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1	In vitro differentiation of porcine Aortic Vascular Precursor Cells to Endothelial and
2	Vascular Smooth Muscle cells
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25	
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- 30
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 32 Muscle Differentiation, Pig Animal Model
- 33

34 Abstract

35 Recent findings suggested that progenitor cells and multipotent mesenchymal stromal cells (MSCs) 36 are anatomically and functionally associated to vascular niches and proposed that cells isolated 37 from a vessel that display mesenchymal proprieties and that are able to differentiate to the whole 38 component of a functional blood vessel (including endothelial and smooth muscle cells), can be 39 defined Vascular Stem Cells (VSCs).

40 We recently isolated and characterized a population of MSC-like cells from porcine aorta (porcine 41 Aortic Vascular Precursor Cells pAVPCs) with MSCs and pericyte-like proprieties in vitro. The 42 main aim of the present work was to investigate whether pAVPCs possess VSC-like properties 43 assessing their differentiation potential toward the endothelial and the smooth muscle lineages. 44 pAVPCs have been isolated as previously described by us and cultured in in a medium specified for 45 pericyte to maintain stemness and multipotency. In order to assess if pAVPCs differentiate to 46 smooth muscle cells, pAVPCs were grown for 21 days in high glucose – DMEM + 10% Foetal 47 Bovine Serum (DMEM + 10% FBS) (Long Term Medium – LTM), whilst in order to assess if cells 48 are able to differentiate toward the endothelial phenotype cells were seeded in a 24 well plate (5000 49 cells/cm2) and cultured for 21 days in Endothelial Differentiation Medium (EDM) (human 50 Endothelial Serum Free Medium supplemented with 5% FBS and 50 ng/mL of human recombinant 51 Vascular Endothelial Growth Factor [hVEGF]).

After 21 days of culture in LTM, pAVPCs showed an elongated fibroblast-like shape and seems to organize in cord-like structures. Quantification of smooth muscle cells markers showed a significant increment of the three transcripts (αSMA, CNN1, SMM-hc) and immunofluorescence confirmed the presence of αSMA and SMM proteins. pAVPCs cultured in EDM displayed an endothelial cell-like morphology and revealed the over-expression of the typical endothelial markers: CD31, VE-Cadherin, vWF and eNOS. Immunofluorescence of CD31 confirmed the presence and the typical distribution of this protein. In conclusion we can be defined pAVPCs as a

- 59 population of VSC-like cells considering that they express markers of MSCs, display the classical
- 60 MSC trilineage differentiation and differentiate, *in vitro*, toward smooth muscle and endothelial
- 61 cells phenotypes.

63 Introduction

64 Recent findings and theories suggested that tissue-specific progenitor cells and multipotent

65 mesenchymal stromal cells (MSCs) are anatomically and functionally associated to

66 perivascular/vascular niches (11, 38, 40-43, 52, 53). According to these theories, blood vessels

67 distributed in the whole body, could be considered a systemic reservoir of multipotent

68 stem/progenitor cells (5, 44). In a recent review, Lin and Lue (29) proposed that cells isolated from

a vessel that display MSC proprieties and that are able to differentiate to the whole component of a

70 functional blood vessel (including endothelial and smooth muscle cells), can be defined vascular

71 stem cells (VSCs) (29).

72 The presence of VSCs has been demonstrated in different type of vessels including embryonic, fetal 73 and adult aorta. Undifferentiated mesenchymal cells from human fetal aorta has been isolated by 74 Invernici and colleagues (22, 23). The expression of endothelial and myogenic markers has been 75 demonstrated for these cells which, if opportunely stimulated with VEGF or PDGFB, were able to 76 give rise, respectively, to endothelial and mural cells. Other populations of progenitor cells have 77 been isolated from human fetal aorta; in particular, Fang and colleagues (14) described a population 78 of CD105, VEGF Receptor 2 (VEGFR2, also known as Flk1) positive and CD34 negative vascular 79 cells which were able to differentiate to endothelial and vascular smooth muscle cells, if cultured, 80 respectively, with VEGF or PDGFB, and to the osteogenic and the adipogenic lineages. Both cells 81 isolated by Invernici and colleagues and by Fang and colleagues displayed high angiogenic 82 potential in vivo, too (13, 22, 23).

Adult aorta is a good source of vascular mesenchymal stromal cells, too, as reported by Pasquinelli and colleagues (35) which isolated from tunica media and adventitia, respectively, CD34 positive and c-kit positive progenitor cells. Both cell populations were positive for the expression of MSCs markers *in vitro* and, in particular, a VEGF treatment induced a phenotypical shift to the endothelial lineage with the upregulation of the expression of Flk1 and von Willebrand Factor (vWF).

Moreover, these cells were able to form capillary like structure in an *in vitro* angiogenesis assay
(35).

90 Although the high potential of MSCs and precursor cells from vasculature in field of regenerative 91 medicine (6, 10, 12, 24), several paper described these cells as potentially involved in pathogenesis 92 of different diseases (20). Using a Knock Out (KO) mouse model of Apolipoprotein E (ApoE), Hu 93 and colleagues showed that adventitial progenitor cells contributed to atherosclerotic lesion in vein 94 grafts (21). Instead, in 2010, Juchem and colleagues described an high pro-thrombogenic potential 95 of intimal pericytes in endothelial-denuded vascular region which support the hypothesis that these 96 cells are involved in the pathogenesis of atherosclerosis, thrombosis and saphenous vein graft 97 disease (26). Moreover, in 2012, in a well performed work by Tang and colleagues, it has been 98 described a population of stem cells resident in the blood vessel wall, named multipotent vascular 99 stem cells, that were able to differentiate to a MSC-like phenotype and subsequently to the smooth 100 muscle one, spontaneously. In that work it has been also shown that in response of vascular injuries 101 these cells become proliferative and differentiate into smooth muscle and chondrogenic cells 102 contributing to vascular remodeling and neointimal hyperplasia (42). 103 Considering the potential of perivascular cells in field of regenerative medicine (10) and 104 cardiovascular physio-pathology (20) and the usefulness of pig as animal model in these fields (15, 105 17, 27, 39, 46, 50, 51), we recently isolated and characterized a population of MSC-like cells from

- 106 porcine aorta. We named these cells porcine Aortic Vascular Precursor Cells (pAVPCs) for their
- 107 differentiation potential and for their pericyte-like proprieties *in vitro* (49).
- 108 Despite pAVPCs has been properly characterized for their morphological, phenotypical and
- 109 functional properties as pericytes-like cells (49), we observed (unpublished data) that they tend to
- 110 lose their trilineage multipotency and to undergo senescence during *in vitro* culture.

111	Therefore in the present work we decided to modify the previous protocol (49) and we cultured
112	pAVPCs in a medium specified for pericyte (Percyte Growth Medium) and we verified in this new
113	culture condition MSC-like and pericyte-like properties.
114	The main aim of the present work was to investigate whether pAVPCs cultured in Pericytes Growth
115	Medium possess VSC-like properties assessing their differentiation potential toward the endothelial
116	and the smooth muscle lineages in order to define these cells Vascular Stem Cells
117	
118	Material and methods
119	
120	Cells isolation and culture
121	pAVPCs have been isolated as previously described by us (49) from 3-mo-old pigs, euthanized for
122	other experimental purposes, to generate three primary cell culture replicates. In line with the

123 reduction rule, an animal-sharing approach was used; the aorta donor animals were the controls of

124 an experimental trial conducted according to relevant Italian and international guidelines. All

125 procedures on pigs were reviewed and approved in advance by the Ethics Committee of the

126 University of Bologna (Bologna, Italy) and were then approved by the Italian Ministry of Health.

127 Briefly, the cells were isolated, from the media layer of the aortas through a collagenase IA

128 digestion. The cells were cultured overnight (15-16 h) in high glucose (hg) DMEM (GIBCO – Life

129 Technology Corporation, Carlsbad, CA, USA) to which 10% FBS and 10X antibiotic-antimycotic

130 (hgDMEM-10X) were added in a 5% CO_2 incubator at 38.5°C. The culture medium was then

131 replaced with hgDMEM + 10% FBS (GIBCO) + 1X antibiotic-antimycotic (GIBCO) (hgDMEM-

132 1X).

After 3 days of culture with hgDMEM-1X medium, the cells were serum starved overnight (24 h) with hgDMEM + 1X antibiotic-antimycotic. After serum starvation the cells were cultured in hgDMEM:M199 (GIBCO) (1:1) to which 10% FBS and 1X antibiotic-antimycotic (DM medium) were added until 60-65% confluency was reached. The cells were trypsinized and cultured to passage (P) 6 in Pericytes Growth Medium (PGM – Promocell, Heidelberg, Germany). The cells were expanded to the further passage when a 60-65% confluency was reached. Cell doubling and doubling time between passages were calculated as previously described (49).

140

141 Transcriptional Characterization of pAVPCs cultured in PGM

142 Cultured cells at P3 were transcriptionally analyzed through qPCR for mesenchymal stromal cells

143 (MSCs) (CD105, CD90, CD73, CD56, CD106), pericytes (neural/glial antigen 2 [NG2], Nestin,

144 platelet derived growth factor (PDGF) receptor β [PDGFR β], CD146, α -smooth muscle myosin

145 [αSMA]), hemopoietic (CD45, CD34) markers and for the gene expression of receptor and growth

146 factor as vascular endothelial growth factor (VEGF) receptor 1 (VEGFR1, also known as Flt1),

147 VEGFR2 (also known as Flk1), VEGF and PDGFB.

148 Total RNA was extracted from 2×10^6 cells using a NucleoSpin RNA kit (Macherey Nagel, Düren,

149 Germany) following the manufacturer's instructions. The extracted RNA was quantified using a

150 GeneQuant 1300 (GE Healthcare, Pittsburgh, PA) spectrophotometer, and an A260/A280 ratio was

151 used to evaluate RNA extraction quality. One microgram of RNA was retrotranscribed using an

152 iScript cDNA synthesis kit (Bio-Rad Laboratories Inc., California, USA), following the

153 manufacturer's instructions, in a 20µl final volume to obtain cDNA.

154 Primers were designed using Beacon Designer 2.07 Software (Premier Biosoft International, Palo

155 Alto, CA). Primers sequences and size (bp) of each product are listed in Table 1.

Quantitative PCR was carried out using a CFX96 (Bio-Rad) thermal cycler. A master mix of the
following reaction components was prepared in nuclease free water to the final concentrations
indicated: 0.2 µM forward primer, 0.2 µM reverse primer, 1X iTaq Universal SYBR Green
Supermix (Bio-Rad). One µl of cDNA were added to 19 µl of the master mix. All samples were
analysed in duplicate. The qPCR protocol used for the transcriptional characterization was: 10 min
at 95°C, 40 cycles at 95°C for 15 s and at 61°C for 30 s, followed by a melting step from 55°C to
95°C (80 cycle of 0.5°C increase/cycle).

163 The gene expression was evaluated using the ΔCq method (reference gene Cq – gene of interest

164 Cq). As Cq for the reference gene, the geometric mean of Cq of three different reference genes (β-

165 actin – β Act, Hypoxanthine phosphoribosyltransferase – HPRT, Glyceraldehyde 3-phosphate

166 dehydrogenase – GAPDH) was considered. Reference genes primers sequences and size (bp) of

167 each product are listed in Table 1.

168

169 Phenotypical Characterization of pAVPCs cultured in PGM

Phenotypical characterization of P3 cells was carried out through immunocytochemistry and flowcytometry following the same protocols described by us (49).

172 In particular, P3 cells were analyzed through flow cytometry for the expression of CD105, CD90,

173 CD56, CD44, CD45, CD34, CD31. The antibodies and the concentration used for the analysis are174 listed in Table 2.

175 Moreover, cells were analyzed by immunofluorescence experiments for the expression of PDGFβ,

176 αSMA, NG2, nestin, CD34, CD31 and smooth muscle myosin – heavy chain (SMM-hc). The

177 primary and secondary antibodies and their concentration used for the analysis are listed in Table 3.

179 Mesenchymal Trilineage Differentiation Potential

180 Cells at P3, grown in PGM, were assessed for the mesenchymal trilineage differentiation potential.

181 A StemPro® Adipogenesis Differentiation Kit, a StemPro® Osteogenesis Differentiation Kit and a

182 StemPro® Chondrogenesis Differentiation Kit (all purchased from GIBCO, Life Technologies)

183 were used following the manufacturer's instructions.

184 After 7 and 21 days for osteogenic differentiation and after 21 days for adipogenic and

185 chondrogenic differentiation the expression of typical genes of differentiated osteocytes (Alkaline

186 Phosphatase [ALP], Osteopontin [SSP1]), adipocytes (Peroxisome proliferator-activated receptor γ

187 [PPARγ], Adiponectin) and chondrocytes (Aggrecan [ACAN], Collagen type II alpha 1 [COL2A1])

188 was evaluated through qPCR. Total RNA extraction, retrotranscription and qPCR were performed

as described above. Primers for the analysis were designed with Beacon Designer 2.07 Software.

190 Primers sequences and size (bp) of each product are listed in Table 1. The expression of these genes

191 was calculated using the Δ Cq method (gene of interest Cq – reference gene Cq). As reference gene

192 Cq, the geometric mean of βAct, HPRT and GAPDH Cq value was considered. Relative expression

193 was calculated as fold of increase with the $2^{-\Delta\Delta Cq}$ method ($\Delta\Delta Cq = \Delta Cq$ differentiated cells – ΔCq

194 control cells).

After 21 days, assessment of differentiation was carried out through classical histological staining,
too, (OilRedO, Alizarin Red and Alcian Blue [all purchased by Sigma-Aldrich Co.]) following the

197 manufacturer's protocol.

198 Briefly, for chondrogenic differentiation assessment, pellets were fixed in 10% buffered formalin,

199 routinely processed and embedded in paraffin blocks. Sections of 3 µm thickness were obtained and

200 mounted on polarized slide. Sections were deparaffinized with xylene, rehydrated through passages

201 into an increasing concentration alcoholic ladder. Slides were stained with Alcian blue solution

202 (pH2.5) for 30 minutes, washed in water and counterstained with Harris hematoxylin (Merk).

204 Long term culture and assessing of smooth muscle cells differentiation potential

205 Tang and colleagues (42) reported that long term cultured vessel derived multipotent cells

spontaneously differentiate toward the smooth muscle cells phenotype. In order to verify the effect

207 of the long term culture on pAVPCs, 1500 cells/cm² cells at P3 were seeded in a 24 well plate and

208 cultured in DMEM + 10 % FBS + 1X antibiotic-antimycotic (Long Term Medium – LTM). Porcine

209 Aortic Precursor Cells grown in PGM has been used as control.

210 After 21 days, cells were analyzed for the expression of differentiated smooth muscle cells markers

both with qPCR (αSMA, calponin [CNN1], smooth muscle myosin heavy chain [SMM-hc]) and

212 immunofluorescence (αSMA, SMM-hc).

213 Quantitative PCR analysis and immunofluorescence staining were carried out as described above.

214 Primers for the analysis were designed with Beacon Designer 2.07 Software. Primers sequences and

size (bp) of each product are listed in Table 1. Relative expression was calculated as described

above. The primary and secondary antibodies and their concentration used for the analysis are listedin Table 3.

218

219 Endothelial Differentiation Potential

220 Cells at P3 were seeded in a 24 well plate at density of 5000 cells/cm² and cultured in human

endothelial Serum Free Medium (GIBCO) supplemented with 5% FBS, 1X antibiotic-antimycotic

and 50 ng/mL of human recombinant Vascular Endothelial Growth Factor (hVEGF) (Endothelial

223 Differentiation Medium - EDM). Porcine Aortic Precursor Cells grown in PGM has been used as

control. After 21 days of culture, cells were analyzed for the expression of differentiated endothelial

cells markers both with qPCR (CD31, vascular endothelial [VE]-Cadherin, vWF, endothelial nitric

oxide synthase [eNOS]) and immunofluorescence (CD31) as described above. Primers for the
analysis were designed with Beacon Designer 2.07 Software. Primers sequences and size (bp) of
each product are listed in Table 1. Relative expression was calculated as described above. The
primary and secondary antibody and their concentration used for the analysis are listed in Table 3.

230

231 Statistical analysis

Statistical analysis was carried out with R software (36). Data obtained from gene expression
analysis of adipogenic, chondrogenic, smooth muscle and endothelial differentiation were analyzed
using the Student-*t* test comparing differentiated and undifferentiated (control) cells, while data
obtained from gene expression analysis of osteogenic differentiation were analyzed using one-way
ANOVA, followed by the Tuckey post-hoc comparison, to detect differences among control cells
and cells differentiated for 7 and 21 days.

239 **Results and Discussion**

240

241 Morphology and doubling time of pAVPCs cultured in PGM

242 Porcine Aortic Precursor Cells in vitro cultured using the commercial medium Pericytes Growth 243 Medium (Promocell) showed a small cell body with little thin and elongated arms (Fig. 1 A and B). 244 This morphology has been already described for perivascular stem progenitor cells (8). The same 245 cells cultured in the unspecific medium DM showed a spindle shape/fibroblast-like morphology 246 (49) unlike if cultured in PGM. Moreover, it was not observed the growth of spheroidal structures in 247 PGM, while in the first passages of the culture in DM it was possible observing these kind of 248 structures (49). Doubling time has been calculated for cells cultured between P1 and P6 and 249 increase from 27.5 ± 0.6 hours, between P1 and P2, to 44.4 ± 10.9 hours, between P5 and P6 (Fig. 1 250 C). Interestingly, the growth curve obtained for cells cultured in PGM is parallel to the one obtained 251 for the culture in DM (49) with a downward shift of about 10 hours, so cells cultured in PGM grew 252 more rapidly than cultured in DM. Cells at P6 reached a cumulative cell doubling number of $11.1 \pm$ 253 1.3, comparable to the value (10.7 ± 0.9) obtained for culture in DM (49).

254

255 Transcriptional Characterization of pAVPCs cultured in PGM

256 Porcine Aortic Precursor Cells cultured in PGM express the main transcripts of MSCs (CD105,

CD90, CD73, CD56, CD106), while do not express CD45, a marker of the hemopoietic lineage (13)
(Fig 2).

259 Moreover, pAVPCs do express pericytes main transcripts (NG2, Nestin, CD146, αSMA, PDGFRβ) 260 (11) and CD34 mRNA, considered a markers of Advential Cells, Myoendothelial Cells (8) and mesangioblasts (31). 261

262 Transcripts of the growth factor VEGF and its main receptors (Flt-1 and Flk1) has been detected as 263 well as for the growth factor PDGFB, although at a lower level for the latter. Considering the 264 vascular origin of pAVPCs, VEGF expression could be an important factor for the cross-talk with 265 endothelial cells in physiological and pathological angiogenic processes that involve vascular 266 precursor cells (16).

267

269

268 Phenotypical Characterization of pAVPCs cultured in PGM

Passage 3 pAVPCs cultured in PGM has been characterized for MSC and pericytes markers 270 expression through immunocytochemistry and flow cytometry. Flow cytometry analysis (Fig. 3) 271 revealed that pAVPCs expressed MSC markers as CD105 (66.8 \pm 0.1 %), CD90 (99.5 \pm 0.2 %), 272 CD44 (99.6 \pm 0.3 %) and less than 2% of them expressed the hemopoietic lineage markers as CD45 273 $(1.4 \pm 0.4 \%)$, CD34 $(1.3 \pm 0.1 \%)$, as requested by the International Society of Cell Therapy (ISCT) 274 guide lines (13). Moreover, cells were negative for the expression of CD31($1.5 \pm 0.1 \%$). In 275 particular, as already described for cells cultured in DM, cells expressed CD56 (99.9 ± 0.1 %) that 276 is considered a marker of subsets of MSCs (2, 4, 37). The FACS analysis revealed the presence of a 277 more uniform population of cells when the *in vitro* culture is performed in PGM compared to the 278 culture in DM (49). 279 Immunocytochemical analysis revealed the expression of cell markers typical of pericytes (8, 11, 280 19, 26) as PDGFR^β, NG2 and Nestin (respectively Figure 4 A, B, C) as already described for the 281 culture in DM (49). A clear difference observed among cells cultured in different media was that in

PGM less than 2% of cells expressed aSMA (Figure 4 D), while in DM (49) 100% of cells 282

283 expressed it. The α SMA protein is considered a functional marker of differentiated pericytes (10)

and so it is presumably expressed in pre-committed cells to the fully differentiated pericytes

285 lineage. The observation that the cells cultured in PGM lost the expression of that marker could

indicate that the culture medium was able to maintain cells more undifferentiated than DM.

287 Despite the expression of CD34 mRNA, cells did not express that marker at the protein level (Fig. 4

E). Moreover, CD31 (Fig. 4 F) was not expressed, too, so both these data allowed us to exclude

289 contamination of hemopoietic and endothelial cells (10).

290 Taken together, these data suggested that pAVPCs in PGM could be phenotypically considered

291 pericyte-like cells, as they displayed several MCS and pericytes markers (8) confirming what it has

been observed in our previous work (49)

293

294 Mesenchymal Trilineage Differentiation Potential

295 Passage 3 cells, cultured in PGM, has been cultured in adipo-, osteo-, chondrogenesis induction

296 media in order to investigate the classical trilineage differentiation potential (Fig. 5) requested for

297 MSCs characterization (13) as already shown for culture in DM (49).

Quantitative PCR analysis revealed the over-expression of transcripts typical of each one of thethree lineages described above.

300 For adipogenic differentiation PPARy and Adiponectin has been used as markers of differentiation

301 (30). Transcripts of both genes has been detected significantly increased in differentiated cells

302 compared to control cells respectively about 525 times (p t-test = 6.20 x 10⁻⁵) and about 45000

303 times (p t-test = 0.003973) (Fig. 5 A).

304 For osteogenic differentiation ALPL and SPP1 has been used as markers of differentiation (28). In

305 Fig. 5 D the expression of both markers is represented at 7 and 21 days of cell culture in osteogenic

306 differentiation medium. The expression of ALPL was significantly increased in differentiated cells 307 (about 1300 times) compared to control only after 21 days of culture in osteogenic medium (p 308 ANOVA = 5.65×10^{-5}), while the expression of SPP1 was significantly increased in differentiated 309 cells (about 17 times) compared to control after just 7 days of culture in osteogenic medium (p 310 ANOVA = 0.00188).

For chondrogenic differentiation ACAN and COL2A1 has been used as markers of differentiation (48). Transcripts of both genes has been detected significantly increased in differentiated cells, compared to control cells respectively, about 585 times (p *t-test* = 0.001668) and about 11 times (p t-test = 0.03973) (Fig. 5 G).

Moreover, cells cultured with adipogenic medium for 21 days grew as monolayer showing the accumulation of lipidic droplets stained with OilRedO (Fig. 5 B, C), cells cultured with osteogenic medium for 21 days grew as aggregates and they positively stained for AlizarinRed confirming the presence of calcium deposits (Fig. 5 E, F) and cells, cultured as pellets in a chondrogenic medium for 21 days, positively stained for AlizanBlue confirming the presence of proteoglicans (Fig. 5 H, I).

321 All these data suggested that PGM was able to maintain pAVPCs undifferentiated as them

322 displayed the classical trilineage differentiation potential that is requested for MSCs

323 characterization (13) and that has been already showed by us for pAVPCs cultured using DM (49).

324 Moreover, these data confirm that pAVPCs could be considered MSC-like cells.

325

326 Long term culture and smooth muscle cells differentiation potential

328	In order to evaluate whether pAVPCs spontaneously differentiation towards smooth muscle
329	phenotype cells has been cultured in LTM and then analyzed through qPCR and ICC.
330	After 21 days of culture in LTM, pAVPCs showed an elonged fibroblast-like shape and seems to
331	organize in cord-like structures, while pAVPCs cultured in PGM grew as multilayer maintaining
332	their classical shape (Suppl. 1).
333	Quantification of smooth muscle cells markers through qPCR (Fig. 6 A) showed a sensible and
334	significant increment of each of the three transcripts (α SMA, CNN1, SMM-hc [47]) analysed in
335	LTM cultured cells compared to the control one. In particular, aSMA showed a significant (p <i>t-test</i>
336	= 0.01623) increment of about 12 times, while CNN1 showed a significant (p t -test = 0.0314)
337	increment of about 41 times and SMM-hc a significant (p t -test = 0.0002068) increment of about
338	966 times.

339 The presence of α SMA and SMM proteins has been investigated in long term cultured cells and the 340 results obtained confirmed the lack of expression of both protein in control cells (Fig. 6 B, D) and 341 the expression of both protein in LTM cultured cells (Fig. 6 C, E).

All these data confirm that cells cultured for long time in the same support with a standard culture
 medium spontaneously differentiate to the smooth muscle phenotype, without growth factor

344 stimulation, as reported by Tang and colleagues (42) for other vascular derived multipotent cells.

345

346 Endothelial Differentiation Potential

In order to assess whether pAVPCs were able to differentiate to endothelial cells, a stimulation with
50 ng/µL VEGF has been performed in a culture medium specific for endothelial cells culture. After
21 days of treatment, VEGF cultured cells displayed an endothelial cell-like morphology growing
as a monolayer upon which few spheroidal structures could be observed (Suppl. 2).

351 Gene expression analysis (Fig. 7 A) revealed the over-expression of the following typical markers 352 of endothelial cells: CD31, VE-Cadherin, vWF and eNOS. Respectively, a significant increase of 353 these transcripts of about 22 times (p t-test = 0.01756), 33 times (p t-test = 0.007046), 7 times (p t-354 test = 0.02367) and 20 times (p t-test = 0.0111) has been detected in VEGF cultured cells compared 355 to control cells. CD31 expression has been investigated at the protein level, too, and the marker has 356 been detected just in differentiated cells showing its typical distribution pattern, mainly along the 357 cell membrane (Fig. 7 C). Control cells did not express the CD31 protein (Fig. 7 B). 358 All these data confirm that pAVPCs cultured in human endothelial Serum Free Medium (GIBCO)

supplemented with Vascular Endothelial Growth Factor differentiate to the endothelial phenotype,
in agreement with results obtained for multipotent cells derived from embryonic, fetal and human
aorta (14, 22, 23, 31, 35) and by Pankajakshan et al (2013) that reported the differentiation of
porcine bone marrow derived-MSCs to endothelial cells after stimulation with 50 ng/µL VEGF
(34).

364

Mesenchymal stromal cells in which depletion of PDFGRβ signaling occur have been reported to
have an high angiogenic potential as they produced pro-angiogenic growth factor and expressed
endothelial cells marker *in vitro*, while *in vivo* they potently stimulate neo-vascularization (1).
Greenberg and colleagues (18) described that VEGF, activating its receptor VEGFR2, is able to
suppress PDGFRβ signaling in vascular smooth muscle cells (VSMCs) through the assembly of a
PDGFRβ/VEGFR2 complex.

371 Considering that pAVPCs have been defined as MSC-like cells (49), the expression on
372 endothelial/angiogenic markers in these cells, after the VEGF stimulation, could be explained
373 through the VEGF-mediated inhibition of the PDGFRβ signaling. That could be the first stimulus to
374 induce the endothelial differentiation ensuring that the main receptor (PDGFRβ) involved in

pericytes/VSMCs regulation and differentiation *in vivo* (18) is blocked as the pathway that
underwent its activation. Indeed, further investigations are necessary to confirm this hypothesis.

377

378 Conclusion

In the present paper we described the ability of porcine Aortic Vascular Precursor Cells (pAVPCs)
to differentiate toward the smooth muscle and the endothelial cell phenotypes. In our previous paper
(49) we described a method to isolate these multipotent cells from the tunica media of pig aorta.

382 In this paper we modified the previous protocol culturing cells in a specific culture medium able to

383 maintain pericytes multipotency (3).

384 In particular, in the present paper pAVPCs cultured in Pericytes Growth Medium has been shown to

be rather pure population of cells that express the main markers of MSCs (CD105, CD90, CD73,

386 CD44) and lack the expression of the main markers of hemopioetic stem cells (CD45 and CD34), as

387 requested by the ISCT (13). Moreover, pAVPCs has been shown to express the main markers

388 (PDGFR β , NG2 and α SMA) that characterized pericytes (11). In addition, pAVPCs have been

389 shown to be able to differentiate toward osteo-, adipo- and chondrocyte phenotype.

390 All the data obtained from PGM cultured pAVPCs characterization, associated to their already

391 proved capability to form capillary-like network if co-cultured with Human Umbilical Vein

392 Endothelial Cells (HUVEC) on extracellular matrix (49), lead us to reinforce the definition of them

as MSC/pericyte-like cells.

Based on the recent definition of Vascular Stem Cells (VSCs) by Lin and Lue we wanted, then, to assess whether pAVPCs could be considered a population of VSC-like cells, in particular, whether they were be able to differentiate toward the smooth muscle and the endothelial phenotypes (29).

397 The data we presented in this paper showed that pAVPCs are able to differentiate spontaneously to 398 the smooth muscle phenotype if long term cultured in an unspecific culture medium. This 399 spontaneous differentiation process could lead to think about these cells in a pathological fashion as 400 it has been reported that some populations of vascular wall resident cells are responsible for several 401 vascular pathologies (20, 21, 26) and for some of these cells the shift to the smooth muscle 402 phenotype is a requirement for their involvement in vascular remodeling and neointimal hyperplasia 403 (42). Moreover, in the present paper we presented data that support the endothelial differentiation of 404 pAVCPs as after 21 days of culture in a medium supplemented with VEGF they expressed markers 405 of endothelial cells. The endothelial differentiation of these cells, instead, could lead to think about 406 pAVPCs in a regenerative medicine fashion. In fact, for regenerative medicine purposes it is of considerable importance the improvement of the vascular network that could be damaged in the 407 408 organ that has to be regenerated (6, 12, 24).

Indeed, it is important to remember that cells, like the multipotent pericytes with which pAVPCs
share multiple features, have been recently identified as blood vessel wall resident cells that
physiologically make the vasculature a dynamic reservoir of stem/progenitor cells (5, 10, 44).

So, considering the recent definition of Vascular Stem Cells (VSCs) by Lin and Lue (29) and all the
data we presented on pAVPCs in this paper, we conclude that they can be defined as a population of
VSC-like cells considering that they express markers of MSCs, display the classical MSC trilineage
differentiation and differentiate, *in vitro*, toward smooth muscle and endothelial cells phenotypes.

416 Indeed, both smooth muscle and endothelial differentiations require to be further investigated in *in*

417 *vivo* animal models of pathologies in order to confirm the involvement of pAVPCs in vascular

418 diseases development and/or the possible usefulness of these cells for regenerative medicine studies419 in porcine animal model.

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427 Ball SG, Worthington JJ, Canfield AE, Merry CL, Kielty CM. Mesenchymal stromal 1. 428 cells: inhibiting PDGF receptors or depleting fibronectin induces mesodermal progenitors with endothelial potential. Stem Cells 2014;32(3):694-705. 429 430 2. Battula VL, Treml S, Bareiss PM, Gieseke F, Roelofs H, de Zwart P, Müller I, Schewe 431 B, Skutella T, Fibbe WE and others. Isolation of functionally distinct mesenchymal stem cell 432 subsets using antibodies against CD56, CD271, and mesenchymal stem cell antigen-1. Haematologica 2009;94(2):173-84. 433 434 3. Blocki A, Wang Y, Koch M, Peh P, Beyer S, Law P, Hui J, Raghunath M. Not All 435 MSCs Can Act as Pericytes: Functional In Vitro Assays to Distinguish Pericytes from Other 436 Mesenchymal Stem Cells in Angiogenesis. Stem Cells Dev 2013;22(17):2347-55. 437 4. Bühring HJ, Treml S, Cerabona F, de Zwart P, Kanz L, Sobiesiak M. Phenotypic 438 characterization of distinct human bone marrow-derived MSC subsets. Ann N Y Acad Sci 439 2009;1176:124-34. 440 5. Chen CW, Corselli M, Péault B, Huard J. Human blood-vessel-derived stem cells for tissue repair and regeneration. J Biomed Biotechnol 2012;2012:597439. 441 442 6. Chen CW, Okada M, Proto JD, Gao X, Sekiya N, Beckman SA, Corselli M, Crisan M, 443 Saparov A, Tobita K and others. Human pericytes for ischemic heart repair. Stem Cells 444 2013;31(2):305-16. 7. Corselli M, Chen CW, Crisan M, Lazzari L, Péault B. Perivascular ancestors of adult 445 multipotent stem cells. Arterioscler Thromb Vasc Biol 2010;30(6):1104-9. 446

8. Corselli M, Chen CW, Sun B, Yap S, Rubin JP, Péault B. The tunica adventitia of
human arteries and veins as a source of mesenchymal stem cells. Stem Cells Dev
2012;21(8):1299-308.

450 9. Corselli M, Crisan M, Murray IR, West CC, Scholes J, Codrea F, Khan N, Péault B.
451 Identification of perivascular mesenchymal stromal/stem cells by flow cytometry. Cytometry
452 A 2013;83(8):714-20.

453 10. Crisan M, Corselli M, Chen WC, Péault B. Perivascular cells for regenerative
454 medicine. J Cell Mol Med 2012;16(12):2851-60.

455 11. Crisan M, Yap S, Casteilla L, Chen CW, Corselli M, Park TS, Andriolo G, Sun B,

Zheng B, Zhang L and others. A perivascular origin for mesenchymal stem cells in multiple
human organs. Cell Stem Cell 2008;3(3):301-13.

458 12. Dar A, Domev H, Ben-Yosef O, Tzukerman M, Zeevi-Levin N, Novak A, Germanguz I,
459 Amit M, Itskovitz-Eldor J. Multipotent vasculogenic pericytes from human pluripotent stem
460 cells promote recovery of murine ischemic limb. Circulation 2012;125(1):87-99.

461 13. Dominici M, Le Blanc K, Mueller I, Slaper-Cortenbach I, Marini F, Krause D, Deans
462 R, Keating A, Prockop D, Horwitz E. Minimal criteria for defining multipotent mesenchymal
463 stromal cells. The International Society for Cellular Therapy position statement. Cytotherapy
464 2006;8(4):315-7.

465 14. Fang B, Li Y, Song Y, Li N. Isolation and characterization of multipotent progenitor
466 cells from the human fetal aorta wall. Exp Biol Med (Maywood) 2010;235(1):130-8.

467 15. Forni M, Mazzola S, Ribeiro LA, Pirrone F, Zannoni A, Bernardini C, Bacci ML,

468 Albertini M. Expression of endothelin-1 system in a pig model of endotoxic shock. Regul Pept
469 2005;131(1-3):89-96.

470	16. Geevarghese A, Herman IM. Pericyte-endothelial crosstalk: implications and
471	opportunities for advanced cellular therapies. Transl Res 2014;163(4):296-306.
472	17. Gessaroli M, Bombardi C, Giunti M, Bacci ML. Prevention of neointimal hyperplasia
473	associated with modified stretch expanded polytetrafluoroethylene hemodialysis grafts (Gore)
474	in an experimental preclinical study in swine. J Vasc Surg 2012;55(1):192-202.
475	18. Greenberg JI, Shields DJ, Barillas SG, Acevedo LM, Murphy E, Huang J, Scheppke L,
476	Stockmann C, Johnson RS, Angle N and others. A role for VEGF as a negative regulator of
477	pericyte function and vessel maturation. Nature 2008;456(7223):809-13.
478	19. Howson KM, Aplin AC, Gelati M, Alessandri G, Parati EA, Nicosia RF. The postnatal
479	rat aorta contains pericyte progenitor cells that form spheroidal colonies in suspension
480	culture. Am J Physiol Cell Physiol 2005;289(6):C1396-407.
481	20. Hu Y, Xu Q. Adventitial biology: differentiation and function. Arterioscler Thromb
482	Vasc Biol 2011;31(7):1523-9.
483	21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant
483 484	21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient
483 484 485	21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient mice. J Clin Invest 2004;113(9):1258-65.
483 484 485 486	 Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient mice. J Clin Invest 2004;113(9):1258-65. Invernici G, Emanueli C, Madeddu P, Cristini S, Gadau S, Benetti A, Ciusani E, Stassi
 483 484 485 486 487 	 21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient mice. J Clin Invest 2004;113(9):1258-65. 22. Invernici G, Emanueli C, Madeddu P, Cristini S, Gadau S, Benetti A, Ciusani E, Stassi G, Siragusa M, Nicosia R and others. Human fetal aorta contains vascular progenitor cells
 483 484 485 486 487 488 	 21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient mice. J Clin Invest 2004;113(9):1258-65. 22. Invernici G, Emanueli C, Madeddu P, Cristini S, Gadau S, Benetti A, Ciusani E, Stassi G, Siragusa M, Nicosia R and others. Human fetal aorta contains vascular progenitor cells capable of inducing vasculogenesis, angiogenesis, and myogenesis in vitro and in a murine
 483 484 485 486 487 488 489 	 21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient mice. J Clin Invest 2004;113(9):1258-65. 22. Invernici G, Emanueli C, Madeddu P, Cristini S, Gadau S, Benetti A, Ciusani E, Stassi G, Siragusa M, Nicosia R and others. Human fetal aorta contains vascular progenitor cells capable of inducing vasculogenesis, angiogenesis, and myogenesis in vitro and in a murine model of peripheral ischemia. Am J Pathol 2007;170(6):1879-92.
 483 484 485 486 487 488 489 490 	 21. Hu Y, Zhang Z, Torsney E, Afzal AR, Davison F, Metzler B, Xu Q. Abundant progenitor cells in the adventitia contribute to atherosclerosis of vein grafts in ApoE-deficient mice. J Clin Invest 2004;113(9):1258-65. 22. Invernici G, Emanueli C, Madeddu P, Cristini S, Gadau S, Benetti A, Ciusani E, Stassi G, Siragusa M, Nicosia R and others. Human fetal aorta contains vascular progenitor cells capable of inducing vasculogenesis, angiogenesis, and myogenesis in vitro and in a murine model of peripheral ischemia. Am J Pathol 2007;170(6):1879-92. 23. Invernici G, Madeddu P, Emanueli C, Parati EA, Alessandri G. Human fetal aorta-

492 Cytotechnology 2008;58(1):43-7.

493 24. James AW, Zara JN, Corselli M, Chiang M, Yuan W, Nguyen V, Askarinam A, Goyal
494 R, Siu RK, Scott V and others. Use of human perivascular stem cells for bone regeneration. J
495 Vis Exp 2012(63):e2952.

496 25. Janeczek Portalska K, Leferink A, Groen N, Fernandes H, Moroni L, van Blitterswijk
497 C, de Boer J. Endothelial differentiation of mesenchymal stromal cells. PLoS One
498 2012;7(10):e46842.

499 26. Juchem G, Weiss DR, Gansera B, Kemkes BM, Mueller-Hoecker J, Nees S. Pericytes
500 in the macrovascular intima: possible physiological and pathogenetic impact. Am J Physiol
501 Heart Circ Physiol 2010;298(3):H754-70.

502 27. Krause U, Harter C, Seckinger A, Wolf D, Reinhard A, Bea F, Dengler T, Hardt S, Ho
503 A, Katus HA and others. Intravenous delivery of autologous mesenchymal stem cells limits
504 infarct size and improves left ventricular function in the infarcted porcine heart. Stem Cells
505 Dev 2007;16(1):31-7.

506 28. Kulterer B, Friedl G, Jandrositz A, Sanchez-Cabo F, Prokesch A, Paar C, Scheideler
507 M, Windhager R, Preisegger KH, Trajanoski Z. Gene expression profiling of human
508 mesenchymal stem cells derived from bone marrow during expansion and osteoblast
509 differentiation. BMC Genomics 2007;8:70.

510 29. Lin CS, Lue TF. Defining vascular stem cells. Stem Cells Dev 2013;22(7):1018-26.

30. Menssen A, Häupl T, Sittinger M, Delorme B, Charbord P, Ringe J. Differential gene
expression profiling of human bone marrow-derived mesenchymal stem cells during
adipogenic development. BMC Genomics 2011;12:461.

514 31. Minasi MG, Riminucci M, De Angelis L, Borello U, Berarducci B, Innocenzi A,
515 Caprioli A, Sirabella D, Baiocchi M, De Maria R and others. The meso-angioblast: a

multipotent, self-renewing cell that originates from the dorsal aorta and differentiates into
most mesodermal tissues. Development 2002;129(11):2773-83.

32. Oswald J, Boxberger S, Jørgensen B, Feldmann S, Ehninger G, Bornhäuser M,
Werner C. Mesenchymal stem cells can be differentiated into endothelial cells in vitro. Stem
Cells 2004;22(3):377-84.

33. Pacilli A, Pasquinelli G. Vascular wall resident progenitor cells: a review. Exp Cell Res
2009;315(6):901-14.

34. Pankajakshan D, Kansal V, Agrawal DK. In vitro differentiation of bone marrow
derived porcine mesenchymal stem cells to endothelial cells. J Tissue Eng Regen Med
2013;7(11):911-20.

35. Pasquinelli G, Tazzari PL, Vaselli C, Foroni L, Buzzi M, Storci G, Alviano F, Ricci F,
Bonafè M, Orrico C and others. Thoracic aortas from multiorgan donors are suitable for
obtaining resident angiogenic mesenchymal stromal cells. Stem Cells 2007;25(7):1627-34.

529 36. R Team. R: A language and environment for statistical computing. Vienna, Austria: R
530 Foundation for Statistical Computing,; 2008.

37. Rossignoli F, Caselli A, Grisendi G, Piccinno S, Burns JS, Murgia A, Veronesi E,
Loschi P, Masini C, Conte P and others. Isolation, characterization, and transduction of
endometrial decidual tissue multipotent mesenchymal stromal/stem cells from menstrual
blood. Biomed Res Int 2013;2013:901821.

535 38. Sacchetti B, Funari A, Michienzi S, Di Cesare S, Piersanti S, Saggio I, Tagliafico E,

536 Ferrari S, Robey PG, Riminucci M and others. Self-renewing osteoprogenitors in bone

537 marrow sinusoids can organize a hematopoietic microenvironment. Cell 2007;131(2):324-36.

538	39.	Sato T, Iso Y, Uyama T, Kawachi K, Wakabayashi K, Omori Y, Soda T, Shoji M,
539	Koba	a S, Yokoyama S and others. Coronary vein infusion of multipotent stromal cells from
540	bone	marrow preserves cardiac function in swine ischemic cardiomyopathy via enhanced
541	neova	ascularization. Lab Invest 2011;91(4):553-64.

542 40. Shi S, Gronthos S. Perivascular niche of postnatal mesenchymal stem cells in human
543 bone marrow and dental pulp. J Bone Miner Res 2003;18(4):696-704.

544 41. Tang W, Zeve D, Suh JM, Bosnakovski D, Kyba M, Hammer RE, Tallquist MD, Graff
545 JM. White fat progenitor cells reside in the adipose vasculature. Science 2008;322(5901):583546 6.

547 42. Tang Z, Wang A, Yuan F, Yan Z, Liu B, Chu JS, Helms JA, Li S. Differentiation of
548 multipotent vascular stem cells contributes to vascular diseases. Nat Commun 2012;3:875.

549 43. Tavazoie M, Van der Veken L, Silva-Vargas V, Louissaint M, Colonna L, Zaidi B,

550 Garcia-Verdugo JM, Doetsch F. A specialized vascular niche for adult neural stem cells. Cell
551 Stem Cell 2008;3(3):279-88.

44. Tavian M, Zheng B, Oberlin E, Crisan M, Sun B, Huard J, Peault B. The vascular wall
as a source of stem cells. Ann N Y Acad Sci 2005;1044:41-50.

554 45. Tintut Y, Alfonso Z, Saini T, Radcliff K, Watson K, Boström K, Demer LL.

555 Multilineage potential of cells from the artery wall. Circulation 2003;108(20):2505-10.

46. Vilahur G, Padro T, Badimon L. Atherosclerosis and thrombosis: insights from large
animal models. J Biomed Biotechnol 2011;2011:907575.

558 47. Xie C, Ritchie RP, Huang H, Zhang J, Chen YE. Smooth muscle cell differentiation in

559 vitro: models and underlying molecular mechanisms. Arterioscler Thromb Vasc Biol

2011;31(7):1485-94.

48. Xu J, Wang W, Ludeman M, Cheng K, Hayami T, Lotz JC, Kapila S. Chondrogenic
differentiation of human mesenchymal stem cells in three-dimensional alginate gels. Tissue
Eng Part A 2008;14(5):667-80.

564 49. Zaniboni A, Bernardini C, Alessandri M, Mangano C, Zannoni A, Bianchi F, Sarli G,

565 Calzà L, Bacci ML, Forni M. Cells derived from porcine aorta tunica media show

566 mesenchymal stromal-like cell properties in in vitro culture. Am J Physiol Cell Physiol
567 2014;306(4):C322-33.

568 50. Zannoni A, Giunti M, Bernardini C, Gentilini F, Zaniboni A, Bacci ML, Forni M.

569 Procalcitonin gene expression after LPS stimulation in the porcine animal model. Res Vet Sci
570 2012;93(2):921-7.

571 51. Zaragoza C, Gomez-Guerrero C, Martin-Ventura JL, Blanco-Colio L, Lavin B,

572 Mallavia B, Tarin C, Mas S, Ortiz A, Egido J. Animal models of cardiovascular diseases. J
573 Biomed Biotechnol 2011;2011:497841.

574 52. Zheng B, Cao B, Crisan M, Sun B, Li G, Logar A, Yap S, Pollett JB, Drowley L,

575 Cassino T and others. Prospective identification of myogenic endothelial cells in human

576 skeletal muscle. Nat Biotechnol 2007;25(9):1025-34.

577 53. Zheng B, Li G, Chen WC, Deasy BM, Pollett JB, Sun B, Drowley L, Gharaibeh B, Usas

578 A, Péault B and others. Human myogenic endothelial cells exhibit chondrogenic and

579 osteogenic potentials at the clonal level. J Orthop Res 2013;31(7):1089-95.

Genes	Accession Number	Forward (5'-3')	Reverse (5'-3')	Product size (bp)	
Mesenchymai	Stromal Cell markers				
CD106	NM_213891	GAGGATGGAAGATTCTGGAATTTACG	ATCACTAGAGCAGGTCATGTTCAC	172	
CD105	NM_214031.1	ATACAAAGGGCTCCATCATC	TGAGTGTGAGACTTCCATTC	151	
CD90	NM_001146129.1	GACTGCCGCCATGAGAATAC	GGTAGTGAAGCCTGATAAGTAGAG	180	
CD73	XM_003353250.1	AACTCATCGCTCAGAAGGTG	ATCGGAGGTGACTATGAATGG	131	
CD56	ENSSSCT00000016411	GGAAATCAGCGTTGGAGAGTC	TGTTGGCATTGTAGATGGTGAG	172	
Pericytes Ma	rkers				
CD146	ENSSSCT00000016482	CTCATCTGTGCCTTCCTGCTAG	CCTCCCACTTCCACCTCCAG	110	
NG2	ENSSSCT0000002098	ACCACCTCCTCCTACAACTC	GTCACTCAGCAGCATCTCTG	104	
PDGFRβ	ENSSSCT00000015788	GCAACGAGGTGGTCAACTTC	GCAGGATAGAACGGATGTGG	111	
Nestin	ENSSSCT00000027298	CAGTGGTTCCAAGGCTTCTC	CATAGGTGTGTCAAGGGTATCG	163	
Hemopoietic	Stem Cell Markers				
CD45	XM_003482796.1	CTCACTCGCAAGCATCTCTG	CGGTTGAAGTAGCTGTGTCTG	188	
CD34	NM_214086	CAGCCTCCACACTGTCTCTC	CACCTCTGAAGTTGTAAAGTTGATG	184	
Smooth Musc	le Cell Markers				
αSMA	NM_001164650	CACGGCATCATCACCAACTG	ACCGCCTGAATAGCCACATAC	200	
CNN1		ATGTCCTCTGCTCACTTCAAC	CCTGCTGGTGGTCATACTTC	91	
SMM-hc	XR_131283	AGGAGGTGACGATGCTGAAG	TGCTTGTTCTTGTCCAGGTTG	160	
Endothelial C	Cell Markers				
CD31	NM 213907	CTCATCGCAGTGGTTGTCATC	TGTCTTCTCATTGTTGGAGTTCAG	150	
VE-	- NM 001001649	ACTECTEAGECTETEETEAG	GTCAACACTCAGCACAGCATAG	111	
Cadherin	NM_001246221			102	
ONOS	NM_001240221	CTCTCACCTTCTTCCTC	CACTTCCACTCCTCATAG	144	
Paganton and	Crowth Easter	Generacementerio	CLACTICLACTICICATAO	144	
кесеріог ини	Growin Facior				
Flt1	AY566244	TTGGACTGTTGGCACAAAGAC	GCTGTTGCTCGTCAGAATGG	141	
Flk1	AJ245446	AACGAGTGGAGGTGACAGATTG	CGGGTAGAAGCACTTGTAGGC	104	
VEGF	AF318502	CCTTGCCTTGCTGCTCTACC	CGTCCATGAACTTCACCACTTC	101	
PDGFβ	ENSSSCT00000029224	CTCTGGCTGCTGCAACAACC	TGGCTTCTTCCGCACAATCTC	100	
Adipocyte Ma	ırkers				
PPARγ	AF103946	AGGAAAGACCACAGACAAATCAC	CAGGGATGTTCTTGGCATACTC	200	
Adiponectin	AY135647	GCGAGAAGGGTGAGAAAGG	ACAGTGACCCGAGTCTCCAG	187	
Osteocyte Ma	urkers				
ALP	XM_003361247	GCAAGCAGCACTCTCACTATATC	TCCACCAGCAAGAAGAAGCC	211	
SPP1	NM_214023	AAACAAGAGACCCTGCCAAG	TCATCGGATTCATCGGAGTG	173	
Chondrocyte	Markers				
ACAN	ENSSSCT0000002052	CCATCATCGCCACACCAGAG	CCCGTAGCAACCTTCCCTTG	128	
COL2A	ENSSSCT00000031054	CCTGGTGATGATGGTGAAGC	ACCTGGGTAACCTCTGTGAC	132	
Reference Genes					
GAPDH	AF017079	TGGTGAAGGTCGGAGTGAAC	TGTAGTGGAGGTCAATGAAGGG	120	
HPRT	AF143818	GGACAGGACTGAACGGCTTG	GTAATCCAGCAGGTCAGCAAAG	115	
βAct	AJ312193	GTCCTCCTCCTCCTGG	GTGGTCTCGTGGATGCC	141	

Table 1: List of primer pairs,
amplicon size (bp) and genesP. numberSpeciesSupplierDilution

analyzed in the paper.Antibody

Primary				
CD45-APC	K252-1E4	mouse	AbD Serotec	$10\mu l/10^6$ cells
CD90-APC	Ab139364	mouse	Abcam	$10\mu l/10^6$ cells
CD105-FITC	Ab53318	mouse	Abcam	$20\mu l/10^6$ cells
CD56-PE	304606	mouse	Biolegend	$10\mu l/10^6$ cells
CD44-PerPC	103036	rat	Biolegend	$10\mu l/10^6$ cells
CD34 unconjugated	Ab81289	rabbit	Abcam	1:60
CD31 unconjugated	MCA1746	mouse	AbD Serotec	1:100
Secondary				
Anti Rabbit-PE	Ab97070	goat	Abcam	1:200
Anti Mouse-FITC	420-120-05	sheep	BioFX	1:100

Table 2: Flow cytometry antibodies list used for the immunophenotyping of cells.

Antibody	P. number	Species	Supplier	Dilution
Primary				
NG2	AB-11160	rabbit	Immunological Sciences	1:200
Smooth Muscle Actin (aSMA)	1A4	mouse	Cell Marque	1:500
Nestin	AB5922	rabbit	Chemicon	1:150
PDGFR-b	#3169	rabbit	Santa Cruz	1:100
PECAM-1	sc-1506	goat	Santa Cruz	1:150
CD34 unconjugated	Ab81289	rabbit	Abcam	1:60
SMM-hc	MAB3570	mouse	Chemicon	1:100
Secondary				
Anti mouse RRX	715-295-151	Donkey	Jackson ImmunoResearch	1:100
Anti goat RRX	705-295-147	Donkey	Jackson ImmunoResearch	1:100
Anti rabbit Alexa Fluor 488	A21206	Donkey	Molecular Probes	1:600
Anti mouse FITC	F4143	Goat	Sigma Aldrich	1:800

Table 3: Immunocytochemistry antibodies list used for the cells characterization.

588 Figure 1



590 Porcine Aortic Vascular Precursor Cells (pAVPCs) morphology and doubling time during 591 culture in Pericytes Growth Medium (PGM). A, B: Cells cultured in PGM, after isolation and 592 starvation step, displayed a small cell body and elongated thin arms at their ends. In A it is shown a 593 microphotograph (10X magnification) of a low confluence culture of pAVPCs, while in B it is 594 shown a microphotograph (10X magnification) of a 60% confluence culture of pAVPCs. C: The 595 growth curve of pAVPCs cultured in PGM is represented in the picture. Cells displayed an 596 increasing replication time between P1 and P6.





- *derived growth factor* β). Data are expressed as Δ Cq calculated as Cq value obtained from the
- 611 geometric mean of the reference genes minus Cq value of the gene of interest.







Immunocytochemical characterization of pAVPC cultured in PGM. In each microphotograph
(scale bar 20 micron), P3 pAVPCs cultured in PGM were stained with different antisera; nuclei
were always stained with Hoechst 33258 (blue). A: PDGFRβ; B: NG2; C: Nestin; D: αSMA; E:
CD34; F: CD31.



628 **Trilineage differentiation potential of pAVPCs.** Porcine Aortic Precursor Cells are able to

629 differentiate toward adipo-, osteo- and chondrocyte phenotype if opportunely stimulated *in vitro*. A,

- 630 **D**, **G**: gene expression analysis of genes, respectively, of adipocytes (PPARγ, Adiponectin),
- 631 osteocytes (ALPL, SPP1) and chondrocytes (ACAN, COL2A1) in differentiated pAVPCs. In the Y-
- 632 axis in each graph is represented the relative expression of each transcripts analysed in
- 633 differentiated pAVPCs compared to the control ($2^{-\Delta\Delta Cq}$ method). For adipogenic differentiation (A)
- and chondrogenic differentiation (G) gene expression has been evaluated after 21 days of culture in

635 differentiation media while for osteogenic differentiation (**D**) gene expression has been evaluated

after 7 and 21 days of culture in differentiation medium. Data obtained for every single gene has

637 been statistically analyzed (comparing control and differentiated samples) for adipogenic

638 differentiation (A) and chondrogenic differentiation (G) through Student-t test (*: p<0.05; **:

p<0.01; ***: p<0.001), while for osteogenic differentiation (**D**) through one-way ANOVA followed

640 by Tuckey post-hoc comparison (different letter correspond to statistically different sample).

641 **B**, **C**: OilRedO staining (40X magnification) showed the presence of lipid droplets (red) in the

642 cytoplasm of pAVPCs cultured for 21 days in adipogenesis induction medium (C), while no lipid

643 droplets has been observed in control cells (**B**).

644 **E**, **F**: AlizarinRed staining (10X magnification) showed calcium rich-deposits (red) in pAVPCs

645 cultured for 21 days in osteogenesis induction medium that grew as spheroidal aggregates (F),

646 while no calcium deposits has been observed in control cells (**E**).

H, I: Alcian blue staining of cross sections of pAVPCs pellet. Blue staining of the extracellular
matrix, indicating presence of proteoglycans and suggesting differentiation toward the chondrocyte
phenotype, was present in differentiated pellet (I) and absent in controls (H). Samples has been
counterstained with hematoxylin. Scale bar = 100 μm.

652 **Figure 6**



654

Smooth muscle lineage differentiation potential of pAVPCs. Porcine Aortic Precursor Cells are 655 656 able to differentiate toward smooth muscle cell phenotype if long term cultured in vitro. A: gene 657 expression analysis of smooth muscle cell genes (aSMA, calponin [CNN1] and smooth muscle 658 myosin heavy chain [SMM-hc]) in long term cultured pAVPCs. In the Y-axis in the graph is 659 represented the relative expression of each transcripts analyzed in pAVPCs cultured in LTM compared to control cells ($2^{-\Delta\Delta Cq}$ method). Gene expression has been evaluated after 21 days of long 660 661 term culture in LTM and data obtained for every single gene has been statistically analyzed (comparing control and differentiated samples) through Student-t test (*: p<0.05; ***: p<0.001) 662 **B**, **C**: αSMA immunostaining, respectively of control and long term cultured cells (40 X 663 magnification). **D**, **E**: smooth muscle myosin-heavy chain (SMM-hc) immunostaining, respectively 664 665 of control and long term cultured cells (40 X magnification).Nuclei has been stained with Hoechst 666 33258 (blue).



670 Endothelial lineage differentiation potential of pAVPC. Porcine Aortic Precursor Cells are able 671 to differentiate toward the endothelial cell phenotype if *in vitro* cultured in an endothelial cell 672 medium supplemented with 50 ng/µL VEGF. A: gene expression analysis of endothelial cell genes 673 (CD31, VE-Cadherin, von Willebrand Factor [vWF] and endothelial Nitric Oxide Synthase 674 [eNOS])in differentiated pAVPCs. In the Y-axis in the graph is represented the relative expression 675 of each transcripts analyzed in pAVPCs cultured in EDM compared to control cells ($2^{-\Delta\Delta Cq}$ method). Gene expression has been evaluated after 21 days 676 677 of culture in EDM and data obtained for every single gene has been statistically analyzed (comparing control and differentiated samples) through Student-t test (*: p<0.05; **: p<0.01) 678 679 **B**, **C**: CD31 immunostaining, respectively of control and EDM cultured cells (scale bar 20 micron).

680 Nuclei has been stained with Hoechst 33258 (blue).

682 Supplementary Figure 1



683



morphology. A, C, E: pAVPC grew in Pericyte Growth Medium (Control) at 7, 14 and 21 days,
respectively; pAVPCs cultured in PGM grew as multilayer maintaining their classical shape with
little thin and elongated arms. B, D, F; pAVPC grew in Long Term Medium at 7, 14 and 21 days,

- respectively; pAVPC cultured in LTM showed an elongated fibroblast-like shape and seems to be
- 689 organized in cord-like structure. Magnification: 10X.

691 Supplementary Figure 2



692

693 Endothelial Differentiation Potential of pAVPC - morphology. A, C, E: pAVPC grew in

694 Pericyte Growth Medium (Control) at 7, 14 and 21 days, respectively; pAVPCs cultured in PGM

695 grew as multilayer maintaining their classical shape. **B**, **D**, **F**: pAVPC grew in Endothelial

Differentiation Medium at 7, 14 and 21 days, respectively; pAVPC stimulated with EDM displayed

- an endothelial cell-like morphology growing in cluster and forming an endothelial cell-like
- 698 monolayer upon which some cells organized in few spheroidal structures. Magnification: 10X.