Structural and Functional Alterations of Skeletal Muscle Microvasculature in Dystrophin-Deficient mdx Mice.
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Title: Structural and functional alterations of skeletal muscle microvasculature in dystrophin-deficient mdx mice

List of authors:
Claire Latroche\textsuperscript{1,2,3,4}, Béatrice Matot\textsuperscript{5,6}, Aurea Martins-Bach\textsuperscript{5,6,7}, David Briand\textsuperscript{1}, Bénédicte Chazaud\textsuperscript{2,3,4,5}, Claire Wary\textsuperscript{5,6}, Pierre G. Carlier\textsuperscript{5,6}, Fabrice Chrétien\textsuperscript{1,4,8,9} & Grégory Jouvion\textsuperscript{1,4,9}

Full affiliations of all authors:
\textsuperscript{1}Institut Pasteur, Infection and Epidemiology department, Human Histopathology and Animal Models, Paris, France; \textsuperscript{2}INSERM U1016, Institut Cochin, Paris, France; \textsuperscript{3}CNRS UMR8104, Paris, France; \textsuperscript{4}Paris Descartes University, PRES Sorbonne-Paris-Cité, Paris, France; \textsuperscript{5}Institut de Myologie, NMR Laboratory, Paris, France; \textsuperscript{6}CEA, I²BM, MIRCen, IdM, NMR Laboratory, Paris, France; \textsuperscript{7}Laboratory of Muscle Proteins and Comparative Histology, Human Genome Research Center, Biosciences Institute, University of Sao Paulo, Brazil; \textsuperscript{8}CH Sainte-Anne, Neuropathology Department, Paris, France.

*These authors contributed equally to this work.
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\textsuperscript{9}Present address: Centre de Génétique et de Physiologie Moléculaire et Cellulaire, Claude Bernard Lyon 1 University, Lyon, France; CNRS UMR5534, Paris, France

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Contact:

Fabrice Chrétien. Fax: +33 (0) 1 40 61 31 55. Tel: +33 (0) 1 40 61 31 44. Mail: fabrice.chretien@pasteur.fr

Institut Pasteur, Histopathologie Humaine et Modèles Animaux, 28 rue du Docteur Roux, 75015 Paris.

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ABSTRACT

Duchenne Muscular Dystrophy (DMD) is a progressive neuromuscular disease, caused by an absence of dystrophin, inevitably leading to death. Although muscle lesions are well characterised, blood vessel alterations that would have major impact on muscle regeneration, remain poorly understood. Our aim was to elucidate alterations of the vascular network organisation, taking advantage of Flk1^{GFP/+} crossed with mdx mice (model for human DMD where all blood vessels express GFP) and functional repercussions using \textit{in vivo} nuclear magnetic resonance (NMR), combining arterial spin labeling imaging of perfusion, and $^{31}$P-spectroscopy of phosphocreatine kinetics. For the first time, our study focused on old (12 month-old) mdx mice, displaying marked chronic muscle lesions, very similar to the lesions observed in human DMD, in comparison to young-adult (3 month-old) mdx mice displaying only mild muscle lesions with no fibrosis. Using an original approach combining specific animal model, state of the art histology/morphometry techniques, and functional NMR, we demonstrated (i) that the microvascular system is almost normal in young-adult in contrast to old mdx mice, displaying marked microvessel alterations, and (ii) functional repercussions on muscle perfusion and bioenergetics after a hypoxic stress, that vary depending on stage of pathology. This original approach clarifies disease evolution and paves the way for setting up new diagnostic markers or therapeutic strategies.
INTRODUCTION

Duchenne muscular dystrophy (DMD) is the most frequent genetic neuromuscular disorder affecting 1:3500 school-age boys worldwide. This X-linked muscle disease is characterised by progressive skeletal muscle weakness and cardiomyopathy, leading to premature death generally because of respiratory and/or cardiac failure. The cause of DMD is the absence of dystrophin, a key component of the dystrophin-associated protein complex involved in the linkage between myofiber cytoskeleton and extracellular matrix. When linkage is disrupted, muscle fibers develop normally but are more susceptible to damage due to mechanical stretch. Despite presence of satellite cells (muscle stem cells) and successive regeneration attempts, myofibers undergo necrosis and are eventually replaced by connective and adipose tissue.

Muscle lesions in DMD have been widely investigated, with studies focusing principally on myofibers and/or satellite cells (SC). Although (i) skeletal muscle is one of the most vascularised tissues, (ii) endothelial cells are essential in muscle regeneration process, and (iii) dystrophin is expressed in endothelial/smooth muscle cells, disease impact on blood vessels and effect of blood vessel alteration in disease expression remain poorly understood.

In recent years, interest in DMD vascular network has increased with primary focus on vasculature-related therapeutic strategies such as methods to increase vasculature by modulating VEGF/VEGFR pathways. These strategies were initially based on: (i) observation of “grouped necrosis” in muscles of DMD patients, i.e. simultaneous necrosis of contiguous myofibers, suggesting local failure in capillary blood supply and muscle ischemic necrosis, and (ii) membrane-associated nitric oxide synthase (NOS) deficiency in dystrophin-deficient muscle. The hypothesis of an ischemic process has been strongly discussed, as other studies could not detect any vascular bed abnormality in DMD either
morphologically using electron microscopy or physiologically by studying muscle blood flow. More recent studies carried out in DMD patients confronted blood vessel alteration with tissue fibrosis. They suggested that endomysial fibrosis plays an essential role, causing an increase in capillary-to-myofiber distances, which impairs both muscle fiber mechanical function and gas exchanges. Moreover, increased distances between capillaries and myofibers could potentially impede their reciprocal stimulation by soluble factors secreted during muscle repair.

Rare studies addressed the relevance of muscle vascular network in dystrophinopathy pathophysiology in animal models, focusing on muscle vascular density and characterisation of a possible hypoxic condition in dystrophic muscle. Part of these studies pointed to a decreased vascular density and an impaired angiogenesis in 6 week- to 6 month-old mdx mice the dystrophin-deficient murine model of human DMD. However, contradictory results were also published in mdx mice, showing a higher hindlimb perfusion one week after femoral artery dissection and significant increase in arteriole length density in 2 month-old animals. These discrepancies could be related in part to the effect of aging in disease progression; age appears to be an important parameter to consider when studying vascular changes. Thus, involvement of blood vessels in the pathogenesis of dystrophy is still not completely understood.

In the present study, we investigated both structural organisation and in vivo function of vascular system in young-adult (3 month-old) mdx mice, displaying only moderate subacute muscle lesions with no fibrosis, and old (12 month-old) mdx mice, displaying marked muscle lesions with persistent inflammation and fibrosis, more relevant for the study of DMD pathophysiology in human. We used complementary morphological approaches based on genetically-modified mice that allowed for the first time to reconstruct the 3-dimensional
microvascular network in mdx mouse. These were confronted to innovative histological
techniques and dynamic and non-invasive multiparametric and functional nuclear magnetic
resonance (NMR).

MATERIAL AND METHODS

Mice

C57Bl/6J control mice were obtained from Charles River Laboratory (l’Arbresle, France),
mdx-4Cv with C57Bl/6 background mice, model for human DMD, were kindly provided by
Pr. Gherardi (Hôpital Henri Mondor, France), Flk-1^{GFP+} mice, in which green fluorescent
protein (GFP) is targeted in vascular endothelial growth factor (VEGF) receptor-2 gene locus,
exhibiting a bright GFP signal in all endothelial cells, were kindly provided by A. Medvinsky
(Institute for Stem Cell Research, University of Edinburgh, UK), and Flk-1^{GFP+}:mdx-4Cv
mice were obtained by crossing Flk-1^{GFP+} with mdx-4Cv mice. Only male animals were used
i.e. young-adults (3 month-old) or old (12 month-old).

Animals were housed in animal facilities of the Institut Pasteur licensed by the French
Ministry of Agriculture and complying with European Union regulations. Protocols were
approved by the Institut Pasteur Animal Experimentation Ethics Committee (01332.01).

Microvascular network organisation in three dimensions

Young-adult and old Flk-1^{GFP+} and Flk-1^{GFP+}:mdx-4Cv mice were anesthetised with
isoflurane inhalation (Forene, Abbott, Rungis, France) and killed by cervical dislocation.
Gastrocnemius muscles were removed and imaging of vascular network was carried out in
two conditions: thick cryo-sections or whole muscle. Gastrocnemius muscle was snap frozen
in liquid nitrogen-cooled isopentane before cryosectionning (100 μm-thick sections).

Confocal acquisitions were performed using a spinning disk microscope (Leica, Wetzlar, Germany), laser femto-second was used: Chameleon Ultra, 20x/0.7 and 40x/0.75 objectives and a CoolSnap HQ2 camera. Optical slices were taken every 0.5 or 0.3 μm interval along the z-axis (80 μm).

For whole muscle conditions, images of Gastrocnemius blood vessels were obtained from the entire muscle using multi-photon scanner resonant confocal Leica TCS-SP5 with 20x/0.95 objective. Optical slices were taken every 0.5 μm along the z-axis.

**Histological/Immunohistochemical analysis**

Gastrocnemius muscles were collected from mice after NMR experiments, snap frozen in liquid nitrogen-cooled isopentane and kept at –80°C. Six different levels of 7 μm-thick sections were cut and stained with hematoxylin-eosin (HE) to describe histopathological modifications of muscle tissue, and Sirius red for visualisation of collagen. For immunohistochemistry analyses, muscle cryosections were incubated with antibodies directed against endothelial cells (anti-CD31; Pharmingen), satellite cells (anti-Pax7; DHSB, Iowa city, IA, USA), pericytes (anti-NG2; Millipore), smooth muscle cells (αSMA; Sigma) and basal lamina (anti-laminin; Sigma). Primary antibodies were incubated overnight at 4°C and revealed by cy3- or TRITC-labeled secondary antibodies (Jackson ImmunoResearch Laboratories).
Morphometric analysis

Two-dimension analysis was performed to evaluate distribution of muscle fiber diameter, percentage of centro- or peri-nucleated fibers, microvessel count and distribution around each myofiber using ImageJ (NIH, Bethesda, MD, USA) and NIS-Element (Nikon) softwares. At least 200 fibers were considered for each muscle.

Three-dimensional analysis was performed to evaluate organisation of vascular network. For each muscle, 10 z-stack image reconstructions were achieved on 80 to 150 µm-thick frozen sections. Analysis was carried out using IMARIS (ImarisBitplane, Zurich, Switzerland) software (quantification of vessel density, tortuosity, volume, anastomose count, diameter and distance between microvessels).

Quantitative RT-PCR

We used real-time PCR to determine the level of angiogenesis-related mRNA expression in young-adult and old mdx mice. Total Gastrocnemius muscle RNA was extracted using RNeasy Mini Kit (Qiagen). One µg of total RNA was reverse transcribed into first-strand cDNA using Superscript II Reverse Transcriptase (Life technologies). Quantitative PCR was carried out on StepOne Plus RealTime PCR system (Applied Biosystems, Carlsbad, CA, USA). Reaction mixtures had a final volume of 20 µl, consisting of 1 µl of cDNA, 10 µl of Sybr Green Master (Roche) and 10 µM of primers, listed in Table 1. After initial denaturation, amplification was performed at 95°C (10 s), 60°C (5 s), 72°C (10 s) for 45 cycles. Calculation of relative expression was determined by the StepOnePlus software (Applied Biosystems) and fold change was normalized to 18S rRNA housekeeping gene.
Nuclear Magnetic Resonance analysis

NMR experiments were performed on: 3 month-old mdx-4Cv (n=6) and control C57Bl/6J (n=9) and on 12 month-old mdx-4Cv (n=5) and control C57Bl/6J (n=7).

Hyperaemic response paradigm: To highlight differences between normal and altered muscles we classically applied a stress to increase the global need for perfusion. Ischemia-reperfusion stress was applied to the mouse left hindlimb which provokes maximal vasodilatation and limited resistance of arteries/arterioles after tourniquet release.

In practice, anaesthesia was induced with 4% isoflurane delivered in 1.5 L/min air and maintained with 1.75% isoflurane. During experiments, a water heating pad ensured a constant temperature of 37°C and breathing was monitored. After a 24 min NMR acquisition at rest (baseline), ischemia of the leg was induced by occlusion of femoral artery by two surgical threads placed around the thigh and pulled tight by application of a weight. After 30 min of ischemia, the weight was instantly removed, inducing a hyperaemic response which was monitored over the next 30 min. During whole protocol, dynamic acquisitions of NMR scans of interleaved perfusion imaging and $^{31}$P-spectroscopy ($^{31}$P-NMRS) were collected.

Multiparametric functional NMR (mpf-NMR) acquisitions. In vivo NMR experiments were conducted in a 4 Tesla Biospec system equipped with a 20 cm diameter 200 mT.m$^{-1}$ gradient insert (BrukerBioSpin MRI GmbH, Ettingen, Germany). Mice were placed supine in a 6 cm diameter, 12 cm length volume transmitter $^1$H coil for whole-body signal excitation. An actively decoupled 2 cm diameter surface $^1$H coil, positioned below the left calf, was used for image signal reception. Muscle metabolites were probed by a 10 mm $^{31}$P saddle-shaped coil placed around the left leg.
As described in detail elsewhere, Arterial Spin Labeling (ASL)-NMR imaging and $^{31}$P-NMR acquisitions were interleaved using the dedicated Bruker MultiScanControl software (BrukerBioSpin GmbH) in order to follow simultaneously and non-invasively: (i) muscle perfusion signal by SATuration-Inversion Recovery (SATIR) (time resolution: 10 sec), and (ii) mitochondrial activity by dynamic $^{31}$P-NMRS (time resolution: 2.5 sec). In brief, ASL imaging is based on non-invasive alternate magnetic tagging of blood water spins to provide endogenous markers of muscle perfusion, measured in regions of interest (ROI) drawn in posterior compartment of the leg. Muscle bioenergetics and pH were assessed from ratios of energetic phosphates measurable by $^{31}$P-NMRS at rest, *in vivo* mitochondrial oxidative capacity was directly assessed from the rate of creatine rephosphorylation at the end of ischemia, and intramuscular pH was calculated from chemical shift between phosphocreatine (PCr) and inorganic phosphate (Pi). A minimum of 50% PCr depletion at the end of ischemia was necessary to reliably measure dynamics for PCr recovery, and examinations which did not reach this threshold were rejected.

**NMR perfusion analysis.** Images were acquired after positive or negative labeling alternately. To avoid large vessels, ROI were drawn in the posterior compartment of the leg. Muscle perfusion $f$ was calculated from the normalized difference between consecutive images according to the equation:

$$f = -\frac{\lambda}{T_{ev}} \times \ln \left[ \frac{(M^+ - M^-)}{(M^+ + M^-) \times (1 - \exp(r_1 T_{ev})) + 1} \right]$$

where $r_1$ is the longitudinal relaxation rate for muscle (measured by saturation-recovery acquisition for each mouse at the end of 30 minutes hyperaemic period), $M^+$ and $M^-$ are the signals of positive and negative labelled perfusion images and $\lambda$ is the blood-tissue partition coefficient ($\lambda = 0.9$).
**$^{31}$P-NMR Spectroscopy analysis.** Successive $^{31}$P Free Induction Decays were acquired throughout rest, ischemia and hyperaemia. $^{31}$P-spectroscopy gives access to principal metabolites implicated in energetic metabolism such as phosphocreatine (PCr), the three $\alpha$, $\beta$, $\gamma$ ATP and inorganic phosphate (Pi). Signal intensity of these resonances is directly proportional to their concentrations, which allows the quantitative following of these metabolite variations.

At ischemia and recovery, PCr recovery was fitted by a mono-exponential function with a least mean squares algorithm and pH was calculated from the chemical shift $\delta_{\text{Pi}}$ between PCr and Pi according to the formula $^20$:

$$pH = 6.75 + \log \left( \frac{3.27 - \delta_{\text{Pi}}}{(\delta_{\text{Pi}} - 5.69)} \right)$$

**Statistics**

Perfusion data were analysed by repeated measurements ANOVA. Analyses were performed with NCSS-2007 software (Kaysville, UT, USA). Group comparisons for perfusion parameters and phosphorus spectroscopy analysis were performed using Mann-Whitney test. Statistical analysis of histological data was performed with GraphPad-Prism software (La Jolla, CA, USA). Fiber diameter repartition was evaluated by a chi-square test followed by a multi-t-test corrected for multiple comparisons using Holm-Sidak method. Same multi-t-test was used to evaluate capillary count/fiber repartition.

Statistical significance was taken at $p<0.05$ and $p$-values indicated on figures are $^*p<0.05$, $^{**}p <0.01$, and $^{***}p<0.001$. Numerical NMR and histological data are reported as mean±SD.
**RESULTS**

Microvessels were defined as the small blood vessels located at the periphery of myofibers, in the endomysium, displaying a diameter of less than 20 µm and a wall sometimes containing one layer of αSMA-expressing cells, thus including capillaries, terminal arterioles and terminal venules.21

**Young-adult Flk1<sup>GFP</sup>+/::mdx mice display a normal microvascular network organisation but a mild decrease in terminal arteriole density.**

Polyphasic subacute lesions, characterised by small inflammatory infiltrates and centrally nucleated fibers, were observed in Gastrocnemius muscle (Figure 1A-B). Surprisingly, these lesions had no impact on blood microvascular network organisation. In both Flk1<sup>GFP</sup>+/ and Flk1<sup>GFP</sup>+/::mdx mice, vascular network was indeed well organised with straight microvessels located along myofibers, parallel to each other with few anastomoses oriented perpendicularly to myofibers (Figure 1C-F). Microvessel diameter, measured using diameter of endothelial cell fluorescence, was similar in both groups (13-14 µm), as well as anastomose count (1,200-1,650 anastomoses/mm³) (Figure 1G-H).

Immunofluorescence analyses did not detect any significant difference between young-adult wild-type and mdx mice (Figure 2). Both muscles displayed the same myofiber cross-section diameter, fiber size distribution, and microvessel density, quantified by microvessel count per fiber. No macrovascular modification was detected either (data not shown). To characterise further the microvascular network and identify terminal arterioles, we carried out an immunohistochemistry against αSMA, highlighting perivascular smooth muscle cells. We quantified a 26% loss of αSMA expression in mdx mice, suggesting a decrease in terminal arteriole density (Figure 3I, J, L). Collectively, these results highlighted a normal
microvascular network organisation in muscles of both groups, but a mild decrease in
terminal arteriole density, in mdx mice.

Capillary-to-fiber perimeter exchange index (CFPE) has been used to calculate the contact
surface area between capillaries and myofibers. It provides an indirect quantitative criterion
to evaluate movement of oxygen from capillaries to muscle fibers. CFPE index was not
affected in young-adult mdx mice (Figure 2F).

As almost no alteration of the microvascular network was detected at the morphological
level, we investigated the expression of angiogenesis-related mRNA (VEGF and its receptors
Flk1 and Flt1, CD31, Ang1, Ang2 and Tie1, Tie2 receptors and nNOS) (Figure 4A). We did
not observe any significant modification of these mRNA expression in young-adult mdx
mice, suggesting no stimulation of the angiogenesis process. In contrast, nNOS expression
was significantly decreased in young-adult mdx mice.

Similar pericyte density but increase in satellite cell count in young-adult mdx mice.

Using immunohistochemistry analysis, we focused on important partners of endothelial cells:
pericytes and satellite cells. Concerning pericytes (NG2+ cells located at the periphery of
blood vessels in muscle sections), no difference in density per mm² was detected between
wt (151.6±14.3 pericytes/mm², n=5) and mdx (154.3±21.2 pericytes/mm², n=5) mice (Figure
3I-K). Satellite cells (SC; Pax7+ cells) are in a close relationship with endothelial cells and
coupling between myogenesis and angiogenesis takes place concomitantly during muscle
regeneration. In young-adult mdx mice, we demonstrated using immunofluorescence an
increase in SC count per mm² (wt: 14.5±0.3 SC/mm², n=4; mdx: 29.7±3.5 SC/mm², n=6; p<0.01) and per fibre (wt: 0.04±0.01 SC/fibre, n=4; mdx: 0.07±0.01 SC/fibre, n=5; p<0.05),
in comparison to wt (Figure 3A-D).
Muscle blood perfusion is modified in young-adult mdx mice.

In accordance with our previous observations, profiles of reactive hyperaemia were significantly different in mdx (n=6) and wt (n=9) mice (Figure 5A, p<10⁻⁶ with ANOVA). The release of ischemia provoked an instantaneous increase of perfusion which was lower in wt mice (mdx: 78.7±27.1 ml/min/100 g; wt: 41.3±32.3 ml/min/100 g, 20 s post-release). In wt mice, this first perfusion peak was followed by a drop to reach a plateau around a value of 26.6±9.2 ml/min/100 g peaking at 270 s post-ischemia. In contrast, mdx muscle perfusion slightly increased to a mean perfusion value of 84.8±24.8 ml/min/100 g at 300 s post-ischemia and reduced to 26.3±25.9 ml/min/100 g only 850 s after stress release (Figure 5A). Moreover, the global volume repaid after ischemia was significantly higher in mdx mice (wt: 474.3±216.3 ml/100 g; mdx: 1017.0±369.2 ml/100 g, p<0.05). The response to ischemic stress was therefore different and enhanced in young-adult mdx mice while almost no morphological modification of microvessels was detected.

Muscle bioenergetics in young-adult mice (Table 2).

At rest, mdx mice displayed a slightly higher Pi/PCr ratio compared to wt which reflects an increase in ADP concentration in dystrophic mice. In addition, a lower PCr/ATP ratio was observed in mdx mice, reflecting a decrease in metabolically functional muscle tissue.

The 30 min ischemic stress induced a higher Pi/PCr ratio in mdx mice while PCr depletion tended to be accelerated compared to wt (mdx: ΔPCr = 65±9%; wt: ΔPCr = 58±6%; p=0.09). At reactive hyperaemia, release revealed a significant acceleration of PCr resynthesis rate in mdx mice compared to wt, reflecting higher mitochondrial ATP production in the mdx.
However, contrarily to wt, combined $^{31}$P-NMRS and perfusion results showed a very tight correlation between time of rephosphorylation τPCr and various parameters reflecting perfusion: maximum perfusion ($r^2 = 0.66$, p<0.05), time-perfusion integrals ($r^2 = 0.99$, p<0.001 until 30 sec; $r^2>0.93$, p<0.01 until 150s) (Figure 6). In wt, none of the correlations between τPCr and perfusion were significant.

In summary, phosphate metabolism was accelerated during ischemia in 3 month-old mdx mice. At recovery, mitochondrial oxidative rephosphorylation was unexpectedly faster and perfusion was increased in comparison to age-matched control mice. Moreover, perfusion in mdx was directly related to mitochondrial ATP production. This is unlike normal healthy case where a luxury perfusion is observed and is neither limiting nor correlated to τPCr.

Alteration of microvascular network organisation in old Flk1$^{GFP/+}$::mdx mouse.

Old mdx mice displayed marked histological lesions; some already observed in young-adult as anisocytosis or centrally nucleated myofibers, others included persistence of chronic inflammation, and presence of endomysial/perimysial fibrosis (Figure 7A-D). The microvascular network was as well organised in old Flk1$^{GFP/+}$ as in young-adult mice (Figure 7E). In contrast, Flk1$^{GFP/+}$::mdx mice displayed significant alterations, characterised by a marked increase in tortuosity and irregular scattering of microvessels (Figure 7F).

Microvessel diameter was similar in both groups (12 µm), but we identified a higher anastomose count, from more than 50,000 anastomoses/mm$^3$ for Flk1$^{GFP/+}$::mdx mice to less than 1,000 anastomoses/mm$^3$ for control Flk1$^{GFP/+}$ (p<0.01) (Figure 7J). Collectively, these results pointed to an anarchic blood vessel organisation in this context of dystrophinopathy.

Immunofluorescence analyses showed that (i) myofiber cross-section mean diameter was smaller in mdx mice (mdx: 47.4±4.2 µm; wt: 61.2±3.9 µm; p<0.001) (Figure 8C), (ii) the
smaller myofibers (<50µm) were clearly under-vascularised (Figure 8E), (iii) perinucleated (Figure 8G), and (iv) represented almost 60% of total muscle fibers in mdx mice in contrast to 35% in controls (Figure 8D). Alterations were also detected at the terminal arteriole level, as a loss of 70% of αSMA expression was detected in old mdx mice (wt: 88.6±2.4%, n=7; mdx: 18.0±2.1%, n=8; p<0.001), suggesting a marked decrease in terminal arteriole density (Figure 3M, N, P). These data suggested either a progressive degradation of tissue with no maintenance of microvascular network with time or a defect of neo-angiogenesis. The CFPE was not affected in old mice (Figure 8F).

Angiogenesis-related mRNA expression analysis revealed a collapse of VEGF expression and its decoy receptor Flt1 (Figure 4B). As observed in young-adult mice, nNOS was also significantly decreased in old mdx mice.

Collectively, these results pointed out severe alterations of microvessel organisation, especially around small/atrophic myofibers, associated with alteration of angiogenesis, suggesting chronic alteration of endothelial-myogenic cell interface.

Pericytes and satellite cells in old mdx mice.

The density of pericytes was similar between wt and mdx mice but, in contrast to young-adult mdx mice, a decrease in satellite cell count was observed for old mdx mice (wt: 12.4±0.6 SC/mm², n=8; mdx: 10.0±0.8 SC/mm², n=9; p<0.05) (Figure 3E-H).

Alteration of muscle perfusion in old mdx mice.

Despite severe alterations in mdx muscle microvascular network organisation, at rest, no difference in muscle perfusion was observed between mdx and wt mice (mdx: 12.09±5.90
ml/min/100 g; wt: 8.19±2.19 ml/min/100 g). After tourniquet release, rapid increase of
perfusion was detected in muscles of the posterior hindlimb compartment; this increase was
significantly lower in old mdx mice (mdx perfusion maximal value at 380 s post-ischemia:
60.5±39.3 ml/min/100 g; wt perfusion maximal value at 400 s post-ischemia: 106.1±38.1
ml/min/100 g, p<0.05), in contrast to what was seen in young-adult mice.

Analysis of variance of perfusion time-courses demonstrated differing profiles between wt
and mdx mice (p<10^-4), with specific differences in the early phase of reperfusion. A similar
initial peak of perfusion, as the one observed in young-adult wt mice, was detected in the old
wt group, 20 s post-ischemia, but was absent in mdx mice (Figure 5B).

Thus at 12 months, both mdx and wt showed different profiles from young-adult animals
(ANOVA, p<10^-6), and in contrast to wt and young-adult mdx, old mdx mice displayed a
decrease in muscle perfusion and a modified perfusion profile after an ischemic stress.

Muscle bioenergetics in 12 month-old mice (Table 3).

At rest, no difference in pH was observed between wt (n=7) and mdx (n=5) mice but hypoxic
stress induced a significant acidosis in both groups (p<0.005), more pronounced in mdx (wt:
ΔpH = 0.22±0.06; mdx: ΔpH = 0.30±0.04; p<0.05). Ischemia was associated with a
significant increase in PCr depletion in old dystrophic mice compared to wt, though the
difference in Pi/PCr ratio between the two groups was not significant. Unlike in young-adult
mice, the rephosphorylation rate was comparable in both groups. Indeed τPCr was shorter in
the old compared to the young-adult wt mice (p<0.01), but was unchanged with age in the
mdx mice. Thus no alteration of oxidative capacities was observed in old mdx mice in
response to hypoxic stress compared to age-matched control mice, despite reduced perfusion.

In older mice (wt and mdx), no correlation was found between τPCr and perfusion variables.
DISCUSSION

Our study deciphers lesions of muscle microvascular network, in a model of dystrophinopathy, using combination of state of the art histology/morphometry techniques and totally non-invasive functional approach. This experimental paradigm, combining histopathology and mpf-NMR, clearly relevant for clinical diagnosis and research, allowed us to associate for the first time the fine 3D-alterations of muscle microvascular network with functional repercussions on muscle.

Concerning the animal model, previous studies used 6 week to 6 month-old mdx mice, which display very few chronic lesions with no fibrosis, in contrast to what happens in human. We therefore worked on 12 month-old mdx mice, displaying persistence of endomysial inflammation and fibrosis, more representative of human DMD and thus more relevant for chronic myopathy and DMD pathophysiology study, in contrast to young-adult mdx mice displaying no chronic lesions. We demonstrated (i) strong alterations of microvascular network structure associated with reduced muscle perfusion in old mdx mice, (ii) functional increase in muscle perfusion and mitochondrial oxidative phosphorylation with normal microvascular network organisation in young-adult mdx mice, and (iii) a different impact of age on wild-type and mdx mouse muscles.

In young-adult and old wild-type mice, no alteration could be detected in muscle histology or microvascular network organisation. Perfusion is primarily regulated by smooth muscles that control blood flow distribution and capillary recruitment. Capillary resistance, at rest, plays only a minor role in perfusion regulation. Using ischemia-reperfusion, we provoke maximal arteriolar dilatation in order to limit arteriolar resistance, and thus microvessel network becomes predominant in control of muscle perfusion. Analysis of perfusion profiles
revealed the existence of a “peak” of perfusion in the first seconds after ischemia release, for young-adult and old wild-type mice. This initial “peak” suggests specific regulation of perfusion in early phase after ischemia release, probably coordinated by perivascular smooth muscles and/or pericytes. 

Surprisingly and in contrast to previous studies describing decreased vascular density, we did not detect any alteration in vascular network 3D-organisation in young-adult mdx mice. However, muscle post-ischemic perfusion was higher than in aged-match control mice, and time resolution of mpf-NMR allowed to demonstrate the absence of the initial “peak” of perfusion. In the same time, we also demonstrated a 26% loss of αSMA expression in young-adult mdx muscle (and more than 70% loss in old mdx), suggesting a drop in perivascular smooth muscle cells, responsible for part of these deleterious effects. It has indeed been demonstrated in vivo that the re-expression of dystrophin only in smooth muscle cells significantly ameliorates vasoregulation in mdx mice confirming the importance of perivascular cells (smooth muscle for example) in blood flow regulation. One of the possible key factor is NO production alteration or impairment of neuronal NO synthase (nNOS), very probably explaining the significant decrease in nNOS expression both in young-adult and old mdx mice, in our study. Concerning pericytes, Yemisci et al. demonstrated in the brain, after a 2h ischemic stress, that pericytes remain contracted despite successful re-opening of blood flow, impairing microcirculatory reflow. These experiments were carried out ex vivo, and no functional in vivo validation was done. Our data seem therefore to highlight functional alterations of smooth muscles and/or pericytes after ischemic stress. This alteration is severe enough to significantly impact perfusion profiles between control and mdx mice, and we are currently carrying out new experiments to better understand the effect of an absence of dystrophin in perivascular cells and their involvement in dystrophinopathy pathophysiology.
Old Flk1^{GFP^+}::mdx model allowed us to highlight disorganisation of microvascular network. A marked increase in microvessel tortuosity, an irregular scattering, and an increase in anastomose count were observed. Existence of these highly abundant anastomoses, suggests that “radial” as well as “longitudinal” (parallel to myofibers) blood flow is important, and in turn, that “longitudinal” flow is abnormally heterogeneous, microvessel longitudinal resistance being likely to vary a lot from microvessel to microvessel which would be the driven force for collateral flow.

Considering the close association between microvessels and myofibers, we demonstrated in old mdx mice that more than 60% of myofibres were atrophic with peripheral nuclei and displayed less microvessels at their periphery, resulting in a global undercapillarisation and loss of terminal arterioles. In parallel, NMR analysis revealed a two-fold decrease in perfusion after ischemia release. The significant decrease of microvessel (capillary and terminal arteriole) density around small myofibers is likely responsible for these functional alterations. Our data are thus in agreement with other studies demonstrating the effect of age on dystrophinopathy pathophysiology. Our hypothesis is that interaction between angiogenesis and myogenesis could be affected in old mdx mice; the increasingly scarce microvessels would provoke an alteration of myofiber regeneration that in turn could lead to impairment of remaining microvascular network support, maintaining a vicious circle. With this idea, we focused on the dynamic of satellite cell density in the muscle tissue. While SC density in young-adult mdx mice was increased, very probably because of the stimulation of muscle regeneration, it collapsed in old mdx mice, with pathology evolution, suggesting a worsening of the situation and an increase in the severity of chronic muscle lesions. Decline in SC number and activity has already been observed with age in mdx mice, in association with attenuated Notch signalling transduction. Christov et al. already introduced the idea that angiogenesis and myogenesis are coupled during muscle regeneration, these processes
involving several growth factors, such as VEGF. In our study, we also observed modifications of VEGF and its receptor Flt1 expression profiles. Flt1 is the decoy receptor of VEGF, acting as a negative regulator of endothelial cell growth and differentiation. A previous study demonstrated that mdx mice knock out for the Flt1 receptor (mdx:Flt1−/−) presented an improved muscle histology associated with a better muscle perfusion and force production compared to mdx mice. These data underlines the link between vascular remodelling and muscle regeneration, even in severe chronic diseases. Concerning VEGF, the 2-fold decreased expression, detected in old mdx mice, was in accordance with previous studies demonstrating that treatment with VEGF strongly ameliorates mdx phenotype, with improvement of functional parameters, increase in capillary density, improved muscle regeneration, and decrease in interstitial fibrosis. Fibrosis is incidentally a key parameter influencing perfusion, and is increased in DMD. However, fibrosis might not be the most limiting factor to perfusion, as it only represents 10% of old mdx mice muscle tissue in our study. Moreover, distance between capillaries and myofibers, generally modified with endomysial fibrosis, is similar between young-adult and old wild-type and mdx mice.

The in vivo increase of post-ischemic muscle perfusion with old age in wt animals was found to be reproducible in different mouse strains, but no explanation is currently put forward, while effect of aging on perfusion is still debated even in humans.

In parallel to perfusion analysis, acquisition of 31P-spectroscopy revealed moderate energetic metabolism alterations, in agreement with previous literature, and contrarily to what might be expected from alterations of enzymatic activities or defects of mitochondrial localization in vitro. Compensatory mechanisms must thus exist in dystrophic muscle.

As for perfusion, anomalies in phosphorus metabolites in wild-type animals depended on age, as already evoked in early studies of mdx metabolism. Anomalies at rest were more
marked in younger mdx: Pi/PCr reflecting resting ADP production was increased while
PCr/ATP proportional to functional muscle was reduced (-11%), coherently with other
reports 40-42 (-50% or more in DMD children 44).
Unlike a recent study in 3 month-old mdx mice 45, where 10min ischemia was used as stress
protocol, we found greater depletion after 30min ischemia in both young-adult and old mdx.
During prolonged ischemia, two energetic pathways are activated to supply ATP demand:
ATP-PCr system and glycolytic pathway. Production of ATP directly from PCr consumption
is very small; it is therefore unlikely that PCr would be consumed to compensate for
defective glycolytic pathway. The higher depletion observed in both young-adult and old
mdx could more likely reflect a higher ATP demand to maintain ionic homeostasis.
Confrontation between perfusion and metabolic data obtained simultaneously by NMR
revealed that despite strongly reduced perfusion in old mdx mice, oxidative metabolism was
preserved, suggesting existence of a “luxury perfusion”, i.e. reserve of perfusion that can be
eliminated without impact on muscle physiology, as previously evidenced in a model of
peripheral artery disease 46. It generally explains the absence, or very loose correlation,
between perfusion and PCr recovery rates in wt animals and in old mdx. This contrasts with
the tight correlation between initial perfusion and τPCr of young-adult mdx mice, which
display faster τPCr recovery and stronger perfusion than controls. Using optical spectroscopy
to analyse myoglobin and haemoglobin oxygen desaturation in parallel to 31P-NMRS,
Percival observed strong uncoupling between ATP synthesis and O₂ consumption in 4 month-
old mdx, dystrophic muscle producing 39% ATP less per O₂ consumed than controls 45. We
might thus hypothesize that at this young age, increased perfusion might be a means to
compensate for mitochondrial inefficiency.
In conclusion, we demonstrated strong structural and functional alterations in muscle microvascular network of dystrophin-deficient mdx mice, with an increasing severity in parallel to aging. Our approach combining 3D-morphological analyses with non-invasive functional evaluation, allowed to better characterise the impact of histological lesions on tissue function. Collectively, our data pointed out that vascular network has a key role in dystrophinopathy pathophysiology and would be very important target for the set-up of new innovative therapeutic strategies.

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Statement of author contributions:

GJ, FC, CW, CL and PGC designed experiments. CL, BM, AMB and DB carried out experiments and GJ, FC, PGC, CW, BM and CL analysed data. GJ, CL, BC, BM and CW were involved in writing the paper and all authors gave final approval of the submitted and published versions.

List of abbreviations:
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34. Fong GH, Rossant J, Gertsenstein M, Breitman ML: Role of the Flt-1 receptor tyrosine kinase in regulating the assembly of vascular endothelium, Nature 1995, 376:66-70


46. Vidal G WC, Giacomini E, Emmanuel F, Carlier PG: A truly non-invasive set-up for the study of perfusion and energy metabolism in the rat calf in vivo: application to a model of peripheral arterial disease (Abstract), Magma 2002, 25:15
FIGURE LEGENDS

FIGURE 1. Normal microvascular network organisation in 3 month-old mdx mice.
In contrast to wild-type mice (A), mdx mice (B) display subacute lesions in Gastrocnemius muscle, characterised by small inflammatory infiltrates (star) associated with regenerated myofibers displaying central nuclei (arrows) (HE staining).

Microvessel 3D organisation of Flk1^{GFP/+} (C,E) and Flk1^{GFP/+}:mdx (D,F) mice: normal blood microvessel organisation, with microvessels regularly scattered along myofibers (C-F) (Scale bars: 50 µm). Morphometric analyses revealed similar diameter (G) and anastomose count/mm³ (H) between microvessels from wild-type and mdx mice.

FIGURE 2. Microvessel morphometry in 3 month-old mice.
Young-adult wild-type (n=5) and mdx (n=3) mice display similar: microvessel distribution in Gastrocnemius muscle (A,B) (laminin-FITC and CD31-TRITC immunohistochemistry to label basal lamina (green) and blood vessels (red); Scale bar: 50 µm), fiber size repartition (C), microvessel count per fiber (D), microvessel diameter (E), and capillary to fiber perimeter exchange index (CFPE) (F).

FIGURE 3. Satellite cells, terminal arterioles and pericytes are affected in mdx mice.
Pool of SC was analysed by immunohistochemistry (A, B, E, F, white arrows) (Pax7-FITC and Laminin-Cy3). SC density is increased in young-adult mdx mice (C, D) while we observed a significant depletion of the SC pool in old mdx mice (G, H). Pericyte density, assessed using NG2 immunolabeling (NG2-FITC), was similar for all groups, at all ages (I-
We also evaluated the expression of \( \alpha \text{SMA} \) (Smooth Muscle Actin) by perivascular smooth muscle cells, surrounding terminal arterioles (\( \alpha \text{SMA-Cy3} \)). We observed a decrease of 26\% \( \alpha \text{SMA} \) expression for young-adult mdx mice in comparison to wt animals (I, J, L), reaching 70\% loss for old mdx (M, N, P), suggesting a marked drop in terminal arterioles, increasing with the disease progression (scale bars: 25 \( \mu m \)) (*p<0.05, **p<0.01, ***p<0.001).

**FIGURE 4.** Analysis of angiogenesis-related gene expression by RT-qPCR.

Total RNA was extracted and complementary DNA was analysed by qPCR. Gene expression was measured in the Gastrocnemius of young-adult (A) and old (B) mdx mice. Results were normalized relative to the expression of the 18s rRNA housekeeping gene. Data are presented as fold change mean±SEM. Symbol ** indicates statistical difference (p<0.01) observed between mdx and wild-type mice from the same age.

**FIGURE 5.** Muscle blood perfusion during ischemia-reperfusion.

After release of ischemia, a rapid and important increase in perfusion is detected.

(A) Different profiles of perfusion are obtained in young-adult mice: total perfusion is higher in mdx mice and a first “peak” of perfusion followed by a rapid decrease in muscle perfusion is only detected in wild-type mice.

(B) A first “peak”, similar to what is observed in young-adult wild-type mice, is also observed in 12 month-old control animals. This first “peak” does not exist in mdx mice that
also displayed a reduced perfusion, with a maximum perfusion equivalent to half the value observed in wild-type mice during the hyperaemia phase.

As the release of ischemia induced movements of the leg, images affected by these movement artefacts are removed from analysis of muscle perfusion.

**FIGURE 6.** Correlation between perfusion and time of creatine rephosphorylation ($\tau$PCr) in 3 month-old mice.

Correlation between $\tau$PCr and maximal perfusion (A) or time-perfusion integrals until 30 sec (B) is significant in mdx mice (*p<0.05 and ***p<0.001, respectively) with a coefficient of determination $r^2$ of 0.66 and 0.99, respectively. In wt, none of the correlations were significant.

**FIGURE 7.** Alteration of microvascular network in 12 month-old mdx mice.

Twelve month-old wild-type mice display histologically normal muscles (A), with no fibrosis (C), and regularly scattered microvessels along myofibers, with few anastomoses (E,G,J). In contrast, 12 month-old mdx mice display chronic histological lesions (B), characterised by multifocal inflammatory infiltrates (mostly macrophages), included in endomysial collagen tissue (fibrosis; stars), associated with a marked variation in myofiber size (anisocytosis) and the presence of atrophic and regenerating myofibers displaying centrally-located nuclei (arrows). Sirius red staining and fluorescence microscopy reveal a moderate to marked endomysial fibrosis (D) and microvascular network alterations (F,H), characterised by irregularly scattered tortuous microvessels (Scale bars: 50 µm). Even if microvessel diameter is similar between mdx and wild-type mice (I), a clear increase in anastomose count/mm$^3$ is
detected for old mdx mice (J). A, B: HE staining. C, D: Sirius red staining (specific for collagen staining). **p<0.01.

**FIGURE 8.** Twelve month-old mdx mice display atrophic myofibers with low capillarisation and terminal arteriole density.

Anisocytosis is more pronounced in old mdx mice, with the presence of small atrophic myofibers (A-C) (Laminin-FITC and CD31-TRITC immunohistochemistry; Scale bars: 50 µm). Atrophic myofibers (with a diameter up to 60 µm) represent more than 60% of the total myofibers in mdx mice (D). These atrophic myofibers display less microvessels at their periphery (E). The distance between microvessels and myofibers, calculated using the capillary to fiber perimeter exchange index (CFPE) is similar in both groups (F). Small myofibers with low capillarisation are mostly perinucleated myofibers (G) (**p<0.01, ***p<0.001).
Table 1. Oligonucleotide Primers Used for qPCR

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>18S rRNA F</td>
<td>5'-CGGACAGGATGTGACGATTG-3'</td>
</tr>
<tr>
<td>18S rRNA R</td>
<td>5'-CAATCGCTCCACCAACTAA-3'</td>
</tr>
<tr>
<td>Flk1 F</td>
<td>5'-CAGTGGTACTGCGCAGCTAGAAG-3'</td>
</tr>
<tr>
<td>Flk1 R</td>
<td>5'-ACAAGCATAACGGGCTTGTGTTT-3'</td>
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<td>Flt1 F</td>
<td>5'-GGCCCGGGATATTTTATAAAGAC-3'</td>
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<tr>
<td>Flt1 R</td>
<td>5'-CCATCCATTTTAGGGGAAGTC-3'</td>
</tr>
<tr>
<td>VEGF F</td>
<td>5'-GCGGTGTGGTGGTGACATGGTT-3'</td>
</tr>
<tr>
<td>VEGF R</td>
<td>5'-ACCTCACAAAGCCAGCACA-3'</td>
</tr>
<tr>
<td>CD31 F</td>
<td>5'-CGGTGTTTCAGCGAGATCC-3'</td>
</tr>
<tr>
<td>CD31 R</td>
<td>5'-ACTCGACAGGATGGAAATC-3'</td>
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<td>Ang1 F</td>
<td>5'-GACAGTAATACAACACCGGGAAGA-3'</td>
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<td>Ang2 F</td>
<td>5'-ACTACGACGACTCAGTGCAAAG-3'</td>
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<td>Tie1 F</td>
<td>5'-AGGGCAGCTCCAGAGTATG-3'</td>
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Tie1 R  5'-GGTTGGCCAGCAATGTTAAG-3'
Tie2 F  5'-GGCTATAAGGATACGGACCATA-3'
Tie2 R  5'-TCCCCTGTCCACGGTCATA-3'
nNOS F  5'-GGCGTTCGTGATTACTGTGA-3'
nNOS R  5'-TCTTCCTCATGTCCAAATCCA-3'

741

742
Table 2. Energetic metabolism analysis from $^{31}$P-spectroscopy in young-adult mice.

<table>
<thead>
<tr>
<th></th>
<th>wt (n=9)</th>
<th>mdx (n=6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta$PCr (%)</td>
<td>58 ± 6</td>
<td>65 ± 9</td>
</tr>
<tr>
<td>$\tau$PCr (s)</td>
<td>118 ± 34</td>
<td>76 ± 34*</td>
</tr>
<tr>
<td>pH at rest (pH$_{\text{rest}}$)</td>
<td>7.20 ± 0.04</td>
<td>7.17 ± 0.03</td>
</tr>
<tr>
<td>pH end ischemia (pH$_{\text{end}}$)</td>
<td>7.00 ± 0.08</td>
<td>6.98 ± 0.03</td>
</tr>
<tr>
<td>Pi/PCr at rest (Pi/PCr$_{\text{rest}}$)</td>
<td>0.08 ± 0.03</td>
<td>0.10 ± 0.01*</td>
</tr>
<tr>
<td>Pi/PCr end ischemia (Pi/PCr$_{\text{end}}$)</td>
<td>0.93 ± 0.25</td>
<td>1.71 ± 0.52*</td>
</tr>
<tr>
<td>PCr/ATP$<em>{\gamma}$ at rest (PCr/ATP$</em>{\gamma\text{rest}}$)</td>
<td>3.39 ± 0.25</td>
<td>3.01 ± 0.27*</td>
</tr>
<tr>
<td>PCr/ATP$<em>{\gamma}$ end ischemia (PCr/ATP$</em>{\gamma\text{end}}$)</td>
<td>1.80 ± 0.62</td>
<td>1.39 ± 0.51</td>
</tr>
</tbody>
</table>

Ischemia stress was sufficient as the mean depletion of phosphocreatine (PCr) for wt and mdx mice was above 50%. Pi/PCr at rest and after ischemia were higher in mdx compared to wt mice while PCr/ATP$_{\gamma}$ at rest was lower (*p<0.05).
Table 3. Energetic metabolism analysis from $^{31}$P-spectroscopy in old mice.

<table>
<thead>
<tr>
<th></th>
<th>wt (n=7)</th>
<th>mdx (n=5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta$PCr (%)</td>
<td>54 ± 4</td>
<td>63 ± 2**</td>
</tr>
<tr>
<td>$\tau$ PCr (s)</td>
<td>66 ± 25</td>
<td>80 ± 20</td>
</tr>
<tr>
<td>pH at rest (pH$_{rest}$)</td>
<td>7.16 ± 0.07</td>
<td>7.18 ± 0.04</td>
</tr>
<tr>
<td>pH end ischemia (pH$_{end}$)</td>
<td>6.94 ± 0.04</td>
<td>6.87 ± 0.04*</td>
</tr>
<tr>
<td>Pi/PCr at rest (Pi/PCr$_{rest}$)</td>
<td>0.08 ± 0.04</td>
<td>0.10 ± 0.01</td>
</tr>
<tr>
<td>Pi/PCr end ischemia (Pi/PCr$_{end}$)</td>
<td>1.24 ± 0.40</td>
<td>1.34 ± 0.20</td>
</tr>
<tr>
<td>PCr/ATP$<em>{\gamma}$ at rest (PCr/ATP$</em>{\gamma rest}$)</td>
<td>2.98 ± 0.35</td>
<td>3.05 ± 0.16</td>
</tr>
<tr>
<td>PCr/ATP$<em>{\gamma}$ end ischemia (PCr/ATP$</em>{\gamma end}$)</td>
<td>1.20 ± 0.35</td>
<td>1.36 ± 0.30</td>
</tr>
</tbody>
</table>

Ischemia stress was sufficient as the mean depletion of phosphocreatine (PCr) for wt and mdx mice was above 50%. $\Delta$PCr is higher in mdx mice; pH decreased for both wt and mdx mice after ischemia, but mdx mice suffered a more severe acidosis. Others energetic parameters did not change in our experimental conditions (*p<0.05, **p<0.01).
Satellite cells

A young-adult wt  B young-adult mdx

C SC/mm²

wt (n=4)  mdx (n=6)

D SC/iber

wt (n=4)  mdx (n=5)

Pericytes and terminal arterioles

E old wt  F old mdx

G SC/mm²

wt (n=8)  mdx (n=9)

H SC/iber

wt (n=6)  mdx (n=5)

I young-adult wt  J young-adult mdx

K Pericytes/mm²

wt (n=5)  mdx (n=5)

L % NG2+ (terminal arterioles)

wt (n=4)  mdx (n=5)

M old wt  N old mdx

O Pericytes/mm²

wt (n=7)  mdx (n=8)

P % NG2+ (terminal arterioles)

wt (n=7)  mdx (n=8)
A Correlation maximal perfusion/τPCr

Young adult wt mice

![Graph showing correlation between maximal perfusion and τPCr for young adult wt mice.](image)

Young adult mdx mice

![Graph showing correlation between maximal perfusion and τPCr for young adult mdx mice.](image)

r² = 0.66

B Correlation perfusion integral until 30 sec/τPCr

Young adult wt mice

![Graph showing correlation between perfusion integral until 30 sec and τPCr for young adult wt mice.](image)

Young adult mdx mice

![Graph showing correlation between perfusion integral until 30 sec and τPCr for young adult mdx mice.](image)

r² = 0.99

***