Association of Apolipoprotein B and Adiponectin Receptor 1 Genes with Carcass, Bone Integrity and Performance Traits in a Paternal Broiler Line

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Abstract

Apolipoprotein B (APOB) and Adiponectin Receptor 1 (ADIPOR1) are related to the regulation of feed intake, fat metabolism and protein deposition and are candidate genes for genomic studies in birds. In this study, associations of two single nucleotide polymorphisms (SNPs) g.102A>T (APOB) and g.729C>T (ADIPOR1) with carcass, bone integrity and performance traits in broilers were investigated. Genotyping was performed on a paternal line of 1,454 broilers. The SNP detection was carried out by PCR-RFLP technique using the restriction enzymes HhaI for the SNP g.729C>T and MslI for the SNP g.102A>T. The association analyses of the two SNPs with 85 traits were performed using the restricted maximum likelihood (REML) and Generalized Quasi-Likelihood Score (GQLS) methods. For REML the model included the random additive genetic effect of animal and fixed effects of sex, hatch and SNP genotypes. In the GQLS method, a logistic regression was used to associate the genotypes with phenotypes adjusted for fixed effects of sex and hatch. The SNP g.729C>T in the ADIPOR1 gene was associated with thickness of the femur and breast skin yield. Thus, the ADIPOR1 gene seems implicated in the metabolism and/or fat deposition and bone integrity in broilers.

Introduction

The chicken (Gallus gallus) was one of the first domestic animals sequenced and its genome assembly was completed in 2004 [1] by the Washington University Genome Sequence Center and the National Human Genome Research Institute of the United States of America. Gallus gallus genome has 38 pairs of autosomal chromosomes (5 macrochromosomes, 5 intermediates...
and 28 microchromosomes) and one pair of sex chromosomes, Z and W [2], with a total of 1.2 Gbp. At 2.5–21 cM/Mb, depending on the chromosome, recombination rates are higher in chickens than in humans or rats, which average 1 cM/Mb and 0.5 cM/Mb respectively [3]. This, plus the great diversity among breeds and strains, has made Gallus gallus an animal model for studying the genetic basis of phenotypic traits, capitalizing on the possibility of detecting more segregation than in other species.

Many traits of economic interest are complex and determined by a number of unknown genes. Loci that control quantitative traits (QTL) have been associated with a number of traits in chickens, including growth, body composition, egg production, antibody production, disease resistance, and behavior. However, determining causative genes of phenotypic variation is difficult, since each locus controls only a fraction of the phenotypic variance of a given trait [3].

Among the molecular markers, single nucleotide polymorphisms (SNPs) are promising for association studies. These markers are abundant and exhibit low mutation rates, which facilitates genotyping [4]. Many studies have been done on the association between SNPs in candidate genes and metabolic pathway in several species [5–10]. Wong et al. [11] identified 2.8 million SNPs in the chicken genome. This abundance of available SNPs may aid in the mapping of causative polymorphisms underlying complex traits in chickens in the future.

Studies on Apolipoprotein B have identified its role in lipid metabolism. This glycoprotein plays an important part in the absorption, assembly and secretion of lipids, including triglycerides and cholesterol. In chickens, Apolipoprotein B is the main component of IDL (intermediate density lipoprotein) and VLDL (very low density lipoproteins) present in plasma cholesterol [12]. In rats and mice, Apolipoprotein B was found in several forms and with diverse functions at different ages by Mcleod et al. [13], who also reported that Apolipoprotein B48, a protein only half the size of Apolipoprotein B100, could mediate both assembly and secretion of chylomicrons (in the intestine) and VLDL (in the liver). In chickens, the Apolipoprotein B (APOB) gene may be related to early body growth rates and fat deposition [14].

Another candidate gene associated with fat metabolism in chickens is Adiponectin (ADIPO). Adiponectin exerts its action by binding to two specific receptors, ADIPOR1 and ADIPOR2. Both ADIPOR1 and ADIPOR2 are seven transmembrane receptors that are structurally and functionally distinct from G-protein-coupled receptors [15]. Adiponectin receptor 1, encoded by the ADIPOR1 gene, is a major adiponectin receptor that mediates the glucose and lipid metabolism-related effects of adiponectin on target cells [16]. Research based on animal models has shown that ADIPOR1 overexpression can increase the biological effects of adiponectin [17].

There is evidence that visceral adiposity or age influences the adiponectin plasma levels in chickens. According to Kershaw and Flier [18], the biological effects of adiponectin depend not only on the relative concentrations in circulation and the properties of the different adiponectin subtypes, but also the specific expression of the subtypes on their respective tissue receptors. Maddineni et al. [19] confirmed their presence in skeletal muscle, diencéphalon, pituitary gland, liver, ovary and kidney. In addition, ADIPO gene may be associated with the initiation and growth processes of adipose tissue deposition in chickens [20, 21].

There are several studies investigating metabolic and genetic mechanisms regulating fat deposition in chickens [22]. Broilers are genetically selected for body weight and muscle growth. However, this genetic selection also leads to an unwanted increase in visceral adiposity [23, 24]. In human studies, fat has been associated to bone disorders [25] and cardiovascular diseases [26] and there is evidence of the interconnection of both these metabolic problems in modern broiler production.

Given the genetic selection process that modern broilers are subject to, it is important to understand both the direct and indirect influence of APOB and ADIPOR1 genes on the
development of birds. Therefore, the object of this study was to investigate the association of
the SNPs in the APOB and ADIPOR1 genes with carcass, bone integrity and performance traits
in broilers. In addition, two different statistical methodologies were used in order to confirm
possible associations.

**Material and Methods**

**Ethics Statement**

This study was performed with the approval of the Embrapa Swine and Poultry Ethical Com-
mittee for Animal Use (CEUA) under protocol number 011/201, following international guide-
lines for animal welfare.

**Population and the collection of data**

Phenotypic records were obtained from 1,454 animals from a paternal lineage of broilers. This
line has been developed and is owned by the Poultry Genetic Improvement Program from
EMBRAPA Swine and Poultry (Brazilian Agricultural Research Corporation, [https://www.
embrapa.br/en/home](https://www.embrapa.br/en/home)). The experimental research centre is located in the city of Concórdia
(27° 14' 03" S—52° 01' 40" W), Santa Catarina state, Brazil. This line of broilers has been under
development since 1992 and aims to: increase body weight and carcass yield; improve viability,
fertility, hatchability, feed conversion, and reduce abdominal fat. The original population was
randomly sampled and then 20 males and 100 females were mated to produce an initial popu-
lation of approximately 1,500 animals.

The birds were housed collectively until 35 days of age and then, in order to evaluate feed
conversion, were moved to individual cages from 35 to 41 days. The birds were banded for
identification, and fed a three-phase diet; starter from the 1st to 21st day (21% crude protein
and 3,150 kcal metabolizable energy), grower from the 22nd to 34th day (20% protein gross and
3,200 kcal metabolizable energy) and finisher from the 35th to 41st day (18.5% crude protein
and 3,200 kcal metabolizable energy).

The birds were slaughtered at 42 days of age following a 6 hour fasting. Cervical dislocation
was manually applied before bleeding of the neck. Blood samples were taken for DNA extrac-
tion during bleeding. Approximately 2 mL of blood was collected in microtubes with 100 mL
(10% v/v) of 0.5 M EDTA anticoagulant. The samples were immediately packed in ice and then
stored in a freezer at -20°C.

Eighty-five traits related to performance, carcass composition, organs and bone integrity
were evaluated (Tables 1 and 2). Yields were expressed as a percentage of the dressed weight
relative to the 42 days live weight. An arcsin transformation was applied to the yield traits to
normalize theirs distributions. Feed conversion ratio was calculated by dividing feed intake by
weight gain in the period from 35 to 41 days of age.

After bleeding, the animals were scalded in a hot water bath (60°C for 45 s) and the feathers
removed mechanically. The carcass weight was calculated by removing the feathers, blood,
head, feet, and organs, except the lungs and kidneys. The carcass cuts (breast, drumsticks and
thighs with bones and without skin, wings, neck, and back, which corresponds to the dorsal
portion of the carcass) were individually weighed. Weight of muscle cuts, separated from skin
and bones, and weight of skin with fat were also recorded.

Tibia and femur length were measured between the distal and proximal ends, and thickness
in the central region of the bones. Both traits were measured with manual caliper (0.01 mm).
The bones were kept at about 0°C for 48 hours and then left at room temperature for about an
hour to determine flexural strength. The bending test was performed on a TA—XT Plus Tex-
ture Analyzer (Texture Technologies Corporation), using the probe TA-92 (Texture
Technologies Corporation). The bones were placed in the same position, with the ends resting on two supports spaced 30 mm. The probe travelled 20 mm after touching the sample at 2.00 mm/s test and pretest speed, and 20.00 mm/s post-test speed. The probe touched the sample with programmed weight force of 500 mg. This force was applied to the central area (diaphysis) to determine the flexural strength and rupture modulus. After breaking, the bone fragments were placed in plastic bags, labeled with the sample number and stored at 0°C for 24 hours to determine dry matter and ash.

Table 1. Carcass traits and corresponding abbreviations used for weight and yields recorded in the broiler line.

| Carcass Traits                                      | Abbreviation | Weight | Yield |
|-----------------------------------------------------|--------------|--------|-------|
| Weight post bleeding and plucking                   | WPBP         | —      | YBP   |
| Weight of blood and feathers                        | WBF          | YBF    |       |
| Chilled carcass weight                              | WCC          | YCC    |       |
| Weight of abdominal fat                             | WAF          | YAF    |       |
| Weight of head                                      | WHD          | YHD    |       |
| Weight of feet                                      | WFT          | YFT    |       |
| Weight of liver                                     | WLI          | YLI    |       |
| Weight of heart                                     | WHT          | YHT    |       |
| Weight of gizzard                                   | WGZ          | YGZ    |       |
| Weight of wing                                      | WW           | YWW    |       |
| Weight of wing drummettes                           | WWDD         | YWWD   |       |
| Weight of wing middles                              | WM           | YWM    |       |
| Weight of wing tips                                 | WWT          | YWT    |       |
| Weight of chilled tibia                             | WCT          | YCT    |       |
| Weight of thigh                                     | WTH          | YTH    |       |
| Weight of thigh skin                                | WTHS         | YTHS   |       |
| Weight of thigh meat                                | WTHM         | YTHM   |       |
| Weight of drumstick                                 | WDS          | YDS    |       |
| Weight of drumstick skin                            | WDSS         | YDSS   |       |
| Weight of drumstick meat                            | WDSM         | YDSM   |       |
| Weight of thigh and drumstick meat                  | WTHDSM       | YTHDSM |       |
| Weight of breast                                    | WBT          | YBT    |       |
| Weight of breast skin                               | WBTS         | YBTS   |       |
| Weight of breast meat                               | WBTM         | YBTM   |       |
| Weight of breast fillet                             | WBTF         | YBTF   |       |
| Weight of breast bone                               | WBTB         | YBTB   |       |
| Weight of back                                      | WBAC         | YBAC   |       |
| Weight of neck                                      | WNEC         | YNEC   |       |
| Weight of lungs                                     | WLNG         | YLNG   |       |
| Weight of tibia                                     | W_TIB        | YW_TIB |       |
| Weight of the femur                                 | W_FEM        | YW_FEM |       |

Weight in grams and yield in %.

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To determine percent dry matter, the bone fragments were left at room temperature for about one hour. Subsequently, they were placed in pre-weighed porcelain crucible and kept in an oven at 105°C for 16 hours, and then, placed in a desiccator until they reach room temperature and
weighed. The dry matter percentage was determined as the ratio of the dry weight to the wet weight of the samples.

Ash was determined immediately after obtaining the dry matter, using the bone fragments. The samples were incinerated in a muffle for six hours. The initial temperature was 350°C for one hour and increased gradually to 450°C and 550°C for one hour and, finally, to 600°C for three hours. Then, the crucibles with the samples were left in the desiccator until reaching room temperature and then weighed. The ash percentage was determined by dividing ash weight by the dry weight of the sample.

Extraction and quantification of DNA

DNA extraction was performed using the DNAzol reagent (Invitrogen) according to manufacturer recommendations. After extraction, the DNA was resuspended in 150 μL ultrapure water, incubated at 37°C for one hour in a water-bath and stored in a freezer at -20°C. A spectrophotometer (Eppendorf BioPhotometer) was used to quantitate the DNA and then, the samples were diluted with ultrapure water to the concentration of 25 ng/μL.

Table 2. Bone integrity and bird performance traits recorded in the broiler line and corresponding abbreviations used.

| Trait                                           | Abbreviation |
|------------------------------------------------|--------------|
| Bone integrity traits                           |              |
| Tibia length                                    | LG_TIB       |
| Thickness of the tibia                          | THK_TIB      |
| Femur length                                    | LNG_FEM      |
| Thickness of the femur                         | THK_FEM      |
| Curvature of the femur                         | SCORE        |
| Force required to break the femur              | FEM_FORCE    |
| Length of the femoral break                    | FEM_LNG      |
| Area of the femoral break                      | FEM_AREA     |
| Force required to break the tibia              | TIB_FORCE    |
| Length of the tibial break                     | TIB_LNG      |
| Break area of the tibia                        | TIB_AREA     |
| Dry matter content of the femur                | FEM_DM       |
| Dry matter content of the tibia                | TIB_DM       |
| Ash content of the femur                       | FEM_ASH      |
| Ash content of the tibia                        | TIB_ASH      |
| Performance traits                              |              |
| Weight at hatching                             | WHTC         |
| Weight at 21 days of age                       | W21          |
| Weight at 35 days of age                       | W35          |
| Weight at 41 days of age                       | W41          |
| Weight at 42 days of age                       | W42          |
| Feed intake 35 to 41 days of age               | FI35_41      |
| Weight gain 35 to 41 days of age               | WG35_41      |
| Feed conversion 35 to 41 days of age           | FC35_41      |

Weight in grams; thickness, distance and length in centimeters; force in kgf/mm².

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Gene amplification and genotyping

Specific regions of the APOB and ADIPOR1 genes were amplified with primers for the identification of SNPs. One pair of primers, including regions of exons and introns, was designed for each gene (Table 3). To identify SNPs, gene fragments were sequenced on ABI 3130XL (Applied Biosystems) and subsequently analyzed with Phred/Phrap/Consed/PolyPhred software [27–29]. One SNP per gene was chosen to be genotyped. The SNPs were selected based on the highest sequencing quality and the highest informativeness, i.e. the most polymorphic ones. The PCR protocols for amplification of DNA fragments were standardized as described in Table 3.

To amplify DNA fragment of APOB gene, a PCR reaction was performed in 25 μL containing 1X reaction buffer, 2.5 mM of MgCl₂, 0.4 mM dNTPs, 0.24 μM of each primer, 1.5 U of Taq DNA Polymerase enzyme (Invitrogen, San Diego, CA), and 50 ng of genomic DNA, and finally adjusting the volume to 25 μL by adding ultrapure water. A similar reaction was used for the primer ADIPOR1, but 1.6 mM MgCl₂ was used instead. The PCR reactions were carried out in BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA) using a touchdown PCR protocol under the following condition: denaturing at 94°C for 5 minutes, followed by 35 cycles of denaturing at 94°C for 1 minute, annealing at 59°C for 1 minute and extension at 72°C for 1 minute. The final extension was performed at 72°C for 10 minutes.

Electrophoresis using 1% agarose gel was done to visualize the final product.

After PCR, a cleavage reaction was performed with specific restriction enzymes. The NEB-cutter program [30] was used to identify the restriction sites and specific enzymes for the selected SNP. Enzyme MslI was selected for the evaluation of the SNPs in fragment amplified by the primers for APOB, whereas enzyme HhaI was selected for analysis of the SNP in the fragment amplified by the primers for ADIPOR1.

The cleavage protocol for each enzyme was:

MslI: Each reaction used 2 μL of the PCR product with 1 μL of 10X buffer; 1U enzyme MslI, and ultrapure water was added to reach 10 μL. The thermal cycler conditions were: one cycle at 65°C for 3.5 hours, followed by one cycle at 80°C for 20 minutes.

HhaI: Each reaction used 2 μL of PCR product in 2 μL of 10X buffer; 1U enzyme HhaI, and ultrapure water added to a final volume of 10 μL. The reaction occurred in a water bath at a temperature of 37°C overnight.

Finally, the digestion reaction was visualized using a 2% agarose gel stained with 0.01% ethidium bromide. The samples and molecular markers (100 bp DNA Ladder—Promega Corporation) were loaded on the gel and subjected to electric current at 100 V for about a half hour in 1X TBE (Tris-Borate-EDTA). A UV transilluminator was used to compare the sample band pattern to the 100 bp marker bands.

Both phenotypic and genotypic data used in this study are available upon request to Dr. Mônica Côrreia Ledur (Embrapa Swine and Poultry. Address: Rodovia BR-153, Km 110, Distrito de Tamanduá Caixa Postal: 21 CEP: 89700–000, Concórdia, Santa Catarina, Brazil).

| Primers | Primer Sequence | GenBank accession number | Chromosome (position) | Amplicon |
|---------|----------------|-------------------------|-----------------------|----------|
| APOB-F  | 5’-CTGCCAAAGACCTTGTGTGGGTTCT-3’ | NC_006090.3 | GGA3 (101882031–101916285 bp) | 1,099 bp |
| APOB-R  | 5’-TCTGTGAGGCCTGTAACAGTCA-3’ | | | |
| ADIPOR1-F | 5’-CCATGCCACACAAAATGTTGGTTCCT-3’ | NC_006113.3 | GGA26 (1088478–1094154 bp) | 1,023 bp |
| ADIPOR1-R | 5’-TGATGTGACTGGAACTGCAGGGA-3’ | | | |

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Statistical analyses

Preliminary analyses, including descriptive statistics, normality test and testing of fixed effects to be included in the models, were done using the **UNIVARIATE** and **GLM** procedures in the Statistical Analysis System (SAS) software [31]. Outliers were identified using Box-plot and were removed from the data set. The descriptive statistics of the studied traits are in Table 4. Genotypic and allelic frequencies were calculated by counting the genotypes and alleles for each SNP and Hardy-Weinberg equilibrium was tested using a chi-square test at 5% significance level.

The association of SNPs with the traits was investigated using two different methods: Restricted maximum likelihood via ASREML [32] and Generalized quasi-likelihood scoring (GQLS; [33]).

**Restricted maximum likelihood method.** The ASREML software was used to fit the following univariate linear model for all the traits, which included additive and dominant effects of the SNP markers:

\[
y_{ijk} = \mu + \text{sex}_j + \text{inc}_k + \sum_{l=1}^{2} \beta_1 x_{1l} + \sum_{l=1}^{2} \beta_2 x_{2l} + a_i + e_{ijk},
\]

where:
- \(y_{ijk}\) is observation for the trait on the \(i^{th}\) animal,
- \(\mu\) is the overall mean,
- \(\text{sex}_j\) is the fixed effect of the \(j^{th}\) sex (\(j = 1, 2\)),
- \(\text{inc}_k\) is the fixed effect of the \(k^{th}\) hatch (\(k = 1, 2, 3, 4, 5\))
- \(x_{1l}, x_{2l}\) are the recoded genotypes (-1, 0, or 1) and (0, 1 or 0) for the \(l^{th}\) SNP, i.e. genotypes AA, AT and TT of the **APOB** gene and CC, CT and TT of the **ADIPOR1** gene, respectively,
- \(\beta_{1l}, \beta_{2l}\) are the linear regression coefficients (additive and dominance effects, respectively) for the \(l^{th}\) SNP,
- \(a_i\) is the random additive genetic (polygenic) effect of the \(i^{th}\) animal,
- and \(e_{ijk}\) is the residual random effect for the \(i^{th}\) animal.

An alternate model was also fit using ASREML to directly estimate the allele substitution effect of the SNPs:

\[
y_{ijk} = \mu + \text{sex}_j + \text{inc}_k + \sum_{l=1}^{2} \beta_i x_l + a_i + e_{ijk},
\]

where:
- \(y_{ijk}\) is observation for the trait on the \(i^{th}\) animal,
- \(\mu\) is the overall mean,
- \(\text{sex}_j\) is the fixed effect of the \(j^{th}\) sex (\(j = 1, 2\)),
- \(\text{inc}_k\) is the fixed effect of the \(k^{th}\) hatch (\(k = 1, 2, 3, 4, 5\))
- \(x_l\) is the number of copies of a given allele in the genotype of the \(l^{th}\) SNP (counted as -1, 0, 1) for genotypes AA, AT and TT of the **APOB** gene and for genotypes CC, CT and TT of the **ADIPOR1** gene, respectively,
- \(\beta_i\) is the linear regression coefficient (allele substitution effect) for the \(i^{th}\) SNP,
- \(a_i\) is the random additive genetic (polygenic) effect of the \(i^{th}\) animal,
- and \(e_{ijk}\) is the residual random effect for the \(i^{th}\) animal.

All the available pedigree information (1,567 animals) was used to build the numerator relationship matrix for the animal additive genetic effect.
Table 4. Descriptive statistics for the studied traits.

| Trait | N   | Mean | SD  | CV (%) | Minimum | Maximum |
|-------|-----|------|-----|--------|---------|---------|
| WHTC  | 1448| 47.66| 3.70| 7.77   | 37.40   | 61.80   |
| W21   | 1426| 648.43| 133.91| 20.65 | 256.00  | 1034.00 |
| W35   | 1450| 1730.96| 202.59| 11.70 | 776.00  | 2444.00 |
| W41   | 1443| 2219.20| 251.91| 11.35 | 1026.00 | 2922.00 |
| W42   | 1452| 2223.86| 260.24| 11.70 | 988.00  | 2971.00 |
| F35_41| 1443| 1091.45| 152.48| 13.97 | 508.00  | 1590.00 |
| WG35_41| 1439| 488.77| 106.57| 21.80 | 128.00  | 802.00  |
| FC35_41| 1439| 2.31 | 0.47| 20.16 | 1.42    | 5.25    |
| WPBP  | 1445| 2055.89| 247.42| 12.03 | 901.00  | 2764.00 |
| WBF   | 1441| 168.55| 26.37| 15.65 | 82.00   | 293.00  |
| WCC   | 1436| 1639.06| 202.05| 12.33 | 661.20  | 2212.00 |
| WAF   | 1435| 47.36 | 14.08| 29.73 | 2.50    | 94.00   |
| WHD   | 1428| 52.55 | 7.31| 13.91 | 27.90   | 77.20   |
| WFT   | 1424| 74.66 | 13.71| 18.36 | 34.30   | 110.10  |
| WLI   | 1422| 52.34 | 8.73| 16.68 | 25.40   | 82.40   |
| WHT   | 1421| 12.40 | 2.15| 17.35 | 6.30    | 19.70   |
| WGZ   | 1423| 32.00 | 6.04| 18.86 | 17.80   | 56.10   |
| WW    | 1422| 167.40| 20.00| 11.95 | 72.00   | 236.40  |
| WWD   | 1432| 85.71 | 11.17| 13.03 | 35.80   | 121.00  |
| WWM   | 1425| 61.91 | 7.98| 12.89 | 26.20   | 87.80   |
| WWT   | 1433| 19.78 | 2.94| 14.87 | 10.00   | 31.80   |
| WCT   | 1432| 55.65 | 10.24| 18.40 | 25.40   | 85.40   |
| WDS   | 1421| 205.87| 31.24| 15.17 | 86.20   | 306.60  |
| WDSS  | 1427| 17.30 | 4.38| 25.31 | 5.40    | 36.60   |
| WDSM  | 1429| 32.51 | 5.57| 17.14 | 18.00   | 56.60   |
| WTH   | 1430| 46.23 | 9.77| 21.14 | 11.60   | 87.60   |
| WTHS  | 1427| 310.49| 46.11| 14.85 | 113.60  | 464.40  |
| WTHM  | 1431| 231.75| 37.78| 16.30 | 81.40   | 351.20  |
| WBT   | 1426| 500.76| 63.48| 12.68 | 211.30  | 710.80  |
| WBTB  | 1431| 31.38 | 6.78| 21.60 | 8.40    | 61.70   |
| WBTM  | 1431| 294.07| 42.66| 14.51 | 123.70  | 428.60  |
| WBTF  | 1434| 77.55 | 11.85| 15.28 | 35.70   | 119.10  |
| WBTB  | 1435| 97.91 | 15.02| 15.34 | 43.50   | 152.70  |
| WUBAC | 1425| 263.95| 35.84| 13.58 | 120.40  | 370.70  |
| WNEC  | 1432| 119.37| 21.52| 18.02 | 35.60   | 200.40  |
| WLNG  | 1430| 15.31 | 3.06| 19.98 | 6.60    | 24.60   |
| WTHDS | 1414| 516.41| 73.98| 14.33 | 200.20  | 736.60  |
| WSTHM | 1426| 364.69| 55.91| 15.33 | 137.20  | 529.40  |
| W_TIB | 673 | 11.75 | 2.11| 17.99 | 7.14    | 17.82   |
| W_FEM | 672 | 8.51  | 1.37| 16.08 | 4.41    | 12.68   |
| LG_TIB| 673 | 95.29 | 3.86| 4.05  | 80.00   | 106.20  |
| THK_TIB| 673 | 8.76  | 0.92| 10.51 | 6.00    | 12.90   |
| THK_FEM| 673 | 8.86  | 0.73| 8.26  | 7.00    | 11.43   |
| LNG_FEM| 672 | 69.56 | 3.09| 4.45  | 59.30   | 78.00   |
| SCORE | 673 | 1.13  | 0.34| 29.94 | 1.00    | 2.00    |
| FEM_FORCE| 672 | 28.69 | 5.78| 20.15 | 13.45   | 52.68   |

(Continued)
The Generalized Quasi-Likelihood Score (GQLS) method developed by Feng et al. [33], which uses a logistic regression model to associate the genotypes (treated as the response variable) with the traits phenotypes adjusted for fixed effects (sex and hatching) was also used as an alternative statistical method. In the GQLS method, a logistic regression is used to associate the expected frequency of a marker allele ($\mu_j$) with the trait phenotypes ($X_i$):

$$\mu_j = E(Y_j|X_i) = \frac{e^{\beta_0 + \beta_1 X_i}}{1 + e^{\beta_0 + \beta_1 X_i}},$$

where:
- $\beta_0$ is the intercept and $\beta_1$ is the linear regression coefficient,
- $\mu_j$ is expected allele frequency of the $j^{th}$ SNP,
- $Y_j$ is proportion of the allele 1 in the observed genotype of the $i^{th}$ animal for $j^{th}$ SNP,
- $X_i$ is the phenotypic observation of the $i^{th}$ animal.

The GQLS method accounts for the relationships among animals through the numerator relationship matrix [33].

The significance of statistical tests needs to be adjusted for multiple hypotheses tests, such as in the case of a large number of traits tested in this study. Adjustments can be made by the Bonferroni correction [34]. However, Bonferroni adjustment assumes that all tests are independent. A principal component analysis of the correlation matrix among the 85 traits analyzed in this study was performed using the PRINCOMP procedure in SAS software [31]. The first 31 principal components explained 99% of the total variance. The comparison-wise significance level for Bonferroni adjustment was then $\alpha/31$, where 31 was considered the effective number of independent tests [35]. An experimental(trait)-wise significant level of 5% was considered to declare a significant association, while a 15% level was considered as a trend for association.

### Results

The two SNPs were found segregating with intermediate allele frequencies and showed high heterozygosity in the broiler line considered (Table 5). The heterozygote frequency was 57.4% (762 animals) for the SNP in the APOB gene and 56.5% (763 animals) for the SNP in the ADIPOR1 gene. Test for Hardy-Weinberg equilibrium showed an excess of heterozygotes.
(P<0.01), indicating that the population was not in Hardy-Weinberg equilibrium. For the APOB SNP, the T allele was more frequent than the A allele, while for the ADIPOR1 SNP, the C allele was more frequent than the T allele (Table 5).

The traits WAF, YWW, YBAC, WNEC, YBTB, and YNEC were not significantly affected by sex (P>0.05) and the traits LG_TIB, LNG_FEM, THK_FEM and TIB_ASH were not significantly affected by hatch (P>0.05). All other traits were significantly affected by sex and hatch (P<0.05).

Tables 6 and 7 shows the results for traits for which at least a trend of association with SNPs in APOB and ADIPOR1 genes was found (P<0.10) at a comparison-wise level. These Tables also present results adjusted for multiple tests. After Bonferroni correction, only the SNP in ADIPOR1 gene was significantly associated (P<0.05) with two traits (YBTS and THK_FEM) and showed suggestive association (P<0.15) with WBTS and FEM_FORCE using the REML method (Table 6).

According to the analysis using the GQLS method, no significant associations with the SNP in APOB gene were found too. The SNP in the ADIPOR1 gene remained associated with THAK_FEM, but also was significantly associated with FEM_FORCE (P<0.05), another bone integrity trait (Table 7).

Even though not significantly at an experimental-wise level, several associations were found at a comparison wise level of 5%, including associations of the SNP in ADIPOR1 gene with TIB_FORCE, WDSS and YDSS (Tables 6 and 7).

**Discussion**

Sex had no significant effect on abdominal fat, although Mignon-Grasteau et al. [36] reported that body weight selection contributes to an increase of sexual dimorphism. Fat depots are greater in females than in males, but this is caused mainly by the larger adipocytes found in females [37]. Differences between sexes for abdominal fat at 28, 35, 42 and 49 days were observed by Dalanezi et al. [38].

There were no reports in literature of sex effect on YWW, YBAC, WNEC, YBTB, and YNEC; and hatch effect on LG_TIB, LNG_FEM, THK_FEM and TIB_ASH traits.

The SNP in the ADIPOR1 gene showed a significant additive effect on YBTS and THK_FEM, suggesting that this polymorphism may have a direct influence on fat deposition in the breast and bone integrity. This SNP showed an opposite additive effect on breast skin and in thickness of the femur (Table 6). This indicates that skin fat deposition in carcass cuts (breast) and bone quality may be altered in opposite directions if selection is conducted based on this SNP marker, what potentially might be beneficial, i.e. selecting for the favourable allele for bone integrity would lead to decrease in skin fat deposition. The suggestive associations (P<0.15) of SNP in the ADIPOR1 gene with FEM_FORCE and WBTS also suggest that this gene may directly influence on fat deposition and bone integrity.

**Table 5. Marker, number of animals (N), genotypic and allelic frequencies for the SNPs studied in the broiler line.** Allelic and genotypic frequencies, percentages are between parentheses.

| Marker       | N  | Genotype | Allele |
|--------------|----|----------|--------|
| APOB g.102A>T SNP | 1,328 | AA | AT | TT | A | T |
|              |     | 188(14.1) | 762(57.4) | 378(28.5) | 42.9 | 57.1 |
| ADIPOR1 g.729C>T SNP | 1,351 | CC | CT | TT | C | T |
|              |     | 342(25.3) | 763(56.5) | 246(18.2) | 53.6 | 46.4 |

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Even though not significantly at an experimental-wise level associations of the SNP in the *ADIPOR1* gene with force required to break the tibia (TIB_FORCE), weight and yield of drumstick skin (WDSS and YDSS) were observed, which corroborate with the significant associations found with yield of breast skin (YBTS) and thickness of the femur (THK_FEM) and suggestive associations with weight of breast skin (WBTS) and force required to break the femur (FEM_FORCE).

No significant SNP association with live weights at different ages and with weights of various cuts was found. However, Hendricks et al. [20] found that young chickens (4 weeks old) had a set amount of plasma adiponectin and that age-related changes or fast growth could lead to the decline of circulating adiponectin levels.

Pisto et al. [39] reported that reduction in plasma adiponectin concentration is indicative of an increase in muscle fiber size. Fast-growing broilers have larger muscle fibers [40] and, therefore, have a smaller amount of adiponectin in the blood. This may be the result of muscle
glucose uptake and nutrient deposition. Increasing dietary energy increases adiponectin gene expression in abdominal adipose tissue at 32 days of age, but there is no associated increase at 49 days of age [21]. This may explain the absence of an additive effect of SNP g.729C>T on abdominal fat, if at 42 days of age the adiponectin concentration is lower, then it cannot have an effect on the deposition and/or metabolism of abdominal adipose tissue.

Studies in other species have observed relationships between adiponectin, adipose tissue and bone [41]. The exact way adiponectin works is not yet fully understood. However, it is known that other adipokines, such as leptin, which has a positive correlation with adipose deposition and metabolism, inhibits adiponectin production [25]. According to some authors, such as Sinsigalli et al. [42], Barbato et al. [24] and Gonzales et al. [43], the deposition of subcutaneous and abdominal fat and skeletal disorders in broilers are associated with varying concentrations of hormones. Furthermore, genes such as APOB and ADIPOR1 and the neural control mechanisms of satiety and hunger that regulate feed intake also influence those traits.

The lack of association between ADIPOR1 and abdominal fat deposition may be due to its expression occurs at an early age, as reported in the literature [14, 20, 21] and, in this study, fat deposition was measured at 42 days of age. Another reason for the non association of ADIPOR1 with abdominal fat may be due to the fact that this gene influences the deposition of different fatty acids present in breast and drumstick fat. According Crespo and Garcia [44], abdominal fat contains more oleic acid, while the breast and thigh fat contains more stearic acid.

While the results of this study provided essential information and found that the SNP in the ADIPOR1 gene was significantly associated with bone integrity and fat deposition in the breast its role in the chicken’s metabolism needs to be further studied before it can be used for genetic screening. The role of the adiponectin gene on metabolism must be better understood, as the intense selection pressure for increased body weight affects this hormone. Excess of adipose tissue increases the production of many adipokines that influence several bodily functions in birds.

Table 7. P-values for associations of SNPs in the APOB and ADIPOR1 genes with broiler traits, using a Generalized Quasi-Likelihood Score method (traits are defined in Tables 1 and 2).

| Trait      | SNP g.102A>T (APOB gene) | SNP g.729C>T (ADIPOR1 gene) |
|------------|---------------------------|-----------------------------|
| THK_FEM    | 0.679005                  | 0.00014**                   |
| FEM_FORCE  | 0.784031                  | 0.00161**                   |
| YBTS       | 0.73904                   | 0.0023*                     |
| WBTS       | 0.2934                    | 0.0056                      |
| TIB_FORCE  | 0.3811                    | 0.0161                      |
| WDSS       | 0.0867                    | 0.0337                      |
| YDSS       | 0.4914                    | 0.0500                      |
| WHT        | 0.3863                    | 0.0603                      |
| YHT        | 0.1591                    | 0.0617                      |
| YAF        | 0.1706                    | 0.0732                      |
| WAF        | 0.9691                    | 0.0885                      |
| FC3S_41    | 0.3437                    | 0.0895                      |
| YNEC       | 0.2455                    | 0.0898                      |
| YFT        | 0.0408                    | 0.8276                      |

**Significance at 5% experiment-wise level by Bonferroni correction.
*A trend of significance at 15% experiment-wise level by Bonferroni correction.

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The SNP in the ADIPOR1 gene is located in an intron region of the gene. According to Ninov et al. [45], when a SNP occurs in the intron region, it may not be involved directly with the associated trait, but may be connected to other polymorphisms located in or around the coding and regulatory regions of the gene. This needs to be further investigated.

Conclusions
The SNP g.729C>T in the ADIPOR1 gene was found associated with thickness of the femur and breast skin yield. Thus, the ADIPOR1 gene seems implicated in the metabolism and/or fat deposition and bone integrity in broilers. Further studies are warranted to elucidate if this SNP could be used as molecular genetic markers in broiler selection.

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Author Contributions
Conceived and designed the experiments: MCL JOP. Performed the experiments: JOP AMGI. Analyzed the data: VARC. Contributed reagents/materials/analysis tools: MS FSS. Wrote the paper: VARC. Results interpretation and manuscript revision: VARC DPM MS FSS MCL JOP AMGI RPS NBS NVG.

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