A length extensible implantable device includes a porous member and a longitudinal constraining member. The longitudinal constraining member can constrain at least a portion, up to the entire length, of the porous member in the longitudinal direction. The length of the longitudinally constrained portion can be expanded by applying force to the porous member. The porous member may be a porous tubular member.

FIG. 1A
LENGTH EXTENSIBLE IMPLANTABLE DEVICE AND METHODS FOR MAKING SUCH DEVICES

TECHNICAL FIELD

[0001] This disclosure relates to length extensible implantable devices and methods for making such devices that may be used for providing a lumen for fluid flow in bodily cavities, organs, and vessels within a patient.

BACKGROUND

[0002] Medical devices are frequently used to treat the anatomy of patients. Such devices can be permanently or semi-permanently implanted in the anatomy to provide treatment to a patient. Frequently, these devices, including stents, grafts, stent-grafts, filters, valves, occluders, markers, mapping devices, therapeutic agent delivery devices, prostheses, pumps, bandages, and other endoluminal and implantable devices, are inserted into the body at an insertion point and delivered to a treatment site using a catheter.

[0003] Implantable devices such as grafts and stent-grafts are used in a variety of places in the human body to repair, support, and/or replace anatomical lumens, such as blood vessels, respiratory ducts, gastrointestinal ducts, and the like. Such devices can, for example, provide lumens for blood flow. In such configurations, flexible and durable devices are needed.

[0004] The selection of such implantable devices can pose potential issues. For example, the particularities of the anatomy of one patient may require a device having a different length than a device suitable for another patient. As a result, it may be difficult to determine the necessary size of a device, and, in many instances, the desired device size may be difficult to obtain.

[0005] As such, there is an ongoing need to provide devices, such as grafts and/or stent-grafts, which have adjustable length properties to provide a range of available lengths. Such devices may improve the ability of a treatment provider to properly size a device for the anatomy of a patient.
SUMMARY

[0006] In a first general aspect, a length extensible implantable device for supporting, repairing, and/or replacing a lumen in the body of a patient includes a porous tubular member capable of being extended to a desired length. The porous tubular member comprises a longitudinally compressed portion covered and maintained in the compressed configuration by a longitudinal constraining member. The longitudinal constraining member can comprise a film wrap or a perforated tube, among other structures.

[0007] In various implementations, a length extensible implantable device is formed by longitudinally compressing a porous tubular member, surrounding a portion of the tubular member with a longitudinal constraining member, and releasing the compressive force. In such implementations, the longitudinally constraining member constrains the portion of the tubular member in the longitudinally compressed configuration. The longitudinal constraining member can optionally be secured to the portion of the porous tubular member by, for example, an adhesive. More than one longitudinal constraining member can used. Further, more than one portion of the porous tubular member can be surrounded by one or more longitudinal constraining members.

[0008] The details of one or more embodiments are set forth in the accompanying drawings and the description below. Other features, objects, and advantages will be apparent from the description and the drawings, and from the claims.

BRIEF DESCRIPTION OF THE DRAWINGS

[0009] Figs. 1A - 1C are perspective views of various length extensible implantable devices in accordance with the present disclosure;

[0010] Figs. 2A and 2B are schematic representations of a microstructure of ePTFE material of the prior art;

[0011] Fig. 3 is a perspective view of a length extensible implantable device in accordance with the present disclosure;
[0012] Fig. 4 is a perspective view of another length extensible implantable device in accordance with the present disclosure; and

[0013] Figs. 5A - 5D are perspective views of a length extensible implantable device in various stages.

[0014] Like reference symbols in the various drawings indicate like elements.

DETAILED DESCRIPTION OF THE DRAWINGS

[0015] This disclosure describes devices, systems, and methods that are useful, for example, for repairing, supporting, and/or replacing anatomical lumens. Several implantable medical devices are described herein, and in general any of the features described with respect to a particular device may also be used with any of the other devices described herein. In some examples, one or more features described with respect to a particular device may be added to or included with another device. Also, various combinations or sub-combinations of any of the features described herein may generally be used with any of the devices described herein.

[0016] In general, any of the implantable devices described herein can be delivered to, and deployed at, an in vivo deployment site within a body of a patient using variously minimally invasive surgical techniques. Likewise, these devices may also be surgically implanted via vascular surgical techniques.

[0017] Further, any of the implantable medical devices described herein can be delivered to, and deployed at, an in vivo deployment site within a body of a patient using various minimally invasive transcatheter deployment techniques. For example, any of the implantable medical devices described herein may be releasably attached to a delivery catheter, and the device and delivery catheter may be loaded into a delivery sheath. The delivery sheath may be introduced to the vasculature of the patient and advanced through the vasculature, until a distal end of the delivery sheath is located at or near the target in vivo deployment site. The implantable medical device may be deployed at the deployment site, for example, by retracting the delivery sheath and/or advancing the delivery catheter and the implantable medical device and detaching the implantable medical device from the delivery catheter. The delivery catheter and delivery sheath can then be withdrawn or retracted from the body of the patient.
Any of the implantable medical devices discussed herein can be used to repair, replace, and/or provide support to a body lumen. In various embodiments, implantable medical devices of the present disclosure can be used in a body lumen, including those within the circulatory and gastrointestinal systems.

As used herein, "implantable" means implanted in the body of a patient for more than 29 days.

As used herein, the term "constrain" means: (i) to limit extension, occurring either through self-expansion or assisted expansion, of the length of an implantable device; or (ii) to cover or surround, but not otherwise restrain, an implantable device such as for storage or biocompatibility reasons and/or to provide protection to the implantable device and/or the vasculature.

Figs. 1A-1C describe perspective views of various example length extensible implantable devices 100 comprising a porous tubular member 102 and a longitudinal constraining member 104. Length extensible implantable device 100 can be implanted in the body of a patient either alone or in combination with one or more other components. For example, length extensible implantable device 100 can be combined with a suitable stent, forming a stent-graft. Further, length extensible implant 100 can be combined with other grafts and/or stent-grafts. In other embodiments, the length extensible graft 100 may be provided with a stent (or stent graft) on only one end or alternatively on more than one end or even each end of the length extensible graft 100. A stent graft is considered to be a stent provided with a graft covering all or a portion of the inner or outer surfaces of the stent or both the inner and outer surfaces of the stent. Devices with more than two ends are also contemplated, such as bifurcated devices. Any combination of length extensible implantable device 100 with any suitable medical device is within the scope of the present disclosure.

In various embodiments, porous tubular member 102 comprises a compressible, porous polymeric material, preferably an open celled material. For example, member 102 can comprise a porous expanded polymer, including expanded polytetrafluoroethylene ("ePTFE"), expanded modified PTFE (e.g. coated materials as described further below), expanded copolymers of PTFE, nylons, polycarbonates, polyethylenes, polypropylenes, polyurethanes and the like. These materials may also include materials having a porous fibrillated microstructure. It is
also appreciated that these types of materials may be provided with coatings such as elastomeric coatings and coatings including therapeutic agents (e.g., heparin). Coatings may be provided as surface coatings or alternatively may partially or entirely impregnate the porous materials. Any suitable compressible porous polymer material is within the scope of the present disclosure.

[0023] Porous tubular member 102 can, for example, comprise an ePTFE construct. In various embodiments, porous tubular member 102 comprises a longitudinally extruded and longitudinally expanded ePTFE tube, such as the tubes described in U.S. Pat. Nos. 3,953,566 and 4,187,390. In other embodiments, polymeric tubular member 102 comprises a wrapped ePTFE film tube. For example, member 102 can comprises a tube made from an ePTFE film that has been cigarette wrapped on the surface of a mandrel or, alternatively, has been helically wrapped on the surface of a mandrel. Such ePTFE films of this type can be made generally as taught by U.S. Pat. Nos. 3,953,566 and 4,187,390. Likewise, conventional longitudinally extruded and expanded ePTFE tubes may be usefully reinforced with an external wrap of ePTFE film, typically, a helical wrap. However, any suitable porous ePTFE tubular member is within the scope of the present disclosure.

[0024] In various embodiments, porous tubular member 102 comprises an ePTFE tube having a multiplicity of fibrils which in turn can be connected to a multiplicity of nodes. The microstructure of porous tubular member 102 can comprise a multiplicity of fibrils having a mean fibril length. Mean fibril length can be determined, for example, by examining a photomicrograph of the surface of porous tubular member 102 and by taking the mean of ten measurements made in the predominant direction of the fibrils between nodes connected by fibrils. First, a photomicrograph is made of a representative region of the sample surface, of adequate magnification to show at least five sequential fibrils within the length of the photomicrograph. A series of five measurements are taken along a straight line drawn across the surface of the photomicrograph in the predominant direction of the fibrils followed by a second series of five measurements made along a second line drawn parallel to the first. A measurement constitutes the distance between adjacent nodes connected by at least one fibril. The ten measurements obtained by this method are meant to obtain the mean fibril length of the region.
For example, as illustrated in Figs. 2A and 2B, porous tubular member 102 can comprise a microstructure of nodes 212, 222 interconnected by fibrils 214, 224.

In various embodiments, in the longitudinally uncompressed configuration, porous tubular member 102 can comprise a multiplicity of straight or unbent fibrils 214. Similarly, visual observation of a magnified longitudinal cross section of porous tubular member 102 indicates that a majority of the fibrils straight or unbent.

For example, after longitudinal compression, portion 110 of porous tubular member 102 comprises a multiplicity of bent fibrils 224. Similarly, visual observation of a magnified longitudinal cross section of portion 110 can indicate that a majority of the fibrils 224 connected to nodes 222 are relatively straight or unbent.

In various embodiments, at least a portion of porous tubular member 102 is held in a longitudinally compressed configuration by longitudinal constraining member 104. As illustrated in Figs. 1A-1C, in such configurations, longitudinal constraining member 104 can surround a portion 110 of an abluminal surface of porous tubular member 102 and maintain portion 110 in the longitudinally compressed configuration. In various embodiments, portion 110 is the entire length of porous tubular member 102.

In various embodiments, portion 110 of porous tubular member 102, when compressed to the laterally compressed configuration, comprises a multiplicity of bent fibrils. In such embodiments, the mean fibril length in portion 110 is shorter than the mean fibril length of porous tubular member 102 in the initial, longitudinally uncompressed configuration. Further, visual observation of a magnified surface of portion 110 can indicate that a majority of the fibrils are relatively non-parallel and bent in relation to the longitudinal axis of the tubular member.

Longitudinal constraining member 104 can be capable of rupturing when force is applied in a particular direction. For example, in configurations in which a portion 110 of porous tubular member 102 is held in the longitudinal compressed configuration, applying tension to one or both ends of porous tubular member can cause longitudinal constraining member 104 to rupture. Rupture of longitudinal constraining member 104 can permit portion 110 to extend from the longitudinally compressed configuration to a less compressed configuration having fibrils that are less bent.
In other embodiments, longitudinal constraining member 104 can be ruptured by applying a radial force. For example, a balloon can be used to apply radial force to porous tubular member 102, rupturing longitudinal constraining member 104 and permitting extension of portion 110 to a lesser compressed configuration having fibrils that are less bent.

With reference to Figs. 1A - 1C, in various embodiments, longitudinal constraining member 104 can comprise a variety of different tubular forms. For example, longitudinal constraining member 104 can comprise an ePTFE film. In various embodiments, longitudinal constraining member 104 comprises an ePTFE film having a multiplicity of nodes connected by fibrils, such as those taught by U.S. Pat. Nos. 3,953,566, 4,187,390, and 5,814,405. However, any film suitable of constraining portion 110 of porous tubular member 102 in a longitudinally compressed configuration is within the scope of the present disclosure.

Fig. 1A illustrates a film wrapped around the surface of porous tubular member 102 at a low angle in relation to a longitudinal axis of the porous tubular member. For example, the film can be wrapped between about 0° and 45° relative to the longitudinal axis of porous tubular member 102.

In various embodiments, as illustrated in Fig. 1B, longitudinal constraining member 104 can comprise a film wrapped around the surface of porous tubular member 102 at a higher angle in relation to the longitudinal axis of the porous tubular member. For example, the film can be wrapped between about 45° and 90° relative to the longitudinal axis of porous tubular member 102.

In yet other embodiments, longitudinal constraining member 104 can comprise a tubular member capable of rupturing upon the application of a sufficiently large force. Such a tubular member can comprise a tubular wall having a multiplicity of slits, holes, and/or perforations that facilitate rupturing. As illustrated in Fig. 1C, longitudinal constraining member 104 can comprise, for example, a perforated tube. Although described in relation to particular examples and embodiments, any tubular member capable of maintaining porous tubular member 102 in a longitudinally compressed configuration and rupturing upon application of sufficient force is within the scope of the present disclosure.

As illustrated in the perspective view of Fig. 3, implantable device 100 can comprise a first longitudinal constraining member 104 and a second longitudinal
constraining member 334. For example, first longitudinal constraining member 104 can surround a first portion 110 of porous tubular member 102, and second longitudinal constraining member 334 can surround a second portion 330 of porous tubular member 102.

[0037] In various embodiments, first portion 110 and second portion 330 can comprise at least a part of the same portion, such that second longitudinal constraining member 334 surrounds first longitudinal constraining member 104. For example, the perspective view of Fig. 4 illustrates second longitudinal constraining member 334 surrounding second portion 330 and a part of first portion 110. Any configuration of first and second longitudinal constraining members, including partial or complete overlap of the two constraining members, is within the scope of the present disclosure. Further, the use of any number of longitudinal constraining members is within the scope of the present disclosure.

[0038] First longitudinal constraining member 104 and/or second longitudinal constraining member 334 can optionally be secured to porous tubular member 102, for example, to maintain the longitudinal constraining members in a desired orientation and position relative to porous tubular member 102. For example, first longitudinal constraining member 104 and/or second longitudinal constraining member 334 can be secured to porous tubular member 102 by applying an adhesive to a segment of an abiuminal surface of porous tubular member 102 and/or the inner surface of the longitudinal constraining members. In various embodiments, a thermoplastic polymer adhesive, including a tetrafluoroethylene and perfluoromethyl vinyl ether copolymer, such as those described in U.S. Pat. No. 7,462,675, can be used. In other embodiments, a fluoroelastomer adhesive, such as a FEP, can be used. Any means capable of securing first longitudinal constraining member 104 and/or second longitudinal constraining member 334 to first porous tubular member 102 is within the scope of the present disclosure.

[0039] A method for making a length extensible implantable device of the present disclosure is described as follows. A porous tubular member in a longitudinally uncompressed configuration is obtained and fitted coaxially over a mandrel having an outside diameter the same as or slightly larger than the inside diameter of the porous tubular member. The tubular member is longitudinally compressed by a compressive force so that the length of the tube is reduced to a desired length. A
longitudinal constraining member is placed over at least a portion of the porous tubular member to maintain the portion of the member in the longitudinally compressed configuration. The longitudinal constraining member can optionally be secured to the porous tubular member. The compressive force on the porous tubular member is released, and the longitudinally compressed porous tubular member is removed from the mandrel.

[0040] Figs. 5A - 5D illustrate a porous tubular member in various stages of a method for forming the porous tubular member into a length extensible implantable device. For example, Fig. 5A illustrates porous tubular member 102 in an initial, longitudinally uncompressed configuration. In the longitudinally uncompressed configuration, porous tubular member 102 can comprise a length £1.

[0041] Fig. 5B illustrates porous tubular member 102 after a compressive force is applied. As the compressive force is applied, porous tubular member 102 is compressed from the initial, longitudinally uncompressed configuration to the longitudinally compressed configuration. In the longitudinally compressed configuration, porous tubular member 102 has a length £2, which is shorter than £1. In various embodiments, porous tubular member 102 is biased such that, upon release of the compressive force, it will at least partially back to £1.

[0042] In various embodiments, £2 can comprise a length that is between about 50% and 75% of £1, such that compression from £1 to £2 reduces the length of porous tubular member 102 to between 50% and 75% of its uncompressed length. In other embodiments, £2 can comprise a length that is between about 25% and 50% of £1. In yet other embodiments, £2 can comprise a length that is between about 5% and 25% of £1. Any relationship between £2 and £1 is within the scope of the present disclosure.

[0043] After porous tubular member 102 is compressed to a desired length £2, at least one longitudinal constraining member 104 is applied around the abluminal surface of porous tubular member 102 to maintain at least a portion 110 of porous tubular member 102 in the longitudinally compressed configuration. For example, Fig. 5C illustrates porous tubular member 102 covered by longitudinal constraining member 104. In various embodiments, and as illustrated in Fig. 5C, portion 110 covered by longitudinal constraining member 104 can comprise the entire length (£2).
of porous tubular member 102. In other embodiments, portion 110 is less than the entire length of porous tubular members.

[0044] In various embodiments, longitudinal constraining member 104 comprises a film. In such embodiments, the film is wrapped around portion 110 of porous tubular member 102 in the longitudinally compressed configuration. As previously discussed, the film can be wrapped at a relatively low (about 0° to 45°) or a relatively high (about 45° to 90°) wrap angle relative to a longitudinal axis of porous tubular member 102. The film can also be wrapped at multiple angles, such as embodiments in which multiple layers of film are wrapped in multiple directions along the abluminal surface of porous tubular member 102.

[0045] In other embodiments, longitudinal constraining member 104 comprises a tubular element, such as a perforated tube. In such configurations, the tubular element is fitted along the surface of portion 110 of porous tubular member 102 in the longitudinally compressed configuration.

[0046] Longitudinal constraining member 104 can optionally be secured to porous tubular member 102. For example, an adhesive can be applied to the abluminal surface of porous tubular member 102. In other examples, an adhesive can be applied to the inner surface of longitudinal constraining member 104. However, as mentioned above, any manner of securing a longitudinal constraining member to a porous tubular member is within the scope of the present disclosure.

[0047] After portion 110 of porous tubular member 102 has been secured in the longitudinally compressed configuration by at least one longitudinal constraining member 104, the compressive force used to shorten porous tubular member 102 from ℓ1 to ℓ2 can be relieved while longitudinal constraining member 104 maintains portion 110 in a compressed configuration, forming length extensible implantable device 100. If portion 110 comprises less than the entire length of porous tubular member 102, upon release of the compressive force, the segment of porous tubular member 102 not constrained can expand to its original length, leaving only portion 110 in the longitudinally compressed configuration. In embodiments in which the entirety length of porous tubular member 102 is covered by longitudinal constraining member 104 (in other words, where portion 110 is equal to ℓ2), all of porous tubular member 102 remains in the longitudinally compressed configuration.
In various embodiments, a second porous tubular member can be positioned around portion 110, portion 330, and/or all of porous tubular member 102. In such configurations, longitudinal constraining members 104 and/or 334 are sandwiched between porous tubular member 102 and a second porous tubular member, such that longitudinal constraining members 110 and/or 330 cannot be seen when visually examining the outer surface of length extensible implantable device 100.

After length extensible implantable device 100 is formed, it can be adjusted and configured for use within the body of a patient. In various embodiments, as illustrated in Fig. 5D, the length of length extensible implantable device 100 can be expanded to a length \( t3 \), which is greater than \( \frac{12}{10} \) and less than or equal to the length of porous tubular member 102 in the initial, laterally uncompressed configuration (having a length of \( t1 \)). In various embodiments, as force is applied to porous tubular member 102, longitudinal constraining member 104 can rupture or tear, forming one or more ruptures 540. Once sufficient force is applied, porous tubular member 102 can continue expanding until it has expanded back to \( t1 \).

In various embodiments, portion 110 of porous tubular member 102 can be extended from the longitudinally compressed configuration to a longer length (such as \( t3 \)) by applying a force parallel to the longitudinal axis of porous tubular member 102. In other embodiments, portion 110 of porous tubular member 102 can be extended from the longitudinally compressed configuration to \( t3 \) by applying a radial force to portion 110.

For example, a treatment provider can determine a desired length of length extensible implantable device 100 before implanting the device into the vasculature of a patient. In other cases, the treatment provider can determine the desired length of length extensible implantable device 100 during the course of implanting the device into the vasculature and delivering the device to a treatment area of the patient.

Several characteristics and advantages have been set forth in the preceding description, including various alternatives together with details of the structure and function of the devices and methods. The disclosure is intended as illustrative only and as such is not intended to be exhaustive or limiting. It will be
evident to those skilled in the art that various modifications may be made, especially
in matters of structure, materials, elements, components, shapes, sizes, and
arrangements of parts including combinations within the principles described herein,
to the full extent indicated by the broad, general meaning of the terms in which the
appended claims are expressed. To the extent that these various modifications
depart from the spirit and scope of the appended claims, they are intended to be
encompassed therein.
WHAT IS CLAIMED IS:

1. An implantable device comprising:
   a porous member comprising a microstructure including a multiplicity of fibrils, wherein the porous member comprises a longitudinally uncompressed configuration and a longitudinally compressed configuration, wherein a mean fibril length in the longitudinally uncompressed configuration is greater than a mean fibril length in the longitudinally compressed configuration; and
   a first longitudinal constraining member covering and maintaining at least a portion of an abluminal surface of the porous member in the longitudinally compressed configuration.

2. The implantable device of claim 1, further comprising a second porous member covering the longitudinal constraining member.

3. The implantable device of claim 1, wherein the porous member is a tubular member.

4. The implantable device of claim 1, wherein the first longitudinal constraining member comprises a porous ePTFE film.

5. The implantable device of claim 4, wherein the porous ePTFE film is wrapped over the at least a portion of the abluminal surface of the porous member at an angle relative to a longitudinal axis of the porous member.

6. The implantable device of claim 1, wherein the first longitudinal constraining member covers more than one portion of the abluminal surface of the porous member.

7. The implantable device of claim 1, wherein the first longitudinal constraining member covers the entire abluminal surface of the porous member.

8. The implantable device of claim 1, wherein the microstructure of the porous member includes a multiplicity of nodes.
9. The implantable device of claim 1, further comprising at least a second longitudinal constraining member.

10. The implantable device of claim 9, wherein the first and second longitudinal constraining members cover different portions of the abluminal surface of the porous member.

11. The implantable device of claim 1, wherein the longitudinal constraining member is secured to the at least a portion of the abluminal surface of the porous member by an adhesive.

12. The implantable device of claim 1, wherein at least a segment of the first longitudinal constraining member is ruptured by applying a longitudinal force to the longitudinal constraining member and the porous member, wherein upon rupturing, at least a portion of the porous member covered by the segment of first longitudinal constraining member longitudinally extends.

13. A method for making an implantable device comprising:
   longitudinally compressing a porous member to a longitudinally compressed configuration, wherein a microstructure of the porous member includes a multiplicity of fibrils, wherein a mean fibril length in the longitudinally compressed configuration is less than a mean fibril length prior to longitudinally compressing the porous member; and
   wrapping a film around at least a portion of a length of an abluminal surface of the porous member when the porous member is in the longitudinally compressed configuration, wherein the film maintains the portion of the abluminal surface of the porous member in the longitudinally compressed configuration.

14. The method of claim 13, wherein the film is secured to the at least a portion of the abluminal surface of the porous member by an adhesive.

15. The method of claim 14, wherein the adhesive is a thermoplastic polymer.
16. The method of claim 15, wherein the adhesive is a tetrafluoroethylene and perfluoromethyl vinyl ether copolymer.

17. The method of claim 14, wherein the adhesive is a fluoroelastomer.

18. The method of claim 17, wherein the adhesive is FEP.

19. The method of claim 13, wherein the film is secured to the at least a portion of the abluminal surface of the porous member by heating the porous member after wrapping the film around at least a portion of the abluminal surface of the porous member.

20. The method of claim 13, wherein the film comprises a porous ePTFE film comprising a plurality of nodes interconnected by fibrils, wherein the nodes are generally aligned in parallel.

21. The method of claim 20, wherein the step of wrapping a film around at least a portion of the abluminal surface of the porous member comprises wrapping the film such that the nodes are generally perpendicular to a longitudinal axis of the porous member.

22. The method of claim 20, wherein the step of wrapping a film around at least a portion of the abluminal surface of the porous member comprises wrapping the film such that the nodes are generally parallel to a longitudinal axis of the porous member.

23. The method of claim 13, wherein the film is wrapped around more than one portion of the abluminal surface of the porous member.

24. The method of claim 13, wherein the film surrounds the entire length of the abluminal surface of the porous member.
25. The method of claim 13, wherein multiple layers of film are wrapped around the abluminal surface of the porous member.

26. The method of claim 25, wherein the multiple layers of film are wrapped around more than one portion of the abluminal surface of the porous ePTFE member.

27. The method of claim 13, wherein the porous member is an ePTFE tubular member.

28. A method of treating a target region of a vasculature of a patient comprising:
   providing a porous member having a longitudinally compressed configuration and a longitudinally uncompressed configuration and a longitudinally constraining member covering and maintaining at least a portion of an abluminal surface of the porous member in the longitudinally compressed configuration;
   applying tension in a longitudinal direction to the porous member to rupture at least a segment of the longitudinal constraining member and elongating at least a portion of the porous member; and
   inserting the porous member into a vasculature of a patient.

29. The method of claim 28, wherein the porous member comprises a plurality of fibrils and wherein a mean fibril length in the longitudinally uncompressed configuration is greater than a mean fibril length in the longitudinally compressed configuration.

30. The method of claim 28, wherein the longitudinally constraining member comprises a film wrapped around at least a portion of the abluminal surface of the porous member at an angle relative to a longitudinal axis of the porous member.

31. The method of claim 30, wherein the film comprises a porous ePTFE film comprising a plurality of nodes interconnected by fibrils.
32. The method of claim 30, wherein the film is wrapped around more than one portion of the abluminal surface of the porous member.

33. The method of claim 28, wherein the longitudinal constraining member surrounds the entire length of the abluminal surface of the porous member.

34. An implantable device comprising:
   a porous ePTFE member comprising a longitudinally compressed portion including bent fibrils; and
   a constraining member covering at least a portion of the longitudinally compressed portion thereby maintaining that portion in a longitudinally compressed configuration.

35. The implantable device according to claim 34 wherein the porous ePTFE member is a tubular member.

36. The implantable device of claim 34, wherein the constraining member comprises a porous ePTFE film.

37. The implantable device of claim 36, wherein the porous ePTFE film has a plurality of nodes interconnected by fibrils, wherein the nodes are generally aligned so as to be substantially parallel to each other.

38. The implantable device of claim 34, wherein upon application of an elongating force to the porous ePTFE member, the constraining member ruptures and permits elongation of the compressed portion, wherein upon release of the elongating force, the elongated compressed portion recovers to a shorter length.

39. The implantable device of claim 35, wherein the constraining member is a perforated sleeve.
40. The implantable device of claim 35, wherein the constraining member comprises a film wrapped around at least a portion of an abluminal surface of the porous ePTFE member at an angle relative to a longitudinal axis of the porous ePTFE tubular member.

41. The implantable device of claim 34, further comprising a stent.

42. The implantable device of claim 34 wherein the porous ePTFE member comprises the longitudinally compressed portion and a longitudinally uncompressed portion.

43. The implantable device of claim 42, wherein a mean fibril length in the longitudinally uncompressed portion is greater than a mean fibril length in the longitudinally compressed portion.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. A61F2/06 B29D23/00

According to International Patent Classification (IPC) onto both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

A61F B29D B29K B29C

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
<td></td>
<td>column 1, line 10 - line 11</td>
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<td>column 7, line 1 - line 10</td>
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<td>column 7, line 22 - line 27; figure 5</td>
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<td>column 5, line 47 - page 6, line 5; figures 11,12</td>
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Further documents are listed in the continuation of Box C. See patent family annex.

Special categories of cited documents:

*A* document defining the general state of the art which is not considered to be of particular relevance.

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*L* document which may throw doubts on priority claim(s) one of which is cited to establish the publication date of another citation or other special reason (as specified).

*O* document referring to an oral disclosure, use, exhibition or other means.

*P* document published prior to the international filing date but later than the priority date claimed.

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention.

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone.

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

"A" document member of the same patent family.

Date of the actual completion of the international search

13 February 2015

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