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(54) SYSTEMS AND METHODS FOR DRUG DELIVERY, TREATMENT, AND MONITORING

SYSTEME UND VERFAHREN ZUR WIRKSTOFFFREISETZUNG, BEHANDLUNG UND ÜBERWACHUNG

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Description

FIELD

[0001] The present disclosure relates to systems and methods for drug delivery, treatment, and monitoring.

BACKGROUND

[0002] In convection-enhanced delivery (CED), drugs are infused locally into tissue through a needle, cannula, or microcatheter inserted into the tissue. Transport of the infused material is dominated by convection, which enhances drug penetration into the target tissue compared with diffusion-mediated delivery or systemic delivery.

[0003] CED has emerged as a leading investigational delivery technique for the treatment of several disorders. Clinical trials using existing devices show mixed results and suggest that the outcome of the therapy depends strongly on the extent of penetration and distribution of the drug into the target tissue, which is determined by infusion velocity, the relative rates of convection and elimination during CED, and various properties of the target tissue.

[0004] As infusion velocity increases, there can be a tendency for the infused fluid to flow back along the insertion pathway, between the exterior of the microcatheter designs have been constructed to reduce this backflow of the drug-containing fluid. However, fluid backflow during CED treatment still remains a critical problem in clinical practice. This is particularly true in the case of CED within the brain, as the poroelastic nature of the brain tissue contributes to backflow or reflux. There is therefore a need for improved CED devices, e.g., CED devices that reduce or eliminate backflow of the infused fluid between the exterior of the device and the surrounding tissue.

[0005] In some instances, it can be desirable to deliver drugs or other therapy (via convection-enhanced delivery or otherwise) over an extended period of time (e.g., several hours, days, weeks, months, years, and so forth). It can also be desirable to monitor various parameters associated with the treatment of a patient over an extended period of time. Accordingly, a need exists for improved systems and methods for drug delivery, treatment, and monitoring.

SUMMARY

[0006] Systems and methods for delivering a drug or other therapy over an extended period of time (e.g., several hours, days, weeks, months, years, and so forth) are disclosed herein, as are systems and methods for monitoring various parameters associated with the treatment of a patient. Systems and methods are also disclosed herein that generally involve CED devices with various features for reducing or preventing backflow. In some

embodiments, CED devices include a tissue-receiving space disposed proximal to a distal fluid outlet. Tissue can be compressed into or pinched/pinned by the tissuereceiving space as the device is inserted into a target region of a patient, thereby forming a seal that reduces or prevents proximal backflow of fluid ejected from the outlet beyond the tissue-receiving space. In some embodiments, CED devices include a bullet-shaped nose proximal to a distal fluid outlet. The bullet-shaped nose forms a good seal with surrounding tissue and helps re-

duce or prevent backflow of infused fluid. [0007] In some embodiments, an implantable delivery system includes a percutaneous access device through which drug-containing fluid can be delivered, a trunk line

¹⁵ having a plurality of independent fluid lumens extending therethrough, the plurality of fluid lumens being in fluid communication with a corresponding plurality of ports formed in the access device, a manifold configured to route the plurality of fluid lumens in the trunk line into a
²⁰ plurality of branch lines, each branch line including one or more corresponding fluid lumens disposed therein, and a skull anchor configured to be secured to the skull of a patient and being configured to couple the branch lines to corresponding microfluidic catheters configured
²⁵ to extend into the brain of the patient.

[0008] The system can include one or more inline filters disposed in the branch lines and configured to remove air, gas, bacteria, or particulates from fluid flowing through the branch lines. The trunk line, manifold, branch 30 lines, and skull anchor can be configured for long-term implantation beneath the skin of a patient. The access device can include one or more electrical connections for coupling extracorporeal electrical conductors to implanted electrical conductors. The trunk line, the mani-35 fold, at least one of the branch lines, the skull anchor, and at least one of the catheters include electrical conductors configured to provide an electrical path between the access device and a sensor or electrode of the at least one catheter. The skull anchor can be disposable 40 over first and second burr holes formed in the skull of the patient such that a first catheter coupled to the skull anchor extends through the first burr hole and a second catheter coupled to the skull anchor extends through the second burr hole. The skull anchor can be disposable 45 over a first burr hole such that first and second catheters coupled to the skull anchor extend through the first burr

[0009] At least one of the catheters can include an array of sensors at a distal end thereof. The array of sensors
can be printed on a substrate of the catheter. The array of sensors can be formed on a ribbon affixed to the catheter. The array of sensors can include at least one electrode. At least one of the catheters can include a sensor. The sensor can include at least one of an interrogatable sensor, a pressure sensor, a glutamate sensor, a pH sensor, a carbon dioxide sensor, an oxygen sensor, a neurotransmitter sensor, and a lactate sensor. The sensor can

hole.

be disposed in a fluid lumen of the at least one catheter adjacent an outlet port of the at least one catheter such that fluid flowing through the outlet port washes over the sensor. The at least one catheter can include at least one drug delivery channel and a dedicated patency channel through which fluid can be directed to clean the sensor. The at least one catheter can include a dedicated electrical conductor channel through which an electrical conductor coupled to the sensor extends, the electrical conductor channel being separate from a fluid delivery channel of the at least one catheter. The fluid delivery channel and the electrical conductor channel can intersect at a chamber in which the sensor is disposed.

[0010] At least one of the catheters can include a biodegradable scaffold on which one or more fluid channels are formed. The scaffold can be configured to biodegrade after being implanted in a patient, leaving behind only the one or more fluid channels. The scaffold can extend to a proximal end of the at least one catheter and can be coupled to the skull anchor. The scaffold can include an upper layer and a lower layer, and the one or more fluid channels can be sandwiched between the upper and lower scaffold layers.

[0011] At least one of the catheters can include a microtip having a proximal portion, a central portion, a distal portion, and at least one fluid channel extending along said proximal, central, and distal portions, the at least one fluid channel having an outlet port at a distal end thereof and an inlet port at a proximal end thereof, a first outer sheath disposed coaxially over the distal portion of the micro-tip such that the distal portion of the micro-tip protrudes from a distal end of the first outer sheath, a first tissue-receiving space defined between an exterior surface of the micro-tip and an interior surface of the distal end of the first outer sheath, a catheter body extending proximally from the micro-tip such that the at least one fluid channel of the micro-tip is in fluid communication with a respective inner lumen of the catheter body, and a nose portion disposed over at least the central portion of the micro-tip and extending between the first outer sheath and the catheter body such that the nose portion defines an exterior surface that tapers from a reduced distal diameter corresponding to the outside diameter of the first outer sheath to an enlarged proximal diameter corresponding to the outside diameter of the catheter body.

[0012] In some embodiments, a treatment method includes implanting a first catheter in a first location in a patient's brain, implanting a second catheter in a second location in the patient's brain, attaching a skull anchor to which the first and second catheters are coupled to the patient's skull, coupling at least one line to the first and second catheters and routing the at least one line beneath the patient's skin to couple the at least one line to a crosscutaneous access device, the at least one line including at least one fluid lumen and at least one electrical conductor, and delivering fluid through the access device, the at least one of

the first and second catheters into the patient's brain. [0013] The method can include delivering energy to an electrode disposed in at least one of the first and second catheters via the access device and the at least one electrical conductor. The method can include delivering fluid through a fluid lumen of the first catheter and delivering energy through an electrode of the first catheter. The method can include delivering fluid lumen of the first catheter and delivering fluid lumen of the first catheter and delivering herefy through an electrode of the first catheter.

¹⁰ trode of the second catheter. The method can include communicating the output of a sensor disposed in at least one of the first or second catheters via the at least one electrical conductor and the access device. The method can include delivering fluid through a fluid lumen of the

¹⁵ first catheter and monitoring a parameter using a sensor of the first catheter. The method can include delivering fluid through a fluid lumen of the first catheter and monitoring a parameter using a sensor of the second catheter. The method can include adjusting the delivery of energy

²⁰ or fluid via the first catheter based on the output of a sensor disposed in the first or second catheters. The method can include maintaining the patency of a sensor disposed in the first or second catheters by flushing fluid through a fluid outlet port in which the sensor is disposed.

²⁵ The method can include fluid through a dedicated patency channel of the first catheter to maintain the patency of a sensor disposed in the first catheter while delivering a drug-containing fluid through a drug-delivery channel of the first catheter.

30 [0014] The method can include at least one of: monitoring the movement of infusate delivered through the first catheter using a sensor disposed in the second catheter; monitoring the spread of a viral vector administered through the first catheter using a sensor disposed in the

³⁵ second catheter; and monitoring the effects of neurostimulation applied via the first catheter using a sensor disposed in the second catheter. The method can include aspirating tissue through at least one of the first and second catheters, the at least one line, and the access de-

40 vice. The first and second catheters can be implanted through a single burr hole in the patient's skull. The first and second catheters can be implanted through first and second burr holes in the patient's skull over which the skull anchor is disposed. The at least one line can include

⁴⁵ a branch line coupled to a trunk line by a manifold. The method can include filtering fluid through an in-line filter disposed in the at least one line. Delivering the fluid can include delivering the fluid via convection-enhanced delivery. Delivering the fluid can include delivering the fluid

to a target site within a patient over a period of hours, days, weeks, months, or years. The method can include a support scaffold of the first catheter to biodegrade leaving behind only fluid conduits of the first catheter. The method can include allowing upper and lower scaffolds
 of the first catheter to biodegrade.

[0015] Implanting the first catheter can include advancing a fluid conduit having a first outer sheath disposed therearound into tissue to compress tissue into a first

tissue-receiving space defined between an exterior surface of the fluid conduit and an interior surface of the distal end of the first outer sheath; and delivering the fluid under positive pressure through the fluid conduit and into a portion of the tissue adjacent to a distal end of the fluid conduit. Advancing the fluid conduit can include urging a nose portion into contact with tissue, the nose portion extending between the first outer sheath and a proximal catheter body such that the nose portion tapers from a reduced distal diameter corresponding to the outside diameter of the first outer sheath to an enlarged proximal diameter corresponding to the outside diameter of the catheter body.

[0016] In some embodiments, a convection-enhanced-delivery (CED) device is provided that includes a micro-tip having a proximal portion, a central portion, a distal portion, and at least one fluid channel extending along said proximal, central, and distal portions, the at least one fluid channel having an outlet port at a distal end thereof and an inlet port at a proximal end thereof. The device also includes a first outer sheath disposed coaxially over the distal portion of the micro-tip such that the distal portion of the micro-tip protrudes from a distal end of the first outer sheath, a first tissue-receiving space defined between an exterior surface of the micro-tip and an interior surface of the distal end of the first outer sheath, and a catheter body extending proximally from the micro-tip such that the at least one fluid channel of the micro-tip is in fluid communication with a respective inner lumen of the catheter body. The device also includes a nose portion disposed over at least the central portion of the micro-tip and extending between the first outer sheath and the catheter body such that the nose portion defines an exterior surface that tapers from a reduced distal diameter corresponding to the outside diameter of the first outer sheath to an enlarged proximal diameter corresponding to the outside diameter of the catheter body.

[0017] The tissue-receiving space can be configured to compress tissue received therein as the device is advanced through the tissue. Tissue compressed by the tissue-receiving space can form a seal that reduces proximal backflow of fluid ejected from the outlet port of the at least one fluid channel beyond the tissue-receiving space. The device can include a second outer sheath disposed over the first outer sheath such that a second tissue-receiving space is defined between an exterior surface of the first outer sheath and an interior surface of a distal end of the second outer sheath. The interior surface of the distal end of the first outer sheath can be shaped to compress tissue received therein as the device is advanced through the tissue. The interior surface of the distal end of the first outer sheath can be conical, convex. and/or concave.

[0018] An inside diameter of the distal end of the first outer sheath can be about 1 μ m to about 200 μ m greater than an outside diameter of the distal portion of the microtip. An inside diameter of the distal end of the first outer

sheath can be about 10 percent to about 100 percent greater than an outside diameter of the distal portion of the micro-tip. The first outer sheath can have a circular outside cross-section. The at least one fluid channel can be formed from at least one of a parylene composition, a silastic composition, a polyurethane composition, and a PTFE composition. The device can include a fluid res-

ervoir in fluid communication with the inner lumen of the catheter body and configured to supply a fluid thereto
 under positive pressure. The micro-tip can be flexible. The micro-tip can include an embedded microsensor.

[0019] The embedded microsensor can include at least one of an interrogatable sensor, a pressure sensor, a glutamate sensor, a pH sensor, a temperature sensor,

¹⁵ an ion concentration sensor, a carbon dioxide sensor, an oxygen sensor, and a lactate sensor. The distal end of the micro-tip can have an atraumatic shape configured to penetrate tissue without causing trauma. The micro-tip can contain a quantity of a drug, can be coated with

²⁰ a drug, and/or can be impregnated with a drug. The drug can include at least one of an antibacterial agent, an antiinflammatory agent, a corticosteroid, and dexamethasone. The micro-tip can include a substrate having the at least one fluid channel formed thereon. The substrate

can have a rectangular transverse cross-section. The catheter body can be formed from a rigid material. Each inner lumen of the catheter body can be defined by a sleeve formed from a flexible material. The catheter body can be formed from at least one of ceramic, PEEK, and
 polyurethane. Each sleeve can be formed from at least

one of polyimide, pebax, PEEK, polyurethane, silicone, and fused silica. The catheter body can be formed from a flexible material. The device can be assembled by forming the nose portion by molding the nose portion over the

³⁵ first outer sheath, inserting the micro-tip into a proximal end of the nose portion, coupling the proximal portion of the micro-tip to the catheter body, and injecting a flowable material through an inlet port formed in the nose portion to fill the interior of the nose portion and secure the microtip and catheter body to the nose portion.

[0020] In some embodiments, a convection-enhanced-delivery (CED) device is provided that includes a fluid conduit having proximal and distal ends, a first outer sheath disposed coaxially over the fluid conduit

⁴⁵ such that the fluid conduit extends out of a distal end of the first outer sheath, and a first tissue-receiving space defined between an exterior surface of the fluid conduit and an interior surface of the distal end of the first outer sheath.

50 [0021] In some embodiments, a micro-molding device is provided that includes a mold cavity sized and configured to receive a catheter body and a catheter micro-tip therein such that at least one fluid channel of the microtip is at least partially disposed within a corresponding fluid line of the catheter body. The device also includes one or more mold channels though which a mold fluid can be injected to fill the mold cavity and secure the microtip to the catheter body such that the at least one fluid

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channel of the micro-tip is in fluid communication with the at least one fluid line of the catheter body. The device can be transparent to allow UV light to pass therethrough to cure mold fluid disposed within the mold cavity. The mold cavity can be sized and configured to form a bullet nose portion over the micro-tip and over at least a portion of an outer sheath received in the mold cavity.

[0022] In some embodiments, a method of delivering a therapeutic agent to a patient is provided. The method includes advancing a fluid conduit having a first outer sheath disposed therearound into tissue to compress tissue into a first tissue-receiving space defined between an exterior surface of the fluid conduit and an interior surface of the distal end of the first outer sheath. The method also includes delivering fluid containing the therapeutic agent under positive pressure through the fluid conduit and into a portion of the tissue adjacent to a distal end of the fluid conduit.

[0023] The method can include delivering a sealing gel through the fluid conduit, before delivering the fluid containing the therapeutic agent, to fill one or more voids that exist between the fluid conduit and the tissue. Tissue compressed into the first tissue-receiving space can form a seal that reduces proximal backflow of fluid ejected from the distal end of the fluid conduit beyond the tissuereceiving space. The method can include advancing a second outer sheath disposed over the first outer sheath into the tissue such that tissue is compressed into a second tissue-receiving space defined between an exterior surface of the first outer sheath and an interior surface of the distal end of the second outer sheath. The interior surface of the distal end of the first outer sheath can be at least one of cylindrical, conical, convex, and concave. The method can include controlling delivery of fluid through the fluid conduit based on an output of a microsensor embedded in the fluid conduit. The method can be used to treat at least one condition selected from central-nervous-system (CNS) neoplasm, intractable epilepsy, Parkinson's disease, Huntington's disease, stroke, lysosomal storage disease, chronic brain injury, Alzheimer's disease, amyotrophic lateral sclerosis, balance disorders, hearing disorders, and cavernous malformations. Advancing the fluid conduit can include urging a nose portion into contact with tissue, the nose portion extending between the first outer sheath and a proximal catheter body such that the nose portion tapers from a reduced distal diameter corresponding to the outside diameter of the first outer sheath to an enlarged proximal diameter corresponding to the outside diameter of the catheter body. The fluid conduit can be coupled to a distal end of a flexible catheter and the method can include inserting the catheter through an incision, positioning the fluid conduit in proximity to the portion of the tissue using stereotactic targeting, removing a stylet inserted through the catheter, tunneling a proximal end of the catheter beneath the scalp of the patient, and coupling one or more proximal fluid connectors of the catheter to a fluid delivery system.

[0024] The present invention further provides systems as claimed.

BRIEF DESCRIPTION OF THE DRAWINGS

[0025] The invention will be more fully understood from the following detailed description taken in conjunction with the accompanying drawings, in which:

¹⁰ FIG. 1 is a perspective view of one exemplary embodiment of a CED device;

FIG. 2 is a cross-sectional view of the device of FIG. 1, taken in a plane normal to the longitudinal axis of the device;

FIG. 3 is a schematic view of a fluid delivery system that includes the device of FIG. 1;

FIG. 4 is a schematic view of the device of FIG. 1 inserted into tissue;

FIG. 5 is a perspective view of another exemplary embodiment of a CED device;

FIG. 6A is a plan view of another exemplary embodiment of a CED device;

FIG. 6B is a plan view of another exemplary embodiment of a CED device;

FIG. 6C is a plan view of another exemplary embodiment of a CED device;

FIG. 7 is a perspective view of another exemplary embodiment of a CED device;

FIG. 8 is another perspective view of the CED device of FIG. 7;

FIG. 9 is a perspective view of the CED device of FIG. 7 with a depth stop and tip protector;

FIG. 10 is a plan view of the CED device of FIG. 7 with a length of extension tubing;

FIG. 11 is a perspective view of a micro-tip of the CED device of FIG. 7;

FIG. 12 is a perspective view of an exemplary embodiment of a molding system;

FIG. 13 is a perspective view of the CED device of FIG. 7 being manufactured using the molding system of FIG. 12;

FIG. 14 is a top view of the CED device of FIG. 7 being manufactured using the molding system of

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FIG. 12;

FIG. 15 is another perspective view of the CED device of FIG. 7 being manufactured using the molding system of FIG. 12;

FIG. 16 is a partially-exploded sectional perspective view of another exemplary embodiment of a CED device;

FIG. 17 is a partially-exploded perspective view of the CED device of FIG. 16;

FIG. 18 is a perspective view of the CED device of FIG. 16;

FIG. 19 is a map of mold filling time for the nose portion of the CED device of FIG. 16;

FIG. 20 is a perspective view of an exemplary em-²⁰ bodiment of a molding system for forming the nose portion of the CED device of FIG. 16;

FIG. 21 is a scale drawing of an exemplary embodiment of the nose portion of the CED device of FIG. ²⁵ 16;

FIG. 22 is a series of images showing infusion of dye using a CED device into a gel designed to simulate tissue;

FIG. 23 is another series of images showing infusion of dye using a CED device into a gel designed to simulate tissue;

FIG. 24 is a magnetic resonance image of a pig brain in which a CED device is inserted and a gadolinium dye is infused;

FIG. 25 is a series of magnetic resonance images showing infusion of gadolinium into white matter of a pig's brain at flow rates of 1, 3, 5, 10, and 20 μ L/min using a CED device;

FIG. 26 is a series of magnetic resonance images 45 showing infusion of gadolinium into the thalamus of a pig's brain at flow rates of 1, 3, 5, 10, and 20 μ L/min using a CED device;

FIG. 27 is a series of magnetic resonance images 50 showing infusion of gadolinium into the putamen of a pig's brain at flow rates of 1, 2, 5, 10, and 15 μ L/min using a CED device;

FIG. 28 is a series of magnetic resonance images showing infusion of gadolinium into the white matter of a pig's brain at a flow rate of 5 μ L/min using a CED device after infusion periods of 1, 9, 16, 24, and 50

minutes;

FIG. 29 is a magnetic resonance image and an in vivo imaging system image of the thalamus of a pig's brain when a CED device is used to simultaneously infuse galbumin and IVIS dye;

FIG. 30 is a comparison of infusate concentration using a CED device of the type described herein to simulated infusate concentration using a traditional catheter;

FIG. 31 is a comparison of tissue expansion using a CED device of the type described herein to simulated tissue expansion using a traditional catheter;

FIG. 32 is perspective view of a delivery and/or monitoring system;

FIG. 33 is a perspective view of a portion of the system of FIG. 32;

FIG. 34 is a perspective view of another portion of the system of FIG. 32;

FIG. 35 is a perspective view of another portion of the system of FIG. 32;

FIG. 36 is a perspective view of another portion of the system of FIG. 32;

FIG. 37 is a perspective view of an exemplary CED device and burr hole adapter;

FIG. 38 is a perspective view of the proximal end of the CED device and the burr hole adapter of FIG. 37;

FIG. 39 is a perspective view of another exemplary CED device;

FIG. 40 is a perspective view of a proximal end of the CED device of FIG. 39;

FIG. 41 is a perspective view of an exemplary CED device after a support scaffold biodegrades;

FIG. 42 is a perspective view of another exemplary CED device after a support scaffold biodegrades;

FIG. 43 is a perspective view of an exemplary CED device with an array of sensors and/or electrodes;

FIG. 44 is another perspective view of the CED device of FIG. 43;

FIG. 45 is a perspective view of a CED device with a sensor disposed in a fluid outlet port;

FIG. 46 is a perspective view of a CED device with a dedicated electrical conductor channel, a lid portion of the CED device being shown in phantom;

FIG. 47 is another perspective view of the CED device of FIG. 46; and

FIG. 48 is another perspective view of the CED device of FIG. 46.

DETAILED DESCRIPTION

[0026] Certain exemplary embodiments will now be described to provide an overall understanding of the principles of the structure, function, manufacture, and use of the methods, systems, and devices disclosed herein. One or more examples of these embodiments are illustrated in the accompanying drawings. Those skilled in the art will understand that the methods, systems, and devices specifically described herein and illustrated in the accompanying drawings are non-limiting exemplary embodiments and that the scope of the present invention is defined solely by the claims. The features illustrated or described in connection with one exemplary embodiment may be combined with the features of other embodiments. Such modifications and variations are intended to be included within the scope of the present invention.

[0027] Systems and methods for delivering a drug or other therapy over an extended period of time (e.g., several hours, days, weeks, months, years, and so forth) are disclosed herein, as are systems and methods for monitoring various parameters associated with the treatment of a patient. Systems and methods are also disclosed herein that generally involve CED devices with various features for reducing or preventing backflow. In some embodiments, CED devices include a tissue-receiving space disposed proximal to a distal fluid outlet. Tissue can be compressed into or pinched/pinned by the tissuereceiving space as the device is inserted into a target region of a patient, thereby forming a seal that reduces or prevents proximal backflow of fluid ejected from the outlet beyond the tissue-receiving space. In some embodiments, CED devices include a bullet-shaped nose proximal to a distal fluid outlet. The bullet-shaped nose forms a good seal with surrounding tissue and helps reduce or prevent backflow of infused fluid.

[0028] FIG. 1 illustrates one exemplary embodiment of a CED device 10. The device 10 generally includes a fluid conduit 12 and an outer sheath 14. The outer sheath 14 can be disposed coaxially over the fluid conduit 12 such that the fluid conduit 12 extends out of a distal end 16 of the outer sheath 14. The fluid conduit 12 and the outer sheath 14 can be sized and dimensioned such that a tissue-receiving space 18 is formed between an exterior surface of the fluid conduit 12 and an interior surface of the distal end 16 of the outer sheath 14.

[0029] The fluid conduit 12 can define one or more fluid

lumens that extend generally parallel to the central longitudinal axis of the device 10. The fluid conduit 12 can include a fluid inlet port (not shown in FIG. 1) and a fluid outlet port 20. While a single fluid outlet port 20 is shown in the illustrated embodiment, it will be appreciated that the device can include a plurality of fluid outlet ports, as well as a plurality of fluid inlet ports and a plurality of fluid lumens extending therebetween. The fluid inlet port can be positioned at a proximal end of the device 10, and can

¹⁰ allow the fluid conduit 12 to be placed in fluid communication with a fluid reservoir, e.g., via one or more catheters, pumps, meters, valves, or other suitable control devices. Such control devices can be used to regulate the pressure at which fluid is supplied to the device 10, or

¹⁵ the rate or volume of fluid that is supplied to the device 10. [0030] Fluid supplied to the conduit 12 though the fluid inlet port can be directed through one or more inner lumens of the conduit 12 and released through the one or more fluid outlet ports 20. The fluid outlet ports 20 can

²⁰ be sized, shaped, and/or positioned to control various release parameters of the fluid. For example, the fluid outlet ports 20 can be configured to control the direction in which fluid is released from the device 10, the distribution of the fluid within the target tissue, and the velocity

or pressure at which the fluid is released. In exemplary embodiments, the size of the fluid outlet ports can progressively increase towards the distal end of the device 10, which can advantageously compensate for pressure loss that occurs along the length of the device such that
fluid is released from each of the plurality of fluid outlet ports at substantially the same pressure. The fluid outlet ports can also be positioned at various points around the circumference of the fluid conduit 12 or can be shaped to control the release direction of the fluid.

³⁵ [0031] The fluid conduit 12 and/or the outer sheath 14 can have circular outside cross-sections, which can advantageously allow the device 10 to rotate within the tissue without causing trauma or forming large gaps between the exterior of the device and the surrounding tissue that might increase backflow. The fluid conduit 12

can also be flexible to allow it to move with the tissue in which it is inserted. While a generally-cylindrical fluid conduit 12 is shown, the fluid conduit 12 can also have a non-cylindrical or polygonal cross-section. For example,

⁴⁵ as described below with respect to FIG. 7, the fluid conduit 12 can be a microfabricated tip that includes a substrate having a square or rectangular cross-section with one or more fluid channels disposed thereon. The interior of the outer sheath 14 can be shaped to substantially

⁵⁰ correspond to the cross-section of the fluid conduit 12. Alternatively, the outer sheath 14 can have an interior cross-sectional shape that differs from the exterior cross-sectional shape of the fluid conduit 12. For example, the outer sheath 14 can have a substantially cylindrical inte-⁵⁵ rior cross-sectional shape at its distal end, while the fluid conduit 12 can have a substantially square or rectangular exterior cross-sectional shape, thereby defining the tissue-receiving space 18 between the exterior of the fluid

conduit 12 and the interior of the outer sheath 14.

[0032] As noted above, the outer sheath 14 can be disposed coaxially over the fluid conduit 12 such that the fluid conduit 12 extends out of the distal end 16 of the outer sheath 14. A clearance space between the exterior surface of the fluid conduit 12 and the interior surface of the sheath 14 can define the tissue-receiving space 18. For example, as shown in FIG. 2, the fluid conduit 12 can have an outside diameter D1 that is less than an inside diameter D2 of the outer sheath 14. The degree to which the diameter D2 exceeds the diameter D1 can dictate the amount of tissue that is compressed into or pinched by the tissue-receiving space 18.

[0033] In some embodiments, an adhesive or other filler can be disposed between the fluid conduit 12 and the sheath 14 to hold the fluid conduit in a fixed longitudinal position relative to the sheath and to maintain the fluid conduit in the center of the sheath (e.g., such that the tissue-receiving space 18 has a uniform width about the circumference of the fluid conduit). For example, the tissue-receiving space 18 can extend proximally a first distance from the distal end 16 of the sheath 14, after which point the clearance space between the fluid conduit 12 and the sheath 14 can be filled. In some embodiments, the sheath 14 can have a stepped, tapered, or other similarly-shaped interior such that a clearance space exists along a distal portion of the sheath 14 and no clearance space exists along a proximal portion of the sheath 14. [0034] In exemplary embodiments, the inside diameter of the distal end 16 of the outer sheath 14 can be about 1 μ m to about 1000 μ m, about 1 μ m to about 500 μ m, about 1 μ m to about 200 μ m, or about 1 μ m to about 20 μ m greater than the outside diameter of the fluid conduit 12. In exemplary embodiments, the inside diameter of the distal end 16 of the outer sheath 14 can be about 5 percent to about 500 percent, about 5 percent to about 250 percent, about 10 percent to about 100 percent, or about 10 percent to about 20 percent greater than the outside diameter of the fluid conduit 12. In exemplary embodiments, the diameter D1 can be about 50 μm to about 2000 μ m, about 50 μ m to about 1000 μ m, or about 50 μ m to about 200 μ m. In exemplary embodiments, diameter D2 can be about 51 μ m to about 5000 μ m, about 55 μ m to about 1000 μ m, or about 55 μ m to about $200 \,\mu\text{m}$. The tissue-receiving space 18 can extend along the entire length of the outer sheath 14, or along only a portion of the outer sheath (e.g., along about 1 mm to about 100 mm, about 1 mm to about 50 mm, or about 1 mm to about 10 mm of the distal-most portion of the outer sheath).

[0035] The fluid conduit 12 and the outer sheath 14 can be formed from any of a variety of materials, including parylene compositions, silastic compositions, polyurethane compositions, PTFE compositions, silicone compositions, and so forth.

[0036] In some embodiments, the device 10 can be mounted on a support scaffold (not shown) to provide structural rigidity to the device and facilitate insertion into

the target tissue. Exemplary support scaffolds are illustrated and described in U.S. Publication No. 2013/0035560, filed on August 1, 2012, entitled "MULTI-DIRECTIONAL MICROFLUIDIC DRUG DELIVERY DE-VICE," the entire contents of which are incorporated herein by reference. To assist with tissue penetration and navigation, the distal end of the fluid conduit 12 and/or the distal end of the scaffold can be tapered, pointed, and/or sharpened. In some embodiments, the fluid con-

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¹⁰ duit 12 and/or the scaffold can be provided with a rounded atraumatic tip so as to facilitate insertion through tissue without causing trauma to the tissue. The support scaffold can be rigid or semi-rigid and can be formed from a degradable thermoplastic polymer, for example, a degra-

¹⁵ dable thermoplastic polyester or a degradable thermoplastic polycarbonate. In some embodiments, the support scaffold can be formed from poly(lactic-co-glycolic acid) (PLGA) and can be configured to biodegrade within the target tissue. This can advantageously eliminate the need to remove the support scaffold once the device 10

- is positioned within target tissue, thereby avoiding the potential to disrupt the positioning of the fluid conduit 12. Any of a variety of other materials can also be used to form the support scaffold, including silicon or various ce-
- 25 ramics, metals, and plastics known in the art. The scaffold can have a width of approximately 100 μm to approximately 200 μm and can have a length that varies depending on the target tissue (e.g., depending on the depth at which the target tissue is situated). In one embodiment,
- the scaffold is between 2 cm and 3 cm long. A variety of techniques can be used to couple the fluid conduit 12 and/or the outer sheath 14 to the support scaffold, such as surface tension from a water drop, adhesives, and/or a biocompatible petroleum jelly.

³⁵ [0037] Any of the fluid conduit 12, the outer sheath 14, and/or the support scaffold can contain or can be impregnated with a quantity of a drug. Alternatively, or in addition, a surface of these components can be coated with a drug. Exemplary drugs include anti-inflammatory components, drug permeability-increasing components, delayed-release coatings, and the like. In some embodi-

ments, one or more components of the device 10 can be coated or impregnated with a corticosteroid such as dexamethasone which can prevent swelling around the in ⁴⁵ jection site and disruptions to the fluid delivery pattern

[0038] The device 10 can also include one or more sensors 22 mounted in or on the fluid conduit 12, the sheath 14, or the scaffold. The sensors 22 can include
temperature sensors, pH sensors, pressure sensors, oxygen sensors, tension sensors, interrogatable sensors, glutamate sensors, ion concentration sensors, carbon dioxide sensors, lactate sensors, neurotransmitter sensors, or any of a variety of other sensor types, and can provide feedback to a control circuit which can in turn regulate the delivery of fluid through the device 10 based on one or more sensed parameters. One or more electrodes 24 can also be provided in or on the fluid conduit

12, the sheath 14, or the scaffold, which can be used to deliver electrical energy to target tissue, e.g., to stimulate the target tissue or to ablate the target tissue. In one embodiment, electrical energy is delivered through an electrode 24 while a drug is simultaneously delivered through the fluid conduit 12.

[0039] FIG. 3 is a schematic illustration of a drug delivery system 26 that includes the device 10. The system 26 includes a reservoir 28 of a drug-containing fluid that is coupled to a pump 30 via a control valve 32. When the control valve 32 is opened, fluid in the reservoir 28 is supplied under pressure by the pump 30 to a pressure regulator 34, which can adjust a pressure at which the fluid is supplied to the device 10. The control valve 32, pump 30, and regulator 34 can be operatively coupled to a controller 36 which can include a microprocessor and a memory and can be configured to execute a drugdelivery control program stored in a non-transitory computer-readable storage medium. The controller 36 can be configured to open or close the valve 32, to turn the pump 30 on or off, to change an output pressure of the pump 30, and/or to adjust a pressure set point of the regulator 34. The controller 36 can also receive information indicative of a sensed parameter via a feedback loop that includes one or more sensors 22 mounted in or on the device 10. Thus, in response to feedback from one or more sensors 22 implanted with the device 10, the controller 36 can start or stop the flow of fluid to the device 10, increase or decrease the pressure at which fluid is supplied to the device 10, etc. In one embodiment, the device 10 includes a pressure sensor 22 that measures a fluid pressure in the vicinity of the device 10 and the controller 36 is configured to maintain the fluid supply pressure at a substantially constant level based on feedback from the pressure sensor 22. It will be appreciated that some or all of the components of the drug delivery system 26 can be implanted in a patient and that some or all of the components can be disposed external to a patient.

[0040] The device 10 can be used for CED of drugs to treat disorders of the brain, spine, ears, neural tissue, or other parts of a human or animal body. When used in the brain, the device 10 can circumvent the blood-brain barrier (BBB) by infusing drugs under positive pressure directly into tissue. The device 10 can provide a number of advantages, such as 1) a smaller cross-sectional area compared with conventional needles used in CED; 2) less disturbance to tissue when inserted into the brain than conventional needles; 3) the reduction or elimination of backflow or reflux along the outside of the inserted part, which in turn, permits higher rates of drug delivery in the device 10 compared with conventional needles; 4) minimal or no occlusion of the fluid delivery conduit 12 during insertion into the brain; 5) multiple lumens can be provided through the fluid conduit 12, each conducting a distinct fluid (drug), which allows simultaneous, sequential, or programmed delivery of multiple agents; 6) the device 10 has the potential to serve simultaneously as a

drug delivery system and as a sensor-equipped probe to measure local tissue characteristics such as, but not limited to, pressure, pH, ion-specific concentrations, location, and other parameters; and 7) the device 10 allows for directional control of the drug release pattern.

[0041] In use, as described further below, the device 10 can be functionally attached to the distal end of a long, thin insertion vehicle such as a cannula or a needle in or on which a fluid attachment can be made to the fluid inlet

¹⁰ port of the device's fluid conduit 12. This can be especially advantageous in applications involving penetration of relatively thick tissue, e.g., insertion through a human skull. [0042] In addition to delivering a drug-containing fluid, the device 10 can also be used to deliver enzymes or

 other materials to modify tissue permeability and improve drug distribution in the targeted tissue. For example, penetration of drug-containing nanoparticles into brain tissue can be enhanced by enzymatic digestion of at least one brain extracellular matrix component and intracranial in fusion of the nanoparticle into the brain tissue. In another embodiment, at least one enzyme can be immobilized to

a surface of the nanoparticle during the step of enzymatic digestion. The device 10 can provide the ability to deliver enzymatic and/or other materials that can, e.g., modify
 the drug delivery site, and therapeutic materials, in virtually any order, sequencing, and/or timing without the need to use different delivery devices and the potential

complications involved in doing so.

[0043] The device 10 can also be used to biopsy tissue,
for example by passing a stylet or a grasping tool through the fluid conduit 12 to a target site and then withdrawing the stylet or grasping tool from the target site with a biopsy specimen therein. In some embodiments, the fluid conduit 12 can have a larger-diameter lumen extending
therethrough for biopsy purposes, with smaller fluid lumens formed therearound.

[0044] The device 10 can be used to deliver a drug-containing fluid under positive pressure to a target tissue region. FIG. 4 illustrates an exemplary method for convection-enhanced delivery of a drug to target tissue 40 in a patient's brain. After appropriate site preparation and cleaning, a tissue opening can formed through the patient's scalp and skull to expose the brain tissue 40. Before or after forming the tissue opening, a pedestal can

optionally be mounted to the patient to support the device10 while it is inserted, which can be particularly useful inlong-term implantations.

[0045] The device 10 can optionally be coupled to a cannula (not shown) with a microfabricated interface for
⁵⁰ mating with the device 10. Any of a variety of cannulas can be used, including standard cannulas configured to mate to a stereotactic frame in guided surgery. In some embodiments, the cannula can include a flexible catheter suitable for extended (e.g., 30 day) implantation. The
⁵⁵ catheter can be about 15 cm long and about 2 cm in diameter. The cannula can include a tubing portion that is approximately 6 feet in length with connectors for fluid and biosensor interface at the proximal end.

[0046] The device 10 can be advanced through the tissue opening and into the brain tissue 40. As shown, the tissue-receiving space 18 can be configured to compress or pinch tissue received therein as the device 10 is advanced through the tissue 40. Tissue compressed by the tissue-receiving space 18 can form a seal that reduces proximal backflow of fluid ejected from the outlet 20 of the fluid conduit 12 beyond the tissue-receiving space 18. In particular, as fluid ejected from the outlet 20 of the fluid conduit 12 flows back proximally between the exterior surface of the fluid conduit 12 and the surrounding tissue 40, it encounters a shoulder of tissue 38 that is compressed into the tissue-receiving space 18. Compression of the tissue 38 against the walls of the tissuereceiving space 18 forms a seal that resists flow of the fluid further in the proximal direction, thereby reducing or preventing undesirable backflow of injected fluid away from the target region of the tissue.

[0047] As explained above, the device 10 can include a support scaffold to facilitate penetration through the brain tissue towards the target region. One or more radiopaque markers can be included in the device 10 to permit radiographic imaging (e.g., to confirm proper placement of the device 10 within or in proximity to the target tissue). In embodiments in which a degradable scaffold is used, the scaffold can degrade shortly after insertion to leave behind only the fluid conduit 12 and outer sheath 14. In some embodiments, the fluid conduit 12 and/or the sheath 14 can be flexible to permit the device 10 to move with the brain tissue 40 if the brain tissue 40 shifts within the skull. This can advantageously prevent localized deformation of brain tissue adjacent to the device 10 that might otherwise occur with a rigid device. Such deformation can lead to backflow of the pressurized fluid along the surface of the device, undesirably preventing the fluid from reaching the target tissue.

[0048] Once the device 10 is positioned within or adjacent to the target tissue, injected media (e.g., a drugcontaining fluid) can be supplied under positive pressure to the device 10 through its fluid inlet port(s). The injected media then flows through the fluid conduit 12 and is expelled under pressure from the outlet port(s) 20 in the target region of tissue. The delivery profile can be adjusted by varying parameters such as outlet port size, outlet port shape, fluid conduit size, fluid conduit shape, fluid supply pressure, fluid velocity, etc. In some embodiments, the device 10 can be configured to deliver fluid at a flow rate between about 5 µl per minute and about 20 µl per minute. In some embodiments, the device 10 can be configured to deliver 50-100 μ l per minute per channel, and each channel can be configured to support greater than 100 psi of pressure.

[0049] In some embodiments, prior to injecting the drug-containing fluid, a gel or other material can be injected through the device 10 to augment the tissue seal. For example, a sealing gel can be injected through the device 10 and allowed to flow back along the exterior of the device, filling and sealing any voids that may exist

between the device and the surrounding tissue, particularly within the tissue-receiving recess 18. Exemplary sealing materials include cyanoacrylate, protein glues, tissue sealants, coagulative glues (e.g., fi-5 brin/thrombin/protein based coagulative glues), and materials such as those disclosed in U.S. Publication No. 2005/0277862, filed on June 9, 2004, entitled "SPLITA-BLE TIP CATHETER WITH BIORESORBABLE ADHE-SIVE," the entire contents of which are incorporated here-10 in by reference.

[0050] It will be appreciated from the foregoing that the methods and devices disclosed herein can provide convection-enhanced delivery of functional agents directly to target tissue within a patient with little or no backflow.

¹⁵ This convection-enhanced delivery can be used to treat a broad spectrum of diseases, conditions, traumas, ailments, etc. The term "drug" as used herein refers to any functional agent that can be delivered to a human or animal patient, including hormones, stem cells, gene ther-

²⁰ apies, chemicals, compounds, small and large molecules, dyes, antibodies, viruses, therapeutic agents, etc.
 [0051] In some embodiments, central-nervous-system (CNS) neoplasm can be treated by delivering an antibody (e.g., an anti-epidermal growth factor (EGF) receptor

²⁵ monoclonal antibody) or a nucleic acid construct (e.g., ribonucleic acid interference (RNAi) agents, antisense oligonucleotide, or an adenovirus, adeno-associated viral vector, or other viral vectors) to affected tissue. Epilepsy can be treated by delivering an anti-convulsive

³⁰ agent to a target region within the brain. Parkinson's disease can be treated by delivering a protein such as glial cell-derived neurotrophic factor (GDNF) to the brain. Huntington's disease can be treated by delivering a nucleic acid construct such as a ribonucleic acid interfer-

ence (RNAi) agent or an antisense oligonucleotide to the brain. Neurotrophin can be delivered to the brain under positive pressure to treat stroke. A protein such as a lysosomal enzyme can be delivered to the brain to treat lysosomal storage disease. Alzheimer's disease can be
treated by delivering anti-amyloids and/or nerve growth factor (NGF) under positive pressure to the brain. Amyotrophic lateral sclerosis can be treated by delivering a protein such as brain-derived neurotrophic factor (BDNF) or ciliary neurotrophic factor (CNTF) under positive pressure

⁴⁵ sure to the brain, spinal canal, or elsewhere in the central nervous system. Chronic brain injury can be treated by delivering a protein such as brain-derived neurotrophic factor (BDNF) and/or fibroblast growth factor (FGF) under positive pressure to the brain.

50 [0052] It will be appreciated that use of the devices disclosed herein and the various associated treatment methods is not limited to the brain of a patient. Rather, these methods and devices can be used to deliver a drug to any portion of a patient's body, including the spine. By 55 way of further example, balance or hearing disorders can be treated by injecting a drug-containing fluid directly into a portion of a patient's ear. Any of a variety of drugs can be used to treat the ear, including human atonal gene.

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The methods and devices disclosed herein can also be used to deliver therapeutics (such as stem cells) to a fetus or to a patient in which the fetus is disposed. The methods and devices disclosed herein can be used to treat a cavernous malformation, for example by delivering one or more antiangiogenesis factors thereto.

[0053] Any of the various treatments described herein can further include delivering a cofactor to the target tissue, such as a corticosteroid impregnated in the device, a corticosteroid coated onto the device, and/or a propagation enhancing enzyme. In addition, any of the various treatments described herein can further include longterm implantation of the device (e.g., for several hours or days) to facilitate long-term treatments and therapies. [0054] A number of variations on the device 10 are set forth below. Except as indicated, the structure and operation of these variations is identical to that of the device 10, and thus a detailed description is omitted here for the sake of brevity.

[0055] In some embodiments, the device 10 can include a plurality of tissue-receiving spaces 18. FIG. 5 illustrates an embodiment with a first tissue-receiving space 18A and a second tissue-receiving space 18B. As shown, a first outer sheath 14A is disposed over the fluid conduit 12 to define the first tissue-receiving space 18A. A second outer sheath 14B is disposed over the first outer sheath 14A to define the second tissue-receiving space 18B. Specifically, the second tissue-receiving space 18B is formed between an exterior surface of the first outer sheath 14A and an interior surface of the distal end 16B of the second outer sheath 14B. While two tissue-receiving spaces are shown, it will be appreciated that any number of tissue-receiving spaces can be provided (e.g., three, four, five, or more) by adding additional sheath layers. A single sheath layer can also be configured to provide multiple tissue-receiving spaces, for example by forming the sheath layer with one or more stepped regions, each stepped region defining a tissue-receiving space therein. Multi-stage devices such as that shown in FIG. 5 can provide additional sealing regions proximal to the distal-most, primary sealing region. The provision of these secondary, tertiary, etc. sealing regions can augment the primary seal or act as a backup in case the primary seal is compromised.

[0056] As shown in FIGS. 6A-6C, the internal wall of the distal end 16 of the outer sheath 14 can be shaped to alter the dimensions of the tissue-receiving space 18 and the type of seal provided when tissue is compressed therein. FIG. 6A illustrates a device 100 in which the interior surface of the distal end 116 of the sheath 114 has a concave curvature. FIG. 6B illustrates a device 200 in which the interior surface of the distal end 216 of the sheath 214 is conical. FIG. 6C illustrates a device 300 in which the interior surface of the distal end 316 of the sheath 314 has a convex curvature. These configurations can provide for a sharper leading edge at the periphery of the sheath as compared with the cylindrical tissue-receiving space 18 of the device 10, and can increase

the amount of tissue compressed into or pinched/pinned by the tissue-receiving space, as well as the degree of compression. A more-robust seal can thus be obtained in some instances using the configurations of FIGS. 6A-

6C. It should be noted, however, that even in the case of a cylindrical tissue-receiving space, the leading edge of the sheath can be sharpened to deflect tissue into the tissue-receiving space and thereby form a better seal. The size and shape of the tissue-receiving space can be

¹⁰ selected based on a variety of parameters, including the type of tissue in which the device is to be inserted. In embodiments with a plurality of tissue-receiving spaces, each of the tissue receiving spaces can have the same configuration (e.g., all cylindrical, all conical, all convex,

¹⁵ or all concave). Alternatively, one or more of the plurality of tissue-receiving spaces can have a different configuration. Thus, for example, one or more tissue-receiving spaces can be cylindrical while one or more other tissue receiving spaces are convex.

20 [0057] The tissue-receiving recesses of the devices disclosed herein can include various surface features or treatments to enhance the seal formed between the device and the surrounding tissue or gel. For example, the tissue-receiving recesses can be coated with a biocom-

²⁵ patible adhesive or can have a textured surface to form a tighter seal with the tissue or gel.

[0058] FIG. 7 illustrates an exemplary embodiment of a CED device 400 that generally includes a fluid conduit in the form of a micro-tip 412 and an outer sheath 414. The micro-tip 412 includes a substrate 442, which can be formed from a variety of materials, including silicon. The substrate 442 can have any of a variety of crosssectional shapes, including a square or rectangular

cross-section as shown. One or more fluid channels 444
can be formed on the substrate 442. The fluid channels 444 can be formed from a variety of materials, including parylene. Additional details on the structure, operation, and manufacture of microfabricated tips such as that shown in FIG. 7 can be found in U.S. Publication No.

40 2013/0035560, filed on August 1, 2012, entitled "MULTI-DIRECTIONAL MICROFLUIDIC DRUG DELIVERY DE-VICE," the entire contents of which are incorporated herein by reference.

[0059] The outer sheath 414 can be disposed coaxially 45 over the micro-tip 412 so as to form a tissue-receiving space 418 therebetween. In some embodiments, the micro-tip 412 can have a substantially rectangular exterior cross-section and the outer sheath 414 can have a substantially cylindrical interior cross-section. In other em-50 bodiments, the micro-tip 412 and the outer sheath 414 can have corresponding cross-sectional shapes with a clearance space defined therebetween. The proximal end of the outer sheath 414 can be coupled to a catheter 446. The catheter 446 can be rigid or flexible, or can 55 include rigid portions and flexible portions. A nose portion 448 (sometimes referred to herein as a "bullet nose" or a "bullet nose portion") can be disposed between the outer sheath 414 and the catheter 446, or can be dis-

posed over a junction between the outer sheath 414 and the catheter 446. As shown, the nose portion 448 can taper from a reduced distal diameter corresponding to the outside diameter of the sheath 414 to an enlarged proximal diameter corresponding to the outside diameter of the catheter 446. The tapered transition provided by the nose portion 448 can advantageously provide stressrelief as it can act as a smooth transition from the sheath 414 to the catheter body 446, avoiding any uneven stresses on the surrounding tissue that may create paths for fluid backflow. The nose portion 448 can be conically tapered, as shown, or can taper along a convex or concave curve. Various compound shapes can also be used that include conical portions, convex portions, and/or concave portions. The nose portion 448 can also be replaced with a blunt shoulder that extends perpendicular to the longitudinal axis of the device 400. Any of a variety of taper angles can be used for the nose portion 448. For example the nose portion 448 can taper at an angle in a range of about 10 degrees to about 90 degrees relative to the longitudinal axis of the device 400, in a range of about 20 degrees to about 70 degrees relative to the longitudinal axis of the device, and/or in a range of about 30 degrees to about 50 degrees relative to the longitudinal axis of the device. For example, the nose portion 446 can taper at an angle of approximately 33 degrees relative to the longitudinal axis of the device 400. In some embodiments, additional sheaths can be provided, e.g., as described above with respect to FIG. 5.

[0060] As shown in FIG. 8, the catheter 446 can include length markings or graduations 450 to indicate the insertion depth of the device 400. In some embodiments, the catheter 446 can be a straight rigid catheter sized and configured for acute stereotactic targeting. The catheter 446 can be formed from any of a variety of materials, including flexible materials, rigid materials, ceramics, plastics, polymeric materials, PEEK, polyurethane, etc. and combinations thereof. In an exemplary embodiment, the catheter 446 has length of about 10 cm to about 40 cm, e.g., about 25 cm. The catheter 446 can include one or more fluid lines extending therethrough. The fluid lines can be defined by the catheter body itself or can be defined by one or more inner sleeves or linings disposed within the catheter body. Any of a variety of materials can be used to form the inner sleeves or linings, such as flexible materials, rigid materials, polyimide, pebax, PEEK, polyurethane, silicone, fused silica tubing, etc. and combinations thereof.

[0061] As shown in FIG. 9, one or more standard Luer or other connectors 452 can be coupled to the proximal end of the catheter 446 to facilitate connection with a fluid delivery system of the type shown in FIG. 3. In the illustrated embodiment, the system 400 includes two connectors 452, one for each of the two fluid channels formed in the catheter 446 and the micro-tip 412. It will be appreciated, however, that any number of fluid channels and corresponding proximal catheter connectors can be provided. The system 400 can also include a collar 454 disposed over the catheter 446 to act as a depth stop for setting the desired insertion depth and preventing overinsertion. The collar 454 can be longitudinally slidable with respect to the catheter 446 and can include a thumb screw 456 for engaging the catheter to secure the collar in a fixed longitudinal position with respect thereto. The system 400 can also include a tip protector 458 for preventing damage to the micro-tip 412 during insertion into stereotactic frame fixtures. Exemplary tip protectors are

 disclosed in U.S. Patent Application No. 14/306,925, filed on June 17, 2014, entitled "METHODS AND DEVICES FOR PROTECTING CATHETER TIPS AND STEREO-TACTIC FIXTURES FOR MICROCATHETERS," the entire contents of which are incorporated herein by refer ence.

[0062] As shown in FIG. 10, the system 400 can include a length of extension tubing 460 to provide a fluid pathway between the proximal connectors 452 of the catheter 446 and a fluid delivery system of the type shown in FIG. 3.
²⁰ In the illustrated embodiment, dual-channel peel-away extension lines 460 are shown. In an exemplary method of using the system 400, an incision can be formed in a

of using the system 400, an incision can be formed in a patient and the catheter 446 can be inserted through the incision and implanted in a target region of tissue (e.g., a region of the patient's brain or central nervous system).

The catheter 446 can be left in the target region for minutes, hours, days, weeks, months, etc. In the case of a flexible catheter 446, the proximal end of the catheter can be tunneled under the patient's scalp with the proximal connectors 452 extending out from the incision. The catheter 446 can be inserted through a sheath to keep the catheter stiff and straight for stereotactic targeting. Alternatively, or in addition, a stylet can be inserted

through the catheter to keep the catheter stiff and straight
for stereotactic targeting. In some embodiments, the stylet can be inserted through an auxiliary lumen formed in the catheter such that the primary fluid delivery lumen(s) can be primed with fluid during catheter insertion. Thus, in the case of a catheter with first and second fluid
lumens, a third lumen can be included for receiving the

stylet. [0063] FIG. 11 is a close-up view of the exemplary micro-tip 412. As shown, the micro-tip 412 generally includes a central body portion 462 with first and second 45 legs or tails 464 extending proximally therefrom and a tip portion 466 extending distally therefrom. First and second microfluidic channels 444 are formed in or on the micro-tip 412 such that they extend along the proximal legs 464, across the central body portion 462, and down 50 the distal tip portion 466. The channels 444 can each include one or more fluid inlet ports (e.g., at the proximal end) and one or more fluid outlet ports (e.g., at the distal end). As noted above, additional details on the structure, operation, and manufacture of microfabricated tips such 55 as that shown in FIG. 11 can be found in U.S. Publication No. 2013/0035560, filed on August 1, 2012, entitled "MULTI-DIRECTIONAL MICROFLUIDIC DRUG DELIV-ERY DEVICE," the entire contents of which are incorpo-

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rated herein by reference.

[0064] Systems and methods for manufacturing and/or assembling the CED device 400 are shown in FIGS. 12-15. Generally speaking, after the micro-tip 412 is fabricated, it can be positioned in a molding or casting system to couple the one or more sheaths 414 to the micro-tip, to form the nose portion 448, and/or to couple fluid lines in the catheter 446 to the fluid channels 444 of the micro-tip.

[0065] FIG. 12 illustrates an exemplary embodiment of a molding system 500. The system 500 includes a base plate 502 with a cradle 504 in which a proximal portion of the catheter 446 is supported. Upper and lower mold blocks 506, 508 are coupled to the base plate 502 by a clamping block 510 with one or more screws 512. The screws 512 can be tightened to lock the mold blocks 506, 508 in position during an injection process and can be removed to allow the mold blocks to be opened for insertion or removal of the CED device components. The system 500 also includes an inlet port 514 through which flowable material can be injected, pumped, etc. into the mold.

[0066] As shown in FIGS. 13-15, the lower mold block 508 includes a recess in which the lower half of the catheter body 446 can be disposed and a recess in which the lower half of the sheath 414 can be disposed. A mold cavity 516 which is substantially a negative of the lower half of the nose portion 448 is formed between the recesses. The recesses can be sized such that the catheter body 446 and the sheath 414 form a seal with the mold block 508 to prevent flowable material injected into the mold cavity 516 from escaping. One or more injection ports or channels 514 are formed in the mold block 508 to allow flowable material to be injected into the cavity 516. While not shown, it will be appreciated that the upper mold block 506 is configured in a similar manner to the lower mold block 508, with recesses that can receive the upper halves of the catheter body 446 and the sheath 414 and a mold cavity 516 which is substantially a negative of the upper half of the nose portion 448.

[0067] In use, the micro-tip 412 is positioned such that the proximal legs 464 are disposed within respective fluid lines formed in the catheter body 446 and such that the distal tip portion 466 of the micro-tip is positioned within the inner lumen of the sheath 414. As noted above, in some embodiments, the catheter fluid lines can be formed by inner linings (e.g., fused silica tubes) encased in an outer housing (e.g., a ceramic housing) that defines the catheter body 446. The inner linings can prevent leaks and hold the catheter body 446 together in the event that the outer housing is cracked or damaged. The micro-tip 412, catheter body 446, and sheath 414 are sandwiched between the upper and lower mold blocks 506, 508 and a flowable material is injected through the mold channels 514 to form the nose portion 448 within the mold cavity 516, and to couple the fluid lines in the catheter 446 to the fluid channels 444 of the micro-tip. Exemplary flowable materials include UV resins, polymers such as polyurethanes, acrylics, PTFE, ePTFE, polyesters, and so forth.

[0068] The flowable material can be injected at low rates to fill the cavity 516. In embodiments in which UV resin is used, the upper and lower mold blocks 506, 508

can be made of a clear material to allow UV light to cure the UV resin. As the UV resin is injected into the micromold cavity 516, it can start to wick/flow up over the microtip tails 464 and under the fluid lines that sit over the tails.

Once the resin flows into the fluid lines, it can be flashed with UV light to "freeze" it in place and avoid wicking/flowing too much (and not completely encapsulating the tails 464 and the inlet holes on the tips of the tails). After the material cures, the mold blocks 506, 508 can be sepa-

¹⁵ rated and the CED device 400 can be removed from the molding system 500.

[0069] It will be appreciated that the above systems and methods can be varied in a number of ways without departing from the scope of the present disclosure. For example, the molding process can be used only for cou-

pling the fluid lines, and the bullet nose portion can be formed using a different process once the fluid connections are made. Also, while wicking is described herein as the mechanism by which the fluid line bonds are

formed, it will be appreciated that these bonds can also be controlled by fill pressure, timing, and other molding variables. The bullet nose can be over-molded directly onto the micro-tip. While an exemplary micro-tip and an exemplary catheter body are shown, it will be appreciated that the micro-molding methods and devices disclosed herein can be used with any of a variety of tips and/or patheters.

catheters.
[0070] Alternative systems and methods for manufacturing and/or assembling the CED device 400 are shown
³⁵ in FIGS. 16-21. As shown in FIGS. 16-19, the bullet nose and the one or more sheaths or over tubes can be as-

sembled separately using an over-molding process as described below to create a molded part 470. To assemble the system 400, the proximal legs 464 of the microtip 412 are inserted into the distal end of the catheter

body 446 (e.g., by inserting each leg into a respective lining disposed within an outer catheter housing). A flowable material (e.g., an adhesive such as a UV curable adhesive) can then be applied to the legs 464 to bond

45 the fluid channels on each leg to a corresponding fluid line of the catheter body 446. The molded part 470 can then be slid over the distal end of the micro-tip 412 such that the central body portion 462 of the micro-tip is disposed in a hollow interior of the molded part and such 50 that the tip portion 466 of the micro-tip extends through the molded part and protrudes from the distal end thereof. [0071] The molded part 470 can include a shoulder that defines a proximal male portion 472 that mates into a female counterbore 474 formed in the distal tip of the 55 catheter body 446. Alternatively, the catheter body 446 can define a male portion and the molded part 470 can include a female counterbore. It will also be appreciated that other ways of mating the catheter body 446 to the

molded part 470 can be used, such as a threaded interface, a snap-fit interface, a key and slot interface, or any other interlocking interface that provides alignment and/or overlap between the molded part and the catheter body. In some embodiments, the counterbore 474 can be formed by machining a recess into the distal end of a ceramic catheter body 446. The inner linings of the catheter can then be inserted into the ceramic outer housing such that the terminal ends of the inner linings are flush with the floor of the counterbore 474. The molded part 470 can be attached to the catheter body 446 using a flowable material (e.g., a UV adhesive), which can be applied to the counterbore 474 and/or the male portion 472 prior to assembling the components or which can be applied through one or more openings 476 formed in the sidewall of the molded part after the components are assembled or dry fit. The flowable material is allowed to cure to form a seal between the fluid lines and to secure the components of the CED device 400 to one another. [0072] An exemplary over-molding system 600 for forming the bullet nose and coupling the bullet nose to one or more over-tubes to form the molded part 470 is shown in FIG. 20. The molding system 600 includes upper and lower plates 602, 604 that sandwich the one or more over-tubes and together define a negative of the bullet nose. The plates 602, 604 also define a plug for forming the bullet nose as a hollow structure which can later be filled as described above during final assembly. A flowable material can be injected through injection ports 606 formed in the plates 602, 604 using a syringe or pump to form the hollow bullet nose over the one or more over-tubes. In some embodiments, the flowable material is a hot resin injected under pressure which forms a strong hold with the over-tube upon curing. The over-tube can be formed from any of a variety of materials, including fused silica tubing.

[0073] A scale drawing of an exemplary molded part 470 is shown with representative dimensions in FIG. 21. Any of the nose portions and/or sheaths described herein can be formed to the same or similar external dimensions. Unless otherwise indicated, the dimensions shown in FIG. 21 are specified in inches.

[0074] FIGS. 22-23 illustrate exemplary results of a gel study conducted by infusing dye through a CED device of the type described herein having first and second fluid channels into a gel designed to simulate tissue. As shown in FIG. 22, little or no backflow occurs at flowrates of 5, 10, and 12 µL/min (total flowrate of both channels combined). As shown in FIG. 23, a flowrate of 5 µL/min resulted in a uniform distribution of the dye over time with little or no backflow.

[0075] FIGS. 24-29 illustrate exemplary results of an animal study conducted using an in-vivo pig model in which multiple anatomies were infused using CED devices of the type described herein. Little or no backflow along the catheter track was observed at flow rates which are much higher than typical clinical flow rates for CED. The study demonstrated the capability to infuse small,

medium, and large molecules using CED devices of the type disclosed herein, and confirmed the functionality of independent flow channels. No blockages or introduction of air bubbles occurred during a multi-hour acute infusion.

- The device was found to be compatible with magnetic 5 resonance imaging and other stereotactic surgical procedures. No leaks, bond breakages, or other catheter issues were observed.
- [0076] As shown in FIG. 24, when inserted into a pig 10 brain, the ceramic catheter body and the bullet nose appear as a thick black line in a magnetic resonance (MR) image. Infused gadolinium (Gd) appears as a bright cloud in the MR image. The micro-tip is not readily visible in the MR image due to its small size.

15 [0077] FIG. 25 illustrates a series of MR images showing infusion of gadolinium into white matter of a pig's brain at flow rates of 1, 3, 5, 10, and 20 µL/min. As shown, no backflow of infusate occurs along the ceramic catheter shaft track. When the infusion cloud becomes too large,

20 the infusate overflows into surrounding anatomy, rather than flowing back along the catheter track, highlighting the capability for the system to reduce or prevent backflow. While flow rates of up to 20 μ L/min are shown, it is expected that similar results would be obtained for flow

25 rates of 30 μ L/min or more. These higher flow rates could not be tested during the animal study because the subject brain(s) became saturated with gadolinium.

[0078] FIG. 26 illustrates a series of MR images showing infusion of gadolinium into the thalamus of a pig's 30 brain at flow rates of 1, 3, 5, 10, and 20 $\mu L/min.$ As shown, no backflow of infusate occurs along the ceramic catheter shaft track. While there is slight backflow across the bullet nose at approximately 20 µL/min, this is a flowrate that is significantly higher than typical clinical CED flowrates (generally about 5 µL/min).

[0079] FIG. 27 illustrates a series of MR images showing infusion of gadolinium into the putamen of a pig's brain at flow rates of 1, 2, 5, 10, and 15 µL/min. As shown, no backflow of infusate occurs along the ceramic catheter shaft track as the infusate stays spherical throughout the

ramped infusion. [0080] The above-described backflow study showed that there is minimal backflow along the catheter shaft at high flow rates (up to 20µL/min for white matter,

45 $5\text{-}20\mu L/\text{min}$ for the thalamus, and $5\text{-}15\mu L/\text{min}$ for the putamen). These flow rates are much higher than typical clinical CED flow rates (e.g., about 5 µL/min). The determination as to whether backflow occurred was made using a 3D analysis of the infusion, not solely based on the 50 MR images included herein. In a total of eleven infusions conducted in various anatomies, zero incidences of back-

flow were observed.

[0081] FIG. 28 illustrates a series of MR images showing infusion of gadolinium into the white matter of a pig's brain at a flow rate of 5 µL/min after infusion periods of 1, 9, 16, 24, and 50 minutes. The lower set of images includes a distribution overlay. As shown, a uniform distribution with no backflow is observed even for long-du-

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ration infusions and when a large volume of infusate is delivered. Similar results were observed in infusions into the thalamus and putamen of the pig's brain.

[0082] FIG. 29 illustrates an MR image and an in vivo imaging system (IVIS) image of the thalamus of a pig's brain when a CED device of the type described herein is used to simultaneously infuse galbumin (gadolinium-labeled albumin laced with europium) through a first fluid channel and IVIS dye through a second fluid channel. As shown, the two different infusates were successfully infused from the two independent channels. A uniform distribution of the two infusates indicates mixing at the tip outlet as desired. No evidence of subarachnoid leakage was observed. This demonstrates that the system can be used to deliver Gd tracer and a drug or other molecule while monitoring the Gd tracer under MR to monitor the distribution of the drug or other molecule.

[0083] FIGS. 30-31 illustrate comparisons between measurements taken with CED devices of the type described herein and simulated measurements for a traditional 0.3 mm catheter. As shown in FIG. 30, CED devices of the type described herein achieve a more uniform concentration of infused colloidal Gd in white matter than traditional 0.3 mm catheters. As shown in FIG. 31, when using CED devices of the type described herein, extracellular expansion of white matter tissue is confined to the tip area by the bullet nose and tube-step, which prevents backflow along the catheter track. With traditional 0.3 mm catheters, on the other hand, increased extracellular expansion occurs along the catheter track due to the infusion pressure and backflow results.

[0084] The above-described infusion studies showed that 150 μL of infusate could be delivered into white matter and thalamus with no backflow along the catheter track. It also showed that the concentration profile of infusate distribution in tissue was within typical ranges for intraparenchymal drug delivery. Successful colloidal Gd (large molecule 30-50 nm) infusion was also demonstrated.

[0085] FIG. 32-36 illustrate an exemplary embodiment of a delivery and/or monitoring system 700 which can be used with any of the catheters or CED devices disclosed herein. As shown, the system 700 can include a crosscutaneous or percutaneous access device 702, a trunk line 704, a manifold 706, one or more branch lines 708, one or more filters 710, a skull anchor 712, and one or more microfluidic catheters or CED devices 714 (shown in FIG. 36). The system 700 is configured for long-term implantation beneath the skin 716 of a patient, with the catheters 714 extending into the brain, spinal column, or other target region of the patient and the access device 702 extending at least partially through the skin. In use, fluids containing drugs or other therapeutic agents can be supplied through the exposed portion of the access device 702 and delivered to the target site within the patient, e.g., via convection-enhanced delivery. In addition, electrical connections can be made through the access device 702 to apply energy to one or more electrodes on

the catheters 714 or to read sensor information from one or more sensors on the catheters 714.

- [0086] As shown in greater detail in FIG. 35, the access device 702 is configured to facilitate fluid communication
 ⁵ between one or more fluid lumens of the trunk line 704 and one or more extracorporeal fluid lumens, fluid sources, pumps, filters, and so forth. The access device 702 can be a bio-feedback and delivery access device. In the illustrated embodiment, the access device 702 includes
- ¹⁰ eight female ports 720 through which fluid can be supplied to eight independent lumens extending through the trunk line 704. The access device 702 can also include electrical connections (e.g., pins, receptacles, contacts, etc.) 722 for coupling extracorporeal electrical conduc-

¹⁵ tors to implanted leads (e.g., sensor or electrode leads). The access device 702 can be positioned just behind the patient's ear as shown, or in any other location on the patient's skin. The access device 702 can include various features to reduce the risk of infection and improve the

²⁰ long term viability of the system 700. For example, the access device 702 can include surface features to promote tissue ingrowth to form a better seal with the surrounding skin. The access device 702 can also be coated with an antibacterial agent.

25 [0087] Referring again to FIG. 32, the trunk line 704 can extend from the access device 702 to the manifold 706. The trunk line 704 can include any number of independent fluid lumens (e.g., 1, 2, 4, 8, 16, etc.) extending therethrough. In the illustrated embodiment, the trunk line 30 includes eight independent fluid lumens. The trunk line 704 can also include one or more electrical conductors 724 coupled to an exterior thereof, disposed within an inner lumen thereof, or embedded in a wall thereof. The electrical conductors 724 can be coupled to the access 35 device 702 and to downstream components to provide a conductive path between the access device and one or more sensors or electrodes of the catheters 714. As the trunk line 704 includes both fluid lumens and electrical conductors, it can be considered a dual-communicating 40 line.

[0088] The manifold 706, shown in greater detail in FIG. 33, includes at least one input port and a plurality of output ports, and is configured to route fluid lumens extending through the trunk line 704 to fluid lumens extending through one or more branch lines 708. In the illustrated embodiment, the manifold 706 divides the eight fluid lumens of the trunk line 704 into four lumens in each of the two branch lines 708. The manifold 706 can also route electrical conductors 724 of the trunk line to corresponding electrical conductors 724 of the branch

lines 708. The low-profile and contoured shape of the manifold 706 can advantageously reduce tissue irritation and patient discomfort during long-term implantation. The manifold 706 can be fixedly mounted to the patient's skull 718, for example using bone screws or anchors.

[0089] The branch lines 708 can include in-line filters or bio-filters 710 configured to remove air, gas, bacteria, and/or particulates from fluid passing through the system

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700 before such contaminants enter the patient's brain or other target treatment area. The branch lines 708 and filters 710 can also include electrical conductors 724 for completing a conductive path between the access device 702 and one or more sensors or electrodes of the catheters 714.

[0090] As shown in FIG. 34, the branch lines 708 extending out of the filters 710 can be secured to the skull anchor or burr hole adapter 712, which in turn can be securely mounted to the patient's skull 718 (e.g., using bone screws or anchors). The skull anchor 712 can be disposed over first and second burr holes formed in the patient's skull, through which microfluidic catheters or CED devices 714 coupled to the branch lines 708 can extend into a target treatment site (e.g., within the patient's brain). In some embodiments, a plurality of catheters 714 can extend from the skull anchor 712 through a single burr hole. As with the manifold 706, the lowprofile, ergonomically-efficient, small form-factor, and contoured shape of the skull anchor 712 can advantageously reduce tissue irritation and patient discomfort during long-term implantation and make the entire system 700 generally less intrusive. The skull anchor 712 can also include electrical conductors for completing a conductive path between the access device 702 and one or more sensors or electrodes of the catheters 714. Alternatively, electrical conductors in the catheters 714 can be coupled directly to the electrical conductors 724 in the branch lines 708.

[0091] Each branch line 708, which includes four independent fluid lumens in the illustrated embodiment, can be coupled to a pair of microfluidic catheters 714. Each of the catheters 714 can include first and second discrete fluid channels. It will be appreciated that, given the dimensions of the catheters 714, which can be microfabricated catheters or CED devices of the type disclosed above, it is possible to insert a plurality of catheters through a single burr hole. This can desirably reduce the number of burr holes that must be formed in order to carry out the desired treatment. As shown in FIG. 36, the catheters 714 can be positioned such that one or more fluid outlet ports formed in the fluid channels of the catheters are disposed in proximity to a target treatment site within the patient. It will be appreciated that the trunk line 704, the branch lines 708, and the catheters 714 can include any number of fluid lumens or channels, and that the specific numbers discussed herein are merely exemplary.

[0092] Any of a variety of catheters can be used with the system 700, including those described above. For example, the catheters 714 can include bullet nose and tube-over-tube features which can advantageously reduce backflow of infusate along the exterior of the catheter.

[0093] By way of further example, FIG. 37 illustrates an exemplary catheter 800 that can be used independently or with the system 700. As shown, the catheter 800 generally includes one or more (e.g., first and second) fluid conduits 802 and an elongate support scaffold 804 to provide structural rigidity to the device and facilitate insertion into the target tissue. The fluid conduits 802 can be formed directly on the support scaffold 804 or on an intermediate substrate (not shown) to which the support scaffold is coupled. To assist with tissue penetration and navigation, the distal end of the support scaffold 804 and/or the substrate can be tapered, pointed, and/or sharpened. In some embodiments, the support scaffold

10 804 and/or the substrate can be provided with a rounded atraumatic tip so as to facilitate insertion through tissue without causing trauma to the tissue.

[0094] The support scaffold 804 can be rigid or semirigid and can be formed from a degradable thermoplastic

¹⁵ polymer, for example, a degradable thermoplastic polyester or a degradable thermoplastic polycarbonate. In some embodiments, the support scaffold 804 can be formed from poly(lactic-co-glycolic acid) (PLGA) and can be configured to biodegrade within the target tissue. This

20 can advantageously eliminate the need to remove the support scaffold 804 once the catheter 800 is positioned within target tissue, thereby avoiding the potential to disrupt the positioning of the catheter. In some embodiments, the scaffold 804 can biodegrade within the target 25 tissue leaving behind only the one or more fluid conduits.

5 tissue, leaving behind only the one or more fluid conduits 802, which can be flexible and can adapt to natural movement of the target tissue.

[0095] Any of a variety of other materials can also be used to form the support scaffold 804, including silicon 30 or various ceramics, metals, and plastics known in the art. The scaffold 804 can have a width of approximately 100 μ m to approximately 10,000 μ m and can have a length that varies depending on the target tissue (e.g., depending on the depth at which the target tissue is sit-35 uated). In one embodiment, the scaffold 804 is between 2 cm and 15 cm long. The one or more fluid lumens 802 can be formed from any of a variety of materials, including at least one of a parylene composition, a silastic composition, a polyurethane composition, a polyamide compo-40 sition, and a PTFE composition.

[0096] The catheter 800 can optionally include an outer sheath or over-tube of the type described above to form a tissue-receiving space about the perimeter or circumference of the catheter. The catheter 800 can optionally 45 include a bullet nose of the type described above. The catheter 800 can be formed using any of a variety of techniques, including the micro-fabrication methods disclosed in U.S. Publication No. 2013/0035560, filed on August 1, 2012, entitled "MULTI-DIRECTIONAL MI-50 CROFLUIDIC DRUG DELIVERY DEVICE," the entire contents of which are incorporated herein by reference. [0097] As best shown in FIG. 38, the proximal end of the catheter 800 can be coupled directly to a burr hole adapter or skull anchor 806. In the illustrated embodi-55 ment, the burr hole adapter 806 includes a distal-facing surface 808 with a recess 810 in which the proximal end of the support scaffold 804 is received. The adapter 806 also includes first and second inlet ports 812 which are

coupled to the first and second fluid lumens 802 of the catheter 800. The adapter 806 provides a smooth flow transition between the inlet ports 812 and the catheter 800, which can be oriented approximately perpendicular to one another. The adapter 806 can also include electrical conductors that provide a connection between the catheter 800 (e.g., one or more sensors or electrodes disposed on or in the catheter) and other portions of the system 700. It will be appreciated that the catheter 800 can also be used with the skull anchor 712 shown in FIG. 32.

[0098] FIG. 39 illustrates another embodiment of a catheter 900 that includes a layered or sandwich configuration. The catheter 900 is substantially the same as the catheter 800, except that it includes first and second scaffolds 904A, 904B disposed on opposite sides of the substrate 914 and the fluid lumens 902. In some embodiments, the substrate 914 can be omitted and the fluid lumens 902 can be disposed in direct contact with the upper and/or lower scaffolds 904A, 904B. The fluid lumens 802 can extend a distance D beyond the distal end of the support scaffolds 904A, 904B. In some embodiments, the distance D can be in the range of about 0.1 cm to about 15 cm. The scaffolds 904A, 904B can include ramped or tapered surfaces to define a smooth transition between the sandwich portion of the catheter and the distal tip. In some embodiments, a bullet nose can be provided as described above at the transition from the sandwich portion of the catheter to the distal tip.

[0099] A proximal end of the catheter 900 is illustrated in FIG. 40. As shown, the substrate 914 of the catheter 900 can include first and second legs 916, 918 along which first and second fluid lumens 902 extend. The fluid lumens 902 can have one or more fluid inlet ports configured to be in fluid communication with the interior of the adapter or skull anchor 806, 712 when the catheter 900 is coupled thereto. As shown in FIG. 41, the scaffolds 904A, 904B can be configured to completely degrade, leaving behind only the substrate 914 and the one or more fluid lumens 902 disposed thereon. Alternatively, as shown in FIG. 42, the substrate 914 can be omitted such that, once the scaffold degrades, all that is left behind is the one or more fluid lumens 902. In addition, the substrate 914 can also be configured to biodegrade such that the scaffolds 904A, 904B and the substrate 914 are bioabsorbed or biodegraded after implantation leaving behind only the one or more fluid lumens 902.

[0100] As noted above, any of the catheters or CED devices disclosed herein can include one or more electrodes or sensors. The electrodes can be used to deliver electrical energy to target tissue, e.g., to stimulate the target tissue or to ablate the target tissue. The sensors can be used to measure one or more parameters associated with treatment of a patient. The sensors can include temperature sensors, pH sensors, pressure sensors, oxygen sensors, tension sensors, interrogatable sensors, glutamate sensors, ion concentration sensors, carbon dioxide sensors, lactate sensors, neurotransmit-

ter sensors, or any of a variety of other sensor types, and can provide feedback to a control circuit which can in turn regulate the delivery of fluid or other treatment through the device based on one or more sensed parameters.

The sensor output can also be displayed to a user, e.g., using an electronic display device, to provide feedback for facilitating treatment decisions, etc.

[0101] FIGS. 43-44 illustrate an exemplary embodiment of a catheter micro-tip 1000 (e.g., of the type described above with respect to FIG. 7) that includes an array 1002 of sensors and/or electrodes. As shown, the micro-tip 1000 generally includes a substrate 1004, which can be formed from a variety of materials, including silicon. The substrate 1004 can have any of a variety of

¹⁵ cross-sectional shapes, including a square or rectangular cross-section as shown. One or more fluid channels 1006 can be formed in or on the substrate 1004. The fluid channels 1006 can be formed from a variety of materials, including parylene and polyamide. Additional details on

the structure, operation, and manufacture of microfabricated tips such as that shown can be found in U.S. Publication No. 2013/0035560, filed on August 1, 2012, entitled "MULTI-DIRECTIONAL MICROFLUIDIC DRUG DELIVERY DEVICE," the entire contents of which are incorporated herein by reference.

[0102] The sensor / electrode array 1002 can be disposed on a tape or ribbon 1008 which can be adhered or otherwise affixed to the substrate 1004. The array 1002 can include one or more sensors and/or one or more
 ³⁰ electrodes. Thus, the array 1002 can include only a single sensor, or can include only a single electrode. In the illustrated embodiment, the array includes eight sensors or electrodes 1010. Each sensor or electrode 1010 can include one or more electrical conductors 1012 that ex ³⁵ tend along the length of the ribbon 1008 to a proximal

connector 1014. The proximal connector 1014 can be sized, shaped, and otherwise configured to electricallycouple the electrical conductors 1012 of the ribbon 1008 to corresponding electrical conductors in the skull anchor

40 712 or the branch lines 708 of the system 700. Alternatively, or in addition, one or more of the sensors / electrodes 1010 can be wireless and can include a wireless antenna to facilitate communication and/or power transmission. In some embodiments, the fluid lumens 1006

⁴⁵ formed in the substrate 1004 can be open channels and the ribbon 1008 can define the ceiling of the fluid lumens. In other embodiments, the channels 1006 can be enclosed and the ribbon 1008 can be disposed over the top of the enclosed channels. In still other embodiments, the

ribbon 1008 can be omitted and the sensors / electrodes 1010 and accompanying electrical conductors 1012 can be formed directly on or in the channels 1006 or the substrate 1004. For example, the sensors / electrodes 1010 and the electrical conductors 1012 can be printed or otherwise formed on the substrate 1004 using a lithography or other micro-fabrication process.

[0103] As shown in FIG. 45, an exemplary catheter 1100 can include a sensor and/or an electrode 1102 dis-

posed directly within a fluid lumen 1104 of the catheter. In such embodiments, fluid flow through the catheter 1100 can be used to maintain the patency of the sensor / electrode 1102. For example, the catheter 1100 can be continuously, intermittently, or periodically flushed with fluid to clear obstructions or debris from the sensor / electrode 1102. This can advantageously allow for long-term or chronic use of the sensor / electrode 1102, e.g., in connection with the system 700 described above. In some embodiments, the catheter 1100 can include one or more dedicated patency channels used only for cleaning a sensor or electrode of the catheter. Alternatively, the primary drug or fluid delivery channels can be used also to maintain the patency of a sensor or electrode disposed therein. Flushing of the sensor or electrode can be performed continuously, intermittently, periodically, etc.

[0104] FIGS. 46-48 illustrate an exemplary catheter 1200 which is similar to the catheter 1100 shown in FIG. 45, except that a dedicated channel 1206 is formed in the substrate 1208 for housing an electrical conductor 1210 of the sensor / electrode 1202. The illustrated arrangement can facilitate easier routing of the electrical and fluid lines at the proximal end of the catheter, since the electrical conductor 1210 is routed separately from the fluid channels 1204. As best shown in FIG. 46, the catheter includes a substrate 1208 with first and second fluid lumens 1204 and a first electrical conductor channel 1206 formed therein. The first fluid lumen makes a 90 degree turn at its distal end to define a chamber 1212 in which the sensor / electrode 1202 is disposed. The first electrical conductor channel 1206 runs parallel to the fluid lumens 1204, and intersects the chamber 1212 to couple with the sensor or electrode 1202. As shown in FIG. 47, the proximal end of the micro-tip 1200 can include first and second legs 1214, 1216 on or in which the fluid lumens 1204 are formed and a third leg 1218 on or in which the electrical conductor channel 1206 is formed. Again, this can facilitate easier routing of the electrical and fluid connections. The fluid lumens 1204 and the conductor channel 1206 can be formed in the substrate 1208 or can be formed as separate structures on top of the substrate. As shown in FIG. 48, when formed in the substrate, a ribbon or lid layer 1220 can be formed on top of the channels 1204, 1206 to close the channels. Alternatively, the ribbon can be omitted and the channels can be formed as enclosed structures.

[0105] The system 700 can be used to treat any of a variety of conditions, diseases, and so forth, as described above. In addition, the system 700 can be used for long-term monitoring of one or more parameters associated with treatment of a patient.

[0106] In an exemplary method, drug-containing fluid can be introduced through the access device 702 and can flow through the lumens of the system 700 to a target treatment site within the patient. Each fluid lumen can carry a different drug or drug combination, or all of the fluid lumens can deliver the same drug or drug combina-

tion. The system 700 can remain implanted for extended periods such that the fluid delivery can take place over a period of days, weeks, months, years, etc.

[0107] In another exemplary method, electrical energy
⁵ can be delivered or introduced through the access device
702 and can flow through the electrical conductors of the system 700 to one or more electrodes to deliver energy to a target treatment site within the patient. A single catheter can be used to deliver fluid and to deliver energy,

¹⁰ either simultaneously or sequentially. In addition, a first catheter can be used to deliver energy and a second catheter can be used to deliver fluid, either simultaneously or sequentially. The first and second catheters can be substantially co-located within the patient or can be

¹⁵ disposed in entirely different regions of the patient (e.g., different regions of the patient's brain). The system 700 can remain implanted for extended periods such that the energy delivery and/or fluid delivery can take place over a period of days, weeks, months, years, etc.

20 [0108] In another exemplary method, data measured by one or more sensors can be delivered via the electrical conductors of the system to the access device 702 and an attached controller, or to an internal or external wireless antenna for communication to the controller. The

sensor data can be used to monitor various parameters, including parameters associated with treatment of the patient. A single catheter can be used for monitoring and to deliver fluid, either simultaneously or sequentially. In addition, a first catheter can be used for monitoring and
 a second catheter can be used to deliver fluid, either si-

multaneously or sequentially. Either or both of the first and second catheters, or optionally a third catheter, can be used to deliver energy to the patient. The first and second catheters can be substantially co-located within ³⁵ the patient or can be disposed in entirely different regions

the patient or can be disposed in entirely different regions of the patient (e.g., different regions of the patient's brain).
 The method can include continuously, intermittently, or periodically maintaining the patency of one or more sensors used for the monitoring, for example by flushing fluid
 across a sensor disposed in a fluid lumen of the catheter.

In some embodiments, as described above, fluid or energy delivery can be performed simultaneously with sensor monitoring and the volume, frequency, etc. of fluid or energy delivery can be controlled based on the measured

⁴⁵ sensor data. In some embodiments, the method can include delivering treatment from a first catheter disposed in a first location in the patient and monitoring using a second catheter disposed in a second location in the patient that is spaced a distance apart from the first location.

⁵⁰ In such embodiments, the monitoring catheter can be used to sense how therapy in one region of the patient is affecting other regions. For example, the movement of infusate, the spreading of viral vector, the effects of neuro-stimulation, etc. can be gauged using the remote ⁵⁵ monitoring catheter. The system 700 can remain implanted for extended periods such that the monitoring and/or fluid delivery can take place over a period of days, weeks, months, years, etc.

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[0109] In another exemplary method, an insertion scaffold of an implanted micro-catheter can be allowed to biodegrade, leaving behind only one or more fluid channels. Fluid can be delivered via the one or more fluid channels using the system 700 described above. The system 700 can remain implanted for extended periods such that the fluid delivery can take place over a period of days, weeks, months, years, etc.

[0110] In another exemplary method, tissue can be biopsied or other materials can be aspirated through the fluid lines of the system 700. For example, a vacuum pump can be coupled to the access device 702 to aspirate tissue, fluid, etc. from a treatment site adjacent a distal end of the catheter. The method can include infusing fluid through a first catheter while simultaneously aspirating fluid or tissue from a second catheter.

[0111] The devices disclosed herein can be manufactured using any of a variety of techniques. For example, the devices can be manufactured by assembling lengths of tubing over one another, by micro-machining lengths of tubing, by molding steps or nose features containing tissue-receiving spaces onto a fluid conduit, or by constructing one or more portions of the device on a substrate using a lithographic microfabrication process.

25 [0112] Additional information (e.g., CED methods and devices, as well as related manufacturing techniques, exemplary micro-tips, and exemplary catheters) are disclosed in the following references, the entire contents of each of which are hereby incorporated by reference herein:

U.S. Publication No. 2013/0035560, filed on August 1, 2012, entitled MULTI-DIRECTIONAL MICRO-FLUIDIC DRUG DELIVERY DEVICE;

U.S. Publication No. 2013/0035574, filed on August 1, 2012, entitled MICROFLUIDIC DRUG DELIVERY DEVICES WITH VENTURI EFFECT;

U.S. Publication No. 2013/0035660, filed on August 40 1, 2012, entitled MULTIDIRECTIONAL MICROFLU-IDIC DRUG DELIVERY DEVICES WITH CON-FORMABLE BALLOONS;

45 U.S. Patent Application No. 14/306,925, filed on June 17, 2014, entitled METHODS AND DEVICES FOR PROTECTING CATHETER TIPS AND STER-EOTACTIC FIXTURES FOR MICROCATHETERS;

U.S. Publication No. 2014/0171760, filed on December 18, 2013, entitled SYSTEMS AND METHODS FOR REDUCING OR PREVENTING BACKFLOW IN A DELIVERY SYSTEM;

U.S. Publication No. 2010/0098767, filed on July 31, 2009, entitled CONVECTION ENHANCED DELIV-ERY APPARATUS, METHOD, AND APPLICA-TION; and

U.S. Publication No. 2013/0046230, filed on November 7, 2012, entitled ULTRASOUND-ASSISTED CONVECTION ENHANCED DELIVERY OF COM-POUNDS IN VIVO WITH A TRANSDUCER CAN-NULA ASSEMBLY.

[0113] Although the invention has been described by reference to specific embodiments, it should be understood that numerous changes may be made within the scope of the inventive concepts described. Accordingly, it is intended that the invention not be limited to the described embodiments, but that it have the full scope defined by the language of the following claims.

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Claims

- 1. An implantable delivery system, comprising:
 - a percutaneous access device (702) through which drug-containing fluid can be delivered; a trunk line (704) having a plurality of independent fluid lumens extending therethrough, the plurality of fluid lumens being in fluid communication with a corresponding plurality of ports (720) formed in the access device (702); a manifold (706) configured to route the plurality

of fluid lumens in the trunk line (704) into a plurality of branch lines (708), each branch line (708) including a plurality of corresponding fluid lumens disposed therein and being coupled to a plurality of microfluidic catheters (714), each of the microfluidic catheters (714) having first and second discrete fluid channels and being configured to extend into the brain of a patient in use; and

a skull anchor (712) configured to be secured to the skull of the patient, the plurality of branch lines (708) and the plurality of microfluidic catheters (714) extending from the skull anchor (712).

- The system of claim 1, further comprising one or 2. more inline filters (710) disposed in the branch lines (708) and configured to remove air, gas, bacteria, or particulates from fluid flowing through the branch lines (708).
- 3. The system of claim 1, wherein the access device (702) includes one or more electrical connections for coupling extracorporeal electrical conductors to implanted electrical conductors.
- 4. The system of claim 1, wherein the trunk line (704), the manifold (706), at least one of the branch lines (708), the skull anchor (712), and at least one of the catheters (714) include electrical conductors config-

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ured to provide an electrical path between the access device (702) and a sensor or electrode of the at least one catheter.

- 5. The system of claim 1, wherein the skull anchor (712) is disposable over first and second burr holes formed in the skull of the patient such that a first catheter coupled (714) to the skull anchor (712) extends through the first burr hole and a second catheter coupled to the skull anchor (712) extends through the 10 second burr hole, or wherein the skull anchor (712) is disposable over a first burr hole such that first and second catheters coupled to the skull anchor (712) extend through the first burr hole.
- 6. The system of claim 1, wherein at least one of the catheters (714) includes a sensor.
- 7. The system of claim 6, wherein the sensor is disposed in a fluid lumen of the at least one catheter 20 (714) adjacent an outlet port of the at least one catheter (714) such that fluid flowing through the outlet port washes over the sensor.
- 25 8. The system of claim 6, wherein the at least one catheter (714) includes at least one drug delivery channel and a dedicated patency channel through which fluid can be directed to clean the sensor.
- 9. The system of claim 6, wherein the at least one cath-30 eter (714) includes a dedicated electrical conductor channel (1206) through which an electrical conductor (1210) coupled to the sensor (1202) extends, the electrical conductor channel (1206) being separate from a fluid delivery channel of the at least one cath-35 eter (714).
- 10. The system of claim 9, wherein the fluid delivery channel and the electrical conductor channel (1206) intersect at a chamber (1212) in which the sensor (1202) is disposed.
- 11. The system of claim 1, wherein at least one of the catheters (714) includes a biodegradable scaffold (804) on which one or more fluid channels are formed.
- 12. The system of claim 11, wherein the scaffold (804) is configured to biodegrade after being implanted in a patient, leaving behind only the one or more fluid 50 channels.
- **13.** The system of claim 11, wherein the scaffold (804) extends to a proximal end of the at least one catheter (714) and is coupled to the skull anchor (712).
- 14. The system of claim 11, wherein the scaffold (804) comprises an upper layer and a lower layer, and

wherein the one or more fluid channels are sandwiched between the upper and lower scaffold layers.

15. The system of claim 1, wherein at least one of the catheters (714) comprises:

> a micro-tip (412) having a proximal portion, a central portion, a distal portion, and at least one fluid channel extending along said proximal, central, and distal portions, the at least one fluid channel having an outlet port at a distal end thereof and an inlet port at a proximal end thereof:

a first outer sheath (414) disposed coaxially over the distal portion of the micro-tip such that the distal portion of the micro-tip (412) protrudes from a distal end of the first outer sheath (414); a first tissue-receiving space (418) defined between an exterior surface of the microtip (412) and an interior surface of the distal end of the first outer sheath (414);

a catheter body (446) extending proximally from the micro-tip (412) such that the at least one fluid channel of the micro-tip (412) is in fluid communication with a respective inner lumen of the catheter body (446); and

a nose portion (448) disposed over at least the central portion of the micro-tip (412) and extending between the first outer sheath (414) and the catheter body (446) such that the nose portion (448) defines an exterior surface that tapers from a reduced distal diameter corresponding to the outside diameter of the first outer sheath (414) to an enlarged proximal diameter corresponding to the outside diameter of the catheter body (446).

Patentansprüche

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1. Implantierbares Freisetzungssystem, umfassend:

eine perkutane Zugangsvorrichtung (702), durch die wirkstoffhaltige Flüssigkeit freigesetzt werden kann:

eine Hauptleitung (704) mit einer Mehrzahl von sich hindurch erstreckenden unabhängigen Flüssigkeitslumen, wobei die Mehrzahl von Flüssigkeitslumen in Flüssigkeitskommunikation mit einer entsprechenden Mehrzahl von in der Zugangsvorrichtung (702) gebildeten Öffnungen (720) steht;

einen Verteiler (706), der dafür konfiguriert ist, die Mehrzahl von Flüssigkeitslumen in der Hauptleitung (704) in eine Mehrzahl von Zweigleitungen (708) zu führen, wobei jede Zweigleitung (708) eine Mehrzahl von entsprechenden Flüssigkeitslumen beinhaltet, die darin ange-

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ordnet und an eine Mehrzahl von mikrofluidischen Kathetern (714) gekoppelt sind, wobei jeder der mikrofluidischen Katheter (714) erste und zweite diskrete Flüssigkeitskanäle aufweist und dafür konfiguriert ist, sich bei Gebrauch in das Gehirn eines Patienten zu erstrecken; und einen Schädelanker (712), der dafür konfiguriert ist, am Schädel des Patienten gesichert zu sein, wobei sich die Mehrzahl von Zweigleitungen (708) und die Mehrzahl von mikrofluidischen Kathetern (714) von dem Schädelanker (712) erstrecken.

- System nach Anspruch 1, ferner umfassend einen oder mehrere Inline-Filter (710), die in den Zweigleitungen (708) angeordnet und dafür konfiguriert sind, Luft, Gas, Bakterien oder Partikulate aus durch die Zweigleitungen (708) fließender Flüssigkeit zu entfernen.
- 3. System nach Anspruch 1, worin die Zugangsvorrichtung (702) eine oder mehrere elektrische Verbindungen zum Koppeln extrakorporaler elektrischer Leiter an implantierte elektrische Leiter beinhaltet.
- 4. System nach Anspruch 1, worin die Hauptleitung (704), der Verteiler (706), mindestens eine der Zweigleitungen (708), der Schädelanker (712) und mindestens einer der Katheter (714) elektrische Leiter beinhalten, die dafür konfiguriert sind, einen elektrischen Weg zwischen der Zugangsvorrichtung (702) und einem Sensor oder einer Elektrode des mindestens einen Katheters bereitzustellen.
- 5. System nach Anspruch 1, worin der Schädelanker (712) über ersten und zweiten im Schädel des Patienten hergestellten Trepanationslöchern anordbar ist, so dass sich ein an den Schädelanker (712) gekoppelter erster Katheter (714) durch das erste Trepanationsloch erstreckt und sich ein an den Schädelanker (712) gekoppelter zweiter Katheter durch das zweite Trepanationsloch erstreckt, oder worin der Schädelanker (712) über einem ersten Trepanationsloch anordbar ist, so dass sich erste und zweite an den Schädelanker (712) gekoppelte Katheter durch das erste Trepanationsloch erstrecken.
- **6.** System nach Anspruch 1, worin mindestens einer der Katheter (714) einen Sensor beinhaltet.
- System nach Anspruch 6, worin der Sensor in einem Flüssigkeitslumen des mindestens einen Katheters (714) neben einer Auslassöffnung des mindestens einen Katheters (714) angeordnet ist, so dass durch die Auslassöffnung fließende Flüssigkeit den Sensor überspült.
- 8. System nach Anspruch 6, worin der mindestens eine

Katheter (714) mindestens einen Wirkstofffreisetzungskanal und einen dedizierten Durchgängigkeitskanal, durch den Flüssigkeit gerichtet werden kann, um den Sensor zu reinigen, beinhaltet.

- 9. System nach Anspruch 6, worin der mindestens eine Katheter (714) einen dedizierten elektrischen Leiterkanal (1206) beinhaltet, durch den sich ein an den Sensor (1202) gekoppelter elektrischer Leiter (1210) erstreckt, wobei der elektrische Leiterkanal (1206) separat von einem Flüssigkeitsfreisetzungskanal des mindestens einen Katheters (714) ist.
- **10.** System nach Anspruch 9, worin sich der Flüssigkeitsfreisetzungskanal und der elektrische Leiterkanal (1206) an einer Kammer (1212) schneiden, in der der Sensor (1202) angeordnet ist.
- 11. System nach Anspruch 1, worin mindestens einer der Katheter (714) ein biologisch abbaubares Gerüst (804) beinhaltet, auf dem ein oder mehrere Flüssigkeitskanäle gebildet sind.
- System nach Anspruch 11, worin das Gerüst (804)
 dafür konfiguriert ist, sich nach Implantation in einem Patienten biologisch abzubauen, wobei nur der eine oder mehrere Flüssigkeitskanäle zurückgelassen werden.
 - System nach Anspruch 11, worin sich das Gerüst (804) zu einem proximalen Ende des mindestens einen Katheters (714) erstreckt und an den Schädelanker (712) gekoppelt ist.
 - **14.** System nach Anspruch 11, worin das Gerüst (804) eine obere Schicht und eine untere Schicht umfasst, und worin der eine oder mehrere Flüssigkeitskanäle sandwichartig zwischen der oberen und unteren Gerüstschicht angeordnet sind.
 - **15.** System nach Anspruch 1, worin mindestens einer der Katheter (714) Folgendes umfasst:

eine Mikrospitze (412) mit einem proximalen Abschnitt, einem zentralen Abschnitt, einem distalen Abschnitt und mindestens einem Flüssigkeitskanal, der sich entlang dem proximalen, zentralen und distalen Abschnitt erstreckt, wobei der mindestens eine Flüssigkeitskanal eine Auslassöffnung an einem distalen Ende davon und eine Einlassöffnung an einem proximalen Ende davon aufweist;

eine erste Außenhülle (414), die koaxial über dem distalen Abschnitt der Mikrospitze angeordnet ist, so dass der distale Abschnitt der Mikrospitze (412) aus einem distalen Ende der ersten Außenhülle (414) hervorsteht;

einen ersten gewebeaufnehmenden Raum

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(418), der zwischen einer Außenfläche der Mikrospitze (412) und einer Innenfläche des distalen Endes der ersten Außenhülle (414) definiert ist:

einen Katheterkörper (446), der sich proximal von der Mikrospitze (412) erstreckt, so dass der mindestens eine Flüssigkeitskanal der Mikrospitze (412) in Flüssigkeitskommunikation mit einem jeweiligen inneren Lumen des Katheterkörpers (446) steht; und

einen Nasenabschnitt (448), der über mindestens dem zentralen Abschnitt der Mikrospitze (412) angeordnet ist und sich zwischen der ersten Außenhülle (414) und dem Katheterkörper (446) erstreckt, so dass der Nasenabschnitt (448) eine Außenfläche definiert, die sich von einem reduzierten distalen Durchmesser, der dem Außendurchmesser der ersten Außenhülle (414) entspricht, zu einem vergrößerten proxi-20 malen Durchmesser, dem Außendurchmesser des Katheterkörpers (446) entspricht, konisch aufweitet.

Revendications

1. Système d'administration implantable, comprenant :

> un dispositif d'accès percutané (702) par leguel un fluide contenant un médicament peut être administré ;

> une ligne principale (704) comportant une pluralité de lumières de fluide indépendantes s'y étendant, la pluralité de lumières de fluide étant en communication fluidique avec une pluralité correspondante d'orifices (720) formés dans le dispositif d'accès (702);

> un collecteur (706), configuré pour acheminer la pluralité de lumières de fluide dans la ligne principale (704) dans une pluralité de lignes secondaires (708), chaque ligne secondaire (708) comprenant une pluralité de lumières de fluide correspondantes qui y sont disposées et qui sont accouplées à une pluralité de cathéters microfluidiques (714), chacun des cathéters microfluidiques (714) ayant des premier et second canaux de fluide distincts et étant configuré pour s'étendre dans le cerveau d'un patient lors de l'utilisation ; et

> un ancrage de crâne (712), configuré pour être fixé au crâne du patient, la pluralité de lignes secondaires (708) et la pluralité de cathéters microfluidiques (714) s'étendant à partir de l'ancrage de crâne (712).

2. Système selon la revendication 1, comprenant en outre un ou plusieurs filtres en ligne (710) disposés dans les lignes secondaires (708) et configurés pour éliminer l'air, les gaz, les bactéries ou les particules du fluide circulant dans les lignes secondaires (708).

- Système selon la revendication 1, dans lequel le dis-3. positif d'accès (702) comprend une ou plusieurs connexions électriques pour accoupler des conducteurs électriques extracorporels à des conducteurs électriques implantés.
- 10 4. Système selon la revendication 1, dans lequel la ligne principale (704), le collecteur (706), au moins une des lignes secondaires (708), l'ancrage de crâne (712) et au moins un des cathéters (714) comprennent des conducteurs électriques configurés pour 15 fournir une voie électrique entre le dispositif d'accès (702) et un capteur ou une électrode de l'au moins un cathéter.
 - 5. Système selon la revendication 1, dans lequel l'ancrage de crâne (712) peut être disposé sur des premier et second trous de trépan formés dans le crâne du patient de sorte qu'un premier cathéter (714) accouplé à l'ancrage de crâne (712) s'étende dans le premier trou de trépan et qu'un second cathéter accouplé à l'ancrage de crâne (712) s'étende dans le second trou de trépan, ou dans lequel l'ancrage de crâne (712) peut être disposé sur un premier trou de trépan de sorte que les premier et second cathéters accouplés à l'ancrage de crâne (712) s'étendent dans le premier trou de trépan.
 - 6. Système selon la revendication 1, dans lequel au moins un des cathéters (714) comprend un capteur.
- 35 7. Système selon la revendication 6, dans lequel le capteur est disposé dans une lumière de fluide de l'au moins un cathéter (714) adjacent à un orifice de sortie de l'au moins un cathéter (714) de sorte que le fluide s'écoulant par l'orifice de sortie passe sur le capteur.
 - 8. Système selon la revendication 6, dans lequel l'au moins un cathéter (714) comprend au moins un canal d'administration de médicament et un canal de perméabilité réservé par lequel un fluide peut être dirigé pour nettoyer le capteur.
 - Système selon la revendication 6, dans lequel l'au 9. moins cathéter (714) comprend un canal de conducteur électrique (1206) réservé par lequel s'étend un conducteur électrique (1210) accouplé au capteur (1202), le canal de conducteur électrique (1206) étant séparé d'un canal d'administration de fluide de l'au moins un cathéter (714).
 - 10. Système selon la revendication 9, dans lequel le canal d'administration de fluide et le canal de conducteur électrique (1206) se coupent au niveau d'une

chambre (1212) dans laquelle est disposé le capteur (1202).

- Système selon la revendication 1, dans lequel au moins un des cathéters (714) comprend un échafaudage biodégradable (804) sur lequel sont formés un ou plusieurs canaux de fluide.
- Système selon la revendication 11, dans lequel l'échafaudage (804) est configuré pour se biodégrader après avoir été implanté chez un patient, laissant derrière lui uniquement le ou les canaux de fluide.
- Système selon la revendication 11, dans lequel l'échafaudage (804) s'étend vers une extrémité ¹⁵ proximale de l'au moins un cathéter (714) et est accouplé à l'ancrage de crâne (712).
- 14. Système selon la revendication 11, dans lequel l'échafaudage (804) comprend une couche supérieure et une couche inférieure, et dans lequel le ou les canaux de fluides sont intercalés entre les couches d'échafaudage supérieure et inférieure.
- **15.** Système selon la revendication 1, dans lequel au ²⁵ moins un des cathéters (714) comprend :

une micropointe (412) comportant une partie proximale, une partie centrale, une partie distale et au moins un canal de fluide s'étendant le long 30 desdites parties proximale, centrale et distale, l'au moins un canal de fluide comportant un orifice de sortie au niveau d'une extrémité distale correspondante et un orifice d'entrée au niveau d'une extrémité proximale correspondante ; 35 une première gaine externe (414) disposée coaxialement sur la partie distale de la micropointe de sorte que la partie distale de la micropointe (412) fasse saillie par rapport à une ex-40 trémité distale de la première gaine externe (414);

un premier espace de réception de tissu (418) défini entre une surface extérieure de la micropointe (412) et une surface intérieure de l'extrémité distale de la première gaine externe (414) ; ⁴⁵ un corps de cathéter (446) s'étendant de manière proximale par rapport à la micropointe (412) de sorte que l'au moins un canal de fluide de la micropointe (412) soit en communication fluidique avec une lumière intérieure respective ⁵⁰ du corps de cathéter (446) ; et

une partie nez (448) disposée sur au moins la partie centrale de la micropointe (412) et s'étendant entre la première gaine externe (414) et le corps de cathéter (446) de sorte que la partie nez (448) définisse une surface extérieure à section croissante d'un diamètre distal réduit, correspondant au diamètre extérieur de la première gaine externe (414), à un diamètre proximal élargi, correspondant au diamètre extérieur du corps de cathéter (446).













FIG. 4









FIG. 6C









FIG. 13



FIG. 14











FIG. 22







FIG. 24









FIG. 29







| TIME (min) | 9 | 17 | 25 |
|-------------|----|----|-----|
| INFUSE (µ1) | 45 | 85 | 125 |

FIG. 31



SIMULATED TISSUE EXPANSION





FIG. 33



FIG. 34









FIG. 38





FIG. 41



FIG. 42



FIG. 43









REFERENCES CITED IN THE DESCRIPTION

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