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(54) **ENDOVASCULAR GRAFT WITH SEPARABLE SENSORS**

ENDOVASKULÄRES IMPLANTAT MIT TRENNBAREN SENSOREN

GREFFE ENDOVASCULAIRE A CAPTEURS SEPARABLES

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Description

[0001] This invention relates to the treatment of body lumens and, more particularly, to the endovascular placement of a prosthetic graft within vasculature for the purpose of repairing the same.

[0002] Ruptured abdominal aortic aneurysms (AAA) are a leading cause of death in the United States. Treatment options to repair AAA include conventional open surgery and implantation of an endovascular graft. Conventional open surgical repair of AAA involves major abdominal surgery with associated high rates of morbidity. Endovascular grafts have been developed to endoluminally bypass abdominal aortic aneurysms through minimally invasive surgery. Many patients that are unacceptable surgical risks for open repairs are eligible for endovascular graft implantation. Deployment of transfemoral, endovascular grafts to treat AAA is appealing for many reasons: avoidance of an abdominal incision, lack of aortic cross clamping, the potential for regional anesthesia, and a shortened hospital stay.

[0003] A bifurcated graft formed from a series of individual components which are intraluminally delivered apart from one another and then assembled to form a fully supported structure is described in US-A-5,676,696. The modular system includes a base member and one or more grafts connected thereto. The base member preferably includes a portion which gradually increases in diameter. A tubular device for inserting the components of the modular system and a method employing the modular system for repairing an abdominal aortic aneurysm are also described.

[0004] Untreated AAA have been shown to continue to expand until rupture, with an associated high mortality rate. Implantation of endovascular grafts have also been associated with high complication rates, including perioperative death, conversion to open repair, the need for further intervention, the need for hemodialysis, a failure to cure the AAA, and wound complications.

[0005] The inability to obtain or maintain a secure seal between the vessel wall and the endovascular graft is a complication unique to endovascular aneurysm exclusion. Because the term "leak" has been associated with aneurysm rupture following conventional surgery, the term "endoleak" has been proposed as a more definitive description of this complication. It is believed that persistent endoleaks result in continued aneurysm expansion, which may eventually lead to aneurysm rupture. Aneurysms that have been successfully excluded have shown a tendency towards a reduction in aneurysm diameter. Failure to properly exclude the aneurysm from systemic arterial blood pressure keeps the patient at risk of impending rupture. Endoleaks have been classified according to the source of the leaks. Current classifications of endoleaks include four categories. Type I endoleaks are "perigraft" or "graft-related" leaks that involve a persistent channel of blood flow due to inadequate or ineffective sealing at the ends of the endovascular graft,

or between overlapping components of a modular system. Type II endoleaks are retrograde flow into the aneurysm sac from patent lumbar arteries, the inferior mesenteric artery, or other collateral vessels. Type III endoleaks result from fabric tears, graft disconnection, or graft disintegration. Finally, Type IV endoleaks are flow through the graft fabric associated with graft wall porosity or permeability. It has been recognized that preoperative patent side branches are not a good predictor of postoperative endoleaks.

[0006] There have been a number of reported cases of aneurysm rupture following implantation of an endovascular graft. Some of the ruptures occurred in patients without a documented endoleak.

[0007] A number of studies have focused on measurement of pressure within the aneurysm sac following implantation of an endovascular graft, both in the human patient, an animal model, or an in vitro model. Properly implanted endovascular grafts have been shown to reduce the pressure within the aneurysm sac while an endoleak, with or without detectable blood flow, continues to pressurize the sac at pressures equivalent to the systemic arterial pressure. Animal studies utilizing a predictable rupturing aneurysm model have shown that non-excluded aneurysms will rupture. Thrombosed aneurysm sacs may still receive pressurization from a sealed endoleak and this continued pressurization keeps the aneurysm at risk for rupture.

[0008] Current methods of patient follow-up include arteriography, contrast-enhanced spiral computed tomography (CT), duplex ultrasonography, abdominal X-ray, and intravascular ultrasound. All of these methods are costly and involve invasive procedures with associated morbidity that may need to be performed in a hospital. None of the imaging methods are completely successful in detecting endoleaks. Therefore, the potential exists for an endoleak to go undetected until eventual rupture. An increase in aneurysm diameter is detectable, and should be considered an indication of endoleak. To avoid aneurysm rupture an increase in aneurysm diameter must be detected in a timely fashion to identify patients in need of corrective endovascular procedures.

[0009] An endovascular graft with the ability to measure pressure within the aneurysm sac and provide feedback to the physician could provide acute confirmation of a procedure and identify those patients with persistent pressurization of their aneurysm, and subsequent risk of rupture. Some physicians are advocating that the follow-up examinations of AAA patients focus on pressure measurements, but that this is not currently clinically feasible. Furthermore, follow-up examinations may be performed in the physician's office as opposed to a hospital. Moreover, clinicians will have a new method to study the pathology of post-endovascularly treated AAA disease.

[0010] Accordingly, there exists a need for an endovascular graft that facilitates non-invasive measurement of pressure, as well as other pertinent parameters, within the aneurysm sac and along the endovascular

graft itself as a means for confirming the success of a procedure as well as identifying patients at risk for aneurysm rupture after the endovascular graft is implanted.

[0011] BP-A-1,132,044 describes a device for providing an indication concerning the pressure existing in an environment, particularly in human or animal tissue. A closed capsule is implanted having a wall which is partly flexible. The capsule is partly filled with a quantity of material which is displaced as a result of movements of the wall. The surface level of the material is detected with respect to one or more radiopaque markers on the capsule so that, by means of repeated observations, changes in the tissue can be detected.

[0012] Likewise, US-B1-6,416,474 describes a biosensor for monitoring pressure or other physical parameters at an aneurysm site which is mountable to a tubular prosthesis that is expandable between contracted and enlarged conditions. A loop, having the biosensor attached thereto, is securable around the prosthesis. An apparatus is used to deliver the loop to the aneurysm site that includes a catheter having a connector on its distal end for detachably securing the loop. The prosthesis is advanced in a contracted state to the aneurysm, and the apparatus, with the loop connected thereto, is advanced to the aneurysm site. The loop and the prosthesis are positioned coaxially with respect to one another, and the prosthesis is expanded towards its enlarged condition, thereby engaging the loop around the prosthesis and engaging the prosthesis with a wall of the blood vessel at the treatment site.

[0013] In EP-A-0,897,690 there is described a device for introduction into a human or animal body to be positioned in an aneurysmal sac in an artery between the wall of the artery and the wall of an endoprosthesis, comprising at least a pressure sensor and a transponder for wireless transmitting data available from the pressure sensor.

[0014] However, providing devices, on an endovascular graft to facilitate the measurement of pertinent parameters poses problems. The measurement device increases bulk, which can significantly effect the delivery profile of the endovascular graft and increase the force necessary to deploy the device, such as jacket or release wire retraction forces. Increased bulk is a significant issue for an endovascular graft. Furthermore, attachment of measurement devices to an endovascular graft may require sutures and the suture knots not only provide increased bulk, but are also potential graft wear points. Additionally, tissue growth around a measuring device attached to an implanted endovascular graft may interfere with its function and inaccurate data may result. The present invention addresses these problems and other needs.

SUMMARY OF THE INVENTION

[0015] According to the present invention there is provided a prosthesis for repairing vasculature, comprising:

a prosthesis component; a plurality of tethers and a plurality of devices having measuring or transmitting capabilities, each device being attached to the prosthesis component by a respective tether, **characterised in that** the devices occupy multiple locations along the prosthesis and at least one tether is bio-reabsorbable.

[0016] The prosthesis or endovascular graft has the ability to be delivered endovascularly and measure pertinent parameters within the lumen in which it is implanted. The endovascular graft has the ability to transmit data about intra-lumen parameters to an external monitoring device. Confirmation of a successful implant procedure is quickly and easily obtained. Patient follow-up is less costly (conducted in the physician office), non-invasive, and more accurate, allowing prompt intervention in those patients most at risk for acute AAA rupture. The invention would also allow for more frequent patient follow-up, increasing the potential to diagnose and treat aneurysms at risk before acute rupture.

[0017] The devices attached to the prosthesis component may include an integrated sensor/transmitter, a sensor, a satellite sensor or a monument device incorporating one or more identifying, observing, transmitting or measuring functions. These devices are attached to the prosthesis component via a bioresorbable suture or tether, The tether functions to place the devices in positions separate from the prosthesis and in desired circumstances, maintain a space between the prosthesis component and the identifying, observing, transmitting or measuring device.

[0018] There is also described an arrangement not falling within the scope of the present claims in which a modular endovascular graft having a main body component and one or more limb components is provided. One or more sensors are attached to the limb component(s). By attaching at least one sensor near the superior end of the limb component and sizing the main body component such that the sensor is adjacent to the aneurysm sac when the endovascular graft is implanted, measurement of pertinent parameters within the aneurysm sac is facilitated. The meaning of the term "adjacent" as used herein encompasses the sensor being located within the aneurysm sac or at a location where the parameters or properties being detected indicate conditions within the aneurysm sac. Measurements of pertinent parameters within the aneurysm, sac may allow early confirmation of a successful procedure and identification of areas of the patient's vasculature at risk for aneurysm rupture, thrombus formation, infection, inflammation or other anomalies without the need for invasive procedures.

[0019] An antenna or other data transmitter and a power source also may be attached to the limb component adjacent to the aneurysm sac, allowing a physician or technician to monitor graft and vessel health without the need for an invasive procedure. The transmitter transmits measurements made by the sensors to a receiver located outside the patient's body. With the main body component unencumbered with sensors, transmitters or power

sources, the bulk of the main body component is minimized and thereby, catheter diameter, jacket retraction and deployment complications are kept to a minimum.

[0020] In another arrangement an endovascular graft is provided that has attached thereto at least one integrated sensor/transmitter device capable of measuring a pertinent parameter and transmitting the measurements to an external monitoring device. Although having more bulk than a sensor, the integrated sensor/transmitter device has less total bulk than a sensor and independent transmitter device, thereby facilitating less total bulk for the endovascular graft.

[0021] Furthermore, the integrated sensor/transmitter device may be designed to allow one or more "satellite" sensors, having no function other than measurement, to be connected thereto. A single integrated sensor/transmitter device and smaller "satellite" sensors facilitate a smaller total bulk than multiple integrated sensor/transmitter devices. Attaching the integrated sensor/transmitter at a central location such as the graft crotch and "satellite" sensors at various locations on the endovascular graft facilitates measurement and transmission to an external monitoring device of pertinent parameters at multiple locations along the endovascular graft and within the lumen. The "satellite" sensors allow a complete profile of pertinent parameters to be obtained and may provide more accurate identification of anomalies. Measurement of pertinent parameters at multiple locations along the endovascular graft or within the aneurysm sac may allow early detection of a defective seal between endovascular graft components, graft wear or changes in aneurysm geometry. The smaller "satellite" sensors may also allow pertinent parameters to be measured from locations on the endovascular graft where local graft bulk is a constraint of the design, such as the graft contra limb or near the superior attachment system that holds the graft in the patient's aorta.

[0022] Additionally, it is contemplated that "satellite" sensors may be attached directly to the lumen of a patient. The integrated sensor/transmitter device also may be attached directly to the lumen or attached to an implanted endovascular graft.

[0023] Moreover, it is contemplated that passive devices, or monuments, which perform no sensing function may be attached to the implanted endovascular graft or directly attached to the aneurysm sac. By tracking the location of the monuments with a monitoring device, changes in the position of the endovascular graft within the lumen or changes in the geometry of the tissue outside the endovascular graft may be detected without the problems of encapsulation or thrombus isolation associated with the measurement of pertinent parameters. Such changes may provide early detection of endovascular graft displacement or aneurysm re-dilation due to an endoleak.

[0024] Sensors with pressure measurement capability may be used to detect pressure changes in the aneurysm sac indicative of graft failure or endoleaks due to an in-

adequate seal between the endovascular graft and the vasculature. Sensors with temperature measurement capability may be used to detect temperature differentials associated with "hot spots" related to inflammation, infection or thrombus formation in the vessel. Sensors with the capability to measure oxygen and other blood constituents such as enzymes, proteins, and nutrients, may be used to detect minute blood flow indicative of endoleak. Sensors with the capability to measure electrical potential or magnetic fields may be used to detect differences in potential associated with areas of the vessel at risk for thrombus formation. Sensors also may be provided to facilitate other sensing applications such as blood oxymetry, blood glucose, blood or fluid flow, biochemical or hormonal balance, blood chemistry, positional data, dynamic displacement data, ocular pressure, respiration, electro physiology, tissue stress, venous return and body acoustics.

[0025] In yet another arrangement sensors are attached to an endovascular graft using one continuous suture. Starting at one location on the sensor, a running stitch around the sensor is used to attach the sensor to the graft fabric, thereby minimizing the number of knots necessary for attachment, in this case a single knot. It is contemplated that the running stitch may start at any location on the sensor depending on the location of the sensor on the endovascular graft. Minimizing the number of knots is advantageous because knots are potential graft wear points and add bulk. Additionally, a single knot attachment design may allow the sensor to be placed close to the graft crotch due to the lack of a knot at the end closest the crotch.

[0026] In another arrangement the sensors may be covered in a coating that either inhibits tissue growth or promotes a known or controlled amount and/or type of tissue growth. Because tissue growth may interfere with the ability of a sensor to perform its measurement function, inhibiting tissue growth or restricting tissue growth to a known type and/or amount may increase the reliability of measurements obtained.

[0027] The invention is applicable to all applications of endovascular grafts to treat aneurysmal segments of blood vessels. Furthermore, the invention and methods disclosed herein may be applied any time it is desired to measure intra-luminal parameters in a non-invasive manner. It is contemplated that the invention may be used with all shapes of endovascular grafts known within the art.

[0028] Other features and advantages of the present invention will become apparent from the following detailed description taken in conjunction with the accompanying drawings, which illustrate, by way of example, the principle of the invention.

55 BRIEF DESCRIPTION OF THE DRAWINGS

[0029]

FIG. 1 is a partial cross-sectional view showing a partially assembled bifurcated endovascular graft implanted across an aneurysm sac;

FIG. 2 is a perspective view of another bifurcated graft with an integrated sensor/transmitter device and "satellite" sensors;

FIG. 3 is a perspective view of a typical integrated sensor/transmitter device;

FIG. 4 is a partial cross-sectional view of an alternate arrangement showing a bifurcated graft implanted across an aneurysm sac into which an integrated sensor/transmitter device and "satellite" sensors have been attached;

FIG. 5 is a schematic view showing an endovascular graft with monuments attached thereto implanted in a patient's body and an external source/receiver;

FIG. 6 is an enlarged view of a portion of graft fabric showing a sensor attached using a single suture and running stitch;

FIG. 7 is a partial cross-sectional view, depicting an identifying, observing or measuring device attached to a graft component by a bio-reabsorbable or hard wire tether or suture; and

FIG. 8 is a partial cross-sectional view, depicting the tether of FIG. 7 partially being reabsorbed.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

[0030] The invention is embodied in a prosthetic endovascular graft having the ability to measure pertinent parameters inside the lumen into which it is implanted and transmit the measurements to a receiver located external to the patient.

[0031] Referring to FIG. 1, there is shown a modular bifurcated endovascular graft 10 of the type known within the art implanted in a body vessel 40 across an aneurysm sac 42 in the area of the contra-lateral 44 and ipsi-lateral 46 iliac arteries using methods known within the art (only the contra-lateral limb is shown). The bifurcated endovascular graft 10 may be assembled in-vivo from a tubular trunk component 20 and two limb components 30. The trunk component 20 has a superior end 22 adapted to be secured above the aneurysm and an inferior end 24 adapted to accept the limb components 30. The limb component 30 has a transmitter 12, power source 14, and sensors 16 attached external the graft material. The transmitter 12, power source 14 and sensors 16 can be integrated into one device as described below. The limb component 30 has a superior end 32 adapted to mate with the trunk component 20 inferior end 24 and an inferior end 34 adapted to be secured to the ipsi-lateral 44 iliac artery.

[0032] The sensors 16 measure pertinent parameters outside the endovascular graft 10 and the power source 14 provides power for the transmitter 12 which transmits the measurements to a receiver (not shown) located outside the patient's body. The transmitter 12, power source

14 and receiver (not shown) may be of any type known in the art of surgical implants or other systems utilizing miniaturized power sources and transmitters. The power source 14 and transmitter 12, for example, may be of the type used in pacemaker technology or passive power sources such as ultrasonic chargeable capacitors.

[0033] One or more sensors 16 are located near the superior end 32 of the limb component 30 but inferior the attachment/sealing area 26 between the trunk component 20 and limb component 30 such that the seal between the trunk component 20 and limb component 30 is not jeopardized. Note that the attachment/sealing area 26 is such that one or more of the sensors 16 on the limb component 30 is adjacent the aneurysm sac 42, thereby facilitating measurement of pertinent parameters within the aneurysm sac 42 without encumbering the trunk component 20 with sensors 16. Note also that the transmitter 12 and power source 14 are attached to the limb component 30 such that they are also adjacent the aneurysm sac 42. Minimizing the bulk of the trunk component facilitates a smaller delivery profile for the endovascular graft 10.

[0034] The sensors 16 may measure pressure, with the measurements used as an aid in endovascular graft 10 placement or to identify anomalies that occur after endovascular graft 10 implantation but before aneurysm rupture occurs. The sensors 16 may detect changes in pressure resulting from blood leakage between the endovascular graft 10 and the vessel wall 40, an endoleak resulting from an inadequate seal between them. Furthermore, the sensors 16 may detect changes in pressure resulting from leakage from an endoleak between the trunk 20 and vessel wall 40. Because sensors 16 are located in the area of the aneurysm sac 42, there may be multiple sensors 16 disbursed over the graft material outer wall since local thrombus or calcification may shield one or more of the sensors 16 from blood flow and render their measurements erroneous. Moreover, the sensors 16 may allow pressure differences throughout the "excluded" aneurysm sac 42 to be mapped. It is contemplated that pressure measurements in the aneurysm sac may be obtained to an accuracy range of +/- 0.133 to 4.400 kPa (+/- 1 to 30 mm Hg) and preferably of +/- 1.333 kPa (+/- 10 mm Hg).

[0035] Alternatively, the sensors 16 may measure temperature. Differences in temperature may identify "hot spots" associated with infection, inflammation, thrombus formation or other anomalies that indicate an increased risk for aneurysm rupture. Methods known in the art of pathology and physiology may be used to relate temperature to changes in the vessel walls within which the endovascular graft is implanted.

[0036] Alternatively, the sensors 16 may detect blood flow by measuring oxygen or other constituents, such as enzymes, proteins and nutrients, which are altered by the presence of blood flow. Such sensors may allow detection of minute blood flow, often missed by conventional imaging modalities, and, therefore, allow endoleaks to

be detected earlier. One method is to obtain a baseline of the constituents upon implantation of the endovascular graft. Thereafter, changes in the amount of the measured constituents may be used to identify anomalies.

[0037] Alternatively, the sensors 16 may measure electrical potential or magnetic field strength. Changes in electrical potential may identify areas of the patient's vasculature that are at risk for thrombus formation. Induced magnetic fields indicate motion at a charged portion of the aneurysm such as would occur from pulsatile pressure.

[0038] Sensors also may be provided to facilitate other sensing applications such as blood oxymetry, blood glucose, blood or fluid flow, biochemical or hormonal balance, blood chemistry, positional data, dynamic displacement data, ocular pressure, respiration, electro physiology, tissue stress, venous return and body acoustics.

[0039] Although shown external the limb component 30 in FIG. 1, it is contemplated that the transmitter 12, power source 14 and sensors 16 may be located internal the graft material of the limb component 30. It is further contemplated that the number of transmitters 12, power sources 14 and sensors 16 shown in FIG. 1 may be varied to meet the requirements of the individual patient. It is further contemplated that sensors 16 which measure different pertinent parameters may be used together. Moreover, the arrangement shown in FIG. 1 may be utilized in any type of endovascular graft implant known in the art.

[0040] Referring to FIG. 2, another arrangement is shown in which an integrated sensor/transmitter 50 is attached to the endovascular graft 110 and "satellite" sensors 52 are attached at various locations on the endovascular graft 110. The integrated sensor/transmitter 50 is capable of measuring a pertinent parameters as well as transmitting measurements to a receiver (not shown) outside that patient's body. The "satellite" sensors 52 only measure pertinent parameters and their measurements are transmitted by the integrated sensor/transmitter, to which they are connected by leads 54. The "satellite" sensors 52 may provide a complete profile of pertinent parameters over the surface of the endovascular graft 110 and, therefore, facilitate better identification of anomalies,

[0041] Because the "satellite" sensors 52 are smaller than the integrated sensor/transmitter 50, the overall bulk of the endovascular graft 110 is smaller than if multiple integrated sensor/transmitter devices 50 were utilized. Furthermore, the smaller "satellite" sensors 52 may be placed such that measurements may be obtained from multiple locations, such as near the superior attachment area or within the aneurysm sac 42. Moreover, the smaller "satellite" sensors 52 facilitate measurements from places on the endovascular graft 110 where local graft bulk is a constraint of the design, such as the contra limb, or near the superior attachment system that holds the graft in the patient's aorta.

[0042] A separate power source 114 may be provided.

The power source can be integrated into the sensor/transmitter as described previously and further below. The leads 54 which connect the integrated sensor/transmitter 50 to the "satellite" sensors 52 may be woven into the graft fabric or attached external to the endovascular graft 110.

[0043] It is contemplated that the location and number of integrated sensor/transmitter devices 50, power sources 114 and "satellite" sensors 52 shown in FIG. 2 may be varied to meet the requirements of the individual patient. In the arrangement shown, the integrated sensor/transmitter device 50 is located at the crotch, thereby allowing it to be near the center of the aneurysm sac 42 (see FIG. 2) while still being located furthest away from the aneurysm wall during shrinkage. It is further contemplated that the arrangement shown in FIG. 2 may be utilized in any type of endovascular graft implant known in the art.

[0044] Moreover, the integrated sensor/transmitter 50 and "satellite" sensors 52 may be of any type known in the art used to measure pressure, temperature, oxygen and other blood constituents, electrical potential or any other pertinent parameter indicative of endovascular graft or lumen health. One such integrated sensor/transmitter 50 to measure pressure is disclosed in U.S. Pat. Application Publication No. 2002/0045921 (Wolinsky et al.), the contents of which are hereby incorporated by reference. Again, other sensing applications may be supported such as blood oxymetry, blood glucose, blood or fluid flow, biochemical or hormonal balance, blood chemistry, positional data, dynamic displacement data, ocular pressure, respiration, electro physiology, tissue stress, venous return and body acoustics.

[0045] FIG. 3 shows a typical integrated sensor/transmitter 50. The integrated sensor/transmitter 50 includes a control chip 55 with a transmitter/receiver, energy exchanger 56, capacitor 57 and sensor 58. An external receiver (not shown) may contain a transducer, computer, LCD display and measurement display devices, such as barometers if the sensor 58 measures pressure. In operation, the transducer in the external receiver charges the capacitor 57 using ultrasonic energy and activates the sensor 58 to measure a pertinent parameter and ultrasonically transmit the measured parameter. The external receiver receives the transmitted pertinent parameter, assesses the measurement and displays the measurement, for example, as a pressure pulse curve.

[0046] Although FIG. 2 shows an integrated sensor/transmitter 50 and "satellite" sensors 52 connected by leads 54 and attached to the endovascular graft 110, alternate arrangements may utilize an integrated sensor/transmitter 50 as the sole implant. Furthermore, the "satellite" sensors 52 may be attached to the vessel 40 with the integrated sensor/transmitter 50 either attached to the vessel 40 or attached to the endovascular graft 110. FIG. 4 shows an endovascular graft 110 implanted across an aneurysm sac 42 into which an integrated sensor/transmitter 50 connected to "satellite" sensors 52 by

leads 54 have been attached. It is contemplated that pressure measurements in the aneurysm sac 42 may be obtained to an accuracy range of +/- 0.133 to 4.000 kPa (+/- 1 to 30 mm Hg) and preferably +/- 1.333 kPa (+/- 10 mm Hg) utilizing either an integrated sensor/transmitter 50 alone or in conjunction with one or more "satellite" sensors 52 or integrated sensor/transmitters 50.

[0047] In an alternate arrangement micro devices having the capability to sense their location relative to each other are either attached to the endovascular graft 110 or attached to the vessel 40. The micro devices may be excited by an external energy source, sense their relative location and transmit the data to a receiver located outside the patient's body. The data may then be interpreted to determine if displacement of the endovascular graft 110 or changes in the geometry of the vessel 40 has occurred. Such changes may provide early detection of endovascular graft 110 displacement or aneurysm 42 re-dilation due to an endoleak. Additional micro devices could be provided for better resolution. Detecting location of the micro devices rather than measuring a pertinent parameter minimizes reliability problems due to encapsulation or thrombus isolation. It is contemplated that the micro devices may be an ultrasonic crystal attached or integral to a MEM chip having the ability to read, interpret and transmit data. The micro devices also may be ultrasonic or other energy reflectors utilized in conjunction with an energy source. Using ultrasound to show or image reflector location would be much easier than creating an ultrasound image of the tissue.

[0048] In a further arrangement devices or monuments which perform no sensing function may be attached to the implanted endovascular graft 110 or directly attached to the aneurysm sac 42. The location of the monuments may be tracked with a monitoring device to detect changes in the position of the endovascular graft 110 within the vessel 40 or changes in the geometry of the tissue outside the endovascular graft 110. It is contemplated that the monitoring device may be located external the patient's body or attached to an implanted endovascular graft. FIG. 5 illustrates an external monitoring device 54 being used to sense the location of an endovascular graft 310 having monuments 60 attached thereto which has been implanted in a patient's body. It is contemplated that the monuments may be electrically passive devices or magnetic sensors. The use of RF or magnets to sense the position of monuments is contemplated.

[0049] In one example, the position of magnetic sensors attached to the aneurysm sac 42 are measured using an external magnet of known magnetic field strength and shape. By moving the external magnet to multiple positions and orientations outside the body and polling the magnetic sensors with ultrasound, a baseline of magnetic sensor positions is established. Subsequent polling of the magnetic sensor positions allows changes in aneurysm sac 42 geometry to be detected.

[0050] In yet another arrangement shown in FIG. 6, sensors 16 or integrated sensor/transmitters 52 are at-

tached to the endovascular graft 110 fabric using one continuous suture 70 with a running stitch, from a starting point 72 on sensor 16, which continues around the sensor 16 and back to an end point 74 (the dashed lines in FIG. 6 indicate where the suture 70 is on the inner diameter of the endovascular graft 110 fabric). Suture loops 71 are provided on the sensor 16 to facilitate attachment. A single knot 75 may be used to join the ends of the suture 70. It is contemplated that the running stitch may start and end at either end of the sensor and utilize more or less stitches depending on the location of the sensor 16 on the endovascular graft 110 and the attachment requirements. Minimizing the number of knots not only reduces bulk but also reduces potential graft wear points and may allow the sensor to be placed close to the graft crotch due to the lack of a knot at the distal end. It is further contemplated that the continuous suture 70 attachment method may be utilized with sensors 16 having suture holes or any other attachment mechanism.

[0051] In yet another arrangement, the sensors 16 may be covered in a coating, such as Teflon or heparin, to inhibit tissue growth or covered in a coating, such as Thrombin, to promote a known or controlled amount of tissue growth. Inhibiting or controlling tissue growth, which may interfere with performance, increases the reliability of the measurements made by the sensors 16.

[0052] With reference to FIGS. 7 and 8, there is shown alternative structure for attaching one or more of the previously described sensor/transmitter devices, sensors, satellite sensors, monuments or other devices 202 to a graft component 204. Although shown attached to a limb component 206 via a tether or suture 208, the identifying, observing, transmitting or measuring devices 202 also can be attached by way of a tether 208 to a main body component 210 or other component. In fact, in certain circumstances, the particular device 202 can be implanted in a vessel wall 220. Moreover, multiple devices 202 can be attached to a medical device in any desired pattern.

[0053] In one arrangement, the tether or suture 208 can embody a hard wire designed to maintain a distance between the identifying, observing, transmitting or measuring device 202 and the graft component 206, 210 itself. In this way, the device 202 can be maintained in an aneurysmal sac 212, for example, allowing the graft 206, 210 to repair vasculature without interference from the device 202. The hard wire tether 208 can be configured to resist any forces that may urge the device 202 against the graft 206, 210, thereby minimizing possible wear on the graft 206, 210 from the device 202.

[0054] In another arrangement, a bio-reabsorbable suture or tether 208 can be employed to attach the identifying, observing, transmitting or measuring device 202 to a graft 206, 210 or other component. After deployment of the graft 206, 210 within vasculature, the graft 206, 210 becomes covered in clot and as the clot organizes, the bio-reabsorbable tether 208 breaks down. After the tether breaks down, the identifying, observing, transmit-

ting or measuring device 202 is left in an aneurysmal sac 212 or in a position separate from the graft. The device 202 is therefore held in place by tissue that is generated as a result of the interventional procedure. Again, the device 202 is held separate from the graft 206, 210 thereby minimizing wear to the graft.

[0055] As suggested, the device 202 can embody any of the previously described devices such as a sensor/transmitter, a sensor or satellite sensor, a monument or a power supply. The device 202 can therefore work independently or in cooperation with any of the disclosed identifying, monitoring, sensing or transmitting devices.

[0056] While particular forms of the invention have been illustrated and described, it will be apparent that various modifications can be made without departing from the scope of the invention. For example, the structures and sensors described can be used with any medical device or in any acceptable area of a patient's body. Accordingly, it is not intended that the invention be limited, except as by the appended claims.

Claims

1. A prosthesis for repairing vasculature, comprising:
 - a prosthesis component (206);
 - a plurality of tethers (208); and
 - a plurality of devices (202) having measuring or transmitting capabilities, each device (202) being attached to the prosthesis component (206) by a respective tether (208), **characterised in that** the devices (202) occupy multiple locations along the prosthesis and at least one tether (208) is bio-reabsorbable, wherein each tether (208) is configured to maintain a distance between the prosthesis component (206) and the respective device (202) after the device (202) is placed within the vasculature.
2. The prosthesis of claim 1, wherein at least one tether (208) is made of suture.
3. The prosthesis of claim 1, wherein the prosthesis comprises first and second prosthesis components (206,210).
4. The prosthesis of claim 3, wherein the second prosthesis component (210) is configured to mate with the first prosthesis component (206) after the first prosthesis component (206) is placed within the vasculature.
5. The prosthesis of claim 3, wherein the first prosthesis component (206) is configured to mate with the second prosthesis component (210) after the second prosthesis component (210) is placed within the vasculature.
6. The prosthesis of claim 1, wherein the prosthesis is a modular graft assembly comprising a plurality of graft components (206,210).
7. The prosthesis of claim 6, wherein only a first graft prosthesis component (206) has the devices (202) attached thereto.
8. The prosthesis of claim 6, wherein the devices (202) are attached to more than one component of the plurality of graft components (206,210).
9. The prosthesis of claim 1, wherein the prosthesis is configured to treat an aneurysm and the tethers (208) maintain the devices (202) within an aneurysm sac.
10. The prosthesis of claim 1, wherein at least one device (202) is a sensor/transmitter assembly.
11. The prosthesis of claim 1, wherein at least one device (202) is a sensor.
12. The prosthesis of claim 1, wherein at least one device (202) is a satellite sensor.
13. The prosthesis of claim 1, wherein at least one device (202) is a monument.
14. The prosthesis of claim 1, wherein at least one device (202) is a power supply.
15. The prosthesis of claim 1, wherein at least one device (202) measures or cooperates to measure or identify blood constituents.
16. The prosthesis of claim 1, wherein at least one device (202) measures or cooperates to measure or identify pressure.
17. The prosthesis of claim 1, wherein at least one device (202) measures or cooperates to measure temperature.
18. The prosthesis of claim 1, wherein at least one device (202) measures or cooperates to measure oxygen.
19. The prosthesis of claim 1, wherein the prosthesis component is bifurcated and includes a first limb and a second limb, at least one device (202) being attached to one of the first and second limbs.

Patentansprüche

1. Prothese zur Gefäßreparatur, welche aufweist:
 - eine Prothesenkomponente (206);

- mehrere Halteseile (208); und mehrere Vorrichtungen (202) mit Mess- oder Sendefähigkeiten, wobei jede Vorrichtung (202) durch ein jeweiliges Halteseil (208) an der Prothesenkomponente (206) befestigt ist, **dadurch gekennzeichnet, dass** die Vorrichtungen (202) sich an mehreren Stellen entlang der Prothese befinden und zumindest ein Halteseil (208) bio-reabsorbierbar ist, wobei jedes Halteseil (208) so ausgebildet ist, dass es einen Abstand zwischen der Prothesenkomponente (206) und der jeweiligen Vorrichtung (202) aufrechterhält, nachdem die Vorrichtung (202) innerhalb des Gefäßsystems angeordnet ist.
2. Prothese nach Anspruch 1, bei der zumindest ein Halteseil (208) aus Nahtmaterial besteht.
 3. Prothese nach Anspruch 1, bei der die Prothese eine erste und eine zweite Prothesenkomponente (206, 210) aufweist.
 4. Prothese nach Anspruch 3, bei der die zweite Prothesenkomponente (210) ausgebildet ist zum Verbinden mit der ersten Prothesenkomponente (206), nachdem die erste Prothesenkomponente (206) innerhalb des Gefäßsystems angeordnet ist.
 5. Prothese nach Anspruch 3, bei der die erste Prothesenkomponente (206) ausgebildet ist zum Verbinden mit der zweiten Prothesenkomponente (210), nachdem die zweite Prothesenkomponente (210) innerhalb des Gefäßsystems angeordnet ist.
 6. Prothese nach Anspruch 1, bei der die Prothese eine modulare Implantatanordnung ist, die mehrere Implantatkomponenten (206, 210) aufweist.
 7. Prothese nach Anspruch 6, bei der nur an einer ersten Implantatprothesenkomponente (206) die Vorrichtungen (202) angebracht sind.
 8. Prothese nach Anspruch 6, bei der die Vorrichtungen (202) an mehr als einer Komponente der mehreren Implantatkomponenten (206, 210) angebracht sind.
 9. Prothese nach Anspruch 1, bei der die Prothese ausgebildet ist zum Behandeln eines Aneurismas und die Halteseile (208) die Vorrichtungen (202) innerhalb eines Aneurismasackes halten.
 10. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) eine Sensor-/Senderanordnung ist.
 11. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) ein Sensor ist.
 12. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) ein Satellitensensor ist.
 13. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) ein Monument ist.
 14. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) eine Stromversorgungsquelle ist.
 15. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) Blutbestandteile misst oder zu deren Messung oder Identifizierung zusammenwirkt.
 16. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) Druck misst oder zu dessen Messung oder Identifizierung zusammenwirkt.
 17. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) die Temperatur misst oder zu deren Messung zusammenwirkt.
 18. Prothese nach Anspruch 1, bei der zumindest eine Vorrichtung (202) Sauerstoff misst oder zu dessen Messung zusammenwirkt.
 19. Prothese nach Anspruch 1, bei der die Prothesenkomponente gegabelt ist und ein erstes Glied und ein zweites Glied enthält, wobei zumindest eine Vorrichtung (202) entweder an dem ersten oder dem zweiten Glied angebracht ist.
- Revendications**
1. Prothèse destinée à réparer la vasculature, comprenant :
 - un composant de prothèse (206) ;
 - une pluralité d'amarres (208) ; et
 - une pluralité de dispositifs (202) ayant des fonctions de mesure ou de transmission, chaque dispositif (202) étant attaché au composant de prothèse (206) par une amarre respective (208), **caractérisée en ce que** les dispositifs (202) occupent de multiples emplacements le long de la prothèse et au moins une amarre (208) est bio-résorbable,
 - dans laquelle chaque amarre (208) est configurée pour maintenir une distance entre le composant de prothèse (206) et le dispositif respectif (202) après que le dispositif (202) est placé dans la vasculature.
 2. Prothèse selon la revendication 1, dans laquelle au moins une amarre (208) est faite de suture.
 3. Prothèse selon la revendication 1, dans laquelle la prothèse comprend un premier et un second com-

- posants de prothèse (206, 210).
4. Prothèse selon la revendication 3, dans laquelle le second composant de prothèse (210) est configuré pour s'emboîter avec le premier composant de prothèse (206) après que le premier composant de prothèse (206) est placé dans la vasculature.
 5. Prothèse selon la revendication 3, dans laquelle le premier composant de prothèse (206) est configuré pour s'emboîter avec le second composant de prothèse (210) après que le second composant de prothèse (210) est placé dans la vasculature.
 6. Prothèse selon la revendication 1, dans laquelle la prothèse est un assemblage de greffe modulaire comprenant une pluralité de composants de greffe (206, 210).
 7. Prothèse selon la revendication 6, dans laquelle seul un premier composant de prothèse à greffer (206) a les dispositifs (202) attachés à celui-ci.
 8. Prothèse selon la revendication 6, dans laquelle les dispositifs (202) sont attachés à plusieurs composants de la pluralité de composants de greffe (206, 210).
 9. Prothèse selon la revendication 1, dans laquelle la prothèse est configurée pour traiter un anévrisme et les amarres (208) maintiennent les dispositifs (202) à l'intérieur d'une cavité anévrismale.
 10. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) est un assemblage de capteur/transmetteur.
 11. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) est un capteur.
 12. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) est un capteur satellite.
 13. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) est une prise.
 14. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) est une alimentation électrique.
 15. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) mesure ou contribue à mesurer ou identifier les constituants sanguins.
 16. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) mesure ou contribue à mesurer ou identifier la pression.
 17. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) mesure ou contribue à mesurer la température.
 18. Prothèse selon la revendication 1, dans laquelle au moins un dispositif (202) mesure ou contribue à mesurer l'oxygène.
 19. Prothèse selon la revendication 1, dans laquelle le composant de prothèse présente une bifurcation et comporte une première jambe et une seconde jambe, au moins un dispositif (202) étant attaché à l'une de la première et de la seconde jambes.

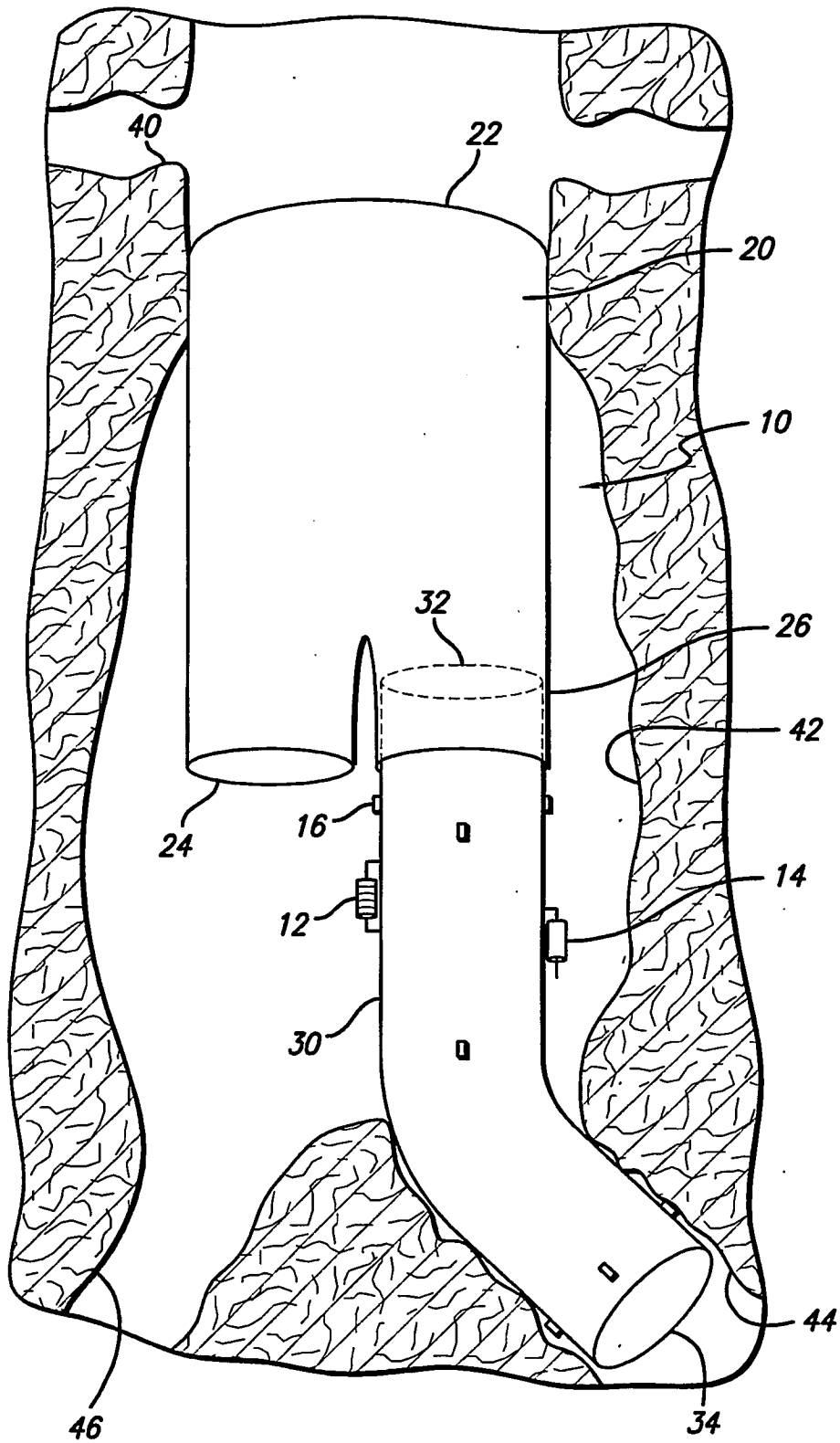


FIG. 1

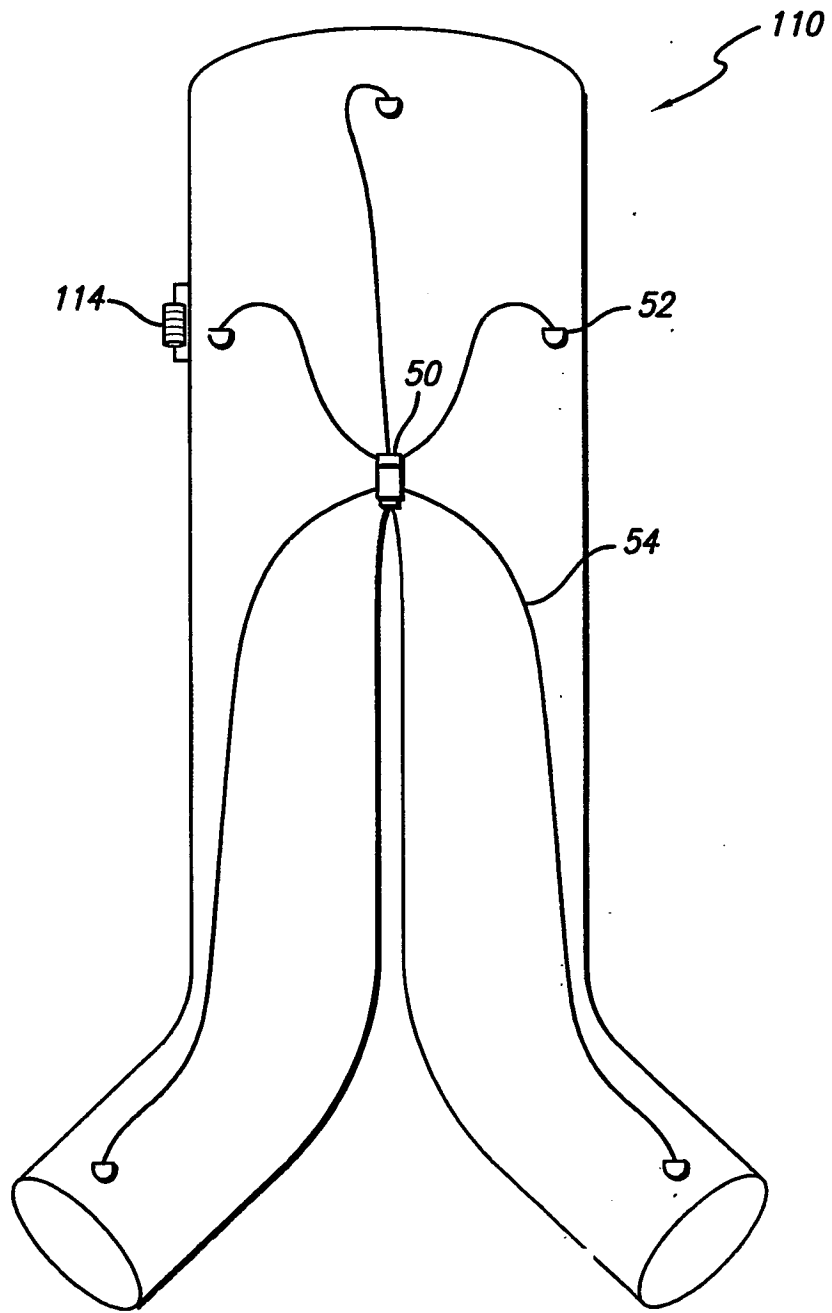


FIG. 2

FIG. 3

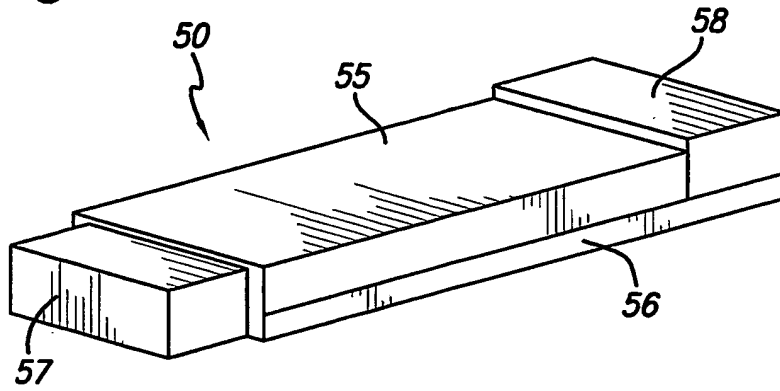
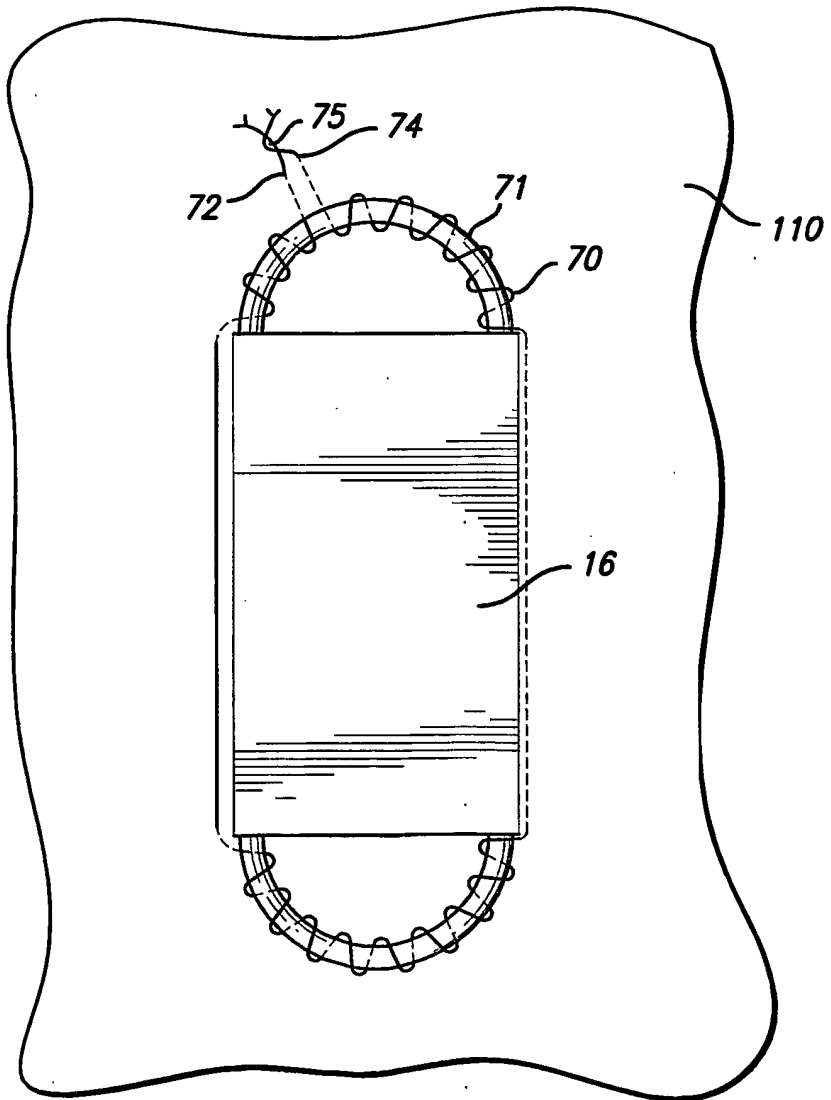


FIG. 6



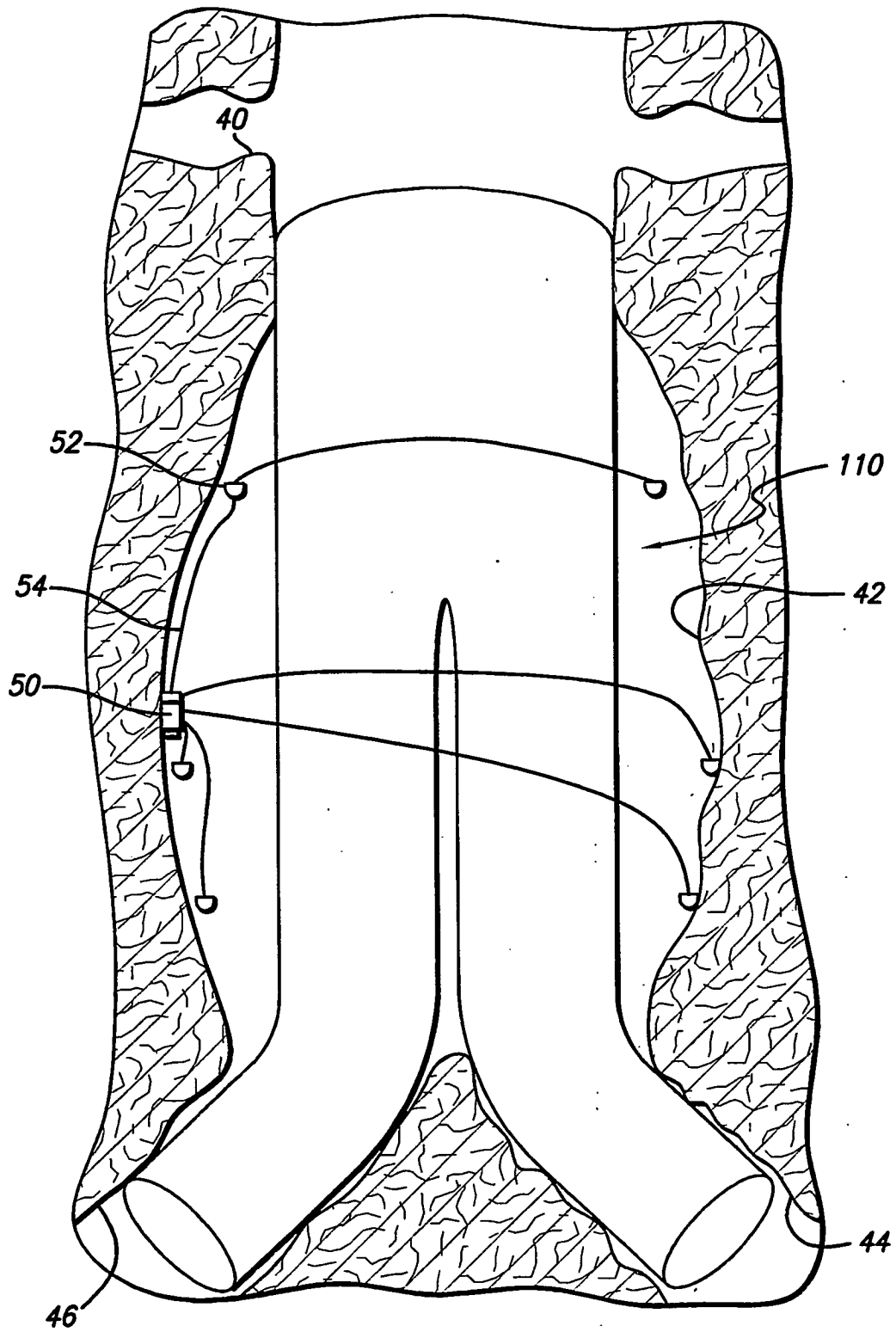


FIG. 4

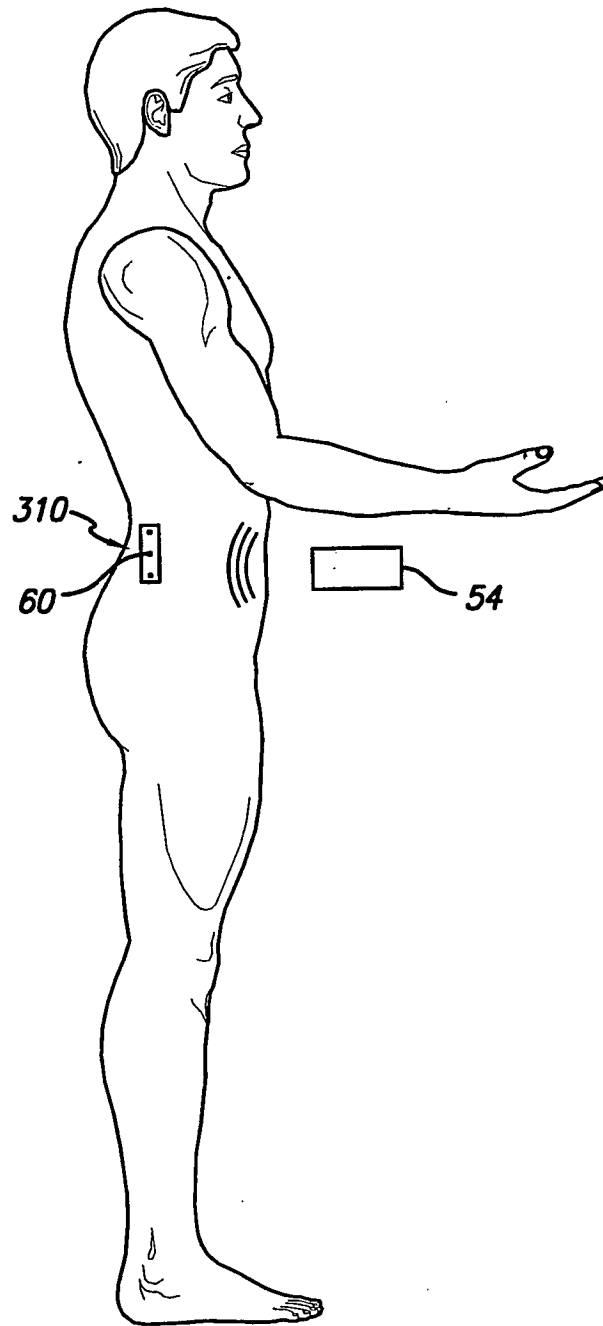


FIG. 5

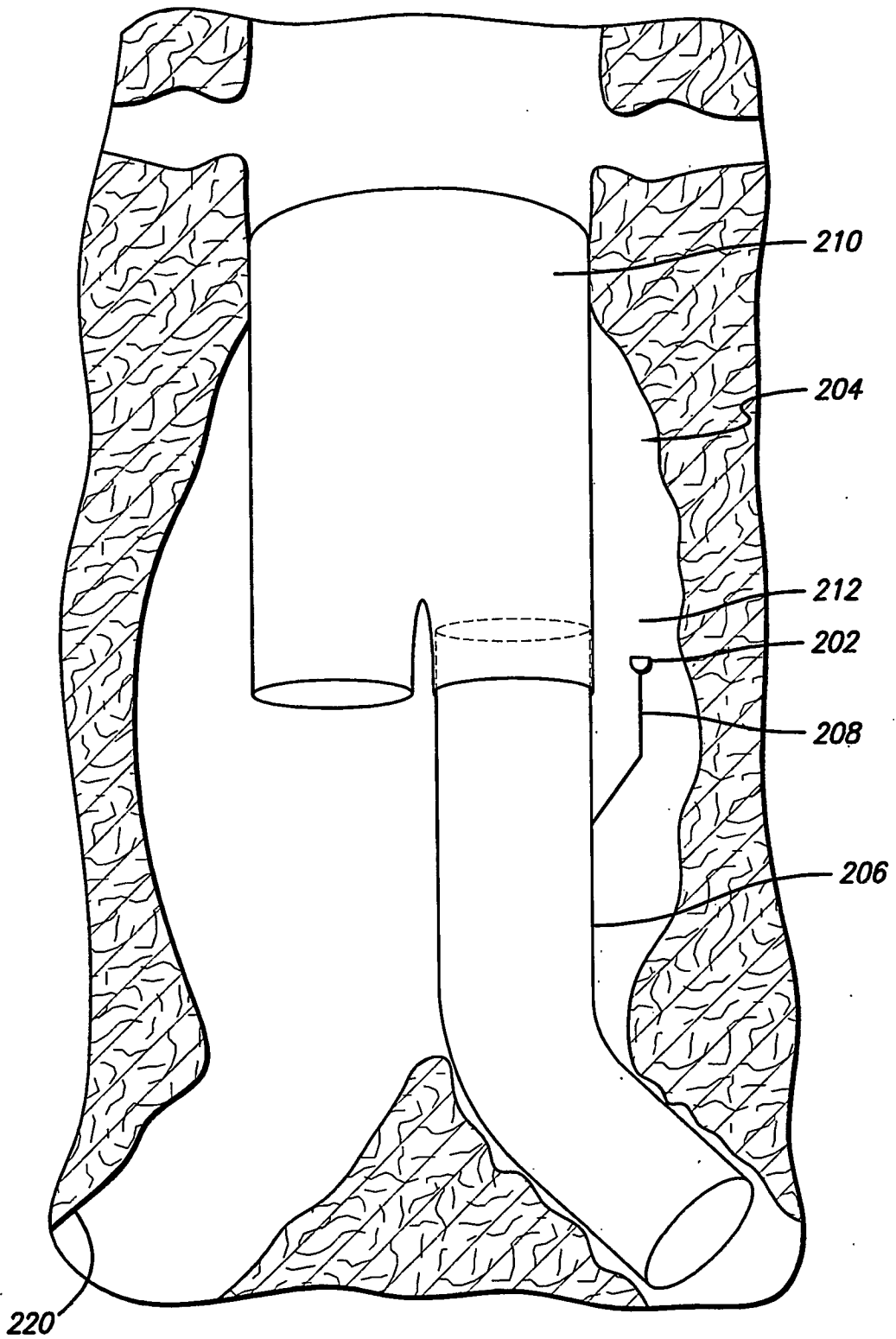


FIG. 7

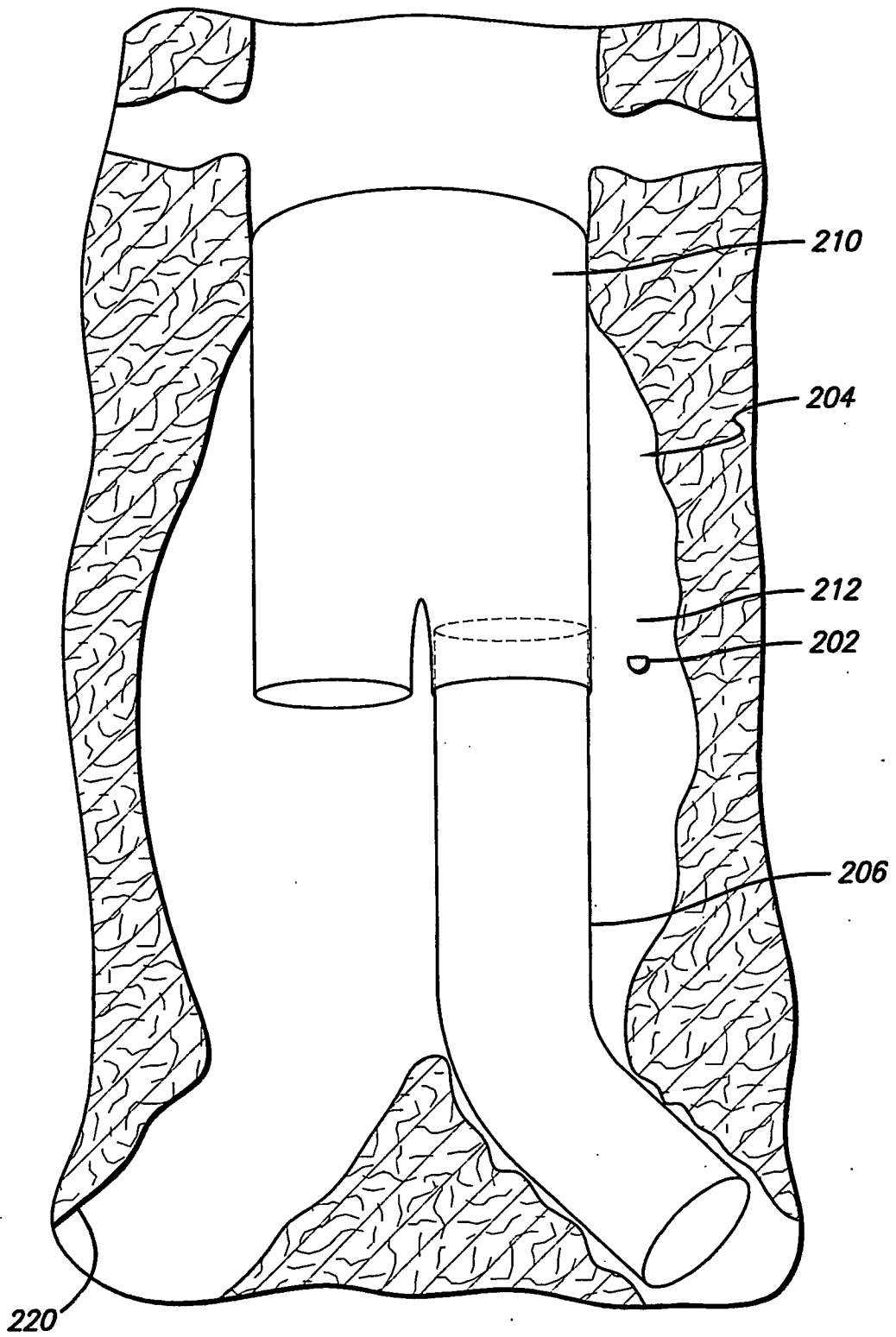


FIG. 8

REFERENCES CITED IN THE DESCRIPTION

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专利名称(译)	带可分离传感器的血管内移植物		
公开(公告)号	EP1626678A4	公开(公告)日	2006-12-06
申请号	EP2004753343	申请日	2004-05-26
[标]申请(专利权)人(译)	血管内TECH		
申请(专利权)人(译)	血管内TECHNOLOGIES, INC.		
当前申请(专利权)人(译)	血管内TECHNOLOGIES, INC.		
[标]发明人	BROWN PETER S KOVAC TIM WEI MICHAEL ECKERT ROBIN W		
发明人	BROWN, PETER, S. KOVAC, TIM WEI, MICHAEL ECKERT, ROBIN, W.		
IPC分类号	A61F2/06 A61F2/02 A61B5/00 A61B5/0215 A61B5/026 A61B5/05 A61F		
CPC分类号	A61B5/0031 A61B5/01 A61B5/02014 A61B5/0215 A61B5/026 A61B5/05 A61B5/145 A61B5/14532 A61F2/07 A61F2002/065 A61F2002/067 A61F2250/0002		
优先权	10/445723 2003-05-27 US		
其他公开文献	EP1626678B1 EP1626678A2		
外部链接	Espacenet		

摘要(译)

附有传感装置的血管内移植物，以便于测量植入该移植物的脉管系统内的相关参数。电源和发射器可以连接到移植植物上，以利于将测量结果传输到患者体外的接收设备。感测设备可以是具有测量和传输能力的电无源或集成设备。感测装置可以附接到移植材料上的特定位置或附接到内腔，从而从脉管系统内部的临界点提供相关参数，或者可以散布在移植材料的表面上或内腔内以提供轮廓。相关参数。感测装置可以使用连续缝线通过一根缝合线附接到移植植物材料，以使移植植物体积最小化，并且可以涂覆有抑制或控制组织生长的材料。可以采用生物可吸收的或硬线的系绳或缝合线将装置连接至一个或多个移植植物组件。

