A novel approach for the isolation and long-term expansion of pure satellite cells based on ice-cold treatment

Benedetti Anna¹, Cera Gianluca²,³, De Meo Daniele²,³, Villani Ciro²,³, Bouche Marina¹,⁴ and Lozanoska-Ochser Biliana¹,⁴*

¹Department of Anatomical, Histological, Forensic and Orthopedic Sciences, Section of Histology and Embryology, Sapienza University of Rome, Rome, Italy
²Department of Anatomical, Histological, Forensic and Orthopedic Sciences, Section of Orthopedics, Sapienza University of Rome, Rome, Italy
³Department of Orthopaedics and Traumatology, Policlinico Umberto I, Rome, Italy
⁴Co-senior authors

*Correspondence: biliana.lozanoska-ochser@uniroma1.it
Abstract

Satellite cells (SCs) are muscle stem cells capable of regenerating injured muscle. The study of their functional potential depends on the availability of methods for the isolation and expansion of pure SCs with preserved myogenic properties after serial passages in vitro. Here, we describe the ice-cold treatment (ICT) method, which is a simple, economical and efficient method for the isolation and in vitro expansion of highly pure mouse and human SCs. It involves a brief (15-30 min) incubation on ice (0 °C) of a dish containing a heterogeneous mix of adherent muscle mononuclear cells, which leads to the detachment of only the SCs, and gives rise to cultures of superior purity compared to other commonly used isolation methods. The ICT method doubles up as a gentle passaging technique, allowing SC expansion over extended periods of time without compromising their proliferation and differentiation potential. Moreover, SCs isolated and expanded using the ICT method are capable of regenerating injured muscle in vivo. The ICT method involves minimal cell manipulation, does not require any expertise or expensive reagents, it is fast, and highly reproducible, and greatly reduces the number of animals or human biopsies required in order to obtain sufficient number of SCs. The cost-effectiveness, accessibility and technical simplicity of this method, as well as its remarkable efficiency, will no doubt accelerate SC basic and translational research bringing their therapeutic use closer to the clinic.

Keywords: Satellite cell isolation, Satellite cells in vitro expansion, Skeletal muscle regeneration
Background

The muscle is endowed with an exceptional regenerative ability primarily due to a resident population of stem cells called satellite cells (SCs)(1–3). Ensnconed between the basal lamina and the plasma membrane of muscle fibers, SCs respond to injury or various stress stimuli by becoming activated, and undergoing proliferation, self-renewal, and differentiation to form new myofibers(4,5). As SCs become activated, a proportion of them identified as Pax7⁺MyoD⁻ replenish the stem cell pool, while others acquire the expression of MyoD (Pax7⁺MyoD⁺), differentiate into myoblasts and enter the myogenic program. After several rounds of division myoblasts give rise to Myogenin⁺ myocytes, which fuse together to form new myofibers (1,4,6).

A major stumbling block in the study of the functional potential of SCs has been the lack of isolation methods involving minimal cell manipulation and allowing the isolation and expansion of highly pure SCs with preserved myogenic properties after serial passages in vitro (7–9). Moreover, the success of SC transplantation therapy depends on having an efficient method to isolate and expand these cells in vitro in undifferentiated state and in sufficient numbers (10).

Presently, there are three main methods commonly used for the isolation of SCs, the pre-plating method, Fluorescence Activated Cell Sorting (FACS), and magnetic bead isolation method.

The pre-plating method is based on the differing adhesive properties of muscle cells, with SCs being the least adherent. Following enzymatic digestion, a heterogeneous mix of skeletal muscle cells is plated onto uncoated culture dishes and after a 1-24 h incubation at 37°C, the non-adherent cells are collected and plated onto new collagen coated dishes (7–9,11). The resulting cell culture contains both SCs and fibroblasts in variable proportion. To improve SC purity, the pre-plating step can be repeated every 24 hours over 6 days(7). Although cheap and straightforward to perform, this method’s main disadvantage is that it is time consuming, and gives rise to cultures of variable purity, with fibroblast contamination and overgrowth by day 7 of culture, leading to early senescence and detachment of myotubes (7).

The FACS sorting method sorts muscle mononuclear cells pre-labelled with SC specific antibodies. Following digestion of muscle with various enzymes the resulting mixture of cells is labelled with specific antibodies to facilitate the identification of SCs, which are then sorted using a FACS sorter instrument (12–14). There are several disadvantages to this method including high cost and the requirement for a FACS sorter instrument. Moreover, this method is time consuming, requires
expertise to perform and cell purity can be variable. The cell labelling step followed by the sorting procedure can potentially stress or damage the cells, decrease their viability or alter their functional properties in vitro (9).

Finally, the third method is based on magnetic cell separation (MACS) and uses magnetic columns and SC specific magnetic bead kits (15). It is based on negative selection of SCs by magnetically labelling and removing other cell lineages. Because this method assumes that all the other cell types are successfully removed from the muscle cell preparation, it is less precise than the FACS sorting method. This method is expensive to perform, time consuming and stressful for the cells. As for the other two methods, cell purity is variable and often the SC cultures become overgrown by fibroblasts by day 7 (7,9).

There is therefore a need for new and improved methods for the isolation, expansion and culture of SCs. Here we describe a simple, inexpensive and efficient method for the isolation of highly pure mouse and human SCs that can be serially expanded in vitro to obtain sufficient number of SCs with preserved proliferation potential, capable of regenerating injured muscle in vivo.

**Methods**

**Mice**

C57BL/10ScSn-\textit{Dmd}^{mdx}, C57BL/10ScSn, and C57BL/6J mice were purchased from the Jackson laboratory (Bar Harbor, ME, USA). Both male and female mice were used. The mice were housed in the Histology Department–accredited animal facility at the University of Sapienza. All the procedures were approved by the Italian Ministry for Health and were conducted according to the EU regulations and the Italian Law on Animal Research.

**Human Muscle Sourcing**

Muscle biopsies (gluteus maximus) were obtained from patients (8 male and 7 female, age range 50-90 years) undergoing surgery at the Department of Orthopaedics and Traumatology, Umberto I Hospital in Rome, Italy. According to the Italian law, the authors are not required to ask for approval from an institutional review board or ethical committee for this type of study. In any case, all patients gave their approval to undergo intraoperative muscle biopsy and to publish the clinical and laboratory data obtained.

**Satellite cell isolation with the ice-cold treatment method**
SCs were isolated from hind-limb muscles of 4-8-week-old mice, or from human biopsies. Muscles were dissected with scissors and finely diced with a scalpel in a dish containing DMEM (Sigma-Aldrich). This was followed by enzymatic digestion with 10 ml/g of muscle of Collagenase type II (Sigma-Aldrich) at a concentration of 0.4 mg/ml in PBS (Sigma-Aldrich), for 45 min in a water bath at 37°C with agitation. Digestion was blocked with DMEM 10% FBS and after centrifuging the muscle preparation and removing the supernatant, a second digestion was performed with 10 ml/g of muscle of Collagenase/Dispase at a concentration of 1 mg/ml (Roche, Basel, Switzerland) in PBS Calcium-Magnesium free (Sigma-Aldrich), for 30 min at 37°C in a water bath with agitation. The digested muscle was then passed first through a 70µm cell strainer followed by 40µm cell strainer to obtain single cell suspension. Next, the cells were centrifuged, resuspended in DMEM 10% FBS, counted and plated at 2x10^6 cells /100mm dish (uncoated), and incubated at 37°C for 1h. Non-adherent cells were collected, centrifuged and the cell pellet was resuspended in DMEM 10% FBS, plated again and incubated for another 1h at 37°C. After the second pre-plating non-adherent cells were collected, centrifuged, counted, resuspended in SC Growth Medium (GM) (DMEM, 20% Horse Serum, 3% Chicken Embryo Extract), and plated into 0.1% gelatin coated 100 mm dishes, at 10^6 cells / dish. The next day, the dishes containing a heterogeneous mix of adhered muscle cells were washed 3 times with PBS, and 10 ml of ice-cold PBS was added into each dish. The dishes were then placed on ice (0 °C) for 15-30 min with occasional gentle manual shaking. The detached cells were collected, centrifuged, resuspended in GM and plated into 0.1% gelatin coated 35 mm dishes at a density of 10^3 cells/dish. To induce differentiation, proliferating cells (day 3 after adding GM) were cultured in differentiating medium containing DMEM 5% Horse Serum, 1% Chicken Embryo Extract.

**Satellite cell isolation with magnetic bead labelling**

Mouse SC isolation by magnetic bead labelling was performed by using a SC Isolation Kit (Miltenyi Biotech, Bergisch Gladbach, Germany) as previously described(16). Briefly, minced muscle was digested as described above. The digested muscle was passed through 70µm and 40 µm cell strainers, and the resulting single cell suspension was centrifuged, resuspended in 80 µl buffer (PBS pH 7.2, 0.5% FBS, 2mM EDTA) and incubated with 20 µl of Satellite Cell Isolation Kit per gram of muscle, for 15 min at 4°C. Next, the cell suspension was passed through a LS column placed in a magnetic field of a MACS Separator. Unlabeled SCs were collected in the flow-through, counted, washed, resuspended in growth medium (GM) and plated into 35 mm dishes at a density of 10^3 cells/dish.

Mouse and human SCs were cultured either in Growth Medium containing DMEM, 20% Horse Serum (Sigma-Aldrich, St. Louis, MS, USA), 3% Chicken Embryo Extract, or in Differentiation Medium containing DMEM 5% Horse Serum, 1% Chicken Embryo Extract.
**SC transplantation**

Acute muscle injury was induced the day before SC transplantation. To induce muscle injury tibialis muscles were injected with 0.01 ml of Cardiotoxin from Naja Pallida (10 µM) (Latoxan ZA Les Auréats, France), using a 30 Gauge micro-syringe (16–19).

For cell transplantation, 15,000 SCs were resuspended in 20µl of DMEM 2% FBS (Sigma-Aldrich, St. Louis, MS, USA) and injected into the TA muscle of one leg with a single injection by using a 30 Gauge micro-syringe. Contralateral TA muscle was injected with only PBS and used as control.

**Immunofluorescence and microscopy**

For immunofluorescence analysis cultured SCs were fixed in PFA 4% for 10 min RT, permeabilized in cold methanol at –20°C for 6 min, blocked in 5% Goat Serum in PBS for 30 min RT, and incubated overnight at 4°C in 4% BSA in PBS with the following primary antibodies: mouse anti-Pax7 (1:10 DSHB, Iowa City, IA, USA), rabbit anti-MyoD (1:50 Santa Cruz C20: sc-304, Dallas, TX, USA), mouse anti-myogenin (1:20, DSHB), mouse anti-Myosin Heavy Chain (1:20 DSHB), and mouse anti-desmin (1:20, DSHB). The next day, SCs were washed 3 times in PBS for 15 min, and then incubated with secondary antibodies goat anti-rabbit Alexa Fluor 488 (1:1000, Abcam) and goat anti-mouse Alexa Fluor 555 (1:1000, Thermo Fisher Scientific, Waltham, MA, USA) diluted in 1% BSA in PBS, for 1h RT. Nuclei were counterstained with Hoechst.

For immunofluorescence analysis of mdx TA muscle transplanted with WT SCs, 8 µm thick muscle cryosections were fixed in 4% PFA for 10 min at room temperature (RT), and then permeabilized in cold methanol for 6 min at –20°C. Sections were then blocked in 5% Goat Serum (Sigma-Aldrich St. Louis, MS, USA) in PBS for 30 min at RT. Next, sections were incubated with primary rabbit anti-dystrophin antibody (1:200 Abcam Cambridge, UK) overnight at 4°C. The next day, sections were washed and incubated with a secondary antibody goat anti rabbit Alexa Fluor 488 (1:1000 Abcam). Nuclei were counterstained with Hoechst.

Samples were analyzed under an epifluorescence Zeiss Axioskop 2 Plus microscope (Carl Zeiss, Oberkochen, Germany).

Bright field images were acquired with an inverted phase-contrast microscope (Nikon Eclipse, TS100). Images were acquired with a Nikon DS-Fi2 camera and NIS Elements version 4.0 Imaging System.

**CFSE staining**
Isolated SCs were stained with CFSE (ThermoFisher Scientific, Waltham, MA, USA) at a concentration of 1µM for 10 min at 37°C in the dark prior to culture. After 4 days of culture, the cells were detached with Accutase Solution (Sigma-Aldrich). Samples were acquired with a CyAn ADP (DAKO) flow cytometer and acquired data were analysed using FlowJo software version 10 (FlowJo LLC, Ashland, OR, USA).

**RNA isolation and Quantitative real-time PCR**

For RNA preparation, cells were lysed with TRI reagent (Sigma-Aldrich) and processed as previously described (16). The following primers were used for amplification: Pax7 (FW: 5’ GTCCCACTCTTACTGGCCAC 3’, RV: 5’ TGTGGACAGGCTACCTAG 3’), Myogenin (FW: 5’ GCATGGAGTCCGACTG3’, RV: 5’ TATCCTCCACCGTGATGCTG 3’), GAPDH (FW: 5’ ACCCAGAAGACTGTGGATGG 3’, RV: 5’ CACATTGGGGGTAGGAACAC 3’). All Real Time PCR results were normalized to GAPDH and expressed as fold-change in expression levels. Data analysis was performed using 7500 Software v2.0.6 provided by Applied Biosystems (Foster City, California).

**Clonal myogenicity assay**

For the clonal myogenicity assay SCs were plated into 0.1% gelatin coated 96-well plates, at a density of 1 cell/well, and grown in growth medium. Colony formation and number of cells were assessed at 24, 48 and 72h of culture.

**Statistical analysis**

All statistical analyses were performed using GraphPad Prism software version 8 (La Jolla, CA, USA). Data are presented as mean ± SEM. Statistical significance was determined using unpaired 2-tailed Student’s t test with Welch’s correction for unequal variances. A P value of ≤ 0.05 was considered statistically significant.

**Results**

**Isolation and characterisation of muscle SCs using the ice-cold treatment method.** Previous studies have demonstrated that cold temperature causes a reduction in cell adhesion, likely due to the downregulation of adhesion receptors (20,21). Compared to other cells, such as fibroblasts, which typically contaminate SC cultures, SC are considered to be less adherent (8,11). Taking this into consideration, as well as the notion that like all stem cells, SCs are sensitive to stress signals and are
among the first muscle resident cells to respond to injury (4,5), we hypothesised that subjecting a heterogeneous culture of muscle cells to a mild stress stimulus such as ice-cold temperature will lead to the detachment of only the SCs. To test this hypothesis, we obtained a mix of muscle cells following enzymatic digestion, and after 2 hours of pre-plating on uncoated dishes, cultured them overnight on gelatin-coated dishes. The next day, after washing and removing the non-adhered cells and debris, we placed the dishes of heterogeneous muscle cells on ice for 30’ (Figure 1A and B). This time point was chosen based on the purity and number of SCs obtained following 15, 30, 45 and 60’ on ice. Although the number of SCs obtained increased with longer incubations, the purity of SCs decreased from 100 % at 15-30’ to 95 and 90 % at 45 and 60’ on ice, respectively (data not shown). The ice-cold treatment (ICT) method can be used to harvest SCs from the original dish containing the heterogeneous muscle cells for at least the first 3 days of culture. Placing the heterogeneous muscle culture dish on ice for 30’ resulted in the detachment of only the SCs giving rise to a highly pure culture of SCs that proliferated and differentiated into myotubes upon culture in differentiating medium (Figure 1B-D). Satellite cells isolated using the ICT method were 100 % pure as determined by the expression of the SC markers Pax7 and MyoD (Figure 1C). At day 3 of proliferation, 100 % of the cells were positive for the satellite cell marker Pax7 and of these 97 % were activated and expressed MyoD (Pax7*MyoD+) (Figure 1C). To examine the myogenic capacity of ICT-isolated SCs we performed a clonal myogenicity assay, by plating a single cell per well and analysing the formation of myogenic colonies. Satellite cells isolated with the ICT method displayed a similar clonal myogenicity of 40 % and a doubling time of 17h to SCs isolated using the magnetic beads method (Figure S1A and B). The myogenic identity of the cultured cells was further confirmed by the expression of myosin heavy chain (MHC) and the formation of myotubes. After 3 days in differentiating medium the SCs differentiated into myoblasts that fused into MHC expressing myotubes with a fusion index of 90 % (Figure 1D). The purity of the isolated SCs at the beginning of culture, as well as the fusion index and number of myonuclei per myotubes were similar between SCs isolated with the ICT and magnetic cell separation (MACS) method (Figure S1C-E). However, as the cells grew and differentiated into myotubes the culture obtained with the MACS method became overgrown by fibroblasts at day 7 after plating causing premature myotube detachment, whereas cultures obtained with the ICT method remained 100% free of contaminating cells (Figure 1E).

Overall, these results confirm our hypothesis that ICT leads to the preferential detachment of SCs and using this method allows the isolation of 100 % pure SCs.
The ICT approach can also be used for serial passaging and long-term expansion of SCs. A major obstacle in SC research has been that cultured SCs lose their proliferation potential after a couple of passages and begin to differentiate into myotubes thereby limiting the number of SCs that can be serially expanded in vitro. Loss of differentiation potential has also been noted with increasing number of passages (7–10,22). We reasoned that, since the ICT leads to SC detachment, it can also be used to passage growing cultures of proliferating SCs. Indeed, placing the dishes of proliferating SCs on ice for 30 min led to the detachment of around 30% of the proliferating cells and we were able to serially passage and expand the proliferating SCs without compromising their proliferation and differentiation capacity (Figure 2A). We used the ICT approach on proliferating SCs and then successively on each established culture of proliferating SCs (for more than 10 passages). This approach yielded on average 2.5 x10^6 cells / g of muscle (Figure 2B). The SCs passaged using the ICT method did not lose their proliferation and differentiation potential and displayed a minimally altered Pax7 and myogenin gene expression (Figure 2C and D). While ICT-passaged SCs retained their proliferation and differentiation capacity even after 10 passages, SCs passaged with the most commonly used passaging reagent trypsin, lost the ability to form new myogenic colonies after just 2 passages and instead begun to differentiate into myotubes (Figure 2E-G). Similar to SCs detached by trypsin, SCs detached using a gentler detachment solution such as Accutase, exhibited accelerated differentiation after more than 3 passages (data not shown).

Previous studies have demonstrated that based on their rate of proliferation, SCs can be divided into two subpopulations: fast dividing and slow dividing (23). The slow-dividing are a subset of SC that have been shown to retain stemness and long-term self-renewal ability (24). Unlike the fast-dividing SCs, which have a limited ability to form secondary myogenic colonies after passage and instead undergo differentiation, the slow dividing SCs form secondary myogenic colonies when passaged (23,24). To examine the prevalence of fast and slow dividing SCs within our population, we labelled the ICT-passaged SCs with CFSE prior to plating and analysed the rate of proliferation 3 days later. We compared the proliferation rate of ICT-isolated and passaged SCs with that of SCs isolated using magnetic beads and passaged with trypsin. In line with previous observations, we found that most of the activated SCs isolated using the commercial kit and passaged with trypsin were fast-dividing cells (CFSE^lo), with slow-dividing cells (CFSE^hi) representing less than 10% of the total. Interestingly, while similarly heterogeneous, the SCs isolated and passaged using the ICT method were enriched in the slow dividing SC population (Figure 2H and I).

Next, we examined the longevity of differentiated ICT-isolated SCs in culture. The highly pure cultures of SCs isolated using the ICT method gave rise to myotubes that could be maintained in
culture for up to 1 month, compared to just 7 days when isolated using the magnetic beads (Figure S2A-C), or the pre-plating method (Figure S2D).

Overall, these data show that the ICT method can be used for the serial expansion of SCs with preserved proliferation and myogenic potential over an extended period of time compared to other methods.

**Satellite cells isolated and expanded with the ICT method efficiently engraft and regenerate injured muscle.** Satellite cells play a central role in the maintenance of muscle homeostasis and regeneration after injury(4,25–27). To examine the potential of ICT-isolated and expanded SCs to regenerate injured muscle, we injected 15,000 freshly isolated or in vitro expanded SCs from wild type mice into the tibialis anterior of mdx mice (lacking dystrophin) CTX injured 24 h previously. Transplantation of both freshly isolated and in vitro expanded SCs contributed to the regeneration process to a similar extent as evidenced by the appearance of newly formed dystrophin positive fibers (Figure 3A and B). There was no difference in the ability of transplanted ICT-isolated SCs and SCs isolated by magnetic beads to regenerate injured muscle (Figure S3A and B). These experiments confirmed that SCs isolated with the ICT method successfully engraft after transplantation and do not lose their potential to regenerate injured muscle after expansion in vitro.

**Efficient isolation, serial expansion and long-term culture of human satellite cells with the ICT method.** The study of human SCs (hSCs) has generally lagged behind that of mouse SCs due to the difficulties associated with obtaining muscle tissue, as well as the lack of methods for the isolation of pure hSCs that can be expanded in vitro without altering their myogenic potential (10). Having demonstrated the remarkable efficiency of the ICT method in the isolation of pure mouse SCs, we set to reproduce these findings using human muscle biopsies. We obtained gluteus maximus specimens from patients undergoing surgery, aged between 50 and 90 years. Using the same approach as described above, we consistently and reproducibly obtained a highly pure culture of hSCs (Figure 4A, D-G) from a heterogeneous population of human muscle cells (Figure S4), that could be serially expanded for more than 10 passages (Figure 4A). As previously reported (28) we found that hSCs proliferated slower than mouse SCs, with a doubling time of 46h (Figure 4B), reaching a peak around day 10 post isolation, and slowing down thereafter. On average we isolated around 20 x 10³ SCs /g of muscle. Using the ICT method, these hSCs could be expanded 300-fold over a period of 2 months to a final total of 6 x10⁶ / g of muscle (Figure 4C). Previous studies have shown that hSCs rapidly
downregulate Pax7 expression in culture (29). In agreement with others (29) we found that the expression of Pax7 was variable and ranged between 45 and 50% at day 2 of culture in GM (Figure 4D). Almost 100% of the hSCs were myogenin positive at day 5 of culture in GM, suggesting that most of them have activated their myogenic program (Figure 4E). The myogenic purity of the hSC culture was further confirmed by desmin at day 5 (Figure 4F) and MHC expression at day 10 after shifting to differentiating medium (Figure 4G).

These data show that the ICT method performs equally well when used for the isolation of human SCs.

Discussion

Over the past decade, a considerable progress has been made in the development of new methodologies for the isolation of SCs (7,11–13,28,30–34). Nevertheless, each of the available methods suffers from at least one disadvantage be it purity, cost, expertise, or a combination of these(8). In this study we describe a novel method for the isolation of pure mouse and human SCs, that is inexpensive, simple to perform and reproducibly efficient. The ICT method takes advantage of the differing adhesive properties of muscle cells as well as the ability of SCs to rapidly respond to stress stimuli (5,8,11). Thus, the combination of a mild cold-stress stimulus and cold-induced reduction in adhesion, leads to the detachment of only the SCs. Exposure of mammalian cells to cold stress can slow down the progression through the cell cycle and inhibit protein synthesis. Moreover, depending on the intensity and duration, cold stress can activate the apoptotic program, or lead to necrosis (35). Thus, it is conceivable that prolonged exposure to cold temperature could interfere with the myogenic properties of SCs. Nevertheless, we found that the brief period of exposure to ice-cold temperature did not interfere with the ability of SCs to proliferate and differentiate in vitro or with their in vivo regeneration potential, suggesting that ICT does not alter SC function. Indeed, SCs isolated by ICT behaved similarly to those isolated using the pre-plating method or the magnetic bead isolation kit. Of note, both the magnetic bead and the FACS sorting isolation methods involve far lengthier incubation times on ice compared to our method without interfering with SC behaviour and function in vitro and in vivo (12,13,15). Interestingly, Marg A et al recently found that storing human muscle biopsies at 4°C, in low serum medium and no O₂ for up to 35 days and subsequent culture at 37°C and 21% oxygen led to SCs expansion outside the fiber fragments. Surprisingly, the purity of the outgrowing colonies of SCs was 100% myogenic cells, since non-myogenic cells such as fibroblasts did not survive prolonged storage at hypothermic conditions (29). Therefore, it is likely
that SCs tolerance for cold stress is high compared to other cell types, and this trait can be exploited as we did with our ICT method, to improve the purity of SCs grown in vitro.

In addition to performing remarkably well for SCs isolation, the ICT method doubles up as a very gentle passaging technique, allowing long-term serial expansion of SCs ex vivo, without altering their proliferation and differentiation properties. After each ICT passage, the already expanded SCs can be cryopreserved, and stored until needed. With the ICT method we were able to passage proliferating mouse and human SCs for at least 10 times, expanding their number 150- and 300-fold respectively. This represents a clear advantage over the most commonly used passaging reagent trypsin, which we and others have shown, typically accelerates the differentiation of passaged SCs after only two passages (8). Apart from being a relatively harsh enzymatic passaging reagent, trypsin leads to the detachment of all the cells in the dish including SCs that are already committed to differentiate into myotubes, as well as any contaminating cells, which might contribute towards the loss of SC proliferative potential and accelerated differentiation. Indeed, even a gentler detachment solution like Accutase leads to loss of myogenic proliferative properties and accelerates differentiation. By contrast, the ICT approach favours the detachment of only the SCs that have not yet committed to differentiate, and in particular the slow dividing population which has previously been shown to retain stemness and long-term self-renewal ability (24). It is conceivable that being ‘true’ stem cells, the slow dividing SCs detaching in response to cold temperature are the so-called first responders to stress or injury in vivo (5), a hypothesis that will be the subject of future investigation in our laboratory.

In a recent study, Gregory WC et al demonstrated that huSCs differentiate and lose their proliferative potential when maintained in high mitogen conditions ex vivo. They used inhibition of p38 signalling to prevent the differentiation of SCs and promote their expansion (32). Using our method, we were able to achieve the same but with minimal manipulation, maintaining the proliferative capacity of hSC ex vivo for an extended period of time, to a similar degree using muscle biopsies taken from a wide range of ages (between 50 and 90 years old). This is an important technical advance for both basic and clinical research since it will allow researchers to obtain sufficient number of cells for transplantation or intervention studies, while reducing the number of human biopsies required. Indeed, a major obstacle to stem cell-based therapies has been the scarcity of human muscle tissue specimens and the limited number of cells that can be obtained for transplantation. Similarly, with the ICT method the number of animals needed to be sacrificed is dramatically reduced.

Moreover, successful transplantation requires the use of freshly isolated SCs because culturing and expanding them in vitro greatly reduces their engraftment capacity (10). Here, we show that SCs isolated and expanded using the ICT method do not lose their regenerative capacity.
Another advantage of the ICT method is the improved longevity of cultured myotubes. Generally, SCs differentiate into myotubes by day 7 of culture and shortly after, begin to detach (8,9). Notably, SCs isolated and passaged with the ICT method could be maintained in culture for up to 2 months, even once they have differentiated into myotubes, likely due to the lack of contaminating cells such as fibroblasts. The purity of the isolated SCs is of paramount importance for in vitro studies since even 97 percent purity is insufficient to prevent overgrowth by fibroblasts, as we demonstrated. While important for cell growth, growth factors produced by fibroblasts, have been linked to senescence induction in long term cultures of mesenchymal stem cells(36).

Conclusions

In the quest for new and improved SC isolation methods, the ideal technique would permit the isolation of pure SCs with minimal manipulation, that can be expanded ex vivo without losing their stemness and regenerative capacity. In terms of purity of the isolated cell population, the ICT method outperforms others such as the pre-plating method or the magnetic beads isolation method. Compared to other commonly used methods, it is fast and easy to perform, and apart from the time required for enzymatic digestion (1.5 h), it involves minimal manipulation of the cells. Finally, using the ICT approach, SCs can be expanded for extended periods of time without losing their proliferation and differentiation potential. This in turn drastically reduces the number of mice or muscle biopsies required to obtain sufficient number of cells.

Overall, the cost-effectiveness, accessibility and technical simplicity of this method, as well as its remarkable efficiency, represent major improvements over existing methods, and will no doubt accelerate SC basic and translational research bringing their therapeutic use closer to the clinic.

Declarations

Ethics approval and consent to participate

All procedures involving mice were approved by the Italian Ministry for Health and were conducted according to the EU regulations and the Italian Law on Animal Research.

According to the Italian law, the authors are not required to ask for approval from an institutional review board or ethical committee for the use of human tissue removed during surgical procedures. All patients gave their approval to undergo intraoperative muscle biopsy and to publish the clinical and laboratory data obtained.
Consent for publication

Not applicable.

Availability of data and materials

All data generated or analysed during this study are included in this published article [and its supplementary information files].

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

Funding

This work was supported by a grant from The Dutch Duchenne Parent Project NL (DPP NL) to BLO; research grants from Parent Project Italy (PP, Italy) to MB and from the University of Rome (# RP11715C7D238352, RM118164275C7EBE and RM11916B7E20311C to MB, and # AR11715C7F9E158E, AR11816436905518 and AR11916B7E2A7B64 to AB).

Authors’ contributions

BLO conceived, developed and standardized the method. BLO, AB and MB designed the experiments. AB performed experiments. AB and BLO analysed data and prepared the figures. BLO and MB supervised the study. CG, DMD and VC provided the human muscle biopsy specimens. BLO wrote the manuscript. All authors critically read, edited and approved the final manuscript.

Acknowledgements

We thank Dr Luca Madaro for critical reading of the manuscript and helpful discussions.
References

1. Chang NC, Rudnicki MA. Satellite Cells: The Architects of Skeletal Muscle. In: Current Topics in Developmental Biology. 2014.

2. Wang YX, Dumont NA, Rudnicki MA. Muscle stem cells at a glance. Journal of Cell Science. 2014;

3. MAURO A. Satellite cell of skeletal muscle fibers. The Journal of biophysical and biochemical cytology. 1961;

4. Wang YX, Rudnicki MA. Satellite cells, the engines of muscle repair. Nature Reviews Molecular Cell Biology. 2012.

5. Evano B, Tajbakhsh S. Skeletal muscle stem cells in comfort and stress. npj Regenerative Medicine. 2018.

6. Feige P, Rudnicki MA. Muscle stem cells. Current Biology. 2018.

7. Keire P, Shearer A, Shefer G, Yablonka-Reuveni Z. Isolation and culture of skeletal muscle myofibers as a means to analyze satellite cells. Methods in Molecular Biology. 2013;

8. Danoviz ME, Yablonka-Reuveni Z. Skeletal muscle satellite cells: Background and methods for isolation and analysis in a primary culture system. Methods in Molecular Biology. 2012;

9. Jonah D Lee BCS, Lisa M Larkin KWV. Isolation and Purification of Satellite Cells for Skeletal Muscle Tissue Engineering. Journal of Regenerative Medicine. 2015;

10. Rinaldi F, Perlingeiro RCR. Stem cells for skeletal muscle regeneration: Therapeutic potential and roadblocks. Translational Research. 2014.

11. Gharaibeh B, Lu A, Tebbets J, Zheng B, Feduska J, Crisan M, et al. Isolation of a slowly adhering cell fraction containing stem cells from murine skeletal muscle by the preplate technique. Nature Protocols. 2008;

12. Pasut A, Oleynik P, Rudnicki MA. Isolation of muscle stem cells by fluorescence activated cell sorting cytometry. Methods in Molecular Biology. 2012;

13. Liu L, Cheung TH, Charville GW, Rando TA. Isolation of skeletal muscle stem cells by fluorescence-activated cell sorting. Nature Protocols. 2015;

14. Chapman MR, Balakrishnan KR, Li J, Conboy MJ, Huang H, Mohanty SK, et al. Sorting single satellite cells from individual myofibers reveals heterogeneity in cell-surface markers and myogenic capacity. Integrative Biology (United Kingdom). 2013;

15. Blanco-Bose WE, Yao CC, Kramer RH, Blau HM. Purification of mouse primary myoblasts based on α7 integrin expression. Experimental Cell Research. 2001;

16. Benedetti A, Fiore PF, Madaro L, Lozanoska-Ochser B, Bouché M. Targeting pkcθ promotes satellite cell self-renewal. International Journal of Molecular Sciences. 2020;
17. Fiore PF, Benedetti A, Sandonà M, Madaro L, de Bardi M, Saccone V, et al. Lack of PKCθ promotes regenerative ability of muscle stem cells in chronic muscle injury. International Journal of Molecular Sciences. 2020;

18. Rizzo G, di Maggio R, Benedetti A, Morroni J, Bouche M, Lozanoska-Ochser B. Splenic Ly6Chi monocytes are critical players in dystrophic muscle injury and repair. JCI Insight. 2020;

19. Lozanoska-Ochser B, Benedetti A, Rizzo G, Marrocco V, di Maggio R, Fiore P, et al. Targeting early PKCθ-dependent T-cell infiltration of dystrophic muscle reduces disease severity in a mouse model of muscular dystrophy. Journal of Pathology. 2018;

20. Juliano RL, Gagalang E. The adhesion of Chinese hamster cells. I. Effects of temperature, metabolic inhibitors and proteolytic dissection of cell surface macromolecules. Journal of Cellular Physiology. 1977;

21. Rico F, Chu C, Abdulreda MH, Qin Y, Moy VT. Temperature modulation of integrin-mediated cell adhesion. Biophysical Journal. 2010;

22. Machida S, Spangenburg EE, Booth FW. Primary rat muscle progenitor cells have decreased proliferation and myotube formation during passages. Cell Proliferation. 2004;

23. Tierney MT, Sacco A. Satellite Cell Heterogeneity in Skeletal Muscle Homeostasis. Trends in Cell Biology. 2016.

24. Ono Y, Masuda S, Nam HS, Benezra R, Miyagoe-Suzuki Y, Takeda S. Slow-dividing satellite cells retain long-term self-renewal ability in adult muscle. Journal of Cell Science. 2012;

25. Sambasivan R, Yao R, Kissenpfennig A, van Wittenberghhe L, Paldi A, Gayraud-Morel B, et al. Pax7-expressing satellite cells are indispensable for adult skeletal muscle regeneration. Development. 2011;

26. Relaix F, Zammit PS. Satellite cells are essential for skeletal muscle regeneration: The cell on the edge returns centre stage. Development (Cambridge). 2012.

27. Tedesco FS, Delvalle A, Diaz-Manera J, Messina G, Cossu G. Repairing skeletal muscle: Regenerative potential of skeletal muscle stem cells. Journal of Clinical Investigation. 2010.

28. Charville GW, Cheung TH, Yoo B, Santos PJ, Lee GK, Shrager JB, et al. Ex vivo expansion and in vivo self-renewal of human muscle stem cells. Stem Cell Reports. 2015;

29. Marg A, Escobar H, Gloy S, Kufeld M, Zacher J, Spuler A, et al. Human satellite cells have regenerative capacity and are genetically manipulable. Journal of Clinical Investigation. 2014;

30. Montarras D, Morgan J, Collins C. Direct Isolation of Satellite Cells for Skeletal Muscle Regeneration. Molecular and Cellular Biology. 2005;

31. Garcia SM, Tamaki S, Lee S, Wong A, Jose A, Dreux J, et al. High-Yield Purification, Preservation, and Serial Transplantation of Human Satellite Cells. Stem Cell Reports. 2018;
Figure Legends

Figure 1. Isolation and characterisation of muscle SCs using the ICT method. A. Schematic representation of the ICT method. B. Representative bright field images of the heterogeneous muscle mononuclear cell culture from which SCs were isolated by ICT, and representative images of the ICT-isolated SCs at D2 and D4 in growth medium (GM) and at D3 after adding differentiation medium (DM). C. Representative immunofluorescence images of ICT-isolated SCs stained with Pax7 and MyoD, and a graph showing the percentage of cells positive for Pax7 and/or MyoD at day 3 of culture in GM (n=3 independent experiments). D. Representative immunofluorescence images of ICT-isolated SCs stained with myosin heavy chain (MHC) and a graph showing percent fusion after
differentiation (4 days in GM followed by 3 days in differentiating medium (DM)) (n= 3 independent experiments). E. Representative bright field images of ICT- and magnetic beads (MB)- isolated SCs and a quantification graph showing percentage of myogenic cells in ICT- and MB-isolated SCs after differentiation (n= 3 independent experiments). Scale bar=100µm. Error bars represent mean ± sem. *P< 0.05 by Student’s t-test.

Figure 2. Serial expansion and long-term proliferation potential of SCs using the ICT method. A. Representative bright field images of SCs at day 3 of culture in GM following ICT 1, ICT 2 and ICT 3, and differentiated cells at day 3 of culture in DM following ICT 3. B. Total number of SCs at day 3 of culture in GM, after 1 and 10 ICTs. C. Pax7 gene expression in SCs at day 2 of culture in GM after ICT 1 and ICT 2 analysed by quantitative real time PCR (n=3 independent experiments). D. Myogenin expression in SCs at day 5 of culture in GM after ICT1 and ICT 2 analysed by quantitative real time PCR (n=3 independent experiments). E. Representative images of SCs at day 2 of culture in GM, after 1-3 detachments with ICT (top panels) and trypsin (bottom panels). F. Pax7 gene expression in SCs detached with ICT or trypsin, at day 2 of culture in GM, analysed by quantitative real time PCR (n=2 independent experiments). G. Myogenin gene expression in SCs detached with ICT or trypsin, at day 5 of culture in GM, analysed by quantitative real time PCR (n=2 independent experiments). H. Representative overlay of histogram plots of CFSE labeled ICT and trypsin detached SCs. I. Graph showing percent CFSE-low or slow-proliferating SCs after ICT or trypsin detachment, at Day 4 of culture in GM (n=3 independent experiments). Error bars represent mean ± sem, *p < 0.05, **p from 0.01 to 0.05 calculated by Student’s t-test.

Figure 3. In vivo functional validation of SCs isolated using the ICT method. A. Representative immunofluorescence images of dystrophin positive fibers (green) and nuclei (blue) in mdx tibialis muscle at 30 days following intra-muscular injection of 15 x 10^3 SCs freshly isolated or expanded (3 days in culture) using the ICT method. B. Quantification of the number of dystrophin positive fibers per TA muscle section in mdx mice (Freshly isolated, n=5 mdx mice; Cultured, n=4 mdx mice). Scale bar=100µm. Error bars represent mean ± sem.

Figure 4. Isolation and in vitro expansion of human satellite cells using the ICT method. A. Representative bright field images of ICT- isolated human SCs following 1, 3, 5 and 10 ICTs, at day 3 and 10 of culture in GM, and at day 10 of culture in DM. B. Number of human SCs at 24, 48 and 72h following ICT 1. C. Total number of human SCs at day 3 of culture in GM after 1 and 10 ICTs.
D. Representative immunofluorescence images of human SCs stained for Pax7 (red) and nuclei (blue). Graph shows percentage of cells positive for Pax7 at day 2 of culture in GM. E. Representative immunofluorescence images of human SCs stained for myogenin (red) and nuclei (blue). Graph shows percentage of cells positive for myogenin at day 5 of culture in GM. F. Representative immunofluorescence images of human SCs stained for desmin (red) and nuclei (blue) after differentiation (10 days in GM + 5 days in DM). G. Representative immunofluorescence images of human SCs stained for myosin heavy chain (MHC) (red) and nuclei (blue). Graph shows percent fusion after differentiation (10d GM + 10d DM). (n=10 images analysed). Scale bar=100μm. Error bars represent mean ± sem.

**Figure S1. Myogenic properties of SCs isolated with the ICT method.** A. Percent of myogenic colony formation among ICT- and MB-isolated SCs. B. Number of cells per clone in single clone-derived ICT- and MB-isolated SCs at 48 and 72 h of culture in GM. (n=2 independent experiments). C. Percent of ICT- and MB-isolated SCs positive for Pax7 at day 2 of culture in GM. (n=3 independent experiments). D. Percent fusion of single clone-derived ICT and MB-isolated SCs after differentiation (4 days in GM followed by 3 days in DM). Fusion index: number of nuclei within myotubes divided by total number nuclei. E. Number of nuclei per myotube in single clone-derived ICT and MB-isolated SCs after differentiation (4 days in GM followed by 3 days in DM). Error bars represent mean ± sem.

**Figure S2. Increased longevity in culture of ICT-isolated SCs.** A. Representative bright field images of ICT-isolated SCs at day 3, 5, 11 and 17 of culture in DM. B. Representative bright field image of MB isolated SCs at day 3 of culture in DM. C. Total number of days in culture of ICT- and MB-isolated SCs. D. Representative bright field images showing the heterogeneous muscle cell culture after pre-plating, at 3 and 5 days of culture in GM. Error bars represent mean ± sem. *P < 0.05 by Student’s t-test.

**Figure S3: In vivo functional validation of SCs isolated using the ICT and MB method.** A. Representative immunofluorescence images of dystrophin positive fibers (green) and nuclei (blue) in mdx tibialis muscle at 30 days following intra-muscular injection of 15 x 10^3 ICT-freshly isolated SCs (left), or MB-freshly isolated SCs (right). B. Quantification of the number of dystrophin positive
fibers per TA section in mdx mice (ICT, n=5 mdx mice; MB, n=5 mdx mice). Scale bar=100µm. Error bars represent mean ± sem.

**Figure S4. Characterisation of the human muscle-derived cells obtained after pre-plating and prior to ICT.** A. Representative bright field images of human muscle-derived cells at day 3 and 10 of culture in GM, and at day 10 of culture in DM. B. Representative immunofluorescence images of the heterogeneous culture of human muscle-derived cells stained for Pax7 (red) and nuclei (blue). Graph shows percentage of cells positive for Pax7 at day 2 of culture in GM. C. Representative immunofluorescence images of the heterogeneous human muscle cell culture stained for myogenin (red) and nuclei (blue). Graph shows percent of cells positive for myogenin at day 5 of culture in GM. D. Representative immunofluorescence images of the heterogeneous culture of human muscle-derived cells stained for MHC (red) and nuclei (blue). Graph shows percent cell fusion after differentiation (10 days in GM followed by 10 days in DM). E. Graph shows percent myogenic cells in the human heterogeneous muscle cells obtained after pre-plating or after ICT at day 5 of culture in GM, calculated by IF staining for myogenin (n=10 images analysed). Scale bar=100µm. Error bars represent mean ± sem. ****P < 0.0001 by Student’s t-test.