Review: Bioengineering approach for the repair and regeneration of peripheral nerve

Joshua Moskow\textsuperscript{a,b,1}, Bryan Ferrigno\textsuperscript{a,1}, Nikhil Mistry\textsuperscript{a}, Devina Jaiswal\textsuperscript{a,b}, Ketan Bulsara\textsuperscript{c}, Swetha Rudra\textsuperscript{a,d}, Sangamesh G. Kumbar\textsuperscript{a,b,∗}

\textsuperscript{a}Department of Orthopaedic Surgery, University of Connecticut Health, 263 Farmington Ave., Farmington, CT 06030, USA
\textsuperscript{b}Department of Biomedical Engineering, University of Connecticut, 260 Glenbrook Road, Unit 3247, Storrs, CT 06269, USA
\textsuperscript{c}Department of Surgery, University of Connecticut Health, 263 Farmington Ave., Farmington, CT 06030, USA
\textsuperscript{d}Department of Pharmaceutical Sciences, School of Pharmacy, University of Saint Joseph, 229 Trumbull St., Hartford CT 06103, USA

\textbf{ARTICLE INFO}

\textbf{Keywords:}
Peripheral nerve regeneration
Composite materials
Growth factor
Electrical stimulation

\textbf{ABSTRACT}

Complex craniofacial surgeries of damaged tissues have several limitations, which present complications and challenges when trying to replicate facial function and structure. Traditional treatment techniques have shown suitable nerve function regeneration with various drawbacks. As technology continues to advance, new methods have been explored in order to regenerate damaged nerves in an effort to more efficiently and effectively regain original function and structure. This article will summarize recent bioengineering strategies involving biodegradable composite scaffolds, bioactive factors, and external stimuli alone or in combination to support peripheral nerve regeneration. Particular emphasis is made on the contributions of growth factors and electrical stimulation on the regenerative process.

1. Introduction

The craniofacial skeleton and related tissues, including nerve and bone, are involved in several major functions ranging from protecting the brain to speaking, hearing, and breathing [1]. Specially, craniofacial nerves are involved in mechanisms to detect and react to changes in one’s internal and external environment and are integral in maintaining proper overall function [2]. Due to their complex structure and networking, craniofacial repair surgeries have limitations and often result in insufficient restoration of facial function and structure. As a consequence, there are limited strategies for nerve repair and regeneration thereby over 20 million Americans experience nerve impairment exceeding a cost of $150 billion [3]. This creates a critical need for nerve regeneration, especially for craniofacial nerves and their peripheral extensions.

The nervous system is divided into two sections: the central nervous system (CNS) and the peripheral nervous system (PNS). The PNS is comprised of both sensory and motor neurons which carry information from all the parts of the body to and from the CNS. Cranial nerves are also categorized under peripheral nervous system and emerge from the brain or brain stem to innervate areas of head and neck. The facial cranial nerves innervate facial muscles that are responsible for expressions. A peripheral nerve consists of a cell body which gives out extensions, called axons that are crucial for targeting distant tissues and organs [3]. These axons are coated with myelin sheath membranes, formed by Schwann cells, and are arranged together in bundles called fascicles [3] (Fig. 1). Due to the complex anatomical structure of nerve bundles and its mechanism of degeneration, nerve repair and regeneration strategies for critical defects have often resulted in failure to re-establish sufficient nerve function.

Various strategies for repair have been implemented depending on the type of trauma experienced [4,5]. In the event of peripheral nerve injury (PNI), a regulated sequence of events involving cellular bodies in the spinal cord and ganglia occur in order to regenerate damaged nerves. Nerve injuries are classified into three broad categories depending on the severity of the injury: neuropaesthesia, axonotmesis, and neurotmesis [4]. Neuropaesthesia is the least severe type of injury and is not associated with long term loss of function [6]. In contrast, axonotmesis occurs when there is a disruption of the nerve axon and the surrounding myelin sheath. The most severe type of nerve injury is neurotmesis,
which involves a complete disconnection of the nerve and will be the focus for this review. This type of nerve injury results in a complete loss of function and axonal regrowth is limited due to potential formation of scar tissue, known as neuroma [4]. For neurotmesis repair, current methods include direct end to end repair, grafts and synthetic nerve conduits (Fig. 2). These nerve replacement strategies are used clinically with direct end to end suturing, considered the current preferred standard [6]. Direct end to end suturing is the easiest method but cannot be used with critically sized injuries as the tension induced from stretching the nerve can cause functional failure. On the other hand, the use of grafts for nerve repair is dependent on availability of the tissue, location from which the tissue is obtained and immune response from the host. These factors can limit the success of grafts for nerve regeneration. Nerve grafts can be autografts or allografts. Autografts are derived from the healthy part of patient’s body thereby it reduces the chances of immunological rejection. However, this causes tissue damage at the donor site, termed donor site morbidity, and therefore limits its applicability [6]. For allografts, the tissue is harvested from another donor of the same species. While this increases availability, it introduces a minute risk of disease transmission and immunological response. Due to the limitations of the above two methods, nerve conduits made from biocompatible polymers such as polyglycolic acid (PGA), collagen, and polycaprolactone (PCL) are currently being used [7,8]. However, current conduits can only be used for defects of a few centimeters and primarily function as structural support and concentrates of clotting factors between the distal and proximal stumps [6]. Alternate methods from the gold standards of current clinical practices involve tissue engineering strategies using composite biomaterials.

The basic paradigm of tissue engineering is based on design and fabrication of novel biomaterial scaffolds that can house cells and deliver biomolecular and physical signals to cells for successful tissue regeneration. A variety of biomaterials are available from synthetic to natural sources for tissue engineering applications. Main categories of biomaterials include metals, ceramics, and polymers. Among these, polymers have ideal properties such as high ductility, superior tensile strength, and ease of suturing for soft tissues such as nerve. Polymers are also able to affect the physicochemical and mechanical properties of a biomimetic matrix. Generally, polymers can be characterized as synthetic or natural. Currently, most common biodegradable synthetic polymers used in medicine are polyactic acid (PLA), polyglycolic acid (PGA), copolymers of PLA and PGA (PLGA), polycaprolactone (PCL), polyanhydrides, polyorthoesters, polycarbonates, and polyfumarates [9,10]. These polymers are synthesized from monomer units in a laboratory through various form of polymerization such as free radical polymerization. On the other hand, natural polymers, such as collagen, chitosan and silk fibroin [6,11,12], are derived from natural sources such as rat tail, crab shells and silk worm, respectively.

The success of any scaffold used in surgery is dictated by its bio-compatibility, porosity, bioreabsorbability, and mechanical strength [1]. First, a scaffold with adequate biocompatibility will successfully integrate with the native tissue without producing extreme foreign body response. This ensures successful long and short term implantation of these materials. Tissue integration is also dependent on porosity of the scaffold. Porous scaffolds with interconnected pores allow cellular infiltration and proliferation to enable tissue regeneration and integration [13,14]. A scaffold used for soft tissue must reabsorb to be slowly replaced by regenerated tissue. For being bioreabsorbable, a material must break down over time in the body into non-toxic subunits. This material property ensures prevention of necrosis of the native and newly regenerated tissue. Last, the success of a scaffold is largely dependent on its mechanical properties. Mechanical properties such as tissue comparable tensile strength is essential in order to ensure proper load transfer and stability during implantation of the scaffold [1,15,16]. If the scaffold is not mechanically competent it will break under load and if it is too strong then it can lead to phenomena such as stress shielding leading to atrophy of surrounding natural tissue [17]. While this phenomenon is classic to bone, the principle applies to all tissues in the body. Many natural and synthetic materials possess some of these ideal properties, however, none are perfect. Therefore, biomaterial composite sites of both synthetic and natural materials provides an ideal solution by combining the optimal properties of each component into one system [13,18–20].

While polymeric materials provide effective biomimetic scaffolds for endogenous nerve cell attachment and proliferation, other biomolecular and physical signals are needed to stimulate cells to regenerate damaged tissue. Various biochemical molecules have shown to be effective in promoting regeneration of damaged tissue by supporting...
differentiation of recruited cells to mature neuronal fates. For example, fibroblast growth factor and insulin like growth factor support peripheral nerve regeneration [21–23]. Additionally, growth factors such as neural growth factor (NGF) induce differentiation of mesenchymal stem cells into mature neural lineages [24]. Aside from chemical cues, electrical stimulation is also known to not only promoting regeneration of axons but also provide signals for native cells to differentiate [25,26].

This review will focus on current biomaterial composites being developed for craniofacial nerve tissue engineering in the hopes of providing an improved treatment method to the current standard. Additionally, an in-depth look into the complimentary chemical and electrical cues that accompany these strategies will be investigated.

2. Peripheral nerve injuries

Neuronal axons in the PNS require axoplasmic flow from the main cell body for its survival. In an event of trauma, the axon distal to the point of injury suffers from Wallerian degeneration. The distal nerve end below the site of the lesion degenerates, while the proximal end (axon on the same side as the cell body) regrows outwards, eventually re-forming synaptic connections with nearby tissues [27]. Wallerian degeneration is a process unique to the PNS and Schwann cells (SC) play an important role in axonal regeneration.

Upon peripheral nerve injuries, an immune response occurs through activated Schwann cells and macrophages that leads to clearing the neuron and myelin debris at the distal nerve stump [28]. Schwann cells (SCs) are capable of denervation and provide a variety of functions during Wallerian degeneration. They grow and migrate towards the site of injury and serve as physical guides for the regenerating axon along the proximal side, forming the well documented “bands of Bungner,” serving as a scaffold [29]. In addition, SCs secrete a variety of neurotrophic factors and cytokines to assist in the regeneration process, including NGF, cell adhesion molecules, and extracellular matrix components [29].

As mentioned previously, myelin found on neurons in the PNS is formed by Schwann cells whereas CNS myelin is generated by oligodendrocytes. While oligodendrocytes and Schwann cells are often compared to each other in terms of function, a major difference between the two is in their ability to repair neurons after injury. Schwann cells promote nerve regeneration and repair, whereas oligodendrocytes inhibit neuron repair after an injury [27,30,31].

3. Bioinstructive nerve conduits

Recent advances in tissue engineering and regenerative medicine have promoted the use and study of manufactured nerve conduits. Nerve conduits are an alternative to allografts or autografts and are used for treatment of large nerve defects. These nerve guidance conduits (NGCs) have several advantages over donor nerve grafts such as ease of availability, limited scar tissue formation, reduced immune response and no donor site morbidity. Furthermore, chances of adverse immune responses can be significantly decreased with the use of appropriately sterilized biomaterials through techniques such as gamma irradiation and ethylene oxide treatment. The surface of the NGCs can also be altered to support axonal elongation between from the proximal to the distal stump [32]. Current tissue engineering strategies utilize 3D-scaffolds made from biomaterials designed for better biocompatibility, biodegradation and porosity [33,34]. Scaffolds serve as a bio-mimicking microenvironment and support for cells that eventually results in tissue regeneration. With time, the scaffold degrades which eliminates the need for removal of the implant from the body [35,36].

Based on the biomaterial used, manufactured nerve conduits can be categorized as: synthetic, natural, or composite. In general, synthetic nerve conduits provide higher degree of controllability, better mechanical properties, and poor bioactivity compared to their natural counterpart. Commonly used, U.S. Food and Drug Administration (FDA) approved, synthetic biomaterial such as polyactic acid (PLA) and polylactic-co-glycolic acid (PLGA) are known for low inflammatory response and ease of processing. PLGA has been shown to give control over its rate of degradation by altering the ratio of its monomer components [37]. Additionally, PLGA scaffolds have the unique ability of adhering to Schwann cells and directing their growth [38,39,39]. Recently, rapid prototyping techniques such as 3D bio-printing are being employed for synthetic nerve conduit fabrication. This technology constructs individualized synthetic nerve grafts with an anatomy and physiology similar to natural nerve [40]. Since 3D bio-printing uses non-thermal extrusion method, biomaterials combined with various cells and neural growth factors can be printed which has improved axonal regeneration of long-segmented nerve defects [40]. These scaffolds are utilized for their biocompatibility, tunable physiochemical properties, and ability to promote cell attachment.

Natural biopolymers used for the fabrication of NGCs typically have regenerative bioactivity along with good mechanical properties. Clinically, FDA approved collagen type I nerve conduits have been implemented for defects smaller than 3 cm [6]. For critically sized injuries other materials such as fibrin and silk have been investigated [41]. A multichannel electrospun silk nerve conduit, mimicking natural tissue ECM structure, was created by Dinis et al., which closely resembled the tubular structure of epineurial tube [42]. This scaffold possessed an ultimate tensile strength extremely close to that of a sciatic nerve [42]. A defect size of 15 mm is considered a limiting factor for nerve regeneration in rats compared to over 3 cm in humans [43]. An in vivo study compared autografts, synthetic and natural biomaterial conduits for critically sized defects [40]. The results showed that even though chitosan tubes were biocompatible and showed better regeneration than synthetic silicone tube, they were inferior to the autograft. Nerve regeneration in rats with autografts was 50% more compared to rats implanted with chitosan nerve conduits [40].

Even though natural and synthetic polymers have been used for tissue engineered scaffolds, research is also being done on composite polymers to enhance material properties. Composite polymers embody the material properties of two or more polymers prepared for a specific application. In nerve tissue engineering, composite nerve conduits can contain a blend of both natural and synthetic polymers. An example of a composite nerve conduit is PCL blended with collagen through electrospinning [44,45]. PCL is a synthetic polymer used for scaffolds for its porous nature, biocompatibility, cell proliferation, and cell adhesion properties [46]. Whereas, collagen is a structural protein present within natural ECM that activates integrin receptors on the ends of the distal axons and glial cells [35,44]. When fabricated together and seeded with cells, this conduit promotes neurite outgrowth, extension and glial migration. Comparing PCL to the PCL/collagen blend (C/PCL), it was seen that C/PCL increased Schwann cell migration, neurite outgrowth and fibroblast sheathing cells, which can be an appropriate material for nerve regeneration [44]. Biomaterial composites can also be formed by mixing two synthetic polymers with distinct properties such as polypyrrole (PPy) and poly(D, L-lactic acid) (PDLLA) composite conduit made and tested by Xu et al. [47]. Since nerve tissue is electrically active, PPy, a conductive polymer, promotes regeneration and differentiation of re-growing nerves. However, PPy is very brittle and non-degradable which makes it a poor biomaterial candidate. Addition of PDLLA, a biodegradable and non-cytotoxic polymer, improved the degradation rate and healing properties of the composite biocompatible conductive matrix [47].

While the properties of biomaterial clearly plays a vital role in peripheral nerve regeneration, one also has to account for the elasticity, geometry, and topology of the final scaffold. Specifically, aligned and grooved topologies on scaffold surfaces has shown enhanced extension compared to non-textured counterparts [48]. Two experiments done by Mobasser et al. highlight this effect. In both of their studies the group utilized non-grooved, sloped, square, and V-shaped morphologies for aligned PCL/PVA NGF films [32,49]. In the first study, hybrid
neuroblastoma glioma rat cells displayed enhanced cellular proliferation and elongation in the alignment direction on both the V-shaped and sloped geometries [32]. The group then tested their conduits in a 10 mm Sprague Dawley rat sciatic nerve defect. In vivo, the grooved samples showed a significantly higher number of regenerated axons at a 3 week time point compared to non-grooved conduits. Whereas the sloped topology conduits were not statistically different from autograft controls in terms of final target innervation and compound motor action potential amplitude [32].

Likewise, the elasticity of a scaffold is an important aspect to consider when utilizing it for nerve regeneration. As mentioned above, it is critical for biomaterials to exhibit specific properties in order to serve as suitable nerve conduits – namely, to be porous, adhesive, biodegradable, and have the appropriate amount of compliance/flexibility amongst other things [50]. The elastic modulus of the scaffold has been shown to impact neuronal stem cell differentiation and growth, thus it is an important parameter to consider [51]. Blending polymers together can maximize and optimize the properties needed to serve as a successful scaffold. Studies have shown that combining certain polymers together can optimize the elasticity, as demonstrated by the blending of PLGA and Poly ethyleneglycol [52,53].

4. Utilization of growth factors

Growth factors are a potential therapeutic option for supporting neuronal growth and enhancing peripheral nerve regeneration. They can be used synchronously with biomaterial composites to promote cell differentiation and proliferation. Extensive research within the literature reports the positive effects of certain neurotrophins that act on subpopulations of neurons in specific stages of development [54]. The three classic Neurotrophins are nerve growth factor (NGF), brain-derived neurotrophic factor (BDNF), and neurotrophin-3 (NT-3). All have affinity to different Receptor Tyrosine Kinases Receptors (RTKs) on target cells. RTKs are trans-membrane proteins that ultimately cause a sequence of cellular responses or a signal transduction cascade following substrate binding.

The mechanism of activation of RTK post neurotrophin binding can be based on an already existing model for epithelial growth factor receptor (EGFR) as shown in Fig. 3. Inactive RTK monomers are present across the plasma membrane comprising of three major domains: ligand binding extracellular domain, hydrophobic transmembrane domain and cytosolic domain [27]. As seen in Fig. 3, following substrate binding, the two monomers aggregate and dimerize. The tyrosine kinase domains on the cytosolic domain of the receptor then auto-phosphorylates the other monomer, with the phosphate groups derived from breaking down of ATP into ADP and Phosphate. After auto-phosphorylation, the active tyrosine kinase domains facilitates binding of other proteins to the phosphorylated tyrosine domains [27]. This triggers a conformational change within them, leading to a transduction cascade which ultimately results in cellular response, as depicted in Fig. 3.

Similar to EGRF model, Receptor Tyrosine Kinases serve multiple roles during nerve regeneration. As mentioned above, RTKs play a key role serving as a membrane receptor for major neurotrophic growth factors promoting axonal regeneration, survival, and neurite outgrowth. Additionally, RTKs are transported through retrograde mechanisms from neuronal projections back to the cell body, where they promote transcription and translation of proteins that serve to enhance long-term neuronal growth and survival [55-57].

4.1. Nerve growth factor

NGF in its mature form has high affinity for Tropomyosin Receptor Kinase A (TrKA). It promotes neuronal survival through its anti-apoptotic effects [58]. NGF only supports a limited set of neuronal populations [59] of the central and peripheral nervous system, specifically supporting the growth and survival of peripheral sympathetic and neural crest-derived sensory neurons [60,61]. Additionally, NGF can act on sympathetic (cholinergic) neurons as well as sensory neurons [62]. It promotes differentiation, survival and synaptic connections amongst neurons via transduction pathways including the Ras and PLC (phospholipase C) pathways [27]. NGF has been abundantly characterized which makes it an attractive candidate for in vivo and in vitro studies.

4.2. Brain-derived neurotrophic factor

BDNF supports neuronal growth and differentiation via Tropomyosin Receptor Kinase B (TrKB) activation [63]. Like other neuronal growth factors, BDNF’s effects are only on certain subpopulations of neurons. Brain-derived neurotrophic factors have been shown to promote survival of a subpopulation of sensory dorsal root ganglion neurons [60] while its effect on sympathetic neurons are minimal. BDNF’s basic functions also includes induction of neurite outgrowth of neurons [27,63] by altering local levels of Ca$^{2+}$ signaling in growth cones, thus enhancing directional growth and the neurite extension process [27]. In addition to supporting sensory neurons in dorsal root ganglia (DRGs) and neurons found in the inferior vagus ganglion, BDNF also supports the survival and outgrowth of motor neurons and their axons. A study confirmed that continuous administration of BDNF in vitro within chick embryos supported 40% of motor neurons that typically undergo degeneration and apoptosis during embryonic development [64].

4.3. Neurotrophin-3

Neurotrophin-3 has overlapping neurotrophic activity to NGF and...
BDNF and has high affinity receptor to Tropomyosin Receptor Kinase C (TrkC) [27], but is able to act on a specific subgroup of neurons. NT-3 promotes neurite outgrowth of both neural placode derived nodose ganglion and paravertebral chain sympathetic ganglia suggesting a broader specificity than either NGF or BDNF [59]. Additionally, NT-3 has been shown to support specifically neurons within the trigeminal ganglion early in development and neurons within the superior cervical sympathetic ganglia early in development [64]. Promotion of growth within the trigeminal ganglion is of particular interest for craniofacial nerve injuries, given the widespread distribution of the trigeminal nerve and its branches. In chick embryos, continuous administration of NT-3 in vivo supported 36% of motor neurons that are typically lost during early embryonic development between day 6 and day 10 [64].

4.4. Vascular endothelial growth factor

Other than the three major neurotrophins responsible for neuronal cell survival, differentiation and apoptosis, there are growth factors that can effectively promote neuronal regeneration. One such growth factor is vascular endothelial growth factor (VEGF). Naturally, VEGF promotes angiogenesis which is a crucial step towards tissue growth and repair [65]. Increased vascularization promoted by VEGF has been shown to also promote regeneration of nerve fibers, axonal outgrowth from the DRG and SCG (superior cervical ganglia) and even neuronal survival in vitro [66,67].

Specifically, VEGF secretes a 165 amino acid binding variant that is seen to bind to neuropilin-1, a neuronal cell surface molecule on neurons, which plays a crucial role in chemorepulsive signaling during axonal outgrowth [66]. In studies conducted by Sondell et al., when VEGF was added to nerve grafts, Schwann cell invasion and neo-vascularization were promoted, both of which are important processes of the nerve regeneration process [65,67]. These treated nerves were seen to have increased tissue organization, vascularization, angiogenesis and myelinated nerve fibers [67]. Additionally, VEGF causes activation of the flk-1 receptor, the MAPK pathway, and a receptor on Schwann cells, which can stimulate axonal outgrowth from the DRG and SCG [67].

5. Utilization of electrical stimulation

Cellular differentiation can be guided by either growth factors that trigger cell signaling pathways or by external physical stimulus. Physical stimulus such as stretch, compression routines [68] or nanotopologies [69] have shown effective outcomes for musculoskeletal and nerve tissue engineering, respectively. These physical stimulus are sensed by the mechanosensors of the cell and transduced to initiate cell signaling cascade. Likewise, application of external electrical stimulation using suitable medium such as conducting polymers has been shown to affect cellular behavior due to the changes in ion influx across the cellular membrane, which propagates intracellular transduction pathways [70]. Generally, the cell membrane maintains a steady state potential, called resting potential. Small electrical activity called action potential can alter the transmembrane potential from negative to positive. Information is transferred in neurons along the axons by means of series of action potentials that elicits a growth-controlling transport processes across the plasma membrane [71]. Severed neurons must be able to switch from a transmitting mode to a growth mode in order to express growth-associated proteins such as GAP-43, tubulin, and actin [72]. As shown in Elzinga et al., electrical stimulation promotes axonal outgrowth through promotion of intracellular cAMP, increased expression of neurotrophic factors (including glial cell line-derived neurotrophic factor (GDNF) and brain-derived neurotrophic factor (BDNF)) [73] along with their receptors (tyrosine kinase B (Trk B)), and an increased expression of genes associated with nerve regeneration such as actin, tubulin, galectin-1, growth associated protein-43 (GAP-43), and neurotrophin-4/5 [73-75].

Electrical stimulation (ES) has shown to have a wide range of positive effects on different tissues. Many of these effects have been observed and studied in vivo as well as in vitro. Utilization of electrical stimulation has shown an increase in nerve regeneration and decreased atrophy of axotomized nerves [70,76]. The effects of ES on neurons and within peripheral nerves following injury has been well documented [77]. ES accelerates both sensory and motor nerve regeneration and healing after an injury [77]. When ES is performed within 30 days following injury, it increases myelin formation at the location of the damaged nerve, prevents Schwann cell apoptosis and promotes their activity—crucial both for myelin production and nerve regeneration [77]. ES also increases the expression of BDNF, its receptor TrkB activation, and Growth Associated Protein 43 (GAP-43) within nervous tissue, promoting regeneration. Additionally, ES promotes muscle growth, regeneration and remodeling following an injury and can assist in preventing disuse atrophy following lower motor neuron (LMN) injury [78].

Interestingly, ES also increases intra-muscular levels of BDNF and glial cell derived neurotrophic factor (GDNF) mRNA [79]. This has been shown to promote neuronal regeneration in surrounding nervous tissue and enhanced synaptic connections at the neuromuscular junction [79]. A proposed mechanism for this phenomenon (intra-muscular neuronal growth factors influencing extra-cellular neurons) is that diffusion of BDNF and GDNF out of the muscle fibers and into the surrounding damaged neurons promotes neuronal regeneration.

ES also increases levels of VEGF mRNA and VEGF protein [80,81]. As discussed previously, VEGF promote angiogenesis in vivo and endothelial cell proliferation in vitro [80]. Angiogenesis is critical for both tissue regeneration and remodeling [81]. It is hypothesized that the metabolic imbalance created by ES results in increased VEGF to promote angiogenesis. ES causes continuous muscle contraction which requires nutrients and oxygen (O2). O2 is perhaps the limiting factor of all nutrients required by muscle and VEGF functions to increase O2 delivery to hypoxic tissues [81]. ES helps prevent disuse atrophy by conserving type-1 twitch muscle fibers [78]. This is perhaps indirectly due to the increased VEGF and angiogenesis allowing for the increased O2 delivery which is utilized by type-1 muscle fibers via the electron transport chain for ATP synthesis and subsequent muscle contraction. ES also increases proliferation and differentiation of myoblasts into myocytes which is due to an increase in expression of genes associated with myogenic differentiation [82-86].

In order for electrical stimulation to elicit electrical surge through the scaffold, the nerve conduit needs to be made of electrically conductive polymers. Electrical stimulation is seen to promote cell proliferation and growth in vitro using minimum amount of electrical stimulation, time, or voltage threshold in order to initiate a cellular cascade [35,87].

For in vitro experiments, one common method to deliver DC electrical stimulation to cells within a culture plate is through the use of salt bridges immersed within the cells culture media [88]. This allows the salt bridges to separate the cells from the electrodes in order to prevent any fluctuations in pH or the formation of chemical byproducts [88,89]. Although this method is useful, there are several drawbacks to using this type of chamber setup: the wells and working area are small which limit the number of cells that can be simultaneously electrically stimulated, the amount of time that the cells can be exposed to the electrical stimulation is limited because of the concentration and heat difference between the salt bridge and the culture media, and there are issue with maintaining sterile conditions [88]. Studies using carbon and metallic electrodes have also been used, however, these studies used single petri dishes, which reduced the amount of cells that could be studied at once, increased the time for the experiments, and reduced replication of the experiments [88,90]. In order to overcome all the described problems, a new plate design was created consisting of platinum electrodes secured to the lid of a well culture plate. This design uses a 1 mm gauge platinum wire, 50 mm in length, and bent into an L.
shape. By attaching the electrodes to the lid of the culture plate, the ES design allows for easy handling, negligible media evaporation, and easy sterilization. The individual electrode tips are soldered to copper wires coated in silver run in a parallel circuit [88]. The silver-coated wires are then attached to a DC power supplier, which can change the voltage, duration, and frequency. Table 1 provides a list of commercially available machines that are used for electrical stimulation and compares their mode of action.

ES has shown to work therapeutically in vivo and mechanistically in vitro. Although literature is lacking regarding the precise mechanisms on a cellular level through which ES manipulates tissues, the clinical argument is strong supporting the benefits of ES in a wide variety of tissues [78,91,92]. ES could become the future standard of care but more research is needed to define the parameters necessary to maximize its benefits.

The parameters used in ES studies in the literature vary greatly both from tissue to tissue, and also within each group. The parameters vary in terms of voltage, frequency, time of stimulation, and delay after injury before ES is used [78,80,91]. More studies and clinical trials are needed to standardize parameters for voltage, frequency, and time of stimulation to achieve maximum benefits of ES.

6. Conclusion

The complex anatomy and physiology of the craniofacial region make incidents that damage nervous tissue very difficult to repair and regenerate. Surgeries utilizing suturing, grafts, and conduits are present, but pose several limitations when trying to recuperate facial tissue. ES has shown to work therapeutically in vivo. Although literature is lacking regarding the precise mechanisms in vivo, the clinical and experimental therapies, BioMed Res. Int. 2014 (2014) 698256.

Table 1

| Machine | Voltage Range | Pulse Length | Electrodes | Reference |
|---------|---------------|--------------|------------|-----------|
| Transcutaneous Electrical Nerve Stimulation (TENS) | 0–350 V | 1–250 μs | Electrode pads | [93] |
| The Jouan PS10/15 Electropulsator | Variable | Variable | Parallel Plates | [94] |
| BTX ECM 830 | 0–500 V | 5 μs–24 msec | Parallel Plates or electrode needles | |
| 20–3000 V | 0.3 μs–99 msec | | |
| Variable | 5 μs–99 μs | | |

Acknowledgements

The authors acknowledge funding support from the National Institute of Biomedical Imaging and Bioengineering of the National Institutes of Health (R01EB020640), the Connecticut Regenerative Medicine Research Fund (15-RMBUCHC-08), and the Department of Defense (ORI21040).

References

[1] R. Tevlin, A. McArdle, D. Athanou, et al., Biomaterials for craniofacial bone engineering, J. Dent. Res. 93 (2014) 1187–1195.
[2] R. Birch, The Peripheral Nervous System: Gross Anatomy, (2010), pp. 1–41.
[3] D. Grinsell, C.P. Keating, Peripheral nerve reconstruction after injury: a review of clinical and experimental therapies, BioMed Res. Int. 2014 (2014) 698256.
[4] M.G. Burnett, E.L. Zager, Pathophysiology of peripheral nerve injury: a brief review, Neurosurg. Focus 16 (2004) 1–7.
[5] E.A. Huerbner, S.M. Strittmatter, Axon Regeneration in the Peripheral and central nervous systems, (2009), pp. 305–360.
[6] J. Inoue, Treatment of acute peripheral nerve injuries: current concepts, J. Hand Surg. 35 (2010) 491–497.
[7] M.F. Meek, J.H. Goert, US Food and Drug Administration/Conform Europe-approved nerve conduits for clinical repair of peripheral and cranial nerves, Ann. Plast. Surg. 60 (2008) 110–116.
[8] A.J. Reid, A.C. de Luca, A. Faroni, S. Downes, M. Sun, G. Terenghi, P.J. Kingham, Long term peripheral nerve regeneration using a novel PCL nerve conduit, Neurosci. Lett. 440 (2008) 125–128.
[9] C.M. Agrawal, R.B. Ray, Biodegradable polymeric scaffolds for musculoskeletal tissue engineering, J. Biomed. Mater. Res.: Off. J. Soc. Biomater. 55 (2001) 141–150. The Japanese Society for Biomaterials and The Australian Society for Biomaterials and the Korean Society for Biomaterials.
[10] R. James, O.S. Manoukian, S.G. Kumbar, Poly (lactic acid) for delivery of bioactive macromolecules, Adv. Drug Deliv. Rev. 107 (2016) 277–288.
[11] Y. Gu, J. Zhu, C. Xue, Z. Li, P. Ding, Y. Yang, X. Gu, Chitosan/silk fibroin-based, Schwann cell-derived extracellular matrix-modified scaffolds for bridging rat sciatic nerve gaps, Biomaterials 35 (2014) 2253–2263.
[12] M. Farokhi, F. Mottaghitalab, M.A. Shokrgozar, D.L. Kaplan, H. Kim, S.C. Kudou, Prospects of peripheral nerve tissue engineering using nerve guide conduits based on silk fibroin protein and other biopolymers, Int. Mater. Rev. 62 (2017) 367–391.
[13] A. Gloria, R. De Santis, A. Ambrosio, Polymer-based composite scaffolds for tissue engineering, J. Appl. Biomater. Biomech. 8 (2010) 57–67.
[14] S.I. Jeong, N.A. Burns, C.A. Bonico, I.K. Kwon, S.A. Khan, E. Alisberg, Improved cell infiltration of highly porous 3D nanofibrous scaffolds formed by combined fiber–fiber charge repulsions and ultra-sonication, J. Mater. Chem. B 2 (2014) 8116–8122.
[15] S. Rose, M. Roy, A. Bandypadhyay, Recent advances in bone tissue engineering scaffolds, Trends Biotechnol. 30 (2012) 546–554.
[16] M.J. Otsuka, X. Cheng, S.S. Lee, R. Kumar, Y. Kim, M.J. Kaufman, E.P. Douglas, L.B. Gover, Bone structure and formation: a new perspective, Mater. Sci. Eng. R 58 (2007) 77–116.
[17] S. Arameshbad, J. Johnston, M. Tanzer, D. Pasini, Fully porous 3D printed titanium femoral stem to reduce stress-shielding following total hip arthroplasty, J. Orthop. Res. 35 (2017) 1774–1783.
[18] W. Daly, L. Yao, D. Zeghibis, A. Windebank, A. Pandit, A biomaterials approach to peripheral nerve regeneration: bridging the peripheral nerve gap and enhancing functional recovery, J. R. Soc. Interface 9 (2012) 202–221.
[19] A.B. Nectow, K.G. Maass, D.L. Kaplan, Development of the peripheral nerve guidance conduits, Tissue Eng. B 18 (2011) 104–117.
[20] G. Li, Y. Kong, Y. Zhao, Y. Zhao, L. Zhang, Y. Yang, Fabrication and characterization of polycrylamide/silk fibroin hydrogels for peripheral nerve regeneration, J. Biomater. Sci. Polym. Ed. 26 (2015) 899–916.
[21] M.I. Hobson, C.J. Green, G. Terenghi, VEGF enhances intraneural angiogenesis and improves nerve regeneration after axotomy, J. Anat. 197 (2000) 591–605.
[22] P. Aebischer, P. Saliegos, I. Wizn, Basic Fibroblast growth factor released from Schwann cell-derived extracellular matrix mediates nerve regeneration across large nerve gaps, J. Neurosci. Res. 23 (1989) 282–289.
[23] M. Kanje, A. Skottner, J. Sjo, Insulin-like growth factor I (IGF-I) stimulates regeneration of the rat sciatic nerve, Brain Res. 486 (1989) 396–398.
[24] N. Hamada, Y. Fujita, T. Kojima, A. Kitamoto, Y. Akao, Y. Nozawa, M. Ito, MicroRNA expression profiling of NGF-treated PC12 cells revealed a critical role for miR-211 in neuronal differentiation, Neurochem. Int. 60 (2012) 743–755.
[25] F. Fiores, P. Ferreira, C.A. Rodrigues, J. Morgado, F.C. Ferreira, Neural stem cell differentiation by electrical stimulation using a cross-linked PEDOT substrate: expanding the use of biocompatible conducting polymers for neural tissue engineering, Biochim. Biophys. Acta Gen. Subj. 1850 (2015) 1158–1168.
[26] A.W. English, G. Schwartz, W. Meador, M.J. Sabatier, A. Mulligan, Electrical stimulation promotes peripheral axon regeneration by enhanced neuronal neurotrophin signaling, Dev. Neurobiol. 67 (2007) 158–172.
[27] D. Purves, R. Cabeza, S.A. Huettel, K.S. LaBar, M.L. Platt, M.G. Woldoff, E.M. Brannon, Cognitive Neuroscience, (2008).
[28] A.D. Gaudet, P.G. Popovich, M.S. Ramey, Wallerian degeneration: gaining perspective on inflammatory events after peripheral nerve injury, J. Neuroinflammation 8 (2011) 110.
[29] K.R. Jensen, R. Minsky, C.L. Lloyd, Schwann cells: development and role in nerve repair, Cold Spring Harb. Perspect. Biol. 7 (2015), p. a020487.
[30] P. Caroni, M.E. Schwab, Antibody against myelin associated inhibitor of neurite growth neutralizes nonpermissive substrate properties of CNS white matter, Neuron 1 (1988) 85–96.
[31] S. Rotteniker, Wallerian degeneration: the innate-immune response to traumatic nerve injury, J. Neuroinflammation 8 (2011) 109.
[32] A. Mabasozi, A. Faroni, B.M. Mignon, S. Downes, G. Terenghi, A.J. Reid, Polymer scaffolds with preferential parallel grooves enhance nerve regeneration, Tissue Eng.
neurotrophic factor which stimulates axonal outgrowth through the flk-1 receptor, Eur. J. Neurosci. 12 (2000) 4243–4254.

M. Sondell, Vascular Endothelial Growth Factor (VEGF) and Peripheral Nerve Regeneration, (2000).

S. Bhatnagar, R. Basu, Vascular Endothelial Growth Factor (VEGF) in Peripheral Nerve Repair and Regeneration, J. Neurosci. Res. 64 (2001) 270–278.

S. Bhatnagar, L. Escobar, P. Saha, Vascular endothelial growth factor is angiogenic in the peripheral nervous system, J. Peripher. Nerv. Syst. 22 (2017) 252–259.

F. Yang, Y. Muraoka, Y. Kojima, Y. Kita, B. Kita, Role of vascular endothelial growth factor in peripheral nerve regeneration, Neurosci. Lett. 645 (2016) 135–140.

M. Sondell, Vascular Endothelial Growth Factor (VEGF) and Peripheral Nerve Regeneration, (2000).

S. Bhatnagar, R. Basu, Vascular Endothelial Growth Factor (VEGF) in Peripheral Nerve Repair and Regeneration, J. Neurosci. Res. 64 (2001) 270–278.

S. Bhatnagar, L. Escobar, P. Saha, Vascular endothelial growth factor is angiogenic in the peripheral nervous system, J. Peripher. Nerv. Syst. 22 (2017) 252–259.