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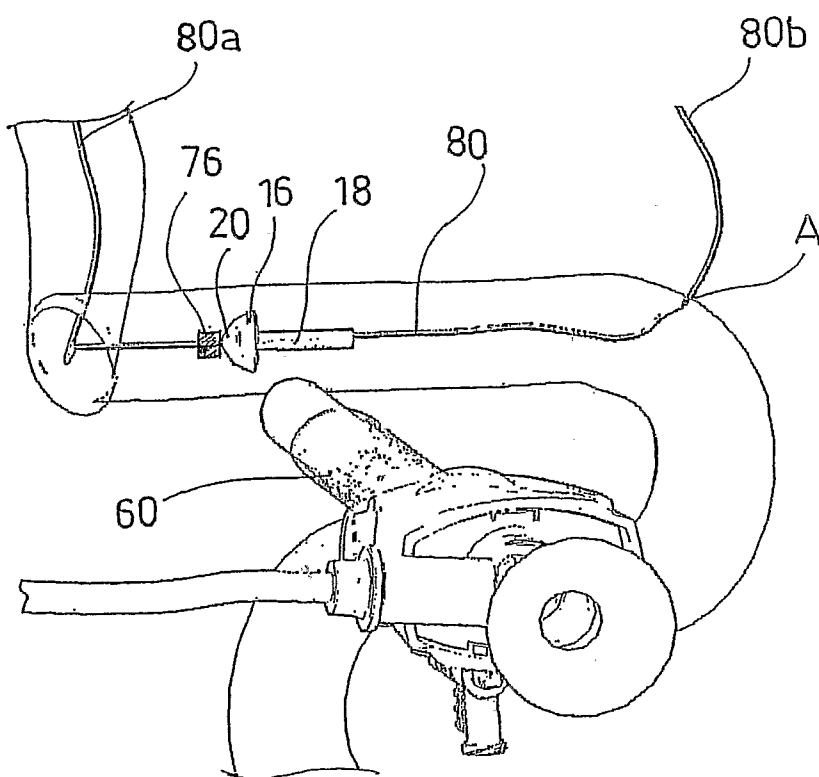
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(54) Title: A DEVICE AND METHOD FOR THE THERAPY OF OBESITY



(57) Abstract: A device for drawing tissues together and creating an anastomosis comprises a main guide wire (66, 86) suitable to receive and draw together portions (A; B) of tissues to be joined by means of anastomosis. The main guide wire (66, 86) is suitable to be shaped like an open ring crossing the portions (A, A'; B, B') of tissues to be joined by means of anastomosis (66, 86), wherein the ends of the main guide wire (66, 86) are different and distinguishable from each other. The main guide wire (66, 86) has a tubular or hollow structure suitable to house at least one needle for a push or radiofrequency perforation of the tissues.

**"A device and method for the therapy of obesity"**

The present invention relates to devices and methods for the therapy of obesity in general. 5 Particularly, the present invention relates to devices for drawing together tissues that are suitable to be used in a method for carrying out anastomosis in tracts of the digestive tube.

The present invention further relates to a method 10 for carrying out anastomosis in tracts of the digestive tube.

At present, surgical anastomoses are very difficult to carry out via endoluminal access. Most of anastomoses, in fact, are created by using open or 15 laparoscopic surgical techniques.

Accordingly, no effective surgical instruments are available which offer the guide and control required to suitably drawing together the tissue surfaces and/or connecting the surfaces with a passage (anastomosis) 20 through the body cavities.

The problem at the heart of the present invention is to provide devices capable of drawing together tissues and that can be used in a method for carrying out anastomosis in tracts of the digestive tube with 25 endoluminal access.

This problem is solved by means of a device for drawing together tissues and create an anastomosis according to claim 1.

Further characteristics and advantages of the 5 device and method according to the invention will result from the description below of preferred exemplary embodiments, which are given as a non-limiting indication, with reference to the attached figures, wherein:

10 Fig. 1 illustrates a perspective view of a circular stapler;

Fig. 2 illustrates a perspective view of a device to be associated with the circular stapler from Fig. 1;

15 Fig. 2a illustrates a perspective view of the stapler from Fig. 1, guide wire and device from Fig. 2 during an assembly step for performing a suture;

Fig. 3 illustrates a perspective view of a possible embodiment of a positioning device;

20 Fig. 4 illustrates a sectional view along a plane containing a longitudinal axis of the device from Fig. 3;

Fig. 5 illustrates a perspective view of a detail of the device from Fig. 3;

25 Fig. 6 illustrates a perspective view of a detail of the device from Fig. 3;

Fig. 7 and 8 illustrate perspective views of the detail from Fig. 5 from different points of view;

Fig. 9 illustrates a perspective view of a possible embodiment of a positioning device;

5 Fig. 10 illustrates a sectional view along a plane containing a longitudinal axis of the device from Fig. 9;

Fig. 11 illustrates a partially sectioned, perspective view of a detail of the device from Fig. 9;

10 Fig. 12 illustrates a perspective view of a detail of the device from Fig. 9;

Fig. 13 illustrates a perspective view of a possible embodiment of a positioning device;

15 Fig. 14 illustrates a sectional view along a plane containing a longitudinal axis of the device from Fig. 13;

Fig. 15 illustrates a perspective view of a detail of the device from Fig. 13;

20 Fig. 16 illustrates a perspective view of a detail of the device from Fig. 13;

Fig. 17-40 illustrate several steps of a method according to the present invention;

25 Fig. 41 illustrates a partial perspective view of a circular stapler, a positioning device and a guide wire, an anchoring ring being inserted thereon;

Fig. 42 illustrates the circular stapler, positioning device and guide wire from Fig. 41 while the positioning device is being inserted in the circular stapler;

5 Fig. 43 illustrates the Fig. 41 in a longitudinal section;

Fig. 44 illustrates Fig. 41 from a different point of view.

With reference to Fig. 2A and Fig. 41-44, there is  
10 generally illustrated a device for drawing together tissues and create an anastomosis comprising a main guide wire suitable to receive and drawing anastomotic devices in order to drawing together tissue portions to be joined by means of anastomosis. With reference to  
15 the examples that will be described below, the main guide wire has been designated in Fig. 17-40 with the numeral 66 or 86. Furthermore, the anastomotic device can be a circular stapler 10 provided with an anvil 16 (such as illustrated in the Fig. 2a and 41-44) or a  
20 positioning device 24 such as will be described below.

The main guide wire is suitable to be shaped as an open ring crossing the tissue portions to be joined by means of anastomosis. Advantageously, the ends of the main guide wire are different and distinguishable from  
25 each other.

In accordance with a possible embodiment, the main guide wire has a tubular or hollow structure suitable to house at least one needle for the push or radiofrequency perforation of the tissues.

5 An anchoring ring 76 is suitable to be made integral with the main guide wire in order to abut against the anastomotic device and draw the latter by drawing an end of the main guide wire. In the case of the positioning device 24, the anchoring ring 76 abuts 10 against a proximal component 26 as will be described below. In the case of the circular stapler 10 the anchoring ring 76 abuts against the anvil 16.

With reference to Fig. 1, with 10 has been indicated a circular stapler comprising a handle 12 and 15 a stem 14 as a whole. The structure of the circular stapler is similar to the known circular staplers, which are conventionally used to carry out circular anastomosis, such as of the intestine. The structure of the circular stapler is changed compared with the 20 conventional ones in that, in a preferred embodiment thereof, it has a channel suitable to receive the guide wire. In Fig. 2A, the circular stapler 10 has a channel crossing the stem 14 thereof from the distal end to an area at the proximal end from which it protrudes 25 outside, for example on one side. In accordance with

different embodiments, not illustrated, the guide wire runs all along the length of the circular stapler or only a distal portion thereof.

The channel is suitable to receive a guide wire 5 (not illustrated in Fig. 1) such that the stapler can slide therealong and be placed in the site requiring anastomosis. An exemplary use of the circular stapler 10 will be described below with particular reference to Fig. 29. The length of the stem 14 and the diameter 10 thereof are sufficient to carry out the method and reach the desired site.

Advantageously, the stem is made of a flexible material, such as to facilitate reaching the site requiring anastomosis.

15 The stapler 10 advantageously comprises an anvil 16 illustrated for example in Fig. 2. The anvil 16 defines an exemplary device for drawing tissues together, particularly a device that, besides drawing the tissues together, is suitable to be associated with the circular 20 anvil 10 such as illustrated in Fig. 1 in order to carry out anastomosis.

The anvil 16 comprises a stem 18 and a head 20. The stem 18 has such a longitudinal and cross size that makes it suitable to be fit on the end of the stem 14 of 25 the circular stapler 10 opposite the handle 12 (Fig. 2A

and 41-44).

Advantageously, a channel 22 crosses the anvil 16 in the longitudinal direction and is suitable to receive a guide wire, not illustrated in Fig. 2. An exemplary 5 use of the anvil 16 for the circular stapler 10 will be described below with particular reference to the Fig. 26-29.

In Fig. 3, with 24 has been generally designated a positioning device suitable to draw tissues together 10 which have been subjected to enterostomy, and to place means for providing a passage (anastomosis) between the tissues that have been drawn together.

The positioning device 24 comprises a first component, or proximal component, designated with 15 numeral 26 and a second component, or distal component, designated with reference 28. Preferably, the positioning device 24 extends along a longitudinal axis 30. Fig. 5, 7 and 8 illustrate perspective views of the proximal component 26, whereas Fig. 6 illustrates a 20 perspective view of the distal component 28.

In accordance with a possible embodiment, the proximal component 26 is suitably shaped to be abutted against the edge of a first enterostomy in order to draw the tissues adjacent thereto against the tissues 25 adjacent to a second enterostomy. The distal component

28 is suitably shaped to be inserted through the enterostomies.

For clarity purpose, the first enterostomy will be also called herein below as the proximal enterostomy, 5 whereas the second enterostomy will be also called the distal enterostomy. With reference to a possible embodiment, the first enterostomy can be a gastrostomy and the second enterostomy can be a jejunostomy. With reference to a different embodiment, the first 10 enterostomy can be a proximal jejunostomy and the second enterostomy can be a distal jejunostomy.

In accordance with a possible embodiment, the proximal component 26 has a substantially cylindrical outer structure. A cavity 32 being formed at one of the 15 bases of the cylindrical structure and longitudinally thereto, preferably has a first portion defined by a surface having the shape of a truncated cone 32a and a second portion defined by a cylindrical surface 32b. The cavity 32 does not run through the entire length of 20 the proximal component 26, leaving a base wall 34. Furthermore, the cross size of the cavity 32 and the proximal component 26 are preferably such as to leave an abutment surface, for example a plane annular surface 26a contouring the cavity.

25 According to a possible embodiment, a lug 36,

preferably cylinder shaped, extends along the longitudinal axis 30 from the bottom of the cavity 32 towards the outside of the cavity, preferably such that a free end 36a of the lug 36 is completely out of the 5 cavity 32. In other words, the length of the lug 36 from the base of cavity 32 to the free end 36a thereof is preferably greater than the depth of the cavity 32. The lug 36 has a preferably cylindrical cavity 38 extending along the longitudinal axis 30 and crossing the base 10 wall 34 leading to the opposite surface of the proximal component 26. In other words, the cavity 38 involves the lug 36 and base wall 34 thereby generating a duct open at the ends thereof and suitable to receive a guide wire not illustrated in Fig. 3-8. Preferably, the proximal 15 component 26 comprising the lug 36 is made as one piece.

According to a possible embodiment, the proximal component 26 comprises holes 40 for example for a suture, which can be used for separating the proximal component from the distal component, to be passed 20 therethrough.

In accordance with a possible embodiment, the distal component 28 comprises a head 42 and a stem 44, which are preferably made as one piece, that develop along the longitudinal axis 30.

25 The head 42 preferably has a shape of a truncated

cone and, according to a possible embodiment, comprises holes 46 for a suture, which can be used for example to separate the distal component from the proximal component, to be passed therethrough.

5 In accordance with a possible embodiment, the stem 44 preferably has a cylindrical structure and a free end thereof, i.e. opposite the head 42, widens to form a preferably annular base 48.

A channel 50 extends along the longitudinal axis 30  
10 from the end of head 42 to the base 48 and is suitable  
to receive a guide wire therein, not illustrated in Fig.  
4 or 6. The cross size of channel 50, at least in the  
portion at the base 48, are such as to receive the lug  
36 of the proximal component 26 therein. In other words,  
15 the channel 50 preferably has a larger section at the  
area in which it receives the lug 36. Preferably, the  
remaining part of channel 50 has the same cross size as  
the cavity 38.

Fig. 3 and 4 illustrate the positioning device 24  
20 when assembled. The proximal component 26 and the distal  
component 28 are joined such that the cavity 38 and  
channel 50 define a channel running all along the  
assembly for introducing a guide wire, not illustrated  
in Fig. 3 and 4. Particularly, Fig. 4 illustrates the  
25 positioning device 24 sectioned along a plane comprising

the longitudinal axis 30. In the assembly position, the proximal component 26 and the distal component 28 lock an elastic ring 52 therebetween, which is hold in a compressed/deformed configuration, and suitable to be 5 placed by the positioning device 24 in a desired anastomotic site in which, after it has been positioned, the elastic ring 52 takes a preset non-compressed rest shape (see for example Fig. 37). The elastic ring can be made of Nitinol, stainless steel or other satisfying 10 materials.

In accordance with a possible embodiment, the elastic ring 52 in its deformed configuration, has an end, such as the proximal (designated with numeral 52a), that is locked between the base 48 of the distal 15 component 28 and the inner diameter of the cavity 32 of the proximal component 26, i.e. inside the cylindrical portion 32b of the cavity 32. The opposite end of the elastic ring 52, i.e. the distal end designated with the numeral 52b, is preferably unfastened and abuts beneath 20 the head 42 of the distal component 28. In this case, it is advantageously provided that the cross size of the distal end 52b of the elastic ring 52 does not exceed the cross size of the head 42 of the distal component 28.

25 An exemplary use of the positioning device 24 will

be described below with particular reference to the Fig. 34-37. Upon use, the outer diameter of the proximal component, and particularly with the flat annular surface 26a, is intended to act as a striker against the 5 wall of the tissue to be drawn together, or in other words, abut against the proximal enterostomy while it minimizes the risk of penetration in the wall.

The distal component 28, with its head 42, is intended to penetrate in the proximal and distal 10 enterostomies and protects the elastic ring 52 while being inserted and positioned, such as will be described in the following.

Fig. 9-12 illustrate a possible variant embodiment of the positioning device 24 and the proximal and distal 15 components thereof according to the present invention. The elements in common have been designated with the same numeral used in Fig. 3-8 and will be described below with reference to the differences from the above embodiment.

20 The proximal component 26 is substantially similar to the one illustrated in Fig. 3-5, 7 and 8. As regards the distal component 28, the stem 44 extends straight up to its free end opposite the head 42 that does not widen to form a base similar to the base 48 of the distal 25 component described above. Furthermore, the head 42,

preferably having the shape of a cone or truncated cone, comprises a flange 54 extending from the outer perimeter of the major base of the head to form a circular wall substantially parallel to the longitudinal axis 30.

5       Fig. 9 and 10 illustrate the positioning device 24 when assembled, in which the channel 50 and the cavity 38 define a channel extending along the longitudinal axis 30 through the entire length of the assembled positioning device 24 to receive a guide wire, not  
10 illustrated in the Fig. 9-12. In the assembled configuration of the positioning device 24, the elastic ring 52 is held between the proximal component 26 and the distal component 28 in a compressed/deformed configuration. After the elastic ring has been  
15 positioned, it takes a preset, uncompressed rest shape as described above. In the deformed configuration, the proximal end 52a of the elastic ring 52 is fastened by the inner diameter of the cavity 32; particularly by the cylindrical portion 32b of the cavity 32, whereas the  
20 distal end 52b of the elastic ring 52 is fastened within the circular flange 54 of the distal component 28.

The cross size of channel 50, at least in the portion at the base 48, are such as to receive the lug 36 of the proximal component 26 therein. In other words,  
25 the channel 50 has a larger section at the area where it

receives the lug 36. Preferably, the remaining part of channel 50 has the same cross size as the cavity 38. Also in this case, the elastic ring 52 can be made of Nitinol (Ni-Ti alloy), stainless steel or other 5 satisfying materials.

The exemplary use of the positioning device is similar throughout the various embodiments described.

Fig. 13-16 illustrate a possible variant embodiment of the positioning device and the proximal and distal 10 components thereof according to the present invention. The elements in common have been designated with the same numeral used in the figures above and will be described below with reference to the differences from the above embodiments.

15 The proximal component 26 has a prismatic outer structure, preferably having a rectangular base. The cavity 32 is formed at one of the bases of the structure and does not run through the entire length of the proximal component 26, leaving a base wall 34. The sizes 20 of the cavity 32 and proximal component 26 are such as to leave a peripheral flat surface 26a.

In the base wall 34, preferably in the middle thereof, there is provided a preferably cylindrical cavity 38 extending along the longitudinal axis 30 and 25 crossing the entire thickness of the base wall.

Ribs 56, preferably on opposite parts of cavity 38 and parallel to the long sides of the rectangular base, extend from the bottom of the cavity 32 by a height preferably less than the depth of cavity 32.

5 According to a possible embodiment, the proximal component 26 comprises holes 40 for example for a suture, which can be used for separating the proximal component from the distal component, to be passed therethrough.

10 The distal component 28 comprises a head 42, which according to a possible embodiment, comprises holes (not illustrated) for a suture, which can be used for example to separate the distal component from the proximal component, to be passed therethrough.

15 The channel 50 extends along the longitudinal axis 30 throughout the solid thickness of the head 42, preferably in the middle thereof, and is suitable to receive a guide wire therein, not illustrated in Fig. 14 or 15.

20 The head 42 has a tract having a substantially pyramidal or having a truncated-pyramid shape, preferably with a rectangular base. Two flanges 54 that preferably involve the short sides of the rectangular base and a limited portion of the long sides extend from  
25 the outer periphery of the major base of the truncated-

pyramidal portion, in a direction substantially parallel to the longitudinal axis 30.

According to a possible embodiment, there is provided a preferably flat extension 58, arranged at a middle portion of each long side of the rectangular base and projecting in the direction substantially parallel to the longitudinal axis 30 along a preferably longer tract than the flanges 54.

Fig. 13 and 14 illustrate the positioning device 24 when assembled, in which the channel 50 and the cavity 38 are arranged along the longitudinal axis 30 to receive a guide wire, not illustrated in the Fig. 13-16. In the assembled configuration of the positioning device 24, the elastic ring 52 is held between the proximal component 26 and the distal component 28 in a preferably flattened, compressed/deformed configuration. After the elastic ring has been positioned, it takes a preset, uncompressed rest shape as described above. In the deformed configuration, the proximal end 52a of the elastic ring 52 is fastened by the inner periphery of the cavity 32. Particularly, the ribs 56 fasten the elastic ring 52 in a flat deformed configuration, or in other words, the elastic ring 52 is arranged between the wall of the cavity 32 and the ribs 56. Furthermore, the distal end 52b of the elastic ring 52 is fastened by the

flanges 54 and extensions 58, when the latter are provided.

On the one hand, the assembled configuration of the positioning device 24 is ensured by the interference 5 between the proximal end 52a of the elastic ring 52 and the walls of the proximal component 26 defining the cavity 32, and on the other hand by the interference between the distal end 52b of the elastic ring 52 and the flanges 54 and the extensions 58, when the latter 10 are provided.

Also in this case, the elastic ring 52 can be made of Nitinol, stainless steel or other satisfying materials.

The exemplary use of the positioning device is 15 similar throughout the various embodiments described. In this latter case, the peripheral wall 26a of the proximal component 26 is the one intended to abut against the wall of the tissue to be drawn together while minimizing the risk that the wall may be 20 penetrated. Furthermore, the angled head 42 of the distal component 28 is intended to penetrate the proximal and distal enterostomies and protects the elastic ring 52 when being introduced and positioned by fastening the ring within the flanges 54 and the 25 extensions 58, such as will be described in the

following.

The channel 50 and the cavity 38 are intended to house a guide wire for transporting the positioning device.

5       The present invention further relates to a method for the therapy of obesity and particularly a method for carrying out anastomosis in tracts of the digestive tube. Fig. 17-40 illustrate several steps of a possible embodiment of the method according to the present  
10 invention. The examples illustrated particularly relate to a method for carrying out an endoluminal/transluminal gastrojejunostomy (G-J) and a jejunojejunostomy (J-J) via transoral access.

In general terms, the method according to the  
15 present invention advantageously provides to draw tissues together and carry out anastomosis via endoluminal access by introducing, through a natural orifice (such as nose, mouth, ears, anus) or other luminal structures, guide or rail means within the  
20 tissues to be drawn together. Suitable components or devices can be thereby carried to the anastomotic site such that the surfaces of the tissues are suitably drawn together and connected with a channel (anastomosis).

Advantageously, the guide or rail means,  
25 particularly a main guide wire or first guide wire, are

introduced such as to generate an open ring that can begin and end in natural orifices, such as the mouth, nose, anus or other natural orifices, such as colostomy, trocar, abdomen incisions, wounds, fistulae. The 5 components or devices provided to draw the tissues together are advantageously moved by locking the device on the guide wire and pulling one of the guide wire ends.

In accordance with a possible embodiment, the ends 10 of the open ring and accordingly of the main guide wire are different from each other, and hence distinguishable. Advantageously, the guide wire is internally hollow, i.e. it has a tubular structure suitable to receive needles for perforating the tissues 15 and carrying out proximal and distal enterostomies. Perforation can take place for example either by pushing the needle through the tissues, or applying a radiofrequency through the needle.

The open ring then crosses the proximal enterostomy 20 and then the distal enterostomy, for example by using a gripping device.

The positioning device 24 is locked on the guide wire by means of an anchoring ring 76 and drawn by the guide wire until it is partially inserted in the 25 proximal enterostomy and abutted against a first tissue

portion to be joined. The positioning device 24 is further drawn until it is partially inserted in a distal enterostomy by drawing together the tissue portions to be joined. Finally, the positioning device 24 partially 5 inserted in the proximal and distal enterostomy releases an elastic ring 52 riding the proximal and distal enterostomies to hold the tissue portions joined to each other thereby generating a passage or anastomosis.

With reference to the above example, Fig. 17-29  
10 illustrate a gastrojejunostomy (G-J) step that is advantageously carried out by introducing guide or rail means through a natural orifice (such as the nose or mouth). Subsequently, the guide means, a guide wire in this case, form an open ring crossing the points of the  
15 tissues to be joined.

Fig. 17 illustrates a first step, which is designated as the step 1, in which a substantially conventional laparoscope 60 has been introduced in the abdominal cavity to view the areas to be treated. This  
20 step can be potentially eliminated after a certain degree of skill in the method has been achieved, thereby the method can be made completely endoluminal and transluminal. The laparoscope 60 is illustrated in Fig.  
17 and also in the subsequent steps, but it can be  
25 omitted as well. Alternatively or in addition to the

laparoscopic control, a gastroscopic control can be provided, i.e. by introducing a secondary gastroscope for example through the esophagus or mouth having the function of controlling the method steps. In case a 5 gastroscope is required to carry out several steps of the method, a main gastroscope carrying out the method steps and a secondary gastroscope monitoring the operation will be introduced.

Fig. 18 illustrates a step of the method according 10 to the present invention, which is also designated as the step 2, in which a substantially conventional main gastroscope 62 is introduced through the esophagus, stomach, passing through the pylorus and subsequently the duodenum to reach the jejunum. Particularly, the 15 gastroscope 62 is advanced by approximatively 20-40 cm beyond the pylorus.

Fig. 19 illustrates a detail of the jejunum and the end of the gastroscope 62. The latter conventionally comprises several channels 64 crossing the entire length 20 thereof and that can be used for tools or the like to be passed therethrough. The step from Fig. 19, also designated as the step 3, provides that a first guide wire 66 or main guide wire being suitable to provide the open ring is advanced along one of the channels 64 of 25 the gastroscope 62. The guide wire is advanced until a

pointed end 66a thereof or a needle sliding along the tubular structure of the guide wire protrudes from the gastroscope. The end 66a of the guide wire 66 perforates the jejunum wall from the inside and creates a 5 jejunoostomy (proximal enterostomy). The laparoscope 60 is optionally provided. When this is provided, the guide wire 66 is advanced and the jejunoostomy is created under the laparoscope visual control.

The jejunoostomy can be carried out by pushing the 10 guide wire directly through the jejunum wall. Alternatively, or in addition thereto, radiofrequency energy may be applied to perforate the jejunum wall and then advance the guide wire 66.

In other words, a first guide wire 66 being part of 15 guide or rail means which will be subsequently indicated in greater detail, is positioned within the tissue to be joined and passed through one of the tissue portions to be joined. The jejunum tissue portion to be drawn near and joined to the stomach thereby forming an anastomosis 20 has been designated with A.

Fig. 20 illustrates a step which has been designated as the step 4 in which the gastroscope 62 is removed and the guide wire 66 is left in the abdomen within the stomach and along a jejunum tract with the 25 end 66a protruding from the jejunum at the tissue

portion A to be joined. The step 4 can be carried out under laparoscopic control (laparoscope 60), when provided.

Fig. 21 illustrates a step which is designated as 5 the step 5, in which a main gastroscope 62, of a substantially conventional type, has been introduced again in the stomach through the esophagus, in order to create a gastrostomy (distal enterostomy). Also in this case, one may directly push either a secondary guide wire 77 with a pointed end 77a or a needle sliding 10 within the tubular structure of the guide wire. Alternatively, or in addition thereto, radiofrequency energy can be applied in order to perforate the stomach wall and advance the guide wire.

15 The gastrostomy is carried out in a stomach portion corresponding to the area to be joined. This portion has been designated with A'.

Step 5 may also be carried out under laparoscopic control.

20 Fig. 22 illustrates a step of the method according to the present invention, which has been designated as the step 6, in which the gastrostomy is enlarged by means of a balloon catheter 72. The catheter is inserted 25 in the gastroscope 62 until a balloon-end 72a thereof reaches the gastrostomy which is enlarged by inflating

the balloon.

Step 6 may also be carried out under laparoscopic control.

Fig. 23 illustrates a step of the method according 5 to the present invention, which has been also designated as the step 7, in which the gastroscope 62 is advanced through the gastrostomy enlarged by the balloon, within the abdominal cavity. Step 7 can be carried out either under gastroscopic (secondary gastroscope) and/or 10 laparoscopic (laparoscope 60) control. As stated above, by gastroscopic control is meant a control carried out by means of a secondary gastroscope, not illustrated in Fig. 23, which is introduced through the esophagus having only a control function. This secondary 15 gastroscope can be provided in every step whenever a gastroscopic control as an alternative or in addition to the laparoscopic control is required.

Fig. 24 illustrates a step which has been designated as the step 8, in which a gripping device 74 20 (forceps or the like) is advanced through the gastroscope 62 and the end 66a of the guide wire 66 protruding from the jejunostomy is coupled therethrough. Gripping the guide wire point is not required.

The gripping device 74 can be for example a loop-shaped endoscopic instrument for polypectomies. 25

Fig. 25 illustrates a step which has been designated as the step 29, in which the guide wire 66 of the jejunum is pulled through the gastrostomy in order to provide a first ring 80 open at the ends thereof, or 5 gastro-jejunum ring (ring 1), the ends thereof protruding from the orifice used (the mouth, esophagus, ...). In Fig. 25, the end of ring 80 corresponding to the jejunum (jejunum end), i.e. the end passing through the stomach and jejunum and protruding from portion A has 10 been designated with 80a, whereas the end of the ring 80 corresponding to the stomach (stomach end), i.e. the end passing through the stomach and protruding therefrom at portion A' has been designated with 80b. Both ends are advantageously different from each other in order to be 15 distinguished.

The ring 80 can be now used as a guide means or rail system in order to introduce and carry suitable anastomotic devices suitable to draw the tissues together and carry out the anastomosis in the site of 20 interest. The anastomotic devices are advantageously locked on the guide wire for example by means of an anchoring ring 76 and one of the ends of the ring is pulled until the anastomotic device partially enters the proximal enterostomy, draws the tissues thereof near the 25 distal enterostomy and partially enters the same.

Fig. 26 illustrates a step which has been designated as the step 10, wherein the selected anastomotic device (anvil 16, positioning device 24, etc.) is inserted on the guide wire from the jejunum end 5 80a and pulled along the ring 80 of guide wire through the esophagus, stomach, duodenum and jejunum. Drawing is allowed by an anchoring ring 76 which is made integral with the guide wire such as to push against the proximal part of the selected anastomotic device.

10 Though an anvil 16 has been illustrated in Fig. 26, a positioning device 24 or other similar devices can be used as well.

By pulling the guide wire from the end of stomach 80b, the anastomotic device can be pulled until the 15 portion A of the jejunum (proximal enterostomy).

As illustrated in Fig. 26, the end of the jejunum part of the guide wire 80a is inserted in the channel 22 of the anvil 16 from the side of the stem 18. In the case of the positioning device 24, the end of the 20 jejunum part of the guide wire 80a would be inserted in the channel 50 from the side of the distal component 28.

Fig. 27 illustrates a step which has been also designated as the step 11, in which the anastomotic device and particularly the anvil 16 is pulled until it 25 has partially passed the jejunostomy (proximal

enterostomy). As already stated above, one can act under laparoscopic control. The stem 18 of the anvil 16 crosses the jejunostomy and protrudes in the abdominal cavity, whereas the head 20 contacts the tissue to be drawn together. When a positioning device 24 is used, the head would enter the jejunostomy whereas the flat peripheral surface 26a would abut against the contouring tissues.

Fig. 28 illustrates a step which has been also designated as the step 12, in which by keeping on pulling the end of the guide wire 80b from the side of the stomach, the anvil 16 and particularly the head 20 acts as a striker against the inner wall of the portion A of the jejunum and draws the jejunum until the portion A' is drawn near the stomach and particularly portion A'. The stem 18 of the anvil 16 (anastomotic device) also partially enters the gastrostomy (distal enterostomy). The operation can be carried out under laparoscopic control (laparoscope 60). When a positioning device 24 is used, the head would enter the gastrostomy whereas the flat peripheral surface 26a would draw the relative contouring tissues together.

Fig. 29 illustrates a step which has been indicated as the step 13, in which the traction on the stomach end 80b of the guide wire is maintained in order to maintain

the portions A and A' near each other. Furthermore, a circular stapler 10 is caused to slide on the guide wire from the stomach end 80b until it reaches the inside of the stomach, and until the stem 18 of the anvil 16 5 connects to the end of the stem 14 of the stapler 10 (such as illustrated in greater detail in Fig. 2a and 41-44). The stapler 10 carries out the anastomosis between the portion A and the portion A' by cutting and suturing the tissue in a circular manner. A passage 84 10 is thereby formed (Fig. 30) which directly communicates the stomach and jejunum. When a positioning device 24 is used, the passage 84 is obtained by detaching the proximal component from the distal component and releasing the elastic ring 52 riding both enterostomies.

15       When the gastrojejunostomy (G-J) has been completed, the guide wire 66 is removed by drawing one end thereof.

With reference to the above example, Fig. 30-40 illustrate a jejunojejunostomy (J-J) step that is 20 advantageously carried out by introducing guide or rail means through a natural orifice (such as the esophagus or mouth). Subsequently, the guide means, a guide wire in this case, form an open ring crossing the points of the tissues to be joined.

25       Fig. 30 illustrates a step designated as the step

14, in which a substantially conventional gastroscope 62 is introduced through the esophagus, stomach, passing through the pylorus and subsequently the duodenum to reach the jejunum. Particularly, the gastroscope 62 is 5 advanced to a portion to be joined that is designated with B in order to carry out a proximal jejunostomy, which portion is proximally arranged relative to channel 84 (anastomosis) that has already been created.

A guide wire 86, or main guide wire, intended to 10 form the open ring is advanced along a channel 64. The guide wire is advanced until a pointed end 86a thereof or a needle sliding within the guide wire protrudes from the gastroscope. The end 86a of the guide wire 66 perforates the jejunum wall from the inside and creates 15 a jejunostomy (proximal enterostomy).

The laparoscope 60 is optionally provided. When this is provided, the guide wire 86 is advanced and the jejunostomy is created under the laparoscope visual control.

20 The jejunostomy can be carried out by pushing the guide wire directly through the jejunum wall. Alternatively, or in addition thereto, radiofrequency energy may be applied to perforate the jejunum wall and then advance the guide wire 86.

25 In other words, a guide wire 86 being part of guide

or rail means which will be subsequently indicated in greater detail, is positioned within the tissue to be joined and passed through one of the tissue portions B to be joined (proximal jejunostomy).

5 Fig. 31 illustrates a step which has been indicated as the step 15, in which the gastroscope 62 is removed and the guide wire 86 is left within the stomach and jejunum, with the end 86a protruding from the walls of the jejunum (proximal jejunostomy). The gastroscope 62  
10 is then advanced through the stomach, the previously accomplished gastrojejunostomy (step 84) and a tract of the distal jejunum until a sufficient distance to create the jejunumjejunum (J-J) anastomosis. The latter portion has been indicated with the reference B'.  
15 Analogously to Fig. 21 and 22 (steps 5 and 6) a secondary guide wire is advanced along the gastroscope 62 until a pointed end thereof protrudes from the end of the gastroscope 62. This pointed end is then passed through the tissue walls at the portion B' in order to  
20 create a distal jejunostomy. The distal jejunostomy can be also carried out either by directly pushing the pointed end through the jejunum wall or applying radiofrequency energy in order to perforate the wall, subsequently advancing the guide wire.

25 The creation of the distal jejunostomy can be

monitored through the laparoscope.

A catheter with balloon-end may be optionally inserted along the gastroscope. When the balloon end is near the distal jejunostomy, the balloon is inflated in 5 order to dilate the distal jejunostomy and the gastroscope is pushed in the abdominal cavity. This dilation may be required when the laparoscope is not used and monitoring is carried out through the gastroscope such that the latter can view the end 86a of 10 the guide wire 86.

Fig. 32 illustrates a step which has been designated as the step 16, in which a gripping device 74 (endoscopic forceps or the like) similar to the one used in step 8, is advanced through the gastroscope to lock 15 the end 86a of the main guide wire 86 protruding from the proximal jejunostomy site. Gripping the guide wire end is not required.

Fig. 33 illustrates a step which has been indicated as the step 17, in which the gastroscope has been removed and the main guide wire 86 has been pulled 20 through the distal jejunostomy to form a ring 88 open at the ends thereof, or jejunumjejunum ring (the ring 2), the ends thereof protruding from the orifice used. In Fig. 33 the end of the jejunum, i.e. the end passing 25 through the stomach and the jejunum and protruding

therefrom at portion B has been designated with 88a, whereas the end of the stomach, i.e. the end passing through the stomach, protruding from the gastrojejunostomy (passage 84) and protruding from the 5 jejunum at portion B' has been designated with 88b.

The ring 88 can be now used as a guide means or rail system in order to introduce and carry suitable anastomotic devices suitable to draw the tissues together and carry out the anastomosis in the site of 10 interest (J-J). As described above, the anastomotic device is locked on the guide wire, an end thereof being pulled in order to advance the anastomotic device.

Fig. 34 illustrates a step which has been designated as the step 18, wherein an anastomotic device 15 such as a positioning device 24 is inserted from the jejunum end 88a and pulled along the ring 88 of guide wire through the esophagus, stomach, duodenum and jejunum. The operation may be carried out under laparoscopic and/or gastroscopic control in order to 20 view the movement.

The anastomotic device may be caused to slide on the guide wire from the jejunum end 88a. Traction is permitted due to an anchoring ring similar to that described above, which is made integral with the guide 25 wire pushing against the proximal part of the selected

anastomotic device. By pulling the guide wire from the end of stomach 88b, the anastomotic device can be pulled until the portion B of the jejunum.

As illustrated in Fig. 34, the positioning device 5 24 (channel 50 and cavity 38) is inserted on the end 88a of the guide wire from the side of the distal component 28.

The anastomotic device and particularly the positioning device 24 is pulled until it has partially 10 passed the proximal jejunostomy. The head 42 of the distal component 28 and a part of the elastic ring 52 cross the proximal jejunostomy and protrude in the abdominal cavity, whereas the other part of the elastic ring 52 remains within the jejunum. The proximal 15 component also remains within the jejunum and abuts, for example with the surface 26a, against the tissue wall to act as a striker.

Fig. 35 illustrates a step which has been designated as the step 19, in which the two jejunum 20 branches are drawn together, optionally under gastroscopic and/or laparoscopic vision by keeping on pulling the anastomotic device (positioning device 24).

Particularly, the head 42 of the distal component 28 with a part of the elastic ring 52 penetrate in the 25 distal jejunostomy.

Fig. 36 illustrates a step which has been designated as the step 20, in which the elastic ring 52 that takes its uncompressed configuration, such as illustrated enlarged for example in Fig. 37 (step 21) is positioned. The ends of the elastic ring 52 fold between the proximal jejunostomy and the distal jejunostomy thereby maintaining the portion B and portion B' joined to each other thereby creating a circular anastomosis. The elastic ring 52 can be released from the positioning device 24 by simultaneously uncoupling the distal and proximal components 26 and 28. Alternatively, one of the two components can be uncoupled by pulling the same while keeping unchanged the position of the elastic ring 52 relative to the anastomotic site.

To uncouple the distal component and the proximal component, one can use the suture threads protruding from the holes 40 of the proximal component 26 and the holes 46 of the distal component 28 by coupling them by means of a suitable tool inserted in a gastroscope 62. The suture stitches are thus gripping points for uncoupling the proximal and distal components of the positioning device 24 from each other.

During positioning, the enterostomy can be viewed by means of a gastroscope or laparoscope.

When the jejunojejunostomy (J-J) has been

completed, the guide wire 86 is removed by drawing one end thereof.

Fig. 38 illustrates a step which has been designated as the step 22, in which the 5 gastrojejunostomy (G-J) and the jejunojejunostomy (J-J) have been completed and in which there is illustrated the route followed by the food along the digestive tract after it has been changed.

To complete the method discussed above, either a 10 gastric partition obtained with a gastric bandage such as illustrated in Fig. 39 (step 23.1) or a gastric partition obtained with an endoscopic stapler such as illustrated in Fig. 40 (step 23.2) can be provided.

From what has been described above, one may 15 appreciate how the provision of guide means carrying components or devices to the desired anastomotic site through natural orifices (such as the nose, mouth, ear, anus) or other luminal structures in order to carry out anastomosis greatly simplifies the procedure, shortens 20 the patient's convalescence and eliminates the drawbacks of traditional surgery.

The provision of components and devices that suitably draw the tissue surfaces together and/or connect the surfaces by means of a passage is 25 particularly advantageous and allows to carry out a

completely endoluminal method.

It should be understood that variations and/or additions to what has been described and illustrated above may be provided.

5 In addition to the method described above, there may provided alternative procedures (ERCP, Chole duct, colo-proctostomy, jejunum-colostomy).

The order of the steps of the method illustrated in the annexed drawings and described above,  
10 (gastrojejunostomy or G-J, jejunojejunostomy or J-J, sectioning) can be readapted. For example, with patients that have already been subjected to gastric bandage, the steps G-J and J-J can be completed as described above. Subsequently, the gastric bandage can  
15 be completely restricted by carrying out a gastric partition thereby completing the procedure.

Alternatively to the use of the circular stapler 10 and anvil 16, the gastrojejunostomy G-J according to the steps described above (Fig. 17-Fig. 29) can be carried  
20 out by means of a positioning device 24 as described above (similarly to the jejunojejunostomy J-J steps corresponding to Fig. 31-38).

To the preferred embodiment of the device, stapler or method described above, those skilled in the art,  
25 aiming at satisfying contingent and specific

requirements, may carry out a number of modifications, adaptations and replacement of elements with others functionally equivalent, without however departing from the scope of the claims below.

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**CLAIMS**

1. A device for drawing tissues together and creating an anastomosis comprising a main guide wire (66, 86) suitable to receive and draw anastomotic devices (10; 16; 24) to draw together portions (A; B) of tissues to be joined by means of anastomosis.
2. The device for drawing tissues together and creating an anastomosis according to claim 1, wherein said main guide wire (66, 86) is suitable to be shaped as an open ring crossing the portions (A, A'; B, B') of tissues to be joined by means of anastomosis.
3. The device for drawing tissues together and creating an anastomosis according to claim 1 or 2, wherein the ends of the main guide wire (66, 86) are different and distinguishable from each other.
4. The device for drawing tissues together and creating an anastomosis according to one of the preceding claims, wherein said main guide wire (66, 86) has a tubular or hollow structure that is suitable to house at least one needle for the push or radiofrequency perforation of the tissues.
5. The device for drawing tissues together and creating an anastomosis according to one of the preceding claims, comprising an anchoring ring (76) suitable to be made integral with said main guide wire

(66, 86) in order to be abutted against said anastomotic device (16, 24) and drawing the same by drawing an end of the main guide wire (66, 86).

6. The device for drawing tissues together and  
5 creating an anastomosis according to one of the preceding claims, wherein said anastomotic device is a positioning device (24) comprising an elastic ring (52), said positioning device (24) comprising a channel (50) where this main guide wire (66, 86) has to be inserted.

10 7. The device for drawing tissues together and creating an anastomosis according to one of claims 1 to 5, wherein said anastomotic device is a circular stapler (10) comprising an anvil (16), said circular stapler (10) and said anvil (16) comprising a channel where this  
15 main guide wire (66, 86) has to be inserted.

8. A method for carrying out anastomosis in tracts of the digestive tube comprising the steps of:

- introducing a guide or rail means within the tissues to be drawn near and joined, through a natural  
20 orifice or other luminal structures,

- inserting an anastomotic device (10, 16; 24) along the guide means and drawing the same until the anastomotic site, by drawing near portions (A, A'; B, B') of the tissues to be joined and carrying out an  
25 anastomosis.

9. The method according to claim 8, wherein the step of introducing the guide or rail means, particularly a main guide wire (66, 86), is such as to generate an open ring (80, 88) with the ends matching 5 natural orifices or other orifices and being such as to cross the portions (A, A'; B, B') of the tissues to be joined.

10. The method according to claim 9, wherein the main ring-shaped guide wire (66, 86) crosses a proximal 10 enterostomy and a distal enterostomy, said enterostomies being obtained by causing a needle to slide within the tubular structure of a guide wire, said needle perforating the tissue by pushing or via radiofrequency.

11. The method according to claim 10, wherein the 15 main guide wire (66, 86) suitable to be ring-shaped is introduced until matching the first portion (A, B) of the tissue to be joined and inserted through a proximal enterostomy that is carried out by causing a needle to slide within the tubular structure of said main guide 20 wire, said needle perforating the tissue by pushing or via radiofrequency.

12. The method according to claim 11, wherein a secondary guide wire (77) is introduced until matching the second portion (A', B') of the tissue to be joined 25 in order to carry out a distal enterostomy by causing a

needle to slide within the tubular structure of said secondary guide wire, said needle perforating the tissue by pushing or via radiofrequency.

13. The method according to claim 12, wherein a gripping device is introduced through said distal enterostomy to grip the end of the main guide wire (66, 86) suitable to be ring-shaped and draw the same to obtain said open ring.

14. The method according to one of claims 9 to 13, wherein at least one portion (16, 24) of the anastomotic device is inserted and locked on the main guide wire defining the open ring and drawn by pulling one of the free ends of the open ring.

15. The method according to claim 14, wherein the anvil (16) is locked on the guide means by means of an anchoring ring (76).

16. The method according to one of claims 8 to 15, wherein said portion (16, 24) of the anastomotic device is drawn until it is partially introduced in a proximal enterostomy thereby abutting against one first portion to be joined (A, B).

17. The method according to claim 16, wherein said portion (16, 24) of the anastomotic device is further drawn until it is partially introduced in a distal enterostomy by drawing near the portions (A, A'; B, B')

of the tissues to be joined.

18. The method according to claim 17, wherein said anastomotic device (10, 16; 24) partially introduced in the proximal and distal enterostomies joins the portions 5 (A, A'; B, B') of the tissues thereby generating a passage or anastomosis.

19. The method according to one of claims 8 to 18, comprising a step of carrying out a gastrojejunostomy (G-J) comprising the steps of:

10 - introducing the guide or rail means, particularly a main guide wire (66) through a natural orifice or other orifice within the stomach and jejunum, and

15 - shaping the guide means in a first open ring (80) with ends (80a, 80b) leading to said orifice, said first ring crossing the portions (A, A') to be joined, respectively a jejunostomy and a gastrostomy.

20. The method according to claim 19, wherein a main guide wire (66) is introduced through the esophagus, stomach and a tract of the jejunum and an end thereof (66a) is passed through the jejunum wall in order to form a jejunostomy at the portion (A) of the tissue to be joined.

21. The method according to claim 20, wherein the jejunostomy is carried out by causing a needle to slide 25 within the tubular structure of the main guide wire,

said needle perforating the tissue by pushing or via radiofrequency.

22. The method according to claim 20 or 21, wherein said main guide wire (66) is introduced through 5 a main gastroscope (62) that has been previously introduced through the esophagus, stomach and jejunum.

23. The method according to claim 22, wherein after said main gastroscope (62) has been introduced, it is removed and the main guide wire (66) is left within 10 the stomach and along a tract of the jejunum with an end thereof (66a) protruding from the jejunum at the tissue portion (A) to be joined.

24. The method according to one of claims 19 to 23, wherein a gastrostomy is created at a portion (A') 15 of the tissue of the stomach to be joined to the corresponding portion (A) of the jejunum.

25. The method according to claim 24, wherein said gastrostomy is created by a needle that is slidingly inserted in the tubular structure of a secondary guide 20 wire (77), said needle perforating the tissue by pushing or via radiofrequency.

26. The method according to claim 24 or 25, wherein said gastrostomy is enlarged, for example by using a balloon catheter (72).

25 27. The method according to one of claims 24 to

26, wherein a gripping device (74) is advanced through the gastrostomy in order to couple the end (66a) of the main guide wire (66) protruding from the jejunostomy.

28. The method according to one of claims 24 to 5 27, wherein the steps of carrying out said gastrostomy or introducing said balloon catheter (72) or introducing said gripping device (74) are carried out by means of a main gastroscope (62) that is introduced in the stomach until a portion (A') of the tissue to be joined and 10 optionally beyond gastrostomy.

29. The method according to claim 27 or 28, wherein said main guide wire (66) is pulled to form an open ring (80) crossing the portions (A, A') of the tissues to be joined and defines guide means for 15 introducing anastomotic devices (24) suitable to join the tissues and carry out the gastrojejunostomy (G-J).

30. The method according to one of claims 19 to 29, wherein an anastomotic device (10, 16; 24) is introduced on the main guide wire (66) and locked by an 20 anchoring ring (76) that is made integral with the guide wire, and wherein a free end of this guide wire (66) is pulled until the anastomotic device reaches both the jejunostomy and portion (A) of the jejunum tissue to be joined, by partially passing through the jejunostomy and 25 abutting against the inner walls at the jejunostomy.

31. The method according to claim 30, wherein said main guide wire (66) is further drawn until the anastomotic device draws the jejunum thereby drawing together the portion to be joined (A) of the jejunum to 5 the portion to be joined (A') of the stomach, said anastomotic device partially entering the gastrostomy.

32. The method according to claim 31, wherein the tissue portions (A, A') are joined thereby creating an anastomosis riding the portions (A, A') of the jejunum 10 and stomach that have been drawn together, respectively, whereby a passage (84) is formed which directly connects the stomach to the jejunum.

33. The method according to one of claims 7 to 32, comprising a step of carrying out a jejunojejunostomy 15 (J-J) comprising the steps of:

- introducing the guide or rail means, particularly a main guide wire (86) through a natural orifice or other orifice within the stomach and jejunum, and

- shaping the guide means in an open ring (88) with 20 ends (88a, 88b) leading to said orifice, said ring crossing the portions (B, B') to be joined, respectively a proximal jejunostomy and a distal jejunostomy.

34. The method according to claim 33, when depending on one of claims 19 to 32, wherein the step of 25 carrying out a jejunojejunostomy (J-J) is subsequent to

the gastrojejunostomy (G-J) whereby a passage (84) is formed which directly connects the stomach to the jejunum.

35. The method according to claim 34, wherein the  
5 main guide wire (86) is introduced through the esophagus, stomach and a tract of the jejunum and an end thereof (86a) is passed through the jejunum wall in order to form a proximal jejunostomy at a portion (B) of the tissue to be joined which is located between the  
10 stomach and the passage (84).

36. The method according to claim 35, wherein the proximal jejunostomy is carried out by causing a needle to slide within the tubular structure of the main guide wire, said needle perforating the tissue by pushing or  
15 via radiofrequency.

37. The method according to claim 35 or 36, wherein said main guide wire (86) is introduced through a main gastroscope (62) that has been previously introduced through the esophagus, stomach and jejunum.

20 38. The method according to claim 37, wherein after said main gastroscope (62) has been introduced, it is removed and the main guide wire (86) is left within the stomach and along a tract of the jejunum with an end thereof (86a) protruding from the jejunum at the tissue  
25 portion (B) to be joined.

39. The method according to one of claims 35 to 38, wherein a distal jejunostomy is created at a portion (B') of the tissue of the jejunum to be joined which is located beyond the passage (84).

5 40. The method according to claim 39, wherein said distal jejunostomy is created by a needle that is slidingly inserted in the tubular structure of a secondary guide wire (77), said needle perforating the tissue by pushing or via radiofrequency.

10 41. The method according to claim 39 or 40, wherein said distal jejunostomy is enlarged, for example by using a balloon catheter (72).

15 42. The method according to one of claims 39 to 41, wherein a gripping device (74) is advanced through the passage (84) and the distal gastrostomy in order to couple the end (86a) of the main guide wire (86) protruding from the proximal jejunostomy.

20 43. The method according to one of claims 39 to 42, wherein the steps of carrying out said gastrostomy or introducing said balloon catheter (72) or introducing said gripping device (74) are carried out by means of a main gastroscope (62) that is introduced in the stomach and jejunum through said passage (84) until a portion (B') of the tissue to be joined and optionally beyond 25 distal jejunostomy.

44. The method according to claim 42 or 43, wherein said main guide wire (86) is pulled to form said open ring (88) crossing the portions (B, B') of the tissues to be joined and defines the guide means for 5 introducing anastomotic devices (16, 10) suitable to draw the tissues together and carry out the jejunojejunostomy (J-J).

45. The method according to one of claims 34 to 44, wherein an anastomotic device is introduced on the 10 main guide wire (86) and locked by a anchoring ring (76) that is made integral with the guide wire, and wherein said main guide wire (86) is pulled until the device reaches both the proximal jejunostomy and portion (B) of 15 the tissue to be joined, by partially passing through the proximal jejunostomy and abutting against the inner walls at the jejunostomy.

46. The method according to claim 45, wherein said main guide wire (86) is further pulled until the anastomotic device draws the jejunum thereby drawing 20 near the portion to be joined (B) at the proximal jejunostomy to the portion to be joined (B') at the distal jejunostomy, said anastomotic device partially entering the distal jejunostomy.

47. The method according to claim 46, wherein the 25 anastomotic device joins the portions (B, B') of the

tissues thereby creating an anastomosis riding the portions (B, B') that are drawn near whereby a passage is formed which connects two portions of the jejunum.

48. The method according to claim 47, wherein a 5 gastric partition obtained by means of a gastric bandage applied by laparoscope is carried out after the anastomosis has been sutured.

49. The method according to claim 47, wherein a gastric partition obtained by means of an endoscopic 10 stapler is carried out after the anastomosis has been sutured.

50. The method according to any claim 8 to 49, wherein the use of a laparoscope (60) is provided for monitoring at least one of the steps of the method.

15 51. The method according to any claim 8 to 49, wherein the use of a secondary gastroscope is provided for monitoring at least one of the steps of the method.

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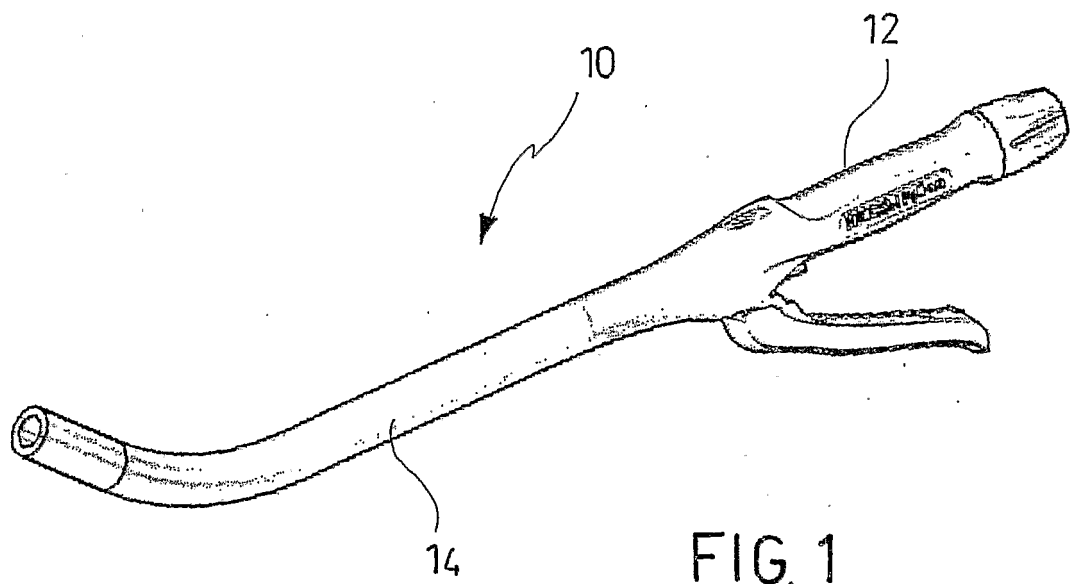


FIG. 1

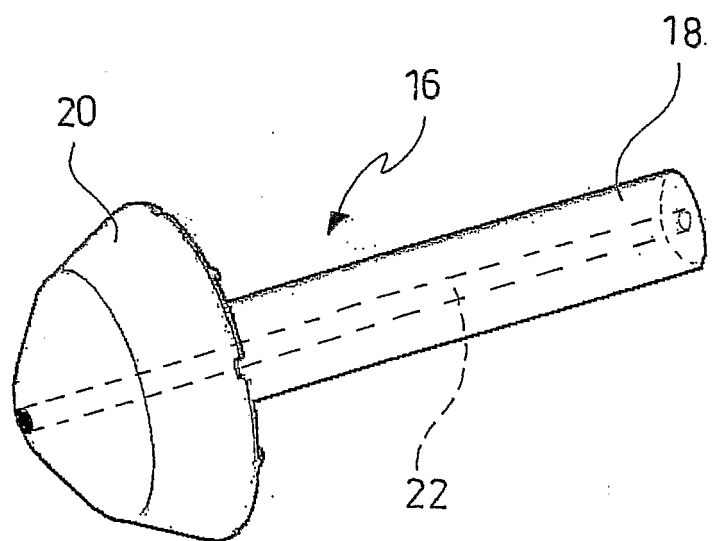
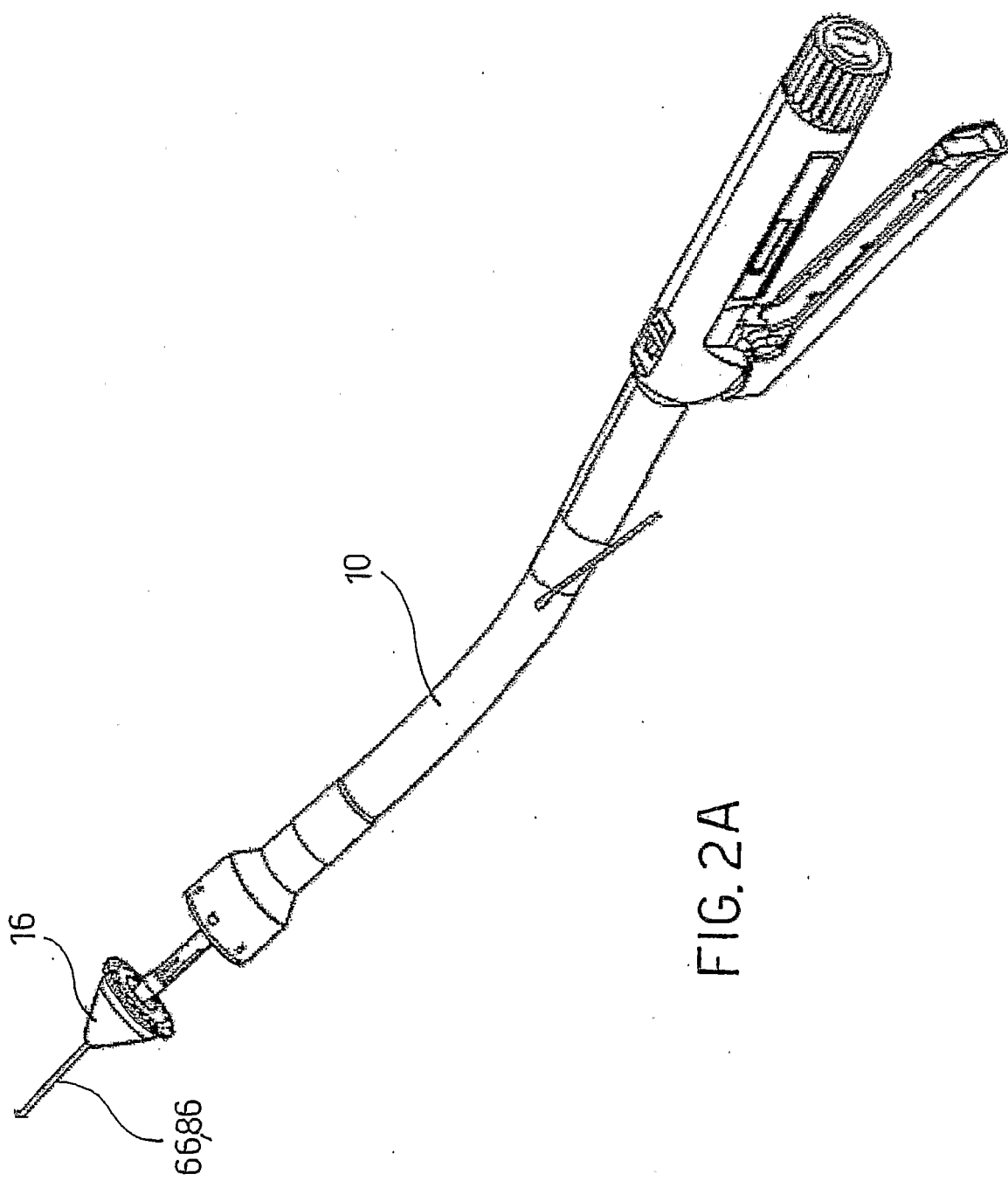


FIG. 2

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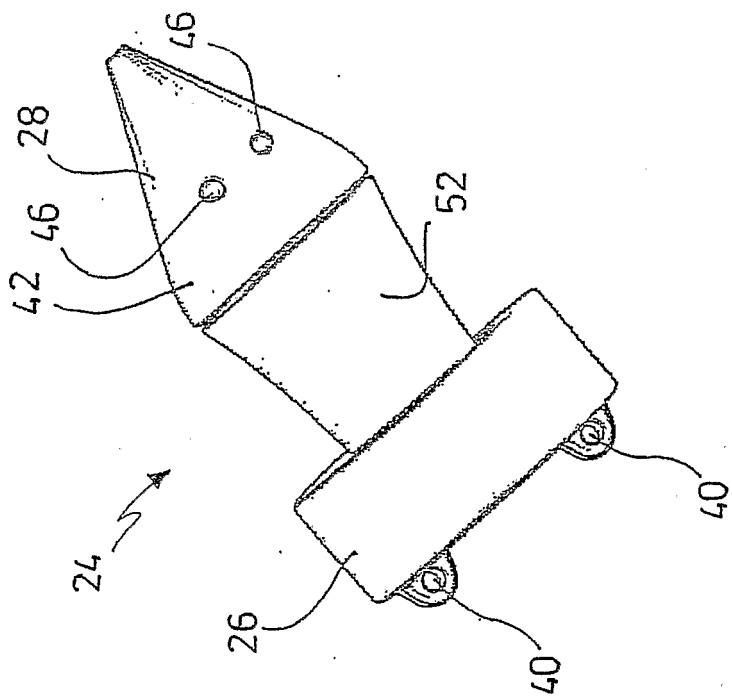


FIG. 3

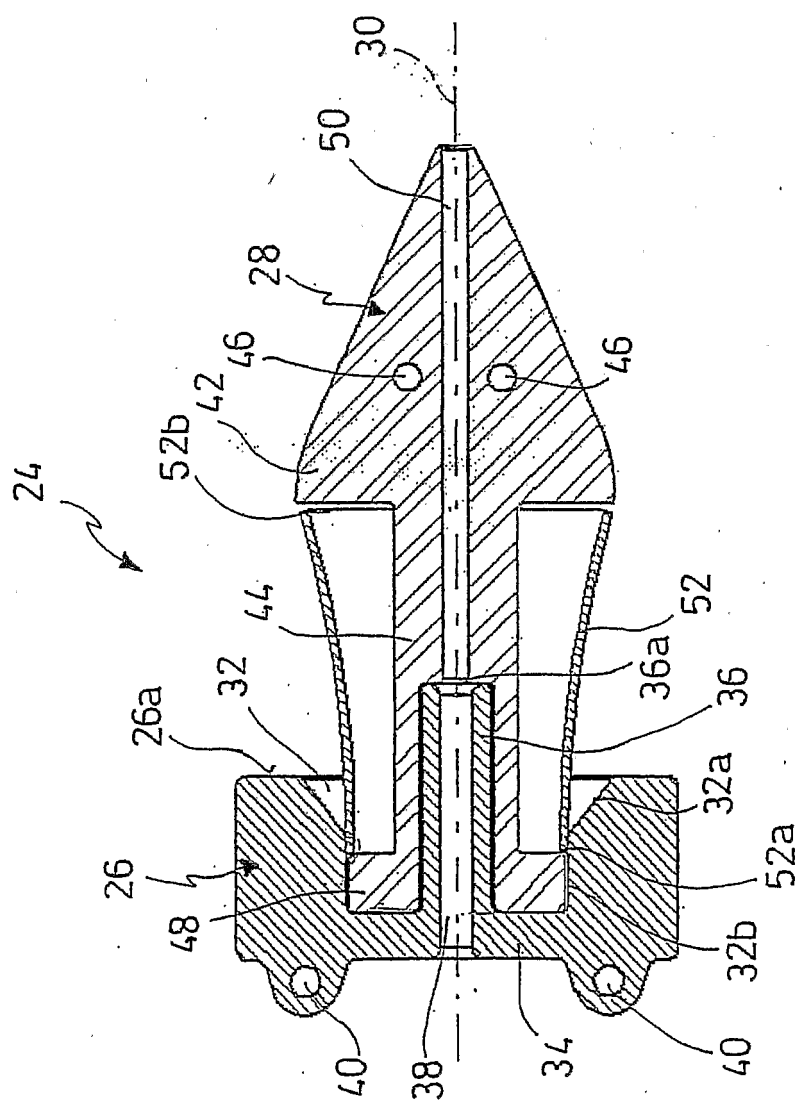
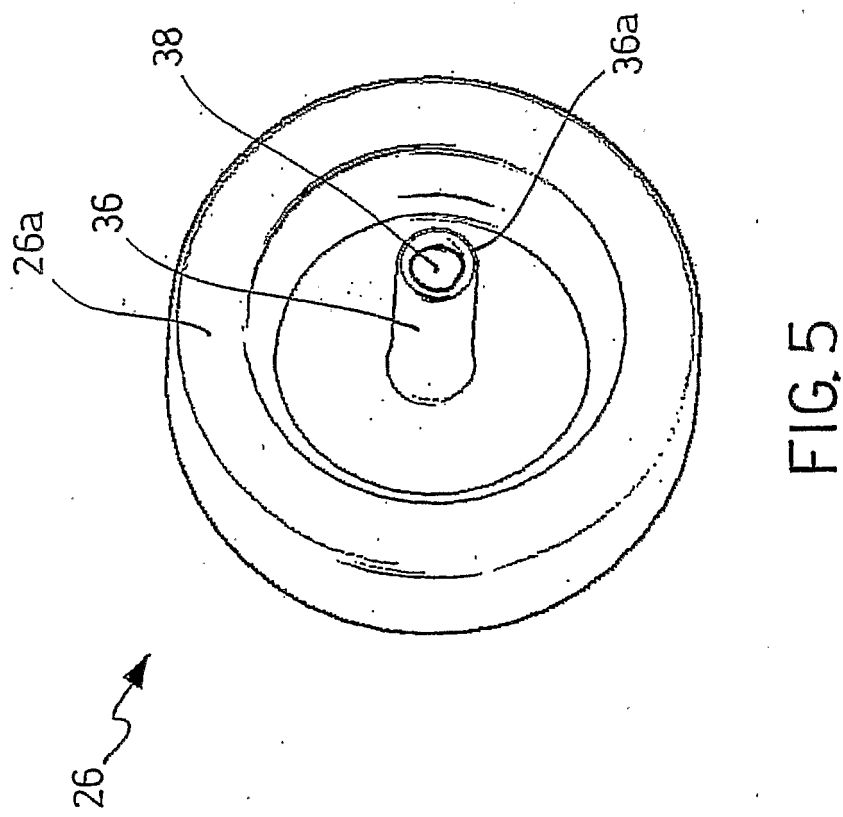
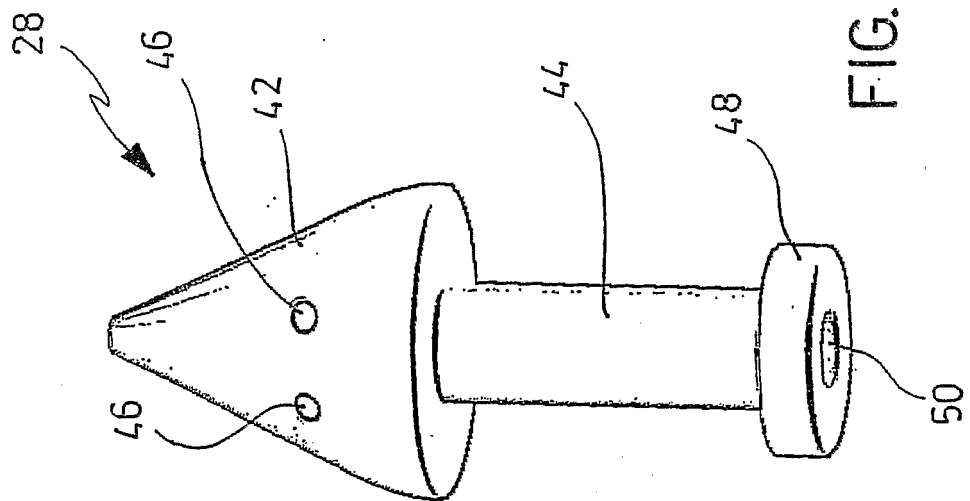
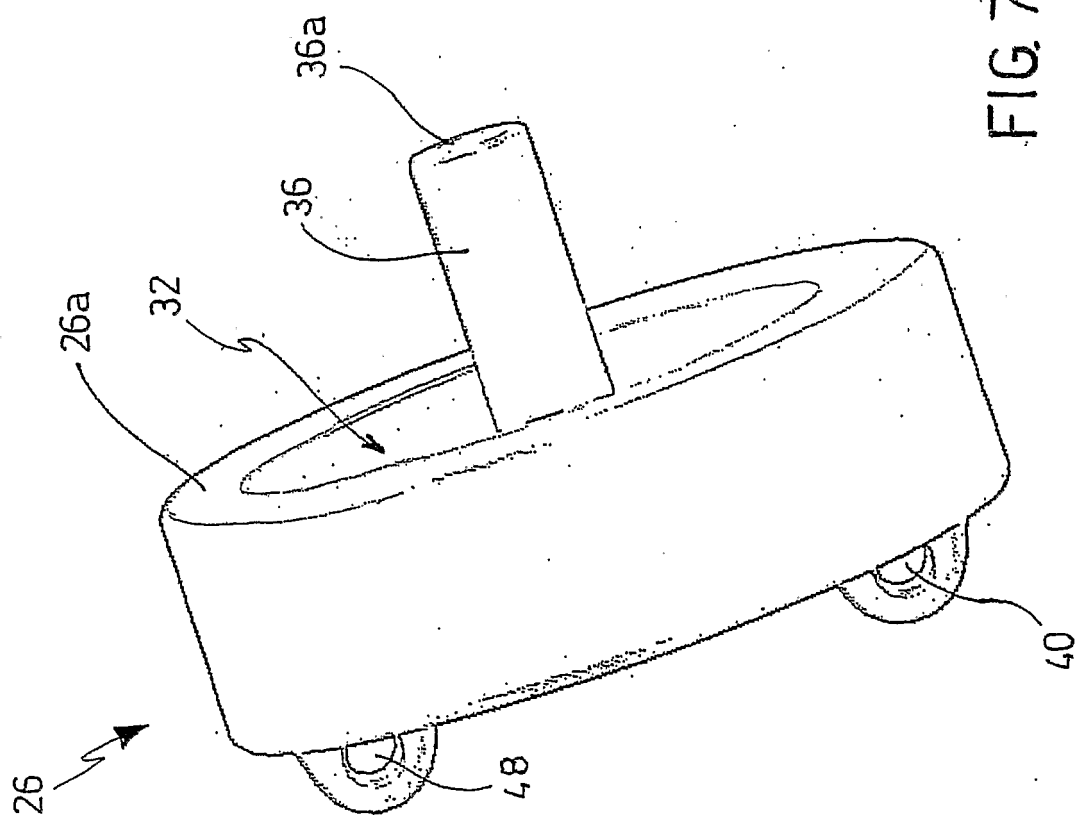
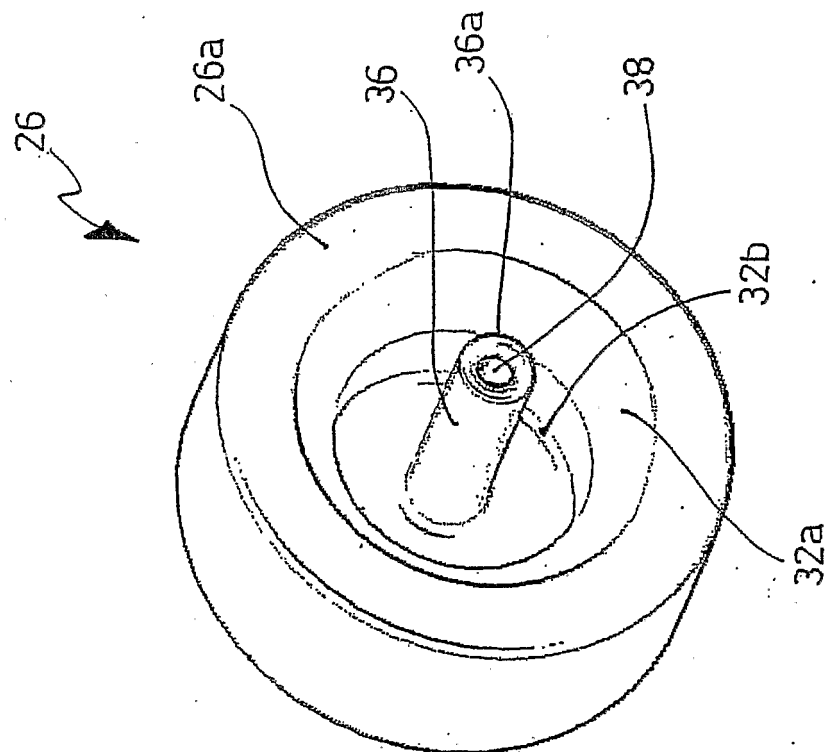


FIG. 4

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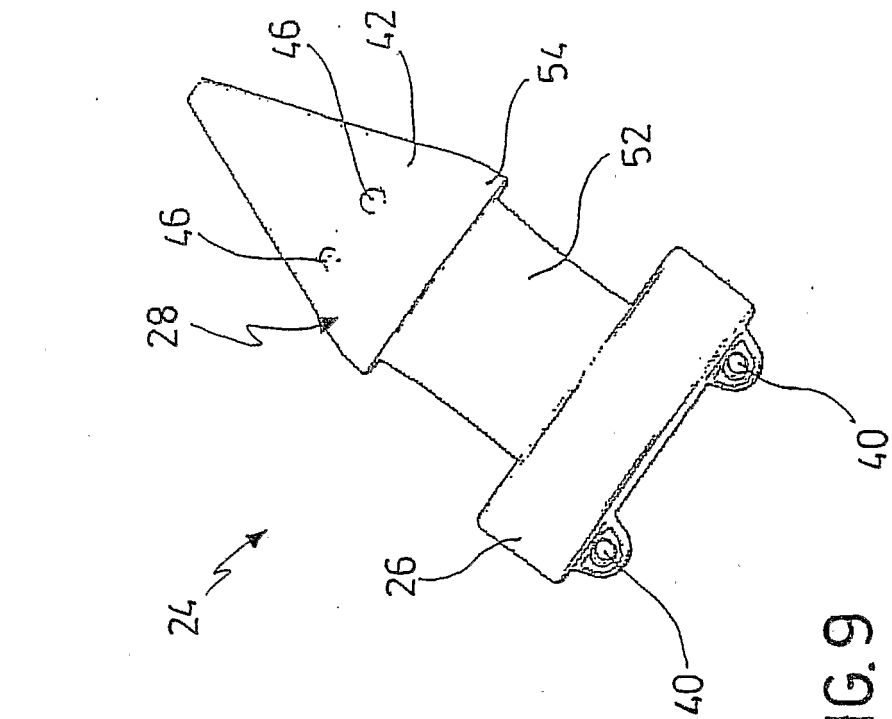


FIG. 9

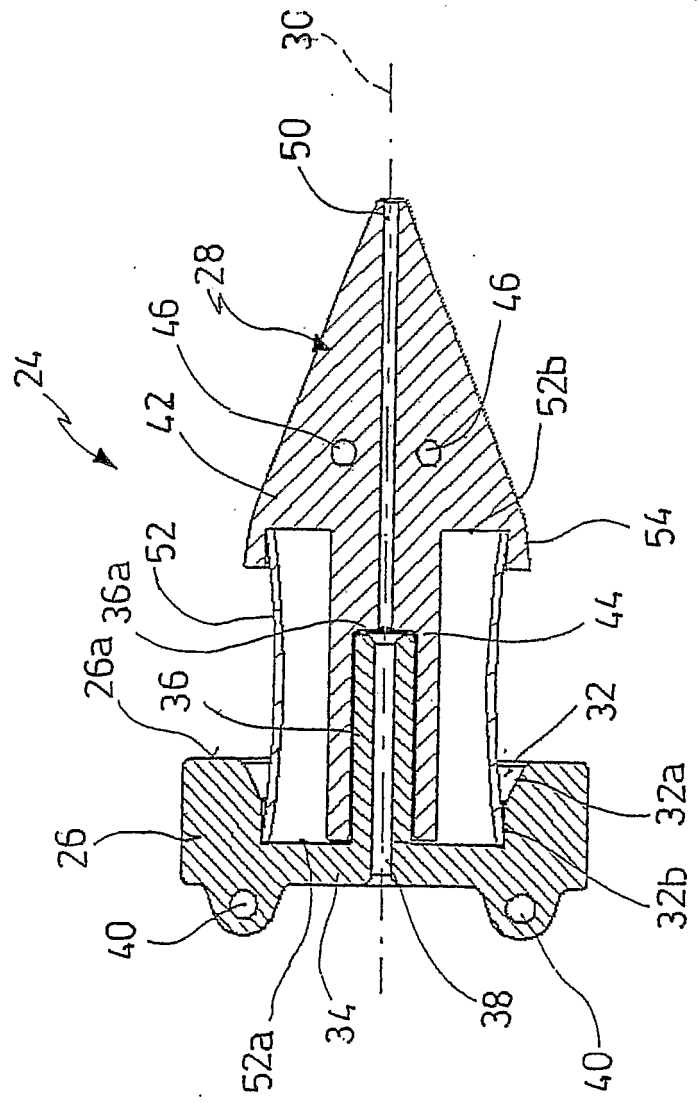


FIG. 10

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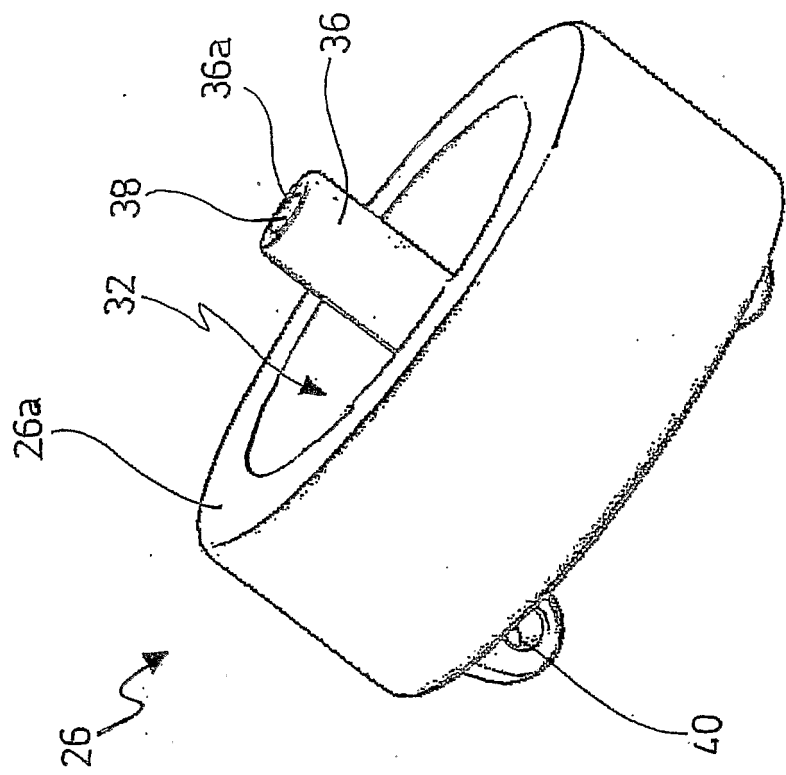


FIG. 12

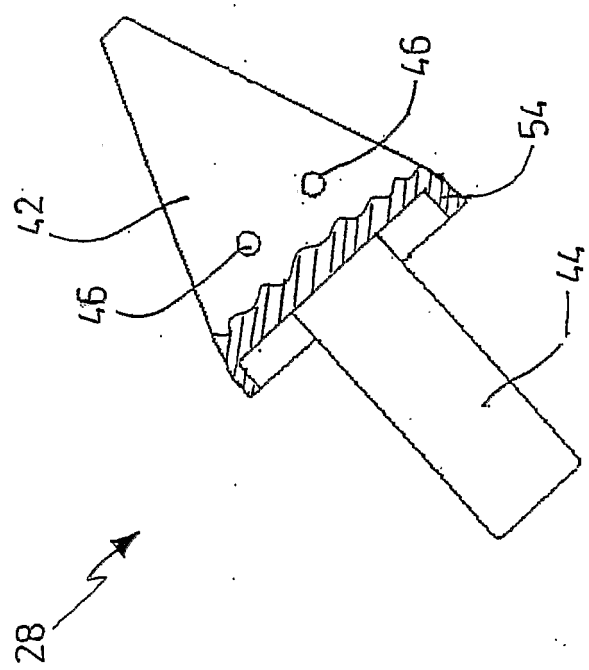


FIG. 11

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FIG.13

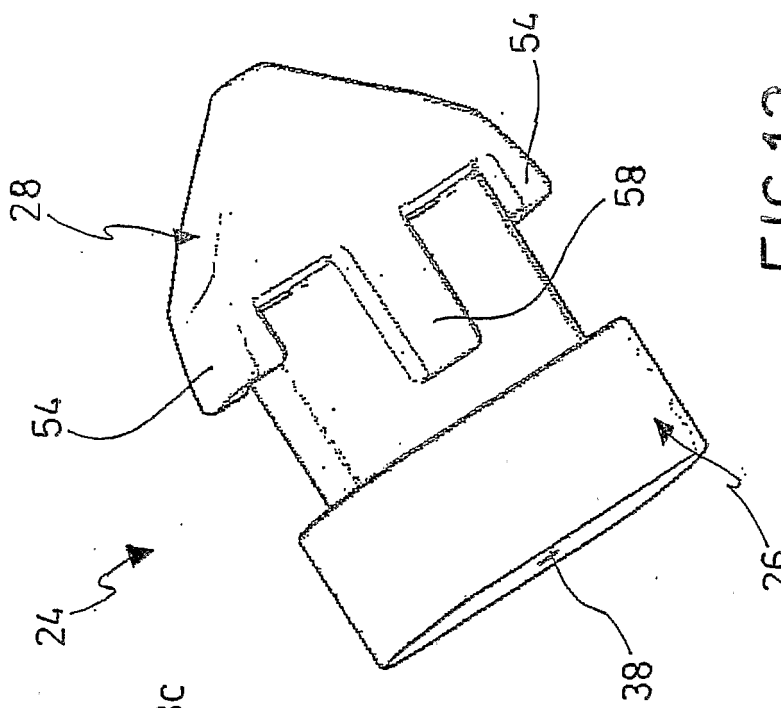
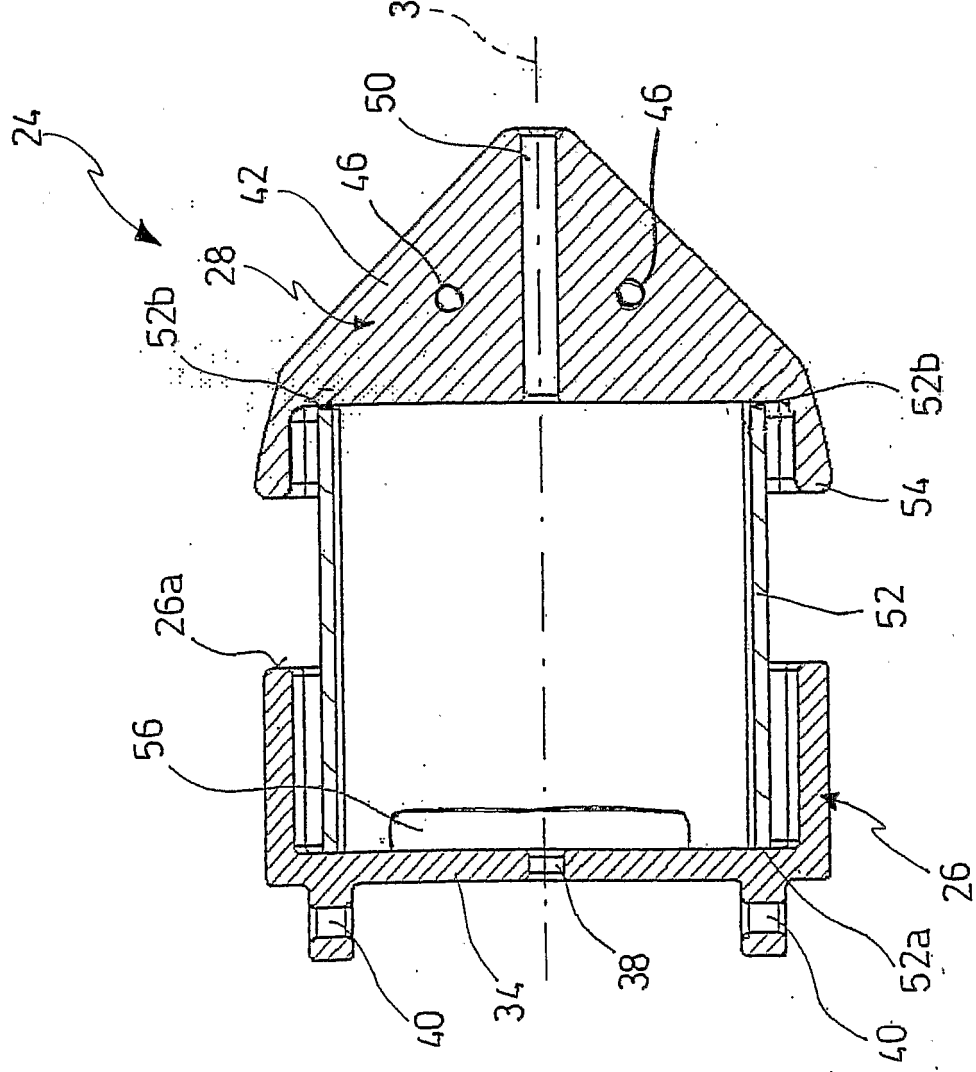


FIG.14



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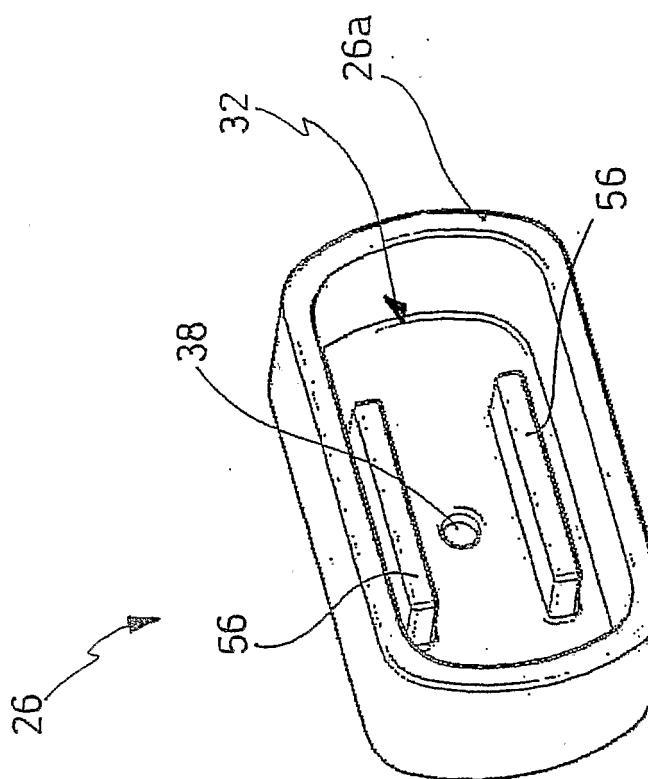


FIG. 15

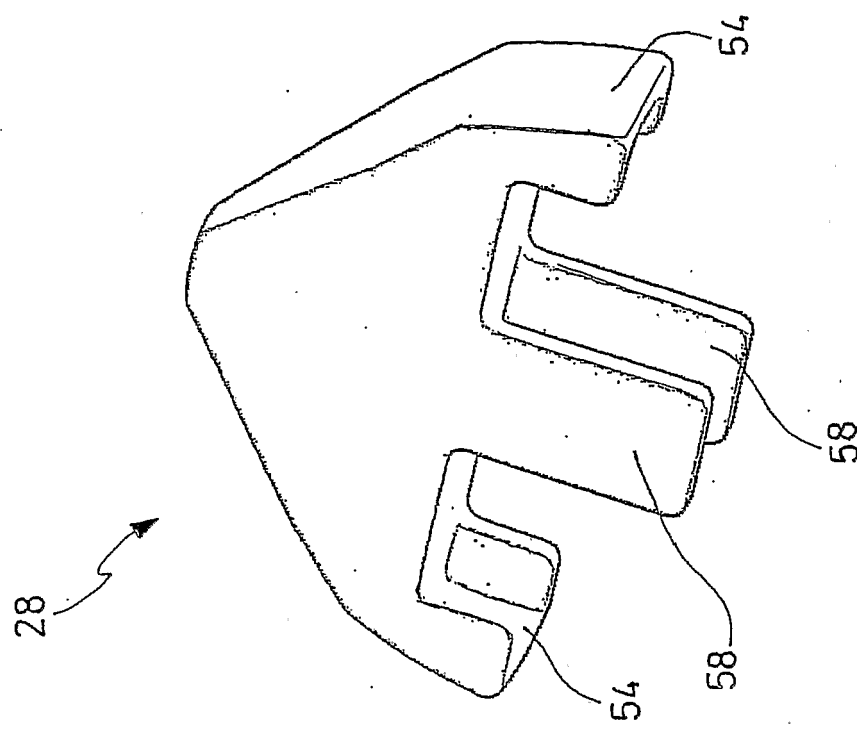


FIG. 16

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FIG. 18

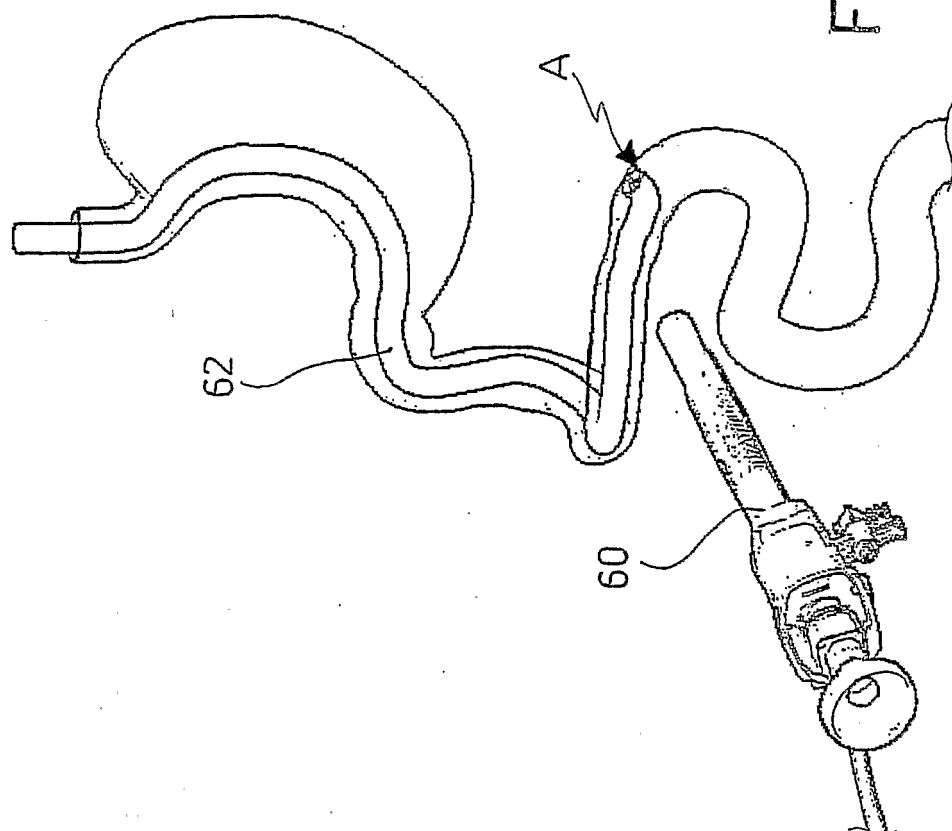
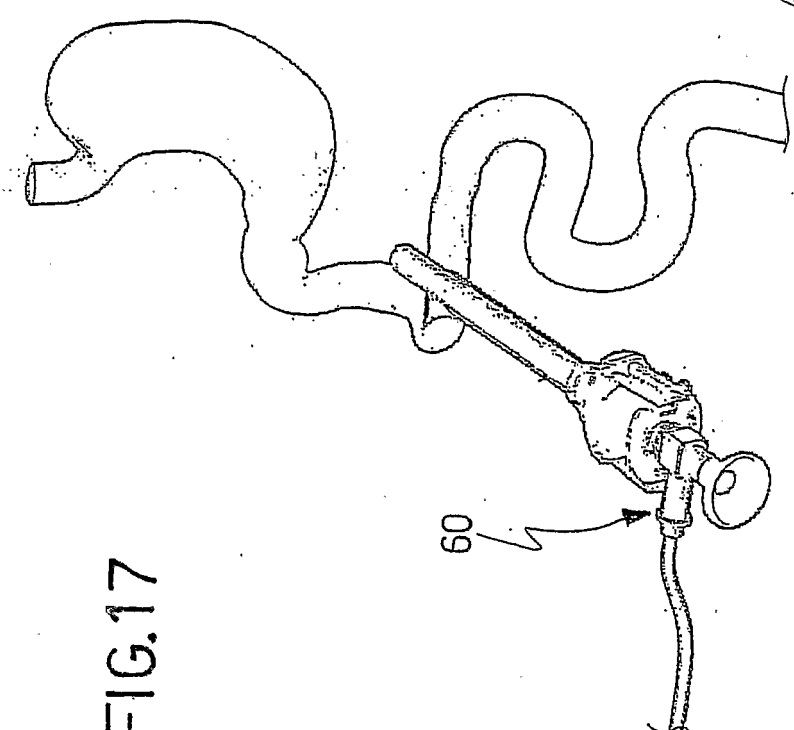


FIG. 17



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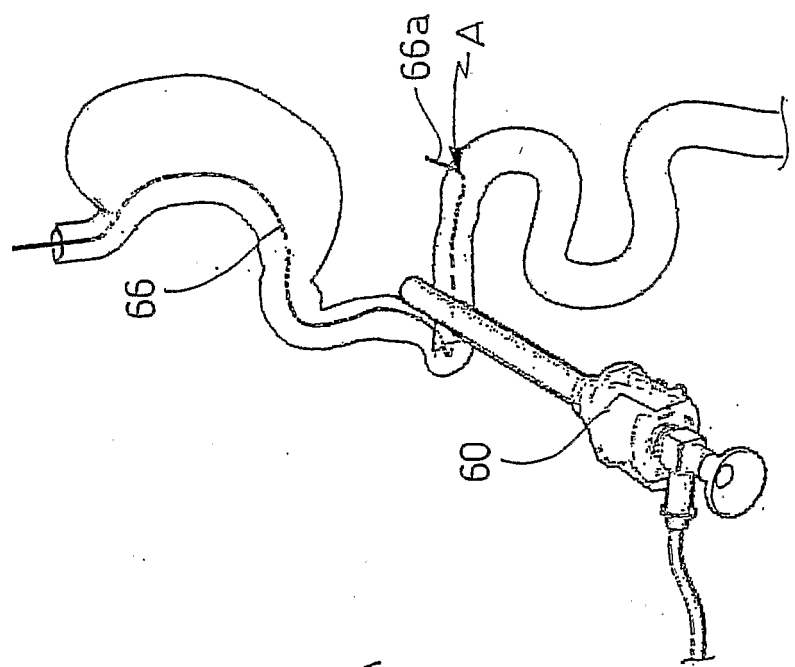


FIG. 20

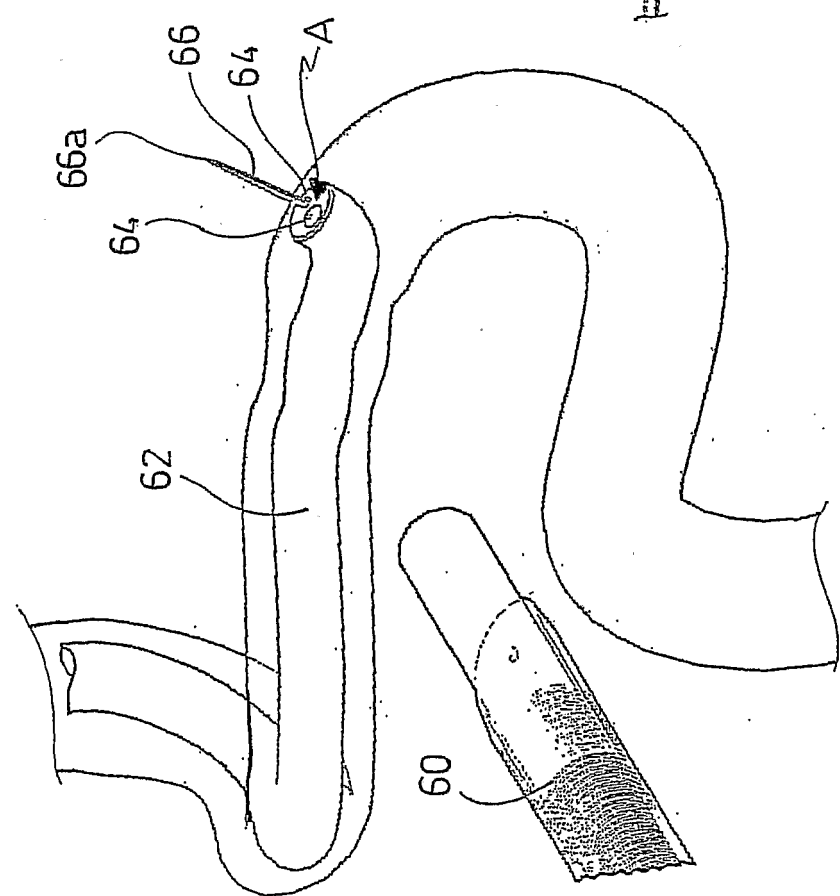


FIG. 19

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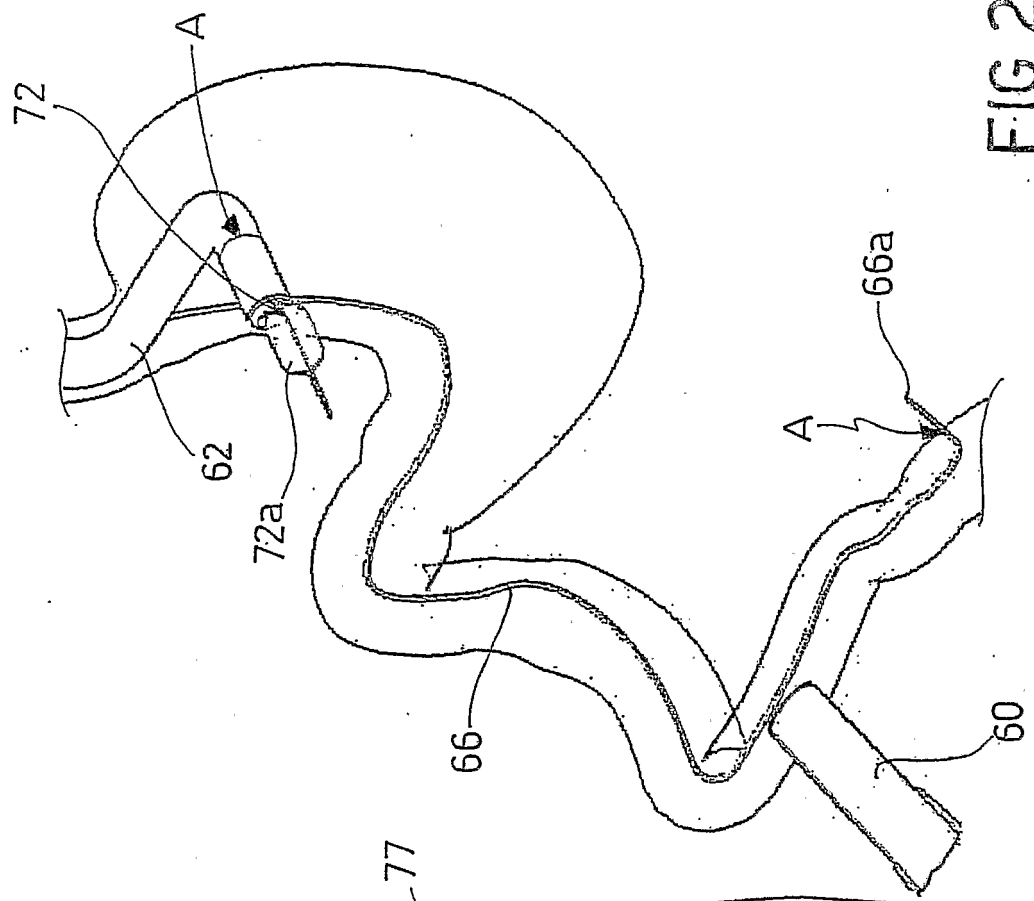


FIG. 22

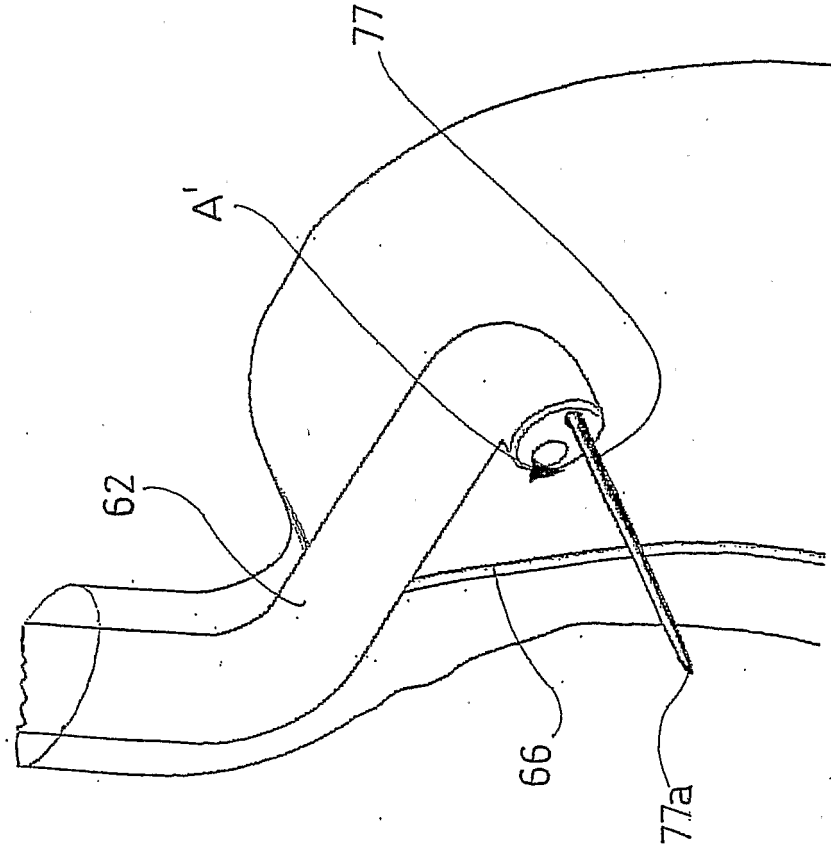
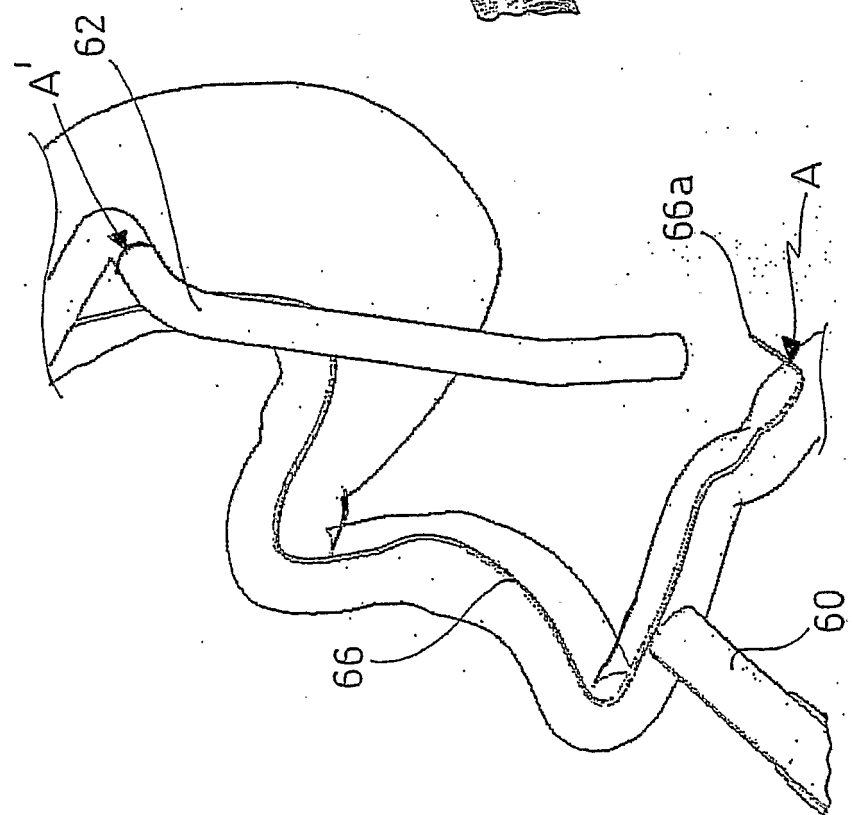
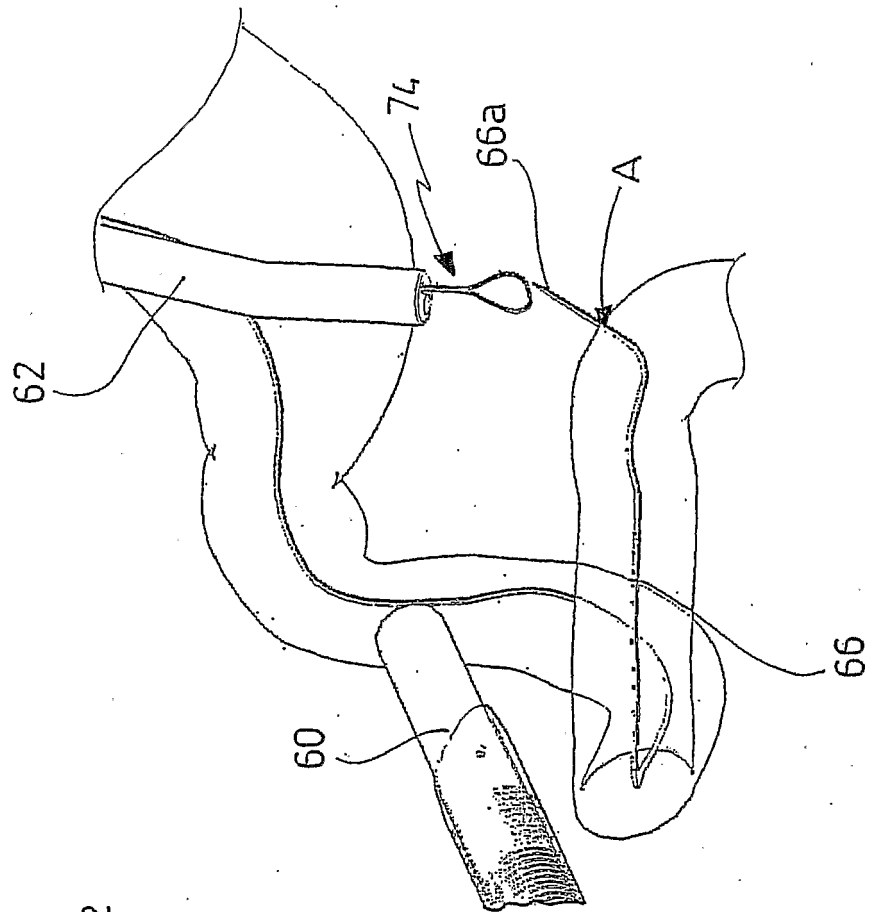
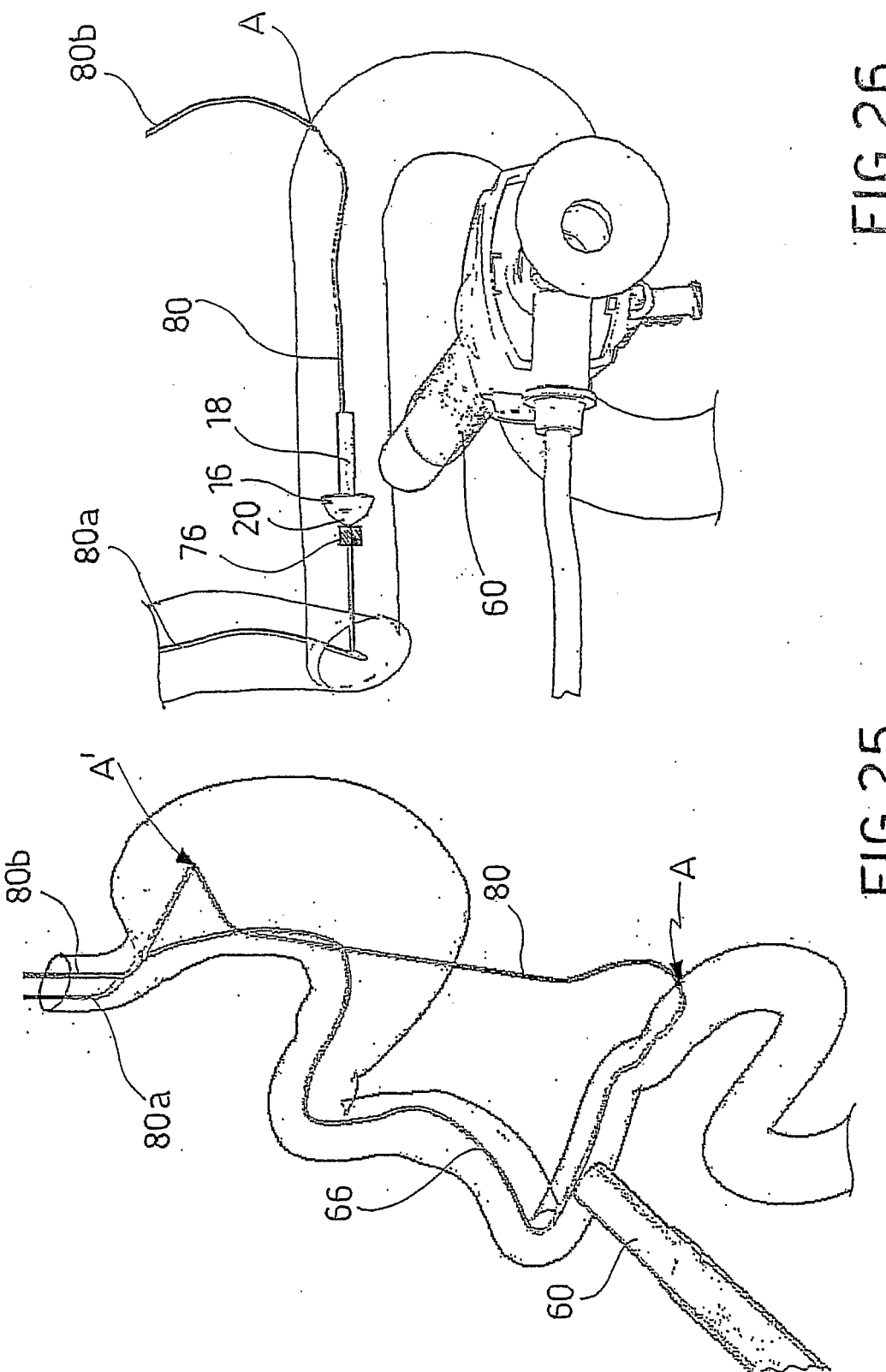


FIG. 21

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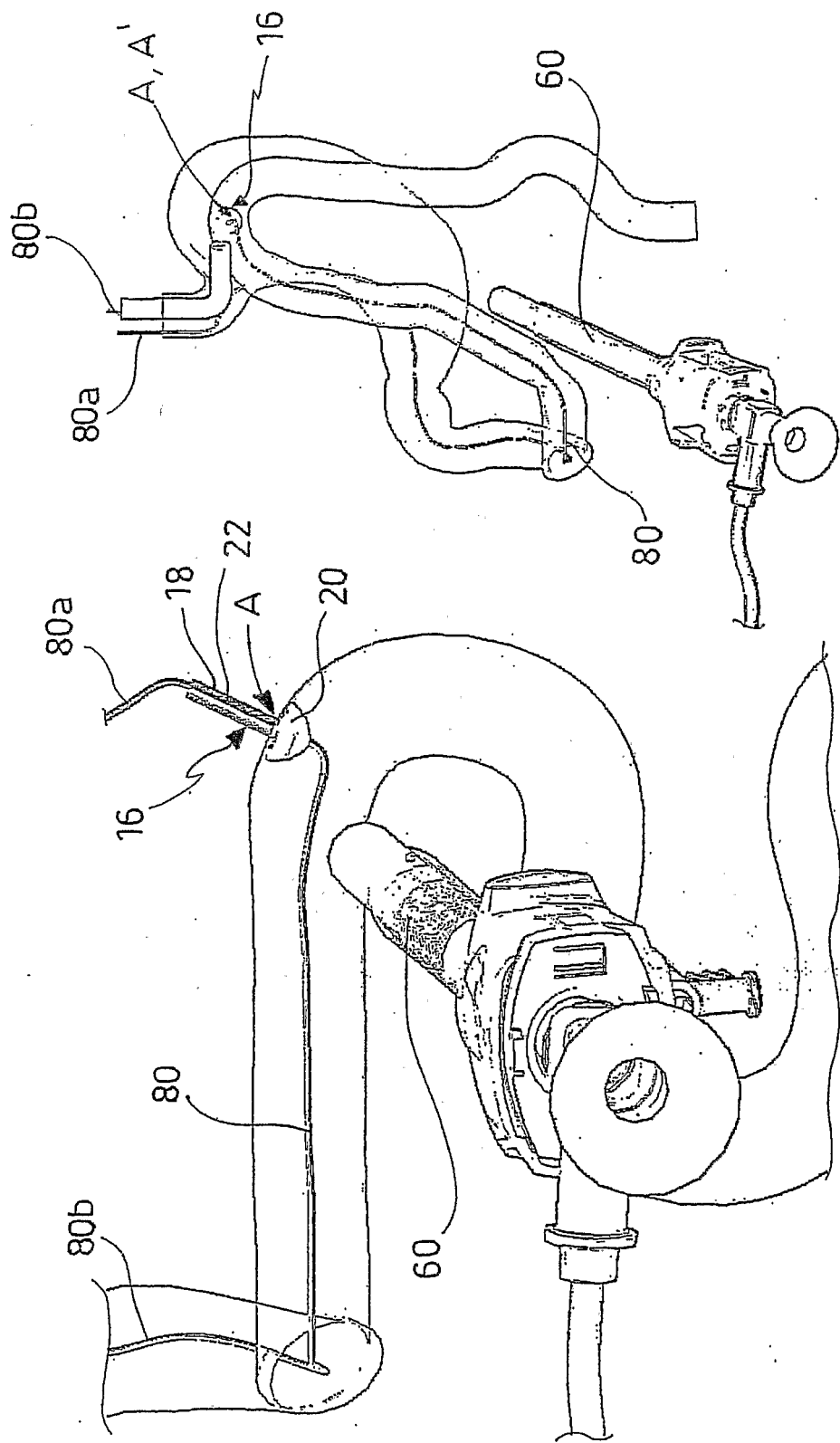


FIG. 27

FIG. 28

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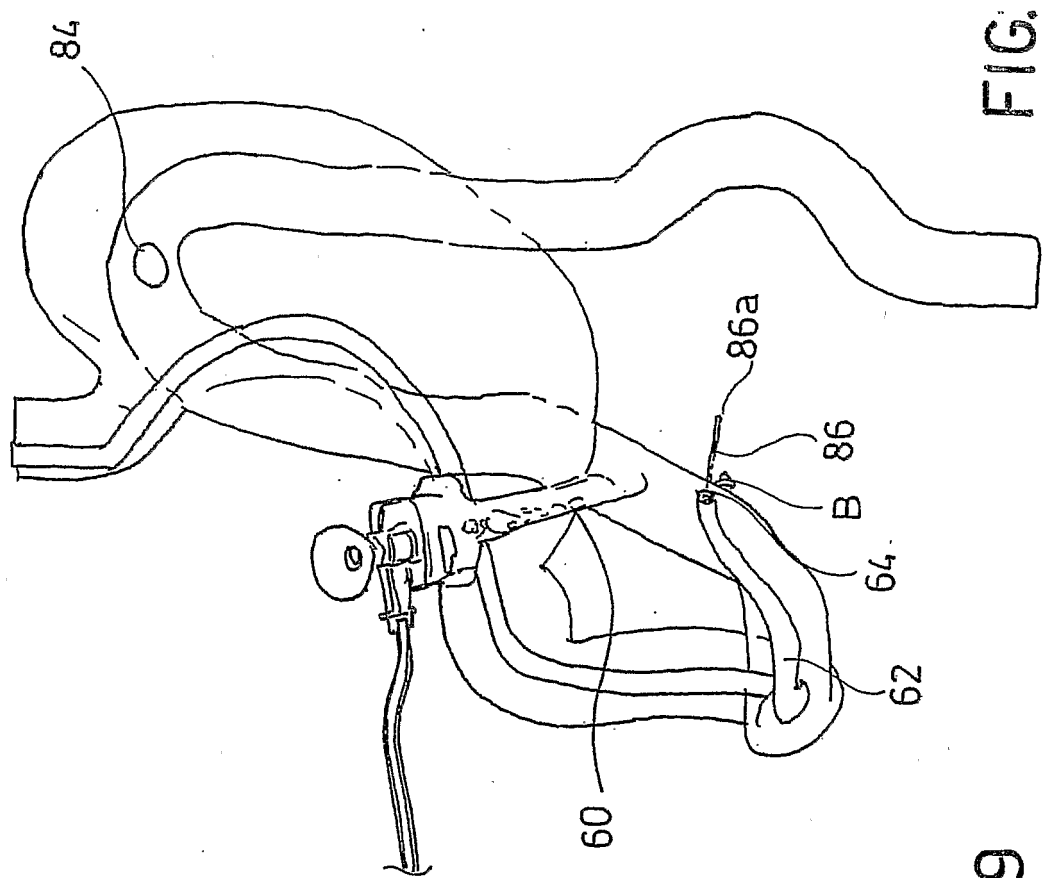
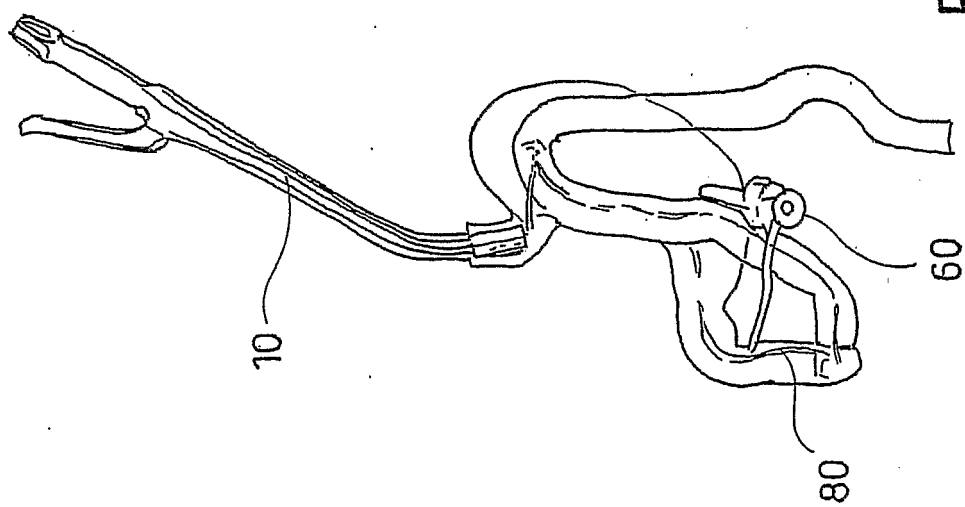


FIG. 29



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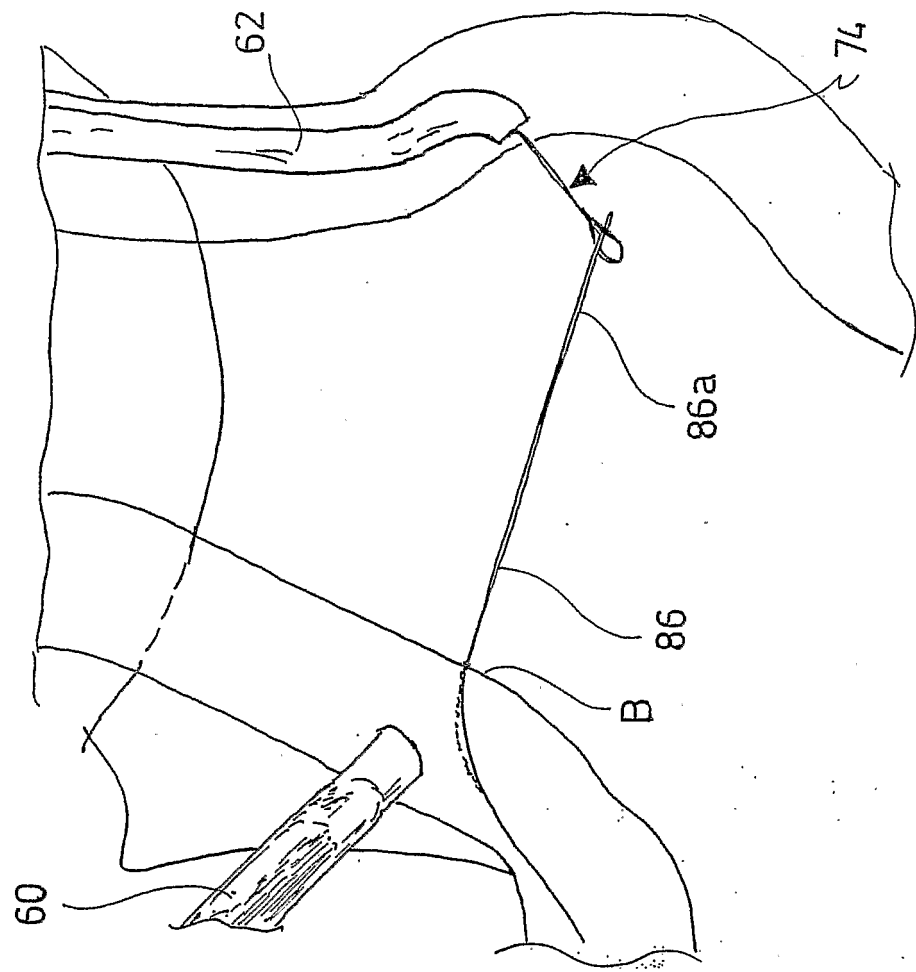


FIG. 32

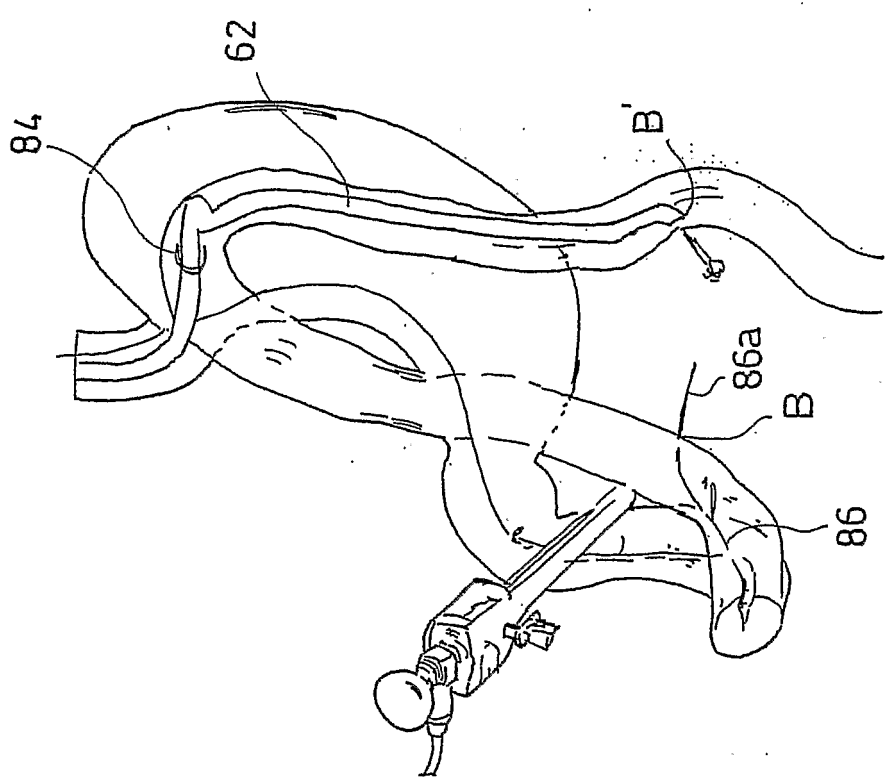


FIG. 31

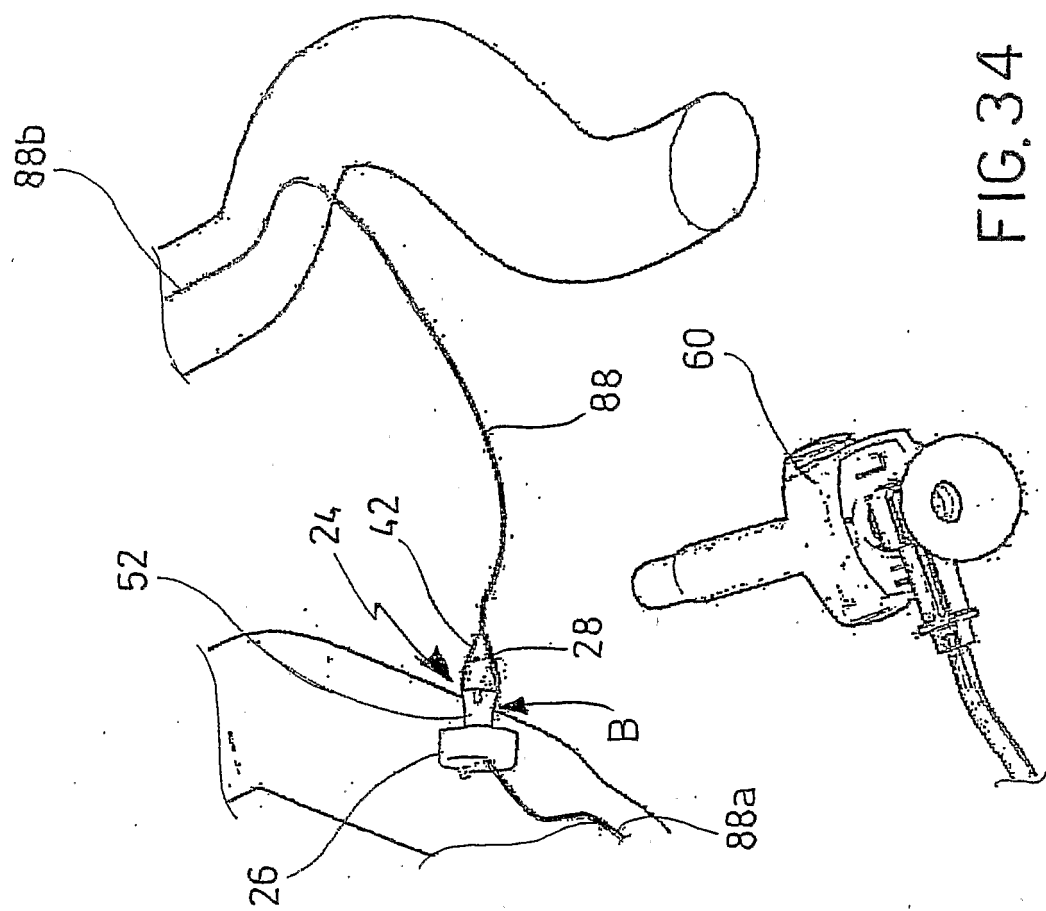


FIG. 34

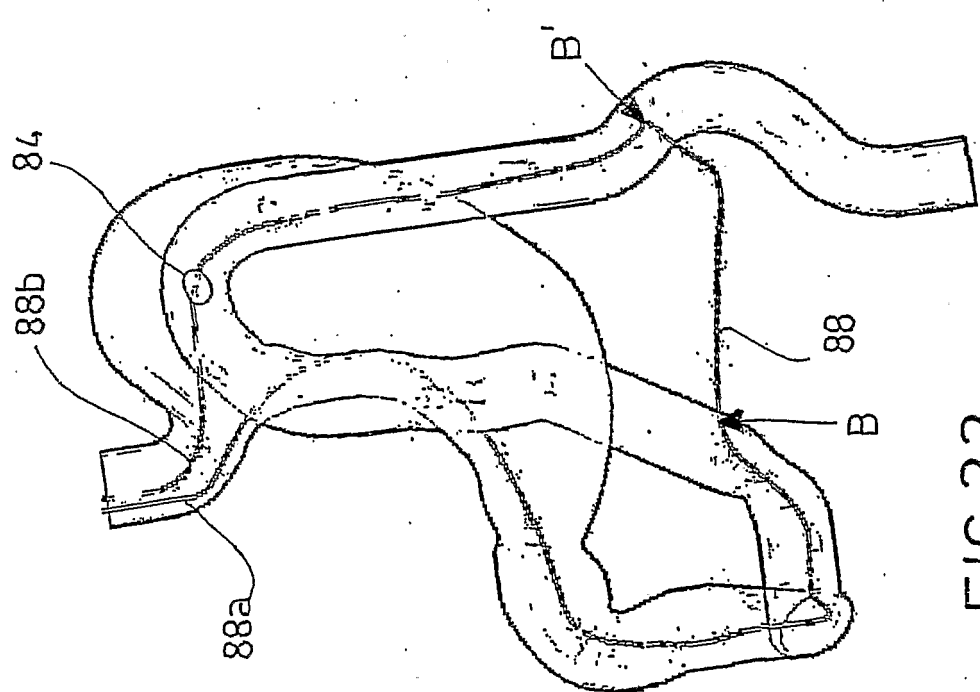


FIG. 33

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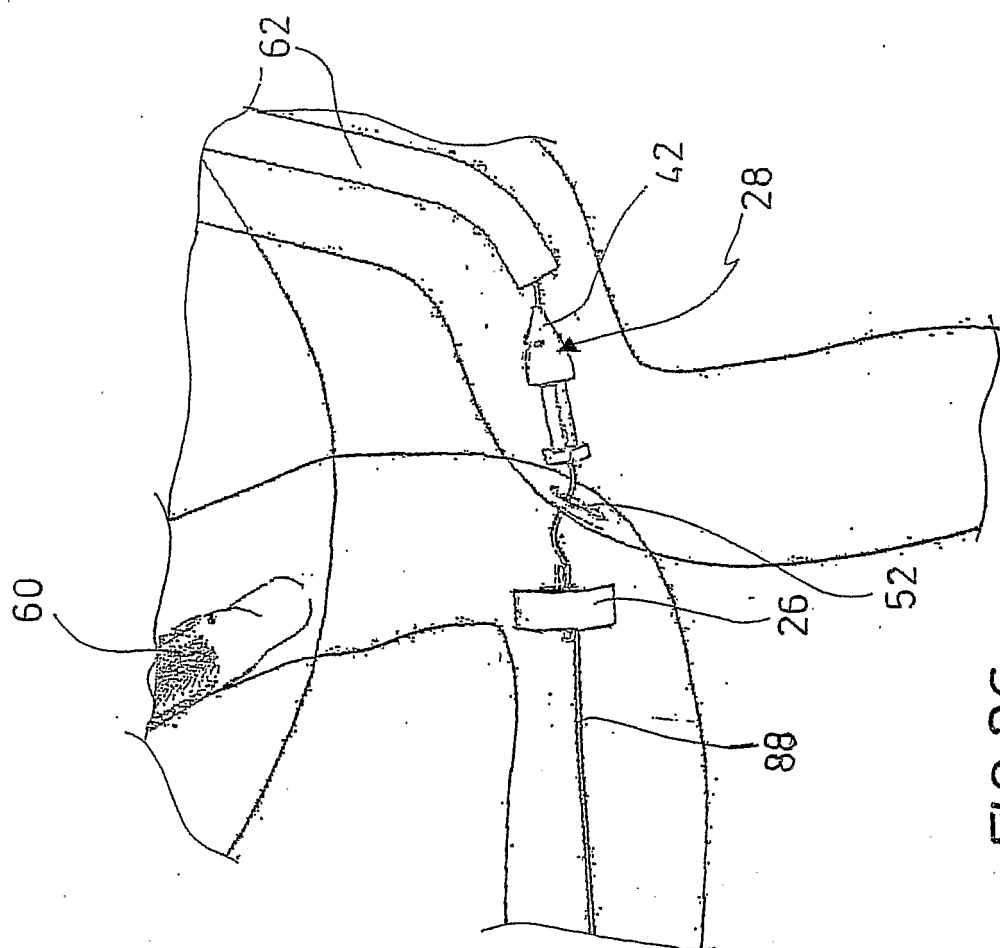


FIG. 36

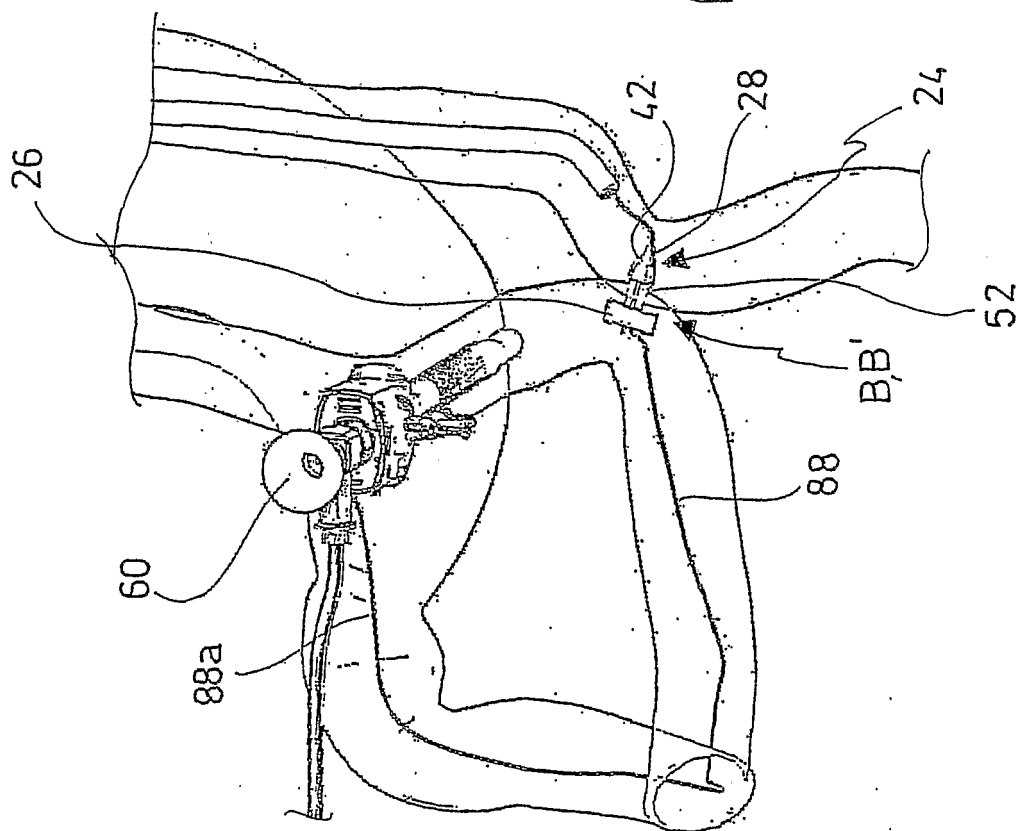


FIG. 35

FIG. 38

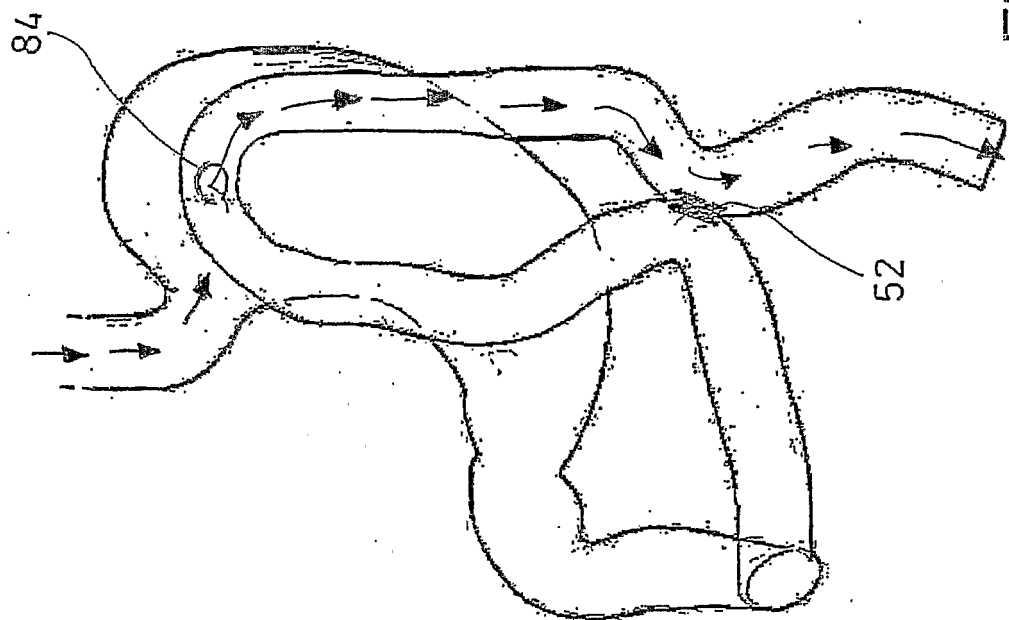


FIG. 37

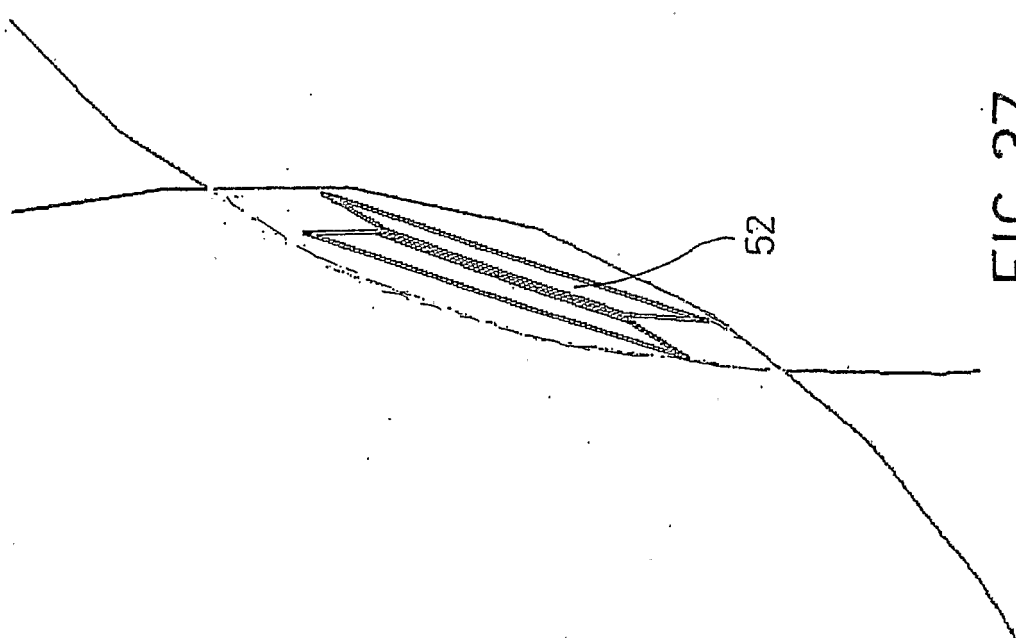


FIG. 40

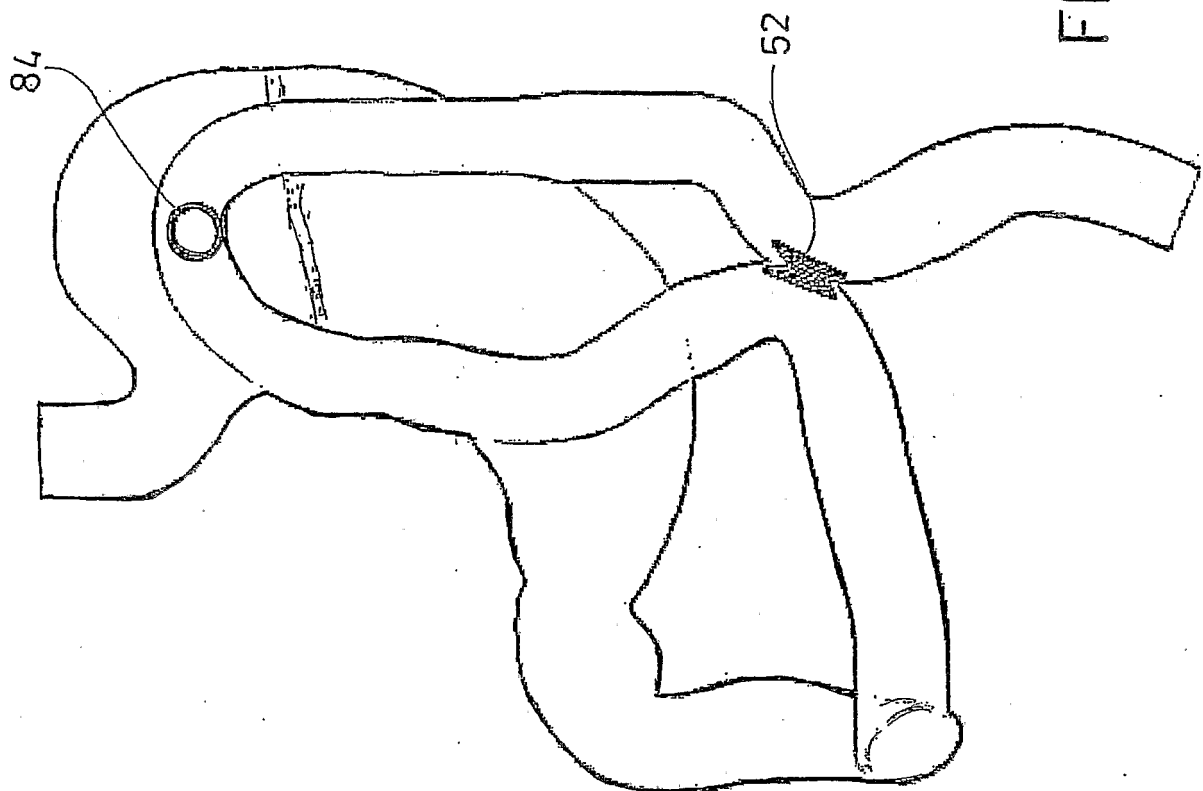
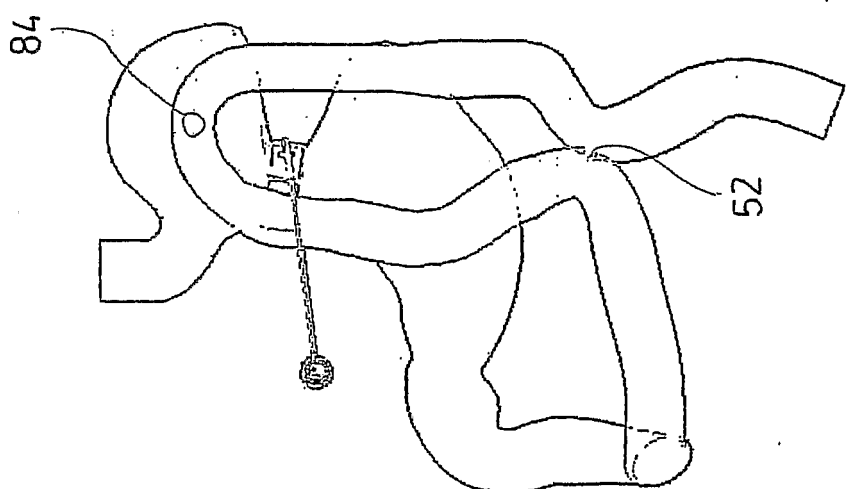


FIG.39



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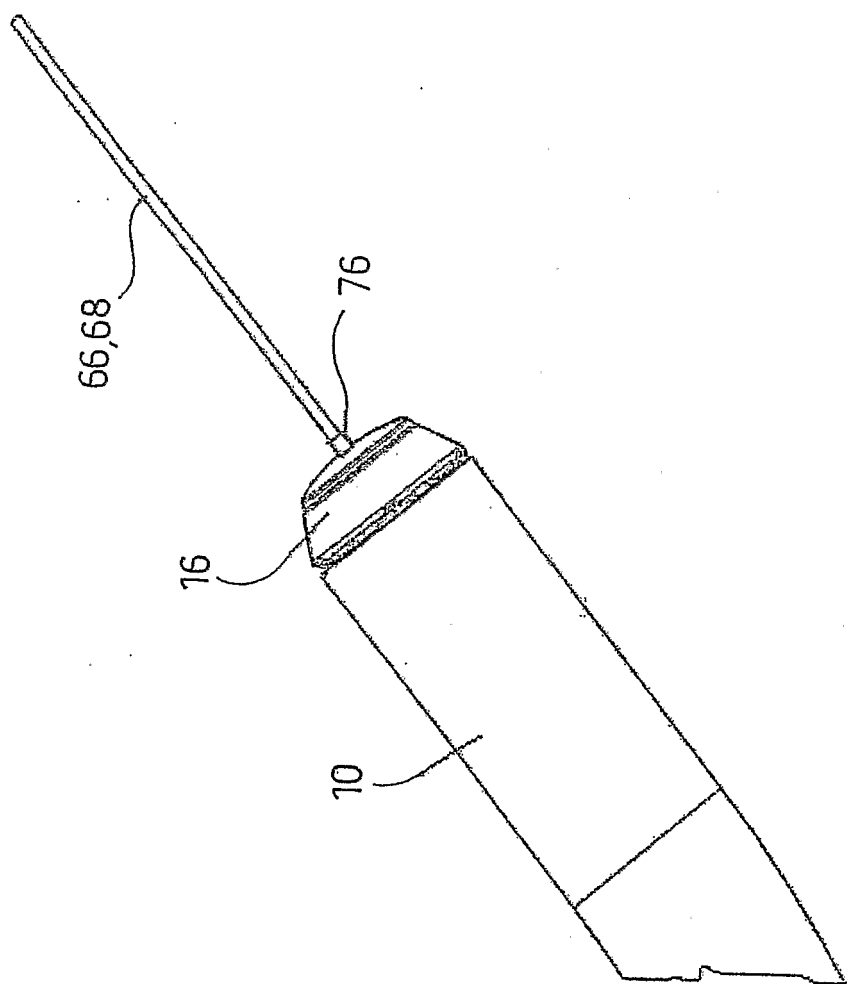
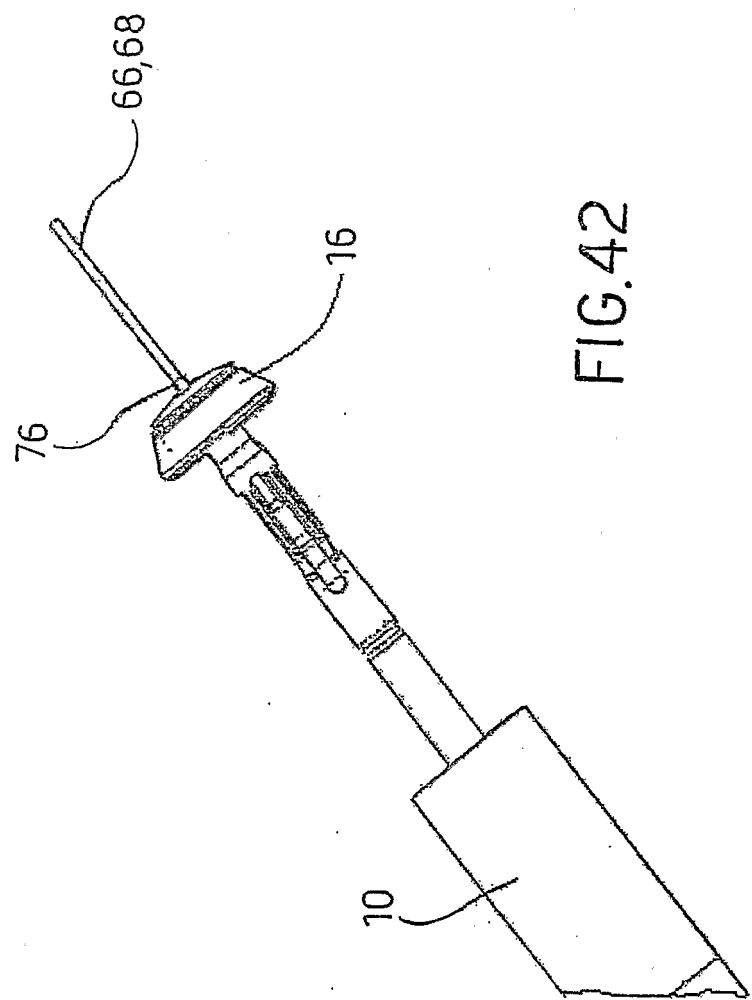


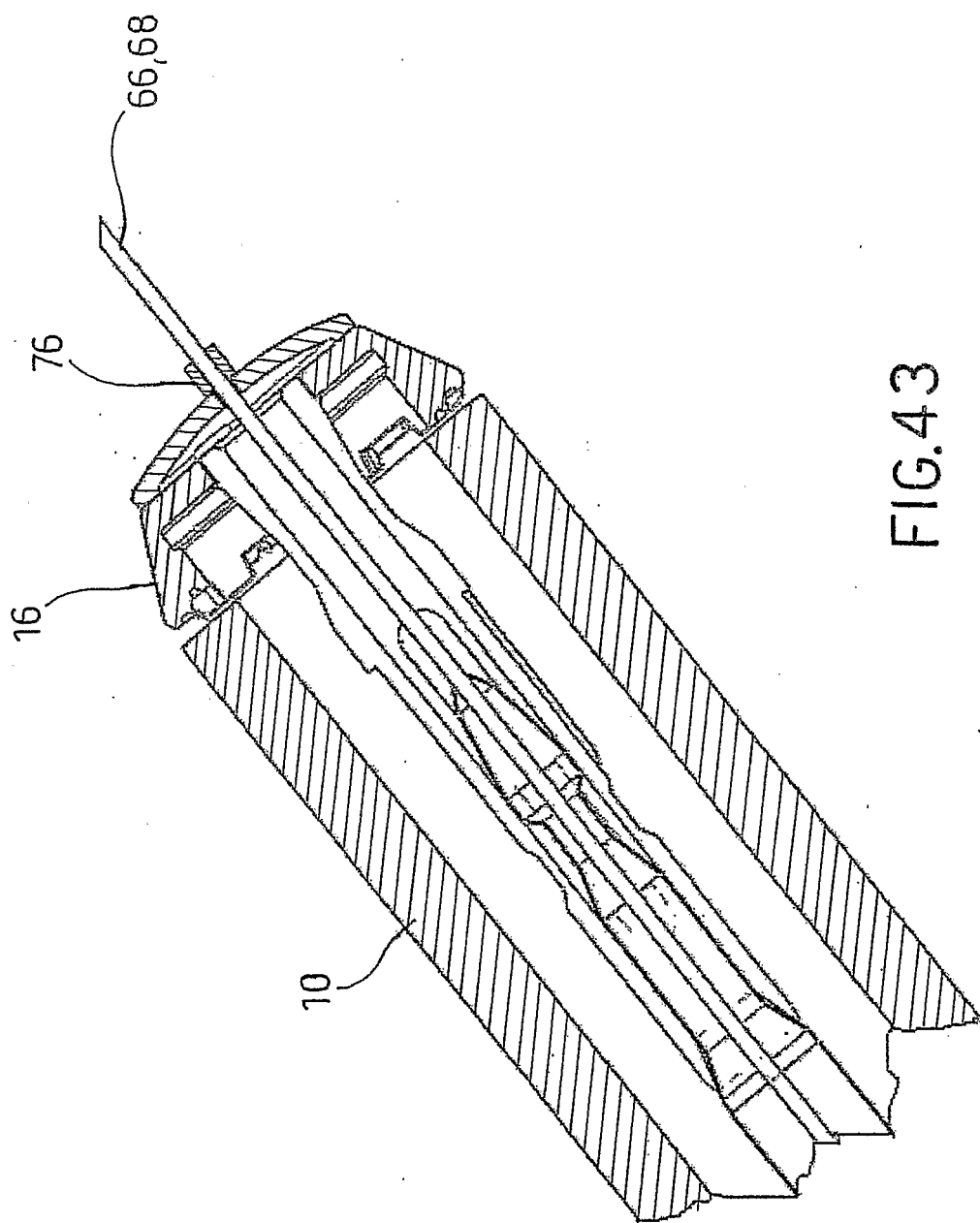
FIG. 41

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FIG.42



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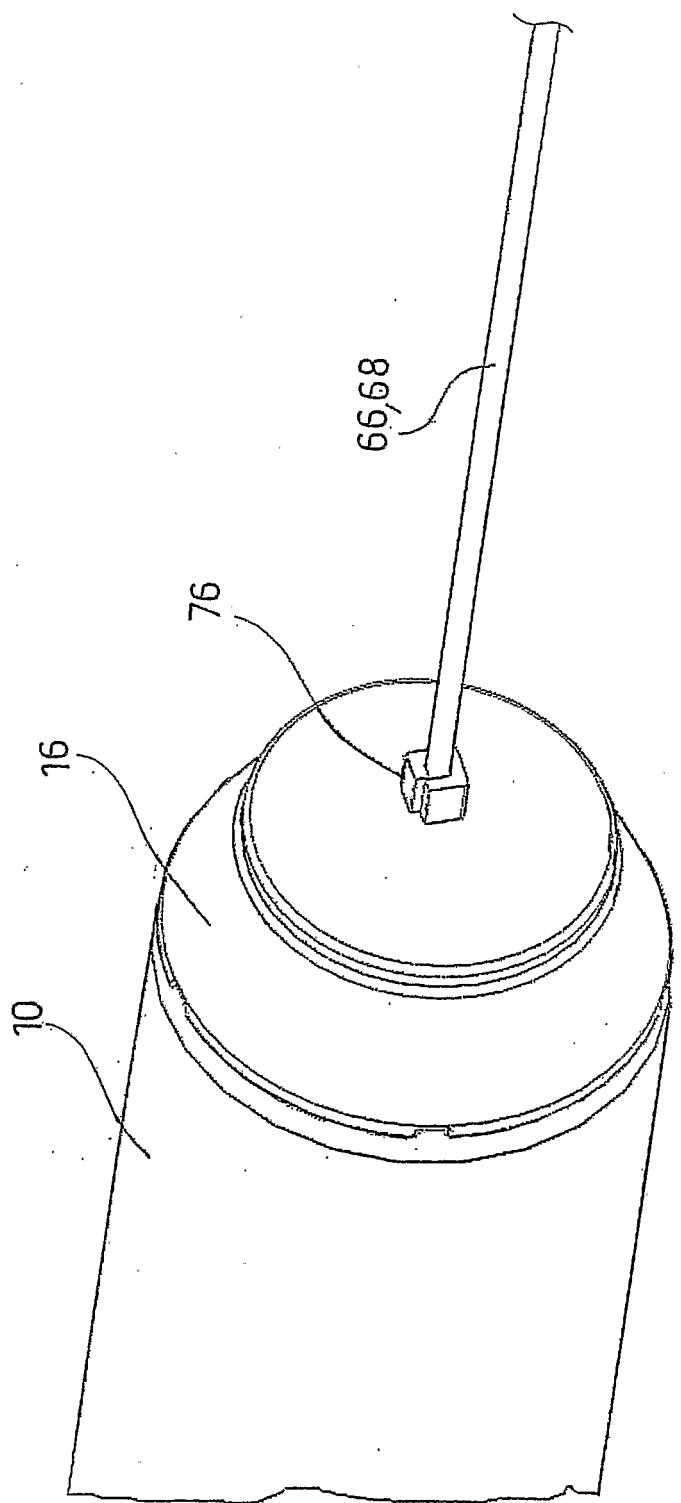


FIG. 44

# INTERNATIONAL SEARCH REPORT

International Application No  
PCT/IT2005/000346

**A. CLASSIFICATION OF SUBJECT MATTER**

IPC 7 A61B17/11

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)

IPC 7 A61B

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 practical, search terms used)

EPO-Internal

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 2003/216749 A1 (ISHIKAWA MASAHIRO ET AL.) 20 November 2003 (2003-11-20) paragraphs '0046!, '0055!, '0059!; figures 1,2B,3A,10,11,13 -----	1-7
X	US 6 482 224 B1 (MICHLER ET AL.) 19 November 2002 (2002-11-19) figures 1,3 -----	1-3,6,7
X	EP 0 478 949 A (UNITED STATES SURGICAL CORPORATION) 8 April 1992 (1992-04-08) figures 1A,3A,11 -----	1-4,6,7
X	US 6 685 716 B1 (FLAHERTY J. CHRISTOPHER ET AL) 3 February 2004 (2004-02-03) figures 5A,5B,7,11 ----- -----	1-3,6,7
		-/-

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

° Special categories of cited documents :

- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier document but published on or after the international filing date
- "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)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed

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

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

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

"&" document member of the same patent family

Date of the actual completion of the international search

26 October 2005

Date of mailing of the international search report

04/11/2005

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2  
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Authorized officer

Assion, J-C

**INTERNATIONAL SEARCH REPORT**International Application No  
PCT/IT2005/000346**C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT**

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 2002/123698 A1 (GARIBOTTO JOHN ET AL) 5 September 2002 (2002-09-05) figures 11A,12A -----	1-4,6,7

## INTERNATIONAL SEARCH REPORT

International application No.  
PCT/IT2005/000346

### Box II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1.  Claims Nos.: 8-51 because they relate to subject matter not required to be searched by this Authority, namely:  
**Rule 39.1(iv) PCT – Method for treatment of the human or animal body by surgery**
2.  Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.  Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

### Box III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1.  As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.  As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.  As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.  No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

#### Remark on Protest

- The additional search fees were accompanied by the applicant's protest.
- No protest accompanied the payment of additional search fees.

## INTERNATIONAL SEARCH REPORT

International Application No  
PCT/IT2005/000346

Patent document cited in search report		Publication date		Patent family member(s)		Publication date
US 2003216749	A1	20-11-2003	US	2005043720 A1		24-02-2005
US 6482224	B1	19-11-2002	US	2002099389 A1		25-07-2002
EP 0478949	A	08-04-1992	CA	2049103 A1		07-03-1992
			DE	69124900 D1		10-04-1997
			DE	69124900 T2		24-07-1997
			ES	2098288 T3		01-05-1997
			US	5382257 A		17-01-1995
US 6685716	B1	03-02-2004		NONE		
US 2002123698	A1	05-09-2002		NONE		

专利名称(译)	用于治疗肥胖症的装置和方法		
公开(公告)号	<a href="#">EP1811905A1</a>	公开(公告)日	2007-08-01
申请号	EP2005761295	申请日	2005-06-16
[标]申请(专利权)人(译)	伊西康内外科公司		
申请(专利权)人(译)	爱惜康内镜手术 , INC.		
当前申请(专利权)人(译)	爱惜康内镜手术 , INC.		
[标]发明人	TACCHINO ROBERTO BILOTTI FEDERICO DARCANGELO MICHELE KUHNS JESSE J CLEM MICHAEL F		
发明人	TACCHINO, ROBERTO BILOTTI, FEDERICO D'ARCANGELO, MICHELE KUHNS, JESSE J. CLEM, MICHAEL, F.		
IPC分类号	A61B17/11		
CPC分类号	A61B17/1114 A61B2017/00862 A61B2017/0404 A61B2017/1103 A61B2017/1139 A61B2017/22038		
优先权	MI2004002129 2004-11-05 IT		
其他公开文献	<a href="#">EP1811905B1</a>		
外部链接	<a href="#">Espacenet</a>		

**摘要(译)**

一种用于将组织拉到一起并产生吻合的装置，包括主导丝（66,86），其适于通过吻合接收和牵拉要连接的组织的部分（A; B）。主导丝（66,86）的形状类似于开口环，其通过吻合术（66,86）与待连接的组织的部分（A，A&#39;; B，B&#39;）交叉，其中末端主导线（66,86）的不同并且可以彼此区分。主导丝（66,86）具有管状或中空结构，适于容纳至少一个针，用于组织的推动或射频穿孔。