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Stem Cells and Spinal Fusion

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ABSTRACT

Background: This manuscript is a review of the literature investigating the use of mesenchymal stem cells (MSCs) being applied in the setting of spinal fusion surgery. We mention the rates of pseudarthrosis, discuss current bone grafting options, and examine the preclinical and clinical outcomes of utilizing MSCs to assist in successfully fusing the spine.

Methods: A thorough literature review was conducted to look at current and previous preclinical and clinical studies using stem cells for spinal fusion augmentation. Searches for PubMed/MEDLINE and ClinicalTrials.gov through January 2021 were conducted for literature mentioning stem cells and spinal fusion.

Results: All preclinical and clinical studies investigating MSC use in spinal fusion were examined. We found 19 preclinical and 17 clinical studies. The majority of studies, both preclinical and clinical, were heterogeneous in design, due to different osteoconductive scaffolds, cells, and techniques used. Preclinical studies showed promising outcomes in animal models when using appropriate osteoconductive scaffolds and factors for osteogenic differentiation. Similarly, clinical studies have promising outcomes, but differ in their methodologies, surgical techniques, and materials used, making it difficult to adequately compare between the studies.

Conclusion: MSCs may be a promising option to use to augment grafting for spinal fusion surgery. MSCs must be used with appropriate osteoconductive scaffolds. Cell-based allografts and the optimization of their use have yet to be fully elucidated. Further studies are necessary to determine the efficacy of MSCs with different osteoconductive scaffolds and growth/osteogenic differentiation factors.

Level of Evidence: 3.

Special Issue

Keywords: stem cells, mesenchymal stem cells, MSC, spinal fusion, biologics, regenerative medicine, autograft, bone graft, scaffold

INTRODUCTION

Spinal fusion surgery is one of the most common procedures performed in the United States, with over 1 million cases performed annually.¹ In 2020, it is still reported as one of the most costly procedures, with an average cost of \$120,000 per hospitalization.² Conditions treated with spinal fusion include various degenerative disorders, fractures, spinal tumors, and deformities of scoliosis, kyphosis, and more. Spinal fusions are performed when any structural or neurologic component of the spine is compromised, typically affecting abnormal motion, and likely producing pain and disability. Successful fusion involves new bone formation between 2 or more adjacent vertebrae, returning stability to the diseased segment of the spine.

A wide variety of spinal fusion procedures exist, with the anatomic location and pathology directing which surgical approach, stabilizing instrumenta-

tion, and procedure may maximize stability and rapid healing, while minimizing surgical trauma. As novel technologies progress influencing instrumentation, biomaterials, implants, and grafting techniques, new and less tissue-destructive approaches are being discovered/designed.

The incidence of pseudarthrosis, or nonunion, can be as high as approximately 25%–35% in spinal fusion surgery, which is highly dependent on the type of procedure, approach used, and patient factors, such as bone quality, health status, and comorbidities.³ This is an extremely high incidence for such a widely and commonly performed, as well as expensive, procedure. When there is a failure of bone formation, unsuccessful fusion leads to the following: pain, instability, implant failure, reoperation, patient stress, and drastically increased costs.⁴ Risk factors include cigarette smoking, age, female sex, excessive thoracolumbar kyphosis, and various

bone diseases, such as osteopenia and osteoporosis.⁵⁻⁸

Methods to prevent pseudarthrosis have become some of the most researched and invested-in aspects of spine surgery today. Traditional gold standard for bone grafting has been autologous bone harvest, collected from a donor site or the surgical site. Other options include allograft, synthetics, and growth factors such as recombinant human bone morphogenetic protein (rhBMP)-2. With the acceleration of regenerative medicine and technology, we have seen the emergence of mesenchymal stem cells (MSCs) as a possible option for increasing fusion rates, as well as decreasing complications. In this manuscript, we present current bone grafting options and then focus on the use of stem cells to augment grafting options to reduce the potential for pseudarthrosis in spinal fusion surgery.

MATERIALS AND METHODS

This manuscript is a review of the literature, performed to be up to date up until January 2021. PubMed/MEDLINE databases were searched, as well as ClinicalTrials.gov, for any literature with relevant information pertaining to stem cells and their use in spinal fusion. Keywords that were used were as follows: spine fusion, spinal fusion, stem cells, MSCs, adipose derived stem cells, autologous bone, allogeneic stem cells. Pertinent studies that were included were largely focused on preclinical and clinical trials investigating the rate of fusion with the use of stem cells in spine surgery. The heterogeneity between studies did not allow for data and statistical analysis to show whether or not fusion rates differed, but the studies are summarized and left for conclusions to be made by the readers.

Bone Graft Review

The gold standard to achieve successful fusion is currently autologous bone (autograft) from either the iliac crest or local bone graft (LBG), found in or near the surgical site. Studies have shown both grafts to be equally effective for single-level fusions, but LBG was found to be unsatisfactory for multilevel procedures.⁹ Autograft contains all 3 key elements to provide for successful spine fusion: osteoinductive factors such as cytokines and growth factors, osteoconductive materials such as collagen and minerals for a structural support scaffolding,

and osteogenic components such as osteoblastic/preosteoblastic cells and bone marrow stem cells.¹⁰

Although successful fusion rates with autograft may be as high as 95%, bone available for harvest is limited, and quality varies depending on the patient bone health, age, smoking, diabetes, and other comorbidities. Additionally, autograft harvesting may lead to infection, donor site pain, blood loss, and risk of fracture.^{11,12} These complications have been reported to be as high as 39% with iliac crest bone graft harvest.¹³

For these reasons, other modalities for grafting have been explored. Biologics and synthetics, osteogenic differentiation factors such as BMP-2, demineralized bone matrix (DBM), hydroxyapatite, provide combinations of osteoinduction and osteoconduction, but fail fundamentally as they are not osteogenic. Additionally, these products also have limitations and side effects of their own. Using these biological and synthetics as stand-alone substitutes for bone graft has not been fully explored, and would likely not provide the adequate stability and fusion ability as in conjunction with other modalities.¹⁴⁻¹⁶ The unmet need for commonly used autograft substitutes has paved the way for the investigation of using stem cells for spinal fusion.

Stem Cells

Stem cells, first described by Friedenstein¹⁷ in 1968, are defined as immature tissue precursor cells, which can differentiate into muscle, bone, tendons, fat, and other various stromal tissues.¹⁸⁻²¹ They can be categorized into embryonic stem cells, induced pluripotent cells, and adult stem cells.

MSCs, which fall under the category of adult stem cells, have benefited from advancements in the field of regenerative medicine, and are the focus for use in spinal fusion procedures. The multipotent nature of individual MSCs was first described by Pittenger et al²² in 1999. Research has shown that MSCs have osteogenic properties, can be modified to secrete osteoinductive factors, and can be implemented on an osteoconductive scaffold to successfully provide the 3 components for optimizing fusion and osteogenesis.¹⁶ MSCs are able to differentiate into osteogenic cells and also exhibit paracrine effects. Additionally, they can be easily cultured and have a high ex-vivo expansive potential.^{23,24}

MSCs can be derived from numerous adult tissues, including bone marrow, muscle, and subcu-

taneous fat,^{25,26} and have been shown to resist immunologic rejection.^{18,27} The most common source for MSCs in spinal fusion is bone marrow aspirate (BMA), followed by adipose tissue.^{28–30} Clinical studies have explored BMA harvest sites, with vertebral bodies and iliac crests having robust harvests.³¹ BMA is easily obtained in the supine position from the posterior iliac bone, or can be obtained easily through the surgical site, as well.³² Studies have looked at regenerative capacity over time with various stem cell subtypes,^{33–35} as well as regenerative capacity in older patients. Mazini et al³³ reported a maintained regenerative capacity in in vitro studies with adipose-derived stem cells, whereas bone marrow MSCs may start to lose expression of specific surface antigens in later passage.^{33,36} Although demonstrated with in vitro studies, this information is difficult to understand and demonstrate in vivo and in the clinical setting.

Bone marrow MSCs have been studied to provide new treatment methods for arthritis, periodontal disease, intrinsic muscular dystrophy, and cardiac disease because of their ability to differentiate into different cell types.^{18,22} Similarly, they have been shown to differentiate into cells of the osteogenic lineage within the appropriate conditions.^{37,38} The adipose-derived MSCs can also be extracted via liposuction, which is typically less painful than bone marrow aspiration.^{39,40} Lastly, allogeneic MSCs from matching donors have been used for patients with low bone volume, who are unable to produce enough of their own MSCs. Concerns have been expressed with allogeneic MSCs due to immune reactions in patients.⁴¹

MSCs present a lesser fraction of the total population of nucleated cells, under 0.01%^{42,43} of cells when isolated from BMA, and an in-vitro expansion phase may be necessary to obtain sufficient stem cell numbers prior to implantation.²² Several techniques for expansion exist, but problems such as sterility technique, culture time, medium used, as well as number of MSCs required are still yet to be established. Additionally, this source of cells may vary and not be as reliable in an elderly population, due to dissipation of the potency of the MSCs.⁴⁴

Preclinical Results

To date, there has been a great deal of preclinical trials investigating the efficacy of stem cells in bony fusion with various animal spinal fusion models.^{45,46}

Many of these studies have investigated variations in growth factors and scaffold options to promote optimized bony fusion, with some studies even looking at genetically modified MSCs. As common in emerging topics, the results of these studies are variable, but the majority of studies are able to replicate outcomes between autograft and MSCs with supporting scaffolds in spinal fusion.^{16,45,47–57} These studies are summarized in Table 1.

Numerous studies have shown the addition of MSCs to achieve superior rates of fusion when compared with autograft.^{45,47,51,58} Nakajima et al⁵¹ studied rabbit spines treated with MSCs cultured in osteogenic differentiation medium versus without differentiation medium and autograft, showing higher fusion rates in the first group. Minamide et al⁴⁵ demonstrated increased fusion rates in rabbits with bone marrow cells when compared with BMP and autograft. Similarly, the same group also showed higher fusion rates with bone marrow derived MSCs cultured in rhBMP-2 and fibroblast growth factor when compared with autograft. Bae et al⁵⁴ showed increased posterolateral intertransverse process fusion rates to 89% in rats treated with BMA on collagen sponges and subeffective concentrations of rhBMP-2 compared with 33%–50% with rhBMP-2 and collagen sponges alone. Additionally, Crowley et al⁵⁹ demonstrated various preclinical and clinical studies investigating MSCs implanted on biologic or synthetic scaffolds with effective results in promoting bony union.

Other studies have shown comparable results with engineered MSCs to autograft. Sheyn et al⁵⁸ and Hasharoni et al⁶⁰ demonstrated genetically modified MSCs expressing BMP-2 when placed in the paraspinal musculature induced spinal fusion in mice that were comparable with the fusion achieved with instrumentation, in regards to segment rigidity. Similar results were demonstrated when looking at MSCs seeded on an alginate scaffold with low doses of BMP-2.¹⁶

Although the majority of the preclinical models have focused on bone marrow derived MSCs, Miyazaki et al⁵⁵ compared bone marrow derived MSCs to adipose-derived MSCs, demonstrating no significant difference in fusion rates between the 2 types of MSCs in a rat model of posterolateral fusion. Similarly, Ammerman et al⁶¹ demonstrated increased fusion rates with adipose derived MSCs in a posterolateral spinal fusion rabbit model.

Table 1. Preclinical studies of spinal fusion using stem cells.

Study	Animals	Conditions	Fusion, %
Fu et al ¹⁶	24 rabbits	Autograft	92
		Alginate + MSC + BMP2	92
		Alginate + MSC	67
Bae et al ⁵⁴	53 rats	Alginate + BMP2	0
		rhBMP2/ACS + fresh syngeneic BMA transplant	89
		rhBMP2/2ACS only	50
		rhBMP2/1ACS only	33
		ACS + fresh syngeneic BMA transplant	0
		ACS only	0
Gupta et al ⁴⁷	24 sheep	TCP + bone marrow cells	33
		TCP + whole marrow	8
		TCP	0
Minamide et al ⁴⁸	30 rabbits	Autograft	25
		MSC-BMP-FGF	86
		MSC-FGF	43
		MSC-BMP	28
		MSC	0
Cinotti et al ⁴⁹	40 rabbits	Autograft	57
		Ceramic + MSC	85
		Ceramic + bone marrow	50
		Ceramic	30
Kai et al ⁵⁰	30 rabbits	Autograft	25
		Ceramic + cells	100
		Ceramic + cells + BMP	100
		Ceramic	50
Valdes et al ⁵⁶	53 rabbits	Autograft	67
		60M rhBMP6 stimulated OPC	62
		30M rhBMP6 stimulated OPC	54
		Autograft	55
Minamide et al ⁴⁵	36 rabbits	DBM	40
		Decortication alone	0
		BMP-HA	100
		High marrow cells	71
		Low marrow cells	0
Nakajima et al ⁵¹	24 rabbits	Autograft	57
		Osteogenic MSC	80
		MSC	33
		Hydroxyapatite	0
Wang et al ⁵²	9 monkeys	Autograft	67
		MSC+ ceramic	67
		Ceramic	17
		Autograft	83
Cui et al ⁵⁷	52 rats	Bone marrow	50
		DI-BAG cells	100
		Matrix only	0
Cuenca-López et al ⁵³	34 sheep	HA + MSC	35
		HA	22
		Allograft	70
		Autograft	70
Miyazaki et al ⁵⁵	48 rats	Collagen + adipose-derived MSC + adeno-BMP2	100
		Collagen + marrow-derived MSC + adeno-BMP2	100
		Collagen + BMP2	100
		Collagen + adipose-derived MSC + adeno-LacZ	0
		Collagen + marrow-derived MSC + adeno-LacZ	0
		Collagen alone	0

Abbreviations: ACS, absorbable collagen sponges; Adeno-LacZ, study specific; BMA, bone marrow aspirate; BMP2, bone morphogenic protein; DBM, demineralized bone matrix; DI-BAG, study specific; FGF, fibroblast growth factor; HA, hydroxyapatite; MSC, mesenchymal stem cell; OPC, osteoprogenitor cells; TCP, tricalcium phosphate.

To summarize, preclinical trials demonstrate promising outcomes for MSCs in spinal fusion. It is difficult to compare different trials and perform statistical analysis, as these results are highly dependent on the use of specific growth factors and differentiation mediums to aid in bone formation, as well as the use of appropriate scaffoldings

and animals. The efficacy of MSCs to promote spinal fusion without the addition of genetic engineering or additional growth factors has been less than ideal.^{54,60,62} With the appropriate techniques, MSCs either approach or match the fusion rates achieved with autograft in preclinical models. As with any type of treatment, the question of how

Table 2. Clinical studies using stem cells for spinal fusion.

Study	Patients	Approach	Type	Conditions	Fusion, %
Gan et al ⁴²	41	PLF/TLF	Autologous	Enriched BMA + β -TCP	95.1
Hostin et al ⁶³	22	AIF	Autologous	Collagen + BMA w/cage	87
Kitchel ⁶⁴	25	PLF, IF	Autologous	Collagen + BMA	80
				Iliac crest bone graft	84
Neen et al ⁶⁵	50	PLF, TLF, 360	Autologous	Collagen/hydroxyapatite + BMA	PLF 93, IF 85
				Iliac crest bone graft	PLF 93, IF 92
Niu et al ⁶⁶	21	PLF	Autologous	LBG + BMA	85.7
				Iliac crest bone graft	90.5
Vaccaro et al ⁶⁷	73	PLF	Autologous	BMA + DBM	63
				Iliac crest bone graft	67
Bansal et al ⁶⁸	30	PLF	Autologous	Hydroxyapatite + TCP + BMA	100
				Iliac crest bone graft	96
Taghavi et al ⁶⁹	62	PLF	Autologous	Collagen + BMA	100
				LBG	100
Odri et al ⁷⁰	15	PLF	Autologous	BMC + BPCG + autologous bone	100
Hart et al ⁷¹	40	PLF	Autologous	BMC + allograft	80
Ammerman et al ⁶¹	23	TLIF	Allogeneic	Osteocel + DBM	91.3
McAfee et al ⁷²	25	XLIF	Allogeneic	Autograft/Osteocel	85
Caputo et al ⁷³	30	XLIF	Allogeneic	Osteocel + DBM	89.6
Tohmeh et al ⁷⁴	40	XLIF	Allogeneic	Osteocel + DBM	90.2
Kerr et al ⁶²	52	ALIF, TLIF, 360	Allogeneic	Osteocel	92.3
Peppers et al ⁷⁵	40	ACDF	Allogeneic	Trinity	91.4
Eastlack et al ⁷⁶	182	ACDF + plating	Allogeneic	Osteocel and PEEK interbody	87

Abbreviations: ACDF, anterior cervical discectomy and fusion; AIF, anterior interbody fusion; ALIF, anterior lumbar interbody fusion; BMA, bone marrow aspirate; BMC, bone marrow concentrate; BPCG, bisphasic phosphate ceramics graft; β -TCP, tricalcium phosphate scaffolding; DBM, demineralized bone matrix; IF, interbody fusion; LBG, local bone graft; PEEK, polyetheretherketone; PLF, posterolateral fusion; TCP, tricalcium phosphate; TLF, transforaminal lumbar fusion; TLIF, transforaminal lumbar interbody fusion; XLIF, extreme lateral interbody fusion.

these results will translate to clinical results is what is most important.

Clinical Results

Clinical studies examining the efficacy of MSCs on spinal fusion are more limited than preclinical trials. Most studies look at MSCs isolated from BMA, as it can be harvested from the iliac crest or vertebral body intraoperatively and then transplanted to the fusion site. Additionally, many of the clinical studies use different carrier scaffolds, making it difficult to compare between trials, but in general, utilizing MSCs for spinal fusion show fusion rates from 63%–100%.^{42,61,61–76} Studies are shown in Table 2.

There are multiple prospective trials and also systemic reviews investigating outcomes of stem cells and spinal fusion, mostly with the use of computed tomography or plain radiographs. In a prospective study, Gan et al⁴² reported on 41 patients with enriched BMA on a β -tricalcium phosphate scaffold achieving 95.1% fusion at 24-month follow-up. Another study reported an 87% successful fusion rate with 182 patients in a multicenter prospective trial.⁵¹ Odri et al⁷⁰ achieved 100% fusion on 15 patients receiving MSCs with macroporous biphasic phosphate ceramic scaffolds and autologous bone.

A systematic review by Khashan et al³¹ compiled results from 7 different clinical studies, 6 prospective and 1 randomized control trial. Studies contained at least 20 patients each and compared BMA on a scaffold to iliac crest and/or LBG.³¹ Fusion rates for BMA with scaffolds ranged from 63%–100%, whereas LBG or iliac crest bone graft ranged from 67%–100%. The majority of these studies examined posterolateral fusion. The review concluded that there is still insufficient evidence to support the use of MSCs or BMA over autologous bone graft.

There are also studies involving allogeneic MSCs (Table 2). Peppers et al⁷⁵ reported on 40 patients undergoing anterior discectomy and fusion with Trinity Evolution Viable Cellular Bone Matrix, an allogeneic stem cell source, with 91.4% fusion rates. Another study reported 92.3% fusion rates for 52 patients with Osteocel, an allograft-based tissue containing live stem cells.⁶² This fusion rate is higher in comparison with Osteocel when being used with DBM (89.6%) and autograft (85%). Osteocel has also shown promising outcomes in other studies, with fusion rates ranging between 87% and 92%.^{61,62,73,76} These studies and products demonstrate high rates of fusion and may be options for patients who are unable to use their own MSCs for fusion.

Currently, as of January 2021, there are 11 active studies listed on ClinicalTrials.gov investigating the

Table 3. Current clinical trials.

ClinicalTrials.gov Identifier	Description	Design	Outcomes
NCT01552707 ⁷⁷	Isolation and ex-vivo expansion of MSCs with Xcelia, then fixed to allogenic bone compared with bone iliac crest alone.	Prospective, RCT	Safety of Xcelia, feasibility of Xcelia, efficacy of spinal fusion
NCT02297256 ⁷⁸	Bone marrow aspirate concentrate (BMAC) + allograft compared with iliac crest bone graft during posterior lumbar/lumbosacral spine fusion.	Prospective, RCT	Fusion status, Oswestry Disability Index (ODI), Short Form Health Survey (SF-12), numeric pain rating scale, length of stay
NCT02924571 ⁷⁹	BMAC and allograft compared with rh-BMP2 for thoracolumbar spine fusion with interbody support.	Prospective, blinded, nonrandomized	ODI, SF-12, numeric pain rating scale
NCT03827096 ⁸⁰	Bone marrow cell aspiration from iliac crest cultivated for 3 passages to expand and suspended on β -tricalcium phosphate foam to lumbar spine.	Single group assignment	Demonstrate absence of complications at the site of spinal fusion, ODI, efficacy of spinal fusion on x-ray and CT
NCT00996073 ⁸¹	Allogeneic mesenchymal precursor cells (NeoFuse) combined with MasterGraft Matrix compared with use of autologous iliac crest bone graft in lumbar interbody fusion site.	Prospective, multicenter, randomized	Determine safety, evaluate overall fusion success
NCT01097486 ⁸²	Allogeneic mesenchymal precursor cells (NeoFuse) combined with MasterGraft Matrix compared with use of autologous iliac crest bone graft in multilevel anterior cervical discectomy and fusion.	Prospective, multicenter, randomized, single-blinded	Determine safety, evaluate overall fusion success
NCT02070484 ⁸³	Stem cell allograft (NuCel) compared with demineralized bone matrix.	Randomized, parallel assignment	ODI, evaluation of fusion via CT
NCT00941980 ⁸⁴	Stem cells attached to allograft bone matrix (Osteoecel Plus) in subjects undergoing posterior lumbar interbody fusion surgery compared with historic autograft control.	Prospective, nonrandomized multicenter	Evaluate fusion rates, complications, radiographic outcome, surgical time, blood loss
NCT00948831 ⁸⁵	Osteoecel Plus in subjects undergoing anterior lumbar interbody fusion.	Prospective, nonrandomized multicenter	Evaluate fusion rates, complications, radiographic outcome, surgical time, blood loss
NCT00947583 ⁸⁶	Osteoecel Plus in subjects undergoing transforaminal lumbar interbody fusion.	Prospective, nonrandomized multicenter	Evaluate fusion rates, complications, radiographic outcome, surgical time, blood loss
NCT00951938 ⁸⁷	Allogeneic cancellous bone matrix with viable osteoprogenitor cells, MSCs, and demineralized cortical bone (Trinity Evolution) in patients undergoing ACDF.	Case-only, prospective	Fusion rates, pain, complications

Abbreviations: BMP2, bone morphogenic protein; CT, computed tomography; MSC, mesenchymal stem cell; RCT, randomized controlled trial.

effects of stem cells on spinal fusion (Table 3). As time goes on, with expanded popularity of stem cells increasing fusion rates, we will likely see more studies investigating clinical outcomes associated with MSCs compared with autologous bone graft. Ultimately, with the heterogeneity of clinical studies at this time, it is difficult to directly compare fusion rates with autologous bone graft, but the studies are overall promising.

DISCUSSION

Successfully achieving spinal fusion is highly dependent on having osteogenic, osteoconductive, and osteoinductive factors available. Traditionally, autologous bone harvesting from the surgical site or iliac crest have been the standard of care, but complications, donor site morbidity, and limited

quantities of bone have brought other options into the spotlight.

MSCs have the potential to become widely used as bone graft augmentation and for achieving successful spinal fusion. It is believed that stem cells contribute to the fusion process and improve union through their osteogenic and osteoinductive properties within the fusion site, although the primary contribution is still unknown. Although they are not yet on the path of becoming the gold standard for achieving successful fusion, MSCs have gained interest due to ease of use, ability to harvest intraoperatively, and the regenerative capabilities. At this time, multiple factors need to be optimized, such as: intrinsic and extrinsic expression of growth factors and cytokines, the optimization of choosing the material and construction of scaffoldings for the cells to be supported, and finding the most beneficial

area of stromal cell harvest, while also minimizing the morbidity of the procedure. MSCs cannot be used alone to promote spinal fusion; they must be used in conjunction with, at minimum, a scaffolding to hold them in place.

There are a significant amount of preclinical studies, which show comparable outcomes when using MSCs to achieve successful spinal fusion. These studies are heterogeneous in nature, using different animal models, various scaffolds, and also various growth factors and harvesting mediums for the cells. Although many types of combinations have been examined, we do not know the optimal combination that will translate to success in the clinical realm. Even if this perfect combination of factors, cells, and scaffold is discovered, we also may not be sure if it is a one-size-fits-all for every patient.

In clinical models, we have seen studies with fusion rates of 63%–100%. Most studies demonstrated at least approximately equal fusion rates, but again, just as with preclinical models, different studies used different combinations of growth factors and scaffolds. Additionally, numerous studies look at the use of BMA, rather than just the MSCs isolated from BMA. Clinical trials involving allogeneic MSCs have shown promise in patients, such as elderly, who may be limited in the number of viable cells available from liposuction or BMA. Lastly, we also are not sure how many of these cells continue to be viable after placement. Newer technologies and techniques are required to quantify viable cells and ensure their survival after implantation.

As future studies are started and planned, we should look to standardize certain aspects, that way we can compare between different studies. Similarly, we can start looking at patient demographics, approaches, and the pathology behind the reason for spinal fusion, to further understand what combination of scaffold and MSCs will work in each setting. Additionally, as we start to understand these factors, cost analyses and other outcome studies will become just as important as fusion rates. Lastly, future studies need to address the regenerative capacity of stem cells over time. Although in vitro studies exist, it would be important to the field to understand the temporal benefit of in vivo use of MSCs in spinal fusion.

In summary, the future of spinal fusion may be heading in the direction of using MSCs, synthetics,

and proteins such as BMP-2 to improve outcomes. Preclinical results with various proteins, growth factors, and scaffolds have shown promising results. Clinical trials, including prospective studies, have shown that MSCs with the appropriate harvesting, growth factors, and scaffoldings can provide comparable fusion rates to autograft. As new studies begin to emerge, and as regenerative medicine and technology advance, we may see MSCs becoming a staple of spinal fusion surgeries.

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