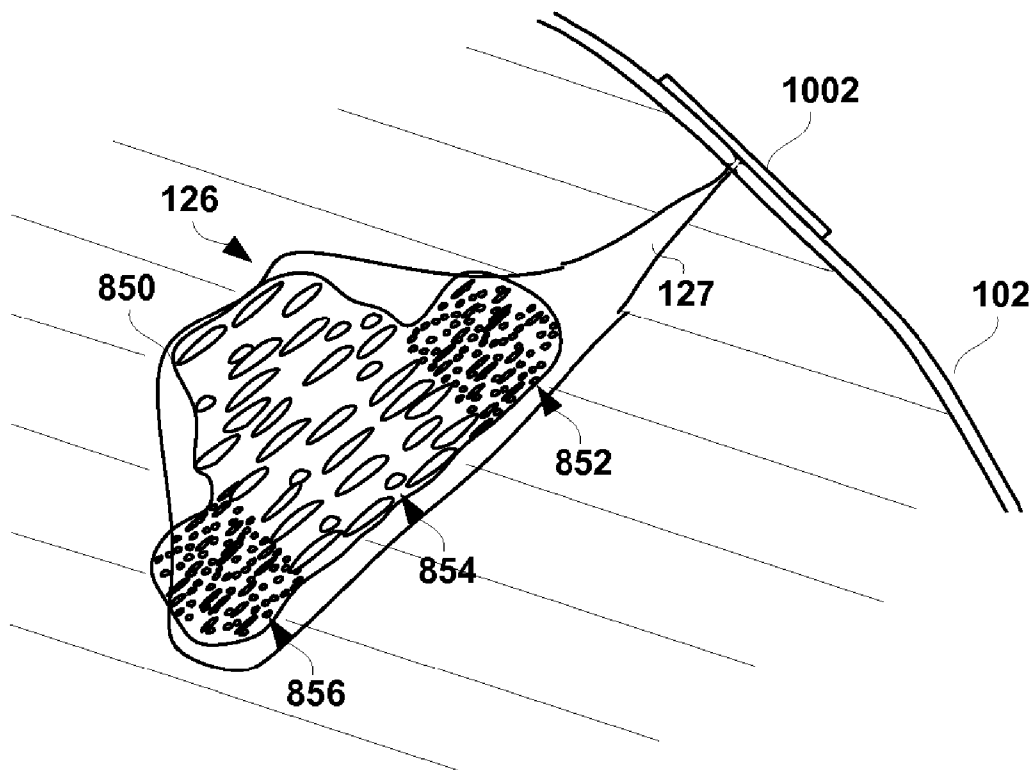


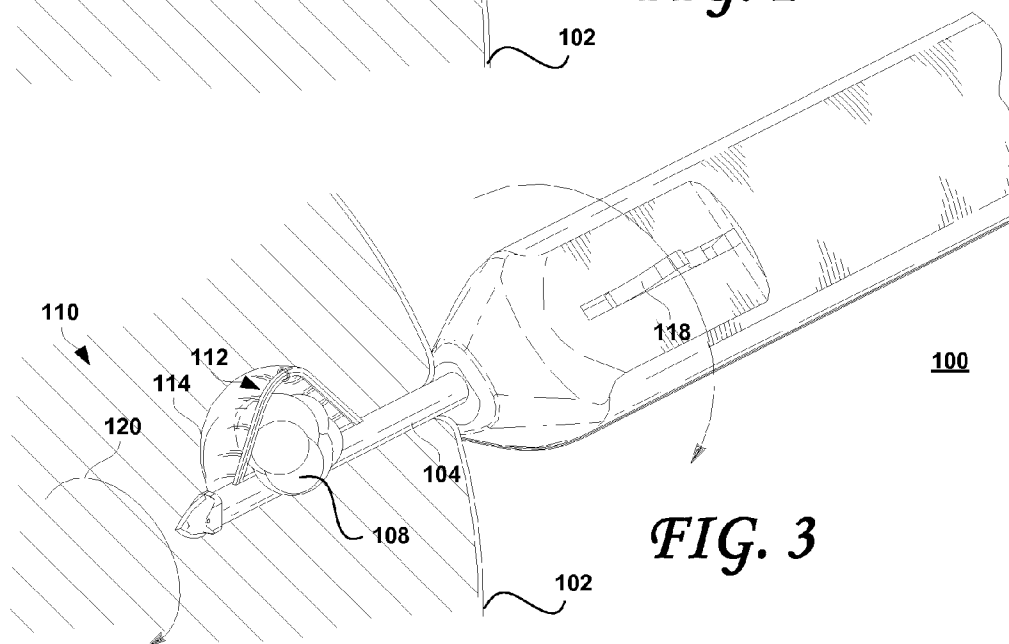
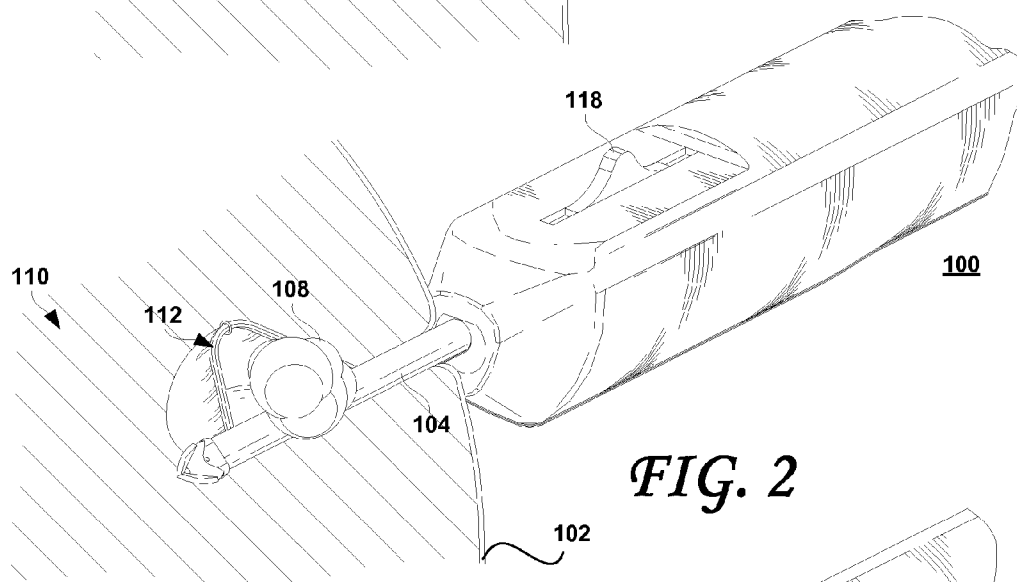
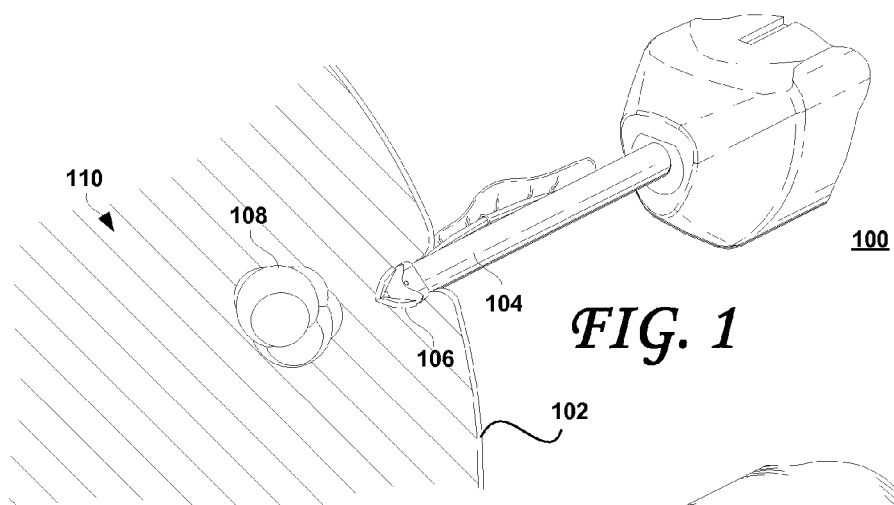


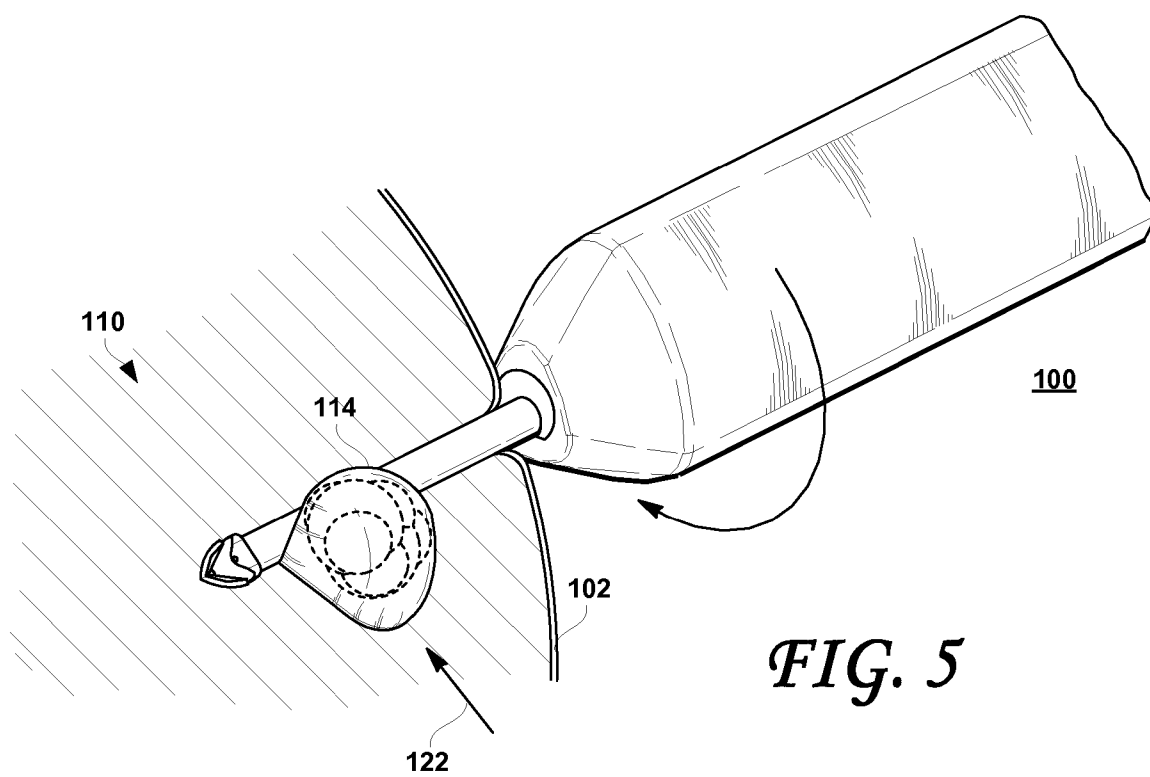
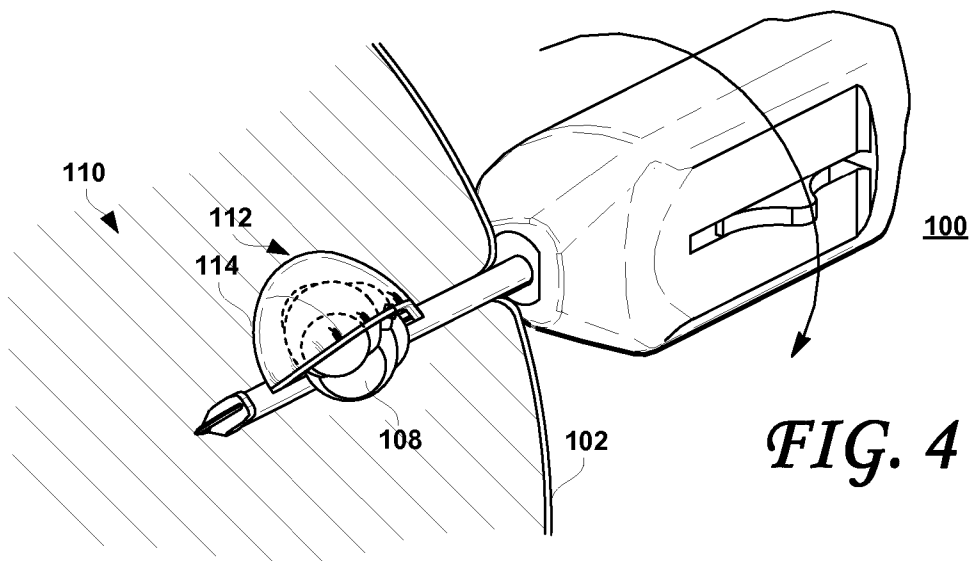
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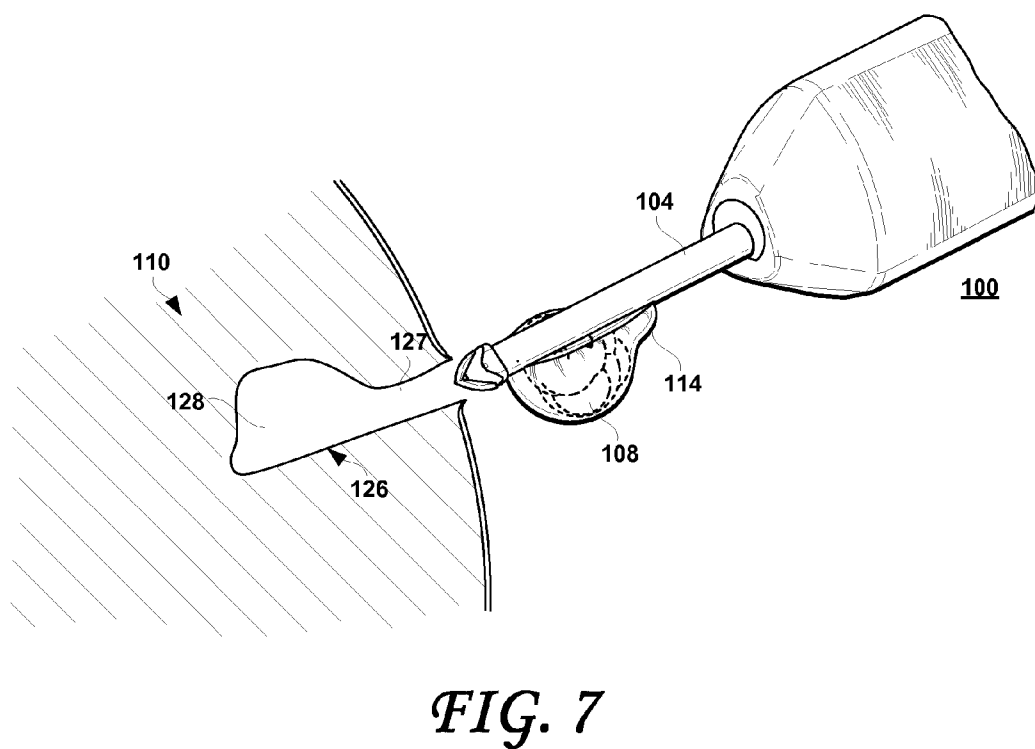
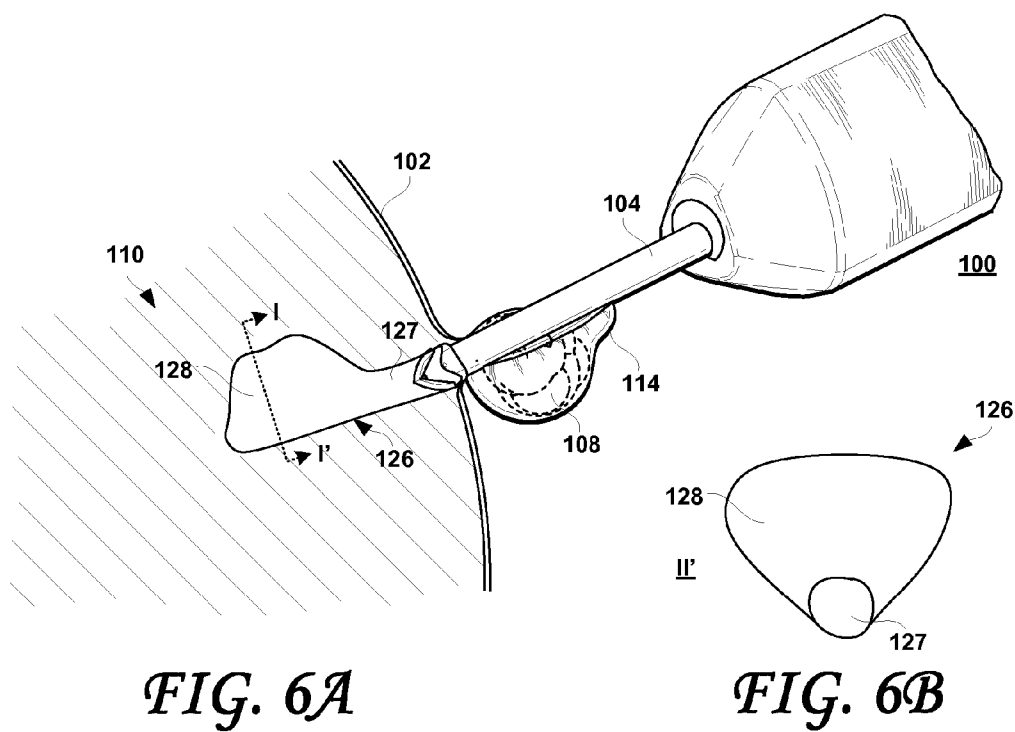
(19) **United States**(12) **Patent Application Publication**
CHERNOMORSKY et al.(10) **Pub. No.: US 2007/0299541 A1**(43) **Pub. Date: Dec. 27, 2007**(54) **POST-BIOPSY CAVITY TREATMENT
IMPLANTS AND METHODS****Publication Classification**(75) Inventors: **Ary S. CHERNOMORSKY**, Walnut
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Portola Valley, CA (US); **Simon**
Chernomorsky, Orinda, CA (US)(51) **Int. Cl.**
A61F 2/02 (2006.01)
(52) **U.S. Cl.** **623/23.72**Correspondence Address:
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PORTOLA VALLEY, CA 94028 (US)(57) **ABSTRACT**

An implant for filling a cavity created by an excisional procedure includes first and second portions. The first portion may include a first collagenous matrix that defines a first selected crosslinking density and the second portion may include a second collagenous matrix that defines a second selected cross-linking density that is different than the first cross-linking density. The first and second cross-linking densities may be selected so as to cause the first and second portions to swell in such a manner that the implant swells into a size and a shape that is similar to the predetermined size and shape of the cavity when the implant is implanted. An aqueous solution may be added to the cavity if the cavity is not sufficiently aqueous to cause the implant to swell.

(73) Assignee: **RUBICOR MEDICAL, INC.**, Redwood
City, CA(21) Appl. No.: **11/850,033**(22) Filed: **Sep. 5, 2007****Related U.S. Application Data**(62) Division of application No. 10/627,960, filed on Jul.
25, 2003.







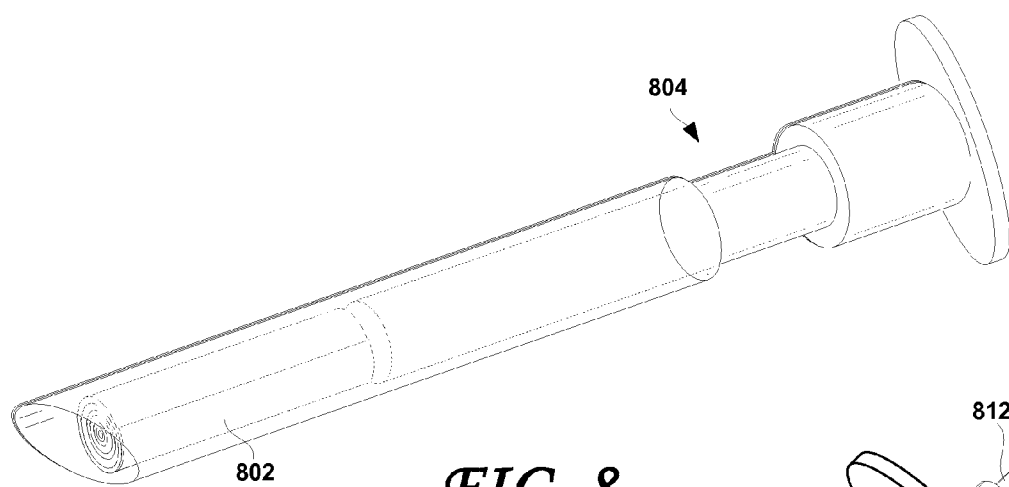


FIG. 8

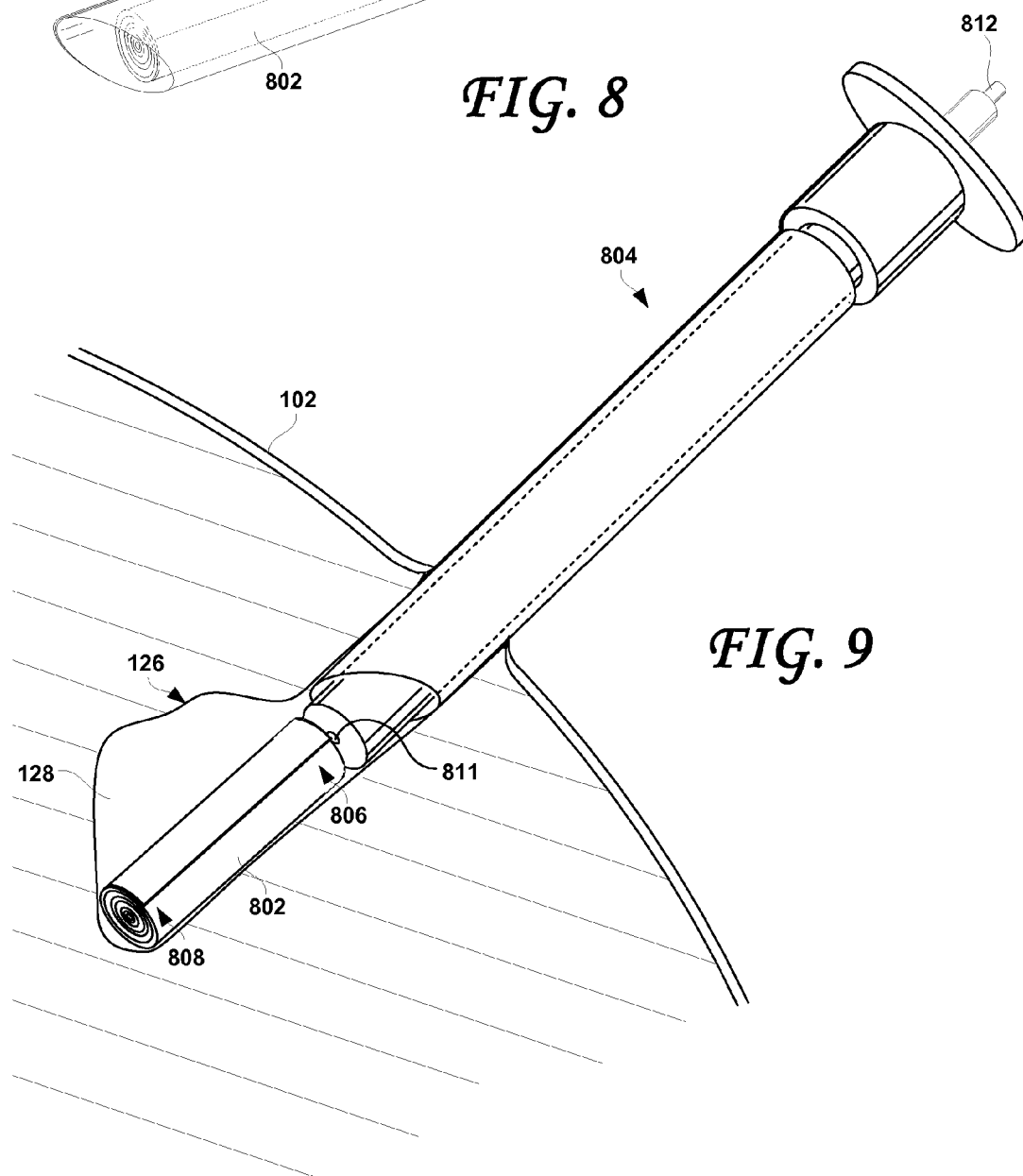


FIG. 9

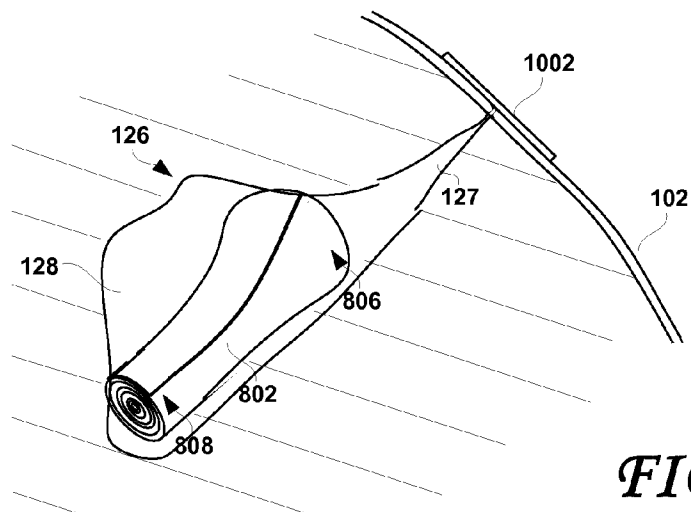


FIG. 10A

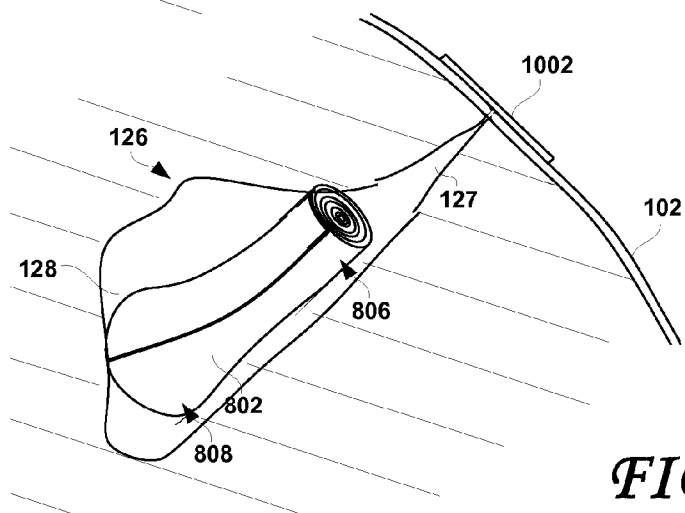


FIG. 10B

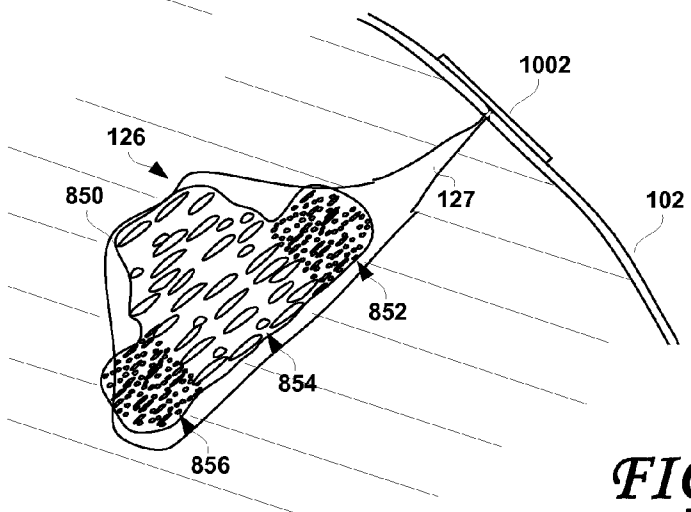
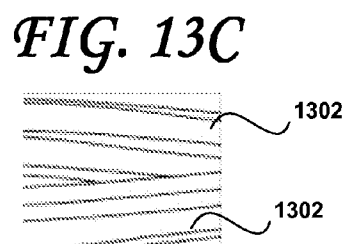
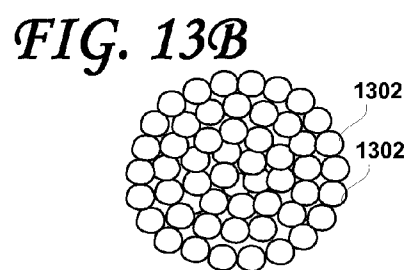
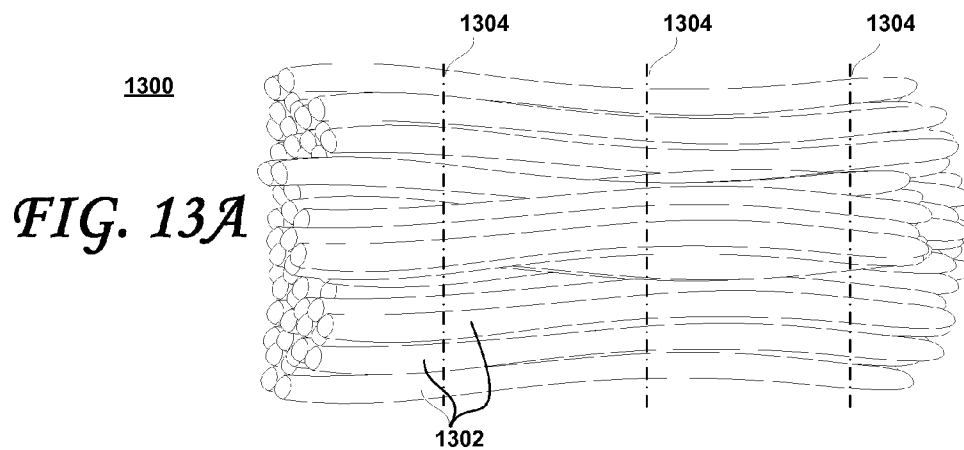
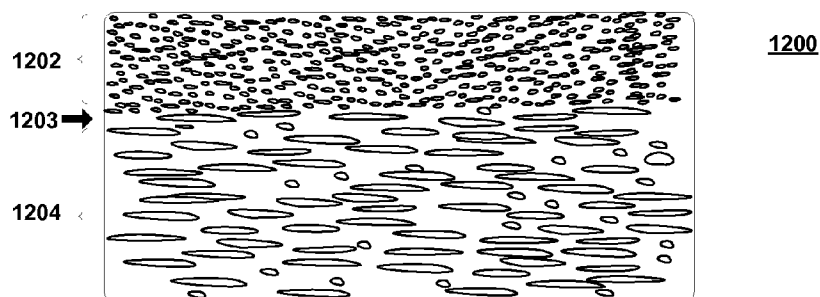
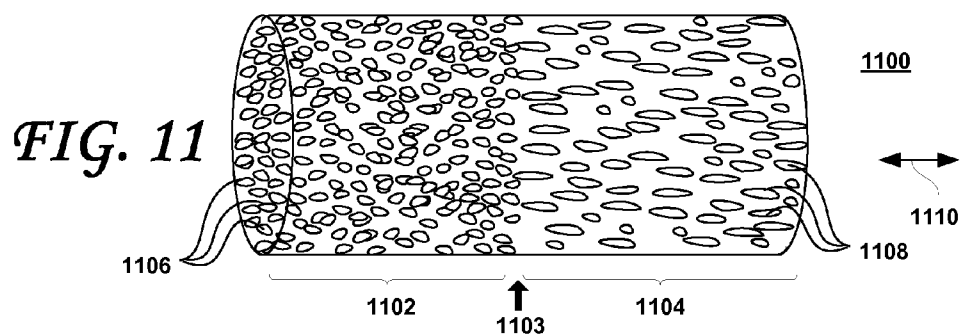


FIG. 10C



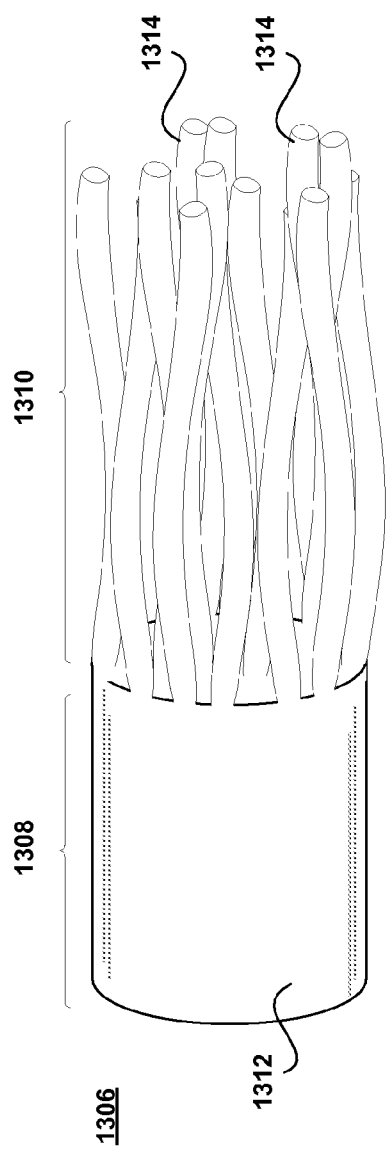


FIG. 13D

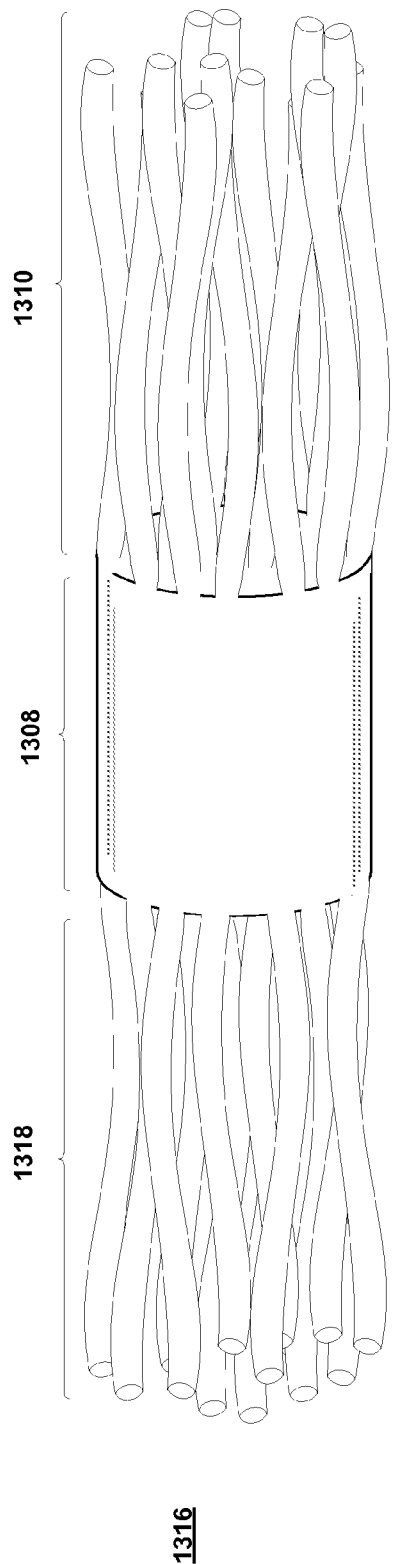


FIG. 13E

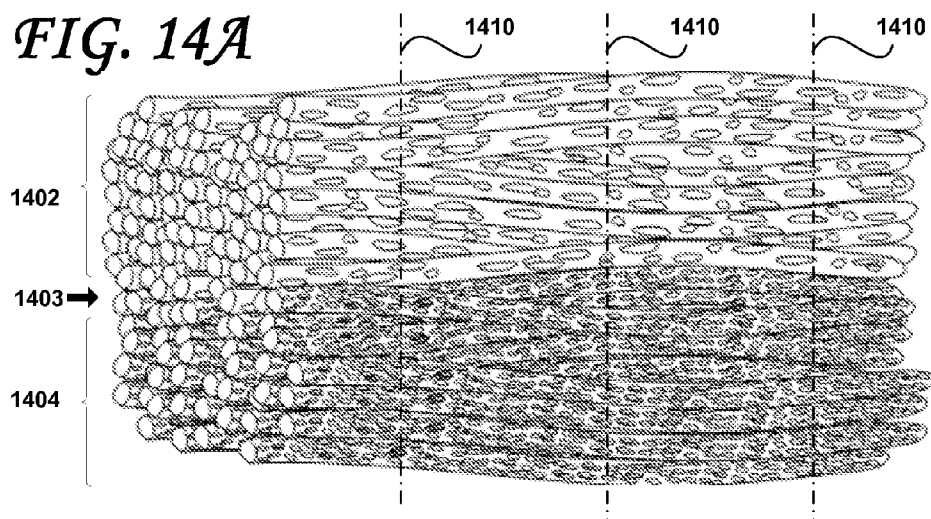


FIG. 14B

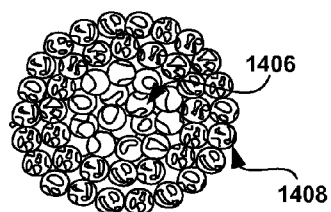


FIG. 14C

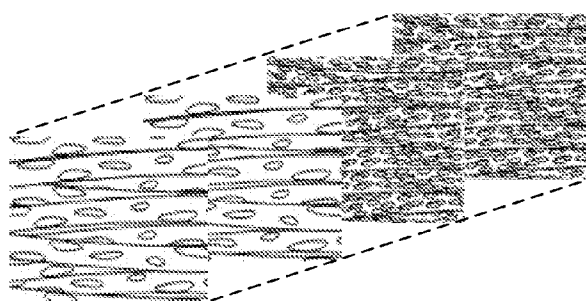
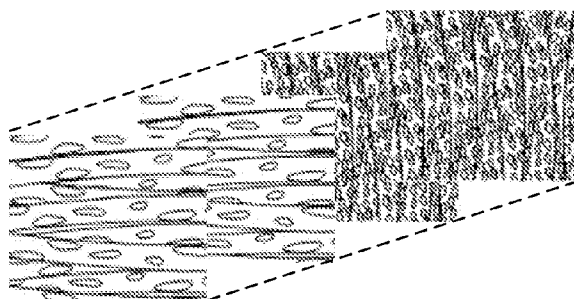


FIG. 14D

FIG. 14E



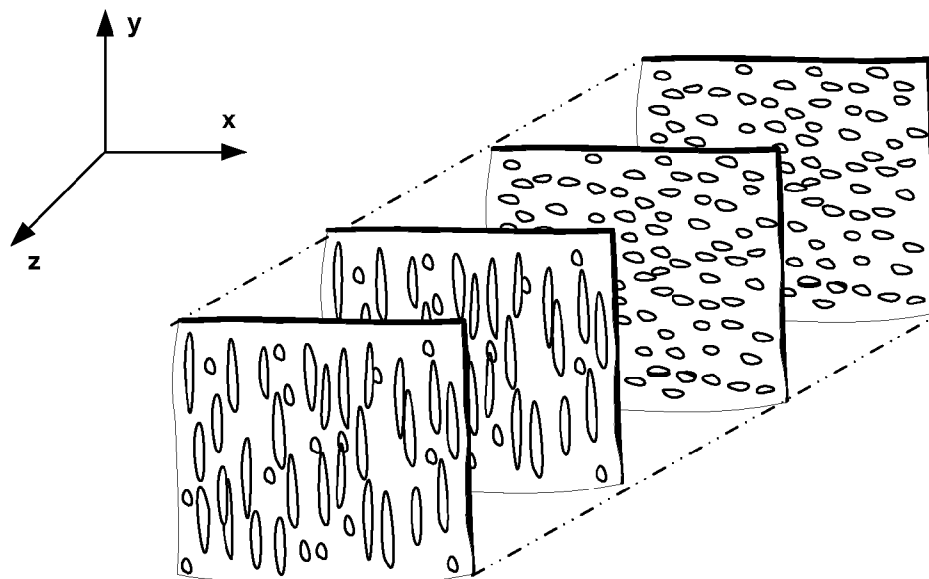


FIG. 14F

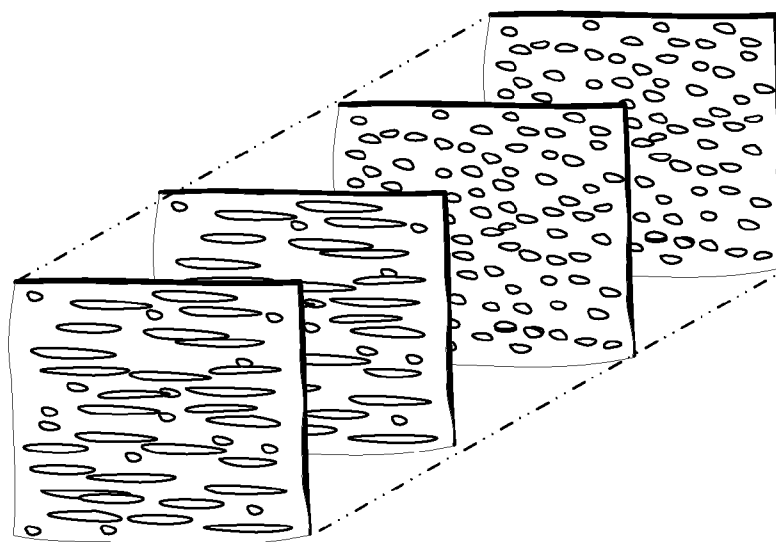


FIG. 14G

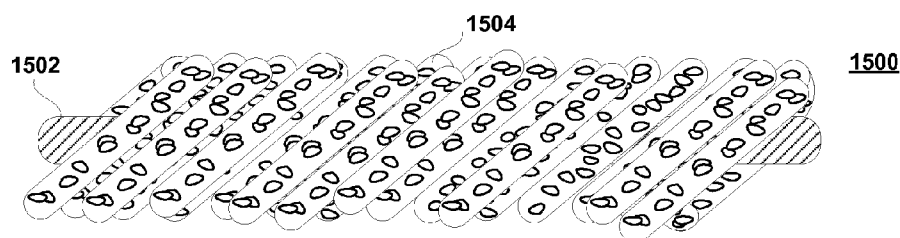


FIG. 15A

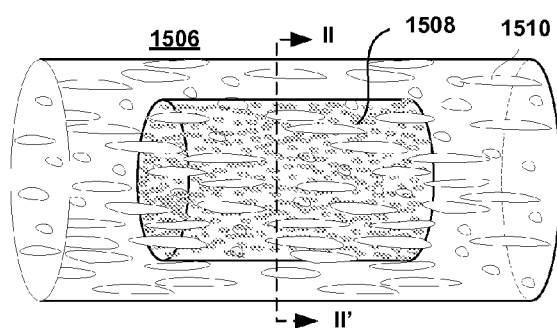


FIG. 15B

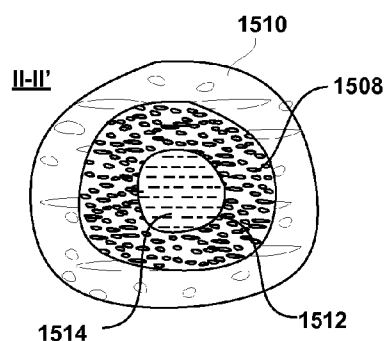


FIG. 15C

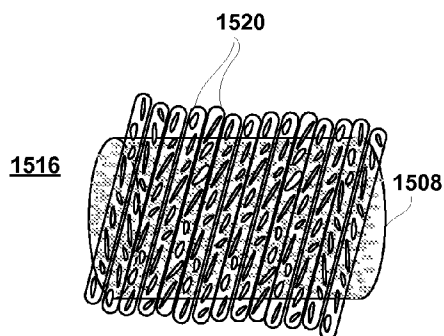


FIG. 15D

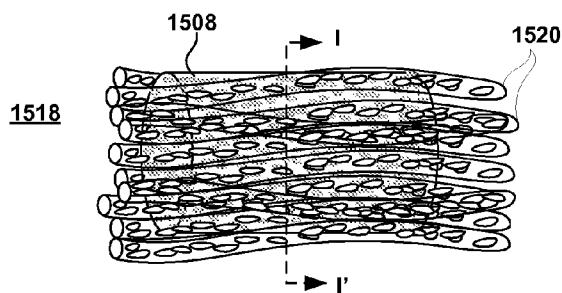


FIG. 15E



FIG. 15F



FIG. 16



FIG. 17



FIG. 18

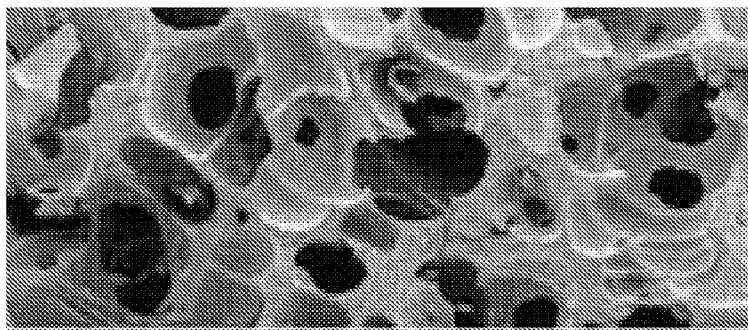


FIG. 19

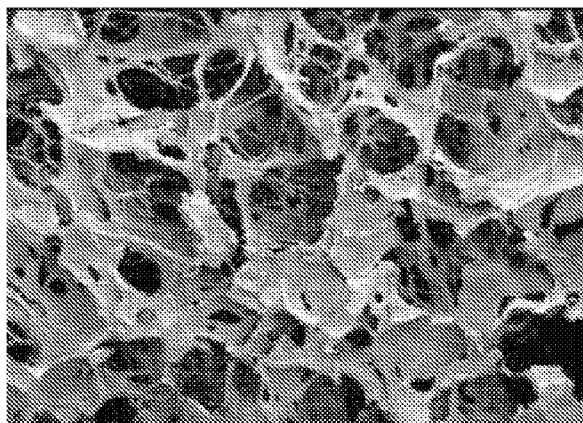


FIG. 20

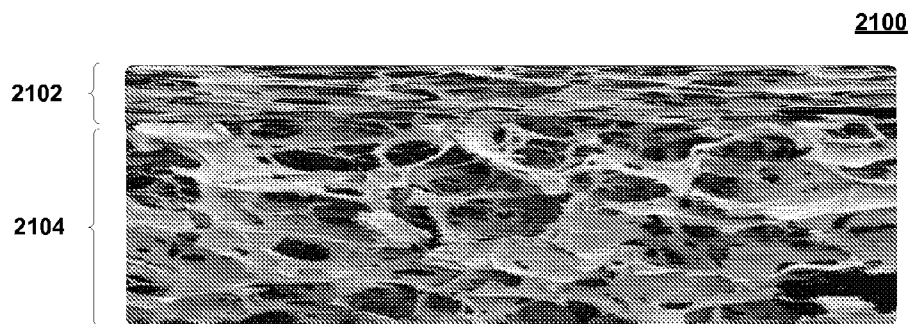
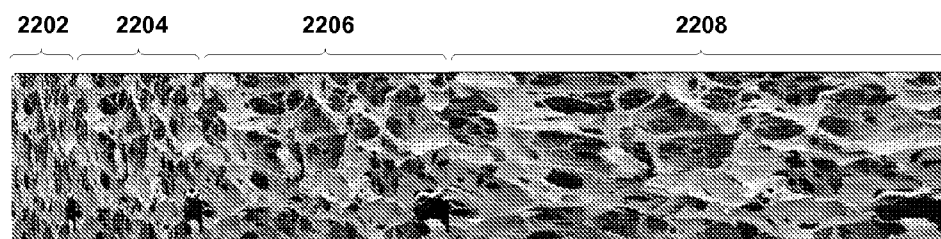


FIG. 21



2200

FIG. 22

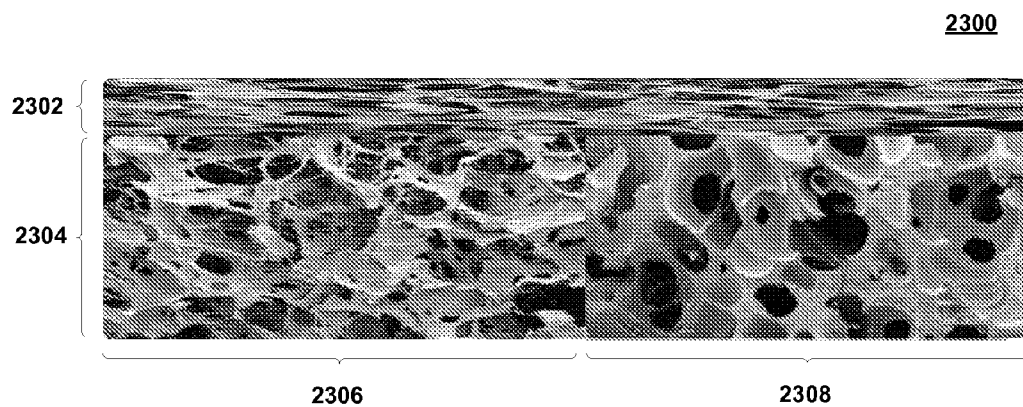


FIG. 23

POST-BIOPSY CAVITY TREATMENT IMPLANTS AND METHODS

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This is a divisional of copending application Ser. No. 10/627,960, filed Jul. 25, 2003, which application is hereby incorporated herein by reference in its entirety.

BACKGROUND OF THE INVENTION

[0002] 1. Field of the Invention

[0003] The present invention relates to post-biopsy cavity treatment methods and implants. More particularly, the present inventions relates to post-biopsy cavity treatment implants inserted into cavities formed in soft tissue that may be created during a biopsy or therapeutic excisional procedure.

[0004] 2. Description of the Related Art

[0005] Breast biopsies are routinely performed in the United States following a detection of abnormalities discovered through mammographic visualization, manual palpation or ultrasound examination. There are a number of traditional methods to obtain breast biopsy tissue samples, including surgical excisional biopsies and stereotactic and ultrasound guided needle breast biopsies. Recently, methodologies have emerged that are based upon percutaneous minimally invasive large intact tissue sample collection. The use of these devices results in a unique cavity connected to the skin by a narrow neck. However, it is becoming apparent, therefore, that the post-biopsy biopsy cavities left by these procedures may both offer and require different post procedural treatments, as compared to the cavities left by needle, core biopsy procedures or open surgical procedures, due to the different nature, size and shape of the cavity created by conventional biopsy devices, as well as the narrow connection to the skin characterized by percutaneous approaches.

[0006] In certain cases, locating a previously biopsied area is highly desirable. Therefore, to mark the biopsy site, a variety of biopsy site markers and identifiers have been developed, ranging from metal clips to pellets and sponges placed during or right after the biopsy procedure. Usually, these markers contain radiopaque and/or echogenic articles and include features such as metal clips and air or gas bubbles incorporated in a biodegradable matrix. From the foregoing, it is apparent that improved methods and devices are needed to treat the cavities left by biopsy devices that are configured to retrieve large intact specimens.

SUMMARY OF THE INVENTION

[0007] The present invention, according to an embodiment thereof, is a method of filling a cavity created by an excisional procedure, the cavity having a predetermined size and shape. The method may include steps of providing an implant, the implant including at least a first portion and a second portion, the first portion comprising a first collagenous matrix that defines a first selected crosslinking density, the second portion comprising a second collagenous matrix that defines a second selected cross-linking density that is different than the first cross-linking density. The first and second cross-linking densities may be selected so as to

cause the first and second portions to swell in such a manner that the implant swells into a size and a shape that is similar to the predetermined size and shape of the cavity when the implant is implanted. The implant may then be implanted within the cavity through an incision. An aqueous solution may be added to the cavity if the cavity is not sufficiently aqueous to cause the implant to swell. The incision may then be closed with the implant implanted in the cavity.

[0008] The providing step may be carried out with the first portion comprising a plurality of first collagenous fibers, each of the plurality of first collagenous fibers having the first selected crosslinking density. The providing step may be carried out with the second portion comprising a plurality of second collagenous fibers, each of the plurality of second collagenous fibers having the second selected cross-linking density.

[0009] According to another embodiment thereof, the present invention is an implant for filling a cavity created by an excisional procedure, the cavity having a predetermined size and a shape. The implant may include a first portion including a first collagenous matrix that defines a first selected crosslinking density, and a second portion including a second collagenous matrix that defines a second selected cross-linking density that is different than the first cross-linking density. The first and second cross-linking densities may be selected so as to cause the first and second portions to swell in such a manner that the implant assumes a size and a shape that is similar to the predetermined size and shape of the cavity when the implant is implanted.

[0010] The first portion may include a plurality of first collagenous fibers, each of the plurality of first collagenous fibers having the first selected crosslinking density. The second portion may include a plurality of second collagenous fibers, each of the plurality of second collagenous fibers having the second selected cross-linking density.

BRIEF DESCRIPTION OF THE DRAWINGS

[0011] For a further understanding of the objects and advantages of the present invention, reference should be made to the following detailed description, taken in conjunction with the accompanying figures, in which:

[0012] FIG. 1 shows an exemplary large intact specimen percutaneous biopsy device in operation.

[0013] FIG. 2 shows further aspects of the exemplary large intact specimen percutaneous biopsy device of FIG. 1 in operation.

[0014] FIG. 3 shows further aspects of the exemplary large intact specimen percutaneous biopsy device of FIG. 1 in operation.

[0015] FIG. 4 shows still further aspects of the exemplary large intact specimen percutaneous biopsy device of FIG. 1 in operation.

[0016] FIG. 5 shows further aspects of the exemplary large intact specimen percutaneous biopsy device of FIG. 1 in operation.

[0017] FIG. 6A shows further aspects of the exemplary large intact specimen percutaneous biopsy device of FIG. 1 in operation, and illustrates the creation of a cavity within the soft tissue from which the excised specimen was taken.

[0018] FIG. 6B is a cross sectional view of the post treatment cavity of FIG. 6A, taken along cross-sectional line II'.

[0019] FIG. 7 shows further aspects of the exemplary large intact specimen percutaneous biopsy device of FIG. 1 in operation, and further illustrates the creation of a cavity within the soft tissue from which the specimen was taken, with the aforementioned narrow neck or access path connecting the cavity to the skin.

[0020] FIG. 8 shows an exemplary delivery device for a post-biopsy cavity treatment implant, according to an embodiment of the present invention.

[0021] FIG. 9 shows the delivery device of FIG. 8 in operation, delivering a post-biopsy cavity treatment implant according to an embodiment of the present invention within the cavity of FIG. 7.

[0022] FIG. 10A shows the cavity of FIG. 7, after the implantation of the post-biopsy cavity treatment implant shown in FIGS. 8 and 9, with the percutaneous incision closed.

[0023] FIG. 10B shows the cavity of FIG. 7, after the implantation of the post-biopsy cavity treatment implant shown in FIGS. 8 and 9 in another orientation, with the percutaneous incision closed.

[0024] FIG. 10C shows the cavity of FIG. 7, after the implantation of a post-biopsy cavity treatment implant according to another embodiment of the present invention, with the percutaneous incision closed.

[0025] FIG. 11 shows a post-biopsy cavity treatment implant having a predetermined pore architecture, according to an embodiment of the present invention.

[0026] FIG. 12 shows another post-biopsy cavity treatment implant having another predetermined pore architecture, according to another embodiment of the present invention.

[0027] FIG. 13A shows a post-biopsy cavity treatment implant that includes a plurality of fibers, according to another embodiment of the present invention.

[0028] FIG. 13B shows a cross-section of a post-biopsy cavity treatment implant, according to another embodiment of the present invention.

[0029] FIG. 13C shows a portion of another post-biopsy cavity treatment implant, according to a further embodiment of the present invention.

[0030] FIG. 13D shows another post-biopsy cavity treatment implant, according to still another embodiment of the present invention.

[0031] FIG. 13E shows another post-biopsy cavity treatment implant, according to still another embodiment of the present invention.

[0032] FIG. 14A shows a post-biopsy cavity treatment implant that includes a plurality of fibers having predetermined pore architectures, according to another embodiment of the present invention.

[0033] FIG. 14B shows a front view of a post-biopsy cavity treatment implant, according to another embodiment of the present invention.

[0034] FIG. 14C shows a portion of another post-biopsy cavity treatment implant, according to a further embodiment of the present invention.

[0035] FIG. 14D illustrates the stacked structure of a post-biopsy cavity treatment implant, according to a further embodiment of the present invention.

[0036] FIG. 14E illustrates the stacked structure of another post-biopsy cavity treatment implant, according to a further embodiment of the present invention.

[0037] FIG. 14F illustrates the stacked structure of a post-biopsy cavity treatment implant, according to a further embodiment of the present invention.

[0038] FIG. 14G illustrates the stacked structure of another post-biopsy cavity treatment implant, according to a further embodiment of the present invention.

[0039] FIG. 15A shows a post-biopsy cavity treatment implant that includes a radiopaque and/or echogenic member around which one or more fibers are wound, according to another embodiment of the present invention.

[0040] FIG. 15B shows a post-biopsy cavity treatment implant that includes a core portion surrounded by an outer shell portion, each of the core and shell portions having a predetermine core architecture, according to another embodiment of the present invention.

[0041] FIG. 15C is a cross-sectional representation of the implant of FIG. 15B, taken along cross-sectional line II-II'.

[0042] FIG. 15D shows a post-biopsy cavity treatment implant that includes a core portion having a first predetermine core architecture surrounded by an outer shell portion formed by a plurality of wound collagenous fibers having a second predetermined pore architecture, according to another embodiment of the present invention.

[0043] FIG. 15E shows a post-biopsy cavity treatment implant that includes a core portion having a first predetermine core architecture surrounded by an outer shell portion formed by a plurality of collagenous fibers having a second predetermined pore architecture, according to another embodiment of the present invention.

[0044] FIG. 15F is a cross-sectional view of the embodiment of FIG. 15D, taken along cross-sectional line I-I'.

[0045] FIG. 16 is a photomicrograph of a collagen matrix having a predetermined pore architecture with post-biopsy cavity treatment implants according to embodiments of the present invention may be constructed.

[0046] FIG. 16 is a photomicrograph of a collagen matrix having a predetermined pore architecture with post-biopsy cavity treatment implants according to embodiments of the present invention may be constructed.

[0047] FIG. 17 is a photomicrograph of a collagen matrix having another predetermined pore architecture with post-biopsy cavity treatment implants according to embodiments of the present invention may be constructed.

[0048] FIG. 18 is a photomicrograph of a collagen matrix having still another predetermined pore architecture with post-biopsy cavity treatment implants according to embodiments of the present invention may be constructed.

[0049] FIG. 19 is a photomicrograph of a collagen matrix having a still further predetermined pore architecture with post-biopsy cavity treatment implants according to embodiments of the present invention may be constructed.

[0050] FIG. 20 is a photomicrograph of a collagen matrix having yet another predetermined pore architecture with post-biopsy cavity treatment implants according to embodiments of the present invention may be constructed.

[0051] FIG. 21 combination of photomicrographs of collagen matrices illustrating the formation of a stacked laminate structure including a first layer having a first predetermined pore architecture and a second layer having a second predetermined pore structure, according to an embodiment of the present invention.

[0052] FIG. 22 is a combination of photomicrographs of collagen matrices that collectively illustrate a post-biopsy cavity treatment implant having a predetermined pore density gradient and/or predetermined graduated crosslinking gradient, according to a further embodiment of the present invention.

[0053] FIG. 23 is a combination of photomicrographs of collagen matrices that collectively illustrate a post-biopsy cavity treatment implant according to another embodiment of the present invention.

DETAILED DESCRIPTION OF THE INVENTION

[0054] FIGS. 1-7 show aspects of a percutaneous method for cutting, collecting and isolating a tissue specimen and the subsequent creation of a cavity within which embodiments of the present inventions may be implanted. The excisional device shown in FIGS. 1-7 is described in commonly assigned U.S. Pat. No. 6,022,362 and in commonly assigned and co-pending application Ser. No. 10/189,277 filed Jul. 3, 2002, the patent and application of which are hereby incorporated herein by reference in their entireties. Although embodiments of the present invention are described relative to a cavity created by the excisional device shown in FIGS. 1-7, it is to be understood that the present inventions are not to be limited thereby. Indeed, embodiments of the present invention may be advantageously utilized to treat cavities of various shapes and sizes created by other devices, including devices that obtain tissue specimen through coring or ablation techniques, for example.

[0055] As shown in FIG. 1, the excisional device 100 is introduced into a mass of tissue 110 through the skin 102, with the integrated cut and collect assembly 112 thereof in a retracted position. The device 100 is then advanced such that the assembly 112 is adjacent to the target lesion 108. The assembly 112 may then be energized and expanded as shown in FIG. 2 by acting upon the actuator 118. As the assembly 112 is RF energized and expanded, it cuts the tissue through which it travels. As shown at FIG. 3, the excisional device 100 may then be rotated, while the assembly 112 remains energized, causing the leading edge thereof to cut through the tissue, preferably with clean margins. The expanded integrated cut and collect assembly 112 deploys the membrane 114 and the cut specimen 108 is collected in the open bag formed by the close-ended deployed flexible membrane 114. As shown in FIGS. 4 and 5, the rotation of the device 100 may then be continued as needed, preferably

under ultrasonic guidance. To fully sever the specimen 108 from the surrounding tissue 110, the assembly 112, while still energized, is retracted to capture, encapsulate and isolate the specimen 108 within the flexible membrane 114. As shown in FIG. 6A, the specimen 108 may then be recovered by retracting the device 100 through the retraction path 127, stretching it as necessary. FIG. 7 shows a fully retracted device 100, containing a collected and isolated specimen 108.

[0056] As shown in FIGS. 6A-7, after the procedure described above or after any procedure in which a substantial volume of tissue specimen is taken, a void or cavity 126 is created where the tissue specimen 108 used to be. Cavities as shown at 126 may require different post procedural treatments, as compared to procedures such as needle biopsies due to the different nature, size and shape created by the biopsy device. As shown in FIGS. 6A and 6B, the exemplary cavity 126 is characterized by a relatively narrow access path 127 that emerges into a larger cavity chamber 128 formed by the extension and rotation of the cut and collect assembly 112 during the above-described procedure. After the device 100 is withdrawn from the patient as shown in FIG. 7, portions of the cavity 126 and/or access path 127 may settle and collapse somewhat, as the interior tissue walls defining the cavity 126 are no longer supported by the tissue previously occupying that space.

[0057] Treating the post-biopsy cavity 126 is desirable for a variety of reasons. One such reason is to accommodate the unique size and shape of the cavity 126 created by the device 100. It is desirable to influence and/or promote the healing process of the cavity, and to do so in a predictable manner. One aspect of influencing the healing process of the cavity 126 is promoting the growth of new connective tissue within the cavity 126 in a predictable manner. Indeed, it is desirable to influence and promote both tissue ingrowth within the cavity and to influence the formation of hematomas and seromas. Another reason for treating the post-biopsy cavity 126 is to modify it in such a manner as to render it recognizable immediately and preferably long after the procedure that created the cavity 126. The cavity 126, left untreated, may be visible under ultrasound. However, that may not be the case and it is believed to be desirable to at least partially fill the cavity 126 with a cavity treatment implant that will render the cavity 126 clearly visible under various imaging modalities, including modalities such as ultrasound, x-ray, magnetic resonance imaging, elastography, microwave and the unaided eye, for example. Such visibility may be due to the structure of a cavity treatment implant or devices implanted within the cavity and/or a recognizable pattern of tissue ingrowth caused or influenced by the continuing or past presence of post-biopsy cavity treatment implants disclosed herein. Other desirable attributes of embodiments of the implantable post-biopsy cavity treatment implant of the present invention include hemostasis, and the ability to deliver one or more therapeutic agents to the patient at the post-biopsy cavity treatment implant site such as, for example, lido/epi, Non-Steroidal Anti-Inflammatory Drugs (NSAIDs), tissue growth factors, anti-neoplastic medications (to name a few) or combinations of the above and/or others. Filling the cavity 126 may have other benefits, including cosmetic. Indeed, filling the cavity and promoting a smooth, gradual, recognizable and orderly tissue ingrowth pattern may prevent dimpling, skin depressions and the like sometimes associated with the removal of

a large intact specimen during the biopsy procedure. Embodiments of the present invention may also find utility in augmentation or reconstructive procedures for the breast or other soft tissue.

[0058] According to an embodiment of the present invention, the post-biopsy cavity treatment implant may have a size and a shape that at least partially fills the cavity. Advantageously, the present post-biopsy cavity treatment implant, after insertion, may have a characteristic shape that is readily perceptible and recognizable through various modalities, including, for example, ultrasound, x-ray or magnetic resonance imaging. The shape of the present post-biopsy cavity treatment implant may also influence the manner in which tissue growths therein. Preferably, embodiments of the present post-biopsy cavity treatment implant should be shaped and dimensioned so as to uniquely accommodate the size and shape of the cavity 126 created by the device 100 of FIGS. 1-7. However, embodiments of the present invention may be readily sized and shaped to specifically accommodate cavities of any shape and size created by other devices and/or biopsy or therapeutic surgical procedures.

[0059] According to an embodiment thereof, the present invention may include an implantable post-biopsy cavity treatment implant having one or more of the structures, characteristic and properties described herein. As shown in FIG. 8, the implantable post-biopsy cavity treatment implant 802, in a pre-implanted state, may be loaded into an introducer, an illustrative example of which is shown at reference numeral 804. The introducer 804 may then be inserted into the tissue 110 through the access path 127 and at least partially into the cavity chamber 128 of the cavity 126. The post-biopsy cavity treatment implant 802 may then be delivered to the cavity 126 and thereafter be left in place and the introducer 804 withdrawn. The pre-implanted state of the post-biopsy cavity treatment implant 802 is preferably in a state in which the post-biopsy cavity treatment implant occupies its minimum volume. According to an embodiment of the present invention, the pre-implanted state of the post-biopsy cavity treatment implant 802 is a lyophilized (e.g., dehydrated) state and the post-biopsy cavity treatment implant may be configured to swell when placed within aqueous environment such as the cavity 126. The post-biopsy cavity treatment implant 802 may define a proximal portion 806 that is closest to the access path 127 and a distal portion 808 that is relatively further away from the access path 127 than is the proximal portion 806.

[0060] Whereas FIG. 9 shows the present post-biopsy cavity treatment implant 802 immediately after implantation in tissue (i.e., still in a state in which it occupies its minimum volume), FIG. 10A shows the state of the present post-biopsy cavity treatment implant 802 a short period of time after implantation. As shown, the post-biopsy cavity treatment implant 802 is no longer in its pre-implanted state. Indeed, the post-biopsy cavity treatment implant 802, having been placed in an aqueous environment (such as the patient's tissue), begins to swell. According to an embodiment of the present invention, the post-cavity treatment implant 802 may be configured to swell in a uniform manner. In another example, the surgeon may inject fluids after placing the device with the intent to "wet" the present post-cavity treatment implant. Substances such as saline, fibrin solution or other catalyst or activator may be used for

that purpose. The activator or swelling fluid could be injected preferentially at the proximal portion 806 or selectively at points in the post-cavity treatment implant to cause it to secure itself in position inside the cavity 126. Alternatively, as part of the insertion device (such as, for example, the introducer 804), an integral vial may be crushed by the surgeon to release the activating fluid (for example, an aqueous solution, dye/pigment) in the area of the proximal portion 806 for example, thus causing rapid swelling of that region. Alternately, the introducer 804 may define an internal lumen 811 over its length and may include a fluid injection port 812 at the proximal end of the device. Fluids such as the aforementioned saline or fibrin may then be introduced into the cavity 126 through the fluid injection port 812 and the internal lumen 811 to cause the rapid swelling of the implant or for any other reason. Delivering such fluids can be especially useful if the field within the cavity is relatively dry as can occur in the ideal case. According to another embodiment of the present invention, the post-biopsy cavity treatment implant 802 may be configured to swell non-uniformly. Such non-uniform swelling rates may be advantageous in insuring that the post-biopsy cavity treatment implant 802 stays where it is placed during the implantation procedure. In the embodiment shown in FIG. 10A, the post-biopsy cavity treatment implant 802 is structured such that the rate at which the proximal portion 806 swells faster than the rate at which the distal portion 808 swells. When implanted in a cavity 126 such as shown in FIGS. 6A, 6B, 7, 9 and 10, the proximal portion 806 swells faster than the distal portion 808, thereby serving to maintain the post-biopsy cavity treatment implant 802 within the cavity chamber 128 of the cavity 126. This may be achieved by, for example, controlling the crosslinking densities or creating a gradient of crosslinking densities within the post-biopsy cavity treatment implant 802, where certain regions of the post-biopsy cavity treatment implant 802 are controlled to have a greater crosslinking density than other regions, resulting in a non-uniform swelling pattern over the extent of the device 802. For example, the distal portion 808 may be configured to be relatively more crosslinked than the proximal portion 806 thereof, resulting in the proximal portion 806 swelling more and/or faster than the distal portion 808. As the proximal portion 806 of the post-biopsy cavity treatment implant 802 swells, it preferably swells from a shape in which it is easily implantable through the access path 127 to a shape and size wherein at least the proximal portion 806 thereof no longer fits through the access path 127. As this swelling occurs rapidly after the post-biopsy cavity treatment implant 802 comes into contact with the fluids present within the cavity 126, the surgeon may retract the introducer 804 from the cavity 126, close the initial incision and be confident that the post-biopsy cavity treatment implant 802 has remain in its intended position, squarely within the cavity chamber 128 of the cavity 126, and has not migrated back into the access path 127.

[0061] The post-biopsy cavity treatment implant 802 may alternatively be structured such that its distal 808 portion swells faster than its proximal portion 806 such as shown in FIG. 10B, such that both the proximal and distal portions 806, 808 of the post-biopsy cavity treatment implant swell relatively faster than the portion thereof between the proximal and distal portions or such that the proximal and distal portions 852, 856 of the implant 850 swell relatively slower than a middle portion 854, as shown in FIG. 10C. Alternatively

still, the post-biopsy cavity treatment implant **802** may not have well defined proximal and distal portions and the post-biopsy cavity treatment implant **802** may be structured such that one portion thereof swells at a different rate than another portion thereof, for the purpose outlined above or for different purposes altogether—such as cavity shaping, for example. As suggested in FIGS. **9** and **10A**, **10B**, the post-biopsy cavity treatment implant **802** may be formed from a tightly rolled up sheet of swellable material. Alternatively, the post-biopsy cavity treatment implant **802** may be formed of stacked layers of swellable material as shown in FIG. **10C**. Alternatively still, the post-biopsy cavity treatment implant **802** may be formed as a single unitary and homogeneous mass of swellable material and molded or cut (stamped) into the desired shape. Other embodiments include post-biopsy cavity treatment implants formed of or including fibers, fibrils and/or bundles of fibers and/or fibrils.

[**0062**] According to embodiments of the present invention, the present post-biopsy cavity treatment implant may include or be formed of biocompatible and water swellable material, such as collagen, for example. The collagen molecule is rod-shaped triple helix and consists of a three polypeptide chains coiled about each other. Besides the central triple helical region of the collagen molecule, there are terminal peptides regions known as telopeptides. These telopeptides are non-helical and are subdivided into two groups; namely, amino terminals and carboxyl terminals. Intermolecular crosslinking between triple helical molecules of collagen occurs in the telopeptides regions. Crosslinking may also occur within the central triple helical region of the collagen molecule, and is known as intramolecular crosslinking. It is the control of the formation and density of such crosslinks that is responsible for some of the mechanical, physicochemical and biological properties of the embodiments of the present post-biopsy cavity treatment implant disclosed herein.

[**0063**] The embodiments of the present post-biopsy cavity treatment implant may be selectively biodegradable and/or bio-absorbable such that it degrades and/or is absorbed after its predetermined useful lifetime is over. An effective way of controlling rate of biodegradation of embodiments of the present post-biopsy cavity treatment implant is to control and selectively vary the number and nature (e.g., intermolecular and/or intramolecular) of crosslinks in the implant material. Control of the number and nature of such collagen crosslinks may be achieved by chemical and/or physical means. Chemical means include the use of such bifunctional reagents such as aldehyde or cyanamide, for example. Physical means include the application of energy through dehydrothermal processing, exposure to UV light and/or limited radiation, for example. Also, a combination of both the chemical and the physical means of controlling and manipulating crosslinks may be carried out. Aldehydes such as glutaraldehydes, for example, are effective reagents of collagenous biomaterials. The control and manipulation of crosslinks within the collagenous matrix of the present post-surgery cavity treatment implant may also be achieved, for example, through a combination of dehydrothermal crosslinking and exposure to cyanamide. For example, the present post-surgery cavity treatment implant may, through proper control of the crosslinking density within the collagen matrix thereof, be designed and implemented to remain long term in situ at the implant site within the cavity

126. Crosslinking density may be indirectly measured, for example, via measurement of the swelling ratio where identical dry and wetted samples are weighted and weight is compared.

[**0064**] According to further embodiments of the present invention, the post-biopsy cavity treatment implant may be formed of or include other biomaterials such as, for example, bioresorbable poly(ester)s such as polylactide (PLA), polyglycolide (PGA), poly(lactide-co-glycolides) (PLGA), polyglyconate, polyanhydrides and their co-polymers, PEG, cellulose, gelatins, lipids, polysaccharides, starches and/or polyorthoesters and the like. According to an embodiment of the present invention, the present post-biopsy cavity treatment implant may be formed of or include collagen having a predetermined structure. Such predetermined structure refers not only to the overall shape of the implant, but also to the structure of its internal collagen matrix. Indeed, embodiments of the present invention include a macroporous cross-linked polymer matrix having a predetermined pore architecture. A “pore”, as the term is used herein, includes a localized volume of the present post-biopsy cavity treatment implant that is free of the material from which the post-biopsy cavity treatment implant is formed. Pores may define a closed and bounded volume free of the material from which the post-biopsy cavity treatment implant is formed. Alternatively, pores may not be bounded and many pores may communicate with one another throughout the internal matrix of the present post-biopsy cavity treatment implant. The pore architecture, therefore, may include closed and bounded voids as well as unbounded and interconnecting pores and channels. The internal structure of the post-biopsy cavity treatment implant according to embodiment of the present invention defines pores whose dimensions, shape, orientation and density (and ranges and distributions thereof), among other possible characteristics are tailored so as to maximize the visibility of the resultant post-biopsy cavity treatment implant **802** under various modalities, notably ultrasound and x-ray, for example. Unlike polymeric matrices that contain bubbles of gas through a process in which gas is forced through a dispersion in a hydrated state, embodiments of the present post-biopsy cavity treatment implant have an internal structure that defines internal voids without requiring such gas to be forced therethrough. There are numerous methods and technologies available for the formation collagenous matrices of different pore architectures and porosities. By tailoring the dimensions, shape, orientation and density of the pores of the present implant, a recognizable pattern of post-biopsy cavity treatment implant material may be formed that may be readily visualized under, for example, ultrasound, x-ray, elastography or microwave radiation. This recognizable pattern may then influence the pattern of tissue ingrowth within the cavity **126**, forming a porous scaffold on and within which tissue may infiltrate and grow. In turn, this pattern of tissue ingrowth may be readily recognizable under ultrasound and/or other imaging modalities discussed above long after the post-biopsy cavity treatment implant has been absorbed by the body or has degraded.

[**0065**] According to an embodiment of the present invention, the post-biopsy cavity treatment implant may be formed of or include a collagen matrix having a predetermined pore architecture. For example, the post-biopsy cavity treatment implant may include one or more sponges of lyophilized collagen having a predetermined pore architec-

ture. Suitable collagen material for the post-biopsy cavity treatment implant may be available from, for example, DEVRO, Integra Life Sciences, Collagen Matrix and Kensey Nash, among others. The present post-biopsy cavity treatment implant, after implantation in the cavity **126**, swells on contact with various body fluids therein and substantially fills a predetermined portion or the entire biopsied cavity, and does so in predictable manner.

[0066] Such a post-biopsy cavity treatment implant may be configured to have a hemostatic functionality to stop bleeding within the cavity **126** through a biochemical interaction with blood (such as coagulation) and/or other bodily fluids. The post-biopsy cavity treatment implant may, according to further embodiments, also be used to medically treat the patient. That is, the porous matrix of the present post-biopsy cavity treatment implant may be imbibed or loaded with a therapeutic agent to deliver the agent through elution at the cavity **126**. Such a therapeutic agent may include, for example, an antibiotic agent, an analgesic agent, a chemotherapy agent, an anti-angiogenesis agent or a steroidal agent, to name but a few of the possibilities.

[0067] The post-biopsy cavity treatment implant **802** shown in FIGS. **9** and **10** may be formed of one or more thin sheets of collagen material having a predetermined (and controlled) pore architecture that has been rolled up into a cylinder shape. As the post-biopsy cavity treatment implant **802** swells with water from the cavity **126**, it may unroll partially or entirely, and at least partially fill the cavity **126**, including at least a portion of the cavity chamber **128**. Some of the access path **127** may also be filled as the post-biopsy cavity treatment implant **802** swells. The post-biopsy cavity treatment implant **802**, according to embodiments of the present invention, has a predetermined pore architecture or a combination of predetermined pore architectures, as will be described hereunder with reference to the drawings. The description of the figures below assumes that the post-biopsy cavity treatment implant is formed of or contains collagen, it being understood that the embodiments of the present invention disclosed herein are not limited to collagen and that aspects of the present inventions may readily be applied to such non-collagen containing post-biopsy cavity treatment implants.

[0068] FIG. **11** shows a post-biopsy cavity treatment implant **1100** having predetermined pore architectures, according to an embodiment of the present invention. As shown therein, the post-biopsy cavity treatment implant **1100** may include a first portion **1102** and a second portion **1104**. The collagen matrix of the first portion **1102** of the device **1100** defines a plurality of pores **1106** having a first predetermined pore architecture and the collagen matrix of the second portion **1104** of the device **1100** defines a plurality of pores **1108** having a second predetermined pore architecture. The dimensions of the layers or portions may be selected at will, preferably accounting for the dimensions of the cavity into which the device is to be inserted. As shown, the first pore architecture features pores **1106** that are relatively small, have a narrow pore size distribution and are substantially randomly oriented. In contrast, the second pore architecture features pores **1108** that have a relatively larger size, have a wider pore size distribution, are predominantly oriented along the axis indicated by double-headed arrow **1110** and are less densely distributed than the pores **1106** of the first portion **1102** of the post-biopsy cavity treatment

implant **1100**. Between the first and second portions **1102** and **1104** lies the interface **1103**. As shown, the post-biopsy cavity treatment implant **1100** may be formed of a first collagen matrix having a first predetermined pore architecture and a second collagen matrix having a second pore architecture. The two collagen matrices may each be formed from separate collagen dispersions, each of which may be caused to form pores having predetermined characteristics and may each be lyophilized and formed (e.g., molded, cut or stamped) into the desired shape (in the illustrative case of FIG. **11**, a substantially cylindrical shape). The two collagen plugs formed thereby may then be stacked one on the other, re-wetted and again lyophilized through a lyophilization process in a specifically shaped mold (for example) to form the stacked laminate structure shown in FIG. **11**. Other methods of making the post-biopsy cavity treatment implant **1100** may occur to those of skill in this art. Not only may the predetermined pore architectures of the first portion **1102** and of the second portion **1104** cause these portions to be visible under, for example, ultrasound, but the interface **1103** therebetween may also be visualizable and recognizable under, for example, ultrasound as the boundary between two regions having a pronounced density differential. As can be seen, the post-biopsy cavity treatment implant **1100** is not formed of a rolled up sheet of material, as is the post-biopsy cavity treatment implant **802** in FIGS. **9** and **10**. Instead, the post-biopsy cavity treatment implant **1100** is formed of solid matrices of collagenous material. It is to be understood that the pore architecture of the first and second portions **1102**, **1104** may be varied at will by, for example, changing the porosity and/or crosslinking of the collagen chains, the pore density, the distribution of pore size, the orientation of the pores and the shape of the pores, to mention a few of the possible pore parameters. By judiciously choosing the pore architectures of the first and second portions **1102**, **1104**, one end of the post-biopsy cavity treatment implant **1100** may be caused to swell at a faster rate than the other end thereof. This is the case illustrated in FIG. **10**.

[0069] Moreover, the cross-sectional characteristics of the post-biopsy cavity treatment implant **1100** may be changed. For example, the first portion **1102** may form a cylindrical inner core of collagenous material having a first predetermined pore architecture and the second portion **1104** may form a cylindrical outer shell around the inner core and may define a second pore architecture. In this manner, the outer surface of the post-biopsy cavity treatment implant **1100** may swell at a different rate (e.g., faster) than the rate at which the inner core swells. Moreover, the pore architectures may be chosen to maximize not only water absorption, but also to promote tissue ingrowth, to facilitate imaging and/or may be tailored to contain and release a pharmaceutical agent at a controllable rate and/or under predetermined conditions. Alternatively, the inner core may be formed of or include a non-collagenous material (such as a polylactic or polyglycolic material, for example) and the outer shell may include a collagenous material, for example. The outer shell may include a solid matrix of collagenous material having a predetermined pore architecture and/or may include wound fibers of collagenous material having a predetermined pore architecture, for example.

[0070] FIG. **12** shows a post-biopsy cavity treatment implant **1200** having predetermined pore architectures, according to another embodiment of the present invention. As shown, the post-biopsy cavity treatment implant **1200**

includes a first portion **1202** and a second portion **1204**, each of which has a predetermined pore architecture. It is to be noted that the present post-biopsy cavity treatment implants may have more than the two portions shown in both FIGS. **11** and **12** (or may define only a single portion). As shown, the post-biopsy cavity treatment implant **1200** is shaped as a substantially rectangular sponge. The first portion **1202** is stacked on the second portion **1204**. As with the embodiment shown in FIG. **11**, the first and second portions may have pore architectures that facilitates tissue ingrowth, wound healing and are readily visualizable and/or recognizable under one or more imaging modalities. The different pore architectures of post-biopsy cavity treatment implants according to embodiments of the present invention may also be chosen so as to maximize the visibility of the interface (such as reference numeral **1203** in FIG. **12**) therebetween under the desired imaging modality such as, for example, ultrasound.

[0071] Post-biopsy cavity treatment implants according to embodiments of the present invention need not be formed as a solid mass of collagen (FIGS. **11**, **12**) or as a rolled up sheet of collagen (FIGS. **9**, **10**). FIGS. **13A**, **13B** and **13C** show various other configurations for the present implant. As shown therein, embodiments of the present invention may include or be formed of a bundle of fibers or fibrils **1302** of (for example) collagenous material having one or more predetermined pore architectures. The pores defined within the collagen matrix of all or some of the fibers are not shown in FIGS. **13A-13C**, but are nevertheless present. The bundle **1300** of fibers shown in FIG. **13A** may be used to form post-biopsy cavity treatment implants by, for example, forming them into a rope-like structure as shown in FIG. **13B**. In the cross-sectional representation of FIG. **13B**, the longitudinal axis of the individual constituent fibers is perpendicular to the plane of the page on which they are printed. Post-biopsy cavity treatment implants may also be formed from the bundle **1300** of FIG. **13A** by cutting (at **1304**, for example) the bundle **1300** into a plurality of sections at an angle that is (for example) perpendicular to the longitudinal axis of the fibers **1302**, so as to form a post-biopsy cavity treatment implant whose constituent fibers run from one end of the post-biopsy cavity treatment implant to the other end thereof, as shown in the detail representation of FIG. **13C**. According to an embodiment of the present invention, a post-biopsy cavity treatment implant may be formed of a volume of collagenous material having a predetermined pore architecture or a combination of several bounded volumes of collagenous materials, each with a predetermined pore architecture. For example, several sponges having the structure shown in FIG. **13C** may be stacked onto each other to define a laminate structure having a layered, composite pore architecture.

[0072] FIG. **13D** shows another embodiment of the present post-biopsy cavity treatment implant. As shown, the implant **1306** includes a first portion **1308** and a second portion **1310**. According to an embodiment of the present invention, the first portion **1308** may include a solid matrix of collagenous material **1312** having a first predetermined pore architecture. The second portion **1310** may include a plurality of fibers or fibrils **1314**. The plurality of fibers may also be formed of or include collagenous material, and this collagenous material may have the same pore architecture as the first portion **1308** or a different pore architecture. The plurality of fibers may be formed or include non-collagenous

material, such as polylactic or polyglycolic acid, for example. In the case wherein the plurality of fibers **1314** are formed of a collagenous material, after implantation in an aqueous environment such as a cavity within a patient, the second portion **1310** may swell at a faster rate than the first portion **1310**, as the constituent fibers **1314** thereof may be exposed to the aqueous environment of the cavity over their entire surface. This swelling rate differential between the first and second portions **1308**, **1310** may serve to further secure the implant **1306** within the cavity. In the case wherein the cavity is relatively dry, the physician may choose to introduce a volume of an aqueous solution, such as saline, into the cavity to speed the swelling of the implant **1306**. The implant **1306** may be formed from a collagen dispersion in a mold configured to form the first portion **1308** and the second portion **1310** and lyophilized. Alternatively, the fibers **1314** may be formed after lyophilization by cutting the implant **1306** so as to form the plurality of fibers **1314**. Alternatively still, the first and second portions **1308**, **1310** may be formed by superimposition of the first and second portions **1310**, **1310**, as discussed above. Other means of forming the first and second portions **1310**, **1312** may occur to those of skill in this art.

[0073] FIG. **13E** shows another embodiment of the present post-biopsy cavity treatment implant. As shown therein, the implant **1316** is similar to the embodiment of FIG. **13D**, but for the addition of a third portion **1318** on another surface of the first portion **1308**. The third portion **1318** may be formed as detailed above relative to second portion **1310**. The pore architecture of the third portion **1318** may be the same as that of the first portion **1308** and the second portion **1310**, or may be different therefrom. It should be noted that various modifications to the embodiments of FIGS. **13D** and **13E** may be envisaged. For example, the embodiment of the implant **1316** of FIG. **13E** may be modified to include additional fibers or fibrils projecting from other surfaces of the first portion **1308**. Other modifications may occur to those of skill in this art, and all such modifications are deemed to fall within the scope of the present invention.

[0074] FIG. **14A** through **14E** show other illustrative embodiments of the post-biopsy cavity treatment implants according to the present invention. As shown in FIG. **14A**, two or more bundles of fibers of collagenous material (for example—the fibers may be made of or include other materials) may be used in the formation of post-biopsy cavity treatment implants according to embodiments of the present invention. As shown, the pores within the fibers of the first bundle **1402** may collectively define a first pore architecture, whereas the pores within the fibers of a second bundle **1404** may collectively define a second pore architecture that is different from the first pore architecture. The two bundles **1402**, **1404** may then be joined together, for example, by re-wetting the bundles, stacking them and lyophilizing the composite structure. The length and diameter of the fibers may be selected and varied at will. The fibers or bundles thereof may even be woven together. From this composite structure, post-biopsy cavity treatment implants may be formed. As shown in FIG. **14B**, the bundles of fibers may be arranged in a cylindrical shape, for example. Such a cylindrical shape may include an inner core **1406** of fibers having a first pore architecture and an outer shell **1408** surrounding the inner core **1406**. The outer shell may include fibers having a second pore architecture that is different from the pore architecture of the inner core **1406**.

FIG. 14C shows a detail of a post-biopsy cavity treatment implant having a first portion **1402** of fibers having a first pore architecture and a second portion **1404** having a second pore architecture, formed, for example, by cutting the composite structure of FIG. 14A at **1410**. Alternatively still, the fibers may be arranged such that the constituent fibers thereof closer to the center of the post-biopsy cavity treatment implant conform to a first pore architecture whereas the outside constituent fibers thereof conform to a second pore architecture that is different from the first pore architecture. As shown in the exploded views of FIGS. 14D and 14E, the post-biopsy cavity treatment implant may be have a layered laminate structure in which sheets formed of fibers (or woven fibers) having a first pore architecture are stacked onto sheets formed of fibers having a second pore architecture. As shown in FIG. 14E, many variations on this theme are possible. As shown therein, the orientation of the fibers (and thus of the pores defined by the collagenous matrix thereof) may be varied. For instance, whereas the fibers of the first (top or outer, for example) portion of the post-biopsy cavity treatment implant may be oriented in a first direction, whereas the fibers of the second (bottom or inner, for example) portion of the post-biopsy cavity treatment implant may be oriented along a direction that is different from the first direction (perpendicular thereto, for example). Imaging such post-biopsy cavity treatment implants within a cavity (such as shown at **126** in FIGS. 9 and 10, for example) using sonography may yield an image in which any fluids contained in the cavity **126** may appear substantially black, because the sound waves travel directly through such anechoic media, and a gradation of visible structures defined by comparatively hypoechoic layers or portions of the post-biopsy cavity treatment implant whose echogenicity is lower than the surrounding area and defined by hyperechoic layers or portions of the post-biopsy cavity treatment implant whose echogenicity is higher than the surrounding area.

[0075] FIGS. 14F and 14G illustrate the stacked structure of a post-biopsy cavity treatment implant, according to further embodiments of the present invention. The embodiments of FIGS. 14F and 14G are similar to the embodiments shown in FIGS. 14D and 14E, but for the structure of the stacked sheets of collagenous material. In FIGS. 14F and 14G, the stacked sheets of collagenous material are not formed of fibers or fibrils, but instead are each formed of a solid mass of collagenous material. The sheets may have the same or different pore architectures. Moreover, the sheets of collagenous material may define pore architectures in which the predominant orientation of the pores is varied. For example, some of the sheets may have a pore architecture in which the pores are predominantly oriented along the y-axis (FIG. 14F) or along the x-axis (FIG. 14G), for example. Alternatively, the constituent sheets of collagenous materials may define pore architectures in which other pore characteristics (size, shape, density, for example) are varied according to a predetermined pattern to influence tissue growth, visualization, etc. The resulting laminate structure may be formed (e.g., molded or cut) in the desired shape of the implant. For example, the resulting laminate structure may then be rolled into a cylindrical shape, as suggested in FIGS. 8 and 9, for example.

[0076] FIG. 15A shows a post-biopsy cavity treatment implant according to another embodiment of the present invention. As shown, the post-biopsy cavity treatment

implant **1500** includes an inner portion **1502** and an outer portion **1504**. The inner portion **1502** may be radiopaque. For example, the inner portion **1502** may be or include a metallic element. The metallic element may have a simple bar shape as shown, or may have a more complex shape such as, for example, a ring. The inner portion **1502** may have other structures to, for example, adhere or hook onto the walls of the cavity **126**. Wound around the inner portion **1502** is one or more fibers **1504** of swellable (collagenous, for example) material having one or more predetermined pore architectures and/or one or more controlled crosslinking densities. The inner portion may be completely encased within the wound bundles of fibers or fibrils **1504** or may be only partially encased, as shown in FIG. 15A. The inner element **1502**, rather than being radiopaque, may have a predetermined echogenicity so as to be immediately recognizable under ultrasound. The inner element **1502**, moreover, may include an inner reservoir configured to contain a volume of therapeutic agent. For example, the inner element **1502** may be bioabsorbable and may be configured to release the contained pharmaceutical agent at a controlled rate. A plurality of fibers **1504** (having the same or different pore architectures) may be wound about the inner element **1502**, the windings thereof being oriented at a given inclination or mutually different inclinations. Moreover, the embodiments of FIGS. 11 through 14E may advantageously be provided with an inner element as shown at **1502** and/or as described immediately above.

[0077] FIG. 15B and the cross-sectional representation of FIG. 15C show another embodiment of the present post-biopsy cavity treatment implant. As shown therein, the implant **1506** may include a first inner portion **1508** forming an inner core and a second outer portion **1510** forming an outer shell around the first inner portion **1508**. Both the first and second portions may be formed of or include a collagenous material. The first portion **1508** may have a first predetermined pore architecture and the second portion **1510** may have a second predetermined pore architecture that is different from pore architecture of the first portion **1508**. For example, the first portion **1508** may have a greater pore density (number of pores per unit volume) than the second portion **1510**. In the exemplary implant **1506** shown in FIGS. 15B and 15C, the pore architecture of the first portion **1508** is such that the collagenous material thereof defines pores that are both smaller and more densely packed than those defined by the collagenous material of the second portion **1510**. Although FIGS. 15B and 15C show the implant **1506** as shaped as a right cylinder, the implant **1506** may be molded into most any shape, to accommodate most any cavity shape. In this manner, the implant may be configured such that its ultimate size and shape after implantation and swelling, substantially matches the size and shape of the cavity in which it is implanted. As shown in the cross-sectional representation of FIG. 15C, the first portion **1508** of the implant **1506** may define an inner reservoir **1512** (created as a void within the first portion **1508** or as a discrete biocompatible reservoir or pouch having a predetermined biodegradability rate). The inner reservoir **1512** may be pre-loaded with a dye/pigment and/or a pharmaceutical agent, as indicated at **1514** in FIG. 15C. The pharmaceutical agent may be configured to slowly release into the cavity **126** as soon as the implant is inserted therein and/or may be configured to require a physician or a RN to pinch or squeeze (or otherwise breach) the implant **1506** to rupture

the reservoir **1512** to release the dye/pigment and/or pharmaceutical agent **1514** contained therein.

[0078] FIG. **15D** shows another embodiment of the implant according to the present invention. The implant **1516** includes a first portion **1508** defining a first pore architecture, such as described above relative to FIGS. **15B** and **15C**. The implant **1516** may include a reservoir **1512**, and the reservoir **1512** may contain a volume of dye/pigment and/or one or more therapeutic agents. Wound around the first portion **1508** is one or more fibers or fibrils of collagenous material defining a second pore architecture that may be different from the first pore architecture. The fibers or fibrils **1520** may completely encase the first portion **1508** or may do so only partially, as shown in FIG. **15D**. FIGS. **15E** and **15F** show another embodiment of the present invention. In this embodiment, the implant **1518** also includes a first portion **1508** defining a first pore architecture, as described relative to FIGS. **15B-15D** above. At least partially surrounding the first portion **1508** are a plurality of fibers or fibrils **1508** that define a second pore architecture that may be different from the first pore architecture. Several layers of such fibers or fibrils **1508** may be disposed around the first portion **1508**, as suggested by the cross-sectional view of FIG. **15F**.

[0079] Most any of the portions or layers of the embodiments disclosed herein may be configured to contain one or more dyes/pigments and/or pharmaceutical agents. The post-biopsy cavity treatment implants discussed herein may be rendered selectively radiopaque by the selective mechanical, chemical or physical incorporation of a radiopaque articles or particles into the collagenous matrix of embodiments of the present post-biopsy cavity treatment implant. For example, the post-biopsy cavity treatment implant may define pores having a predetermined and recognizable architecture and may incorporate some radiopaque compound or particles such as, for example barium sulfate or other commonly used radiopaque or radioactive materials.

[0080] Embodiments of the present invention may also include recognizable articles or substances within the collagenous matrix such as, for example, dyes and/or pigments (i.e., including both synthetic dyes and natural pigments). The dyes/pigments may be incorporated within the collagenous dispersion that forms the constituent layers or portions of the embodiments of the post-biopsy cavity treatment implants disclosed herein. Such dyes/pigments may form mapping compounds that may be gradually released into the body upon implantation of the present post-surgery cavity treatment implant and may form the basis of lymphatic mapping in the future. In this manner, lymphatic mapping may be carried out immediately after a biopsy procedure via elution of the mapping compound (e.g., dyes/pigments and/or radioactive agent) deposited into the collagenous matrix of the implant. In the case wherein a cancer is detected or suspected in the tissue specimen retrieved by the biopsy procedure, this elution of mapping compound from the post-biopsy cavity treatment implant may enable the physician to skip the conventional step of injecting dyes/pigments into the patient, which dye/pigment injection step is conventionally carried out prior to a (sentinel) lymph node status evaluation procedure. Embodiments of the post-biopsy cavity treatment implant according to present invention may include metal-less dyes/pigments as well radiopaque, radioactive or paramagnetic metal-containing dyes/pig-

ments such as, for example, porphyrins and/or porphyrin derivatives (such as chlorophyll and/or chlorophyll derivatives, for example) that are bound to the collagenous matrix. The porphyrins and/or porphyrin derivatives may be tailored, for example, to enhance crosslinking and enhance wound healing and/or to control biodegradation, among other reasons. A metal with paramagnetic properties (such as Mn, for example) may be placed within the porphyrins or porphyrin derivatives so that another mode of recognition may be achieved. Impregnation of the present post-biopsy cavity treatment implant with porphyrins or porphyrin derivatives (for example, copper chlorophyllin) gives the post-biopsy cavity treatment implant a lymphatic mapping functionality due to the elution of the porphyrins or porphyrin derivatives into the surrounding tissue lymphatic drainage system.

[0081] According to other embodiments of the present invention, the present post-biopsy cavity treatment implants may define or include an internal reservoir configured to contain a volume of a mapping compound and/or a beneficial therapeutic agent. Following the biopsy procedure and the subsequent implantation of the present post-biopsy cavity treatment implant having a predetermined pore architecture into the biopsy cavity and following a histopathology report on the excised biopsy specimen, the physician or RN may pinch or squeeze the post-biopsy cavity treatment implant to express the mapping compound(s) and/or agent(s) into the surrounding tissue via lymphatic system to the sentinel node and other lymphatics. In the absence of such squeezing or pinching, the mapping compound and/or therapeutic agent may much more gradually find its way into the surrounding tissue through elution following a gradual biodegradation of the reservoir.

[0082] FIGS. **16-20** are photomicrographs of collagenous matrices having various pore architectures. As shown, the porosity of the collagenous material is not formed by bubbles forced through the collagen dispersion prior to lyophilization thereof. Indeed, it is the structure of the collagen material itself that creates and defines the voids or pores (anechoic regions that appear black in the photomicrographs) within the material. FIGS. **17** and **19** show relatively round pores having a wide size distribution, whereas FIGS. **16** and **18** show a relatively denser collagen matrix having a smaller pore size distribution. FIG. **20** shows an example of a collagenous matrix that is relatively less dense than, for example, the matrix shown in FIG. **18**.

[0083] FIGS. **21-23** are combinations of photomicrographs to illustrate further embodiments of the post-biopsy cavity treatment implants according to the present invention. FIG. **21** shows a post-biopsy cavity treatment implant **2100** that includes a first portion **2102** having a first pore architecture and, stacked thereon, a second portion **2104** having a second pore architecture. As shown, the pore architecture of the first portion **2102** may be characterized as being relatively denser than the pore architecture of the second portion **2104**. Alternatively, the post-biopsy cavity treatment implant **2100** may be structured such that the first portion has a higher porosity (is less dense) than that of the second portion **2104**. The thicknesses of the first and second portions **2102**, **2104** may be varied at will. More than two layers of collagenous material may be provided.

[0084] FIG. **22** shows a post-biopsy cavity treatment implant **2200** having a graduated porosity profile. Such a

post-biopsy cavity treatment implant **2200** may be formed by lining up a plurality of collagen matrices having of progressively lower densities. That is, matrix **2202** has the highest density (amount of collagen per unit volume), matrix **2204** has the next highest density, matrix **2206** has the next to lowest porosity and matrix **2208** has the lowest porosity of the entire post-biopsy cavity treatment implant **2200**. Alternatively, the degree to which each matrix is crosslinked may be varied and controlled. For example, each matrix may be crosslinked to a different degree through the use of, for example, glutaraldehyde. For example, matrix **2202** may be configured to have about 0.0085% glutaraldehyde, matrix **2204** may be configured with about 0.0075% glutaraldehyde, matrix **2206** may be configured with about 0.0065% glutaraldehyde and matrix **2208** may be configured with about 0.0055% glutaraldehyde, for example. Other concentrations are possible, as are different reagents. After superimposing all four such matrices **2202**, **2204**, **2206** and **2208**, a (in this case, piece-wise linear) cross-linking and/or porosity gradient may be achieved across the embodiment of the present post-biopsy cavity treatment implant shown at **2200**.

[0085] FIG. 23 shows a composite post-biopsy cavity treatment implant **2300** having a more complex structure, according to another embodiment of the present invention. The post-biopsy cavity treatment implant **2300** includes three distinct collagen matrices, as shown at **2302**, **2306** and **2308**. As shown, each of the matrices **2302**, **2306** and **2308** has a unique pore architecture. Indeed, the portion of the post-biopsy cavity treatment implant referenced at numeral **2302** has a dense appearance, in which the pores have a high aspect ration and are aligned substantially parallel to the length of the device **2300**. The post-biopsy cavity treatment implant **2300** also includes a second portion **2304** that includes two unique collagenous matrices referenced at **2306** and **2308**, each having different pore architectures. Whereas matrix **2306** features a wide distribution of pore shapes and sizes, matrix **2308** features comparatively larger, generally rounder pores than those of matrix **2306**. Each of these matrices **2302**, **2306** and **2308** may have a unique ultrasonic or x-ray signature and/or contain dyes/pigments or radiopaque materials or compounds. Moreover, not only may the various matrices be visible under selected modalities, the interfaces therebetween may also provide the physician with position and orientation information of the post-biopsy cavity treatment implant within the cavity. Indeed, there are distinct interfaces between dissimilar materials between matrices **2302** and **2306**, between matrices **2302** and **2308** as well as a distinct interface between adjoining matrices **2306** and **2308**, each of which may be readily visible under, for example, ultrasound. It is to be noted that the interfaces between the external surfaces of all three matrices **2302**, **2306** and **2308** with the surrounding tissue may also provide the physician with additional visual clues as to the position and orientation of the post-biopsy cavity treatment implant **2300** within the cavity in which it is implanted. The interfaces described herein, as well as the different rates of swelling may be achieved through control of the porosity and/or as through the control of crosslinking. A single post-biopsy cavity treatment implant may include constituent portions controlled to have a predetermined pore architecture and/or predetermined portions having controlled crosslinking. Although the irregular closed features within the drawings are intended to suggest pores of various configurations and densities, they are alternatively intended

to indicate crosslinking. Therefore, illustrated differences in these irregular closed features between adjacent portions of an implant may also be interpreted as being differences in crosslinking densities between adjacent portions in the implant.

[0086] Use of the post-biopsy cavity treatment implants disclosed herein is not limited to filling post biopsy cavities. Indeed, the present post-biopsy cavity treatment implants also find utility in the correction of defects caused by poorly healed cavities, whatever their origin or cause. The present post-biopsy cavity treatment implants may be placed in cavities in which it is desired that the collagen matrices be replaced, over time, with (human or animal) autogenous tissue. Hence, the embodiments of the present invention may be used for the repair of tissue that has been damaged due to tissue removal, thereby providing a favorable tissue scaffold in which autogenous tissue may infiltrate and grow. In addition, embodiments of the post-biopsy cavity treatment implants according to the present invention may serve to absorb exudates within the cavity, thereby further facilitating the healing process.

[0087] While the foregoing detailed description has described preferred embodiments of the present invention, it is to be understood that the above description is illustrative only and not limiting of the disclosed invention. For example, the post-biopsy cavity treatment implants disclosed herein may be configured to have a unique "signaturing" capability, in which a specific code appears under a given imaging modality. The specific code may be formed within or molded into the structure of the collagen matrix or matrices. For example, a combination of the elements with different crosslinking patterns (e.g., bundles of cylindrical fibers or layers of collagen sponges) may be used for both pattern recognition and predictable filling of the post biopsy procedure cavity. Alternatively, the code may be embodied as a discrete echogenic or radiopaque constituent element of the implant. The codes may confer information to the radiologist or treating physician when viewed under x-ray or ultrasound. Alternatively still, the post-biopsy cavity treatment implants having predetermined pore architectures and/or controlled crosslinking densities according to the disclosed embodiments may include a biocompatibly-sealed integrated circuit that may be interrogated electronically to convey information to the physician. Those of skill in this art may recognize other alternative embodiments and all such alternative embodiments are deemed to fall within the scope of the present invention.

What is claimed is:

1. A method of filling a cavity created by an excisional procedure, the cavity having a predetermined size and shape, the method comprising the steps of:

providing an implant, the implant including at least a first portion and a second portion, the first portion comprising a first collagenous matrix that defines a first selected crosslinking density, the second portion comprising a second collagenous matrix that defines a second selected cross-linking density that is different than the first cross-linking density, the first and second cross-linking densities being selected so as to cause the first and second portions to swell in such a manner that

the implant swells into a size and a shape that is similar to the predetermined size and shape of the cavity when the implant is implanted;

implanting the implant within the cavity through an incision;

adding an aqueous solution to the cavity if the cavity is not sufficiently aqueous to cause the implant to swell, and

closing the incision with the implant implanted in the cavity.

2. The method of claim 1, wherein the providing step is carried out with the first portion comprising a plurality of first collagenous fibers, each of the plurality of first collagenous fibers having the first selected crosslinking density.

3. The method of claim 1, wherein the providing step is carried out with the second portion comprising a plurality of second collagenous fibers, each of the plurality of second collagenous fibers having the second selected cross-linking density.

4. An implant for filling a cavity created by an excisional procedure, the cavity having a predetermined size and a shape, the implant comprising:

a first portion including a first collagenous matrix that defines a first selected crosslinking density, and

a second portion including a second collagenous matrix that defines a second selected cross-linking density that is different than the first cross-linking density,

wherein the first and second cross-linking densities are selected so as to cause the first and second portions to swell in such a manner that the implant assumes a size and a shape that is similar to the predetermined size and shape of the cavity when the implant is implanted.

5. The implant of claim 4, wherein the first portion comprises a plurality of first collagenous fibers, each of the plurality of first collagenous fibers having the first selected crosslinking density.

6. The implant of claim 4, wherein the second portion comprises a plurality of second collagenous fibers, each of the plurality of second collagenous fibers having the second selected cross-linking density.

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