Descriptive and cytogenetic study on testicular hypoplasia in a buffalo bull

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Abstract
Phenotypic, andrologic, and cytogenetic investigations were performed on a 2.5 years old buffalo-bull that produced azoospermic semen from puberty. Also, ultrasonographic and histopathologic examinations were conducted on its testis. Phenotypic examinations showed atypical male features. Andrological investigations revealed small non-pendulous scrotum and reduced testicular size as indicated by lowered scrotal circumference measurement and ultrasound-detected testicular breadth. Testicular hypoplasia was confirmed by the histopathological findings represented by a marked reduction in the number and diameter of seminiferous tubules and absence of mitotic or meiotic activities. Cytogenetic examination of peripheral leucocytes revealed the presence of trisomy-24. It could be concluded that identifying trisomy-24 as possible cause of testicular hypoplasia in buffalo bulls is being useful for earlier diagnosis of the affected cases before reaching the age of puberty when the condition could be diagnosed and thus reducing the costs of keeping them till puberty.

Introduction
Keeping buffalo in small numbers among large number of breeders and wide spread use of uncontrolled natural service interfere with its genetic improvement and tracing up of genetic causes of infertility. Farm animals with spermatogenic impairment are usually not identified until the age they are expected to breed [1]. Among the non-infectious causes, genetic defects constitute an important factor for reproductive failure in cattle [2]. Cytogenetic studies in cattle, horses, sheep and pigs have been clearly identified chromosomal aberrations as one of the important factors contributing to infertility problems [3]. In cattle, testicular hypoplasia has been associated with chromosomal aberration such as mosacism [4] or 61xxy; the counterpart of human Klinefelter’s syndrome [5]. Abnormal distribution of sex chromosomes or autosomes bearing sex influencing genes may affect the development of the testis and male masculinity [6]. Trisomy of chromosomes-22 results in sterility and complete spermatogenic arrest in man [7]. In contrast to human, chromosomal aberrations in domestic animals may be more common than they appear probably because of lack of cytogenetic investigators, facilities or difficulty in obtaining materials containing chromosomal aberrations [8]. The main aim of this work was to describe testicular hypoplasia in a buffalo bull and investigate its possible genetic cause.

Material and method
The animal and its phenotype
The buffalo bull used in the current study was born as singleton for two apparently normal parents of Egyptian buffalo; it was 2.5 years in age and weighed 500 kg. The bull has been recorded to ejaculate azoospermic semen from puberty. Five bulls having the similar age and body weight were used as a control group. All were reared in Mahallet Mousa Buffalo Experimental Station. The secondary sex characters of the affected bull were studied in comparison with control group.

Sexual desire and semen evaluations
Libido was assessed by estimation of reaction time. Semen was collected by means of artificial vagina twice weekly for 6 weeks. The collected semen was directly evaluated [9].

Serum testosterone concentration
Blood samples were collected once weekly for 6 weeks from the affected and control bulls. Sera were obtained by centrifugation (3000 rpm) for 10 minutes and stored at -20°C until assessment of testosterone. Serum testosterone concentrations were measured by Enzyme-linked immunosorbantassay kits (ELISA).

Scrotal circumference (SC)
Scrotal circumferences for the affected bull and control ones were measured at the largest diameter of the testis according to Almiquist et al., [10].

Ultrasound examination of testis
The breadth and ultrasound features of the testis were detected by ultrasound scanning of each testes, while the animal was controlled in lateral recumbent position and one of its hind limb was pulled to the...
anterior and the other to the posterior. The ultrasound examination was done using a real time B-mode scanner (Ultrascan, 900, Alliance, Inc., Quebec, Canada) equipped with a 5 MHz linear transducer (L118 x D23 mm).

**Histopathological examination**

At slaughter, the testis of the affected bull and one of the control bulls were dissected, photographed and weighed. Several sections were taken from the core, peripheral parts of the 2 poles and middle part of the testis and fixed for histopathological examination [11].

**Cytogenetic examination**

Blood samples were collected from jugular vein by means of 5 ml heparinized vacutainer tubes. Leucocytes were cultured according to the modified method reported by Basrur and Gilman [12]. Air and flame dried slides were prepared. Some were stained using Geimsa stain [13]. The stained slides were examined for metaphase plates. Suitable plates were examined and photographed for preparation of karyotypes.

**Result**

**Phenotypic characters**

The affected bull showed atypical male phenotype. Its face shifted to the female type when compared with that of a normal bull (Figure 1). Unlike those of the normal, the scrotum of the affected bull was not pendulous in hot weather (Figure 2).

**Sexual desire and semen characteristics**

The affected bull had a reduced sexual desire as indicated by mounting after hesitation with feeble holding and longer reaction time compared with normal bulls (Table 1). At all times of semen collections, the ejaculate of the affected bulls was clear, watery and azoospermic. The overall mean of the ejaculate volume of the affected bull was greatly reduced compared with that of the controls (Table 1).

**Serum testosterone concentration**

The overall mean of the serum testosterone concentration in the affected bull was lower than that in normal bulls (Table 1).

**Scrotal circumference and ultrasound-detected testicular breadth**

The overall means of scrotal circumferences were 20.5 cm and 32.8 cm for the affected and a normal comparable bull, respectively. The weights and ultrasound-detected testicular-breadths of both testes in the affected bull were numerically lower than those of the normal ones (Table 2 and Figure 3).

**Ultrasonographic features**

The testes of one of the normal control bulls appeared moderate and homogenous echogenic with a central hyperechoic line, representing mediastinum testis while that of the affected one appeared hypoechoic with hyperechoic central line (Figure 4).

**Histopathological findings**

Histopathological examination of the testes of the affected bull revealed a marked reduction in the number of the seminiferous tubules with massive interstitium in-between (Figure 5b) compared with that of the normal bull (Figure 5a). The interstitium is formed from large amount of collagen fibers and numerous blood vessels and islets of Leydig cells with some of them containing pyknotic nuclei (Figure 6a). Also, the diameters of seminiferous tubules were markedly reduced with no evidence of spermatogenesis as indicated by absence of both mitotic and meiotic activities (Figure 6b). They were lined by only one layer of spermatogonia. In contrast to the normal (Figure 5a), the seminiferous tubules were collapsed or folded (Figure 5b).

**Cytogenic findings**

Examination of Geimsa stained metaphase spreads of the affected bull revealed four lines of cells: 50, xy/48, xy/49, xy/ and 51, xy with testicular hypoplasia was mainly due to absence of the germ cells. Such suggestion may be supported by the findings of Moura and Erickson [1] who found that the difference in the testicular weight and seminiferous tubules diameter between normal and bull affected with testicular hypoplasia was mainly due to absence of the germ cells. This suggestion was confirmed by the histopathological findings where there was few numbers of the seminiferous tubules that were lined with one layer of the spermatogonia with absence of any mitotic and meiotic activities. These findings are similar to those reported by Bongso et al., [2]. In contrast to the moderate homogenous echogenic feature with central hyperechoic line characteristic of the normal testes [17] in the control bull, the testes of the affected bull showed a relative hypoechoic feature. However, the hypoechoic feature may be explained in the light of marked reduction in the number of the seminiferous tubules and presence of one layer of spermatogonia within the tubules.

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**Table 1.** The reaction time, semen characteristics and serum testosterone concentration in the affected and normal control bulls.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Group</th>
<th>Affected bull</th>
<th>Normal bulls (n=5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reaction time (sec)</td>
<td></td>
<td>564 ±15.1</td>
<td>56.9±8.5</td>
</tr>
<tr>
<td>Ejaculate volume (ml)</td>
<td></td>
<td>1.8±0.39</td>
<td>3.1±0.23</td>
</tr>
<tr>
<td>Individual motility (%)</td>
<td></td>
<td>ND</td>
<td>79.8±4.8</td>
</tr>
<tr>
<td>Sperm cell concentration (x10^6/ml)</td>
<td></td>
<td>ND</td>
<td>112±48.17</td>
</tr>
<tr>
<td>Serum testosterone concentration (ng/ml)</td>
<td></td>
<td>0.13±0.2</td>
<td>0.89±0.6</td>
</tr>
</tbody>
</table>

ND= not detected.

**Table 2.** The scrotal circumferences, testicular breadths and weights for the affected bull and a normal control one.

<table>
<thead>
<tr>
<th>Group</th>
<th>Ultrasound detected testicular-breadth</th>
<th>Testicular weight (gm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Right tests (mm)</td>
<td>Left tests (mm)</td>
</tr>
<tr>
<td>AFFECTED BULL</td>
<td>17.9</td>
<td>13.0</td>
</tr>
<tr>
<td>CONTROL BULL RATIO</td>
<td>53.4</td>
<td>43.4</td>
</tr>
</tbody>
</table>

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Also, the lower serum testosterone concentration in the bull with testicular hypoplasia compared with normal bulls explains the presence of both feminine characters and low libido. Such low serum testosterone concentration may be attributed to the lower rate of testosterone production by the hypoplastic testes compared with the normal ones [18]. Veeramachaueni et al., [19] attributed the lowered serum testosterone concentration in bulls with testicular hypoplasia to the impairment in the function of the Leydig's cells by local factors either causing tubular damage or those consequent to the tubular damage.

Identification of trisomy in the autosome 24 in affected bull suggests a correlation between testicular hypoplasia and such chromosomal aberration. According to the standard Karyotype of the river buffalo [20,21] this chromosome resembles chromosome 25 in cattle. Chromosome number 25 in cattle carrying many genes some of them are reported to play a role in fertility.

Testicular organogenesis involves a cascade of gene activation and differentiation of the component cell types [22]. When cell organization is abnormal, the testes is described as dysgenetic, which could be resulted from chromosomal abnormalities [23].

According to the available literature this may be the first record to correlate between testicular hypoplasia and trisomy-24 in buffalo bulls. A comparable study in human, reported that trisomy-21 resulted in sterility and complete spermatogenetic arrest in man [7]. However, the atypical male features of the affected bull may be attributed to the abnormal distribution of the autosome-24 that might be a one of those bearing masculinity determining genes. Pineda [6] cited that genes for masculinity on both autosomes and on Y- chromosome must overcome and prevail over the female determining genes on the single x-chromosomes of an XY individual. However further studies are needed to determine if such chromosome is carrying the concerned gene.

Figure 1. Faces of both normal and affected bulls showing the feminine characters on the face of the affected bull.

Figure 2. Scrotum of both affected and normal bulls showing non-pendulous scrotum in the affected bull.

Figure 3. Photograph of the right and left testis in both affected (AF) bull and (NT) normal control one.

Figure 4. Ultrasonographic feature of affected (a) and normal testes (b)

Figure 5. Cross section in the testies of the normal (a) and affected (b) bulls stained with H&E (x100).

Figure 6. Cross section in the testies of the normal (a) and affected (b) bulls stained with H&E (x 400).
Shah et al. [7] attributed such condition to the reduced proliferations of primordial germ cells during migration to the gonadal ridge and accelerated rate of apoptosis.

It could be concluded that identifying trisomy of autosomal chromosome 24 as a possible cause of testicular hypoplasia in buffalo bulls may aid in elimination of the affected cases in young ages and not wait the age of puberty when the semen could be collected, and the affection could be diagnosed.

References

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