Morphometric Traits, Serum Chemistry and Antibody Response of Three Chicken Genotypes under Free-Range, Semi-Intensive and Intensive Housing Systems

ABSTRACT

The present study evaluated the effect of housing system on the morphometrics, serum chemistry and antibody response of dual-purpose chicken genotypes. A total of 156 pullets and 39 cockerels were randomly picked from 18 treatment block groups (3 housing system × 3 genotypes × 2 sexes) according to Randomized Complete Block Design (RCBD). Three genotypes, purebred Naked Neck (NN) and two crossbred Rhode Island Red × Naked Neck (RIR × NN = RNN) and Black Australorp × Naked Neck (BAL × NN = BNN), were compared. Morphometric traits were recorded during rearing period, thereafter, serum chemistry and antibody response were evaluated in pullets. Intensive and semi-intensive chickens were heavier (males, \( p = 0.0012 \); females, \( p < 0.0001 \)) on week 21. Body length was maximum (\( p < 0.0001 \)) for free-range female chicken. Maximum (\( p < 0.0001 \)) keel length was found in semi-intensive female chickens. Regarding genotypes, RNN and BNN chickens were heavier than NN (males, \( p = 0.0015 \); females, \( p < 0.0001 \)). Keel length was maximum (\( p = 0.0002 \)) in BNN and NN female chickens. Drumstick circumference were maximum (males, \( p < 0.0001 \); females, \( p < 0.0001 \)) in NN chickens, shank circumference was maximum (\( p = 0.0150 \)) in RNN and BNN male chickens. Wingspan was maximum (\( p = 0.0029 \)) in NN female chickens. Plasma glucose level was higher (\( p = 0.0008 \)) in intensive female chickens whereas cholesterol levels was higher (\( p = 0.0123 \)) in NN female chicken. Antibody titer against ND was higher (\( p = 0.0204 \)) in RNN female chickens while higher (\( p = 0.0001 \)) antibody titer against IB was found in free-range chickens. Overall, housing system did not impact morphometric traits or serum chemistry. Only a few differences were observed regarding body weight, body and keel length, plasma glucose, cholesterol and antibody response against ND and IB.

INTRODUCTION

Crossbreeding is an effective tool for the development of modern-day commercial chickens and equally important for the improvement of rural chickens (Sheridan, 1981). There are different types of crossbreeding comprising two-way, three-way, and four-way rotational crosses or back crosses. Crossbreeding also maximizes the expression of hybrid vigor, improves fitness characteristics that are generally reflected in the resultant cross. Three-way or four-way crosses has to be employed in order to retain the heterosis in material traits (Hoffmann, 2005). In general, crossbreeding involves a two-way cross between an exotic breed and a local one. The aim of these crosses is to combine the characteristics of both genotypes and produce individuals that are more productive, have higher resistance to disease and better adapted to harsh climatic conditions than the parent genotypes (Khawaja et al., 2013).
**MATERIALS AND METHODS**

This study was conducted under practical conditions at Indigenous Chicken Resource Centre (ICGRC), Department of Poultry Production, University of Veterinary and Animal Science (UVAS), Ravi Campus, A-Block, Pattoki, Pakistan. Pattoki is located at 31°1’0N, 73°50’60E and at an altitude of 186 m (610 ft). This city normally experiences hot and humid tropical climate, with maximum temperatures ranging from 13°C in winter and 43°C in summer.

Despite having enormous potential, limited research work has been conducted for the improvement of indigenous chickens in developing countries. Some attempts have been made to improve the productive of indigenous chickens by crossbreeding or upgrading with known exotics breeds and then leaving the offspring to natural selection (Njenga, 2005). In Pakistan, a dual-purpose chicken genotype was developed by adopting four-way crossbreeding programs in which local chicken (desi = non-descript) was crossed with three exotic breeds: White Cornish, New Hampshire and White Leghorn. The resultant breed, named Lyallpur Silver Black (LSB), was developed that have better productive performance and livability in harsh climatic conditions (Siddiqi et al., 1979).

Blood biochemical profile is generally considered as an ideal indicator of health status, and frequently applied by avian pathologists to determine birds’ immune status and to obtain basic knowledge on specific poultry diseases. Regarding blood chemistry, total serum protein is useful to draw inferences about the quality of dietary protein (Bonadiman et al., 2009; Alikwe et al., 2010). Likewise, triglyceride and glucose level indicate the energy requirements for physiological responses and allow proper body biochemical functions (Kral & Suchy, 2000).

In order to understand infection outcomes and bird’s performance, the knowledge of the immune response is essential. In this regard, indigenous poultry may be the most efficient model to study the immune response against bacterial and viral infections (Haunshi et al., 2011). There are still limited data on the maternal effects or reference values of distinct crosses. The lack of reference serum chemistry levels and antibody response against diseases motivates scientists to establish these references for particular crossbreds. Therefore, present study aimed at investigating if there are differences in morphometric traits, serum chemistry and antibody response in dual-purpose chicken genotypes reared in free-range, semi-intensive or intensive systems.

**Ethics**

Birds’ care and use of bird were in accordance with the laws and regulations of Pakistan, and the experimental procedures were approved by the Committee of Ethical Handling of Experimental Birds (No. DR/124), UVAS, Pakistan.

**Experimental Birds**

Four hundred and eighty one-day-old chicks hatched at Avian Research and Training (ART) Centre, UVAS, Lahore, Pakistan, were transported to ICGRC, A-Block, UVAS, Ravi Campus, Pattoki, Pakistan for evaluation. Chicks belong to the genotypes (160 birds each): Rhode Island Red × Naked Neck (RNN) crossbreds, Black Australorp × Naked Neck (BNN) crossbreds, and Naked Neck × Naked Neck (NN) purebreds.

Chicks were brooded in well-ventilated open-sided house, and submitted to standard management practices until six weeks old (June to July, 2018.) Birds were fed a commercial broiler breeder diet (16% crude protein, 2900 kcal metabolizable energy/kg). During the brooding period, birds were vaccinated against Newcastle disease and infectious bronchitis, according to local schedule of area.

At 6 weeks of age, 60 (30 males and 30 females) from each genotype (RNN, BNN and NN) were transferred to three housing systems (free-range, semi-intensive or intensive), totaling 360 birds (3 genotypes × 3 housing systems × 2 sexes × 20 birds = 360). Weekly body weight and behavioral repertoires were recorded for the duration of 10 weeks (6 to 16 weeks).

At 16 weeks of age, out of the 260 birds (156 female and 104 male), 52 pullets and 13 cockerels from each genotype were used in rearing phase (17 to 21 weeks). For this, 156 females and 39 males were randomly picked from 18 treatment groups (3 genotypes × 3 housing systems × 2 sexes) according to Randomized Complete Block Design (RCBD). Furthermore, males were reared separately.

**Free-Range, Semi-intensive and Intensive Systems**

All the experimental birds were individually tagged. In the free-range and semi-intensive systems, birds were kept in open sided shed (6.1 m L × 6.1 m W × 3.66 m H) oriented east to west. A range area of fertile land (10 m L × 2.99 W, at a stocking density of 0.23m² / bird) located in front of the shed was used. Free range area enriched with grasses and platsns (Mung (Vigna radiate L.), Black eyed Pea (Vigna unguiculata L.), French Pea (Phaseolus vulgaris L.) and Lucerne
The birds under intensive system were maintained in well-ventilated poultry shed equipped with three-tiered battery cage system (FACCO, Poultry Equipment-C3, Italy), during rearing phase, 17 cages were used comprising four birds each; 0.14 m²/bird floor space was provided. Birds were offered a broiler breeder developer diet formulated according to the recommendations of the NRC (1994) (Table 2) and daily feed allowance was increased corresponding to their growth and requirement (Table 3).

### Parameters Studied

#### Morphometric traits

Morphometric traits were weekly measured, including body weight, beak length, drumstick length, shank length, drumstick circumference, shank circumference, body length and wing spread.

#### Serum Chemistry and Antibody Response

At the end of the experiment (21 weeks of age), 3 mL of blood were collected from the brachial wing vein of three females per treatment using a syringe with anticoagulant. After blood centrifugation, the serum was collected in Eppendorf tubes and stored at -15°C to -20°C until analyses (Gunes et al., 2002). Serum was analyzed for albumin, globulin, uric acid, glucose, total protein, creatinine and cholesterol contents using serum analysis kits (Kumar and Kumbhakar, 2015). One week prior to slaughter, birds were vaccinated against Newcastle Disease and Infectious Bronchitis and antibody titer were evaluated at the end of experimentation (Xie et al., 2008).

### Statistical Analysis

Collected data regarding morphometric traits, serum chemistry and antibody response were analyzed by two-way analysis of variance assuming
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Morphometric traits differed among genotypes, housing systems and their interaction (Tables 4, 5, 6).

Intensive and semi-intensive reared chickens were heavier (males, $p=0.0012$; females, $p=0.00001$) than free-range chickens. Regarding genotypes, RNN and BNN chickens were heavier (males, $p=0.0015$; females, $p<0.0001$) than NN chickens. The interaction between housing systems and genotypes showed that RNN and BNN male chickens reared in the intensive system and BNN female chickens in the semi-intensive system were heavier (males, $p=0.0009$; females, $p<0.00001$).

The body length of males did not differ among housing systems ($p=0.5539$) or genotypes ($p=0.3783$), and their interaction was not significant ($p=0.6835$). In females, longer bodies ($p<0.0001$) were determined in free-range and semi-intensive systems relative to the intensive system, whereas no body length differences were detected among genotypes. However, a significant interaction was detected between housing system and genotype, with RNN and BNN females in free-range and semi-intensive systems presenting the longest bodies ($p<0.0001$).

The keel length of males did not differ among housing systems ($p=0.0910$) or genotype ($p=0.4783$), and their interaction was not significant ($p=0.4278$). Maximum ($p<0.0001$) keel length was found in females reared in the semi-intensive compared with intensive and free-range systems. Regarding genotype, BNN and NN females had longer keels ($p=0.0002$) than

### RESULTS

**Morphometric traits**

Morphometric traits differed among genotypes, housing systems and their interaction (Tables 4, 5, 6).

### Table 4 – Effect of genotype and housing system on body weight, body length and keel length of chickens at 21 weeks of age. $^1$

| Genotype x Housing System | Body Weight (g) | Body Length (cm) | Keel Length (cm) |
|---------------------------|-----------------|------------------|-----------------|
|                           | Male            | Female           | Male            | Female           | Male            | Female           |
| RIR × NN                  | 1817.25 ± 45.32 | 1425.17 ± 18.35  | 69.64 ± 1.81    | 63.27 ± 1.67    | 11.68 ± 0.37    | 10.04 ± 0.35    |
| BAL × NN                  | 1811.17 ± 63.10 | 1456.22 ± 25.26  | 68.49 ± 2.39    | 62.27 ± 1.78    | 12.27 ± 0.36    | 10.89 ± 0.32    |
| NN                        | 1616.05 ± 30.99 | 1256.79 ± 34.92  | 69.41 ± 0.30    | 65.29 ± 0.76    | 12.27 ± 0.29    | 11.58 ± 0.17    |
| Free-range                | 1619.60 ± 20.88 | 1273.80 ± 31.76  | 69.53 ± 1.39    | 65.49 ± 1.39    | 11.54 ± 0.28    | 9.86 ± 0.31     |
| Semi-intensive            | 1774.89 ± 57.21 | 1412.12 ± 30.15  | 70.45 ± 1.57    | 66.76 ± 0.71    | 12.64 ± 0.33    | 11.75 ± 0.21    |
| Intensive                 | 1849.97 ± 56.20 | 1452.26 ± 19.65  | 67.56 ± 2.44    | 58.58 ± 1.85    | 12.03 ± 0.36    | 10.89 ± 0.32    |
| BAL × NN                  | 1639.64 ± 48.80 | 1558.90 ± 9.62   | 69.50 ± 3.40    | 65.48 ± 2.99    | 11.28 ± 0.53    | 8.30 ± 0.44     |
| RIR × NN                  | 1875.10 ± 53.89 | 1388.32 ± 26.57  | 69.64 ± 2.16    | 66.06 ± 1.50    | 12.07 ± 0.85    | 11.80 ± 0.52    |
| RIR × NN                  | 1936.80 ± 30.57 | 1328.28 ± 23.82  | 69.77 ± 4.44    | 58.28 ± 3.57    | 11.68 ± 0.66    | 10.01 ± 0.57    |
| BAL × NN                  | 1645.64 ± 28.13 | 1242.62 ± 10.87  | 72.68 ± 1.64    | 68.63 ± 2.32    | 11.37 ± 0.67    | 9.56 ± 0.50     |
| BAL × NN                  | 1822.51 ± 132.75 | 1663.10 ± 14.25  | 68.50 ± 3.16    | 67.49 ± 1.21    | 13.16 ± 0.47    | 12.24 ± 0.27    |
| BAL × NN                  | 1965.38 ± 90.89 | 1462.93 ± 11.51  | 64.31 ± 2.62    | 50.70 ± 2.19    | 12.27 ± 0.48    | 10.86 ± 0.64    |
| NN                        | 1573.33 ± 23.43 | 1019.88 ± 8.68   | 66.41 ± 0.72    | 62.37 ± 1.62    | 11.98 ± 0.22    | 11.73 ± 0.28    |
| NN                        | 1627.07 ± 63.94 | 1184.93 ± 20.37  | 73.22 ± 2.93    | 66.73 ± 0.94    | 12.70 ± 0.30    | 11.20 ± 0.24    |
| NN                        | 1647.75 ± 70.30 | 1565.57 ± 33.90  | 68.59 ± 5.15    | 66.76 ± 1.08    | 12.14 ± 0.82    | 11.81 ± 0.35    |

Source of Variation $p$-value

| Genotype          | 0.0015 | <0.0001 | 0.9044 | 0.2510 | 0.3783 | 0.0002 |
|-------------------|--------|---------|--------|--------|--------|--------|
| Housing System    | 0.0012 | <0.0001 | 0.5539 | <0.0001 | 0.0910 | <0.0001 |
| Genotype × Housing System | 0.0009 | <0.0001 | 0.6835 | <0.0001 | 0.4278 | <0.0001 |

$^1$ Means in the same column with no common superscript differ significantly at $p<0.05$.

$^\text{a}$ Values are mean ± standard error.

RIR = Rhode Island Red; NN = Naked Neck; BAL = Black Australorp.

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[1] eRBCA-2019-0921
The interaction between factors showed that RNN and BNN females reared in the semi-intensive system and NN female chicken reared in the free-range and intensive systems had maximum keel length. Drumstick and shank lengths were not influenced by housing system (males, \( p=0.1755 \), females, \( p=0.3391 \)), genotype (males, \( p=0.4638 \), females, \( p=0.4550 \), \( p=0.0939 \)), or their interaction (males, \( p=0.5830 \), females, \( p=0.8618 \), \( p=0.7999 \), \( p=0.6373 \)).

Table 5 – Effect of genotype and housing system on drumstick, shank length and drumstick circumference of chickens at 21 weeks of age.¹

| Genotype  | Housing System | Drumstick Length (cm) | Drumstick Circumference (cm) | Shank Length (cm) |
|-----------|----------------|-----------------------|-------------------------------|-----------------|
|           | Male           | Female                | Male                          | Female          |
| RIR × NN² | 13.69 ± 0.41   | 12.45 ± 0.49          | 8.00± 0.20                    | 6.70± 0.22      | 11.05 ± 0.55 | 8.42 ± 0.34 |
| BAL × NN² | 14.37 ± 0.31   | 12.64 ± 0.42          | 8.01± 0.39                    | 7.03± 0.24      | 11.10 ± 0.90 | 8.79 ± 0.44 |
| NN        | 13.76 ± 0.51   | 13.15 ± 0.26          | 10.13± 0.23                   | 9.82± 0.11      | 9.85 ± 0.27 | 9.46 ± 0.16 |
|           | Free-range     | 13.38 ± 0.42          | 12.56 ± 0.35                  | 8.69± 0.44      | 7.56 ± 0.26 | 10.06 ± 0.33 | 8.66 ± 0.22 |
|           | Semi-intensive | 14.53 ± 0.45          | 12.57 ± 0.39                  | 8.70± 0.41      | 7.65 ± 0.31 | 11.35 ± 0.91 | 9.04 ± 0.39 |
|           | Intensive      | 13.92 ± 0.33          | 13.10 ± 0.46                  | 8.75 ± 0.39     | 8.04 ± 0.27 | 10.60 ± 0.51 | 8.97 ± 0.38 |
| RIR × NN  | Free-range     | 13.09 ± 0.50          | 12.31 ± 0.80                  | 8.19± 0.51      | 6.94± 0.30  | 10.04 ± 0.78 | 8.00 ± 0.43 |
| RIR × NN  | Semi-intensive | 14.29 ± 1.05          | 12.01 ± 0.87                  | 7.81± 0.25      | 6.27± 0.49  | 12.07 ± 1.28 | 8.79 ± 0.66 |
| RIR × NN  | Intensive      | 13.69 ± 0.49          | 13.02 ± 0.92                  | 8.00± 0.32      | 6.88± 0.34  | 11.05 ± 0.62 | 8.47 ± 0.66 |
| BAL × NN  | Free-range     | 14.23 ± 0.70          | 12.07 ± 0.62                  | 7.97± 0.94      | 7.00± 0.46  | 10.73 ± 0.34 | 8.55 ± 0.41 |
| BAL × NN  | Semi-intensive | 14.51 ± 0.65          | 12.70 ± 0.71                  | 8.05± 0.70      | 6.87± 0.39  | 11.47 ± 2.63 | 8.98 ± 0.97 |
| BAL × NN  | Intensive      | 14.38 ± 0.35          | 13.14 ± 0.85                  | 8.01± 0.55      | 7.22± 0.39  | 11.10 ± 1.35 | 8.85 ± 0.83 |
| NN        | Free-range     | 12.80 ± 0.92          | 13.31 ± 0.24                  | 9.90± 0.48      | 9.62± 0.15  | 9.40 ± 0.44 | 9.44 ± 0.20 |
| NN        | Semi-intensive | 14.77 ± 0.79          | 13.00 ± 0.40                  | 10.24± 0.36     | 9.82± 0.15  | 10.50 ± 0.44 | 9.34 ± 0.22 |
| NN        | Intensive      | 13.70 ± 0.85          | 13.14 ± 0.66                  | 10.24± 0.45     | 9.65± 0.46  | 9.59 ± 0.40 | 9.59 ± 0.40 |

Source of Variation

| Source of Variation | p-value |
|--------------------|---------|
| Genotype           | 0.4638  |
| Housing System     | 0.1755  |
| Genotype × Housing System | 0.5830 |

¹Means in the same column with no common superscript differ significantly at \( p<0.05 \).

Values are least square mean ± standard error.

Table 6 – Effect of genotype and housing system on shank circumference and wingspan of chickens at 21 weeks of age.¹

| Genotype  | Housing System | Shank circumference (cm) | Wingspan (cm) |
|-----------|----------------|--------------------------|---------------|
|           | Male           | Female                   | Male          | Female        |
| RIR × NN² | 4.25 ± 0.21    | 3.56 ± 0.12              | 10.25 ± 0.48  | 8.30 ± 0.34   |
| BAL × NN² | 4.06 ± 0.10    | 3.56 ± 0.11              | 11.04 ± 0.46  | 8.94± 0.26    |
| NN        | 3.58 ± 0.11    | 3.35 ± 0.06              | 10.01 ± 0.23  | 9.62± 0.14    |
|           | Free-range     | 3.98 ± 0.21              | 3.44 ± 0.10   | 10.10 ± 0.40  | 9.01 ± 0.24 |
|           | Semi-intensive | 3.69 ± 0.12              | 3.67 ± 0.09   | 10.87 ± 0.47  | 8.96 ± 0.28 |
|           | Intensive      | 4.01 ± 0.10              | 3.36 ± 0.09   | 10.34 ± 0.38  | 8.90 ± 0.29 |
| RIR × NN  | Free-range     | 4.47 ± 0.51              | 3.57 ± 0.23   | 10.04 ± 0.89  | 8.53 ± 0.57 |
| RIR × NN  | Semi-intensive | 4.03 ± 0.39              | 3.74 ± 0.19   | 10.47 ± 0.98  | 8.21 ± 0.62 |
| RIR × NN  | Intensive      | 4.25 ± 0.17              | 3.37 ± 0.19   | 10.25 ± 0.87  | 8.16 ± 0.62 |
| BAL × NN  | Free-range     | 3.98 ± 0.16              | 3.35 ± 0.19   | 10.46 ± 0.90  | 9.07 ± 0.40 |
| BAL × NN  | Semi-intensive | 4.14 ± 0.21              | 3.96 ± 0.14   | 11.63 ± 0.99  | 8.95 ± 0.51 |
| BAL × NN  | Intensive      | 4.06 ± 0.16              | 3.37 ± 0.18   | 11.05 ± 0.60  | 8.80 ± 0.48 |
| NN        | Free-range     | 3.50 ± 0.17              | 3.40 ± 0.11   | 9.79 ± 0.23   | 9.42 ± 0.17 |
| NN        | Semi-intensive | 3.51 ± 0.47              | 3.31 ± 0.09   | 10.51 ± 0.47  | 9.71 ± 0.22 |
| NN        | Intensive      | 3.73 ± 0.12              | 3.33 ± 0.11   | 9.72 ± 0.43   | 9.75 ± 0.34 |

Source of Variation

| Source of Variation | p-value |
|--------------------|---------|
| Genotype           | 0.0150  |
| Housing System     | 0.8485  |
| Genotype × Housing System | 0.2115 |

¹Means in the same column with no common superscript differ significantly at \( p<0.05 \).

Values are least square mean ± standard error.

RIR = Rhode Island Red; NN = Naked Neck; BAL = Black Australorp.
interaction (males, $p=0.5830$, females, $p=0.8618, p=0.6373$). Larger drumstick circumference (males, $p<0.0001$; females, $p<0.0001$) in NN chickens than those of RNN and BNN chickens. The interaction between housing system and genotype determined the largest (males, $p=0.0039$; females, $p<0.0001$) drumstick circumference in NN chickens reared in the free-range and intensive systems. Larger shank length was not affected by the treatments. No differences in wingspan was determined in males ($p>0.05$), however, NN females had longer wingspan ($p=0.0029$) compare with RNN females.

Serum Chemistry and Antibody Response

There were no differences in total protein, albumin, globulin, uric acid and creatinine blood levels among genotypes and housing systems and no significant interactions were detected (Table 7, 8).

There was no influence of housing system or genotype on total protein, albumin, globulin, uric acid or creatinine levels ($p>0.05$). However, serum glucose and cholesterol levels, as well as antibody responses against ND and IB differed among treatments (Table 8). Serum glucose level was higher ($p=0.0008$) in females reared in the intensive system relative to the semi-intensive and free-range systems, and in NN birds than in RNN birds ($p<0.0123$). The interaction between housing system and genotype showed the highest ($p=0.0164$) plasma glucose level in NN females reared in the intensive system. Higher cholesterol levels ($p=0.0123$) were detected in NN birds compared with BNN. The interaction between housing system and genotype was significant ($p=0.0103$), with the highest cholesterol level measured in BNN birds reared in the intensive system. Relative to antibody titers against ND, higher ($p=0.0204$) titer were determined in RNN birds than in BNN. Furthermore, higher ($p=0.0001$) antibody titer against IB was found in free-range chickens followed by those reared in the semi-intensive and intensive systems. The interaction showed that NN birds reared in the free-range system had the highest ($p=0.0067$) titer against IB.

DISCUSSION

The present study aimed at comparing morphometric traits, serum chemistry and antibody response among different genotypes and housing systems.

When housing systems were compared, although no differences were detected in drumstick length and circumference, shank length and circumference, and wingspan, males were 9-14% and females were 11-14% heavier when reared in the intensive and semi-intensive systems at market age (21 weeks) compared...
with those reared in the free-range system. This may be attributed to the active behavior of free-range chickens. In general, these birds do more exercise during their life span, ultimately spending more calories. These results are in agreement with the findings of Rehman et al. (2016) who found higher body weight of Aseel chicken varieties when reared under intensive and semi-intensive housing systems. Likewise, Olaniyi et al. (2012) reported higher body weight of Harco black and Novogen cockerels when reared under deep litter system as compared to free-range reared birds. Similarly, reduced body weight in slow-growing broilers exposed to free-range access was reported by Stadig et al. (2016). In the present study, the longest body length was measured in free-range females, and keel length females reared in the semi-intensive system.

Differences among genotypes were also detected. Both male and female RNN and BNN chickens were heavier at 21 weeks of age, and larger Shank circumference than NN chickens. Longer keels were measured in BNN and NN females, whereas higher drumstick circumference and wingspan values were determined in NN chickens.

The observed differences in morphological traits agree with the findings of Qureshi et al. (2018), who found variation among different phenotypes of Aseel chickens in Pakistan. Similarly, Adekoya et al. (2013) and Fadare (2014) reported variation in morphological traits among five indigenous chicken genotypes in Nigeria.

The higher glucose level obtained in female reared in the intensive system relative to semi-intensive and free-range systems are consistent with the reports of Gunes et al. (2002) and Rehman et al. (2016), who evaluated alternative housing systems and determined higher blood glucose levels in intensively-reared layers and in Aseel chickens, respectively. It is possible that the lower plasma glucose level determined in free-range chickens may be due to their intense exercise, which ultimately increases insulin level and stimulates glucose metabolism.

There was no influence of housing systems on cholesterol level, in agreement with other studies (Elerogly et al. 2011; Diktas et al. 2015; Eleroglu et al. 2015) that found negligible effects of housing system on cholesterol level among different chicken genotypes. However, higher cholesterol level was determined in NN than in BNN birds. This may be attributed to their specific genetic makeup.

Antibody titers against ND were not influenced by housing system, but higher titers were determined in RNN than in BNN birds. This result may be attributed to distinct genetic resistance against the disease, which was more pronounced in RNN chickens compared with BNN chickens. On the other hand, genotype did not affect antibody titers against IB, whereas higher titers were measured in free-range chickens followed

| Genotype × Housing System | Cholesterol (mg/dL) | ND (HI titer) | IB (ELISA titer) |
|--------------------------|---------------------|---------------|-----------------|
| RIR × NN Free-range      | 134.48 ± 3.50       | 5.10 ± 0.06   | 3629.91 ± 53.88 |
| BAL × NN Free-range      | 127.11 ± 5.85       | 4.70 ± 0.10   | 3629.89 ± 70.91 |
| NN                       | 143.87 ± 3.13       | 4.95 ± 0.11   | 3599.70 ± 87.39 |
| Free-range               | 128.96 ± 5.41       | 4.98 ± 0.10   | 3823.56 ± 30.79 |
| Semi-intensive           | 138.01 ± 4.44       | 4.79 ± 0.14   | 3598.62 ± 31.44 |
| Intensive                | 138.48 ± 4.21       | 4.97 ± 0.05   | 3437.32 ± 65.58 |
| RIR × NN Free-range      | 131.84 ± 3.06       | 5.13 ± 0.07   | 3801.17 ± 51.87 |
| RIR × NN Semi-intensive  | 140.83 ± 4.92       | 5.08 ± 0.15   | 3588.95 ± 68.15 |
| RIR × NN Intensive       | 130.75 ± 9.09       | 5.07 ± 0.09   | 3499.61 ± 59.20 |
| BAL × NN Free-range      | 112.83 ± 9.41       | 4.80 ± 0.10   | 3801.14 ± 69.18 |
| BAL × NN Semi-intensive  | 123.11 ± 4.79       | 4.40 ± 0.13   | 3640.87 ± 22.34 |
| BAL × NN Intensive       | 145.38 ± 4.69       | 4.92 ± 0.10   | 3447.66 ± 154.03|
| NN                       | 142.21 ± 5.60       | 5.02 ± 0.28   | 3868.35 ± 48.92 |
| NN Semi-intensive        | 150.09 ± 1.04       | 4.89 ± 0.25   | 3566.04 ± 72.42 |
| NN Intensive             | 139.30 ± 7.34       | 4.94 ± 0.08   | 3364.70 ± 140.52|

Source of Variation

| p-value |
|---------|
| Genotype | 0.0123 | 0.0204 | 0.8858 |
| Housing System | 0.1274 | 0.2546 | 0.0001 |
| Genotype × Housing System | 0.0103 | 0.1001 | 0.0067 |
by those reared in the semi-intensive and intensive systems. Similar differences in antibody response against ND and IB among different chicken and duck genotypes were obtained by Shini (2003), Arbona et al. (2011), Shi et al. (2011), and Rehman et al. (2016).

CONCLUSIONS

In general, morphometric traits and serum chemistry were not affected by housing system, except for a few differences observed regarding body weight, body and keel length, plasma glucose, cholesterol and antibody response against ND and IB. Therefore, alternative housing systems (semi-intensive and free-range) can successfully be adopted for dual-purpose chicken genotypes.

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CONFLICT OF INTEREST

No potential conflict of interest was found by the authors.

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