



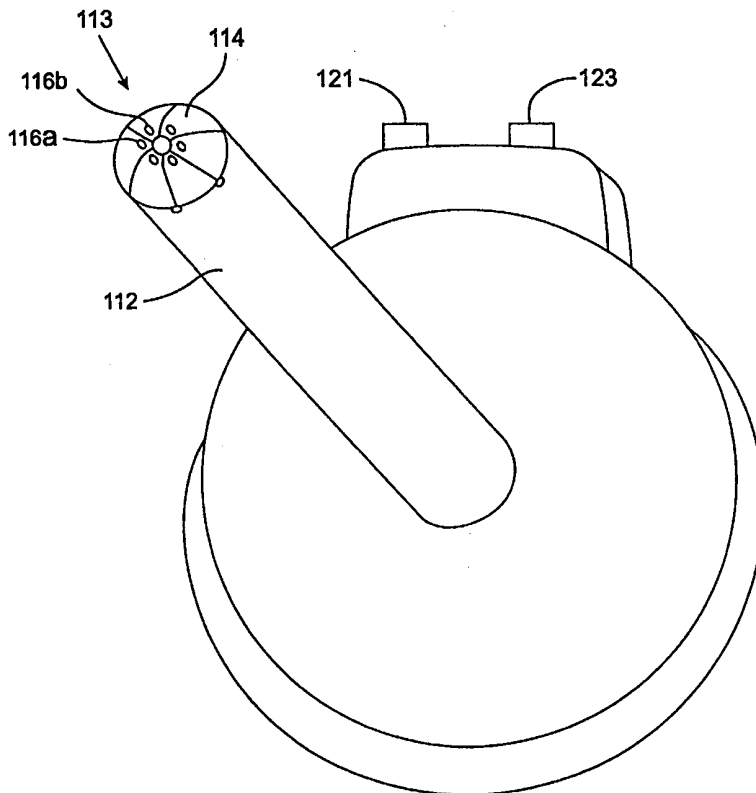
INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

<p>(51) International Patent Classification <sup>6</sup> : <b>A61B 5/0492, 17/39</b></p>	<p><b>A1</b></p>	<p>(11) International Publication Number: <b>WO 00/38574</b> (43) International Publication Date: 6 July 2000 (06.07.00)</p>
<p>(21) International Application Number: PCT/US99/12651 (22) International Filing Date: 4 June 1999 (04.06.99) (30) Priority Data: 60/113,651 23 December 1998 (23.12.98) US 60/120,663 19 February 1999 (19.02.99) US 60/123,268 8 March 1999 (08.03.99) US 09/325,998 4 June 1999 (04.06.99) US (71) Applicant: NUVASIVE, INC. [US/US]; 10065 Old Grove Road, San Diego, CA 92131 (US). (72) Inventors: MARINO, James, F.; 2620 St. Tropez Street, La Jolla, CA 92307 (US). STONE, Corbett, W.; 12212 Misty Blue Court, San Diego, CA 92131 (US). CHRISTOPHER, Troy, K.; 1002 Knight Drive, San Diego, CA 92126 (US). BLEWETT, Jeffrey, J.; 9950 Erma Road #104, San Diego, CA 92131 (US). KELLEHER, Brian, S.; 16999 Sky Valley Drive, Ramona, CA 92065 (US). (74) Agents: HECKADON, David, R. et al.; Townsend and Townsend and Crew LLP, 8th floor, Two Embarcadero Center, San Francisco, CA 94111 (US).</p>		<p>(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).  <b>Published</b> <i>With international search report.</i></p>

(54) Title: NERVE SURVEILLANCE CANNULAE SYSTEMS

(57) Abstract

This invention is an expandable tip cannula system (110), comprising a hollow cannula shaft (112) having a proximal end and a distal end; and an expandable tip (113) mounted at the distal end of the hollow cannula shaft, the expandable tip comprising a plurality of generally triangular shaped petals (114) held together in a radially inwardly tapered arrangement between adjacent petals, each petal comprising a nerve sensing electrode (116) disposed therein.



**FOR THE PURPOSES OF INFORMATION ONLY**

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav Republic of Macedonia	TM	Turkmenistan
BF	Burkina Faso	GR	Greece	ML	Mali	TR	Turkey
BG	Bulgaria	HU	Hungary	MN	Mongolia	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MR	Mauritania	UA	Ukraine
BR	Brazil	IL	Israel	MW	Malawi	UG	Uganda
BY	Belarus	IS	Iceland	MX	Mexico	US	United States of America
CA	Canada	IT	Italy	NE	Niger	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NL	Netherlands	VN	Viet Nam
CG	Congo	KE	Kenya	NO	Norway	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NZ	New Zealand	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's Republic of Korea	PL	Poland		
CM	Cameroon	KR	Republic of Korea	PT	Portugal		
CN	China	KZ	Kazakstan	RO	Romania		
CU	Cuba	LC	Saint Lucia	RU	Russian Federation		
CZ	Czech Republic	LI	Liechtenstein	SD	Sudan		
DE	Germany	LK	Sri Lanka	SE	Sweden		
DK	Denmark	LR	Liberia	SG	Singapore		
EE	Estonia						

## **NERVE SURVEILLANCE CANNULAE SYSTEMS**

### **CROSS-REFERENCES TO RELATED APPLICATIONS**

5 The present application is a regular application  
claiming benefit under 35 USC §119(e) from U.S. Provisional  
Patent Applications Serial Nos. 60/120,663 filed February  
19, 1999; and 60/113,651, filed December 23, 1998; the  
complete disclosures of which are hereby incorporated herein  
by reference in their entirety for all purposes.

10

### **TECHNICAL FIELD**

The present invention relates to nerve  
surveillance systems and to cannulae systems for use in  
minimally invasive spinal surgery.

15

### **BACKGROUND OF THE INVENTION**

A significant danger of performing intervertebral  
operations or accessing an intervertebral space during spine  
surgery is that of inadvertently contacting or damaging the  
20 para-spinal nerves, including the exiting nerve roots,  
traversing nerves and the nerves of the cauda equina. The  
exact location of these para-spinal nerves can not be  
determined prior to the commencement of surgery. Moreover,  
intervertebral spaces in the spine have other sensitive  
25 nerves disposed at locations which are not entirely  
predictable prior to insertion of the surgical tool into the  
intervertebral area. Accordingly, the danger of pinching or  
damaging spinal nerves when accessing an intervertebral  
space has proven to be quite limiting to the methods and  
30 devices used during minimally invasive spinal surgery. In  
addition, as cannulae are received through the patient's  
back, such as when performing minimally invasive spinal  
surgery, minor blood vessels are ruptured, thereby blocking

the surgeon's vision inside the intervertebral region after the cannula has been inserted.

#### SUMMARY OF THE INVENTION

5           The present invention provides nerve surveillance probes which are adapted to assist the surgeon in identifying the presence and location of para-spinal nerves as the probe is advanced during minimally-invasive surgery, thus providing a device for guiding the path of other  
10 surgical instruments to be inserted into this intervertebral space. In a preferred aspect of the present invention, an expandable tip cannula system is provided which functions both as an access portal for spinal surgery and as a system for nerve surveillance such that the presence and relative  
15 position of para-spinal nerves can be detected as the expandable tip cannula is inserted through the patient's facia and para-spinal musculature. An advantage of determining the position of a para-spinal nerve with respect to the distal tip of the cannula in particular is that the  
20 para-spinal nerve can be avoided or gently moved out of the surgeon's way while inserting the cannula. Accordingly, in a preferred aspect, the present invention provides a cannulated system which is adapted to assist the surgeon in guiding the path of surgical instruments received into the  
25 intervertebral space, while identifying the presence and location of para-spinal nerves as the cannula is advanced to a patient's intervertebral space during minimally invasive surgery.

          Optionally, the present nerve surveillance  
30 expandable tip cannula may also be adapted to selectively electrically induce cauterization of severed blood vessels when the cannula or other surgical instruments sever small blood vessels when they are inserted percutaneously into the

patient and are advanced along a path into the patient's intervertebral space. An additional advantage of the present cannula system therefore is that, prior to piercing the annulus of an intervertebral disc, vessels on the surface of the disc may be cauterized to assure clear vision inside the disc after surgical entry is made.

In one embodiment, the present expandable tip nerve surveillance cannula preferably comprises a hollow tubular body with a expandable tip portion mounted at its distal end. In a preferred aspect of the invention, the expandable tip portion comprises a plurality of generally triangular shaped petals which are held together in a radially-inwardly tapering arrangement by breakable seals disposed between adjacent petals. Since the expandable tip portion of the cannula tapers to a narrow blunt end, the cannula can be easily pushed through the patient's fascia and spinal musculature using blunt dissection, while minimizing the amount of cutting and tearing of such structures.

Alternatively, a central electrode can be disposed on a central obturator passing through the cannula and a second electrode can be disposed on a distal end of a second cannula, wherein the second cannula is used to open the petals.

An obturator shaft which is slidably received within the hollow tubular cannula body provides support for the cannula, giving the cannula sufficient strength such that the cannula can be inserted percutaneously through the patient's fascia and para-spinal musculature. Preferably, the obturator has a large solid handle which allows the surgeon to grasp and push the cannula through the resistance of the fascia and para-spinal musculature.

After the cannula has been inserted and is resting on the patient's annulus, a inner cannula or rod which is



Fig. 2 is a sectional side elevation view of the first nerve surveillance probe positioned adjacent the spinal nerve with the first probe received within a first cannula which is itself received with an expandable mesh.

5 Fig. 3 shows the probe of Fig. 2, but with the mesh expanded and a second cannula received thereover, (after the distal ends of the first cannula and expandable mesh have been advanced past the nerve).

10 Fig. 4 is a sectional side elevation corresponding to Fig. 3, but with the first probe and first cannula removed.

Fig. 5 is an end view corresponding to Fig. 4.

Fig. 6 is a side perspective view of a second nerve surveillance probe of the present invention.

15 Fig. 7 is a sectional side elevation view of a second nerve surveillance probe received within the second cannula.

Fig. 8 is an end view corresponding to Fig. 7.

20 Fig. 9A, 9B and 9C sequentially show a schematic view of an expandable mesh system as moved from a contracted position (Fig. 9A) to an expanded position (Fig. 9B), and with an outer cannula received thereover (Fig. 9C).

25 Fig. 10 is an end view of the nerve surveillance probe of Fig. 6 pushing a nerve out of the way of an advancing cannula.

Fig. 11 is an illustration of an expandable tip nerve surveillance probe of the present invention.

Fig. 12 is a perspective distal view of the system of Fig. 11.

30 Fig. 13 is a view of the distal tip of the system of Fig. 12, with the petals in a closed position.

Fig. 14 is a view corresponding to Fig. 13, but with the petals in an open position.

Fig. 15 is a sectional view of the system of Fig. 11, with an obturator received therein and the petals in a closed position.

5 Fig. 16 is a schematic illustration of the electrodes at the distal tip of the present invention, the electrodes being used to sense the position of a para-spinal nerve.

10 Fig. 17 is a sectional view of the system of Fig. 11 with a inner cannula received therein and the petals in an open position.

Fig. 18 is a side view of an alternate embodiment of the distal tip region of the present invention having truncated petals.

15 Fig. 19 is an end view corresponding to Fig. 18.

Fig. 20 is a top plan view of a peel back expandable tip cannula.

Fig. 21 is a side elevation view of the peel back cannula Fig. 20.

20 Fig. 22 is a side sectional view of the peel back cannula of Fig. 20 in a sealed position.

Fig. 23 is a sectional side elevation view of the peel back cannula of Fig. 20 in an open position.

Fig. 24 is a top plan view corresponding to Fig. 23.

25 Fig. 25 is a side elevation view of a curved petal nerve surveillance probe.

Fig. 26 is a side elevation view corresponding to Fig. 25, but with the petals in an open position.

30 Fig 27 is a view corresponding to Fig. 26, but with an expandable elastomer shown wrapped around the distal end of the curved petals.

Fig. 28 is a sectional elevation view of the distal end of an alternate nerve surveillance cannula.



Fig. 29 is a perspective view an alternative nerve surveillance probe.

Fig. 30 shows the surveillance probe of Fig. 29 with the petals opened by an inner cannula.

5 Fig. 31 corresponds to Fig. 30, but with the internal obturator removed.

Fig. 32 corresponds to Fig. 30, but with the internal obturator advanced distally.

10 Fig. 33 corresponds to Fig. 31, but with the internal cannula advanced distally.

#### DESCRIPTION OF THE SPECIFIC EMBODIMENTS

As will be set forth herein, the present invention encompasses both nerve surveillance probes which are received through cannulae, and various expandable tip cannulae comprising nerve surveillance probes at their distal ends.

In a first preferred embodiment, as is seen in Fig. 1, an electromyography nerve surveillance probe 10 having a blunt end 11 is provided. Electrode 13 is disposed at the distal end of probe 10 and is charged by electrical contacts 15. As electrode 13 approaches nerve 20 (as seen in Fig. 2), the minimal threshold depolarization value elicited by the electrode will result in corresponding electromyography activity, such that the presence of nerve 20 can be sensed by standard electromyographic techniques, thus indicating the presence of the nerve. Specifically, using standard electromyographic techniques, the presence of nerve 20 will be sensed by appropriate needles or patches attached to the appropriate muscle as electrode 13 stimulates, and thereby depolarizes electrode 13.

In an exemplary method of application, (as is shown in Fig. 2), the present nerve surveillance probe 10

can be advanced percutaneously through the patient's back in a posterolateral approach towards the patient's intervertebral space using the arrangement in which a first cannula 30 surrounds probe 10 as the probe is advanced. As probe 10 is advanced, it will then become positioned proximal nerve 20. When this occurs, the presence of nerve 20 relative to probe 10 will be determined by the signal generated by electrode 13 as set forth above.

In one preferred aspect of the present invention, an expandable mesh 32 is received over first cannula 30 such that expansion of this mesh from the contracted position shown in Fig. 2 to the expanded position shown in Fig. 3 will gently move nerve 20 out of the way.

Also in a preferred aspect as shown in Fig. 3, a second cannula 34 can thereafter be received over expanded mesh 32, thereby providing a large passageway 40 for intervertebral access when probe 10, first cannula 30, and expanded mesh 32 are removed as shown in Figs. 4 and 5. Accordingly, the large passageway 40 into the intervertebral area provided by cannula 34 protects sensitive nerve 20 while providing access for surgical instruments therethrough, including such surgical instruments as intervertebral inserts, bone decorticators, cameras, articulating forceps, intervertebral inserts and intervertebral positioning systems.

As is seen in Fig. 6, a second nerve surveillance probe 9 is also provided. Nerve surveillance probe 9 has a plurality of electrodes 12, 14, 16 and 18 disposed at radial locations adjacent to blunt distal end 8, as is seen in Figs. 6, 7 and 8. Radially-disposed electrodes 12, 14, 16, and 18 perform a variety of useful functions, as follows.

Referring to Fig. 8, as electrodes 12, 14, 16, and 18 are disposed at radial locations around the tip of probe

10, the electrodes which are closest to nerve 20, (in this case electrode 14, and to a lesser degree electrodes 12 and 16), will operate to depolarize the nerve such that the presence of nerve 20 can be detected by standard  
5 electromyographic techniques. As such, a signal will be generated telling the operating surgeon that nerve 20 is proximal to electrode 14. As can be appreciated, should nerve 20 instead be positioned in another orientation, the signal from electrodes 12, 14, 16 and 18 would instead  
10 indicate the presence of the nerve at a different location. Accordingly, probe 9 can be operated as a tool for inspecting the interior passageway of cannula 34 to determine if nerve 20 had become inadvertently trapped therein as cannulae 34 is advanced over expanded mesh 32.  
15 Moreover, as the electrodes 12, 14, 16, and 18 are disposed at radial locations around the distal end of the probe, it is possible to determine the exact location of nerve 20. Preferably as well, each of electrodes 12, 14, 16, and 18 will be activated in a repeating sequence with a sufficient  
20 delay time therebetween to detect an electromyographic response.

In another aspect of the invention, radially disposed electrodes 12, 14, 16, and 18 can be used for electrocoagulation of blood vessels, for example, blood  
25 vessels on the patient's annulus when accessing the patient's intervertebral region. Specifically, as a plurality of electrodes are disposed at the distal end of probe 9, it is possible to pass current between various electrodes, thus cauterizing adjacent blood vessels.

30 In another aspect of the invention, radially disposed electrodes 12, 14, 16, and 18 can be used to assist in avoiding, (or alternatively in moving), nerve 20 as follows. Referring to Fig. 10, nerve 20 will be determined

to be adjacent to electrode 14 using the above set forth method. Probe 10 can then be gently moved in a radial direction away from electrode 14, as is shown by arrow D, such that nerve 20 can then be gently pushed out of the way, providing safe access to the patient's intervertebral space. Alternatively, the movement of probe 10 in a direction opposite direction D will push the nerve out of the way such that a cannula can then be advanced past nerve 20 without damaging the nerve.

In another aspect of the present invention as shown in Figs. 9A, 9B and 9C, the expansion of mesh 32 is controlled as follows. As is shown in Fig. 9A, expandable mesh 32 is in a contracted position and is mounted on the end of a cannula 35. A distal end of mesh 32 is positioned against the patient's annulus 40 or any other suitably hard bone structure. Pushing rod or cannula 35 in direction D2 will compress mesh 34, causing it to expand radially and shorten. This movement will displace nerve 20 (shown here in cross section). Following this, cannula 37 can be slid over expanded mesh 32 is seen in Fig. 9B. Following this, cannula 37 can be advanced past nerve 20, gently pushing nerve 20 still further out of the way, as shown in Fig. 9C. Lastly, rod or cannula 35 and attached mesh 32 can be removed, leaving a large cannulated passageway to the annulus or intervertebral space.

It is to be understood that the present nerve surveillance probes can be used without the expandable mesh system of Figs. 9A, 9B and 9C. Moreover, it is to be understood that the present method and apparatus of minimally invasive nerve surveillance can be used in any arthroscopic procedure.

As can also be appreciated the present nerve surveillance probes are able to detect the presence of any

other efferent skeletal motor nerve in addition to the spinal nerve and can thus be used in various surgical procedures. Alternatively, using evoked potential electromyography, the present nerve surveillance probes are  
5 also adapted to sense the presence of afferent sensory nerves in response to signals received in the spinal cord or cerebral cortex.

In a second preferred embodiment, the present invention provides an expandable tip nerve surveillance  
10 cannula system 110 comprising an endoscopic hollow cannula shaft 112 having an expandable tip 113 comprised of a plurality of petals 114, (the details of petals 114 are better shown in Figs. 12, 13, and 14). System 110 further comprises an obturator 120 which is slidably received within  
15 cannula shaft 112. As is shown in Fig. 15, obturator 120 is a rigid structure which provides internal support to cannula shaft 112 such that cannula shaft 112 can be received percutaneously. Shaft 112 can have a cross section which is circular, oval, racetrack-shaped or any other design. By  
20 holding obturator handle 122, the surgeon is able to advance cannula shaft 112 through the patient's para-spinal musculature and dock expandable tip 113 at the patient's annulus.

As seen in Figs. 12 and 13, expandable tip 113 is  
25 comprised of a plurality of petals 114, held together by breakable seals 115. Breakable seals 115 can be formed by an elastomeric material with predictable failure segments between the petals, which fracture with radial expansion of the petals. In one preferred aspect each of petals 114 has  
30 an electrode 116 disposed therein as shown. Electrodes 116 serve the following important functions.

First, electrodes 116 can be used for electromyography, and in particular to sense the presence

and relative position of para-spinal nerves as cannula shaft 112 is advanced. Referring to Fig. 16, as can be seen electrodes 116a, 116b, 116c, 116d, 116e and 116f are disposed radially about cannula shaft 112, with one  
5 electrode disposed in each of petals 114, as has been described. Electrodes 116a, 116b, 116c, 116d, 116e and 116f assist in sensing the presence and location of para-spinal nerve 160 as follows. The electrodes closest to nerve 160, (in this case electrodes 116b and 116d, and to a lesser  
10 degree, electrodes 116a and 116d), will operate to depolarize nerve 160 such that the presence of nerve 160 can be detected by electromyography. As such, shaft 112 can be moved in direction D, thereby avoiding nerve 160 as shaft 112 is inserted. Alternatively, of course, shaft 112 can be  
15 moved in the opposite direction to D, such that cannula shaft 112 gently moves nerve 112 out of the way. Moreover, when none of electrodes 116a, 116b, 116c, 116d, 116e and 116f sufficiently stimulate to depolarize the nerve, (and thereby assist in its detection), shaft 112 can be safely  
20 advanced toward the patient's intervertebral space. Should each one of electrodes 116a, 116b, 116c, 116d, 116e and 116f depolarize the nerve, this would indicate that the nerve is directly in front of the advancing cannula shaft 112. Accordingly, the cannula shaft could be moved such that  
25 contact with the nerve is avoided.

Alternatively, when none of electrodes 116a, 116b, 116c, 116d, 116e and 116f indicate the presence of a nerve, electrodes 116a, 116b, 116c, 116d, 116e and 116f can be powered to a higher level such that cauterization of minor  
30 blood vessels can be achieved by passing increased electric current between each of the various adjacent electrodes, thus cauterizing adjacent blood vessels. Preferably, the present invention comprises a safety system such that

cauterization power levels for electrodes 116 are not activated when any of electrodes 116 sense the presence of a para-spinal nerve thereby.

Preferably, each of electrodes 116a, 116b, 116c, 5 116d, 116e and 116f are operated in sequence, affording a sufficient latency period therebetween for the detection of an electromyographic signal.

As seen in Fig. 11, button 121 can be used to activate the nerve sensing functions and button 123 can be 10 used to activate the blood vessel cauterization functions. Buttons 121 and 123 are conveniently located on the near handle 122 such that they may be activated while the surgeon grips obturator handle 122.

Subsequent to being positioned at the patient's 15 annulus, obturator 120 is removed from cannula shaft 112. As seen in Fig. 17, inner cannula 130 is then inserted into cannula shaft 112. Inner cannula 130 is dimensioned to be of a size that, when fully inserted into shaft 112, inner cannula 130 breaks apart seals 115, forcing petals 114 to be 20 displaced radially outwards to a distance of at least the internal diameter of shaft 112 as shown. Inner cannula 130 can alternately comprise a solid rod or obturator which is dimensioned to be received within shaft 112 to open petals 114.

As can be seen in Fig 13, a notch 118 is found 25 between adjacent petals 114 where petals 114 are mounted to the distal end 113 of cannula shaft 112. Notches 118 operate to facilitate breakage of seals 115 by providing a stress relief region at the base of breakable seals 115.

In an alternate design, as shown in Figs. 18 and 30 19, distal tip 113 comprises truncated petals 114a which, when sealed together by way of breakable seals 115, meet at their distal end to define a small opening 117 at distal tip

113 of cannula shaft 112. In this design, an obturator 120a is slidably received within cannula shaft 112. Obturator 120a has a narrow distal end 113a which protrudes through opening 117. Electrodes 119a, 119b, 119c, 119d, 119e and  
5 119f are disposed radially about the narrow distal end 113a of obturator 120a, functioning similar to the probe design shown in Fig. 6.

In this alternate design of Figs. 18 and 19, nerve surveillance and blood vessel cauterization functions as  
10 described above and as performed by electrodes 116 on petals 114 are instead performed by electrodes 119 on obturator 120a. In this aspect of the invention, petals 114a are truncated and obturator 120a protrudes therethrough.

In another alternate embodiment, a peel back  
15 cannula having an expandable tip is provided. Referring to Fig. 20, cannula 150 is provided. Cannula 150 has a tapered narrow distal end 152 and a tear away line 153 which is formed in the preferred polymeric material of cannula 150. Tear away line 153 will split under tension as will be  
20 explained. Cannula 150 may also comprise electrodes 153 which perform a similar function to the electrodes 116 described herein. Electrodes 153 can be disposed axially along the length of cannula 150, or radially around the distal end of cannula 150, or some combination thereof.

25 An advantage of being disposed axially along the cannula is that electrodes 153 will be able to sense the position of a nerve relative to the cannula in an axial dimension. Similarly, an advantage of being disposed radially around the cannula is that the electrodes will be  
30 able to sense the position of a nerve relative to the cannula in a radial dimension. It is to be understood that all embodiments of the present invention comprise the concept of nerve surveillance electrodes disposed both



radially around and axially along the nerve surveillance cannula or obturator, and that the radial electrode placement shown in the design of Figs. 7, 8 and 11 to 19, and the axial electrode placement shown in the design of  
5 Figs. 20 to 23 is not limiting.

In a preferred method of operation, cannula 150 is advanced such that its tapered end 152 is adjacent nerve 160 as is seen in Fig. 22. An obturator 155 is positioned within cannula 150. Obturator 155 provides structural support for  
10 the cannula as it is being inserted or as it is moving a nerve. Obturator 155 is thereafter removable such that cannula 150 operates as an open passageway as will be explained.

A narrow inner cannula 157 may also be provided.  
15 Cannula 157 is received around obturator 155 and within cannula 150. When the operator has determined it is safe and desirable to open cannula 150, inner cannula 157 is advanced to the position shown in Figs. 23 and 24. Specifically, inner cannula 157 pushes against the tapered  
20 end of 152 of cannula 150 causing cannula 150 to split open along tear away line 153. Accordingly, inner cannula 157 can be used to provide a cannulated passageway when obturator 157 has been withdrawn therefrom. Alternatively, inner cannula 157 can be replaced by a suitably dimensioned  
25 obturator for opening cannula 150 along tear away line 153.

Tear away line 153 may be formed by scribing the polymeric material forming cannula 150. Tear away line 153 preferably runs some distance along opposite sides of the open 152 of cannula 150. Alternatively, tear away line 153  
30 can be disposed along the top and bottom of cannula 150 as shown.

Figure 25 is a side view of a curved petal design of the present invention in a closed position with cannula

220 having outwardly curved petals 212 at distal end 215. A nerve 230 is disposed adjacent the ends of closed petals 212 as shown. Petals 212 are then opened, (using methods described herein), as shown in Fig. 26. The opening of  
5 petals 212 causes nerve 230 to be generally displaced upward away from an operative site which may preferably comprise a patient's intervertebral disk 240.

As shown in Fig. 27, an elastomer 250 can be wrapped around the petals 212 such that nerves are not  
10 pinched in gaps 213 between the adjacent petals either when the petals are first opened or when the petals are closed during the removal of the cannula from the patient. It is to be appreciated that elastomer 250 could also be wrapped around the ends of any of the straight petal designs shown  
15 in Figs. 11 to 19.

The operative site or target site may comprise a patient's intervertebral disk 240 when the present invention is used in minimally invasive spinal surgery. It is to be understood, however, that the present expandable tip cannula  
20 can be used in all manner of minimally invasive surgery and is especially useful for approaching any target site having sensitive nerves adjacent thereto since the present invention is specifically adapted to gently push the nerve out of the way as the petals are opened, thereby providing a  
25 cannulated access portal for the insertion and removal of various surgical devices through cannula 220.

Fig. 28 shows an alternate design of the distal end 302 of a nerve surveillance cannula 300. Cannula 300 has a plurality of expanding petals 314, with each petal 314  
30 comprising an electrode 316 adapted for nerve surveillance or blood vessel cauterization as described above. In this aspect of the invention, an obturator 310 protrudes through an opening between petals 314, as shown. As can be seen,

obturator 310 may preferably be tapered to a narrow distal end 302, which assists in easing cannula 300 through the patient's fascia and para-spinal musculature and into the patient's intervertebral space. In addition, distal end 302  
5 of obturator 310 can be shaped to latch against the ends of petals 314, as shown, thereby assisting in holding together petals 314 as cannula 300 is advanced.

Preferably, obturator 310 further comprises a centrally disposed electrode 320. Electrode 320, being  
10 axially displaced from electrodes 316 is adapted to sense the position of a nerve in the axial direction as probe 300 approaches the nerve. Subsequent to placement at the patient's intervertebral space, an internal cannula 315 can be advanced distally to open petals 314 with obturator 310  
15 being advanced slightly to first un-latch the distal ends of petals 314 and then withdrawn from cannula 300, providing a cannulated access to the patient's intervertebral space.

Fig. 29 through 33 show an alternative nerve surveillance cannula and probe system 400, comprising a  
20 cannula 402 having a plurality of radially outwardly extending petals 404. An internal obturator 500 is received within cannula 402. Obturator 500 has a electrode 502 disposed at its distal end as shown in Fig. 30. Electrode 502 can also be seen at distal end of cannula 402 in Fig.  
25 29. Electrode 502 operates to stimulate and thereby, depolarize a nerve as cannula 402 is advanced towards the patient's intervertebral space. Fig. 29 shows cannula 402 with petals 404 closed around electrode 502 as the cannula is advanced.

30 Fig. 30 shows an inner cannula 550 which is advanced through cannula 402 to open petals 404 as shown. Inner cannula 550 preferably comprises an electrode 510 which is disposed around the distal end of the cannula, as

shown. After inner cannula 550 has opened petals 404, as shown, electrode 502 is turned off and obturator 500 is removed from inner cannula 550 as is shown in Fig. 31. Electrode 510 remains turned on such that it is adapted to  
5 detect whether a nerve is positioned close to entering within cannula 550, or whether a surgical instrument advanced through cannula 550 would contact a nerve proximal electrode 510.

As is shown in Fig. 32, obturator 500 can  
10 thereafter be advanced through cannula 550 to bluntly divide and dilate the annulus of a disc. In this aspect of the invention, electrode 502 is turned off as the annulus is divided and dilated. Annular electrode 510 may preferably be turned on during this procedure to sense the presence of  
15 nerves adjacent the distal end of cannula 550.

As is seen in Fig. 33, after the annulus has been divided and dilated, obturator 500 can be withdrawn from cannula 550 with cannula 550 advanced distally into the hole cut into the annulus. As such, a safe cannulated access way  
20 into the annulus or other region of the patient's body is provided.

WHAT IS CLAIMED IS:

- 1           1. A nerve surveillance system, comprising:  
2                 a cannula having an expandable distal end with a  
3 plurality of nerve surveillance electrodes disposed thereon.
  
- 1           2. The nerve surveillance system of claim 1, wherein  
2 the expandable distal end comprises:  
3                 a plurality of inwardly tapering petals.
  
- 1           3. The nerve surveillance system of claim 2, wherein,  
2                 at least one of the plurality of nerve surveillance  
3 electrodes are disposed in each of the petals.
  
- 1           4. The nerve surveillance system of claim 1, wherein,  
2                 the nerve surveillance electrodes are adapted for  
3 electromyography.
  
- 1           5. The nerve surveillance system of claim 1, wherein,  
2                 the nerve surveillance electrodes are adapted to  
3 cauterize blood vessels.
  
- 1           6. The nerve surveillance system of claim 2, further  
2 comprising,  
3                 an slidable member received within the cannula, the  
4 slidable member dimensioned to cause the petals to deflect  
5 radially outwards when advanced therethrough.
  
- 1           7. The nerve surveillance system of claim 6, wherein,  
2 the slidable member comprises an inner cannula.
  
- 1           8. The nerve surveillance system of claim 6, wherein,  
2 the slidable member comprises an obturator.

1           9. The nerve surveillance system of claim 6, further  
2 comprising:  
3           an obturator dimensioned to be received within the  
4 inner cannula.

1           10. The nerve surveillance system of claim 2, wherein,  
2           the petals are curved radially outwards at their  
3 distal ends.

1           11. The nerve surveillance system of claim 2, further  
2 comprising,  
3           an expandable elastomer wrapped around the petals.

1           12. The nerve surveillance system of claim 2, wherein,  
2           the petals are held together by breakable seals.

1           13. The nerve surveillance system of claim 2, wherein,  
2           the petals are generally triangular in shape.

1           14. The nerve surveillance system of claim 1, wherein,  
2           the plurality of nerve surveillance electrodes are  
3 disposed radially around the distal end of the cannula.

1           15. The nerve surveillance system of claim 1, wherein,  
2           at least two of the plurality of nerve surveillance  
3 electrodes are disposed axially with respect to one another  
4 along the length of the cannula.

1           16. The nerve surveillance system of claim 9, wherein,  
2 the obturator protrudes from the distal end of the cannula.

1           17. The nerve surveillance system of claim 16, wherein,  
2 the obturator further comprises a central electrode at its  
3 distal end.

1           18. A method of accessing an intervertebral space while  
2 protecting adjacent spinal nerves, comprising:

3                 advancing a cannula through a patient's fascia and  
4 para-spinal musculature, the cannula comprising an  
5 expandable tip with a plurality of nerve sensing electrodes  
6 disposed thereon;

7                 sensing the presence of a nerve with the nerve  
8 sensing electrodes;

9                 inserting a slidable member though the cannula to  
10 open the expandable tip of the cannula; and

11                 removing the slidable member from the expandable  
12 tip cannula, thereby providing a cannulated access to the  
13 intervertebral space through the expandable tip cannula.

1           19. The method of claim 18, wherein,  
2 the expandable tip comprises a plurality of generally-  
3 triangular shaped petals, each petal having a nerve sensing  
4 electrode disposed thereon.

1           20. The method of claim 18, wherein sensing the  
2 presence of a nerve with the nerve sensing electrodes,  
3 comprises:

4                 sensing the axial position of the nerve relative to the  
5 expandable tip cannula.

1           21. The method of claim 18, wherein sensing the  
2 presence of a nerve with the nerve sensing electrodes,  
3 comprises:

4                 sensing the radial position of the nerve relative to  
5 the expandable tip cannula.

1           22. The method of claim 18, wherein, the cannula is  
2 inserted percutaneously.

1           23. The method of claim 19, wherein, the cannula is  
2 inserted posterolaterally.

1           24. The method of claim 18, further comprising,  
2           avoiding an adjacent spinal nerve by moving the  
3 cannula in a direction away from the nerve.

1           25. The method of claim 18, further comprising,  
2           activating the electrodes to cauterize blood  
3 vessels.

1           26. A system for accessing an intervertebral space  
2 comprising,  
3           a nerve surveillance probe having a plurality of  
4 electrodes disposed thereon;  
5           a first cannula dimensioned to be slidably  
6 received over the nerve surveillance probe;  
7           an expandable covering dimensioned to be slidably  
8 received over the; and  
9           a second cannula dimensioned to be slidably  
10 received over the expandable covering.

1           27. The system of claim 26, wherein the plurality of  
2 electrodes are disposed radially around the distal end of  
3 the probe.

1           28. The system of claim 26, wherein the plurality of  
2 electrodes are disposed axially along the distal end of the  
3 probe.

1           29. The system of claim 26, wherein the expandable  
2 covering comprises an expandable mesh.



1           30. A method of providing a passageway to a patient's  
2 intervertebral space, comprising:

3                 advancing a nerve surveillance probe in a cannula  
4 towards the patient's intervertebral space;

5                 sensing the presence of a nerve with at least one  
6 electrode disposed on the nerve surveillance probe.

1           31. The method of claim 30, wherein sensing the  
2 presence of a nerve with at least one electrode comprises:

3                 sensing the presence of the nerve in a radial  
4 direction relative to a central axis of the cannula with a  
5 plurality of electrodes radially disposed around the end of  
6 the probe.

1           32. The method of claim 30, further comprising:

2                 expanding an expandable covering disposed over the  
3 cannula;

4                 receiving a second cannula over the expandable  
5 covering after the expanded covering is in an expanded  
6 position; and

7                 removing the nerve surveillance probe and  
8 expandable covering from within the second cannula, thereby  
9 providing a passageway to the intervertebral region.

1           33. A nerve surveillance system, comprising:

2                 a cannula having an expandable distal end;

3                 an inner cannula received within the cannula  
4 having an expandable distal end;

5                 the inner cannula actuating the opening of the  
6 expandable distal end; and

7                   an obturator received within the inner cannula,  
8 the obturator having a centrally disposed electrode at a  
9 distal end thereof.

1           34. The nerve surveillance system of claim 33, wherein  
2           the inner cannula has an annular electrode at its  
3 distal end.

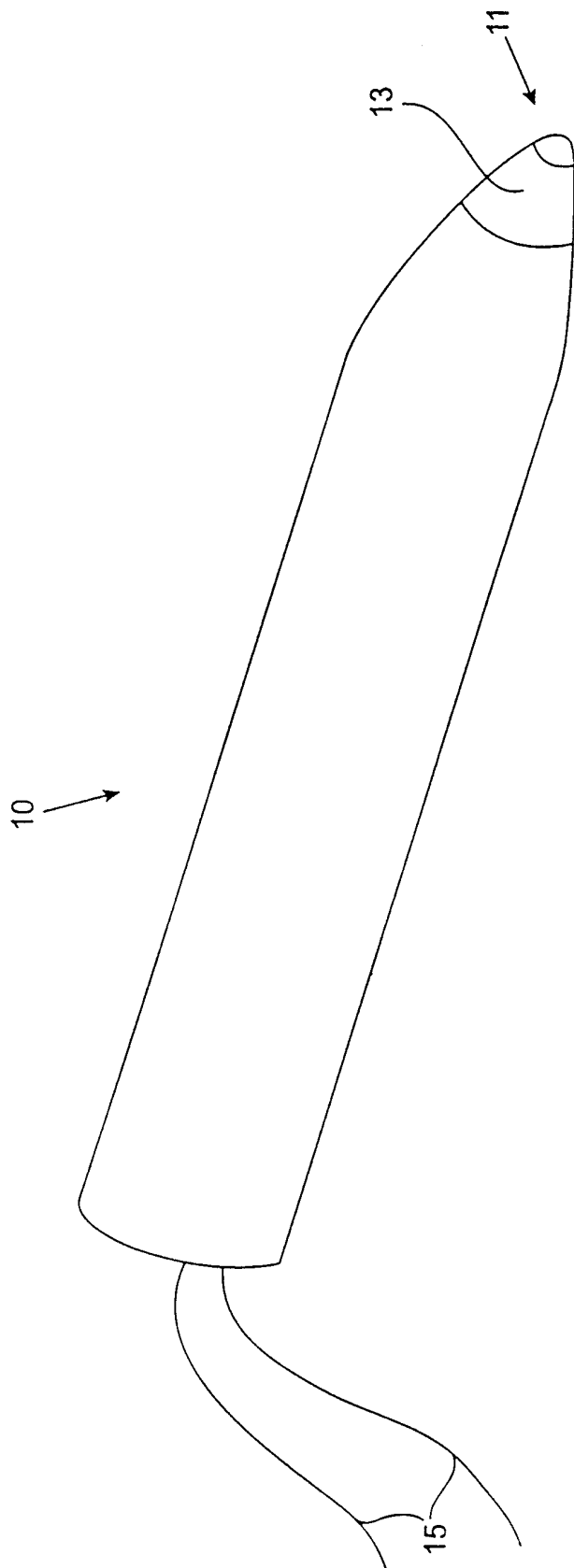


FIG. 1

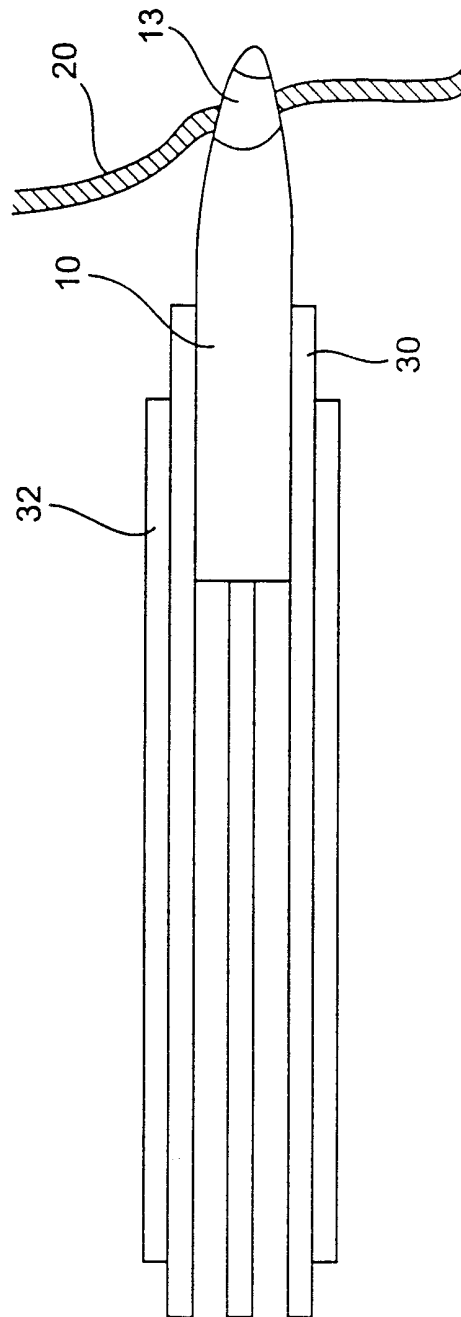


FIG. 2

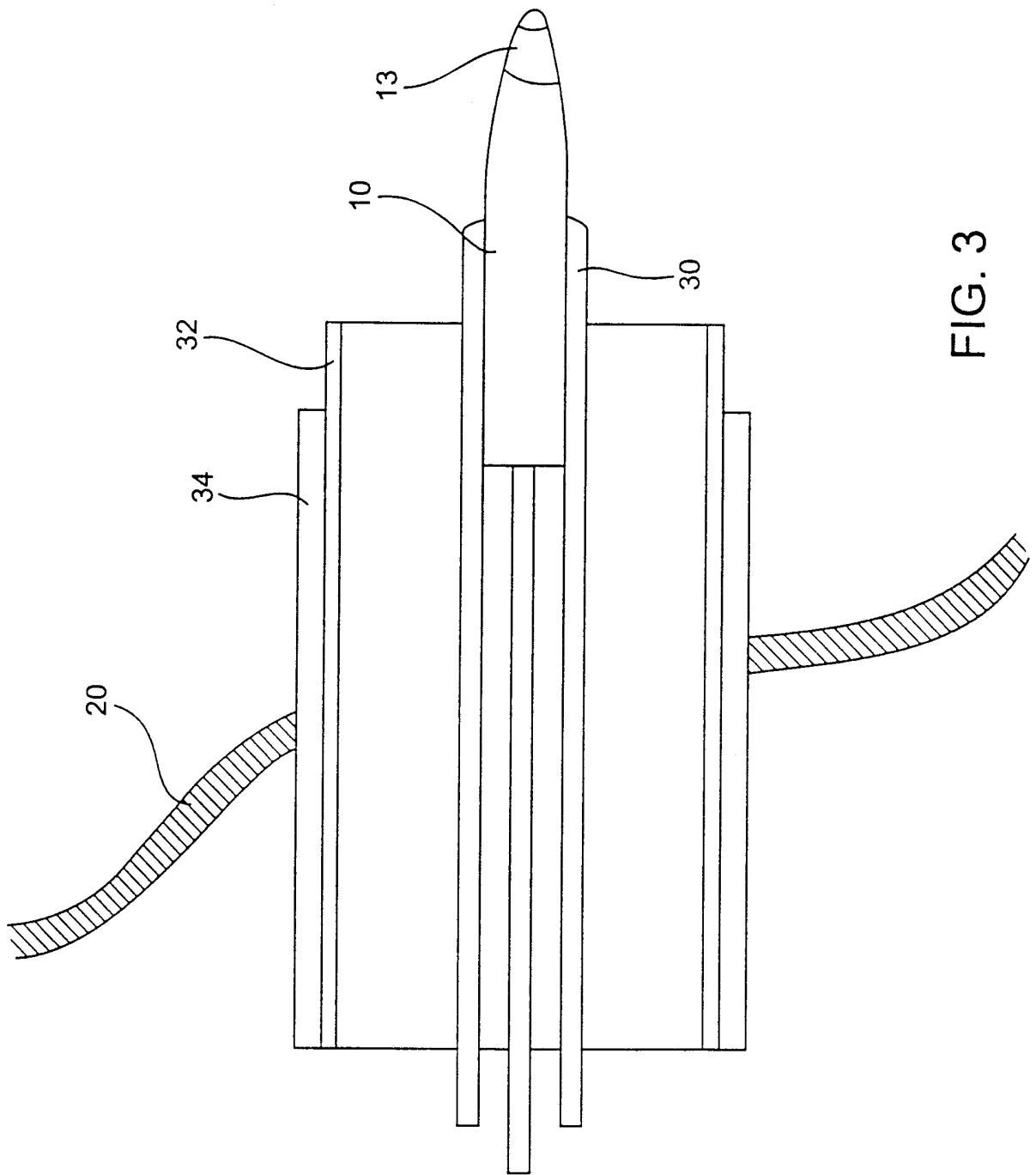
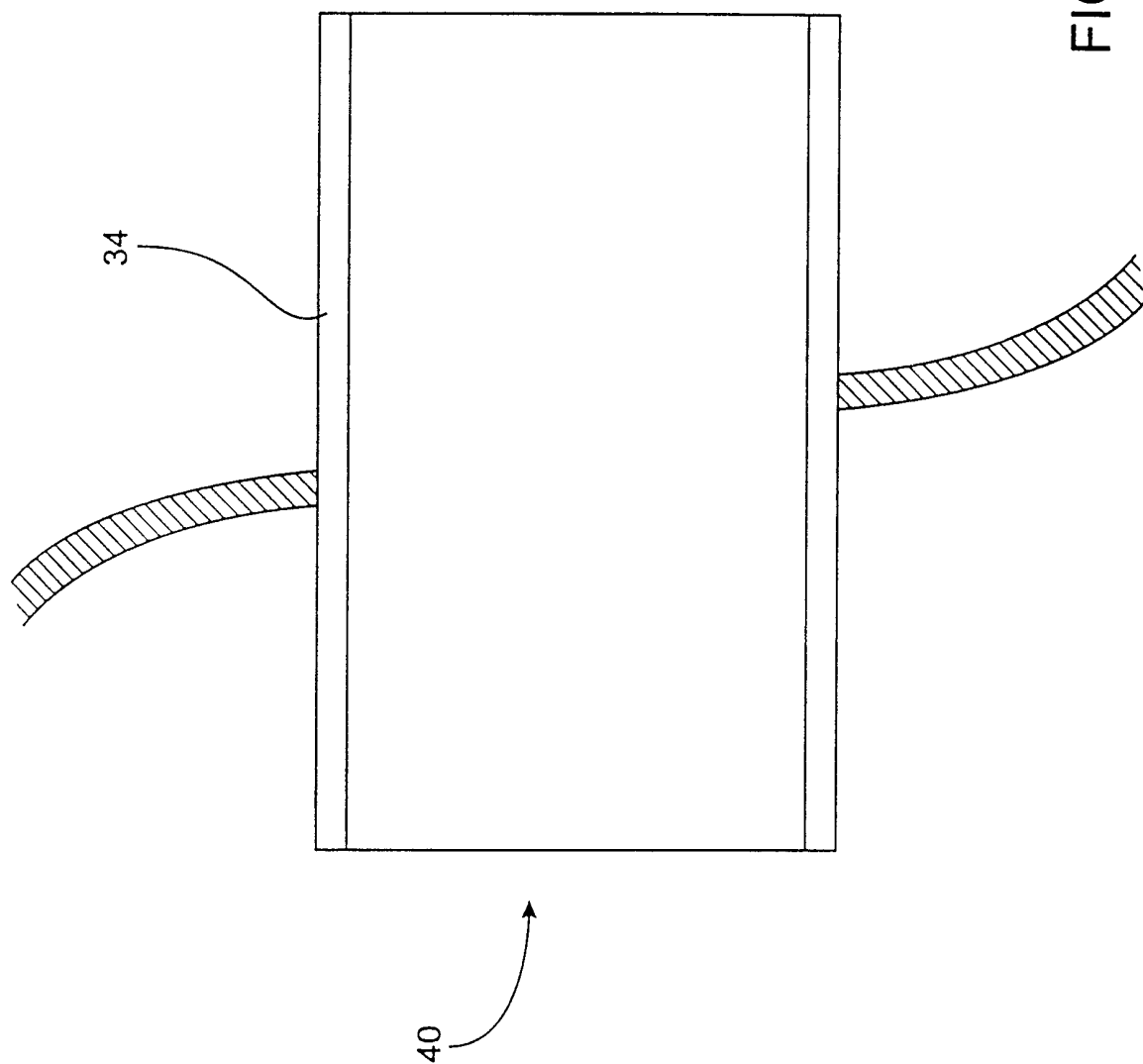


FIG. 3



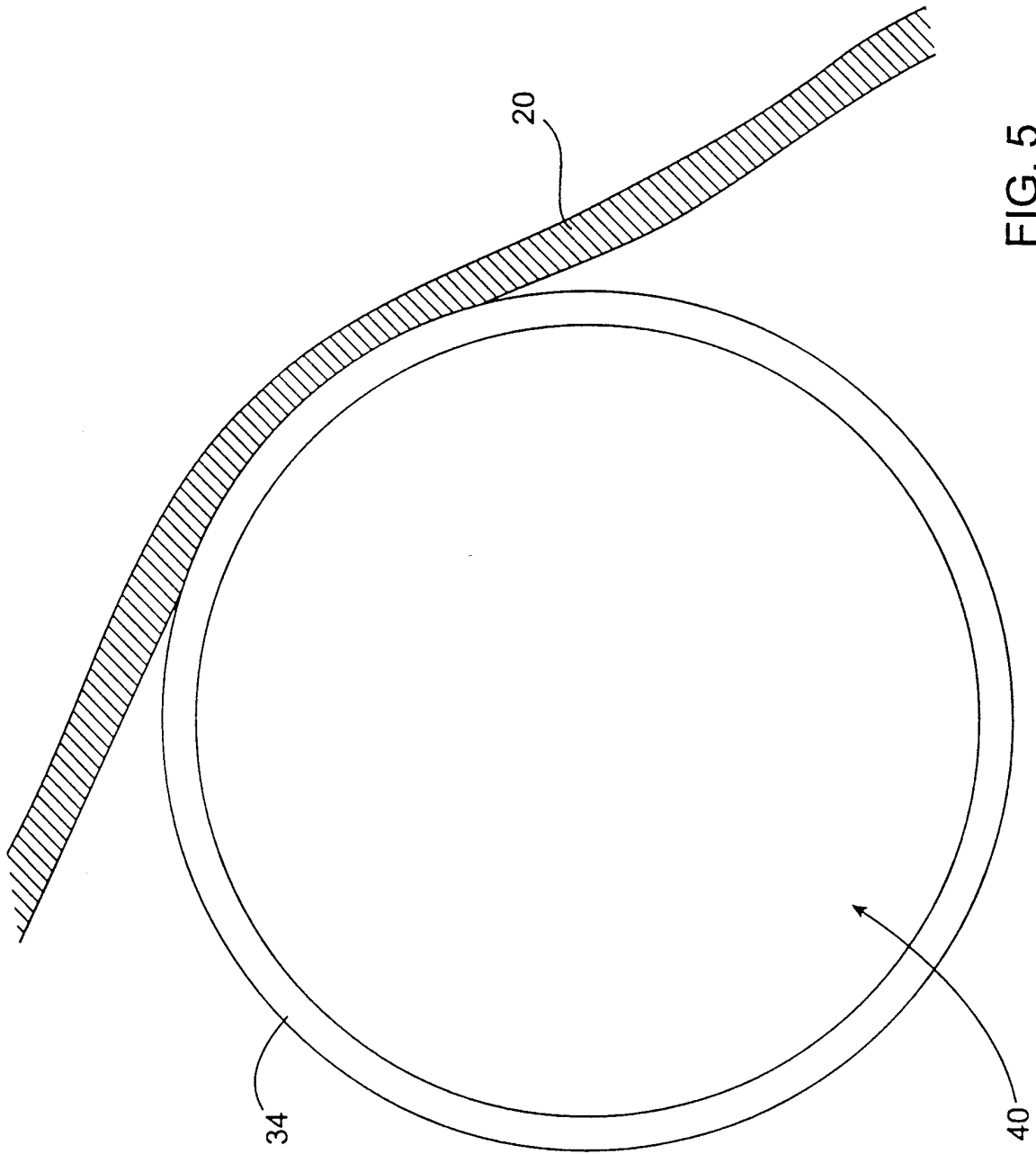


FIG. 5

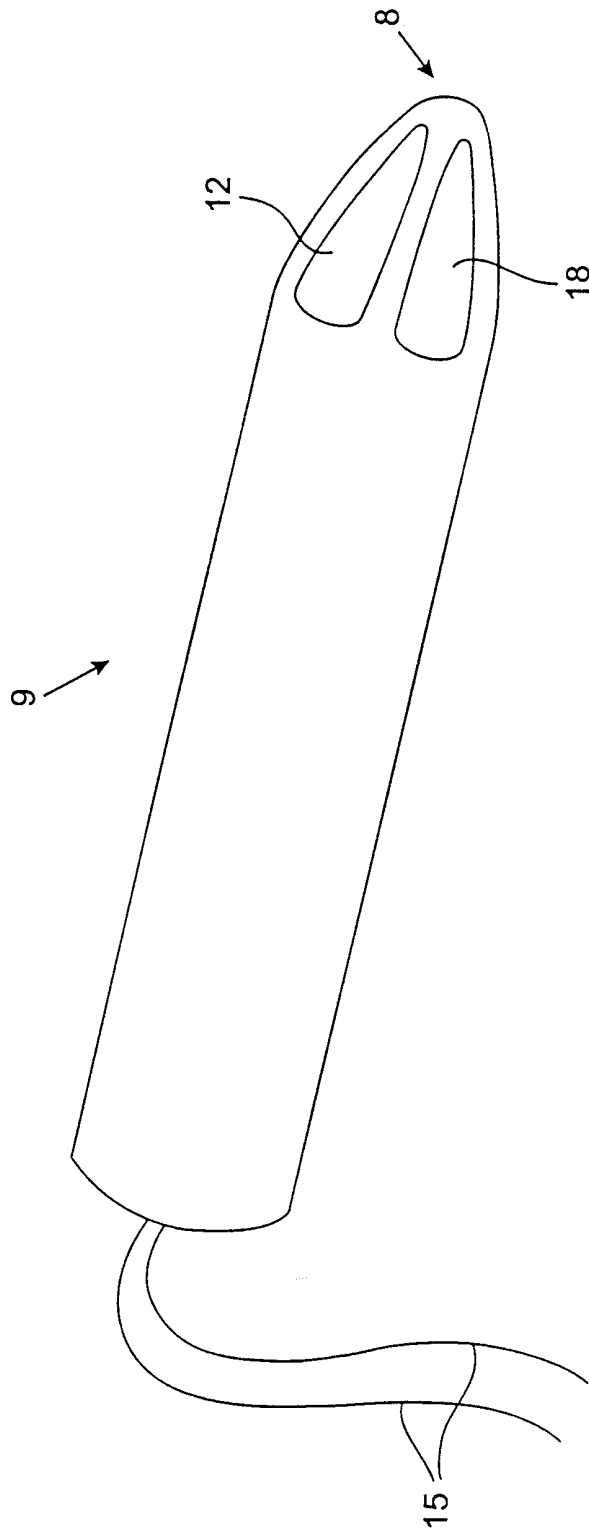


FIG. 6



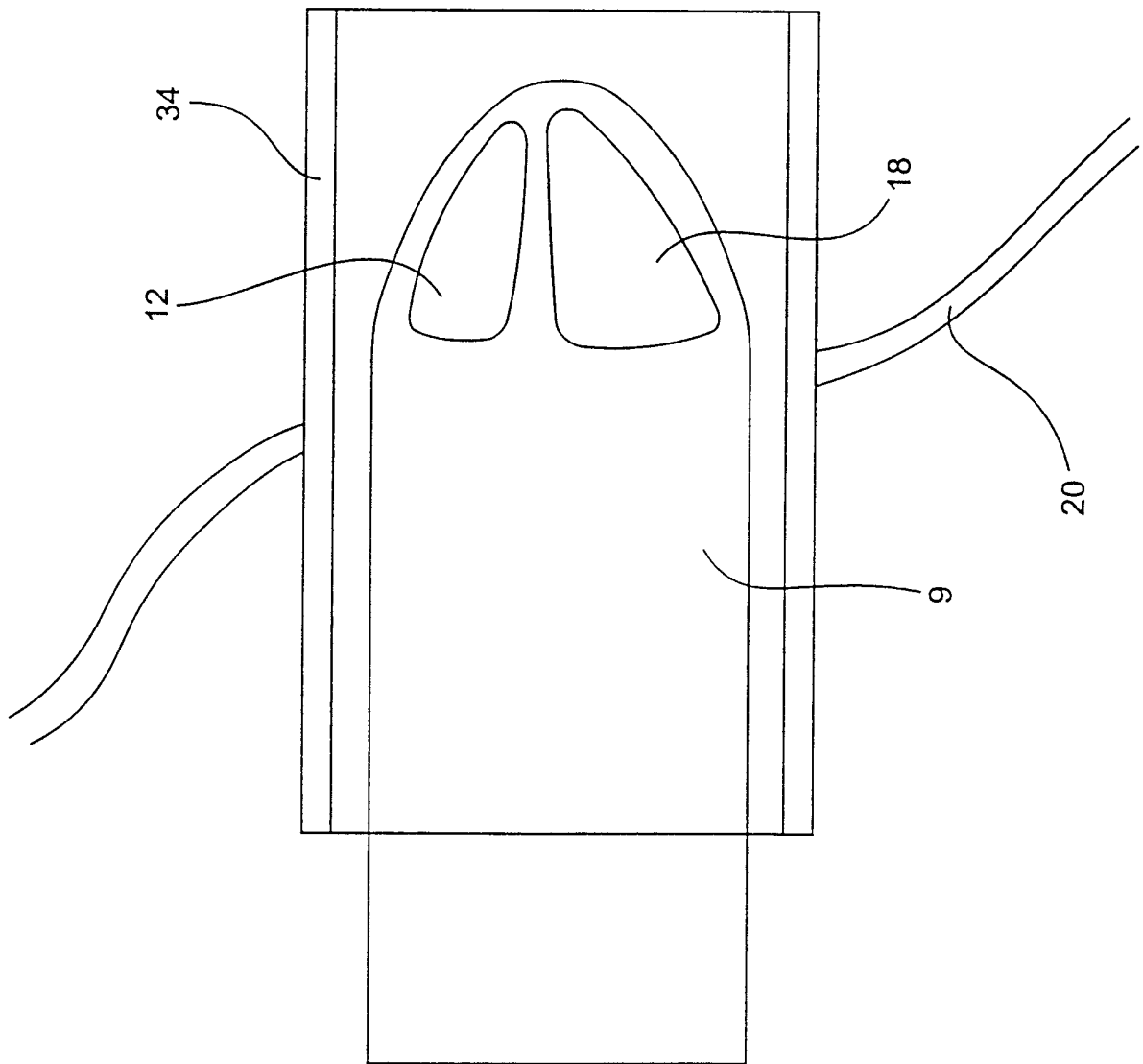


FIG. 7

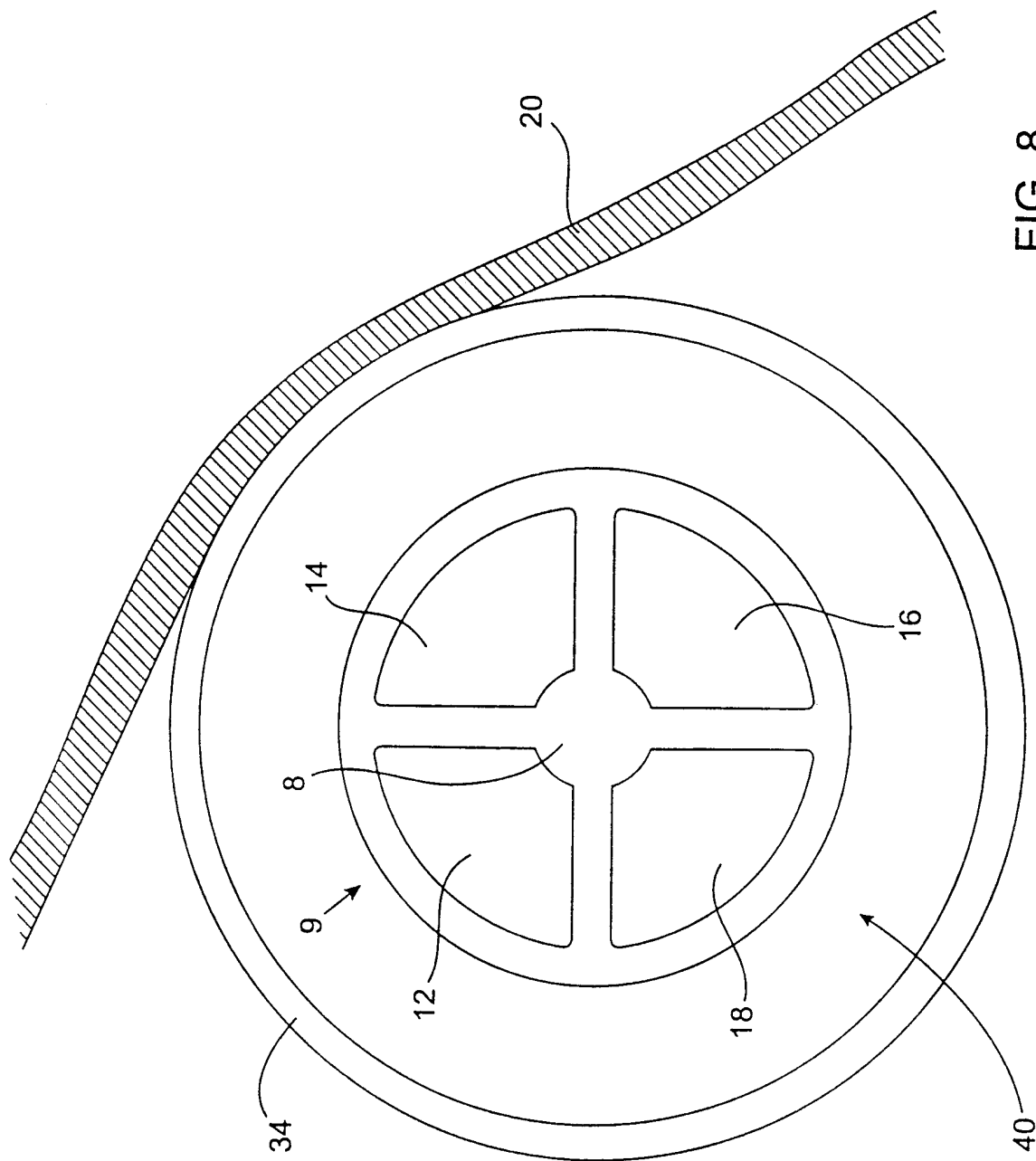


FIG. 8

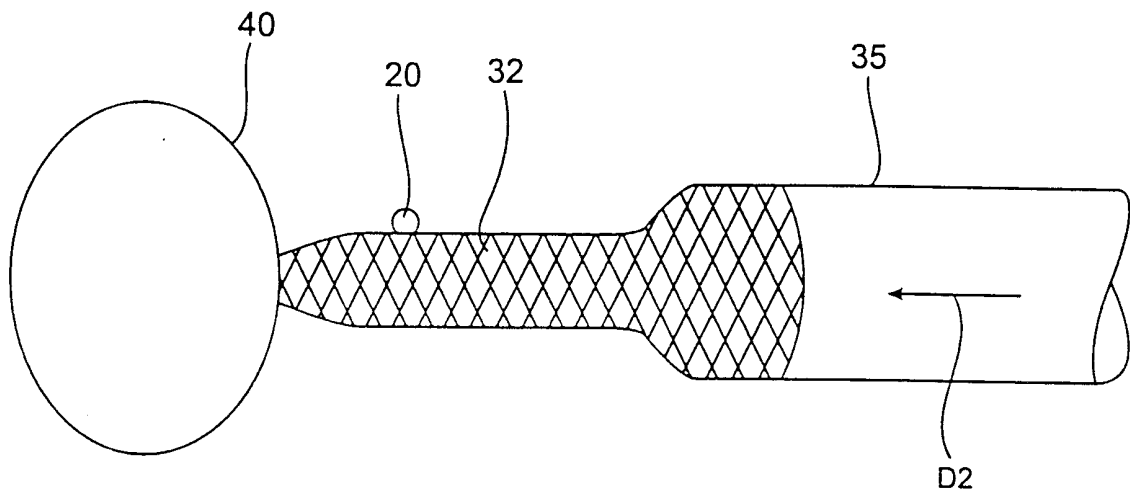


FIG. 9A

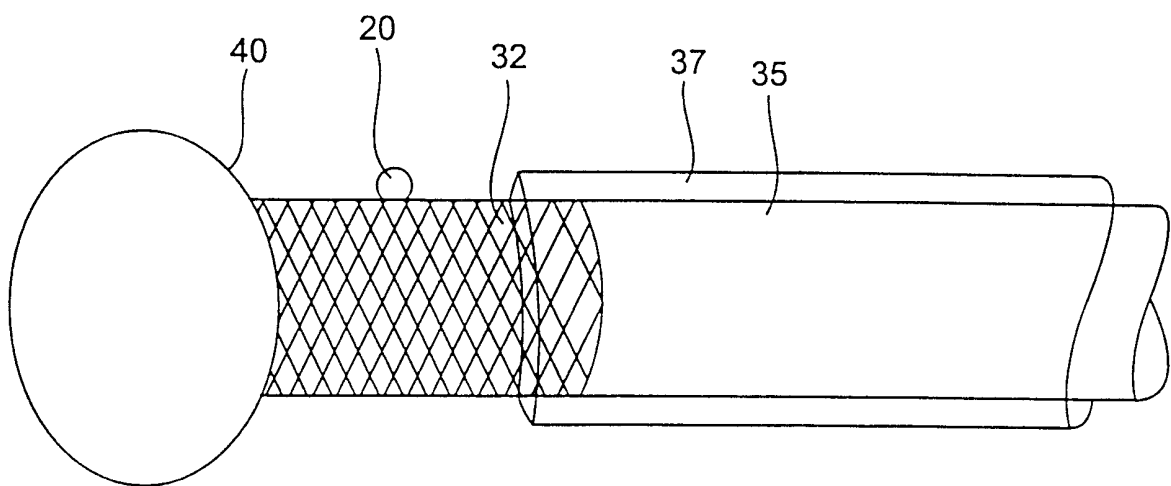


FIG. 9B

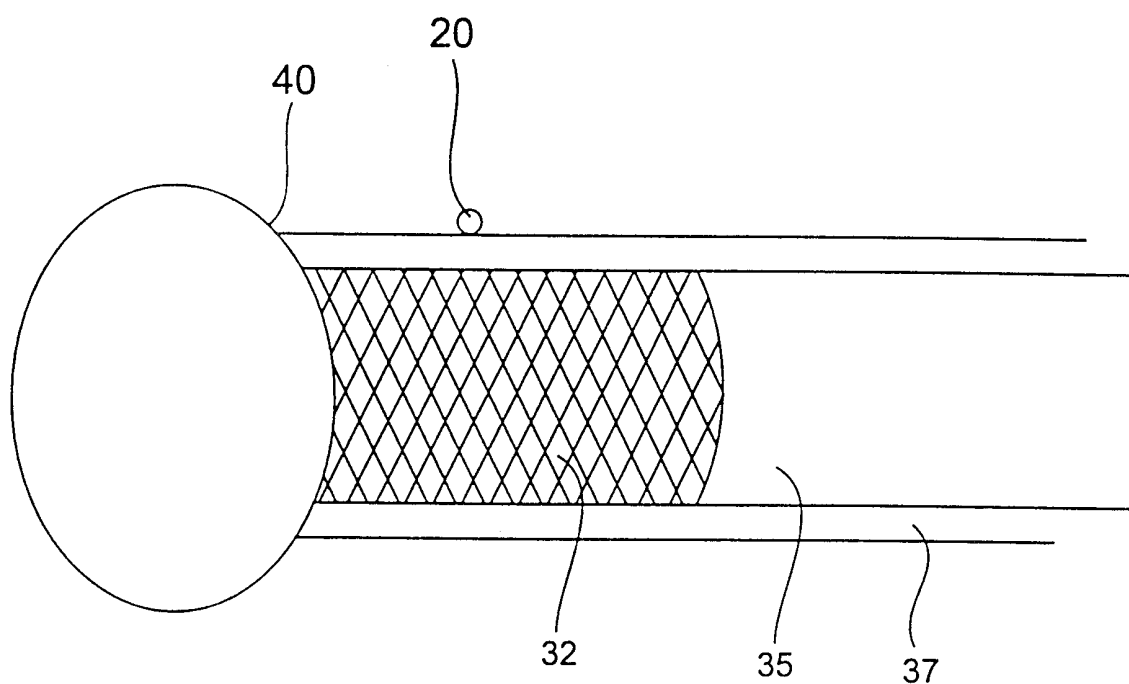


FIG. 9C

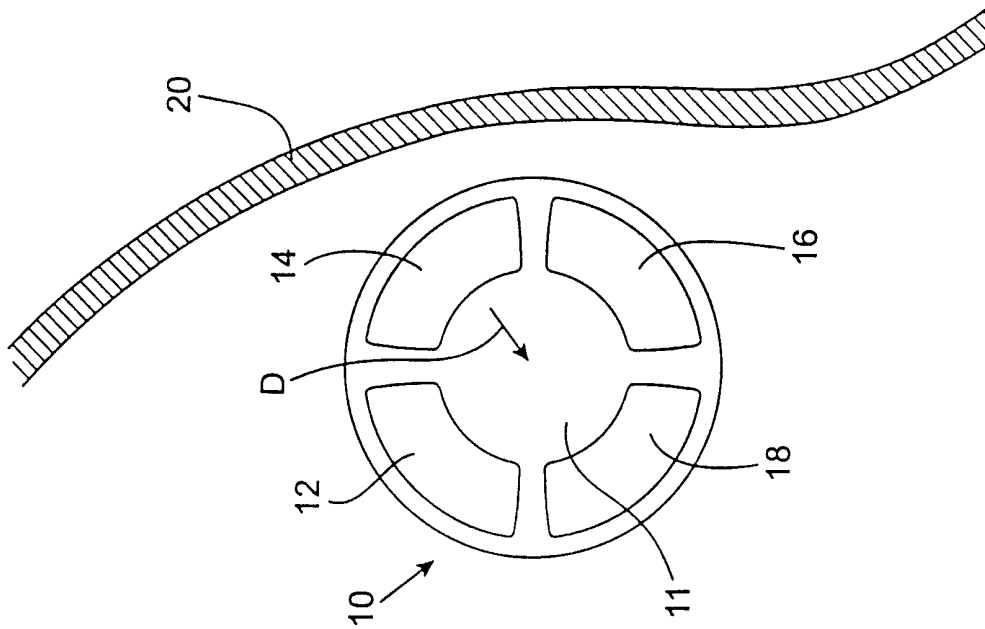


FIG. 10

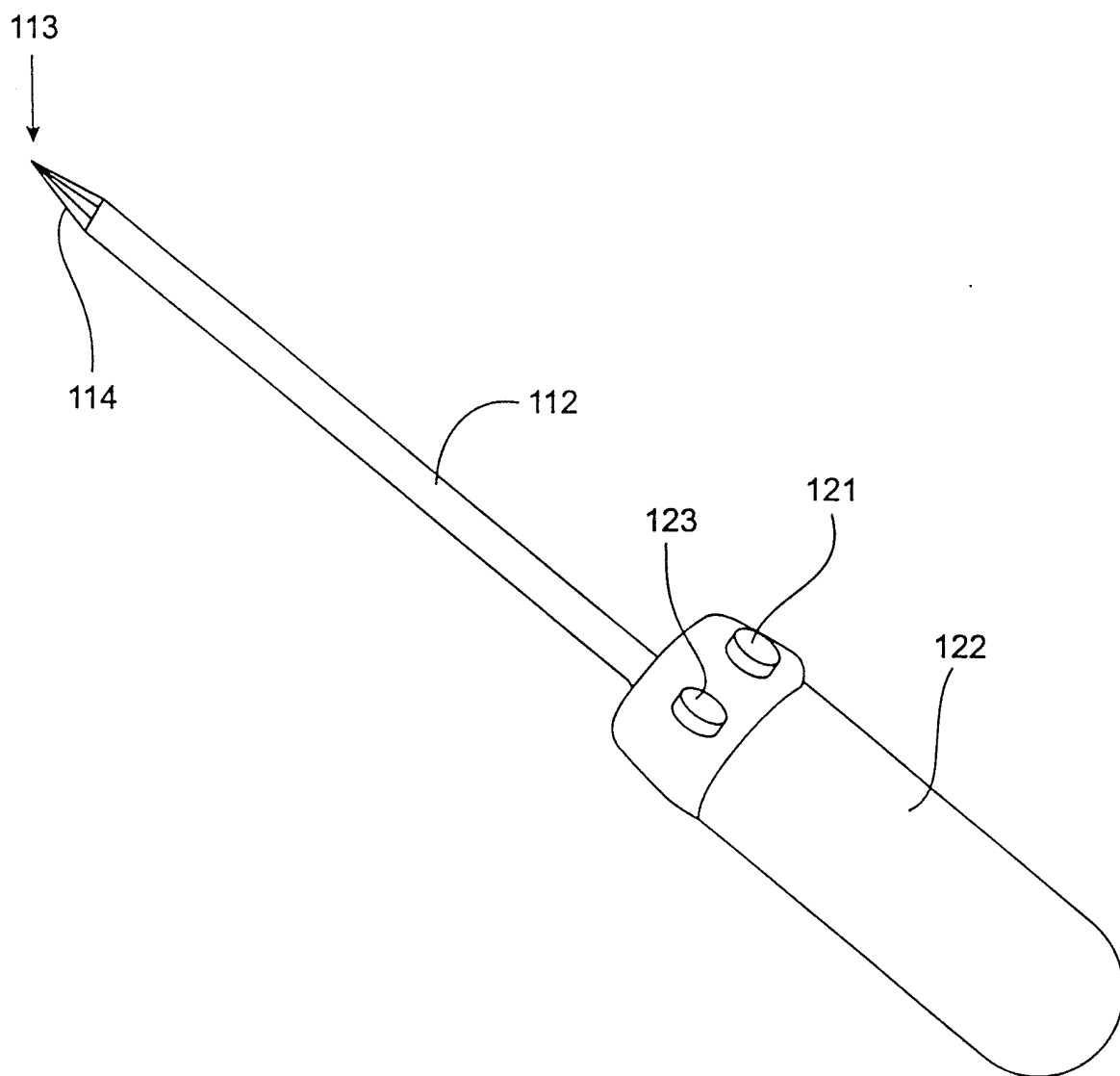


FIG. 11

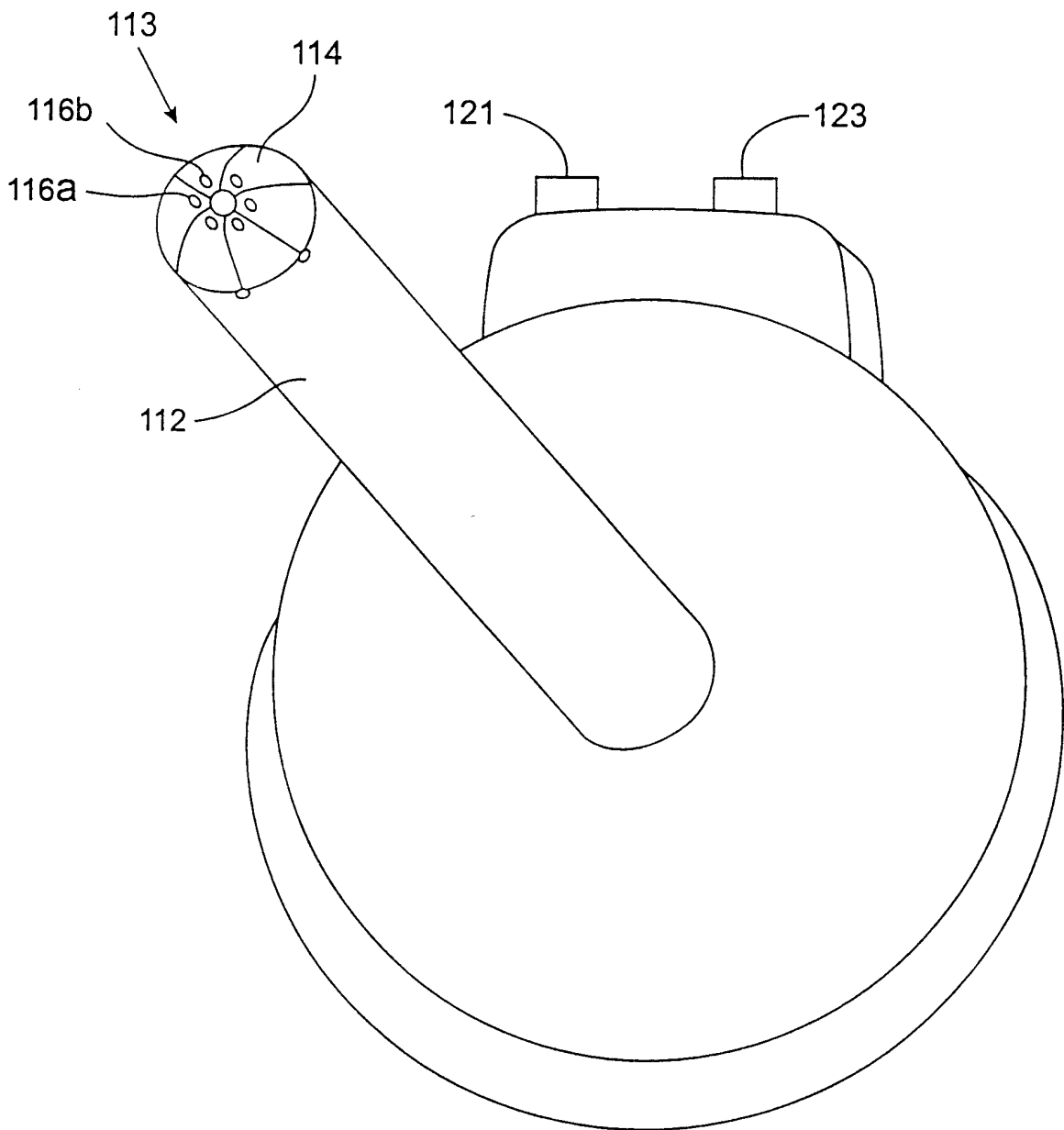


FIG. 12

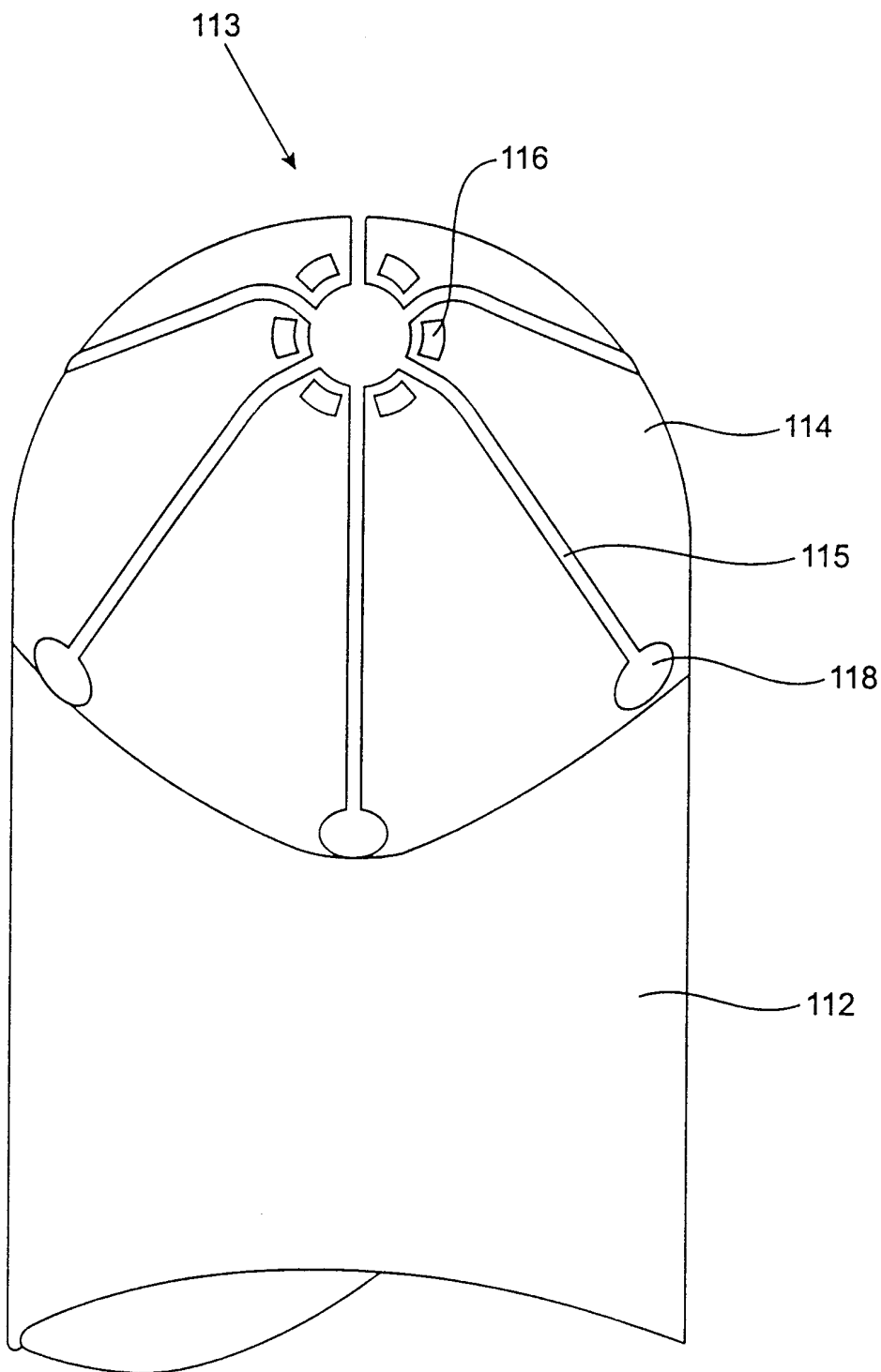


FIG. 13



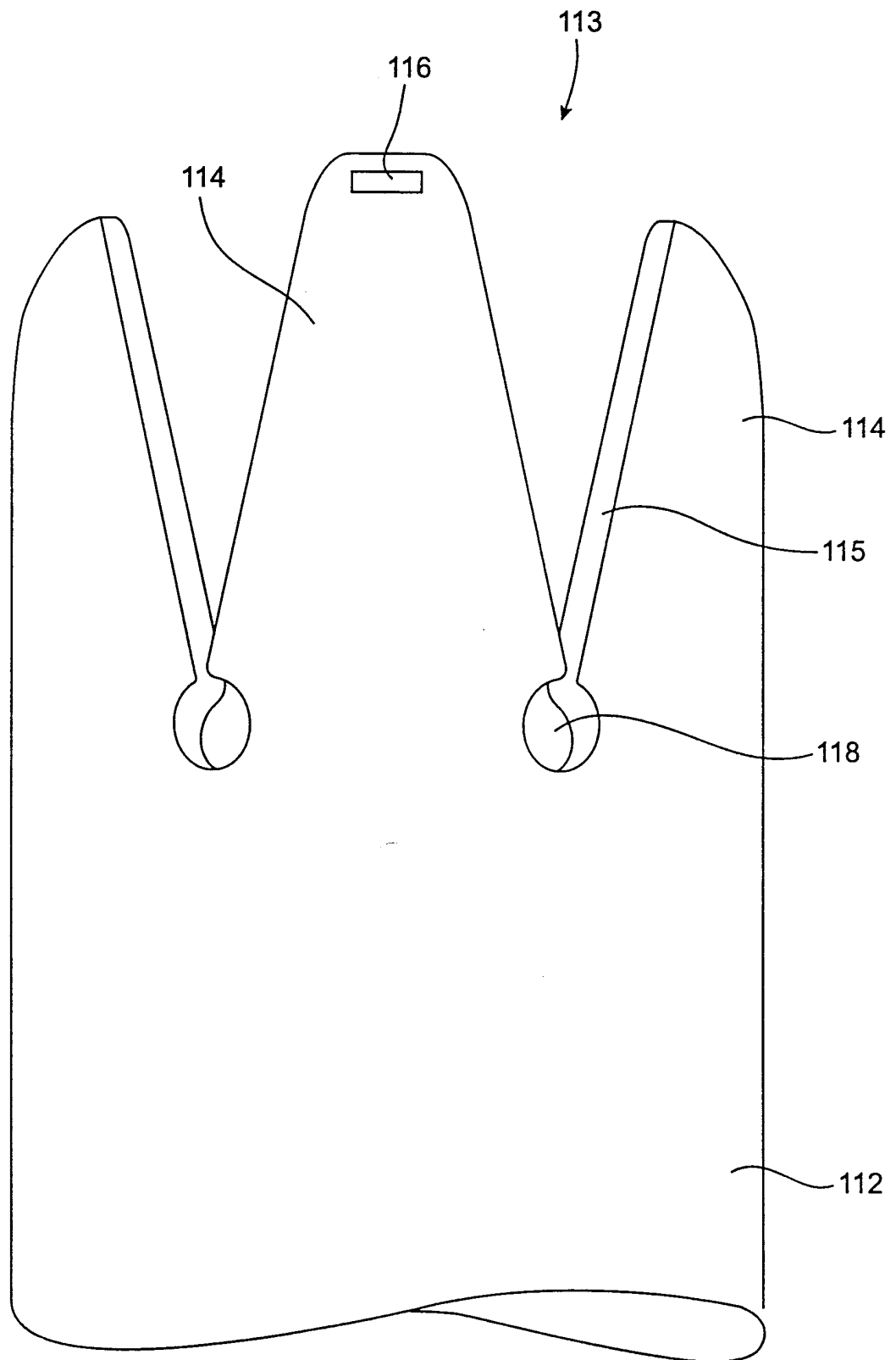


FIG. 14

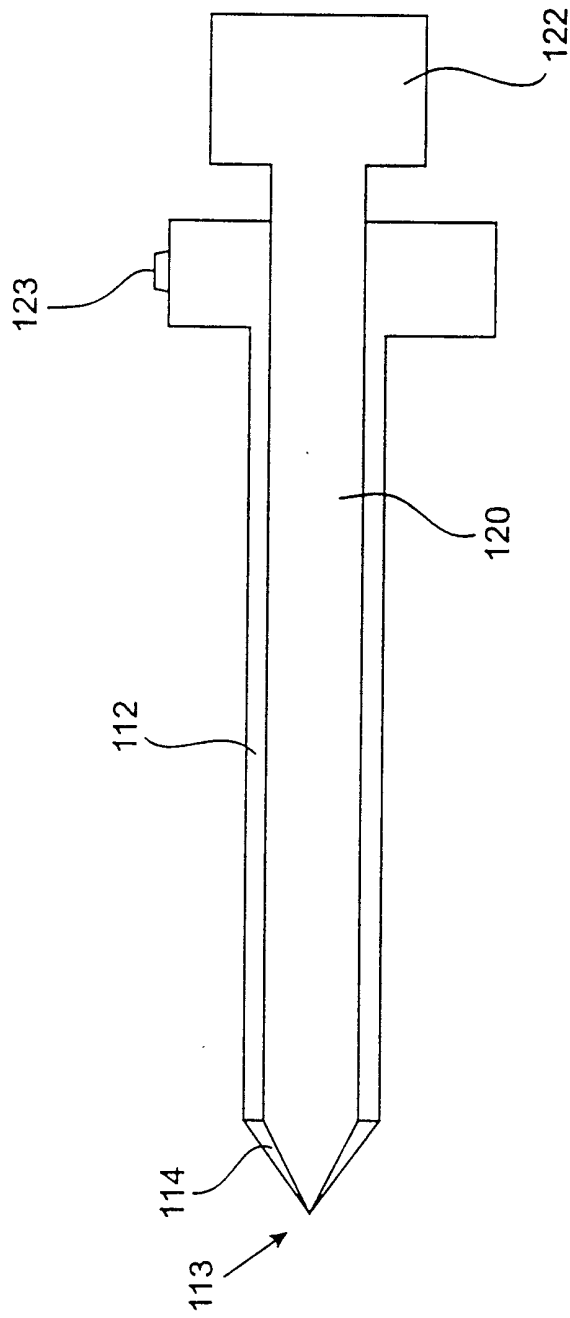


FIG. 15

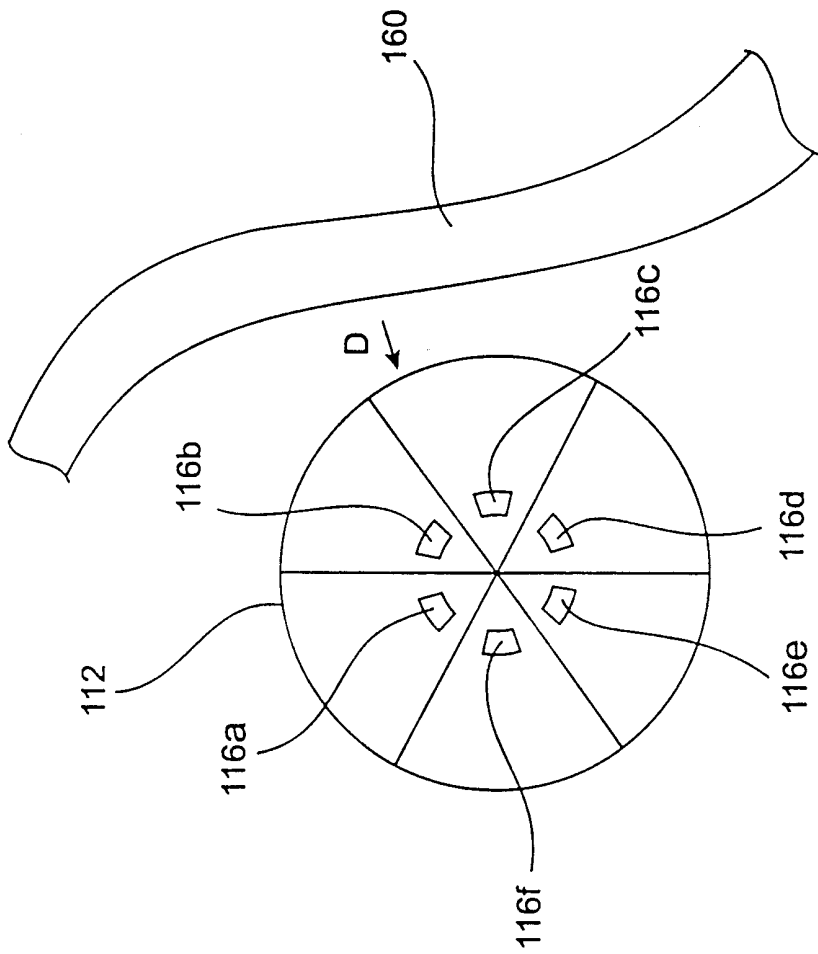


FIG. 16

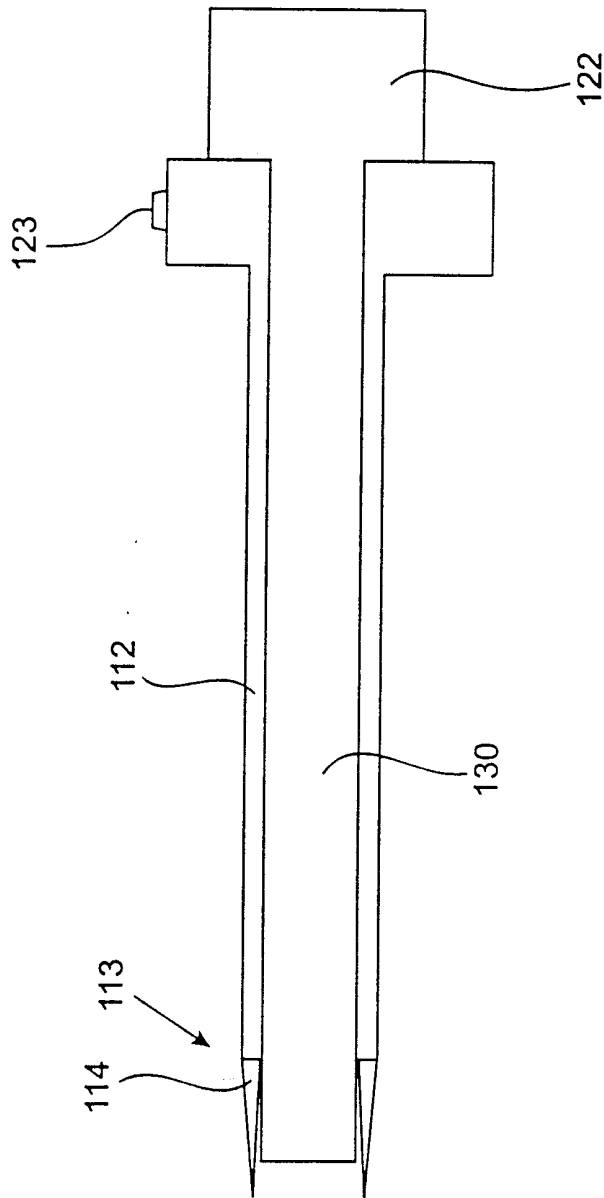


FIG. 17

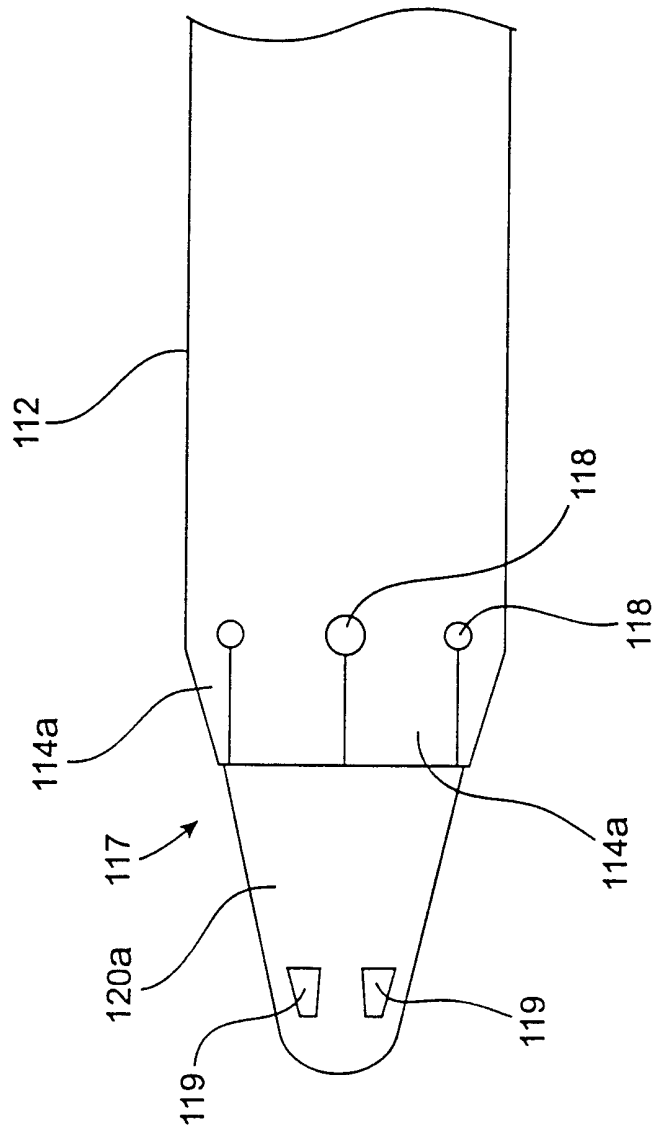


FIG. 18

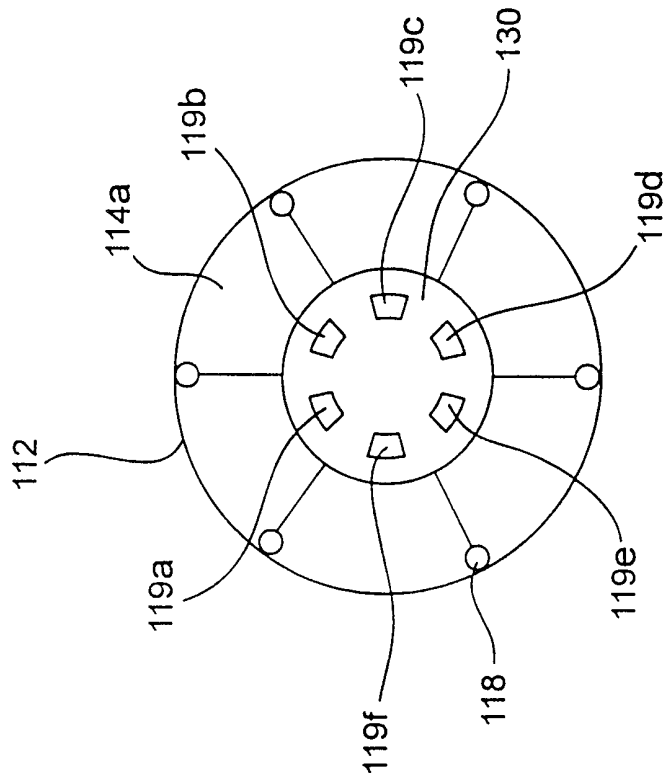


FIG. 19

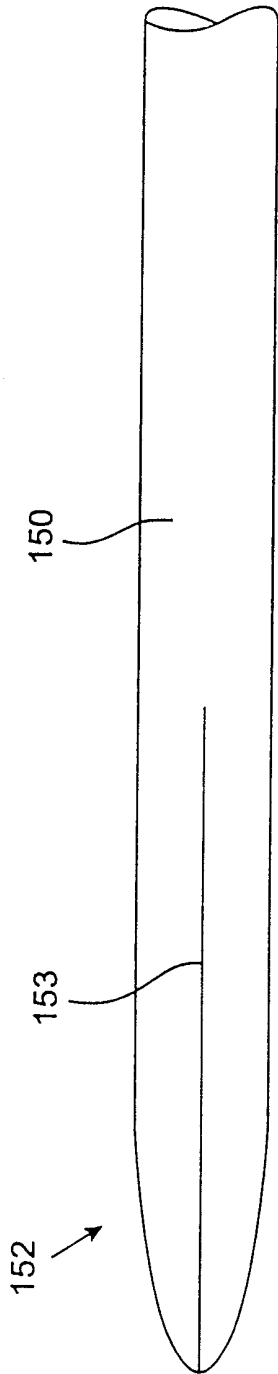


FIG. 20

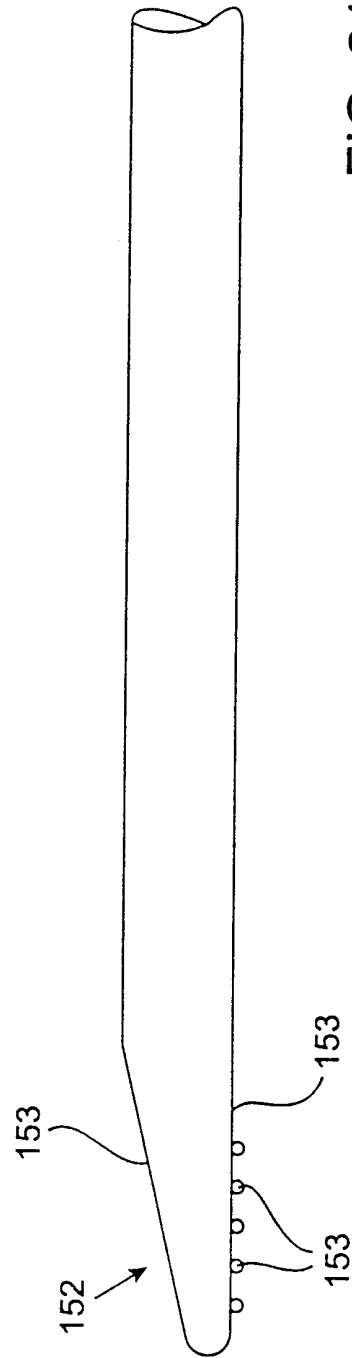


FIG. 21

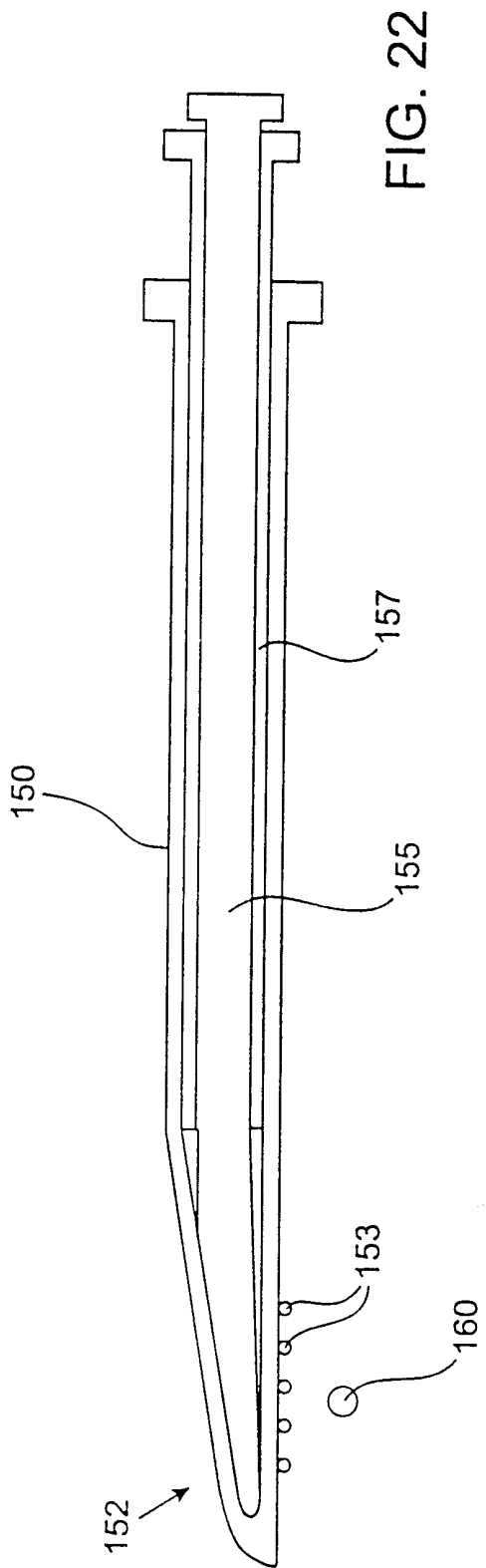


FIG. 22

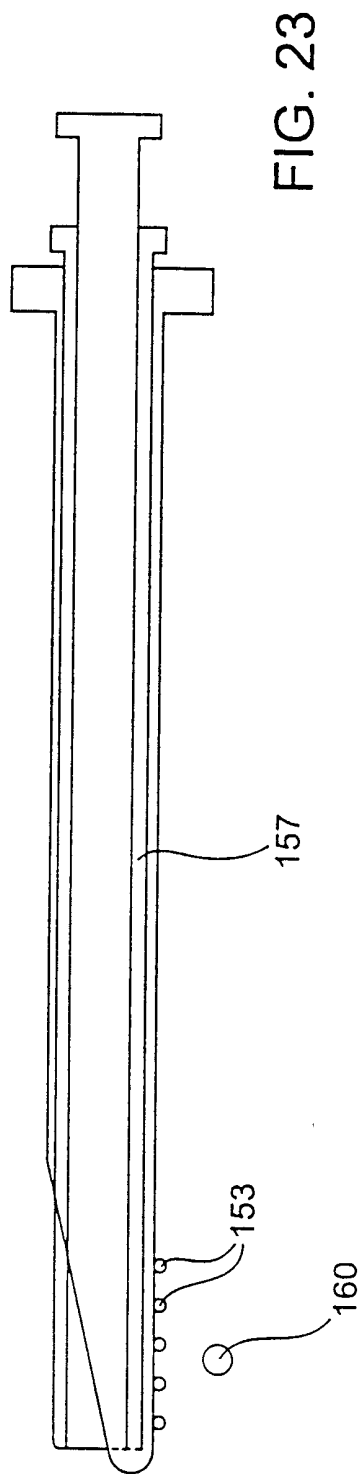


FIG. 23



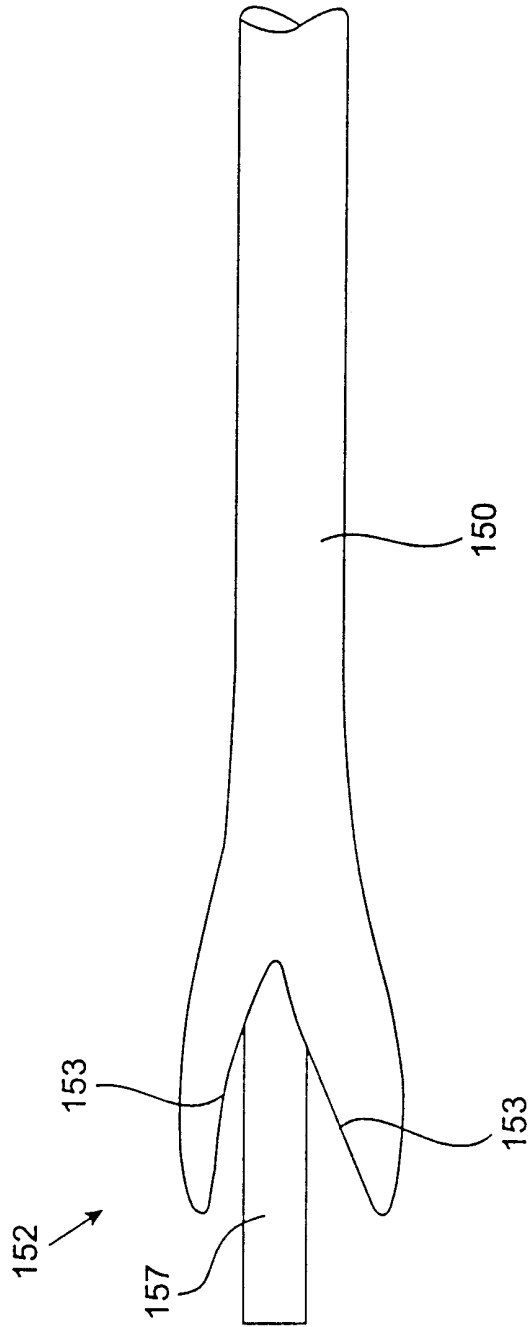
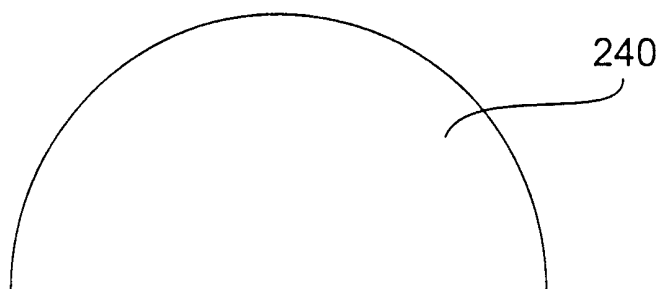
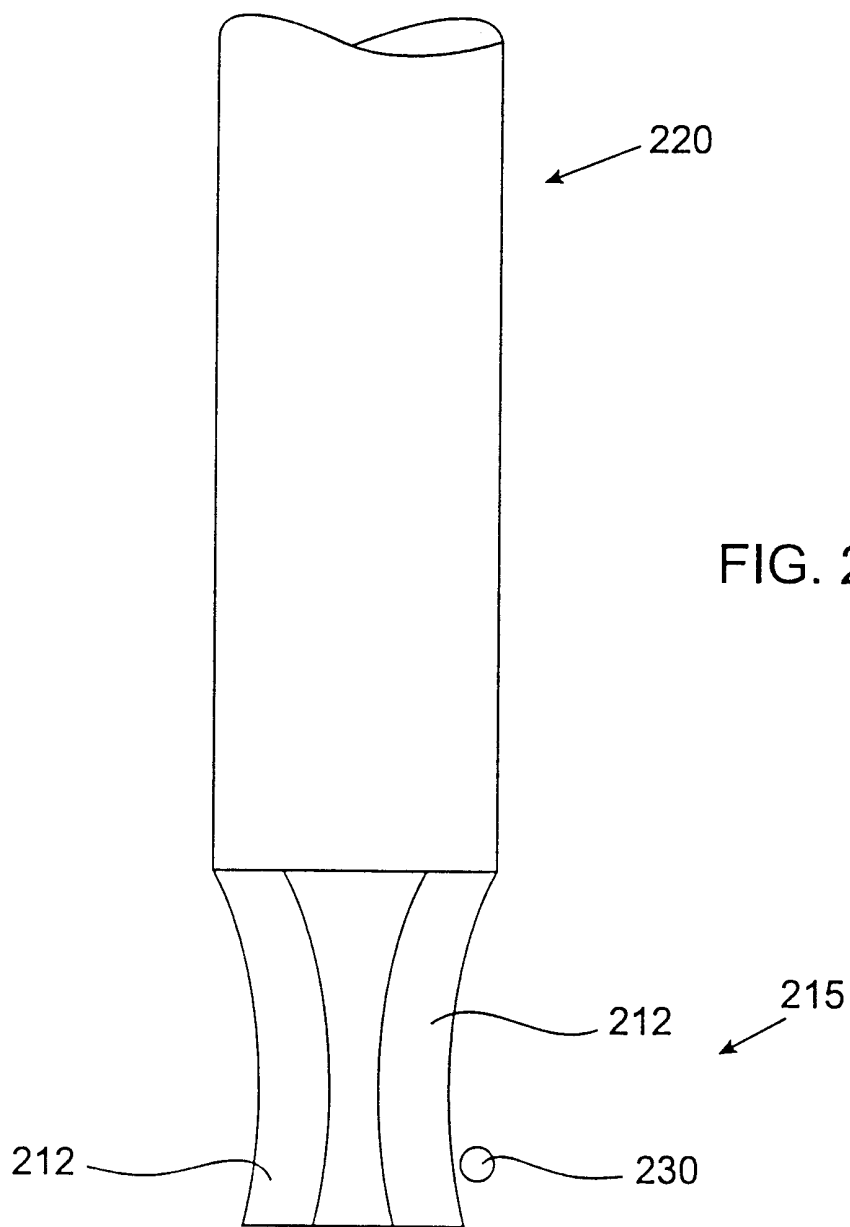


FIG. 24



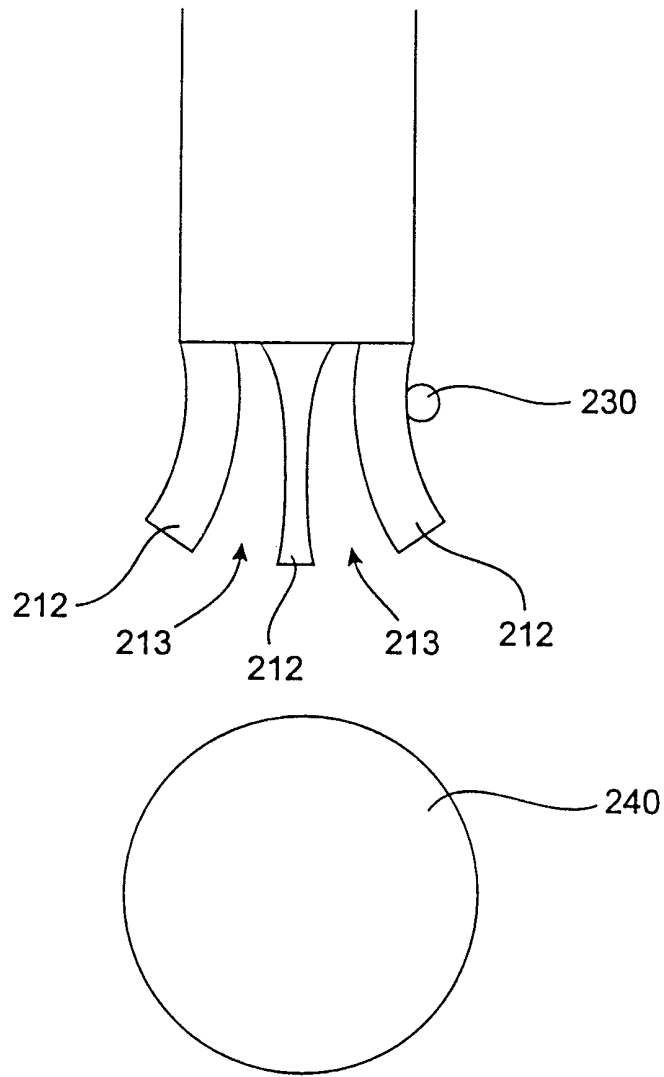


FIG. 26

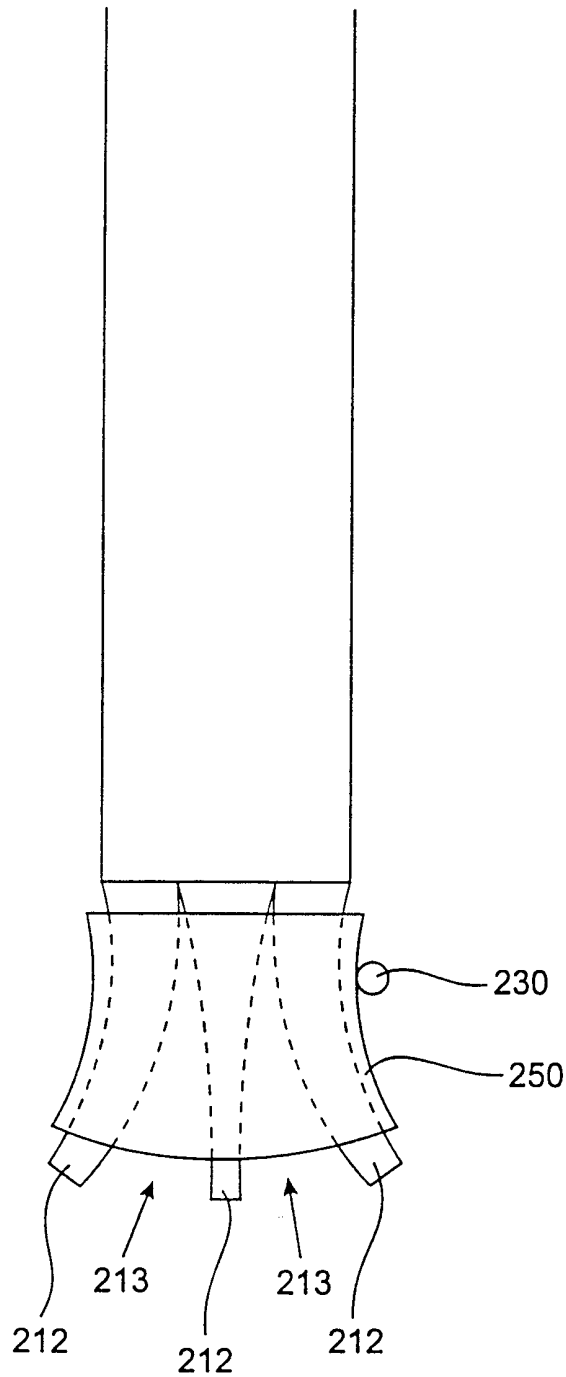


FIG. 27

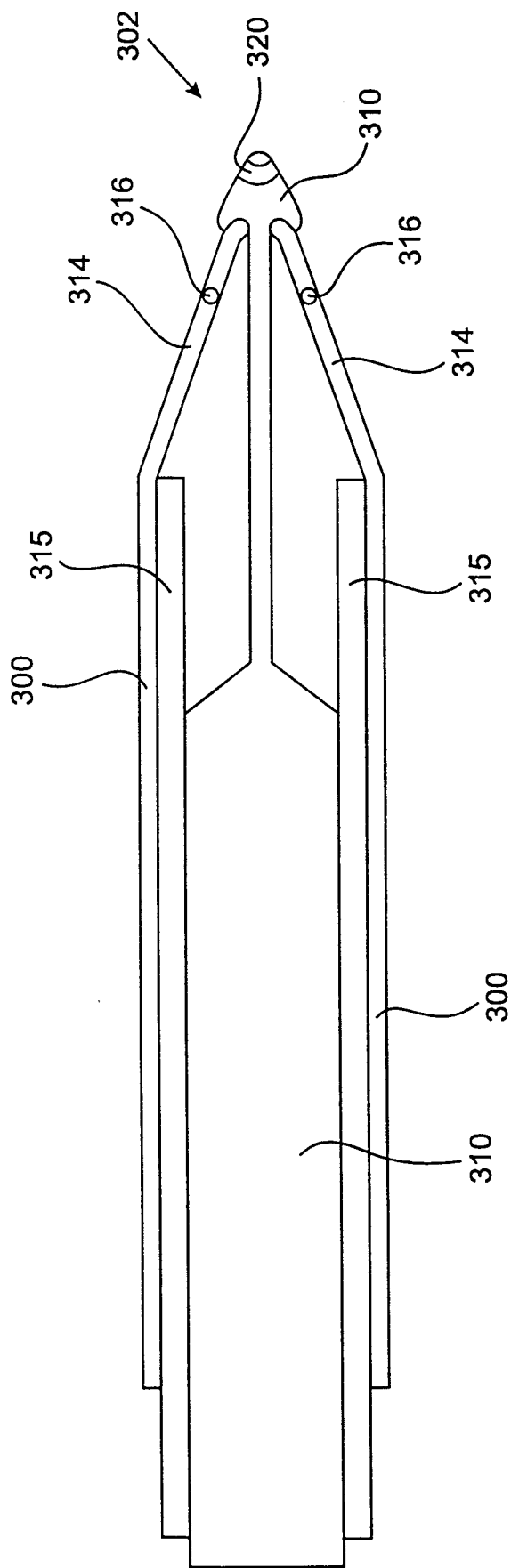


FIG. 28

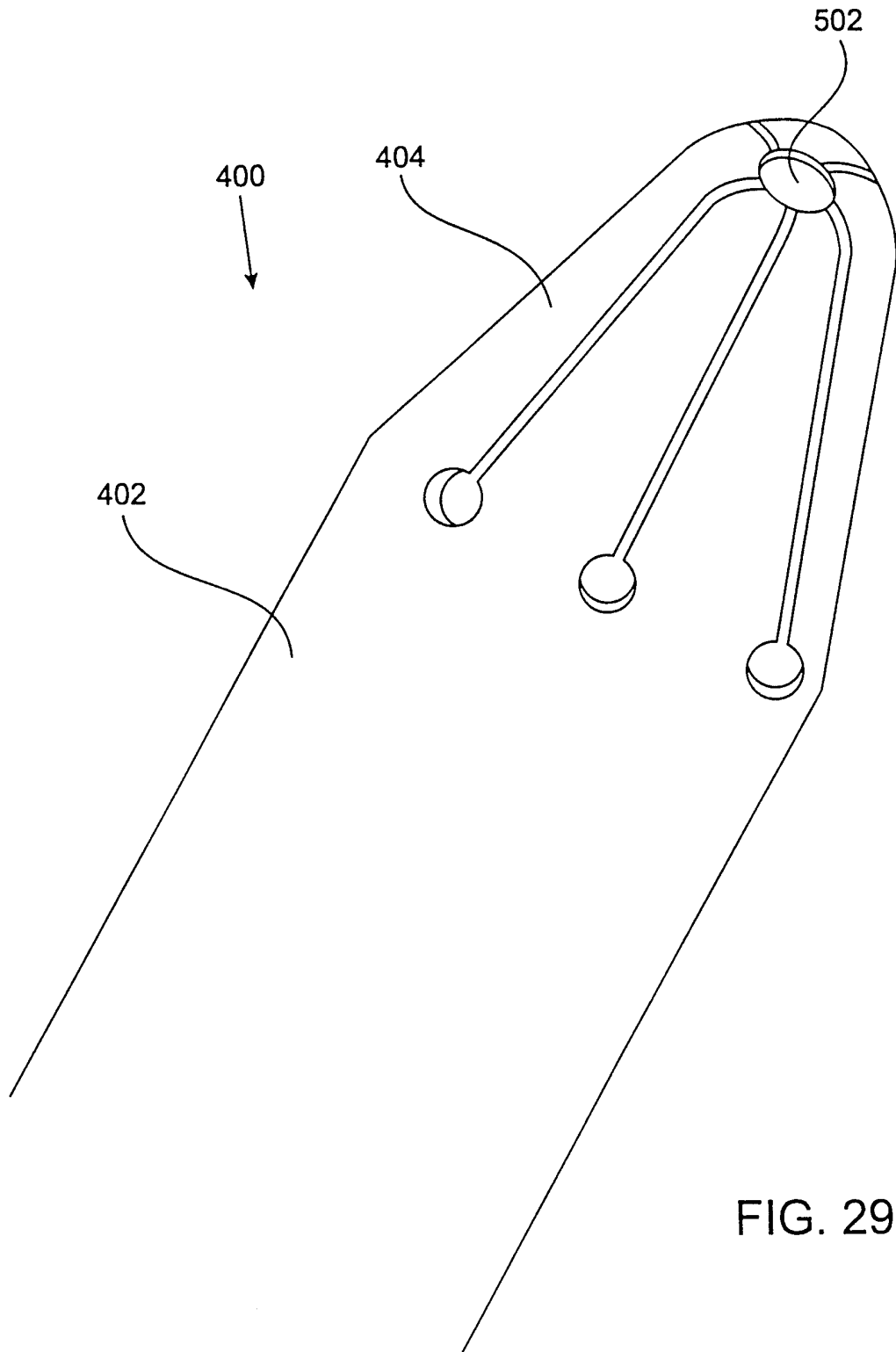


FIG. 29

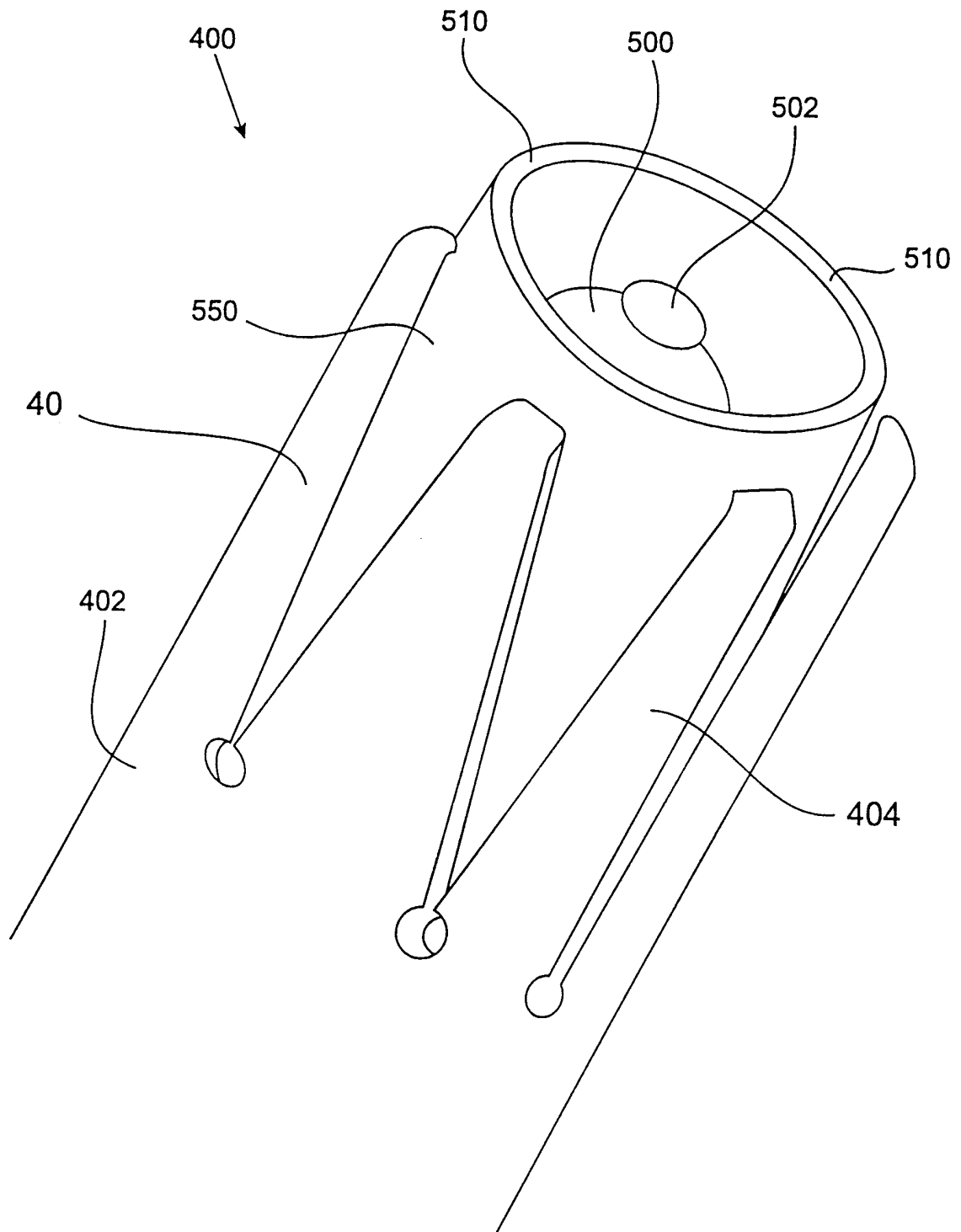


FIG. 30

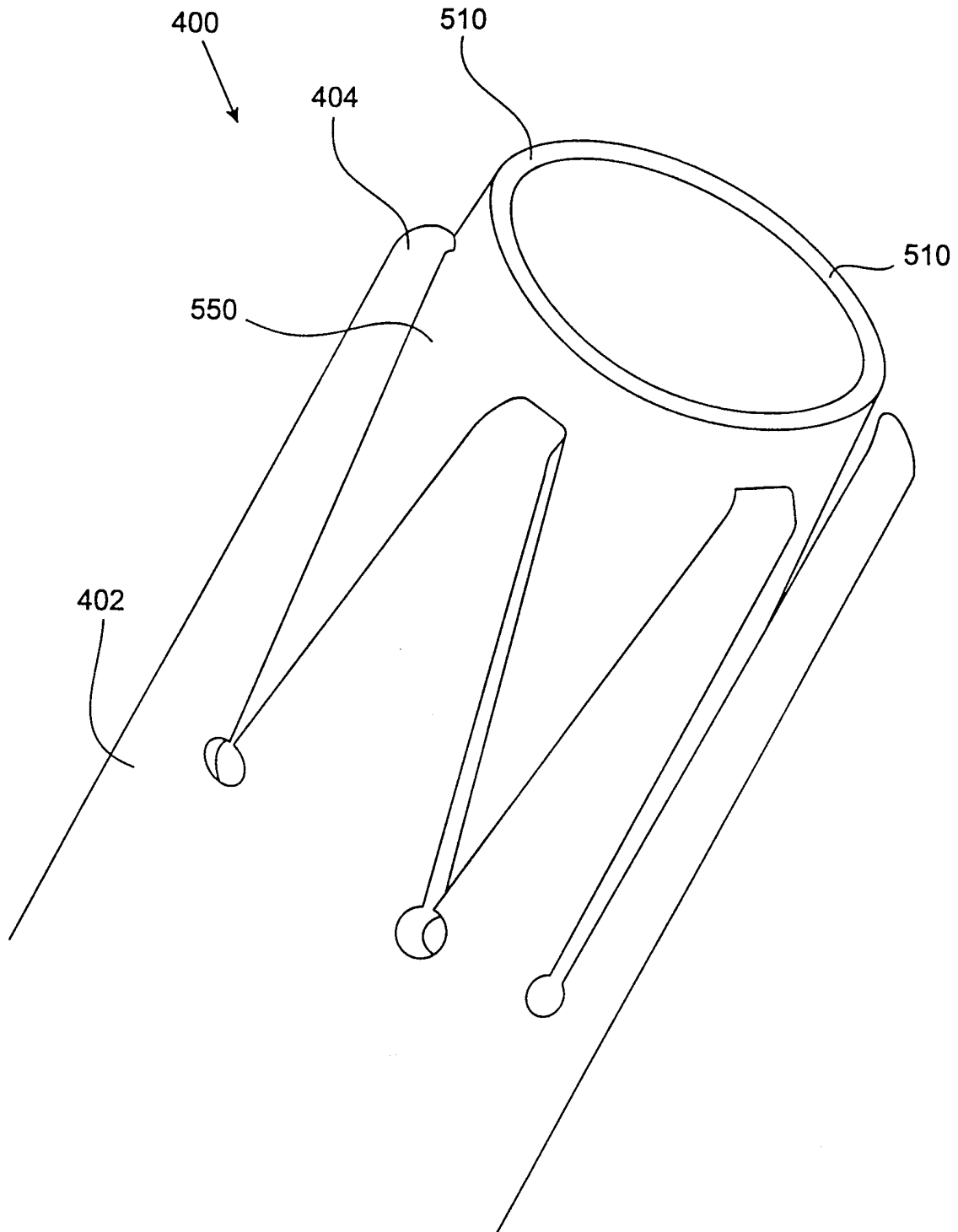


FIG. 31



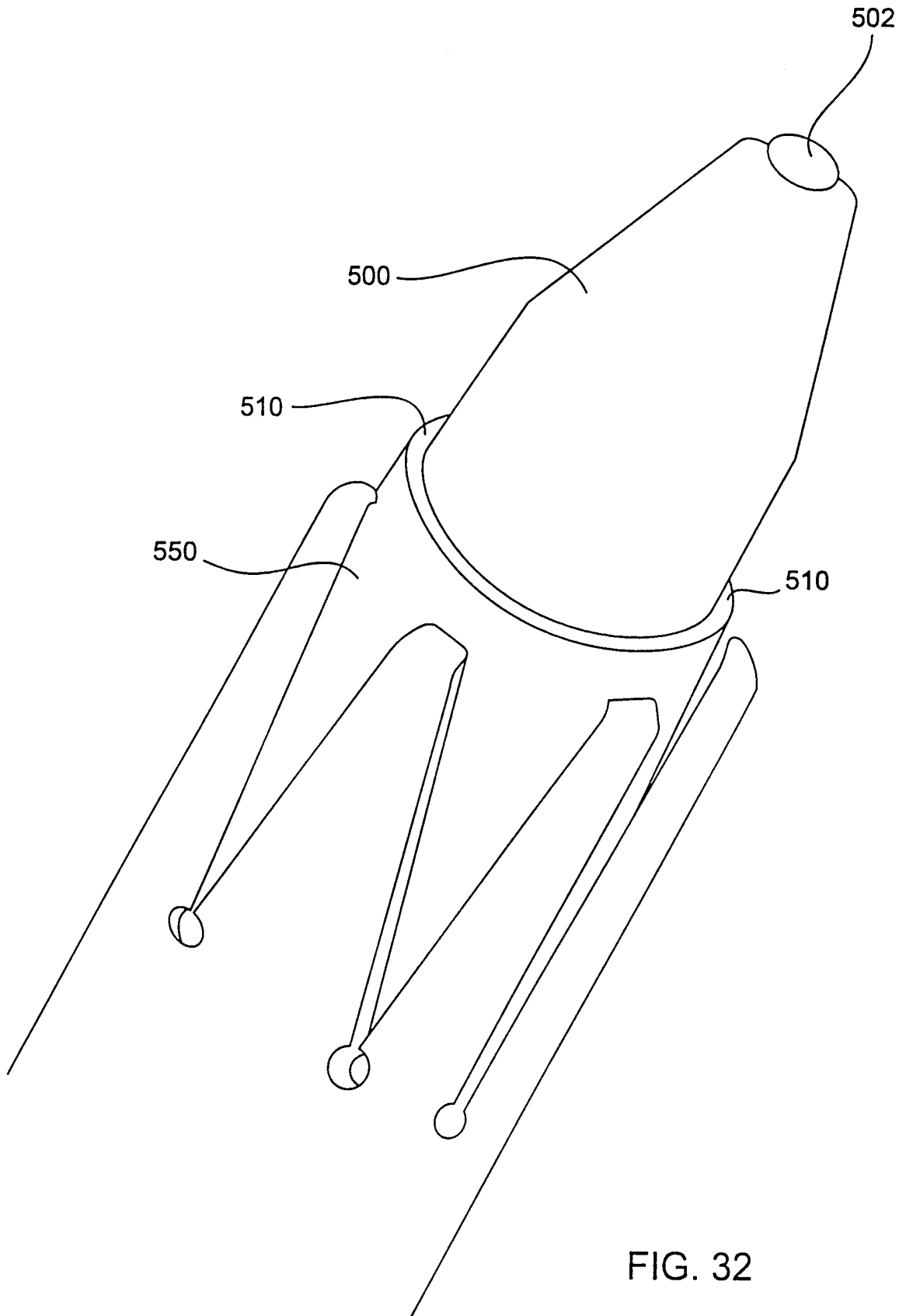


FIG. 32

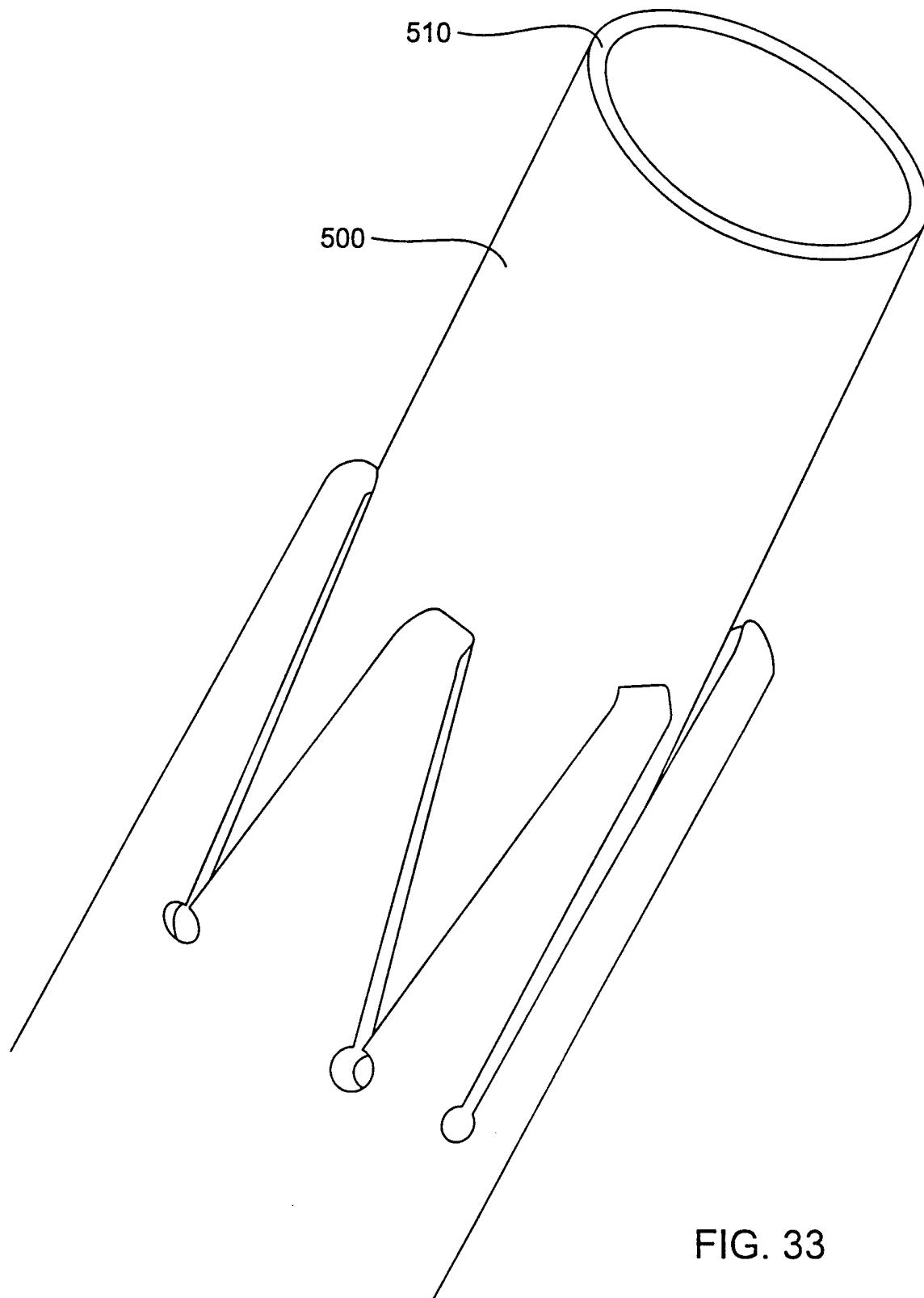


FIG. 33

INTERNATIONAL SEARCH REPORT

International application No.  
PCT/US99/12651

**A. CLASSIFICATION OF SUBJECT MATTER**  
 IPC(6) :A61B 5/0492, 17/39  
 US CL :600/373, 393, 546; 606/49, 50  
 According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)  
 U.S. : 600/373, 377, 393, 546; 606/46, 49, 50; 607/116-118

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)

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 3,957,036 A (NORMANN) 18 May 1976, entire document.	1-34
X	US 5,630,813 A (KIETURAKIS) 20 May 1997, col, 6 lines 1-58.	1, 4, 5, 15

Further documents are listed in the continuation of Box C.  See patent family annex.

* Special categories of cited documents:	"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
"A" document defining the general state of the art which is not considered to be of particular relevance	"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
"E" earlier document published on or after the international filing date	"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
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"&" document member of the same patent family
"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	

Date of the actual completion of the international search 13 SEPTEMBER 1999	Date of mailing of the international search report 17 NOV 1999
--	---

Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230	Authorized officer <i>Lee S. Cohen</i> LEE S. COHEN Telephone No. (703) 308-2998
---	---