

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization

International Bureau



(10) International Publication Number

WO 2013/022491 A1

(43) International Publication Date  
14 February 2013 (14.02.2013)

(51) International Patent Classification:  
*B01D 33/15 (2006.01)*

(74) Agent: ALTMANN, Curtis, R.; Arnold & Porter LLP,  
555 12th Street, N.W., Washington, D.C. 20004 (US).

(21) International Application Number:  
PCT/US2012/030930

(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

(22) International Filing Date:  
28 March 2012 (28.03.2012)

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, ZA, ZM, ZW).

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:  
61/468,377 28 March 2011 (28.03.2011) US

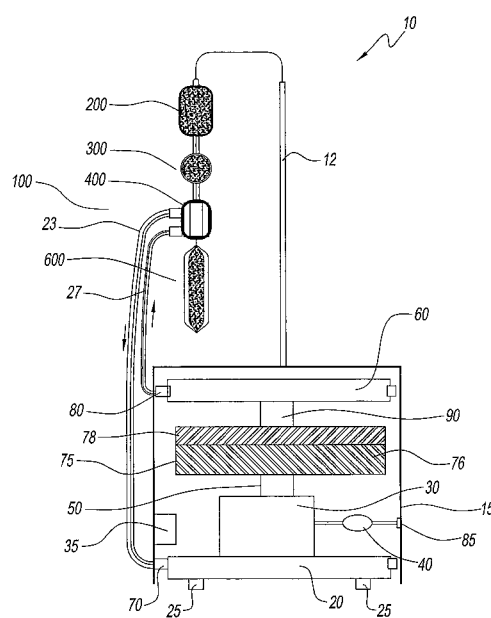
(71) Applicant (for all designated States except US): NEW HEALTH SCIENCES, INC. [US/US]; 6903 Rockledge Drive, Suite 230, Bethesda, MD 20817-1818 (US).

(72) Inventors; and

(75) Inventors/Applicants (for US only): VERNUCCI, Paul [US/US]; 7 Dyer Street, Billerica, MA 01862 (US). YOSHIDA, Tatsuro [JP/US]; 1763 Commonwealth Avenue, West Newton, MA 02465 (US).

[Continued on next page]

(54) Title: METHOD AND SYSTEM FOR REMOVING OXYGEN AND CARBON DIOXIDE DURING RED CELL BLOOD PROCESSING USING AN INERT CARRIER GAS AND MANIFOLD ASSEMBLY



(57) Abstract: A portable assembly for processing red blood cells RBCs including a disposable blood collection set including a blood bag, an anaerobic storage bag and an oxygen and/or oxygen and carbon dioxide depletion device disposed between the blood collection bag and anaerobic storage bag. The portable assembly further provides for a gas circulation device in fluid communication with the oxygen or oxygen and carbon dioxide depletion device, The gas circulation device includes a pressure source that is able circulate flushing gas through the depletion device as RBCs pass from the blood collection bag, through the depletion device and into the anaerobic storage bag.

FIG. 1a



SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, **Published:**  
GW, ML, MR, NE, SN, TD, TG).

— *with international search report (Art. 21(3))*

METHOD AND SYSTEM FOR REMOVING OXYGEN AND CARBON DIOXIDE  
DURING RED CELL BLOOD PROCESSING USING AN INERT CARRIER GAS AND  
MANIFOLD ASSEMBLY

5

BACKGROUND OF THE INVENTION

1. Field of the Invention

10 The present disclosure relates to a portable blood treatment manifold assembly. More, particularly, the present disclosure relates to a portable blood treatment manifold assembly for leukoreduction and oxygen and/or carbon dioxide depletion of blood in preparation for blood storage and/or transfusion to a recipient.

15 2. Background of the Art

20 The supplies of liquid blood in are currently limited by storage systems used in conventional blood storage practice. Using current systems, stored blood expires after about 42 days of refrigerated storage at a temperature above freezing (i.e. 1-6°C) as packed blood cell preparations. Red blood cells (RBCs) may be concentrated from whole blood with separation of the liquid blood component (plasma). Expired blood cannot be used and is discarded.

25 There are periodic shortages of blood that occur due to donation fluctuation, emergencies and other factors. The logistics of blood supply and distribution impact the military, especially during times of combat and remote hospitals or medical facilities making blood processing or transfusions very difficult. Accordingly, there is a need to be able to rapidly prepare RBCs for storage or for transfusions in remote locations.

30 Storage of frozen blood is known in the art but such frozen blood has limitations. For a number of years, frozen blood has been used by blood banks and the military for certain high-demand and rare types of blood. However, frozen blood is difficult to

handle. It must be thawed which makes it impractical for emergency situations. Once blood is thawed, it must be used within 24 hours. United States Patent No. 6,413,713 to Serebrennikov is directed to a method of storing blood at temperatures below 0° C.

5 U.S. Patent No. 4,769,318 to Hamasaki *et al.* and U.S. Patent No. 4,880,786 to Sasakawa *et al.* are directed to additive solutions for blood preservation and activation. U.S. Patent No. 5,624,794 to Bitensky *et al.*, U.S. Patent No. 6,162,396 to Bitensky *et al.*, and U.S. Patent No. 5,476,764 are directed to the storage of red blood cells under oxygen-depleted conditions. U.S. Patent No. 5,789,151 to Bitensky *et al.* is directed to 10 blood storage additive solutions.

Additive solutions for blood preservation and activation are known in the art. For example, Rejuvesol (available from enCyte Corp., Braintree, MA) is added to blood after cold storage (i.e., 4 °C) just prior to transfusion or prior to freezing (i.e., at -80°C with 15 glycerol) for extended storage. U.S. Patent No. 6,447,987 to Hess *et al.* is directed to additive solutions for the refrigerated storage of human red blood cells.

In light of current technology, there is a need for a portable and cost effective 20 apparatus and methodology for the preparation of RBCs that removes leukocytes and oxygen and/or carbon dioxide in advance of transfusion or in preparation for anaerobic storage.

## SUMMARY OF THE INVENTION

25 Accordingly, the present disclosure provides a system that is capable of removing oxygen and/or carbon dioxide and/or leukocytes from RBCs in advance of transfusion or for further storage in an anaerobic environment.

The present disclosure also provides for a system and methodology for the 30 preparation of RBCs in advance of transfusion or for further storage in an anaerobic environment.

It is a further object of the present disclosure to provide a stand-alone portable system that has an oxygen or an oxygen/carbon dioxide depletion (OCDD) device that removes oxygen or oxygen and or carbon dioxide from RBCs passing through the 5 device. The OCDD device operates with a gas exchange system that pumps gas into the device through which RBCs that first passes through an oxygen or oxygen/carbon dioxide (OCDD) device to remove oxygen or oxygen/carbon dioxide from such RBCs. The RBCs are thereby depleted of oxygen or oxygen/carbon dioxide and deposited in a blood storage bag for extended storage or storage in advance of transfusion.

10

It is a still further object of the present disclosure to provide a stand alone portable system that pumps gas into the device through which RBCs pas through a leukoreduction filter and an oxygen and/or carbon dioxide (OCDD) device to remove leukocytes and oxygen or oxygen/carbon dioxide from such RBCs, respectively. The 15 RBCs are thereby free of leukocytes and depleted of oxygen or oxygen/carbon dioxide and deposited in a blood storage bag for extended storage or storage in advance of transfusion.

It is still a further object of the present disclosure to provide a standalone portable 20 system that circulates oxygen depleted and or/carbon dioxide adjusted air air or inert gas mixtures through an OCDD device to remove such gases from RBCs flowing through the filter in preparation for anaerobic storage or transfusion. Such system contains oxygen, carbon dioxide and/or partial pressure sensors between an inlet 25 manifold that receives oxygen and/or carbon dioxide rich air or inert gas from an OCDD device and an outlet manifold. The sensors monitor and regulate oxygen and or carbon dioxide levels in air or inert gas mixtures received in the outlet manifold and monitor oxygen and carbon dioxide partial pressure of filtered gas that is pumped pumped back to OCDD device.

30 It is still a further object of the present disclosure to provide a standalone portable

system that reduces leukocytes and circulates oxygen and/or carbon dioxide adjusted air or inert gas mixtures through an OCDD device to remove such gases from RBCs in preparation for anaerobic storage or transfusion. Such system contains oxygen, carbon dioxide and/or partial pressure sensors between an inlet manifold that receives oxygen and/or carbon dioxide rich air or inert gas mixtures from an OCDD device and an outlet manifold that feeds oxygen and carbon dioxide depleted air or inert gas mixtures back to the OCDD device. The sensors monitor and regulate oxygen and or carbon dioxide levels in gas received in the outlet manifold and monitor oxygen and carbon dioxide partial pressure of gas that is pumped back to OCDD device.

10

A portable assembly for processing red blood cells RBCs including a disposable blood collection set including a blood bag, an anaerobic storage bag and an oxygen and/or oxygen and carbon dioxide depletion device disposed between the blood collection bag and anaerobic storage bag. The portable assembly further provides for a gas circulation device in fluid communication with the oxygen or oxygen and carbon dioxide depletion device, The gas circulation device includes a pressure source that is able circulate flushing gas through the depletion device as RBCs pass from the blood collection bag, through the depletion device and into the anaerobic storage bag.

20 A portable assembly for processing red blood cells (RBCs) including an oxygen or oxygen and carbon dioxide depletion (OCDD) device. The OCDD device includes a cartridge having an inlet and an outlet and a plurality of hollow fibers disposed between the inlet and the outlet for transporting RBCs through the OCDD device. The plurality of hollow fibers are surrounded by a continuous space. The portable assembly includes a gas exchange device in fluid communication with the OCDD device. The gas exchange device includes a pressure source that is able to circulate a flushing gas through the continuous space and remove oxygen and/or carbon dioxide from RBCs passing through the OCDD device.

25 30 These and other objects and advantages of the present invention and

equivalents thereof, are achieved by the methods and compositions of the present invention described herein and manifest in the appended claims.

#### BRIEF DESCRIPTION OF THE DRAWINGS

5

Fig. 1a illustrates a portable blood processing system according to the present disclosure;

10 Fig 1b illustrates an alternative embodiment of the present disclosure in which red blood cells are processed using a load cell;

Figure 1c illustrates the OCDD device of the embodiment of Fig. 1b directly connected to the processing system;

15 Fig. 1d illustrates a collection system that incorporates a flow regulator according to the embodiment of Fig. 1b;

Fig. 1e illustrates a collection system that incorporates a leukoreduction filter with an OCDD device;

20

Figs. 2a through 2c illustrate a leukoreduction filter incorporated into an OCDD device according to the embodiment of Fig. 1e;

Fig. 2d illustrates an OCDD device of the embodiment of Fig. 1a;

25

Fig. 3 illustrates an OCDD device according to a further embodiment of the present disclosure having OCCD device, leukoreduction filter and plasma separation device in a unitary structure.

30

## DETAILED DESCRIPTION OF THE DISCLOSURE

Referring to Fig. 1, a stand alone blood processing system is shown and referenced using reference numeral 10. System 10 includes a housing 15 and supports a blood collection and depletion system 100 (hereinafter "collection system 100"). Collection system 100 includes a blood bag 200, a leukoreduction filter 300, an oxygen and/or carbon dioxide depletion (OCDD) device 400 and an anaerobic storage bag 600. Device 400 is able to deplete oxygen or alternatively, oxygen and carbon dioxide from gas from RBCs. Collection system 100 is suspended within system 10 to enable convenient movement and transport of blood preparation processes in locations that may be remote from a standard hospital or clinical setting. The orientation of system 100, permits RBCs in blood bag 200 to flow under the force of gravity to anaerobic storage bag 600. Although a single collection system 100 is shown, stand 12 of housing 15 could carry as many as ten or more such systems for processing. Housing 15 includes a gas circulation device including a pressure source such as a pump 30 or a vacuum or a pressurized container, a valve/pressure regulator 40 and further components that will be discussed further that enable gas to circulate and pass through OCDD device 400. Inlet 410 and outlet 415 (Fig. 2d) that area connected to tubing 427 and 426, respectively.

20

Collection system 100 includes a blood bag 200 that contains RBCs that have been collected from whole blood. Generally, whole blood is collected from a donor using traditional methods and processed using centrifugation to separate plasma and RBCs. Blood bag 200 is a standard blood collection bag. RBCs are collected in a blood bag 200 that may contain an additive. An additive solution, such as, for example, OFAS3, includes adenine, dextrose, mannitol,  $\text{NaH}_2\text{PO}_4$ , and optionally  $\text{NaCl}$  and/or  $\text{NH}_4\text{Cl}$ . Additive solution OFAS3 preferably comprises ingredients having the following ranges: about 0.5-4.0 mmole/liter of adenine, about 50-150 mmole/liter of dextrose, about 20-70 mmole/liter of mannitol, about 0-100 mmole/liter of  $\text{NaCl}$ , about 2-20 mmole/liter of  $\text{NaH}_2\text{PO}_4$ , and about 0-30 mmole/liter  $\text{NH}_4\text{Cl}$ . Preferably, OFAS3, has an adjusted pH from about 5.5-7.5 and includes about 2 mmole/liter adenine, about 110

mmole/liter dextrose, about 55 mmole/liter NaCl, and about 12 mmole/liter NaH<sub>2</sub>PO<sub>4</sub> and an adjusted pH of about 6.5. Additives such as SAGM, PAGG-SM, AS-1, AS-3, AS-5, SOLX, MAPS, PAGG-GM or any additive approved for blood storage may also be used in this system.

5

RBCs contained in blood bag 200 flow under the force of gravity to leukoreduction filter 300 and through OCDD device 400. Leukoreduction is the process of removing white blood cells from the whole blood or RBCs. Leukocytes in blood products can cause immunosuppressive effects and can pre-dispose patients to an 10 increased risk of viruses, fevers, and have deleterious effects on RBCs. Leukoreduction reduces RBC storage lesions, reduces primary alloimmunization and reduces total number of transfusion reactions.

15 The process of leukoreducing RBCs preferably occurs after the RBCs have been separated from the plasma and can occur before or after removal of oxygen and carbon dioxide have been removed from the RBCs. In either case, leukoreduction should occur before storage of RBCs and anaerobic storage bag 600.

20 Referring to FIGS. 2a, 2b, and 2c leukoreduction filter 300 is incorporated into OCDD device 500. OCDD device 500 includes a cartridge 505, an inlet 510, a leukoreduction filter 520, a plurality of hollow fibers 530, and a fiber support 540 to hold the plurality of hollow fibers 530. OCDD device 500 also includes an outlet 515 for 25 passage of RBCs. Leukoreduction filter 520 is preferably a fibrous or a felt-like filtering material that captures leukocytes, prior to such leukocytes travelling through plurality of hollow fibers 530. Fiber support 540 supports the plurality of hollow fibers 530 in a vertical configuration and may be made from a material such as polyurethane or a similar material. Either whole blood or pRBC flow through filter 520 during leukoreduction process. OCDD device 500 is in communication with gas from pump 30 via an inlet 524 and an outlet 528.

30

OCDD cartridge 500 contains approximately 5000 fibers for the passage of

RBCs. More or fewer fibers may be used to generate a sufficient surface area for gas exchange to reduce the oxygen and/or carbon dioxide concentrations to the desired levels. Plurality of hollow fibers 530 are for the purpose of removing oxygen or oxygen and carbon dioxide from RBC and will be discussed further below. Gas spaces 550, 5 outside of hollow fibers and inside of cartridge 505, that surround plurality of hollow fibers 530 and are filled with a carrier gas. Gas permeable material or porous materials of plurality of hollow fibers 530 enable oxygen and carbon dioxide to pass from RBCs to carrier gas when such gas is circulated through OCDD device 500. OCDD device 500 depletes, O<sub>2</sub> and CO<sub>2</sub>, or O<sub>2</sub>, or CO<sub>2</sub> alone, or O<sub>2</sub> with specific levels of CO<sub>2</sub> by 10 supplying an appropriate composition of flushing gas. Gases appropriate for depletion for use in OCDD devices are any inert gasses that will not cause harm to the RBCs or blood recipient, for example, Ar, He, N<sub>2</sub>, Ar/CO<sub>2</sub>, He/CO<sub>2</sub> or N<sub>2</sub>/CO<sub>2</sub>.

15 RBCs flow into OCDD device 500 to be depleted of oxygen or oxygen and carbon dioxide. OCDD device 500 reduces the degree of RBC hemoglobin oxygen saturation levels to less than 3 % and the carbon dioxide partial pressure to less than 50 Torr at 37 °C. OCDD device 500 is a combination oxygen and carbon dioxide filter that removes oxygen and carbon dioxide from RBCs to enhance the storage life of such RBCs and promotes optimal transfusion. OCDD device 500 is used with housing 115 20 and stand 12 of Fig. 1e and contains same components as embodiment of Fig. 1a.

25 Alternatively, as shown in Fig. 2d, an OCDD device 400 does not contain the leukoreduction capability and is only capable of depleting oxygen or oxygen and carbon dioxide from RBCs passing there through. Fig. 2d illustrates an OCDD device 400 that has an inlet 410 for the entry of RBCs, an outlet 415 for the passage of RBCs, and a plurality of fibers 430 through which such RBCs pass to be deleted of oxygen and/or carbon dioxide gas. OCDD device 400 also contains an entry port 424 for flushing gas and an exit port 428 for the egress of flushing gas and a plurality of spaces 450 that surround plurality of fibers 430 that are inside of cartridge 405 and where gas exchange 30 from RBCs to flushing gas occurs. The circulation of gas through OCDD device 400 via entry port 424, exit port 428 and plurality of spaces 450 ensures that the partial

pressure of oxygen and carbon dioxide in RBCs stored in bags 600 is at acceptable levels for optimal storage of RBCs.

Referring to Fig. 1a, again, housing 15 includes an inlet manifold 20, a pump 30, 5 an outlet manifold 60 and an inlet valve/pressure regulator 40. OCDD cartridge 400 is connected to inlet manifold 20 and outlet manifold 60 by tubing 27 and 13 or direct connections 128 and 124 (Fig. 1c) respectively. A first oxygen/carbon dioxide sensor 50 and a second oxygen/carbon dioxide sensor 90 are disposed between inlet manifold 20 and outlet manifold 60. System 10 is connectable to an AC outlet or other supply of 10 power for operation of pump 30. Alternatively, system 10 can connect to a battery for remote operation of system 10.

Housing 15 contains a disposable or re-usable sorbent cartridge 75 that is disposed between inlet manifold 20 and outlet manifold 60 to purify and air or inert gas 15 mixture that has passed through OCDD device 400. Sorbent cartridge 75 is a large cartridge that is preferably iron based or other inorganic and/or organic compound that can physically or chemically absorb oxygen or oxygen/carbon dioxide. Sorbent cartridge 75 contains an oxygen and/or a carbon dioxide sorbent 76. As an alternative to a large sorbent pack or organic and inorganic compounds, oxygen and carbon 20 dioxide can also be depleted from oxygen and carbon dioxide rich air or inert gas mixture by using membrane filters designed for gas separation, such as those found in nitrogen generator systems. In addition to oxygen or oxygen/carbon dioxide sorbent 76, sorbent cartridge 75 also includes activated charcoal filter 78 to absorb volatiles 25 produced by oxygen or oxygen/carbon dioxide sorbent. Charcoal filter 78 also includes a HEPA filter to remove any particulates.

System 10 also includes various sterilization filter sensor assemblies 70, 80 and 85. Sterilization filter sensor assembly 70 are disposed between tubing 23 and inlet manifold 20. Sterilization filter sensor assembly 80 is disposed between outlet manifold 30 60 and tubing 27. Filters 70 and 80 capture any pathogens and/or particulates that could enter gas flow between respective tubing and manifold and compromise filtration

and or purification of RBCs. Filters in 70 and 80 filter sensor assemblies monitor levels partial pressures of oxygen and carbon dioxide for an individual OCDD 400 (or 500). Sterilization filter 85 is disposed between external portion of housing 15 and inlet valve pressure regulator 40. Sterilization filter filter sensor assembly 85 monitors gas entering 5 pump 30. Filter in filter sensor assembly 85 capture pathogens and particulates between system 10 and ambient air or inert gas mixture and are also able to sense levels of oxygen, carbon dioxide, temperature and pressure and humidity. Filter sensor assemblies 70, 80 and 85 also function as sensors and are in communication with controller 35. Controller 35 is programmed with predetermined set points to monitor 10 and control concentration and flow ratre of oxygen and carbon dioxide, temperature, humidity and total pressure of the gas mixtures. Should levels not be appropriate, a warning signal, such as a light or alarm, informs an operator that sorbent cartridge, sterilization filter or HEPA filter should be replaced.

15 Housing 15 includes casters 25 to permit movement and positioning of system  
10. System 10 also includes a large sorbent cartridge 75 or hollow fiber gas separation module.

20 In operation, and as shown in Fig. 1, RBCs flow from collection bag 200 into OCDD cartridge directly or via leukoreduction filter. Flushing gas is simultaneously 25 circulated through OCDD cartridge 400. The flow of oxygen or oxygen/carbon dioxide adjusted gas and oxygen/carbon dioxide rich gas to and from OCDD cartridge 400 is carried by tube 27 and tube 23, respectively. Tube 23 is connected to inlet manifold 20 and tube 27 is connected to outlet manifold 60. Tube 23 is connected to inlet manifold 25 by a sterilization filter sensor assembly 70. Similarly, outlet manifold 60 is connected to tube 27 by sterilization filter 80.

30 After oxygen rich air or inert gas mixture egressing from OCDD device 400 via tubing 23, such air or inert gas mixture is received at inlet manifold 20, and pumped via pump 30 through sensor 50. Pump 30 operates to maintain gas flow through system 10. Pump 30 is preferably an electrically driven pump that regulates pressures and

flows. Pump 30 is connected to a valve 40, preferably a one way valve and pressure regulator that accepts ambient air or inert gas mixture at ambient pressure or insert gasses at elevated pressures. Sensor 50 and sensor 90 measure partial pressure of oxygen and carbon dioxide, in addition to gas partial pressure, temperature, flowrate 5 total pressure and humidity of the entire portable assembly. Air or inert gas is purified in cartridge 75 and returned to OCDD 400 to continue to depletion RBCs before such RBCs flow into anaerobic storage bag 600.

Figs. 1b through 1d show an alternative embodiment of a housing 115. Housing 10 115 contains similar gas exchange components as housing 15. Namely, housing 115 also contains an inlet manifold 20, a pump 30, an outlet manifold 60 and an inlet valve/pressure regulator 40 contained within housing 115. Housing 115 also contains a load cell 6 that is connected to bag 200 and a flow regulator valve 470. Load cell 6 measures the unit weight in bag 200 and communicates change in mass in bag to a 15 controller 35 that communicates with flow regulator valve 470 to monitor flow of RBCs through OCDD device 400. By monitoring change of mass of RBCs in bag 200, valve 470 can be adjusted to ensure that RBCs remain in OCDD device 400 for adequate oxygen or oxygen and carbon dioxide removal. Controller 35 is in electrical communication with load cell 6, flow regulator valve 470 and oxygen saturation sensor 20 475. Oxygen saturation sensor 475 measures oxygen saturation levels in RBCs. Controller 35 receives signals indicative of oxygen saturation levels and in turn sends signal to adjust flow regulator valve 470 to assure adequate oxygen depletion levels in RBCs. The several bags 200 (Fig. 1b) can be connected to housing 115 and be similarly equipped with a flow regulator valve 470 although only one flow regulator 470 25 is shown. Housing 115 has an outside surface to which OCDD devices 400 can be directly connected via couplings. By configuring OCDD devices 400, as shown in Figs. 1b through 1d, so that they are directly connected to housing 115 via couplings 124 and 128, the need for tubing of the embodiment of Fig. 1a is eliminated. The configuration of housing 115 can also be used with devices 500 that include leukoreduction 30 capability.

Referring to Fig. 3, a multifunction OCDD device 700 is a combination leukoreduction filter 710, OCDD device 720, in combination with a plasma separator 730. Multifunction OCDD device 700 eliminates the need for separation of the whole blood, received from donor, which is currently a separated by using a centrifuge. By 5 combining these three devices into a single device, the need for a separate centrifuge, a highly costly and cumbersome device, is eliminated. This embodiment contains a leukoreduction portion 710, a OCDD device 720 and a plasma separator 730. Plasma flows through port 740 to a further collection bag for further processing. Accordingly, in this embodiment, whole blood can be collected from a donor, leukocytes can be 10 removed, oxygen, or oxygen and carbon dioxide can be removed and plasma and platelets can be removed to pass RBCs through device. The RBCs are then deposited into collection bag 600 for storage or transfusion to a recipient. Multifunction OCDD 700 as part of collection system 100 and system 10 permit rapid transformation of whole blood to stored RBCs for immediate storage or transfusion to a recipient.

15 Although the present disclosure describes in detail certain embodiments, it is understood that variations and modifications exist known to those skilled in the art that are within the disclosure. Accordingly, the present disclosure is intended to encompass all such alternatives, modifications and variations that are within the scope of the disclosure as set forth in the disclosure. \*

20

25

30

We Claim:

1. A portable system for processing red blood cells (RBCs) comprising:
  - 5 a disposable blood collection set comprising a blood bag, an anaerobic storage bag and an oxygen and/or oxygen and carbon dioxide depletion device disposed between said blood collection bag and said anaerobic storage bag; and
  - 10 a gas circulation device in fluid communication with said oxygen/oxygen and carbon dioxide depletion device, said gas circulation device comprising a pressure source that is able circulate flushing gas through said depletion device as RBCs pass from the blood collection bag, through the depletion device and into the anaerobic storage bag.
- 15 2. The portable system according to claim 1, wherein said depletion device comprises a cartridge, a plurality of hollow fibers extending within the cartridge from an inlet to an outlet of the cartridge, and a contiguous space surrounding the plurality of hollow fibers for the passage of flushing gas,
- 20 3. The portable system of claim 2, wherein the plurality of hollow fibers comprise a gas permeable material or a porous material and are adapted for receiving and conveying red blood cells.
- 25 4. The portable system according to claim 1, further comprising a housing containing said gas circulation device, wherein said housing further comprises an inlet manifold for receiving flushing gas from said depletion device, an outlet manifold for providing flushing gas to said depletion device and a sorbent disposed between said pressure source and said outlet manifold for removing oxygen and/or oxygen and carbon dioxide from the flushing gas.

5. The portable system according to claim 4, further comprising a sensor disposed between said pressure source and said sorbent and a second sensor disposed between sorbent and said outlet manifold for detecting a level of oxygen and/or carbon dioxide in the flushing gas in the OCDD.

10. 6. The portable system according to claim 1, wherein said depletion device further comprises a leukoreduction filter.

15. 7. The portable system device according to claim 1, wherein said depletion device further comprises a leukoreduction filter and a plasma separator.

20. 8. The portable system according to claim 1, further comprising a load cell connected to said collection bag, a flow regulator valve disposed between said depletion device and the anaerobic storage bag, and a controller.

25. 9. The portable system according to claim 8, wherein said load cell measures the weight of RBCs in the collection bag, and said controller receives a signal indicative of the weight of the RBCs and communicates a signal to said flow regulator valve to adjust said flow regulator valve to restrict or facilitate the flow of RBCs and, thereby controlling the exposure of the RBCs to flushing gas in the depletion device.

30. 10. The portable system according to claim 8, further comprising a oxygen saturation sensor disposed between said depletion device and said flow regulator valve, wherein said controller receives a signal indicative of the level of oxygen saturation in the RBCs and sends a signal to said flow regulator valve to adjust said flow regulator valve to restrict or facilitate the flow of RBCs and thereby control the exposure of the RBCs to flushing gas in the depletion device.

11. The portable system according to claim 1, wherein said flushing gas comprises Ar, He, N<sub>2</sub>, Ar/CO<sub>2</sub>, He/CO<sub>2</sub> or N<sub>2</sub>/CO<sub>2</sub>, or any combination of inert gasses and/or CO<sub>2</sub>.

12. The portable system according to claim 1, further comprising a plurality of disposable blood collection sets connected to said housing and in fluid communication with said gas circulation device.

5

13. The portable system according to claim 1, wherein said pressure source is selected from the group consisting of a pump, a vacuum or a pressurized container.

10

14. A portable system for processing red blood cells (RBCs) comprising:

15

an oxygen or oxygen and carbon dioxide depletion (OCDD) device comprising a cartridge having an inlet and an outlet and a plurality of hollow fibers disposed between said inlet and said outlet for transporting RBCs through said OCDD device, wherein said plurality of hollow fibers are surrounded by a continuous space; and

20

a gas circulation device in fluid communication with said OCDD device; wherein said gas circulation device comprises a pressure source that is able to circulate a flushing gas through the continuous space and remove oxygen and/or carbon dioxide from RBCs passing through said OCDD device.

25

15. The portable system according to claim 14, further comprising a blood bag and an anaerobic storage bag, and wherein said blood bag is connected to said inlet and said anaerobic storage bag is connected to said outlet.

30

16. The portable system according to claim 14, further comprising a housing having an outer surface, said housing containing said gas circulation device, wherein said depletion device is directly coupled to said outer surface of said housing.

17. The portable assembly according to claim 14, further comprising a housing having an outer surface, said housing containing said gas exchange device and

a plurality of depletions devices, wherein said plurality of depletion devices are directly coupled to said outer surface of said housing.

18. The portable system according to claim 14, further comprising a load cell, a flow

5 regulator valve, and a controller, wherein said load cell measures the weight of RBCs in the blood bag and said controller receives a signal indicative of the weight of the RBCs and communicates a signal to said flow regulator valve to adjust said flow regulator valve to restrict or facilitate the flow of RBCs, thereby decreasing or increasing the exposure of the RBCs to flushing gases in the

10 depletion device.

19. The portable system according to claim 18, further comprising a oxygen saturation sensor disposed between said depletion device and said flow regulator valve, wherein said controller receives a signal indicative of the level of oxygen saturation in the RBCs and sends a signal to said flow regulator valve to adjust said flow regulator valve to restrict or facilitate the flow of RBCs and increase or decrease the exposure of the RBCs to flushing gas in the depletion device.

20. The portable system according to claim 17, wherein said housing further contains

an inlet manifold for receiving flushing gas from said depletion device, an outlet manifold for providing flushing gas to said depletion device and a sorbent disposed between said pressure source and said outlet manifold for removing oxygen or oxygen and carbon dioxide from the flushing gas.

25 21. The portable system according to claim 20, further comprising a system sensors on opposite sides of said sorbent to sense levels of oxygen or oxygen and carbon dioxide in flushing gas that is received and filtered in said sorbent.

22. The portable system according to claim 14, wherein said depletion device further 30 comprises a leukoreduction filter.

23. The portable assembly device according to claim 14, wherein said depletion device further comprises a leukoreduction filter and a plasma separator.

5 24. The portable system according to claim 20, further comprising a filter sensor assembly disposed between said depletion device and said inlet manifold and a filter sensor assembly disposed between said depletion device and said outlet manifold for monitoring levels of gas parameters of gas flowing through said inlet and said outlet.

10 25. The portable system of claim 24, wherein said filter sensor assembly disposed between said depletion device and said inlet manifold and said depletion device, said filter sensor assembly disposed between said depletion device and said outlet manifold and said system sensors measure partial pressure of oxygen, partial pressure of carbon dioxide, temperature, pressure humidity and gas flow rate of gas flowing through the system.

15 26. The portable system according to claim 14, wherein said flushing gas comprises Ar, He, N<sub>2</sub>, Ar/CO<sub>2</sub>, He/CO<sub>2</sub> or N<sub>2</sub>/CO<sub>2</sub>, or any combination of inert gasses and/or CO<sub>2</sub>.

20 27. The portable system according to claim 14, wherein said pressure source is selected from the group consisting of a pump, a vacuum or a pressurized container.

25

30

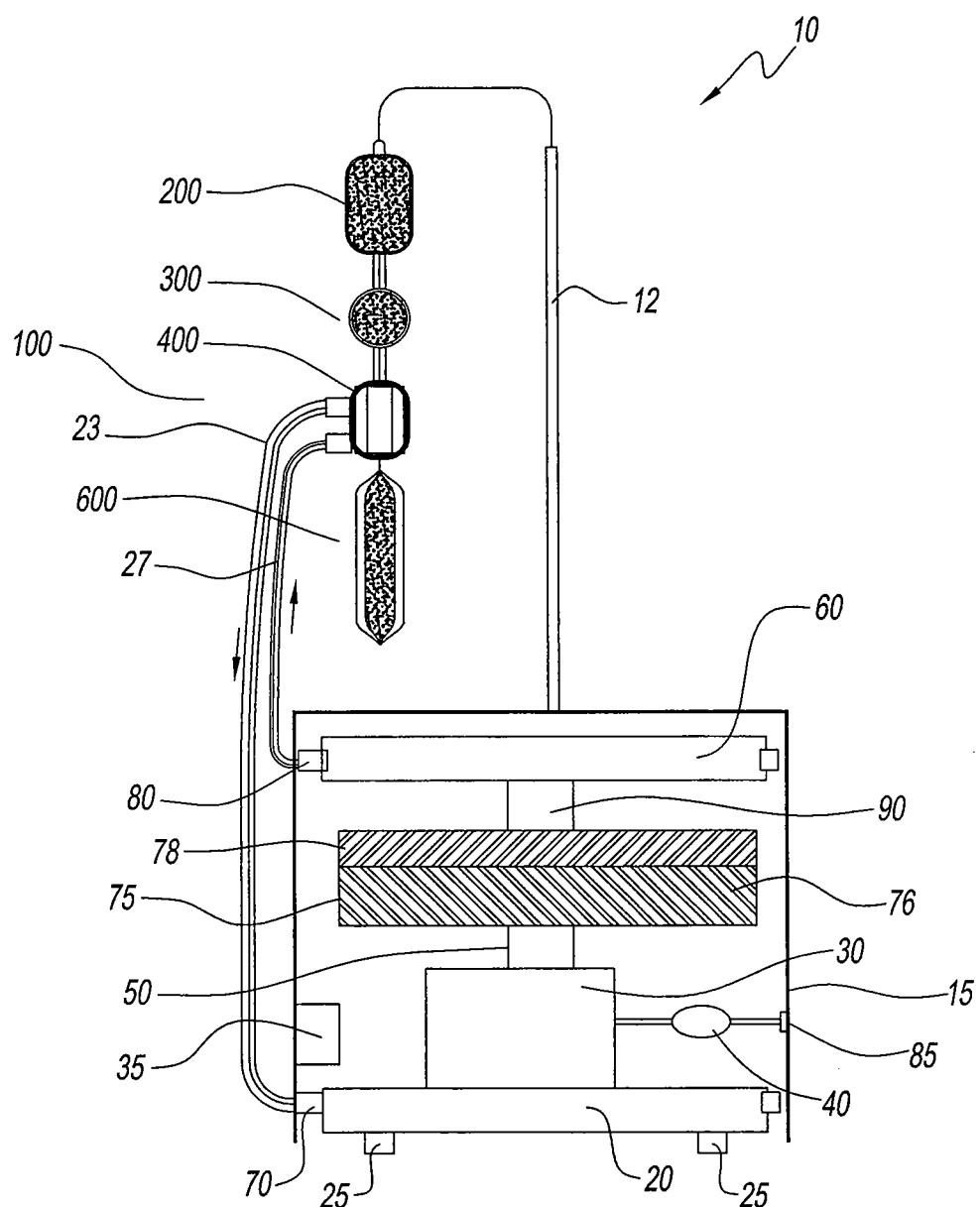


FIG. 1a

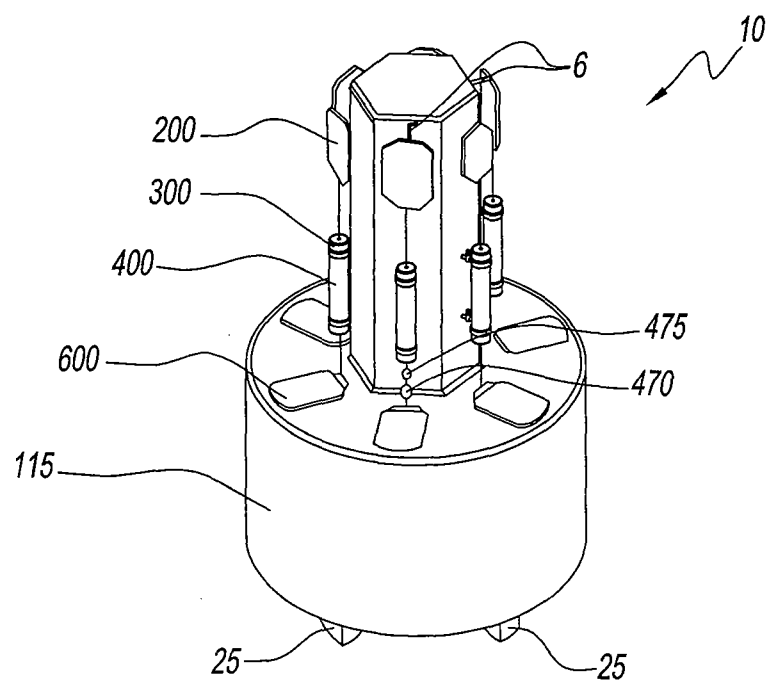


FIG. 1b

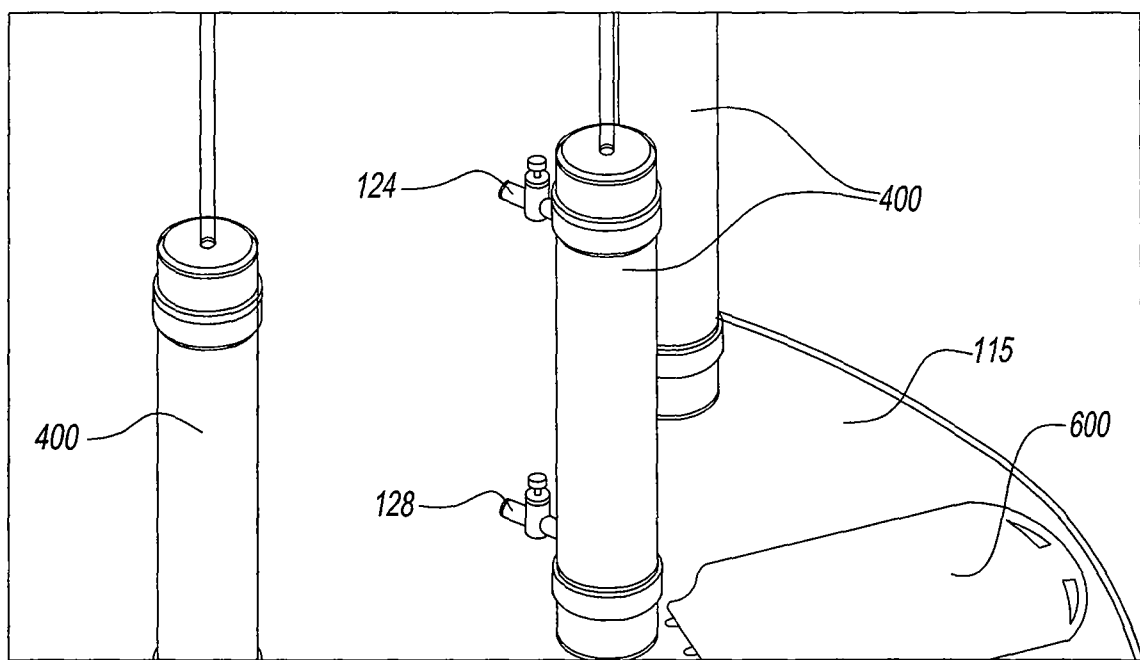
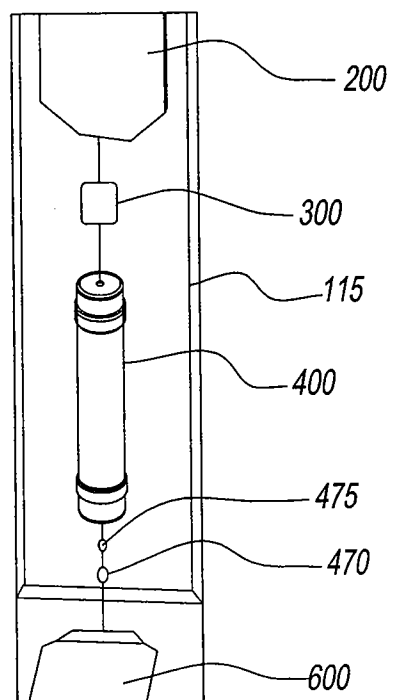


FIG. 1c



*FIG. 1d*

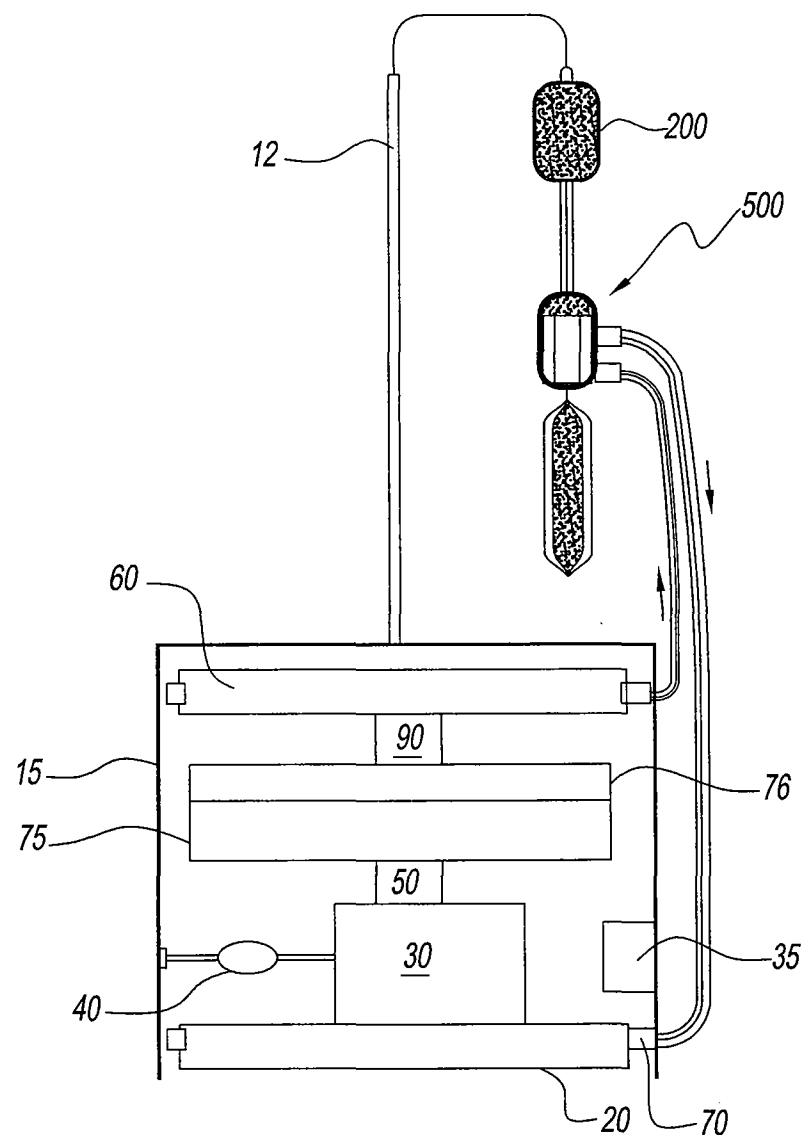


FIG. 1e

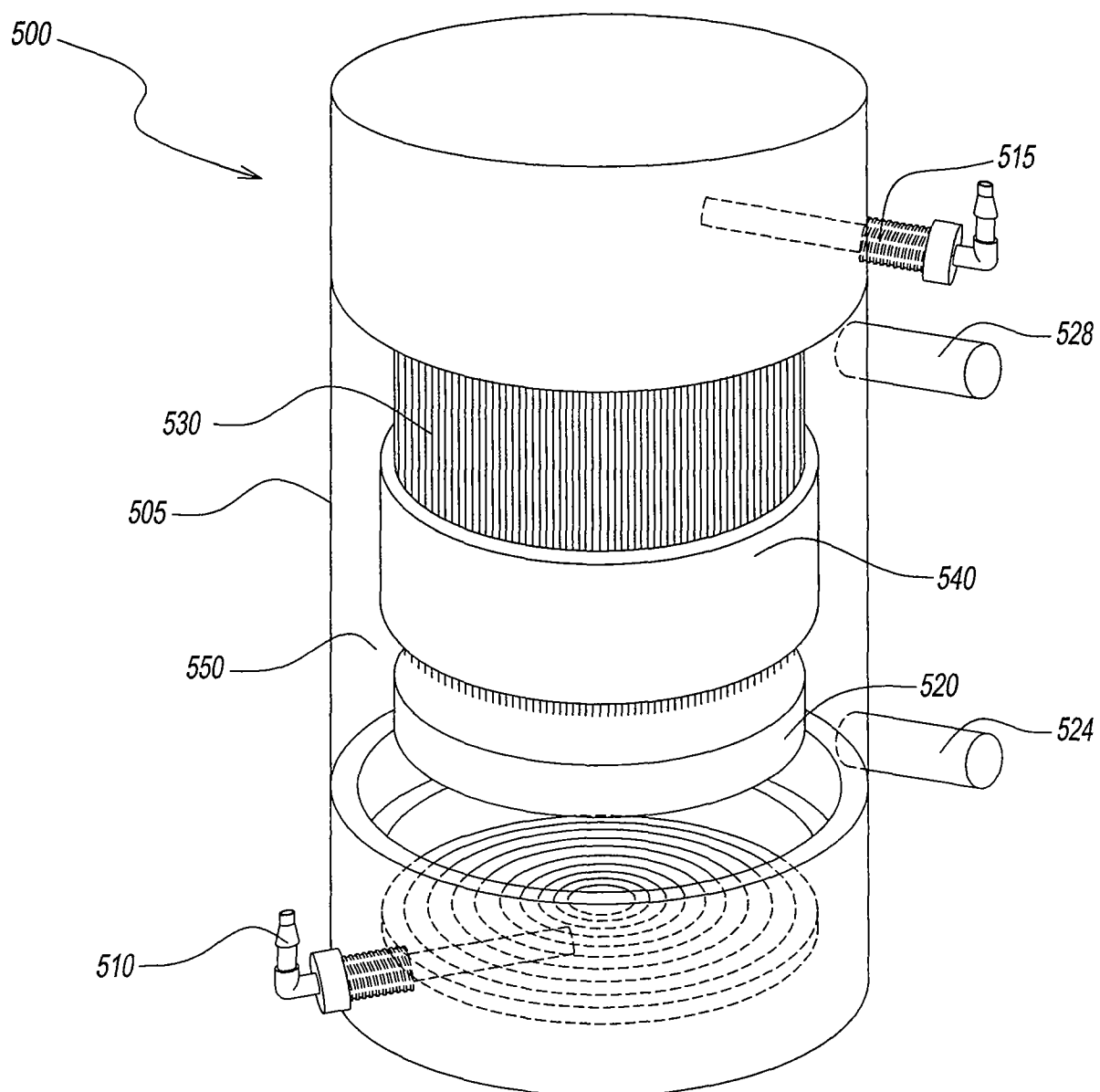


FIG. 2a

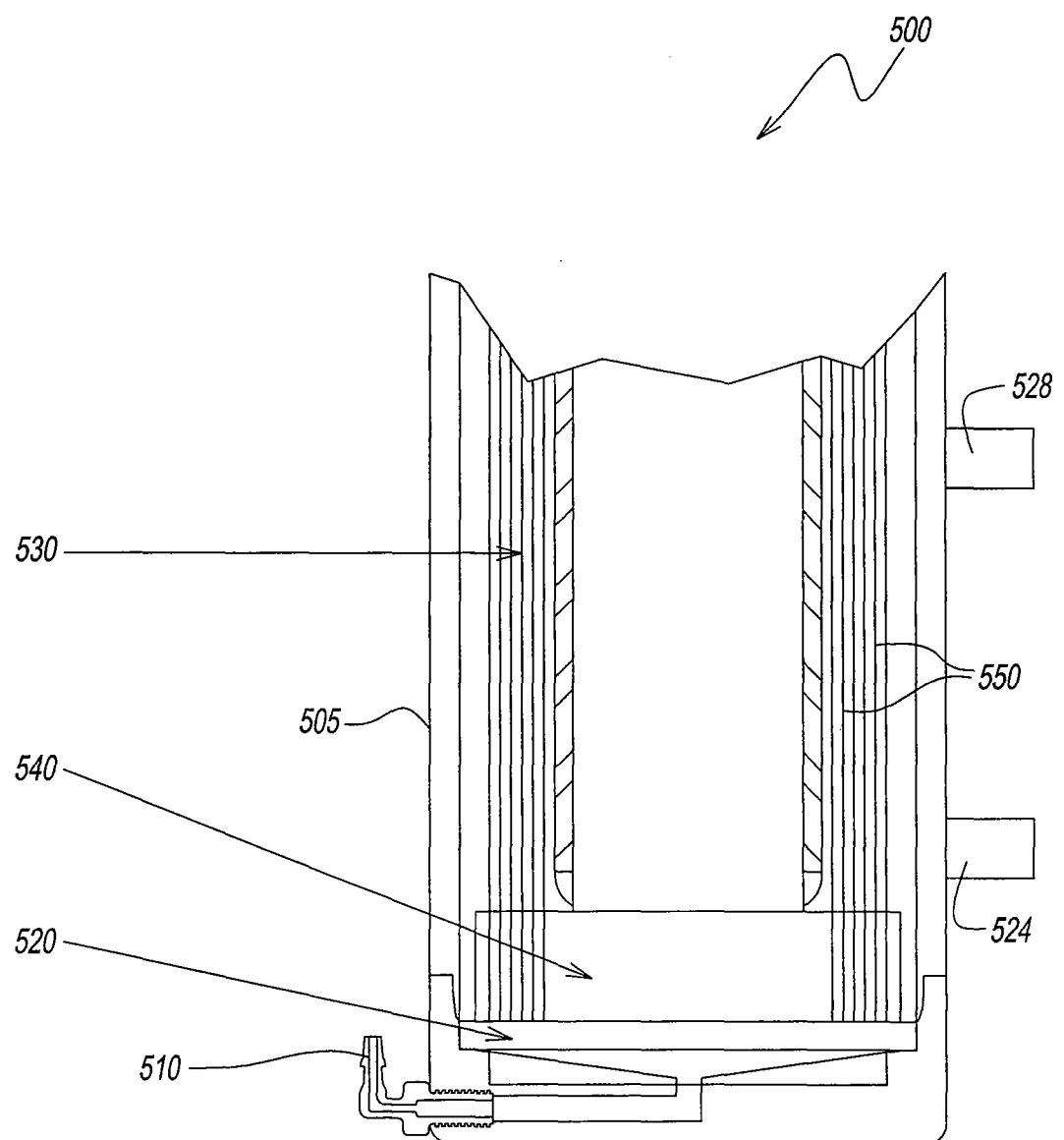


FIG. 2b

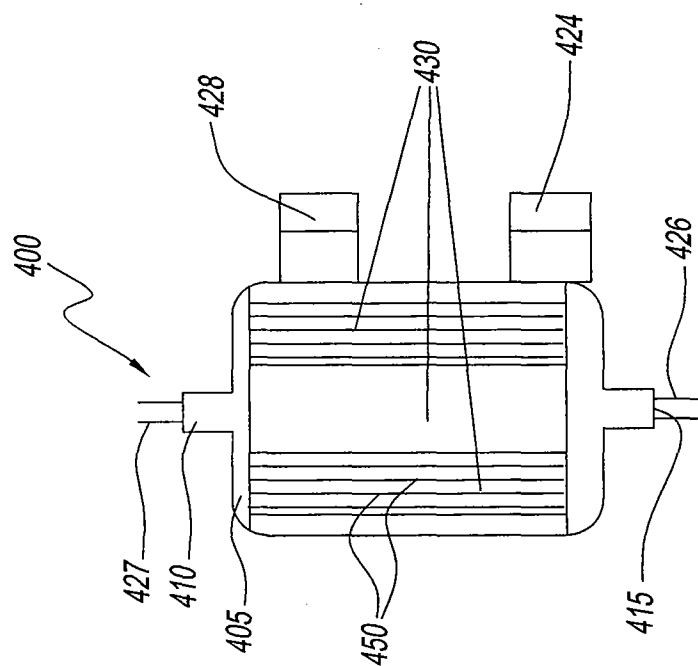


FIG. 2d

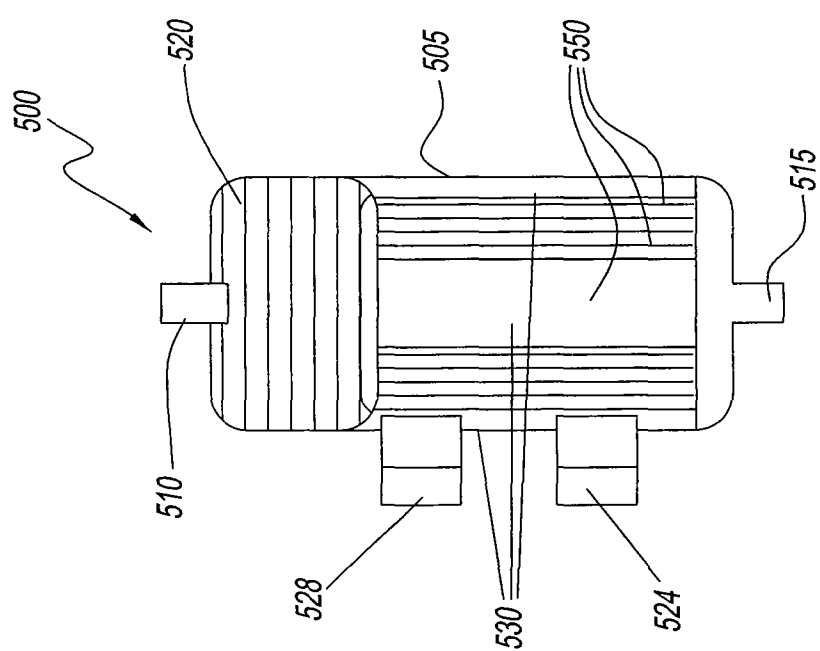


FIG. 2c

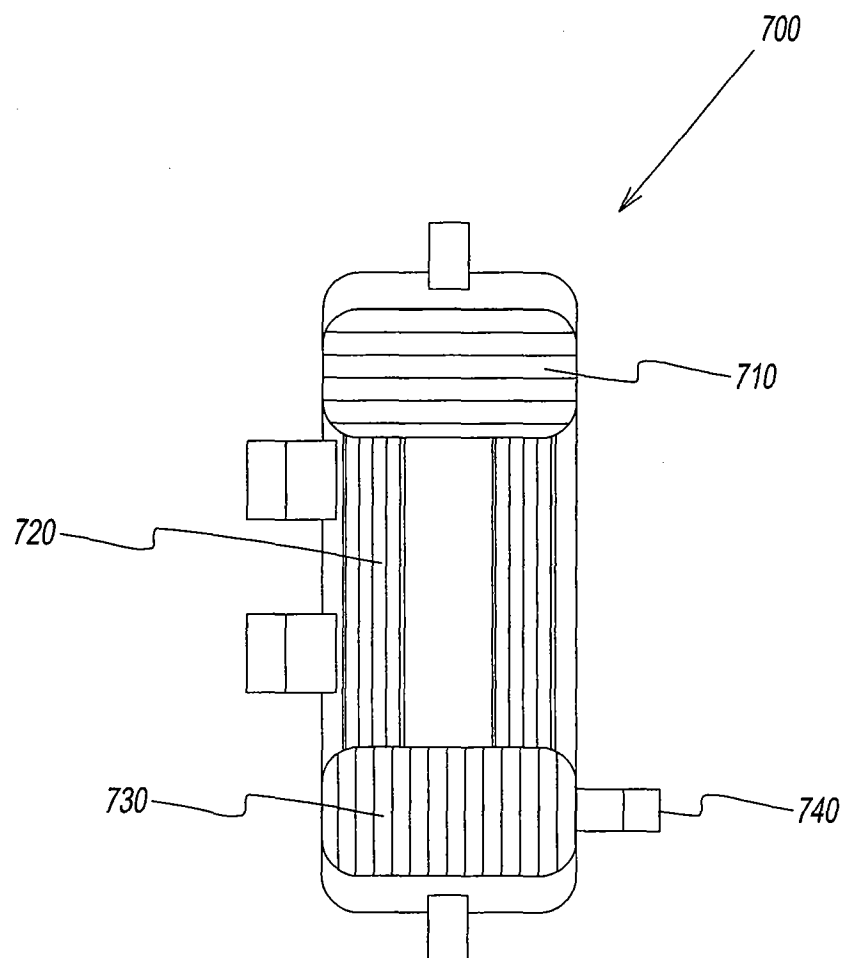


FIG. 3

**INTERNATIONAL SEARCH REPORT**

International application No. PCT/US 12/30930	
--	--

**A. CLASSIFICATION OF SUBJECT MATTER**  
IPC(8) - B01D 33/15 (2012.01)  
USPC - 210/782

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

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)  
IPC(8): B01D 33/15 (2012.01); USPC: 210/782

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched  
USPC: 210/782,645,194,512,1,787,512,3

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)  
Google Scholar, Google Patents, PubWEST (PGPB,USPT,EPAB,JPAB) (Red, blood, cell, collect, bag, carbon, dioxide, deplete, anaerobic, storage, oxygen, circulate, erythrocyte, housing, filter, valve, inlet, outlet, tubing, leukocyte, reduce, load cell, OCDD, nitrogen, argon, sorbent, sensor, temperature, pressure, humidity)

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 2007/0078113 A1 (Roth, et al.) 05 Apr 2007 (05.04.2007) Figures 11, 19, paragraphs [0179], [0377], [0489]-[0491], [0533], [0553], [0596], [0631], [0678]	1-27
Y	US 2006/0081524 A1 (Sengupta, et al.) 20 Apr 2006 (20.04.2006) Figures 1, 6, 10, paragraphs [0005]-[0006], [0024]-[0030], [0039]-[0042], [0046]-[0049]	1-27
Y	US 2010/0221697 A1 (Sehgal) 02 Sep 2010 (02.09.2010) paragraph [0146]	6-7, 22-23
Y	US 2002/0085952 A1 (Ellingsboe, et al.) 04 Jul 2002 (04.07.2002) Figures 1, 3B, paragraphs [0092], [0097], [0419]-[0420], [0473]	8-10, 18-19

Further documents are listed in the continuation of Box C.

* Special categories of cited documents:	
"A" document defining the general state of the art which is not considered to be of particular relevance	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E" earlier application or patent but published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"O" document referring to an oral disclosure, use, exhibition or other means	"&" document member of the same patent family
"P" document published prior to the international filing date but later than the priority date claimed	

Date of the actual completion of the international search  18 Jun 2012 (18.06.2012)	Date of mailing of the international search report  <b>03 JUL 2012</b>
Name and mailing address of the ISA/US  Mail Stop PCT, Attn: ISA/US, Commissioner for Patents P.O. Box 1450, Alexandria, Virginia 22313-1450 Facsimile No. 571-273-3201	Authorized officer:  Lee W. Young  PCT Helpdesk: 571-272-4300 PCT OSP: 571-272-7774