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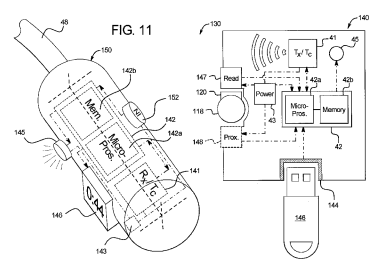
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(54) Title: SMART PATIENT TRANSFER SET FOR PERITONEAL DIALYSIS



(57) Abstract: A peritoneal dialysis system includes: a dialysis fluid container associated with an identifier (118); a reader for reading the identifier (147); a signal transmitter (41) for communicating data based on the identifier; and a patient transfer (150) set for semi-permanent connection to a patient, the patient transfer set including: (i) a signal receiver (141) for receiving the data sent by the signal transmitter, (ii) an output device (146) for communicating with the patient, and (iii) electronics configured to received the data and command the output device to communicate with the patient accordingly.



SMART PATIENT TRANSFER SET FOR PERITONEAL DIALYSIS

BACKGROUND

[0001] The present disclosure relates generally to medical fluid delivery systems and methods. More particularly, this disclosure includes systems, methods and apparatuses for selecting a dwell time for peritoneal dialysis based on an individual patient's response to dialysis, and also based on one or more peritoneal dialysis input parameters. The dwell time is selected to yield the best therapy outcome for that patient based on the dialysis parameters.

[0002] Due to various causes, a person's renal system can fail. Renal failure produces several physiological impairments and difficulties. The balance of water, minerals and the excretion of daily metabolic load is no longer possible and toxic end products of nitrogen metabolism (urea, creatinine, uric acid, and others) can accumulate in blood and tissue. Kidney failure and reduced kidney function have been treated with dialysis. Dialysis removes waste, toxins and excess water from the body that would otherwise have been removed by normal functioning kidneys. Dialysis treatment for replacement of kidney functions is critical to many people because the treatment is life saving.

[0003] Hemodialysis and peritoneal dialysis are two types of dialysis therapies used commonly to treat loss of kidney function. A hemodialysis ("HD") treatment utilizes the patient's blood to remove waste, toxins and excess water from the patient. The patient is connected to a hemodialysis machine and the patient's blood is pumped through the machine. Catheters are inserted into the patient's veins and arteries so that blood can flow to and from the hemodialysis machine. The blood passes through a dialyzer of the machine, which removes waste, toxins and excess water from the blood. The cleaned blood is returned to the patient. A large amount of dialysate, for example about 120 liters, is consumed to dialyze the blood during a single hemodialysis therapy. Hemodialysis treatment lasts several hours and is generally performed in a treatment center about three or four times per week.

[0004] Another form of kidney failure treatment involving blood is hemofiltration ("HF"), which is an alternative renal replacement therapy that relies on a convective transport of toxins from the patient's blood. This therapy is accomplished by adding substitution or replacement fluid to the extracorporeal circuit during treatment (typically ten to ninety liters of such fluid). That substitution fluid and the fluid accumulated by the patient in between

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treatments is ultrafiltered over the course of the HF treatment, providing a convective transport mechanism that is particularly beneficial in removing middle and large molecules.

[0005] Hemodiafiltration ("HDF") is another blood treatment modality that combines convective and diffusive clearances. HDF uses dialysate to flow through a dialyzer, similar to standard hemodialysis, providing diffusive clearance. In addition, substitution solution is provided directly to the extracorporeal circuit, providing convective clearance.

[0006] Peritoneal dialysis uses a dialysis solution, also called dialysate, which is infused into a patient's peritoneal cavity via a catheter. The dialysate contacts the peritoneal membrane of the peritoneal cavity. Waste, toxins and excess water pass from the patient's bloodstream, through the peritoneal membrane and into the dialysate due to diffusion and osmosis, i.e., an osmotic gradient occurs across the membrane. The spent dialysate is drained from the patient, removing waste, toxins and excess water from the patient. This cycle is repeated.

[0007] Peritoneal dialysis machines are used to accomplish this task. Such machines are described, for example, in the following U.S. Patents, all of which are incorporated by reference in their entirety, as though each patent were set forth herein, page by page, in its entirety: 5,350,357; 5,324,422; 5,421,823; 5,431,626; 5,438,510; 5,474,683; 5,628,908; 5,634,896; 5,938,634; 5,989,423; 7,153,286; and 7,208,092.

[0008] There are various types of peritoneal dialysis therapies, including continuous ambulatory peritoneal dialysis ("CAPD"), automated peritoneal dialysis ("APD"), tidal flow APD and continuous flow peritoneal dialysis ("CFPD"). CAPD is a manual dialysis treatment. The patient manually connects an implanted catheter to a drain, allowing spent dialysate fluid to drain from the peritoneal cavity. The patient then connects the catheter to a bag of fresh dialysate, infusing fresh dialysate through the catheter and into the patient. The patient disconnects the catheter from the fresh dialysate bag and allows the dialysate to dwell within the peritoneal cavity, wherein the transfer of waste, toxins and excess water takes place. After a dwell period, the patient repeats the manual dialysis procedure, for example, four times per day, each treatment lasting about an hour. Manual peritoneal dialysis requires a significant amount of time and effort from the patient, leaving ample room for improvement. There is room for improvement in the selection of dwell times for each patient.

[0009] Automated peritoneal dialysis ("APD") is similar to CAPD in that the dialysis treatment includes drain, fill, and dwell cycles. APD machines, however, perform the cycles

automatically, typically while the patient sleeps. APD machines free patients from having to manually perform the treatment cycles and from having to transport supplies during the day. APD machines connect fluidly to an implanted catheter, to a source or bag of fresh dialysate and to a fluid drain. APD machines pump fresh dialysate from a dialysate source, through the catheter, into the patient's peritoneal cavity, and allow the dialysate to dwell within the cavity, and allow the transfer of waste, toxins and excess water to take place. The source can be multiple sterile dialysate solution bags.

[0010] APD machines pump spent dialysate from the peritoneal cavity, though the catheter, to the drain. As with the manual process, several drain, fill and dwell cycles occur during APD. A "last fill" occurs at the end of CAPD and APD, which remains in the peritoneal cavity of the patient until the next treatment.

[0011] Both CAPD and APD are batch type systems that send spent dialysis fluid to a drain. Tidal flow systems are modified batch systems. With tidal flow, instead of removing all of the fluid from the patient over a longer period of time, a portion of the fluid is removed and replaced after smaller increments of time.

[0012] Continuous flow, or CFPD, systems clean or regenerate spent dialysate instead of discarding it. These systems pump fluid into and out of the patient, through a loop. Dialysate flows into the peritoneal cavity through one catheter lumen and out another catheter lumen. The fluid exiting the patient passes through a reconstitution device that removes waste from the dialysate, e.g., via a urea removal column that employs urease to enzymatically convert urea into ammonia. The ammonia is then removed from the dialysate by adsorption prior to reintroduction of the dialysate into the peritoneal cavity. Additional sensors are employed to monitor the removal of ammonia. CFPD systems are typically more complicated than batch systems.

[0013] In each of the kidney failure treatment systems discussed above, it is important to control ultrafiltration, which is the process by which water (with electrolytes) moves across a membrane, such as a dialyzer or peritoneal membrane. For example, ultrafiltration in peritoneal dialysis is a result of transmembrane and osmotic pressure differences between blood and dialysate across the patient's peritoneal membrane. It is also important to control the concentration of metabolic substances in the patient's bloodstream, such as urea concentration, β_2 -microglobulin, creatinine concentration, and so forth. Each of these, along with many other variables, constitutes a peritoneal dialysis outcome.

[0014] Each patient is different, possessing for instance, a unique peritoneal membrane, its own separation characteristics, and its unique response to peritoneal dialysis. Each patient is also different with respect to body surface area (BSA) and total body water volume, which also have an effect on transport characteristics. Each patient is different in terms of transport characteristics that relate to the ultrafiltration rate. Each patient is also different in terms of response to dialysis, that is, the amount of water and waste removed in a given time period, using a given fill volume, a particular dialysis fluid, and so forth. What is needed is a way to better control the particular dialysis therapy offered to each patient, so that the treatment will yield the best therapy outcome for that patient, for one or more dialysis input parameters

[0015] While APD frees the patient from having to manually performing the drain, dwell, and fill steps, a need still exists for CAPD. Some patients prefer the control that CAPD offers. Since the patient is awake during CAPD, the patient can adjust himself/herself during drain to produce more complete drains. Further, many patients who perform APD also perform a midday exchange using a CAPD technique.

[0016] Since CAPD does not typically use a machine, advantages of using a machine are not available to the CAPD patient, such as features intended to optimize therapy for the patent. It is therefore desirable to provide a "smart" system that is applicable to both APD and CAPD systems.

SUMMARY

[0017] One embodiment is a method for accomplishing peritoneal dialysis. The method includes steps of administering a peritoneal equilibration test (PET) to a patient, determining and recording a patient status as a result of the PET test, and calculating a peritoneal dialysis dwell time based on the patient status and at least one peritoneal dialysis therapy outcome, wherein the dwell time optimizes the at least one peritoneal dialysis therapy outcome for the patient.

[0018] Another embodiment is a method for accomplishing peritoneal dialysis. The method includes steps of determining peritoneal transport properties of a patient, determining a classification of the peritoneal transport properties of the patient, and calculating a peritoneal dialysis dwell time based on the classification, a plurality of peritoneal dialysis input parameters, and at least one desired peritoneal dialysis therapy outcome, wherein the

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dwell time is calculated to maximize the at least one desired peritoneal dialysis therapy outcome.

[0019] Another embodiment is a system for calculating a peritoneal dialysis dwell time and conducting peritoneal dialysis. The system includes a processor for operating a peritoneal dialysis machine, a memory of the processor or a memory accessible to the processor, the memory storing a look-up table containing peritoneal dialysis input parameters, peritoneal dialysis therapy outcomes, and peritoneal dialysis dwell times corresponding to the input parameters and dwell times, and a software program stored in the memory of the processor or the memory accessible to the processor for receiving a selection or an input of at least one desired therapy outcome of a patient and calculating a dwell time for the patient for optimizing the at least one peritoneal dialysis therapy outcome for the patient.

[0020] Another embodiment is a computer program embodied on a computer readable medium for calculating a peritoneal dialysis dwell time. The computer program includes a code segment for accessing data of a correspondence between a plurality of peritoneal dialysis input parameters, a plurality of peritoneal dialysis therapy outcomes, and a plurality of peritoneal dialysis dwell times. The computer program also includes a code segment that allows a user to input or to select at least one peritoneal dialysis input parameter from the plurality of peritoneal dialysis input parameters and at least one desired therapy outcome; a code segment that receives an indication of the at least one input parameter and at least one desired therapy outcome selected by the user; a code segment that calculates a dwell time corresponding to the at least one desired therapy outcome selected by the user; and a code segment that inputs the dwell time to the dialysis machine.

[0021] Another embodiment is a peritoneal dialysis system. The peritoneal dialysis system includes a dialysis cassette and a housing suitable for receiving the cassette, the cassette including at least one pump for pumping dialysis fluid to and from a patient; a microcontroller suitable for operating the peritoneal dialysis system; a memory of the microcontroller or accessible to the microcontroller, the memory including data of a plurality of dialysis input parameters, a plurality of dialysis dwell times, and a plurality of therapy outcomes corresponding to the input parameters and dwell times, wherein a user instructs the microcontroller to select or calculate a dwell time for optimizing at least one dialysis outcome for a patient; and a patient transfer device in communication with the microcontroller.

[0022] In yet another embodiment, a system including a smart transfer set is provided. The system includes a docking unit having a docking port that receives and holds the fill or solution bag line, and in one implementation a connector located at an end of the fill line. The connector (or perhaps the fill line itself) bears an identifier that in one embodiment identifies the dialysate solution type, solution volume and solution expiration date.

[0023] The docking unit is provided with a reader that reads the identifier. The docking port and the fill line connector are configured to mate, such that the reader can read the identifier. The identifier can be a barcode, in which case the reader is a barcode reader. The identifier is alternatively a radio frequency identifier ("RFID") tag, the reader an RFID reader, in which case the orientation of the connector within the docking port may not be as critical.

[0024] The docking unit in one embodiment also includes a computer memory device port, such as a universal serial bus ("USB") port. The port allows the patient to insert a memory device, such as a USB flash drive into the USB port, which allows the patient to download patient specific data to a memory located within the docking unit. That memory can also be used to store information read from the identifier. The docking unit in one embodiment includes processing that processes the information gleaned from the identifier and the patient memory device and develops therapy parameters that are sent from the docking unit to the transfer set. The transfer set is configured to communicate the parameter information to the patient, who uses the information to control a peritoneal dialysis therapy, such as a continuous ambulatory peritoneal dialysis ("CAPD") therapy. To this end, the docking unit is provided with a transmitter (or transceiver) that transmits the parameter information, e.g., wirelessly to a receiver (or transceiver) located within the transfer set.

[0025] The transfer set also includes memory and processing, which interface between the receiver and at least one output device for communicating with the patient, such as a light, buzzer or video display. The at least one output device communicates therapy parameter information to the patient, such as when to begin filling from the dialysate supply, when to attempt to have the fill completed, and when to drain spent dialysate, e.g., based on a determined dwell time. While the majority of the processing is done in the docking unit in one embodiment, it is also contemplated to let the transfer set do the majority of the processing, in which case the docking unit serves mainly to transfer information to the transfer set for processing.

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[0026] The therapy parameter information can be used in a number of ways. In one embodiment, the transfer set also includes an input device that the patient can activate once the patient is done filling from the dialysis fluid or dialysate container. Here, the docking unit can determine an optimum dwell duration and send it to the transfer set. Once the patient activates the input device, the transfer set beings a running of the optimal dwell duration. The transfer set can include a digital time remaining readout or display that counts down to zero, for example. The transfer set additionally or alternatively includes an alarm or buzzer that communicates when the dwell duration has lapsed, signaling a patient drain.

[0027] In another embodiment, the docking unit sends a fill duration and a dwell duration to the smart transfer set. The fill duration is determined principally from the dialysate volume gleaned from the identifier. The dwell duration is determined from the volume, the solution type (e.g., glucose level) and patient parameters, such as PET parameters. The fill duration begins to run, giving the patient an adequate time to fill from the dialysis fluid or dialysate container. When the patient fill times out, the dwell duration begins to run. The transfer set communicates the running of the dwell duration according to any of the ways discussed above. This embodiment does not require an input device or patient activation.

[0028] A third embodiment includes a fill duration and a dwell duration, like the last embodiment. The docking unit also includes a sensor, such as a proximity sensor (capacitive or inductive), which senses the presence or absence of the fill line connector received within the docking port. The proximity sensor is therefore located near the identifier reader. The sensor senses when the connector is removed from the docking unit, at which point it is assumed that the patient is about to connect the connector to the transfer set to begin filling. The sensed removal of the connector is communicated to the transfer set to begin (or begin after a short delay) the running of the fill duration. The running of the fill and dwell durations then proceeds according to the previous embodiment.

[0029] It is therefore an advantage of the present disclosure to provide peritoneal dialysis ("PD") systems and methods that optimize therapy dwell times for a patient.

[0030] It is another advantage of the present disclosure to provide peritoneal dialysis ("PD") systems and methods that streamline therapy time.

[0031] It is a further advantage of the present disclosure to provide peritoneal dialysis ("PD") systems and methods that operate with CAPD as well as APD.

[0032] It is still another advantage of the present disclosure to provide peritoneal dialysis ("PD") systems and methods that operate with different types and volumes of dialysate supplies or solutions.

[0033] It is still a further advantage of the present disclosure to provide peritoneal dialysis ("PD") systems and methods that preclude the use of an expired solution.

[0034] Additional features and advantages are described herein, and will be apparent from the following Detailed Description and the figures.

BRIEF DESCRIPTION OF THE FIGURES

[0035] Fig. 1 is a prior art peritoneal dialysis system.

[0036] Fig. 2 is a control system for a peritoneal dialysis system according to the present disclosure.

[0037] Fig. 3 is schematic view of a more detailed control system for a peritoneal dialysis system.

[0038] Figs. 4A and 4B are two embodiments of remote connectors that may be used with the control system.

[0039] Figs. 5 to 8 are graphs of peritoneal parameters and values of peritoneal parameters as they relate to dwell times for the peritoneal dialysis fluid.

[0040] Fig. 9 is a flow chart for a method of determining a dwell time for an optimal therapy outcome.

[0041] Fig. 10 is a perspective view of an alternative embodiment of the present disclosure, which employs a smart transfer set.

[0042] Fig. 11 is a schematic view of one software and electrical layout for the system and method of Fig. 10.

[0043] Fig. 12 is a flow chart for one method for implementing a dialysis treatment using a smart patient transfer set.

[0044] Fig. 13 is a flow chart for a second method (having two versions) for implementing a dialysis treatment using a smart patient transfer set.

DETAILED DESCRIPTION

Optimizing Therapy

[0045] Patients respond differently to peritoneal dialysis. There are a host of variables or parameters involved in peritoneal dialysis. One object of the present disclosure

is to discuss these therapy variables or parameters and show how they can be used in tailoring the therapy, and to show how that therapy can be tailored for the best possible outcome for that patient. Optimal execution of peritoneal dialysis therapy can help patients avoid numerous inefficiencies. These inefficiencies include the unnecessary loss of ultrafiltrate due to excessive dwell times, unnecessary carbohydrate absorption from long dwell times, and suboptimal urea ("Kt/V") and creatinine ("Ccr") clearances.

[0046] Rather than using a standard 14-15 hour daytime automated peritoneal dialysis ("APD") regimen, or a standard 9-10 nighttime continuous ambulatory peritoneal dialysis ("CAPD") regimen, a dwell time is calculated that is better suited to each patient. The dwell time depends on the patient's transmembrane transport capabilities, usually expressed as the patient's status as determined by a peritoneal equilibration test ("PET"). As an example of one benefit, carbohydrate ("CHO") absorption does not peak during peritoneal dialysis dwell times, but continues at a steady pace. The carbohydrate (e.g., glucose or icodextrin) in peritoneal dialysis solutions creates an osmotic gradient, enabling mass transport through the peritoneum. It is well known that this osmotic gradient decreases drastically during a long dwell time, most likely caused by absorption of the CHO itself into the peritoneum. Accordingly, shorter dwell times will reduce CHO absorption and free the patient for other activities.

[0047] One basic difference among patients is the rate at which water and metabolic wastes pass from the patient's bloodstream through the peritoneal membrane. Once the water and wastes pass through the peritoneal membrane, they are absorbed into the dialysis therapy fluid that has been placed into the patient's peritoneal cavity, and then removed from the patient. A peritoneal equilibration test (PET) determines the relative rate of transmembrane transport. Patients can then be classed as high-rate transporters, high-average transporters, low-average transporters, or low-rate transporters, depending on the speed of waste removal. Other classification schemes may also be used, such as simply high, average, and low transporters. Patients may also be classified in terms of their total body surface area (BSA), e.g., a high, average, or low BSA. The total body water volume may also be used as an input parameter to help predict transport characteristics of the patient.

[0048] In general, the rate of water removal is different from the rate of waste removal, and both depend on the patient transporter type. For example, fast transporters can quickly pass metabolic waste, but glucose from the dialysis solution is rapidly absorbed into the body. As a result, glucose concentration in the dialysate decreases and the osmotic

gradient diminishes within a relatively short period of time, depending on the patient transporter type. For instance, high transporters may benefit more from short dwell times, such as those used in automated peritoneal dialysis (APD), where the effect of high osmotic gradients is still present.

[0049] Conversely, the osmotic gradient will be sustained for a longer period of time in the case of a low transporter patient, resulting in a larger volume of ultrafiltrate removal. Such a patient will likely benefit from a longer dwell time, such a continuous ambulatory peritoneal dialysis (CAPD) and with perhaps only a single nighttime exchange. Much useful information about a patient's response to therapy can be learned from administering the PET test to the patient. The results of the PET test can then be used to administer the therapy that would lead to the best outcome for that patient.

[0050] Another variable in peritoneal dialysis is the fill volume, that is, the volume infused into the patient's peritoneum at the beginning of the dwell. The fill volume should be tailored to the comfort of the patient and the efficacy of the therapy. Fill volumes typically range from about 1.5 liters to about 3 liters, i.e., from about 1500 ml to about 3000 ml.

[0051] Therapy outcomes differ based on the fill volume, and they also differ based on the particular therapy fluid or peritoneal dialysis fluid used. For example, peritoneal dialysis fluids, such as Dianeal® from Baxter International, Deerfield, Illinois, U.S.A., may contain from 1.5% to 4.25% glucose. Other solutions may also be used. The glucose is used to provide a large osmotic pressure gradient between the infused dialysate solution and the patient's bloodstream, in order to draw excess water from the patient, i.e., ultrafiltrate. Other fluids may have other osmotic agents, such as icodextrin, e.g., 7.5% icodextrin in Extraneal® from Baxter International, which is typically used for longer dwell times.

[0052] The inputs to a particular dialysis therapy are thus seen to include at least the patient characteristics, i.e., his or her unique response to therapy, the therapy solution used, the total volume of therapy solution, and the dwell time used for the therapy. As discussed above, the outputs of the therapy are the results of the therapy. These results include the amount of water removed, typically expressed as net ultrafiltrate volume.

[0053] Other results or therapy outcomes include urea clearance, sometimes expressed as Kt/V, creatinine clearance (Ccr), and total carbohydrate absorption (CHO, also known as glucose or icodextrin absorption). Other therapy outcomes may also be measured, such as sodium removal, phosphate clearance, and middle molecule clearances, e.g., β_2 -microglobulin. Note that an optimum therapy outcome requires high removal of urea and

creatinine, as well as other wastes and ultrafiltrate. However, carbohydrate absorption should be minimized. As noted above, water can be transported both ways across the peritoneal membrane. Thus, net ultrafiltrate should be positive, with water removed, rather than having water absorbed from the peritoneal dialysis fluid, which would constitute negative ultrafiltration, and which is possible in a high transporter with a long dwell time.

[0054] In order to optimize therapy outcomes for individual patients, kinetic modeling has been undertaken using the above variables. Kinetic modeling software, PD Adequest 2.0TM, is from Baxter International, Deerfield, Illinois, U.S.A. This program uses a three-pore model of a patient's peritoneal membrane, and accepts choices of high, high-average, low-average, and low patient parameters, and uses a body surface area (BSA) input from 1.7 to 2.0 m². In developing the data presented herein, fill volume inputs of 1.5 L, 2 L, 2.5 L and 3 L were used, as were therapy fluid inputs of 1.5%, 2.5% and 4.25% glucose Dianeal, and Extraneal with 7.5% icodextrin.

[0055] The results were tabulated in tables and plotted on graphs. An example of different dwell times and the modeled results is depicted in Table 1 below.

Dwell time,	UF, ml	Urea	Creatinine	Glucose	Urea
hours		removal, g	removal,	absorption,	clearance,
			L/wk/1.73	g.	Kt/V
2.5	133	1.84	10.7	30.5	0.32
5	69	2.01	12.6	38.6	0.35
6	24	1.99	12.7	40.1	0.34
10	-184	1.81	11.87	42.6	0.31

[0056] Table 1, high transporter, 2 L fill, 2.5% Dianeal®

[0057] Table 1 clearly shows that a shorter dwell time is better for a high-transporter patient. After 2.5 hours, the net ultrafiltrate (UF) is highest, 133 ml net ultrafiltrate, with high levels of urea and creatinine removal as shown. Allowing the therapy fluid to dwell for five hours has a deleterious effect: the net ultrafiltrate has decreased by about half, to abut 69 ml, and glucose absorption has increased by about 25% to 38.6 g. Because the volume of ultrafiltrate continues to decrease, this patient may already have reached a point of diminishing returns, at least for ultrafiltrate volume.

[0058] The only benefit from increased dwell time is a small increase in urea and creatinine removal. However, if there were some medical reason to do so, one could select

the urea removal or creatinine removal as the outcome of interest, and select the appropriate dwell time, 5 hours for urea removal or 6 hours for creatinine removal. This selection would optimize the value of the particular desired outcome, whether net ultrafiltrate, urea removal, or creatinine removal, or the removal of other solutes, such as phosphate or β_2 -microglobulin that are not quantified here.

[0059] Another example for a high-transporter patient is depicted in Table 2 below, in which the principal change is to use 2.5 L fill volume rather than 2 L.

Dwell time,	UF, ml	Urea	Creatinine	Glucose	Urea	
hours		removal, g	g removal, absorption,		clearance,	
			L/wk/1.73	g.	Kt/V	
3	165	2.30	13.32	38.0	0.40	
6	86	2.52	15.78	48.4	0.44	
7	40	2.50	15.91	50.0	0.43	

[0060] Table 2, high transporter, 2.5 L fill, 2.5% Dianeal®

[0061] Table 2 depicts results of using additional fill volume and slightly longer dwell times. Using the additional 500 ml of fill volume has caused an increase in net ultrafiltrate, to 165 ml, with increases in urea and creatinine removal over the amounts removed with 2.5 hours dwell and a 2 L fill volume. While these are desirable, there has been an increase in glucose absorption. The caregiver or medical professional can decide whether the increased ultrafiltration, urea removal and creatinine removal is sufficient to justify an increase in glucose absorption. The desired outcome is then used to select the dwell time for the patient, as well as whether it is desirable to use 2 L fill volume or 2.5 L fill volume. Other examples are given below for other transporter conditions.

[0062]	Table 3. ł	nigh-average i	transporter.	2. L	fill.	2.5% D	ianeal® -
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Dwell time,	UF, ml	Urea	Creatinine Glucose		Urea	
hours		removal, g	removal,	absorption,	clearance,	
			L/wk/1.73	g.	Kt/V	
3.5	181	2.21	10.04	30.04	0.33	
6	125	2.35	11.87	36.93	0.35	
8	40	2.30	12.26	39.82	0.34	

Dwell time,	UF, ml	Urea	Creatinine	Glucose	Urea	
hours		removal, g	removal,	absorption,	clearance,	
			L/wk/1.73	g.	Kt/V	
4	222	2.74	12.29	36.77	0.41	
7	160	2.94	14.82	46.00	0.44	
10	29	2.86	15.41	50.41	0.43	

[0063] Table 4, high-average transporter, 2.5 L fill, 2.5% Dianeal®

[0064] Tables 3 and 4, for high-average transporters, demonstrate a shift of peak time points to longer dwell times. The ultrafiltration volumes are higher at these longer dwell times, higher than the ultrafiltration volumes for the high-transporter Tables 1 and 2. There are also greater urea and creatinine removals.

[0065] Table 5, low-average transporter, 2 L fill, 2.5% Dianeal®

Dwell time,	UF, ml	Urea	Creatinine	Glucose	Urea	
hours		removal, g	removal, g removal, absorption,		clearance,	
			L/wk/1.73	g.	Kt/V	
4	209	2.45	9.14	26.80	0.32	
7	173	2.67	11.34	34.50	0.35	
11	28	2.58	12.08	39.64	0.33	

[0066] Table 6, low-average transporter, 2.5 L fill, 2.5% Dianeal®

Dwell time,	UF, ml	Urea	Creatinine	Glucose	Urea
hours		removal, g	removal,	absorption,	clearance,
			L/wk/1.73	g.	Kt/V
5	258	3.10	11.69	34.37	0.40
9	193	3.35	14.50	44.57	0.43
13	41	3.23	15.17	49.64	0.42

[0067] Tables 5 and 6 demonstrate shift of peak time points to even longer dwell times for low-average transporter patients. The ultrafiltration volumes are higher at these longer dwell times, higher than the ultrafiltration volumes for the high-transporter Tables 1 and 2 and for high-average transporter Tables 3 and 4, especially with a 2.5 L fill. There are also greater urea and creatinine removals, urea removal peaking at about 9 hours.

[0068] Table 7,	low transporter,	2 L fill, 2.5%	Dianeal®

Dwell time,	UF, ml	Urea	Creatinine	Glucose	Urea	
hours		removal, g removal, abso		absorption,	clearance,	
			L/wk/1.73	g.	Kt/V	
6	335	2.24	9.01	26.94	0.37	
9	287	2.33	10.66	32.83	0.38	
15	49	2.16	11.63	39.42	0.35	

[0069] Table 8, low transporter, 2.5 L fill, 2.5% Dianeal®

Dwell time,	UF, ml	Urea	Creatinine Glucose		Urea	
hours		removal, g	removal,	absorption,	clearance,	
			L/wk/1.73	g.	Kt/V	
7	409	2.79	11.14	33.42	0.46	
11	342	2.91	13.48	41.76	0.48	
16	148	2.78	14.58	47.97	0.45	

[0070] Tables 7 and 8 show that the low transporter patients benefit from longer dwell times, in clear contrast with the high transporter patients. The ultrafiltration volumes are significantly increased, and may already have peaked since the volumes are decreasing. Urea removal peaks at 9 and 11 hours respectively

[0071] These data can be used to select a dwell time for the best possible outcome of a dialysis therapy session for a specific patient. Dialysis therapy is typically conducted with a peritoneal dialysis machine, such as the machine depicted in Fig. 1. One suitable peritoneal dialysis machine is the HomeChoice® peritoneal dialysis machine from Baxter International, Deerfield, IL, U.S.A. A patient P is connected to a dialysis machine 1, shown within the dashed lines, with a patient access device 5, such as an implanted catheter as shown. The catheter may be a single lumen or double lumen catheter, or another type of access device may be used. A plurality of containers 2 of dialysis solution is connected to the dialysis machine, as shown, through valves or other connectors. A pump 3 is used to transport dialysis fluid from the containers 2, one at a time, through a balance chamber 4 to the peritoneal cavity of the patient P through the access device. After the peritoneal dialysis solution has remained within the patient for the desired dwell time, the same pump 3 or another pump 6 may be used to pump the spent dialysis solution through the balance chamber 4 and then to a drain 7.

[0072] In embodiments discussed herein, a dialysis machine 1 may be used with a dialysis control system 10 as depicted in Fig. 2. Dialysis control system 10 includes an operating portion, such as the peritoneal dialysis machine depicted in Fig. 1, including fluid lines 12 for connection to patient access device 15. The operating section 11 performs dialysis for the patient under the supervision of a control unit 13. Control unit 13 in one embodiment has at least an input keypad 14, control panel 14a, which may be a touch screen, input number pad 14b, and mouse 14c. The control unit will also include input drive 15a, which may be suitable for a floppy drive or for a CD drive. The computer in this embodiment is configured with a port for Internet access 15b, as well as additional inputs and outputs, including ports 16. The additional input ports may be any combination of serial ports, such as USB ports, or parallel ports.

[0073] In some embodiments, the control unit will be adapted to receive commands from a remote control unit, and will include an IR receiver 15c for a hand-held remote. Inputs/outputs may include an optical input or output 15d and other digital or analog inputs. Control portion 15e includes a series of controls knobs or switches for operating the dialysis machine. A speaker output 17 can alert the patient or a caregiver if there is an emergency or other malfunction of the dialysis machine. There is also a visual alarm 15f for alerting the patient or caregiver. The control section includes an antenna 19 for receiving remote commands or information. The antenna may be used for communication with a wireless device for the patient, as discussed below. The antenna may also be used for wireless (WiFi) internet access or may be used for remote, but closer, commands.

[0074] Fig. 3 depicts a closer view of the control portions 30 of the dialysis machine 10. Machine control portion 30 is in communication with a "smart" patient control portion 40. As seen in Fig. 3, the communication is wireless, for convenience and mobility of patients, such as mobile CAPD patients. However, those with skill in the art will recognize that a wire harness or cable could also connect the two portions. Dialysis machine control portion 30 includes a supervisory microcontroller 31, which receives power from a power supply 32. The microcontroller receives inputs from at least a keypad 33, and may also receive data and commands from a wired connection 34, such as from a clinic or hospital information system. Inputs may also be received from the patient via wireless connection and radio 35. The microcontroller has outputs to a video monitor 36, a speaker 37, as well as controls to the dialysate pumps 38 and a heater 39 for the dialysate. The machine control

system includes at least one memory as a part of the microcontroller 31 or accessible by the microcontroller 31.

[0075] The patient control portion 40, as noted above, is not attached to the dialysis machine, enabling a mobile patient to move about without a wire harness or other connecting cable. Of course, other embodiments may include a cable, infrared (IR) or RF communications instead of the radio described herein. The patient control portion includes a separate microcontroller 42 and power supply 43, such as a battery 42. The controller 42 receives input from the radio 41, with outputs through the radio and to an audio alarm or speaker 45 and a small video monitor 46. In some embodiments, the patient control portion may also include switches or other electromechanical inputs for signaling the microcontroller 42 or for controlling the operation of the patient control portion 40.

[0076] The signal processing circuitry and radio 41 or wireless receiver/ transmitter are small and compact, and are easily placed on the patient at the access site, such as in a "smart" module or connector. One radio that works is a wireless module in accord with ZigBee/IEEE 805.15.4. This is a standard for a very low power radio system with a very limited range, about 10-20 feet. Modules made in accordance with this standard may be purchased from Maxstream, Inc., Lindon, UT, U.S.A., Helicomm, Inc., Carlsbad, CA, U.S.A., and ANT, Cochrane, Alberta, Canada. The module is very small, and may be about 2 cm square (about 1 inch square), and about 3 mm thick (1/8 inch). The patient control portion 40, as noted, is intended for close proximity, within range of the ZigBee module, of about 10-20 feet, of the dialysis machine. Thus, the local portion or signal module is conveniently small and unobtrusive for the patient, but fully capable of communication and control with the machine control portion 30.

[0077] The patient may use the patient control portion or may simply use the dialysis machine, such as the embodiment depicted in Fig. 2. In one embodiment, shown in Fig. 4a, the patient P is connected to the dialysis machine through patient line 18, through patient control device and transfer set 50, and a catheter serving as a peritoneal access device 47. Patient control device 50 is connected via luer connectors 49, or other suitable connectors. The present day transfer set, into which the patient control device can be integrated, includes a length of tubing with a clamp, the length of tubing including one luer connector, such as a titanium luer, for connecting to the patient access device and a second luer for connecting to the patient line. Those who have skill in the art will recognize that patient transfer sets vary in regards to the connecters used, and also vary in the clamp (not shown) or clamps used.

[0078] In this embodiment, the patient access device 47 is a double-lumen catheter and the patient line 18 includes two lengths of tubing. Patient control device 50 includes an audio alarm or speaker 45 and a lamp 52, such as an LED, to alert a patient when the therapy session has begun or has ended. Two lamps may be used, such as a green lamp when therapy has begun and a red lamp to alert the patient that the therapy session has ended. In one embodiment, patient control device 50 can be disengaged and separated from the tubing or transfer set for cleaning, replacement, and so forth. In another embodiment, the device cannot be disengaged and is embedded within the transfer set in such a manner as to extend the functions of the transfer set, without significantly impacting the volume or area (footprint) of the transfer set.

[0079] Another embodiment 40 of the patient control device, and its application, is depicted in Fig. 4B. Patient P is connected to the dialysis machine through a single-lumen patient line 18, patient control device and transfer set 40, and a single-lumen catheter 48 serving as a patient access device. Patient control device 40 is connected via luer connectors 50 or other suitable connectors. The patient control device 40 includes a small video output 46 and a lamp 52. An audio alarm 45 may be used to signal the patient to begin or end a therapy session. The video output 46 is suitable for displaying a time remaining on the dialysis session. Lamp 52 may be used to signal the patient, as discussed above. The patient control device 40 also includes two switches 51, suitable for allowing the patient to respond to queries from the microcontroller 42. The switches, in this embodiment, are "yes" and "no" switches that are suitable for responding to queries from the controller, such as "shall we start the dialysis session?" or "please enter a start time for the dialysis session."

[0080] In addition to tabular data, as might be expected in a look-up table, the correspondence between dwell times and therapy outcomes can be expressed as equations, and can be presented in graphical format, as shown in Figs. 5-8. Each of these figures depicts performance for a patient according to the PET test status of the patient, as explained above for the tables. Fig. 5, for instance, for a high-transporter patient (H), depicts therapy outcomes for patients that have been dialyzed with 2 L or 2.5 L of 2.5% glucose Dianeal® peritoneal dialysis solution. The solid lines depict 2.5 L results and the dotted lines depict 2 L results, for net ultrafiltrate (read on the scale on the left side of the graph), urea removed (read on the scale on the right side of the graph). The data demonstrate that net ultrafiltrate peaks at about 2.5 or 3 hours for both a 2 L or a 2.5 L fill for a high-transporter patient. Urea removal peaks

at about 4 hours with a 2 L fill (dotted line), or at about 6 hours for a 2.5 L fill (solid line). For creatinine, clearance peaks at about 6 hours for a 2 L fill and at about 7 hours for a 2.5 L fill.

[0081] Using this data, a patient or a caregiver, such as a medical professional, can select a dwell time based on the desired outcome, e.g., 3 hours dwell time with a 2.5 L fill for maximum ultrafiltrate. If the desire is to accommodate more than one desired outcome, a compromise can be reached by interpolating between the desired outcomes. For instance, both high ultrafiltrate and high urea removal may be desired. For a 2.5 L fill, maximum urea removal occurs at about 6.5 hours. A dwell time between 3 and 6.5 hours may be selected, based on averaging the two times, such as 3 + 6.5 = 9.5, and then dividing by two, to arrive at about 4.75 hours. If one outcome or therapy result is deemed more important, a weighted average may be used. For example, if ultrafiltration is more important, the time for the best ultrafiltration outcome may be multiplied by a weighting factor of 2, while using a weighting factor of 1 for the creatinine dwell time. In this case, the result would be $(3 \times 2) + 6.5 = 12.5$, and then dividing the result by three, to arrive at a dwell time of about 4.2 hours. Weighting factors may be pre-selected and may be programmed into a computer program to calculate a resultant dwell time.

[0082] Fig. 6 depicts therapy outcomes for a patient whose PET status is that of a high-average (HA) transporter. The kinetic modeling was accomplished using parameters of 2.5% glucose Dianeal®, with 2 L and 2.5 L fill volumes. In these results, the creatinine has a very subtle peak removal at about 9 hours, with very little effect from adding or subtracting two hours from the dwell time. The ultrafiltration and urea removal curves are shifted somewhat to the right, with greater dwell times, in comparison with those of Fig. 5 for H transporters. In other words, HA transporters act more slowly and need more time than H transporters, which is to be expected.

[0083] Fig. 7 depicts results for low-average (LA) transporters, with the same input parameters of 2.5% glucose Dianeal®, with 2 L and 2.5 L fill volumes. The curves for creatinine now have no peak, suggesting that longer dwell times result in additional cumulative creatinine transport. The curves for urea removal now have more subtle peaks, and the curves for both urea and ultrafiltrate are shifted toward even longer dwell times.

[0084] Fig. 8 demonstrates great differences in transport properties for low (L) transporter patients. In comparison with patients having higher transport status, low transporters have much higher rates of ultrafiltration and slower rates of waste removal. The

ultrafiltration curves, the highest two curves on the left side of the chart, have much higher rates (note that the left-hand scale is changed from the previous graphs). After about 6-7 hours, depending on the fill, the ultrafiltration reverses, i.e., dialysis solution is being absorbed into the patient's peritoneum, rather than ultrafiltrate being expressed into the dialysis solution. The ultrafiltration curves are thus the lowest two curves at the longest dwell times (right side of chart). Creatinine removal does not peak, even at 16 hours, while urea removal peaks at about 9 hours (2 L fill) and about 10 hours (2.5 L fill).

[0085] Many additional data may also be used to understand the transport behavior of peritoneal dialysis patients, as shown in the tables below, which concern respectively, performance at fill volumes of 1500 ml (1.5 L), 2000 ml (2 L), 2500 ml (2.5 L), and 3000 ml (3 L). These data are depicted respectively in Tables 9-12.

[0086] Table 9, 1500 ml (1.5 L) volume fill

1.5 % Dianeal Solution 2.5% Dianeal solution 4.25% Dianeal Solution Extraneal Solution

	Dwell	Net	Kt/V									
		UF			UF			UF			UF	
Н	3.0	18		2.0	103		3.0	294		16	534	
	3.5		.24	4.0		.26	5.0		.29	16		.35
HA	1.5	25		2.5	140		4.0	394		16	594	
	4.0		.24	5.0		.28	6.0		.31	16		.35
LA	2.0	28		3.5	164		5.0	463		16	548	
	5.0		.29	6.0		.26	7.0		.31	16		.34
L	2.5	5.6		5.0	260		7.0	693		16	665	
	6.0		.25	7.0		.29	9.0		.36	16		.37

[0087] Table 10, 2000 ml (2 L) volume fill

1.5 % Dianeal Solution 2.5% Dianeal solution 4.25% Dianeal Solution Extraneal Solution

	Dwell	Net	Kt/V									
		UF			UF			UF			UF	
Н	1.5	24		2.5	133		3.5	375		16	651	
	4.0		.32	5.0		.35	6.0		.38	16		.46
HA	2.0	34		3.5	181		5.0	500		16	700	
	5.0		.32	8.0		.35	7.0		.41	16		.46
LA	2.5	37		4.0	209		7.0	586		16	630	
	7.0		.31	7.0		.35	9.0		.41	16		.45
L	3.5	75		6.0	335		9.0	877		16	750	
	8.0		.33	9.0		.38	11.0		.48	16		.47

[0088] Table 11, 2500 ml (2.5 L) volume fill

1.5 % Dianeal Solution 2.5% Dianeal solution 4.25% Dianeal Solution Extraneal Solution

	Dwell	Net	Kt/V	Dwell	Net	Kt/V	Dwell	Net	Kt/V	Dwell	Net	Kt/V
		UF			UF			UF			UF	
Н	1.5	31		3.0	165		4.0	457		16	739	
	5.0		.41	6.0		.44	7.0		.49	16		.56
HA	2.0	42		4.0	222		6.0	606		16	777	
	6.0		.40	7.0		.44	8.0		.50	16		.59
LA	3.0	47		5.0	258		8.0	712		16	689	
	8.0		.39	9.0		.43	11.0		.51	16		.53
L	4.0	93		7.0	408		10.0	1065		16	825	
	9.0		.42	11.0		.48	13.0		.58	16		.57

[0089] Table 12, 3000 ml (3 L) volume fill

1.5 % Dianeal Solution 2.5% Dianeal solution 4.25% Dianeal Solution Extraneal Solution

	Dwell	Net	Kt/V	Dwell	Net	Kt/V	Dwell	Net	Kt/V	Dwell	Net	Kt/V
		UF			UF			UF			UF	
H	2.0	38		3.5	197		5.0	545		16	806	
	5.0		.49	7.0		.52	8.0		.59	16		.65
HA	2.5	52		5.0	264		7.0	719		16	836	
	7.0		.48	3.0		.52	10.0		.60	16		.65
LA	3.5	57		6.0	302		9.0	839		16	730	
	9.0		.47	10.0		.52	12.0		.81	16		.61
L	5.0	112		8.0	465		12.0	1249		16	872	
	10.0		.50	12.0		.57	15.0		.70	16		.66

[0090] The tables demonstrate that some of the trends discussed above hold true across many variables, e.g., ultrafiltration increases with decreasing transport properties, and ultrafiltration also increases with increasing fill volume. Other trends can also be discerned, but the point remains the same: the data can be used to select a dwell time to optimize the dialysis session for a particular patient, based on the patient's transport properties, fill volume, and dialysis solution used.

[0091] It is understood that the tables are simply an easy way to present data. The correspondence between dialysis input parameters, desired therapy outcomes, and dwell times may reside as data in one or more tables, such as look-up tables. The data may also

take the form of graphs, or may be reduced to equations. There are many embodiments of the invention, including all of these methods of presenting, storing, and using the data.

[0092] A flowchart for a method of optimizing a therapy session is depicted in Fig. 9. The classification of the patient as a high transporter, high-average transporter, and so forth, is determined by administering 91 a PET test. In other embodiments, other tests may be used, and other classifications may be used. The point is to determine how each individual patient can benefit from the dialysis conditions best suited for him or her. In any event, the transport properties of the patient with respect to ultrafiltration and waste transport are determined 92. The patient or a caregiver, such as a medical professional, determines 93, e.g., selects, at least one desired outcome of a dialysis treatment, such as ultrafiltrate volume or urea clearance.

[0093] A dwell time for optimizing the selected outcome is then calculated or estimated 94. The selected dwell time is then entered 95 into the controller of the dialysis machine. For dialysis treatments that do not necessarily involve a peritoneal dialysis machine, such as ambulatory peritoneal dialysis, the controller may simply be a timer on the transfer set. The therapy session is then conducted 96 using the entered dwell time. At the end of the dwell time, the timer or other alerting device alerts 97 the patient that the dwell time has expired and the therapy session may be ended.

[0094] It should be understood that various changes and modifications to the presently preferred embodiments described herein will be apparent to those skilled in the art. Such changes and modifications can be made without departing from the spirit and scope of the present subject matter and without diminishing its intended advantages. It is therefore intended that such changes and modifications be covered by the appended claims.

Smart Patient Transfer Set

[0095] In a further embodiment, the present disclosure sets forth a peritoneal dialysis "smart" system and method for performing peritoneal dialysis, which is applicable to both continuous ambulatory peritoneal dialysis ("CAPD") and automated peritoneal dialysis ("APD") therapies. The system and method take advantage of one common component for both CAPD and APD, namely, that both types of systems include or connect to a patient's transfer set. As discussed above, the patient's transfer set is a permanent item worn and carried around by the patient. The set is connected to a tube that transitions to an indwelling catheter located within the patient's peritoneal cavity.

[0096] The patient's transfer set connects to a connector located at the end of a patient fill tube. The fill tube can extend from a manual flow control device for CAPD or a disposable cassette for APD. The manual flow control device connects to a single supply bag typically. The disposable cassette connects to multiple supply bags typically. Thus another constant with CAPD and APD systems is the use of at least one supply bag.

[0097] The system places a readable identifier on a connector located at the end of a tube or pigtail extending from the supply bag. Alternatively, the identifier is placed on the tube or pigtail itself. Further alternatively, if the bags do not include tubes or pigtails and are instead spiked, the bags themselves may be provided with identifiers that are read. Still further alternatively, in an APD embodiment the identifier can be placed on a connector connected to a patient fill line running from the APD machine. The identifiers in any case can be a barcode or radio frequency identifier, for example.

[0098] A reader is provided that reads the identifier. The reader can be integrated into the APD machine or be provided in a standalone unit. It is contemplated to use the standalone unit in both APD and CAPD therapies. The reader reads the identifier, processes the information obtained from the identifier, and sends a signal based on the processed information.

[0099] The patient's transfer set receives the signal sent from the reader. The signal may be a wireless signal sent from a transmitter (or transceiver) of the reader to a receiver (or transceiver) of the transfer set. The transceivers are provided alternatively for two-way communication between the reader and the transfer set. The transfer set is also provided with microprocessing and memory that are programmed to act on the received information. The microprocessor outputs to an output device, such as an alarm and/or readout, that communicates information to the patient. In one example, the supply container identifier supplies information regarding the volume of fluid in the dialysis fluid or supply container and a dextrose or glucose level of the dialysate residing in the container. The identifier may also include expiration date information for the supply.

[00100] Referring now to Figs. 10 and 11, one embodiment of a smart transfer set system 130 and corresponding method is illustrated. System 130 includes two primary components, namely, an alternative control portion 140, which is hereafter called a remote docking unit, and an alternative or smart transfer set 150. Remote docking unit 140 can be configured for operation with control portion 30 of the dialysis machine 10 discussed above

for APD. Alternatively, docking unit 140 operates solely with smart transfer set 150 to run a CAPD therapy.

[00101] Docking unit 140 accepts a supply line connector 118 having a tag or identifier 120, which can be a barcode or radio frequency identification ("RFID") tag. Tag or identifier 120 is placed alternatively on the supply line itself, e.g., near supply line connector 118. Further alternatively, in APD application, supply line connector 118 can instead be a connector for the patient line 18 coming from APD machine 30.

[00102] Docking unit 140 includes a reader 147 positioned in suitable proximity to tag or identifier 120 to read the information from the tag. Reader 147 can for example be a barcode scanner that faces barcode 120 to read its information. Reader 147 is alternatively an RFID reader that may not have to be placed as directly adjacent to RFID tag 120 to read the information.

[00103] The information in one embodiment includes (i) the volume, (ii) the type and (iii) the expiration date of solution or dialysis fluid held by the supply bag connected to the supply line connected to supply line connector 118. If the dialysis fluid has expired, an alarm will be posted and if used with APD machine 10, system 130 will not allow therapy to continue. The volume and type of dialysis solution is used to set fill and dwell times as discussed in detail below.

[00104] Docking unit 140 also includes a memory card or drive port 144, such as a universal serial bus ("USB") port for receiving a memory storage member 146, such as a flash drive or disk drive. The memory storage member 146 stores patient specific data, such as data shown above in Tables 1 to 12, which is combined with data from tag or identifier 120 to determine an optimal dwell time for the patient. In one embodiment, memory storage member 146 needs to be inserted into port 144 for each treatment. In another embodiment, memory storage member 146 needs to be inserted only once into port 144 until the data on the memory storage member 146 is changed.

[00105] In one alternative embodiment discussed below in connection with Fig. 13, a separate proximity sensor 148 is provided. Proximity sensor 148 can be a capacitive or inductive proximity sensor. The proximity sensor senses the presence or absence of connector 118 for reasons discussed below.

[00106] Each of the reader 147, memory member receiving port 144 and proximity sensor 148 is linked to a controller 42 provided within docking unit 140. In the illustrated embodiment (and for any of the embodiments discussed herein), controller 42 includes

processing 42a and memory 42b. Processing 42a and memory 42b are also linked in communication with a wireless transmitter (Tx) or transceiver (Tc) 41, which communicates wirelessly with a receiver (Rx) or transceiver (Tc) 141 located within smart transfer set 150.

[00107] Controller 42 also commands one or more output device 45, such as a light and/or buzzer, which can for example communicate to the patient whenever docking unit 140 is processing or transmitting data. For example, an alarm output device 45 can be provided to indicate when that the solution of a bag connected to connector 118 has expired. Output device 45 can alternatively be a display which indicates "expired" in this instance.

[00108] A power supply 43, which can be AC sourced, be a rechargeable battery or be a replaceable battery supplies the appropriate power to each of controller 42, output device 45, reader 147, transmitter (Tx) or transceiver (Tc) 41 and proximity switch 148 (if provided). Low power can be indicated to the patient via light indication or via a readout.

[00109] Processing 42a and memory 42b receive the solution data from reader 147 and patient data from memory storage member 146 and process the data to arrive at an optimal dwell time according to the methods described herein. That optimal dwell time is then sent via transmitter (Tx) or transceiver (Tc) 41 wirelessly (possibly with other information as discussed below) to receiver (Rx) or transceiver (Tc) 141 located within smart transfer set 150. Alternatively, docking unit 140 serves mainly as an information transfer device, which transfers solution data from reader 147 and patient data from memory storage member 146 to smart transfer set 150, which then uses its processing 142a and memory 142b to compute the optimal dwell duration (and other needed information as discussed below). It may be possible under one of these scenarios to remove or limit the processing and memory from one of docking unit 140 and smart transfer set 150.

[00110] As seen in Fig. 11, smart transfer set 150 includes a controller 142 having processing 142a and memory 142b, which accept information from receiver (Rx) or transceiver (Tc) 141 and command operation of one or more patient output device, such as a light and/or buzzer 145 and a small (e.g., liquid crystal display ("LCD")) readout 146. In one embodiment, discussed below in connection with Fig. 12, smart transfer set 150 is provided with a patient input device 152, such as a pushbutton or switch. Input device 152 inputs a signal to controller 142, for example, to indicate when a fill of the solution from the bag to the patient has been completed.

[00111] Transfer set 150 also includes a power supply 143, which can be a rechargeable battery or a replaceable battery, and which supplies the appropriate power to

each of controller 142, receiver (Rx) or transceiver (Tc) 141 and output devices 145 and 146. Again, low power can be indicated to the patient via light indication or via a readout.

[00112] Referring now to Fig. 12, a method 160 for operating system 130 is illustrated. Upon beginning method 160 as seen at oval 162, dwell duration is determined as seen at block 164. Dwell duration is calculated using the solution data from reader 147 and patient data from memory storage member 146 and the methodology set forth herein by either the controller 42 of docking unit 140 or controller 142 of smart transfer set 150 as discussed above. Dwell time duration is thus either determined at smart transfer set 150 or sent to smart transfer set 150 from docking unit 140, and in any case is known by smart transfer set 150 at block 164.

[00113] At diamond 166, method 160 waits for the patient to press input device 152 indicating that a fill from a supply bag connected to connector 118 has been completed. Method 160 allows for the patient to fill as fast or slow as the patient desires and is independent of fill time. When the patient presses input device 152 indicating that the fill is complete, method 160 proceeds to block 168, at which time the optimal dwell duration beings to run. Small display 146 at Fig. 11 shows one embodiment in which dwell duration is counted backwards from the starting duration down to zero. It is contemplated to build false input device 152 activation protection into method 160, e.g., requiring a confirm press of the input device or allowing a second press of the input device to undue the original input and start over.

[00114] At diamond 170, method 160 waits for the dwell duration to run out completely, at which time the patient is alerted that dwell is finished and that the patient should begin to drain the spent dialysate, as seen at block 172. Different scenarios are contemplated. For example, light/buzzer 145 could flash with five or ten minutes before dwell is completed to give the patient a heads-up the he/she needs to get to a place appropriate for draining. All the while clock 146 is counting down to zero. At zero, light/buzzer 145 lights/sounds to indicate that dwell is complete and that drain needs to start as soon as possible. The buzzer can be for a predetermined duration, and input device 152 can be set to stop the buzzer immediately when pressed in case the patient does not want noise.

[00115] At oval 174, method 160 ends. Method 160 is then repeated for each supply bag of the therapy.

[00116] Referring now to Fig. 13, another method 180 (actually two versions of this methods as discussed below) for operating system 130 is illustrated. Upon beginning method 180 as seen at oval 182, fill and dwell durations are determined as seen at block 184. Dwell duration is again calculated using the solution data from reader 147 and patient data from memory storage member 146 and the methodology set forth herein by either the controller 42 of docking unit 140 or controller 142 of smart transfer set 150 as discussed above. Dwell duration is in any case is known by smart transfer set 150 at block 184. Both fill and dwell durations are affected by the volume of the dialysis solution in the supply container. Dwell duration is also affected by the type or dextrose level of the dialysis solution. Fill duration is also effected by the patient's fill position relative to the supply container if the fill is a gravity fill. Position can play a role even when fluid is pumped from the supply bag to the patient. Thus, the fill duration may be for a particular head height level, which is either known generally by the patient or communicated via system 130 to the patient via the APD machine 10, docking unit 140 or smart transfer set 150.

[00117] At block 186a, the patient is alerted that the patient should begin filling fresh dialysate from the supply container, through connector 118 and transfer set 150, to the patient's peritoneum. Different scenarios are contemplated for the fill duration run-out indicated at block 188. For example, light/buzzer 145 could flash and or buzz, while display 146 counts the fill duration down to zero. Display 146 could also display the word "fill" to indicate that the current countdown is for filling. Alternatively or additionally, light 145 could be lighted a different color for fill (e.g., green) and dwell (e.g., yellow).

[00118] At diamond 190, method 180 waits for the fill duration to run out completely, at which time the patient is alerted that fill is supposed to be finished and that the dwell duration is beginning. Different scenarios are contemplated dwell duration run-out indicated at black 192. For example, light/buzzer 145 could flash and or buzz for a period to indicate the transition from fill to drain, while display 146 resets itself and counts now the dwell duration down to zero. Display 146 could also display the word "dwell" to indicate that the current countdown is for filling. Alternatively or additionally, light 145 could be changed from, e.g., green to yellow. Method 180 accordingly does not require input device 152 because the method transitions automatically from fill to dwell.

[00119] At diamond 190, method 180 waits for the dwell duration to run out completely, at which time the patient is alerted that dwell is finished and that the patient should begin to drain the spent dialysate, as seen at block 198. As with method 160,

impending dwell completion can be indicated by the flashing of light/buzzer 145 with five or ten minutes before dwell is done to give the patient a heads-up the he/she needs to get to a place appropriate for draining. All the while clock 146 is counting down to zero. At zero, light/buzzer 145 lights/sounds to indicate that dwell is complete and that drain needs to start as soon as possible. The buzzer can be again be for a predetermined duration, and input device 152 can be set to stop the buzzer immediately when pressed in case the patient does not want noise.

[00120] Block 186b illustrates a modification of method 180. If there is going to be a substantial error in estimating fill duration, it is likely going to involve an instance in which system 130 assumes the patient is diligently following the time run-outs to meet the deadlines or time expirations. It may occur however that the patient becomes distracted or forgets that the fill time is underway. In either case, it is likely that the patient will have left connector 118 positioned in its holding port of docking unit 140. The modification via block 186b assumes that once the patient has undertaken to remove connector 118 from docking unit 140, that the patient will thereafter diligently connect connector 118 to the transfer set for filling. Here, proximity sensor 148 senses a removal of connector 118 from the docking unit, sends a signal to controller 42, which commands Tx/Tc 41 to send a corresponding wireless signal to Rx/Tc 141 of transfer set 150, which is routed to controller 142, which then initiates the fill duration run-out sequence indicated at block 188. If space permits, proximity sensor 148 can be located alternatively in transfer set 150, which then looks for the presence of connector 118 to begin the fill duration run-out. In either case, method 180 is immune to patient delay in removing connector 118 from docking unit 140.

[00121] It should be understood that various changes and modifications to the presently preferred embodiments described herein will be apparent to those skilled in the art. Such changes and modifications can be made without departing from the spirit and scope of the present subject matter and without diminishing its intended advantages. It is therefore intended that such changes and modifications be covered by the appended claims.

CLAIMS

The invention is claimed as follows:

- 1. A peritoneal dialysis system comprising:
 - a dialysis fluid container associated with an identifier;
 - a reader for reading the identifier;
 - a signal transmitter for communicating data based on the identifier; and
- a patient transfer set for semi-permanent connection to a patient, the patient transfer set including:
 - (i) a signal receiver for receiving the data sent by the signal transmitter,
 - (ii) an output device for communicating with the patient, and
 - (iii) electronics configured to received the data and command the output device to communicate with the patient accordingly.
- 2. The peritoneal dialysis system of Claim 1, wherein the reader and the signal transmitter are housed in a common unit.
- 3. The peritoneal dialysis system of Claim 2, wherein the unit further includes a port configured to receive a patient data memory device.
- 4. The peritoneal dialysis system of Claim 2, wherein the unit includes a docking area configured to receive and hold a tube extending from the dialysis fluid container, the reader reading the identifier from the received tube.
- 5. The peritoneal dialysis system of Claim 4, the docking area configured to receive and hold a connector located at an end of the tube, the reader reading the identifier from the receiving connector.
- 6. The peritoneal dialysis system of Claim 1, wherein the signal transmitter and signal receiver are configured to communicate the data wirelessly.
- 7. The peritoneal dialysis system of Claim 1, wherein the signal transmitter and the signal receiver are each part of a transceiver for transmitting and receiving data.

8. The peritoneal dialysis system of Claim 1, wherein the identifier identifies at least one of: (i) dialysis fluid type, (ii) dialysis fluid volume, and (iii) dialysis fluid expiration data.

- 9. The peritoneal dialysis system of Claim 1, wherein the data includes a dwell duration, the patient transfer set including a start input that the patient activates upon completion of a fill from the dialysis fluid container, the dwell duration initiated upon activation of the start input.
- 10. The peritoneal dialysis system of Claim 9, the output device configured to alert the patient when the dwell duration has ended or is about to end.
- 11. The peritoneal dialysis system of Claim 9, wherein the unit is configured to determine the dwell duration based on the identifier and at least one patient-specific parameter.
- 12. The peritoneal dialysis system of Claim 1, wherein the data includes (i) a fill duration, after which it is assumed a fill from the dialysis fluid container is completed, and (ii) a dwell duration that begins after the fill duration.
- 13. The peritoneal dialysis system of Claim 12, the output device configured to alert the patient of at least one of: (i) the fill duration and (ii) when the dwell duration has ended or is about to end.
- 14. The peritoneal dialysis system of Claim 12, the output device configured to alert the patient that the fill duration is about to begin.
- 15. The peritoneal dialysis system of Claim 12, wherein the unit is configured to determine the dwell duration based on the identifier and at least one patient-specific parameter.

16. The peritoneal dialysis system of Claim 12, which includes a sensor positioned adjacent to the reader, the sensor sensing the presence or absence of a member bearing the identifier, and which is configured to begin a running of the fill duration upon a sensed removal of the member and the identifier from the reader.

- 17. The peritoneal dialysis system of Claim 1, the data including solution type, solution volume and at least one patient-specific parameter, the electronics configured to determine at least one of a fill duration, a dwell duration and a start drain time from the data.
 - 18. A peritoneal dialysis system comprising:
 - a dialysis fluid container associated with an identifier;
 - a reader for reading the identifier;
 - a signal transmitter configured to communicate a dwell duration determined at least in part from the identifier; and
 - a patient transfer set for semi-permanent connection to a patient, the patient transfer set including:
 - (i) a signal receiver for receiving the dwell duration sent by the signal transmitter; and
 - (ii) an output device configured to communicate the dwell duration to the patient.
- 19. The peritoneal dialysis system of Claim 18, the patient transfer set further including an input device and electronics configured to begin a running of the dwell duration upon a patient activating the input.
- 20. The peritoneal dialysis system of Claim 18, the identifier identifying at least one of (i) dialysis fluid type and (ii) dialysis fluid volume, which are used to determine the dwell duration in combination with at least one patient specific parameter.
- 21. The peritoneal dialysis system of Claim 18, which includes a unit housing the reader and the signal transmitter, the unit further including electronics configured to determine the dwell duration from the identifier.

22. The peritoneal dialysis system of Claim 18, the signal transmitter further configured to communicate a fill duration, after which it is assumed that a fill from the dialysis fluid container is completed, the dwell duration beginning automatically after the fill duration.

- 23. A peritoneal dialysis system comprising:
 - a dialysis fluid container associated with an identifier;
 - a reader for reading the identifier;
- a signal transmitter configured to communicate (i) a fill duration, after which it is assumed that a fill from the dialysis fluid container is completed and (ii) a dwell duration based, at least in part, on the identifier;
- a sensor positioned to sense the presence of a member bearing the identifier; and
- a patient transfer set for semi-permanent connection to a patient, the patient transfer set including a signal receiver for receiving the fill duration and the dwell duration, the patient transfer set configured to begin a running of the fill duration upon receiving a signal from the transmitter indicating that the sensor sensed a removal of the member from a location of the reader and the sensor.

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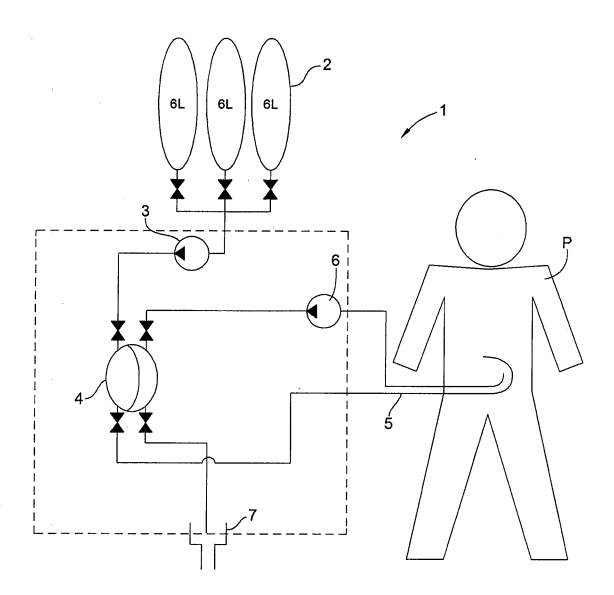


FIG. 1 PRIOR ART

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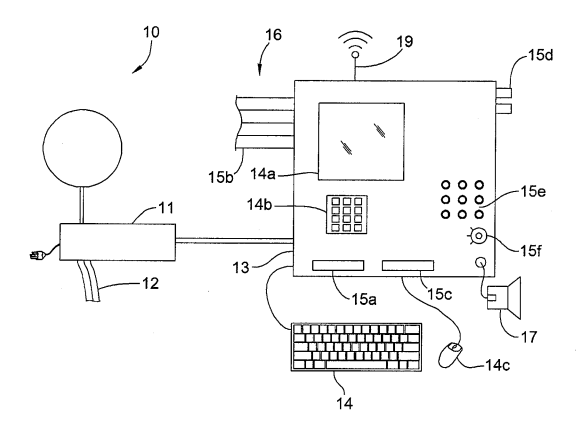
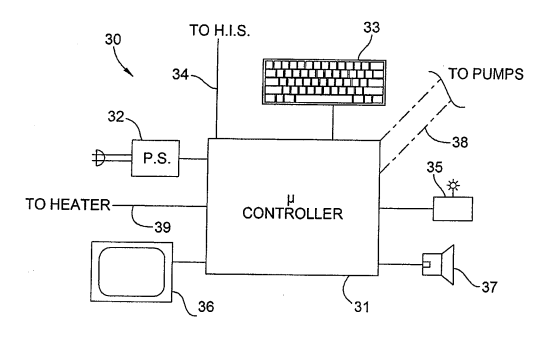


FIG. 2



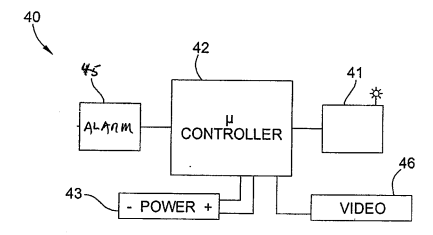


FIG. 3

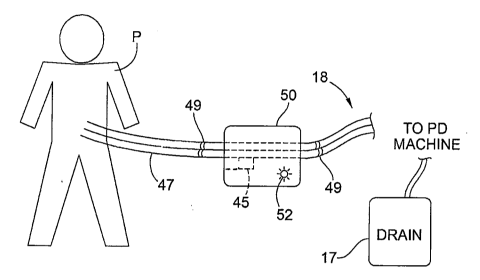


FIG. 4a

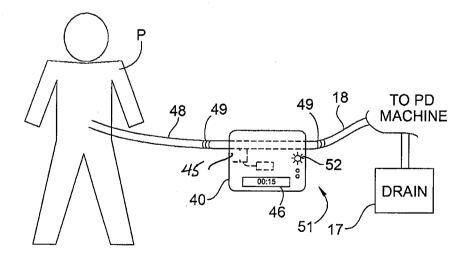
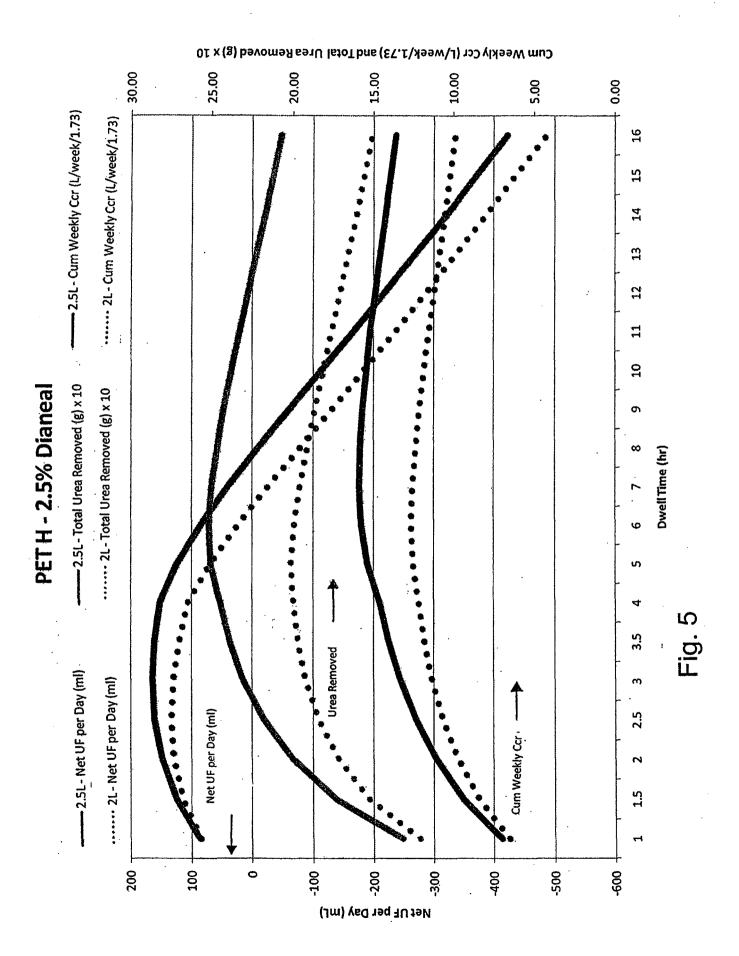
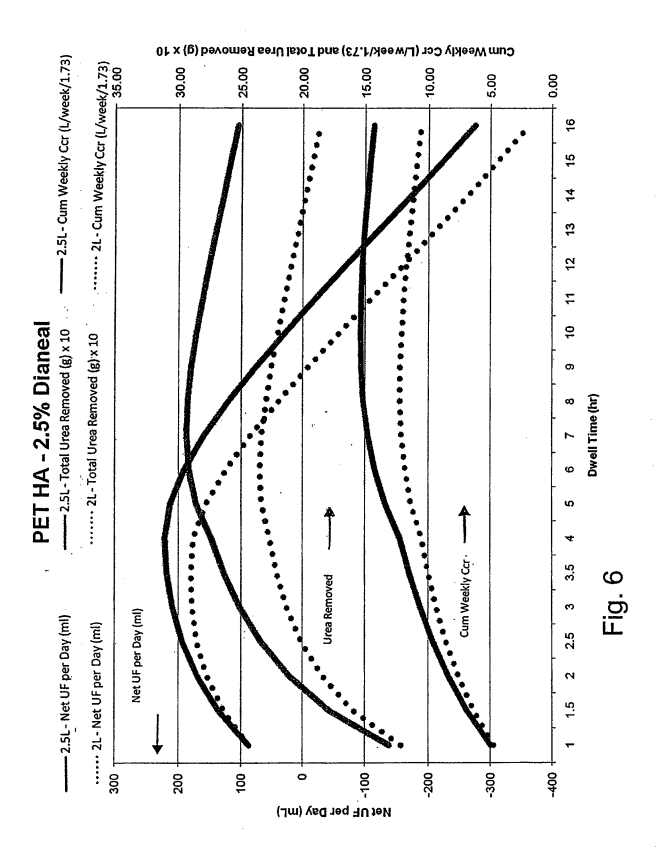
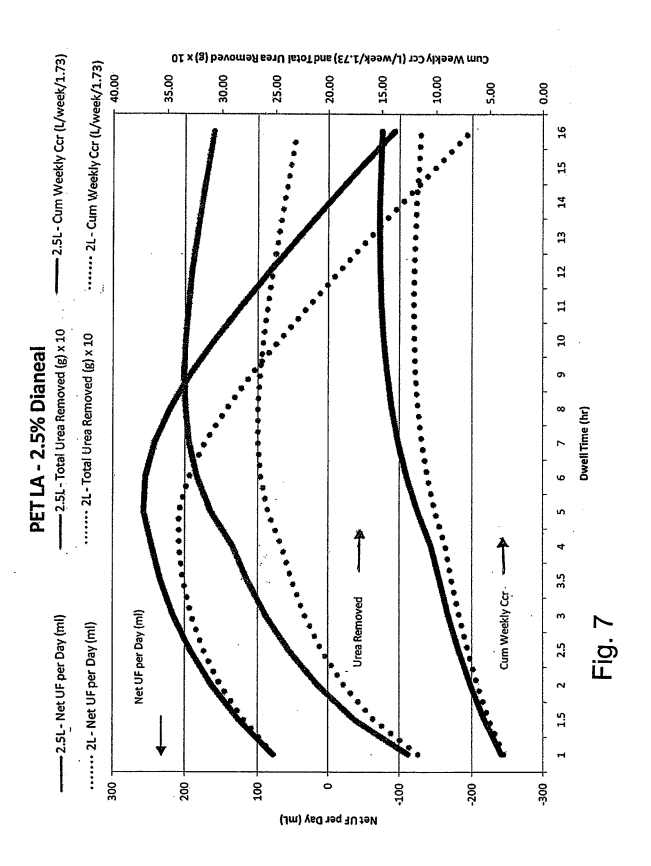
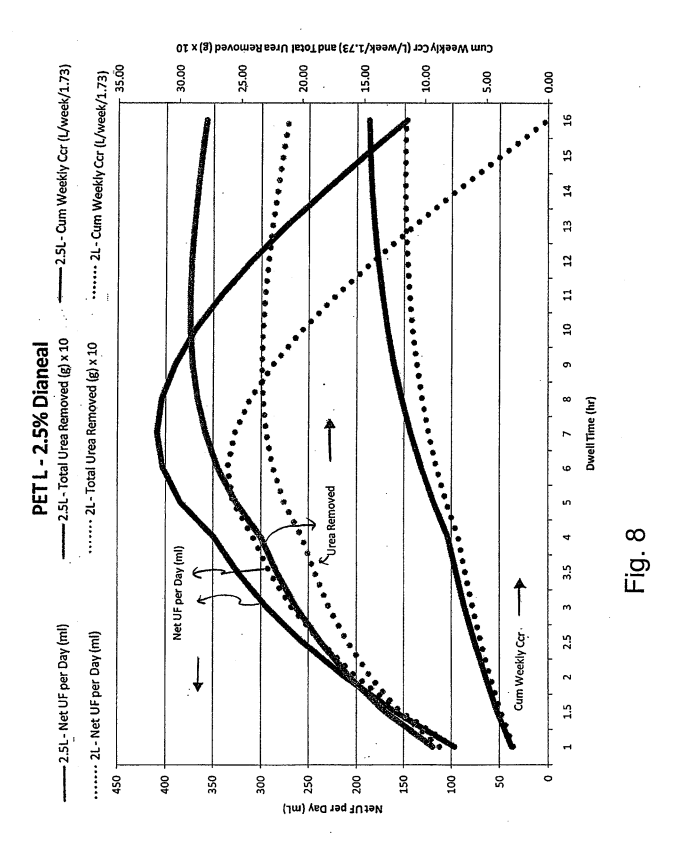


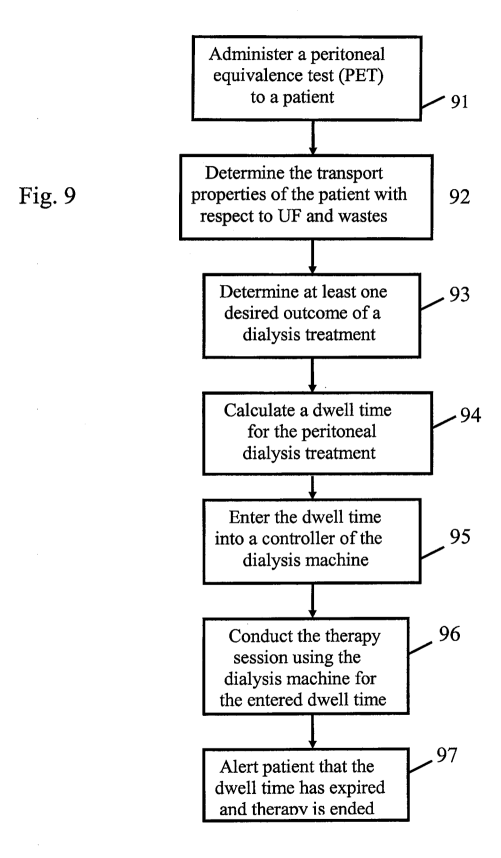
FIG. 4b



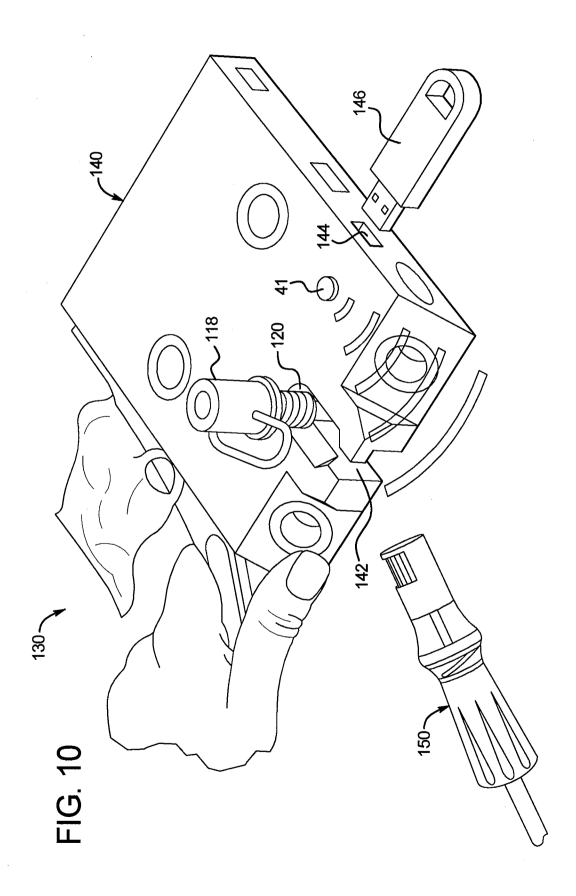


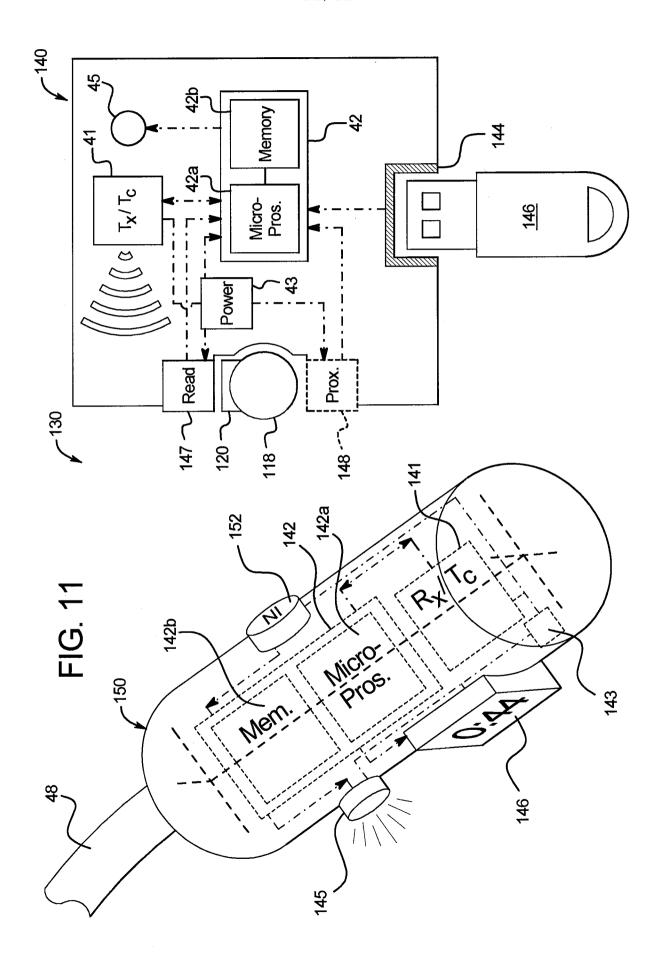


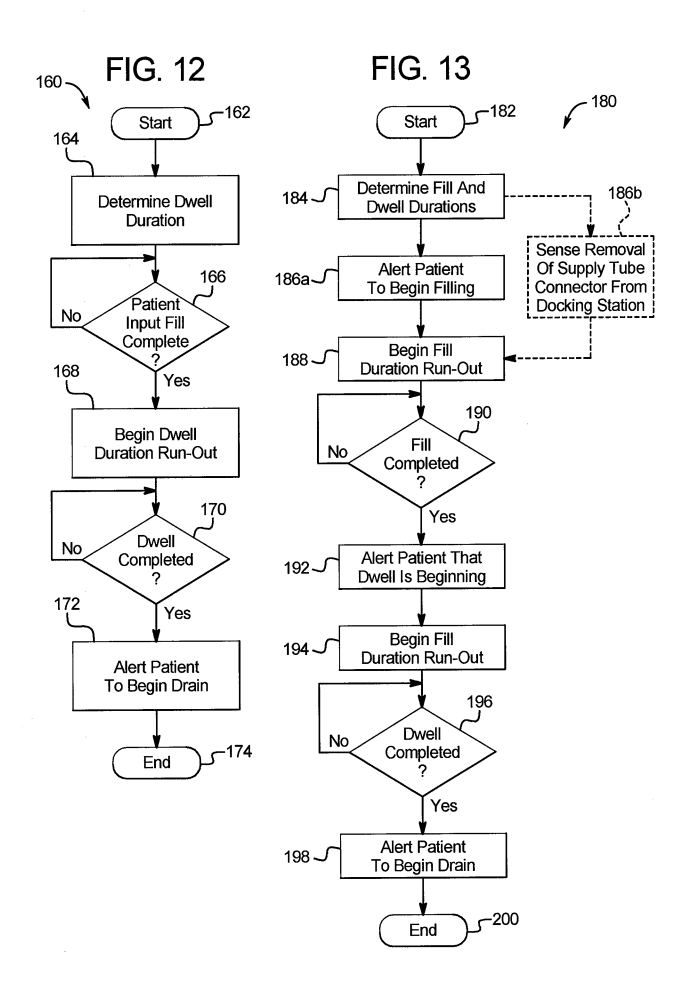




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INTERNATIONAL SEARCH REPORT

International application No PCT/US2009/042103

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AUU. A	A61M205/35 A61M205/60		
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Electronic da	ata base consulted during the international search (name of data bas	e and, where practical, search terms used)	
EPO-In	ternal		
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to claim No.
		, , , , , , , , , , , , , , , , , , ,	, iolovali to claim (40.
X	EP 1 872 814 A (DEBIOTECH SA [CH] 2 January 2008 (2008-01-02))	1,2,4-23
Υ	the whole document		3
Υ	WO 2008/027967 A (FRESENIUS MEDIC HOLDING [US]; PLAHEY KULWINDER S TRI [) 6 March 2008 (2008-03-06) paragraph [0025] - paragraph [002 figure 2	[US]; LY	3
А	US 2001/040127 A1 (DONIG RAINER [DOENIG RAINER [DE] ET AL) 15 November 2001 (2001-11-15) the whole document	DE] ET AL	1-23
r	<u> </u>		
Furt	her documents are listed in the continuation of Box C.	X See patent family annex.	
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Date of the	actual completion of the international search	Date of mailing of the international sea	rch report
2	25 August 2009	31/08/2009	
Name and	mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk Tel. (+31–70) 340–2040, Fax: (+31–70) 340–3016	Authorized officer Louarn, Arzhur	

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No PCT/US2009/042103

Patent document cited in search report		Publication date	•	Patent family member(s)	Publication date
EP 1872814	Α	02-01-2008	NONE		
WO 2008027967	A	06-03-2008	CA EP	2007289206 A1 2661670 A1 2059277 A1 2008097283 A1	06-03-2008 06-03-2008 20-05-2009 24-04-2008
US 2001040127	A1	15-11-2001	NONE		