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(54) **PORTRABLE REGULATED VACUUM PUMP FOR MEDICAL PROCEDURES**

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(52) **U.S. Cl.** **606/123; 606/201**

(58) **Field of Classification Search** **600/37, 600/202; 601/6, 7, 43; 602/32; 604/35, 604/129, 176; 606/115, 123, 201–203**

See application file for complete search history.

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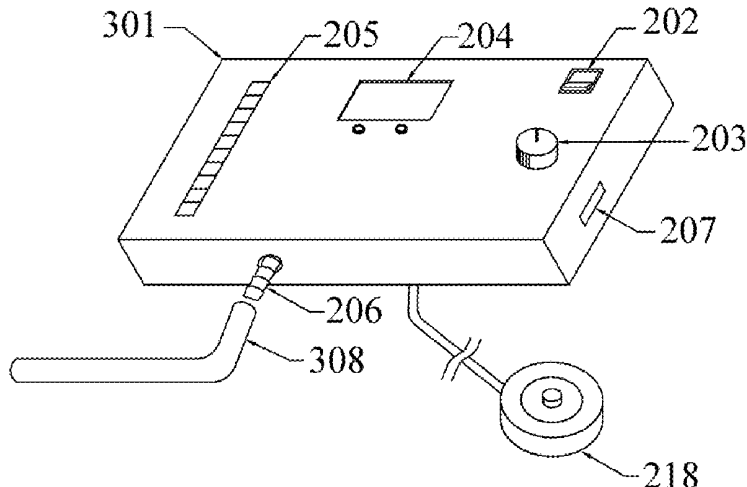
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(57) **ABSTRACT**

In various embodiments of the invention, an electro-vacuum pump can be used either as a stand alone unit to create, monitor and control a vacuum or to maintain, assist or as a back up for a hand operated pump in a variety of surgical procedures. The vacuum pump is controlled by a processor and is automatically activated when the vacuum is below a pre-selected parameter or a parameter selected on the fly by the surgeon. Visual or audio feedback is used to allow the surgeon increased control of the vacuum device while carrying out the surgical procedure.

20 Claims, 10 Drawing Sheets



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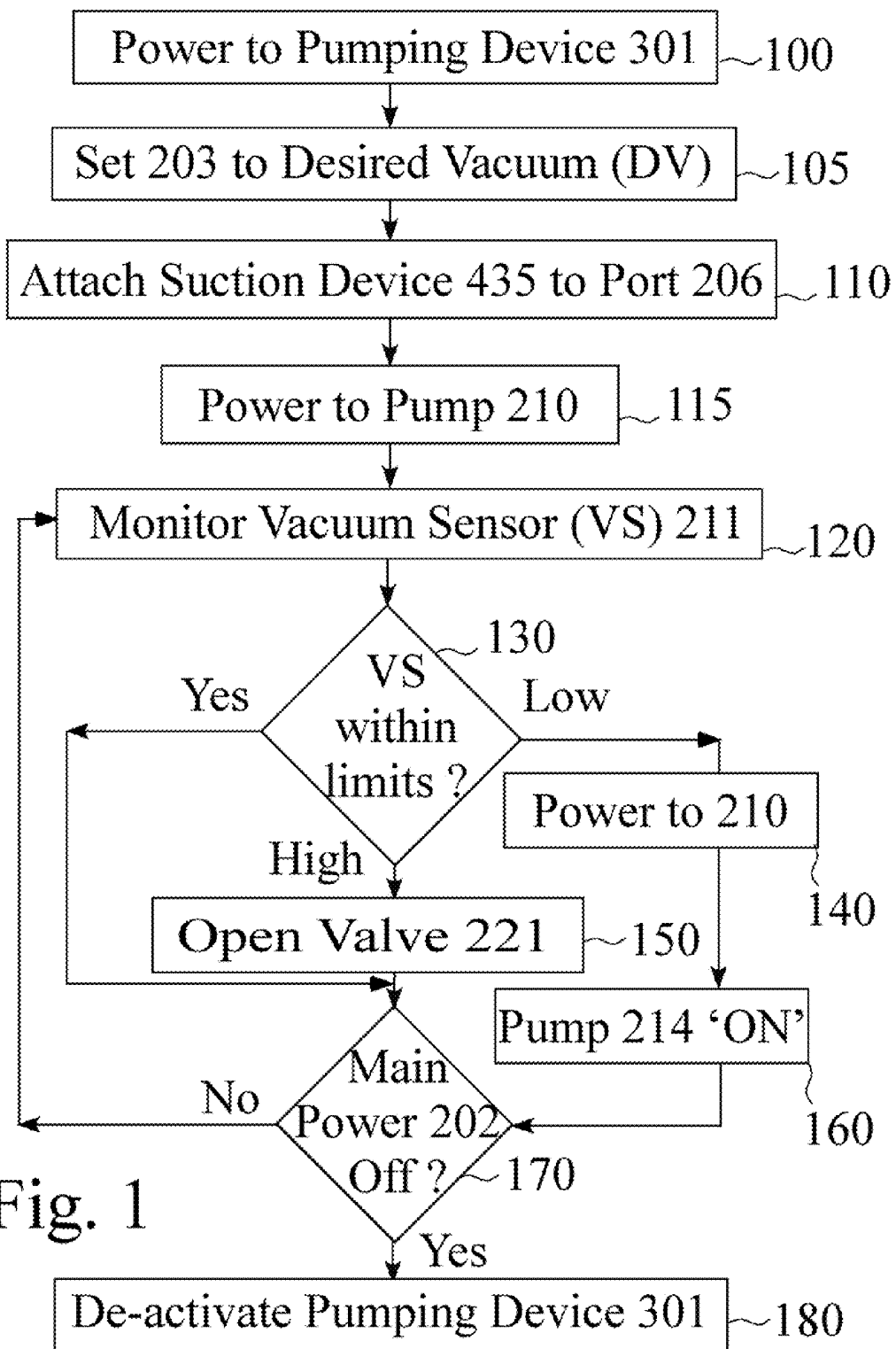


Fig. 1

Fig. 2

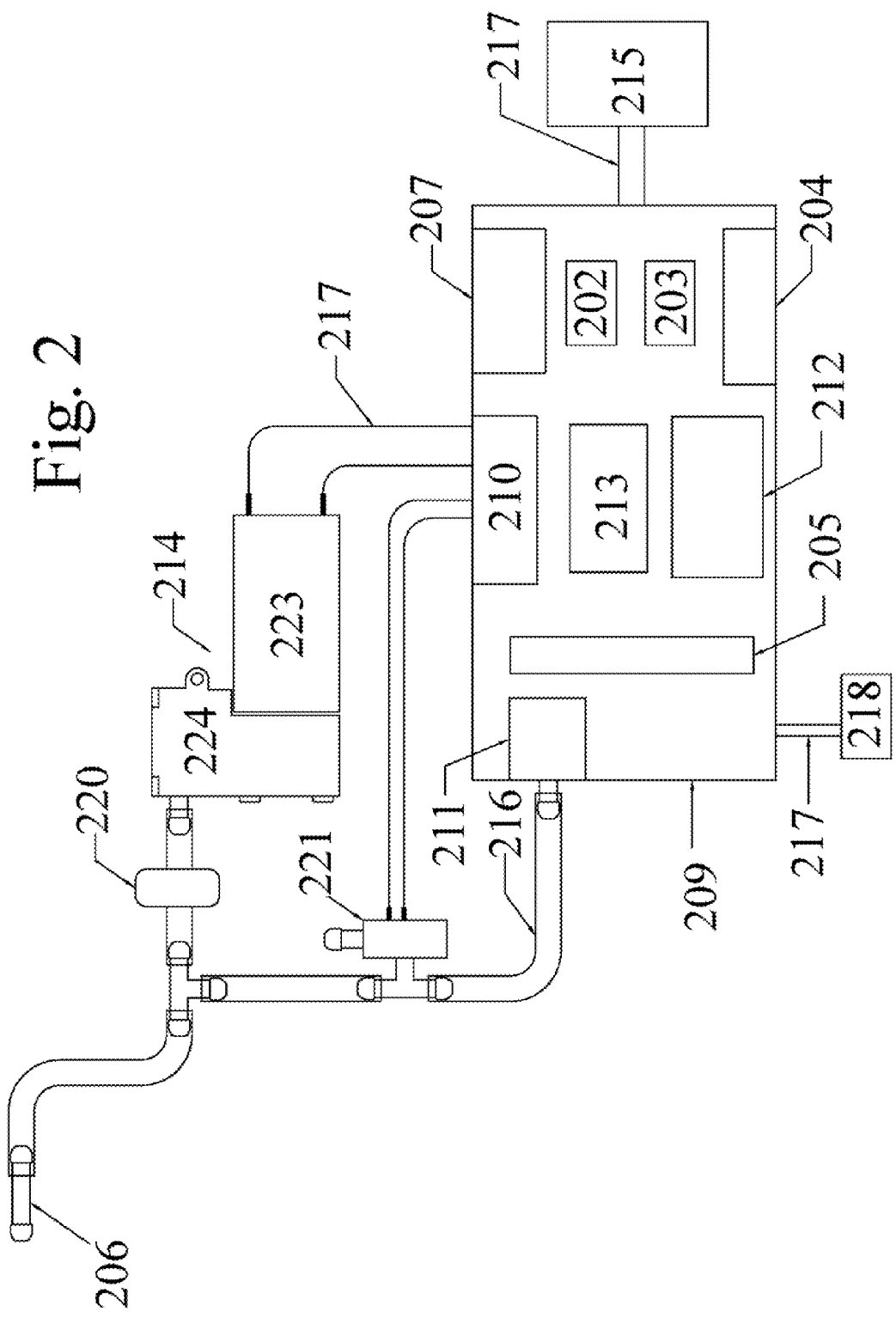


Fig. 3

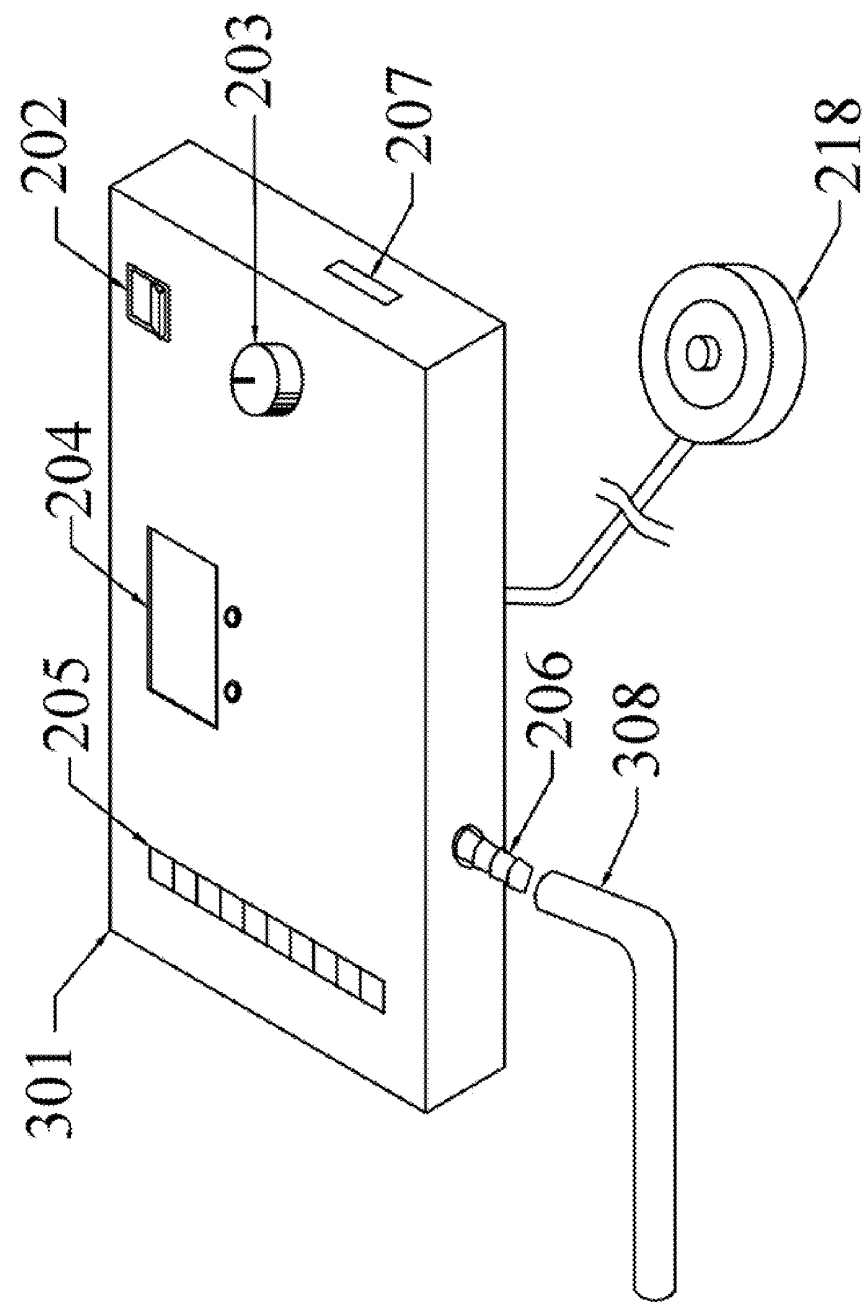


Fig. 4

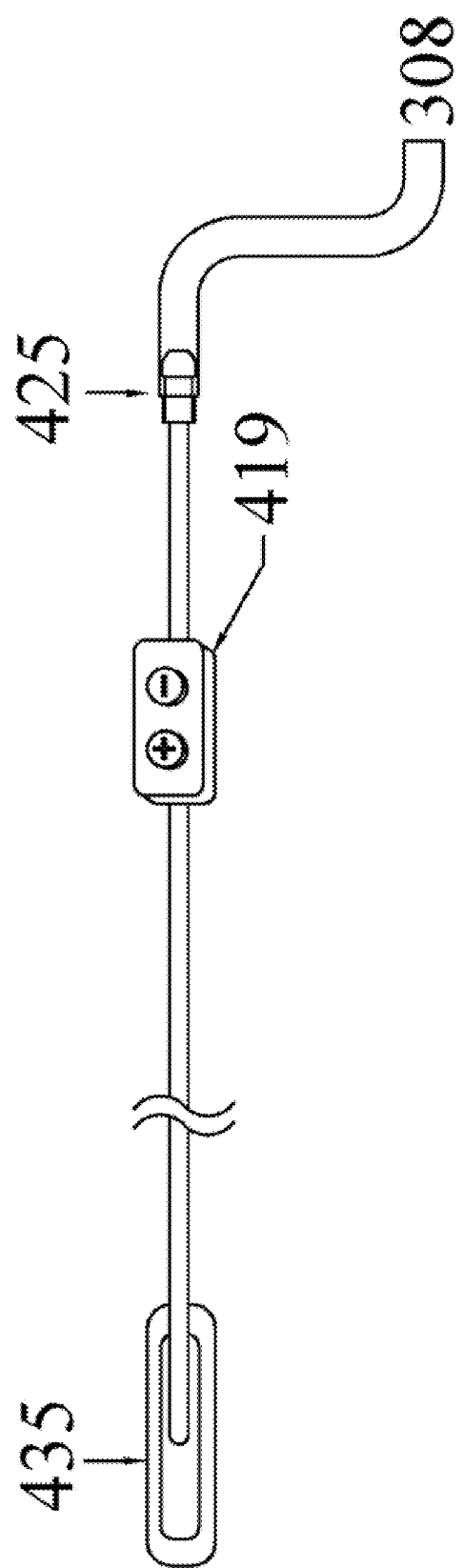
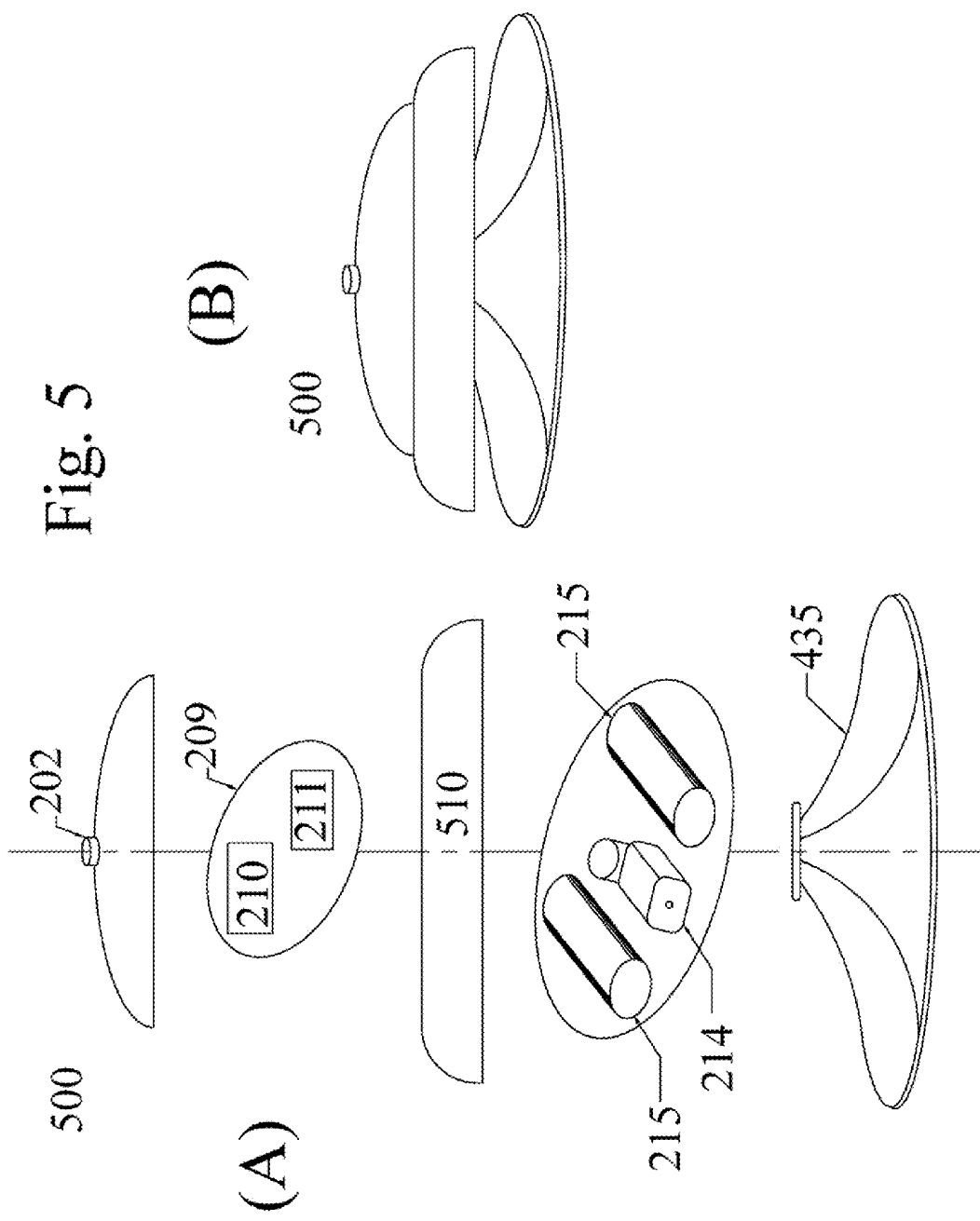
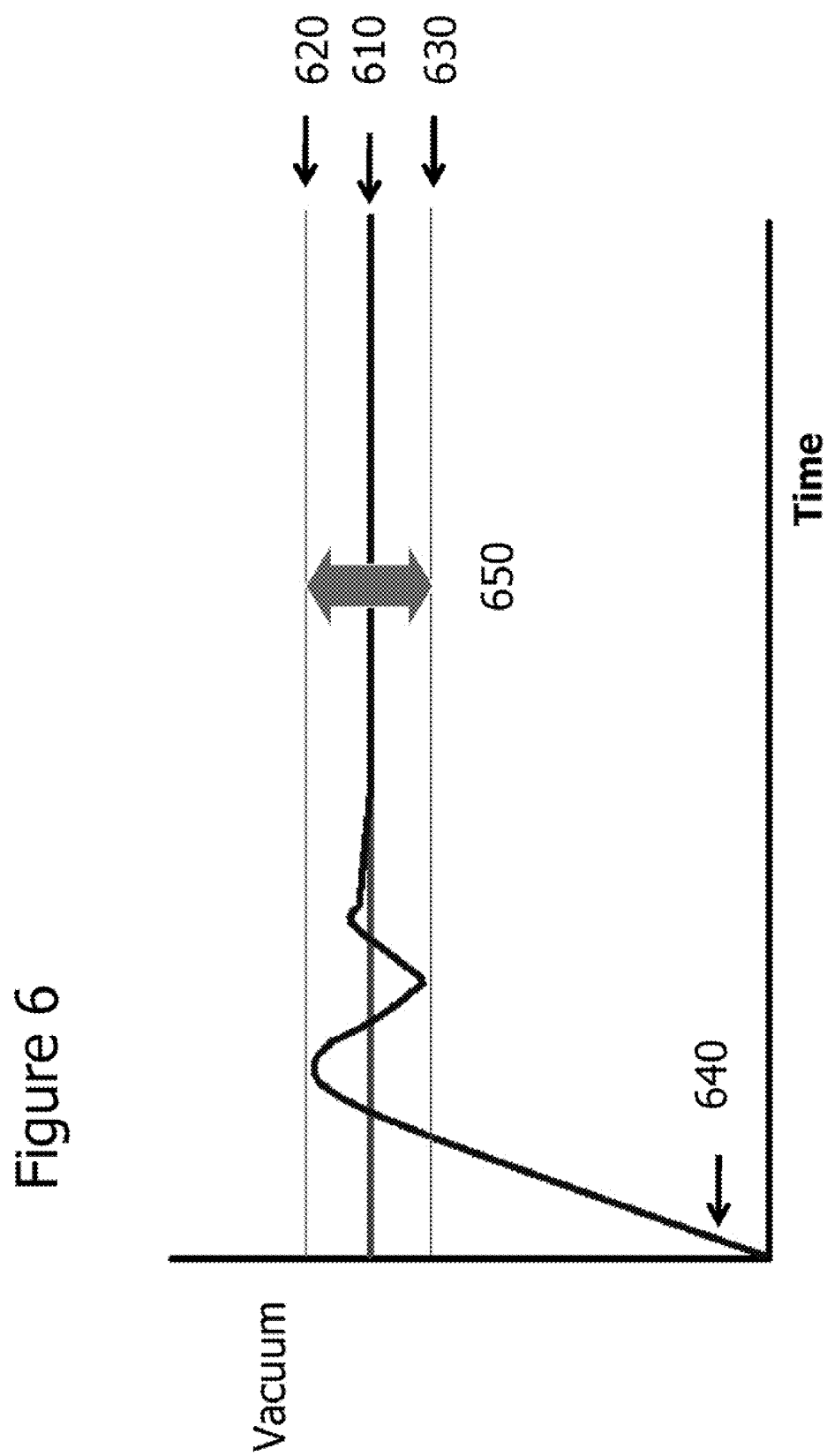


Fig. 5





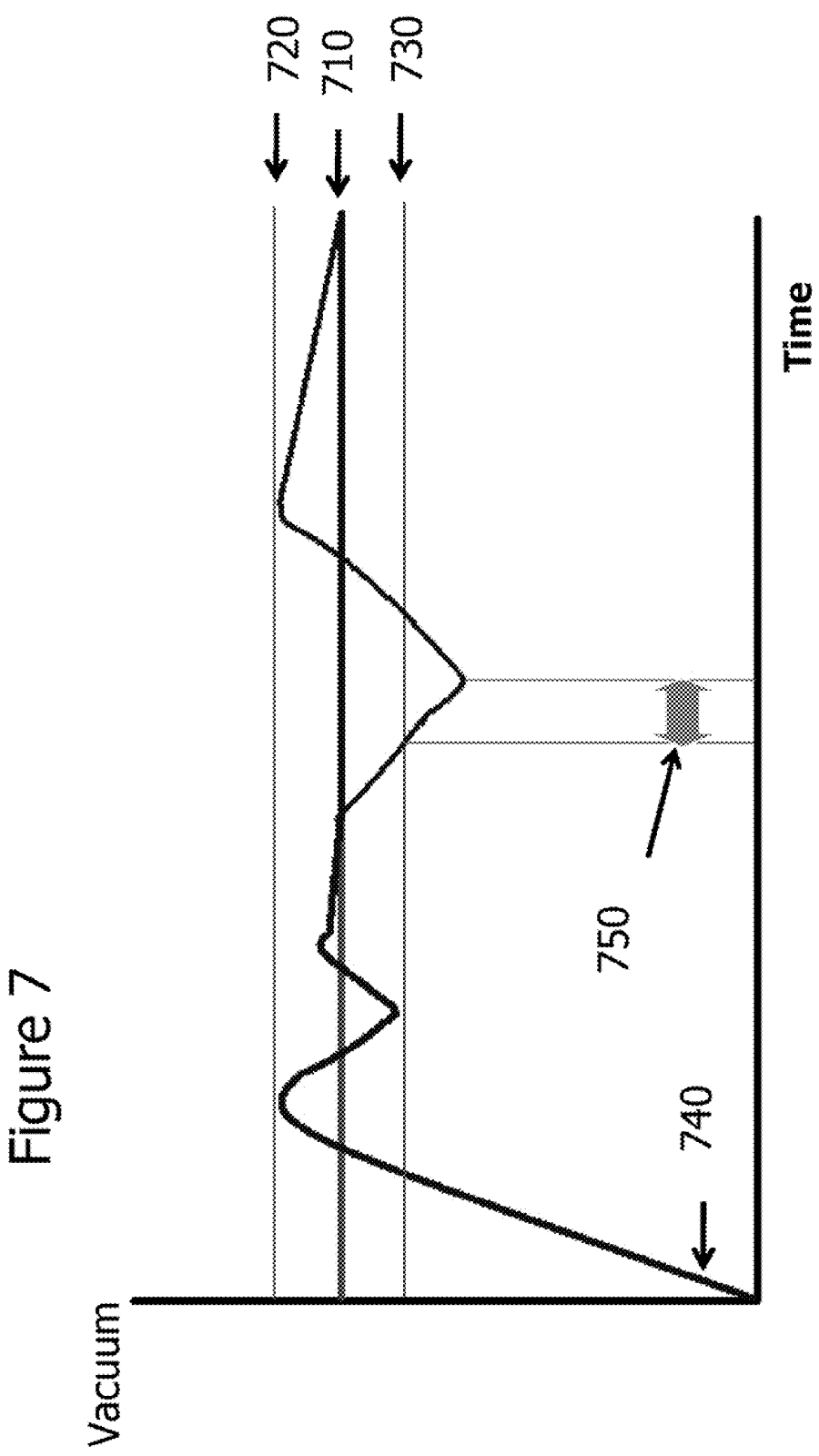


Fig. 8(A)

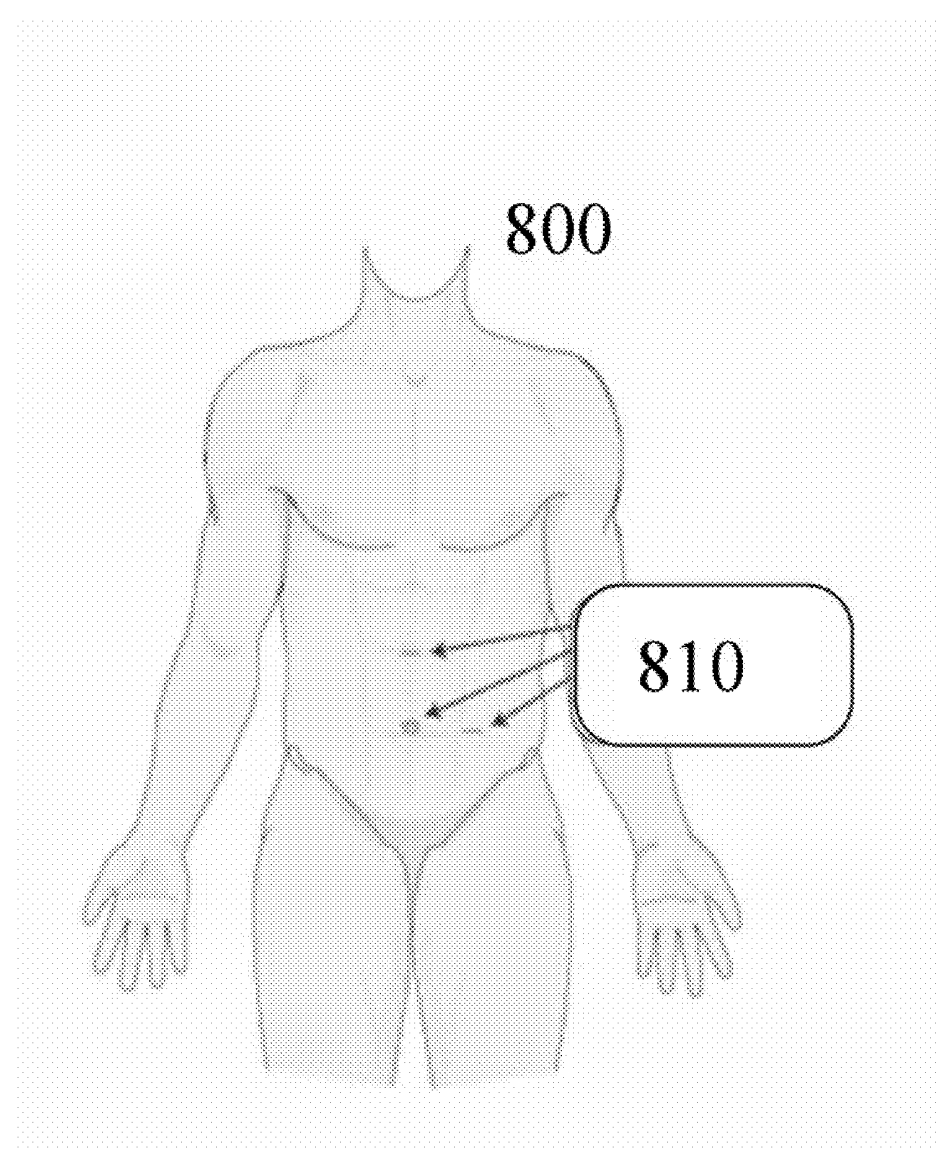


Fig. 8(B)

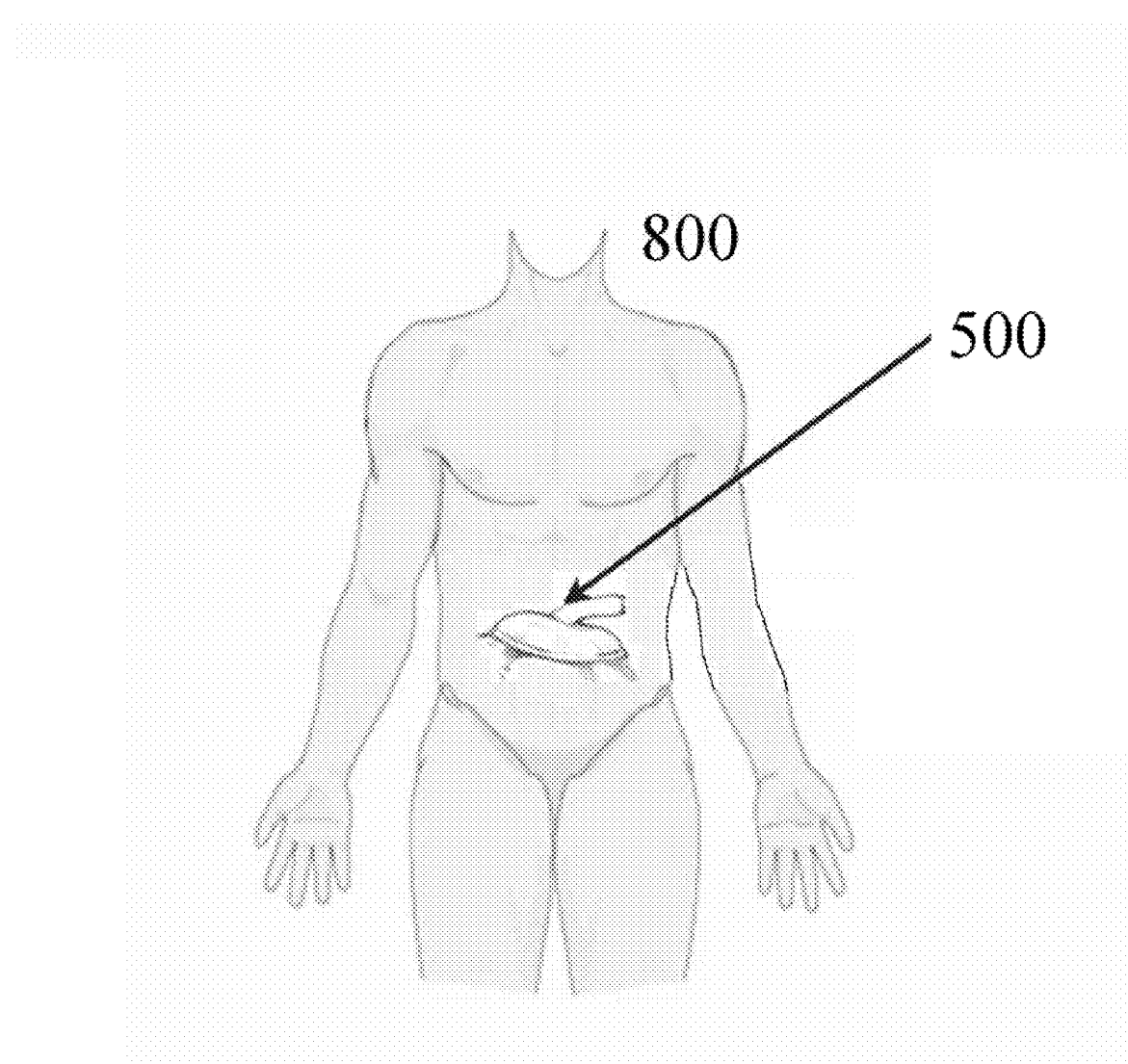
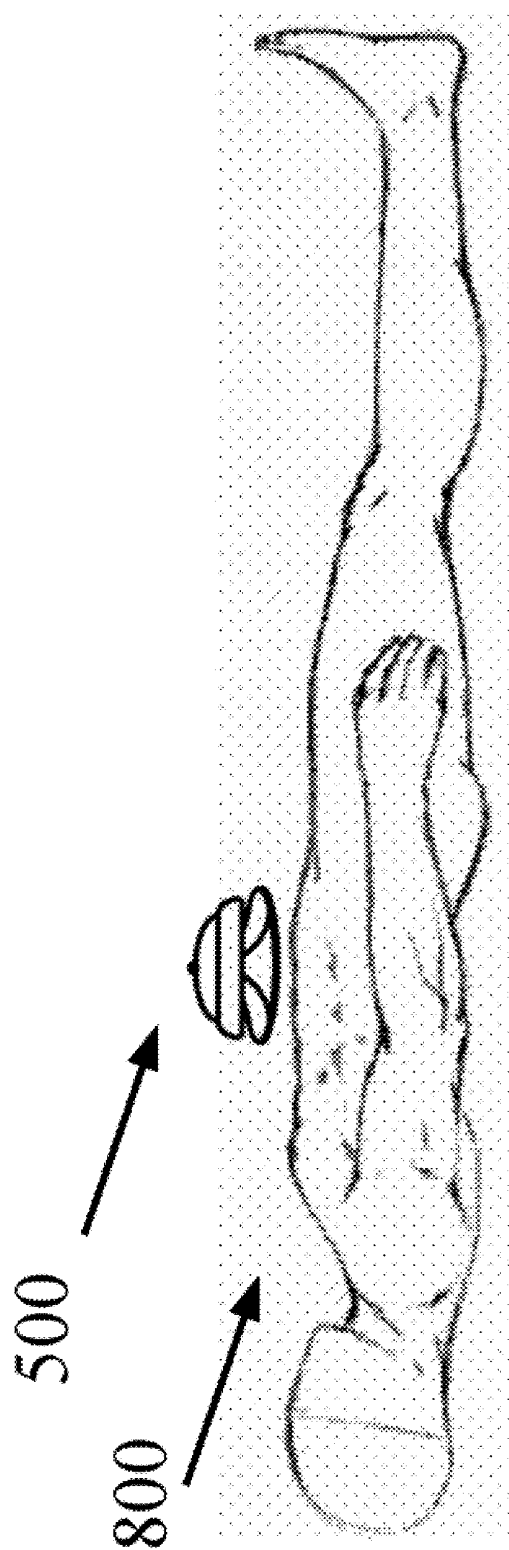


Fig. 8(C)



**PORTABLE REGULATED VACUUM PUMP
FOR MEDICAL PROCEDURES**

PRIORITY CLAIM

This application claims priority to: (1) U.S. Provisional Patent Application Ser. No. 61/146,429, entitled: "PORTABLE REGULATED VACUUM PUMP FOR MEDICAL PROCEDURES", inventors: A. Steve Gurganian and Neal M. Lonky, filed Jan. 22, 2009, which application is incorporated herein by reference in its entirety.

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is related to the following applications:

(2) "SURGICAL VACUUM INSTRUMENT FOR RETRACTING, EXTRACTING, AND MANIPULATING TISSUE" by Neal M. Lonky, application Ser. No. 09/489, 632, filed on Jan. 24, 2000, which issued on Nov. 4, 2003 as U.S. Pat. No. 6,641,575;

(3) "VACUUM INSTRUMENT FOR SLOWING OR ARRESTING THE FLOW OF BLOOD" by Neal M. Lonky, application Ser. No. 10/677,848, filed on Oct. 2, 2003; and

(4) "VACUUM INSTRUMENT FOR LAPAROTOMY PROCEDURES" by Neal M. Lonky, application Ser. No. 11/067,512, filed on Feb. 25, 2005, now abandoned, which applications (2)-(4) are explicitly incorporated herein by reference in their entireties.

FIELD OF THE INVENTION

This invention relates to vacuum assisted devices that require the use of a vacuum in order to attach to body parts and/or tissue.

BACKGROUND OF THE INVENTION

A grasping cup, obstetric bonnet, or ventouse cap are some of the names given to the devices used in the field of obstetrics. These devices use a suction cup to assist in the vaginal and operative delivery of newborns. Typically, the suction cup is applied to the fetal scalp during the second stage (pushing through the birth canal) of labor. The obstetrician then applies traction to the infant's head via a flexible or rigid shaft with a handle coupled to the suction cup. The traction is applied in an outward fashion while the delivering mother pushes, thereby assisting in the delivery of the newborn. These suction cups are typically made of silicone, rubber, vinyl or other plastic, or combinations of plastic and rubber. Suction is generally applied through suction tubing which is coupled to a nipple on the vacuum cup, the nipple communicating with the interior of the cup. The method by which the suction is produced can vary from large stationary mechanical vacuum/suction devices to hand-held pumps similar to those which are used to bleed brake fluid from brake lines of automobiles. The suction cup has been reported to be safer than forceps and is intended to more gently ease a baby from the womb than forceps.

Suction devices can also be used to manipulate tissue during surgery. For example, U.S. Pat. No. 5,196,003 to Bilweis discloses an endoscopic surgical instrument which includes a tube with a suction cup at one end and a bulb at its opposite end. The cup is placed on a target tissue and the bulb is compressed and released in order to apply suction to the tissue. The tissue is released by again compressing the bulb.

SUMMARY OF THE INVENTION

In an embodiment of the present invention, a pump is used to apply a vacuum to a vacuum device attached to a body part.

The vacuum device comprises the suction device (or suction cup), a sensor for monitoring the vacuum applied to the suction device, a pump for increasing the vacuum to the suction device, a tube for applying the vacuum to the suction device and a control circuit for activating the pump. In an embodiment of the present invention, the control circuit is operated by the surgeon holding the suction device. In an embodiment of the present invention, the surgeon controls the vacuum applied while performing the surgical procedure. In an embodiment of the present invention, the suction device can be manipulated and the vacuum device can be controlled by a surgeon alone while the surgical procedure is carried out.

BRIEF DESCRIPTION OF THE DRAWINGS

This invention is described with respect to specific embodiments thereof. Additional features can be appreciated from the Figures in which:

FIG. 1 is a flowchart showing the use of the vacuum device 301 in a procedure in accordance with an embodiment of the invention;

FIG. 2 illustrates the components of a circuit 209 and vacuum pump 214 for use in a vacuum device 301, in accordance with an embodiment of the invention;

FIG. 3 is a vacuum device 301 in accordance with an embodiment of the invention that can be used to implement the flowchart of FIG. 1;

FIG. 4 is a top view of a wireless control unit 419 with two buttons in accordance with an embodiment of the invention for controlling the vacuum device 301 and applying a vacuum to a suction device 435;

FIG. 5A is a disassembled integral suction and vacuum device 500 ready for assembly in accordance with an embodiment of the invention;

FIG. 5B is an assembled integral suction and vacuum device 500 in accordance with an embodiment of the invention;

FIG. 6 is a plot of vacuum vs time illustrating how vacuum hysteresis is minimized in accordance with an embodiment of the invention;

FIG. 7 is a plot of vacuum vs time illustrating how vacuum dwell time is minimized in accordance with an embodiment of the invention;

FIG. 8A is a frontal diagram showing a patient with a number of laceration bleeding sites;

FIG. 8B is a frontal diagram showing the utilization of an integral suction and vacuum device 500 on the patient shown in FIG. 8A in accordance with an embodiment of the invention; and

FIG. 8C is a cross section diagram showing the utilization of an integral suction and vacuum device 500 on a patient in accordance with an embodiment of the invention.

DETAILED DESCRIPTION OF THE INVENTION

In the following description, various aspects of the present invention will be described. However, it will be apparent to those skilled in the art that the present invention may be practiced with only some or all aspects of the present invention. For purposes of explanation, specific numbers, materials, and configurations are set forth in order to provide a thorough understanding of the present invention. However, it will be apparent to one skilled in the art that the present invention may be practiced without the specific details. In other instances, well-known features are omitted or simplified in order not to obscure the present invention.

Parts of the description will be presented in data processing terms, such as data, selection, retrieval, generation, and so forth, consistent with the manner commonly employed by those skilled in the art to convey the substance of their work to others skilled in the art. As is well understood by those skilled in the art, these quantities (data, selection, retrieval, generation) take the form of electrical, magnetic, or optical signals capable of being stored, transferred, combined, and otherwise manipulated through electrical, optical, and/or biological components of a processor and its subsystems.

Various operations will be described as multiple discrete steps in turn, in a manner that is most helpful in understanding the present invention; however, the order of description should not be construed as to imply that these operations are necessarily order dependent.

Various embodiments will be illustrated in terms of exemplary classes and/or objects in an object-oriented programming paradigm. It will be apparent to one skilled in the art that the present invention can be practiced using any number of different classes/objects, not merely those included here for illustrative purposes. Furthermore, it will also be apparent that the present invention is not limited to any particular software programming language or programming paradigm.

Systems and methods in accordance with embodiments of the present invention can provide for improved presentation and interaction with digital content and representations of digital content. Representation as used herein includes, but is not limited to, any visual and/or audible presentation of digital content. By way of a non-limiting example, digital images, web pages, digital documents, digital audio, and other suitable content can have corresponding representations of their underlying content. Moreover, interfaces such as graphical user interfaces can have corresponding representations of their underlying content.

The invention is illustrated by way of example and not by way of limitation in the figures of the accompanying drawings in which like references indicate similar elements. It should be noted that references to 'an' or 'one' embodiment in this disclosure are not necessarily to the same embodiment, and such references mean at least one.

The treatment of open wounds too large to spontaneously close have long been a troublesome area of medical practice. The initial stage of healing is characterized by the formation of granulation tissue which can take several hours. When an open wound is present or formed during surgery, it is not feasible to wait for normal wound healing. In order to help address this situation, application of vacuum has been proposed as a sealing device for the abnormal tissue. If a tissue displays a site which is bleeding, a suction cup may be applied at that site to tamponade active bleeding until the surgery can progress sufficiently to repair or remove the structure in question.

When a vacuum is applied to a suction cup on a structure having a blood flow, blood can typically pool inside the site, which may lead to the formation of a hematoma. When an obstetrical vacuum extractor is applied to a fetal scalp during vacuum extraction-assisted deliveries of newborns, for example, occasionally a small bruise or hematoma occurs on the scalp. Additionally, some soft tissue edema and swelling may occur on the area of the scalp where the vacuum was applied.

There is a significant need for vacuum-assisted devices in surgical and obstetrical delivery procedures. For example procedures that involve the use of vacuum cups, grasping cups, a tamponade apparatus, or suction cups that attach to body tissue require the use of external vacuum generation pumps.

Before describing the invention in detail, it is helpful to describe an example environment in which the invention may be implemented. The present invention is particularly useful in obstetrical delivery where the surgical procedure requires that a grasping cup or suction cup be attached to the fetal skull during the obstetrical delivery. Under these circumstances, monitoring of the suction cup can be partially obscured as the cup is attached in utero during the second stage of labor. The presence or absence of hair on the fetus scalp, the weight of the fetus and amount of fatty tissue on the scalp, the position of the fetus in the womb and the location that the suction device 435 is applied to the fetus scalp can all affect the effectiveness of this vacuum device 301 and the procedure for its use. In this manner, the vacuum device 301 can considerably assist the surgeon. The surgeon is able to diagnose on a continually unfolding basis the important features affecting the delivery procedure in a timely and accurate fashion and use this diagnosis to set appropriate vacuum levels for the vacuum device 301. Further, the ability of the surgeon to be able to make use of real-time changes to the delivery procedure in changing the vacuum levels applied to the vacuum device 301 to maintain the fixed vacuum to the suction device 435, can allow the surgeon to compensate for many of the variations related to these important features.

Surgical and obstetrical delivery procedures require suction devices 435 including vacuum cups, grasping cups, a tamponade apparatus, or other fixtures that attach to body tissue. Each of these devices requires a vacuum be applied to attach the device to tissue. The vacuum can be applied by a pump. Further, these procedures require continual monitoring, manipulation and re-activation of the pump by the physician. As such, the monitoring and manipulation create additional workload that distracts the physician from the primary surgical task.

Suction devices to manipulate tissue during surgery have exhibited various disadvantages. The suction devices are difficult to utilize in that the surgeon has very little control over the level of vacuum applied to the tissue. There is a check release valve on the hand-held pump, and the surgeon has the ability to re-pump up the vacuum. However, these tasks are difficult when the physician is concentrating on other facets of the surgical procedure. Importantly, the surgeon's ability to monitor the vacuum level that is applied through a suction cup and fine tune manipulations to increase or decrease vacuum pressure is very imprecise. Further, although the hand pumps can include a gauge, the physician does not have the time to visually monitor the gauge. More importantly, even if the physician had the ability to visually monitor the pressure of the gauge, the physician's manual reflexes would not allow the physician to react to the pressure fluctuations in time to avoid, for example, separation of the suction device. In an embodiment of the invention, the vacuum can prevent separation at the margin, i.e., at the suction device rim contact site. Further, releasing the vacuum on the tissue can only be carried out by releasing the check relief valve. Moreover, manipulation of the tissue using the suction device is limited by the direct application of force along the tube. In addition, the field of movement is not versatile, and may be inadequate for purposes of a given surgery. Additionally, applying the vacuum requires the assistance of a second medical professional.

A vacuum of atmospheric pressure is 1 atmosphere=760 torr. Generally, "approximately" in this pressure range encompasses a range of pressures from below 7.6×10^{-3} torr to 7.6×10^{-1} torr. A pressure of below 10^{-3} torr would constitute a high vacuum. A pressure of below 10^{-2} torr would constitute a medium vacuum. A pressure between atmospheric pressure

and a medium vacuum constitutes a low vacuum. In an embodiment of the invention a vacuum is sensed at a suction device 439. In an alternative embodiment of the invention a vacuum is sensed at a wound 510. In another embodiment of the invention a vacuum is sensed at a tissue. In an embodiment of the invention the vacuum sensed at the tissue compensates for out-gassing and leaks when the suction device is applied to the tissue. Generally, "approximately" in a low vacuum pressure range encompasses a range of pressures from between 5×10^{-1} torr to 5×10^{-2} torr. Generally, "approximately" in a medium vacuum pressure range encompasses a range of pressures from between 5×10^{-2} torr to 5×10^{-3} torr. In an embodiment of the invention, a marginal increase in vacuum is an increase of vacuum from 10^{-1} torr to 5×10^{-2} torr. In an alternative embodiment of the invention, a marginal increase in vacuum is an increase of vacuum from 5×10^{-2} torr to 10^{-2} torr. The vacuum device can operate to lower the pressure from atmospheric pressure to generate a high vacuum. In an embodiment of the invention, a low vacuum can be applied in for example obstetric suction devices. In an embodiment of the invention, a medium vacuum can be applied in for example orthopedic suction devices. In an embodiment of the invention, immediate release of a low vacuum reduces the vacuum to approximately atmospheric pressure in between a lower limit of approximately 0.1 second and an upper limit of approximately 0.5 second. In an embodiment of the invention, immediate release of a medium vacuum reduces the vacuum to approximately atmospheric pressure in between a lower limit of approximately 0.3 second and an upper limit of approximately 1.0 second. In an embodiment of the invention, a force of approximately 10^2 Newton can be applied to tissue with the vacuum device 301 through a suction device 439. In an alternative embodiment of the invention, a force of approximately 10^3 Newton can be applied to tissue with the vacuum device 301 through a suction device 439.

In various embodiments of the invention, the vacuum device can allow grasping of structures through a suction device during various types of surgery. Any organ or structure that has a smooth surface to which a vacuum device can attach and manipulate can be a candidate for use with the vacuum device. U.S. Pat. No. 6,641,575 "SURGICAL VACUUM INSTRUMENT FOR RETRACTING, EXTRACTING, AND MANIPULATING TISSUE" by Neal M. Lonky outlines an "open cup grasper" suction device 435 as envisaged in various embodiments of the invention attached to the vacuum device 301. The "open cup graspers" can be adapted for use with the vacuum device for open abdominal (laparotomy), chest (thoracotomy), brain (craniotomy) or orthopedic uses (vacuum traction or manipulation of bony surfaces). In addition, there is a more retroperitoneal surgical approach that can benefit from the vacuum device 301. Urology surgeons use an incision in the back to get to the kidneys. During this surgery the kidneys or other surfaces can be manipulated using the vacuum device.

Unexpectedly, it was found that application and maintenance of a fixed constant vacuum from a vacuum device prevented separation of the margin of the suction device from tissue when traction was applied or when patient movement jarred the margin of the device in the setting of tamponade of a bleeding site. In particular, it was found that the degree and range of motion, and amount of force which could be applied in traction, extraction, tamponade, and manipulation were all enhanced by the application of a constant vacuum.

In discussions with surgery assistants during procedures where only the hand pump was available to a number of surgeons, it became apparent that a mechanical pump led to

procedural failures due to surgeon fatigue. The effort required in constantly monitoring the pressure gauge and reinitiating the mechanical pumping by grasping and compressing the handle exacerbated psychological pressure and physical fatigue due to the surgical procedure. In experiments with an animal model or simulations with human tissue it was observed that "pop off" detachments were significantly minimized with the automated vacuum device 301. This result was unexpected based on the informal questioning of a group of five surgeons who were shown a prototype vacuum device 301 in these model or simulation environments. This result was also unexpected inasmuch as all surgeons consulted indicated that they did not know marginal separation was a factor that could be addressed by a constant vacuum pump until shown the prototype and/or the results.

If traction force exceeds the pressure under the suction device (the attachment force), the suction device edge furthest from the fraction point will lift and separate from the tissue, causing the leakage leading to cup "pop off". Essentially, when that equilibrium is reached and the two pressures equilibrate, there is a slow leak. It was learned after consulting with surgical assistants that the surgeon does not sense this release and thereby cannot manually pump the hand-held vacuum pump to counteract the leakage in time before the suction device releases completely. Unexpectedly, it was found that a machine-driven pump sensor can detect the release as measured by the drop in vacuum, and compensate for the release by activating the automated pump and compensating for the partial release, thereby allowing application of greater traction force without experiencing "pop off".

When a suction device 435 is applied to a fetal scalp on which lanugo, vellus or terminal hair is present, the hair can disturb the seal between the suction device and the scalp and thereby result in a vacuum leak. Application of a constant vacuum can compensate for the vacuum leak allowing for traction, extraction, and manipulation of the fetal head. Once the head has exited the vagina the surgeon needs to be able to release the suction device and manually deliver the fetus. Complete venting of the relief valve 221 via a solenoid allows instantaneous release of the suction device 435. When a suction device is applied to an organ containing an uneven or obstructed tissue surface, the uneven surface can result in a vacuum leak. Application of a constant vacuum can compensate for the vacuum leak allowing for traction, extraction, and manipulation of the organ.

Application of a suction device at a tissue site which is bleeding, with a constant vacuum, i.e., an applied vacuum that compensates for the changed clotting behavior of the wound with time, can allow enhanced clotting and cessation of the bleeding from the site. It could be expected that a vacuum device with a vacuum tube delivering a vacuum to a suction device applied to the wound would accelerate the flow of blood. Contrary to this preconceived idea, application of vacuum constricts the tissue at the margin of the bleeding site into/toward the bleeding site. As a result, the margin collapses, allowing pooling of blood and clotting factors aiding clotting and cessation of blood flow.

In an embodiment of the invention, the timing and periodicity of the application of the vacuum can be applied in a pulsatile flow opposite that of a patient's pulse. In an embodiment of the invention, the vacuum can be applied to tissue. The tissue can include, epithelial layers, organs, osseous tissue, bony tissue and connective tissue exposed by an open incision during a laparotomy, thoracotomy, craniotomy, retroperitoneal surgery and orthopedic vacuum traction or manipulation of bony surfaces. In an embodiment of the invention, the vacuum can be applied to a wound when the

surgeon senses diastoly to more actively collapse the wound. The wound can be one or more of a subcutaneous hematoma, an incision made by the surgeon, a site of removal of tissue by the surgeon, a site of removal of an organ made by the surgeon, one or more puncture holes being treated by the surgeon or one or more laceration sites being treated by the surgeon. Treating the wound can include stopping the blood flow from a site, reducing the blood from a site, assisting in the arrest of the flow of blood to a site, stopping the blood flow to a subcutaneous hematoma, reducing the blood flow to a subcutaneous hematoma, assisting in the arrest of the flow of blood to a subcutaneous hematoma limiting its size and scope. In an embodiment of the invention, the vacuum can be increased to the wound when the surgeon senses diastoly to more actively collapse the wound. In effect the surgeon can use the patient's own tissue and the suction device as a tourniquet and the vacuum can be increased to tighten the tourniquet. In an embodiment of the invention, marginally increased vacuum pressure can be applied in diastoly, and thereby more easily applied, and then the vacuum can be maintained to overcome the systolic pressure pulse, thus slowing or arresting the flow of blood.

In an embodiment of the invention, the vacuum can be adjusted to compensate for the pulse pressure variation, maintaining or slightly exceeding the suction device vacuum against the higher systolic pressure to secure placement at or over the bleeding site. Maintenance of the suction device on the bleeding site will ultimately more actively collapse the wound. In various embodiments of the invention, the vacuum device 301 can increase or decrease the vacuum to the wound in synchronization with a patient's heart changing from diastole to systole to more actively collapse the wound. In this manner, the clotting and cessation of blood flow of a wound can be further improved. It was unexpected that vacuum constriction applied when the patient's blood pressure is low could reduce bleeding. It was also unexpected that the ability of a surgeon to simultaneously perform surgery and monitor the pulse pressure would be near impossible, which was observed when we interviewed 5 surgeons in a simulation of a bleeding site event.

Further, it is noted that undetected vacuum loss may result in the untoward premature detachment of the vacuum cup device from the body surface under manipulation. Improper or negligent operation of the vacuum pump, or the manipulation of the tissue in contact with the various procedure cups/devices, may hamper the medical procedure or damage the affected tissue.

In the past, vacuum assisted devices or vacuum generating devices have not recorded or maintained a record of operation of parameters including the applied vacuum, vacuum leak being compensated, time of application, temperature, change in conditions during procedure, or notes regarding the classification of the surface and the nature of the problem addressed that can be reviewed and archived for inclusion in medical records. As such, these vacuum assist devices do not allow the physician to utilize insight and intuition to choose conditions for the procedure that are similar to conditions previously utilized. Finally, these vacuum assist devices do not allow the surgeon to select conditions based not only on the similarity of the procedure but also based on the surgeon performing the procedure.

Post-surgical examination of the parameters used during a number of surgical procedures including vacuum applied, nature of tissue, evenness of surface, vacuum leak due to uneven surface, force of extraction/manipulation applied, time duration of vacuum applied and variation of vacuum applied, allow review and establishment of guidelines that

advance the safety and efficacy of the various surgical procedures being undertaken for a given patient sub-population. By recording all the data obtained during surgery pertaining to the use of the vacuum device 301 and making this data available together with other details of the patient and the operating surgeon, a surgeon can store useful data and proactively select parameters for future use in similar circumstances based on stored data. For example, ultrasound analysis of the fetus prior to birth can be performed to determine how much hair is present on the fetal scalp. This information can be used to select appropriate conditions for use in the childbirth, should a suction device be required. In addition to the absence or presence of hair and the relative amount of hair, the surgeon can also select the size and type of the specific suction device 435, and the specific type of vacuum device 301, based on the performance characteristics of the specific suction device 435, and the specific type of vacuum device 301. During specific surgical procedures a variable vacuum can be desirable, where the vacuum applied to the suction device 435 is varied depending on the stage of the surgical procedure. In such circumstances, the surgeon can manually indicate the different stages of the surgical procedure through a single depression of a control unit 419 button or through voice-activated recognition and response by the vacuum device 301. In an alternative embodiment of the invention, the different stages of the surgical procedure can be communicated to or noted by an assistant who enters the appropriate information in the vacuum device 301. In an embodiment of the invention, the control unit 419 is attached to the suction device 439. In an embodiment of the invention, the control unit 419 can be held by the surgeon while holding and controlling the suction device 439. In an embodiment of the invention, the control unit 419 can be held by the surgeon with the same hand holding and controlling the suction device 439.

In various embodiments of the present invention, an electromechanical compact pump provides a sustained vacuum force to a variety of suction devices 435 applied to body tissue and surfaces for the purpose of grasping, manipulating, tamponade, and/or traction of body parts of a patient or a fetal skull during obstetrical delivery. In an embodiment of the present invention, the vacuum can be automatically regulated and monitored continuously by the vacuum device 301 to effect the uninterrupted generation of a constant vacuum pressure to prevent untoward release of the suction device from the affected body part or tissue. In an embodiment of the present invention, the vacuum can be automatically regulated and monitored continuously by the vacuum device 301 to effect the uninterrupted generation of a variable vacuum pressure to prevent untoward release of the suction device 435 from the affected body part or tissue. In an embodiment of the present invention, the desired vacuum provided by the vacuum device 301 can be preset by the operator and can be adjusted by the surgeon as necessary during a medical procedure. In an embodiment of the present invention, an electronic record of the vacuum pressure(s) vacuum leak, and time duration of the vacuum can be stored internally within the device. In an embodiment of the present invention, the electronic records can be retrieved by an accessory memory transfer device or transmitted by wired or wireless (RF/IR frequency) connection to an external computer or other receiving equipment (i.e. printer, mass storage device). In an embodiment of the present invention, the vacuum device 301 can operate as a primary vacuum pump. In an embodiment of the present invention, the vacuum device 301 can assist continuously or during designated times in parallel with existing hand-operated vacuum pumps. In an embodiment of the

present invention, the vacuum device **301** can operate as a back-up vacuum pump in parallel with existing hand-operated vacuum pumps.

In various embodiments of the present invention, improved and unique operation is provided when compared to non-regulated vacuum pumps. In an embodiment of the present invention, the vacuum device **301** includes an electromechanical pump (electro-vacuum pump). In an embodiment of the present invention, the vacuum device **301** is hand held. In an embodiment of the present invention, the vacuum device **301** is portable. An electro-vacuum pump for assisting or regulating the vacuum obtained by hand-operated pumps has not previously been proposed or used. In an embodiment of the present invention, the vacuum device **301** improves performance of hand-operated pumps currently in use by providing sustained and regulated vacuum. In an embodiment of the invention, the surgeon selects the desired vacuum by applying the vacuum with a hand-held pump and the vacuum device **301** is used to maintain this pressure. In an embodiment of the present invention, the vacuum regulation provided by the invention prevents untoward release ("pop-offs") of grasping cups and other suction devices **435** by automatically compensating for air leaks at the interface between the grasping cup and the adhered surface, or between the grasping cup and the vacuum device **301**.

In various embodiments of the invention, the portable regulated vacuum device **301** shown in FIG. 3 is a self-contained vacuum-producing device that provides preset regulated pressure(s) and electronically stores operational performance data for retrieval by external means. The circuit for the vacuum device is shown schematically in FIG. 2. The power switch **202** activates the vacuum device **301** for operation utilizing the energy storage device **215**. In an embodiment of the present invention, the power switch **202** remains indefinitely in whichever state it is placed. In an embodiment of the invention, the vacuum device is activated by contact between the suction device and tissue. In an embodiment of the present invention, the vacuum selector switch **203** is a multiple position on-off switch that remains indefinitely at whichever position is selected. In an embodiment of the present invention, the vacuum selector switch **203** can provide three (3) positions. In an embodiment of the present invention, the vacuum selector switch **203** can provide ten (10) positions. In an embodiment of the present invention, the vacuum selector switch **203** can provide any number of positions between a minimum of three (3) positions and a maximum of ten (10) positions. In an embodiment of the present invention, the vacuum selector switch **203** can provide two (2) positions. In an embodiment of the present invention, the vacuum selector switch **203** can provide more than ten (10) positions. In an embodiment of the invention, the vacuum selector switch **203** can be used to select a "maintain" vacuum setting, wherein a vacuum generated by a hand-held vacuum pump is maintained by the electro-vacuum pump during the surgical procedure. In an embodiment of the present invention, the vacuum selector switch **203** position can be selected based on the surgical procedure being undertaken. In an embodiment of the present invention, the vacuum selector switch **203** position can be selected based on the identity of the surgeon performing the surgical procedure.

FIGS. 2-4 depict objects as logically separate. Such depiction is merely for illustrative purposes. It will be apparent to those skilled in the art that the objects portrayed in FIGS. 2-4 can be arbitrarily combined or divided into separate software, firmware or hardware components. Furthermore, it will also be apparent to those skilled in the art that such objects, regardless of how they are combined or divided, can execute on the

same device or can be distributed among different devices connected by one or more connections.

In an embodiment of the present invention, the operator display and control **204** can be comprised of an alpha-numeric display panel and associated push-button controls to effect the change and selection of displayed captions and data. In various embodiments of the present invention, the display panel can be an LCD (liquid crystal display), LED (light emitting diode), electro-luminescent or any other component used to provide visual text or graphical representation of the vacuum device's **301** operational condition, current or stored in memory. In an embodiment of the present invention, the push-button controls can be typical momentary switches such as membrane, snap-dome, or any other mechanically actuated switch. In an embodiment of the present invention, the operator display and control **204** are attached to the device circuit board **209** and are interfaced to the device processor circuitry **213** for control.

In an embodiment of the present invention, the LED bar-graph indicator **205** can be comprised of multi-color LEDs or any other illuminating display device(s) that provide a sequential lighted indicator. In various embodiments of the present invention, any number of indicators can be used. In an embodiment of the invention, ten indicators can be used. In an embodiment of the invention, the indicators can use green, yellow and red colors to indicate a displayed parameter. In an embodiment of the invention, green indicators can be used at the bottom extending past the middle followed by yellow, with the red color at the top section of a vertically arranged stack of indicators.

In an embodiment of the present invention, the vacuum attachment port **206** can be a barbed hose stem made of any resilient material such as metal or plastic, or other flexible or rigid polymer. In various embodiments of the present invention, the length and diameter of the vacuum attachment port **206** can be any size. In an embodiment of the invention, the vacuum attachment port **206** extends approximately 5 mm in length from the vacuum device **301** and is 2 mm in diameter. In an alternative embodiment of the invention, the vacuum attachment port **206** extends approximately 25 mm in length from the vacuum device **301** and is 6 mm in diameter. In an embodiment of the invention, an external suction device **435** can be connected to the vacuum device **301** at the vacuum attachment port **206** location via the vacuum assisted device tubing **308**.

In an embodiment of the present invention, the external memory and interface port **207** can be an electrical connector used for small computer communication interfaces. The external memory and interface port **207** can be an RS-232, RS-422/485, Ethernet or Universal Serial Bus (USB) connector. The interface port **207** can provide for the wired connection of the vacuum device **301** to external peripherals including personal computers, printers or other portable computing devices such as personal digital assistants (PDA), mass storage devices, flash memory devices (thumb drives) or digital telecom devices such as cellular telephones.

In an embodiment of the present invention, the vacuum assisted device tubing **308** can be a section of flexible tubing of surgical quality manufactured of any typical material such as PVC, rubber (natural or synthetic), silicone or others. The tubing **308** can be any length or diameter necessary to provide for the inter-connection of external vacuum assisted devices to the vacuum attachment port **206**. As shown in FIG. 4, the vacuum assisted device tubing **308** can have identical or differing internal or outer diameters (ID and OD) that facilitate connection to a suction device **435** or other target medical device that requires vacuum pressure for operation.

In an embodiment of the present invention, the device circuit board 209 holds the electronic components for the vacuum device 301 system control. The operator display and control 204, LED bargraph indicator 205, external memory and interface port 207, pump control circuitry 210, vacuum sensor 211, wireless communications circuitry, and the device processor circuitry 213 can be attached to the circuit board 209. The device circuit board 209 can be externally powered by the energy storage device 215.

In an embodiment of the present invention, the pump control circuitry 210 can be comprised of typical electronic components to effect the switching of power to the vacuum pump 214. These components include, but are not limited to: electromechanical relays, transistors, MOSFETS, diodes, or other devices capable of controlling the electrical current necessary to power the vacuum pump 214. The pump control circuitry 210 can be controlled through an interface with the device processor circuitry 213.

In an embodiment of the present invention, the vacuum sensor 211 can be a semiconductor device utilizing typical variable resistance or capacitance technologies that provides a proportional voltage signal output based upon the vacuum pressure applied to the sensor. The vacuum sensor 211 voltage signals can be interfaced to the device processor circuitry 213 to provide real-time vacuum pressure readings necessary to regulate the vacuum pump 214 operation.

In an embodiment of the present invention, the wireless communications circuitry 212 can provide modulated waveform signals in either the infrared (IR) or radio frequency (RF) signal range. In an embodiment of the invention, the wireless communications circuitry 212 can provide RF inputs/outputs. The modulated signals can be transmitted using any typical small computer Ethernet system WiFi standard such as EIA-802.11G, Bluetooth® methodology or discrete signaling utilizing a non-linear code encryption algorithm for secure control. The wireless communications circuitry 212 can be controlled through an interface with the device processor circuitry 213 to provide remote control and data transfer between the vacuum device 301 and peripheral devices as: remote control units (pushbuttons, foot switches), personal computers, printers or other portable computing devices such as personal digital assistants (PDA), mass storage devices, or digital telecom devices such as cellular telephones.

In an embodiment of the present invention, the device processor circuitry 213 can be comprised of typical micro-processor electronic components necessary to provide pre-programmed and user-selected operation of the portable regulated vacuum device 301. In various embodiments of the present invention, components of the processor circuitry 213 include but are not limited to: volatile and non-volatile memory, real-time clocking, and peripheral interface and logic devices. In an embodiment of the present invention, an internal system processor firmware program can provide the direction to operate the various circuit blocks that reside on the device circuit board 209. Functionality provided by the firmware can include:

a) comparison of desired and actual vacuum pressure supplied by the vacuum device 301 to the user-selected suction device 435. In an embodiment of the present invention, a control algorithm can be used to control hysteresis (dead-band, see FIG. 6) and dwell (delay timing, see FIG. 7) and thereby minimize or eliminate oscillatory cycling of the vacuum pump 214 that: minimizes pump run duration time, maintains vacuum pressure within a desired range, and minimizes power consumption;

b) activation of the pump control circuitry 207 to control the vacuum pump 214 to reach and maintain the user-selected vacuum pressure;

c) logic output signal control to effect a proportional indication of the vacuum sensor 211 signal at the LED bargraph indicator 205;

d) interface logic for the operator display and controls 204 to provide user adjustments of the vacuum device 301 and to provide status messaging and data during operation;

e) control of the wireless communication circuitry 212 to implement the properly formatted exchange of data with other peripheral devices; and

f) control of the external memory and interface port 207 to provide the proper transfer of stored histogram data to external memory storage devices, and the bi-directional communication required to effect remote control of the vacuum device 301 using an external host computing device.

In an embodiment of the present invention, the vacuum pump 214 can be a miniature pump utilizing any typical pressure building chamber-type mechanism such as diaphragm, bellows, piston or WOB-L® technology. FIG. 2 shows an embodiment of the invention where the vacuum pump 214 includes the pump motor 223 connected to a pump head 224. In an embodiment of the present invention, the pump motor can be a rotational electromechanical type. In an alternative embodiment of the present invention, the pump motor can be of a reciprocating electro-mechanical type. In an embodiment of the present invention, the pump motor can use an alternating current (AC) electrical power source. In an alternative embodiment of the present invention, the pump motor can use a direct current (DC) electrical power source.

In an embodiment of the present invention, the energy storage device 215 can be a DC energy or charge storage device that is configured to provide power to the device circuit board 209. The energy storage device 215 can include any battery or cell known in the field, including general purpose batteries, alkaline batteries, lithium ion batteries, nickel-cadmium batteries, nickel metal hydride batteries, lead acid batteries, deep cycle batteries, rechargeable batteries, or any other batteries. In an embodiment of the invention, the voltage or energy content of the energy storage device 215 is in the range from a minimum voltage of approximately 6 volts to a maximum voltage of approximately 24 volts. In an embodiment of the invention, the voltage or energy content of the energy storage device 215 is in the range from a minimum voltage of approximately 9 volts to a maximum voltage of approximately 18 volts. In an embodiment of the invention, the voltage or energy content of the energy storage device 215 is approximately 12 volts.

In an embodiment of the present invention, the flexible tubing 216 can be flexible tubing of surgical quality manufactured of any typical material such as PVC, rubber (natural or synthetic), silicone or others. The flexible tubing 216 can be sized to provide an airtight connection between the vacuum pump 214, the vacuum sensor 211, the check valve 220, the solenoid relief valve 221 and the vacuum attachment port 206.

In an embodiment of the present invention, the wiring 217 can comprise a current-conducting material that electrically connects the components of the vacuum device 301. The wiring 217 can include any material or wiring known in the art, including but not limited to copper and other metals having low electrical resistance. The wire gauge may be chosen to reduce costs while simultaneously carrying current with a minimum of resistant power dissipation.

In an embodiment of the present invention, a foot switch 218 can be used with the vacuum device 301. In an embodiment,

ment of the present invention, the foot switch 218 can be a push-on/push-off latching switch that remains in the last selected position indefinitely. The foot switch 218 can be electrically connected to the vacuum device 301 with a wired tether or through wireless means utilizing the wireless communications circuitry 212. The foot switch 218 can allow the vacuum pump 214 of the vacuum device 301 to be operated remotely. The electrical switch mechanism is typically a mechanically actuated push-button switch; when in the 'on' condition, the vacuum pump 214 is allowed to operate. Conversely, when in the 'off' condition the operation of the vacuum pump 214 is disabled.

In an embodiment of the present invention, the wireless control unit 419 can be a small form factor self-contained RF transmitter with an integral storage battery power source. In an embodiment of the present invention, the wireless control unit 419 can be wirelessly connected to the vacuum device 301 utilizing the wireless communications circuitry 212 with discrete signaling utilizing a non-linear code encryption algorithm for secure control. In an embodiment of the present invention, the wireless control unit 419 control mechanism can be comprised of two (2) or more mechanically actuated push-button switches with tactile surfaces allowing the operator to differentiate between the buttons and thereby allowing each button to be identified without visual verification. Tactile differentiation can be accomplished by unique button size, shape, embossed or raised symbols, or any combination of these methods. The wireless control unit 419 can allow the vacuum device 301 to be operated remotely. Through the actuation of a single or predefined combination of buttons, the operation of the vacuum device 301 can be controlled. Control functions include: vacuum pump 214 on/off, release of vacuum, real-time adjustment of regulated vacuum operating pressure and adjustment of other operating parameters as afforded by the device processor circuitry 213.

The check valve 220 can be an in-line, 2-port miniature air valve utilizing swing disk diaphragm, ball, or other seating methods typical in the art. The check valve 220 allows air flow in only one direction, toward the vacuum pump 214. Disk or ball check valves may be free moving or utilize levers or springs to assist in seating faster to eliminate air flow shock and/or inhibit air flow based upon applied vacuum. The check valve 220 can be positioned to eliminate vacuum leakage through the vacuum pump 214 when the vacuum pump 214 is not operating.

The solenoid relief valve 221 can be an electro-mechanically operated 2-port valve utilizing moving plunger or plate technology to open and close an air passage between the two air ports. Plunger or plate valve actuators can be driven by a magnetic field created when electrical current is applied to a valve coil. The solenoid coil can be AC or DC current types as typical in the art. The valve can be held in a normally-closed condition, inhibiting air flow at and through each port. Upon electrical actuation by the pump control circuitry 210, the valve can be opened allowing vacuum pressure present at vacuum attachment port 206 to be vented to the atmosphere, releasing vacuum holding pressure at the external vacuum assisted device.

FIG. 2 shows the pressure chamber of vacuum pump 214 connected through the check valve 220 to both the vacuum attachment port 206 and the solenoid relief valve 221, then to the vacuum sensor 211 through the use of the flexible tubing 216. The interconnection of these items can also be accomplished with other methods such as face to face port connections using threaded fasteners or all items can be integrated into the pump chamber housing forming a single unit requiring no external connections.

In an embodiment of the present invention, the circuitry sections on the device circuit board 209: pump control 210, vacuum sensor 211, external memory and interface 207, wireless communication 212, device processor, operator display and controls 204 and the LED bargraph indicator 205 can all be soldered in place and interconnected as required with the printed wiring that forms a part of the device circuit board 209. In an embodiment of the present invention, the vacuum pump 214 and the energy storage device 215 can be wired 217 to the device circuit board 209. In an alternative embodiment of the present invention, the vacuum pump 214 and the energy storage device 215 can be connected and soldered directly to the device circuit board 209.

In an embodiment of the present invention, the power switch 202 and vacuum selector switch 203 can be connected to the device circuit board 209. In an alternative embodiment of the present invention, the power switch 202 and vacuum selector switch 203 can be relocated with alternatively wired 217 connections.

In an embodiment of the present invention, the foot switch 218 can be wired 217 to the vacuum device 301 with a cable of a suitable length to position the foot switch 218 near the operator. In an alternative embodiment of the present invention, the foot switch 218 can be used wirelessly through an RF link with the wireless communications 212 circuitry to accomplish the same functionality afforded by a wired 217 connection with the device circuit board 209.

FIG. 1 is a schematic flow diagram used to exemplify the use of an embodiment of the present invention in a surgical procedure. The vacuum device 301 is initially in an 'off' state, with no power applied to the internal components. Prior to operation, the user attaches the suction device 435 via connector 425 (see FIG. 4) to the vacuum attachment port 206 of the vacuum device 301 using the vacuum assisted device tubing 308 at step 110. The suction device 435 is one of one or more devices that require vacuum assistance. Then the user applies power to the vacuum device 301 with power switch 202 at step 100 and positions selector switch 203 to the desired operating vacuum setting at step 105.

When enabled with the foot switch 218 or the suction device control 419 (at step 115), the device processor circuitry 213 determines if the vacuum pump 214 is required to operate, or not, based upon the relationship between the user setting of the vacuum selector switch 203 and the signal magnitude of the vacuum sensor 211 (at steps 120 and 130). Once activated the vacuum device begins to monitor and store the data including the vacuum applied to the suction device, the vacuum at the vacuum pump, the vacuum loss (not shown in FIG. 1). If the vacuum sensor signal is lower than the desired vacuum, the pump control circuitry 210 is activated powering the vacuum pump 214 until the desired vacuum is reached (steps 140 and 160); as determined by the signal from the vacuum sensor 211 and the device processor circuitry 213. The relative vacuum pressure is annunciated by the LED bargraph indicator 205, with the value of the vacuum sensor 211 signal proportionately indicated by sequential LED indicator illumination from the bottom of the bargraph to the top. If the vacuum sensor signal is higher than the desired vacuum, the pump control circuitry 210 is not activated and the relief valve 221 is opened (step 150) to reduce the vacuum. If the vacuum sensor signal is within the limits the vacuum sensor continues to monitor the pressure (step 120) until the power switch 202 is turned 'off' (steps 170 and 180).

During the operation of the vacuum device 301, vacuum pressure data can be stored in non-volatile memory that forms a part of the device processor circuitry 213. The vacuum pressure data can be stored sequentially with associated real-

time (time of day) values that provide a direct correlation of the operating vacuum at a given time. The stored memory data can be retrieved by the user. In various embodiments of the invention, the stored memory data can be retrieved by:

- (a) negotiation and data download to an external device using the wireless communication circuitry 212;
- (b) insertion of a non-volatile memory device (flash/thumb drive) at the external memory and interface port 207; or
- (c) by request (serial communication) of a host computer connected to the external memory and interface port 207.

With the operator display and controls 204, the user may perform various functions that affect the operation of the vacuum device 301; these functions include, but are not limited to:

- (i) adjust dead-band (+/- error range of vacuum signal when compared to desired setting) and dwell (delay time before/after allowable error is exceeded to activate/de-activate the pump) parameters;
- (ii) adjust preset vacuum values associated with the user vacuum selector switch 203;
- (iii) monitor vacuum pressure in real-time with display provided in selectable engineering units including mm/Hg, bar, mbar, torr, mtorr, PSI and kPa;
- (iv) retrieve and view stored time-stamped vacuum histogram values; and
- (v) adjust other system parameters associated with the functionality of the LED bargraph indicator 205 and communication protocols used by the external memory and interface port 207 and the wireless communication circuitry 212.

In an embodiment of the present invention, the vacuum selector switch 203 can be expanded in functionality to include: more selections (positions), factory pre-set operational values. In an alternative embodiment of the present invention, the vacuum selector switch 203 can be eliminated completely and replaced by a single pre-set or adjustable setting through the use of the operator display and controls 204. In another embodiment of the present invention, the vacuum selector switch 203 can be set according to the surgical procedure being undertaken. In an embodiment of the present invention, the minimum and maximum pressure which can be selected during the surgical procedure can be based on the surgical procedure being undertaken. In an embodiment of the present invention, the minimum and maximum pressure which can be selected can be based on the stored identity of a surgeon carrying out the procedure. The identity of the surgeon carrying out the procedure can be detected through a RFID reader in the vacuum device 301 and a tag associated with the surgeon. In an embodiment of the present invention, the minimum and maximum pressure which can be selected during the surgical procedure can be restricted based on the surgical procedure being undertaken.

The functionality of the LED bargraph indicator 205 can be expanded to allow user selected indications that provide: 0-100% indication of the vacuum sensor 211 signal, scaled to provide 0-100% indication of the vacuum selector switch pre-set range, or other user selected range of indication.

Attachment of the external vacuum assisted device to the vacuum attachment port 206 can be accomplished using custom configured vacuum assisted device tubing 308 sections that are pre-fabricated and sized to provide the simple interface of devices currently used in the art.

In an embodiment of the present invention, the vacuum device 301 can be reconfigured to utilize simple electronic circuit components eliminating the need for the device processor circuitry 213. Electronic circuitry that provides an analog of the processor control algorithm, which performs the

hysteresis and dwell functions, can be used to provide simple dedicated operation of the portable regulated vacuum device 301.

In various embodiment of the present invention, the unique operational characteristics of the vacuum device 301 can be utilized with vacuum pumps 214 of any size or type by modifying the pump control circuitry 210 and the vacuum sensor 211 expanding the size and types of vacuum assisted devices that can be supported.

In an embodiment of the present invention, a vacuum device 301 to enable a specified vacuum to be applied to a suction device 435 attached to tissue of a mammal, comprises a sensor for monitoring a vacuum applied to the suction device. A pump 214 for increasing the vacuum to the suction device 435. A valve 221 for immediately releasing the vacuum applied to the suction device and a control circuit. Wherein the control unit 419 controls one or more of activating the pump, opening the valve 221 and adjusting operating parameters of the vacuum device. Wherein activating the pump 214 applies the specified vacuum to the suction device. Wherein opening the valve 221 immediately releases the vacuum applied to the suction device 435 to release the tissue grasped by the suction device 435. Wherein adjusting operating parameters of the vacuum device modifies the specified vacuum, wherein the control circuit is positioned to be operated with the hand holding the suction device.

In an embodiment of the present invention, the vacuum device 301 can be made by providing the individual electronic components that comprise the functional capability for the circuitry blocks shown on the device circuit board 209. These items can be assembled using automated or manual means to the circuit board and then soldered in place. The energy storage device 215 and the vacuum pump 214 can then be connected as shown in FIG. 2, such as by soldering the electrical connections. The flexible tubing 216 can be installed by manual means to inter-connect the vacuum related devices as represented on the drawing. The foot switch 218 can be assembled separately and can be wired directly to the vacuum device 301 or provided as a wireless accessory to the vacuum device 301.

In an embodiment of the present invention, to use the vacuum device 301, an operator can select the suction device 435 necessary for the intended medical procedure and attach such to the vacuum device 301 using the vacuum assisted device tubing 308 that can be then attached to the vacuum attachment port 206. In an embodiment of the present invention, the invention can be used as an ancillary/back-up vacuum pump, in support of hand-operated vacuum devices. In an alternative embodiment of the present invention, the vacuum device 301 can be used as a stand-alone primary vacuum pump.

In an embodiment of the present invention, the operator then decides the desired vacuum pressure required for the selected suction device 435 and uses either the vacuum selector switch 203 to choose an operating vacuum, adjusts the vacuum pressure selection using the functionality afforded by the operator display and controls 204 or uses the wireless control unit 419.

In an embodiment of the present invention, once the portable regulated vacuum device 301 is set by the operator and powered on, the vacuum pump 214 can operate to reach and maintain the vacuum pressure setting when selected to do so by the foot switch 218 condition (on/off) or wireless control 419. In an embodiment of the present invention, the operator can monitor the relative performance of the pump and its delivered vacuum by viewing the LED bargraph indicator 205, using the indicator colors green, yellow and red to

readily determine if the vacuum pressure is acceptable for the medical procedure task. In an alternative embodiment of the present invention, the operator can be apprised of the relative performance of the pump through audible signals. In an embodiment of the present invention, audible signals can be used to warn the surgeon of critical parameters throughout the surgical procedure. In an embodiment of the present invention, audible signals can be used to warn the surgeon of critical vacuum or critical vacuum in relation to time parameters. In an embodiment of the present invention, the vacuum device 301, by applying a pulsating vacuum can inhibit bleeding at a wound. In an embodiment of the present invention, the vacuum device 301, by applying a pulsating vacuum based on a patient's pulse can inhibit bleeding at a wound.

In an embodiment of the present invention, the vacuum pressure records can be stored in the processor memory during the execution of the medical procedure. In an embodiment of the present invention, upon the conclusion of the procedure, the vacuum pressure records, stored in the processor memory during the execution of the medical procedure can be retrieved. In an embodiment of the invention, the vacuum pressure records can be in the form of a histogram. The retrieval of the stored data can be performed by the simple insertion of a flash memory device into the external memory and interface port 207 or by connection of this port to a host computer for subsequent download. Once the medical procedure is completed, and/or the stored data is retrieved, the vacuum device 301 can then be powered off using the power switch 202.

In an embodiment of the invention, the system control functionality of the vacuum device 301 can be used to provide positive pressures, through the simple exchange of pump hose connectivity, sensor selection and processor firmware control algorithm changes. The device application can then be reversed, providing positive pressure, rather than vacuum, to devices requiring such positive pressure assistance.

In another embodiment of the invention, a Radio Frequency IDentification (RFID) tag is imbedded in one or more of the vacuum devices 435. In an embodiment of the invention, the RFID tag is used to identify the vacuum device and thereby determine the parameters for operation of the vacuum device 301. In one embodiment of the invention, the RFID tag operates using an Ultra High Frequency (UHF) signal. In another embodiment of the invention, the RFID tag operates using a microwave frequency signal.

In an embodiment of the present invention, a RFID reader is present in the vacuum device 301 which can then read the RFID tags in the individual vacuum devices. In an embodiment of the invention, the RFID reader can be positioned so that the RFID tag antenna is least affected by any conducting material.

In one embodiment the RFID tag is read only. In another embodiment, the RFID tag contains an Electrically Erasable Programmable Read-Only Memory (EPROM), which enables both read and write functions. In an embodiment of the invention, the RFID tag is passive. In another embodiment of the invention, the RFID tag is semi passive containing a source of energy such as a battery to allow the tag to be constantly powered. In a further embodiment of the invention, the RFID tag is active, containing an internal power source, such as a battery, which is used to power any Integrated Circuits (ICs) in the tag and generate the outgoing signal. In another embodiment, the tag has the ability to enable location sensing through a photo sensor.

In an embodiment of the invention, the vacuum device 301 is able to monitor the type, previous use data and condition of the vacuum device 435. In this manner, a surgeon can choose

when a surgical procedure warrants using the same vacuum device 301 that has previously been used for a similar surgical procedure using similar parameters and under the same or similar conditions.

5 In one embodiment of the invention, means of communication with a base station is embedded in the vacuum device 301.

In one embodiment of the invention, the communication means utilizes one or more of a wireless local area network; a 10 wireless wide area network; a cellular network; a satellite network; a Wi-Fi network; and a pager network. In one embodiment of the invention, a modem capable of communicating with one or more of the aforementioned networks is embedded in the vacuum device 301. In the following discussion the term 'cellular modem' will be used to describe the device embedded. The term 'cellular modem' will be herein used to identify any device of comparable size capable of communicating over one or more of the aforementioned networks. In one embodiment of the invention, the cellular modem can be a Code Division Multiple Access (CDMA) 15 modem. In an embodiment of the invention, a RFID reader and associate integrated circuit processor can be embedded together with the cellular modem in the vacuum device 301. In such an embodiment, the RFID tags and RFID reader can be positioned to optimize the RFID read of the RFID tags from the available devices 435.

In an embodiment of the invention, a RFID reader and a 20 cellular modem can be positioned in the vacuum device 301; the RFID reader is in communication with one or more RFID readers, associated cellular modems and the RFID tags of one or more vacuum devices 301. Through communications with the RFID reader and associated integrated circuit processor of the plurality of vacuum devices 301, a RFID reader and associated integrated circuit processor is able to distinguish 25 the RFID tag from vacuum devices 301 in the vicinity based on one or more of location, strength of signal, variation of RFID tag signal with position, variation of RFID tag signal with time and prior input data. In an embodiment of the invention, one or more antenna inserted can be used to help 30 discriminate the location of the vacuum devices 301. In an embodiment of the invention, the RFID reader and associate processor can be in communication with the cellular modem. In an embodiment of the invention, the cellular modem is in communication with a base station and can transmit one or 35 more parameters selected from the group consisting of one or more RFID tag location, one or more RFID tag identification code, vacuum device 301, suction device 435, vacuum device conditions, suction device conditions and time stamp.

In an embodiment of the invention, the vacuum device can 40 be set to allow for a preset hysteresis. As shown in FIG. 6, when the vacuum device can be set at a preset vacuum 610, a range of vacuum 650 can be acceptable, given by the maximum acceptable vacuum 620 and the minimum acceptable vacuum 630. The vacuum 640 increases above the preset 45 vacuum 610 and the trigger point to turn off the pump 214 occurs when the vacuum 640 exceeds the maximum acceptable vacuum 620. Once the pump 214 is turned off or the relief valve 221 opened, leakage and out gassing of adsorbed molecules result in an increase in the pressure (loss of vacuum) 50 until the pump 214 is turned on when the vacuum 640 falls below the minimum acceptable vacuum 630. As the pump 214 is turned off and on the hysteresis is reduced and the 55 preset vacuum 610 can be attained.

In an embodiment of the invention, the vacuum device can 60 be set to allow for a preset dwell time 750. As shown in FIG. 7, when the vacuum device can be set at a preset vacuum 710, given by the maximum acceptable vacuum 720 and the mini-

mum acceptable vacuum 730 and preset dwell time 750. After the vacuum 740 has attained a stable preset vacuum 710 (see FIG. 6), breaking the suction device seal, leakage and/or out-gassing of adsorbed molecules can result in an increase in the pressure (loss of vacuum). The dwell time 750 sets the wait time after the minimum acceptable vacuum has been triggered to turn on the pump 214. After the pump 214 is turned on the vacuum continues to fall until the pump 214 has compensated for the leak. Thereafter, the pump 214 increases the vacuum 740 above the preset vacuum 710 and the trigger point to turn off the pump 214 occurs when the vacuum 740 exceeds the maximum acceptable vacuum 720.

In one embodiment of the invention the RFID code uses the IEEE format and is Electronic Product Code (EPC) readable. In another embodiment of the invention the RFID code uses the UCC format and is Universal Product Code (UPC) readable. In another embodiment, the format is compatible for EPC, European Article Number (EAN) and UPC read and write functions.

In an embodiment of the invention, the device method or system can be used for the treatment of humans. In an embodiment of the invention, the device method or system can be used for the treatment of animals. In an embodiment of the invention, the device method or system can be used in veterinary applications. In an embodiment of the invention, the device method or system can be used in medical applications.

In an embodiment of the invention, a vacuum device and suction device are assembled as an integral unit (see FIGS. 5A and B). As shown in FIG. 5A, the integral suction and vacuum device 500 can have an accessible on off switch 202, located on or adjacent to the vacuum device circuit board 209, which activates the pump control circuitry 210, wherein the vacuum sensor can monitor the vacuum applied to the suction device 435. These components can be attached within or onto integral suction and vacuum device housing 510. The energy storage device 215 and the vacuum pump 214 can also be housed in the integral suction and vacuum device housing 510, onto which the suction device can be attached 435. The integral suction and vacuum device 500 can be a compact unit as shown in FIG. 5B. In an embodiment of the invention, an integral suction and vacuum device 500 used for tamponade can be “joined” or can be assembled into one unit as shown in FIG. 5. In an embodiment of the invention, an integral suction and vacuum device 500 is portable, ergonomic, and superior for in situ use. FIG. 8A is a frontal diagram showing a patient 800 with a number of wound or laceration sites 810. FIG. 8B is a frontal diagram showing the utilization of an integral suction and vacuum device 500 on a patient 800 in accordance with an embodiment of the invention. FIG. 8C is a cross section diagram showing the utilization of an integral suction and vacuum device 500 on a patient 800 in accordance with an embodiment of the invention. As shown in FIGS. 8B-8C the integral suction and vacuum device 500 can be utilized with a patient 800. As the patient moves, the device can move to compensate for the movement. In contrast, if the vacuum device was separate, or relied upon a tether to hold the device to the patient, then movement could jar and detach the suction device. In an embodiment of the invention, in an integral suction and vacuum device 500, the suction device is integrated into the vacuum device for use by a first-responder to treat a wound. In an embodiment of the invention, in an integral suction and vacuum device 500, the suction device is integrated into the vacuum device for use by a first-responder to curtail bleeding. In an embodiment of the invention, an integral suction and vacuum device 500 can be used in a battlefield to stem bleeding from laceration sites 810. In an

embodiment of the invention, an integral suction and vacuum device 500 can be used in a battlefield to treat one or more wound sites 810. In an embodiment of the invention, a sterilized material is used to clean the wound prior to applying the integral suction and vacuum device 500 which can be positioned with the suction device 435 fully encompassing multiple wound sites 810. In an embodiment of the invention, a sterilized material is used to temporarily remove blood from the laceration sites in order to identify the site(s) of bleeding, so that the integral suction and vacuum device 500 can be positioned with the suction device 435 fully encompassing the multiple laceration sites 810. In an embodiment of the invention, the integral suction and vacuum device 500 can be packaged in a sterilized container so that the wound is not contaminated when the integral suction and vacuum device 500 is applied to the mammal. This will help ensure that the wound site(s) can be kept clean and free of bacteria. In an embodiment of the invention, the integral suction and vacuum device 500 can be packaged in a sterilized container so that the laceration sites are not contaminated when the integral suction and vacuum device 500 is applied to the mammal. This will help ensure that the laceration site(s) can be kept clean and free of bacteria. In an embodiment of the invention, the integral suction and vacuum device 500 can be sealed in a hermetically sealed package which is opened directly prior to the integral suction and vacuum device 500 being applied to the mammal. In an embodiment of the invention, once the wound or laceration site can be established an integral suction and vacuum device 500 can be applied to treat the mammal. In an embodiment of the invention, an integral suction and vacuum device 500 can be used once and then discarded to minimize the risk of contamination. In an embodiment of the invention, an integral suction and vacuum device 500 can be used once and then discarded to eliminate the risk of blood or tissue contamination. In an embodiment of the invention, the integral suction and vacuum device 500 can replace traditional dressings and/or a tourniquet.

In an embodiment of the invention, a vacuum device to enable a specified vacuum to be applied to a suction device attached to tissue, comprises a sensor for monitoring a vacuum applied to the suction device, a pump for increasing the vacuum to the suction device, a valve for immediately releasing the vacuum applied to the suction device to release the tissue and a control circuit. Wherein the control circuit controls two or more functions selected from the group consisting of activating the pump, opening the valve and adjusting operating parameters of the vacuum device. Wherein activating the pump applies the specified vacuum to the suction device. Wherein opening the valve immediately releases the vacuum applied to the suction device. Wherein adjusting operating parameters of the vacuum device modifies the specified vacuum. Wherein the control circuit is operated while holding the suction device.

In an embodiment of the invention, the vacuum device further comprises a control unit mounted one or both on and adjacent to the suction device, wherein the control unit allows one or more of control of the vacuum applied to the suction device, control of the vacuum device and adjustment of the vacuum device settings during the procedure.

In an embodiment of the invention, the vacuum device further comprises a foot switch, wherein the foot switch allows one or more of control of the vacuum applied to the suction device, control of the vacuum device and adjustment of the vacuum device settings during the procedure.

In an embodiment of the invention, the vacuum device further comprises one or both visual and audio feedback that allows one or more methods of control of the vacuum applied

to the suction device, control of the vacuum device and adjustment of the vacuum device settings during the procedure.

In an embodiment of the invention, the vacuum device is portable. In an embodiment of the invention, the vacuum device is hand held. In an embodiment of the invention, the vacuum device is portable and hand held.

In an embodiment of the invention, a method of adjusting and monitoring a vacuum device during a surgical procedure, comprises receiving the suction device attached to tissue and receiving a vacuum device for applying a vacuum to the suction device during the surgical procedure, wherein the vacuum device one or both receives and has preset parameters to control the vacuum to be applied to the suction device. Activating the vacuum device, wherein the vacuum device functions include monitoring the vacuum applied to the suction device via a sensor, comparing the vacuum applied to the suction device and the parameters at regular time intervals using a processor and automatically increasing the vacuum to the suction device when the comparison indicates an increased vacuum is required. The method further comprises monitoring the vacuum device during the surgical procedure using one or both audio or visual feedback from the vacuum device and adjusting the parameters selected in response to changed conditions of the surgical procedure while the vacuum is applied to the tissue. In an embodiment of the invention, the vacuum is adjusted to control against separation of the suction device from the tissue at the rim of the suction device where it contacts the tissue.

In an embodiment of the invention, a system for adjusting and monitoring a suction device comprises a vacuum device for applying a vacuum to the suction device, wherein the vacuum device includes an RFID tag reader, wherein the RFID tag reader can read an RFID tag on the suction device, an input module for selecting parameters for use with the vacuum device, wherein the input module selects parameters based at least in part on the suction device selected, a sensor for monitoring the vacuum applied by the vacuum device to the suction device and a processor for comparing the vacuum applied to the suction device and the selected parameters and automatically activating the vacuum device to increase the vacuum to the suction device when the comparison indicates an increased vacuum is required, the suction device including the RFID tag and a control unit, wherein the RFID tag identifies parameters associated with the suction device, wherein the control unit adjusts the vacuum device parameters in response to changed conditions and audio or visual feedback of the vacuum applied to the suction device during the surgical procedure, wherein based on audio or visual feedback the control unit is used to adjust the vacuum device parameters.

In an embodiment of the invention, a system for a first responder to minimize bleeding of a wound in a mammal comprises an integral suction and vacuum device for applying a vacuum to the wound. The integral suction and vacuum device includes a switch to activate the integral suction and vacuum device, a suction cup, a relief valve, a vacuum pump, wherein the vacuum pump generates a vacuum at the suction cup, a sensor for monitoring the vacuum at the suction cup and a processor. The processor compares the vacuum applied to the suction cup and selected parameters and automatically activating the vacuum pump to increase the vacuum to the suction cup when the comparison indicates an increased vacuum is required and one or both deactivating the vacuum pump and automatically opening the relief valve when the comparison indicates a decreased vacuum is required. Application of the suction cup and the vacuum to the wound attaches the integral suction and vacuum device to the mammal such that movement of the mammal does not detach the

integral suction and vacuum device. The system further comprises a material to one or both identify the wound and wipe blood away from the wound. After the wound site(s) are identified the integral suction and vacuum device is applied to the wound and activated to one or more of treat the wound, stop or minimize bleeding of a wound.

In an embodiment of the invention, a system for adjusting and monitoring a suction device during a surgical procedure, comprises a vacuum device for applying a vacuum to the suction device. The vacuum device includes an input module for selecting parameters for use with the vacuum device, wherein the input module selects parameters based at least in part on the suction device selected, a sensor for monitoring the vacuum applied by the vacuum device to the suction device, a processor for comparing the vacuum applied to the suction device and the selected parameters and automatically activating the vacuum device to increase the vacuum to the suction device when the comparison indicates an increased vacuum is required and audio or visual feedback to monitor the vacuum applied to the suction device during the surgical procedure. The system also comprises the suction device attached to a subject's tissue and a device to monitor the subject's pulse, wherein the subject's pulse is used to adjust the vacuum device parameters.

In an embodiment of the invention, the vacuum device 301 can be applied in any scientific, manufacturing, or industrial apparatus that requires the use of a regulated constant or variable vacuum. This can include laboratory equipment that requires vacuum assisted grasping, reactions, sampling, storage or any other clinical procedure. The manipulation of components, fluids or assemblies used in a manufacturing process, including: precision handling, clean-room transport, and material transport can also be supported.

Various embodiments can be implemented using a conventional general purpose or specialized digital computer(s) and/or processor(s) programmed according to the teachings of the present disclosure, as will be apparent to those skilled in the computer art. Appropriate software coding can readily be prepared by skilled programmers based on the teachings of the present disclosure, as will be apparent to those skilled in the software art. The invention can also be implemented by the preparation of integrated circuits and/or by interconnecting an appropriate network of component circuits, as will be readily apparent to those skilled in the art.

Embodiments of the present invention can include a computer readable medium, such as computer readable storage medium. The computer readable storage medium can have stored instructions which can be used to program a computer to perform any of the features present herein. The storage medium can include, but is not limited to, any type of disk including floppy disks, optical discs, DVD, CD-ROMs, micro drive, and magneto-optical disks, ROMs, RAMs, EPROMs, EEPROMs, DRAMs, flash memory or any media or device suitable for storing instructions and/or data. The present invention can include software for controlling both the hardware of a computer, such as general purpose/specialized computer(s) or microprocessor(s), and for enabling them to interact with a human user or other mechanism utilizing the results of the present invention. Such software may include, but is not limited to, device drivers, operating systems, execution environments/containers, and user applications.

Embodiments of the present invention can include providing code for implementing processes of the present invention. The providing can include providing code to a user in any manner. For example, the providing can include transmitting

digital signals containing the code to a user; providing the code on a physical media to a user; or any other method of making the code available.

Embodiments of the present invention can include a computer-implemented method for transmitting the code which can be executed at a computer to perform any of the processes of embodiments of the present invention. The transmitting can include transfer through any portion of a network, such as the Internet; through wires, the atmosphere or space; or any other type of transmission. The transmitting can include initiating a transmission of code; or causing the code to pass into any region or country from another region or country. A transmission to a user can include any transmission received by the user in any region or country, regardless of the location from which the transmission is sent.

The foregoing description of embodiments of the methods, systems, and components of the present invention has been provided for the purposes of illustration and description. It is not intended to be exhaustive or to limit the invention to the precise forms disclosed. Many modifications and variations will be apparent to one of ordinary skill in the relevant arts. For example, steps performed in the embodiments of the invention disclosed can be performed in alternate orders, certain steps can be omitted, and additional steps can be added. The embodiments were chosen and described in order to best explain the principles of the invention and its practical application, thereby enabling others skilled in the art to understand the invention for various embodiments and with various modifications that are suited to the particular uses contemplated. Other embodiments are possible and are covered by the invention. Such embodiments will be apparent to persons skilled in the relevant art(s) based on the teachings contained herein. The breadth and scope of the present invention should not be limited by any of the above-described exemplary embodiments, but should be defined only in accordance with the following claims and their equivalents.

What is claimed is:

1. A method to attach an integrated vacuum device to a patient's tissue, comprising:

- (a) receiving a vacuum device including a valve, a radio frequency identification (RFID) tag reader, a sensor, and a processor;
- (b) inserting a suction device including a programmable RFID tag into the vacuum device to form an integrated vacuum device;
- (c) reading the programmable RFID tag to identify the suction device and prior use data of the suction device;
- (d) receiving one or more operating parameters;
- (e) activating the integrated vacuum device to generate a vacuum provided the prior use data indicates no possibility of contamination from prior use of the suction device; and
- (f) attaching the integrated vacuum device to the patient's tissue.

2. The method of claim 1, where the vacuum is pulsatile and varied opposite the patient's pulse.

3. The method of claim 1, further comprising grasping the patient's tissue with the integrated vacuum device to manipulate the patient's tissue.

4. The method of claim 1, where the vacuum is increased when the patient's blood pressure to the patient's tissue is in diastole.

5. The method of claim 1, where the vacuum is marginally increased in diastole and maintained to overcome the systolic pressure pulse.

6. A method to monitor for a 'pop-off' event during a surgical procedure on a patient, comprising:

(a) receiving a vacuum device including a valve, a radio frequency identification (RFID) tag reader, a processor, a sensor for monitoring a vacuum and a sensor for monitoring a patient's pulse;

(b) inserting a suction device including a programmable RFID tag into the vacuum device to form an integrated vacuum device;

(c) reading the programmable RFID tag to identify the suction device;

(d) receiving one or more operating parameters;

(e) activating the integrated vacuum device to generate a vacuum to the suction device, where the vacuum is pulsatile and varied opposite the patient's pulse;

(f) attaching the integrated suction device to the patient's tissue during the surgical procedure;

(g) monitoring for a 'pop-off' event based on the change in vacuum over time; and

(h) recording one or both the 'pop-off' event and the release of the vacuum through the valve at completion of the surgical procedure.

7. The method of claim 6, where reading the programmable RFID tag includes reading prior use data of the suction device, where the prior use data is used to eliminate the possibility of contamination from prior use of the suction device.

8. The method of claim 6, further comprising grasping with the integrated vacuum device to manipulate the tissue.

9. The method of claim 6, where the vacuum is increased when the blood pressure to the tissue is in diastole.

10. The method of claim 9, where the vacuum is marginally increased in diastole and maintained to overcome the systolic pressure pulse.

11. A system for a first responder to grasp tissue comprising:

a suction device adapted to attach to a mammal for grasping a tissue of the mammal including a programmable radio frequency identification (RFID) tag;

a vacuum device adapted to receive the suction device, where upon insertion of the vacuum device into the suction device an integrated vacuum device is formed with no intermediate connections, where the integrated vacuum device further includes:

- (i) an RFID tag read/write device, where the vacuum device is adapted to read the RFID tag of the suction device and one or both recognize the suction device and retrieve previous use data of the suction device, where the vacuum device is adapted to apply a vacuum to the tissue of the mammal sufficient for grasping of the tissue of the mammal, where the vacuum generated is based on one or more of the tissue being grasped, the suction device recognized and the previous use data;
- (ii) a switch to activate the integrated vacuum device;
- (iii) a relief valve;
- (iv) one or more sensors adapted to determine two or more sensor parameters selected from the group consisting of the vacuum at the tissue of the mammal, the temperature of the mammal, the blood pressure of the mammal and the change in blood pressure of the mammal over time; and

- (v) a processor programmed to carry out steps including: reading the programmable RFID tag to retrieve one or more device parameters, adjusting the vacuum based on one or more of the one or more device parameters and the two or more sensor parameters such that the vacuum at the tissue of the mammal is maintained during one or both transport and movement of the mammal, monitoring the one or more sensors, deactivating the integrated

25

vacuum device and automatically opening the relief valve as required by one or both safety guidelines and the completion of grasping treatment, and updating the previous use data of the programmable RFID tag.

12. The system of claim 11, where the suction device is selected from the group consisting of an obstetrical grasping cup, a retraction device, a suction cup, a manipulating device, a tamponade device, a traction device, open cup graspers and a tourniquet.

13. The system of claim 11, where the integrated vacuum device is used in one or both laparoscopic and thorascoscopic surgical procedures.

14. The system of claim 11, where the integrated vacuum device is used in procedures employing an open incision selected from the group consisting of laparotomy, thoracotomy, craniotomy, retroperitoneal surgical approaches and orthopedic vacuum traction or manipulation of bony surfaces.

15. The system of claim 11, where the integrated vacuum device is used on a bony surface or on the structures under or within an incision in bone or aperture in bone including but not limited to craniotomy and orthopedic surgery.

10 16. The system of claim 11, where the integrated vacuum device applies a pulsatile vacuum based on a mammal's pulse to inhibit bleeding at the tissue of the mammal.

5 17. The system of claim 11, where the vacuum device alone provides the vacuum to the suction device.

18. The system of claim 11, where the vacuum device is used to one or both maintain the vacuum and assist in generating the vacuum of a hand-operated vacuum pump.

19. The system of claim 11, further comprising a remote device which allows control of one or more functions selected from the group consisting of the vacuum applied to the suction device, the vacuum device and adjustment of the vacuum device settings while the vacuum device is activated.

15 20. The system of claim 11, further comprising one or both a visual and an audio feedback unit that allows one or more functions selected from the group consisting of control of the vacuum applied to the suction device, control of the vacuum device and adjustment of the vacuum device settings while the vacuum device is activated.

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专利名称(译)	便携式调节真空泵，用于医疗程序		
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[标]申请(专利权)人(译)	LONKY NEAL MARC GURGANIAN ALBERT 史蒂夫		
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当前申请(专利权)人(译)	MEDITECH发展INCORPORATED		
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摘要(译)

在本发明的各种实施例中，电动真空泵可以用作独立单元来创建，监视和控制真空，或者用于在各种外科手术中维护，辅助或作为手动泵的备用。真空泵由处理器控制，并且当真空低于预先选择的参数或外科医生在运行中选择的参数时自动启动。视觉或音频反馈用于允许外科医生在执行外科手术时增加对真空装置的控制。

