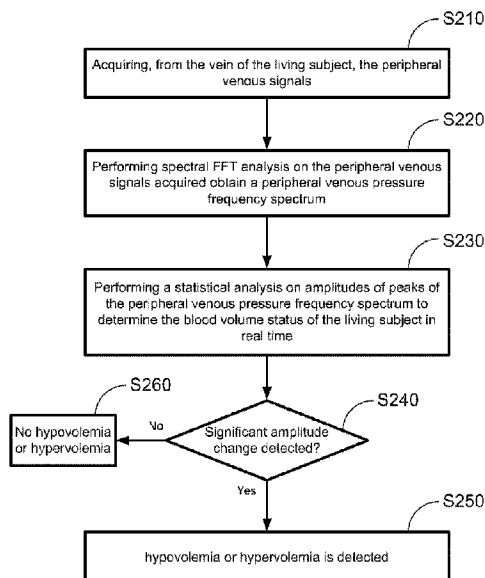




(86) Date de dépôt PCT/PCT Filing Date: 2015/09/14
(87) Date publication PCT/PCT Publication Date: 2016/03/17
(45) Date de délivrance/Issue Date: 2023/01/17
(85) Entrée phase nationale/National Entry: 2017/03/13
(86) N° demande PCT/PCT Application No.: US 2015/050001
(87) N° publication PCT/PCT Publication No.: 2016/040947
(30) Priorités/Priorities: 2014/09/12 (US62/049,829);
2015/09/14 (US14/853,504)

(51) Cl.Int./Int.Cl. *A61B 5/021* (2006.01),
A61B 5/00 (2006.01), *A61B 5/02* (2006.01),
A61B 5/0215 (2006.01)
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(54) Titre : DETECTION D'HYPOVOLEMIE/D'HYPERVOLEMIE EN UTILISANT UNE ANALYSE DE FORME D'ONDE
INTRAVEINEUSE PERIPHERIQUE (PIVA) ET APPLICATIONS ASSOCIEE
(54) Title: HYPOVOLEMIA/HYPERVOLEMIA DETECTION USING PERIPHERAL INTRAVENOUS WAVEFORM
ANALYSIS (PIVA) AND APPLICATIONS OF SAME



(57) **Abrégé/Abstract:**

Aspects of the invention relates to systems and methods for hypovolemia and/or hypervolemia detection of a living subject using peripheral intravenous waveform analysis. In one embodiment, the method includes: acquiring, from a vein of the living subject, peripheral venous signals; performing a spectral analysis on the acquired peripheral venous signals to obtain a peripheral venous pressure frequency spectrum; and performing a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum to determine the blood volume status of the living subject in real time. Specifically, at least two peaks, respectively corresponding to a first frequency and a second frequency, are obtained on the peripheral venous pressure frequency spectrum. Amplitude change of the second peak is used to determine the blood volume status of the living subject. Hemorrhage may be detected when a significant amplitude decrease is detected from the second baseline peak to the second peak.

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property
Organization
International Bureau



(10) International Publication Number
WO 2016/040947 A1

(43) International Publication Date
17 March 2016 (17.03.2016)

(51) International Patent Classification:

A61B 5/021 (2006.01) A61B 5/00 (2006.01)
A61B 5/02 (2006.01)

(21) International Application Number:

PCT/US2015/050001

(22) International Filing Date:

14 September 2015 (14.09.2015)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

62/049,829 12 September 2014 (12.09.2014) US
14/853,504 14 September 2015 (14.09.2015) US

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(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, ST, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, KM, ML, MR, NE, SN, TD, TG).

Published:

— with international search report (Art. 21(3))

(54) Title: HYPOVOLEMIA/HYPERVOLEMIA DETECTION USING PERIPHERAL INTRAVENOUS WAVEFORM ANALYSIS (PIVA) AND APPLICATIONS OF SAME

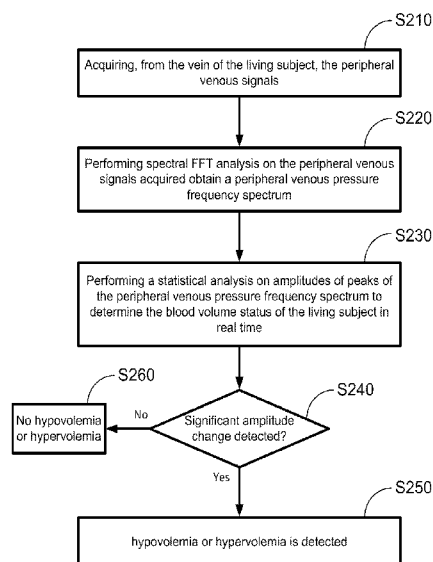


FIG. 2

(57) Abstract: Aspects of the invention relates to systems and methods for hypovolemia and/or hypervolemia detection of a living subject using peripheral intravenous waveform analysis. In one embodiment, the method includes: acquiring, from a vein of the living subject, peripheral venous signals; performing a spectral analysis on the acquired peripheral venous signals to obtain a peripheral venous pressure frequency spectrum; and performing a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum to determine the blood volume status of the living subject in real time. Specifically, at least two peaks, respectively corresponding to a first frequency and a second frequency, are obtained on the peripheral venous pressure frequency spectrum. Amplitude change of the second peak is used to determine the blood volume status of the living subject. Hemorrhage may be detected when a significant amplitude decrease is detected from the second baseline peak to the second peak.

WO 2016/040947 A1

HYPOVOLEMIA/HYPERVOLEMIA DETECTION USING PERIPHERAL INTRAVENOUS WAVEFORM ANALYSIS (PIVA) AND APPLICATIONS OF SAME

10

Some references, which may include patents, patent applications and various
20 publications, are cited and discussed in the description of this invention. The citation and/or
discussion of such references is provided merely to clarify the description of the present
invention and is not an admission that any such reference is “prior art” to the invention
described herein.

25 In terms of notation, hereinafter, “[n]” represents the n-th reference cited in the reference list. For example, [1] represents the first reference cited in the reference list, namely, Wilson, M., D.P. Davis, and R. Coimbra, *Diagnosis and monitoring of hemorrhagic shock during the initial resuscitation of multiple trauma patients: a review*. The Journal of Emergency Medicine, 2003. **24**(4): p. 413-422.

FIELD OF THE INVENTION

detecting hypovolemia and/or hypervolemia using peripheral IV waveform analysis (PIVA) to assess blood volume status of a living subject, and applications of the same.

BACKGROUND OF THE INVENTION

5 Hemorrhagic shock remains the leading preventable cause of death in the casualty care setting [42, 43]. Trauma is the leading cause of death in patients less than 40 years of age and poses a significant economic burden worldwide. Survival is contingent upon early recognition of hemorrhage, appropriate triage, and goal-directed transfusion therapy [43, 44].

Early recognition of hemorrhage and guided fluid administration is critical for
10 providing timely intervention and maintaining end organ viability [1] of human beings and other living animals. For example, Timely damage control surgery (DCS) and restrictive fluid resuscitation (RFR) for wounded soldiers have been shown to significantly improve mortality [45]. While early resuscitation is warranted in patients with hypovolemic shock, over-resuscitation can also have deleterious effects, such as decreased end-organ perfusion,
15 pulmonary edema, and increased mortality [2] [3] [4]. However, there remains a critical technological void for detecting early hemorrhage and resuscitation efficacy, particularly in the clinical setting. Specifically, recognition of subclinical hemorrhage and proper fluid resuscitation has remained elusive, resulting in delayed triage and poor management of the wounded soldier [46]. Often, continuous occult bleeding is not recognized until the onset of
20 hemorrhagic shock and hemodynamic collapse, particularly in a young, healthy soldier with good compensatory mechanisms [47, 48]. Unrecognized hemorrhage leads to delayed triage and DCR, resulting in preventable end-organ damage [49, 50, 35].

Early recognition of hemorrhage is the mainstay of therapy to prevent end-organ damage and to improve survival [34]. Hypovolemic shock and end organ damage are
25 challenging diagnoses in patients with normal heart rate and blood pressure, particularly in patients with good compensatory mechanisms [35]. Specifically, advanced hemorrhagic shock often precedes significant changes in standard vital sign monitoring, particularly in young healthy individuals [5] [1]. In fact, significant vital sign changes do not occur until the patient has lost 15-30% of blood volume, characteristic for Stage II hemorrhage [5]. Even
30 invasive monitoring modalities such as central venous pressure and pulmonary arterial pressure are poor determinants of volume status [6]. Pulmonary artery occlusion pressure and central venous pressure fail to predict ventricular filling volume, cardiac performance, or the response to volume infusion in normal subjects [6]. Furthermore, central monitoring is associated with major vascular complications and central line associated bloodstream

infections [7]. More importantly, central venous monitoring offers no survival benefit in critically-ill patients [8] [9].

Therefore, a heretofore unaddressed need exists in the art to address the aforementioned deficiencies and inadequacies.

5

SUMMARY OF THE INVENTION

In one aspect, the present invention relates to a method for determining hypovolemia, hypervolemia and vascular tone of a living subject based on an intravascular volume status of the living subject. In certain embodiments, the method includes: acquiring, continuously for
10 a time period from T_0 to T_2 , peripheral venous signals from a vein of the living subject, wherein the time period is divided into a first time period from T_0 to T_1 , and a second time period from T_1 to T_2 ; processing the peripheral venous signals acquired at the first time period to obtain a baseline peripheral venous pressure frequency spectrum; obtaining a plurality of baseline peaks $\{B_{N-1}\}$ on the baseline peripheral venous pressure frequency
15 spectrum, wherein N is a positive integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to a plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} ; processing the peripheral venous signals acquired at the second time period to obtain a peripheral venous pressure frequency spectrum; obtaining a plurality of peaks $\{P_{N-1}\}$ on the peripheral venous
20 pressure frequency spectrum, wherein the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$; and determining the intravascular volume status of the living subject at the second time period by comparing amplitudes of the peaks $\{P_{N-1}\}$ to that of the baseline peaks $\{B_{N-1}\}$ respectively. The intravascular volume status of the living subject at the second time period
25 indicates hypovolemia or hypervolemia when amplitude changes greater than a threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$.

In certain embodiments, the intravascular volume status of the living subject at the second time period indicates hypovolemia when amplitude decreases are detected greater than a first threshold from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$. In certain
30 embodiments, the intravascular volume status of the living subject at the second time period indicates hypervolemia when amplitude increases greater than a second threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$.

In certain embodiments, the method is performed to the living subject during resuscitation of the living subject. In one embodiment, the intravascular volume status of the

living subject at the second time period indicates a return of euvoemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at the first time period, and amplitude increases greater than a third threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$. In one embodiment, the intravascular volume status of the living subject at the second time period indicates over-resuscitation when the living subject is determined to be in an euvoemic state at the first time period, and amplitude increases greater than a fourth threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$.

In certain embodiments, the peripheral venous signals are acquired by: inducing anesthesia on the living subject; inserting a peripheral intravenous (IV) catheter into the vein of the living subject, wherein the vein is an upper extremity vein; and capturing and recording the peripheral venous signals from the peripheral IV catheter at a sampling rate of about 1 kHz. In one embodiment, the peripheral IV catheter is a peripherally-inserted central catheter (PICC). In certain embodiments, a pressure transducer is used to be directly connected to the peripheral IV catheter, and the peripheral venous signals are captured and recorded by the pressure transducer.

In certain embodiments, the peripheral venous signals are processed by a spectral fast Fourier transform (FFT) analysis to obtain the baseline peripheral venous pressure frequency spectrum and the peripheral venous pressure frequency spectrum, respectively.

In certain embodiments, the plurality of peaks $\{P_{N-1}\}$ includes a first peak corresponding to a first frequency F_0 and a second peak corresponding to a second frequency F_1 . In one embodiment, the first peak corresponding to the first frequency F_0 is associated with a respiratory rate of the living subject; and the second peak corresponding to the second frequency F_1 is associated with a heart rate of the living subject.

Another aspect of the present invention relates to a method for determining a blood volume status of a living subject, which includes: acquiring, from a vein of the living subject, peripheral venous signals; performing a spectral analysis on the acquired peripheral venous signals to obtain a peripheral venous pressure frequency spectrum; and performing a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum to determine the blood volume status of the living subject in real time.

In certain embodiments, the intravascular volume status of the living subject indicates hypovolemia when amplitude decreases greater than a first threshold are detected from the peaks of the peripheral venous pressure frequency spectrum; and the intravascular volume status of the living subject indicates hypervolemia when amplitude increases greater than a

second threshold are detected from the peaks of the peripheral venous pressure frequency spectrum. In certain embodiments, the method is performed to the living subject during ultrafiltration/dialysis or diuresis of the living subject. In one embodiment, the method further includes: generating an alert message when the intravascular volume status of the

5 living subject indicates hypovolemia.

In certain embodiments, the method is performed to the living subject during resuscitation of the living subject. In one embodiment, the intravascular volume status of the living subject indicates a return of euolemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at an earlier time period, and amplitude

10 increases greater than a third threshold are detected from the peaks of the peripheral venous pressure frequency spectrum. In one embodiment, the intravascular volume status of the living subject indicates over-resuscitation when the living subject is determined to be in an euolemic state at the earlier time period, and amplitude increases greater than a fourth threshold are detected from the peaks of the peripheral venous pressure frequency spectrum.

15 In certain embodiments, the method further includes: detecting efficacy of treatment and the return to euolemia in the living subject based on the intravascular volume status of the living subject.

In certain embodiments, the peripheral venous signal is acquired by: inserting a peripheral intravenous (IV) catheter into the vein of the living subject; and capturing and

20 recording the peripheral venous signal from the peripheral IV catheter at a sampling rate. In one embodiment, the peripheral IV catheter is a peripherally-inserted central catheter (PICC). In one embodiment, a pressure transducer is directly connected to the peripheral IV catheter, and the peripheral venous signals are captured and recorded by the pressure transducer.

In certain embodiments, the spectral analysis is a spectral fast Fourier transform

25 (FFT) analysis. In certain embodiments, the statistical analysis may include: obtaining a plurality of baseline peaks $\{B_{N-1}\}$ on a baseline peripheral venous pressure frequency spectrum, wherein N is a positive integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to a plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} ; obtaining a

30 plurality of peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum, wherein the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$; and determining the intravascular volume status of the living subject in real time by comparing the amplitudes of the peaks $\{P_{N-1}\}$ to that of the baseline peaks $\{B_{N-1}\}$ respectively.

In certain embodiments, the baseline peripheral venous pressure frequency spectrum is obtained by: acquiring the peripheral venous signals from the vein of the living subject at an earlier time period; and processing the peripheral venous signals acquired at the earlier time period by the spectral FFT analysis to obtain the baseline peripheral venous pressure
5 frequency spectrum.

In certain embodiments, the method may be capable of detecting at least 6% of blood loss or at least 5.9% of blood volume overload of the living subject.

In certain embodiments, the plurality of peaks $\{P_{N-1}\}$ includes a first peak P_0 corresponding to a first frequency F_0 and a second peak P_1 corresponding to a second
10 frequency F_1 . In certain embodiments, the first peak P_0 corresponding to the first frequency F_0 is associated with a respiratory rate of the living subject; and the second peak P_1 corresponding to the second frequency F_1 is associated with a heart rate of the living subject.

In a further aspect of the present invention, a peripheral intravenous (IV) waveform analysis (PIVA) system includes: a peripheral IV device configured to acquire, from a vein of
15 a living subject, peripheral venous signals; and an processing device communicatively connected to the peripheral IV device, configured to: receive the peripheral venous signals from the peripheral IV device; perform a spectral process and analysis on the peripheral venous signal to obtain a peripheral venous pressure frequency spectrum; and perform a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency
20 spectrum to determine the blood volume status of the living subject in real time.

In certain embodiments, the intravascular volume status of the living subject indicates hypovolemia when amplitude decreases greater than a first threshold are detected from the peaks of the peripheral venous pressure frequency spectrum; and the intravascular volume status of the living subject indicates hypervolemia when amplitude increases greater than a
25 second threshold are detected from the peaks of the peripheral venous pressure frequency spectrum. In certain embodiments, the system is applied to the living subject during ultrafiltration/dialysis or diuresis of the living subject. In one embodiment, the processing device is further configured to: generate an alert message when the intravascular volume status of the living subject indicates hypovolemia.

30 In certain embodiments, the system is applied to the living subject during resuscitation of the living subject. In one embodiment, the intravascular volume status of the living subject indicates a return of euolemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at an earlier time period, and amplitude increases greater than a third threshold are detected from the peaks of the peripheral venous

pressure frequency spectrum. In one embodiment, the intravascular volume status of the living subject indicates over-resuscitation when the living subject is determined to be in an euvoletic state at the earlier time period, and amplitude increases greater than a fourth threshold are detected from the peaks of the peripheral venous pressure frequency spectrum.

5 In certain embodiments, the processing device is further configured to: detect efficacy of treatment and the return to euvoletic state in the living subject based on the intravascular volume status of the living subject.

In certain embodiments, the processing device is communicatively connected to the peripheral IV device through a wireless connection.

10 In certain embodiments, the peripheral IV device includes: a peripheral IV catheter being inserted into the vein of the living subject; and a monitoring device connected to the peripheral IV catheter, configured to capture and record the peripheral venous signals from the peripheral IV catheter at a sampling rate. In one embodiment, the peripheral IV catheter is a peripherally-inserted central catheter (PICC). In certain embodiments, the monitoring
15 device comprises a pressure transducer directly connected to the peripheral IV catheter, wherein the peripheral venous signals are captured and recorded by the pressure transducer.

In certain embodiments, the processing device is a computing device, which may be a desktop computer, a laptop computer, a smartphone, a tablet device, or any other computing devices with processors to perform the processing functions.

20 In certain embodiments, the spectral analysis is a spectral fast Fourier transform (FFT) analysis. In certain embodiments, the statistical analysis may include: obtaining a plurality of baseline peaks $\{B_{N-1}\}$ on a baseline peripheral venous pressure frequency spectrum, wherein N is a positive integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to a plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is
25 a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} ; obtaining a plurality of peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum, wherein the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$; and determining the intravascular volume status of the living subject in real time by comparing the amplitudes of the peaks $\{P_{N-1}\}$ to that of the baseline peaks $\{B_{N-1}\}$ respectively.
30

In certain embodiments, the baseline peripheral venous pressure frequency spectrum is obtained by: acquiring the peripheral venous signals from the vein of the living subject at an earlier time period; and processing the peripheral venous signals acquired at the earlier time period by the spectral FFT analysis to obtain the baseline peripheral venous pressure

frequency spectrum.

In certain embodiments, the plurality of peaks $\{P_{N-1}\}$ includes a first peak P_0 corresponding to a first frequency F_0 and a second peak P_1 corresponding to a second frequency F_1 . In certain embodiments, the first peak P_0 corresponding to the first frequency F_0 is associated with a respiratory rate of the living subject; and the second peak P_1 corresponding to the second frequency F_1 is associated with a heart rate of the living subject.

In certain embodiments, the PIVA system further includes: a pump connected to the living subject to perform liquid exchange to the living subject; and a pump controlling mechanism communicatively connected to the processing device, configured to control the pump by intermittently pausing the pump or subtract the pump signal when the peripheral IV device acquires the peripheral venous signals, and restarting the pump when the peripheral IV device does not acquire the peripheral venous signals. The processing device is further configured to send a signal to the pump controlling mechanism to notify the pump controlling mechanism to control the pump. In certain embodiments, the pump may be a dialysis pump, a cardiopulmonary bypass pump, an extracorporeal membrane oxygenation (ECMO), or an infusion pump.

In a further aspect, the present invention relates to a method for determining hypovolemia, hypervolemia and vascular tone of a living subject based on an intravascular volume status of the living subject using the PIVA system as described above.

- In certain embodiments, the system and method as described above may be used for:
- (1) point-of-care detection of sepsis or other low systemic vascular resistance states;
 - (2) point-of-care detection of increased right ventricular pressure/afterload or right heart failure due to pulmonary embolus, idiopathic pulmonary hypertension, venous air embolus, or amniotic fluid embolus;
 - (3) point-of-care detection of high systemic vascular resistance states, such as pre-eclampsia/eclampsia;
 - (4) detection of volume status in spontaneously breathing as well as mechanically ventilated individuals, as the signal does not depend on intrathoracic changes due to positive pressure ventilation;
 - (5) automated algorithm and closed loop system for intravascular volume management – this may include, but is not limited to, automated dialysis ultrafiltration rates; and
 - (6) automated algorithm may use peak detection in the time domain, peak

detection in the Fourier transform, neural network analysis of signals, wavelet analysis, deconvolution of the time domain signal, or additional signal processing tools.

In certain embodiments, detection of hypovolemia through the system and method as
5 described above may be performed through the analysis of amplitude increases in the peaks with higher frequencies, such as F_2 , as well as the amplitude changes in F_1 .

These and other aspects of the present invention will become apparent from the following description of the preferred embodiments taken in conjunction with the following drawings, although variations and modifications thereof may be affected without departing
10 from the spirit and scope of the novel concepts of the disclosure.

BRIEF DESCRIPTION OF THE DRAWINGS

The accompanying drawings illustrate one or more embodiments of the invention and, together with the written description, serve to explain the principles of the invention.
15 Wherever possible, the same reference numbers are used throughout the drawings to refer to the same or like elements of an embodiment.

FIG. 1 shows a PIVA system according to certain embodiments of the present invention.

FIG. 2 shows a flowchart of a method for detecting hypovolemia/hypervolemia of a
20 living subject according to certain embodiments of the present invention.

FIG. 3 shows charts of the peripheral venous waveforms and the Fourier transformation of the signals in states of hypovolemia (A), euvolemia (B), and hypervolemia (C) in the porcine model, according to certain embodiments of the present invention.

FIG. 4A shows a chart of the F_1 amplitude of the hemorrhage and transfusion of blood
25 in the porcine model for volume status according to certain embodiments of the present invention.

FIG. 4B shows a chart of mean arterial pressure (MAP), heart rate (HR), and shock index (SI) to the blood volume of the hemorrhage and transfusion of blood in the porcine model for volume status according to certain embodiments of the present invention.

FIG. 5A shows a chart of the F_1 amplitude of fluid administration to mild
30 hypervolemia in the porcine model for volume status according to certain embodiments of the present invention.

FIG. 5B shows a chart of MAP, HR, and SI to the blood volume of fluid administration to mild hypervolemia in the porcine model for volume status according to

certain embodiments of the present invention.

FIG. 6 shows the receiver operator curves (ROC) for detection of (A) hypovolemia (>200mL hemorrhage, 5.9%) and (B) hypervolemia (>200mL fluid administration, 5.9%) for the peripheral venous signal, HR, MAP, and SI according to certain embodiments of the

5 present invention.

FIG. 7 shows a table of patient demographics in a controlled human hemorrhagic model according to certain embodiments of the present invention.

FIG. 8 shows charts of the peripheral venous waveform and the Fourier transformation of the signals at baseline and after autologous blood removal in the controlled human hemorrhagic model according to certain embodiments of the present invention.

10

FIG. 9 shows charts of hemodynamic measurements for (A) the F_1 amplitude, (B) HR, (C) diastolic pulmonary artery pressure (dPAP) and (D) MAP in the controlled human hemorrhagic model at baseline and following hemorrhage at 250 mL and 500mL according to certain embodiments of the present invention.

15

FIG. 10A shows the ROC curves for detection of the F_1 amplitude, dPAP, HR and SI according to certain embodiments of the present invention.

FIG. 10B shows a table of the area under the curve (AUC), standard error (SE) and 95% confidence interval for the data as shown in FIG. 8A according to certain embodiments of the present invention.

20

FIG. 11A shows the F_1 amplitudes with (+PPV) and without (-PPV) positive pressure ventilation according to certain embodiments of the present invention.

FIG. 11B shows the F_0 amplitudes with (+PPV) and without (-PPV) positive pressure ventilation according to certain embodiments of the present invention.

25

FIG. 12 shows (A) PIVA signal and (B) shock index for detecting hemorrhage in a porcine animal model (n=8), according to certain embodiments of the present invention.

FIG. 13 shows (A) PIVA signal and (B) shock index for detecting hemorrhage in a porcine animal model (n=8), according to certain embodiments of the present invention.

DETAILED DESCRIPTION OF THE INVENTION

30

The invention will now be described more fully hereinafter with reference to the accompanying drawings, in which exemplary embodiments of the invention are shown. This invention may, however, be embodied in many different forms and should not be construed as limited to the embodiments set forth herein. Rather, these embodiments are provided so that this disclosure will be thorough and complete, and will fully convey the scope of the

invention to those skilled in the art. Like reference numerals refer to like elements throughout.

The terms used in this specification generally have their ordinary meanings in the art, within the context of the invention, and in the specific context where each term is used.

5 Certain terms that are used to describe the invention are discussed below, or elsewhere in the specification, to provide additional guidance to the practitioner regarding the description of the invention. For convenience, certain terms may be highlighted, for example using italics and/or quotation marks. The use of highlighting has no influence on the scope and meaning of a term; the scope and meaning of a term are the same, in the same context, whether or not
10 it is highlighted. It will be appreciated that the same thing can be said in more than one way. Consequently, alternative language and synonyms may be used for any one or more of the terms discussed herein, nor is any special significance to be placed upon whether or not a term is elaborated or discussed herein. Synonyms for certain terms are provided. A recital of one or more synonyms does not exclude the use of other synonyms. The use of examples
15 anywhere in this specification including examples of any terms discussed herein is illustrative only, and in no way limits the scope and meaning of the invention or of any exemplified term. Likewise, the invention is not limited to various embodiments given in this specification.

It will be understood that when an element is referred to as being “on” another
20 element, it can be directly on the other element or intervening elements may be present there between. In contrast, when an element is referred to as being “directly on” another element, there are no intervening elements present. As used herein, the term “and/or” includes any and all combinations of one or more of the associated listed items.

It will be understood that, although the terms first, second, third, etc. may be used
25 herein to describe various elements, components, regions, layers and/or sections, these elements, components, regions, layers and/or sections should not be limited by these terms. These terms are only used to distinguish one element, component, region, layer or section from another element, component, region, layer or section. Thus, a first element, component, region, layer or section discussed below could be termed a second element, component,
30 region, layer or section without departing from the teachings of the invention.

The terminology used herein is for the purpose of describing particular embodiments only and is not intended to be limiting of the invention. As used herein, the singular forms “a”, “an” and “the” are intended to include the plural forms as well, unless the context clearly indicates otherwise. It will be further understood that the terms “comprises” and/or

“comprising”, or “includes” and/or “including” or “has” and/or “having” when used in this specification specify the presence of stated features, regions, integers, steps, operations, elements, and/or components, but do not preclude the presence or addition of one or more other features, regions, integers, steps, operations, elements, components, and/or groups thereof.

Furthermore, relative terms, such as “lower” or “bottom” and “upper” or “top”, may be used herein to describe one element's relationship to another element as illustrated in the Figures. It will be understood that relative terms are intended to encompass different orientations of the device in addition to the orientation depicted in the Figures. For example, if the device in one of the figures is turned over, elements described as being on the “lower” side of other elements would then be oriented on “upper” sides of the other elements. The exemplary term “lower” can, therefore, encompass both an orientation of “lower” and “upper”, depending on the particular orientation of the figure. Similarly, if the device in one of the figures is turned over, elements described as “below” or “beneath” other elements would then be oriented “above” the other elements. The exemplary terms “below” or “beneath” can, therefore, encompass both an orientation of above and below.

Unless otherwise defined, all terms (including technical and scientific terms) used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. It will be further understood that terms, such as those defined in commonly used dictionaries, should be interpreted as having a meaning that is consistent with their meaning in the context of the relevant art and the present disclosure, and will not be interpreted in an idealized or overly formal sense unless expressly so defined herein.

As used herein, “around”, “about”, “substantially” or “approximately” shall generally mean within 20 percent, preferably within 10 percent, and more preferably within 5 percent of a given value or range. Numerical quantities given herein are approximate, meaning that the term “around”, “about”, “substantially” or “approximately” can be inferred if not expressly stated.

As used herein, the terms “comprise” or “comprising”, “include” or “including”, “carry” or “carrying”, “has/have” or “having”, “contain” or “containing”, “involve” or “involving” and the like are to be understood to be open-ended, *i.e.*, to mean including but not limited to.

As used herein, the term “hemodynamic” generally refers to blood movement, and “hemodynamic resuscitation” generally refers to increasing blood movement (or blood pressure) in a patient experiencing symptoms of compensated shock (e.g., based on a

“hemodynamic score” or “resuscitation score”).

As used herein, the term “peripheral intravenous waveform analysis” or its abbreviation “PIVA” refers to an analysis of the peripheral venous waveforms measured from a vein of a living subject through a standard peripheral intravenous (IV) catheter.

5 As used herein, the term “hypovolemia” refers to a medical condition of decreased blood volume, and more specifically a decrease in volume of blood plasma. In certain embodiments, hypovolemia stems from loss of blood volume due to hemorrhage, dehydration or intravascular water loss.

As used herein, the term “hypervolemia” refers to a medical condition of fluid
10 overload (i.e., having too much fluid) in the blood. In certain embodiments, hypervolemia stems from compromised regulatory mechanisms for sodium handling, such as congestive heart failure (CHF) or renal failure, or due to iatrogenic fluid administration.

OVERVIEW OF THE INVENTION

15 The ideal hemodynamic monitoring system for guiding fluid therapy would predict patient volume status accurately with minimal risks [36]. Unfortunately, existing dynamic systems and methods for assessing volume status, such as pulse pressure variation (PPV), stroke volume variation (SVV), and plethysmographic wave respiratory variation, may predict fluid responsiveness, but do not directly measure volume status, and they have not
20 been proven to detect iatrogenic volume overload during resuscitation [37, 38]. These existing non-invasive dynamic monitoring systems, such as SVV and PPV, rely on intrathoracic effects on left ventricular stroke volume during mechanical ventilation [10, 11]. Furthermore, these techniques depend on heart-lung interactions during mechanical ventilation for detection of hypovolemia. [37, 51] This critical limitation renders the
25 techniques ineffective in the spontaneously breathing patient [52].

In an effort to address the pitfalls associated with the existing central venous monitoring modalities, peripheral venous pressure (PVP) monitoring has been increasingly explored as an alternative for determining intravascular volume status. There is a good correlation between trends in absolute PVP and central venous pressure (CVP) measurements
30 in critically ill patients [13] [14]. However, even trends of CVP, and therefore PVP, are not reliable indicators of volume status, resulting in a need to modify this approach. One such method, cuff-occlusion rate of rise of PVP (CORRP), has been shown to correlate with volume status in critically-ill patients, but is limited by operator-dependent, non-continuous measurements to assess volume status [15]. Another method used peripheral venous

waveform spectral analysis for continuous beat-to-beat monitoring of intravascular volume status to quantify changes associated with lower body negative pressure, a simulation of hemorrhage, in spontaneously breathing subjects [16]. However, lower body negative pressure results in vasodilation and may not truly represent the physiological responses to hemorrhage. In other words, there is no existing non-invasive method of accurately assessing patient blood volume status in intrathoracic pressure changes without ventilation-induced or negative pressure. The current gold-standard measurement requires either echocardiography or a pulmonary artery catheter, and both of these measurements have wide variability in their accuracy.

Recently, *Sileshi et al.* reported that peripheral intravenous waveform analysis (PIVA) can detect early Stage 1 hemorrhage, during perioperative autologous blood donation in patients undergoing cardiac surgery [17]. Moreover, this study demonstrated that PIVA was more sensitive to acute changes in circulating blood volume than standard vital signs and invasive pulmonary arterial monitoring.

Moreover, since not all wounded soldiers require endotracheal intubation and positive pressure ventilation, there is an acute need for accurate and sensitive volume status monitoring in wounded, spontaneously breathing soldiers. There is a need for a point of care monitor that detect small direct changes in volume status and warns of impending hemodynamic collapse in the combat casualty care setting.

Accordingly, aspects of the present invention relates to systems and methods of detecting early stage hemorrhage using PIVA to assess blood volume status of a living subject, which may include human beings and/or other animals, and applications of the same.

FIG. 1 shows a PIVA system according to certain embodiments of the present invention. As shown in FIG. 1, the PIVA system 100 includes: a peripheral IV device 110 and a processing device 120. The processing device 120 is communicatively connected to the peripheral IV device 110. In certain embodiments, the connection between the peripheral IV device 110 and the processing device 120 may be through a network, which may be implemented by a wired connection or a wireless connection. Examples of the network may include without being limited to, a local area network (LAN), a wide area network (WAN), the Internet, or any other types of network.

The peripheral IV device 110 is configured to acquire, from a vein of a living subject 130, peripheral venous signals. In certain embodiments, the living subject may be a human being, or may be other animals. In one embodiment, the living subject may be a human patient, and the patient may be awake and spontaneously breathing/moving, or may be

induced with anesthesia such that the patient is not spontaneously moving. In certain embodiments, the peripheral IV device 110 may include a peripheral IV catheter being inserted into the vein of the living subject 130, and a monitoring device connected to the peripheral IV catheter. The monitoring device is configured to capture and record the peripheral venous signals from the peripheral IV catheter at a sampling rate. In certain embodiments, the monitoring device may include a pressure transducer directly connected to the peripheral IV catheter, such that the peripheral venous signals are captured and recorded by the pressure transducer.

The processing device 120 is configured to: receive the peripheral venous signals from the peripheral IV monitoring device; perform a spectral process and analysis on the peripheral venous signal to obtain a peripheral venous pressure frequency spectrum; and perform a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum to determine the blood volume status of the living subject in real time. In certain embodiments, the processing device 120 may be a computing device, which may be a desktop computer, a laptop computer, a smartphone, a tablet device, or any other computing devices with processors to perform the processing functions. In certain embodiments, the spectral analysis may be a spectral fast Fourier transform (FFT) analysis.

In certain embodiments, the PIVA system 100 further includes: a pump (not shown) connected to the living subject to perform liquid exchange to the living subject; and a pump controlling mechanism (not shown) communicatively connected to the processing device, configured to control the pump by intermittently pausing the pump or subtract the pump signal when the peripheral IV device acquires the peripheral venous signals, and restarting the pump when the peripheral IV device does not acquire the peripheral venous signals. In certain embodiments, the pump may be a dialysis pump, a cardiopulmonary bypass pump, an extracorporeal membrane oxygenation (ECMO), or an infusion pump. In this case, the processing device 120 is further configured to send a signal to the pump controlling mechanism to notify the pump controlling mechanism to control the pump.

FIG. 2 shows a flowchart of a method for detecting hypovolemia/hypervolemia of a living subject according to certain embodiments of the present invention. As shown in FIG. 2, at step S210, the peripheral IV device 110 acquires the peripheral venous signals from the vein of the living subject. At step S220, upon receiving the peripheral venous signals from the peripheral IV device 110, the processing device 120 performs a spectral process and analysis, such as the spectral FFT analysis, on the peripheral venous signal to obtain a peripheral venous pressure frequency spectrum. At step S230, the processing device 120

performs a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum to determine the blood volume status of the living subject in real time. At step S240, the processing device 120 determines whether a significant amplitude change of the peaks is detected. If so, at step S250, the processing device 120 determines that the

5 living subject has hypovolemia or hypervolemia, depending on the amplitude change. If not, at step S260, the processing device 120 determines that the living subject has no hypovolemia or hypervolemia.

Specifically, the steps S210 and S220 may be performed continuously, such that at two different time period, two sets of the peripheral venous pressure frequency spectrums

10 may be obtained. For example, for a time period from T_0 to T_2 , the time period may be divided into a first time period from T_0 to T_1 , and a second time period from T_1 to T_2 , and each of the first time period and the second time period may be used to obtain a separate set of peripheral venous pressure frequency spectrums. In certain embodiments, the time period may be divided into more than two time periods, and multiple sets of peripheral venous

15 pressure frequency spectrums may be obtained. In certain embodiments, the peripheral venous pressure frequency spectrum obtained at an earlier time may be used as a baseline peripheral venous pressure frequency spectrum. Thus, the statistical analysis at step S230 may be performed by obtaining a plurality of baseline peaks $\{B_{N-1}\}$ from a lower frequency side on a baseline peripheral venous pressure frequency spectrum, where N is a positive

20 integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to a plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} . In other words, the baseline peaks may include a first baseline peak B_0 corresponding to a first frequency F_0 , a second baseline peak B_1 corresponding to a second frequency F_1 , a third baseline peak B_2 corresponding to a third

25 frequency F_2 ..., and the second frequency F_1 is greater than the first frequency F_0 . Then, a plurality of peaks $\{P_{N-1}\}$ may be obtained on the peripheral venous pressure frequency spectrum currently obtained, where the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$. For example, the peaks may include a first peak P_0 corresponding to the first frequency F_0 , a

30 second peak P_1 corresponding to the second frequency F_1 , a third peak P_2 corresponding to the third frequency F_2 In certain embodiments, the number of peaks on the peripheral venous pressure frequency spectrum equals to the number of baseline peaks on the baseline peripheral venous pressure frequency spectrum. In this way, the intravascular volume status of the living subject may be determined in real time by comparing the amplitudes of the

peaks to that of the corresponding baseline peaks, respectively.

In certain embodiments, the intravascular volume status of the living subject at the second time period indicates hypovolemia when amplitude decreases are detected greater than a first threshold from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$. In certain

5 embodiments, the intravascular volume status of the living subject at the second time period indicates hypervolemia when amplitude increases greater than a second threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$. For example, if there is a significant decrease in the F_1 amplitude of the second peak of the peripheral venous pressure frequency spectrum that reaches the first threshold, the processing device 120 will determine

10 that hypovolemia (e.g., hemorrhage or intravascular water loss, etc.) has occurred in the living subject. On the other hand, if there is a significant increase in the F_1 amplitude of the second peak of the peripheral venous pressure frequency spectrum that reaches the second threshold, the processing device 120 will determine that hypervolemia (e.g., congestive heart failure, renal failure, iatrogenic fluid administration, etc.) has occurred in the living subject.

15 It should be noted that, in addition to the F_1 amplitude of the second peak P_1 , the amplitude of other peaks, such as the first peak P_0 , the third peak P_2 , and any subsequent peaks, may also be used to determine hypovolemia or hypervolemia of the living subject.

In certain embodiments, the method is performed to the living subject during resuscitation of the living subject. In one embodiment, the intravascular volume status of the

20 living subject at the second time period indicates a return of euolemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at the first time period, and amplitude increases greater than a third threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$. In one embodiment, the intravascular volume status of the living subject at the second time period indicates over-resuscitation when the living subject is

25 determined to be in an euolemic state at the first time period, and amplitude increases greater than a fourth threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$.

In certain embodiments, the actual volume calculation may be performed based on a large logic table, with interpolation, that will be generated based on a population study and

30 can be further fit to the specific patient in a chronic-use device. An example of the logic table is provided as follows.

Example Logic Table

H1- amplitude	H2- amplitude	H3- amplitude	Volume Status Ratio
0.05	0.023	0.018	1.1
0.071	0.035	0.025	1.1
0.119	0.652	0.017	0.8

Further, in certain embodiments, a regression model may be provided to calculate the volume status ratio. For example:

$$\text{Volume Status Ratio} = b_0 + b_1 \cdot p_1 + b_2 \cdot p_2 + \dots + b_N \cdot p_N$$

- 5 where $b_0, b_1, b_2, \dots, b_N$ are linear regression coefficients determined in a multivariate regression model for each independent parameter p_i . In certain embodiments, the parameters may include:

1. Amplitudes (or powers) of each harmonic in the frequency domain
2. Noise floor amplitude
- 10 3. HR
4. Time domain amplitude of peripheral venous pressure

In certain embodiments, the method may be capable of detecting at least 6% of blood loss of the living subject.

- In certain embodiments, the system 100 may be implemented by an instrumented IV
- 15 catheter interfaced with a smart phone device for early hemorrhage detection and guided fluid therapy to improve survival in the casualty care setting. The device uses: (1) A ruggedized miniature pressure transducer integrated into an IV catheter, (2) real-time algorithm for early hemorrhage detection and guided goal-directed fluid resuscitation (3) wireless immediate data transfer for remote or centralized monitoring and intervention. In certain embodiments,
- 20 the device may involve a portable, lightweight (<50 grams) sensor that attaches directly to a standard peripheral IV catheter interfaced to a Bluetooth radio for transmission to a Smartphone device. The form factor of the device will be comparable to a USB stick. The cost of the device may be USD <\$25 making it feasible to integrate the device into every IV catheter used in casualty care and trauma settings. Further, the device may utilize immediate
- 25 data transfer and storage to a secure database and mobile application for remote Damage Control Resuscitation and emergency telemedicine. In addition to immediate graphic display

for field medics, the smart IV will have Bluetooth technology allowing data transfer to a Smart phone mobile application. Bidirectional data transfer and feedback from remote clinical practitioners provides emergency telemedicine assistance to field medics in the battlefield. Further, physiologically data can be uploaded to a server, providing an electronic medical record for all care providers throughout the resuscitation period. Such device may be potentially relevant in military use, as the military needs a robust point-of-care method for early hemorrhage detection in the wounded soldier. This device will be rugged, wireless, lightweight and portable with minimal, self-contained energy requirements, conducive to austere environments.

As described above, the PIVA system 100 includes the processing device 120, which serves as a controller device that is configured to receive the peripheral venous signals from the monitoring device of the peripheral IV device 110. The processing device 120 may then perform the Fourier transform on the peripheral venous signals of the time domain data to process the signal to obtain a spectrum. Once the fourier spectrum is generated, the second peak at the frequency F_1 that correspond to the heart rate or another hemodynamic parameter's amplitude or power is measured and each resonant frequency of that hemodynamic parameter's amplitude or power is also measured. The amplitudes and/or powers are input into the algorithm that weights each resonant frequency and outputs a measurement of a hemodynamic parameter. Additional inputs may also be required, including but not limited to the following: age, weight, gender, and height, these variables may be input into the algorithm to determine a more accurate depiction of the patient's hemodynamic state. Based on the Fourier transform of the piezoelectric signal the processor is used to generate a resuscitation score based on the hemodynamic parameter.

The invention relates to methods for peripheral venous pressure analysis algorithm that uses spectral analysis to estimate intravascular volume status, and its applications. In certain aspects, the invention recites, among other things:

- 1) Harmonic peripheral venous pressure waveform analysis algorithm.
- 2) Method of measuring peripheral venous pressure frequency spectra for determination of real-time volume status.
- 3) A venous pressure monitor algorithm that can distinguish between euvoolemia and hypervolemia (all current technologies stop at euvoolemia).
- 4) A method for assessing volume status in a spontaneously breathing patient.
- 5) A volume status monitor that uses a peripheral IV.
- 6) A closed loop system for controlling volume status with a peripheral venous

pressure monitor and intravenous fluid pump.

In the following examples, the inventors have utilized PIVA in different models, including a porcine hemorrhage-resuscitation model and a controlled human hemorrhagic model, to analyze and study dynamic volume changes and shifts in the peripheral venous waveforms. The tests in the examples are performed in standardized settings in order to test the hypothesis that PIVA is more sensitive and specific than standard and invasive monitors for detecting and quantitating acute hemorrhage, as well as resuscitation to euvolemia and iatrogenic fluid overload. Further, the accuracy and linearity of the PIVA estimation of blood volume has been evaluated as compared to absolute volume removal and administration.

EXAMPLE 1

TEST IN PORCINE HEMORRHAGE-RESUSCITATION MODEL

In this example, a test has been performed using PIVA in a porcine hemorrhage-resuscitation model to study dynamic volume shifts in a standardized setting. Under an approved Institutional Animal Care and Use Committee protocol, 8 adult Yorkshire pigs, each weighing 45+/-0.8 kg, were monitored non-invasively with a noninvasive blood pressure cuff, 5-lead electrocardiogram, and pulse oximeter (SurgiVet, Norwell, Boston, MA). Each animal was induced with general anesthesia Telazol 2 mg, Ketamine 50 mg, and Xylazine 2 mg given via an ear vein. After intubation with a cuffed 5.0 ID endotracheal tube the pigs were ventilated with a volume controlled ventilator (Hallowell EMC, MA, USA) with a volume-controlled mode of 8mL/kg tidal volume with a positive end-expiratory pressure of 5 cm H₂O, I:E ratio 1:2, and a FiO₂ 1.0. Respiratory rate (16-22 breaths/minute) was titrated to maintain an end-tidal CO₂ of 35-40 mmHg.

Anesthesia was maintained with 1% isoflurane (Primal Healthcare, Boston, MA). Surgical exposure of the femoral artery and vein was obtained. A 6Fr catheter (Mila International, Erlanger, KY) was inserted directly into the femoral vein and sutured in place. A 20g angiocatheter (Mila International, Erlanger, KY) was placed in the femoral artery for continuous blood pressure measurements. A 20g peripheral IV (Smiths Medical, Dublin, OH) was inserted into the front extremity of each animal. Details of the test are provided as follows:

Peripheral Venous Waveform Acquisition

After induction of anesthesia, a 20 gauge peripheral intravenous (IV) catheter was inserted into an upper extremity vein and directly connected to a pressure transducer

(ADInstruments, Colorado Springs, CO) via standard high- pressure tubing. The IV catheter was dedicated to obtaining venous waveforms and was not used for infusion of fluids or drug delivery during the test. Peripheral venous waveform tracings were recorded continuously throughout the procedure and analyzed on LabChart (ADInstruments, Colorado Springs, CO). Peripheral intravenous waveform data was captured at 1 kHz to allow adequate sampling to perform spectral analysis of the waveform data.

Hemorrhage Protocol

After induction of anesthesia, baseline measurements of the peripheral venous signal were obtained. Standard IV tubing was attached proximally to the femoral venous line and distally to a sterile collection bag, producing a closed system. Intravenous Heparin 10 units/kg was administered immediately prior to hemorrhage. Venous blood was removed via gravity in a stepwise fashion of 50 mL per minute over a 10-minute period. Vital signs including heart rate, arterial blood pressure, and peripheral venous waveforms were continuously recorded throughout the procedure. The procedure was terminated at 400 mL blood removal or mean arterial pressure (MAP) < 40mmHg.

Volume Resuscitation

Immediately following the hemorrhage protocol, the entire autologous blood was returned to the pig at a continuous rate of 50 mL/min over 10 minutes through the femoral venous catheter. Vital signs were continuously recorded throughout this process. Following return of the autologous blood, each pig was considered to be in an euvoletic state.

Iatrogenic Volume Overload

Following volume resuscitation to a euvoletic state, warm (40 °C) 500 mL Plasmalyte (Baxter International, Deerfield, IL) balanced crystalloid solution was administered through the femoral venous catheter over a 10-minute period. Continuous hemodynamic monitoring was performed as previously described.

Spectral Analysis

The spectral fast Fourier transform (FFT) analysis of peripheral venous pressure was performed using an 8K sampling window with no window overlap. Data was recorded at a sampling rate of 1 kHz necessitating 8 seconds of continuous time-domain signal to perform the 8K-FFT spectral analysis. Following Fourier transformation, the amplitude of each

frequency peak was calculated in LabChart. The peak amplitude associated with the heart rate was used to acquire F_1 (the fundamental frequency) amplitude of the signal. Data was captured in triplicate for each point used in analysis.

5 Results

All 8 pigs successfully underwent hemorrhage, resuscitation, and volume overload without complications. Conversion of the time domain signal into the frequency domain was performed to determine the amplitude of each of the fundamental frequency recorded during the experiment.

10 FIG. 3 shows charts of the peripheral venous waveforms and the Fourier transformation of the signals in states of hypovolemia (A), euvolemia (B), and hypervolemia (C) in the porcine model, according to certain embodiments of the present invention. As shown in FIG. 3, the top panels of each section (A), (B) and (C) show the peripheral waveforms recorded from LabChart in the time domain, and the bottom panels of each
15 section show the Fourier transformation. Specifically, in the Fourier transformation charts, there are distinct peaks at frequencies F_0 and F_1 , where F_0 represents the porcine's respiratory rate and its contribution to the signal, and F_1 represents the porcine's heart rate. The amplitude of the signal was correlated directly with the volume status.

20 Hemorrhage and Autologous Blood Transfusion

FIG. 4A shows a chart of the F_1 amplitude of the hemorrhage and transfusion of blood in the porcine model for volume status according to certain embodiments of the present invention. As shown in FIG. 4A, blood loss of 200, 300, and 400 mL of crystalloid created significant differences in the F_1 amplitude ($P < 0.05$). When measuring the F_1
25 amplitude alone, both the hemorrhage and autologous blood transfusion resulted in significantly non-zero slopes of -0.000223 and -0.000242 and with P values of 0.0049 and 0.0008, respectively. There was a significant change in F_1 amplitude following 200 mL ($P < 0.01$), 300 mL ($P < 0.001$), and 400 mL ($P < 0.001$) of blood loss. There was also a significant incremental change in F_1 between 100 and 400 mL blood loss.

30 FIG. 4B shows a chart of mean arterial pressure (MAP), heart rate (HR), and shock index (SI) to the blood volume of the hemorrhage and transfusion of blood in the porcine model for volume status according to certain embodiments of the present invention. Specifically, FIG. 4B shows the arterial pressure, heart rate, and shock index for the pigs over a hemorrhage of 0 to 400 mL. As shown in FIG. 4B, there were no significant differences in

the heart rate or the blood pressure after 400 mL blood loss (11.8% estimated blood volume), and the shock index did not significantly change until 300 mL blood loss (8.85% estimated blood volume) ($p < 0.05$).

Taken together, these results show that changes in F_1 amplitude is a more sensitive
 5 measure for hemorrhage and transfusion of blood than HR, MAP, or SI.

Iatrogenic volume overload

FIG. 5A shows a chart of the F_1 amplitude of fluid administration to mild
 hypervolemia in the porcine model for volume status according to certain embodiments of the
 10 present invention. Specifically, FIG. 5A shows the F_1 amplitude and how it is affected by
 additional IV fluid being given. As shown in FIG. 5A, the fluid administration of crystalloid
 beyond the euvolumic state resulted in significant increase in the F_1 amplitude of the signal.
 Specifically, the F_1 amplitude changed with fluid administration resulting in a significantly
 nonzero slope ($p = 0.0017$). Additions of 200, 300, and 400 mL of crystalloid created
 15 significant differences in the F_1 amplitude ($P < 0.05$), between increments of 0 and 200
 ($P < 0.05$), 0 and 300 ($P < 0.05$), and 0 and 400 ($P < 0.01$) mL of crystalloid.

FIG. 5B shows a chart of MAP, HR, and SI to the blood volume of fluid
 administration to mild hypervolemia in the porcine model for volume status according to
 certain embodiments of the present invention. Specifically, FIG. 5B shows the HR, MAP and
 20 SI for the pigs over volume status ranging from 0 to +400mL. As shown in FIG. 5B, there
 were no significant changes in the heart rate and MAP with volume overload. Shock index
 changed significantly between 0 and 300 mL ($P < 0.05$) of fluid administration and 0 and 400
 mL of fluid administration ($P < 0.05$).

Taken together, these results show that changes in F_1 amplitude is a more sensitive
 25 measure for hypervolemia than HR, MAP, or SI.

Further, HR, MAP, SI and F_1 amplitude were measured during hemorrhage and fluid
 administration to the porcine volume status model. FIG. 6 shows the receiver operator curves
 (ROC) for detection of (A) hypovolemia (>200mL hemorrhage, 5.9%) and (B) hypervolemia
 (>200mL fluid administration, 5.9%) for the peripheral venous signal, HR, MAP, and SI
 30 according to certain embodiments of the present invention. As shown in FIG. 6,
 measurement of F_1 amplitude showed improved sensitivity and specificity for detection of
 hypovolemia and hypervolemia in the porcine model. For the detection of hypovolemia as
 shown in FIG. 6(A), the F_1 amplitude generated an ROC curve with an area under the curve
 (AUC) of 0.93, HR generated an ROC curve with an AUC of 0.61, MAP generated an ROC

curve with an AUC of 0.48, and SI generated an ROC curve with an AUC of 0.72. For the detection of hypervolemia as shown in FIG. 6(B), the F_1 amplitude generated an ROC curve with an area under the curve (AUC) of 0.85, HR generated an ROC curve with an AUC of 0.62, MAP generated an ROC curve with an AUC of 0.63, and SI generated an ROC curve with an AUC of 0.65. In sum, MAP demonstrated the weakest ROC curves for the hypovolemia, and HR was the least sensitive at detecting hypervolemia in the porcine model. In all situations the F_1 amplitude obtained by PIVA had the greatest sensitivity and specificity at detecting volume status.

The test results are in agreement with findings of Sileshi, et al. which demonstrated hemorrhage detection after approximately ~6% estimated blood loss in a cardiac surgery population [31]. The ROCs illustrate that PIVA had a significantly greater sensitivity and specificity for detecting hemorrhage compared to HR, MAP, and SI, as shown in FIG. 2B. Further, PIVA is measured independently of intrathoracic pressure changes produced by mechanical ventilation and may be able to fill a void in dynamic monitoring.[16, 17]

Further, the significant changes in the F_1 amplitude has been demonstrated not only during hemorrhage, but also a return of the F_1 amplitude to baseline with transfusion of a matched quantity of autologous blood during resuscitation, as shown in FIGS. 4A and 4B. Detecting a return to euolemia has great potential to enhance accurate goal directed volume resuscitation to prevent fluid overload in the hemorrhaging patient.

In addition, the test simulated iatrogenic volume overload with balanced crystalloid, an inadvertent, yet common event during acute resuscitation. PIVA detected iatrogenic volume overload following only 200mL (5.9% estimated total blood volume) of crystalloid administration beyond the euolemic state, as shown in FIGS. 5A and 5B. ROC showed PIVA was significantly more sensitive and specific in detecting hypervolemia as compared to SI, HR, and MAP, as shown in FIG. 6. With fluid administration beyond the euolemic state, the F_1 amplitude significantly increases, which provides a novel method for detecting and quantifying fluid overload. This is likely because the additional fluid administration increased wall tension in the vein, thereby increasing the fundamental frequency amplitude.

EXAMPLE 2

TEST IN CONTROLLED HUMAN HEMORRHAGIC MODEL

In this example, a test has been performed using PIVA in a controlled human hemorrhagic model to analyze dynamic changes in the peripheral venous waveforms to assess volume status. The test was approved by the Vanderbilt University Institutional Review

Board (IRB), and informed written consent was obtained preoperatively in select patients scheduled for elective cardiac surgery. Any patient undergoing elective cardiac surgery met the inclusion criteria. Patients with a history of moderate or severe right ventricular dysfunction, severe anemia (hemoglobin <8 g/dl) or patients who presented with arrhythmias or hemodynamic instability were excluded. A total of 12 patients were studied.

Anesthesia and Mechanical Ventilation

All patients were induced with an opiate and propofol and received non-depolarizing neuromuscular blockade so that there was no evidence of spontaneous respirations. All patients were intubated with an endotracheal tube and received mechanical ventilation with tidal volumes of 7 to 9 mL/kg, positive end-expiratory pressure of 4 mmHg, and a fraction of inspired oxygen of 1.0. Patients were maintained with 0.8 to 1 MAC of isoflurane. In five patients, mechanical ventilation was halted for brief intervals (<10 seconds) for reasons unrelated to this protocol, such as surgeon's request. This opportunity was used to evaluate the effects of mechanical ventilation on the peripheral venous signal. Comparisons have been conducted between the peripheral venous waveform tracings during breath holding and immediately after mechanical ventilation resumed. The ventilator circuit remained connected to the endotracheal tube throughout this period.

Hemodynamics

All patients were monitored with standard non-invasive monitors including 5-lead electrocardiogram, non-invasive blood pressure cuff, and pulse oximetry. Prior to induction of anesthesia, a 20g arterial catheter (Arrow International, Reading, PA) was inserted into the radial artery. Following induction of anesthesia, all patients received a right 9Fr central catheter (Cook Critical Care, Bloomington, IN), and a pulmonary artery catheter (Edwards Lifesciences, Irvine, CA). To minimize complications, the pulmonary artery catheter was not placed in the wedge position. The pulmonary artery diastolic pressure was used as a surrogate for pulmonary artery occlusion pressure. All pressure transducers were placed at the level of the right atrium and zeroed to atmospheric pressure.

Peripheral Venous Waveform Acquisition:

After induction of anesthesia, an 18 or 16 gauge peripheral intravenous catheter (IV) was inserted into an upper extremity vein and directly connected to a pressure transducer (ADInstruments, Colorado Springs, CO) via standard high- pressure tubing. The IV catheter

was dedicated to obtaining venous waveforms and was not used for infusion of fluids or drug delivery during the study. Peripheral venous waveform tracings were recorded continuously throughout the procedure and analyzed on LabChart (ADInstruments, Colorado Springs, CO). Peripheral intravenous waveform data was captured at 1kHz for adequate sampling to perform spectral analysis on the data.

Hemorrhage Protocol

Following induction of anesthesia and central line placement, the hemorrhage protocol was initiated. Standard IV tubing was attached proximally to the central venous catheter and distally to a citrate bag, producing a closed system. Venous blood was removed from the central venous catheter via gravity over a 10-minute period. Up to 500ml or 10% of blood volume was removed from each patient. All were successfully completed within 10 minutes without protocol interruption. Vital signs including heart rate, arterial blood pressure, central venous pressure, pulmonary artery pressure, and PVA were continuously recorded throughout the procedure.

Spectral Analysis

The spectral FFT analysis of peripheral venous pressure was performed using an 8K sampling window with no window overlap. Since data was recorded at a sampling rate of 1 kHz, 8 seconds of continuous time-domain signal was required to perform the 8K-FFT spectral analysis. Once the data is transformed into the frequency domain the amplitude of each frequency is calculated in LabChart. The lowest frequency peak is associated with the patient's respiratory rate (F_0) followed by the patient's heart rate (F_1). Data was captured in triplicate for each point used in analysis. Peripheral waveform is sensitive to patient movement and electrocautery signal. To minimize signal noise, data was captured during periods of minimal patient movement and no electrocautery use.

Statistical analysis:

Data for physiologic measurements and their associated blood loss was analyzed using a one-way ANOVA analysis with a post-test of Tukey's multiple comparison with paired analysis. Data was also entered as baseline measurements and blood loss of 500 mL in patients into MedCalc to determine a pairwise comparison of ROC curves. Area under the curve, standard error, and a 95% confidence interval for each data set was acquired. ROC curves were plotted on the same graph to easily identify differences between the sensitivity

and specificity of physiologic parameters. To determine the effect of positive pressure ventilation, F_0 and F_1 values were compared using a paired student's t-test with values of $P < 0.05$ being considered statistically significant. Statistical analysis was performed using GraphPad Prism and MedCalc.

5 FIG. 7 shows a table of patient demographics in a controlled human hemorrhagic model according to certain embodiments of the present invention. As shown in FIG. 7, twelve patients who underwent elective cardiac surgery were enrolled in this test. Mean age of the patients was 65.5 years, with 9 (75%) being male. The patients underwent variety of cardiac surgical procedures with 6 of 12 (50%) of patients presenting for mitral valve
10 repair/replacement for mitral regurgitation. All patients with the exception of one had normal right ventricular function. Seven patients (58%) presented with a normal left ventricular ejection fraction (LVEF); two patients (16%) had a mild LVEF depression; two patients (16%) had a moderate LVEF depression; and one (8%) patient had a severely depressed LVEF of 15-20%. One patient had severe pulmonary hypertension from chronic pulmonary
15 emboli. In 10 patients, autologous blood donation was completed prior to sternal incision. In four patients, autologous blood was removed following a right thoracotomy for minimally invasive mitral surgery. These patients remained on two-lung ventilation throughout the protocol.

 FIG. 8 shows charts of the peripheral venous waveform and the Fourier
20 transformation of the signals at baseline and after autologous blood removal in the controlled human hemorrhagic model according to certain embodiments of the present invention. As shown in FIG. 8, the top panels show the peripheral waveforms, and the bottom panels show the corresponding Fourier transformation of data at baseline and after blood removal. There was a notable amplitude decrease in F_0 , F_1 , and higher frequencies following hemorrhage.

25 FIG. 9 shows charts of hemodynamic measurements for (A) the F_1 amplitude, (B) HR, (C) diastolic pulmonary artery pressure (dPAP) and (D) MAP in the controlled human hemorrhagic model at baseline and following hemorrhage at 250 mL and 500mL according to certain embodiments of the present invention. Specifically, FIG. 9 is provided for a comparison of different indices of volume status during blood loss. As shown in FIG. 9,
30 there were statistically significant changes in the F_1 amplitude with blood removal between baseline (0mL) and 250mL ($p=0.0019$), baseline and 500mL ($p=0.0042$), and 250mL to 500mL ($p=0.0382$). MAP, HR, and dPAP did not significantly change between baseline and 250mL or 500mL blood volume removal.

 FIG. 10A shows the ROC curves for detection of the F_1 amplitude, dPAP, HR and SI

according to certain embodiments of the present invention, and FIG. 10B shows a table of the area under the curve (AUC), standard error (SE) and 95% confidence interval for the data as shown in FIG. 10A according to certain embodiments of the present invention. As shown in FIG. 8A, at 6% estimated blood loss, F_1 had the greatest area under the ROC curve (AUC = 0.90, 95%CI [0.67, 0.99]), compared to dPAP (AUC = 0.62), heart rate (AUC = 0.54) and MAP (AUC = 0.512).

In order to demonstrate independence of F_1 signal from effects of positive pressure ventilation, the F_1 and F_0 amplitudes are recorded with and without mechanical ventilation. FIG. 11A shows the F_1 amplitudes with (+PPV) and without (-PPV) positive pressure ventilation according to certain embodiments of the present invention, and FIG. 11B shows the F_0 amplitudes with (+PPV) and without (-PPV) positive pressure ventilation according to certain embodiments of the present invention. As shown in FIG. 11A, there was no significant change in the F_1 amplitude with and without mechanical ventilation ($n=5$, $P=0.21$). In comparison, the F_0 amplitude, which represents the respiratory frequency as reflected by changes in the peripheral venous waveform, was significantly greater in ventilated patients compared to non-ventilated patients ($n=5$, $P=0.0059$), as shown in FIG. 11B.

Based on the test results, PIVA is able to detect as little as 6% blood volume loss, and the signal was independent of mechanical ventilation. This amount of blood loss is well within the definition of Stage I or subclinical hemorrhage. In contrast, heart rate, MAP, CVP, and dPAP were poor predictors of early stage hemorrhage, a finding that is consistent with previously published studies [39] [6] [40].

EXAMPLE 3

ADDITIONAL TESTS

As shown in the previous examples, the inventors have found that venous waveform analysis overcomes many critical barriers associated with arterial-based monitoring. The inventors discovered and confirmed with tests that peripheral intravenous waveform analysis (PIVA) obtained via a pressure transducer in a standard intravenous catheter detects hemorrhage in humans and porcine models.

The inventors have conducted additional tests on PIVA detections of blood loss in pigs: In the test, the PIVA device is applied to intubated and sedated pigs ($n=4$). All pigs were monitored in real time with intra-arterial blood pressure, heart rate, pulse oximeter, and a 5-lead electrocardiogram. The PIVA device was interfaced with LabChart (ADInstruments,

Colorado Springs, CO, USA) software for continuous, real-time data collection. Up to 15% of blood volume was incrementally removed during a 20-minute period.

FIG. 12 shows (A) PIVA signal and (B) shock index for detecting hemorrhage in a porcine animal model (n=8), according to certain embodiments of the present invention. As shown in FIG. 12, PIVA waveform data analyzed with Fourier transformation techniques was found to be more sensitive than invasive arterial blood pressure, heart rate, and even shock index for detecting hemorrhage and determining fluid status.

Fluid overload is responsible for increased mortality in injured patients. However, standard vital sign monitoring fails to detect thresholds for euvoolemia and hypervolemia, contributing to unnecessary and potentially harmful over-resuscitation. We studied a porcine animal resuscitation model to determine intravascular volume status during fluid administration. Animals were considered euvolemic upon presentation based on weight and hemodynamic stability. After completion of the hemorrhagic stage, pigs (n=8) underwent fluid resuscitation. Autologous blood was re-administered followed by an additional 10 mL/kg of balanced crystalloid solution over 20 minutes.

FIG. 13 shows (A) PIVA signal and (B) shock index for detecting hemorrhage in a porcine animal model (n=8), according to certain embodiments of the present invention. As shown in FIG. 13, PIVA continues to detect intravascular changes beyond the euvolemic state during fluid administration in a porcine model n=8 pigs. Notably, heart rate and blood pressure remained unchanged even after 10 mL/kg of crystalloid administration beyond the euvolemic state (data not shown). It is shown from these data that hypervolemia detection with PIVA is superior to standard and invasive monitoring for guiding resuscitation. Development of an algorithm for detecting fluid overload is an extension of the method. In addition to resuscitation, these methods may be useful for managing patients with congestive heart failure.

In summary, the ability of PIVA to detect subclinical hemorrhage and goal-directed fluid therapy offers significant advantages over standard and possibly dynamic monitoring modalities. Further, PIVA is minimally invasive, requires minimal training, and it averts infectious and major vascular complications associated with central venous and pulmonary artery catheters. The PIVA signal is independent of effects of mechanical ventilation and may be useful in patients receiving lung protective ventilator strategies and possibly in spontaneously breathing individuals. PIVA has the potential to provide a low-cost, minimally invasive method for detecting and quantitating subclinical hemorrhage. As shown in the examples, PIVA is more sensitive than standard and invasive monitoring for detecting

subclinical hemorrhage, and may provide a powerful alternative to central venous catheterization. Further, the method is independent of effects of mechanical ventilation, a potential advancement for monitoring spontaneously breathing patients.

The foregoing description of the exemplary embodiments of the invention has been presented only for the purposes of illustration and description and is not intended to be exhaustive or to limit the invention to the precise forms disclosed. Many modifications and variations are possible in light of the above teaching.

The embodiments were chosen and described in order to explain the principles of the invention and their practical application so as to enable others skilled in the art to utilize the invention and various embodiments and with various modifications as are suited to the particular use contemplated. Alternative embodiments will become apparent to those skilled in the art to which the present invention pertains without departing from its spirit and scope. Accordingly, the scope of the present invention is defined by the appended claims rather than the foregoing description and the exemplary embodiments described therein.

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What is claimed is:

1. A method for determining, by a peripheral intravenous (IV) waveform analysis (PIVA) system, hypovolemia, hypervolemia and vascular tone of a living subject based on an intravascular volume status of the living subject, the PIVA system including a peripheral IV device and a processing device and the method comprising:

acquiring, by the peripheral IV device, continuously for a time period from T_0 to T_2 , peripheral venous signals from a vein of the living subject, wherein the time period is divided into a first time period from T_0 to T_1 , and a second time period from T_1 to T_2 ;

processing, by the processing device, the peripheral venous signals acquired at the first time period to obtain a baseline peripheral venous pressure frequency spectrum;

obtaining, by the processing device, a plurality of baseline peaks $\{B_{N-1}\}$ on the baseline peripheral venous pressure frequency spectrum, wherein N is a positive integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to a plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} ;

processing, by the processing device, the peripheral venous signals acquired at the second time period to obtain a peripheral venous pressure frequency spectrum;

obtaining, by the processing device, a plurality of peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum, wherein the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$; and

determining, by the processing device, the intravascular volume status of the living subject at the second time period by comparing amplitudes of the peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum to that of the baseline peaks $\{B_{N-1}\}$ on the baseline peripheral venous pressure frequency spectrum respectively,

wherein the intravascular volume status of the living subject at the second time period indicates hypovolemia or hypervolemia when amplitude changes greater than a threshold are detected from the baseline peaks $\{B_{N-1}\}$ on the baseline peripheral venous pressure frequency spectrum to the peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum.

2. The method of claim 1, wherein:
the intravascular volume status of the living subject at the second time period indicates hypovolemia when amplitude decreases are detected greater than a first threshold from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$; and
the intravascular volume status of the living subject at the second time period indicates hypervolemia when amplitude increases greater than a second threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$.
3. The method of claim 1, being performed to the living subject during resuscitation of the living subject, wherein:
the intravascular volume status of the living subject at the second time period indicates a return of euolemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at the first time period, and amplitude increases greater than a third threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$; and
the intravascular volume status of the living subject at the second time period indicates over-resuscitation when the living subject is determined to be in an euolemic state at the first time period, and amplitude increases greater than a fourth threshold are detected from the baseline peaks $\{B_{N-1}\}$ to the peaks $\{P_{N-1}\}$.
4. The method of claim 1, wherein the peripheral venous signals are acquired by:
capturing and recording the peripheral venous signals from a peripheral intravenous (IV) catheter in an upper extremity vein at a sampling rate of about 1 kHz.
5. The method of claim 4, wherein the peripheral IV catheter is a peripherally-inserted central catheter (PICC).
6. The method of claim 4, wherein a pressure transducer is directly connected to the peripheral IV catheter, and the peripheral venous signals are captured and recorded by the pressure transducer.

7. The method of claim 1, wherein the peripheral venous signals are processed by a spectral fast Fourier transform (FFT) analysis to obtain the baseline peripheral venous pressure frequency spectrum and the peripheral venous pressure frequency spectrum, respectively.

8. The method of claim 7, wherein:

the plurality of peaks $\{P_{N-1}\}$ comprises a first peak P_0 corresponding to a first frequency F_0 and a second peak P_1 corresponding to a second frequency F_1 ;

the first peak P_0 corresponding to the first frequency F_0 is associated with a respiratory rate of the living subject; and

the second peak P_1 corresponding to the second frequency F_1 is associated with a heart rate of the living subject.

9. A method for determining an intravascular volume status of a living subject by a peripheral intravenous (IV) waveform analysis (PIVA) system, the PIVA system including a peripheral IV device and a processing device and the method comprising:

acquiring, from a vein of the living subject and by the peripheral IV device, peripheral venous signals;

performing, by the processing device, a spectral analysis on the acquired peripheral venous signals to obtain a peripheral venous pressure frequency spectrum; and

performing, by the processing device, a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum by comparing the amplitudes of the peaks on the peripheral venous pressure frequency spectrum to amplitudes of a plurality of baseline peaks on a baseline peripheral venous pressure frequency spectrum respectively to determine the intravascular volume status of the living subject in real time, wherein the plurality of peaks corresponds to a plurality of frequencies.

10. The method of claim 9, wherein:
the intravascular volume status of the living subject indicates hypovolemia when amplitude decreases greater than a first threshold are detected from the peaks of the peripheral venous pressure frequency spectrum; and
the intravascular volume status of the living subject indicates hypervolemia when amplitude increases greater than a second threshold are detected from the peaks of the peripheral venous pressure frequency spectrum.

11. The method of claim 10, being performed to the living subject during ultrafiltration/dialysis or diuresis of the living subject, wherein the method further comprises:
generating an alert message when the intravascular volume status of the living subject indicates hypovolemia.

12. The method of claim 9, being performed to the living subject during resuscitation of the living subject, wherein:
the intravascular volume status of the living subject indicates a return of euolemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at an earlier time period, and amplitude increases greater than a third threshold are detected from the peaks of the peripheral venous pressure frequency spectrum; and
the intravascular volume status of the living subject indicates over-resuscitation when the living subject is determined to be in an euolemic state at the earlier time period, and amplitude increases greater than a fourth threshold are detected from the peaks of the peripheral venous pressure frequency spectrum.

13. The method of claim 12, further comprising:
detecting the return to euolemia in the living subject based on the intravascular volume status of the living subject.

14. The method of claim 9, wherein the peripheral venous signal is acquired by:
capturing and recording the peripheral venous signal from a peripheral intravenous (IV) catheter in the vein of the living subject catheter at a sampling rate,
wherein the peripheral IV catheter is a peripherally-inserted central catheter (PICC).

15. The method of claim 14, wherein a pressure transducer is directly connected to the peripheral IV catheter, and the peripheral venous signals are captured and recorded by the pressure transducer.

16. The method of claim 9, wherein the spectral analysis is a spectral fast Fourier transform (FFT) analysis, wherein the statistical analysis comprises:

obtaining the plurality of baseline peaks $\{B_{N-1}\}$ on the baseline peripheral venous pressure frequency spectrum, wherein N is a positive integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} ;

obtaining a plurality of peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum, wherein the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$; and

determining the intravascular volume status of the living subject in real time by comparing the amplitudes of the peaks $\{P_{N-1}\}$ to that of the baseline peaks $\{B_{N-1}\}$ respectively.

17. The method of claim 16, wherein the baseline peripheral venous pressure frequency spectrum is obtained by:

acquiring the peripheral venous signals from the vein of the living subject at an earlier time period; and

processing the peripheral venous signals acquired at the earlier time period by the spectral FFT analysis to obtain the baseline peripheral venous pressure frequency spectrum.

18. The method of claim 16, wherein:
the plurality of peaks $\{P_{N-1}\}$ comprises a first peak P_0 corresponding to a first frequency F_0 and a second peak P_1 corresponding to a second frequency F_1 ;
the first peak P_0 corresponding to the first frequency F_0 is associated with a respiratory rate of the living subject; and
the second peak P_1 corresponding to the second frequency F_1 is associated with a heart rate of the living subject.
19. A peripheral intravenous (IV) waveform analysis (PIVA) system, comprising:
a peripheral IV device configured to acquire, from a vein of a living subject, peripheral venous signals; and
a processing device communicatively connected to the peripheral IV device, configured to:
receive the peripheral venous signals from the peripheral IV device;
perform a spectral analysis on the peripheral venous signal to obtain a peripheral venous pressure frequency spectrum; and
perform a statistical analysis on amplitudes of peaks of the peripheral venous pressure frequency spectrum by comparing the amplitudes of the peaks on the peripheral venous pressure frequency spectrum to amplitudes of a plurality of baseline peaks on a baseline peripheral venous pressure frequency spectrum respectively to determine an intravascular volume status of the living subject in real time, wherein the plurality of peaks corresponds to a plurality of frequencies.
20. The PIVA system of claim 19, wherein:
the intravascular volume status of the living subject indicates hypovolemia when amplitude decreases greater than a first threshold are detected from the peaks of the peripheral venous pressure frequency spectrum; and
the intravascular volume status of the living subject indicates hypervolemia when amplitude increases greater than a second threshold are detected from the peaks of the peripheral venous pressure frequency spectrum.

21. The PIVA system of claim 19, being applied to the living subject during ultrafiltration/dialysis or diuresis of the living subject, wherein the processing device is further configured to:

generate an alert message when the intravascular volume status of the living subject indicates hypovolemia.

22. The PIVA system of claim 19, being applied to the living subject during resuscitation of the living subject, wherein:

the intravascular volume status of the living subject indicates a return of euvoolemia from a hypovolemic state when the living subject is determined to be in the hypovolemic state at an earlier time period, and amplitude increases greater than a third threshold are detected from the peaks of the peripheral venous pressure frequency spectrum; and

the intravascular volume status of the living subject indicates over- resuscitation when the living subject is determined to be in an euvolemic state at the earlier time period, and amplitude increases greater than a fourth threshold are detected from the peaks of the peripheral venous pressure frequency spectrum.

23. The PIVA system of claim 22, wherein the processing device is further configured to: detect efficacy of treatment and the return to euvoolemia in the living subject based on the intravascular volume status of the living subject.

24. The PIVA system of claim 19, wherein the processing device is communicatively connected to the peripheral IV device through a wireless connection.

25. The PIVA system of claim 19, wherein the peripheral IV device comprises:
a peripheral IV catheter to be inserted into the vein of the living subject; and
a monitoring device connected to the peripheral IV catheter, configured to capture and record the peripheral venous signals from the peripheral IV catheter at a sampling rate.

26. The PIVA system of claim 25, wherein the monitoring device comprises a pressure transducer directly connected to the peripheral IV catheter, wherein the peripheral venous signals are captured and recorded by the pressure transducer.

27. The PIVA system of claim 19, wherein the processing device is a computing device.

28. The PIVA system of claim 19, wherein the spectral analysis is a spectral fast Fourier transform (FFT) analysis, wherein the statistical analysis comprises:

obtaining the plurality of baseline peaks $\{B_{N-1}\}$ on the baseline peripheral venous pressure frequency spectrum, wherein N is a positive integer, and the plurality of baseline peaks $\{B_{N-1}\}$ respectively corresponds to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that B_{N-1} is a function of F_{N-1} satisfying $B_{N-1} = B_{N-1}(F_{N-1})$, wherein F_N is greater than F_{N-1} ;

obtaining a plurality of peaks $\{P_{N-1}\}$ on the peripheral venous pressure frequency spectrum, wherein the plurality of peaks $\{P_{N-1}\}$ correspond to the plurality of frequencies $\{F_0, F_1, \dots, F_N\}$, such that P_{N-1} is a function of F_{N-1} satisfying $P_{N-1} = P_{N-1}(F_{N-1})$; and

determining the intravascular volume status of the living subject in real time by comparing the amplitudes of the peaks $\{P_{N-1}\}$ to that of the baseline peaks $\{B_{N-1}\}$ respectively.

29. The PIVA system of claim 28, wherein the baseline peripheral venous pressure frequency spectrum is obtained by:

acquiring, by the peripheral IV monitoring device, the peripheral venous signals from the vein of the living subject at an earlier time period; and

processing the peripheral venous signals acquired at the earlier time period by the spectral FFT analysis to obtain the baseline peripheral venous pressure frequency spectrum.

30. The PIVA system of claim 28, wherein:

the plurality of peaks $\{P_{N-1}\}$ comprises a first peak P_0 corresponding to a first frequency F_0 and a second peak P_1 corresponding to a second frequency F_1 ;

the first peak P_0 corresponding to the first frequency F_0 is associated with a respiratory rate of the living subject; and the second peak P_1 corresponding to the second frequency F_1 is associated with a heart rate of the living subject.

31. The PIVA system of claim 19, further comprises:

a pump to be connected to the living subject to perform liquid exchange to the living subject; and

a pump controlling mechanism communicatively connected to the processing device, configured to control the pump by intermittently pausing the pump or subtract the pump signal when the peripheral IV device acquires the peripheral venous signals, and restarting the pump when the peripheral IV device does not acquire the peripheral venous signals;

wherein the processing device is further configured to send a signal to the pump controlling mechanism to notify the pump controlling mechanism to control the pump.

32. The PIVA system of claim 31, wherein the pump is a dialysis pump, a cardiopulmonary bypass pump, an extracorporeal membrane oxygenation (ECMO), or an infusion pump.

33. Use of the PIVA system of claim 19 for determining hypovolemia, hypervolemia and vascular tone of a living subject based on an intravascular volume status of the living subject.

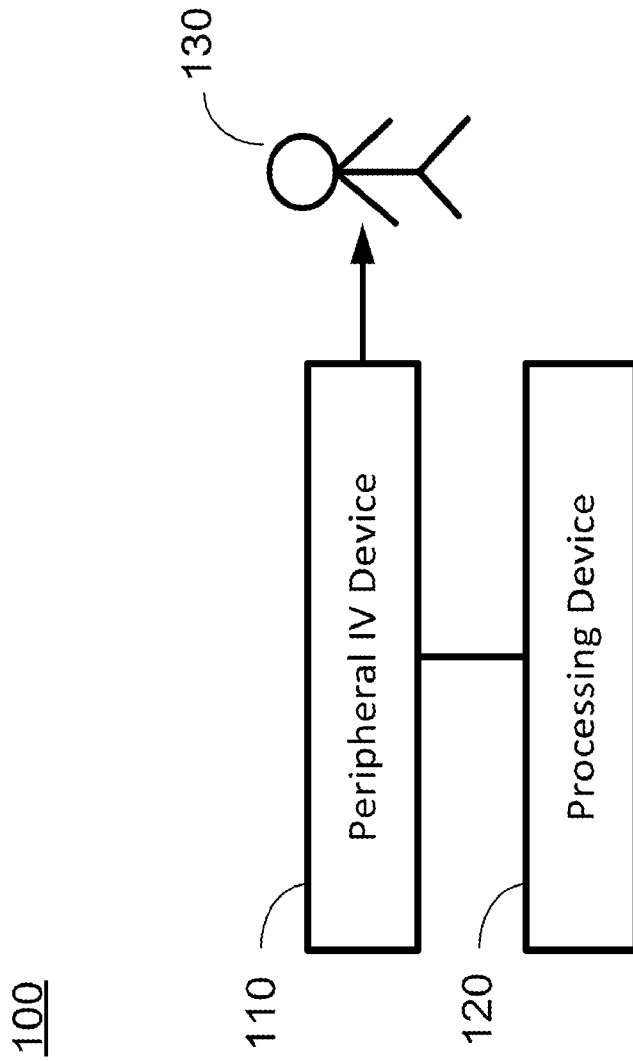


FIG. 1

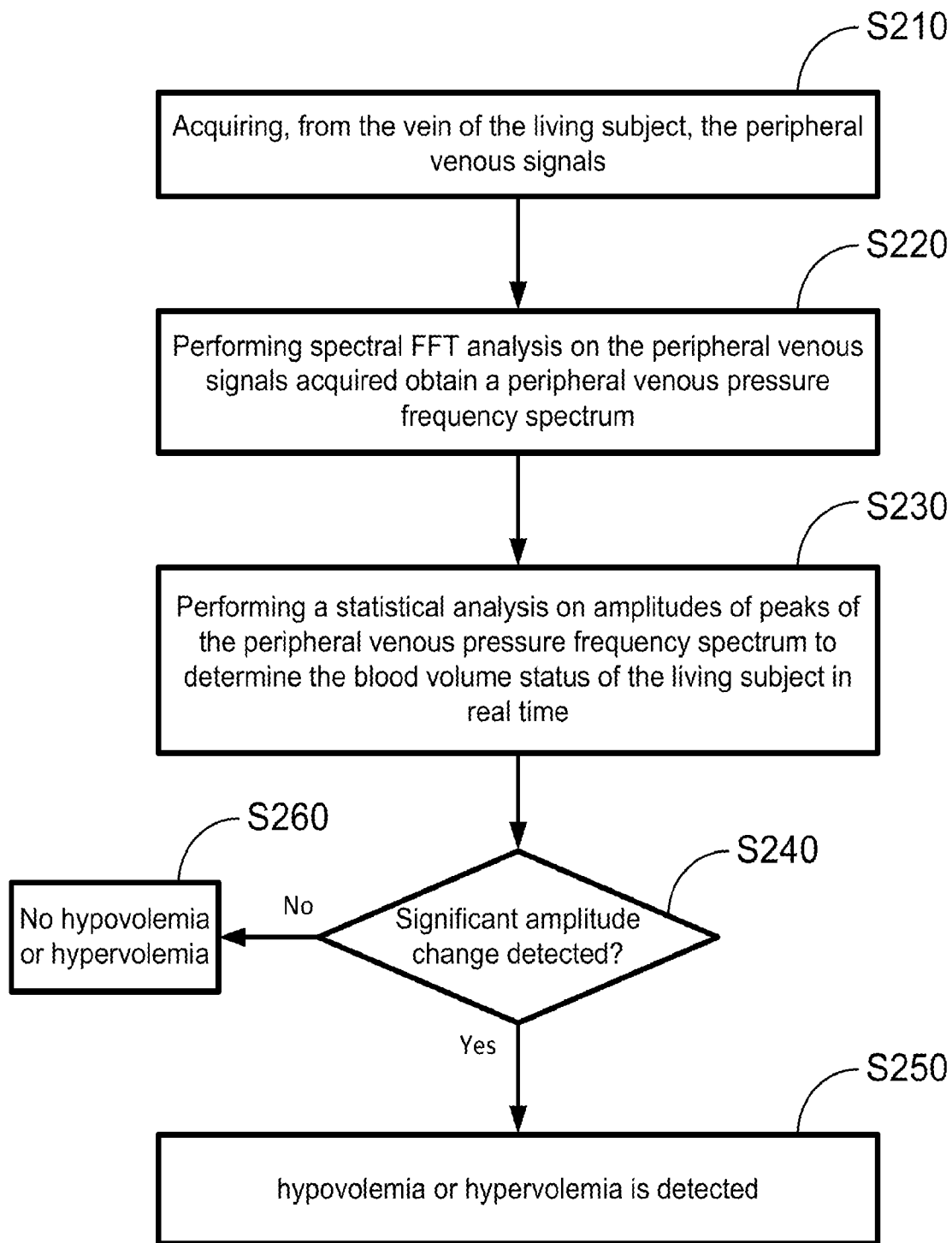


FIG. 2

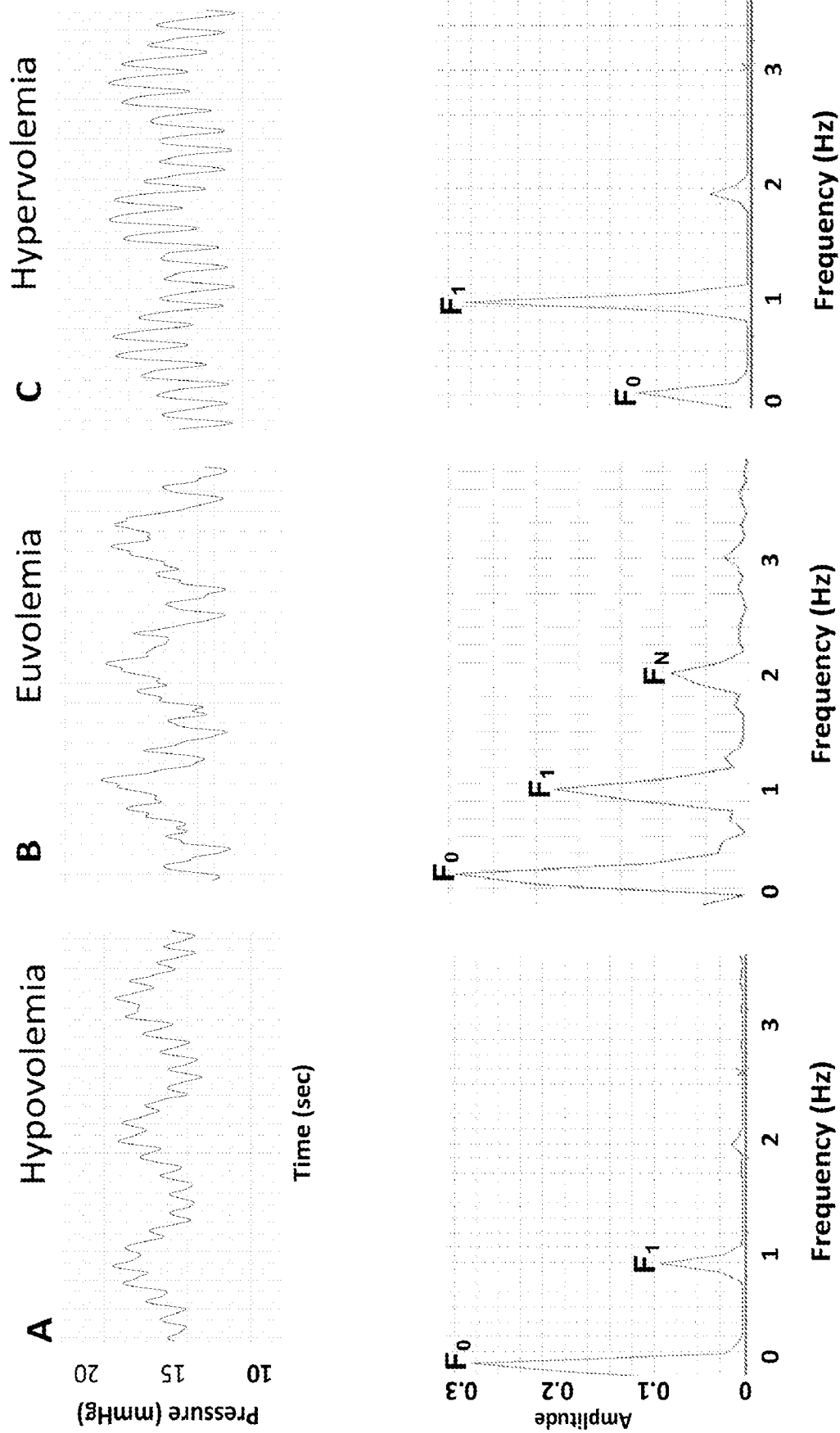


FIG. 3

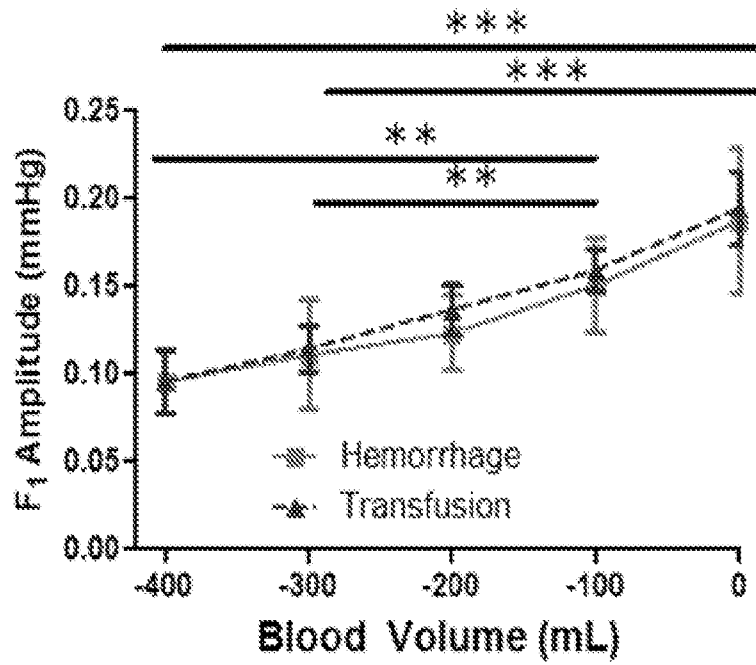


FIG. 4A

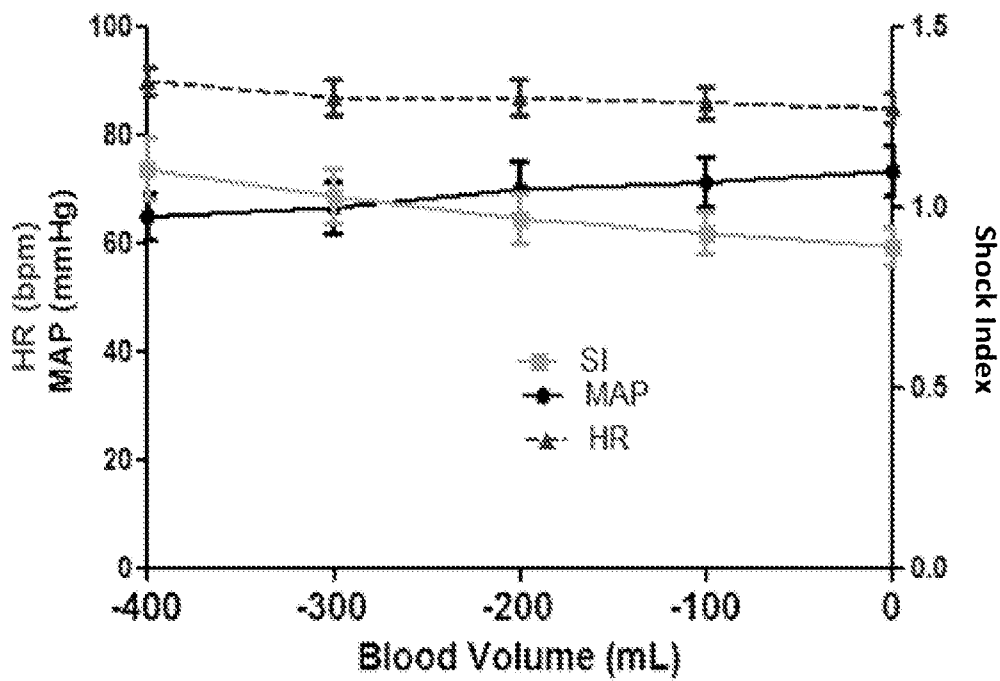


FIG. 4B

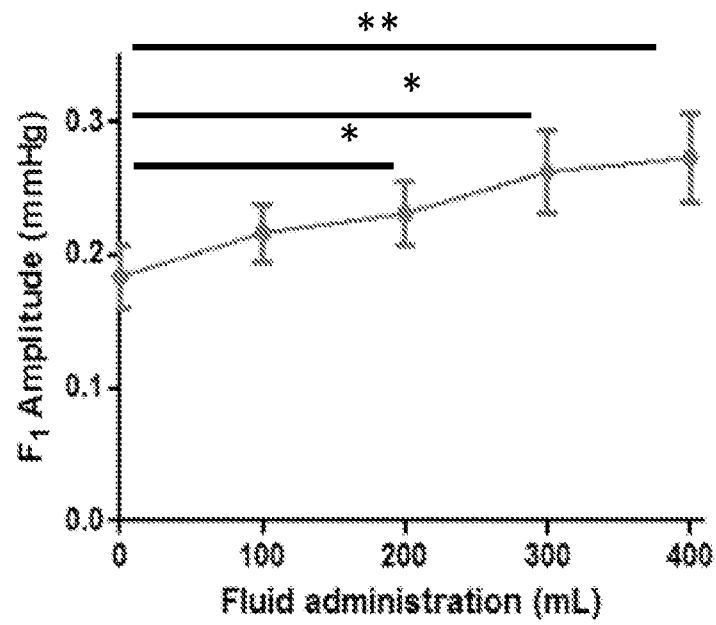


FIG. 5A

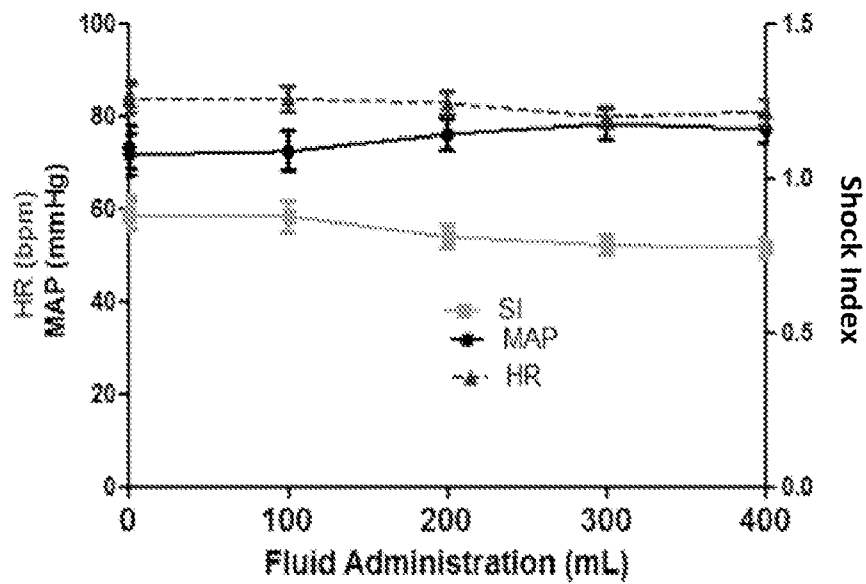


FIG. 5B

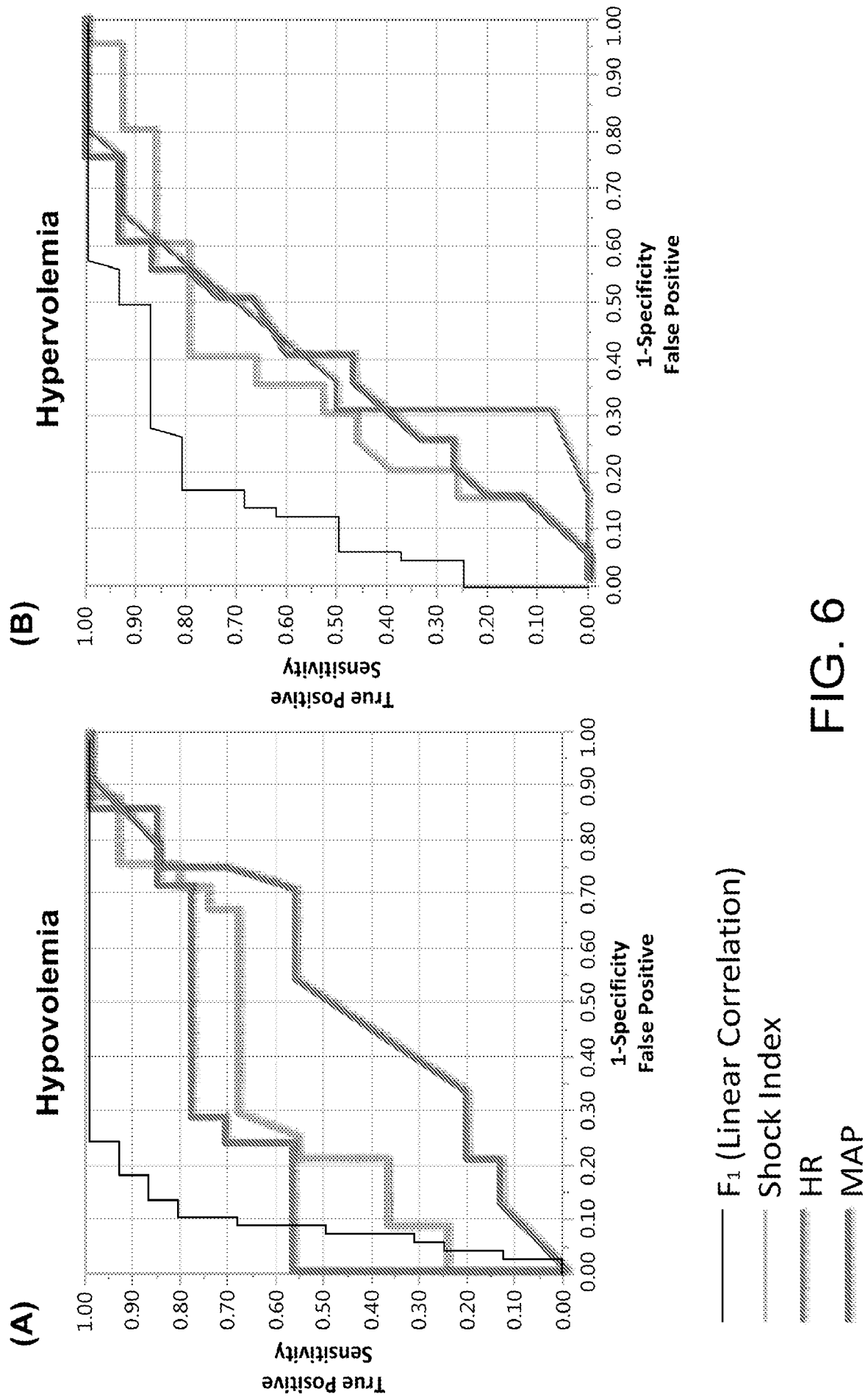


FIG. 6

Table 1. Patient Demographics.

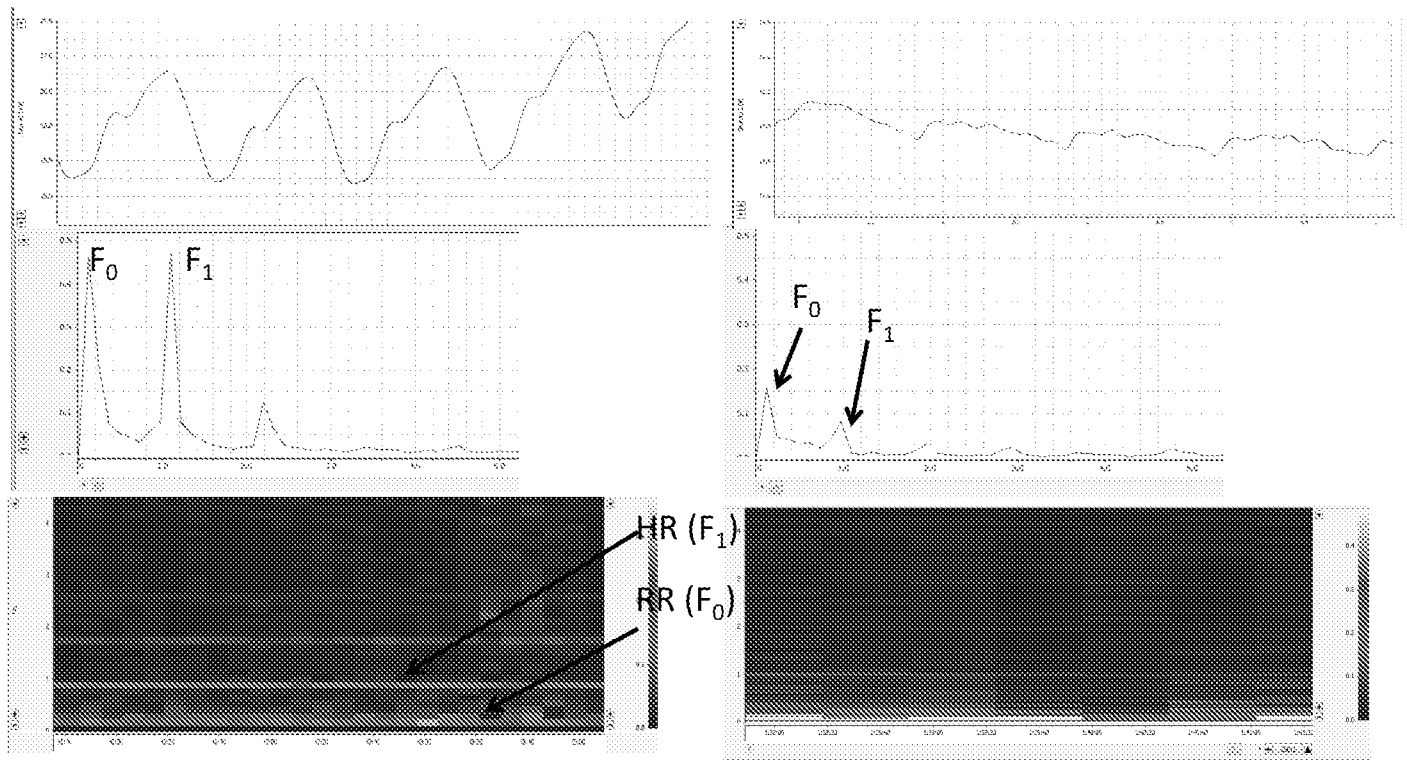
Patient number	Age	Gender	Preoperative Diagnosis	Surgery	Wt. (kg)	BSA	Diabetes	Creatinine	Preoperative Hgb	Ejection Fraction	RV/dysfunction	RV size (gauge)	Baseline PAP (mean)
1	70	F	Severe MR	M/R	85	1.9	Yes	1.52	12.7	45-50	Nbre	18	34/12 (20)
2	63	M	Severe MR, CAD	M/R	93	2.03	Yes	2.01	12.3	15-20	Nbre	18	31/14 (20)
3	54	M	Severe MR	M/R/repair	96	2.21	No	1.2	14.4	>55	Nbre	18	22/15 (17)
4	85	M	Severe AS, CAD	A/R, CABG	89	2.1	Yes	0.88	12.5	30-40	Nbre	18	25/17 (20)
5	33	M	Severe AI	A/R	73	1.9	No	0.82	15.4	>55	Nbre	16	19/11 (14)
6	68	M	Severe AS	A/R	98	2.25	No	1.79	13.5	35-44	Nbre	18	18/10 (14)
7	74	F	CAD	CABG	92	2.07	No	1.07	9.8	>55	Nbre	18	31/15 (21)
8	71	F	Severe MR, moderate TR	M/R, TV/repair	71	1.78	No	0.76	14.2	45-55	Nbre	18	32/20 (19)
9	76	M	Severe MR	M/R/repair	90	2.03	No	0.83	14.2	>55	Nbre	18	44/20 (23)
10	58	M	Chronic PE	Pulmonary endarterectomy	80	1.94	No	1.15	13.9	>55	Mild	16	115/42 (68)
11	57	M	Aortic dissection	Aortic dissection repair	88	2.11	No	0.79	12.8	>55	Nbre	18	46/22 (15)
12	77	M	Severe MR	M/R	103	2.27	Yes	1.28	15.6	>55	Nbre	18	68/32 (44)

Abbreviations: AI – Aortic insufficiency, AS – Aortic stenosis, AVR – aortic valve replacement, BSA – body surface area, CABG – Coronary artery bypass grafting, CAD – Coronary artery disease, Hgb – hemoglobin, MR – mitral regurgitation, MV – mitral valve, MVR – mitral valve replacement, PAP – pulmonary artery pressure, PE – Pulmonary embolus, PIV – peripheral intravenous catheter, RV – Right ventricle, TR – tricuspid regurgitation, TV – tricuspid valve, Wt. – weight

FIG. 7

Baseline

Hypovolemia



time →

FIG. 8

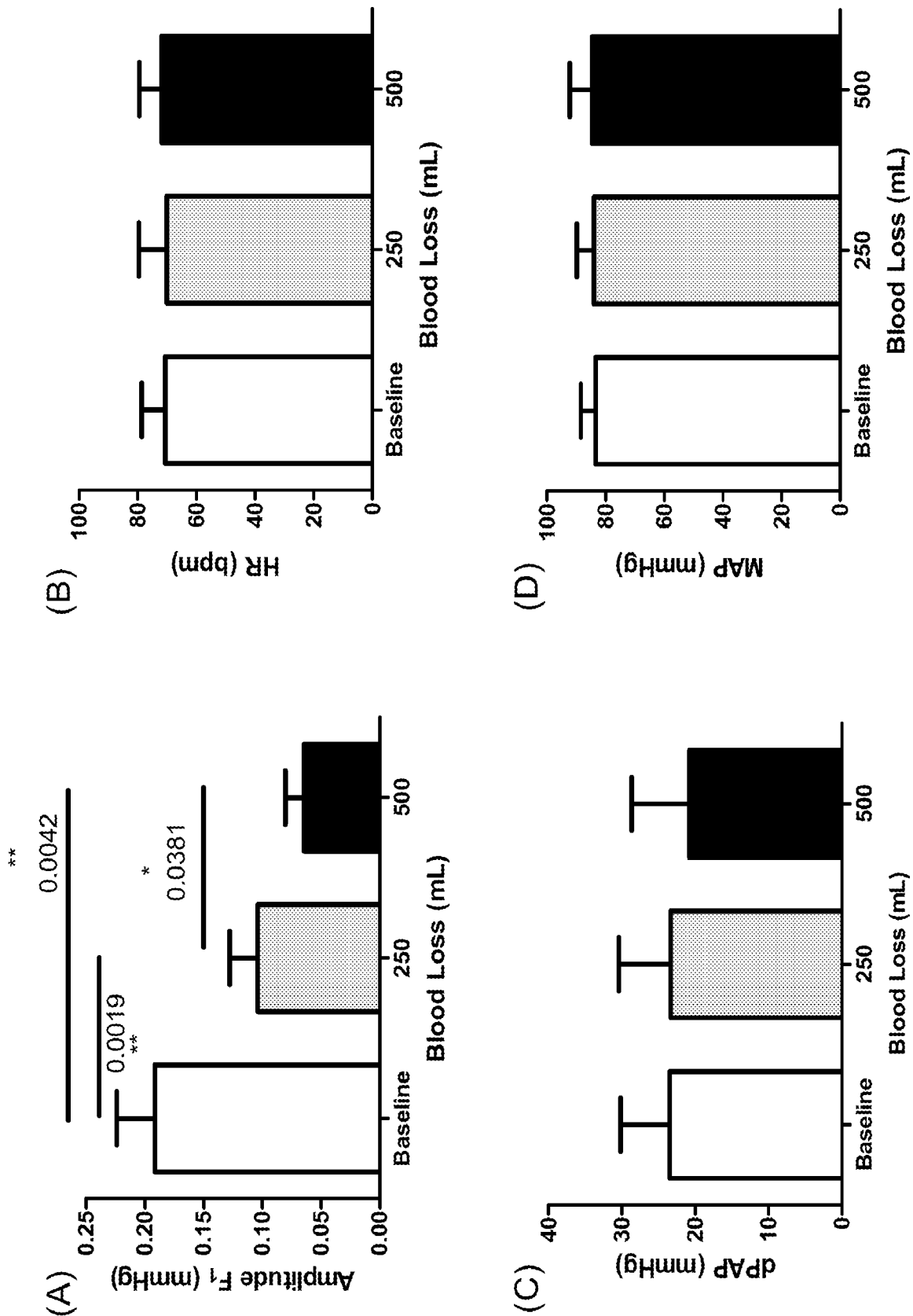


FIG. 9

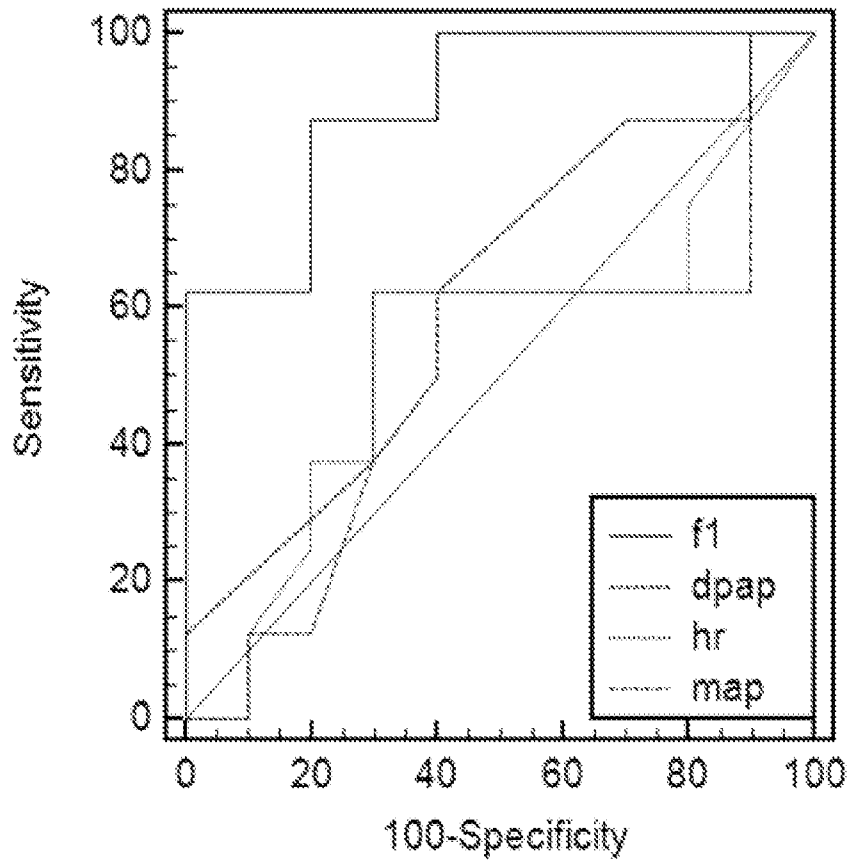


FIG. 10A

Variable	AUC	SE	95% CI
f1	0.900	0.0722	0.667 to 0.990
dpap	0.619	0.137	0.365 to 0.833
hr	0.544	0.152	0.297 to 0.775
map	0.512	0.155	0.271 to 0.750

FIG. 10B

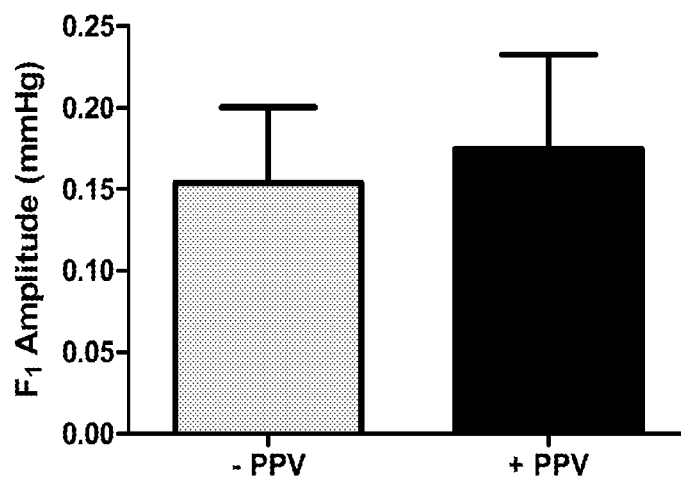


FIG. 11A

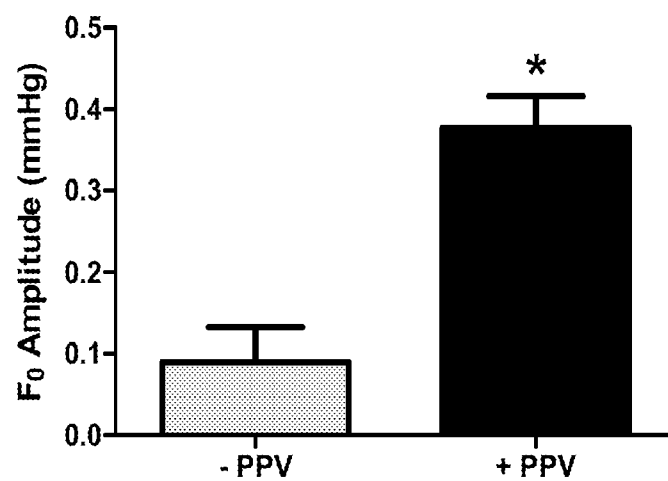


FIG. 11B

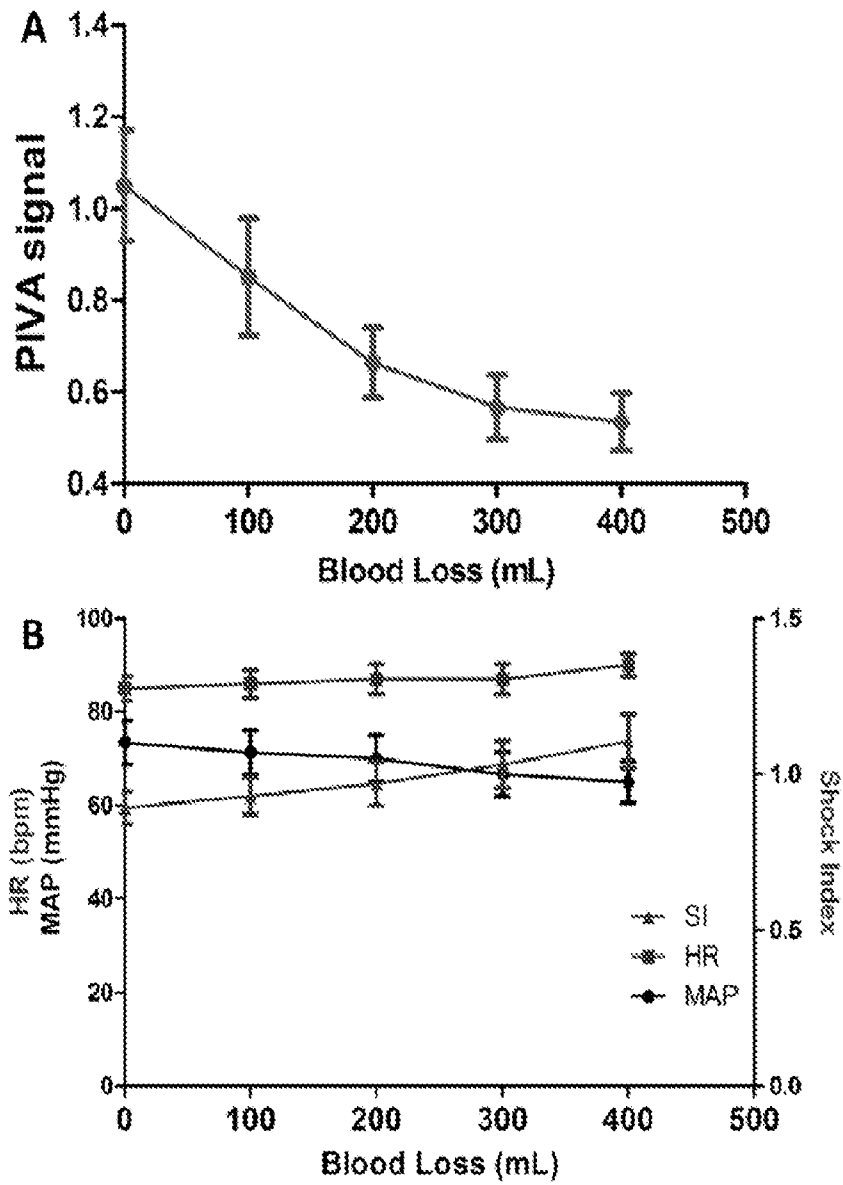


FIG. 12

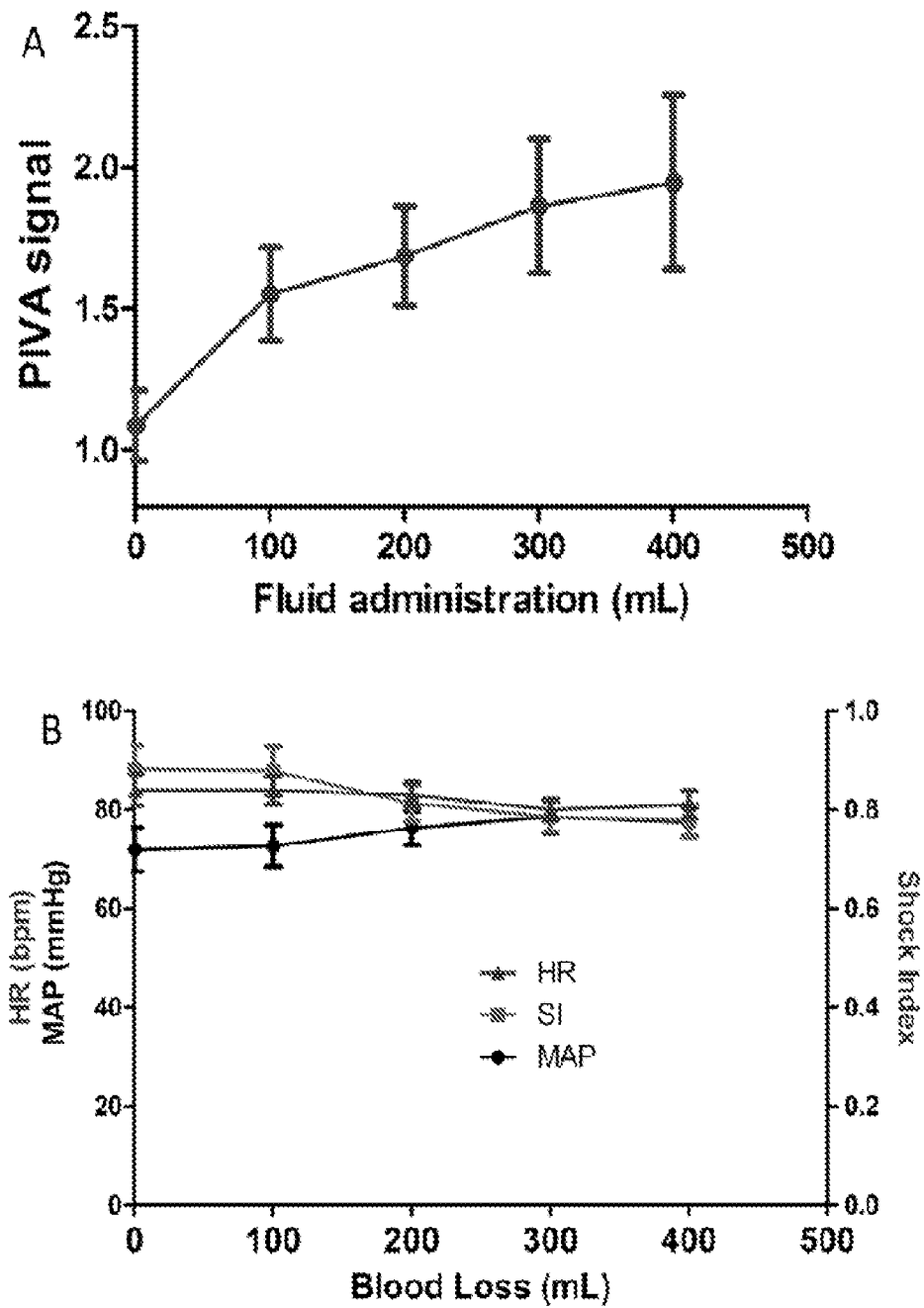


FIG. 13

