Incorporation of Ascorbic Acid, Caffeine and Chloroquine Diphosphate in Dilutor Improves Structural and Functional Status of Frozen Semen

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Abstract

Effect of supplementation of ascorbic acid, caffeine and chloroquine diphosphate in the dilutor on post-thaw quality of spermatozoa was studied. Thirty-two semen ejaculates (8 each from 2 Holstein Friesian and 2 crossbred; 50% Friesian x 50% Haryana) were studied. Ascorbic acid, caffeine and chloroquine diphosphate were added in Tris-egg yolk-glycerol (control) extender at the concentration of 10 mM, 0.54 mM and 7 mM, respectively. The post-thaw semen parameters studied were structural (acrosomal integrity and livability), functional (motility, per cent sperm responding to hypo-osmotic swelling test, i.e. HOS positive sperm and penetrability of sperm in bovine cervical mucus, i.e. sperm penetration distance-SPD) and measurement of activities of Glutamic oxaloacetic transaminase (GOT), Glutamic pyruvic transaminase (GPT), Alkaline transaminase (AKP) and Acid transaminase (ACP). Certain enzymes in the seminal plasma. The semen from crossbred bull had significantly lower mass activity (P ≤ 0.01), initial progressive motility (P ≤ 0.01), sperm penetration distance (SPD value) (P ≤ 0.01), HOS positive sperm and seminal plasma GOT activity, but higher abnormal sperm. Incorporation of additives in the dilutor significantly improved the post-thaw semen quality, especially the post-thaw motility, post-thaw per cent live sperm, sperm penetration distance and sperm responding to hypo-osmotic solution. The leakage of all the 4 enzymes (GOT, GPT, ACP and AKP) was significantly less in the samples frozen in presence of ascorbic acid and chloroquine as compared to control. Ascorbic acid resulted in maximum improvement to the quality of frozen-thawed semen followed by chloroquine and caffeine. The study concludes that post-thaw quality of frozen semen of HF and crossbred bulls significantly improves if the dilutor is supplemented with 10 mM concentration of ascorbic acid.
Keywords
Semen; Freezing; Ascorbic Acid; Caffeine; Chloroquine; Sperm Penetration Distance; Hypo-Osmotic Swelling

1. Introduction
Artificial insemination has attained a stage of being an indispensable and integral component of animal reproduction technologies. Its adoption has made a rapid progress in livestock production, mainly aimed at increasing milk production of bovines in many parts of the world. The population of exotic and their crossbred cattle in India has been ever increasing since last few decades; therefore demand for frozen semen of pure and crossbred dairy cattle is also increasing. However, the production and supply of frozen semen with an acceptably predictable fertility has always been a challenge because of poor fertility of cryopreserved semen [1] [2].

Cryopreservation is known to degrade the potential fertility of the sperm cells by causing death of about 50 per cent of the cells and altered characteristics of many of the remaining cells [3]. The damages generally are the consequences of mechanical and osmotic phenomenon, oxidative stress, increased membrane permeability, lipid peroxidation and subsequent membrane damage during cooling, freezing and thawing. Reactive oxygen species (ROS) and other intermediate free radicals are produced in more quantities during cooling [4]. Moreover, the levels of antioxidant defenses are decreased in bovine spermatozoa after a cycle of freezing and thawing [5]. Free oxygen radicals play a major role in lipid peroxidation as well as in protein damages that leads to cell death [6] or altered characteristics (as if the spermatozoa are capacitated or acrosome reacted) of the surviving cells [7]. Recent observations show that addition of antioxidants such as ascorbate and alfa-tocopherol to dilutors improves sperm quality by exerting protective effect on both metabolic activity and cellular viability of frozen bovine semen [8] [9].

Ascorbic acid is known to improve post-thaw motility and live percent of buffalo spermatozoa [10]. Caffeine, a metabolic stimulant enhances motility, respiration and fructolysis of epididymal and ejaculated spermatozoa of cattle and buffalo bulls [11]-[13]. It has also been reported to improve progressive motility of both fresh and preserved sperms of various species [14] [15]. Chloroquine diphosphate, a membrane stabilizer is known to stimulate the respiration and motility of fresh and aged spermatozoa in vitro [16] and to improve the post thaw quality of frozen semen [17].

Increasing evidences of beneficial effects of above compounds inspired us to investigate further in a more comprehensive way their effect on structural and functional status of the frozen semen of Holstein Friesian and crossbred (Holstein Friesian x Haryana) bull with an objective to ascertain their role, if any, in improving the potential fertility of frozen-thawed sperm.

2. Materials and Methods
Source of Semen: Semen samples for the experiment were collected using artificial vagina from 2 purebred Holstein Friesian and 2 crossbred (Holstein Friesian x Haryana) bulls maintained at Germ Plasm Center, Indian Veterinary Research Institute, Izatnagar, (I.V.R.I.) India. Thirty-two ejaculates (8 from each bull) were collected, processed, frozen and evaluated for various seminal characteristics.

Semen Evaluation: Each semen sample was examined for routine semen parameters (volume, color, per cent live sperm, total sperm concentration and abnormal sperm as per the standard methods described [18]. The selected ejaculates (having + 3 mass activity) were divided into three parts; part one was used for dilution and freezing, part two for in vitro fertility tests [Bovine cervical mucus penetration test (BCMPT) and Hypo—osmotic swelling test (HOST)] and part three for the separation of the seminal plasma for assaying fresh stage activity of GOT, GPT, ACP and AKP. The seminal plasma was separated by centrifugation (1100 g for 20 min.) both from fresh and frozen-thawed samples preserved in sterilized micro centrifuge tubes at −20°C until the assessment of the activities of GOT, GPT, ACP and AKP were measured.

Percent Intact Acrosome: The percent acrosomal integrity of sperm was determined in fresh and post—thawed semen (200 sperm cells) by preparing smears stained with Giemsa. In Figures 1, 2, 3 and 4 respectively, the denuded/lost acrosome, elongated sperm head with lost acrosome and ruffled acrosome using Giemsa stain
have been shown. Various types of acrosome reaction with different stages with live and dead sperms are shown in plate 4 using FITC-PSA and PI stain.

**Dilution and Freezing:** Semen aliquot was further divided in four groups with and without additives and extended in Tris-fructose-yolk-glycerol dilutor (TFYG); [Tris Hydroxy-methyl-aminomethane—254 mM, citric acid monohydrate—78 mM, Fructose anhydrous—70 mM, Glycerol—7.0 ml, Penicillin—1000 IU/ml and Streptomycin 1000 microgram/ml containing 10% egg yolk added at the time of use]. Thus prepared dilutor gave an osmolality of ~320 milli osmol. This dilutor without additives was used as a control one (T₀) and treatment dilutors were prepared as follows:

- T1—TFYG + ascorbic acid 10 mM. (Merck Chemicals, Mol. Wt. 176.13)
- T2—TFYG + caffèine 7 mM (Sigma Chemicals, Mol. Wt.)
- T3—TFYG + Chloroquine diphosphate 0.54 mM (Sigma Chemicals, Mol. Wt. 319.88)

These concentrations of additives were determined based on a pilot study taken up in a comprehensive project work at our Laboratory at I.V.R.I., Izatnagar; using many other concentrations of these additives separately in TFG (control) on post-thaw quality of semen (motility and live percent).

The semen samples were extended at 35˚C to the extent that each ml of diluted semen contained 50 million motile spermatozoa. 0.5 ml medium French straws for the filling of semen were used. The filled straws were kept in a bread box half filled with water at 35˚C. This box was kept in a refrigerator [freezer chamber] for 2 hr. And when the temperature of straws reached 5˚C they were transferred to a cold cabinet (4˚C) where these straws were dried on a blotting paper, spread evenly over the rack and kept horizontally in liquid nitrogen vapor till the temperature of the straws reached ~120˚C. The straws were held for 10 min. At this temperature and then plunged into the liquid nitrogen.

**Post-Thaw Evaluation:** After 24 hr, storage in liquid nitrogen straws were thawed at 37˚C for 30 sec. Post-thaw per cent live sperm and per cent intact acrosome were determined. A part of the post-thawed semen from all the treatment groups was used for assessment of in vitro fertilizing ability using cervical mucus penetration test and HOS test. The enzymatic activities were determined after separating the seminal plasma from the frozen semen.

**2.1. Tests for in Vitro Fertilizing Ability**

**Cervical mucus penetration test:** Bovine cervical mucus was collected aseptically from estrus cows brought for insemination at Institute Polyclinic. Mucus samples showing typical crystallization fern pattern and showing negative results in white side test, indicating absence of obvious infection [19] were taken for the experiment.

Non-heparinized hematocrit capillary tubes (8 cm in length) were loaded with the mucus with the help of a syringe and a rubber tube. One end of the capillary tube was sealed with hemoseal (Shandilya Chemicals, India). In a separate micro centrifuge tube, 0.5 mL of semen samples (fresh and post-thawed) were placed (in duplicate) and the open end of the loaded tubes was dipped in the semen vertically. After 60 min of incubation at 37˚C the capillary tubes were cleaned, wiped, placed on a graduated glass slide and observed under the microscope for the distance traveled in mucus by the vanguard spermatozoa. The semen samples were graded on the basis of the distance traveled by the vanguard spermatozoa in 60 min as follows [20].

- Excellent more than 30 mm.
- Good more than 20 mm but less than 30 mm.
- Medium 12 to 20 mm.
- Poor less than 12 mm.

**Hypo-osmotic swelling test (HOST):** In micro-tubes containing 1.0 ml of hypo-osmotic solution (sod. citrate 25 mM, fructose 75 mM added to 100 mL double distilled water with an osmolality of approximately 100 mM) 0.1 mL of fresh and frozen-thawed semen samples were incubated at 37˚C for 60 min. After incubation, a small drop from the bottom of the tube was placed on a slide, covered with a coverslip and examined for different swelling pattern as follows [21]:

- Pattern A: no swelling, no membrane reaction
- Pattern B: swelling of the tip of the sperm tail
- Pattern C: different types of hairpin like swellings of the mid piece
- Pattern D: complete swelling of the tail region

A total of 200 cells were counted. The sperm cells showing Pattern B, C and D were considered positive for the HOST test.
2.2. Enzymes Assay

Seminal plasma from fresh and frozen-thawed samples was separated by centrifugation at 1100 g for 20 min. and preserved at −20°C till the enzyme activity was determined. The activity of glutamic oxaloacetic and pyruvic transaminases (GOT and GPT) was estimated according to the method described by [22]. The activity of alkaline phosphatase (AKP) and acid phosphatase (ACP) was determined following the method of [23] using Qualigen’s Kit.

2.3. Statistical Analysis

Means and standard errors were calculated. Analysis of variance (ANOVA) was used for comparing the effects of different factors (bulls, breeds, treatments etc.) and means were compared using Duncan’s Multiple Range Test (DMRT) as per [24] on Wipro-Pentium computer of the Institute.

3. Results

3.1. Fresh Semen Quality

The mean (± SE) cytomorphological seminal characteristics and functional attributes of fresh semen of both Holstein Friesian (HF) and crossbred bulls are presented in Table 1. The crossbred bull semen had a significantly lower mass activity (P ≤ 0.01), individual motility (P ≤ 0.01), sperm penetration distance in cervical mucus (P ≤ 0.01), per cent HOS positive sperm (P ≤ 0.01) and plasma GOT level (P ≤ 0.01) and higher abnormal sperm per cent (P ≤ 0.01) than those of HF bull.

Volume of semen differed significantly (P < 0.01) between two HF bulls. Mass motility (0 - 5 scale) differed significantly (P < 0.01) between two breeds. The per cent individual motility differed significantly (P < 0.01) between 2 HFxH (P < 0.01) and 2 HF bulls (P < 0.05). Similarly the per cent live sperm differed significantly between the two breeds (P < 0.01) and between two HF bulls (P < 0.05). A significant difference (P < 0.01) in the total per cent abnormal sperm was also found between 2 HF bulls (P < 0.01). The population of sperm with intact acrosome differed significantly (P < 0.01) between breeds and between 2 HF bulls (P < 0.05). Besides these seminal characteristics the total sperm concentration per unit did not differ between breeds and bulls (Table 1).

Analysis of variance revealed a significant difference (P < 0.01) in the GOT activity of fresh semen between

<table>
<thead>
<tr>
<th>Table 1. Cyto-morphological and functional attributes (Mean ± SE) of fresh semen of HF and crossbred bulls.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Seminal characteristics</strong></td>
</tr>
<tr>
<td>Mass motility (%)</td>
</tr>
<tr>
<td>Volume (ml)</td>
</tr>
<tr>
<td>Individual motility (%)</td>
</tr>
<tr>
<td>Live sperm count (%)</td>
</tr>
<tr>
<td>Total abnormal sperm count (%)</td>
</tr>
<tr>
<td>Intact acrosome (%)</td>
</tr>
<tr>
<td>Sperm concentration (millions/ml)</td>
</tr>
<tr>
<td>BCMPT distance (nm)</td>
</tr>
<tr>
<td>HOST positive sperms (%)</td>
</tr>
<tr>
<td>GOT (micro mole/L)</td>
</tr>
<tr>
<td>GPT (micro mole/L)</td>
</tr>
<tr>
<td>AKP (KAU/100ml)</td>
</tr>
<tr>
<td>ACP (KAU/100ml)</td>
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</tbody>
</table>

Mean bearing superscript a, b differ significantly (P < 0.01). Means bearing superscript A, B differ significantly (P < 0.05).
the 2 breeds, but within the breeds between the bulls the difference was non-significant (Table 2). The GPT activity, however, did not differ significantly between breeds and bulls. The AKP activity in the fresh seminal plasma did not differ between the breeds but it differed significantly (P < 0.05) between 2 HFxH bulls. No variation in the ACP activity in the fresh seminal plasma of either between breeds or between bulls within the breeds was found.

Comparison of distance traveled by vanguard sperm (the sperm moving farthest distance) in the cervical mucus, by the sperm of both the breeds revealed that the SPD varied significantly (P < 0.01) between breeds and bulls. The HF sperm traveled a longer distance than HFxH sperm. The per cent total sperm responding to hypo-osmotic swelling test was significantly (P < 0.01) higher for HF bull semen than HFxH semen. This parameter also differed significantly (P < 0.01) between the 2 HFxH bulls.

3.2. Post-Thaw Quality

Analysis of data for the pooled means of various seminal characteristics of HF and HFxH bulls after freezing thawing using DMRT Test revealed that post-thaw motility between the two breeds of bulls did not differ significantly (Table 2) but within the breed between bulls it differed significantly (P < 0.01; Table 3).

The pooled data on various seminal characteristics showing the effect of different additives are presented in Table 4. Perusal of this data indicates that almost all the parameters which are responsible for adjudging the quality of semen were significantly (P < 0.05) higher in the treatment group of ascorbic acid than other two additives. Similarly the leakage of enzymes was significantly low in this group of ascorbic acid than other two additives (Table 4). More interestingly it was noted that as compared to control group, in the caffeine treated group the enzymes leakage was higher (although not significantly) but in the ascorbic acid and chloroquine

<table>
<thead>
<tr>
<th>Table 2. Cytomorphological and functional attributes (Mean ± SE) of frozen thawed semen of HF and crossbred bulls.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Seminal characteristics</strong></td>
</tr>
<tr>
<td>Post-thaw motility %</td>
</tr>
<tr>
<td>Post-thaw livability %</td>
</tr>
<tr>
<td>Post-thaw intact acrosome (%)</td>
</tr>
<tr>
<td>Post-thaw BCMPT distance (mm)</td>
</tr>
<tr>
<td>Post-thaw HOST positives sperms (%)</td>
</tr>
<tr>
<td>Post-thaw GOT (micro mole/L)</td>
</tr>
<tr>
<td>Post-thaw GPT (micro mole/L)</td>
</tr>
<tr>
<td>Post-thaw AKP (KAU/100ml)</td>
</tr>
<tr>
<td>Post-thaw ACP (KAU/100ml)</td>
</tr>
</tbody>
</table>

The means bearing superscript a, b differ significantly (P < 0.01).

<table>
<thead>
<tr>
<th>Table 3. Mean sum of squares showing influence of breeds, bulls and treatments on cytomorphological and functional characteristics and seminal plasma enzymes level of frozen-thawed spermatozoa.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sources of variation</strong></td>
</tr>
<tr>
<td>Between breeds</td>
</tr>
<tr>
<td>Between treatments</td>
</tr>
<tr>
<td>Between HF bulls</td>
</tr>
<tr>
<td>Between FxH bulls</td>
</tr>
<tr>
<td>Breed x treatment</td>
</tr>
</tbody>
</table>

**Significant (P < 0.01); *Significant (P < 0.01).
group the enzyme leakage was less than the control. This table further reveals that the increase in the post-thaw motility was almost 8% with ascorbic acid, 10% with caffeine and 7% with chloroquine. The increase in the percent post-thaw live sperm was 17%, 11% and 8%, respectively for ascorbic acid, caffeine and chloroquine. The percent increase in the intact acrosome was about 11%, 5% and 3% in presence of ascorbic acid, caffeine and chloroquine, respectively. The sperm traveled an increased distance of about 9%, 5% and 4% in presence of ascorbic acid, caffeine and chloroquine, respectively. The response of sperm cells to HOST was also on the similar lines and the increased percent for ascorbic acid, caffeine and chloroquine, respectively were 9%, 5% and 4%.

3.3. Correlation among Semen Parameters (Post-Thawed)

Post-thaw motility showed a strong positive correlation with the parameters like livability (r = 0.86, P < 0.01), intact acrosome (r = 0.74, P < 0.01), SPD (r = 0.73, P < 0.01) and HOST response (r = 0.72, P < 0.01) (Table 5) as expected. This relationship of motility along with the positive correlation of post-thaw intact acrosome (PTIA) with SPD (r = 0.67, P < 0.01) and HOST response (r = 0.66, P < 0.01) supports the approach of using three independent parameters of sperm structure and function as an indicator of cryopreservation success in examining the quality of frozen semen. The AKP was also positively correlated with ACP value (r = 0.88, P < 0.01).

Table 4. Cytomorphological, functional and some biochemical attributes (Mean ± SE) of frozen-thawed semen irrespective of breed and bulls.

<table>
<thead>
<tr>
<th>Seminal characteristics</th>
<th>Control</th>
<th>Ascorbic acid</th>
<th>Caffeine</th>
<th>Chloroquine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Motility (%)</td>
<td>29.2 ± 5.2c</td>
<td>47.8 ± 5.8a</td>
<td>38.9 ± 4.7b</td>
<td>36.1 ± 5.0b</td>
</tr>
<tr>
<td>Livability (%)</td>
<td>55.2 ± 4.6c</td>
<td>72.6 ± 3.1a</td>
<td>66.6 ± 3.6e</td>
<td>63.1 ± 4.2e</td>
</tr>
<tr>
<td>Intact acrosome (%)</td>
<td>69.2 ± 4.4f</td>
<td>80.3 ± 3.5a</td>
<td>74.1 ± 3.9g</td>
<td>72.8 ± 3.9b</td>
</tr>
<tr>
<td>BCMPT distance (mm)</td>
<td>19.0 ± 2.2e</td>
<td>28.3 ± 2.4a</td>
<td>24.7 ± 2.3d</td>
<td>23.6 ± 2.6b</td>
</tr>
<tr>
<td>HOS positive sperms (%)</td>
<td>21.4 ± 2.0f</td>
<td>30.4 ± 3.0a</td>
<td>26.8 ± 2.4d</td>
<td>25.7 ± 2.4b</td>
</tr>
<tr>
<td>GOT (micro mole/l)</td>
<td>384.0 ± 100.2c</td>
<td>285.1 ± 86.4a</td>
<td>399.6 ± 106.6e</td>
<td>323.4 ± 87.6a</td>
</tr>
<tr>
<td>GPT (micro mole/l)</td>
<td>203.1 ± 80.4a</td>
<td>134.6 ± 55.4b</td>
<td>214.0 ± 102.3s</td>
<td>153.7 ± 71.0a</td>
</tr>
<tr>
<td>AKP (KAU/100ml)</td>
<td>383.9 ± 164.8d</td>
<td>314.6 ± 141.2b</td>
<td>420.4 ± 186.1a</td>
<td>345.7 ± 146.7b</td>
</tr>
<tr>
<td>ACP (KAU/100ml)</td>
<td>515.0 ± 246.1</td>
<td>432.1 ± 195.3</td>
<td>561.4 ± 250.3</td>
<td>464.1 ± 181.7</td>
</tr>
</tbody>
</table>

Means bearing different superscripts (a–h) in a row are significantly different (P < 0.05), separately for each treatment.

Table 5. Correlation coefficients among cytomorphological, functional and biochemical attributes of post thaw semen in tris dilutor (HF and FXH pooled) without additives.

<table>
<thead>
<tr>
<th>Seminal characteristics</th>
<th>PTM</th>
<th>PTL</th>
<th>PTIA</th>
<th>PT-SPD</th>
<th>PT HOST</th>
<th>PT-GOT</th>
<th>PT-GPT</th>
<th>PT-AKP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Motility % (PTM)</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Livability % (PTL)</td>
<td>0.86**</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intact acrosome % (PTIA)</td>
<td>0.74**</td>
<td>0.65**</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BCMPT dist. mm. (PT SPD)</td>
<td>0.73**</td>
<td>0.69**</td>
<td>0.67**</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Host Positive % (PT HOST)</td>
<td>0.72**</td>
<td>0.71**</td>
<td>0.66**</td>
<td>0.92**</td>
<td>1.00</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GOT Level umole/L</td>
<td>0.13</td>
<td>0.02</td>
<td>0.24</td>
<td>0.26</td>
<td>0.18</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GPT level umole/L</td>
<td>−0.02</td>
<td>−0.12</td>
<td>0.06</td>
<td>0.20</td>
<td>0.17</td>
<td>0.18</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>AKP level KAU/100ml</td>
<td>0.28</td>
<td>0.10</td>
<td>0.09</td>
<td>−0.02</td>
<td>0.08</td>
<td>−0.14</td>
<td>−0.09</td>
<td>1.00</td>
</tr>
<tr>
<td>ACP level KAU/100ml</td>
<td>0.03</td>
<td>−0.07</td>
<td>−0.02</td>
<td>−0.24</td>
<td>−0.13</td>
<td>−0.45**</td>
<td>−0.11</td>
<td>0.76**</td>
</tr>
</tbody>
</table>

PT = Post-Thaw, KAU = King and Armstrong Unit. *Significant at 5% level; **Significant at 1% level.
In general, the trend of correlations of seminal parameters, even after addition of these substances in the dilu-
tor (i.e. with treatments) remained the same as in control dilutor (presented in Table 4), but the values differed
to some extent. The post-thaw motility (PTM) was significantly positively correlated with post-thaw livability
(PTL) \( r = 0.86, 0.57, 0.59 \) and 0.59 for \( T_0, T_1, T_2 \) and \( T_3 \) treatments respectively), PTIA \( r = 0.74, 0.78, 0.73 \) and 0.58 for \( T_0, T_1, T_2 \) and \( T_3 \) treatments respectively) and SPD value \( r = 0.73, 0.60, 0.58 \) and 0.45 for \( T_0, T_1, T_2 \) and \( T_3 \) treatments respectively). PTM was also significantly positively correlated with HOST response \( r = 0.72, 0.67 \) and 0.55 \((P < 0.01) \) for \( T_0, T_1, T_2 \) and \( r = 0.38 \) \((P < 0.05) \) for \( T_3 \) group].

4. Discussion
Crossbred bulls of different genetic make-up are usually reported to have different semen characteristics [25]-
[28]. Total sperm abnormalities were higher in crossbred bull than HF bull semen which had also been reported
earlier [29]. The perpetuation of this problem is inherent which can be attributed to the fact that even in well or-
ganized dairy farms, sperm parameters as a trait receive less priority in the selection criteria for the cross breed-
ing of animals under Indian management conditions. The differences in seminal cyto-morphological characteris-
tics between the two breeds of bulls are normally expected and our values are in agreement with the earlier re-
ports from bulls of different genetic make-up [26]-[28]. These findings confirm the inferiority of crossbred bull
semen in many respects as compared to semen from pure bred bulls.

Perusal of data of the frozen thawed semen from these bulls revealed that post-thaw parameters such as motil-
ity, livability, intact acrosome, BCMPT distance and percent HOST positive sperm were lower than desirable
(Table 2). On these samples (control) intact acrosome per cent was significantly lower in crossbred (HFxH)
bulls than that of pure HF bulls. Acrosomal changes are highly correlated with fertility of frozen-thawed semen
[30] and are simpler and easier method to evaluate the effect of many variables on the spermatozoa. A signifi-
cant relationship between sperm motility and acrosomal integrity with fertility has been reported [31]. However,
Cumming [32] observed that despite the effect that the bulls could be classified into three statistically significant
groups according to their non return rates, there was no significant correlation between the percentage of intact
acrosome and the non return rates of bulls. In previous report on cattle and buffalo bulls [33] acrosomal intact-
ness is reported to be significantly affected by the dilutor composition in frozen semen. Possibly this explains
our findings of differences in the per cent intact acrosome in presence of different additives, which definitely al-
tered the composition of dilutor. Ascorbic acid was superior in maintaining better acrosomal intactness in frozen
samples than caffeine and chloroquine.

Variation in post-thaw per cent intact acrosome (PTIA) could be due to differences in response of the cell
membranes to freezing that can be related to the defects in spermatogenesis [34]. A decline in the intact acro-
some percent in the frozen thawed spermatozoa has also been reported earlier in cattle bull [29] [35] [36] and
buffalo bull semen [33] [37].

The leakage of enzymes from sperm cell into the seminal plasma was very severe. Compared to fresh samples
more than 8 fold increases in enzymatic activities of the GOT and GPT and almost 2 folds of ACP and AKP in
the frozen-thawed seminal plasma was observed. This increase in enzymatic activity in the frozen-thawed se-
mental plasma indicates increased cell membrane permeability and damages to the sperm cells during the pro-
cess of freezing and thawing. However, minimum leakage of these enzymes during the process of freezing-
thawing in presence of ascorbic acid than caffeine and chloroquine suggests that the detrimental actions of ROS
were prevented especially on the membrane structure by ascorbic acid.

Assessment of post-thaw semen based on cytomorphology, sperm function (motility, HOST response and
SPD and some enzymatic constituents; phosphatases and transminases) in the seminal plasma after the addition
of ascorbic acid, caffeine or chloroquine indicated that there was a significant improvement in the quality of se-
men both in HF and crossbred bull semen. Ascorbic acid when added, in the semen, is known to enhance motil-
ity of bull sperm [40] and buffalo sperms under refrigerated temperature for 120 hours increase post-thaw sperm
motility and livability [38]. However, Sanchez-Partida et al. [39] observed no effect on motility after addition of
ascorbic acid at the concentration of 25 mM. Addition of ascorbic acid resulted in about 18% increase in the
post-thaw motility in the present study. Improvement in the post-thaw motility after adding ascorbic acid in the
dilutor has been reported [40]. The better effect of ascorbic acid than caffeine and chloroquine, in the present
study, might be because of low sperm damage in presence of ascorbic acid as it has been reported to protect the
spermatozoa from the peroxidase damage [41].
Reactive oxygen species (ROS) are continuously produced in the ejaculates by spermatozoa and accompanying leukocytes [42] [43] impairing spermatozoal metabolism and affecting their fertilizing ability [44]. Any substance present in the dilutor, which can prevent the action of ROS, may improve the quality of semen. Ascorbic acid protects spermatozoa from free radical damage and improves their motility and viability [45]. On these lines the better effect of ascorbic acid in improving the post-thaw quality of frozen semen of both the breeds in the present study are justifiable.

One of the objectives of present study was to find out the effects of the additives on both HF and crossbred bull semen and see if there was any interaction between the breeds and the treatment. There was no any interaction between the breed and additives for all the investigated parameters (Table 3). The analysis of data of all the four bulls pooled together (Table 4) revealed that incorporation of all the three additives resulted in significantly higher (P < 0.01) values for percent motility, livability, intact acrosome, BCMPT distance and HOST response than that of control; ascorbic acid showing the best and caffeine and chloroquine moderate improvement in the quality of the frozen semen. The effect of chloroquine and caffeine is comparable for all the parameters. Caffeine inhibits cAMP and cGMP which appears to be responsible for enhancing sperm motility [11]. cAMP acts directly on the plasma membrane of sperm cells and thereby regulates the exchange of inorganic phosphate and calcium ions resulting in improvement in sperm motility and integrity [34]. Incorporation of caffeine also improved the per cent post-thaw intact acrosome in present study, but the values were lower than ascorbic acid. Since considerable damage is inflicted on the sperm plasma membrane during cryo-storage, caffeine may compensate for this damage by stimulating the activity of an outwardly directed plasma membrane calcium pump through the mediation of cAMP [34]. However, prevention of damages to plasma membrane in caffeine group was poor than ascorbic acid and chloroquine (Table 4), possibly because caffeine is a stimulant and once spermatozoal metabolism is increased in presence of caffeine such results are expected.

The incorporation of chloroquine in the extender has been found to increase sperm viability after cryopreservation [17]. However, chloroquine did not improve post-thaw motility of boar semen [46]. The findings that chloroquine diphosphate might be useful to prevent the leakage of vital enzymes [47] support our results. In presence of chloroquine diphosphate in the dilutor although an increase in post-thaw motility in the present study, as well as reported earlier [16], was recorded but this increase was less as compared to ascorbic acid and caffeine. [16] reported an increase in the post-thaw motility after incorporating chloroquine. Chloroquine a membrane stabilizer and anti-inflammatory agent is reported to stimulate respiration and motility [16] through directly or indirectly acting on adenyl cyclase system.

Incorporation of additives resulted in lower enzyme leakage in the extracellular fluid as evident from reduced activities of GOT, GPT, AKP and ACP in frozen-thawed seminal plasma of the treatment group than that of in the control group (Table 4). In the seminal plasma minimum enzymatic activities were observed in the presence of ascorbic acid followed by chloroquine and caffeine when compared with the control group. The difference in values for transaminaes (GPT and GOT) for caffeine added and control groups was non-significant but the former had a slightly higher value. The leakage of and ACP was also maximum in the caffeine added group implying that enhanced metabolic activity during cooling, freezing and post-thawing causes the membrane to be more permeable and prone to damages.

It is now known that cryopreservation induces formation of reactive oxygen species (ROS) that are detrimental to spermatozoa [48] [49]. Ascorbic acid through its reducing properties [50] might have minimized the probable oxidative damages to the membranes by ROS thus preventing to some extent the enzyme leakage.

It is apparent from the study that ascorbic acid, chloroquine diphosphate and caffeine treated dilutor altered the structural and functional status of the frozen spermatozoa when compared with that of control. All the three compounds used as additives produced desirable effect on frozen thawed spermatozoa by increasing live, motile sperm and HOST responsive cell percentage and by reducing the enzyme leakage from the sperm cells during cryopreservation. Among the three compounds ascorbic acid brought the best desirable changes followed by chloroquine di-phosphate. It could, therefore, be concluded that addition of ascorbic acid at the concentration of 10 mM exerts beneficial effect and help improve the quality of frozen semen of pure HF and crossbred (HFxH) bulls.

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Appendix

Figure 1. Denuded/lost acrosome.

Figure 2. Elongated head with lost acrosome.

Figure 3. Ruffled crosome.

Figure 4. Sperm cells showing different stages of acrosome reacted live, acrosome reacted dead, intact acrosome live and intact acrosome dead sperm cell.