Debrided Skin as a Source of Autologous Stem Cells for Wound Repair

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Key Words. Debrided skin • Adipose stem cells • Burn wounds • Perivascular niche

ABSTRACT

Major traumatic injuries to the body, such as large surface area burns, limit the availability of autologous stem cell populations for wound repair. This report demonstrates that even after severe burn trauma to the body, resident stem cells present within the subcutaneous adipose tissue survive and are available for therapeutic uses. Debrided skin from wounded areas contains subcutaneous adipose tissue and can yield approximately $1.5 \times 10^5$ to $2.5 \times 10^5$ cells per milliliter of tissue. This observation indicates that tissue, which is normally discarded, could be a valuable source of stem cells. Initial immunohistochemistry of the debrided tissue localized platelet-derived growth factor receptor beta$^+$ (PDGFR-$\beta^+$) cells to perivascular niches of vascular beds. It was immunophenotypically confirmed that the cell isolates are stem cells and designated as debrided skin adipose-derived stem cells (dsASCs). Gene expression analysis of stem cell specific transcripts showed that the dsASCs maintained their stemness over serial passages. Furthermore, dsASCs were able to differentiate into adipogenic, osteogenic, and vascular cell lineages. Finally, an in vivo excision wound model in athymic rats demonstrated that the dsASCs are engrafted within a wound bed after 12 days. These data provide the first evidence that subcutaneous adipose tissue from discarded burned skin contains a viable population of stem cells that can be used for wound repair and skin regenerative therapies.

BACKGROUND

Burn wounds constitute 5%-10% of military casualties, involving 40%-60% of total body surface area (TBSA) [1–3], and require extensive reconstruction [4, 5]. Stem cells, a valuable tool in regenerative medicine, have demonstrated the capacity to help heal extensively injured tissues [6, 7]. Among adult mesenchymal stem cells (MSCs), adipose-derived stem cells (ASCs) have gained particular attention due to their ease of isolation, relative abundance [8–12], and multilineage differentiation potential [13–19]. Recent studies of various tissues, including adipose tissue, indicate that their resident stem cell niches localize to perivascular regions and express markers of both MSCs (CD105, STRO-1, and CD34) and pericytic (NG2 and platelet-derived growth factor receptor beta (PDGFR-$\beta^+$)/CD140b) lineages [20–24]. These perivascular cells are in direct contact with the inner endothelium and help initiate a stable vessel network during angiogenic responses. Furthermore, this perivascular-endothelial cell interaction is necessary for restoring an appropriate microenvironment for tissue remodeling during skin regeneration [25].

INTRODUCTION

Burn wounds constitute 5%-10% of military casualties, involving 40%-60% of total body surface area (TBSA) [1–3], and require extensive reconstruction [4, 5]. Stem cells, a valuable tool in regenerative medicine, have demonstrated the capacity to help heal extensively injured tissues [6, 7]. Among adult mesenchymal stem cells (MSCs), adipose-derived stem cells (ASCs) have gained particular attention due to their ease of isolation, relative abundance [8–12], and multilineage differentiation potential [13–19]. Recent studies of various tissues, including adipose tissue, indicate that their resident stem cell niches localize to perivascular regions and express markers of both MSCs (CD105, STRO-1, and CD34) and pericytic (NG2 and platelet-derived growth factor receptor beta (PDGFR-$\beta^+$)/CD140b) lineages [20–24]. These perivascular cells are in direct contact with the inner endothelium and help initiate a stable vessel network during angiogenic responses. Furthermore, this perivascular-endothelial cell interaction is necessary for restoring an appropriate microenvironment for tissue remodeling during skin regeneration [25].

After a burn injury, the host immediately responds by coordinating molecular signals to initiate healing of the wounded area [26]. Adipose tissue in conjunction with the host immune system plays an important role in initiating the signaling events in the healing process [27–29]. ASCs have demonstrated their capacity to elicit a “regenerative” response through autocrine, paracrine, or direct cell-cell interactions [30–34]. Unfortunately, after a severe burn injury, the source of autologous stem cells from a patient is restricted due to limited clinical access of normal stem cell sources (e.g., subcutaneous liposapirate) and the lack of uninjured viable tissue, requiring an alternate source of stem cells for regenerative approaches.

Current standard of care for burn injuries involves surgical debridement of necrotic tissue associated with eschar, followed by the application of skin grafts or tissue-engineered skin substitutes using allogeneic and/or autologous cells [35, 36]. The process of wound debridement typically involves the removal of subcutaneous layers and associated tissue structures, including portions of intact hypodermal adipose tissue. Based on this observation, we hypothesize that the intact adipose tissue associated with the debrided skin could be a source of viable autologous stem cells for use in wound repair and skin regeneration.
**Title:** Debrided skin as a source of autologous stem cells for wound repair

**Authors:** Natesan S., Wrice N. L., Baer D. G., Christy R. J.

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### Abstract
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treatments. This study determined that (a) the hypodermal adipose region of the discarded human skin, after burn wound debridement, contains viable stem cells; (b) the cells appear to be of perivascular origin; and (c) the cells can be easily isolated and expanded in vitro and retain their ability to differentiate into multilineages. Furthermore, using an athymic rat excision wound model, we demonstrate that these cells can engraft into healing wound tissue.

**Figure 1.** Discarded human skin after burn wound debridement. (A): Gross morphology of debrided tissue. (B–E): Histological analysis of sections stained with Mason’s trichrome showing regions of (B): Necrotic dermis and hyalinized collagen (bold arrows). (C): Viable subcutaneous hypodermis regions with intact interlobular septae (asterisk) and other regions of completely necrotized tissue (serrated circle). (D): Thrombi were present at the dermal-hypodermal junction (serrated circle). (E): Other regions exhibited collapsed (open arrow) and patent blood vessels (closed arrow) in hypodermis. (F–I): Immunohistochemical analysis of microvessels in a discarded burn tissue stained with (F): Hoechst for nuclei. Primary antibodies for (G): CD31 (Alexa fluor 594) and (H): PDGFRβ (Alexa fluor 488). (I): Overlay of CD31, PDGFRβ, and Hoechst. Original magnification: ×40 (B), ×100 (C, D), ×200 (E), and ×400 (F–I). Abbreviations: *, viable subcutaneous hypodermis regions with intact interlobular septae; d, necrotic dermis; hd, hypodermis; hd(n), necrotic subcutaneous hypodermis; hd(v), viable subcutaneous hypodermis; PDGFRβ, platelet-derived growth factor beta.

**MATERIALS AND METHODS**

**Discarded Human Burned Skin**

Discarded human skin samples (Fig. 1A) from severely burned patients undergoing wound debridement were obtained from the Brooke Army Medical Center Burn Unit (Fort Sam Houston, TX). Samples were collected from 21 male burn patients with a mean age of 33 years. This study focused on the 20- to 30-year-old age group, which is the average age of military male burn patients [37]. From this age group, three samples were randomly selected for extensive analysis; and a sample from an older patient (age 49 years) was analyzed for age comparison (Table 1). This study was conducted under a protocol reviewed and approved by the U.S. Army Medical Research and Materiel Command Institutional Review Board and in accordance with the approved protocol (HSC20080290N). In this study, the authors were blinded from the patient information regarding TBSA and the status of the wound during surgical debridement. Before tissue processing, punch biopsies of approximately 5 mm in diameter were taken from various regions of the debrided sample and fixed with 10% neutral-buffered formalin for histological analysis or were cryopreserved for immunohistochemical analysis using the gradient sucrose cryopreservation technique [38].
Table 1. Debrided tissue samples

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age (years)</th>
<th>Percentage of tissue viability</th>
<th>dsASC yield floating fraction (cells per milliliter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BH1</td>
<td>24</td>
<td>45</td>
<td>1.85 x 10^4</td>
</tr>
<tr>
<td>BH2</td>
<td>26</td>
<td>75</td>
<td>2.48 x 10^4</td>
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<tr>
<td>BH3</td>
<td>40</td>
<td>40</td>
<td>1.5 x 10^4</td>
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<tr>
<td>BH4</td>
<td>24</td>
<td>80</td>
<td>2.2 x 10^4</td>
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Histological Analysis

Histological analysis was performed on 5–7 μm sections of formalin-fixed biopsies from the discarded human skin samples and the granulation tissue collected from the wound bed of athymic rats. A “blinded” trained pathologist analyzed Masson’s trichrome and/or H&E stained sections of the discarded human skin biopsies to assess tissue viability and burn depth and the overall wound healing pattern of the athymic rat tissue.

Isolation and Cell Culture

Discarded human skin samples were washed three times with Hank’s balanced saline solution (HBSS; Invitrogen, Carlsbad, CA, http://www.invitrogen.com/) to remove adherent blood clots. The hypodermal layer was dissected from the dermal region along with the areolar connective tissue, transferred to a petri dish, and then finely minced with scissors. The minced tissue was then suspended in HBSS (5 ml of minced tissue per 25 ml HBSS), centrifuged for 10 minutes at 500 x g, and the floating fraction carefully pooled into sterile flasks. For every 1 ml of minced floating tissue, 3,500 units of collagenase type II (Sigma-Aldrich, St. Louis, MO, http://www.sigmaaldrich.com/) was added and incubated in an orbital shaker (125 rpm) for 45–60 minutes at 37°C, the floating fraction carefully pooled into sterile flasks. For every 1 ml of minced floating tissue, 3,500 units of collagenase type II (Sigma-Aldrich, St. Louis, MO, http://www.sigmaaldrich.com/) was added and incubated in an orbital shaker (125 rpm) for 45–60 minutes at 37°C, the floating fraction carefully pooled into sterile flasks. For every 1 ml of minced floating tissue, 3,500 units of collagenase type II (Sigma-Aldrich, St. Louis, MO, http://www.sigmaaldrich.com/) was added and incubated in an orbital shaker (125 rpm) for 45–60 minutes at 37°C, the floating fraction carefully pooled into sterile flasks. For every 1 ml of minced floating tissue, 3,500 units of collagenase type II (Sigma-Aldrich, St. Louis, MO, http://www.sigmaaldrich.com/) was added and incubated in an orbital shaker (125 rpm) for 45–60 minutes at 37°C, the floating fraction carefully pooled into sterile flasks.

Cryosections of 10 μm from frozen biopsies were washed once with HBSS and fixed with 4% paraformaldehyde (PFA; EMS, Hatfield, PA, http://www.emsdiasum.com/) for 20 minutes at room temperature. After three washes with HBSS, nonspecific Fc receptor mediated sites were blocked by incubating the sections for 40 minutes with human Fc receptor blocking solution (Innovex Biosciences, Richmond, CA, http://www.innovx.com/), for 2 hours with 1% bovine serum albumin (BSA, Sigma) in HBSS, or 5% donkey serum (Sigma) for cytoskeletal markers. To localize cytosolic markers, sections were permeabilized with 0.01% Triton X-100 in HBSS and blocked for 2 hours with 1% BSA or 5% donkey serum. The human burn skin tissue sections were incubated with 10 μg/ml of anti-human monoclonal antibody directed against PDGFRβ (CD140b; BD Bioscience) and/or CD31 (R&D Systems, Minneapolis, MN, http://www.rndsystems.com/); tissue sections obtained on day 8 and 12 from rats treated with dsASC-Matrigel were incubated with 10 μg/ml human monoclonal antibody against mitochondrial protein antibody (Millipore, Billerica, MA, http://www.millipore.com/) and 5 μg/ml mouse specific antilamin monoclonal antibody (Millipore) to detect dsASCs and Matrigel within the wound bed, respectively. Tissue sections were incubated with primary antibodies overnight at 4°C. Primary antibodies were detected by incubating 5 μg/ml Alexafluor 488 and/or 555 (Invitrogen) tagged secondary antibodies of corresponding Ig class for 45 minutes at 4°C. Antibodies with product numbers are listed in Supporting Information Table 1.

Passage 1 (P1) dsASCs (BH1, BH2, and BH3) were cultured separately overnight in two-well chamber slides at 20,000 cells per well (Nalgene Nunc, LabTek chamber slide, Naperville, IL, http://www.nuncbrand.com/). The cells were washed twice with HBSS, fixed with 4% PFA for 20 minutes, and incubated overnight at 4°C with anti-human monoclonal antibodies targeted against transferrin receptor (CD71; 10 μg/ml R&D Systems), intercellular cell adhesion molecule-1 (CD54; 15 μg/ml R&D Systems), endoglin (CD105; 40 μg/ml R&D Systems), Thy-1 (CD90; 10 μg/ml BD Bioscience), and PDGFβR (50 μg/ml, BD Bioscience). The specificity of human-specific antimitochondrial protein antibody (Millipore) was assessed in vitro by labeling dsASCs cocultured with rat ascites (rASCs). rASCs were isolated by using a protocol described previously [12]. The tissue isolation was performed in compliance with the Animal Welfare Act and Animal Welfare Regulations and in accordance with the principles of the Guide for the Care and Use of Laboratory Animals. rASCs were cytoplasmically labeled with quantum dot (Qdot) nanocrystals 625 using Qracker cell labeling kit (Invitrogen) according to the manufacturer’s instructions. The rASCs and dsASCs were cultured in two-well chamber slides (20,000 cells per well), either alone or mixed in equal numbers (10,000 cells of each) for 48 hours. The cells were fixed with 4% PFA and immunolabeled with 5 μg/ml of monoclonal human-specific antimitochondrial protein antibody (Millipore). Primary antibody was detected using 5 μg/ml of Alexafluor 488 labeled secondary antibodies. After tissue sections and cells were processed for immunohistochemistry, their nuclei were stained with Hoechst 33,342 (10 μg/ml, Invitrogen) for 20 minutes at room temperature. Nonspecific fluorescence was determined by incubating cell preparations/tissue sections with isotype controls or solely with secondary labeled antibodies.

Flow Cytometry

P1 dsASCs from patients BH1–BH4 were washed twice with HBSS, trypsinized, and resuspended in fluorescence-activated cell sorting (FACS) cell staining buffer (Biologend, San Diego, CA, http://www.biologend.com/) to a final concentration of 5 x 10^5 cells per milliliter. The dsASCs were then incubated for 45 minutes (4°C) with fluorophore-labeled mouse anti-human antibodies (5–10 μg/ml; CD45-PE, CD90-PE, CD105-PE, and STRO-1-Alexa fluor 647 (Biologend); CD54-FTC and CD71-PE (R&D Systems); PDGFβR-PE (BD Bioscience); and CD19 and CD44 (R&D Systems). Experiments were controlled by incubating the dsASCs with their respective fluorophore-labeled isotype antibodies (10 μg/ml, R&D Systems). FACS analysis was performed with an FACS Aria flow cytometer (BD Bioscience). Data collected from P1 dsASC cultures were controlled by analyzing P0 cells from one preparation (BH2) to eliminate any biased results that may have occurred due to selective culture adherence of P1 cells. Commercially available ASCs from subcutaneous fat tissue (ZenBio Inc., Research Triangle Park, NC, http://www.zen-bio.com/) was used as a positive control. Before FACS analysis, forward and side scatter were adjusted for each sample using appropriate unstained cells to eliminate dead cells and cell debris. Autofluorescence signals were eliminated by adjusting the signal outputs from designated channels, and the sensitivity was adjusted to collect a gated population of cells. Total percentage
of cells staining positive for individual markers within the gated populations was determined. Results were quantitated by FACS Diva software (BD Biosciences). Approximately 20,000 events were collected and analyzed to determine the total population of cells staining positive for cell surface specific markers.

Adipogenic, Osteogenic, and Vasculogenic Differentiation of dsASCs

P1 dsASCs were cultured to confluence and induced to differentiate toward adipogenic and osteogenic lineages, using methods described previously [39]. Adipogenic cells were stained for neutral lipids using oil red O (Sigma) and osteogenic cells for calcium phosphate deposition with Alizarin red S (Sigma). Tube formation assays were performed using 1.5 ml of growth factor-reduced Matrigel (BD Bioscience) in six-well culture inserts (BD Falcon). P2 dsASCs (50,000 cells per well) were added on top of the three-dimensional Matrigel and cultured for 10 days using the Cascade 200 medium containing growth supplements (Invitrogen), with/without 50 ng/ml of vascular endothelial growth factor (VEGF; Peprotech, NJ, http://www.peprotech.com/) and 10 ng/ml of basic fibroblast growth factor (bFGF; Peprotech); and a phase-contrast photomicrograph was taken. After differentiation, RNA was isolated to analyze the cells’ ability to express adipogenic, osteogenic, and vasculogenic specific mRNA targets. Undifferentiated dsASCs were used as controls.

RNA Isolation and RT-PCR

Total RNA was isolated using Triozol LS reagent (Invitrogen) [40]. The concentration of RNA was determined at OD260nm, using a NanoDrop spectrometer (NanoDrop Technologies, Inc., Wilmington, DE, http://www.nanodrop.com/). cDNA was synthesized in duplicate samples using 150 ng of total RNA to begin first-strand synthesis (SuperScript III first-strand synthesis supermix with oligo-dT primers, Invitrogen). The following primer sets were used to evaluate the mRNA expression of specific markers for stem cells: CD31, CD90, CD105, PDGFRβ, and CD45 (negative control); adipogenic cells, insulin-regulated glucose transporter-4 (GLUT-4), adiponectin, peroxisome proliferator-activated receptor gamma (PPARγ), and leptin; and osteogenic cells, alkaline phosphatase (ALP), and runt-related transcription factor 2 (Runtx2). Master mixes with 200 nM forward/reverse primers and SYBR GreenER, qPCR supermix (Invitrogen), and each sample of synthesized cDNA template were analyzed by real-time polymerase chain reaction (RT-PCR) using a Bio-Rad CFX96 thermal cycler system (Bio-Rad, Hercules, CA, http://www.bio-rad.com/). Non-template control and no reverse transcriptase controls were run for each reaction. Gene expression was normalized to glyceraldehye-3-phosphate dehydrogenase, and runt-related transcription-4 (Runtx4) as an endogenous control. Expression levels for each mRNA transcript were normalized to glyceraldehye-3-phosphate dehydrogenase, and runt-related transcription-4 (Runtx4) as an endogenous control. The fold change in expression levels for each mRNA transcript was determined by the 2^-△△Ct method [41]. For the Matrigel (BD Bioscience) assays, RNA from five gels was pooled, and cDNA was synthesized as described. Expression levels of endothelial (CD31) and pericyte (NG2) specific genes were analyzed. The fold change in expression levels for stem cell specific genes from P1 dsASCs was normalized to the expression levels of control ASCs (ZenBio Inc.). The fold change in expression levels for stem cell specific genes from P2 to P7 dsASCs was normalized to the expression levels of the corresponding P1 dsASCs. The fold change in expression of adipogenic, osteogenic, and vasculogenic differentiation specific genes was normalized to the expression levels of undifferentiated P2 dsASCs. All primer sets were designed by, and are proprietary to, SA Biosciences (Frederick, MD, http://www.sabiosciences.com/).

Engraftment of dsASCs In Vivo

Male nmu nude athymic rats (175–250 g) were obtained from Harlan Laboratories (Indianapolis, IN, http://www.harlan.com/) and housed in the United States Army Institute of Surgical Research animal care facility. Rats were allowed access to water and chow ad libitum. Before the wound was created, a silicone splinting ring was stapled around the delineated wound margin; and a 1.4-cm-diameter full-thickness excision wound was created on the dorsum down to the panniculus of rat [42, 43]. Animals were randomly divided into three groups of five: saline control (250 µl), Matrigel (1 ml), and Matrigel (1 ml) + 100,000 dsASCs (P2) isolated from patient BH3. All the wounds were dressed with Tegaderm transparent film dressing (3M, St. Paul, MN, http://www.3m.com/) and observed for up to 12 days. The wound beds, including the healed area of skin surrounding the wound, were harvested on days 8 and 12 and samples prepared for histological and immunohistochemical analysis.

RESULTS

Viability of Discarded Human Skin

Histological examination of Masson’s trichrome and/or H&E stained biopsies of each debrided skin sample indicated that the debrided samples ranged from 40%–80% tissue viability (Table 1). Representative light micrographs of Masson’s trichrome-stained sections (Fig. 1) show complete loss of epidermis and reticular dermal regions (Fig. 1B). The remnant dermal tissue shows the presence of hyalinized collagen (bold arrows) with loss of individual collagen bundles and cellular necrosis. The hypodermal region consisted of intact adipocytes separated by intact interlobular septae and thermally collapsed areas with complete necrosis of both fat cells and dermis (Fig. 1C). Some regions of tissue exhibited a loss of vascular patency and the presence of thrombi in the dermal-hypodermal junction (Fig. 1D). Still other regions of the tissue contained completely intact blood vessels (Fig. 1E).

Immunofluorescence analysis of frozen tissue biopsies showed the presence of PDGFRβ+ cells (Fig. 1H) in the perivascular space of CD31+ microvascular structures (Fig. 1G) in all four samples studied. Figure 1I shows a representative immunofluorescence image of PDGFRβ+ cells specifically wrapping CD31+ cells, which is observed throughout the hypodermal region, regardless of the burn severity. However, most of the PDGFRβ+ cells appeared to be associated with intact microvessels. Although positively stained cells were observed in areas of collapsed lobular regions, a spatially distinct staining was not observed (data not shown).

Stem Cells from the Hypodermis of Debrided Skin

During cell isolation, hypodermis from all the debrided burn samples were separated into two fractions: a floating fraction, comprising most of the viable fat tissue, and a pellet fraction, composed of mostly extracellular components associated with some hypodermis and remnant dermal debris. The cells isolated from the floating fraction of debrided tissue were plated overnight, and all the adherent cells (dsASCs) were considered P0. A portion of the P0 cells were grown in chamber slides for 24–48 hours (P1; Fig. 2A) for morphological and immunocytochemical analysis. The cells exhibited characteristic fibroblast-like morphology and stained positive for stem cell-associated surface markers including CD54, CD71, CD90, CD105, STRO-1, and PDGFRβ (Fig. 2C). This demonstrates that the floating fraction of isolated cells is originated from the perivascular space of blood vessels surrounding the adipocytes. The P1 cells were negative for endothelial (CD31) and leukocytic (CD45) cell markers, indicating selective adhesion of stem cells during subsequent subculture (data not shown). The floating hypodermal fraction, after collagenase digestion, yielded between 1.5 × 10^5 and 2.5 × 10^5 stem cells per milliliter, shows that cells could still be harvested irrespective of the skin tissue viability (Table 1). When the
adherent cells obtained from the floating hypodermal fraction were subjected to serial passage, the dsASCs showed a four- to fivefold increase in cell number every 48 hours up to P4, after which the proliferation rate marginally decreased to three- to fourfold every 48 hours up to P8 (Fig. 2B). FACS analysis (Fig. 3A) of P1 cells exhibited forward and side scatter plots consistent with the cell isolate having minimal contaminating CD45* cells (1%–3% ± 0.25, n = 4), which demonstrated that the isolation technique eliminated leukocytic contamination. The gated R1 population was CD54*CD71*CD90*CD105*STRO-1* for each patient sample. These cells were also PDGFRβ* (BH1, 87.8%; BH2, 79.6%; BH3, 69.4%; and BH4, 80.4%), suggesting their perivascular origin (compare with Fig. 1I). Overall, the percentage expression of CD71, CD90, CD105, and PDGFRβ markers was consistently high in all the samples analyzed (Fig. 3B) and not related to the percentage viability of the debrided tissue. CD54*STRO-1* cell fractions were significantly lower for all the samples, when compared with CD54*CD71*CD90*CD105*PDGFRβ* panel. All patient tissues used in this study were obtained from combat-associated burns; therefore, the patient pool consisted of mainly

Figure 2. Immunophenotype characterization of isolated debrided skin adipose-derived stem cells (dsASCs). (A): Micrograph depicting a confluent monolayer of dsASCs at Passage 1 (P1) isolated from the remnant hypodermis of debrided skin. (B): Cumulative cell number of dsASCs serially passed with an initial seeding density of 6 × 10^3 cells per square centimeter surface area. (C): Micrographs of dsASCs at P1, immunolabeled for stem cell surface markers CD54, CD71, and CD90 (Alexa fluor 594), CD105, STRO-1, and PDGFRβ (Alexa fluor 488). Nuclei were stained with Hoechst’s reagent. Original magnification: ×200 (A) and ×400 (C).
young males. For this reason, positive control ASCs (Zenbio, Inc.) were isolated from subcutaneous fat of a younger, healthy adult male (age 32 years). In addition, dsASCs isolated from a 49-year-old patient was used for age comparison. Further investigation into the factors responsible for the age-related percentage change in number of stem cells for the panel of CD54^+ CD71^+ CD90^+ CD105^+ PDGFR^b^ markers is warranted.

**Figure 3.** Fluorescence-activated cell sorting (FACS) analysis of debrided skin adipose-derived stem cells (dsASCs). The dsASC isolates from four different donors (BH1, BH2, BH3, and BH4) and control ASCs derived from a 32-year old healthy individual were analyzed for stem cell marker expression. (A): The top panel depicts scatter plots of Passage 1 (P1) dsASCs and FACS histograms for labeled antibodies specific for stem cell surface markers: CD45, CD71, CD90, CD105, PDGFR^b^ (phycoerythrin), CD54 (flourescein isothyocynate), and STRO-1 (Alexa flour 647-APC). (B): Comparative bar graph of the percentage (±SD) gated population of dsASCs positive for CD54, CD71, CD90, CD105, PDGFR^b^, and STRO-1 in all four donor isolates and control ASCs. (C): FACS scatter plots showing the overall distribution of P0 and P1 dsASCs. (D): Bar graph of CD54, CD71, CD90, CD105, PDGFR^b^, and STRO-1 positive cells that were present within the total gated population of P0 and P1 dsASCs. For all FACS analysis, 20,000 events were collected (n = 2, percentage population ± SD) and analyzed for the total population positive for specific cell surface markers. The red line plot in all the FACS histograms represents isotype-matched antibody controls. Abbreviations: APC, allophycocyanin; FITC, flourescein isothyocynate; FSC, forward scatter; PE, phycoerythrin; PDGFR^b^, platelet-derived growth factor beta; SSC, side scatter.
dsASCs Maintain Stemness

To preliminarily assess the ability of the dsASCs to maintain their stemness after passaging, P0 and P1 cells were compared by FACS analysis for stem cell marker expression (Fig. 3C). A twofold higher percentage of CD54⁺CD71⁺CD90⁺CD105⁺PDGFRβ⁺ cells was present within the total gated population of P1 cells (Fig. 3D). P0 cells had a noticeable number of side-scattered events due to a mixture of leukocytes (6.2%), which were significantly low in P1 cultures (<1%), suggesting that after plating CD45⁺ cells are substantially eliminated. Further, the P1 cells were CD19⁻CD44⁺ (Supporting Information Fig. S1A–S1C) [44], providing evidence for preferential expansion of stem cells. When expanded in culture, a high percentage of dsASCs was PDGFRβ⁺ (Supporting Information Fig. S1D, S1E), suggesting that the cells originated from the perivascular space, while maintaining their stem cell characteristics.

For a more extensive analysis of dsASCs to maintain their stemness, gene expression levels of stem cell markers were studied using RT-PCR. Relative expression levels of CD71⁺CD90⁺CD105⁺PDGFRβ⁺ panel were investigated in dsASCs at different passages (P1–P7) for all the debrided tissue samples (Fig. 4A–4C). P1 cells exhibited little to no change in the expression levels for all the markers (less than 0.2-fold) regardless of age or percentage of tissue viability. In contrast, there was an apparent increase in the expression levels of CD71⁺CD90⁺CD105⁺PDGFRβ⁺ transcripts, with respect to tissue samples and passage number (P2–P7). dsASCs from BH1 and BH2, with 45% and 40% skin tissue viability, had variable gene expression levels during early passage (P2 and P3); whereas later-passage cells (P4–P7) showed fairly consistent levels of expression. In comparison, cells from BH2, which had 80% viability, showed consistent levels of elevated expression of stem cell markers throughout analysis.

P2 dsASCs from BH1, BH2, and BH3 also exhibited multilineage differentiation potential. When induced with adipogenic media, cytoplasmic accumulation of lipids began by...
day 6, and by day 14, when stained with oil red O, the cells showed intense red staining for lipids (Fig. 4G). Upon differentiation, the expression level of PPARγ, an early transcription promoting receptor factor [45], significantly increased in all samples (20–40-fold) and is similar to those observed in control ASCs (e.g., BH2, 25.6-fold, and control ASCs, 23.1-fold). Furthermore, the increase in adiponectin transcript levels (approximately 1,000-fold) demonstrates the ability of dsASCs to undergo adipogenesis through the activation of a glucose-regulated signaling pathway [46]. In addition, expression of GLUT-4 and leptin from all patient samples was significantly higher relative to noninduced dsASCs (Fig. 4D).

When dsASCs were differentiated into an osteogenic lineage, all samples showed increased expression levels of ALP, indicative of stromal cell commitment to osteogenesis [39], and showed comparable levels of expression as control ASCs. Matrix mineralization appeared by day 8 postinduction and increased over time, with the cells positively staining for calcium by day 21 (Fig. 4H). Terminal differentiation was confirmed by the expression of Runx2, a master osteogenic transcription factor responsible for mature osteoblast formation (Fig. 4E). Induced cells from all the patient samples and control ASCs expressed Runx2 by 21 days with subtle differences in each expression level. Still, faster osteogenic induction was observed in cells isolated from higher viable tissue (BH2) in comparison to those derived from less viable tissue (BH1 and BH3). Expression of Runx2 significantly increased by 28 days (approximately 70-fold) along with ALP (approximately sevenfold), indicating the commitment of dsASCs toward the osteogenic lineage (data not shown).

The ability of dsASCs to differentiate toward a vascular phenotype was also investigated. P2 dsASCs were seeded over a three-dimensional matrix of Matrigel and induced with endothelial cell-specific media. The dsASCs spontaneously joined, forming tubular structures covering the entire surface of the Matrigel, with a dense cellular network forming within 10 days (Fig. 4I). When analyzed by RT-PCR, the differentiated dsASCs from BH1, BH2, and BH3 exhibited a significant increase in expression of CD31 (Fig. 4F) when compared with undifferentiated cells. Furthermore, the expression levels of PDGFRβ did not change, revealing that under endogenic induction conditions, dsASCs differentiate into endothelial-like cells rather than pericytic cells. The differentiated cells from all the samples, regardless of percentage viability, show negligible increases in expression levels of the pericyte specific marker NG2 (less than 0.1-fold, data not shown). These results are comparable to recent studies showing that ASCs seeded on Matrigel are capable of forming capillary tube-like networks in vitro but are unstable over time [23, 25].

**In Vivo Engraftment of dsASCs**

Engraftment efficiency of dsASCs (P2; BH2) was assessed using an excision wound splinting model [42] in immunodeficient rats (Fig. 5A). After 8 and 12 days postgrafting of dsASC-Matrigel composites, the wound bed was photographed (Fig. 5B, 5C) and harvested for analysis using routine histological examination. The dsASC-Matrigel composite integrated within the healing wound bed as depicted within the serrated black circles in Figure 5D, 5F, which corresponds to days 8 and 12, respectively. This was further confirmed...
through positive staining of the wound bed for laminin (a major component of Matrigel) observed on day 8 (Supporting Information Fig. S2A–S2C) and on day 12 (Supporting Information Fig. S2D–S2F).

On day 8, the wound-composite region was rich in cells and lacked collagen bundles. Re-epithelialization was evident at the epidermal region (cells above the dotted white line in Fig. 5D) and loosely associated with the acellular amorphous matrix (Matrigel). The dermis showed the presence of multifocal amorphous matrices with organized fibrous granulation tissue along with spindle-shaped cells and nascent collagen (Fig. 5E), eventually turning into a mature vascularized wound.

**Figure 6.** Tracking of human debrided skin adipose-derived stem cells (dsASCs) in vivo using a rat model of skin wound healing. To qualify the specificity of the human-specific anti-mitochondrial protein antibody, in vitro coculture experiments were performed using mixtures of rat ASCs (rASCs) and dsASCs. Immunofluorescence labeling of pure human dsASCs with the anti-human mitochondrial antibody (A) (green), and cultures of pure rASCs labeled with Q-dots (B) (red) were also subjected to the anti-human mitochondrial antibody (B) (negative). (C): Cocultures of an equal number of dsASCs and Qdot labeled rASCs immunostained with anti-human specific mitochondrial protein antibody demonstrates that the antibody specifically detects dsASCs and not rASCs. To examine the behavior of dsASCs in vivo in a rat model, wound beds were treated with a mixture of Passage 2 (P2) dsASCs in Matrigel (100,000 per wound) and tracked by staining with human-specific anti-mitochondrial protein antibody (F, G, K) (red), and total cells tracked by Hoechst stain (rat host and human dsASCs) in the wound bed (D, E, J) (blue). To simultaneously view both total cells and transplanted dsASCs in the same region, photomicrographs were superimposed as follows: (D, F) = (H), (E, G) = (I), and (J, K) = (L). (D–I) Day 8 images; (J–L) day 12 images. Human cells were able to be detected in the wound bed (white arrows in (H) and (I)) and at the re-epithelializing surface (white arrow heads in (H)), surrounded by an acellular region (white serrated circle in (H)) on day 8. Higher magnification shows the dermal matrix populated with human dsASCs (I) (inset). Panel (L) inset shows multifocal punctate cytoplasmic staining of human mitochondrial staining observed on day 12. Serrated white line on panel (D), (F), and (H) shows the region of re-epithelialization (above the line). Original magnification: ×200 (D, F, H), ×600 (E, G, I), and ×900 (J–L).
dsASCs from all samples were able to maintain stemness as multiligneage differentiation capabilities [44, 50]. Furthermore, PDGFR indicate that the debrided skin biopsies possess both NG2, and 3G5) cell-specific markers [48, 49]. Our results perivascular space and are immunopositive for stromal/C2 bridement. The dissected hypodermal layer is a practical stem cells derived from discarded burn tissue after wound debridement. The viable cells isolated from debrided tissue and those observed were mostly localized in the wound bed, and those observed were mostly confined to the dermal region with multifocal punctate cytoplasmic fluorescence (Fig. 6L and inset).

**DISCUSSION**

This study provides evidence for a new source of autologous stem cells derived from discarded burn tissue after wound debridement. The dissected hypodermal layer is a practical source of autologous ASCs (1.5 × 10^5 and 2.5 × 10^5 stem cells per milliliter), which could be used for developing burn wound therapies. The viable cells isolated from debrided tissue were identified as stem cells (dsASCs) and have a similar panel of cell surface markers as ASCs isolated from a normal subcutaneous lipoaspirate [8]. Previous studies have shown that the cells comprising capillary vessels in adipose tissue exhibit spatially separated expression of CD31 and PDGFR/β markers [20]. These PDGFR/β cells have been identified as the point of origin for isolated ASCs. These ASCs occupy the perivascular space and are immunopositive for stromal (CD44, CD90, CD105, and STRO-1) and pericytic (PDGFR/β, NG2, and 3G5) cell-specific markers [48, 49]. Our results indicate that the debrided skin biopsies possess both PDGFR/β and CD31+ cells and that isolated dsASCs (P1) are a relatively homogenous cell population when cultured in vitro. These dsASCs consistently express CD54+CD71+CD90+CD105+PDGFR/β markers in high percentages (>65%) regardless of tissue viability, with discernible variation in CD54 and STRO-1 expression. This observation is analogous to other studies, where culture expansion of clonally selected STRO-1+CD31+CD45+ cells increases their multiligneage differentiation capabilities [44, 50]. Furthermore, dsASCs from all samples were able to maintain stemness as demonstrated by their gene expression profile for stem cells markers when P1 dsASCs were compared with P1 ASCs derived from a healthy individual. The expression of these markers remained consistent across P1–P7 passages with little to no variability resulting from age difference of donors. This finding illustrates that the stemness of the isolated dsASCs remains unaffected after thermal injury to the surrounding tissue. Cells isolated from the youngest donor (BH1: approximately 45% tissue viability), when compared with older donors, exhibited higher expression levels of panel markers during initial passages than in later passages. These results imply that the expandability and the ability of dsASCs to maintain their stemness are directly dependent on patient age, an observation corroborated by other studies [51]. Similarly, of the various markers tested, CD71 has been shown to have a one- to twofold higher expression level in proliferating cells [52, 53]; this observation is dependent on passage number, patient age, and viability of debrided skin tissue. In our dsASCs cultures, relative expression levels of CD71 could be considered as an index to characterize the ability of culture-expanded dsASCs to maintain their stemness between donors. However, any effect that age may have on dsASCs stemness properties requires further investigation due to the lack of older patient burn samples in this study.

Furthermore, dsASCs exhibited adipogenic, osteogenic, and endothelial differentiation potential, confirming their stem cell-like phenotype. Regardless of age or viability of skin, dsASCs displayed adipogenesis through the classical pathway of transcription activation of adipogenic-specific genes involved in early (PPARγ) and terminal (leptin and adiponectin) stages of differentiation [45]. Increases in leptin, GLUT-4, and adiponectin transcript levels between days 14 and 21 were consistent with ASCs differentiation isolated from healthy individuals. When dsASCs were induced with osteogenic media, calcium deposition and increased levels of ALP and Runx2 were observed, demonstrating commitment of dsASCs to undergo osteogenesis [54, 55]. It is noteworthy that Jackson et al. [56] and Nesti et al. [57] recently showed that mesenchymal progenitor cells isolated from traumatized muscle expressed similar sets of stem cell markers (expressed by dsASCs) in uninjured muscle tissue. Finally, we also show that dsASCs can be induced toward an endothelial phenotype when seeded on Matrigel, a characteristic feature of MSCs of pericytic origin [23]. This differentiation potential of dsASCs could be stimulated with exogenous mitogens, such as vascular endothelial growth factor and basic fibroblast growth factor, further confirming their ability to be induced toward an endothelial phenotype (data not shown). In this study, dsASCs were able to form tube-like networks across the three-dimensional Matrigel but became unstable after several days. This phenomenon demands further investigation to determine the role of dsASCs in forming both endothelial cells and pericytes during neovascularization both in vitro and in vivo.

When the dsASC-Matrigel mixture was applied to an excision wound in athymic rats, they engrafted into the granulating wound bed after 8 days, with only a few cells identified on day 12. The embedded dsASCs may be contributing significantly to the wound healing and skin regeneration process in this model, which complements the recent finding that ASCs do contribute to healing of chronic wounds [58]. Overall, the data reported in this study supports the hypothesis that dsASCs can be a source of stem cells for skin regenerative therapies, especially in patients with burn wounds involving large TBSA. In these patients, accessibility of autologous MSCs from normal tissue sources (e.g., bone marrow and lipoaspirates) is not feasible clinically. Therefore, in such cases, the hypodermal layer of discarded burn tissue may be an invaluable source of stem cells for wound repair and skin regeneration.
This study provides evidence that stem cells in the hypodermis are preserved during and after severe thermal injury. These autologous stem cells can be isolated from the debrided skin that is surgically removed from burn-injured patients and in quantities that could be useful clinically for burn repair and regeneration. Considering that large TBSA burn wounds require extensive debridement, this waste tissue represents an invaluable stem cell source. Promising results from both in vitro and in vivo perspectives support the use of deASCs alone or in combination with appropriate bioscaffolds as a useful therapeutic tool in the field of regenerative medicine.

We thank Cpt. Laura McGhee and Thomas Garza for collection of debrided tissue; Col. Seung Kim and Lt. Scot J.

The authors indicate no potential conflicts of interest.