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(54) **SYSTEM AND METHOD FOR DETECTING FLUID RESPONSIVENESS OF A PATIENT**

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“Derivation of Respiratory Signals from Multi-lead EGGs, Moody,” et al. (1985).
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Related U.S. Application Data

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(57) **ABSTRACT**

A system is configured to determine a fluid responsiveness index of a patient from a physiological signal. The system may include a sensor configured to be secured to an anatomical portion of the patient, and a monitor operatively connected to the sensor. The sensor is configured to sense a physiological characteristic of the patient. The monitor is configured to receive a physiological signal from the sensor. The monitor may include an index-determining module configured to determine the fluid responsiveness index through formation of a ratio of one or both of amplitude or frequency modulation of the physiological signal to baseline modulation of the physiological signal.

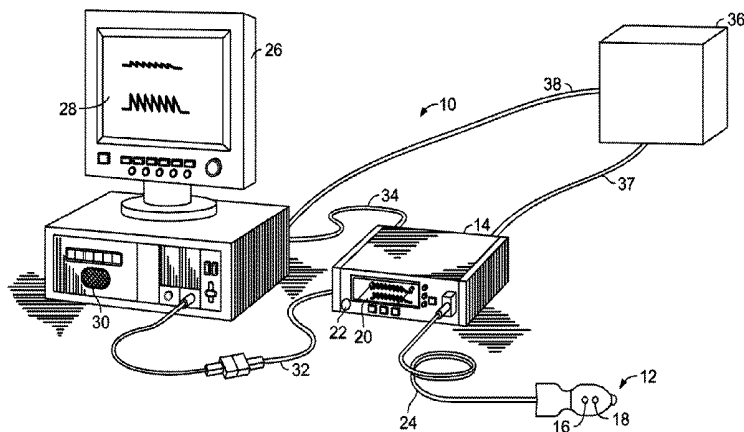
(52) **U.S. Cl.**

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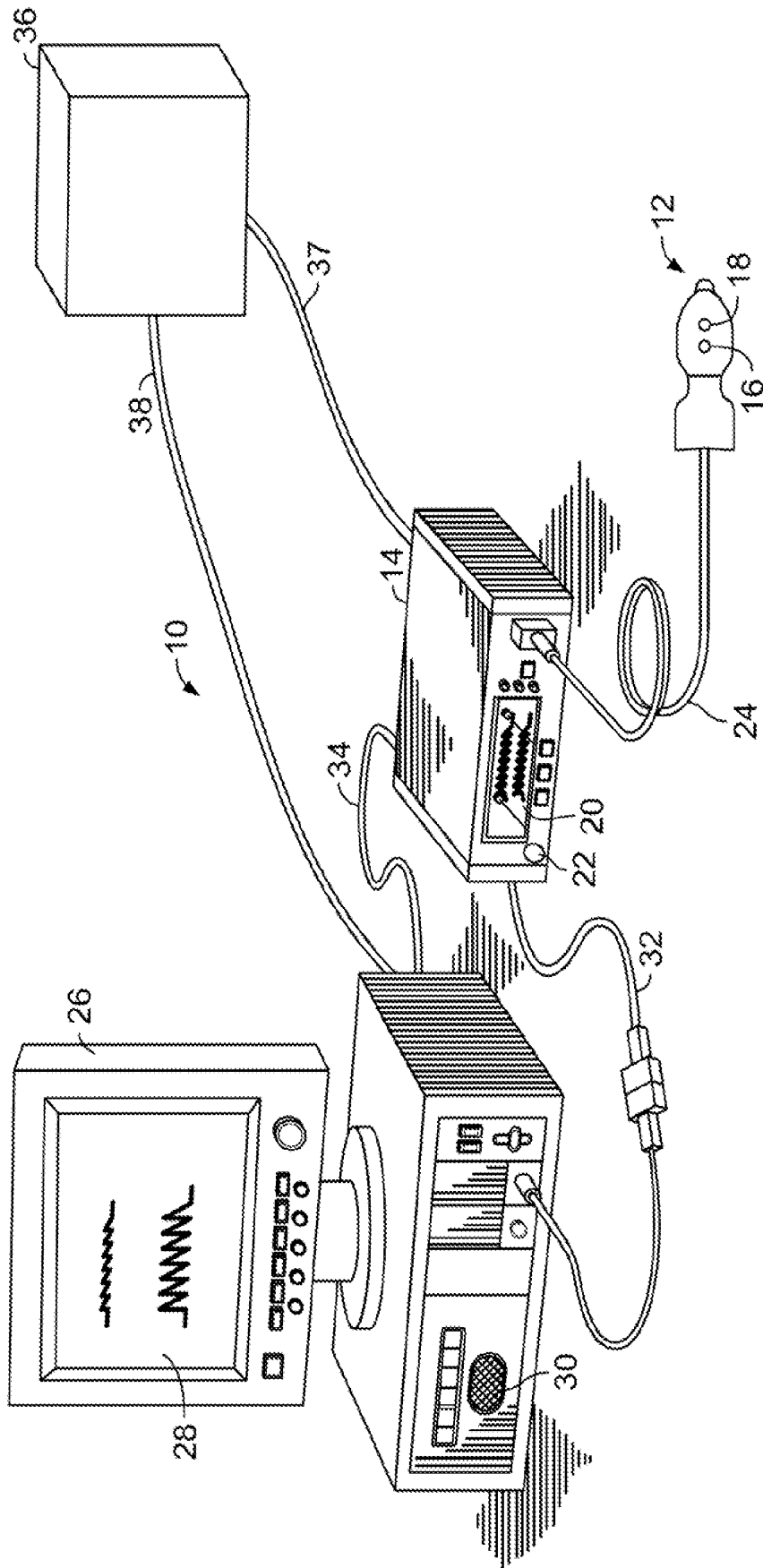


FIG. 1

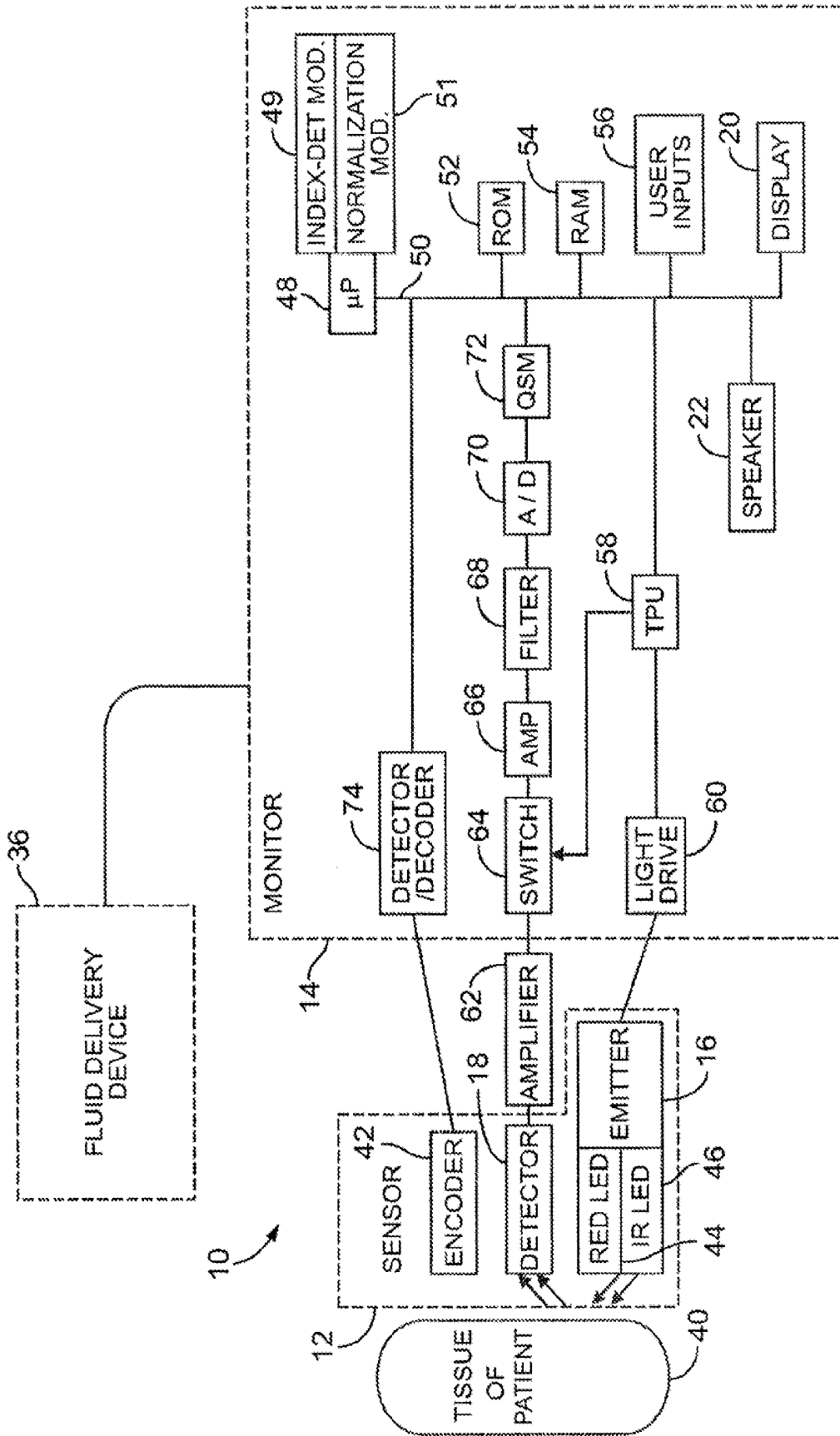


FIG. 2

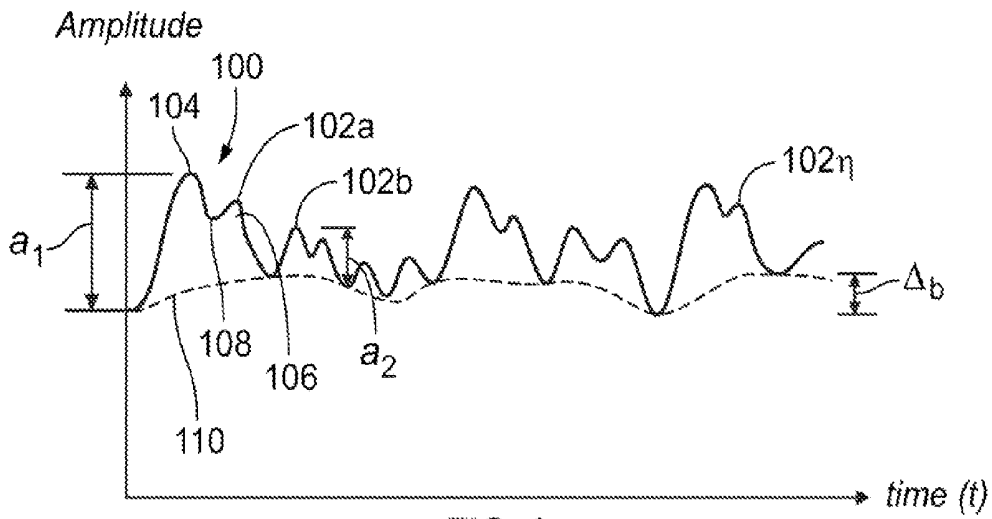


FIG. 3

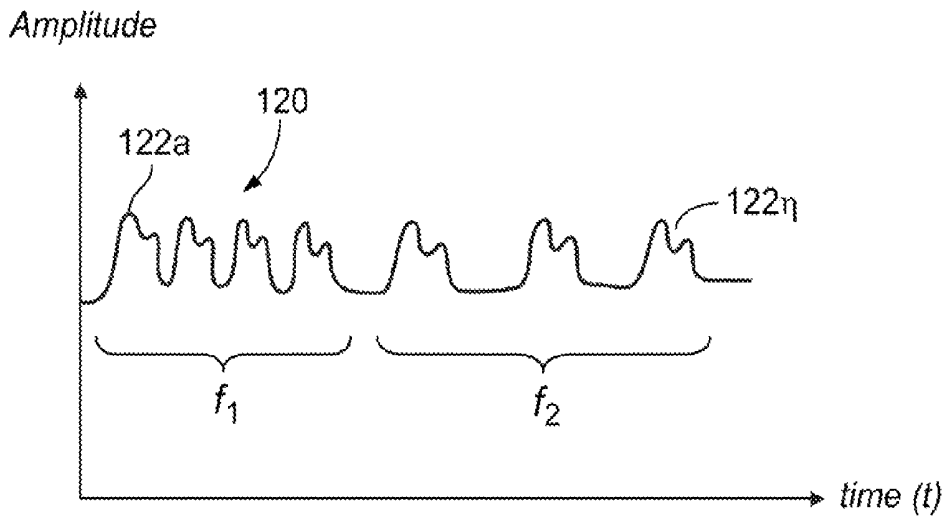


FIG. 4

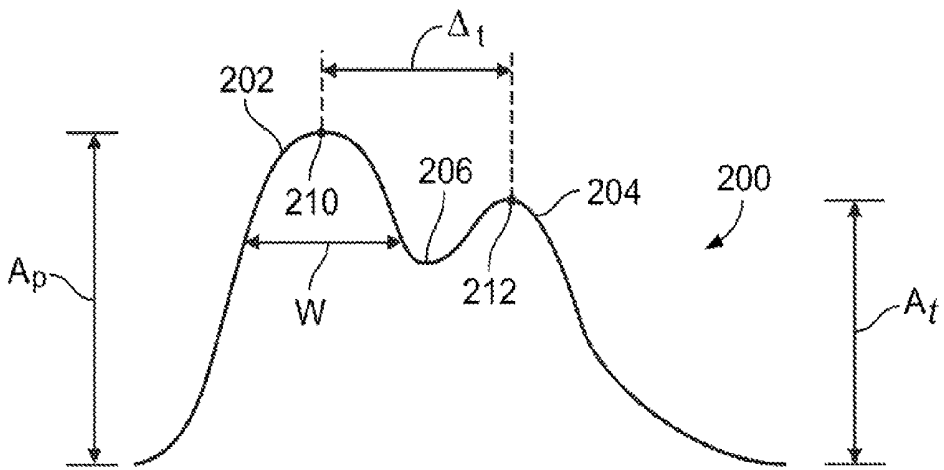


FIG. 6

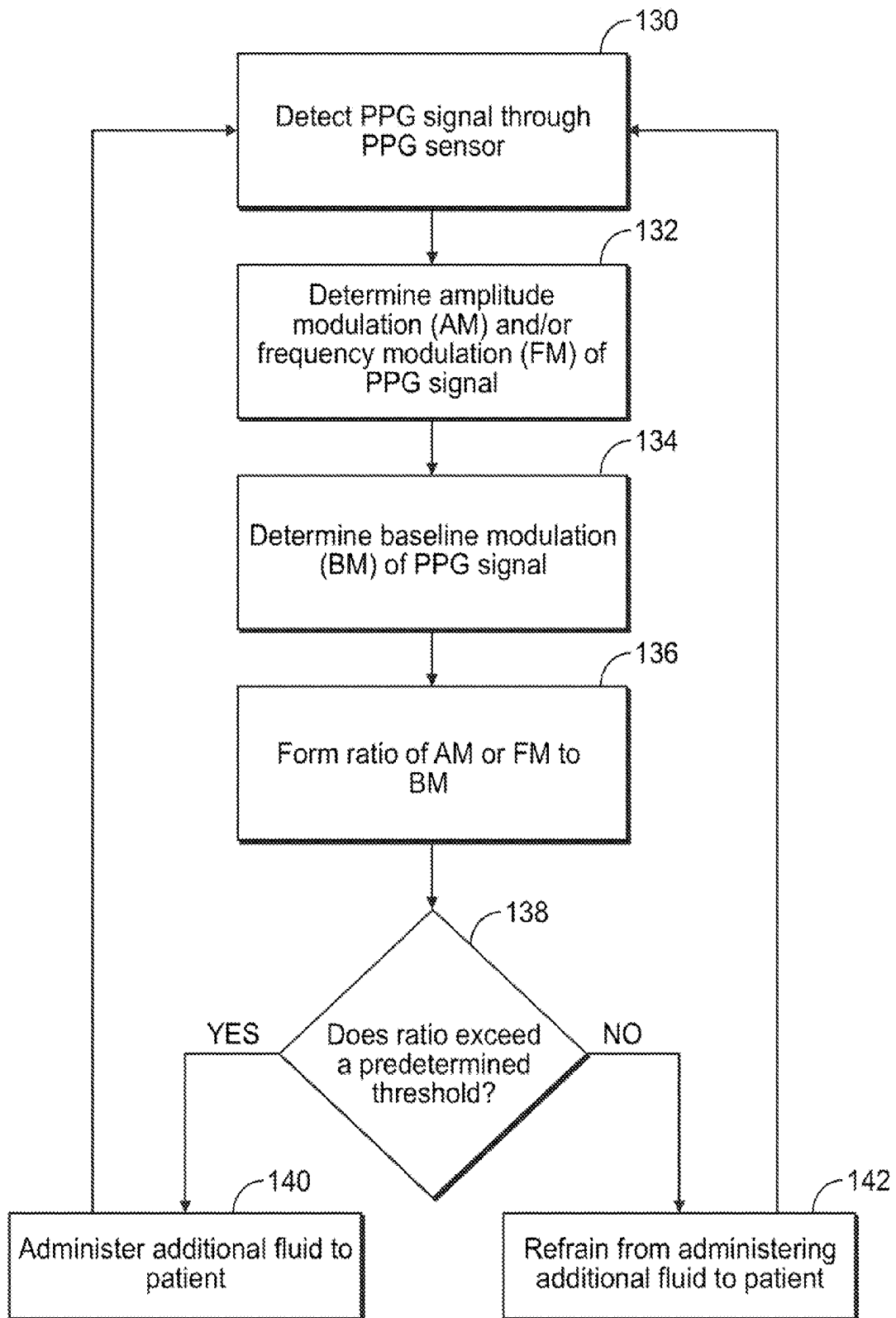


FIG. 5

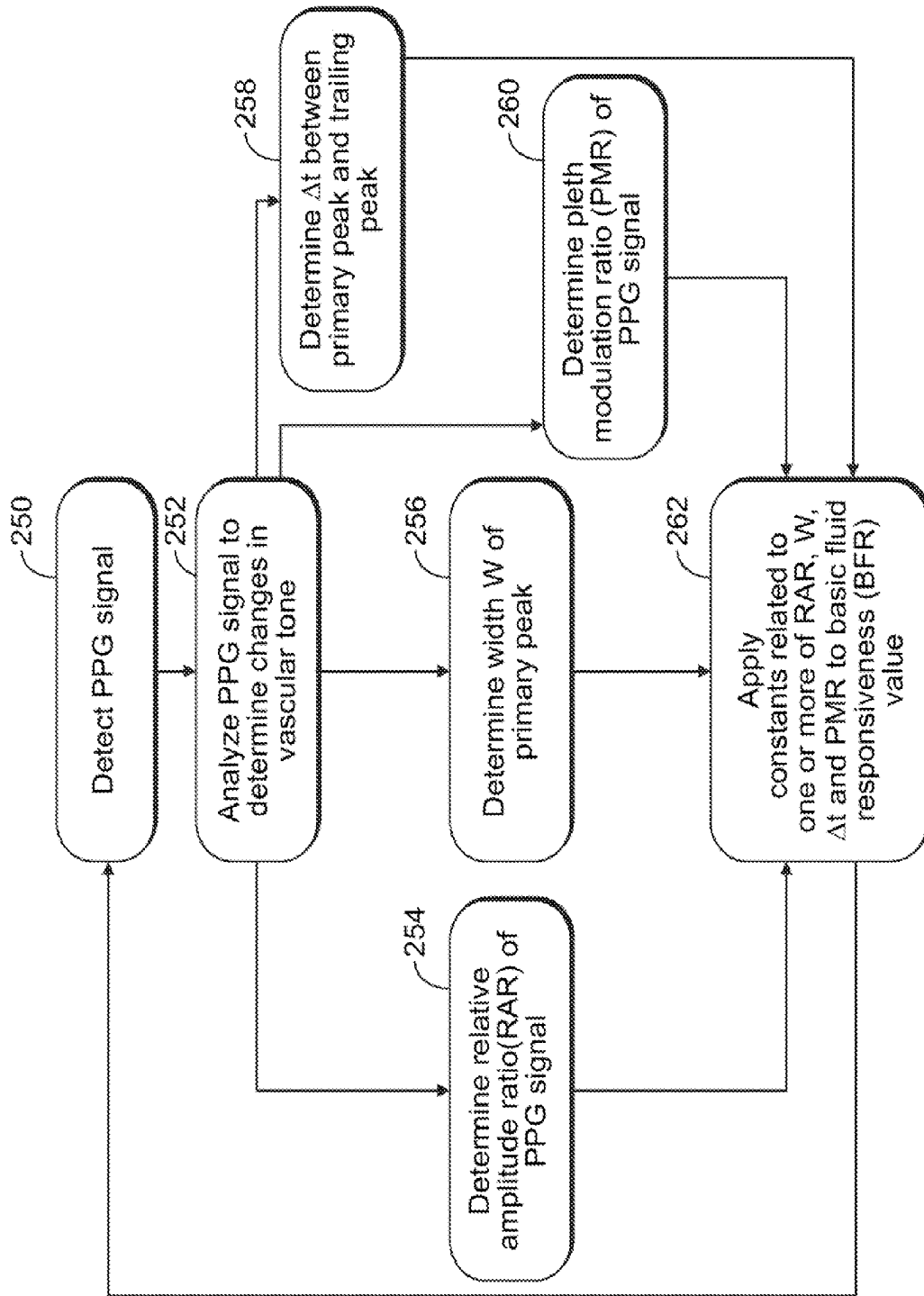


FIG. 7

SYSTEM AND METHOD FOR DETECTING FLUID RESPONSIVENESS OF A PATIENT

CROSS REFERENCE TO RELATED APPLICATION

This application is a continuation of U.S. patent application Ser. No. 13/591,455, filed Aug. 22, 2012, now U.S. Pat. No. 9,060,745, which is hereby incorporated by reference herein in its entirety.

FIELD

Embodiments of the present disclosure generally relate to physiological signal processing and more particularly, to processing physiological signals to determine the fluid responsiveness of a patient.

BACKGROUND

Photoplethysmography (PPG) is a non-invasive, optical measurement that may be used to detect changes in blood volume within tissue, such as skin, of an individual. PPG may be used with pulse oximeters, vascular diagnostics, and digital blood pressure detection systems. Typically, a PPG system includes a light source that is used to illuminate tissue of a patient. A photodetector is then used to measure small variations in light intensity associated with blood volume changes proximal to the illuminated tissue.

In general, a PPG signal is a physiological signal that includes an AC physiological component related to cardiac synchronous changes in the blood volume with each heartbeat. The AC component is typically superimposed on a DC baseline that may be related to respiration, sympathetic nervous system activity, and thermoregulation.

SUMMARY

Certain embodiments provide a system for determining a fluid responsiveness index of a patient from a physiological signal, such as a PPG signal. The system may include a sensor configured to be secured to an anatomical portion of the patient, and a monitor operatively connected to the sensor. The sensor is configured to sense a physiological characteristic of the patient. The monitor is configured to receive a physiological signal from the sensor. The monitor includes an index-determining module configured to determine the fluid responsiveness index through formation of a ratio of one or both of amplitude modulation or frequency modulation of the physiological signal with respect to baseline modulation of the physiological signal.

The monitor may include a normalization module configured to normalize the fluid responsiveness index. The normalization module may be configured to normalize the fluid responsiveness index to correct for changes in one or both of vascular tone or motion artifacts. The normalization module may be configured to normalize the fluid responsiveness index based on one or more of a relative amplitude ratio of the physiological signal, a width of at least one primary peak of the physiological signal, a time difference between the at least one primary peak and at least one trailing peak, or a pleth modulation ratio of the physiological signal.

The index-determining module may determine the fluid responsiveness index by applying a calibration coefficient or constant to the ratio.

The physiological signal may include a photoplethysmogram (PPG) signal. The sensor may include a PPG sensor.

The monitor may include a PPG monitor. The sensor may include a pulse oximetry sensor. The monitor may include a pulse oximetry monitor.

The system may also include a fluid delivery device operatively connected to the monitor. The monitor may automatically determine whether or not to administer fluid to a patient from the fluid delivery device based on the fluid responsiveness index.

Certain embodiments provide a method for determining a fluid responsiveness index of a patient from a physiological signal. The method may include securing a sensor to an anatomical portion of the patient, sensing a physiological characteristic of the patient with the sensor, receiving a physiological signal from the sensor at a monitor comprising an index-determining module, and determining the fluid responsiveness index with the index-determining module. The determining the fluid responsiveness index operation may include forming a ratio of one or both of amplitude modulation or frequency modulation of the physiological signal with respect to baseline modulation of the physiological signal.

Certain embodiments provide a tangible and non-transitory computer readable medium that includes one or more sets of instructions configured to direct a computer to receive a physiological signal from a sensor secured to an anatomical portion of a patient, and determine a fluid responsiveness index of a patient by forming a ratio of one or both of amplitude modulation or frequency modulation of the physiological signal with respect to baseline modulation of the physiological signal.

Embodiments of the present disclosure allow for quick and simple determination of fluid responsiveness through analysis of a physiological signal, such as a PPG signal. In contrast to previous systems and methods, embodiments may not require an arterial line ("A-line") having a pressure transducer to detect fluid responsiveness. Moreover, embodiments may be used to detect fluid responsiveness with patients that are spontaneously breathing and/or those that are on lung-protective low-tidal volume ventilation, in contrast to previous systems and methods that typically require high-tidal ventilation in order to determine fluid responsiveness. Further, embodiments provide systems and methods for normalizing a basic fluid responsiveness index or value, such as by accounting for changes in vascular tone and/or motion artifacts.

Certain embodiments may include some, all, or none of the above advantages. One or more other technical advantages may be readily apparent to those skilled in the art from the figures, descriptions, and claims included herein. Moreover, while specific advantages have been enumerated above, various embodiments may include all, some, or none of the enumerated advantages.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 illustrates an isometric view of a photoplethysmogram (PPG) system, according to an embodiment.

FIG. 2 illustrates a simplified block diagram of a PPG system, according to an embodiment.

FIG. 3 illustrates a PPG signal over time, according to an embodiment.

FIG. 4 illustrates a PPG signal over time, according to an embodiment.

FIG. 5 illustrates a flow chart of a method of determining fluid responsiveness, according to an embodiment.

FIG. 6 illustrates a pulse of a PPG signal, according to an embodiment.

FIG. 7 illustrates a flow chart of a method normalizing a fluid responsiveness index, according to an embodiment.

DETAILED DESCRIPTION

FIG. 1 illustrates an isometric view of a PPG system 10, according to an embodiment. While the system 10 is shown and described as a PPG system 10, the system may be various other types of physiological detection systems, such as an electrocardiogram system, a phonocardiogram system, and the like. The PPG system 10 may be a pulse oximetry system, for example. The system 10 may include a PPG sensor 12 and a PPG monitor 14. The PPG sensor 12 may include an emitter 16 configured to emit light into tissue of a patient. For example, the emitter 16 may be configured to emit light at two or more wavelengths into the tissue of the patient. The PPG sensor 12 may also include a detector 18 that is configured to detect the emitted light from the emitter 16 that emanates from the tissue after passing through the tissue.

The system 10 may include a plurality of sensors forming a sensor array in place of the PPG sensor 12. Each of the sensors of the sensor array may be a complementary metal oxide semiconductor (CMOS) sensor, for example. Alternatively, each sensor of the array may be a charged coupled device (CCD) sensor. In another embodiment, the sensor array may include a combination of CMOS and CCD sensors. The CCD sensor may include a photoactive region and a transmission region configured to receive and transmit, while the CMOS sensor may include an integrated circuit having an array of pixel sensors. Each pixel may include a photodetector and an active amplifier.

The emitter 16 and the detector 18 may be configured to be located at opposite sides of a digit, such as a finger or toe, in which case the light that is emanating from the tissue passes completely through the digit. The emitter 16 and the detector 18 may be arranged so that light from the emitter 16 penetrates the tissue and is reflected by the tissue into the detector 18, such as a sensor designed to obtain pulse oximetry data.

The sensor 12 or sensor array may be operatively connected to and draw power from the monitor 14. Optionally, the sensor 12 may be wirelessly connected to the monitor 14 and include a battery or similar power supply (not shown). The monitor 14 may be configured to calculate physiological parameters based at least in part on data received from the sensor 12 relating to light emission and detection. Alternatively, the calculations may be performed by and within the sensor 12 and the result of the oximetry reading may be passed to the monitor 14. Additionally, the monitor 14 may include a display 20 configured to display the physiological parameters or other information about the system 10. The monitor 14 may also include a speaker 22 configured to provide an audible sound that may be used in various other embodiments, such as for example, sounding an audible alarm in the event that physiological parameters are outside a predefined normal range.

The sensor 12, or the sensor array, may be communicatively coupled to the monitor 14 via a cable 24. Alternatively, a wireless transmission device (not shown) or the like may be used instead of, or in addition to, the cable 24.

The system 10 may also include a multi-parameter workstation 26 operatively connected to the monitor 14. The workstation 26 may be or include a computing sub-system 30, such as standard computer hardware. The computing sub-system 30 may include one or more modules and control units, such as processing devices that may include one or more microprocessors, microcontrollers, integrated circuits, memory, such as read-only and/or random access memory, and the like.

The workstation 26 may include a display 28, such as a cathode ray tube display, a flat panel display, such as a liquid crystal display (LCD), light-emitting diode (LED) display, a plasma display, or any other type of monitor. The computing sub-system 30 of the workstation 26 may be configured to calculate physiological parameters and to show information from the monitor 14 and from other medical monitoring devices or systems (not shown) on the display 28. For example, the workstation 26 may be configured to display an estimate of a patient's blood oxygen saturation generated by the monitor 14 (referred to as an SpO₂ measurement), pulse rate information from the monitor 14 and blood pressure from a blood pressure monitor (not shown) on the display 28.

The monitor 14 may be communicatively coupled to the workstation 26 via a cable 32 and/or 34 that is coupled to a sensor input port or a digital communications port, respectively and/or may communicate wirelessly with the workstation 26. Additionally, the monitor 14 and/or workstation 26 may be coupled to a network to enable the sharing of information with servers or other workstations. The monitor 14 may be powered by a battery or by a conventional power source such as a wall outlet.

The system 10 may also include a fluid delivery device 36 that is configured to deliver fluid to a patient. The fluid delivery device 36 may be an intravenous line, an infusion pump, any other suitable fluid delivery device, or any combination thereof that is configured to deliver fluid to a patient. The fluid delivered to a patient may be saline, plasma, blood, water, any other fluid suitable for delivery to a patient, or any combination thereof. The fluid delivery device 36 may be configured to adjust the quantity or concentration of fluid delivered to a patient.

The fluid delivery device 36 may be communicatively coupled to the monitor 14 via a cable 37 that is coupled to a digital communications port or may communicate wirelessly with the workstation 26. Alternatively, or additionally, the fluid delivery device 36 may be communicatively coupled to the workstation 26 via a cable 38 that is coupled to a digital communications port or may communicate wirelessly with the workstation 26.

FIG. 2 illustrates a simplified block diagram of the PPG system 10, according to an embodiment. When the PPG system 10 is a pulse oximetry system, the emitter 16 may be configured to emit at least two wavelengths of light (for example, red and infrared) into tissue 40 of a patient. Accordingly, the emitter 16 may include a red light-emitting light source such as a red light-emitting diode (LED) 44 and an infrared light-emitting light source such as an infrared LED 46 for emitting light into the tissue 40 at the wavelengths used to calculate the patient's physiological parameters. For example, the red wavelength may be between about 600 nm and about 700 nm, and the infrared wavelength may be between about 800 nm and about 1000 nm. In embodiments where a sensor array is used in place of single sensor, each sensor may be configured to emit a single wavelength. For example, a first sensor may emit a red light while a second sensor may emit an infrared light.

As discussed above, the PPG system 10 is described in terms of a pulse oximetry system. However, the PPG system 10 may be various other types of systems. For example, the PPG system 10 may be configured to emit more or less than two wavelengths of light into the tissue 40 of the patient. Further, the PPG system 10 may be configured to emit wavelengths of light other than red and infrared into the tissue 40. As used herein, the term "light" may refer to energy produced by radiative sources and may include one or more of ultrasound, radio, microwave, millimeter wave, infrared, visible,

ultraviolet, gamma ray or X-ray electromagnetic radiation. The light may also include any wavelength within the radio, microwave, infrared, visible, ultraviolet, or X-ray spectra, and that any suitable wavelength of electromagnetic radiation may be used with the system 10. The detector 18 may be configured to be specifically sensitive to the chosen targeted energy spectrum of the emitter 16.

The detector 18 may be configured to detect the intensity of light at the red and infrared wavelengths. Alternatively, each sensor in the array may be configured to detect an intensity of a single wavelength. In operation, light may enter the detector 18 after passing through the tissue 40. The detector 18 may convert the intensity of the received light into an electrical signal. The light intensity may be directly related to the absorbance and/or reflectance of light in the tissue 40. For example, when more light at a certain wavelength is absorbed or reflected, less light of that wavelength is received from the tissue by the detector 18. After converting the received light to an electrical signal, the detector 18 may send the signal to the monitor 14, which calculates physiological parameters based on the absorption of the red and infrared wavelengths in the tissue 40.

In an embodiment, an encoder 42 may store information about the sensor 12, such as sensor type (for example, whether the sensor is intended for placement on a forehead or digit) and the wavelengths of light emitted by the emitter 16. The stored information may be used by the monitor 14 to select appropriate algorithms, lookup tables and/or calibration coefficients stored in the monitor 14 for calculating physiological parameters of a patient. The encoder 42 may store or otherwise contain information specific to a patient, such as, for example, the patient's age, weight, and diagnosis. The information may allow the monitor 14 to determine, for example, patient-specific threshold ranges related to the patient's physiological parameter measurements, and to enable or disable additional physiological parameter algorithms. The encoder 42 may, for instance, be a coded resistor that stores values corresponding to the type of sensor 12 or the types of each sensor in the sensor array, the wavelengths of light emitted by emitter 16 on each sensor of the sensor array, and/or the patient's characteristics. Optionally, the encoder 42 may include a memory in which one or more of the following may be stored for communication to the monitor 14: the type of the sensor 12, the wavelengths of light emitted by emitter 16, the particular wavelength each sensor in the sensor array is monitoring, a signal threshold for each sensor in the sensor array, any other suitable information, or any combination thereof.

Signals from the detector 18 and the encoder 42 may be transmitted to the monitor 14. The monitor 14 may include a general-purpose control unit, such as a microprocessor 48 connected to an internal bus 50. The microprocessor 48 may be configured to execute software, which may include an operating system and one or more applications, as part of performing the functions described herein. A read-only memory (ROM) 52, a random access memory (RAM) 54, user inputs 56, the display 20, and the speaker 22 may also be operatively connected to the bus 50. The control unit and/or the microprocessor 48 may include an index-determining module 49 that is configured to determine a basic fluid responsiveness index or value, as described below, and a normalization module 51 that is configured to normalize the basic fluid responsiveness index or value based on changes in vascular tone, as described below.

The RAM 54 and the ROM 52 are illustrated by way of example, and not limitation. Any suitable computer-readable media may be used in the system for data storage. Computer-

readable media are configured to store information that may be interpreted by the microprocessor 48. The information may be data or may take the form of computer-executable instructions, such as software applications, that cause the microprocessor to perform certain functions and/or computer-implemented methods. The computer-readable media may include computer storage media and communication media. The computer storage media may include volatile and non-volatile media, removable and non-removable media implemented in any method or technology for storage of information such as computer-readable instructions, data structures, program modules or other data. The computer storage media may include, but are not limited to, RAM, ROM, EPROM, EEPROM, flash memory or other solid state memory technology, CD-ROM, DVD, or other optical storage, magnetic cassettes, magnetic tape, magnetic disk storage or other magnetic storage devices, or any other medium which may be used to store desired information and that may be accessed by components of the system.

The monitor 14 may also include a time processing unit (TPU) 58 configured to provide timing control signals to a light drive circuitry 60, which may control when the emitter 16 is illuminated and multiplexed timing for the red LED 44 and the infrared LED 46. The TPU 58 may also control the gating-in of signals from the detector 18 through an amplifier 62 and a switching circuit 64. The signals are sampled at the proper time, depending upon which light source is illuminated. The received signal from the detector 18 may be passed through an amplifier 66, a low pass filter 68, and an analog-to-digital converter 70. The digital data may then be stored in a queued serial module (QSM) 72 (or buffer) for later downloading to RAM 54 as QSM 72 fills up. In an embodiment, there may be multiple separate parallel paths having amplifier 66, filter 68, and ND converter 70 for multiple light wavelengths or spectra received.

The microprocessor 48 may be configured to determine the patient's physiological parameters, such as SpO₂ and pulse rate, using various algorithms and/or look-up tables based on the value(s) of the received signals and/or data corresponding to the light received by the detector 18. The signals corresponding to information about a patient, and regarding the intensity of light emanating from the tissue 40 over time, may be transmitted from the encoder 42 to a decoder 74. The transmitted signals may include, for example, encoded information relating to patient characteristics. The decoder 74 may translate the signals to enable the microprocessor 48 to determine the thresholds based on algorithms or look-up tables stored in the ROM 52. The user inputs 56 may be used to enter information about the patient, such as age, weight, height, diagnosis, medications, treatments, and so forth. The display 20 may show a list of values that may generally apply to the patient, such as, for example, age ranges or medication families, which the user may select using the user inputs 56.

The fluid delivery device 36 may be communicatively coupled to the monitor 14. The microprocessor 48 may determine the patient's physiological parameters, such as a change or level of fluid responsiveness, and display the parameters on the display 20. In an embodiment, the parameters determined by the microprocessor 48 or otherwise by the monitor 14 may be used to adjust the fluid delivered to the patient via fluid delivery device 36.

As noted, the PPG system 10 may be a pulse oximetry system. A pulse oximeter is a medical device that may determine oxygen saturation of blood. The pulse oximeter may indirectly measure the oxygen saturation of a patient's blood (as opposed to measuring oxygen saturation directly by analyzing a blood sample taken from the patient) and changes in

blood volume in the skin. Ancillary to the blood oxygen saturation measurement, pulse oximeters may also be used to measure the pulse rate of a patient. Pulse oximeters typically measure and display various blood flow characteristics including, but not limited to, the oxygen saturation of hemoglobin in arterial blood.

A pulse oximeter may include a light sensor, similar to the sensor **12**, that is placed at a site on a patient, typically a fingertip, toe, forehead or earlobe, or in the case of a neonate, across a foot. The pulse oximeter may pass light using a light source through blood perfused tissue and photoelectrically sense the absorption of light in the tissue. For example, the pulse oximeter may measure the intensity of light that is received at the light sensor as a function of time. A signal representing light intensity versus time or a mathematical manipulation of this signal (for example, a scaled version thereof, a log taken thereof, a scaled version of a log taken thereof, and/or the like) may be referred to as the photoplethysmograph (PPG) signal. In addition, the term "PPG signal," as used herein, may also refer to an absorption signal (for example, representing the amount of light absorbed by the tissue) or any suitable mathematical manipulation thereof. The light intensity or the amount of light absorbed may then be used to calculate the amount of the blood constituent (for example, oxyhemoglobin) being measured as well as the pulse rate and when each individual pulse occurs.

The light passed through the tissue is selected to be of one or more wavelengths that are absorbed by the blood in an amount representative of the amount of the blood constituent present in the blood. The amount of light passed through the tissue varies in accordance with the changing amount of blood constituent in the tissue and the related light absorption. Red and infrared wavelengths may be used because it has been observed that highly oxygenated blood will absorb relatively less red light and more infrared light than blood with lower oxygen saturation. By comparing the intensities of two wavelengths at different points in the pulse cycle, it is possible to estimate the blood oxygen saturation of hemoglobin in arterial blood.

The PPG system **10** and pulse oximetry are further described in United States Patent Application Publication No. 2012/0053433, entitled "System and Method to Determine SpO₂ Variability and Additional Physiological Parameters to Detect Patient Status," United States Patent Application Publication No. 2010/0324827, entitled "Fluid Responsiveness Measure," and United States Patent Application Publication No. 2009/0326353, entitled "Processing and Detecting Baseline Changes in Signals," all of which are hereby incorporated by reference in their entireties.

Respiratory variation of the PPG signal may correlate with fluid responsiveness. Fluid responsiveness relates to the volume of fluid, such as blood, in the arteries, veins, and vasculature of an individual. In general, fluid responsiveness may include a measurement of the response of stroke volume, the volume of blood passing out of the heart with each heartbeat, to venous return, the volume of blood entering the heart with each heartbeat, caused by the clinical administration of fluid into the vasculature, such as through an intravenous injection. With each heartbeat, a certain amount of blood is pumped out of the heart. The more blood that fills the heart, the more blood the heart can pump out with each heartbeat. If blood volume is too low, the heart may not fully fill with blood. Therefore, the heart may not pump out as much blood with each heartbeat. Consequently, low blood volume may lead to low blood pressure, and organs and tissues may not receive enough blood to optimally and/or properly function. Monitoring fluid responsiveness allows a physician to determine whether addi-

tional fluid should be provided to a patient, such as through an intravenous fluid injection. In short fluid responsiveness represents a prediction of whether or not additional intravenous fluid may improve blood flow within a patient. Fluid responsiveness may be viewed as a response of a heart in relation to overall fluid within a patient.

Fluid responsiveness may be monitored in, for example, critically-ill patients because fluid administration plays an important role in optimizing stroke volume, cardiac output, and oxygen delivery to organs and tissues. However, clinicians need to balance central blood volume depletion and volume overloading. Critically-ill patients are generally at greater risk for volume depletion and severe hypotension is a common life-threatening condition in critically-ill patients. Conversely, administering too much fluid may induce life-threatening adverse effects, such as volume overload, systemic and pulmonary edema, and increased tissue hypoxia. Therefore, obtaining reliable information and parameters that aid clinicians in fluid management decisions may help improve patient outcomes.

FIG. **3** illustrates a PPG signal **100** over time, according to an embodiment. The PPG signal **100** is an example of a physiological signal. However, embodiments may be used in relation to various other physiological signals, such as electrocardiogram signals, phonocardiogram signals, ultrasound signals, and the like. The PPG signal **100** may be determined, formed, and displayed as a waveform by the monitor **14** (shown in FIG. **1**) that receives signal data from the PPG sensor **12** (shown in FIG. **1**). For example, the monitor **14** may receive signals from the PPG sensor **12** positioned on a finger of a patient. The monitor **14** processes the received signals, and displays the resulting PPG signal **100** on the display **28** (shown in FIG. **1**).

The PPG signal **100** may include a plurality of pulses **102a-102n** over a predetermined time period. The time period may be a fixed time period, or the time period may be variable. Moreover, the time period may be a rolling time period, such as a 5 second rolling timeframe.

Each pulse **102a-102n** may represent a single heartbeat and may include a pulse-transmitted or primary peak **104** separated from a pulse-reflected or trailing peak **106** by a dichrotic notch **108**. The primary peak **104** represents a pressure wave generated from the heart to the point of detection, such as in a finger where the PPG sensor **12** (shown in FIG. **1**) is positioned. The trailing peak **106** may represent a pressure wave that is reflected from the location proximate where the PPG sensor **12** is positioned back toward the heart.

As shown in FIG. **3**, each pulse **102a-102n** has a particular amplitude. For example, the pulse **102a** has an amplitude a_1 , while the pulse **102b** has an amplitude a_2 . The amplitudes a_1 and a_2 may differ, as shown. Indeed, the amplitudes of each pulse **102a-102n** may vary with respect to one another. In general, the overall amplitude of the PPG signal **100** over time t may modulate. The index-determining module **49** (shown in FIG. **2**) of the monitor **14** may track and store the magnitude of the amplitude modulation of the PPG signal **100** over time t . Optionally, the index-determining module **49** of the monitor **14** may track and store the magnitude of the amplitude modulation of the PPG signal **100** over time periods of varying lengths. For example, the index-determining module **49** of the monitor **14** may compare the amplitudes of neighboring pulses **102a** and **102b** and store the change in amplitude between the two pulses **102a** and **102b** as a magnitude of amplitude modulation. The index-determining module **49** of the monitor **14** may continually track and store amplitude modulation between neighboring pulses **102a-102n** over the time period t . Optionally, the index-determining module **49** of

the monitor **14** may determine an average modulation of the pulses **102a-102n** over the time period *t* and store the average modulation as a magnitude of amplitude modulation.

The PPG signal **100** also includes a baseline **110**, which may modulate over the time period *t*. The index-determining module **49** of the monitor **14** may monitor and store the baseline modulation Δb over the time period *t*. Optionally, the index-determining module **49** of the monitor **14** may determine the baseline modulation between neighboring peaks **102a-102n**. As shown in FIG. 3, however, the magnitude of baseline modulation Δb may be summed or averaged over the time period *t*.

The index-determining module **49** of the monitor **14** may determine an index of fluid responsiveness determined through the modulation of the PPG signal. For example, the index of fluid responsiveness may be determined through a ratio of amplitude modulation to baseline modulation (AM/BM). The amplitude modulation and baseline modulation may be calculated and/or otherwise determined by the index-determining module **49** or otherwise by the monitor **14** as discussed above.

It has been found that amplitude modulation relates to stroke volume. For example, amplitude modulation may be directly proportional to stroke volume. The higher the magnitude of amplitude modulation, the higher the stroke volume.

Similarly, it has been found that baseline modulation relates to venous return. For example, baseline modulation may be directly proportional to venous return. The higher the magnitude of baseline modulation, the higher the venous return.

The ratio of amplitude modulation to baseline modulation (AM/BM) provides an index of fluid responsiveness.

$$\text{BFR} = c(\text{AM}/\text{BM}) \quad \text{Equation (1)}$$

where BFR is Basic Fluid Responsiveness or a fluid responsiveness index, AM is amplitude modulation, BM is baseline modulation, and *c* represents an empirically-determined constant and/or calibration coefficient. The empirically-determined constant *c* may be a calibration value that may be determined through clinical examinations of patients. For example, a group of patients may be monitored to determine fluid responsiveness levels through traditional detection methods, such as with pressure transducers secured to arterial catheters. The traditional detection methods yield fluid responsiveness values that are correlated with particular AM and BM, which may also be monitored at the same time. The empirically-determined constant *c* may then be based on the relation between the fluid responsiveness values and AM and BM. The calibrated value *c* may then be used by the index-determining module **49** of the monitor **14** with respect to a determination of basic fluid responsiveness in other patients based on detection of AM and BM. The index-determining module **49** of the monitor **14** may store and use the value *c* as a calibration coefficient, and/or in relation to a look-up table. Optionally, the BFR may be determined simply through the ratio of AM/BM. Thus, the BFR or fluid responsiveness index may include the ratio of AM/BM and the empirically-determined coefficient *c*. As shown above, the BFR or fluid responsiveness index may be a product of the ratio of AM/BM and the empirically-determined coefficient *c*.

The BFR may be a dimensionless scalar index value. Based on Equation (1), Basic Fluid Responsiveness (BFR) is higher when the AM is high and the BM is low (relative to a predetermined neutral threshold). As AM decreases and/or BM increases, BFR decreases. If the BFR is relatively high, for example, a healthcare professional such as physician, and/or the monitor **14** through the fluid delivery device **36**, may

determine that the patient may benefit from additional fluid, such as saline, plasma, blood, water, any other fluid suitable for delivery to a patient, or any combination thereof. If the BFR is low, the physician may determine that no additional fluid is to be provided to the patient. In short, the BFR allows one to quickly and easily determine whether or not additional fluid should be administered to a patient. Alternatively, the ratio may be inverted to BM/AM, in which case fluid would be administered to a patient when the BFR is low, and fluid is not administered when the BFR is high.

Embodiments allow for a quick and simple determination of fluid responsiveness through analysis of a physiological signal, such as the PPG signal **100**. In contrast to previous systems and methods, embodiments may not require an arterial line ("A-line") having a pressure transducer to detect fluid responsiveness. Moreover, embodiments may be used to detect fluid responsiveness with patients that are spontaneously breathing and/or those that are on lung-protective low-tidal volume ventilation, in contrast to numerous previous systems and methods that typically require high-tidal ventilation in order to determine fluid responsiveness.

As explained above, fluid responsiveness may be determined through a ratio of AM/BM of the PPG signal **100**. Additionally, as described above, fluid responsiveness may also be determined through a ratio of frequency modulation to baseline modulation of a PPG signal.

FIG. 4 illustrates a PPG signal **120** over time, according to an embodiment. The PPG signal **120** is an example of a physiological signal, but various other physiological signals may be used. The PPG signal **120** includes a plurality of pulses **122a-122n**. The frequency of the pulses **122a-122n** may vary. For example, the frequency f_1 of the pulses **122a-122n** over a first period of time may vary from a frequency f_2 over a later period of time. The monitor **14** (shown in FIG. 1) may monitor and determine the frequencies f_1 and f_2 . The index-determining module **49** of the monitor **14** may then detect a magnitude of frequency modulation FM over the time period *t*. The index-determining module **49** of the monitor **14** may determine the FM between neighboring pulses **122a-122n** and calculate an aggregate FM over the time period *t*. Optionally, the index-determining module **49** of the monitor **14** may average the FM between neighboring pulses **122a-122n** over the time period *t*. The index-determining module **49** of the monitor **14** determines the magnitude of FM, which may then be used to determine fluid responsiveness, as explained below.

$$\text{BFR} = c(\text{FM}/\text{BM}) \quad \text{Equation (2)}$$

where BFR is Basic Fluid Responsiveness or a fluid responsiveness index, FM is a magnitude of frequency modulation, BM is a magnitude of baseline modulation, and *c* represents an empirically-determined constant. The empirically-determined constant *c* may be a calibration value that may be determined through clinical examinations of patients. For example, a group of patients may be monitored to determine fluid responsiveness levels through traditional detection methods, such as with pressure transducers secured to arterial catheters. The traditional detection methods yield fluid responsiveness values that are correlated with particular FM and BM, which may also be monitored at the same time. The empirically-determined calibration coefficient or constant *c* may then be based on the relation between the fluid responsiveness values and FM and BM. The calibrated value *c* may then be utilized by the monitor **14** with respect to a determination of basic fluid responsiveness in other patients based on detection of AM and BM. The index-determining module **49** of the monitor **14** may store and use the value *c* as a calibration

coefficient, and/or in relation to a look-up table. Optionally, the BFR may be determined simply through the ratio of FM/BM. Thus, the BFR or fluid responsiveness index may include the ratio of FM/BM and the empirically-determined coefficient *c*. As shown above, the BFR or fluid responsiveness index may be a product of the ratio of FM/BM and the empirically-determined coefficient *c*.

The FM may be directly proportional to the stroke volume. For example, FM from respiratory sinus arrhythmia may be directly proportional to stroke volume because heart rate changes to maintain the same cardiac output from reduced stroke volume. Thus, as stroke volume increases, the magnitude of FM may increase. BM may be directly proportional to venous return. Thus, as venous return increases, the magnitude of BM may increase.

The BFR may be a dimensionless scalar index value. Based on Equation (2), Basic Fluid Responsiveness (BFR) is higher when the FM is high and the BM is low (relative to a predetermined neutral threshold). As FM decreases and/or BM increases, BFR decreases. If the BFR is relatively high, a physician may determine that the patient may benefit from additional fluid. If the BFR is low, the physician may determine that no additional fluid is to be provided to the patient. In short, the BFR allows one to quickly and easily determine whether or not additional fluid should be administered to a patient. Alternatively, the ratio may be inverted to BM/FM, in which case fluid would be administered to a patient when the BFR is low, and fluid is not administered when the BFR is high.

Embodiments allow for a quick and simple determination of fluid responsiveness through analysis of the PPG signal **120**. In contrast to previous systems and methods, embodiments do not require an arterial line ("A-line") having a pressure transducer to detect fluid responsiveness. Moreover, embodiments may be used to detect fluid responsiveness with patients that are spontaneously breathing and/or those that are on lung-protective low-tidal volume ventilation, in contrast to numerous previous systems and methods that typically require high-tidal ventilation in order to determine fluid responsiveness.

Referring to FIGS. **3** and **4**, embodiments provide a system and method of determining fluid responsiveness through analysis of a physiological signal, such as a PPG signal. A fluid responsiveness index may be determined through a ratio of AM/BM, BM/AM, FM/BM, and/or BM/FM. The fluid responsiveness index may be a dimensionless scalar value. A fluid responsiveness index spectrum may range from low, such as 1, to high, such as 10. A physician may determine whether or not additional fluid should be provided to a patient through a review of the fluid responsiveness index. Alternatively, the monitor **14** may automatically determine whether fluid should be provided to a patient through an automated analysis of the fluid responsiveness index. For example, if the fluid responsiveness index is high (such as between 6 and 10), then the physician and/or monitor **14** may determine that the patient should be provided with additional fluid. For example, the monitor **14** may automatically detect the fluid responsiveness index is high, and then automatically provide fluid to the patient through the fluid delivery device **36** (shown in FIG. **1**). If the fluid responsiveness signal is low (such as between 1-4), then the physician may refrain from administering additional fluid to the patient.

As noted, the fluid responsiveness index may be determined simply and quickly through an analysis of the PPG signal, without the need for an A-line, or any other sensor (such as an intra-arterial or intra-venous pressure transducer). The system and method may detect a signal through the PPG

sensor **12**, and analyze the PPG signal to determine AM (and/or FM) and BM to yield a fluid responsiveness index value.

FIG. **5** illustrates a flow chart of a method of determining fluid responsiveness, according to an embodiment. At **130**, the process begins by detecting a PPG signal through a PPG sensor. Then, at **132**, the PPG signal is analyzed to determine AM and/or FM. Similarly, at **134**, the PPG signal is analyzed to determine BM. Then, at **136**, a ratio of (AM or FM)/BM is formed. At **138**, a monitor, such as the monitor **14** of FIG. **1**, and/or a physician, for example, determines whether the ratio exceeds (or is below) a predetermined threshold. For example, if the index spectrum ranges 1-10, with 1 being low, and 10 being high, the predetermined neutral threshold may be 5. If the index value does exceed the threshold, at **140**, additional fluid is administered to the patient. The process then returns to **130**. If, however, the index value does not exceed the threshold, at **142**, additional fluid is not administered to the patient, and the process returns to **130**.

Referring to FIGS. **1-5**, the magnitude of respiratory modulation, whether AM, FM, or BM, may also be affected by changes in vascular tone, which relates to the elasticity and hardness of vasculature, such as the arteries. Vascular resistance relates to the resistance to flow that is overcome to push blood through the circulatory system. Arterial compliance, for example relates to a measure of the ability of arteries to allow fluid to pass therethrough.

Vascular tone may vary at different points of the body. For example, the vascular tone at the fingertips, where the PPG sensor **12** (shown in FIG. **1**) may be located, may be different than at a location closer to the heart. Ambient temperature and drugs, for example, may affect vascular tone in that both may affect the degree of constriction of the blood vessels.

In cold conditions, for example, peripheral vascular tone may be tight and resistant to blood flow. As the ambient temperature increases, the vascular tone may open and be less resistant to blood flow. If blood vessels are closed and tight, vascular resistance increases, and a reflected pulse is stronger as a greater amount of the reflected pulse is reflected back, as compared to when blood vessels are open and flexible.

Additionally, a PPG signal may be sensitive to motion artifacts. The motion artifacts may be caused by patient and/or PPG sensor movement. Motion artifacts may be caused by hand or finger tremors, coughing, changes in breathing patterns, such as a yawn, a PPG sensor being tugged by a connected cable, and/or the like.

FIG. **6** illustrates a pulse **200** of a PPG signal, according to an embodiment. While the pulse **200** is described as a component of a PPG signal, the pulse **200** may be a component of various other physiological signals. The pulse **200** represents a single heartbeat and may include a primary peak **202** separated from a trailing peak **204** by a dichrotic notch **206**. The primary peak **202** represents a pressure wave generated from the heart to the point of detection, such as in a finger where the PPG sensor **12** (shown in FIG. **1**) is positioned. The trailing peak **204** may represent a pressure wave that is reflected from the location proximate where the PPG sensor **12** is positioned back toward the heart.

The magnitude of AM, BM, and/or FM may be affected by changes in vascular tone and/or motion artifacts. Changes in vascular tone and/or motion artifacts may affect the ratio of AM/BM or FM/BM. Accordingly, changes in the BFR may be due to changes in vascular tone and/or motion artifacts, rather than changes in fluid responsiveness. For example, when arterial compliance in large arteries decreases, the AM of the PPG signal increases relative to the BM due to increased steepening of the propagating pressure wave. When

peripheral vascular resistance increases, the AM of the PPG signal may also increase relative to the BM due to peak amplification from an increased reflected pulse.

The BFR or fluid responsiveness index may be corrected or normalized to account for changes in vascular tone and/or motion artifacts through analysis of the PPG signal, as described below. Several metrics of a PPG pulse may be related to arterial compliance, peripheral vascular resistance, and/or motion artifacts.

The normalization module **51** (shown in FIG. 2) of the monitor **14** may compare amplitudes of the peaks **202** and **204** to determine peripheral vasculature resistance. For example, the ratio of the amplitude A_t of the trailing peak **204** to A_p of the primary peak **202** may yield a relative amplitude ratio (RAR).

$$\text{RAR} = A_t / A_p \quad \text{Equation (3)}$$

The RAR may be directly proportional to peripheral vascular resistance. Thus, as peripheral vascular resistance increases, A_t increases and/or A_p decreases. Conversely, as peripheral vascular resistance decreases, A_t decreases and/or A_p increases. The RAR provides an indication of peripheral vascular resistance. The normalization module **51** may use the RAR to normalize the fluid responsiveness index, or BFR.

Further, the width W of the primary peak **202** may be directly proportional to arterial compliance. For example, arterial compliance increases with increasing width W . Increased arterial compliance is directly proportional to vascular resistance. Thus, a high degree of arterial compliance yields a low degree of vascular resistance. Conversely, a low degree of arterial compliance yields a high degree of vascular resistance. Therefore, the greater the width W of the primary peak **202**, the higher degree of arterial compliance and the lower degree of vascular resistance. The W may be used to normalize the fluid responsiveness index, or BFR. The normalization module **51** may use W to normalize the fluid responsiveness index, or BFR.

Additionally, the time Δt between the primary peak **202** and the trailing peak **204** may be directly proportional to arterial compliance. The time Δt may be measured from a fiducial **210** of the primary peak **202** to a fiducial **212** of the trailing peak **204**. The fiducial **210** may be selected as the highest point on the primary peak **202**, while the fiducial **212** may be selected as the highest point on the trailing peak **204**. However, the fiducials **210** and **212** may be at various other locations on the peaks **202** and **204**, respectively. Alternatively, the time Δt may be determined by a time delay between a fiducial on the PPG and another fiducial on: another PPG at a different site (such as a forehead or foot sensor), a phonocardiogram, an electrocardiogram, a pressure transducer on a radial artery, and/or the like.

Because the time Δt may be directly proportional to arterial compliance, as Δt increases, so, too, does arterial compliance. Conversely, as Δt decreases, so too does arterial compliance. The Δt may be used to normalize the fluid responsiveness index, or BFR. The normalization module **51** may measure Δt and use Δt to normalize the fluid responsiveness index, or BFR.

Additionally, a pleth modulation ratio (PMR) may be expressed by AC/DC. The AC physiological component may generally be a variable component related to cardiac synchronous changes in the blood volume with each heartbeat. The AC component is typically superimposed on a constant DC baseline. The PMR may be inversely proportional to vascular resistance. Thus, as PMR increases, vascular resistance decreases. Conversely, as PMR decreases, vascular resistance increases. The PMR may be used to normalize the fluid

responsiveness index, or BFR. The normalization module **51** may determine PMR and use PMR to normalize the fluid responsiveness index, or BFR.

The normalization module **51** of the monitor **14** may utilize the RAR, Δt , PMR, and/or W to normalize the BFR or fluid responsiveness index. As shown below, the vascular tone and/or motion artifact corrected fluid responsiveness index (VTC) may be determined as follows:

$$\text{VTC} = c(\text{AM}/\text{BM} \times G/H) \quad \text{Equation (4)}$$

$$\text{VTC} = c(\text{FM}/\text{BM} \times H) \quad \text{Equation (5)}$$

where c is a constant, as described above, AM is Amplitude Modulation, BM is Baseline Modulation, FM is frequency modulation, and G and H may be constants and/or calibration coefficients. As shown above, VTC may be determined through AM/BM or FM/BM. G may be a function of W and/or Δt , while H may be a function of RAR and/or 1/PMR. For example, G may be a calibrated value that is a function of W or Δt , while H may be a calibrated value that is a function of RAR and/or 1/PMR. Optionally, G and H may each be functions of one or more of RAR, Δt , PMR, and/or W .

In an embodiment, G may be directly or inversely proportional to W and/or Δt . In an embodiment, H may directly or inversely proportional to RAR and/or 1/PMR. Each of c , G , and H may be empirically-derived, non-linear functions.

Thus, as shown above, the BFR or fluid responsiveness index may be normalized through calculations of RAR, Δt , PMR, and/or W . The normalization of the BFR or fluid responsiveness index accounts for differences in vascular tone and/or motion artifacts.

The normalization module **51** of the monitor **14** may analyze a plurality of PPG pulses **200** to determine RAR, Δt , PMR, and/or W . For example, a plurality of PPG pulses **200** over a particular time period may be averaged to yield an average pulse, which the normalization module **51** of the monitor **14** analyzes to determine RAR, Δt , PMR, and/or W . Alternatively, the normalization module **51** may analyze each and every pulse over a particular time period to determine RAR, Δt , PMR, and/or W in relation to each and every pulse.

FIG. 7 illustrates a flow chart of a method normalizing a fluid responsiveness index, according to an embodiment. At **250**, a PPG signal is detected, as explained above. At **252**, the PPG signal is analyzed to determine changes in vascular tone and/or motion artifacts. Changes in vascular tone, for example may be determined by determining one or more of a relative amplitude ratio (RAR) of the PPG signal at **254**, a width W of the primary peak of the PPG signal at **256**, Δt between the primary peak and the trailing peak at **258**, and/or the pleth modulation ratio (PMR) of the PPG signal at **260**. Then, the process continues to **262**, in which constants, functions, calibrated values, or the like related to one or more of PAR, W , Δt , and PMR are applied to the basic fluid responsiveness (BFR) value or index in order to normalize the value or index. The process may then return to **250**.

It will be understood that the present disclosure is applicable to any suitable physiological signals and that PPG signals are used for illustrative purposes. Those skilled in the art will recognize that the present disclosure has wide applicability to other signals including, but not limited to other physiological signals (for example, electrocardiogram, electroencephalogram, electrogastrogram, electromyogram, heart rate signals, pathological sounds, ultrasound, or any other suitable biosignal) and/or any other suitable signal, and/or any combination thereof.

Various embodiments described herein provide a tangible and non-transitory (for example, not an electric signal)

machine-readable medium or media having instructions recorded thereon for a processor or computer to operate a system to perform one or more embodiments of methods described herein. The medium or media may be any type of CD-ROM, DVD, floppy disk, hard disk, optical disk, flash RAM drive, or other type of computer-readable medium or a combination thereof.

The various embodiments and/or components, for example, the control units, modules, or components and controllers therein, also may be implemented as part of one or more computers or processors. The computer or processor may include a computing device, an input device, a display unit and an interface, for example, for accessing the Internet. The computer or processor may include a microprocessor. The microprocessor may be connected to a communication bus. The computer or processor may also include a memory. The memory may include Random Access Memory (RAM) and Read Only Memory (ROM). The computer or processor further may include a storage device, which may be a hard disk drive or a removable storage drive such as a floppy disk drive, optical disk drive, and the like. The storage device may also be other similar means for loading computer programs or other instructions into the computer or processor.

As used herein, the term “computer” or “module” may include any processor-based or microprocessor-based system including systems using microcontrollers, reduced instruction set computers (RISC), application specific integrated circuits (ASICs), logic circuits, and any other circuit or processor capable of executing the functions described herein. The above examples are exemplary only, and are thus not intended to limit in any way the definition and/or meaning of the term “computer”.

The computer or processor executes a set of instructions that are stored in one or more storage elements, in order to process input data. The storage elements may also store data or other information as desired or needed. The storage element may be in the form of an information source or a physical memory element within a processing machine.

The set of instructions may include various commands that instruct the computer or processor as a processing machine to perform specific operations such as the methods and processes of the various embodiments of the subject matter described herein. The set of instructions may be in the form of a software program. The software may be in various forms such as system software or application software. Further, the software may be in the form of a collection of separate programs or modules, a program module within a larger program or a portion of a program module. The software also may include modular programming in the form of object-oriented programming. The processing of input data by the processing machine may be in response to user commands, or in response to results of previous processing, or in response to a request made by another processing machine.

As used herein, the terms “software” and “firmware” are interchangeable, and include any computer program stored in memory for execution by a computer, including RAM memory, ROM memory, EPROM memory, EEPROM memory, and non-volatile RAM (NVRAM) memory. The above memory types are exemplary only, and are thus not limiting as to the types of memory usable for storage of a computer program.

As described, embodiments disclose systems and methods for determining a basic fluid responsiveness index or value through analysis of a physiological signal, such as a PPG signal. Embodiments allow for quick and simple determination of fluid responsiveness through analysis of the PPG signal. In contrast to previous systems and methods, embodi-

ments may not require an arterial line (“A-line”) having a pressure transducer to detect fluid responsiveness. Moreover, embodiments may be used to detect fluid responsiveness with patients that are spontaneously breathing and/or those that are on lung-protective low-tidal volume ventilation, in contrast to previous systems and methods that typically require high-tidal ventilation in order to determine fluid responsiveness. Further, embodiments provide systems and methods for normalizing a basic fluid responsiveness index or value, such as by accounting for changes in vascular tone and/or motion artifacts.

While various spatial and directional terms, such as top, bottom, lower, mid, lateral, horizontal, vertical, front, and the like may be used to describe embodiments, it is understood that such terms are merely used with respect to the orientations shown in the drawings. The orientations may be inverted, rotated, or otherwise changed, such that an upper portion is a lower portion, and vice versa, horizontal becomes vertical, and the like.

It is to be understood that the above description is intended to be illustrative, and not restrictive. For example, the above-described embodiments (and/or aspects thereof) may be used in combination with each other. In addition, many modifications may be made to adapt a particular situation or material to the teachings without departing from its scope. While the dimensions, types of materials, and the like described herein are intended to define the parameters of the disclosure, they are by no means limiting and are exemplary embodiments. Many other embodiments will be apparent to those of skill in the art upon reviewing the above description. The scope of the disclosure should, therefore, be determined with reference to the appended claims, along with the full scope of equivalents to which such claims are entitled. In the appended claims, the terms “including” and “in which” are used as the plain-English equivalents of the respective terms “comprising” and “wherein.” Moreover, in the following claims, the terms “first,” “second,” and “third,” etc. are used merely as labels, and are not intended to impose numerical requirements on their objects. Further, the limitations of the following claims are not written in means-plus-function format and are not intended to be interpreted based on 35 U.S.C. §112, sixth paragraph, unless and until such claim limitations expressly use the phrase “means for” followed by a statement of function void of further structure.

What is claimed is:

1. A medical monitor for determining a fluid responsiveness index of a subject, the monitor comprising:
 - an electrical input to receive a physiological signal from a sensor;
 - a processor configured to:
 - determine a value indicative of fluid responsiveness based on modulations of the physiological signal;
 - determine a value indicative of changes in vascular tone of the subject based on one or more metrics of the physiological signal; and
 - determine a fluid responsiveness index of the subject based on the value indicative of fluid responsiveness and the value indicative of changes in vascular tone.
2. The monitor of claim 1, wherein the one or more metrics comprise a relative amplitude ratio of the physiological signal.
3. The monitor of claim 1, wherein the one or more metrics comprise a width of at least one primary peak of the physiological signal.

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4. The monitor of claim 1, wherein the one or more metrics comprise a time difference between at least one primary peak of the physiological signal and a trailing peak of the physiological signal.

5. The monitor of claim 1, wherein the one or more metrics comprise a pleth modulation ratio of the physiological signal.

6. The monitor of claim 1, wherein the processor is configured to determine the fluid responsiveness index of the subject by normalizing the value indicative of fluid responsiveness based on the value indicative of changes in vascular tone of the subject.

7. The monitor of claim 1, wherein the monitor comprise a pulse oximetry monitor, wherein the physiological signal comprises a photoplethysmogram (PPG) signal, and wherein the sensor comprises a pulse oximetry sensor.

8. The monitor of claim 7, wherein the one or more metrics comprise at least one of a relative amplitude ratio of the PPG signal, a width of at least one primary peak of the PPG signal, a time difference between at least one primary peak of the physiological signal and a trailing peak of the PPG signal, and a pleth modulation ratio of the PPG signal.

9. The monitor of claim 1, further comprising a fluid delivery device operatively connected to the monitor, wherein the monitor determines whether or not to administer fluid to the subject from the fluid delivery device based on the fluid responsiveness index.

10. The monitor of claim 1, further comprising a display operatively connected to the processor, wherein the processor is further configured to output a prompt to the display regarding whether or not to administer fluid to the subject, based on the fluid responsiveness index.

11. A method for determining a fluid responsiveness index of a subject from a physiological signal, the method comprising:

receiving a photoplethysmograph (PPG) signal from a pulse oximetry sensor at a pulse oximetry monitor;

determining, using the pulse oximetry monitor, a fluid responsiveness value based on the PPG signal;

determining, using the pulse oximetry monitor, a value indicative of changes in vascular tone of the subject based on the PPG signal; and

determining, using the pulse oximetry monitor, a fluid responsiveness index of the subject based on the fluid responsiveness value and the value indicative of changes in vascular tone of the subject.

12. The method of claim 11, wherein the value indicative of changes in vascular tone of the subject comprises at least one of a relative amplitude ratio of the PPG signal, a width of at least one primary peak of the PPG signal, a time difference between at least one primary peak of the physiological signal and a trailing peak of the PPG signal, and a pleth modulation ratio of the PPG signal.

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13. The method of claim 11, wherein the value indicative of changes in vascular tone comprises a relative amplitude ratio of the PPG signal.

14. The method of claim 11, wherein the value indicative of changes in vascular tone comprises a width of at least one primary peak of the PPG signal.

15. The method of claim 11, wherein the value indicative of changes in vascular tone comprises a time difference between at least one primary peak of the physiological signal and a trailing peak of the PPG signal.

16. The method of claim 11, wherein the value indicative of changes in vascular tone comprises a pleth modulation ratio of the PPG signal.

17. The method of claim 11, wherein determining, using the pulse oximetry monitor, the fluid responsiveness index of the subject based on the fluid responsiveness value and the value indicative of changes in vascular tone of the subject comprises normalizing the value indicative of fluid responsiveness based on the value indicative of changes in vascular tone of the subject.

18. The method of claim 11, further comprising:

determining, using the pulse oximetry monitor, whether or not to administer fluid to the subject based on the fluid responsiveness index; and

administering, using a fluid delivery device coupled to the pulse oximetry monitor, fluid to the subject when the pulse oximetry monitor determines to administer fluid.

19. A tangible and non-transitory computer readable medium that includes one or more sets of instructions configured to direct a computer to:

receive a photoplethysmograph (PPG) signal from a pulse oximetry sensor secured to a subject;

determine a fluid responsiveness value based on the PPG signal;

determine a value indicative of changes in vascular tone of the subject based on the PPG signal; and

determine a fluid responsiveness index of the subject based on the fluid responsiveness value and the value indicative of changes in vascular tone of the subject.

20. The tangible and non-transitory computer readable medium of claim 19, further configured to direct the computer to determine whether or not to administer fluids based on the fluid responsiveness index.

21. The tangible and non-transitory computer readable medium of claim 19, wherein the value indicative of changes in vascular tone of the subject comprises at least one of a relative amplitude ratio of the PPG signal, a width of at least one primary peak of the PPG signal, a time difference between at least one primary peak of the physiological signal and a trailing peak of the PPG signal, and a pleth modulation ratio of the PPG signal.

* * * * *

专利名称(译)	用于检测患者的流体反应性的系统和方法		
公开(公告)号	US9402573	公开(公告)日	2016-08-02
申请号	US14/730150	申请日	2015-06-03
[标]申请(专利权)人(译)	柯惠有限合伙公司		
申请(专利权)人(译)	COVIDIEN LP		
当前申请(专利权)人(译)	COVIDIEN LP		
[标]发明人	SU MARK WATSON JAMES NICHOLAS ADDISON PAUL STANLEY		
发明人	SU, MARK WATSON, JAMES NICHOLAS ADDISON, PAUL STANLEY		
IPC分类号	A61B5/1455 A61B5/02 A61B5/024 A61B5/00 A61B5/026 A61B5/1495 A61B5/0245 A61B5/0205 A61B5/0295 A61B6/00 A61B5/05 A61B8/06 A61B5/0432		
CPC分类号	A61B5/7214 A61B5/02416 A61B5/0245 A61B5/026 A61B5/14552 A61B5/1495 A61B5/7225 A61B5/7278 A61B5/486 A61B5/4839 A61B5/0295 A61B5/0261 A61B5/0205 A61B5/02007 A61B5/742 A61B6/507 A61B8/06 A61B5/0059 A61B5/0432 A61B5/0507 A61B5/6826		
审查员(译)	门德斯MANUEL		
其他公开文献	US20150265196A1		
外部链接	Espacenet USPTO		

摘要(译)

系统被配置为根据生理信号确定患者的流体响应指数。该系统可包括配置成固定到患者的解剖部分的传感器，以及可操作地连接到传感器的监视器。传感器配置为感测患者的生理特征。监视器配置为从传感器接收生理信号。监测器可包括指数确定模块，其配置成通过形成生理信号的幅度或频率调制与生理信号的基线调制中的一者或两者的比率来确定流体响应性指数。

