Review Article:
Immunomodulatory Effect of Mesenchymal Stem Cells in Multiple Sclerosis and Experimental Autoimmune Encephalomyelitis: A Review Study

Shaghayegh Pezeshki Naraghi1, Seyed Mahmoud Hashemi1*

1. Department of Immunology, School of Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran.

ABSTRACT

Multiple Sclerosis (MS) is a chronic inflammatory and demyelinating disease of the central nervous system that may lead to disability of the patient. Current MS treatment regimens are still insufficient and research is conducted for developing more effective therapies capable of targeting neurodegeneration, inflammation, and demyelination. Recent results of experimental and clinical studies in cell-based therapy suggest Mesenchymal Stem Cells (MSCs) as potential candidates in autoimmune disorders. Immunomodulatory and neuroprotective properties of mesenchymal stem cells might offer the valuable therapeutic potential to modulate inflammatory responses associated with MS and repairing damaged nervous tissue. This review summarizes the main experimental studies and clinical trials using MSCs in MS.

Introduction

Multiple Sclerosis (MS) is an autoimmune and common disease that affects central nervous system in humans [1]. It is a chronic disease more common in women and some parts of the world such as Northern Europe, America and Canada [2]. In MS, myelin that covers the nerves and is crucial for nerve transmission [3] is destroyed. Those parts of the nerve that lost their myelin cover due to MS is called plates. With the expansion of the plates, transmission of neural messages along the nerve slows down creating MS symptoms [4]. Symptoms usually start with unexplained and vague fatigue followed by general weakness. However, this weakness may occur only in one leg or one arm. Some patients experience blurred or double
vision (diplopia). Moreover, symptoms like numbness and prickling of the face, arms, legs, and the whole body; balance disorder; chills; confusion; difficulty in swallowing or trouble speaking; and loss of bladder control can be seen in patients with MS [5].

From immunology perspective, because of an unknown factor myelin antigens are initially delivered to the T cells by the antigen-presenting dendritic cells in the lymphatic system, causing proliferation of reactive and active T cells in the lymph nodes [6]. Then the reactive T cells pass the blood brain barrier and enter into the central nervous system. Here they react with myelin antigens and produce inflammatory cytokines and chemokines, which in turn stimulate further migration of the immune cells to the site and progressive destruction of myelin in the central nervous system, causing sclerosis plaques in the brain and spinal cord [7, 8].

Various Methods for Classifying MS

According to the latest classification, four different courses are identified for the MS disease: 1. Relapsing Remitting MS (RRMS) [9]; 2. Secondary Progressive MS (SPMS) [10]; 3. Primary Progressive MS (PPMS) [11]; and 4. Progressive Relapsing MS (PR) [12].

Immunotherapy and Cell Therapy in MS

Intravenous Immunoglobulin

In cases where the attack does not have an appropriate response to corticosteroid therapy, the doctor may use Intravenous Immunoglobulin (IVIG) [13, 14].

Plasmapheresis

Plasmapheresis is used in cases of disease attacks that do not give a favorable response to corticosteroid therapy. In this type of treatment, a given volume of blood is replaced during 3 to 4 days [15]. Some medications are in this category that can reduce the number of attacks and slow down the course of the disease. The following drugs are in this group:

Interferons

Interferons proteins are produced naturally in the body and defend the body against viruses. There are three types of natural interferons: alpha, beta, and gamma; of which beta has a treatment role in MS. This type of interferon exists in the market in three injection forms [16]:

- 1. Interferon beta-1a including Avonex (Sinnovex) [17];
- 2. Rebif (Resijen) [18];
- 3. Interferon beta-1b (Betaferon) [19].

Glatiramer acetate (brand Copaxone or Copamer and Isomer)

Glatiramer acetate is currently used only in MS. This drug is a chain of four molecules of amino acids called glutamic acid, lysine, alanine, and tyrosine. It is naturally found in myelin protein molecules (nerve sheath). Thus, the drug is similar to the myelin around the nerve in terms of chemical structure. The drug is used by the American Food and Drug Administration (FDA) since 1996 to reduce the number of attacks of MS, although the observations show its effectiveness in preventing the progression of disability. This medicine is used daily as a subcutaneous injection [20].

The mechanism of drugs action is not well understood, but it seems that it inhibits inflammatory reactions, and perhaps due to its similarity to the myelin sheath, it perverts the immune responses generated against this sheath [18]. In studies where the drug was compared with Rebif and Betaferon (BEYOND), no difference was found between this drug and others in terms of the effectiveness [21]. In addition to definite MS forms, the drug can be also used to cure Clinically Isolated Syndrome (CIS) (first attack of MS). However, it has no effect in progressive forms (primary and without attack) [22]. The drugs like interferon beta can be used for the disease attack forms with low failure rate.

Mitoxantrone

Mitoxantrone is usually among the chemotherapy drugs that, in certain cases of the disease, is injected once every three months. It usually does not continue more than two years. Until 2006, these drugs were the main treatment of MS [23, 24]. Since then, newer drugs were added to the treatment list. These include:

Natalizumab (Tysabri)

Natalizumab was confirmed by both American and Europe (EU), FDA in 2006. It has a very good effect in controlling MS and used once a month as intravenous injection. Unfortunately, due to its lethal complication called progressive multifocal leukoencephalopathy, the drug should be used with close monitoring and in the context of specific systems [25].
Fingolimod (Gilenya)

Fingolimod is the first oral medication for MS disease that has received approval from FDA in 2010, and has been licensed by the same organization in Europe in April 2011. The drug, taken by mouth daily, alleviates the number of attacks and slows down the disease progression [26, 27].

Here are two essential things. First, all of these drugs can control MS disease and do not have the ability to eradicate it, and second, a large group of medicines are now under investigation; some of them certainly will pass the tests successfully in the coming years [28].

Introduction of Mesenchymal Stem Cells

Mesenchymal Stem Cells (MSCs) have been identified as factors regulating the immune system [29], and the immunomodulatory role of these cells on a variety of immune system cells such as T and B lymphocytes [30][31], Natural Killer (NK) cells [32], macrophages [33] and Dendritic Cells (DCs) [34] has led to their use in the treatment of inflammatory diseases, particularly autoimmune diseases such as MS [35] and rheumatoid arthritis [36]. Bone marrow is the main source of mesenchymal stem cells [37], but they are isolated from other body tissues including adipose tissue [38]. Since T lymphocytes are considered as the most important cellular immune system cells, MSCs effect on the activity of these cells is of the utmost importance in directing the immune response [39].

T lymphocytes have different phenotypes and function in different conditions. According to prevailing conditions, they are distinct to certain subgroups with their special function and phenotype [40]. MSCs result in distinction of T cells from T regulatory cells that play an important role in maintaining tolerance and preventing autoimmune diseases [31]. Mesenchymal stem cells were first used to create the bone cells in the bone marrow in 1960. Later, researchers found that these cells, in addition to the ability to differentiate into bone, cartilage and other connective tissues in laboratory conditions, can be isolated from a variety of sources including placenta, umbilical cord, adipose tissue and teeth [41].

Mesenchymal stem cell do not have the surface hematopoietic cell markers including cluster of differentiation (CD) 45, CD34, and CD14. But they express specific molecules such as SH2 (CD105), SH3, SH4 (CD73), CD44, CD22, intercellular adhesion protein (ICAM) -1 (CD54) and vascular cell adhesion protein (VCAM) -1 (CD106) as well as CD13, CD90, CD166 and STR-1. Also they are negative in terms of having CD40, CD80 and CD86 and express low levels of human leukocyte antigen (HLA) -I and LFA-3; however, They do not express HLA-III [42]. These stem cells have the immunomodulatory property; this property applies on B, T, NK and DC cells and reduces the immune response [31, 30].

MSCs Immunomodulatory Effects

MSCs have immunoregulatory role both in the innate and adaptive immune system. The unique properties of these cells suppress and modulate immune responses [43, 42]. MSCs reduce NKGD2D, NKP44 and NKP30 on the surface of Natural Killer (NK) cells and lead to reduced production of interferon gamma (INFγ) in these cells [32]. As said previously, they have an inhibitory effect on neutrophil cells and prevent the production of hydrogen peroxide by activated neutrophils [44]. Many studies indicate that MSCs have an inhibitory effect on proliferation of T cell induced by polyclonal mitogens, allogeneic cells or specific antigens [45]. These cells do their inhibitory effect on the proliferation of lymphocytes by stopping the cells in the G1/G0 phase of cell cycle [46].

Due to the secretion of their cytokines, MSCs are capable of directing differentiation of T lymphocytes subsets to reduce the secretion of proinflammatory cytokines (IFNγ, Tumor Necrosis alpha [TNFα] and Interleukin [IL] -6 and IL-17) produced by activated T cells in the extracellular environment [47], and in contrast, secretion of anti-inflammatory cytokines (IL-4 and IL-10) increases [45]. T+CD4 lymphocytes have few main subgroups or subsets, each of which has its own specific phenotypic and performance characteristics; among these subsets, Th1 and Th17 are identified as inflammatory and directing inflammatory pathways subsets. On the other hand, Th2 and regulatory T cells (Treg) subsets are considered as a population of immune system suppressor or regulator cells [48, 49]. Th1 cells are the main cells in acute inflammatory reactions [50]. Abundant evidence has shown that MSCs reduce the production of INFγ by Th1 and lead to the inhabitation of this subset [51]. MSCs isolated from the bone marrow also inhibit allergic inflammation of the airways caused by Th2 cells by inducing IFNγ production [52].

Th17 cells produce IL-17A, IL17-F and IL-6, and are involved in autoimmunity and inflammation of the tissue [51]. Th17 cells as subsets of Th1 and have a central role in the process of acute graft rejection mediated by the cells, and with the production of inflammatory cytokines
decreases the survival of the transplanted organ in the host body [53]. Various reports regarding the effect of MSCs on the Th17 cells have shown that MSCs inhibit Th17 response and inhibit inflammation caused by these cells, while other studies indicate that the MSCs induce and enhance Th17 response [54, 55].

Regulatory T cells (Treg) suppress and inhibit immune responses, and have an important role in maintaining tolerance and preventing autoimmune diseases [56]. Treg cells produce large amounts of IL-10 and Transforming Growth Factor (TGF)-β. MSCs lead to differentiation of T naive cells to Treg cells with high production capability of IL-10 and TGF-β [57]. On the other hand, with suppression of Th1 response and decreasing the production of INF-γ, the result is intense inhabitation of immune response [58]. Furthermore MSCs also cause adsorption and proliferation of Treg cells, and consequently, result in suppression of immune responses [59]. Therefore, the immunosuppressive characteristic of MSCs is currently used in the treatment of many autoimmune diseases such as MS [60], rheumatoid Arthritis [36], Type 1 diabetes [61], allergic diseases [62] and other hypersensitivities. It can also be used in organ transplantation as specific immunity suppressor to increase graft survival and coping T regulatory responses toward other lymphocyte subsets [63].

MSCs with secretion of indoleamine 2, 3-dioxygenase can catalyze an essential amino acid called tryptophan to kynurenine, and impair the synthesis of various cellular proteins leading to inhibition of cell proliferation [64]. They inhibit the differentiation and function of B cells, too. This effect plays its role by reducing the chemokine receptors CXCR4, CXCR5, and CCR7 [65].

Because of MSCs properties, especially their immunomodulatory function, they have been used as an immunomodulator for the treatment of diseases caused by the immune system. In recent years, the MSC therapy on the models of autoimmune diseases has been investigated [60].

**MSCs Immunomodulatory Effect in Autoimmune and Inflammatory Diseases**

Systemic injection of MSCs isolated from the bone marrow to a rat model (Experimental autoimmune encephalomyelitis, EAE) reduces the symptoms and severity of the disease (Table 1), suppresses demyelination in the brain and spinal cord, slows down the arrival of macrophages and T lymphocytes into the Central Nervous System (CNS) tissue, and reduces the antibody production against pathogenic protein-lipid proteins (one of the components of myelin) compared with the control group [66, 67]. In addition, the risk of a relapse and demyelination is reduced as compared with the control group. With the studies on the migration characteristics of MSCs in the rat model of EAE, it was found that MSCs remain in the lymph organs and the CNS inflamed areas.
several weeks after systemic injection [68]. In addition to the immunomodulatory activity, MSCs have a protective effect on neurons by secreting nutritional factors and call for tissue precursor cells and their differentiation into different types of nerve cells [39, 69].

In connection with rheumatoid arthritis, the medical function of MSCs in the animal model of rheumatoid arthritis was studied and it was revealed that clinical symptoms and inflammatory cytokines decreases, but the anti-inflammatory cytokines in the lymph nodes and joints increases, and thus preventing severe damages to the tissues [36]. Studies on the MSCs therapeutic activity on the experimental models of muscle weakness (Myasthenia) showed that clinical signs and proliferation of specific lymphocytes of acetylcholine receptor (AchR) decreases, and the body weight in mice and rats with myasthenia is increases [70].

The rat model of Systemic Lupus Erythematosus (SLE) improved the illness in the models, and allogeneic MSC infusion reduces serum antibody, glomerular IgG level, and proteinuria, but increases bone formation and restores osteoblast niche [71].

**MS and Mesenchymal Stem Cells**

There are different types of stem cells: hematopoietic stem cells, mesenchymal cells, and neural cells that are used in the treatment of MS. Of which hematopoietic stem cells and mesenchymal cells are commonly used [75]. There are three ways to inject these cells: intravenous injection, spinal injection or tissue injection. Of course, the immunity of tissue injection has not yet been proved because a large amount of stem cells enter the body; it has also the risk of developing tumors in the tissues. For this reason, spinal and intravenous injections are done frequently [33, 76].

**Stem Cell Injections**

According to Figure 2, mesenchymal stem cells intravenously enter into the lungs. Many of these cells are destroyed by the macrophages of the lungs, and only a limited number of cells enter into the peripheral lymph nodes, heart, and brain [77].

**Injecting Mesenchymal Stem Cells in EAE Rats**

Experimental Autoimmune Encephalomyelitis (EAE) mouse is an animal model suitable for MS studies. In terms of clinical signs and tissue damages, it can well imitate MS because, in this model, the chronic neurologic disease is created along with the phases of attack and suppress, and like MS, hard sclerosis plaques are formed in the central nervous system [48, 49].

---

**Table 1. Systemic injection of MSCs isolated from the bone marrow to a rat**

<table>
<thead>
<tr>
<th>Disease</th>
<th>Animal Model</th>
<th>Mesenchyme Type</th>
<th>Result</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myasthenia Gravis</td>
<td>Rat</td>
<td>Human bone marrow</td>
<td>Improved symptoms and reduced acetylcholine level</td>
<td>[70]</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>Rat</td>
<td>Human adipose tissue</td>
<td>Improved clinical symptoms, Reduced inflammatory cytokines in the joints</td>
<td>[36]</td>
</tr>
<tr>
<td>Diabetes</td>
<td>Rat</td>
<td>Human bone marrow</td>
<td>Increased number and size of islets of Langerhans</td>
<td>[72]</td>
</tr>
<tr>
<td>Colitis</td>
<td>Rat</td>
<td>Rat bone marrow</td>
<td>Increased number and size of islets of Langerhans</td>
<td>[73]</td>
</tr>
<tr>
<td>Systemic lupus erythematosus</td>
<td>Rat</td>
<td>Rat bone marrow</td>
<td>Reduced rickets, and bone cell niche repair, and improved performance of damaged organisms</td>
<td>[71]</td>
</tr>
<tr>
<td>MS</td>
<td>Rat</td>
<td>Rat bone marrow</td>
<td>Improved symptoms reduced inflammatory cell infiltration and reduced demyelination</td>
<td>[46]</td>
</tr>
<tr>
<td>MS</td>
<td>Rat</td>
<td>Human bone marrow</td>
<td>Delayed the outbreak of disease, and improved symptoms</td>
<td>[73]</td>
</tr>
</tbody>
</table>
Measurement of IL4, IL10 and IL17 CytoKines in EAE Rat With MSC Injection

In this study, the rate of important cytokines IL-4 (TH2 response), IL-17FA (inflammatory response) and IL-10 (anti-inflammatory response) has been studied. Production of IL-4 and IL-10 cytokines from mouse spleen cells has increased while the production of inflammatory cytokines IL-17AF decreased [66, 78].

In fact, these findings indicate that mesenchymal stem cells migrate from the immune cells into the CNS tissue and thus prevent further destruction of myelin and suppress the inflammatory response by increasing anti-inflammatory IL-10 and IL-4 cytokines [79, 80]. Furthermore, mesenchymal stem cells, by increasing IDO1 (Indoleamine 2, 3-dioxygenase 1), ADP (antioxidant enzymes), ribose polymerase-1 and P53, oligodendrocyte progenitor cells, reduce the migration of inflammatory cells into the CNS and also induces remyelination [64, 81, 82]. Also the pathologist examination on stained samples shows the reduced number of immune cells infiltrating into the nerve tissue [39, 83].

Because of their immunomodulatory and neuroprotective properties, mesenchymal cells hold great promises in the treatment of autoimmune diseases such as MS. Studies on animals show the positive impact of MSCs in reducing the symptoms of EAE (Table 2) [84, 85]. Transplanting human adipose-derived MSCs (AdMSCs) expressing the anti-inflammatory cytokine Interleukin (IL)-4 in EAE causes reduction in peripheral MOG-specific T-cell responses and a shift from pro- to anti-inflammatory cytokine response [86].

In another study using human bone marrow-derived mesenchymal stem cells (hBM-MSCs) to secrete IFN-β (MSCs-IFNβ), the MSCs-IFNβ treatment enhanced the immunomodulatory effects. This, in turn, suppresses proinflammatory cytokines (IFN-ϒ and TNF-α) and conversely increases anti-inflammatory cytokines (IL-4 and IL-10). Importantly, the injected MSCs-IFNβ migrates into the inflamed CNS and significantly reduces further injury of the Blood-Brain Barrier (BBB) permeability in EAE mice [87]. Moreover, Payne, N. L. et al. engineered the expression of IL-10 in human adipose-derived mesenchymal stem cells (Adi-IL-10-MSCs) and transplanted these cells early in the disease course to mice with EAE. They showed that when administered at the time of CD4+T-cell priming, Adi-MSCs overexpressing human IL-10 significantly attenuates EAE by modulating APC function and inhibiting Th17-mediated neuro-inflammation [88].

In a recent study, a comparative analysis of the immunomodulatory properties of Adipose Tissue Mesenchymal stem cells (AT-MSCs) and their Conditioned Media (CM), derived from C57/BL6 mice, was done for mitigating the adverse clinical course of EAE. There was no significant difference in the clinical scores and the body weight of EAE mice treated with AT-MSCs and CM. The reduction in proliferative responses and brain cell infiltration was more pronounced in the mice injected with CM than in other groups. It was found that the percentage of splenic CD4+CD25+FOXP3+population and the level of IL4 production in the mice administrated with AT-MSCs more increased compared to other animals [89].

In another study, they compared the immune regulatory properties of adipose tissue MSCs (AT-MSCs) in two independent routes of injection; namely Intraperitoneal...
<table>
<thead>
<tr>
<th>Disease</th>
<th>Injection Way</th>
<th>Cell Source</th>
<th>Result</th>
<th>Cause</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Syngeneic (C57BL6)</td>
<td>Syngeneic (C57BL6) Environmental toleranceReduction of inflammation CNS</td>
<td>Decreased INFγ Decreased TNFα</td>
<td>[84]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Xenogenic (Human)</td>
<td>Environmental toleranceReduction of inflammation CNS</td>
<td>Induced oligodendrocyte progenitor cells (NG2)</td>
<td>[85]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Xenogenic (Human)</td>
<td>Neuroprotection</td>
<td>Production of NGF</td>
<td>[85]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Allogeneic (C57BL6 SJL)</td>
<td>Neuroprotection Environmental tolerance</td>
<td>Inhibition of Th17, Th1 and NK cells production</td>
<td>[78]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intraventricular</td>
<td>Syngeneic (C57BL6)</td>
<td>Neuroprotection (reduced inflammation) and neuroprotection</td>
<td>Immunomodulatory and reducing inflammatory cytokine</td>
<td>[39]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intraperitoneal</td>
<td>Syngeneic (C57BL6)</td>
<td>Immunomodulatory</td>
<td>Reduced inflammatory cytokines Th17</td>
<td>[67]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Syngeneic adipose derived MSC (C57BL6)</td>
<td>Immunomodulatory and neuroprotection</td>
<td>Increased cytokines type Th2Decreased Th17</td>
<td>[79]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Xenogenic (Human)</td>
<td>Immunomodulatory and neuroprotection</td>
<td>Decreased Th17Increased IL4</td>
<td>[80]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intraperitoneal</td>
<td>Syngeneic (C57BL6)</td>
<td>Immunomodulatory</td>
<td>Increased IDO1Indoleamine 2, 3-dioxygenase 1</td>
<td>[64]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Syngeneic (C57BL6)</td>
<td>Neuroprotection</td>
<td>Activities of ADP antioxidant enzymes, Ribose polymerase-1 and P53</td>
<td>[82]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intraperitoneal</td>
<td>AllogeneicBALB (C57BL)</td>
<td>Immunomodulatory</td>
<td>Increase INFγ Reduced IL-17</td>
<td>[92]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intraperitoneal</td>
<td>Immunomodulation and neuroprotection</td>
<td>Xenogenic (Human)</td>
<td>Decreased inflammation</td>
<td>[83]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Adipose-derived MSCs (Ad-MSCs) engineered to overexpress the anti-inflammatory cytokine interleukin (IL-4)</td>
<td>Immunomodulation and neuroprotection</td>
<td>Reduction in peripheral MOG-specific T-cell responses and a shift from a pro- to anti-inflammatory cytokine response</td>
<td>[86]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Interferon β-secreting human bone marrowmesenchymal stem cells</td>
<td>Suppressing demyelination and immunomodulatory effects</td>
<td>Promoting a shift from the Th1 to the Th2 cytokine balance</td>
<td>[87]</td>
</tr>
</tbody>
</table>
(IP) and Intravenous (IV), and reported that the IP route has a more pronounced effect in maintaining the splenic CD4+CD25+FOXP3+T cell population and increase of IL-4 secretion. They also showed that IP injection of cells resulted in lower IFN-γ secretion and reduced cell infiltration in brain more effectively as compared to the IV route. The effects of AT-MSCs on down-regulation of splenocyte proliferation, IL-17 secretion and alleviating the severity of clinical scores were similar in IP and IV routes [90]. Furthermore, it was reported that Adipose-Derived Mesenchymal Stem Cells (AD-MSCs) expressed murine interferon beta (MSCs-VP/IFN-β) in the animal model of MS (EAE) to induce Tregs and IL-10 and reduce IL-17 [91].

### Treatment of Patients With MS Using Mesenchymal Stem Cells (MSCs) (Table 3)

<table>
<thead>
<tr>
<th>Disease</th>
<th>Injection Way</th>
<th>Cell Source</th>
<th>Result</th>
<th>Cause</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Human adipose-derived mesenchymal stem cells engineered to secrete IL-10</td>
<td>Immunomodulation and neuroprotection</td>
<td>Inhibit APC function and limit CNS autoimmunity</td>
<td>[88]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Autologous adipose tissue mesenchymal stem cells (AT-MSCs) and their conditioned media (CM)</td>
<td>Reduction in proliferative responses and brain cell infiltration was more pronounced in mice injected with CM</td>
<td>Induced CD4 + CD25 + FOXP3 + regulatory T cells after in vitro co-culture with naïve T cells</td>
<td>[89]</td>
</tr>
<tr>
<td>EAE</td>
<td>Intravenous</td>
<td>Adipose-derived mesenchymal stem cells (AD-MSCs) expressing murine interferon beta (MSCs-VP/IFN-β)</td>
<td>Immunomodulation-reduces the migration of inflammatory cells into the CNS</td>
<td>Induction of Tregs and IL-10 and reduction of IL-17</td>
<td>[91]</td>
</tr>
</tbody>
</table>

Karussis et al. conducted studies on 15 patients with MS using intrathecal and intravenous injection of autologous stem cells and showed the MSCs immunomodulatory effect as well as the safety of the treatment of MS by injecting stem cells [96]. Another study done in the phase IIA of clinical studies, confirmed the safety and efficacy of autologous stem cells in 10 patients with MS [96]. In addition, treatment of 50 patients with MS in Iraq with intrathecal injection of purified peripheral blood cells through aphaeresis with Granulocyte Colony Stimulating Factor (G-CSF) yielded promising results [97].

Connick et al. in a clinical trial study injected autologous stem cells intravenously to 10 patients with MS who had disorders in optic nerves and monitored them for 10 months. They observed that the patients had achieved relative improvement in their optical nerves, and no side effects were observed during the use of stem cells [97]. In another study, autologous stem cells were intravenously injected to 8 patients with MS. The outcome was promising together with some positive effects without significant side effects [98].

Bonab et al. reported that intrathecal injection of stem cells that was expanded in vitro, have reduced the disease symptoms without the side effects [99]. Furthermore intrathecal treatment of MS patients with MSC resulted in increased expression of FoxP3 [100]. On the other hand, intrathecal injection of stem cells to 25 patients with MS had no significant effect on the cytokines such as IFN-γ, TGF-B, IL-4, IL-10, and IL-6, as well as FoxP3 in peripheral blood cells [101]. In addition, in a study over four years of treatment, a patient with progressive MS with both allogeneic MSCs derived from human umbilical cord and cells derived from autologous bone marrow, no significant side effects were observed [102]. Several clinical trials are underway around the world that exam-
Table 3. Treatment of patients with MS using mesenchymal stem cells (MSCs)

<table>
<thead>
<tr>
<th>Disease Type</th>
<th>Number of Patients</th>
<th>Cells Injected into the Patient</th>
<th>Injection Route</th>
<th>Clinical Results</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Progressive MS</td>
<td>10</td>
<td>MSCs expanded in vitro</td>
<td>Intrathecal</td>
<td>Improvement of symptoms</td>
<td>[93]</td>
</tr>
<tr>
<td>MS</td>
<td>4</td>
<td>MSCs derived from endometrial cells</td>
<td>Intravenous and intrathecal injection</td>
<td>Improved symptoms, no side effects</td>
<td>[94]</td>
</tr>
<tr>
<td>MS</td>
<td>32</td>
<td>MSCs derived from umbilical cord</td>
<td>Intrathecal</td>
<td>Suppression of neuro-inflammation</td>
<td>[58]</td>
</tr>
<tr>
<td>MS</td>
<td>10</td>
<td>MSCs derived from autologous bone marrow</td>
<td>Intrathecal injection</td>
<td>Improved symptoms</td>
<td>[95]</td>
</tr>
<tr>
<td>MS</td>
<td>15</td>
<td>MSCs derived from autologous bone marrow</td>
<td>Intravenous injection-intrathecal injection</td>
<td>Improved symptoms, no side effects</td>
<td>[96]</td>
</tr>
<tr>
<td>Secondary progressive MS</td>
<td>10</td>
<td>MSC derived from autologous bone marrow</td>
<td>Intravenous injection</td>
<td>Improved symptoms, no side effects</td>
<td>[97]</td>
</tr>
<tr>
<td>MS in Iraq</td>
<td>50</td>
<td>The peripheral blood cells purified through aphaeresis with G-CSF</td>
<td>Intrathecal injection</td>
<td>Improvement of clinical symptoms, no adverse-clinical effects</td>
<td>[98]</td>
</tr>
<tr>
<td>MS with optic nerve disorders</td>
<td>10</td>
<td>Autologous MSC</td>
<td>Intravenous</td>
<td>Relative improvement in optic nerve, no clinical side effects</td>
<td>[99]</td>
</tr>
<tr>
<td>MS</td>
<td>8</td>
<td>Autologous MSC</td>
<td>Intravenous</td>
<td>Improved clinical symptoms, no clinical side effects</td>
<td>[100]</td>
</tr>
<tr>
<td>Progressive MS</td>
<td>25</td>
<td>Autologous MSC</td>
<td>Intrathecal</td>
<td>MSC therapy can improve/stabilize the course of the disease in progressive MS in the first year after injection with no serious adverse effects.</td>
<td>[101]</td>
</tr>
<tr>
<td>MS</td>
<td>7</td>
<td>Autologous MSC</td>
<td>Intrathecal</td>
<td>Increased expression of FoxP3, and thus increasing Treg</td>
<td>[102]</td>
</tr>
<tr>
<td>MS</td>
<td>25</td>
<td>Autologous MSC</td>
<td>Intrathecal</td>
<td>It has no effect on the cytokines such as IFN-γ, TGF-β, IL-4, IL-10, IL-6 as well as Foxp3 in the peripheral blood</td>
<td>[102]</td>
</tr>
<tr>
<td>Progressive MS</td>
<td>1</td>
<td>1) Allogeneic MSCs derived from human umbilical cord(2) MSCs derived from autologous bone marrow</td>
<td>Intrathecal injection</td>
<td>The use of both types of MSCs improves clinical symptoms with no clinical side effects</td>
<td>[103]</td>
</tr>
<tr>
<td>3 SPMS 3 RRMS 2 PPMS</td>
<td>8</td>
<td>Autologous MSC</td>
<td>Intravenously</td>
<td>MRI lesion relapse and improved clinical symptoms</td>
<td>[98]</td>
</tr>
</tbody>
</table>
ine the effectiveness and side effects of treatment of MS patients using stem cells [33]. However, despite some studies in phase I/II, no significant positive result indicating complete remission of the disease using stem cells is available and more research is needed.

The use of stem cells and MSCs for the treatment of autoimmune diseases has been much considered in the past decade. Now MSCs are widely used in clinical trials, and initial promising results are obtained with regard to the control and prevention of MS. Several studies have shown that these cells are in close contact with various immune cells. Therefore, they can be considered a valuable tool for the treatment of various inflammatory diseases such as MS. However, clarification of the exact mechanisms of these cells in the treatment of patients with MS is required. The identification of these mechanisms in future could be useful in selecting the appropriate dose and injection time. It is also valuable in the selection of cells derived from different tissues and sources for the treatment of MS.

Ethical Considerations

Compliance with ethical guidelines

There is no ethical principal to be considered doing this research.

Funding

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Conflict of interest

The authors declare no conflict of interest.

References


[49] Steinman L. A brief history of T(H)17, the first major revision in the T(H)1/T(H)2 hypothesis of T-cell-mediated tissue damage. Nature Medicine. 2007; 13(2):139-45. [DOI:10.1038/nm1551]


