SCA-1 micro-heterogeneity in the fate decision of dystrophic fibro/adipogenic progenitors

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Abstract
The term micro-heterogeneity refers to non-genetic cell to cell variability observed in a bell-shaped distribution of the expression of a trait within a population. The contribution of micro-heterogeneity to physiology and pathology remains largely uncharacterised. To address such an issue, we investigated the impact of heterogeneity in skeletal muscle fibro/adipogenic progenitors (FAPs) isolated from an animal model of Duchenne muscular dystrophy (DMD), the mdx mouse. FAPs play an essential role in muscle homoeostasis. However, in pathological conditions or ageing, they are the source of intramuscular infiltrations of fibrotic or adipose tissue. By applying a multiplex flow cytometry assay, we characterised and purified from mdx muscles two FAP cell states expressing different levels of SCA-1. The two cell states are morphologically identical and repopulate each other after several growth cycles. However, they differ in their in vitro behaviour. Cells expressing higher levels of SCA-1 (SCA1-High-FAPs) differentiate more readily into adipocytes while, when exposed to a fibrogenic stimulation, increase the expression of Col1a1 and Timp1 mRNA. A transcriptomic analysis confirmed the adipogenic propensity of SCA1-High-FAPs. In addition, SCA1-High-FAPs proliferate more extensively ex vivo and display more proliferating cells in dystrophic muscles in comparison to SCA1-Low-FAPs. Adipogenesis of both FAP cell states is inhibited in vitro by leucocytes from young dystrophic mice, while leucocytes isolated from aged dystrophic mice are less effective in limiting the adipogenesis of SCA1-High-FAPs suggesting a differential regulatory effect of the microenvironment on micro-heterogeneity. Our data suggest that FAP micro-heterogeneity is modulated in pathological conditions and that this heterogeneity in turn may impact on the behaviour of interstitial mesenchymal cells in genetic diseases.

Introduction
The human body is estimated to contain approximately 200 different cell types1. However, this estimate relies on the experimental tools used to investigate cell populations and to define cell types. Developments in single-cell technologies have challenged this view revealing that cell populations often display a higher molecular heterogeneity than originally reported2–4. This new perspective raises several fundamental questions regarding the origin of cell phenotypic or molecular heterogeneity and its impact on physiological and pathological processes5. Two different types of cell heterogeneities may be distinguished, macro-heterogeneity and micro-heterogeneity. The former refers to discrete cell subpopulations, while the latter can be viewed as a spread in the distribution of the expression of a trait in an seemingly homogeneous cell population6. This kind of cell-to-cell variability has been reported to modulate some important physiological mechanisms as, for instance, the differentiation decisions of a multipotent mouse hematopoietic cell line and the asymmetric DNA segregation during muscle satellite cells (MuSCs) division7,8. However, the
involvement of micro-heterogeneity in disease is only poorly explored.

We asked whether the heterogeneity observed in muscle interstitial cells has any impact on and is affected by myopathies such as Duchenne muscular dystrophy (DMD). DMD is an X-linked genetic disease caused by mutations in the dystrophin gene. Sarcolemma instability originating from this condition leads to a chronic damage, progressive muscle wasting and deposition of ectopic tissues, such as intramuscular adipose tissue (IMAT) and fibrotic tissue. The source of ectopic tissues is a population of interstitial cells known as fibro/adipogenic progenitors (FAPs) that develop from embryonic mesenchymal cells. FAPs reside in the endomysium of skeletal muscle and are characterised by the expression of the surface proteins stem cell antigen-1 (SCA-1 also known as lymphocyte antigen 6A-2/6E-1, Ly6a) and CD140a (also known as platelet-derived growth factor receptor alpha, PDGFR-alpha). Recently, single-cell approaches have revealed heterogeneity in FAPs and have provided some evidence of its involvement in muscle development and dystrophy in mouse models.

Here, we characterise two dynamic cell states in FAPs from muscles of mdx mice and describe how this heterogeneity impacts their properties ex vivo and in vivo. We show that the two FAP cell states, characterised by different levels of SCA-1 expression, differ in differentiation and proliferation potential. Furthermore, we show that the muscle microenvironment differentially modulates the behaviour of the two FAP cell states by affecting their propensity to differentiate into adipocytes in an age-dependent manner. These results suggest that micro-heterogeneity takes part in the fate decision of a mesenchymal cell population involved in the pathogenesis of a genetic disease.

Materials and methods

Mouse strains

C57BL/6J (wild type) and C57BL10ScSn-Dmdmdx/J (mdx) mice were purchased from the Jackson Laboratories (Ben Harbor, Maine, United States). We employed young (1.5 months old) and old (15 months old) mice. Whenever possible, we balanced gender of the mice in each experiment. Animals were bred and maintained according to the standard procedures of the animal facility. The experimental procedures were conducted according to the rules of good animal experimentation 1. A.C.U.C. no. 432 of March 12 2006 and under the approval released from the Italian Ministry of Health.

Isolation of primary cells

Wild type and mdx mice were sacrificed through cervical dislocation and were washed with 70% ethanol. After an incision through the skin, hind limbs were excised and placed into cold Hank’s Balanced Salt Solution without Calcium and Magnesium (HBSS, Biowest, Nuaille, France L0605-500) supplemented with 0.2% bovine serum albumin (BSA, Applichem PanReac, Darmstadt, Germany A1391) and 100 U/ml penicillin, 100 mg/ml streptomycin (Gibco, Waltham, Massachusetts, United States 15140122). Hind limb muscles were removed from bones under sterile hood and mechanically minced using a scalpel. Minced tissue was washed with HBSS and centrifuged at 700 × g for 10 min at 4 °C. Pelleted tissue was weighted and resuspended in the enzymatic digestion mix composed by 2.4 U/ml dispase II (4 ml/g of muscles) (Roche, Basel, Switzerland 04942078001) dissolved in Dulbecco’s phosphate buffered saline (D-PBS) with Calcium and Magnesium (Biowest L0625-500), 0.01 mg/ml DNase I (Roche 04716728001) and 2 μg/ml collagenase A (Roche 10103586001). Tissue preparations were incubated at 37 °C in a water bath (in gently shaking and not in immersion) for 1 h vortexing every 30 min. Digestion was stopped by the addition of HBSS and samples were centrifuged at 700 × g for 10 min at 4 °C. Pellets of cells were resuspended in 10 ml of HBSS and filtered through a 100 μm cell strainer (Falcon, New York, United States 352360). Cell suspensions were centrifuged at 700 × g for 10 min at 4 °C, pellets were resuspended in 10 ml of HBSS and filtered through a 70 μm cell strainer (Falcon 352350). After a further centrifuge red blood cells were lysed in 1 ml of RBC lysis buffer (Santa Cruz Biotechnology, Dallas, Texas, United States sc-296258) in ice for 2.5 min. Lysis was stopped adding HBSS and cell suspensions were filtered through a 40 μm cell strainer (Falcon 352340). Cell strainers were always washed before and after the use with 5 ml of HBSS. Cell suspensions were centrifuged at 700 × g for 10 min at 4 °C and resuspended in 500 μl of Magnetic Beads Buffer (MBB) composed by D-PBS without Calcium and Magnesium, 0.5% BSA and 2 mM Ethylenediaminetetraacetic acid (EDTA). Cells were filtered through a 30 μm cell strainer (Miltenyi, Bergish Gladbach, Germany 130-041-407) which was washed 3 times with MBB. Mononuclear cells in this suspension were counted and centrifuged at 700 × g for 10 min at 4 °C. The isolation protocol proceeds with the magnetic activated cell sorting (MACS) of the CD45−CD31−cells. Pellets were resuspended in MBB and incubated with a microbead conjugated antibody against CD45 (Miltenyi 130-052-301) according to the manufacturer’s instructions. After 15 min, cells were washed with 2 ml of MBB and centrifuged. Pellets were resuspended in 500 μl of MBB and cells were separated with MS columns (Miltenyi 130-042-201) according to the manufacturer’s instructions to collect CD45−cells. Protocol proceeds with the selection of CD31−cells applying the same procedures described for CD45−cells. Next, FAPs and muscle satellite cells (MuSCs) were isolated through fluorescent
activated cell sorting (FACS). CD45−CD31− cell suspensions were incubated with the following primary antibodies for 30 min at 4°C in D-PBS without Calcium and Magnesium, 2% BSA and 2 mM EDTA at 1 × 10^6 cells per 100 μl: anti-ITGA7 APC 1:500 (Invitrogen, Waltham, Massachusetts, United States MA5-23555) and anti-SCA-1 FITC 1:50 (BD Pharmingen, Franklin Lakes, New Jersey, United States 557405). Gates were strictly designed using proper fluorescence minus one (FMO) controls for each antibody. Debris and cell clusters were excluded using side scatter (SSC) and forward scatter (FSC) as shown in supplementary figures. Moreover, we applied two criteria to enhance the reproducibility of the sorting strategy of FAP cell states: FAP cell states must represent at least 80% of the total FAPs (these criteria also increase the yield of the isolation); the mean intensity of SCA-1 must differ of at least 3-fold between the two cell states. FAPs and MuSCs were sorted directly in their growth media as ITGA7−SCA-1− and ITGA7+ cells respectively. The purity of sorted cells was assessed immediately after sorting.

Sorting was performed on FACSARia™ III (Becton Dickinson) cell sorter equipped with 5 lasers (18 parameters). Analysis was carried out using FlowJo™ Software.

FAP cell states in vivo transplantation

A total of 75,000 freshly isolated SCA1-High-FAPs and SCA1-Low-FAPs were separately stained by incubating in the cell proliferation dye eFluor 670 (eBioscience 65-0840-85) following the manufacturer’s instruction. SCA1-High-FAPs and SCA1-Low-FAPs were then resuspended in 30 μl of PBS and injected into the left and right TAs of mdx mice, respectively. After three days, the TAs and a gastrocnemius were harvested and total mononuclear cells were isolated as already described. Cells were then incubated 30 min in ice with the anti-SCA−1 FITC antibody and analysed by flow cytometry. Gastrocnemius was used to determine the gate identifying labelled cells.

EdU administration

5-ethylneyl-2’-deoxyuridine (EdU) (Invitrogen, C10418) was dissolved with 2 ml sterile PBS at the concentration of 5 mg/ml (20 mM). Mice were weight and injected intraperitoneally with 40 mg/kg EdU21. After 24 h, CD45−CD31− were isolated by MACS and staining was performed using Click-i™ EdU Pacific Blue™ Flow Cytometry Assay Kit (Invitrogen, C10418). Briefly, 1 × 10^6 CD45−CD31− were incubated with anti-ITGA7 APC 1:500 (Invitrogen MA5-23555) and anti-SCA-1 FITC 1:50 (BD Pharmingen 557405) antibodies for 30 min in ice. The subsequent fixation, permeabilization and Click-i™ reaction were performed as indicated by the manufacturer’s instructions. Unstained cells and FMO controls were prepared. Samples were resuspended at the concentration of 1 × 10^6 cells per ml in PBS 1% BSA and analysed as described in the flow cytometry section.

Flow cytometry

Cell suspensions were incubated with primary antibodies in ice for 30 min in PBS containing 2 mM EDTA and 2% BSA at the concentration of 1 × 10^6 cells/ml. We used the following antibodies and dilutions: anti-ITGA7 APC 1:500 (Invitrogen MA5-23555), anti-SCA-1 FITC 1:50 (BD Pharmingen 557405) and anti-CD140a APC 1:50 (eBioscience, San Diego, California, United States 17-1401). To stop the incubation cell suspensions were diluted with 2 ml of PBS 2 mM EDTA 2% BSA and centrifuged at 4°C for 10 min at 700 × g. Pellets were resuspended in PBS 2 mM EDTA 2% BSA at the concentration of 1 × 10^6 cells/ml. Unstained samples and suitable FMO controls were prepared when necessary. Approximately 10,000 events per samples were acquired by CytoFLEX S (Beckman Coulter, Brea, California, United States) equipped with three lasers (488 nm, 405 nm and 638 nm) and 13 detectors. Quality control of the cytometer was assessed daily using CytoFLEX Daily QC Fluorospheres (Beckman Coulter B52320). Data were collected by CytExpert (Beckman Coulter) software. If needed, a compensation matrix was calculated using VersaComp Antibody Capture Kit (Beckman Coulter B22804) according to the manufacturer’s instructions. FCS files were analysed using FlowJo™ Software.

Multiplex flow cytometry assay and analysis

Total mononuclear cells were isolated from wild type and mdx mice as described. 1 × 10^6 cells were stained in PBS containing 2 mM EDTA and 2% BSA at the concentration of 1 × 10^6 cells/ml in ice for 30 min with antibodies reported in Supplementary Table 1. 2 ml of PBS 2 mM EDTA 2% BSA were added to stop the incubation, cell suspensions were centrifuged and cell pellets were suspended in PBS containing 2 mM EDTA and 2% BSA. At least 300,000 events were acquired by CytoFLEX S (Beckman Coulter) and collected by CytExpert (Beckman Coulter) software. Wild type and mdx mice samples were acquired at different days therefore we created two different compensation matrices. Next FCS files of these two datasets were simultaneously uploaded on Cytobank platform to perform multidimensional analysis. Cells were identified by SSC-A and FCS-A and single cells by FCS-A and FSC-H applying ‘tailored’ gates for each sample.

Dimensional reduction was carried out applying the tSNE22 algorithm as followed: 50,000 events per sample; 2000 iterations; perplexity = 30; theta = 0.5. All 9 channels were selected applying compensation matrices.

To identify main muscle populations firstly we ran the FlowSOM algorithm with the following settings: all
markers were selected; 50,000 events per sample; 100 clusters; 25 metaclusters; 10 iterations with scale normalisation. A hierarchical consensus clustering method was used to obtain the metaclusters. Next, metaclusters with similar expression profiles were merged together using the ‘automated cluster gates’ function.

FAP heterogeneity was studied applying the FlowSOM algorithm as followed: 600 events per samples; selected markers were specified in the results section; 100 clusters; 4 metaclusters; 10 iterations with scale normalisation. A hierarchical clustering method was used to obtain the metaclusters. We merged the two metaclusters expressing higher levels of SCA-1 and the two metaclusters expressing lower levels of SCA-1 obtaining two groups of FAPs. Self-organising maps (SOM) were uploaded in Cytoscape23 and graph layouts were modified.

Mass cytometry
Isolation of FAPs by MACS technology from mdx mice was performed according to the protocol described in Cerquone Perpetuini et al., 2020.

FAPs were incubated with Cell-ID-Cisplatin-194Pt (Fluidigm, South San Francisco, California, United States, 201194) at the concentration of 1 μM to stain dead cells. Next, samples from three independent biological replicates were barcoded with a combination of palladium isotopes of the Cell-ID 20-Plex Pd Barcoding kit (Fluidigm 201067). Finally, cells were quenched by adding Maxpar cell staining buffer (Fluidigm 201060) at the concentration of 5 μM. Staining was quenched by adding Maxpar cell staining buffer (Fluidigm 201068). Barcoded samples were mixed and stained with lanthanides-conjugated antibodies according to the manufacturer’s instructions. The full list of antibodies purchased from Fluidigm is reported in Supplementary Table 2. Next, the suspension was incubated with Cell-ID Intercalator-Ir (Fluidigm, 2011928) at the concentration of 125 nM for 1 h in Maxpar fix and Perm Buffer (Fluidigm 201067). Finally, cells were filtered with 30 μm cell strainer.

Samples were acquired using CyTOF2 tuned and calibrated according to the manufacturer’s instructions. Rate of acquisition was < 400 events per second. Dataset was converted in fcs files using Debarcoding software (Fluidigm) and normalised.

Analysis of mass cytometry data
Debarcoded fcs files were uploaded to Cytobank platform. Cells were identified using DNA1 and DNA2 content identified by Ir intercalator, while single cells were identified using the event length parameter and DNA1 (or DNA2). Next, dead cells were gated out using Pt channel.

For the experiment with FAPs isolated by MACS and induced to differentiate into adipocyte, dimensional reduction was carried out applying the tSNE algorithm as followed: 35,000 events per sample; 1000 iterations; perplexity = 30; theta = 0.5. All parameters were selected as clustering channels. To identify FAP cell states we ran the FlowSOM algorithm with the following settings: SCA-1 and phospho-CREB-1 were selected as clustering channels; all events were selected; 100 clusters; 3 metaclusters; 10 iterations. A k-Means clustering method was used to obtain the metaclusters. Next, metaclusters with similar expression profiles were merged together using the ‘automated cluster gates’ function and mapped onto viSNE maps to facilitate results interpretation.

Cell culture
Freshly isolated mdx FAPs cell states were seeded on 96 well plates (Falcon 353072) in growth medium (GM) at the density of 15,000 cell/cm² for differentiation assays and at the density of 6200 cells/cm² for the proliferation assay. GM is composed by Dulbecco’s Modified Eagle Medium, High Glucose, GlutaMAX™ (Gibco 61965-026) supplemented with 20% fetal bovine serum (FBS) (Euroclone, Pero, Milan, Italy, ECS0180L), 100 U/ml penicillin and 100 μg/ml streptomycin (Gibco 15140122), 1 mM sodium pyruvate (Sigma-Aldrich, St. Louis, Missouri, United States, S8636) and 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) (HEPES) (Sigma H8087).

To induce adipogenic differentiation cells were exposed to the adipogenic induction medium (AIM) consisting of GM complemented with 1 μg/ml insulin (Sigma-Aldrich I9278), 1 μM dexamethasone (Sigma D4902) and 0.5 mM of 3-isobutyl-1-methylxanthine (IBMX) (Sigma, I5879). After three days, cells were exposed to the maintenance medium (MM) composed by GM and 1 μg/ml insulin for additional two days. For the adiponectin experiment, the two FAP cell states were treated with 25 μM AdipoRon (Selleckchem, Munich, Germany S7365) for 24 h in GM before adipogenic induction.

To induce fibrogenic differentiation cells were cultured in a medium composed by Dulbecco’s Modified Eagle Medium, High Glucose, GlutaMAX™, 100 U/ml penicillin and 100 μg/ml streptomycin, 1 mM sodium pyruvate and 10 mM HEPES, 5% horse serum (HS) (Euroclone ECS0090D) and 1 ng/ml TGF-beta-1 (PreproTech, Cranbury, New Jersey, United States 100-21). To study the expression of collagen genes, cells were plated in 24 well plates at the concentration of 15,000 cells/cm².

For intravital expansion, FAPs cell states were cultured at the concentration of 3400 cells/cm² for 9 days in Cytogrow (Resnova, Genzano di Roma, Italy TGM-9001-B) medium. Afterwards, cell monolayers were washed twice in sterile D-PBS and cells were detached with Trypsin 0.5 g/l EDTA 0.2 g/l (Lonza, Basel, Switzerland # BE17-161E) for 5 min. Next, cells were processed for flow cytometry analysis.

Muscle satellite cells (MuSCs) were cultured in a growth medium composed by Dulbecco’s Modified Eagle
Medium, High Glucose, GlutaMAX™, 100 U/ml penicillin and 100 mg/ml streptomycin, 1 mM sodium pyruvate, 10 mM HEPES, 20% FBS, 10% Horse serum (Euroclone ECS090D), 2% Chicken embryo extract (Biotrend, Koln, Germany CE-650-J).

CD45⁺ leucocytes were isolated from mdx mice with MACS technology. They were cultured for 24 h in DMEM Dulbecco's Modified Eagle Medium, High Glucose, GlutaMAX™, 100 U/ml penicillin and 100 mg/ml streptomycin with 0.2% BSA at 375,000 cells/ml. Next, media were harvested, centrifuged at 700 × g for 10 min at 4°C and finally supernatant was filtered to remove cell debris. Conditioned media from different biological replicates were mixed just before the treatment. One volume of the resulting conditioned medium was mixed with one volume of 2X AIM to induce adipogenesis. The same procedure was applied to prepare MM.

**Immunofluorescence**

Indirect immunolabelling was performed as followed. Cells were fixed with 2% paraformaldehyde (Santa Cruz sc-281692) for 20 min at room temperature (RT). Paraformaldehyde was removed by inversion and cells were washed three times in PBS and one time in PBS 0.5% Triton X-100. Permeabilization was performed incubating cells for 5 min with 0.5% Triton X-100 (Sigma T9284) in PBS. Permeabilization solution was removed by inversion and cells were washed with 0.1% Triton X-100 in PBS. Unspecific sticky sites were blocked with a blocking solution consisting of 10% FBS 0.1% Triton X-100 in PBS for 30 min at RT. Next, samples were incubated with the following primary antibodies for 1 h at RT: rabbit anti-PPAR-gamma 1:200 (Cell Signaling, Danvers, Massachusetts, United States 2443S), mouse anti-SMA 1:300 (Southern Biotech, Birmingham, Alabama, United States AF1062) diluted in PBS 2.5% BSA: goat anti-CD140a 1:80 (R&D Systems, Minneapolis, Minnesota, United States AF1062) and rabbit anti-PPAR-gamma 1:200 (Cell Signaling 2443S). The following day section was incubated overnight at 4°C with the anti-CD140a (R&D Systems, Minneapolis, United States AF1062) diluted 1:100 in blocking solution and incubated with sections overnight at 4°C. Sections were washed twice in 0.2% Triton X-100 1% BSA in PBS (washing solution) for 15 and 5 min at RT. Incubation with goat anti-rabbit Alexa Fluor 488 1:200 (Southern Biotech 4050-30) diluted in blocking solution was performed at RT for 30 min. At the end of incubation, sections were washed 15 min in washing solution and 5 min in PBS. Finally, nuclei were stained with 2 μg/μl Hoechst 33342 (Invitrogen H3570) in PBS for 10 min at RT. Sections were washed twice in PBS for 5 min. Finally, slides were covered with coverslips using Aqua-Poly/ Mount (Tebu-bio 18606-20) and stored at 4°C.

For the on-section detection of FAPs, TAs section were fixed in 2% PFA and washed three times with PBS. Permeabilization was performed incubating sections with 0.3% Triton X-100 in PBS for 30 min. This step was directly followed by an incubation with Protein Block Serum Free (Dako X0909) for 30 min to block the sticky sites. After a wash in PBS for 5 min, sections were incubated overnight at 4°C with the following antibodies diluted in PBS 2.5% BSA: goat anti-CD140a 1:80 (R&D Systems AF1062) and rabbit anti-PPAR-gamma 1:200 (Cell Signaling 2443S). The following day section was washed three times for 10 min with PBS 0.1% Triton X-100. Next, samples were incubated for 30 min with the following secondary antibodies diluted in PBS 2.5% BSA: donkey anti-goat Alexa Fluor 488 1:200 (Southern Biotech 3425-30) and donkey anti-rabbit Alexa Fluor 555 1:200 (Thermo fisher scientific A32794). After two wash for 5 min each with PBS 0.1% Triton X-100 and three wash in PBS, sections were incubated with 2 μg/μl Hoechst 33342 (Invitrogen H3570) in PBS for 10 min at RT. Sections were washed three times in PBS for 5 min. Finally, slides were covered with coverslips using Aqua-Poly/Mount (Tebu-bio, Le Perray-en-Yvelines, France 18606-20) and stored at 4°C.

**Oil red O staining**

Oil red O (ORO) (Sigma-Aldrich O0625) was dissolved in isopropanol at the concentration of 3.5 mg/ml. This
stock was diluted to a working solution composed of 3 volumes of ORO and 2 volumes of distilled water. This ORO solution was filtered two times just before the use. Fixed cells were washed three times in PBS for 5 min and were incubated with ORO for 10 min at RT. ORO was removed by inversion and cells were washed three times in PBS for 5 min. Nuclei were counterstained with 2 μg/μl Hoechst 33342 in PBS 0.1% Triton X-100 for 5 min at RT. Finally, cells were washed three times in PBS for 5 min and were stored at 4 °C in PBS containing 0.02% of sodium azide.

During immunofluorescence, ORO staining was performed after incubation with secondary antibodies. After three washes in PBS protocol continues with the Hoechst 33342 incubation as described in the immunofluorescence section.

Alkaline phosphatase assay
The two FAP cell states were isolated from mdx mice and cultured for 5 days in GM at 15,000 cell/cm². Samples were fixed using 2% paraformaldehyde and then washed three times with PBS. NBT/BCIP stock solution (Roche 11681451001) was diluted 1:50 in a buffer composed of 0.1 M Tris, 0.05 M MgCl₂, 0.1 M NaCl (pH = 9.5) and added to cell monolayers. After 10 min this buffer was removed and samples were washed three times in PBS. Finally, nuclei were stained with Hoechst 33342 as already described.

CFSE staining and analysis of cell morphology
CarboxyFluorescein succinimidyl ester (CFSE) (Abcam, Cambridge, United Kingdom ab113853) staining was carried out according to the manufacturer’s instructions. Culture medium was removed and then 10 μM of CFSE in PBS was overlaid onto cells. After 15 min of incubation at 37 °C, staining was quenched adding an equal volume of GM and allow to sit for 5 min at 37 °C. Next medium was removed and cells were washed one time in PBS and fixed with 2% paraformaldehyde. Samples were washed three times in PBS for 5 min. Nuclei were counterstained with 2 μg/μl Hoechst 33342 in PBS 0.1% Triton X-100 for 5 min at RT. After three wash in PBS samples were at 4 °C in PBS containing 0.02% of sodium azide.

Images were manually acquired. Images were analysed using Fiji. Background signal was removed using the subtract background command and an image specific threshold was applied to obtain a binary image. Then we run the following series of commands to define cell borders: median filter, erode and fill holes. Finally, we measured area occupied, roundness and aspect ratio. We analysed at least 50 cells from a total of three independent biological replicates.

Acquisition of images and analysis
Images were acquired with a LEICA microscope (DMi6000B) using the LAS X software. For experiments using ex vivo cell culture, a matrix of 25 non-overlapping field per well was acquired. Experiments were performed in technical duplicate. ORO positive cells, Ki-67 positive nuclei and PPAR-gamma positive nuclei were manually counted using Fiji and were divided by the number of nuclei per field. SMA-positive area per cell was assessed using a dedicated pipeline of CellProfiler10. Area expressed in pixel was converted in square micron and normalised over the total number of nuclei in each field.

Microphotographs of TA sections immunolabelled for perilipin were acquired using a 10x objective. At least three sections for each mouse were acquired. These overlapped images were used to reconstruct the whole TA section through the grid/collection stitching algorithm available in Fiji. To identify perilipin positive area we applied a threshold (0-100) and we divided this area by the whole section area.

Microphotographs of TA section labelled for CD140a were acquired with Leica microscope for the quantification and representative images were acquired using the confocal microscope Olympus IX-81 with a 40x objective. Images of CD140a and PPAR-gamma positive FAPs on the TA section were acquired using the confocal microscope Olympus IX-81 with a 40x objective.

RNA extraction, retrotranscription and Real-Time PCR
Cell monolayers were washed in PBS and lysed in TRIzol™ Reagent (Invitrogen 15596-018). For extraction from tissue, TAs were homogenised in liquid nitrogen using a pestle and then resuspended in TRIzol™. Isolation of total RNA was performed according to the manufacturer’s instruction. RNA precipitation was performed at −20 °C overnight with glycogen. Total RNA was resuspended in nuclease free water, concentration and 260 nm/280 nm ratio were determined by Nanodrop Lite Spectrophotometer (Thermo Fisher Scientific, Waltham, Massachusetts, United States). Samples were diluted to the concentration of 50 ng/μl and stored at −80 °C. cDNA was generated by PrimeScript RT Reagent Kit (Takara, Kusatsu, Japan RR037A). Real-time PCR was performed on 15 ng of cDNA using SYBR Premix Ex Taq (Tli RNaseH Plus) (Takara RR420W) in a reaction volume of 20 μl. Tubulin was used as housekeeping gene. We used primers in Supplementary Table 3. Comparative expression was performed using the 2−ΔΔCt method27 where ΔΔCt is (Ct, target − Ct, tubulin) − (Ct, mean target − Ct, mean tubulin).

RNA-sequencing experiment
FAP cell states were cultured 5 days in GM at the density of 15,000 cell/cm². Next, media were removed, cell monolayers were washed twice in PBS and detached using Trypsin 0.5 g/l EDTA 0.2 g/l. Total RNA was isolated from the harvested cells using RNeasy micro kit (Qiagen, Hilden, Germany 74004) and was quantified
using the Qubit 2.0 fluorimetric Assay (Thermo Fisher Scientific). Libraries were prepared from 100 ng of total RNA using the QuantSeq 3′ mRNA-Seq Library Prep Kit FWD for Illumina (Lexogen GmbH, Vienna, Austria) and their qualities were assessed by using screen tape High sensitivity DNA D1000 (Agilent Technologies, Santa Clara, California, United States). Libraries were sequenced on a NovaSeq 6000 sequencing system using an S1, 100 cycles flow cell (Illumina Inc., San Diego, California, United States). Illumina novaSeq base call (BCL) files were converted into fastq file by bcl2fastq (version v2.20.0.422). Sequence reads were trimmed using bbduk software (bbmap suite 37.31) in order to remove adapter sequences, poly-A tails and low-quality end bases (regions with average quality below 6). Alignment was performed with STAR 2.6.0a28 on mm10 reference assembly obtained from cellRanger website (Ensembl assembly release 93). The expression levels of genes were determined with htseq-count 0.9.15 (Ref. 29) by using mm10 Ensembl assembly (release 93) downloaded from cellRanger website. The expression data was analysed by Rosalind, with a HyperScale architecture developed by OnRamp BioInformatics. Individual sample counts were normalised via Relative Log Expression (RLE) using DESeq2 R library30. Multidimensional scaling (MDS) plot was generated as part of the QC step. DESeq2 was also used to calculate fold changes and p-values and perform covariate correction. Clustering of genes for the final heatmap of differentially expressed genes was done using the PAM (Partitioning Around Medoids) method using the fpc R library. Gene set enrichment analysis was performed using DAVID31. PPAR-gamma transcriptionally regulated genes have been retrieved from TRRUST32. Cystoscape has been used to create the networks.

The RNA-seq data have been deposited in GEO with the identifier GSE163016.

Single-cell RNAseq data analysis

Briefly, single-cell RNA-seq data of hind limb muscle have been retrieved from Tabula Muris Senis project and re-processed with the Seurat R package33. After log normalisation, we used the 2000 most variable features to proceed to downstream analyses. Data have been scaled and the first 10 principal components used to run the t-distributed stochastic neighbour embedding (tSNE) algorithm. Whenever possible, biomarkers have been retrieved from Myo-REG database to assign cell line ontology identifications to each cluster, on the basis of the expression level of each biomarker for each cluster, according to their distribution. Once cell IDs have been assigned, FAPs population subset has been further analysed for the expression of SCA-1 and downstream analyses.

Differentially expressed genes were identified by comparing the mRNA expression profile of the two FAPs clusters. Only genes whose expression difference had an adjusted p-value < 0.05 were considered as sub-population biomarkers. EnrichR34 has been used to retrieve the enriched GO Molecular Function terms for each population with p-value < 0.05.

Statistical analysis

Experiments were performed with primary cells isolated from at least three different mice (here defined biological replicates). The number of biological replicates was specified in the figure legends. No statistical methods were applied to estimate sample size and the investigator was not blinded to the group of allocation during the experiments. Statistical analysis was performed by a two-tailed Student’s t test between two groups, while one-way ANOVA or two-way ANOVA were applied for comparison among more than three groups. Results were presented as mean ± standard error of the mean (SEM). Differences were considered significant when p-value is less than 0.05 (*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001). Plots and statistical analysis were produced using the GraphPad Prism 7 software.

Results

SCA-1 expression defines two FAP populations

We have recently shown that purified FAPs are a heterogeneous population including several sub-populations19. Here we use a multiplex flow cytometry approach to characterise FAP heterogeneity from a suspension of mixed populations derived from skeletal muscles. We first isolated mononuclear cells from hind limb muscles of 6-8 weeks old wild type and mdx mice. In this time window, muscles of dystrophic mice are in a regeneration phase35,36. Cells were stained with fluorescent antibodies against 9 surface antigens (CD45, CD31, Integrin alpha-7, SCA-1, CD140a, CD146, CD54, CD90.2) to distinguish the main mononuclear cell populations residing in the skeletal muscle: leucocytes (CD45+), endothelial cells (CD45-CD31+), muscle satellite cells (MuSCs) (CD45-CD31-Integrin alpha-7-), and FAPs (CD45-CD31-Integrin alpha-7-SCA-1+). The antigen expression profiles of single cells were analysed by a mixed approach whereby an automatic clustering is followed by expert manual refinement (Fig. 1a and details in supplementary materials). This approach yielded 4 cell clusters corresponding to the main muscle mononuclear cell populations (Fig. 1b) and 11 unassigned clusters (Supplementary Fig. 1d), which were characterised by an antigen profile that could not be assigned to any of the main mononuclear cell populations in the muscle niche. FAPs amount to 2.6% of the mononuclear cells in the wild type mouse, whereas their abundance increases to 5.7% in the mdx mouse (Supplementary Fig. 1e).
We next focused on FAPs and we chose to study the expression variability of 6 antigens (SCA-1, CD140a, CD140b, CD146, CD34, CD90.2) associated to the FAP immunophenotype. By applying the same clustering strategy used for the total mononuclear populations, we obtained 4 metaclusters represented by red, blue, green and black outlined nodes of the self-organising maps (SOMs) produced by the FlowSOM algorithm.
dynamic cell states. Terminology in the cell plasticity of SCA-1 represents stable sub-populations or rather a cell state rather than a cell type model (Fig. 1d). The intensity distribution of SCA-1 shows a single peak and a large standard deviation (Fig. 1c), and is a feature that characterises the cell to cell variability dubbed micro-heterogeneity.

SCA1-High-FAPs and SCA1-Low-FAPs are cell states in dynamic equilibrium

The observation that FAPs display micro-heterogeneity prompted us to design a sorting strategy to isolate, from the mdx mouse model, FAPs (CD45<sup>-</sup> CD31<sup>-</sup> Integrin alpha-7<sup>-</sup> SCA-1<sup>+</sup>) expressing high and low levels of SCA-1, henceforth called SCA1-High-FAPs and SCA1-Low-FAPs respectively (Fig. 1f; refer to Supplementary Fig. 3a for the complete gating strategy, to the Supplementary File). Similar sorting strategies have already been used to resolve micro-heterogeneity. Sorted FAP populations are significantly enriched for CD140a, as assessed by immunofluorescence (Supplementary Fig. 3d) and by flow cytometry (Supplementary Fig. 3e, f). Both populations can be stained for alkaline phosphatase (Supplementary Fig. 3g) but they do not show any myogenic potential.

Next, we asked whether FAPs expressing high and low levels of SCA-1 represent stable sub-populations or rather dynamic cell states. Terminology in the cell plasticity field is still debated. Here we define a “sub-population” of a given cell type as a group of cells identified by a discrete expression of specific markers and capable of maintaining identity when grown for several duplications ex vivo. Instead, considered a seemingly homogeneous cell population, we talk of a “cell state” when referring to a group of cells with an expression profile skewed when compared to the symmetrical expression distribution of the parent population. Cell states are in a dynamic equilibrium therefore they lose their identity after in vitro expansion.

To address this issue, we performed a repopulation experiment in which SCA1-High-FAPs and SCA1-Low-FAPs were allowed to proliferate ex vivo. After 9 days, we analysed SCA-1 expression by flow cytometry. The SCA-1 intensity distributions of freshly isolated cells are clearly separated, whereas in expanded cells they overlap and restore the expression profile of the whole FAP compartment (Fig. 2a, gating strategies and controls in Supplementary Fig. 4a–e). We also explored the dynamics of the two populations in vivo by injecting purified and fluorescently labelled FAP populations into the tibialis anterior (TAs) of mdx mice. Three days after transplantation, total mononuclear cells were isolated from TAs and the distribution of SCA-1 expression was evaluated by flow cytometry (Fig. 2b). Even though a repopulation process is observed in the muscle niche, its dynamic is different from that characterised in vitro (Fig. 2c). These observations suggested that SCA1-High and SCA1-Low-FAPs are dynamic states capable of repopulating each other with different dynamics in vitro and in vivo.

We next investigated whether SCA1-High-FAPs and SCA1-Low-FAPs could be distinguished in vitro according to their morphology. To this end, we stained their cytoplasms with Carboxyfluorescein succinimidyl ester (CFSE) and we selected three suitable descriptors to characterise cell shape. Roundness and aspect ratio are parameters that describe round cells and elongated cells respectively (Supplementary Fig. 4f). The third descriptor is cell size, which was estimated by the CFSE positive area. As a control we included in this analysis a population enriched for MuSCs since they acquire characteristic shapes during myogenic differentiation. Indeed,
Fig. 2 (See legend on next page.)
in the MuSC sample we are able to identify cells that exhibit high roundness/low AR and cells with low roundness/high AR (Supplementary Fig. 4g–j). The former round cells are proliferating myoblasts, the latter elongated cells are differentiating myocytes39. Moreover, these myogenic cells are smaller than SCA1-High-FAPs or SCA1-Low-FAPs (Supplementary Fig. 4i). Focusing on SCA1-High-FAPs (Fig. 3b, d) we stained the lipid droplets using Oil Red O (ORO). We noticed little differentiation in both cell states when incubated in GM in the absence of adipogenic stimuli. Interestingly in cells incubated with the adipogenic medium (AM), the percentage of adipocytes in SCA1-High-FAPs is significantly higher than the one observed in SCA1-Low-FAPs (Fig. 3b, d). This different behaviour may be the result either of a different propensity to undergo adipocyte commitment or to a delay in adipogenic differentiation. We addressed this point by evaluating the expression of peroxisome proliferator-activated receptor gamma (PPAR-gamma) by immunofluorescence. This transcription factor is a master regulator of adipogenesis and is not expressed in FAPs soon after isolation14,15. PPAR-gamma expression is significantly higher in SCA1-High-FAPs (Fig. 3c, d) suggesting that adipocyte commitment may be impaired in SCA1-Low-FAPs. To further support this conclusion, we allowed a longer time for FAPs to differentiate into mature adipocytes by extending the incubation in MM to 11 days (Fig. 3e). This long-term insulin stimulation allows cells to display their full adipogenic potential. In this condition, both cell states differentiate more efficiently and adipocytes are more mature compared to what observed in the standard protocol (Fig. 3g). However, also in this condition, a larger percentage of SCA1-High-FAPs yielded mature adipocytes (Fig. 3f). At the end of the differentiation process, SCA1-High-FAPs reach a higher confluence when compared to SCA1-Low-FAPs (Supplementary Fig. 5a-b). To rule out the possibility that the observed differences in the adipogenic differentiation of FAP cell states could be a consequence of a different kinetics in reaching cell confluence, we plotted the total number of nuclei per field over the total number of adipocytes per field from the...
Fig. 3 (See legend on next page.)
FAPs were isolated from muscles of CREB-1, a marker of early adipogenic differentiation, and CXCR-4, SCA-1, CD31, c-Kit, Actin, phospho-CREB-1). Vimentin, CD90.2, phospho-STAT3, Integrin alpha-7, of antibodies against 18 proteins (CD45, CD146, cleaved single cells were analysed by mass cytometry using a panel and then were incubated in either GM or AIM. After 72 h path more readily.

Low-FAPs are characterised by a reduced adipogenesis, correlate with a different adipogenic potential. SCA1-SCA-1 expression levels in the different FAP cell states and SCA1-High-FAPs (Supplementary Fig. 5d).

addition, we found that SCA-1 expression decreases clearly reveals a correlation between cell con...

Collectively, these results indicate that the different SCA-1 expression levels in the different FAP cell states correlate with a different adipogenic potential. SCA1-Low-FAPs are characterised by a reduced adipogenesis, whereas SCA1-High-FAPs engage in this differentiation path more readily.

FAP cell states differ in the expression of fibrogenic genes

Abnormal deposition of extracellular matrix (ECM) is another pathogenic hallmark of muscular dystrophies. Differently from fat deposition this condition also develops in young dystrophic mice. Myofibroblasts differentiate from several cell types and, among these, FAPs are considered the main source of fibrotic scars. Upon differentiation, myofibroblasts synthesise alpha-smooth muscle actin (alpha-SMA) to form positive stress-fibres and increase the production of ECM proteins. Transforming growth factor beta-1 (TGF-beta-1) promotes fibrogenic differentiation of FAPs in vitro and takes part in fibrosis in vivo.

We asked whether FAP heterogeneity, in addition to modulating adipogenesis, could also impact on fibrogenesis. After 5 days in growth medium (GM), FAPs were exposed for 5 additional days to a fibrogenic medium (FM) containing TGF-beta-1 (Fig. 4a). Next, the expression of alpha-SMA was monitored by immunofluorescence. Both cell states respond to FM by increasing the alpha-SMA-positive area. The alpha-SMA-positive area surrounding SCA1-Low-FAP cells was twice as large as that of SCA1-High-FAP cells (Fig. 4b, c, nuclei count in Supplementary Fig. 5e). The SMA-positive area in samples exposed only to GM also tends to be higher in SCA1-Low-FAPs.

We also monitored the mRNA levels of Col3a1 and Col1a1 genes upon fibrogenic induction. These collagen chains are components of the skeletal muscle ECM and are produced by FAPs upon TGF-beta-1 treatment in vitro. Both cell states respond to FM by increasing the expression of Col3a1 mRNA (Fig. 4d). However, only SCA1-High-FAPs significantly increased the expression of Col1a1 mRNA upon induction of differentiation (Fig. 4e).

By looking at the mRNA levels in the absence of induction of fibrogenesis (GM), SCA1-Low-FAPs tend to express higher level of Col1a1. This difference was significant in freshly isolated cells (Fig. 4h). We found no difference in Col3a1 mRNA levels when the two cell states were compared in freshly isolated cells or after growth in GM (Fig. 4g). Moreover, we monitored the expression of the Timp1 mRNA, another transcript expressed by FAPs upon in vitro fibrogenic stimulation. SCA1-High-FAPs but not SCA1-Low-FAPs respond to the FM enhancing their expression of the Timp1 mRNA (Fig. 4f).
Fig. 4 (See legend on next page.)
We conclude that both FAPs cell states are able to differentiate into myofibroblasts when exposed to a fibrogenic microenvironment, but they exhibit a differential expression of at least two genes coding an ECM component.

SCA1-High-FAPs are characterised by a transcriptional profile responsive to adipogenic stimuli

We next set out to compare the transcriptional landscapes of the two FAP cell states as it may shed light on the molecular determinants of their different responsiveness to adipogenic and fibrogenic stimuli. Therefore, we studied their transcriptional profiles by RNA-sequencing after 5 days of incubation in GM. The multidimensional scaling (MDS) plot of the transcriptional data shows that the two cell states at this time, before differentiation induction, have distinct transcriptomes (Fig. 5a), a difference which is most noticeable along coordinate 1. The differentially expressed genes are 704 (adjusted p-value < 0.05) 266 of which are up regulated (log2 fold change > +0.5) and 438 are downregulated (log2 fold change < −0.5) in SCA1-Low-FAPs (Fig. 5b). Samples are grouped into 2 different clusters that discriminate SCA1-High-FAPs (the orange cluster) and SCA1-Low-FAPs (the blue cluster) (Fig. 5c). Moreover, differentially expressed genes are grouped into 2 clusters of upregulated and downregulated genes (Fig. 5c, d). A gene set enrichment analysis of the “Biological Process” GO terms revealed that in SCA1-High-FAPs the genes involved in adipocyte differentiation (GO terms “brown fat differentiation” and “fat cell differentiation”), among others, are over-represented in the list of upregulated genes (Supplementary Table 4) (Fig. 5d). This is consistent with the observation that SCA1-High-FAPs are more prone to differentiate into adipocytes. In agreement with this hypothesis, some of the genes that are transcriptionally regulated by PPAR-gamma are upregulated in SCA1-High-FAPs (Fig. 5e) and PPAR-gamma is more expressed in SCA1-High-FAPs (Fig. 5f, g).

The second most upregulated mRNA in SCA1-High-FAPs is Adipoq which encodes for adiponectin (log2 fold change = −4.07). We confirmed this differential expression by Real-Time PCR (Supplementary Fig. 6a). Adiponectin is a well-known sensitiser of the insulin pathway47,48. For this reason, we hypothesise that the spontaneous adipogenic propensity of SCA1-High-FAPs is likely due to an adiponectin-mediated autocrine circuit, which is not active in SCA1-Low-FAPs (Supplementary Fig. 6b). According to this hypothesis, adiponectin sensitises SCA1-High-FAPs to the adipogenic mix which contains insulin. Indeed, a 24 h pre-treatment of SCA1-Low-FAPs with AdipoRon, a small molecular which binds and activates the adiponectin receptors49, induces a three-fold increase of their sensibility to the adipogenic mix (Supplementary Fig. 6c, d).

Regarding fibrogenic differentiation, we found that several collagen genes are differentially regulated. Among these, the Col3a1 mRNA is upregulated in SCA1-High FAPs (log log2 fold change = −1.03), while Col12a1 mRNAs are upregulated in SCA1-Low-FAPs (log2 fold change = 0.82). Consistent with the observation that SCA-1 activates metalloproteinases (MMPs) expression50, three MMPs genes are upregulated in SCA1-High-FAPs: Mmp11 (log2 fold change = −0.84), Mmp23 (log2 fold change = −0.91) and Mmp19 (log2 fold change = −0.81) (Supplementary Fig. 6e, g). Finally, SCA1-High-FAPs express higher levels of Alpl mRNA than SCA1-Low-FAPs (Supplementary Fig. 6h).

In conclusion, the two FAP cell states differentially express several genes that participate in ECM assembly and remodelling. At the same time, SCA1-High-FAPs spontaneously acquire a transcriptional profile that sensitise them to induction of adipogenic differentiation, possibly under an autocrine signalling circuit mediated by adiponectin.

SCA1-High-FAPs proliferate more extensively than SCA1-Low-FAPs

A coordinated modulation of FAP proliferation and apoptosis is a key requisite to achieve efficient muscle regeneration17. Indeed, the deregulation of these processes observed in DMD leads to FAP accumulation and the ensuing deposition of ectopic tissues23. We investigated whether the observed FAP heterogeneity would also affect cell proliferation. Cells were sorted from mdx mice and the two cell states were plated at the same low density. Starting at the first time point (T0), we collected samples every 24 h (Fig. 6a) and we...
counted the number of nuclei and the percentage of Ki-67 expressing cells to identify proliferating cells. Already at 48 h, the number of SCA1-High-FAPs nuclei in a field is significantly higher than in the SCA1-Low-FAPs (Fig. 6b). However, the percentage of cells expressing Ki-67 is comparable in freshly purified cells and decreases more
Fig. 6 (See legend on next page.)
rapidly at later times in SCA1-High-FAPs (Fig. 6c) as this cell state reaches confluence earlier and stops growing. In the exponential growth phase, the doubling time of SCA1-High-FAPs is about 10 h shorter (Fig. 6d and Supplementary Fig. 7a). Taken together these results suggest that ex vivo SCA1-High-FAPs have a shorter duplication time reaching at late time points a higher confluence and a lower number of Ki-67 positive cells.

In order to lend support to an in vivo relevance of these observations, we injected 5-ethyl-2'-deoxyuridine (EdU) into the peritoneum of mdx mice. After 24 h we isolated mononuclear cells (Fig. 6e) and we selected CD45−CD31− cells by magnetic activated cell sorting (MACS). Next, we identified FAP cell states in the Integrin alpha-7−SCA-1+ compartment and monitored EdU incorporation using flow cytometry (Fig. 6g; for the unstained sample and FMO controls see Supplementary Fig. 7b, c). A larger fraction of SCA1-High-FAPs have incorporated EdU in proliferating FAPs (n = 5). g Gating strategy to evaluate EdU incorporation. For the unstained sample and FMO controls see Supplementary Fig. 7b, c. Data are presented as mean ± SEM. Statistical analysis was performed using Two-way ANOVA for the in vitro experiment and Student’s t test for the in vivo experiment. * p ≤ 0.05, ** p ≤ 0.01, *** p ≤ 0.0001.

Single-cell transcriptomic analysis of FAP cell states from wild type mice

Till now we have used the expression of a single antigen (SCA-1) to characterise two FAP cell states displaying distinct differentiation and proliferation characteristics. In these experiments, cells were purified from muscles of mdx mice as in this model the two cell states are more evenly represented. However, heterogeneity in the expression of SCA-1 is observed both in wild type and mdx muscles (Fig. 1). To obtain a more general picture of the observed heterogeneity by an independent experimental approach we made use of the Tabula Muris Senis dataset. The Tabula Muris Senis is a comprehensive compendium of single-cell RNA-sequencing datasets in 23 organs from wild type mice at different ages. We focused on the skeletal muscle and performed an unbiased automated clustering using markers from the Myo-REG resource (Supplementary Fig. 8a) to identify the main skeletal muscle populations. We obtained 11 clusters (Fig. 7a) with an expression profile characteristic of each population (Supplementary Fig. 8b–l). Next, we carried out an additional clustering analysis on the FAP population and observed four clusters (Fig. 7b) characterised by different levels of SCA-1 expression (Fig. 7c). Cluster 0 (red), cluster 2 (blue) and cluster 3 (purple) express similar low level of SCA-1 and where collapsed into a single group, thus obtaining two populations expressing high and low SCA-1 levels (Fig. 7d). SCA1-Low-FAPs are more numerous than SCA1-High-FAPs (Fig. 7e), similarly to what was observed in young mice using the flow cytometry assay (Fig. 1c). Moreover, their ratio only slightly increases in ageing.

We performed a Gene Ontology (GO) term enrichment analysis on differentially expressed genes which did not provide any robust clue regarding FAP cell states properties. However, the list of differentially expressed genes revealed that SCA1-High-FAPs express higher levels of Dpp4 mRNA which is a marker of mesenchymal progenitors residing in the reticular interstitium of inguinal white adipose tissue (IWAT). In this tissue, Dpp4 characterise progenitors able to differentiate into adipocytes suggesting an analogy between such progenitors and SCA1-High-FAPs. Next, we asked if the differential expression of the Col1a1 gene observed in mdx FAP cell states (Fig. 4) is conserved in the wild type counterpart. Col1a1 mRNA is slightly downregulated in SCA1-High-FAPs (average log2 fold change = −0.341, adjusted p-value < 0.05), in accord with our findings in FAP cell states freshly isolated from mdx mice.

We conclude that single-cell transcriptional profiling of wild type FAPs allow us to distinguish two cell states on the basis of SCA-1 transcriptional levels. The analysis of genes that are differentially expressed in the two cell states provides hints about possible mechanisms underlying the observed phenotypic differences.

Signals secreted from cells of the immune compartment control adipogenesis of FAP cell states in an age-dependent fashion

In vivo the FAP proliferation and differentiation traits are controlled by the muscle microenvironment. The prolonged and sustained chronic inflammation of mdx...
muscles is characterised by a subset of unconventional macrophages. It was shown that in an in vivo model recapitulating muscle aging in \textit{mdx} mice, these macrophages fail to induce FAP apoptosis and promote their excessive growth\cite{21}. Others report that, in wild type mice, ageing determines a decrease in the number of FAPs associated with a reduced adipogenic potential\cite{53}.

To find out how ageing affects FAPs and FAP cell states in the \textit{mdx} mouse we determined the number of FAPs in the TA of young and aged dystrophic mice by immunofluorescence (Fig. 8a, b). We observed a reduction in the number of FAPs within old dystrophic muscles and this condition is followed by an increase of FAPs expressing PPAR-gamma in the tibialis anterior (TA) of aged \textit{mdx} mice (Fig. 8c). These results suggest that FAPs in the microenvironment of aged \textit{mdx} mice are more committed to their gene expression profile. The four clusters (0, 1, 2,3) were mapped onto the \textit{viSNE} map of FAPs. The expression of SCA-1 antigen has been mapped to each cell of the dataset. d \textit{viSNE} map of FAPs in which clusters 0, 2, 3 are collapsed together to obtain two clusters expressing high level (SCA-1 High) and low level (SCA-1 Low) of SCA-1. e Stacked bar plot showing the percentage of the two population on the basis of SCA-1 expression levels across different ages.

To verify this hypothesis we set out to investigate the effect of the immune cell compartment of young and old \textit{mdx} mice on the adipogenic differentiation of FAP cell states. For this purpose, we isolated CD45$^+$ cells (henceforth called leucocytes) from young (6–8 weeks) and old (60 weeks) \textit{mdx} mice and after 24 h of in vitro culture we harvested their supernatants. Next, we added each of these conditioned media (CM) to AIM and MM to test their effect on young FAP cell states differentiation (Fig. 8d). Adipocytes were identified by ORO staining. The CM of leucocytes from young \textit{mdx} mice significantly inhibits adipogenesis of SCA1-High-FAPs (~7-fold decrease) (Fig.8e, f). We observed a less important inhibition using the medium conditioned by leucocytes from old \textit{mdx} mice (~3-fold decrease). In general, the differentiation potential of FAPs in both cell states is inhibited by the supernatants conditioned by leucocytes from young \textit{mdx} mice and less so by supernatants of leucocytes from old \textit{mdx} mice. While in these conditions the differentiation of SCA1-Low-FAPs is cut down to negligible levels, the differentiation of SCA1-High-FAPs still remains significant especially when exposed to conditioned media of leucocytes from old \textit{mdx} mice. These observations suggest that the microenvironment of young \textit{mdx} mice is more efficient in inhibiting FAP adipogenesis and probably sufficient to keep in check the differentiation of both FAP cell states. With aging, the negative control exerted by the CD45$^+$ cell compartment
Fig. 8 (See legend on next page.)
becomes less effective, selectively releasing the adipogenic potential of SCA1-High-FAPs. These findings offer a possible mechanism to explain the appearance of IMAT in the TA of aged mdx mice as demonstrated by the increase of immunostaining against Perilipin (Fig. 8g, h) and the increase of Cebpb mRNA in the whole TA (Fig. 8i).

**Discussion**

Non-genetic heterogeneity in clonal cell populations is suggested to underlie fundamental developmental and pathological processes. However, as a consequence of the dearth of appropriate technologies, this concept has not been widely addressed as many experimental approaches still rely on bulk analysis of cell populations. The emergence, over the past decade, of powerful single-cell approaches has permitted to uncover a common hidden heterogeneity in biological systems. To address the relation between heterogeneity and disease, we applied a multiplex flow cytometry approach to study the expression of nine surface antigens in a suspension of primary cells from a solid tissue. The single-cell expression profiles were used in a clustering approach to identify fibro/adipogenic progenitors (FAPs) in wild type and dystrophic mice. We confirmed the inherent heterogeneity of this cell population (Supplementary Fig. 2) and by focusing on the expression of the SCA-1 antigen we identified two cell clusters that are present with different abundance in the muscles of wild type and mdx mouse models. Interestingly an unsupervised clustering of the single-cell gene expression dataset of the Tabula Muris Consortium also leads to similar conclusions identifying clusters of FAPs characterised by different expression levels of SCA-1. To characterise FAP micro-heterogeneity we have used flow cytometry to monitor the expression of FAP surface markers. However, as our antibody panel was limited, the combination of surface markers used in our analysis is not sufficient to clearly distinguish all the interstitial populations involved in muscle regeneration, such as for instance mesoangioblasts (MABs), Pvl1+/Pax7+ interstitial cells (PICs) and Twist2+ progenitors. The 11 unassigned clusters (Supplementary Fig. 1d) that we did not pursue in our analysis may contain these cell types. In addition, the SCA-1 marker is not unique for FAPs, as it is also expressed by mesoangioblasts (MABs). As single-cell RNA-seq analysis also indicates a degree of overlap between MABs and FAPs we wanted to exclude that our cell preparation contained a mixture of FAPs and MABs. To further characterise our preparation, we monitored the expression of alkaline phosphatase, a marker of the perivascular niche which includes MABs and pericytes. While we observed that both cell types express alkaline phosphatase activity, we noticed that SCA1-High-FAPs express higher level of Alpl mRNA. However, the cells in our FAP preparation do not have myogenic potential, a feature that characterise type 2 pericytes and MABs. In addition they express high level of PDGFR-alpha, a surface marker that differentiate FAPs from MABs and pericytes.

Different kinds of cell to cell variability within a monoclonal cell population have been described using single-cell technologies. In particular, the term micro-heterogeneity refers to a condition in which the distribution of the expression of a given trait, an mRNA or a protein, in a cell population is represented by a bell-shaped curve whose standard deviation is larger than expected by random noise or experimental error. We identified such a condition in the expression distribution curve of SCA-1 in FAPs. SCA-1 micro-heterogeneity was already shown to modulate lineage choice in a Erythroid myeloid lymphoid (EML) cell line. This consideration encouraged us to further investigate the role of SCA-1...
heterogeneity in fate decision of dystrophic FAPs. To this purpose, based on the results of our multiplex flow cytometry approach and prior knowledge\(^7\), we resolved the continuity of the distribution of SCA-1 expression, a process defined decomposition\(^3\), into two discrete groups of cells by a fluorescence activated sorting strategy based on SCA-1 expression levels.

Although efforts have been made, clear definitions and terminologies regarding sub-populations and cell states are still lacking. Here we use two in vitro criteria to distinguish these conditions: the nature of their heterogeneity and the ability to maintain their identity once they are separated and cultured in vitro. By applying these criteria, the two groups of dystrophic FAPs were found to be morphologically identical cell states able to repopulate each other in vitro. Although we have shown that also in vivo FAP cell states are not static and repopulate each other to a certain degree, the dynamics and the final equilibrium of such a process is affected by the muscle microenvironment. The extent of this repopulation and its effect on in vivo FAP lineage choice as well as the signals from the niche that determine its dynamics remain to be established.

The involvement of micro-heterogeneity in the decision of cell fate has been addressed in different biological contexts. In the pre-implantation embryo, blastomeres express variable level of CDX2 (Ref.\(^{55}\)). In HeLa cells, the probability and timing of cell death in response to TRAIL are determined by the variance in protein levels of its signalling cascade\(^54\). The role of stem cell heterogeneity during muscle regeneration was recognised only lately\(^8\) and its contribution to FAP phenotype in Duchenne muscular dystrophy (DMD) has been highlighted even more recently\(^8\). However, its role in the development of ectopic tissues in DMD is unexplored. Here we have reported our investigation on the impact of the observed FAP heterogeneity on differentiation and proliferation in FAPs isolated from the muscles of the mdx mouse, an animal model of DMD\(^65\). FAPs are exposed in vivo to a complex and dynamic microenvironment which adds further complexity\(^66\), hence initially we performed our investigation in vitro to take into account only their intrinsic SCA-1 heterogeneity and their response to a controlled microenvironment. We demonstrated that SCA-1 micro-heterogeneity affects the expression of PPAR-gamma in response to an adipogenic medium resulting in a higher propensity of SCA1-High-FAPs to fully differentiate into adipocytes. Moreover, despite FAP cell states showing a similar propensity to differentiate into myofibroblasts, SCA1-Low-FAPs do not increase their expression of Col1a1 and Timp1 mRNAs after fibrogenic stimulation. Interestingly, these cells express higher amount of Col1a1 mRNA in both mdx and wild type mice. We surmise that the expression of Col1a1 gene in SCA1-Low-FAPs do not respond to the fibrogenic medium probably because of an already high basal transcription. Analysis of the transcriptional profile of SCA1-High-FAPs revealed that they preferentially express genes that are likely to make them more sensitive to adipogenic induction by insulin and uncover a possible pro-adipogenic autocrine circuit mediated by adiponectin. We also checked if SCA-1 micro-heterogeneity has an influence on FAP proliferation, a central process in muscle regeneration. The increase of approximately 10 h in the doubling time of SCA1-Low-FAPs decrease their proliferation rate and lengthen the period in which cells are in proliferation. Overall, our results demonstrate that micro-heterogeneity in SCA-1 expression is involved in the fate decision of dystrophic FAPs in a controlled in vitro environment and raise the possibility that this type of cell to cell variability may play a role in the differentiation and proliferation of mesenchymal-like cells in other pathological contexts.

What is known of the function of SCA-1 is not sufficient to rationalise the observed difference in the differentiation potential of the SCA1-High and SCA1-low cell states. Alternatively, and more likely, SCA1 micro-heterogeneity only correlates with the observed phenotypes without being their direct cause. The molecular cause of micro-heterogeneity was attributed to a transcriptome-wide noise originating via a plethora of mechanisms\(^6,7,64\). We have used SCA1 expression mainly as a tool to identify FAPs with divergent phenotypes. A future challenge is the clarification of the origin of this heterogeneity. An additional challenge consists in the elucidation of the physiological relevance of these observations in vivo where further complexity is added by the interaction of micro-heterogeneity with a dynamic microenvironment. To provide some evidence in this respect, we analysed the proliferation of FAPs by looking at the incorporation of EdU after injection into the peritoneum of mdx mice. The results confirmed that in vivo, as in isolated cells, FAPs expressing different level of SCA1 proliferate to a different extent.

Our results show an increase of Perilipin and Cebp\(h\) mRNA in old mdx mice muscles associated with a decreased number of FAPs. These FAPs are also more committed to adipogenic differentiation as demonstrated by the increase of PPAR-gamma positive FAPs. We considered the possibility that the microenvironment under the stress caused by the muscle chronic damage could affect FAP micro-heterogeneity and differentially promote or inhibit the adipogenic potentials of FAP expressing high and low levels of SCA-1. To this end, we repeated the differentiation experiment using adipogenic media conditioned by leucocytes (CD45\(^+\) cells) from young and old mdx mice. In this condition, the two FAP cell states respond differently to the stimuli in the leucocyte
supernatants. Adiogenesis of both cell states is inhibited by leucocyte secreted factor(s). The supernatant of leucocytes from young *mdx* mice is more efficient in restraining adiogenesis of both FAP cell states consistent with the observation that no fat infiltrations are observed in young *mdx* mice. With aging, however, leucocytes become less efficient in inhibiting adiogenesis of SCA1-High-FAPs. These conditions are still far from closely matching the in vivo microenvironment of young and old dystrophic muscles, as for instance the regulatory mechanisms provided by the muscle fibres are not included. However, these observations, suggesting a different age-dependent response of FAP cell states to the anti-adiogenic signals of the immune compartment may explain the increase of IMAT in muscles of old dystrophic mice.

In conclusion, we surmise that micro-heterogeneity in SCA-1 expression plays a role during the decision-making of FAP FAPs in vitro and it is affected by the environment of dystrophic muscles.

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Experiments has been designed, performed and analysed by G.G. with the contribution of S.V., C.F., G.M. and F.R. Sorting strategy has been designed and performed by G.G., E.G. and C.F. Analysis of the Tabula Muris Senis dataset has been carried out by A.P. Mass cytometry experiments was performed by A.R., M.R. and C.F. and was analysed by G.G. Project has been supervised by C.F., C.G., M.V., L.C. and G.C. G.G. and G.C. wrote the manuscript. All authors reviewed the manuscript.

Data availability

The RNA-seq data have been deposited in GEO with the identifier GSE163016.

Code availability

The code for single-cell RNA-seq analysis is available in GitHub at the following link: https://github.com/Alessandro-Palma/Sca1/blob/main/analyse.R.

Ethics statement

This work does not involve human samples.

Conflict of interest

The authors declare that they have no conflict of interest

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