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Highlights (for review)

- *In vivo* matured oocytes have higher competence than *in vitro* matured oocytes.
- A single FSH injection prior to OPU can improve oocyte competence.
- *In vitro* pre- maturation enables to improve oocyte competence without FSH priming.

1 Review Article

2

3 Title: Follicle priming by FSH and pre-maturation culture to improve oocyte quality *in vivo* and *in*
4 *vitro*

5

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19

20 **Abstract**

21 Nowadays there is strong demand to produce embryos from premium quality cattle, and we
22 can produce embryos using oocytes collected from living premium animals by ovum-pick up (OPU)
23 followed by *in vitro* fertilization (IVF). However, the developmental competence of IVF oocytes to
24 form blastocysts is variable. The developmental competence of oocytes depends on the size and stages
25 of follicles, and follicle-stimulating hormone priming (FSH-priming) prior to OPU can promote
26 follicular growth and improve the developmental competence of oocytes. Furthermore, following the
27 induction of ovulation using an injection of luteinizing hormone or gonadotropin-releasing hormone
28 after FSH-priming, we can collect *in vivo* matured oocytes from ovulatory follicles, which show higher
29 developmental competence than oocytes matured *in vitro*. However, the conventional protocols for
30 FSH-priming consist of multiple FSH injection for 3 to 4 days, which is stressful for the animal and
31 labor-intensive for the veterinarian. In addition, these techniques cannot be applied to IVF of oocytes
32 collected from bovine ovaries derived from slaughterhouses, which are important sources of oocytes.
33 Here, we review previous research focused on FSH-priming, especially for collecting *in vivo* matured
34 oocytes and a simplified method for superstimulation using a single injection of FSH. We also
35 introduce the previous achievements using *in vitro* pre-maturation culture, which can improve the
36 developmental competence of oocytes derived from non-stimulated animals.

37 **Keywords:** FSH; Ovum pick-up; *In vivo* maturation; Single injection; *In vitro* pre-maturation

38 **1. Introduction**

39 *In vitro* fertilization (IVF) technology is now widely used commercially for producing
40 embryos in cattle [1]. Ultrasound-guided ovum-pick up (OPU) combined with IVF is used to produce
41 embryos in cattle for genetic improvement [1], and the efficiency of embryo production by OPU-IVF
42 is higher than that by *in vivo* embryo production [2, 3]. Although there has been significant progress
43 in the technology of IVF procedures, the rate of development to blastocysts from oocytes recovered
44 from non-superstimulated donors is rarely consistently exceeds 40 to 50% [4, 5]. When donors are
45 superstimulated, a significant increase in the rate of blastocyst is seen but can vary from 50 to 80%
46 [6]. However, *in vivo* maturation will remain the gold standard resulting in the highest rate of
47 blastocyst.

48 In mono-ovulatory species including cattle, the emergence of follicular growth is induced
49 by a surge-like secretion of follicle-stimulating hormone (FSH). Then, a dominant follicle is selected
50 as the level of FSH decreases due to the inhibitory effect of estradiol-17 β and inhibin secreted by
51 follicles themselves. The dominant follicle continues to grow in response to stimulation with
52 luteinizing hormone (LH), and this results in ovulation [7, 8]. In other words, most follicles degenerate
53 at immature stages, and only a small proportion of follicles will develop fully and ovulate in the life
54 time of a cow [7, 8]. During folliculogenesis, the diameter of the oocyte will increase during follicular
55 growth phases. Developmental competence is defined as the ability of an oocyte to acquire nuclear,
56 cytoplasmic and molecular maturation and thus the competence to produce a blastocyst following

57 fertilization that will result in a healthy offspring after embryo transfer [6]. Although most oocytes in
58 immature antral follicles (≥ 2 mm, diameter of the oocyte: approximately 110 μm) acquire competence
59 for nuclear maturation [9], most oocytes will not have acquired cytoplasmic and/or molecular
60 competence when compared to larger oocytes (120 μm) from follicles grown sufficiently *in vivo* [10,
61 11].

62 To enhance follicular growth and the developmental competence of oocytes, administration
63 of FSH before OPU (FSH-priming) has been conducted in many studies [12, 13]. However,
64 conventional FSH treatments require multiple intramuscular (im) administrations, which are stressful
65 for the animal and time-consuming for the veterinarian. In conventional *in vivo* embryo production by
66 superovulation followed by uterine flushing, many researchers have tried to simplify FSH treatment
67 [14]. Application of these methods for OPU should be evaluated.

68 Although the quality of oocytes before IVM is considered the critical factor for the outcome
69 of *in vitro* embryo production [15-17], there are some cases where exogenous FSH is not applicable
70 to promote the developmental competence of oocytes, such as oocytes derived from slaughterhouses
71 or low-responding animals [18]. Previous studies have suggested that the developmental competence
72 of oocytes can be improved by adding a pre-maturation *in vitro* step (pre-IVM) prior to IVM. Pre-
73 IVM inhibits germinal vesicle breakdown and holds the oocytes at the germinal vesicle (GV) stage to
74 acquire full developmental competence of oocytes *in vitro* during meiotic arrest [19].

75 In this review, we describe FSH-priming to improve developmental competence before

76 collection of oocytes. In particular, we focused on collecting *in vivo* matured oocytes and simplifying
77 the priming protocol using a single injection of FSH. In addition, we also examine pre-IVM to improve
78 the developmental competence of oocytes *in vitro*.

79

80 **2. FSH-priming**

81 **2.1. Collecting *in vivo* matured oocytes by FSH-priming following induction of an LH surge**

82 Several studies have suggested that *in vivo* matured oocytes are more developmentally
83 competent than *in vitro* matured oocytes (Table 1) [20-26]. Evidence of an increased developmental
84 competence following *in vivo* maturation include differences in mRNA transcription in oocytes [27-
85 29], size of the meiotic spindle [30], cytoplasmic maturation (distribution of cortical granules [26],
86 ATP content [30]), and developmental kinetics of embryos [26]. Bordignon *et al.* [20] reveal that
87 heifers treated with gonadotropin-releasing hormone (GnRH) after superstimulation exhibited an LH
88 surge within 3 h after treatment (34 h after prostaglandin F_{2α} (PGF_{2α}) injection). In the control group
89 subjected to superstimulation and PGF_{2α} injection without a GnRH injection, 40% of heifers also
90 exhibited an LH surge within 47 h (between 41 and 45 h) after PGF_{2α} treatment. Most of the oocytes
91 recovered from control heifers were in metaphase I (MI) stage regardless of the state of expansion of
92 the cumulus, whereas most oocytes (97%) with expanded cumulus cells in the GnRH-treated group
93 were at the metaphase II (MII) stage. As a result, the developmental competence of oocytes with
94 expanded cumulus cells was higher in the GnRH-treated group than the control when oocytes were

95 directly subjected to IVF without IVM after OPU (control: 40%, GnRH: 60%, Table 1) [20]. Rizos *et*
96 *al.* [22] showed that *in vivo* matured oocytes had higher developmental competence than oocytes from
97 follicles just before LH surge. These results indicate that a post-LH follicular environment induced by
98 a GnRH injection is essential to improve the developmental competence of bovine oocytes.

99 The maturation of bovine oocytes is known to initiate 6 h after the LH surge *in vivo* [31];
100 therefore, some researchers collected oocytes 6 h after an injection of LH or human chorionic
101 gonadotropin [32-34]. Although they suggested promotive effects of exogenous LH injections on
102 developmental competence [33, 34], waiting 6 h after LH was not long enough to permit oocytes to
103 complete maturation *in vivo* as expressed by non-expanded cumulus cells surrounding the oocytes.
104 Matoba *et al.* [24] reported that ovulation occurred at 29 to 32 h after a GnRH injection (average 30.0
105 h), and *in vivo* matured oocytes could be successfully collected 25 to 26 h after a GnRH injection.
106 Other researchers collected oocytes after 20 to 26 h after GnRH injection, and succeed in collecting *in*
107 *vivo* matured oocytes with higher developmental competence [20-22, 25, 26], whereas Sprícigo *et al.*
108 [25] reported that they could collect more MII oocytes 24 h after GnRH injection (85%) than at 20 h
109 after GnRH injection (31%). These results indicated that the suitable duration between GnRH injection
110 and OPU for collecting *in vivo* matured oocytes should be 24 to 26 h. Although *in vivo* matured oocytes
111 show higher developmental competence, handling of them will be difficult due to stickiness of
112 expanded cumulus investments. Matoba *et al.* [24] changed conical tubes for collecting oocytes after
113 5 to 6 follicle aspirations, making it easier to collect individual oocytes instead of clumps.

114 *In vivo* matured oocytes with expanded cumulus cells were directly subjected to IVF after
115 OPU in some studies [20, 22, 25], while oocytes in other studies were cultured in IVM medium for 3
116 to 24 h [20, 21, 23, 24, 26] (Table 1). Bordignon *et al.* [20] reported that when oocytes with expanded
117 cumulus cells were subjected to IVM for 24 h, 29.2% of them underwent spontaneous activation, and
118 caused lower developmental competence than *in vivo* matured oocytes directly subjected to IVF after
119 OPU (Table 1). The total maturation period initiated from LH surge in Bordignon *et al.*'s study was
120 48 h (*in vivo* maturation for 24 h and *in vitro* maturation for 24 h). Previously, we suggested that IVM
121 for longer than 30 h caused aging of oocytes, and these oocytes showed lower developmental
122 competence than oocytes matured *in vitro* properly for 22 h [35]. Taken together, 22 to 24 h of IVM
123 was too long for *in vivo* matured oocytes. In cattle, OPU for collecting oocytes with FSH treatment
124 was conducted at 25 to 26 h after GnRH injection because ovulation occurred at 29 to 32 h after GnRH
125 injection (average 30.0 h) [24]. Some researchers matured *in vitro* the oocytes for another 3 h after
126 OPU to synchronize the total maturation period *in vivo* and *in vitro* with the time after GnRH injection
127 and ovulation [24, 26]. Further studies are needed to optimize the duration for IVM for *in vivo* matured
128 oocytes.

129

130 **2.2. Simplification of the regimen for FSH-priming**

131 Superovulation after a single injection of FSH has been developed in conventional *in vivo*
132 embryo production by superovulation followed by uterine flushing [14]. This approach can be done in

133 one of two ways; (1) using a solvent that enabled FSH to be released slowly, such as
134 polyvinylpyrrolidone (PVP) [36, 37], aluminum hydroxide gel mixes [38], or hyaluronan-based slow
135 release formulation [39, 40], or (2) a single subcutaneous administration of high-dose FSH dissolved
136 in saline [36, 41, 42]. In both treatments, blood FSH is slowly absorbed into the general circulation
137 and induces the growth of multiple ovulatory follicles (PVP [36], aluminum hydroxide gel [38],
138 hyaluronan [43], and FSH dissolved in saline by subcutaneous (sc) injection [42]). The injection
139 methods tested for FSH-priming before OPU to simplify the regimen of FSH injection included PVP
140 [44], aluminum hydroxide gel [26], hyaluronan [43], FSH dissolved in saline by sc injection [33, 45,
141 46], im injection [33, 47, 48], or epidural injection [49] (Table 2). There is very limited information
142 about the efficiency of a single injection of FSH prior to OPU, but Vieira *et al.* [43] reported that a
143 single im injection of FSH dissolved in 0.5% hyaluronan resulted in similar plasma FSH profiles as
144 twice-daily FSH treatments, and a single injection increased the number of embryos per OPU-IVF. In
145 some studies, researchers conducted a single FSH injection prior to OPU using saline as a solvent of
146 FSH (sc [45], im [47], or a simultaneous injection of im and sc [33, 50]). The efficiency of these
147 treatments on the acquisition of oocyte developmental competence was not reported to be different
148 with that of non-stimulated animals [45, 47] or lower than that of conventional multiple FSH injection
149 [33].

150 It is known that the interval between the last FSH injection of multiple FSH injections and
151 OPU (“coasting period”) critically affects the developmental competence of oocytes [34, 51]. Nivet *et*

152 *al.* [51] suggested that a coasting period between 44 and 68 h showed better developmental
153 competence of blastocysts than a shorter period (20 h), although a longer coasting period (92 h)
154 decreased the developmental competence [51]. During the coasting period, a progressive hypoxia
155 occurs in follicles, which is related to the increase in apoptosis and inflammation of follicles [52]. This
156 follicular environment is similar to several preovulatory changes in the dominant follicle and
157 associated with improvement in the developmental competence of oocytes [19]. Reduction of the FSH
158 level during the coasting period was similar to the growth of a dominant follicle [53], because FSH
159 levels decrease for several days before ovulation in the natural estrous cycle [54]. After the reduction
160 in FSH level, the basal level of LH is supposed to maintain growth and prevent atresia of follicles
161 during the coasting period [13]. Although the coasting period is important to improve developmental
162 competence, there is little information about the suitable interval between a single FSH injection and
163 the OPU. Blondin *et al.* [48] conducted a study where a single bolus FSH injection was administered
164 to beef heifers, and oocytes were collected from animals soon after slaughter at 24, 48, and 72 h after
165 the FSH injection (Table 2). In that case, the developmental competence of oocytes was higher in
166 animals slaughtered 48 h after the FSH injection than those after 24 or 72 h. Furthermore, the
167 developmental competence of oocytes was higher when oocytes were collected 4 to 5 h after slaughter
168 than oocytes collected soon after slaughter (Table 2). This result cannot be applied directly to other
169 single-injection regimens, because the plasma circulation of FSH after a single FSH injection was
170 different between im and sc injections [55], and the type of animal and solvent can affect the transition

171 of FSH to general circulation from the injection site. For example, some researchers successfully
172 induced superstimulation using a single sc injection of FSH in beef cows [41, 42], but Takedomi *et al.*
173 [36] failed to induce superstimulation using a single sc injection in Holstein heifers. When FSH
174 dissolved in saline was subcutaneously administered into Holstein heifers, the plasma concentration
175 of FSH markedly increased within 3 h and was maintained until 9 h after administration [36]. Plasma
176 concentration of FSH decreased to the basal level after 36 h, and superovulation was not induced [36].
177 However, a FSH solution dissolved in PVP [36] results in a gradual increase in FSH plasma
178 concentrations that peak 12 h after administration. Then, FSH plasma concentrations decreased
179 gradually but were maintained at a higher level than the basal level for more than 48 h, and
180 superstimulation can be induced in Holstein heifers [36]. For the optimization of OPU followed by a
181 single FSH injection, further studies are needed to find out the appropriate coasting periods based on
182 the plasma dynamics of circulating FSH after the injection.

183

184 **2.3. Epidural area as an injection site of FSH to induce superstimulation**

185 Although there are some effective methods for superstimulation, the effectiveness of these
186 different treatments varies considerably, probably because of differences in the amount of
187 subcutaneous fat tissue in the animals [36, 41, 42]. To develop a more efficient method to simplify the
188 regimen of FSH injection, we took an idea from a study of human pharmacokinetics, which suggested
189 that alfentanil (an opioid analgesic drug) was slowly absorbed into the general circulation after

190 epidural administration in humans [56]. Therefore, we firstly compared the outcome of *in vivo* embryo
191 production by superovulation followed by uterine flushing between a conventional multiple FSH
192 injection and a single epidural FSH injection. We collected embryos from five Japanese black cows
193 given twice-daily im FSH administration (totally 20 armour units of Antrin R-10, approximately 200
194 international units, Kyoritsu Seiyaku, Tokyo, Japan) for 3 d (control) or a single epidural FSH injection
195 (30 armour units, approximately 300 international units). The number of transferable blastocysts after
196 epidural treatment (9.0 ± 6.0) was similar to that in the control group (4.7 ± 3.5 , $P = 0.10$). Furthermore,
197 we confirmed the efficiency of a single epidural FSH injection for OPU-IVF of cattle with low
198 productivity by *in vivo* embryo production. We conducted OPU for three Japanese black cows with
199 low embryo productivity given twice daily im FSH administration (total 30 armour units) or a single
200 epidural FSH injection (30 armour units). Although most follicles were less than 6 mm in diameter,
201 and the numbers of follicles and collected oocytes were similar between treatments, the rate of
202 transferable blastocysts in the epidural group was higher than that of the control (Table 3, $P < 0.0001$).
203 Further study is needed to reveal the mechanism of improved developmental competence of oocytes
204 after a single epidural FSH injection, and plasma FSH dynamics after epidural injection to optimize
205 the protocol such as the coasting period.

206

207 **3. Pre-maturation**

208 **3.1. Application of cyclic adenosine monophosphate (cAMP) modulators for pre-maturation**

209 In conventional *in vitro* embryo production including ultrasound-guided OPU-IVF, oocytes
210 derived from antral follicles larger than 2 mm in diameter are used [6, 57], in which the oocytes acquire
211 competence for meiotic resumption [9]. One of the reasons for the lower competence of oocytes is
212 precocious meiotic resumption. If meiotically competent oocytes are isolated from follicles, they can
213 resume meiosis spontaneously without ovulatory stimulation such as an LH surge [58]. However,
214 oocytes collected from living cattle without FSH-priming or from slaughterhouse-derived ovaries
215 originate from follicles of varied developmental stages [6, 9, 59]. This means that all oocytes are not
216 growing enough to acquire developmental competence, resulting in lower developmental competence
217 to the blastocyst stage. Meiotic arrest is caused by an increase in cAMP in oocytes, which is produced
218 by oocytes themselves [60, 61] or supplied from cumulus cells via gap junctions [62, 63]. To improve
219 the acquisition of developmental competence of oocytes, many researchers have cultured oocytes from
220 non-stimulated slaughterhouse-derived ovaries in conditions that prevent meiotic resumption by
221 controlling cAMP concentration before IVM (pre-IVM; Table 3) [64-71]. Reduction of cAMP
222 concentration can be achieved through the addition of phosphodiesterase (PDE), the enzyme that
223 degrades cAMP to 5'-AMP. Thus, inhibition of PDE activity has been applied in pre-IVM culture. 3-
224 isobutyl-1-methylxanthine (IBMX) is a non-specific PDE inhibitor that prevents a reduction in cAMP
225 levels and inhibits meiotic resumption in bovine oocytes [72-75]. Forskolin (FSK) is an activator of
226 adenylate cyclase, which promotes the synthesis of cAMP. Culture conditions with IBMX and
227 forskolin increase cAMP levels in bovine oocytes [76]. A culture system called the “simulated

228 physiological oocyte maturation” (SPOM) system consists of pre-IVM culture with 500 μ M IBMX
229 and 100 μ M FSK for 2 h before IVM, which promoted the blastocyst rate [64, 71] and cell numbers
230 in blastocysts [64]. More recently, extending pre-IVM culture from 2 to 6 h was reported to increase
231 the proportion of hatched blastocysts on day 8 and yielded a highest ratio of inner cell mass to total
232 cells on day 8 after IVF by increasing intra-oocyte reduced glutathione, which has important roles as
233 an antioxidant agent in oocyte maturation, fertilization, and embryonic development [66].

234

235 **3.2. Effect of the diameter of oocytes on the outcome of pre-IVM culture**

236 Otoi *et al.* [11] collected oocytes from follicles 1 to 7 mm in diameter in slaughterhouse-
237 derived ovaries, and morphologically healthy oocytes (three or more dense layers of cumulus cells,
238 evenly granulated cytoplasm) were divided into groups based on their diameters and subjected to *in*
239 *vitro* embryo production. They showed that the developmental competence of oocytes became higher
240 as the diameter of oocytes became larger [11]. Based on Otoi *et al.*'s studies [11], we speculated that
241 the most suitable duration of pre-IVM culture is dependent on the diameter of oocytes. We collected
242 bovine oocytes from slaughterhouse-derived ovaries and divided them into small-sized (110 to < 115
243 μ m) and large-sized (\geq 115 μ m) oocytes and subjected them to pre-IVM culture for 0, 5, or 10 h in
244 medium containing IBMX (500 μ M) and a low dose of FSH (2×10^{-6} units/mL, from porcine pituitary)
245 [69]. Before pre-IVM culture, all oocytes were at GV stage in both groups. Although approximately
246 90% of oocytes were still GV stage after pre-IVM for 5 h in both groups, half of the oocytes reached

247 metaphase I after pre-IVM for 10 h, indicating the spontaneous meiotic resumption of oocytes during
248 an extending pre-IVM culture in both groups. In large oocytes ($\geq 115 \mu\text{m}$), the percentage of
249 blastocysts after IVF was not different between the different duration of pre-IVM culture (31%).
250 However, pre-IVM culture for 5 h showed a higher blastocyst rate (16%) than for 0 h (9%) or 10 h
251 (8%) (Table 3). Although the mechanism underlying improved developmental competence in pre-IVM
252 for 5 h is unclear, we previously reported that the mitochondrial activity of *in vitro* grown oocytes
253 (105.9 to 122.7 μm) increased at 10 h of pre-IVM, then decreased after 20 h of pre-IVM [77]. Changes
254 in mitochondrial activity during pre-IVM were accompanied by developmental competence to form
255 blastocysts [77]. Similarly, mitochondrial activity may increase during the first 5 h of pre-IVM, then
256 decrease after 10 h of pre-IVM in *in vivo* grown small-sized oocytes (110 to $< 115 \mu\text{m}$). Further studies
257 are needed to define the appropriate duration of pre-IVM treatment in more detail.

258

259 **3.3. Culturing oocyte with natriuretic peptide precursor type C**

260 During follicular development, natriuretic peptide precursor type C (NPPC or CNP) derived
261 from mural granulosa cells, and its receptor derived from cumulus cells (natriuretic peptide receptor
262 2; NPR2), play important roles for inhibiting meiotic resumption [78]. In mice, NPPC derived from
263 granulosa cells promotes production of cyclic guanosine monophosphate (cGMP) by NPR2 in
264 cumulus cells [78]. Cumulus cell-derived cGMP inhibits the reduction of cAMP concentration by the
265 inhibition of PDE3A, an oocyte-specific phosphodiesterase, which is a trigger for meiotic resumption

266 [78]. As with mice, bovine cumulus cells express NPR2 mRNA [65, 68, 79] and protein [68], and
267 meiotic resumption of oocytes can be arrested during culture of cumulus-oocyte complexes (COCs)
268 with NPPC [65, 67, 68, 70, 80]. Although NPR2 was not expressed in oocytes in mice [68], some
269 studies reported the expression of NPR2 mRNA [68, 79] and protein [68] in bovine oocyte membranes,
270 and meiotic resumption of denuded oocytes was arrested by NPPC [68]. Some studies suggested that
271 pre-IVM with NPPC (100 nM or 200 nM) for 6 h improved the blastocyst rate [67, 68], the blastocysts
272 cell number [65, 67], and the blastocyst hatching rate [65] (Table 3). The combination of NPPC (100
273 nM) and IBMX (500 μ M) in pre-IVM culture followed by 20 h IVM also showed a higher blastocyst
274 rate [70] (Table 3).

275

276 **4. Conclusion**

277 *In vivo* matured oocytes have higher developmental competence than *in vitro* matured
278 oocytes, but there is still a need for further research to optimize IVM conditions that will result in the
279 acquisition of developmental competence for immature bovine oocytes. Although a single injection of
280 FSH prior to OPU can improve the developmental competence of oocytes similar to conventional
281 multiple FSH injection, further study, such as of the coasting period, is needed to maximize the
282 potential of oocytes. For oocytes collected from cattle without FSH-priming, pre-IVM is a candidate
283 method to improve the developmental competence of oocytes. In addition, the diameter of oocytes is
284 an important criterion to affect the optimal duration of pre-IVM culture and the developmental

285 competence of oocytes.

286

287 **Competing interests**

288 The authors have no competing interest in publishing findings of this research.

289

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Table 1. Summary of data published on the developmental competence of *in vivo* matured bovine oocytes.

Authors	Injection manner of FSH or equine chorionic gonadotropin (eCG)	Time between GnRH or LH treatment and OPU	IVM duration for <i>in vivo</i> matured oocytes	Developmental competence (%) *
Bordignon <i>et al.</i> 1997 [20] **	FSH injection in eight decreasing doses 12 hours apart for 4 d	26 h	0 h or 24 h	Blastocysts on Day 7 FSH-only-IVM 0 h: 40 ^a FSH-GnRH-IVM 0 h: 60 ^b FSH-only-IVM 24 h: 20 FSH-GnRH-IVM 24 h: 13
van de Leemput <i>et al.</i> 1999 [21]	A single injection of eCG followed by an injection of eCG antibody (112 h later)	24 h	Maximally 2 h 54 min	Blastocysts on Day 11 Slaughterhouse: 26.4 ^a eCG-GnRH: 49.3 ^b
Rizos <i>et al.</i> 2002 [22] ***	FSH injection in eight decreasing doses twice daily for 4 d	20 h	0 h	Blastocysts on Day 7 Slaughterhouse (2 to 6 mm): 31.7 ^a (> 6 mm): 38.4 ^{ab} FSH-only: 35.3 ^a FSH-GnRH: 48.5 ^b
Dias <i>et al.</i> 2013 [23] ****	FSH injection in eight or 14 consistent doses twice daily for 4 or 7 d (Short FSH or Long FSH)	24 h	6 h	Morulae and blastocysts on Day 9 Short FSH: 24.7 ^{ab} FSH starvation: 18.1 ^b Long FSH: 36.6 ^a
Matoba <i>et al.</i> 2014 [24] *****	FSH injection in eight decreasing doses twice daily for 4 d	25 to 26 h	3 h	Good-quality blastocyst until Day 9 Non-stimulated: 36.1 Dominant follicle ablation: 54.9 ^a GnRH: 21.5 ^b
Sprcigo <i>et al.</i> 2015 [25]	FSH injection in eight decreasing doses twice daily for 4 d	24 h	0 h	Blastocysts on Day 7 Slaughterhouse: 37.9 ^a Non-stimulated: 50.6 ^a FSH only: 58.8 ^b FSH-GnRH: 62.4 ^b
Egashira <i>et al.</i> 2019 [26]	A single sc FSH injection using aluminum hydroxide gel	25 to 26 h	3 h	Good-quality blastocyst on Day 8 Non-stimulated: 29.5 ^a FSH-GnRH: 45.6 ^b

*: The definitions of each experimental group are described below (Day 0 = Day of IVF).

FSH-only: Groups of animals subjected to FSH-priming, but without GnRH injection later.

FSH (or eCG)-GnRH: Groups of animals given GnRH after FSH (or eCG)-priming for collecting *in vivo* matured oocytes.

Slaughterhouse: Oocytes were collected from slaughterhouse-derived ovaries to serve as a control.

Non-stimulated: Groups of animals with collected oocytes without FSH-priming and GnRH injection.

** : Collected oocytes were subjected to IVM for 0 or 24 h.

*** : Collected oocytes from slaughterhouse-derived ovaries were classified by the diameter of follicles (2 to 6 mm or > 6 mm)

**** : Cows were treated with three different FSH treatments as described below.

Short FSH: FSH was administered (im) in eight consistent doses twice daily for 4 d.

FSH starvation: FSH was administered (im) in eight doses twice daily for 4 d, and OPU was conducted 4.5 d after the final FSH injection.

Long FSH: FSH was administered (im) in eight consistent doses twice daily for 4 d.

***** : Developmental competence was compared between two methods (Dominant follicle ablation or GnRH injection) for the follicular wave control 1.5 d prior to FSH-priming.

^{a, b}: $P < 0.05$

Table 2. Summary of data published on the developmental competence of bovine oocytes after a single FSH injection.

Authors	Breeds	Injection site for single FSH	Solvent for single FSH	Coasting period	Developmental competence (%) *
Bungartz <i>et al.</i> 1994 [47]	Lactating Holstein cow	im	Saline	4 d	Morulas and blastocysts on Day 7 Non-stimulated: 2.9 Single FSH: 3.8
Ooe <i>et al.</i> 1996 [44] **	Cyclic lactating Holstein cows	im	30% PVP (10 mL)	48 h	Blastocysts on Day 8 Single FSH on Day 1: 25 Single FSH on Day 7: 28 Single FSH in pregnant cows: 33 Single FSH on Day 8 to 14: 29 Embryos (≥ 64 cells) on Day 6
Blondin <i>et al.</i> 1997 [48] ***	Beef heifers	im	Saline	Experiment 1: 24, 48, or 72 h Experiment 2: 48 h + 1 to 2 h or 4 to 5 h after slaughter	Experiment 1: Non-stimulated: 15 ^a FSH + 24 h: 5 ^b FSH + 48 h: 25 ^a FSH + 72 h: 7 ^b Experiment 2: Non-stimulated: 18 ^a FSH 48 h + 1 to 2 h: 24 ^{ab} FSH 48 h + 4 to 5 h: 41 ^b
Goodhand <i>et al.</i> 2000 [45]	Cyclic beef \times Friesian cows	sc	Saline	Multiple: 1 d Single: 3 d	Transferable embryos on Day 7 Non-stimulated: 43 Single FSH: 33 Multiple FSH: 35
Chaubal <i>et al.</i> 2007 [33] ****	Angus cross cows	im and sc	Saline	54 h	Blastocysts on Day 7 Multiple FSH-LH: 21.7 Multiple FSH: 18.7 Single FSH-LH: 18.8 Single FSH: 17.2
Vieira <i>et al.</i> 2016 [43]	Non-lactating Holstein cows	im	0.5% hyaluronan	Multiple FSH: 1.5 d Single FSH: 3 d	Blastocysts on Day 6 Non-stimulated: 25.9 Multiple FSH: 30.3 Single FSH: 30.3
Sakaguchi <i>et al.</i> 2018 [49]	Japanese black cows	Epidural	Saline (5 mL)	Multiple FSH: 21 to 23 h Single FSH: 75 to 78 h	Blastocysts on Day 7 Multiple FSH: 10.5 ^a Single FSH: 26.2 ^b
Egashira <i>et al.</i> 2019 [26] *****	Japanese black cows	sc	Aluminum hydroxide gel	4.5 d	Good-quality blastocyst on Day 8 Non-stimulated: 29.5 ^a Single FSH-GnRH: 45.6 ^b
Sakagami <i>et al.</i> 2019 [46] *****	Japanese black cows	sc	Saline (50 mL)	72 h	Blastocysts on Day 8 Single FSH: 22.1 ^a Single FSH-pre-IVM: 39.1 ^b

*: The definition of each experimental group is described below (Day 0 = Day of IVF).

Non-stimulated: Groups of animals with collected oocytes without FSH-priming and GnRH injection.

Single FSH: Groups of animals subjected to a single FSH injection.

Multiple FSH: Groups of animals subjected to multiple FSH injections for few days.

** : There were four experimental groups as described below.

Single FSH on Day 1: FSH was administered (im) on Day 1 (Day 0 = ovulation), and OPU was conducted on Day 3.

Single FSH on Day 7: DFA was conducted on Day 6, FSH was administered (im) on Day 7, and OPU was conducted on Day 9

Single FSH in pregnant cows: FSH was administered (im) on 70, 75, 80, 85, and 90 d of pregnancy and OPU was conducted 48 h later (5 times at 5 d intervals).

Single FSH on d 8 to 14: FSH was administered (im) on Days 8 to 14 and OPU was conducted 48 h later

***: Oocytes were collected from ovaries after slaughter. In experiment 1, animals were slaughtered at 24, 48, or 72 h after a single FSH injection then oocytes were collected soon after (FSH + 24, 48, or 72 h groups). In experiment 2, animals were slaughter at 48 h after a single im FSH injection, and oocytes were collected 1 to 2 h or 4 to 5 h after slaughter (FSH 48 h + 1 to 2 h or 4 to 5 h groups).

****: In the single FSH group, FSH was given simultaneously by two routes (im and sc). In Multiple FSH-LH and Single FSH-LH groups, LH was injected 6 h prior to OPU (48 h after the end of FSH treatment).

*****: In single FSH-GnRH group, GnRH was administered (sc) 25 to 26 h prior to OPU (2 d after a single FSH injection).

*****: In single FSH-pre-IVM group, collected oocytes were subjected to pre-IVM for 2 h before IVM.

^{a, b}: $P < 0.05$

Table 3. Summary of data published on the developmental competence of bovine oocytes subjected to pre-IVM culture.

Authors	cAMP modulator, NPPC, or FSH in pre-IVM medium	Duration of pre-IVM	Duration of IVM	Developmental competence (%) *
Albuz <i>et al.</i> 2010 [64]	FSK (100 µM), IBMX (500 µM)	0 (control) or 2 h	24 h or 30 h	Blastocysts/cleaved on Day 8 Control IVM 24 h: 22 ^a Pre-IVM IVM 24 h: 48 ^b Control IVM 30 h: 27 ^a Pre-IVM IVM 30 h: pre-IVM: 69 ^c
Franciosi <i>et al.</i> 2014 [65]	Recombinant human FSH (10 ⁻⁴ unit/mL) + NPPC (100 nM) or cilostamide (10 µM) or without NPPC and cilostamide (control)	8 h	22 h	Expanded and hatched blastocysts on Day 9 Control: 78 ^a Cilostamide ^{**} : 94 ^b NPPC: 93 ^b
Li <i>et al.</i> 2016 [66]	FSK (100 µM), IBMX (500 µM)	0 (control), 2, 4, or 6 h	Control: 24 h Pre-IVM 2 h: 24 h Pre-IVM 4 h: 22 h Pre-IVM 4 h: 20 h	Blastocysts/cleaved on Day 8 Control: 26.3 ^a Pre-IVM 2 h: 39.2 ^b 4 h: 35.2 ^b 6 h: 34.2 ^b
Zhang <i>et al.</i> 2017 [67]	NPPC (200 nM)	0 (control) or 6 h	24, 28, or 32 h	Blastocysts on Day 7 Control IVM 24 h: 32.2 ^b IVM 28 h: 15.0 ^a IVM 32 h: 0 Pre-IVM IVM 28 h: 51.6 ^c
Xi <i>et al.</i> 2018 [68]	NPPC (200 nM)	0 (control) or 6 h	24, 26, or 28 h	Blastocysts on Day 7 Control IVM 24 h: 23.5 ^a 26 h: 24.1 ^a 28 h: 21.7 ^a Pre-IVM IVM 24 h: 26.9 ^a 26 h: 45.2 ^b 28 h: 41.6 ^b
Abdel-Ghani <i>et al.</i> 2018 [69]	Porcine pituitary FSH (2 × 10 ⁻⁶ units/mL), IBMX (500 µM)	0, 5, or 10 h	22 h	Blastocysts on Day 7 (110 to < 115 µm) *** Control: 9 ^a Pre-IVM 5 h: 16 ^b 10 h: 8 ^{ab}
Soto-Heras <i>et al.</i> 2019 [70]	NPPC (200 nM), IBMX (500 µM)	0 (control) or 6 h	24 (control) or 20 h	Blastocysts on Day 8 Control: 34.5 Pre-IVM: 45.1
Hashimoto <i>et al.</i> 2019 [71]	FSK (100 µM), IBMX (500 µM)	0 (control) or 2 h	24 h	Blastocysts on Day 7 Control: 16.7 ^a Pre-IVM: 27.5 ^b Blastocysts on day 8 Control: 25.0 ^a Pre-IVM: 33.3 ^b

*: Day 0 = Day of IVF

** : Specific inhibitor of PDE3 (specific PDE of oocytes).

***: Collected oocytes from slaughterhouse-derived ovaries were classified by their diameter (110 to < 115 μm or $\geq 115 \mu\text{m}$)

a, b, c: $P < 0.05$