Translational products of adipose tissue-derived mesenchymal stem cells: Bench to bedside applications

Shilpa Sharma, Sathish Muthu, Madhan Jeyaraman, Rajni Ranjan, Saurabh Kumar Jha

ORCID number: Shilpa Sharma 0000-0001-8695-8372; Sathish Muthu 0000-0002-7143-4354; Madhan Jeyaraman 0000-0002-9045-9403; Rajni Ranjan 0000-0003-2324-6970; Saurabh Kumar Jha 0000-0002-7437-0755.

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Abstract

With developments in the field of tissue engineering and regenerative medicine, the use of biological products for the treatment of various disorders has come into the limelight among researchers and clinicians. Among all the available biological tissues, research and exploration of adipose tissue have become more robust. Adipose tissue engineering aims to develop by-products and their substitutes for their regenerative and immunomodulatory potential. The use of biodegradable scaffolds along with adipose tissue products has a major role in cellular growth, proliferation, and differentiation. Adipose tissue, apart from being the powerhouse of energy storage, also functions as the largest endocrine organ, with the release of various adipokines. The progenitor cells among the heterogeneous population in the adipose tissue are of paramount importance as they determine the capacity of regeneration of these tissues. The results of adipose-derived stem-cell assisted fat grafting to provide numerous growth factors and adipokines that improve vasculogenesis, fat graft integration, and survival within the recipient tissue and promote the regeneration of tissue are promising. Adipose tissue gives rise to various by-products upon processing. This article highlights the
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significance and the usage of various adipose tissue by-products, their individual characteristics, and their clinical applications.

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Core Tip: Promising evidence supports clinical application of derivatives of adipose-derived stem cells enriched with numerous growth factors and adipokines for improved vasculogenesis, graft integration, and survival within the recipient tissue. Analysis of its differential characteristics and practical applications became a necessity. In this review, we highlight the significance and usage of various adipose tissue by-products, their individual characteristics, and their clinical applications along with the evidence supporting its use.

INTRODUCTION

With developments in the field of tissue engineering and regenerative medicine, the use of biological products for the treatment of various disorders has come into the limelight among researchers and clinicians. Among all the biological products, keen interest has been shown in adipose tissue and its by-products for translation from bench to clinical applications, with due consideration to its various unique properties. Because of the heterogeneous cellular population of adipose tissue, adipose tissue-derived products possess a greater advantage of regeneration compared with bone marrow-derived products[1,2].

Traditionally adipose tissue or fat graft transfer was practiced for elective cosmetic procedures and plastic and reconstructive surgery[3]. Because of the regeneration potential possessed by adipose tissue, use has become widespread in cosmetic-dermatological procedures, facial rejuvenation, breast and buttocks augmentation, and genital aesthetics[4-6]. Cellular viability and survival at the transplanted site depends on various factors involving the recipient site, donor site, laboratory processing, and manipulation[7,8].

Adipose tissue engineering aims to develop by-products and their substitutes for regeneration and restoring function[9]. That further eliminates the need for organ transplants and mechanical device placement. The use of biodegradable scaffolds along with adipose tissue products has a major role in cellular growth, proliferation, and differentiation[10,11]. Once implanted at an appropriate site, scaffolds degrade, progenitor cells proliferate with the help of growth factors and cytokines to form new tissue. Preclinical studies have investigated the ability of biomolecules and biodegradable three-dimensional scaffolds to interact with adipose tissue products to promote the adipogenesis of stem cells in vitro[12-15]. In this review, we discuss the differential characteristics of adipose tissue derivatives and elucidate their applications in clinical scenarios.

ADIPOSE TISSUE BIOLOGY

An adipocyte is usually 50-150 mm in diameter, it dies from ischemia if it grows larger, and its life expectancy varies from few months to 10 years in humans[16]. Adipose tissue, a specialized connective tissue with lipid-rich adipocytes, it contains a heterogeneous population of cells but adipocytes represents only 20% or less of the cellular mixture[17]. Based on adipocyte morphology, the two types of adipose tissue are
white adipose tissue found in adults and brown adipose tissue found in newborns [18]. Adipose tissue, apart from being the powerhouse of energy storage, also functions as the largest endocrine organ, with the release of various adipokines. Adipose tissue-derived adipokines include leptin, adiponectin, apelin, chemerin, interleukin (IL)-6, 8, and 10; monocyte chemoattractant protein (MCP)-1; plasminogen activator inhibitor (PAI)-1, retinol binding protein (RBP)-4, tumor necrosis factor (TNF)-α, progranulin, complement C1q tumor necrosis factor-related protein (CTRP)-4, interferon (IFN)-γ, and interferon-γ-inducible protein (IP)-10, which are readily available sources to induce stem cells [19-22]. The adipokines work in a paracrine fashion when transplanted as a cellular therapeutic tool [23,24]. The components of adipose tissue are lipid-laden adipocytes, fibroblasts, neural and vascular progenitor cells, multipotent progenitor cells, pericytes, extracellular matrices, cytokines, growth factors, and immune cells such as CD4+ T cells, as shown in Figure 1. The progenitor cells in adipose tissue are of paramount importance, as they have the capacity for regeneration of these tissues. The progenitor cells differentiate into mesodermal, ectodermal, and endodermal lineages [25-27]. Tissue engineering experts focus on adipose tissue and its products for their plasticity, relative ease of harvest, and potential autologous usage. Stem cells and progenitor cells from freshly prepared SVFs constitute up to 3%, which represents 2500-fold more than the stem cells isolated from bone marrow source (0.002%) [28].

The literature on adipose tissue survival and regeneration depicts “cell survival theory” and “cell replacement theory” [29-32]. Various studies have shown the promising results of adipose-derived stem cell (ADSC)-assisted fat grafting, which provides numerous growth factors and adipokines for improved vasculogenesis, fat graft integration, and survival within the recipient tissue [8,33]. Adipocytes contain three zones namely, (1) the outer surviving layer (100–300 microns), (2) middle regenerating layer (600–1200 microns), and (3) inner necrotic layer, as shown in Figure 2 [30-32,34].

Yoshimura et al. [35] demonstrated the sequence of changes that happen after grafting or transplantation of adipose tissue. In preclinical studies, it is shown that all adipocytes undergo apoptosis in the initial few days of grafting. Activation of adipogenesis was by adipose-derived progenitor cells, which were augmented by the adipokines at 3 mo after grafting. By the end of 9 mo, lipid droplets were absorbed by macrophages. The final fat graft retention at the recipient site was determined by the rate of successful replacement of the adipocytes [35]. ADSCs possess various advantages over bone marrow-derived mesenchymal stem cells (BM-MSCs) [36-38]. Harvesting adipose tissue by liposuction is less painful than bone marrow aspiration [39,40]. The quantity of stem/stromal cells obtained from adipose tissue than that obtained from bone marrow [41]. In long-term culture, ADSCs are more genetically stable [42,43].

**Characterization of ADSCs**

Researchers demonstrated that ADSCs have a consistent phenotyp and reproductive capacity, based on cellular yield, cellular viability, adipocyte differentiation, and cell surface markers [44-46]. During initial culture, ADSCs are polygonal cells that adhere to the flask surface. They exhibit fibroblastic plastic morphology and expand in in vitro cultures. within 2 d of primary culture, 90% of cells become confluent when subcultured within 2 d of primary culture with a demonstrated ADSC yield of 87% and viability of 94% [47]. Luna et al. [47] recovered 1 × 10⁶ adipocytes, 1 × 10⁶ ADSCs, 1 × 10⁶ vascular endothelial cells, and 1 × 10⁶ other cells from 1 g of adipose tissue.

During in vitro culture, the ADSCs immunophenotype changes from CD34+, major histocompatibility complex (MHC) I and II molecules, CD80, CD86, CD45, CD11a, CD14, CD117, human lymphocyte antigen (HLA)-DR+, NOG+, undifferentiated embryonic cell transcription factor (UTF)1, WNT6, and WNT8A to increased expression of CD9, CD13, CD29, CD44, CD63, CD73, CD90, CD105, CD166, bone morphogenetic protein receptor (BMPR)2, collagen type VI alpha 2 chain (COL6A2), transforming growth factor (TGF)-βR1, and vascular endothelial growth factor (VEGF)-A [48-52]. The lack of expression of HLA class I and II molecules in coculture and serial passages, confers ADSCs the property of immunosuppression and make them suitable for allogenic transplantation [53-55].

MSCs from adipose tissue have strong positive expression of STRO-1, CD29, CD73, CD90, CD105, CD166 & CD44 and weak positive expression for CD34 and CD45 [56-57]. MSCs possess enhanced angiogenesis associated with the increased expression of CD105 and CD34. Proliferation and differentiation of MSCs is enhanced by CD9, CD29, CD44, CD49d, and CD106 [58-61]. Grontos et al. [62] demonstrated that adipose tissue-derived stroma-vascular fraction (SVF) support...
hematopoiesis in vitro. After about 8–12 cellular doublings in culture, ADSCs express CD34+ [63]. Various theories are available for the attribution of stem cell properties to pericytes. Szöke et al [64] stated that pericytes are present in both MSCs and ADSCs whereas Traktuev et al [65] and Crisan et al [66] stated that CD34+ and CD34- pericytes are identical to adipose-derived stem cells.

**Immunomodulation of ADSCs**

The cellular components of ADSC induce and activate quiescent native MSCs to secrete biological micromolecules at the site of injury to establish local homeostasis by increasing the permeability of cells at the injury site, downregulating inflammatory processes, and recognition of host-tissue progenitor cells for final differentiation into the cells of interest in the injured tissue [67]. Therefore, ADSCs activate adaptive cellular responses and secrete IL-1Ra, IL-1β, PGE2, IDO, IL-4 & -10, and TGF-β that modulate and prime the native immune cells. The micromolecular interactions lead to a cascade of events responsible for immunotolerance of engraftment to a foreign site.

The immunotolerant/immunosuppressive/immunomodulatory activity of ADSCs is caused by the interplay of regulatory T cells, cytotoxic T cells, and B cells as shown in Figure 3. Immunomodulatory activity includes the inhibition of INF-γ and TNF-α production by effector T lymphocytic cells, upregulation of IL-4 and IL-10 by native immune cells, and inhibition of proliferation, migration, and differentiation of B cells that produce immunoglobulins [68]. The activity of natural killer cells is suppressed by the production of indolamine 2,3-dioxygenase by MSC-like cells [69,70]. Dendritic cells enhance the expression of IL-4 IL-10 and suppressed INF-γ and TNF-α [71,72]. The cascade of events directly suppresses tissue inhibitors of matrix metalloproteinases (TIMPs) and matrix metalloproteinases (MMPs), resulting in the conversion of the proinflammatory environment to an anti-inflammatory environment. Adipose-derived MSCs secrete extracellular vesicles such as exosomes as a channel of communication with neighboring cells [73]. They act not only via direct cell-to-cell interaction but also via paracrine mechanisms [74] and are key mediators of signaling molecules such as TGF-β1 [75], IL-10 [76], PGE2 [77], NO [78] and IDO [79,80]. The bioactive factors in-turn help to promote tissue regeneration and repair at the site of action.
Derivatives and applications of adipose tissue

The various derivatives of adipose tissue that are of practical utility, with therapeutic potential are shown in Figure 4 and are discussed below:

**Adipose stem cells:** ADSCs are one of the forms of MSCs of adipose origin with an inherent property of self-renewal and multipotent differentiation[73,81]. Advantages of easy accessibility of the source, abundant availability, and fewer ethical concerns than embryonic stem cells, render ADSCs more suitable for use in regenerative medicine and tissue engineering. Tissue-engineered 3D scaffolds with ADSCs and biomolecules such as growth factors and extracellular matrix materials play a robust role in treating various disorders by cellular proliferation and differentiation[4]. Studies found that ADSCs have more grounded immunomodulatory impact than BM-MSCs[82-84]. Adipose tissue contains stem and progenitor cells in amounts of up to 3% of the uncultured SVF, which is 2500 times more than the stem cells obtained from bone marrow[85,86]. Depending on the donor and tissue harvesting site, lipoaspirates yield 1% to 10% stem cells, which is approximately 0.5 × 10^4 to 2 × 10^5 MSC/g adipose tissue[87]. ADSCs act in paracrine fashion by releasing adipocytokines, cytokines, and growth factors that form a secretome[88].

Following liposuction, adipose tissue is washed in phosphate-buffered saline containing 5% antibiotic, followed by tissue digestion by collagenase-1[89]. After the disintegration of adipose tissue, the resultant sample is transferred to tubes and centrifuged at 2000 rpm for 5 min to obtain a SVF that contains the ADSCs. The resultant cellular pellet is resuspended in 1 mL lysis buffer, recentrifuged at 2000 rpm for 5 min and the resulting cellular suspension is passed through a 70 μm cell strainer. The cellular mixture is transferred to lysine-coated culture plates and incubated at 37 °C with 5% CO_2_. To obtain ADSCs, about 500 mg of the adipose tissue cell suspension is inoculated in the wells[87,90]. According to International Fat Applied Technology Society, uncultured ADSCs express CD34^+, CD45^−, CD235a^− and CD31^−[63,91], whereas cultured ADSCs express CD73^+, CD90^+, CD105^+, CD44^+, CD45^− and CD31^−[91,92].

The short- and long-term storage of ADSCs have been investigated. The cellular proliferative capacity of ADSCs decreases with the length of storage[43]. Hence, they must be supplemented with 10% human serum or platelet-rich plasma (PRP) in 0.9% saline at 4 °C for the first 2 h and not more than 4 h[1,94]. For long-term storage, ADSCs can be stored at −80 °C in liquid nitrogen for up to 6 mo[95,96]. Certain strategies can be followed to enhance and optimize the potential of the ADSC, such as culturing them in hypoxic conditions, which not only enhances the immunomodulatory effect but also reduces the risk of chromosomal aberration and tumorigenesis[97]. Further, the cells can be cryopreserved in the early passages for future usage.

Figure 3 Immunomodulatory effects of mesenchymal stem cells via T and B lymphocyte system.
Figure 4 Derivatives of the adipose tissue complex.

ADSCs are used in cardiac tissue engineering where they enhance regeneration of myocardial tissues, improve left ventricular ejection fraction, and reduce the scar volume in ventricular wall in rodent models[99]. ADSCs are seeded as bioscaffold in the healing of cutaneous ulcers and soft tissue injuries[100,101]. Several studies have proven the beneficial role of ADSCs in multiple sclerosis, diabetes mellitus, and rheumatoid arthritis. The anti-inflammatory and immunomodulatory effects of ASCs have been demonstrated in various preclinical models of autoimmune diseases[102]. ADSCs have regenerative potential for bone and cartilage healing in addition to electrical stimulation of cells and tissues[103]. Various researchers across the globe have been working on the therapeutic efficacy of ADSCs in osteoarthritis knees and hips[104-106]. Agostini et al[107] developed a protocol for ADSC application in osteoarthritis in terms of isolation, dose, frequency, analysis, and follow-up. There is no consensus on the use of cultured vs uncultured cells in the management of osteoarthritis[106,108]. ADSCs have been used in sports injuries of ligaments and tendons[109,110], but evidence to support the safety and efficacy is lacking.

The potency of the stem cells to differentiate into various lineages including nerve cells and nervous tissue makes them a good candidate for use in neurological disorders. Cultured and uncultured cells have been used in the treatment of various neurological diseases such as Alzheimer’s disease, Parkinson’s disease, intervertebral disc, amyotrophic lateral sclerosis, multiple system atrophy, post-polio residual paralysis and traumatic brain injury[111-114]. The cells enhanced neurovasculogenesis, counterfeit fibrosis, oxidative stress, anti-inflammation, and neuromodulation. In neurological diseases, the research toward the use of ADSCs is ongoing in animal models.

**Microfat:** Compared with nanofat, microfat appears deep yellow, with a fine granular structure and intact three-dimensional adipose tissue architecture[115]. Microfat is composed of mature adipocytes, SVF cells, pericytes, capillary fragments, and fibrous scaffolds are well preserved[116-118]. Hence, they are a natural recipient site for survival of grafts, and provide a niche for SVF cellular mixtures for tissue regeneration and rejuvenation. Yang et al[117] harvested microfat from adipose tissue with cannulas that had side holes of less than 1 mm. Examination of the harvests revealed intact fat lobular structures without need of any inter-syringe passages of emulsification procedures. Side holes of < 0.8 mm allowed the processed microfat to easily pass through the needles without any blockage. Such harvests, used for skin and facial rejuvenation through 27 gauge needles, revealed the preservation of micro-functional units of fat tissue that retained the regenerative properties of ADSCs[117]. Caggiati et al[119] demonstrated a higher yield of ADSCs from lipoaspirates harvested by barbed compared with blunt cannulas, but they found that barbed cannulas cut the fibrous septum and produced higher quantities of coarse fibers that increased the probability of molecular blockage in the needle. The cellular components in the microfat induced tissue regeneration through the recruitment of monocyte/macrophage differentiation at the recipient site[35,120]. The cellular yield in microfat (2.28 ± 1.90) × 10⁵ cells/mL is higher than the yield in nanofat (4.12 ± 1.37) × 10⁴ cells/mL, indicating that nanofat involves mechanical emulsification[117]. Microfat components integrate into the local environment because of the intact microintegrity of cellular transplant, with inherent thereby making them available for future use[98].
resistance to attack by host immune-mediated cells[121].

Microfat possess various advantages. (1) No mechanical emulsification or centrifugation is involved in preparation, other than the shearing forces during harvesting. (2) Intact vascular fragments with viable cellular components are present in the mixture. (3) There is less ischemic exposure time during adipose tissue harvest and while preparing the microfat graft. (4) Preservation of viable adipocytes that restore the degenerated and atrophied skin and subcutaneous tissues[117]. Compared with other adipose derivatives, microfat retains the three-dimensional architecture of the native tissue essential for the survival of mature adipocytes, thereby providing a natural niche for ADSC cells to ensure optimal tissue regeneration. Microfat is used in esthetic procedures. Because the three-dimensional architecture is retained in microfat, it is used as a lipofiller in breast reconstruction and facial and gluteal augmentation[122-125]. Microfat is also being tried in scar revision, burn injuries, lipodystrophies, rhinoplasty and wrinkles[3,40,117,126,127].

Nanofat: In 2013, Tonnard et al[128] developed nanofat, which is an ultrapurified adipose tissue-derived product that is devoid of mature adipocytes but contains CD34+ ADSCs, microvascular fragments, growth factors, biological peptides, and cytokines[93,94]. It is a liquefied, autologous injectable product with the property of biological integration with adjacent cells and tissues because of its homogenous consistency[129]. The size of nanofat components is approximately 400 to 600 μm[130]. Nanofat behaves much like adipose tissue-derived mesenchymal stromal cells. At the site of injury, stromal cells initiate a site-specific reparative response comprised of remodeling the extracellular matrix (ECM), enhanced and sustained angiogenesis, and immune system modulation. Because of the multi- and pluripotent nature of the cellular components in nanofat, it possesses the ability to differentiate into multiple lineages. Hence, nanofat can be used in preclinical and translational research in tissue engineering. Various preclinical and clinical studies have demonstrated antiﬁbrotic, proangiogenic, neuroregenerative properties, and enhanced collagen deposition potential of adipose tissue-derived nanofat[131-134]. Apart from being an adipogenic derivative with ADSC, the proportion of ADSCs in nanofat is higher than that in microfat. The differences might be attributable to the method of preparation of the microfat, where the fibers and their accompanying capillaries that were the location of ADSCs are removed. In contrast, the ADSCs were mechanically separated from the native site and concentrated in nanofat, thereby making it more effective in terms of the number of ADSCs delivered to the target site. Sesé et al[135] estimated the total cellular load in mechanically prepared nanofat as 6.63 million cells/g of lipoaspirate whereas in enzymatically disintegrated SVF it was 0.68 million cells/g of lipoaspirate. The nucleated cellular count was 70% in nanofat and 7.3% in SVF. The cellular burden in nanofat contains predominantly the stromal cellular population[136].

Nanofat grafting enhances neoangiogenesis without producing any visible scars, and provides a favorable outcome in esthetic medicine for breast, buttocks, and genital augmentation, facial rejuvenation, and facial volume augmentation[137,138]. Nanofat injections retract the atrophic scars because of the presence of adipose tissue-derivedstromal cells, and avoids the need for surgical procedures. Nanofat components can regenerate dermis and subcutaneous fatty tissues and enhance the dermo-epidermal junction[139]. They regenerate by laying down adipose tissue-derived ECM, collagen deposition, and neoangiogenesis. Klinger et al[140] reported that autologous fat grafting allowed the regeneration of skin that was soft, flexible, and matched the color of neighboring skin. This concept of skin rejuvenation can be extrapolated to scars present in joints, eyelids, the face, and mouth.

Nanofat grafting beneath and within the substance of the scar improves the quality, integrity, and texture of burn scars[141]. Improved skin texture, elasticity, skin moisture, facial rejuvenation, and anti-aging properties can be achieved by combining nanofat (autocrine and paracrine effects) with platelet-rich fibrin (PRF)[142]. In a preclinical trial, nanofat injection improved the thickness of the dermal layer and promoted angiogenesis in the photoaged skin of nude mice[143]. A wide range of improvements in wrinkles, discolorations, and burn scars have been seen with nanofat application[141]. Esthetically, nanofat grafting is used for the correction of dark circles[144,145], malar bags[3], hollow eyes[145], and blepharoectomy[146]. Because of fat atrophy in the aging process, nanofat has emerged as a plausible technique for facial rejuvenation[126,147,148]. Nanofat is also being increasingly used in primary rhinoplasty procedures[149]. Nanofat is being used to correct slight skin irregularities that do not require cartilage grafting[127,150]. Segreto et al[151] evaluated the role of a combination of nanofat grafting with autologous PRP in chronic nonhealing infected
wounds, where it enhanced the regeneration of soft tissue. The multi-differentiation potential of adipose tissue, which is a component in nanofat grafting, allows evaluation in avascular necrosis of femoral head, mild to moderate grades of osteoarthritis of the knees, tendinopathies, and nonunion of fractures. Preclinical and clinical studies of nanofat use have proven its regenerative capacity in various clinical settings.

Microvascular fragments: Microvascular fragments (MVF), the byproduct of adipose tissue, range from 40–180 mm, and are composed of arteriolar, capillary, and vein segments. MVFs release VEGF and basic fibroblast growth factor (bFGF) under culture conditions. They are the richest source of proangiogenic factors that induce vasculogenesis in a paracrine fashion. MVF contains Sca 1/VEGF-2+ endothelial progenitor cells and MSCs expressing CD44+, CD73+, CD90+ and CD117+. The components of MVF exhibit the morphology of intact lumen, endothelium, and perivascular stabilizing cells. MVF are the building blocks for therapeutic vasculogenesis.

It was initially speculated that the high vascularization potential of MVFs is mainly derived from stem cell populations. McDaniel et al. compared the regenerative properties of conventionally isolated adipose-derived stem cells and multipotent cells derived from an explant culture of microvascular fragments. They found that the latter source exhibited a higher proliferation rate, an increased expression of genes involved in differentiation, and an improved ability to form capillary-like structures. In line with the concept of the “stem cell niche,” the findings indicated that compared with single-cell isolates, microvascular fragments including stem cell components provides a more physiological environment that maximizes the regenerative activity. Transplanted or injected MVFs rapidly integrate with native tissues to promote neoangiogenesis in the physiological environment. The three phases of MVF-induced neoangiogenesis are (1) immature vascular segments with high proliferation capacity; (2) vascular remodeling with a high rate of apoptosis; and (3) vascular maturation with microvascular network organization. The complex pattern of neoangiogenesis is associated with the upregulation of angiogenic genes.

Researchers have observed the regenerative potential in cartilage defects and skeletal muscle injury, myocardial infarction, partial- or complete-thickness skin defects, and diabetes mellitus. MVF loaded scaffolds can reverse lymphatic network disorders by reducing edema formation and promoting vasculogenesis in the area of repair. With the available, diverse potentials of MVF, the application of MVF in the clinical setting is plausible with the imperative question of technical and regulatory compliance from bench to bedside.

SVF: SVF is an ultra-byproduct of adipose tissue through various processing methods. The development of biocellular regenerative medicine and cellular biology, has increased the use of SVF by regenerative surgeons and researchers. The components of SVF are MSCs, HSCs, T-regulatory cells, pericyte endothelial cells, mast cells, complex microvascular beds with fibroblasts, WBC, dendritic cells, intra-adventitial smooth muscular-like cells, and others, and extracellular matrix. A sufficient number of SVF cells can be obtained without culture. Current restrictions on the use of cultured cells in humans cite the possibilities of contamination, tumorigenesis, unexpected cell differentiation, and limited cell sources. SVFs behave much like BM-MSCs. SVF mixtures contain 30% MSCs, 3% endothelial cells, and 14% endothelial precursor cells. BM-MSCs contain 0.001% MSCs, 0.1% endothelial cells, and 2% endothelial precursor cells. About 2% of isolated SVF express the hematopoietic markers CD34+ and CD 45+ and 7% express the MSC markers CD105+, and CD146+/CD133+. SVF cells express cell surface markers similar to those of BM-MSCs, such as CD105+/SH2+, CD90+, CD29+, CD44+, CD71+, and SH3+, along with low expression of CD31+, CD45+, and CD24+[184-186].

Various methods of isolation of SVF have been described in the literature. SVF isolation needs fat obtained by liposuction and transportation to a cGMP certified laboratory for further processing. The global researchers and clinicians widely use the enzymatic method for isolating and harvesting SVF by collagenase enzymes. After the addition of collagenase enzyme to the liposapate, digestion and disintegration of adipose cells take place and result in the formation of an aqueous mixture including two phases, a floating adipocyte fraction and precipitated cellular components. The separation can be enhanced by density gradient centrifugation and filtration. The lower portion of the tube contains a yellowish red pellet which is the SVF mixture that can be used for in vitro expansion.
Sharma S et al. Adipose tissue-derived MSCs

[196]. Because of the ethical issues associated with SVF isolation by the enzymatic method, researchers opt for alternative methods for SVF isolation[197,198]. Researchers proposed mechanical agitation and disruption of adipose tissue, which releases a mixture of cellular components and stromal cells. The proportion of cells in the SVF obtained by mechanical disruption is lower that obtained by enzymatic degradation, but the regenerative potential of SVF mixtures obtained by mechanical disruption and enzymatic degradation are the same[198]. The anti-inflammatory effects of SVF result from the presence of higher amounts of IL-1 and 10 receptor antagonists, which are expressed by TNF-α and leptin released by monocytes and macrophages in the SVF mixture[199]. SVF mixtures provoke adaptive immunity by producing T-reg cells by suppressing the activation of dendritic cells[200,201].

Krawiec et al.[202] demonstrated that SVF components seeded with biodegradable porous scaffolds in vivo for 8 wk resulted in the generation of tissue-engineered vascular grafts when populated with primary vascular components such as smooth muscle cells, endothelial cells, collagen, and elastin. ADSCs or SVF combined with PRP enhanced the regenerative potential in the form of neoangiogenesis, cellular proliferation, and differentiation[203]. The combination of SVF with PRP enhanced the stemness of MSCs, and the growth factors present in PRP induced locally available stem cells and prolonged the survival time and survival rates of cells present in the PRP[204]. Such combination treatments have been evaluated in preclinical and clinical trials of wound healing, osteoarthritis of the knees, bone and tendon regeneration, periodontal engineering, fat grafting procedures, and vascular diseases[205-208].

Osteochondral defects of the knees in goats were managed by the application of autologous type 1 and 3 collagen scaffolds along with SVF cells[209]. In rat models of acute kidney injury, the transplantation of autologous SVF cells induced antiapoptotic effects and the release of growth factors like VEGF and HGF to regenerate kidney cells [210]. In preclinical studies, the therapeutic effect, safety and functional outcome of autologous SVF uncultured or culture-expanded cells were observed in acute myocardial infarction, skin flap necrosis, and erectile dysfunction with cavernous nerve injury[211-213]. In human clinical studies, the use of SVF cells was extensively studied in conditions like breast reconstruction surgery, traumatic calvaria defects, types 1 and 2 diabetes mellitus, Crohn’s disease with enterocutaneous fistula, and burn injuries[214-218]. Zhao et al[219] evaluated the efficacy of SVF in diabetic feet in terms of cellular survival, proliferation rate and differentiation and they determined the characteristics of transplanted cells. Various researchers demonstrated endobronchial administration of autologous SVF cells for treating idiopathic pulmonary fibrosis[220]. Autologous SVF is also being evaluated in COVID-19 individuals[221-223]. When cocultured with 5% or 20% PRP, the viability, motility, proliferation rate, and differentiation of ADSCs were increased[224]. In the case of fat grafting, clinicians are challenged by the fat graft rejection, with a reported resorption rate ranging from 25%-80%, which is mostly the result of apoptosis of mature adipocytes[225]. When the lipoaspirate was given along with SVF, a 35% greater graft retention was noted[194]. Further, more prominent microvasculature was noted compared with normal graft tissue, suggesting its clinical potential as a nourishing medium that enhanced the efficacy of the native therapy. Moreover, in situations such as osteoarthritis, an ultrafiltrate fraction of an adipose derivative such as SVF would be the ideal medium of choice to tap the benefits of ADSCs in cartilage regeneration[226].

**ADSC exosomes:** Exosomes are the cell-free regenerative tool in the field of tissue engineering. Exosomes are extracellular vehicles that are endosome-derived lipid bilayer spherical vesicles of 40 to 150 nm in size. They are found in all cell types and body fluids and are comprised of cytokines, proteins, lipids, DNA, and RNA from the parent cell. Exosomes are an integral part of both the diagnostic and therapeutic methods used in various diseases. ADSC exosomes (EXOs) differ from other MSC EXOs in proliferation and differentiation abilities and immunosuppressive pathways. Compared with ADSCs, ADSC-derived exosomes possess a high biosafety profile with low immunogenicity[227]. They protect cargoes from degradation, have tissue and target specificity, good tissue permeability, intercellular signaling and communication, immune function, tissue homeostasis, and development of cell fate [227].

Ogawa et al[228] observed that mRNAs of adipose tissue-derived exosomes have a significant role in metabolic, immunological, and cellular responses. Various studies demonstrated that adipose tissue-derived exosomes possess the capability of regeneration of muscles and bones, promotion of wound healing, and enhancement of cellular proliferation, and neoangiogenesis. ADSCs-EXOs enhance the proliferation and migration of vascular endothelial cells and promote vasculogenesis. They retain
the fat graft volume by the stimulation of angiogenesis and regulating inflammatory responses[229,230]. ADSC EXOs were tested by Wang et al[231] for the promotion of wound healing in diabetic mice by enhancing angiogenesis, proliferation of fibroblasts, and collagen synthesis in the later stages. The evidence also supports the use of ADSC EXOs for treating diabetic foot patients[232]. To stabilize the ADSC EXOs concentration when used for local application, bioscaffolds like hydrogels or fibrin are augmented with exosomes to delay release at the therapeutic site[233,234]. ADSC EXOs are used for scarless cutaneous repair by retracting the size of scars, increasing the collagen 3 to collagen 1 ratio, and by regulating the migration, proliferation, differentiation, and gene expression of fibroblasts[235]. Kim et al[236] demonstrated that key cytokines and growth factors in ADSC EXOs facilitated tissue regeneration and repair through anti-oxidation, anti-wrinkle and skin-whitening activity. ADSC EXOs with multiple bioactive molecules for the management of aging warrants further extensive research. Zhao et al[237] demonstrated crosstalk that facilitated immune and metabolic homeostasis, providing a vital therapy for obesity and diabetes mellitus. ADSC EXOs combined with poly(lactide-co-glycolide) scaffolds improved the osteoinductive effects, MSC migration, and homing abilities in bone regeneration[238]. Chen et al[239] demonstrated that exosomes derived from miR-375-overexpressing ADSCs incorporated with hydrogel enhanced bone regeneration in a rat model of calvarial defect. Direct stem cell transplantation with ADSC EXOs primed by TNF-α, promoted the proliferation and differentiation of human osteoblasts promoted through the Wnt signaling pathway, which further widened the application of exosomes in bone regeneration[240]. Pers et al[241,242] reported that ADSC EXOs were a safe, effective, and inexpensive therapy for osteoarthritic knees. ADSC EXOs downregulated inflammation and oxidative stress in osteoarthritic knees[243]. Zhang et al[244] confirmed that intra-articular injection of ADSC EXOs inhibited cartilage and subchondral bone degradation and osteophyte formation and slowed the progression of the disease process in osteoarthritis. ADSC EXOs participated cellular communications and applications in plastic and cosmetic surgery. Although applications in clinical practice are lacking, ADSC EXOs have an increasing role in maximizing the therapeutic effectiveness for dermopathies and for tissue reconstruction[245]. There is a future for exosomes as a diagnostic tool for early identification of pathological processes involved in disease and as a therapeutic, targeted acellular conduit to optimize the pathological milieu at the site of interest.

**Ethical concerns**

An early therapy and drug development are dependent on translational phases, the best strategy is to standardize the process, and regularly acquire data to validate and certify the technologies being developed. The safety and efficacy of the developed therapy must be proven from various perspectives, including the donor, recipient, product, manufacturing, clinical application, and biovigilance. The procedures involved in making adipose-derived products involve the collection and preparation of fat tissue. The procedures require that the cells are obtained by safe methods without any adverse events. Clinicians who use the cells in their practice are obliged to abide by existing laws that control the use in clinical practice. The European Medicines Agency (EMA), United States Food and Drug Administration (FDA), and others consider adult human cells as biological products of two classes. The first involves cells processed with minimal manipulation techniques such as filtration, centrifugation, and mechanical disruption done in a single surgical window within a sterile operating environment. The second class of products undergoes significant manipulation such as expansion in cell culture, characterization, cultivation, and other manipulation techniques used outside of a single surgical window or the operating room. As per FDA and EMA regulations, adipose products are considered medicinal products that have to be harvested using validated procedures adhering to stringent regulatory protocols. The clinical quality attributes of therapeutic products ensure safety along with the maintenance of identity, purity, and potency as per the FDA Guidance for industry [Pharmaceutical development. Q8(R2). Current Step 4 version dated August 2009.Q8 (R2)]. In agreement with the reflection paper EMA/CAT/-600280/2010 Rev 1, 20 June 2014, it is presumed that for autologous use under a single therapeutic window with minimal manipulation in aseptic conditions, adipose-derived products would not require ethical committee underwriting for clinical use. However, there are no standardized methods or means to isolate ADSCs with predetermined critical quality attribute, making isolation of ADSC for targeted therapies difficult[246]. Hence, a modular system configured as a single-use kit that contained all the essential components in the exact proportions needed would be ideal and is recommended. Such kits would allow following a specific protocol to
standardize isolation of adipose derivatives with targeted action of ADSCs and comply with specific critical quality attributes.

Having discussed the practical constraints of the use of ADSCs for practical applications, the use of ADSC-derived exosomes as a therapeutic conduit is being explored [247]. Exosomes do not carry the risk of genetic instability and immune activation following their administration in the host environment. The differential advantage of using them vs their parent cell of origin include immune privilege in the host environment, which helps them to evade the native phagocytosis, their size in nanoscale enabling them to move in and out of cells with ease, their homing molecules on their surface enabling them to migrate to their site of interest [248]. Hence, with the above-mentioned advantages in using ADSC-derived exosomes, they are the principal focus of research in advancing regenerative therapy in the future.

CONCLUSION

Of late, interest in adipose derivatives for regenerative therapies and constructive tissue engineering is on the rise. Having elaborated the biology, characteristics, immunology, and clinical applications of adipose-derived products, it is imperative that evidence to strengthen the clinical applicability of these therapeutic products is needed to warrant an official recommendation from regulatory authorities. Unraveling that evidence to strengthen the clinical applicability of these therapeutic products is imperative in advancing regenerative therapy in the future.

REFERENCES

1. Chu DT, Nguyen Thi Phuong T, Tien NLB, Tran DK, Minh LB, Thanh VV, Gia Anh P, Pham VH, Thi Ngoc V. Adipose Tissue Stem Cells for Therapy: An Update on the Progress of Isolation, Culture, Storage, and Clinical Application. *J Clin Med* 2019; 8 [PMID: 31247996] DOI: 10.3390/jcm8070197
2. Han Y, Li X, Zhang Y, Han Y, Chang F, Ding J. Mesenchymal Stem Cells for Regenerative Medicine. *Cells* 2019; 8 [PMID: 31412678] DOI: 10.3390/cells8080886
3. Simonacci F, Bertozzi N, Griceo MP, Grignaffini E, Raposio E. Procedure, applications, and outcomes of autologous fat grafting. *Ann Med Surg (Lond)* 2017; 20: 49-60 [PMID: 28702187] DOI: 10.1016/j.amsu.2017.06.059
4. Chen FM, Liu X. Advancing biomaterials of human origin for tissue engineering. *Prog Polym Sci* 2016; 53: 86-168 [PMID: 27022202] DOI: 10.1016/j.progpolymsci.2015.02.004
5. Langer R, Vacanti JP. Tissue engineering. *Science* 1993; 260: 920-926 [PMID: 8493529] DOI: 10.1126/science.8493529
6. Feinberg AW. Engineered tissue grafts: opportunities and challenges in regenerative medicine. *Wiley Interdiscip Rev Syst Biol Med* 2012; 4: 207-220 [PMID: 22012681] DOI: 10.1002/wsbm.164
7. Hsu VM, Stramsky CA, Bucky LP, Percec I. Fat grafting’s past, present, and future: why adipose tissue is emerging as a critical link to the advancement of regenerative medicine. *Aesthet Surg J* 2012; 32: 892-899 [PMID: 22942117] DOI: 10.1177/1090820X12455658
8. Tan SS, Ng ZY, Zhan W, Rozen W. Role of Adipose-derived Stem Cells in Fat Grafting and Reconstructive Surgery. *J Cutan Aesthet Surg* 2016; 9: 152-156 [PMID: 27761084] DOI: 10.4103/0974-2077.191672
9. Flynn L, Woodhouse KA. Adipose tissue engineering with cells in engineered matrices. *Organogenesis* 2008; 4: 228-235 [PMID: 19337402] DOI: 10.4161/org.4.4.7082
10. Nikolova MP, Chavali MS. Recent advances in biomaterials for 3D scaffolds: A review. *Bioact Mater* 2019; 4: 271-292 [PMID: 31709311] DOI: 10.1016/j.bioactmat.2019.10.005
11. Song R, Murphy M, Li C, Ting K, Soo C, Zheng Z. Current development of biodegradable polymeric materials for biomedical applications. *Drug Des Devel Ther* 2018; 12: 3117-3145 [PMID: 30288019] DOI: 10.2147/DDDT.S165440
12. Bahmad HF, Daouk R, Azar J, Sapudom J, Teo JCM, Abou-Kheir W, Al-Sayegh M. Modeling Adipogenesis: Current and Future Perspective. *Cells* 2020; 9 [PMID: 33092038] DOI: 10.3390/cells9102326
13. Young DA, Christian KL. Injectable biomaterials for adipose tissue engineering. *Biomed Mater* 2012; 7: 024104 [PMID: 22456805] DOI: 10.1088/1748-6041/7/2/024104
14. Klingelhutz AJ, Gouronne FA, Chaly A, Wadkins DA, Burand AJ, Markan KR, Idiga SO, Wu M, Pothoff MJ, Ankrom JA. Scaffold-free generation of uniform adipose spheroids for metabolism research and drug discovery. *Sci Rep* 2018; 8: 523 [PMID: 29323267] DOI: 10.1038/s41598-018-19024-z
15. Turner PA, Gurumurthy B, Bailey JL, Elks CM, Janokar A. Adipogenic Differentiation of Human Adipose-Derived Stem Cells Grown as Spheroids. *Process Biochem* 2017; 59: 312-320 [PMID: 28966553] DOI: 10.1016/j.procbio.2017.02.003
Sharma S et al. Adipose tissue-derived MSCs

Stenkula KG, Erlanson-Albertsson C. Adipose cell size: importance in health and disease. *Am J Physiol Regul Integr Comp Physiol* 2018; 315: R284-R295 [PMID: 29641234 DOI: 10.1152/ajpregu.00257.2017]

Church C, Horowitz M, Rodeheffer M. WAT is a functional adipocyte? *Adipocyte* 2012; 1: 38-45 [PMID: 23708509 DOI: 10.4161/adip.19132]

Berry DC, Stenesen D, Zeve D, Graff JM. The developmental origins of adipose tissue. Development 2013; 140: 3393-3394 [PMID: 24043615 DOI: 10.1242/dev.080549]

Fasshauer M, Blüher M. Adipokines in health and disease. *Trends Pharmacol Sci* 2015; 36: 461-470 [PMID: 26029234 DOI: 10.1016/j.tips.2015.04.014]

Zalechowska P, Kozlowska E, Pastwińska J, Agier J, Brzezińska-Blaszczyk E. Adipocytokine involvement in innate immune mechanisms. *J Interferon Cytokine Res* 2018; 38: 527-538 [PMID: 30431386 DOI: 10.1089/jir.2018.0102]

Coelho M, Oliveira T, Fernandes R. Biochemistry of adipose tissue: an endocrine organ. *Arch Med Sci* 2013; 9: 191-200 [PMID: 23671428 DOI: 10.5111/aoms.2013.33181]

Kershaw EE, Flier JS. Adipose tissue as an endocrine organ. *J Clin Endocrinol Metab* 2004; 89: 2548-2556 [PMID: 15181022 DOI: 10.1210/jc.2004-0395]

Mohamed-Ali V, Pinkney JH, Coppack SW. Adipose tissue as an endocrine and paracrine organ. *Int J Obes Relat Metab Disord* 1998; 22: 1145-1158 [PMID: 9877249 DOI: 10.1038/sj.oj.ijo.0800770]

Fantuzzi G. Adipose tissue, adipokines, and inflammation. *J Allergy Clin Immunol* 2005; 115: 911-919; quiz 920 [PMID: 15867843 DOI: 10.1016/j.jaci.2005.02.023]

Ullah I, Subbarao RB, Rho GJ. Human mesenchymal stem cells - current trends and future prospective. *Biosci Rep* 2015; 35 [PMID: 25797907 DOI: 10.1042/BSR20150025]

Moon SH, Kim JM, Hong KS, Shin JM, Kim J, Chung HM. Differentiation of hESCs into Mesodermal Subtypes: Vascular-, Hematopoietic- and Mesenchymal lineage cells. *Int J Stem Cells* 2011; 4: 24-34 [PMID: 22493334 DOI: 10.15283/ijsc.2011.4.1.24]

Ballini A, Sacco S, Coletti D, Pluchino S, Tatullo M. Mesenchymal Stem Cells as Promoters, Enhancers, and Playmakers of the Translational Regenerative Medicine. *Stem Cells Int* 2017; 2017: 3292810 [PMID: 28740512 DOI: 10.1155/2017/3292810]

Fraser JK, Zhu M, Wuhr I, Alfonso Z. Adipose-derived stem cells. *Methods Mol Biol* 2008; 449: 59-67 [PMID: 18370083 DOI: 10.1007/978-1-60327-169-1_4]

Peer LA. Cell survival theory versus replacement theory. *Plast Reconstr Surg (1946)* 1955; 16: 161-168 [PMID: 13266544 DOI: 10.1097/00006534-195509000-00001]

Pu LL. Mechanisms of Fat Graft Survival. *Ann Plast Surg* 2016; 77 Suppl 1: S84-S86 [PMID: 26808753 DOI: 10.1097/SAP.0000000000000730]

Bellini E, Grieco MP, Raposio E. The science behind autologous fat grafting. *Ann Med Surg (Lond)* 2017; 24: 65-73 [PMID: 29188051 DOI: 10.1016/j.amsu.2017.11.001]

Harrison BL, Malafa M, Davis K, Rohrich RJ. The discordant histology of grafted fat: a systematic review of the literature. *Plast Reconstr Surg* 2015; 135: 542e-555e [PMID: 25719719 DOI: 10.1097/PRS.0000000000006973]

Hong KY. Fat grafts enriched with adipose-derived stem cells. *Arch Craniofac Surg* 2020; 21: 211-218 [PMID: 32687409 DOI: 10.7181/acfs.2020.00525]

Eto H, Kato H, Suga H, Aoi N, Doi K, Kuno S, Yoshimura K. The fate of adipocytes after nonvascularized fat grafting: evidence of early death and replacement of adipocytes. *Plast Reconstr Surg 2012; 129*: 1081-1092 [PMID: 22261362 DOI: 10.1097/PRS.0b013e318242ab19]

Yoshimura K, Eto H, Kato H, Doi K, Aoi N. In vivo manipulation of stem cells for adipose tissue repair/reconstruction. *Regen Med* 2011; 6: 33-41 [PMID: 21999260 DOI: 10.2217/rmc.11.62]

Stringa M, Viswanathan S, Darinskas A, Slaby O, Michalek J. Same or not the same? *Stem Cells Dev* 2012; 21: 2724-2752 [PMID: 22468918 DOI: 10.1089/scd.2011.0722]

Lee SH. The advantages and limitations of mesenchymal stem cells in clinical application for treating human diseases. *Osteoporos Sarcopenia* 2018; 4: 150 [PMID: 30775559 DOI: 10.1016/j.jafs.2018.11.083]

Musial-Wysoka A, Kot M, Majka M. The Pros and Cons of Mesenchymal Stem Cell-Based Therapies. *Cell Transplant* 2019; 28: 801-812 [PMID: 31018669 DOI: 10.1177/0963689719837897]

Schneider S, Unger M, van Griensven M, Balmayer ER. Adipose-derived mesenchymal stem cells from liposuction and resected fat are feasible sources for regenerative medicine. *Eur J Med Res* 2017; 22: 17 [PMID: 28526089 DOI: 10.1186/s40001-017-0258-9]

Simonacci F, Bertozzi N, Grieco MP, Raposio E. From liposuction to adipose-derived stem cells: indications and technique. *Acta Biomed* 2019; 90: 197-208 [PMID: 31124996 DOI: 10.23750/abm.v90i2.6619]

Mizuno H. Adipose-derived stem cells for tissue repair and regeneration: ten years of research and a literature review. *J Nippon Med Sch* 2009; 76: 56-66 [PMID: 19443990 DOI: 10.1272/jnms.76.56]

Meza-Zepeda LA, Noer A, Dahl JA, Micci F, Myklebost O, Collas P. High-resolution analysis of genetic stability of human adipose tissue stem cells cultured to senescence. *J Cell Mol Med* 2008; 12: 553-566 [PMID: 18419597 DOI: 10.1111/j.1582-4934.2007.00146.x]

Lindroos B, Suuronen R, Miettinen S. The potential of adipose stem cells in regenerative medicine. *Stem Cell Rev Rep* 2011; 7: 269-291 [PMID: 20853072 DOI: 10.1007/s12015-010-9193-7]

Gimble JM, Katz AJ, Bunnell BA. Adipose-derived stem cells for regenerative medicine. *Circ Res* 2007; 100: 1249-1260 [PMID: 17405232 DOI: 10.1161/01.RES.0000265074.83288.09]
Sharma S et al. Adipose tissue-derived MSCs

45 Alonso-Goulart V, Ferreira LB, Duarte CA, Isabel Lemos de Lima, Enza Rafaela Ferreira, Bárbara Candido de Oliveira, Luna Nascimento Vargas, Dayane Dotto de Moraes, Isaura Beatriz Borges Silva, Rafael de Oliveira Faría, Aline Gomes de Souza, Letícia de Souza Castro-Filipe. Mesenchymal stem cells from human adipose tissue and bone repair: a literature review. Biotechnol Res Innov 2018; 2: 74-80 [DOI: 10.1016/j.bire.2017.10.005]

46 Mohamed-Ahmed S, Fristad I, Lie SA, Suliman S, Mustafa K, Vindenes H, Idris SB. Adipose-derived and bone marrow mesenchymal stem cells: a donor-matched comparison. Stem Cell Res Ther 2018; 9: 168 [PMID: 29921311 DOI: 10.1186/s13287-018-0941-1]

47 Luna AC, Madeira ME, Conceição TO, Moreira JA, Laiso RA, Maria DA. Characterization of adipose-derived stem cells of anatomical region from mice. BMC Res Notes 2014; 7: 552 [PMID: 25138545 DOI: 10.1186/1756-0500-7-552]

48 Mitchell JB, McIntosh K, Zvonie S, Garrett S, Floyd ZE, Kloster A, Di Halvorsen Y, Storms RW, Goh B, Kilroy G, Wu X, Gimble JM. Immunophenotype of human adipose-derived cells: temporal changes in stromal-associated and stem cell-associated markers. Stem Cells 2006; 24: 376-385 [PMID: 16322640 DOI: 10.1634/stemcells.2005-0234]

49 Screven R, Kenyon E, Myers MJ, Yancy HF, Skasko M, Boxer L, Bigley EC 3rd, Borch Jessl DN, Zhu M. Immunophenotype and gene expression profile of mesenchymal stem cells derived from canine adipose tissue and bone marrow. Vet Immunol Immunopathol 2014; 161: 21-31 [PMID: 25026887 DOI: 10.1016/j.vetimm.2014.06.002]

50 Rady D, Abbass MMS, El-Rashidy AA, El Moshy S, Radwan IA, Dörfir CE, Fawzy El-Sayed KM. Mesenchymal Stem/Progenitor Cells: The Prospect of Human Clinical Translation. Stem Cells Int 2020; 2020: 8837654 [PMID: 33953753 DOI: 10.1155/2020/8837654]

51 Bogdanova A, Berzins U, Nikulshin S, Sindrastina D, Ezerza A, Ledzidina D, Kozlovskaya T. Characterization of human adipose-derived stem cells cultured in autologous serum after subsequent passage and long term cryopreservation. J Stem Cells 2014; 9: 135-148 [PMID: 25177468]

52 Zhu M, Dörfer CE, Fawzy El-Sayed KM, Brinchmann JE. Concise review: therapeutic potential of adipose tissue-derived angiogenic cell populations. J Stem Cells 2010; 3: 27778035 [DOI: 10.4172/09749578.10001157]

53 Hoogduijn MJ, Roemeling-van Rhijn M, Korevaar SS, Engela AU, Weimar W, Baan CC. Immunological aspects of allogeneic and autologous mesenchymal stem cell therapies. Hem Gene Ther 2011; 22: 1587-1591 [PMID: 21732766 DOI: 10.1089/hum.2011.039]

54 Haddad R, Saldanha-Araujo F. Mechanisms of T-cell immunosuppression by mesenchymal stromal cells: what do we know so far? Biomed Res Int 2014; 2014: 216806 [PMID: 25025040 DOI: 10.1155/2014/216806]

55 Mazini L, Rochette L, Amine M, Malka G. Regenerative Capacity of Adipose Derived Stem Cells (ADSCs), Comparison with Mesenchymal Stem Cells (MSCs). Int J Mol Sci 2019; 20: 31121953 DOI: 10.3390/ijms20102523]

56 Zhong YC, Wang SC, Han YH, Wen Y. Recent Advance in Source, Property, Differentiation, and Applications of Infrapatellar Fat Pad Adipose-Derived Stem Cells. Stem Cells Int 2020; 2020: 2560174 [PMID: 32215015 DOI: 10.1155/2020/2560174]

57 Zhao Y, Waldman SD, Flym LE. The effect of serial passaging on the proliferation and differentiation of bovine adipose-derived stem cells. Cells Tissues Organs 2012; 195: 414-427 [PMID: 21893933 DOI: 10.1159/000329255]

58 Wai E, Chen Y, Wang X, Chen X, Xiong C. Characterization of human adipose-derived stem cells derived from adipose tissue and bone marrow. Vet Immunol Immunopathol 2014; 161: 21-31 [PMID: 25026887 DOI: 10.1016/j.vetimm.2014.06.002]

59 Wei W, Xu C, Ye ZY, Huang XJ, Yuan JE, Ma TB, Lin HB, Chen XQ. Biological characteristics of mesenchymal stem cell and hematopoietic stem cell in the co-culture system. Sheng Li Xue Bao 2016; 68: 691-698 [PMID: 27778036]

60 Perucca S, Di Palma A, Piccaluga PP, Gemelli C, Zoratti E, Bassi G, Giacopuzzi E, Lojano M, Brinchmann JE. Concise review: therapeutic potential of adipose tissue-derived angiogenic cell populations. J Stem Cells 2010; 3: 27778035 [DOI: 10.4172/09749578.10001157]

61 Gritti A, Judson RJ, Stockwell TR, Corbett AB, Begley CG, Zivin JA, Hynynen K. Progress toward clinical translation of mesenchymal stem cells: an update. Stem Cells Int 2020; 2020: 2560174 [PMID: 32215015 DOI: 10.1155/2020/2560174]

62 Bourin P, Bunnell BA, Castellani L, Dominici M, Katz AJ, March KL, Redl H, Rubin JP, Yoshinura K, Gimble JM. Stromal cells from the adipose tissue-derived stromal vascular fraction and culture expanded adipose tissue-derived stromal/stem cells: a joint statement of the International Federation for Adipose Therapeutics and Science (IFATS) and the International Society for Cellular Therapy (ISCT). Cytotherapy 2013; 15: 641-648 [PMID: 23570660 DOI: 10.1016/j.jcyt.2013.02.006]
Characterization and Expansion of Human Adipose-Derived Stem Cells (ASCs): An Overview.

Mol Sci Characterization and Expansion of Human Adipose-Derived Stem Cells (ASCs): An Overview.

Baer PC, Geiger H. Adipose-derived mesenchymal stromal/stem cells: tissue localization, characterization, and heterogeneity. Cell Stem Cells 2012; 8: 154-158 [PMID: 23110271] DOI: 10.4252/wjsc.v12.i1.1

Leto Barone AA, Croitoru A, Siragusa G, Cifone MG, Cinque B, Giuliani M. Methods of Isolation, Characterization and Expansion of Human Adipose-Derived Stem Cells (ASCs): An Overview. Int J Mol Sci 2018; 19 [PMID: 29958391] DOI: 10.3390/ijms19071897

Sharma S et al. Adipose tissue-derived MSCs.
Dubey NK, Mishra VK, Dubey R, Deng YH, Tsai FC, Deng WP. Revisiting the Advances in Isolation, Characterization and Secretome of Adipose-Derived Stromal/Stem Cells. *Int J Mol Sci* 2018; 19 [PMID: 30060511 DOI: 10.3390/ijms19082200]

Bunnell BA, Flaat M, Gagliardi C, Patel B, Ripoll C. Adipose-derived stem cells: isolation, expansion and differentiation. *Methods* 2008; 45: 115–120 [PMID: 18593669 DOI: 10.1016/j.ymeth.2008.03.006]

Mailey B, Hosseini A, Baker J, Young A, Alfonso Z, Hicok K, Wallace AM, Cohen SR. Adipose-derived stem cells: methods for isolation and applications for clinical use. *Methods Mol Biol* 2014; 1210: 161–181 [PMID: 25171368 DOI: 10.1007/978-1-4939-1435-7_11]

Baer PC. Adipose-derived mesenchymal stromal/stem cells: An update on their phenotype in vivo and in vitro. *World J Stem Cells* 2014; 6: 256–265 [PMID: 25126376 DOI: 10.4252/wjsc.v6.i3.256]

Zimmerlin L, Donnenberg VS, Rubin JP, Donnenberg AD. Mesenchymal markers on human adipose stem/progenitor cells. *Cytometry A* 2013; 83: 134–140 [PMID: 23184564 DOI: 10.1002/cyto.a.22227]

Brooks AES, Inimitoff M, Williams E, Damanti T, Jackson-Patel V, Fan V, James J, Dunbar PR, Feist V, Sheppard HM. *Ex Vivo Human Adipose Tissue Derived Mesenchymal Stromal Cells (ASC) Are a Heterogeneous Population That Demonstrate Rapid Culture-Induced Changes*. *Front Pharmacol* 2019; 10: 1695 [PMID: 32153389 DOI: 10.3389/fphar.2019.01695]

Wu YD, Li M, Liao X, Li SH, Yan JX, Fan L, She WL, Song JX, Liu HW. Effects of storage culture media, temperature and duration on human adipose-derived stem cell viability for clinical use. *Mol Med Rep* 2019; 19: 2189–2201 [PMID: 30664198 DOI: 10.3892/mmr.2019.9842]

Shaik S, Wu X, Gimble J, Devireddy R. Effects of Decade Long Freezing Storage on Adipose Derived Stem Cells Functionality. *Sci Rep* 2018; 8: 8162 [PMID: 29023253 DOI: 10.1038/s41598-018-26546-7]

De Rosa A, De Francesco F, Tirino V, Ferraro GA, Desiderio V, Paino F, Pirozzi G, D’Andrea F, Papaccio G. A new method for cryopreserving adipose-derived stem cells: an attractive and suitable large-scale and long-term cell banking technology. *Tissue Eng Part C Methods* 2009; 15: 659–667 [PMID: 19254116 DOI: 10.1089/tenc.TEC.2008.0674]

Choi JR, Yong KW, Wan Safwani WKZ. Effect of hypoxia on human adipose-derived mesenchymal stem cells and its potential clinical applications. *Cell Mol Life Sci* 2017; 74: 2587–2600 [PMID: 28224204 DOI: 10.1007/s00018-017-2484-2]

Yong KW, Choi JR, Dolbashid AS, Wan Safwani WKZ. Biosafety and bioefficacy assessment of human mesenchymal stem cells: what do we know so far? *Regen Med* 2018; 13: 219–232 [PMID: 29509072 DOI: 10.2217/rme-2017-0078]

Bai X, Yan Y, Song YH, Seidensticker M, Rabinovich B, Metzele R, Bankson JA, Vykoukal D, Alt Y. Both cultured and freshly isolated adipose tissue-derived stem cells enhance cardiac function after acute myocardial infarction. *Eur Heart J* 2010; 31: 489–501 [PMID: 20037143 DOI: 10.1093/eurheartj/ehp568]

Hassan WU, Greiser U, Wang W. Role of adipose-derived stem cells in wound healing. *Wound Repair Regen* 2014; 22: 313–325 [PMID: 24844331 DOI: 10.1111/wrr.12173]

Shingyochi Y, Orbay H, Mizuno H. Adipose-derived stem cells for wound repair and regeneration. *Expert Opin Biol Ther* 2015; 15: 1285–1292 [PMID: 26037027 DOI: 10.1517/14712598.2015.1053867]

Sabol RA, Bowles AC, Côté A, Wise R, Pashos N, Bunnell BA. Therapeutic Potential of Adipose Stem Cells. *Adv Exp Med Biol* 2018 [PMID: 30051318 DOI: 10.1007/978-1-4939-1435-7_11]

Björninen M, Gilmore K, Pelto J, Seppänen-Kaijansinkko R, Kellomäki M, Miettinen S, Wallace G, Grijpma D, Haimi S. Electrically Stimulated Adipose Stem Cells on Polypryrole-Coated Scaffolds for Smooth Muscle Tissue Engineering. *Ann Biomed Eng* 2017; 45: 1015–1026 [PMID: 27844175 DOI: 10.1007/s10439-016-1755-7]

Dall’Oca C, Breda S, Elena N, Valenti R, Samaila EM, Magnan B. Mesenchymal Stem Cells injection in hip osteoarthritis: preliminary results. *Acta Biomed* 2019; 90: 75–80 [PMID: 30715002 DOI: 10.23750/abm.v90i1-S.8084]

Jones IA, Wilson M, Togashi R, Han B, Mircheff AK, Thomas Vangness C Jr. A randomized, controlled study to evaluate the efficacy of intra-articular, autologous adipose tissue injections for the treatment of mild-to-moderate knee osteoarthritis compared to hyaluronic acid: a study protocol. *BMC Musculoskelet Disord* 2018; 19: 383 [PMID: 30355133 DOI: 10.1186/s12891-018-2300-7]

Lee WS, Kim HH, Kim KJ, Kim GB, Jin W. Intra-Articular Injection of Autologous Adipose Tissue-Derived Mesenchymal Stem Cells for the Treatment of Knee Osteoarthritis: A Phase Ib, Randomized, Placebo-Controlled Clinical Trial. *Stem Cells Transl Med* 2019; 8: 504–511 [PMID: 30835956 DOI: 10.1002/stem.18-0122]

Agostini F, Rossi FM, Aldinucci D, Battiston M, Lombardi E, Zanolin S, Massarut S, Parodi PC, Da Ponte A, Tessitori G, Pivetta B, Durante C, Mazzuccato M. Improved GMP compliant approach to manipulate liposapirates, to cryopreserve stromal vascular fraction, and to expand adipose stem cells in xeno-free media. *Stem Cell Res Ther* 2018; 9: 130 [PMID: 29751821 DOI: 10.1186/s13287-018-0886-1]

Rozato I, Belisario DC, Compago M, Lena A, Bistolfi A, Maccari L, Mussano F, Genova T, Godio L, Peralle G, Formica M, Cambieri I, Castagnoli C, Robba T, Felli L, Ferracini R. Concentrated adipose tissue infusion for the treatment of knee osteoarthritis: clinical and histological observations. *Int Orthop* 2019; 43: 15–23 [PMID: 30311059 DOI: 10.1007/s00264-018-4192-4]
Serratrice N, Bruzzese L, Magalon J, Véran J, Giraudo L, Aboudou H, Ould-Ali D, Nguyen PS, Sharma S et al. Adipose tissue-derived MSCs
Bausset O, Daumas A, Casanova D, Granel B, Andrac-Meyer L, Sabatier F, Magalon G. New fat-derived products for treating skin-induced lesions of scleroderma in nude mice. Stem Cell Res Ther 2014; 5: 138 [PMID: 25519759 DOI: 10.1186/scrt5528]

Memar O, Nezamabadi A, Milani BY, Milani FY, Djalilian A. Nanofat grafting: basic research and clinical application. Plast Reconstr Surg 2014; 133: 728e [PMID: 24776592 DOI: 10.1097/PRS.0000000000000135]

Frijtt MT. Nanofat grafting: basic research and clinical applications. Plast Reconstr Surg 2014; 134: 33e-33ae [PMID: 25068362 DOI: 10.1097/PRS.0000000000000333]

Gentile P, Scioli MG, Bielli A, Orlandi A, Cervelli V. Comparing different nanofat procedures on scars: role of the stromal vascular fraction and its clinical variation. Regen Med 2017; 12: 939-952 [PMID: 29236575 DOI: 10.2217/mae-2017-0076]

Sesé B, Sammartini JM, Ortega B, Matas-Palau A, Lull R. Nanofat Cell Aggregates: A Nearly Constitutive Stromal Cell Inoculum for Regenerative Site-Specific Therapies. Plast Reconstr Surg 2019; 144: 1079-1088 [PMID: 31454336 DOI: 10.1097/PRS.0000000000002615]

Copcu HE, Oztan S. New Mechanical Fat Separation Technique: Adjustable Regenerative Adipose-tissue Transfer (ARAT) and Mechanical Stromal Cell Transfer (MEST). Aesthet Surg J Open Forum 2020; 2: oja035 [PMID: 33791661 DOI: 10.1093/asjof/oja035]

Menkes S, Luca M, Soldati G, Polla L. Subcutaneous Injections of Nanofat Adipose-derived Stem Cell Grafting in Facial Rejuvenation. Plast Reconstr Surg Glob Open 2020; 8: e2550 [PMID: 32095390 DOI: 10.1097/GOX.0000000000002550]

Cohen SR. Hewett S, Ross L, Delaunay F, Goodacre A, Ramos C, Leong T, Saad A. Regenerative Cells For Facial Surgery: Biofilling and Biocontouring. Aesthet Surg J 2017; 37: S16-S32 [PMID: 29025218 DOI: 10.1093/asjgjx078]

Fakhii-Gomez N, Steward E, Orte Aldea M del C. Nanofat in Facial Rejuvenation: Step-by-Step Procedure, Patient Evaluation and Recovery Process. The American Journal of Cosmetic Surgery 2021; 38: 27-35 [DOI: 10.1177/0748480082091931]

Klinger M, Cavigioli F, Klingler FM, Giannasi S, Bandi V, Banzatti B, Forcellini D, Maione L, Catania B, Vinci V. Autologous fat graft in scar treatment. J Craniofac Surg 2013; 24: 1610-1615 [PMID: 24036737 DOI: 10.1097/SCS.0b013e3182a24548]

Jan SN, Bashir MM, Khan FA, Hidayat Z, Ansari HH, Sohail M, Bajwa AB, Shami HB, Hanif A, Oztan S. New Mechanical Fat Separation Technique: Adjustable Regenerative Adipose-tissue Transfer (ARAT) and Mechanical Stromal Cell Transfer (MEST). Aesthet Surg J Open Forum 2020; 217-222 [PMID: 32453929 DOI: 10.1097/PRS.0000000000006155]

Xu P, Yu Q, Huang H, Zhang WJ, Li W. Nanofat Increases Dermis Thickness and Neovascularization in Photoaged Nude Mouse Skin. Aesthetic Plast Surg 2018; 42: 343-351 [DOI: 29380024 DOI: 10.1007/s00266-018-1091-4]

Ziaide G, Karam D. Emulsified fat and nanofat for the treatment of dark circles. Dermatol Ther 2020; 33: e14100 [PMID: 32725706 DOI: 10.1111/dht.14100]

Oh DS, Kim DH, Roh TS, Yun IS, Kim YS. Correction of Dark Coloration of the Lower Eyelid Skin with Nanofat Grafting. Arch Aesthetic Plast Surg 2014; 20: 92-96 [DOI: 10.14730/aaps.2014.20.2.92]

Tonnard P, Verpaele A, Carvass M. Fat Grafting for Facial Rejuvenation with Nanofat Grafts. Clin Plast Surg 2020; 47: 53-62 [PMID: 31739897 DOI: 10.1016/j.cps.2019.08.006]

Boureaux EJ. Fat grafting for facial rejuvenation. J Craniomaxillofac Surg 2019; 47: 374-380 [PMID: 29878136 DOI: 10.1016/j.jcms.2015.12.013]

Le TP, Peckinpaugh J, Naficy S, Amadi AJ. Effect of autologous fat injection on lower eyelid position. Ophthamol Plast Reconstr Surg 2014; 30: 504-507 [PMID: 24814272 DOI: 10.1097/IOP.0000000000000160]

Yuksek E, Spina M, Yazgan H. Role of Fat Grafting in Primary Rhinoplasty. Plast Reconstr Surg 2012; 130: 44-45 [DOI: 10.1097/PRS.0b013e3182478e8111]

Kao WP, Lin YN, Lin TY, Huang YH, Chou CK, Takahashi H, Shieh TY, Chang KP, Lee SS, Lai CS, Lin SD, Lin TM. Microautologous Fat Transplantation for Primary Augmentation Rhinoplasty: Long-Term Monitoring of 198 Asian Patients. Aesthet Surg J 2016; 36: 648-656 [PMID: 26764261 DOI: 10.1093/asjgjx025]

Segreto F, Marangi GF, Nobile C, Alessandri-Bonetti M, Gregory C, Cerbone V, Gratteri M, Caldara E, Tirindelli MC, Persichetti P. Use of platelet-rich plasma and modified nanofat grafting in infected ulcers: Technical refinements to improve regenerative and antimicrobial potential. Arch Plast Surg 2020; 47: 217-222 [PMID: 32453929 DOI: 10.5999/aps.2019.01571]

Fruh FS, Später T, Scheuer C, Menger MD, Laschke MW. Isolation of Murine Adipose Tissue-derived Microvascular Fragments as Vascularization Units for Tissue Engineering. J Vis Exp 2017 [PMID: 28518106 DOI: 10.3791/55721]

Hoying JB, Boswell CA, Williams SK. Angiogenic potential of microvessel fragments established in three-dimensional collagen gels. In Vitro Cell Dev Biol Anim 1996; 32: 409-419 [PMID: 8856341 DOI: 10.1007/BF02723003]
Kamat P, Frueh FS, McLuckie M, Sanchez-Macedo N, Wolint P, Lindenblatt N, Plock JA, Calcagni M, Buschmann J. Adipose tissue and the vascularization of biomaterials: Stem cells, microvascular fragments and nanofat-a review. Cytotherapy 2020; 22: 400-411 [PMID: 32507607 DOI: 10.1016/j.jcyt.2020.03.433]

Sun XT, Ding YT, Yan XG, Wu LY, Li Q, Cheng N, Qiu YD, Zhang MY. Angiogenic synergistic effect of basic fibroblast growth factor and vascular endothelial growth factor in an in vitro quantitative microcarrier-based three-dimensional fibrin angiogenesis system. World J Gastroenterol 2004; 10: 2524-2528 [PMID: 15300897 DOI: 10.3748/wjg.v10.i17.2524]

Frueh FS, Spitzer T, Lindenblatt N, Calcagni M, Giovanoli P, Scheuer C, Menger MD, Laschke MW. Adipose Tissue-Derived Microvascular Fragments Improve Vascularization, Lymphangiogenesis, and Integration of Dermal Skin Substitutes. J Invest Dermatol 2017; 137: 217-227 [PMID: 27574793 DOI: 10.1016/j.jid.2016.08.010]

Laschke MW, Menger MD. Adipose tissue-derived microvascular fragments: natural vascularization units for regenerative medicine. Trends Biotechnol 2015; 33: 442-448 [PMID: 26137863 DOI: 10.1016/j.tibtech.2015.06.001]

Klein D, Weisshardt P, Kleff V, Jastrow H, Jakob HG, Ergün S. Vascular wall-resident CD44+ multipotent stem cells give rise to pericytes and smooth muscle cells and contribute to new vessel maturation. PLoS One 2011; 6: e20540 [PMID: 21637782 DOI: 10.1371/journal.pone.0020540]

Laschke MW, Spitzer T, Menger MD. Microvascular Fragments: More Than Just Natural Vascularization Units. Trends Biotechnol 2021; 39: 24-33 [PMID: 32593437 DOI: 10.1016/j.tibtech.2020.06.001]

McDaniel JS, Pilia M, Ward CL, Pollot BE, Rathbone CR. Characterization and multilineage potential of cells derived from isolated microvascular fragments. J Surg Res 2014; 192: 214-222 [PMID: 24690547 DOI: 10.1016/j.sjs.2014.05.047]

Shepherd BR, Chen HY, Smith CM, Gruiou G, Williams SK, Hoying JB. Rapid perfusion and network remodeling in a microvascular construct after implantation. Arterioscler Thromb Vasc Biol 2004; 24: 898-904 [PMID: 14988090 DOI: 10.1161/01.ATV.000014103.86943.1e]

Karschnia P, Scheuer C, Heß A, Spitzer T, Menger MD, Laschke MW. Erythropoietin promotes network formation of transplanted adipose tissue-derived microvascular fragments. Eur Cell Mater 2018; 35: 268-280 [PMID: 29761823 DOI: 10.22203/eCM.v035a19]

Nunes SS, Krishna M, Gerard CS, Dale JR, Maddie MA, Benton RL, Hoying JB. Angiogenic potential of microvessel fragments is independent of the tissue of origin and can be influenced by the cellular composition of the implants. Microcirculation 2010; 17: 557-567 [PMID: 21040121 DOI: 10.1111/j.1549-8719.2009.0052.x]

Pilia M, McDaniel JS, Guda T, Chen XK, Rhoads RP, Allen RE, Corona BT, Rathbone CR. Transplantation and perfusion of microvascular fragments in a rodent model of volumetric muscle loss injury. Eur Cell Mater 2014; 28: 11-23; discussion 23 [PMID: 25017641 DOI: 10.22203/eCM.v028a02]

Li MT, Ruehle MA, Stevens HY, Serves N, Willett JK, Karlhockeyanak S, Warren GL, Guldberg RE, Krishna L. Skeletal Myoblast-Seeded Vascularized Tissue Scaffolds in the Treatment of a Large Volumetric Muscle Defect in the Rat Biceps Femoris Muscle. Tissue Eng Part A 2017; 23: 989-1000 [PMID: 28372522 DOI: 10.1089/ten.TEA.2016.0523]

Orth M, Altmeinter MAB, Scheuer C, Braun BJ, Holstein JH, Eglin D, DE'Este M, Histing T, Laschke MW, Pohlemann T, Menger MD. Effects of locally applied adipose tissue-derived microvascular fragments by thermoresponsive hydrogel on bone healing. Acta Biomater 2018; 77: 201-211 [PMID: 30030175 DOI: 10.1016/j.actbio.2018.07.029]

Ruehle MA, Li MA, Cheng A, Krishnan L, Willett NJ, Guldberg RE. Decorin-supplemented collagen hydrogels for the co-delivery of bone morphogenetic protein-2 and microvascular fragments to a composite bone-muscle injury model with impaired vascularization. Acta Biomater 2019; 93: 210-221 [PMID: 30685477 DOI: 10.1016/j.actbio.2019.01.045]

Shepherd BR, Hoying JB, Williams SK. Microvascular transplantation after acute myocardial infarction. Tissue Eng 2007; 13: 2871-2879 [PMID: 17883324 DOI: 10.1089/ten.2007.0025]

Frueh FS, Spitzer T, Körbel C, Scheuer C, Simson AC, Lindenblatt N, Giovanoli P, Menger MD, Laschke MW. Prevascularization of dermal substitutes with adipose tissue-derived microvascular fragments enhances early skin grafting. Sci Rep 2018; 8: 10977 [PMID: 30030486 DOI: 10.1038/s41598-018-29252-6]

Spitzer T, Frueh FS, Menger MD, Laschke MW. Potentials and limitations of Integra® flowable wound matrix seeded with adipose tissue-derived microvascular fragments. Eur Cell Mater 2017; 33: 268-278 [PMID: 28378876 DOI: 10.22203/eCM.v033a20]

Hiscox AM, Stone AL, Limesand S, Hoying JB, Williams SK. An inlet-stabilizing implant constructed using a prevascularized construct. Tissue Eng Part A 2008; 14: 433-440 [PMID: 18333795 DOI: 10.1089/tea.2007.0099]

Saberianpour S, Heidarzadeh M, Geramanyeh MH, Hosseinikhani H, Rahbarhaziri R, Nouri M. Tissue engineering strategies for the induction of angiogenesis using biomaterials. J Biol Eng 2018; 12: 36 [PMID: 30603044 DOI: 10.1186/s13036-018-0133-4]

Rouwendaal J, Khademhosseini A. Vascularization and Angiogenesis in Tissue Engineering: Beyond Creating Static Networks. Trends Biotechnol 2016; 34: 733-745 [PMID: 27032730 DOI: 10.1016/j.tibtech.2016.03.002]

Spitzer T, Ampofo E, Menger MD, Laschke MW. Combining Vascularization Strategies in Tissue
Adipose tissue-derived MSCs

Engineering: The Faster Road to Success? Front Bioeng Biotechnol 2020; 8: 592095 [PMID: 33364230 DOI: 10.3389/fbioe.2020.592095]

175 Bora P, Majumdar AS. Adipose tissue-derived stromal vascular fraction in regenerative medicine: a brief review on biology and translation. Stem Cell Res Ther 2017; 8: 145 [PMID: 28619097 DOI: 10.1186/s13287-017-0596-z]

176 Muller S, Adler I, Creff J, Leminger H, Achard P, Castella L, Sensebli L, Carrière A, Deschaseaux F. Human adipose stromal-vascular fraction self-organizes to form vascularized adipose tissue in 3D cultures. Sci Rep 2019; 9: 7250 [PMID: 31076601 DOI: 10.1038/s41598-019-43624-6]

177 Vezzani R, Shaw I, Lesme H, Yong L, Khan N, Tremolada C, Paulte B. Higher Pericyte Content and Secretory Activity of Microfragmented Human Adipose Tissue Compared to Enzymatically Derived Stromal Vascular Fraction. Stem Cells Transl Med 2018; 7: 876-886 [PMID: 30255987 DOI: 10.1002/sctm.18-0051]

178 Alexander RW. Understanding Adipose-derived Stromal Vascular Fraction (AD-SVF) Cell Biology and Use on the Basis of Cellular, Chemical, Structural and Paracrine Components: A Concise Review. Journal of Proliferation Therapy. 2012; 4: e855-e869

179 Ramakrishnan VM, Boyd NL. The Adipose Stromal Vascular Fraction as a Complex Cellular Source for Tissue Engineering Applications. Tissue Eng Part B Rev 2018; 24: 289-299 [PMID: 28316259 DOI: 10.1089/ten.TEB.2017.0061]

180 Zimmerlin L, Donnenberg VS, Pfeifer ME, Meyer EM, Pernaut R, Rubin JP, Donnenberg AD. Stromal vascular progenitors in adult human adipose tissue. Cytotherapy 2010; 12: 22-30 [PMID: 19852056 DOI: 10.1002/cyt.v.a.20813]

181 Roh JD, Sawh-Martinez R, Brennan MP, Jay SM, Devine L, Rao DA, Yi T, Mirenksy TL, Nalbandian A, Udelsman B, Hibino N, Shinoaka T, Saltzman WM, Snyder E, Kyriakides TR, Pober JS, Breuer CK. Tissue-engineered vascular grafts transform into mature blood vessels via an inflammation-mediated process of vascular remodeling. Proc Natl Acad Sci USA 2010; 107: 4669-4674 [PMID: 20209749 DOI: 10.1073/pnas.0911465107]

182 Cawthorn WP, Scheller EL, MacDougall OA. Adipose tissue stem cells meet preadipocyte commitment: going back to the future. J Lipid Res 2012; 53: 227-246 [PMID: 22140268 DOI: 10.1194/jlr.R021089]

183 Zannettino AC, Paton S, Arthur A, Khor F, Itsescu S, Gimble JM, Grontos S. Multipotential human adipose-derived stromal stem cells exhibit a perivascular phenotype in vitro and in vivo. J Cell Physiol 2008; 214: 413-421 [PMID: 17654479 DOI: 10.1002/jcp.21210]

184 Maleki M, Ghanbarvar F, Reza Behvarz M, Ejtemaei M, Ghadirkhomi E. Comparison of mesenchymal stem cell markers in multiple human adult stem cells. Int J Stem Cells 2014; 7: 118-126 [PMID: 25473449 DOI: 10.15283/ijsc.2014.7.2.118]

185 Rojewski MT, Weber BM, Schrenzenmeier H. Phenotypic Characterization of Mesenchymal Stem Cells from Various Tissues. Transfus Med Hemother 2008; 35: 168-184 [PMID: 21547115 DOI: 10.1159/000120913]

186 Thillerîtrechea P, Lohširivat W, Poungpairoj P, Tantithavorn V, Onnamlo N. Extensive Characterization of Mesenchymal Stem Cell Marker Expression on Freshly Isolated and In Vitro Expanded Human Adipose-Derived Stem Cells from Breast Cancer Patients. Stem Cells Int 2020; 2020:8237197 [PMID: 32655648 DOI: 10.1155/2020/8237197]

187 Arenowitz JA, Lockhart RA, Hakakian CS. A Method for Isolation of Stromal Vascular Fraction Cells in a Clinically Relevant Time Frame. Methods Mol Biol 2018; 1773: 11-19 [PMID: 29687377 DOI: 10.1007/978-1-4939-7799-4_2]

188 van Dongen JA, Harmesen MC, Stevens HP. Isolation of Stromal Vascular Fraction by Fractionation of Adipose Tissue. Methods Mol Biol 2019; 1993: 91-103 [PMID: 31148081 DOI: 10.1007/978-1-4939-9473-1_8]

189 Senesi L, De Francesco F, Farinelli L, Manzotti S, Gagliardi G, Papulia GF, Riccio M, Gigante A. Mechanical and Enzymatic Procedures to Isolate the Stromal Vascular Fraction From Adipose Tissue: Preliminary Results. Front Cell Dev Biol 2019; 7: 88 [PMID: 31231649 DOI: 10.3389/fcell.2019.00088]

190 Condé-Green A, Kotamarti VS, Sherman LS, Keith JD, Lee ES, Granick MS, Rameshwar P. Shift toward Mechanical Isolation of Adipose-Derived Stromal Vascular Fraction: Review of Upcoming Techniques. Plast Reconstr Surg Glob Open 2016; 4: e1017 [PMID: 27757339 DOI: 10.1097/GOX.0000000000001017]

191 SundarRaj S, Deshmukh A, Priya N, Krishnan VS, Cherat M, Majumdar AS. Development of a System and Method for Automated Isolation of Stromal Vascular Fraction from Adipose Tissue Liposapirate. Stem Cells Int 2015; 2015: 109353 [PMID: 26167182 DOI: 10.1186/1599-3130-2015-109353]

192 Rodriguez J, Pratta AS, Abbassi N, Fabre H, Rodriguez F, Debard C, Adobati J, Boucher F, Mallein-Gerin F, Aucamps F, Damour O, Mojallal A. Evaluation of Three Devices for the Isolation of the Stromal Vascular Fraction from Adipose Tissue and for ASC Culture: A Comparative Study. Stem Cells Int 2017; 2017: 9289213 [PMID: 28321259 DOI: 10.1155/2017/9289213]

193 Doi K, Tanaka S, Iida H, Eto H, Kato H, Aoi N, Kuno S, Hiroti T, Yoshimura K. Stromal vascular fraction isolated from lipo-aspirates using an automated processing system: bench and bed analysis. J Tissue Regen Med 2013; 7: 864-870 [PMID: 22438241 DOI: 10.1002/term.1478]

194 Matsumoto D, Sato K, Gonda K, Takaki Y, Shigeura T, Sato T, Aiba-Kojima E, Iizuka F, Inoue K, Suga H, Yoshimura K. Cell-assisted lipotransfer: supportive use of human adipose-derived cells for soft tissue augmentation with lipoinjection. Tissue Eng 2006; 12: 3375-3382 [PMID: 17518674 DOI: 10.1089/ten.2006.12.3375]
Wu AY, Zou M, Mizuno H, Huang J, Futrell JW, Katz AJ, Benhaim P, Lorenz HP, Hedrick MH. Multilineage cells from human adipose tissue: implications for cell-based therapies. *Tissue Eng* 2001; 7: 211-228. [PMID: 11304456 DOI: 10.1089/10763270130062839]

Riis S, Zachar V, Boucher S, Vemuri MC, Pennisi CP, Fink T. Critical steps in the isolation and expansion of adipose-derived stem cells for translational therapy. *Expert Rev Mol Med* 2015; 17: e11. [PMID: 26052798 DOI: 10.1017/erm.2015.10]

Shah FS, Wu X, Dietrich M, Rood J, Gimble JM. A non-enzymatic method for isolating human adipose tissue-derived stromal stem cells. *Cytotherapy* 2013; 15: 979-985. [PMID: 23725689 DOI: 10.1016/j.jcyt.2013.04.001]

Aronowitz JA, Lockhart RA, Hakakian CS. Mechanical versus enzymatic isolation of stromal vascular fraction cells from adipose tissue. *Springerplus* 2015; 4: 713. [PMID: 26636001 DOI: 10.1186/s40064-015-1509-2]

Kamy S, Vollrath JT, Relja B. Cytokines in Inflammatory Disease. *Int J Mol Sci* 2019; 20: [PMID: 31795299 DOI: 10.3390/ijms20236008]

Bowles AC, Wise RM, Gerstein BY, Thomas RC, Ogelman R, Febo I, Bunnett NA. Immunomodulatory Effects of Adipose Stromal Vascular Fraction Cells Promote Alternative Activation Macrophages to Repair Tissue Damage. *Stem Cells* 2017; 35: 2198-2207. [PMID: 28801931 DOI: 10.1002/stem.2689]

Dong Z, Peng Z, Chang Q, Lu F. The survival condition and immunoregulatory function of adipose stromal vascular fraction (SVF) in the early stage of nonvascularized adipose tissue transplantation. *PLoS One* 2013; 8: e80364. [PMID: 24266375 DOI: 10.1371/journal.pone.0080364]

Krawiec JT, Liao HT, Kwan LL, D’Amore A, Weinbaum JS, Rubin JP, Wagner WR, Vorp DA. Evaluation of the stromal vascular fraction of adipose tissue as the basis for a stem cell-based tissue-engineered vascular graft. *J Vasc Surg* 2017; 66: 883-890.e1. [PMID: 28017585 DOI: 10.1016/j.jvs.2016.09.034]

Karina, Samudra MF, Rosadi I, Afni I, Widyastuti T, Sobariah M, Puspitasari RL, Rosliana I, Tunngadewi TI. Combination of the stromal vascular fraction and platelet-rich plasma accelerates the wound healing process: pre-clinical study in a Sprague-Dawley rat model. *Stem Cell Invest* 2019; 6: 18. [PMID: 31463311 DOI: 10.21037/sci.2019.06.08]

Tohita M, Tajima S, Mizuno H. Adipose tissue-derived mesenchymal stem cells and stem cell-rich plasma: stem cell transplantation methods that enhance stemness. *Stem Cell Res Ther* 2015; 6: 215. [PMID: 26541973 DOI: 10.1186/s13287-015-0217-8]

Rigotti G, Charles-de-Sá L, Gontijo-de-Amorim NF, Takiya CM, Amable PR, Borojevic R, Benati D, Bernardi P, Sabrati A. Expanded Stem Cells, Stromal-Vascular Fraction, and Platelet-Rich Plasma Enriched Fat: Comparing Results of Different Facial Rejuvenation Approaches in a Clinical Trial. *Aesthet Surg J* 2016; 36: 261-270. [PMID: 26879294 DOI: 10.1093/asj/sjv231]

Butt G, Hassain I, Ahmad FJ, Choudhery MS. Stromal vascular fraction-enriched platelet-rich plasma therapy reverses the effects of androgenetic alopecia. *J Cosmet Dermatol* 2020; 19: 1078-1085. [PMID: 31541565 DOI: 10.1111/jocd.13149]

Van Pham P, Hong-Thien Bui K, Quoc Ngo D, Tan Khuat L, Kim Phan N. Transplantation of Nonexpanded Adipose Stromal Vascular Fraction and Platelet-Rich Plasma for Articular Cartilage Injury Treatment in Mice Model. *J Med Eng* 2013; 2013: 832396. [PMID: 27006923 DOI: 10.1155/2013/832396]

Gentile P, Orlandi A, Scioli MG, Di Pasquali C, Bocchini I, Cervelli V. Concise review: adipose-derived stromal vascular fraction cells and platelet-rich plasma: basic and clinical implications for tissue engineering therapies in regenerative surgery. *Stem Cells Transl Med* 2012; 1: 230-236. [PMID: 23197782 DOI: 10.5966/scitm.2011-0054]

Jurgens WJ, Kroeze RJ, Zandieh-Doulabi B, van Dijk A, Renders GA, Smit TH, van Milligen FJ, Karina, Samudra MF, Rosadi I, Afni I, Widyastuti T, Sobariah M, Puspitasari RL, Rosliana I, Tunngadewi TI. One-step surgical procedure for the treatment of osteochondral defects with adipose-derived stem cells in a caprine knee defect: a pilot study. *Biorex Open Access* 2013; 2: 315-325. [PMID: 23914338 DOI: 10.1089/biores.2013.0024]

Zhou L, Xu L, Shen J, Song Q, Wu R, Ge Y, Xin H, Zou J, Wu J, Jia R. Preischemic Administration of Nonexpanded Adipose Stromal Vascular Fraction Attenuates Acute Renal Ischemia/Reperfusion Injury and Fibrosis. *Stem Cells Transl Med* 2016; 5: 1277-1288. [PMID: 27365485 DOI: 10.5966/scitm.2015-0223]

van Dijk A, Naaijkens BA, Jurgens WJ, Nallahk S, Sairras S, van der Pijl RJ, Vo K, Vonk AB, van Rossum AC, Paulus WJ, van Milligen FJ, Niessen HW. Reduction of infarct size by intravenous injection of uncultured adipose derived stromal cells in a rat model is dependent on the time point of application. *Stem Cell Res* 2011; 7: 219-229. [PMID: 21907165 DOI: 10.1016/j.scr.2011.06.003]

Sheng L, Yang M, Li H, Du Z, Yang Y, Li Q. Transplantation of adipose stem cells promotes neovascularization of random skin flaps. *Tohoku J Exp Med* 2011; 224: 229-234. [PMID: 21701129 DOI: 10.1620/tjem.224.229]

Qiu X, Fandell TM, Ferretti L, Albermes M, Orabi H, Zhang H, Lin G, Lin CS, Schroeder T, Lue TF. Both immediate and delayed intraoperative injection of autologous adipose-derived stromal vascular fraction enhances recovery of erectile function in a rat model of cavernous nerve injury. *Eur Urol* 2012; 62: 720-727. [PMID: 22397847 DOI: 10.1016/j.eururo.2012.02.003]

Wu AY, Morrow DM. Autologous fat transfer with in-situ maturation (AIM): a novel and compliant method of adult mesenchymal stem cell therapy. *J Transl Med* 2013; 11: 136. [PMID: 23725573]
Sharma S et al. Adipose tissue-derived MSCs

DOI: 10.1186/1479-5876-11-136

Gentile P, Orlandi A, Scioli MG, Di Pasquali C, Bocchini I, Curcio CB, Floris M, Fiaschetti V, Floris R, Cervell V. A comparative translational study: the combined use of enhanced stromal vascular fraction and platelet-rich plasma improves fat grafting maintenance in breast reconstruction. Stem Cells Transl Med 2012; 1: 341-351 [PMID: 23197813 DOI: 10.5966/sctm.2011-0665]

Castella L, Planat-Beurard V, Laharrague P, Cousin B. Adipose-derived stromal cells: their identity and uses in clinical trials, an update. World J Stem Cells 2011; 3: 25-33 [PMID: 21607134 DOI: 10.4255/wjsc.v3.i2.25]

García-Olmo D, García-Arranz M, Herrerós D, Pascau I, Peiro C, Rodriguez-Montes JA. A phase I clinical trial of the treatment of Crohn's fistula by adipose mesenchymal stem cell transplantation. Dis Colon Rectum 2005; 48: 1416-1423 [PMID: 15933795 DOI: 10.1007/s10350-005-0052-6]

Atalay S, Coruh A, Deniz K. Stromal vascular fraction improves deep partial thickness burn wound healing. Burns 2014; 40: 1375-1383 [PMID: 24572074 DOI: 10.1016/j.burns.2014.01.023]

Zhao X, Guo J, Zhang F, Zhang J, Liu D, Hu W, Yin H, Jin L. Therapeutic application of adipose-derived stromal vascular fraction in diabetic foot. Stem Cell Res Ther 2020; 11: 394 [PMID: 32928305 DOI: 10.1186/s13287-020-01825-1]

Tzouvelekis A, Pasqualiari V, Koliakos G, Ntosios P, Boursos N, Dardzinska B, Giritzalis D, Antoniades A, Froudarakis M, Kolios G, Boursos D. A prospective, non-randomized, no placebo-controlled, phase Ib clinical trial to study the safety of the adipose derived stromal cells-stromal vascular fraction in idiopathic pulmonary fibrosis. J Transl Med 2013; 11: 171 [PMID: 23855653 DOI: 10.1186/1479-5876-11-171]

Gentile P, Sterodimas A. Adipose Stem Cells (ASCs) and Stromal Vascular Fraction (SVF) as a Potential Therapy in Combating (COVID-19)-Disease. Aging Dis 2020; 11: 465-469 [PMID: 32488692 DOI: 10.14336/AD.2020.0422]

Mazini L, Ezzouiti M, Malka G. Overview of current adipose-derived stem cell (ADSCs) processing involved in therapeutic advancements: flow chart and regulation updates before and after COVID-19. Stem Cell Res Ther 2021; 12: 1 [PMID: 33397467 DOI: 10.1186/s13287-020-02006-w]

Jeyaraman M, Ranjan R, Kumar R, Arora A, Chaudhary D, Ajay SS, Jain R. Cellular Therapy: Shafts of Light Emerging for COVID-19. Stem Cell Investig 2020; 7: 11 [PMID: 32695804 DOI: 10.21037/sci-2020-020]

D’Esposito V, Passaretti F, Perruolo G, Ambrosio MR, Valentino R, Oriente F, Raciti GA, Nigro C, Miele C, Sammartino G, Beguinot F, Formisano P. Platelet-Rich Plasma Increases Growth and Motility of Adipose Tissue-Derived Mesenchymal Cells and Controls Adipocyte Secretory Function. J Cell Biochem 2015; 116: 2408-2418 [PMID: 26012576 DOI: 10.1002/jcb.25235]

Kolle SF, Fischer-Nielsen A, Mathiasen AB, Elberg JJ, Oliveri RS, Glovinski PV, Kastrup J, Kirchhoff M, Rasmussen BS, Talman ML, Thomsen C, Dickmeiss E, Drzewiecki KT. Enrichment of adipocytes in adipose tissue improves fat grafting maintenance in breast reconstruction. J Orthop Surg Res 2020; 15: 137 [PMID: 32272946 DOI: 10.1186/s13073-020-01664-2]

Wu J, Yang Q, Wu S, Yuan W, Zha N. Adipose-Derived Stem Cell Exosomes Promoted Hair Regeneration. Tissue Eng Regen Med 2021; 18: 685-691 [PMID: 34173219 DOI: 10.1007/s13770-021-00347-y]

Ogawa R, Tanaka C, Sato M, Nagasahi K, Sugimura K, Nakamura Y, Aoki N. Adipocyte-derived microvesicles contain RNA that is transported into macrophages and might be secreted into blood circulation. Biochim Biophys Acta 2010; 398: 723-729 [PMID: 20621060 DOI: 10.1016/j.bbr.2010.07.008]

Mou S, Zhou M, Li Y, Wang J, Yuan Q, Xiao P, Sun J, Wang Z. Extracellular Vesicles from Human Adipose-Derived Stem Cells for the Improvement of Angiogenesis and Fat-Grafting Application. Plast Reconstr Surg 2019; 144: 869-880 [PMID: 31568294 DOI: 10.1097/PRS.0000000000001806]

Han Y, Ren J, Bai Y, Pei X, Han Y. Exosomes from hypoxia-treated human adipose-derived mesenchymal stem cells enhance angiogenesis through VEGF/VEGF-R. Int J Biochem Cell Biol 2019; 109: 59-68 [PMID: 30710751 DOI: 10.1016/j.biocel.2019.01.017]

Wang J, Yi Y, Zhu Y, Wang Z, Wu S, Zhang J, Hu X, Nie J. Effects of adipose-derived stem cell released exosomes on wound healing in diabetic mice. Zhongguo Xiu Fu Chong Jian Wai Ke Za Zhi 2020; 44: 124-131 [PMID: 31393247 DOI: 10.7507/1002-1892.201903058]

Li X, Xie X, Lian W, Shi R, Han S, Zhang H, Lu L, Li M. Exosomes from adipose-derived stem cells overexpressing Nr2f2 accelerate cutaneous wound healing by promoting vascularization in a diabetic foot ulcer rat model. Exp Mol Med 2018; 50: 1-14 [PMID: 29651102 DOI: 10.1038/s12276-018-0058-5]

Shafei S, Khanmohammadi M, Heidari R, Ghanbari H, Taghdiri Nooshabadi V, Farzamfar S, Akbariromi M, Sanikhihi NS, Absalan M, Tavoosidana G. Exosome loaded alginate hydrogel promotes tissue regeneration in full-thickness skin wounds: An in vivo study. J Biomed Mater Res A 2020; 108: 545-556 [PMID: 31702867 DOI: 10.1002/jbm.a.36835]

Taverna S, Pucci M, Alessandro R. Extracellular vesicles: small bricks for tissue repair/regeneration. Ann Transl Med 2017; 5: 83 [PMID: 28275628 DOI: 10.21037/atm.2017.01.53]
Wang L, Hu L, Zhou X, Xiong Z, Zhang C, Shehada HMA, Hu B, Song J, Chen L. Exosomes secreted by human adipose mesenchymal stem cells promote scarless cutaneous repair by regulating extracellular matrix remodelling. Sci Rep 2017; 7: 13321 [PMID: 29042658 DOI: 10.1038/s41598-017-12919-x].

Kim WS, Park BS, Kim HK, Park JS, Kim KJ, Choi JS, Chung SJ, Kim DD, Sung JH. Evidence supporting antioxidant action of adipose-derived stem cells protection of human dermal fibroblasts from oxidative stress. J Dermatol Sci 2008; 49: 133-142 [PMID: 17870415 DOI: 10.1016/j.jdermsci.2007.08.004].

Zhao H, Shang Q, Pan Z, Bai Y, Li Z, Zhang H, Zhang Q, Guo C, Zhang L, Wang Q. Exosomes From Adipose-Derived Stem Cells Attenuate Adipose Inflammation and Obesity Through Polarizing M2 Macrophages and Beiging in White Adipose Tissue. Diabetes 2018; 67: 235-247 [PMID: 29133512 DOI: 10.2337/db17-0356].

Li W, Liu Y, Zhang P, Tang Y, Zhou M, Jiang W, Zhang G, Zhou Y. Tissue-Engineered Bone Immobilized with Human Adipose Stem Cells-Derived Exosomes Promotes Bone Regeneration. ACS Appl Mater Interfaces 2018; 10: 5240-5254 [PMID: 29359912 DOI: 10.1021/acsami.7b17620].

Chen S, Tang Y, Liu Y, Zhang P, Lv L, Zhang X, Jia L, Zhou Y. Exosomes derived from miR-375-overexpressing human adipose mesenchymal stem cells promote bone regeneration. Cell Prolif 2019; 52: e12669 [PMID: 31380594 DOI: 10.1111/cpr.12669].

Lu Z, Chen Y, Dunstan C, Roohani-Esfahani S, Zeerag H. Priming Adipose Stem Cells with Tumor Necrosis Factor-Alpha Preconditioning Potentiates Their Exosome Efficacy for Bone Regeneration. Tissue Eng Part A 2017; 23: 1212-1220 [PMID: 28346708 DOI: 10.1089/ten.tea.2016.0548].

Pers YM, Quentin J, Feirreira R, Abdellaoui N, Erkilic N, Cren M, Dufourcq-Lopez E, Pullig O, Nöth U, Jorgensen C, Louis-Plence P. Injection of Adipose-Derived Stromal Cells in the Knee of Patients with Severe Osteoarthritis has a Systemic Effect and Promotes an Anti-Inflammatory Phenotype of Circulating Immune Cells. Theranostics 2018; 8: 5519-5528 [PMID: 30555561 DOI: 10.7150/thno.27674].

Riau AK, Ong HS, Yam GHF, Mehta JS. Sustained Delivery System for Stem Cell-Derived Exosomes. Front Pharmacol 2019; 10: 1368 [PMID: 31798457 DOI: 10.3389/fphar.2019.01368].
