A device suitable for use in a body lumen and comprises an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter (i.e., thrombus) from a wall of the body lumen (i.e., a vein or an artery). The radially expansible member has a proximal end and a distal end and is adjustable between a contracted orientation and an expanded orientation. The elongated control member comprising a proximal arm connected at or adjacent to the proximal end of the radially expansible member and a distal arm connected at or adjacent to the distal end of the radially expansible member such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member. The device comprises a control mechanism operatively connected to both arms and adapted to provide resistance to the movement of one arm relative to the other. The control mechanism may be a biasing means, a brake means, or both.

**Figure 10.**

**Abstract:**
A device suitable for use in a body lumen and comprises an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter (i.e., thrombus) from a wall of the body lumen (i.e., a vein or an artery). The radially expansible member has a proximal end and a distal end and is adjustable between a contracted orientation and an expanded orientation. The elongated control member comprising a proximal arm connected at or adjacent to the proximal end of the radially expansible member and a distal arm connected at or adjacent to the distal end of the radially expansible member such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member. The device comprises a control mechanism operatively connected to both arms and adapted to provide resistance to the movement of one arm relative to the other. The control mechanism may be a biasing means, a brake means, or both.

**Designated States (unless otherwise indicated, for every kind of national protection available):**
Title
A device suitable for removing matter from inside the lumen and the wall of a body lumen

Technical Field
The invention relates to a device suitable for removing matter from inside the lumen and the wall of a body lumen. In particular, the invention relates to a thrombectomy device for removing thrombus from the walls of a vein or artery of a human.

Background to the Invention
There have been many methods and devices to extract a blockage from vessels. Thrombectomy devices exist that macerate and extract the clot rather than pulling out the clot intact. The followings are some of the related patents.

US patent number 5972019 introduces an embolism treatment device having a core element and a cage. The cage may include an expandable braid which is rotatably attached to the core element and can be opened to separate the clot from the vessel wall or be expanded beyond the clot to pull it out. The core element has a rotatable part which is non removable from the cage.

Patent 6066158 discloses an embolectomy device having a core wire element and a spiral collector to collect the embolism. The device may also have an actuator to allow expansion of the cage after delivery.

US patent 5795322 presents a device for reducing the size of thrombus and removing it from the vessel. The device is a tube-like catheter with a filter formed from longitudinal strip-shape of the catheter. The filter opens when the distal and the proximal sided are pushed towards each other respectively. The device comprises a jet flow and lumen to extract the clot.

US patent 6660014 presents a catheter for removing occlusive material from vessel lumen. The catheter comprises a radially expandable positioning cage and a radially expandable macerator within the cage. The diameter of the cage is adjustable with predetermined unconstrained diameter. Patent number 6454775 discloses an expansible macerator.
Patent application US 2006/0229645 presents a radially expansible cage for the removal of hardened and aged thrombus from the vessel wall. The cage opening and closing is controlled at the proximal/user end by moving the cage ends closer together or further apart; this is done either manually, or with a threaded tube to define the radial expansion of the cage.

US patent 6383205 presents a mechanism including a double filter device to extract the clot with minimum risk of embolism.

Patent number 6616679 is a vascular device for emboli and thrombus removal. It includes a blood permeable sac which collects the emboli and can be collapsed. This is a fast way to extract small emboli but for large clot the sac need to be extremely long.

In patent number 6652548 a thrombectomy device has been claimed. The catheter based device comprises shearing members located at the distal ends of the catheter and the inner shearing member is rotatable.

Patent numbers 6656203, 7083633, 7014647, 6558405, EP1310219A3, EP1310219B1, 20070173883, US6179859, US7922741 and US20110125181 are examples of different structures for embolic filters. The filter may be placed in the vessel prior to an operation such as stenting or PTCA. After the operation, the filter is collapsed and taken out of the body. Neither of the filters are suitable for large tapered veins with thrombosis. Because filters are not designed to be moved along vessels after deployment, this makes it difficult to remove long thrombus. The existing devices and methods described above do not have a high performance in harvesting large volumes of thrombus and also thrombus in large and/or tapered and/or branched vessels. Limitations associated with these devices include procedural duration and thrombus removal efficiency.

It is an object of the invention to overcome at least one of the above-referenced problems.

**Statements of Invention**

Broadly, the invention provides a device suitable for use in a body lumen comprising an elongated control member and a radially expansible member (i.e. a cage, funnel, or a ring) disposed at or adjacent to a distal end of the elongated control arm and that is adjustable.
between a contracted orientation and an expanded (deployed) orientation. The radially expansible member is adapted to remove matter (i.e. thrombus) from the walls of a body lumen (i.e. veins, arteries or other lumens such as the urethra), for example scrape thrombus from vessel walls, collect matter removed from the walls of a lumen, and optionally both, when in a deployed orientation. The control member comprises two arms, one of which is connected to or adjacent to a proximal end of the radially expansible member and the other of which is connected to or adjacent to a distal end of the radially expansible member. Movement of one of the arms relative to the other effects adjustment of the diameter or radial strength of the radially expansible member, for example adjustment of the diameter from a contracted orientation to an expanded (deployed) orientation, or adjustment of the radial force from a first force to a greater force. The movement is preferably longitudinal movement. The device additionally includes a control mechanism that is ideally operatively connected to both arms and provides resistance to movement of one of the arms relative to the other. The control mechanism may comprise biasing means for biasing the radially expansible member into, or in the direction of, an expanded or contracted orientation (Fig. 11), it may include brake means which clamp the two arms in one orientation with a certain force such that movement of one of the arms relative to the other only occurs when a specific predetermined force is applied to the radially expansible member (Fig. 9), or it may comprise a combination of biasing means and brake means (Fig. 12).

A device of the invention is ideally suited for use in tapering lumens with obstructions such as valves where movement of the cage in a deployed shape along the lumen requires the diameter of the cage to change. When the lumen is tapering inwardly (in the direction of travel of the cage), the control mechanism exerts a force on the radially expansible member resisting contraction of the radially expansible member (by biasing the radially expansible member into an expanded orientation or by clamping the two arms in a specific disposition) until the force exerted on the radially expansible member by the lumen exceeds the total resistance force including that exerted by the resistance mechanism at which point the diameter of the radially expansible member will reduce. This means that the radially expansible member can be moved along the walls of the lumen exerting radial force against the walls, thereby effecting a scraping / collecting action.

Likewise, when the tube is tapering outwardly (in the direction of travel of the cage), the resistance mechanism comprises biasing means that exerts a force on the cage by biasing the
cage into an expanded orientation. This means that cage will scrape along the walls of the vasculature exerting radial force against the walls, thereby effecting a scraping action along the outwardly tapering walls.

According to the invention, there is provided a device suitable for use in a body lumen and comprising:

- an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,
- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,
- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter and/or radial strength of the radially expansible member,

the device being characterised in that it comprises a control mechanism operatively connected to both arms and adapted to provide resistance to the movement of one arm relative to the other.

The distal arm is generally connected to the radially expansible member distally of the proximal arm connection. Thus, the distal arm may be connected at or adjacent to the distal end of the radially expansible member, and the proximal arm may be connected to the radially expansible member at a point between the distal arm connection and the proximal end of the cage.

Preferably, the radially expansible member is a cage, although other radially expansible members such as, for example, expansible rings, funnels are envisaged. The radially expansible member may be formed from one, two, or more struts, which may be arranged in a helical arrangement to form the cage.

The distal or proximal arm may be, for example, a wire or a tube. In one embodiment, one of the arms is a tube (for example the proximal arm) and the other arm (for example the distal arm) is a wire, in which the wire is suitably disposed within a lumen in the tube. Suitably, the distal and proximal arms are co-extensive along most of their length (for example they are coextensive up to the cage). The control member comprising the two arms may be disposed
within a tube, typically a catheter tube. In a preferred embodiment, the proximal arm is tubular and the distal arm is disposed within the proximal arm. Preferably, the distal arm is tubular (so as to facilitate the device being delivered over a guidewire).

Typically, the control mechanism comprises biasing means adapted to bias the radially expansible member into, or in the direction of, the expanded or contacted orientation. Ideally, the control mechanism comprises biasing means adapted to bias the cage into, or in the direction of, the expanded orientation.

Suitably, the control mechanism comprises brake means adapted to resist movement of one arm relative to the other arm. One embodiment of a brake means is a friction screw, typically an adjustable friction screw. Preferably, the control mechanism is adjustable so that the level of resistance to movement can be adjusted.

In one embodiment, the control mechanism comprises biasing means adapted to bias the radially expansible member into, or in the direction of, the expanded or contacted orientation and brake means adapted to resist movement of one arm relative to the other arm, for example brake means adapted to prevent expansion or retraction of the radially expansible member.

Suitably, the control mechanism comprises a first part connected to one of the arms, a second part connected to the other of the arms and movable relative to the first part, and force controlled resistance means for resisting movement of the first part relative to the second part. The force controlled resistance means may be biasing means adapted to bias the cage into, or in the direction of, the expanded or contacted orientation, and/or brake means adapted to resist movement of the radially expansible member or movement or one arm relative to the other arm, typically both.

In this specification, the term "force controlled" as applied to the resistance mechanism should be understood to mean that the diameter of the radially expansible member is not pre-determined or controlled, but is dependent on the force applied.

In one preferred embodiment, the invention provides a device suitable for use in a body lumen and comprising:
an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,

- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,

- the elongated control member comprising a proximal arm connected to the radially expansible member and a tubular distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter and/or radial strength of the radially expansible member,

- a control mechanism operatively connected to both arms and adapted to provide resistance to the movement of one arm relative to the other, in which the control mechanism comprises a first part connected to one of the arms, a second part connected to the other of the arms that is movable relative to the first part, and force controlled resistance means for resisting movement of the first part relative to the second part, characterised in that the force controlled resistance means comprises (a) biasing means for biasing the radially expansible member into the expanded orientation, or (b) a brake means adapted to resist movement of one of the first and second parts relative to the other of the first and second parts.

In another embodiment, the invention provides a device suitable for use in a body lumen and comprising:

- an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,

- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,

- a thrombus extractor, at least a part of which is preferably disposed within the radially expansible member

- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter and/or radial strength of the radially expansible member,
- a control mechanism operatively connected to both arms and adapted to provide resistance to the movement of one arm relative to the other, in which the control mechanism comprises a first part connected to one of the arms, a second part connected to the other of the arms that is movable relative to the fixed first part, and force controlled resistance means for resisting movement of the first part relative to the second part, characterised in that the force controlled resistance means comprises (a) biasing means for biasing the radially expansible member into the expanded orientation, or (b) a brake means adapted to resist movement of one of the first and second parts relative to the other of the first and second parts.

Typically, the force controlled resistance means comprises biasing means, for example a resiliently deformable or displacable member such as for example a spring or a pneumatic or hydraulic member, generally disposed between the first and second parts of the control mechanism. The spring may be any type of spring, for example a compression, tension, flat, constant force, or adjustable constant spring.

In one embodiment, the force controlled resistance means comprises biasing means, for example a resiliently deformable member disposed between the first and second parts of the resistance mechanism, and a brake means adapted or configured to resist movement of one of the first and second parts relative to the other of the first and second parts.

The control mechanism may be disposed at any point along the device, for example proximal to the radially expansible member, distal to the radially expansible member, or adjacent to or within the radially expansible member. In a preferred embodiment, the control mechanism is disposed at a proximal end of the catheter (for example, on the handle) such that, in use, it is exposed proud of the body.

In a preferred embodiment of the invention, the first part of the control mechanism is connected to the proximal end of the radially expansible member by means of the proximal arm, and the adjustable second part of the control mechanism is attached to the distal end of the radially expansible member by means of the distal arm. Thus, the distal arm typically passes through the cage.
Preferably, the proximal arm is a tube comprising a lumen and the distal arm is disposed within the lumen of the first arm, typically coaxially with the first arm. Preferably, the distal arm is a tube, ideally adapted for receipt of a guidewire.

Movement of one arm relative to the other arm effects adjustment of the diameter and/or radial strength of the radially expansible member. Preferably, the movement is longitudinal movement, although other movement is envisaged, for example lateral, radial, circumferential rotational or combinations thereof.

In one embodiment, the brake means comprises a friction screw fixed to one of the parts of the control means and adjustable to engage the other part of the control means (or the arm that is operably connected to the other part). Preferably, the friction screw is fixed to the second part, for example a movable stop forming part of the second part, and is adjustable to engage the first part or the control means or the distal arm that is operable connected to the first part.

In one embodiment of the control mechanism, the first part comprises a barrel, and the second part is adapted for sliding movement within the barrel, wherein the resistance means comprises a friction screw, preferably an adjustable friction screw, disposed on the barrel and adapted for engagement with the second part. Suitably, the resistance means additionally comprises biasing means suitably adapted to bias the cage into, or in the direction of, the expanded orientation.

Typically, the control mechanism is disposed proximally of the radially expansible member such that in use it is located outside of the body. Suitably, the first part is operably connected to the proximal arm and comprises a guide path, and the second part is operably connected to the distal arm and is associated with the first part for movement along the guide path, and wherein the force controlled resistance means comprises a resiliently deformable member disposed along the guide path between the first and second parts. Preferably, the first part comprises a barrel and a second part comprises a block configured for sliding and controlled movement within the barrel.

Typically, the first part comprises a movable stop which is movable to vary the length of the guide path and/or the resiliently deformable member.
Suitably, the brake means comprises a friction screw operably connected to the movable stop and configured for adjustable engagement with the distal arm.

Suitably, the control mechanism is disposed within the radially expansible member and comprises a first part connected to the distal arm, a second part connected to the proximal arm, and a helical spring operatively connected to the first and second parts and configured to provide force controlled resistance to movement of the first and second parts together.

Preferably, the radially expansible member comprises a cage. Suitably, the cage comprises a proximal portion having apertures for receipt of thrombus into the cage, and a distal portion having a fine mesh for capturing thrombus, in which the cage optionally comprises a cut tube. Alternatively, the cage comprises a distal portion having apertures for receipt of thrombus into the cage, and a proximal portion having a fine mesh for capturing thrombus, in which the cage optionally comprises a cut tube.

Suitably, the control mechanism is disposed distally of the radially expansible member. Suitably, the control mechanism comprises a resiliently deformable member (i.e. a spring) having a distal end (first part) operably connected to the distal arm and a proximal end (second part) operably connected to a distal end of the radially expansible member, and wherein the distal end of the distal arm is operably connected to the distal end of the radially expansible member by means of the resiliently deformable member.

Preferably, the device comprises an extractor at least a part of which is disposed within the radially expansible member. Suitably, the extractor comprises holes or apertures dimensioned to allow blood, but prevent thrombus, pass out of the extractor.

Suitably, the extractor comprises a helical formation adapted to rotate. The rotation is adapted to remove thrombus from the radially expansible member or optionally deliver agents into the body lumen, for example a thrombolytic agent. The helical formation may comprises a single, double or triple helical formation. Typically, the pitch of the helix on the helical formation is configured to squeeze blood from thrombus during use.
Typically, the helical formation is disposed within an extractor tube. Ideally, a portion of the helical formation within the radially expansible member is exposed by means of, for example, one or more windows or cut-away portions.

Suitably, a leading edge of the extractor tube comprises a cutting edge. Preferably, the helical formation comprises a cutting edge, ideally disposed at or adjacent a distal end of the helical formation. In one embodiment, a leading edge of the extractor tube comprises a cutting edge and the extractor comprises an aspirator tube.

In one embodiment, the thrombus extractor is an aspirator tube.

Typically, the force controlled resistance means is self-adjusting. In this specification, the term "self adjusting" as applied to the force controlled resistance means should be understood to mean that the resistance means adjusts itself without any user input other than the actions of the user to move the device along a vessel.

Typically, the elongated control member is contained within a tubular sheath. Ideally the tubular sheath extends along all or most of the length of the control member. Suitably, the longitudinal position of the sheath is adjustable. Suitably, the tubular sheath can be adjusted to cover the radially expansible member and maintain the radially expansible member in a contracted orientation.

Typically, the tubular sheath comprises a plurality of holes for perfusion of a liquid.

Suitably, the device comprises a liquid administration lumen configured to deliver a liquid into the body. Preferably, the liquid administration lumen comprises a lumen formed within the distal arm, between the distal arm and proximal arm, and between the proximal arm and an external sheath. Ideally, the device comprises an injection port for delivery of liquid into the liquid administration lumen.

In a preferred embodiment, the invention provides a device suitable for use in a body lumen and comprising:

- an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,
- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,
- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member,

the device being characterised in that it comprises biasing means for biasing the radially expansible member into, or in the direction of, an expanded orientation.

In another embodiment, the invention provides a device suitable for use in a body lumen and comprising:
- an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,
- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,
- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member,

the device being characterised in that it comprises biasing means for biasing the radially expansible member into, or in the direction of, an expanded orientation and brake means brake means adapted to resist movement of one arm relative to the other arm.

In another embodiment, the invention provides a device suitable for use in a body lumen and comprising:
- an elongated control member having a distal and a proximal end, and a radially expansible member disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,
- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,
- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member, the device being characterised in that it comprises a resistance mechanism comprising a fixed first part connected to one of the arms, a second part connected to the other of the arms and movable relative to the fixed first part, and resistance means for resisting movement of the first part relative to the second part, in which the resistance means optionally comprises biasing means adapted to bias the cage into the expanded or contacted orientation or brake means adapted to prevent expansion of the cage.

In one embodiment, the radially expansible member is a cage. Typically, the cage is a filtering cage (i.e. it is adapted to collect or filter thrombus from blood that passes through the cage). In another embodiment, the cage is a shearing cage (i.e. a cage adapted to shear or scrape thrombus from a wall of the vasculature). In a preferred embodiment, the cage is both a shearing and filtering cage. Suitably, the cage has a proximal end that is at least partially open (for receipt of thrombus). The cage may comprise a braided material, for example a braided wire, or it may comprise a cut tube, for example a cut tube (i.e. Polymeric, metal such as stainless steel Nitinol or cobalt chromium, or ceramic; or a combination of these materials) or a laser cut tube, or it may comprise a braided material and a cut tube. When formed from a braided material, the density of the braid may be greater towards the distal end of the cage that the proximal end of the cage. In a preferred embodiment, the cage is formed from a shape memory material such as Nitinol. Preferably, the cage has an at least partially open proximal end and tapers inwardly towards its distal end.

In one embodiment, the cage comprises a distal section having a fine mesh, and an intermediate section having large apertures. Thus, the fine mesh is suited for filtering/capturing thrombus, and the large apertures adapted for allowing passage into the cage of thrombus. Typically, the cage is cut from a tube, suitably a polymeric or metal tube.

In one embodiment, the cage comprises a cutting element disposed circumferentially around the cage. This may be a wire, or a cutting edge. Typically, at least a part of the cutting element is exposed proud of the cage. Suitably, at least a part of the cutting element is disposed within
the cage. Typically, the cutting element is disposed with respect to the cage such that the cutting element expands and contracts with the expansion and contraction of the cage, respectively.

In one embodiment, the invention provides a device suitable for use in a body lumen and comprising:

- an elongated control member having a distal and a proximal end, and a cage disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,

- the radially expansible member having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,

- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member

the device being characterised in that the cage comprises a distal section having a fine mesh for capturing thrombus, optionally a proximal section having a coarse mesh adapted for crimping attachment to the proximal arm, and an intermediate section having apertures dimensioned to allow passage of thrombus into the cage.

In another embodiment, the invention provides a device suitable for use in a body lumen and comprising:

- an elongated control member having a distal and a proximal end, and a cage disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,

- the cage having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,

- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member,
the device being characterised in that the cage has an at least partially open proximal end and tapers inwardly towards its distal end, and wherein a helical extractor is optionally disposed within the cage towards a distal end of the cage.

In one embodiment, the invention provides a device suitable for use in a body lumen and comprising:

- an elongated control member having a distal and a proximal end, and a cage disposed at or near the distal end and adapted for collection and/or shearing of matter from a wall of the body lumen,
- the cage having a proximal end and a distal end and being adjustable between a contracted orientation and an expanded orientation,
- the elongated control member comprising a proximal arm connected to the radially expansible member and a distal arm connected to the radially expansible member distally of the proximal arm connection such that movement of one arm relative to the other arm effects adjustment of the diameter or radial strength of the radially expansible member,

the device being characterised in that the cage comprises a cutting or separation element disposed circumferentially around the cage.

Preferably, the device of the invention comprises an extractor for matter (i.e. debris such as thrombus) at least a part of which is preferably disposed within the radially expansible member. The extractor may comprise either aspiration, a helical formation adapted to rotate and remove thrombus from the radially expansible member, or a combination of both. The helical formation may comprise a screw or auger, that is preferably arranged co-axially about the second (distal) arm, but optionally may be arranged eccentrically to the radially expansible member axis. Alternatively, the helical formation may comprise a helical wire or the like arranged, for example, around the distal arm and adapted to rotate. Preferably, the helical formation is disposed within an extractor tube, wherein a distal end of the helical formation that is disposed within the cage is exposed proud of the extractor tube (i.e. the tube may be cut-away leaving a part of the helical formation within the radially expansible member exposed. In one embodiment of the invention, the extractor comprises a wire arranged helically about the distal arm for rotation about the arm, and an extractor tube, wherein the helical wire and distal arm are disposed within the extractor tube with a portion of the distal end of the helical wire exposed proud of the extractor tube. Suitably, the extractor tube comprises a window disposed within
the cage, typically disposed towards the distal end of the cage. Preferably, a leading edge of the extractor tube comprises a sharp or cutting edge.

In the device of the invention, the proximal arm is generally connected at or adjacent to the proximal end of the radially expansible member. Typically, the distal arm is generally connected at or adjacent to the distal end of the radially expansible member.

Preferably, the device is a thrombectomy device, ideally a thrombectomy catheter.

The invention also relates to a method for removing matter from a target area of a wall of a body lumen, for example thrombus from a wall of a vein or artery, comprising a step of providing a device according to the invention with the radially expansible member is a contracted orientation, inserting the device into the target lumen such that the radially expansible member is positioned distally of the target area of the wall, adjusting the radially expansible member from a contracted orientation to an expanded (deployed) orientation, and withdrawing the radially expansible member along the lumen such that the radially expansible member scrapes some / all matter from the target area of the wall of the lumen.

The invention also provides a method of removing thrombus from a blood vessel comprising the steps of placing the device of the invention in a blood vessel, and moving the device along the blood vessel. Typically, the device is moved along the blood vessel in an inwardly tapering direction or an outwardly tapering direction. Suitably, the blood vessel is a vein, typically a large vein.

Brief Description of the Figures
The invention will be more clearly understood from the following description of some embodiments thereof, given by way of example only, in which:

Figure 1. Shows a braided cage in its expanded configuration, with a filtered end (201) and an open end (202).

Figure 2. An example of a cage made from laser cut Nitinol. The distal side is solid tube and the proximal side is a closed mesh to fit over the proximal arm tube.
Figure 3. An example of a laser cut cage after expansion. The circumferential elements act as filter and separate the thrombus from the vessel wall.

Figure 4. Cage with cutting element external to the cage.

Figure 5. Cage with cutting element internal to the cage.

Figure 6. Cage with cutting element as part of the cage.

Figure 7. The perimeter of the cage may be tapered at an angle.

Figure 8. Schematic of the cage and control mechanism, operated using a friction screw.

Figure 9. Schematic of the cage and control mechanism, operated using a friction screw.

Figure 10. Schematic of the cage and control mechanism, operated using a friction screw and a spring.

Figures 11A and 11B. Schematic of the cage and control mechanism, operated using a spring, and a movable stop.

Figure 12. Schematic of the cage and control mechanism, operated using a flat spring.

Figure 13. Schematic of the cage and control mechanism, operated using a flat spring.

Figure 14. Schematic of the control mechanism, operated using a flat spring.

Figure 15. The extractor is made of a wire wrapped around a tube. It transports and macerates the thrombus.

Figure 16. The wire may be displaced above the tube surface.
Figure 17. The proximal arm may act as the outer tube of the extraction mechanism and provide a passage way to let material from the blood vessel into the lumen of the macerator.

Figure 18. The proximal arm of extractor with cutting edge and larger extraction path.

Figure 19. The proximal arm of extractor may have more than one leading edge.

Figure 20. Image of the macerator with a rotating tip/cutting edge. The tip rotates with the helical formation.

Figure 21. Shows the cage in its open configuration. The passageway provided by the proximal arm, may be located inside the cage.

Figure 22. The extractor may be eccentric to the centreline of the cage.

Figure 23. The tip of the extractor may be eccentric and positioned away from the centreline of the cage.

Figure 24. Section view - the extractor may contain a region with a shorter pitch. This may compress the debris being extracted and allow fluid to flow back through the cavities and into the vessel.

Figure 25. Section view - Combination of the cage and the extractor. The cage is opened and closed by the relative movement of the proximal and distal arms. The rotating extractor rotates over the distal arm and within the proximal arm. A sheath can run over the device to ensure the cage is closed during delivery and retrieval.

Figure 26. Section view - The distal end of the device showing an embodiment of the cage with the extractor tube acting also as the distal arm. As the distal arm moves proximally, the cage opens. As the distal arm moves distally, the cage closes.

Figure 27. Section view - The tip of the device. In this embodiment the extractor tube also acts at the distal arm. The extractor tube rotates at high speed while cage connector does not rotate. The bearing facilitates this relative movement.
Figure 28. Schematic of a device of the invention with a control mechanism located within the cage.

Figure 29. Schematic of a device of the invention with a control mechanism located distally of the cage.

Figure 30. Schematic of the device having a sheath covering the elongated control member and adjustable to cover the cage.

Figure 31. Schematic of the device of the invention showing how thrombolytic agents may be infused through the extractor.

Figure 32. Schematic of the device of the invention showing how thrombolytic agents may be infused between the extractor and the sheath.

Figure 33. Schematic of the device of the invention showing how thrombolytic agents may be infused through holes or perforations formed in the sheath.

Figure 34. Schematic of a radially expansible member formed from a single strut.

Figure 35. Schematic of a radially expansible member formed from two struts.

Figure 36. An example of the radially expansible member made from a series of wires, and fused or twisted for connection (shown as a flat pattern).

Figure 37. An example of a braided expansible member with the wires twisted at the intersections to limit the movement of the wires with respect to each other (shown as a flat pattern).

Figure 38. An example of a braid configuration for the creation of a braided radially expansible member, with twisted wires (for struts) and fused ends (shown as a flat pattern).
**Figure 39.** An example of a braid configuration for the creation of a braided radially expansible member, with twisted wires to create struts.

**Detailed Description of the Invention**

**Cage and Filter**

The cage can be made in many different ways, such as from a braid, a series of wires, laser cut tubes or a combination of them. The cage also may act as a filter or be a structure for the filter at the distal part. The cage can be made of different materials such as, but not limited to, polymeric, metal such as stainless steel Nitinol or cobalt chromium, or ceramic; or a combination of these materials. The proximal side of the cage is generally open and allows thrombus into the cage. The distal part of the cage is suitably constrained onto a tube or wire with small diameter and the proximal part of the cage is connected to a tube with larger diameter. A sheath may cover the entire device at delivery and at retrieval.

A connector, which can be a wire or tube and connects the distal end of the cage to user or controls the distal movement of the cage, is called the distal arm. Another connector, which can be a wire or tube and connects the proximal end of the cage to user, is called the proximal arm.

Once the cage is assembled and opened, section 202 provides a passageway for thrombus into the cage, while section 201 prevents large thrombus passing the cage (1).

In Figs 2 and 3, an example of a cage which has been laser cut from Nitinol tube is shown. The laser cuts have the same feature as the braided mesh. It provides a passageway on the proximal side and acts as filter on the distal side. It may also carry circumferential elements for cutting through thrombus and separating thrombus from the wall.

By pushing the proximal and distal ends of the cage axially towards each other the diameter of the cage and its radial force may increase. By pulling the sides away from each other the diameter and the radial force of the cage may decrease.
Relative movement of the distal and proximal arms controls the distance between the distal and proximal sides of the cage; this adjusts the diameter and radial strength of the cage. If the distal arm is fixed and the proximal arm is pulled proximally by the user, the diameter of the cage becomes smaller. If the proximal arm is fixed and the distal arm is being pulled, the diameter of the cage increases.

The cage may also have a circumferential cutting element for removing/scraping thrombus from the vessel wall. This may be contained around or within the cage and may be composed of a round wire, flat wire, blade or a combination of these. Fig. 4 shows the cage with a flat wire attached the outside of the cage; the wire can scrape thrombus from the vessel wall. Fig 5 shows the cutting element internal to the cage, while Fig. 6 shows the cutting element of the cage as one part (in this case from a laser cut tube). The cutting/scraping element may also be tapered at an angle (Fig. 7).

A Cage with Self Adjustable Diameter Control Mechanism:
In the case of treating a tapered vessel, the cage diameter should be adjustable while it is pulled along the lumen. Also in the case of obstruction such as vascular valve it is desirable that the device be able to manoeuvre through it. Therefore a mechanism which can control the diameter of the cage by changing the resistance force is desirable. Presented here is a mechanism to control the diameter of the cage based on resistance forces.

As the cage moves through a reducing tapered vessel (pulling the proximal arm), the vessel exerts a force on the cage. When the force from the vessel exceeds the total force including the preset force from the resistance mechanism (set by the user), the distal arm moves distally relative to the proximal arm; this closes the cage. The next section includes some embodiments of the self-adjustment mechanisms.

The control mechanism may contain friction elements, springs, pneumatics, hydraulics, simple weight, or a combination of these elements.

Resistance mechanism: Sliding
In this embodiment of the invention (Figs 8 and 9), the cage (3) is made from a laser cut self-expanding tube. The cage opens when 1b and 2b move closer together, and closes when 1b and
2b move apart. The mechanism for controlling the diameter and force exerted by the cage on the vessel wall is the basis for the current invention.

The device, as shown in Fig. 8, consists of a cage (3) which has a distal end (lb) and a proximal end (2b), a distal arm (1), a proximal arm (2), and a handle. The control mechanism is comprised in the handle, and comprises a first part in the form of a housing (2a) having a guidance path, a second part in the form of a sliding block (la) configured for sliding movement along the guidance path of the housing (2a). Force controlled resistance means is provided in the form of a brake adapted to resist movement of one or the arms relative to the other. In this embodiment, the brake comprises a friction screw (4) mounted on the housing (2A) and adjustable to apply a compression force against the sliding block (la) to provide resistance to movement of one arm relative to the other, and in the case of the device being passed along a vessel that tapers inwardly, resistance to the compression of the cage which has the effect of keeping the periphery of the cage in contact with the vessel wall.

The process begins in an expanded state in the vessel as shown in Fig. 8(A). The opening and closing of the cage is governed by the relative movement of the sliding block (la) to the handle (2a). As the cage is pulled by the handle through an obstruction or the tapered section of the vessel (5), force from the vessel wall is exerted to the cage. This force is transferred to the sliding block (la). If the force applied to the block 1a is larger than the preset friction force, then the block 1a slides forward from its position 2(A) to 2(B). This allows the cage to conform to the shape of the narrower vessel. The force exerted by the cage on the vessel is therefore dictated in part by the ease of movement of the sliding block relative to the handle; this is controlled by the friction screw.

Resistance mechanism: sliding and spring

Figs. 11A and 12B show an alternative embodiment of the resistance mechanism in which parts described with reference to Fig. 10 are assigned the same reference numerals. In this embodiment, a compression spring (6) is disposed along the guide path between the housing (2a) and the sliding block (la). The housing (2a) is connected to the proximal arm (2) and the block (la) is connected to the distal arm (1). Thus, compression of the cage (3) into a contracted orientation causes the distal arm (1) to extend, causing compression of the spring. In this manner, the cage is biased into an expanded orientation when it is being passed along a vessel that is narrowing, thereby maintaining contact between the cage and the vessel wall. A friction
screw (4) is mounted on the housing (2A) and is adjustable to apply a compression force against the block (1A) and thereby provides further resistance to movement of one arm relative to the other, and in the case of the device being passed along a vessel that tapers inwardly, again provides resistance to the compression of the cage which has the effect of keeping the periphery of the cage in contact with the vessel wall.

Figs. 11A and 11B show an alternative embodiment of the resistance mechanism in which parts described with reference to Figs. 8 to 10 are assigned the same reference numerals. In this embodiment, the self adjusting mechanism comprises a movable stop (7) that is movable to adjust the length of the guidance path and, thus, the degree of compression of the spring. Thus, the force that the spring applies can be varied by changing the position of the movable stop (7) along the guidance path.

**Resistance mechanism: Spring with adjustable spring constant**

In this embodiment the distal arm (1) is attached to a flat spring (22). The other side of the spring (22) is fixed respect to proximal arm (2) through a solid handle (20). Since the spring constant of a flat spring changes with its length, a mechanism for adjusting its length (21) can slide over the spring to control its deformation. When the sliding part (21) is down (Fig 12) the spring constant is the lowest which means it allows higher displacement of inner tube (1) at a lower force. When the sliding tube (21) rises (Fig. 13), the spring constant (22) increases which allows less displacement of inner tube (1). This effectively reduces the cage diameter (3). Fig. 14 shows a 3D representation of the mechanism.

**The Extraction mechanism:**

The extraction system may consist of suction, or a rotating tube/wire with a means of transformation and/or maceration on the outer surface; the extraction system may also contain a combination of these mechanisms. The rotating extractor may be manufactured from one part or made from several attachments. For example, the extraction mechanism may comprise a wire (42) wrapped around an extraction core tube (43) (Fig. 15). This tube may be placed over the distal arm (1) of the cage and inside the proximal arm (2). Alternatively, the distal arm may be used as the extraction core tube. The distal side of the extraction core tube may be close to the distal end of the proximal arm (2). As the extraction tube turns at high speed thrombus is forced proximally between the extraction tube and the proximal arm. The distal end of the extraction mechanism may be located inside the cage (Fig. 1).
The wire may have varying cross sections along the device for different means; from circular, rectangular or triangular cross sections. The wire can be from different materials such as stainless steel, Nitinol, or polymers. Once the wire is wrapped around the tube, it might be completely fit around the tube (41) or be partially fit at a distance (43) from the rotating tube (41) surface (Fig. 16).

The proximal arm may also contain a side window (Fig. 17) to improve access of the extractor to thrombus. The extractor may also be formed from one machined part rather than a wire wrapped around a tube. The rotating extractor will likely contain sharp cutting edges at its distal end to break up matter prior to extracting it. The non-rotating proximal arm may also have a leading edge/cutting element (Fig. 18) which helps to break down thrombus, while also increasing the size of the path at the entrance of the extractor. The proximal arm may have more than one leading edge/cutting element to break down matter; Figure 19 shows the proximal arm with two leading edges. The leading edge/cutting element may form part of the proximal arm (as shown) or alternatively be a separate part attached to the non-rotating proximal arm. The extractor may also contain an additional port at the proximal end where suction can be added to enhance thrombus extraction, and to attract thrombus towards the extractor. Fig. 20 shows an embodiment in which the rotating tip and helical wire are attached to the distal arm for rotation therewith. Fig. 21 shows the extractor with the cage. The extractor may also be eccentric to the centre of the cage; Fig. 22 shows the extractor eccentric to the centre cage and adjacent to it while Fig. 23 shows the tip of the cage away from the centre of the cage. In both Figs 22 and 23 the extractor may be stationary or rotate around the centreline.

The extractor may have a varied pitch along its length. One embodiment of this is shown in Fig. 24, where a section of the extractor has a reduced pitch, and the non-rotating tube has a number of small holes (acting as a filter) in it. This may allow any debris to be squeezed, forcing residual fluid through the holes and into the vessel while the remaining debris continues to be extracted.

The extraction mechanism in combination with the cage may have various functions along the device. In one embodiment, the extractor breaks down the thrombi into relatively big pieces which are small enough to enter the inlet of tube and big enough not to pass through the filter. Once the thrombus is inside the tube, the extractor, breaks them down into smaller pieces, make
it easier to transfer along the catheter. Then, the extraction system may have a means of extraction such as helical wire or vacuum. The matter removed will be collected in a suitable collection means, for example a blood bag, or syringe or similar.

The helical formation may have a cutting edge, and which is adapted to rotate with the helical formation and cut or break up thrombus for extraction. The cutting edge may be a blade or the like, and may be disposed at or close to an end of the helical formation. The extractor tube may also have a cutting edge, and may be adapted to rotate.

Combination of the cage and extractor
The device contains both a cage with a self adjustment mechanism, and an extractor. Fig. 25 shows an embodiment of the distal end of the device. The device is opened and closed by the relative movement of the proximal and distal arms, while the extractor rotates over the distal arm and within the proximal arm. The proximal arm may also contain a window for extraction. A sheath covers the entire device during delivery and retrieval.

Fig. 26 shows another embodiment of the distal end of the device, with the cage and extractor combined. In this embodiment, the rotating extractor also acts at the distal arm; the relative movement of which opens and closes the cage. A ball bearing has been included to facilitate contact of the rotating extractor/distal arm and non-rotating proximal arm and cage (27).

Device with control mechanism within cage
Fig. 28 shows an embodiment of the device in which the control mechanism is disposed within the cage. In this embodiment, a control mechanism is provided within the cage (3) and comprises a first part operably connected to the proximal arm (2), a second part operably connected to the distal arm (1), and a helical spring (6) connecting the first and second parts. In use, as the cage passes along a vessel that is tapering inwardly, the control mechanism ensures that a force controlled resistance is applied to the cage as it contracts, thereby ensuring that the cage remains in contact with the walls of the vessel.

Device with control mechanism mounted distally of cage
Fig. 29 shows an embodiment of the device in which the control mechanism is disposed distally of the cage. In this embodiment, the device comprises a proximal arm (2), which is connected to a proximal part (2b) of the cage and which extends through, and distally beyond, the cage,
and a distal arm (1) which extends distally of the cage (3). The control mechanism comprises
a first part (block lb) operably connected to an end of the proximal arm, a second part operably
connected to the distal arm (1), and a helical spring (6). In use, as the cage passes along a vessel
that is tapering inwardly, the control mechanism ensures that a force controlled resistance is
applied to the cage as it contracts, thereby ensuring that the cage remains in contact with the
walls of the vessel.

Device with outer sheath
Specifically, Fig. 30 shows an embodiment of the device of the invention in which a sheath (8)
is provided that covers the elongated control member. In this embodiment, the device can be
manipulated such that the cage (3) is contracted and withdrawn within the sheath, which will
thus keep it in a contracted orientation

Delivery of liquid agent to vessel lumen
The device of the invention may also be employed to deliver liquid agent, for example a
thrombolytic agent which can break down thrombus, to the vessel lumen. This may be
achieved in a number of different ways including:
• The direction of rotation of the extractor screw can be changed to infuse rather than
  extract.
• Inject through the hollow distal arm.
• Inject through a lumen between the distal arm and the extractor tube (proximal arm)
  (Fig. 31)
• Injected in between the extractor tube (proximal arm) and the sheath (Figs. 32).
• Injected through cavities in the sheath (Fig. 33).
• The location of the sheath and cavities can be adjusted along the catheter length.
• One of, or a combination of, the above methods of infusion.
Generally, the liquid agent would be injected into the delivery lumen, which may be any of the
above. Alternatively, the liquid agent may be delivered slowly by means of a drip feed. As
indicated above, the liquid agent may be delivered in a number of different ways, for example
through a hollow distal arm (which has the advantage of being capable of delivering liquid
agent distally of the cage), through a lumen formed between the distal arm and the proximal
arm (also referred to as the extractor tube), or through a lumen formed between the proximal
arm and the outer sheath.
Design of radially expansible member

Figs 34 to 39 describe embodiments of the radially expansible member forming part of the device of the invention, and specifically braid configurations that may be employed in radially expansible members.

The invention is not limited to the embodiments hereinbefore described which may be varied in construction and detail without departing from the spirit of the invention.
CLAIMS

1. A device suitable for use in a body lumen (5) and comprising:
   - an elongated control member having a distal and a proximal end, and a radially
     expansible member (3) disposed at or near the distal end and adapted for collection
     and/or shearing of matter from a wall of the body lumen,
   - the radially expansible member having a proximal end (2B) and a distal end (IB) and
     being adjustable between a contracted orientation and an expanded orientation,
   - the elongated control member comprising a proximal arm (2) operatively connected to
     the radially expansible member (3) and a tubular distal arm (1) operatively connected
     to the radially expansible member distally of the proximal arm connection such that
     movement of one arm relative to the other arm effects adjustment of the diameter and/or
     radial strength of the radially expansible member,
   - a control mechanism operatively connected to both arms and adapted to provide
     resistance to the movement of one arm relative to the other, in which the control
     mechanism comprises a first part (2A) connected to one of the arms, a second part (1A)
     connected to the other of the arms that is movable relative to the first part, and a force
     controlled, resistance means (4, 6) for resisting movement of the first part relative to
     the second part,

characterised in that the force controlled resistance means comprises (a) biasing means (6) for
biasing the radially expansible member (3) into the expanded orientation, or (b) a brake means
(4) adapted to resist movement of one of the first and second parts relative to the other of the
first and second parts.

2. A device suitable for use in a body lumen (5) and comprising:
   - an elongated control member having a distal and a proximal end, and a radially
     expansible member (3) disposed at or near the distal end and adapted for collection
     and/or shearing of matter from a wall of the body lumen,
   - the radially expansible member having a proximal end (2B) and a distal end (IB) and
     being adjustable between a contracted orientation and an expanded orientation,
   - a thrombus extractor (42.43),
   - the elongated control member comprising a proximal arm (2) operatively connected to
     the radially expansible member and a distal arm (1) operatively connected to the
     radially expansible member distally of the proximal arm connection such that
movement of one arm relative to the other arm effects adjustment of the diameter and/or radial strength of the radially expansible member,
- a control mechanism operatively connected to both arms and adapted to provide resistance to the movement of one arm relative to the other, in which the control mechanism comprises a first part (2A) connected to one of the arms, a second part (1A) connected to the other of the arms that is movable relative to the fixed first part, and force controlled, resistance means (4, 6) for resisting movement of the first part relative to the second part,
characterised in that the force controlled resistance means comprises (a) biasing means (6) for biasing the radially expansible member into the expanded orientation, or (b) a brake means (4) adapted to resist movement of one of the first and second parts relative to the other of the first and second parts.

3. A device as claimed in Claim 1 or 2 in which the force controlled resistance means comprises (a) biasing means (6) for biasing the radially expansible member into the expanded orientation, and (b) a brake means (4) adapted to resist movement of one of the first and second parts relative to the other of the first and second parts.

4. A device as claimed in any preceding Claim in which the biasing means (6) comprises a resiliently deformable or displacable member.

5. A device as claimed in any preceding Claim in which the resiliently deformable member comprises a helical spring (6).

6. A device as claimed in Claim 4 or 5 in which the resiliently deformable member is disposed between the first and second parts and configured to provide force controlled resistance to movement of the first and second parts together.

7. A device as claimed in claimed in any preceding Claim in which the brake means comprises a friction screw (4).

8. A device as claimed in any preceding Claim in which the control mechanism is disposed proximally, distally, or within, the radially expansible member.
9. A device as claimed in Claim 8 in which the control mechanism is disposed proximally of the radially expansible member such that in use it is located outside of the body, wherein the first part (2A) is operably connected to the proximal arm (2) and comprises a guide path, and the second part (1A) is operably connected to the distal arm (1) and is associated with the first part for movement along the guide path, and wherein the force controlled resistance means comprises a resiliently deformable member (6) disposed along the guide path.

10. A device as claimed in Claim 9 in which the first part (2A) comprises a movable stop (7) which is movable to vary the length of the guide path and/or the resiliently deformable member.

11. A device as claimed in Claim 10 in which the brake means comprises a friction screw (4) operably connected to one of the distal or proximal arms and configured for adjustable engagement with the other of the distal or proximal arms.

12. A device as claimed in Claim 11 in which the brake means comprises a friction screw (4) operably connected to the distal arm and configured for adjustable engagement with the proximal arm.

13. A device as claimed in Claim 8 in which the control mechanism is disposed within the radially expansible member and comprises a first part connected to the distal arm, a second part connected to the proximal arm, and a helical spring operatively connected to the first and second parts and configured to provide force controlled resistance to movement of the first and second parts together.

14. A device as claimed in any preceding Claim in which the radially expansible member comprises a cage (3).

15. A device as claimed in Claim 14 in which the cage comprises a proximal portion having apertures for receipt of thrombus into the cage, and a distal portion having a fine mesh for capturing thrombus, in which the cage optionally comprises a cut tube.

16. A device as claimed in Claim 14 in which the cage comprises a distal portion having apertures for receipt of thrombus into the cage, and a proximal portion having a fine mesh for capturing thrombus, in which the cage optionally comprises a cut tube.
17. A device as claimed in any preceding Claim and comprising an extractor at least a part of which is disposed within the radially expansible member.

18. A device as claimed in Claim 18 in which the extractor comprises a helical formation (42) adapted to rotate.

19. A device as claimed in Claim 18 in which the helical formation (42) is adapted to rotate to (a) remove thrombus from the radially expansible member, and/or (b) deliver an agent to the body lumen.

20. A device as claimed in Claim 18 or 19 in which the helical formation (42) shortens in pitch such that in use it squeezes blood from solid thrombus.

21. A device as claimed in Claim 18, 19 or 20 in which the helical formation (42) is disposed within an extractor tube (43), wherein one or more portion of the helical formations within the radially expansible member is exposed.

22. A device as claimed in Claim 21 in which the extractor tube (43) comprises holes to allow blood pass out of the extractor tube.

23. A device as claimed in Claim 21 or 22, in which a leading edge of the extractor tube comprises a cutting edge.

24. A device as claimed in any of Claims 18 to 23 in which the helical formation comprises a cutting edge.

25. A device as claimed in Claim 17 in which the extractor is an aspirator tube.

26. A device as claimed in Claim 25 in which the device comprises a rotating cutting edge disposed within the radially expansible member.

27. A device as claimed in any preceding Claim, in which the force controlled resistance means is self-adjusting.
28. A device as claimed in any preceding Claim in which the proximal arm (2) comprises a lumen and the distal arm (1) is at least partly disposed within the lumen of the first arm, optionally coaxially with the first arm.

29. A device as claimed in any preceding Claim and comprising a tubular sheath, in which the elongated control member is contained within the tubular sheath.

30. A device as claimed in Claim 29 in which the tubular sheath can be adjusted to cover the radially expansible member and maintain the radially expansible member in a contracted orientation.

31. A device as claimed in any of Claims 29 or 30 in which the tubular sheath comprises a plurality of holes for perfusion of a liquid.

32. A device as claimed in any preceding Claim and comprising a liquid administration lumen configured to deliver a liquid into the body.

33. A device as claimed in Claim 33 in which the liquid administration lumen comprises a lumen formed within the distal arm, between the distal arm and proximal arm, and between the proximal arm and an external sheath.

34. A device as claimed in Claim 32 or 33 and including an injection port for delivery of liquid into the liquid administration lumen.

35. A device as claimed in any of Claims 14 to 34 in which the cage comprises a single strut, two struts, or a multiplicity of struts.

36. A device as claimed in any preceding in which the radially expansible member comprises a leading edge.

37. A device as claimed in any preceding Claim which is a thrombectomy device suitable for removing thrombus from large and/or tapering blood vessels.

38. A method of removing thrombus from a blood vessel (5) comprising the steps of placing the device of any of Claims 1 to 37 in the blood vessel, and moving the device along the blood vessel.
39. A method as claimed in Claim 38 in which the device is moved along the blood vessel in an inwardly tapering direction or an outwardly tapering direction.

40. A method as claimed in Claim 38 or 39 in which the blood vessel is a vein.

41. A method of removing thrombus from a blood vessel comprising steps of placing a device of any of Claims 1 to 37 into the blood vessel in an expanded configuration, injecting thrombolytic agent into the lumen of the blood vessel, and then operating the device to remove thrombus from the blood vessel.

42. A method as claimed in Claim 41 in which the device is operated to remove thrombus by moving the device along the blood vessel.

43. A method as claimed in Claim 41 in which the device comprises an extractor, and the device is operated to remove thrombus by actuation of the extractor.

44. A method as claimed in any of Claims 41 to 43 in which the thrombolytic agent is injected through the liquid administration lumen.
Figure 4.
Figure 6.

Figure 7.
Figure 8.

Figure 9.
Figure 10.

Figure 11
Figure 12.

Figure 13.
Figure 14.

Figure 15.
Figure 18.
Figure 19.

Figure 20.
Figure 24.
Figure 25.

Figure 26.
Figure 27.
Figure 36

Connection point/ fused

Struts of the expansible member
Figure 39

2x twisted wires

4x twisted wires
A. CLASSIFICATION OF SUBJECT MATTER
INV. A61B17/221 A61B17/3207

ADD.

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

B. FIELDS SEARCHED

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

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

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

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
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<tr>
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<th>Relevant to claim No.</th>
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<td>X</td>
<td>US 2013/066346 Al (PIGOTT) 14 March 2013 (2013-03-14) abstract; figures 1,5 paragraphs [0023] - [0032]</td>
<td>1, 2, 7, 8, 14, 28-37</td>
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<td>X</td>
<td>US 6 146 396 A (KONYA ET AL.) 14 November 2000 (2000-11-14)</td>
<td>1-6, 8-12, 14, 27, 28, 35-37</td>
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<td>Y</td>
<td>figures 7,9A,9B, 10 column 11, line 57 - column 14, line 17</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 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

"I" 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
"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"S" document member of the same patent family

Date of the actual completion of the international search: 19 June 2014

Date of mailing of the international search report: 07/07/2014

Name and mailing address of the ISA:
European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040
Fax: (+31-70) 340-3018

Authorized officer
Gimenez Burgos, R
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<td>X</td>
<td>WO 01/87168 A1 (LAMY-PERRET) 22 November 2001 (2001-11-22)</td>
<td>1-8, 11, 12, 14-16, 27, 28, 35-37</td>
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<td>Y</td>
<td>figures 1-9 page 3, line 8 - page 5, line 7</td>
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<td>A</td>
<td>US 6 454 775 B1 (DEMARIAIS ET AL.) 24 September 2002 (2002-09-24) abstract; figures</td>
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<td>A</td>
<td>US 4 909 789 A (TAGUCHI ET AL.) 20 March 1990 (1990-03-20) figures 1-4 column 1, line 57 - column 2, line 5 column 2, line 62 - column 3, line 42</td>
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<td>A</td>
<td>US 5 573 530 A (FLEURY ET AL.) 12 November 1996 (1996-11-12) figures 1-3 column 2, line 59 - column 3, line 60 column 5, lines 48-58</td>
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<td>X, P</td>
<td>WO 2013/109756 A2 (PERFLOW MEDICAL LTD.) 25 July 2013 (2013-07-25)</td>
<td>1, 2, 4-6, 8, 9, 14, 27-30, 36, 37</td>
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<td>L</td>
<td>figures 1A-1C, 3A-4,5D paragraphs [0006], [0007], [0120] - [0122]</td>
<td>1-6, 8-14, 27-34, 36, 37</td>
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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. ☐ Claims Nos.: 38-44  
   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

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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