Abstract: A mounting tool (20) for mounting an anastomotic device (10, 200) to an end (220, 221) of a tubular structure (90) comprises a receiving portion for receiving a πigid part (11, 13) of said device (10, 200) and a conical portion with a large end facing said πigid part (11, 13) having a diameter larger than or equal to the rigid part (11, 13) and a small end. The conical portion comprises a releasable portion and is insertable into the elastic part (12) arranged inside the tubular structure (90) for expanding the diameter thereof and for passing the elastic part (12) beyond the large end and onto a rigid part (11, 13) arrangeable at the receiving portion.

A method for mounting a device (10, 200) to an end (120, 121) of a tubular structure (90) comprises arranging a πigid part (11, 13) of said device (10, 200) at a receiving portion of a mounting tool (20), inserting a small end of a conical portion of said mounting tool (20) into an elastic part (12) of said device (10, 200), wherein the elastic part (12) is arranged at the end (120, 121) of the tubular structure (90), and passing the elastic part (12) along said conical portion, thereby expanding the elastic part (12) until passing over the large end to be received in the πigid part (11, 13) arranged at the receiving portion.
A MOUNTING TOOL AND A METHOD FOR A DEVICE FOR ANASTOMOSIS

Technical field

The present invention relates to a mounting tool for a device for anastomosis of a living tissue, and more particularly to a mounting tool for a device for compression anastomosis of tubular structures. Furthermore, the invention relates to a method for mounting the device to a tubular structure by means of the mounting tool.

Background of the invention

Colorectal cancer is the third most frequent type of cancer in the world having an occurrence of about 1 million new cases every year. The incidents of cancer are considerably more frequent in the industrial part of the world.

Current techniques for mechanically performing anastomosis of hollow organs use circular mechanical staplers, which execute the connection of the tissue edges of the dissected hollow organ by metallic or plastic staples. A wide variety of surgical staplers have been developed for gastric, oesophageal and intestinal surgery. In performing surgical anastomotic stapling, generally two pieces of the hollow organ are joined by a ring of staples with a closed loopstapler. End to end anastomoses are generally performed by intraluminal surgical staplers that deliver a pair of staggered rings of staples. During this process, a circular knife blade is used to separate the tissue that is held within the circular ring. The separated tissue is then removed with the stapler to form a circular opening within the lumen along the stapling line.

A major issue regarding anastomosis healing is the blood circulation of the anastomosis during the healing process. Despite substantial development of surgical techniques during the last decades, morbidity and mortality after resections in the gastrointestinal tract, e.g. due to anastomotic leakage, remain as serious problems. Ischemia and inflammation, which are natural parts of the healing process, may cause leakage and secondary infection that may be fatal for the patient in the stapling area. Therefore, it has become common practice to relieve the pressure from the anastomosis by performing a deviating stoma, especially when the anastomosis is carried out in the lower part of colon and in rectum. By relieving pressure and faecal stream from the anastomosis during the healing process, the leakage incident may be reduced and fatal consequences of anastomotic dehiscence can be avoided. The inconvenience for the patient is obvious, since the patient must have a temporary stoma for a time period of about 3-6 months, and then has to undergo a second surgery in order to close the stoma. Unfortunately in many cases,
the closure of the stoma cannot be reversed and the patient is forced to live with a permanent stoma leading to lower quality of life associated with increased costs.

Another problem arising from stapling of anastomoses is anastomotic stenosis. The critical area for healing is the contact area between the two ends of the hollow structure to be connected. The connection has to be liquid proof, and the cross section of the lumen should be as wide and flexible as the original lumen. The size of the stapler determines the size of the lumen and thus the contact area between the ends. Surgical staplers create a smaller and more rigid opening compared to the cross section of the original lumen due to the staples inside the hollow structure connecting the two ends thereof, i.e. a collar may be formed that may lead to stenosis. For solving this problem repeated need for dilatation is required.

Another disadvantage associated with mechanical staplers, is that there is no fast, simple and reliable method to control anastomotic insufficiency, which at late discovery can result in abdominal sepsis.

Furthermore, staplers require an incision in the intestine in order to insert the instrument into the bowel lumen. This additional incision increases the duration of the operation and the risks associated with surgery, e.g. secondary infections and anastomotic leakage.

The stapler itself is a critical link, since there are several severe problems connected with the use of mechanical staplers in surgical anastomotic stapling, such as anastomotic leakage and anastomotic stenosis. Other disadvantages are high consumption of time and expensive instruments for the performance.

US 5 697 943 discloses a surgical instrument for carrying and attaching separate components of an anastomosis device to the end of tissue of a tubular hollow organ. The instrument includes an elongated housing having a proximal end and a distal end, first and second compression anastomosis device components, a supporting member operatively associated with the distal end of the housing, and an assembling structure associated with the supporting member and operable from the proximal end of the housing for assembling the first and second compression anastomosis device components within tubular body tissue.

**Disclosure of the invention**

A first object of the present invention is to eliminate at least one of the drawbacks mentioned above, which is achieved by assigning to the mounting tool the characteristics according to claim 1.

A second object of the invention is to provide a method for mounting a device to a tubular structure by means of the mounting tool.
A third object of the invention is to provide a method for performing an anastomosis.

According to a first aspect of the invention, there is provided a mounting tool for mounting an anastomotic device to an end of a tubular structure, comprising a receiving portion for receiving a rigid part of said anastomotic device, and a conical portion with a large end facing said rigid part having a diameter larger than or equal to the rigid part and a small end insertable in an elastic part arranged inside the tubular structure. The conical portion comprises a releasable portion, whereby said conical portion is insertable into the elastic part for expanding the diameter of the elastic part and for passing said elastic part beyond the large end and onto a rigid part arrangable at the receiving portion. The conical portion comprises a top cone portion a truncated cone portion comprising several cone segments. The top cone is integral with a central axle threaded at one end of the mounting tool, and is forming a first end of the tool. A handle is forming a second end of the tool. The tool further comprises a cylinder having a bore and an end-hat having a threaded bore for locking the handle. The receiving portion comprises a profiled section at the large end of the truncated cone. The truncated cone is formed by arranging the side cone segments on the axle between the top cone abutting the proximal end thereof and the rigid part abutting the cylinder, which is fixed by the handle. The top cone and the cone segments are made of a polymeric material. The axle and the cylinder are made of a metal. According to a second embodiment of the invention the axle of the mounting tool is provided with recesses to accommodate catheters provided at holes arranged around the periphery of one rigid part of the device.

According to a second aspect of the invention, there is provided a method for mounting a device to an end of a tubular structure, comprising arranging a rigid part of said device at a receiving portion of a mounting tool, inserting a small end of a conical portion of said mounting tool into a hollow elastic part of said device, said elastic part being arranged at said end of said tubular structure, said conical portion having a large end facing said rigid part and having a diameter larger than or equal to the receiving portion, and passing said elastic part along said conical portion, thereby expanding the elastic portion until passing over the large end to be received in said rigid part arranged at the receiving portion. The method further comprises removing a releasable portion of said conical portion for making free the interconnected elastic and rigid parts and for removing the mounting tool from the tubular structure.

According to a third aspect of the invention there is provided a method of performing an anastomosis, comprising arranging rigid and elastic parts at a first and a second end of a tubular structure to be interconnected by the anastomosis, passing said ends towards each other, engaging a connection member for the
interconnection of the rigid parts each having an elastic part arrange thereto of each end of the tubular structure for arranging outer surfaces of said tubular structure of each end into contact with each other supported by said elastic parts of each end, and forming a necrosis line adjacent said contact area for initiating tissue growth by squeezing a portion of said tubular structure between said elastic and rigid parts. A pressure is exerted between the ends of the tubular structure, when the ends are arranged into the interlocked rigid and elastic parts, which is essential uniform around the periphery of the ends of the tubular structure.

Further objects, features and advantages of the present invention will appear from the following detailed description, from the attached drawings as well as from the dependent claims.

**Brief description of the drawings**

In order to explain the invention, a number of embodiments of the invention will be described below with reference to the drawings, in which:

Fig 1A is a schematic exploded perspective view of an anastomotic device according to a first embodiment,

Fig 1B is a schematic perspective view of a first rigid part of the device in Fig 1A provided with catheters,

Fig 2A is a schematic perspective view of a mounting tool according to a first embodiment for mounting a rigid part of the device in Fig 1A to a tubular structure,

Fig 2B is a schematic perspective view of a mounting tool according to a second embodiment for mounting the rigid part in Fig 1B to a tubular structure,

Fig 3A is a cross sectional view of a first rigid part of the device in Fig 1A,

Fig 3B is a schematic perspective view of the first rigid part in Fig 3A provided with catheters,

Fig 4A is a cross sectional view of a second rigid part of the device in Fig 1A,

Fig 4B is a schematic perspective view of the second rigid part of the device in Fig 4A,

Fig 5A is a perspective view of an axle and a top cone of the mounting tool according to the first embodiment before mounting side cone segments, and before arranging a rigid part thereto,

Fig 6A is a perspective view of the axle and the top cone in Fig 5A having the side cone segments mounted in position and the rigid part arranged thereto,

Fig 7A a perspective view of the mounting tool according to the first embodiment having a rigid part mounted thereto and ready for use,
Fig 8A shows a cross sectional view of a truncated cone comprising a receiving portion to receive a rigid part of the device, before the arrangement of the rigid part,

Fig 5B is a perspective view of an axle and a top cone of the mounting tool according to the second embodiment before mounting side cone segments, and before arranging a rigid part provided with catheters thereto,

Fig 6B is a perspective view of the axle and top cone in Fig 5B and the catheters of the rigid part arranged thereto, before mounting side cone segments in position,

Fig 7B is a perspective view according to Fig 6B after mounting the side cone segments in position and arranging the rigid part thereto,

Fig 8B is a perspective view of the mounting tool according to the second embodiment having a rigid part arranged thereto and provided with a guide before mounting an end tip to the guide,

Fig 9 is a schematic perspective view of an elastic part arranged inside a tubular structure at one end thereof, the edge of this end being folded over the elastic part,

Fig 10 is a schematic perspective view according to Fig 9 showing the mounting tool according to the second embodiment inserted into the tubular structure before mounting the elastic part on the first rigid part,

Fig 11 is a schematic perspective view according to Fig 10 showing the elastic part mounted on the first rigid part,

Fig 12 is a schematic perspective view showing a first member of the device arranged at one end of the tubular structure,

Fig 13 is a schematic perspective view showing the first member arranged at one end of the tubular structure and a second member arranged at the other end of the tubular structure,

Fig 14 is a schematic perspective view according to Fig 13 when the first and second members are interlocked,

Fig 15 is a schematic perspective view showing the device passing through the tubular structure when released after healing up the ends thereof,

Fig 16A is a side view of a device according to a second embodiment without the elastic parts mounted thereto,

Fig 16B is a cross sectional view along line A-A of the device in Fig 16A,

Fig 17 is a cross sectional view of a portion of the device in Fig 16A before interlocking the first and second rigid parts,

Fig 18 shows the device in Fig 17 after interlocking the first and second rigid parts,
Fig 19 is a cross sectional view of a device according to an alternate embodiment.

Fig 20A is a schematic cross sectional view showing the device according to the first embodiment provided with catheters and arranged to the tubular structure fusing the ends thereof, and

Fig 20B is an enlarged view of the encircled area in Fig 20A showing the contact area and the point of necrosis.

Same reference numerals have been used to indicate the same parts in the figures to increase the readability of the description and for the sake of clarity.

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Description of embodiments of the invention

Dehiscence of intestinal anastomosis is associated with high morbidity and mortality rates. Rapid and effective wound healing of intestinal anastomosis is critical for a safe and prompt recovery of patients, which are undergoing anastomotic surgery. Wound healing is a relatively stereotypical tissue reaction that follows a predictive sequence of events including acute inflammation, proliferation (cell division and matrix protein synthesis), and over time tissue remodeling for adaptation of the new tissue to the mechanical demands.

The unique property of the anastomotic device is that the wound healing process is initiated by local tissue ischemia and necrosis resulting in that previously intact tissue, i.e. the intestinal serosa of each of the divided segments of the tubular structure that are joined together by the anastomotic ring, fuse by a healing response initiated by the construction of the anastomotic device. It should be noted that in the abdominal cavity of a healthy living, the serosa of one segment of the intestine is in constant contact with the serosa of other segments of the intestine without being fused together, which otherwise would be potentially dangerous, e.g. if causing intestinal obstruction.

In contrast, the pressure maintained by the device creates a circumferential area of ischemia in the intestine, which provokes a tissue response leading to the healing process that seals together the two divided serosal sides of the intestine, and further leading to completion of the anastomosis.

On a molecular level, the ischemia provokes a local inflammatory reaction, including the recruitment of leukocytes and tissue oedema. Thus, local cells, such as macrophages and mast cells, which secrete cytokines and chemokines that provoke the extravasations of leukocytes into healing area, detect necrotic cells. Moreover, the local ischemia per se also stimulates macrophages and mast cells to secrete pro-inflammatory mediators.

Leukocyte recruitment is critical for the compression induced healing of intestinal anastomosis due to the compounds released by accumulated inflammatory
cells, including radical oxygen species and metalloproteinase (MMP), which are necessary for the breakdown of the ischemic tissue as well as for breaking the integrity of the serosal surface.

MMPs are necessary for the breakdown of matrix proteins; especially important is collagen type I, which is abundant in the intestinal submucosa, but also collagen of type III and V. The inflammatory cells predominate secrete MMP-2 and MMP-9, which both are active in the early phase of the compression induced healing of intestinal anastomoses.

The breakdown of the ischemic tissue is important for the tissue disintegration and release of the intestinal segment from the device. In addition, the serosal surface (visceral peritoneum) contains numerous mesothelial cells, which also respond to the ischemia by secreting potent MMPs and angiogenetic factors.

The broken integrity of the serosal surface is a precondition for the unique healing process induced by the device knowing that the serosal side of the intestines normally exerts anti-adhesive functions avoiding adhesions between intestinal segments in the abdominal cavity. In parallel, intestinal fibroblasts and epithelial cells respond to the ischemic necrosis induced by the CARP-device by secreting angiogenetic factors, including vascular endothelial growth factor, which stimulates the growth of new blood vessels in the healing tissue.

Moreover, in this complex reaction, intestinal fibroblasts and epithelial cells also secrete metalloproteinases (MMP-I, MMP-7 and MMP-10), which contribute to the breakdown of matrix proteins and resolution of the tissue paving the way for the compression induced healing of anastomoses.

Fig 1A shows an exploded view of an anastomotic device 10 according to a first embodiment, which is to be mounted to a tubular structure 90 by means of a mounting tool 20, 21 according to the invention illustrated in Fig 2A, 2B. The device 10 comprises a first member comprising a first rigid part 11 and a first elastic part 12, a second member comprising a second rigid part 13 and a second elastic part 12, a sealing 14 and a connection member 15. The rigid parts 11, 13 and the elastic parts 12 have a generally hollow open configuration.

According to the first embodiment of the device 10, the connection member 15 is integral with the first rigid part 11. The first member and the second member are interlockable to each other, as a male-female component, by the connection member 15.

Fig 1B shows the first rigid part 11 provided with four catheters 16 mounted to four holes 17 (see Fig 3B) symmetrically arranged around the annular wall of the first rigid part 11 opening at the outer side thereof, forming a free passage through each catheter, which further elongates through the hole.
The rigid parts 11, 13 and the elastic parts 12, 14 are made of a polymeric material, more specifically a biocompatible material and most specifically a biodegradable material.

The elastic parts 12, 14 have a cross section, which may be of any shape, for example circular, elliptic, rectangular or flat, and has a diameter of about 2 to 9 mm, or more specifically 4 to 8 mm, or most specifically 5 to 7 mm. The elastic parts 12, are substantial circular symmetric rings and are made as a compact body or as a tube, e.g. filled with air, gas or fluid, and are made of an elastic polymeric material of for example 40 to 70 Shore. The elastic parts 12 have an inner diameter that is smaller than the smallest outer diameter of the rigid parts 11, 13 as will be explained below.

Fig 2A shows a mounting tool 20 according a first embodiment, and Fig 2B shows a mounting tool 21 according to a second embodiment. The mounting tools 20, 21 are required for arranging the device 10 to a tubular structure, which will be described in detail below.

As shown in Fig 3A, the first rigid part 11 according to the first embodiment has a substantially cylindrical inner surface, which is flared out at a non-connecting end 30. The outer surface is profiled and forms a partly semi-circular surface between the non-connecting end 30 and a connecting end, and the diameter at the non-connecting end 30 is larger than or equal to the diameter at the connecting end. A radial surface is provided at the connecting end delimiting the partly circular contour. The edge 35 of the non-connecting end 30 is slightly bevelled, and the surface 34 ends in an edge 36 connecting the surface 34 with the semi-circular surface.

The connection member 15 is integral with the first rigid part 11, as shown in Fig 3A, and is provided with transversal slits 37 from the free end thereof along the periphery forming tongues 38 between the slits 37. At least one of the tongues 38 is provided with an outward protrusion 39 arranged adjacent or at a distance from the free ends of the tongues. The number of slits 37 may vary, and the length of the slits 37 can be as long as the entire width of the connection member 15, as illustrated in Fig 3A, or be shorter. The slits 37 are either symmetrical or unsymmetrical provided around the periphery forming tongues with a similar or varying width. Fig 3B shows as an example, the connection member having six tongues without protrusions and six tongues with protrusions, arranged symmetrical around the periphery, wherein the tongues with protrusions have a width smaller than the tongues without protrusions. The number of tongues can vary, for example 2-10 may be provide, which can be arranged symmetrical or unsymmetrical around the periphery. The connection member 15 has a circumferential recess 31 around its outer diameter to accommodate the sealing 14.
The second rigid part 13 has a substantially cylindrical inner surface that is flared out at a non-connecting end 40, as seen in Fig 4A. The outer surface is profiled and forms a partly semi-circular surface between the non-connecting end 40 and a connecting end 41, the outermost edge 47 at the non-connecting end 40 being slightly bevelled. The diameter at the non-connecting end 40 is larger than or equal to the diameter at the connecting end 41. A bevel 42 is provided at the connecting end 41 to facilitate insertion of the connection member 15 into the second rigid part 13, as will be explained below. A surface 44 is provided at the connecting end 40 due to the partly circular contour. The surfaces 34 and 44 are equal in size and ends in the edge 36 and an edge 46, respectively. Grooves 45 or threads can be provided on the internal annular surface of the second rigid part 13, for example at the portion adjacent the connecting end 41, to obtain a locking function when interlocking the first member and the second member by the connection member 15.

In an alternative embodiment, the first rigid part has a circumferential notch around the circumference, as shown in Fig 16B.

With reference to Fig 5A, the mounting tool 201 according to the first embodiment comprises a rigid rod 50 forming a central axle having a top cone 51 at one end and provided with threads 52 at the other end. The top cone has a closed distal end and is integral with the axle. The tool 20 further comprises a truncated cone which is formed by several side cone segments 54, for example 6-8 segments. The truncated cone has a small diameter end abutting the top cone and a broad diameter end having a profiled section 55. Furthermore, the mounting tool 20 comprises a cylinder 56 and a handle 57 with an end-hat 58. The elements of the tool 20 are coaxially held together by the rod due to their symmetrically arranged bores. The mounting tool 20 comprises a receiving portion 48” for receiving the rigid parts 11, 13 comprising the profiled section of the large end of the truncated cone, which is dimensioned to receive the non-connecting end 30, 40 having a bevelled edge 48, as shown in Fig 8A for the rigid part 13. The tool 20 also comprises a conical portion comprising the top cone and the truncated cone including the cone segments. The conical portion has a releasable portion comprising the cone segments. Fig 5A shows a rigid part 11, 13 that is to be arranged at the receiving portion, and cone segments, which are to be arranged on the central axle between the top cone and the rigid part for forming a circular cone having a smooth transition to the rigid part, as shown in Fig 6A. The broad diameter end of the truncated cone is equal or slightly wider than the outer diameter of the rigid part. The cylinder has a recess (not shown) on the side facing the rigid part to accommodate the connection member 15 when this is integral with the first rigid part 11. The side cone segments between the top cone and the rigid part abutting the
cylinder are firmly fastened in position, since all the elements mentioned are held together by the handle, which is non-rotational fixed by the end-hat. When the mounting tool 20 is used to arrange the second rigid part 13, the cylinder is turned 180° around the rod so that the circular recess is facing the handle.

In an alternate embodiment, the cylinder may be integral with the handle. In another alternate embodiment, a cylindrical section may be provided for the arrangement of the side cone segments around the rod.

The mounting tool 21 according to a second embodiment is used for mounting a device 10 having a first rigid part 11 of the first embodiment provided with catheters 16. The mounting tool 21 of the second embodiment differs from the first one 20A in that the central axle 50' is provided with recesses 59 symmetrically arranged along the axle at the distal end, and the recesses are elongating through the top cone to each accommodate one catheter, as shown in Fig 5B and 6B. The recesses are opening into a hole 60 at the upper part of the top cone 51', where the catheters 16 will exit, see Fig 7B. A flexible annular guide 61 with an end-tip 62 is provided and arranged at the hole to accommodate and stabilize the catheters, as shown in Fig 8. The rest of the description of the mounting tool 21 according to the second embodiment is similar to the description above of the mounting tool 20 according to the first embodiment.

The rod is for example made of steel. The top cone and the cone segments are for example made of a polymeric material, stainless steel or another metallic material. The rest of the elements of the tool 20 are made of a polymeric material, stainless steel or another metallic material.

Reference is made to Fig 9-11 for describing the mounting of the device 10 to a tubular structure 90 having a first end 120 and a second end 121, which are to be fused together. In the example described, the device 10 has a first rigid part 11 provided with catheters 16, hence the tool 21 according the second embodiment is used for performing the mounting. The rigid part 11 is initially arranged at the receiving portion of the tool 21 having the connection member 15 arranged into the circular recess of the cylinder 56, as described above. The elastic part 12 is arranged inside the tubular structure 90 at one end 120, and the edge of this end 120 is folded over the elastic part 12, as shown in Fig 9. The tool 21, with the rigid part 11 arranged into the profile 48" of the truncated cone and with the catheters arranged into the recesses 59 along the axle 50' and elongating through the guide 61, is inserted into the tubular structure 90 from the folded end 120, as shown in Fig 10, expanding the elastic part 12, which is surrounded or wrapped on three sides by the tubular structure 90, until it is snapped around the circumference of the rigid part 11. The elastic part 12 is then locked to the rigid part 11 due to its partly semi-circular contour forming a recess, as illustrated in Fig 11. This snapping-operation
requires of an operator to keep firm hold of the end 120 with the elastic part 12 arranged thereto by one hand and of the tool 21 by the other hand. For releasing the first member 11 from the tool 21, firstly, the end-hat 58 is released. In order to prevent that the tool 21 and the rigid part 11 rotates, damaging the tubular structure, it is necessary to keep a firm grip of the handle 57, which is locked for rotation in relation to the axle 50’due to plane surface on the axle at the threading, then the handle is removed from the axle. Thereafter the cylinder 56 is released and removed. Now when the axle 50’ is free the truncated cone comprising the side one segments 54 is falling apart and are removed by a pair of forceps through the opening formed due to the rigid part 11. Then, the first member is pushed off the axle 50’ over the top cone.

The operation described is repeated for arranging the second rigid part 13 to the other end 121 of the tubular structure 90 by using either the mounting tool of the first 20 or second 21 embodiments. Thus, the first and second members of the device 10 is formed with the ends 120, 121, respectively, having the tubular structure 90 arranged between the rigid parts 11, 13 and the elastic parts 12. Thus, a single layer of the tubular structure 90 is squeezed between a rigid part 11, 13 and an elastic part 12.

The extremity of the first end 120, when folded and arranged between the first rigid part 11 and the first elastic part 12, forms a first lip or a contact surface 130 that is essentially circular, as seen in Fig 13. Similarly, the second end 121 forms a second lip or contact surface 131. An essentially uniform and equal pressure is exerted on the contact surfaces 130, 131 all over the periphery.

The final action for forming the device 10 is to connect the first member and the second member by the connection member 15, which is performed by a simple press action by hand by the operator. In the first embodiment the connection member 15 is integral with the first rigid part 11, and the first member including the connection member and the second member are joined together by interlocking the members by hand. The tongues 38, of which some or all, have protrusions 39 arranged on the connection member 15 interact with the grooves 45 arranged on the inner surface of the second rigid part 13 and perform a locking action. Thus, the connection member 15 engages the second rigid part 13, and the first and second members are fixedly mounted to each other.

The first and second members are dimensioned so that a gap appears between the surfaces 34 and 44 (see Figs. 3 and 4) when the members are interlocked without arrangement of the two ends 120, 121 of the tubular structure 90 between the rigid parts 11, 13 and the elastic parts 12, 14, respectively. When interlocking the first member and the second member with the tubular structure 90 arranged thereto, the gap still exists between the surfaces 34 and 44. The contact surfaces
130, 131 are pressed against each other and the end 120, 121 of the tubular structure
90 are compressed between the elastic part 12, 14 and the rigid parts 11, 12,
respectively, resulting in that the ends 120, 121 will slightly creep into the gap and
hang over the edges 44 and 36, respectively.

A contact area 201 is created between the contact surfaces 130 and 131, and
a point of necrosis 202, or rather line of necrosis, is defined as the point where the
tubular structure 90 is pressed against the edges 45 and 36 by the pressure from the
elastic parts 12, 14, respectively, as seen in Fig 20b. The contact area 201 and the
point of necrosis 202 have a major importance in the necrosis process.

A second embodiment of an anastomotic device 200 will now be described
with reference to Fig 16A, 16B, 17, and 18. The device 200 comprises a first
member, a second member and a connection member 160, each of a hollow open
configuration. The first and second members are identical, each comprising a rigid
part 13 and an elastic part 12, wherein the rigid parts 13 have the same
configuration as the second rigid part 13 according to the first embodiment of the
device 10. The connection member 160 is made as a substantially cylindrical
separate piece, i.e. separated from the first and second member, and has notches 170
(see Fig 17) arranged on the outer surface thereof for engagement with the grooves
45, or threads, provided on the internal cylindrical surface of the rigid parts 13. The
connection member 160 is dimensioned for interlocking the first and second
member when arranged inside the rigid parts 13, as illustrated in Fig 18. Recesses
172 may be provided, for example symmetrically, on the outer surface of the
connection member 160, each to accommodate a circular sealing member (not
shown), such as an O-ring. The connection member 160 has optionally at least one
hole 161 passing through the annular wall of the connection member 160, for
example at the centre symmetry line. At least one nipple (not shown) can engage
this/these hole(s) 162 for the connection of at least one tube or catheter, which will
be explained below. Fig 17 shows the first and second rigid part 13 and the
connection member 160 before interlocking. Fig 18 shows the identical first and
second rigid parts 13 interlocked by the connection member 160. A distance or a
gap 180 is provided between the surfaces 87 of the rigid parts 13.

The connection member 160 is made as a substantially cylindrical separate
device piece, i.e. separated from the first and second member, and has notches 171 (see Fig
17) arranged on the outer surface thereof for engagement with the grooves or
threads 171 provided on the inner annular surface of the rigid parts 13. The
connection member 160 is dimensioned for interlocking the first and second
member when arranged inside the rigid parts 13, as illustrated in Fig 18. Recesses
172 are provided, for example symmetrically, on the outer surface of the connection
member 160, each to accommodate a circular sealing member (not shown), such as
an O-ring. The connection member 160 has optionally at least one hole 161 passing through the annular wall of the connection member 160, for example at the centre symmetry line. At least one nipple (not shown) can engage this/these hole(s) 162 for the connection of at least one tube or catheter, which will be explained below. Fig 5 shows the first and second rigid part 13 and the connection member 160 before interlocking. Fig 18 illustrates the rigid parts 13 locked to each other by the connection member 160. A distance or a gap 180 is provided between the surfaces 44 of the rigid parts 13.

The pressure that is exerted on the contact surfaces 130, 131 at the ends 120, 121 of the tubular structure 90, when arranged in the device 10, 200, can be increased or decreased by adjusting the size of the gap 203, 180. In the case with the device of the first embodiment 10, the gap 203 may be varied by arranging the notches 39 on the connection member 15 closer or less closer to the free ends of the tongues 38. When using the device according to the second embodiment 200, the gap 180 may be varied by arranging the connection member 160 deeper or less deep inside the rigid parts 13 in relation to a reference position. The hollow open configuration of the first and second members and the connection member 160 are shown as being essentially circular from a side view, see Fig 18, but can be of any other shape, e.g. elliptic, oval, partly rectangular or triangular, or can be segmented as shown in Fig 19.

Fig 20A illustrates the device 10 of the first embodiment arranged to a tubular structure 90. The catheters 16 are arranged to the holes 17 and are elongating through the tubular structure 90. The first ends of the catheters 16 that are arranged to the holes are opening into the cavity 203, and the second ends of the catheters 16 are exiting through rectum. The guide 61 is removed when the second ends of the catheters have exit. These ends of the catheters can be connected to for example syringes, pumps or other devices used for supervising or controlling the healing up of the ends of the tubular structure. It is also possible to supply different fluids to the cavity 203, such as grow stimulating substances or contrast media, as will be explained below.

The tubular structure 90 will swell up at the ends 120, 121, thus closing the gap between the surfaces 34 and 44., forming a closed cavity 203 defined by the compressed tubular structure 90, the connection member 15 and the surfaces 34 and 44 of the rigid parts 11, 13. A contact area 201 is created between the contact surfaces 130, 131, and a point or line of necrosis 202, which is defined as the point 202 where the tubular structure 90 is pressed against the edges 36 and 46 by the pressure from the elastic part 12. The blood stream or circulation at the ends 120, 121 of the tubular structure 90 is cut off (strangled) and ceases right up to the point of necrosis 202. Tissue regeneration takes place at the contact area 201, which will
result in fusing together the two ends 120, 121 of the tubular structure 90. When the two ends of the tubular structure are fused, i.e. have healed up, the device 10, 200 is automatically released and leaves the tubular structure following the faecal stream through rectum. Fig 20A shows the catheters 16 fastened to the device 10 and arranged inside and along the tubular structure 90. By means of the guide 61, the ends of the catheters are passing out of the tubular structure 90 through rectum.

It is possible to supply different fluids to the cavity 203 by means of the catheters 16. For example, a certain selected liquid may be supplied to the cavity 203, in a continuous or an intermittent flow. The liquid can be tissue growing stimulating, or perform any other actions for accelerating the healing, such as stimulating the recruitment of leukocytes and/or secretion of cytokines and chemokines.

Furthermore, contrast medium may be supplied through the catheters 16 for performing a radiological control of the anastomosis, for example regarding closeness or contrary leakage, which is especially important directly after mounting the device to the tubular structure. By supplying a certain pressure of the supplied liquid to one catheter, while keeping the other catheters closed, a measure of the pressure for leakage is obtained. In this manner, a possibility is provided to continuously supervise the healing process of the two ends of the tubular structure. In case of a minor leakage of the anastomosis, it is possible to apply a sub-pressure or slight vacuum to the cavity 203 by simply connecting one catheter to a vacuum air pump (not shown).

Herein above, several embodiments of the invention are described with reference to the drawings in order to enable a skilled person to perform the invention. However, the features and method steps included in these embodiments do not limit the invention. Moreover, the features and method steps may be combined in other manners than specifically described.

The cross sections of the elastic parts 12 are shown as substantially circular. However, other shapes may be used, such as rectangular, triangular, hexagonal, octagonal etc. The outer surface of the rigid parts 11, 13 comprises a recess intended to receive the elastic parts 12 respectively. This recess has a shape that is at least partly complementary to the shape of the elastic part 12. Thus, the recess may be rectangular, triangular, hexagonal, octagonal, etc.

The configuration or outer shapes of the elastic parts 12 are shown to be substantially cylindrical having a circular outer contour. However, other shapes are possible, such as rectangular, triangular, hexagonal, octagonal etc.

The elastic parts 12 can be only partially elastic. The elasticity is used for clamping the tubular structure 90 between the elastic part 12 and the rigid part 11.
13 with a certain force. Other means performing the same function is possible to use.

In an alternate embodiment, the top cone may be snapped onto the central axle, as well as the end-hat of the handle. Furthermore, the truncated cone can be made as a whole piece, which can be released from the axle through rectum at low anastomoses.

In the claims, the term "comprises/comprising" does not exclude the presence of other elements or steps. Furthermore, although individually listed, a plurality of means, elements or method steps may be implemented. Additionally, although individual features may be included in different embodiments, these may possibly be combined in other ways, and the inclusion in different embodiments does not imply that a combination of features is not feasible. In addition, singular references do not exclude a plurality. The terms "a", "an" does not preclude a plurality.

Reference signs in the claims are provided merely as a clarifying example and shall not be construed as limiting the scope of the claims in any way.
Claims

1. A mounting tool (20, 21) for mounting an anastomotic device (10, 200, 190) to an end (120, 121) of a tubular structure (90), characterized by
   a receiving portion (48") for receiving a rigid part (11, 13, 191, 193) of said anastomotic device (10, 200);
   a conical portion with a large end having a diameter larger than or equal to the rigid part (11, 13, 191, 193) and a small end insertable in an elastic part (12) arranged inside the tubular structure (90);
   said conical portion comprising a releasable portion, whereby said conical portion is insertable into the elastic part (12) for expanding the diameter of the elastic part (12) and for passing said elastic part (12) beyond the large end and onto a rigid part (11, 13, 191, 193) arrangeable at said receiving portion.

2. The mounting tool (20, 21) according to claim 1, characterized in that the conical portion comprises a top cone (51) and a truncated cone.

3. The mounting tool (20, 21) according to claim 1 or 2, characterized in that the truncated cone comprises one or several cone segments (54).

4. The mounting tool (20, 21) according to claim 2, characterized in that the top cone (51) is integral with a central axle (50) of the mounting tool (20, 21).

5. The mounting tool (20, 21) according to any of the previous claims, characterized in that the central axle is threaded at one end (52).

6. The mounting tool (20, 21) according to any one of the previous claims, characterized in that the top cone (51) is forming a first end of the tool (20, 21) and a handle (57) is forming a second end of the tool (20, 21).

7. The mounting tool (20, 21) according to any one of the previous claims, characterized in that further comprising a cylinder (56) having a bore for arrangement on the axle (50, 50'), and an end-hat (58) having a threaded bore for locking the handle to the axle (50, 50').

8. The mounting tool (20, 21) according to claim 7, characterized in that said receiving portion (55) is a profiled section arranged at the large end of the conical portion.

9. The mounting tool (20, 21) according to any one of the previous claims, characterized in that the truncated cone is formed by arranging the side cone segments (54) on the central axle (50, 50') between the top cone (51) abutting the proximal end thereof and the rigid part (11, 13, 191, 193) abutting the cylinder (56), which is fixed by the handle (57).
10. The mounting tool (20, 21) according to any one of the previous claims, characterized in that the top cone (51) and the cone segments (54) are made of a polymeric material.

11. The mounting tool (20, 21) according to any one of the previous claims, characterized in that the axle (50, 50’) and the cylinder (56) are made of a metallic or polymeric material.

12. The mounting tool (21) according to any one of the previous claims, characterized in that the axle (50, 50’) are provided with recesses (59) which are elongating through the top cone (51).

13. The mounting tool (21) according to claim 12, characterized in that recesses are opening into a hole (60) at the upper part of the top cone.

14. A method for mounting a device (10, 200) to an end (120, 121) of a tubular structure (90), comprising:
   arranging a rigid part (11, 13) of said device (10, 200, 190) at a receiving portion of a mounting tool (20);
   inserting a small end of a conical portion of said mounting tool (20) into a hollow elastic part (12) of said device (10, 200), said elastic part (12) being arranged at said end (120, 121) of said tubular structure (90), wherein said conical portion has a large end which is larger than or equal to said rigid part (11, 13, 193);
   passing said elastic part (12) along said conical portion, thereby expanding the elastic part (12) until passing over the large end to be received in said rigid part (11, 13) arranged at the receiving portion.

15. The method of claim 14, further comprising:
   arranging catheters (16) along recesses (59) of an axle (50, 50’) of said mounting tool (20, 21) each catheter (16) arranged with a first end to holes (17) provided around the periphery of the rigid part (11, 160, 192) of the device (10, 200) and with a second end arranged along the tubular structure exiting through rectum.

16. The method of claim 14 or 15, further comprising:
   removing a releasable portion of said conical portion for releasing the interconnected elastic (12) and rigid (11, 13) parts from the mounting tool and for removing the mounting tool (20) from the tubular structure (90).

17. A method for performing an anastomosis, comprising:
   arranging rigid (11, 13, 191, 193) and elastic (12) parts of a device (10, 200) at a first end (120) and a second end (121) of a tubular structure (90) to be interconnected by the device (10, 200);
   passing said ends (120, 121) towards each other;
   engaging a connection member (15, 160) for the interconnection of the rigid parts (11, 13) each having an elastic part (12) arranged thereto of each end (120,
121) of the tubular structure (90) for arranging outer surfaces (130, 131) of said tubular structure (90) of each end (120, 121) into contact with each other supported by said elastic parts (12) of each end (120, 121);

forming a necrosis point (202) adjacent said contact area (201) for initiating tissue growth by squeezing a portion of said tubular structure (90) between said elastic (12, 14) and rigid (11, 13) parts.

18. The method according to claim 17, characterized by exerting a pressure between the ends (120, 121) of the tubular structure (90) when the ends (120, 121) are arranged into the interlocked rigid (11, 13) and elastic (12) parts.

19. The method according to claim 18, characterized in that the pressure is essential uniform around the periphery of the ends (120, 121) of the tubular structure (90).
Fig 2A

Fig 2B