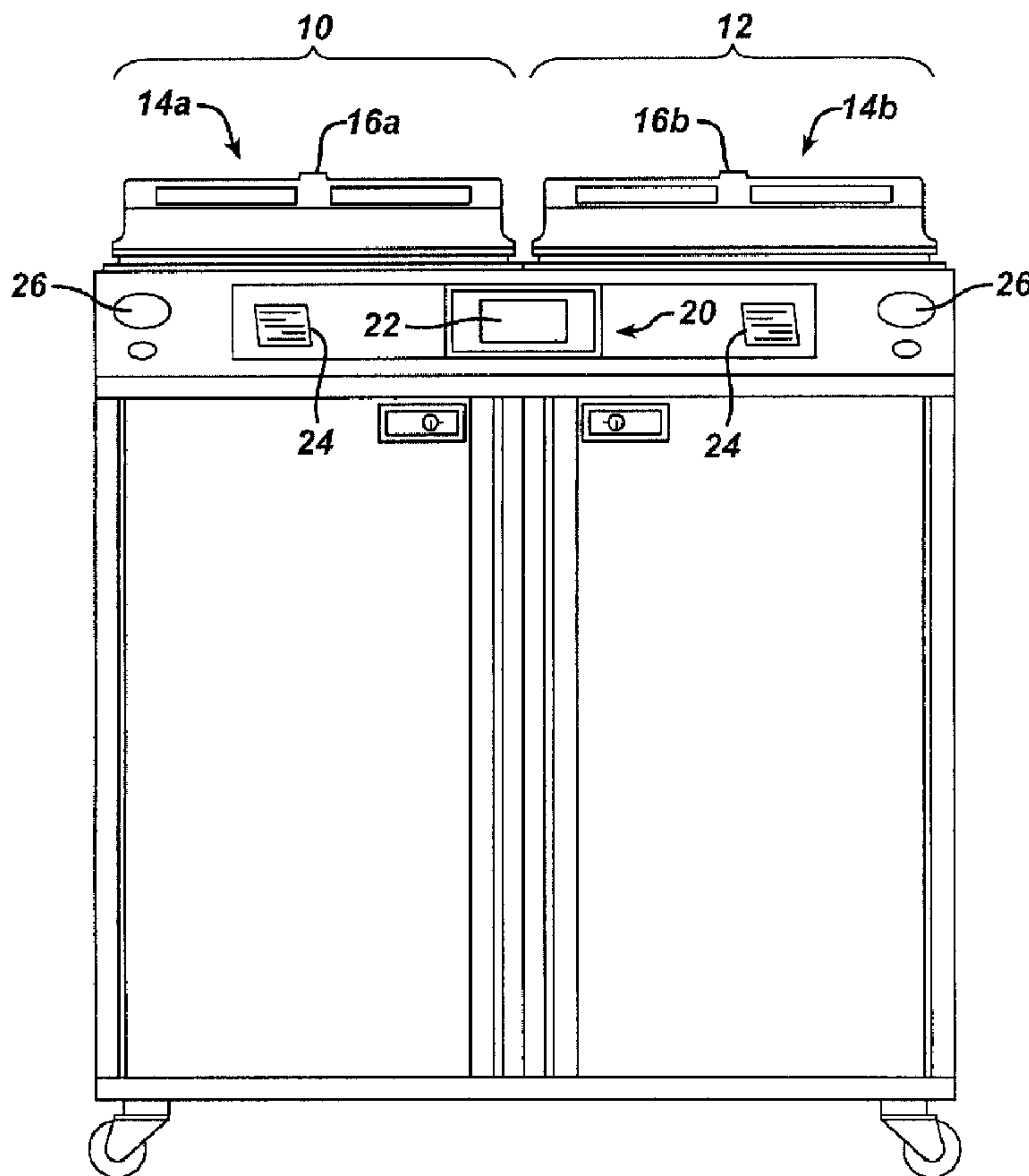




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(54) Titre : METHODE DE MISE A L'ESSAI DE SOLUTIONS AU MOYEN D'UNITES DE RETRAITEMENT AUTOMATISE  
 POUR ENDOSCOPES  
 (54) Title: AUTOMATED ENDOSCOPE REPROCESSOR SOLUTION TESTING



(57) Abrégé/Abstract:

A method measures a property of a solution spectroscopically by placing a quantity of the solution into a cuvette and passing a light therethrough. A reservoir receives a quantity of the solution and bubbles are driven off prior to putting a sample therefrom into the cuvette for measuring.



## ABSTRACT

A method measures a property of a solution spectroscopically by placing a quantity of the solution into a cuvette and passing a light therethrough. A reservoir  
5 receives a quantity of the solution and bubbles are driven off prior to putting a sample therefrom into the cuvette for measuring.

AUTOMATED ENDOSCOPE REPROCESSOR  
SOLUTION TESTING

5 BACKGROUND OF THE INVENTION

The present invention relates to the decontamination arts including the sterilization arts. It finds particular application in conjunction with the decontamination of medical devices, especially medical devices such as endoscopes and other devices  
10 having channels or lumens that must be decontaminated after use.

Endoscopes and similar medical devices having channels or lumens formed therethrough are being used on an ever increasing basis in the performance of medical procedures. The popularity of these devices has led to calls for improvements in the  
15 decontamination of these devices between use, both in terms of the speed of the decontamination and the effectiveness of the decontamination.

One popular method for cleaning and disinfection or sterilization of such endoscopes employs an automated endoscope reprocessor which both washes and then  
20 disinfects or sterilizes the endoscope. Typically such a unit comprises a basin with a selectively opened and closed cover member to provide access to the basin. Pumps connect to various channels through the endoscope to flow fluid therethrough and an additional pump flows fluid over the exterior surfaces of the endoscope. Typically, a detergent washing cycle is followed by rinsing and then a sterilization or disinfection  
25 cycle and rinse.

To insure adequate washing and sterilization it may be desirable to measure the strength of fluids used for washing and sterilization. In particular, it is desirable to make sure that the proper concentration has been achieved in the circulating fluid.  
30

## SUMMARY OF THE INVENTION

An endoscope processor according to the present invention incorporates a solution measuring system. The measuring system comprises a cuvette for holding a sample of the solution, a light source for passing a light through the cuvette and the sample, and a light sensing mechanism for sensing light passing through the cuvette and the sample. A reservoir is provided for receiving a quantity of the solution containing bubbles. A pump associated with the reservoir allows pumping of a quantity of solution out of the reservoir through a first path from the reservoir and/or a second path from the reservoir to the cuvette. A control system associated with the pump is programmed to first direct the pump to move a portion of the quantity of fluid in the reservoir out through the first path, whereby to drive bubbles therein out of the reservoir, and then to direct a sample of the liquid into the cuvette.

Preferably, the solution comprises an aldehyde., as for instance orthophthalaldehyde.

Preferably, a light path through the sample in the cuvette is between 1 mm and 5 mm, more preferably between 1 mm and 3 mm.

Preferably, the first path leaves the reservoir from an upper portion thereof. The first path can be the same as the second path with the bubbles being removed through cuvette. The control system is then preferably programmed to pump fluid out of the reservoir for a time period sufficient to pump substantially all of the bubbles in the solution out through the cuvette whereby to leave a quantity of solution in the cuvette substantially free of bubbles.

Preferably, the control system is programmed to delay pumping of fluid out of the reservoir for a time period after the reservoir is filled sufficient to allow the bubbles in the solution to float to the surface.

A method according to the present invention in an endoscope processor provides for measuring a property of a solution to be applied to the endoscope. The

method comprises: collecting a quantity of the solution in a reservoir; directing a portion of the solution out from the reservoir through a first path to carry bubbles in the solution out of the reservoir; then directing a sample of the solution from the reservoir to a cuvette; measuring the property of the solution in the sample in the cuvette by  
5 passing light through the cuvette and the sample and reading said light passing through said light and said sample.

Preferably, the property of the solution being measured is the level of sterilant therein.

10

#### BRIEF DESCRIPTION OF THE DRAWINGS

The invention may take form in various components and arrangements of components and in various steps and arrangements of steps. The drawings are for  
15 purposes of illustrating preferred embodiments only, and are not to be construed as limiting the invention.

FIG. 1 is a front elevational view of a decontamination apparatus in accordance with the present invention;

20

FIG. 2 is a diagrammatic illustration of the decontamination apparatus shown in FIG. 1, with only a single decontamination basin shown for clarity;

FIG. 3 is a cut-away view of an endoscope suitable for processing in the  
25 decontamination apparatus of FIG. 1;

FIG. 4 is a diagrammatic illustration of spectroscopic fluid measuring subsystem of the decontamination apparatus of FIG. 2; and

30 FIG. 5 is a perspective view of the spectroscopic fluid measuring subsystem of FIG. 4.

## DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

FIG. 1 shows a decontamination apparatus for decontaminating endoscopes and other medical devices which include channels or lumens formed therethrough; FIG. 2 shows the apparatus in block diagram form. The decontamination apparatus generally includes a first station 10 and a second station 12 which are at least substantially similar in all respects to provide for the decontamination of two different medical devices simultaneously or in series. First and second decontamination basins 14a, 14b receive the contaminated devices. Each basin 14a, 14b is selectively sealed by a lid 16a, 16b, respectively, preferably in a microbe-blocking relationship to prevent the entrance of environmental microbes into the basins 14a, 14b during decontamination operations. The lids can include a microbe removal or HEPA air filter formed therein for venting.

A control system 20 includes one or more microcontrollers, such as a programmable logic controller (PLC), for controlling decontamination and user interface operations. Although one control system 20 is shown herein as controlling both decontamination stations 10, 12, those skilled in the art will recognize that each station 10, 12 can include a dedicated control system. A visual display 22 displays decontamination parameters and machine conditions for an operator and at least one printer 24 prints a hard copy output of the decontamination parameters for a record to be filed or attached to the decontaminated device or its storage packaging. The visual display 22 is preferably combined with a touch screen input device. Alternatively, a keypad or the like is provided for input of decontamination process parameters and for machine control. Other visual gauges 26 such as pressure meters and the like provide digital or analog output of decontamination or medical device leak testing data.

FIG. 2 diagrammatically illustrates one station 10 of the decontamination apparatus. Those skilled in the art will recognize that the decontamination station 12 is preferably similar in all respects to the station 10 illustrated in FIG. 2. However, the station 12 has not been shown in FIG. 2 for clarity. Further, the decontamination apparatus can be provided with a single decontamination station or multiple stations.

The decontamination basin 14a receives an endoscope 200 (see FIG. 3) or other medical device therein for decontamination. Any internal channels of the endoscope 200 are connected with flush lines 30. Each flush line 30 is connected to an outlet of a pump 32. The pumps 32 are preferably peristaltic pumps or the like that pump fluid, such as liquid and air, through the flush lines 30 and any internal channels of the medical device. Specifically, the pumps 32 either can draw liquid from the basin 14a through a filtered drain 34 and a first valve S1, or can draw decontaminated air from an air supply system 36 through a valve S2. The air supply system 36 includes a pump 38 and a microbe removal air filter 40 that filters microbes from an incoming air stream. It is preferable that each flush line 30 be provided with a dedicated pump 32 to ensure adequate fluid pressure and to facilitate the individual monitoring of the fluid pressure in each flush line 30. A pressure switch or sensor 42 is in fluid communication with each flush line 30 for sensing excessive pressure in the flush line. Any excessive pressure sensed is indicative of a partial or complete blockage, e.g., by bodily tissue or dried bodily fluids, in a device channel to which the relevant flush line 30 is connected. The isolation of each flush line 30 relative to the others allows the particular blocked channel to be easily identified and isolated, depending upon which sensor 42 senses excessive pressure.

The basin 14a is in fluid communication with a water source 50 such as a utility or tap water connection including hot and cold inlets and a mixing valve 52 flowing into a break tank 56. A microbe removal filter 54, such as a 0.2  $\mu\text{m}$  or smaller absolute pore size filter, decontaminates the incoming water which is delivered into the break tank 56 through the air gap to prevent backflow. A pressure type level sensor 59 monitors liquid levels within the basin 14a. An optional water heater 53 can be provided if an appropriate source of hot water is not available.

The condition of the filter 54 can be monitored by directly monitoring the flow rate of water therethrough or indirectly by monitoring the basin fill time using a float switch or the like. When the flow rate drops below a select threshold, this indicates a partially clogged filter element that requires replacement.

A basin drain 62 drains liquid from the basin 14a through an enlarged helical tube 64 into which elongated portions of the endoscope 200 can be inserted. The drain 62 is in fluid communication with a recirculation pump 70 and a drain pump 72. The recirculation pump 70 recirculates liquid from the basin drain 62 to a spray nozzle assembly 60 which sprays the liquid into the basin 14a and onto the endoscope 200. Coarse and fine screens 71 and 73, respectively, filter out particles in the recirculating fluid. The drain pump 72 pumps liquid from the basin drain 62 to a utility drain 74. A level sensor 76 monitors the flow of liquid from the pump 72 to the utility drain 74. The pumps 70 and 72 can be simultaneously operated such that liquid is sprayed into the basin 14a while it is being drained to encourage the flow of residue out of the basin and off of the device. Of course, a single pump and a valve assembly could replace the dual pumps 70, 72.

An inline heater 80, with temperature sensors 82, downstream of the recirculation pump 70 heats the liquid to optimum temperatures for cleaning and disinfection. A pressure switch or sensor 84 measures pressure downstream of the circulation pump 70.

Detergent solution 86 is metered into the flow upstream of the circulation pump 70 via a metering pump 88. A float switch 90 indicates the level of detergent available. Typically, only a small amount of disinfectant 92 is required. To more accurately meter this, a dispensing pump 94 fills a pre-chamber 96 under control of a hi/low level switch 98 and of course the control system 20. A metering pump 100 meters a precise quantity of disinfectant as needed.

Endoscopes and other reusable medical devices often include a flexible outer housing or sheath surrounding the individual tubular members and the like that form the interior channels and other parts of the device. This housing defines a closed interior space, which is isolated from patient tissues and fluids during medical procedures. It is important that the sheath be maintained intact, without cuts or other holes that would allow contamination of the interior space beneath the sheath.

Therefore, the decontamination apparatus includes means for testing the integrity of such as sheath.

An air pump, either the pump 38 or another pump 110, pressurizes the interior  
 5 space defined by the sheath of the device through a conduit 112 and a valve S5.  
 Preferably, a HEPA or other microbe-removing filter 113 removes microbes from the  
 pressurizing air. An overpressure switch 114 prevents accidental over pressurization of  
 the sheath. Upon full pressurization, the valve S5 is closed and a pressure sensor 116  
 looks for a drop in pressure in the conduit 112 which would indicate the escape of air  
 10 through the sheath. A valve S6 selectively vents the conduit 112 and the sheath  
 through an optional filter 118 when the testing procedure is complete. An air buffer  
 120 smoothes out pulsation of pressure from the air pump 110.

Preferably, each station 10 and 12 each contain a drip basin 130 and spill sensor  
 15 132 to alert the operator to potential leaks.

An alcohol supply 134 controlled by a valve S3 can supply alcohol to the  
 channel pumps 32 after rinsing steps to assist in removing water from the endoscope  
 channels.

20  
 Flow rates in the supply lines 30 can be monitored via the channel pumps 32  
 and the pressure sensors 42. The channels pumps 32 are peristaltic pumps which  
 supply a constant flow. If one of the pressure sensors 42 detects too high a pressure the  
 associated pump 32 cycles off. The flow rate of the pump 32 and its percentage on  
 25 time provide a reasonable indication of the flow rate in an associated line 30. These  
 flow rates are monitored during the process to check for blockages in any of the  
 endoscope channels. Alternatively, the decay in the pressure from the time the pump  
 32 cycles off can also be used to estimate the flow rate, with faster decay rates being  
 associated with higher flow rates.

30  
 A more accurate measurement of flow rate in an individual channel may be  
 desirable to detect more subtle blockages. A metering tube 136 having a plurality of  
 level indicating sensors 138 fluidly connects to the inputs of the channel pumps 32.

One preferred sensor arrangement provides a reference connection at a low point in the metering tube and a plurality of sensors 138 arranged vertically thereabove. By passing a current from the reference point through the fluid to the sensors 138 it can be determined which sensors 138 are immersed and therefore determine the level within the metering tube 136. Other level sensing techniques can be applied here. By shutting valve S1 and opening a vent valve S7 the channel pumps 32 draw exclusively from the metering tube. The amount of fluid being drawn can be very accurately determined based upon the sensors 138. By running each channel pump in isolation the flow therethrough can be accurately determined based upon the time and the volume of fluid emptied from the metering tube.

In addition to the input and output devices described above, all of the electrical and electromechanical devices shown are operatively connected to and controlled by the control system 20. Specifically, and without limitation, the switches and sensors 42, 59, 76, 84, 90, 98, 114, 116, 132 and 136 provide input I to the microcontroller 28 which controls the decontamination and other machine operations in accordance therewith. For example, the microcontroller 28 includes outputs O that are operatively connected to the pumps 32, 38, 70, 72, 88, 94, 100, 110, the valves S1-S7, and the heater 80 to control these devices for effective decontamination and other operations.

Turning also to FIG. 3, an endoscope 200 has a head part 202, in which openings 204 and 206 are formed, and in which, during normal use of the endoscope 200, an air/water valve and a suction valve are arranged. A flexible insertion tube 208 is attached to the head part 202, in which tube a combined air/water channel 210 and a combined suction/biopsy channel 212 are accommodated.

A separate air channel 213 and water channel 214, which at the location of a joining point 216 merge into the air/water channel 210, are arranged in the head part 202. Furthermore, a separate suction channel 217 and biopsy channel 218, which at the location of the joining point 220 merge into the suction/biopsy channel 212, are accommodated in the head part 202.

In the head part 202, the air channel 213 and the water channel 214 open into the opening 204 for the air/water valve. The suction channel 217 opens into the opening 206 for the suction valve. Furthermore, a flexible feed hose 222 connects to the head part 202 and accommodates channels 213', 214' and 217' which via the  
5 openings 204 and 206, are connected to the air channel 213, the water channel 214 and the suction channel 217, respectively. In practice, the feed hose 222 is also referred to as the light-conductor casing.

The mutually connecting channels 213 and 213', 214 and 214', 217 and 217' will  
10 be referred to below overall as the air channel 213, the water channel 214 and the suction channel 217.

A connection 226 for the air channel 213, connections 228 and 228a for the water channel 214 and a connection 230 for the suction channel 217 are arranged on the  
15 end section 224 (also referred to as the light conductor connector) of the flexible hose 222. When the connection 226 is in use, connection 228a is closed off. A connection 232 for the biopsy channel 218 is arranged on the head part 202.

A channel separator 240 is shown inserted into the openings 204 and 206. It  
20 comprises a body 242, and plug members 244 and 246 which occlude respectively openings 204 and 206. A coaxial insert 248 on the plug member 244 extends inwardly of the opening 204 and terminates in an annular flange 250 which occludes a portion of the opening 204 to separate channel 213 from channel 214. By connecting the lines 30 to the openings 226, 228, 228a, 230 and 232, liquid for cleaning and disinfection can be  
25 flowed through the endoscope channels 213, 214, 217 and 218 and out of a distal tip 252 of the endoscope 200 via channels 210 and 212. The channel separator 240 ensures the such liquid flows all the way through the endoscope 200 without leaking out of openings 204 and 206 and isolates channels 213 and 214 from each other so that each has its own independent flow path. One of skill in the art will appreciate that  
30 various endoscopes having differing arrangements of channels and openings will likely require modifications in the channel separator 240 to accommodate such differences while occluding ports in the head 202 and keeping channels separated from each other

so that each channel can be flushed independently of the other channels. Otherwise a blockage in one channel might merely redirect flow to a connected unblocked channel.

5 A leakage port 254 on the end section 224 leads into an interior portion 256 of the endoscope 200 and is used to check for the physical integrity thereof, namely to ensure that no leakage has formed between any of the channels and the interior 256 or from the exterior to the interior 256.

10 Turning also now to FIGS. 4 and 5, a concentration monitor 300 monitors concentration of the disinfecting solution circulating through the basin 14a or 14b. An inlet valve 302 connects through its A port 304 to the circulating fluid downstream of the main circulation pump 70. Its B port 306 leads either to waste or back to the basin 14a or 14b such as through the air gap 56. Its C port 308 leads to a sampling valve 310 through its A port 312. Its B port 314 leads to a piston chamber 316 liquid side 318 and 15 its C port 320 leads to a drain valve 322. A piston 324 operates within the piston chamber 316. With the piston 324 all the way down, the liquid side 318 should have a volume of about 15 to 50 ml or larger to promote flotation of entrained bubbles. A reservoir size of 30 to 35 ml has been shown to work well with OPA. Its diameter should be 13 to 26 mm, ore preferably 18 to 20 mm, to promote bubble flotation. A 20 larger size could also be used. The air pump 38 connects to an air side 326 of the piston chamber 316 through an air valve 328 at its A port 330. The air valve 328 B port 332 connects to the piston chamber air side 326 and its C port 334 opens to atmosphere.

25 On the drain valve 322, its A port 336 leads to the sampling valve 310, its B port 338 to drain and its C port 340 to an inlet 342 of a cuvette 344. An outlet 346 of the cuvette 344 leads preferably to drain, but can lead to a sample collection container (not shown) for further periodic testing of the fluid, or back to the basin 14a or 14b.

30 The cuvette 344 is preferably holds a sample of about 5 ml and is 2mm wide having optical grade glass or quartz side windows 348 through which light may pass for spectroscopically measuring a property of the liquid in the cuvette 344. A UV lamp 350 passes light through a filter 352, collimator 353 and a beam-splitter 354 passes a

portion of the light through the cuvette 344 and liquid therein to a first detector 356 and reflects another portion of the light toward a second, reference, detector 358. The lamp emits in the 150 nm to 600 nm range and the filter passes light at 254 nm for measuring concentration of OPA. Other wavelengths would be appropriate for different solutions and are easily determined by those of skill in the art. A controller 360 ties into the valves, lamp and detectors to control the operation thereof, and it itself links into the main controller 28.

In use, a sample of the circulating liquid is drawn in through the inlet valve 302 and sample valve 310 into the liquid side 318 of the piston chamber 316. The air side 326 of the piston chamber 316 is open to atmosphere through the air valve 328 to allow the piston 324 to move as the liquid enters the piston chamber 316. After filling the liquid side 318 and moving the piston all the way down, the liquid is allowed to rest to allow any bubbles therein to float to the surface. For an OPA solution a rest time of 30 to 40 seconds should be sufficient. Then the sample valve 310 and air valve 328 are cycled allowing air to enter the air side 326 driving the piston upwards and expelling the bubbles out of the liquid side 318 toward the drain valve 322 and out its B port 338 to drain. After a time period sufficient to expel the bubbles, the drain valve 322 is cycled to direct the liquid out of the A port 336 to the cuvette 344. Alternatively, the drain valve 322 can be omitted with the bubbles being passed out through the cuvette 344 by passing a sufficient quantity of liquid therethrough to obtain a bubble free sample within the cuvette 344. With a sample in the cuvette 344, light is passed through to spectroscopically measure the concentration of the OPA or other component therein.

25

The entire cleaning and sterilization cycle in detail comprises the following steps.

### **Step 1. Open the Lid**

Pressing a foot pedal (not shown) opens the basin lid 16a. There is a separate foot pedal for each side. If pressure is removed from the foot pedal, the lid motion stops.

30

**Step 2. Position and connect the endoscope**

The insertion tube 208 of the endoscope 200 is inserted into the helical circulation tube 64. The end section 224 and head section 202 of the endoscope 200 are situated within the basin 14a, with the feed hose 222 coiled within the basin 14a  
5 with as wide a diameter as possible.

The flush lines 30, preferably color-coded, are attached, one apiece, to the endoscope openings 226, 228, 228a, 230 and 232. The air line 112 is also connected to the connector 254. A guide located on the on the station 10 provides a reference for the color-coded connections.

**10 Step 3. Identify the user, endoscope, and specialist to the system**

Depending on the customer-selectable configuration, the control system 20 may prompt for user code, patient ID, endoscope code, and/or specialist code. This information may be entered manually (through the touch screen) or automatically such as by using an attached barcode wand (not shown).

**15 Step 4. Close the basin lid**

Closing the lid 16a preferably requires the user to press a hardware button and a touch-screen 22 button simultaneously (not shown) to provides a fail-safe mechanism for preventing the user's hands from being caught or pinched by the closing basin lid 16a. If either the hardware button or software button is released  
20 while the lid 16a is in the process of closing the motion stops.

**Step 5. Start Program**

The user presses a touch-screen 22 button to begin the washing / disinfection process.

**Step 6. Pressurize the endoscope body and Measure the Leak Rate**

25 The air pump is started and pressure within the endoscope body is monitored. When pressure reaches 250 mbar, the pump is stopped, and the pressure is allowed to stabilize for 6 seconds. If pressure has not reached 250 mbar in 45 seconds the

program is stopped and the user is notified of the leak. If pressure drops to less than 100 mbar during the 6-second stabilization period, the program is stopped and the user is notified of the condition.

5 Once the pressure has stabilized, the pressure drop is monitored over the course of 60 seconds. If pressure drops more than 10 mbar within 60 seconds, the program is stopped and the user is notified of the condition. If the pressure drop is less than 10 mbar in 60 seconds, the system continues with the next step. A slight positive pressure is held within the endoscope body during the rest of the process to prevent fluids from leaking in.

#### 10 **Step 7. Check Connections**

A second leak test checks the adequacy of connection to the various ports 226, 228, 228a, 230, 232 and the proper placement of the channel separator 240. A quantity of water is admitted to the basin 14a so as to submerge the distal end of the endoscope in the helical tube 64. Valve S1 is closed and valve S7 opened and the pumps 32 are run in reverse to draw a vacuum and to ultimately draw liquid into the endoscope channels 210 and 212. The pressure sensors 42 are monitored to make sure that the pressure in any one channel does not drop by more than a predetermined amount in a given time frame. If it does, it likely indicates that one of the connections was not made correctly and air is leaking into the channel. In any event, in the presence of an unacceptable pressure drop the control system 20 will cancel the cycle and indicate a likely faulty connection, preferably with an indication of which channel failed.

#### **PRE-RINSE**

25 The purpose of this step is to flush water through the channels to remove waste material prior to washing and disinfecting the endoscope 200.

**Step 8. Fill basin**

The basin 14a is filled with filtered water and the water level is detected by the pressure sensor 59 below the basin 14a.

**Step 9. Pump water through channels**

- 5 The water is pumped via the pumps 32 through the interior of the channels 213, 214, 217, 218, 210 and 212 directly to the drain 74. This water is not recirculated around the exterior surfaces of the endoscope 200 during this stage.

**Step 10. Drain**

- 10 As the water is being pumped through the channels, the drain pump 72 is activated to ensure that the basin 14a is also emptied. The drain pump 72 will be turned off when the drain switch 76 detects that the drain process is complete.

**Step 11. Blow air through channels**

During the drain process sterile air is blown via the air pump 38 through all endoscope channels simultaneously to minimize potential carryover.

**15 WASH****Step 12. Fill basin**

The basin 14a is filled with warm water (35 °C ). Water temperature is controlled by controlling the mix of heated and unheated water. The water level is detected by the pressure sensor 59.

**20 Step 13. Add detergent**

The system adds enzymatic detergent to the water circulating in the system by means of the peristaltic metering pump 88. The volume is controlled by controlling the delivery time, pump speed, and inner diameter of the peristaltic pump tubing.

**Step 14. Circulate wash solution**

The detergent solution is actively pumped throughout the internal channels and over the surface of the endoscope 200 for a predetermined time period, typically of from one to five minutes, preferably about three minutes, by the channel pumps 32 and the external circulation pump 70. The inline heater 80 keeps the temperature at about 35 °C.

**Step 15. Start block test**

After the detergent solution has been circulating for a couple of minutes, the flow rate through the channels is measured. If the flow rate through any channel is less than a predetermined rate for that channel, the channel is identified as blocked, the program is stopped, and the user is notified of the condition. The peristaltic pumps 32 are run at their predetermined flow rates and cycle off in the presence of unacceptably high pressure readings at the associated pressure sensor 42. If a channel is blocked the predetermined flow rate will trigger the pressure sensor 42 indicating the inability to adequately pass this flow rate. As the pumps 32 are peristaltic, their operating flow rate combined with the percentage of time they are cycled off due to pressure will provide the actual flow rate. The flow rate can also be estimated based upon the decay of the pressure from the time the pump 32 cycles off.

**Step 16. Drain**

The drain pump 72 is activated to remove the detergent solution from the basin 14a and the channels. The drain pump 72 turns off when the drain level sensor 76 indicates that drainage is complete.

**Step 17. Blow air**

During the drain process sterile air is blown through all endoscope channels simultaneously to minimize potential carryover.

## **RINSE**

### **Step 18. Fill basin**

The basin 14a is filled with warm water (35 °C). Water temperature is controlled by controlling the mix of heated and unheated water. The water level is detected by  
5 the pressure sensor 59.

### **Step 19. Rinse**

The rinse water is circulated within the endoscope channels (via the channel pumps 32) and over the exterior of the endoscope 200 (via the circulation pump 70 and the sprinkler arm 60) for 1 minute.

### 10 **Step 20. Continue Block test**

As rinse water is pumped through the channels, the flow rate through the channels is measured and if it falls below the predetermined rate for any given channel, the channel is identified as blocked, the program is stopped, and the user is notified of the condition.

### 15 **Step 21. Drain**

The drain pump is activated to remove the rinse water from the basin and the channels.

### **Step 22. Blow air**

20 During the drain process sterile air is blown through all endoscope channels simultaneously to minimize potential carryover.

### **Step 23. Repeat rinse**

Steps 18 through 22 are repeated to ensure maximum rinsing of enzymatic detergent solution from the surfaces of the endoscope and the basin.

## **DISINFECT**

### **Step 24. Fill basin**

The basin 14a is filled with very warm water (53 °C). Water temperature is controlled by controlling the mix of heated and unheated water. The water level is  
5 detected by the pressure sensor 59. During the filling process, the channel pumps 32 are off in order to ensure that the disinfectant in the basin is at the in-use concentration prior to circulating through the channels.

### **Step 25. Add disinfectant**

A measured volume of disinfectant 92, preferably CIDEX OPA orthophalaldehyde  
10 concentrate solution, available from Advanced Sterilization Products division Ethicon, Inc., Irvine, CA, is drawn from the disinfectant metering tube 96 and delivered into the water in the basin 14a via the metering pump 100. The disinfectant volume is controlled by the positioning of the fill sensor 98 relative to the bottom of the dispensing tube. The metering tube 96 is filled until the upper  
15 level switch detects liquid. Disinfectant 92 is drawn from the metering tube 96 until the level of the disinfectant in the metering tube is just below the tip of the dispensing tube. After the necessary volume is dispensed, the metering tube 96 is refilled from the bottle of disinfectant 92. Disinfectant is not added until the basin is filled, so that in case of a water supply problem, concentrated disinfectant is not  
20 left on the endoscope with no water to rinse it. While the disinfectant is being added, the channel pumps 32 are off in order to insure that the disinfectant in the basin is at the in-use concentration prior to circulating through the channels.

### **Step 26. Disinfect**

The in-use disinfectant solution is actively pumped throughout the internal channels  
25 and over the surface of the endoscope, ideally for a minimum of 5 minutes, by the channel pumps and the external circulation pump. The temperature is controlled by the in-line heater 80 to about 52.5 °C. During this process a sample of the circulating liquid is taken and tested for proper concentration using the concentration monitor 300. If the concentration is low, additional sterilant can be  
30 added and the timer for this step reset.

**Step 27. Flow check**

During the disinfection process, flow through each endoscope channel is verified by timing the delivering a measured quantity of solution through the channel. Valve S1 is shut, and valve S7 opened, and in turn each channel pump 32 delivers a  
 5 predetermined volume to its associated channel from the metering tube 136. This volume and the time it takes to deliver provides a very accurate flow rate through the channel. Anomalies in the flow rate from what is expected for a channel of that diameter and length are flagged by the control system 20 and the process stopped.

**Step 28. Continue Block Test**

10 As disinfectant in-use solution is pumped through the channels, the flow rate through the channels is also measured as in Step 15.

**Step 29. Drain**

The drain pump 72 is activated to remove the disinfectant solution from the basin and the channels.

**Step 30. Blow air**

15 During the drain process sterile air is blown through all endoscope channels simultaneously to minimize potential carryover.

**FINAL RINSE****Step 31. Fill basin**

20 The basin is filled with sterile warm water (45 °C ) that has been passed through a 0.2 μ filter.

**Step 32. Rinse**

25 The rinse water is circulated within the endoscope channels (via the channel pumps 32) and over the exterior of the endoscope (via the circulation pump 70 and the sprinkler arm 60) for 1 minute.

**Step 33. Continue Block test**

As rinse water is pumped through the channels, the flow rate through the channels is measured as in Step 15.

**Step 34. Drain**

- 5 The drain pump 72 is activated to remove the rinse water from the basin and the channels.

**Step 35. Blow air**

During the drain process sterile air is blown through all endoscope channels simultaneously to minimize potential carryover.

10 **Step 36. Repeat rinse**

Steps 31 through 35 are repeated two more times (a total of 3 post-disinfection rinses) to ensure maximum reduction of disinfectant residuals from the endoscope 200 and surfaces of the reprocessor.

**FINAL LEAK TEST**

15 **Step 37. Pressurize the endoscope body and measure leak rate**

Repeat Step 6.

**Step 38. Indicate program completion**

The successful completion of the program is indicated on the touch screen.

**Step 39. De-pressurize the endoscope**

- 20 From the time of program completion to the time at which the lid is opened, pressure within the endoscope body is normalized to atmospheric pressure by opening the vent valve S5 for 10 seconds every minute.

**Step 40. Identify the user**

Depending on customer-selected configuration, the system will prevent the lid from being opened until a valid user identification code is entered.

**Step 41. Store program information**

- 5 Information about the completed program, including the user ID, endoscope ID, specialist ID, and patient ID are stored along with the sensor data obtained throughout the program.

**Step 42. Print program record**

- 10 If a printer is connected to the system, and if requested by the user, a record of the disinfection program will be printed.

**Step 43. Remove the endoscope**

- 15 Once a valid user identification code has been entered, the lid may be opened (using the foot pedal as in step 1, above). The endoscope is then disconnected from the flush lines 30 and removed from the basin 14a. The lid can then be closed using both the hardware and software buttons as described in step 4, above.

- 20 The invention has been described with reference to the preferred embodiments. Obviously, modifications and alterations will occur to others upon reading and understanding the preceding detailed description. It is intended that the invention be construed as including all such modifications and alterations insofar as they come within the scope of the appended claims or the equivalents thereof.

## WHAT IS CLAIMED IS:

1. An endoscope processor having a solution measuring system, the measuring system comprising:
  - 5 a cuvette for holding a sample of the solution;
  - a light source for passing a light through the cuvette and the sample;
  - a light sensing mechanism for sensing light passing through the cuvette and the sample;
  - a reservoir for receiving a quantity of the solution containing bubbles;
  - 10 a pump associated with the reservoir for pumping a quantity of solution out of the reservoir;
  - a first path from the reservoir;
  - a second path from the reservoir to the cuvette;
  - a control system associated with the pump and programmed to first direct the
  - 15 pump to move a portion of the quantity of fluid in the reservoir out through the first path, whereby to drive bubbles therein out of the reservoir, and then to direct a sample of the liquid into the cuvette.
2. An endoscope processor according to claim 1 wherein the solution comprises an aldehyde.
- 20 3. An endoscope processor according to claim 2 wherein the solution comprises orthophthalaldehyde.
4. An endoscope processor according to claim 1 wherein a light path through the sample in the cuvette is between 1 mm and 5 mm.

5. An endoscope processor according to claim 4 wherein a light path through the sample in the cuvette is between 1 mm and 3 mm.

6. An endoscope processor according to claim 1 wherein the first path leaves the reservoir from an upper portion thereof.

5 7. An endoscope processor according to claim 1 wherein the first path is the same as the second path and the bubbles are removed through the cuvette.

8. An endoscope processor according to claim 7 wherein the control system is programmed to pump fluid out of the reservoir for a time period sufficient to pump substantially all of the bubbles in the solution out through the cuvette  
10 whereby to leave a quantity of solution in the cuvette substantially free of bubbles.

9. An endoscope processor according to claim 1 wherein the control system is programmed to delay pumping of fluid out of the reservoir for a time period after the reservoir is filled sufficient to allow the bubbles in the solution to float to the surface.

15 10. A method for measuring a property of a solution to be applied to an endoscope in an endoscope processing procedure using the endoscope processor according to any one of claims 1 to 6, the method comprising:

collecting a quantity of the solution in the reservoir;

10 directing a portion of the solution out from the reservoir through the first path to carry bubbles in the solution out of the reservoir;

then directing a sample of the solution from the reservoir to the cuvette;

measuring the property of the solution in the sample in the cuvette by passing light through the cuvette and the sample and reading said light passing through said light and said sample.

25 11. A method according to claim 10 wherein the bubbles are removed through the cuvette.

12. A method according to claim 10 wherein after collecting a quantity of the solution in the reservoir there is a delay sufficient to allow the bubbles in the solution to float to the surface before performing the step of directing a portion of the solution out from the reservoir through the first path.

5 13. A method according to any one of claims 10 to 12 wherein the property of the solution being measured is the level of sterilant therein.

**FIG. 1**

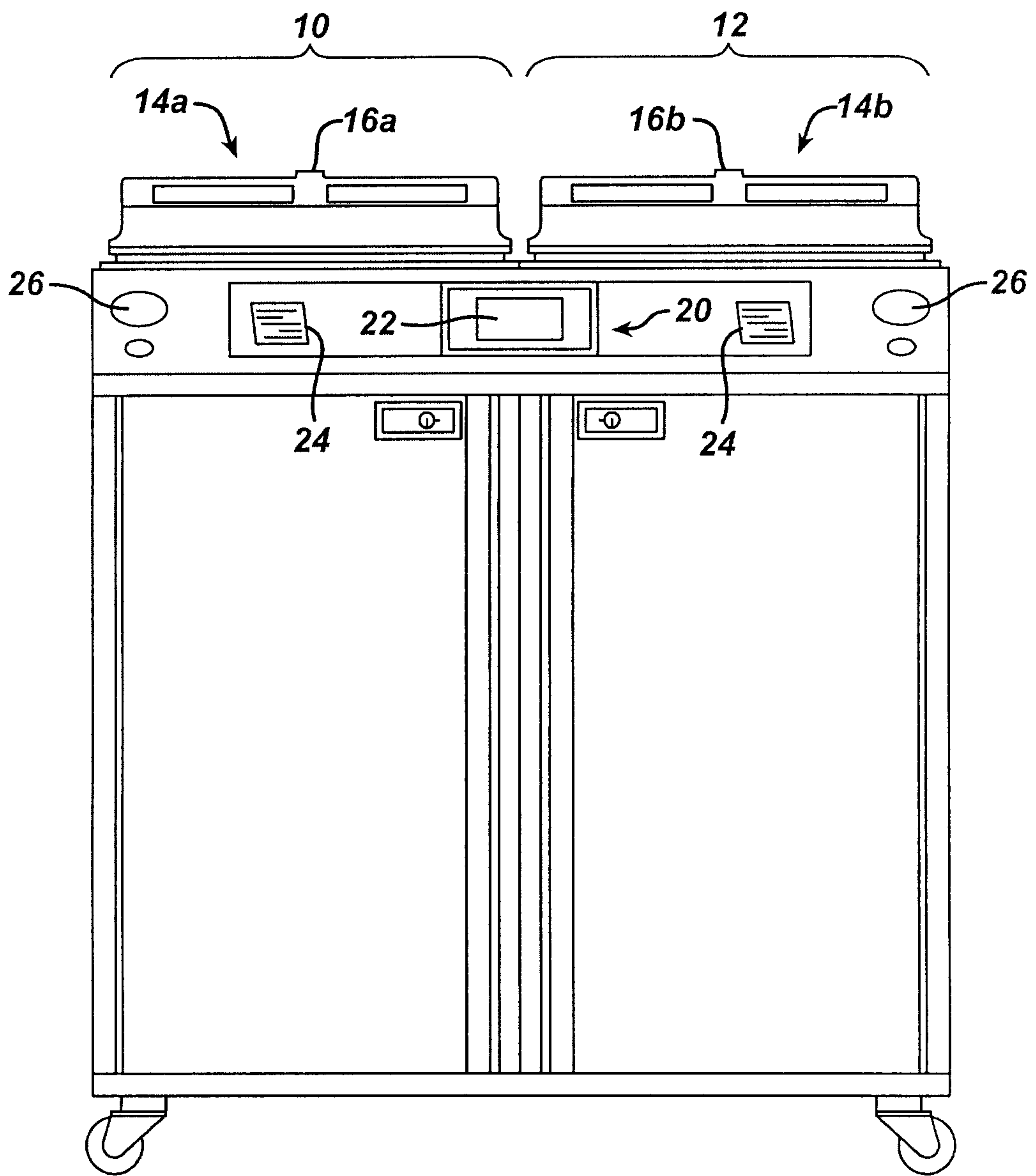
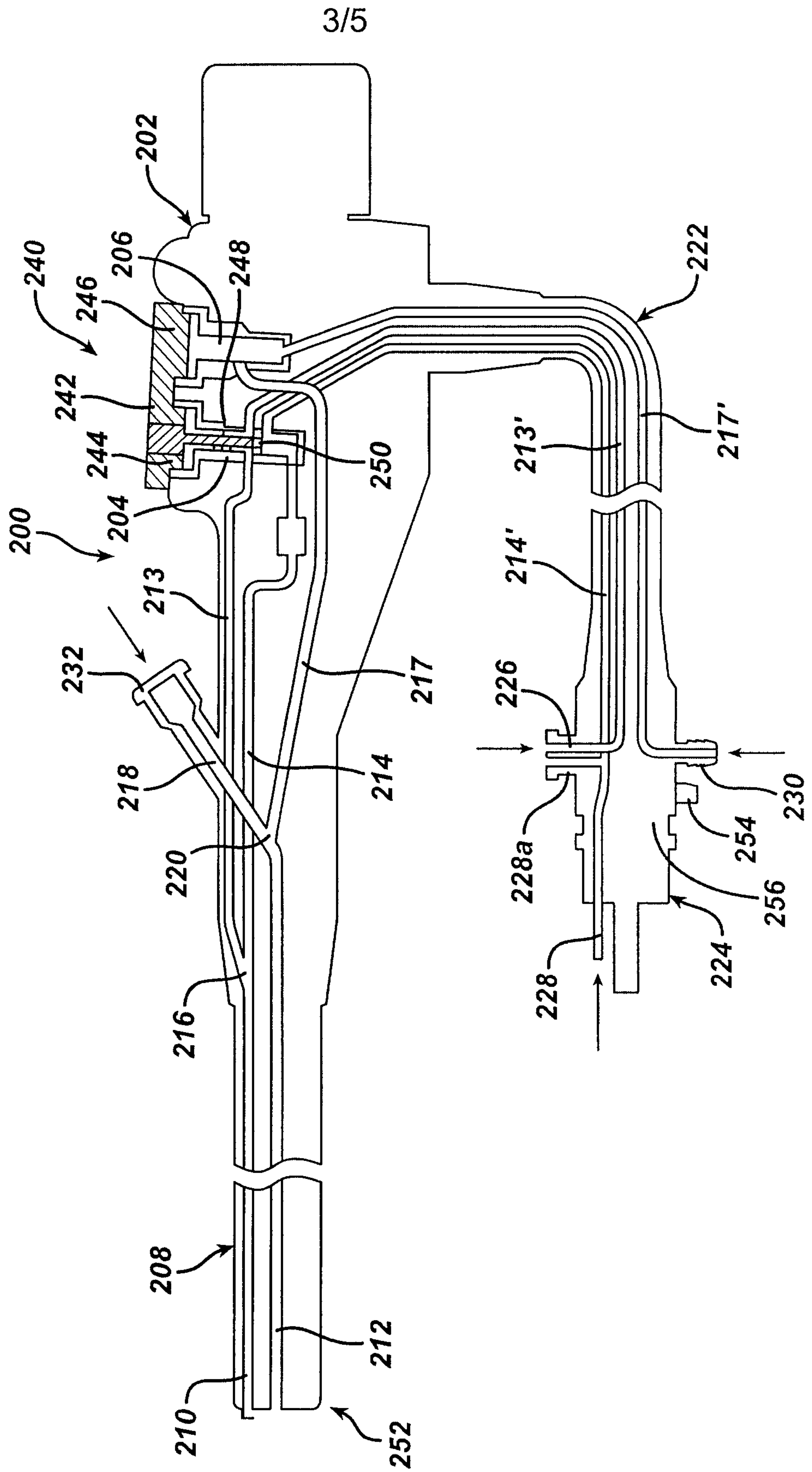


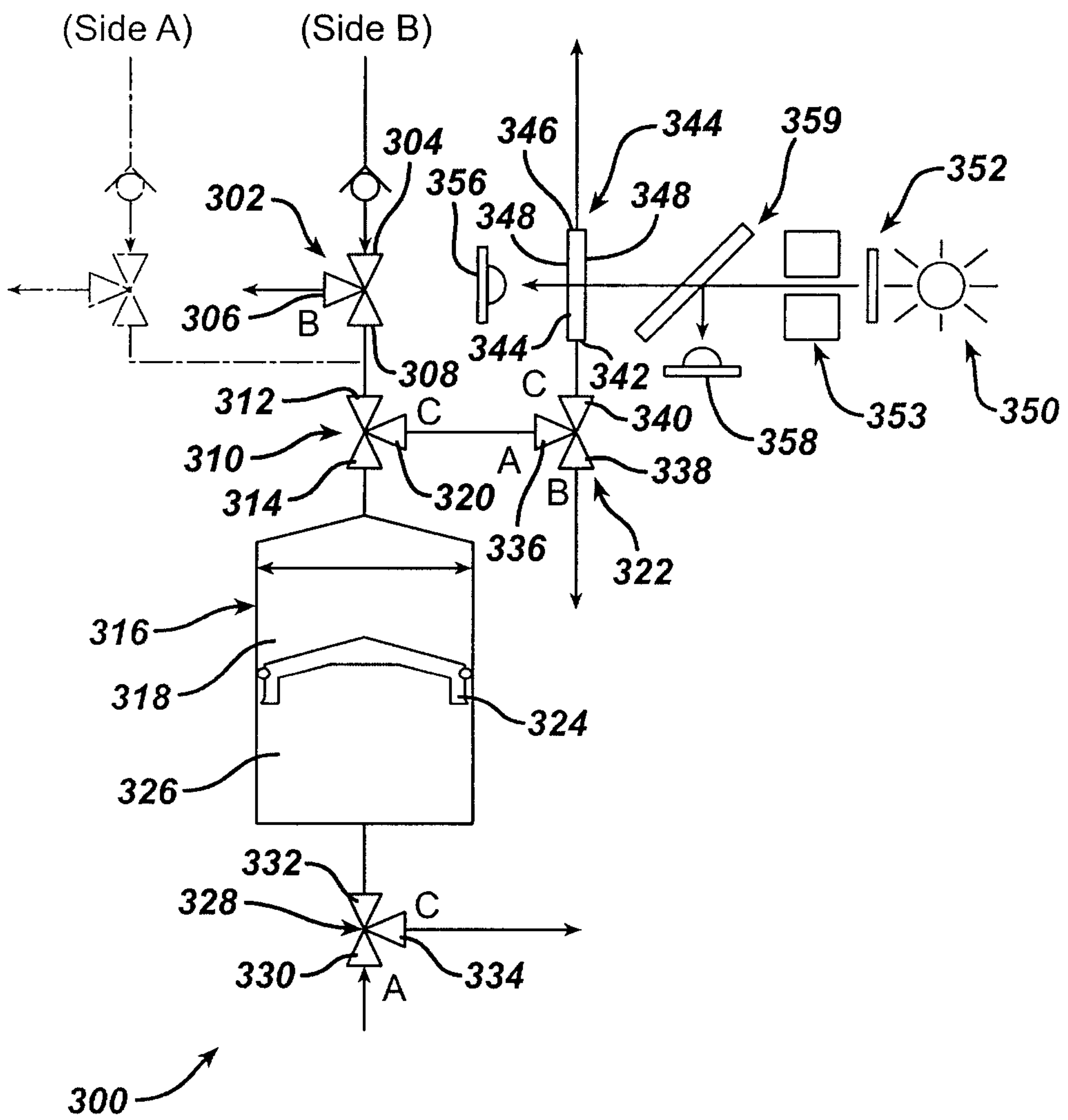


FIG. 3

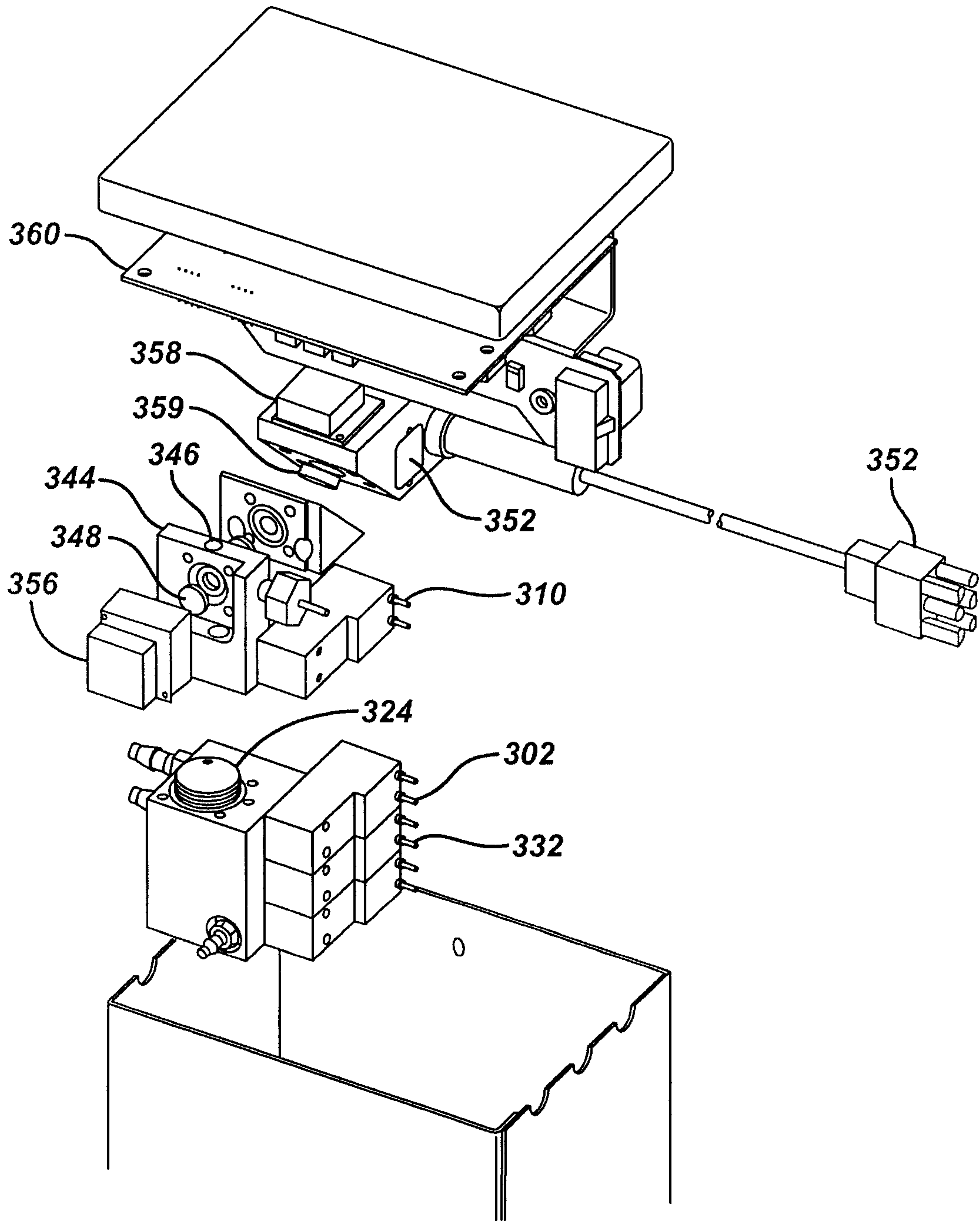


**FIG. 4**

OPAC In-use Solution



5/5



**FIG. 5**

