sequenced reads were mapped to the reference genome (Gallus_gallus_4.0) by BWA software (v0.7.10) [23]. PCR duplications were removed with -rmdup argument in Samtools (version 0.1.1.18) [24]. SNPs were identified and genotyped for each data set with mpileup function in Samtools, then called by VarScan [25]. Only those highly confident variants supported by both methods were kept for downstream analyses. The SNPs calling details parameter were described by Liu et al [16]. The SNPs with MAF < 0.05 and the INDELs in each breed/line were filtered by vcf tools [26]. In Beijing-you chicken, Jingxing-Huang chicken, Sanhuang chicken, and the two lines of cyan-shank partridges minus the MAFs of Cobb paternal line, as well as the MAFs of Recessive White chicken, and the paternal and Cobb paternal line is three to four times larger.
maternal generation of Cobb minus the MAFs of Beijing-You chicken, respectively. The SNPs with low ΔF were excluded. The value of ΔF was adjusted for 140 K SNPs reserved in local breeds and commercial lines to generate the first group of candidate SNPs. The threshold of ΔF in local breeds and commercial lines are 0.609 and 0.731, respectively. The SNPs acquired through genome re-sequencing of eight breeds/lines supplied the major data for the first group of SNPs in the array. SNPs specific for chromosome W were removed and were not considered in current designing. There are also 25 INDELs for special interest, which were defined as priority.

Selection of the second group of candidate SNPs based on GWAS analysis of 15 traits
The second group of candidate SNPs was selected according to a GWAS analysis of 15 traits. Phenotype and genotype data were generated from the CAAS chicken F2 resource population as described in Sun’s report [27]. Briefly, the population was derived from a cross between local Beijing-You chickens and commercial Cobb broilers (Cobb-Vantress, Inc.). The weight, carcass, immune and meat quality traits were measured from 367 F2 chickens. The 15 traits were as follows, (a.) body weight of day 28 and day 42, (b.) carcass traits including total weight percentage after slaughtering, breast muscle weight percentage, leg muscle weight percentage, abdominal fat percentage, (c.) meat quality traits including the breast muscle intramuscular fat ratio, ultimate pH (24 h), meat lightness, redness value and yellowness value of breast muscle, (d.) immune traits including IgY level to sheep red blood cell, the heterophil and lymphocyte ratio, IgY level in serum, and the average red blood cell backlog.

SNPs were genotyped by using Illumina 60 K SNP Bead chip for chicken [9]. All description of the phenotypes had been reported by Sun et al. in 2013 [27]. To maximize the polymorphism resources for SNP array, the GLM procedures were used for the GWAS analysis and was performed by PLINK software (version 1.07) [28] with 42,585 SNPs passed quality control. The details were described by Sun et al. [27]. The SNPs with top 1% lowest p-values were used in the following procedures.

Selection of the third group of candidate SNP based on the associated genes for target traits
Known candidate genes for economic traits were collected and used for the SNP array design. All genes were identified through previous researches by our group [12, 13, 29, 30]. We retrieved total 861 genes related to skeletal muscle and intramuscular fat development, chicken fat metabolism, salmonella enteritidis resistance etc. (Additional file 2). The SNPs were annotated by the Ensembl tool VEP [31]. Mutations and the SNPs in the exons, splicing region, and UTRs were firstly selected out. A maximum of 5 candidate SNPs were selected out for each gene.

In addition, the SNPs in this group also included a batch SNPs detected from a set of capture sequencing of Chr. 11, Chr. 16, and Chr. 19 of White Leghorns and Beijing-You chickens with low or high serum IgY (Liu et al., unpublished, Supplement Table S3).

Selection of the fourth group of candidate SNPs for RFI
The fourth group candidate SNPs were selected from a whole genomic re-sequencing research of low- and high-RFI Cobb and Beijing-You chickens. SNPs calling results showed that 8,505,214 and 8,479,041 single nucleotide polymorphisms (SNPs) were detected in low- and high-RFI Beijing-You chickens, respectively; 8,352,008 and 8,372,769 SNPs were detected in low- and high-RFI Cobb chickens, respectively. The SNPs with Fst value < 5% in each breed were excluded followed by SNPs with mean ΔF < 0.35 between low- and high-RFI chickens. Through the above filtering processes, 3.74 K SNPs assigned to 1137 candidate genes in Beijing-You chickens and 0.58 K SNPs (448 genes) in Cobb chickens were reserved [16].

Selection of the SNPs from chicken SNPs database
The first four groups cannot cover the whole genome evenly. In the fifth group, SNPs were selected from chicken SNPs database from NCBI (ftp://ftp.ncbi.nih.gov/snp/organisms/archive/chicken_9031/).

SNP screening according to the scoring of probes
All the SNPs’ positions were transformed from WASHUC2.1 (Illumina 60 K), and Gallus_gallus-4.0 (Affy 600 K) to Gallus_gallus-5.0 (Affy 55 K) by the Lift-Over tool on UCSC Genome Browser. Take utility of all SNPs from the five candidate groups above, in silico validation, was performed using the AxiomGTv1 algorithm of APT, which generated an output score file containing p-convert values, signifying the SNP array quality and list of recommended and non-recommended SNP probes. For a high-quality SNP array design, non-recommended SNP probes were all excluded in the following procedure.

SNPs selection procedure for the final 55K array
The final SNPs selection was done in multiple steps using several criteria. The roadmap is shown in Fig. 1.

A custom-made algorithm was applied as described below. According to the Gallus_gallus-5.0, the chicken genome length is about 1.2 Gb. To ensure the probe position evenly distributed in the chicken genome, the whole genome was distributed by windows with 22 Kb
length. The backward window started from the probe position of the forward probe position. The selection of the final array was performed on each chromosome separately. The first four groups SNPs were divided as 2 priorities. The SNPs in group 2, group 3, group 4, and the INDELs in group 1 were defined as priority 1, and the SNPs in group 1 were defined as priority 2.

1. a) The selection of the SNPs in priority 1. If there is no SNP in a 22 kb window, the window will be reserved. b) If there are one or two SNPs in the window, the SNP(s) was reserved. c) If there are 3 or more SNPs in a window, only 2 SNPs in this window will be reserved, which can make the SNPs even distributed in this window according to the following formula. SD² = \[\frac{1}{4} (S - x)² + (N_i - x)² + (N_j - x)² + (E - x)²\]. In the formula above, the S and E are the start position and the end position of the window respectively; and N_i and N_j are the target SNPs position in the window. The SNPs N_i and N_j which can minimum the SD², will be reserved.

2. The selection of priority 2 SNPs. The windows reserved 1 or 2 SNPs will be skipped. The windows without SNP will be filled by one SNP of priority 2 according to the formula described above.

3. The windows without any SNP will be filled by 1 SNP from the NCBI SNPdb of chicken, while the validated SNPs will have a priority for filling.

The final array contains 55K probes for 52 K SNPs, which were manufactured by Affymetrix® using photolithography. The redundant probes are used for interrogating each SNPs [32, 33]. The final 52 K SNPs were annotated by the online tool Ensembl VEP [34].

The comparisons of the 55K Affy array with the existing arrays (Affy 600 K array, and Illumina 60 K)
All the SNPs’ positions were transformed from WASHUC2.1 (Illumina 60 K), Gallus_gallus-4.0 (Affy 600 K) and Gallus_gallus-5.0 (Affy 55 K) to GRCh38a by the LiftOver tool on UCSC Genome Browser. All the SNP positions of the three genotyping arrays were compared. The SNPs on 600 K array and 60 K array were also performed by Ensembl VEP [31]. Overlapping Venn plot was performed by the Calculate and draw custom Venn diagrams website (http://bioinformatics.psb.ugent.be/webtools/Venn/).

Validation of the 55K array in 13 chicken breeds/lines
The genomic DNA from 12 breeds/lines (Chahua, Dagu, Liyang, Luhua, Qingyuan, Silkie, Wenchang, Bai’er, and Xianju, Hubbard, Cobb, and White Leghorn) and two lines with larger populations (Jingxing-Huang and Cobb) were isolated as mentioned above. The genotyping was done on Axiom® arrays using the Affymetrix® GeneTitan® system according to the procedure described by Affymetrix (https://assets.thermofisher.com/TFS-Assets/LSG/manuals/702899_PI.pdf) in the Beijing Compass Biotechnology Co., Ltd. (Beijing, China).

Basic genotype statistics for each marker, including call rate, MAF, Hardy-Weinberg Equilibrium (HWE), allele and genotype counts were calculated using the Quality Assurance Module from the SNP Variation Suite version 7 (SVS; Golden Helix Inc., Bozeman, Montana: www.goldenhelix.com). The following quality control criteria (filtering) were used to remove SNPs with less than 95% call rate for further analysis. The SNPs with less than 0.05 MAF. SNPs were tested for HWE (P < 0.001) to identify possible typing error. Samples with more than 10% missing genotypes were removed from the study.

The MDS was performed using the genotype data of the SNPs from the 55K panel on all the breeds samples (n = 226) to assess the utility of the panel in detecting population structure. Population structure between 12 breeds was carried out using PLINK software (version 1.90b3) [28] with the MDS method on, and the plot was performed by ggplot2 [35]. The linkage disequilibrium in 2 populations were performed by the GAPIT [36]. The LD decay plot performed by PopLDdecay software are presented as whole genome levels and as chromosome levels with the parameter of smaller break point size of 5 Kb and bigger break point size of 40 Kb [37].

Additional files

Additional file 1: The summary of each sequencing pools. The raw reads number, clean reads number, sequencing depth, Q30 percentage, and coverage et al. for each sequencing library were provided. (XLSX 14 kb)

Additional file 2: The second group of SNPs which related to the economic traits. The locus and the p-value of SNPs which related to 15 economic traits were provided. (XLSX 998 kb)

Additional file 3: The third group of SNPs which related to 861 candidate genes. The information of 861 candidate genes and 118.4 K SNPs selected were provided. (XLSX 6645 kb)

Additional file 4: The third group of SNPs which related to serum IgY. The loci and allele information of 0.8 K SNPs related to serum IgY were provided. (XLSX 32 kb)

Additional file 5: The loci information for the 55K array. The loci, allele information, SNPs frequencies in each breed/line, and genotype counts were calculated using the Quality Assurance Module from the SNP Variation Suite version 7 (SVS; Golden Helix Inc., Bozeman, Montana: www.goldenhelix.com). The following quality control criteria (filtering) were used to remove SNPs with less than 95% call rate for further analysis. The SNPs with less than 0.05 MAF. SNPs were tested for HWE (P < 0.001) to identify possible typing error. Samples with more than 10% missing genotypes were removed from the study.

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Additional file 5: The loci information for the 55K array. The loci, allele information, SNPs frequencies in each breed/line, and overlap information were provided. (XLSX 7761 kb)

Additional file 6: The LD decay in whole genome level in Cobb population. (JPEG 563 kb)

Additional file 7: The LD decay in whole genome level in Jingxing-Huang population. (JPEG 578 kb)

Abbreviations
CAAS: Chinese Academy of Agricultural Sciences; Chr: Chromosome; Da: inter-population net nucleotide divergence; GLM: general liner model; GWAS: Genome-Wide Association Study; HWE: Hardy-Weinberg Equilibrium; INDEL: insert/deletion.; LD: Linkage Disequilibrium; MAF: Minor Allele
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Availability of data and materials
The whole genome sequencing clean data reported in this paper have been deposited in the Genome Sequence Archive [38] in BIG Data Center [39] under accession number CRA001289 which can be publicly accessed at http://bigd.big.ac.cn/gsa.

Authors' contributions
RL contributed to the design and performing of the study, the analysis and interpretation of data and writing of the manuscript; SX contributed to the design and performing of the study, the analysis and interpretation of data were supported by the earmarked fund for the modern agro-industry technology research system (CARS-41) and Agricultural Science and Technology Innovation Program (ASTIP-IASOM; ASTIP-IAS-TS-15); the interpretation of data and writing the manuscript were supported by the National Nonprofit Institute Research Grant (2017ywf-zd-2).

Ethics approval and consent to participate
All experimental procedures with chickens were performed according to the Guidelines for Experimental Animals established by the Ministry of Science and Technology (Beijing, China). Ethical approval on animal survival was given by the animal ethics committee of the Institute of Animal Sciences and Technology (Beijing, China). Ethical approval on animal survival was given by the animal ethics committee of the Institute of Animal Sciences and Technology (Beijing, China). Ethical approval on animal survival was given by the animal ethics committee of the Institute of Animal Sciences and Technology (Beijing, China).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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Author details
1Institute of Animal Sciences, Chinese Academy of Agricultural Sciences, No. 2 Yuanmingyuan West Road, Beijing 100193, People's Republic of China.
2Animal Breeding and Genomics, Wageningen University & Research, Wageningen, The Netherlands.
3State Key Laboratory of Animal Nutrition, Ministry of Agriculture and Rural Affairs, Beijing 100193, People's Republic of China.
4Key Laboratory of Animal (Poultry) Genetics Breeding and Reproduction, Ministry of Agriculture and Rural Affairs, Beijing 100193, People's Republic of China.

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References
1. Resources CNCoAG: animal genetic resources in China: poultry: China agriculture press; 2011.
2. Liu et al. BMC Genomics (2019) 20:410
3. Polotsky YV, Wright WE, Wittenberg B. Evolution of the vertebrate genome with implications for the origin of the human genome. Annu Rev Genet. 1994;28:109–31.
4. Liu H, Durbin R. Fast and accurate read alignment with burrows-wheeler transform. Bioinformatics. 2009;25(1):1754–60.
5. R Development Core Team. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. 2018-04-03.
6. Madsen B, Nordborg M. Genome-wide association studies and whole-genome selection. Nat Rev Genet. 2009;10(4):259–68.
7. Patil NG. Large-scale association mapping of quantitative traits. Nature. 2006;441(7091):616–20.
8. O'Connell JR, Purcell SM. Genotyping arrays and the genetic architecture of complex traits. Nat Rev Genet. 2010;11(11):765–74.
9. Johnson WE, Livak KJ, Quinlan AR, Schierup MH, Blumenstiel L. Genotype dosage calls with PLINK. Nat Genet. 2006;38(9):968–9.
10. Zhang Y,软件开发团队. Analyzing next-generation sequencing data: tools and strategies. Brief Bioinform. 2009;10(2):217–30.
11. Liu Y, Tian J, Wang Y, Hu Y, Yang F, Zhang L, et al. Identification of candidate genes for the chicken slow growth phenotype by using whole-genome resequencing. G3. 2015;5(10):2119–26.
25. Koboldt DC, Chen K, Wylie T, Larson DE, McLellan MD, Mardis ER, Weinstock GM, Wilton RK, Li D. VarScan: variant detection in massively parallel sequencing of individual and pooled samples. Bioinformatics. 2009;25(17):2283.

26. Petr D, Adam A, Goncalo A, Albers CA, Eric B, Depristo MA, Handsaker RE, Gerton L, Marth GT, Sherry ST. The variant call format and VCFtools. Bioinformatics. 2011;27(15):2156–8.

27. Sun Y, Zhao G, Liu R, Zheng M, Hu Y, Wu D, Zhang L, Li P, Wen J. The identification of 14 new genes for meat quality traits in chicken using a genome-wide association study. BMC Genomics. 2013;14(1):458.

28. Purcell S, Neale B, Todd-Brown K, Thomas L, Ferreira MAR, Bender D, Maller J, Sklar P, Bakker P, Daly MJ. PLINK: a tool set for whole-genome association and population-based linkage analyses. Am J Hum Genet. 2007; 81(3):556–75.

29. Liu J, Fu R, Liu R, Zhao G, Zheng M, Cui H, Li Q, Song J, Wang J, Wen J. Protein profiles for muscle development and intramuscular fat accumulation at different post-hatching ages in chickens. PLoS One. 2016;11(8):e0159722.

30. Cui HX, Liu RR, Zhao GP, Zheng MQ, Chen J, Wen J. Identification of differentially expressed genes and pathways for intramuscular fat deposition in pectoralis major tissues of fast- and slow-growing chickens. BMC Genomics. 2012;13(1):213.

31. McLaren W, Gil I, Hunt SE, Riat HS, Ritchie GR, Thorman A, Flicek P, Cunningham F. The Ensembl variant effect predictor. Genome Biol. 2016;17(1):122.

32. Gunderson K, Steemers F, G, Mendoza L, Chee M. A genome-wide scalable SNP genotyping assay using microarray technology. Nat Genet. 2005; 37(5):S49–554.

33. Syv, Auml AC, nen. Toward genome-wide SNP genotyping. Nat Genet. 2005; 37 Suppl:37 Suppl55.

34. Zerbino DR, Achuthan P, Akanni W, Amode MR, Barrell D, Bhai J, Billis K, Cummins C, Gall A, Giron CG, et al. Ensembl 2018. Nucleic Acids Res. 2018; 46(D1):D754–61.

35. Wickham H. ggplot2: Elegant Graphics for Data Analysis: Springer Publishing Company, Incorporated, 2009.

36. Alexander EL, Feng T, Qishan W, Jason P, Meng L, Peter JB, Michael AG, Edward SB, Zhiwu Z. GAPIT: genome association and prediction integrated tool. Bioinformatics. 2012;28(18):2397.

37. Zhang C, Dong SS, Xu JF, He WM, Yang TL. PopLDdecay: a fast and effective tool for linkage disequilibrium decay analysis based on variant call format files. Bioinformatics. 2017;33(15):2156–64.

38. Wang Y, Song F, Zhu J, Zhang S, Yang Y, Chen T, Tang B, Dong L, Nan D, Qian Z. GSA: genome sequence archive. Genomics Proteomics & Bioinformatics. 2017;15(1):14–8.

39. Members BDC. Database resources of the BIC data center in 2018. Nucleic Acids Res. 2018;46(Database issue):D14–20.