Title: HIGH PRESSURE BALLOON SHOCKWAVE CATHETER AND METHOD

Abstract: A system and method for breaking obstructions in body lumens includes a catheter including an elongated carrier, a balloon at one end of the carrier in sealed relation thereto, the carrier including a channel arranged to receive a fluid that fills and pressurizes the balloon to an internal pressure of greater than two atmospheres, and an arc generator including at least one electrode within the balloon that forms a mechanical shock wave within the balloon. The system further includes a power source that provides electrical energy to the arc generator.
HIGH PRESSURE BALLOON SHOCKWAVE CATHETER AND METHOD

PRIORITY CLAIM

[1] The present application claims the priority of U.S. Application No. 13/267,383 filed on October 6, 2011, which application claims the benefit of United States Provisional Patent Application No. 61/439,633, filed February 4, 2011, which application are incorporated herein by references in their entireties.

BACKGROUND OF THE INVENTION

[2] The present invention relates to a treatment system for percutaneous coronary angioplasty or peripheral angioplasty in which a dilation catheter is used to cross a lesion in order to dilate the lesion and restore normal blood flow in the artery. It is particularly useful when the lesion is a calcified lesion in the wall of the artery. Calcified lesions require high pressures (sometimes as high as 10-15 or even 30 atmospheres) to break the calcified plaque and push it back into the vessel wall. With such pressures comes trauma to the vessel wall which can contribute to vessel rebound, dissection, thrombus formation, and a high level of restenosis. Non-concentric calcified lesions can result in undue stress to the free wall of the vessel when exposed to high pressures. An angioplasty balloon when inflated to high pressures can have a specific maximum diameter to which it will expand but the opening in the vessel under a concentric lesion will typically be much smaller. As the pressure is increased to open the passage way for blood the balloon will be confined to the size of the opening in the calcified lesion (before it is broken open). As the pressure builds a
A tremendous amount of energy is stored in the balloon until the calcified lesion breaks or cracks. That energy is then released and results in the rapid expansion of the balloon to its maximum dimension and may stress and injure the vessel walls.

**SUMMARY OF THE INVENTION**

[3] The invention provides a catheter comprising an elongated carrier and a balloon at one end of the carrier in sealed relation thereto. The carrier includes a channel arranged to receive a fluid therein that inflates the balloon to an internal pressure of greater than two atmospheres. The catheter further comprises an arc generator within the balloon that forms a mechanical shock wave within the balloon.

[4] The balloon may be formed of non-compliant material. Alternatively, the balloon may be formed of compliant material. The catheter may further comprise a sensor that senses reflected energy.

[5] The invention further provides a system comprising a catheter including an elongated carrier and a balloon at one end of the carrier in sealed relation thereto. The carrier includes a channel arranged to receive a fluid therein that inflates the balloon to an internal pressure above two atmospheres. An arc generator within the balloon forms a mechanical shock wave within the balloon. The system further includes a power source that provides electrical energy to the arc generator. The balloon may be formed of non-compliant material or a compliant material. The system may further comprise a sensor that senses reflected energy.

[6] The invention still further provides a method comprising providing a catheter including an elongated carrier, a balloon at one end of the carrier in sealed
relation thereto, the carrier including a channel arranged to receive a fluid therein that inflates the balloon, and an arc generator including at least one electrode within the balloon that forms a mechanical shock wave within the balloon, inserting the catheter into a body lumen of a patient adjacent to an obstruction of the body lumen, admitting fluid into the carrier channel to inflate the balloon to an internal pressure above two atmospheres, and applying high voltage pulses to the arc generator to form a series of mechanical shocks within the balloon. The method may further include the step of sensing reflected energy within the catheter.

**BRIEF DESCRIPTION OF THE DRAWINGS**

[7] The features of the present invention which are believed to be novel are set forth with particularity in the appended claims. The invention, together with further features and advantages thereof, may best be understood by making reference to the following description taken in conjunction with the accompanying drawings, in the several figures of which like reference numerals identify identical elements, and wherein:

[8] **Fig. 1** is a view of the therapeutic end of a typical prior art over-the-wire angioplasty balloon catheter;

[9] **Fig. 2** is a side view of a dilating angioplasty balloon catheter with two electrodes within the balloon attached to a source of high voltage pulses according to one embodiment of the invention;

[10] **Fig. 3** is a schematic of a high voltage pulse generator;
Fig. 3A shows voltage pulses that may be obtained with the generator of Fig. 3;

Fig. 4 is a side view of the catheter of Fig. 2 showing an arc between the electrodes and simulations of the shock wave flow;

Fig. 5 is a side view of a dilating catheter with insulated electrodes within the balloon and displaced along the length of the balloon according to another embodiment of the invention;

Fig. 6 is a side view of a dilating catheter with insulated electrodes within the balloon displaced with a single pole in the balloon and a second being the ionic fluid inside the balloon according to a further embodiment of the invention;

Fig. 7 is a side view of a dilating catheter with insulated electrodes within the balloon and studs to reach the calcification according to a still further embodiment of the invention;

Fig. 8 is a side view of a dilating catheter with insulated electrodes within the balloon with raised ribs on the balloon according to still another embodiment of the invention;

Fig. 8A is a front view of the catheter of Fig. 8;

Fig. 9 is a side view of a dilating catheter with insulated electrodes within the balloon and a sensor to detect reflected signals according to a further embodiment of the invention;

Fig. 10 is a pressure volume curve of a prior art balloon breaking a calcified lesion;

Fig. 10A is a sectional view of a balloon expanding freely within a vessel;
[21] Fig. 10B is a sectional view of a balloon constrained to the point of breaking in a vessel;

[22] Fig. IOC is a sectional view of a balloon after breaking within the vessel;

[23] Fig. 11 is a pressure volume curve showing the various stages in the breaking of a calcified lesion with shock waves according to an embodiment of the invention;

[24] Fig. 11A is a sectional view showing a compliant balloon within a vessel;

[25] Fig. 11B is a sectional view showing pulverized calcification on a vessel wall;

[26] Fig. 12 illustrates shock waves delivered through the balloon wall and endothelium to a calcified lesion;

[27] Fig. 13 shows calcified plaque pulverized and smooth a endothelium restored by the expanded balloon after pulverization;

[28] Fig. 14 is a schematic of a circuit that uses a surface EKG to synchronize the shock wave to the "R" wave for treating vessels near the heart;

[29] Fig. 15 is a side view, partly cut away, of a dilating catheter with a parabolic reflector acting as one electrode and provides a focused shock wave inside a fluid filled compliant balloon; and

[30] Fig. 16 is a chart illustrating relative Shockwave energy delivered versus internal balloon pressure above ambient pressure for a fixed Shockwave creating voltage.

DETAILED DESCRIPTION OF THE INVENTION

[31] Fig. 1 is a view of the therapeutic end of a typical prior art over-the-wire angioplasty balloon catheter 10. Such
catheters are usually non-compliant with a fixed maximum dimension when expanded with a fluid such as saline.

[32] Fig. 2 is a view of a dilating angioplasty balloon catheter 20 according to an embodiment of the invention. The catheter 20 includes an elongated carrier, such as a hollow sheath 21, and a dilating balloon 26 formed about the sheath 21 in sealed relation thereto at a seal 23. The balloon 26 forms an annular channel 27 about the sheath 21 through which fluid, such as saline, may be admitted into the balloon to inflate the balloon. The channel 27 further permits the balloon 26 to be provided with two electrodes 22 and 24 within the fluid filled balloon 26. The electrodes 22 and 24 are attached to a source of high voltage pulses 30. The electrodes 22 and 24 are formed of metal, such as stainless steel, and are placed a controlled distance apart to allow a reproducible arc for a given voltage and current. The electrical arcs between electrodes 22 and 24 in the fluid are used to generate shock waves in the fluid. The variable high voltage pulse generator 30 is used to deliver a stream of pulses to the electrodes 22 and 24 to create a stream of shock waves within the balloon 26 and within the artery being treated (not shown). The magnitude of the shock waves can be controlled by controlling the magnitude of the pulsed voltage, the current, the duration and repetition rate. The insulating nature of the balloon 26 protects the patient from electrical shocks.

[33] The balloon 26 may be filled with water or saline in order to gently fix the balloon in the walls of the artery in the direct proximity with the calcified lesion. The fluid may also contain an x-ray contrast to permit fluoroscopic viewing of the catheter during use. The carrier 21 includes a lumen 29 through which a guidewire (not shown) may be inserted to guide the catheter into position. Once positioned the physician or operator can start with low energy shock waves.
and increase the energy as needed to crack the calcified plaque. Such Shockwaves will be conducted through the fluid, through the balloon, through the blood and vessel wall to the calcified lesion where the energy will break the hardened plaque without the application of excessive pressure by the balloon on the walls of the artery.

[34] Fig. 3 is a schematic of the high voltage pulse generator 30. Fig. 3A shows a resulting waveform. The voltage needed will depend on the gap between the electrodes and generally 100 to 10,000 volts. The high voltage switch 32 can be set to control the duration of the pulse. The pulse duration will depend on the surface area of the electrodes 22 and 24 and needs to be sufficient to generate a gas bubble at the surface of the electrode causing a plasma arc of electric current to jump the bubble and create a rapidly expanding and collapsing bubble, which creates the mechanical shock wave in the balloon. Such shock waves can be as short as a few microseconds.

[35] Fig. 4 is a cross sectional view of the Shockwave catheter 20 showing an arc 25 between the electrodes 22 and 24 and simulations of the shock wave flow 28. The shock wave 28 will radiate out from the electrodes 22 and 24 in all directions and will travel through the balloon 26 to the vessel where it will break the calcified lesion into smaller pieces.

[36] Fig. 5 shows another dilating catheter 40. It has insulated electrodes 42 and 44 within the balloon 46 displaced along the length of the balloon 46.

[37] Fig. 6 shows a dilating catheter 50 with an insulated electrode 52 within the balloon 56. The electrode is a single electrode pole in the balloon, a second pole being the ionic fluid 54 inside the balloon. This unipolar
configuration uses the ionic fluid as the other electrical pole and permits a smaller balloon and catheter design for low profile balloons. The ionic fluid is connected electrically to the HV pulse generator 30.

[38] Fig. 7 is another dilating 60 catheter with electrodes 62 and 64 within the balloon 66 and studs 65 to reach the calcification. The studs 65 form mechanical stress risers on the balloon surface 67 and are designed to mechanically conduct the shock wave through the intimal layer of tissue of the vessel and deliver it directly to the calcified lesion.

[39] Fig. 8 is another dilating catheter 70 with electrodes 72 and 74 within the balloon 76 and with raised ribs 75 on the surface 77 of the balloon 76. The raised ribs 75 (best seen in Fig. 8A) form stress risers that will focus the Shockwave energy to linear regions of the calcified plaque.

[40] Fig. 9 is a further dilating catheter 80 with electrodes 82 and 84 within the balloon 86. The catheter 80 further includes a sensor 85 to detect reflected signals. Reflected signals from the calcified plaque can be processed by a processor 88 to determine quality of the calcification and quality of pulverization of the lesion.

[41] Fig. 10 is a pressure volume curve of a prior art balloon breaking a calcified lesion. Fig. 10B shows the buildup of energy within the balloon (region A to B) and Fig. IOC shows the release of the energy (region B to C) when the calcification breaks. At region C the artery is expanded to the maximum dimension of the balloon. Such a dimension can lead to injury to the vessel walls. Fig. 10A shows the initial inflation of the balloon.
Fig. 11 is a pressure volume curve showing the various stages in the breaking of a calcified lesion with shock waves according to the embodiment. The balloon is expanded with a saline fluid and can be expanded to fit snugly to the vessel wall (Region A) (Fig. 11A) but this is not a requirement. The pressurization of the balloon may be provided by the physician using a commonly available insulflator as is well known in the art. As the High Voltage pulses generate shock waves (Region B and C) extremely high pressures, extremely short in duration will chip away the calcified lesion slowly and controllably expanding the opening in the vessel to allow blood to flow un-obstructed (Fig. 11B).

Fig. 12 shows, in a cutaway view, shock waves 98 delivered in all directions through the wall 92 of a saline filled balloon 90 and intima 94 to a calcified lesion 96. The shock waves 98 pulverize the lesion 96. The balloon wall 92 may be formed of non-compliant or compliant material to contact the intima 94.

Fig. 13 shows calcified plaque 96 pulverized by the shock waves. The intima 94 is smoothed and restored after the expanded balloon (not shown) has pulverized and reshaped the plaque into the vessel wall.

Fig. 14 is a schematic of a circuit 100 that uses the generator circuit 30 of Fig. 3 and a surface EKG 102 to synchronize the shock wave to the "R" wave for treating vessels near the heart. The circuit 100 includes an R-wave detector 102 and a controller 104 to control the high voltage switch 32. Mechanical shocks can stimulate heart muscle and could lead to an arrhythmia. While it is unlikely that Shockwaves of such short duration as contemplated herein would stimulate the heart, by synchronizing the pulses (or bursts of pulses) with the R-wave, an additional degree of safety is
provided when used on vessels of the heart or near the heart. While the balloon in the current drawings will provide an electrical isolation of the patient from the current, a device could be made in a non-balloon or non-isolated manner using blood as the fluid. In such a device, synchronization to the R-wave would significantly improve the safety against unwanted arrhythmias.

[46] Fig. 15 shows a still further dilation catheter 110 wherein a shock wave is focused with a parabolic reflector 114 acting as one electrode inside a fluid filled compliant balloon 116. The other electrode 112 is located at the coaxial center of the reflector 114. By using the reflector as one electrode, the shock wave can be focused and therefore pointed at an angle (45 degrees, for example) off the center line 111 of the catheter artery. In this configuration, the other electrode 112 will be designed to be at the coaxial center of the reflector and designed to arc to the reflector 114 through the fluid. The catheter can be rotated if needed to break hard plaque as it rotates and delivers Shockwaves.

[47] In accordance with further aspects of the invention, improved therapeutic effect may be obtained if the fluid within the balloon not only fills the balloon, but pressurizes it. Fig. 16 is a graph illustrating relative Shockwave energy delivered versus balloon pressure (pressure within the balloon above ambient pressure) for a fixed Shockwave creating voltage. More particularly, as may be seen in the chart of Fig. 16, as a balloon, such as balloon 26 of Fig. 2, is pressurized, the amount of energy transmitted varies for a fixed Shockwave creating voltage. The transmitted energy decreases between zero and two atmospheres of balloon pressure. Above two atmospheres the transmitted Shockwave energy improves. At six atmospheres of balloon pressure the transmitted Shockwave energy is nearly equal to the energy
that would be transmitted if there were no balloon at all and
at eight atmospheres of balloon pressure the transmitted
Shockwave energy is actually higher than if there were no
balloon material being in the path of the Shockwave energy.
Beyond eight atmospheres of balloon pressure, the transmitted
energy rises.

[48] Hence, as may be seen from the foregoing, the
combination of high pressure, above two atmospheres, and
delivering a Shockwave by electrical discharge in a field
inside a balloon is desirable. Further, by creating a
Shockwave inside of a pressurized field, one can release more
energy from the same Shockwave discharge than from a free
field and thus enable increased therapeutic effect at an
equivalent Shockwave discharge level.

[49] The above runs counter to intuitive thinking. It has
long been thought that the balloon can adversely affect the
amount of Shockwave energy that is transmitted through it.
Twenty percent of the Shockwave strength can be absorbed or
reflected by the balloon material. Softer more pliable
materials may absorb less but such materials are less
effective at dilation of a vessel. Generally, more
noncompliant materials are desired. Unfortunately, the
materials adversely affect the transmitted energy. For these
reasons, the effect of increasing the transmitted Shockwave
energy for a given Shockwave discharge energy is a most
unexpected and desirable result.

[50] While particular embodiments of the present
invention have been shown and described, modifications may be
made. It is therefore intended in the appended claims to
cover all such changes and modifications which fall within the
true spirit and scope of the invention as defined by those
claims.
What is claimed is:

1. A catheter comprising:
   an elongated carrier;
   a balloon at one end of the carrier in sealed relation thereto,
   the carrier including a channel arranged to receive a fluid therein that inflates the balloon to an internal pressure of greater than two atmospheres; and
   an arc generator within the balloon that forms a mechanical shock wave within the balloon.

2. The catheter of claim 1, wherein the balloon is formed of non-compliant material.

3. The catheter of claim 1, wherein the balloon is formed of compliant material.

4. The catheter of claim 1, further comprising a sensor that senses reflected energy.

5. A system comprising:
   a catheter including an elongated carrier, a balloon at one end of the carrier in sealed relation thereto, the carrier including a channel arranged to receive a fluid therein that inflates the balloon to an internal pressure above two atmospheres, and an arc generator within the balloon that forms a mechanical shock wave within the balloon and
   a power source that provides electrical energy to the arc generator.
6. The system of claim 5, wherein the balloon is formed of non-compliant material.

7. The system of claim 5, wherein the balloon is formed of compliant material.

8. The system of claim 5, further comprising a sensor that senses reflected energy.

9. A method comprising:
   providing a catheter including an elongated carrier, a balloon at one end of the carrier in sealed relation thereto, the carrier including a channel arranged to receive a fluid therein that inflates the balloon, and an arc generator including at least one electrode within the balloon that forms a mechanical shock wave within the balloon;
   inserting the catheter into a body lumen of a patient adjacent to an obstruction of the body lumen;
   admitting fluid into the carrier channel to inflate the balloon to an internal pressure above two atmospheres; and
   applying high voltage pulses to the arc generator to form a series of mechanical shocks within the balloon.

10. The method of claim 9, including the further step of sensing reflected energy within the catheter.
FIG. 14

SKIN EKG

R WAVE DETECTOR

CONTROLLER FOR HV SWITCH

VARIABLE HV SOURCE

ARC