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(54) SELF-EXPANDING STENT DELIVERY SYSTEM WITH TWO SHEATHS

EINFÜHRVORRICHTUNG MIT DOPPELHÜLLE FÜR SELBSTEXPANDIERBAREN STENT

SYSTEME DE POSE DE STENT A AUTO-EXPANSION A DEUX GAINES

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Description

Field of the Invention

[0001] This invention relates to a stent delivery catheter system, such as the kind used in percutaneous transluminal coronary angioplasty (PTCA) procedures. More particularly, it relates to a stent delivery catheter employing two retractable sheaths which may be retracted to release a self-expanding stent, a balloon assisted expandable stent or a balloon expandable stent.

Background of the Invention

[0002] In typical PTCA procedures, a guiding catheter is percutaneously introduced into the cardiovascular system of a patient and advanced through the aorta until the distal end is in the ostium of the desired coronary artery. Using fluoroscopy, a guide wire is then advanced through the guiding catheter and across the site to be treated in the coronary artery. An over the wire (OTW) balloon catheter is advanced over the guide wire to the treatment site. The balloon is then expanded to reopen the artery. The OTW catheter may have a guide wire lumen which is as long as the catheter or it may be a rapid exchange catheter wherein the guide wire lumen is substantially shorter than the catheter. Alternatively, a fixed wire balloon catheter could be used. This device features a guide wire which is affixed to the catheter and cannot be removed.

[0003] To help prevent arterial closure, repaired dissection, or prevent restenosis, a physician can implant an intravascular prosthesis, or a stent, for maintaining vascular patency inside the artery at the lesion. The stent may either be a self-expanding stent, a balloon assisted expandable stent or a balloon expandable stent. For the latter type, the stent is often delivered on a balloon and the balloon is used to expand the stent. The self-expanding stents may be made of shape memory materials such as nitinol or constructed of regular metals but of a design which exhibits self-expansion characteristics.

[0004] In certain known stent delivery catheters, a stent and an optional balloon are positioned at the distal end of the catheter, around a core lumen. The stent and balloon are held down and covered by a sheath or sleeve. When the distal portion is in its desired location of the targeted vessel the sheath or sleeve is retracted to expose the stent. After the sheath is removed, the stent is free to self-expand or be expanded with a balloon.

[0005] In a coronary stent deployment system which utilizes a retractable sheath one problem which is encountered is the interaction of the sheath and the stent upon retraction of the sheath. Typically, as the sheath slides off of the stent, the stent is subjected to potential marring by the sheath. While this problem can be minimized by making the sheath of soft materials, such materials are often unsuitable for use with a self-expanding stent where prolonged storage results in creep deformation of the inner sheath.

Summary of the Invention

[0006] It is desirable to provide a medical device delivery system which provides a protective, non-marring inner sheath for the medical device and is capable of retaining the medical device for brief periods of time and which further has an additional outer sheath over the inner sheath which is capable of retaining the medical device for lengthy periods of time, thereby allowing the device to have a suitable shelf life.

[0007] EP 0 696 447 discloses an apparatus for a deployment release of intraluminal prosthesis. The known apparatus uses in one embodiment a retractable outer sheath together with an inner membrane attached thereof.

[0008] WO 98/39056 discloses a balloon catheter having a first removable sleeve and a second removable outer sleeve positioned over the inner sleeve. The outer sleeve is connected to the inner sleeve. A retraction device is provided for both sleeves.

[0009] The technical problem of the invention is to provide an improved delivery system for a medical device having a first and a second sheath.

[0010] The problem is solved according to claim 1.
at least a portion of inner sheath.

[0014] In one embodiment of the invention, the distal inner sheath is a tear away sheath and the distal outer sheath is retractable by means of an outer sheath retraction device. The outer sheath retraction device extends in a distal direction from the manifold. The distal outer sheath extends from the distal end of the retraction device.

[0015] In another embodiment of the invention, the distal inner sheath is retractable by means of an inner sheath retraction device. The inner sheath retraction device extends in a distal direction from the manifold. The distal inner sheath extends from the distal end of the inner sheath retraction device. The outer sheath retraction device extends in a distal direction from the manifold. The distal outer sheath extends from the distal end of the outer sheath retraction device.

[0016] In another embodiment of the invention, the distal inner sheath is retractable by means of an inner sheath retraction device. The inner sheath retraction device extends in a distal direction from the manifold. The distal inner sheath extends from the distal end of the inner sheath retraction device. The outer sheath retraction device extends in a distal direction from the manifold. The distal outer sheath extends from the distal end of the outer sheath retraction device.

[0017] In all of the embodiments, the delivery system may further comprise the medical device mounted on the medical device mounting region. Among the contemplated medical devices for use with this system are stents and grafts. Desirably, the stent will be self-expanding or a balloon assisted expandable stent.

Brief Description of the Figures

[0018] Figure 1 shows a longitudinal cross-sectional view of a medical device delivery system.

Figure 2 shows a partial cut-away perspective view of the medical device delivery system of Fig. 1.

Figure 3 shows an enlarged view of region 3 of the medical device delivery system of Fig. 1.

Figure 4 shows a longitudinal cross-sectional view of an embodiment of the inventive medical device delivery system.

Figure 5 shows a longitudinal cross-sectional view of an embodiment of the inventive medical device delivery system.

Figure 6 shows a perspective view of a stent for use with the inventive medical device delivery system.

Figure 7 shows a perspective view of a graft for use with the inventive medical device delivery system.

Figure 8 shows a side elevational view of a vena cava filter for use with the inventive medical device delivery system.

Detailed Description of the Invention

[0019] While this invention may be embodied in many different forms, there are described in detail herein specific preferred embodiments of the invention. This description is an exemplification of the principles of the invention and is not intended to limit the invention to the particular embodiments illustrated.

[0020] The present invention provides a medical device delivery system in which the medical device is contained and/or surrounded and/or protected by both an inner sheath and an outer sheath.

[0021] Turning to the figures, Fig. 1 shows one medical device delivery system generally at 110. The medical device delivery system 110 has a proximal end and a distal end and comprises a manifold 120 located at the proximal end of delivery system 110. Extending in a distal direction from manifold 120 is an inner tube 124 having a distal region 128 and a proximal region 132. At the distal region of inner tube 124 is a medical device mounting region 136 for concentrically mounting a medical device thereon. Delivery system 110 further comprises a distal inner sheath 140 attached to inner tube 124 at distal region 128. Distal inner sheath 140 is concentrically disposed about inner tube 124 and covers at least a portion, desirably a substantial portion and more desirably the entirety of medical device mounting region 136 to retain a medical device about the medical device mounting region. Inner sheath 140 is fixedly attached to the medical device delivery system and desirably is mounted to the inner tube. As shown in Fig. 2, distal inner sheath 140 is a perforated 137 or scored tear away sheath. A distal outer sheath 144 is concentrically disposed about inner tube 124 and at least a portion of distal outer sheath 144 is disposed about at least a portion of distal inner sheath 140. The device further comprises an outer sheath retraction device 148 which extends in a distal direction from manifold 120. Distal outer sheath 144 is seen to extend from the distal end of outer sheath retraction device 148.

[0022] Although there are a variety of outer sheath retraction devices that may be used in the practice, as shown in Figs. 1 and 3, a preferred outer sheath retraction device 148 comprises a proximal outer tube 152, a collapsible sheath 156 extending from the distal end of proximal outer tube 152 and a distal outer tube 160. The proximal end of distal outer tube 160 extends from the distal end of the collapsible sheath 156. A slidable pull wire 164 extends from manifold 120 to distal outer sheath 144. In use, the distal end of retraction device 148 moves in a proximal direction upon sliding pull wire 164 proximally thereby retracting distal outer sheath 144. Tear away inner sheath 140 may then open as a result of the force of the expansion of the expandable medical device. Because the tear away inner sheath is fixedly attached to the medical device delivery system, the tear away inner sheath is withdrawn from the body of the medical device delivery system.
The inner sheath retraction device 341 having a distal end of medical device mounting region 336. As in the embodied about inner tube 324 and covers at least a portion sheath 340. Distal inner sheath 340 is concentrically disposed region 332 and a medical device mounting region 336 for concentrically mounting a medical device thereon as in the previous embodiment. Similarly, as in the previous embodiment, delivery system 210 further comprises a distal inner sheath 240. Distal inner sheath 240 is concentrically disposed about inner tube 224 and covers at least a portion of medical device mounting region 236. Unlike in the previous embodiment, delivery system 210 further comprises a inner sheath retraction device 241 having a distal end and a proximal end. Inner sheath retraction device 241, consists of pull collar 243 mounted on proximal end of distal end of inner sheath 240 and a pull wire 245 extending distally from manifold 220 to pull collar 243.

As in the previous embodiment, a distal outer sheath 244 is concentrically disposed about inner tube 224. At least a portion of distal outer sheath 244 is disposed about at least a portion of distal inner sheath 240. Also, the device comprises an outer sheath retraction device 248 comprising a proximal outer tube 252, a collapsible sheath 256 and a distal outer tube 260 as described for the embodiment of Figs. 1-3. The collapsible sheath section of the medical device delivery system is similar to that shown in Fig. 3, differing only in the presence of an additional wire, corresponding to a slidable pull-wire operably associated with the inner sheath. Slidable wire 264 extends from manifold 220 to distal outer sheath 244 and in use, the outer sheath retraction device works in a manner identical to that described for the outer sheath retraction device described above. Alternatively, although not shown in Fig. 4, inner sheath 240 may also be retracted via a collapsible retraction device similar to retraction device 248 used to retract outer sheath 244.

Also shown is an optional medical device in the form of a self-expanding stent 268 mounted on the medical device mounting region 236. In another embodiment, the invention comprises a medical device delivery system shown generally at 310 in Fig. 5. Medical device delivery system 310, as in the previous embodiments, comprises a manifold 320, an inner tube 324 having a distal region 328, a proximal region 332 and a medical device mounting region 336 for concentrically mounting a medical device thereon. Delivery system 310 further comprises a distal inner sheath 340. Distal inner sheath 340 is concentrically disposed about inner tube 324 and covers at least a portion of medical device mounting region 336. As in the embodiment of Fig. 4, delivery system 310 further comprises an inner sheath retraction device 341 having a distal end and a proximal end. Inner sheath retraction device 341, consists of pull collar 343 mounted on proximal end of distal end of inner sheath 340 and a pull wire 345 extending distally from manifold 320 to pull collar 343.

As in the previous embodiments, a distal outer sheath 344 (in sock form) is concentrically disposed about inner tube 324. At least a portion of distal outer sheath 344 is disposed concentrically about at least a portion of distal inner sheath 340. Unlike in any of the previous embodiments, distal outer sheath 344 extends proximally from the distal end of the inner tube and is removable by sliding the distal outer sheath in a distal direction. Distal outer sheath 344 is in contact with distal inner sheath 340. Mounted concentrically about the inner tube and carrying the pull wire is outer tube 360. Although distal outer sheath 344 is depicted in Fig. 5 as being closed at the distal end, it may optionally be open at the distal end. In use, distal outer sheath 344 is removed prior to insertion of the medical device delivery system into the body. A removable sheath such as that disclosed in US 5,800,517 to Anderson et al. may be used.

Also shown is an optional medical device in the form of a self-expanding stent 368 mounted on the medical device mounting region 336.

In another embodiment, not shown, the medical device delivery system is substantially similar to that shown in Fig. 5 differing only in that the retraction device for retraction the inner sheath is a collapsible sheath as shown in Figs. 1 and 3.

In the various embodiments of the invention, suitable manifolds, as are known in the art, may be employed. The manifold must be able to accommodate two retraction mechanisms.

The inner tubes employed in the various embodiments may be made of suitable materials as are known in the art including. Preferably, the inner tubes are made of flexible, but incompressible construction such as a polymer encapsulated braid or coil. Such construction is known in the art. The braid/coil may be comprised of stainless steel encased in a polymer such as Polymide with an inner layer of Teflon.

The pull collars attached to the retractable sheaths may suitably be ringshaped members made of stainless steel affixed to the interior of the retractable sheaths by an appropriate adhesive such as Loctite 4011, a cyanoacrylate. Desirably, the pull collar will be made of a radio-opaque material such as gold.

The outer sheath, desirably will be made of a material which has sufficient strength to contain a self-expanding stent in the stent’s unexpanded configuration. It is desirable that the outer sheath be constructed so as to be creep resistant. It is also highly desirable that the inner sheath be constructed to be less creep resistant than the outer sheath.

Some of the benefits of the present invention may also be realized in a system wherein the outer sheath is made of a thicker material than the inner sheath.

Suitable materials for the outer sheath include...
polyimide, Pebax, polyethylene, Nylon, and metal for the embodiments in which the outer sheath is retractable via a retraction device extending to the manifold. Suitable materials for the sock-like distal outer sheath include polyimide, Pebax, polyethylene, Nylon, and metal. As for the distal inner sheath, suitable materials include PTFE, Pebax, polyurethane, polyethylene, and polyimide for the tear away embodiments and for the retractable distal inner sheath embodiments.

[0037] The invention also contemplates the use of porous materials for the inner and outer sheaths thereby allowing for the inflow of bodily fluids into the medical device mounting region. This can be helpful in priming the medical device by forcing out any air in the region of the medical device. Suitable porous materials include Suitable porous materials include expanded polytetrafluoroethylene (ePTFE), polyester and silicone. Desirably, the materials will have a pore size ranging from 0.01 mm to 5.0 mm.

[0038] Although the tear away sheath has been described as being mechanically released by the force of the expanding medical device, the invention also contemplates the use of a tear away sheath which is hydrolytically released. The sheath may be ‘glued’ shut via a bio-compatible water soluble material. The sheath may then be opened by supplying water thereto so as to dissolved the ‘glue’. Optionally, the glue may be chosen such that it is stable in the presence of fluids at bodily temperatures by dissolves upon exposure to a fluid of slightly elevated temperature such as water at a temperature of 42°C. Alternatively, the sheath may be glued shut via a material which is releasable via actinic energy such as ultraviolet radiation or gamma radiation supplied thereto.

[0039] The distal inner and outer sheaths may be bonded to the inner tube and/or retraction devices by the use of suitable adhesives including Loctite 4011, a cyanoacrylate as well as methacrylate, or H.B. Fuller 3507, a urethane adhesive. Other suitable bonding methods include pressure welding, heat welding and laser welding.

[0040] The invention also contemplates the use of various lubricants on at least a portion of one or more of the inner and outer sheaths to facilitate the relative motion of the inner and outer sheaths upon retraction or removal of the outer sheath. As seen in Fig. 2, distal inner sheath 140 has an inner surface facing the inner tube and an outer surface 138 facing distal outer sheath 144. Similarly, distal outer sheath 144 has an inner surface 146 facing distal inner sheath 140 and an outer surface facing outward. A lubricant may applied to at least a portion of at least one of outer surface 138 of inner sheath 140 or inner surface 146 of outer sheath 144 so as to reduce frictional forces between the two sheathes. The lubricant may be applied selectively to the surfaces or, alternatively, may be applied to the entirety of the surfaces.

[0041] Although the inner surface and outer surface to which lubricants may be applied have been highlighted in Fig. 2, it is understood that the invention provides for the similar use of such lubricants on the outer surface of the inner sheath and the inner surface of the outer sheath in the other embodiments as well.

[0042] In all of the above embodiments, a lubricant may, optionally, be applied to at least a portion of the inner wall and/or outer wall. Suitable lubricants include silicones, PVP (polyvinyl pyrrolidone), PPO (polypropylene oxide) and PEO. Additionally, BioSlide™ coating produced by SciMed made be used as well. BioSlide™ is a hydrophilic, lubricious coating comprising polyethylene oxide and neopentyl glycol diacrylate polymerized in a solution of water and isopropyl alcohol in the presence of a photoinitiator such as azobisisobutynitrile.

[0043] Additional details of the design of embodiments of the inventive medical device delivery system which employ collapsible sheaths, in particular the portion of the device proximal to the inner sheath may be found in the various embodiments disclosed in US 5,534,007 to St. Germain and Olson.

[0044] In addition to the use of a collapsible sheath retraction device for retracting the outer sheath, the invention also contemplates the use of other suitable retraction means as are known in the art including slidably sealed retractable sheaths and midshaft seals as described in co-pending commonly assigned US Patent Application No. 08/722,834 filed September 27, 1996, and a continuation-in-part application 09/071,484 filed May 1, 1998. Other contemplated retraction means include sheaths activated directly by pull-collars as described in US Patent Application No. 09/071,484 filed May 27, 1998, and screw-like retraction devices as described in US Patent 5,201,757 to Heyn et al.

[0045] Although the medical device shown in the figures have all been described as self-expanding stents, other mechanically expandable stents may be used as well, including balloon expandable stents. A perspective view of one suitable stent is shown in Fig. 6 at 668. Other medical devices suitable for delivery with the present delivery system include implants such as grafts and vena cava filters. A suitable graft is shown in Fig. 7 at 768 while a suitable vena cava filter is shown in Fig. 8 at 868.

[0046] As shown in the figures, the medical device delivery systems may further comprise other optional features, as are known in the art, such as bumpers 172, 272, 372 and 472 and markers 176, 276, 376 and 476. Bumpers 172-472 may be made of polyethylene and are affixed to inner tube 124 by adhesive such as H.B. Fuller 3507. Marker bands 176-476 are preferably made of a radio-opaque material such as gold although other materials such as stainless steel may be used as well. The markers are included to aid in positioning and may be affixed to inner tube 124 by adhesive such as Loctite 4011.

[0047] While several specific embodiments of the present invention have been described, the invention is directed more generally toward the inclusion of two sheaths in any other suitable catheter design not specifically described herein including fixed wire, over-the-wire.
and rapid-exchange catheters.

[0048] In the case of the fixed-wire design, the guidewire is fixedly attached to the medical device delivery system. A fixed-wire delivery system is described in U.S. 5,702,364 to Euteneuer et al., and may be suitably modified for use with the inventive medical device delivery system.

[0049] In an over-the-wire embodiment, the inner tube extends proximally to a manifold and a guide wire may be inserted into the inner tube from the proximal end, the guide wire extending to the distal end of the system. The medical device delivery system may then ride on the guidewire.

[0050] Similarly, a rapid exchange delivery system is described in U.S. 5,534,007 to St. Germain et al., and may be suitably modified for use with the inventive medical device delivery system. Specifically, the rapid-exchange version may be realized by terminating the inner tube in a guide wire port in a location along the system distal to the proximal end of the system to allow for insertion of a guide wire therein. In the rapid-exchange embodiment, only a usable length of the medical device delivery system is approximately 135 cm. For a rapid-exchange medical device delivery system, the distance from where the guidewire accesses the inner tube to the distal tip will be approximately 5 cm to 35 cm.

[0051] The above examples and disclosure are intended to be illustrative and not exhaustive. These examples and description will suggest many variations and alternatives to one of ordinary skill in this art.

Claims

1. A medical device delivery system (210, 310) having a proximal end and a distal end comprising:

   • an inner tube (224, 324) having a distal region and a proximal region and a medical device mounting region at the distal region for mounting a medical device thereabout;
   • a distal inner sheath (240; 340) disposed about at least a portion of the medical device mounting region (236, 336);
   • an inner sheath retraction device (241, 341) extending from the distal inner sheath to the proximal end of the medical device delivery system;
   • a distal outer sheath (244, 344) disposed about at least a portion of the medical device and about at least a portion of the distal inner sheath (140, 240, 340),

   characterized in that it further comprises an outer sheath retraction device extending from the distal outer sheath to the proximal end of the medical device delivery system, so that the distal inner sheath and the distal outer sheath are separately retractable from over the medical device.

2. The medical device delivery system of claim 1 wherein the outer sheath is constructed to be creep resistant.

3. The medical device delivery system of claim 1 wherein the inner sheath is constructed to be less creep resistant than the outer sheath.

4. The medical device delivery system of claim 1 wherein the outer sheath is made of a thicker material than the inner sheath.

5. The medical device delivery system of claim 1 wherein the inner sheath is a perforated tear away sheath.

6. The medical device delivery system of claim 1 wherein the inner sheath has an inner surface facing the outer sheath and an outer surface facing outward and a lubricant is applied to at least a portion of at least one of the outer surface of the inner sheath or the inner surface of the outer sheath so as to reduce frictional forces between the two sheathes.

7. The medical device delivery system of claim 1 wherein at least one of the inner sheath and the outer sheath is made of a porous material.

8. The medical device delivery system of claim 1 further comprising a medical device selected from the group consisting of stents, grafts and vena cava filters, the medical device mounted on the medical device mounting region.

9. The medical device delivery system of claim 8 wherein the medical device is a stent.

Patentansprüche

1. Einführvorrichtung für medizinisches Gerät (210, 310), die ein nähles und ein fernes Ende besitzt und die umfasst:

   - ein Innenrohr (226, 326), das einen fernen Bereich und einen nahen Bereich und einen Montagebereich für medizinisches Gerät im fernen Bereich besitzt, um etwa da ein medizinisches Gerät zu montieren,
   - eine ferne innere Hülle (240, 340), die um zumindest einen Teil des Montagebereichs für medizinisches Gerät (236, 336) herum angeordnet ist,
   - eine Vorrichtung zum Rückzug der inneren
Hülle (261, 361), wobei sich die Vorrichtung von der fernen inneren Hülle zum nahen Ende der Einführvorrichtung für medizinisches Gerät erstreckt, und
dadurch gekennzeichnet, dass sie außerdem eine Vorrichtung zum Rückzug einer äußeren Hülle umfasst, wobei sich die Vorrichtung von der fernen äußeren Hülle bis zum nahen Ende der Einführvorrichtung für medizinisches Gerät erstreckt, so dass die ferne innere Hülle und die ferne äußere Hülle getrennt voneinander vom medizinischen Gerät zurückziehbar sind.

2. Die Einführvorrichtung für medizinisches Gerät gemäß Anspruch 1, wobei die äußere Hülle so konstruiert ist, dass sie kriechresistent ist.

3. Die Einführvorrichtung für medizinisches Gerät gemäß Anspruch 1, wobei die innere Hülle so konstruiert ist, dass sie weniger kriechresistent ist als die äußere Hülle.

4. Die Einführvorrichtung für medizinisches Gerät gemäß Anspruch 1, wobei die äußere Hülle aus einem dickeren Material angefertigt ist als die innere Hülle.

5. Die Einführvorrichtung für medizinisches Gerät gemäß Anspruch 1, wobei die innere Hülle eine perforierte Abreißhülle ist.

6. Die Einführvorrichtung für medizinisches Gerät gemäß Anspruch 1, wobei die innere Hülle eine Innenschicht besitzt, die dem Innenrohr zugewandt ist, und eine Außenfläche, die der äußeren Hülle zugewandt ist, und wobei die äußere Hülle eine Innenschicht besitzt, die der inneren Hülle zugewandt ist, und eine Außenfläche, die nach außen gewandt ist, und wobei ein Gleitmittel auf zumindest einen Teil von zumindest einer Außenfläche der inneren Hülle oder auf zumindest eine Innenschicht der äußeren Hülle aufgetragen ist, um die Reibungs Kräfte zwischen den zwei Hüllen zu reduzieren.

7. Die Einführvorrichtung für medizinisches Gerät gemäß Anspruch 1, wobei zumindest die innere Hülle oder die äußere Hülle aus porösem Material hergestellt ist.


Revendications

1. Système de pose de dispositif médical (210, 310) présentant une extrémité proximale et une extrémité distale, comprenant :

- un tube intérieur (224, 324) présentant une région distale et une région proximale et une région de montage de dispositif médical au niveau de la région distale, permettant de monter un dispositif médical autour de celle-ci ;
- une gaine intérieure distale (240, 340) disposée autour d’au moins une partie de la région de montage de dispositif médical (236, 336) ;
- un dispositif de rétraction de gaine intérieure (241, 341) s’étendant de la gaine intérieure distale à l’extrémité proximale du système de pose de dispositif médical ; et
- une gaine extérieure distale (244, 344) disposée autour d’au moins une partie du dispositif médical et autour d’au moins une partie de la gaine intérieure distale (140, 240, 340),
caractérisé en ce qu’il comprend en outre un dispositif de rétraction de gaine extérieure s’étendant de la gaine extérieure distale à l’extrémité proximale du système de pose de dispositif médical, de sorte que la gaine intérieure distale et la gaine extérieure distale peuvent être rétractées séparément de sur le dispositif médical.

2. Système de pose de dispositif médical selon la revendication 1, dans lequel la gaine extérieure est fabriquée de manière à être résistante au fluage.

3. Système de pose de dispositif médical selon la revendication 1, dans lequel la gaine extérieure est fabriquée de manière à être moins résistante au fluage que la gaine extérieure.

4. Système de pose de dispositif médical selon la revendication 1, dans lequel la gaine extérieure est constituée d’un matériau plus épais que la gaine intérieure.

5. Système de pose de dispositif médical selon la revendication 1, dans lequel la gaine intérieure est une gaine perforée déchirable.

6. Système de pose de dispositif médical selon la revendication 1, dans lequel la gaine intérieure pré-
sente une surface intérieure orientée vers le tube intérieur et une surface extérieure orientée vers la gaine extérieure et la gaine extérieure présente une surface intérieure orientée vers la gaine intérieure et une surface extérieure orientée vers l’extérieur et un lubrifiant est appliqué à au moins une partie d’au moins une surface parmi la surface extérieure de la gaine intérieure ou la surface intérieure de la gaine extérieure de manière à réduire les forces de frottement entre les deux gaines.

7. Système de pose de dispositif médical selon la revendication 1, dans lequel au moins une gaine parmi la gaine intérieure et la gaine extérieure est constituée d’un matériau poreux.

8. Système de pose de dispositif médical selon la revendication 1 comprenant en outre un dispositif médical sélectionné dans le groupe constitué des stents, des greffes et des filtres pour veine profonde, le dispositif médical étant monté sur la région de montage de dispositif médical.

9. Système de pose de dispositif médical selon la revendication 8, dans lequel le dispositif médical est un stent.