A review on stem cell therapy for multiple sclerosis: special focus on human embryonic stem cells

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Abstract: Multiple sclerosis (MS), a complex disorder of the central nervous system (CNS), is characterized with axonal loss underlying long-term progressive disability. Currently available therapies for its management are able to slow down the progression but fail to treat it completely. Moreover, these therapies are associated with major CNS and cardiovascular adverse events, and prolonged use of these treatments may cause life-threatening diseases. Recent research has shown that cellular therapies hold a potential for CNS repair and may be able to provide protection from inflammatory damage caused after injury. Human embryonic stem cell (hESC) transplantation is one of the promising cell therapies; hESCs play an important role in remyelination and help in preventing demyelination of the axons. In this study, an overview of the current knowledge about the unique properties of hESC and their comparison with other cell therapies has been presented for the treatment of patients with MS.

Keywords: multiple sclerosis, stem cells, human embryonic stem cells, remyelination, axonal loss, neurological disorder

Introduction
Promising advances have been made in the recent years in stem cell transplantation to treat and cure damaged tissues, injuries, and various diseases. This is especially true with the various autoimmune disorders, specifically for the ones which are associated with nervous system, where damage often appears from the degeneration of cells.1

Treatment using cell therapy involves a form of immune suppression but unlike standard immune-based drugs, it is designed to reset rather than suppress the immune system. Scientists define three ways to use stem cell therapy: the first involves the use of stem cells to replace damaged myelin-forming oligodendrocytes within the central nervous system (CNS);2,3 the second aims to replace the individual's malfunctioning immune system, making use of hematopoietic stem cells (HSCs);4,5 and the third seeks to utilize the endogenous stem cell populations by mobilization with or without in vitro expansion, exploiting their reparative and neuroprotective properties.6,7

Studies have been carried out for various type of stem cells: autologous hematopoietic stem cells (AHSCs), mesenchymal stem cells (MSCs), neural stem cells (NSCs), induced pluripotent stem cells (iPSCs), and human embryonic stem cells (hESCs).8

hESC transplantation is a controversial yet a powerful strategy for the treatment of patients with autoimmune diseases. hESC lines can differentiate into neural precursor cells and neurons, astrocytes, and oligodendrocytes that show their potential in treating several incurable neurological disorders like spinal cord injury,9,10 cerebral palsy,11
multiple sclerosis (MS), Parkinson’s disease, and many more. The remarkable properties of hESC therapy in treating various terminal conditions both in vitro and in vivo could be used as the first-line therapy in the future.

Of the various neurological diseases, MS is one of the most complicated in which the underlying pathological mechanism remains unclear, and the incompetence of a large number of promising treatments for the disease makes it an ideal target for use in regenerative medicine. Pluripotency, self-renewal, and ability to regenerate and differentiate into neuronal cells are the promising factors to use hESCs in treating MS. These properties make them an ideal resource of unlimited supply of neural derivatives. This review provides an account on the role of hESC lines in treating MS and their comparison with other cellular therapies. Subsequently, this will discuss the development of hESC line at the Department of Stem Cell Therapy, Nutech Mediworld, India that has been able to treat various terminal diseases.

**MS and its pathogenesis**

MS is a chronic inflammatory demyelinating disease of the CNS which involves the loss of myelin-forming oligodendrocytes that can be followed by a spontaneous and an efficient regenerative process called remyelination. It affects the people of almost all the ages in many parts of the world, mostly young people, especially more prevalent in women, and among those in northern latitudes. Relapsing–remitting multiple sclerosis (RRMS) is the most common form of MS. The disease course is biphasic; initially, there are alternating episodes of acute neurological deficits or worsening episodes of relapses followed by a complete or partial recovery (i.e., remission). Within a span of 15–20 years, most (70%) of the patients with RRMS develop secondary progressive MS which is characterized by a progressive neurological decline independent of relapses (inflammation). MS has a genetic susceptibility, but it is not directly inherited. The disease has an unknown etiology with no cure; only symptomatic treatment is available.

**Current treatment approach for MS and their limitations**

Since 2010, various oral therapies were approved which offer added convenience, but all these therapies provided only symptomatic relief. The currently available treatments mainly target the underlying immunologic etiology of MS. Most of the therapies are composed of medications that are either immunomodulatory or immunosuppressive and are aimed at reducing the frequency and intensity of the relapses. These treatments are undoubtedly quite efficient in preventing the frequency of relapses but are not able to make up the axon damage loss, further progression of disease, and irreversible disability. Current disease modifying therapies (DMTs) including interferons, immunosuppressants, corticosteroids, and monoclonal antibodies mainly target the cells and mediators of the adaptive immune system which are the triggering factors to cause the immune response to elicit the disease; however, this only helps to slow down the progression. Also, these treatments lack in providing the neuroprotective benefits to the patients, resulting in further damage/loss.

The treatment for MS is expensive. With the progression of the disease and increasing disability level, there has been a further increase in the direct and indirect costs involved. This increment in cost is generally related to relapses and productivity costs rather than the direct cost involved in using DMTs. Direct and indirect health care costs range from US$8,528 to $54,244 per patient per year in the US. MS ranks second only to congestive heart failure in terms of price when compared with other chronic conditions.

First-generation DMTs, originally costing US$8,000 to $11,000, now cost about US$60,000 per year. Newer DMTs commonly entered the market with a cost 25%–60% higher than existing DMTs. Cellular therapies in MS

Generally for most of the stem cell therapies, the first approach to treat MS is to start with high dose of immunosuppressants followed by stem cell transplantation. The rationale behind this approach is that immunosuppression helps to eliminate autoreactive T and B cells, therefore allowing installation of new and tolerant immune system.

Despite the standard therapies that are available for MS, majority of the patients eventually enter a secondary progressive phase for which no therapy has demonstrated efficacy. Reversible relapses due to the potent compensatory mechanisms in the brain, which cause extensive axonal loss and demyelination, are the bottlenecks of this disease. Remyelination is one of the strategies to recover the damage and prevent the further relapses. It is a regenerative process by which demyelinated axons are reinvested with new myelin sheaths. Spontaneous and robust remyelination occurs at the early stages of MS.

Stem cell therapy is an effective therapy in regeneration medicine and a new approach to overcome the limitations of currently available therapies. Stem cell treatment is now available all over the world to treat MS. However,
the cost of treatment varies in every country as per its own regulations and policies. In US, the average cost per treatment is US$7,000 to $10,000. Costs for treatments outside of the US are usually far higher than in the US, ranging from US$20,000 up to $100,000.\textsuperscript{37,38} A cost-effectiveness survey of AHSC transplantation for MS treatment in the UK conducted by Tappenden et al reported that the cost-effectiveness of AHSC transplantation is expected to be around £2,800 per additional quality-adjusted life year gained.\textsuperscript{39} Table 1 shows the various stem cell studies in MS.

Different cellular therapies and their mechanism of actions in treating neurological disorders

Studies have been carried out for various types of stem cells: HSCs, adult stem cells that are found in bone marrow and blood; MSCs, adult stem cells found in several places in the body, including the bone marrow, skin, and fat tissue; NSCs, specialized stem cells responsible for repairing nerve-insulating myelin in the brain. These can be derived from other types of stem cells such as MSCs, iPSCs (engineered from adult cells to produce many types of cells), and hESCs (stem cells derived from donated embryos that can naturally produce every type of cell in the body).\textsuperscript{8}

Autologous hematopoietic stem cell treatment

High-dose immunosuppression followed by autologous hematopoietic stem cell treatment has been investigated in the past years as treatment for MS patients who are refractory to first-line therapy. The treatment is based on the rationale “high-dose immunosuppression is able to eliminate autoreactive T and B cells, therefore allowing installation of a new and tolerant immune system.”\textsuperscript{40,41} It uses harvested stem cells to reset patients’ immune systems and reverse some of the symptoms of MS. During the treatment, bone marrow stem cells are collected from patient’s blood and are frozen. These “HSCs” are at an early stage of development and lack the flaws that trigger MS. The faulty immune system is then destroyed using chemotherapy. The thawed-out stem cells are reinfused into the blood to reboot the immune system.\textsuperscript{42}

MSCs

Immunomodulatory effects of MSCs include suppression of T cell proliferation, induction of regulatory T cells, influencing dendritic cell maturation and function, suppression of B-cell proliferation and terminal differentiation, and inhibition of natural killer cell function. Yamout et al reported that intrathecal administration of ex vivo-expanded autologous bone marrow-derived MSCs showed therapeutic benefit in 10 patients with advanced MS.\textsuperscript{43}

NSCs

Remyelination is the regenerative process by which demyelinated axons are reinvested with new myelin sheaths. NSCs in the adult brain differentiate into oligodendrocytes in a context-specific manner and get involved in the remyelination of MS patients. NSCs may exert their beneficial effects not only through oligodendrocyte replacement but also by providing trophic support and immunomodulation, a phenomenon now known as “therapeutic plasticity.” For the NSC therapy to be successful in MS, the cells need to be plastic enough to accommodate and survive in the nonpermissive inflammatory environment.\textsuperscript{24} In the adult CNS, tissue-specific germinal niches, such as the subventricular zone of the lateral ventricles and the subgranular zone of the dentate gyrus of the hippocampus, contain multipotent NSCs which have the capacity of self-renewal and differentiation into functional neurons and glial cells.\textsuperscript{54,45}

Induced pluripotent stem cells

These cells can be differentiated into oligodendrocyte precursor cells (OPCs) which are essential to prevent axonal and subsequent retrograde neuronal degeneration in demyelinating lesions in MS. Studies conducted by Thiruvalluvan et al in non-human primate model showed that human-induced pluripotent stem cell (hiPSC)-derived OPCs migrate toward the MS-like lesions in the corpus callosum where they myelinate the denuded axons; this new approach indicates the potential use of hiPSC-derived OPCs for progressive forms of MS.\textsuperscript{46}The therapeutic potential of iPSCs has been extensively assessed by transplantation of iPSC-derived specific cells into different animal disease models.\textsuperscript{47,48} Safety and efficacy are the most important concerns for clinical development of iPSCs, as certain cells derived from iPSCs have tendency to elicit antigen-specific immune rejection responses after transplantation into the autologous recipients.\textsuperscript{49,50}

hESCs

The remarkable properties of hESCs have shown their clinical usefulness over the other stem cells studied. hESC-based therapies can give rise to specific specialty cells such as dermatomes from undifferentiated ESCs or incompletely differentiated neural cell. hESCs provide immunosuppressive neuroprotective mechanism.\textsuperscript{51} Research on hESCs is obligated
Table 1 Studies using stem cells to treat MS

<table>
<thead>
<tr>
<th>Type of stem cells</th>
<th>Authors, year</th>
<th>Preclinical/clinical</th>
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<tr>
<td>Neural stem cells (NSCs) – glial cells</td>
<td>Haffernan et al, 2012⁴⁹</td>
<td>Clinical human model</td>
<td>The study involved the designing of CPP sequence that selectively transduced human glial cells (excluding nonglial cell types), and conducted preliminary screens of purified, recombinant CPPs with immature and matured human oligodendrocytes and astrocytes and two nonglial cell types to treat MS.</td>
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<td>Embryonic stem cells derived NSCs</td>
<td>Payne et al, 2012⁵¹</td>
<td>Preclinical mouse model</td>
<td>The study evaluated the therapeutic potential of NSCs derived from ES cells by two different neural differentiation protocols that utilized adherent culture conditions and compared their effect with primary NSCs derived from the subventricular zone (SVZ).</td>
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<tr>
<td>Hematopoietic stem cells (HSCs)</td>
<td>Fassas and Kazis, 2003⁴¹</td>
<td>Clinical Phase I and II studies in human</td>
<td>The study was based on concept of immune ablation by high-dose therapy and reconstitution of normal immunity from transplant-derived lymphocyte progenitors. It gave way to the concept of “resetting” the immune system and of bringing the disease to a lower level of activity.</td>
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<tr>
<td>iPSC-derived neural progenitor cells (NPCs)</td>
<td>Nicaise et al, 2007⁵²</td>
<td>iPSC cell lines were obtained from humans and NPCs were obtained from mice models</td>
<td>The study involved comparison of treating PPMS cases with NPCs and iPS-derived NPCs. The results showed that NPCs from PPMS cases provided no neuroprotection against active CNS demyelination compared to NPCs from control iPS lines which were capable of completely preventing injury.</td>
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<tr>
<td>Neural precursor stem cells (NPSCs)</td>
<td>Donega et al, 2014⁵³</td>
<td>Preclinical mouse model</td>
<td>The study involved injecting NPSCs in mice model of experimental autoimmune encephalomyelitis (EAE), via iv and icv route of administration, with chronic CNS inflammatory demyelination, and envisage the systemic stem cell delivery as a valuable technique for the selective targeting of the inflamed brain in regenerative neurology.</td>
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<td>Autologous hematopoietic stem cells (AHSCs)</td>
<td>Chen et al, 2012¹</td>
<td>Clinical human studies</td>
<td>This was a retrospective evaluation for the safety and long-term clinical outcome of AHSC therapy in MS patients in China. Twenty-five patients with various types of MS were treated with AHSC therapy. Peripheral blood stem cells were derived by leukapheresis after mobilized with granulocyte colony-stimulating factor. Then CD34⁺ cell selection of the graft was performed and anti-thymocyte globulin was given for T-cell depletion, with the conditioning regimen BEAM adopted and early and late toxicities recorded. Long-term responses were evaluated by the expanded disability status scale (EDSS), progression-free survival, and gadolinium-enhanced magnetic resonance imaging scans. Ten, seven, and eight patients experienced neurological improvement, stabilization, and progression, respectively.</td>
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<tr>
<td>Mesenchymal stem cells (MSCs)</td>
<td>Al Jumah et al, 2012⁵⁴</td>
<td>Preclinical: EAE mouse model</td>
<td>The study evaluated the immunomodulatory and neuroprotective effects of MSCs in EAE for MS. The study concluded that MSCs can rescue neural cells via a mechanism that is mediated by soluble factors, which provide a suitable environment for neuron regeneration, remyelination, and cerebral blood flow improvement.</td>
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<tr>
<td>Autologous MSCs – bone marrow-derived cells</td>
<td>Connick et al, 2012⁵⁵</td>
<td>Clinical human studies</td>
<td>This was an open label phase 2a proof of concept study in patients with secondary progressive MS with characteristic visual pathways (expanded disability status score: 5.5–6.5). The intravenous infusion of autologous bone marrow-derived MSCs were injected, and after the period of 10 months, improvement was noted after treatment in visual acuity (difference in monthly rates of change −0.02 logMAR units, 95% CI: −0.03 to −0.01; p=0.003) and visual evoked response latency (−1.33 ms, 95% CI: −2.44 to −0.21; p=0.020), with an increase in optic nerve area (difference in monthly rates of change 0.13 mm², 95% CI: 0.04 to 0.22; p=0.006). There were no significant effects on color vision, visual fields, macular volume, retinal nerve fiber layer thickness, or optic nerve magnetization transfer ratio.</td>
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<tr>
<td>Embryonic-derived oligodendrocyte progenitor cells (OPCs) – MSCs</td>
<td>Cristofanilli et al, 2011⁵⁶</td>
<td>Preclinical mouse model</td>
<td>The study investigated the effect of syngeneic MSCs on the survival and remyelination abilities of allogeneic OPCs in adult non-immunosuppressed shiverer mice. At all time points examined, cotransplantation with MSCs increased OPC engraftment, migration, and maturation in myelinating oligodendrocytes, which produced widespread remyelination in the host corpus callosum. In addition, MSCs reduced microglia activation and astrogliosis in the brain of transplanted animals as well as T-cell proliferation in vitro.</td>
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**Abbreviations:** CPP, cell penetrating peptides; MS, multiple sclerosis; ES, embryonic stem; PPMS, primary progressive multiple sclerosis; CNS, central nervous system; iv, intravenous; icv, intracerebroventricular; logMAR, logarithm of the minimum angle of resolution; BEAM, BCNU, etoposide, arabinosylcytosine, melphalan.
as there is a limitation in the preparation of sufficient human OPCs to get tissue-specific progenitor cells, though some preclinical studies have tried to differentiate mouse ESCs (mESCs) into oligodendrocyte with myelogenic properties.52,53 Aharonowiz et al transplanted hESC-derived neural progenitors into the mice with experimental autoimmune encephalomyelitis. Histological evaluation revealed that the transplanted neural progenitors migrate to the mice brain, especially to the host white matter. However, remyelination and production of mature oligodendrocytes were not clearly observed.54

How do hESCs work in treating neurological disorders?

For the therapeutic action of the cells, it is necessary that the transplanted cells should reach the target site or the site of injury, also referred to as “homing.” Although, no study till date has assessed the mechanism of action of hESCs, various hypotheses have been suggested and discussed to explain the mechanism of action of hESCs for treating MS and other autoimmune disorders (Figure 1), which are discussed in the following section.

Differentiation

Human ESCs differentiate efficiently to form trophoblast, the outer layer of the placenta that mediates implantation and connects the conceptus to the uterus. hESCs have also been observed to differentiate into cells expressing genes characteristic of germ cells.55–57 In particular, the ability of hESCs to differentiate into defined neural lineages, neurons, astrocytes, and oligodendrocytes is a fundamental characteristic that has been exploited for the treatment of neurological disorders.58,59

The studies also observed that the transplanted stem cells act by combining with local stem cells in the injured tissue to accomplish the healing process. Liu et al reported that the immune system of the patient releases signals (interferon [IFN]-γ and TNF-α) to communicate with the transplanted MSCs during the repair of the injured tissue. Thus, the stem cells get attracted to the site of injury.60 Other studies identifying the homing of MSCs to the target site via the SDF-1/CXCR4 pathway reported that SDF-1, a chemoattractant, is released by the injured tissue which attracts the transplanted stem cells.61,62 VEGF is another key mediator of mobilization.63 It is hypothesized that hESCs might follow the same pattern to migrate to the target site and initiate regeneration and repair processes. Angiogenesis, improving cerebral blood flow to treat the neurological disorder, is another mechanism by which hESCs work. The more likely mechanisms of action include the release of VEGF and increase in the endogenous levels of other factors (such as brain-derived neurotrophic factor and fibroblast growth factor) that play a role in neovascularization.64

Various studies of cell transplantation have reported the mechanism of angiogenesis to treat the disorders. Studies conducted by Taguchi et al had reported that systemic administration of human cord blood-derived CD34+ cells to immunocompromised mice subjected to stroke 48 hours earlier induced neovascularization in the ischemic zone and provided a favorable environment for neuronal regeneration.65

Reduced apoptosis and release of trophic factors

The neuroprotective response is elicited by cell transplantation by reducing the apoptotic cells, particularly in penumbra tissue. This is mediated by the secretion and upregulation of certain trophins, such as basic fibroblast growth factor, brain-derived neurotrophic factor, VEGF, glial cell-line-derived factor, and nerve growth factor. It helps in reducing apoptosis in injured tissue and aids in neurological recovery.66

Reduced inflammation

Immunomodulatory effects of the transplanted cells play a major role in neuroprotection by reducing infiltration of macrophages and microglial activation and inhibiting T-cell activation. Preclinical studies have shown evidence for upregulation of anti-inflammatory cytokines and attenuation of expression of proinflammatory cytokines in both ischemic and hemorrhagic stroke.67,68

Small size

Another remarkable characteristic is that these cells are very small in size (<1 um) and have very high multiplication rate. Their small size makes it possible for them to cross the blood–brain barrier and reach the target sites easily. These cells resemble and behave like previously very small embryonic stem cells.43

Comparison of other cellular therapies with hESCs

Current cellular therapies for MS are largely palliative, not curative. MSCs are regenerative and immunosuppressive, indicating a potential therapy for MS, yet the variability and low potency of MSCs from adult sources hinder their therapeutic potential. Similarly, NSCs and neural progenitor cells (NPCs) are associated with limitations of appropriate microenvironment needed for differentiation into oligodendrocytes. Various studies have reported the neuroprotective and immunomodulatory effects of transplanted exogenous
Factors causing autoimmune responses: genetic and environmental factors, molecular mimicry responses, Epstein–Barr virus infection of B cells, TLR signal activation

Blood–brain barrier

Antigen presentation

Auto reactive T cell (CD4+ or CD6+)
activation

TLR signal activation

Lessons formation

Clonal expansion

Vasular alterations

Chemotactic factors and cytokine production

Astrocytes, oligodendrocytes, and microglial cell activation and causing demyelination

Healthy neuron

Myelin sheath

Axon

Demyelination in progress

CD20 B cell

Antibody production

Mechanism of action of hESCs

Stimulate angiogenesis-Increase cerebral blood vessels

Anti-inflammatory-reducing infiltration of macrophages and microglial activation and inhibiting T-cell activation

Human embryonic stem cells

Antiapoptopic-reduce cell death

Produce trophic factors-secretion and upregulation of certain trophins, such as basic fibroblast growth factor, brain-derived neurotrophic factor, VEGF, glial cell-line-derived factor, and nerve growth factor

NSCs on T-cell activation, microglial activation, and endogenous remyelination and their effects on the pathological process and prognosis in animal models of MS. However, a lack of neural progenitor sources and immune rejection are the two hindrances restricting the development of this new MS treatment strategy. HSCs are obtained from bone marrow, peripheral blood, or umbilical cord blood. They are suitable for both autologous as well as allogeneic use, but are associated with the issues regarding consistency of numbers and potency of cells obtained from the bone marrow, as well as the need for ex vivo expansion of cells when using umbilical cord blood as the source.

The therapeutic potential of iPSCs has been studied in various animal models. However, at the same time, there is a risk of tumor formation due to the residual undifferentiated iPSCs. Also, some studies reported the cases of antigen-specific immune rejection responses after transplantation into the autologous recipients. hESCs have shown substantial results in treating MS than stem cells derived from adult bone marrow. A recent research conducted by Wang et al, where they compared eight lines of adult bone marrow stem cells to four lines of hESCs, showed that all of the bone marrow-related stem cells expressed high levels of a protein molecule called cytokine that stimulates autoimmunity and can worsen the disease, whereas all of the hESC-related lines expressed low levels of the inflammatory cytokine.

hESC therapy is not only a safe, effective, and efficient treatment for MS, but is also a cost-effective treatment with better results of recovery. Since remyelination is an irreversible process, the frequency of relapses is quite low or negligible as compared to the conventional therapies where the treatment has to be continued for rest of the life of patient. Comparatively, hESC treatment is generally a one-time treatment plan, and the patient gets rid of further relapses with single use of therapy.
hESC cell line developed at our institute and its uniqueness

The hESC line at our institute was developed using patented technology (United States patent US 8592, 208, 52) in a good manufacturing practice, good laboratory practice, and good tissue practice compliant laboratory (United States patent WO 2007/141657A PCT/IB 2007, 2007 Dec 13).71

The cell lines used in our study were obtained from a single fertilized egg after 24–48 hours of fertilization. Since a single fertilized egg is able to provide therapeutic amounts of hESCs and/or their derivatives to treat multitudes of patients, there was no repeated exploitation of human embryos, and the number of ethical issues associated with the use of hESC therapy could be avoided. The use of hESCs at our institute has also been accepted and confirmed by the House of Lords, Regenerative Medicine, Science, and Technology Committee.72

Xeno-free culture medium and appropriate size

While maintaining the culture conditions of hESCs, there are chances of chromosomal aberrations, which are difficult to recognize by conventional karyotypic analysis techniques. These chromosomal alterations seem to be more common when hESCs are grown in feeder-free conditions and maintained using enzymatic rather than mechanical dissociation methods.73–75 The culture media used was free of any antioxidants, growth factors, insulin or insulin substitutes, collagen precursors or collagen precursor substitutes, residues or “conditioned media,” trace elements, animal products, and feeder cells. The absence of any xeno-product makes it suitable for clinical cell therapy. The cells thus obtained are very small in size (50 nm–2.5 µm) and procured 24 hours after fertilization.71,76

Unique characterization

The cell lines obtained at our institute harbor all the properties of hESCs and blastocysts and express PSC markers like octamer-binding transcription factor 4, (sex determining region Y)-box 2, Nanog, stage-specific embryonic antigen-4, trophoectoderm marker; keratin 18, beta-human chorionic gonadotropin (negative), immune-regulatory marker; human leukocyte antigen G (negative), gene activating marker 5-methylcytosine, and other markers like telomerase and α fetoprotein.76

Genomic stability and cell viability

Nonenzymatic passaging method was used and regular checks on spontaneous differentiation were conducted. All these considerations have resulted in maintenance of stable cell lines over the years on repeated passaging. These cell lines are in clinical use for more than 14 years and have undergone >4,000 passages. The cell lines have been characterized at cellular, molecular, and functional levels and established for its safety and efficacy. These have been evaluated for their viability as well as genomic integrity. A special cryopreservation technique was used where a relatively low concentration of DMSO (0.2%–2%) has been used, which is helpful in preventing any possible toxicity to patients. Our analyses of viability of these cells showed that 95% of the cells were alive after being thawed for more than 9 years.76,77

No immune rejections

Most of the previous studies have targeted either four single blastomeres (four-cell and five-cell stage) or morula embryos (eight-cell stage) as a source to derive hESCs.78–80 hESCs prepared at our institute are unique in their procedure of development, are harvested at very initial stage of blastocyst formation, i.e., two-celled stage blastocyst, post-pronuclear fertilization and first cell division, a stage at which cells initiate to differentiate or rather are in the transition phase and do not acquire any antigenic property. At this embryonic stage of harvesting, genesis is not activated, thus these cells are safe for transplantation which declines the risks of immune rejection.71

Other clinical applications

The cell lines have been used safely in various clinical applications including neurological disorders such as MS, Parkinson’s disease, cerebral palsy, Friedrich ataxia, amyotrophic lateral sclerosis, Duchenne muscular dystrophy, Lyme disease, autism, traumatic brain injury, and many more.81–87 Table 2 presents the summary of our hESC lines used in various clinical applications and their results in patients.

hESC lines used in MS

The cells are of two types: predominantly neuronal and non-neuronal (menenchymal, ocular, germ, etc.) progenitor cells with their undifferentiated states. This study was performed on the generation of different neuronal cells from an hESC line produced from biopsied blastomeres at the two-celled cleavage stage from a discarded embryo during an in vitro fertilization process. The differentiation of neuronal cells by determining the lineage-specific neuronal marker, Neu N, was also explored. It was found that these multipotent NPCs differentiated into all three types of cells of the nervous system.
system, i.e., neurons, oligodendrocytes, and astrocytes, as observed under a phase contrast microscope.88

The uniqueness of these cells is that they are in a ready-to-inject form and can be used in a number of otherwise incurable conditions. In patients with MS, hESC therapy was given as per the standard treatment protocol which included treatment phases (8–12 weeks) with gap phases (4 months) wherein 0.25 mL (<4 million cells) of hESCs was administered intramuscularly twice a day to ensure that the recipient’s immune system does not reject the stem cells (“prime” the body). About 1 mL of hESCs (<16 million cells) was administered every 10 days intravenously (to “home in” to the required area) and 1–5 mL of hESCs was administered every 7 days. It was observed that these hESC cell lines have the homing capability because of their therapeutic effect at the injured or inflamed tissue. After 2 months of treatment, patients showed remarkable improvement in overall stamina, muscle strength, functional skills, and cognitive abilities; the patients experienced an increased energy level and power of upper limbs. The body weight was also increased.

The radiological investigation using MRI and tractography suggested that there was a significant improvement in the condition of the patient. Tractography studies

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<th>Clinical application</th>
<th>Type of study</th>
<th>Results</th>
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<td>Cerebrovascular accident</td>
<td>Case series (2015)</td>
<td>Twenty two patients were evaluated for improvement on the basis of European Stroke Scale (ESS) at baseline and at the end of each treatment period. A total of 22 patients were included and all received intensive dosing with hESCs. All affected patients showed an improvement in gait (22 patients), speech (15 patients), level of consciousness (two patients), comprehension and gaze (one patient each) by at least one point at the end of T6. In addition, patients showed improvement in walking, balance (sitting and standing), and spasticity after receiving hESC therapy. Overall, 11 patients (50%) experienced adverse events (AEs) during the study. No serious adverse events (SAEs) and deaths were reported. Conclusion: All the patients showed improved cognitive skills and regained their functional ability. No severe AEs or SAEs were reported during the study. hESC therapy was well tolerated among all the patients included in the study.</td>
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<tr>
<td>Parkinson's disease</td>
<td>Case report (2015)</td>
<td>The study was on a 65-year-old patient suffering with Parkinson’s disorder. With the help of hESC treatment, the patient showed significant improvement in his heath following the treatment, such as reduction in tremors, bradykinesia, muscle rigidity, pain and stiffness in the neck, shoulder and low back, and improvement in the range of movements of the neck. The patient was able to balance himself while walking, and had reduction in numbness in his left hand and both of his legs. He was able to speak louder and had improved writing skills. No AE or teratoma formation was observed.</td>
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<td>Spinocerebellar ataxia</td>
<td>Case series (2015)</td>
<td>This was a case study of three patients. All received hESC treatment for 2 years. After therapy, all showed improvement in overall stamina, endurance, coordination, sitting balance, standing and walking ability, speech, and flexibility. There was a reduction in tremors and no head nodding was observed.</td>
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<tr>
<td>Cerebral palsy (CP)</td>
<td>Case series (2015)</td>
<td>The study included two patients suffering from CP, after undergoing treatment with hESC therapy. The treatment consisted of phases in which 0.25 mL of hESCs (&lt;4 million cells) was administered through intramuscular route once daily and 1 mL of hESCs (&lt;16 million cells) was administered through intravenous route twice every 7 days. A marked improvement in cognitive and motor functions was seen in the patients after the treatment.</td>
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<tr>
<td>Retrospective study</td>
<td>Retrospective study (2015)</td>
<td>The study included 91 patients; SPECT scan showed improved perfusion in all patients. Cognitive skills improved in 69% of the patients; there were no seizures (n=90) and hearing improvement was observed (n=8).</td>
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<tr>
<td>Multiple sclerosis (MS)</td>
<td>Case report (2015)</td>
<td>A case report of a 34-year-old female with MS. After treatment, tractography showed a mild reduction in the size of lesions in bilateral periventricular white matter and in the right occipital white matter. Improvement in muscle bulk, tone, and power had increased energy level and power of upper limbs and gained weight.</td>
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<tr>
<td>MS and Lyme disease (LD)</td>
<td>Case series (2016)</td>
<td>The study included one male (42 years) and one female (30 years) with both MS and LD. The hESCs were administered via different routes to regenerate the injured cells. Both the patients showed remarkable improvement in their functional skills, overall stamina, cognitive abilities, and muscle strength. Furthermore, the improvements in the patients’ conditions were assessed by magnetic resonance tractography and SPECT.</td>
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</table>

Abbreviations: hESCs, human embryonic stem cells; SPECT, single photon emission computed tomography.
interpreted the extent of improvement in perfusion, which showed that there was a mild reduction in the size of lesions in bilateral periventricular white matter and another lesion in the right occipital white matter. The remyelination in the spine was clearly visible in the tractography of spine and MRI performed showed improvement in baseline and subcortical regions of the brain after the therapy. The patients were also scored using expanded disability status scale. The scores obtained before and after the treatment showed remarkable improvement in muscle power, bulk, and tone after the hESC therapy. Neither of the patients reported any AEs following the hESC therapy. Further follow-up studies are required to be done with patients treated with hESCs to evaluate their use. Since the use of stem cell treatments for patients with MS is at a much earlier stage of development, it is not yet carried out routinely and treatment protocols have not been standardized. Treatment may be available through a clinical trial or, rarely, outside a trial if the clinical need is clear. Prior to our studies to treat MS with hESC, there is no study reporting the management of chronic MS in humans with hESC therapy.

**Conclusion**

Stem cell therapy is undoubtedly one of the remarkable achievements in medical history. This technology has revolutionized the laboratory cell biology and will provide much improved cell culture models for discovery and development of drugs, and fundamental studies of the genetic basis of disease. hESC therapy with its unique properties has shown promising results to treat complex neurological disorders like MS for which other cellular therapies have failed. MS, which is considered an incurable and lifelong disability, can be cured with hESC treatment.

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**Disclosure**

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