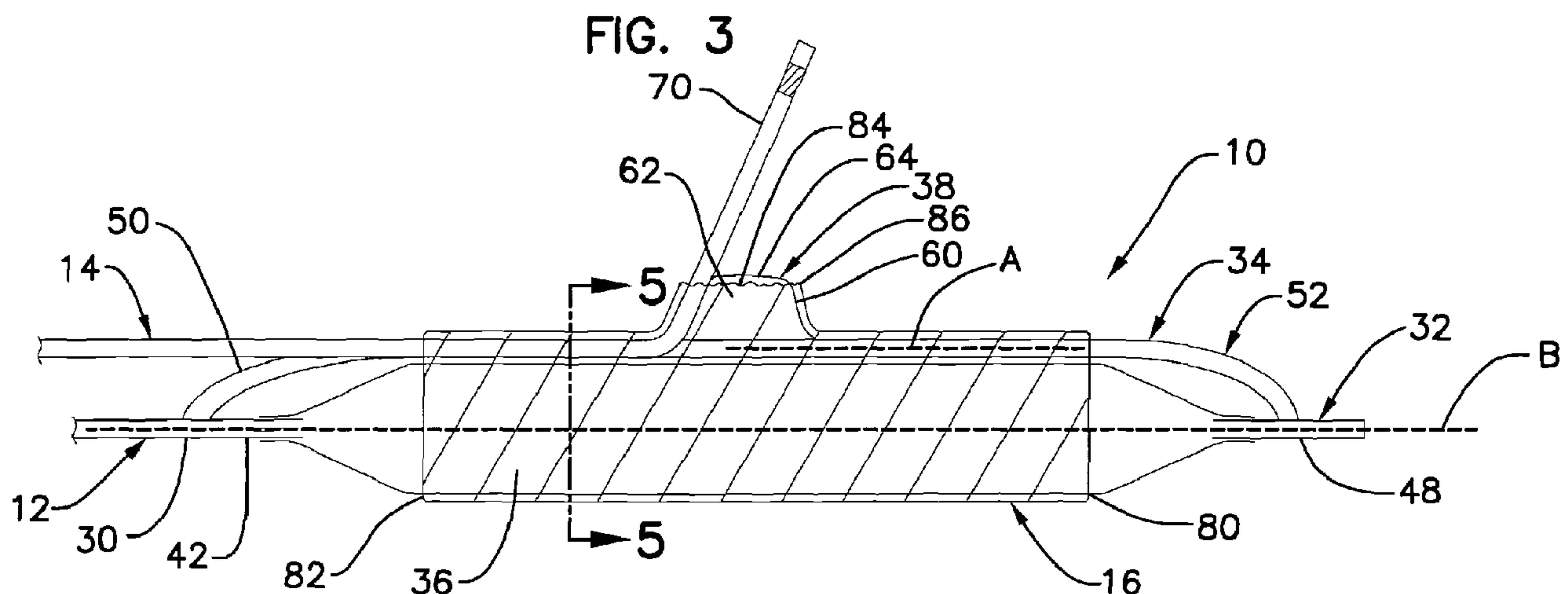




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(57) **Abrégé/Abstract:**

A catheter assembly (10) includes a main catheter branch (12) and a side catheter branch (14). The main catheter branch includes a main balloon (36), a side balloon (38) and a side inflation member (34). The side inflation member intersects the side balloon at a location on the side balloon that is offset laterally from a central line passing from a distal most point on the side balloon to a proximal most point on the side balloon. The side balloon is configured to extend radially outward relative to the main balloon when the side balloon is inflated. The side catheter branch can be centrally aligned with the side balloon central line and be positioned laterally adjacent to the side inflation lumen.



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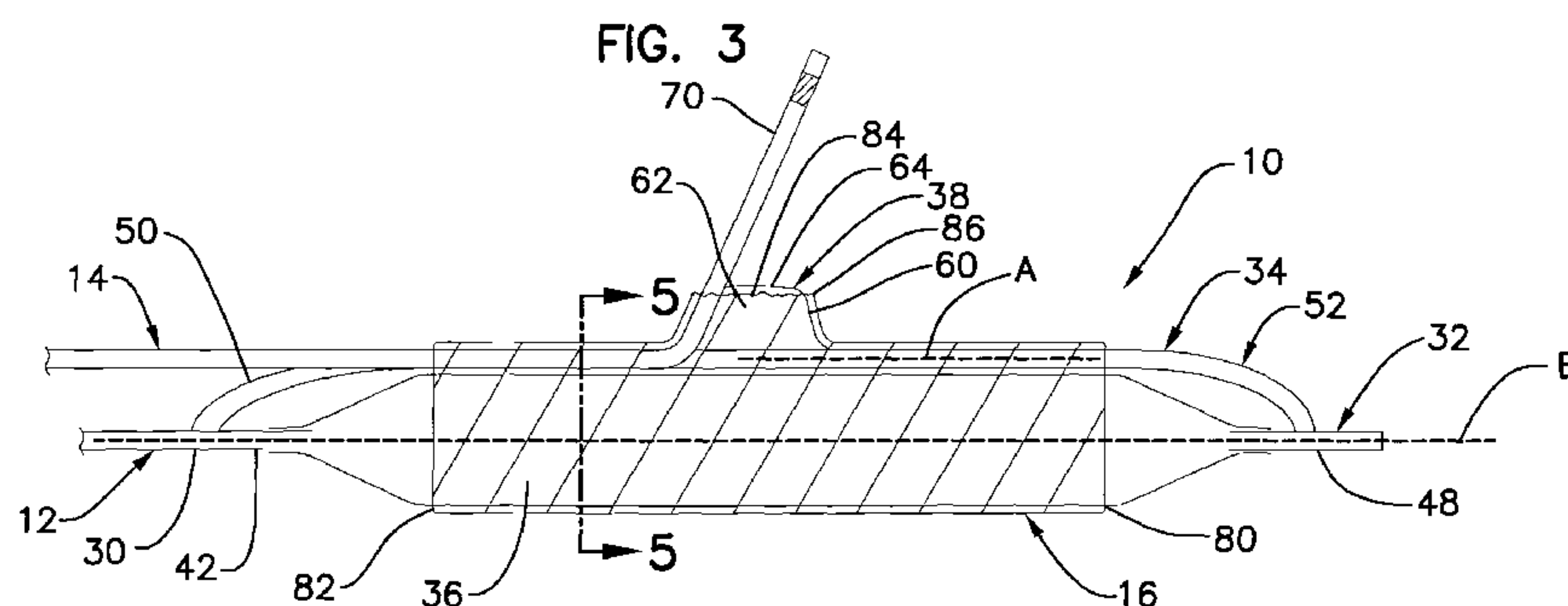
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BIFURCATION CATHETER ASSEMBLY AND METHODS

Technical Field

This disclosure relates to catheter systems and methods for treating vessel
5 bifurcations. Preferred arrangements relate to inflation lumen and balloon
configurations for bifurcation catheter systems.

Background

Catheters are used with stents and inflatable structures to treat conditions such
10 as strictures, stenoses, and narrowing in various parts of the body. Various catheter
designs have been developed for the dilatation of stenoses and to deliver and deploy
stents at treatment sites within the body.

Stents are typically intraluminally placed by a catheter within a vein, artery, or
other tubular shaped body organ for treating conditions such as, for example,
15 occlusions, stenoses, aneurysms, dissections, or weakened, diseased, or abnormally
dilated vessels or vessel walls, by expanding the vessels or by reinforcing the vessel
walls. Once delivered, the stents can be expanded using one or more inflatable
members such as balloons. Stents can improve angioplasty results by preventing
elastic recoil and remodeling of the vessel wall and treating dissections in blood
20 vessel walls caused by balloon angioplasty of coronary arteries. Stents can also be
used as a drug delivery medium for treatment of damaged portions of a vessel.

While conventional stent technology is relatively well developed, stent
technologies related to treatment of the region of a vessel bifurcation are still being
developed.

25

Summary of the Disclosure

The present disclosure relates generally to catheter assemblies for treatment of
bifurcated lumens in a patient, such as vessel bifurcations. In one example, a catheter
assembly includes a main catheter branch and a side catheter branch. The main
30 catheter branch includes a main balloon, a side balloon and a side inflation member.
The side inflation member intersects the side balloon at a location on the side balloon
that is offset laterally from a central line passing from a distal-most point on the side
balloon to a proximal-most point on the side balloon. The side balloon is configured
to extend radially outward relative to the main balloon when the side balloon is

inflated. The side catheter branch can be centrally aligned with the side balloon central line and be positioned laterally adjacent to the side inflation lumen.

The side inflation lumen can include a proximal segment that extends proximally from the side balloon, and a distal segment that extends distally from the side balloon. Typically, at least the proximal segment intersects the side balloon laterally offset from the side balloon central line. In some arrangements, the distal segment can also intersect the side balloon at a laterally offset location from the side balloon central line.

The catheter assemblies can be used to deliver a stent to a vessel treatment site. An example stent delivered by the catheter assemblies includes a lateral branch opening through which the side catheter branch extends. The side balloon is aligned with the lateral branch opening, wherein inflation of the side balloon can help further open the lateral branch opening of the stent.

There is no requirement that an arrangement or method include all features characterized herein to obtain some advantage according to this disclosure.

Brief Description of the Drawings

Figure 1 is a schematic representation of an exemplary catheter assembly for treatment of a bifurcation, the assembly constructed having a proximal end portion and distal end portion, wherein the distal end portion includes main and side balloons and main and side catheter branches.

Figure 2 is a schematic side view of the distal end portion of the catheter assembly shown in Figure 1 with the balloons in the un-inflated state.

Figure 3 is a schematic side view of the distal end portion of the catheter assembly shown in Figure 1 with the balloon portions inflated and the stent expanded.

Figure 4 is a schematic cross-sectional view taken along cross-sectional indicators 4-4 in Figure 2.

Figure 5 is a schematic cross-sectional view of the catheter assembly shown in Figure 3.

Figure 6 is a schematic top view of the catheter assembly shown in Figure 3 with the stent removed.

Figure 7 is a schematic cross-sectional view taken along cross-sectional indicators 7-7 in Figure 5.

Figure 8 is a schematic top view of an example side balloon with an offset side

inflation member segment.

Figure 9 is a schematic top view of another example side balloon with two offset side inflation member segments.

Figure 10 is a schematic top view of another side balloon and an alternative
5 arrangement of two side offset inflation member segments.

Figure 11 is a schematic rear view of an example side balloon with a side inflation member segment centrally positioned.

Figure 12 is a schematic rear view of an example side balloon with an offset side inflation member segment.

10 Figure 13 is a schematic front view of an example side balloon with an offset side inflation member segment.

Figure 14 is a schematic top view of the side balloon and side inflation member segments shown in Figure 8 with a centrally oriented side catheter branch.

Figure 15 is a schematic top view of the side balloon and side inflation
15 member arrangement shown in Figure 9 with a centrally oriented side catheter branch.

Figure 16 is a schematic top view of the side balloon and side inflation member arrangement shown in Figure 10 with a centrally located side catheter branch.

Figure 17 is a schematic rear view of the side balloon and side inflation member shown in Figure 28 with an offset side catheter branch.

20 Figure 18 is a schematic rear view of the side balloon and side inflation member arrangement shown in Figure 15 with a centrally located side catheter branch.

Figure 19 is a schematic front view of the side balloon and side inflation member arrangement shown in Figure 15 with a centrally located side catheter branch.

Figure 20 is a schematic cross-sectional side view of an example two-piece
25 mold used to generate a side balloon from a side inflation member.

Figure 21 is a schematic cross-sectional top view of the mold shown in Figure 20.

Figure 22 is a schematic top view of an example side balloon with offset side inflation member segments and a centrally oriented side catheter branch, wherein the
30 side balloon includes an angled surface.

Figure 23 is a schematic side view of the side balloon, side inflation member, and side catheter branch shown in Figure 22.

Figures 24-26 illustrate the catheter assembly shown in Figures 1-7 in use treating a vessel bifurcation.

Figure 27 illustrates the catheter assembly having a proximal side inflation member segment that is centrally located on a rear side of the side balloon.

Figure 28 illustrates the catheter assembly shown in Figure 27 with the main and side balloons inflated.

5 Figure 29 is a schematic cross-sectional view taken along cross-sectional indicators 29-29 in Figure 27.

Figure 30 is a schematic cross-sectional view taken along cross-sectional indicators 30-30 in Figure 28.

10 Figure 31 is a schematic top view of the catheter assembly shown in Figure 27 with the stent removed.

Figure 32 is a schematic top view of the catheter assembly shown in Figure 28 with the stent removed.

Detailed Description

15 This disclosure relates to bifurcation treatment systems, catheter assemblies, and related methods of treating bifurcations in a patient's body. The term bifurcation means a division location from one unit into two or more units. Generally, two types of bifurcations of a body organ include: 1) a main tubular member defining a main lumen and a branch tubular member defining a branch lumen that extends or branches
20 off from the main tubular member, wherein the main and branch lumens are in fluid communication with each other, and 2) a primary or main member defining a primary or main lumen (also referred to as a parent lumen) that splits into first and second branch members defining first and second branch lumens. The term lumen means the cavity or bore of a tubular structure such as a tubular organ (e.g., a blood vessel).

25 An example bifurcation is a vessel bifurcation that includes a continuous main vessel and a branch vessel, wherein the vessels define a main lumen and a branch lumen, respectively that are in fluid communication with each other. Alternatively, a vessel bifurcation can include a parent vessel that divides into first and second branch vessels, wherein the vessels define a parent lumen and first and second branch lumens,
30 respectively, which lumens are all in fluid communication with each other.

Example applications of the inventive principles disclosed herein include cardiac, coronary, renal, peripheral vascular, gastrointestinal, pulmonary, urinary, and neurovascular systems. The catheter assemblies, systems and methods disclosed herein can be used for locating a branch vessel of the vessel bifurcation and for

placement of a stent relative to the vessel bifurcation for treatment of the vessel bifurcation.

While alternatives are possible, the example catheter assemblies disclosed herein generally include at a distal end portion thereof a main catheter branch and a
5 side catheter branch. The side catheter branch typically includes a side guidewire housing that defines a side guidewire lumen. A distal end portion of the side catheter branch is configured to extend into a branch vessel at a vessel bifurcation. The side catheter branch is used to align features of a stent carried by the proximal end portion of the vessel bifurcation treatment system with an ostium (also referred to as a branch
10 vessel opening) into the branch vessel.

The main catheter branch includes a catheter shaft having a distal end portion. A main balloon and a side balloon are positioned at the distal end portion of the catheter shaft. A main catheter branch includes a main guidewire housing that defines a main guidewire lumen. The main guidewire housing extends through the main
15 balloon. The side balloon is positioned on a side inflation member. The side inflation member can extend in parallel with a longitudinal dimension of the main balloon. The side inflation member defines a side inflation lumen. The side inflation member can include proximal and distal segments that are connected in fluid communication with the side balloon. One aspect of the present disclosure relates to the intersection
20 point between the side inflation member and the side balloon. The side inflation member can intersect the side balloon at a location that is laterally offset from a central proximal-distal line of the side balloon. The central proximal-distal line of the side balloon extends from a proximal-most point on the side balloon to a distal-most point on the side balloon. Typically, the central proximal-distal line of the side
25 balloon is also located centrally between opposing laterally offset sides of the side balloon (e.g., sides 61-63 in Figures 8-10). The location of the central proximal-distal line can also be defined as the line that divides the side balloon in half in a projection view of the side balloon, wherein the projection view is that view from a direction in which the side balloon extends radially outward from the main balloon (e.g., see
30 Figures 8-10).

Another aspect of the present disclosure relates to various configurations of the side balloon (e.g., shape and size) and the interface between the side balloon and the side catheter branch when the side balloon is inflated.

Referring now to Figures 27-31, an example catheter assembly 200 is shown

and described. The catheter assembly 200 includes a main catheter branch 12, a side catheter branch 14, and a stent 16. The main catheter branch 12 includes a catheter shaft 30 having a distal end portion 42, a guidewire housing 32, a main balloon 36, a side inflation member 34, and a side balloon 38. The side catheter branch 14 has a distal end portion 70. The stent 16 is operably mounted to the main balloon 36, side inflation member 34, and the side catheter branch 14 using, for example, crimping techniques. The distal end portion 70 of the side catheter branch 14 extends into the stent 16 through a proximal end 82 thereof, and protrudes out of a lateral branch opening 84. The lateral branch opening is positioned on the stent at a location between the proximal and distal open ends 82, 80.

The catheter assembly has a first distal profile P1 at a location distal the lateral branch opening 84, and a second distal profile P2 at a location proximal of the lateral branch opening 84 and distal of the proximal end 82 of the stent. As shown in Figure 29, the profile P2 results from stacking the proximal segment of the side inflation member on top of the main balloon, and stacking the side catheter branch 14 on top of the side inflation member 34. Such stacking of the side inflation member 34 and the side catheter branch 14 helps maintain the side catheter branch 14 centrally aligned with a central proximal-distal line A of the side balloon 38 (see Figure 32). The profiles P1, P2 can generally be the same, wherein a difference between P1 and P2 being that the side catheter branch is positioned outside of the stent for the profile P1 versus inside the stent for the profile P2. In one example, the profiles P1, P2 are each in the range of about 0.055 in. to about 0.062 in., inclusive.

When the main and branch balloons 36, 38 are inflated, the side catheter branch 14 typically moves laterally into a position from on top of the proximal segment of the side inflation member to a position laterally adjacent to the proximal segment of the side inflation member 34 as shown in Figure 30. This lateral shifting of the side catheter branch results in the side catheter branch not being aligned centrally with the central proximal-distal line A of the side balloon 38 as shown in Figure 32. Figure 31 illustrates how the side catheter branch 14 is aligned with the side inflation member 34 and side balloon 36 along the central proximal-distal line A prior to inflation of the main and side balloons 36, 38. Figure 32 illustrates how the side catheter branch 14 is shifted laterally (laterally to the right in this arrangement) relative to the central proximal-distal line A of the side balloon 38.

When the side catheter branch 14 shifts laterally in this manner, the side

catheter branch 14 typically is offset from being centrally aligned with the lateral branch opening 84 of the stent 16 as shown in Figure 30. Typically, the side catheter branch 14 shifts around to a right or left side of the side balloon 38, which results in the side catheter branch 14 extending out of the lateral branch opening 84 along the left or right side of the lateral branch opening 84. A side catheter branch that is not aligned centrally with the lateral branch opening of the stent can affect proper alignment of the lateral branch opening 86 with an opening into a branch vessel of a vessel bifurcation as will be discussed in further detail below with reference to Figures 24-26.

Referring now to Figure 11, an example side balloon 38 is shown from a rear or proximal view. The proximal segment of the side inflation member 34 is positioned at a center location at a portion across the width of the side balloon 38 along the proximal side 62. When the side balloon 38 and side inflation lumen 34 are used with a side catheter branch 14 as described above with reference to Figures 27-32, the side catheter branch 14 tends to shift laterally as shown in Figure 17 so as to be positioned laterally adjacent to the proximal segment 50 of the side inflation member 34 at a location proximal of the side balloon 38. Figure 17 illustrates the side catheter branch 14 positioned to the right of the side inflation member 34 such that the distal end portion 70 of the side catheter branch 14 extends along the right side 63 and across the radially outward facing surface 64 of the side balloon 38 when the side balloon 38 is inflated. In other instances, the side catheter branch 14 shifts laterally to the left so as to be aligned along the left side 61 of the side balloon 38.

Other example catheter assemblies disclosed herein with reference to Figures 1-26 include various configurations for attachment of the side inflation member distal and proximal segments to the side balloon. These arrangements can improve alignment of the side catheter branch 14 centrally with the central proximal-distal line of the side balloon when the side and main balloons are inflated. By maintaining proper orientation of the side catheter branch relative to the side balloon, the catheter assemblies disclosed herein can provide improved consistency in aligning the lateral branch opening of the stent with the ostium of the branch vessel when treating a vessel bifurcation.

The Example Illustrated in Figures 1-7

An example catheter assembly 10 is shown schematically with reference to

Figures 1-7. The catheter assembly 10 is configured for treatment of a vessel bifurcation such as the vessel bifurcation 22 shown in Figures 24-26. While alternatives are possible, the catheter assembly 10 generally includes a main catheter branch 12, a side catheter branch 14, and a stent 16. The main catheter branch 12 includes a catheter shaft 30 having a proximal end portion 40 (see Figure 1) and a distal end portion 42. The catheter shaft 30 defines an inflation lumen 44 (see Figure 7) that extends from the proximal end portion 40 to the distal end portion 42.

The main catheter branch 12 further includes a main guidewire housing 32 having a distal end portion 48. The main guidewire housing 32 defines a main guidewire lumen 46 as shown in Figure 7 that is sized to advance over a main guidewire. The main guidewire housing 32 can extend within the catheter shaft 30 between the proximal and distal end portions 40, 42. Alternatively, the main guidewire housing 32 can be configured as a rapid exchange arrangement wherein the main guidewire housing extends along only a portion of the length of the catheter shaft 30.

The main catheter branch can also include a main balloon 36 extending along the distal end portion 48 of the guidewire housing 32. A proximal end of the main balloon 36 extends from the proximal end portion of the catheter shaft 30, and a distal end of the main balloon 36 is secured to the main guidewire housing 32 at the distal end portion 48 (see Figure 7).

The main catheter branch 12 further includes a side inflation member 34. The side inflation member 34 includes a proximal segment 50 having proximal and distal ends 51, 53, and a distal segment 52 having proximal and distal ends 55, 57 (see Figure 7). The side inflation member defines a side inflation lumen 54 through which inflation fluid is provided to a side balloon 38. The side balloon includes distal and proximal sides 60, 62, left and right sides 61, 63 (also referred to as opposing laterally offset sides), and a radially outward facing surface 64. When un-inflated, the side balloon 38 maintains a generally collapsed profile (e.g., see side balloon 38 in the collapsed state shown in Figures 27 and 31). When inflated, the side balloon extends radially outward relative to the side inflation member 34 and the main balloon 36 (see Figures 3, 5 and 7). The side balloon 38 typically has a width W (see Figure 6), a length L (see Figure 6), and height H (see Figure 7). The side balloon 38 illustrated in Figures 1-26 has a generally cylindrical shape with a circular cross-section. In other examples, the side balloon can have a variety of different shapes, sizes and cross-

sections.

The side balloon can extend from the side inflation member 34 and the main balloon 36 (i.e. a longitudinal axis or a longitudinal dimension of the side balloon and main balloon (e.g. axis B shown in Figure 3)) at an angle different from the generally 90° (perpendicular) angle shown in the Figures. While alternatives are possible, the side balloon 38 can generally extend from the side inflation member 34 and the main balloon 36 at an angle in the range of about 30° to about 90°, inclusive.

Figures 1 and 2 illustrate the catheter assembly 10 with the main and side balloons 36, 38 in an un-inflated state. Figure 4 illustrates the catheter assembly 10 with the main and branch balloons 36, 38 in an inflated state. Figure 4 illustrates the relative arrangement of the side inflation member 34, side catheter branch 14, and main balloon 36 proximal of the lateral branch opening 84 in the stent 16. The side inflation member 34 and side catheter branch 14 are positioned laterally adjacent to each other and resting in contact with an outer surface of the balloon 36.

In the inflated state shown in Figures 3, 5, 6 and 7, the side inflation member 34 is positioned offset laterally to the left relative to the central proximal-distal line A of the side balloon 38. The side inflation member 34 can also be offset from a central plane C (see Figure 5) that extends through the central proximal-distal line A and a longitudinal axis B of the main balloon 36 (see Figure 3). The offset orientation of the side inflation member 34 provides space for the side catheter branch 14 to remain aligned with the central proximal-distal line A of the side balloon 38 as well as being centrally oriented relative to the lateral branch opening 84 of the stent 16.

Figure 6 illustrates a particular arrangement for connection of the proximal and distal segments 50, 52 of the side inflation member 34 to the proximal and distal sides 62, 60 of the side balloon 38, respectively. The distal end 53 of the proximal segment 50 is connected to the proximal side 62 of the side balloon at a position offset towards the left side 61 of the side balloon. The proximal end 55 of the distal segment 52 is connected to the side balloon 38 along the distal side 60 at a orientation offset laterally from the central proximal-distal line A to the left side 61. This laterally offset arrangement of the proximal and distal segments 50, 52 relative to the central proximal-distal line A of the side balloon 38 permits orientation of the side catheter branch 14 along the central proximal-distal line A before and after inflation of the main and side balloons 36, 38. This arrangement typically provides at least two possible advantages.

One possible advantage of intersecting the side inflation member 34 with the side balloon 38 at a location laterally offset from the central proximal-distal line A is that the profiles P3, P4 of the catheter assembly 10 in the un-inflated state shown in Figure 2 can be reduced as compared to an arrangement wherein the side catheter branch 14 is stacked on top of the side inflation member 34. Typically, the profiles P3, P4 (representing the maximum outer dimension of the catheter assembly along the stent 16 at locations distal and proximal of the lateral branch opening 84, respectively), are each less than the profiles P1, P2 of the catheter assembly 200 described above. This reduced profile results from the ability to position the side catheter branch 14 laterally adjacent to the side inflation member 34 in the catheter assembly 10 instead of vertically on top of the side inflation member 34 (see catheter assembly 200 in Figures 27-30).

Another possible advantage of intersecting the side inflation member 34 with the side balloon 38 at a location laterally offset from the central proximal-distal line A relates to maintaining alignment of the side catheter branch 14 along the central proximal-distal line A of the side balloon 38 before, during and after inflation of the main and side balloons 36, 38.

The Example Illustrated in Figures 8-19

Figures 8-10 illustrate several connection arrangements of the distal and proximal segments of the side inflation member 34 to a side balloon 38. Figures 14-16 illustrate the arrangements of Figures 8-10 in combination with a side catheter branch 14 arranged along a central proximal-distal line A of the side balloon 38. Figures 8 and 14 illustrate intersection of the proximal segment 50 at its distal end 53 with the proximal side 62 of the side balloon at a location offset laterally towards the right side 63 of the side balloon. The distal segment 52 intersects at its proximal end 55 with the distal side 60 of the side balloon and is aligned along the central proximal-distal line A of the side balloon 38.

Figures 9 and 15 illustrate the proximal segment 50 intersects at the proximal side 62 offset towards the right side 63 of the side balloon. The distal segment 52 intersects at the distal side 60 of the side balloon offset to the left side 61 of the balloon. Figures 10 and 16 illustrate the proximal and distal segments 50, 52 intersect with the proximal and distal sides 62, 60 of the side balloon and laterally offset towards the right side 63 of the side balloon 38.

The arrangement of Figures 8 and 14 can be modified in other examples to offset the proximal segment 50 to the left side 61 rather than towards the right side 63 as shown. Likewise, in Figures 9 and 15 the proximal segment 50 can be offset towards the left side 61 and the distal segment 52 can be offset towards the right side 63 in other examples. The mirror image of the arrangement in Figures 10 and 16 is illustrated in Figure 6.

Figures 12 and 13 illustrate the distal and proximal sides 60, 62 of the side balloon 38 and the various intersection points of the proximal and distal segments 50, 52 at offset locations to the left or right sides 61, 63 of the side balloon 38. Figures 18 and 19 illustrate the arrangements of Figures 12 and 13 and further illustrate positioning of the side catheter branch 14 centrally relative to the side balloon 38 between the left and right sides 61, 63. In one example, the maximum width dimension of the proximal and distal segments 50, 52 of the side inflation member 34 is substantially the same as the maximum width dimension of the side catheter branch 14 in the area of the side balloon 38. Therefore, offsetting the segments 50, 52 to either the left or right side relative to a position of the segments 50, 52 aligned with the central proximal-distal line A a distance equal to half of that maximum width dimension will permit orientation of the side catheter branch 14 along the central proximal-distal line A.

The maximum width W of the balloon 38 (see Figures 12 and 18) in one example can be about three times the maximum width dimension of the side catheter branch 14 or the side inflation member 34. With these dimensions for the side balloon 38, the side catheter branch 14 can remain centrally aligned with the central proximal-distal line A and the segments of the side inflation member 34 can be positioned on either the left or the right side of the side catheter branch 14 without the side inflation member 38 protruding laterally beyond the width of the side balloon 38. In some preferred arrangements, the side inflation member 34 does not extend laterally beyond the left or right side 61, 63 beyond the width W of the side balloon 38. In other examples, the side balloon can have any width W or other dimension that is unrelated to the maximum width dimensions of either the side catheter branch 14 or the side inflation member 34.

Referring again to Figures 9 and 15, the intersection of the proximal and distal segments 50, 52 at laterally offset orientations to the right or left (or vice versa) of the central proximal-distal line A of side balloon 38 can provide additional advantages

when inflating the main and branch balloons 36, 38. The catheter assembly 10 shown in Figures 1-7 includes a side inflation member 34 that is connected at its proximal-most and distal-most ends to the catheter shaft 30 and main guidewire housing 32, respectively (i.e., at the distal end of the distal segment 52 and the proximal end of the proximal segment 50). Typically, the length of the side inflation member 34 between these proximal and distal connection points is such that prior to inflation of the main and side balloons 36, 34, there is relatively little slack or extra length in the side inflation member 34. While additional advantages are possible, minimizing the length of the side inflation member between the proximal and distal connection points can help, for example, minimize the profiles P3, P4, improve alignment of the lateral branch opening with the side balloon during assembly of the catheter assembly 10 (e.g., crimping the stent 16 in place on the main balloon 36 with the side balloon 38 aligned with the lateral branch opening 34), and reduce the incidence of loose material and structure at the distal end portion of the catheter assembly to improve ease of advancing the catheter assembly 10 to a bifurcation treatment site.

Providing the side inflation member 34 with a minimum length between the proximal-most and distal-most connection points can result in additional tension forces in the side inflation member generated when inflating the main balloon 36. Inflation of the main balloon 36 increases the length of the path that the side inflation member 34 must traverse between the proximal-most and distal-most connection points. These additional tension forces in the side inflation member can be problematic. For example, the additional tension forces can weaken the connection between the side inflation member segments 50, 52 and the side balloon 38, weaken the connection point at the proximal-most and distal-most connection points of the side inflation member 34 to the main guidewire housing 32 at a point distal of the main balloon 36 and the catheter shaft 30 at a point proximal of the main balloon 36, create deformations in the main balloon 38 outer profile, or create some misalignment between the side balloon 38 and the lateral branch opening 84 of the stent 16.

Referring again to Figures 9 and 15, laterally offsetting the proximal and distal segments 50, 52 in opposing directions relative to the central proximal-distal line A of the side balloon 38 can result in a rotational force that rotate the side balloon 38 when the main balloon 36 is inflated and tension is generated along the length of the side inflation member 34. This rotational force can slightly lengthen the side inflation member or at least provide some flexibility in the side inflation member 34 that can

minimize the generation of the potential tension induced concerns discussed above. The rotation effect can be further magnified if the intersection point between the distal end 57 of the proximal segment is along the side 63 further around towards the distal side 60 of the side balloon. Likewise, intersecting the proximal end 51 of the distal segment 52 with the balloon at a location further along the side 61 of the balloon around towards the proximal side 62 of the balloon can create further rotation of the side balloon 38 and flexibility during inflation of the main and side balloons 36, 38 that can be advantageous.

10 The Example Illustrated in Figures 20 and 21

Figures 20 and 21 illustrate an example mold used in creating a side balloon 38 from the side inflation member 34. The mold 90 includes a proximal portion 92 and a distal portion 94 that together define a balloon cavity 96, a proximal segment cavity 98, and a distal segment cavity 100. The mold 90 can be configured such that the balloon cavity defines any desired balloon shape, size and orientation relative to the inflation member 34. The arrangement of the proximal and distal segment cavities 98, 100 relative to the balloon cavity 96 results in a desired laterally offset intersection of one or more of the proximal and distal segments 50, 52 of the side inflation member with the side balloon 38.

20 In operation, a side inflation member 38 is positioned within the cavities 96, 98, 100. Heat is applied to the mold 90 and pressure is applied internal the side inflation member 34 that results in expansion of portions of the side inflation member 34 to fill the balloon cavity 96. Typically, the mold is heated using an external source of heat such as hot water or hot air. Pressure is applied internally in the side inflation member 34 by filling the side inflation member 34 with a fluid such as heated water. The resulting wall thickness of the side balloon 38 can be modified by, in addition to changing the size of the balloon cavity 96, increasing or reducing the thickness of portions of the side inflation member 34 prior to the molding process. The thickness of the side inflation member 34 can also be reduced after the molding process to reduce the wall thickness of the balloon 38 (e.g., using grinding or other material removal techniques).

The two-part design of the mold 90 is advantageous for removing the molded product from the mold 90 since the side balloon 38 is now molded integral with the side inflation member 34 and would be difficult to remove otherwise. The mold 90

shape and two-piece configuration is exemplary only. Other configurations, such as molds having three or more pieces are possible in other examples.

The Example Illustrated in Figures 22 and 23

5 Figures 22 and 23 illustrate a side balloon design that can help maintain the side catheter branch 14 in a position oriented along the central proximal-distal line A of the side balloon 38 during treatment of a vessel bifurcation. The side balloon 138 includes an angled surface 66 that extends from the proximal side 62 to the radially outward facing surface 64. The angled surface 66 is generally planar in this
10 arrangement but can have other shapes and sizes in other arrangements. For example, the angled surface 66 can include a contoured shape portion such as a concave shaped portion. The angled surface 66 extends generally at an angle θ relative to a plane extending parallel with the central proximal-distal line A. Although alternatives are possible, the angle θ is typically an acute angle (less than 90°) facing in a distal
15 direction. The angle θ is typically in the range of about 15° to about 90° , and preferably in the range of about 30° to about 75° .

 The orientation of angled surface 66 relative to the side balloon 38 can be altered depending on the orientation of the side balloon 38 relative to the side inflation member 34 and the main balloon 36. For example, if the side balloon 38 extends
20 radially away from the side inflation member 34 and main balloon 36 at an angle less than 90° (e.g., in the range of about 30° to about 80°), the angle θ may be smaller while still providing the desired function.

 The angled surface 66 can have advantages related to keeping the side catheter branch 14 from moving laterally as the side balloon and main balloon 38, 36 inflate.
25 Typically, the side balloon 38 can include a generally cylindrical, spherical, hemispherical shape with many different cross-sectional shapes possible. The side balloon 38 can also typically includes a rounded or convex shaped outward facing surface 64. Since many surfaces of such side balloon constructions are contoured in a convex direction, contact between the side catheter branch 14 and the side balloon 38
30 is usually a point (assuming no concave deformation of the side catheter branch 14 and side balloon 38). This type of minimum contact on a convex surface can make it more difficult to maintain a single position of the side catheter branch aligned along the central proximal-distal line A of the side balloon 38. Including the generally planar angled surface 66 along the proximal side 62 provides a more substantial

contact surface between the side catheter branch 14 and the side balloon 38. Further, lack of curvature across the width of the surface 66 can make it easier to maintain alignment of the side catheter branch 14 along the central proximal-distal line A.

The addition of other structure such as convex portions, protrusions, or
5 multiple planar surfaces in addition to or in place of the angled surface 66 can provide the same or similar effect of helping maintain alignment of the side catheter branch 14 with the central proximal-distal line A.

The Example Illustrated in Figures 24-26

10 The catheter assembly 10 described above with reference to Figures 1-7 can be used for treatment of a vessel bifurcation 22 as shown in Figures 24-26. Typically, a main vessel guidewire 18 is first inserted into a main vessel 24 of the vessel bifurcation 22 to a point distal of the vessel bifurcation. A branch vessel guidewire 20 is advanced to the vessel bifurcation and inserted through an ostium or opening 28 of
15 a branch vessel 26. Proximal ends of the guidewires 18, 22 are then inserted into the main guidewire lumen 46 and branch guidewire lumen 72, respectively. The catheter assembly 10 is advanced over the guidewires 18, 22 to the vessel bifurcation as shown in Figure 24. The catheter assembly 10 is advanced further distally until the distal end portion 70 of the side catheter branch 14 is positioned within the branch vessel 26. A
20 marker system can be used to help confirm proper rotational (radial) and axial alignment of the lateral branch opening 84 of the stent 16 relative to the ostium 28 into the branch vessel 26. An example marker system is described below.

After proper axial and rotational positioning of the catheter assembly 10 is confirmed, the main and branch balloons 36, 38 are inflated. The laterally offset
25 orientation of at least the proximal segment 50 of the side inflation member 34 provides for positioning of the side catheter branch 14 aligned with the central proximal-distal line A of the side balloon 38 prior to and during inflation of the balloons 36, 38. Central alignment of the side catheter branch 14 with the side balloon 38 can improve consistency in aligning the lateral branch opening 84 with the
30 ostium 28 of the vessel bifurcation. Typically, inflation of the side balloon 38 can also result in expansion of expandable structure 86 surrounding the lateral branch opening 84. The expanded expandable structure 86 can extend through the ostium 28 and at least partially into the branch vessel 26.

In a follow-up step, after the balloons 36, 38 have been deflated and the

catheter branches 12, 14 retracted proximally, a separate balloon member (not shown) can be advanced through the lateral branch opening to treat the branch vessel 26 and further open the expandable structure 86 into the branch vessel 26. In a still further step, an additional branch stent can be advanced through the lateral branch opening 84
5 and into the branch vessel and expanded to treat further the branch vessel 26.

The particular method steps described above can be altered in other example treatment methods. For example, one of the guidewires 18, 20 can be advanced with the catheter assembly 10 to the vessel bifurcation. In another example, the balloons 36, 38 can be inflated sequentially rather than simultaneously for purposes of, for
10 example, improving alignment of the lateral branch opening 84 with the ostium into the branch vessel.

Materials and Other Considerations

A wide variety of stents, catheters, and guidewire configurations can be used
15 with the catheter assembly embodiments of the present disclosure. The inventive principles disclosed herein should not be limited to any particular design or configuration. Some example stents that can be used with the catheter assemblies disclosed herein can be found in, for example, U.S. Pat. Nos. 6,210,429, 6,325,826 and 6,706,062, 7,220,275, and U.S. Published Patent Application No. 2004/0176837 titled
20 SELF-EXPANDING STENT AND CATHETER ASSEMBLY AND METHOD FOR TREATING BIFURCATIONS, the entire contents of which are incorporated herein by reference. In general, the aforementioned stents include a lateral branch opening located between distal and proximal open ends of the stent. The lateral branch opening defines a path between an inner lumen or inner volume of the stent and an
25 area outside of the stent. The stent lateral branch opening is distinct from the cell openings defined between strut structures from which the stent sidewall is constructed. In some stents, the lateral branch opening can be surrounded by expandable structure. The expandable structure can be configured to extend radially into the branch lumen of the bifurcation upon expansion of, for example, an inflatable
30 portion of the bifurcation treatment system. Typically, the stent is expanded after being positioned in the main lumen with the lateral branch opening aligned with an opening into the branch lumen. Alignment of the lateral branch opening with the opening into the branch lumen includes both radial and axial alignment. The stent, including the expandable structure surrounding the lateral branch opening, can be

expanded with a single expansion or multiple expansions using one or more inflatable members.

The main and side balloons, and all other balloons disclosed herein, can be made of any suitable balloon material including compliant and non-compliant materials and combinations thereof. Some example materials for the balloons and catheters disclosed herein include thermoplastic polymers, polyethylene (high density, low density, intermediate density, linear low density), various copolymers and blends of polyethylene, ionomers, polyesters, polycarbonates, polyamides, poly-vinyl chloride, acrylonitrile-butadiene-styrene copolymers, poly ether-polyester copolymers, and polyetherpolyamide copolymers. One suitable material is Surlyn[®], a copolymer polyolefm material (DuPont de Nemours, Wilmington, Del.). Stil further suitable materials include thermoplastic polymers and thermoset polymeric materials, poly(ethylene terephthalate) (commonly referred to as PET), thermoplastic polyamide, polyphenylene sulfides, polypropylene. Some other example materials include polyurethanes and block copolymers, such as polyamide-polyether block copolymers or amide-tetramethylene glycol copolymers. Additional examples include the PEBAX[®] (a polyamide/polyether/polyester block copolymer) family of polymers, e.g., PEBAX[®] 70D, 72D, 2533, 5533, 6333, 7033, or 7233 (available from Elf AtoChem, Philadelphia, Pa.). Other examples include nylons, such as aliphatic nylons, for example, Vestamid L2101 IF, Nylon 11 (Elf Atochem), Nylon 6 (Allied Signal), Nylon 6/10 (BASF), Nylon 6/12 (Ashley Polymers), or Nylon 12. Additional examples of nylons include aromatic nylons, such as Grivory (EMS) and Nylon MXD-6. Other nylons and/or combinations of nylons can also be used. Still further examples include polybutylene terephthalate (PBT), such as CELANEX[®] (available from Ticona, Summit, N.J.), polyester/ether block copolymers such as ARNITEL[®] (available from DSM, Erionspilla, Ind.), e.g., ARNITEL[®] EM740, aromatic amides such as Trogamid (PA6-3-T, Degussa), and thermoplastic elastomers such as HYTREL[®] (Dupont de Nemours, Wilmington, Del.). In some embodiments, the PEBAX[®], HYTREL[®], and ARNITEL[®] materials have a Shore D hardness of about 45D to about 82D. The balloon materials can be used pure or as blends. For example, a blend may include a PBT and one or more PBT thermoplastic elastomers, such as RITEFLEX[®] (available from Ticona), ARNITEL[®], or HYTREL[®], or polyethylene terephthalate (PET) and a thermoplastic elastomer, such as a PBT thermoplastic elastomer. Additional examples of balloon material can be found in U.S. Pat. No.

6,146,356, which is incorporated herein by reference.

The catheter assembly 10 can include marker material that is visible under X-ray or in fluoroscopy procedures. Figure 2 illustrates markers 1-4 positioned along the distal end portions of the main and side catheter branches 12, 14. Any features of the system 10 that include marker material can be more easily identified and distinguished under X-ray or in fluoroscopy procedures. Some example marker materials include gold, platinum and tungsten. In one embodiment, the marker material can be included in a band structure that is secured to at least one of the main and side catheter branches 12, 14. In other embodiments, the marker material is part of the material composition of portions of the main and side catheter branches 12, 14. Viewability of features of the catheter assembly 10 under X-ray or fluoroscopy can assist the physician operating the system 10 to more easily adjust a position of the system 10 relative to the vessel bifurcation 40. Example markers and marker materials suitable for use with system 10 are described in U.S. Pat. No. 6,692,483 to Vardi, et al., and co-pending U.S. Published Patent Application No. 2007/0203562, filed on February 22, 2007, and titled MARKER ARRANGEMENT FOR BIFURCATION CATHETER, which matters are incorporated herein by reference.

Alternative catheter assemblies to those described above are configured for use with stents having self-expanding features. Self-expanding stents and self-expanding features of a stent typically do not require the use of an inflatable member such as a balloon to expand the stent or stent feature. Typically, self-expanding stents, such as those stents described in U.S. Published Patent Application No. 2004/0176837, are held in a constricted state using a sheath that fits over the stent. In the constricted state, the stent is able to navigate through a body lumen to the treatment site. Once the stent and sheath are positioned at the treatment site, the sheath is retracted proximally to release the stent for expansion of the stent into a radially expanded state.

One aspect of the present disclosure relates to a catheter assembly that includes a stent, a main catheter branch, and a side catheter branch. The stent includes a proximal open end, a distal open end, and a lateral branch opening. The main catheter branch includes a proximal end portion and a distal end portion. The distal end portion includes a main balloon, a side balloon, and a side inflation lumen. The main balloon includes a proximal end portion, a distal end portion, and a longitudinal dimension extending between the proximal and distal end portions. The

side balloon is arranged at a location between the proximal and distal end portions of the main balloon and in alignment with the lateral branch opening of the stent. The side balloon is configured to extend radially outward relative to the main balloon when the side balloon is inflated, and defines a central proximal-distal line. The side inflation member intersects the side balloon at a location offset laterally from the side balloon central proximal-distal line. The side catheter branch defines a branch guidewire lumen and extends through the lateral branch opening of the stent. The side catheter branch is centrally aligned with the side balloon central proximal-distal line.

Another aspect of the present disclosure relates to a catheter assembly that includes a main balloon, a side balloon, and a side inflation member. The main balloon includes a proximal end portion, a distal end portion, and a longitudinal dimension extending from the proximal end portion to the distal end portion. The side balloon is positioned at a location between the proximal and distal end portions of the main balloon and is configured to extend radially outward relative to the main balloon when the side balloon is inflated. The side balloon has a proximal-most point, a distal-most point, and a central proximal-distal line that extends from the proximal-most point to the distal-most point of the side balloon. The side inflation member intersects the side balloon at a location offset laterally from the side balloon central proximal-distal axis.

A still further aspect of the present disclosure relates to a method of assembling a stent delivery system. The stent delivery system includes a stent, a main catheter branch, and a side catheter branch. The stent has a proximal open end, a distal open end, and a lateral branch opening. The main catheter branch includes a main balloon, a side balloon and a side inflation member, wherein the side balloon defines a central proximal-distal line, and the side inflation member intersects the side balloon at a location offset laterally from the side balloon central proximal-distal line. The side catheter branch defines a branch guidewire lumen. Steps of the method include extending the main catheter branch into the stent with the main balloon and the side balloon positioned in the stent and the side balloon aligned with the lateral branch opening of the stent, extending the side catheter branch into the stent with a portion of the side catheter branch extending through the lateral branch opening of the stent and the side catheter branch centrally aligned with the side balloon central proximal-distal line, and crimping the stent into releasable engagement with the main

catheter branch and the side catheter branch.

It is noted that not all of the features characterized herein need to be incorporated within a given arrangement, for the arrangement to include improvements according to the present disclosure.

WE CLAIM:

1. A catheter assembly, comprising:
 - a stent having a proximal open end, a distal open end, and a lateral branch opening;
 - a main catheter branch having a proximal end portion and a distal end portion, the distal end portion including:
 - a main balloon, the main balloon having a proximal end portion, a distal end portion, and a longitudinal dimension extending between the proximal and distal end portions;
 - a side balloon arranged at a location between the proximal and distal end portions of the main balloon and in alignment with the lateral branch opening of the stent, the side balloon configured to extend radially outward relative to the main balloon when the side balloon is inflated, the side balloon defining a central proximal-distal line;
 - a side inflation member that intersects the side balloon at a location offset laterally from the side balloon central proximal-distal line; and
 - a side catheter branch defining a branch guidewire lumen, the side catheter branch extending through the lateral branch opening of the stent, the side catheter branch being centrally aligned with the side balloon central proximal-distal line.
2. The assembly of claim 1, wherein the side inflation member includes a proximal segment and a distal segment, the proximal segment extending from the side balloon in a proximal direction, and the distal segment extending from the side balloon in a distal direction.
3. The assembly of claims 1 or 2, wherein the main catheter branch further includes:
 - a catheter shaft having a distal end portion and defining a main inflation lumen; and
 - a main guidewire housing defining a main guidewire lumen, the main guidewire housing extending through at least a portion of the main inflation lumen and through the main balloon from the proximal end portion to the distal end portion of the main balloon.

4. The assembly of any of claims 1-3, wherein the side balloon includes a distal side and a proximal side, and the side inflation lumen intersects the side balloon on the proximal side of the side balloon at a location offset laterally from the side balloon central proximal-distal line.

5. The assembly of any of claims 1-3, wherein the side balloon has a generally cylindrical shape with a radially outward facing surface, a distal side and a proximal side, wherein the side inflation member intersects the side balloon at least at the proximal side of the side balloon.

6. The assembly of any of claims 1-5, wherein the side balloon central proximal-distal axis divides the side balloon in half in a projection view of the side balloon.

7. A catheter assembly, comprising:
a main balloon, the main balloon having a proximal end portion, a distal end portion, and a longitudinal dimension extending from the proximal end portion to the distal end portion;
a side balloon positioned at a location between the proximal and distal end portions of the main balloon, the side balloon configured to extend radially outward relative to the main balloon when the side balloon is inflated, the side balloon having a proximal most point, a distal most point, and a central proximal-distal line that extends from the proximal-most point to the distal-most point of the side balloon; and
a side inflation member that intersects the side balloon at a location offset laterally from the side balloon central proximal-distal axis.

8. The assembly of claim 7, further comprising a side catheter branch centrally aligned with the side balloon central proximal-distal line, the side catheter branch defining a branch guidewire lumen sized to advance over a branch guidewire.

9. The assembly of claims 7 or 8, wherein the side balloon central proximal-distal axis divides the side balloon in half in a projection view of the side balloon, the projection view arranged in a direction in which the side balloon extends radially outward from the main balloon.

10. The assembly of any of claims 7-9, further comprising a stent, the stent having opposing proximal and distal open ends, and a lateral branch opening, the lateral branch opening positioned at a location between the proximal and distal open ends, wherein the stent is operably mounted to the main balloon with the side balloon oriented in alignment with the lateral branch opening.

11. The assembly of any of claims 7-10, further comprising a main guidewire member that extends through the main balloon, the main guidewire member defining a main guidewire lumen sized to advance over a main guidewire.

12. The assembly of any of claims 7-11, wherein the side inflation member includes a distal segment and a proximal segment, the distal segment extending distally from the side balloon and the proximal segment extending proximally from the side balloon, and at least the proximal segment intersects the side balloon at a location offset laterally from the side balloon central proximal-distal axis.

13. The assembly of claim 12, wherein the proximal segment is in fluid communication with the side balloon and the main balloon.

14. The assembly of claims 12 or 13, wherein the distal segment and the proximal segments intersect the side balloon at a location offset laterally from the side balloon central proximal-distal axis.

15. The assembly of any of claims 7-14, wherein the side balloon central proximal-distal axis is arranged in parallel with the longitudinal dimension of the main balloon.

16. The assembly of any of claims 7-15, wherein the side balloon includes an angled surface defined in part in a proximal side of the side balloon and a radially outward facing surface of the side balloon, the angled surface being arranged at an angle less than 90° relative to the longitudinal dimension of the main balloon.

17. A method of assembling a stent delivery system, the stent delivery system including a stent, a main catheter branch, and a side catheter branch, the stent having a proximal open end, a distal open end, and a lateral branch opening, the main catheter branch including a main balloon, a side balloon and a side inflation member, the side balloon defining a central proximal-distal line, and the side inflation member intersects the side balloon at a location offset laterally from the side balloon central proximal-distal line, the side catheter branch defining a branch guidewire lumen, the method comprising:

extending the main catheter branch into the stent with the main balloon and the side balloon positioned in the stent and the side balloon aligned with the lateral branch opening of the stent;

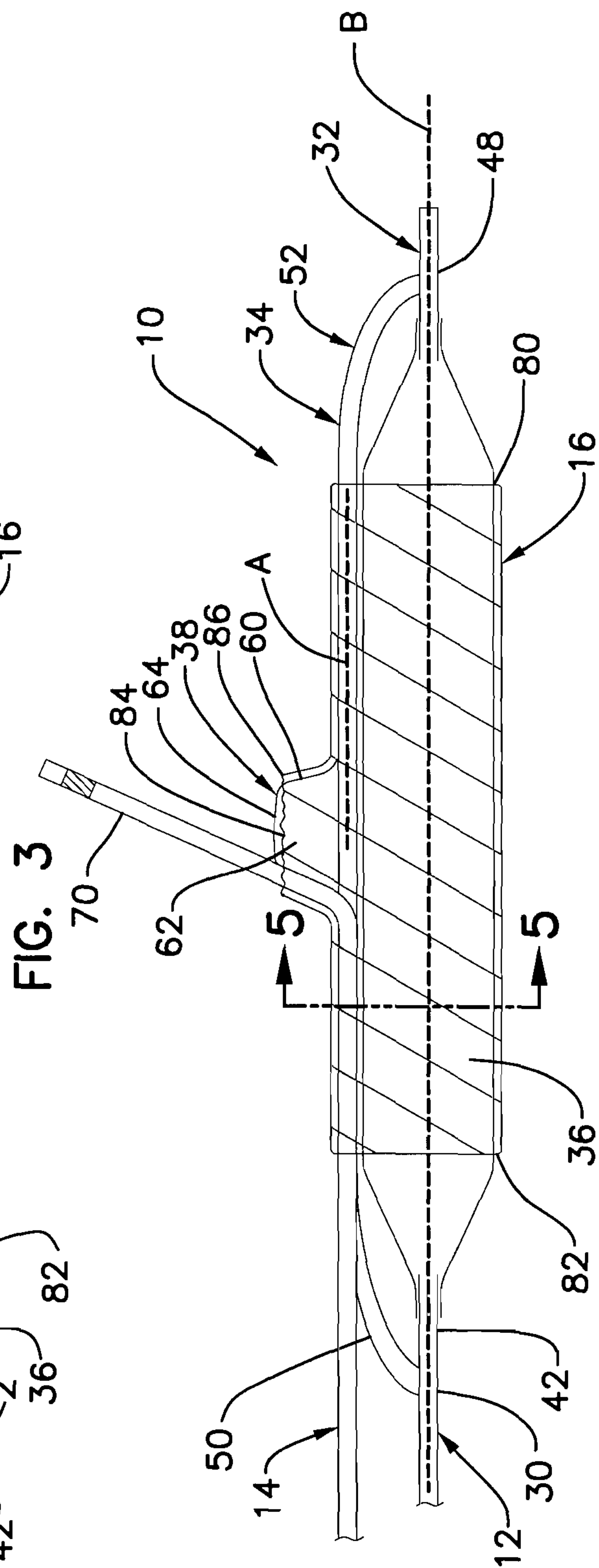
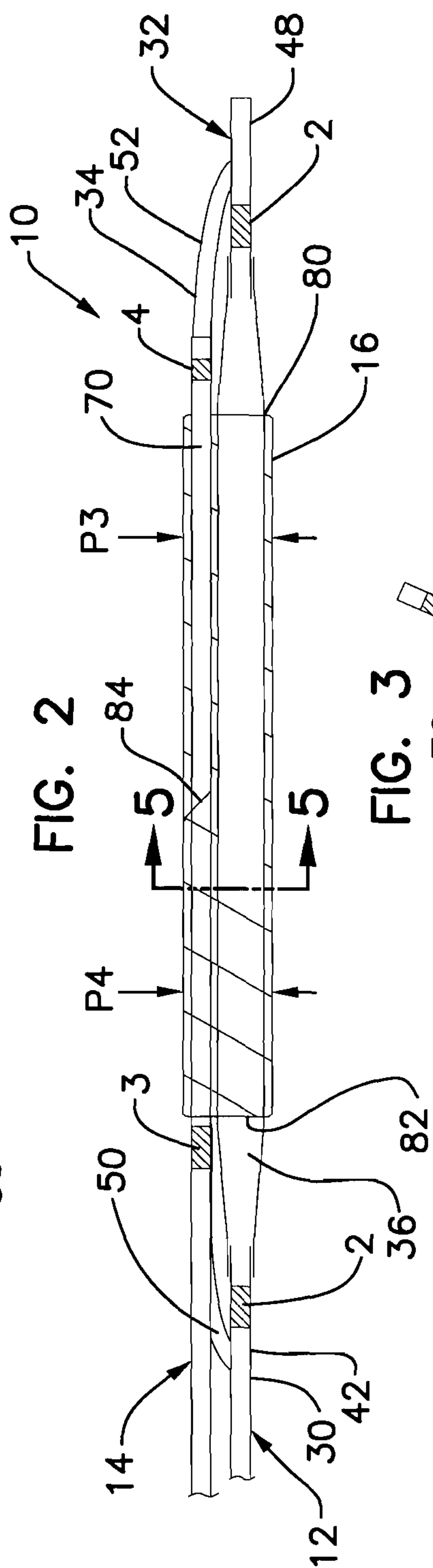
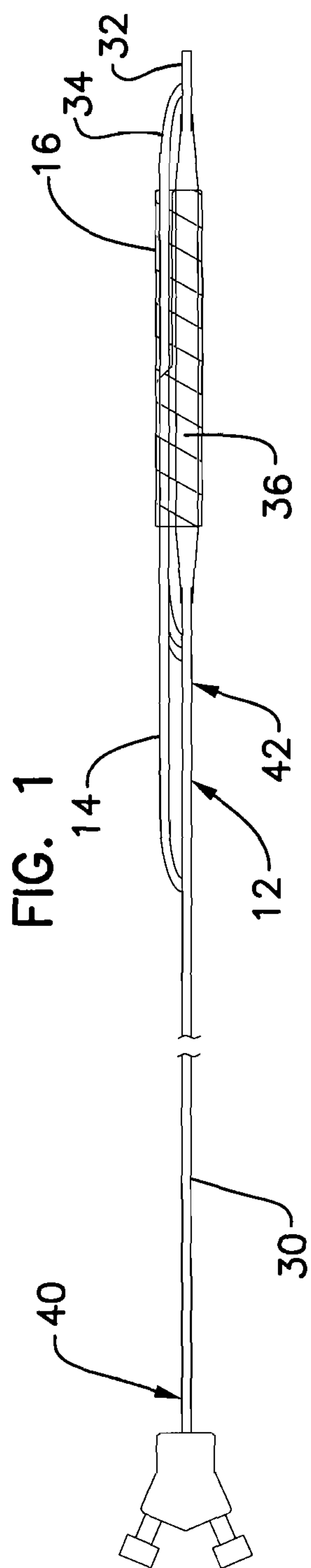
extending the side catheter branch into the stent with a portion of the side catheter branch extending through the lateral branch opening of the stent and the side catheter branch centrally aligned with the side balloon central proximal-distal line;

and

crimping the stent into releasable engagement with the main catheter branch and the side catheter branch.

18. The method of claim 17, wherein the side inflation lumen includes a proximal segment that extends proximally from the side balloon and a distal segment that extends distally from the side balloon, and extending the side catheter branch into the stent includes positioning the side catheter branch laterally adjacent to the proximal segment of the side inflation member.

19. The method of claims 17 or 18, wherein crimping the stent includes reducing an outer profile of the stent while maintaining the side balloon in alignment with the lateral branch opening and while maintaining the side catheter branch centrally aligned with the side balloon central proximal-distal line.



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FIG. 5

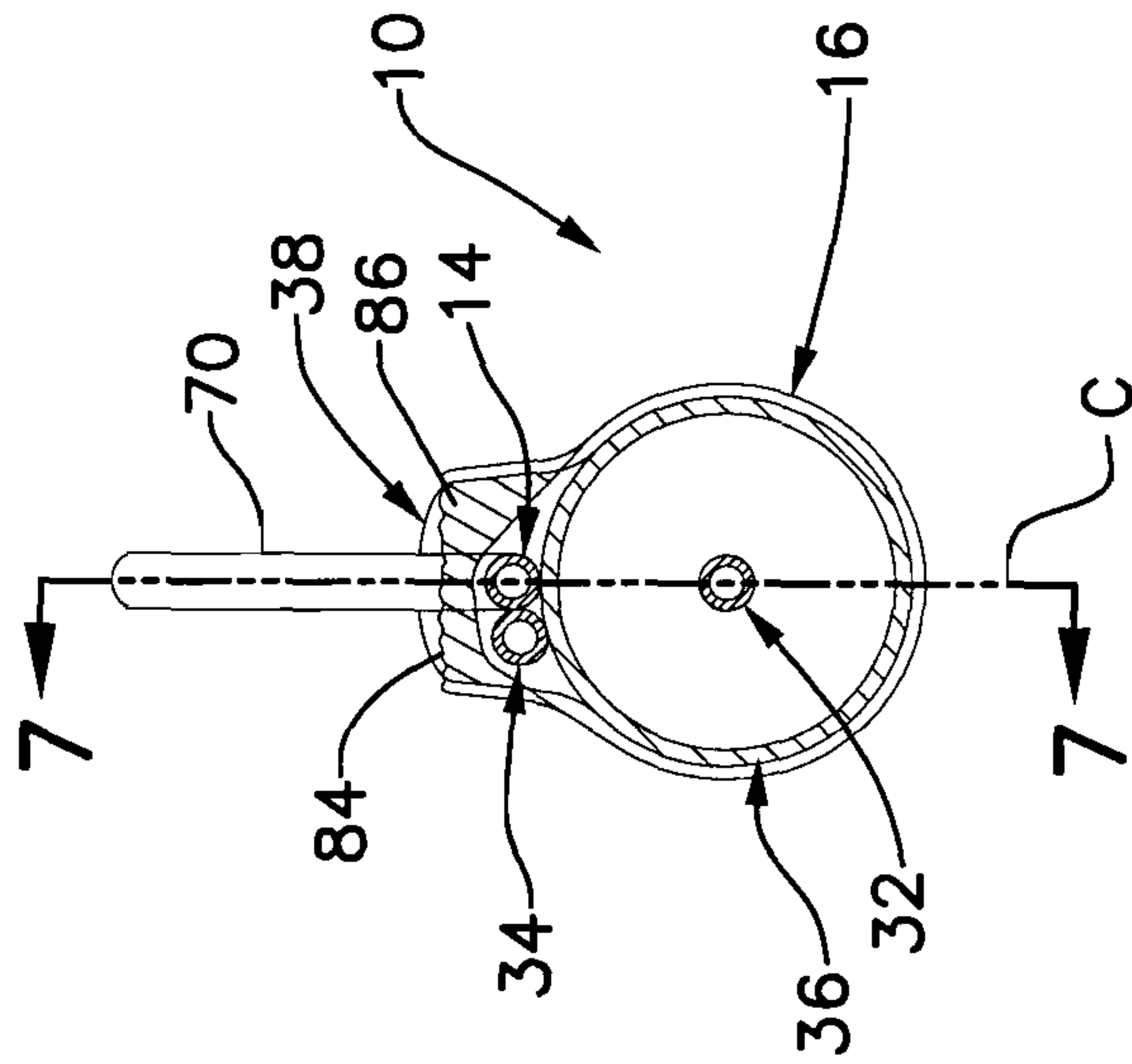


FIG. 4

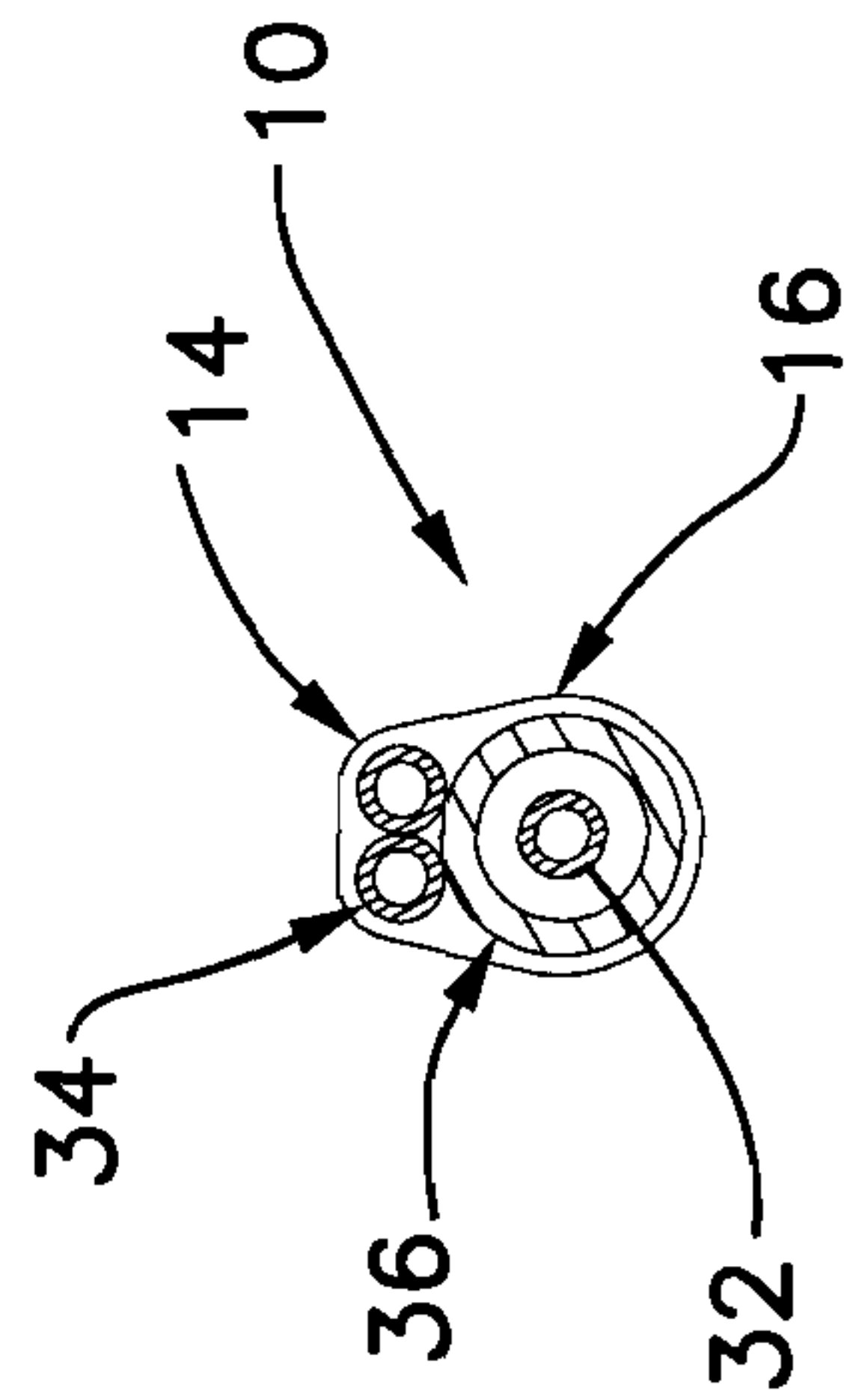
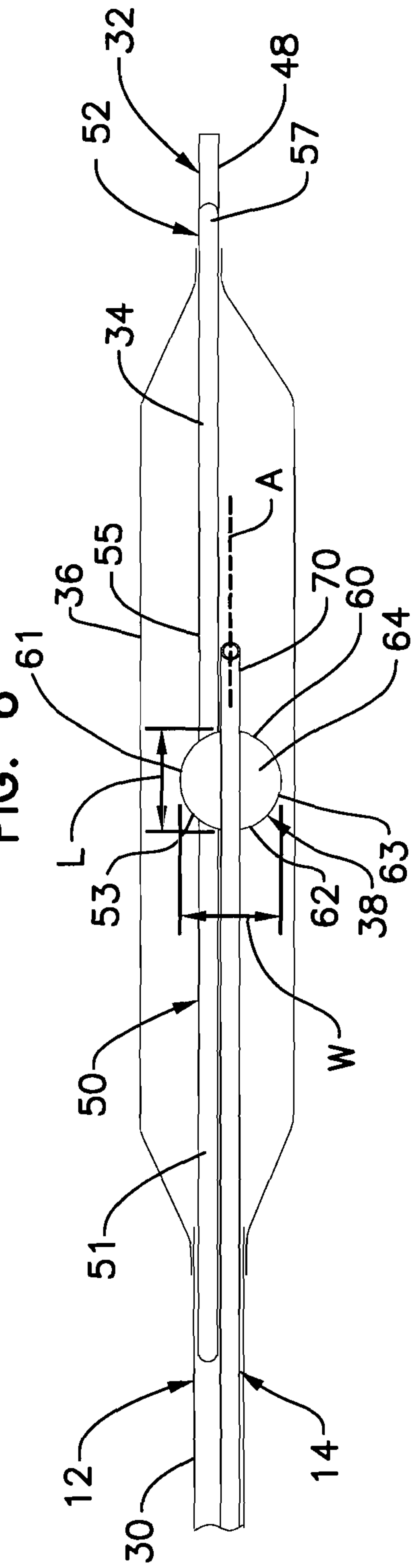


FIG. 6



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FIG. 7

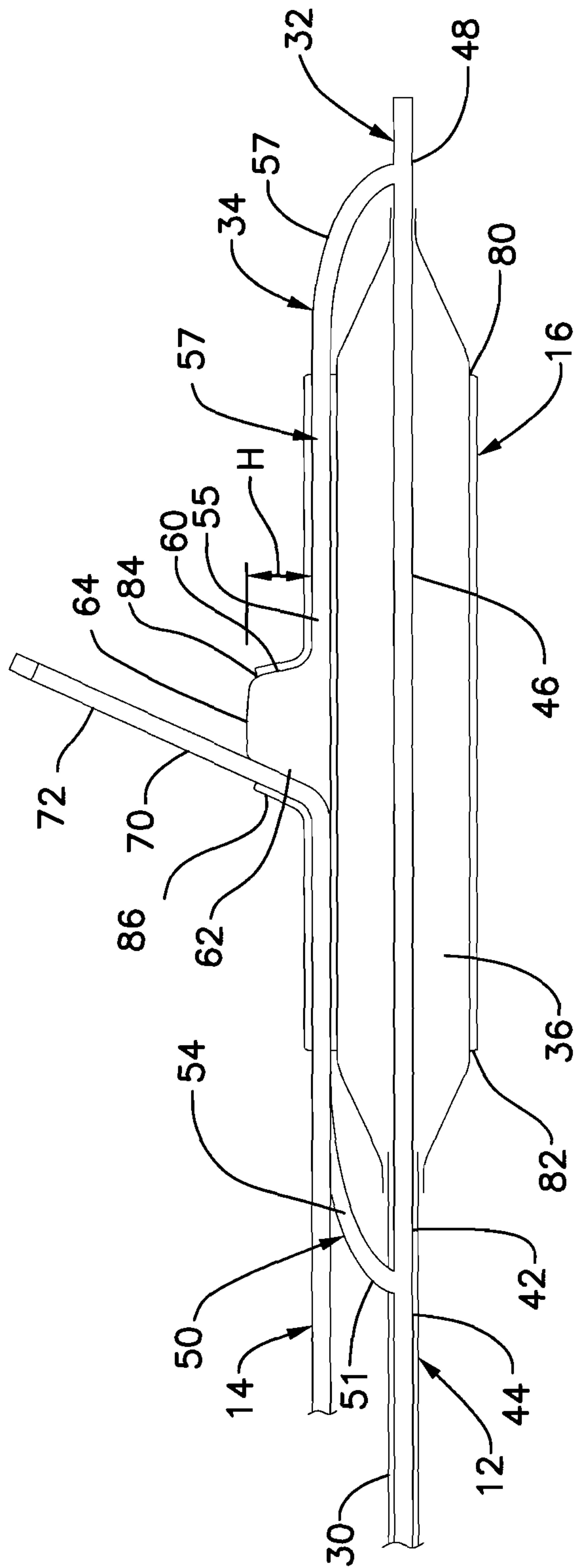
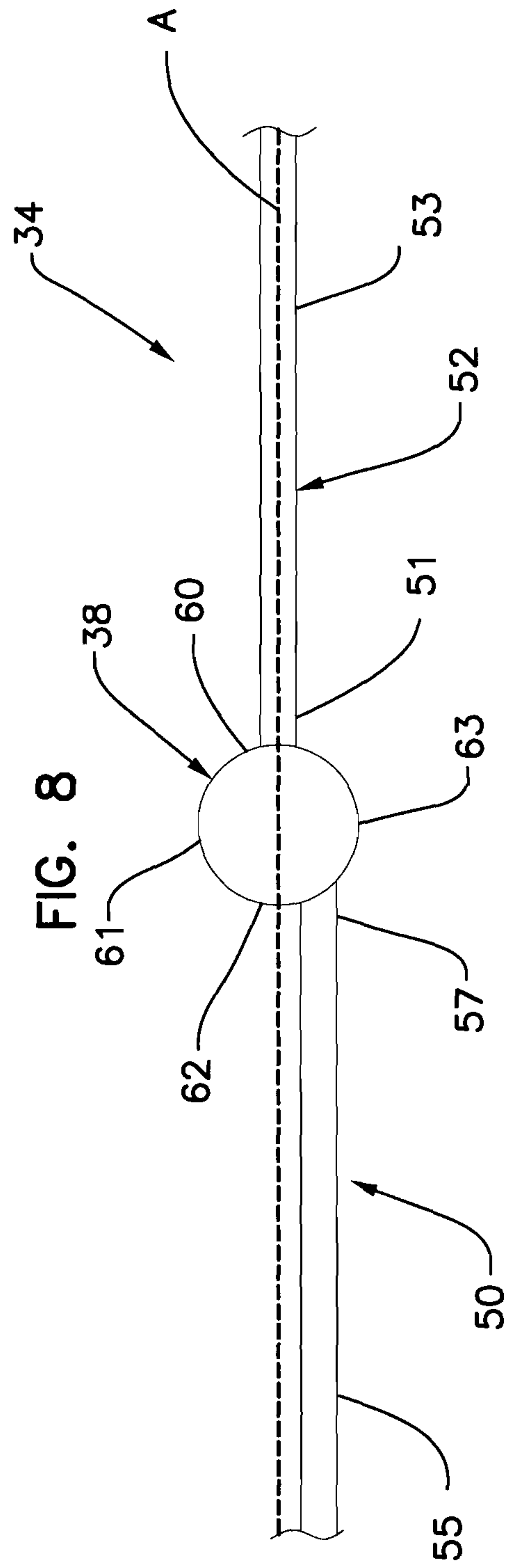
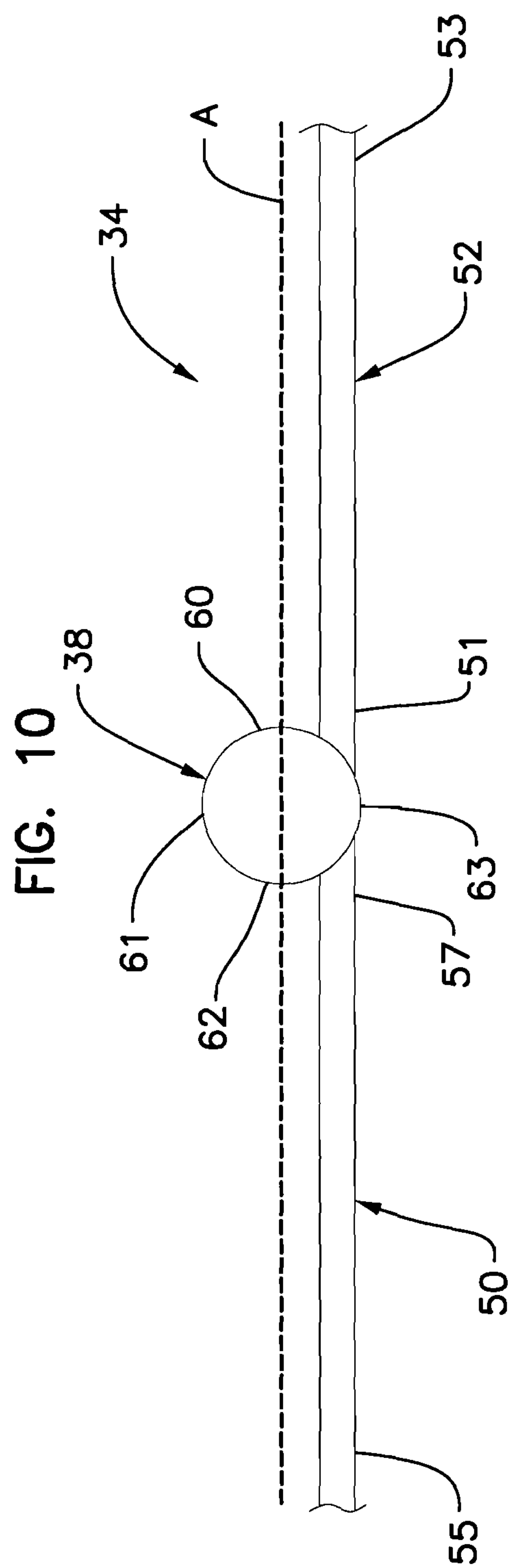
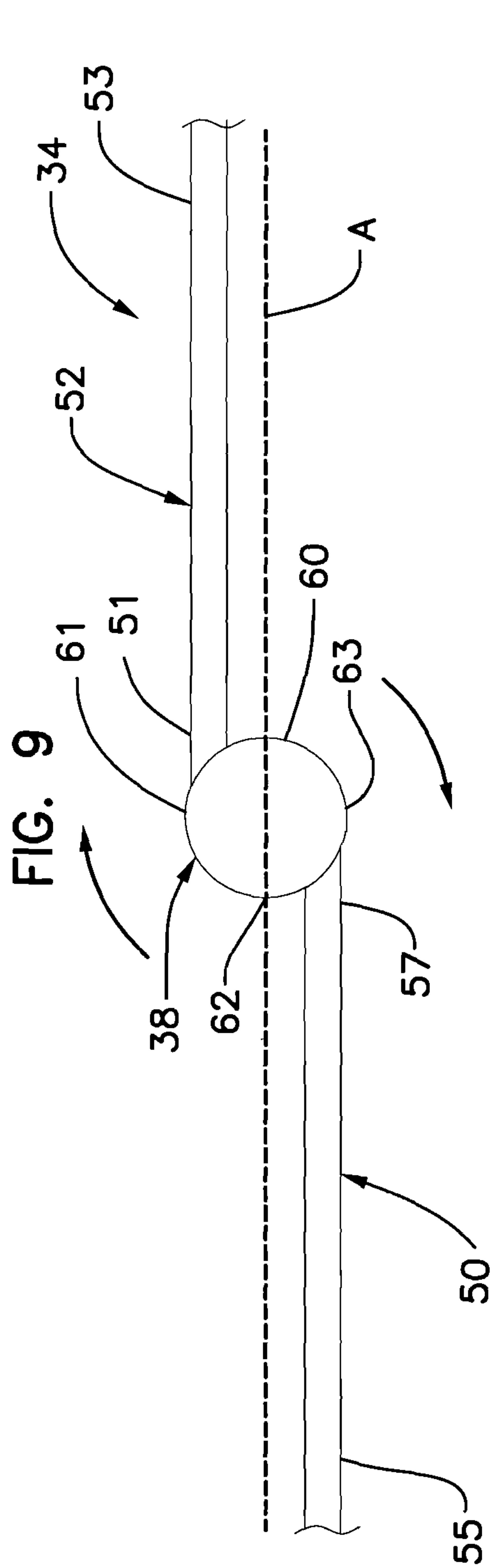
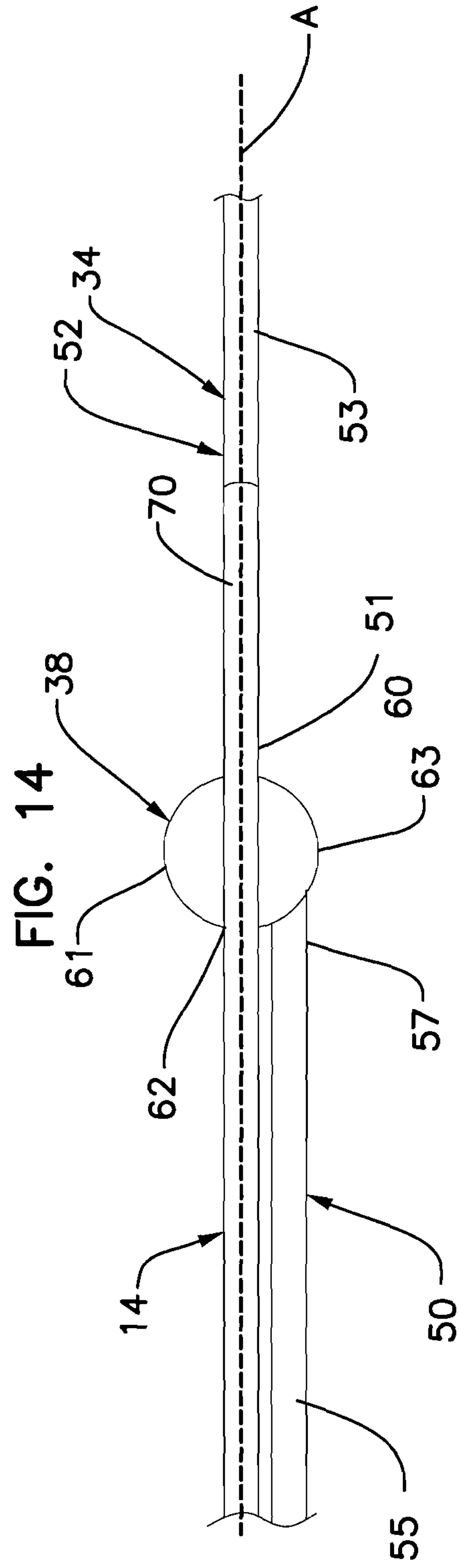
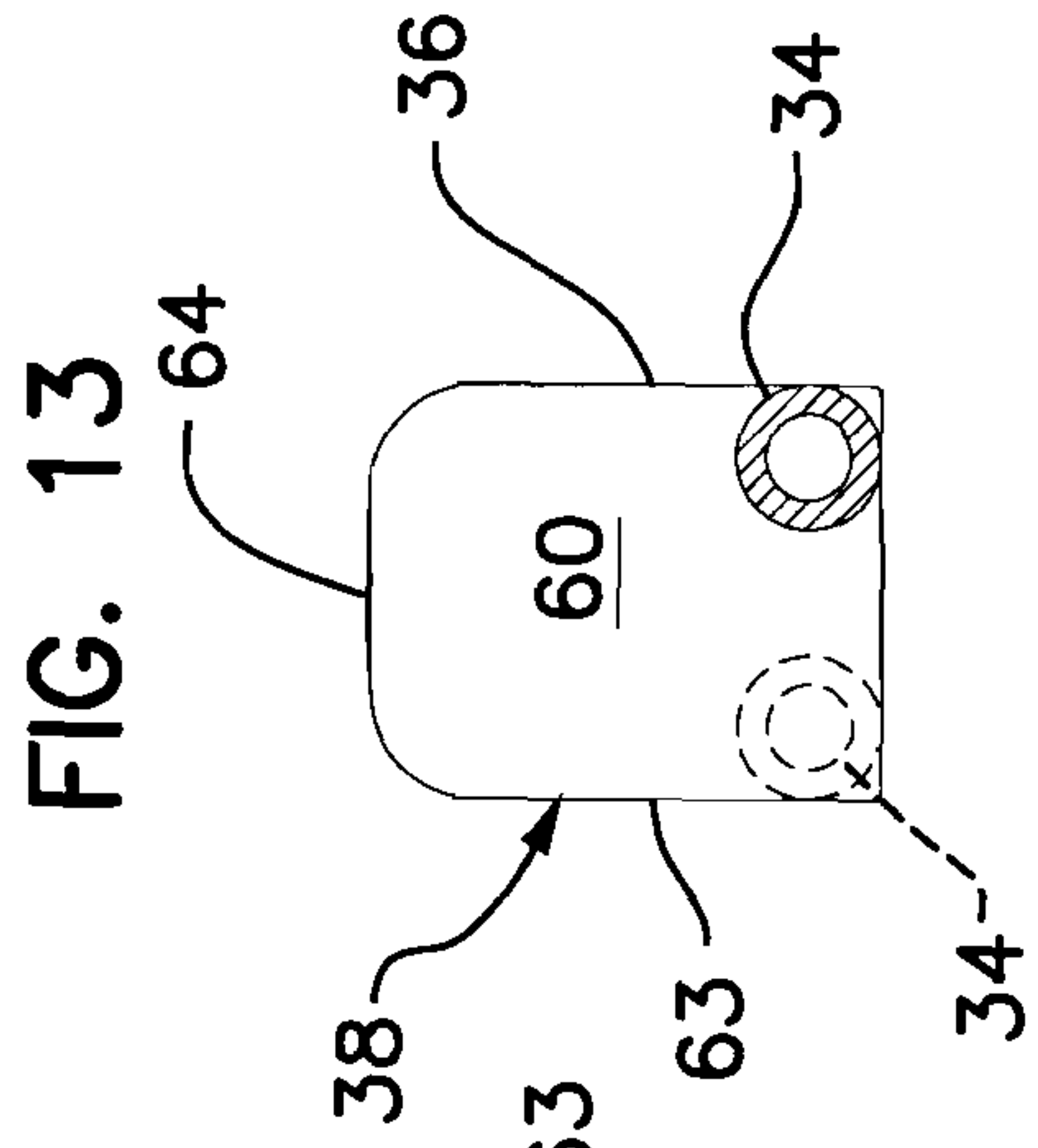
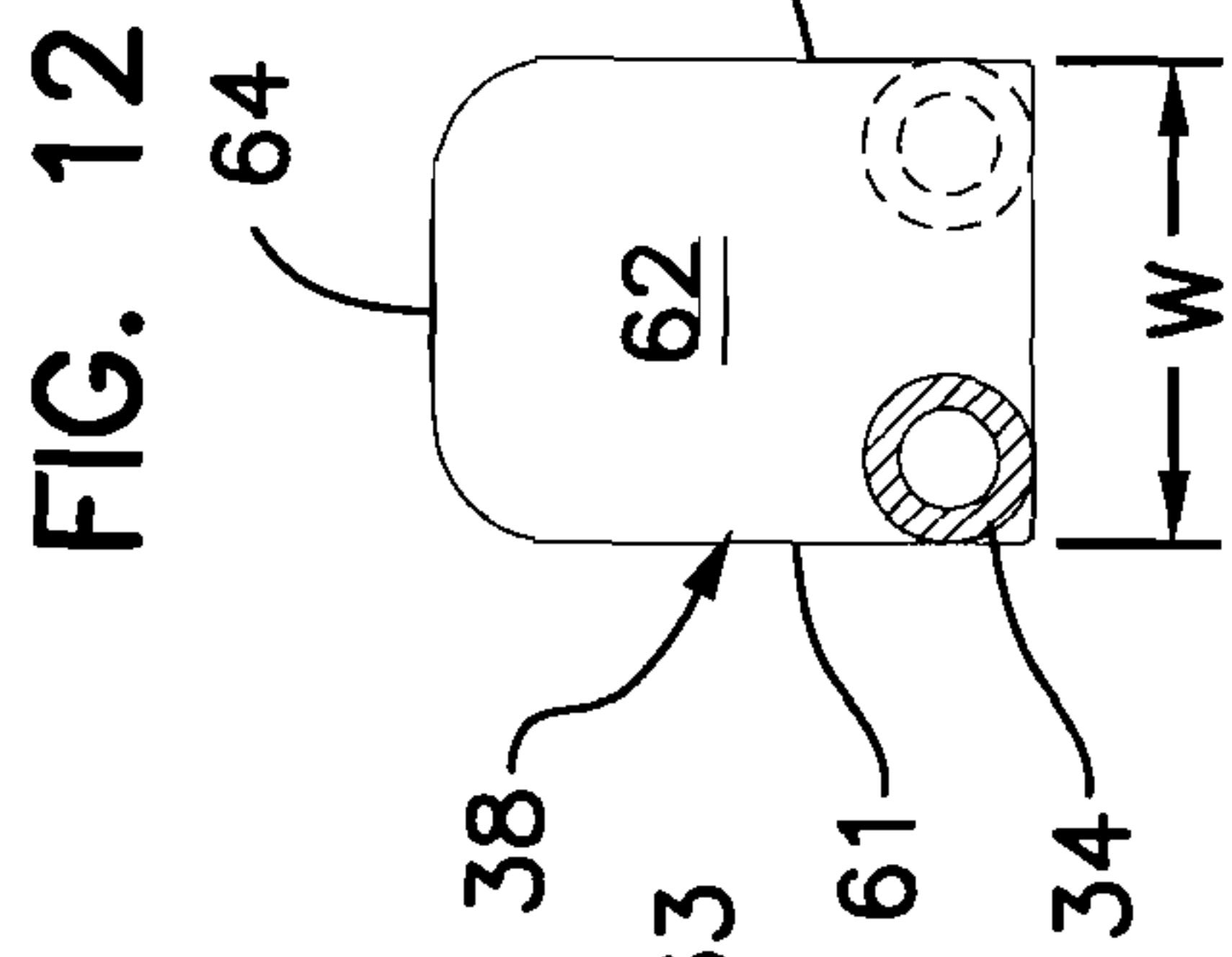
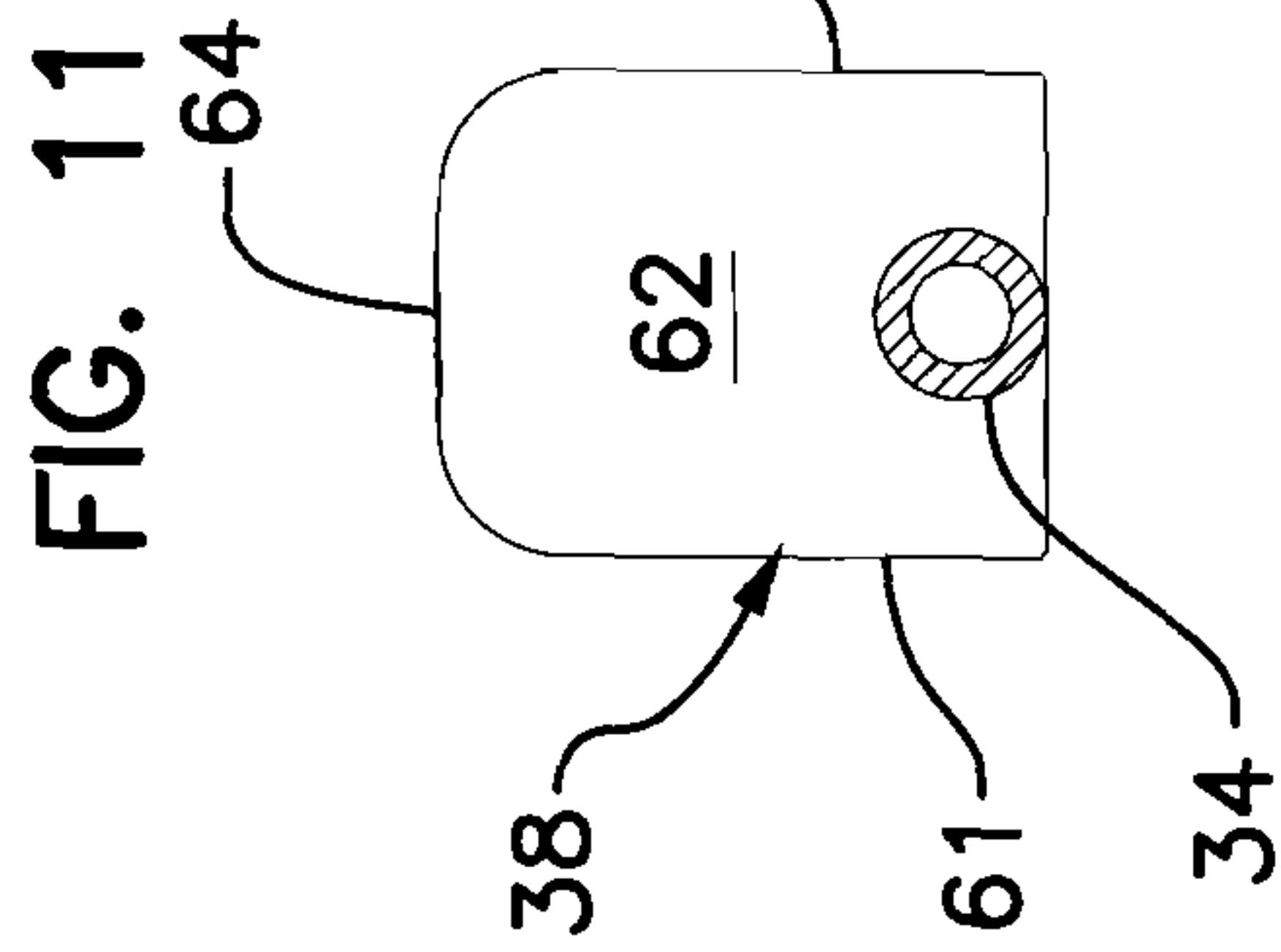


FIG. 8





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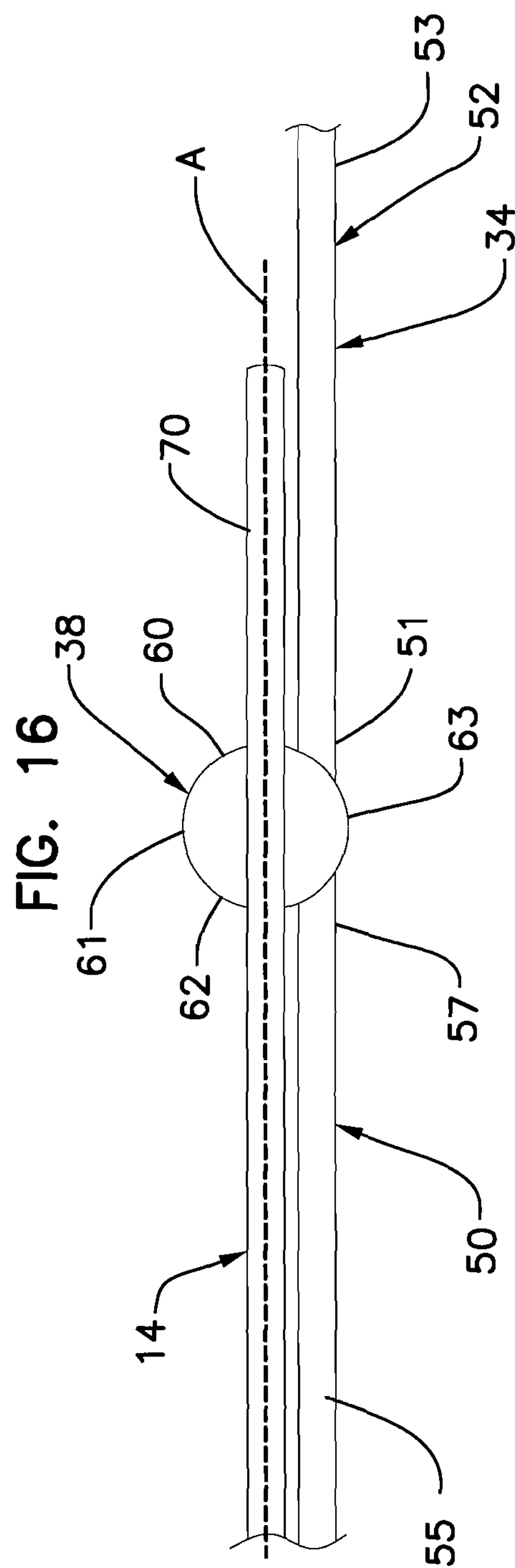
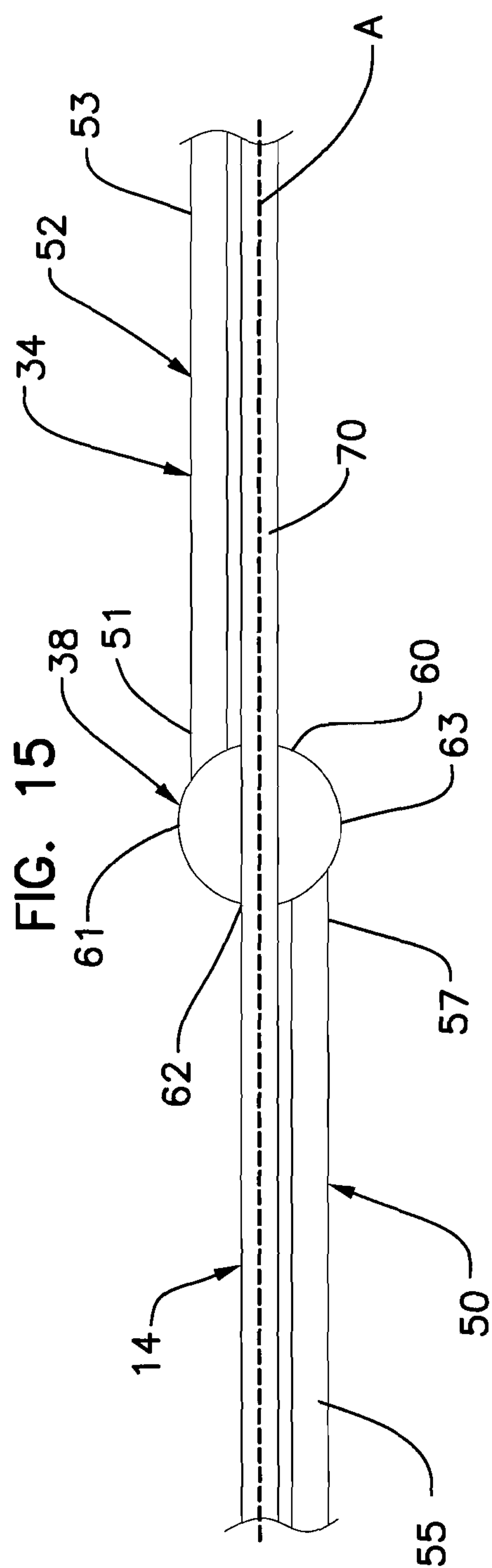


FIG. 17

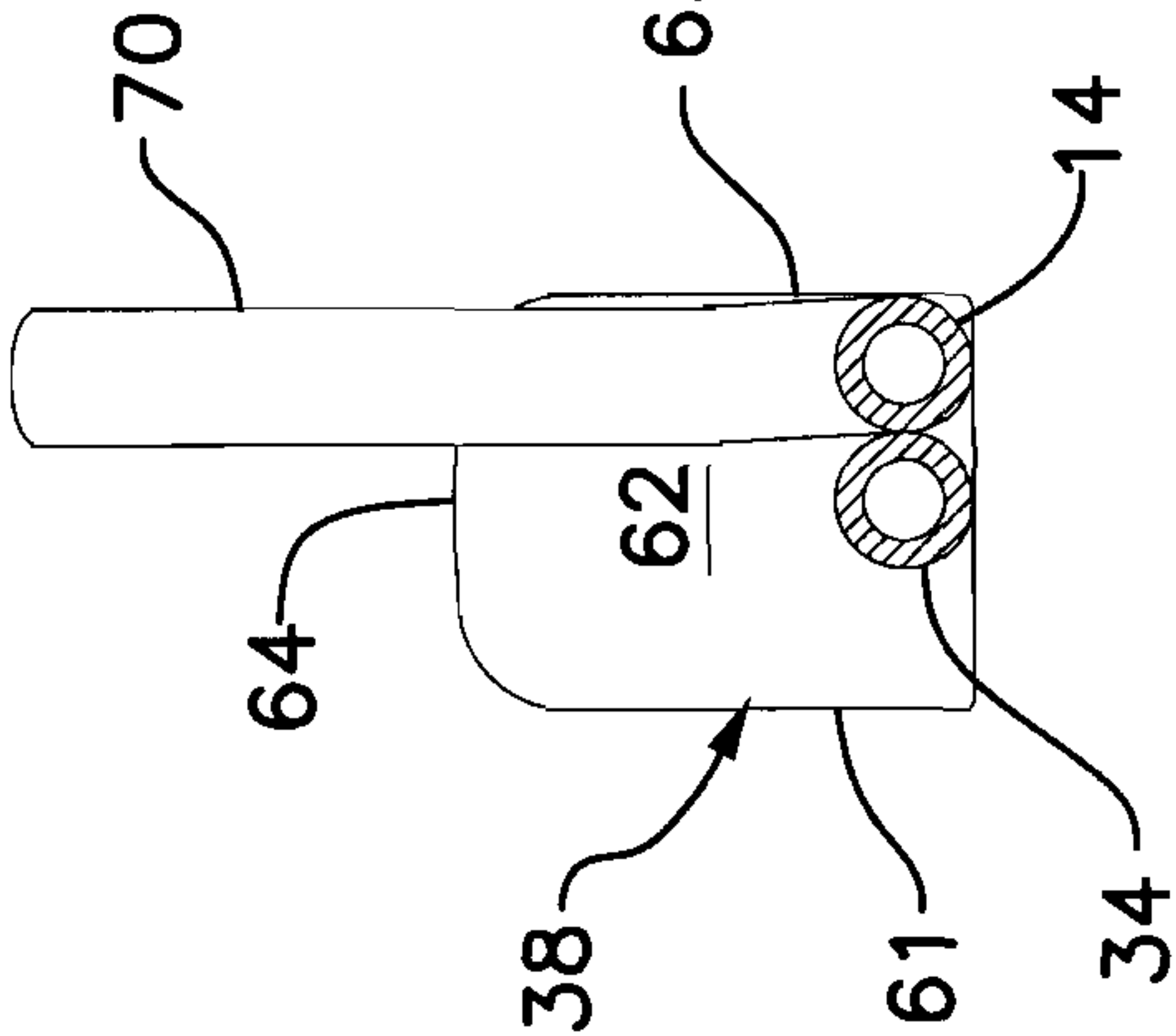


FIG. 18

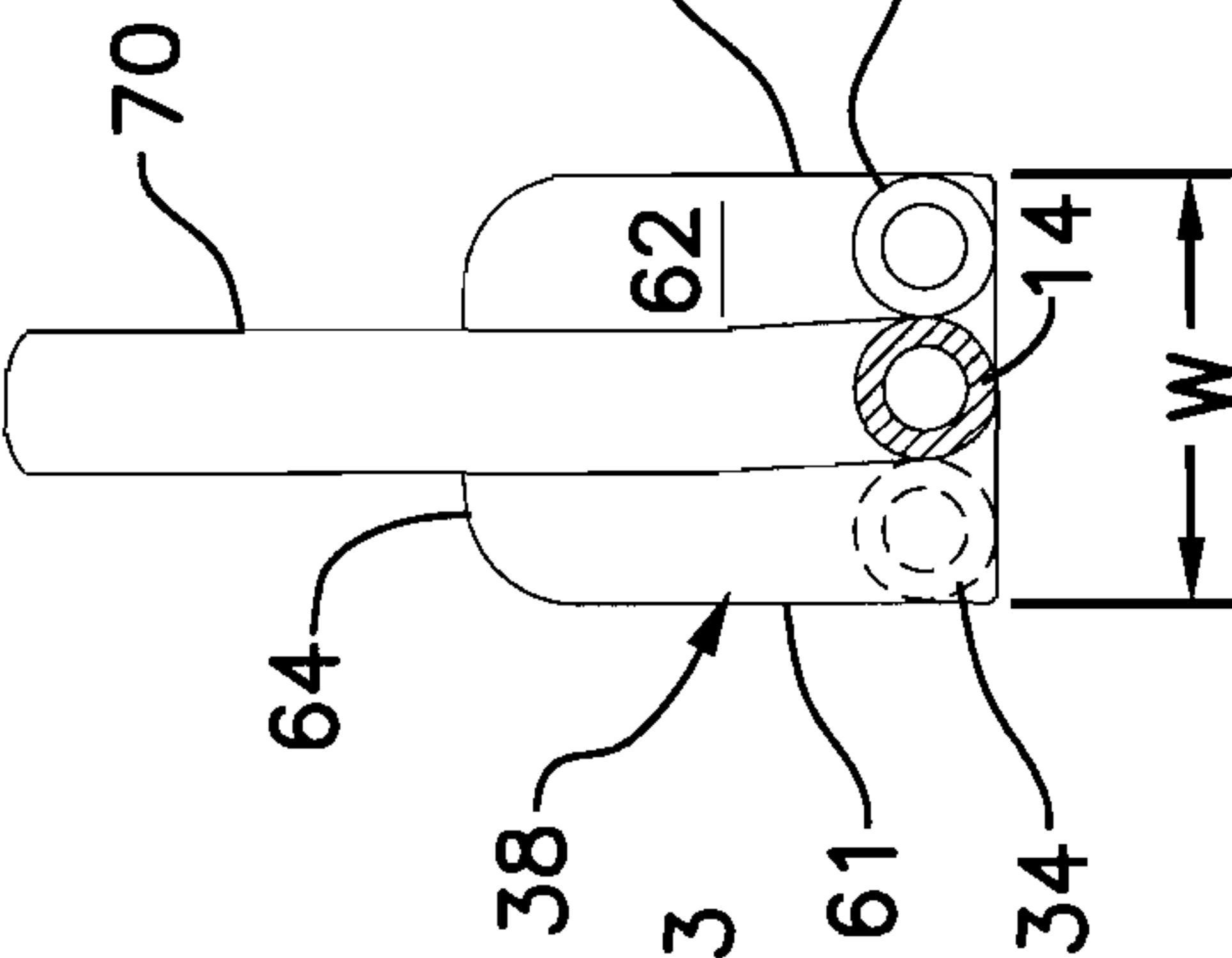


FIG. 19

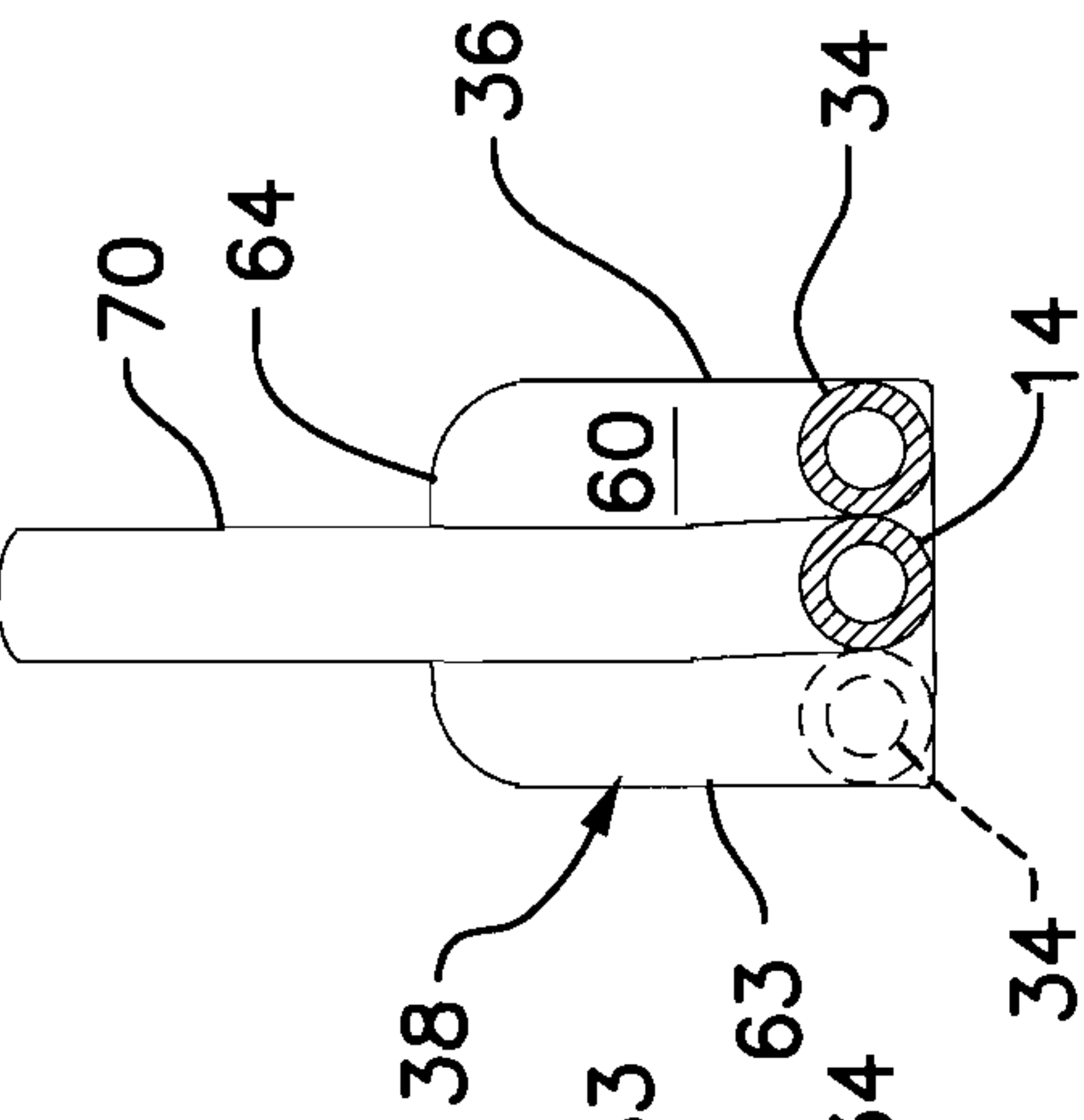


FIG. 20

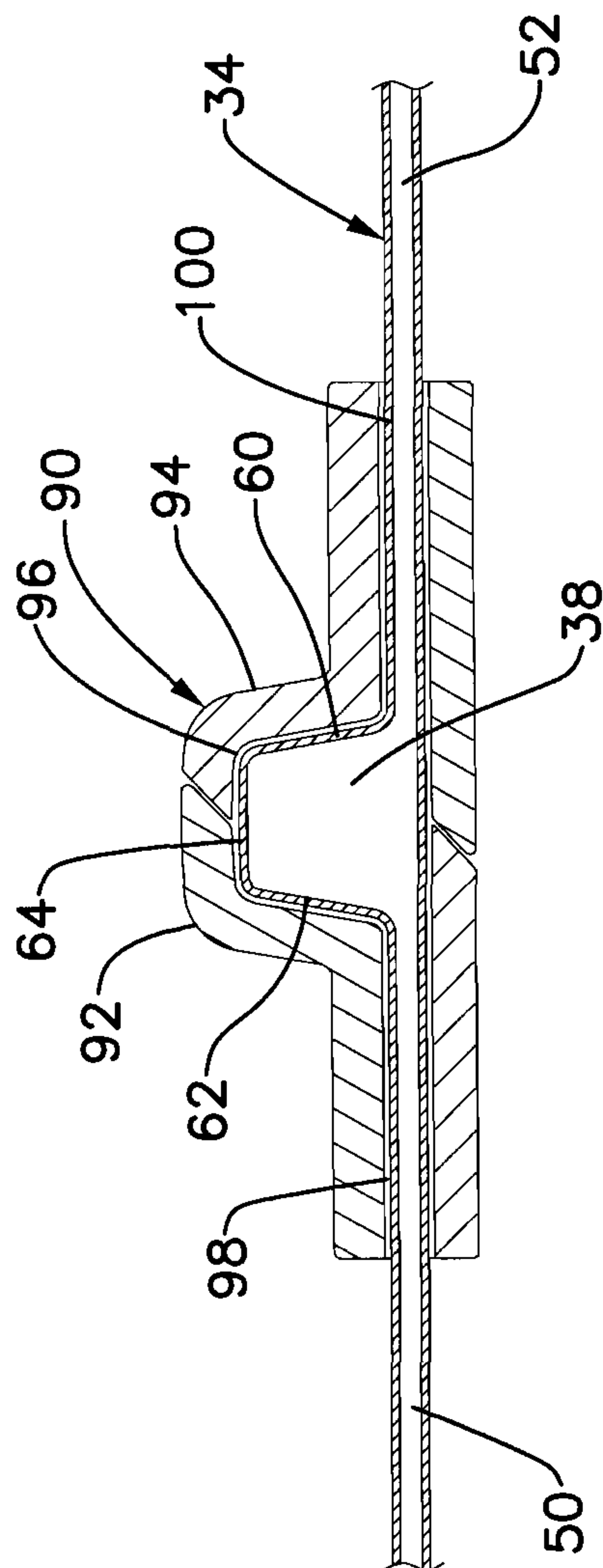


FIG. 21

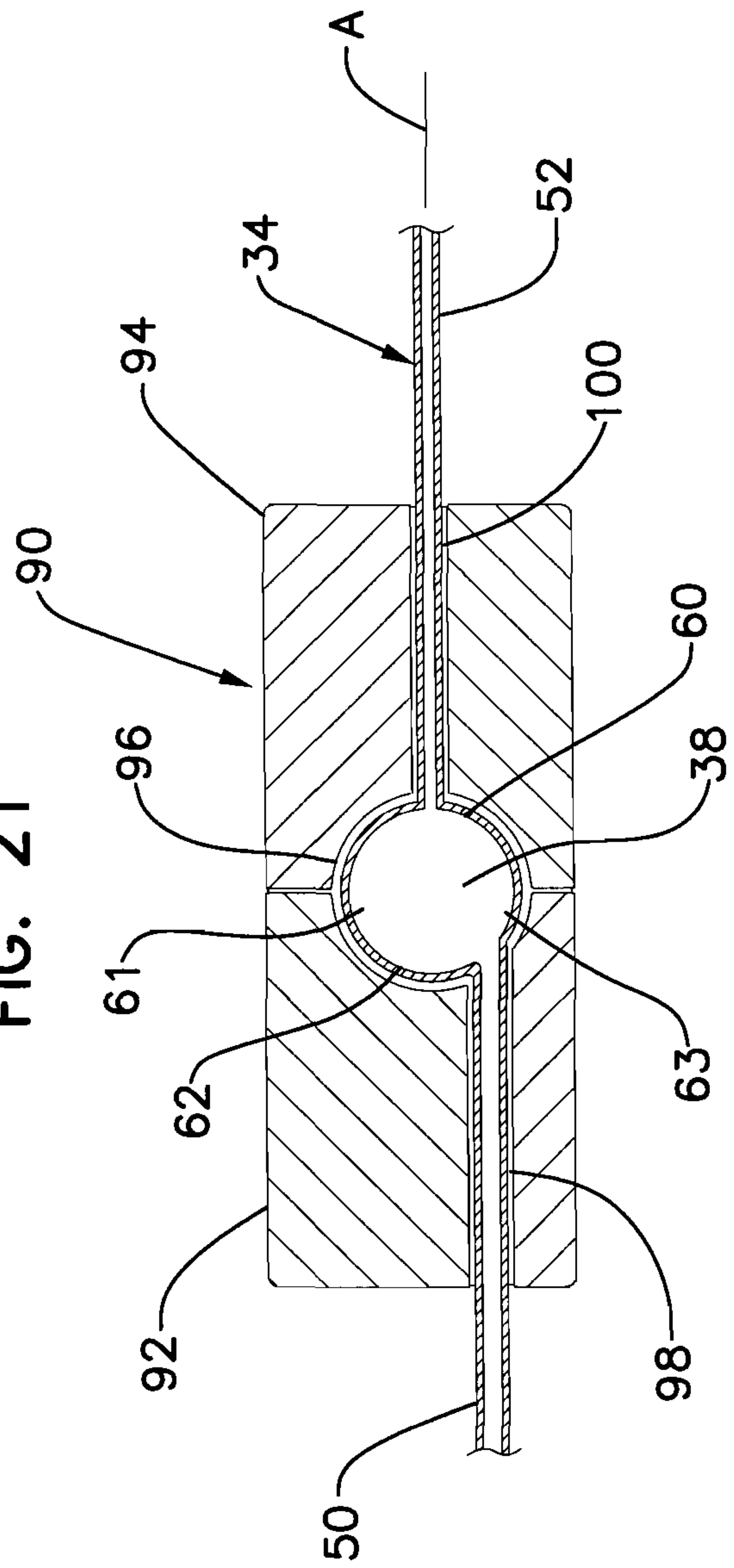


FIG. 22

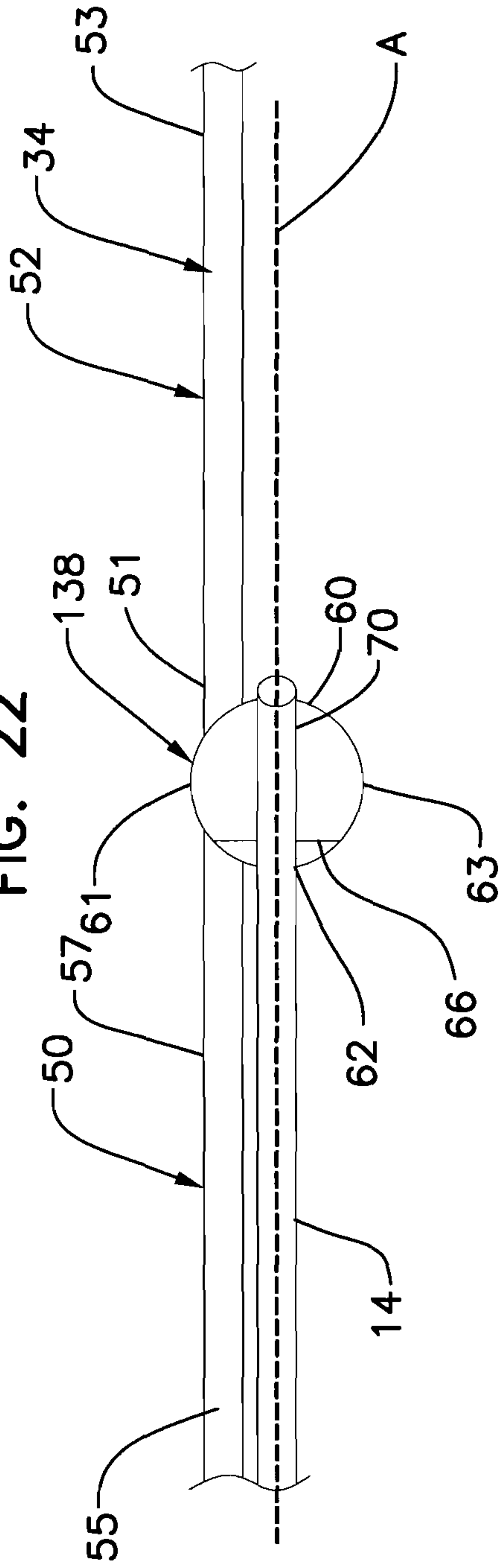


FIG. 23

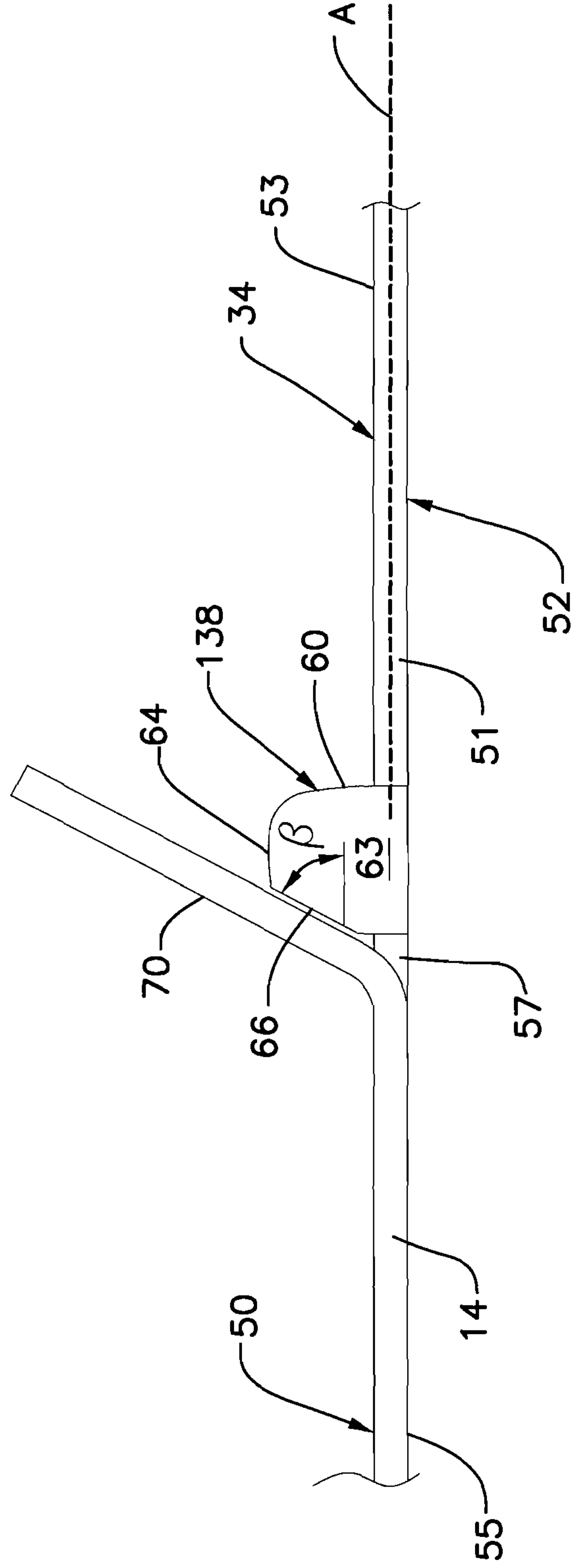


FIG. 24

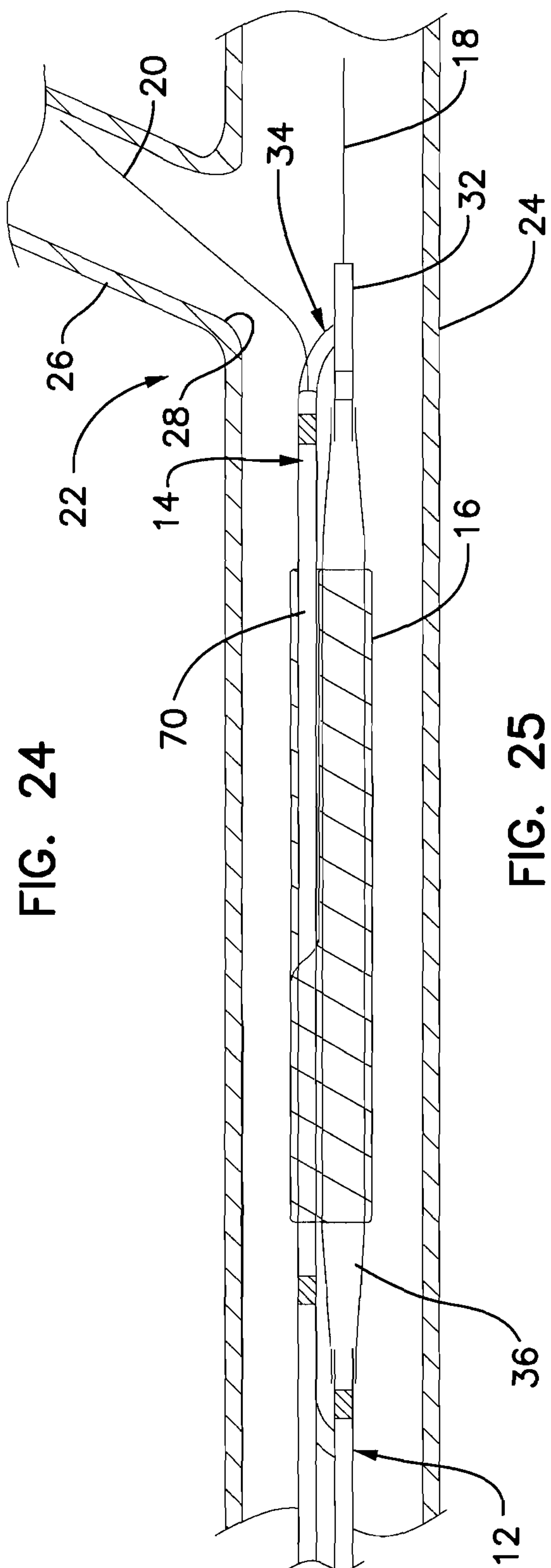


FIG. 25

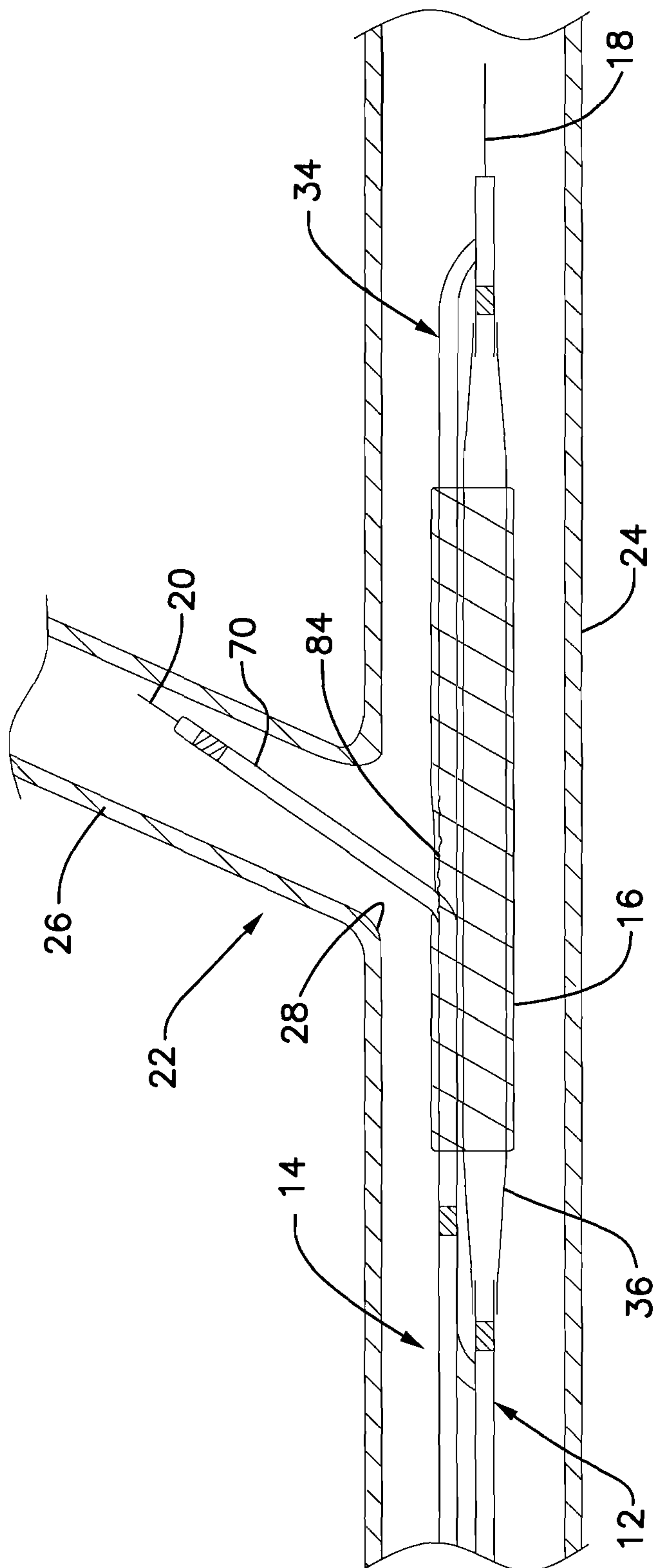


FIG. 26

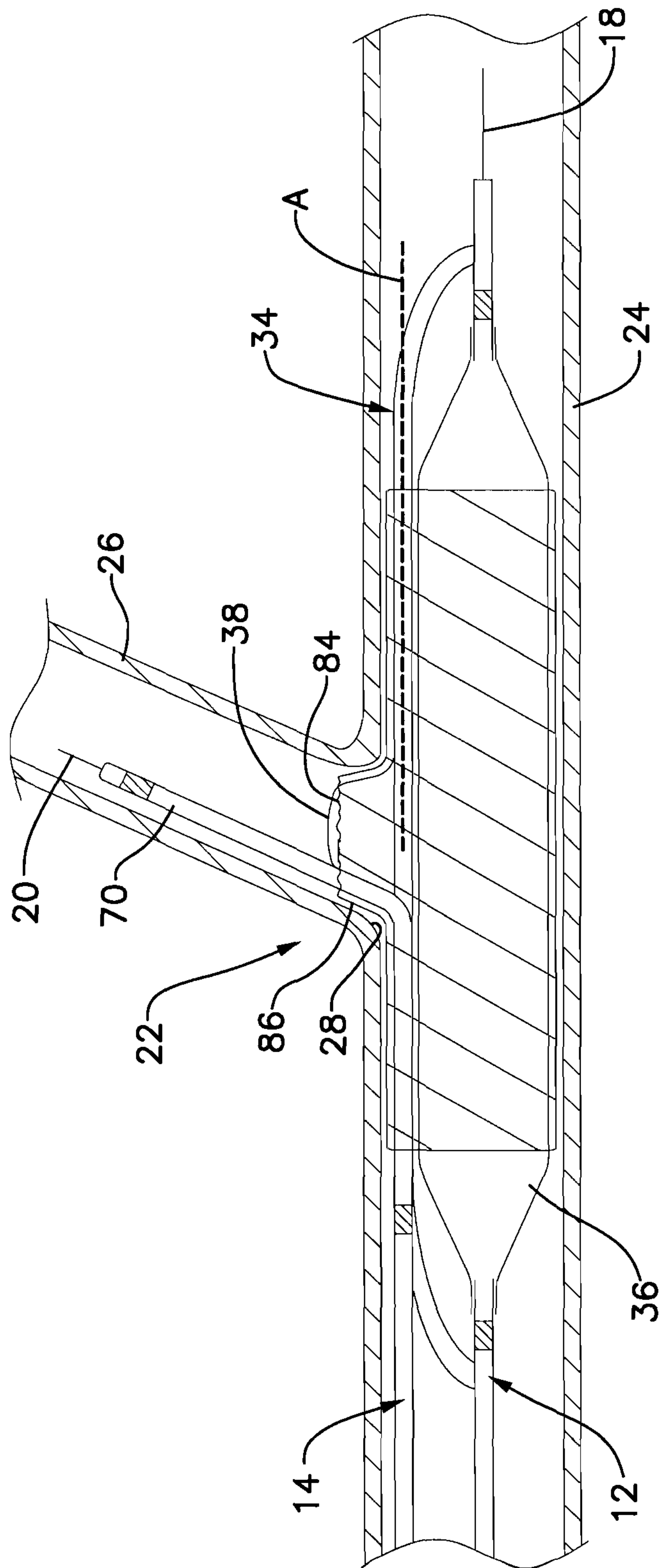


FIG. 27

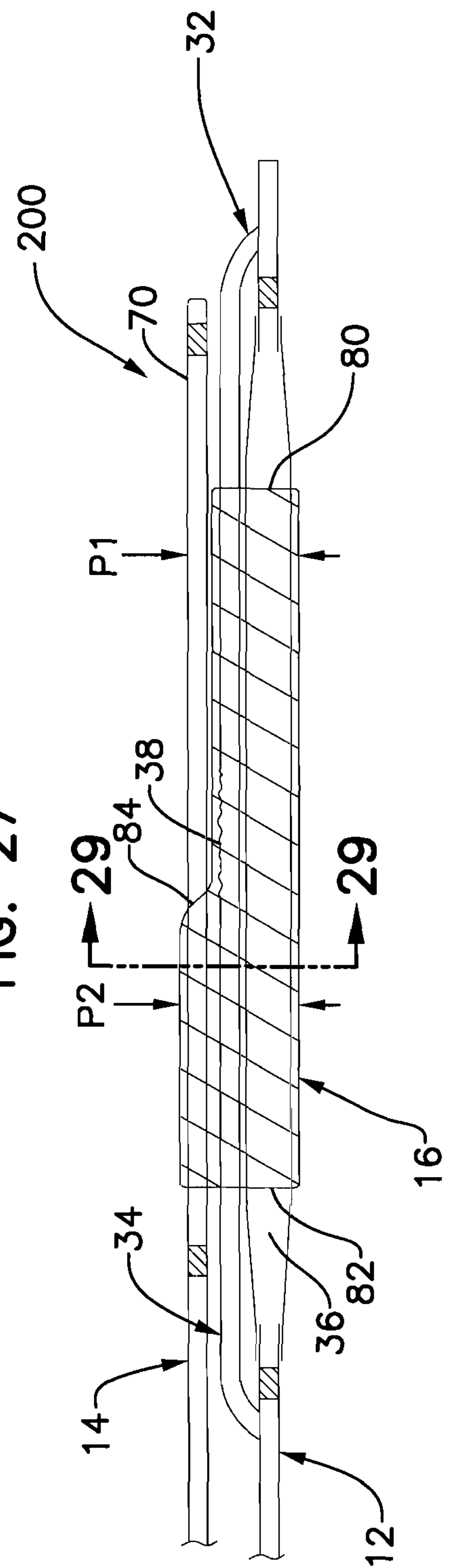


FIG. 28

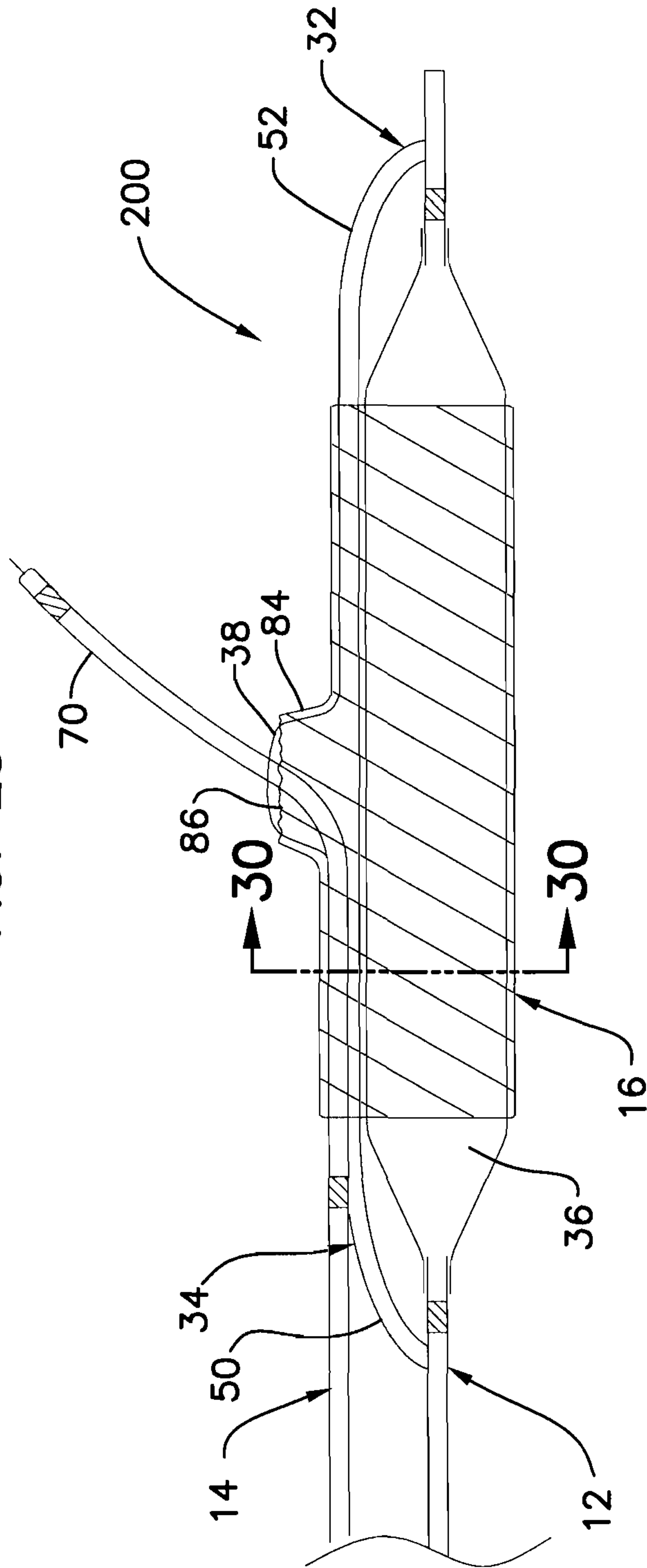


FIG. 30

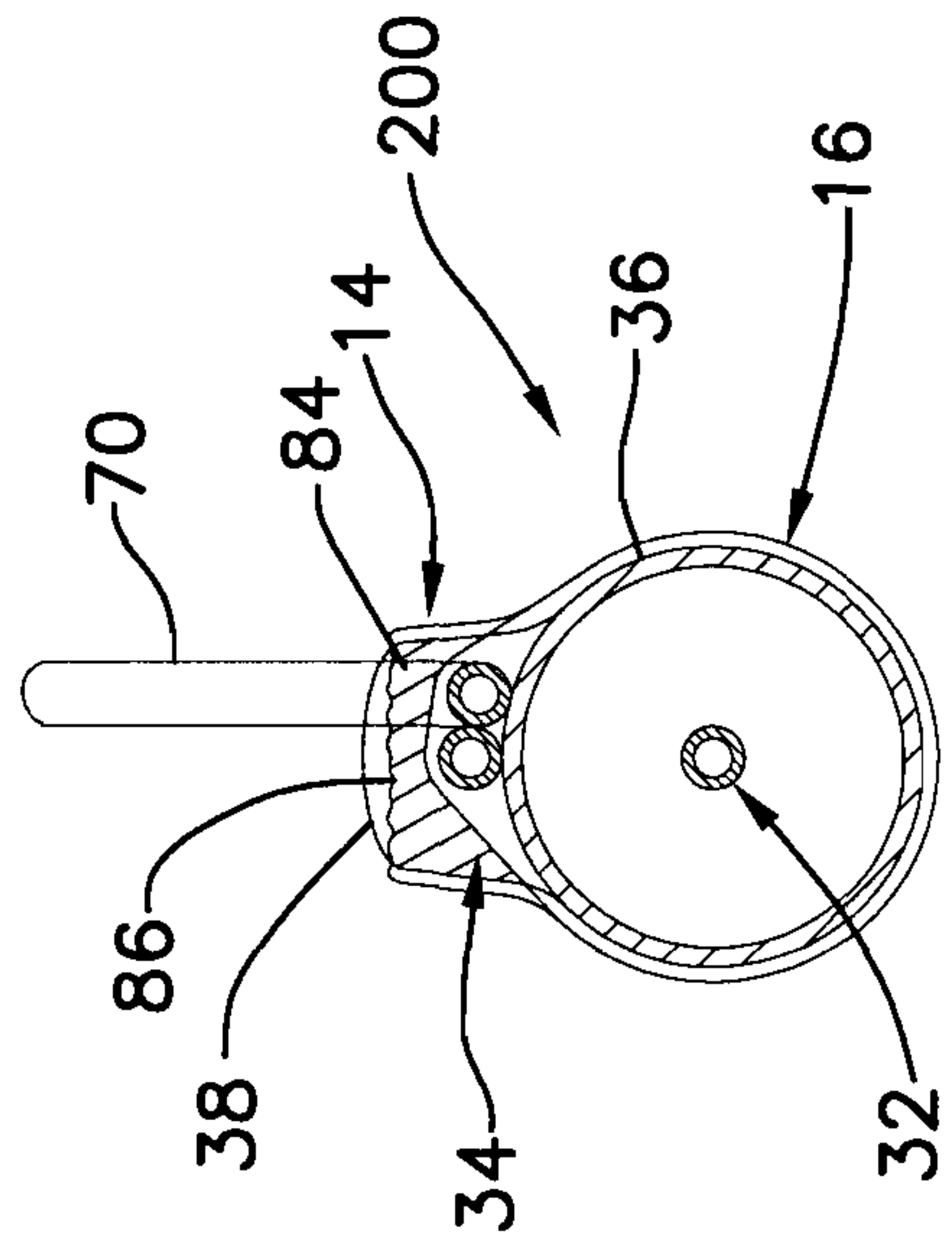
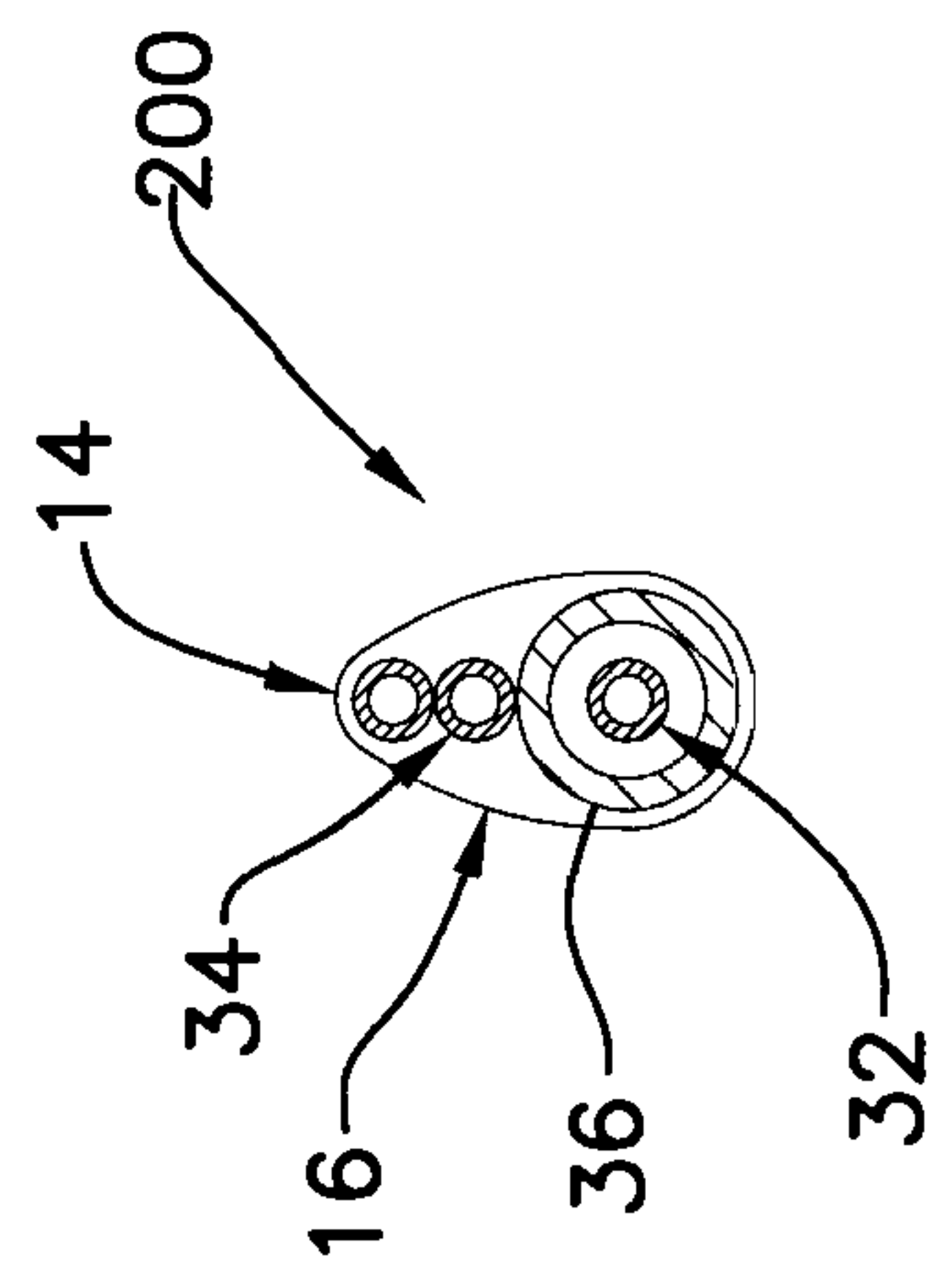


FIG. 29



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