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(54) **INSUFFLATING OPTICAL SURGICAL INSTRUMENT**

INSUFFLIERENDES OPTISCHES OPERATIONSINSTRUMENT

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DescriptionBACKGROUND OF THE INVENTIONField of the Invention

[0001] This invention generally relates to surgical instruments and, in particular, to surgical instruments providing visual entry and visual insufflation.

Discussion of the Prior Art

[0002] Laparoscopic surgery of the abdominal area typically requires the introduction of an insufflation gas into the peritoneal cavity of the patient. The insufflation gas is usually pressurized to about 10 mm Hg above atmospheric pressure. This in turn lifts the abdominal wall away from the organs underlying it. Cannulas having seals are then placed at various locations through the abdominal wall to allow the use of a laparoscope and operating instruments. It is well known that establishing access to a non-inflated peritoneal cavity can be a very dangerous part of any laparoscopic procedure. The most common method to achieve insufflation is to pass a sharp needle through the abdominal wall and into the abdominal region, and then inject a gas through the needle and into the region thereby creating an enlarged or ballooned cavity to accommodate a laparoscopic procedure. Unfortunately, insertion of the needle has been required without any visual aid to facilitate location of the sharp needlepoint. In order to reduce the probability of inadvertent penetration of delicate internal organs in this "blind" procedure, the sharp insufflation needle has been provided with a spring-loaded and retractable safety mechanism. Such a needle, commonly referred to as a Veress needle, is disclosed in WO 96/01132.

[0003] The safety mechanisms associated with most insufflation needles consist of a blunt or rounded member disposed within the lumen of the needle, and biased by a spring to an extended position beyond the needle tip. This spring must be responsive to the insertion pressure during placement of the needle but must be capable of immediately moving forward when that pressure is relieved. This is a highly mechanical event at best and offers less than optimal arrangement. As pointed out above, a drawback of this procedure is it is normally performed blindly. A consequence of this blind insertion is the surgeon may inadvertently damage the organs and tissues underlying the abdominal wall such as major blood vessels and the intestinal tract. Once access is gained, it can take several minutes for the gas to insufflate the abdomen and while this is happening the surgeon may be unaware of any complications caused by the insertion of the needle. WO 96/01132 discloses a Veress needle that optionally provides for a physician, once the needle has been inserted, to subsequently insert an optical viewing scope into an obturator of the Veress needle, to enable the location of the distal end to be determined.

[0004] Another commonly used method of gaining initial access to the peritoneal cavity is by using a procedure known as the Hasson technique. This method involves making a mini-laparotomy and using the fingers to bluntly dissect the tissues of the abdominal wall and thereby creating an access similar to an open surgical procedure. This method is generally considered to be safer but not without risks, and results in an access site that is not well suited for the subsequent introduction and use of a laparoscopic cannula. The cannula is typically held in place with an additional device that allows the cannula to be tied down with sutures to prevent it from slipping out of the abdominal wall. This also leaves a large defect and is difficult to perform in large abdominal walls.

[0005] Some surgeons have used trocars designed for use with laparoscopes for the initial entry into the peritoneal cavity. These devices allow the placement of a laparoscope through the internal diameter of the trocar and have a trocar tip that is made of clear plastic to allow the surgeon to visualize the passage of the tip through the abdominal wall. However, in order to allow the subsequent introduction of insufflation gas through the cannula, the trocar and cannula must be inserted all the way through the wall of the abdomen and this in turn can be potentially dangerous as the tip of the trocar may have to advance as much as one inch beyond the distal surface of the abdominal wall and into the underlying anatomical structures. As such, there remains a need in the art for an improved surgical instrument that provides visual entry and visual insufflation, and that minimizes the risks of damaging organs, tissues and vessels underlying a body wall.

SUMMARY OF THE INVENTION

[0006] The invention is directed to surgical instruments providing visual entry and visual insufflation with minimal risks of injury to organs, tissues and vessels underlying a body wall. It is appreciated that the concept of the invention may be applied to any surgical instrument that provides the ability to insufflate under direct vision of the site of insufflation, regardless of the size of the instrument and the type of insufflation fluid. More specifically, the surgical instrument provides the ability to transfer an insufflation fluid such as CO₂ or saline from outside a patient to inside a surgical cavity under vision. The insufflation fluid may be transferred inside a lumen, along a body channel or through a coiled tube of a surgical instrument or scope used for vision.

[0007] According to the present invention there is provided an insufflating surgical instrument as recited in Claim 1.

[0008] In one aspect, at least one of the tip and the shaft is formed of a transparent material to facilitate visualization of the abdominal wall and the abdominal region. With this aspect, the shaft and the tip are configured to enable insertion of a laparoscope. In particular, the shaft includes a lumen extending along the axis between

the proximal end and the distal end to enable insertion of the laparoscope. The lumen and insufflation channel may be formed as separate channels or as one shared channel. The insufflating surgical instrument may further comprise a second vent hole being in connection with the insufflation channel and formed along the shaft. It is appreciated that the tip may be blunt, the shaft and the tip may be integrally formed, and the vent hole may be of any geometric shape including round, oval, square and rectangular. With this aspect, at least one of the tip and the shaft may be formed of a translucent or a transparent material such as polycarbonate. The blunt tip may further comprise a marker to indicate when the vent hole has been positioned for insufflation. More specifically, the marker indicates the point where the vent hole has penetrated the abdominal wall. The shaft of the invention may further comprise a scope lock to prevent the laparoscope from being inserted too far into the shaft and block at least one of the insufflation channel and the vent hole. In another aspect, the tip may be sharp, pointed or bladed to facilitate penetration of body tissue.

[0009] The seal housing may comprise a septum seal and a plurality of leaflets forming an instrument seal in the presence of a laparoscope, and providing a zero seal in the absence of an instrument. The thickness of the leaflets may be formed to a desired dimension to create a pressure release mechanism that inverts and releases pressure if the abdominal pressure within the patient undergoes a sudden spike. The septum seal may be formed of an elastomeric material including Kraton, silicone and the like. The seal housing may further comprise a duckbill or a double duckbill valve distal of the leaflets to further limit gas or fluid escape.

[0010] These and other features of the invention will become more apparent with a discussion of the various embodiments in reference to the associated drawings.

DESCRIPTION OF THE DRAWINGS

[0011] The accompanying drawings, which are included in and constitute a part of this specification, illustrate the embodiments of the invention and, together with the description, explain the features and principles of the invention. In the drawings:

FIG. 1 illustrates a typical laparoscopic abdominal surgery of the prior art;

FIG. 2 illustrates a perspective view of an insufflation needle of the prior art;

FIG. 3 illustrates a perspective view of an insufflating optical trocar of the invention;

FIGS. 4(a) and 4(b) illustrate cross-sectional views of an alternative insufflating optical trocar of the invention;

FIG. 5 illustrates a perspective view of a septum seal for use with the insufflating optical trocar of the invention;

FIG. 6(a)-6(d) illustrate cross-sectional views of the

septum seal for use with the insufflating optical trocar of the invention;

FIGS. 7(a) and 7(b) illustrate the insufflating optical trocar and cannula of the invention;

FIGS. 8(a)-8(f) illustrate different geometric shapes and patterns of the vent hole of the invention;

FIGS. 9(a)-9(e) illustrate additional tip designs of an insufflating optical surgical instrument in accordance with additional aspects of the invention;

FIGS. 10(a)-10(c) illustrate flip-top or flip-tip designs of insufflating optical surgical instruments in accordance with additional embodiments of the invention;

FIGS. 11(a)-11(c) illustrate cross-sectional views of insufflating valve vents in accordance with additional embodiments of the invention;

FIGS. 12(a) and 12(b) illustrate an insufflating surgical instrument including an insufflating optical trocar and a cannula having a gas channel for transferring insufflation gas to the trocar in accordance with another embodiment of the invention; and

FIGS. 13(a)-13(i) illustrate additional tip designs in accordance to other aspects of the invention to facilitate penetration of body tissue.

DESCRIPTION OF THE INVENTION

[0012] Referring to FIG. 1, there is shown a typical laparoscopic abdominal surgery where an inflation needle 10 is inserted through a body or abdominal wall 15 and into an abdominal cavity 25. A gas is passed through the needle 10 to create a space within the abdominal cavity 25. This procedure is referred to as insufflation. The needle 10 is referred to as an insufflation needle and the gas supply is referred to as an insufflation gas. The insufflation needle 10 is placed through the body wall 15 blindly. In other words, there is no direct visualization of the procedure from the inside of the body wall 15. As explained earlier, the current procedure may inadvertently damage organs and tissues underlying the body or abdominal wall 15 such as major blood vessels and the intestinal tract. It is not uncommon for there to be internal structures attached to the internal side of the body wall 15. This is especially so in the case of the abdominal cavity 25. Portions of the intestines, colon and bowel may be attached to the abdominal wall 15. These attachments are referred to as adhesions.

[0013] Adhesions represent a potential complication in laparoscopic surgery. This is especially the case as the procedure is initiated using a sharp or pointed instrument such as the insufflation needle 10. The delicate internal structures that may be attached by adhesions may inadvertently be pierced by the introduction of the insufflation needle 10. This can be very serious and may go undetected for some time.

[0014] Referring to FIG. 2, the typical insufflation needle 10 comprises an elongate tubular body 12, a proximal connecting housing 14, a sharp, pointed distal tip 16 and a spring, biased internal blunt core 18 with a blunt end

24 that extends beyond the sharp distal tip 16 under the influence of an extended compression spring 20. A typical placement of the insufflation needle 10 requires a user to push the sharp distal tip 16 into the abdominal wall 15, which pushes the blunt core 18 proximally, then continues to push until the distal end 22 is through the body wall 15. At that point, the blunt end 24 moves forward and thereby protects delicate structures from being inadvertently punctured by the sharp distal tip 16 of the needle 10. The safety of such a device depends to a large extent on the design and integrity of the spring 20 since the sharp distal tip 16 remains within the established internal region or body cavity 25.

[0015] Referring to FIG. 3, there is shown a perspective view of an insufflating optical instrument or trocar 30 forming part of the surgical instrument of the invention. The insufflating optical trocar 30 is designed to separate tissue fibers during insertion through the body wall 15. The insufflating optical trocar 30 includes a shaft 34 having a lumen extending substantially along an axis between a proximal end and a distal end, a handle 35 disposed at the proximal end of the shaft 34, and a blunt tip 32 disposed at the distal end of the shaft 34. The shaft 34 of the insufflating optical trocar 30 is sized and configured for disposition within a working channel of a cannula system. With this disposition, the insufflating optical trocar 30 functions to penetrate a body or abdominal wall to provide the cannula with access across the body wall 15 and into the body cavity 25, such as the peritoneal or abdominal cavity.

[0016] The shaft 34 and tip 32 may be integrally formed of a transparent material to enable visualization of tissue during the insertion of the insufflating optical trocar 30 through the body wall 15. The insufflating optical trocar 30 is configured to enable the insertion of a conventional laparoscope, which typically includes an imaging element and fiber optic light fibers. The tip 32 further includes at least one vent hole 36, and preferably two or more vent holes one on each side of the tip 32, for the insufflation gas to transfer from the inside of the trocar 30 into the body or abdominal cavity 25. The vent hole 36 may be chamfered on the proximal side such that the vent hole does not core tissue as the insufflating optical trocar 30 enters through the body wall 15.

[0017] The shaft 34 includes at least one shaft vent 38 and preferably a plurality of shaft vents 38 along the axis between the proximal end and the distal end. It is appreciated that vent holes 36 and shaft vents 38 may be of any geometric shape including round, oval, square, rectangular, etc., as illustrated in FIGS. 8(a)-8(c) and may be configured in different patterns such as a waffle pattern as illustrated in FIG. 8(f). Furthermore, the tip 32 may be an open tip 180 or a non-coring tip 182 as illustrated in FIGS. 8(d) and 8(e), respectively, to allow the transfer of insufflation gas into the body cavity 25. Advantages of the shaft vents 38 include supporting a plurality of core pins during the injection molding process of the shaft 34 to provide a uniform part thickness, and al-

lowing the insufflation gas to transfer from the inner diameter of the cannula and seal housing into the inner diameter of the insufflating optical trocar 30 and consequently out of the vent holes 36 at tip 32.

[0018] Referring to FIGS. 4(a) and 4(b), there are shown side cross-sectional views of an example of an insufflating optical trocar 40 of a surgical instrument of the invention. The insufflating optical trocar 40 includes a shaft 34b having a lumen extending substantially along an axis between a proximal end and a distal end, and a blunt tip 32b disposed at the distal end. The insufflating optical trocar 40 further includes at least one gas channel 42 extending along the length of the shaft 34b to provide rapid gas transfer through the insufflating optical trocar 40 when a scope has been placed within the inner diameter. It is appreciated that there may be more than one gas channel 42 extending along the length of the shaft 34b to provide rapid gas transfer through the trocar 40. It is further appreciated that the gas channel 42 may be formed as a separate channel or as the same channel for inserting the scope, that is, by increasing the inner diameter of the shaft 34b to be bigger than the diameter of the scope. More specifically, even if the scope and gas share the same channel, the gas channel 42 assures that there is sufficient cross-sectional area for the gas to travel along the side of the scope and down to the vent hole(s) 36b even when the scope is in place.

[0019] The tip 32b may further include a marker 46 to be used as a visible reference point. The marker 46, together with an area indicated by reference number 44 as shown in FIG. 4A, depict the down vent of the insufflating optical trocar 40 that transfers the gas from the gas channel 42 of the trocar 40 to the vent holes 36b at tip 32b. For example, as the trocar 40 is being placed through the abdominal wall 15, at some point the tip 32b of the trocar 40 will penetrate the peritoneum. Once the peritoneum can be seen through the tip 32b and once the peritoneum is above the visible marker 46, the insufflation gas can be turned on and insufflation can begin. As such, this marks the point where the vent hole 36b is within the abdominal cavity. Once the insufflation gas has created sufficient space between the abdominal wall and the organ bed, the remainder of the insufflating optical trocar 40 including the cannula system can then be fully inserted to an operative position.

[0020] The insufflating optical trocar 40 may further include a scope stop 48 as illustrated in FIG. 4(b) to keep a scope from being inserted into the taper of the inner diameter of the trocar 40. The scope stop 48 may further include a ledge that further prevent the possibility of the scope from being inserted too far into the trocar and consequently block the distal portion of the gas channel 42.

[0021] Referring to FIG. 5, there is shown a perspective view of a septum seal 50 to be used with the insufflating optical trocar of the invention. The septum seal 50 includes a tubular body 52, a septum ring 53 and a plurality of leaflets 54 formed by a slit 56 providing an instrument seal when a scope is inserted into the insufflating optical

trocar and a zero seal when the scope is withdrawn from the trocar. In addition, the thickness of the leaflets 54 can be controlled such that a pressure release mechanism can be created and consequently allowing the leaflets 54 to invert and release pressure if the abdominal pressure within the patient undergoes a sudden spike. Referring to FIGS. 6(a)-6(d), there are shown cross-sectional views of the septum seal 50 of FIG. 5. The septum seal 50 further includes a retaining ledge 58, which allows the septum seal 50 to be attached to the cap or handle 35 of the insufflating optical trocar, and also serves as a sealing surface. Reference number 59 illustrates a sealing surface between the septum seal 50 and the cap or handle of the insufflating optical trocar. The septum seal 50 may further comprise a duckbill or double duckbill valve placed distally of the leaflets 54 to further limit gas or fluid escape.

[0022] During use, the insufflating optical trocar 30 is first inserted into a seal housing 84 and cannula 70 as illustrated in FIG. 7(a). A conventional laparoscope 72 is then inserted into the proximal end of the insufflating optical trocar 30 and advanced to the distal end of the trocar 30. An endoscopic video camera is attached to the proximal end of the laparoscope 72. The trocar 30 is then axially advanced by the surgeon through the body wall 15. As the surgeon advances the cannula 70 and trocar 30 through the body wall 15, the surgeon can visually observe tissue of the body wall 15 as it is being separated via a video monitor, which is connected to the endoscopic video camera. The surgeon can also readily determine when the body wall 15 has been completely traversed by the trocar 30. Once the trocar 30 has traversed the body wall 15, the trocar 30 and laparoscope 72 may be removed which leaves the cannula 70 disposed across the body wall 15 to provide an access channel into the body cavity 25 for the insertion of laparoscopic instrumentation.

[0023] As illustrated in FIG. 7(a), the insufflating optical trocar 30 is designed for use with the seal housing 84 and cannula 70 of the invention. The tip 32 may be blunt and not include any sharp edges, piercing points or blades. The tip 32 of the bladeless insufflating optical trocar 30 may be transparent and generally hollow. This enables a clear view through the distal tip of the insufflating optical trocar 30 and increases the visibility of tissue as it is being traversed. The obturator shaft 34 with the integral tip 32 may be formed of a transparent material such as polycarbonate. The septum seal 50, which may be formed of a material such as Kraton, silicone and the like, may be snap fitted onto the proximal end of the obturator shaft 34. The seal housing 84 may further include a handle attachment 35 including a cannula seal, which may be formed of a plastic material such as polycarbonate, that operates to attach the trocar 30 to the cannula 70 so as to maintain axial position during insertion. The diameter of the shaft 34 can range from about 2 mm to 50 mm and is designed to fit within the seal housing 84 and cannula 70.

[0024] Referring to FIG. 7(b), the cannula 70 is designed to releasably attach to the seal housing 84 via cannula seal 80a, 80b. As the shaft 34 is inserted into the seal housing 84 and cannula 70, the cannula seal 80a, 80b passively engages the seal housing 84 and serves to axially lock the shaft 34 to the seal housing 84 and cannula 70. To release the shaft 34 from the seal housing 84 and cannula 70, outboard tabs on shaft 34 are depressed inwardly and the shaft 34 is then free to be slidably removed. The shaft 34 includes axial key members 37 (see FIG. 3) at its proximal end which are designed to mate with axial keyways on the seal housing 84. As the shaft 34 is inserted into the seal housing 84 and cannula 70, the shaft 34 is rotated slightly to align key members 37 with the keyways and then advanced until the cannula seal 80a, 80b engages the seal housing 84. The key members 37 serve to rotationally index the shaft 34 to the seal housing 84. In another aspect, cannula 70 may further include distal cannula seal 82 formed at a distal portion of cannula 70 and shaft 34 so as to further limit gas or fluid escape.

[0025] The insufflating optical trocar 30 may include a laparoscope lock 86 having an elastomeric element. The addition of the elastomeric element would enhance the frictional engagement with the laparoscope 72. An example of an elastomeric element would be a silicone O-ring sized with an inside diameter smaller than the outside diameter of the laparoscope 72. The laparoscope lock 86 could either rotate freely to enable the laparoscope 72 to rotate freely relative to the shaft 34 or the laparoscope lock 86 could be rotationally fixed to prevent the laparoscope 72 from rotating relative to the shaft 34.

[0026] In use, first the skin around the area to be operated on is incised appropriately for the size of the cannula 70. An insufflation gas line 90, which is attached to the seal housing 84, the insufflating optical trocar 30 and the laparoscope 72 are then inserted into the cannula 70. At this point the gas supply is still turned off. The assembled device is then advanced through the body or abdominal wall 15 under direct vision until it is observed that just the tip 32 of the device has penetrated the peritoneal cavity. The device is then held in place and the flow of insufflation gas is begun. The gas will flow through the tip 32 and into the peritoneal cavity until the cavity is sufficiently distended by gas pressure. The surgeon then completes the insertion of the insufflating optical trocar 30 until the cannula 70 is in an appropriate or desired position. The insufflating optical trocar 30 and laparoscope 72 may then be removed. At this point, the surgeon may elect to reinsert just the laparoscope 72 through the seal housing 84 and thereby allow observation of the abdominal cavity and subsequent insertions of additional laparoscopic instrumentation.

[0027] As explained earlier, an indicator line or marker 46 as shown in FIG. 4(a) may be located on tip 32 to be viewed by laparoscope 72 to indicate when the device has advanced far enough into the body cavity to begin insufflation. The coincidence of anatomical features with

the indicator line or marker 46 may indicate the correct position to begin insufflation. The indicator line or marker 46 could be circumferential in nature and when the peritoneal layer, as it is being penetrated, forms a coincident circle with respect to the indicator line 46, the surgeon can begin insufflating. Another method is to employ an O-ring seal 86. Additionally, it is preferred that a zero seal be present on the trocar to prevent escape of the gas when the trocar is used to place cannulas without the laparoscope 72. A double duckbill valve 88 would work well too for this application as would a single duckbill, a flapper valve or a slit valve.

[0028] It is appreciated that the above-described concept may be applied to any surgical instruments providing visual entry and visual insufflation, regardless of size or type of fluid transfer as further described in the following exemplary embodiments of the invention. x

[0029] Referring to FIGS. 9(a)-9(e), there are shown additional tip designs 32o-32s to facilitate penetration of a body tissue. FIG. 9(a) illustrates a spoon-shaped or asymmetric tip 32o having at least one vent hole 36o; FIG. 9(b) illustrates a generally domed or conical shaped tip 32p having plastic or metal blades 33p along an axis of the shaft and at least one vent hole 36p; FIG. 9(c) illustrates a blunt tip 32q having at least one vent hole 36q; FIG. 9(d) illustrates a generally domed or conical shaped tip 32r having at least one bladed fin 33r and at least one vent hole 36r; and FIG. 9(e) illustrates a generally conical shaped tip 32s having at least one vent hole 36s at the distal tip. It is appreciated that tips 32o, 32p, 32r and 32s have a sharp, pointed or bladed tip and/or edge to facilitate penetration of body tissue. In yet other embodiments of the invention, the surface of the tip may have at least one tissue engaging raised pattern on the surface. The surface operates to facilitate insertion of the insufflating surgical instrument or optical trocar with a reduced penetration force and minimize tenting of the body wall. The surface may further facilitate separation of different layers of the body wall and provides proper alignment of the tip between the layers. In another embodiment of the invention, the tip may have an outer surface extending distally to a blunt point and includes a pair of side sections separated by an intermediate section, and wherein the side sections extend from the blunt point radially outwardly with progressive positions proximally along the axis. The side sections may include a distal portion in proximity to the blunt point and a proximal portion in proximity to the tubular body, and the distal portion of the side sections being twisted radially with respect to the proximal portion of the side sections.

[0030] Referring to FIGS. 10(a)-10(c), there are shown additional tip designs 32v-32x in accordance to other aspects of the invention. For example, the tip 32v as illustrated in FIG. 10(a) comprises a flip-top 272 and a conical body 270 that operates to move from a first, penetrating position to a second, insufflating position when the body wall has been traversed. The tip 32v may further comprise a retention member for connecting the flip-top 272

and the conical body 270. The retention member may be one of a spring, a spring wire, an offset hinge or a "living" hinge. Other flip-top or flip-tip designs as described in co-pending U.S. Patent Application Serial No. 10/805,864, entitled "Surgical Access Port and Method of Using Same," filed March 22, 2004 may also be used with the insufflating concept of the invention. In yet another aspect of the invention, the tip 32w as illustrated in FIG. 10(b) comprises a two-piece flip-top 282a, 282b that operates to move from a first, penetrating position to a second, insufflating position when the body wall has been traversed. In particular, the tip 32w may comprise at least two or more parts or petals that reposition to the side of the shaft 34w in the second, insufflating position. FIG. 10(c) illustrates the tip 32x in accordance with another aspect of the invention comprising a two-stage flip-top 290 that operates to move from a penetrating position to an insufflating position and then to an instrument access position. In particular, the two-stage flip-top 290 comprises a distal flip portion 292 and a proximal flip portion 294. In the first stage, the distal flip portion 292 moves from a penetrating position to an open or insufflating position once the body wall has been traversed. Once insufflation has been achieved, the proximal flip portion 294 moves to an open or instrument access position in the second stage. The tip 32x may further comprise retention members for connecting between the distal flip portion 292 and the proximal flip portion 294, and between the proximal flip portion 294 and the shaft 34x.

[0031] FIGS. 11(a)-11(c) illustrate insufflating valve vents. More specifically, FIG. 11 (a) illustrates an insufflating valve vent 300 formed at the distal end of the shaft 34. The insufflating valve vent 300 is formed of an elastic material to allow gas such as CO₂ to be introduced from the inside of the shaft 34 to a body cavity. It is appreciated that when there is no gas, the elastic material of the insufflating valve vent 300 causes it to close so as to provide an airtight seal. FIG. 11 (b) illustrates an insufflating flapper valve 310 formed at tip 32y of an insufflating optical trocar. The insufflating flapper valve 310 comprises at least one flapper valve vent 312 that operates to open when a gas such as CO₂ is introduced in the shaft 34. It is appreciated that when there is no gas, the flapper valve vent 312 closes to provide a tight seal. Similarly to FIG. 11 (b), FIG. 11 (c) illustrates an insufflating reverse flapper valve 320 formed at tip 32z of an insufflating optical trocar. The insufflating reverse flapper valve 320 comprises at least one flapper valve vent 322 that remains close or shut by tissue during insertion, and once peritoneum is passed, pressure by a gas such as CO₂ would then open the reverse flapper valve 320 to allow the transfer of the gas into a body cavity. It is appreciated that each of the above flapper valve vents may be spring loaded to operate like a **Veress** needle.

[0032] Referring to FIGS. 12(a) and 12(b), there are shown illustrations of an alternative insufflating surgical instrument 500 in accordance with the invention. The insufflating surgical instrument 500 comprises an insufflat-

ing optical trocar 502 and a cannula 520. The insufflating optical trocar 502 comprises a shaft 504 having a lumen extending along an axis between a proximal end and a distal end, a tip 506 disposed at a distal end of the shaft, at least one vent hole 508 to introduce gas from the cannula 520 into the body or abdominal cavity as further discussed below, and a gas channel 510 formed in the shaft 504 and operably connected to the at least one vent hole 508 to allow gas transfer from the cannula 520 to the insufflating optical trocar 502. The cannula 520 comprises at least one cannula gas channel 522 extending along its longitudinal axis to transfer gas to the trocar gas channel 510 after insertion of the insufflating optical trocar 502 into the cannula 520. In other words, the cannula gas channel 522 is encased as a lumen in the cannula wall. During operation, the transfer of gas only takes place if there is an alignment between the cannula gas channel 522 and the trocar gas channel 510 as illustrated in FIGS. 12(a) and 12(b). A scope 525 may be inserted at the proximal end of the insufflating optical trocar 502 and then advanced to the distal end of the trocar 502 as the trocar 502 is placed through an abdominal wall.

[0033] Referring to FIGS. 13(a)-13(i), there are shown additional tip designs 32aa-32ii. Each of these tip designs includes at least one vent hole (36aa-36ii) at the distal tip to introduce insufflation gas into a body cavity. It is appreciated that some of these tips have a sharp, pointed or bladed tip and/or edge to facilitate penetration of body tissue.

[0034] It is appreciated that the above described surgical instruments and devices can be used to access not only the peritoneal cavity but can be used for preperitoneal hernia repair, retroperitoneal operations including back and kidney operations, percutaneous kidney operations, thoracic surgery and arthroscopic access. In addition to gas such as carbon dioxide, it is appreciated that other fluids such as air, water and saline can also be introduced into a body cavity with the technique of the invention. It is appreciated that operating scopes may be modified such that a lumen may be used to introduce insufflation fluid. Accordingly, it is understood that many other modifications can be made to the various disclosed embodiments without departing from the scope of the invention. For these reasons, the above description should not be construed as limiting the invention, but should be interpreted as merely exemplary embodiments.

Claims

1. An insufflating surgical instrument adapted for movement across an abdominal wall to insufflate an abdominal region of a patient, comprising:

a cannula (70, 520) comprising a wall forming a first lumen extending along an axis between a proximal end and a distal end and adapted for

connection to a source of fluid under pressure at the proximal end; and

a seal housing (84) releasably attached to the proximal end of the cannula via a cannula seal (80a, 80b) that passively engages the seal housing (84), the seal housing having axial keyways; a trocar (30, 40, 250, 502) comprising:

a shaft (34, 504) arranged such that it may be inserted into the seal housing (84) and cannula (70, 520), the shaft (34, 504) having outboard tabs at a proximal end and an insufflation channel (42, 224, 430, 510) extending along an axis between a proximal end and a distal end, the insufflation channel (42, 224, 430, 510) being operably connected to allow fluid transfer from the cannula (70, 520) to the trocar (30, 40, 250, 502);

a tip (32, 194, 506) at the distal end of the shaft (34, 504);

at least one vent hole (36, 196, 216, 236, 508) formed at the tip (32, 194, 506) or along the shaft (34, 504), the vent hole (36, 196, 216, 236, 508) being in connection with the insufflation channel (42, 224, 430, 510) and being adapted to expel the fluid under pressure to insufflate the abdominal region; and axial key members (37) configured to mate with the axial keyways on the seal housing, the axial key members being designed to rotationally index the shaft (34) to the seal housing (84),

wherein, in an assembled operational configuration, the trocar (30, 40, 250, 502) is arranged to be insertable into and removable from the first lumen of the cannula (70, 520) wherein: as the shaft (34) is inserted into the seal housing (84) and cannula (70) the shaft (34) is arranged to be axially locked to the seal housing (84) and cannula (70); the shaft is arranged (34) to be released from the seal housing (84) by depressing the outboard tabs on the shaft (34); the trocar (30, 40, 250, 502) is longer than the cannula (70, 520); and the instrument is configured to permit a user to apply a force to the shaft to cause the tip to penetrate tissue ahead of the tip of the cannula (70, 520).

2. The insufflating surgical instrument of claim 1, wherein at least one of the tip (32, 194, 506) and the shaft (34, 504) is formed of a transparent material to facilitate visualization of the abdominal wall and the abdominal region.

3. The insufflating surgical instrument of claim 1,

wherein the insufflation channel is operably connected to allow fluid transfer from the cannula (70) to the trocar (30) by at least one shaft vent (38) in the shaft (34).

4. The insufflating surgical instrument of claim 1, wherein the shaft (34, 504) has a lumen extending along the axis between the proximal end and the distal end to enable insertion of the laparoscope.
5. The insufflating surgical instrument of claim 1, wherein the shaft (34) is arranged to be rotated slightly to align key members (37) with the keyways.
6. The insufflating surgical instrument of claim 4, wherein the lumen of the shaft is also the insufflation channel.
7. The insufflating surgical instrument of claim 1 wherein the trocar (30) includes a laparoscope lock (86) having an elastomeric element configured to enhance frictional engagement with a laparoscope.
8. The insufflating surgical instrument of claim 7, wherein the laparoscope lock (86) is a silicone O-ring sized with an inside diameter smaller than the outside diameter of a laparoscope.
9. The insufflating surgical instrument of claim 7, wherein the laparoscope lock (86) is rotationally fixed to prevent the laparoscope from rotating relative to the shaft (34, 504).
10. The insufflating surgical instrument of claim 1, further comprising at least one second vent hole (36, 196, 216, 236, 508) being in connection with the insufflation channel (42, 224, 430, 510).
11. The insufflating surgical instrument of claim 1, wherein the blunt tip (32, 194, 506) further comprises a marker (46) to indicate when the vent hole (36, 196, 216, 236, 508) is positioned for insufflation.
12. The insufflating surgical instrument of claim 11, wherein the marker (46) indicates the point where the vent hole (36, 196, 216, 236, 508) has penetrated the abdominal wall.
13. The insufflating surgical instrument of claim 1, wherein the seal housing (84) comprises a septum seal (50) and a plurality of leaflets (54) forming an instrument seal in the presence of a laparoscope, and providing a zero seal in the absence of an instrument.
14. The insufflating surgical instrument of claim 1, wherein the shaft (34) has a diameter ranging from about 2 mm to about 5 mm.

15. An insufflating surgical instrument of claim 1 further including a second insufflation channel (522) formed in the wall of the cannula extending along the axis between the proximal end and the distal end and being adapted for connection to the source of fluid under pressure at the proximal end; the insufflation channel (510) at the distal end of the shaft (504) being adapted for alignment with the second insufflation channel (522) of the cannula (520) to allow the fluid under pressure to be expelled from the at least one vent hole (508) to insufflate the abdominal region.

16. The insufflating surgical instrument of claim 3 wherein the insufflation channel (42) is configured such that fluid under pressure flows from the cannula (70), through the at least one shaft vent (38) in the shaft, into the trocar, along at least a portion of the insufflation channel (42) through the at least one vent hole (36) at the distal end of the tip (32) or along the shaft and out of the trocar (30) to insufflate an abdominal region of the patient.

Patentansprüche

1. Ein chirurgisches Insufflationsinstrument, das zur Bewegung über eine Bauchwand hinweg angepasst ist, um eine Bauchregion eines Patienten aufzublasen, das Folgendes beinhaltet:

eine Kanüle (70, 520), die eine Wand beinhaltet, die ein erstes Lumen bildet, das sich entlang einer Achse zwischen einem proximalen Ende und einem distalen Ende erstreckt und zum Anschluss an eine Fluidquelle unter Druck an dem proximalen Ende angepasst ist; und ein Dichtungsgehäuse (84), das lösbar an dem proximalen Ende der Kanüle über eine Kanülendichtung (80a, 80b) befestigt ist, die passiv Eingriff mit dem Dichtungsgehäuse (84) nimmt, wobei das Dichtungsgehäuse axiale Keilnuten aufweist; einen Trokar (30, 40, 250, 502), der Folgendes beinhaltet:

einen Schaft (34, 504), der so angeordnet ist, dass er in das Dichtungsgehäuse (84) und die Kanüle (70, 520) eingeführt werden kann, wobei der Schaft (34, 504) Außenklappen an einem proximalen Ende aufweist, und einen Insufflationskanal (42, 224, 430, 510) der sich entlang einer Achse zwischen einem proximalen Ende und einem distalen Ende erstreckt, wobei der Insufflationskanal (42, 224, 430, 510) betriebsfähig verbunden ist, um den Fluidtransfer von der Kanüle (70, 520) zum Trokar (30, 40, 250,

- 502) zu erlauben;
 eine Spitze (32, 194, 506) an dem distalen Ende des Schafts (34, 504);
 mindestens ein Entlüftungsloch (36, 196, 216, 236, 508), das an der Spitze (32, 194, 506) oder entlang des Schafts (34, 504) ausgebildet ist, wobei das Entlüftungsloch (36, 196, 216, 236, 508) in Verbindung mit dem Insufflationskanal (42, 224, 430, 510) steht und zum Ausstoßen des Fluids unter Druck angepasst ist, um die Bauchregion aufzublasen; und
 axiale Keilelemente (37), die dazu konfiguriert sind, mit den axialen Keilnuten auf dem Dichtungsgehäuse zusammenzupassen, wobei die axialen Keilelemente zur Rotationsindexierung des Schafts (34) zum Dichtungsgehäuse (84) konzipiert sind, wobei der Trokar (30, 40, 250, 502) in einer zusammengebauten betriebsfähigen Konfiguration so angeordnet ist, dass er in das erste Lumen der Kanüle (70, 520) eingeführt und aus diesem entfernt werden kann, wobei: wenn der Schaft (34) in das Dichtungsgehäuse (84) und die Kanüle (70) eingeführt wird, der Schaft (34) dazu angeordnet ist, axial an dem Dichtungsgehäuse (84) und der Kanüle (70) arretiert zu werden; der Schaft dazu angeordnet (34) ist, von dem Dichtungsgehäuse (84) gelöst zu werden, indem die Außenklappen auf dem Schaft (34) nach unten gedrückt werden; der Trokar (30, 40, 250, 502) länger als die Kanüle (70, 520) ist; und das Instrument so konfiguriert ist, dass es einem Benutzer gestattet, eine Kraft auf den Schaft anzuwenden, um die Spitze zu veranlassen, vor der Spitze der Kanüle (70, 520) liegendes Gewebe zu penetrieren.
2. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei mindestens eines von der Spitze (32, 194, 506) und dem Schaft (34, 504) aus einem transparenten Material gebildet ist, um die Sichtbarmachung der Bauchwand und der Bauchregion zu erleichtern.
 3. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei der Insufflationskanal betriebsfähig verbunden ist, um den Fluidtransfer von der Kanüle (70) zum Trokar (30) durch mindestens eine Schaftentlüftung (38) in dem Schaft (34) zu erlauben.
 4. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei der Schaft (34, 504) ein Lumen aufweist, das sich entlang der Achse zwischen dem proximalen Ende und dem distalen Ende erstreckt, um das Einführen des Laparoscops zu ermöglichen.
 5. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei der Schaft (34) dazu angeordnet ist, geringfügig rotiert zu werden, um die Keilelemente (37) mit den Keilnuten auszurichten.
 6. Das chirurgische Insufflationsinstrument gemäß Anspruch 4, wobei das Lumen des Schafts auch der Insufflationskanal ist.
 7. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei der Trokar (30) eine Laparoskoparretierung (86) einschließt, die ein elastomeres Element aufweist, das dazu konfiguriert ist, den Reibeingriff mit einem Laparoskop zu verbessern.
 8. Das chirurgische Insufflationsinstrument gemäß Anspruch 7, wobei die Laparoskoparretierung (86) ein Silikon-O-Ring ist, der mit einem Innendurchmesser dimensioniert ist, der kleiner ist als der Außendurchmesser eines Laparoscops.
 9. Das chirurgische Insufflationsinstrument gemäß Anspruch 7, wobei die Laparoskoparretierung (86) rotational fixiert ist, um ein Rotieren des Laparoscops im Verhältnis zum Schaft (34, 504) zu verhindern.
 10. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, das ferner mindestens ein zweites Entlüftungsloch (36, 196, 216, 236, 508) beinhaltet, das in Verbindung mit dem Insufflationskanal (42, 224, 430, 510) steht.
 11. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei die abgestumpfte Spitze (32, 194, 506) ferner einen Marker (46) beinhaltet, um anzuzeigen, wann das Entlüftungsloch (36, 196, 216, 236, 508) zum Aufblasen positioniert ist.
 12. Das chirurgische Insufflationsinstrument gemäß Anspruch 11, wobei der Marker (46) den Punkt anzeigt, an dem das Entlüftungsloch (36, 196, 216, 236, 508) die Bauchwand penetriert hat.
 13. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei das Dichtungsgehäuse (84) eine Trennwanddichtung (50) und eine Vielzahl von Segelklappen (54) beinhaltet, die eine Instrumentendichtung bilden, wenn ein Laparoskop vorhanden ist, und eine Nulldichtung bereitstellen, wenn kein Instrument vorhanden ist.
 14. Das chirurgische Insufflationsinstrument gemäß Anspruch 1, wobei der Schaft (34) einen Durchmesser im Bereich von etwa 2 mm bis etwa 5 mm aufweist.
 15. Ein chirurgisches Insufflationsinstrument gemäß Anspruch 1, das ferner einen zweiten Insufflationskanal (522) aufweist, der in der Wand der Kanüle

ausgebildet ist, die sich entlang der Achse zwischen dem proximalen Ende und dem distalen Ende erstreckt und zum Anschluss mit der Fluidquelle unter Druck an dem proximalen Ende angepasst ist; wobei der Insufflationskanal (510) an dem distalen Ende des Schafts (504) dazu angepasst ist, mit dem zweiten Insufflationskanal (522) der Kanüle (520) ausgerichtet zu werden, um das Ausstoßen von Fluid unter Druck aus dem mindestens einen Entlüftungsloch (508) zu erlauben, um die Bauchregion aufzublasen.

16. Das chirurgische Insufflationsinstrument gemäß Anspruch 3, wobei der Insufflationskanal (42) so konfiguriert ist, dass Fluid unter Druck aus der Kanüle (70), durch die mindestens eine Schaftentlüftung (38) in den Schaft, in den Trokar, entlang mindestens eines Teils des Insufflationskanals (42) durch das mindestens eine Entlüftungsloch (36) an dem distalen Ende der Spitze (32) oder entlang des Schafts und aus dem Trokar (30) heraus strömt, um eine Bauchregion des Patienten aufzublasen.

Revendications

1. Un instrument chirurgical d'insufflation adapté afin de se déplacer au travers de la paroi abdominale pour insuffler une région abdominale d'un patient comprenant :

une canule (70, 520) comprenant une paroi formant une première lumière s'étendant le long d'un axe entre une extrémité proximale et une extrémité distale et adaptée pour être reliée à une source de fluide sous pression au niveau de l'extrémité proximale ; et

un boîtier d'étanchéité (84) fixé de manière détachable à l'extrémité proximale de la canule par l'intermédiaire d'un joint de canule (80a, 80b) qui entre en contact passivement avec le boîtier d'étanchéité (84), ce dernier pourvu de rainures axiales ;

un trocart (30, 40, 250, 502) comprenant :

une tige (34, 504) disposée de manière à ce qu'elle puisse être insérée à l'intérieur du boîtier d'étanchéité (84) et de la canule (70, 520), la tige (34, 504) étant pourvue de languettes extérieures au niveau d'une extrémité proximale et un canal d'insufflation (42, 224, 430, 510) s'étendant le long d'un axe entre une extrémité proximale et une extrémité distale, le canal d'insufflation (42, 224, 430, 510) étant relié de manière fonctionnelle pour permettre un transfert de fluide depuis la canule (70, 520) vers le trocart (30, 40, 250, 502) ;

une pointe (34, 194, 506) au niveau de l'extrémité distale de la tige (34, 504) ; au moins un trou d'aération (36, 196, 216, 236, 508) formé au niveau de la pointe (32, 194, 506) ou le long de la tige (34, 504), le trou d'aération (36, 196, 216, 236, 508) étant relié au canal d'insufflation (42, 224, 430, 510) et étant adapté pour expulser le fluide sous pression afin d'insuffler la région abdominale ; et

des éléments de clavette axiale (37) conçus pour s'adapter aux rainures axiales du boîtier d'étanchéité, les éléments de clavette axiale étant conçus pour indexer en rotation la tige (34) au boîtier d'étanchéité (84), dans lequel, dans une configuration fonctionnelle assemblée, le trocart (30, 40, 250, 502) est disposé pour pouvoir être inséré à l'intérieur et retiré de la première lumière de la

canule (70, 520) dans laquelle: lorsque la tige (34) est insérée à l'intérieur du boîtier d'étanchéité (84) et de la canule (70), la tige (34) est disposée pour être verrouillée de manière axiale dans le boîtier d'étanchéité (84) et à la canule (70) ; la tige (34) est disposée pour être détachée du boîtier d'étanchéité (84) en enfonçant les languettes extérieures sur la tige (34) ; le trocart (30, 40, 250, 502) est plus long que la canule (70, 520) ; et l'instrument est conçu pour permettre à un utilisateur d'exercer une force sur la tige afin d'amener la pointe à pénétrer le tissu situé devant la pointe de la canule (70, 520).

2. L'instrument chirurgical d'insufflation de la revendication 1, dans lequel au moins l'une des pointes (32, 194, 506) et la tige (34, 504) sont constituées d'un matériau transparent afin de faciliter la visualisation de la paroi abdominale et de la région abdominale.

3. L'instrument chirurgical d'insufflation de la revendication 1, dans lequel le canal d'insufflation est relié de manière fonctionnelle afin de permettre un transfert de fluide depuis la canule (70) vers le trocart (30) au moyen d'au moins un conduit d'air (38) dans la tige (34).

4. L'instrument d'insufflation de la revendication 1, dans lequel la tige (34, 504) est dotée d'une lumière s'étendant le long d'un axe entre l'extrémité proximale et l'extrémité distale pour permettre l'insertion d'un laparoscope.

5. L'instrument chirurgical d'insufflation de la revendication 1, dans lequel la tige (34) peut être légèrement tournée afin d'aligner les éléments de clavette (37)

- avec les rainures.
6. L'instrument d'insufflation de la revendication 4, dans lequel la lumière de la tige est également le canal d'insufflation. 5
7. L'instrument chirurgical d'insufflation de la revendication 1 dans lequel le trocart (30) comprend un dispositif de verrouillage de laparoscope (86) doté d'un élément élastomère conçu pour améliorer le contact par friction avec un laparoscope. 10
8. L'instrument chirurgical d'insufflation de la revendication 7, dans lequel le dispositif de verrouillage de laparoscope (86) est un joint torique en silicium dimensionné pour avoir un diamètre intérieur plus petit que le diamètre extérieur d'un laparoscope. 15
9. L'instrument chirurgical d'insufflation de la revendication 7, dans lequel le dispositif de verrouillage de laparoscope (86) est fixé en rotation afin d'empêcher le laparoscope de tourner par rapport à la tige (34, 504). 20
10. L'instrument chirurgical d'insufflation de la revendication 1, comprenant en outre au moins un second trou d'aération (36, 196, 216, 236, 508) étant relié au canal d'insufflation (42, 224, 430, 510). 25
11. L'instrument chirurgical d'insufflation de la revendication 1, dans lequel la pointe émoussée (32, 194, 506) comprend en outre un repère (46) indiquant si le trou d'aération (36, 196, 216, 236, 508) est positionné pour l'insufflation. 30
- 35
12. L'instrument chirurgical d'insufflation de la revendication 11, dans lequel le repère (46) indique le point où le trou d'aération (36, 196, 216, 236, 508) a pénétré la paroi abdominale. 40
- 45
13. L'instrument chirurgical d'insufflation de la revendication 1, dans lequel le boîtier d'étanchéité (84) comprend un joint septum (50) et une pluralité d'ailettes (54) formant un joint d'instrument en présence du laparoscope, et fournissant un joint invisible en l'absence d'instrument. 45
- 50
14. L'instrument chirurgical d'insufflation de la revendication 1, dans lequel la tige (34) a un diamètre compris entre 2 mm et 5 mm environ. 50
- 55
15. Un instrument chirurgical d'insufflation de la revendication 1 comprenant en outre un second canal d'insufflation (522) formé dans la paroi de la canule s'étendant le long de l'axe entre l'extrémité proximale et l'extrémité distale et étant adapté pour être relié à la source de fluide sous pression au niveau de l'extrémité proximale ;
- le canal d'insufflation (510) au niveau de l'extrémité distale de la tige (504) étant adapté pour s'aligner avec le second canal d'insufflation (522) de la canule (520) afin de permettre au fluide sous pression d'être expulsé à partir d'au moins un trou d'aération (508) pour insuffler la région abdominale.
16. L'instrument chirurgical d'insufflation de la revendication 3, dans lequel le canal d'insufflation (42) est conçu de sorte que le fluide sous pression s'écoule à partir de la canule (70), à travers au moins un conduit d'air (38) dans la tige, à l'intérieur du trocart, le long d'au moins une partie du canal d'insufflation (42) à travers au moins un conduit d'air (36) au niveau de l'extrémité distale de la pointe (32) ou le long de la tige et hors du trocart (30) afin d'insuffler une région abdominale du patient.

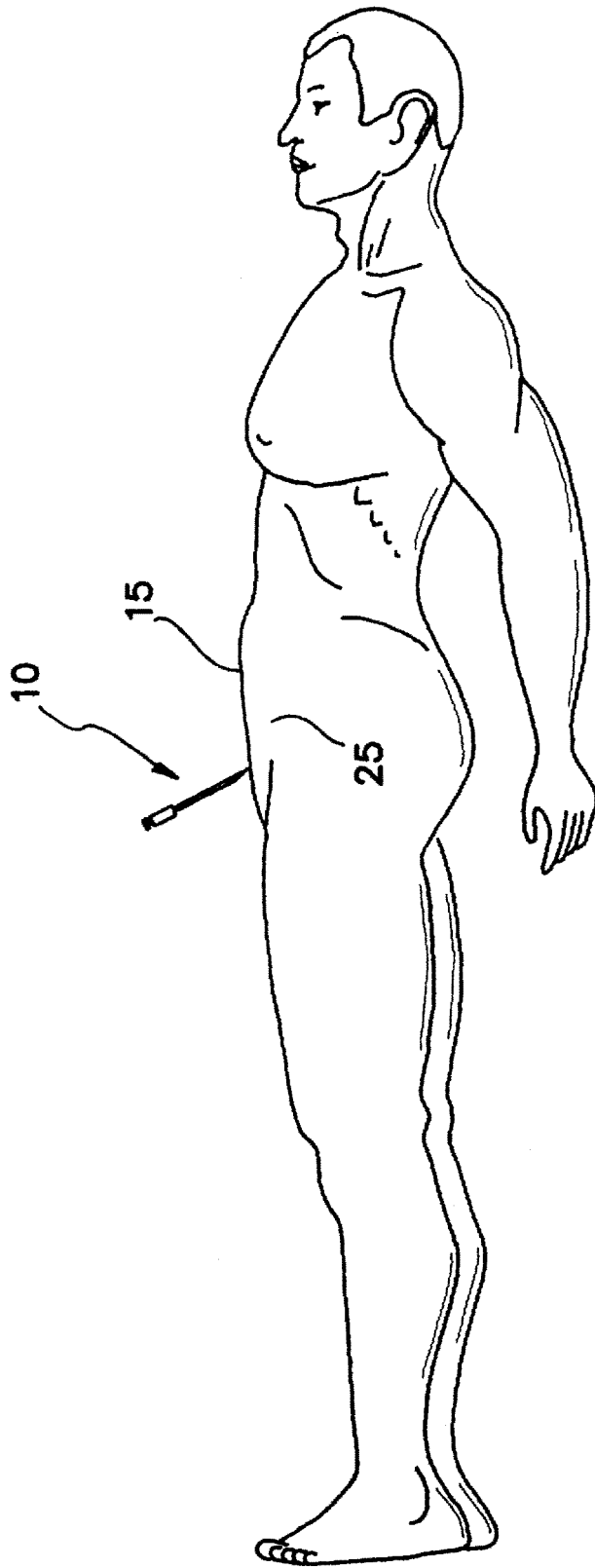


FIG. 1
PRIOR ART

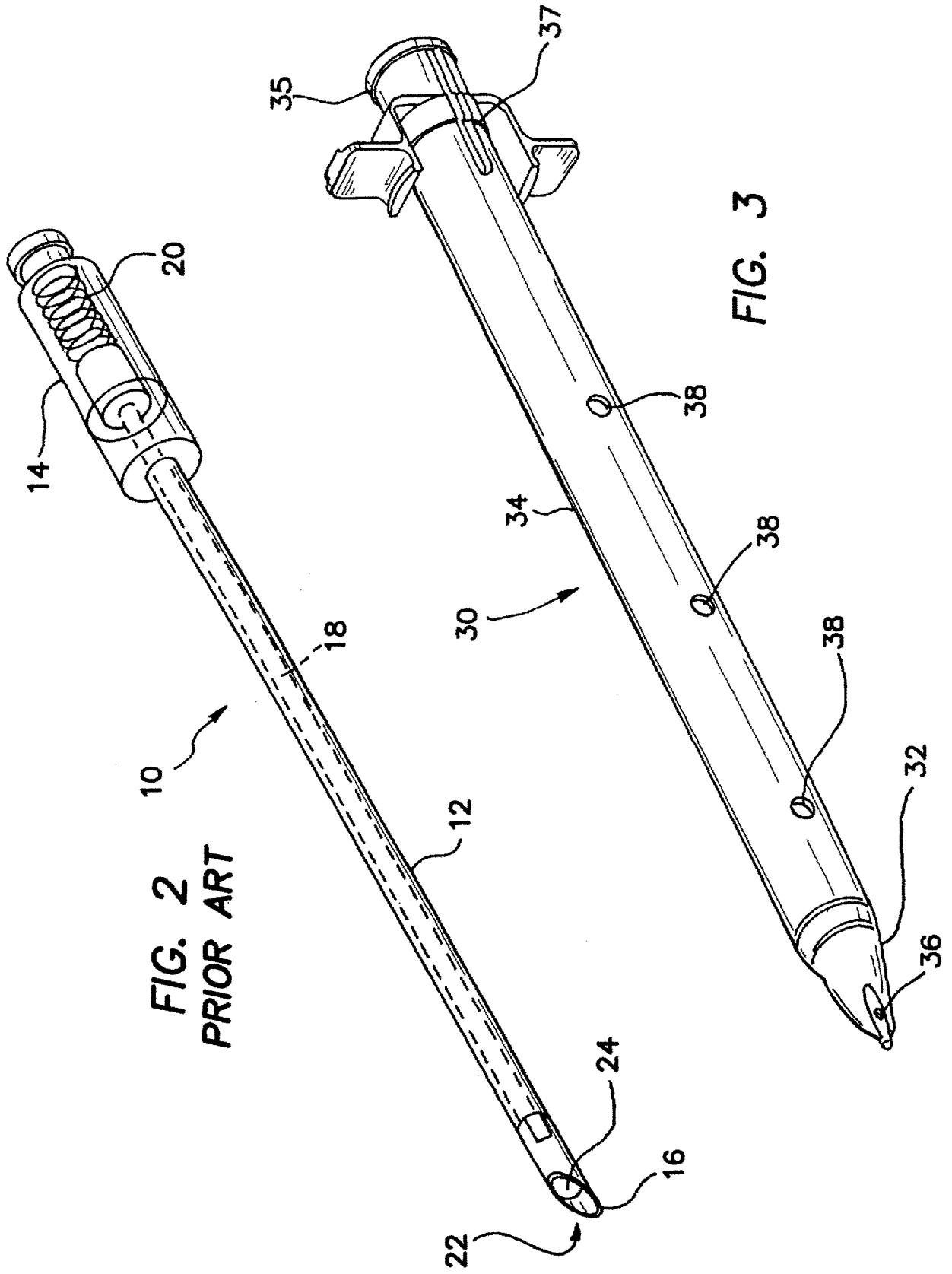
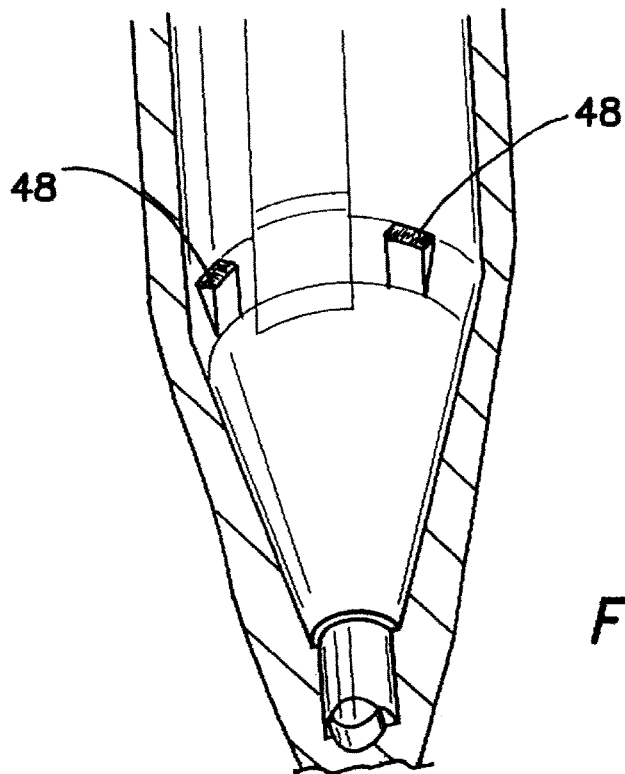
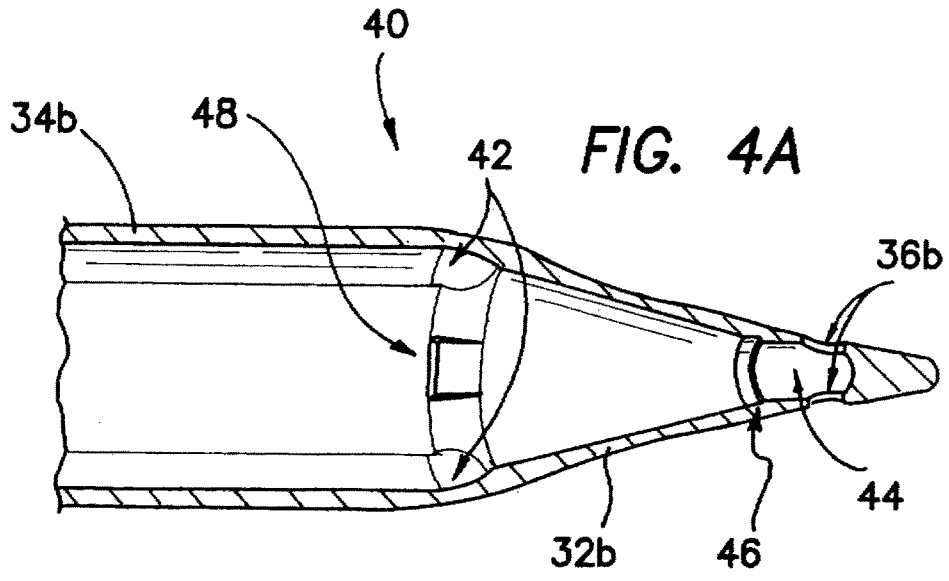


FIG. 2
PRIOR ART

FIG. 3



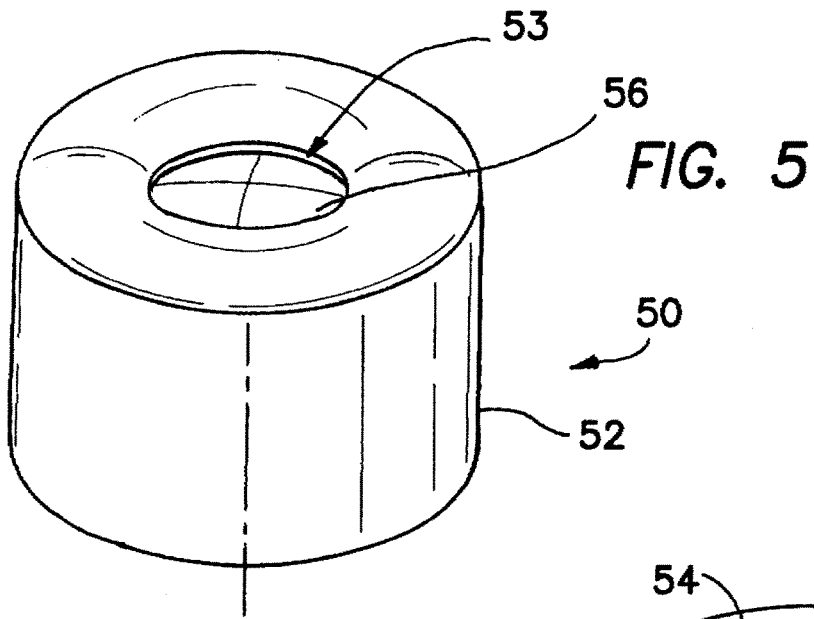
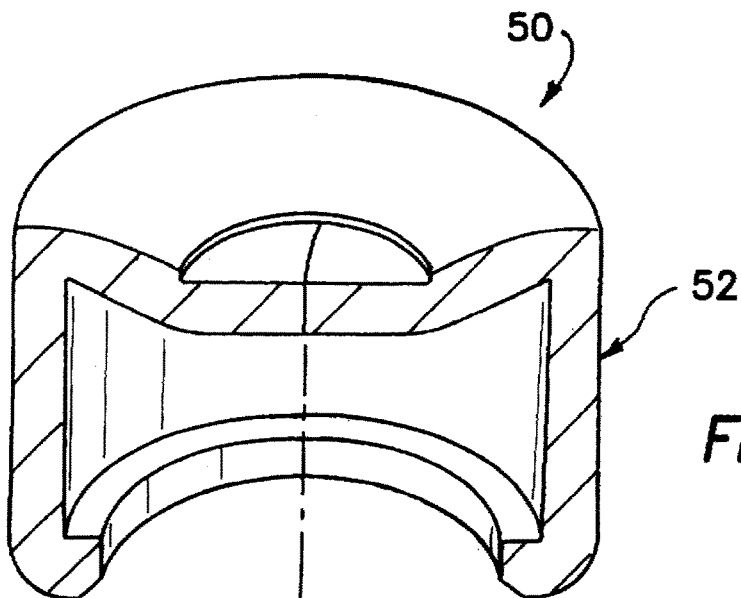
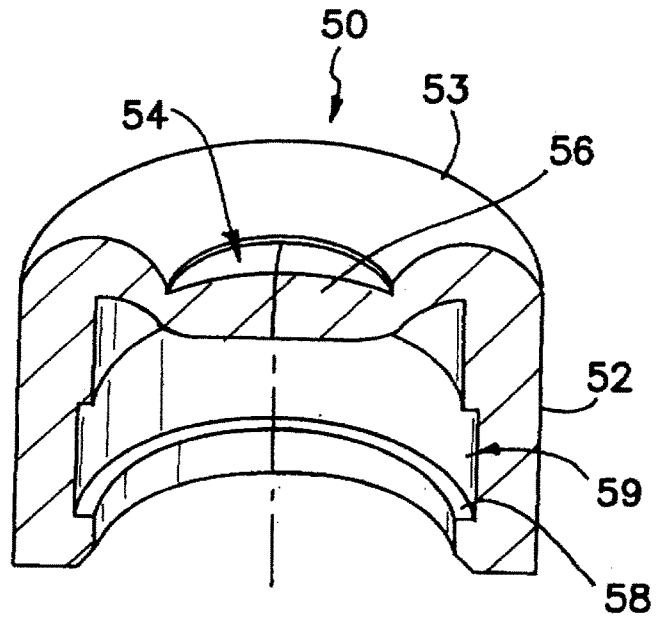


FIG. 6A



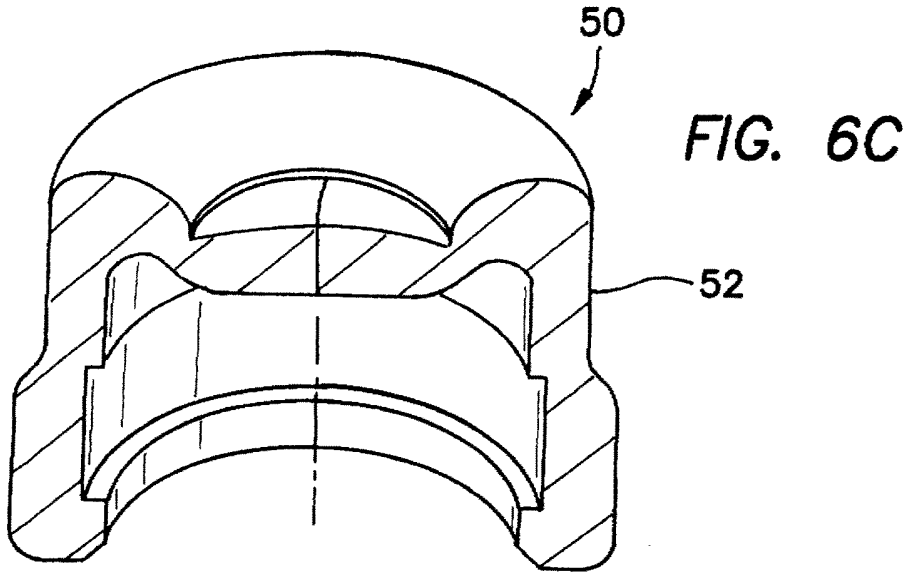


FIG. 6C

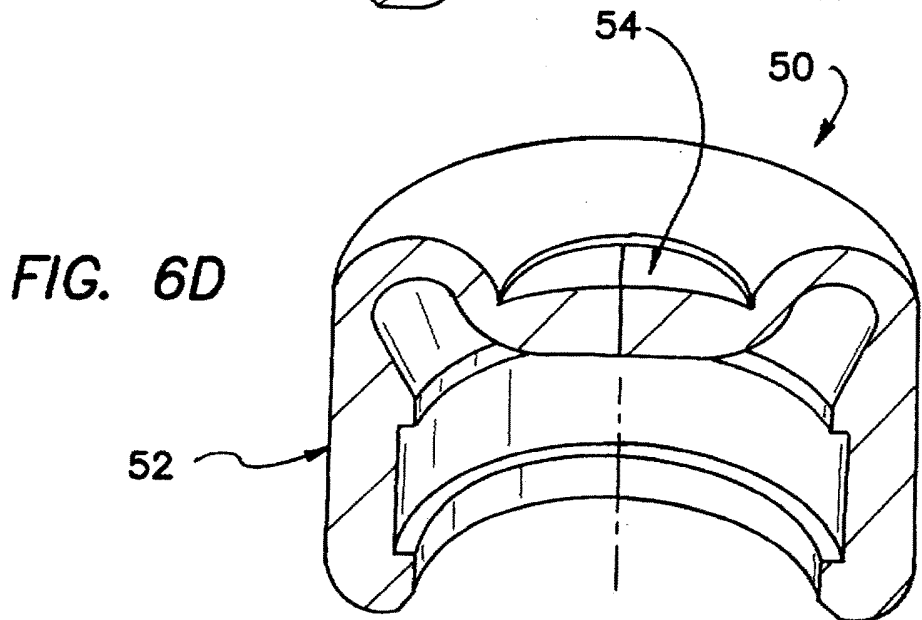


FIG. 6D

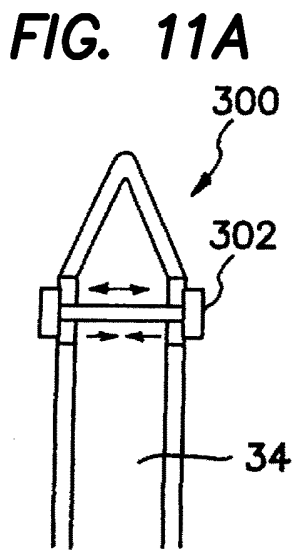


FIG. 11A

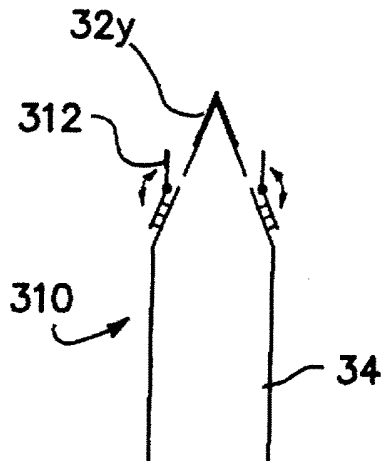


FIG. 11B

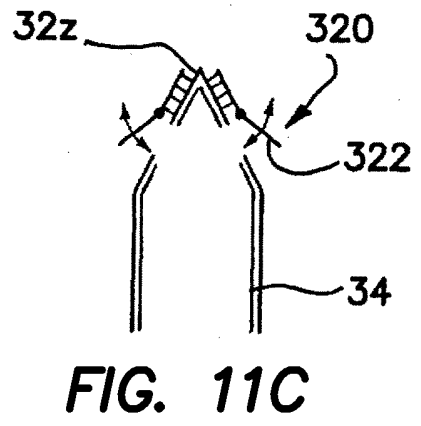


FIG. 11C

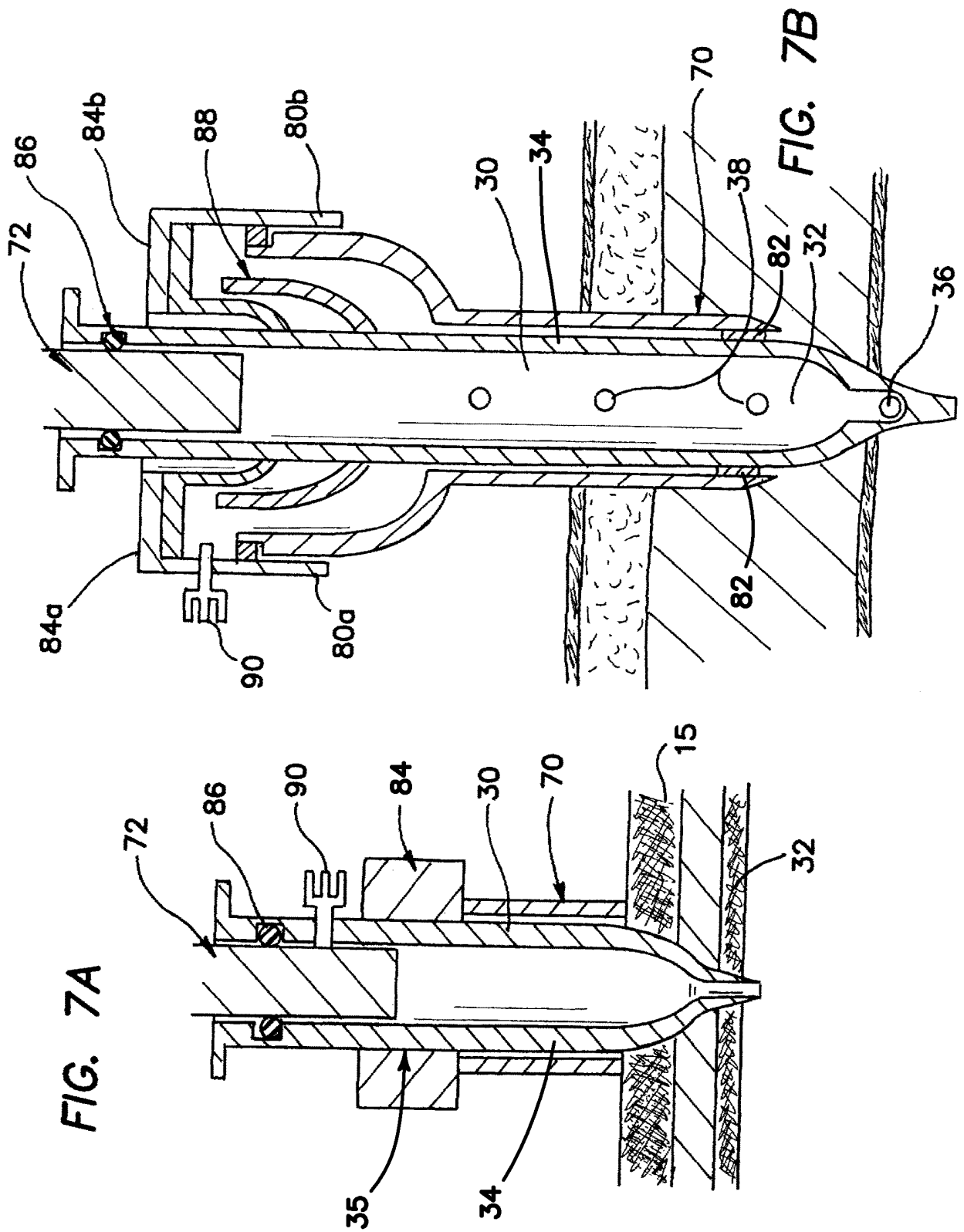


FIG. 7A

FIG. 7B

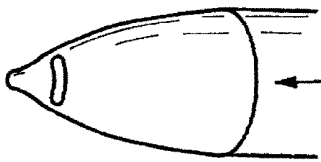


FIG. 8A

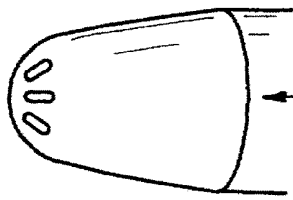


FIG. 8B

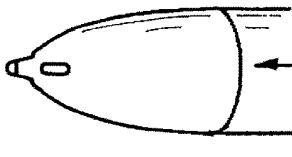


FIG. 8C

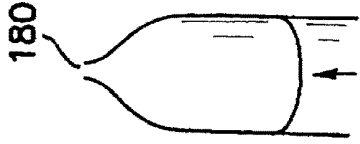


FIG. 8D

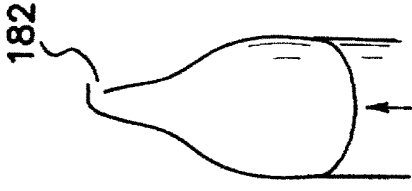


FIG. 8E

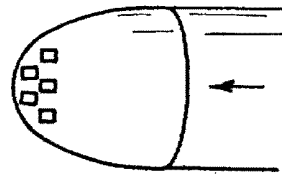


FIG. 8F

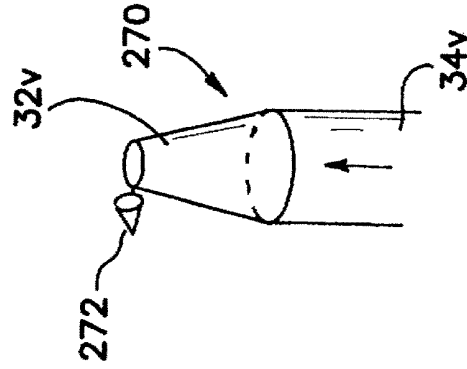


FIG. 10A

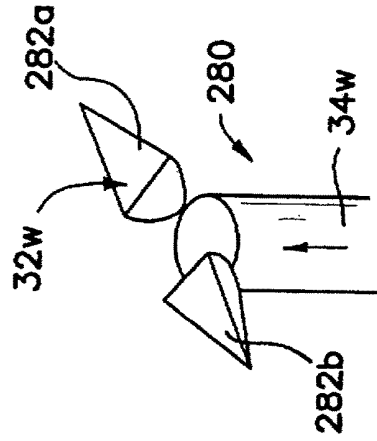


FIG. 10B

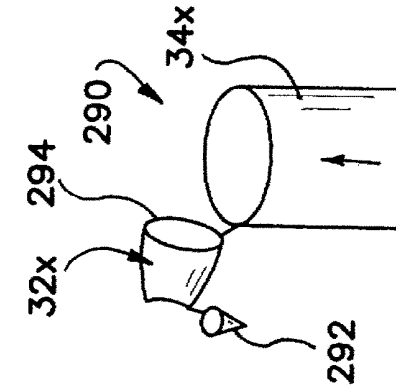


FIG. 10C

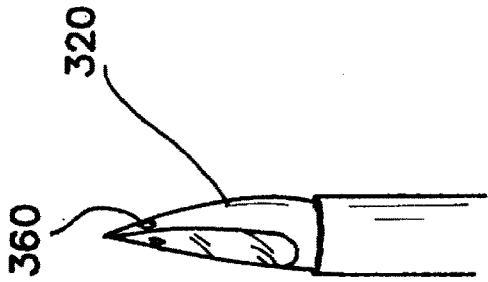


FIG. 9A

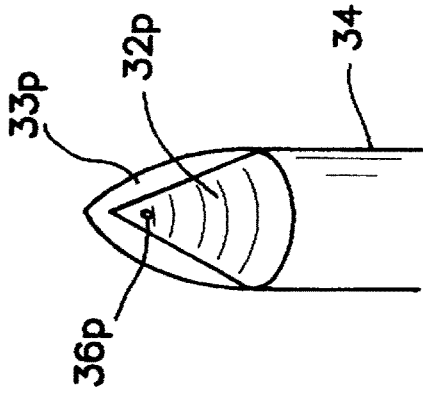


FIG. 9B

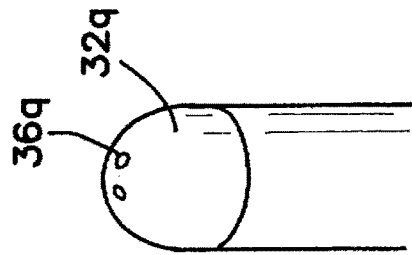


FIG. 9C

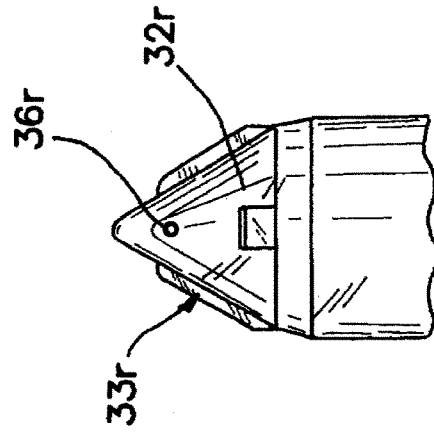


FIG. 9D

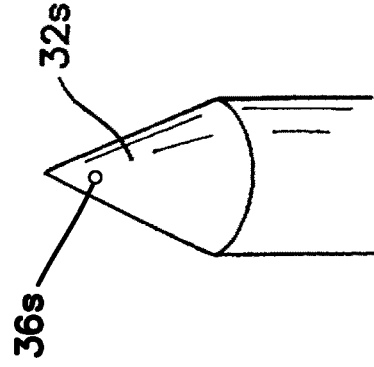


FIG. 9E

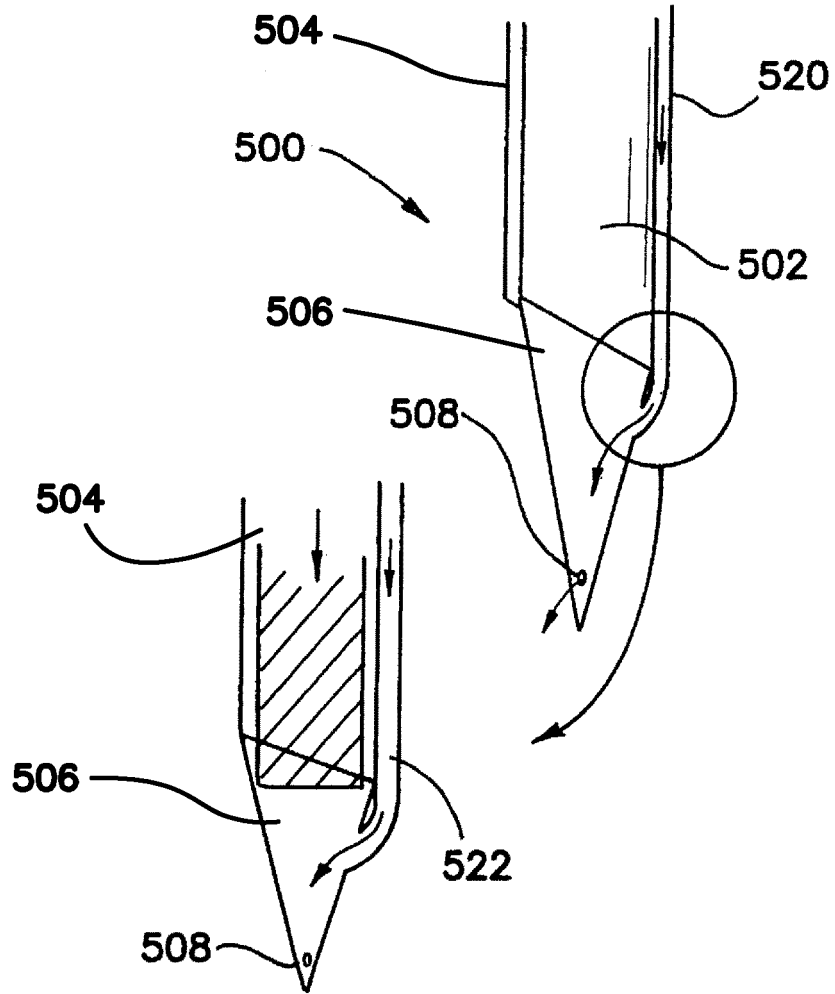
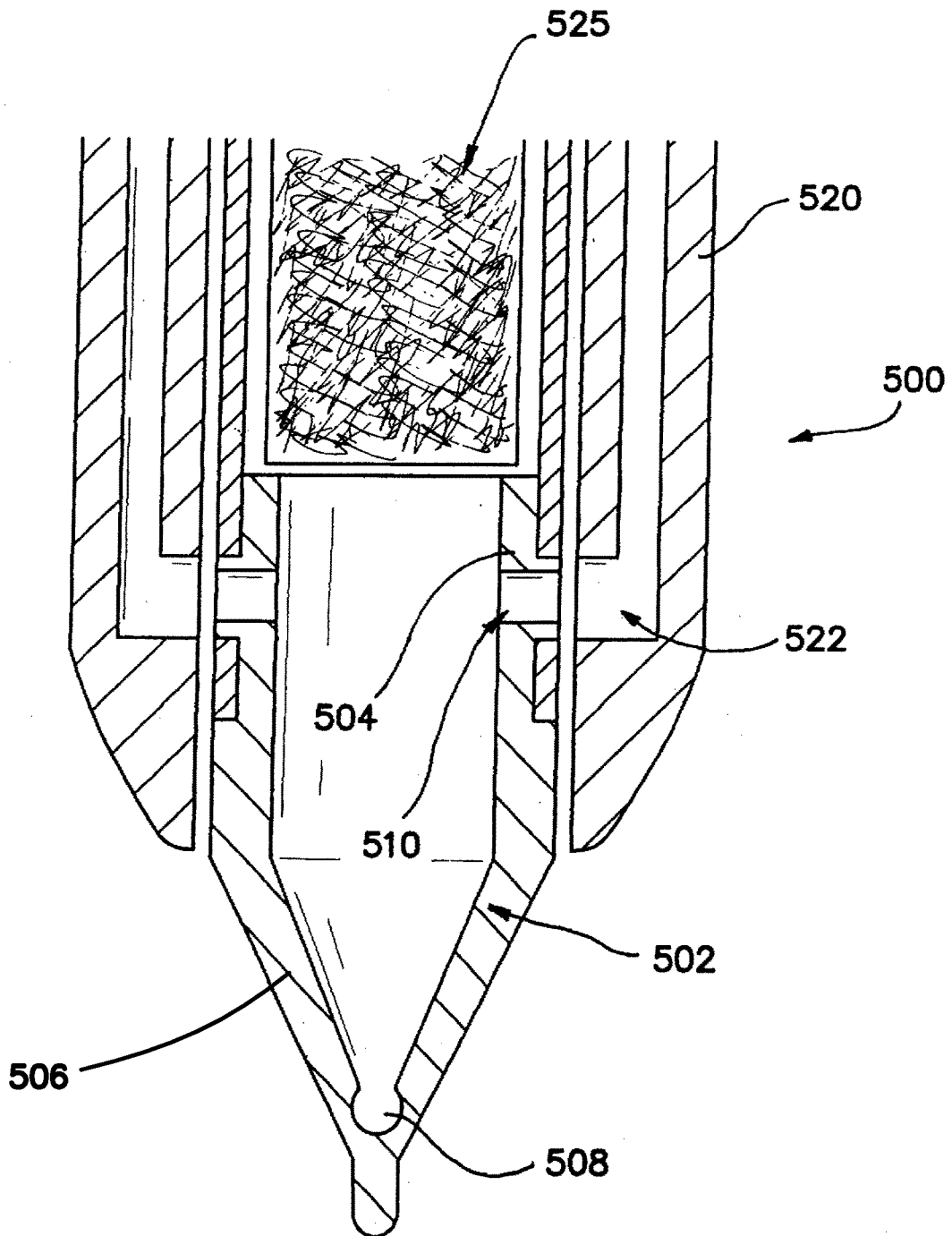


FIG. 12A

FIG. 12B



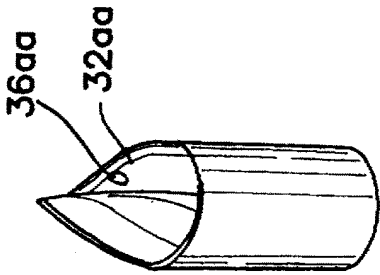


FIG. 13A

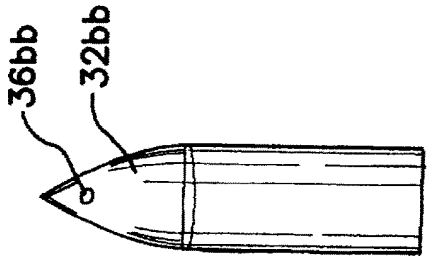


FIG. 13B

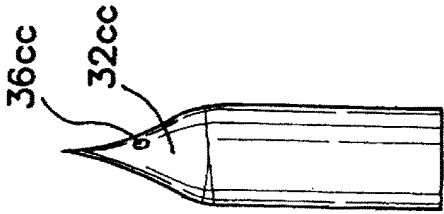


FIG. 13C

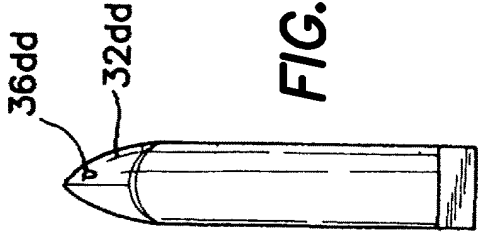


FIG. 13D

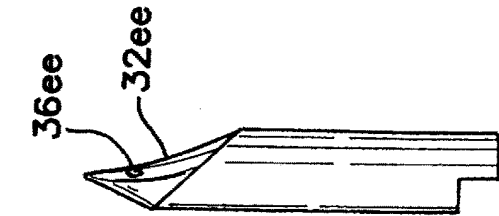


FIG. 13E

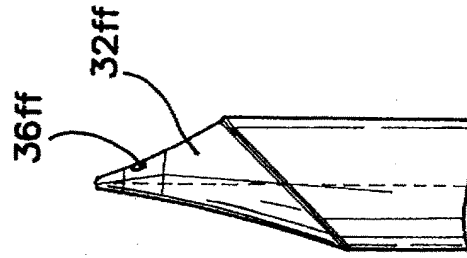


FIG. 13F

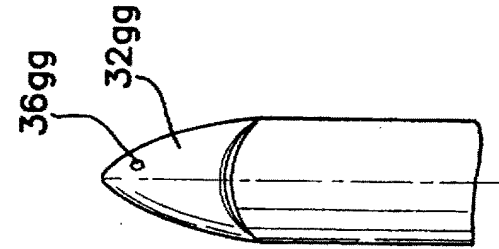


FIG. 13G

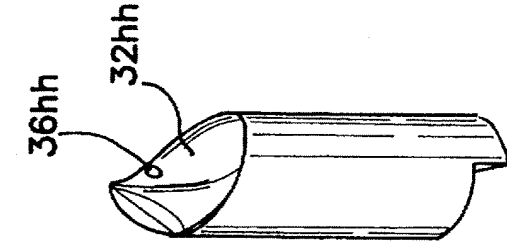


FIG. 13H

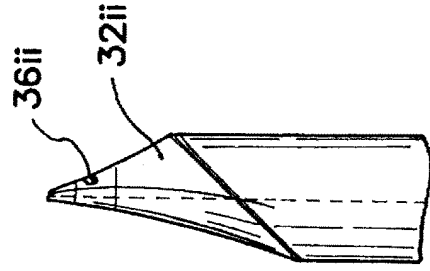


FIG. 13I

REFERENCES CITED IN THE DESCRIPTION

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专利名称(译)	充气光学手术器械		
公开(公告)号	EP1765197B1	公开(公告)日	2017-03-29
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申请(专利权)人(译)	应用医疗资源		
当前申请(专利权)人(译)	应用医疗资源CORPORATION		
[标]发明人	ALBRECHT JEREMY J BRUSTAD JOHN R TAYLOR SCOTT V JOHNSON GARY M HILAL NABIL		
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其他公开文献	EP1765197A1		
外部链接	Espacenet		

摘要(译)

一种吹气手术器械，适于穿过腹壁移动以吹入患者的腹部区域。该器械包括轴，该轴具有沿近端和远端之间的轴线延伸的吹入，吹入通道适于连接到近端压力下的流体源，轴的远端处的尖端，在所述尖端处或所述轴处形成的至少一个通气孔与所述吹气通道连接并且适于在压力下排出所述流体以吹入所述腹部区域。尖端和轴中的至少一个由诸如聚碳酸酯的半透明或透明材料形成，以便于腹壁和腹部区域的可视化。

EP 1 765 197 B1

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(54) INSUFFLATING OPTICAL SURGICAL INSTRUMENT
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INSTRUMENT CHIRURGICAL OPTIQUE INSUFFLATEUR

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US-A: 2002 103 715 US-A: 2003 023 201
US-B1: 6 228 963

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