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Oxidative stress in donor mares for ovum pick-up delays embryonic development

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1 **Abstract**

2 The *in vitro* production of equine embryos via ovum pick-up (OPU) and intracytoplasmic
3 sperm injection (ICSI) has increased rapidly. There is a marked effect of the individual mare on the
4 outcome of OPU-ICSI, but little is known about the influence of the mare's health condition. This
5 study aimed to investigate the potential associations between the concentrations of interleukin-6 (IL-
6 6), reactive oxygen metabolites (d-ROMs), and biological antioxidant potential (BAP) in serum of
7 oocytes' donor mares and the subsequent embryonic development. Just before OPU, a blood sample
8 was collected from 28 Warmblood donor mares, that were subjected to a routine OPU-ICSI program.
9 The serum concentrations of IL-6, d-ROMs, and BAP were assayed photometrically. The maturation,
10 cleavage and blastocyst rate as well as the kinetics of blastocyst development were recorded. The
11 average blastocyst rate was 24.68 ± 5.16 % and the average concentrations of IL-6, d-ROMs, and BAP
12 were 519.59 ± 157.08 pg/mL, 171.30 ± 4.55 carratelli units (UCARR), and 2711.30 ± 4.55 μ mol/L,
13 respectively. Serum concentrations of IL-6, d-ROMs, and BAP were not significantly different
14 between mares yielding at least one blastocyst (552.68 ± 235.18 pg/mL, 168.36 ± 5.56 UCARR, and
15 2524.80 ± 159.55 μ mol/L) and mares yielding no blastocysts (468.47 ± 179.99 pg/mL, 175.85 ± 7.89
16 UCARR, and 2999.50 ± 300.13 μ mol/L, respectively). Serum concentrations of d-ROMs were
17 significantly lower in mares with fast growing (at day 7-8 post ICSI; 148.10 ± 8.13 UCARR) compared
18 to those with slow growing blastocysts (\geq day 9 post ICSI; 179.41 ± 4.89 UCARR; $P= 0.003$). Taken
19 together, the serum concentration of IL-6, d-ROMs, and BAP do not determine the mare's ability to
20 produce blastocysts *in vitro*. Although it may be questioned whether a single sample is representative
21 of the mare's health status, changes in serum metabolites related to oxidative stress at the time of
22 oocyte retrieval were linked to a delayed blastocyst development in a clinical OPU-ICSI outcome.

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26 **Keywords:** IL-6; Oxidative stress; d-ROMs; BAP; OPU-ICSI; Mares

27 **1. Introduction**

28 Ovum pick-up (OPU) and intracytoplasmic sperm injection (ICSI) are substantially used to
29 produce equine embryos *in vitro* [1-3]. The OPU-ICSI program is multi-advantageous and is as
30 effective as embryo flushing when measured by the number of day 45 pregnant recipients per mare
31 [4]. Regardless the stage of the ovarian cycle, follicular health and season, OPU-ICSI allows the
32 production of a high number of foals, even from old, subfertile [5], and euthanized mares [6]. The
33 success rate of OPU-ICSI is mainly evaluated by the mare's ability to produce a blastocyst and by the
34 rate of (transferable) blastocysts [2], which are repeatable for an individual mare between two
35 consecutive sessions [7].

36 There are several known mare related factors that can affect the success rate of OPU-ICSI program.
37 Aged mares (> 20 y) have a relatively low number of ovarian follicles [8]. As such, the number of
38 embryos per OPU session declines in old mares, but the mare's age does not have a significant effect
39 on the developmental competence of the oocytes [7, 9]. A second factor which markedly affects the
40 success rate of OPU-ICSI is the mare's breed [2]. The oocytes of Arabian donor mares show
41 significantly lower cleavage and blastocyst rates compared to those of Warmblood mares [10]. Still,
42 maternal age and breed are constant factors, and they cannot explain short-term fluctuations in the
43 success rates of OPU-ICSI for an individual mare. The relationship between maternal health, oocyte
44 quality, and OPU-ICSI outcome has been scarcely investigated in mares. On the one hand, it has been
45 shown that the physiological status (transitional vs. cycling; [7, 11]) and the presence of reproductive
46 disorders [11] are not significantly affecting the blastocyst rate. On the other hand, mares engaged in
47 intense sporting activities [11] and obese mares [12, 13] display a decreased oocyte developmental
48 competence *in vitro*. Nevertheless, more research is needed to study the impact of the mare's health
49 condition on the success rate of OPU-ICSI.

50 Female gametes are vulnerable to oxidative stress [14]. Estimation of the systemic oxidative stress
51 index (OSI), measured by derivatives of reactive oxygen metabolites (d-ROMs) and biological

52 antioxidant potential (BAP) has been well established in women undergoing *in vitro* fertilization [15,
53 16]. There is a direct association between serum and follicular fluid values of d-ROMs and OSI in
54 women [17], where higher values in serum were accompanied with abnormal fertilization, while
55 increased values in follicular fluid were associated with diminished embryo quality [18]. In horses, we
56 recently showed that the values of OSI in serum and follicular fluid are correlated too [19].

57 Pro-inflammatory cytokines play a vital role in maintaining the ovarian physiology during
58 folliculogenesis, oocyte maturation and ovulation [20]. There is a strong association between serum
59 and follicular fluid concentrations of IL-6 both in women [21] and mares [19]. Excess IL-6 has been
60 associated with decreased estradiol synthesis and aromatase activity in granulosa cells of women *in*
61 *vitro* [22]. Higher IL-6 concentrations inhibited the expression of luteinizing hormone receptor mRNA
62 during the maturation and differentiation of cultured rat granulosa cells [23]. Higher follicular fluid
63 IL-6 values in women were associated with decreased clinical pregnancy rate [24]. In mares, higher
64 concentrations of IL-6 within the preovulatory follicle have been correlated with diminished oocyte
65 quality [25].

66 The relationship between maternal inflammation or oxidative stress, and the OPU-ICSI outcome
67 has not been previously investigated in mares. We hypothesize that there is an association between the
68 serum concentrations of oxidative stress markers (d-ROMs, BAP, and OSI) and the pro-inflammatory
69 cytokine IL-6 at the time of oocytes retrieval (OPU) and the oocyte developmental competence in
70 mares. Therefore, the objective of the present study was to investigate the associations between the
71 serum concentrations of d-ROMs, BAP, OSI, and IL-6 and the OPU-ICSI outcome in mares.

72 **2. Materials and methods**

73 For this study, no specific samples were acquired from or extra procedures were performed with
74 the mares included in this study as analyses were performed during routine clinical OPU-ICSI services.
75 For this reason, no extra ethical clearance was necessary for the present study.

76 **2.1. Animals**

77 Twenty-eight Warmblood mares, with a body condition score ranged between 3 to 6 [26] and aged
78 2-23 years old were used between mid-January and mid-March 2022. These mares regularly
79 participated in the OPU-ICSI program at the equine reproduction clinic, Faculty of Veterinary
80 Medicine, Ghent University.

81 **2.2. OPU procedures**

82 Just before conducting the OPU and after blood sampling, a preoperative regime [9] of
83 benzylpenicillin (20000 IU/ kg intramuscular; Penikel[®], Kela, Sint Niklaas, Belgium) and flunixin
84 meglumin (1.1 mg/ kg intravenous; Wellicox[®], Ceva Santé Animale, Naaldwijk, The Netherlands) was
85 used. During the OPU, detomidine hydrochloride (0.01 mg/ kg intravenous; Domidine[®], Eurovet
86 Animal Health BV, Bladel, The Netherlands) and butorphanol tartrate (0.01 mg/ kg intravenous;
87 Dolorex[®], MSD Animal Health, Sint-Lambrechts-Woluwe, Belgium) were used for sedation. To
88 subside intestinal contractions, N-butylscopolammonium bromide (0.3 mg/ kg intravenous;
89 Buscopan[®], Boehringer Ingelheim, Brussel, Belgium) was injected. Urinary bladder catheterization
90 and epidural anesthesia were not applied. After proper aseptic preparation for the perineal region, the
91 transvaginal transducer (7.5 MhZ linear probe, MyLabOne, Esaote, Genoa, Italy; [9]) equipped by a
92 12-G double-lumen needle attached via a double way tube system to a prewarmed collection bottle of
93 flushing medium (Equiplus[®], Mintube, Tiefenbach, Germany). All visible antral follicles were
94 punctured, aspirated, scraped, and flushed 8 times.

95 **2.3. In vitro embryo production**

96 The collection of oocytes [9] was carried out under sterile conditions using a laminar air flow
97 equipped with a stereomicroscope (Olympus SZX7[®], Olympus Corp., Japan). The whole contents of
98 the collection bottle (follicular fluid, flushing medium, and scrapped follicular cells) were filtrated
99 through a sterile 70 µm filter (Cell strainer[®], BD Biosciences, Falcon, Erembodegem, Belgium) and
100 the COCs were recovered from the filtrated contents in medium 199 with Hank's salts (Gibco, Life

101 Technologies, Merelbeke, Belgium) supplemented with 10% fetal bovine serum (FBS; Gibco).
102 According to the schedule, the recovered COCs were either directly transferred to maturation medium
103 (medium 199 with Earl's salts (Gibco) containing 10% (v/v) FBS (Gibco), 9.4 µg/mL follicle
104 stimulating hormone, and 1.88 g/ml luteinising hormone (Stimufol, Reprobiol, Ouffet, Belgium)) or
105 were kept overnight in a commercial embryo holding medium (Emcare[®], Agtech, Zulte, Belgium) at
106 room temperature (~22 °C) prior to maturation. *In vitro* maturation was carried out in groups of 2-18
107 COCs in 100-500 µl maturation medium under oil (CooperSurgical, Venlo, The Netherlands) at 38.2
108 °C in 5% CO₂ containing air for 28-32 h. A small piece of straw with frozen semen was thawed in 1
109 mL G-MOPS (38.2 °C ; Vitrolife, Londerzeel, Belgium) and centrifuged twice (400 × g/ 3 min at
110 room] temperature; ~22 °C). After the first centrifugation, the supernatant was discarded and the pellet
111 was re-suspended in 1 mL G-MOPS. After the second centrifugation, the supernatant was discarded
112 and the pellet was resuspended in 200 µl G-MOPS. Immediately before ICSI, a small amount of the
113 resuspended sperm was added to a 5 µl droplet of 7% polyvinylpyrrolidone (CooperSurgical, Venlo,
114 The Netherlands). Intracytoplasmic sperm injection, *in vitro* culture of presumptive zygotes, and the
115 evaluation of embryonic development were performed until day 13 post ICSI [9].

116 **2.4. Blood collection and laboratory analyses**

117 A single blood sample per mare was collected from the jugular vein into vacutainer tubes with clot
118 activator (BD Vacutainer[®], BD-Plymouth, UK). To separate the serum, samples were centrifuged at
119 2460 × g for 20 min at 4 °C. Serum was aliquoted into sterile 2 ml Eppendorf tubes and stored at -80
120 °C until further biochemical analysis.

121 Colorimetric kits (Diacron[®]; Diacron International, Italy) were used to measure the serum
122 concentrations of d-ROMs and BAP, according to the manufacturer's guidelines [19]. A Multiskan
123 GO spectrophotometer (Thermo Fisher Scientific, Finland; at 37° C) was used to estimate the
124 photometric measurements for both kits at 505 nm. The coefficient of variation was 1.72 % for d-
125 ROMs and 2.32 % for BAP. The lowest limit of detection for d-ROMs and BAP was 11 UCARR and

126 150 $\mu\text{mol/L}$, respectively. The OSI was determined from the concentrations of d-ROMs and BAP
127 using the formula $\text{d-ROMs/BAP} \times 100$ [27].

128 Serum concentrations of IL-6 were measured spectrophotometrically using an equine IL-6 ELISA
129 kit (Nori®, Genorise Scientific, USA), according to the manufacturer's guidelines. A Multiskan GO
130 spectrophotometer (Thermo Fisher Scientific, Finland; room temperature) was used to determine the
131 optical density at 450 and 540nm, which was followed by a wavelength correction. The average
132 coefficient of variation was 6.49 % and the lowest detection limit was 16 pg/mL.

133 **2.5. Study design**

134 Blood sampling was performed just before OPU. Immediately after OPU, the cumulus-oocyte
135 complexes (COCs) were recovered, matured, and fertilized by ICSI.

136 At each OPU-ICSI session, (a) the number of aspirated follicles, (b) the number of recovered
137 oocytes, (c) the recovery rate ($\text{b/a} \times 100$), (d) the number of mature oocytes, (e) the maturation rate (d/b
138 $\times 100$), (f) the number of cleaved presumptive zygotes, (g) the cleavage rate ($\text{f/d} \times 100$), (h) the number
139 of produced blastocysts, (i) the blastocyst rate ($\text{h/d} \times 100$), (j) the proportion of the cleaved zygotes that
140 developed to blastocysts ($\text{h/f} \times 100$), (k) the time of blastocyst formation, and (l) the serum
141 concentrations of d-ROMs, BAP, OSI, and IL-6 were recorded. The mares were divided into different
142 groups according to (1) their ability to produce blastocysts: blastocyst producing (≥ 1 blastocyst; $n=17$)
143 and non-producing (0 blastocyst; $n=11$) mares, (2) the required time for embryonic development:
144 mares with fast growing (first blastocyst developed at day 7-8 post ICSI; $n=6$) and mares with slow
145 growing (first blastocyst developed at day ≥ 9 post ICSI; $n=11$) embryos, and (3) age: young (≤ 14 y),
146 middle-aged (15-19 y), and old (≥ 20 y) mares [28].

147 **2.6. Statistical analyses**

148 The assessment of the normality of data was performed using a Shapiro-Wilk test. The mean values
149 of d-ROMs, BAP, and OSI, but not IL-6, were normally distributed. For the blastocyst producing

150 mares, Spearman's correlation coefficients between the blastocyst rate, the proportion of cleaved
151 zygotes that developed to blastocysts, and the serum concentrations of d-ROMs, BAP, OSI, and IL-6
152 were calculated. Differences in serum parameters between groups based on the mare's ability to
153 produce embryos (blastocyst producing vs. non-producing mares) and the onset of embryonic
154 development (fast vs. slow growing blastocysts) were determined using the independent t-test or
155 Mann-Whitney U-test. Differences between groups based on maternal age were explored using Welch
156 one-way ANOVA followed by Games-Howell or Kruskal-Wallis test. The data were analyzed using
157 the statistical package for social science SPSS® (SPSS Inc., version 16.0, Chicago, IL. USA), and a *P*-
158 value <0.05 was considered significant. Data are presented as mean ±SEM.

159 **3. Results**

160 The mean ±SEM values of all the studied parameters are presented in Table 1. There were no
161 significant correlations between the blastocyst rate, the proportion of cleaved zygotes that produced
162 blastocysts, and the serum concentrations of d-ROMs, BAP, OSI, and IL-6.

163 As shown in Table 2, serum concentrations of BAP were significantly higher in old mares
164 (3544.60±218.07 µmol/L) compared to the young ones (2461.40±133.56 µmol/L). Values of OSI were
165 significantly increased in young mares (7.41±0.52) compared to old ones (4.95±0.24). Serum
166 concentrations of d-ROMs and IL-6 were not significantly different between young, middle-aged, and
167 old mares.

168 Serum concentrations of d-ROMs, BAP, OSI, and IL-6 were not significantly different between
169 the blastocyst producing and non-producing mares (Table 3). Serum concentrations of d-ROMs (Table
170 4) were significantly (*P*= 0.003) higher in mares with slow growing blastocysts (179.41±4.89 UCARR)
171 compared to those with fast growing ones (148.10±8.13 UCARR).

172

173

174 **4. Discussion**

175 In this study, we found an association between the serum concentrations of oxidative stress markers
176 (d-ROMs, BAP, and OSI) and the pro-inflammatory cytokine IL-6 at the time of OPU and kinetics of
177 embryo development. More specifically, high concentrations of d-ROMs at the time of OPU are linked
178 to delayed embryonic development in mares. This may point out that a disturbance in maternal health
179 related to oxidative stress can affect the OPU-ICSI outcome in mares.

180 Overall, the OPU-ICSI results in this study were consistent with those reported previously [2, 9].
181 Serum concentrations of d-ROMs, BAP, and IL-6 were within the previously reported range in
182 Warmblood mares during the non-breeding season [19]. In agreement with literature, ageing had no
183 effect on the serum concentrations of d-ROMs in horses [29]. In our study, the serum concentrations
184 of BAP were significantly higher and the values of OSI were significantly lower in old mares compared
185 to young ones. In humans, the total antioxidant capacity in serum also increases with advancing age,
186 which may be related to diet and daily routine [30]. The effect of age on the serum antioxidant status
187 in horses is not clear. Some studies did not report any significant effect of ageing on the serum total
188 antioxidant status [31] and BAP [29] in horses. On the other hand, Andriichuk et al. [32] found that
189 physical exercise increases the plasma concentrations of thiobarbituric acid reactive substrates,
190 catalase, and glutathione reductase in Warmblood horses. While our study indicates an effect of ageing
191 on oxidative stress, further influence of diet and physical activity remains to be determined.

192 Although oxidative stress markers in serum did not determine the mare's ability to produce
193 embryos, mares with slow growing blastocysts showed significantly higher serum concentrations of
194 d-ROMs. In a previous study, we found that serum and follicular fluid values of OSI (d-ROMs/BAP
195 $\times 100$) are directly correlated in Warmblood mares [19]. Speed of embryo development affects both
196 pregnancy and foaling rates in mares. Ducheyne et al. [33] found that fast growing embryos (formed
197 before day 9 post ICSI) yield significantly higher pregnancy rates compared to the slow growing ones

198 (formed after day 9 post ICSI). Foaling rate was significantly higher for day 7 and day 8 embryos
199 (71.7% and 53.3%) compared to day 9 and day 10 embryos (38.5% and 25%; [34]). In mammals, the
200 oocyte developmental competence is linked to maternal health [35]. Maternal oxidative stress is
201 increasing the concentrations of reactive oxygen species (ROS) in oocytes, which reduces their
202 viability [36, 37]. Oocytes with higher levels of ROS show delayed two-cell, four-cell, and blastocyst
203 development in mice [36, 38]. Oxidative stress in serum and follicular fluid significantly decreases the
204 clinical pregnancy rate in women [15, 39]. Higher values of d-ROMs in follicular fluid of women have
205 been associated with abnormal fertilization and production of bad quality embryos [18]. Here, we
206 hypothesize that the higher serum concentrations of d-ROMs may be reflected by increased d-ROMs
207 in the follicular fluid, which may affect oocytes quality, resulting in delayed embryo development.
208 More studies are necessary to further explore the effect of oxidative stress and antioxidants on the
209 oocyte developmental competence in horses.

210 In this study, serum concentrations of IL-6 neither affected the mare's ability to produce embryos
211 nor the blastocyst rate. There is a positive association between serum and follicular fluid concentrations
212 of IL-6 in mares [19, 25]. In the follicle, IL-6 is responsible for extracellular matrix formation and
213 stabilization, which regulates cumulus cells expansion and increases the oocyte competence [40].
214 Several studies have been conducted, but there is no conclusive answer regarding the role of IL-6 in
215 oocytes and subsequent embryos. Higher concentrations of IL-6 in FF can either increase [41] or
216 decrease [24] the pregnancy outcome in women. Supplementation of culture media with IL-6 improved
217 fetal development of IVF-embryos in cows [42] and supported embryonic compaction, blastulation
218 and hatching in mice [43]. The expression of IL-6 and IL-6 signal transducer genes in granulosa cells
219 was upregulated with advancing maternal age in mares [28]. In women with infertility, there was a
220 downregulation in the expression of IL-6 signal transducer and IL-6 receptor genes in granulosa cells
221 of older patients [44]. However, maternal age did not affect the concentrations of IL-6 in mares' serum

222 (this study) and women's FF [45]. Therefore, it seems that the role of IL-6 in oocytes and embryos is
223 species-specific and dose-dependent.

224 **5. Conclusions**

225 In conclusion, mares with higher serum concentrations of d-ROMs at the time of oocyte retrieval
226 (OPU) show a delayed embryonic development. More studies should be conducted to investigate
227 underlying mechanisms and potential therapy by antioxidants supplementation during the *in vitro*
228 maturation of the oocytes collected from mares with oxidative stress. The measured ranges of d-ROMs,
229 BAP, and IL-6 concentrations in serum at the time of OPU cannot be used to predict the mare's ability
230 to produce embryos *in vitro*.

231 **CRedit authorship contribution statement**

232 Conceptualization: MH, KS, JG; Animals work: JG, MP, SP, IG, EVB; Lab work: MH, DAV,
233 TDC, KS; Data curation: MH, DAV; Original draft writing: MH; Editing, Review, and Supervision:
234 JL, KS, AVS; Providing fund: JL, JG, KS.

235 **Conflict of interest**

236 The authors declare no competing interests.

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