Regenerative capacity of bone marrow stem cells with or without superparamagnetic iron oxide nanoparticles after facial nerve degeneration: A narrative review

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ABSTRACT

Facial palsy can be defined as a kind of paralysis affecting facial muscles. It is termed Bell's palsy if it is unilateral. It may occur due to trauma to the facial nerve, infections as herpes zoster, neoplastic lesions, or unknown cause. It may be also associated with metabolic and systemic diseases as hypertension, toxicity, amyloidosis, alcoholism, auto-immune diseases and diabetes mellitus. Mesenchymal stem cells (MSCs) are multipotent adult stromal cells that have many benefits as an evolving treatment modality. Bone marrow stem cells (BMSCs) divide progressively in culture, and differentiate into neurons exclusively with use of a simple protocol. Most ongoing preclinical and clinical cell treatment modalities composed of local or systemic transplantation of stem or progenitor cells. In addition, they depend on the migration and retention of transplanted cells at insult areas. Nevertheless, one of the main obstacles against this modality is how to detect the fate and exact location of these cells inside the body, and how to maintain the cells at this specific site. Magnetic targeting systems, which depends on cells labelled by magnetic carriers, have been assessed as a more efficient technique for stem cell delivery to target sites. These systems depend on loading stem cells with magnetic nanoparticles and attracting them to the exact intended area within the body by placing an external magnetic field. Superparamagnetic iron oxide nanoparticles (SPIONs) have been introduced in the last few years as a rising applicant of nanoparticles in a vast variety of medical fields as magnetic separation, drug delivery, magnetic resonance imaging (MRI) and magnetic hyperthermia. In addition, applications of SPIONs, as a site-specific drug carrier, diagnostic agent and stem cell delivery agent, receive most attention of researchers in that field. In this review, up-to-date information about Magnetic targeting of degenerated facial nerve by BMSCs labelled with SPIONs may suggest its capacity of better regeneration than injection of BMSCs alone.

Keywords: bone marrow stem cells, facial nerve, magnetic targeting, superparamagnetic iron oxide nanoparticles

INTRODUCTION: FACIAL NERVE

Facial nerve is a cranial nerve that passes in the parotid gland to provide nerve supply to the muscles of the face. It has five main branches which are: the temporal, zygomatic, buccal, mandibular, and cervical branches. Normal activity of facial muscles performs a very important function in a person’s physiological, psychological, and emotional reactions. Social and vocational disabilities, which is formed from all these components, can be caused by facial disfigurement (1).

FACIAL NERVE DISORDERS

Facial nerve carries motor, sensory, and parasympathetic fibers, therefore, both functional and cosmetic impairment can be caused by facial palsy. Facial nerve palsy is clinically detected when weakness of the facial muscles is diagnosed. It can be diagnosed by pain around the ear, curving in the angle of the mouth, improper mouth closure, dryness of the eye, hyperacusis, improper taste sensation, incomplete lid closure or eye brow immobility (2).
ETIOLOGY OF FACIAL NERVE DISORDER

Facial nerve diseases can arise from a wide variety of disorders. The origin of these disorders may be traumatic (as basal skull fractures and facial injuries) (3), neurologic (4) or infectious (5). Facial nerve degeneration can also arise from metabolic causes (as diabetes mellitus (6), hyperthyroidism (7) and hypertension (8)) , neoplastic causes (9), and iatrogenic causes (as mandibular block anesthesia and other dental procedures) (10). Some chemicals can cause toxic degeneration of the facial nerve as thalidomide (11), ethylene glycol (12), arsenic intoxication (13) and local anesthesia (14).

TISSUE REGENERATION

Insults to any living tissues are counteracted by a natural sequence of actions for healing through repair or regeneration. However, natural repair and regeneration can lose adequacy and regulation in case of major trauma or surgery, leading to undesirable results as fibrosis and aging. So, recent researches are aiming to use regenerative cells to restore the living tissue, this is what is called tissue regeneration technology (15).

STEM CELLS

Stem cells are defined as the cells which have less or undifferentiated phenotype, and have the ability of self-renewal and production of different types of cells. Stem cells can be classified according to their lineage into embryonic and adult stem cells. Embryonic stem cells (ESCs) are pluripotent cells derived from the embryoblasts of the blastocyst that can be propagated endlessly in an undifferentiated state. When ESCs are cultured in a convenient environment, they can differentiate into ectodermal, endodermal and mesodermal cells. Adult stem cells are part of tissue-related cells of organism after birth. They can differentiate to the cells of their original tissue. Adult stem cell plasticity term, which evolved in the past years, means that stem cells of specific organ can differentiate into cells of other diversity of organs. This term has greatly changed the idea that tissue-specific stem cells could only differentiate into cells of the tissue source (16).

NERVE TISSUE ENGINEERING

Establishment of a more supportive environment for regenerating axons, in addition to maintaining this support for an extended period of time, is considered the main target of any stem cell treatment advance to peripheral nerve trauma. Increasing amount of stem cells around motor and sensory neurons would probably provide higher numbers of neurons for regeneration, which could aid regeneration in an additive fashion to the establishment of a favorable environment (17).

Transplantation of stem cell into muscle deprived of nerve supply may present a promising method to allow muscle reinnervation and activity. Moreover, muscle atrophy and leaving the tissue without nerve supply can be prevented over extended time intervals (18,19).

BONE MARROW STEM CELLS

BMSCs are form of adult stem cells that can yield variety of cells, even if they are not mesenchymal, by differentiation, dedifferentiation, and transdifferentiation. It has been validated that they are able to produce cardiomyocytes (20), hepatocytes (21), osteocytes (22), neurons and Schwann cells (23).

Animals with peripheral nerve injury, which received topical or systemic transplantation of BMSC, showed BMSCs transfer and specific occupying in traumatized neural areas (spinal cord and injured nerves). This is associated with an intense decrease in pain manifestations, reduction of occurrence risk of any neurochemical variation often detected in neurons during peripheral nerve injury, and a faster degeneration-regeneration process, reflected upon electromyographic readings (24). On this base, a number of researches depended on BMSCs for regeneration of the facial nerve as listed in table 1.

CHALLENGES FACING STEM CELL THERAPY

Most ongoing cell treatment modalities (whether in vivo or in vitro) composed of local or systemic transplantation of stem or progenitor cells. In addition, they depend on the transfer and habitation of transplanted cells at insult areas. Nevertheless, one of the main obstacles against this modality is how to detect the fate and exact location of these cells inside the body, and how to maintain the cells at this specific site. To overcome these obstacles, researchers were trying to find out methods to increase the occupation of transplanted cells at the intended areas for treatment in the body (32).

NANOTECHNOLOGY

Nanotechnology refers to the formulation and usage of materials, which have components of very small size, measured in the nanoscale (i.e., their sizes are up to 100 nm). Nanotechnology is used nowadays in a wide variety of fields as electrical, optical, and magnetic fields as well as structural performance at the molecular and submolecular level. Their characteristics of being safe, cheap, easily carried and administrated, give them the capacity to
TABLE 1. Researches collected from PubMed database estimating the impact of BMSCs on facial nerve regeneration in vitro and in vivo

| Author (publication year) | Study design | Aim | Result |
|---------------------------|--------------|-----|--------|
| Grosheva et al., 2008 (25) | In vivo | Transected facial nerve model | To evaluate whether manual stimulation could be beneficial after a surgical procedure for treatment of large peripheral nerve defects | BMSCs failed to diminish the degree of collateral axonal branching at the injury region and did not correct the physiological impairment |
| Aggarwal et al., 2012 (26) | In vivo | Facial nerve paralysis model | To study the safety profile and role of BMSCs in the rehabilitation of posttraumatic facial nerve paralysis, which is not treated by conventional therapy | Stem cell therapy can be used safely in human beings without any adverse effects on humans |
| Costa et al., 2013 (27) | In vivo | Animal rat model | To estimate histological and physiological impact of BMSCs with polyglycolic acid tube in autografted rat facial nerves | Schwann-like cells were associated with superior outcomes. BMSC introduced in neural tissue allowed maintenance of former cell phenotype for six weeks |
| Salomone et al., 2013 (28) | In vivo | Animal rat model | To evaluate (BMSC) in a silicone conduit for rat facial nerve regeneration from isolated stumps. | Regeneration improvement by both Ubmsc and dBMSC in rats, however, uBMSC was associated with superior functional results |
| Lucena et al., 2014 (29) | In-vitro | Quantitative phenotypic analysis | To evaluate the plasticity of BMSCs of mice in the existence of culture medium conditioned with facial nerve explants and fibroblast growth factor-2 (FGF-2) | BMSCs showed typical morphology and fast expansion, especially the cells that were subjected to a medium conditioned by the facial nerve |
| Ge, Yining et al., 2018 (30) | In-vivo | Animal rat model | To detect the technique of the immunomodulation impact of BMSCs on facial nerve trauma | BMSCs can secrete cytokines IL-6, HGF, PGE2, iNOS, and TGF-β1 in an independent manner. These cytokines might set coordination among subsets of CD4+ T cells, inhibiting neuron apoptosis, thus preserving the integrity of facial nerves |
| Li Wu et al., 2020 (31) | In vivo | Animal rat model | To study the impact of co-transplantation of BMSCs and monocytes in facial nerve axotomy. | Significant improvement in the facial nerve nucleus in axotomy rats, enhancing the coordination of the chemotaxis, homing, differentiation of BMSCs, with diminishing neuronal apoptosis |

remodel an array of medical and biotechnology applications (33).

MAGNETIC TARGETING OF MSCS

Last few decades have witnessed the emergence of magnetic targeting systems as a prominent technique for site-specific targeting of various pharmacological agents or stem cells. With the effect of a magnetic field, it avoids reticuloendothelial system and directs the drugs or stem cells, to reach the target accurately (34). Magnetic carriers could be nanoparticles, microspheres, liposomes and emulsion. Magnetic targeting of MSCs includes the following processes: 1) isolation, growth, and preservation of MSCs in culture media; 2) magnetization of MSCs; and 3) path control of magnetized MSCs via static magnetic fields (32).

SUPERPARAMAGNETIC IRON OXIDE NANOPARTICLES

Superparamagnetic nanoparticles are those which have size less than 30 nm, i.e., their magnetization happens only when an external magnetic field is present. This is a beneficial property for biological applications. Therefore, SPIONs are crucial materials for several medical purposes of MSCs with magnetic targeting technology. In addition, they can also be used as contrast agents for MSC labeling and tracking in the body due to the powerful signal they generate in magnetic resonance imaging (MRI) (35).

SPIONs has many beneficial properties as easy synthesis, biocompatibility, multifunctionality, and possibility of further surface modification with various chemical agents, SPIONs can aid in many domains of medicine. SPIONs have also some disadvantages, such as their high uptake by macrophages. However, they sound to be very promising in stem cell treatment modalities depending on the results the current researches (35).

THE USE OF BMSCS WITH SPIONS IN NERVE REGENERATION

Zhang et al. (2016) studied the impact of magnetic targeting of NT3 gene-transfected BMSCs through lumbar puncture in a rat study design of SCI. They
proved that delivery of transplanted BMSCs to the exact injury site, in addition to performing cell imaging via MRI, could be achieved using magnetic targeting cells delivery system. Moreover, the study also reported that this treatment modality could markedly enhance functional recovery and nerve regeneration, in comparison with transplanting NT3 gene-transfected BMSCs without magnetic targeting system (36). These outcomes may put in mind that the use of BMSCs with SPIONs could be a promising method of treating other degenerated nerves (i.e. facial nerve).

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CONCLUSIONS

BMSCs therapy is considered as a promising choice for treatment of facial nerve degeneration. However, localization of the injected cells within the injury site remains a challenge against the effectiveness of this treatment. We suggest that the use of BMSCs with SPIONs as a magnetic targeting system may allow definite delivery of stem cells to the injured facial nerve, and so, enhancing the effectiveness of the treatment.
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