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Muscle satellite cells and endothelial cells: close neighbours and privileged partners.

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Genetically engineered mice (Myf5^{nLacZ/+}, Myf5^{GFP-P/+}) allowing direct muscle satellite cell (SC) visualization indicate that, in addition to being located beneath myofiber basal laminae, SCs are strikingly close to capillaries. After GFP$^+$ bone marrow transplantation, blood-borne cells occupying SC niches previously depleted by irradiation, were similarly detected near vessels, thereby corroborating the anatomic stability of juxtavascular SC niches. BrdU pulse-chase experiments also localise quiescent and less quiescent SCs near vessels. SCs, and to a lesser extent myonuclei, were non-randomly associated with capillaries in humans. Significantly, they were correlated with capillarisation of myofibers, regardless to their type, in normal muscle. They also varied in paradigmatic physiologic and pathologic situations associated with variations of capillary density, including amyopathic dermatomyositis, a unique condition in which muscle capillary loss occurs without myofiber damage, and in athletes, in whom capillaries increase in number. Endothelial cell (EC) cultures specifically enhanced SC growth, through IGF-1, HGF, bFGF, PDGF-BB and VEGF, and, accordingly, cycling SCs remained mainly juxtavascular. Conversely, differentiating myogenic cells were both proangiogenic in vitro and spatiotemporally associated with neoangiogenesis in muscular dystrophy. Thus, SCs are largely juxtavascular and reciprocally interact with ECs during differentiation to support angio-myogenesis.

Muscle tissue repair is a complex biological process that crucially involves activation of stem cells. Skeletal muscle contains two different stem cell types: (i) myogenic stem cells, so-called satellite cells (SCs), that reside beneath the basal lamina of muscle fibers (Mauro, 1961) and express both NCAM/CD56 and early myogenic cell markers such as M-cadherin, Pax7 and Myf5 or Myf5^{nlacZ} (Bischoff and Franzini-Armstrong, 2004;
Hawke and Garry, 2001; Beauchamp et al., 2000; Seale et al., 2000); (ii) interstitial multipotent stem cells, that are extralaminal, exhibit fibroblastic morphology and do not express myogenic markers (Asakura et al., 2001; Tamaki et al., 2002). SCs are primarily quiescent in skeletal muscle, can self-renew (Collins et al., 2005) and, upon activation, proliferate and further differentiate to become fusion-competent myoblasts and ensure muscle regeneration (Bischoff and Franzini-Armstrong, 2004; Hawke and Garry, 2001). Interstitial "muscle-derived" stem cells give rise to several lineages after transplantation, and, in this setting, contribute to synchronized reconstitution of blood vessels (pericytes, smooth muscle cells, and endothelial cells), peripheral nerve (Schwann cells), and muscle cells (myofibers, and SCs) (Tamaki et al., 2005). However, participation of multipotent interstitial stem cells in physiologic muscle repair appears to be limited. Instead, it is widely accepted that sublaminal SCs represent the pre-eminent muscle stem cell type used for muscle growth, repair and regeneration (Dhawan & Rando, 2006).

Understanding how stem cell niches are organized in vivo and what interactions their progeny develop with neighbouring cell types is a critical issue in stem cell biology (Suda et al., 2005). The sublaminar location of SCs led to their identification, but little is known about the anatomical organization of the surroundings and the constituents of the niche. Moreover, signals that are likely conferred to myogenic cells by either direct contacts with the adjacent myofiber or soluble factors released by neighbouring non-muscle cells, constitute one of the major unexplored areas of SC biology (Dhawan & Rando, 2006). It seems plausible that cell-cell contacts between myofibers and SCs play a role in SC maintenance in a quiescence state (Dhawan & Rando, 2006). Such quiescent stem cells anchored in their niche exhibit a low requirement for growth factors (Suda et al., 2005).

Besides the quiescent stem cell niche, a maturation compartment may exist, involving progenitor cells and supportive stromal cells (Suda et al., 2005). We previously showed that activated SCs crucially interact with macrophages recruited at the site of muscle regeneration, and receive mitogenic signals from macrophages,
mediated by the sequential release of different soluble factors (Chazaud, 2003b). Myoblasts, and to a higher extent myotubes, also receive cell-contact-mediated pro-survival signals from macrophages (Sonnet et al., 2006).

In the present study, we focussed on the microvascular bed as another partner of SCs. Endothelial cells and myogenic cells may derive from common progenitors at development stages (Kardon, 2002) and vascular endothelial progenitor cells are probably essential for muscle organogenesis (Solursh 1987, Le Grand 2004). Moreover, different types of adult stem/progenitor cells crucially interact with the microvascular bed and the term of vascular niche was coined to define this compartment in the bone-marrow (Suda et al., 2005) and the hippocampus (Palmer et al., 2000). In bone marrow, the vascular niche is anatomically distinct from the endosteal niche of quiescent stem cells and is required for proliferation and terminal differentiation of hematopoietic progenitor cells (Suda et al., 2005). In the hippocampus, neural stem cells reside and self renew near fine capillaries, receive EC instructive cues, and form neurovascular units involved in an intimate combination of neurogenesis and angiogenesis (Palmer et al., 2000, Shen et al., 2004). In both cases, stem/progenitor cells were shown to secrete and respond to angiogenic factors (Tordjman et al., 2001; Maurer et al., 2003).

The close association of SCs to microvessels is intriguing (Schmalbruch and Hellhammer, 1977; Chazaud et al., 2003b), but it has been ignored or not acknowledged (Bischoff and Franzini-Armstrong, 2004; Carpenter and Karpati, 2001). In the present study, we documented this association using genetically engineered mice allowing SC visualization, and two methods conventionally used to mark stem cell niches in vivo (Lie and Xie, 2005), including bone marrow transplantation and label retention studies. Furthermore, we examined human muscle at steady state and in paradigmatic situations of microvascular loss or increase, to assess the interdependence of SCs and capillaries. Finally, coculture experiments, angiogenesis tests, protein and mRNA array-based detection of growth factors, and functional tests with blocking antibodies allowed us to establish that endothelial cells specifically enhance satellite cell
growth, and that, in turn, differentiating myogenic cells are proangiogenic.

These findings suggest that quiescent satellite cells, tightly associated with the myofiber in their sublaminal niche, are pre-positioned near capillaries and can, therefore, easily interplay with endothelial cells upon activation to set up coordinated angio-myogenesis in a functional manner.

RESULTS

Genetically engineered mice reveal juxtavascular location of most SCs

Neither electron microscopy nor teased fiber preparations are appropriate for the study of the spatial relationships between SCs and capillaries. Because immunocytochemical detection of mouse SCs is suboptimal, we used genetically engineered mice to visualize SCs in tibialis anterior (TA) muscle cryosections. We first used the heterozygous *Myf5*^{nlacZ/+} mouse, which has a reporter gene encoding nuclear-localizing β-Galactosidase (β-Gal) targeted to the *Myf5* locus such that expression of endogenous Myf5 is reported by β-Gal activity, thus allowing detection of SC nuclei (Tajbakhsh et al., 1996; Beauchamp et al., 2000). SCs appeared strikingly associated with capillaries when β-gal activity revelation was combined with either immunohistochemistry for basal lamina-specific collagen IV (Fig. 1A) or histoenzymatic reaction for microvascular alkaline phosphatase activity (Fig. 1B). Then we used the *Myf5*^{GFP-P/+} mouse which has a reporter gene encoding cytoplasmic green fluorescent protein (GFP) targeted to the *Myf5* locus (Kassar-Duchossoy et al., 2004). For unknown reasons, this mouse allows detection of a smaller proportion of SC than the *Myf5*^{nlacZ/+} mouse, but offers the advantage of visualizing SC cytoplasmic processes. Cryosections immunostained for laminin 1 showed that the mean distance separating GFP⁺ SCs (n=100) from capillaries was 2.6± 3.3 µm: 82% of SCs were at <5 µm from endothelial cells (ECs), 12% at 5-10 µm, and 7% at >10 µm (Fig. 1C).

It has been suggested that SCs form an heterogeneous population (Schultz and Lipton 1982; Molnar et al., 1996; Rouger et al., 2004), raising the possibility that SC
subpopulations might have different spatial relationships with capillaries, according, for example, to ontogenic or cycling rate specificities. We addressed this point by using two commonly used stem cell niche spotting methods (Li and Xie, 2005) and detection of quiescent stem cells retaining the BrdU label.

**Juxtavascular niches can incorporate bone marrow-derived cells**

We first tracked bone marrow-derived cells in mice in which stem cell niches are first depleted through irradiation (Labarge and Blau, 2002). Although sublaminal SCs are derived mainly from somites (Gros et al., 2005), a small numbers of SCs may be generated from bone marrow grafts (Labarge and Blau, 2002; Dreyfus et al., 2004). These ectopic SCs, express canonic SC markers (Dreyfus et al., 2004; Chretien et al, 2005) but exhibit very limited myogenic potential once re-isolated from muscle (Sherwood et al., 2004). Thus, they might represent foreign stem cells squatting in SC niches. If BMT experiments are of unknown significance in terms of SC lineage, they uniquely allow demonstration that, in the absence of the resident SC, the niche may be occupied by another cell. This information is important since adult stem cell niches are currently conceived as specific anatomic stem cell anchoring sites, which are strategically placed in the tissue and stable enough to welcome a novel stem cell (Li and Xie, 2005); this view is opposed to that of a labile microenvironment, freely generated at the stem cell request anywhere in the tissue, strictly dependent upon signals from a given stem cell, and, thus disappearing with it.

Five 4 week-old, 9.0 Gy-irradiated, wild C57BL/6 mice were transplanted with 3.10^7 BM-derived cells from Tg:CAG-GFP transgenic mice in which the GFP transgene is expressed under the control of a ubiquitous promoter (Okabe et al., 1997) and serves as an unambiguous marker for donor-derived cells in host tissues (Labarge and Blau, 2002; Dreyfus et al., 2004). Six months post transplantation, TA muscle transections immunostained for basal lamina markers (laminin or collagen IV) showed a total number of 94 GFP+ cells, 38 of which were sublaminal (Fig. 1D). Co-immunostaining showed nuclear Pax7 expression in a proportion of GFP positive cells (n=7), as
previously reported (Dreyfus et al., 2004; Chretien et al., 2005). Vascular proximity of these cells became obvious when collagen IV immunostaining was done as a supplementary reaction and merged with initial images of Pax7+ and GFP+ cells (Fig. 1 E-G).

The mean distance separating subliminal GFP+ cells from capillaries was 2.2 ± 3.8 µm: 89% were at <5 µm from ECs, 8% at 5-10 µm, 3% at >10 µm. This result is in keeping with that obtained for SCs from Myf5GFP-P/+ mice, suggesting that juxta-vascular SC niches are stable anchoring sites housing stem cells regardless of their origin.

**Juxtavascular niches contain slowly cycling BrdU-retaining SCs**

The thymidine analogue BrdU can label newly synthesized DNA in cycling cells. It is assumed that, after a long period of chase, slowly cycling cells retain a concentration of label sufficiently high to allow their detection (Fuchs et al., 2004). To readily discriminate SC nuclei from both myonuclei and interstitial cell nuclei that can also retain BrdU, we used Myf5nlacZ/+ mice. BrdU was administered intraperitoneally to 5 mice, twice daily from post-natal day 3 to 8, an age at which about 90% of TA muscle SCs incorporate BrdU (Shinin et al., 2006), presumably because they actively divide to increase muscle mass. Those cells that did not divide subsequently were detected in TA muscle as label-retaining cells after a chase period of 6 weeks. In pilot studies, due to HCl pre-treatment required for BrdU immunohistochemistry, it proved difficult to simultaneously produce high quality signals for SCs, capillaries and BrdU. Thus we combined chromogenic and fluorescence revelation approaches on the same preparations as shown in Fig. 1H-I. Among 100 consecutive β-gal+ SCs, 34 were BrdU-positive SCs. The mean distance separating SCs from capillaries was 2.5 ± 3.3 µm for BrdU-retaining SCs and 2.9 ± 3.3 µm for BrdU-negative SCs: 80 vs 74% of SCs were at <5 µm from capillaries, 10 vs 19% at 5-10 µm, 10 vs 7% at >10 µm (NS). After one dose of notexin in a control animal subjected to the same BrdU exposure regime, the percentage of BrdU-positive SCs dropped to 8% in the injected TA muscle, suggesting that the signal diluted during cell division of the progeny of previously quiescent SCs.
Thus, juxtavascular niches house quiescent SCs but also label non-retaining cells.

**SCs are juxtavascular but physically separated from ECs in mammal species**

SC proximity to capillaries was similarly detected in other mammal species, including rats (Fig. 2 A,B), dogs and humans (Fig. 2 C-E). Electron microscopy routinely showed SCs separated from ECs by the respective basal laminae. Two distinct patterns were found: close cell body apposition or SCs projecting a cytoplasmic process toward EC from a more distal site (Fig. 2 A,B). In human deltoid muscle, confocal microscopy of 30 µm-thick sections immunostained for SC markers (NCAM or MYF5) and either vonWillebrand (vW) factor, an EC marker, or collagen IV, allowed 3D reconstructions of 134 SCs. Even those SCs most closely associated with capillaries remained separated from ECs by a patent interstitial space in at least one plane of observation. Under this angle, SC-to-capillary distance was: <2µm (53%), 2-5µm (15%), 5-10µm (10%), 10-20µm (5%), >20µm (17%). A similar proximity was found at all ages from 24 to 73 years (n=17, no correlation with age) and in all tested muscle groups (data not shown).

**SCs are more frequently associated with capillaries than myonuclei**

We then determined whether the proximity of capillaries to any structure situated at the periphery of myofibers could be due to a chance event. We used large reconstructions of 4 normal human deltoid muscle cross-sections to explore the overall distribution of 337 SCs, 2972 myonuclei, and 2796 capillaries, detected by immunostaining with nuclear counterstaining. We first determined whether spatial distribution deviates from randomness, a pattern corresponding to the Poisson distribution, towards either clustering, i.e. attraction of points, or regularity, a pattern characterized by a minimal inviolable distance between points. We used Ripley’s K function (Ripley, 1988), a spatial point pattern analysis method that relies on systematic computation of the position of neighbours of each object present in the field by successive targeting. This method uniquely allows the formal demonstration of non-randomness of a spatial distribution. It demonstrated both regularity due to interposition of myofibers between points, and clustering of both SCs and myonuclei.
with capillaries unrelated to their peripheral position (see supplementary data). Highly different numbers of myonuclei and SCs precluded comparison of their respective proximity to capillaries by Kr analysis. Therefore, we used the quadrat test, with grid squares calibrated (21 µm diagonally) to enclose the largest clusters (18 µm) detected by Kr analysis. Crosstabulation tables and Fisher's exact test allowed 2 by 2 comparisons of respective frequencies of colocalization with capillaries within the same square of SCs, myonuclei, and virtual sarcolemmal points randomly distributed at the periphery of fibers. The proportion of colocalization with capillaries was much higher for SCs than myonuclei (88±6% vs 54±3%, p<0.0001) and for myonuclei than virtual sarcolemmal points (54±3% vs 35±3%, p<0.003).

**SC numbers linearly correlate with capillarisation independently from the myofiber type**

In another morphometric approach, we determined the number of capillaries immediately adjacent to each myofiber in 3 normal human deltoid muscle samples (2332 myofibers). Histogram of myofiber capillarisation frequency was close to Gaussian (Fig.2F). In each case, a majority of SCs (51, 66, 67%) was associated with myofibers ringed by 5 or more capillaries, SC detection rate increased linearly with myofiber capillarisation ($\chi^2$ test for trend, p<0.0001; Fig. 2F).

Fast-myosin immunostaining was used to discriminate fast (type II) from slow (type I) muscle fibers. In each case, type II fibers had less surrounding capillaries than type I fibers (Fig. 2G), the average capillarisation being 3.6±0.7 and 4.9±0.8 per fiber, respectively (p<0.0001). Type II fibers also showed a lower number of SCs, largely proportionate to their capillarisation (Fig. 2G).

Due to mosaicism of human muscle, a number of capillaries are shared by fibers of different types. Capillaries contacting type I, type II, or both type I and II fibers were similarly associated with SCs (range 2.4-5.2% vs 2.4-5.8% vs 2.8-6.0%, $\chi^2$ test: NS in each case). Accordingly, capillaries contacting both type I and II fibers accounted for 67-78% of capillaries surrounding type II fibers, and were associated with 63-77% of
SCs in this fiber type. A similar correlation was found in type I myofibers (42-73% and 53-74%) pointing out capillarisation as a plausible factor governing SC frequency in myofibers regardless of their type.

**Muscle capillary loss is associated with proportionate SC decrease**

As muscle blood vessels can proliferate or regress under the control of a variety of physiological and pathological stimuli (Duscha et al., 1999; Emslie-Smith and Engel, 1990; Jensen et al., 2004), we explored how SCs react to the reduction or increase of the microvascular bed. We deliberately chose situations in which microvascular changes occur without interfering pathologic myofiber alterations.

We used amyopathic dermatomyositis (aDM) as a paradigm of pure capillary loss, typically observed in the absence of both myofiber damage and inflammation (Emslie-Smith and Engel, 1990; Cosnes et al., 1995). As compared to matched controls, 3 aDM patients selected from a previous study (Cosnes et al., 1995) had myofiber capillarisation decreased by 45%, as previously reported (Emslie-Smith and Engel, 1990). Importantly a roughly proportionate decrease (53%) in the mean SC number per myofiber (Fig. 2H) was also observed. Undepleted SCs remained strongly colocalised with capillaries (quadrat test: 91±2% in aDM vs 88±6% in controls), indicating joint loss of SCs and capillaries in the same areas (Fig. 2I).

**Capillary increase is associated with both SC increase and perivascular accumulation in trained muscle**

As workload is known to increase muscle capillary density (Bischoff and Franzini-Armstrong, 2004; Carpenter and Karpati, 2001; Jensen et al., 2004), we analysed deltoid muscle samples from 3 heavy-training athletes. Athletes showed increased mean numbers of both capillaries (33%) and SCs (56%) per myofiber (Fig. 2I). SCs remained close to capillaries (quadrat test: 87±5%) where they often formed perivascular accumulations, appearing as pairs or triplets of SCs facing each other in adjacent myofibers (Fig. 2J). Similar SC accumulations around vessels are routinely detected in pathologic muscle biopsies showing repair or regeneration (data not shown). This
prompted us to examine possible EC effects on myogenic cell proliferation, in vitro.

**EC monolayers specifically increase myogenic cell growth through soluble factors**

We further investigated EC interactions with myogenic cells, using cocultures under conditions avoiding cell:cell contacts between cell types. *In vitro*, SCs are released from quiescence and undergo exponential cell growth mimicking an injury model rather than the stable state of quiescent sublaminal SCs. In the same way, the standard EC monolayer culture is characterized by both inherent dedifferentiation of ECs and lack of contact with mural cells, i.e. pericytes and smooth muscle cells, that tightly control EC maturation (Korff et al 2001). Therefore, EC monolayers mimick a situation of immature microvessels (Korff et al 2001). These characteristics imply good relevance of EC:myogenic cell cocultures to dynamic states at work during muscle tissue formation and regeneration.

Human myogenic cells were plated in the lower chamber of culture wells while the upper transwell compartment, separated by a porous filter, was seeded with human umbilical vascular ECs (HUVEC), human microvascular ECs (HMEC), human smooth muscle cells (SMC), human fibroblast cell line (MRC-5) or, as a control, other human myogenic precursor cells (mpc) derived from different primary cultures. Mpc growth was strongly promoted in the presence of EC monolayers (Fig. 3A). As compared to controls, both HUVECs and HMECs increased mpc growth, at day 3 (158%, \(p<0.02\); 227%, \(p<0.008\)), day 5 (245%, \(p<0.004\); 203%, \(p<0.003\)), and day 7 (259%, \(p<0.0001\); 212%, \(p=0.0005\)), respectively. In contrast, no increase of mpc growth was observed with SMC or MRC-5 fibroblasts. These findings demonstrate that EC monolayers specifically release soluble factors which promote myogenic cell growth.

**EC-derived IGF-1, HGF, bFGF, PDGF-BB and VEGF promote SC growth**

To detect the main EC-derived molecules increasing SC growth, we used both mRNA profiling by DNA macroarray to screen mpc expression of growth factor receptors and
protein array to assess EC secretion of soluble effectors (see methods). Of the 397 genes represented on our DNA macroarray membrane, 84 were receptors for soluble factors. Among them, receptors of 5 factors known to mediate mpc proliferation and survival signals were constitutively expressed by human mpc (Fig. 3B). The protein array allowed detection of the corresponding ligands, including IGF-1, HGF, bFGF, PDGF-BB and VEGF, in HUVEC conditioned medium (Fig. 3B).

Functional involvement of the detected molecules was assessed using specific blocking antibodies (Fig. 3C). Mpc were incubated for 3 days with antibodies deposited at saturating concentrations in the lower chamber of the co-culture insert. Whereas no changes in cell growth were detected when mpc were grown alone, significant effects of blocking antibodies were observed in co-cultures. HUVEC-sustained mpc growth decreased by 41 to 62.5% after IGF-1, HGF, bFGF, PDGF-BB and VEGF inhibition (all p<0.01). Global inhibition obtained by pooling all 5 blocking antibodies resulted in a 90% inhibition of mpc growth (p<0.01). Incubation with immunoglobulins or non relevant antibodies (MCP-1 and dysferlin) did not affect growth of mpc cocultured with ECs. These results strongly implicate the 5 tested effectors in EC paracrine stimulation of mpc growth.

Unlike other effectors, VEGF has not been definitely demonstrated to directly stimulate muscle cell growth, although previous in vivo observations gave support for this view (Arsic et al. 2004). Thus, we incubated mpc cultures with 25 ng/mL recombinant VEGF and measured cell density every 2 days. Mpc growth increased at all evaluated time points (paired Student t test : p<0.05), the increase reaching 58% at day 4 (Fig. 3D), thus confirming SC responsiveness to VEGF.

**Differentiating myogenic cells are proangiogenic in vitro**

Resident cells of many tissues can signal to ECs and influence the angiogenic process (Cleaver and Melton, 2003). We examined if muscle cells do so, using primary cultures. Angiogenesis was assessed by measuring capillary-like structure formation of HUVECs plated on Matrigel™ (Jones et al., 1999). A 24h incubation of HUVECs with day
14 human mpc-conditioned medium increased the proportion of connected ECs by 169% (42.4±13.2 in basal medium vs 71.5±8.7 in conditioned medium, p<0.04) and the number of junctions between vascular tubes by 156% (10.5±1.2 per field vs 16.4±1.4, p<0.03). A proangiogenic effect of myogenic cells was observed from the proliferation stage (day 7), increased with differentiation, as assessed by expression of the late myogenic factor Myogenin, and culminated at the time of myoblast fusion into myotubes, i.e. at day 14 in our assay (day 7 vs day 14, p<0.02) (Fig. 4I).

This effect is in agreement with our previous observation that human mpc produce VEGF, the key angiogenic factor, during their differentiation (Chazaud et al., 2003b). Therefore, we assayed for VEGF expression, \textit{in vivo} (Fig. 4II). Normal human muscle showed no VEGF expression by SCs, but faint diffuse microvascular immunostaining (Fig. 4A), as previously illustrated (Rissanen et al, 2002). aDM muscle was almost completely negative (Fig. 4B). In contrast, athlete muscles showed VEGF immunostaining in a small proportion of SCs with elongated processes suggestive of some degree of activation (Fig. 4C). Consistently, VEGF was detected in NCAM+ myogenic cells in routine muscle biopsies showing necrotic/regenerating fibers (Fig. 4D-F).

\textbf{Myogenesis and angiogenesis are spatio-temporally correlated in Duchenne muscular dystrophy (DMD)}

To further investigate possible relationships between myofiber repair and angiogenesis we analyzed muscle biopsies of 3 patients with DMD, a severe myopathy characterised by repeated cycles of intense myofiber degeneration/regeneration occurring in small clusters (Carpenter and Karpati, 2001).

We first immunolabelled the Ki-67 nuclear antigen which is associated with all cell proliferation phases but is absent in resting cells (G0). A maximum of 8% of NCAM/CD56+ cells were Ki-67+ in DMD muscles. Double staining for Ki-67 and Laminin 1 showed that 82% of Ki-67+ sublaminal SCs were colocalised with capillaries (Fig. 5A). Ki-67 was occasionally detected in SC pairs. Thus, SCs does not need to leave their
juxtavascular location to undergo active proliferation, and they can benefit from EC supportive cues in this position.

Moreover, myogenic cell differentiation assessed by Myogenin expression was associated with neoangiogenesis (Fig. 5BC). DMD muscle cross-sections contained a total number of 175 undifferentiated MYF5\(^+\) SCs, and 211 Myogenin\(^+\) cells undergoing myogenic differentiation. The quadrat test showed that 97.4% Myogenin\(^+\) nuclei (vs 88.5% MYF5\(^+\) nuclei) colocalized with CD31\(^+\) ECs. Moreover, the area occupied by ECs was about two-fold higher in grid squares enclosing Myogenin\(^+\) cells than in those enclosing MYF5\(^+\) cells (48±28µm\(^2\) vs 19±18µm\(^2\), 59±33µm\(^2\) vs 32±18µm\(^2\), 44±25µm\(^2\) vs 23±18µm\(^2\), all p<0.0001) (Fig. 5II). In addition, most capillaries associated with Myogenin\(^+\) cells were sectioned longitudinally, indicating a tortuous and highly interconnected capillary network (Fig. 5II) typical of neoangiogenesis in skeletal muscle (Hansen-Smith et al., 1996). These data reveal strong spatio-temporal correlation between myogenic cell differentiation and neoangiogenesis.

The multifocal pattern of angio-myogenesis was consistent with the distribution of VEGF immunostaining. In DMD, discrete foci of intense VEGF positivity were separated by nearly negative areas (Fig. 5D-F). It was difficult to unambiguously identify cell sources of VEGF in these foci, but the observed pattern was recently predicted by a computational model in which VEGF focally produced by muscle cells remains sequestered near sources of VEGF secretion bound to both VEGF receptors and heparan sulfate proteoglycans present at high concentrations in the muscle microenvironment (Mac Gabhann et al, 2006). Centronucleation, the hallmark of \textit{in vivo} muscle regeneration, was detected next to positive areas, and orientation of interstitial cells at this level was suggestive of angiogenic sprouts growing at nearly 90 degrees from parent vessels against the steep VEGF gradient (Fig. 5D-F).

**DISCUSSION**

In the present report we show that SC niches are juxtavascular, that they can incorporate ectopic stem cells without loosing this anatomical characteristic, that most
SCs remain in close proximity to capillaries regardless of their state of quiescence, proliferation and differentiation, and that individual myogenic cell differentiation is spatio-temporally associated with new vessel formation. In light of our *in vitro* experiments, these previously unappreciated morphologic characteristics likely favour bi-directional signalling between ECs and differentiating SCs at work during muscle regeneration.

Factors that govern SC frequency are not known (Bischoff and Franzini-Armstrong, 2004). Although the frequency of SCs appears to be much higher in slow compared to fast muscles (Zammit and Beauchamp, 2001), there is little difference at birth, and the metabolic properties *per se* may not be crucially involved (Bischoff and Franzini-Armstrong, 2004). Capillaries are more numerous in slow muscles than in fast ones (Hudlicka et al., 2004). This may represent a key factor if one considers the linear increase of SCs with capillaries we observed at the individual myofiber level, regardless of the myofiber type. Notably, to avoid influence of the different workload of slow and fast individual muscles, a single type I: type II balanced muscle was chosen to compare fiber types. Conversely, in aDM, where focal capillary loss is insidious enough to cause no myofiber damage (Emslie-Smith and Engel, 1990), SC loss is proportionate to capillary loss and selectively affects capillary-depleted myofibers. Of note, this is not the case in full-blown dermatomyositis, where strong SC proliferation may be observed, despite marked capillary loss, in reaction to conspicuous ischemic myofiber damage and inflammation (Carpenter and Karpati, 2001). SCs are more resistant to acute ischemia than myonuclei (Schultz et al., 1988; Snow, 1977), suggesting that disappearance of SCs in areas of capillary dropout in aDM may reflect, at least in part, loss of supportive paracrine EC signals. Investigating factors involved in SC maintenance and self-renewal would likely require complex assays integrating EC and myofiber-to-SC signalling. Detection of perivascular SC accumulations in trained muscle led us to examine the role of ECs in more dynamic states associated with SC proliferation. We used indirect EC:mpc coculture as an *in vitro* counterpart of muscle repair. EC monolayers that mimick immature capillaries supported mpc growth through release of soluble factors.
Notably, mpc growth was not supported by fibroblasts and smooth muscle cells, two other cell types that may be found in the endomysium close to SCs. A direct role for paracrine signaling between ECs and surrounding target cells has been described during embryonic development and cell differentiation of several organs (Cleaver and Melton, 2003) but data in muscle are scant. Endothelial progenitors have been isolated from fetal tissues and adult blood (Le Grand et al., 2004; Shmelkov et al., 2005) and can be instructed to co-differentiate into functional angiomyogenic colonies (Shmelkov et al., 2005) and restore dystrophin in dystrophin-deficient mice and dogs (Le Grand et al., 2004; Torrente et al., 2004, Sampaiolesi et al., 2006). As shown here-in, circulating stem cells can incorporate juxtavascular SC niches of irradiated mice (Labarge and Blau, 2002; Dreyfus et al., 2004) but the physiologic relevance of this phenomenon remains far from clear (Sherwood et al, 2004). Nevertheless, vascular precursors enter the quail limb buds around the time of muscle precursors (Nimmagadda et al., 2004), and vessels develop before the appearance of SCs in the mouse limb (Le Grand et al., 2004). In wing buds of avian embryos, migrating myogenic cells may be found in close proximity to a prepatterned vascular network (Solursh et al., 1987). In line with these observations, spatio-temporal correlations between the ingrowth of blood vessels into a formerly ischemic area and detection of cells with myogenic capacities in that area has been repeatedly reported (Kardon et al., 2002; Phillips et al., 1987). Although the lineage relationships between ECs and muscle progenitors is not firmly agreed upon in the literature, our present studies do strongly support a functional relationship.

Post-injury muscle regeneration is associated with an increase in both capillarisation and cross-sectional area of individual regenerating myofibers (Luque et al., 1995). In the same way, most data on compensatory muscle hypertrophy due to overload reveal an almost linear relationship between capillary-to-fiber ratio and fiber area (Snow, 1977). Our anatomic and functional results indicate what cellular events may underpin these structural changes. Human myogenic cells undergoing differentiation secrete VEGF (Chazaud et al., 2003b). Conversely, we and others have shown that human ECs can produce a series of mitogens for myogenic cells (Hawke and Garry, 2001), and our
co-culture studies indicated that bFGF, IGF-1, HGF, VEGF, and PDGF-BB mediate the bulk of paracrine EC sustaining effect on mpc growth. Recently, attention has been paid to the role of VEGF and its receptors in muscle biology (Williams and Annex, 2004). Under normal in vivo conditions, mature ECs do not produce VEGF but express VEGF receptors R1 (Flt-1) and R2 (KDR/Flk-1) which may bind low-levels of VEGF uniformly released by normal muscle tissue (Mac Gabhann et al, 2006). Exercise-induced VEGF upregulation in muscle cells operate physiological angiogenesis (Egginton et al, 2001; Ameln et al, 2005). Myoblasts manipulated to produce VEGF specifically promote HUVEC proliferation (Kim et al, 2005) and induce pronounced localized angiogenesis in their immediate vicinity after i.m. transplantation (Springer et al, 2003). A steep gradient of VEGF concentration emanating from the implantation site into neighboring host muscle tissue has been repeatedly predicted, but was not previously documented by immunolocalization (Springer et al, 2003; Mac Gabhann et al, 2006). It was visualized herein in DMD, possibly because VEGF release occurs at a particularly high rate in dystrophin deficient tissue (Nico et al, 2002).

VEGF receptor expression is not restricted to ECs and VEGF effects extend to a variety of non-endothelial cell types, including myogenic cells (Rissanen et al, 2002, Germani et al. 2003, Motoike et al, 2003, Arsic et al. 2004). We showed that VEGF stimulates in vitro myogenic cell growth. In addition, VEGF stimulates their migration (Germani et al. 2003), protects them from apoptosis (Germani et al. 2003, Arsic et al. 2004), up-regulate myoglobin expression (van Weel et al., 2004), and promotes formation of centronucleated myofibers (Arsic et al. 2004).

On these grounds, evidence that, upon activation, SCs accumulate and proliferate close to capillaries, receive efficient support from ECs for their growth, are proangiogenic, and colocalize with new vessel formation during their myogenic differentiation, strongly suggests that angiogenesis and myogenesis reciprocally signal. It is likely that angiogenesis and myogenesis share VEGF as a co-regulatory factor (Williams and Annex, 2004) probably in combination with other factors known to stimulate both ECs and SCs, such as bFGF or IGF1 (Bischoff and Franzini-Armstrong,
2004; Hawke and Garry, 2001). We assume that, following workload or more pronounced injury, angiogenesis is tightly coordinated with SC proliferation, differentiation, and fusion to underlying myofibers. Newly SC-derived myonuclei may remain close to capillaries, as suggested by the frequent colocalization of myonuclei with capillaries. Occasional observation of SCs immediately adjacent to a myonucleus (Fig. 2J) is even reminiscent of the model proposed by Moss and Leblond (1971) in which asymmetrical SC division generates a new SC and a daughter cell fated to become a myonucleus. Newly incorporated myonuclei likely produce muscle-specific proteins causing increase of the myofiber cross-section size. Harmonious increase of myofiber capillarisation and caliber ensues, allowing larger myofibers to benefit from an appropriately enhanced blood supply.

Such a myo-vascular unit may share similarities with the vascular neural stem cell niche that supports intimate combination of neurogenesis and angiogenesis in the brain (Palmer et al., 2000; Shen et al., 2004).

The knowledge that muscle SCs evolve into a juxtavascular niche may be critical for several reasons. It provides a novel clue to investigate poorly understood muscle pathologic processes combining capillary bed and myofiber size variations (Emslie-Smith and Engel, 1990; Duscha et al., 1999; Hudlicka et al., 2004; Jensen et al., 2004). Moreover, it could pave the way for improved myoblast transfer therapy of heart, sphincter and skeletal muscle diseases. Dramatic non-mechanic transplanted cell death is a major limitation of this therapeutic approach (Chazaud et al., 2003a). The present results suggest that early cell death results from extrinsic growth factor deprivation of mpc massively implanted in an inappropriate microanatomic environment (Chazaud et al., 2003b). Recreating an appropriate endothelial cell or molecular support to the transplanted stem cells thus represents a promising way to improve striated muscle cell therapies.

**MATERIAL AND METHODS**

**Transgenic mouse strains.**
Mice models allowing visualization of satellite cells

Constructions of both Myf5\textsuperscript{nlacZ} and Myf5\textsuperscript{GFP-P} knock-in mice have been described (Tajbakhsh et al., 1996; Kassar-Duchossoy et al., 2004). Both mice have a ScaI-BstYI deletion in exon 1 (122 amino-acids) of Myf5 gene, thus lacking the basic-HLH domain required for Myf5 transcription factor activity, and a reporter gene targeted to this exon. Myf5\textsuperscript{GFP-P/P} are reduced in size, whereas both heterozygote mice, Myf5\textsuperscript{nlacZ/+} and Myf5\textsuperscript{GFP-P/+}, are phenotypically normal and were used for the experiments.

Mice used for bone-marrow transplantation

C57BL/6 mice were transplanted with BM-derived cells from Tg:CAG-GFP transgenic mice (C57BL/6 Tg[actEGFP]Osb YO1) in which the GFP transgene is expressed under the control of a non-tissue specific promoter, chicken beta-actin with cytomegalovirus enhancer, as a cytoplasmic protein (Okabe et al., 1996).

Animal experiments.

Mouse handling

All mice were housed in our level 2 biosafety animal facility, and received food and water \textit{ad libitum}. Prior to manipulations, animals were anaesthetized using intraperitoneal injection of chloral hydrate. This study was conducted in accordance with the EC guidelines for animal care [Journal Officiel des Communautés Européennes, L358, december 18, 1986].

Bone marrow transplantation

Briefly, donor BM cells were obtained by flushing femurs of Tg:CAG-GFP mice with DMEM medium (Invitrogen, Paisley, UK), and washed twice in cold PBS. Retro-orbital injection of 3 \times 10^7 BM cells in 0.1 ml mouse serum and PBS (1:1), was done in 9.0 Gy-irradiated, 4 week-old B6 mice (\textsuperscript{60}Co \textgamma-rays within 1 day before BM transplantation) as previously reported (Dreyfus et al., 2004). After transplantation, mice received 10 mg/kg/day ciprofloxacin for 10 days to prevent infection during the aplastic phase. Blood chimerism >95% was controled by flow cytometry 4 weeks post-transplantation and TA muscles were examined 6 months post-transplantation.
BrdU pulse-chase experiments

*Myf5*nlacZ/+ pups received 50 mg/kg BrdU diluted in 20 µL PBS by I.P. route twice daily from post-natal day 3 to 8. After a chase period of 6 weeks, mice were sacrificed and TA muscles were examined to evaluate BrdU retention by β-Gal+ satellite cell nuclei. As a control, one *Myf5*nlacZ/+ mouse from the series received notexin injection (25µg/ml in 10 µl PBS) in TA muscle 6 weeks post-exposure and had the injected TA examined 2 weeks later to assess dilution of BrdU in the progeny of the previously quiescent satellite cells.

Human muscle samples.

Normal muscle samples were obtained in 20 human adults (aged from 24 to 73 years, from various muscle groups, including 12 deltoid, 4 vastus lateralis, 2 gastrocnemius and 2 peroneus brevis muscles), in 3 adult C57/B6 mice (tibialis anterior muscle), in 2 adult retriever dogs (sartorius muscle) and, for electron microscopy, in 10 adult Wistar rats (extensor digitorum longus muscle). Muscle samples were also collected from deltoid muscle of 3 heavy-training athletes who had no significant myopathologic alterations (males aged from 32 to 39 years practicing intense daily push-up training), of 3 patients with amyopathic dermatomyositis collected from a previous study (Cosnes et al., 1995) (2 females, 1 male aged from 32 to 39 years), and of 3 boys with Duchenne muscular dystrophy (males aged from 2 to 14 years). Muscle samples were frozen and kept at -80°C before processing.

Immunohistochemistry.

In mouse tissues, vessels were immunostained using anti-laminin 1 (1:50, polyclonal, DakoCytomation, Glostrup, Denmark) or anti-collagen IV (1:50, polyclonal, Chemicon, Temecula, CA) antibodies revealed by Cy3 conjugated secondary antibodies (Jackson Immunoresearch Laboratories, West Grove, PA). Due to HCl pre-treatment required for BrdU immunohistochemistry, we combined chromogenic and fluorescence revelation approaches on the same preparation. First, endogenous alkaline phosphatase activity of microvascular cells (Pearce et al., 2000) was detected using fast red (forming both
chromogenic and fluorescent red precipitate) followed by chromogenic immunohistochemistry against β-Gal (using a polyclonal domestic antibody, from Pasteur Institute, 1:2000e revealed by a peroxidase-conjugated secondary antibody) forming a brown precipitate. Afterwards, sections were treated with 2N HCl for 30 min at 37°C and BrdU was detected using a sheep polyclonal antibody (1:200e, Abcam Inc, Cambridge, MA) revealed by FITC-conjugated secondary antibody (Jackson Immunoresearch lab).

In human tissues, several antibodies were used alone or in combination to localize and identify SCs and their progeny, ECs and VEGF. They included antibodies against laminin-1 (1:500, ref L9393, Sigma Aldrich, St-Louis, MO) for basal laminae, MYF5 (1:50, ref sc302, Santa Cruz, Santa Cruz Biotechnology, CA) for undifferentiated SCs, CD56/NCAM (1:100, ref PN6602705, Beckman Coulter, Fullerton, CA) for both quiescent and differentiating myogenic cells, myogenin (1:200, ref M3559, DakoCytomation) for differentiating myogenic cells, CD31 (1:50, ref M0823, DakoCytomation) and vWF (1:500, ref A0082, DakoCytomation) for ECs, Ki-67 antigen (1:50, ref M7240, DakoCytomation) for cycling cells and VEGF (1:100, AF293NA, R&D Systems, Minneapolis, MN, for immunofluorescence; or 1:50, sc-152, Santa Cruz Biotechnology, with SuperPicTure™ polymer detection kit, Zymed, and AEC as a chromogen for light microscopy).

Dogs samples were labelled using the same antibodies to CD56/NCAM and vWF.

**Electron and confocal microscopy.** Transmission electron microscopy was performed after conventional processing of muscle samples using a Philips-410 microscope. Confocal microscopy (Zeiss LSM 410 and a x63 PlanApo NA 1.4 oil objective, Carl Zeiss, Oberkochen, Germany) on 10-30 µm-thick sections and deconvolution of stacks of optical sections obtained along the Z-axis were used to obtain high resolution images and three-dimensional volume reconstructions. Steps along the optical axis were set at 0.2-0.3 µm and pixel size in X,Y at 0.11-0.14 µm thus respecting Nyqvist’s criteria optimal for subsequent deconvolution. Z-stacks acquired separately for each channel (green or red) were subjected to iterative optical deconvolution by the AxioVision 4.2
software package defining appropriate parameters of a theoretical Point Spread Function and settings to correct for optical aberration. Deconvolution (15–25 iterations) was performed on a personal computer equipped with a Pentium 4 processor (3.0 GHz, 2 Go RAM). In rare cases, better results were obtained by the Nearest Neighbour algorithm of AxioVision 4.2. 3D reconstructions combining both channels were then obtained in the transparency option of the 4D module of AxioVision 4.2. Measurement of the minimal distance between SC and the neighbouring capillary were carried out on the projection (rotation) that excluded spurious colocalization of channels that could potentially result from the considerable Z-thickness of the reconstruction (10-20 µm).

**Morphometric data collection.** Muscle areas showing motor endplates were excluded from analysis. Immunostained sections were digitized with a Hamamatsu CDD device (Hamamatsu Photonics KK, Hamamatsu, Japan) and subjected to morphometric analysis using different modules of the KS4003.0 (Carl Zeiss) and Simple PCI image analysis software (C-Imaging Compix Inc, Cranberry Township, PA). Each object (SC, Cap, myonuclei or VSP) was defined as a point with X and Y coordinates in a study area including at least 1000 myofibers per specimen. Because of either difficulty to discriminate between closely associated structures on double-immunostainings, or limited use of antibodies of the same species for double-immunostainings, some quantitative evaluations were conducted using single immunostainings on alternate muscle cross-sections. For example, alternate sections were immunostained for MYF5, CD31 and myogenin, myogenic cell positions being manually reported in the graphic plane of image reconstruction of the alternate immunostaining of capillaries.

**Univariate spatial point pattern analysis.** To explore if point distributions deviate from complete spatial randomness (CSR) We calculated spatial point pattern statistics with the ADE4 software package, using Ripley’s K-function for univariate point patterns, and both the K12 function and a quadrat test for bivariate patterns (Ripley, 1988). Basically, the K function assesses for each point the cumulative number of neighbouring points within increments of a predefined distance t as compared with the expected number of neighbours under the null hypothesis of CSR. These increments define "n"
concentric rings so that the outer radius of the external ring \( r = nt \). According to recommendations of the ADE4 manual, \( t \) and \( n \) values were determined for each set of data, taking into account both size of the study area \( R \) and the observed minimal distances between points. The definition of Ripley’s \( K \) is \( K(r) = N(r)/\lambda \), where \( N(r) \) is the number of neighbours within distance \( r \) and \( \lambda \) is the intensity of the pattern. Under CSR, \( K(r) = \pi r^2 \), in case of clustering, \( K(r) > \pi r^2 \), and in case of regularity, \( K(r) < \pi r^2 \). By convention, \( K(r) \) is substituted by \( L(r) = \sqrt{K(r)/\pi} - t \), this transformation offering the advantage of \( L(r) = 0 \) under CSR, \( L(r) > 0 \) for clustered, and \( L(r) < 0 \) for regular patterns. Edge correction was carried out as proposed by Ripley (Ripley, 1988). Deviation from CSR was tested by plotting \( L(r) \) values against the envelope of significance at \( p<0.0001 \) for the null hypothesis of CSR. This envelope was built using the Monte Carlo method that consists in the realization of 9999 CSR patterns of the same intensity as the observed pattern. Graphically, values above the upper limit of the envelope indicate clustering whereas values below its lower limit indicate regularity.

**Bivariate point pattern analysis.** The K12 function, and its L12 transformation, for bivariate patterns is identical to Ripleys' \( K \) with the exception that points on which the function is centered and neighbour points are of two different types, i.e. correspond to different objects. Graphic expression of results was similar to that used for Ripleys'K. The distribution of real objects, such as SC and myonuclei, relative to capillaries was compared to that of randomly distributed virtual sarcolemmal objects, one object per myofiber being randomly inserted following a clock dial scheme.

**Quadrat test.** A quadrat test grid was superimposed in the graphic plane of each image. The grid square size (21 µm diagonally, 225 µm\(^2\)) was chosen to enclose the largest clusters of points detected by bivariate analysis, as deduced from the graphic expression of the \( K \) function. Each square was examined for the presence of SC, myonuclei, VSP and Cap and colocalization was estimated by a Fisher's exact test comparing the relative number of squares containing a capillary and either SC, myonuclei, or VSP. In some tests, the capillary area (µm\(^2\)) in each square was measured after color segmentation (DAB, brown) of CD31 labelling of vessels using KS400 3.0.
**Myofiber capillarisation evaluation.** Muscle fiber capillarisation in normal, aDM, and athlete muscles was assessed by the number of capillaries bordering each individual fiber, as previously described (Emslie-Smith and Engel, 1990). Cap numbers and frequency distribution in aDM and control patients of our study were closely similar to those previously reported (Emslie-Smith and Engel, 1990).

**Cell cultures.** Unless indicated, culture media components were from Gibco (Paisley, Scotland) and culture plastics from TPP AG (Trasadingen, Switzerland). Human mpc were cultured from muscle samples as previously described (Chazaud et al., 2003b). Mpc were grown in HAM-F12 medium containing 15% FCS. Only cultures presenting over 95% NCAM/CD56\(^+\) (1/20, 123C3, Sanbio/Monosan, Uden, Netherlands) cells were used. Mpc differentiation was assessed by myogenin immunoblotting. MRC-5 human fibroblastic cell line was obtained from ATCC (LGC-Promochem, Molsheim, France) and cultured in MEM containing 10% FCS and 1% NEAA. HUVEC and HMEC were obtained from Promocell GmbH (Heidelberg, Germany) and cultured in EC growth medium (Promocell). SMC were obtained from human pulmonary arteries kindly provided by Dr S. Eddahibi (INSERM U651, Créteil, France) and cultured in RPMI containing 10% FCS and 1% NEAA.

**In vitro angiogenesis assay.** HUVEC were seeded (30000 cells/well) in 24-well tissue culture plates coated with growth factor-reduced Matrigel (0.3 ml/well, Becton Dickinson) that was allowed to solidify at 37°C for 30 min. Cells were incubated in serum-free medium or in 24h mpc-conditioned medium for 24 h at 37°C. The cells were photographed using an inverted photomicroscope (DMIRB Leica, Leica Microsystems AG, Wetzlar Germany) with a CCD camera (CoolSnap, Photometrics, Roper Scientific Inc. Tucson, AZ). Tube formation was assessed in 5 randomly selected x20 powerfields, using the number of branching points in the network per field and the proportion of connected cells (number of connected cells among the total number of cells in the same field) (Jones et al., 1999).

**Indirect cocultures.** MRC-5 fibroblasts were seeded at 15000 cells/cm\(^2\), SMC and mpc at 20000 cells/cm\(^2\) in Falcon inserts (0.4 µm diameter pores) (Falcon, BD
Biosciences, Franklin Lakes, NJ); EC were seeded at 30000 cells/cm$^2$ and 3 days after confluence, EC monolayer integrity was assessed in 2 wells by absence of Trypan blue (0.2% in 0.1% BSA) translocation from upper to lower chamber after 3 h incubation. Mpc were seeded into 24 well plates at 3000 cells/cm$^2$ and allowed to adhere in their culture medium for 6 h. Then, inserts were placed over the mpc-containing well and the medium of both chamber was changed for MCDB-131 medium containing 2% FCS and 1.5 µg/ml ECGS. The medium was changed twice weekly. At each time point, mpc density was determined by counting cells after trypsinization, and the number of cell present in the upper chamber was also evaluated. In these conditions, the number of EC remains constant at each time point as they form impermeable monolayers. In contrast, the number of SMC, MRC-5 and control mpc increased during time of experiment to reach approximatively 60000, 60000 and 40000 cells/cm$^2$, respectively. To avoid bias due to the different numbers of cells in the upper chamber at each time point, calculation of the effect on mpc growth was normalized to 30000 cells/cm$^2$, thus allowing comparison of individual cell biological activity, as previously reported (Chazaud et al., 2003b). All these in vitro experiments were performed using at least 3 different cultures. In some experiments, blocking antibodies were added in the well at saturating concentrations (calculated from IC50): anti-IGF-1 (6 µg/ml, AF291NA, R&D), anti-PDGF-BB (5 µg/ml, AF220NA, R&D), anti-bFGF (6 µg/ml, AF233NA, R&D), anti-HGF (5 µg/ml, AF294NA, R&D), anti-VEGF (6 µg/ml, AF293NA, R&D). Controls included addition of whole mice IgGs (6 µg/ml, Vector Laboratories, Burlingame, CA), antibodies against muscle membrane protein dysferlin (3 µg/ml, Novocastra laboratories, Newcastle, UK), and antibodies against MCP-1 (5 µg/ml, P500-P34, Abcys, Paris, France), a chemokine that is secreted by EC (Cleaver and Melton, 2003) and the receptor of which is expressed by human mpc (DNA array, data not shown).

Mpc grown alone were subjected to similar blocking antibody experiments and to recombinant human VEGF (RandD systems) added at 25 ng/ml.

**DNA Array.** Total RNA was prepared from mpc at day 14 of culture using the RNeasy mini kit (Qiagen, Hilden, Germany). All further steps were performed according to the
manufacturer's instructions in Human cytokine array kit (R&D Systems). Analysis was performed using Image Quant software (Amersham, Buckinghamshire, UK), that allows background noise subtraction, correction for the variation of density for housekeeping genes (all genes showed the same intensity variation between the 2 membranes), and finally, densitometric analysis of signals above internal negative controls. Results were expressed in arbitrary units.

**Protein Array.** 24h-HUVEC- conditioned medium was subjected to protein array according to the manufacturer's instructions in RayBio® Human Cytokine Antibody Array (RayBiotech Inc., Norcross, GA). Densitometric analysis was performed as described above for DNA array.

**Statistical analyses.** Fischer's exact test was used to compare colocalizations in the quadrat test. $\chi^2$ test was used to compare frequency of SCs associated with variously capillarized myofibers, and with various Cap types. Student's t test was used in in vitro experiments. A p value <0.05 was considered significant.
FIGURE LEGENDS

Figure 1. SC proximity to capillaries in mouse models

(A) TA muscle cryosections from Myf5\textsuperscript{\textasciitilde lacZ/+} mouse in which SC nuclei expressing nuclear β-Gal activity (blue pseudo-color) are observed close to microvessels which thick basal lamina is visualized by enhanced collagen IV immunoreactivity (red) bar: 10 µm.

(B) TA muscle cryosections from Myf5\textsuperscript{\textasciitilde lacZ/+} mouse in which most SC nuclei expressing nuclear β-Gal activity (blue) are found close to capillaries expressing alkaline phosphatase activity (red) bar: 10 µm.

(C) TA muscle cryosections from Myf5\textsuperscript{GFP-P/+} mouse in which one SC expressing cytoplasmic GFP (green) is closely associated with a capillary. Laminin 1 is labeled in red. Bar 10: µm.

(D) TA muscle cryosections from a WT mouse transplanted with GFP+ bone marrow-derived cells (BMDC) showing a blood borne GFP+ cell housed in a juxtavascular SC niche. Bar 10: µm.

(E) The nucleus (blue) that expresses the satellite cell marker Pax7 (red) belongs to a blood borne GFP+ cell (green). Collagen IV immunostaining performed as a second step indicates that the cell is subliminal and juxtavascular.

(F-H) TA muscle cryosections from a Myf5\textsuperscript{\textasciitilde lacZ/+} mouse perinatally infused with BrdU showing one label-retaining SC located close to a capillary: BrdU immunostaining appears in fluorescent green in both dark field (F) and merge with DIC (G), microvascular alkaline phosphatase activity appears in red at both fluorescent (E,G) and photonic (H) microscopy, SC nuclear β-Gal antigen is revealed in brown by immunoperoxidase (H). Bar: 10 µm.
Figure 2. Topological relations between SCs and Capillaries

(A-B) Transmission electron microscopy of rat SCs. Both apposition of SC body to EC (A) and apposition of a cytoplasmic process elongated from a distant SC toward EC (B) are observed. For clarity, SC plasma membrane was underlined by red dots. Please note the absence of direct SC-to-EC contact. Bars: 2 µm.

(C-E) Chromogenic or fluorescent immunostainings of human deltoid muscle showing extreme proximity of SCs and capillaries. In C, SCs (black arrowheads) are immunostained for NCAM and capillaries (red arrowheads) are identified without immunostaining. In D, a SC nucleus is immunostained for Myf 5 (black arrowhead) and capillaries (red arrowheads) are immunostained for CD31. In E, NCAM of SC plasma membrane is green and the EC marker vWF is red. Bar: 10 µm.

(F-G) Correlations between SCs and capillaries in normal muscle. Left graph (F): Histogramm of myofiber capillarisation frequency (top) is close to Gaussian while SC frequency (bottom) increases linearly with myofiber capillarisation (data pooled from 3 normal deltoid muscles). Right graph (G): numbers of both capillaries (●) and SCs (+) are different and largely proportionate in type I and type II myofibers (individual results from 3 normal deltoid muscles identified by different colors).

(H) Correlations between SCs and capillaries in aDM muscle. A proportionate loss of muscle capillaries (●) and SCs (+) was found in 3 patients with aDM (in red, grey and white, respectively) as compared to 3 age, sex, and muscle-matched normal controls (in black).

(I-J) Heavy-trained muscle. (I): as compared to their controls (in black), 3 athletes (in red, grey and white, respectively) show an increase of both muscle capillaries (●) and SCs (+). (J): SC clusters are formed by accumulations of SCs (NCAM⁺, brown) belonging to different myofibers, often adjacent to a myonucleus (haematoxylin nuclear counterstaining in blue), and surrounding the same capillary (red dots). Bar: 10 µm.
Figure 3. ECs stimulate SC growth

(A) *In vitro* mpc growth in presence of various cell types. Mpc cocultures were done using a bicameral insert avoiding direct cell:cell contacts between cell types. Coculture with either HUVEC or HMEC markedly increased mpc growth, whereas other cell types, including SMC, MRC-5 fibroblasts or heterologous mpc had no effect. Photos show representative mpc cultures with and without HUVEC at day 5 of coculture.

(B) Selection of candidate effectors. DNA macroarray and protein array showed expression of 5 growth factor receptor mRNAs and their ligands by human mpc and HUVEC, respectively, assessed by signals above internal negative controls. Signal intensity is conventionally expressed in arbitrary densitometric units.

(C) Functional involvement of the candidate effectors. Mpc grown alone and incubated with blocking antibodies against IGF-1, HGF, bFGF, PDGF-BB and VEGF showed no significant change in their growth. In contrast, in cocultures, the same antibodies induced reduction of HUVEC-induced growth effect. This was not observed with non-relevant antibodies (whole Igs, anti-MCP-1 or anti-Dysferlin antibodies) assessing implication of the five effectors in EC paracrine effects on mpc.

(D) Incubation of mpc cultures with recombinant VEGF. Human mpc cultured with VEGF (25 ng/ml) showed significant enhancement of their growth as compared to mpc cultured without VEGF (p<0.05). Results are mean + or - of 5 experiments run in duplicate.
Figure 4. Myogenic cells are proangiogenic

(I) In vitro mpc proangiogenic activity. As compared to control (top), mpc (day 14) conditioned-medium (bottom) stimulated the formation of tubular-like structures by HUVEC in matrigel™ (24h exposure, phase contrast). Pro-angiogenic activity significantly increased with mpc differentiation, assessed by Myogenin immunoblotting.

(II) In vivo localization of VEGF. (A) Normal human muscle show weak microvascular immunostaining; (B) amyopathic dermatomyositis shows virtually no signal; (C) athlete muscle shows VEGF immunostaining in SCs with elongated processes; (D-F) Immunofluorescence shows a myofiber undergoing post-necrotic regeneration enclosing marginally located CD56+ myogenic cells (green) coexpressing VEGF (red). Bar = 5 µm (except C where magnification is about 4-fold higher).
Figure 5. Combined myogenesis and angiogenesis in DMD

(I) Myogenesis and angiogenesis in DMD. (A) Double immunostaining for nuclear Ki-67 antigen (green) and laminin 1 (red) shows a sublaminal SC undergoing cell cycling close to a capillary; (B, C) alternate sections immunostained for Myogenin and CD31, showing differentiating myogenic cells directly adjacent to transversely oriented neovessels; (D–F) VEGF immunostaining showing two discrete foci of intense positivity visualizing VEGF, sequestered next to myofibers with a central nucleus, directing transverse angiogenic sprouting.

(II) Spatial association of myogenesis and angiogenesis in DMD. Selected squares enclosing both CD31+ ECs (brown) and either MYF5+ (green points) or myogenin+ (yellow points) cells from quadrat test applied to a DMD muscle. Myogenin+ cells undergoing late myogenic differentiation were associated with larger EC cytoplasmic areas than undifferentiated MYF5+ positive SCs (these illustrations were built from alternate sections immunostained for MYF5, CD31 and Myogenin, myogenic cell positions being manually reported in the graphic plane of image reconstruction of CD31 immunostaining).

Supplementary Figure. Non-random proximity of SCs and capillaries in human skeletal muscle

Graphic expression of univariate (red curves) and bivariate (blue curves) point pattern analyses obtained by L(r) transformation of Ripleys K function in human deltoid muscle (n=4, age range 20-33 years). Values above the upper limit of the envelope (in between black curves) of complete spatial randomness built using the Monte Carlo method indicate clustering whereas values below its lower limit indicate regularity:

- univariate analysis of all objects taken together, including myonuclei, SCs, and capillaries (left graph, red curve) shows both point clustering (0 to 18 µm) and regularity (25 to 50 µm). Univariate analyses of myonuclei and capillaries studied alone (right graphs, red curves) show no clustering whereas regularity imposed by the
myofiber cytoplasmic area is still observed.

- bivariate analysis of both SCs vs capillaries and myonuclei vs capillaries (left graphs, blue curves) indicate clustering of both SCs and myonuclei with capillaries, a finding not observed with virtual points introduced at random along sarcolemma (VSPs vs capillaries, right graph, blue curve). All graphs were obtained from cross-sections double-immunostained for NCAM/CD56 and CD31. (Mn means myonuclei, caps means capillaries, SCs mean satellite cells, VSPs mean virtual sarcolemmal objects).
REFERENCES


SUPPLEMENTARY DATA

SCs, myonuclei, and capillaries are not randomly distributed in human muscle

We first explored spatial distribution of SCs, myonuclei, and capillaries in muscle cross cryosections immunostained for SC markers, NCAM/CD56 or MYF5, and EC markers, CD31 or Von Willebrand factor (vWF), with nuclear counterstaining (Fig 1). Whether their distribution deviated from complete spatial randomness (CSR) towards either clustering or regularity, was assessed by Ripley's K function (Kr). This spatial point pattern analysis method relies on the systematic computation of the neighbours of each point in the field (Ripley, 1988). Kr is conventionally used in geostatistics and material science, and was preferred to usual histologic morphometric methods since it uniquely allows the formal demonstration of spatial non-randomness. Normal human deltoid muscle samples (n=4, age range 20-33 years) contained a total number of 2972 myonuclei, 337 SCs, and 2796 capillaries. In each case, overall distribution of myonuclei, SCs and capillaries (univariate Kr) showed a particular pattern characterized by point clustering within a distance of 0 to 18 µm from targeted points, followed by regularity at a distance of 25 to 50 µm, and by randomness at longer distances (Fig 1a). Regularity was imposed by the myofiber cytoplasmic area interposed between peripherally situated subsarcolemmal myonuclei, sublaminal SCs, and interstitial capillaries. Consistently, regularity was similarly observed when myonuclei or capillaries were separately analyzed (Fig 1a), and when virtual sarcolemmal points (VSPs, n=567) were introduced at random into images. In contrast, clustering was not found when each type of point was separately analyzed. Thus, in normal muscle, concentrations of homologous points could not account for clustering detected when all points were analyzed together (Fig. 1a).

Both SCs and myonuclei show clustering with capillaries

To explain clustering in the overall pattern, we examined distributions of points corresponding to different structures by bivariate point pattern analysis. Bivariate Kr
showed marked clustering between capillaries and both SCs and myonuclei (Fig 1a). Capillaries involved in these clusters were within a distance of 0 to 20 µm from SC or myonuclei (Fig 1a). A similar analysis carried out with VSPs in place of SCs or myonuclei showed that peripheral position of SCs and myonuclei does not account for their pronounced clustering with capillaries (Fig 1a). Highly different numbers of myonuclei and SCs precluded comparison between respective frequency of myonuclei and SC clustering with capillaries by Kr analysis.
normal muscle

F

G

E

aDM muscle

athlete muscle

H

I

J

Figure 2
A - *in vitro* mpc growth in presence of various cell types

![Graph showing mpc density in lower chamber with and without EC](image)

B - selection of candidate effectors

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C - effector inhibition by blocking antibodies

![Bar chart showing mpc density with and without antibodies](image)

D - effect of VEGF on mpc growth

![Graph showing mpc density over days with and without VEGF](image)
I - angiogenesis

control

mpc supernatant

Tube formation (% of control)

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myogenin expression

II - VEGF expression

A

B

C

CD56

VEGF

merge

FIGURE 4
I - Myogenesis and angiogenesis in DMD

II - Spatial association of myogenesis and angiogenesis in DMD

FIGURE 5
Point pattern analysis (Kr)

all objects

Mn

Caps

SCs vs Caps

Mn vs Caps

VSPs vs Caps

Supplementary figure