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12

13 **Stem cells from foetal adnexa and fluid in domestic animals: an update on their features**
14 **and clinical application.**

15

16 **Running Title:** MSCs from foetal adnexa in domestic animals

17

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26

27 **Summary**

28 Over the past decade stem cell research has emerged as an area of major interest for its
29 potential in regenerative medicine applications. This is in constant need of new cell sources to
30 conceive regenerative medicine approaches for diseases that are still without therapy.
31 Scientists drew the attention toward alternative sources such as foetal adnexa and fluid, since
32 these sources possess many advantages: first of all, cells can be extracted from discarded
33 foetal material and it is noninvasive for the patient and inexpensive, secondly abundant stem
34 cells can be obtained and finally, these stem cell sources are free from ethical considerations.
35 Cells derived from foetal adnexa and fluid preserve some of the characteristics of the
36 primitive embryonic layers from which they originate. Many studies have demonstrated the
37 differentiation potential in vitro and in vivo toward mesenchymal and non-mesenchymal cell
38 types; in addition the immune-modulatory properties make these cells a good candidate for
39 allo- and xenotransplantation. Naturally occurring diseases in domestic animals can be more
40 ideal as disease model of human genetic and acquired diseases and could help to define the
41 potential therapeutic use efficiency and safety of stem cells therapies. This review offers an
42 update on the state of the art of characterization of domestic animals MSCs derived from
43 foetal adnexa and fluid and on the latest findings in pre-clinical or clinical setting of the stem
44 cell populations isolated from these sources.

45 **Key words:** Amniotic fluid, Amniotic Membrane, Umbilical cord blood, Wharton's Jelly,
46 Mesenchymal Stem Cells

47

48 **Introduction**

49

50 A stem cell is an undifferentiated cell capable of differentiating into specialized cells and to
51 originate other stem cells. In mammals, there are three types of stem cells (SCs): embryonic
52 stem cells (ESCs), foetal stem cells, and adult stem cells. ESCs derive from the inner cell
53 mass (ICM) of a blastocyst. They are pluripotent but highly tumorigenic after transplantation.
54 Foetal stem cells are distinguished into two types: foetal proper stem cells (from the tissue of
55 the fetus), and extra-embryonic foetal stem cells (from extra-embryonic membranes). Adult
56 SCs are multipotent cells, designate to tissue repairing, and they are present in almost all adult
57 tissues.

58 Mesenchymal stem cells (MSCs) are a population of multipotent stem cells that meet the
59 following criteria: 1) plastic-adherence when maintained in standard culture conditions; 2)
60 expression of CD105, CD73 and CD90, and lack of CD45, CD34, CD14 or CD11b, CD79 α
61 or CD19 and HLA-DR surface molecules; 3) differentiation to osteoblasts, adipocytes and
62 chondroblasts in vitro (Dominici et al. 2006).

63 Since these properties, MSCs offer a great chance for cell-based therapies and tissue
64 engineering applications. The effective management of companion animals, such as dog, cat
65 or horse, requires sophisticated new treatments and preventive strategies. This, combined with
66 their role as models for human diseases, led to an increase in the number of research focused
67 on isolation and clinical use of MSCs in these animals.

68 Bone marrow (BM) is the common source of autologous MSCs for clinical applications in
69 veterinary medicine. Alternatively, adipose tissue-derived MSCs can be used, since they have
70 a higher proliferation potential. Anyway, for both sources, an invasive procedure is required
71 and there is a large variability in the cell yield related to the donor (Colleoni et al. 2009).

72 Placental tissues and foetal fluids represent a source of cells for regenerative medicine, and
73 are readily available and easily procured without invasive procedures. MSCs from foetal

74 fluids and adnexa are defined as an intermediate between ESCs and adult SCs, due to the
75 preservation of some characteristics typical of the primitive native layers. Among foetal
76 adnexal tissues, the major sources of MSCs are amnion, amniotic fluid, umbilical cord and
77 umbilical cord blood. These cells can be obtained in large numbers without risks for donors
78 and cryopreserved for future use in regenerative veterinary medicine.

79 The markers required for defining a MSC were determined for human cells, and the lack of
80 specific antibodies for different animal species lead to a great variability of approaches and
81 results for molecular characterization of animal MSCs. The aim of this review is to provide an
82 overview of the advances in characterization and clinical applications of domestic animal
83 MSCs derived from foetal adnexa and fluid.

84

85 **Developmental biology of foetal adnexa and amniotic fluid**

86

87 Extra-embryonic or foetal membranes are structures that develop from the zygote, do not
88 form part of the embryo itself, and are of functional importance only in embryonic life. At the
89 end of cleavage, the blastocysts, enclosed within the zona pellucida, consists of an ICM and a
90 trophoblastic layer. During the second week of development, hatched blastocyst leaves its
91 confined space and elongates. In particular this process involves the trophoblast, and,
92 partially, the inner endodermal lining. The head and tail of growing embryo, progressively,
93 push deeper in to the trophoblast, which become lined by a layer of somatic mesoderm
94 forming the extra-embryonic somatopleure (McGeady et al. 2006). The outer somatopleure
95 membrane forms chorion, while the inner somatopleure membrane, attached to the embryos at
96 umbilicus, forms amnion. Due to its early development, amnion could be a reservoir of
97 pluripotent stem cells, derived from the contiguity of embryonic epiblast, even at full term
98 pregnancy. Amnion is a thin, tough, membranous and fluid-filled sac which surrounds and
99 protects the embryo and then the foetus (McGeady et al. 2006). When first formed, it is in

100 contact with the embryo's body, but, after few weeks of pregnancy, fluid begins to
101 accumulate within it. This fluid increases in quantity relatively rapidly up to mid-pregnancy
102 and then gradually decreases. In early pregnancy, the composition of amniotic fluid and
103 plasma are similar and may be considered a dialysate of the maternal or foetal extracellular
104 fluid. Later in gestation, when the foetus urethra becomes patent, foetal urine passes in
105 amniotic cavity. Amniotic fluid (AF) contains less than 2% of solids, consisting of urea and
106 other extractives, inorganic salts, a small amount of protein, a trace of sugar and multiple cell
107 types from embryonic and extra-embryonic tissues (Gosden 1983).

108 With the progress of gestation, the embryo sinks into the amniotic membrane (AM), which
109 wraps the mesoblastic axis in which allantois and umbilical vessels developed, giving rise to
110 the umbilical cord (UC) (Barone 1994). Based on structural and functional studies, at least
111 four distinctive zones are now recognized in the UC: surface amniotic epithelium,
112 subamniotic stroma, perivascular stroma, vessels. All together these structures are called
113 umbilical cord matrix (UCM). The connective tissue included between subamnion and
114 perivascular regions consists of mucous, mesenchymal connective tissue, and it is called
115 Wharton's jelly (WJ). It contains a very low number of cells and a high amount of
116 extracellular matrix components, such as collagen, hyaluronic acid and proteoglycans
117 (Sobolewski et al. 1997). These substances make this tissue highly hydrated and resistant to
118 extension and compression evoked by foetal movements and uterine contraction. Stromal
119 cells of WJ, previously described as myofibroblast (Takechi et al. 1993), have properties of
120 potentially multipotent SCs (Mitchell et al. 2003).

121

122 **Amniotic membrane**

123

124 After mechanically separating amnion from allantois, as in human medicine, it is possible to
125 harvest two cellular types, amniotic epithelial (AECs) or mesenchymal cells (AMSCs). AECs

126 have a typical polygonal morphology while AMSCs are fibroblast-like. Both AMSCs and
127 AECs from different species have been investigated for proliferative potential, differentiation
128 ability, and phenotypically characterized.

129 AECs showed similar patterns to AMSCs in the sheep (Gloria et al. 2010; Mauro et al. 2010;
130 Mattioli et al. 2012), but the latter were superior in promoting interbody fusion in induced
131 anterior cervical discectomy after allogenic implantation combined with bone grafts
132 (Goldschlager et al. 2011). AECs phenotype, methylation status, immunomodulatory and
133 stemness properties are affected by gestational age (Barboni et al. 2014). In the horse (Lange-
134 Consiglio et al. 2012) and in the bovine (Corradetti et al., 2013), AECs showed a lower
135 number of doublings compared to AMSCs but the same phenotype and plasticity at
136 differentiation. Feline AECs were fully characterized and were able to differentiate into
137 adipogenic, chondrogenic, osteogenic and neurogenic lineages (Rutigliano et al. 2013).
138 Porcine AECs were successfully used for construction of tissue-engineered cornea (Luo et al.
139 2013).

140 Focusing on MSCs, they express pluripotency and mesenchymal markers, have a multi-
141 lineage differentiation potential and possess a privileged immunogenic status related to the
142 low expression or even absent levels of MHC I and MHC II (data are summarized in Table 1).
143 AMSCs were capable of inhibiting peripheral blood mononuclear cell proliferation after
144 allogenic stimulation either when co-cultured in cell-to-cell contact or when separated by a
145 transwell membrane (Lange-Consiglio et al. 2013b). Furthermore, it was demonstrated that
146 AMSCs possess nonteratogenicity 4 weeks post injection into immunodeficient mice (Vidane
147 et al. 2014).

148 Comparing results from different species, among pluripotency markers, a difference was
149 found for NANOG expression. While in the horse (Violini et al. 2012) and buffalo (Dev et al.
150 2012; Mann et al. 2013) AMSCs expressed NANOG at RT-PCR, canine AMSCs were

151 negative (Filioli Uranio et al. 2011, 2014). Anyway, Park et al. (2012) could quantified it in
152 canine AMSCs using qPCR.

153 Mesenchymal markers were expressed in all species while haematopoietic markers were not
154 found. In the horse, CD34 was not expressed in early stages of culture, but began to be
155 expressed at later passages (P5) (Lange-Consiglio et al. 2013a). By the way, the consensus
156 about absence of CD34 expression on MSCs has been challenged. The CD34 expression
157 status in haematopoietic and MSCs appears to depend on the environment, and can change
158 from positive to negative, and vice versa (Lin et al. 2012).

159 Karyotype of cultured AMSCs was confirmed to be stable in the dog (Filioli Uranio et al.
160 2011, 2014), buffalo (Dev et al. 2012c; Mann et al. 2013) and horse (Seo et al. 2013). In the
161 dog, a high telomerase activity was found (Filioli Uranio et al. 2011). Telomerase activity and
162 telomere maintenance characterize all cells with self-renewal capability, such as ESCs, MSCs,
163 and most cancer cells (Colosimo et al. 2013a).

164

165 **Amniotic fluid**

166 In the veterinary field, as in the human, AF contains a heterogeneous population of cells:
167 cuboidal epitheloid (E-cells), round (R-cells) and spindle-shaped fibroblastic (F-cells) cells.
168 E-cells probably derive from foetal skin and urine and they are lost during the first passages
169 of culture; R-cells are supposed to originate from foetal membranes and trophoblasts; F-cells
170 are generated from mesenchymal tissues and are supposed to be the MSC population of the
171 AF (Klemmt et al. 2011). In domestic animals, the prevalent cell population that persist
172 during culture is the F-type, except for one study on porcine AFMSCs (Sartore et al. 2005)
173 and one on bovine AFMSCs (Rossi et al. 2014), where both F-cells and R-cells persist
174 through passages. Other groups, that studied bovine AF cell populations (Corradetti et al.
175 2013; Gao et al. 2014; da Cunha et al. 2014), found R-cells only at the very first passages,
176 maybe due to the addition in the culture medium of growth factors like EGF and bFGF.

177 AFMSCs generally express MSC markers and mesodermal markers like Vimentin and α -
178 Smooth Muscle Actin (α SMA) and are negative for haematopoietic markers (Table 2). In the
179 horse, expression of CD73 is controversy. Positivity was found only for AD2 clone, while
180 clone 5F/B9 did not cross-react (Gulati et al. 2013). Anyway, for AD2 clone cross-reactivity
181 was not found in equine (Iacono et al. 2012a) and also in feline (Iacono et al. 2012b) and
182 bovine AFMSCs (Rossi et al. 2014). Like AMSCs, also AFMSCs express pluripotency
183 markers and only in one study in the bovine (Rossi et al. 2014) expression of OCT4, NANOG
184 and SOX2 was not found.

185 Different authors verified that AFMSCs show telomerase activity and that telomere length is
186 not affected by culture conditions (Filioli Uranio et al. 2011; Colosimo et al. 2013a), such as
187 karyotype (Filioli Uranio et al. 2011; Dev et al. 2012a; Dev et al. 2012b; Colosimo et al.
188 2013b; Weber et al. 2013; da Cunha et al. 2014).

189

190 **Umbilical cord blood (UCB)**

191

192 In human medicine, the use of UCB as a source of MSCs can be traced back to 2000: in this
193 year Erices et al. showed that UCB cells were able to give rise to two types of adherent cells,
194 one of which expressed antigens typical of MSCs. Also in veterinary medicine cells with
195 characteristics similar to human UCBMSCs have been isolated in different species (Table 3).
196 Usually, an isolation rate of fibroblast-like, spindle-shaped cells, similar to human, between
197 50 and 70% is reported (Koch et al. 2007; Iacono et al. 2012a; Mohanty et al. 2014), except
198 for feline UCB where cells at P0 comprised a heterogeneous population, including fibroblast-
199 like, spindle-shaped, and small round morphologies (Jin et al. 2008). Reported data may be
200 due to the limited number of MSCs in UCB as well as suboptimal isolation and expansion
201 conditions. In horses was demonstrated that samples cultured at 5% O₂ proliferated rapidly to
202 the desired cell number. The effect of mild hypoxia may be attributable in part to the

203 induction of signaling pathways but also to a decrease in survival of contaminating leukocytes
204 in UCB, thus increasing the isolation rate up to 80% (Schuh et al. 2009). Furthermore,
205 particularly in equine species, the UCB recovered volume is often low, because of a wide
206 spread belief that the mare continues to pass blood to the foal through the umbilicus
207 immediately post foaling (Rossdale 1958). However, using Doppler ultrasound, Doarn (1985)
208 showed that blood flow through the cord ceases within few minutes following delivery. All
209 authors reported that UCBMSCs cells from domestic animals sources were positive for CD44
210 (Table 3), an antigen expressed by a variety of cells types, not part of the ISCT MSC
211 definition (Dominici et al. 2006). Given its constant expression and considering the difficulty
212 in UCBMSCs isolation, sorting of CD44+ cells prior to plastic adherence isolation could lead
213 to more homogenous populations in domestic animals (Paebst et al. 2014). A variability of the
214 other MSCs marker expression exists particularly for CD73. Similarly to AFMSCs, in the
215 horse only Mohanty et al. (2014) showed that UCBMSCs expressed CD73, both with RT-
216 PCR and flow cytometry. Authors explained their finding with the use of different CD73
217 clones (5F/B9), compared with a previous study reporting its negative expression (De
218 Schauwer et al. 2012; 10fl clone). However, Mohanty's research group stated that equine
219 AFMSCs reacted with AD2 clone, but not with 5F/B9 clone (Gulati et al. 2013), showing
220 opposing results for the same species. Anyway, differences in the culture medium, such as the
221 content of FBS, the timeframe of plastic-adherent culture and the cell harvesting technique
222 could influence the cell characteristics (Paebst et al. 2014).

223 UCBMSCs from different species possess an intermediate phenotype that more closely
224 resembles ES cells: Tra1-60, Tra1-81 and Oct4 are present (Reed et al. 2008; Seo et al. 2009;
225 Raoufi et al. 2011; Mohanty et al. 2014), but conflicting data are reported for Nanog and Sox2
226 (Reed et al. 2008; Mohanty et al. 2014), particularly in the horse.

227

228 **Umbilical Cord**

229

230 UC is routinely discarded at parturition and its extracorporeal nature facilitates isolation by
231 eliminating the invasive and discomfort extraction procedures as well as patient risks that
232 attend adult stem cell isolation. Most significantly, the comparatively large volume of UC and
233 ease of physical manipulation theoretically increase the number of stem cells that can be
234 extracted, which make it possible to get substantial number of cells in several passages
235 without need of long term culture and extensive expansion *ex vivo*. Probably due to the
236 possible absence of this structure at birth, because of reduction of the water and substances
237 content towards the end of pregnancy, different authors, particularly in veterinary medicine,
238 named interchangeably WJ or UCM. Due to this reason, in Table 4, we reported data
239 regarding WJ and UCM MSCs together. Usually, spindle-shaped cells have been isolated;
240 however few authors described a heterogeneous cell population (fusiform or spindle form and
241 small round cells with large and prominent nucleus), from porcine and caprine UC,
242 characterized by a plateau or a stationary phase (Mitchell et al. 2003; Babaei et al. 2008). In
243 these species, such as in canine (Filioli Uranio et al. 2011) and bovine (Cardoso et al. 2012),
244 UC cells expressed high levels of telomerase activity, and this phenomenon could explain the
245 long term of cell culture observed by the authors. Moreover, tissue culture procedure does not
246 alter chromosomal organization (Filioli Uranio et al. 2011; Cardoso et al. 2012; da Cunha et
247 al. 2014). As reported above, also cell isolated from UC are negative for hematopoietic
248 markers and express a number of antigens associated with adult MSCs (Table 4). Conflicting
249 data are reported regarding their expression of embryonic markers. Cells from porcine,
250 bubaline and equine UC resulted positive for pluripotency genes OCT4, NANOG, SOX2 at
251 different culture passages (Mitchell et al. 2003; Carlin et al. 2006; Hoynowsky et al. 2007;
252 Singh et al. 2013; Sreekumar et al. 2014). In canine, all authors agree that during *in vitro*
253 passages cells lost OCT4 expression. However, while Lee et al. (2013a) stated that expression
254 of NANOG and SOX2 is negatively correlated with the number of passages but is always

255 present, most recently Filioli Uranio et al. (2014) showed that NANOG gene is present only
256 in cells recovered at late gestational age. This discrepancy is likely due to different culture
257 conditions and to different selection of cord tissue to be processed. The immature cells that
258 retain the ability to proliferate were located close to the amniotic surface, whereas highly
259 differentiated, non-proliferating fibroblasts were located in closer proximity to the umbilical
260 vessels (Nanaev et al. 1997).

261

262 **Clinical Applications**

263

264 Clinical application of foetal MSCs in domestic animals had an increase in the last five years.
265 The majority of the studies are on AFMSCs, mainly used in animal-model diseases (heart,
266 diaphragm, trachea and tendon induced injuries in different species) (Table 5). Among tissue-
267 engineering studies in the sheep, diaphragmatic repair with a mesenchymal amniocyte-based
268 engineered tendon led to improved structural outcomes when compared with equivalent fetal
269 myoblast-based and acellular graft (Kunisaki et al. 2006c). Investigating the ability of
270 AFMSCs for phenotypic conversion to vascular cells and cardiomyocytes (CM) when
271 autotransplanted in a porcine model acute ischemic myocardium (AMI), MSCs were able to
272 transdifferentiate to cells of vascular cell lineages but not to CM. Thus, porcine AFMSCs may
273 require further ex vivo re-programming to be suitable for therapeutic use in AMI (Sartore et
274 al. 2005). On the other hand, intramuscular injection of porcine AFMSCs reduced scar size
275 and preserved heart function after induced myocardial infarction in mice (Peng et al. 2014).
276 Only in the horse AFMSCs were used, combined with PRP (platelet rich plasma), to treat
277 spontaneous decubitus ulcers (Iacono et al. 2012c). When comparing different treatments
278 (aloe gel, PRP alone, MSCs+PRP), mean regression of ulcer treated with MSCs+PRP was
279 higher than others.

280 Despites animal UC cells molecular characterization, in vitro differentiation and their
281 immunosuppressive effects on T-cells in vitro (Cardoso et al. 2012), there are sparse
282 literatures documenting their therapeutic applications (Table 5). Equine UCBMSCs were
283 successfully used as allogenic therapy in race-horses with spontaneous tendon lesions (Kang
284 et al. 2013; Van Loon et al. 2014). Canine UCBMSCs and WJMSCs were mainly applied to
285 treat induced spinal cord injuries or bone defects. Dog WJMSCs can replace adult MSCs in
286 clinical bone engineering procedures (Kang et al. 2012) and promote functional recovery in
287 spinal cord injured dogs after allogenic transplantation (Ryu et al. 2012). Caprine WJMSCs
288 allogenic (Azari et al. 2011) and xenogenic transplantation (Pratheesh et al. 2014) in a rabbit
289 model, could be successfully performed for treating induced skin wounds. In not-treated
290 wounds, incomplete re-epithelialization and thick granulation tissue were observed, whereas,
291 in treated wounds intact skin with complete re-epithelialization, no inflammation, and thin
292 granulation tissue were seen. Furthermore, in both studies, histomorphological evaluation
293 demonstrated better epithelization, lower neovascularisation, and denser collagen fibres in
294 treated wounds.

295 Although implantation of human AMSCs into animals resulted in successful and persistent
296 engraftment in multiple organs and tissues (Lindenmair et al. 2012), their application in
297 veterinary medicine are still limited (Table 5). In fact, except for a study to enhance cervical
298 interbody fusion in an ovine model (Goldschlader et al. 2011) and histological and functional
299 improvement after transplantation in a porcine model of chronic myocardial ischemia
300 (Kimura et al. 2012), AMSCs were only applied in equine spontaneous tendon injuries.
301 Allogenic transplantation resulted in a quick reduction in tendon size and ultrasonographic
302 cross-sectional area (Lange-Consiglio et al. 2012) and the re-injury rate was lower compared
303 to autologous BMMSCs transplantation (Lange-Consiglio et al. 2013c). The hypothesis of a
304 paracrine mechanism of tendon repair when AMSCs are injected was confirmed in vivo;
305 when horse tendon injuries were treated with conditioned medium, a lower rate of re-injuries

306 was observed compared to untreated animals (Lange-Consiglio et al. 2013b). AMSCs and
307 their conditioned medium were recently considered candidates for equine uterine regenerative
308 therapy, due to their ability to increase in vitro endometrial cell proliferation rate (Corradetti
309 et al. 2014). So far, AMSCs are promising not only for cell-based therapies, but also for novel
310 therapeutic biological cell-free products.

311

312 **Conclusion**

313 Altogether, these studies offer authoritative views on markers expression and therapeutic
314 potential of MSCs from foetal tissues and fluids in domestic animals. These sources are easily
315 available and relatively abundant, so MSCs may have an attraction compared to other
316 established SCs in different clinical approaches. Anyway, more clinical applications are
317 needed to fully understand their properties and to establish the future clinical use in the
318 treatment of various diseases.

319

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322

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689 **Table 1.** AMSCs characterization.

Species	Phenotype	<i>In vitro</i> differentiation	Ref.
Sheep	ICC: Oct4+,TERT+	Osteogenic, myogenic	(Mauro et al. 2010)
Horse	FCA: CD44+,CD90+,CD14+,CD45+,CD29-,MHCI-,MHCII- ICC: cKit+,Oct4+,CD105+	Adipogenic, chondrogenic, osteogenic	(Coli et al. 2011)
	RT-PCR: OCT4+,SOX2+,NANOG+,CD105+,CD34-	Adipogenic, chondrogenic, osteogenic	(Violini et al. 2012)
	ICC: Tra1-60+,SSEA3+,SSEA4+,Oct4+ RT-PCR: CD29+,CD105+,CD44+,CD166+,CD34-,MHCI+,MHCII-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Lange-Consiglio et al. 2012)
	FCA: CD44+,CD90+,CD105+,CD19-,CD20-,CD28-,CD31-,CD34-,CD38-,CD41a-,CD62L-,CD62P-,CD200-	Adipogenic, chondrogenic, osteogenic	(Seo et al. 2013)
	AP+ ICC: Tra1-60+, SSEA4+,Oct4+ RT-PCR: CD29+,CD105+,CD44+,CD166+,CD34-,MHCI+,MHCII-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Lange-Consiglio et al. 2013a)
	FCA: Oct4+,SSEA4+,cMyc+,CD34- RT-PCR: CD29+,CD44+, CD166+, CD105+, CD34-MHCI+,MHCII-		(Corradetti et al. 2014)
Dog	RT-PCR: OCT4+,NANOG-,CD44+,CD184+,CD29+,CD34-,CD45-,DLA79-,DLA88-,DLA-DRA1-,DLA-DQA1-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Filioli Uranio et al. 2011)
	FCA: CD90+,CD105+,CD3-,CD11c-,CD28-,CD34-,CD38-,CD41a-,CD45-,CD62L- qPCR: OCT4+,SOX2+,NANOG+,KLF4+	Adipogenic, chondrogenic, osteogenic, neurogenic	(Park et al. 2012)

	RT-PCR: OCT4+,NANOG-,CD29+,CD44+,CD184+,CD34-,CD45-,DLA79-,DLA88-,DLA-DRA1-,DLA-DQA1-	Osteogenic, neurogenic	(Filioli Uranio et al. 2014)
Cat	FCA: CD44+,CD90+,CD105+,CD14-,CD34-,CD45-,CD73-	Adipogenic, chondrogenic, osteogenic	(Iacono et al. 2012b)
	FCA: CD73+,CD90+,CD34±,CD45-,CD79-	Adipogenic, chondrogenic, osteogenic	(Vidane et al. 2014)
Buffalo	AP+ RT-PCR: OCT4+,SOX2+,NANOG+,AP+,SCF+,NESTIN+		(Dev et al. 2012c)
	AP+ RT-PCR: OCT4+,SOX2+,NANOG+	Osteogenic	(Mann et al. 2013)
Bovine	RT-PCR: OCT4+,Myc+,CD44+,CD166+,CD73+,CD29+,CD105-,CD45-,CD14-,CD34-,MHCI+,MHCII-	Adipogenic, chondrogenic, neurogenic	(Corradetti et al. 2013)

690 +: positive, -: negative; ±: weakly positive; ICC: immunocytochemistry.

691 **Table 2.** AFMSCs characterization.

Species	Phenotype	<i>In vitro</i> differentiation	Ref.
Sheep	ICC: FSP+,CD31-,Vimentin+, α SMA+,Desmin-,CK8+,CK18+		(Kaviani et al. 2001)
	ICC: Vimentin+,CK8+,CK18+,CD31-	Chondrogenic	(Kunisaki et al. 2006a)
		Chondrogenic	(Kunisaki et al. 2006b)
	FCA: CD29+,CD44+,CD90 \pm ,CD105 \pm ,CD31-	Adipogenic, osteogenic	(Kunisaki et al. 2007)
	ICC: Oct4+,TERT+	Osteogenic, myogenic	(Mauro et al. 2010)
	FCA: CD44+,CD58+,CD166+,CD14-,CD31-,CD45-	Adipogenic, osteogenic	(Shaw et al. 2011)
	FCA: CD29+,CD44+,CD105+,CD166+,CD31-		(Gray et al. 2012)
	ICC: SSEA4+,STRO4+,CD29+,CD44+,CD166+ RT-PCR: OCT4+,NANOG+,SOX2+		(Weber et al. 2012)
	ICC: TERT+,Sox2+,Nanog+ semiqPCR: TERT+,NANOG+,SOX2 \pm ,OCT4 \pm	Osteogenic	(Colosimo et al. 2013a)
	FCA: Sox2+,Nanog+,TERT+,CD166+,CD29 \pm ,CD58 \pm ,CD14-,CD31- ,CD45-,c-Kit- ICC: Sox2+,Nanog+,TERT+,c-Kit-	Adipogenic, osteogenic, tenogenic	(Colosimo et al. 2013b)
	FCA: TERT+,Oct3/4+,Nanog+,Sox2+,CD29+,CD58+,CD90+,CD166 \pm ,CD14- ,CD31-,CD45-,CD49f-,CD117-,HLA-ABC+, HLA-DR-	Osteogenic	(Di Tomo et al. 2013)
	FCA: CD44+,CD166+,CD29+,STRO4+ ICC: CD29+,CD44+,STRO4+,CD166 \pm ,CD11b-	Chondrogenic, osteogenic, endothelial, myogenic/myofibroblasts	(Weber et al. 2013)

	, Vimentin+, α SMA \pm , Desmin \pm qPCR: NANOG+, STAT3+		
Swine	ICC: SSEA4 \pm , Oct4 \pm , CD90+, Vimentin+, vWf+, SM22+, VE- Cadherin \pm , α SMA \pm , CD31 \pm , CD105-, CD117-, CD34-, CD45-, Sca-1- , CardiacTnI-		(Sartore et al. 2005)
	RT-PCR: OCT4+, SOX2+, THY+	Adipogenic, osteogenic, neurogenic, myogenic, endothelial	(Zheng et al. 2009)
	RT-PCR: OCT4+, SOX2+, THY+ FCA: CD44+, CD166+, CD34-, CD45-, CD54- ICC: Oct4+, Nanog+, Tra1-60+, Tra1-81+, SSEA4+, SSEA1- RT-PCR: CD90+, CD117+, HLA-ABC+, CD45-	Neurogenic Adipogenic, neurogenic, cardiomyogenic; Multilineage- differentiation through EBs	(Zheng et al. 2010) (Chen et al. 2011)
	FCA: CD44+, CD90+, CD31-, CD4a-		(Peng et al. 2014)
Horse	ICC: Oct4+, SSEA4+, Tra1-60 \pm RT-PCR: CD29+, CD44+, CD105+, CD34-, ELA-ABC+, ELA-DR-	Adipogenic, chondrogenic, osteogenic	(Lovati et al. 2011)
	FCA: CD44+, CD90+, CD105+, CD14-, CD34-, CD45-, CD73-	Adipogenic, chondrogenic, osteogenic	(Iacono et al. 2012a)
	AP+ FCA: CD29+, CD44+, CD90+, CD73(clone 5F/B9)-, CD73(clone AD2)+, CD34-, CD45- ICC: CD73+, CD90+, CD34-, CD45- RT-PCR: CD73+, CD90+, CD105+, CD14-, CD34-, CD45-	Adipogenic, chondrogenic, osteogenic, tenogenic	(Gulati et al. 2013)

Dog	RT-PCR: OCT4+,NANOG-,CD44+,CD184-,CD29-,CD34-,CD45-,DLA79+,DLA-DRA1+,DLA-DQA1-,DLA 88-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Filioli Uranio et al. 2011)
	ICC: Vimentin+, Nestin+, CK18+	Adipogenic, chondrogenic, osteogenic, neurogenic	(Fernandes et al. 2012)
	FCA: CD29+,CD44+,CD90+,CD34- ICC: Oct4+,Nanog+,Sox2+,CD29+,CD44+,CD90+,CD34- RT-PCR: OCT4+,NANOG+,SOX2+	Adipogenic, chondrogenic, osteogenic, hepatogenic	(Choi et al. 2013)
		Neurogenic	(Kim et al. 2014)
Cat	FCA: CD44+,CD90+,CD105+,CD34±,CD14-,CD45-,CD73-	Adipogenic, chondrogenic, osteogenic	(Iacono et al. 2012b)
Buffalo	AP+ RT-PCR: OCT4+,NESTIN+,FGF5+		(Dev et al. 2012a)
	AP+ RT-PCR: OCT4+,NANOG+,SOX2+,AP+,SCF+,CyclinA+,NESTIN+,FGF5+	Neurogenic	(Dev et al. 2012b)
	AP+ qPCR: OCT4+,NANOG+,SOX2+		(Yadav et al. 2012)
Caprine	FCA: Nanog+,Sox2+ ICC: Oct4+,Nanog+,Sox2+,SSEA4+,SSEA1+ RT-PCR: CD73+,CD90+,CD105+,CD34-	Adipogenic, chondrogenic, osteogenic	(Pratheesh et al. 2013)

Bovine	RT-PCR: OCT4+,MYC+,CD29+,CD44+,CD73+,CD166+,CD14- ,CD34-,CD45-,CD105-,CD117+,MHCI+,MHCII-	Adipogenic, chondrogenic, neurogenic	(Corradetti et al. 2013)
	ICC: Oct4+,CD29+,CD44+,CD71+,CD34- RT-PCR: OCT4+,CD29+,CD44+,CD73+,CD166+,CD34-,CD45-	Adipogenic, chondrogenic, neurogenic	(Gao et al. 2014)
	FCA: CD34+,CD44+,CD90+,CD105+,CD14- ICC: Oct4-,SSEA4±,αSMA+,Vimentin+,N-Cadherin+,E-Cadherin-,CK- qPCR: OCT4-,NANOG-,SOX2-	Adipogenic, chondrogenic, osteogenic	(Rossi et al. 2014)

692 +: positive, -: negative; ±: weakly positive; ICC: immunocytochemistry.

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694 **Table 3.** UCBMSCs characterization.

Species	Phenotype	<i>In vitro</i> differentiation	Ref.
Sheep		Chondrogenic	(Fuchs et al. 2005)
		Adipogenic, chondrogenic, osteogenic	(Jager et al. 2006)
	FCA: CD44+,CD38-,CD45-,CD41/61-		(Fadel et al. 2011)
Horse		Adipogenic, chondrogenic, osteogenic	(Koch et al. 2007)
	ICC: Oct4-,SSEA1-,SSEA3-,SSEA4-,Tra1-60-,Tra1-81-,AP+,CD29+,CD44+,CD90+,CD14-,CD79 α -,Coll I+,Coll II+,Coll III+,Coll IX/XI+,MHCI \pm ,MHCII-	Adipogenic, chondrogenic, Osteogenic	(Guest et al. 2008)
	AP +	Adipogenic, chondrogenic, hepatogenic, myogenic	(Reed and Johnson, 2008)
	ICC: Oct4+,Tra1-60+,Tra1-81+,SSEA1+,SSEA4 \pm	Chondrogenic	(Berg et al. 2009)
	ICC: Vimentin \pm , α SMA \pm ,CD18-,PanCK-,Factor VIII-,Osteonectin-,Osteocalcin-	Adipogenic, chondrogenic, osteogenic	(Schuh et al. 2009)
		Osteogenic	(Toupadakis et al. 2010)
		Osteogenic	(Figueroa et al. 2011)
	FCA: CD86-,MHCI-,MHCII-		(Carrade et al. 2011)
FCA: CD29+,CD44+,CD90+,CD86-,MHCI+,MHCII-,F6B-		(Carrade et al. 2012)	
FCA: CD29+,CD44+,CD90+,CD79 α -,MHCII-	Adipogenic, chondrogenic,	(De Shauwer et al. 2011)	

		osteogenic	
	FCA and Confocal Microscopy: CD29+,CD44+,CD90+,MHCII-,CD105±,CD45- ,CD73±,CD79α-,monocyte marker-	Adipogenic, chondrogenic, osteogenic	(De Schauwer et al. 2012)
		Adipogenic, chondrogenic, osteogenic	(Burk et al. 2013)
	FCA: CD29+,CD44+,CD73+,CD90+,CD105+,CD45-CD79α- ,MHCI+,MHCII-,monocyte marker- RT-qPCR: CD40+,CD80±,CD86-,TGFβ+,HGF±,IDO-,TNFα-	Adipogenic, chondrogenic, osteogenic,	(De Schauwer et al. 2013)
	FCA: CD90+,CD105+,CD20-,CD28-,CD38-,CD62L-CD200- ,CD31-,CD62P-,CD34-,CD41a-	Adipogenic, chondrogenic, osteogenic	(Kang et al. 2013)
	FCA: CD29+,CD44+,CD73+,CD90+,CD105+,CD45-CD79α- ,MHCI+,MHCII-,monocyte marker- qPCR: CD40+,CD80+,CD86-,TGFβ+,HGF±,IDO-,TNFα-	Adipogenic, chondrogenic, osteogenic,	(De Schauwer et al. 2014)
	FCA: CD29+,CD44+,CD73+,CD90+,CD34-,CD45- ICC: CD73+,CD90+,CD34-,CD45- RT-PCR: OCT4+,NANOG+,SOX2+,CD73+,CD90+,CD105+,CD14- ,CD34-,CD45-	Tenogenic	(Mohanty et al. 2014)
	FCA: CD29±,CD44+,CD73-,CD90-,CD105-,CD14-CD34- ,CD45-,CD79α-,MHCII-		(Paebst et al. 2014)
Dog		Osteogenic	(Jang et al. 2008)
	FCA: Oct4+,CD29+,CD33+,CD44+,CD105+,CD184+,CD4- ,CD8a-,CD10-,CD14-,CD24-,CD31-,CD34-,CD38-,CD41a- ,CD45-,CD49b-,CD41/61-,CD62p,CD73-,CD90-,CD133-,HLA-	Chondrogenic, osteogenic, neurogenic	(Seo et al. 2009)

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	FCA: CD44+,CD73+,CD90+,CD105+,CD14-,CD34-,CD45-	Osteogenic	(Kang et al. 2012)
Swine	ICC: CD29+,CD49b+,CD105+,CD45-,CD133-,SCF+,LIF+,G-CSF+	Adipogenic, chondrogenic, osteogenic	(Kumar et al. 2007)
Cat	FCA: CD9+,CD44+,CD18-,CD45- ICC: Vimentin+	Neurogenic	(Jin et al. 2008)
Bovine	RT-PCR: OCT4+,CD73+	Adipogenic, chondrogenic, osteogenic	(Raoufi et al. 2011)

695 + :positive; - : negative; ± : weakly positive; ICC: immunocytochemistry

696 **Table 4.** WJMSCs and UCMSCs characterization.

Species	Phenotype	<i>In vitro</i> differentiation	Ref.
Swine	Immunoblotting: c-kit+, α SMA+	Neurogenic	(Mitchell et al. 2003)
	AP+		(Carlin et al. 2006)
	ICC: Oct4+,Nanog+		
	RT-PCR: OCT4+,NANOG+,SOX2+		
	qPCR: OCT4+,NANOG+,SOX2+		
Horse	FCA: CD54+,CD90+,CD105+,CD146+,Oct4+,SSEA4+,c-kit+, SSEA3 \pm ,Tra1-60 \pm ,HLA-ABC-,HLA-1AG-,MHCII-	Adipogenic, chondrogenic, osteogenic	(Hoynowski et al. 2007)
	RT-PCR: Oct4+, Sox2+		(Cremonesi et al. 2008)
		Adipogenic, chondrogenic, osteogenic	(Passeri et al. 2009)
		Osteogenic	(Toupadakis et al. 2010)
	FCA: Oct4+,c-myc+,CD34-,CD105-,SSEA4-,SSEA3-,Tra1-60-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Corradetti et al. 2011)
	RT-PCR: CD29+,CD44+,CD105+,CD166+,CD14-CD34-, MHCI+,MHCII-		
	RT-PCR: CD29+,CD44+,CD105+,CD166+,CD14-CD34-, MHCI+,MHCII-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Lange-Consiglio et al. 2011)
	RT-PCR: CD29+,CD44+,CD105+,CD34-,MHCI+,MHCII-	Adipogenic, chondrogenic, osteogenic	(Lovati et al. 2011)
	ICC: Oct4+,SSEA4+,Tra1-60 \pm		
	RT-PCR: CaSR+		(Martino et al. 2011)
	FCA: CD86-,MHCI-,MHCII-		(Carrade et al. 2011)

	FCA: CD29+,CD44+,CD90+,CD86-,MHCI+,MHCII-,F6B-		(Carrade et al. 2012)
		Adipogenic, chondrogenic, osteogenic	(Burk et al. 2013)
	FCA: CD29+,CD44+,CD90+,CD45-,CD73-,CD79 α -,CD105-,MCHII-,Monocyte marker-		(De Schauwer et al. 2013)
	FCA: CD44+,CD90+,CD105+,CD34-,MHCII- ICC: MHCII-	Adipogenic, chondrogenic, osteogenic	(Barberini et al. 2014)
	FCA: CD29+,CD44+,CD73+,CD90+,CD105+,CD45-,CD79 α - MHCI+,MHCII-,monocyte marker- qPCR: CD40+,CD80+,CD86-,TGF β +,HGF \pm ,IDO-TNF α -	Adipogenic, chondrogenic, osteogenic,	(De Schauwer et al. 2014)
	FCA: CD29+,CD44 \pm ,CD73-,CD90-,CD105-,CD14-,CD34-, CD45-,CD79 α -,MHCII-		(Paebst et al. 2014)
Caprine	AP+ RT-PCR: NANOG-	Chondrogenic	(Co et al. 2014) (Babaei et al. 2008)
	AP+ FCA: CD44+,CD34- ICC: α SMA+		(Azari et al. 2011)
	FCA: STRO1+,CD73+,CD105+,CD34- ICC: STRO1+,CD73+,CD105+,CD34-	Adipogenic, chondrogenic, osteogenic	(Pratheesh et al. 2014)
Dog	RT-PCR: CD29+,CD44+,CD184+,OCT4+,CD34-,CD45-, DLA-DQA1-,DLA79-,DLA88-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Filioli Uranio et al. 2011)

	FCA: CD44+,CD73+,CD90+,CD105+,CD14-,CD34-,CD45-	Osteogenic	(Kang et al. 2012)
	FCA: CD44+,CD73+,CD90+,CD105+,CD14-,CD34-,CD45-	Adipogenic, osteogenic, neurogenic	(Ryu et al. 2012)
	FCA: CD90+,CD105+,CD11c-,CD34-,CD3-,CD45-,CD28-,CD38-,CD62L,CD41a-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Seo et al. 2012)
	RT-PCR: CD44+,CD54+,CD61+,CD80+,CD90+,CD105+	Chondrogenic	(Lee et al. 2013a)
	FCA: CD44+,CD90+,CD105+,CD184+,CD29-,CD33-,CD34-,CD45- ICC: Oct3/4+,Nanog+,Sox2+,SSEA4+,CD44+,CD90+,CD105+,CD184+ RT-PCR: OCT3/4+,NANOG+,SOX2+,CD44+,CD90+,CD105+,CD184+,CD29-,CD34-,CD45-	Adipogenic, chondrogenic, osteogenic, neurogenic	(Lee et al. 2013b)
	FCA: CD29+,CD44+,CD184+,OCT4+,CD34-,CD45-,DLA-DRA-,DLA-DQA1-,DLA79-,DLA88-	Osteogenic, neurogenic	(Filioli Uranio et al. 2014)
Bovine	RT-PCR: OCT4+,ITSN1+	Adipogenic, chondrogenic, osteogenic, neurogenic	(Cardoso et al. 2012a)
		Neurogenic	(Cardoso et al. 2012b)
Buffalo	FCA: CD29+,CD73+,CD90+,CD105+,CD34-,CD36-,CD45- RT-PCR: OCT4+,NANOG+,SOX2+	Adipogenic, osteogenic	(Singh et al. 2013)
	AP+ ICC: Oct4+,Nanog+,Sox2+,CD73+,CD90+,CD105+,CD34- RT-PCR: CD73+,CD90+,CD105+ qPCR: OCT4+,NANOG+,SOX2+	Adipogenic, chondrogenic, osteogenic	(Sreekumar et al. 2014)

698 **Table 5.** Clinical applications of foetal MSCs in domestic animals.

MSCs source	Species	Clinical application	Ref.
AM	Sheep	Discectomy with bone graft	(Goldschlaget et al. 2011)
	Horse	Spontaneous tendon and ligament injuries	(Lange-Consiglio et al. 2012; 2013b; 2013c)
	Pig	Myocardial ischemia	(Kimura et al. 2012)
AF	Sheep	Diaphragmatic repair with graft	(Fuchs et al. 2004; Turner et al. 2011)
		Prenatal tracheal reconstruction with scaffold	(Kunisaki et al. 2006b; Gray et al. 2012)
		Diaphragmatic repair	(Kunisaki et al. 2006c)
		Prenatal treatment with lentiviral transduced cells	(Shaw et al. 2011)
		Prenatal cell-based heart valves implantation	(Weber et al. 2012)
		Tendon injuries with nucleofected cells	(Colosimo et al. 2013a)
	Swine	Cardiac ischemia	(Sartore et al. 2005)
		Myocardial infarction in mouse	(Peng et al. 2014)
	Horse	Spontaneous decubitus ulcers with PRP	(Iacono et al. 2012c)
UCB	Horse	Intra-articular injection	(Carrade et al. 2011)
		Spontaneous tendinitis of superficial digital flexor tendon	(Kang et al. 2013)
		Spontaneous tendon and ligament disorders	(Van Loon et al. 2014)

	Dog	Spinal cord injury	(Lim et al. 2007; Park et al. 2011)
		Diaphyseal defect in the radius	(Jang et al. 2008)
		Bone regeneration in ectopic implantations	(Byeon et al. 2010)
		Segmental bone defects	(Kang et al. 2012)
WJ/UCM	Swine	Transplantation into rat brains	(Weiss et al. 2003)
		Allogeneic engraftment in the neonatal intestine	(Miller et al. 2012)
	Caprine	Skin wounds	(Azari et al. 2011)
		Skin wounds in rat	(Pratheesh et al. 2014)
	Horse	Intra-articular injection	(Carrade et al. 2011)
	Dog	Segmental bone defects	(Kang et al. 2012)
		Spinal cord injuries	(Ryu et al. 2012)