

## Stem Cells Therapy as a Treatment for Discogenic Low Back Pain: A Systematic Review

Randy Randy, Khandar Yosua, Aswin Guntara and Nicko P. Hardiansyah

*Int J Spine Surg* 2025, 19 (2) 156-167

doi: <https://doi.org/10.14444/86717>

<https://www.ijssurgery.com/content/19/2/156>

This information is current as of May 13, 2025.

---

**Email Alerts** Receive free email-alerts when new articles cite this article. Sign up at:  
<http://ijssurgery.com/alerts>

# Stem Cells Therapy as a Treatment for Discogenic Low Back Pain: A Systematic Review

RANDY RANDY, MD<sup>1</sup>; KHANDAR YOSUA, MD<sup>2</sup>; ASWIN GUNTARA, MD<sup>3</sup>; AND NICKO P. HARDIANSYAH, MD<sup>4</sup>

<sup>1</sup>Emergency Department, Mayapada Hospital Kuningan, Jakarta, Indonesia; <sup>2</sup>Emergency Department, Betang Pabelum Hospital, Palangka Raya, Indonesia;

<sup>3</sup>Emergency Department, Esnawan Antariksa Air Force Hospital, Jakarta, Indonesia; <sup>4</sup>Spine Consultant, Department of Orthopedic and Traumatology, Mayapada Hospital Kuningan, Jakarta, Indonesia

 RR, 0009-0003-6554-9950

## ABSTRACT

**Background:** Low back pain (LBP) is 1 of the most common problems that present in 80% of people. LBP can be caused by some pathologies, with discogenic pain being 1 source. Pain from LBP can become chronic and also cause disability. Treatment options for LBP varied from conservative to operative, and a novel treatment nowadays is using stem cells therapy to treat with pain from LBP.

**Methods:** Database searches from Pubmed and ScienceDirect from inception to 13 September 2023. A total of 283 discogenic LBP cases from 8 articles. This study measured clinical outcomes using a visual analog scale (VAS) and Oswestry Disability Index (ODI) obtained from each study.

**Results:** Functional outcomes in patients treated with stem cell therapy showed significant improvement ODI and VAS ( $P < 0.00001$ ). Improvement also showed in Pfirrmann grade before and after treatment with stem cells ( $P = 0.005$ ). Subgroup analyses using bone marrow aspirate concentrate also showed significant differences in both ODI and VAS ( $P < 0.00001$ ).

**Conclusion:** Stem cells therapy could be beneficial as an option of treatment for discogenic LBP in improving pain and activity of daily living.

**Clinical Relevance:** Intradiscal stem cell therapy is a promising alternative for managing discogenic low back pain, offering improvements in pain and function.

**Level of Evidence:** 4.

Biologics

Keywords: discogenic low back pain, stem cells therapy, oswestry disability index, visual analog scale

## INTRODUCTION

Low back pain (LBP) is the most common musculoskeletal symptom caused by degenerative joint disease in the lumbar spine.<sup>1</sup> LBP is defined as pain and discomfort between the costal margin and inferior gluteal fold with or without leg pain. An estimated 80% of the normal population will experience LBP in their lifetime, and this has enormous socioeconomic consequences.<sup>2</sup> One of the conditions leading to degenerative joint disease in the lumbar spine is degenerative disc disease, in which this condition will lead to herniation of the intervertebral disc (IVD).<sup>1</sup>

Discogenic pain is a common source of LBP, with an overall prevalence of 26% to 42%. In younger populations, this increases to over 80%. Discogenic LBP is typically persistent with chronic pain and disability, and most treatment resources are directed toward refractory pain symptoms.<sup>3</sup> The pathophysiology of discogenic back pain involves an imbalance in the anabolic and

catabolic environments of the extracellular matrix in favor of catabolism. The resultant alteration in disc height affects the biomechanics of the involved spinal segment, which often results in segmental instability.<sup>4</sup>

It is estimated that the annual worldwide LBP incidence in adults is 15%, and the point prevalence is 30%.<sup>5</sup> In 2020, LBP affected 619 million (95% uncertainty interval 554–694) people globally, with a projection of 843 million (759–933) prevalent cases by 2050.<sup>6</sup> The high rate of LBP prevalence observed in all regions globally could have some important social and economic consequences, especially considering the substantial cost of care for this condition. For instance, from 2012 to 2014, the direct aggregate costs for all individuals with a spine condition in the USA were US\$315 billion, with a substantial proportion of costs attributed to surgical procedures.<sup>6</sup>

Makkiyah et al said that the prevalence of LBP in middle-aged adults in Indonesia was 44.29% at 12

months.<sup>7</sup> The Ministry of Health of the Republic of Indonesia stated that the number of LBP incidents in Indonesia in 2018 was 18%. Based on Indonesia's 2017 national and subnational disease burden analysis data, LBP complaints were the main cause of loss of productive years due to disability due to illness and injury in Jakarta in 2017.<sup>8</sup>

Treatment modalities in the management of chronic lumbosacral pain include conservative management with physical therapy, pharmacological therapy, interventional and intradiscal, as well as surgical intervention through fusion or disc replacement.<sup>9</sup> At present, nonsurgical treatment based on physiotherapy and pharmacological interventions remains the first-line treatment option for lumbar discogenic pain.<sup>10</sup>

In recent years, there has been an increasing shift toward regenerative therapies for several pathologies across the entire spectrum of medicine.<sup>11</sup> Among the biological disc repair therapies, cell therapy has gained interest as it offers a disc regenerative potential while being minimally invasive.<sup>12</sup> A cell therapy approach aims to address disc inflammation by inhibiting aberrant cytokine production and to promote disc rehydration and height restoration by initiating matrix anabolism, as well as repopulating and stimulating the native cells.<sup>12</sup> As a result of these efforts, number of different regenerative modalities are being considered as treatment options for LBP due to DDD; these include platelet-rich plasma, stem cells, and bone marrow concentrate (BMC). Among these options, stem cell and BMC have shown promising results in the treatment of discogenic LBP.<sup>11</sup>

Based on all this, we conduct a systematic review of current literature on the role of stem cell therapy treatment for patients with discogenic LBP aiming to evaluate clinical improvement on LBP treated with stem cell therapy.

## METHODS

### Search Strategy and Study Selection

This study was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-Analysis guidelines.<sup>13</sup> A comprehensive search across 2 databases (PubMed and ScienceDirect) from inception to 13 September 2023. The search terms used were “stem cell therapy” AND “discogenic LBP” (Figure 1). The inclusion criteria were (1) patients with discogenic LBP treated with stem cells therapy, (2) a study presents the effectiveness of intradiscal stem cell therapy, and (3) in English. Exclusion criteria were (1) patients had

history of spinal fusion, (2) case report and review, and (3) incomplete data.

### Eligibility Criteria and Quality Appraisal

The included studies were of any design reporting on the usage of stem cells for the treatment of discogenic LBP. The inclusion criteria used in this study were (1) published in English, (2) concerned discogenic LBP, and (3) patients had no history of prior surgery at the affected lower back. The exclusion criteria used in this study were (1) review articles, (2) animal studies, and (3) articles unavailable in English. Study quality was assessed with the Cochrane risk of bias tool for randomized controlled trials (RCTs) and the methodological index for non-randomized studies score for non-RCTs (Figure 2, Table 1).<sup>14</sup> Each included study stated that there was no source of funding for their research.

### Data Extraction

Three independent reviewers (R., K.Y., and A.G.) selected the articles included in the study through title and abstract screening and conducting full-text reviews of the selected articles. Any disagreements were resolved by a fourth, independent reviewer (N.P.H.). Initially, in the literature searching and cross-referencing, 86 articles were found, 19 articles were excluded before screening because of duplication. After titles and abstracts were screened, 11 articles met the eligibility criteria. After a full-text review, 3 articles were excluded (prestudy protocol and nonusable data), and 8 articles were included in the systematic review.

### Statistical Analyses

Mean difference and 95% confidence intervals were calculated using the inverse variance method, and study heterogeneity was assessed using  $I^2$  with a value of >50% marked as significant heterogeneity. If the  $I^2 > 50\%$ , subgroup analysis was carried out. Statistical significance was defined with a  $P$  value of <0.05. Statistical analyses were performed by Review Manager 5.4 analysis software.

## RESULTS

### Patient Characteristics and Demographics

A total of 86 articles were identified, with 8 studies meeting the inclusion criteria after screening. Details of the studied, such as the number of patients, gender, study design, outcomes, and follow-up period, are presented in Table 2. Seven studies including a total of 283

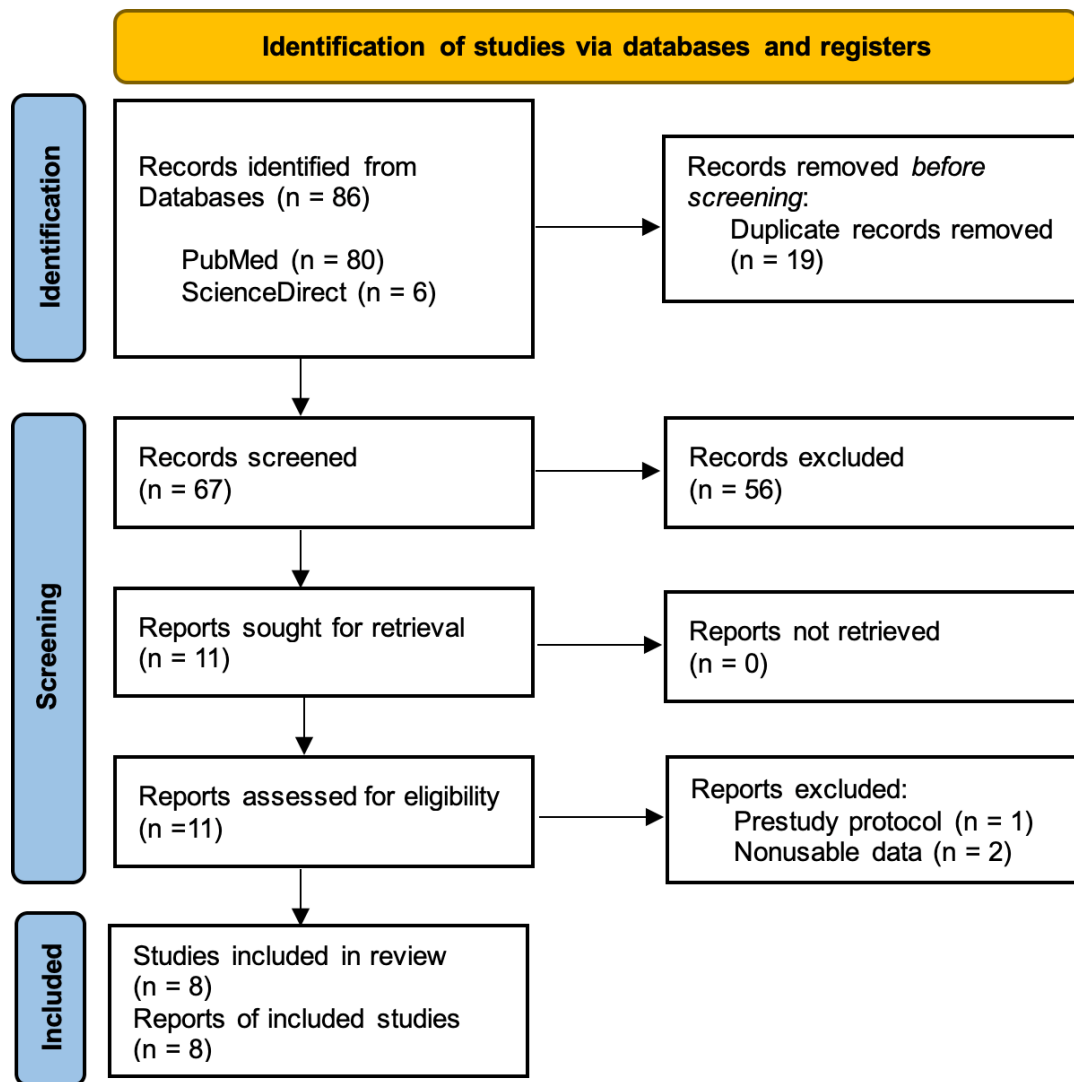


Figure 1. Preferred Reporting Items for Systematic Reviews and Meta-Analysis flowchart.

patients were used in statistical analysis. One study was not included in statistical analysis because the results in the study were grouped. Patients' ages ranged from 35 to 60 years. The follow-up period after stem cell therapy ranged from 12 to 36 months.

### Outcome Measurement and Results

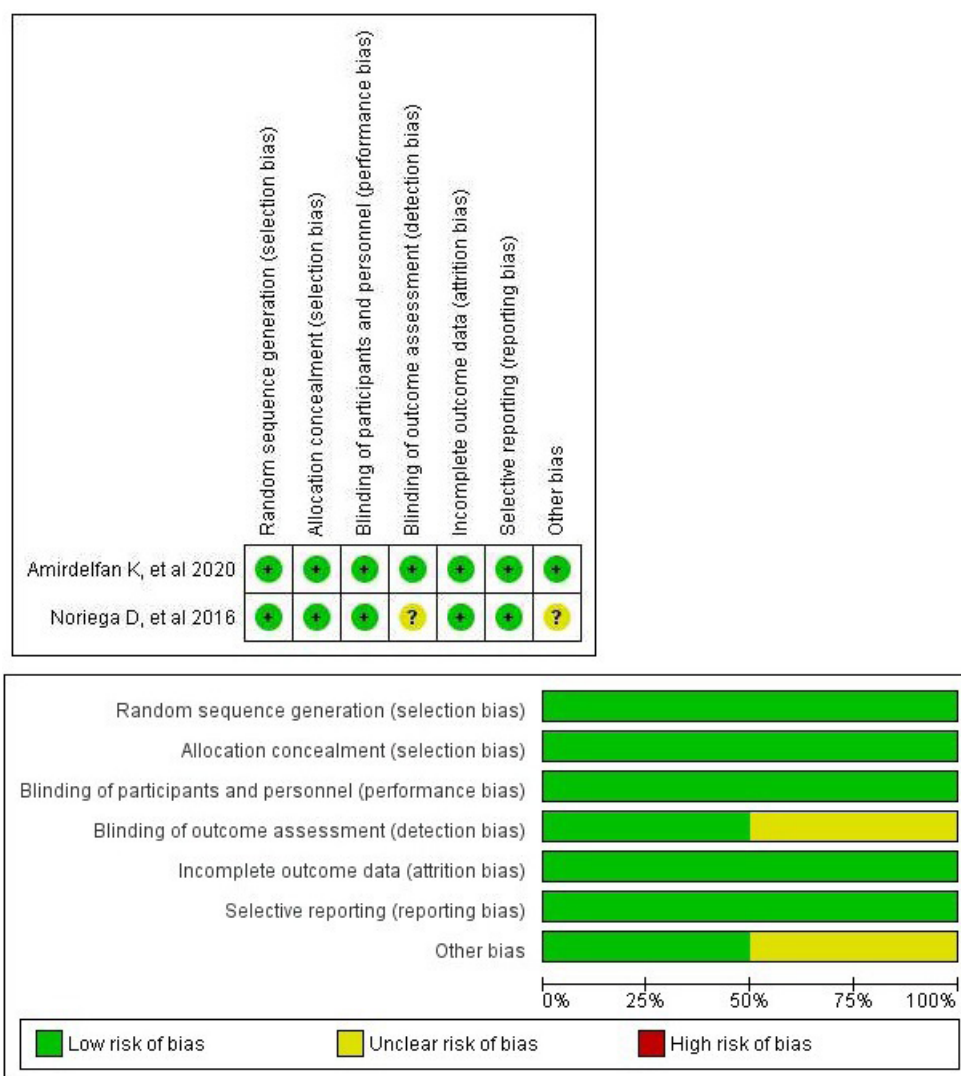
The improvement in the Oswestry Disability Index (ODI) in 7 studies resulted in a significant difference ( $P < 0.00001$ ) in patients treated with stem cell therapy. Subgroup analysis based on follow-up periods revealed significant reductions in ODI observed across all time points; however, heterogeneity was high ( $I^2 = 94\%$ ; Table 3; Figure 3).

Pain scale using a visual analog scale (VAS) or numeric rating scale in 7 studies before and after treatment using stem cells therapy resulted in significant

difference ( $P < 0.00001$ ). Subgroup analysis by follow-up period similarly indicated consistent improvement across time points, though heterogeneity remained high ( $I^2 = 94\%$ ; Table 3; Figure 4).

A subgroup analysis of ODI and VAS outcomes in studies using bone marrow aspirate concentrate was conducted across 4 studies. ODI improvement was significant across follow-up periods of 1, 3, 6, and 12 months, with moderate heterogeneity ( $I^2 = 58\%$ ; Table 3; Figure 5). For VAS, it was significantly different in follow-up periods 3, 6, and 12 months. In 1-month follow-up, it included 2 studies but was insignificant ( $P = 0.17$ ), and the pooled heterogeneity was moderate ( $I^2 = 76\%$ ; Table 3; Figure 6).

Magnetic resonance imaging assessments using Pfirrmann grading scale were available in 5 studies. Two studies included in the meta-analysis showed



**Figure 2.** Risk of bias of included randomized controlled trials.

**Table 1.** Methodological index for non-randomized studies score.

| Criteria                                             | Lewandrowski et al, 2023 <sup>15</sup> | Wolff et al, 2020 <sup>16</sup> | Pettine et al, 2017 <sup>17</sup> | Atluri et al, 2021 <sup>18</sup> | Orozco et al, 2011 <sup>19</sup> | Kumar et al, 2017 <sup>20</sup> |
|------------------------------------------------------|----------------------------------------|---------------------------------|-----------------------------------|----------------------------------|----------------------------------|---------------------------------|
| A clearly stated aim                                 | 2                                      | 1                               | 2                                 | 2                                | 2                                | 2                               |
| Inclusion of consecutive patients                    | 2                                      | 2                               | 2                                 | 2                                | 2                                | 2                               |
| Prospective collection of data                       | 0                                      | 0                               | 2                                 | 2                                | 2                                | 2                               |
| Endpoint appropriate to the aim of the study         | 2                                      | 2                               | 2                                 | 2                                | 2                                | 2                               |
| Unbiased assessment of the study endpoint            | 1                                      | 1                               | 0                                 | 0                                | 1                                | 1                               |
| Follow-up period appropriate to the aim of the study | 2                                      | 2                               | 2                                 | 2                                | 2                                | 2                               |
| Loss of follow-up < 5%                               | 0                                      | 1                               | 2                                 | 1                                | 2                                | 2                               |
| Prospective calculation of the study size            | 2                                      | 2                               | 2                                 | 2                                | 1                                | 2                               |
| Additional criteria for comparative studies          |                                        |                                 |                                   |                                  |                                  |                                 |
| An adequate control group                            | NA                                     | NA                              | NA                                | 2                                | NA                               | NA                              |
| Contemporary group                                   | NA                                     | NA                              | NA                                | 2                                | NA                               | NA                              |
| Baseline equivalent of groups                        | NA                                     | NA                              | NA                                | 2                                | NA                               | NA                              |
| Adequate statistical analysis                        | NA                                     | NA                              | NA                                | 2                                | NA                               | NA                              |
| Total                                                | 11                                     | 11                              | 14                                | 21                               | 14                               | 15                              |

Abbreviation: NA, not applicable.

Note: Low risk of bias : 13–16 (noncomparative studies), 20–24 (comparative studies); moderate risk of bias: 9–12 (noncomparative studies), 15–19 (comparative studies); high risk of bias: 0–8 (noncomparative studies), 0–14 (comparative studies).

Table 2. Characteristics of included studies.

| Study                                 | Study Design (Level of Evidence)              | Sample Size and Demographic                                     | Follow-Up, mo | Type and Site of Insertion                                                              | Dose, Cells (mean CFU-F)                                         | Main Results                                                                                                                                                                                                                                                                                                                   | Radiological Results                                                                                                                                          |
|---------------------------------------|-----------------------------------------------|-----------------------------------------------------------------|---------------|-----------------------------------------------------------------------------------------|------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Lewandowski et al, 2023 <sup>15</sup> | Retrospective observational cohort (II)       | N = 33, mean age = 47.6 y, and M:F = 18:15                      | 24            | Allogeneic polyclonal MSC, intradiscal                                                  | 5 × 10 <sup>6</sup> allogeneic polyclonal MSC + 1% HA            | ODI score<br>● Baseline 44.81; 1 mo 19.7; 3 mo 15.38; 6 mo 13.48; 12 mo 11.8; 24 mo 6.07 (P < 0.001)<br>VAS score<br>● Baseline 82.2; 1 mo 33.1; 3 mo 25.7; 6 mo 20.3; 12 mo 18.1; 24 mo 17.4 (P < 0.001)<br>1 patient experienced severe LBP for 1 d, but otherwise, no adverse treatment effects were mentioned in the study | Baseline: Pfirrmann grade (mean ± SD): 4.06 ± 0.72<br>Result: 3.65 ± 0.81                                                                                     |
| Wolff et al, 2020 <sup>16</sup>       | Retrospective pilot study (II)                | N = 33, mean age = 45 y, and M:F = 19:14                        | 12            | BMC, Intradiscal                                                                        | 3 mL, cells NA                                                   | ODI improvement from baseline<br>● 2 wk 4.2%, 6–8 wk 26.7%; 3 mo 36.4%; 6 mo 55.6%; 12 mo 30.8%<br>NRS improvement from baseline<br>● 2 wk 13.8%, 6–8 wk 45.8%; 3 mo 41.1%; 6 mo 23.5%; 12 mo 38.9%<br>No remarkable adverse event reported                                                                                    | NA                                                                                                                                                            |
| Pettine et al, 2017 <sup>17</sup>     | Prospective, nonrandomized (II)               | N = 26, mean age = 40 y, and M:F = 11:15                        | 36            | Autologous BMC, Intradiscal                                                             | 2–3 mL, 121 × 10 <sup>6</sup> per mL (2,713 per mL)              | ODI score<br>● Baseline 56.7; 36 mo 17.5 (P < 0.001)<br>VAS score<br>● Baseline 82.1; 36 mo 21.9 (P < 0.001)<br>No adverse events related to the study                                                                                                                                                                         | Baseline:<br>Pfirrmann Grade IV = 4,<br>Grade V = 11,<br>Grade VI = 15, and<br>Grade VII = 10.<br>Result: 8 of 20 patients had improved Pfirrmann grade<br>NA |
| Atluri et al, 2021 <sup>18</sup>      | Open label, prospective controlled study (II) | N = (40 intervention/40 control), mean age = 60 y, and M:F = NA | 12            | Autologous bone marrow MSC; intradiscal, facet intraepidural, facet joint, and SI joint | 2 mL (each level), 239.3 × 10 <sup>6</sup> per mL (4,987 per mL) | ODI score, intervention vs control<br>● Baseline 46.1 vs 44.3; 1 mo 33.4 vs 45.9; 3 mo 28.8 vs 47.3; 6 mo 29.9 vs 48.9; 12 mo 31.1 vs 49.5 (P < 0.001)<br>NRS score, intervention vs control<br>● At baseline 71 vs 66; 1 mo 38 vs 68; 3 mo 31 vs 69; 6 mo 37 vs 66; 12 mo 42 vs 71 (P < 0.001)                                | NA                                                                                                                                                            |
| Orozco et al, 2011 <sup>19</sup>      | Pilot study (II)                              | N = 10, mean age = 35 y, and M:F = 4:6                          | 12            | Autologous bone marrow cells, intradiscal                                               | 10 <sup>7</sup> cells per disc                                   | ODI score<br>● Baseline 25; 3 mo 13; 6 mo 9.4; 12 mo 7.4 (P < 0.0001)<br>VAS score<br>● Baseline 68.9; 3 mo 26.5; 6 mo 21.6; 12 mo 20 (P < 0.0001)                                                                                                                                                                             | NA                                                                                                                                                            |
| Noriega et al, 2016 <sup>21</sup>     | RCT (I)                                       | N = 24, mean age = 38 y, and M:F = 17:7                         | 12            | Allogeneic mesenchymal bone marrow cells, Intradiscal                                   | 25 × 10 <sup>6</sup> in 2 mL saline per disc                     | ODI score, intervention vs control<br>● Baseline 34 vs 24; 1 wk 27 vs 20; 3 mo 16 vs 25; 6 mo 20 vs 30; 12 mo 22 vs 34<br>VAS score, intervention vs control<br>● Baseline 67 vs 62; 1 wk 63 vs 45; 3 mo 43 vs 46; 6 mo 40 vs 51; 12 mo 47 vs 47                                                                               | Baseline (control vs intervention): Pfirrmann 3.15 ± 0.76 vs 3.68 ± 0.67<br>Results:<br>3.78 ± 0.82 vs 3.18 ± 0.87                                            |

Table 2. Continued.

| Study                                | Study Design (Level of Evidence)                    | Sample Size and Demographic                                | Follow-Up, mo | Type and Site of Insertion                                                                 | Dose, Cells (mean CFU-F)                                                                   | Main Results                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | Radiological Results                                                                                                                                                                                                                                               |
|--------------------------------------|-----------------------------------------------------|------------------------------------------------------------|---------------|--------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Kumar et al, 2017 <sup>20</sup>      | Phase 1 clinical trial, open label, single arm (II) | N = 10, mean age = 43.5 y, and M:F = 6:4                   | 12            | Adipose-derived MSC + HA, Intradiscal                                                      | Low dose $2 \times 10^7$ cells/disc (n = 5); high dose $4 \times 10^7$ cells/disc (n = 5)  | ODI score<br>● Baseline 42.8; 1 mo 31.2; 3 mo 31.7; 6 mo 21.3; 12 mo 16.8 (P = 0.002)<br>VAS score<br>● Baseline 65; 1 mo 46; 3 mo 43; 6 mo 32; 12 mo 29 (P = 0.002)                                                                                                                                                                                                                                                                                                                                                         | Baseline: Pfirrmann Grade IV = 10.<br>Result: No increase in Pfirrmann grade                                                                                                                                                                                       |
| Amirdelfan et al, 2020 <sup>22</sup> | RCT (I)                                             | N = 100 (6M 30/18M 30); age = 37.9–44.5 y, and M:F = 53:47 | 36            | Allogeneic MPC + 1% HA 1 mL (6 million and 18 million) control: saline and HA; Intradiscal | Low dose: $6 \times 10^6$ MPC + 1 mL 1% HA<br>High dose: $18 \times 10^6$ MPC + 1 mL 1% HA | No adverse event during 1 y follow-up<br>ODI score<br>● Baseline 52.07; 1 mo 36.83; 3 mo 32.59; 6 mo 28.25; 12 mo 31.85; 24 mo 29.69; 36 mo 30.69 (P < 0.05)<br>VAS score<br>● Baseline 69.67; 1 mo 40.93; 3 mo 30.28; 6 mo 25.93; 12 mo 31.48; 24 mo 33; 36 mo 35.69 (P < 0.05)<br>ODI score<br>● Baseline 50.67; 1 mo 37.33; 3 mo 32.89; 6 mo 31.7; 12 mo 29.59; 24 mo 28.75; 36 mo 24.95 (P < 0.05)<br>VAS score<br>● Baseline 71.47; 1 mo 40.97; 3 mo 35.26; 6 mo 34.6; 12 mo 32.37; 24 mo 37.63; 36 mo 28.24 (P < 0.05) | Baseline (control vs intervention):<br>Grade II (0 vs 4),<br>Grade III (12 vs 13),<br>Grade IV (19 vs 32),<br>Grade V (3 vs 4),<br>Grade VI (2 vs 3),<br>Grade VII (3 vs 3), and<br>Grade VIII (1 vs 1).<br>Results: No significant improvement in Pfirrmann score |

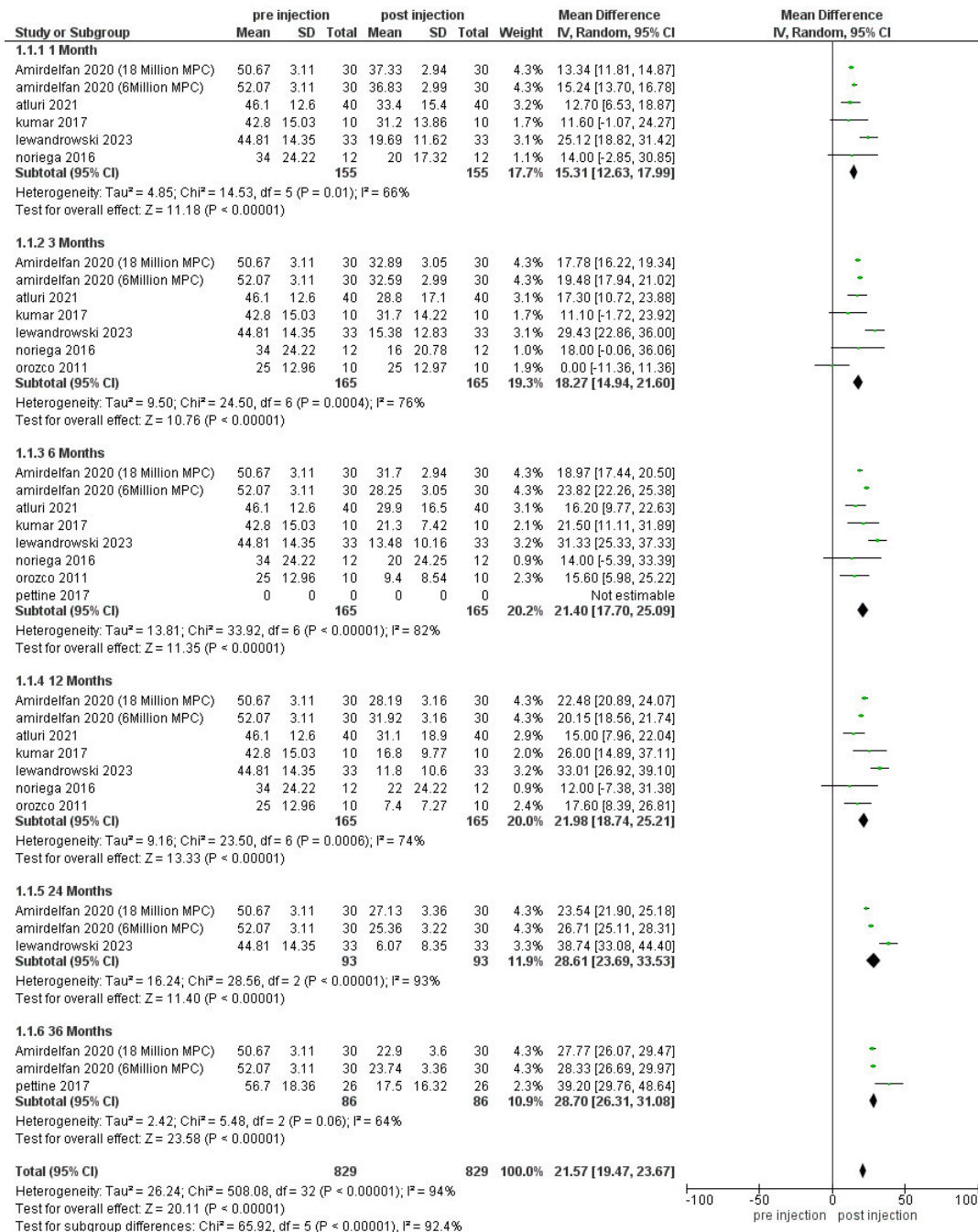
Abbreviations: BMC, bone marrow concentrate; CFU-F, colony forming unit-fibroblast; HA, hyaluronic acid; LBP, low back pain; M:F, male:female; MPC, mesenchymal precursor cell; MSC, mesenchymal stem cell; NA, not applicable or not available; NRS, numeric rating scale; RCT, randomized controlled trial; SI, sacroiliac; VAS, visual analog scale.

**Table 3.** Statistical analysis results.

| Outcome Measure          | No of studies | Mean Difference (95% CI) | P                  | I <sup>2</sup> (Heterogeneity) |
|--------------------------|---------------|--------------------------|--------------------|--------------------------------|
| ODI                      | 7             | 21.57 (19.47, 31.08)     | <b>&lt;0.00001</b> | 94% (high)                     |
| VAS                      | 7             | 38.97 (36.01, 41.93)     | <b>&lt;0.00001</b> | 94% (high)                     |
| ODI after BMAC treatment | 4             | 16.99 (12.65, 21.33)     | <b>&lt;0.00001</b> | 58% (moderate)                 |
| VAS after BMAC treatment | 4             | 36.06 (28.35, 43.76)     | <b>&lt;0.00001</b> | 76% (moderate)                 |
| Pfirmann improvement     | 2             | 0.44 (0.13, 0.75)        | <b>0.005</b>       | 0% (low)                       |

Abbreviations: BMAC, bone marrow aspirate concentrate; ODI, Oswestry Disability Index; VAS, visual analog scale.

Note: Boldface indicates statistical significance.

**Figure 3.** Oswestry Disability Index forest plot.

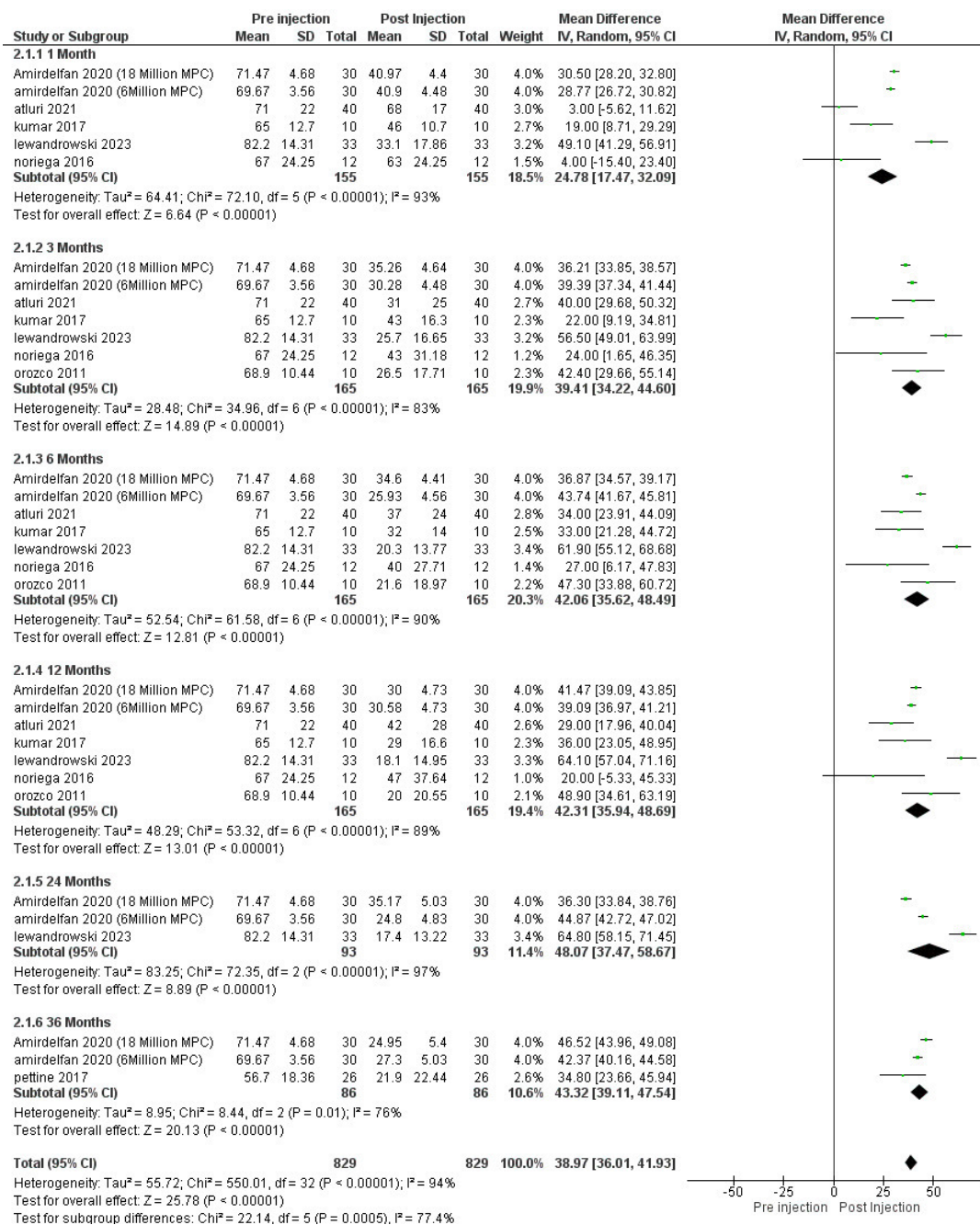


Figure 4. Visual analog scale forest plot.

statistically significant improvement in Pfirrmann grade ( $P = 0.005$ ) with low heterogeneity ( $I^2 = 0\%$ ; Table 3; Figure 7). Other studies showed varied results. In the study by Pettine et al, 8 of 20 patients showed improved Pfirrmann grade. In the study by Noriega et al,<sup>21</sup> the control group had worsened Pfirrmann grade at the end of the follow-up. Although limited, these results suggest the potential of stem cell therapy to influence disc morphology, warranting further investigation with larger samples.

## DISCUSSION

Regenerative treatments using stem cell therapy sourced from the human body are gaining popularity across medical fields. In orthopedics, it offers a promising alternative for patients with degenerative conditions who opt to avoid surgery. This therapy aims to stimulate the body's natural regenerative mechanisms, especially in the IVDs, potentially alleviating pain in patients affected by degenerative disc disease over time.

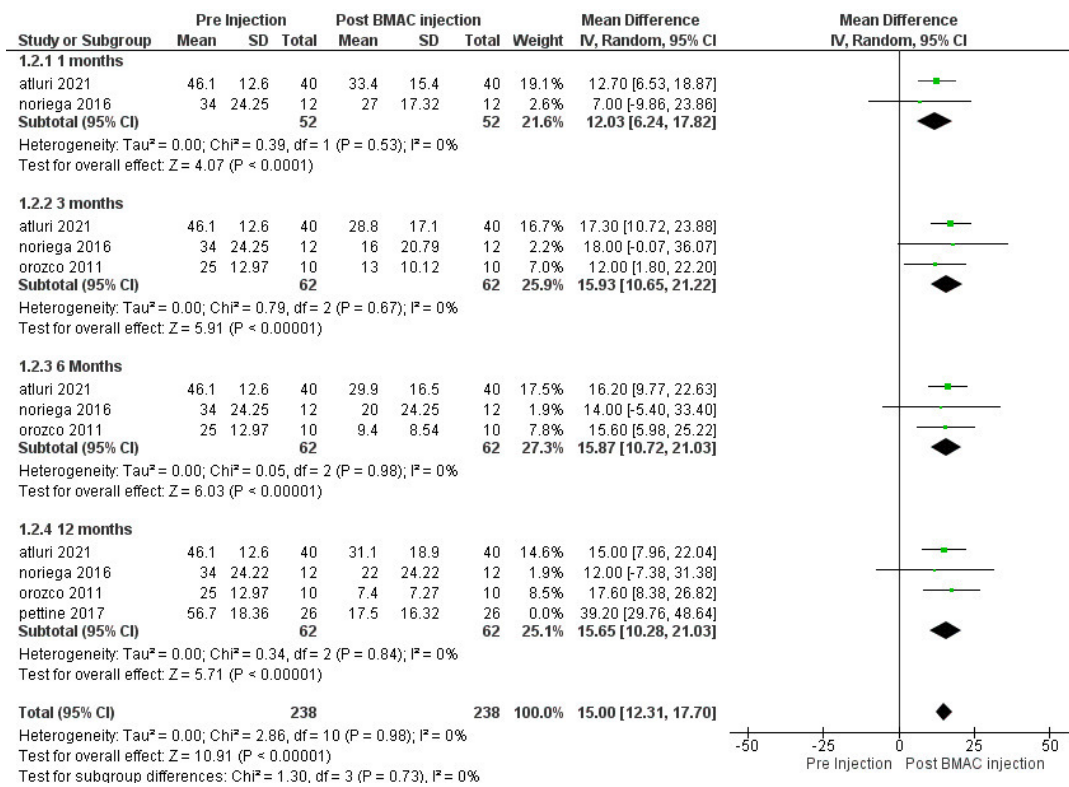


Figure 5. Oswestry Disability Index improvement with bone marrow aspirate concentrate.

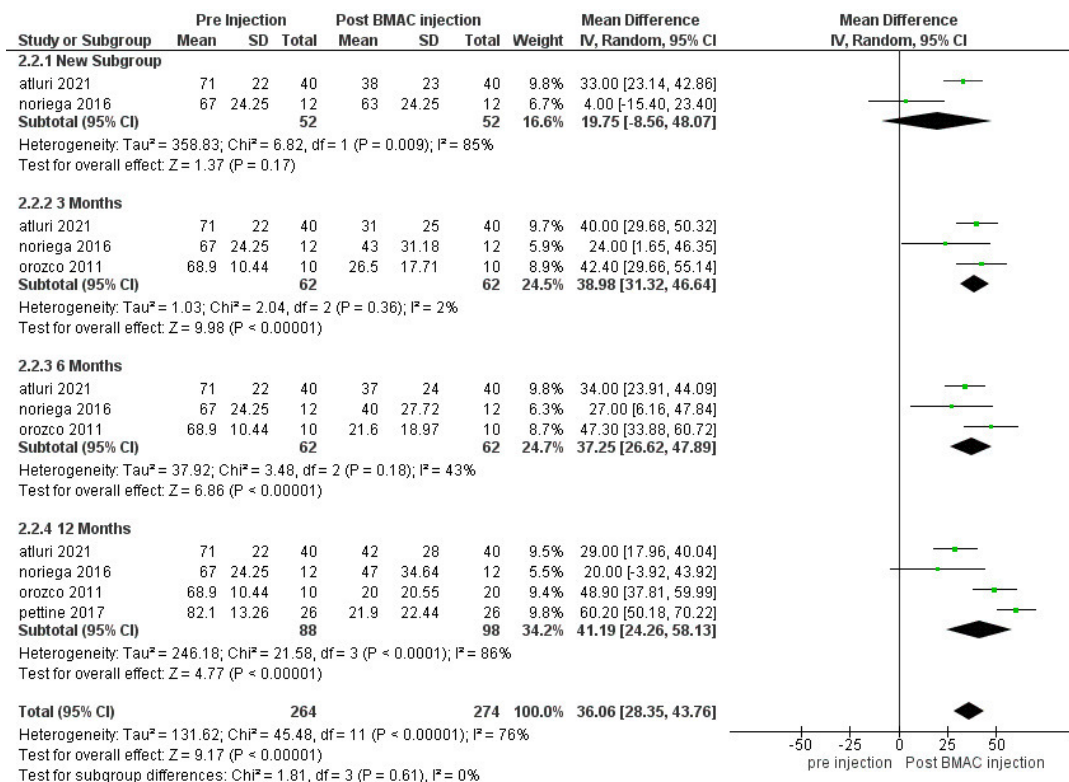


Figure 6. Visual analog scale improvement with bone marrow aspirate concentrate.

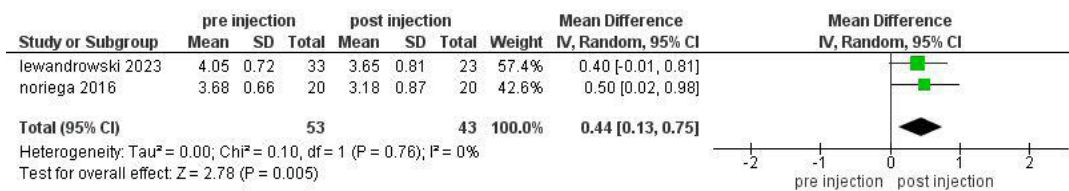


Figure 7. Pfirrmann improvement.

Regenerative medicine therapies have shown potential for sustained relief and, in certain cases, may provide curative outcomes. In recent years, allogeneic stem cell transplantation has gained increasing interest as an alternative to autologous BMC transplantation.<sup>15</sup> Another source of mesenchymal stem cells (MSCs) is BMC, which can be derived from either autologous or allogeneic bone marrow aspirate. BMC can be harvested from the posterior superior iliac spine of the patient<sup>16–18</sup> or obtained from an allogeneic donor and subsequently processed following good manufacturing practice standards.<sup>21</sup>

Multiple studies<sup>3,4,9–11,23–25</sup> investigating the potential of MSC therapy for degenerative disc disease have consistently concluded that MSCs may offer a promising treatment approach for this condition. Although the precise mechanisms by which MSCs can alleviate disc degeneration remain unclear, evidence suggests that intradiscal injection of MSCs could serve as a potential treatment option for patients with chronic LBP. MSCs are considered strong candidates for regenerating IVDs, as they aim to replenish disc tissue and rejuvenate its functionality by promoting matrix synthesis through the implanted cells. Furthermore, MSCs may exert positive effects on the surrounding native cells.<sup>26</sup> Animal studies have shown that MSCs injected into the nucleus pulposus not only survive but also proliferate, potentially leading to improvements in the condition of degenerated discs.<sup>4</sup>

This systematic review and meta-analysis evaluated the effects of stem cell therapy on VAS, ODI, and disc morphology (Pfirrmann grade) in patients with discogenic LBP. Across studies, stem cell therapy showed statistically significant improvement in VAS/Numeric Rating Scale, ODI, and Pfirrmann improvement, suggesting substantial benefits for pain relief and functional outcomes in patients with discogenic LBP. High heterogeneity in overall analyses indicates variability in study outcomes, likely due to differences in intervention types, follow-up periods, and patient populations. Subgroup analyses by follow-up period and intervention type (bone marrow aspirate concentrate) reduced heterogeneity to some extent, suggesting the effects

of stem cell therapy on disability may be durable over time.

The magnetic resonance imaging findings, although limited in scope, provide preliminary evidence that stem cell therapy may positively affect disc degeneration, as indicated by changes in Pfirrmann grade. However, varied results across studies underscore the need for standardization in imaging protocols and extended follow-up to ascertain the true impact on disc morphology. These findings, while promising, are derived from a small subset of studies, highlighting the need for more robust evidence with larger, standardized sample sizes.

Wu et al,<sup>24</sup> Zhang et al,<sup>10</sup> and Yolcu et al<sup>11</sup> reported improvements in discogenic LBP after stem cell therapy. Yolcu et al observed improvement at 3 months, 6 months, and 12 months, although the quantitative gains were slightly lower at the 12-month follow-up. In contrast, Wu et al and Zhang et al compared only the baseline data with final follow-up outcomes.

The observed outcomes may be linked to the introduction of anabolic growth factors and stem cells into the degenerated IVD, effectively counteracting its catabolic environment. Increased levels of growth factors and cytokines have been shown to enhance cellular proliferation of both annulus fibrosus and nucleus pulposus cells, enhance glycosaminoglycan content, stimulate collagen synthesis, and upregulate gene expression related to extracellular matrix proteins that are essential for IVD function. This synergistic effect likely plays a significant role in improving the overall condition of the disc.<sup>16</sup>

Sanapati et al<sup>9</sup> also stated in their study that regenerative therapies, including MSCs, may be effective in treating discogenic LBP, with the potential to decelerate or even halt the degenerative process of the IVD. However, they also suggested that the effectiveness of MSC injections could be improved by combining them with growth factors present in platelet-rich plasma.

Although no adverse events were reported following MSC injections, Meisel et al<sup>12</sup> emphasized that evidence regarding the efficacy and safety of cell therapy remains limited due to potential biases and small sample sizes. Similarly, Schneider et al<sup>3</sup> found that the quality

of evidence for the effectiveness of intradiscal biological treatments was very low. This finding highlights the need for further research, particularly focusing on the efficacy and safety of MSC injections.

The authors acknowledge the limitations of this study. Variations in stem cell dose and cell types may influence the findings. Diverse reporting tools and follow-up times posed challenges in harmonizing results, and the small sample sizes in some studies may have contributed to bias. Different standards for patient selection across studies may potentially lead to a good outcome bias. Studies included in this review clearly stated no direct funding for their research, but several authors were affiliated with companies that manufactured the device used for their interventions, which may introduce a potential motivational bias. Future studies could benefit from more disclosure on author affiliations and potential conflicts of interest to further enhance objectivity.

Further research is necessary to compare the outcomes of stem cell therapy with standard treatments in long-term follow-up studies. Additionally, the optimal dose and variations in stem cell types required further investigation, as did establishing consistent patient selection criteria to ensure more accurate assessments of treatment efficacy.

## CONCLUSION

Stem cells therapy could be beneficial as an option for the treatment for discogenic LBP in improving pain and activity of daily living. Future prospective studies with control subjects as a comparison of the effectiveness are strongly recommended to be conducted to verify this finding and also explore more effects regarding the use of stem cells.

## REFERENCES

1. Salter RB. *Textbook of Disorders and Injuries of the Musculoskeletal System: An Introduction to Orthopaedics, Fractures, and Joint Injuries, Rheumatology, Metabolic Bone Disease, and Rehabilitation*. 3rd ed. Williams & Wilkins; 1999.
2. Apley SL. *Apley & Solomon's System of Orthopaedics and Trauma*. 10th ed. (Blom A, Warwick D, Whitehouse M, eds.). CRC Press; 2018.
3. Schneider BJ, Hunt C, Conger A, et al. The effectiveness of intradiscal biologic treatments for discogenic low back pain: a systematic review. *Spine J*. 2022;22(2):226–237. doi:10.1016/j.spinee.2021.07.015
4. Soufi KH, Castillo JA, Rogdriguez FY, DeMesa CJ, Ebinu JO. Potential role for stem cell regenerative therapy as a treatment for degenerative disc disease and low back pain: a systematic review. *Int J Mol Sci*. 2023;24(10):8893. doi:10.3390/ijms24108893
5. Ganesan S, Acharya AS, Chauhan R, Acharya S. Prevalence and risk factors for low back pain in 1,355 young adults: a cross-sectional study. *Asian Spine J*. 2017;11(4):610–617. doi:10.4184/asj.2017.11.4.610
6. Ferreira ML, de Luca K, Haile LM. Global, regional, and national burden of low back pain, 1990–2020, its attributable risk factors, and projections to 2050: a systematic analysis of the global burden of disease study 2021. *Lancet Rheumatol*. 2023;5(6):e316–e329. doi:10.1016/S2665-9913(23)00098-X
7. Makkiyah FA, Sinaga TA, Khairunnisa N. A study from a highly populated country: risk factors associated with lower back pain in middle-aged adults. *J Korean Neurosurg Soc*. 2023;66(2):190–198. doi:10.3340/jkns.2021.0278
8. Yudawijaya A, Hayati Mutmainah RR, Pariama G. Profile of low back pain patients at the neurology polyclinic. *Int J Health Sci Res*. 2022;12(12):187–199. doi:10.52403/ijhsr.20221228
9. Sanapati J, Manchikanti L, Atluri S, et al. Do regenerative medicine therapies provide long-term relief in chronic low back pain: a systematic review and metaanalysis. *Pain Physician*. 2018;21(6):515–540.
10. Zhang W, Wang D, Li H, et al. Mesenchymal stem cells can improve discogenic pain in patients with intervertebral disc degeneration: a systematic review and meta-analysis. *Front Bioeng Biotechnol*. 2023;11:1155357. doi:10.3389/fbioe.2023.1155357
11. Yolcu YU, Moinuddin FM, Wahood W, Alvi MA, Qu W, Bydon M. Use of regenerative treatments in treatment of lumbar degenerative disc disease: a systematic review. *Clin Neurol Neurosurg*. 2020;195:105916. doi:10.1016/j.clineuro.2020.105916
12. Meisel H-J, Agarwal N, Hsieh PC, et al. Cell therapy for treatment of intervertebral disc degeneration: a systematic review. *Glob Spine J*. 2019;9(1 Suppl):39S–52S. doi:10.1177/2192568219829024
13. Page MJ, McKenzie JE, Bossuyt PM, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. *BMJ*. 2021;372:n71. doi:10.1136/bmj.n71
14. Slim K, Nini E, Forestier D, Kwiatkowski F, Panis Y, Chipponi J. Methodological index for non-randomized studies (minors): development and validation of a new instrument. *ANZ J Surg*. 2003;73(9):712–716. doi:10.1046/j.1445-2197.2003.02748.x
15. Lewandrowski KU, Dowling A, Vera JC, Leon JFR, Telefean AE, Lorio MP. Pain relief after allogenic stem cell disc therapy. *Pain Physician*. 2023;26(2):197–206.
16. Wolff M, Shillington JM, Rathbone C, Piasecki SK, Barnes B. Injections of concentrated bone marrow aspirate as treatment for discogenic pain: a retrospective analysis. *BMC Musculoskelet Disord*. 2020;21(1):135. doi:10.1186/s12891-020-3126-7
17. Pettine KA, Suzuki RK, Sand TT, Murphy MB. Autologous bone marrow concentrate intradiscal injection for the treatment of degenerative disc disease with three-year follow-up. *Int Orthop*. 2017;41(10):2097–2103. doi:10.1007/s00264-017-3560-9
18. Atluri S, Murphy MB, Dragella R, et al. Evaluation of the effectiveness of autologous bone marrow mesenchymal stem cells in the treatment of chronic low back pain due to severe lumbar spinal degeneration: a 12-month, open-label, prospective controlled trial. *Pain Physician*. 2022;25(2):193–207.
19. Orozco L, Soler R, Morera C, Alberca M, Sánchez A, García-Sancho J. Intervertebral disc repair by autologous mesenchymal bone marrow cells: a pilot study. *Transplantation*. 2011;92(7):822–828. doi:10.1097/TP.0b013e3182298a15
20. Kumar H, Ha D-H, Lee E-J, et al. Safety and tolerability of intradiscal implantation of combined autologous adipose-derived mesenchymal stem cells and hyaluronic acid in patients with chronic

discogenic low back pain: 1-year follow-up of a phase I study. *Stem Cell Res Ther.* 2017;8(1):262. doi:10.1186/s13287-017-0710-3

21. Noriega DC, Ardura F, Hernández-Ramajo R, et al. Intervertebral disc repair by allogeneic mesenchymal bone marrow cells: a randomized controlled trial. *Transplantation.* 2017;101(8):1945–1951. doi:10.1097/TP.0000000000001484

22. Amirdelfan K, Bae H, McJunkin T, et al. Allogeneic mesenchymal precursor cells treatment for chronic low back pain associated with degenerative disc disease: a prospective randomized, placebo-controlled 36-month study of safety and efficacy. *Spine J.* 2021;21(2):212–230. doi:10.1016/j.spinee.2020.10.004

23. Urits I, Capuco A, Sharma M, et al. Stem cell therapies for treatment of discogenic low back pain: a comprehensive review. *Curr Pain Headache Rep.* 2019;23(9):65. doi:10.1007/s11916-019-0804-y

24. Wu T, Song HX, Dong Y, Li JH. Cell-based therapies for lumbar discogenic low back pain: systematic review and single-arm meta-analysis. *Spine (Phila Pa 1976).* 2018;43(1):49–57. doi:10.1097/BRS.0000000000001549

25. Zeckser J, Wolff M, Tucker J, Goodwin J. Multipotent mesenchymal stem cell treatment for discogenic low back pain and disc degeneration. *Stem Cells Int.* 2016;2016. doi:10.1155/2016/3908389

26. Richardson SM, Kalamegam G, Pushparaj PN, et al. Mesenchymal stem cells in regenerative medicine: focus on articular cartilage and intervertebral disc regeneration. *Methods.* 2016;99:69–80. doi:10.1016/j.ymeth.2015.09.015

**Funding:** The authors received no financial support for the research, authorship, and/or publication of this article.

**Declaration of Conflicting Interests:** The authors report no conflicts of interest in this work.

**Author Contributions:** R., K.Y., A.G., and N.P.H. contributed to conceptualization, analysis, and critical review of the study. R., K.Y., and A.G. searched the literature and wrote the manuscript. N.P.H. reviewed and supervised the study.

**Corresponding Author:** Randy Randy, Emergency Department, Mayapada Hospital Kuningan, Palembang 5, A3 no.8, Cengkareng, West Jakarta 11830, Indonesia; dr.randysamsul@gmail.com

Published 08 January 2025

Copyright © 2025 ISASS. The IJSS is an open access journal following the Creative Commons Licensing Agreement CC BY-NC-ND. To learn more or order reprints, visit <http://ijssurgery.com>.