Mesenchymal Stromal Cells and Micro Fragmented Adipose Tissue: New Horizons of Effectiveness of Lipogems

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Abstract

In the last three years there has been an explosion of published in vivo and clinical data discussing micro fragmented adipose tissue and its regenerative properties. In fact, the paracrine activity of micro fragmented adipose tissue has been well described, showing the importance of the maintenance of an intact microvascular environment of fat tissue during the processing in order to guarantee cellular activation and long-term viability there by maintaining their signaling capacity in order to effectively modulate inflammation and immune response.

Lipogems®, patented in 2010 and clinically available since 2013, is a novel technology that reproducibly and simply obtains, micro fragmented adipose tissue with enhanced regenerative properties with somewhat astonishing results that are now confirmed from recent clinical trials.

In this review we have summarized all the new evidence published, focusing on possible new clinical indications from intriguing new in vivo studies.

Keywords: Lipogems®; Medicinal signaling cells; Mesenchymal stromal cells; Micro fragmented adipose tissue; Pain; Plastic surgery; Regenerative medicine

Introduction

Since its first clinical use in 1995 [1], we have seen an exponential increase of the literature pertaining to Mesenchymal Stromal Cells (MSCs) during the last twenty years. Their activity has been deeply characterized in several articles [2] both in humans and in animals [3]. It has been also proven in vitro that MSCs can differentiate into different adult cells according to the environment in which they have been injected [1].

MSCs have been harvested from several different tissues [4,5] (and using a variety of different methods), even though the most easily accessible site is from adipose tissue [6,7]. Importantly, MSCs derived from either/any site have common biological characteristics, with some biological differences in vitro but no discernable differences in activity or behavior in clinical practice [8]. The most promising source of MSCs, adipose tissue, presents a significantly higher concentration of MSCs compared to bone marrow (1% versus >0.01%), and also other sources such as umbilical cord, dental pulp or menstrual blood [9]. Furthermore harvesting from adipose tissue is less invasive than bone marrow with less risk of severe complications.

Considering the multipotent properties of MSCs derived from adipose tissue [10-16], they have been proposed as potential reparative and regenerative candidates for the treatment of diseases where an angiogenic, anti-inflammatory, anti-fibrotic or anti-apoptotic mechanism was a critical component needed to treat the pathophysiology of the disease (covers most major disease processes potentially).

A recent review [17] analyzed all of the currently available methods of isolation of MSCs from adipose tissue and there in vitro and in vivo applications. Autologous adipose tissue has been used as intact lipo-aspirate, enzymatically derived Stromal Vascular Fraction (SVF) or Micro Fragmented Fat (MFAT). Vezzani et al., [18] has demonstrated that the “regenerative properties” of these three methods are quite different; with the MFAT preparations being significantly more productive in the release of grow factor and cytokines concomitant with notably superior effects on repairing tissues, induction and modification of immunomodulatory activity and in supporting vascular angiogenesis.

These results may be attributed to the importance of maintaining the perivascular environment/niche, with MSCs being a critical component of this and of perivascular origin. The pericytes and other perivascular cells have a key role in the regenerative medicine as they can be reprogrammed into “regenerative cells” that act as immunomodulatory and anti-inflammatory cells to restore and repair damaged tissue [19]. Pericytes are multi-potent structural cells localized within the vascular basement membrane and present on all parts of the vascular system or ‘stromal vascular fraction’ of adipose tissue. Fascinatingly, when activated during tissue damage or inflammation, they transform into MSCs with the capability to create a pro-regenerative environment and have the ability to differentiate into mesodermal lineage terminal cells [20].

The literature also provides evidence that tissue or cellular treatments beyond a ‘minimal’ mechanical manipulation of adipose tissue (for example adipose fat or other derived MSC containing cell mixture), such as enzymatic dissociation, leads to a different gene expression pattern [21] and exosome content of the MSCs [22]. Currently,
researchers are studying the cytokine retention and delivery capacity and safety profile of exosomes derived from MSCs (MEX) [23,24]. Even though the MEX regulatory properties are well demonstrated, the most important being that their activity is highly variable depending upon the tissue they are isolated from-making tissue sourcing an important consideration before extraction, banking or use.

A more detailed characterization of MEX will be of high value for further research in the field of regenerative medicine and in the future development of MSC-based drug delivery protocols [24].

One example where MSCs derived from adipose tissues have been shown to be effective is in the treatment of complex-non-healing wounds. In vivo studies in mice demonstrated a significantly better healing response and final outcome compared with controls treated using a standard protocol [25]. As with all the experimental data, large differences in the experimental findings are seen when comparing the capability of MSCs derived from different sources. Hence, it is important to adopt, both in experimental models and in a clinical settings, a fully standardized Standard Operating Procedures (SOPs) to harvest and to manipulate adipose tissue that will minimize the damage to the perivascular environment and maximize the regenerative effect concomitantly reducing the risk of contamination [26]. Lipogems® technology is a closed sterile system that allows the collection of MSCs from MFAT without contamination.

In this review we present an update of previously published data demonstrating a clear rationale, based on in vivo results and clinical studies produced using this technology and justifying it as an important tool for regenerative medicine [27].

**Lipogems®: Technology and New Insights in its Activity**

Lipogems® is a new device that efficiently harvests, processes and transfers adipose tissue with unique characteristics including long-term expression and secretion of physiologically active concentrations of angiogenic, immunomodulatory, anti-inflammatory, anti-apoptotic and anti-fibrotic secretome [25].

The resulting Lipogems product is composed of small specifically sized ‘adipose’ clusters of adipocytes maintaining an intact perivascular environment and a pericytes’ activation through ball-bearing-induced mechanical shock. Lipogems®, creates a minimally manipulated fat-derived product according to and in line with the regulations set forth by US Food and Drug Administration (FDA).

It has received FDA clearance as a class II medical device for processing of autologous adipose tissue. According FDA we could defined Lipogems® as (1) autologous; (2) minimally manipulated; (3) intended for homologous use; (4) enzyme-free; (5) not dependent on the metabolic activity of the cells for its primary function; (6) used in the same surgical procedure and (7) not combined with anything other than saline.

The harvest technique has been already described in several other articles [27,28], and here it will be described briefly.

The procedure is divided into three phases:

• The first phase is the harvest and process of a small quantity of fat tissue after local anesthesia and Klein solution subcutaneous fat tissue injection (mini-liposuction). Klein solution is a tumescent anesthesia that uses large volumes of saline with local anesthetic (usually lidocaine at 0.05-0.1% concentration) and epinephrine (1:500,000-1:1,000,000).

• The second phase is carried out (Figure 1) in a closed full immersion low pressure cylindrical system in order to obtain two parts: a micro fragmentated fat tissue (fat tissue clusters of 0.3-0.7 mm) and a fluid containing oil, all the excreted proinflammatory cytokines and dead/dying or apoptotic cells, that are subsequently eliminated. The system obtains the micro-fragmented fat only through mechanical forces without any disruption of the integrity of the fat tissue and/or stromal vascular fraction and with “activation” of pericytes and within the microvascular environment of MSCs [29].

• In the third phase milliliter quantities of Lipogems® are injected into the area to be treated, resulting in a transplant of MFAT containing a complex spectrum of bioactive molecules. As previously described, Lipogems® has a significantly higher content and concentration of MSCs and exosomes compared to other fat-derived products including those derived from enzymatic methods [as there is no any digestions of extracellular matrix, the exosomes are not altered and the adipose structural niche is not damaged, but reduced in its size increasing the surface area of the active tissue] [22,31-35].

The whole process is performed in less than twenty minutes in a sterile environment and with minimal manipulation.

Through this autologous adipose tissue graft we have created a fast, reproducible and regulatory compliant technology that effectively supports the natural healing of joints or tissues, through its capacity to cushion, lubricate and ‘stick’ or remain viable in the area of interest for significant length of time via a self-regenerative mechanism.

In addition, as already described in our previous review [27], and as a secondary potential mechanism of support, MFAT has been demonstrated to be a useful source of growth factors and cytokines that may assist natural repair and regeneration of tissues due to its content of mesenchymal stromal cells and high paracrine activity.

Interestingly, very recently, Nava et al., demonstrated that micro-fragmented fat tissue displayed not only a higher, but also a longer anti-inflammatory activity compared to normal fat tissue graft [36]. In fact, its anti-inflammatory activity persists for more than 1 month.
as defined by measurement of paracrine activity and its ability to inhibit monocyte and macrophage migration.

Probably, this phenomenon relates to the long term survival of MSCs within the in vivo environment where they are injected, due to the optimized Lipogems® adipoclasts functioning to stabilize their survival and allowing targeted therapeutic responses following mechanical activation of bound pericytes following Lipogems extraction [37].

These results underline that micro-fragmented fat could represent not only a regenerative technique, but an autologous long-lasting drug able to modulate chronic inflammation in clinical conditions in which inflammation can be difficult to control, such as chronic joint inflammation or chronic systemic inflammation (see last chapter).

**Clinical Applications of Micro Fragmented Adipose Tissue**

In this chapter we will summarize the diseases where micro-fragmented fat tissue has demonstrated its effectiveness in clinical practice. In fact, in several clinical applications there a growing interest has been shown in the literature regarding clinical effectiveness and safety over the last decade [27].

**Plastic surgery**

Initially, MFAT has been used for plastic surgery not only for lipofilling but also associated with traditional aesthetic surgery where a fat tissue graft is needed, such as blepharoplasty, facelift or breast augmentation. In this setting Lipogems® demonstrated not only an “augmentation” activity, but also a real anti-inflammatory activity with better short- (reduced postoperative pain and swelling) and long-term results [27].

In a randomized controlled trial [38] the effectiveness of micro-fragmented fat tissue versus conventional treatment has been investigated in a series of patients destined for diabetic foot amputation. At 6 months, 80%, of the group treated with Lipogems MFAT healed compared to 46% of those in the group of conventional treatment. This randomized controlled trial underlined the potential wound healing and tissue regenerative properties of MFAT injection.

Lipogems is also frequently used also for management of treatment of ulcers and systemic sclerosis, -as have other types of fat tissue graft [39,40], but with significantly better outcomes and possible indications also for the treatment of chronic non-healing wounds as recently demonstrated [41]. These authors demonstrated, using an in vitro model, the pathophysiological basis that defines how micro-fragmented fat tissue, such as Lipogems®, is effective in treating chronic non-healing wounds, identifying a vascular stabilization activity (with inhibition of endothelial expression) and negative modulation of macrophage migration.

**General and urological gynecological surgery**

MFAT was demonstrated to be effective after a single injection in a case series of 15 patients with perianal fistulas with Crohn’s disease [42]. Patients were treated with a single local injection of MFAT. After 6 weeks 10 patients showed a complete clinical and instrumental remission, 4 patients slightly improved and only 1 patient did not have any improvement.

In addition, effective relief of symptoms has also been reported in case series of patients with for both anal [42,43] and urinary [44] incontinence.

Lipogems® has also been successfully demonstrated to be effective in the relief of symptoms of postmenopausal genitourinary atrophy [45] in three patients who remain symptom free and showing both clinical and histopathological complete recovery for three years. Biopsy of the area demonstrated a completely normal looking epithelial mucosa consistent with a premenopausal condition. The authors are currently confirming this data in a larger cohort of women (submitted paper).

**Maxillofacial surgery**

Also in maxilla surgery MFAT has been found effective in regenerative bone atrophy [46] and in 120 patients with double-jaw procedures where soft tissue had to be restored [47]. In these patients microfragmented fat tissue demonstrated to produce less local inflammatory reaction compared to usual fat tissue graft.

**Orthopedic surgery**

In 2019 a group of researchers [48] used rabbit model osteoarthritic, to investigate the pattern of cell migration of three different injected fat-derived materials: expanded-Adipose Stromal Cells (ASCs) and adipose niches after enzymatic (Stromal Vascular Fraction - SVF) and mechanical processes (Micro Fragmented Fat Tissue - MFAT). They found a significantly improved long-term (30 days) migration of MFAT toward cartilage while the cells prepared using the other two methods migrated toward the synovia. Furthermore, Paolella et al. [49] demonstrated that freshly prepared MFAT had a greater effect in reducing inflammation also in synovial tissue compared to the adipose-derived mesenchymal stromal cells obtained from the same samples of MFAT.

As already underlined in our previous review, Lipogems® is currently used within the clinic in several orthopedic contraindications, where intra-articular injections are used to for the management of knee, hip and shoulder osteoarticular degeneration (more than 35000 patients over the world). Initiated more than three years ago, these results have now been confirmed in several case series [50,51], and prospective clinical studies with follow up [52,53], and the “regenerative” pathophysiological mechanism of Lipogems®, when injected in the joint, have now been more carefully elucidated [54,55].

MFAT has also been demonstrated effective also in four patients with osteochondral lesions of the talus [56] who underwent to arthroscopic injection of MFAT of the osteochondral lesion after a first phase of microperforation of the cartilage. After six months all patients have a recovery of the lesion.

Jannelli E et al., [57], in an interesting position paper, proposed micro-fragmented fat tissue transplantation to treat delamination and first and second-degree chondral lesions also in knee arthroscopic surgery in order to regenerate cartilage and prevent severe grade of osteoarthrits. Schiavone Panni et al., [50] underlined in a recent case series in fifty-two patients also that MFAT treatment as an adjunct to arthroscopic debridement could be very effective also in patient with early knee osteoarthrits. They found a significative improvement after one year both of the pain scores and of the functional activity.

In a retrospective study on 76 patients (and 106 knees) no statistical difference in functional outcome or quality of life was seen
between intra articular MFAT and bone marrow aspirate concentrate knee injection in osteoarthritises patients [58]. MFAT was easier and less invasive to harvest compared to bone marrow and with no any side effects.

Finally, there are some ongoing randomized clinical trials [59] that are currently investigating effectiveness of micro-fragmented fat tissue injection compared to other techniques (clinicaltrials.gov NCT03117608, NCT03379168 and NCT03922490).

Recently, MFAT was demonstrated effective also in severe osteoarthritis in dogs. After 6 months 88% of dogs treated have an improvement of osteoarthritises signs and 92% of owners declared a significant improvement of pain and functional activity of their dogs [60]

**New Possible Indications for the Future**

In this chapter we will review findings from investigations in animal together with other clinical data from further regenerative techniques that could suggest additional and interesting potential uses for micro-fragmented fat.

**Regenerative medicine and axial back pain**

Chronic inflammatory flare is also one of the main pathophysiological processes that can cause back pain [61]. Unfortunately, chronic inflammation of the spinal column’s joints (i.e. sacroiliac and facet joint) or of the intervertebral disc is difficult to treat with current drugs and a mechanism-based diagnosis is mandatory [62]. As discussed above, it has been demonstrated that Mesenchymal Stromal Cells (MSCs) contained in liposipirate have important anti-inflammatory and immunomodulatory properties, that can last for several months in Micro-Fragmented Fat Tissue (MFAT) and that could therefore potentially be useful for back pain. Furthermore, MFAT in cellular experiments has displayed more anti-inflammatory activity than enzymatically derived stromal vascular fraction as it contains more pericytes that are responsible for modulation of the immune (especially NK cells that are strictly related with chronic pain) and inflammatory response [18]. These results are the basis to justify the use of regenerative medicine for chronic pain not only to “regenerate” cartilage or joint structures but also (and maybe most critically) to modulate inflammation more effectively than current pharmacological agents.

Hence, regenerative medicine in Low Back Pain (LBP) could be used not only for its “regenerative” properties, but also for its long lasting “immunomodulatory and antiinflammatory” properties. This possibility has been confirmed in a recent metanalysis [63] that summarizes all recent clinical trials and observational studies where regenerative medicine has been proved effective in treating back pain for all pain generators (facet joints, sacroiliac joint, discogenic pain).

Currently the studies are more about Platelet Rich Plasma (PRP) and not about micro-fragmented fat. However, as recently discussed by American Society of Interventional Pain Physicians (ASIPP) guidelines [64], the level of scientific evidence still remains low, as randomized clinical trials are needed to demonstrate not only the effectiveness but also its effectiveness over time, as some studies have underlined that analgesia can last for several months after a single injection.

Finally, despite the promising results, we need more studies and data on who could benefit from regenerative medicine and who probably will not. Regenerative medicine could hence represent a mechanism-based treatment in many LBP patients with chronic inflammatory pain and it is only a matter of time before we get there.

**Promising animal studies**

Considering the great potential of immunomodulatory and regenerative activity of microfragmented fat tissue, in the last two years new important animal studies have suggested possible indications that need further exploration in order to attain clinical trial status.

Researchers from the Pasteur Institute [65] have analyzed in a Duchenne muscular dystrophy murine model the effect of micro fragmented fat injected in the muscle. Interestingly they found a reduction of fibrosis and necrosis with concomitant reduction of cytokines productions with final increase in muscle strength.

The same group has also investigated the antiinflammatory and immunomodulatory activity of micro fragmented fat, compared not only to placebo but also to liposipirate, if administered systemically in a murine sepsis model [66]. They found not only a significant (compared both to placebo and liposipirate) reduction of systemic inflammatory response in the presence of micro-fragmented fat tissue but also an important improvement of outcome with reduction in mortality.

Finally, MFAT has been recently studied as a possible interesting new scaffold for drug delivery especially in cancer treatments as a possible vehicle of Paclitaxel to increase chemotherapy effectiveness (delivered directly to the tumor) whilst reducing systemic side effects [67]. The authors demonstrated also that in the drain bag during Lipogems® preparation, it is possible to find single isolated cells that can be expanded and loaded with Paclitaxel to administer chemotherapy [68]

**Conclusion**

This review updates the incredible amount of new scientific data about micro-fragmented fat tissue published in the last three years [27]. In fact evidence that justifies the stronger regenerative effect of micro-fragmented fat tissue is growing exponentially. Furthermore, also the clinical evidence, especially for orthopedic surgery, is increasing with new prospective and randomized clinical trials that are not only confirming previous results but are indicating new possible solutions for diseases that have not at this moment valid therapeutic solutions.

Lipogems® appears to be really promising in this arena and it is already approved by the FDA as an autologous fat tissue graft. More randomized clinical studies are needed but the rational of its paracrine activity could justify its use also in new clinical settings.

Finally, in order to improve even more effectiveness of regenerative medicine, it is important to understand that MSCs have to be considered Medicinal Signaling Cells that could help the modulation of inflammatory flare (even chronic) and of the immune system in order to restore physiological conditions in the area affected [69].

**Conflicts of Interest**

Carlo Tremolada is a founder of Lipogems International SpA. Mark Slevin is a consultant of Lipogems. Massimo Allegri declines any conflict of interest relative this topic.
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