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The full details of the published version of the article are as follows:

TITLE: Effect of melatonin supplementation in the long-term preservation of the sheep ovaries at different temperatures and subsequent in vitro embryo production

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JOURNAL: THERIOGENOLOGY

PUBLISHER: Elsevier

PUBLICATION DATE: January 2018

DOI: 10.1016/j.theriogenology.2017.10.009



# Accepted Manuscript

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PII: S0093-691X(17)30483-1

DOI: 10.1016/j.theriogenology.2017.10.009

Reference: THE 14292

To appear in: Theriogenology

Received Date: 14 April 2017

Revised Date: 1 October 2017

Accepted Date: 7 October 2017

Please cite this article as: Goodarzi A, Zare Shahneh A, Kohram H, Sadeghi M, Moazeni zadeh MH, Fouladi-Nashta A, Dadashpour Davachi N, Effect of melatonin supplementation in the long-term preservation of the sheep ovaries at different temperatures and subsequent in vitro embryo production, *Theriogenology* (2017), doi: 10.1016/j.theriogenology.2017.10.009.

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# Effect of melatonin supplementation in the long-term preservation of the sheep ovaries at different temperatures and subsequent in vitro embryo production.

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- 11 Abstract

Investigations in the past decades have shown that oocytes developmental competence following in vitro 12 13 fertilization is greatly influenced by an interval between isolation of the ovaries immediately after 14 death/slaughter and oocytes recovery from the visible follicles. In order to determine the optimal 15 conditions for long-term preservation of ovaries, an experiment was conducted with adding different 16 doses of melatonin (0 (C), 500 (M1), 600 (M2), 700 (M3) and 800 (M4) µM) as an antioxidant to sheep 17 ovaries preservation medium (PBS) maintained at 4 and 20°C for 24 hours. The effects on in vitro embryo production (IVEP) parameters including maturation, fertilization, cleavage, and blastocyst rates 18 19 and the total number of blastomere were evaluated after the ovaries preservation. Melatonin reduced the 20 decline in fertilization rate as an indicator of success in vitro maturation ( $P \le 0.05$ ). Furthermore, ovarian 21 storage time had significant negative effect ( $P \le 0.05$ ) on IVEP parameters. Supplementation with 22 melatonin increased the total cell number of blastocysts as an indicator of embryo quality (i.e. mean 23 blastomeric cells in 4°C groups: 86.00±3.00, 98.50±3.5, 111.5±1.5, 125.5±2.00 and 126.50±5.5 for C, M1, M2, M3 and M4. respectively). Overall, the results showed that the use of melatonin antioxidant in 24

- the ovaries storage medium had beneficial effects on sheep oocytes development and embryos quality.
- 26 Key words: melatonin, ovary, preservation medium, sheep
- 27 1. Introduction

Sheep is an important species in animal husbandry industry, and a critical research model to development 28 29 of assisted reproductive technologies (ARTs) in human and endangered species[1, 2].Cumulus-oocyte-30 complexes (COCs) from both small and large antral follicles can be isolated directly from ovarian tissue and matured in vitro to obtain mature gametes. However, collection of oocytes for ART from live animals 31 32 is costly. The use of ovaries from slaughtered animals has provided an affordable source of oocytes for researchers and laboratories involved in animal ART [2, 3]. However, integrating ovarian transport into in 33 34 vitro embryo production (IVEP) protocols has been an important challenge in large countries and/or in 35 low resource settings where the slaughterhouse is located far from the laboratory [4]. The long distance transportation of ovaries to the laboratory has adverse effects on oocyte quality in terms of nuclear 36 maturation and developmental competence after the in vitro maturation (IVM) and fertilization (IVF) [5]. 37 38 Ovaries need to be collected and returned to the laboratory instantly after the slaughter in order to make 39 effective use of the oocytes contained within [6]. The type of transportation medium [3, 6-8], storage time [3-16], as well as, its temperature during transportation of the ovaries [4, 5, 10-16] are among the factors 40 affecting subsequent follicular and oocyte survival, and oocyte developmental competence [7]. 41 Consequently, many studies have been conducted in order to improve oocyte preservation by modifying 42 43 the transportation solutions. These studies have demonstrated that the transportation of ovaries is possible 44 without a considerable harm to oocyte and follicles [4, 7, 10, 17]. However, there seems to be differences 45 between animal species in sensitivity of the oocytes to the transportation conditions. For example in 46 bovine, when the ovaries were stored for 7 hours at 4°C, 20°C, and 39°C, none of the oocytes from the 47 ovaries stored at 4°C and 39°C developed to blastocyst stage compared to the other group [13]. Whereas in mice, storage of ovaries at 4°C for up to 24 hours did not affect the number of mature gametes which 48 could be collected or fertilized post-orthotropic transplantation [10]. 49

50 Normal cellular metabolism produces reactive oxygen species (ROS) and reactive nitrogen species 51 (RNS), which regulate diverse cell functions. Reactive species, nevertheless are highly reactive with lipid, protein, and nucleic acid resulting in a loss of membrane integrity, structural, or functional changes in 52 53 proteins, and damage in nucleic acids referred to as oxidative stress [18]. During the ovary transportation 54 to laboratory, the stoppage of blood flow reduces oxygen and energy supply, and put ovaries under ischemic conditions [19]. Ischemia damages follicles viability and luteal function in ovaries; oxygen free 55 56 radicals in particular, are major contributors to organ damage during preservation [20]. Furthermore, the 57 antioxidant system of ovary cells is compromised during the preservation. To prevent the damage due to the reactive species, the cells possess a number of antioxidant enzymes, such as superoxide dismutase 58 59 (SOD), glutathione peroxidase (GPX) and catalase (CAT) [7]. The balance between ROS and antioxidants within the follicle seems to be critical to the function of oocyte and granulosa cells [4, 7, 21]. 60 61 During the ovary transportation, ROS and RNS generation in the follicular microenvironment by oocyte and somatic cells accompanied by reduced levels of antioxidant enzymes may cause oxidative stress-62 63 mediated apoptosis in follicles [22].

64 The main goal in ovaries preservation is to maintain the function of the oocyte enclosed in follicle for in vitro maturation and fertilization. Accordingly, many studies investigated the effects of different 65 66 antioxidants to reduce the damage caused by free oxygen radicals during ovaries preservation [6-8, 15]. 67 Several studies indicated that the imbalance between ROS production and antioxidant activities could 68 cause oxidative stress [7, 23]. In cow, oocyte meiotic competence could be maintained when ovaries 69 were stored in saline supplemented with epigallocatechingallate compared to glutathione [6]. 70 Experimental data from the feline model showed that oocytes from ovaries preserved in PBS 71 supplemented with SOD reduced the percentage of apoptosis in COCs compared to control group [7].

The protective effects of melatonin as a powerful direct scavenger of free radicals are well documented in 72 73 the recent years [21, 23-29]. Melatonin, because of its amphiphilic nature, diffuses broadly in diverse 74 subcellular compartments barriers. It is also a highly effective antioxidant and anti-apoptotic agent, which 75 due to its direct scavenging of toxic oxygen derivatives and its ability to reduce of ROS and RNS, 76 prevents oxidative and nitrosative damages to all macromolecules in all compartments of a 77 cell[30]. Therefore, we hypothesized that the use of this antioxidant in preservation medium may also have 78 beneficial effects by reducing oxidative stress in ovaries during long term preservation. In the present 79 study, we used the sheep as an experimental model to examine the effects of supplementing the ovaries 80 transport medium with melatonin at two different temperatures on blastocyst rate and quality after IVEP.

#### 81 2. Materials and methods

- 82 Except otherwise indicated, chemicals were purchased from Sigma Chemical Co. (Saint Louis, MO,
- USA). Adult testes and ovaries (without distinguished corpus luteum) were obtained from slaughtered
  Lory-Bakhtiary sheep aged 2 to 3 years without apparent abnormality in reproductive tract. Due to the
- 85 low solubility of melatonin in water, it was dissolved in ethanol before adding to PBS.
- 86 2.1 Experimental design
- 87 There were two experimental groups in the present study design. Two different temperatures (4
- and 20 °C) and 4 concentrations of melatonin (M1: 500, M2: 600, M3: 700 and M4: 800  $\mu$ M) were
- 89 considered as the treatments. In experiment 1 we analyzed the effect of maintaining ovaries in PBS
- supplemented with four concentrations of melatonin (M1- M4) at 4°C for a period of 24 h prior to oocyte
- 91 retrieval. As the control group(C) ovaries were maintained in PBS without melatonin supplementation for
- a period of 24 hours before oocyte collection. Fertilization was carried out using the freshly collected
- 93 sperm from ram testicles of the same breed at the time of ovaries collection.
- In the experiment 2 we assessed the effect of maintaining ovaries at 20°C for a period of 24h in PBS
  supplemented with (M1-M4) or without (C) melatonin.
- To evaluate the accuracy of the laboratory procedure during this study we also had considered IVF cycleswith standard condition (SC) concurrent with experiment 1, 2.
- 98 2.2 Ovaries collection

99 Ovine ovaries were collected immediately after slaughter and transferred into PBS containing 100  $\mu$ g/mL 100 penicillin–streptomycin (Gibco; Grand Island, NY, USA) as a primary repository. Afterwards, the 101 collected ovaries were washed three times in sterile PBS and then randomly divided between the 102 treatment conditions in thermos flask containing PBS (C) and PBS ± melatonin (M1-M4). The insulation 103 efficiency of the thermos flask in maintaining a constant temperature was tested prior to conducting the 104 experiments. After arrival in laboratory the SC group was instantly conformed for embryo production and 105 the other groups were preserved for 24 h in the experimental conditions.

- 106 2.3 In vitro maturation of oocyte
- Oocyte collection and in vitro maturation method were carried out as previously described [31], with 107 some modifications. In brief, ovarian antral follicles 2-6 mm in diameter were aspirated (45 ovaries per 108 treatment group) using an aspiration pump (MEDAP Sekretsauger P7040; Tilburg, The Netherlands) 109 fitted with a disposable vacuum line (length-35 cm, the internal diameter of 3 mm). The flow rate was set 110 at 10 mL H<sub>2</sub>O/min using an attached disposable 20-gauge needle. Next, oocytes with more than three 111 layers of cumulus cells and uniform ooplasm were selected for IVM [31-33]. The TCM199 medium used 112 for IVM was supplemented with 10% fetal bovine serum (FBS) (Cat. Number: A6003), 0.2 mM sodium 113 pyruvate, 5  $\mu$ g/mL of gentamicin, 10  $\mu$ g/mL of ovine follicle-stimulating hormone (oFSH), and 1  $\mu$ g/mL 114 of estradiol. COCs were cultured for 24 hours in a 50 µl droplet of maturation medium (approximately 10 115 116 oocytes per drop) under mineral oil at 38.5 °C in an atmosphere of 5% CO2 with maximum humidity.
- 117 After 24 hours oocyte with expanded cumulus considered as mature and selected for next stage.

118 2.4 Sperm preparation, in vitro fertilization (IVF) and in vitro culture (IVC), and staining 119 methods

In vitro fertilization was carried out as previously described [34] with some modifications. Briefly, fresh 120 spermatozoa were obtained from the slaughtered Lory-Bakhtiary rams (2-3 years old) in fertilization day. 121 After transport of the testicles to the laboratory in a cool box (5°C), all the blood and connective tissues 122 123 were removed aseptically in a cold room with the temperature of 5°C. For sperm recovery, 1 mL 124 tuberculin syringe attached with 22-gauge needle was inserted into the vas deference. The content of the vas deference was aspirated gently and the recovered spermatozoa was diluted 1:100 in sperm-TALP and 125 storage in 4°C less dan one houre. Samples with more than 60% progressive motility, which had normal 126 appearance, were selected for IVF. Motile spermatozoa were separated by percoll gradient (45% over 127 90%). The fertilization medium included 12 mM KCL, 25 mM NaHCO, 90 mM NaCl, 0.5 mM 128 NaH<sub>2</sub>PO<sub>4</sub>, 0.5 mM MgSO<sub>4</sub>, 10 mM sodium lactate, 3 mg/mL BSA (fatty acid free), 50 µg/mL gentamicin. 129 At least 15 min prior to insemination the sperm suspension was transferred into a droplet of fertilization 130 medium immediately prior to co-culture adjusted to a final concentration of  $1 \times 10^6$  cells/ml. Co-131 incubation of gametes (day 0; IVF) was carried out under 5% CO2/5% O2 in air for 18 hours at 38.5 °C. 132 133 After co-incubation, cumulus cells and attached spermatozoa to zona pellucida were removed by 134 repeatedly pipetting. The presumptive fertilized eggs were cultured at 38.5°C under 5% CO, 5% O<sub>2</sub>, 90% N<sub>2</sub> for the next 7 days in synthetic oviduct fluid (SOF), used as in vitro culture (IVC) medium[31]. Two 135 days after IVC the cleavage rate of the embryo were recorded. At day 8 after insemination, the rate of 136 embryo development to the blastocyst stage was recorded. 137

Examination of the IVF results carried out according to the previously described procedures [31]. Briefly, 138 24 hours after insemination one fourth of zygote were removed randomly from the culture medium and 139 washed twice in PBS-PVP (Polyvinylpyrrolidone, 1mg/mL). Then the zygote were fixed on a 100 µL 140 141 drop of paraformaldehyde solution [4% (w/v) in PBS, pH 7.4], at room temperature. In the final step, the 142 zygote were removed from the fixation drop and washed three times in PBS-PVP. Afterwards, zygotes 143 were transferred to 100µl drop of PBS-PVP containing 1mg/mL Hoechst 33342 for assessment of fertilization rate. The slides were examined under a fluorescent microscope with a UV filter showing the 144 sperm head and pronuclei having a blue appearance. The same method was used for counting the total 145 146 cells in blastocysts day 8 after insemination.

147 2.5 Statistical analysis

148 The GLM procedure of the SAS (SAS, Inc., Cary, NC, USA) was used for the analysis of 149 variance (ANOVA). The Duncan test was taken into account for comparisons of mean values with a 150 significant main effect. P values <0.05 were considered statistically significant. Data are presented as 151 mean  $\pm$  SEM, total number of blastomeric are presented as number.

152 3. Results

As expected, the standard condition group which was used as a control to assure the accuracy of IVEP procedure, resulted in higher maturation, fertilization, blastocyst rates and mean blastomeric cells  $(93.12\pm0.47, 84.62\pm0.75, 28.27\pm1.73 \text{ and } 171.5\pm4.50, \text{ respectively})$ . The 156 oocytes in this group were retrieved from ovaries which were transported to the laboratory in less 157 than three hours in PBS at  $37.5\Box$ ,

158 3.1 Effects of storage condition at 4°C

159 The effect of melatonin supplementation in ovaries preservation for 24 h at 4°C is presented in Table 1. 160 The results show that oocyte maturation rate was affected by treatment (P < 0.05). Only M3 (700) and M4 (800) increased maturation rates compared to the control, and the rates were similar to M1 and M2 161 groups. Higher concentration of melatonin (M3 and M4) improved fertilization and cleavage rates 162 compared to M1 and control group (p <0.05). Moreover, the highest melatonin supplementation (M4: 163 164 800) increased morula rates compared to all groups. Interestingly, some of the cleaved embryos in control group developed to blastocyst stage. The percentages of zygotes undergoing blastocyst in oocytes 165 obtained from ovaries of theM3 and M4 groups were similar (16.95±0.59 vs. 18.19±0.33, respectively) 166 but in all melatonin-treated groups were significantly higher (P<0.05) than C group. The total cell number 167 was significantly greater for M3-M4 compared with M1 and C blastocysts (Figure 1. a-f). 168

169 3.2 Effects of storage condition at 20°C

170 As shown in Table 2, supplementation of different concentration of melatonin to ovaries preservation medium led to significant increase in oocyte maturation rate evaluated based on cumulus expansion (p < p171 172 0.05). There were no significant differences between 500, 700, and 800  $\mu$ M treatment groups. No differences were observed in the maturation rate of oocytes from PBS alone (control) with 600 µM 173 melatonin (M2) group. Also, fertilization rate were not significant between M3 and M4 groups (p < 0.05). 174 Higher concentration of melatonin (M4; 800 µM) resulted in the best fertilization rate. Oocytes from 175 176 control group and low melatonin groups had inferior fertilization rate (p < 0.05). Similarly, the percent of 177 cleaved zygote in melatonin-treated groups was higher than the control group and the highest 178 concentration of melatonin resulted in higher cleavage rate (p<0.05), except compared to M3 group. 179 Moreover, the highest melatonin concentration showed the highest morula rates compared to all groups (P<0.05). No blastocyst was formed from the control group. Melatonin supplementation resulted in 180 significant increase in blastocyst formation with the higher doses (700 and 800 µM) resulting in higher 181 blastocyst rate than the lower doses (500 and 600 µM melatonin). Similar results were obtained for 182 blastocyst quality as compared by the mean number of blastomeric cells being significantly higher in M3 183 and M4 groups than the low melatonin groups (p < 0.05) (Figure 1. B-F). 184

185 4. Discussion

Preservation of ovarian tissue is challenging because there are several cell types within the tissue that 186 need to be maintained in a normal state. During the ovaries transportation to the laboratory, the stoppage 187 of blood flow reduces energy supply, and put ovaries under ischemic conditions. As well as, storage of 188 ovary at atmospheric level of oxygen (21%) could generate higher amounts of free radicals in follicle 189 190 microenvironment, which is toxic to oocyte and other cells surrounding it. Most likely due to the 191 formation of superoxide anion (O<sub>2</sub><sup>•-</sup>), this leads to the ulterior formation of HO<sup>•</sup>, H<sub>2</sub>O<sub>2</sub>, NO<sup>•</sup>, and ONOO<sup>-</sup>, 192 which results in oxidative stress [24, 35, 36]. The oocyte oxidative stress caused by ROS must be limited in order for a good embryo to be produced [36]. Free radicals spreads within follicle microenvironment, 193 and react with proteins, lipids and DNA of oocyte and cumulus cells leading to cell membrane-lipid 194 peroxidation, DNA damage and apoptosis [27, 37]. The results reported here show that the 195

supplementation of ovaries transport medium (PBS) with melatonin significantly maintained the developmental potential of the oocytes than PBS for 24 hours at 4 and 20°C. In particular, the oocyte competence for development the blastocyst stage was significantly improved as compared to control. Furthermore, high-dose melatonin (700 and 800  $\mu$ M) during 24 h storage in both 4, 20°C resulted in significant improvement in blastocyst quality as evidenced by greater numbers of blastomeric cells in these groups(Tables 1 and 2).

202 The melatonin directly suppresses free radicals derived from superoxide anion. In addition, metabolites that are formed during interaction with free radicals (i.e., cyclic 3-hydroxmelatonin, N1-acetyl-N2-203 formyl-5-methoxykynuramine, and N1-acetyl-5-methoxykynuramine) are high potent scavengers of toxic 204 reactants [24]. Furthermore, it is well known that melatonin has an important indirect function in up 205 regulation of antioxidant and down regulation of provident enzymes [30]. Brzezinski et al [38] reported 206 207 that melatonin concentration in human follicular fluid is significantly higher than serum samples. 208 Melatonin in follicular fluid besides their physiological role in steroidogenic mechanism, acts as free 209 radical scavenger [21]. The antioxidant activities of melatonin in oxidative stress has been reported in different species. In the mouse oocyte, melatonin promote the development rate of mouse two-cell 210 embryos to blastocysts stage [26]. Moreover, melatonin treatment increases intrafollicular concentration 211 212 of melatonin and improved fertilization rate embryo transfer [39]. A recent study presented that supplementation of IVM medium with melatonin-loaded lipid-core Nano capsules (Mel-LNC) during 213 214 IVEP improved cleavage and blastocyst rates of bovine embryo. Also, significantly decreased ROS 215 levels, and down-regulated the genes involved apoptosis caspase 3 (CASPA3) B and BAX [40]. Rodriguez-Osorio et al. [37]reported that, in IVC medium melatonin supplementation improved cleavage 216 217 rates in stressed conditions (H<sub>2</sub>O<sub>2</sub> or 40°C) and increased blastocyst cell numbers in usual terms. Our results showed a significant effect of melatonin on developmental potential of oocytes derived from 218 ovaries preserved in supplemented medium. In addition, our analysis demonstrated that the resultant 219 blastocysts in the M3 (700µM) and M4 (800µM) groups were better considering the healthy blastomeric 220 221 cells (Figure 1. E, e, F and f). These findings demonstrated that melatonin significantly reduced the toxic 222 effect of ROSs that generated in ovaries during storage period.

Although, improvement was noted when ovaries stored in melatonin supplemented medium, this positive 223 224 effects in low temperature  $(4^{\circ}C)$  were more evident than that of high temperature  $(20^{\circ}C)$ . The oocytes from the ovaries stored at 20°C without melatonin (control) lost their competence to develop to blastocyst 225 stages. Whereas, when ovaries stored at 4°C, obtained COCs in untreated group developed to blastocyst 226 stage, although, had significantly lower rate than in melatonin treated groups (Tables 1, 2). It is well 227 known that preservation of organs in hypothermia suppresses metabolic requirements. Consequently, it 228 229 results in reduced tissue damage caused by loss of ATP, increase in pH, lipid peroxidation and proteolysis 230 associated with ischemic condition [12, 20, 40]. Several studies investigated effects of temperature on pool of follicles in farm animal ovaries. In the Iberian red deer, preservation of ovaries in hypothermia 231 232 condition significantly elevated cleavage rate [4]. Nakao et al. [13]reported that oocyte obtained from 233 ovaries stored at 20°C had higher competence than that of stored at 39°C or 4°C. Results showed that 234 temperature of preservation media affected the quality of oocytes, in which cytoplasmic membrane, microtubule, cytoskeleton and zona pellucida might be sensitive to low temperatures [5, 12]. Canine 235 ovaries were transported for up to 4 hours at 4 and 35°C, obtained oocyte from the ovaries stored at 4°C 236 237 had higher MII maturation rate compared with the other group [16]. In the present study, the storage of sheep ovaries in low temperature media exhibited better results in terms of the developmental competenceof oocytes and the cleavage and production of the morula and blastocysts in contrast to high temperature.

240 In conclusion, the present study showed that (i) supplementation of phosphate buffer saline as ovine ovaries storage media with melatonin (500, 600, 700 and 800 µM), reduced the oocytes competence 241 decline to develop to blastocyst stage. (ii) for long preservation of ovaries (24 hours), low storage 242 243 temperature (4°C) can maintain oocyte competence but higher temperature (20°C) has a detrimental effect on oocyte viability, maturation, fertilization and subsequent development after IVF; and (iii) storage of 244 245 ovaries in high concentration (700, 800 µM) for 24 hours significantly maintain the oocyte quality and obtained blastocysts were better than other group, but this protective effect of melatonin were improved in 246 cool condition. Beneficial effect of melatonin in ovine ovaries storage medium may be employed to 247 reduce injuries mediated by ROS during ovaries preservation. 248

- 249 Acknowledgments
- 250 The authors would like to thank Helia Shamsi for her assistance during the laboratory procedure. We

251 would also like to express our appreciation to the University of Tehran for financial support of for this

252 project under grant number: 7108011/6/44. Similarly, the authors would like to acknowledge the financial

support of Iran National Science Foundation under grant number: 94810027.

254	Table 1: Effects of ovaries preservation for 24 h at 4°	C in melatonin supplemented PBS on oocyte and embryo
255	development.	

antioxidant dosage (µM)	Cumulus cells expansion%(n)	Fertilization%(n*)	Cleaved zygote on day 2 %(n)	Morula rate%(n)	Blast rate%(n)	Mean blastomere (n)
0	84.19±1.52 <sup>c</sup> (122/145)	72.12±2.88 <sup>c</sup> (18/25)	61.98±3.98 <sup>e</sup> (60/97)	34.09±2.09 <sup>c</sup> (33/97)	7.19±0.81° (7/97)	86.00±3.00 <sup>c</sup>
500	86.83±2.21 <sup>bc</sup> (120/138)	71.67±1.66 <sup>c</sup> (18/25)	68.34±1.66 <sup>de</sup> (65/95)	44.11±1.89 <sup>b</sup> (42/95)	10.56±0.55 <sup>b</sup> (10/95)	98.50±3.5 <sup>bc</sup>
600	90.18±2.68 <sup>abc</sup> (128/142)	76.79±1.78 <sup>bc</sup> (20/26)	72.28±0.27 <sup>cd</sup> (73/101)	44.57±1.43 <sup>b</sup> (45/101)	$10.90 \pm 1.10^{b}$ (11/101)	111.5±1.5 <sup>b</sup>
700	91.20±1.28 <sup>ab</sup> (142/156)	80.00±0.75 <sup>ab</sup> (24/30)	75.9±0.46 <sup>bc</sup> (85/112)	49.99±0.89 <sup>b</sup> (56/112)	16.95±0.59 <sup>a</sup> (19/112)	125.00±2.00 <sup>a</sup>
800	91.22±0.50 <sup>ab</sup> (135/148)	80.13±3.20 <sup>ab</sup> (20/25)	80.89±1.26 <sup>ab</sup> (89/110)	56.35±0.79 <sup>a</sup> (62/110)	18.19±0.33 <sup>a</sup> (20/110)	126.50±5.5 <sup>a</sup>

**256** Each value represents the means  $\pm$  SEM.

257 \* randomly selected from zygotes for assessment IVF parameters

Table 2: . Effects of ovaries preservation for 24 h at 20°C in melatonin supplemented PBS on oocyte and embryo
 development. No melatonin was added to the control group.

Antioxidant dosage (µM)	Cumulus cells expansion %(n)	Fertilization%(n *)	Cleaved zygote on day 2 %(n)	Morula rate%(n)	Blast rate%(n)	Mean blastomere (n)
0	75.50±1.64 <sup>b</sup> (119/158)	53.57±3.57 <sup>c</sup> (14/26)	32.42±5.59 <sup>c</sup> (30/93)	21.53±5.74 <sup>c</sup> (20/93)	0.00 <sup>c</sup> (0/93)	-

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500	83.13±1.88	62.08±1.19 <sup>b</sup>	52.53±3.47 <sup>b</sup>	33.38±1.24 <sup>b</sup>	5.83±0.17 <sup>b</sup>	83.50±4.50
	<sup>a</sup> (133/160)	(18/29)	(54/104)	(35/104)	(6/104)	c
600	79.75±0.93 <sup>b</sup> (138/173)	65.48±109 <sup>b</sup> (19/29)	60.67±2.98b <sup>b</sup> (65/109)	33.81±0.48 <sup>b</sup> (37/109)	7.48±0.21 <sup>b</sup> (8/109)	89.00±4.00
700	80.37±0.88	75.13±1.80 <sup>ab</sup>	70.82±1.18 <sup>a</sup>	41.24±2.35b <sup>b</sup>	10.36±0.36 <sup>a</sup>	105.50±5.5
	<sup>a</sup> (127/158)	(21/28)	(75/99)	(40/99)	(11/99)	0 <sup>ab</sup>
800	80.29±1.04	77.38±5.95 <sup>a</sup>	68.66±1.34 <sup>a</sup>	45.72±2.86 <sup>a</sup>	12.72±0.73 <sup>a</sup>	119.00±6.0
	<sup>a</sup> (130/158)	(20/26)	(70/104)	(47/104)	(13/104)	0 <sup>a</sup>

 $260 \qquad \ \ Each value represents the means \pm SEM.$ 

261 \* Randomly selected from zygotes for assessment IVF parameters

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Figure 1. Example of blastocysts 7 days. Stained with Hoechst fluorescent dye. SC: Blastocyst obtained in conventional method (up to 3h storage in 37.5°C). C: from control group (ovaries stored without melatonin

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- supplementation) at 4°C [B] or20°C [b]. M1-M4: Blastocysts from ovaries stored with 500, 600, 700 or 800 μM
- melatonin at 4 [C, D, E and F] or 20oC [c, d, e and f]. A high-resolution version of panels A, B, b, C, c, D, d, E, e, F
  and f of this image are available as Virtual Microscope eSlides: VM04430, VM04431, VM04432, VM04433,
- $200 \qquad \text{and 1 of this image are available as virtual interoscope condes. vivio++50, vivio++51, vivio++52, 200 \qquad \text{NM}04424 \text{ NM}04425 \text{ NM}04427 \text{ NM}04428 \text{ NM}04420 \text{ and NM}04440 \text{ measure stimulation}$
- **269** VM04434, VM04435, VM04436, VM04437, VM04438, VM04439 and VM04440, respectively.
- 270
- 271 References
- [1] Menchaca A, Anegon I, Whitelaw CBA, Baldassarre H, Crispo M. New insights and current tools for genetically
   engineered (GE) sheep and goats. Theriogenology. 2016;86:160-9.
- [2] Wani NA. In vitro maturation and in vitro fertilization of sheep oocytes. Small Ruminant Research. 2002;44:89 95.
- [3] Gonçalves R, Cavalcante A, Gouveia B, Lins T, Barberino R, Menezes V, et al. Lower apoptosis rate in ovine
   preantral follicles from ovaries stored in supplemented preservation media. Zygote. 2015;23:943-50.
- [4] García-Álvarez O, Maroto-Morales A, Berlinguer F, Fernández-Santos M, Esteso M, Mermillod P, et al. Effect
   of storage temperature during transport of ovaries on in vitro embryo production in Iberian red deer (Cervus elaphus
   hispanicus). Theriogenology. 2011;75:65-72.
- [5] Wongsrikeao P, OTOI T, KARJA NWK, AGUNG B, NII M, NAGAI T. Effects of ovary storage time and
   temperature on DNA fragmentation and development of porcine oocytes. Journal of Reproduction and
   Development. 2005;51:87-97.
- [6] Nagao Y, Harada Y, Yamaguchi M, Igarashi A, Ooshima Y, Kato Y. Antioxidant treatment during preservation
   of bovine ovaries increased the development potential of embryos. Zygote. 2010;18:315-21.
- [7] Cocchia N, Corteggio A, Altamura G, Tafuri S, Rea S, Rosapane I, et al. The effects of superoxide dismutase
  addition to the transport medium on cumulus–oocyte complex apoptosis and IVF outcome in cats (Felis catus).
  Reproductive biology. 2015;15:56-64.
- [8] Gomes R, Andrade E, Lisboa L, Ciquini A, Barreiros T, Fonseca N, et al. Effect of holding medium, temperature
   and time on structural integrity of equine ovarian follicles during the non-breeding season. Theriogenology.
   2012;78:731-6.
- [9] Hornick J, Duncan F, Shea L, Woodruff T. Isolated primate primordial follicles require a rigid physical
   environment to survive and grow in vitro. Human reproduction. 2012;27:1801-10.
- [10] Kamoshita K, Okamoto N, Nakajima M, Haino T, Sugimoto K, Okamoto A, et al. Investigation of in vitro
   parameters and fertility of mouse ovary after storage at an optimal temperature and duration for transportation.
   Human Reproduction. 2016:dew023.
- [11] Lucci CM, Kacinskis MA, Rumpf R, Báo SN. Effects of lowered temperatures and media on short-term
   preservation of zebu (Bos indicus) preantral ovarian follicles. Theriogenology. 2004;61:461-72.
- [12] Matsushita S, Tani T, Kato Y, Tsunoda Y. Effect of low-temperature bovine ovary storage on the maturation
   rate and developmental potential of follicular oocytes after in vitro fertilization, parthenogenetic activation, or
   somatic cell nucleus transfer. Animal reproduction science. 2004;84:293-301.
- [13] Nakao H, Nakatsuji N. Effects of storage conditions of bovine ovaries and oocytes on the success rate of in vitro fertilization and culture. Journal of Reproduction and Development. 1992;38:11-3.
- [14] Naoi H, Otoi T, Shimamura T, Karja NWK, Agung B, Shimizu R, et al. Developmental competence of cat
   oocytes from ovaries stored at various temperature for 24 h. Journal of Reproduction and Development.
   2007;53:271-7.

307 [15] Silva J, Lucci C, Carvalho F, Báo S, Costa S, Santos R, et al. Effect of coconut water and Braun-Collins
 308 solutions at different temperatures and incubation times on the morphology of goat preantral follicles preserved in
 309 vitro. Theriogenology. 2000;54:809-22.

- [16] Taş M, Evecen M, Özdaş ÖB, Cirit Ü, Demir K, Birler S, et al. Effect of transport and storage temperature of
   ovaries on in vitro maturation of bitch oocytes. Animal reproduction science. 2006;96:30-4.
- [17] Onions V, Mitchell M, Campbell B, Webb R. Ovarian tissue viability following whole ovine ovary
   cryopreservation: assessing the effects of sphingosine-1-phosphate inclusion. Human Reproduction. 2008;23:606 18.
- [18] Devine PJ, Perreault SD, Luderer U. Roles of Reactive Oxygen Species and Antioxidants in Ovarian Toxicity
   1. Biology of reproduction. 2011;86:Article 27, 1-10.
- 317 [19] Iwata H, Hayashi T, Sato H, Kimura K, Kuwayama T, Monji Y. Modification of ovary stock solution with
- magnesium and raffinose improves the developmental competence of oocytes after long preservation. Zygote.
   2005;13:303-8.
- [20] Guibert EE, Petrenko AY, Balaban CL, Somov AY, Rodriguez JV, Fuller BJ. Organ preservation: current concepts and new strategies for the next decade. Transfusion Medicine and Hemotherapy. 2011;38:125-42.
- 322 [21] Adriaens I, Jacquet P, Cortvrindt R, Janssen K, Smitz J. Melatonin has dose-dependent effects on
   323 folliculogenesis, oocyte maturation capacity and steroidogenesis. Toxicology. 2006;228:333-43.
- [22] Cruz M, Leal C, Cruz J, Tan D, Reiter R. Essential actions of melatonin in protecting the ovary from oxidative
   damage. Theriogenology. 2014;82:925-32.
- [23] Jiang J, Cheung C, Wang Y, Tsang B. Regulation of cell death and cell survival gene expression during ovarian
   follicular development and atresia. Frontiers in bioscience: a journal and virtual library. 2003;8:d222-37.
- 328 [24] Bonnefont Rousselot D, Collin F, Jore D, Gardès Albert M. Reaction mechanism of melatonin oxidation by
   329 reactive oxygen species in vitro. Journal of pineal research. 2011;50:328-35.
- [25] Galano A, Tan DX, Reiter RJ. On the free radical scavenging activities of melatonin's metabolites, AFMK and
   AMK. Journal of pineal research. 2013;54:245-57.
- [26] Ishizuka B, Kuribayashi Y, Murai K, Amemiya A, Itoh MT. The effect of melatonin on in vitro fertilization and
   embryo development in mice. Journal of pineal research. 2000;28:48-51.
- [27] Izadpanah G, Zare-Shahneh A, Zhandi M, Yousefian I, Emamverdi M. Melatonin has a beneficial effect on
   stallion sperm quality in cool condition. Journal of Equine Veterinary Science. 2015;35:555-9.
- [28] Mondal P, Hasan KN, Pal PK, Maitra SK. Influences of exogenous melatonin on the oocyte growth and
   oxidative status of ovary during different reproductive phases of an annual cycle in carp <em>Catla catla</em>.
   Theriogenology.87:349-59.
- [29] Tamura H, Nakamura Y, Korkmaz A, Manchester LC, Tan D-X, Sugino N, et al. Melatonin and the ovary:
   physiological and pathophysiological implications. Fertility and sterility. 2009;92:328-43.
- [30] R Ramis M, Esteban S, Miralles A, Tan D-X, J Reiter R. Protective effects of melatonin and mitochondria targeted antioxidants against oxidative stress: a review. Current medicinal chemistry. 2015;22:2690-711.
- [31] Dadashpour Davachi N, Zare Shahneh A, Kohram H, Zhandi M, Shamsi H, Hajiyavand AM, et al. Differential
  influence of ampullary and isthmic derived epithelial cells on zona pellucida hardening and in vitro fertilization in
  ovine. Reproductive Biology. 2016;16:61-9.
- [32] Dadashpour Davachi N, Kohram H, Zainoaldini S. Cumulus cell layers as a critical factor in meiotic
   competence and cumulus expansion of ovine oocytes. Small Ruminant Research. 2012;102:37-42.

- [33] Dadashpour Davachi N, Kohram H, Zare Shahneh A, Zhandi M, Goudarzi A, Fallahi R, et al. The effect of
   conspecific ampulla oviductal epithelial cells during in vitro maturation on oocyte developmental competence and
   maturation-promoting factor (MPF) activity in sheep. Theriogenology. 2017;88:207-14.
- [34] Dashti S, Zare Shahneh A, Kohram H, Zhandi M, Dadashpour Davachi N. Differential influence of ovine
  oviduct ampullary and isthmic derived epithelial cells on in vitro early embryo development and kinetic. Small
  Ruminant Research. 2016;136:197-201.
- [35] Fischer B, Bavister B. Oxygen tension in the oviduct and uterus of rhesus monkeys, hamsters and rabbits.
  Journal of reproduction and fertility. 1993;99:673-9.
- [36] Thompson J, Simpson A, Pugh P, Donnelly P, Tervit H. Effect of oxygen concentration on in-vitro
   development of preimplantation sheep and cattle embryos. Journal of reproduction and fertility. 1990;89:573-8.
- [37] Rodriguez Osorio N, Kim I, Wang H, Kaya A, Memili E. Melatonin increases cleavage rate of porcine
   preimplantation embryos in vitro. Journal of pineal research. 2007;43:283-8.
- [38] Brzezinski A, Seibel MM, Lynch HJ, Deng M-H, Wurtman RJ. Melatonin in human preovulatory follicular
   fluid. The Journal of Clinical Endocrinology & Metabolism. 1987;64:865-7.
- [39] Tamura H, Takasaki A, Miwa I, Taniguchi K, Maekawa R, Asada H, et al. Oxidative stress impairs oocyte
   quality and melatonin protects oocytes from free radical damage and improves fertilization rate. Journal of pineal
   research. 2008;44:280-7.
- 365 [40] Remião MH, Lucas CG, Domingues WB, Silveira T, Barther NN, Komninou ER, et al. Melatonin delivery by 366 nanocapsules during in vitro bovine oocyte maturation decreased the reactive oxygen species of oocytes and
- 367 embryos. Reproductive Toxicology. 2016;63:70-81.
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## 1 Highlight

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3 Effect of melatonin supplementation in ovaries storage medium for up 24 hours in 4 and 20°C

- 4 was studied on oocyte quality and embryo production in sheep.
- 5 Oocyte competence, fertilization rate, morula and blastocyst improved by melatonin (500, 600,
- 6~~700 and 800  $\mu M)$  supplementation in ovaries storage medium.
- 7 Storage of ovaries in high concentration (700, 800  $\mu$ M) for 24 hours significantly maintain the
- 8 oocyte quality and obtained blastocysts were better than other groups.