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1 *Original Research Paper*

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2 **Evaluation of pigeon egg yolk for post thaw quality, enzyme leakage and**  
3 **fertility of buffalo (*Bubalus bubalis*) bull spermatozoa**

4  
5 S. Akhter<sup>a</sup>, B.A. Rakha<sup>b</sup>, M.S. Ansari<sup>c</sup>, S. Iqbal<sup>d</sup>, M. Khalid<sup>e</sup>

6  
7 <sup>a</sup> *Department of Zoology, Pir Mehr Ali Shah Arid Agriculture University Rawalpindi-46300,*  
8 *Pakistan*

9 <sup>b</sup> *Department of Wildlife Management Pir Mehr Ali Shah Arid Agriculture University*  
10 *Rawalpindi-46300, Pakistan*

11 <sup>c</sup> *Department of Zoology, University of Sargodha-Lyallpur Campus-38000, Pakistan*

12 <sup>d</sup> *Semen Production Unit, Qadirabad, Sahiwal, Pakistan*

13 <sup>e</sup> *Department of Pathobiology and Population Sciences, Royal Veterinary College, University*  
14 *of London, UK*

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16 **Running Title:** Pigeon egg yolk for cryopreservation of buffalo semen

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19 **\* Corresponding Author:**

20 Dr. Shamim Akhter

21 Associate Professor

22 Department of Zoology

23 Pir Mehr Ali Shah Arid Agriculture University, Rawalpindi-46300, Pakistan

24 Email: [sashraf1993@gmail.com](mailto:sashraf1993@gmail.com)

25 Cell: + 923335206056

## 26 ABSTRACT

27 Egg yolk is used as a cryoprotectant in semen preservation. However, its composition varies  
28 according to the species which may influence its effectiveness during the freeze-thaw process.  
29 Therefore, study was conducted to identify the optimum level of pigeon egg yolk (PEY) in *Tris*  
30 citric acid (TCA) extender for freezability and *in vivo* fertility of buffalo semen. Semen was  
31 collected at weekly intervals for a period of three weeks (replicates) from 6 Nili Ravi buffalo  
32 bulls (2 ejaculates/bull/ replicate) and diluted with TCA extender ( $50 \times 10^6$  motile spermatozoa  
33  $\text{ml}^{-1}$ ) containing 5%, 10%, 15% and 20% PEY or 20% CEY (control) and cryopreserved. Post-  
34 thaw sperm quality and extracellular enzymes leakage was assessed after thawing. Sperm  
35 motility, plasma membrane integrity, livability and viability was significantly higher in  
36 extenders containing 10% and 15 % PEY compared to 5% PEY, 20% PEY or 20 % CEY  
37 (controls). A dose-dependent decrease was recorded in the chromatin damage for the PEY,  
38 being lowest for the 15% and 20% PEY which was significantly less compared to controls  
39 (20%CEY). The extracellular GOT and LDH leakage was significantly lower ( $P < 0.05$ ) in  
40 extender containing 10% and 15 % PEY compared to the controls. Semen collected from 2  
41 bulls and cryopreserved in extenders containing 15% PEY or 20% chicken egg yolk was  
42 assessed after AI. A total of 400 buffaloes were inseminated (100 inseminations/extender/bull).  
43 The overall fertility rate was significantly higher ( $P < 0.05$ ) with semen cryopreserved in  
44 extender containing 15% PEY (56 %) compared to 20% CEY (42 %; controls). In conclusion,  
45 pigeon egg yolk at 15 % offers advantages over 20% chicken egg yolk in terms of *in vitro* post-  
46 thaw semen quality and *in vivo* fertility of buffalo.

47

48 **Keywords:** *Columba livia domestica*; *bubalus bubalis*; extender; Lactic dehydrogenase,  
49 Glutamic oxaloacetic transaminase

50

## 51 **1. Introduction**

52 Egg yolk is a very valuable source of vital nutrients usable in medical, pharmaceutical,  
53 nutraceutical and biotechnological industries. Its cryoprotective properties were identified for  
54 the first time by Philips and Lardy [1] and since then it has been used as impermeable  
55 cryoprotectant for sperm across the species. The cryoprotectant role of egg yolk has been  
56 attributed to its low density lipoproteins (LDLs) fraction that includes proteins, lipids,  
57 phospholipids, cholesterol, and both saturated and unsaturated fatty acids [2]. In addition to  
58 LDLs, the other vital components in the egg yolk include vitamins, minerals/ trace elements,  
59 antibodies and antioxidants [3,4], all of which are of immense significance for sperm.

60 Egg yolks from different avian species differ in their composition, especially in the  
61 content of cholesterol, fatty acids, phospholipids [5,6], amino acids [7] and trace elements [8].  
62 Levels of omega-3 polyunsaturated fatty acids, vitamin E, carotenoids, and Se in eggs not only  
63 show profound interspecies variations, but they are also markedly higher in free-living species  
64 compared with their domesticated or captive counterparts [8,9]. Moreover, a substantial (up to  
65 three fold) variation for the trace element content has been reported in eggs from different  
66 domestic avian species that were offered same feed [8]. As, such differences in the composition  
67 of egg yolks from different species may potentially affect their effectiveness for sperm  
68 cryoprotection [10], attempts have been made to identify egg yolk sources most suitable for  
69 sperm cryopreservation. It is imperative to mention that egg yolk from avian species other than  
70 chicken were found more beneficial for cryopreservation of stallion [6,11], jackass [12], bull  
71 [13,14], ram [15] and buffalo sperm [16]. Very recently turkey and quail egg yolk have been  
72 tested as cryoprotectants for buffalo sperm cryopreservation; they not only improved the sperm  
73 cryopreservation but were also able to improve in vivo fertility compared to traditionally used  
74 chicken egg yolk. Moreover, the results obtained in this study showed that in the presence of  
75 turkey and quail egg yolk there was a significant decrease in the extracellular enzyme leakage

76 from buffalo sperm which led the authors to suggest that the cryoprotective abilities of the  
77 turkey and quail egg yolk are linked to their variable cholesterol or phospholipid contents [17].

78 The principal protective component of the egg yolk is the lecithin that can replace some  
79 of phospholipids in sperm plasma membrane and form a protective film thereby decreasing its  
80 damage during cooling [18]. Pigeon egg yolks have higher lecithin content compared to  
81 chicken egg yolk (4.86 % Vs 2.94%) [19]. In addition, the high density lipoproteins (HDLs)  
82 that are antagonistic to cryoprotection and the yolk granules that hinder sperm motility are less  
83 in pigeon compared to other egg yolks [18]. When the trace elements were studied in different  
84 avian egg yolks, interestingly, the pigeon egg yolk was found superior in having Selenium (Se)  
85 [8] that has been positively correlated with fertility in humans and other species [20,21]. Pigeon  
86 egg yolk LDLs had the best cryoprotective effects on frozen-thawed boar sperm than the hen,  
87 duck, quail, pigeon and ostrich egg yolks' LDLs [18] and when whole egg yolk was compared  
88 with other avian species, pigeon egg yolk offered better protection to bull [13] and ram sperm  
89 [22]. Conventionally, chicken egg yolk is used at 20 % in TCA extender for buffalo semen [23-  
90 26], while few studies investigated optimal levels of other avian egg yolks in comparison to 20  
91 % chicken egg [17,27]. Therefore, present study was designed to investigate the optimal level  
92 of pigeon egg yolk in in *Tris* citric acid extender by assessing the post thaw sperm quality,  
93 extracellular enzyme leakage and *in vivo* fertility of cryopreserved buffalo bull semen.

## 94 **2. Materials and methods**

### 95 *2.1. Preparation of extenders*

96 Tris-citric acid buffer was used for the semen extender. It was prepared by dissolving  
97 1.56g citric acid (Fisher Scientific, UK) and 3.0g *Tris*-(hydroxymethyl)-aminomethane  
98 (Research Organics, USA) in 73 mL distilled water. The pH of buffer was 7.0 and the osmotic  
99 pressure was 320 mOsmol kg<sup>-1</sup>. Apart from the buffer, the semen extender contained 0.2%

100 (wt/v) Fructose (Scharlau, Spain); 7% (v/v) glycerol (Riedel-deHaen, Germany) and a  
101 combination of antibiotics consisting of streptomycin sulphate (1 mg/mL), procaine penicillin  
102 (300 IU/mL) and benzyl penicillin (Sinbiotic<sup>®</sup>, China) (100 IU/mL). The pigeon egg yolk  
103 (PEY) was added in the semen extender at 5%, 10%, 15% and 20%, while CEY was added at  
104 20% which served as controls.

### 105 2.2. *Semen collection and evaluation*

106 Semen from buffalo bulls were collected with artificial vagina (42°C), and transferred  
107 to the laboratory for initial evaluation (volume, sperm motility and sperm concentration).

108 Sperm progressive motility was assessed with phase contrast microscope at 200X at 37  
109 °C by placing a drop of diluted semen sample on a pre-warmed glass slide covered with a cover  
110 slip [25]. Sperm concentration was measured by taking 1 µL of semen and 200 µL of formal  
111 citrate solution (1 mL of 37% formaldehyde in 99 mL of 2.9% sodium citrate) using Neubauer  
112 haemocytometer (Marienfeld, Germany). Only those ejaculates were selected for further  
113 processing that qualified a minimum standard of 1 mL volume, 60% motility and 0.5 billion  
114 spermatozoa/ml.

### 115 2.3. *Semen processing and cryopreservation protocol*

116 The qualifying semen ejaculates were split into five aliquots for dilution in  
117 experimental extenders containing PEY (5%, 10%, 15% and 20%) or 20% chicken egg yolk  
118 (CEY) (controls). Semen aliquots were diluted in a single step at 37°C with each of the  
119 experimental extenders at  $50 \times 10^6$  motile spermatozoa mL<sup>-1</sup>. Diluted semen was cooled to 4°C  
120 for 2 hours and equilibrated for 4 hours at 4°C before being filled in 0.5 mL French straws  
121 (IMV, France) with a suction pump at 4°C in a cold cabinet (Minitub, Germany). The semen  
122 filled straws were then kept over liquid nitrogen vapours about 5cm above the level of liquid  
123 nitrogen<sub>2</sub> for 10 minutes before plunging into liquid nitrogen (-196 °C). The frozen-straws

124 for each treatment were thawed at 37 °C for 30 seconds in water bath and assessed for post-  
125 thaw semen quality parameters.

## 126 2.5. *Post-thaw sperm assays*

### 127 2.5.1. *Sperm motility*

128 Sperm progressive motility was assessed with phase contrast microscope at 200X at  
129 37°C by placing a drop (5 µL) of semen sample on a pre-warmed glass slide and covered with  
130 a cover slip [25].

### 131 2.5.2. *Sperm plasma membrane integrity*

132 Sperm plasma membrane integrity was assessed by hypo-osmotic swelling (HOS) assay  
133 [28]. Solution for HOS assay consisted of 0.73g sodium citrate and 1.35g fructose dissolved in  
134 100 mL distilled water (osmotic pressure ~190 mOsmol kg<sup>-1</sup>). For assessment, 50 µL of frozen-  
135 thawed semen sample was mixed with 500 µL of HOS solution and incubated for 30-40 min at  
136 37 °C. After that, 5 µL of mixture was placed on a glass slide, covered with cover-slip and  
137 examined using phase contrast microscope (400X). Two hundred spermatozoa per  
138 experimental extender per replicate were examined for their swelling characterized by coiled  
139 tail indicating intact sperm plasma membrane [25].

### 140 2.5.3. *Sperm viability and live/dead ratio*

141 Sperm viability and live/dead ratio were studied by dual staining procedure [26]. Equal  
142 drops of Trypan-blue (MP Biomedicals, Eschwege, Germany) and semen sample were placed  
143 on a glass slide at room temperature, mixed and made into a smear. The smear was air-dried  
144 and fixed with formaldehyde-neutral red for 5 min. The slides were then rinsed with distilled  
145 water after which 7.5% Giemsa stain (Sigma) was applied for 4 hours. The slides were rinsed  
146 with water, air dried and mounted with mounting media. Transparent or light blue sperm were  
147 considered as live while those stained dark blue were considered as dead. Transparent or light

148 blue sperm with clear acrosome were considered viable (live with intact acrosome), while  
149 sperm having a clear dark blue demarcation and blunt ended acrosome were considered non-  
150 viable (dead with damaged acrosome). A total of two hundred spermatozoa per experimental  
151 extender per replicate were evaluated in each smear using a phase contrast microscope (1000X;  
152 Olympus BX20, Tokyo, Japan) separately for live/dead ratio and sperm viability.

#### 153 2.5.4. *Sperm chromatin damage*

154 Sperm chromatin damage was assessed using acridine orange assay [29,30]. Smears of  
155 semen were prepared on glass slides, air-dried and fixed in Carnoy's solution (methanol and  
156 glacial acetic acid in a 3:1 proportion) overnight. The slides were air-dried and incubated in  
157 tampon solution (80 mmol/L citric acid and 15 mmol/L Na<sub>2</sub>HPO<sub>4</sub>, pH 2.5) at 75°C for 5  
158 minutes to test DNA integrity. The slides were then stained with acridine orange (0.2 mg/mL),  
159 washed with water to remove background staining and while still wet, covered with cover slips  
160 and evaluated with a epifluorescence microscope (480/550 nm excitation/barrier filter). Sperm  
161 with normal DNA presented green, whereas those with an abnormal/damaged DNA presented  
162 fluorescence that varied from yellow-green to red in spectrum. One hundred sperm cells were  
163 analyzed for each semen sample.

#### 164 2.5.5. *Biochemical tests*

165 Sperm cells with damaged membranes lose their essential metabolites and enzymes. To  
166 check this damage, the levels of two intracellular enzymes Lactic dehydrogenase (LDH) and  
167 Glutamic oxaloacetic transaminase (GOT) were studied as described by Dhimi and Sahni [31].  
168 For this purpose, the 2 mL thawed semen sample was centrifuged at x 166g for 20 min and the  
169 supernatant was separated to analyze for the extra cellular release of LDH and GOT. For LDH  
170 (IU/l) analysis, 20 µL of supernatant was mixed with 400 µL lactate and 100 µL reagent NDH  
171 (Merckmillipore®) in a 5 mL tube and allowed to stand for 10 seconds to complete the reaction.  
172 For GOT, 50 µL of the supernatant was mixed with 400 µL of TRIS, L-Aspartate of MDH



173 (malate dehydrogenase) in a 5 mL tube. For LDH (lactate dehydrogenase) 50  $\mu$ L of the  
174 supernatant was mixed with 100  $\mu$ L of 2-Oxoglutarate and NADH (Merck millipore®) in a 5 mL  
175 tube. Both the mixtures were then allowed to stand for 60 seconds to complete the reaction.  
176 After the completion of reaction, absorbance was measured at 405 and 340 nm for LDH and  
177 GOT, respectively, using a spectrophotometer (Microlab 300, ELITech Group, France).

#### 178 2.6. *Evaluation of the best evolved extenders by assessing the in vivo fertility rate of buffalo* 179 *sperm*

180 Based on the *in vitro* sperm quality assays, the best evolved levels of pigeon egg yolk  
181 in extender were evaluated for in vivo fertility rate of cryopreserved semen. The semen was  
182 collected from two buffalo bulls of similar age and known fertility. The ejaculates were split,  
183 with one portion cryopreserved in the chicken egg yolk (20%, Control) containing extender  
184 and the other portion in the pigeon egg yolk (15%) containing extender. The inseminations  
185 were performed under field conditions over the three months of period during the peak breeding  
186 season. All the experimental inseminations were performed approximately 24 hours after onset  
187 of oestrus. Four hundred artificially bred animals (100 buffaloes/bull/extender) were examined  
188 for pregnancy through rectal palpation at least 90-days post-insemination.

#### 189 2.7. *Experimental design and data analysis*

190 Semen was collected from 6 Nili Ravi buffalo bulls (2 ejaculates/bull/ replicate) at weekly  
191 intervals for a period of three weeks (replicates). Semen from each bull was processed  
192 separately and each ejaculate was split into five aliquots and cryopreserved with TCA extender  
193 containing 5%, 10%, 15% and 20% PEY or 20% CEY (controls). Total numbers of ejaculates  
194 were 36 (three replicates per each of the six bulls and two ejaculates per replication). The data  
195 on semen quality parameters and enzyme leakages were analyzed using analysis of variance in  
196 randomized complete block design. When F-ratio was found significant ( $P < 0.05$ ), Least  
197 significant difference (LSD) test was used to compare the treatment means. For *in vivo* fertility

198 rates: semen ejaculates collected from two buffalo bulls was split into two aliquots and  
199 cryopreserved in the PEY (15%)  
200 containing extender or CEY (20 %) containing extender (control). Four hundred (100  
201 buffaloes/bull/extender) were examined for pregnancy through rectal palpation at least 90-days  
202 post-insemination. The data on conception rate were analyzed using Chi-square test.

203

### 204 **3. Results**

205 The data on effect of different levels of pigeon egg yolk in extender on post-thaw  
206 quality of buffalo bull spermatozoa are presented in figures 1-5. The percentages of progressive  
207 motility, plasma membrane integrity, live/dead ratio and sperm viability were higher ( $P < 0.05$ )  
208 in extenders having 10% and 15% PEY compared to 5% PEY, 20% PEY or 20% CEY  
209 (controls;). Sperm chromatin damage did not differ in extenders having PEY (5%, 10%, 15 %)  
210 and CEY (20%; controls). However, the least damage was observed with 20% PEY egg-yolk  
211 compared to CEY (controls;  $P < 0.05$ ).

212 The data on the effects of different concentrations of pigeon egg yolk in extender on  
213 the leakage of LDH and GOT are shown in figures 6-7. It is evident that the extender containing  
214 10% PEY and 15% PEY egg yolk had less LDH and GOT leakage compared to 5%PEY, 20%  
215 PEY and 20 % CEY (controls;  $P < 0.05$ ).

216 The extender having 15% PEY yielded higher ( $P < 0.05$ ) fertility rate in bull 1 (57% vs.  
217 41%), while the difference was not significant in bull 2 (55vs. 43%;  $P > 0.05$ ). The overall  
218 fertility rate was higher ( $P < 0.05$ ) with spermatozoa cryopreserved in extender containing 15%  
219 PEY (56% vs. 42.0%) compared to 20% CEY (controls) ( $P < 0.05$ ; Table 1).

220

### 221 **4. Discussion**

222 The pigeon egg yolk possesses higher lecithin content [19], lower cryoprotectant  
223 antagonists [18] and higher levels of useful trace elements. Therefore, present study was  
224 conducted to investigate the optimum levels of pigeon egg yolk for cryopreservation of buffalo  
225 bull semen.

226 The sperm membrane is considered to be primary site of cryodamages owing to its  
227 higher PUFA content that makes it more vulnerable to the intracellular ice crystals formed  
228 during the freezing process. The damage to the membrane results in the loss of lipids,  
229 phospholipids [32], cholesterol and ultimately the ratio of PUFAs: SFAs is disturbed. Since  
230 egg yolks from different avian species vary considerably in their fatty acid, cholesterol and  
231 phospholipid content, their use in the semen extender may result in to different levels of  
232 cryoprotection for the sperm. In the present study, pigeon egg yolk at 10% and 15% levels in  
233 tris citric acid extender improved the post thaw quality of buffalo sperm compared to 20%  
234 CEY. In a similar study, quail egg yolk that is high in saturated fatty acids and turkey egg yolk  
235 that have higher cholesterol content were reported to have better cryoprotective effect on  
236 buffalo sperm compared to chicken egg yolk [17]. It is relevant to mention that the major  
237 cryoprotectant viz; lecithin content being higher in the pigeon compared to the chicken egg  
238 yolk [19] might have played a role to ameliorate the cryodamages possibly through replacing  
239 the phospholipids in sperm plasma membrane and/or by forming a protective film around the  
240 sperm membrane as reported earlier [18]. On the basis of results obtained in the present study  
241 it is clear that 10-15% of pigeon egg yolk in the semen extender had enough lecithin to provide  
242 better protection to the sperm plasma membrane compared to 20% chicken egg yolk. Further,  
243 decreased levels of cryoprotectant antagonists like HDLs and yolk granules that are  
244 interestingly present in low concentration in pigeon egg yolk [18] might also have contributed  
245 to the better sperm protection observed in this study.

246 The avian egg yolks have different levels of micronutrients such as vitamins, minerals/  
247 trace elements [8], antibodies [33] and antioxidants [34]. Among trace elements, Selenium (Se)  
248 is reported to have significant impact on male reproduction in rats, mice, chickens, pigs, sheep,  
249 and cattle [35]. Selenium is an integral part of glutathione peroxidase (GSH-PX), an enzyme  
250 which has a major role in providing protection to cell internal structures and cell membrane  
251 lipids against free radicals [36,37]. In the present study, improvement observed in sperm  
252 motility, plasma membrane, livability, and viability and especially chromatin damage is  
253 considered might be associated with the higher levels of Se present in the pigeon egg yolk  
254 along with a reduction in oxidative stress induced during cryopreservation. Not only Se  
255 supplementation of the semen extender results in a dose-dependant increase in the total  
256 antioxidant capacity and post thaw quality of buffalo sperm [38] but also dietary Se  
257 supplementation in mice, sheep and cattle has been reported to improves the semen quality [35,  
258 39-41].

259 The cryopreservation induces extracellular enzymes leakages possibly by damaging/  
260 deteriorating the sperm membrane [17]. The extracellular enzyme leakage is considered to be  
261 a marker of fertilizing ability of ram [42], buck [43] bull [44] and buffalo bull sperm [17].  
262 Lactate dehydrogenase (LDH) is involved in metabolic processes which provide energy for  
263 survival, motility and fertility of sperm [17,43]. Its extracellular leakage has negatively been  
264 correlated with fertilizing ability of sperm and could be used as a marker of membrane damage  
265 during freezing. Similarly, the release of glutamic oxalacetic transaminase (GOT) is associated  
266 with cryodamages in buffalo sperm [45]. In the present study, extracellular leakages of LDH  
267 and GOT were recorded lowest in buffalo sperm cryopreserved in the extender with 10% and  
268 15 % pigeon egg yolk compared to 20 % chicken egg yolk (controls). The higher levels of  
269 lecithin present in the pigeon egg yolk might have formed a protective layer and/or replenished  
270 the phospholipids damaged/lost during cryopreservation resulting into comparatively less

271 enzyme leakage [19] and therefore, a better protection of the sperm. Moreover, higher levels  
272 of Se in pigeon egg yolk might have enhanced the antioxidant potential that resulted in  
273 reduction of lipid peroxidation of sperm phospholipids [8,38]. It is pertinent to mention that  
274 the data on motility, plasma membrane integrity and viability are very much supported with  
275 that of enzyme leakages, i.e., a similar pattern of effects of egg yolk levels are observed on  
276 motility, viability and plasma membrane integrity of cryopreserved buffalo semen.

277         The higher fertility rates were recorded with extender having 15% pigeon egg yolk  
278 compared to 20% chicken egg yolk. The real test to evaluate a semen sample is to check the  
279 fertility rate after its *in vivo* insemination in a routine artificial insemination programme under  
280 field conditions [46-48]. The higher fertility rate observed with 15% pigeon egg yolk observed  
281 in the present study are in line with the higher sperm quality viz; motility, plasma membrane  
282 integrity, livability, viability, chromatin integrity and extracellular enzyme leakage suggesting  
283 that all the sperm quality parameters used to assess the semen quality do have essential role in  
284 the fertilization process [49].

285         On the basis of existing information on the PEY, better sperm quality and higher  
286 fertility rates observed post-insemination in this study could be linked to the higher lecithin  
287 contents and higher Se and Mo concentrations present in the PEY. However, for proper  
288 evaluation, a precise bio-chemical composition of the pigeon egg yolk is required along with  
289 its mechanism for sperm membrane protection. In conclusion, the results of this study have  
290 shown that 15% pigeon egg yolk in the semen extender significantly improves the post-thaw  
291 semen quality and *in vivo* fertility in the buffalo.

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295

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