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(54) Patient support apparatus with redundant identity verification

Patientenliegevorrichtung mit redundanter Identitätsverifizierung

Appareil de support de patient avec vérification d'identité redondante

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Description

[0001] The present disclosure is related to patient support apparatuses, such as hospital beds, stretchers, and the like. More specifically, this disclosure relates to a patient support apparatus that can verify the identity of a person or an object and/or monitor the person's vital signs, the patient support comprising a support frame configured to support a patient in a plurality of positions including a horizontal position, and a control module supported on a component of the patient support apparatus, the control module including a first sensing device.

[0002] In healthcare, efforts are often needed to make sure that the patients under care are receiving the proper therapy, medication and/or treatments. To do so, the patient's identity and/or the identity of the person providing or overseeing the therapy, treatment, or medication delivery, may need to be verified. If the patient is to receive medication, the healthcare professional may need to confirm the identity of the medication and/or other details of the patient's prescription and dosing regimen, prior to dispensing the medication to the patient.

[0003] US 2004/0249673 discloses an integrated point-of-care system which includes a medical monitoring device, a medical care device, a computing system, and a structure. The medical monitoring device monitors patient information for a patient. The medical care device provides medical care to the patient. The computing system receives patient information from the medical monitoring device and transmits control instructions to the medical care device to control the medical care to the patient. The computing system also exchanges data with a central data repository through a communication network. The structure supports the patient, the medical monitoring device, the medical care device, and the computing system.

[0004] The invention is defined by the claims.

[0005] The invention will now be further described by way of example with reference to the accompanying drawings, in which:

FIG. 1 is a simplified perspective view of at least one embodiment of a patient support apparatus including a siderail with a control unit having sensing devices for identity verification;

FIG. 2 is a simplified side view of at least one embodiment of another siderail for the patient support apparatus of FIG. 1, including a control unit having sensing devices for identity verification and a display;

FIG. 3 is a simplified partial perspective view of the patient support apparatus of FIG. 1, including at least one embodiment of a siderail with a control unit having a barcode reader device disposed therein;

FIG. 4 is a simplified perspective view of at least one embodiment of the barcode reader device of FIG. 3, and showing in relation thereto illustrative objects having labels that may be scanned by the barcode reader device for identity verification;

FIG. 5 is a simplified partial perspective view of the patient support apparatus of FIG. 1, including at least one embodiment of a siderail, showing a view of a control unit having a pulse oximeter module and pulse oximetry sensor apparatus, from the vantage point of a patient positioned on the patient support apparatus;

FIG. 6 is a simplified perspective view of the control unit of FIG. 5, showing a view of the control unit from the vantage point of a caregiver positioned adjacent to the patient support apparatus;

FIG. 7 is a simplified perspective view of at least one embodiment of another control unit for the patient support apparatus of FIG. 1, including a pulse oximeter module and pulse oximetry sensor apparatus; FIG. 8 is a simplified flow diagram of at least one embodiment of a method for handling data communications with sensing devices of a patient support apparatus for identity verification; and

FIG. 9 is a simplified block diagram of at least one embodiment of a control system for a patient support apparatus.

[0006] Referring now to FIG. 1, an embodiment of a patient support apparatus 100 includes a control unit 132, which has embodied therein an identity verification module 134. The illustrative identity verification module 134 includes a plurality of sensing devices 136, 138 supported thereby. The sensing devices 136, 138 can be used to verify the identity of a person, such as a patient or a caregiver, and/or an object, such as a medication. In this way, redundant identity verification can be performed directly at the point of care, e.g., the location at which the control unit 132 is mounted to the patient support apparatus 100. To do this, each of the sensing devices 136, 138 detects a different form of identifying information, and one or more of the multiple forms of identifying information may be used to authenticate a person or an object. For instance, as illustrated in FIG. 1, the sensing device 136 may be embodied as a non-biometric sensing device such as a barcode reader or radio-frequency identification (RFID) receiver, while the sensing device 138 may be embodied as a biometric sensing device such as a fingerprint sensor. As described in more detail below, other combinations of different types of sensing devices, which are configured to detect different kinds of identifying information, may also be used.

[0007] In the embodiment of FIG. 1, the control unit 132 is removably coupled to a siderail 130 of the patient support apparatus 100, in that the control unit 132 may be physically displaced from the siderail and used as a handheld device. As explained further below, however, the control unit 132 and/or other embodiments thereof, such as those shown in the subsequent figures and described below, may be integrated with the siderail (e.g., in a non-removable fashion), or may be similarly removably or fixedly installed in or mounted to another component of the patient support apparatus 100 (such as a

headboard 120, a footboard 122, or another siderail 124, 126, 128) or to another architectural structure located at the point of care (a headwall, support column, cart, or point-of-care computer system, for example). In general, any of the embodiments of the control units described herein may include any one or more of the features or functions of any of the other control units herein described, whether or not specifically stated.

[0008] The illustrative patient support apparatus 100 is embodied as, for example, a hospital bed, a stretcher, or a similar device that can support a person in a horizontal position and/or one or more non-horizontal positions. The illustrative patient support apparatus 100 is of a type that is typically used in hospitals and/or other facilities in which health care is provided. However, aspects of this disclosure can be applied to any type of person support apparatus or similar structure, including but not limited to beds, mattresses, cushions, tables, stretchers, chairs, wheelchairs and other similar structures, whether or not all of the features of the illustrated patient support apparatus 100 are included in such structure, and whether or not such person support structure includes other features not mentioned herein.

[0009] The patient support apparatus 100 has a base 110, which is movably supported by wheels or casters 116, 118 (view of one caster obstructed in FIG. 1). A frame 112 is coupled to and supported by the base 110. In some embodiments, the frame 112 is supported by a lift mechanism, which operates to raise, lower, and tilt the frame 112 relative to the base 110. In some versions of the patient support apparatus 100, a deck may be coupled to and supported by the frame 112. The deck may have a number of sections, some of which may pivot relative to the frame 112 or to each other to, for example, elevate the head section of the bed, lower the foot section of the bed, or a combination thereof (e.g., to attain a chair position). In some embodiments, movement of the frame 112 and/or deck sections may be driven by actuators (not shown), which may be electronically-controlled based on, for example, user and/or sensor inputs.

[0010] The frame 112 supports a support surface 114 (e.g., a mattress), which, in turn, may support a patient positioned thereon. The support surface 114 may include a number of air bladders, foam, a combination thereof and/or other suitable materials. In embodiments where the support surface 114 includes air bladders, the inflation and deflation of the air bladders may be electronically-controlled, based on, for example, user and/or sensor inputs.

[0011] The electronically-controlled features and functions of the patient support apparatus 100 are managed by a patient support system 900, which is described further below with reference to FIG. 9. Aspects of the system 900 may be implemented at the control unit 132 and/or at a frame- or base-mounted control unit 140, for example. The control unit 140 may include, among other things, computer circuitry for the control of the frame 112, control of the surface 114, and/or network connectivity

(including wired and/or wireless connectivity). The electronically-controlled features and functions of the patient support apparatus 100 are powered by a power supply, such as a battery and/or a connector that can connect the patient support apparatus 100 to a supply of electrical current (e.g. a wall outlet).

[0012] Referring now to FIG. 2, a siderail 200 for the patient support apparatus 100 is shown, from the vantage point of a caregiver facing the patient support apparatus 100 if the siderail were installed in the position of the siderail 130 of FIG. 1. The siderail 200 has mounted thereto a number of bed function controls 210, which can be activated by the caregiver or another person to, for example, raise or lower the frame 112 or adjust the inflation parameters of the surface 114. The siderail 200 has installed therein or mounted thereto a control unit 212, which includes a number of sensing devices 214, 216, 218 and a display 220, which are supported by a housing 228. The housing 228 may be integrated with the siderail 200 (e.g., as a single molded piece) in some embodiments. In other embodiments, the housing 228 may be pivotably or removably coupled to the siderail 200. The sensing devices 214, 216, 218 are each configured to capture a different type of identifying information from a person or an object. For instance, in the illustrative control unit 212, the sensing device 214 includes an RFID receiver, which may be used to detect RFID signals transmitted by, for example, an ID badge or bracelet worn by a person, such as a caregiver or a patient. The sensing device 216 includes an acoustic sensor or microphone, which can be used to capture spoken words or phrases uttered by a caregiver or patient. For example, the sensing device 216 may capture a person's speech in response to a prompt for the person to state their name. The sensing device 218 includes an optical sensor or camera, which can be used to capture digital images or representations of, for example, bar codes, facial features, and/or other identifying features of a person or an object.

[0013] The illustrative display 220 is embodied as a touch screen display, but may be embodied as any suitable type of interactive display, in other embodiments. In response to inputs received by one or more of the sensing devices 214, 216, 218, information about the person whose identity has been verified by the control unit 212 based on the sensor inputs, and/or information about medication whose identity has been verified by the control unit 212 based on the sensor inputs, is displayed on the display 220. For instance, as shown, a digital image 222 of a caregiver is displayed in response to a determination by the control unit 212 that the caregiver's identity has been verified based on a combination of the sensor inputs (e.g., RFID data transmitted by a tag worn by the caregiver and detected by the RFID sensor 214, voice data captured by the microphone 216, barcode data printed on a tag or ID card carried by the caregiver and scanned by the barcode reader/camera 218, and/or facial recognition data detected by the camera 218). The car-

egiver's identity may be verified by the control unit 212 executing computer logic to match each of the multiple sensor inputs to corresponding data stored in one or more computer systems, such as a hospital personnel database. Similarly, in the case where the sensing devices 214, 216, 218 may be used to detect identifying information of a patient, the identity data (e.g., bar code, facial features, voice data, and/or RFID data) may be matched against patient information that may be stored in, for example, a hospital admission, discharge, and transfer (ADT) system.

[0014] A medication container or packaging having a barcode (e.g., a two-dimensional or three-dimensional bar code or Quick Response (QR) code), may similarly be scanned by the sensing device 218. In such event, the control unit 212 may verify the identity of the medication (e.g., the medication type and dosage form) by matching the medication identification data (e.g., bar code data) received by the sensing device 218 to information stored in a medication database (such as a "global" medication database that may be maintained by an external entity or service, such as a pharmacy or health-care network). Such a database may be accessed by the control unit 212 via a connection to a hospital communication network and/or a secure connection to an external network such as the Internet. Once the control unit 212 has verified a medication, information such as the medication name and dosage form may be presented to the caregiver on the display 220, so that the caregiver can input corresponding information about a dose of medication actually dispensed to the patient at the point of care. As shown in FIG. 2, the medication information that may be entered by the caregiver at the display 220 may include the actual amount of medication dispensed to the patient (e.g., number of milligrams, number of tablets, etc.), the site or location at which the medication was dispensed to the patient (e.g., the room number and/or bed number), and the date and/or time at which the medication was dispensed to the patient. As shown in FIG. 2, a history of the patient's previous doses of the medication and/or other medications may be obtained from, for example, the patient's electronic medical record, and also displayed. For example, the display may indicate any other doses of medication that the patient has already received on the current day. Once the caregiver has finished entering the patient's current dose information, the information may be saved to the patient's electronic medical record by the activation of a user interface control 226 (e.g., a physical or virtual button, switch, slide, dial, or the like). Activation of the control 226 may cause the patient's current dose information to be stored locally in memory at the patient support apparatus 100 and/or transmitted to, for example, a hospital's electronic medical records (EMR) system, over a hospital communication network.

[0015] Referring now to FIGS. 3 and 4, perspective views of a handheld or portable barcode reader 314 are shown. In FIG. 3, the barcode reader 314 is shown cou-

pled to a siderail 300 that may be used in connection with the patient support apparatus 100. Although not specifically shown, the siderail 326 may similarly be configured. In FIG. 4, the barcode reader 314 is shown in relation to objects having labels or tags that may be scanned by the barcode reader 314 for identity verification. In some embodiments, the barcode reader 314 may be a dedicated barcode-scanning device. In other embodiments, the barcode reader 314 may be a multifunctional device such as a mobile computing device (e.g., a smart phone or tablet computer equipped with an integrated camera and barcode-scanning software).

[0016] Referring to FIG. 3, the siderail 300 includes a control unit 310 which includes a docking area 312. The docking area 312 is shaped to receive at least a portion of the body of the barcode reader 314, and includes a retainer (e.g., a latch, clip, or detent) to secure the barcode reader 314 therein when the barcode reader 314 is not in use. Although not specifically shown, charging contacts may be supplied within the docking area 312 so that when not in use, the barcode reader 314 may be stored in the docking area 312 in such a way that its charging contacts align with the charging contacts of the docking area 312, so that the barcode reader 314 may be electrically charged by the connection of the charging contacts to the electrical system of the patient support apparatus 100 (e.g., by electrical wiring routed through the siderail 400 to a frame- or base-mounted control unit or power supply).

[0017] As shown in FIG. 4, the barcode reader 314 may be coupled to the control unit 310 by a cord or tether 410, which may include electrical wiring or cables, in some embodiments. In such embodiments, the tether 410 may have at its distal end an electrical connector that connects with an electrical port disposed in the docking area 312, for electrical power and/or data communication with the control unit 310. The illustrative barcode reader 314 includes a user control 412 (e.g. a button or switch) and an optical reader or sensor 414 configured to read or scan or capture an image of a bar code label or tag when the sensor 414 is directed at the label and the user control 412 is activated. For instance, the barcode reader 314 may be used to capture barcode data printed on a label 422 attached to a container or package of medication 420, a label 432 attached to an identification badge or tag 430 that may be worn by a person (e.g., a patient or hospital personnel), or a label 442 attached to a bracelet 440 that may be worn by a person (e.g., a patient, caregiver or other hospital personnel). Electrical circuitry transmits the barcode data received by the barcode reader 314 to the control unit 310, either by the electrical contacts or outlet (to which the tether 410 may be connected) located in the docking area 312. In some embodiments, the control unit 310 may include a display similar to that shown in FIG. 2, to display medication or person information associated with the scanned bar code at the siderail 300. In some embodiments, the control unit 310 may communicate the scanned bar code data

to the patient support system 900 via an internal bed network, such as a Controller Area Network (CAN), which may then interface with an external system through a hospital communication network to receive therefrom medication and/or person data for display at the control unit 310.

[0018] Referring now to FIGS. 5 and 6, a control unit 510 is installed in a siderail 500, which may be mounted to the patient support apparatus 100 adjacent an end-board 522, as shown. A patient-facing side 512 of the control unit 510 is shown in FIG. 5; that is, a view of the control unit 510 from the vantage point of a person positioned on the support surface 114 of the patient support apparatus 100. The patient-facing side 512 supports an electrical data communications port to which a communications link 516 (e.g., insulated wiring or cable) connects a vital signs sensing device 518. The sensing device 518 is, in the illustrated embodiment, a pulse oximetry sensor apparatus or "finger clasp." In other embodiments, other types of physiological sensors may be similarly used and connected to the control unit 510. Vital signs data is communicated from the sensing device 518 to the control unit 510, which displays the vital signs data as shown, for example, in FIG. 6. The control unit 510 may communicate the vital signs data to the patient support system 900 via the internal bed network, which may then communicate the vital signs data to another computer system (such as an EMR system), via a secure network connection as mentioned above.

[0019] In FIG. 6, a caregiver-facing side 610 of the siderail 500 and the control unit 510 is shown. That is, the side 610 is the view of the siderail 500 that would be seen by a person standing near the side of the patient support apparatus 100 to which the siderail 500 is mounted. The caregiver-facing side of the control unit 510 includes a display 614 and a number of user controls 618. The illustrative display 614 displays a graphical representation and a textual representation of vital signs data detected by the sensing device 518. For instance, as shown, the display 614 shows a graphical depiction and a numerical value indicative of the patient's vital signs data (e.g., blood oxygenation levels) over time, while the patient has their finger inserted in the finger clasp 518. The user controls 618 include physical and/or virtual controls similar to those described above. Some of the user controls 618 may be configured to control one or more of the electronically-controllable features or functions of the patient support apparatus 100. Other controls 618 may control the display 614 to, e.g., change a display parameter such as the time scale or an aspect of the graphical or textual display.

[0020] In some embodiments, vital signs data collected by the sensing device 518, or the lack thereof, may be used to determine whether a person is present or absent from the patient support apparatus 100. For instance, if the control unit 510 is receiving vital signs data, it may be inferred that a patient is positioned on the patient support apparatus 100, and vice versa. In response, the con-

trol unit 510 may automatically update the configuration of the patient support apparatus 100. For instance, if the control unit 510 infers from the vital signs data that a patient is present on the patient support apparatus 100, the control unit 510 may initiate a patient weighing function, a pressure relief function, or some other electronically-controllable feature or function of the patient support apparatus 100, or may evaluate the vital signs data and issue an alert to inform healthcare personnel if it appears from the vital signs data that the patient may need assistance. Likewise, if the control unit 510 infers from the absence of vital signs data that the patient has exited the patient support apparatus 100, the control unit 510 may terminate or suspend a bed feature or function that was previously in use, or may issue an alert to inform healthcare personnel of the patient's status.

[0021] In some embodiments, the vital signs data collected by the sensing device 518 may be used to verify the identity of the patient. For instance, a sampling of such vital signs data captured over time by the sensing device 518 may reveal a vital signs "signature" that can be compared to vital signs signatures of a number of different persons, which may be stored in a hospital or healthcare database. If the patient's vital signs signature is found to match a stored vital signs signature, the patient may be identified. If a match is not found, another form of identification may be requested by the patient support apparatus 100.

[0022] Referring now to FIG. 7, a control unit 700 includes similar features and functionality as the control unit 510; however, the control unit 700 is embodied as a portable pulse oximeter module that is configured to be removably coupled to a siderail of the patient support apparatus 100. As such, the control unit 700 may be equipped with a wireless transceiver and associated circuitry, in order to communicate vital signs data wirelessly to the patient support system 900. The illustrative control unit 700 is embodied in a housing, which includes a caregiver-facing panel 710, a pair of opposing side panels 712 (view of one panel obstructed), and top and bottom portions 714, 716. The illustrative portions 714, 716 are concavely formed or otherwise configured to cooperate with or be received by corresponding mating portions of a siderail (e.g., the siderail 130) so that the control unit 700 may be removably received and secured within a recess or window of the siderail (e.g., the window 142 of the siderail 130). A data communications port 718 is illustratively supported by the side panel 712, but may be placed in any suitable location on the control unit 700 according to the requirements of a particular design. The vital signs sensor or finger clasp 518 is connected to the data communications port 718 by a cable 720. As such, vital signs data (e.g., pulse oximetry data) collected from a patient whose finger is inserted in the finger clasp 518 may be communicated to the control unit 700. The control unit 700 displays graphical 726 and textual/numerical information 728 relating to or representative of the vital signs data collected by the sensing device 518, on a dis-

play 724. User controls 722 may be used to configure the display 724 according to the needs of a particular healthcare environment or healthcare situation, in a similar fashion as discussed above.

[0023] Referring now to FIG. 8, a method 800, which may be implemented as executable instructions, modules, routines, or logic units, for example, executed by the system 900, and may be embodied in one or more control modules or units of the patient support apparatus 100, is shown. At block 810, the patient support apparatus 100 receives a combination of at least two different types of person identity data from one or more of the sensing devices described above. For instance, the patient support apparatus 100 may receive bar code data from the bar code reader 136 and fingerprint data from the fingerprint sensor 138. In some embodiments, the patient support apparatus 100 may prompt the user for a secondary type of identity data upon having detected a first, different, type of identity data. For instance, in some embodiments, the patient support apparatus 100 may prompt the user for voice or bar code input upon having received biometric or vital signs data, or vice versa. At block 812, the patient support apparatus 100 authenticates each of the received forms of the person identity data according to its type. For instance, if fingerprint data is received, the patient support apparatus 100 may analyze and/or compare the fingerprint data to a database of stored fingerprint data, which may be accessible to the patient support apparatus 100 by a hospital communication network. Similarly, if voice, barcode, RFID, or facial feature data is received, such data may be analyzed and/or compared to stored samples of that data type. If a match is found (e.g., within reasonable tolerances) for each of the received types of identity data, the patient identity is verified at block 814. Such analyses and comparisons may be carried out by computer logic of one or more modules of the patient support system 900 or, e.g., "in the cloud," at one or more computing devices to which the person support apparatus 100 is communicatively coupled via the hospital communication network 950. If a match is not found for at least two different types of identity data, then additional identifying data may be requested. If the person's identity is not verified by at least two different types of identifying data, then at block 828 the person support apparatus 100 may send an alert to a caregiver or other healthcare personnel, or display the alert at a point-of-care display (such as the display 220 of the control unit 212 of FIG. 2), as needed according to the requirements of a particular design. If the person's identity is verified (e.g., by at least two forms of identifying data), then at block 816, the person's identity is associated with the patient support apparatus and data relating to the person's identity (e.g., name, title, digital image, etc.) may be displayed on a display located at the point of care (e.g., at a siderail of the patient support apparatus 100). If the person whose identity has been verified is a caregiver or other hospital personnel, then the person's identity may be associated

with the patient support apparatus 100 for, e.g., healthcare facility workflow monitoring or other reasons. If the person whose identity has been verified is a patient, the person's identity may be associated with the patient support apparatus 100 in order to associate medication, treatment, and/or therapy data generated at the patient support apparatus 100 with the patient's electronic medical record, to track the patient's progress through the healthcare facility, or for other reasons. Data relating to the person's identity may be displayed at the patient support apparatus 100 as shown in FIG. 2, for example.

[0024] In some embodiments, at block 818, the configuration or status of an electronically-controllable function or feature of the patient support apparatus 100 may be modified or updated, or an alarm or an alert may be issued (e.g., to a nurse call system) based on the verification of the person's identity, whether such verification is made by biometric sensors, vital signs signatures, or otherwise. For example, if the person is a caregiver, the available bed functions or features may be updated to reflect only those functions or features that the caregiver is able to activate, based on the caregiver's role or access privileges. If the person is a patient, the configuration of the bed functions or features may be configured based on aspects of the patient's medical condition, which may be known to the patient support apparatus 100 through an interface with an EMR system or based on vitals signs data collected at the patient support apparatus 100 from, e.g., a pulse oximeter or other physiological sensor.

[0025] At block 820, the patient support apparatus 100 determines whether medication identifying information has been received by one of the identity verification sensing devices. For instance, the patient support apparatus 100 determines whether a barcode scanned by, e.g., the barcode reader 136, 314 relates to a medication. If medication identifying information has not been received, then at block 826, the updated patient information (e.g., the fact that the patient's identity has been verified and any resulting changes to the bed features or functions) may be transmitted to an EMR system or other healthcare system as needed. If medication identifying information has been received at block 820, then at block 822 the patient support apparatus 100 determines whether the medication is verified for the person whose identity was verified at block 814. For instance, if the person whose identity was verified at block 814 is a caregiver, then at block 822 the patient support apparatus 100 determines whether the caregiver is authorized to dispense the medication. To do so, the patient support system 900 may interface with a hospital personnel database, for example, via the hospital communication network 950. If the person whose identity was verified at block 814 is a patient, then the patient support apparatus 100 determines whether the patient's physician has authorized the dispensing of the identified medication to the patient and if so, in what amount. To do so, the patient support system 900 may interface with a hospital EMR system or prescription medication database, for example, via the hos-

pital communication network 950.

[0026] If the identified medication is not verified for the identified person (e.g., a caregiver or patient), then the patient support apparatus 100 may send an alert to a healthcare communication system or display a notification at a point of care location, as needed. If the identified medication is verified for the identified person, then at block 824 medication dispensing information may be displayed and/or received at the patient support apparatus 100. For instance, in a first loop of the method 800, a caregiver's identity may be verified at block 814, and at block 822, the caregiver may be approved to dispense a medication identified at block 820. Then, at block 824, information about the medication may be displayed to the caregiver at the patient support apparatus 100. For example, a digital image of the type of medication associated with a barcode received at block 820 may be displayed, so that the caregiver can visually verify that the medication is correct. At block 826, information relating to the caregiver authorization and/or the medication authorization may be sent to an appropriate healthcare system (such as a hospital workflow tracking system) at block 826.

[0027] In a second or subsequent loop of the method 800, a patient's identity may be verified at block 814, and at block 822, the patient may be approved to receive the medication identified at block 820. Then, at block 824, the display (e.g., the display 220) may permit the caregiver to enter data relating to the medication actually dispensed and received by the identified patient, as described above. At block 826, the patient support apparatus 100 may send the newly received information about the medication actually dispensed to the patient to the patient's medical record or to a healthcare system (e.g., an EMR system), as needed.

[0028] Referring now to FIG. 9, a patient support system 900 including an illustrative patient support apparatus control system 910 is shown in greater detail. As noted above, portions of the patient support apparatus control system 910 may be embodied in different structural units of the patient support apparatus 100. For example, some modules and components of the patient support apparatus control system 910 may be embodied in a siderail- or endboard-mounted control unit, while other modules or components may be embodied in a frame- or base-mounted control unit, which may be in communication with the siderail- or endboard-mounted control unit by a communications network. Such a network may be configured according to a Controller Area Network or Echelon protocol, for example, or according to any other suitable bed network communications protocol.

[0029] In the illustrative patient support system 900, the patient support apparatus control system 910 is in communication with the electronically-controllable features and functions of the patient support apparatus 100 as mentioned above, and with one or more other computing systems 952, 954, 958, 964 via the hospital communication network 950. Portions of the patient support

apparatus control system 910 may be local to the patient support apparatus 100, however, some portions thereof may be distributed across one or more of the other computing systems 952, 954, 958, 964. For instance, in some cases, identity verification algorithms and/or databases may be located on one or more other computing systems 964.

[0030] In general, the patient support apparatus control system 910 includes at least one computing device (such as an embedded device, a desktop computer, a portable computer, a wall-mounted unit, or a server, for example) that includes one or more processors or processor cores (e.g., microprocessors, microcontrollers, digital signal processors, etc.) 912, memory 914, and an input-output subsystem 916. For instance, portions of the control system 910 may be embodied in one or more physical control units of the patient support apparatus 100 and/or any type of computing device including a point of care device such as a "nurse's station" or "patient station" of a nurse-call system, a server, a network of computers, or a combination of computers and/or other electronic devices.

[0031] The I/O subsystem 916 typically includes, among other things, an I/O controller, a memory controller, and one or more I/O ports. In some embodiments, the I/O subsystem 916 may include a CAN bus and/or other types of communication links. The processor 912 and the I/O subsystem 916 are communicatively coupled to the memory 914. The memory 914 may be embodied as any type of suitable computer memory device (e.g., volatile memory such as various forms of random access memory).

[0032] The I/O subsystem 916 is communicatively coupled to a number of hardware and/or software components including user interface/controls 918 (e.g., any of the user controls described above), an identity verification module 920 (which includes a person identity verification module 922 and a medication identity verification module 924), a patient monitoring module 926 (which includes a vital signs monitoring module 928 and a patient presence/absence monitoring module 930), one or more frame/deck/surface control module(s) 932 (which control features of the bed frame, deck or support surface of the patient support apparatus 100, as mentioned above), one or more machine-readable storage media 934 (which may store patient data 936 and/or bed data 938), one or more sensing devices 940 (e.g., the biometric sensing devices and/or non-biometric sensing devices described above), one or more readers/receiver(s) 924 (e.g., RFID receivers, wireless communication transceivers, etc.), a display 944 (e.g., any of the displays described above), and a communication interface 946.

[0033] The storage media 934 may include one or more hard drives or other suitable data storage devices (e.g., flash memory, memory cards, memory sticks, and/or others). In some embodiments, portions of the modules 920, 926, 932 may reside at least temporarily in the storage media 934. Portions of these modules 920,

926, 932 may be copied to the memory 914 during operation of the patient support apparatus control system 910, for faster processing or other reasons.

[0034] The hospital communication network 950 may communicatively couple the person support apparatus control system 910 to one or more hospital or healthcare facility computer systems, for example. Accordingly, the communication interface 946 may include one or more wired or wireless network interface cards or adapters, for example, as may be needed pursuant to the specifications and/or design of the particular system 900.

[0035] Computing systems in communication with the hospital communication network 950 may include, for example, a healthcare communication system 952 (e.g., a patient-nurse communication system or "nurse call" system), a locating and tracking system 954 (e.g., a system that monitors the location of caregivers, patients, and/or equipment in a healthcare facility), an EMR system 958, and other system(s) 964 (e.g., an admission, transfer and discharge system, hospital workflow system, etc.). The patient support system 900 may include other components, sub-components, and devices not illustrated in FIG. 9 for clarity of the description. In general, the components of the person support system 900 are communicatively coupled as shown in FIG. 9 by electronic signal paths, which may be embodied as any type of wired or wireless signal paths capable of facilitating communication between the respective devices and components.

[0036] In more detail, the identity verification module 920 processes identity data received by the sensing devices 940 and/or readers/receivers 942. If the identity data relates to a person, the person identity verification module 922 verifies the received identity data. For example, the person identity verification module 922 may access person identity data 966, and compare the received identity data to corresponding data stored in a person identity database, depending on the data type of the identity data. If the identity data is received from a fingerprint reader, for example, the person identity verification module 922 compares the received fingerprint data to data stored in a fingerprints database 968. Similarly, the person identity verification module 922 compares received voice data to stored voice recordings 970, compares received ID codes (e.g., RFID codes or barcodes) to stored ID codes 972, compares received digital images (e.g., facial features) to stored images 974, and compares received vitals signs signatures to stored vital signs signatures 976, in order to verify a person's identity. Likewise, if the received identity data relates to a medication, the medication identity verification module 924 compares the received identity information (e.g., barcode) to medication data 978, in order to verify the identity of the medication. The identity verification module 920 or portions thereof may be embodied as software stored and executed locally, e.g., at a siderail-mounted control unit of the patient support apparatus, or portions thereof may be distributed across one or more of the other systems 952, 954, 958, 964 connected to the hospital com-

munication network 950.

[0037] When a person's identity is verified, the person (e.g., a caregiver or a patient) may be associated with the patient support apparatus 100 as described above. To do this, the identity verification module 920 may communicate with the locating/tracking system 954 to obtain identifying information about the patient support apparatus 100 and create an association between the person identity data and the identifying information for the patient support apparatus 100. Such information about the person and the patient support apparatus 100, and the association therebetween, may be stored in the storage media 934, e.g., in a patient data structure 936 and/or a bed data structure 938.

[0038] When a patient's identity is verified, the identity verification module 920 may access or update medical records 960 and/or prescription data 962 that are associated with the identified patient, as described above. Similarly, when a medication is identified, the identity verification module 920 may access or update the prescription data 962.

[0039] The patient monitoring module 926 receives data from vital signs monitoring devices such as the pulse oximetry device described above and/or other physiological sensors. As mentioned above, such vital signs data may be used to determine the presence or absence of a patient on the patient support apparatus 100, or to identify the patient by comparison to the vital signs signatures 976. Accordingly, a vital signs monitoring module 928 receives the vital signs data from the physiological sensors 940 and evaluates the comparison of the received data to the stored vital signs signatures 976, to determine whether there is a match. A presence/absence monitoring module 930 monitors the vital signs data or lack thereof and draws therefrom inferences as to whether the patient is present or absent on the patient support apparatus 100. For example, the module 930 may compare the level and/or frequency of the vital signs signals to one or more threshold values or ranges of values and based on that comparison, determine whether the patient is likely present or absent.

[0040] Embodiments in accordance with the disclosure may be implemented in hardware, firmware, software, or any combination thereof. Embodiments may also be implemented as instructions stored using one or more machine-readable media, which may be read and executed by one or more processors. A machine-readable medium may include any mechanism for storing or transmitting information in a form readable by a machine. For example, a machine-readable medium may include any suitable form of volatile or non-volatile memory.

[0041] Modules, data structures, and the like defined herein are defined as such for ease of discussion, and are not intended to imply that any specific implementation details are required. For example, any of the described modules and/or data structures may be combined or divided into sub-modules, sub-processes or other units of computer code or data as may be required by a particular

design or implementation of the system 900.

[0042] In the drawings, specific arrangements or orderings of schematic elements may be shown for ease of description. However, the specific ordering or arrangement of such elements is not meant to imply that a particular order or sequence of processing, or separation of processes, is required in all embodiments. In general, schematic elements used to represent instruction blocks or modules may be implemented using any suitable form of machine-readable instruction, and each such instruction may be implemented using any suitable programming language, library, application programming interface (API), and/or other software development tools or frameworks. Similarly, schematic elements used to represent data or information may be implemented using any suitable electronic arrangement or data structure. Further, some connections, relationships or associations between elements may be simplified or not shown in the drawings so as not to obscure the disclosure. Although certain illustrative embodiments have been described in detail above, variations and modifications exist.

Claims

1. A patient support apparatus (100) comprising:

a support frame (112) configured to support a patient in a plurality of positions including a horizontal position;
a first sensing device (136, 214, 218) supported on a component (120, 122, 124, 126, 228, 200, 300, 500) of the patient support apparatus;
a second sensing device (138, 518) supported on the component, the second sensing device (138) being a different type of sensing device than the first sensing device (136, 214, 218); and
a control module (132, 134, 212, 310, 510, 700, 900), removably coupled to a siderail (200, 300, 500), in communication with the first sensing device (136) and the second sensing device (138, 518) to verify the identity of a person based on first identity information received by the first sensing device (136, 214, 218) and second identity information received by the second sensing device (138, 518), the first identity information being different than the second identity information, wherein:

the first sensing device (136, 214, 218) and the second sensing device (138, 518) are coupled to the control module (132, 134, 212, 310, 510, 700, 900);
the siderail (200, 300, 500) has mounted thereto a plurality of bed function controls (210);
the control module (132, 134, 212, 310, 510, 700, 900) is configured to analyse vital signs

data produced by the second sensing device 138, 518, the second sensing device being a pulse oximeter sensor apparatus, to determine a vital signs signature for the person for comparison to stored vital signs signatures to verify the identity of the person; and
the control module (132, 134, 212, 310, 510, 700, 900) initiates at least one electronically-controllable function of the patient support apparatus in response to the vital signs data.

2. The patient support apparatus of claim 1, wherein the control module (132, 134, 212, 310, 510, 700, 900) is in communication with the first sensing device (214, 218) to verify the identity of a medication based on medication identity information received by the first sensing device.
3. The patient support apparatus of claim 2, wherein the control module (212) comprises a display (220), in communication with the control module (212) to display content relating to a medication whose identity has been verified by the control module (212) and to display content relating to a person whose identity has been verified by the control module (212).
4. The patient support apparatus of claim 3, wherein the content relating to the medication comprises one or more of dosing information relating to the medication and a digital image of the medication.
5. The patient support apparatus of either claim 3 or claim 4, wherein the content relating to the person comprises one or more of prescription information relating to the person and a digital image of the person.
6. The patient support apparatus of any one of claims 2 to 5, wherein the control module (212, 226) is configured to send data relating to a medication verified by the control module to an electronic medical records system.
7. The patient support of any one of claims 2 to 6, wherein the control module (212) is configured to send an alert to a healthcare communication system (950) if the control module (212) is unable to verify the identity of the medication.
8. The patient support apparatus of any preceding claim, wherein the control module (212, 226) is configured to send data relating to a person verified by the control module to an electronic medical records system.

9. The patient support of any preceding claim, wherein the control module is configured to send an alert to a healthcare communication system if the control module is unable to verify the identity of the person.

Patentansprüche

1. Eine Patientenliegevorrichtung (100), umfassend:

einen Stützrahmen (112), der so konfiguriert ist, dass ein Patient in einer Vielzahl an Positionen, einschließlich einer horizontalen Position, darauf liegen kann;
eine erste Sensorvorrichtung (136, 214, 218), die von einer Komponente (120, 122, 124, 126, 228, 200, 300, 500) der Patientenliegevorrichtung getragen wird;
eine zweite Sensorvorrichtung (138, 518), die von der Komponente getragen wird, wobei sich der Typ der zweiten Sensorvorrichtung (138) vom Typ der ersten Sensorvorrichtung (136, 214, 218) unterscheidet; und
ein Steuerungsmodul (132, 134, 212, 310, 510, 700, 900), das beweglich an eine Seitensicherung (200, 300, 500) gekoppelt ist, in Kommunikation mit der ersten Sensorvorrichtung (136) und der zweiten Sensorvorrichtung (138, 518), um die Identität einer Person, basierend auf der ersten Identitätsinformation, die von der ersten Sensorvorrichtung (136, 214, 218) empfangen wird, und der zweiten Identitätsinformation, die von der zweiten Sensorvorrichtung (138, 518) empfangen wird, zu verifizieren, wobei sich die erste Identitätsinformation von der zweiten Identitätsinformation unterscheidet, wobei:

die erste Sensorvorrichtung (136, 214, 218) und die zweite Sensorvorrichtung (138, 518) an das Steuerungsmodul (132, 134, 212, 310, 510, 700, 900) gekoppelt sind;
an der Seitensicherung (200, 300, 500) eine Vielzahl an Bedienelementen für die Bettfunktionen (210) montiert ist;
das Steuerungsmodul (132, 134, 212, 310, 510, 700, 900) so konfiguriert ist, dass es Vitalparameter-Daten analysiert, die von der zweiten Sensorvorrichtung (138, 518) erzeugt werden, wobei die zweite Sensorvorrichtung eine Pulsoximeter-Sensorvorrichtung ist, um für die Person eine Vitalparameter-Signatur zum Vergleich mit gespeicherten Vitalparameter-Signaturen zu bestimmen, um die Identität der Person zu verifizieren; und
das Steuerungsmodul (132, 134, 212, 310, 510, 700, 900) mindestens eine elektronisch steuerbare Funktion der Patientenlie-

gevorrichtung als Reaktion auf die Vitalparameter-Daten in Betrieb setzt.

2. Die Patientenliegevorrichtung nach Anspruch 1, wobei das Steuerungsmodul (132, 134, 212, 310, 510, 700, 900) in Kommunikation mit der ersten Sensorvorrichtung (214, 218) steht, um die Identität einer Medikation basierend auf einer Medikations-Identitätsinformation, die von der ersten Sensorvorrichtung empfangen wird, zu verifizieren.
3. Die Patientenliegevorrichtung nach Anspruch 2, wobei das Steuerungsmodul (212) eine Anzeige (220) umfasst, die in Kommunikation mit dem Steuerungsmodul (212) steht, um Inhalte bezüglich einer Medikation anzuzeigen, deren Identität vom Steuerungsmodul (212) verifiziert wurde, und um Inhalte bezüglich einer Person anzuzeigen, deren Identität vom Steuerungsmodul (212) verifiziert wurde.
4. Die Patientenliegevorrichtung nach Anspruch 3, wobei die Inhalte bezüglich der Medikation eine oder mehrere Dosierungsinformationen bezüglich der Medikation und ein digitales Bild der Medikation umfassen.
5. Die Patientenliegevorrichtung nach entweder Anspruch 3 oder Anspruch 4, wobei die Inhalte bezüglich der Person eine oder mehrere Verschreibungsinformationen bezüglich der Person und ein digitales Bild der Person umfassen.
6. Die Patientenliegevorrichtung nach einem der Ansprüche 2 bis 5, wobei das Steuerungsmodul (212, 226) so konfiguriert ist, dass es Daten bezüglich einer Medikation, die vom Steuerungsmodul verifiziert wurden, an ein System für elektronische Patientenakten sendet.
7. Die Patientenliege nach einem der Ansprüche 2 bis 6, wobei das Steuerungsmodul (212) so konfiguriert ist, dass es eine Warnung an ein Kommunikationssystem für das Gesundheitswesen (950) sendet, wenn das Steuerungsmodul (212) die Identität der Medikation nicht verifizieren kann.
8. Die Patientenliegevorrichtung nach einem der vorstehenden Ansprüche, wobei das Steuerungsmodul (212, 226) so konfiguriert ist, dass es Daten bezüglich einer Person, die vom Steuerungsmodul verifiziert wurden, an ein System für elektronische Patientenakten sendet.
9. Die Patientenliege nach einem der vorstehenden Ansprüche, wobei das Steuerungsmodul so konfiguriert ist, dass es eine Warnung an ein Kommunikationssystem für das Gesundheitswesen sendet, wenn das Steuerungsmodul die Identität der Person

nicht verifizieren kann.

Revendications

1. Appareil de support de patient (100) comprenant :

un cadre de support (112) conçu pour supporter un patient dans une pluralité de positions, y compris une position horizontale ;
un premier dispositif de détection (136, 214, 218) supporté par un composant (120, 122, 124, 126, 228, 200, 300, 500) de l'appareil de support de patient ;

un deuxième dispositif de détection (138, 518) supporté par le composant, le deuxième dispositif de détection (138) étant un type de dispositif de détection différent du premier dispositif de détection (136, 214, 218) ; et

un module de commande (132, 134, 212, 310, 510, 700, 900), couplé de façon amovible à une ridelle (200, 300, 500), en communication avec le premier dispositif de détection (136) et le deuxième dispositif de détection (138, 518) pour vérifier l'identité d'une personne sur la base de premières informations d'identité reçues par le premier dispositif de détection (136, 214, 218) et de deuxième informations d'identité reçues par le deuxième dispositif de détection (138, 518), les premières informations d'identité étant différentes des deuxième informations d'identité, dans lequel :

le premier dispositif de détection (136, 214, 218) et le deuxième dispositif de détection (138, 518) sont couplés au module de commande (132, 134, 212, 310, 510, 700, 900) ; une pluralité de commandes de fonction de lit (210) sont montées sur la ridelle (200, 300, 500) ;

le module de commande (132, 134, 212, 310, 510, 700, 900) est conçu pour analyser les données de signes vitaux produites par le deuxième dispositif de détection (138, 518), le deuxième dispositif de détection étant un capteur d'oxymètre de pouls, pour déterminer une caractéristique de signes vitaux pour la personne pour la comparer à des caractéristiques de signes vitaux stockées afin de vérifier l'identité de la personne ; et

le module de commande (132, 134, 212, 310, 510, 700, 900) lance au moins une fonction pouvant être commandée électriquement de l'appareil de support de patient en réponse aux données de signes vitaux.

2. Appareil de support de patient selon la revendication 1, dans lequel le module de commande (132, 134, 212, 310, 510, 700, 900) est en communication avec le premier dispositif de détection (214, 218) pour vérifier l'identité d'un médicament sur la base des informations d'identité de médicaments reçues par le premier dispositif de détection.

3. Appareil de support de patient selon la revendication 2, dans lequel le module de commande (212) comprend un afficheur (220), en communication avec le module de commande (212) pour afficher un contenu relatif à un médicament dont l'identité a été vérifiée par le module de commande (212) et pour afficher le contenu relatif à une personne dont l'identité a été vérifiée par le module de commande (212).

4. Appareil de support de patient selon la revendication 3, dans lequel le contenu relatif au médicament comprend une ou plusieurs informations de dosage relatives au médicament et une image numérique du médicament.

5. Appareil de support de patient selon la revendication 3 ou la revendication 4, dans lequel le contenu relatif à la personne comprend une ou plusieurs informations de prescription relatives à la personne et une image numérique de la personne.

6. Appareil de support de patient selon l'une quelconque des revendications 2 à 5, dans lequel le module de commande (212, 226) est conçu pour envoyer des données relatives à un médicament vérifié par le module de commande à un système de dossiers médicaux électroniques.

7. Support de patient selon l'une quelconque des revendications 2 à 6, dans lequel le module de commande (212) est conçu pour envoyer une alerte à un système de communication de services de santé (950) si le module de commande (212) n'est pas en mesure de vérifier l'identité du médicament.

8. Appareil de support de patient selon l'une quelconque des revendications précédentes, dans lequel le module de commande (212, 226) est conçu pour envoyer des données relatives à une personne vérifiée par le module de commande à un système de dossiers médicaux électroniques.

9. Support de patient selon l'une quelconque des revendications précédentes, dans lequel le module de commande est conçu pour envoyer une alerte à un système de communication de services de santé si le module de commande n'est pas en mesure de vérifier l'identité de la personne.

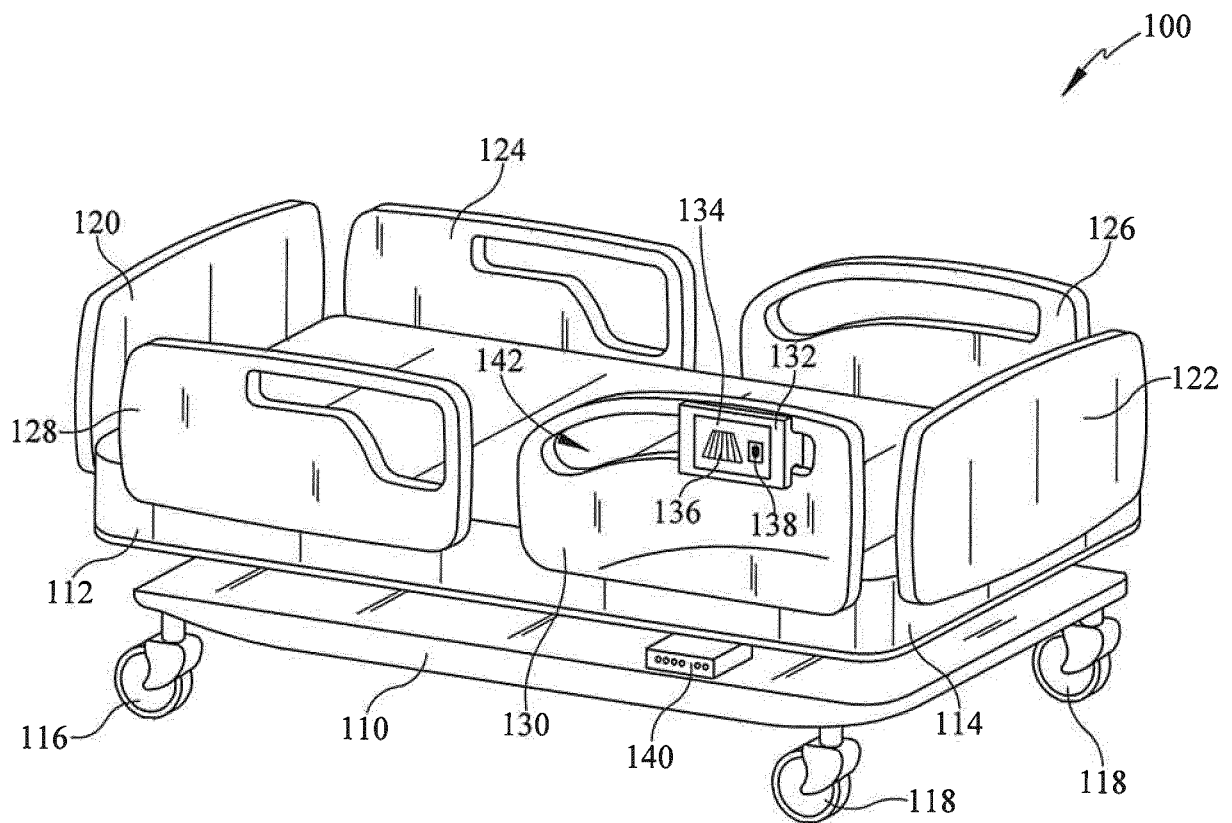


FIG. 1

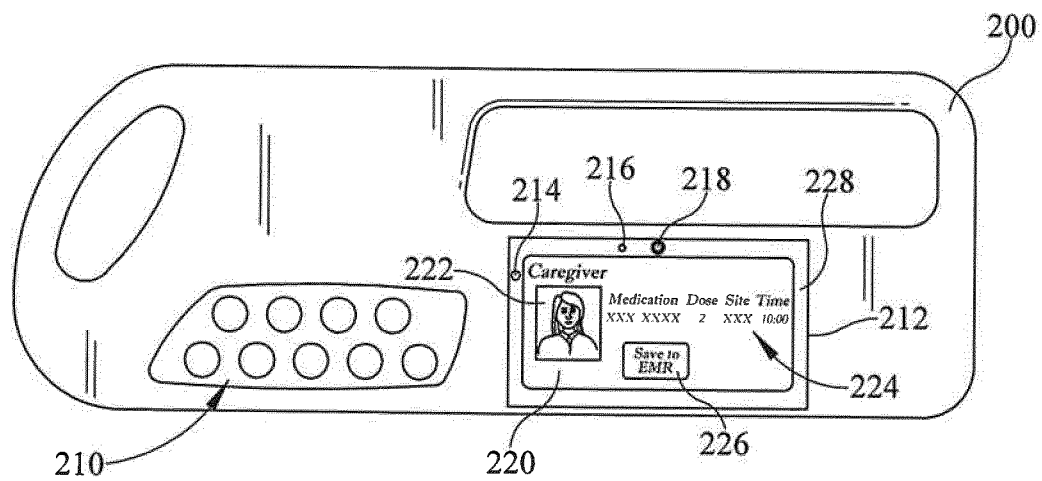


FIG. 2

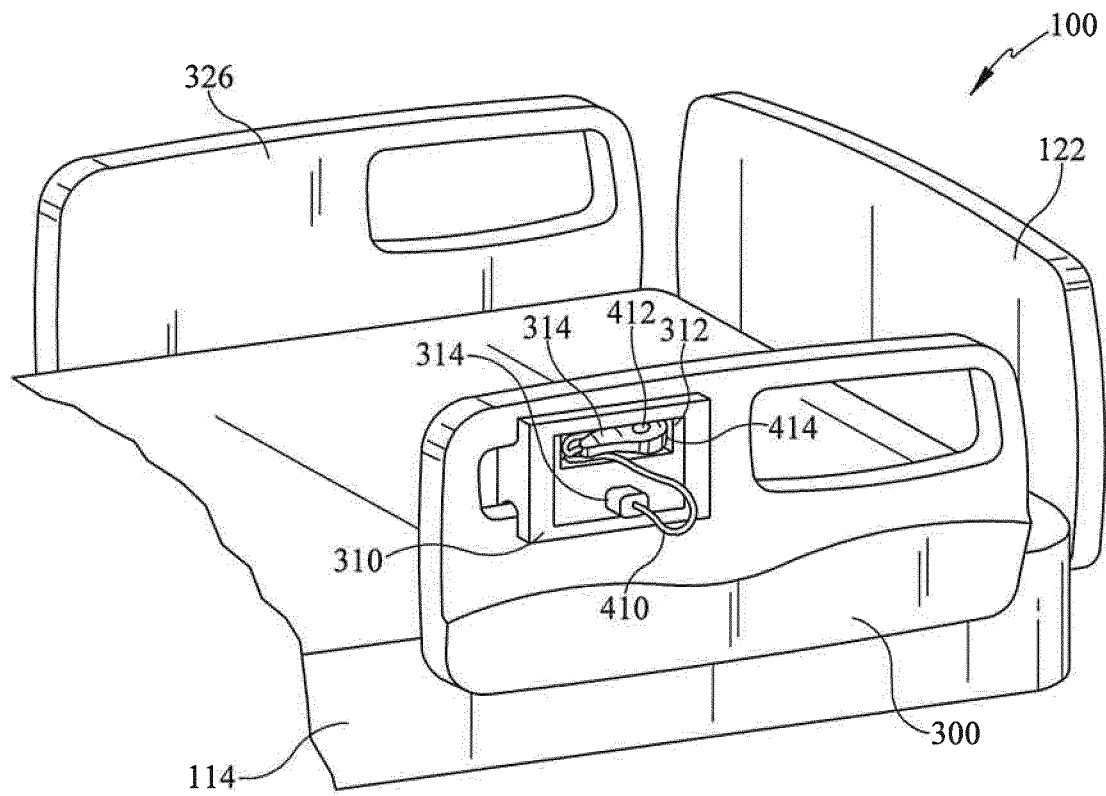


FIG. 3

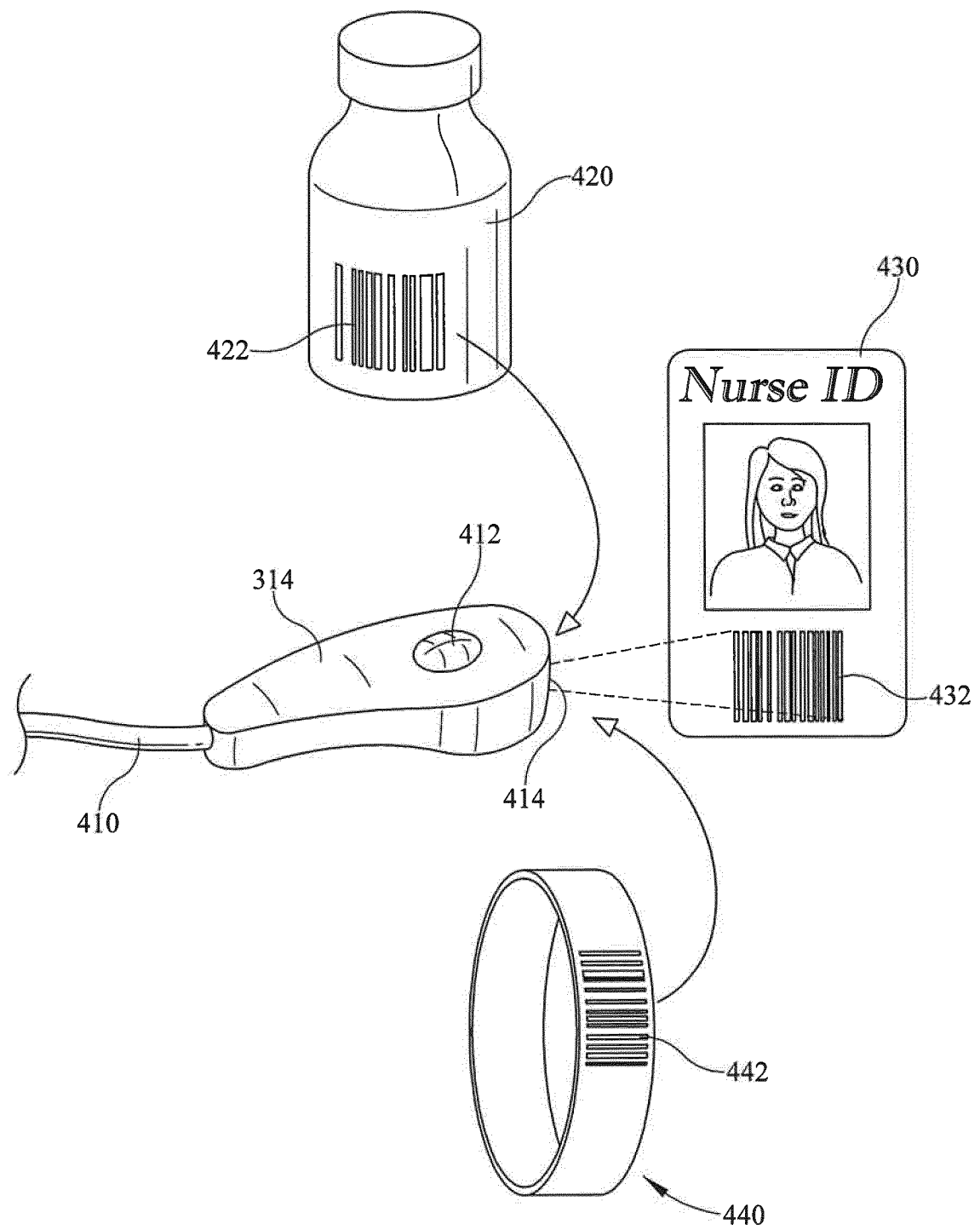


FIG. 4

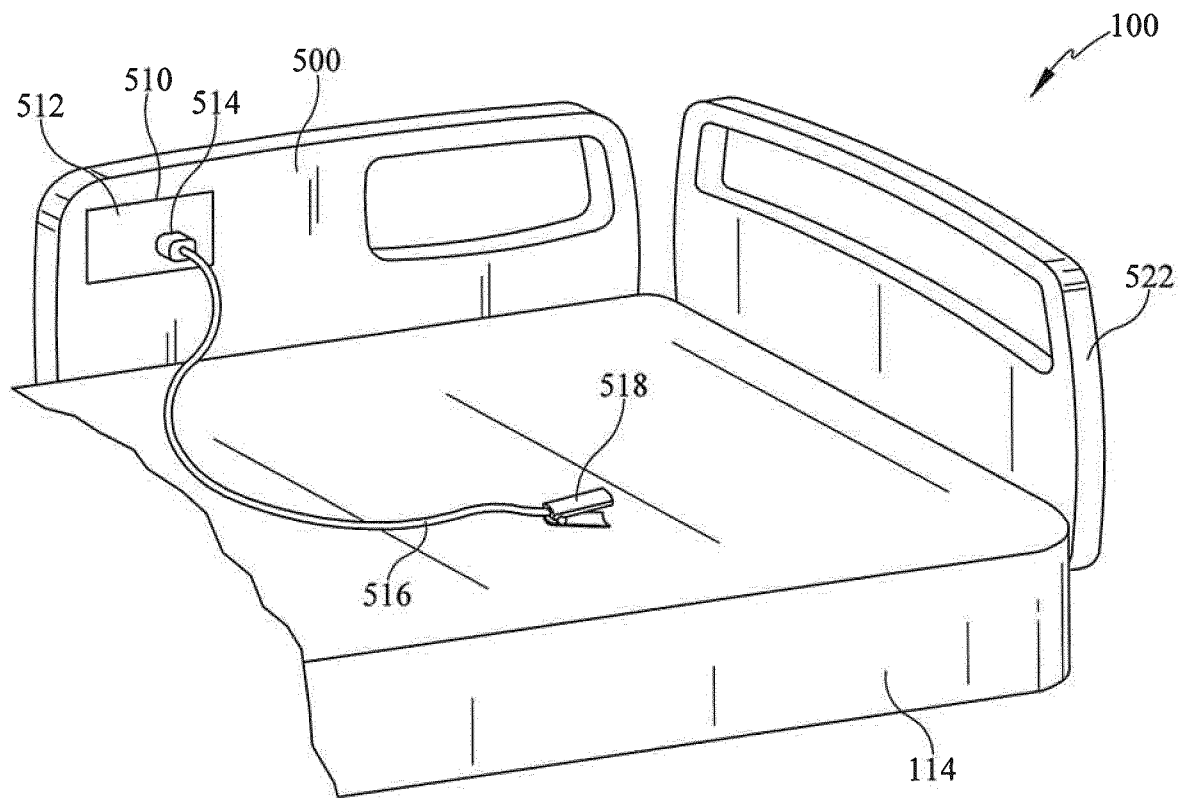


FIG. 5

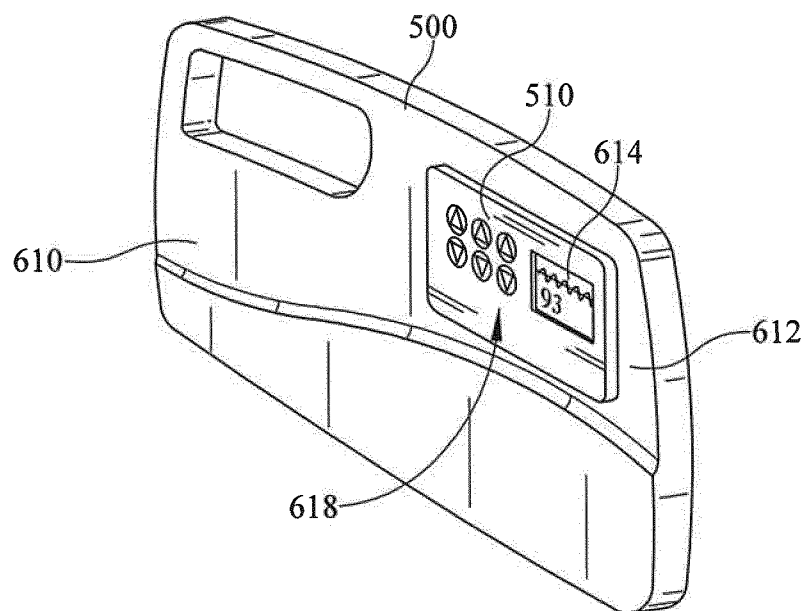


FIG. 6

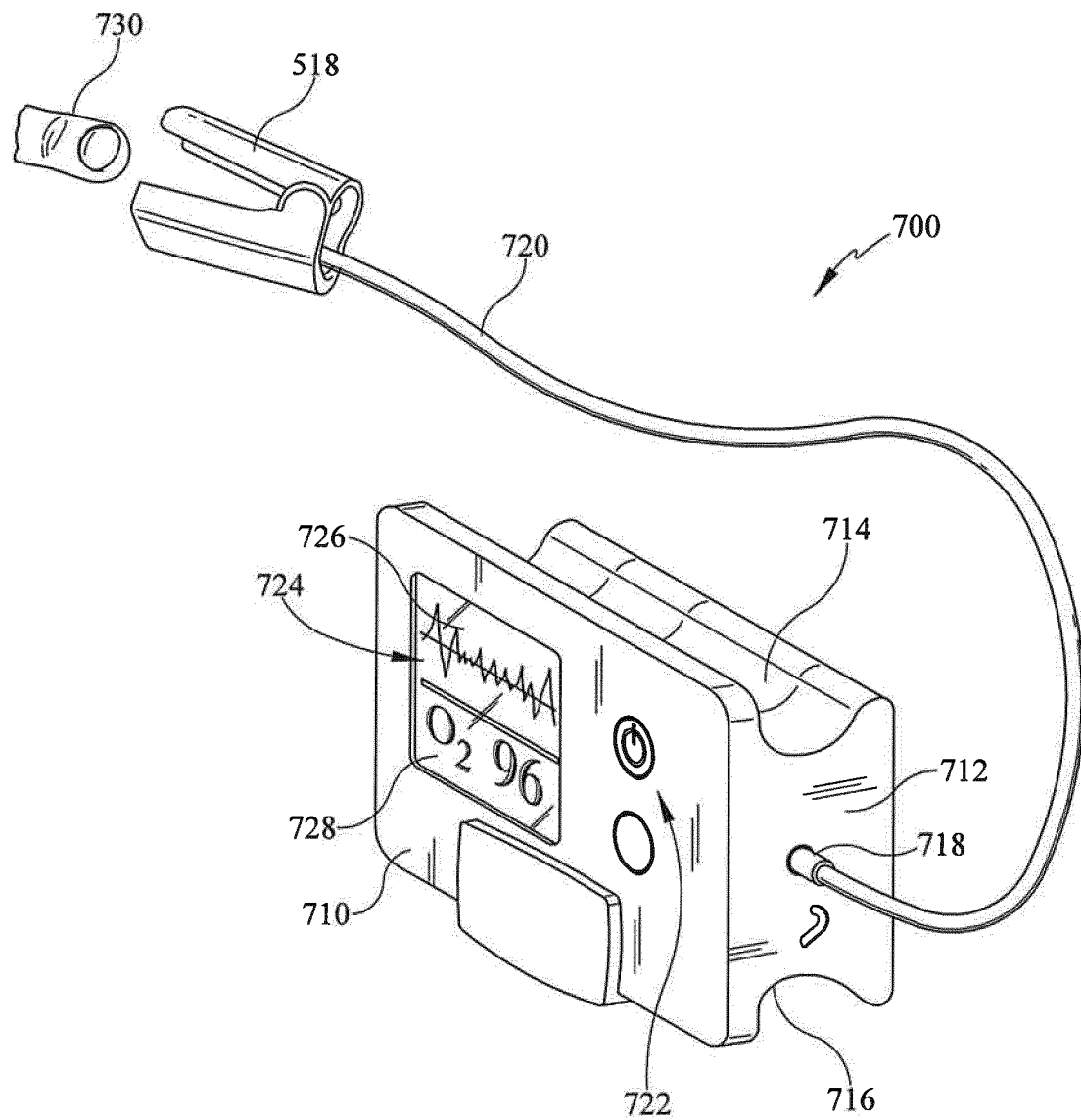


FIG.7

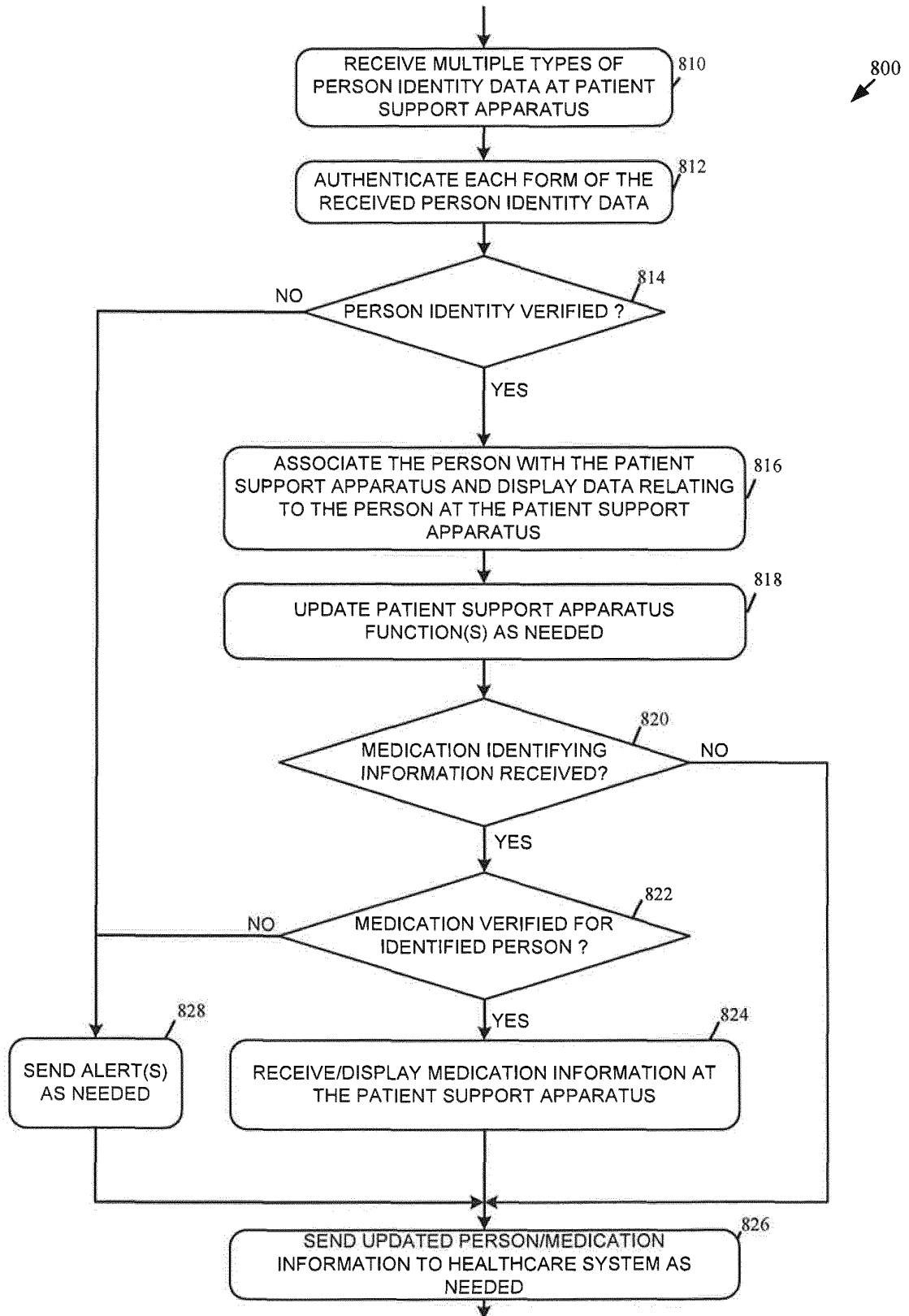


FIG. 8

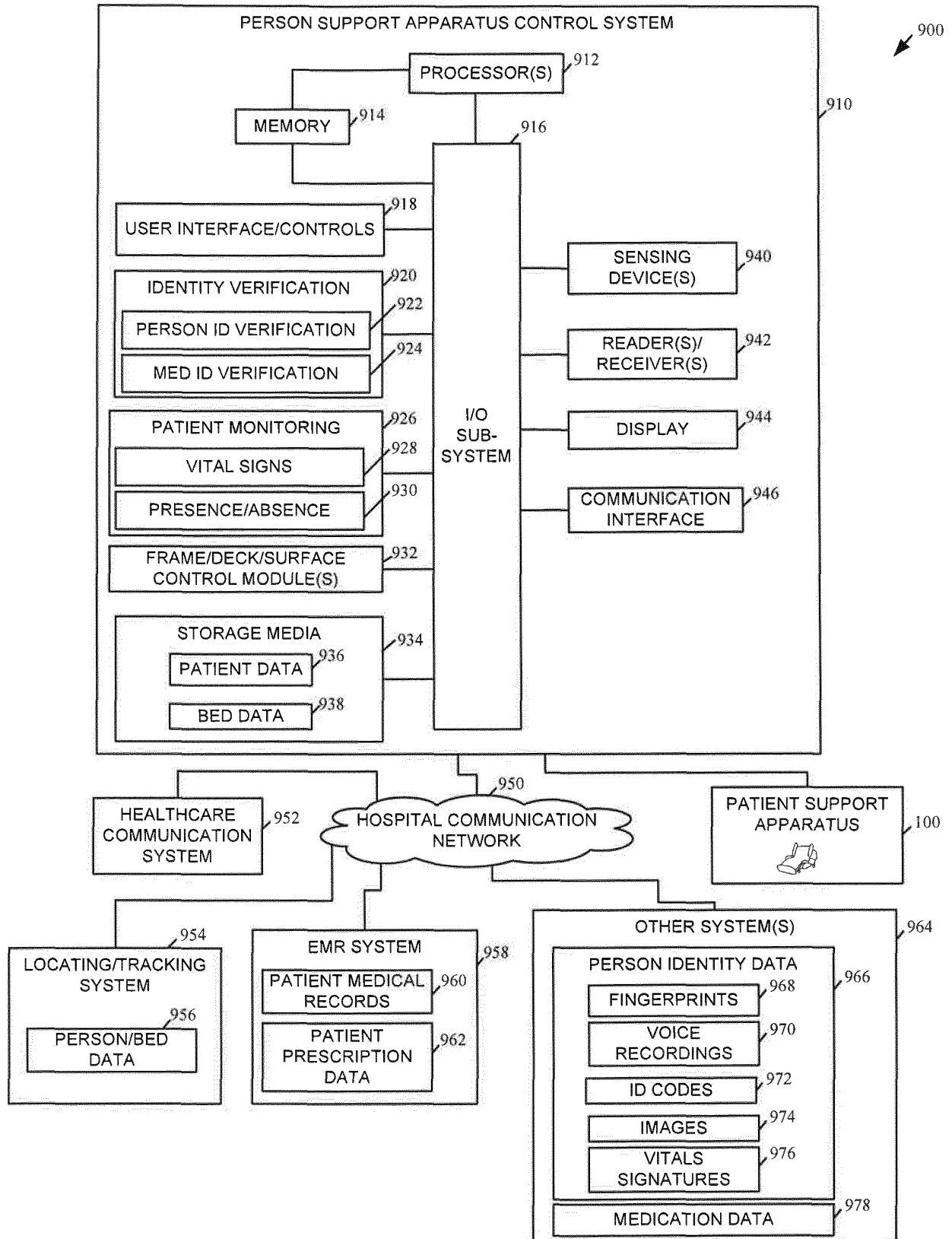


FIG. 9

REFERENCES CITED IN THE DESCRIPTION

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Patent documents cited in the description

- US 20040249673 A [0003]

专利名称(译)	具有冗余身份验证的患者卧榻设备		
公开(公告)号	EP2641577B1	公开(公告)日	2019-11-13
申请号	EP2013160141	申请日	2013-03-20
[标]申请(专利权)人(译)	希尔 - 罗姆服务股份有限公司		
申请(专利权)人(译)	HILL-ROM SERVICES , INC.		
当前申请(专利权)人(译)	HILL-ROM SERVICES , INC.		
[标]发明人	DIXON STEVEN ALAN RIBBLE DAVID LANCE ONEAL TODD P LACK CHRISTOPHER K ALLEN JAMES MAURICE HEIL THOMAS F		
发明人	DIXON, STEVEN ALAN RIBBLE, DAVID LANCE O'NEAL, TODD P. LACK, CHRISTOPHER K. ALLEN, JAMES MAURICE HEIL, THOMAS F.		
IPC分类号	A61G7/018 A61G7/05 A61B5/00 A61B5/1455 A61B90/96 A61B90/98 G16H40/63 G06Q50/22 A61B5/1172		
CPC分类号	A61B5/1172 A61B5/14551 A61B5/7495 A61B90/96 A61B90/98 A61G7/018 A61G7/0507 A61G2203/20 A61G2203/30 A61G2205/10 A61G2205/60 G06F19/3462 G06Q50/22 G16H40/63 A47C21/00 A47C21/08 A61B5/14552		
优先权	61/613961 2012-03-21 US 13/796697 2013-03-12 US		
其他公开文献	EP2641577A2 EP2641577A3		
外部链接	Espacenet		

摘要(译)

一种患者支撑设备，包括一个或多个读取器和/或传感设备，以接收可用于验证人或物体（例如患者，护理人员和药物）身份的数据。感测设备可以包括例如生物特征传感器，生理传感器，条形码读取器，照相机，麦克风或其组合。患者支撑设备可以将这些设备获得的身份验证信息用于多种不同目的，包括例如管理对与患者支撑设备相关联的患者的治疗，药物或治疗的递送。

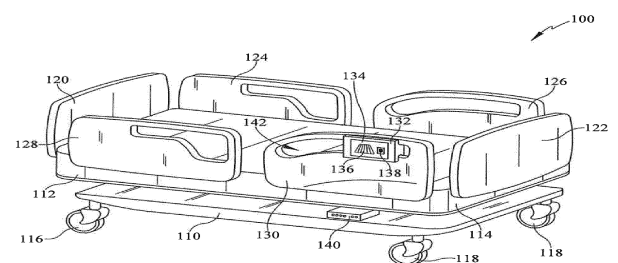


FIG. 1

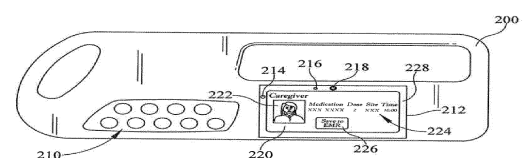


FIG. 2