Abstract:

A surgical instrument and method of manipulating tissue of a patient includes a body having a firing mechanism configured to be selectively manipulated by an operator, a shaft assembly extending distally from the body, and an end effector extending distally from the shaft assembly and configured to receive a cartridge for manipulating tissue of a patient. The end effector includes a distal end portion, a proximal end portion, a gap between the distal and proximal end portions, and a guide pin. The guide pin extends from the distal end portion to the proximal end portion and is connected thereto. Thereby, the guide pin is configured to secure the distal end portion relative to the proximal end portion and inhibit deflection of the distal end portion relative to the proximal end portion.
SURGICAL STAPLER WITH FIXED JAW SUPPORT PIN

BACKGROUND

Some surgical staplers are operable to clamp down on one or more layers of patient tissue, form staples through the layers of tissue to substantially seal the layers of tissue together near the formed staples, and cut through the layers of tissue for forming severed ends of operatively sealed tissue. An exemplary stapling instrument may include a pair of cooperating elongate jaw members, where each jaw member may be adapted to be inserted into a patient and positioned relative to tissue that is to be stapled and/or incised. One of the jaw members may support a staple cartridge with at least two laterally spaced rows of staples contained therein, and the other jaw member may support an anvil with staple-forming pockets aligned with the rows of staples in the staple cartridge. Generally, the stapling instrument may further include a pusher bar and a knife blade that are slidable relative to the jaw members to sequentially or simultaneously eject the staples from the staple cartridge via camming surfaces on the pusher bar and/or camming surfaces on a wedge sled that is pushed by the pusher bar. The camming surfaces may be configured to activate one or more staple drivers carried by the cartridge and associated with the staples in order to push the staples against the anvil and form laterally spaced rows of deformed staples in the tissue gripped between the jaw members. Such rows may be arranged as linear rows and/or arcuate rows for sequentially or simultaneously stapling and cutting the tissue of the patient in the form of a predetermined pattern. The knife blade may trail the camming surfaces and cut the tissue along a linear or arcuate line between the rows of staples formed in the tissue.

Merely exemplary surgical staplers are disclosed in U.S. Pat No. 6,988,650, entitled "Retaining Pin Lever Advancement Mechanism for a Curved Cutter Stapler," issued January 24, 2006; U.S. Pat No. 7,134,587, entitled "Knife Retraction Arm for a Curved Cutter Stapler," issued November 14, 2006; U.S. Pat No. 7,147,139, entitled "Closure Plate Lockout for a Curved Cutter Stapler," issued December 12, 2006, U.S.
A surgical stapler may be inserted into a patient to perform colorectal surgery. Such procedures may include the use of the stapler to operatively seal, sever, and remove the colon of the patient, in whole or in part. For instance, a proctocolectomy may be performed during a lower anterior resection ("LAR") for treating and inhibiting the spread of colorectal cancer cells. Of course, surgical staplers may be used in various other settings and procedures.

While various kinds of surgical stapling instruments and associated components have been made and used, it is believed that no one prior to the inventor(s) has made or used the invention described in the appended claims.

BRIEF DESCRIPTION OF THE DRAWINGS

The accompanying drawings, which are incorporated in and constitute a part of this specification, illustrate embodiments of the invention, and, together with the general description of the invention given above, and the detailed description of the embodiments given below, serve to explain the principles of the present invention.

FIG. 1A depicts a right front perspective view of an exemplary surgical stapling
instrument with a pin actuation mechanism in an open position and a staple cartridge in open position;

[0007] FIG. 1B depicts a right front perspective view of the surgical stapling instrument of FIG. 1A with the pin actuation mechanism in a closed position and the staple cartridge in the open position;

[0008] FIG. 1C depicts a right front perspective view of the surgical stapling instrument of FIG. 1A with the pin actuation mechanism in the closed position and the staple cartridge in a closed position via actuation of a closure mechanism;

[0009] FIG. 1D depicts a right front perspective view of the surgical stapling instrument of FIG. 1A with the pin actuation mechanism and the staple cartridge in the closed positions and a firing trigger in a fired position for stapling and cutting tissue of a patient;

[0010] FIG. 2A depicts a right side view of a handle assembly of the surgical stapling instrument of FIG. 1A, with various components removed for clarity, and with the pin actuation mechanism in a closed position and the staple cartridge in the open position;

[0011] FIG. 2B depicts a right side view of the handle assembly of the surgical stapling instrument of FIG. 1A, with various components removed for clarity, and with the pin actuation mechanism in the closed position and the staple cartridge in the closed position via actuation of the closure mechanism;

[0012] FIG. 2C depicts a right side view of the handle assembly of the surgical stapling instrument of FIG. 1A, with various components removed for clarity, and with the pin actuation mechanism and the staple cartridge in the closed positions and the firing trigger in the fired position for stapling and cutting tissue of a patient;

[0013] FIG. 3 depicts a partially exploded right front perspective view of the surgical stapling instrument of FIG. 1A showing the staple cartridge removed from a remainder of an end effector;

[0014] FIG. 4 depicts a right front perspective view of the staple cartridge of FIG. 3;
FIG. 5 depicts a rear perspective view of the staple cartridge of FIG. 3;

FIG. 6 depicts an exploded rear perspective view of the staple cartridge of FIG. 3;

FIG. 7A depicts a left side view of the end effector of FIG. 1A with various components removed for clarity;

FIG. 7B depicts a left side view of the end effector of FIG. 1A with various components removed for clarity, and with the pin actuation mechanism in a closed position and the staple cartridge in the open position;

FIG. 7C depicts a left side view of the end effector of FIG. 1A with various components removed for clarity, and with the pin actuation mechanism in the closed position and the staple cartridge in the closed position via actuation of the closure mechanism;

FIG. 7D depicts a left side view of the end effector of FIG. 1A with various components removed for clarity, and with the pin actuation mechanism and the staple cartridge in the closed positions and the firing trigger in the fired position for stapling and cutting tissue of a patient;

FIG. 8 depicts a cross-sectional view of the end effector of FIG. 7D, taken along section line 8-8 of FIG. 7D;

FIG. 9 depicts an enlarged cross-sectional view of a portion of the end effector of FIG. 8;

FIG. 10A depicts a left side view of the end effector of FIG. 1A, with various components removed for clarity, with the staple cartridge returned to the open position after actuating the firing trigger;

FIG. 10B depicts a left side view of the end effector of FIG. 1A, with various components removed for clarity, with the staple cartridge removed from the remainder of the end effector;
The drawings are not intended to be limiting in any way, and it is contemplated that various embodiments of the invention may be carried out in a variety of other ways, including those not necessarily depicted in the drawings. The accompanying drawings incorporated in and forming a part of the specification illustrate several aspects of the present invention, and together with the description serve to explain the principles of the invention; it being understood, however, that this invention is not limited to the precise arrangements shown.

DETAILED DESCRIPTION

The following description of certain examples of the invention should not be used to limit the scope of the present invention. Other examples, features, aspects, embodiments, and advantages of the invention will become apparent to those skilled in the art from the following description, which is by way of illustration, one of the best modes contemplated for carrying out the invention. As will be realized, the invention is capable of other different and obvious aspects, all without departing from the invention. Accordingly, the drawings and descriptions should be regarded as illustrative in nature and not restrictive.
It is further understood that any one or more of the teachings, expressions, embodiments, examples, etc. described herein may be combined with any one or more of the other teachings, expressions, embodiments, examples, etc. that are described herein. The following-described teachings, expressions, embodiments, examples, etc. should therefore not be viewed in isolation relative to each other. Various suitable ways in which the teachings herein may be combined will be readily apparent to those of ordinary skill in the art in view of the teachings herein. Such modifications and variations are intended to be included within the scope of the claims.

For clarity of disclosure, the terms "proximal" and "distal" are defined herein relative to a human or robotic operator of the surgical instrument. The term "proximal" refers the position of an element closer to the human or robotic operator of the surgical instrument and further away from the surgical end effector of the surgical instrument. The term "distal" refers to the position of an element closer to the surgical end effector of the surgical instrument and further away from the human or robotic operator of the surgical instrument. It will be further appreciated that for convenience and clarity, spatial terms such as "vertical," "horizontal," "lower," "upper," "front," and "rear" are used herein with respect to the drawings. However, surgical instruments are used in many orientations and positions, and these terms are not intended to be limiting and/or absolute.

I. Exemplary Surgical Stapler

FIG. 1A depicts an exemplary surgical stapling and severing instrument (10) that includes a handle assembly (12), a shaft assembly (14), and an end effector (16) distally projecting from shaft assembly (14). It should be understood that terms such as "proximal," "distal," "right," and "left" are used herein with reference to a clinician gripping handle assembly (12) of surgical stapling instrument (10). Thus, end effector (16) is distal with respect to the relatively proximal handle assembly (14). Except as otherwise described herein, instrument (10) may be configured and operable in accordance with at least some of the teachings of U.S. Pat Pub. No. 2005/0143759,
entitled "Curved Cutter Stapler Shaped for Male Pelvis," published on June 30, 2005, the disclosure of which is incorporated by reference herein; and/or U.S. Pat Application No. 14/813,242 entitled "Surgical Instrument Comprising Systems for Assuring the Proper Sequential Operation of the Surgical Instrument," filed on July 30, 2015, the disclosure of which is incorporated by reference herein.

[00036] Handle assembly (12) includes several actuation mechanisms for operating end effector (16) during the surgical procedure. To this end, exemplary handle assembly (12) includes a saddle shaped slide (18), a closure trigger (20), and a firing trigger (22) in communication with end effector (16) via shaft assembly (14). As shown in FIG. 1A, slide (18) and closure trigger (20) are in open configurations such that end effector (16) is configured to receive tissue laterally within a gap (25) between an anvil (26) and a cartridge (28) of end effector (16). Translating slide (18) distally toward end effector (16) slides a retaining pin (30) of end effector distally as shown in FIG. 1B for capturing the tissue between anvil (26) and cartridge (28). With respect to FIGS. 1C and ID, sequentially actuating closure trigger (20) and firing trigger (22) respectively compresses the tissue between anvil (26) and cartridge (28) in a closed configuration and then forms a plurality of staples (not shown) within the tissue and severs the tissue with a knife (32) (see FIG. 6) for treatment. Additional details regarding these exemplary actuation mechanisms will be provided below in greater detail.

[00037] A. Exemplary Handle Assembly and Shaft Assembly

[00038] As shown in FIGS. 1A and 2A, handle assembly (12) has a handle housing (34), a pair of handle frame plates (35, 36) within handle housing (34) extending along shaft assembly (14), saddle shaped slide (18), closure trigger (20), and firing trigger (22) as briefly discussed above. Handle housing (34) defines a hand grip (38), which the operator, such as a surgeon, grasps with the palm of at least one hand. Handle housing (34) is formed by a right shroud handle portion (40) and a left shroud handle portion (42). Closure trigger (20) is proximally positioned relative to firing trigger (22) and each are pivotally mounted to frame plates (35, 36) to extend underneath a remainder of handle
assembly (12) for manipulation by the fingers of the operator. Closure and firing triggers (20, 22) are shown in unactuated positions prior to closing end effector (16) and firing staples (not shown) and/or knife (32) (see FIG. 6). Consequently, cartridge (28) is spaced from anvil (26) for receiving tissue within gap (25) therebetween.

Surgical stapling instrument (10) captures tissue via a tissue retaining pin actuation mechanism (37) prior to actuation of the closure and firing triggers (20, 22). FIG. 1A shows retaining pin actuation mechanism (37), which includes slide (18), in the open configuration, whereas FIG. 2A shows retaining pin actuation mechanism (37) in the closed configuration in greater detail. With respect to FIG 2A, slide (18) is mounted on an upper surface of handle housing (34) and is configured to linearly translate between proximal and distal positions. Slide (18) connects to posts (44), which extend laterally outwardly from a push rod driver (46), through slots (48) (see FIG. 1A). Push rod driver (46) is restrained within handle housing (34) along longitudinal movement by slots (48). Push rod driver (46) is connected to a proximal end of a push rod (50). A distal end of push rod (50) connects to retaining pin (30) (see FIG. 6) such that distal movement of slide (18) causes push rod (50) to similarly slide proximally along shaft assembly (14) for moving retaining pin (30) (see FIG. 6) to the closed configuration, which will be discussed below in greater detail.

A closure mechanism (52), which includes closure trigger (20), is configured to selectively move cartridge (28) toward the tissue positioned between anvil (26) and cartridge (28) in the closed configuration in anticipation of stapling and/or cutting the tissue. Closure mechanism (52) further includes an elongated closure member (54), with a generally U-shaped cross-section, extending distally from handle assembly (12), through shaft assembly (14), and into end effector (16) for receiving a cartridge (28) (see FIG. 3) at a distal end portion thereof as discussed below. A proximal end portion of closure member (54) is operatively connected to closure trigger (20) by a plurality of linkages configured to convert pivoting motion of closure trigger (20) into translation of closure member (54). More particularly, the intermediate and proximal end portions of
closure member (54) extend through handle assembly (12) between left and right handle frame plates (35, 36). Right and left closure links (56) are respectively pivotally attached at the right and left proximal ends of closure member (54) by an integral closure link pin (58). At an opposite end of the closure links (56), closure links (56) are pivotally attached to another integral closure link pin (60). Closure link pin (60) connects closure links (56) to a slotted closure arm link (62), which is pivotally mounted to handle frame plates (35, 36) at a closure trigger pin (64). Closure trigger (20) descends from the slotted closure arm link (62) for pivotal rotation about closure trigger pivot pin (64) both toward and away from hand grip (38). A closure spring (66) housed within hand grip (38) is secured to the slotted closure arm link (62) to provide a desired resistance when the operator squeezes closure trigger (20) toward hand grip (38), and to bias closure trigger (20) toward the open position.

Closure member (54) is further configured for directing movement of tissue retaining pin actuation mechanism (37) to automatically direct movement of the retaining pin (30) to the closed configuration while the operator squeezes closure trigger (20). Such automation may be useful in the event that the operator did not manually move the slide (18) to the distal position before actuating trigger (20). Closure member (54) includes posts (68), which extend laterally on each opposing side of closure member (54) within handle housing (34). Posts (68) slidably connect to a yoke (70) via L-shaped slots (72). Yoke (70) is pivotally mounted within handle housing (34) by a pivot pin (74). Yoke (70) further includes cam pins (76) that are configured to push camming surfaces (78) on push rod driver (46). Thus, actuating closure trigger (20) to an intermediate position shown in FIG. 2A directs the closure member (52) distally and, in turn, causes yoke (70) to engage push rod driver (46) and force retaining pin (30) (see FIG. IB) to the closed position. Slide (18) is thereby dragged along handle housing (34) from the proximal position to the distal position in the event that the operator did not manually manipulate slide (18) to the distal position before actuating trigger (20).

The operator further squeezes the closure trigger (20) to the hand grip (38) as
shown in FIGS. 1C and 2B to effectively set surgical stapling instrument (10) in the
closed configuration prior to forming the staples (not shown) and severing the tissue as
discussed briefly above. Exemplary handle assembly (12) is configured to form the
staples (not shown) and sever the tissue via a firing mechanism (80) upon operator
manipulation of firing trigger (22) toward closure trigger (20) as shown in FIGS. 1D and
2C. With respect to FIGS. 1C, ID, 2B, and 2C, firing mechanism (80), which includes
firing trigger (22), has a firing bar (82) extending distally from handle assembly (12) and
within end effector (16). A distal end of firing bar (82) cooperates with cartridge (28) as
discussed below in greater detail, whereas a proximal end of firing bar (82) is operatively
connected to firing trigger (80) for selective firing thereof.

Firing bar (82) has a rectangular receiving slot (84) [see FIG. 2A] in a portion of
firing bar (82) positioned within handle housing (34). Integral closure link
pin (58) extends through receiving slot (84). The underside of the proximal end
portion of firing bar (82) has a sliding surface (86). The proximal end portion of firing
bar (82) also has a terminal side engagement surface (82) extending from sliding
surface (86). Firing trigger (22) is pivotally mounted to handle frame plates (35, 36) by a
firing trigger pin (90) spaced from closure trigger pin (64) such that each of pins (90, 64)
pivot about mutually independent axes. Firing trigger (22) includes an arcuate firing
trigger link (92) extending from firing trigger (22) at firing trigger pin (90) to an
apex (94), which rests on sliding surface (86) of the proximal end portion of firing
bar (82). Within handle assembly (12), firing trigger (22) is attached to firing trigger
spring arms (95, 96), respectively. Firing trigger spring arms (95, 96) support a torsion
spring (not shown) on the right half of firing trigger (22). Finally, a firing bar return
spring (98) is secured to the underside of firing bar (82) at the portion of firing
bar (82) within handle assembly (12) to bias firing bar (82) toward its unactuated
position.

As the operator squeezes closure trigger (20) toward hand grip (38), slotted
closure arm link (62) and closure links (56) move distally within receiving slot (84) of
firing bar (82). This distal movement causes closure member (54) to correspondingly move distally. Likewise, firing bar (82) concurrently moves distally with closure member (54), because integral closure link pin (58), to which closure links (56) are attached, extends through receiving slot (84) in firing bar (82) (see FIG. 2A). Thereby, firing bar (82) is forced distally to form the staples (not shown) in the tissue and/or sever the tissue with knife (32) (see FIG. 6). Finally, the operator may fully squeeze firing trigger (22) toward hand grip (38) to "fire" surgical stapling instrument (10) and force firing bar (82) further distally to form the staples (not shown) and sever the tissue. This distal movement of firing bar (82) may also be referred to herein as "firing" the firing bar (82) to the actuated or "fired" position.

Upon operator release of one or both of closure and firing triggers (20, 22) while one or both of triggers (20, 22) is/are in a fired position, or in an intermediate position between the unactuated and fired positions, surgical stapling instrument (10) may be further configured to releasably lock in one of a variety of configurations. The operator may then release the hand grip (38) to free one or more hands for another task during the surgical procedure and, when desired, release surgical stapling instrument (10) from its locked position by release button (24). By way of example, surgical stapling instrument (10) has an intermediate closure detent position and a closure detent position. With respect to FIGS. 2A-2C, the top side of the slotted closure arm link (62) has a clamp sliding surface (102) that displays an intermediate detent (104) and a closure detent (106). A release pawl (108) slides on clamp sliding surface (102) and may engage intermediate and closure detents (104,106). Release pawl (108) has a laterally extending pawl lug (110) at its distal end.

Release pawl (108) is located within handle assembly (12) and is integrally formed with release button (24), which is situated exterior of handle housing (34) for manipulation by the operator. Release button (24) has a thumb rest (112) pivotally attached to handle housing (34) by a release trunnion (114). Release button (24) is biased outwardly from handle housing (34) and, therefore, release pawl (108) is biased
downwardly toward clamp sliding surface (102) by a release spring (116). Release
spring (116) is mounted to handle housing (34) by a spring retention pin (118) and is
mounted to release button (24) by a button spring post (120). Slotted closure arm
link (62) has an arcuate recess (122) located between intermediate and closure
detents (104, 106). Resting within arcuate recess (122) for rotational movement are
integrally connected left and right hand toggles (124). Each toggle (124) has a toggle
arm (126) that is engageable with pawl lug (110).

[00047] In order to releasably lock handle assembly (12), toggle arms (126) from pawl
lug (110) disengage from pawl lug (110) as closure trigger (20) is squeezed toward hand
grip (38). Consequently, as toggle (124) continues to rotate in a clockwise direction,
release pawl lug (108) rides up toggle arms (126) and, with continued motion of closure
trigger (20), falls into one of intermediate and closure detents (104, 106), depending on
the position of closure trigger (20) in use. As release pawl (108) rides up toggle
arm (126), release pawl (108) rotates release button (24) clockwise. Release pawl
(108) thereby falls into one of intermediate and detents (104, 106) and generates an
audible clicking sound alerting the surgeon that one of the intermediate and closure
positions have been reached.

[00048] In order to release handle assembly (12) from the intermediate or closure
positions discussed herein, the surgeon depresses release button (24). In turn, release
pawl (108) pivots about release trunnion (114) in a clockwise direction to dislodge pawl
lug (110) from one of the intermediate and closure detents (104, 106). As pawl lug (110)
is dislodged, pawl lug (110) rides on toggle arms (126) to another position, such as the
unactuated position. Therefore, the operator may release closure and firing triggers (20,
22) such that each may return to the unactuated positions FIG. 1A and FIG. 3.

[00049] Surgical stapling instrument (10) of the present example includes each of handle
frame plates (35, 36), push rod (50), closure member (54), and firing bar (82) extending
continuously from handle assembly (12) to end effector (16), thereby defining shaft
assembly (14) extending therebetween. Handle frame plates (35, 36), push rod (50),
closure member (54), and firing bar (82) of surgical stapling instrument (10) provide merely a subset of elongated components extending distally from handle assembly (12) as shaft assembly (14). Alternatively, shaft assembly (14) may include additional components, such as an articulating joint, or may include a rearrangement of various components such that shaft assembly (14) may be modular relative to handle assembly (12). In any case, it will be appreciated that the invention is not intended to be limited to shaft assembly (14) described herein, and may include various alternative arrangements for operatively connecting end effector (16) to handle assembly (12). Of course, handle assembly (12) and shaft assembly (14) may have a variety of other components, features, and operabilities, in addition to or in lieu of any of those noted above. Other suitable configurations for handle and shaft assemblies (12, 14) will be apparent to those of ordinary skill in the art in view of the teachings herein.

[00050] B. Exemplary End Effector

[00051] As also shown in FIGS. 3-5 and discussed briefly above, end effector (16) of the present example includes anvil (26), replaceable cartridge (28) including a plurality of staples (not shown) and knife (32) (see FIG. 6), and retainer pin (30). While end effector (16) of the present example is adapted for use in conjunction with replaceable cartridge (28) having various components, it will be appreciated that the concepts underlying the present invention could be applied to a variety of end effector and cartridge constructions for treating the patient.

[00052] End effector (16) provides a surgical fastening assembly that includes cartridge (28) received within a C-shaped supporting structure (128). The term C-shaped is used throughout the specification to describe the concave nature of supporting structure (128) and cartridge (28). The C-shaped construction facilitates enhanced functionality and access to tissue within the patient. The term "C-shaped" as used herein should be construed to include a variety of concave shapes that would similarly enhance the functionality of surgical stapling and cutting instruments. By way of example only, the C-shape of supporting structure (128) may be sized to promote access to the lower
colon within the pelvic bowl of a patient, such as to perform a LAR in a proctocolectomy procedure.

Supporting structure (128) of end effector (16) is respectively attached to handle frame plates (35, 36) of shaft assembly (14) by a shoulder rivet (129) and posts (130) which extend from supporting structure (128) into receiving holes in handle frame plates (35, 36). The distal end of closure member (54) is disposed to receive cartridge (28) thereon for directing cartridge (28) to the closed configuration. Upon return of cartridge (28) from the closed configuration to the open configuration, cartridge (28) further includes a safety lockout mechanism (131) (see FIG. 7A) configured to inhibit inadvertently re-firing cartridge (28). Safety lockout mechanism (131) will be discussed below in additional detail.

Cartridge (28) includes anvil (26) coupled to a cartridge housing (132). Cartridge (28) also includes retaining pin (30) and a tissue contacting surface (34), which defines a plurality of staple-containing slots (136) in staggered formation in one or more rows on either side of knife (32) (see FIG. 6). Staples (not shown) are fired from cartridge housing (132) against a staple-forming surface (138) of anvil (26) that faces tissue-contacting surface (134) of cartridge housing (132). Cartridge (28) may also include a removable retainer (not shown) for storage between anvil (26) and tissue contacting surface (34) prior to and/or after use in order to inhibit unintended contact with various portions of cartridge (28).

As shown in FIGS. 4-6, cartridge (28) includes a staple driver assembly (140) within cartridge housing (132) and proximally positioned behind the plurality of staples (not shown) within staple-containing slots (136). Driver assembly (140) of the present example is formed as a unitary structure of a plurality of staple drivers (141). Thus, the term "assembly" is not intended to be limited to an assembly of individual components, but may also include integrally formed components with unitary structures. Driver assembly (140) is configured to push the staples (not shown) respectively out of staple containing slots (136) and toward anvil (26) for formation. A knife holder (142) is
disposed immediately proximal of driver assembly (140) in cartridge housing (132) and defines a slot (144) and ledge (146) for interaction with a knife retractor hook (148) (see FIG. 10B), which is discussed below in greater detail. Knife holder (142) is attached to knife (32) such that knife (32) extends distally from knife holder (142) through a slot (150) in driver assembly (140) and through another slot (152) in cartridge housing (132). Although knife (32) is disclosed as being within cartridge housing (132) in the present example, other configurations may also be used. For example, it will be appreciated that cartridge (28) may alternatively not include knife (32) for alternative treatments.

Knife holder (142) has a detent post (154) that extends through a slot (155) in cartridge housing (132). Detent post (154) is positioned in order to contact a detent protrusion (156) of cartridge slot (155) during the longitudinal travel of knife (132) and knife holder (142). Similarly, driver assembly (140) has a detent post (158) positioned in order to contact proximal and distal detent protrusions (159, 160) of cartridge slot (155).

Knife (32) and slots (150, 152) are positioned such that there is at least one row of staples (not shown) on either side of knife (132). In some versions, two rows of staple slots (136) containing respective rows of staples (not shown) are provided on each side of slot (152) of cartridge housing (132).

Cartridge housing (132) defines two longitudinally extending, generally circular holes (162, 164) at respective ends of knife slot (152). More particularly, hole (162) at a lower portion of cartridge housing (132) is shaped and dimensioned to receive a guide pin (166) through cartridge housing (132). Hole (164) at an upper portion of cartridge housing (132) is shaped and dimensioned to slidably receive retaining pin (30) through cartridge housing (132). Staple slots (136) of the present example are arranged such that the staples (not shown) laterally extend past the generally circular holes (162, 164).

Anvil (26) of the present example includes a plastic cutting washer (168) and a metallic staple-forming surface (138). Anvil (26) is disposed to maintain staple-forming surface (138) in alignment with the staples (not shown) to receive and form the staples
Retaining pin (30) is connected to a couplet (170) by a circumferential slot (172) in retaining pin (30) and a groove (not shown) in couplet (170). Couplet (170) is disposed within an arm (176) of cartridge housing (132) and is secured to arm (176) by an end cap (178).

Guide pin (166) and retaining pin (30) include respective slots (180, 182) into which lower and upper ends (184, 186) of knife (32) are slidably disposed. A proximal end (188) of guide pin (166) is connected to anvil (26), whereas a distal end (190) of guide pin (166) extends from cartridge housing (132) and extends through a slot (192) in anvil (26). Cutting washer (168) slips onto anvil (26) via groove (194). Thereby, cutting washer (168) is configured to trap guide pin (166) in the opening formed by slot (192) in anvil (26) and a cutting surface (157) of anvil (26) for connecting anvil (26) to cartridge housing (132).

Lockout mechanism (131) is shown in FIG. 7A in greater detail. Lockout mechanism (131) is configured to inhibit full proximal movement of cartridge housing (132) to its unactuated position after firing. To this end, lockout mechanism (131) of the present example includes a lockout lever (196) that is pivotally mounted to the distal end of closure member (54) by a pin (198). Lockout lever (196) is spring biased toward the proximal end portion of supporting structure (128) by a spring (not shown). A proximal end portion of lockout lever (196) has a cam surface (200) and a locking groove (202). Supporting structure (128) of end effector (16) also has a ledge (204) that is configured to cooperate with locking groove (202) when lockout mechanism (131) is engaged. In contrast, supporting structure (128) has a base surface (206) configured to cooperate with cam surface (200) when lockout lever (131) is not engaged.

C. Exemplary Actuation of Cartridge

In the present example, cartridge (28) is driven toward anvil (26) via closure member (54) until reaching the closed configuration with tissue positioned between cartridge (28) and anvil (26) as discussed above with respect to handle assembly (12). From the closed configuration, knife (32) and staple driver assembly (140) are further
moved toward anvil (26) via firing bar (82) to form staples (not shown) in the tissue, fluidly seal the tissue, and sever the tissue for treating the patient. While actuation of cartridge (28) includes stapling and severing tissue in this example, it will be appreciated that one or more of these steps may be omitted from treatment as desired by the operator. Moreover, it will be appreciated that surgical stapling instrument (10) may be reconfigured to perform these steps simultaneously or sequentially as desired. For example, actuation of firing bar (82) causes driver assembly (140) and knife (32) to move distally toward anvil (26) in the present example. Alternatively, surgical stapling instrument (10) may be reconfigured to selectively fire one of staples (not shown) or knife (32), or selectively fire staples (not shown) and then knife (32), or vice versa. It should therefore be understood that the invention is not intended to be limited to the particular operation of surgical stapling instrument (10) or the associated treatment.

As shown in FIG. 7A, cartridge (28) is spaced proximally from anvil (26) to receive tissue within gap (25) in the open configuration. With tissue received between cartridge (28) and anvil (26), the operator manually directs push rod (50) distally via slide (18) as discussed above and shown in FIG. 7B. Push rod (50) is operatively connected to couplet (70) (see FIG. 6), which is connected to retaining pin (30). Thus, distally translating push rod (50) similarly translates retaining pin (30) to extend from cartridge (28) to anvil (26) and capture tissue between retaining pin (30) and guide pin (166).

As shown in FIG. 7C, manipulation of closure trigger (20) (see FIG. 1C) forces closure member (54) to translate distally relative to supporting structure (128) of end effector (16). Closure member (54) supports cartridge (28) thereon such that distal translation of closure member (54) similarly moves firing bar (82) and cartridge (28) toward anvil (26). With cartridge (28) in the closed configuration and the tissue effectively captured in the end effector (16), the operator manipulates firing trigger (22) (see FIG. 1D) toward anvil (26) to the fired position. Distal translation of firing bar (82) causes firing bar (82) to engage knife holder (142), which supports both driver assembly
(140) and knife (32) extending through driver assembly (140) as shown in FIG. 7D. In
turn, driver assembly (140) directs staples (not shown) from staple slots (136) and against
staple-forming surface (138) to form the staples (not shown) within the tissue for fluidly
sealing the tissue. As the staples (not shown) are formed, knife (32) continues to
translate distally through tissue and into anvil (26) to sever the fluidly sealed tissue.
FIGS. 8-9 illustrate the fired cartridge (28) in greater detail, with knife (32) guided along
cartridge housing slot (152), guide pin slot (180); and with retaining pin slot (182)
between rows of staple slots (136) toward anvil (26).

[00066] Once fired, the operator may depress release button (24) (see FIG. 2C) and
withdraw closure member (54) and firing bar (82) proximally from the actuated, fired
position to the unactuated position shown in FIGS. 10A-10B. More particularly, retractor hook (148) engages knife holder (142) to pull knife (32) proximally. At
approximately the same time, as cartridge (28) translates proximally with closure
member (54), lockout lever (196) of lockout mechanism (131) engages cartridge housing
(132) to hold cartridge housing (132) in position. Thereby, the continued pull of knife
(32) retracts knife (32) within cartridge housing (132) to inhibit unintended contact by
operator with knife (32). Cartridge (28) may then be removed from supporting structure
(128) of end effector (16), discarded, and replaced for further treatment if so desired. Of
course, various suitable settings and procedures in which surgical stapling instrument
(10) may be used will be apparent to those of ordinary skill in the art in view of the
teachings herein.

[00067] It should also be understood that any other components or features of surgical
stapling instrument (10) may be configured and operable in accordance with any of the
various references cited herein. Additional exemplary modifications that may be
provided for surgical stapling instrument (10) will be described in greater detail below.
Various suitable ways in which the below teachings may be incorporated into surgical
stapling instrument (10) will be apparent to those of ordinary skill in the art. Similarly,
various suitable ways in which the below teachings may be combined with various
teachings of the references cited herein will be apparent to those of ordinary skill in the art. It should also be understood that the below teachings are not limited to surgical stapling instrument (10) or devices taught in the references cited herein. The below teachings may be readily applied to various other kinds of instruments, including instruments that would not be classified as surgical staplers. Various other suitable devices and settings in which the below teachings may be applied will be apparent to those of ordinary skill in the art in view of the teachings herein.

[00068] II. Exemplary Surgical Stapling Instruments with Alternative End Effectors

[00069] While the above surgical stapling instrument (10) provides one example of end effector (16) projecting distally from handle assembly (12), it will be appreciated that the operator may desire an alternative end effector depending on one of a variety particular treatments. Various tissues may be more or less difficult to treat depending on size and/or density within the patient. Despite the operator properly positioning end effector (16) relative to tissue for accurately and precisely stapling and severing the tissue, thicker and/or denser tissues often require added force to be transmitted along surgical stapling instrument (10) and, in turn, may cause one or more components of instrument (10) to deform during use. For example, compressing a portion of the colon between anvil (26) and cartridge (28) in the closed configuration may deform supporting structure (128) of end effector (16), particularly as the tissue is stapled and severed. Thus, the particular location in which the staples form and the knife (32) cuts may vary or deviate a small, but relatively meaningful, amount that may negatively impact the effectiveness of the treatment. It may therefore be desirable to provide surgical stapling instrument (310) with end effector (316) having a supporting structure (328) that is configured to reduce deformation within end effector (316) during treatment. By way of example, end effector (316) may have supporting structure (448) configured to cooperate with exemplary guide and pin (466) for providing greater structural rigidity during use for reducing deformation within end effector (316); and further increasing the accuracy and precision of the treatment.
End effector (316) is described below in the context of a proctocolectomy surgical procedure. While the following description of end effector (316) and methods of treatment is provided in the context of stapling and/or cutting colon tissue, it will be appreciated that surgical stapling instrument (310) and end effectors (316) may be alternatively configured to treat any tissue in the human body with similar features. It should also be understood that the features discussed below may be readily incorporated into surgical stapling instrument (10) discussed above. To this end, like numbers indicate like features described above in greater detail.

A. Exemplary End Effector with Structural Guide Pin

As shown in FIGS. 13-15, end effector (316) has a structural guide pin (466) configured to receive tissue thereagainst for inhibiting lateral positioning of tissue beyond cartridge (28) and anvil (26) and guide movement of cartridge (332) from the closed configuration toward the open configuration as discussed above. In addition, structural guide pin (466) also rigidifies distal end portion (564), which receives anvil (26). Structural guide pin (446) extends through hole (162) in cartridge (328) and is configured to guide longitudinal movement of cartridge (328) between distal and proximal positions as discussed above in greater detail with respect to surgical stapling instrument (10) (see FIGS. 1A-1D). In addition, structural guide pin (466) extends from distal end portion (564) of supporting structure (448) and into a proximal end portion (571) to traverse the opening between anvil (26) and cartridge (328), providing structural rigidity therealong to inhibit distal end portion (564) from deflecting relative to proximal end portion (571) during treatment.

As shown in FIGS. 14 and 15, structural guide pin (466) of the present example is in the form of a threaded fastener having a distal head end (466a) and a proximal threaded end (466b). Supporting structure (448) has a distal pin bore (572) positioned offset from a proximal pin bore (573) such that distal and proximal pin bores (572, 573) receive structural guide pin (466) from the distal direction. Distal pin bore (572) is thus configured such that distal head end (466a) seats therein; whereas proximal pin bore
(572) receives a threaded nut (574) seated opposite the distal head end (466a). Proximal threaded end (466b) rotatably threads into threaded nut (574) such that structural guide pin (466) is held in tension between pin bores (572, 573) as shown in FIG. 15.

[00074] In use, structural guide pin (466) receives tissue thereagainst and guides cartridge (328) between open and closed configurations as discussed above and shown in FIGS. 1A and 1C, respectively. In addition, structural guide pin (466) braces supporting structure (448) in tension between distal and proximal end portions (564, 571) as tissue is compressed between anvil (26) and cartridge (328). In other words, structural guide pin (466) rigidifies supporting structure (448) to inhibit distal end portion (564) from deflecting relative to proximal end portion (571) of end effector (316). For example, structural guide pin (466) also inhibits distal end portion (564) from deflecting relative to proximal end portion (571) as staples are pressed into staple-forming surface (138) of anvil (26) from cartridge (328) for improved accuracy and precision of staple formation and tissue severing.

[00075] III. Exemplary Combinations

[00076] The following examples relate to various non-exhaustive ways in which the teachings herein may be combined or applied. It should be understood that the following examples are not intended to restrict the coverage of any claims that may be presented at any time in this application or in subsequent filings of this application. No disclaimer is intended. The following examples are being provided for nothing more than merely illustrative purposes. It is contemplated that the various teachings herein may be arranged and applied in numerous other ways. It is also contemplated that some variations may omit certain features referred to in the below examples. Therefore, none of the aspects or features referred to below should be deemed critical unless otherwise explicitly indicated as such at a later date by the inventors or by a successor in interest to the inventors. If any claims are presented in this application or in subsequent filings related to this application that include additional features beyond those referred to below,
those additional features shall not be presumed to have been added for any reason relating to patentability.

Example 1

A surgical instrument comprising: (a) a body having a firing mechanism configured to be selectively manipulated by an operator; (b) a shaft assembly extending distally from the body; and (c) an end effector extending distally from the shaft assembly and configured to receive a cartridge for manipulating tissue of a patient, the end effector including: (i) a distal end portion, (ii) a proximal end portion, (iii) a gap between the distal end portion and the proximal end portion, and (iv) a guide pin extending from the distal end portion to the proximal portion, wherein the guide pin is configured to receive the tissue thereagainst for positioning tissue within the gap, wherein the guide pin is connected to each of the distal and proximal end portions to secure the distal end portion relative to the proximal end portion and inhibit deflection of the distal end portion relative to the proximal end portion.

Example 2

The surgical instrument of Example 1, wherein the distal end portion extends distally from the proximal end portion in the form of a hook-shape supporting structure with the gap therebetween configured to receive the tissue.

Example 3

The surgical instrument of any one or more of Examples 1 through 2, wherein the guide pin is connected to each of the distal and proximal end portions in tension to secure the distal end portion relative to the proximal end portion.

Example 4

The surgical instrument of any one or more of Examples 1 through 3, wherein the guide pin is in the form of a threaded fastener.
The surgical instrument of any one or more of Examples 1 through 4, wherein the distal end portion has a distal pin bore and the proximal end portion has a proximal pin bore, wherein the distal and proximal pin bores are positioned opposite each other and receive the guide pin therein.

Example 6

The surgical instrument of Example 5, wherein the proximal pin bore includes a threaded nut, and wherein the guide pin is threaded into the threaded nut to be connected therein.

Example 7

The surgical instrument of Example 6, wherein the guide pin is in the form of a threaded fastener.

Example 8

The surgical instrument of any one or more of Examples 4 through 7, wherein the threaded fastener is connected to each of the distal and proximal end portions in tension to secure the distal end portion relative to the proximal end portion.

Example 9

The surgical instrument of any one or more of Examples 1 through 8, further comprising a cartridge received against the guide pin, wherein the cartridge is configured to be guided from the proximal end portion toward the distal end portion along the guide pin.

Example 10

The surgical instrument of Example 9, wherein the cartridge includes at least one
of a knife or a plurality of staples, wherein the knife is configured to cut the tissue, and
wherein the plurality of staples are configured to fasten the tissue.

[0009] Example 11

[00098] The surgical instrument of Example 10, wherein the cartridge includes each of the
knife and the plurality of staples.

[00099] Example 12

[00100] The surgical instrument of any one or more of Examples 1 through 11, wherein
the distal end portion of the end effector includes an anvil, and the guide pin is
configured to inhibit deflection of the anvil relative to the proximal end portion.

[00101] Example 13

[00102] The surgical instrument of any one or more of Examples 1 through 12, wherein
the guide pin is a rigid guide pin.

[00103] Example 14

[00104] An end effector of a surgical instrument for manipulating tissue of a patient with a
cartridge, comprising: (a) a distal end portion; (b) a proximal end portion; (c) a gap
between the distal end portion and the proximal end portion; and (d) a guide pin
extending from the distal end portion to the proximal portion, wherein the guide pin is
configured to receive the tissue thereagainst for positioning the tissue within the gap,
wherein the guide pin is connected to each of the distal and proximal end portions to
secure the distal end portion relative to the proximal end portion and inhibit deflection of
the distal end portion relative to the proximal end portion.

[00105] Example 15

[00106] The surgical instrument of Example 14, wherein the distal end portion extends
distally from the proximal end portion in the form of a hook-shape supporting structure
with the gap therebetween configured to receive the tissue.

[000107] Example 16

[000108] The surgical instrument of any one or more of Examples 14 through 16, wherein the guide pin is connected to each of the distal and proximal end portions in tension to secure the distal end portion relative to the proximal end portion.

[000109] Example 17

[000110] The surgical instrument of any one or more of Examples 14 through 16, wherein the guide pin is in the form of a threaded fastener.

[000111] Example 18

[000112] The surgical instrument of any one or more of Examples 14 through 17, further comprising a cartridge received against the guide pin, wherein the cartridge is configured to be guided from the proximal end portion toward the distal end portion along the guide pin.

[000113] Example 19

[000114] A method of manipulating tissue of a patient with a surgical instrument, the surgical instrument including a body having a firing mechanism configured to be selectively manipulated by an operator, a shaft assembly extending distally from the body, and an end effector extending distally from the shaft assembly and configured to receive a cartridge for manipulating the tissue of the patient, wherein the end effector has a distal end portion, a proximal end portion, a gap between the distal and proximal end portions, and a guide pin extending from the distal end portion to the proximal end portion, wherein the guide pin is configured to receive the tissue thereagainst for positioning the tissue within the gap, wherein the guide pin is connected to each of the distal and proximal end portions to secure the distal end portion relative to the proximal end portion and inhibit deflection of the distal end portion relative to the proximal end
portion, the method comprising: (a) receiving the tissue of the patient within the gap between the distal and proximal end portions of the end effector; (b) positioning the tissue against the guide pin; (c) inhibiting deflection of the distal end portion of the end effector relative to the proximal end portion of the end effector with the guide pin; and (d) manipulating the firing mechanism to thereby manipulate the tissue of the patient.

[000115] Example 20

[000116] The method of Example 19, further comprising increasing tension in the guide pin such that guide pin further inhibits deflection of the distal end portion of the end effector relative to the proximal end portion of the end effector.

[000117] IV. Miscellaneous

[000118] It should be understood that any one or more of the teachings, expressions, embodiments, examples, etc. described herein may be combined with any one or more of the other teachings, expressions, embodiments, examples, etc. that are described herein. The above-described teachings, expressions, embodiments, examples, etc. should therefore not be viewed in isolation relative to each other. Various suitable ways in which the teachings herein may be combined will be readily apparent to those of ordinary skill in the art in view of the teachings herein. Such modifications and variations are intended to be included within the scope of the claims.

[000119] It should be appreciated that any patent, publication, or other disclosure material, in whole or in part, that is said to be incorporated by reference herein is incorporated herein only to the extent that the incorporated material does not conflict with existing definitions, statements, or other disclosure material set forth in this disclosure. As such, and to the extent necessary, the disclosure as explicitly set forth herein supersedes any conflicting material incorporated herein by reference. Any material, or portion thereof, that is said to be incorporated by reference herein, but which conflicts with existing definitions, statements, or other disclosure material set forth herein will only be
incorporated to the extent that no conflict arises between that incorporated material and the existing disclosure material.

[000120] The surgical instrument systems described herein have been described in connection with the deployment and deformation of staples; however, the embodiments described herein are not so limited. Various embodiments are envisioned which deploy fasteners other than staples, such as clamps or tacks, for example. Moreover, various embodiments are envisioned which utilize any suitable means for sealing tissue. For instance, an end effector in accordance with various embodiments can comprise electrodes configured to heat and seal the tissue. Also, for instance, an end effector in accordance with certain embodiments can apply vibrational energy to seal the tissue.

[000121] Versions of the devices described above may be designed to be disposed of after a single use, or they can be designed to be used multiple times. Versions may, in either or both cases, be reconditioned for reuse after at least one use. Reconditioning may include any combination of the steps of disassembly of the device, followed by cleaning or replacement of particular pieces, and subsequent reassembly. In particular, some versions of the device may be disassembled, and any number of the particular pieces or parts of the device may be selectively replaced or removed in any combination. Upon cleaning and/or replacement of particular parts, some versions of the device may be reassembled for subsequent use either at a reconditioning facility, or by an operator immediately prior to a procedure. Those skilled in the art will appreciate that reconditioning of a device may utilize a variety of techniques for disassembly, cleaning/replacement, and reassembly. Use of such techniques, and the resulting reconditioned device, are all within the scope of the present application.

[000122] By way of example only, versions described herein may be sterilized before and/or after a procedure. In one sterilization technique, the device is placed in a closed and sealed container, such as a plastic or TYVEK bag. The container and device may then be placed in a field of radiation that can penetrate the container, such as gamma radiation, x-rays, or high-energy electrons. The radiation may kill bacteria on the device
and in the container. The sterilized device may then be stored in the sterile container for later use. A device may also be sterilized using any other technique known in the art, including but not limited to beta or gamma radiation, ethylene oxide, or steam.

Having shown and described various embodiments of the present invention, further adaptations of the methods and systems described herein may be accomplished by appropriate modifications by one of ordinary skill in the art without departing from the scope of the present invention. Several of such potential modifications have been mentioned, and others will be apparent to those skilled in the art. For instance, the examples, embodiments, geometries, materials, dimensions, ratios, steps, and the like discussed above are illustrative and are not required. Accordingly, the scope of the present invention should be considered in terms of the following claims and is understood not to be limited to the details of structure and operation shown and described in the specification and drawings.
I/we claim:

1. A surgical instrument comprising:
   (a) a body having a firing mechanism configured to be selectively manipulated by an operator;
   (b) a shaft assembly extending distally from the body; and
   (c) an end effector extending distally from the shaft assembly and configured to receive a cartridge for manipulating tissue of a patient, the end effector including:
      (i) a distal end portion,
      (ii) a proximal end portion,
      (iii) a gap between the distal end portion and the proximal end portion, and
      (iv) a guide pin extending from the distal end portion to the proximal portion, wherein the guide pin is configured to receive the tissue thereagainst for positioning tissue within the gap, wherein the guide pin is connected to each of the distal and proximal end portions to secure the distal end portion relative to the proximal end portion and inhibit deflection of the distal end portion relative to the proximal end portion.

2. The surgical instrument of claim 1, wherein the distal end portion extends distally from the proximal end portion in the form of a hook-shape supporting structure with the gap therebetween configured to receive the tissue.

3. The surgical instrument of claim 1, wherein the guide pin is connected to each of the distal and proximal end portions in tension to secure the distal end portion relative to the proximal end portion.
4. The surgical instrument of claim 1, wherein the guide pin is in the form of a threaded fastener.

5. The surgical instrument of claim 1, wherein the distal end portion has a distal pin bore and the proximal end portion has a proximal pin bore, wherein the distal and proximal pin bores are positioned opposite each other and receive the guide pin therein.

6. The surgical instrument of claim 5, wherein the proximal pin bore includes a threaded nut, and wherein the guide pin is threaded into the threaded nut to be connected therein.

7. The surgical instrument of claim 6, wherein the guide pin is in the form of a threaded fastener.

8. The surgical instrument of claim 7, wherein the threaded fastener is connected to each of the distal and proximal end portions in tension to secure the distal end portion relative to the proximal end portion.

9. The surgical instrument of claim 1, further comprising a cartridge received against the guide pin, wherein the cartridge is configured to be guided from the proximal end portion toward the distal end portion along the guide pin.

10. The surgical instrument of claim 9, wherein the cartridge includes at least one of a knife or a plurality of staples, wherein the knife is configured to cut the tissue, and wherein the plurality of staples are configured to fasten the tissue.

11. The surgical instrument of claim 10, wherein the cartridge includes each of the knife and the plurality of staples.
12. The surgical instrument of claim 11, wherein the distal end portion of the end effector includes an anvil, and the guide pin is configured to inhibit deflection of the anvil relative to the proximal end portion.

13. The surgical instrument of claim 1, wherein the guide pin is a rigid guide pin.

14. An end effector of a surgical instrument for manipulating tissue of a patient with a cartridge, comprising:
   (a) a distal end portion;
   (b) a proximal end portion;
   (c) a gap between the distal end portion and the proximal end portion; and
   (d) a guide pin extending from the distal end portion to the proximal portion, wherein the guide pin is configured to receive the tissue thereagainst for positioning the tissue within the gap, wherein the guide pin is connected to each of the distal and proximal end portions to secure the distal end portion relative to the proximal end portion and inhibit deflection of the distal end portion relative to the proximal end portion.

15. The surgical instrument of claim 14, wherein the distal end portion extends distally from the proximal end portion in the form of a hook-shape supporting structure with the gap therebetween configured to receive the tissue.

16. The surgical instrument of claim 14, wherein the guide pin is connected to each of the distal and proximal end portions in tension to secure the distal end portion relative to the proximal end portion.

17. The surgical instrument of claim 14, wherein the guide pin is in the form of a threaded fastener.
18. The surgical instrument of claim 14, further comprising a cartridge received against the guide pin, wherein the cartridge is configured to be guided from the proximal end portion toward the distal end portion along the guide pin.

19. A method of manipulating tissue of a patient with a surgical instrument, the surgical instrument including a body having a firing mechanism configured to be selectively manipulated by an operator, a shaft assembly extending distally from the body, and an end effector extending distally from the shaft assembly and configured to receive a cartridge for manipulating the tissue of the patient, wherein the end effector has a distal end portion, a proximal end portion, a gap between the distal and proximal end portions, and a guide pin extending from the distal end portion to the proximal end portion, wherein the guide pin is configured to receive the tissue thereagainst for positioning the tissue within the gap, wherein the guide pin is connected to each of the distal and proximal end portions to secure the distal end portion relative to the proximal end portion and inhibit deflection of the distal end portion relative to the proximal end portion, the method comprising:

(a) receiving the tissue of the patient within the gap between the distal and proximal end portions of the end effector;
(b) positioning the tissue against the guide pin;
(c) inhibiting deflection of the distal end portion of the end effector relative to the proximal end portion of the end effector with the guide pin; and
(d) manipulating the firing mechanism to thereby manipulate the tissue of the patient.

20. The method of claim 19, further comprising increasing tension in the guide pin such that guide pin further inhibits deflection of the distal end portion of the end effector relative to the proximal end portion of the end effector.
Fig. 15
**INTERNATIONAL SEARCH REPORT**

**International application No**
PCT/US2016/066802

**A. CLASSIFICATION OF SUBJECT MATTER**

INVI. A61B17/072

According to International Patent Classification (IPC) into both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)
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 practicable, search terms used)
EPO-Internal, WPI Data

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tbody>
<tr>
<td>X</td>
<td>US 3 494 533 A (GREEN DAVID T ET AL) 10 February 1970 (1970-02-10) col umn 13, line 20 - line 70; figures 1,3,9,15,.31,.38</td>
<td>1-5,9-18</td>
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<tr>
<td>X</td>
<td>CN 103 083 053 A (SUZHOU TIANCHEN INT MED TECH) 8 May 2013 (2013-05-08) abstract; figures 1.2,5</td>
<td>1,2,5, 9-15, 18</td>
</tr>
<tr>
<td>X</td>
<td>WO 86/02254 AI (UNITED STATES SURGICAL CORP [US]) 24 April 1986 (1986-04-24) page 9, paragraph 2; figures 3-8</td>
<td>1-5,9-18</td>
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- Additional documents are listed in the continuation of Box C.
- See patent family 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 application or patent but published on or after the international filing date
  
  *"L"* document which may throw doubts on priority claim(s) or on the correctness of the claimed invention or on some parts thereof
  
  *"O"* document referred to in oral proceedings before the International Bureau

* Additional documents:
  
  *"X"* later document published after the international filing date and not in conflict with the application but cited to understand the principle or theory underlying the invention
  
  *"Y"* 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
  
  *"Z"* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is taken in combination with one or more other such documents, such combination being obvious to a person skilled in the art
  
  *"A"* document member of the same patent family

Date of the actual completion of the international search: 10 March 2017

Date of mailing of the international search report: 23/03/2017

Name and mailing address of the ISA/
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Fax: (+31-70) 340-3016

Authorized officer:
Roudaut, Tanguy

Form PCT/ISA/210 (second sheet) (April 2005)
<table>
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<th>Relevant to claim No.</th>
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<td>X</td>
<td>EP 2 248 474 A2 (TYCO HEALTHCARE [US]) 10 November 2010 (2010-11-10) paragraph [0078] - paragraph [0087]; figures 1-4, 22-26</td>
<td>1-5, 9-18</td>
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<tr>
<td>A</td>
<td>US 4 216 891 A (BEHLKE HAROLD D [US]) 12 August 1980 (1980-08-12) column 3, line 13 - column 4, line 19; figures 1-4</td>
<td>1-18</td>
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<tr>
<td>A</td>
<td>EP 1 550 410 A2 (ETHICON ENDO SURGERY INC [US]) 6 July 2005 (2005-07-06) paragraph [0025] - paragraph [0031]; figures 1, 6, 9, 28-36, 39, 40</td>
<td>1-18</td>
</tr>
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</table>
INTERNATIONAL SEARCH REPORT

Box No. 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.: 19, 20
   because they relate to subject matter not required to be searched by this Authority, namely:
   see FURTHER INFORMATION sheet PCT/ISA/210

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 No. 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 additional fees.

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 and, where applicable, the payment of a protest fee.

□ The additional search fees were accompanied by the applicant’s protest but the applicable protest fee was not paid within the time limit specified in the invitation.

□ No protest accompanied the payment of additional search fees.
Continuation of Box II.1

Claims Nos.: 19, 20

Claims 19 and 20 relate to a method for treatment of the human or animal body by surgery, practiced on the human or animal body (Rule 39.1 (iv) PCT), namely a method of manipulating tissue of a patient with surgical instrument. Consequently, no opinion will be formulated with respect to the novelty, inventive step and industrial applicability of the subject-matter of these claims (Article 17(2)(a)(i) PCT).
<table>
<thead>
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<th>Patent document cited in search report</th>
<th>Publication date</th>
<th>Patent family member(s)</th>
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