A medical fluid drainage system which may be advantageously used to actively absorb excess fluid from an interstitial in a living host and to transport it from an inlet member to a non edematous body part out of an outlet member via pumps built in series, such as a distal area with functional lymphatic vessels, or directly in a lymphatic vessel, or directly in a blood vessel.
MEDICAL FLUID DRAINAGE SYSTEM

FIELD OF INVENTION

The present invention generally relates to the draining of physiological fluids. More specifically, the present invention relates to pump-based drainage systems, capable of absorbing excess fluid distributed in an interstitium (e.g. subcutaneous space) and transporting said fluid to another location such as the bloodstream, directly through a connection to the venous system or indirectly, through the lymphatic system.

BACKGROUND OF THE INVENTION

Edema is the swelling of soft tissues due to accumulation of interstitial fluid. The fluid is predominantly water, but protein and cell-rich fluid can accumulate if there is infection or lymphatic obstruction. The swelling is the result of the accumulation of excess fluid under the skin in the spaces within the tissues. All tissues of the body are made up of cells and connective tissues that hold the cells together. This connective tissue around the cells is known as the interstitium.

The body's organs have interstitial spaces where fluid can accumulate. An accumulation of fluid in the interstitial air spaces (alveoli) in the lungs occurs in pulmonary edema. In addition, excess fluid sometimes collects in what is called the third space, which includes cavities in the abdomen (abdominal or peritoneal cavity - called "ascites") or in the chest (lung or pleural cavity - called "pleural effusion").

Edema results from increased movement of fluid from the intravascular to the interstitial space, or decreased movement of water from the interstitium into the capillaries or lymphatic vessels. Increased movement of fluid from the intravascular to the interstitial space is due to increased capillary permeability that occurs in infections or as the result of toxic or inflammatory damage to the capillary walls. Edema also results from decreased movement of fluid out of the interstitial space into the capillaries or lymphatic vessels due to lack of adequate plasma oncotic pressure as in nephrotic syndrome, protein-losing enteropathy, or starvation. The lymphatic system is responsible for removing protein and white blood cells (along with some water) from the interstitium. Lymphatic obstruction causes these substances to accumulate in the interstitium.

Lymphedema is a highly disabling disease that causes swollen body limbs due to the malfunctioning of the lymphatic system. It can be inherited (primary) or it can be a consequence of cancer treatment (secondary). Moreover, it is widespread in developing countries as a result of filariasis, a disease caused by a parasitic worm infection, transmitted by mosquitoes. Lymphedema affects approximately 140 million people worldwide. Although epidemiologic data are controversial, it is estimated that in the United States around 6.8 million people have developed or have high risk of developing lymphedema. The incidence of lymphedema is estimated to be around 20% for people undergoing invasive cancer surgery, or other invasive surgeries as hip and knee replacement, cellulitis removal and coronary-artery bypass graft.
In normal conditions, lymphatic vessels absorb up to 1.5 liters per day of fluid (lymph) from the peripheral tissues and they bring it back to the blood circulation. If the lymphatic system is damaged, its drainage action is impaired, resulting in the subcutaneous accumulation of fluid in the limbs, and consequent local swelling. The accumulation of stagnant lymph causes in time dermatitis, pain, weight gain, fatigue, tissue fibrosis, loss of mobility, localized immunodeficiency and recurrent infections. Depression often occurs due to the aesthetical impairment.

As of today, lymphedema cannot be cured. However, a series of therapies are commonly performed, based on the combination of decongestion therapy, massages, lymphatic drainage and compression strategies. These treatments do not remove the cause of the problem but aim only at treating the side effects. Indeed, the disease is not cured and the side effects always come back: the patients must undergo lifelong treatments.

One possible solution for lymphedema could be replacing the function of the damaged lymphatic compartment with an implantable medical system. Implantable medical systems suitable for fluid drainage are known per se. For instance, US 8,157,792 describes a wound drainage system for draining fluid from a wound of a patient. The system includes a system that periodically increases and decreases the application of suction at a drain catheter, together with said drain catheter. WO 2009/096852 describes an implantable drain adapted to move body fluid from one part of the body to another part of the body. The drainage system enables a patient to easily move around while still being attached to the drain. The drainage system comprises a drainage system for pumping body fluid and a connecting tube and is adapted to be implanted inside the body of the patient. The system can pump body fluid from a treatment area to another part of the body where the fluid can be absorbed and transported out from the body in a normal way. The drainage system has a pump comprising a bellow and a motor may be adapted to compress the bellow and move fluid. The motor is advantageously adapted to repeat the compression at suitable time intervals whereby the drainage system is enabled to repeat the sucking and moving of fluid to substantially constantly suck fluid to the other part of the body.

WO 2014/062679 describes a medical system for moving lymphatic fluid. The medical system may include an implantable body having an inlet end region and an outlet end region represented by an implantable tubular member having one or more fluid openings formed therein. The implantable tubular member may be configured to be implanted or partially implanted in or adjacent to the lymphatic system (e.g., within a lymphatic vessel). The lymphedema medical system may also include a pump coupled to the implantable tubular member. A pump member may be positioned between the inlet end region and the outlet end region. The pump member may be configured to draw lymphatic fluid into the implantable body through the inlet end region and transfer lymphatic fluid out from the implantable body through the outlet end region.

Many other medical systems for draining a fluid are known in the art, such as those described in US 8,517,973, US 5,762,599, US 3,810,259, incorporated herein by reference.

Although such systems are suitable for draining fluid from one body area, or body cavity, to another, they would not be able to absorb and transport fluids distributed in an interstitium, as typical of edemas.
Therefore, there is a need for a system which is capable of draining fluid from an extended area and actively pumping it to reach, directly or indirectly, the bloodstream.

SUMMARY OF THE INVENTION

Accordingly, it is an object of the present invention to provide a system that is able to avoid accumulation of fluid in an interstitium (e.g. a subcutaneous space), maintaining tissue fluid homeostasis over time.
It is also an object of the invention to provide a system that actively absorbs a body fluid from multiple openings distributed along the system and that effectively transports such fluid in a desired body location (e.g., subclavian vein, saphenous vein, lymphatic duct).
A further object of the invention is to provide a system which is minimally invasive, easy to implant and that requires no or minimal maintenance.
It is another object of the present invention to provide a system of small size, allowing laparoscopic implantation and no or minimal discomfort for the patient.
A further object of the present invention is to provide a drainage system that allows a tailored drainage action based on specific patient’s need.

These objects are achieved thanks to the drainage system and the method as defined in the claims.

Advantageously the input end of the system according to the invention may be introduced in an edematous space and the output end, e.g. the tip of a catheter, may be connected to a non-edematous space, (e.g., a functioning lymphatic vessel or a vein). The draining system is composed by at least two controllable micropumps connected in series, where the number of micropumps, as well as the distance between the micropumps, depends on the extension of the edema.
The idea is to emulate the functioning of the natural draining system: the lymphatic vessels. Multiple functional units, called lymphangions, compose such vessels. Each lymphangion is an actively contracting vessel tract, coupled to a non-return valve. Each lymphangion, when contracting, propels the lymph to the next unit and, when relaxing, drains the fluid from the previous unit and/or from the external space.
The lymphangions contract sequentially, allowing the drainage of the interstitial fluid from very large areas through peristalsis.

In an embodiment of the invention, the system comprises several functional units, in turn composed by a micropump and a porous element. Each functional unit is activated by a controller, and the sequential activation of the micropumps allows the drainage of fluid from the affected tissue through peristalsis, maintaining tissue homeostasis over time. This system may avoid the problems related to current therapies of chronic edemas and in particular of lymphedema, such as the need of recurring of classical approaches (e.g., physiotherapeutic sessions, massages, presso-therapies and the use of compression bandages), the aesthetical impairment for the patient and recurring limb infections due to the stagnant lymph.
In one embodiment, the invention provides for a permanently implantable fluid drainage system comprising \( N \) implantable pumps connected in series, with \( N \geq 2 \), each having an inlet, an outlet, and an actuator activated by an energy source through a controller; a first porous element connecting an edematous space with the inlet of the first pump; \( N-1 \) porous elements implanted in the edematous space, connecting the \( N \) pumps in series; and a last tube connecting the output of the last pump with the subcutaneous space of a distal output, wherein the porous elements are characterized by access pores distributed along their surface and wherein the \( N \) pumps are activated and coordinated through one or more controllers to provide the suction of the edematous liquid and its movement from the access pores of the porous elements to the output of the last tube.

The implantable drainage system in accordance with the present invention can be placed within the body of a patient at a suitable location depending on the particular treatment. For example, and without limitation, the implantable drainage system may be placed subcutaneously via surgery in an edematous area to be treated. Any suitable surgical technique can be used to implant the drainage system according to the invention, such as for instance laparoscopy. The implantable drainage system may be used to treat any condition involving the distributed accumulation of fluid in an interstitium within the body of a living host, such as for instance lymphedema, chronic venous insufficiency, chronic edema, myxedema, pulmonary edema, periorbital edema, lymphatic filariasis, edema which occur in specific organs as part of inflammations (e.g. in tendinitis or pancreatitis) and the like. In preferred embodiments, the conditions to be treated are lymphedema, chronic venous insufficiency or chronic edema. In a further preferred embodiment, the condition to be treated is lymphedema.

The implantable drainage system may comprise several porous elements having two tubular extremities. The porous elements have a size and a shape specifically adapted to the edematous space to be treated. Said porous elements may have a cylindrical shape, a polygonal shape, a flat shape or any other suitable shape. In one embodiment, the porous element has a tubular shape. The porous elements are characterized by access pores distributed along their surface, allowing the entry of the body fluid to be drained within the implantable drainage system. In a preferred embodiment, the body fluid to be drained may flow through the pores of the porous elements in a unidirectional way, i.e. from the interstitial space to be drained inside the porous element, while the opposite is impeded. In one embodiment, the extremity of the tubular portion of the first porous element which is not connected with the first pumping member (i.e., the distal input) is closed, so that no fluid can flow inside the first porous element from said extremity.

The system may comprise a last tube connecting the output of the last pumping member of the drainage system with the subcutaneous space of a distal output. The extremity of the last tube which is not in contact with the last pumping member of the drainage system (i.e., the distal output) provides for the release of the drained fluid in a distal area, which in one embodiment is a non-edematous subcutaneous area. In preferred embodiments, the distal output of the last tube of the drainage system ends in close proximity to the lymphatic system, or is connected with a lymphatic vessel or with a vein. In an alternative embodiment of the invention, the distal output of the last tube is connected to a tubular member of another implanted drainage system. In one embodiment, the last tube of the drainage
system is a catheter. The term "catheter" is used broadly to refer to any suitable tubing or structure that includes a lumen through which body fluids can flow.

The porous elements and the last tube may be constructed of a soft, flexible, biocompatible material. For example, the porous elements may be made of, without limitation, a polymeric material, such as polyethylene, polypropylene, polybutylene, polystyrene, polyurethane, polycarbonate and/or other suitable materials. In various embodiments, the porous elements and the last tube may include an anti-fouling and/or anti-fibrotic coating of the internal and external surfaces in order to avoid the clogging of the access ports and of the lumen of the tubes.

The pumping members are specifically designed to be easily implanted within the body of a living host and coupled to both the porous elements and/or the last distal tube through an inlet and an outlet. When activated, the pumping members are responsible for the suction of the fluid to be drained (e.g. an interstitial fluid) within the porous elements of the drainage system, thus allowing the drainage of the affected area and the transport of the fluid towards the distal output. The number and the position of the pumping members depend on the area to be treated, and can be therefore tailored based on the special needs of the living host.

In some embodiments of the invention, the pumping members are micropumps comprising a controller element which regulates the functioning or other parameters of the same. In the present disclosure, the term "controller" refers to any system which monitors and physically alters the operating conditions of a given dynamical system. In the frame of the invention, a controller generally controls the functioning of the micropumps by activating/deactivating the associated actuator and determines the fluid flow rate of said micropumps.

In one aspect of the invention, each and every pumping member has an integrated controller. In accordance with various embodiments of the invention, a controller periodically increases and/or decreases application of suction from the pumping members at porous elements. In various embodiments, the controller may control both the magnitude and associated time of the applied suction.

In various embodiments, the controller may receive information from one or more sensors. The one or more sensors may include, for example, a conventional pressure transducer that provides a signal representative of pressure within the porous elements and/or the last tube. Based on the signal received from the one or more sensors, the controller may verify and/or adjust the level of suction generated by the micropumps.

The controller may include, without limitation, a circuit, such as a timer circuit, and/or a Central Processor Unit (CPU) that may include memory and be appropriately preprogrammed or configured to be loaded with an appropriate program. In an alternative embodiment, each pumping member is controlled remotely by a controller implanted subcutaneously in a distal location. A wireless remote
control can non-invasively regulate any function of the drainage system. Even more important, several parameters of the drainage system may be programmable by such a remote control.

Each micropump may be driven by an actuator. In the frame of the present disclosure, an “actuator” is any type of motor that is responsible for moving or controlling a mechanism or system. It is operated by a source of energy, typically electric current, hydraulic fluid pressure, or pneumatic pressure, and converts that energy into motion. In at least some embodiment, micropumps of the drainage system can be driven for example by an electromagnetic actuator, by a piezoelectric actuator or by an electrosmotic actuator. The actuators can be provided with an energy source that is chargeable from outside the body.

In one embodiment of the invention, the implanted system may be passive, while the micropumps may be driven by magnetic forces from external actuators. In other embodiments, the energy source supplying the pumps’ actuators may comprise an internal energy source, i.e. the energetic source of the system may be incorporated in each micropump. In another variation of the system, the energy source may be an external source outside the body of the living host and the energy may be transmitted telemetrically via inductive coupling to the implanted active elements, such as for instance actuators having a coil. For example, in a particular embodiment the actuators are magnetically coupled to one or more actuators placed outside the body of the living host. Another variation of the system may include a mechanism to retrieve the energy for the implanted active elements from the living body movements. Alternatively, an energy transmitter transmitting energy wirelessly from an external energy source to charge an internal energy source can be envisaged.

The system preferably comprises feedback means for sending information from inside the host's body to the outside thereof to give feedback information related to at least one functional parameter of the system or a physical parameter of the host, thereby optimizing the performance of the system. Preferred functional parameters to be analyzed are the flow rate or the pressure of the drained fluid. One variation of the system may include a pressure and/or flow sensing element in a feedback loop to regulate the activation of the micropumps. In some embodiments of the invention, at least one pressure sensor and/or at least one flow velocity sensor is connected to at least one of the porous elements and/or the last tube, and the pressure and/or fluid flow rate values measured from the sensors determine the frequency of motion of the actuators.

For example, the sensors can measure pressure or flow rate of the body fluid through the porous elements and/or the last tubular element, and/or other desired measurements associated with body fluid drainage. Pressure sensors can be for instance small electrical sensors positioned along the drainage system.

The implantable drainage system in accordance with the present invention may be available in multiple configurations, depending on the position and of the extension of the edema. For lower limb edema, for example, the catheter connecting the edematous space to the non edematous space may be much longer respect to the case of upper limb lymphedema.
A further object of the present invention is to provide a method for draining excess body fluid from an edematous area to a non-edematous area by using the drainage system described in the present invention. In particular, it is disclosed herewith a method of draining excess fluid from an edematous space to the blood circulation, comprising implanting N pumps, with N>=2, connected in series through N-1 porous elements, an initial porous element and a last tube connecting the output of the last pump with the subcutaneous space of a distal output, and activating the pumps to drain fluid from the edematous space to the lumen of porous elements and to propel the said fluid to the output of the final tube.

In at least some embodiments of the said method, the pumps are activated in an intermittently, asynchronous manner, so that they create a peristaltic pressure wave along the porous elements and the last tube, and they can further be remotely activated. In one embodiment, the edematous space to be drained is a subcutaneous edematous space.

BRIEF DESCRIPTION OF THE DRAWINGS

The present invention will be described in details in the following section, containing the detailed embodiments of the invention, which are presented solely by non-restrictive examples and illustrated by the attached drawings in which:

FIG. 1A to 1C show detailed views of the invention according to an embodiment of the implantable system.

FIG 2A to 2B show detailed cut views of an embodiment of the fluid drainage system.

FIG 3 shows a first possible pumping scheme of the flow drainage system.

FIG 4A to 4B show a second possible pumping scheme of the flow drainage system.

FIG 5A TO 5H show a third possible pumping scheme of the flow drainage system.

FIG 6A TO 6C show detailed cut view of embodiments of the power sources and of the pumping units.

FIG 7A TO 7B show detailed views of possible configurations and placement of the fluid drainage system in the living host in case of lymphedema at the upper right limb.

FIG 8A TO 8B show detailed views of possible configurations and placements of the fluid drainage system in the living host in case of lymphedema at the lower right limb.

DETAILED DESCRIPTION OF THE INVENTION

The present disclosure may be understood more readily by reference to the following detailed description presented in connection with the accompanying drawing figures, which form a part of this disclosure. It is to be understood that this disclosure is not limited to the specific conditions or
parameters described and/or shown herein, and that the terminology used herein is for the purpose of describing particular embodiments by way of example only and is not intended to be limiting of the claimed disclosure.

In one aspect of the invention, the implantable fluid drainage system for distributed edemas is designed to be implanted subcutaneously and to drain fluid accumulated in an interstitium of a living host, actively and continuously transporting it to a distal, non-edematous space. FIG. 1A shows a transverse view of the fluid drainage system according to one embodiment of the invention. The system may comprise N pumping members 5, 6, 7 and N + 1 porous elements, in form of tubular members, 1, 2, 3, 4. The first N tubular members 1, 2, 3 have multiple accesses pores 8 to their lumen on their lateral surfaces, while the last tubular member 4 has accesses only on its extremities.

The first tubular member 1 may have one extremity 9 without access, while its other extremity may be connected to the inlet of the first pumping member 5. The last tubular member 4 may have one extremity 10 connected to the last pumping member 7 and the other extremity 11 connected to a non-edematous space. For example, as shown in FIG. 1B, the tubing member 4 could be sutured to a functional vessel 12, which in one embodiment could be a lymphatic vessel or a vein, through the use of a suturable connector 13. In another embodiment of the invention, depicted in FIG. 1C, the tubing member 4 may be anchored to the internal side of the skin 14 in a distal area, where functional lymphatic vessels 15 are present.

As shown in FIG. 1A, the tubular members 2, 3, may have one extremity 16, 17 connected to the outlet of the foregoing pumping member 5, 6, and the other extremity 18, 19 connected to the inlet of the following pumping member 6, 7.

Both the tubular and the pumping members 1, 2, 3, 4, 5, 6, 7 may be coated with an antifibrotic and/or antifouling coating.

The activation of each pumping member may be controlled by a controller unit, as shown in FIG. 2. In a first embodiment, depicted in FIG. 2A, a controller unit 20 may be connected telemetrically with each pumping unit 5, 6, 7 and placed in a distal location, inside or outside the living host. The controller unit 20 will supply the input signals to the pumping units 5, 6, 7 via an antenna 21 and received by dedicated antennas 22, 23, 24 on the pumping units.

At the same time, the pumping units 5, 6, 7 may supply feedback signals to the controller unit 20 via the antennas 22, 23, 24. In an alternative embodiment, shown in FIG. 2B, the controller unit 20 is coupled to the pumping unit 5, 6, 7 via a wire 25.

In one aspect of the invention, the controller unit 20 may as well receive inputs by one or more sensors 26, 27, which provide information about the pressure or flow rate in the tubing members. As shown in Figure 2B, the sensors 26, 27 can be positioned proximate to the outlet or the inlet of the pumping members. In one embodiment, the first sensor 26 can measure the flow rate and/or the pressure within a porous element before it enters the pumping member 5, and the second sensor 27 can measure the flow rate and/or pressure within the proximal portion of the last tube as it exits the pumping member 7. This information can be used to ensure the pumping members generate the desired drainage rate, to
monitor patient parameters and/or derive other desired measurements or characteristics. In other embodiments, the drainage system can include more or less sensors. In one embodiment, the sensors 26, 27 may provide said information via the antenna of one or more pumping unit. In an alternative embodiment, the sensor 26, 27 may send the inputs via the wire 25.

The controller 20 may activate and coordinate the pumping members 5, 6, 7, according to different pumping schemes, depending on the desired draining action. FIG 3 shows a possible pumping scheme in a case in which the number N of pumping members is 4. In a first embodiment, the pumping members 5, 6, 28, 7 are active at the same time, but the provided flow rates V1, V2, V3, V4, are different between each other. In particular, the following relation may be valid: V1<V2<V3<V4. In this way, pumping member 5 drains fluids from the interstitial space 29 through the accesses of the tubing member 1 and move the fluids towards the tubing member 2; pumping member 6 drains from the interstitial space 30 and from the tubing member 2, avoiding the leakage of the fluids pumped by 5 from the accesses of 2, and move the drained fluids towards the tubing member 31; pumping member 28 drains from the interstitial space 32 and from the tubing member 31, avoiding the leakage of the fluids pumped by 6 from the accesses of 31; In the same way, pumping member 7 will drain from the interstitium 33 and from the tubing member 3, moving the fluids towards the tubing member 4, which is without lateral accesses as shown in FIG 1A.

In case the number of pumping member N is equal to 4, another non-exclusive example of pumping scheme is shown in FIG 4. The pumping scheme may be composed of a two step cycle. The first step is schematized in FIG 4A. Initially, 5 is draining fluid from the interstitial space 29 towards 2; at the same time, 28 is draining fluids from the interstitial space 32 towards 3; meanwhile, 6 and 7 act as an open circuit, allowing the passage of the fluid pumped by 5 and 28. The second step of the cycle is shown in FIG 4B; pump 5 closes; pumping member 6 drains fluid from 30 and from 2, moving it towards 31; at the same time, 7 drains from 3 and from 33, transporting the fluid in the tubing member 4, while the pumping member 28 is open.

Another non-exclusive example of pumping scheme, in case of N equal to 4, is depicted in FIG 5. In this case, the pumping scheme will be composed of 7 consecutive steps, shown in FIG 5A-G. In step 1, shown in Fig. 5A, all the pumping members except 7 will be closed, while pumping element 7 will drain fluid from interstitial space 33 through tubular element 3, transporting it towards the last distal tube 4. In the step 2, shown in FIG 5B, the pumping member 28 is activated, draining from the interstitial space 32 through tubular element 31 and transporting fluids towards porous tube 3. In step 3, depicted in FIG 5C, the pumping member 7 is deactivated, remaining open and allowing the passage of fluid due to the pumping action of the pumping element 28. Step 4, schematized in FIG 5D, contemplate the activation of the pumping member 6, draining from interstitial space 30 through tubular element 2 and pushing fluid towards the porous tube 31. As before, the step 5 (FIG 5E) consists in the deactivation of 28, which remains open. In step 6 (FIG 5F), pumping member 5 is activated, draining from the interstitium 29 through the tubing member 1, towards tubing member 2. In the last step, shown in FIG 5G, pumping member 6 deactivates, remaining open. Next, the cycle continues in the reverse order, with step 6 (FIG 5H) -5-4-3-2-1. Overall, the described pumping scheme provides suction of fluid from the multiple
opening in the tubing member of the draining system, transporting it towards the final tubing member by the formation of a peristaltic wave.

The implantable fluid drainage system can be activated by magnetic forces from an external actuator, or by inductive coupling with an external power source, or by one or more implanted batteries, as shown in FIG 6. In a first embodiment, shown in FIG 6A, one or more power sources 34 are outside the skin 35 of the living host. Power source 34 activates an electromagnetic actuator 36, which is outside the body of the living host, although in close proximity with the skin 35 and with the implanted pumping member 37. Pumping member 37 may include a permanent magnet 38. When the electromagnetic actuator 36 is activated and put in movement, it exerts magnetic forces on 38, which moves as well and activates the pumping member 37.

In a second embodiment, shown in FIG 6B, the power may be transmitted from one external power source 34 to the implanted pumping members 37 via resonant magnetic coupling; a primary magnetic coil 39 may be placed outside 35 and powered by 34. A secondary magnetic coil 40 may be part of the pumping member 37. When activated, magnetic coil 39 transfers energy to 40, which in turn activates the actuator 41. 41 may for instance be an electromagnetic actuator, or a piezoelectric actuator or an electrosmotic actuator.

In FIG 6C is shown a third, non-exclusive embodiment of the invention, in which the energy needed to activate the implanted pumping member 37 may be provided by an implanted battery 42. Battery 42 may be directly connected via wire with the actuator 41.

The implantable fluid drainage system schematized in FIG 1 may be implanted subcutaneously in areas where there is a distributed accumulation of fluids. Non-exclusive example may be lymphedema-affected upper limbs, lower limbs or male genitalia.

In FIG 7 two possible placements of the implantable fluid drainage systems in case of upper right limb lymphedema are shown. FIG 7A illustrates the placement of a single implant 43 in the subcutaneous space 48 of the right upper limb of the living host. In this example, the implant is composed by 4 pumping members 44 and by 5 tubular members 45. In this example, the final output 11 of the drainage system is sutured to a functional vessel 46 (lymphatic or venous) in the lower neck area.

FIG 7B schematizes a second possible configuration, in which two drainage systems 43 and 47 are implanted in the edematous area 48. In this example, the additional drainage system 47 is composed by 3 pumping members and 4 tubular members. In this example, the outlet 49 of 47 is connected to the last tubular member of 43.

FIG 8 illustrates two possible placements of the implantable fluid drainage system in case of lower limb lymphedema. FIG 8A illustrates the placement of a single implant 50 in the edematous area 57. In this example, the implant is composed by 5 pumping members and 6 tubular members 45. In this example, the final output 11 of the system is sutured to the great saphenous vein 51.

FIG 8B schematize a second possible configuration, in which three drainage systems 52, 53, 54 are implanted in the edematous area 57. In this example, drainage systems 52 and 54 are composed by 3 pumping members and 4 tubular members; drainage system 53 is composed by 4 pumping members and 5 tubular members. In this example, the implant 54 is directly connected to the vein 51 through its
output 11; the output 55 of drainage system 53 is connected to 54; the output 56 of drainage system 52 is connected to drainage system 53.
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OTHER PUBLICATIONS


CLAIMS

1. A medical fluid drainage system comprising:
   N pumps, with N>=2, connected in series, each having an inlet and an outlet;
   a first inlet member connected to the inlet of the first pump;
   N-1 inlet members providing fluidic connections between the N pumps;
   an outlet member connected to the outlet of the last pump; said pumps, inlet members and
   outlet member forming a fluid line;
   wherein each inlet member contains at least one entry point through which a fluid may directly
   enter into said fluid line and wherein said outlet member contains at least one exit point through
   which said fluid may directly exit said fluid line.

2. The system of claim 1 wherein each inlet member has a tubular shape with at least one lateral
   entry point.

3. The system of claim 1 wherein said inlet members comprise a porous surface containing several
   entry points.

4. The system of claim 1 wherein said inlet members comprise an absorbing surface containing
   several entry points.

5. The system of anyone of the previous claims wherein said outlet member has a tubular shape
   with an open free end.

6. The system of anyone of the previous claims wherein said outlet member has a tubular shape
   with at least one lateral exit point.

7. The system of anyone of the previous claims comprising a controller intended to provide a
   coordinated operation of said pumps.

8. The system of anyone of the previous claims for draining a physiological fluid from a location
   within a living body.

9. The system of claim 8 for the treatment of edema.

10. The system of claim 8 or 9 characterized by the fact that it is partially or totally implantable.

11. The system of anyone of the previous claims wherein said outlet member is connected in parallel
    to (an)other fluid drainage system(s).

12. The system of anyone of the previous claims comprising one pressure or flow sensor connected
    to at least one inlet member or to the outlet member, and wherein the pressure or fluid flow
    rate values measured from the sensors determine the frequency of motion of the actuators.

13. A method of draining excess fluid from an edematous space to another location, comprising:
    implanting the system as defined in anyone of the previous claims in a living host, positioning
    said first inlet member in an edematous area and locating the end of the outlet member in a
    non-edematous area;
    activating and coordinating the pumps to drain fluid from the edematous area to the non-
    edematous through said fluid line.

14. The method of claim 13, wherein each pump provides a specific flow rate which depends on its
    location along the fluid line.
15. The method of claim 13 or 14, wherein the pumps function in a way as to generate a peristaltic pressure wave along the fluid line.
**INTERNATIONAL SEARCH REPORT**

**International application No**
PCT/IB2015/057394

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### A. CLASSIFICATION OF SUBJECT MATTER

**INV.** A61M27/00

**ADD.**

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

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### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

A61M

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### E. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tbody>
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**X** Further documents are listed in the continuation of Box C. **X** See patent family annex.

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* 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 application or patent but published on or after the international filing date
- **L** document which may throw doubts on priority claim(s) on 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

- **T** later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

- **X** document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

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**Date of the actual completion of the international search**

11 January 2016

**Date of mailing of the international search report**

19/01/2016

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**Name and mailing address of the ISA/Authorize officer**

European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel: (+31-70) 340-2040,
Fax: (+31-70) 340-3016

Roland, Philippe
<table>
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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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| A        | GB 2 350 794 A (HABIB NAGY ADLY [GB])  
13 December 2000 (2000-12-13)  
page 1, line 24 - page 4, line 30; figures 1-3 | 1-12                  |
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. [ ] Claims Nos.: 13-15 because they relate to subject matter 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 and therapy

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 fee, 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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