Abstract: A device for regulating cerebrospinal fluid (10) in a cerebrospinal fluid space (22) includes a cerebrospinal conduit (20) having a distal end (20a) for insertion into the cerebrospinal fluid space in fluid communication with the cerebrospinal fluid (18). An actively oscillatably changeable sealed fluid volume (17) can be in fluid communication with the cerebrospinal conduit. The changeable sealed fluid volume can be in a sealed fluid path extending to the distal end of the cerebrospinal conduit and is capable of actively oscillating in a changing fluid volume size for oscillating the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space.
DEVICE FOR INCREASING CEREBRAL BLOOD FLOW

RELATED APPLICATION

This application claims the benefit of U.S. Provisional Application No. 61/001,795, filed on November 2, 2007. The entire teachings of the above application are incorporated herein by reference.

BACKGROUND

It has been determined that the cerebrospinal fluid (CSF) environment of the brain is important in regulating, in a negative and positive fashion, cerebral blood flow. Control of the pressure and flow pulsatility of CSF can result in improved blood flow in the brain secondary to decreasing cerebrovascular resistance and increasing cranial compliance. In many situations involving brain injury, whether due to stroke, trauma or other causes, a major effort in an intensive care unit (ICU) setting is to prevent secondary, extended brain injury which results from ischemia caused by decreased cerebral blood flow. These efforts can involve increasing systemic arterial pressure, or decreasing intracranial pressure (ICP) to ultimately increase cerebral perfusion. In the case of trauma and stroke for example, large hemi-craniotomies have been performed to decrease intracranial pressure and increase brain compliance allowing more blood flow. Another alternative is the placement of a ventricular catheter to allow drainage of cerebral spinal fluid, allowing more space for blood inflow and increasing brain compliance. Unfortunately, the former technique is a large procedure and has complications. Ventricular catheterization is limited by the amount of fluid which can be drained, also resulting in slit ventricles completely drained without further advantage.
SUMMARY

The present invention can provide a device for regulating cerebrospinal fluid in a cerebrospinal fluid space including a cerebrospinal conduit having a distal end for insertion into the cerebrospinal fluid space in fluid communication with the cerebrospinal fluid. An actively oscillatably changeable sealed fluid volume can be in fluid communication with the cerebrospinal conduit. The changeable sealed fluid volume can be in a sealed fluid path extending to the distal end of the cerebrospinal conduit and is capable of actively oscillating or modulating in a changing fluid volume size for oscillating or modulating the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space.

In particular embodiments, the device can include an actuator coupled to the changeable sealed fluid volume for oscillating the changing fluid volume size of the changeable sealed fluid volume. A sensor system can sense conditions of a patient. The sensor system can be in communication with the actuator for controlling operation of the actuator. The sensor system can include a control system for controlling operation of the actuator. Operation of the actuator can be synchronized with a biorythm of the patient. The biorythm can be related to the patient's heart. The actuator can be synchronized with a signal such as an ECG signal, a pulse signal and a pressure signal. The sensor system can include a sensor for sensing within the cerebrospinal fluid space for forming a feed back loop to control the level of cerebrospinal fluid within the cerebrospinal fluid space. A fluid storage container can be coupled to the changeable sealed fluid volume and controllably fluidly isolated from the changeable sealed fluid volume. In one embodiment, the changeable sealed fluid volume can include a movable piston, and in another embodiment, a bellows device. In other embodiments, the changeable sealed fluid volume can include a deformable fluid tight membrane on which the actuator is capable of applying an oscillating force for oscillating the changing fluid volume size of the changeable sealed fluid volume. The changeable sealed fluid volume can be within a container. The container can have a deformable fluid tight membrane with a first surface that defines a boundary forming at least a portion of the changeable sealed fluid volume. The deformable membrane is capable of deforming to oscillate the changing fluid volume size of the changeable sealed fluid volume.
The deformable membrane can have a second surface which is fluidly isolated from the first surface. Application of an oscillating force on the second surface of the deformable membrane is capable of deforming the deformable membrane to oscillate the changing fluid volume size of the changeable sealed fluid volume. The container can have a first port in communication with the changeable sealed fluid volume. The cerebrospinal conduit can be coupled to the first port. The container can also have a second port in communication with the second surface of the deformable membrane. In one embodiment, the deformable membrane can be a diaphragm extending within the container. In another embodiment, the deformable membrane can be a balloon positioned within the container. An oscillating pump can be coupled to the second port of the container for providing oscillating fluid pressure to the second surface of the deformable membrane.

The present invention can also provide a device for regulating cerebrospinal fluid in a cerebrospinal fluid space including a cerebrospinal conduit having a distal end for insertion into the cerebrospinal fluid space in fluid communication with the cerebrospinal fluid. An actively oscillatable changeable sealed fluid volume can be in fluid communication with the cerebrospinal conduit. The changeable sealed fluid volume can be in a sealed fluid path extending to the distal end of the cerebrospinal conduit and is capable of actively oscillating or modulating in a changing fluid volume size for oscillating or modulating the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space. An actuator can be coupled to the changeable sealed fluid volume for oscillating the changing fluid volume size of the changeable sealed fluid volume. A control system can control operation of the actuator and can be synchronized with a biorythm.

The present invention also provides a method of regulating cerebrospinal fluid in a cerebrospinal fluid space. A distal end of a cerebrospinal conduit can be inserted into the cerebrospinal fluid space in fluid communication with the cerebrospinal fluid. An actively oscillatable changeable sealed fluid volume can be coupled in fluid communication with the cerebrospinal conduit. The changeable sealed fluid volume can be in a sealed fluid path extending to the distal end of the cerebrospinal conduit. The changeable sealed fluid volume can be actively oscillated or modulated in a changing fluid volume size for oscillating or modulating
the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space.

In particular embodiments, the changing fluid volume size of the changeable sealed fluid volume can be oscillated with an actuator coupled to the changeable sealed fluid volume. Conditions of a patient can be sensed with a sensor system. The sensor system can be in communication with the actuator for controlling operation of the actuator. Operation of the actuator can be controlled with a control system associated with the sensor system. The actuator can be operated in synchronization with a biorythm of the patient. The biorythm can be related to the patient's heart. The actuator can be synchronized with a signal such as an ECG signal, a pulse signal and a pressure signal. The level of cerebrospinal fluid within the cerebrospinal fluid space can be controlled with a sensor within the cerebrospinal fluid space that forms a feedback loop. A fluid storage container can be coupled to the changeable sealed fluid volume. The fluid storage container can be controllably fluidly isolated from the changeable sealed fluid volume. In one embodiment, the changing fluid volume size of the changeable sealed fluid volume can be oscillated with a movable piston, and in another embodiment, with a bellows device. In other embodiments, the changeable sealed fluid volume can include a deformable fluid tight membrane. An oscillating force can be applied on the deformable membrane for oscillating the changing fluid volume size of the changeable sealed fluid volume. The changeable sealed fluid volume can be within a container. The container can have a deformable fluid tight membrane with a first surface that defines a boundary forming at least a portion of the changeable sealed fluid volume. The deformable membrane can be deformed to oscillate the changing fluid volume size of the changeable sealed fluid volume. The deformable membrane can have a second surface which is fluidly isolated from the first surface. An oscillating force can be applied on the second surface of the deformable membrane for deforming the deformable membrane to oscillate the changing fluid volume size of the changeable sealed fluid volume. The container can have a first port in communication with the changeable sealed fluid volume, and a second port in communication with the second surface of the deformable membrane. The cerebrospinal conduit can be coupled to the first port. In one embodiment, a
diaphragm can be extended within the container as the deformable membrane. In another embodiment, a balloon can be positioned within the container as the deformable membrane. Oscillating fluid pressure can be provided to the second surface of the deformable membrane with an oscillating pump coupled to the second part of the container.

The present invention can provide use of a device including a cerebrospinal conduit having a distal end for insertion into a cerebrospinal fluid space in fluid communication with a cerebrospinal fluid. An actively oscillatably changeable sealed fluid volume can be in fluid communication with the cerebrospinal conduit. The changeable sealed fluid volume can be in a sealed fluid path extending to the distal end of the cerebrospinal conduit and is capable of actively oscillating or modulating in a changing fluid volume size for oscillating or modulating the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space for regulating the cerebrospinal fluid in the cerebrospinal fluid space.

In embodiments of the present invention, through alteration of CSF space volume in a cardiac synchronous manner, intracranial pressure and flow pulsation can be controlled and cranial compliance and cerebrovascular impedent be changed. Embodiments of the present invention device can be used, for example, in a patient with a decreased cerebral blood flow in an ICU setting and, who can also be candidate for other CSF space catheterization to an external bag. The device can be part of an externalized catheter and bag system and no further invasive procedure is needed. The device can maintain the closed CSF drainage system and allow CSF removal as needed.

Embodiments of the device can be used to increase cerebral blood flow in patients with elevated intracranial pressure, and can be used in a patient (e.g., an ICU patient) who is undergoing external CSF drainage using, for example, a standard ventricular catheter or any other catheterization of a CSF space. It may be used in patients with a variety of decreased cerebral blood flow states including those induced by trauma, stroke, vasospasm, hydrocephalus or congestive heart failure. It can be applied as a device connected to a standard external ventricular
drainage catheter and integrated between the standard catheter and a standard external drainage bag.

**BRIEF DESCRIPTION OF THE DRAWINGS**

The foregoing will be apparent from the following more particular description of example embodiments of the invention, as illustrated in the accompanying drawings in which like reference characters refer to the same parts throughout the different views. The drawings are not necessarily to scale, emphasis instead being placed upon illustrating embodiments of the present invention.

FIG. 1 is a schematic drawing of an embodiment of a device or system for regulating cerebrospinal fluid.

FIGs. 2 and 3 are schematic drawings of the operation of one embodiment of a device or system for regulating cerebrospinal fluid having an oscillating sealed fluid volume with a diaphragm.

FIGs. 4 and 5 are schematic drawings of another embodiment of a device or system for regulating cerebrospinal fluid having an oscillating sealed fluid volume with a balloon.

FIG. 6 is a schematic drawing of another embodiment of an oscillating sealed fluid volume.

FIG. 7 is a schematic drawing of yet another embodiment of an oscillating sealed fluid volume.

FIGs. 8 and 9 are schematic drawings of still other embodiments of oscillating sealed volumes.

**DETAILED DESCRIPTION**

Referring to FIG. 1, in one embodiment of the present invention, cerebrospinal fluid regulating device or system 10, can regulate cerebrospinal fluid 18 within a cerebrospinal fluid space or volume 22, such as in the head or spine of a patient. Device 10 can include a cerebrospinal conduit or catheter 20, which has an open distal end 20a that can be inserted into the cerebrospinal fluid space 22. The
cerebrospinal catheter 20 is a fluid communication with an actively oscillatably changeable sealed fluid supply, reservoir, chamber or volume 17 via port A1. A drainage or storage container or bag 30 can be connected to the volume 17 via port A2 by a conduit or tube 24, a valve 26, and a conduit or tube 28. The valve 26 can be a 3-way stopcock, or other suitable shut off valve device or member, manual or automatically controlled, and in some embodiments can be a clamp. The valve 26 can be opened to allow drainage of cerebrospinal fluid 18 from the cerebrospinal fluid space 22, and into the drainage bag 30 via catheter 20 and volume 17. Once the valve 26 is closed, the volume 17 is isolated from or sealed off from fluid communication with the drainage bag 30 and is in sealed fluid communication with the catheter 20, and the cerebrospinal fluid 18 therein.

The volume 17 can be oscillated or modulated by an actuator 29 that is connected or coupled to volume 17 by a coupling member 31. The volume 17 can be fluid and biologically sealed tight relative to the actuator 29. The actuator 29 can be a pump such as a piston, rotary, centrifugal, peristaltic, roller pump, etc., or a rotary or linear actuator, and can apply or cause forces or pressure on the volume 17 for changing the fluid volume size of the volume 17. In some embodiments, the actuator 29 and volume 17 can be separate units, and in other embodiments, one unit. Since the volume 17 is in a sealed fluid communication path extending through catheter 20 to the cerebrospinal fluid space 22, a change in size to be a smaller fluid volume size can force cerebrospinal fluid 18 from or within the volume 17 and catheter 20 fluid path, into cerebrospinal fluid space 22. A change in size of volume 17 to be a larger fluid volume size, can draw cerebrospinal fluid 18 from the cerebrospinal fluid space 22 into the volume 17 and catheter 20 fluid path. The actuator 29 can be oscillated or modulated to oscillate or modulate the volume 17, which in turn can oscillate or modulate cerebrospinal fluid 18 in and out of the distal end 20a of the catheter 20 and the cerebrospinal fluid space 22, which can oscillate or modulate fluid pressure therein. Such oscillation or modulation can be slow or periodic, to increase or decrease pressure in the cerebrospinal fluid space 22 for long periods of time, for example, hours, days, or weeks, or can be multiple or many times per minute on a continual basis. Drawing cerebrospinal fluid 18 from the
cerebrospinal fluid space 22 can reduce pressure in the cerebrospinal fluid 22 and can increase blood flow, for example cerebral blood flow.

The actuator 29 and volume 17 can be oscillated or modulated continuously in synchronization with aspects, conditions or biorhythms of a patient to aid, assist or increase blood flow. Examples of some biorhythms can be heart rate, cardiac cycle, blood pressure pulse, breathing or respiratory rate. The actuator 29 can be connected to and controlled by signals from a monitor or controller 36 via a control line 37, which can be a physical electrically connected line, or a wireless connection. The monitor 36 can have a sensor 35 connected to a control line 39 (wireless connection or physical electrical line) which can monitor a patient's biorhythm source 33, for example, the heart. In one embodiment, the monitor or controller 36 can be or include an electrical cardiogram (ECG or EKG) monitor for monitoring heart rate or heart activity. The actuator 29 and volume 17 can be oscillated in synchronization with the heart and/or related electrical signals to increase blood flow. Oscillation of the cerebrospinal fluid 18 can increase and decrease pressure in the cerebrospinal fluid 22 in an alternating manner, which can allow blood to flow more easily through blood vessels around or near cerebrospinal fluid space 22. The alternating pressure can in some cases form a sort of pumping action. In another embodiment, the monitor or controller 36 can be or include an intra-cranial pressure monitor which can be connected by a control line 27 (wireless or physical) to a sensor 25 in or near the cerebrospinal fluid space 22, for sensing pressure in or around the cerebrospinal fluid space 22 and providing related electrical signals, thereby enabling sensing of the rhythm or timing of blood flow in the region. Sensor 25 can be an intracranial sensor, and can act as a blood pulse sensor or a fluid pressure sensor. The sensor 25 can be part of a feed back loop for controlling actuator 29 and controlling the pressure or fluid level within the cerebrospinal fluid space 22 to a desired level. Some embodiments can include a pressure sensor 13a for measuring blood pressure, which can be connected to monitor or controller 36 by a wireless or physical line 13. In addition, an attachable pulse sensor 21a for attaching to appendages or other suitable body parts, such as a finger, toe, or ear pulse sensor, can be included for measuring blood pressure pulses from a body part or appendage 23 such as a finger, toe or ear. The sensor 21a can be connected to
monitor or controller 36 by a wireless or physical line 21. In particular embodiments, cerebrospinal fluid 18 can be removed from the cerebrospinal fluid space 22 during systole to decrease intracranial pressure when blood flow is maximum, and deliver the cerebrospinal fluid 18 into the cerebrospinal fluid space 22 during diastole. Such a pumping action can be tied to the heart or heart beat, and synchronized with the ECG or EKG, blood pressure pulse, or signals associated therewith, or other signals tied to the cardiac cycle.

In some embodiments, the monitor or controller 36 and/or actuator 29 can include controls or software that can control the frequency, timing and duration of the operation of the actuator 29 to form a desired oscillating or modulating waveform depending upon the biorhythm sensed, its sensing location, or user input. For example, when the actuator 29 is a pump, there can be a delay between the pumping action waveform and the waveform of a biorhythm, such as a cardiac cycle, as well as morphology of the pumping waveform and duration of each state of the pump. The waveform of the pump 34 and resulting oscillation or modulation of cerebrospinal fluid 18, can be consistent in time, frequency, magnitude, shape (ramp up/down), etc., or can have variations either in a fixed pattern, or include random variations in response to sensed conditions or changes thereof. The monitor or controller 36, can be connected to or include other controllers, monitors or equipment, such as standard hospital monitors and equipment. In addition, the monitor or controller can be a standard hospital monitor or equipment. If desired, the monitor or controller 36 can be incorporated or combined with the actuator 29.

The volume 17 can be sealed in manner that can prevent pathogens surrounding volume 17 from entering the sterile fluid space within the volume 17 and can allow a pressure and pumping action to be formed therein by actuator 29. The volume 17 can have openings in the seal configured to allow the insertion or withdrawal of fluid and still be considered sealed. Although a drainage bag 30 can be connected to the volume 17 via conduit 28, valve 26 and conduit 24, and form an opening into volume 17 which can allow insertion or withdrawal of fluid, these components can also be sterile and sealed so that pathogens do not enter the volume 17 from these components. In addition, components 30, 28, 26 and 24 do not prevent pressure and pumping action to be formed in volume 17. Pressure and
pumping action of volume 17 can be aided, enabled, or enhanced by closing valve
26. In some embodiments, the catheter 20, volume 17 and drainage bag 30 together,
or portions thereof, can form a sterilized disposable 9. Since the volume 17 can be
being in fluid and sterile isolation from the actuator 29, and
oscillation of cerebrospinal fluid 18 in a sterile manner. In other embodiments, the
actuator 29 can be sterile or sterilized and can be or form part of the volume 17.

Referring to FIGs. 2 and 3, cerebrospinal fluid regulating device or system
11 is an embodiment of the present invention which has an actively oscillatably
changeable sealed fluid volume 17 that is part of an oscillating or modulating fluid
assembly 8, and can be positioned or located within an enclosed container or
housing 12. The container 12 can be a rigid housing, tank or reservoir, or
alternatively, a flexible bag. The container 12 can include a deformable membrane
14 separating the container 12 into two fluidly isolated and enclosed regions,
changeable sealed volume 17 or compartment A, and actuation region 15 or
compartment B. The deformable membrane 14 can have first 14a second 14b
surfaces on opposite fluidly isolated sides that can form or define a boundary of at
least a portion of volume 17, and region 15. The container 12 can form the
remaining portions of volume 17 and region 15. The deformable membrane 14 can
be fluid and biologically tight, and can be sealed or secured within container 12 in a
fluid and biologically tight manner. The deformable membrane 14 can be sized,
and/or formed of elastic material to enable movement to one side of a delineation
axis X or to the other side of axis X, to alternately increase and decrease the volume
sizes of volume 17 and region 15. For example, the deformable membrane 14 can
be a flexible membrane having a surface area that is larger than the cross-section of
the container 12, allowing the deformable membrane to move under an applied force
to either side of axis X, and/or can be an elastic membrane, that can resiliently
stretch or expand under an applied force to either side of axis X. In some
embodiments, the deformable membrane 14 can be formed with a resilient bias, for
example, biased for expanding into actuation region 15.

The container 12 can have three inlet/outlet ports A1, A2 and B1. Ports A1
and A2 can be in fluid communication with volume 17. Port A1 can be coupled to
conduit or catheter 20, for example a ventricular drainage catheter, and can include additional tubing and fittings. Port A2 can be coupled to drainage bag 30 via conduit 24, and valve 26. Port B1 can be in fluid communication with actuation region 15 and connected to an oscillating or modulating pump or pump system 34 by a conduit 32.

The pump 34 can serve as an actuator 29 by pumping an actuation fluid 16, such as air or other gases, or a suitable liquid, to and from pump 34 and actuation region 15, through a conduit 32 that serves as a coupling member 31. Pump 34 can be a suitable pump such as a piston, rotary centrifugal, roller, peristaltic pump, etc. The actuation fluid 16 can be a biocompatible liquid such as saline.

Referring to FIG. 2, pumping of the actuation fluid 16 into the actuation region 15 of container 12 as shown by the arrow, can apply a force or fluid pressure P on the surface 14b of deformable membrane 14 which forces the deformable membrane 14 onto the compartment A side of axis X, thereby enlarging or increasing the fluid volume size of region 15 and decreasing the fluid volume size of volume 17. This forces cerebrospinal fluid 18 out of the volume 17 into catheter 20, as shown by the arrow, so that a certain volume of cerebrospinal fluid 18 can exit the distal end 20a of catheter 20 and enter cerebrospinal fluid space 22, which can increase fluid pressure therein.

Conversely, referring to FIG. 3, when pump 34 pumps actuation fluid 16 out of the actuation region 15 as shown by the arrow, this can create a suction or pressure drop within region 15 and can allow the deformable membrane 14 to be drawn onto the compartment B side of axis X, thereby decreasing the fluid volume size of region 15, while at the same time drawing cerebrospinal fluid 18 as shown by the arrow into and increasing the fluid volume size of volume 17. In some embodiments, fluid pressure P of cerebrospinal fluid 18 can act on surface 14a of deformable membrane 14. The increase in fluid volume size of volume 17 can create a suction within volume 17 and catheter 20, causing cerebrospinal fluid 18 in the cerebrospinal fluid space 22 to be drawn into the distal end 20a of the catheter 20, which can decrease or reduce fluid pressure in the cerebrospinal fluid space 22. The deformable membrane 14 can form a fluid and sterile or biological barrier which can allow volume 17 to be sterilized and continue to remain and operate in a
sterile manner whether or not the pump 34 and actuation fluid 16 are sterile. The pump 34 can alternately physically squeeze or compress, and then expand the fluid volume size of volume 17.

The pump 34 can be controlled by monitor or controller 36 to oscillate or modulate actuation fluid 16 in and out of the actuation region 15, thereby oscillating or modulating the deformable membrane 14 and therefore cerebrospinal fluid 18, in and out of the cerebrospinal fluid space 22, which can oscillate or modulate fluid pressure therein. The monitor or controller 36 can be connected by control lines to any or all of sensors 13a, 21a, 35 and 25 as in FIG. 1. In addition, pressure sensors 19a and 19b can be positioned within volume 17 and actuation region 15 for monitoring the pressure of cerebrospinal fluid 18 and actuation fluid 16. Sensors 19a and 19b can be connected to monitor or controller 36 by physical or wireless control lines. The oscillation or modulation sequence, timing duration, magnitude, waveform, etc., can be affected by the pressures sensed by sensors 19a and 19b. The location of the sensors 19a and 19b can be varied, for example, can be located near ports A1 and B1, or in conduits 20 and 32. In some embodiments, sensors that sense volume can be employed, or sensors that sense a parameter associated with volume, for example, the position of deformable membrane 14. Although a pump 34 has been shown connected to actuation region 15 by a conduit 32, in other embodiments, a reciprocating piston of an actuator 29 can extend into region 15 for displacing the actuation fluid 16 and deformable membrane 14 in an oscillating or modulating manner.

Referring to FIGs. 4 and 5, cerebrospinal fluid regulating device or system 40 differs from device 11 in that oscillating or modulating fluid assembly 42 is employed in place of oscillating fluid assembly 8. Assembly 42, as with assembly 8, also includes a container or housing 12 with ports A1, A2 and B1. A deformable membrane 34 in the shape or configuration of a bag, bladder or balloon 38 can be positioned within container 12, and connected and sealed in a fluid tight manner in fluid communication with port B1 for fluid communication with pump 34 via conduit 32. The interior of balloon 38 can form actuation region 15, and the areas within container 12 outside balloon 38 can form volume 17. Balloon 38 can be a fluid and sterile or biologically tight barrier, and the interior, fluidly and biologically
isolated from volume 17, to allow volume 17 to be sterilized and remain sterile. Ballon 38 can be formed of resilient or stretchable material, but alternatively, can be non stretchable. Pumping of actuation fluid 16 into and out of the balloon 38 by pump 34 can expand and deflate balloon 38 to increase and decrease the fluid volume size of balloon 38.

Referring to FIG. 4, pumping actuation fluid 16 into balloon 38 and actuation region 15 as shown by the arrow, can apply a force or fluid pressure P on surface 14b on the interior of balloon 38. Since surface 14b is the inner surface of balloon 38, the balloon 38 expands in volume size within the container 12. The volume 17 is in the regions of container 12 not occupied by balloon 38, so that as balloon 38 expands, volume 17 decreases in fluid volume size, which forces cerebrospinal fluid 18 out of volume 17 into catheter 20, and out the distal end 20a into cerebrospinal space 22, as shown by the arrow, which can increase fluid pressure therein.

Conversely, referring to FIG. 5, pumping actuation fluid 16 out of balloon 38 and actuation region 15 as shown by the arrow, can create a suction or pressure drop within the balloon 38 and against surface 14b, which causes or allows collapse or deflation of the balloon 38, thereby decreasing the internal fluid volume size of the balloon 38 and actuation region 15. The fluid pressure P of cerebrospinal fluid 18 within volume 17 on the exterior of balloon 38 on the surface 14a, can also aid in the collapse of balloon 38. As balloon 38 and actuation region 15 decreases in fluid volume size within container 12, the fluid volume size of volume 17 increases and can create a suction within volume 17 and catheter 20, drawing cerebrospinal fluid 18 into volume 17 as shown by the arrow, causing cerebrospinal fluid 18 in the cerebrospinal fluid space 22 to be drawn into the distal end 20a of catheter 20, which can decrease or reduce the fluid pressure in the cerebrospinal fluid space 22. Device 40 can be operated and controlled in a similar manner as device 11.

Referring to FIG. 6, in other embodiments, actively oscillatable changeable sealed fluid volume 17 can be incorporated in a bellows device 50 which can have a collapsible and expandable bellows membrane 54 extending or positioned between pivoting arms 52. The membrane 54 can extend between and be fluidly sealed to the arms 52 to form volume 17 in the space therebetween, or can be a flexible fluid and biological tight container or bag forming volume 17, positioned between and
connected to arms 52. The membrane 54 can include ports A1 and A2 for connection to catheter 20 and drainage bag 30. Movement of the arms 52 towards and away from each other as indicated by the arrow can increase and decrease the fluid volume size of volume 17 to cause oscillation or modulation of cerebrospinal fluid 18. The arms 52 can pivot about a pivot point 56, and can be moved in an arc by actuator 29 via coupling member 31. The actuator 29 can be for example, a rotary motion actuator, such as a servomotor coupled to the pivot point 56, or a linear actuator such as a piston or fluid cylinder coupled to at least one arm 52. Actuator 29 can also be other suitable types of actuating devices. Bellows device 50 can be a disposable unit or can be a permanent unit that is sterilized.

Referring to FIG. 7, bellows device 60 is another embodiment which differs from bellows device 50 in that collapsible and expandable bellows membrane 64 is positioned between members 62 that can move linearly towards and away from each other as indicated by the arrows for increasing and decreasing the fluid volume size of volume 17. Membrane 64 can be sealed to members 62 or can be a bag connected to members 62. Actuator 29 can be connected to bellows device 60 by coupling member 31 for moving members 62 relative to each other. Actuator 29 can be a linear motion actuator. In some embodiments of FIGs. 6 and 7, the membranes 54 or 64, in a bag configuration, can be sterile disposables and can be attachable between arms 52 or members 62. In other embodiments, the arms 52 or members 62 and the membranes 54 or 64, can form sterile disposable units that can be connected to actuator 29. In addition, the actuator 29 can be part of the bellows devices 50 or 60.

Referring to FIG. 8, oscillating or modulating fluid assembly 70 is an embodiment that differs from assembly 8 in that a reciprocating piston 72 is positioned and movably or slidably sealed within container 12. Piston 72 can be fluidly sealed to the interior walls of container 12 with a sliding seal 74. The container 12 and piston 72 can be cylindrical, and the sliding seal 74 can be an annular sealing ring positioned within an annular groove. If desired, more than one seal 74 can be employed, or other suitable sealing configurations. When actuation fluid 16 is pumped by pump 34 into actuation region 15, the piston 72 moves, increasing the fluid volume size of actuation region 15 and decreasing the fluid
volume size of volume 17, thereby delivering cerebrospinal fluid 18 into cerebrospinal fluid space 22, which can increase fluid pressure therein. Conversely when pump 34 pumps actuation fluid 16 out of region 15, the piston 72 moves to decrease the fluid volume size of region 15 and increases the fluid volume size of volume 17, thereby removing cerebrospinal fluid 18 from cerebrospinal fluid space 22, which can decrease or reduce fluid pressure therein. Oscillation or modulation of cerebrospinal fluid 18 fluid pressure can be achieved by reciprocating piston 72.

Referring to FIG. 9, oscillating or modulating fluid assembly 75 is an embodiment that differs from assembly 70 in that reciprocating piston 72 can be connected to an actuator 29 by a mechanical coupling member 31, such as a reciprocating coupling rod for mechanically driving or moving the piston 72 in an oscillating or modulating manner. The actuator 29 can be a linear motion device, such as a fluid cylinder, or can be a motor which in conjunction with the coupling member 31 and the piston 72, can form a crank slider mechanism. In some embodiments, the actuator 29 and coupling rod can be incorporated into or mounted on container 12.

Referring back to FIGs. 1-3, an example of operation of an embodiment of device 11 now follows. A patient undergoes catherization for removal of cerebrospinal fluid 18 for purposes of relieving pressure and increasing cerebral blood flow, and/or for monitoring the cerebrospinal fluid 18. The catheter 20 is inserted into the cerebrospinal fluid space 22. If the catheter 20 is separate or separated from the container 12, the catheter 20 can be secured to port A1. If desired, a ligature can be used. Likewise if the drainage bag 30 is separate or separated, the drainage bag 30 and valve 26 can be connected to port A2 via conduit 24. Compartment A or volume 17 can then be filled with cerebrospinal fluid 18 from cerebrospinal fluid space 22. If desired, valve 26 can opened to drain cerebrospinal fluid 18 into drainage bag 30, and then closed. If there is not sufficient cerebrospinal fluid 18 to fill volume 17, saline solution can be used to fill volume 17. Fluid within the volume 17, whether cerebrospinal fluid 18, saline or both, is considered cerebrospinal fluid for purposes of describing operation of the devices in the present invention. When pump 34 is separate from container 12, pump 34 can be connected to port B1 via conduit 32. An ECG signal from a
standard monitoring system in the hospital, or an independently placed ECG
monitory system can be obtained from the patient, for example from sensor 35 (FIG. 1). In addition, a signal can be obtained which is related to intracranial pressure (ICP) from a standard intracranial pressure monitor used in a hospital, or a pressure sensing system placed in compartment A and/or the tubing attached to compartment A, such as with sensors 25 (FIG. 1) and 19a. One or both of the ECG and ICP signals can be used by controller 36 to regulate the movement of pump 34, and change the ICP pressure if appropriate, via modulation of actuation fluid 16 in actuation region 15, and the deformable membrane 14 located between actuation region 15 and volume 17. Actuation fluid 16 can be removed from actuation region 15 or compartment B during systoli, and added during diastoli. Modulation can be based on the cardiac cycle, respiratory cycle, ICP and/or CSF pressure monitoring, cerebral blood flow monitory, other suitable biorhythms that can have a relation to ICP, or other after standard intensive care unit monitoring systems. Device 11 can be used in an intermittent or continuous fashion, and can be used during drainage of cerebrospinal fluid 18. Devices 10 and 40 can be used similarly.

The devices in the present invention can be used for treating patients having altered intracranial compliance, decreased cerebral blood flow and/or abnormal intracranial pressure. Such conditions can occur with head injuries, aging, cerebrovascular disease, brain atrophy, post brain hemorrhage and infection, vasospasms, congestive heart failure, carotid endarterectomy, carotid occlusion/stenosis, cardiopulmonary bypass procedure, hydrocephalus, stroke, dementia, or migraine headaches. Hydrocephalus can be chronic hydrocephalus, normal pressure hydrocephalus, pseudotumor, cerebri, or slit ventricle syndrome. Stroke can be acute stroke, chronic stroke, microvascular disease, dementia, moyamoya, multiple infarct disease, posterior circulation insufficiencies or Binswanger disease. Dementia can be vascular dementia, Alzheimer’s disease, and normal pressure hydrocephalus. Migraine headaches can be pediatric migraines, adult migraines, or intractable migraines. In some conditions, increased intracranial pressure may be desired. Depending upon the condition, treatment can be temporary or long term.
While this invention has been particularly shown and described with references to example embodiments thereof, it will be understood by those skilled in the art that various changes in form and details may be made therein without departing from the scope of the invention encompassed by the appended claims.

For example, embodiments of the present invention can be used for draining, adding and/or oscillating fluid in other spaces, cavities, or fluid systems of a patient. In addition, features of the various embodiments shown and described can be combined together, or some features can be omitted.
CLAIMS

What is claimed is:

1. A device for regulating cerebrospinal fluid in a cerebrospinal fluid space comprising:
   a cerebrospinal conduit having a distal end for insertion into the cerebrospinal fluid space in fluid communication with the cerebrospinal fluid; and
   an actively oscillatably changeable sealed fluid volume in fluid communication with the cerebrospinal conduit, the changeable sealed fluid volume being in a sealed fluid path extending to the distal end of the cerebrospinal conduit and capable of actively oscillating in a changing fluid volume size for oscillating the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space.

2. The device of Claim 1 further comprising an actuator coupled to the changeable sealed fluid volume for oscillating the changing fluid volume size of the changeable sealed fluid volume.

3. The device of Claim 2 further comprising a sensor system for sensing conditions of a patient, the sensor system being in communication with the actuator for controlling operation of the actuator.

4. The device of Claim 3 in which the sensor system includes a control system for controlling operation of the actuator.

5. The device of Claim 4 in which the operation of the actuator is synchronized with a biorhythm of the patient.
6. The device of Claim 5 in which the biorythm is related to the patient's heart.

7. The device of Claim 5 in which the actuator is synchronized with a signal that is selected from the group consisting of an ECG signal, a pulse signal and a pressure signal.

8. The device of Claim 4 in which the sensor system includes a sensor for sensing within the cerebrospinal fluid space for forming a feedback loop to control the level of cerebrospinal fluid within the cerebrospinal fluid space.

9. The device of Claim 1 further comprising a fluid storage container coupled to the changeable sealed fluid volume and controllably fluidly isolated from the changeable sealed fluid volume.

10. The device of Claim 1 in which the changeable sealed fluid volume includes a movable piston.

11. The device of Claim 1 in which the changeable sealed fluid volume includes a bellows device.

12. The device of Claim 2 in which the changeable sealed fluid volume includes a deformable fluid tight membrane on which the actuator is capable of applying an oscillating force for oscillating the changing fluid volume size of the changeable sealed fluid volume.

13. The device of Claim 1 in which the changeable sealed fluid volume is within a container, the container having a deformable fluid tight membrane with a first surface that defines a boundary forming at least a portion of the changeable sealed fluid volume, the deformable membrane being capable of deforming to oscillate the changing fluid volume size of the changeable sealed fluid volume.
14. The device of Claim 13 in which the deformable membrane has a second
surface which is fluidly isolated from said first surface, whereby application
of an oscillating force on the second surface of the deformable membrane is
able of deforming the deformable membrane to oscillate the changing
fluid volume size of the changeable sealed fluid volume.

15. The device of Claim 14 in which the container has a first port in
communication with the changeable sealed fluid volume, the cerebrospinal
conduit being coupled to the first port, the container also having a second
port in communication with the second surface of the deformable membrane.

16. The device of Claim 15 in which the deformable membrane is a diaphragm
extending within the container.

17. The device of Claim 15 in which the deformable membrane is a balloon
positioned within the container.

18. The device of Claim 15 further comprising an oscillating pump coupled to
the second port of the container for providing oscillating fluid pressure to the
second surface of the deformable membrane.

19. A device for regulating cerebrospinal fluid in a cerebrospinal fluid space
comprising:
   a cerebrospinal conduit having a distal end for insertion into the
cerebrospinal fluid space in fluid communication with the cerebrospinal
fluid; and
   an actively oscillatably changeable sealed fluid volume in fluid
communication with the cerebrospinal conduit, the changeable sealed fluid
volume being in a sealed fluid path extending to the distal end of the
cerebrospinal conduit and capable of actively oscillating in a changing fluid
volume size for oscillating the cerebrospinal fluid in and out of the distal end
of the cerebrospinal conduit and cerebrospinal fluid space;
an actuator coupled to the changeable sealed fluid volume for
oscillating the changing fluid volume size of the changeable sealed fluid
volume; and
a control system for controlling operation of the actuator
synchronized with a biorhythm.

20. A method of regulating cerebrospinal fluid in a cerebrospinal fluid space
comprising:
inserting a distal end of a cerebrospinal conduit into the cerebrospinal
fluid space in fluid communication with the cerebrospinal fluid;
coupling an actively oscillatably changeable sealed fluid volume in
fluid communication with the cerebrospinal conduit, the changeable sealed
fluid volume being in a sealed fluid path extending to the distal end of the
cerebrospinal conduit; and
actively oscillating the changeable sealed fluid volume, the
changeable sealed fluid volume oscillating in a changing fluid volume size
for oscillating the cerebrospinal fluid in and out of the distal end of the
cerebrospinal conduit and cerebrospinal fluid space.

20 21. The method of Claim 20 further comprising oscillating the changing fluid
volume size of the changeable sealed fluid volume with an actuator coupled
to the changeable sealed fluid volume.

22. The method of Claim 21 further comprising sensing conditions of a patient
with a sensor system, the sensor system being in communication with the
actuator for controlling operation of the actuator.

23. The method of Claim 22 further comprising controlling operation of the
actuator with a control system associated with the sensor system.

24. The method of Claim 23 further comprising operating the actuator in
synchronization with a biorhythm of the patient.
25. The method of Claim 24 further comprising relating the biorythm with the patient's heart.

26. The method of Claim 24 further comprising synchronizing the actuator with a signal selected from the group consisting of an ECG signal, a pulse signal and a pressure signal.

27. The method of Claim 23 further comprising controlling the level of cerebrospinal fluid within the cerebrospinal fluid space with a sensor within the cerebrospinal fluid space that forms a feedback loop.

28. The method of Claim 20 further comprising coupling a fluid storage container to the changeable sealed fluid volume, the fluid storage container being controllably fluidly isolated from the changeable sealed fluid volume.

29. The method of Claim 20 further comprising oscillating the changing fluid volume size of the changeable sealed fluid volume with a movable piston.

30. The method of Claim 20 further comprising oscillating the changing fluid volume size of the changeable sealed fluid volume with a bellows device.

31. The method of Claim 21 in which the changeable sealed fluid volume includes a deformable fluid tight membrane, the method further comprising applying an oscillating force on the deformable membrane for oscillating the changing fluid volume size of the changeable sealed fluid volume.

32. The method of Claim 20 in which the changeable sealed fluid volume is within a container, the container having a deformable fluid tight membrane with a first surface that defines a boundary forming at least a portion of the changeable sealed fluid volume, the method further comprising deforming
the deformable membrane to oscillate the changing fluid volume size of the changeable sealed fluid volume.

33. The method of Claim 32 in which the deformable membrane has a second surface which is fluidly isolated from said first surface, the method further comprising applying an oscillating force on the second surface of the deformable membrane for deforming the deformable membrane to oscillate the changing fluid volume size of the changeable sealed fluid volume.

34. The method of Claim 33 in which the container has a first port in communication with the changeable sealed fluid volume, and a second port in communication with the second surface of the deformable membrane, the method further comprising coupling the cerebrospinal conduit to the first port.

35. The method of Claim 34 further comprising extending a diaphragm within the container as the deformable membrane.

36. The method of Claim 34 further comprising positioning a balloon within the container as the deformable membrane.

37. The method of Claim 34 further comprising providing oscillating fluid pressure to the second surface of the deformable membrane with an oscillating pump coupled to the second port of the container.

38. Use of a device comprising:

   a cerebrospinal conduit having a distal end for insertion into a cerebrospinal fluid space in fluid communication with a cerebrospinal fluid; and

   an actively oscillatably changeable sealed fluid volume in fluid communication with the cerebrospinal conduit, the changeable sealed fluid volume being in a sealed fluid path extending to the distal end of the
cerebrospinal conduit and capable of actively oscillating in a changing fluid volume size for oscillating the cerebrospinal fluid in and out of the distal end of the cerebrospinal conduit and cerebrospinal fluid space for regulating the cerebrospinal fluid in the cerebrospinal fluid space.
FIG. 2
FIG. 5

ECG and/or ICP Data

Oscillating Pump System

Drainage Bag

Stopcock (closed)

Patient CSF Space

Compartment B

Compartment A

Port A1 20

Port A2 18

Port B1 32

Port A1 20

16 15 12

14b 14a

14b 14a

19a

17 18

38 14

24 28

29 30

34 36

37

40

42
A. CLASSIFICATION OF SUBJECT MATTER
INV. A61M27/00

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

B. FILOS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
A61M A61B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic database consulted during the international search (name of data base and, where practical, search terms used)
EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<td>DE 299 21 166 U1 (RYCYK MANFRED [DE]) 2 March 2000 (2000-03-02) the whole document</td>
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Further documents are listed in the continuation of Box C

See patent family annex

* Special categories of cited documents
'A' document defining the general state of the art which is not considered to be of particular relevance
'E' earlier document but published on or after the international filing date
'L' document which may throw doubts on novelty claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
'O' document referring to an oral disclosure, use, exhibition or other means
'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
2 February 2009

Date of mailing of the international search report
10/02/2009

Name and mailing address of the ISA/
European Patent Office, P B 5818 Patentlaan 2 NL - 2280 HV RUISWICK
Tel (+31-70) 340-2040, Fax (+31-70) 340-3016

Authorized officer
Przykutta, Andreas
**DOCUMENTS CONSIDERED TO BE RELEVANT**

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INTERNATIONAL SEARCH REPORT

Box No. II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. [x] Claims Nos.: 20-38
   Because this matter is not required to be searched by this Authority, namely:
   Rule 39.1(iv) PCT - Method for treatment of the human or animal body by surgery

2. [ ] Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. [ ] Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1. [ ] As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. [ ] As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. [ ] As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

4. [ ] No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 

Remark on Protest

- [ ] The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.

- [ ] The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.

- [ ] No protest accompanied the payment of additional search fees.
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