Comparative study of semen traits and histomorphometric features of testes of broiler breeder males with different phenotypic traits

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Article Info

Abstract

Fertility reduction due to sub-fertile males is a major concern in breeder flocks. Phenotypic traits of breeder male flocks and their relationships with fertility can be used as reliable indicators for identification and removal of sub-fertile males from breeder flocks. This study was conducted to investigate semen traits (semen volume, sperm motility, sperm viability and sperm count) and testes histomorphometric features including tubule differentiation index (TDI), spermiation index (SPI), Sertoli cell index (SCI) and mitotic index (MI) of broiler breeder males with the same age but different phenotypic traits. According to phenotypic traits, 12 broiler breeder males (Ross-308 strain) were classified into three equal groups. Group 1: roosters with fertile phenotypic traits (fertile), group 2: roosters with the lowest fertile phenotypic traits (sub-fertile) and group 3: roosters with moderate fertile phenotypic traits (moderate). The results confirmed potential relationship between phenotypic traits and fertility in broiler breeder males. Semen traits and histomorphometric features of broiler breeder males' testis of the group 3 were more similar to those of the fertile roosters. Therefore, it can be concluded that exclusion of these roosters from the breeder flock may have undesirable effects on flock fertility.

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Key words: Breeder male, Histomorphometry, Phenotype, Semen, Testis

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Introduction

Reproduction is the most important requisite of livestock breeding and fertile eggs play a dominant role in assessment of their performance. It is well known that breeder males’ phenotypes have a great impact on the percent of fertile eggs and the genetic of breeder males has also a major influence on their progeny performance.1 In commercial flocks, fertility of broiler breeders typically peaks between 30 to 40 weeks of age2-4 and routinely declines after 45-50 weeks of age.4 It is now well documented that low fertility could be largely due to the males’ features in broiler breeder flocks. As the breeder males age increases, there is a reduction in the number of spermatozoa, volume of semen2,3,5,6 and libido as well as changes in the body conformation that inhibit mating and affecting sperm quality.6 Observable secondary sex features are used not only as a criterion for reproductive maturity and health status in breeder flocks;7 but also can be used as the indicators for high semen physical traits.8 Moreover, there are positive genetic correlations between phenotypic characteristics (mostly comb and wattle measurements) with semen traits which can be used as reliable indices to facilitate the identification and removal of sub-fertile males from the breeder flocks.9

The degree of development of the secondary sexual features could also affect the reproductive vigor of an individual.10-12 The size and color of comb are significant predictors of sperm viability13 as males with larger combs were likely to have higher fertility.9 In naturally mated broiler breeder flocks, the reduction of fertility is mostly due to sub-fertile males in which their behavior, physiological (i.e. development of testes and hormonal secretion) and physical (i.e. size of comb and wattles) characteristics affect their fertility.14 In general, hens may mate with several roosters and choose males based on multiple secondary sexual characteristics.9,13,15,16 As the assessment of semen quality characteristics of roosters give an excellent indicator of their reproductive potentials,17 thus, this study was conducted to investigate the semen traits (semen volume, sperm motility, sperm viability and sperm count) and testis histomorphometric features including tubule differentiation index (TDI), spermiation index (SPI), Sertoli cell index (SCI) and mitotic index (MI) of broiler breeder males with almost the same weights and the same age, but with different phenotypic traits in order to predict the fertility in broiler breeder flocks.

Materials and Methods

Chickens. Twelve broiler breeder males (Ross-308 strain) at 42 weeks of age but with different phenotypic characteristics were selected from a broiler breeder flock and divided into three equal groups according to their phenotypic features including alertness and activity, body condition (shape and softness or hardness of breast muscle tone), head (uniform, intense red color around the comb, wattle and eye area), feathering (partial feather loss, especially around the shoulders and thighs) and vent (some feather wear, be large and moist, with red coloration) as shown in Figure 1.18 Accordingly, roosters with the best phenotypic features were placed in group 1 (fertile group), while roosters with the lowest phenotypic features were placed in group 2 (sub-fertile group), and roosters with moderate phenotypic features were placed in group 3 (moderate group). All roosters were housed in individual pens and were fed with a recommended breeder diet (2700 kcal kg⁻¹ diet, 11.50% crude protein, 0.70% calcium and 0.35% phosphorous) according to Ross 308 parent stock nutritional specification manual 2016.19 The ethical approval was obtained from the Animal Ethics Committee of Urmia University (AECVU-155-2017).

Semen collection. In order to collect semen, the roosters were trained for 10 days before semen collection began. Semen collection was performed by abdominal massage as previously described.20 Briefly, semen was obtained by gently massaging (stroking) the back of roosters with the palm of the hand while the abdomen was massaged towards the tail with the other hand. Two people were involved in performing the semen collection, one holding the rooster by the thigh and the other massaging for collecting the semen. After excitation of the roosters with abdominal massage, the male organ becomes swell and protrude while white semen can be seen in the central furrow of the organ. The semen was milked down by firm finger pressure either side of the vent into the collection tube. Immediately after collection, each ejaculate was evaluated for volume, motility, concentration and live-dead ratio of sperm.

Semen volume. Semen volume (mL) from each rooster using graduated centrifuge tubes was measured as previously described.21

Fig. 1. Head and vent features in different groups of the examined roosters. A and D: head and vent of group 1 (fertile roosters), B and E: head and vent of group 2 (Sub-fertile roosters). C and F: head and vent of group 3 (roosters between groups 1 and 2).
Sperm motility. Immediately after collection, semen was diluted (1:200) in modified ringer solution (NaCl: 68 g, KCl: 17.33 g, CaCl₂: 6.42 g, MgSO₄: 2.50 g, NaHCO₃: 24.50 g and distilled water 10,000 mL) as previously described. For evaluation of motility, one drop of the diluted semen was placed on a slide and covered with a cover slide, then sperm motility was estimated by microscopic observation (400× magnification). Motility was expressed as the percentage of motile spermatozoa with moderate to rapid progressive movement. A minimum of five microscopic fields were assessed to evaluate sperm motility on at least 200 sperm for each sample and percentage of motile sperms were calculated as previously described.

Sperm viability. A 20 μL of sperm suspension (1:10) was mixed with an equal volume of 0.05% eosin-Y on a slide. After 20-30 sec, 20 μL nigrosin was added. Slides were examined using light microscope with 400× magnification following 2 min incubation at room temperature. Dead sperms appeared to be pink and live sperms were not stained. In each sample, 200 sperms were counted and their viability (%) were recorded.

Sperm count. Sperm concentration was determined using the standard hemocytometric method as previously described. Briefly, approximately 10 μL of sperm suspension (1:200) was transferred to each of the counting chambers of the Neubauer hemocytometer (HBG, Berlin, Germany), and was allowed to stand for five min in a humid condition in order to prevent drying. The cells were counted using a light microscope (400× magnification). The counted sperms were expressed as number of sperm per mL.

Histological analysis. At the end of the study, the males were euthanized (cervical dislocation) according to American Veterinary Medical Association guideline and specimens from testis were collected for histological evaluation as previously described. Briefly, after tissue fixation in 10% buffered formalin, specimens were processed through paraffin embedding and cut into 6 μm sections, stained with periodic acid-Schiff (PAS) technique. All specimens were studied under 400× and 1000× magnifications. Tubule differentiation index (TDI: the percentage of seminiferous tubules containing at least three differentiated germ cells) and spermatiation index (SPI: the percentage of seminiferous tubules with normal spermatiation) were calculated. Two hundred cross-sections of seminiferous tubules were randomly analyzed (one hundred per testis) for calculation of TDI and SPI. For the estimation of Sertoli cell index (SCI: the ratio of the number of germ cells to the number of Sertoli cells identified by a characteristic nucleus in all seminiferous tubules) and mitotic index (MI: the number of round spermatids for each pachytene primary spermatocytes, was calculated for determination of cell loss percentage during cell division), sixty seminiferous tubules per group were randomly examined.

Statistical analysis. The results are expressed as the mean ± standard error (Mean ± SE). Differences among the groups were assessed by one-way analysis of variance using SPSS software package for Windows (version 23.0; IBM Corp., Armonk, USA). Statistical significance among the groups was determined by Tukey multiple comparison post-hoc test and the p-values < 0.05 were considered to be statistically significant.

Results

Sperm traits (sperm viability, sperm motility and sperm count) of the examined roosters are shown in Table 1. There was no significant difference in sperm volume among the roosters. As shown in Table 1, the percent of sperm motility in group 1 was significantly (p < 0.05) higher than that of group 2. There was also a significant difference (p < 0.05) in sperm motility between group 2 and group 3. Evaluation of sperm viability among the groups showed that sperm viability rate differed among the groups but the difference was not significant (Fig. 2). Sperm count was significantly higher in group 1 roosters in comparison with those in group 2 (p < 0.05).

Average weights of right and left testes and morphometric features of the testes in different groups are presented in Tables 2 and 3. Weight of both testicles (right and left) of the roosters of group 1 was significantly higher than those of group 2 and 3 (p < 0.05), while there was no significant difference in testes’ weight of roosters belonged to groups 2 and 3 (Table 2).

Table 1. Sperm properties of the examined broiler breeder males. Data are presented as mean ± SE.

| Parameters                        | Group 1            | Group 2            | Group 3            |
|-----------------------------------|--------------------|--------------------|--------------------|
| Motility (%)                      | 78.00 ± 8.87a      | 13.70 ± 8.00b      | 57.75 ± 10.00a     |
| Viability (%)                     | 96.97 ± 1.04a      | 44.37 ± 5.62a      | 90.97 ± 2.43a      |
| Count (10⁶ mL⁻¹)                  | 2.62 ± 0.23a       | 0.59 ± 0.34b       | 1.68 ± 0.37ab      |

Different superscripts indicate significant difference in each row (p < 0.05).

Fig. 2. Live and dead sperm cells. Dead sperm appeared to be pink (DS) and live sperms were not stained (LS) (eosin-nigrosin 1000×).
The SPI, TDI, SCI and MI of both testes in group 1 was significantly higher than those of group 2 (p < 0.05; Tables 2 and 3). Regarding to TDI, SCI and MI, difference were not significant between groups 1 and 3. The histological section of right and left testes in different groups are illustrated in Figure 3. The SPI of the left testes were significantly different between groups 1 and 3 (p < 0.05), however, SPI of the right testes were not significantly different between groups 1 and 3 (p > 0.05). The SPI and TDI calculated for groups 2 and 3 were significantly different (p < 0.05), however, SCI did not differ between these two groups. The results showed that MI of right testes were significantly different between groups 2 and 3 (p < 0.05).

**Discussion**

Fertility problems have a direct effects on performance and profitability of breeder flocks and infertility are mostly contributed to cockles’ failure in birds. On the other hand, semen analysis is the most common way of accessing infertility in males. Previous studies indicated that reduced sperm concentration and semen volume in aging broiler breeder males contributed to reduction of fertility.33,34

As in broiler breeder flocks, fertile eggs are the main products which are used to produce one-day old chicks, therefore reduced fertility leads to infertile eggs and finally less chicks. Tabatabaei et al. reported that sperm mobility and viability are reduced in aging roosters.37 Hence, identification and removal of infertile roosters are one of the main goals of aged breeder flocks’ management. Unfortunately, identification of sub-fertile males which also have a negative impact on flock fertility is a vital issue, too. Relationships between phenotypic characteristics with fertilization (sperm trait and histomorphometric features of testes) in aged roosters provided practically acceptable features to boost the recognition and elimination of sub-fertile males from the breeder flock. As shown in Table 1, roosters of the group 1 had higher sperm motility (almost six folds), sperm viability (almost two folds) and sperm count (almost 5 folds) in comparison with roosters of the group 2, while differences of these features between group 1 and group 3 were not significant (p > 0.05). Regarding to testis histomorphometric features as shown in Tables 2 and 3, there is a significant (p < 0.05) difference between group 1 and group 2 but differences between group 1 and group 3 was not significant (p > 0.05). Also, Figure 3 confirms the

**Table 2.** Histomorphometric features of right testes in the examined broilers. Data are presented as mean ± SE.

| Parameters                              | Group 1                | Group 2                | Group 3                |
|-----------------------------------------|------------------------|------------------------|------------------------|
| Testis weight (g)                       | 13.12 ± 1.11           | 4.78 ± 1.26           | 6.88 ± 1.14           |
| Spermiation index (%)                   | 75.86 ± 6.11           | 11.55 ± 1.9           | 55.45 ± 9.56           |
| Tubule differentiation index (%)        | 86.80 ± 1.90           | 23.38 ± 1.44          | 77.12 ± 5.18           |
| Sertoli cell index                      | 27.90 ± 5.27           | 4.90 ± 0.49           | 18.74 ± 0.63           |
| Mitotic index                           | 4.33 ± 0.61            | 0.72 ± 0.14           | 2.64 ± 0.21           |

ab Different superscripts indicate significant difference in each row (p < 0.05).

**Table 3.** Histomorphometric features of left testes in the examined broilers. Data are presented as mean ± SE.

| Parameters                              | Group 1                | Group 2                | Group 3                |
|-----------------------------------------|------------------------|------------------------|------------------------|
| Testis weight (g)                       | 14.65 ± 1.55           | 4.35 ± 1.15           | 7.09 ± 1.33           |
| Spermiation index (%)                   | 83.47 ± 4.70           | 2.50 ± 0.25           | 54.87 ± 6.26           |
| Tubule differentiation index (%)        | 88.65 ± 2.50           | 14.42 ± 0.85          | 84.87 ± 2.44           |
| Sertoli cell index                      | 31.34 ± 3.14           | 5.02 ± 0.52           | 17.98 ± 1.98           |
| Mitotic index                           | 4.53 ± 0.34            | 0.55 ± 0.13           | 2.07 ± 0.54           |

abc Different superscripts indicate significant difference in each row (p < 0.05).

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Fig. 3. Photomicrographs of testicular sections of the roosters. A) Testes of group 1 roosters exhibit relatively normal features of seminiferous tubules with active spermatogenesis and presence of sperms (SP) in lumen. B) Testes of group 2 roosters had germ cells depletion and impaired spermatogenesis. In some tubules, small numbers of sperms are present in lumen. C) Testes of group 3 roosters showed partial disorganizations and vacuoles (black arrows) in seminiferous tubules (PAS, Bar = 100 µm).
results of histomorphometric evaluations and consistent with the data of Tables 2 and 3. Furthermore, the results of this study (Tables 1, 2, 3 and Figs. 2, 3) had a good relation with phenotypic traits as shown in Figure 1 and confirms that the roosters of group 1 had higher fertility rate when compared with roosters that do not have these features (group 2 and 3). Our results are in agreement with the results of previous studies reporting that the size and color of comb in roosters could be used as significant predictors of their sperm viability, quality and sperms ability to reach and hydrolyze the perivitelline membrane of the ovum. Reportedly, the reproductive potentials of an individual male could be affected by the degree of development of the secondary sexual characters, and testosterone is essential not only for their development but also for normal mating behavior. Moreover, it has been reported that sperm quality of roosters will be improve by the presence of hens and breeding females crouch and mate more frequently with males possessing large comb and wattle. Overall, our results support the hypotheses of correlation between phenotypic aspects with fertility and could be used for maintain of a good fertility management in broiler breeder flocks.

In conclusion the results of this study indicated that the roosters of the group 3 which were placed between fertile and sub-fertile groups, based on semen traits and histomorphometric features of testes, were more similar to fertile roosters and their exclusion may be resulted in lower fertility in the flock.

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Conflict of interest

The authors declare no potential competing conflict of interest.

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