

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
8 July 2010 (08.07.2010)

(10) International Publication Number
WO 2010/078342 A2

(51) International Patent Classification:

<i>A61M 1/00</i> (2006.01)	<i>A61L 27/58</i> (2006.01)
<i>A61M 27/00</i> (2006.01)	<i>A61L 27/24</i> (2006.01)
<i>A61M 37/00</i> (2006.01)	<i>A61L 27/54</i> (2006.01)
<i>A61M 31/00</i> (2006.01)	<i>A61L 27/56</i> (2006.01)

(21) International Application Number:

PCT/US2009/069710

(22) International Filing Date:

29 December 2009 (29.12.2009)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

61/142,065	31 December 2008 (31.12.2008)	US
61/142,053	31 December 2008 (31.12.2008)	US
61/234,692	18 August 2009 (18.08.2009)	US
61/238,770	1 September 2009 (01.09.2009)	US

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(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PE, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

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(54) Title: SYSTEM FOR PROVIDING FLUID FLOW TO NERVE TISSUES

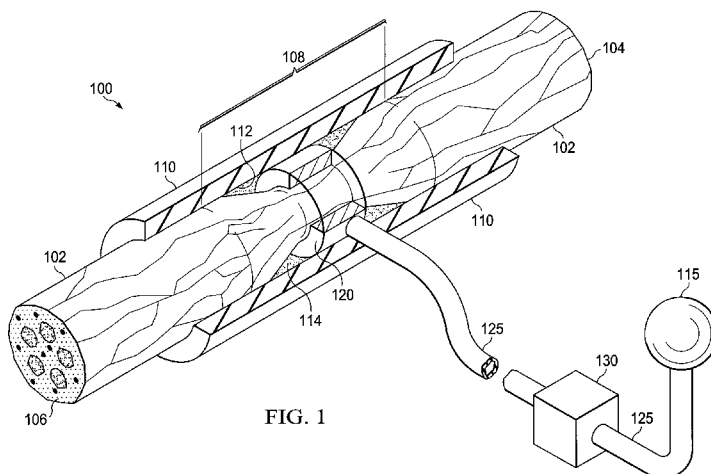


FIG. 1

(57) Abstract: An apparatus and system for providing reduced pressure to a defect in a nerve is disclosed. The apparatus comprises a nerve conduit having a generally tubular shape that has walls including a luminal wall surrounding the tissue site to contain fluids within a luminal space between the tissue site and the luminal wall. The apparatus further comprises a manifold having a porous body and a connector for receiving reduced pressure, wherein the manifold is positioned within the luminal space adjacent the tissue site to distribute the reduced pressure to the defect. Additionally, a method for providing reduced pressure to a defect in a nerve is disclosed that includes implanting the nerve conduit and manifold at a site of damaged nerve tissue and applying a reduced pressure to the manifold thereby stimulating repair or regrowth of nerve tissue.



WO 2010/078342 A2

Published:

— *without international search report and to be republished upon receipt of that report (Rule 48.2(g))*

SYSTEM FOR PROVIDING FLUID FLOW TO NERVE TISSUES

CROSS-REFERENCED TO RELATED APPLICATIONS

5 [0001] This application claims priority to U.S. Provisional Application Nos. 61/142,053 and 61/142,065, each filed on December 31, 2008. This application also claims priority to U.S. Provisional Application No. 61/234,692, filed on August 18, 2009 and U.S. Provisional Application No. 61/238,770, filed on September 1, 2009. Each of the foregoing applications is incorporated herein by reference in their entirety.

10

BACKGROUND

1. Field of the Invention

[0002] The present application relates generally to tissue engineering and in particular to apparatuses and systems suitable for use in treatment of damaged nerve tissue.

2. Description of Related Art

15 [0003] Clinical studies and practice have shown that providing a reduced pressure in proximity to a tissue site augments and accelerates the growth of new tissue at the tissue site. The applications of this phenomenon are numerous, but application of reduced pressure has been particularly successful in treating wounds. This treatment (frequently referred to in the medical community as “negative pressure wound therapy,” “reduced pressure therapy,” or
20 “vacuum therapy”) provides a number of benefits, including faster healing and increased formation of granulation tissue. Typically, reduced pressure has been applied to tissue through a porous pad or other manifold device. The porous pad contains pores that are capable of distributing reduced pressure to the tissue and channeling fluids that are drawn from the tissue. The porous pad often is incorporated into a dressing having other components that facilitate
25 treatment. A scaffold can also be placed into a defect to support tissue growth into the defect. The scaffold is usually bioabsorbable, leaving new tissue in its place.

[0004] Scaffolds for reduced pressure treatment are described in, *e.g.*, WO08/091521, WO07/092397, WO07/196590, WO07/106594. The adequacy of current scaffolds for reduced pressure treatment can be evaluated in light of current knowledge of wound healing. Injury to
30 body tissues results in a wound healing response with sequential stages of healing that include hemostasis (seconds to hours), inflammation (hours to days), repair (days to weeks), and remodeling (weeks to months). A high level of homology exists across most tissue types with regards to the early phases of the wound healing process. However, the stages of healing for various tissues begin to diverge as time passes, with the involvement of different types of

growth factors, cytokines, and cells. The later stages of the wound healing response are dependent upon the previous stages, with increasing complexity in the temporal patterning of and interrelationships between each component of the response.

5 [0005] Strategies to facilitate normal repair, regeneration, and restoration of function for damaged tissues have focused on methods to support and augment particular steps within this healing response, especially the latter aspects of it. To this end, growth factors, cytokines, extracellular matrix (ECM) analogs, exogenous cells and various scaffolding technologies have been applied alone or in combination with one another. Although some level of success has been achieved using this approach, several key challenges remain. One main challenge is
10 that the timing and coordinated influence of each cytokine and growth factor within the wound healing response complicate the ability to add individual exogenous factors at the proper time and in the correct coordination pattern. The introduction of exogenous cells also faces additional complications due to their potential immunogenicity as well as difficulties in maintaining cell viability.

15 [0006] Synthetic and biologic scaffolds have been utilized to provide three-dimensional frameworks for augmenting endogenous cell attachment, migration, and colonization. To date nearly all scaffolds have been designed with the idea that they can be made to work with the biology. Traditional scaffolding technologies, however, rely on the passive influx of endogenous proteins, cytokines, growth factors, and cells into the interstitium of the porous
20 scaffold. As such, the colonization of endogenous cells into the scaffold is limited by the distance away from vascular elements, which provide nutrient support within a diffusion limit of the scaffold, regardless of tissue type. In addition, the scaffolds can also elicit an immunogenic or foreign body response that leads to an elongated repair process and formation of a fibrous capsule around the implant. Taken together, these complications can all lead to
25 less than functional tissue regeneration at the injury site.

[0007] It would therefore be advantageous to provide additional systems for the repair and remodeling of specialized tissues. The present invention provides such systems.

SUMMARY

[0008] The apparatuses, systems and methods of the illustrative embodiments described herein provide active guidance of nerve tissue repair and regeneration through an implanted manifold and nerve conduit. In one embodiment, an apparatus for providing reduced pressure therapy and facilitating growth of nerve tissue in a patient is provided that includes a nerve conduit and nested manifold adaptable for implantation at a damaged nerve site, wherein the manifold provides and distributes reduced pressure at the site of damaged nerve tissue. A manifold according to the invention may also be coupled to a scaffold material which further distributes reduced pressure and provides a structural matrix for growth of the tissue.

[0009] In another embodiment, a system for providing reduced pressure therapy and facilitating growth of nerve tissue in a patient is provided that comprises a source of reduced pressure for supplying reduced pressure and a manifold nested in a nerve conduit adaptable for implantation at the tissue site, where the manifold is in fluid communication with the source of reduced pressure. Such a system may also comprise a scaffold material coupled to the manifold which further distributes reduced pressure and provides a structural matrix for the growth of nerve tissue. In a further embodiment, such a system may further comprise a canister for fluid capture and/or a valve for control of reduced pressure in fluid communication with, and positioned between, the manifold and the reduced pressure source. In still a further embodiment, a system according to the invention further comprises a fluid source in fluid communication with the manifold and the damaged nerve tissue.

[0010] In a further embodiment, a method of providing reduced pressure therapy and facilitating growth of nerve tissue at a site of nerve tissue damage in a patient is provided that includes implanting a nerve conduit and nested manifold at the tissue site, where the manifold provides a reduced pressure to the damaged nerve tissue. The manifold may also be coupled to a scaffold material, wherein the scaffold material provides a structural matrix for the growth of the nerve tissue. In certain embodiments, the method further comprises providing a manifold in fluid communication with a fluid source, wherein the fluid source may be used to deliver a fluid to the manifold and the damaged nerve tissue. In yet a further embodiment, the fluid source may comprise a fluid comprising one or more bioactive compounds including, but not limited to, an antibiotic, an antiviral, a cytokine, a chemokine, an antibody, and a growth factor.

[0011] Other objects, features, and advantages of the illustrative embodiments will become apparent with reference to the drawings and detailed description that follow.

BRIEF DESCRIPTION OF THE DRAWINGS

[0012] FIG. 1 is a schematic, perspective view of a first reduced-pressure system for repairing a pinched nerve including a nerve conduit and a first embodiment of a manifold with a section of the nerve conduit removed to show the manifold;

5 [0013] FIG. 2 is a schematic, perspective view of a second reduced-pressure system for repairing a severed nerve including a nerve conduit and a second embodiment of a manifold with a section of the nerve conduit removed to show the manifold;

[0014] FIG. 3 is a schematic, perspective view of a third embodiment of a manifold for use in the systems shown in FIGS. 1 and 2;

10 [0015] FIG. 4 is a schematic, perspective view of the system shown in FIGS. 1 and 2 showing the nerve conduit enclosing the damaged nerve; and

[0016] FIG. 5 is a schematic view of a fluid control system for the system shown in FIGS. 1 and 2.

15

DETAILED DESCRIPTION

[0017] In the following detailed description of the illustrative embodiments, reference is made to the accompanying drawings that form a part hereof. These embodiments are described in sufficient detail to enable those skilled in the art to practice the invention, and it is understood that other embodiments may be utilized and that logical structural, mechanical, electrical, and chemical changes may be made without departing from the spirit or scope of the invention. To avoid detail not necessary to enable those skilled in the art to practice the embodiments described herein, the description may omit certain information known to those skilled in the art. The following detailed description is, therefore, not to be taken in a limiting sense, and the scope of the illustrative embodiments are defined only by the appended claims.

20 [0018] Referring to FIG. 1, a reduced pressure therapy system 100 for applying reduced pressure at a tissue site in the body of a patient to repair a defect such as, for example, a damaged nerve is disclosed. The damaged nerve may have been pinched, partially disconnected or severed, or partially degenerated as a result of disease. For example, the damaged nerve in FIG. 1 is a pinched nerve 102 having a proximal segment 104 and a distal segment 106 relative to the central nervous system (CNS) of the patient. The pinched nerve 102 has been damaged at a nerve damage site 108 that has been pinched or partially degenerated, but has not been completely severed. The pinched nerve 102 may be branched or unbranched at the nerve damage site 108. The term "nerve damage site" as used herein refers to a wound or defect located on or within any nerve tissue, including, but not limited to, a

disconnected or partially disconnected nerve, a degenerated or partially degenerated nerve, and a compressed or pinched nerve. For example, reduced pressure tissue treatment may be used to enhance repair or regrowth of existing nerve tissue or to facilitate growth or grafted or transplanted nerve tissue and/or cells.

5 [0019] The reduced pressure therapy system 100 comprises a nerve conduit 110 that surrounds the pinched nerve 102 at the nerve damage site 108 with a section of the nerve conduit 110 removed to show the nerve damage site 108. The nerve conduit 110 is substantially tubular in shape and closes the nerve damage site 108 and a portion of the proximal segment 104 and the distal segment 106 that has not been damaged. The nerve conduit 110 has an inner surface
10 112 that forms a nerve gap 114 with the surface of the nerve damage site 108, i.e., a luminal space between the inner surface 112 of the nerve conduit 110 and the surface of the nerve damage site 108. The reduced pressure therapy system 100 also comprises a reduced pressure source 115 for providing a reduced pressure and a manifold 120 fluidly coupled to the pressure source 115 via a first conduit 125. The manifold 120 is generally tubular in shape and
15 positioned within the nerve gap 114 surrounding the surface of the nerve damage site 108. The manifold 120 may have a variety of shapes depending on the type of nerve damage, and in this particular embodiment has a tubular shape to occupy a portion of the nerve gap 114 around the nerve damage site 108. The manifold 120 may also be coupled to a scaffold material or contain scaffold material (not shown) that provides a structure for tissue growth
20 and repair. The reduced pressure therapy system 100 further comprises a canister 130 fluidly coupled between the reduced pressure source 115 and the manifold 120 via the conduit 125 to collect bodily fluids such as blood or exudate that are drawn from the nerve gap 114. In one embodiment, the reduced pressure source 114 and the canister 130 are integrated into a single housing structure.

25 [0020] As used herein, the term “coupled” includes direct coupling or indirect coupling via a separate object. The term “coupled” also encompasses two or more components that are continuous with one another by virtue of each of the components formed from the same piece of material. Also, the term “coupled” may include chemical, mechanical, thermal, or electrical coupling. Fluid coupling means that fluid is in communication with designated parts or
30 locations.

[0021] In the context of this specification, the term “reduced pressure” generally refers to a pressure that is less than the ambient pressure at a tissue site that is subjected to treatment. In most cases, this reduced pressure will be less than the atmospheric pressure of the location at which the patient is located. Although the terms “vacuum” and “negative pressure” may be

used to describe the pressure applied to the tissue site, the actual pressure applied to the tissue site may be significantly greater than the pressure normally associated with a complete vacuum. Consistent with this nomenclature, an increase in reduced pressure or vacuum pressure refers to a relative reduction of absolute pressure, while a decrease in reduced pressure or vacuum pressure refers to a relative increase of absolute pressure. The term “ $-\Delta p$ ” means change in reduced pressure. As used herein, a greater (*i.e.*, more negative) $-\Delta p$ means increased negative pressure (*i.e.*, a greater change in pressure from ambient pressure). Reduced pressure treatment typically applies reduced pressure at -5 mm Hg to -500 mm Hg, more usually -5 to -300 mm Hg, including but not limited to -50, -125 or -175 mm Hg.

Reduced pressure may be constant at a particular pressure level or may be varied over time. For example, reduced pressure may be applied and stopped periodically or ramped-up or -down over time.

[0022] As indicated above, a nerve damage site may be a wound or defect located on or within any nerve tissue including, for example, a completely severed nerve 202 having a proximal segment 204 and a distal segment 206 relative to the CNS of the patient as shown in FIG. 2. The severed nerve 202 has been damaged at a nerve damage site 208 that has been completely severed. Another reduced pressure therapy system 200 for applying reduced pressure at the nerve damaged site 208 comprises similar components as the reduced pressure therapy system 100. The reduced pressure therapy system 200 comprises the nerve conduit 110 that surrounds the nerve damage site 208 and the severed ends of the severed nerve 202. The inner surface 112 of the nerve conduit 110 forms a nerve gap 214 between the severed ends of the severed nerve 202 within the nerve damage site 208, *i.e.*, a luminal space between the inner surface 112 of the nerve conduit 110 and the surface of the nerve damage site 208. The reduced pressure therapy system 200 also comprises a manifold 220 fluidly coupled to the pressure source 115 via the first conduit 125 and positioned within the nerve gap 214 leaving nerve gaps 214a and 214b between each of the severed ends of the severed nerve 202. The manifold 220 may be generally tubular in shape to enhance fluid flow within the nerve gap 214. The manifold 220 may also be generally cylindrical in shape and function additionally as a scaffold material (not shown) that provides a structure for tissue growth and repair between the severed ends of the severed nerve 202.

[0023] The manifolds 120, 220 shown in FIGS. 1 and 2 create flow patterns and gradients in the nerve gaps 114 and 214 within the nerve damage sites 108 and 208, respectively, which can be further modified to enhance nerve tissue growth. Referring to FIG. 3, a manifold 320 having a core portion 322 with a generally cylindrical shape similar to the other manifolds

120, 220 is shown. The manifold 320 may be used for severed nerve damage, e.g., nerve damage site 208. The core portion 322 of the manifold 320 may also include an axial opening 323 extending between the end surfaces of the core portion 322 (one end surface 324 shown) to accommodate a pinched nerve, e.g., nerve damage site 108. The manifold 320 further
5 comprises manifold rods 325, each one having one end in fluid communication with the end surfaces of the core portion 322 of the manifold 320 and the other end extending axially therefrom within the nerve gaps 314a and 314b at the nerve damage site 208. Any number of manifold rods 325 may be used as required for the re-growth nerve tissue. Furthermore, the manifold 320 may include manifold rods 325 extending from only one of its end surfaces (not
10 shown) so that the core portion 322 may be positioned adjacent the distal or proximal end of the damaged nerve site 208. The manifold rods 325 may be generally tubular in shape and formed of the same material as the other manifolds 120, 220 as described above so that they are in fluid communication with the nerve gaps 114 and 214 to promote the formation of new nerve tissue. Hence, the manifold rods 325 may also be bioabsorbable so that there is no need
15 to remove them after new nerve tissue begins forming in the nerve gaps 114 and 214. Alternatively, the manifold rod 325 may comprise a conduit closed at one end and fluidly coupled to the core portion 322 of the manifold 320 at the other end. The walls of this manifold rod 325 may include holes or pores 326 through which the core portion 322 of the manifold 320 fluidly communicates within the nerve gaps 114 and 124.

20 **[0024]** The nerve conduit 110 is shown in FIGS. 1 and 2 with a section removed to show the manifolds 120, 220 and is shown as completely surrounding the nerve damage sites 108, 208 as a closed nerve conduit 310 in FIG. 4. After the manifolds 120, 220 are inserted in the nerve damage sites 108, 208, respectively, the nerve conduit 110 may be sealed by utilizing one or more stitches 415 or any other fastening device known in the art. The nerve conduit 110 may
25 be composed of a bioabsorbable or a bioinert material. Materials that may be used for nerve conduits include, without limitation, polylactic acid (PLA), polyglycolic acid (PGA), polylactide-co-glycolide (PLGA), polyvinylpyrrolidone, polycaprolactone, polycarbonates, polyfumarates, caprolactones, polyamides, polysaccharides (including alginates (*e.g.*, calcium alginate) and chitosan), hyaluronic acid, polyhydroxybutyrate, polyhydroxyvalerate,
30 polydioxanone, polyorthoesters, polyethylene glycols, poloxamers, polyphosphazenes, polyanhydrides, polyamino acids, polyacetals, polycyanoacrylates, polyurethanes, polyacrylates, ethylene-vinyl acetate polymers and other acyl substituted cellulose acetates and derivatives thereof, polystyrenes, polyvinyl chloride, polyvinyl fluoride, poly(vinylimidazole), chlorosulphonated polyolefins, polyethylene oxide, polyvinyl alcohol, Teflon[®], and nylon. In

certain aspects biological (*e.g.*, purified or recombinant) materials may be used form nerve conduits including but are not limited to, fibrin, fibronectin or collagen (*e.g.*, DURAMATRIX™).

5 [0025] A nerve conduit 110 may be an unbroken substantially tubular structure fitted across a gap between a proximal and distal nerve stump such as depicted in FIG. 4. Examples of such substantially tubular nerve conduits, also referred to as nerve guides, include without limitation NEURAGEN® and NEUROFLEX™ collagen conduits. A nerve conduit may also be formed from a wrap that is positioned around a disconnected or damaged (*e.g.*, pinched) nerve and sealed with a closure, such as a suture. Specific examples of wrap-type nerve
10 conduits include, without limitation, NEUROMEND™ and NEURAWRAP™ collagen conduits. In certain aspects, the nerve conduit is made of a material that and encloses the damaged nerve, so as to exclude infiltration of non-nerve cells such as glia. In some embodiments, the nerve conduit material is permeable, thereby allowing fluid and protein factors to diffuse through the conduit. For example, the pores in a nerve conduit may be
15 sufficiently small so as to exclude the entry of cells into the conduit lumen (*e.g.*, pores having an interior diameter or average interior diameter of between about 5 µm and 50 µm, 10 µm and 30 µm or 10 µm and 20 µm). Thus, when reduced pressure is applied to the interior of the conduit fluid and proteins may be drawn to the lumen of the conduit by the pressure gradient. The skilled artisan will recognize that the dimensions of the conduit may be adjusted for any
20 particular nerve application. Generally, the conduits comprise an internal diameter of less than about 6.0 mm, such as about 5, 4, 3, 2.5 or 2 mm.

[0026] Referring to FIG. 5, the reduced pressure therapy system 100 may further comprise a pressure sensor 140 operably connected to the first conduit 125 to measure the reduced pressure being applied to the manifolds 120, 220. The system 100 further includes a control
25 unit 145 electrically connected to the pressure sensor 140 and the reduced pressure source 115. The pressure sensor 140 measures the reduced pressure within the nerve gap 114, 214 and also may indicate whether the first conduit 125 is occluded with blood or other bodily fluids. The pressure sensor 140 also provides feedback to control unit 145 which regulates the reduced pressure therapy being applied by the reduced pressure source 115 through the first conduit
30 125 to the manifolds 120, 220. The reduced pressure therapy system 100 may also comprise a fluid supply 150 fluidly coupled to the first conduit 125 via a second conduit 152 and operatively connected to the control unit 145. The fluid supply 150 may be used to deliver growth and/or healing agents to the nerve damage site 108, 208 including, without limitation, an antibacterial agent, an antiviral agent, a cell-growth promotion agent, an irrigation fluid, or

other chemically active agents. The system 100 further comprises a first valve 154 positioned in the second conduit 152 to control the flow of fluid therethrough, and a second valve 156 positioned in the first conduit 125 between the reduced pressure supply 115 and the juncture between the first conduit 125 and the second conduit 152 to control the flow of reduced pressure. The control unit 145 is operatively connected to the first and second valves 154, 156 to control the delivery of reduced pressure and/or fluid from the fluid supply 150, respectively, to the manifolds 120, 220 as required by the particular therapy being administered to the patient. The fluid supply 150 may deliver the liquids as indicated above, but may also deliver air to the manifolds 120, 220 to promote healing and facilitate drainage at the site of the nerve damage site 108, 208. As used herein, the term “manifold” refers to a substance or structure that is provided to assist in directing reduced pressure to, delivering fluids to, or removing fluids from a tissue site. A manifold can include a plurality of flow channels or pathways that are interconnected to improve distribution of fluids provided to and removed from the area of tissue around the manifold. Examples of manifolds may include, without limitation, devices that have structural elements arranged to form flow channels, cellular foams such as open-cell foam, porous tissue collections, and liquids, gels and foams that include or cure to include flow channels. A detailed description of manifolds and their use according to the invention is provided below.

[0027] The term “scaffold” as used herein refers to a substance or structure applied to or in a wound or defect that provides a structural matrix for the growth of cells and/or the formation of tissue. A scaffold is often a three dimensional porous structure. The scaffold can be infused with, coated with, or comprised of cells, growth factors, extracellular matrix components, nutrients, integrins, or other substances to promote cell growth. A scaffold can take on characteristics of a manifold by directing flow through the matrix. A manifold can transmit flow to the scaffold and tissue; in the context of reduced pressure treatment, the manifold can be in fluid communication with the scaffold. A detailed description of scaffolds and their use according to the invention is provided below.

[0028] As such, the invention disclosed here discloses methods and apparatuses for controlling cellular-level based patterns of fluid flow that would allow for control of patterned protein organization at a microscopic, nanoscopic, or mesoscopic scale amenable to provide a structured manifold and, optionally, a scaffold material for cellular migration, differentiation, and like behavior necessary for functional regeneration of tissues. In comparison to the passive nature of the current state of the art with regards to tissue repair and regeneration, the methods, scaffolds, manifolds, flow sources and systems disclosed herein provide an active

mechanism by which to promote the endogenous deposition of proteins and organization of the provisional matrix with biochemical and physical cues to direct cellular colonization of a scaffold or tissue space. The present invention thus enhances current technology by exploiting the active force of directed fluid flow, providing a framework upon which to design manifolds and scaffolds based upon the need of the biology under the influence of fluid flow. Flow vectors and pathways are utilized to enhance protein deposition and cellular colonization. The systems provided herein are designed to promote establishment of a provisional matrix network with a seamless transition from the healthy tissue edges through a scaffold or tissue site to promote a functional tissue continuum.

10 [0029] Thus, the apparatuses, methods and systems disclosed herein provide a means for active guidance of tissue regeneration through an implanted scaffold or within a tissue site to promote functional recovery. This active guidance occurs through mechanisms of controlled fluid flow, which can be used to initiate or augment the early stages of the body's own natural healing process; a manifold can provide the active guidance necessary to create a controlled fluid flow. Specifically, the controlled flow vectors that the manifolds provide can be used to facilitate the directed influx of cells and proteins into a scaffold. Creation of specific flow pathways within a tissue site or scaffold can lead to patterned deposition of proteins, such as collagen and fibrin within the manifold, scaffold or tissue space. Biochemical cues from cytokines, growth factors, and cells bound within the provisional matrix can work in conjunction with the natural physical cues of the provisional matrix and extracellular matrix to guide the subsequent migration of endogenous cells during the repair stages of healing. These cues act as a form of track that emanates from the healthy tissues and passes through the scaffolding or tissue space to facilitate a continuous guidance pathway for organized tissue regeneration.

25 [0030] To that end, this disclosure provides unique manifold technologies designed for specific biological needs based upon principles of fluid flow. In certain aspects, the invention concerns a new approach to wound healing, flow (or gradient) activated tissue engineering. In rudimentary form, this approach involves a source or generator of flow that forms a gradient for controlled movement of either endogenous or exogenous fluids into, out of, or through a tissue space for the organized deposition of proteins and/or spatial concentration of cytokines and growth factors, with subsequent formation of a directionally oriented provisional matrix. The tissue space being defined here includes, but is not limited to, the region surrounding a site of nerve tissue damage.

[0031] Fluid flow into, through, or out of the nerve tissue space can be refined and directed through the inclusion of additional elements to the system including manifolds and/or scaffolds. The coordinated elements of the system are designed to create flow parameters, pathways, and patterns sufficiently detailed in scale as to be able to influence and direct the controlled adsorption of proteins, the organization of matrix, and organized colonization of specific cell types. Individual elements of the system are as follows.

[0032] Source or Generator of Flow. Flow is induced into, through, or out of the nerve tissue space by methods or apparatuses that introduce changes in mechanical, chemical, and/or electrical potentials. These generators of flow provide either a gradient or a change in potential from the site or reservoir of endogenous or exogenous fluids to the placement position of the flow generator or its extension element (*i.e.*, manifold or scaffold). In one embodiment, the source of flow comprises a source of reduced pressure. Systems and apparatuses according to the invention may also comprise valves or arrays of valves that control the application and amount of negative pressure applied to the manifold. In certain aspects, nerve conduits and/or manifolds described herein comprise a pressure sensor. Thus, in some embodiments, the amount of negative pressure applied by a source is regulated based on the amount of negative pressure that is sensed in the manifold or nerve conduit or at the site of tissue damage.

[0033] Manifold. The flow generators are the driving force for stimulating the flow of fluids. Manifolds are apparatuses for refining the pattern of flow between the source or generator of flow and the tissue space. The macroscale level of flow is refined by specialized manifolds utilized for directed localization to a single point or to a plurality of selectively positioned points for creating initiation sites for microscale flow pathways within the manifold/scaffold and, ultimately, the tissue space. The manifold may also serve as a conduit for the removal of fluids from and as an apparatus for the delivery of exogenous fluids to the tissue space.

[0034] A manifold generally refers to a physical substance or structure that serves to assist in applying and translating a mechanical, chemical, electrical or similar alterations into changes in the flow of a fluid, herein defined as the movement of liquids, gases, and other deformable substances such as proteins, cells, and other like moieties. As such, this physical device includes a single point or plurality of points for the egress or evacuation of pressure, fluids, and like substances capable of translating the movement of fluids in a scaffold, as defined above. This can include, but is not limited to, the introduction of exogenous factors such as cells and/or therapeutic moieties into the scaffold through the lumen or plurality of lumens present in the manifold. In addition, as used herein, a manifold includes a single point or

plurality of points for the ingress or introduction of fluid from the scaffold back towards the point source of flow.

5 [0035] Flow distributed by the manifold can direct the movement of endogenous proteins, growth factors, cytokines, and cells from their resident locations within the host to the tissue space or scaffold in an organized manner. The establishment of flow along these pathways leads to the deposition of proteins and provisional matrix that creates an interfacial endogenous network connecting the host to the scaffold. Extensions of this matrix can be established within the scaffold through selective positioning of the manifold flow initiation sites with flow promoting scaffolding designs. The organized protein deposition and
10 provisional matrix provide a biochemical and physical framework that stimulates the attachment and migration of cells along directed pathways throughout the scaffold and the tissue space. The resulting endogenous network of proteins, growth factors, and cells provides a foundation upon which subsequent phases of the body's own tissue repair and regeneration mechanisms can build.

15 [0036] When in place, the manifold works in conjunction with a flow generating source and a scaffold, if present. Flow generating sources include, but are not limited to generators of negative pressure; generators of positive pressure; and generators of osmotic flow. The flow gradient established in the manifold may be further refined through the scaffold, to deliver a flow gradient to the scaffold to optimize flow through the scaffold as needed for the particular
20 defect. Many of the embodiments disclosed herein are manifolds capable of translating changes in pressure and the like into controlled movement of fluids, optionally through a physical scaffold, for the purposes of directed tissue regeneration. These embodiments are generally specified for a particular application in the regeneration of specific tissues, but are not limited to a particular tissue therein.

25 [0037] In order to realize the goal of inducing flow for the purpose of tissue regeneration, alterations in the aforementioned mechanical, chemical, or electrical impetus must be translated from the singular gradient source toward a physical substrate or scaffold to elicit cellular-level changes in protein adsorption, matrix organization, cell migration, and other tissue regeneration-related behaviors. These alterations are multivariate in nature and can
30 include mechanical changes that elicit a physical change in pressure applied to the scaffold as applied to the site of the wound or desired site of tissue regeneration, chemical changes that elicit a gradient in protein and/or ion concentrations, which result in the creation of osmotic gradients capable of inducing flow, or electrical changes that create a gradient of current/ion exchange allowing for propagation of electrical signals from the point source. It is to be

understood, however, that the applicants are not bound by any particular mechanism through which gradients and fluid flow induce advantageous results in tissue repair or growth. In order to advantageously transmit these gradients to the tissue, a physical device is needed to direct the path of flow from its source to the scaffold or tissue site and vice versa.

5 **[0038]** In some embodiments, the manifold comprises a physical structure in close apposition to or within the contents of a scaffold and serves to propagate an alteration in a physical parameter, whether it be mechanical, chemical, electrical, or something similar in nature, for the means of directing these changes from its point source to the scaffolding material. The placement of this manifold with respect to its location with regard to that of the scaffold may
10 be of crucial importance for facilitating controlled and directed regeneration of specific tissue types. For example, in peripheral nerve where regeneration primarily occurs in a unidirectional manner from the proximal to distal nerve stumps, it may be important to place the manifold along the length of a nerve conduit more towards its distal end to help direct regeneration towards that end. However, it may also be important to not place the manifold at
15 the most distal aspect of the scaffold/conduit as soluble factors derived from the distal stump have been shown to be important for directing nerve regeneration towards its source.

[0039] Manifolds may be composed of a bioabsorbable or bioinert material. Examples include non-bioabsorbable materials such as medical grade silicone polymers, metals, polyvinylchloride (PVC), and polyurethane. Bioabsorbable polymers such as collagen,
20 polylactic acid (PLA), polyglycolic acid (PGA), polylactide-co-glycolide (PLGA), a polysaccharide, a hydroxyapatite, or a polyethylene glycol, or combinations thereof, can also be used. Some manifolds are also a mix of non-bioresorbable and bioresorbable materials. In general material used for a scaffold may also be used to compose a manifold and such materials are further detailed below. In certain aspects, manifold materials are structured to
25 include a high void fraction for improved bioabsorption properties.

[0040] Scaffold. Biologic and synthetic scaffolds are used in the field of tissue engineering to support protein adhesion and cellular ingrowth for tissue repair and regeneration. The current state of the art in scaffold technology relies upon the inherent characteristics of the surrounding tissue space for the adsorption of proteins and migration of cells. A scaffold for
30 use according to the invention is coupled to a manifold, provides physical guidance to direct the pathway of fluid flow in the tissue site, creating avenues for the movement and migration of adhesive proteins and cells, respectively, which are integral to the establishment of a provisional matrix in predetermined patterns of organization within the tissue space. The methods and apparatuses described for fluid flow-induced and gradient-induced generation of

tissues have direct implications into the design of the scaffolds. Within this context, scaffolds serve to refine the pathways of fluid flow within the tissue space to cellular level patterns from the fluid source to the point(s) of flow initiation within the manifold. A scaffold may embody characteristics of a manifold or be combined in conjunction with a manifold for refinement of the flow pathways within the tissue site. In certain aspects, a scaffold is a reticulated structure comprising high void fraction for improved bioabsorption properties.

[0041] Nonlimiting examples of suitable scaffold materials include extracellular matrix proteins such as fibrin, collagen or fibronectin, and synthetic or naturally occurring polymers, including bioabsorbable or non-bioabsorbable polymers, such as polylactic acid (PLA), polyglycolic acid (PGA), polylactide-co-glycolide (PLGA), polyvinylpyrrolidone, polycaprolactone, polycarbonates, polyfumarates, caprolactones, polyamides, polysaccharides (including alginates (*e.g.*, calcium alginate) and chitosan), hyaluronic acid, polyhydroxybutyrate, polyhydroxyvalerate, polydioxanone, polyethylene glycols, poloxamers, polyphosphazenes, polyanhydrides, polyamino acids, polyortho esters, polyacetals, polycyanoacrylates, polyurethanes, polyacrylates, ethylene-vinyl acetate polymers and other acyl substituted cellulose acetates and derivatives thereof, polystyrenes, polyvinyl chloride, polyvinyl fluoride, poly(vinylimidazole), chlorosulphonated polyolefins, polyethylene oxide, polyvinyl alcohol, Teflon[®], and nylon. The scaffold can also comprise ceramics such as hydroxyapatite, coralline apatite, calcium phosphate, calcium sulfate, calcium carbonate or other carbonates, bioglass, allografts, autografts, xenografts, decellularized tissues, or composites of any of the above. In particular embodiments, the scaffold comprises collagen, polylactic acid (PLA), polyglycolic acid (PGA), polylactide-co-glycolide (PLGA), a polyurethane, a polysaccharide, an hydroxyapatite, or a polytherylene glycol. Additionally, the scaffold can comprise combinations of any two, three or more materials, either in separate areas of the scaffold, or combined noncovalently, or covalently (*e.g.*, copolymers such as a polyethylene oxide-polypropylene glycol block copolymers, or terpolymers), or combinations thereof. Suitable matrix materials are discussed in, for example, Ma and Elisseeff, 2005, and Saltzman, 2004.

[0042] Bioactive agents

[0043] In certain aspects, the apparatuses and methods according to the invention concern bioactive agents. Bioactive agents may, in some cases, be incorporated directly onto a manifold or scaffold material (*i.e.*, to generate a bioactive manifold and/or scaffold). For example, agents that facilitate tissue growth such as collagen or fibrin may be directly incorporated onto or into a manifold or scaffold material. Likewise, in applications where

aberrant immune response need be avoided (*e.g.*, tissue grafts) immune regulator agents such as rapamycin may be incorporated into manifold or scaffold structures.

[0044] In further aspects soluble bioactive agents may be introduced at a site of tissue damage by virtue of the flow through the tissue site. For example, a manifold may be in fluid

5 communication with a fluid source and a bioactive agent may be introduced into the fluid source and thereby into the manifold and damaged nerve tissue.

[0045] Nonlimiting examples of useful bioactive growth factors for various applications are growth hormone (GH), a bone morphogenetic protein (BMP), transforming growth factor- α (TGF- α), a TGF- β , a fibroblast growth factor (FGF), granulocyte-colony stimulating factor (G-
10 CSF), granulocyte/macrophage-colony stimulating factor (GM-CSF), epidermal growth factor (EGF), platelet derived growth factor (PDGF), insulin-like growth factor (IGF), vascular endothelial growth factor (VEGF), hepatocyte growth factor/scatter factor (HGF/SF), an interleukin, tumor necrosis factor- α (TNF- α) or nerve growth factor (NGF).

[0046] Nerve tissue repair and engineering. The apparatuses and systems disclosed herein can
15 be used for nerve tissue repair and engineering in various contexts including the following.

[0047] Repair and regeneration of lost tissue. A generator of flow may be combined with manifolds and/or scaffolds to direct the regeneration of lost tissue at a site of injury or
compromised function. Tissues lost from traumatic injury, surgery, burns, or other causes
(*e.g.*, infection or autoimmune disease) can be led to regenerate using the methods, scaffolds,
20 manifolds, flow sources and systems of the invention. Functional nerve tissue is directed to regenerate.

[0048] Retard the progression of a tissue disease state. A generator of flow may be combined
with manifolds and/or scaffolds to retard disease progression of an affected nerve tissue such
as occurs, *e.g.*, in autoimmune disease.

25 [0049] Maintenance of tissue viability. A generator of flow may be combined with manifolds and/or scaffolds to maintain the viability of explanted tissues either for *in vitro* study, *ex vivo* scaffold or implant preparation, or *in vivo* transplant. A generator of flow combined with a manifold may be used to provide nutrient fluid flow to the tissue and to control waste removal from the tissue.

30 [0050] Expansion of tissue. A generator of flow may be combined with manifolds and/or scaffolds to promote the expansion of existing tissues. The methods, scaffolds, manifolds, flow sources and systems of the invention can be used to direct the growth of tissues where additional tissue quantity is needed or desired

[0051] Acceleration of tissue formation. A generator of flow may be combined with manifolds and/or scaffolds to accelerate the rate of tissue formation within a natural healing response. The methods, scaffolds, manifolds, flow sources and systems of the invention may be used to accelerate tissue growth by augmenting formation of provisional matrices, facilitating its stable positioning, and aiding in recruitment of cells to the tissue space.

[0052] Stimulating the differentiation of stem cells along specific pathways. A generator of flow may be combined with manifolds and/or scaffolds to stimulate the differentiation of stem cells or other pluripotent cells into specific lineages. Application of flow using the methods, scaffolds, manifolds, flow sources and systems of the invention may be used to direct pluripotent cells into specific cell lineages needed to foster growth in the tissue space.

[0053] Introducing proteins, matrix, cells, or pharmaceuticals into the *in vivo* environment. A generator of flow may be combined with manifolds and/or scaffolds to introduce exogenous growth factors, proteins, cells, or pharmaceutical agents into the tissue space to augment tissue repair, regeneration, and/or maintenance.

[0054] Creating matrices *in vitro* for implantation *in vivo*. A generator of flow may be combined with manifolds and/or scaffolds to facilitate formation of matrices *in vitro* that may subsequently be used for *in vivo* transplantation.

[0055] Promoting integration of transplanted tissue. A generator of flow may be combined with manifolds and/or scaffolds to promote integration of transplanted tissue into the host environment. This can be applied to autograft, allograft, or xenograft transplants.

[0056] Directing extracellular matrix (ECM) deposition and orientation *in vitro*. A flow generator may be combined with manifolds and/or scaffolds to guide the directed deposition and orientation of ECM expressed by cells and tissues. The directed orientation of ECM has an impact in organizing and directing the attachment and colonization of subsequent cell layers and tissues.

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5 [0058] All references cited in this specification are hereby incorporated by reference. The discussion of the references herein is intended merely to summarize the assertions made by the authors and no admission is made that any reference constitutes prior art. Applicants reserve the right to challenge the accuracy and pertinence of the cited references.

10 [0059] In view of the above, it will be seen that the advantages of the invention are achieved and other advantages attained. As various changes could be made in the above methods and compositions without departing from the scope of the invention, it is intended that all matter contained in the above description and shown in the accompanying drawings shall be interpreted as illustrative and not in a limiting sense.

CLAIMS

We claim:

- 5 Claim 1. An apparatus for providing reduced pressure to a defect at a tissue site of a nerve, the apparatus comprising:
a nerve conduit having a generally tubular shape having walls including a luminal wall surrounding the tissue site to contain fluids within a luminal space between the tissue site and the luminal wall; and
10 a manifold having a porous body and a connector for receiving reduced pressure, and positioned within the luminal space adjacent the tissue site to distribute the reduced pressure to the defect.
- Claim 2. The apparatus of claim 1, wherein the defect is a severed, partially severed, pinched, or degenerated nerve.
- 15 Claim 3. The apparatus of claim 1, wherein the manifold is positioned on the distal side of the nerve relative to the tissue site.
- Claim 4. The apparatus of claim 1, wherein the manifold provides reduced pressure to the distal side of the nerve relative to the tissue site.
- Claim 5. The apparatus of claim 1, wherein the manifold is composed of a bioinert material.
- 20 Claim 6. The apparatus of claim 1, wherein the manifold is composed of a bioabsorbable material.
- Claim 7. The apparatus of claim 1, wherein a portion of the manifold includes a scaffold that facilitates tissue growth.
- 25 Claim 8. The apparatus of claim 7, wherein the scaffold is formed from a foam or gel material.
- Claim 9. The apparatus of claim 7, wherein the scaffold is a biological material selected from fibrin or collagen.

Claim 10. The apparatus of claim 7, wherein the scaffold material comprises a bioactive agent.

Claim 11. The apparatus of claim 10, wherein the bioactive agent is at least one of an antibiotic, an antibody and a growth factor.

5 Claim 12. The apparatus of claim 11, wherein the bioactive agent is a growth hormone (GH), a bone morphogenetic protein (BMP), transforming growth factor- α (TGF- α), a TGF- β , a fibroblast growth factor (FGF), granulocyte-colony stimulating factor (G-CSF), granulocyte/macrophage-colony stimulating factor (GM-CSF), epidermal growth factor (EGF), platelet derived growth factor (PDGF), insulin-like growth factor (IGF), vascular
10 endothelial growth factor (VEGF), hepatocyte growth factor/scatter factor (HGF/SF), an interleukin, tumor necrosis factor- α (TNF- α) or nerve growth factor (NGF).

Claim 13. The apparatus of claim 1, wherein the manifold is substantially cylindrical in shape with a longitudinal axis extending generally axially through the luminal space.

15 Claim 14. The apparatus of claim 13, wherein the defect is a severed nerve having severed ends with the manifold positioned between the severed ends.

Claim 15. The apparatus of claim 14, wherein the manifold having end surfaces further comprises at least one manifold rod having a first end fluidly coupled to one of the end surfaces of the manifold and a second end extending therefrom into the luminal space.

20 Claim 16. The apparatus of claim 15, wherein the manifold includes the at least one manifold rod extending only from one end surface.

Claim 17. The apparatus of claim 15, wherein the at least one manifold rod is composed of a bioabsorbable material.

Claim 18. The apparatus of claim 15, wherein the at least one manifold rod includes a scaffold that facilitates tissue growth.

25 Claim 19. The apparatus of claim 15, wherein the at least one manifold is tubular in shape with porous walls in fluid communication with the luminal space.

- Claim 20. The apparatus of claim 13, wherein the manifold having end surfaces further comprises an opening extending between the end surfaces along the longitudinal axis of the manifold.
- 5 Claim 21. The apparatus of claim 20, wherein the defect is a pinched nerve extending through the opening of the manifold.
- Claim 22. The apparatus of claim 21, wherein the manifold further comprises at least one manifold rod having a first end fluidly coupled to one of the end surfaces of the manifold and a second end extending therefrom into the luminal space.
- 10 Claim 23. The apparatus of claim 22, wherein the manifold includes the at least one manifold rod extending only from one end surface.
- Claim 24. The apparatus of claim 22, wherein the at least one manifold rod is composed of a bioabsorbable material.
- Claim 25. The apparatus of claim 22, wherein the at least one manifold rod includes a scaffold that facilitates tissue growth.
- 15 Claim 26. The apparatus of claim 1, wherein the wall of the nerve conduit includes a slice along the length of the nerve conduit that forms an opening whereby the nerve conduit is implantable around the tissue site and sealable with one or more closure elements.
- Claim 27. The apparatus of claim 1, wherein the nerve conduit is composed of a bioinert material.
- 20 Claim 28. The apparatus of claim 1, wherein the nerve conduit is composed of a bioabsorbable material.
- Claim 29. The apparatus of claim 28, wherein the nerve conduit is composed of collagen.
- Claim 30. The apparatus of claim 1, wherein the nerve conduit comprises pores.

Claim 31. The apparatus of claim 30, wherein the walls of the nerve conduit comprises pores sufficiently small to exclude cells from tissue surrounding the nerve conduit from entering into the luminal space.

5 Claim 32. The apparatus of claim 31, wherein the pores have an interior diameter of between about 5 μm and 50 μm .

Claim 33. A system for providing reduced pressure to a defect at a tissue site of a nerve, the system comprising:

a pressure source for supplying the reduced pressure;

10 a nerve conduit having a generally tubular shape having walls including a luminal wall surrounding the tissue site to contain fluids within a luminal space between the tissue site and the luminal wall; and

a manifold in fluid communication with the pressure source and positioned within the luminal space adjacent the tissue site to distribute the reduced pressure to the defect.

15 Claim 34. The system of claim 33, further comprising a fluid source in fluid communication with the manifold for supplying a fluid to the luminal space.

Claim 35. The system of claim 33, wherein the defect is a severed, partially severed, pinched, or degenerated nerve.

20 Claim 36. The system of claim 33, wherein the manifold is positioned on the distal side of the nerve relative to the tissue site.

Claim 37. The system of claim 33, wherein the manifold provides reduced pressure to the distal side of the nerve relative to the tissue site.

Claim 38. The system of claim 33, wherein the manifold is composed of a bioinert material.

25 Claim 39. The system of claim 33, wherein the manifold is composed of a bioabsorbable material.

Claim 40. The system of claim 33, wherein a portion of the manifold includes a scaffold that facilitates tissue growth.

Claim 41. The system of claim 40, wherein the scaffold is formed from a foam or gel material.

5 Claim 42. The system of claim 40, wherein the scaffold is a biological material selected from fibrin or collagen.

Claim 43. The system of claim 33, wherein the manifold is substantially tubular in shape having a luminal surface forming a channel in fluid communication with the defect.

10 Claim 44. The system of claim 43, wherein the defect is a pinched nerve extending through the channel of the manifold.

Claim 45. The system of claim 33, wherein the defect is a severed nerve and the manifold is substantially cylindrical in shape having a plurality of channels extending generally axially through the manifold in fluid communication with each end of the severed nerve.

15 Claim 46. The system of claim 40, wherein the scaffold material comprises a bioactive agent.

Claim 47. The system of claim 46, wherein the bioactive agent is at least one of an antibiotic, an antibody and a growth factor.

20 Claim 48. The system of claim 47, wherein the bioactive agent is a growth hormone (GH), a bone morphogenetic protein (BMP), transforming growth factor- α (TGF- α), a TGF- β , a fibroblast growth factor (FGF), granulocyte-colony stimulating factor (G-CSF), granulocyte/macrophage-colony stimulating factor (GM-CSF), epidermal growth factor (EGF), platelet derived growth factor (PDGF), insulin-like growth factor (IGF), vascular endothelial growth factor (VEGF), hepatocyte growth factor/scatter factor (HGF/SF), an interleukin, tumor necrosis factor- α (TNF- α) or nerve growth factor (NGF).

25

Claim 49. The system of claim 33, wherein the wall of the nerve conduit includes a slice along the length of the nerve conduit that forms an opening whereby the nerve conduit is implantable around the tissue site and sealable with one or more closure elements.

5 Claim 50. The system of claim 33, wherein the nerve conduit is composed of a bioinert material.

Claim 51. The system of claim 33, wherein the nerve conduit is composed of a bioabsorbable material.

Claim 52. The system of claim 51, wherein the nerve conduit is composed of collagen.

Claim 53. The system of claim 33, wherein the nerve conduit comprises pores.

10 Claim 54. The system of claim 53, wherein the walls of the nerve conduit comprises pores sufficiently small to exclude cells from tissue surrounding the nerve conduit from entering into the luminal space.

Claim 55. The system of claim 54, wherein the pores have an interior diameter of between about 5 μm and 50 μm .

15 Claim 56. A method for providing reduced pressure to a defect at a tissue site of a nerve, the method comprising:

implanting a manifold at the tissue site for receiving the reduced pressure;

implanting a nerve conduit having a luminal wall to surround the tissue site and the manifold for creating a luminal space between the luminal wall and the tissue site;

20 sealing the nerve conduit around the tissue site to contain fluids within the luminal space; and

applying reduced pressure through the manifold to the tissue site.

Claim 57. A method for repairing or regenerating a defect at a tissue site of a nerve, the method comprising:

implanting a manifold at the tissue site for receiving the reduced pressure;

implanting a nerve conduit having a luminal wall to surround the tissue site and the

5 manifold for creating a luminal space between the luminal wall and the tissue site;

sealing the nerve conduit around the tissue site to contain fluids within the luminal space;

applying reduced pressure through the manifold to the tissue site; and

10 wherein the application of reduced pressure facilitates the repair or regeneration of the defect.

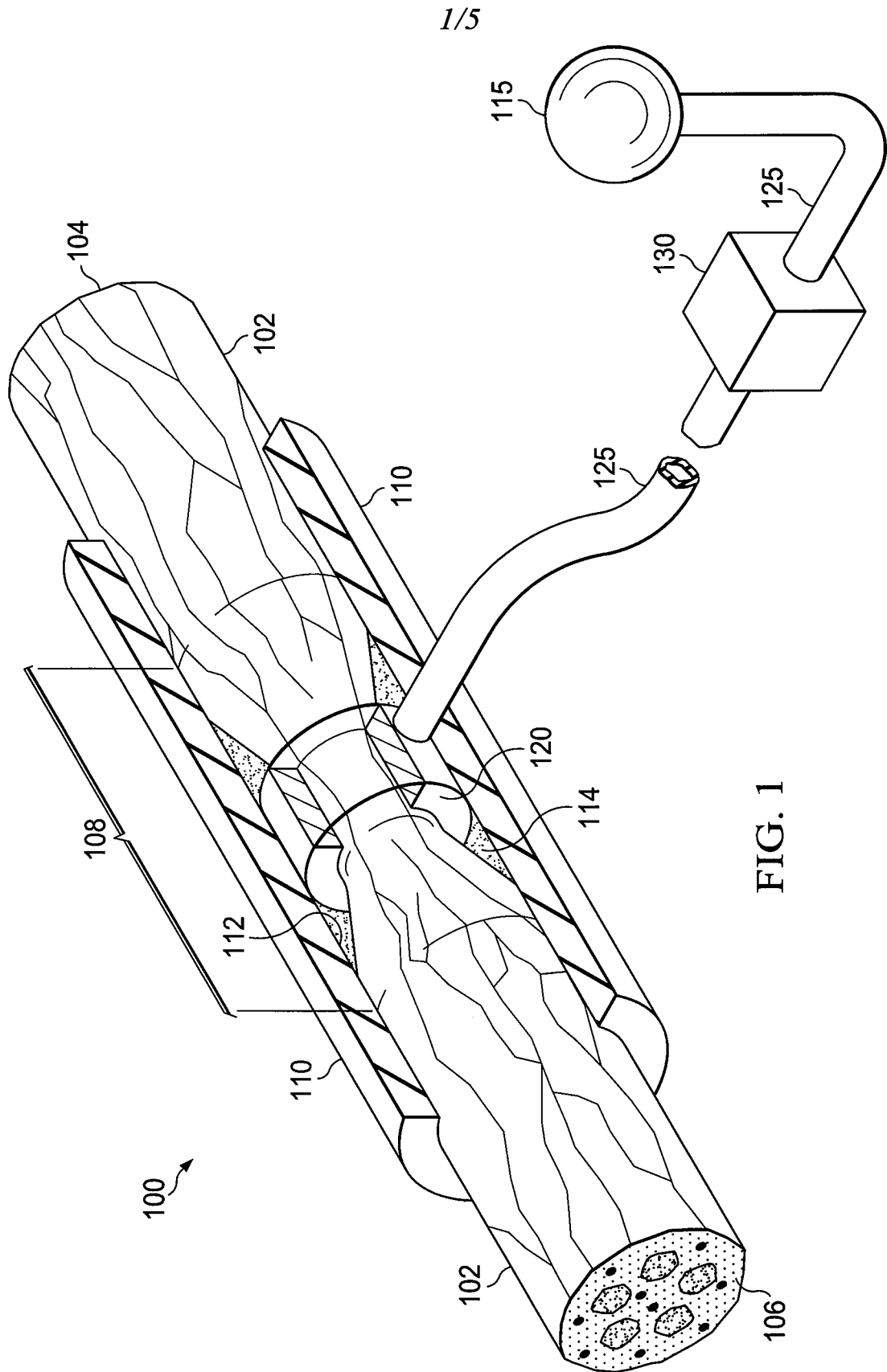


FIG. 1

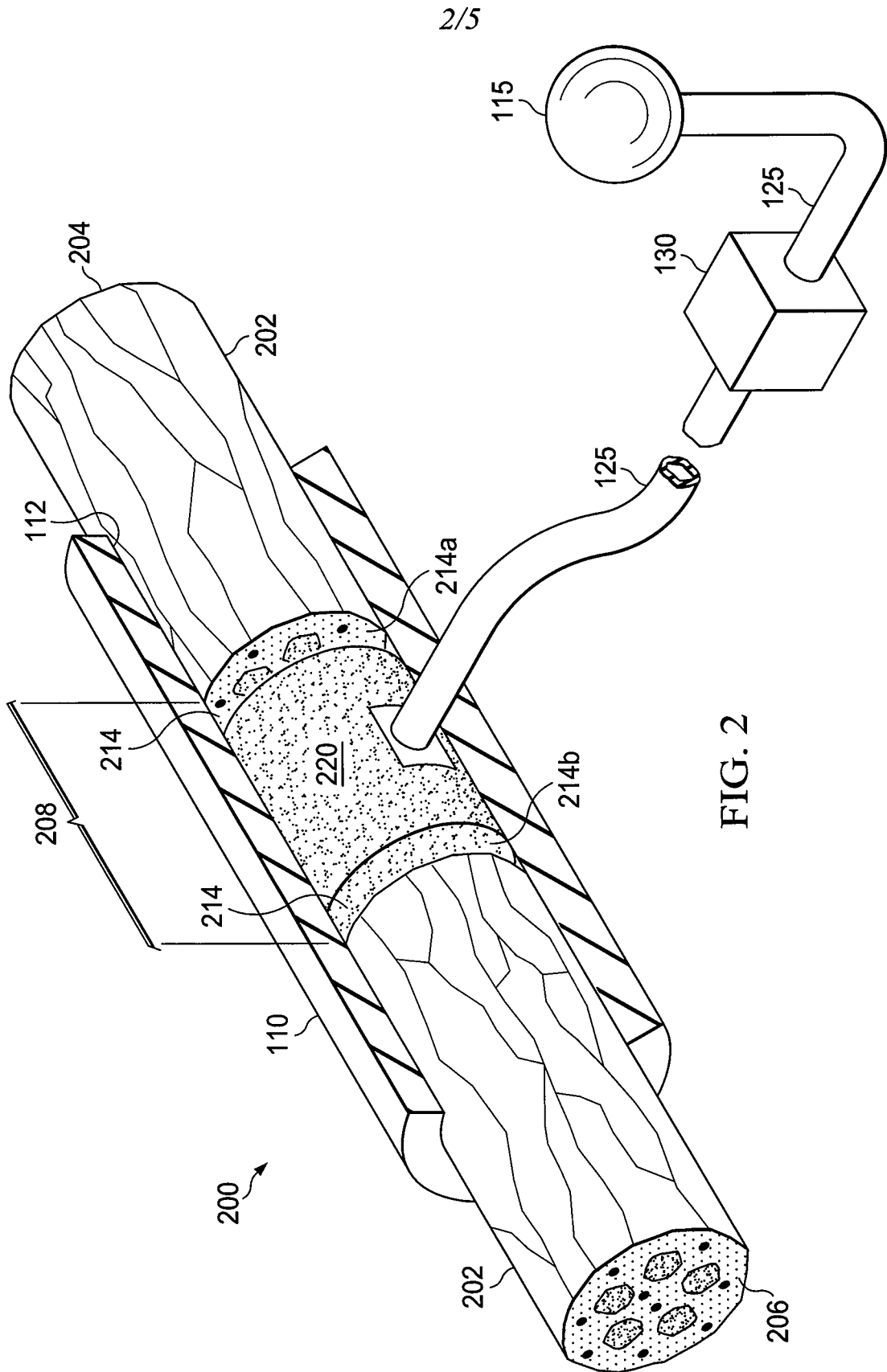


FIG. 2

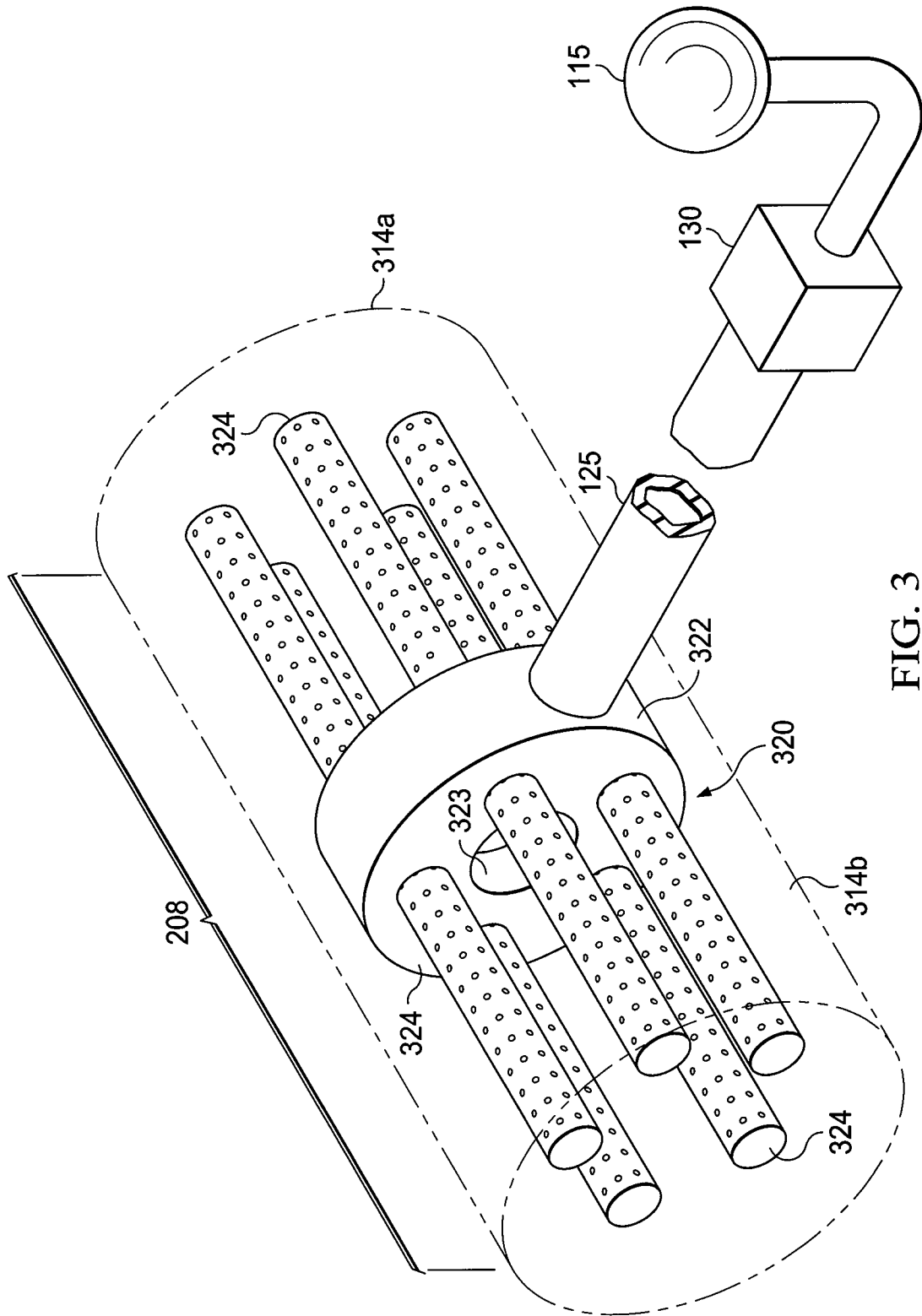


FIG. 3

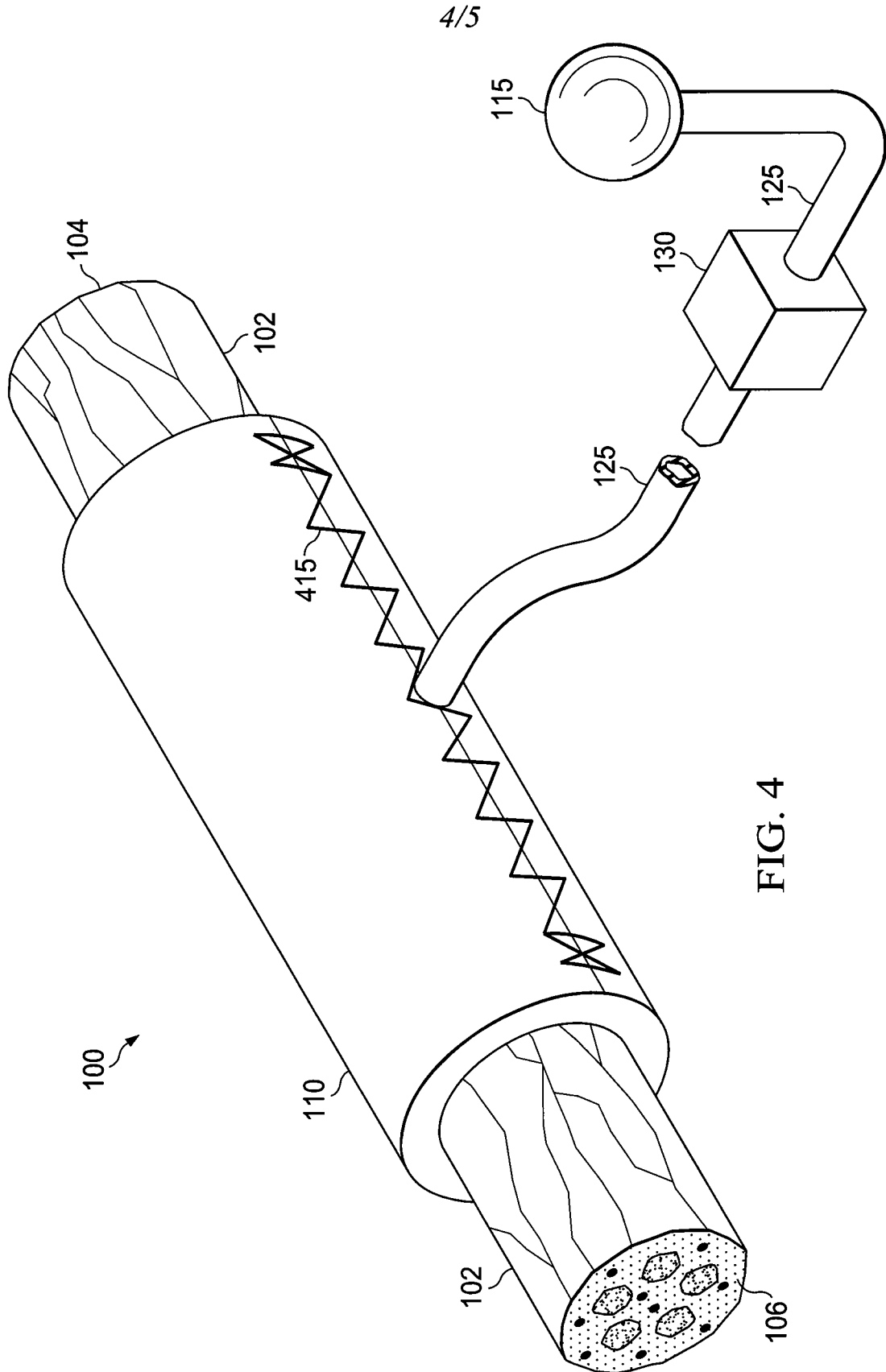


FIG. 4

FIG. 5

