Therapeutic Aspects of Mesenchymal Stem Cell-Based Cell Therapy with a Focus on Human Amniotic Epithelial Cells in Multiple Sclerosis: A Mechanistic Review

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Multiple sclerosis (MS) is an inflammatory disease of central nervous system (CNS). The immune system plays an important role in its pathogenesis. Current treatments are unable to cure patients and prevent the progression of MS lesions. Stem cell-based cell therapy has opened a new window for MS treatment. Stem cells regulate immune responses and improve axonal remyelination. Stem cells can be obtained from different origins such as embryonic, neural, bone marrow, and adipose tissues. But yet there is a challenge for the selection of the best cell source for stem cell therapy. Mesenchymal stem cells (MSCs) are a type of stem cell obtained from different origins and have significant immunomodulatory effects on the immune system. The increasing evidence have suggested that umbilical cord and adipose tissue can be a suitable source for isolation of MSCs. Moreover, human amniotic epithelial cells (hAECs) as novel stem cell origins by having immunoregulatory effects, regenerative effects, and less capacity of antigenicity can be a candidate for MS treatment. This review discussed the mechanistic effects of MSCs with a focus on human amniotic epithelial cells, which can be used to treat and improvement of outcome in MS disease.

Keywords: Multiple sclerosis, Human amniotic epithelial cell, Mesenchymal stem cell, Immunomodulatory effects, Regenerative impacts

Introduction

Neurodegenerative disorders are defined as chronic and progressive processes which result in the deposition of abnormal forms of specific proteins in the nervous system and destruction of neurons in motor, sensory, or cognitive systems (1). These disorders mainly occur in females and subjects with age range of 20 to 30 years (2). Besides, it is estimated that more than 2.5 million subjects have multiple sclerosis (MS) around the world which need to effective treatments to recover disability and improve the central nervous system (CNS) functions (2). The major obstacle of curative treatment of MS is an unsuccessful regeneration of neurons (3). The most common treatments for MS exert therapeutic impacts through inhibiting immune reactions (2, 4). However, the most of these therapeutic approaches are unable to stop the degeneration of nerve tissue in aggressive MS (5). Among these methods, stem cell based-cell therapies have a promising outlook for
the remedy of neurological disorders by differentiation into astrocytes and oligodendrocytes effectively in vivo and in vitro as well as the secretions of neurotrophic agents which have immunoregulatory effects and regenerative roles for remyelination and thereby can decrease neural damages (6). There are some challenges for using stem cells as a curative approach in clinical trials because they can quickly proliferate and subsequently have a capacity for tumorigenicity (7). However, increasing evidence have shown that stem cell therapy can have positive effects on animal models with neurological diseases (7, 8). Clinical applications of adult stem cells, particularly mesenchymal stem cells (MSCs) and human amniotic epithelial cells (hAEC), have been recommended for the treatment of neurological disorders such as MS (9-11). These cells can easily be isolated, expanded, and manipulated ex vivo (12). They can effortlessly differentiate into mesodermal lineages (13). Other advantages of therapeutic applications of MSCs and hAECs include their capability to transport from blood to damaged sites, their relative safety, and low immunogenicity in comparison with other stem cell sources (13, 14). Hereby, the present review aimed to describe and discuss evidence regarding MSC-based therapies with concentrating on hAECs and their mechanisms for treating MS.

**MS and Its Pathogenesis**

Multiple sclerosis (MS) is a chronic and progressive neuro-inflammatory disease of the CNS, leading to axonal lack and myelin disruption. Among this, immune cells play a critical role in the pathogenesis of MS (15, 16). The immune system modulates neural evolution via regulating oligodendrogenesis, neurogenesis, and synaptic organization. In addition to environmental factors, genetic factors such as immune system regulating genes including major histocompatibility complex (MHC), T-cell receptor (TCR), immunoglobulin (Ig), and cytokines can increase the risk of MS (2, 17). Formations of sclerotic plaques and lesions in the CNS and the cerebrospinal cord are the features of MS (15, 16). However, T cells alone cannot form lesions and plaques and other immune cells play important roles in the pathogenesis of MS. Th1 and Th17 produce interleukine-1 (IL-1), IL-17, tumor necrosis factor-alpha (TNF-α), interferon-gamma (IFN-γ), and other pro-inflammatory cytokines (18). The consequence of macrophage activated by Th1 cytokines is the destruction of the myelin and exposure of more CNS antigens. In vivo studies have shown that Th1 cytokines such as IFN-γ, lymphotoxin (LT)-α and IL-2 have the important roles in the pathogenesis of MS. Furthermore, TNF-α is associated with progression of the disease. CD8+ T cells are also observed in MS lesions, especially around the blood vessel. Clonal expansion of CDS+ T cell is more than CDS+ T cell and CD8+ T cell number is largely correlated to axons injury (1). Although it is shown that autoreactive T cells are the main effector cells responsible for the pathogenesis of disease, previous studies have reported that autoreactive B cells also participate in neural damages and their myelin sheaths and axons, through autoantibody productions, antigen presentations, and cytokine secretions (19). Clinical and experimental studies have shown that autoantibodies can be major components of the immune system that can be detected in MS plaques. In this notion, there are some studies showing the associations between immunoglobulin G (IgG) and MS signs. Moreover, it is indicated that IgG, especially IgG against myelin basic proteins (MBP) and proteolipid proteins (PLP), can be considered as the hallmarks of MS. However, these antibodies have the unknown roles in its pathogenesis (Fig. 1) (20).

**Mesenchymal Stem Cells (MSCs)**

MSC can be isolated from all connective tissues including bone marrow, umbilical cord, adipose tissue, dental tissues, brain, and fetal lung (21-23). Several studies have shown that MSCs can differentiate into monocytes and neurons in vitro and in vivo (24). MSCs can migrate to areas of injured tissue via expressions of receptors for chemokines such as CCR1, CXCR4, CXCR5, and CXCR6, and some growth factors (6).

MSCs may support axon and increase neural stability through anti-oxidant and anti-apoptotic impacts and/or trophic factor secretions (25). MSCs can recruit oligodendrocyte precursors to the CNS and induce differentiation of these cells into neuronal cells (26, 27). In addition, MSC inhibits neuron apoptosis, improves neural cell differentiation, and promotes angiogenesis and CNS repair in MS patients (20).
MSC regulates immune responses via suppressing the functions of B, T cells, and other immune cells (28). Studies have shown that intravascular injection of MSC induces T cell tolerance against myelin glycoproteins and improves CNS tissue repair (29). Intravenous injection of syngeneic MSC to animals with experimental autoimmune encephalomyelitis (EAE), as a model of MS, induces tolerance in MOG-specific T cells, which results in the reduction in immune cell infiltrations to the CNS and myelin destruction and improves the clinical course (30, 31). MSCs inhibit immune responses through the secretions of anti-inflammatory cytokines such as TGF-β, indoleamine-pyrrole 2, 3-dioxygenase (IDO), and prostaglandinE-2 (PGE-2) (32).

In vivo studies have indicated that MSCs are able to inhibit the activation and proliferation of Th1 cells and CD8+ T cell-mediated cytotoxicity functions through cell-cell interactions, which consequently lead to an impairment in the activation and proliferation of B cells and production of antibody (33). Moreover, MSC can enhance the activation of suppressor of cytokine signaling 3 (SOCS3) and control Th17 differentiation via IFN-γ pathway (34). IDO production is another mechanism used by MSC for inhibiting immune responses (34). IDO depletes tryptophan from the environment of lymphocytes which has crucial role in lymphocyte activations (35). MSC induces peripheral dendritic cells (DCs) to secret IL-10 and thereby promotes the development of regulatory T cells (Treg) (35).

Recent studies suggest that MSCs improve peripheral tolerance through preventing the differentiation and function of DCs leading to reduction in antigen presentation and subsequently inhibition of clonal expansion of autoreactive T cells (36). MSCs secrete hepatocyte growth factor (HGF) which increases tolerogenic DCs (37). It is reported that injection of MSC with HGF to EAE mice reduced CNS inflammation and immune cell infiltrations (37). Thus, it is likely that HGF derived from MSC is a good candidate for the treatment of MS. In several studies, genetically modified MSCs have been used to over express anti-inflammatory cytokines such as IL-10, and IL-4 (38). These studies have shown that MSCs suppress immune responses, reduce BBB injury, and improve remyelination of neurons in EAE mice (Fig. 2) (39).

**Human Bone Marrow-Derived MSCs (hBM-MSCs)**

The hBM-MSC, as a nonhematopoietic BM cell, includes approximately 0.0001 ~ 0.01% of all BM-nucleated cells (39). MSCs obtained from bone marrow have multiple properties including: I) differentiation into mesodermal lineage cells, II) stromal marker expressions, III) colony formations that contribute to haemopoiesis (40). Regarding therapeutic aspects of hBM-MSCs in neurological disorders, it is stated that these cells can promote functional recovery in relapsing-remitting and chronic types of MS mouse model probably through reducing demyelination regions and inflammatory infiltrates, enhancing brain-derived neurotrophic factor (BDNF) expression, and inducing oligodendrogenesis (41). Some studies have shown that BM-MSCs possess immunomodulatory effects in EAE mice (36). They can exert immunoregulatory impacts through releasing various bioactive agents which suppress the proliferation of B and T cells and the maturation of antigen-presenting cells (APCs) (42). These cells play neuroprotective roles via inducing local progenitor
cells and suppressing scar creation, gliosis, and neuron apoptosis (43). In addition to protective impacts, they are able to differentiate into the neurons and enhance the replacement of the cells (43). Nevertheless, the isolation of BM-MSCs is painful, invasive, and low efficiency (44). In EAE, BM-MSCs are efficient only when they are administered before disease initiation, and have few curative effects in stabilized stage of disease (13). Thus, other sources of MSCs can be useful for MS treatment.

**Human Umbilical Cord (hUC)-MSCs**

Among several sources of MSCs, the utilization of human umbilical cord (hUC)-MSC has significant benefits such as less ethical issues, less invasive methods for its isolation, great proliferation potential, little immunogenicity capacity, and differentiation capacity into various lineages (45). Multiple studies have suggested hUC-MSC administration in autoimmune conditions like encephalomyelitis, type 1 diabetes, and rheumatoid arthritis due to its regulatory impacts on immune reactions (46-48). HUC-MSCs have significant effects on tissue recovery and immunoregulation (49). In comparison with BM-MSC, hUC-MSCs significantly have the decreased expression of HLA-I, increased expansion capacity, and more rapid growth in vitro (50). HUC-MSCs can also increase Treg number in vitro and in vivo (51). Liu et al. (48) showed that hUC-MSCs can improve behavioral activities and diminish the histopathological impairments of encephalomyelitis mouse models. They also declared that this mesenchymal source can dramatically elevate IL-4 and IL-10 and reduce IL-1 and IL-6 levels. In addition, hUC-MSCs can release various nerve growth factors, for example, BDNF, glial cell-derived neurotrophic factor (GDNF). Furthermore, hUC-MSCs can differentiate into oligodendrocyte precursor cells and enhance axonal growth (52).

In line with the immunomodulatory effects of these cells, it is demonstrated that hUC-MSCs cultured with peripheral blood mononuclear cells (PBMCs) from healthy individuals can promote Treg frequency and anti-inflammatory cytokine secretions (Fig. 3) (51, 53).

**Human Adipose-Derived MSCs (AD-MSCs)**

AD-MSC is a subset of MSCs isolated by collagenase digests of adipose tissue. Adipose tissue is available, abundant, easily accessible for use, and contain high levels of MSCs (about 100–1000 MSCs per gram of fat). Therefore, this tissue is considered as a valuable source of the cell for cellular therapy. AD-MSCs have the neurogenic, cardiogenic, myogenic, chondrogenic, osteogenic, and adipogenic capacity in vitro, which can be obtained and used to stem cell therapy by less invasive methods (6, 54). AD-MSCs can express α4 integrin, an adhesive molecule, and thereby have the ability to migrate in different organs (55). Previous studies have demonstrated that autologous and allogeneic AD-MSCs can be useful in the treatment of various diseases such as MS, diabetes, Parkinson’s disease, autoimmune encephalomyelitis (56-59). In line with this notion, Li et al. revealed that intravenously injection of AD-MSC to EAE mice contributed to the reduction in infiltrations of inflammatory cells in the CNS and significantly inhibited demyelination and axonal loss (8, 59). These cells release various growth factors including anti-apoptotic, angiogenic and neurotrophic mediators which have significant roles in many cellular mechanisms such as cell differentiation, proliferation, and maturation (60). In contrast with BM-MSCs, AD-MSCs express integrin α4β1. Thus, it is thought that AD-MSC has more capabilities for stem cell-based cell therapy, due perhaps to its
potential to pass the BBB and exert its effects such as anti-inflammatory, immunoregulatory, and neurodegenerative (Fig. 4) (5). Recently, several studies have been performed in this field (61, 62). Based on their results, the researchers are hopeful to find a standard method for treatment of MS by these cells (62). However, it seems that many concerns regarding the use of MSC such as tumorigenesis and immune-rejection after clinical application must be addressed in future works.

**Human Amniotic Epithelial Cells (hAECs)**

hAEC is a potential source of stem cell which can be easily isolated from the amniotic membrane (AM), the inner layer of the fetal membranes (18, 63-65). These cells can express some markers of pluripotent stem cells, such as Rex-1, FGF-T, SSEA-4, Sox-2, Nanog, and Oct4, which some of them play critical roles in self-renewal and pluripotency properties in induced pluripotent stem (iPS) cells and embryonic stem cells (ESCs) (66). hAEC displays multilineage differentiation potential and has the ability to differentiation into different cells such as the neural cells, pancreatic cells, hepatocytes, adipocytes, cardiomyocytes, and myocytes, which originate from the endoderm, mesoderm, and ectoderm (12). Extensive data from the literature have indicated that hAECs have immunoregulatory effects on adaptive and innate immune systems (14, 63, 67). They have inhibitory impacts on the migrations of neutrophil and macrophage, proliferation of B cell, activations of natural killer (NK) and CD4+ T cells, and se-
cretions of pro-inflammatory cytokines of CD4+ T cells (12, 18, 68, 69). The immunoregulatory features of hAECs are mainly mediated by secretions of immunoinhibitory agents, including interleukin-4 (IL-4), PG-E2, and transforming growth factor-beta (TGF-β) (70-73). Moreover, immunomodulatory impacts of hAECs may be mediated by the increase of Tregs and Th2 cells, IL-2, IL-5, and IL-10 levels, inhibition of pathogenic T cell reactions, and preservation of the peripheral naive CD4+ T cell source (74, 75). Thus, hAECs may be acted as a useful cell source for MS treatment (70, 74). To support this notion, these cells have a significant impact on shift in the immune system away from Th1 immunity towards Th2 responses (74). It is reported that hAECs-treated EAE mice experienced significant reductions in demyelination and infiltration of CD3+ T cell and F4/80+ monocyte/macrophage into the CNS (74). Some reports have indicated that hAECs inhibit the differentiation of Th17 cells through reducing the expressions of TGF-β and IL-6, which have indispensable roles in Th17 cell differentiation (67). Other studies have indicated that hAECs are able to produce alpha-fetoprotein (AFP), which is involved in the reduction of lymphocyte function and neuroinflammation in MS animal model (76, 77). Moreover, they can reside in inflammation locations, such as the brain, and subsequently decrease grey and white matter damages (78). hAEC can release neurotrophic agents such as nerve growth factor (NGF), neurotrophin-3 (NT-3) and brain-derived neurotrophic factor (BDNF) (Fig. 5) (79, 80). These properties along with low antigenicity provide further confirmations to support therapeutic features of hAECs in managing and controlling inflammatory neurological disorders such as MS (70). hAECs have a restricted proliferative potential due to lacking telomerase (66, 70, 74, 81), which may reduce potential tumorigenicity of stem cell-based therapies. Nevertheless, it is thought that additional studies and more information are required to clarify the possible ability of hAECs in treating diseases with immune pathophysiology.

Comparison of hAECs with MSCs from Different Sources

There are some similarities and differences between hAECs and MSCs derived from different sources including morphologic and tumorigenic properties, angiogenesis capacities, immunomodulatory features, and ethical issues related to their isolations and applications (44, 82). In terms of morphology, hAECs indicate a cobblestone-like morphology, and cultured hAMSCs have a spindle fibroblast-like morphology (83). The morphologic feature of BM-MSCs can be ranged from fibroblast-like spindle-shape to large flat cells in culture (84). MSCs from other sources, such as hUC-MSCs and AD-MSCs, show spindle shapes in the culture (85, 86). The collection of amniotic cells can easily be performed by prenatal testing. Moreover, the amniotic membrane can be collected using standard isolation techniques after cesarean section which are noninvasive without detrimental effects on human embryos and ethical issues (82). However, some ethical problems are proposed to clinical applications of MSCs and the isolation of some MSC sources (84, 87). As mentioned previous, the isolation of hBM-MSCs is invasive, painful,
and low efficiency (44). HAECS don’t have tumorigenic capacity, and there is no document about the tumorigenicity of amnion membrane or membrane-originated cells when they are used to clinical applications (14). However, it is reported that MSCs may elevate tumor growth in some cancer mouse models (88). HAECS, BM-MSCs, and AD-MSCs can promote angiogenesis through releasing angiogenic factors such as VEGF, HGF, EGF and some mechanisms associated to protease and cytokine productions (83, 89). Based on evidence, hAECs have better immunoregulatory effects but lesser osteogenic impacts than MSCs derived from BM and human amniotic fluid (hAF) (90). These cells, similar to MSCs, express some MSC markers such as CD44, CD90, and CD105. However, hAECs possess higher levels of SSEA4 and SSEA3 expressions than hBM-MSCs and hAFMSC, indicating more multipotent potential of hAECs (14, 91). Furthermore, hAECs and hAFMSC express higher PD-L1 and PD-L2 levels than hBM-MSCs, thereby these cells may have more effective effects in peripheral tolerance in different cells of the immune system (90, 92).

Mesenchymal Stem Cell-Based Cell Therapy and Clinical Trials

Recently, several clinical trials were preformed using MSCs as a treatment for MS. In a phase II clinical trial, nine relapsing-remitting multiple sclerosis (RRMS) patients were treated with MSCs for six months resulting in the reduction of lesions in magnetic resonance imaging (MRI) (43). In 2010, Yamout et al. (43) injected autologous BM-MSCs to nine secondary progressive multiple sclerosis (SPMS) and one RRMS patients. After three months to one year, they observed improved clinical features in MS patients. In this phase IIa clinical trial, 10 SPMS patients received intravascular MSCs for six months and the researchers observed remyelination and neuroprotection effect of MSCs (43). Moreover, in another study conducted by Mohyeddin Bonab et al. (93) in 2007, MSC was intrathecally injected to 10 MS patients and investigated disease procession. The disease development was gradually decreased in half of patients upon MSC application. In a study on 22 patients with primary progressive multiple sclerosis (PPMS), it was reported that intravenously and intrathecally injections of BM-MSCs had helpful effects in MS treatment (32). In a triple-blind and placebo-controlled investigation on 30 patients with SPMS, it was shown that AD-MSCs infusion is a feasible and safe procedure in SPMS patients (94). In another study by Staff et al. (95) the safety of intrathecal admin-istration of AD-MSCs in amyotrophic lateral sclerosis (ALS) cases was reported. Li et al. (96) assessed the therapeutic influences of another MSC source. They demonstrated that hUC-MSCs transplantation decreased MS symptoms and relapse occurrence compared to control subjects. Furthermore, they revealed a shift from Th1 to Th2 immunity in the hUC-MSCs-administrated group. In addition, in another study, Riordan et al. (97) indicated the safety of hUC-MSCs transplantation and useful impacts of these cells on life quality and brain lesion in MS cases.

Conclusions

Stem cell-based cell therapy is getting to be a treatment for MS and other neurological diseases. However, there is an inconsistency in the results of previous studies. This discrepancy proposes that more studies are needed to demonstrate the exact role of stem cells in MS treatment. Among several types of stem cells, MSCs are more likely to find a place in future treatments for MS because these cells have different mechanisms to regulate immune responses and repair CNS damages. Furthermore, MSCs have a supportive role on axon and the neural stability and have anti-oxidant, anti-apoptotic and trophic factors secretion properties. A growing body of evidence recommends that umbilical cord and adipose tissue can be more effective for stem cell therapy on the basis of MSCs. Moreover, hAECs are considered as a novel stem cell source that can have a potential for differentiation into derivatives of three germinal layers such as neural cells and possess immunoregulatory effects. Therefore, they can also have the ability to MS treatment. However, more experimental investigations should be performed to explain their effectiveness and mechanisms in the treatment of this disease.

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Potential Conflict of Interest

The authors have no conflicting financial interest.

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