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(54) **BODY-WORN VITAL SIGN MONITOR**

AM KÖRPER GETRAGENER LEBENSZEICHENMONITOR

DISPOSITIF À PORTER SUR SOI POUR LA SURVEILLANCE DE SIGNES VITAUX

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(73) Proprietor: **Sotera Wireless, Inc.**
San Diego, CA 92121 (US)

(72) Inventors:
• **MOON, Jim**
Portland
OR 97229-8028 (US)
• **BANET, Matt**
Kihei
HI 96753 (US)

- **TROMMER, Gunnar**
Encinitas
CA 92024 (US)
- **DIBATTISTA, Caroline**
San Diego
CA 92037 (US)
- **WATLINGTON, Tom**
La Mesa
CA 91941 (US)
- **DHILLON, Marshal**
San Diego
CA 92127 (US)
- **McCOMBIE, Devin**
San Diego
CA 92130 (US)

(74) Representative: **Schiweck, Weinzierl & Koch**
Patentanwälte Partnerschaft mbB
Landsberger Straße 98
80339 München (DE)

(56) References cited:
US-A1- 2002 072 859 US-A1- 2004 267 099
US-A1- 2006 265 246 US-A1- 2007 188 323
US-A1- 2007 188 323 US-A1- 2007 193 834
US-A1- 2008 004 904 US-A1- 2008 033 255
US-A1- 2008 195 735 US-A1- 2008 221 399
US-A1- 2008 275 349 US-A1- 2009 112 281
US-A1- 2009 259 113

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Description

BACKGROUND OF THE INVENTION

[0001] The present invention relates to medical devices for monitoring vital signs, e.g., arterial blood pressure.

[0002] Conventional vital sign monitors are used throughout the hospital, and are particularly commonplace in high-acuity areas such as the intensive care unit (ICU), emergency department (ED), or operating room (OR). Patients in these areas are generally sick and require a high degree of medical attention, typically provided by a relatively high ratio of clinicians compared to lower-acuity areas of the hospital. Outside the ICU and OR, clinicians typically measure vital signs such as systolic, diastolic, and mean arterial blood pressures (SYS, DIA, MAP), respiratory rate (RR), oxygen saturation (SpO₂), heart rate (HR), and temperature (TEMP) with portable or wall-mounted vital sign monitors. It can be difficult to effectively monitor patients in this way, however, because measurements are typically made every few hours, and the patients are often ambulatory and not constrained to a single hospital room. This poses a problem for conventional vital sign monitors, which are typically heavy and unwieldy, as they are not intended for the ambulatory population. To make a measurement, a patient is typically tethered to the monitor with a series of tubes and wires. Some companies have developed ambulatory vital sign monitors with limited capabilities (e.g. cuff-based blood pressure using oscillometry and SpO₂ monitoring), but typically these devices only make intermittent, rather than continuous, measurements. And even these measurements tend to work best on stationary patients, as they are easily corrupted by motion-related artifacts.

[0003] Most vital signs monitors feature a user interface that shows numerical values and waveforms associated with the vital signs, alarm parameters, and a 'service menu' that can be used to calibrate and maintain the monitor. Some monitors have internal wireless cards that communicate with a hospital network, typically using protocols such as 802.11b/g.

[0004] One of the most important parameters measured with vital signs monitors is blood pressure. In critical care environments like the ICU and OR, blood pressure can be continuously monitored with an arterial catheter inserted in the patient's radial or femoral artery. Alternatively, blood pressure can be measured intermittently with a cuff using oscillometry, or manually by a clinician using auscultation. Most vital sign monitors perform both catheter and cuff-based measurements of blood pressure. Blood pressure can also be monitored continuously with a technique called pulse transit time (PTT), defined as the transit time for a pressure pulse launched by a heartbeat in a patient's arterial system. PTT has been shown in a number of studies to correlate to SYS, DIA, and MAP. In these studies, PTT is typically measured with a conventional vital signs monitor that includes sep-

arate modules to determine both an electrocardiogram (ECG) and SpO₂. During a PTT measurement, multiple electrodes typically attach to a patient's chest to determine a time-dependent ECG component characterized by a sharp spike called the 'QRS complex'. The QRS complex indicates an initial depolarization of ventricles within the heart and, informally, marks the beginning of the heartbeat and a pressure pulse that follows.

[0005] SpO₂ is typically measured with a bandage or clothespin-shaped sensor that clips to a patient's finger and includes optical systems operating in both the red and infrared spectral regions. A photodetector measures radiation emitted from the optical systems that transmits through the patient's finger. Other body sites, e.g., the ear, forehead, and nose, can also be used in place of the finger. During a measurement, a microprocessor analyzes both red and infrared radiation detected by the photodetector to determine the patient's blood oxygen saturation level and a time-dependent waveform called a photoplethysmograph (PPG). Time-dependent features of the PPG indicate both pulse rate and a volumetric absorbance change in an underlying artery caused by the propagating pressure pulse.

[0006] Typical PTT measurements determine the time separating a maximum point on the QRS complex (indicating the peak of ventricular depolarization) and a foot of the PPG waveform (indicating the beginning the pressure pulse). PTT depends primarily on arterial compliance, the propagation distance of the pressure pulse (which is closely approximated by the patient's arm length), and blood pressure. To account for patient-dependent properties, such as arterial compliance, PTT-based measurements of blood pressure are typically 'calibrated' using a conventional blood pressure cuff and oscillometry. Typically during the calibration process the blood pressure cuff is applied to the patient, used to make one or more blood pressure measurements, and then left for future measurements. Going forward, the calibration measurements are used, along with a change in PTT, to measure the patient's continuous blood pressure (cNIBP). PTT typically relates inversely to blood pressure, i.e., a decrease in PTT indicates an increase in blood pressure.

[0007] A number of issued U.S. Patents describe the relationship between PTT and blood pressure. For example, U.S. Patents 5,316,008; 5,857,975; 5,865,755; and 5,649,543 each describe an apparatus that includes conventional sensors that measure both ECG and PPG waveforms which are then processed to determine PTT. The document US 2008/033255 A1 discloses a method to initiate communication between two medical devices, e.g. by a mechanical manipulation of one of the two devices.

SUMMARY OF THE INVENTION

[0008] To improve the safety of hospitalized patients, particularly those in lower-acuity areas, it is desirable to

have a body-worn monitor that continuously measures all vital signs from a patient, provides tools for effectively monitoring the patient, and wirelessly communicates with a hospital's information technology (IT) network. Preferably the monitor operates algorithms featuring: 1) a low percentage of false positive alarms/alerts; and 2) a high percentage of true positive alarms/alerts. The term 'alarm/alert', as used herein, refers to an audio and/or visual alarm generated directly by a monitor worn on the patient's body, or alternatively a remote monitor (e.g., a central nursing station). To accomplish this, the invention provides a body-worn monitor that measures a patient's vital signs (e.g. cNIBP, SpO2, HR, RR, and TEMP) while simultaneously characterizing their activity state (e.g. resting, walking, convulsing, falling) and posture (upright, supine). The body-worn monitor processes this information to minimize corruption of the vital signs and associated alarms/alerts by motion-related artifacts.

[0009] The body-worn monitor features a graphical user interface (GUI) rendered on a touchpanel display that facilitates a number of features to simplify and improve patient monitoring and safety in both the hospital and home. For example, the monitor features a battery-powered, wrist-worn transceiver that processes motion-related signals generated with an internal motion sensor (e.g. an accelerometer). When the transceiver's battery runs low, the entire unit can be swapped out by simply 'bumping' the original transceiver with a new one having a fully charged battery. Accelerometers within the transceivers detect the 'bump', digitize the corresponding signals, and wirelessly transmit them to a patient data server (PDS) within the hospital's network. There, the signals are analyzed and patient information (e.g. demographic and vital sign data) formerly associated with the original transceiver is re-associated with the new transceiver. A clinician can view the data using a computer functioning as a remote viewing device (RVD), such as a conventional computer on wheels (COW).

[0010] The body-worn monitor additionally includes a speaker, microphone, and software that collectively facilitate voice over IP (VOIP) communication. With these features, the wrist-worn transceiver can be used as a two-way communicator allowing, e.g., the patient to alert a clinician during a time of need. Additionally, during medical procedures or diagnoses, the clinician can enunciate annotations directly into the transceiver. These annotations along with vital sign information are wirelessly transmitted to the PDS and ultimately a hospital's electronic medical records (EMR) system, where they are stored and used for post-hoc analysis of the patient. In a related application, the transceiver includes a barcode scanner that, prior to administering medications, scans barcodes associated with the patient, clinician, and medications. The transceiver sends the decoded barcode information back to the PDS, where a software program analyzes it to determine that there are no errors in the medication or the rate at which it is delivered. A signal is then sent from the PDS to the GUI, clearing the clinician to admin-

ister the medications.

[0011] The body-worn monitor can determine a patient's location in addition to their vital signs and motion-related properties. Typically, the location-determining sensor and the wireless transceiver operate on a common wireless system, e.g. a wireless system based on 802.11a/b/g/ 802.15.4, or cellular protocols. In this case a location is determined by processing the wireless signal with one or more algorithms known in the art. These include, for example, triangulating signals received from at least three different wireless base stations, or simply estimating a location based on signal strength and proximity to a particular base station. In still other embodiments the location sensor includes a conventional global positioning system (GPS).

[0012] VOIP-based communications typically take place between the body-worn monitor and a remote computer or telephone interfaced to the PDS. The location sensor, wireless transceiver, and first and second voice interfaces can all operate on a common wireless system, such as one of the above-described systems based on 802.11 or cellular protocols. In embodiments, the remote computer, for example, can be a monitor that is essentially identical to the transceiver worn by the patient, and can be carried or worn by a clinician. In this case the monitor associated with the clinician features a display wherein the user can select to display information (e.g. vital signs, location, and alarms) corresponding to a particular patient. This monitor can also include a voice interface so the clinician can communicate with the patient.

[0013] The wrist-worn transceiver's touchpanel display can render a variety of different GUIs that query the patient for their pain level, test their degree of 'mentation', i.e. mental activity, and perform other functions to assist and improve diagnosis. Additionally, the transceiver supports other GUIs that allow the patient to order food within the hospital, change the channel on their television, select entertainment content, play games, etc. To help promote safety in the hospital, the GUI can also render a photograph or video of the patient or, in the case of neonatal patients, their family members.

[0014] The body-worn monitor can include a software framework that generates alarms/alerts based on threshold values that are either preset or determined in real time. The framework additionally includes a series of 'heuristic' rules that take the patient's activity state and motion into account, and process the vital signs accordingly. These rules, for example, indicate that a walking patient is likely breathing and has a regular heart rate, even if their motion-corrupted vital signs suggest otherwise.

[0015] The body-worn monitor features a series of sensors that attach to the patient to measure time-dependent PPG, ECG, ACC, oscillometric (OSC), and impedance pneumography (IP) waveforms. A microprocessor (CPU) within the monitor continuously processes these waveforms to determine the patient's vital signs, degree of motion, posture and activity level. Sensors that measure

these signals typically send digitized information to the wrist-worn transceiver through a serial interface, or bus, operating on a controlled area network (CAN) protocol. The CAN bus is typically used in the automotive industry, and allows different electronic systems to effectively and robustly communicate with each other with a small number of dropped packets, even in the presence of electrically noisy environments. This is particularly advantageous for ambulatory patients that may generate signals with large amounts of motion-induced noise.

[0016] Blood pressure is determined continuously and non-invasively using a technique, based on PTT, which does not require any source for external calibration. This technique, referred to herein as the 'Composite Technique', determines blood pressure using PPG, ECG, and OSC waveforms. The Composite Technique is described in detail in the document US 2010/0168589 A1. In other aspects, PTT can be calculated from time-dependent waveforms other than the ECG and PPG, and then processed to determine blood pressure. In general, PTT can be calculated by measuring a temporal separation between features in two or more time-dependent waveforms measured from the human body. For example, PTT can be calculated from two separate PPGs measured by different optical sensors disposed on the patient's fingers, wrist, arm, chest, ear, or virtually any other location where an optical signal can be measured using a transmission or reflection-mode optical configuration. In other embodiments, PTT can be calculated using at least one time-dependent waveform measured with an acoustic sensor, typically disposed on the patient's chest. Or it can be calculated using at least one time-dependent waveform measured using a pressure sensor, typically disposed on the patient's bicep, wrist, or finger. The pressure sensor can include, for example, a pressure transducer, piezoelectric sensor, actuator, polymer material, or inflatable cuff.

[0017] Specifically, in one aspect, the invention provides a method for monitoring a patient according to present claim 1.

[0018] In embodiments, both the first and second motion sensors are accelerometers that generate time-dependent waveforms (e.g. ACC waveforms). Contacting the two transceivers typically generates waveforms that include individual 'pulses' (e.g. a sharp spike) caused by rapid acceleration and deceleration detected by the respective accelerometers. Typically the pulses are within waveforms generated along the same axes in both transceivers. The pulses can be collectively processed (using, e.g., an autocorrelation algorithm) to determine that they are generated during a common period of time. In embodiments, amplitudes of the first and second pulses are required to exceed a pre-determined threshold value in order for the second transceiver to replace the first transceiver. Pulses that meet this criterion are wirelessly transmitted to a remote server, where they are processed as described above. If the server determines that the second transceiver is ready to replace the first transceiver, it

transmits instruction information to the transceivers to guide the replacement process. This instruction information, for example, is displayed by the GUIs of both transceivers. Once the replacement process is complete, vital sign information measured by the second transceiver is stored along with that measured by the first transceiver in a computer memory (e.g. a database) on the remote computer. The vital sign information can include conventional vital signs (e.g. HR, SYS, DIA, RR, and TEMP), along with the time-dependent waveforms used to calculate the vital signs (e.g. PPG, ECG, OSC, IP) and motion-related properties (ACC). Patient demographic information (e.g. name, gender, weight, height, date of birth) can also be associated with both the first and second sets of vital sign information.

[0019] In another aspect, the invention provides a method for pairing a patient monitor with a remote display device (e.g. an RVD) according to claim 9. The display device is typically a portable display device (e.g. a personal digital assistant, or PDA), or a remote computer, such as a COW or central nursing station. Here, the act of contacting the display device with the transceiver generates a pulse in the ACC waveform, as described above. Processing done by the computer analyzes both the pulse and a location of the display device to associate it with the transceiver.

[0020] Several methods can be used to determine the location of the display device. For example, the wireless transmitter within the transceiver is configured to operate on a wireless network, and algorithms operating on the remote computer and can analyze signals between the transceiver and wireless access points within the network (e.g. RSSI signals indicating signal strength) to determine an approximate location of the transceiver and thus the display device which it contacts. In embodiments the algorithms can involve, e.g., triangulating at least three RSSI values, or simply estimating location by determining the nearest access point from a single RSSI value. Triangulation typically involves using a map grid that includes known locations of multiple wireless access points and display devices within a region of the hospital; the map grid is determined beforehand and typically stored, e.g., in a database. For example, the approximate location of the transceiver can be determined using triangulation. Then the nearest display device, lying with a known location within a pre-determined radius, is paired with the transceiver. Typically the pre-determined radius is between 1-5m.

[0021] In another aspect, the invention provides systems for measuring a patient according to claims 2 and 11. Furthermore disclosed is a body-worn monitor including first and second sensors attached to the patient, and a processing component that interfaces to both sensors and processes signals from them to calculate at least one vital sign value. A wireless transmitter receives the vital sign value and transmits it over a wireless interface, and additionally provides a two-way communications system configured to transmit and receive audio signals

over the same wireless interface. In embodiments, the two-way communications system includes a speaker and a microphone, both of which are integrated into the transceiver. Typically the wireless interface is a hospital-based wireless network using an 802.11 protocol (e.g. 802.11a/b/g/n). A VOIP system typically runs on the wireless network to supply two-way voice communications. Alternatively the wireless network is based on a cellular protocol, such as a GSM or CDMA protocol.

[0022] Typically the body-worn monitor features a wrist-worn transceiver that functions as a processing component, and includes a touchpanel display configured to render both patient and clinician interfaces. The touchpanel display is typically a liquid crystal display (LCD) or organic light-emitting diode display (OLED) display with a clear touchpanel utilizing established resistive or capacitive technologies adhered to its front surface. The patient interface is typically rendered by default, and includes a graphical icon that, when initiated, activates the two-way communications system. The clinician interface typically requires a security code (entered using either a 'soft' numerical keypad or through a barcode scanner) to be activated. The transceiver typically includes a strap configured to wrap around the patient's arm, and most typically the wrist; this allows it to be worn like a conventional wristwatch, which is ideal for two-way communications between the patient and a clinician.

[0023] In a related aspect, herein disclosed is a wrist-worn transceiver wherein the two-way communications system described above, or a version thereof, is used as a voice annotation system. Such a system receives audio signals (typically from a clinician), digitizes them, and transmits the resulting digital audio signals, or a set of parameters determined from these signals, over the wireless interface to a computer memory. The audio signals are typically used to annotate vital sign information. They can be used, for example, to indicate when a pharmaceutical compound is administered to the patient, or when the patient undergoes a specific therapy. Typically the voice annotation uses the same speaker used for the two-way communication system. It also may include a speech-to-text converter that converts audio annotations from the clinician into text fields that can be easily stored alongside the vital sign information. In embodiments, both a text field and the original audio annotation are stored in a computer memory (e.g. database), and can be edited once stored. In other embodiments, a pre-determined text field (indicating, e.g., that a specific medication is delivered at a time/date automatically determined by the transceiver) is used to annotate the vital sign information. In still other embodiments, a set of parameters determined from the digital audio signals can include an icon or a numerical value. Annotations in the database can be viewed afterwards using a GUI that renders both the vital sign information (shown, e.g., in a graphical form) and one or more of the annotations (e.g. icon, text field, numerical value, or voice annotation).

[0024] In another aspect, herein disclosed is a wrist-

worn transceiver featuring a GUI that the patient can use to indicate their level of pain. Here, the GUI typically includes a touchpanel display configured to render a set of input fields, with each input field in the set indicating a different level of pain. Once contacted, the input fields generate a signal that is processed to determine the patient's level of pain. This signal can be further processed and then wirelessly transmitted to a remote computer for follow-on analysis.

[0025] It is furthermore disclosed that the touchpanel display features a touch-sensitive area associated with each input field that generates a digital signal (e.g. a number) after being contacted. Each input field is typically a unique graphical icon such as a cartoon or numerical value indicating an escalating level of pain. The transceiver can also include a voice annotation system similar to that described above so the patient can specifically describe their pain (e.g. its location) using their own voice. This information can be wirelessly transmitted to a remote computer (e.g. a PDS) featuring a display device (e.g. an RVD). This system can render both vital sign information and a parameter determined from the pain signal, and can additionally include an alarming system that activates an alarm if the pain signal or a parameter calculated therefrom exceeds a pre-determined threshold.

[0026] In a related aspect, herein disclosed is a wrist-worn transceiver that includes a mentation sensor configured to collect data input characterizing the patient's level of mentation (e.g. mental acuity). This information, along with traditional vital signs and the waveforms they are calculated from, is wirelessly transmitted to a remote computer for analysis. In embodiments, the mentation sensor is a touchpanel display that renders a GUI to collect information characterizing the patient's level of mentation. For example, the GUI can render a series of icons, a game, test, or any other graphical or numerical construct that can be used to evaluate mentation. In a specific embodiment, for example, the GUI includes a set of input fields associated with a numerical value. Here, the mentation 'test' features an algorithm to determine if the input fields are contacted by the patient in a pre-determined numerical order. Upon completion, the test results can be evaluated to generate a mentation 'score'. In this aspect, the wrist-worn transceiver also includes a two-way communication system that receives audio information from the patient. This audio information can be used for conventional communication purposes, and can additionally be analyzed to further gauge mentation. As in previous embodiments, the mentation score can be sent with vital sign information to a PDS/RVD for follow-on analysis. These systems may include an alarming system that generates an alarm if the mentation parameter or a parameter calculated therefrom exceeds a pre-determined threshold.

[0027] In another aspect, herein disclosed is a wrist-worn transceiver featuring a motion sensor (e.g. an accelerometer, mercury switch, or tilt switch) that generates

a motion signal indicating the transceiver's orientation. The processing component within the transceiver processes the motion signal and, in response, orients the GUI so that it can be easily viewed in 'rightside up' configuration, i.e. with text rendered in a conventional manner from left to right. If the transceiver is moved (e.g., so that it is viewed by a clinician instead of a patient), the accelerometers generate new motion signals, and the GUI is 'flipped' accordingly. Typically, for example, the GUI is rendered in either a first orientation or a second orientation, with the two orientations separated by 180 degrees., and in some cases by 90 degrees. In embodiments, the first orientation corresponds to a 'patient GUI', and the second orientation corresponds to a 'clinician GUI'. This allows, for example, the appropriate GUI to be automatically rendered depending on the transceiver's orientation. The clinician GUI typically includes medical parameters, such as vital signs and waveforms, whereas the patient GUI typically includes non-medical features, such as a 'nurse call button', time/date, and other components described in more detail below.

[0028] It is furthermore disclosed that the motion sensor is a 3-axis accelerometer configured to generate a time-domain ACC waveform. During a measurement, the processing component additionally analyzes the waveform to determine parameters such as the patient's motion, posture, arm height, and degree of motion.

[0029] It is disclosed that the wrist-worn transceiver features a display device configured to render at least two GUIs, with the first GUI featuring medical content, and the second GUI featuring non-medical content relating to entertainment, food service, games, and photographs. The photograph, for example, can include an image of the patient or a relative of the patient; this latter case may be particularly useful in neo-natal hospital wards. To capture the photograph, the body-worn monitor may include a digital camera, or a wireless interface to a remote digital camera, such as that included in a portable computer or cellular telephone.

[0030] It is disclosed that the second GUI is configured to render menus describing entertainment content, such as television (e.g. different channels or pre-recorded content), movies, music, books, and video games. In this case, the touchpanel display can be used to select the content or, in embodiments, play a specific game. The wireless transmitter within the transceiver is further configured to transmit and receive information from a remote server configured to store digital representations of these media sources. In still other embodiments, the second GUI is configured to display content relating to a food-service menu. Here, the wireless transmitter is further configured to transmit and receive information from a remote server configured to interface with a food-service system.

[0031] In another aspect, the invention provides a system for monitoring a patient that includes a vital sign monitor configured to be worn on the patient's body, and a remote computer according to claim 2 and 11. The vital

sign monitor features connection means (e.g. a flexible strap or belt) configured to attach a transceiver to the patient's body, and sensor with a sensing portion (e.g. electrodes and an optical sensor) that attaches to the patient to measure vital sign information. A mechanical housing included in the transceiver covers a wireless decoder, processing component, and wireless transmitter, and supports a display component. The wireless decoder (e.g. a barcode scanner or radio frequency identification (RFID) sensor) is configured to detect information describing a medication, a medication-delivery rate, a clinician, and the patient. For example, this information may be encoded in a barcode or RFID tag located on the patient, clinician, medication, or associated with an infusion pump. The processing component is configured to process: 1) the vital sign information to generate a vital sign and a time-dependent waveform; and 2) information received by the wireless decoder to generate decoded information. The wireless transmitter within the mechanical housing receives information from the processing component, and transmits it to a remote computer. In response the remote computer processes the information and transmits an information-containing packet back to the vital sign monitor.

[0032] It is disclosed that the remote computer performs an analyzing step that compares information describing both the medication and the patient to database information within a database. The database may include, for example, a list of acceptable medications and acceptable medication-delivery rates corresponding to the patient. In some cases both the vital sign information and the decoded information are collectively analyzed and compared to values in the database to affect treatment of the patient. For example, this analysis may determine that a patient with a low blood pressure should not receive medications that further lower their blood pressure. Or it may suggest changing a dosage level of the medication in order to compensate for a high heart rate value. In general, the remote computer can analyze one or more vital sign values corresponding to a patient, along with the patient's demographic information, medical history, and medications, and determine acceptable medications and medication-delivery rates based on this analysis. In response, the computer can transmit a packet back to the vital sign monitor, which renders its contents on the display. The packet can include a message confirming that a particular medication and medication-delivery rate are acceptable for the patient, and may also include a set of instructions for delivering the medication and performing other therapies.

[0033] Still other embodiments are found in the claims.

BRIEF DESCRIPTION OF THE DRAWINGS

[0034]

Fig. 1 is a schematic drawing showing the wrist-worn transceiver of the invention attached to a patient's

wrist;

Figs. 2A and 2B show, respectively, schematic drawings of the wrist-worn transceiver of Fig. 1 oriented 'rightside up' so that a patient can view the GUI, and 'upside down' so that a clinician can view the GUI; Fig. 3 shows a schematic drawing of the wrist-worn transceiver of Fig. 1 and a list of features available in both a patient GUI and a clinician GUI;

Fig. 4 shows a schematic drawing of the body-worn monitor featuring sensors for measuring ECG, PPG, ACC, OSC, and IP waveforms, and systems for processing these to determine a patient's vital signs; Fig. 5 shows a schematic drawing of an IT configuration of the invention where the body-worn monitor of Fig. 4 is connected through a wireless network to a PDS and hospital EMR;

Fig. 6A and 6B show, respectively, schematic drawings of a new transceiver having a fully charged battery being swapped with an original transceiver having a depleted battery before and after deploying the 'bump' methodology;

Fig. 7 shows a schematic drawing of transceivers undergoing the 'bump' methodology of Figs. 6A and 6B and wirelessly transmitting their ACC waveforms to the PDS for analysis;

Fig. 8 shows screen captures from a GUI used to guide a clinician through the 'bump' methodology of Figs. 6A, 6B, and 7;

Fig. 9 shows a schematic drawing of a transceiver being 'bumped' against a RVD in order to pair the two devices;

Fig. 10 shows a map indicating how the transceiver and RVD of Fig. 9 are paired to each other;

Fig. 11 shows a schematic drawing of the wrist-worn transceiver of Fig. 1 being used for voice annotation of a patient's vital sign data;

Fig. 12 shows a schematic drawing of the wrist-worn transceiver of Fig. 11 wirelessly transmitting voice annotations to the PDS for analysis;

Fig. 13 shows screen captures from a GUI used to guide a clinician through the voice annotation methodology of Figs. 11 and 12;

Fig. 14 shows screen captures from a GUI used when the wrist-worn transceiver functions as a two-way communicator between the patient and a clinician;

Fig. 15 shows a screen capture from a GUI used to render a 'pain index' on the wrist-worn transceiver;

Fig. 16 shows a screen capture from a GUI used to render a mentation test on the wrist-worn transceiver;

Fig. 17 shows a screen capture from a GUI used to render a photograph of the patient on the wrist-worn transceiver;

Fig. 18 shows a screen capture from a GUI used to render a food menu on the wrist-worn transceiver;

Fig. 19 shows a screen capture from a GUI used to render a menu of television channels on the wrist-

worn transceiver;

Fig. 20 shows a schematic drawing of the barcode scanner in the wrist-worn transceiver scanning barcodes associated with a patient, clinician, and medication, and sending the decoded barcode information to the PDS;

Figs. 21A and 21B show three-dimensional images of the body-worn monitor of Fig. 4 attached to a patient with and without, respectively, a cuff-based pneumatic system used for a calibrating indexing measurement;

Figs. 22A and 22B show, respectively, three-dimensional images of the wrist-worn transceiver before and after receiving cables from other sensors within the body-worn monitor;

Fig. 23A shows a schematic drawing of a patient wearing the body-worn monitor of Fig. 21B and its associated sensors;

Fig. 23B shows graphs of time-dependent ECG, PPG, OSC, ACC, and IP waveforms generated with the body-worn monitor and sensors of Fig. 23A;

Fig. 24 shows screen captures from a GUI used to render vital signs and ECG, PPG, and IP waveforms on the wrist-worn transceiver;

Fig. 25 shows a schematic drawing of the ACC, ECG, pneumatic, and auxiliary systems of the body-worn monitor communicating over the CAN protocol with the wrist-worn transceiver;

Fig. 26 shows an alternate IT configuration of the invention where the wrist-worn transceiver of Fig. 1 communicates with the PDS through a wireless access point connected to the Internet;

Fig. 27 shows an alternate IT configuration of the invention where the wrist-worn transceiver of Fig. 1 communicates with the PDS through a wireless device connected to the Internet; and

Fig. 28 shows an alternate IT configuration of the invention where the wrist-worn transceiver of Fig. 1 communicates with the PDS through an internal cellular modem connected to the Internet.

DETAILED DESCRIPTION OF THE INVENTION

System Overview

[0035] Fig. 1 shows a transceiver 72 according to the invention that attaches to a patient's wrist 66 using a flexible strap 90. The transceiver 72 connects through a first flexible cable 92 to a thumb-worn optical sensor 94, and through a second flexible cable 82 to an ECG circuit and a series of chest-worn electrodes (not shown in the figure). During a measurement, the optical sensor 94 and chest-worn electrodes measure, respectively, time-dependent optical waveforms (e.g. PPG) and electrical waveforms (e.g. ECG and IP), which are processed as described below to determine vital signs and other physiological parameters such as cNIBP, SpO₂, HR, RR, TEMP, pulse rate (PR), and cardiac output (CO). Once

measured, the transceiver 72 wirelessly transmits these and other information to a remote PDS and RVD. The transceiver 72 includes a touchpanel display that renders a GUI 50 which, in turn, displays the vital signs, physiological parameters, and a variety of other features described in detail below. Collectively, the transceiver 72 and GUI 50 incorporate many features that are normally reserved for non-medical applications into a body-worn vital sign monitor that continuously monitors ambulatory patients as they move throughout the hospital.

[0036] The transceiver 72 includes an embedded accelerometer that senses its motion and position, and in response can affect properties of the GUI. Referring to Figs. 2A and 2B, for example, time-resolved ACC waveforms from the accelerometer can be processed with a microprocessor within the transceiver to detect orientation of the touchpanel display. This information can then be analyzed to determine if it is the clinician or patient who is viewing the display. In response, the GUI can 'flip' so that it is properly oriented (i.e. 'rightside up', as opposed to being upside down) for the viewer. For example, as shown in Fig. 2A, when the transceiver 72 is worn on the patient's right wrist 66 the internal accelerometer generates ACC waveforms that are processed by the microprocessor to determine this orientation. The GUI 50A is adjusted according so that it is always oriented with numbers and text arranged rightside up and read from left to right. When the patient's arm is rotated, as shown in Fig. 2B, the ACC waveforms change accordingly because the accelerometer's axes are swapped with respect to gravity. Such a situation would occur, for example, if a clinician were to orient the patient's arm in order read the transceiver's display. In this case, the ACC waveforms are processed to determine the new orientation, and the GUI 50B is flipped so it is again rightside up, and can be easily read by the clinician.

[0037] The internal accelerometer can also detect if the transceiver is 'bumped' by an external object. In this case, the ACC waveform will feature a sharp 'spike' generated by rapid acceleration and deceleration caused by the bumping process. As described in detail below, such a bumping process can serve as a fiducial marker that initiates a specific event related to the transceiver, such as a battery swap or process that involves pairing the transceiver to an external wireless system or display.

[0038] The accelerometer within the transceiver, when combined with other accelerometers within the body-worn monitor, can also be used to determine the patient's posture, activity level, arm height and degree of motion, as described in detail below. Use of one or more accelerometers to detect such motion-related activities is described, for example, in the following patent applications: US 2010/0298656 A1 and US 2010/0298650 A1.

[0039] Referring to Fig. 3, in addition to the GUI 50, the transceiver 72 includes a high-fidelity speaker 120, a microphone 101, and a barcode scanner 102 which, respectively, enunciates audible information, measures voice signals from both the patient and a clinician, and

scans graphical barcodes to decode numerical information describing the patient and their medication. Signals from these and other components are processed to supply information to either a 'patient GUI' 52 or a 'clinician GUI' 54. The patient GUI 52, for example, typically includes features that are decoupled from a standard clinical diagnosis; these include a nurse call button, voice communications, a 'pain' index, a mentation test to estimate the patient's cognitive abilities, meal ordering within the hospital, games, and a controller for entertainment content, e.g. to adjust parameters (e.g. channels, volume) for a standard television set. The clinical GUI 54, in comparison, includes features that are used for clinical diagnoses and for operating the transceiver in a hospital environment. The primary features of this GUI 54 include displaying vital signs (e.g. cNIBP, SpO2, HR, RR, TEMP), other medical parameters (e.g. PR, CO), and waveforms (PPG, ECG, IP). Secondary features of the clinical GUI include voice communications, battery-change and pairing operations using the above-described 'bump' methodology, voice annotation of medical records and diagnoses, a method for checking medications using the barcode scanner 102, and display of a photograph or video describing the patient.

[0040] During normal operation, the GUI renders 50 simple icons indicating that the transceiver is powered on and operational (e.g., a 'beating heart'), the strength of the wireless signal (e.g. a series of bars with escalating height), and the battery level (e.g. a cartoon of a battery with a charge-dependent gauge). The transceiver 72 displays these icons until the touchpanel display is contacted by either the patient or a clinician. This process yields the patient GUI 52, which features a large icon 57 showing a telephone (which is used for nurse call applications, as described below), and a smaller icon 53 showing a lock which, when tapped, enables the clinician to 'unlock' the transceiver and utilize the clinician interface 54. The transceiver 72 immediately renders a GUI that shows vital signs and waveform information if the patient's physiological condition requires immediate medical attention, e.g. in the case of cardiac arrest.

[0041] The clinician interface 54 is password-protected to prevent the patient or any other non-clinician from viewing important and potentially confusing medical information. A password can either be entered as a standard personal identification number (PIN) by tapping keys on a numerical keypad (as shown in Fig. 3), or by simply swiping a barcode printed on the clinician's hospital badge across the barcode scanner 102. The microprocessor within the transceiver unlocks the clinician interface following either of these events, and enables all the features associated with the interface, which are described in detail below. For example, with this interface the clinician can view vital signs and waveforms to make a medical diagnosis, as described with reference to Fig. 24. If the transceiver's battery charge is running low, the clinician can swap in a new transceiver and transfer data from the original transceiver simply by 'bumping' the two

transceivers together, as described with reference to Figs. 6-8. Medical records can be voice-annotated and stored on the PDS or a hospital's EMR using the process shown in Figs. 11-13. The patient's medication can be checked by scanning and processing information encoded in barcodes associated with the patient, clinician, and medication, as shown in Fig. 20. All of this functionality is programmed within the transceiver and the body-worn monitor, and can be accomplished without tethering the patient to a conventional vital sign monitor typically mounted on a wall in the hospital or a rolling stand. Ultimately this allows the patient to wear a single body-worn monitor as they transition throughout the various facilities within the hospital, e.g. the ED, ICU, x-ray facility, and operating room.

Hardware in Body-Worn Monitor

[0042] Figs. 4 and 5 show schematic drawings of a body-worn monitor 100 used to measure vital signs from a patient and render the different GUIs described above (Fig. 4), along with a wireless system over which the transceiver 72 sends information through a hospital network 60 to either a remote RVD, e.g. a computer 62 or hand-held device 64 (Fig. 5). Referring to Fig. 4, the body-worn monitor 100 features a wrist-worn transceiver 72 that continuously determines vital signs and motion-related properties from an ambulatory patient in a hospital. The monitor 100 is small, lightweight, and comfortably worn on the patient's body during their stay in the hospital; its specific form factor is described in detail below with reference to Figs. 21 and 22. It provides continuous monitoring, and features a software framework that determines alarms/alerts if the patient begins to decompensate. Such systems are described in the following co-pending patent applications: US 2010/0298656 A1 and US 2010/0298650 A1. The framework processes both the patient's motion and their vital sign information with algorithms that reduce the occurrence of false alarms.

[0043] A combination of features makes the body-worn monitor 100 ideal for ambulatory patients within the hospital. For example, as shown in Fig. 5, the transceiver 72 features a wireless transmitter 224 that communicates through a collection of wireless access points 56 (e.g. routers based on 802.11 protocols) within a hospital network 60, which includes a PDS. From the PDS 60 data are sent to an RVD (e.g. a portable tablet computer 62) located at a central nursing station, or to a local computer (e.g. a hand-held PDA 64) carried by the clinician. In embodiments, data can be sent to the PDA 64 through a peer-to-peer wireless connection. The specific mode of communication can be determined automatically (using, e.g., a signal strength associated with the wireless connection), or manually through an icon on the GUI.

[0044] The transceiver 72 features a CPU 222 that communicates through a digital CAN interface, or bus, to external systems featuring ECG 216, external accelerometers 215b-c, pneumatic 220, and auxiliary 245 sen-

sors. Each sensor 215b-c, 216, 220, 245 is 'distributed' on the patient to minimize the bulk and weight normally associated with conventional vital sign monitors, which typically incorporate all electronics associated with measuring vital signs in a single plastic box. Moreover, each of these sensors 215b-c, 216, 220, 245 generate digital signals close to where they actually attach to the patient, as opposed to generating an analog signal and sending it through a relatively long cable to a central unit for processing. This can reduce noise due to cable motion which is often mapped onto analog signals. Cables 240, 238, 246 used in the body-worn monitor 210 to transmit packets over the CAN bus typically include five separate wires bundled together with a single protective cladding: the wires supply power and ground to the remote ECG system 216, accelerometers 215b-c, pneumatic 220, and auxiliary systems 245; provide high/low signal transmission lines for data transmitted over the CAN protocol; and provide a grounded electrical shield for each of these four wires. There are several advantages to this approach. First, a single pair of transmission lines in the cable (i.e. the high/low signal transmission lines) can transmit multiple digital waveforms generated by completely different sensors. This includes multiple ECG waveforms (corresponding, e.g., to vectors associated with three, five, and twelve-lead ECG systems) from the ECG circuit, along with ACC waveforms associated with the x, y, and z axes of accelerometers within the body-worn monitor 100. The same two wires, for example, can transmit up to twelve ECG waveforms (measured by a twelve-lead ECG system), and six ACC waveforms (measured by the accelerometers 215b-c). Limiting the transmission line to a pair of conductors reduces the number of wires attached to the patient, thereby decreasing the weight and any cable-related clutter. Second, cable motion induced by an ambulatory patient can change the electrical properties (e.g. electrical impedance) of its internal wires. This, in turn, can add noise to an analog signal and ultimately the vital sign calculated from it. A digital signal, in contrast, is relatively immune to such motion-induced artifacts.

[0045] The ECG 216, pneumatic 220, and auxiliary 245 systems are stand-alone systems that each includes a separate CPU, analog-to-digital converter, and CAN transceiver. During a measurement, they connect to the transceiver 72 through cables 240, 238, 246 and connectors 230, 228, 232 to supply digital inputs over the CAN bus. The ECG system 216, for example, is completely embedded in a terminal portion of its associated cable. Systems for three, five, and twelve-lead ECG monitoring can be swapped in an out simply by plugging the appropriate cable (which includes the ECG system 216) into a CAN connector 230 on the wrist-worn transceiver 72, and the attaching associated electrodes to the patient's body.

[0046] As described above, the transceiver 72 renders separate GUIs that can be selected for either the patient or a clinician. To do this, it includes a barcode scanner

242 that can scan a barcode printed, e.g., on the clinician's badge. In response it renders a GUI featuring information (e.g. vital signs, waveforms) tailored for a clinician that may not be suitable to the patient. So that the patient can communicate with the clinician, the transceiver 72 includes a speaker 241 and microphone 237 interfaced to the CPU 222 and wireless system 224. These components allow the patient to communicate with a remote clinician using a standard VOIP protocol. A rechargeable Li-ion battery 239 powers the transceiver 72 for about four days on a single charge. When the battery charge runs low, the entire transceiver 72 is replaced using the 'bump' technique described in detail below.

[0047] Three separate digital accelerometers 215a-c are non-obtrusively integrated into the monitor's form factor; two of them 215b-c are located on the patient's body, separate from the wrist-worn transceiver 72, and send digitized, motion-related information through the CAN bus to the CPU 222. The first accelerometer 215a is mounted on a circuit board within the transceiver 72, and monitors motion of the patient's wrist. The second accelerometer 215b is incorporated directly into the cable 240 connecting the ECG system 216 to the transceiver 72 so that it can easily attach to the patient's bicep and measure motion and position of the patient's upper arm. As described below, this can be used to orient the screen for viewing by either the patient or clinician. Additionally, signals from the accelerometers can be processed to compensate for hydrostatic forces associated with changes in the patient's arm height that affect the monitor's cNIBP measurement, and can be additionally used to calibrate the monitor's blood pressure measurement through the patient's 'natural' motion. The third accelerometer 215c is typically mounted to a circuit board that supports the ECG system 216 on the terminal end of the cable, and typically attaches to the patient's chest. Motion and position of the patient's chest can be used to determine their posture and activity states, which as described below can be used with vital signs for generating alarm/alerts. Each accelerometer 215a-c measures three unique ACC waveforms, each corresponding to a separate axis (x, y, or z) representing a different component of the patient's motion. To determine posture, arm height, activity level, and degree of motion, the transceiver's CPU 222 processes signals from each accelerometer 215a-c with a series of algorithms, described in the following pending patent applications US 2010/0298656 A1 and US2010/0298650 A1. In total, the CPU 222 can process nine unique, time-dependent signals corresponding to the three axes measured by the three separate accelerometers. Algorithms determine parameters such as the patient's posture (e.g., sitting, standing, walking, resting, convulsing, falling), the degree of motion, the specific orientation of the patient's arm and how this affects vital signs (particularly cNIBP), and whether or not time-dependent signals measured by the ECG 216, optical 218, or pneumatic 220 systems are corrupted by motion.

[0048] To determine blood pressure, the transceiver 72 processes ECG and PPG waveforms using a measurement called with Composite Technique, which is described in the following patent application, US 2010/0168589 A1. The Composite Technique measures ECG and PPG waveforms with, respectively, the ECG 216 and optical 218 systems. The optical system 218 features a thumb-worn sensor that includes LEDs operating in the red ($\lambda \sim 660\text{nm}$) and infrared ($\lambda \sim 900\text{nm}$) spectral regions, and a photodetector that detects their radiation after it passes through arteries within the patient's thumb. The ECG waveform, as described above, is digitized and sent over the CAN interface to the wrist-worn transceiver 72, while the PPG waveform is transmitted in an analog form and digitized by an analog-to-digital converter within the transceiver's circuit board. The pneumatic system 220 provides a digitized pressure waveform and oscillometric blood pressure measurements through the CAN interface; these are processed by the CPU 222 to make cuff-based 'indexing' blood pressure measurements according to the Composite Technique. The indexing measurement typically only takes about 40-60 seconds, after which the pneumatic system 220 is unplugged from its connector 228 so that the patient can move within the hospital without wearing an uncomfortable cuff-based system. The optical waveforms measured with the red and infrared wavelengths can additionally be processed to determine SpO2 values, as described in detail in the following patent application US 2010/0324385 A1.

[0049] Collectively, these systems 215a-c, 216, 218, and 220 continuously measure the patient's vital signs and motion, and supply information to the software framework that calculates alarms/alerts. A third connector 232 also supports the CAN bus and is used for auxiliary medical devices 245 (e.g. a glucometer, infusion pump, system for measuring end-tidal CO2) that is either worn by the patient or present in their hospital room.

[0050] Once a measurement is complete, the transceiver 72 uses the internal wireless transmitter 224 to send information in a series of packets to a PDS 60 within the hospital. The wireless transmitter 224 typically operates on a protocol based on 802.11, and can communicate with the PDS 60 through an existing network within the hospital as described above with reference to Fig. 5. Information transmitted by the transceiver alerts the clinician if the patient begins to decompensate. The PDS 60 typically generates this alarm/alert once it receives the patient's vital signs, motion parameters, ECG, PPG, and ACC waveforms, and information describing their posture, and compares these parameters to preprogrammed threshold values. As described in detail below, this information, particularly vital signs and motion parameters, is closely coupled together. Alarm conditions corresponding to mobile and stationary patients are typically different, as motion can corrupt the accuracy of vital signs (e.g., by adding noise), and induce artificial changes in them (e.g., through acceleration of the patient's

heart and respiratory rates) that may not be representative of the patient's actual physiology.

Swapping and Pairing Transceivers Using 'Bump' Methodology

[0051] Figs. 6A, 6B, 7, and 8 show how a wrist-worn transceiver 72A with a depleted battery can be swapped with a similar transceiver 72B having a fully charged battery using the 'bump' methodology described above. Prior to the swap, as shown in Fig. 6A, both transceivers are readied by activating the appropriate GUI 50C, 50D following the screens shown in Fig. 8. This process activates firmware on each transceiver 72A, 72B indicating that the swap is about to occur. In response, each transceiver sends a packet through the wireless access point 56 and to the hospital network and PDS 60. The packet describes a transceiver-specific address, e.g. a MAC address associated with its wireless transmitter. Once this is done, the GUIs 50C, 50D on both transceivers 72A, 72B indicate to a clinician that they can be 'bumped' together, and that the swap can proceed.

[0052] At this point, as shown in Fig. 6A, the new transceiver 72B (with the fully charged battery) is then bumped against the old transceiver 72A (with the depleted battery). Internal accelerometers within both transceivers 72A, B detect the bumping process and, in response, independently generate ACC waveforms 130, 132, both featuring a sharp spike indicating the rapid acceleration and deceleration due to the bumping process. Typically the ACC waveforms 130, 132 correspond to the same axes in both transceivers. The ACC waveforms are digitized within each transceiver and then transmitted through the wireless access point 56 to the PDS 60, where they are stored in a computer memory and analyzed with a software program that is activated when both devices are 'readied', as described above. The software program compares formatted versions of the ACC waveforms 130', 132' to detect the rapid spikes, as shown by the graph 140 in Fig. 7. The rapid spikes in the waveforms 130', 132' should occur within a few microseconds of each other, as indicated by the shaded window 142 in the graph 140. Other transceivers operating on the network may generate similar motion-related spikes due to movements of the patient wearing them, but the probability that such spikes occur at the exact same time as the transceivers being swapped is extremely low. The software program interprets the concurrence of the spikes as indicating that data stored on the old transceiver 72A is to be transferred to the new transceiver 72B. The data, for example, includes demographic information describing the patient (e.g. their name, age, height, weight, photograph), the medications they are taking, and all the vital sign and waveform information stored in memory in the old transceiver 72A. Following the bump, this information is associated with the address corresponding to the new transceiver 72B. At this time information may also be sent from the PDS so it can be stored

locally on the new transceiver. When all the relevant information is transferred over, the GUIs 50C, 50D on both transceivers 72A, 72B indicate that they can be swapped. At this point, cables connected to the optical sensor and ECG electrodes are unplugged from the old transceiver 72A, and plugged into the new transceiver 72B. The clinician then attaches the new transceiver to the patient's wrist, and commences measuring vital signs from the patient as described above.

[0053] In other embodiments, a time period corresponding to a portion (e.g. a peak value) of the motion-generated spike is determined on each of the wrist-worn transceivers that are bumped together. Each transceiver then sends its time period to the PDS, where they are collectively analyzed to determine if they are sufficiently close in value (e.g. within a few hundred milliseconds). If this criterion is met, software on the PDS assumes that the transceivers are ready to be swapped, and performs the above-described steps to complete this process.

[0054] Fig. 8 shows a sequence of screens within the GUI that describe the process for swapping transceivers to the clinician. The process begins when a screen 158 rendered by Device A indicates that its battery is running low of charge. This is indicated by a standard 'low battery' icon located in the upper right-hand corner of the screen 158, as well as a larger icon located near the bottom of the screen. A time describing the remaining life of the battery appears near this icon when this time is 5 minutes or less. Each transceiver includes a sealed internal Li:ion battery that cannot be easily replaced in the hospital. Instead, the transceiver is inserted in a battery charger that typically includes eight or sixteen ports, each of which charges a separate transceiver. To swap Device A with Device B, the clinician taps the screen 158 to yield a new screen 160 which includes a series of six icons, each related to a unique feature. The icon in the lower left-hand corner shows two interchanging batteries. When tapped, this icon yields a new screen 160 indicating that Device A is ready to be swapped. Device B is then removed from a port in the battery charger, and a sequence of screens 150, 152, 154 are initiated as described with reference to Device A.

[0055] When Devices A and B both show, respectively, screens 162, 154, they are ready to be swapped using the 'bumping' process. At this point, as described above, a clinician 'bumps' Device B into Device A, which in turn generates two ACC waveforms 130, 132 featuring sharp, time-dependent spikes indicating the bump. The waveforms 130, 132 include spikes, as shown by the shaded box 142, which are concurrent in time, and are wirelessly transmitted in a packet that indicates their origin through the pathway shown in Fig. 7 to the PDS. There, they are analyzed by the software program described above to determine that data associated with Device A (e.g. patient information, vital signs) is now associated with Device B. When this association is complete, the PDS transmits a packet back through the pathway shown in Fig. 7 to both Device A and B, indicating that the PDS is ready

to transfer the data. Device B then renders a screen 166 asking the clinician to confirm the process. Data is transferred if the clinician taps the 'check' box in the lower right-hand corner of the screen; during this process Device B renders a screen 168 that shows the patient's name to further confirm with the clinician that the transfer process is valid. When it is complete, Device A is no longer active, meaning it cannot collect data or generate alarms. Device B renders a screen 170 that instructs the clinician to disconnect the optical and electrical sensors from Device A, and to clean this device and insert it into the battery charger. During this process all alarms are paused for Device B. A screen 172 on Device B then instructs the clinician to connect the sensors and attach Device B to the patient's wrist. When this is complete, Device B renders a final confirmatory screen 176, which when checked finalizes the swapping process. At this point Device B is officially associated with the patient, renders a standard screen 178, and commences measuring vital signs from the patient. These vital signs, along with those collected from Device A, are included in a contiguous data file characterizing the patient.

[0056] As an alternative to the 'bumping' process, Device B's barcode can be read and processed to facilitate swapping the transceivers. In this case, an icon on Device A, when tapped, renders a screen 164 indicating that Device A is ready to read the barcode printed on Device B. At this point, Device B's barcode is swiped across Device A's barcode reader, decoded, and wirelessly transmitted to the PDS as indicated in Fig. 7. The PDS uses this information to associate Device B with the patient as described above. Once this is complete, Device B uses the same screens used for the 'bumping' transfer process (screens 166, 168, 170, 172, 176, 178) to associate Device B with the patient. The 'bumping' process shown in Fig. 6 takes place along the long axes of Device A and Device B. Alternatively, it can take place along the short axes of these devices. Or the short axis of one device can be bumped against the long axis of the other device to initiate the process.

[0057] The 'bumping' process described above can also be used for other applications relating to the wrist-worn transceiver. It can be used, for example, to pair the transceiver with an RVD, such as a display located at the patient's bedside, or at a central nursing station. In this embodiment, indicated in Figs. 9 and 10, a clinician selects a transceiver 72 from the battery charger and brings it near an RVD 62. Before attaching the transceiver 72 to the patient, the clinician 'bumps' it against a hard surface proximal to the RVD 62 (or against the RVD itself) to generate a sharp spike in the ACC waveform 133. The waveform 133 is similar in shape to that generated when two transceivers are swapped with the bumping process, as described above. The RVD's location needs to be determined in order to pair it with the transceiver 72. To do this, at a pre-determined time period (e.g. every few minutes) all neighboring wireless access points 56A, 56B, 56C transmit a 'location beacon' 59A, 59B, 59C to the

transceiver, which is received and used to calculate a value for signal strength (typically characterized by an 'RSSI value') between the transceiver 72 and the respective access point 56A, 56B, 56C. The transceiver concatenates values for RSSI and identifiers for the access points into a single 'location packet' 59D, which it then transmits along with the ACC waveform 133 and an identifying code describing the transceiver (not shown in the figure) through a single access point 56B to the PDS 60. The PDS 60 receives the location packet 56D and parses it to arrive at RSSI values for the three wireless access points 56A, 56B, 56C within wireless range of the transceiver 72. In other embodiments, the individual access points 56A, 56B, 56C determine RSSI values characterizing the signal strength between them and the transceiver, and send these as individual packets to the PDS. Software on the PDS then concatenates these packets to determine signals similar to those included in the location packet.

[0058] Referring to Fig. 10, location-determining software operating on the PDS triangulates the signals, along with known locations of each wireless access point 56A, 56B, 56C, to determine an approximate location 71 of the transceiver 72. The known locations of the access points are stored within a map grid 73 in a computer memory associated with the location-determining software. The transceiver's approximate location typically has an accuracy of 1-3m. Using the map grid 73, the software then processes the approximate location 71 and a known location of any RVD 62 lying within a pre-determined radius 75. Typically the pre-determined radius is 3-5m. If the location of the RVD 62 lies within the pre-determined radius 75, the RVD 62 is automatically 'paired' with the transceiver 72. Once paired, the RVD 62 then displays any follow-on waveform, motion, and vital sign information sent by the transceiver.

[0059] In related embodiments, the location-determining software described above uses triangulation algorithms to determine the patient's current and historical location. Such a process can be used to monitor and locate a patient in distress, and is described, for example, in the following issued patent: WIRELESS, INTERNET-BASED, MEDICAL DIAGNOSTIC SYSTEM (U.S. Patent 7,396,330). If triangulation is not possible, the location-determining software may simply use proximity to a wireless access point (as determined from the strength of an RSSI value) to estimate the patient's location. Such a situation would occur if signals from at least three wireless access points were not available. In this case, the location of the patient is estimated with an accuracy of about 5-10m. In embodiments, the RVD may be a central nursing station that displays vital sign, motion-related properties (e.g. posture and activity level) and location information from a group of patients. Such embodiments are described in the following co-pending patent application US 2011/0066050 A1. In other embodiments, the location-determining software determines the location of a patient-worn transceiver, and automatically pairs it to

a RVD located nearby (e.g. within a pre-determined radius, such as that shown in Fig. 10). In this way, the patient's information can be displayed on different RVDs as they roam throughout the hospital.

[0060] In embodiments, the patient's location can be analyzed relative to a set of pre-determined boundaries (e.g. a 'geofence') to determine if they have wandered into a restricted area. Or their speed can be determined from their time-dependent location, and then analyzed relative to a pre-determined parameter to determine if they are walking too fast. In general, any combination of location, motion-related properties, vital signs, and waveforms can be collectively analyzed with software operating on either the transceiver or PDS to monitor the patient. Patients can be monitored, for example, in a hospital, medical clinic, outpatient facility, or the patient's home.

[0061] In the embodiments described above, location of the transceiver can be determined using off-the-shelf software packages that operate on the PDS. Companies that provide such software include, for example, by Cisco Systems (170 West Tasman Drive, San Jose, CA 93134; www.cisco.com), Ekahau (12930 Saratoga Avenue, Suite B-8, Saratoga, CA 95070; www.ekahau.com), and others.

[0062] In still other embodiments, software operating on the transceiver puts it into a 'sleep mode' when it is not attached to the patient. This way the transceiver can determine and transmit a location packet even when it is not used for patient monitoring. Using the above-described location-determining software, this allows the transceiver's location to be determined and then analyzed if it has been lost, misplaced, or stolen. For example, the transceiver's serial number can be entered into the software and then used to send a 'ping' the transceiver. The transceiver responds to the ping by collecting and transmitting a location packet as described above. Or the location of all unused transceivers can be automatically rendered on a separate interface. In still other embodiments, the location-determining software can transmit a packet to a specific transceiver (e.g. one that is stolen) to disable it from operating further.

[0063] In other embodiments, the 'bumping' process described above can be used for a variety of applications involving the body-worn monitor, wrist-worn transceiver, PDS, and RVD. In embodiments, for example, one or more 'bumps' of a transceiver can modulate the ACC waveform, which is then processed and analyzed to initiate a specific application. Applications include turning the transceiver on/off; attaching sensors to the transceiver; pairing the transceiver with a hand-held device (e.g. a cellular phone or personal digital assistant) over a peer-to-peer connection (using, e.g., 802.11 or 802.15.4); pairing the transceiver with a printer connected to a hospital network to print data stored in its computer memory; associating the transceiver with a specific clinician; and initiating display of a particular GUI. In general, the 'bumping' process can be used to initiate any application that can also be initiated with icons on the GUI.

Annotating the Medical Record Using the Wrist-Worn Transceiver

[0064] Figs. 11-13 show how the wrist-worn transceiver can be used to communicate audible information from both the patient and a clinician. Audible information from the clinician 140 can be used, for example, to annotate vital sign information collected with the body-worn monitor. Audible information from the patient 141 can be transmitted to a clinician (e.g. a nurse working at a central station) to alert the clinician of a problem. In both applications, the transceiver 72 is attached to the patient's wrist 66 as described above and used to measure vital signs and waveform information. Audible information is received by a microphone 101 mounted on a circuit board within the transceiver. A speaker 120 mounted to the same circuit board enunciates voice information to the patient. In these and other voice-related applications, voice information is digitized by an internal analog-to-digital converter within the transceiver, and then wirelessly transmitted through a hospital's wireless network using conventional VOIP protocols. Systems that operate these protocols are marketed, for example, by Cisco Systems (170 West Tasman Drive, San Jose, CA 93134; www.cisco.com), Skype (22/24 Boulevard Royal, 6e etage, L-2449, Luxembourg; www.skype.com), and others.

[0065] Fig. 12 describes the annotation process in more detail. In this case, the transceiver 72 within the body-worn monitor is attached to the patient's wrist 66 to measure the patient's vital signs (e.g. blood pressure). During the measurement process, the clinician uses the GUI 50F to activate an 'annotation' function which enables the transceiver to receive audible signals 140 which are used, for example, to annotate different medications administered to the patient. After the annotation function is activated, the clinician orally describes the medications. The microphone 101 within the transceiver 72 detects the voice signals, digitizes them with associated hardware, and then sends them and an associated time/date sample using a VOIP protocol through an access point 56 and to a PDS located within the hospital network 60. Vital signs are transmitted before and after the annotation function is activated, and are stored along with the annotation in a computer memory associated with the PDS. Typically these data are stored within a hospital's EMR.

[0066] As shown in graph 141, annotated vital sign data can be viewed afterwards to determine, for example, how a patient responds to specific medications. In this case, administration of a beta blocker as a means of lowering the patient's blood pressure is recorded on the graph by a written description of the annotation, along with an icon (a black triangle) indicating when it occurred in time. To generate the written description the PDS requires software that performs a speech-to-text conversion. Such software is available, for example, from Nuance Systems (1 Wayside Road, Burlington, MA 01803;

www.nuance.com). Similarly, the graph 141 shows a second annotation indicating that the patient was hydrated with saline to increase their blood pressure.

[0067] Fig. 13 shows a series of screens within the GUI 50F that are used to control the annotation process. As described above, to annotate medical information the clinician taps an icon located in the upper right-hand portion of screen 180. This action readies the voice recording features within the transceiver. Tapping the annotation icon drives the transceiver to render a second screen 182 that includes the type of annotation, e.g. audible content relating to medication, a specific intervention or procedure, a medical assessment, or another subject. Typically annotations are delivered as audible speech, in which case the 'Speech' button is tapped, as shown in screen 184. Alternatively the annotation can be text or numerical; these can be typed in, e.g., using a 'soft' keyboard on the transceiver, or scanned in using the transceiver's barcode scanner. The annotations can also be associated with an alarm condition, such as those shown on screens 181, 183, 185, 187, 189, 191. Prior to recording an annotation, the GUI renders a screen 188 that, once tapped, initiates the recording. The recording can also be paused using screen 186. After it is complete, the clinician taps the 'checkbox' on the screen 188, thus saving the recording. It is then sent to the PDS as shown in Fig. 12, and used to annotate the patient's medical information.

[0068] Other forms of annotation are also possible with the transceiver. For example, it can include a small CCD camera that allows images of the patient or their body (e.g. a wound) to be captured and used to annotate the medical information. In other applications, a barcode printed on medication administered to the patient can be scanned by the transceiver's barcode scanner, and the information encoded therein can be used to annotate vital sign information. In other embodiments, the transceiver can integrate with other equipment in the hospital room (e.g. an infusion pump, ventilator, or patient-controlled anesthesia pump) through a wired or wireless connection, and information from this equipment can be collected and transmitted to the PDS in order to annotate the vital sign information. In other embodiments, text annotations can be stored on the PDS, and then edited afterwards by the clinician.

Other GUI Applications

[0069] As shown in Figs. 11 and 14, the speaker 120 and microphone 101 within the transceiver 72, combined with VOIP software operating on the hospital network, can also function as a nurse call system that communicates both distress signals and voice information. Here, the transceiver enables two-way communication between the patient and a remote clinician. During this application, the transceiver typically operates the 'patient GUI', shown schematically in Fig. 3 and in more detail in Fig. 14. Here, the GUI shows a single screen 192 that

indicates a nurse call function with an icon showing a telephone. When the patient taps on the telephone the transceiver initiates a call to a pre-programmed IP address, corresponding, e.g., to a computer at a central nursing station or a VOIP-enabled phone. Alternatively the transceiver can call a pre-programmed phone number corresponding to a telephone. While the call is being place the GUI renders a screen 194 that shows the telephone's receiver being off the hook. A third screen 196 indicates that the patient is connected to the clinician. The call is terminated when the patient finishes talking to the clinician and taps the screen. Alternatively, the transceiver can include software that detects that no further voice communications are taking place, and then uses this information to terminate the call. In embodiments, the entire call can be stored in a computer memory on either the transceiver or the PDS.

[0070] The GUI operating on the wrist-worn transceiver's touchpanel display can render several other interfaces that facilitate patient monitoring in the hospital. For example, referring to Fig. 15, the GUI can be used to monitor the patient's pain level, a parameter often considered by clinicians to be as important as vital signs for characterizing a patient. The GUI 200 shown in the figure features a simple series of icons that provide a relative indication of the patient's pain level. An index value of 0 (corresponding to a 'happy' face) indicates a low level of pain; an index value of 10 (corresponding to a 'sad' face) indicates a high level of pain. During a measurement, the patient simply touches the icon that best characterizes their pain level. The numerical value corresponding to this level is then wirelessly transmitted back to the PDS and stored in the patient's EMR. The GUI, for example, may be automatically rendered periodically (e.g. every hour) on the transceiver to continuously monitor the patient's pain level. In other embodiments, the GUI could render a graphical display that provides a more sophisticated metric for determining the patient's pain, such as the McGill Pain Questionnaire. This system described in the following journal article: 'The McGill Pain Questionnaire: Major Properties and Scoring Methods', Melzak, Pain, 1:277-299 (1975).

[0071] In a similar manner, the GUI can be used to gauge the patient's level of mentation, i.e. mental activity. Mentation has been consistently shown to be a valuable tool for diagnosing a patient, but is typically determined empirically by a clinician during a check-up or hospital visit. Such a diagnosis is somewhat arbitrary and requires the clinician to meet face-to-face with the patient, which is often impractical. But with the wrist-worn transceiver, diagnosis of mentation can be made automatically at the patient's bedside without a clinician needing to be present. Fig. 16, for example, shows a GUI 202 that provides a simple 'mentation test' for the patient to complete. In this case, the mentation test involves a graphical representation of a series of non-sequential numbers. The patient completes this test by tapping on the numbers rendered by the touchpanel display in their numerical or-

der. An algorithm then 'scores' the test based on accuracy and the time required to complete it. Once determined, the score is wirelessly transmitted back to the PDS, and then stored in the patient's EMR. Other simple tests with varying complexity can be used in place of that shown in Fig. 16. The tests can vary depending on the specific mentation function to be tested. For example, unique tests can be generated for patients with head injuries, cardiac patients, patients in severe pain, Alzheimer's patients, etc. In all cases, the tests are designed to make a quantitative assessment of the patient's mental status; the transceiver sends a numerical value representing this parameter and an identifier for the test back to the EMR for analysis. The transceiver can be programmed so that the GUI 202 for the mentation test, like the GUI 200 for pain level shown in Fig. 15, is automatically rendered at basically any time interval on the touch-panel display. This time interval can be periodic and on an hourly basis, once/day, etc.

[0072] As shown in Fig. 17, the transceiver can include a GUI 204 that displays a photograph or video of the patient. The photograph could be taken by a digital camera within the transceiver, or with an external camera and then transferred to the transceiver through a variety of means, e.g. the hospital's wireless network, a peer-to-peer wireless connection, using a non-volatile memory such as an SD card, or even using a data-transfer process initiated by the 'bump' methodology described above. In general, the same means used to port a photograph from a standard digital camera to a personal computer or other device can be used in this application. Once the photograph is received, software on the transceiver displays it in either a default screen (e.g., in place of the 'beating heart' shown in Figs. 1 and 3), or when the GUI 204 is activated through a tap of a corresponding icon. Displaying the patient's photograph in this manner provides a visual indicator which the clinician can use to correctly identify the patient. In other embodiments, a photograph of someone associated with the patient (e.g. a relative) can also be displayed on the GUI 204. Such an embodiment may be particularly useful for neo-natal hospitals wards, wherein one or more photographs of an infant's parents could be displayed on a transceiver attached to the infant. This way a clinician could check the photograph to ensure that visitors to the neo-natal hospital ward are, in fact, the infant's parents.

[0073] Figs. 18 and 19 show other GUIs 206, 208 that can be rendered on the wrist-worn transceiver's display to carry out basic features in the hospital, such as meal ordering (Fig. 18), and changing the channel on a television or computer (Fig. 19). In these cases, the PDS associated with the transceiver receives a packet describing the function at hand (e.g., the meal that has been ordered, or the channel that is desired), and communicates with another software application in the hospital to complete the transaction. This communication, for example, can take place using a XML-based Web Services operation, such as that described in the following patent

application US 2005/0216199 A1. In related embodiments, a GUI similar to that shown in Fig. 19 can be used to order movies, video games, television programs stored on a digital video recorder, books, and music. Content corresponding to these components is typically stored on a remote server, and then accessed using an XML-based operation, as described above.

[0074] In yet another application, as shown in Fig. 20, the wrist-worn transceiver 72 and its associated barcode scanner 102 can be used to check medication before it is administered to the patient. In this embodiment, barcodes associated with the patient 63, clinician 65, and the medication 67 are read by the barcode scanner 102 within the transceiver 72. The transceiver then wirelessly transmits decoded barcode information through a local access point 56 and to the PDS connected to the hospital network 60. Using a software program, the PDS analyzes these data and communicates with the patient's record in the hospital EMR 58 to determine if the medication is appropriate for the patient. For example, the software program may check to see if the patient is allergic to the medication, if the dosage is correct, or if the patient has previously exhibited any detrimental side effects that may affect the dosage. In related embodiments, the transceiver may also include a GUI wherein the clinician enters ancillary information, such as the dosage of the medication or demographic information describing the patient, using a 'soft' keypad. Or the GUI may include a simple questionnaire that guides the clinician through the process of checking the medication, and then administering it. In still other embodiments, the infusion pump that delivers the medication may include a wireless connection through the access point 56 to the PDS 60 or to the transceiver 72 to automatically supply information related to the medication to the software program.

[0075] Once the software program determines that it is safe to administer the medication, it sends a packet from the PDS 60, through the access point 56, and back to the transceiver 72, which then renders a GUI instructing the clinician to proceed. In other embodiments, the PDS 60 sends the packet through the access point 56 to either a remote computer 62 (e.g. a tablet computer) or a portable device 64 (e.g. a cellular telephone or personal digital assistant).

Form Factor of the Body-Worn Monitor

[0076] Figs. 21A and 21B show how the body-worn monitor 100 described above attaches to a patient 70 to measure RR, SpO₂, cNIBP, and other vital signs. These figures show two configurations of the system: Fig. 21A shows the system used during the indexing portion of the Composite Technique, and includes a pneumatic, cuff-based system 85, while Fig. 21B shows the system used for subsequent measurements. The indexing measurement typically takes about 60 seconds, and is typically performed once every 4-8 hours. Once the indexing measurement is complete the cuff-based system 85 is

typically removed from the patient. The remainder of the time the monitor 100 performs the RR, HR, SpO₂ and cNIBP measurements.

[0077] The body-worn monitor 100 features a wrist-worn transceiver 72, described in more detail in Figs. 22A and 22B, featuring a touch panel interface 73 that displays the various GUIs described above and in Fig. 24. A wrist strap 90 affixes the transceiver 72 to the patient's wrist like a conventional wristwatch. A flexible cable 92 connects the transceiver 72 to an optical sensor 94 that wraps around the base of the patient's thumb. During a measurement, the optical sensor 94 generates a time-dependent PPG waveform which is processed along with an ECG to measure cNIBP, SpO₂, and, in some applications, RR. To determine ACC waveforms the body-worn monitor 100 features three separate accelerometers located at different portions on the patient's arm and chest. The first accelerometer is surface-mounted on a circuit board in the wrist-worn transceiver 72 and measures signals associated with movement of the patient's wrist. As described above, this motion can also be indicative of that originating from the patient's fingers, which will affect the SpO₂ measurement. The second accelerometer is included in a small bulkhead portion 96 included along the span of the cable 82. During a measurement, a small piece of disposable tape, similar in size to a conventional bandaid, affixes the bulkhead portion 96 to the patient's arm. In this way the bulkhead portion 96 serves two purposes: 1) it measures a time-dependent ACC waveform from the midportion of the patient's arm, thereby allowing their posture and arm height to be determined as described in detail above; and 2) it secures the cable 82 to the patient's arm to increase comfort and performance of the body-worn monitor 100, particularly when the patient is ambulatory. The third accelerometer is mounted in the sensor module 74 that connects through cables 80a-c to ECG electrodes 78a-c. Signals from these sensors are then digitized, transmitted through the cable 82 to the wrist-worn transceiver 72, where they are processed with an algorithm as described above to determine RR.

[0078] The cuff-based module 85 features a pneumatic system 76 that includes a pump, valve, pressure fittings, pressure sensor, manifold, analog-to-digital converter, microcontroller, and rechargeable Li:ion battery. During an indexing measurement, the pneumatic system 76 inflates a disposable cuff 84 and performs two measurements according to the Composite Technique: 1) it performs an inflation-based measurement of oscillometry and measurement of a corresponding OSC waveform to determine values for SYS, DIA, and MAP; and 2) it determines a patient-specific relationship between PTT and MAP. These measurements are described in detail in the co-pending patent application US 2009/0018453 A1.

[0079] The cuff 84 within the cuff-based pneumatic system 85 is typically disposable and features an internal, airtight bladder that wraps around the patient's bicep to deliver a uniform pressure field. During the indexing

measurement, pressure values are digitized by the internal analog-to-digital converter, and sent through a cable 86 according to a CAN protocol, along with SYS, DIA, and MAP blood pressures, to the wrist-worn transceiver 72 for processing as described above. Once the cuff-based measurement is complete, the cuff-based module 85 is removed from the patient's arm and the cable 86 is disconnected from the wrist-worn transceiver 72. cNIBP is then determined using PTT, as described in detail above.

[0080] To determine an ECG, the body-worn monitor 100 features a small-scale, three-lead ECG circuit integrated directly into the sensor module 74 that terminates an ECG cable 82. The ECG circuit features an integrated circuit that collects electrical signals from three chest-worn ECG electrodes 78a-c connected through cables 80a-c. As described above, the ECG electrodes 78a-c are typically disposed in a conventional Einthoven's Triangle configuration, which is a triangle-like orientation of the electrodes 78a-c on the patient's chest that features three unique ECG vectors. From these electrical signals the ECG circuit determines up at least three ECG waveforms, each corresponding to a unique ECG vector, which are digitized using an analog-to-digital converter mounted proximal to the ECG circuit and sent through the cable 82 to the wrist-worn transceiver 72 according to the CAN protocol. There, the ECG and PPG waveforms are processed to determine the patient's blood pressure. HR and RR are determined directly from the ECG waveform using known algorithms, such as those described above. More sophisticated ECG circuits (e.g. five and twelve-lead systems) can plug into the wrist-worn transceiver to replace the three-lead system shown in Figs. 21A and 21B.

[0081] Figs. 22A, 22B show three-dimensional views of the wrist-worn transceiver 72 before and after receiving cables 82, 86, 89 from sensors worn on the patient's upper arm and torso, as well as the cable 92 that connects to the optical sensor. The transceiver 72 is sealed in a water-proof plastic casing 117 featuring electrical interconnects (not shown in the figure) on its bottom surface that interface to the terminal ends 111, 119a-c of cables 82, 86, 89, 92 leading to the monitor's various sensors. The electrical interconnects support serial communication through the CAN protocol, described in detail herein, particularly with reference to Fig. 25. During operation, the transceiver's plastic casing 117 snaps into a plastic housing 106, which features an opening 109 on one side to receive the terminal end 111 of the cable 92 connected to the optical sensor. On the opposing side the plastic housing 106 features three identical openings 104a-c that receive the terminal ends 119a-c of cables 82, 86, 89 connected to the ECG and accelerometer systems (cable 82), the pneumatic cuff-based system (cable 86), and ancillary systems (cable 89) described above. In addition to being waterproof, this design facilitates activities such as cleaning and sterilization, as the transceiver contains no openings for fluids common in the hospital, such

as water and blood, to flow inside. During a cleaning process the transceiver 72 is simply detached from the plastic housing 106 and then cleaned.

[0082] The transceiver 72 attaches to the patient's wrist using a flexible strap 90 which threads through two D-ring openings in the plastic housing 106. The strap 90 features mated Velcro patches on each side that secure it to the patient's wrist during operation. A touchpanel display 50 renders the various GUIs described above.

[0083] The electrical interconnects on the transceiver's bottom side line up with the openings 104a-c, and each supports the CAN protocol to relay a digitized data stream to the transceiver's internal CPU, as described in detail with reference to Fig. 25. This allows the CPU to easily interpret signals that arrive from the monitor's body-worn sensors, and means that these connectors are not associated with a specific cable. Any cable connecting to the transceiver 72 can be plugged into any opening 104a-c. As shown in Fig. 22A, the first opening 104a receives the cable 82 that transports digitized ECG waveforms determined from the ECG circuit and electrodes, and digitized ACC waveforms measured by accelerometers in the cable bulkhead and the bulkhead portion associated with the ECG cable 82.

[0084] The second opening 104b receives the cable 86 that connects to the pneumatic cuff-based system used for the pressure-dependent indexing measurement. This connector receives a time-dependent pressure waveform delivered by the pneumatic system to the patient's arm, along with values for SYS, DIA, and MAP determined during the indexing measurement. The cable 86 unplugs from the opening 104b once the indexing measurement is complete, and is plugged back in after approximately 4-8 hours for another indexing measurement.

[0085] The final opening 104c can be used for an auxiliary device, e.g. a glucometer, infusion pump, body-worn insulin pump, ventilator, or end-tidal CO₂ monitoring system. As described with reference to Fig. 25, digital information generated by these systems will include a header that indicates their origin so that the CPU can process them accordingly.

Measuring and Displaying Time-Dependent Physiological Signals

[0086] Figs. 23A and 23B show how a network of sensors 78a-c, 83, 84, 87, 94 within the body-worn monitor 100 connect to a patient 70 to measure time-dependent ECG 261, PPG 262, OSC 263, ACC 264, and RR 265 waveforms. These, in turn, yield the patient's vital signs and motion parameters. Each waveform 261-265 relates to a unique physiological characteristic of the patient 70. For example, each of the patient's heartbeats generates electrical impulses that pass through the body near the speed of light, along with a pressure wave that propagates through the patient's vasculature at a significantly slower speed. Immediately after the heartbeat, the pres-

sure wave leaves the heart 148 and aorta 149, passes through the subclavian artery 150 to the brachial artery 144, and from there through the radial and ulnar arteries 145 to smaller arteries in the patient's fingers. Three disposable electrodes 78a-c attached to the patient's chest measure unique electrical signals which pass to a single-chip ECG circuit 83 that terminates a distal end of the ECG cable. Typically, these electrodes attach to the patient's chest in a conventional 'Einthoven's triangle' configuration featuring three unique 'vectors', each corresponding to a different lead (e.g. LEAD I, II, III). Related configurations can also be used when five and twelve-lead ECG systems are used in place of the three-lead system, as described above with reference to Figs. 21A, 21B. Within the ECG circuit 83 signals are processed using an amplifier/filter circuit and analog-to-digital converter to generate a digital ECG waveform 261 corresponding to each lead. The ECG waveform 261 features a sharp, well-defined QRS complex corresponding to each heartbeat; this marks the initiation of the patient's cardiac cycle. Heart rate is determined directly from the ECG waveform 261 using known algorithms, such as those described in the following journal article, 'ECG Beat Detection Using Filter Banks', Afonso et al., IEEE Trans. Biomed Eng., 46:192-202 (1999).

[0087] To generate an IP waveform 265, one of the ECG electrodes in the circuit 78a is a 'driven lead' that injects a small amount of modulated current into the patient's torso. A second, non-driven electrode 78c, typically located on the opposite side of the torso, detects the current, which is further modulated by capacitance changes in the patient's chest cavity resulting from breathing. Further processing and filtering of the IP waveforms 265 yields respiratory rate. Respiration can also be determined using an adaptive filtering approach that processes both the IP waveform and ACC waveform 264, as described in more detail in the following co-pending patent application, US 2011/0066037.

[0088] The optical sensor 94 features two LEDs and a single photodetector that collectively measure a time-dependent PPG waveform 262 corresponding to each of the LEDs. The sensor and algorithms for processing the PPG waveforms are described in detail in the following co-pending patent application US 2010/0324385 A1. The waveform 262 represents a time-dependent volumetric change in vasculature (e.g. arteries and capillaries) that is irradiated with the sensor's optical components. Volumetric changes are induced by a pressure pulse launched by each heartbeat that travels from the heart 148 to arteries and capillaries in the thumb according to the above-described arterial pathway. Pressure from the pressure pulse forces a bolus of blood into this vasculature, causing it to expand and increase the amount of radiation absorbed, and decrease the transmitted radiation at the photodetector. The pulse shown in the PPG waveform 262 therefore represents the inverse of the actual radiation detected at the photodetector. It follows the QRS complex in the ECG waveform 261, typically by

about one to two hundred milliseconds. The temporal difference between the peak of the QRS complex and the foot of the pulse in the PPG waveform 262 is the PTT, which as described in detail below is used to determine blood pressure according to the Composite Technique. PTT-based measurements made from the thumb yield excellent correlation to blood pressure measured with a femoral arterial line. This provides an accurate representation of blood pressure in the central regions of the patient's body.

[0089] Each accelerometer generates three time-dependent ACC waveforms 264, corresponding to the x, y, and z-axes, which collectively indicate the patient's motion, posture, and activity level. The body-worn monitor, as described above, features three accelerometers that attach to the patient: one in the wrist-worn transceiver 72, one in the ECG circuit 83, and one near the bicep 87 that is included in the cable connecting these two components. The frequency and magnitude of change in the shape of the ACC waveform 264 indicate the type of motion that the patient is undergoing. For example, the waveform 264 can feature a relatively time-invariant component indicating a period of time when the patient is relatively still, and a time-variant component when the patient's activity level increases. Magnitudes of both components will depend on the relationship between the accelerometer and a gravity vector, and can therefore be processed to determine time-invariant features, such as posture and arm height. A frequency-dependent analysis of the time-variant components yields the type and degree of patient motion. Analysis of ACC waveforms 264 is described in detail in the above-mentioned patent applications.

[0090] The OSC waveform 263 is generated from the patient's brachial artery 144 with the pneumatic system and a cuff-based sensor 84 during the pressure-dependent portion of the Composite Technique. It represents a time-dependent pressure which is applied to the brachial artery during inflation and measured by a digital pressure sensor within the pneumatic system. The waveform 263 is similar to waveforms measured during deflation by conventional oscillometric blood pressure monitors. During a measurement, the pressure waveform 263 increases in a mostly linear fashion as pressure applied by the cuff 84 to the brachial artery 144 increases. When it reaches a pressure slightly below the patient's diastolic pressure, the brachial artery 144 begins to compress, resulting in a series time-dependent pulsations caused by each heartbeat that couple into the cuff 84. The pulsations modulate the OSC waveform 263 with an amplitude that varies in a Gaussian-like distribution, with maximum modulation occurring when the applied pressure is equivalent to the patient's MAP. The pulsations can be filtered out and processed using digital filtering techniques, such as a digital bandpass filter that passes frequencies ranging from 0.5 - 20 Hz. The resulting waveform can be processed to determine SYS, DIA, and MAP, as is described in detail in the above-referenced patent applications, the

contents of which have been previously incorporated herein by reference. The cuff 84 and pneumatic system are removed from the patient's bicep once the pressure-dependent component of the Composite Technique is complete.

[0091] The high-frequency component of the OSC waveform 263 (i.e. the pulses) can be filtered out to estimate the exact pressure applied to the patient's brachial artery during oscillometry. According to the Composite Technique, PTT measured while pressure is applied will gradually increase as the brachial artery is occluded and blood flow is gradually reduced. The pressure-dependent increase in PTT can be fit with a model to estimate the patient-specific relationship between PTT and blood pressure. This relationship, along with SYS, MAP, and DIA determined from the OSC waveform during inflation-based oscillometry, is used during the Composite Technique's pressure-free measurements to determine blood pressure directly from PTT.

[0092] There are several advantages to making the indexing measurement during inflation, as opposed to deflation. Measurements made during inflation are relatively fast and comfortable compared to those made during deflation. Inflation-based measurements are possible because of the Composite Technique's relatively slow inflation speed (typically 5 - 10 mmHg/second) and the high sensitivity of the pressure sensor used within the body sensor. Such a slow inflation speed can be accomplished with a small pump that is relatively lightweight and power efficient. Moreover, measurements made during inflation can be immediately terminated once systolic blood pressure is calculated. This tends to be more comfortable than conventional cuff-based measurements made during deflation. In this case, the cuff typically applies a pressure that far exceeds the patient's systolic blood pressure; pressure within the cuff then slowly bleeds down below the diastolic pressure to complete the measurement.

[0093] A digital temperature sensor proximal to the ECG circuit 83 measures the patient's skin temperature at their torso. This temperature is an approximation of the patient's core temperature, and is used mostly for purposes related to trending and alarms/alerts.

[0094] Fig. 24 shows how the above-described ECG, PPG, and IP waveforms, along with vital signs calculated from them, are rendered using different screens 300, 304, 306, 308 within a GUI. In all cases, the waveforms are displayed with a rolling graphical technique, along with a moving bar that indicates the most current point in time. As per the AAMI/ANSI EC-13 reference standard, the ECG waveforms are displayed alongside a bar that indicates a signal intensity of 1 mV. Screen 308 shows different ECG vectors (corresponding to, e.g., Lead I, II, III, aVR, aVF) that are rendered when the clinician taps the ECG waveform on screen 300, and then the corresponding lead on screen 308. Waveforms for a particular vital sign (e.g. a PPG waveform for the SpO2 measurement; an IP waveform for the RR measurement) are ren-

dered when the clinician taps on the value of the corresponding vital sign. During a measurement both waveforms and the vital signs calculated from them are wirelessly transmitted to the PDS, as described above.

Communicating with Multiple Systems Using the CAN Protocol

[0095] As described above, the ECG, ACC, and pneumatic systems within the body-worn system send digitized information to the wrist-worn transceiver through the CAN protocol. Fig. 25 shows a schematic drawing indicating how CAN packets 201a-d, 212a-e transmitted between these systems facilitate communication. Specifically, each of the ACC 215, ECG 216, pneumatic 220, and auxiliary 245 systems include a separate analog-to-digital converter, microcontroller, frequency-generating crystal oscillator (typically operating at 100 kHz), and real-time clock divider that collectively generate and transmit digital data packets 201a-d according to the CAN protocol to the wrist-worn transceiver 72. Each crystal uses the internal real-time clock on the internal microprocessor within the respective system. This allows the microcontroller within each system to be placed in a low-power state in which its real-time operating system (RTOS) dispatch system indicates that it is not ready to run a task. The real-time clock divider is programmed to create an interrupt which wakes up the microcontroller every 2 milliseconds.

[0096] The wrist-worn transceiver 72 features a 'master clock' that generates real-time clock 'ticks' at the sampling rate (typically 500 Hz, or 2 ms between samples). Each tick represents an incremented sequence number. Every second, the wrist-worn transceiver 72 transmits a packet 212e over the CAN bus that digitally encodes the sequence number. One of the criteria for accurate timing is that the time delay between the interrupt and the transmission of the synchronizing packet 212e, along with the time period associated with the CAN interrupt service routine, is predictable and stable. During initialization, the remote CAN buses do not sleep; they stay active to listen for the synchronization packet 212e. The interrupt service routine for the synchronization packet 212e then establishes the interval for the next 2 millisecond interrupt from its on-board, real-time crystal to be synchronized with the timing on the wrist-worn transceiver 72. Offsets for the packet transmission and interrupt service delays are factored into the setting for the real-time oscillator to interrupt synchronously with the microprocessor on the wrist-worn transceiver 72. The magnitude of the correction factor to the real-time counter is limited to 25% of the 2 millisecond interval to ensure stability of this system, which represents a digital phase-locked loop.

[0097] When receipt of the synchronization packet 212e results in a timing correction offset of either a 0, +1, or -1 count on the remote system's oscillator divider, software running on the internal microcontroller declares that the system is phase-locked and synchronized. At this

point, it begins its power-down operation and enables measurement of data as described above.

[0098] Each remote system is driven with a 100 kHz clock, and a single count of the divider corresponds to 20 microseconds. This is because the clock divider divides the real-time clock frequency by a factor of 2. This is inherent in the microcontroller to ensure that the clock has a 50% duty cycle, and means the clock can drift +/- 20 microseconds before the actual divider chain count will disagree by one count, at which time the software corrects the count to maintain a phase-locked state. There is thus a maximum of 40 microseconds of timing error between data transmitted from the remote systems over the CAN bus. Blood pressure is the one vital sign measured with the body-worn monitor that is calculated from time-dependent waveforms measured from different systems (e.g. PPG and ECG waveforms). For this measurement, the maximum 40-microsecond timing error corresponds to an error of +/- 0.04 mmHg, which is well within the error (typically +/- 5mmHg) of the measurement.

[0099] In order to minimize power consumption, the wrist-worn transceiver 72 and remote systems 215, 216, 220, 245 power down their respective CAN bus transceivers between data transfers. During a data transfer, each system generates a sequence number based that is included in the synchronization packet 212e. The sequence number represents the interval between data transfers in intervals of 2 milliseconds. It is a factor of 500 (e.g. 2, 4, 5, 10) that is the number of 2 millisecond intervals between transfers on the CAN bus. Each remote system enables its CAN bus during the appropriate intervals and sends its data. When it has finished sending its data, it transmits a 'transmit complete' packet indicating that the transmission is complete. When a device has received the 'transmit complete' packet it can disable its CAN transceiver to further reduce power consumption.

[0100] Software in each of the ACC 215, ECG 216, pneumatic 220, and auxiliary 245 systems receive the sequence packet 212e and the corresponding sequence number, and set their clocks accordingly. There is typically some inherent error in this process due to small frequency differences in the crystals (from the ideal frequency of 100 kHz) associated with each system. Typically this error is on the order of microseconds, and has only a small impact on time-dependent measurements, such as PTT, which are typically several hundred milliseconds.

[0101] Once timing on the CAN bus is established using the above-described procedure, each of the ACC 215, ECG 216, and pneumatic 220 systems generate time-dependent waveforms that are transmitted in packets 201a-d, each representing an individual sample. Each packet 201a-d features a header portion which includes the sequence number 212a-d and an initial value 210a-d indicating the type of packet that is transmitted. For example, accelerometers used in the body-worn system are typically three-axis digital accelerometers, and

generate waveforms along the x, y, and z-axes. In this case, the initial value 210a encodes numerical values that indicate: 1) that the packet contains ACC data; and 2) the axis (x, y, or z) over which these data are generated. Similarly, the ECG system 216 can generate a time-dependent ECG waveform corresponding to Lead I, II, or III, each of which represents a different vector measured along the patient's torso. Additionally, the ECG system 216 can generate processed numerical data, such as heart rate (measured from time increments separating neighboring QRS complexes), respiratory rate (from an internal impedance pneumography component), as well as alarms calculated from the ECG waveform that indicate problematic cardiovascular states such as VTAC, VFIB, and PVCs. Additionally, the ECG system can generate error codes indicating, for example, that one of the ECG leads has fallen off. The ECG system typically generates an alarm/alert, as described above, corresponding to both the error codes and potentially problematic cardiovascular states. In this case, the initial value 210b encodes numerical values that indicate: 1) that the packet contains ECG data; 2) the vector (Lead I, II, or III) corresponding to the ECG data; and 3) an indication if a cardiovascular state such as VTAC, VFIB, or PVCs was detected.

[0102] The pneumatic system 220 is similar to the ECG system in that it generates both time-dependent waveforms (i.e. a pressure waveform, measured during oscillometry, characterizing the pressure applied to the arm and subsequent pulsations measured during an oscillometric measurement) and calculated vital signs (SYS, DIA, and MAP measured during oscillometry). In some cases errors are encountered during the oscillometric blood pressure measurement. These include, for example, situations where blood pressure is not accurately determined, an improper OSC waveform, over-inflation of the cuff, or a measurement that is terminated before completion. In these cases the pneumatic system 220 generates a corresponding error code. For the pneumatic system 220 the initial value 210c encodes numerical values that indicate: 1) that the packet contains blood pressure data; 2) an indication that the packet includes an error code.

[0103] In addition to the initial values 210a-d, each packet 201a-d includes a data field 214a-d that encodes the actual data payload. Examples of data included in the data fields 214a-d are: 1) sampled values of ACC, ECG, and pressure waveforms; 2) calculated heart rate and blood pressure values; and 3) specific error codes corresponding to the ACC 215, ECG 216, pneumatic 220, and auxiliary 225 systems.

[0104] Upon completion of the measurement, the wrist-worn transceiver 72 receives all the CAN packets 201a-d, and synchronizes them in time according to the sequence number 212a-d and identifier 210a-d in the initial portions 216 of each packet. Every second, the CPU updates the time-dependent waveforms and calculates the patient's vital signs and motion-related proper-

ties, as described above. Typically these values are calculated as a 'rolling average' with an averaging window ranging from 10-20 seconds. The rolling average is typically updated every second, resulting in a new value that is displayed on the wrist-worn transceiver 72. Each packet received by the transceiver 72 is also wirelessly retransmitted as a new packet 201b' through a wireless access point 56 and to both an PDS and RVD within a hospital network 60. The new packet 201b' includes the same header 210b', 212b' and data field information 214b' as the CAN packets transmitted between systems within the body-worn monitor. Also transmitted are additional packets encoding the cNIBP, SpO2, and processed motion states (e.g. posture, activity level, degree of motion), which unlike heart rate and SYS, DIA, and MAP are calculated by the CPU in the wrist-worn transceiver. Upon receipt of the packet 201b', the RVD displays vital signs, waveforms, motion information, and alarms/alerts, typically with a large monitor that is easily viewed by a clinician. Additionally the PDS can send information through the hospital network (e.g. in the case of an alarm/alert), store information in an internal database, and transfer it to a hospital EMR.

25 Alternate IT Configurations

[0105] Fig. 26 shows an alternate configuration of the invention wherein the transceiver 72 transmits both voice and data information through a wireless access point 56A and to the Internet 55, and from there to the hospital network and PDS 60. Such a configuration would be used, for example, when the patient is located outside of the hospital (e.g. at home). It allows clinicians to monitor and care for a patient as if they were located in the hospital. Once information arrives at the PDS 60, it can be transferred to the hospital EMR system 58, or through a wireless access point 56B within the hospital to an external computer 62 or a portable device 64.

[0106] In an alternate embodiment, as shown in Fig. 27, the first wireless access point 56A shown in Fig. 26 is replaced by a wireless modem 64A, such as a cellular telephone or personal digital assistant. Here, the wireless modem 64A receives voice and data information from the transceiver through a peer-to-peer wireless interface (e.g. an interface based on 802.11b/g or 802.15.4). The wireless modem 64A then transmits the voice and data information to the Internet 55 using, e.g., a cellular connection, such as one based on GSM or CDMA. In yet another embodiment, as shown in Fig. 28, the transceiver 72 includes an internal long-range wireless transmitter based on a cellular protocol (e.g. GSM or CDMA), allowing it to transmit voice and data information directly to the Internet 55. In the embodiments shown in both Figs. 27 and 28, information sent through the Internet is ultimately received by the PDS 60, and is sent from there through a wireless access point 56 to either the remote computer 62 or portable device 64.

[0107] In embodiments, the transceiver 72 features

multiple wireless transmitters, and can operate in multiple modes, such as each of those shown in Figs. 26-28. In this case the wireless protocol (based on, e.g. 802.11 or cellular) is manually selected using the GUI, or automatically selected based on the strength of the ambient wireless signal.

Other examples of the invention

[0108] In addition to those methods described above, the body-worn monitor can use a number of additional methods to calculate blood pressure and other properties from the optical and electrical waveforms. These are described in the following co-pending patent applications US 2005/0228297 A1, US 2005/0228296 A1, US 2005/0228300 A1, US 2005/0228301 A1, US 2006/0084878 A1, US 2005/0228299 A1, US 2005/0261598 A1, US 2006/0009697 A1, US 2006/009698 A1, US 2007/018593 A1, US 2007/0276261 A1, US 2007/0276632 A1, US 2008/0082004 A1, US 2008/0114220 A1, US 2008/0221399 A1.

[0109] Other examples are also within the scope of the invention. For example, other measurement techniques, such as conventional oscillometry measured during deflation, can be used to determine SYS, DIA, and MAP for the above-described algorithms. Additionally, processing units and probes for measuring pulse oximetry similar to those described above can be modified and worn on other portions of the patient's body. For example, optical sensors with finger-ring configurations can be worn on fingers other than the thumb. Or they can be modified to attach to other conventional sites for measuring SpO₂, such as the ear, forehead, and bridge of the nose. In these embodiments the processing unit can be worn in places other than the wrist, such as around the neck (and supported, e.g., by a lanyard) or on the patient's waist (supported, e.g., by a clip that attaches to the patient's belt). In still other embodiments the probe and processing unit are integrated into a single unit.

[0110] In examples, the interface rendered on the display at the central nursing station features a field that displays a map corresponding to an area with multiple sections. Each section corresponds to the location of the patient and includes, e.g., the patient's vital signs, motion parameter, and alarm parameter. For example, the field can display a map corresponding to an area of a hospital (e.g. a hospital bay or emergency room), with each section corresponding to a specific bed, chair, or general location in the area.

[0111] Further embodiments of the invention are within the scope of the following claims:

Claims

1. A method for monitoring a patient, comprising the following steps:

- (a) associating a first set of vital sign information with the patient, the first set of vital sign information measured from the patient with a first monitor comprising a first motion sensor;
- (b) storing the first set of vital sign information associated with the patient in a computer memory;
- (c) contacting the first monitor with a second monitor comprising a second motion sensor, the contacting causing the first motion sensor to generate a first motion signal, and the second motion sensor to generate a second motion signal;
- (d) wirelessly transmitting both the first and second motion signals to a remote server
- (e) collectively processing the first and second motion signals on the remote server to determine that the first monitor is to be replaced by the second monitor; and
- (f) associating a second set of vital sign information with the patient, the second set of vital sign information measured from the patient with the second monitor.

2. A system for measuring a patient, comprising:

- a first patient monitor comprising a first motion sensor and a first wireless transmitter;
- a second patient monitor comprising a second motion sensor and a second wireless transmitter;
- a remote server connected to the first patient monitor through the first wireless transmitter, and to the second patient monitor through the second wireless transmitter, the remote server configured to receive a first motion signal generated with the first motion sensor and a second motion signal generated with the second motion sensor after the first and second patient monitors contact each other;
- wherein the remote server is configured to associate a single patient with a first set of information measured with the first patient monitor and a second set of information measured with the second patient monitor.

3. The method of claim 1, wherein the first motion sensor comprises a first accelerometer, and the second motion sensor comprises a second accelerometer.

4. The method of claim 3, wherein the first motion signal is a first time-dependent waveform, and the second motion signal is a second time-dependent waveform, wherein the first time-dependent waveform optionally comprises a first pulse, and the second time-dependent waveform comprises a second pulse, wherein the processing step optionally further comprises determining that the first pulse and the second

pulse are generated during a common period of time or
 wherein the processing step optionally further comprises determining that a first amplitude of the first pulse and a second amplitude of the second pulse both exceed a pre-determined threshold value.

5. The method of claim 1, further comprising the step of wirelessly transmitting instruction information from the remote server to at least one of the first monitor and the second monitor indicating that the first monitor is to be replaced by the second monitor or wherein both the first and second sets of vital sign information are optionally stored in a computer memory on the remote computer, wherein the computer memory on the remote computer optionally is a database.
6. The method of claim 1, wherein both the first and second sets of vital sign information comprise values for at least one of the following properties: blood pressure, heart rate, respiratory rate, SpO₂, temperature.
7. The method of claim 1, wherein both the first and second sets of vital sign information comprise at least one of the following time-dependent waveforms: PPG, ECG, OSC, IP.
8. The method of claim 1, further comprising the step of associating patient demographic information describing the patient with both the first and second sets of vital sign information, wherein the patient demographic information comprises at least one of the following parameters: name, gender, weight, height, date of birth.
9. A method for monitoring a patient, comprising the following steps:
 - (a) contacting at least one of a display device and an area proximal to the display device with a patient monitor comprising a motion sensor and a wireless transmitter, the contacting causing the motion sensor to generate a motion signal;
 - (b) wirelessly transmitting the motion signal to a remote computer over a wireless network using the wireless transmitter;
 - (c) processing the motion signal with the computer to associate the patient monitor with the display device;
 - (d) measuring at least one set of vital sign information from the patient with the patient monitor; and
 - (e) displaying the at least one set of vital sign information on the display device.

10. The method of claim 9, wherein wireless signals received by multiple wireless access points on the wireless network are processed to determine an approximate location of at least one of the patient monitor and the display device, and the motion signal and the approximate location of at least one of the patient monitor and the display device are processed to associate the patient monitor with the display device.

11. A system for measuring a patient, comprising:

a patient monitor comprising a motion sensor and a wireless transmitter;
 a display device;
 a remote server connected to the patient monitor through the wireless transmitter and to the display device through a network or a computer connected to the patient monitor and to the display device through a network, the computer, the remote server and the computer, respectively, configured to receive a motion signal generated with the motion sensor after the patient monitor contacts at least one of the display device and an area proximal to the display device; wherein the remote server is configured to associate the patient monitor with the display device so that the display device receives vital sign information measured by the patient monitor and transmitted through the network, the algorithm comprising collectively analyzing the motion signal and a location of the display device.

12. The method of claim 9, wherein the motion sensor comprises an accelerometer.
13. The method of claim 9, wherein the motion signal is a time-dependent waveform, wherein the time-dependent waveform optionally comprises a pulse.
14. The method of claim 9, wherein the processing step further comprises processing the motion signal and a location of the display device to associate the patient monitor with the display device.
15. The method of claim 14, further comprising a step for determining the location of the display device, wherein the patient monitor optionally further comprises a wireless transmitter, wherein the wireless transmitter is optionally configured to operate on a wireless network, wherein
 - (a) the step for determining the location of the display device optionally further comprises analyzing signals characterizing a signal strength between multiple access points operating on the wireless network and the wireless transmitter, wherein the step for determining the location of

the display device optionally further comprises triangulating at least three values of signal strength, with each value of signal strength characterizing a signal between an individual access point operating on the wireless network and the wireless transmitter, wherein the step for determining the location of the display device optionally further comprises comparing an approximate location value determined by triangulating the at least three values of signal strength with a map grid, wherein the map grid optionally comprises a true location value of the display device, wherein the step for determining the location of the display device optionally further comprises comparing an approximate location value that is proximal to a true location value comprised by the map grid, wherein the step for determining the location of the display device optionally further comprises comparing an approximate location value to a true location value within a pre-determined radius, wherein the pre-determined radius optionally is between 1-5m or (b) the computer optionally is a remote computer configured to receive signals from the patient monitor through the wireless transmitter,

wherein the display device optionally is configured to receive signals from the remote computer through the network.

Patentansprüche

1. Verfahren zum Überwachen eines Patienten, aufweisend die folgenden Schritte:

- (a) Zuordnen einer ersten Menge von Vitalparameterinformationen zu dem Patienten, wobei die erste Menge von Vitalparameterinformationen des Patienten mit einem ersten Überwachungsgerät gemessen wird, das einen ersten Bewegungssensor umfasst;
- (b) Speichern der dem Patienten zugeordneten ersten Menge von Vitalparameterinformationen in einem Rechnerspeicher;
- (c) Kontaktieren des ersten Überwachungsgeräts mit einem zweiten Überwachungsgerät, das einen zweiten Bewegungssensor umfasst, wobei das Kontaktieren bewirkt, dass der ersten Bewegungsfühler ein erstes Bewegungssignal erzeugt und der zweite Bewegungssensor ein zweites Bewegungssignal erzeugt;
- (d) drahtloses Übertragen von sowohl dem ersten als auch dem zweiten Bewegungssignal an einen Remote-Server;

- (e) gemeinsames Verarbeiten des ersten und zweiten Bewegungssignals auf dem Remote-Server, um zu bestimmen, dass das erste Überwachungsgerät durch das zweite Überwachungsgerät zu ersetzen ist; und
- (f) Zuordnen einer zweiten Menge von Vitalparameterinformationen zu dem Patienten, wobei die zweite Menge von Vitalparameterinformationen des Patienten mit dem zweiten Überwachungsgerät gemessen wird.

2. System zum Messen eines Patienten, aufweisend:

- ein erstes Patientenüberwachungsgerät, das einen ersten Bewegungssensor und einen ersten drahtlosen Sender aufweist;
- ein zweites Patientenüberwachungsgerät, das einen zweiten Bewegungssensor und einen zweiten drahtlosen Sender aufweist;
- einen Remote-Server, der mit dem ersten Patientenüberwachungsgerät durch den ersten drahtlosen Sender und mit dem zweiten Patientenüberwachungsgerät durch den zweiten drahtlosen Sender in Verbindung steht, wobei der Remote-Server so eingerichtet ist, ein erstes, mit dem ersten Bewegungssensor erzeugtes Bewegungssignal und ein zweites, mit dem zweiten Bewegungssensor erzeugtes Bewegungssignal zu empfangen, nachdem sich das erste und zweite Patientenüberwachungsgerät kontaktieren;
- wobei der Remote-Server eingerichtet ist, einen einzelnen Patienten einer mit dem ersten Patientenüberwachungsgerät gemessenen ersten Informationsmenge und einer mit dem zweiten Patientenüberwachungsgerät gemessenen zweiten Informationsmenge zuzuordnen.

3. Verfahren nach Anspruch 1, wobei der erste Bewegungssensor einen ersten Beschleunigungsmesser aufweist, und der zweite Bewegungssensor einen zweiten Beschleunigungsmesser aufweist.

4. Verfahren nach Anspruch 3, wobei das erste Bewegungssignal eine erste zeitabhängige Wellenform ist, und das zweite Bewegungssignal eine zweite zeitabhängige Wellenform ist, wobei die erste zeitabhängige Wellenform optional einen ersten Impuls umfasst, und die zweite zeitabhängige Wellenform einen zweiten Impuls umfasst, wobei der Verarbeitungsschritt ferner optional ein Bestimmen aufweist, dass der erste Impuls und der zweite Impuls während einer gemeinsamen Zeitdauer erzeugt werden oder wobei der Verarbeitungsschritt ferner optional ein Bestimmen aufweist, dass eine erste Amplitude des ersten Impulses und eine zweite Amplitude des zweiten Impulses beide einen vorbestimmten Schwellenwert überschreiten.

5. Verfahren nach Anspruch 1, das ferner den Schritt des drahtlosem Übertragens von Anweisungsinformationen von dem Remote-Server an mindestens das erste Überwachungsgerät und/oder das zweite Überwachungsgerät aufweist, welche angezeigt, dass das erste Überwachungsgerät durch das zweite Überwachungsgerät zu ersetzen ist oder wobei sowohl die erste als auch die zweite Menge von Vitalparameterinformationen optional in einem Rechnerspeicher auf dem Remote-Rechner gespeichert werden, wobei der Rechnerspeicher auf dem entfernt angeordneten Rechner optional eine Datenbank ist.
6. Verfahren nach Anspruch 1, wobei sowohl die erste als auch die zweite Menge von Vitalparameterinformationen Werte für mindestens eine der folgenden Eigenschaften umfassen: Blutdruck, Herzfrequenz, Atemfrequenz, SpO2, Temperatur.
7. Verfahren nach Anspruch 1, wobei sowohl die erste als auch die zweite Menge von Vitalparameterinformationen mindestens eine der folgenden zeitabhängigen Wellenformen aufweisen: PPG, EKG, OSC, IP.
8. Verfahren nach Anspruch 1, ferner den Schritt des Zuordnens von demographischen Patienteninformationen aufweisend, die den Patienten mit sowohl der ersten als auch der zweiten Menge von Vitalparameterinformationen beschreiben, wobei die demographischen Patienteninformationen mindestens einen der folgenden Parameter umfassen: Name, Geschlecht, Gewicht, Größe, Geburtsdatum.
9. Verfahren zum Überwachen eines Patienten, aufweisend die folgenden Schritte:
- (a) Kontaktieren einer Anzeigeeinheit und/oder eines zu der Anzeigeeinheit proximalen Bereichs mit einem Patientenüberwachungsgerät, das einen Bewegungssensor und einen drahtlosen Sender aufweist, wobei das Kontaktieren bewirkt, dass der Bewegungssensor ein Bewegungssignal erzeugt;
 - (b) drahtloses Übertragen des Bewegungssignals an einen Remote-Rechner über ein drahtloses Netzwerk durch Verwenden unter Verwendung des drahtlosen Senders;
 - (c) Verarbeiten des Bewegungssignals mit dem Rechner, um das Patientenüberwachungsgerät der Anzeigeeinheit zuzuordnen;
 - (d) Messen mindestens einer Menge von Vitalparameterinformationen des Patienten mit dem Patientenüberwachungsgerät; und
 - (e) Anzeigen der mindestens einen Menge von Vitalparameterinformationen auf der Anzeigeeinheit.
10. Verfahren nach Anspruch 9, wobei durch mehrere drahtlose Zugangspunkte auf dem drahtlosen Netzwerk empfangene drahtlose Signale verarbeitet werden, um einen ungefähren Standort des Patientenüberwachungsgeräts und/oder der Anzeigeeinheit zu bestimmen, und das Bewegungssignal und der ungefähre Standort des Patientenüberwachungsgeräts und/oder der Anzeigeeinheit verarbeitet werden, um das Patientenüberwachungsgerät der Anzeigeeinheit zuzuordnen.
11. System zum Messen eines Patienten, aufweisend:
- ein Patientenüberwachungsgerät, aufweisend einen Bewegungssensor und einen drahtlosen Sender;
 - eine Anzeigeeinheit;
 - einen Remote-Server, der mit dem Patientenüberwachungsgerät durch den drahtlosen Sender und mit der Anzeigeeinheit durch ein Netzwerk verbunden ist, oder einen mit dem Patientenüberwachungsgerät und mit der Anzeigeeinheit durch ein Netzwerk verbundenen Rechner, wobei jeweils der Rechner, der Remote-Server und der Rechner eingerichtet sind, ein mit dem Bewegungssensor erzeugtes Bewegungssignal zu empfangen, nachdem das Patientenüberwachungsgerät die Anzeigeeinheit und/oder einen zu der Anzeigeeinheit proximalen Bereich kontaktiert;
 - wobei der Remote-Server eingerichtet ist, das Patientenüberwachungsgerät der Anzeigeeinheit zuzuordnen, damit die Anzeigeeinheit durch das Patientenüberwachungsgerät gemessene und durch das Netzwerk übertragene Vitalparameterinformationen empfängt, wobei der Algorithmus gemeinsames Analysieren des Bewegungssignals und eines Standorts der Anzeigeeinheit umfasst.
12. Verfahren nach Anspruch 9, wobei der Bewegungssensor einen Beschleunigungsmesser umfasst.
13. Verfahren nach Anspruch 9, wobei das Bewegungssignal eine zeitabhängige Wellenform ist, wobei die zeitabhängige Wellenform optional einen Impuls umfasst.
14. Verfahren nach Anspruch 9, wobei der Verarbeitungsschritt ferner ein Verarbeiten des Bewegungssignals und eines Standorts der Anzeigeeinheit aufweist, um das Patientenüberwachungsgerät der Anzeigeeinheit zuzuordnen.
15. Verfahren nach Anspruch 14, ferner einen Schritt

zum Bestimmen des Standorts der Anzeigeeinheit aufweisend,
wobei das Patientenüberwachungsgerät ferner optional einen drahtlosen Sender umfasst, wobei der drahtlose Sender optional eingerichtet ist, in einem drahtlosen Netzwerk zu funktionieren, wobei

(a) der Schritt des Bestimmens des Standorts der Anzeigeeinheit optional ferner ein Analysieren von Signalen aufweist, die eine Signalstärke zwischen mehreren Zugangspunkten, die in dem drahtlosen Netzwerk in Betrieb sind, und dem drahtlosen Sender kennzeichnen, wobei der Schritt des Bestimmens des Standorts der Anzeigeeinheit optional ferner ein Triangulieren von mindestens drei Signalstärkewerten umfasst, wobei jeder Signalstärkewert ein Signal zwischen einem individuellen Zugangspunkt, der in dem drahtlosen Netzwerk in Betrieb ist, und dem drahtlosen Sender kennzeichnet

wobei der Schritt zum Bestimmen des Standorts der Anzeigeeinheit ferner optional ein Vergleichen eines ungefähren Standortwerts umfasst, der durch Triangulieren der mindestens drei Signalstärkewerte mit einem Kartengitter bestimmt wird,

wobei das Kartengitter optional einen tatsächlichen Standortwert der Anzeigeeinheit umfasst, wobei der Schritt des Bestimmens des Standorts der Anzeigeeinheit ferner optional ein Vergleichen eines ungefähren Standortwerts umfasst, der proximal zu einem tatsächlichen Standortwert ist, der durch das Kartengitter umfasst wird, wobei der Schritt des Bestimmens des Standorts der Anzeigeeinheit ferner optional ein Vergleichen eines ungefähren Standortwerts zu einem tatsächlichen Standortwert innerhalb eines vorbestimmten Radius umfasst, wobei der vorbestimmte Radius wahlweise zwischen 1 bis 5 m ist oder

(b) der Rechner wahlweise ein Remote-Rechner ist, der eingerichtet ist, Signale von dem Patientenüberwachungsgerät durch den drahtlosen Sender zu empfangen,

wobei die Anzeigeeinheit optional eingerichtet ist, durch das Netzwerk Signale von dem entfernt angeordneten Rechner zu empfangen.

Revendications

1. Un procédé de surveillance d'un patient, comprenant les étapes suivantes :

(a) l'association d'un premier ensemble d'infor-

mations de signes vitaux au patient, le premier ensemble d'informations de signes vitaux étant mesuré à partir du patient avec un premier dispositif de surveillance comprenant un premier capteur de mouvement ;

(b) le stockage du premier ensemble d'informations de signes vitaux associé au patient dans une mémoire d'ordinateur ;

(c) la mise en contact du premier dispositif de surveillance avec un second dispositif de surveillance comprenant un second capteur de mouvement, la mise en contact amenant le premier capteur de mouvement à générer un premier signal de mouvement, et le second capteur de mouvement à générer un second signal de mouvement ;

(d) la transmission sans fil des premier et second signaux de mouvement à un serveur distant

(e) le traitement collectivement des premier et second signaux de mouvement sur le serveur distant pour déterminer que le premier dispositif de surveillance doit être remplacé par le second dispositif de surveillance ; et

(f) l'association d'un second ensemble d'informations de signes vitaux au patient, le second ensemble d'informations de signes vitaux étant mesuré à partir du patient avec le second dispositif de surveillance.

2. Un système pour mesurer un patient, comprenant :

un premier dispositif de surveillance de patient comprenant un premier capteur de mouvement et un premier émetteur sans fil ;

un second dispositif de surveillance de patient comprenant un second capteur de mouvement et un second émetteur sans fil ;

un serveur distant connecté au premier dispositif de surveillance de patient par l'intermédiaire du premier émetteur sans fil, et au second dispositif de surveillance de patient par l'intermédiaire du second émetteur sans fil, le serveur distant étant configuré pour recevoir un premier signal de mouvement généré par le premier capteur de mouvement et un second signal de mouvement généré par le second capteur de mouvement après que les premier et second dispositifs de surveillance de patient soient entrés en contact l'un avec l'autre ;

dans lequel le serveur distant est configuré pour associer un patient unique à un premier ensemble d'informations mesurées avec le premier dispositif de surveillance de patient et à un second ensemble d'informations mesurées avec le second dispositif de surveillance de patient.

3. Le procédé selon la revendication 1, dans lequel le premier capteur de mouvement comprend un pre-

mier accéléromètre, et le second capteur de mouvement comprend un second accéléromètre.

4. Le procédé selon la revendication 3, dans lequel le premier signal de mouvement est une première forme d'onde dépendante du temps, et le second signal de mouvement est une seconde forme d'onde dépendante du temps, dans lequel la première forme d'onde dépendant du temps comprend éventuellement une première impulsion, et la seconde forme d'onde dépendante du temps comprend une seconde impulsion, dans lequel l'étape de traitement comprend en outre le fait de déterminer que la première impulsion et la seconde impulsion sont générées pendant une période de temps commune ou dans lequel l'étape de traitement comprend en outre le fait de déterminer qu'une première amplitude de la première impulsion et une seconde amplitude de la seconde impulsion dépassent toutes deux une valeur seuil prédéterminée. 5 10 15 20
5. Le procédé selon la revendication 1, comprenant en outre l'étape de transmission sans fil des informations d'instruction du serveur distant à au moins l'un des premier dispositif de surveillance et second dispositif de surveillance indiquant que le premier dispositif de surveillance doit être remplacé par le second dispositif de surveillance ou dans lequel les premier et second ensembles d'informations de signes vitaux sont éventuellement stockés dans une mémoire d'ordinateur sur l'ordinateur distant, dans lequel la mémoire informatique de l'ordinateur distant est éventuellement une base de données. 25 30 35
6. Le procédé selon la revendication 1, dans lequel les premier et second ensembles d'informations de signes vitaux comprennent des valeurs pour au moins l'une des propriétés suivantes : la pression artérielle, la fréquence cardiaque, la fréquence respiratoire, le SpO2, la température. 40
7. Le procédé selon la revendication 1, dans lequel les premier et second ensembles d'informations de signes vitaux comprennent au moins l'une des formes d'onde dépendant du temps suivantes : PPG, ECG, OSC, IP. 45
8. Le procédé selon la revendication 1, comprenant en outre l'étape consistant à associer des informations démographiques de patient décrivant le patient à la fois aux premier et second ensembles d'informations de signes vitaux, dans lequel les informations démographiques du patient comprennent au moins un des paramètres suivants : le nom, le sexe, le poids, la taille, la date de naissance. 50 55

9. Un procédé de surveillance d'un patient, comprenant les étapes suivantes :

- (a) la mise en contact d'au moins l'un d'un dispositif d'affichage et d'une zone à proximité du dispositif d'affichage avec un dispositif de surveillance de patient comprenant un capteur de mouvement et un émetteur sans fil, la mise en contact amenant le capteur de mouvement à générer un signal de mouvement ;
- (b) la transmission sans fil du signal de mouvement à un ordinateur distant sur un réseau sans fil à l'aide de l'émetteur sans fil ;
- (c) le traitement du signal de mouvement par l'ordinateur pour associer le dispositif de surveillance de patient au dispositif d'affichage ;
- (d) la mesure d'au moins un ensemble d'informations de signes vitaux du patient avec le dispositif de surveillance de patient ; et
- (e) l'affichage du ou des ensembles d'informations de signes vitaux sur le dispositif d'affichage.

10. Le procédé selon la revendication 9, dans lequel des signaux sans fil reçus par de multiples points d'accès sans fil sur le réseau sans fil sont traités pour déterminer un emplacement approximatif d'au moins l'un du dispositif de surveillance de patient et du dispositif d'affichage, et le signal de mouvement et l'emplacement approximatif d'au moins l'un du dispositif de surveillance de patient et du dispositif d'affichage sont traités pour associer le dispositif de surveillance de patient au dispositif d'affichage. 25 30 35

11. Un système pour mesurer un patient, comprenant :

un dispositif de surveillance de patient comprenant un capteur de mouvement et un émetteur sans fil ; un dispositif d'affichage ; un serveur distant connecté au dispositif de surveillance de patient par l'intermédiaire de l'émetteur sans fil et au dispositif d'affichage par l'intermédiaire d'un réseau ou un ordinateur connecté au dispositif de surveillance de patient et au dispositif d'affichage par l'intermédiaire d'un réseau, l'ordinateur, le serveur distant et l'ordinateur, respectivement, étant configurés pour recevoir un signal de mouvement généré par le capteur de mouvement après que le dispositif de surveillance de patient soit entré en contact avec au moins l'un du dispositif d'affichage et d'une zone proximale au dispositif d'affichage ; dans lequel le serveur distant est configuré pour associer le dispositif de surveillance de patient au dispositif d'affichage de telle sorte que le dispositif d'affichage reçoit des informations de signes vitaux mesurées par le dispositif de sur-

- veillance de patient et transmises par l'intermédiaire du réseau, l'algorithme comprenant collectivement l'analyse du signal de mouvement et d'un emplacement du dispositif d'affichage.
- 5
12. Le procédé selon la revendication 9, dans lequel le capteur de mouvement comprend un accéléromètre.
13. Le procédé selon la revendication 9, dans lequel le signal de mouvement est une forme d'onde dépendant du temps, dans lequel la forme d'onde dépendant du temps comprend éventuellement une impulsion.
- 10
14. Le procédé selon la revendication 9, dans lequel l'étape de traitement comprend en outre le fait de traiter le signal de mouvement et un emplacement du dispositif d'affichage pour associer le dispositif de surveillance de patient au dispositif d'affichage.
- 15
15. Le procédé selon la revendication 14, comprenant en outre une étape de détermination de l'emplacement du dispositif d'affichage,
- 20
- dans lequel le dispositif de surveillance de patient comprend éventuellement en outre un émetteur sans fil,
- 25
- dans lequel l'émetteur sans fil est éventuellement configuré pour fonctionner sur un réseau sans fil,
- dans lequel
- 30
- (a) l'étape de détermination de l'emplacement du dispositif d'affichage comprend éventuellement en outre l'analyse de signaux caractérisant une intensité de signal entre de multiples points d'accès fonctionnant sur le réseau sans fil et l'émetteur sans fil,
- 35
- dans lequel l'étape de détermination de l'emplacement du dispositif d'affichage comprend éventuellement en outre la triangulation d'au moins trois valeurs d'intensité du signal, avec chaque valeur d'intensité du signal caractérisant un signal entre un point d'accès individuel fonctionnant sur le réseau sans fil et l'émetteur sans fil,
- 40
- dans lequel l'étape de détermination de l'emplacement du dispositif d'affichage comprend éventuellement en outre la comparaison d'une valeur d'emplacement approximative déterminée par triangulation des au moins trois valeurs d'intensité du signal avec une grille cartographique,
- 45
- dans lequel la grille cartographique comprend éventuellement une valeur d'emplacement réelle du dispositif d'affichage,
- 50
- dans lequel l'étape de détermination de l'emplacement du dispositif d'affichage comprend éventuellement en outre la comparaison d'une valeur d'emplacement approximative qui est
- 55

proche d'une valeur d'emplacement réelle comprise dans la grille cartographique, dans lequel l'étape de détermination de l'emplacement du dispositif d'affichage comprend éventuellement en outre la comparaison d'une valeur d'emplacement approximative à une valeur d'emplacement réelle dans un rayon prédéterminé,

dans lequel le rayon prédéterminé est éventuellement compris entre 1 et 5 m ou

(b) l'ordinateur est éventuellement un ordinateur distant configuré pour recevoir des signaux provenant du dispositif de surveillance de patient par l'intermédiaire de l'émetteur sans fil,

dans lequel le dispositif d'affichage est éventuellement configuré pour recevoir des signaux provenant de l'ordinateur distant par l'intermédiaire du réseau.

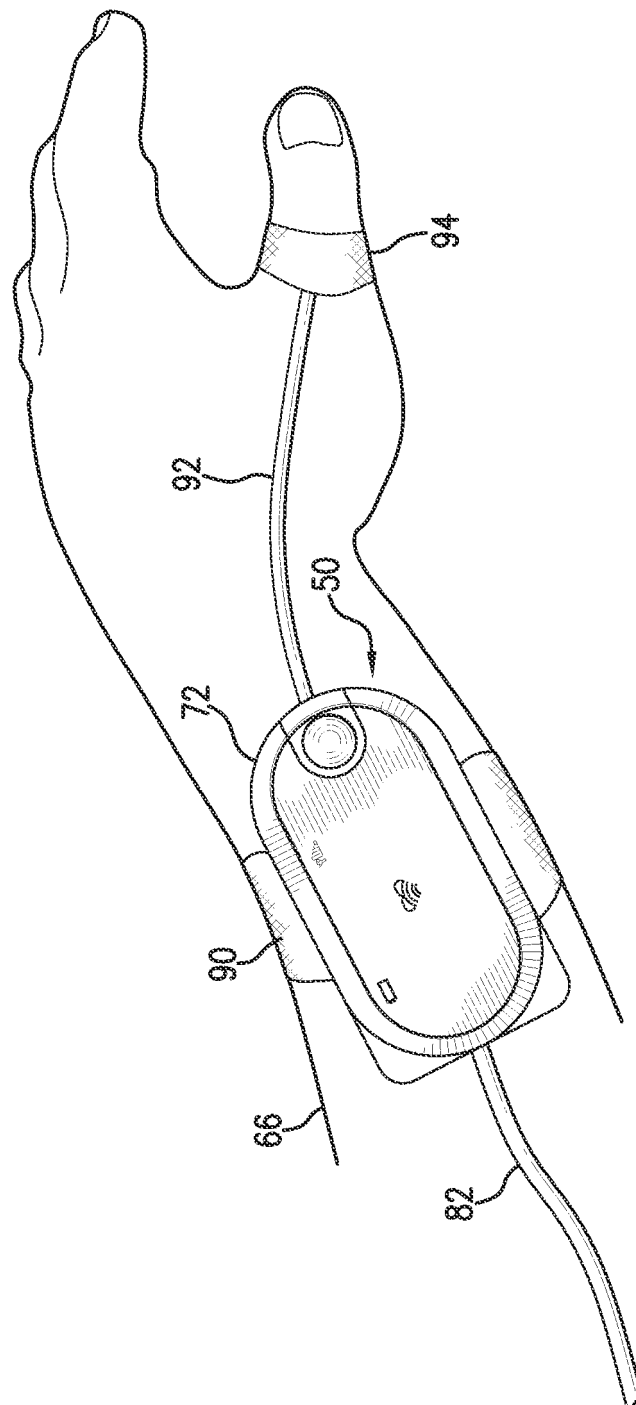


FIG.1

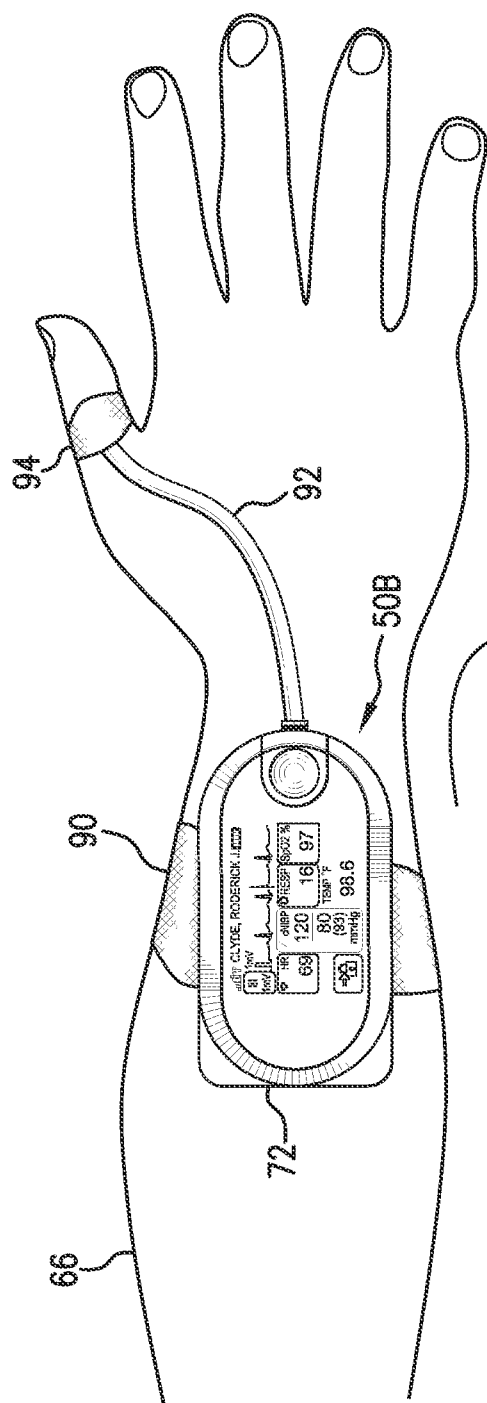
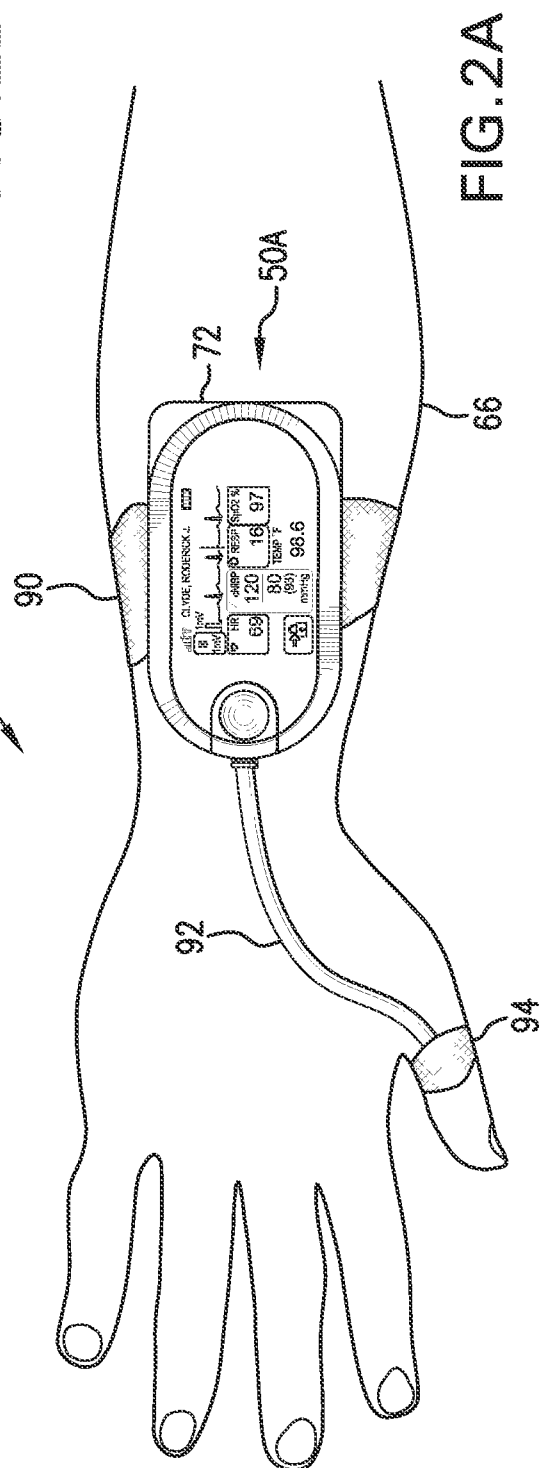
B
2
G
L

FIG. 2A

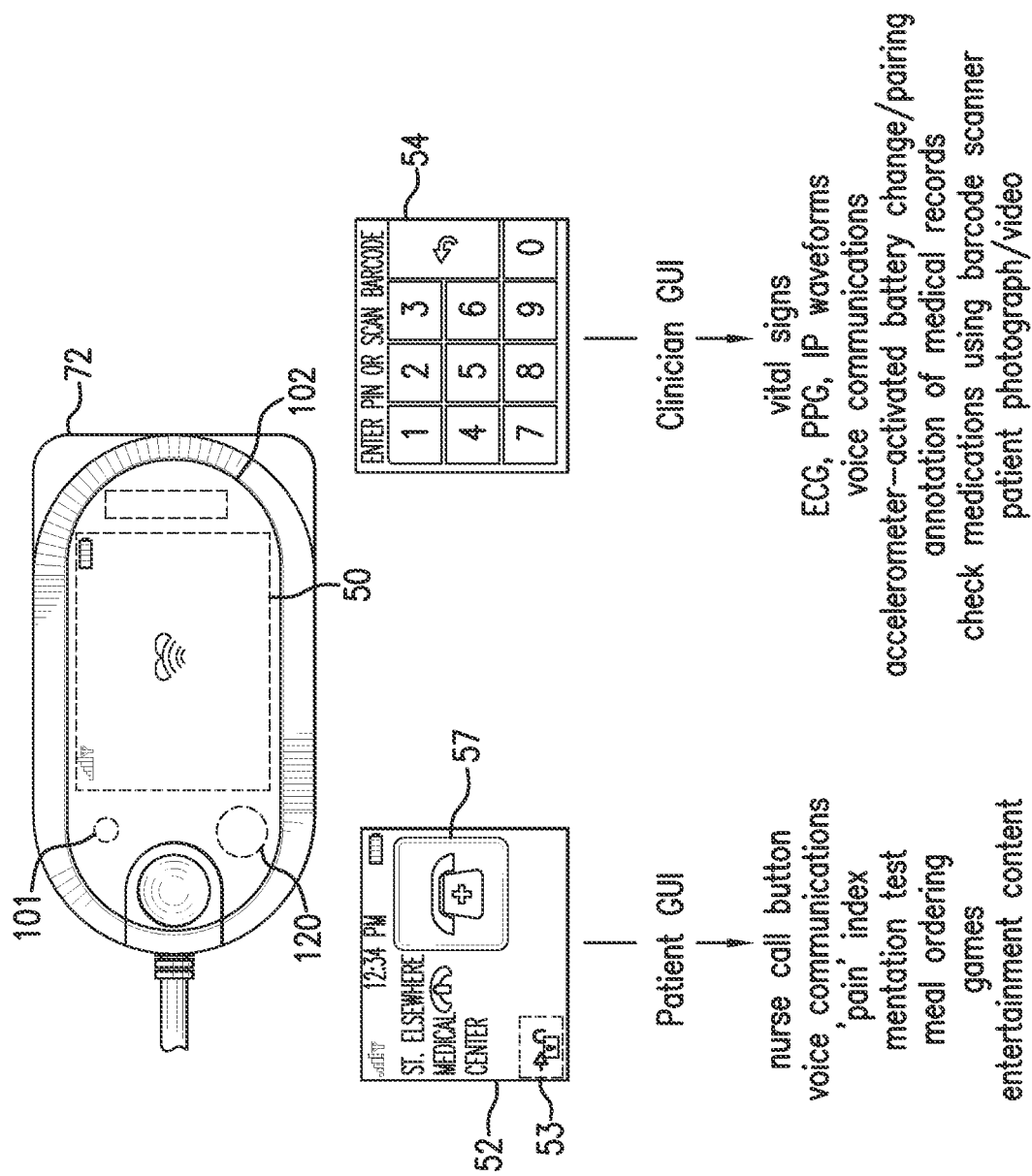


FIG.3

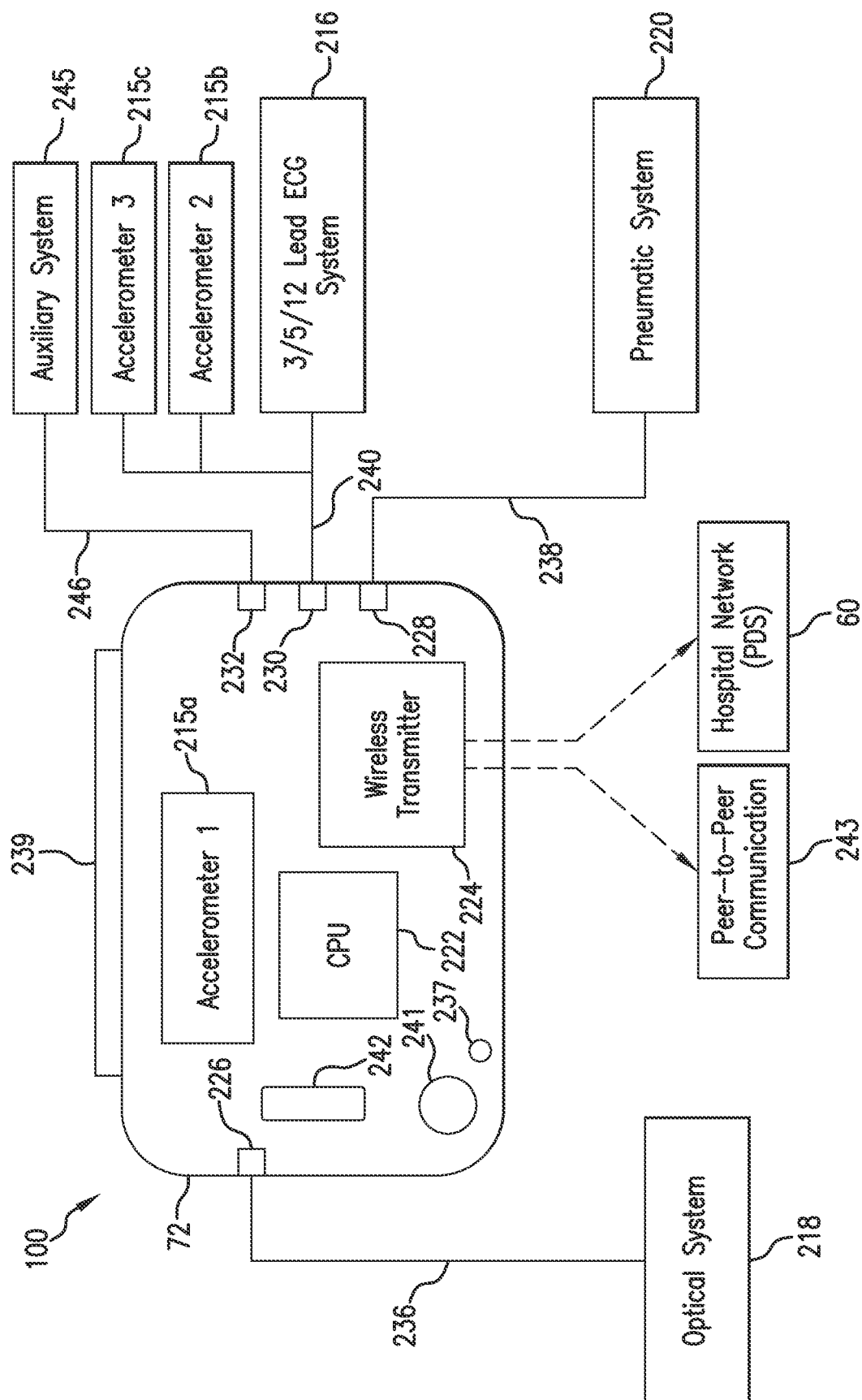


FIG. 4

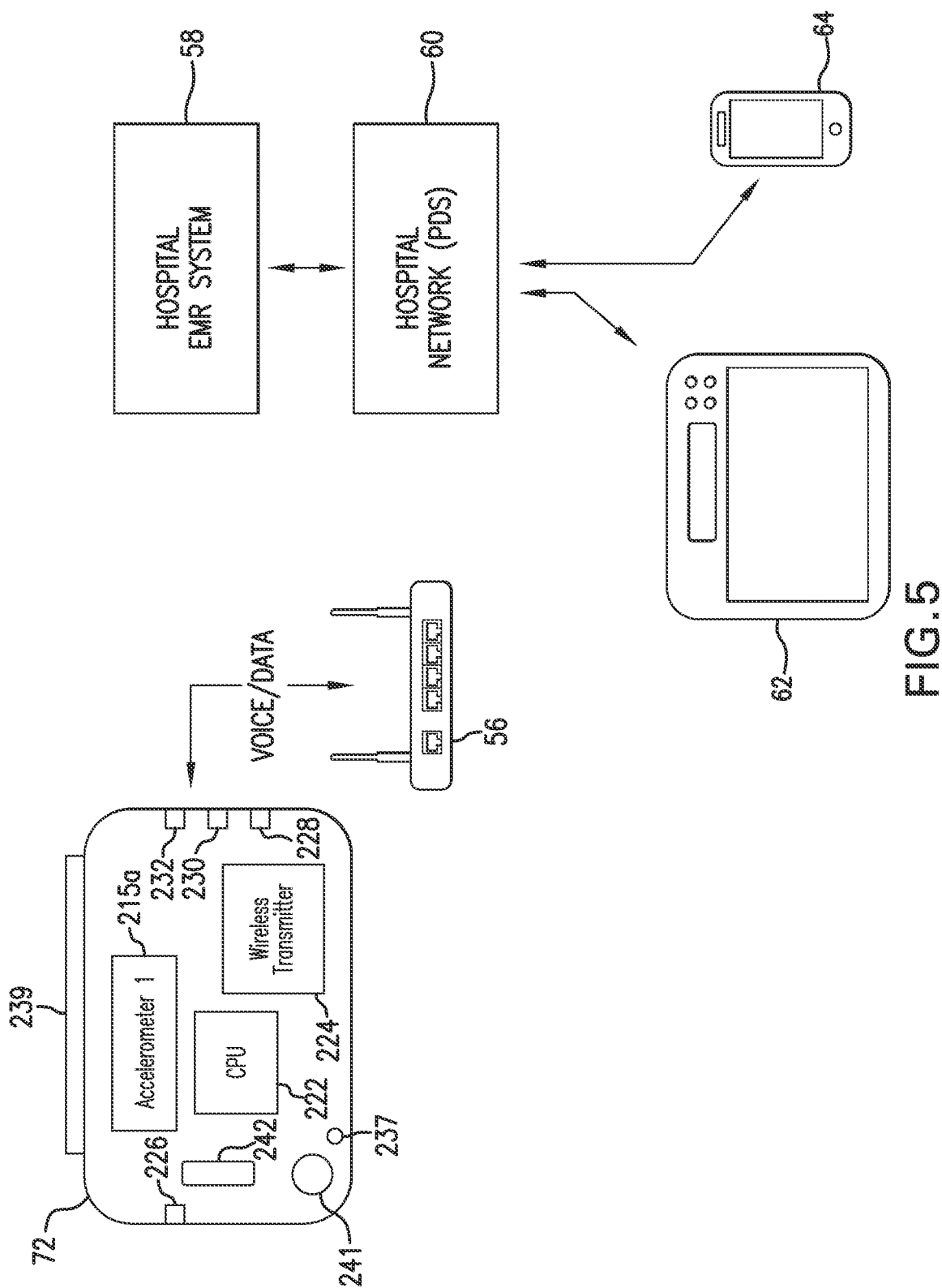
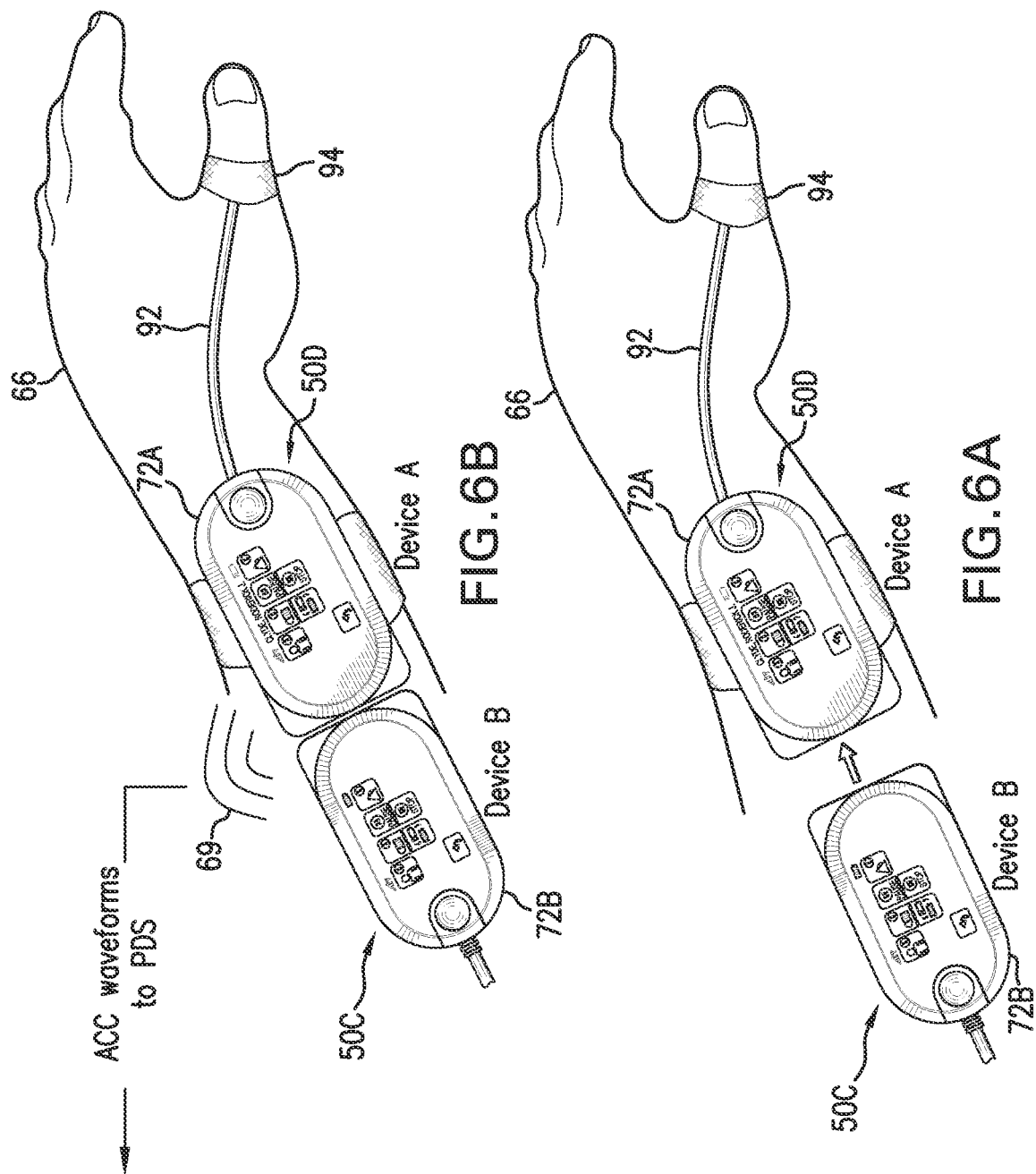
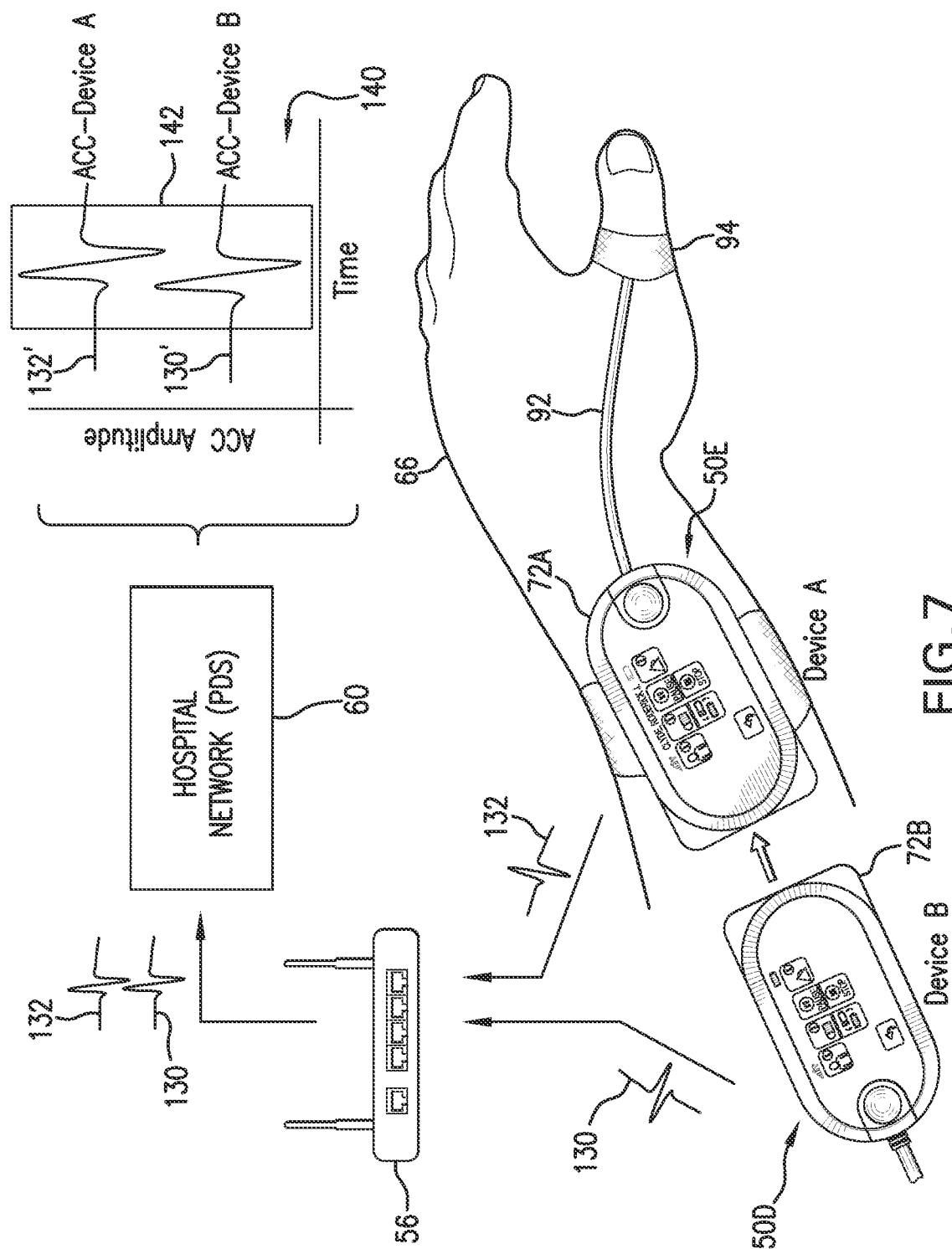
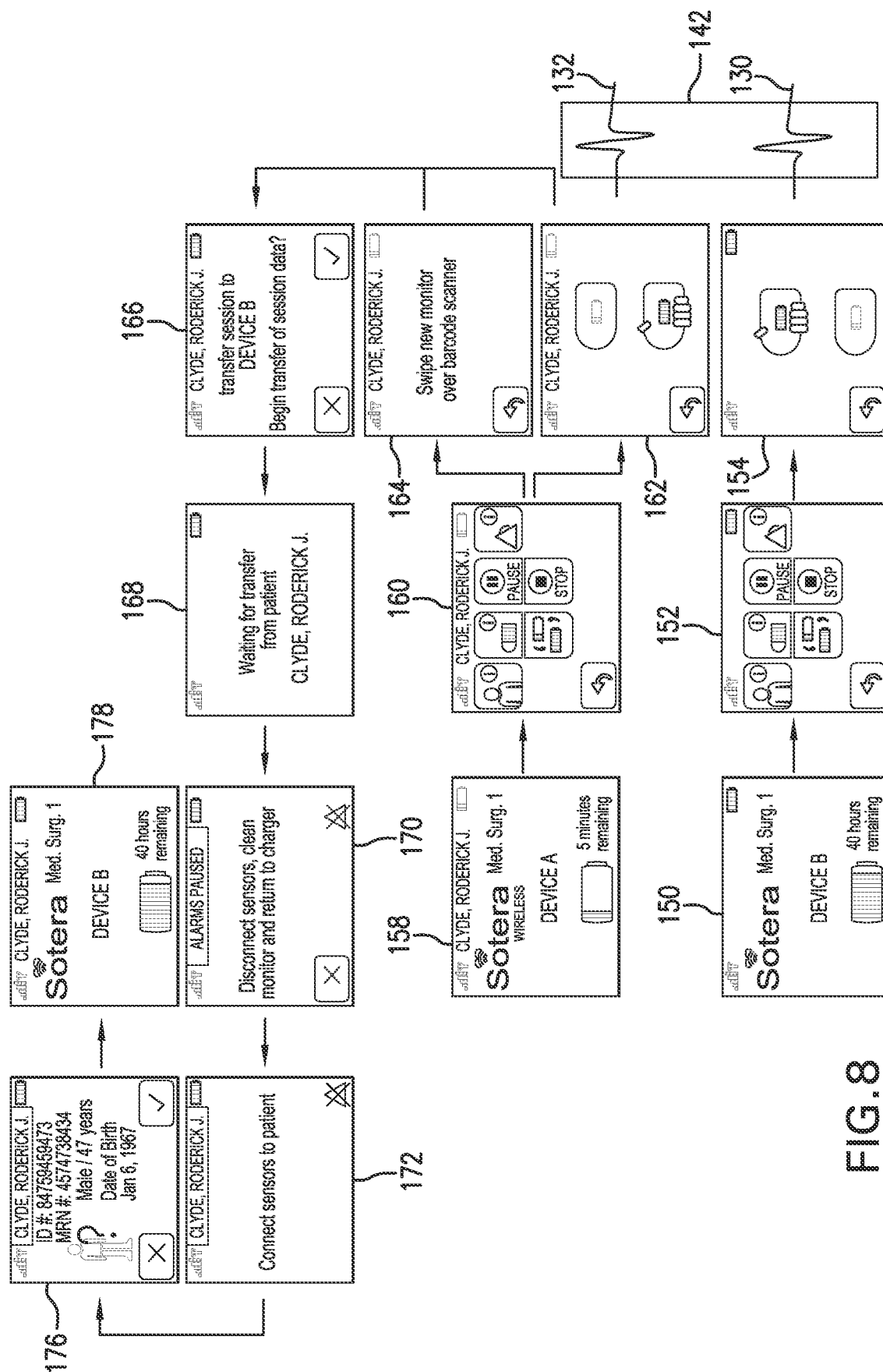


FIG. 5





7
R
G
L



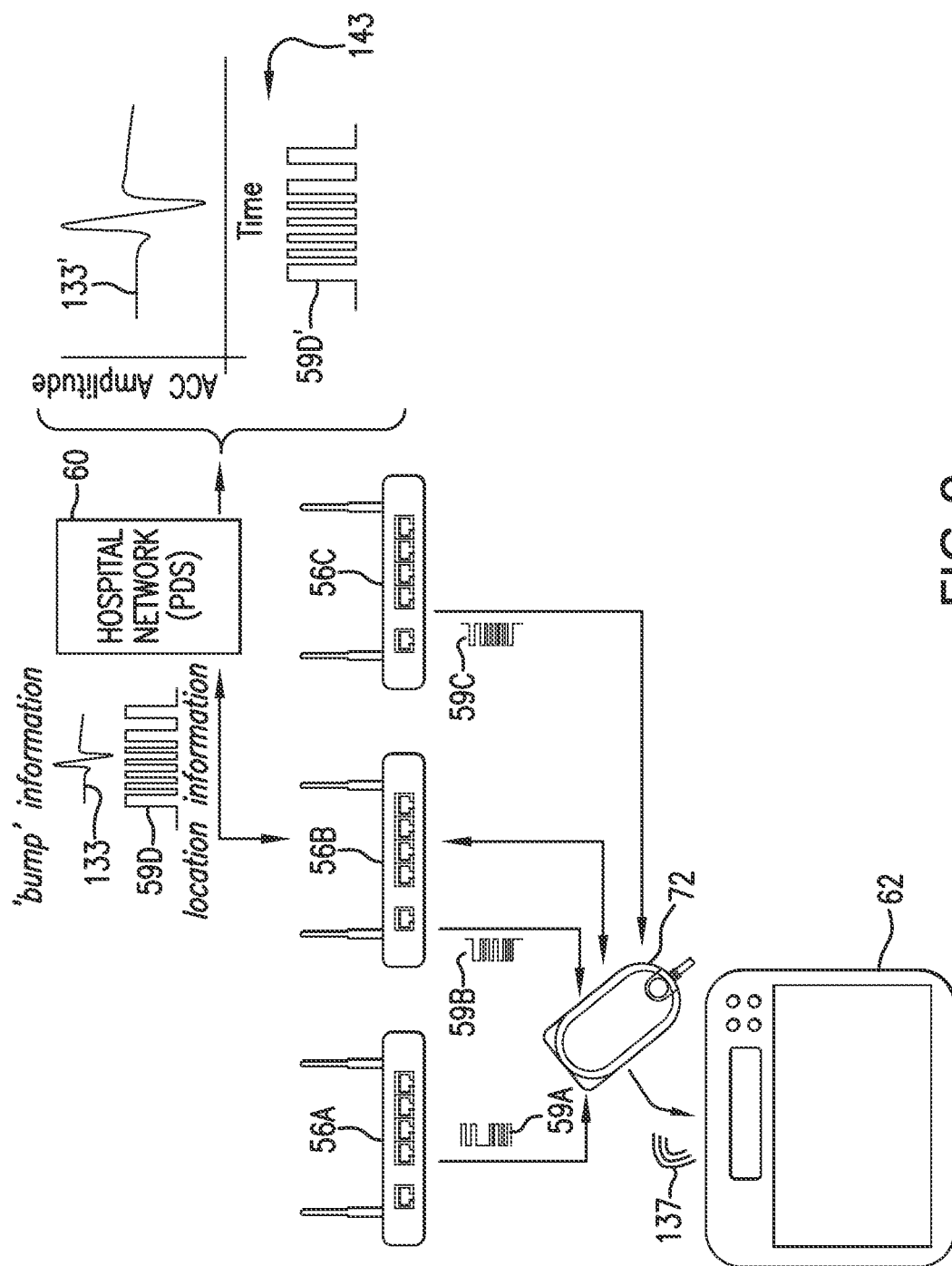


FIG. 9

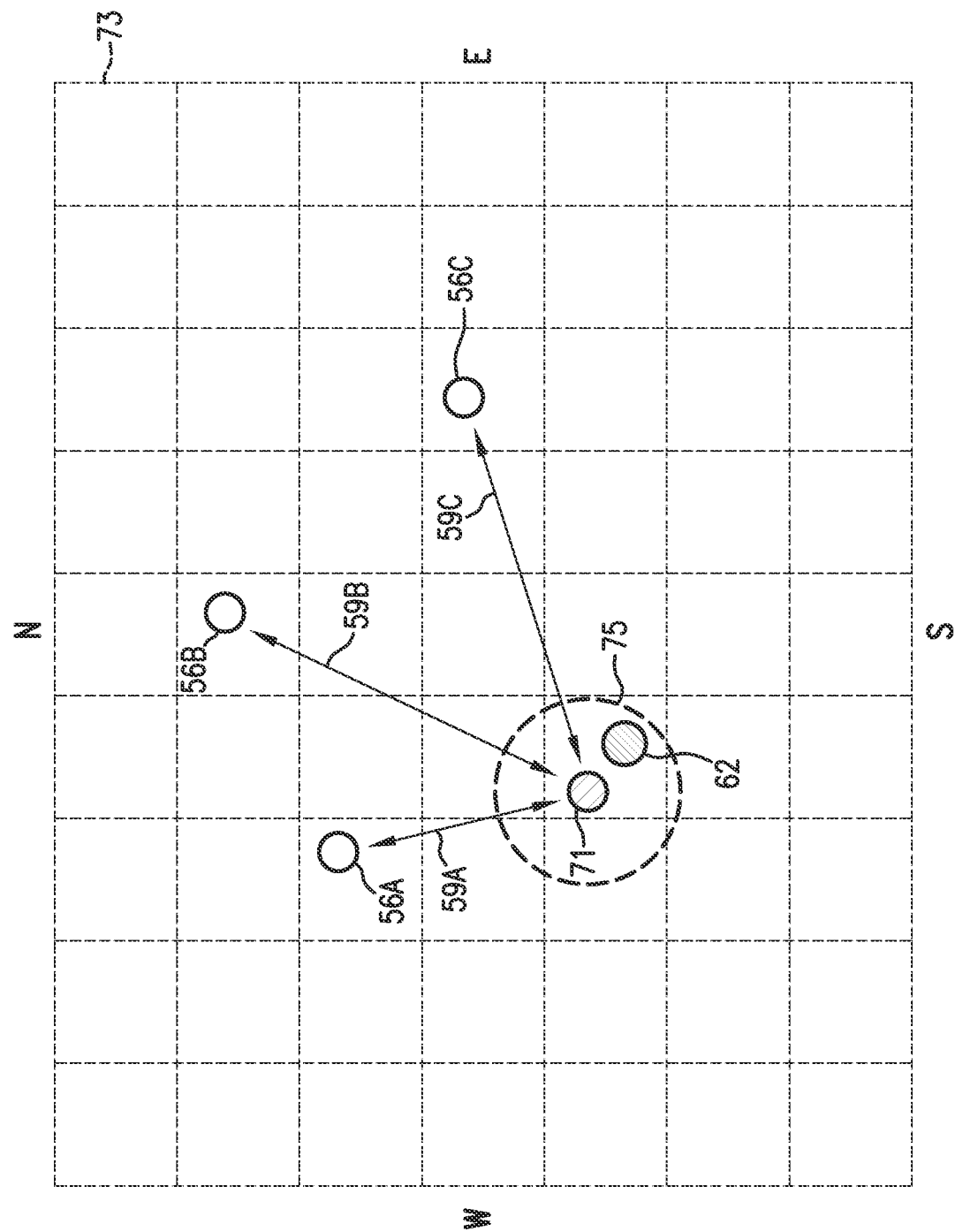
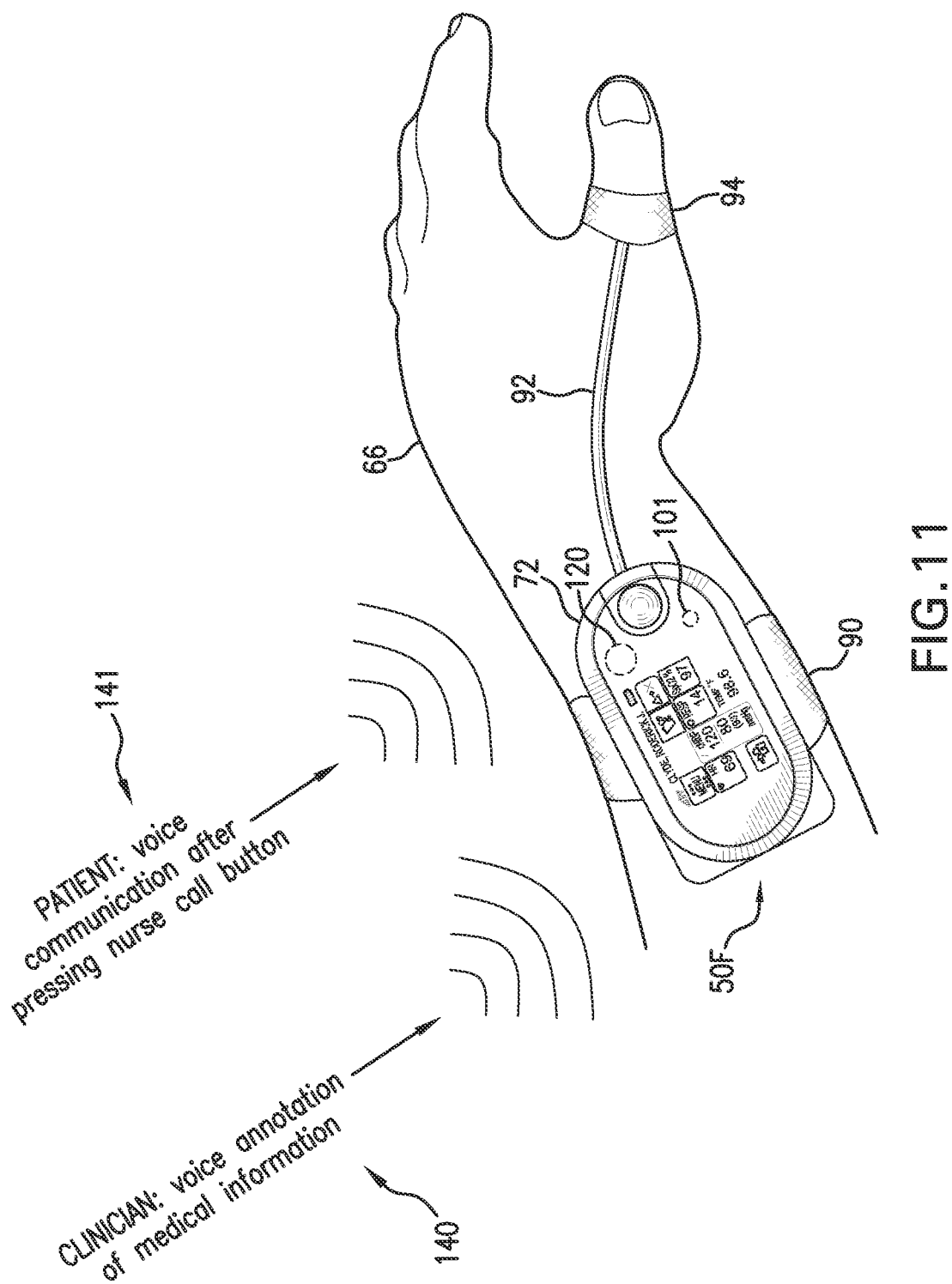
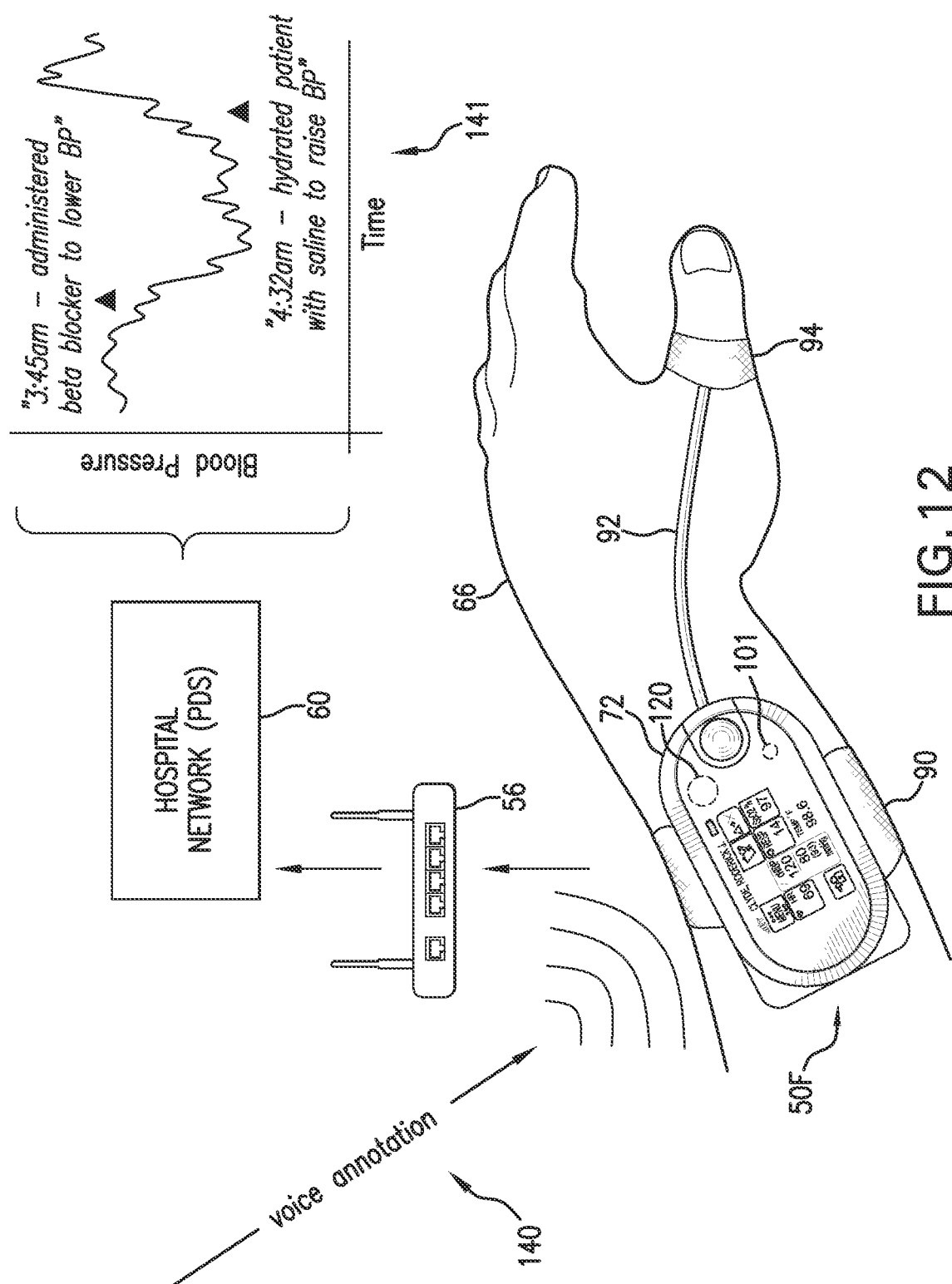


FIG. 10





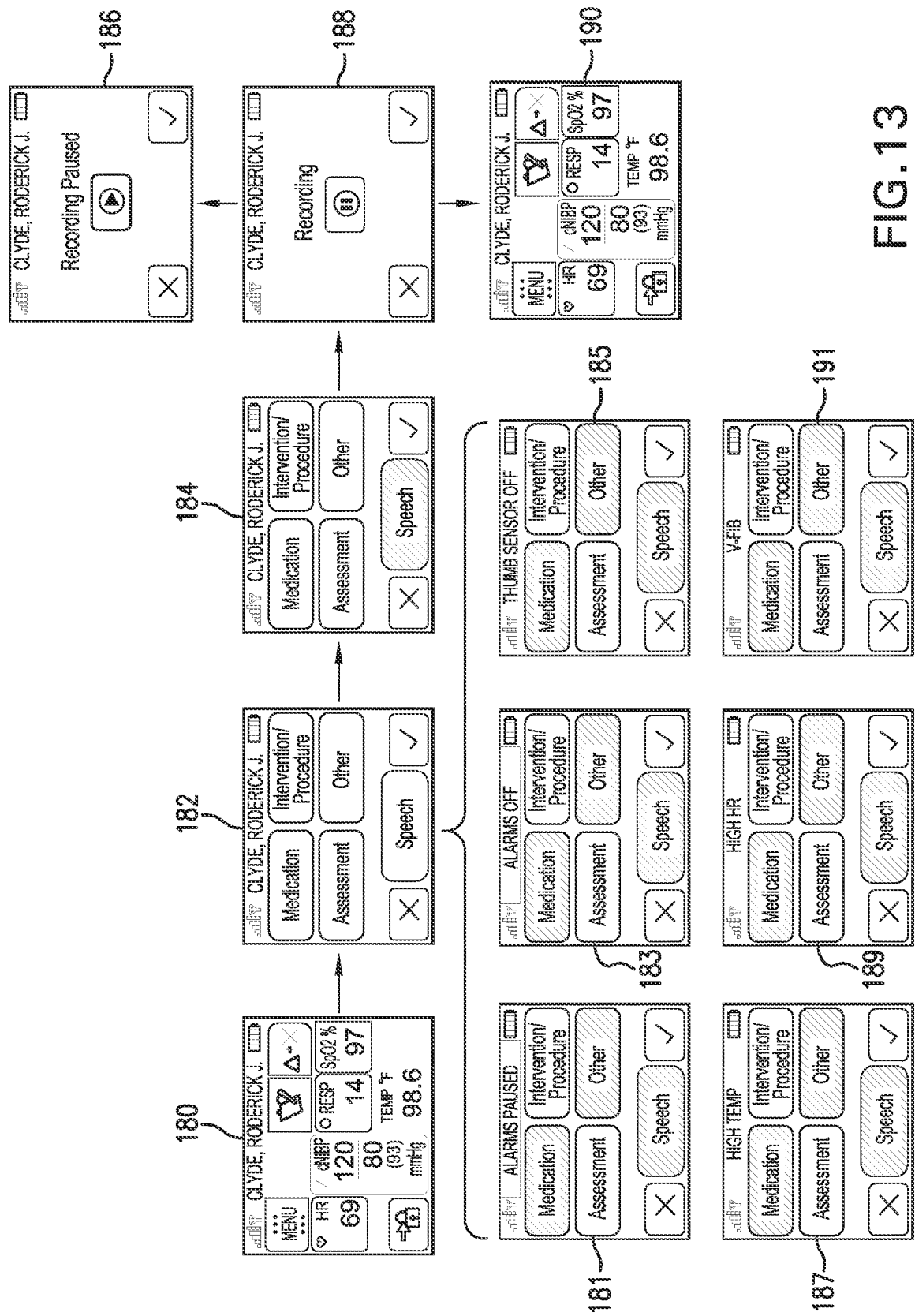


FIG. 13

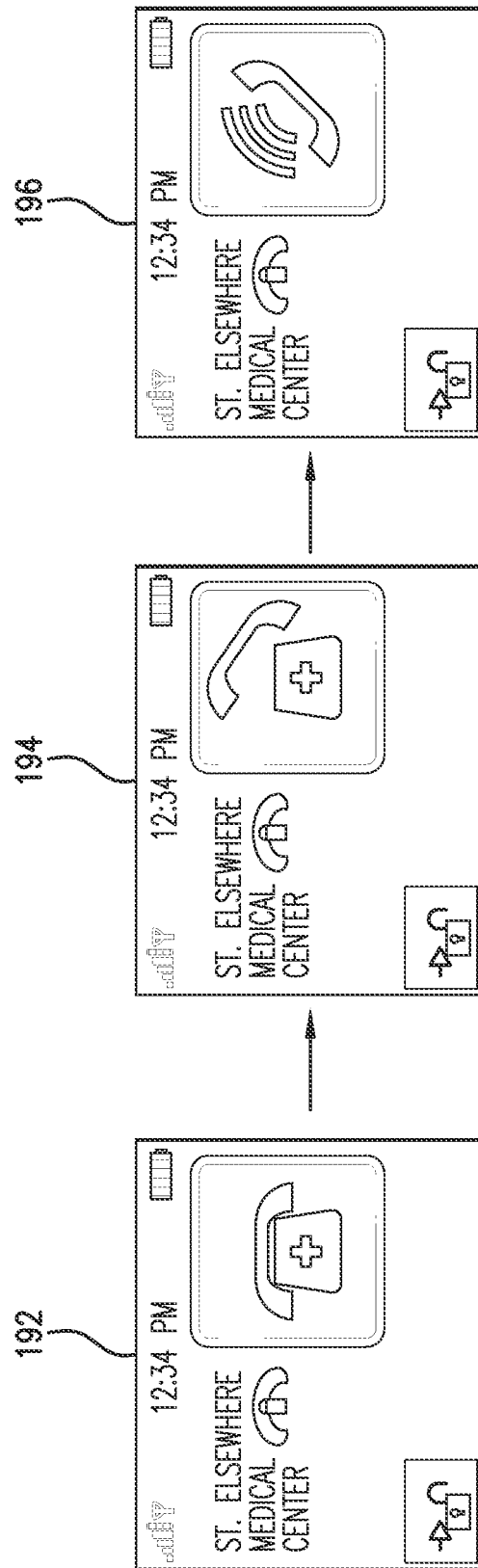


FIG.14

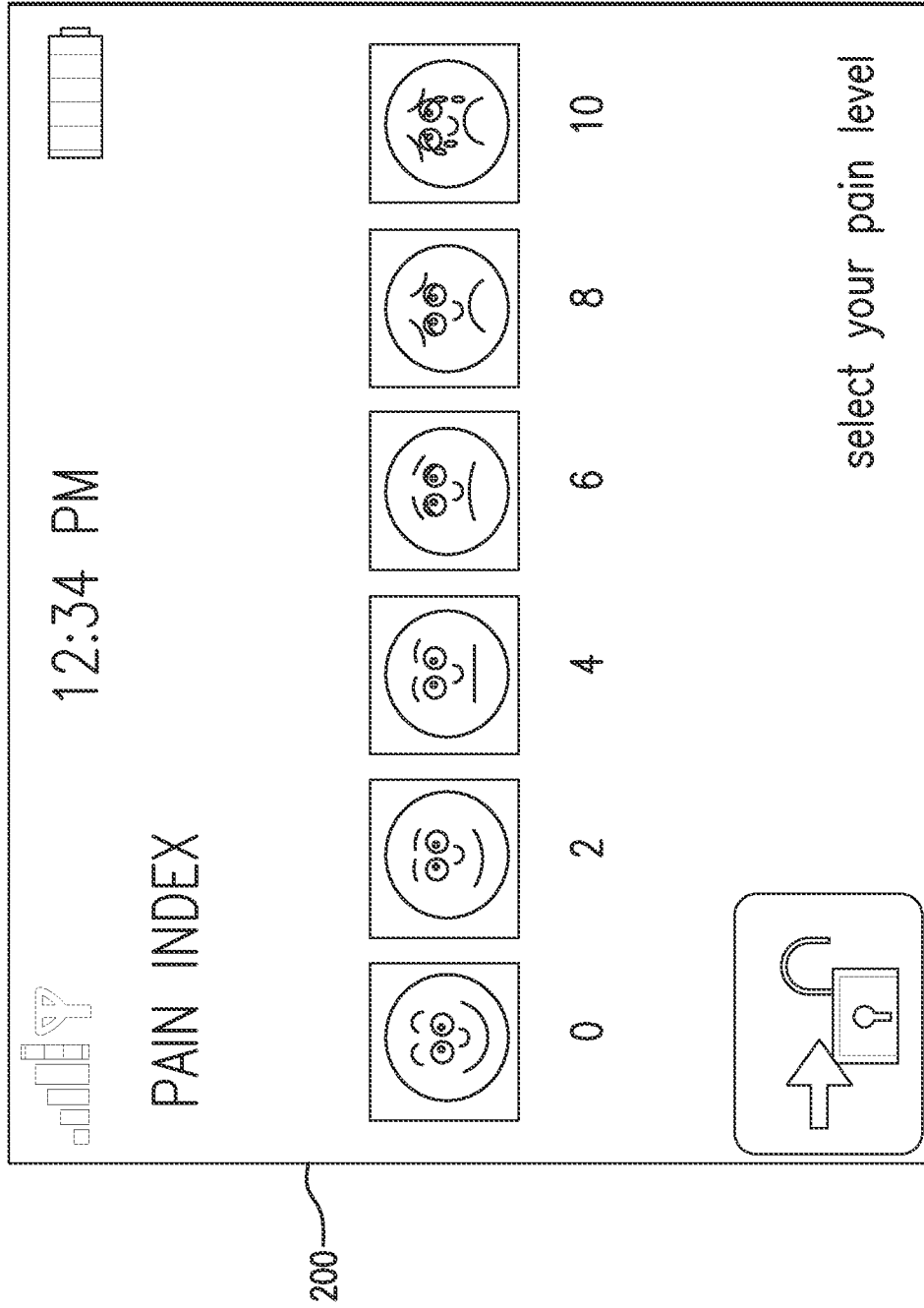


FIG.15

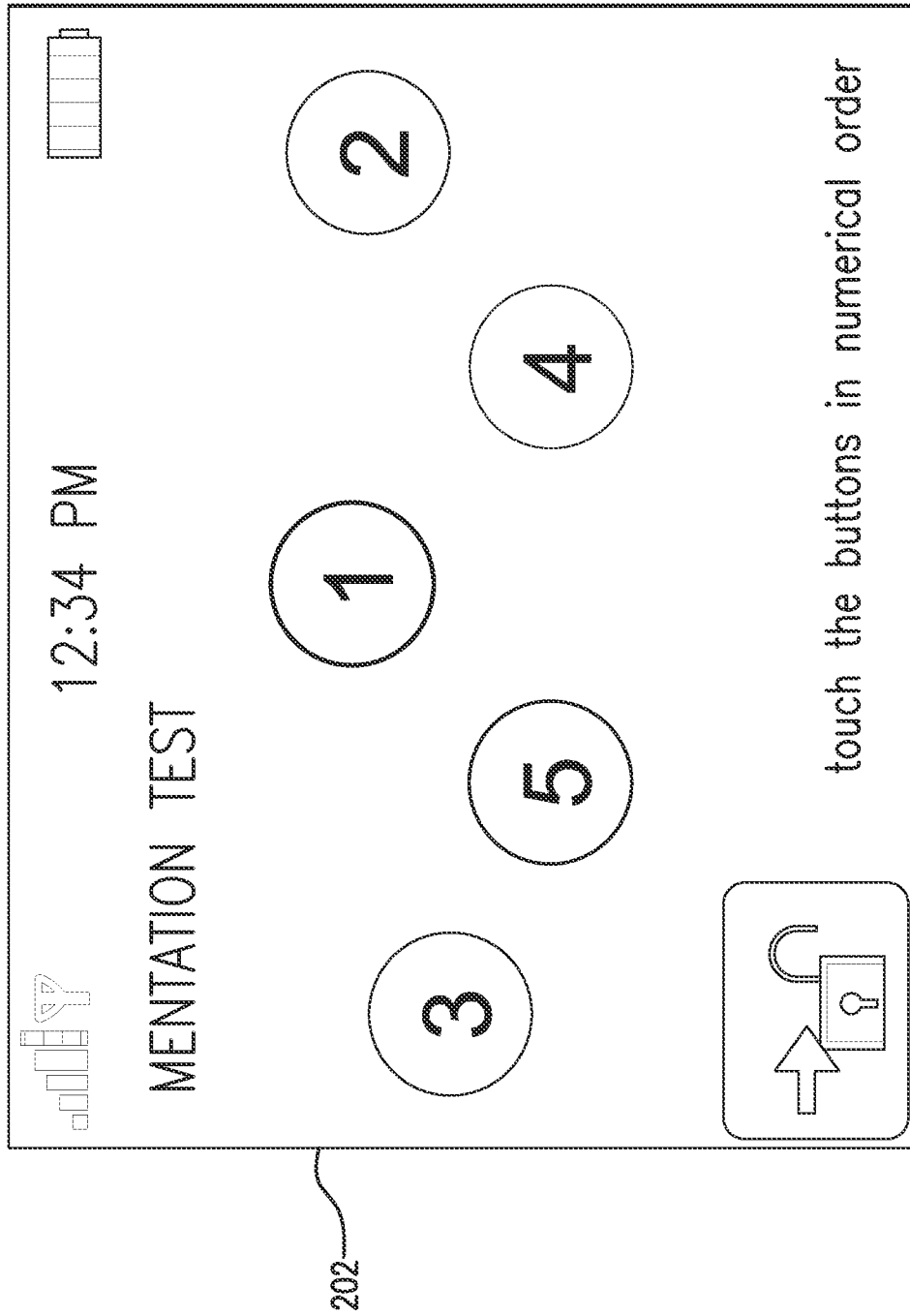


FIG. 16



FIG.17

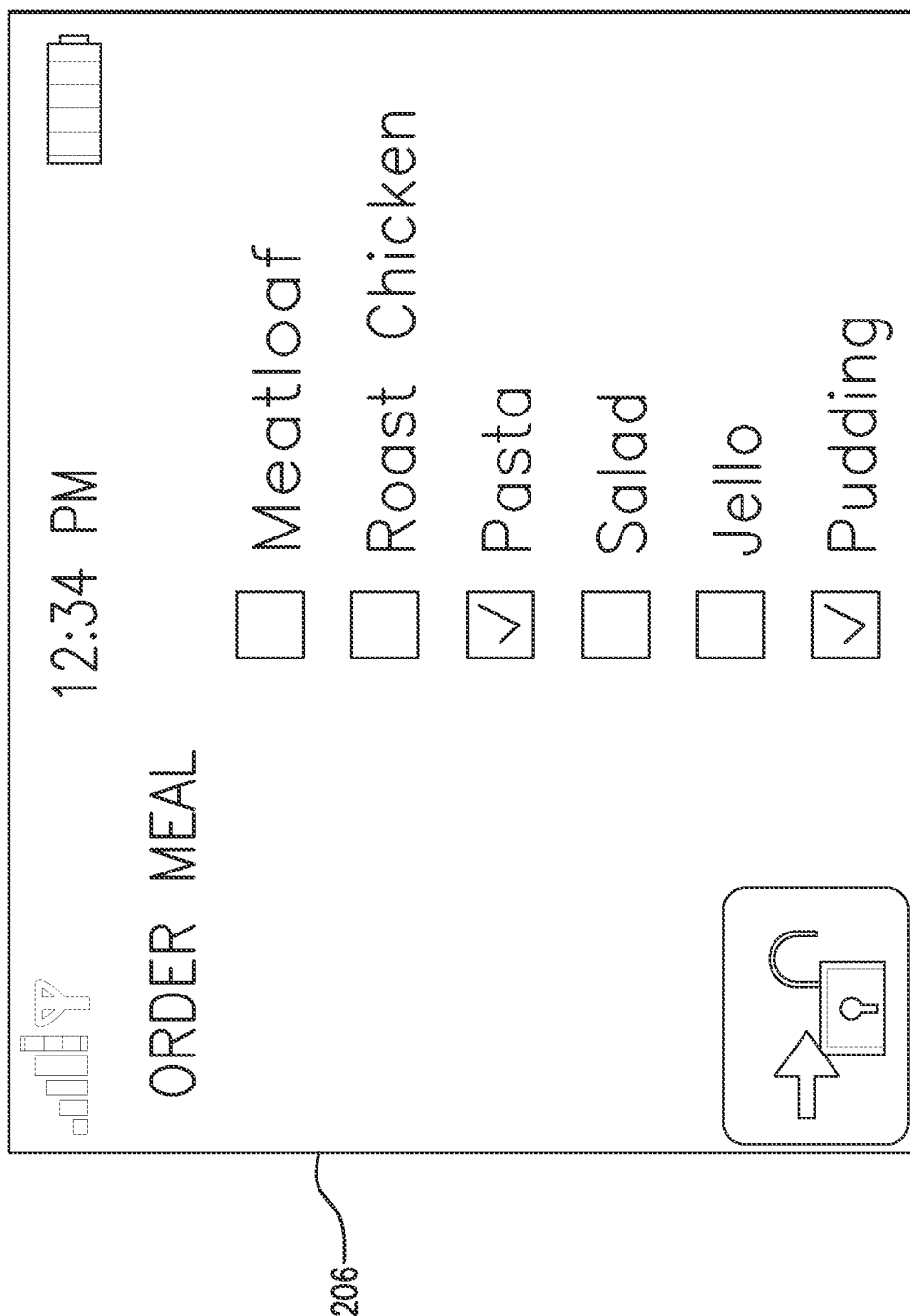


FIG.18

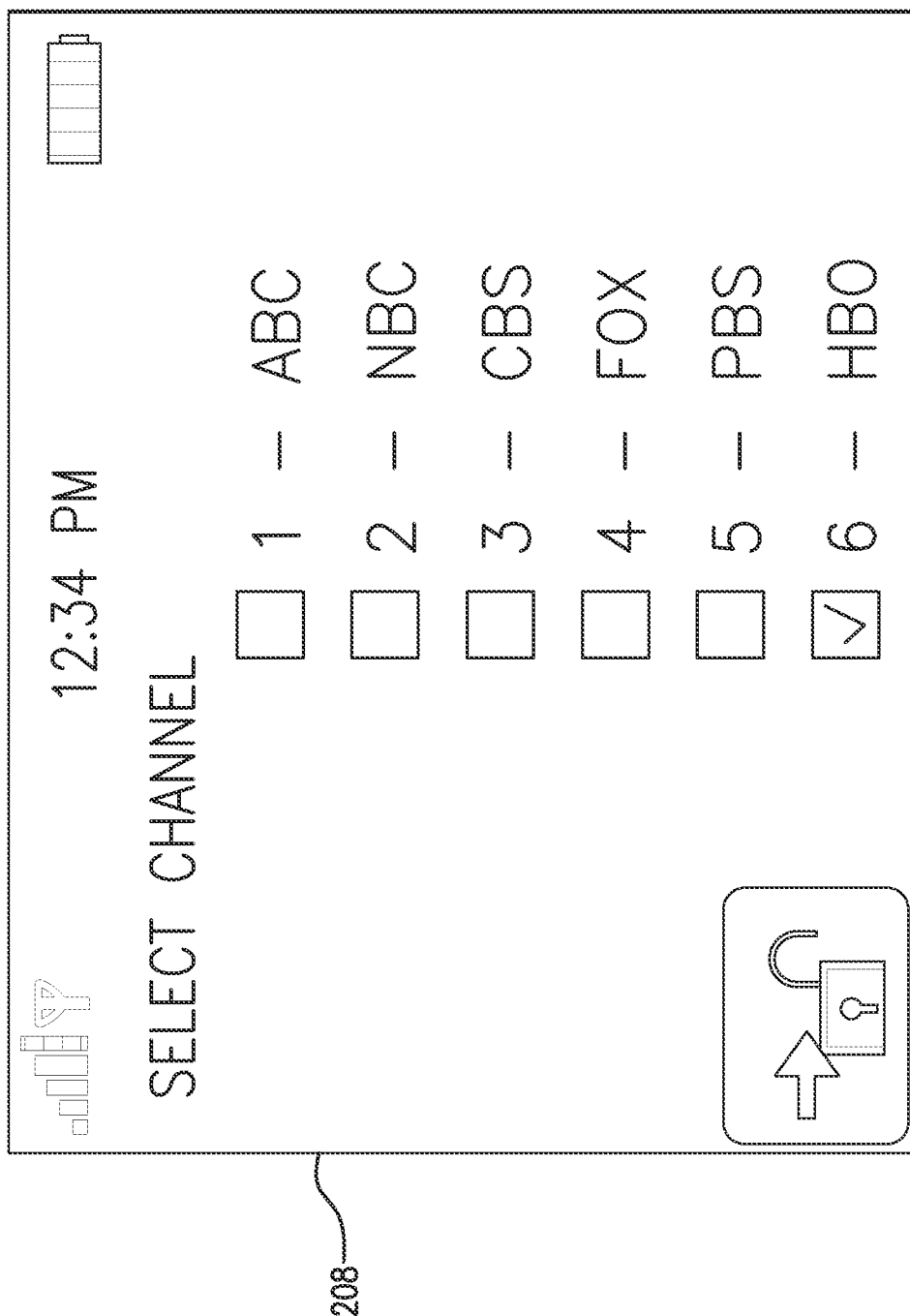


FIG.19

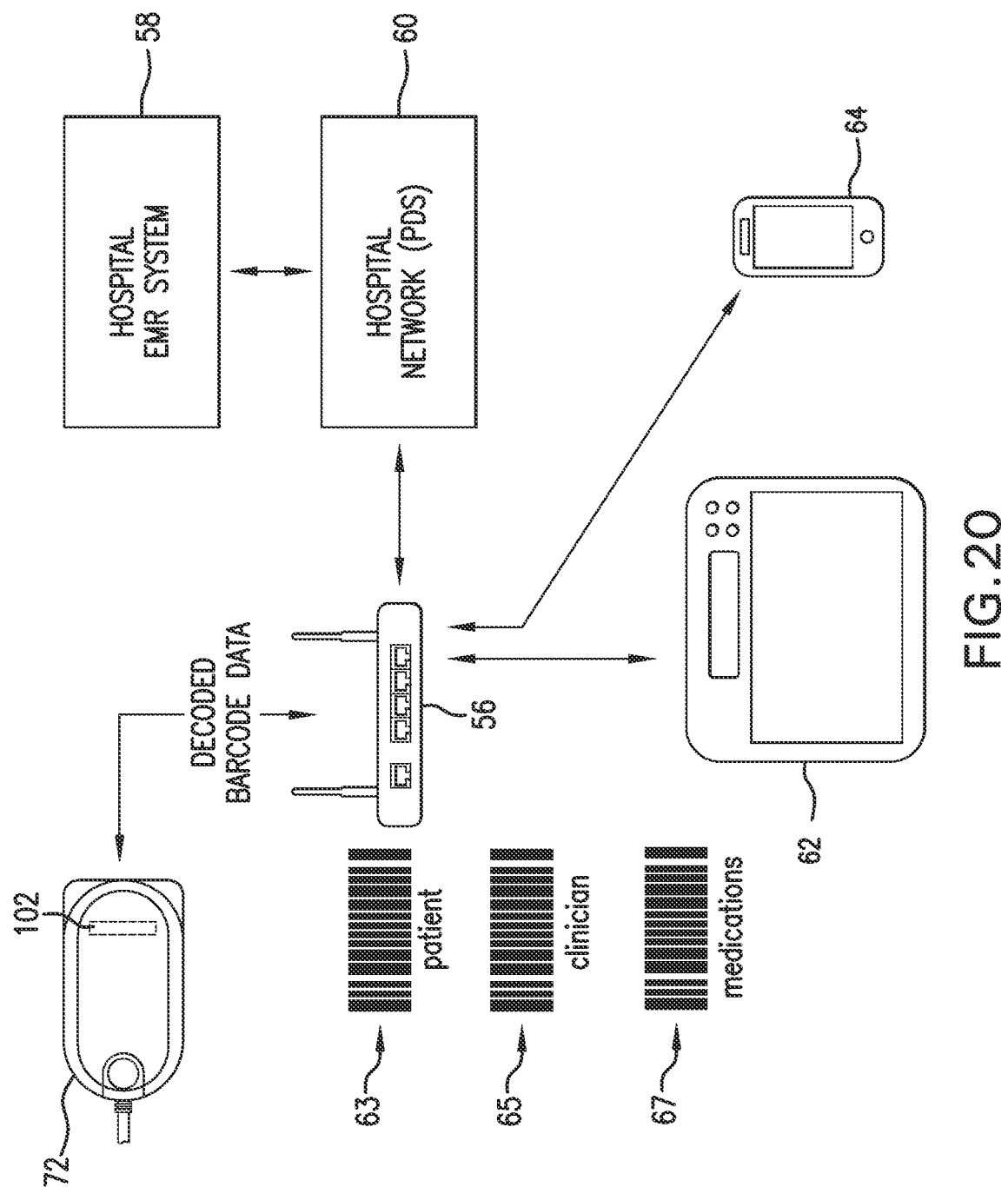


FIG. 20

Configuration for Indexing cNIBP Measurement (first ~60 seconds) Configuration for cNIBP/RR Measurements (remaining 3 hours, 59 minutes)

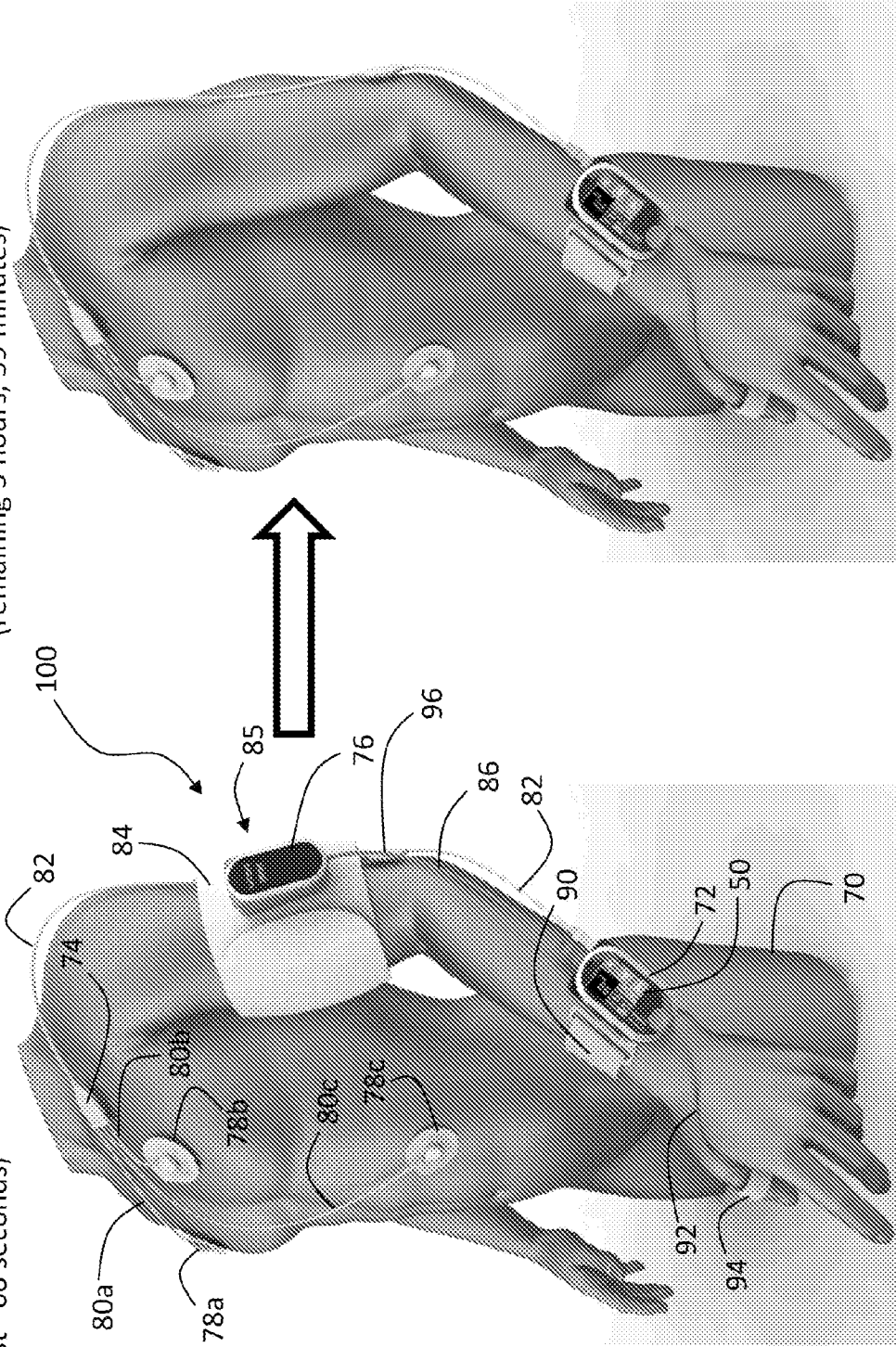
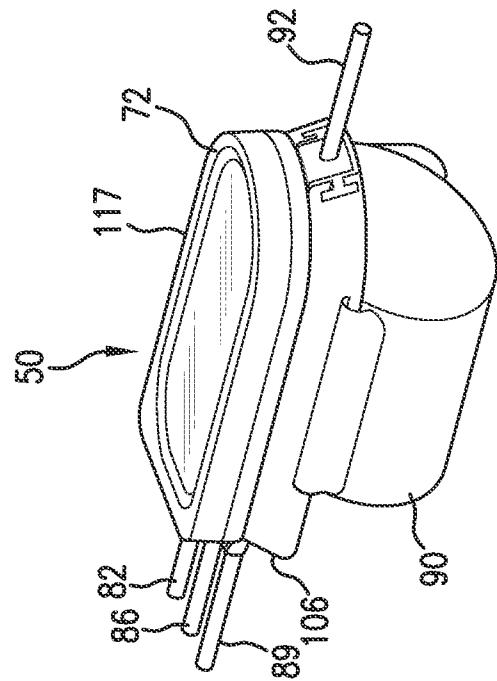
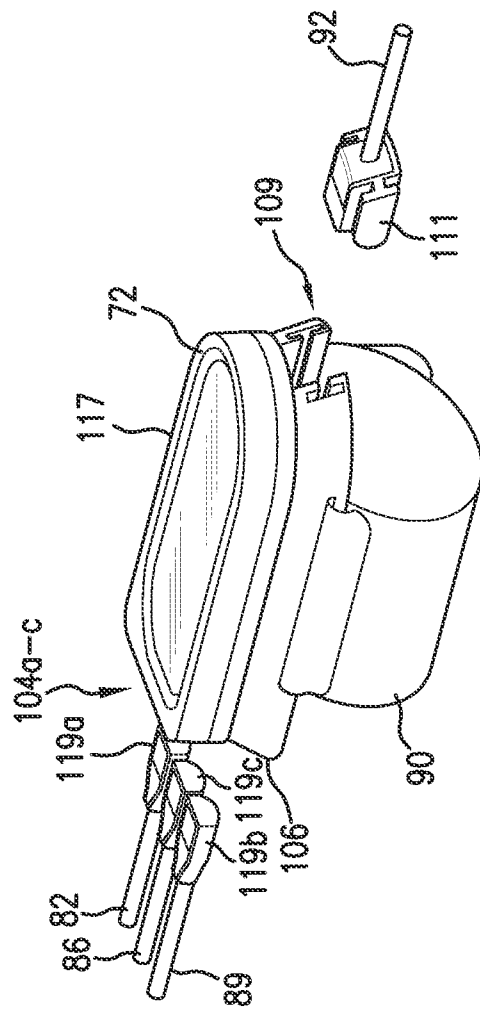


FIG. 21B

FIG. 21A



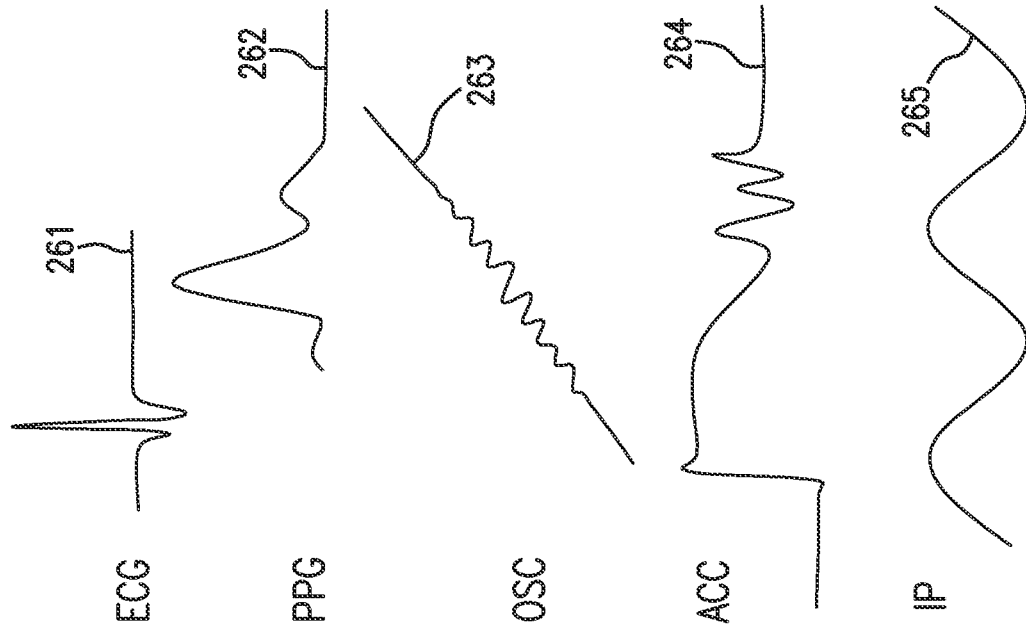


FIG. 23B

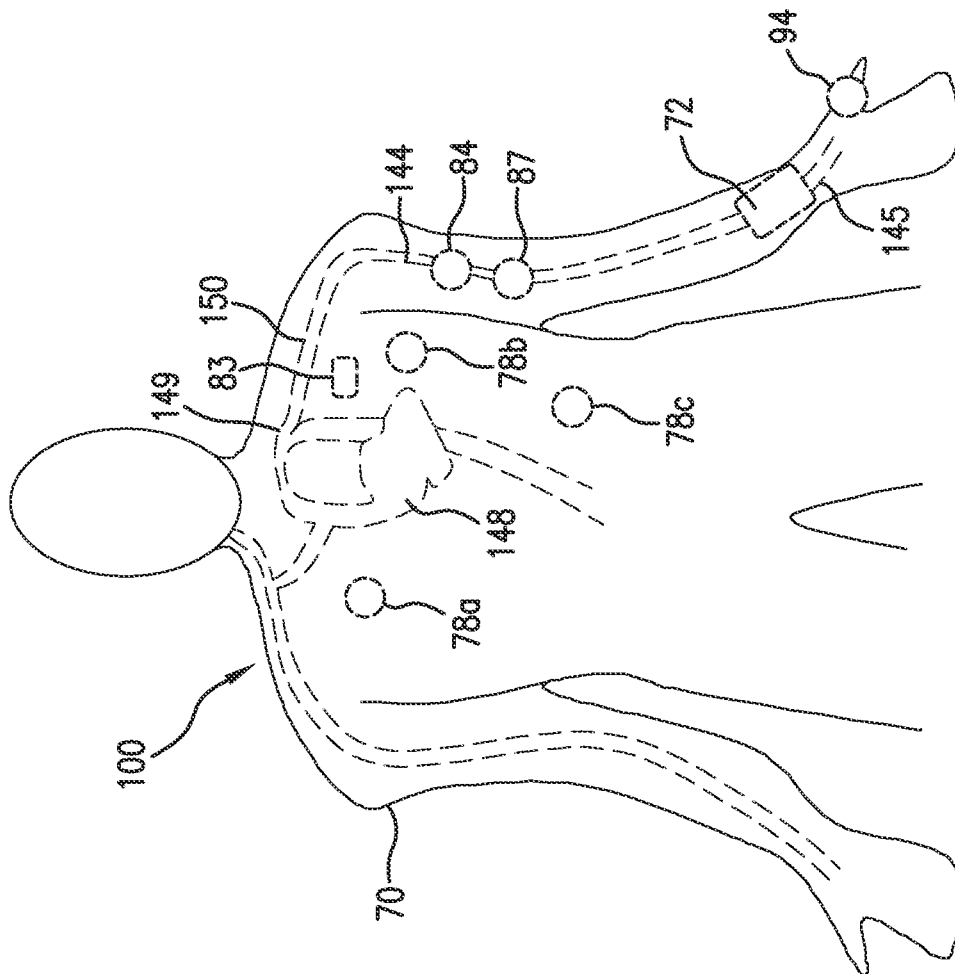


FIG. 23A

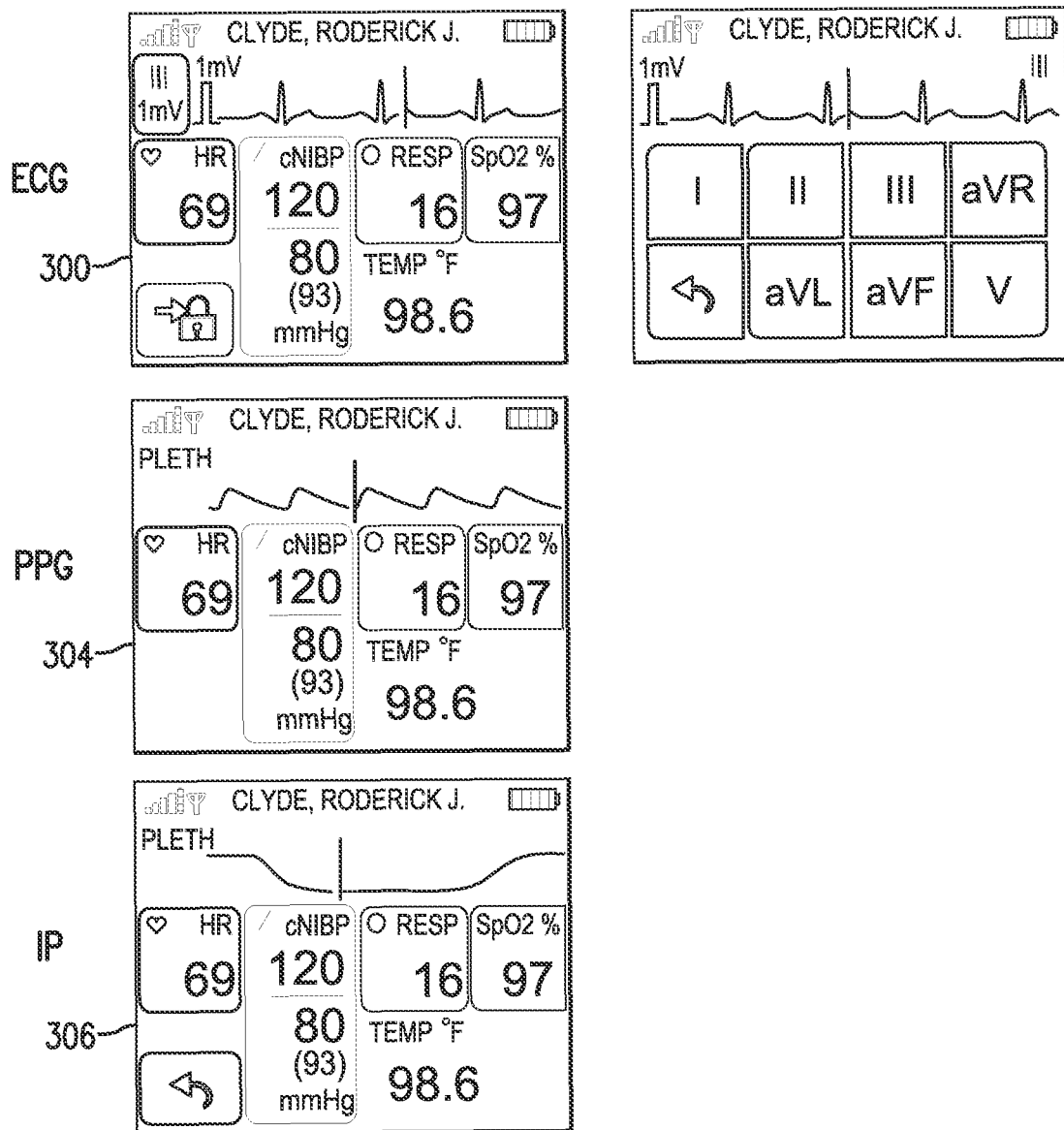


FIG.24

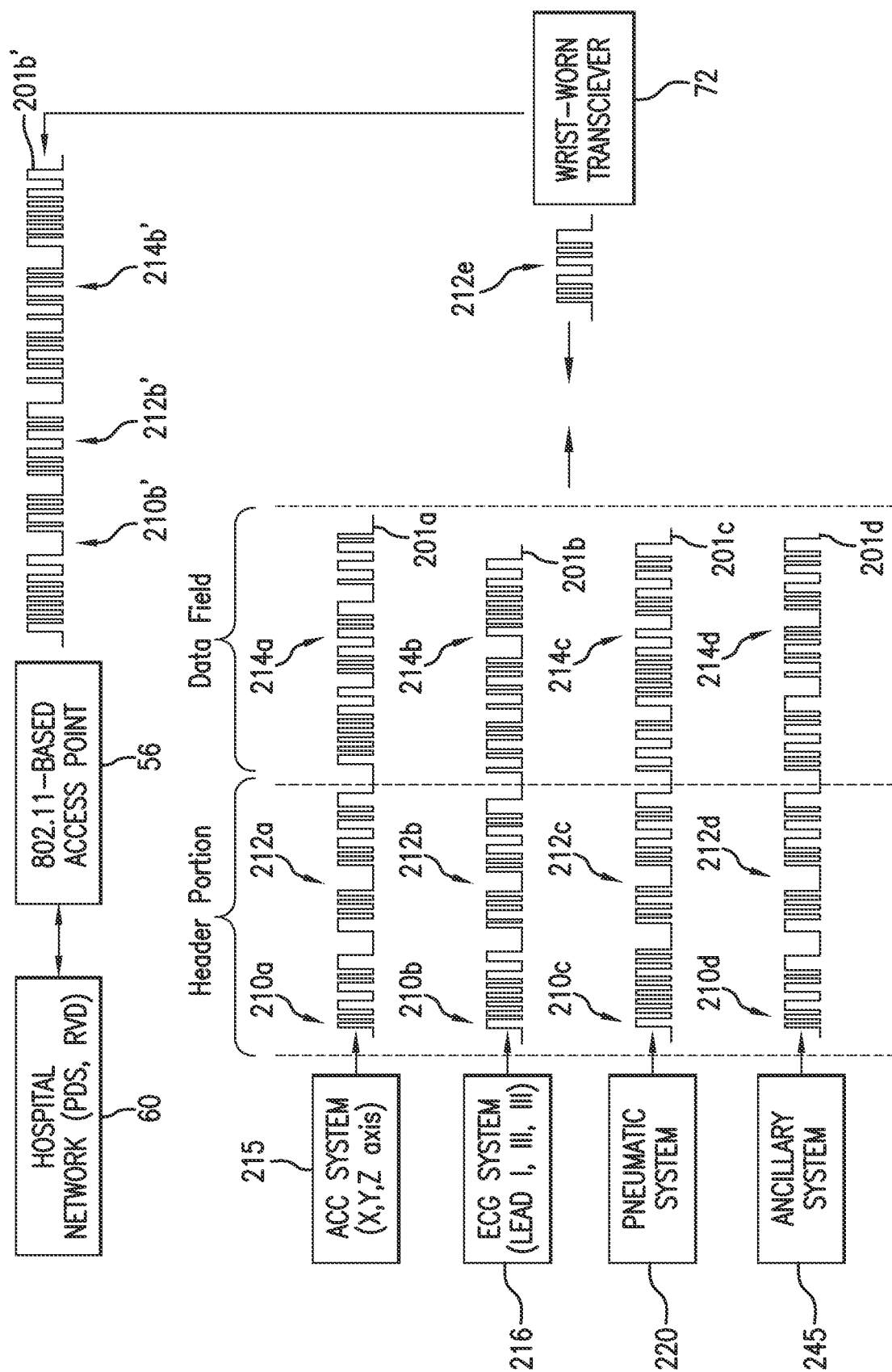


FIG. 25

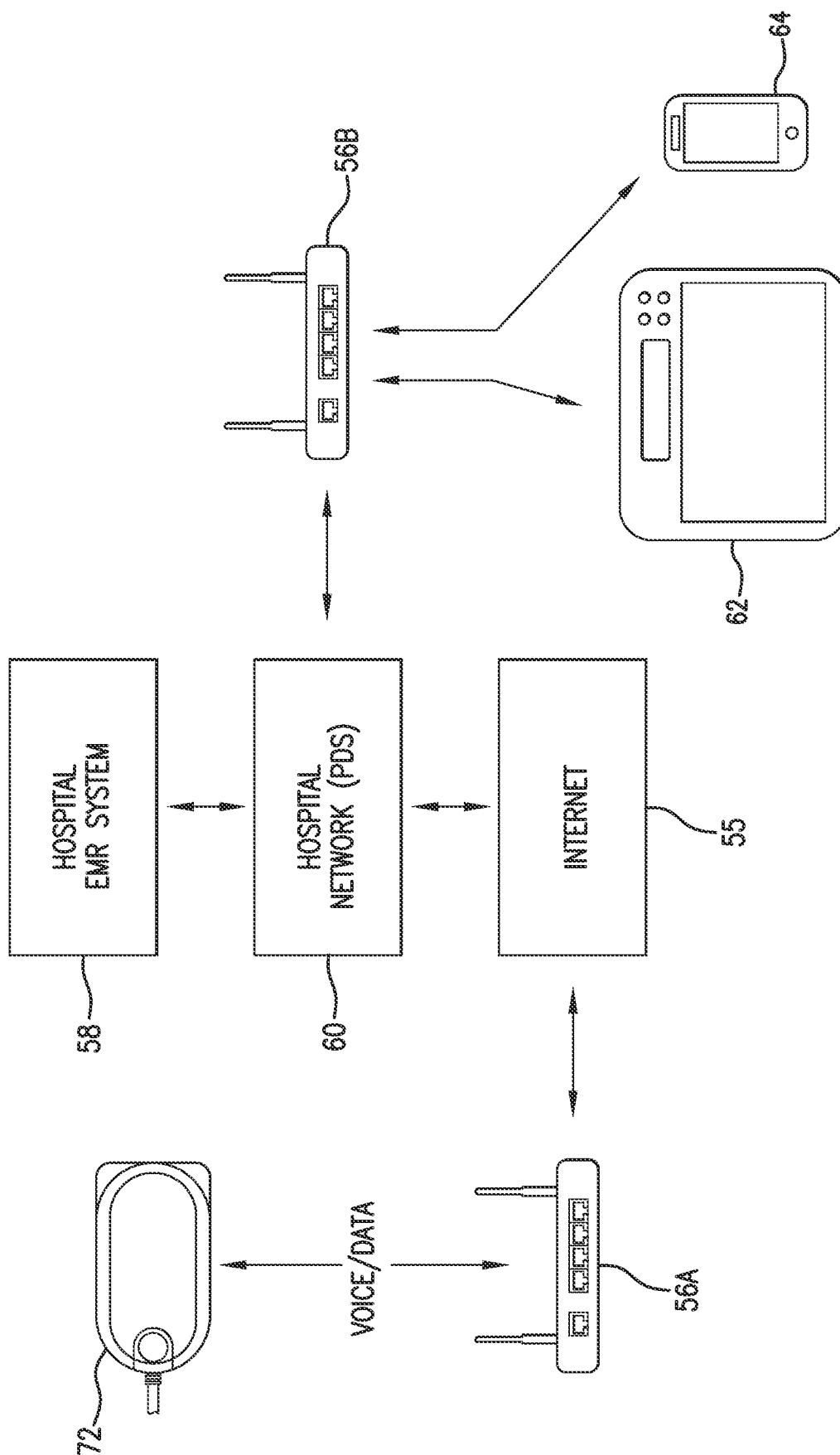


FIG. 26

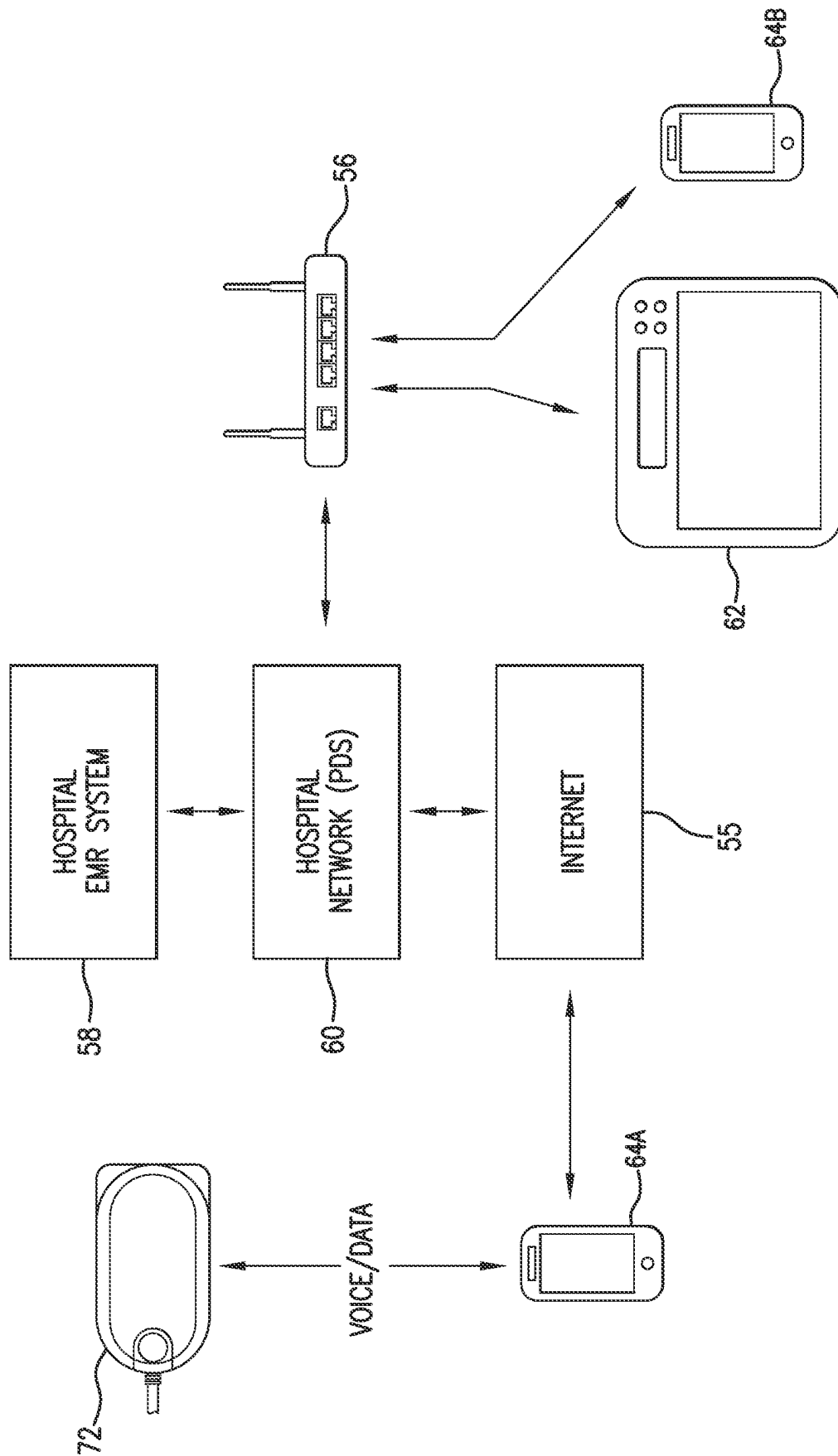


FIG. 27

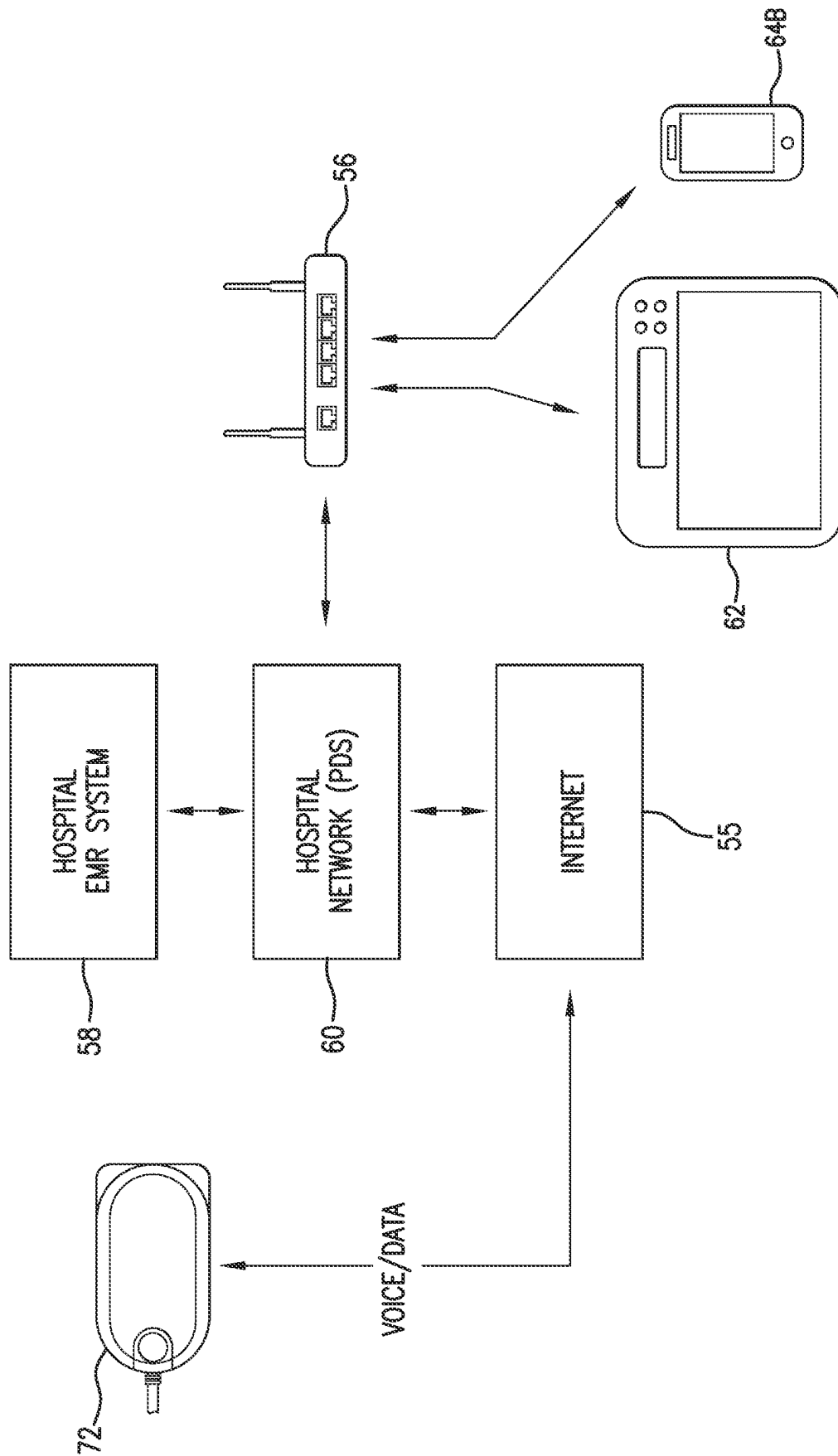


FIG.28

REFERENCES CITED IN THE DESCRIPTION

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Patent documents cited in the description

- US 5316008 A [0007]
- US 5857975 A [0007]
- US 5865755 A [0007]
- US 5649543 A [0007]
- US 2008033255 A1 [0007]
- US 20100168589 A1 [0016] [0048]
- US 20100298656 A1 [0038] [0042] [0047]
- US 20100298650 A1 [0038] [0042] [0047]
- US 20100324385 A1 [0048] [0088]
- US 7396330 B [0059]
- US 20110066050 A1 [0059]
- US 20050216199 A1 [0073]
- US 20090018453 A1 [0078]
- US 20110066037 A [0087]
- US 20050228297 A1 [0108]
- US 20050228296 A1 [0108]
- US 20050228300 A1 [0108]
- US 20050228301 A1 [0108]
- US 20060084878 A1 [0108]
- US 20050228299 A1 [0108]
- US 20050261598 A1 [0108]
- US 20060009697 A1 [0108]
- US 2006009698 A1 [0108]
- US 2007018593 A1 [0108]
- US 20070276261 A1 [0108]
- US 20070276632 A1 [0108]
- US 20080082004 A1 [0108]
- US 20080114220 A1 [0108]
- US 20080221399 A1 [0108]

Non-patent literature cited in the description

- **MELZAK.** The McGill Pain Questionnaire: Major Properties and Scoring Methods. *Pain*, 1975, vol. 1, 277-299 [0070]
- **AFONSO et al.** ECG Beat Detection Using Filter Banks. *IEEE Trans. Biomed Eng.*, 1999, vol. 46, 192-202 [0086]

专利名称(译)	体佩的生命体征监视器		
公开(公告)号	EP2544584B1	公开(公告)日	2017-09-20
申请号	EP2011754055	申请日	2011-03-10
[标]申请(专利权)人(译)	索泰拉无线公司		
申请(专利权)人(译)	SOTERA WIRELESS , INC.		
当前申请(专利权)人(译)	SOTERA WIRELESS , INC.		
[标]发明人	MOON JIM BANET MATT TROMMER GUNNAR DIBATTISTA CAROLINE WATLINGTON TOM DHILLON MARSHAL MCCOMBIE DEVIN		
发明人	MOON, JIM BANET, MATT TROMMER, GUNNAR DIBATTISTA, CAROLINE WATLINGTON, TOM DHILLON, MARSHAL MCCOMBIE, DEVIN		
IPC分类号	A61B5/0205 H04W4/20 A61B5/00		
优先权	12/762777 2010-04-19 US 61/312624 2010-03-10 US 12/762726 2010-04-19 US 12/762790 2010-04-19 US 12/762836 2010-04-19 US 12/762846 2010-04-19 US 12/762751 2010-04-19 US 12/762733 2010-04-19 US 12/762822 2010-04-19 US 12/762810 2010-04-19 US		
其他公开文献	EP2544584A1 EP2544584A4		
外部链接	Espacenet		

摘要(译)

本发明提供一种体戴式生命体征监测器，其测量患者的生命体征（例如，血压，SpO2，心率，呼吸速率和温度），同时表征他们的活动状态（例如，休息，行走，抽搐，下降）和姿势。（直立，仰卧）。监视器处理此信息以最小化生命体征的损坏以及与运动相关的伪像的相关警报/警报。它还具有在触摸屏显示器上呈现的图形用户界面（GUI），有助于简化和改善医院和家庭中的患者监控和安全性。

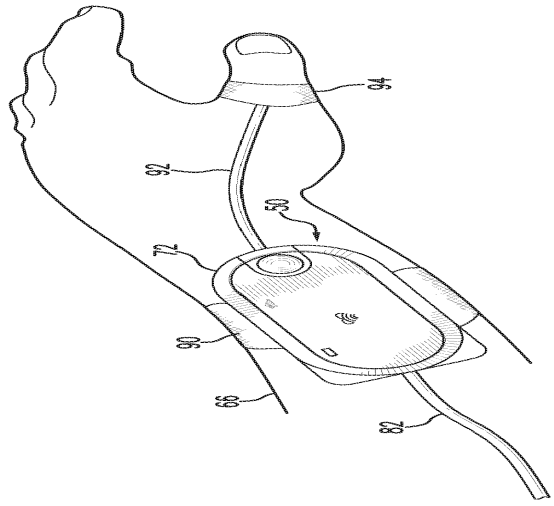


FIG.1