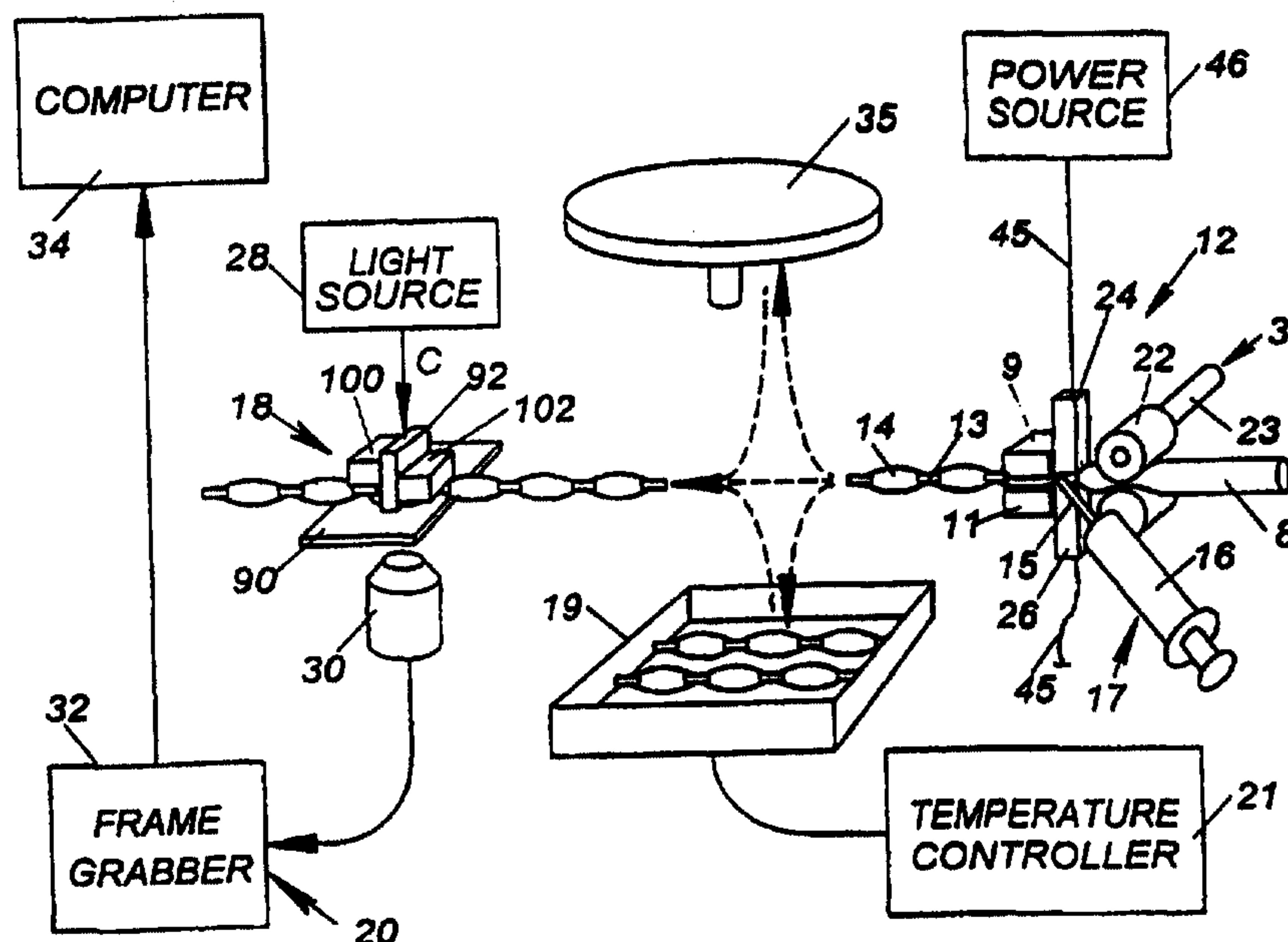


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(54) Title: FLUID SAMPLE TESTING SYSTEM



(57) Abstract

A sample testing system has a chamber sealing apparatus adapted to receive a chamber containing a fluid sample material and to seal portions of a chamber into segments. A reagent injector cartridge actuator is adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber. In certain preferred embodiments, a flow control device induces a flow of sample in a segment. A sensor generates a signal corresponding to a condition of the mixture.

FLUID SAMPLE TESTING SYSTEM

INTRODUCTION

The present invention is directed to a system for testing a fluid sample, and, more particularly, to a fluid sample testing system having improved automation, safety and efficiency.

BACKGROUND

5 Collection, transportation and pretreatment of fluid samples, such as blood samples, are currently done generally in a manual fashion. Blood is commonly collected in test tubes and samples from these test tubes are deposited in reaction chambers for testing. These tubes can be placed in an automated testing machine to perform testing using various assays. This process can be expensive, time consuming, and may lead to
10 human error, possibly leading to false test results. Current automated testing systems require large capital investment; incur high costs for reagents, disposables, operation, maintenance, service and training; and do not provide required sample pretreatment.

SUMMARY

The present invention provides a sample testing system which reduces or wholly
15 overcomes some or all of the aforesaid difficulties inherent in prior known devices. Particular aspects and advantages of the invention will be apparent to those skilled in the art, that is, those who are knowledgeable or experienced in this field of technology, in view of the following disclosure of the invention and detailed description of certain preferred embodiments.

20 The principles of the invention may be used to advantage to provide a sample testing system which is highly automated, thereby increasing efficiency, reducing costs, and increasing safety due to reduced handling of samples. A sample can be collected in a chamber which is then divided into a plurality of sealed segments. A reagent can be added to a segment and the segment can be inspected to detect a condition of the sample.

In accordance with a first aspect, a sample testing system has a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of the chamber. A reagent injector cartridge actuator is adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber. A sensor generates an output signal corresponding to a condition of a fluid sample material within a segment of a chamber.

In accordance with another aspect, a sample testing system has a tube sealing apparatus having a tube compression and sealing member to laterally seal a flexible plastic tube containing a fluid sample material, whereby a fluid-tight tubule containing a portion of the fluid sample material can be formed between axially spaced lateral seals. A reagent injector cartridge actuator is adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a tubule. A flow control device has a contact member movable into contact with a tubule to effect mechanically induced fluid flow within a fluid passageway in the tubule. An inspection system has a light detector to receive light passed through a tubule and to generate an output signal corresponding to a condition of the fluid sample material within a tubule.

In accordance with another aspect, a sample testing system has a tube sealing apparatus having a tube compression and sealing member to laterally seal a flexible plastic tube containing a fluid sample material, whereby a fluid-tight tubule containing a portion of the fluid sample material can be formed between axially spaced lateral seals. A reagent injector has at least one needle in fluid communication with a reagent reservoir, and a needle actuator to insert the needle into a tubule and inject a quantity of reagent into a tubule. A flow control device has a contact member movable into contact with a tubule to effect mechanically induced fluid flow within a fluid passageway in the tubule. An inspection system has a light detector to receive light passed through a tubule and to generate an output signal corresponding to a condition of the fluid sample material within a tubule.

In accordance with another aspect, a reagent cartridge has a housing and at least one reservoir in the housing. At least one needle in the housing is in fluid

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communication with one of the reagent reservoirs. A needle actuator inserts the needle into a tubule and injects a quantity of reagent.

In accordance with yet another aspect, a sample testing tubule has a length of flexible plastic tube having fluid-tight lateral seals at axially spaced locations to define a fluid-tight fluid sample chamber between the lateral seals containing a fluid sample material. A self-sealing injection channel is formed in the tubule, the injection channel being normally substantially free of fluid sample material and capable of fluid communication with the fluid sample material in the tubule.

In accordance with another aspect, a method of performing a sample assay includes the following steps: collecting a sample of fluid material into a length of substantially transparent, flexible, heat-sealable, plastic tube; inserting the tube into a sample testing machine having a tube sealing apparatus, a reagent injector having at least one needle in fluid communication with a reagent reservoir and a needle actuator to insert the needle into a tubule and inject a quantity of reagent, a flow control device having a contact member movable into contact with a tubule to effect mechanically induced fluid flow within the tubule, and an inspection system having a light detector to receive light passed through a tubule and to generate an output signal corresponding to a condition of the sample material within a tubule; actuating the tube sealing apparatus to seal lengths of the tube into tubules; actuating the needle actuator to insert the needle into a selected tubule and inject reagent to form a mixture of sample material and reagent in the selected tubule; actuating the flow control device to mix the mixture of sample material and reagent; and actuating the inspection system to inspect the mixture and to generate an output signal corresponding to a condition of the mixture.

In accordance with another aspect, there is provided a sample testing system comprising, in combination:

a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of the chamber;

a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber;

3a

a flow control device comprising at least one contact member movable into contact with a segment of a chamber to effect mechanically induced fluid flow within a flow passageway in a segment of a chamber; and

a sensor to generate an output signal corresponding to a condition of a fluid sample material within a segment of a chamber.

In accordance with another aspect, there is provided a sample testing system comprising, in combination:

tube sealing apparatus having a tube compression and sealing member to laterally seal a flexible plastic tube containing a fluid sample material, whereby a fluid-tight tubule containing a portion of the fluid sample material can be formed between axially spaced lateral seals; the tube sealing apparatus comprising a first sealing head comprising the tube compression and sealing member; and a second sealing head, at least one of the first and second sealing heads being movable toward the other of the sealing heads to compress a section of flexible plastic tube positioned between the first and second sealing heads to create a sample-free zone in the tube, wherein the tube compression and sealing member is operatively connected to a power source to heat a sealing zone of a tube located in the sample-free zone to form a fluid-tight lateral seal in the tube;

a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a fluid-tight tubule; and

a sensor to generate an output signal corresponding to a condition of a fluid sample material within a fluid-tight tubule.

In accordance with another aspect, there is provided a sample testing system comprising, in combination:

a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of the chamber;

a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber;

a sensor to generate an output signal corresponding to a condition of a fluid sample material within a segment of a chamber;

3b

a pair of electrodes adapted to have a predetermined voltage difference, and
an electrode actuator to insert the pair of electrodes into a segment of a chamber,
wherein the sensor is responsive to electrophoretic light emitted from within a segment of a
chamber.

In accordance with another aspect, there is provided a sample testing system
comprising, in combination:

a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-
tight segments of the chamber;

a segment of the chamber including a coating on an outside surface of the segment to
increase the transmission of light through the segment;

a reagent injector cartridge actuator adapted to receive a reagent injector cartridge
having at least one needle in fluid communication with a reagent reservoir, and to move a
reagent injector cartridge to inject a quantity of reagent into a segment of a chamber; and

a sensor to generate an output signal corresponding to a condition of a fluid sample
material within a segment of a chamber.

In accordance with another aspect, there is provided a sample testing system
comprising, in combination:

a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-
tight segments of the chamber;

a reagent injector cartridge actuator adapted to receive a reagent injector cartridge
having at least one needle in fluid communication with a reagent reservoir, and to move a
reagent injector cartridge to inject a quantity of reagent into a segment of a chamber;

an incubation chamber to retain a segment of a chamber for a predetermined period of
time; and

a sensor to generate an output signal corresponding to a condition of a fluid sample
material within a segment of a chamber.

3c

From the foregoing disclosure, it will be readily apparent to those skilled in the art, that is, those who are knowledgeable or experienced in this area of technology, that the present invention provides a significant technological advance. Preferred embodiments of the fluid sample testing system of the present invention can provide increased efficiency, reduced costs, and increase safety. These and additional features and advantages of the invention disclosed here will be further understood from the following detailed disclosure of certain preferred embodiments.

BRIEF DESCRIPTION OF THE DRAWINGS

Certain preferred embodiments are described in detail below with reference to the appended drawings wherein:

Fig. 1 is a partially schematic perspective view of a sample testing system in accordance with a preferred embodiment of the present invention;

Fig. 2 is a schematic representation of the components of the sample testing system of Fig. 1;

Fig. 3 is a schematic perspective view, partially in phantom, of a tube sealing apparatus of the testing system of Fig. 1;

Fig. 4 is a schematic elevation view, shown partially cut away, of a tube being compressed by the tube sealing apparatus of Fig. 3;

Fig. 5 is a schematic elevation view, shown partially cut away, of a tube being sealed by the tube sealing apparatus of Fig. 3;

Fig. 6 is a schematic plan view of a sealing head of the tube sealing apparatus of Fig. 3;

Fig. 7 is a schematic plan view of a plurality of tubules formed in a length of tube by the tube sealing apparatus of Fig. 3;

Fig. 8 is a schematic plan view of an alternative embodiment of a sealing head of the tube sealing apparatus of Fig. 3;

Fig. 9 is a schematic plan view of another alternative embodiment of a sealing head of the tube sealing apparatus of Fig. 3;

Fig. 10 is a schematic section view of a reagent cartridge suitable for use in the sample testing system of Fig. 1;

Fig. 11 is a schematic section view of an alternative embodiment of a reagent cartridge for the sample testing system of Fig. 1;

Fig. 12 is a schematic section view of the reagent cartridge of Fig. 11 shown injecting reagent into a tubule;

Fig. 13 is a schematic section view of another alternative embodiment of a reagent cartridge of the sample testing system of Fig. 1;

Fig. 14 is a schematic section view of yet another alternative embodiment of a reagent cartridge of the sample testing system of Fig. 1;

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Fig. 15 is a schematic elevation view of a flow control device and inspection system of the sample testing system of Fig. 1;

Fig. 16 is a schematic elevation view of an alternative embodiment of the flow control device of the sample testing system of Fig. 1;

5 Fig. 17 is a schematic elevation view of another alternative embodiment of the flow control device of the sample testing system of Fig. 1;

Fig. 18 is a schematic elevation view of yet another alternative embodiment of the flow control device of the sample testing system of Fig. 1;

10 Fig. 19 is a schematic elevation view of an alternative embodiment of the inspection system of the sample testing system of Fig. 1;

Fig. 20 is a schematic elevation view of another alternative embodiment of the inspection system of the sample testing system of Fig. 1;

Fig. 21 is a schematic elevation view of a coating being applied to a tubule of the present invention;

15 Fig. 22 is a schematic perspective view of a reagent cartridge and a tube divided into tubules, suitable for the sample testing system of Fig. 1;

Fig. 23 is a schematic perspective view of one preferred embodiment of a tube of the present invention and a drawing device into which the tube is placed;

20 Fig. 24 is a schematic elevation view of an alternative embodiment of the tube sealing apparatus of Fig. 1; and

Fig. 25 is a schematic plan view of an alternative embodiment of a tubule of the present invention, shown with a pressure gate between compartments of the tubule.

25 The figures referred to above are not drawn necessarily to scale and should be understood to present a representation of the invention, illustrative of the principles involved. Some features of the sample testing system depicted in the drawings have been enlarged or distorted relative to others to facilitate explanation and understanding. The same reference numbers are used in the drawings for similar or identical components and features shown in various alternative embodiments. Sample testing system as disclosed herein, will have configurations and components determined, in part, by the intended
30 application and environment in which they are used.

DETAILED DESCRIPTION OF CERTAIN PREFERRED EMBODIMENTS

The present invention has many uses which will become readily apparent to those skilled in the art, given the benefit of this disclosure. Sample material to tested may be, e.g., blood, cell suspensions, biofluids or other fluids. Exemplary tests to be performed on fluid samples include clinical diagnosis, therapeutic monitoring, and screening of chemical compounds for discovery of new drugs. The following discussion will discuss blood testing specifically for purposes of illustration.

The present invention provides for a chamber containing a fluid sample to be divided into a plurality of segments, with fluid-tight seals separating adjacent segments from one another. It is considered to be a highly advantageous feature of certain preferred embodiments that a chamber into which a fluid sample is drawn, e.g., a tube into which a patient's blood is drawn, can itself then also be the testing or reaction chamber within which that blood or other fluid sample is tested, without ever having to remove the blood or fluid sample from the chamber.

Referring to Fig. 1, a testing machine according to the present invention is shown generally by the reference numeral 2. Testing machine 2 comprises a housing 4 having an entry port 6 on a front side thereof for receiving a chamber containing a fluid sample. In the illustrated embodiment, the chamber is a tube 8 from a blood bag 10. Tube 8 is preferably a flexible, thermoplastic, substantially transparent tube having an inner diameter of approximately 1 mm to 5 mm, preferably approximately 3-4 mm. Tube 8 may be formed of polyvinylchloride (PVC) or other suitable material. A control panel 7 is located on the front of housing 4 to receive information, such as information read from bar code labels or keyed data, and a monitor 5 displays operating information, such as the results of testing. A tube sealing apparatus 12, described in greater detail below, is contained within housing 4 for sealing portions of tube 8 into tubules 14. Reagent cartridge 60 is loaded into a reagent cartridge actuator 49 in housing 4, with reagent from reservoirs 16 contained within reagent cartridge 60 being added to tubules 14 (described in greater detail below). A sensor 41 in housing 4 reads a bar code label 73 (seen in Fig. 22) on reagent cartridge 60 which provides information identifying the particular reagent or reagents in reagent cartridge 60 as well as information regarding test procedures associated with the particular reagent or reagents. Mixing device or flow control device

18, seen in Fig. 2 and described in greater detail below, is also contained within housing 4 for creating a fluid passageway to allow the flow of cells within tubule 14. Computerized microscopic inspection system 20 is mounted in housing 4 to view and analyze the flow of cells within tubule 14. In certain preferred embodiments, multiple testing machines 2 may be connected to computer analysis and system control components of inspection system 20, either directly, or via a computer network. In certain preferred embodiments, flow control device 18 may not be present, or may not be employed if present. In such an alternative embodiment, inspection system 20 inspects a sample within tubule 14 without a flow of cells within the sample being created.

10 A tube advancement system 3 is provided to support and control forward movement of tube 8 through testing machine 2. Suitable tube advancement systems will become readily apparent to those skilled in the art, given the benefit of this disclosure. In the embodiment illustrated in Fig. 2, tube advancement system 3 comprises a pair of rotating wheels 22 which rotate in opposite directions to advance the tube. At least one wheel 22 is connected to and driven by output shaft 23 of a motor which is not shown. 15 Tube 8 is inserted between rotating wheels 22 and advanced into tube sealing apparatus 12. The volume of sample within each tubule 14 is controlled by compressing tube 8. Specifically, upper plunger 9 and lower plunger 11 are spaced apart from one another and movable toward one another to partially compress a tubule 14 positioned therebetween prior to it being sealed. An upper, or first sealing head 24 and a lower, or second sealing head 26 compress a portion of tube 8 and then use radio frequency energy to seal tube 8, forming lateral seals 13 between adjacent tubules 14. Lateral seals, as used herein, refer to seals which separate axially adjacent portions of tube 8. In a preferred embodiment, the lateral seals extend substantially perpendicular to a longitudinal axis of tube 8. Seals 25 13 are fluid-tight seals, that is, seals 13, under normal operating conditions, prevent the flow of fluid through the seal. Each tubule 14 contains a sample of blood. The length of each tubule 14 is preferably approximately 3 to 15 mm, and more preferably about 5 to 10 mm. Reagent is added to tubule 14 via needle 15 of injector 17.

Tubules 14 then advance to one of an incubation chamber 19, a centrifuge 35, or 30 flow control device 18. Flow control device 18 forms a pair of reservoir zones in tubule 14 with a thin fluid passageway extending between the reservoirs. Light from light

source 28 is projected through the tubule 14 in flow control device 18. A camera with a microscopic lens 30 captures images of blood cell aggregates flowing from one reservoir zone to the other through the thin passageway. It sends the images to a frame grabber 32, which in turn sends the images to programmable control system or computer 34 for analysis. The results of the testing done in computer 34 may be transmitted to display 7, seen in Fig. 1, for reading by an operator. In other preferred embodiments, the results of the testing may be stored for later retrieval, or forwarded to another computer or other device, e.g. a printer for preparing a hard copy of the results.

Centrifuge 35 is provided to separate components of the sample in a length of tube 8 in a known fashion. A length of tube 8, typically longer than a typical tubule 14, is conveyed to centrifuge 35 via suitable conveying means. Once the components of the sample in the length of tube 8 have been separated, the length of tube is sealed into tubules 14 providing a fluid-tight seal between the different components. The length of tube is sealed either by a tube sealing apparatus at centrifuge 35, or it may be advanced to tube sealer 12 by suitable conveying means for sealing. Centrifuge 35 may also be used during testing in order to perform certain assays.

In certain preferred embodiments, selected tubules 14 may be stored in incubation chamber 19 prior to advancing to flow control device 18. Incubation chamber 19 may provide temperature control of tubules 14, and may allow the addition of a second reagent to tubules 14. Temperature controlling means 21 is connected to incubation chamber 19 to heat and/or cool incubation chamber 19. It is to be appreciated that the temperature of tubules 14 may be controlled directly, such as with a temperature sensor detecting the temperature of tubules 14 and maintaining a desired setpoint temperature. Alternatively, the temperature of the tubules could be controlled indirectly by sensing and controlling the temperature of incubation chamber 19. Temperature controlling means 21 may include a heating element and may also include a cooling device. Other suitable temperature controlling means will become readily apparent to those skilled in the art given the benefit of this disclosure.

Turning now to Fig. 3, tube sealing apparatus 12 will be shown in greater detail. Tube sealing apparatus 12 comprises upper, or first sealing head 24 and lower, or second sealing head 26. Upper sealing head 24 has conductors 36 extending from an upper

surface 38 to a lower sealing surface 40. Lower sealing head 26 also has conductors 36 extending from an upper sealing surface 42 to a lower surface 44. Conductors 36 are connected by cables 45 to a power source 46 which creates a radio frequency (RF) electrical field between the conductors 36 of upper sealing head 24 and lower sealing head 26 which heat seals tube 8. Conductors 36 are preferably formed of a material having high electrical and heat conductivity. Suitable materials for conductor 36 are, for example, metals such as copper. Other suitable materials for the sealing heads will become readily apparent to those skilled in the art, given the benefit of this disclosure. Upper sealing head 24 and lower sealing head 26 are preferably formed of a substantially rigid insulating material having high heat conductivity. Suitable materials for the sealing heads include plastics such as nylon. Other suitable materials for the sealing heads will become readily apparent to those skilled in the art, given the benefit of this disclosure. Resilient pads 48 are preferably located at the outer edges of lower sealing surface 40 and upper sealing surface 42. Resilient pads 48 may be formed of rubber, silicone rubbers, teflon, fluoropolymers, or any other suitable resilient material. In certain preferred embodiments, a central bar 50 may be located between a pair of conductors 36. As seen in Fig. 4, both upper sealing head 24 and lower sealing head 26 have a central bar 50. It is to be appreciated that in certain preferred embodiments, only upper sealing head 24 may have a central bar 50, while lower sealing head 26 has a single conductor 36.

As seen in Fig. 4, tube 8, containing fluid sample 51, e.g., whole blood, is passed between upper sealing head 24 and lower sealing head 26. The volume of a portion of tube 8, or tubule 14, is adjusted by compressing upper bar 9 and lower bar 11 together about tubule 14. In certain preferred embodiments, the volume of tubule 14 is approximately 20 μ l. The tubule 14 may contain, for example, approximately 5 μ l of whole blood or approximately 15 μ l of plasma. Upper and lower sealing heads 24, 26 are then squeezed together under pressure, compressing a portion of tube 8 and pushing fluid sample 51 outwardly in the direction of arrows A. As sealing heads 24, 26 compress tube 8, a sample free zone 52 is created, that is, a zone is created within tube 8 which is substantially free of any fluid sample 51. The pressure must be sufficient to squeeze fluid sample 51 out of sample free zone 52 as well as sufficient to prevent pressure in tubule 14 from forcing fluid sample 51 back into sample free zone 52, especially during

sealing. The required pressure forcing sealing heads 24, 26 together is dependent on the material of tube 8, as well as its diameter and wall thickness. In certain preferred embodiments, fluid sample 51 is approximately 2mm away from conductors 36 which provide the sealing of tubule 14.

5 As seen in Fig. 5, central bar 50 is then raised, releasing the pressure in a central area of sample free zone 52 and creating an injection channel 54 which is also free of fluid sample 51. Power source 46 then supplies RF power through cables 45 to conductors 36 which seals tube 8 forming seal 13. In certain preferred embodiments, the frequency of the RF power supplied is approximately 40 MHz. The RF power is
10 supplied for a time period typically less than one second. The power and duration of the supplied RF energy may vary based on the size of tube 8 and the material of which it is constructed. Upper sealing member 24 is then raised, tube 8 is advanced to the left as seen in Fig. 4, and tube 8 is sealed again, forming a tubule 14 between seals 13. By creating sample free zone 52, fluid sample 51 is kept a safe distance from conductors 36
15 when the RF power is applied, thereby reducing negative effects on fluid sample 51 from the RF power and the heat it generates.

 In the embodiment illustrated in Fig. 4, lower sealing head 26 is fixed and upper sealing head 24 moves downwardly in the direction of arrows B toward lower sealing head 26. In other preferred embodiments, upper sealing head 24 may be fixed with lower
20 sealing head 26 moving toward upper sealing head 24, or both upper and lower sealing heads 24, 26 may move toward one another.

 In the embodiment illustrated in Figs. 4, 5, lower sealing surface 40 and upper sealing surface 42 have a substantially convex profile. Thus when sealing heads 24, 26 are brought together, tube 8 is compressed a maximum amount in the central area of
25 heads 24, 26, that is, in sample free zone 52, and compresses to a lesser extent outside of sample free zone 52.

 In certain preferred embodiments, as seen in Fig. 6, central bar 50 has an L shaped, or inverted L shaped profile. In the embodiment illustrated, central bar 50 of first sealing head 24 has an inverted L shape and central bar 50 of second sealing head 26 has
30 an L shape. Conductor 36 is formed of conductor element 36A and conductor element 36B, spaced apart by central bar 50. Conductor element 36A extends along the long leg

of central bar 50 and terminates at its short leg. Conductor element 36 B extends along the length of the long leg of central bar 50. Lines W represent the width of a tube 8 which is sealed by sealing heads 24, 26. It can be seen that the sealing heads extend beyond the edge of the tube such that the seal, when formed, extends across the entire width of the tube. When the RF power is applied, as seen in Fig. 7, seal 13, comprising first portion 13A and second portion 13B is formed only in the areas where conductor elements 36A, 36B lie, creating L shaped injection channel 54 which is capable of being in fluid communication with tubule 14. However, tension in the area of seal 13 prevents fluid sample 51 from entering injection channel 54. Reagent is added to injection channel 54 through needle 15, seen in Fig. 2 and described in greater detail below. The amount of reagent added to tubule 14 is preferably approximately 1-15 μ l depending on the assay being performed. By maintaining injection channel 54 free of fluid sample 51, any leakage from tubule 14 is prevented when a needle punctures the side wall of the tube to inject reagent into the tubule through injection channel 54. In certain preferred embodiments, the needle puncture in injection channel 54 has been found to be able to withstand pressure of up to approximately 3 atm. without leaking.

The specific configuration of injection channel 54 is not critical, except that it must be sufficiently large to receive the reagent injection needle. Also, in accordance with a highly advantageous aspect, indicated above, it is sufficiently small so as to be self-sealing. That is, the bore, length, and configuration of the injection channel are such that the passageway is normally substantially devoid of fluid sample. Given the benefit of this disclosure of the general concept and principles of the injection channel, it will be within the ability of those skilled in the art to select suitable dimensions and configurations for the injection channel, taking into account the size, wall thickness and resiliency of the flexible plastic tube. Thus, while the injection channel is normally closed or collapsed so as to be devoid of fluid sample, it still provides fluid communication into the main fluid chamber within the tubule. That is, reagent or other fluid injected into the injection channel under suitable injection pressure passes through the injection channel to the main chamber. Once the injection needle is withdrawn, however, the injection channel returns to its closed or collapsed condition such that

leakage does not occur during normal operating conditions through the hole in the wall formed at the end of the passageway by the needle.

In another preferred embodiment, seen in Fig. 8, central bar 50' has a T shaped profile with conductor 36 comprising conductor elements 36B, 36C, and 36D. In yet another preferred embodiment, seen in Fig. 9, conductor 36 is formed of a single conductor element 36E. In this embodiment, a single lateral seal 13 is formed across tube 8. Alternatively, tube 8 or tube sealing apparatus 12 can be repositioned after a first seal 13A is formed, creating a second seal 13B as seen in Fig. 7 to form an injection channel 54.

As shown in Fig. 2, needle 15 is inserted into tubule 14, preferably into injection channel 54, to add reagent to fluid sample 51 into tubule 14. In a preferred embodiment, the reagent is added through injection channel 54 prior to upper and lower sealing heads 24, 26 being fully released. In other preferred embodiments, the reagent is added just prior to the tubule 14 entering flow control device 18, so that the inspection of the sample is done soon after the reagent has been added. Reagent can be drawn from reservoir 16 by releasing upper and lower bars 9, 11, creating vacuum pressure within tubule 14 and drawing reagent into tubule 14. Central bar 50 may then be depressed, forcing any reagent remaining in injection channel 54 into tubule 14.

As seen in Fig. 24, tube sealing apparatus 55 may comprise a pair of rotatable wheels 57 having a plurality of circumferentially disposed teeth 59. The outer surface of each tooth 59 is substantially planar or curvoplanar. A conductor 61 operably connected to power source 46 by cables (not shown) is located within each tooth 59. The surface 63 of wheels 57 extending between teeth 59 is substantially concave. Wheels 57 rotate in opposite directions to progress tube 8 through tube sealing apparatus 55, with surfaces 63 preferably being configured to compress each portion of tube 8 between the seals to a desired volume. As an opposed pair of teeth 59 meet, radio frequency energy or heat, etc. is transmitted through conductors 61, forming seal 13 in the manner described above.

In other preferred embodiments, sealing of the chamber or tube 8 can be accomplished by other suitable sealing means. Examples of other sealing means include, for example, mechanical clamps, a fold lock, ultrasound fusion, and direct application of heat to the tube. Tube 8 may, in certain preferred embodiments, be a heat shrinkable tube

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and the tube sealing apparatus may be a device for applying focused heat to each of the seal locations along the length of the tube.

In another preferred embodiment, shown in Fig. 10, reagent reservoir 16 may be contained in a reagent cartridge 60 having housing 62. Bladder 64 is disposed within housing 62 and is secured to an inner wall of housing 62 by ring 66. Reagent is thus contained within bladder 64. Needle 15 extends from housing 62 and is preferably covered by resilient cover 68. Vent 70 is provided in an upper surface of housing 62 and a filler plug 71 is provided in housing 62 for adding reagent. In certain preferred embodiments, magnetic stirrer 72 is positioned in reservoir 16 on a bottom surface of housing 62. A magnetic field generator 74 positioned outside housing 62 creates rotation of magnetic stirrer 72, mixing the reagent, e.g. a cell suspension, prior to injection into tubule 14. The reagent may also be mixed by other means such as shaking. Tube 76 of piezoelectric material surrounds needle 15 and serves as a drop generator as described more fully in U.S. Patent No. 4,329,698. Multiple reservoirs 16 of reagent may be contained within reagent cartridge 60, allowing different reagents to be added to different tubules 14 as they pass through testing machine 2.

testing machine 2.

One preferred embodiment is shown in Fig. 22. In the illustrated embodiment, reagent cartridge 60 contains 12 reservoirs of different reagents, each reservoir having its own needle 15, and each reagent being used for a specific test. A bar code label 73 on reagent cartridge 60 provides information to identify particular reagents contained therein and test procedure necessary for programming the sample test system. Tubules 14 are moved in an axial direction, preferably in step-wise fashion, past reagent cartridge 60. Reagent cartridge 60 is movable in a direction transverse to a longitudinal axis of the tubules in order to position the proper needle 15 corresponding to a desired reagent, at the injection channel of each tubule in turn. Once reagent cartridge 60 is properly positioned, needle 15 is injected into tubule 14 to inject the desired reagent.

Another preferred embodiment is shown in Fig. 11, where reagent cartridge 60A has housing 62A with an adapter 78 located on an upper surface of housing 62A to receive air nozzle 80. In use, as seen in Fig. 12, needle 15 extends through resilient cover 68 and penetrates the wall of tubule 14. In the preferred embodiment illustrated, needle

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15 extends into injection channel 54. Air pressure is introduced onto bladder 64 through air nozzle 80, causing reagent from reservoir 16 to be forced into tubule 14. In the embodiment illustrated, needle 15 is fixed with respect to reagent cartridge 60A, and the entire reagent cartridge 60A is moved vertically by actuator 49 (seen in Fig. 1) in order to inject needle 15 into tubule 14. In other preferred embodiments, needle 15 may be independent of reagent cartridge 60A such that only needle 15 moves in order to inject reagent into tubule 14.

Another preferred embodiment is shown in Fig. 13, where reagent cartridge 60B comprises housing 62B having piston 82 disposed therein above reservoir 16 containing reagent. A pair of resilient annular rings 84 are positioned between piston 82 and an inner wall of housing 62B, providing a seal between piston 82 and housing 62B. Shaft 86 is in contact with the upper surface of piston 82 and pressure is introduced into reservoir 16 as shaft 86 causes piston 82 to be lowered. The pressure in reservoir 16 forces reagent through needle 15 into tubule 14.

Yet another embodiment is shown in Fig. 14, where reagent cartridge 60C comprises housing 62C having resilient sac 88 forming reservoir 16 therein. Shaft 86 engages an outer surface of sac 88, introducing pressure into reservoir 16 in order to force reagent through needle 15.

In other preferred embodiments, multiple reagent cartridges, each having a single reservoir or reagent, may be chained together with a flexible connector such that a large number of reagent cartridges may be connected together. The connected reagent cartridges can then, for example, be rolled up to facilitate storage and delivery.

In certain preferred embodiments, a reagent cartridge with multiple needles in fluid communication with a single, or corresponding multiple reservoirs, may be used to inject, or deposit reagent simultaneously, or sequentially, into multiple different tubules. The reagent cartridge may also be used to inject or deposit reagent into other chambers or containers. For example, a reagent cartridge with multiple needles in fluid communication with a single, or corresponding multiple reservoirs, can be used to simultaneously, or sequentially, inject or deposit reagent into a plurality of containers, such as the recesses of a ninety-six well microplate.

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Flow control device 18 is seen in Fig. 15 and comprises transparent base member 90 upon which tubule 14 is placed. Transparent central plunger 92 is positioned above tubule 14 and lowered onto tubule 14 such that tubule 14 is sandwiched between central plunger 92 and base member 90, creating first and second reservoir zones 94, 96 in tubule 14, with a narrow flow passage 98 extending therebetween through which a thin layer of sample flows. A first outer plunger 100 is positioned above first reservoir zone 94 and a second outer plunger 102 is positioned above second reservoir zone 96. First and second outer plungers 100, 102 are alternately raised and lowered (shown by arrows D), engaging and disengaging tubule 14, creating a flow of fluid sample 51 back and forth through narrow flow passage 98. By sensing the pressure needed to cause the flow of fluid sample 51 through passage 98, the specific molecular binding strength between cells or particles in the sample can be determined. The number of particles or cells in the sample can be counted, and cell properties such as size and light intensity can be measured. In a preferred embodiment, the height of, or gap created by, flow passage 98 is approximately 10 μ m to 100 μ m, depending on the assay performed. Through such a narrow passageway, the flow of fluid sample 51 can be analyzed by computerized microscopic inspection system 20. Light from light source 28, shown by arrows C, is projected through central plunger 92 and passage 98. Images of fluid sample 51 as it flows through passage 98 are captured by camera with microscopic lens 30 which then transfers the images through frame grabber 32 to computer 34 (seen in Fig. 2) for analysis through known signal processing algorithms. It is to be appreciated that operation of flow control device 18 may, in certain preferred embodiments, include portions of time where no flow is generated through passage 98, and camera 30 may capture images of fluid sample 51 during these non-flow periods. Camera 30 is, in certain preferred embodiments, a charged-coupled device (CCD) camera. Cell interaction kinetics can be analyzed by computer 34 by monitoring cell motion and/or location as well as optical properties of the cells such as light scattering.

Cell-cell interaction occurs in tubule 14 when any of certain known reagents are added to a blood sample. Molecular interactions occur when the reagent is added to the sample. Aggregates may be formed in the sample, and the size and distribution of the aggregates varies depending on the type of reagent added to fluid sample 51, the shear

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flow of the sample, and the time period elapsed after injection of the reagent. In a known fashion, the size and quantity of aggregates passing through flow passage 98 allows various types of screening or analysis to be performed on fluid sample 51. For example, immunodiagnosis such as blood typing, antibody screening and infectious disease testing can be performed using the present invention by selecting suitable known reagents to be injected into one or more tubules. Specifically, blood forward typing can be performed by adding a related antibody as the reagent to fluid sample 51 comprising whole blood. Blood reverse typing can be performed by adding a cell suspension as the reagent to fluid sample 51 comprising plasma. Blood reverse typing can also be performed by adding cell suspension as the reagent to fluid sample 51 comprising whole blood. Hematology tests for blood components such as red and white blood cell counts, coagulation and aggregation time testing, and platelet function tests can be performed as well. The reagent may comprise anti-analyte coated beads in order to detect specific analyte in the sample. Other tests such as nucleic acid amplification and DNA analysis may also be performed in the manner disclosed here. Blood chemistry analysis can detect, for example, sugar levels, cholesterol levels, etc. Drug compound testing can also be performed using the present invention. Other testing which can be performed using the present invention will become readily apparent to those skilled in the art, given the benefit of this disclosure.

The present invention provides many advantages. A testing machine can be used cost effectively for many different tests and groups of tests. The testing machine has high throughput and low complexity for ease of operation. Bio-safety is increased due to reduced handling of samples such as blood.

Computer 34, in certain preferred embodiments, may be operably connected to tube advancing system 3, tube sealing apparatus 12, flow control device 18, incubation chamber 19, centrifuge 35, and inspection system 20 by cables (not shown). Computer 34 can provide control and coordination of the operating parameters of the components of testing machine 2 in a known fashion, and further description of the control of the components of testing machine 2 need not be provided here.

In another preferred embodiment, shown in Fig. 16, flow control device 18A comprises transparent cylindrical plunger 92A having a longitudinal axis L and a beveled

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surface 104 formed on lower surface 106 of plunger 92A. A reservoir 94A is formed beneath beveled surface 104 and passage 98A is formed beneath lower surface 106. As plunger 92A is rotated about longitudinal axis L, flow through passage 98A can be observed in the same manner described above.

5 Another preferred embodiment is shown in Fig. 17, where flow control device 18B comprises transparent plunger 92B having first and second beveled surfaces 108, 110 formed on a lower surface thereof. First and second reservoirs 94B, 96B are formed beneath beveled surfaces 108, 110, respectively, with narrow passage 98B extending therebetween. As plunger 92B is rocked back and forth, fluid sample 51 passes back and forth from first reservoir 94B to second reservoir 96B through passage 98B. The flow of fluid sample 51 is observed by camera 30 as described above.

10 Yet another embodiment is shown in Fig. 18, where flow control device 18C comprises transparent plunger 92C whose lower surface 112 has an arcuate profile. The arcuate profile of lower surface 112 creates a narrow flow passage 98C extending between a first reservoir 94C and a second reservoir 96C. Plunger 92C is rolled back and forth, forcing fluid sample 51 back and forth from first reservoir 94C to second reservoir 96C through flow passage 98C. The flow of fluid sample 51 through flow passage 98C is observed by camera 30 as described above.

15 In certain preferred embodiments, as seen in Fig. 19, a first electrode 120 and a second electrode 122 are inserted into tubule 14 and are connected by cables 124 to voltage source 126 which creates a voltage difference between first and second electrodes 120, 122. Red blood cells in fluid sample 51 within tubule 14 are negatively charged so that by electrophoresis they are attracted to the positively charged electrode 122. An electrochemiluminescent reagent is added to tubule 14 by reagent cartridge 60 or other suitable means, creating an electrochemiluminescent reaction near the surface of electrode 122 which causes a particular light to be emitted (shown by arrows E) from electrode 122 based on the type of reagent added to tubule 14. Sensor 128 receives the transmitted light and generates a corresponding electrical signal which is sent to computer 34 for analysis, display, recording, etc. In other preferred embodiments, a current is passed by first and second electrodes 120, 122 through the sample. In this embodiment,

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certain electrochemical properties of the sample can be measured by analyzing the voltage difference between the first and second electrodes 120, 122.

Another preferred embodiment is shown in Fig. 20. First and second electrodes 130, 132 are inserted into tubule 14. Second electrode 132 is a fiberoptic sensor. As described above with respect to Fig. 19, an electrochemiluminescent reaction occurs near the surface of electrode 132 causing light to be generated. The light travels through fiberoptic electrode 132 to a fiber optic sensor, or reader 134 which captures and interprets the information provided by the type of light generated. Second electrode 132 preferably has a diameter between approximately 0.4mm and 1mm. Second electrode 132 is formed of a material or is coated with a material suitable for providing sufficient conductivity.

In certain preferred embodiments, a coating may be deposited on tubule 14 to increase visibility through the wall of tubule 14. As seen in Fig. 21, a coating material 140 is transferred through conduit 142 from coating supply 144 and deposited on the outer surface of tubule 14. If the walls of tubule 14 are translucent, the addition of coating 140 to the outer surface of tubule 14 can make the walls of tubule 14 substantially transparent, increasing the effectiveness of viewing the flow of fluid sample 51 through flow passage 98. Coating 140 preferably has the same optical refractive index as that of the walls of tubule 14. Suitable materials for coating 140 are dependent on the material of tubule 14 and include, for example, oil.

Suitable methods for filling a tube with a sample will be apparent to those skilled in the art, given the benefit of this disclosure. Exemplary methods include injecting sample fluid into one end of a tube or drawing sample into a tube by creating a vacuum in the tube. A suitable tube 150 is shown in Fig. 23, having a self-sealing head 152 at a first end thereof for needle penetration. Tube 150 may have a label 154 to assist in identifying the source of the sample, e.g., a patient's name when the sample is blood. Label 154 may be, e.g., a bar code label. Tube 150 is inserted into a tube-like drawing device 156 through an aperture 158 at a first end of drawing device 156. To draw a sample into tube 150, the tube-like drawing device 156 is plugged into a needle holder commonly used for drawing blood into a vacuum tube, and slide handle 160 is moved downwardly along drawing device 156. A pair of opposed rollers (not shown) within

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drawing device 156 and operably connected to slide handle 160 compress a portion of, and roll downwardly along, tube 150, pumping or drawing a sample of blood into tube 150.

5 In some cases a multiple stage reaction within a segment of a chamber may be desired. In one embodiment, the reagent is injected through an injection channel in the segment, reacted with the contents therein, and then, later, a second reagent is added and reacted with the contents. In an alternative preferred embodiment, the segment may be formed with a pressure gate, separating the volume of the segment into two
10 compartments between which there is fluid communication only at pressure levels achieved by application of external pressure. Pressure for moving sample material from one compartment into an adjacent compartment may be applied, e.g., by hand or by automatic mechanical pressure devices such as those shown in Figs. 2, 4, 5 and adapted to apply pressure to a single compartment.

One preferred example is shown in Fig. 25, where a segment or tubule 168 is
15 separated by a seal 170 into first compartment 172 and second compartment 174. Seal 170 is formed in a manner as described above with respect to seal 13. Seal 170 forms a pressure gate 176, which, under normal operating conditions, provides a fluid-tight seal between first and second sub-segments or compartments 172, 174. In a preferred embodiment, pressure gate 176 opens upon application of pressure greater than a certain
20 value, for example, approximately 2 atm. When external pressure is applied to one of the compartments, pressure gate 176 opens, allowing fluid to flow from the high pressure compartment to the low pressure compartment. One preferred application is in a two stage antibody screening wherein first compartment 172 of tubule 168 is pre-filled with plasma. A first reagent is injected through injection channel 54 into second compartment
25 174. External pressure is then applied to second compartment 174, forcing the first reagent into first compartment 172. A second reagent is added to second compartment 174 through injection channel 54. Tubule 168 is then conveyed by suitable means to incubation chamber 19 for a predetermined time period of incubation. Tubule 168 is then conveyed by suitable means to centrifuge 35 where tubule 168 is spun such that the cells
30 of the first reagent accumulate proximate pressure gate 176. In certain preferred embodiments, the second reagent may be added after tubule 168 has been incubated in

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incubation chamber 19 or spun in centrifuge 35. External pressure is applied to first compartment 172 such that cells of the first reagent are passed to second compartment 174. Tubule 168 is then conveyed to flow control device 18 and inspected by inspection system 20 in the manner described above.

- 5 In light of the foregoing disclosure of the invention and description of the preferred embodiments, those skilled in this area of technology will readily understand that various modifications and adaptations can be made without departing from the true scope and spirit of the invention. All such modifications and adaptations are intended to be covered by the following claims.

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CLAIMS:

1. A sample testing system comprising, in combination:
 - a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of a chamber;
 - a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move the reagent injector cartridge to inject a quantity of reagent into a segment of the chamber; and
 - a sensor to generate an output signal corresponding to a condition of a fluid sample material within the segment of the chamber.
2. The sample testing system of claim 1, further comprising an output device responsive to the output signal.
3. The sample testing system of claim 1 or 2, wherein the sensor comprises a light detector to receive light responsive to a fluid sample in the segment of the chamber and to generate the output signal corresponding to the condition of the fluid sample material within the segment of the chamber.
4. The sample testing system of any one of claims 1 to 3, further comprising a second sensor to receive information from a label of the reagent injector cartridge.
5. The sample testing system according to any one of claims 1 to 4, further comprising a programmable control system coupled with the chamber sealing apparatus, the reagent injector cartridge actuator, and the sensor.
6. The sample testing system according to any one of claims 1 to 5, wherein the chamber is a flexible plastic tube containing the fluid sample material.
7. The sample testing system according to claim 6, wherein the chamber sealing apparatus has a tube compression and sealing member to laterally seal the flexible plastic

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tube and thereby form a fluid-tight tubule containing a portion of the fluid sample material between axial spaced lateral seals.

8. The sample testing system according to claim 7, wherein the chamber sealing apparatus comprises:
 - a first sealing head comprising the tube compression and sealing member; and
 - a second sealing head, at least one of the first and second sealing heads being movable toward the other of the sealing heads to compress a section of flexible plastic tube positioned between the first and second sealing heads to create a sample-free zone in the tube, wherein the tube compression and sealing member is operatively connected to a power source to heat a sealing zone of tube located in the sample-free zone to form a fluid-tight lateral seal in the tube.
9. The sample testing system according to claim 8, wherein the first sealing head and the second head each has at least one electrically conductive portion and at least one of the sealing heads has a substantially convex profile.
10. The sample testing system according to claim 8 or 9, further comprising a radio frequency power generator to transmit radio frequency energy between the first and second sealing heads through the sealing zone to form a lateral seal in the tube.
11. The sample testing system according to any one of claims 8 to 10, wherein the chamber sealing apparatus further comprises a compression apparatus to partially compress a portion of a tube to create a tubule of a predetermined volume.
12. The sample testing system according to any one of claims 6 to 11, further comprising an incubation chamber to retain a segment of a tubule for a predetermined period of time.

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13. The sample testing system according to claim 12, wherein the incubation chamber includes a temperature control device to control the temperature of a segment of a tubule in the incubation chamber.
14. The sample testing system according to any one of claims 6 to 13, wherein the sensor detects properties of particles within a fluid sample in a segment of the tubule.
15. The sample testing system according to any one of claims 6 to 14, further comprising a light source to project light through a segment of the tubule.
16. The sample testing system according to claim 15, wherein the sensor comprises a camera having a microscopic lens to capture images of a fluid sample in an inspection zone of a segment of the tubule as the light source is projecting light through the inspection zone; and
further comprising a computer operably connected to the camera to analyze the images of the fluid sample.
17. The sample testing system according to any one of claims 6 to 16, further comprising:
a pair of electrodes adapted to have a predetermined voltage difference, and
an electrode actuator to insert the pair of electrodes into a segment of the tubule, wherein the sensor is responsive to electrophoretic light emitted from within a segment of the tubule.
18. The sample testing system according to claim 17, wherein one of the electrodes comprises a fiber optic conductor.
19. The sample testing system according to any one of claims 6 to 18, further comprising a segment of a tubule and a coating on an outside surface of the segment to increase the transmission of light through the segment.

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20. The sample testing system according to claim 19, wherein the coating has a refractive index substantially the same as a refractive index of the segment.
21. The sample testing system according to any one of claims 6 to 20, further comprising a segment advancing device to transport a length of tubule within the sample testing system.
22. The sample testing system according to claim 21, wherein the segment advancing device comprises a pair of rotatable wheels spaced apart from one another to receive the tubule therebetween, at least one of the wheels being a driven wheel operatively connected to an output member of an electric motor.
23. A sample testing system comprising, in combination:
a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of a chamber;
a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move the reagent injector cartridge to inject a quantity of reagent into a segment of the chamber;
a flow control device comprising at least one contact member movable into contact with the segment of the chamber to effect mechanically induced fluid flow within a flow passageway in the segment of the chamber; and
a sensor to generate an output signal corresponding to a condition of a fluid sample material within the segment of the chamber.
24. The sample testing system of claim 23, further comprising an output device responsive to the output signal.
25. The sample testing system of claim 23 or 24, wherein the sensor comprises a light detector to receive light responsive to the fluid sample in the segment of the chamber and to generate the output signal corresponding to the condition of the fluid sample material

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within the segment of the chamber.

26. The sample testing system of any one of claims 23 to 25, further comprising a second sensor to receive information from a label of the reagent injector cartridge.

27. The sample testing system according to any one of claims 23 to 26, further comprising a programmable control system coupled with the chamber sealing apparatus, the reagent injector cartridge actuator, the flow control device, and the sensor.

28. The sample testing system according to any one of claims 23 to 26, wherein the chamber is a flexible plastic tube containing the fluid sample material and the chamber sealing apparatus has a tube compression and sealing member to laterally seal the flexible plastic tube and thereby form a fluid-tight tubule containing a portion of the fluid sample material between axial spaced lateral seals.

29. The sample testing system according to claim 28, wherein the chamber sealing apparatus comprises:

a first sealing head comprising the tube compression and sealing member; and
a second sealing head, at least one of the first and second sealing heads being movable toward the other of the sealing heads to compress a section of flexible plastic tube positioned between the first and second sealing heads to create a sample-free zone in the tube, wherein the tube compression and sealing member is operatively connected to a power source to heat a sealing zone of tube located in the sample-free zone to form a fluid-tight lateral seal in the tube.

30. The sample testing system according to claim 29, wherein the first sealing head and the second head each has at least one electrically conductive portion and at least one of the sealing heads has a substantially convex profile.

31. The sample testing system according to claim 29 or 30, further comprising a radio frequency power generator to transmit radio frequency energy between the first and

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second sealing heads through the sealing zone to form a lateral seal in the tube.

32. The sample testing system according to any one of claims 29 to 31, wherein the chamber sealing apparatus further comprises a compression apparatus to partially compress a portion of a tube to create a tubule of a predetermined volume.

33. The sample testing system according to any one of claims 23 to 32, wherein the flow control device further comprises a base member for supporting a segment of a chamber, the contact member being movable toward the base member to partially compress at least one portion of a segment of a chamber against the base member forming a fluid passageway in the segment and further movable in repeating fashion to alternately compress and decompress at least one portion of the segment to induce fluid flow within the fluid passageway.

34. The sample testing system according to claim 33, wherein the sensor detects light passing through a segment of a chamber as the flow control device is operating to induce fluid flow in that segment.

35. The sample testing system according to any one of claims 23 to 34, further comprising an incubation chamber to retain a segment of a chamber for a predetermined period of time.

36. The sample testing system according to claim 35, wherein the incubation chamber includes a temperature control device to control the temperature of a segment of a chamber in the incubation chamber.

37. The sample testing system according to any one of claims 23 to 36, wherein the sensor detects properties of particles within a fluid sample in a segment of a chamber.

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38. The sample testing system according to any one of claims 23 to 37, further comprising a light source to project light through a segment of a chamber.
39. The sample testing system according to claim 38, wherein the sensor comprises a camera having a microscopic lens to capture images of a fluid sample in an inspection zone of a segment of chamber as the light source is projecting light through the inspection zone; and
further comprising a computer operably connected to the camera to analyze the images of the fluid sample.
40. The sample testing system according to any one of claims 23 to 39, further comprising
a pair of electrodes adapted to have a predetermined voltage difference, and an electrode actuator to insert the pair of electrodes into a segment of a chamber, wherein the sensor is responsive to electrophoretic light emitted from within a segment of a chamber.
41. The sample testing system according to claim 40, wherein one of the electrodes comprises a fiber optic conductor.
42. The sample testing system according to any one of claims 23 to 41, further comprising a segment of a chamber and a coating on an outside surface of the segment to increase the transmission of light through the segment.
43. The sample testing system according to claim 42, wherein the coating has a refractive index substantially the same as a refractive index of the segment.
44. The sample testing system according to any one of claims 23 to 43, further comprising a segment advancing device to transport a length of tube within the sample testing system.

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45. The sample testing system according to claim 44, wherein the segment advancing device comprises a pair of rotatable wheels spaced apart from one another to receive the tube therebetween, at least one of the wheels being a driven wheel operatively connected to an output member of an electric motor.
46. A sample testing system comprising, in combination:
tube sealing apparatus having a tube compression and sealing member to laterally seal a flexible plastic tube containing a fluid sample material, whereby a fluid-tight tubule containing a portion of the fluid sample material can be formed between axially spaced lateral seals; the tube sealing apparatus comprising a first sealing head comprising the tube compression and sealing member; and a second sealing head, at least one of the first and second sealing heads being movable toward the other of the sealing heads to compress a section of flexible plastic tube positioned between the first and second sealing heads to create a sample-free zone in the tube, wherein the tube compression and sealing member is operatively connected to a power source to heat a sealing zone of the tube located in the sample-free zone to form a fluid-tight lateral seal in the tube;
a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a fluid-tight tubule; and
a sensor to generate an output signal corresponding to a condition of a fluid sample material within a fluid-tight tubule.
47. The sample testing system according to claim 46, wherein the first sealing head and the second head each has at least one electrically conductive portion and at least one of the sealing heads has a substantially convex profile.
48. The sample testing system according to claim 46 or 47, further comprising a radio frequency power generator to transmit radio frequency energy between the first and second sealing heads through the sealing zone to form a lateral seal in the tube.
49. The sample testing system according to any one of claims 46 to 48, wherein the

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chamber sealing apparatus further comprises a compression apparatus to partially compress a portion of a tube to create a tubule of a predetermined volume.

50. A sample testing system comprising, in combination:
- a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of the chamber;
 - a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber;
 - a sensor to generate an output signal corresponding to a condition of a fluid sample material within a segment of a chamber;
 - a pair of electrodes adapted to have a predetermined voltage difference, and
 - an electrode actuator to insert the pair of electrodes into a segment of a chamber,
- wherein the sensor is responsive to electrophoretic light emitted from within a segment of a chamber.

51. The sample testing system according to claim 50, wherein one of the electrodes comprises a fiber optic conductor.

52. A sample testing system comprising, in combination:
- a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of the chamber;
 - a segment of the chamber including a coating on an outside surface of the segment to increase the transmission of light through the segment;
 - a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber; and
 - a sensor to generate an output signal corresponding to a condition of a fluid sample material within a segment of a chamber.

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53. The sample testing system according to claim 52, wherein the coating has a refractive index substantially the same as a refractive index of the segment.
54. A sample testing system comprising, in combination:
a chamber sealing apparatus to form a plurality of seals defining a plurality of fluid-tight segments of the chamber;
a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a segment of a chamber;
an incubation chamber to retain a segment of a chamber for a predetermined period of time; and
a sensor to generate an output signal corresponding to a condition of a fluid sample material within a segment of a chamber.
55. The sample testing system according to claim 54, wherein the incubation chamber includes a temperature control device to control the temperature of a segment of a chamber in the incubation chamber.
56. A sample testing system comprising, in combination:
tube sealing apparatus having a tube compression and sealing member to laterally seal a flexible plastic tube containing a fluid sample material, whereby a fluid-tight tubule containing a portion of the fluid sample material can be formed between axially spaced lateral seals;
a reagent injector cartridge actuator adapted to receive a reagent injector cartridge having at least one needle in fluid communication with a reagent reservoir, and to move a reagent injector cartridge to inject a quantity of reagent into a tubule;
a flow control device comprising a contact member movable into contact with a tubule to effect mechanically induced fluid flow within a fluid passageway in the tubule;
and

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an inspection system having a light detector to receive light passed through a tubule and to generate an output signal corresponding to a condition of the fluid sample material within a tubule.

57. The sample testing system according to claim 56, further comprising an output device responsive to the output signal.

58. The sample testing system according to claim 56 or 57, further comprising a programmable control system coupled with the tube sealing apparatus, the reagent injector cartridge actuator, the flow control device and the inspection system.

59. The sample testing system according to any one of claims 56 to 58, wherein the tube sealing apparatus comprises:

a first sealing head comprising the tube compression and sealing member; and
a second sealing head, at least one of the first and second sealing heads being movable toward the other of the sealing heads to compress a section of flexible plastic tube positioned between the first and second sealing heads to create a sample-free zone in the tube, wherein the tube compression and sealing member is operatively connected to a power source to heat a sealing zone of tube located in the sample-free zone to form a fluid-tight lateral seal in the tube.

60. The sample testing system according to claim 59, wherein the first sealing head has a substantially convex profile and an electrically conductive central portion, and the second sealing head has a substantially convex profile and an electrically conductive central portion.

61. The sample testing system according to claim 59 or 60, further comprising a radio frequency power generator to transmit radio frequency energy between the first and second sealing heads through the sealing zone to form a lateral seal in the tube.

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62. The sample testing system according to any one of claims 59 to 61, wherein the tube sealing apparatus further comprises a compression apparatus to partially compress a portion of a tube to create a tubule of a predetermined volume.

63. The sample testing system according to any one of claims 56 to 62, wherein the flow control device further comprises a base member for supporting a tubule, the contact member being movable toward the base member to partially compress at least one portion of a tubule against the base member forming a fluid passageway in the tubule and further movable in repeating fashion to alternately compress and decompress at least one portion of the tubule to induce fluid flow within the fluid passageway.

64. The sample testing system according to any one of claims 56 to 63, further comprising an incubation chamber to retain a tubule for a predetermined period of time.

65. The sample testing system according to claim 64, wherein the incubation chamber includes a temperature control device to control the temperature of a tubule in the incubation chamber.

66. The sample testing system according to any one of claims 56 to 65, wherein the light detector of the inspection system detects light passing through a tubule as the flow control device is operating to induce fluid flow in that tubule.

67. The sample testing system according to any one of claims 56 to 66, wherein the light detector of the inspection system detects properties of particles within a fluid sample in a tubule.

68. The sample testing system according to any one of claims 56 to 67, wherein the inspection system further comprises a light source to project light through a tubule.

69. The sample testing system according to claim 68, wherein the inspection system further comprises

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a camera having a microscopic lens to capture images of a fluid sample in an inspection zone of a tubule as the light source is projecting light through the inspection zone; and

a computer operably connected to the camera to analyze the images of the fluid sample.

70. The sample testing system according to any one of claims 56 to 69, further comprising

a pair of electrodes adapted to have a predetermined voltage difference; and

an electrode actuator to insert the pair of electrodes into a tubule, wherein the inspection system light detector is responsive to electrophoretic light emitted from within a tubule.

71. The sample testing system according to claim 70, wherein one of the electrodes comprises a fiber optic conductor.

72. The sample testing system according to any one of claims 56 to 71, further comprising a tubule and a coating on an outside surface of the tubule to increase the transmission of light through the tubule.

73. The sample testing system according to claim 72, wherein the coating has a refractive index substantially the same as a refractive index of the tubule.

74. The sample testing system according to any one of claims 56 to 73, further comprising a tube advancing device to transport a length of tube within the sample testing system.

75. The sample testing system according to claim 74, wherein the tube advancing device comprises a pair of rotatable wheels spaced apart from one another to receive the tube therebetween, at least one of the wheels being a driven wheel operatively connected to an output member of an electric motor.

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76. A sample testing system comprising, in combination:

tube sealing apparatus having a tube compression and sealing member to laterally seal a flexible plastic tube containing a fluid sample material, whereby a fluid-tight tubule containing a portion of the fluid sample material can be formed between axially spaced lateral seals;

a reagent injector having at least one needle in fluid communication with a reagent reservoir, and a needle actuator to insert the needle into a tubule and inject a quantity of reagent into a tubule;

a flow control device comprising a contact member movable into contact with a tubule to effect mechanically induced fluid flow within a fluid passageway in the tubule; and

an inspection system having a light detector to receive light passed through a tubule and to generate an output signal corresponding to a condition of the fluid sample material within a tubule.

77. A method of performing a sample assay comprising the following steps, in combination:

collecting a sample of fluid material into a length of substantially transparent, flexible, heat-sealable, plastic tube;

inserting the tube into a sample testing machine comprising a tube sealing apparatus, a reagent injector having at least one needle in fluid communication with a reagent reservoir and a needle actuator to insert the needle into a tubule and inject a quantity of a reagent, a flow control device comprising a contact member movable into contact with a tubule to effect mechanically induced fluid flow within the tubule, and an inspection system having a light detector to receive light passed through a tubule and to generate an output signal corresponding to a condition of the sample of fluid material within a tubule;

actuating the tube sealing apparatus to seal lengths of the tube into tubules;

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actuating the needle actuator to insert the needle into a selected tubule and inject the reagent to form a mixture of the sample of fluid material and the reagent in the selected tubule;

actuating the flow control device to mix the mixture of the sample of fluid material and the reagent; and

actuating the inspection system to inspect the mixture and to generate an output signal corresponding to a condition of the mixture.

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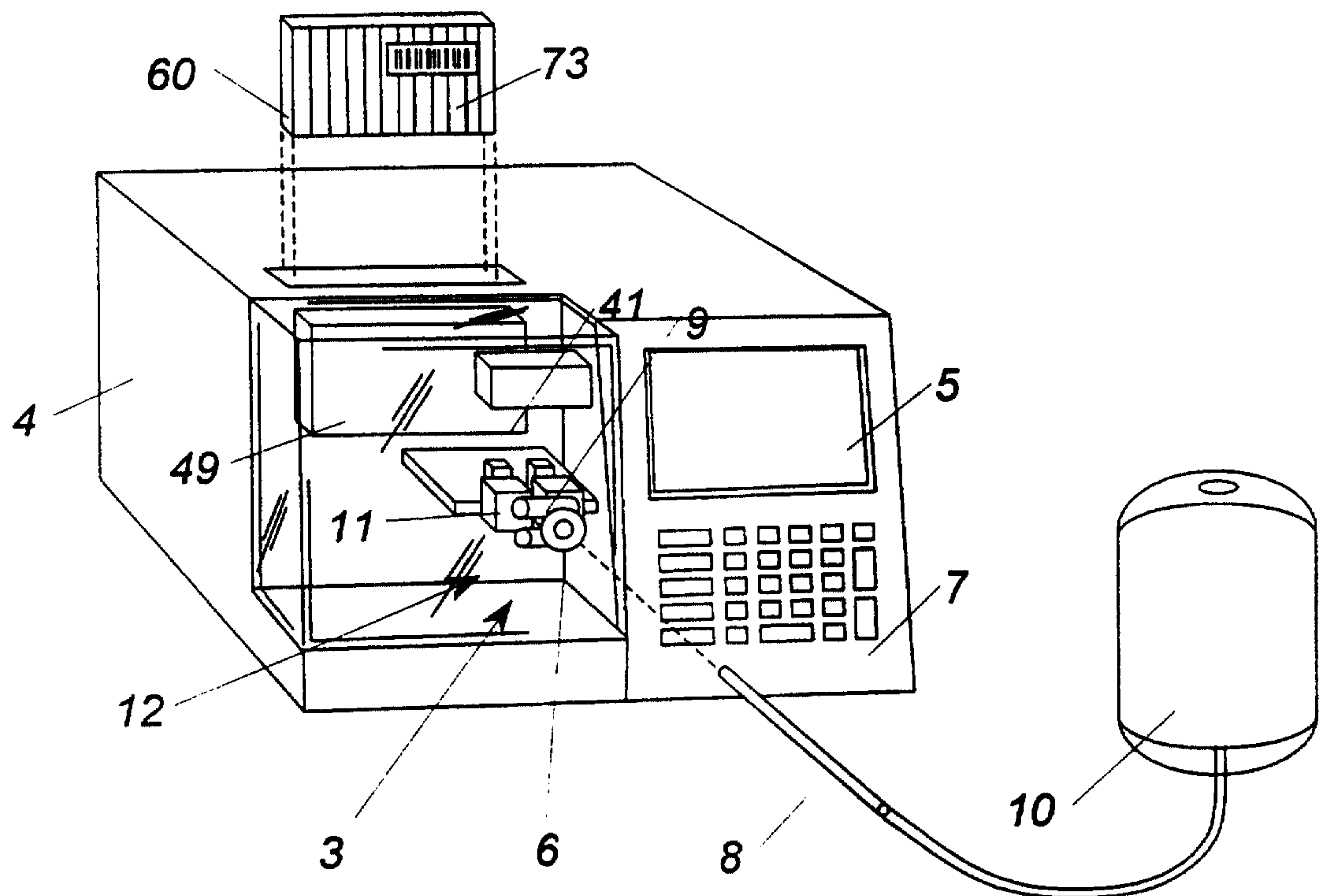


FIG. 1

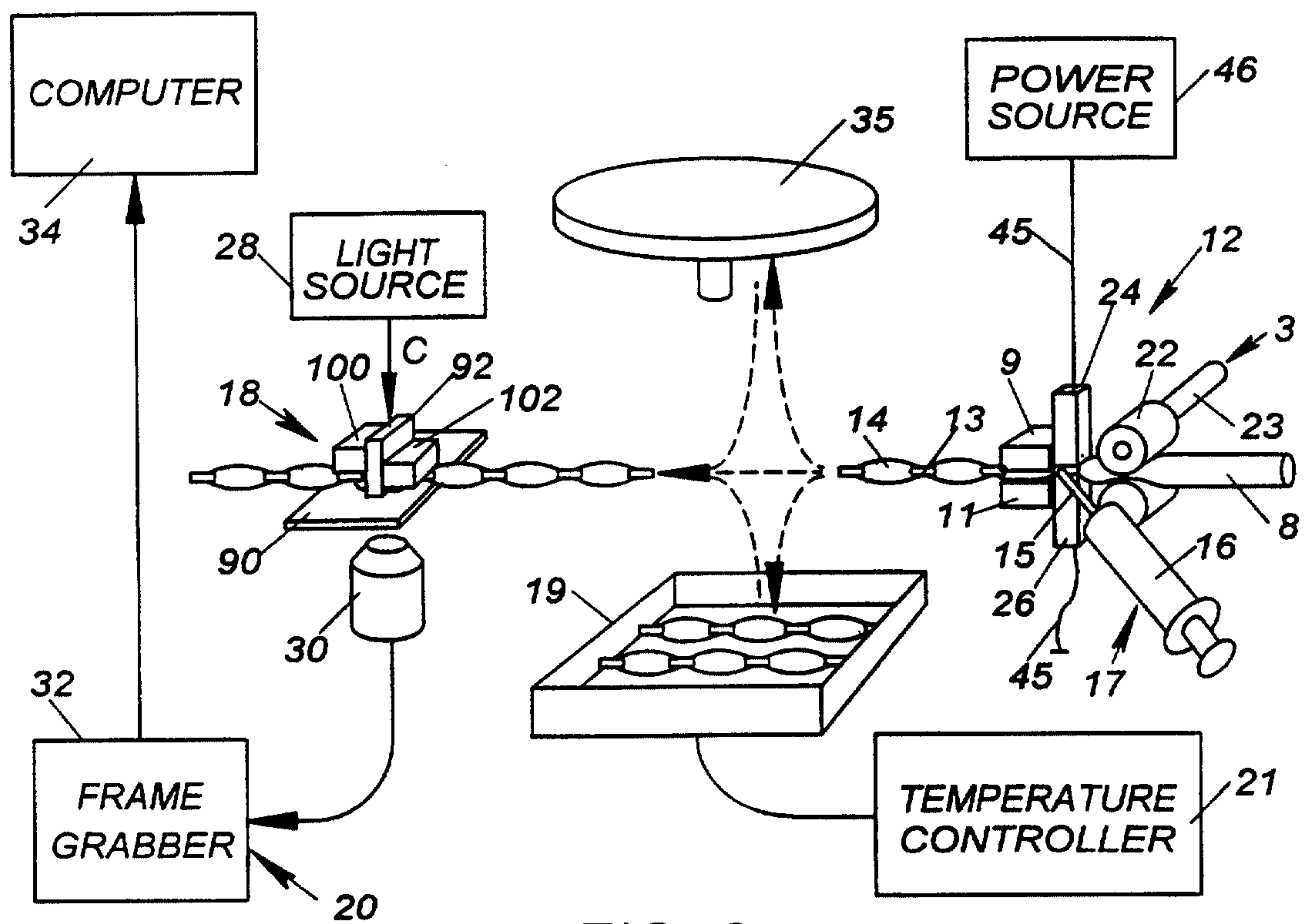


FIG. 2

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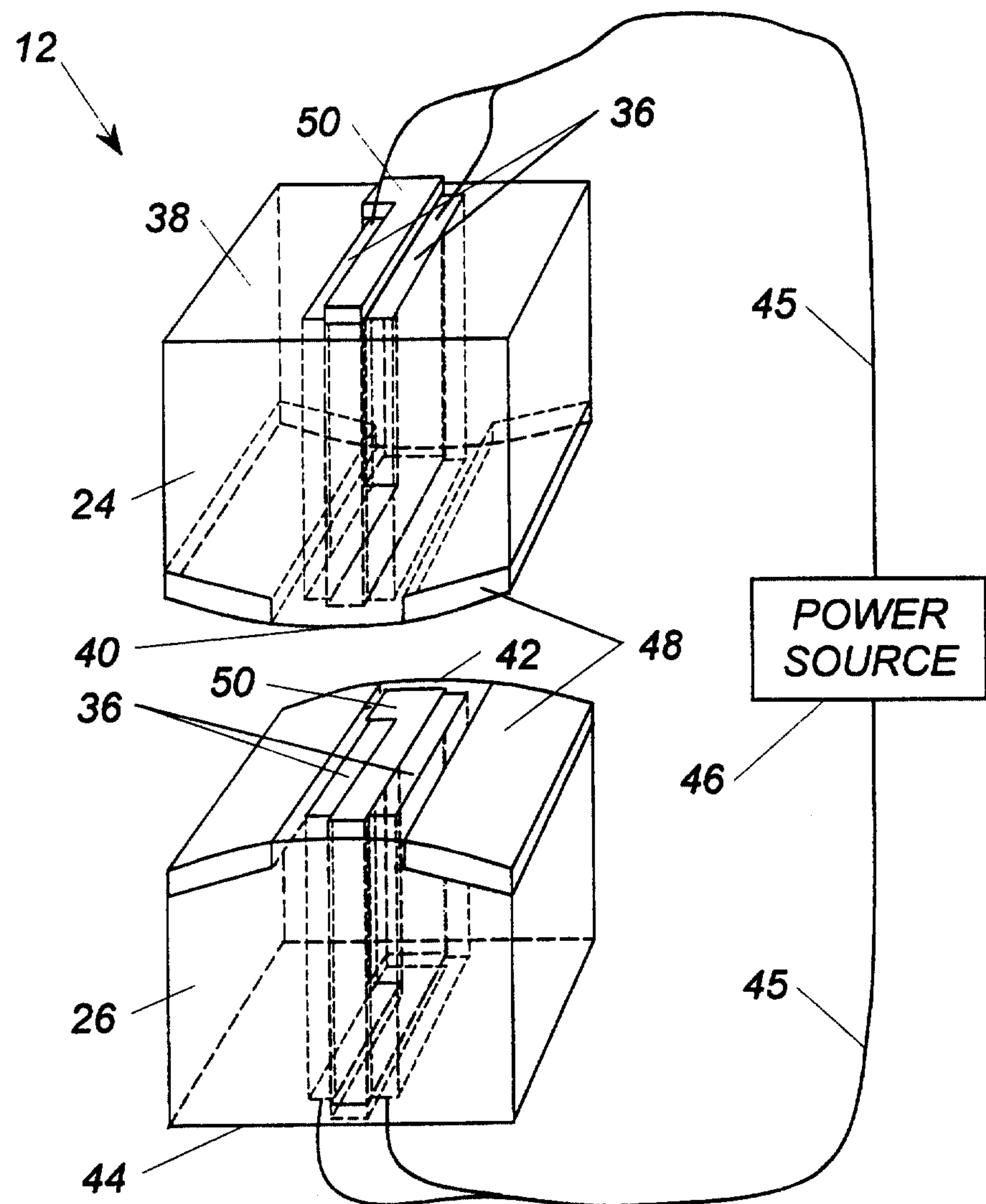


FIG. 3

SUBSTITUTE SHEET (RULE 26)

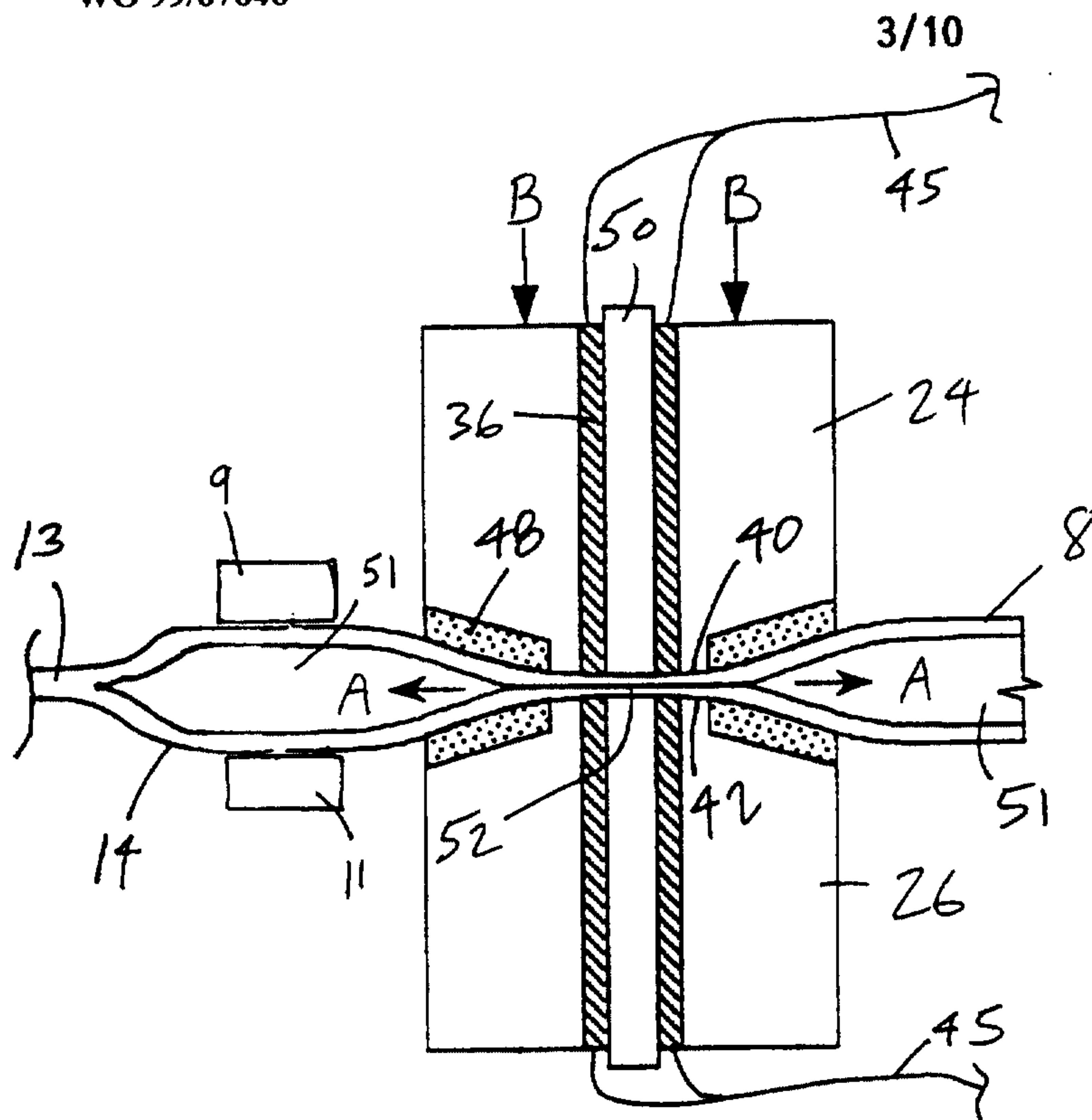


FIG. 4

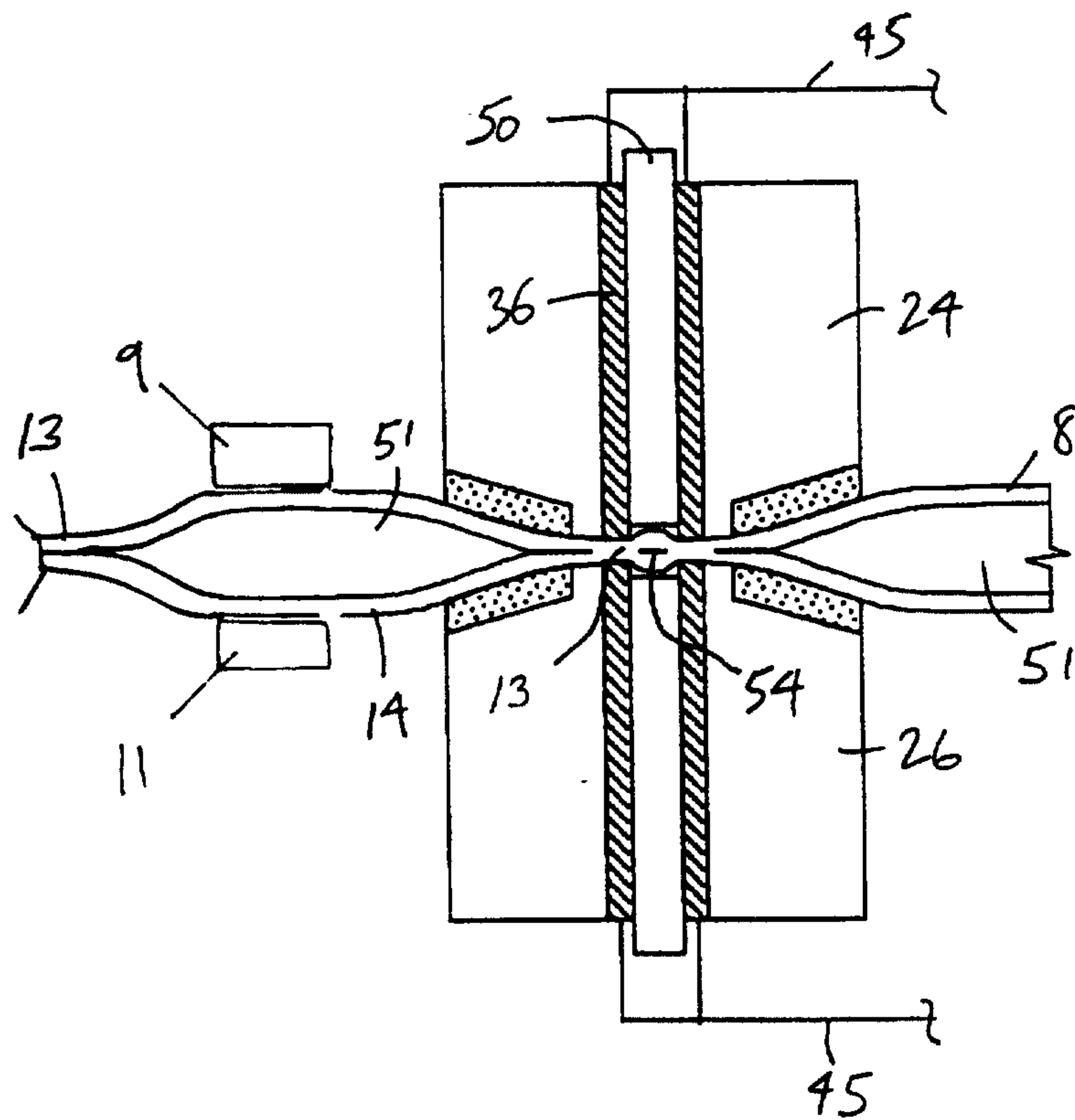


Fig. 5

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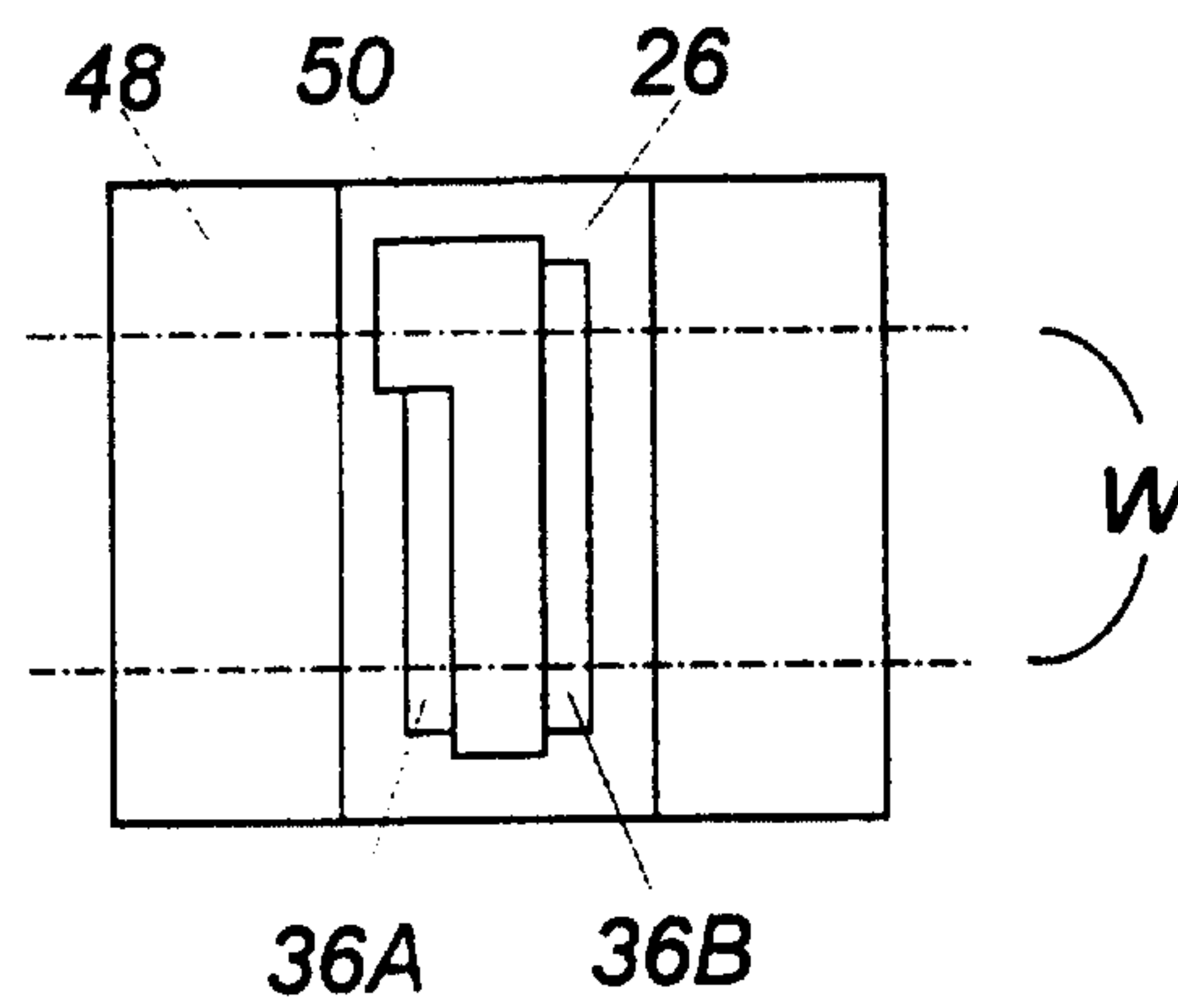


FIG. 6

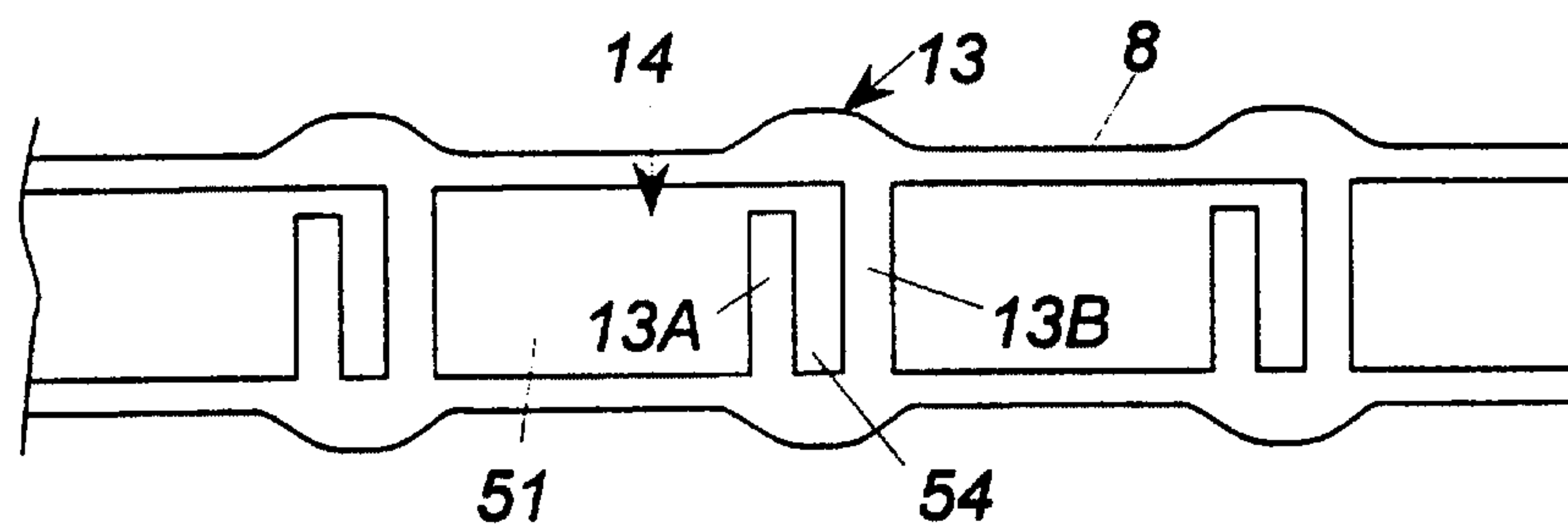


FIG. 7

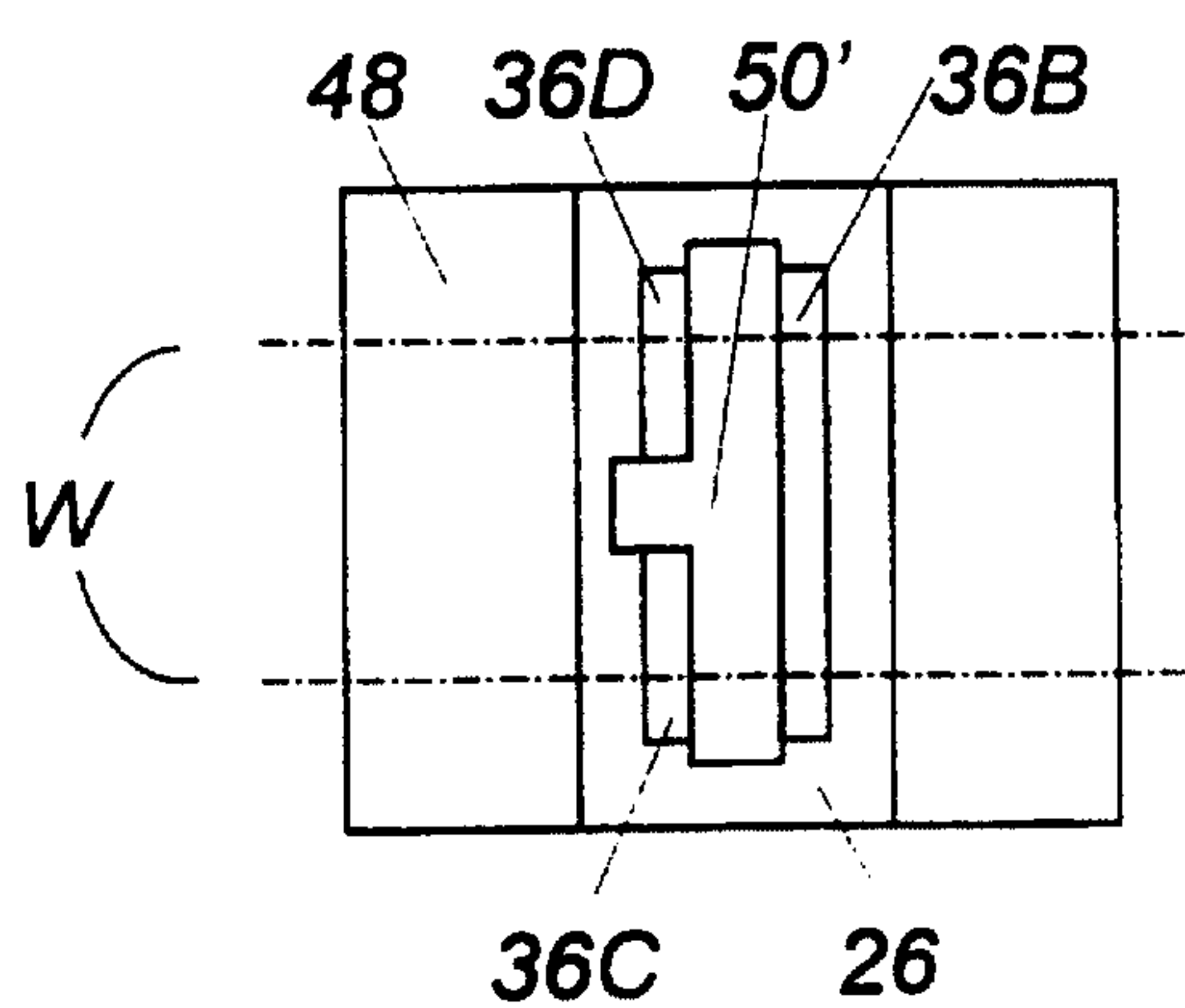


FIG. 8

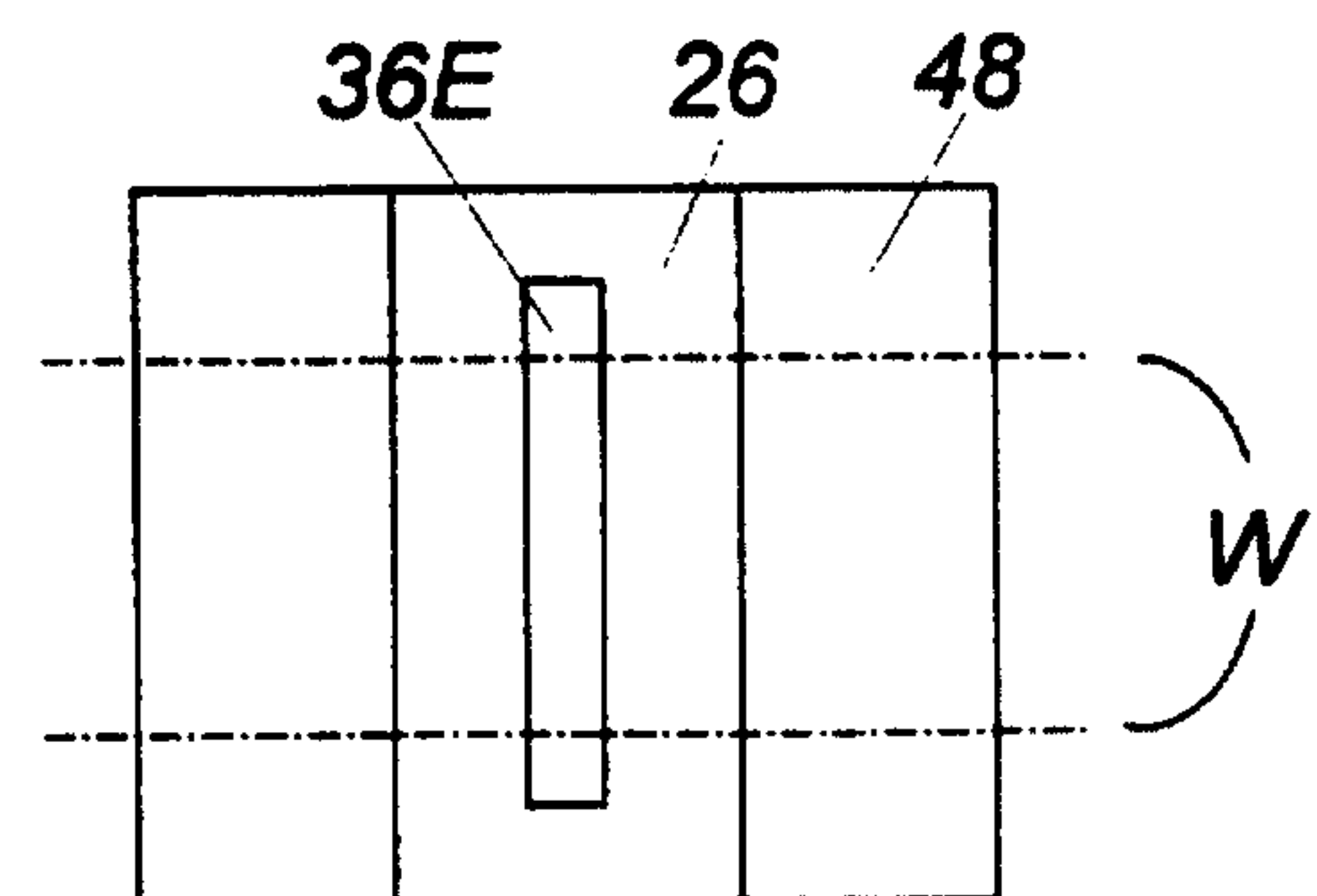


FIG. 9

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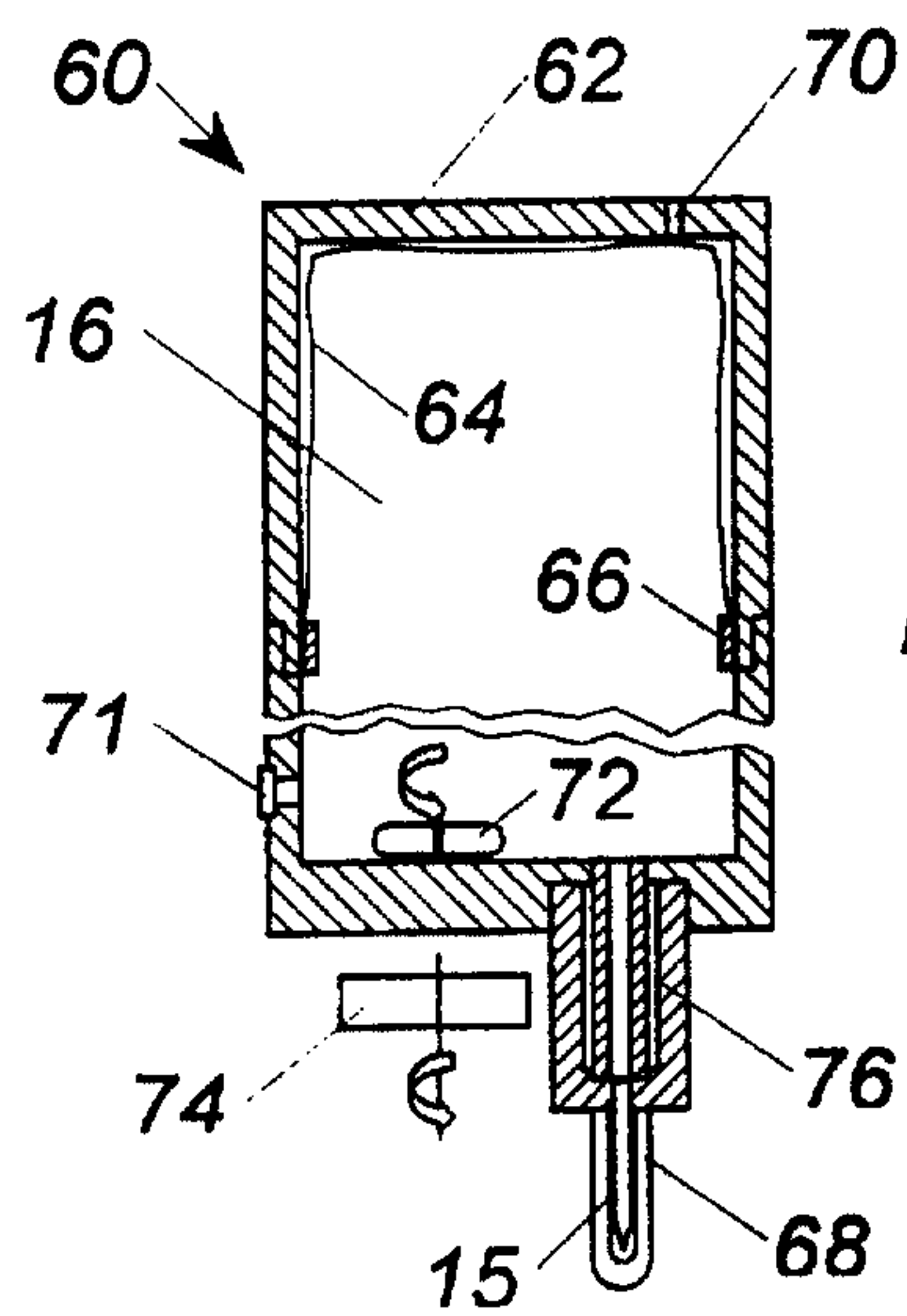


FIG. 10

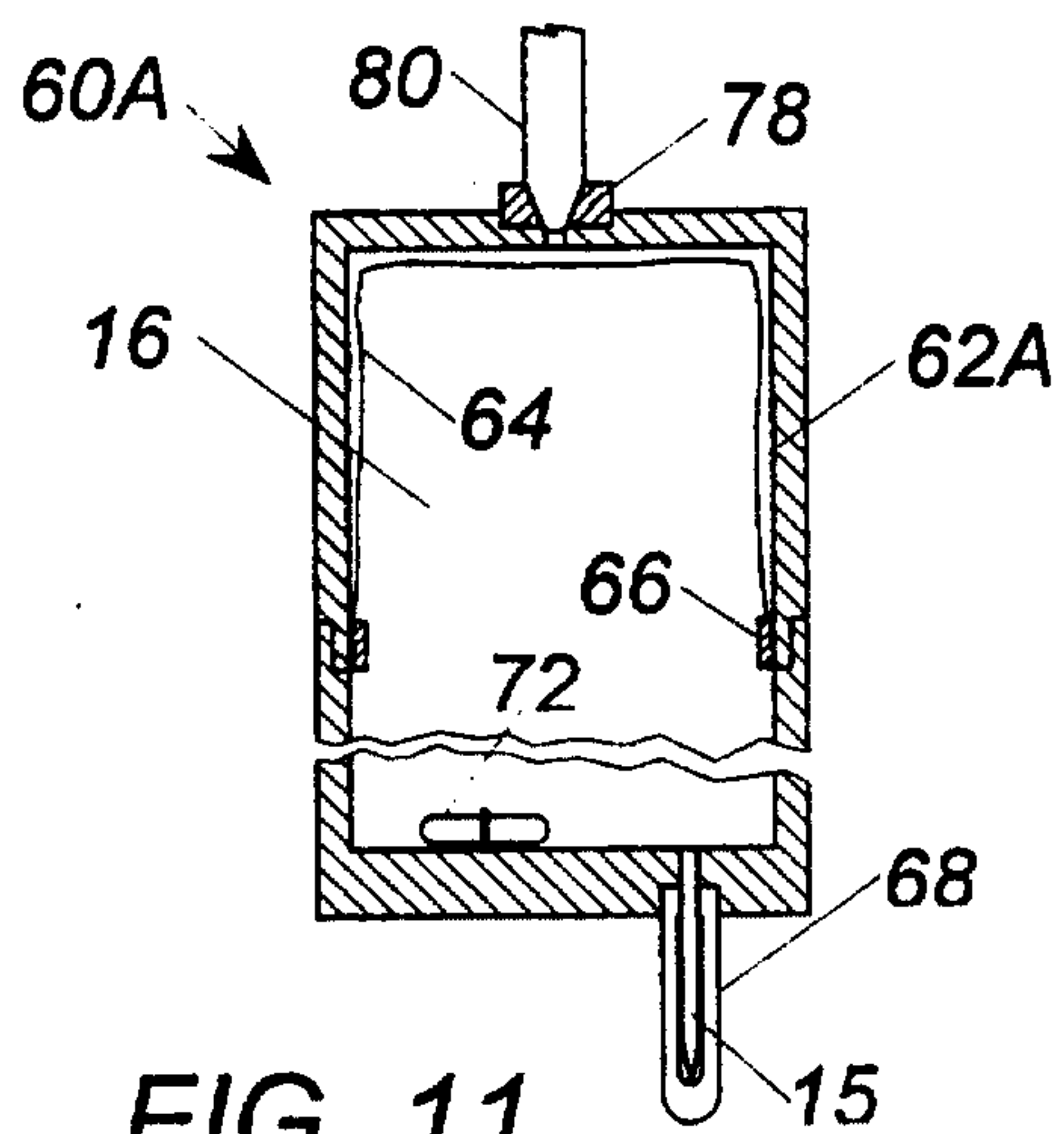


FIG. 11

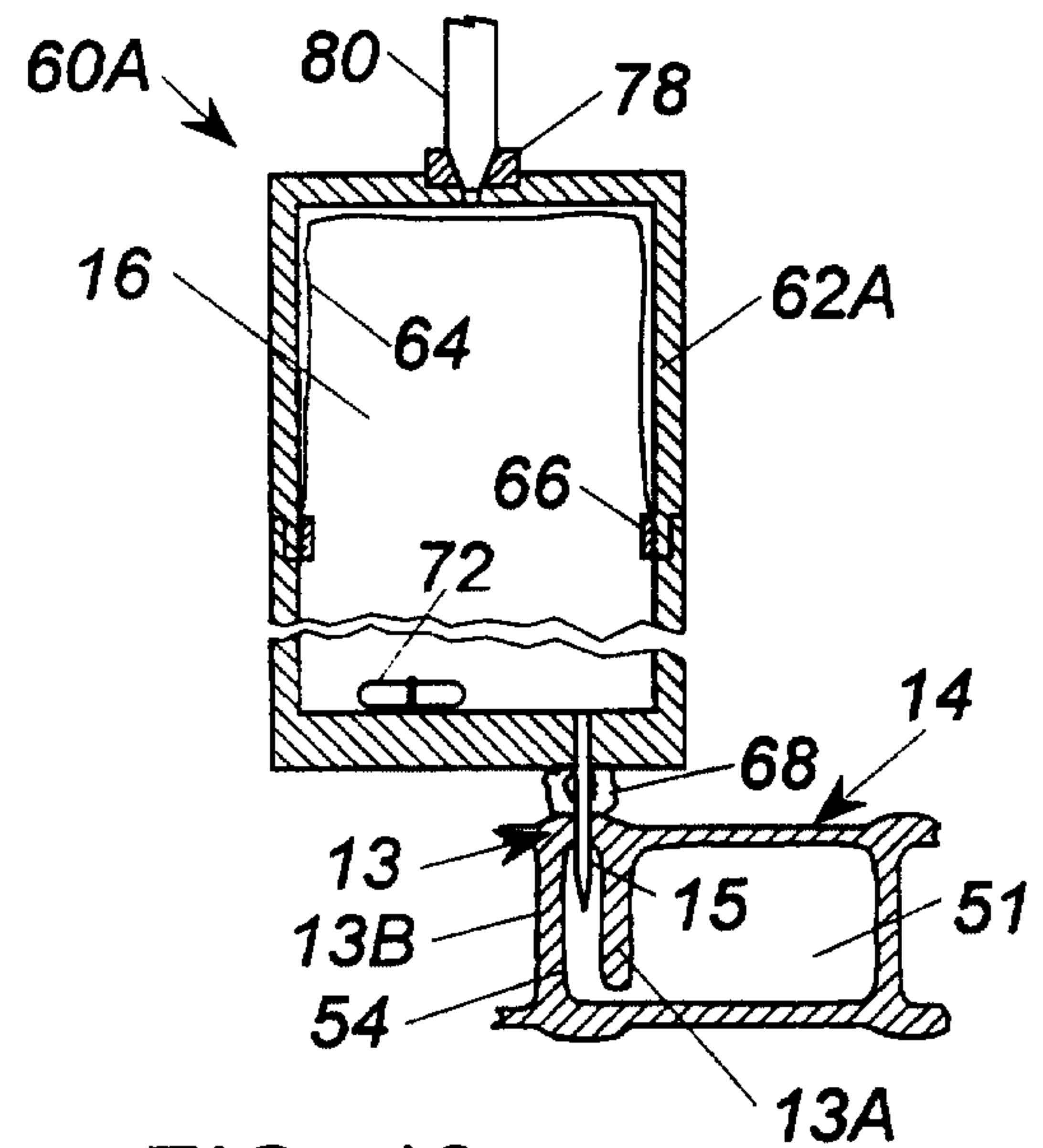


FIG. 12

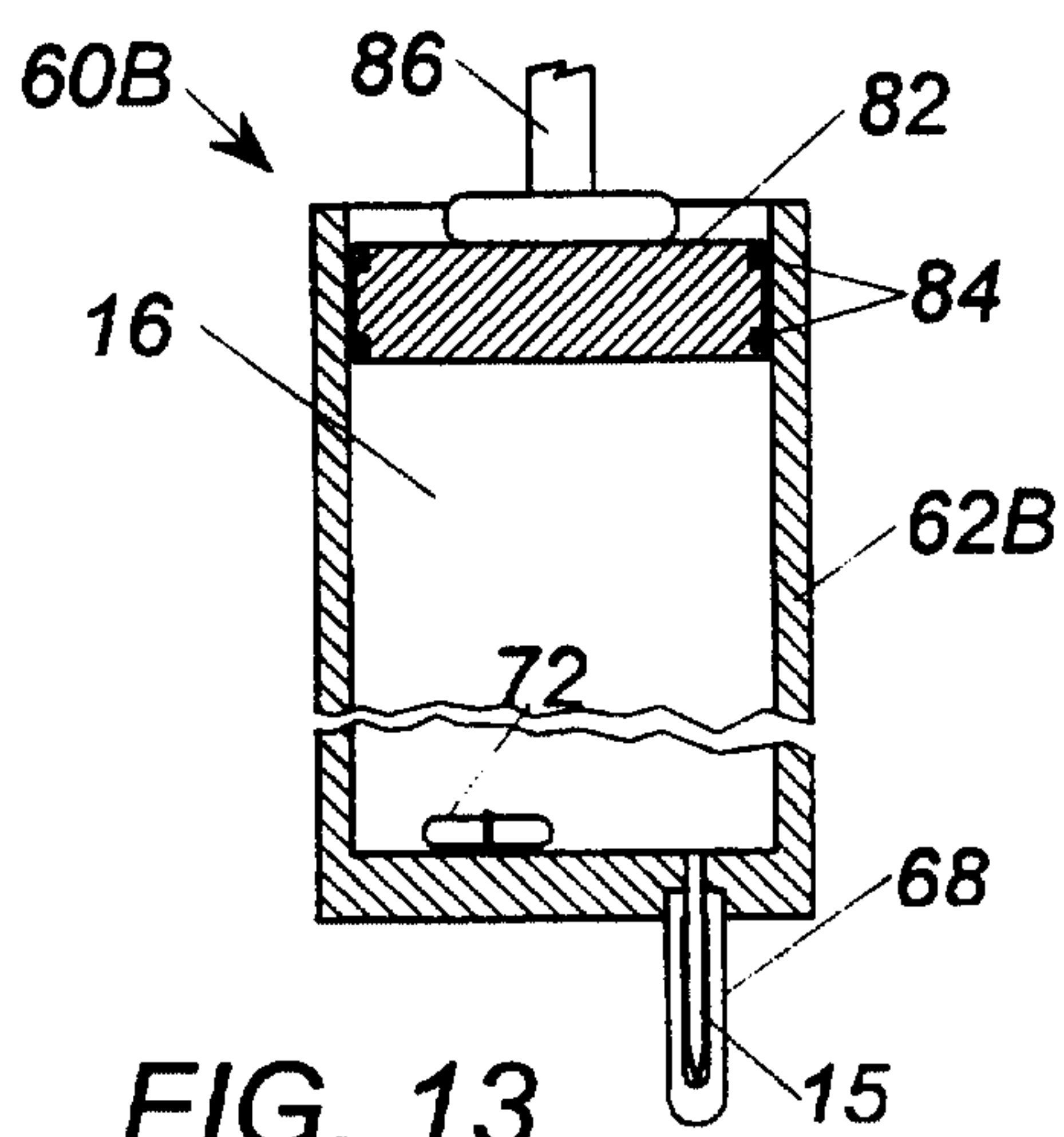


FIG. 13

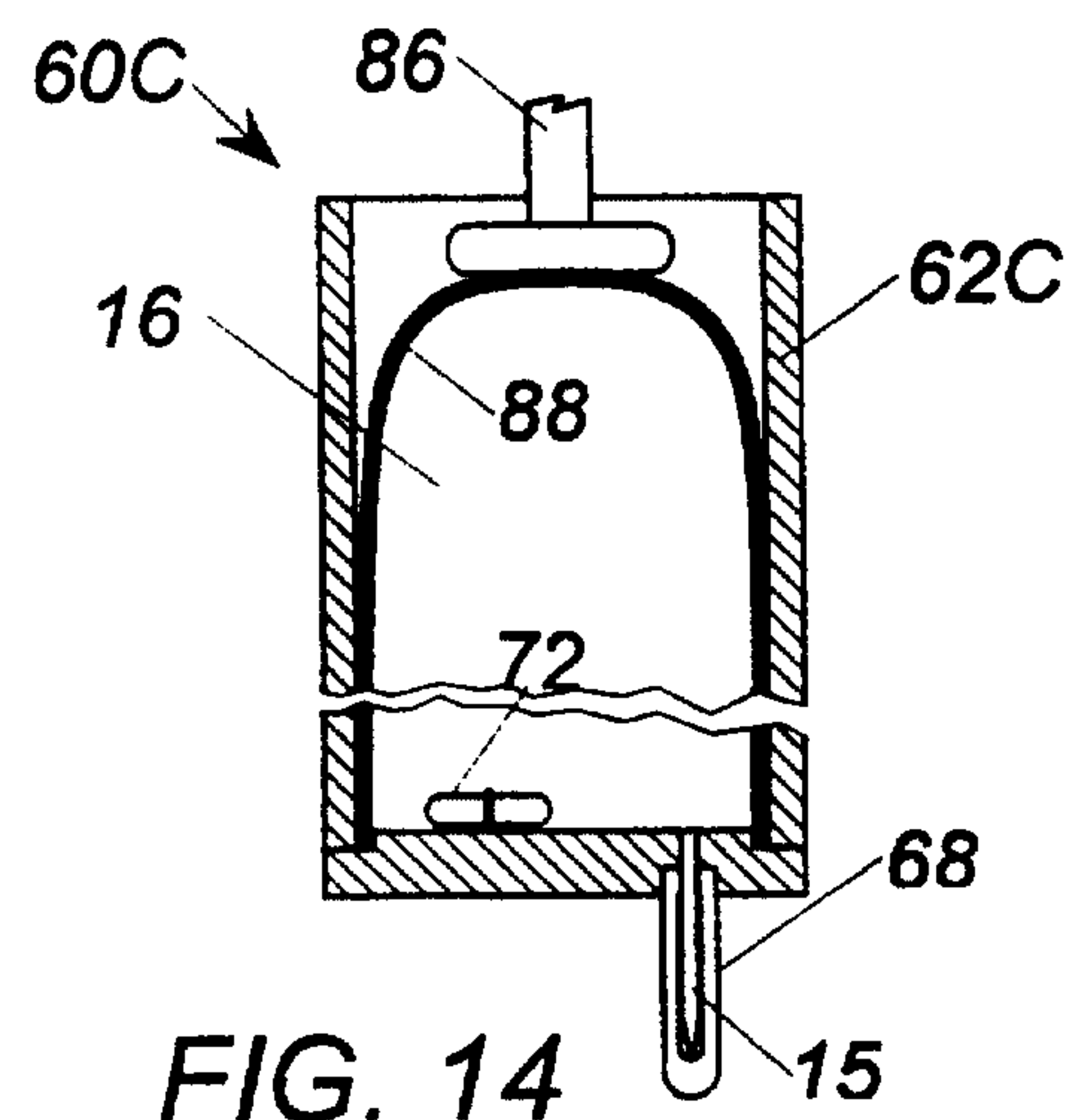


FIG. 14

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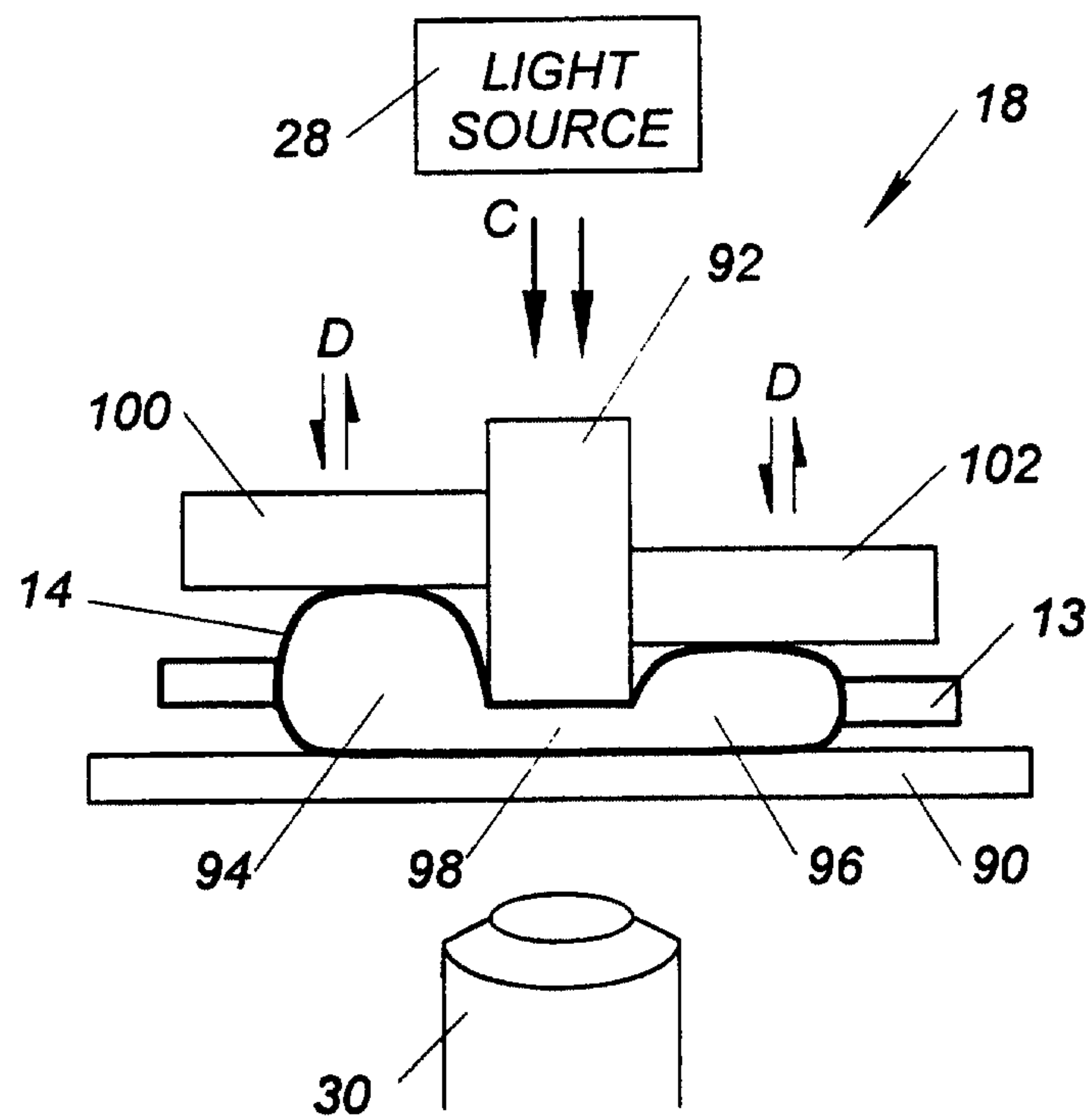


FIG. 15

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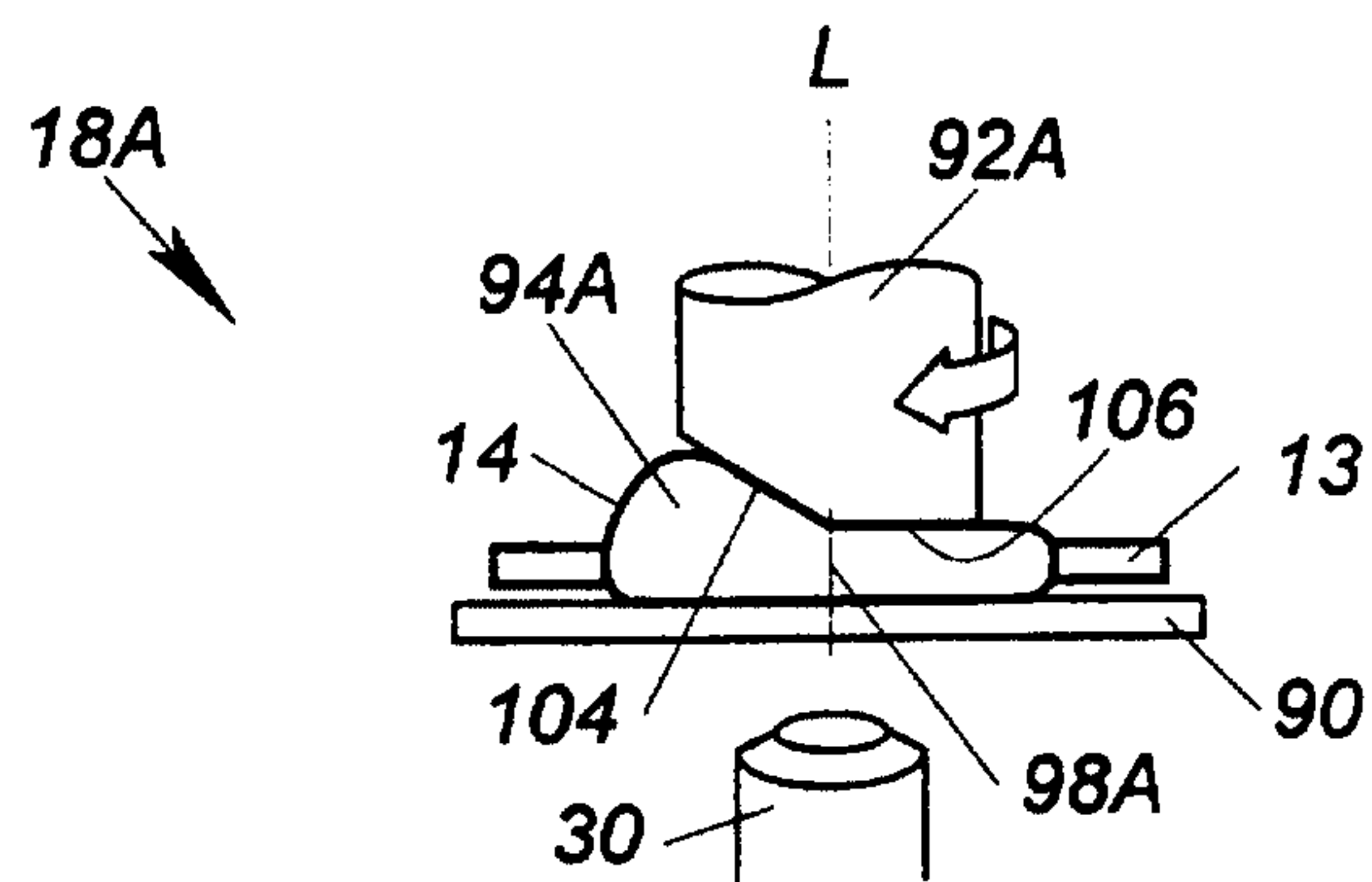


FIG. 16

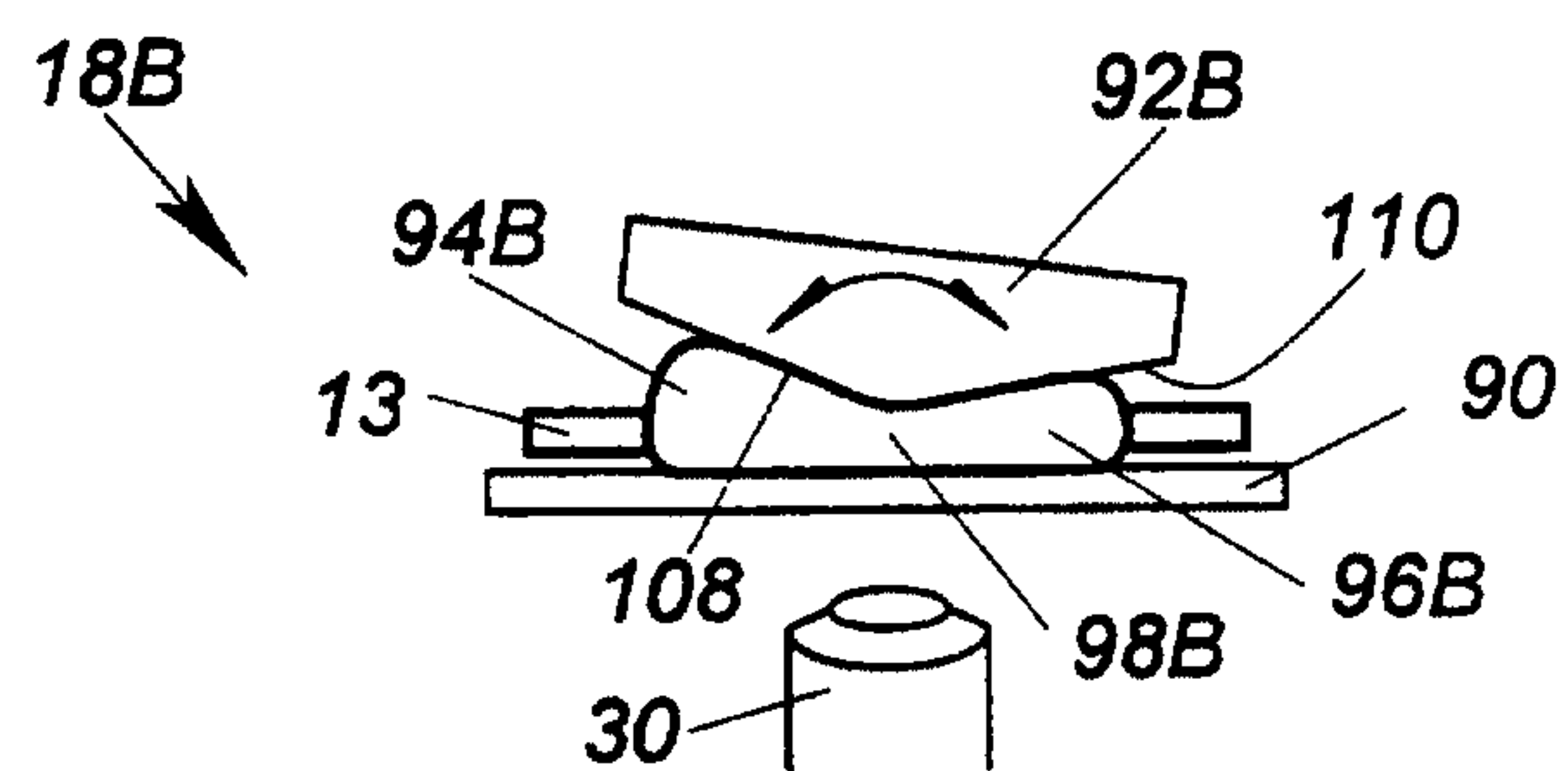


FIG. 17

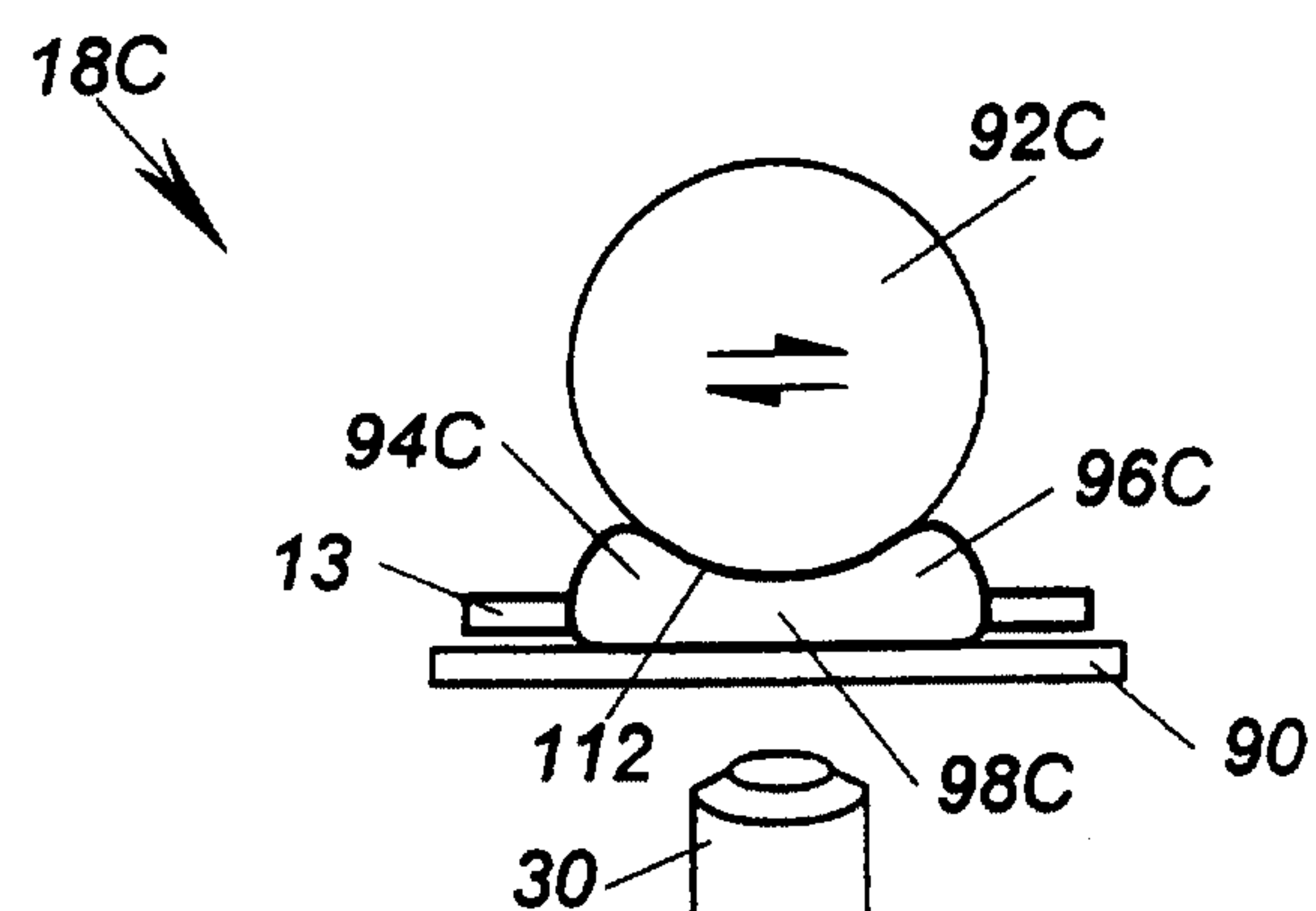
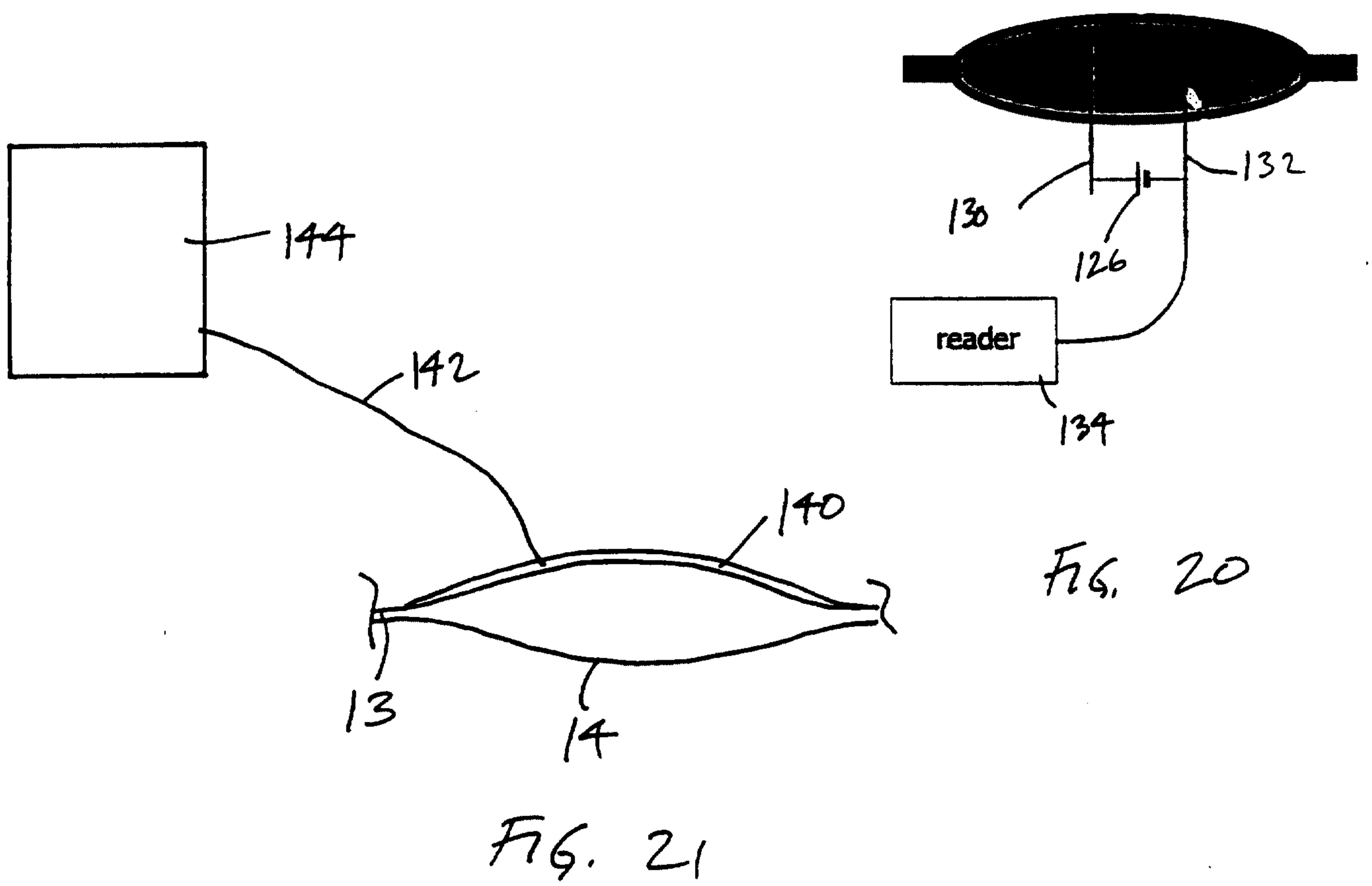
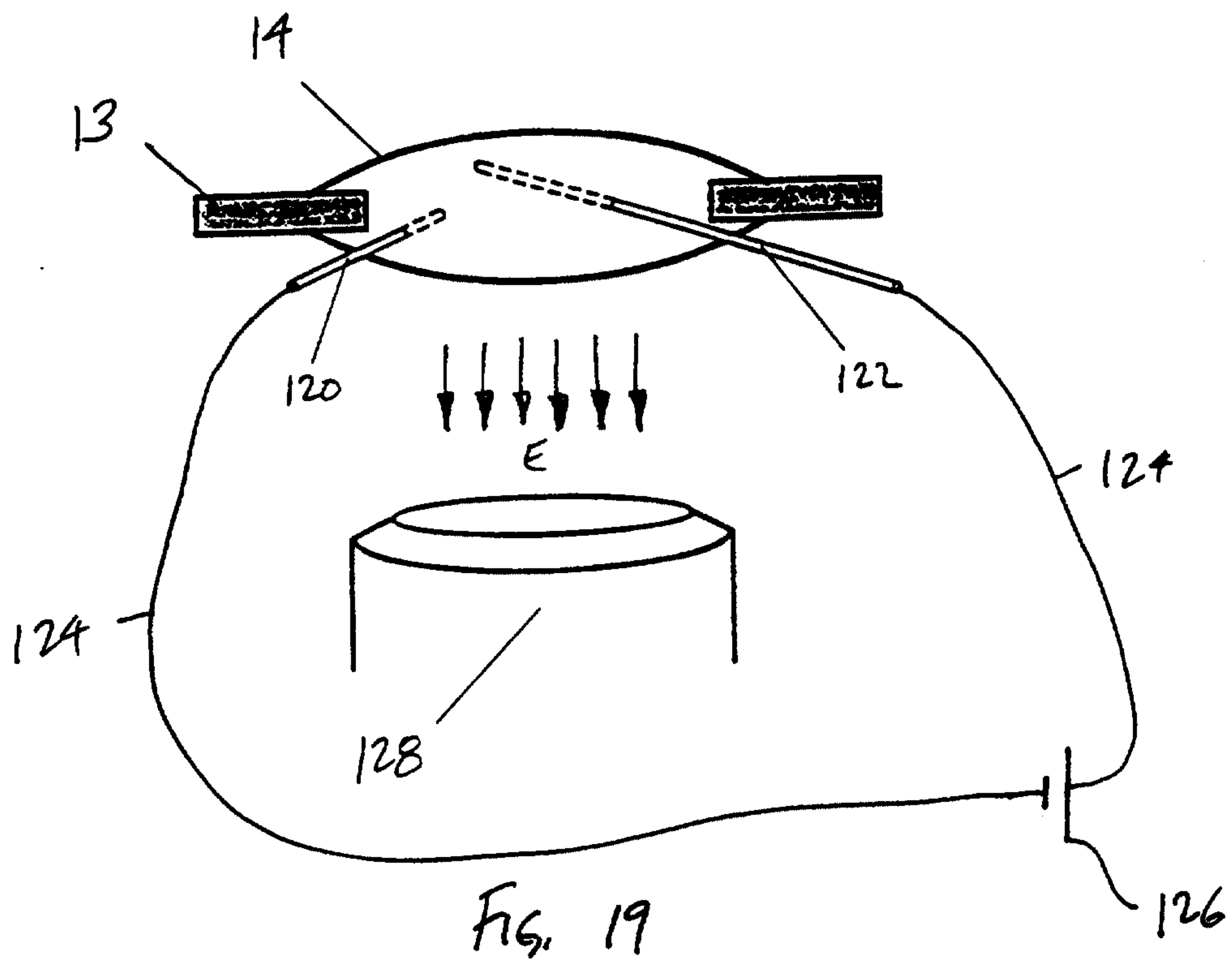


FIG. 18

SUBSTITUTE SHEET (RULE 26)



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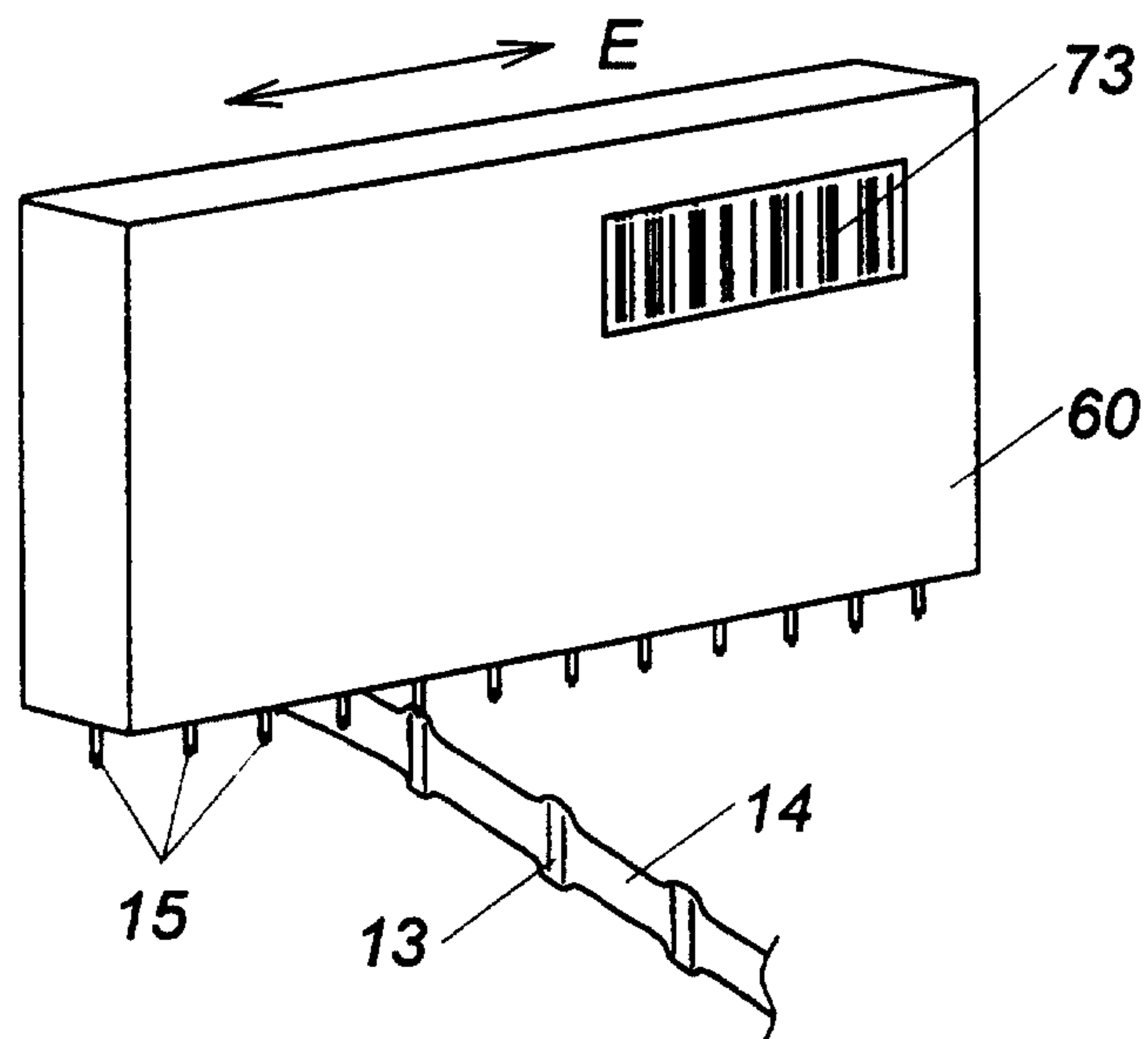


FIG. 22

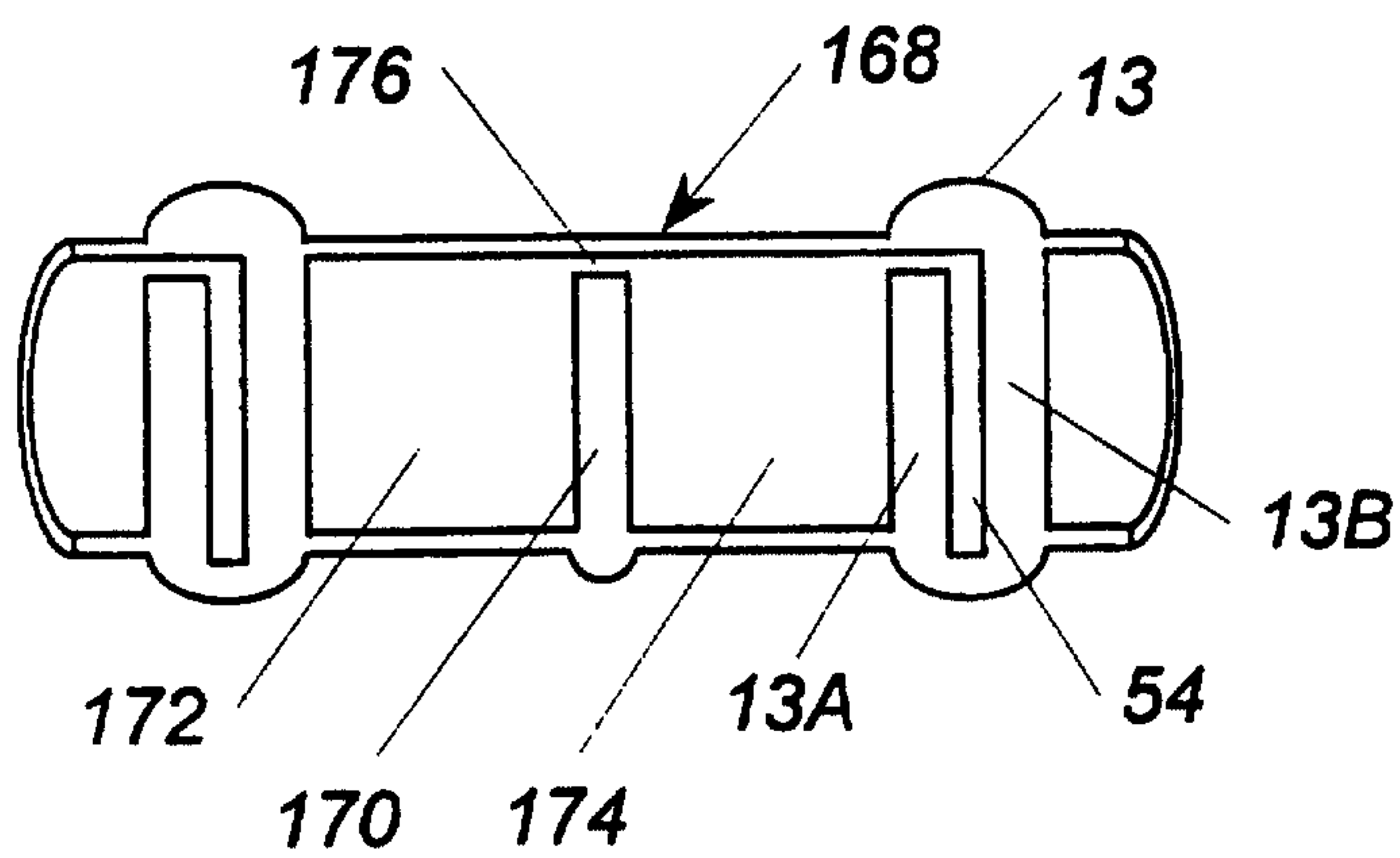


FIG. 25

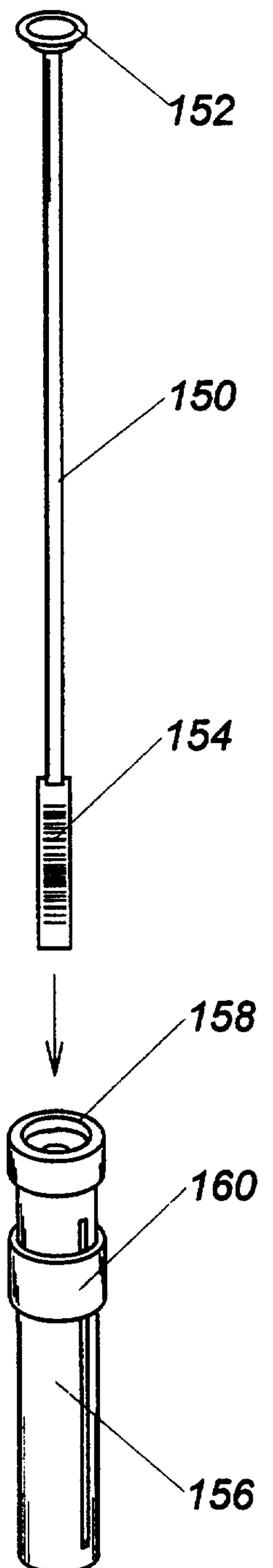
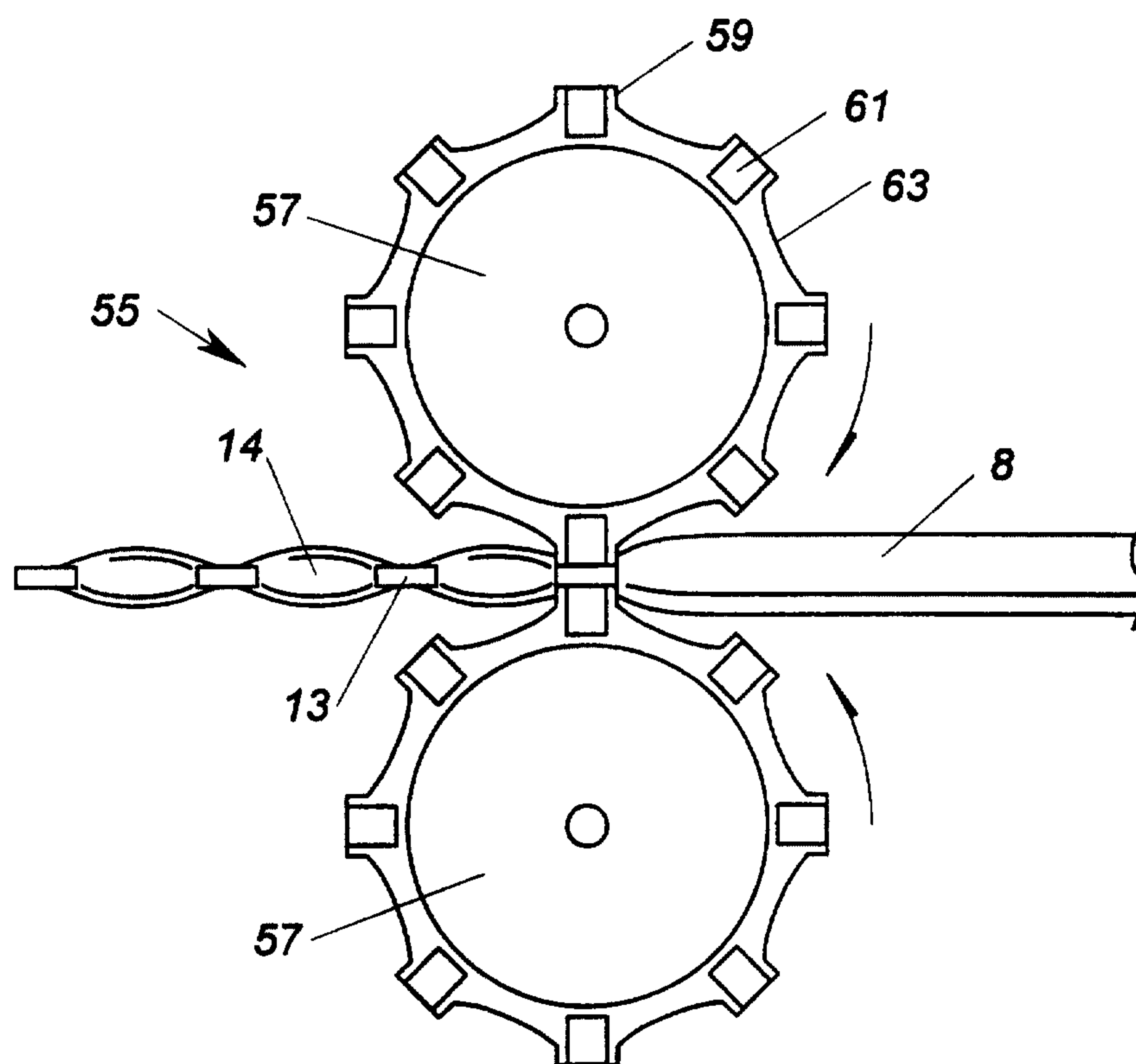


FIG. 23

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**FIG. 24**

