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Exposure to follicular fluid during oocyte maturation and oviductal fluid during post-maturation does not improve *in vitro* embryo production in the horse

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Summary

Most wild equids and many domestic horse breeds are at risk of extinction, so there is an urgent need for 11 genome resource banking. Embryos cryopreservation allows the preservation of genetics from male and 12 13 female and is the fastest method to restore a breed. In the equine, embryo production *in vitro* would allow the production of several embryos per cycle. Intracytoplasmic sperm injection (ICSI) is used 14 to generate horse embryos, but it requires expensive equipment and expertise in micromanipulation, 15 and blastocyst development rates remain low. No conventional in vitro fertilization (IVF) technique 16 for equine embryo production is available. The development of culture conditions able to mimic the 17 maturation of the oocyte in preovulatory follicular fluid (pFF) and the post-maturation in oviductal fluid 18 19 (OF) may improve embryo production *in vitro*. Our aim was to analyse the effect of *in vitro* maturation in pFF and incubation in OF on *in vitro* maturation of equine oocytes, fertilization using conventional 20 21 IVF or ICSI, and embryo development after culture in synthetic oviductal fluid (SOF) or DMEM-F12. Oocytes collected from slaughtered mares or by ovum pick up were matured in vitro in pFF or semi-22 synthetic maturation medium (MM). The in vitro maturation, fertilization and development rates were 23 24 not statistically different between pFF and MM. After in vitro maturation, oocytes were incubated with 25 or without OF. Post-maturation in OF did not significantly improve the fertilization and development rates. Thus, in our study, exposure to physiological fluids for oocyte maturation and post-maturation 26 does not improve *in vitro* embryo production in the horse. 27

28 Keywords: Equine, IVM, IVF, Oocyte, Spermatozoa

Introduction

Most wild equids are currently endangered or threatened in the wild, such as the Asiatic wild ass or Grevy's zebra, as mentioned in the IUCN Red List of endangered animal species of the International Union for the Conservation of Nature (Adams *et al.*, 2009). Moreover, many domestic horse breeds are at risk of

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extinction, with less than 300 active breeding mares, 36 37 and several breeds that are nearly extinct with fewer than 100 active breeding mares, such as the Faer Island 38 39 pony in Denmark, Estonian draught horse in Estonia, Landais pony and Grand Noir du Berry donkey 40 in France (Smits et al., 2012b). The actions that are 41 42 currently undertaken to preserve endangered horse 43 breeds include the creation of a Genome Resource Bank. 44

Genome resource banking requires cryopreservation 45 46 of semen, oocytes and/or embryos. Embryo cryopreservation allows the preservation of genetics from 47 both male and female and is the fastest method to 48 restore a breed. In equids, embryo production in 49 50 vivo is limited, as experimental induction of multiple ovulations has a low efficiency (Meyers-Brown et al., 51 2011) and routine induction of multiple ovulations is 52 still ineffective (Smits et al., 2012b). Embryo production 53 in vitro allows the production of several embryos 54 55 per cycle (Hinrichs, 2012). Intracytoplasmic sperm 56 injection (ICSI) has been widely adopted to generate 57 horse embryos in vitro, both for scientific purposes 58 and in the horse breeding industry (Choi *et al.*, 2011; 59 Hinrichs, 2012). This method allows the investigation 60 of specific aspects of fertilization in the horse, such as sperm chromatin-ooplasm interactions, overcoming 61 62 limited efficiency of *in vitro* sperm penetration through oocyte barriers. However ICSI requires expensive 63 equipment and expertise in micromanipulation, and 64 very few laboratories worldwide routinely produce 65 66 equine embryos following ICSI. Moreover, blastocyst 67 development rates following ICSI do not exceed 40% per cleaved oocyte (Hinrichs, 2012). Over the last 68 decades, several attempts to establish an efficient in 69 70 vitro fertilization (IVF) technique in the equine were performed. Palmer and collaborators published the 71 72 first paper on IVF of equine oocytes (Bézard *et al.*, 1989) and obtained the first and only two IVF-produced 73 74 foals (Palmer et al., 1991) using preovulatory oocytes 75 and fresh sperm treated with calcium ionophore. However, this technique did not yield IVF rates 76 higher than 36% (Palmer et al., 1991; Alm et al., 77 78 2001) and, although low IVF rates have been reported sporadically (Hinrichs *et al.*, 2002), this technique was 79 not repeatable (Mugnier et al., 2009). Dell'Aquila and 80 collaborators reported a 32% rate of equine IVF after 81 82 incubation of in vitro matured oocytes subjected to 83 partial cumulus removal, with frozen sperm treated 84 with heparin (Dell'Aquila et al., 1996) but these results could not be replicated (Dell'Aquila et al., 1997a, b). 85 Low rates of IVF (0-36%) were obtained by Alm 86 and collaborators (Alm et al., 2001) after exposure 87 88 of spermatozoa to calcium ionophore or heparin. 89 Since then, no reports showing the efficiency of one of these IVF techniques have been published. In 90 91 2009, a 60% rate of fertilized oocytes was reported after treatment of fresh spermatozoa with procaine 92 to induce hyperactivated motility (McPartlin et al., 93 2009). Using the same technique, we obtained 37% of 94 fertilized oocytes (Ambruosi et al., 2013), but Leemans 95 and collaborators showed that equine IVF embryos 96 fail to develop beyond the 8-16-cell stage (Leemans 97 et al., 2015). Thus, to date, no efficient conventional 98 IVF technique for equine embryo production in vitro 99 is available. 100

During its journey in the follicle and the oviduct, 101 the oocyte acquires factors necessary for fertilization 102 and early development. The somatic environment 103 is of crucial importance for oocyte preparation for 104 fertilization and development (Coy et al., 2012; Aviles 105 et al., 2010). The development of in vitro culture 106 conditions able to mimic the maturation of the oocyte 107 in follicular fluid and the post-maturation of the 108 ovulated oocyte in OF may help to improve embryo 109 production in vitro. For example, addition of follicular 110 fluid to maturation medium of equine oocytes 111 increases fertilization and cleavage rate (Dell'Aquila 112 et al., 1997b). Pre-incubation of equine oocytes with OF 113 or oviductal cells increases fertilization rate (Mugnier 114 et al., 2009) (Ambruosi et al., 2013). Moreover, several 115 studies have shown the crucial role played by the 116 oviduct in the preparation of equine gametes for 117 fertilization (Goudet, 2011). 118

Several media have been used for in vitro culture of 119 equine zygotes in different labs: Synthetic OF (SOF) 120 medium (Tremoleda et al., 2003), DMEM-F12-51445C 121 medium (in ME Dell'Aquila laboratory), DMEM-F12-122 D8437 medium (in G Goudet lab), DMEM-F12-D8900 123 medium (Choi et al., 2011; Martino et al., 2016). 124 However, these medium have never been used in the 125 same laboratory and conditions. 126

Our aim was to analyse the effect of in vitro 127 maturation in follicular fluid and pre-incubation in OF 128 on in vitro maturation, fertilization using conventional 129 IVF or ICSI, and development of equine oocytes. For 130 this purpose: (1) we compared a semi-synthetic in 131 vitro maturation medium with preovulatory follicular 132 fluid, in which maturation naturally occurs; (2) we 133 analysed the influence of pre-incubation of oocytes 134 with OF before fertilization, which occurs in vivo after 135 ovulation of the oocyte into the oviduct; (3) we tested 136 four culture media for the *in vitro* development of 137 fertilized oocytes using conventional IVF or ICSI. 138

Materials and Methods

All procedures on animals were conducted in ac-
cordance with the guidelines for the care and use of
laboratory animals issued by the French Ministry of
Agriculture and with the approval of the ethical review140
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144 committee (Comité d'Ethique en Expérimentation
145 Animale Val de Loire) under numbers 2011/6 and
146 02701.01. The study was conducted in France and
147 in Italy (48° and 41° North parallel) during two
148 subsequent breeding seasons.

All chemicals were purchased from Sigma-Aldrich
(Milano, Italy and St Quentin Fallavier, France) unless
otherwise indicated.

152 Collection of equine immature oocytes

Equine immature cumulus-oocyte complexes (COCs)
were collected during the breeding season either
from slaughtered mares in commercial abattoirs in
France and Italy or by transvaginal ultrasound-guided
aspiration on experimental mares in France.

158 For COCs collection from slaughtered mares, ovaries from females of unknown reproductive history were 159 obtained at local commercial abattoirs immediately 160 161 after females were slaughtered. They were transported 162 to the laboratory within 2 h in 0.9% (w/v) NaCl 163 at 32-38°C. In the French laboratory, COCs were 164 collected using the aspiration procedure previously 165 described by Goudet and collaborators (Goudet et al., 2000). Briefly, the tunica albuginea was removed 166 167 and all follicles larger than 5 mm were aspirated 168 with an 18-gauge needle at 100 mmHg of vacuum pressure, the ovaries were cut into thick sections 169 170 with a scalpel blade to find other follicles within the ovarian stroma. Follicular fluids were examined under 171 172 a stereomicroscope for COCs recovery. In the Italian 173 laboratory, COCs were collected using the scraping 174 procedure as previously described (Dell'Aquila et al., 2001). All follicles from 5 to 25 mm in diameter 175 176 were opened with a scalpel blade and the granulosa cells layer scraped with a curette, COCs were 177 178 identified in the collected mural granulosa cells by 179 using a dissection microscope. In both laboratories, 180 oocytes denuded of cumulus and degenerated oocytes 181 showing shrunken, dense or fragmented cytoplasm, were discarded. 182

For COCs collection by transvaginal ultrasound-183 184 guided aspiration (ovum pick up: OPU), adult cyclic pony mares from our experimental stud were used. 185 Ovarian activity was assessed by routine rectal 186 ultrasound scanning to choose mares with several 187 follicles from 5 to 25 mm. Follicles were punctured 188 189 by transvaginal ultrasound-guided aspiration with a 190 double-lumen needle (length 700 mm, outer diameter 2.3 mm, internal diameter 1.35 mm, Casmed, Cheam, 191 192 Surrey, England) and a sectorial probe (Aloka SSD900) as previously described (Goudet et al., 1997). After 193 194 follicular fluid aspiration, the follicle was flushed with PBS (Phosphate Buffered Saline, Dulbecco A, 195 Oxoid, Basingstoke, Hampshire, England) and heparin 196 197 (Choay, Sanofi Aventis 5000 IU/ml) at 38°C. All aspirated fluids were examined for oocyte recovery, and 198 oocytes denuded of cumulus or degenerated oocytes 199 showing shrunken, dense or fragmented cytoplasm 200 were discarded. During the collection procedure, 201 mares were injected with detomidine (Medesedan[®], 202 0.25 ml/animal i.v., 10 mg/ml detomidine, Centravet, 203 Plancoet, France) and butorphanol (Dolorex[®], 0.6 204 ml/animal i.v., 10 mg/ml butorphanol tartrate and 205 0.1 mg/ml benzethonium chloride, Centravet) for 206 sedation and analgesia, dipyrone and butylsco-207 polamine (Estocelan[®], 15 ml/animal i.v., 4 mg/ml 208 butylscopolamine and 500 mg sodique metamizole, 209 Centravet) for analgesia and antispasmodia. After 210 puncture, the mares received a preventive antibiotic 211 injection (Depocilline, 20 ml/animal i.m., benzylpeni-212 cillin 170.41 mg/ml Intervet, Beaucouze, France). 213

In vitro maturation (IVM) of equine immature oocytes

Just after collection, COCs were washed in Medium 216 199 with Earle's salts, 25 mM HEPES and NaHCO₃ 217 supplemented with 20% (v/v) fetal calf serum 218 (FCS) and 25 µg/ml gentamycin. They were then 219 cultured in group of 10 to 30 for 27 h in an 220 atmosphere of 5% CO2 in air at 38.5°C in 100% 221 humidity in 500 µl of maturation medium (MM) or 222 100% equine preovulatory follicular fluid (pFF). The 223 maturation medium was Medium 199 with Earle's 224 salts supplemented with 20% (v/v) FCS and 50 ng/ml 225 epidermal growth factor (Goudet et al., 2000). The 226 equine pFF was collected by transvaginal ultrasound-227 guided aspiration on three adult cyclic pony mares 228 from our experimental stud. Ovarian activity was 229 assessed by routine rectal ultrasound scanning. At 230 the emergence of a follicle larger than 33 mm in 231 diameter, the mare was injected with 1500 IU human 232 chorionic gonadotropin (hCG, i.v., Chorulon, Intervet). 233 The preovulatory follicle was punctured 35 h after 234 hCG injection, just before ovulation, by transvaginal 235 ultrasound-guided aspiration with a single-lumen 236 needle (length 600 mm, outer diameter 1.8 mm, 237 Thiebaud Freres, Jouvernex Margencel, France) and a 238 sectorial probe as previously described. The presence 239 of a metaphase II oocyte was ascertained using nuclear 240 chromatin configuration analysis as described below. 241 The pFF was centrifuged at 1500 g for 10 min at 4° C. 242 The supernatants were pooled and kept at -20° C. 243

Post-maturation of equine oocytes in OF

After IVM, equine COCs were partially denuded and245incubated in droplets of 30 μ l of porcine OF for24630 min in an atmosphere of 5% CO2 in air at 38.5°C247in 100% humidity. For porcine OF collection, genital248tracts from gilts were obtained at a commercial abattoir249and transported to the laboratory at room temperature.250

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251 Genital tracts with both ovaries containing several 252 follicles larger than 5 mm were used. The oviducts 253 were dissected free from surrounding tissues. The 254 oviductal content from the ampulla was expelled by 255 gentle squeezing using a sterile microscope slide and 256 collected by introducing the tip of a pipette into 257 the ampulla and aspirating while making a manual 258 ascendant pressure from the isthmus to the ampulla 259 (Carrasco et al., 2008). After centrifugation at 10,000 g for 15 min, the supernatant containing secreted and 260 intracellular components was immediately stored at 261 -20°C until use as 'OF'. 262

In vitro fertilization (IVF) procedure 263

The IVF procedure was performed in France, using 264 oocytes collected from slaughtered mares or by OPU. 265

Preparation of equine sperm and IVF 266

267 Fresh equine semen was collected with a closed 268 artificial vagina from three adult Welsh pony stallions of proven fertility from our experimental stud. Semen 269 270 was filtered through gauze, sperm motility was 271 visually evaluated under light microscopy on a heated 272 stage and sperm concentration was assessed using a 273 spectrophotometer (Ciba-Geigy). Then, 1 ml of semen 274 was diluted in 2 ml of pre-warmed modified Whitten's 275 Medium (MW; 100 mM NaCl, 4.7 mM KCl, 1.2 mM 276 MgCl₂, 22 mM HEPES, 4.8 mM lactic acid hemicalcium salt, 1 mM pyruvic acid) supplemented with 5.5 mM 277 278 glucose (anhydrous), pH 7.25 (McPartlin et al., 2009). 279 Diluted sperm was transported to the laboratory 280 within a few minutes at 37°C and centrifuged in 281 15 ml conical tubes at 100 g for 1 min at 37°C 282 to remove particulate matter and dead sperm. The 283 supernatant was then transferred to a 14 ml round-284 bottom tube and centrifuged at 600 g for 5 min at 37°C. The pellet was re-suspended in 1.5 ml of 285 286 pre-warmed MW supplemented with glucose, and 287 the concentration was determined by counting on a Thoma chamber under a microscope (Olympus, IMT-288 289 2, Paris, France). Spermatozoa were then diluted at 290 10×10^6 /ml in pre-warmed MW supplemented with 5.5 mM glucose, 25 mM NaHCO₃ and 7 mg/ml 291 BSA, pH 7.25 (capacitating MW) (McPartlin et al., 292 2009). Spermatozoa were incubated in 500 µl aliquots 293 294 in polyvinyl alcohol-coated 5 ml round-bottom tubes 295 at 37°C in a humidified atmosphere during 6 296 h. The motility was visually evaluated under a microscope (Olympus, IMT-2, Paris, France) at the 297 298 beginning and at the end of the incubation period. Spermatozoa were then diluted at 1×10^6 /ml in 299 300 capacitating MW supplemented with 5 mM procaine 301 to induce hyperactivated motility. Droplets of 100 µl of spermatozoa suspension were laid down onto culture 302 303 dishes and covered with mineral oil.

Equine oocytes were washed in capacitating MW 304 and groups of 10 were transferred to droplets of 305 100 µl of spermatozoa suspension and co-incubated for 306 18 h in an atmosphere of 5% CO₂ in air at 38.5°C in 307 100% humidity. 308

Control of parthenogenetic activation

Equine oocytes were washed in capacitating MW 310 and transferred to droplets of 100 µl of capacitating 311 MW supplemented with 5 mM procaine without 312 spermatozoa and incubated for 18 h in an atmosphere 313 of 5% CO₂ in air at 38.5°C in 100% humidity. 314

In vitro *culture of equine zygotes*

After 18 h co-incubation with spermatozoa, equine zy-316 gotes were washed in the culture medium and flushed 317 to remove attached spermatozoa. Groups of 10 were 318 transferred to droplets of 30 µl of culture medium for 319 30 h or 54 h (48 h or 72 h post IVF) in an atmosphere of 320 5% CO₂, 5% O₂ and 90% N₂ at 38.5°C in 100% humidity. 321

Four culture media were tested. Synthetic OF 322 (SOF) medium was SOF (Minitub) supplemented 323 with 2% Basal Medium Eagle amino acids solution, 324 1% Minimum Essential Medium non-essential amino 325 acids solution, 0.33 mg/ml sodium pyruvate, 6 mg/ml 326 BSA fatty acid free, 5% FCS and 25 µg/ml genta-327 mycin. DMEM-F12-51445C medium was Dulbecco's 328 Modified Eagle's Medium Ham's Nutrient Mixture 329 F12 with 3151 mg/l dextrose, 2.5 mM L-glutamine, 15 330 mM HEPES, 55 mg/l sodium pyruvate (ref. 51445C) 331 supplemented with 10% FCS and 25 µg/ml gentamy-332 cin. DMEM-F12-D8437 medium was DMEM-F12 with 333 2.5 mM L-glutamine, 15 mM HEPES, 1.2 g/l sodium 334 bicarbonate (ref D8437) supplemented with 10% 335 FCS and 25 µg/ml gentamycin. DMEM-F12-D8900 336 medium was DMEM-F12 with 2.5 mM L-glutamine 337 and 15 mM HEPES (ref D8900) supplemented with 338 1.2 g/l NaHCO₃, 10% FCS and 25 μ g/ml gentamycin. 339 The three references of DMEM-F12 had identical 340 composition but different forms (liquid or powder, 341 sodium bicarbonate included or added subsequently). 342

Assessment of nuclear status

Nuclear status was assessed either after 27 h IVM, 344 after 18 h IVF or after 30 h or 54 h in vitro development. 345 Oocytes and zygotes were washed by aspiration in and 346 out of a pipette in PBS, fixed in 4% paraformaldehyde 347 in PBS for 20 min at room temperature, washed in 348 PBS and processed for DNA and nuclear membrane 349 staining. They were incubated for 30 min at room 350 temperature in 0.2% Triton X-100 in PBS. Non-specific 351 reactions were blocked by incubation for 1 h at room 352 temperature in 10% goat serum in PBS. Oocytes and 353 zygotes were incubated overnight at 4°C or 2 h at 354 room temperature with an anti-lamin A/C antibody 355 (ThermoScientific) diluted 1:100 in PBS containing 356

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357 0.2% BSA and 0.1% Tween. After four washings for 358 5 min in PBS containing 0.2% BSA and 0.1% Tween, 359 they were incubated for 1 h at room temperature with an AlexaFluor 594-conjugated-anti-mouse antibody 360 (Life Technologies) diluted 1:400 in PBS. They were 361 362 then washed five times for 5 min in PBS containing 363 0.1% Tween and two times for 5 min in PBS. They 364 were incubated with 1 µg/ml bis-benzimide (Hoechst 365 33342) in PBS for 5 min and mounted on microscope slides in Mowiol V4-88 (133 mg/ml; Hoechst, Frank-366 367 furt, Germany) and *n*-propyl gallate (5 mg/ml). The slides were kept at 4°C in the dark until observation. 368 369 Oocytes were observed under an epifluorescence 370 microscope (Zeiss). Controls were performed using 371 no primary antibodies to ascertain the absence of non-specific binding or no secondary antibodies to 372 373 ascertain the absence of auto-fluorescence.

374 Intracytoplasmic sperm injection (ICSI) procedure

The ICSI procedure was performed in Italy, using
oocytes collected from slaughtered mares and matured *in vitro* with the procedure described above.

378 *Preparation of equine sperm and ICSI procedure*

379 Fresh semen samples from three mature stallions with a reproductive history of normal fertility were used. 380 381 The stallions were located in the reproductive centre Pegasus (Veterinary Clinics and Animal Productions 382 383 Unit - DETO, Polo di Valenzano, University of Bari Aldo Moro, Valenzano, Bari, Italy) and were 384 385 routinely used in artificial insemination programs. 386 Semen was collected with a Missouri artificial vagina 387 with an in-line gel filter, and was extended with 388 INRA 96 (IMV Technologies, Piacenza, Italy) at a concentration of 20 to 25×10^6 sperm cells/ml and 389 390 used immediately. Sperm cells for ICSI were prepared by the swim-up procedure in Earle's balanced salt 391 392 solution supplemented with 0.4% BSA and 50 µg/ml 393 gentamicin as previously described (Dell'Aquila et al., 394 2001, 2003; Ambruosi *et al.*, 2009).

Intracytoplasmic sperm injection was carried out as
previously reported (Ambruosi *et al.*, 2009; Dell'Aquila *et al.*, 2001, 2003). All procedures were performed
at 38.5°C in Quinn's Advantage Fertilization HTF
Universal medium (Cooper Surgical, Trumbull, CT,
USA) supplemented with 5 mg/ml human serum
albumin (HSA).

402 In vitro *culture of embryos*

In a first embryo culture experiment, aimed to test
the influence of IVM medium (MM versus pFF)
on fertilization and early embryonic development,
injected oocytes were put immediately after ICSI in
one of the two embryo culture medium (either DMEMF12–51445C or DMEM-F12-D8900 supplemented with

1.2 g/l NaHCO₃) plus 10% FCS with 25 µg/ml 409 gentamycin under mineral oil in droplets of 10 µl 410 and cultured individually for 72 h in a humidified 411 atmosphere of 5% CO₂, 5% O₂ and 90% N₂ at 38.5°C. 412 At the end of the culture period, early embryos 413 and uncleaved ova were removed from culture, fixed 414 and evaluated using the procedures as described 415 below. In a second embryo culture experiment, 416 aimed to test the influence of pre-incubation with 417 OF on embryo development, injected oocytes were 418 cultured in DMEM-F12 D8900 for up to 10 days. 419 At day 10 of culture, embryos (developed, delayed 420 and degenerated embryos and uncleaved ova) were 421 morphologically evaluated, fixed and stained for 422 assessing nuclear chromatin, as described below. 423

Nuclear chromatin evaluation

Nuclear chromatin configuration of embryos and un-425 cleaved ova was evaluated under an epifluorescence 426 microscope (Nikon Eclipse 600 equipped with B-2 427 A, 346 nm excitation/460 nm emission filter) after 428 staining with Hoechst 33258, as previously described 429 (Dell'Aquila et al., 2001; Hinrichs et al., 2005; Lange 430 Consiglio et al., 2009). Normally cleaved embryos 431 were defined by the presence of nuclei of regular 432 morphology within each blastomere. The number of 433 morphologically normal nuclei was counted for each 434 embryo. Embryos were classified as morulae when 435 they had more than 32 nuclei. Embryos with more 436 than 64 nuclei and having an outer layer of apparent 437 differentiating trophoblast cells were considered to be 438 blastocysts (Choi et al., 2006). In the group of uncleaved 439 ova, normal fertilization was defined by the presence 440 of two polar bodies with two pronuclei (PN). Oocytes 441 showing one PN with intact sperm cell were classified 442 as activated oocytes. Oocytes showing a metaphase 443 plate and one polar body with an intact sperm 444 cell were classified as unfertilized; oocytes having 445 degenerated, irregularly clustered or faint chromatin 446 were classified as degenerating. 447

Statistical analysis

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The percentages of mature oocytes, fertilized oocytes,
cleaved embryos, morulae and blastocysts were
compared between groups using chi-square analysis.449
450Differences were considered statistically significant at
P < 0.05.451

Results

Maturation rates after IVM

After 27 h IVM in MM or pFF, nuclear status of the
oocytes was assessed. Oocytes with metaphase II and
one polar body were considered mature (Fig. 1a).456457458

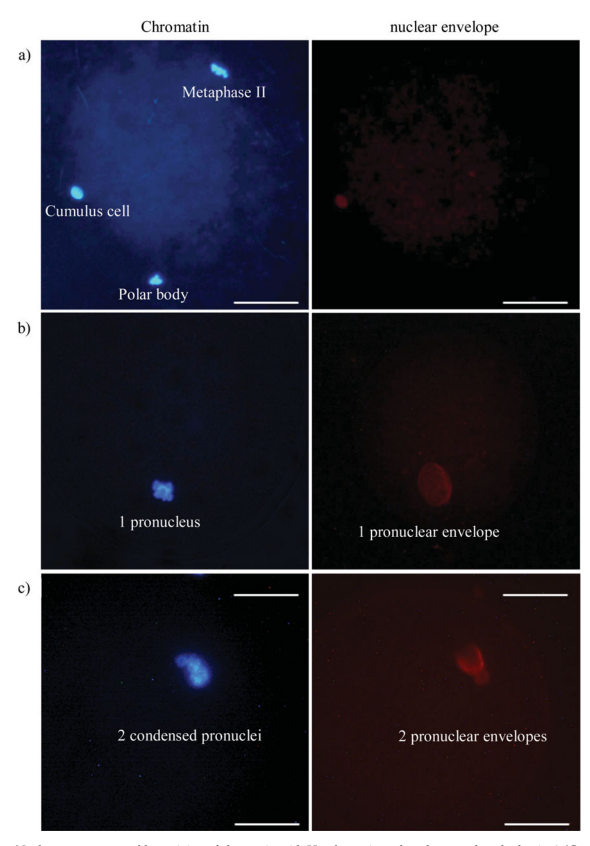


Figure 1 Nuclear status assessed by staining of chromatin with Hoechst stain and nuclear envelope by lamin A/C antibody. (*a*) Mature oocyte with metaphase II and one polar body with no detectable nuclear envelope. (*b*) Oocyte with one pronucleus showing chromatin and a nuclear envelope without sperm cell. (*c*) Oocyte with two condensed pronuclei with a compact mass of chromatin within the nuclear envelope. Scale bar represent 60 μ m.

459 For the oocytes collected in France, three repetitions 460 were performed. After 27 h IVM in MM and pFF, 16 oocytes out of 21 (76%) and 14 oocytes out of 21 (67%) 461 462 were mature. For the oocytes collected in Italy, seven repetitions were performed and in total, 219 oocytes 463 were analysed after IVM. After 27 h IVM in MM and 464 465 pFF, 57 oocytes out of 108 (53%) and 60 oocytes out of 111 (54%) were mature. In both experiments, the 466 467 maturation rates in MM and pFF were not statistically different (P > 0.05). 468

Fertilization and development rates after IVM, IVF and *in vitro* culture

471 Influence of IVM medium on fertilization rates after IVF

472 After oocytes collection from slaughterhouse or OPU, 473 IVM in MM or pFF, post-maturation with OF and 474 gametes co-incubation, nuclear status was assessed. 475 We observed oocytes with metaphase II and one polar body with no detectable nuclear envelope (Fig. 1a), 476 477 oocytes with one pronucleus (PN) showing DNA 478 stained with Hoechst stain and a nuclear envelope 479 stained positive with lamin A/C antibody without sperm cell (Fig. 1b) and oocytes with two PN in 480 481 the cytoplasm, each PN showing the presence of DNA and a nuclear envelope (Fig. 1c). All the 2PN 482 483 oocytes showed condensed PN with a compact mass 484 of chromatin within the nuclear envelope and very 485 few 2PN oocytes showed one or two polar bodies. Degenerated oocytes, having shrunken, dense or 486 fragmented cytoplasm, were discarded. 487

488 For the oocytes collected in a slaughterhouse, three 489 repetitions were performed (Fig. 2a). The percentage of oocytes containing 2PN 18 h post IVF was not 490 491 statistically different between the two maturation 492 conditions (IVM in MM: 22/33, 67%; IVM in pFF: 24/42, 57%; P > 0.05). For the oocytes collected by 493 494 OPU, two repetitions were performed (Fig. 2a). The 495 percentage of oocytes containing 2PN 18 h post IVF was not statistically different (IVM in MM: 4/10, 496 40%; IVM in pFF: 4/9, 44%; *P* > 0.05). When oocytes 497 collected in a slaughterhouse and by OPU were 498 499 pooled, the 2PN rates from MM (26/43, 60%) and pFF 500 (28/51, 55%) were not statistically different (P > 0.05).

Influence of post-maturation with OF on fertilization ratesafter IVF

503 After oocytes collection from slaughtered mares, IVM 504 in MM, pre-incubation or not with OF and gametes 505 co-incubation, nuclear status was assessed. Two repetitions were performed (Fig. 2b). The percentage 506 507 of oocytes containing 2PN 18 h post IVF was not statistically different between post-maturation with 508 OF (19/31, 61%) and without OF (17/34, 50%) (P >509 510 0.05).

Control of parthenogenetic activation

For the control groups for parthenogenetic activation, 512 three repetitions were performed during the time of 513 the experiments. Equine immature oocytes collected 514 in France from slaughtered mares were in vitro 515 matured in MM or pFF, incubated with OF and co-516 incubated with or without spermatozoa. Three oocytes 517 containing 2 PN were observed in the control group 518 incubated without spermatozoa (3/26, 11% and 0/7 519 after IVM in MM and pFF respectively; Fig. 2c). In 520 the other oocytes incubated without spermatozoa we 521 observed either a metaphase II (62%, 16/26 and 57%, 522 4/7) or 1 PN (27%, 7/26 and 43%, 3/7) after IVM 523 in MM and pFF respectively. The percentage of 2PN 524 oocytes was significantly different for incubation with 525 vs without spermatozoa after IVM in MM and pFF (P 526 < 0.05). 527

Development rates after IVM, IVF and in vitro culture After oocytes collection from slaughtered mares or by OPU, IVM in MM or pFF, post-maturation with OF, IVF and *in vitro* culture in one of the four tested media, nuclear status was assessed after 30 h *in vitro* culture, i.e. 48 h post IVF, or after 54 h *in vitro* culture, i.e. 72 h post IVF. For each medium, three repetitions were performed.

In total, 168 oocytes/embryos were analysed 48 h 536 post IVF (39 from SOF, 45 from DMEM-F12-51445C, 537 42 from DMEM-F12-D8437 and 42 from DMEM-538 F12-D8900). In 60 of them (36%), we observed a 539 metaphase II and a polar body or one pronucleus. In 540 89 of them (53%), we observed two fully decondensed 541 pronuclei: the chromatin was diffuse, some nucleoli 542 were observed and the filamentous chromatin filled 543 the whole nuclear envelope (Fig. 3a, c-e). Very few 2PN 544 oocytes showed polar bodies. Moreover, pronuclear 545 apposition was the dominant feature of these 89 546 oocytes (Fig. 3a, c-e). In 19 of the oocytes/embryos 547 (11%), we observed several nuclei (from three to 16) 548 but the quality of these embryonic structures was 549 poor as the cleavage was abnormal: the number 550 of cells and nuclei were different and some cells 551 had no or several nuclei (Fig. 3b, f). The percentage 552 of 2PN oocytes and abnormally cleaved embryos 553 with respect to the number of non-degenerated 554 oocytes for each culture medium is presented 555 in Fig. 4. 556

After culture 48 h post IVF in SOF medium (Figs 3a 557 and 4a), the percentage of 2PN oocytes and abnormal 558 embryos from slaughtered mares was not statistically 559 different between the two maturation media (73% 560 for MM and 67% for pFF; P > 0.05). One embryo 561 (from pFF) contained eight nuclei but no cleavage 562 of the cytoplasm was observed. The percentage of 563 2PN oocytes and abnormal embryos from OPU was 564 not statistically different between the two maturation 565

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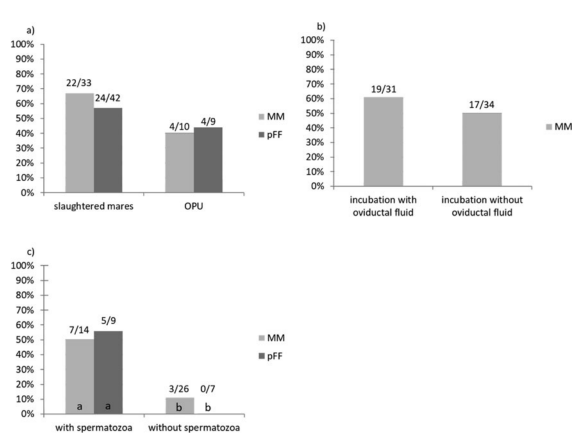


Figure 2 (*a*) Percentage of oocytes containing two pronuclei with respect to the number of non-degenerated oocytes after collection from slaughtered mares or by ovum pick up (OPU), *in vitro* maturation in maturation medium (MM) or preovulatory follicular fluid (pFF), incubation in oviductal fluid and *in vitro* fertilization. The percentages were not statistically different between the two maturation media (P > 0.05). (*b*) Percentage of oocytes containing two pronuclei with respect to the number of non-degenerated oocytes for the oocytes collected from slaughtered mares and incubated with oviductal fluid or not. The percentages were not statistically different between the two conditions (P > 0.05). (*c*) Percentage of oocytes containing two pronuclei with respect to the number of non-degenerated oocytes for the oocytes containing two pronuclei with respect to the number of non-degenerated oocytes for the oocytes containing two pronuclei with respect to the number of non-degenerated oocytes for the oocytes containing two pronuclei with respect to the number of non-degenerated oocytes for the oocytes collected from slaughtered mares and incubated with or without spermatozoa (parthenogenetic controls). The percentages were statistically different between the two conditions: with and without spermatozoa, within a maturation medium. ^{a,b} P < 0.05).

566 media (60% for MM and 75% for pFF, P > 0.05). Six embryos (three from MM and three from pFF) 567 contained from five to 16 nuclei, but the cleavage 568 stopped at two to three cells (Fig. 3b). When oocytes 569 collected in a slaughterhouse and by OPU were 570 571 pooled, the percentages of 2PN oocytes and embryos from MM (69%) and pFF (70%) were not statistically 572 different (P > 0.05). 573

574 After culture in DMEM-F12-51445C medium (Figs 3c and 4b), the percentage of 2PN oocytes and 575 abnormal embryos from slaughtered mares was not 576 statistically different between the two maturation 577 578 media (50% for MM and 71% for pFF, P > 0.05). Three 579 embryos (one from MM and two from pFF) contained from seven to 16 nuclei, however they did not cleave. 580 The percentage of 2PN oocytes and abnormal embryos 581 582 from OPU was not statistically different between the two maturation media (50% for MM and 71% for pFF, 583 584 P > 0.05). No cleavage was observed. When oocytes from slaughterhouse and OPU were pooled, the
percentages of 2PN oocytes and abnormal embryos
from MM (50%) and pFF (71%) were not statistically
different (P > 0.05).585
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After culture in DMEM-F12-D8437 medium (Figs 3d 589 and 4*c*), the percentage of 2PN oocytes and abnormal 590 embryos from slaughtered mares was not statistically 591 different between the two maturation media (75% 592 for MM and 71% for pFF, P > 0.05). One embryo 593 (from MM) contained 12 PN, but no cleavage of 594 the cytoplasm was observed. The percentage of 2PN 595 oocytes and abnormal embryos from OPU was not 596 statistically different between the two maturation 597 media (80% for MM and 75% for pFF, *P* > 0.05, Fig. 4c). 598 No cleavage was observed. When oocytes collected 599 in a slaughterhouse and by OPU were pooled, the 600 percentages of 2PN oocytes and abnormal embryos 601 from MM (76%) and pFF (71%) were not statistically 602 different (P > 0.05). 603

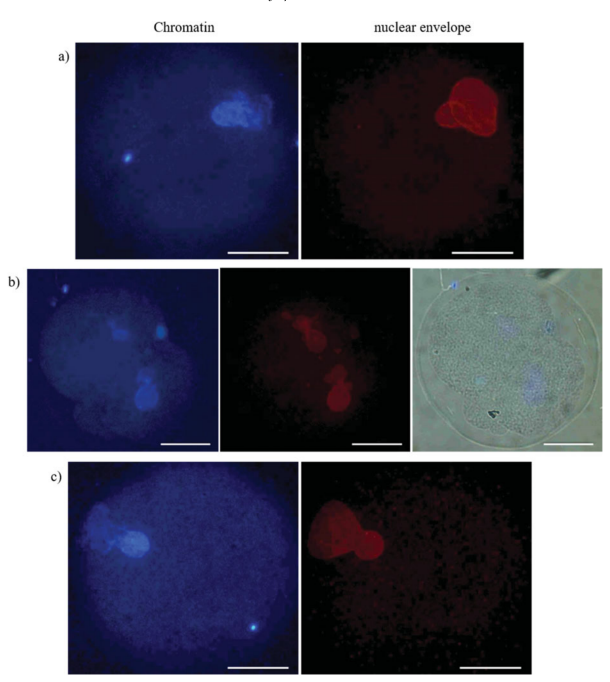


Figure 3 Nuclear status assessed by staining of chromatin with Hoechst and nuclear envelope by lamin A/C antibody. (*a*) An oocyte with two pronuclei fully decondensed after *in vitro* culture in SOF medium. (*b*) An abnormally cleaved embryo with two cells and four nuclei after *in vitro* culture in SOF medium. (*c*) An oocyte with two pronuclei fully decondensed after *in vitro* culture in DMEM-F12–51445C. (*d*) An oocyte with two pronuclei fully decondensed after *in vitro* culture in DMEM-F12-D8437. (*e*) An oocyte with two pronuclei fully decondensed after *in vitro* culture in DMEM-F12-D8430. (*f*) An abnormally cleaved embryo with six cells and five nuclei after *in vitro* culture in DMEM-F12-D8900. Scale bar represent 60 µm.

After culture in DMEM-F12-D8900 medium (Figs 3*e* and 4*d*), the percentage of 2PN oocytes and abnormal embryos from slaughtered mares was not statistically different between the two maturation media (40% for MM and 57% for pFF, P > 0.05). Six embryos (two from MM and four from pFF) contained from 3 to 10 PN but the cleavage stopped at three to six cells (Fig. 3f). The percentage of 2PN oocytes and abnormal embryos611from OPU was not statistically different between the612two maturation media (56% for MM and 67% for pFF,613P > 0.05). Two embryos (from MM) contained three614and nine nuclei, but the cleavage stopped at three615to four cells with fragmentations. When oocytes from616slaughterhouse and OPU were pooled, the percentages617

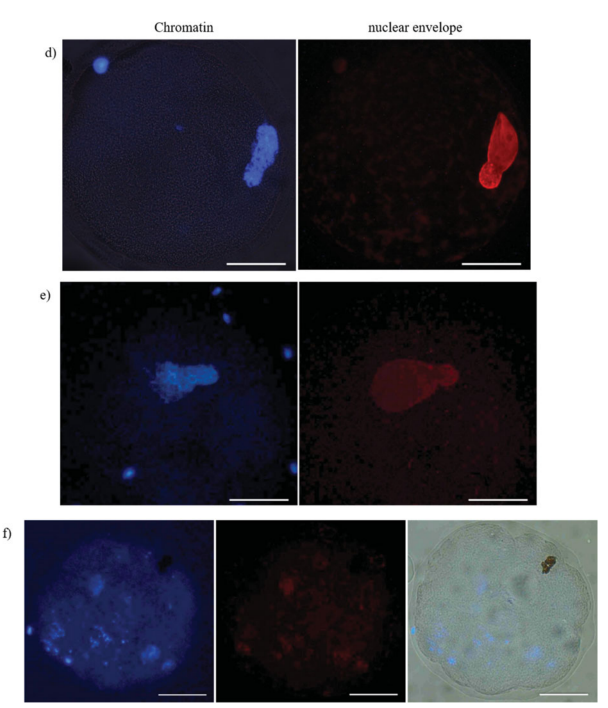


Figure 3 Continued.

of 2PN oocytes and abnormal embryos from MM (47%) 618 and pFF (61%) were not statistically different (P >619 620 0.05).

Finally, when data from the four embryo culture 621 media were pooled, the percentages of 2PN oocytes 622 and abnormal embryos from MM (48/80, 60%) and 623 pFF (60/88, 68%) were not statistically different (P >624 0.05). Moreover, the percentages of abnormally cleaved 625 626 embryos from MM (9/48, 19%) and pFF (10/60, 17%) were not statistically different (P > 0.05). As the four 627 embryo culture media were not tested at the same time, no statistical comparison of these media was performed.

In total, 121 oocytes/embryos were analysed 72 h 631 post IVF (40 from SOF, 26 from DMEM-F12-51445C, 632 37 from DMEM-F12-D8437 and 18 from DMEM-F12-633 D8900). None of them went further in its development 634 during these additional 24 h of culture, and most of 635

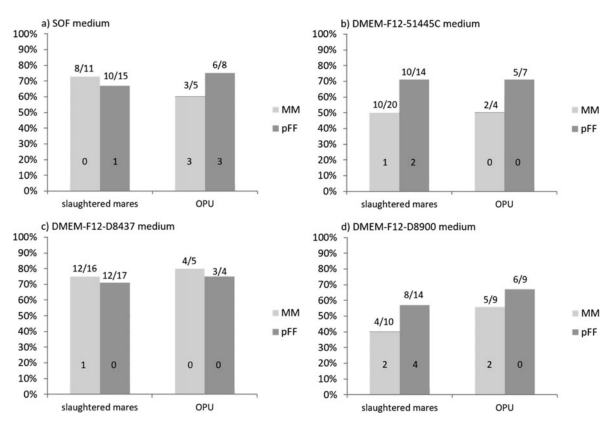


Figure 4 Percentage of 2PN oocytes and abnormally cleaved embryos after collection from slaughtered mares or by ovum pick up (OPU), *in vitro* maturation in maturation medium (MM) or preovulatory follicular fluid (pFF), incubation in oviductal fluid, *in vitro* fertilization and *in vitro* culture 48 h post IVF in (*a*) SOF medium; (*b*) DMEM-F12–51445C medium; (*c*) DMEM-F12-D8437 medium; and (*d*) DMEM-F12-D8900 medium. The fractions at the top represent the number of 2PN oocytes and embryos out of the number of non-degenerated oocytes. The numbers at the bottom represent the number of abnormally cleaved embryos. The percentages were not statistically different between MM and pFF (P > 0.05).

- 636 them degenerated (88% from SOF, 88% from DMEM-
- 637 F12–51445C, 41% from DMEM-F12-D8437 and 100%
- 638 from DMEM-F12-D8900).

639 Fertilization and development rates after IVM, ICSI 640 and *in vitro* culture

641 Influence of IVM medium on fertilization and development642 rates 72 h after ICSI

643 After oocytes collection from slaughtered mares, IVM in MM or pFF, ICSI and in vitro culture for 72 h, nuclear 644 status was assessed. In total, 117 oocytes/embryos 645 were analysed. Among these, 58 (50%) were matured 646 but not fertilized and 59 (50%) were fertilized. Twenty-647 648 five (21%) oocytes remained uncleaved after ICSI and 649 72 h in vitro culture. They showed two polar bodies (PB) and two pronuclei (PN) and were called zygotes. 650

651 In samples cultured in DMEM-F12–51445C, the 652 total fertilization rate (including zygotes and cleaved 653 embryos) did not differ significantly between the 654 two IVM media (18/36, 50% versus 18/38, 47%, 655 for MM and pFF, respectively; P > 0.05, Fig. 5a). 656 In samples cultured in DMEM-F12-D8900, the total fertilization rate was higher in oocytes matured in 657 pFF than in MM (15/22, 68% versus 8/21, 38%; P <658 0.05, Fig. 5b). Independent of culture conditions, all 659 zygotes either derived from in vitro culture in DMEM-660 F12-51445C (Fig. 6a, b) or in DMEM-F12-D8900 (data 661 not shown) showed decondensed pronuclei, their 662 chromatin was diffuse, some nucleoli were observed 663 and the filamentous chromatin filled the whole nuclear 664 area. Pronuclear apposition was found in 37% (7/19) of 665 samples derived from in vitro culture in DMEM-F12-666 51445C (2/9 after IVM in MM and 5/10 after IVM in 667 pFF) whereas it was never found in zygotes derived 668 from *in vitro* culture in DMEM-F12-D8900. 669

The total cleavage rates with respect to the number 670 of metaphase II (MII) injected oocytes for each IVM 671 and culture medium are presented in Fig. 5. In both 672 embryo culture media, the cleavage rates did not 673 differ between maturation conditions: in DMEM-F12-674 51445C, the total cleavage rates were 9/36 (25%) versus 675 8/38 (21%) for MM and pFF respectively (P > 0.05); in 676 DMEM-F12-D8900, the total cleavage rates were 7/21 677 (33%) versus 10/22 (45%) for MM and pFF respectively 678 (P > 0.05).679

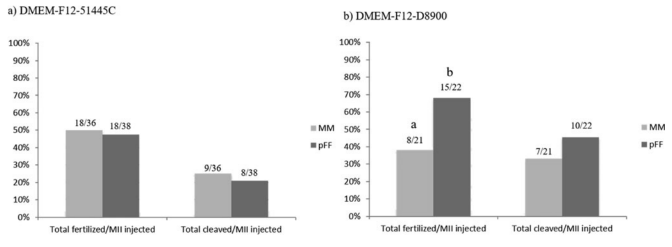


Figure 5 Percentage of fertilized oocytes (with respect to the number of mature oocytes) and cleaved embryos (with respect to the number of mature oocytes) after collection from slaughtered mares, *in vitro* maturation in maturation medium (MM) or preovulatory follicular fluid (pFF), *in vitro* fertilization by intracytoplasmic sperm injection (ICSI) and *in vitro* culture 72 h post-ICSI in (*a*) DMEM-F12–51445C medium; or (*b*) DMEM-F12-D8900 medium. The fractions at the top represent the number of fertilized oocytes out of the number of mature oocytes and the number of cleaved embryos out of the number of mature oocytes. Chi-squared test between maturation media: ${}^{a,b}P < 0.05$.

Representative micrographs of equine early embryos obtained after IVM in MM or in pFF, ICSI
and 72 h *in vitro* embryo culture are presented in
Fig. 6. Independent of culture conditions, most of
the embryos, either derived from *in vitro* culture
in DMEM-F12–51445C (Fig. 6e–h) or in DMEM-F12D8900, showed nuclei of regular morphology.

The percentages of zygotes and embryos at different 687 688 developmental stages observed after IVM in MM or pFF, IVF by ICSI and in vitro culture in DMEM-689 F12-51445C or in DMEM-F12-D8900 are reported in 690 691 Table 1. Within each developmental stage (2PB2PN, 692 2-cell, 4-cell and 8/16-cell stage) no significant differences were found between the two IVM media. 693 694 As the two embryo culture media were not tested at the same time, no statistical comparison of these two 695 696 embryo culture media was performed.

By pooling data of embryo culture media (DMEMF12–51445C + D8900 in Table 1), no significant
differences were found between oocytes matured in
MM or pFF.

Influence of post-maturation with OF on fertilization and development rates 10 days after ICSI

703 After oocytes collection from slaughtered mares, IVM 704 in MM, post-maturation or not with OF, ICSI and in 705 vitro culture, cleavage rates were assessed at day 3 and nuclear status was assessed at day 10 post-ICSI. Data 706 are presented in Table 2. The percentages of cleaved 707 708 embryos at day 3 were not significantly different 709 between oocytes incubated with OF (62%, 29 cleaved embryos/47 injected oocytes) or not (65%, 28/43) 710 711 (P > 0.05). The percentages of morula/MII injected oocytes at day 10 were not significantly different 712 for oocytes incubated with OF (4%, 2/47) compared 713 with oocytes incubated without (0%, 0/43) (P > 0.05). 714 The percentages of blastocyst/MII injected oocytes 715 at day 10 were not significantly different between 716 oocytes incubated with OF (6%, 3/47) or without (7%, 3/47)717 3/43 (P > 0.05). Representative micrographs of equine 718 blastocysts obtained after IVM in MM, incubation or 719 not in OF, ICSI and 10 days in vitro embryo culture in 720 DMEM-F12-D8900 are presented in Fig. 7. They were 721 observed either in culture (Fig. 7a, b) or after fixation 722 and Hoechst staining (Fig. 7c, d). In this experiment, 723 all blastocysts reached a good expansion level with a 724 consistent number of nuclei of regular morphology. 725

Discussion

The aim of our work was to analyse the effect of 727 IVM in follicular fluid and pre-incubation in OF on 728 IVM, fertilization using conventional IVF or ICSI, and 729 development of equine oocytes. We have been faced 730 with the difficulty with equine oocyte collection. As 731 the number of slaughtered mares is very low and 732 the collection rate of equine oocytes is low compared 733 with bovine oocytes for example (Hawley et al., 1995), 734 the number of oocytes from slaughterhouses available 735 for our experiments was low. Moreover, the collection 736 of equine oocytes using OPU is expensive and time-737 consuming, thus a maximum of 4 to 5 OPU sessions 738 per morning were performed, and the number of 739 oocytes collected by OPU was low. The authors are 740 aware of the small sample size in their experiments 741

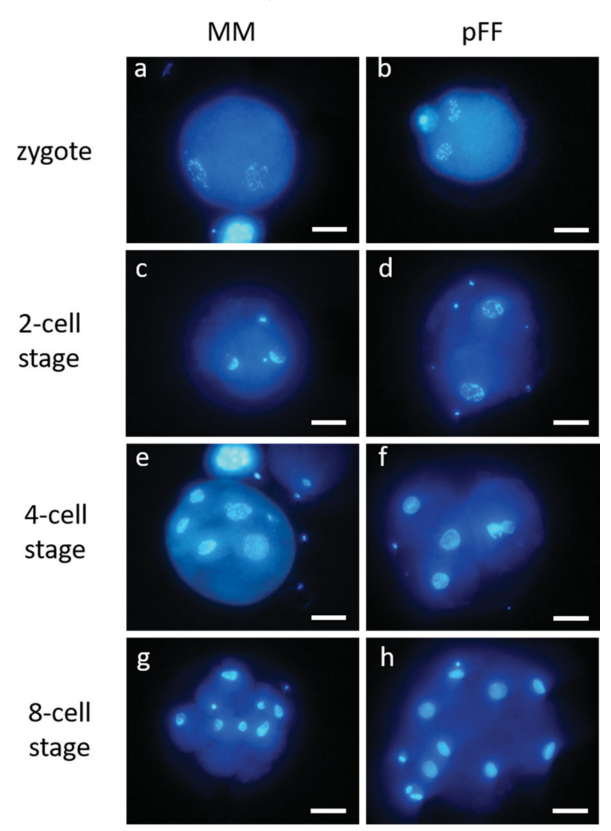


Figure 6 Representative photomicrographs of equine early embryos obtained after IVM in maturation medium (MM) or preovulatory follicular fluid (pFF), fertilized by ICSI and *in vitro* cultured in DMEM-F12–51445C for 72 h. Nuclear status assessed by chromatin staining with Hoechst and examined under ultraviolet (UV) light. (*a*, *b*) Zygotes showing two decondensed pronuclei. (*c*, *d*) Two-cell stage embryo. (*e*, *f*) Four-cell stage embryo. (*g*, *h*) Eight-cell stage embryo. Scale bar represent 60 μ m.

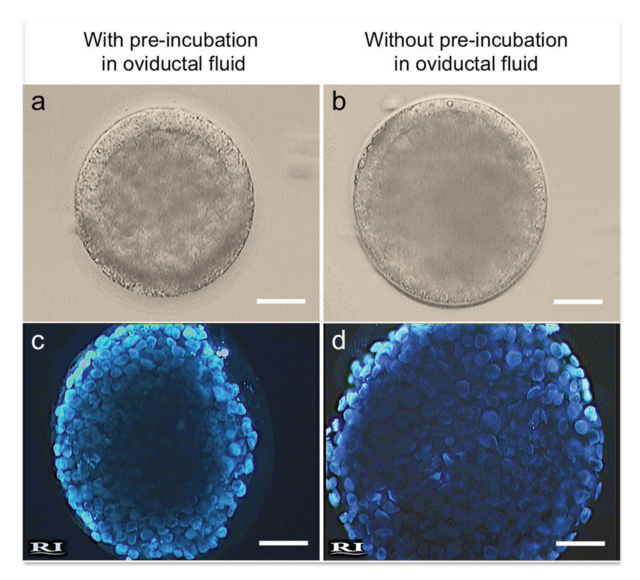


Figure 7 Representative photomicrographs of equine blastocysts obtained after IVM in maturation medium (MM), incubation or not in oviductal fluid and *in vitro* culture in DMEM-F12-D8900 for 10 days and observed under phase contrast (a, b) or UV light (c, d) after Hoechst staining. Two expanded blastocysts with the outer layer of trophoblastic cells and a high number of nuclei are shown. Scale bar represent 60 μ m.

due to the difficulty in collecting equine oocytes, butthe statistical tests have been adapted to small samplesize and our conclusions are relevant.

Our first objective was to compare IVM in MM 745 or preovulatory follicular fluid. Our hypothesis was 746 that the low IVF and development rates observed 747 in the equine may be due, at least partly, to a 748 749 poor cytoplasmic maturation related to unsuited IVM 750 conditions. Preovulatory follicular fluid, in which 751 maturation naturally occurs, may sustain a better cytoplasmic maturation and thus a better competence 752 753 for IVF and development. In our study, the nuclear maturation rates in MM and pFF were not statistically 754 755 different, neither for the oocytes collected in France (76% of the oocytes are in metaphase II in MM 756

versus 67% in pFF) nor for the oocytes collected 757 in Italy (53% in MM versus 54% in pFF). Thus, 758 the semi-synthetic MM provides nuclear maturation 759 rates similar to follicular fluid. In previous studies, 760 the maturation rates of equine oocytes were similar: 761 68% in pFF (Caillaud et al., 2008), 54-58% in pFF 762 (Conforti et al., 2005), 61% in MM (Ambruosi et al., 763 2013), 71% in MM (Deleuze et al., 2009). In our 764 study, the fertilization rates after ICSI were not 765 different between maturation in MM (46%) vs pFF 766 (55%). This result shows that the semi-synthetic MM 767 supports cytoplasmic maturation to a similar extent to 768 preovulatory follicular fluid. Moreover, the percentage 769 of oocytes containing two pronuclei after IVF was 770 not statistically different between maturation in MM 771

Table 1 Percentage of equine zygotes (oocytes showing two polar bodies and two pronuclei, 2PB2PN) and embryos at different developmental stages after oocyte collection from slaughtered mares, IVM in maturation medium (MM) or preovulatory follicular fluid (pFF), intracytoplasmic sperm injection (ICSI) and *in vitro* culture in DMEM-F12–51445C medium or DMEM-F12-D8900 medium for 72 h post-ICSI

13/3.4	Embryo culture medium	N° of cultured oocytes	N° (%) of MII and injected oocytes	N° (%) of total fertilized oocytes ^a	N° (%) of zygotes and cleaved embryos found 72 h post-ICSI b				
IVM medium					2PB2PN	2-cell stage	4-cell stage	8/16-cell stage	Total cleaved
MM	DMEM-F12- 51445C	69	36 (52)	18 (50)	9 (25)	1 (3)	6 (17)	2 (5)	9 (25)
pFF		71	38 (53)	18 (47)	10 (26)	3 (8)	4 (10)	1 (3)	8 (21)
MM + pFF	DMEM-F12-	140	74 (53)	36 (49)	19 (26)	4 (5)	10 (14)	3 (4)	17 (23)
	51445C								
MM	DMEM-F12 D8900	39	21 (54)	8 (38)*	1 (5)	3 (14)	3 (14)	1 (5)	7 (33)
pFF		40	22 (55)	15 (68)*	5 (23)	5 (23)	2 (9)	3 (14)	10 (46)
MM + pFF	DMEM-F12-D8900	79	43 (54)	23 (53)	6 (14)	8 (19)	5 (12)	4 (9)	17 (40)
MM	Total DMEM-F12-	108	57 (53)	26 (46)	10 (17)	4 (7)	9 (16)	3 (5)	16 (28)
	51445C + D8900			. ,	. ,			. /	. ,
pFF		111	60 (54)	33 (55)	15 (25)	8 (13)	6 (10)	4 (7)	18 (30)

^aNumbers and percentages of fertilized oocytes out of the number of MII injected oocytes.

^bNumbers and percentages of zygotes and cleaved embryos out of the number of MII injected oocytes.

Chi-squared test between media, for fertilization and cleavage rates: *P < 0.05.

Table 2 Percentage of equine embryos at morula or blastocyst stages after oocyte collection from slaughtered mares, IVM in maturationmedium (MM), incubation or not with OF, intracytoplasmic sperm injection (ICSI) and *in vitro* culture for 10 days post-ICSI inDMEM-F12-D8900

IVM medium	Pre-incubation with oviductal fluid	No. of cultured oocytes	No. (%) of MII and injected oocytes	No. (%) of morula (day 10) ^a	No. (%) of blastocyst (day 10) ^a	No. (%) of morula + blastocyst (day 10) ^a
MM	Pre-incubation	103	47 (46)	2 (4)	3 (6)	5 (11)
MM	No pre-incubation	109	43 (39)	0 (0)	3 (7)	3 (7)
Total	Total	212	90 (42)	2 (2)	6 (7)	8 (9)

^aNumbers and percentages of embryos out of the number of MII injected oocytes.

772 versus pFF, neither for the oocytes collected in a 773 slaughterhouse (67% in MM versus 57% in pFF) nor 774 for the oocytes collected by OPU (40% in MM versus 44% in pFF). As very few oocytes containing two 775 pronuclei extruded PB, the significance of this nuclear 776 777 stage may be questionable. One could hypothesize that either the PB were extruded and degenerated 778 779 quickly or the expulsion of the PB did not occur 780 as a consequence of an abnormal fertilization. On the one hand, using equine zygotes flushed from 781 oviducts at different time intervals from ovulation and 782 783 in vitro fertilized oocytes, Bézard and collaborators 784 showed that PB degeneration and undetectability may 785 occur quickly after fertilization both in vitro and in vivo (Bézard et al., 1989). On the other hand, 786 Leemans and collaborators showed that the absence 787 of the second polar body extrusion is due to oocyte 788 cytokinesis induced by procaine instead of sperm 789 penetration of equine oocytes (Leemans et al., 2015). In 790 our study, the presence of some equine oocytes con-791 792 taining two pronuclei in the parthenogenetic control 793 group incubated without spermatozoa supports this 794 second hypothesis. Previous studies have shown that 795 spontaneous parthenogenetic division of unfertilized equine oocytes is rare either in vitro or in vivo (Zhang 796 et al., 1989; Zhang et al., 1990). Finally, after culture 797 798 in embryo culture media, we observed no difference 799 between maturation in MM vs pFF for the percentage of zygotes after ICSI (17% vs. 25% respectively) and 800 801 the percentage of cleaved embryos after ICSI (28% 802 versus 30%). Thus, our MM and the preovulatory 803 follicular fluid support the acquisition of competence 804 for maturation, fertilization and development to a similar extent. However, in our conditions, IVM occurs 805 806 in a static medium, whereas in vivo maturation naturally takes place in a dynamic medium in which 807 808 subtle changes occur during final maturation of 809 the preovulatory follicle. IVM in a culture system 810 in which hormones would be added sequentially to mimic changes observed in vivo might provide 811 812 better conditions for equine oocyte maturation. For example, porcine oocytes matured in a medium in 813 814 which hormones were added sequentially exhibited greater developmental competence to blastocyst stage 815 (Kawashima *et al.*, 2008). 816

The influence of oviductal secretions on oocytes 817 has been widely studied in mammals (Aviles et al., 818 819 2010; Coy et al., 2012; Lopera-Vasquez et al., 2015). A 820 beneficial effect on equine oocytes has been observed 821 with incubation of equine oocytes with OF collected 6 h after ovulation (Ambruosi et al., 2013) or co-culture 822 823 of equine oocytes with equine or porcine oviduct epithelial cells (Mugnier et al., 2009). In our study, 824 incubation of equine oocytes with OF collected before 825 826 ovulation, at the end of follicular growth, did not 827 influence fertilization or development rates. This result suggests that oviductal factors with a beneficial effect 828 on oocyte may be present in the oviduct during a 829 specific time interval after ovulation. Several oviductal 830 factors, which influence fertilization rates, have been 831 studied in the equine oviduct: Deleted in malignant 832 brain tumour one is secreted in the oviduct from early 833 follicular phase to post-ovulatory stage (Ambruosi 834 et al., 2013) and osteopontin and atrial natriuretic 835 peptide A are present in the equine oviduct from 836 emergence of the dominant follicle to preovulatory 837 stage (Mugnier et al., 2009). Further studies are in 838 progress to clarify the role and secretion pattern of 839 oviductal factors with a beneficial effect on equine 840 oocvtes. 841

Several culture media have been used for in 842 vitro development of equine fertilized oocytes. SOF 843 medium has been used previously for in vitro culture 844 of equine zygotes after ICSI with a cleavage rate 845 of 61% (Tremoleda et al., 2003), 69% (Galli et al., 846 2002) and 64% (Lazzari et al., 2002) 48 h post-847 ICSI. In our conditions, 26% of the oocytes after 848 IVF were abnormally cleaved and 74% contained 849 two fully decondensed and apposed pronuclei. The 850 pronuclei decondensation and apposition evidence 851 the first step of embryo development. However, 852 the lack of normal embryo cleavage throws doubt 853 on the fertilization technique. The use of procaine 854 during the IVF procedure in order to induce sperm 855 hyperactivation may have a detrimental effect on 856 oocytes and zygotes. Leemans and collaborators have 857 shown that procaine induces oocyte cytokinesis and 858 that cleaved oocytes did not develop beyond 8-16 859 cells with daughter cells containing aberrant DNA 860 fragments (Leemans et al., 2015). DMEM-F12 medium 861 has been used previously for in vitro culture of equine 862 zygotes after ICSI with a cleavage rate of 68% (Smits 863 et al., 2012a), 69% (Choi et al., 2011) and 75% (Choi 864 et al., 2006) 3 days post-ICSI. Moreover, DMEM-F12 865 medium has been used previously for in vitro culture 866 of equine zygotes after IVF with a cleavage rate of 867 66% (McPartlin et al., 2009). In our conditions, three 868 references of DMEM-F12 were tested, with identical 869 composition but different forms (liquid or powder, 870 sodium bicarbonate included or added subsequently). 871 Following ICSI and 72 h culture in DMEM-F12, the 872 percentages of zygotes (26% in DMEM-F12-51445C 873 and 14% in DMEM-F12-D8900) and the percentages 874 of cleaved embryos (23% in DMEM-F12-51445C 875 and 40% in DMEM-F12-D8900) were consistent, and 876 the morphology of the ICSI embryos was normal. 877 Following ICSI and 10 days culture in DMEM-F12-878 D8900, the blastocysts formation rate was correct 879 and repeatable and blastocysts quality was good, as 880 blastocyst expansion and hatching throughout the 881 zona pellucida were observed. Following IVF and 882 culture in DMEM-F12, no normal embryo cleavage and 883 development were observed. As the IVM and embryo
culture media were the same in the IVF and ICSI
experiments, it can be speculated that the IVM and
embryo culture conditions are appropriate and that
the IVF conditions have to be improved to increase
the quality of the embryos and their developmental
competence.

891 Thus, in our study, exposure to physiological fluids for oocyte maturation and post-maturation does not 892 improve in vitro embryo production in the horse. 893 In vitro culture of ICSI fertilized oocytes provides 894 895 morphologically normal embryos. However, when procaine is used during IVF, no embryo able to develop 896 897 properly is observed. Improving the IVF technique 898 will improve the developmental competence of IVF embryos and bring them to a state in which they will 899 900 be transferred into surrogate mares.

901 Author's contributions

MED and GG conceived of the study and participated
in its design. CD, OP, FR, SD and GG performed the
experiments and analysis in France, NAM, MN, GML
and MED performed the experiments and analysis
in Italy. MED and GG wrote the manuscript and all
authors read and approved the final manuscript.

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926 **Conflict of interest**

927 The authors declare no conflict of interest.

References

- Adams, G. P., Ratto, M. H., Collins, C. W. & Bergfelt, D.R.
 (2009). Artificial insemination in South American camelids
 and wild equids. *Theriogenology* 71, 166–75.
 931
- Alm, H., Torner, H., Blottner, S., Nurnberg, G. & Kanitz,
 W. (2001). Effect of sperm cryopreservation and treatment with calcium ionophore or heparin on *in vitro* fertilization of horse oocytes. *Theriogenology* 56, 817–29.
- Ambruosi, B., Accogli, G., Douet, C., Canepa, S., Pascal, G., Monget, P., Moros, C., Holmskov, U., Mollenhauer, J., Robbe-Masselot, C., Vidal, O., Desantis, S. & Goudet, G. (2013). Deleted in malignant brain tumor 1 is secreted in the oviduct and involved in the mechanism of fertilization in equine and porcine species. *Reproduction* 146, 119–33.
- Ambruosi, B., Lacalandra, G. M., Iorga, A. I., De Santis, T., Mugnier, S., Matarrese, R., Goudet, G. & Dell'aquila, M.E. (2009). Cytoplasmic lipid droplets and mitochondrial distribution in equine oocytes: Implications on oocyte maturation, fertilization and developmental competence after ICSI. *Theriogenology* **71**, 1093–104.
- Aviles, M., Gutierrez-Adan, A. & Coy, P. (2010). Oviductal secretions: will they be key factors for the future ARTs? *Mol. Hum. Reprod.* **16**, 896–906.
- Bézard, J., Magistrini, M., Duchamp, G. & Palmer, E. (1989). Chronology of equine fertilisation and embryonic development in vivo and *in vitro*. *Equine Vet. J. Suppl.* 8, 105–10.
- Caillaud, M., Dell'aquila, M. E., De Santis, T., Nicassio, M., Lacalandra, G. M., Goudet, G. & Gerard, N. (2008). *In vitro* equine oocyte maturation in pure follicular fluid plus interleukin-1 and fertilization following ICSI. *Anim. Reprod. Sci.* **106**, 431–9.
- Carrasco, L. C., Romar, R., Aviles, M., Gadea, J. & Coy, P. (2008). Determination of glycosidase activity in porcine oviductal fluid at the different phases of the estrous cycle. *Reproduction* **136**, 833–42.
- Choi, Y. H., Love, C. C., Varner, D. D. & Hinrichs, K. (2006). Equine blastocyst development after intracytoplasmic injection of sperm subjected to two freeze-thaw cycles. *Theriogenology* **65**, 808–19.
- Choi, Y. H., Varner, D. D., Love, C. C., Hartman, D. L. & Hinrichs, K. (2011). Production of live foals via intracytoplasmic injection of lyophilized sperm and sperm extract in the horse. *Reproduction* **142**, 529–38.
- Conforti, V. A., Vanderwall, D. K. & Woods, G.L. (2005). Effect of homologous follicular fluid from medium-sized and large follicles on *in vitro* maturation of equine cumulus–oocyte complexes. *Reprod. Fertil. Dev.* **17**, 651–8.
- Coy, P., Garcia-Vazquez, F. A., Visconti, P. E. & Aviles, M. (2012). Roles of the oviduct in mammalian fertilization. *Reproduction* **144**, 649–60.
- Deleuze, S., Goudet, G., Caillaud, M., Lahuec, C. & Duchamp, G. (2009). Efficiency of embryonic development after intrafollicular and intraoviductal transfer of *in vitro* and in vivo matured horse oocytes. *Theriogenology* **72**, 203–9.
- Dell'Aquila, M. E., Albrizio, M., Maritato, F., Minoia, P. & 985
 Hinrichs, K. (2003). Meiotic competence of equine oocytes and pronucleus formation after intracytoplasmic sperm 987

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982

988 injection (ICSI) as related to granulosa cell apoptosis. *Biol.*989 *Reprod.* 68, 2065–72.

- Dell'Aquila, M. E., Cho, Y. S., Minoia, P., Traina, V., Fusco, S.,
 Lacalandra, G. M. & Maritato, F. (1997a). Intracytoplasmic
 sperm injection (ICSI) versus conventional IVF on
 abattoir-derived and *in vitro*-matured equine oocytes. *Theriogenology* 47, 1139–56.
- 995 Dell'Aquila, M. E., Cho, Y. S., Minoia, P., Traina, V.,
 996 Lacalandra, G. M. & Maritato, F. (1997b). Effects of
 997 follicular fluid supplementation of in-vitro maturation
 998 medium on the fertilization and development of equine
 999 oocytes after in-vitro fertilization or intracytoplasmic
 1000 sperm injection. *Hum. Reprod.* 12, 2766–72.
- 1001 Dell'Aquila, M. E., Fusco, S., Lacalandra, G. M. & Maritato,
 1002 F. (1996). *In vitro* maturation and fertilization of equine oo1003 cytes recovered during the breeding season. *Theriogenology*1004 **45**, 547–60.
- 1005 Dell'Aquila, M. E., Masterson, M., Maritato, F. & Hinrichs,
 1006 K. (2001). Influence of oocyte collection technique on
 1007 initial chromatin configuration, meiotic competence, and
 1008 male pronucleus formation after intracytoplasmic sperm
 1009 injection (ICSI) of equine oocytes. *Mol. Reprod. Dev.* 60, 79–
 1010 88.
- Galli, C., Crotti, G., Turini, P., Duchi, R., Mari, G., Zavaglia,
 G., Duchamp, G., Daels, P. & Lazzari, G. (2002). Frozenthawed embryos produced by ovum pick up of immature
 oocytes and ICSI are capable to establish pregnancies in
 the horse. *Theriogenology* 58, 705–8.
- 1016 Goudet, G. (2011). Fertilisation in the horse and paracrine 1017 signalling in the oviduct. *Reprod. Fertil. Dev.* **23**, 941–51.
- 1018 Goudet, G., Belin, F., Mlodawska, W. & Bezard, J.
 1019 (2000). Influence of epidermal growth factor on *in vitro*1020 maturation of equine oocytes. *J. Reprod. Fertil. Suppl.* 56,
 1021 483–92.
- Goudet, G., Bézard, J., Duchamp, G., Gérard, N. & Palmer,
 E. (1997). Equine oocyte competence for nuclear and
 cytoplasmic *in vitro* maturation: effect of follicle size and
 hormonal environment. *Biol. Reprod.* 57, 232–45.
- Hawley, L. R., Enders, A. C. & Hinrichs, K. (1995). Comparison of equine and bovine oocyte–cumulus morphology
 within the ovarian follicle. *Biol. Reprod. Monograph Series* 1,
 243–52.
- Hinrichs, K. (2012). Assisted reproduction techniques in thehorse. *Reprod. Fertil. Dev.* 25, 80–93.
- Hinrichs, K., Choi, Y. H., Love, L. B., Varner, D. D., Love,
 C. C. & Walckenaer, B.E. (2005). Chromatin configuration
 within the germinal vesicle of horse oocytes: changes post
 mortem and relationship to meiotic and developmental
 competence. *Biol. Reprod.* 72, 1142–50.
- Hinrichs, K., Love, C. C., Brinsko, S. P., Choi, Y. H. &
 Varner, D.D. (2002). *In vitro* fertilization of *in vitro*-matured
 equine oocytes: effect of maturation medium, duration
 of maturation, and sperm calcium ionophore treatment,
 and comparison with rates of fertilization in vivo after
 oviductal transfer. *Biol. Reprod.* 67, 256–62.
- Kawashima, I., Okazaki, T., Noma, N., Nishibori, M.,
 Yamashita, Y. & Shimada, M. (2008). Sequential exposure
 of porcine cumulus cells to FSH and/or LH is critical
 for appropriate expression of steroidogenic and ovulationrelated genes that impact oocyte maturation in vivo and *in vitro. Reproduction* 136, 9–21.

- Lange Consiglio, A., Dell'Aquila, M. E., Fiandanese, N., Ambruosi, B., Cho, Y. S., Bosi, G., Arrighi, S., Lacalandra,
 G. M. & Cremonesi, F. (2009). Effects of leptin on *in vitro*maturation, fertilization and embryonic cleavage after
 ICSI and early developmental expression of leptin (Ob)
 and leptin receptor (ObR) proteins in the horse. *Reprod. Biol. Endocrinol.* 7, 113.
- Lazzari, G., Crotti, G., Turini, P., Duchi, R., Mari, G., Zavaglia,
 G., Barbacini, S. & Galli, C. (2002). Equine embryos
 at the compacted morula and blastocyst stage can be
 obtained by intracytoplasmic sperm injection (ICSI) of *in vitro* matured oocytes with frozen-thawed spermatozoa
 from semen of different fertilities. *Theriogenology* 58,
 709–12.
 Leemans, B., Gadella, B. M., Stout, T. A., Heras, S., Smits,
- Leemans, B., Gadella, B. M., Stout, T. A., Heras, S., Smits, K., Ferrer-Buitrago, M., Claes, E., Heindryckx, B., De Vos, W. H., Nelis, H., Hoogewijs, M. & Van Soom, A. (2015).
 Procaine induces cytokinesis in horse oocytes via a pH-dependent mechanism. *Biol. Reprod.* 93, 23.

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- Lopera-Vasquez, R., Hamdi, M., Maillo, V., Lloreda, V., Coy,
 P., Gutierrez-Adan, A., Bermejo-Alvarez, P. & Rizos, D.
 (2015). Effect of bovine oviductal fluid on development
 and quality of bovine embryos produced *in vitro*.
 1071 *Reprod. Fertil. Dev.* doi: 10.1071/RD15238. [Epub ahead of
 print]
 Martino, N. A., Marzano, G., Nicassio, M., Minervini,
- Martino, N. A., Marzano, G., Nicassio, M., Minervini, F., Cardinali, A., Lacalandra, G. M., Hinrichs, K. & Dell'aquila, M.E. (2016). Effects of verbascoside treatment during oocyte *in vitro* maturation on blastocyst development and bioenergetic/oxidative status after ICSI in the horse. *J. Equine Vet. Sci.* **41**, 66.
- McPartlin, L. A., Suarez, S. S., Czaya, C. A., Hinrichs, K. & Bedford-Guaus, S.J. (2009). Hyperactivation of stallion sperm is required for successful *in vitro* fertilization of equine oocytes. *Biol. Reprod.* 81, 199–206.
- Meyers-Brown, G., Bidstrup, L. A., Famula, T. R., Colgin, M. & Roser, J.F. (2011). Treatment with recombinant equine follicle stimulating hormone (reFSH) followed by recombinant equine luteinizing hormone (reLH) increases embryo recovery in superovulated mares. *Anim. Reprod. Sci.* **128**, 52–9.
- Mugnier, S., Kervella, M., Douet, C., Canepa, S., Pascal,1090G., Deleuze, S., Duchamp, G., Monget, P. & Goudet, G.1091(2009). The secretions of oviduct epithelial cells increase1092the equine *in vitro* fertilization rate: are osteopontin, atrial1093natriuretic peptide A and oviductin involved? *Reprod. Biol.*1094*Endocrinol.* 7, 129.1095
- Palmer, E., Bezard, J., Magistrini, M. & Duchamp, G. (1991). In vitro fertilization in the horse. A retrospective study. J. Reprod. Fertil. Suppl. 44, 375–84.
- Smits, K., Govaere, J., Hoogewijs, M., Piepers, S. & Van Soom, A. (2012a). A pilot comparison of laser-assisted vs piezo drill ICSI for the *in vitro* production of horse embryos. *Reprod. Domest. Anim.* **47**, e1–3.
- Reprod. Domest. Anim. 47, e1–3.1102Smits, K., Hoogewijs, M., Woelders, H., Daels, P. & Van
Soom, A. (2012b). Breeding or assisted reproduction?1103Relevance of the horse model applied to the conservation
of endangered equids. Reprod. Domest. Anim. 47(Suppl. 4),
239–48.1107
- Tremoleda, J. L., Stout, T. A., Lagutina, I., Lazzari, G., Bevers, 1108
 M. M., Colenbrander, B. & Galli, C.(2003). Effects of *in vitro* 1109

- 1110 production on horse embryo morphology, cytoskeletal 1111 characteristics, and blastocyst capsule formation. *Biol.*
- 1112 Reprod. 69, 1895–906.
- 1113 Zhang, J. J., Boyle, M. S., Allen, W. R. & Galli, C. 1114 (1989). Recent studies on in vivo fertilization of *in*

vitro matured horse oocytes. Equine Vet. J. Suppl. 8, 1115 101–4. 1116

Zhang, J. J., Muzs, L. Z. & Boyle, M.S. (1990). In vitro 1117 fertilization of horse follicular oocytes matured in vitro. 1118 Mol. Reprod. Dev. 26, 361–5. 1119