(54) Title: A SYSTEM AND METHOD FOR REDUCING FALSE ALARMS ASSOCIATED WITH VITAL-SIGNS MONITORING

(57) Abstract: Systems and methods of reducing false alarms associated with a vital-sign monitor are disclosed. One or more data samples of a vital sign of a patient are generated at a first sampling rate in a normal mode of operation. Whether the one or more data samples satisfy an alert condition is determined. An alert mode of operation is entered into if the alert condition is satisfied. One or more additional data samples of the vital sign are generated at a second sampling rate higher than the first sampling rate in the alert mode.

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Published:
— without international search report and to be republished upon receipt of that report (Rule 48.2(g))
A SYSTEM AND METHOD FOR REDUCING FALSE ALARMS ASSOCIATED WITH VITAL-SIGNS MONITORING

Cross-References to Related Applications

[0001] The following applications disclose certain common subject matter with the present application: A Vital-Signs Monitor with Encapsulation Arrangement, docket number 080624-0612; A Vital-Signs Monitor with Spaced Electrodes, docket number 080624-0623; A Vital-Signs Monitor with Spaced Electrodes, docket number 080624-0624; A Vital-Signs Patch Having a Strain Relief, docket number 080624-0624; A Temperature Probe Suitable for Axillary Reading, docket number 080624-0625; System and Method for Monitoring Body Temperature of a Person, docket number 080624-0626; A System and Method for Storing and Forwarding Data from a Vital-Signs Monitor, docket number 080624-0627; A System and Method for Conserving Battery Power in a Patient Monitoring System, docket number 080624-0629; A System and Method for Saving Battery Power in a Patient Monitoring System, docket number 080624-0630; A System and Method for Tracking Vital-Signs Monitor Patches, Docket Number 080624-0631; A System and Method for Location Sensing, Docket Number 080624-0634; all of the listed applications filed on July 27, 2010.

Field

[0002] The present disclosure generally relates to systems and methods of physiological monitoring, and, in particular, relates to systems and methods for reducing false alarms associated with a vital-sign monitor.

Description of the Related Art

[0003] Some of the most basic indicators of a person's health are those physiological measurements that reflect basic body functions and are commonly referred to as a person's "vital signs." The four measurements commonly considered to be vital signs are body
temperature, pulse rate, blood pressure, and respiratory rate. Most or all of these measurements are performed routinely upon a patient when they arrive at a healthcare facility, whether it is a routine visit to their doctor or arrival at an Emergency Room (ER).

Vital signs are frequently taken by a nurse using basic tools including a thermometer to measure body temperature, a sphygmomanometer to measure blood pressure, and a watch to count the number of breaths or the number of heart beats, typically 10 seconds, which is then converted to a "per minute" rate. If a patient's pulse is weak, it may not be possible to detect a pulse by hand and the nurse may use a stethoscope to amplify the sound of the patient's heart beat so that she can count the beats.

When a patient is admitted to a hospital, it is common for vital signs to be measured at regular intervals during the patient's stay to monitor their condition. A typical interval is 4 hours, which leads to the undesirable requirement for a nurse to awaken a patient in the middle of the night to take vital sign measurements.

When a patient is admitted to an ER, it is common for a nurse to do an "triage" assessment of the patient's condition, which will determine how quickly the patient receives treatment. During busy times in an ER, a patient who does not appear to have a life-threatening injury may wait for hours until more serious cases have been treated. While the patient may be reassessed at intervals while awaiting treatment, the patient may not be under observation between these reassessments.

Measuring certain vital signs is normally intrusive at best and difficult to do on a continuous basis. Measurement of body temperature, for example, is commonly done by placing an oral thermometer under the tongue or an infrared thermometer in the ear canal such that the tympanic membrane, which shared blood circulation with the brain, is in the sensor's field of view. Other countries report temperatures made by placing a thermometer under the armpit, referred to as an "axillary" measurement as axilla is the Latin word for armpit. Skin temperature can be measured using a stick-on strip that may contain panels that change color to indicate the temperature of the skin below the strip.
Measurement of respiration is easy for a nurse to do, but relatively complicated for equipment to achieve. A method of automatically measuring respiration is to encircle the upper torso with a flexible band that can detect the physical expansion of the rib cage, when a patient inhales. An alternate technique is to measure a high-frequency electrical impedance between two electrodes placed on the torso and detect the change in impedance created when the lungs fill with air. The electrodes are typically placed on opposite sides of one or both lungs, resulting in placement on the front and back or on the left and right sides of the torso, commonly alone or adhesive electrodes connected by wires or by using a torso band with multiple electrodes in the strap.

Measurement of pulse is also relatively easy for a nurse to do and intrusive for equipment to achieve. A common automatic method of measuring a pulse is to use an electrocardiograph (ECG or EKG) to detect the electrical activity of the heart. An EKG machine may use up to six electrodes placed at defined points on the body to detect various signals associated with the heart function. Another common piece of equipment is simply called an "heart rate monitor." Widely sold for use in exercise and training, heart rate monitors commonly consist of a torso band, in which are embedded two electrodes on the skin and a small electronics package. Such heart rate monitors can communicate wirelessly to other equipment such as a small device that is worn like a wrist watch, and that can transfer data wirelessly to a computer.

Nurses are expected to provide complete care to an assigned number of patients. The workload of a typical nurse is increasing, driven by a combination of a continuing shortage of nurses, an increase in the number of formal procedures that must be followed, and an expectation of increased documentation. Replacing the manual measurement and logging of vital signs with a system that measures and records vital signs would enable a nurse to spend more time on other activities and avoid the potential for error that is inherent in any manual procedure.
SUMMARY

[0011] For some or all of the reasons listed above, there is a need for a hospital to be able to continuously monitor its patients in different settings within the hospital. In addition, it is desirable for this monitoring to be done with limited interference with a patient's mobility or interfering with their other activities.

[0012] Embodiments of the patient monitoring system disclosed herein measure certain vital-sign readings of a patient, which include respiratory rate, pulse rate, and body temperature, on a regular basis and compare these measurements to preset limits.

[0013] In addition, certain embodiments of the patient monitoring system disclosed herein can send alarm notifications to a healthcare system if the vital-sign readings satisfy an alarm condition. It is desirable to reduce or eliminate false positives in the alarm notifications so as to avoid unnecessary trips or other actions of a healthcare provider that such false positives can cause.

[0014] In one aspect of the present disclosure, a method of reducing false alarms associated with vital-sign monitor is disclosed. The method can comprise generating one or more data samples of a patient at a first sampling rate in a normal mode of operation. The method can further comprise determining whether the one or more data samples satisfies an alert condition. The method can further comprise entering an alert mode of operation if the alert condition is satisfied. The method can further comprise generating one or more additional data samples of the vital sign at a second sampling rate higher than the first sampling rate in the alert mode.

[0015] In one aspect of the present disclosure, a vital sign monitoring system is disclosed. The system can comprise a vital-sign monitor configured to monitor one or more vital signs of a patient. The system can further comprise a surveillance server configured to gather data relating to the one or more vital signs of the patient from the vital-sign monitor. The vital-sign monitor can be further configured to generate one or more data samples of a vital sign of a patient at a first sampling rate in a normal mode of operation, determine whether the one or more data samples satisfy an alert condition, enter an alert mode if the alert condition is satisfied, and generate one or more data samples of a vital sign of a patient at a second sampling rate in the alert mode.
satisfied, and generate one or more additional data samples of the vital sign at a second
sampling rate higher than the first sampling rate in the alert mode.

It is understood that other configurations of the subject technology will become
readily apparent to those skilled in the art from the following detailed description, wherein
various configurations of the subject technology are shown and described by way of
illustration. As will be realized, the subject technology is capable of other and different
configurations and its several details are capable of modification in various other respects, all
configurations and its several details are capable of modification in various other respects, all
without departing from the scope of the subject technology. Accordingly, the drawings and
detailed description are to be regarded as illustrative in nature and not as restrictive,
detailed description are to be regarded as illustrative in nature and not as restrictive.

BRIEF DESCRIPTION OF THE DRAWINGS

The accompanying drawings, which are included to provide further understanding
and constitute a part of this specification, illustrate disclosed embodiments and
together with the description serve to explain the principles of the disclosed embodiments. In the drawings:

FIG. 1 is a diagram illustrating an exemplary embodiment of a patient monitoring
system according to certain aspects of the present disclosure.

FIG. 2A is a perspective view of the vital sign monitor patch shown in FIG. 1
according to certain aspects of the present disclosure.

FIG. 2B is a cross-sectional view of the vital sign patch shown in FIGS. 1 and 2A
according to certain aspects of the present disclosure.

FIG. 2C is a functional block diagram illustrating exemplary electronic and sensor
components of the monitor patch of FIG. 1 according to certain aspects of the present
disclosure.

FIG. 3A is a functional schematic diagram of an embodiment of the bridge according
to certain aspects of the present disclosure.
Continuous monitoring of patients in a hospital is desirable at least to ensure that patients do not suffer an unnoticed sudden deterioration in their condition or a secondary injury during their stay in the hospital. It is impractical to provide continuous monitoring by a clinician and cumbersome to connect sensors to a patient, which are then connected to a fixed monitoring instrument by wires. Furthermore, systems that sound an alarm when the measured value exceeds a threshold value may sound alarms so often and in situations that are not truly serious that such alarms are ignored by clinicians.

Measuring vital signs is difficult to do on a continuous basis. Accurate measurement of cardiac pulse, for example, can be done using an electrocardiograph (ECG or EKG) to detect the electrical activity of the heart. An EKG machine may use up to 10 electrodes to detect the electrical activity of the heart.
placed at various points on the body to detect various signals associated with the heart.

Another common piece of equipment is termed a "heart rate monitor." Widely sold for use in exercise and physical training, heart rate monitors may consist of a torso band in which are embedded two electrodes held against the skin and a small electronics package. Such heart rate monitors can communicate wirelessly to other equipment such as a small device that is worn like a wrist watch that can transfer data wirelessly to a personal computer (PC).

Certain exemplary embodiments of the present disclosure include a system that

construction of the vital-signs monitor patch is described according to certain aspects of the present disclosure. As the patch may be worn continuously for a period of time that may be several days, as is described in the following disclosure, it is desirable to encapsulate the components of the patch such that the patient can bathe or shower and engage in their normal activities without degradation of the patch function. An exemplary configuration of the construction of the patch to provide a hermetically sealed enclosure about the electronics is disclosed.

In the following detailed description, numerous specific details are set forth to provide a full understanding of the present disclosure. It will be apparent, however, to one ordinarily skilled in the art that embodiments of the present disclosure may be practiced without some of the specific details. In other instances, well-known structures and techniques have not been shown in detail so as not to obscure the disclosure.

FIG. 1 discloses a vital sign monitoring system according to certain embodiments of the present disclosure. The vital sign monitoring system 12 includes vital-signs monitor 19, bridge 90, and surveillance server 80 that can send messages or interact with peripheral devices exemplified by mobile device 90 and workstation 100.
Monitor patch 20 resembles a large adhesive bandage and is applied to a patient when in use. It is preferable to apply the monitor patch 20 to the upper chest of the patient, although other locations may be employed. The monitor patch 20 incorporates one or more electrodes (not shown) that are in contact with the skin of the patient to measure vital signs such as cardiac pulse rate and respiration rate. Monitor patch 20 also may include other sensors such as an accelerometer or a temperature sensor to measure other characteristics associated with the patient. Monitor patch 20 also includes a wireless transmitter that can both transmit and receive signals. This transmitter is preferably a short-range, low-power radio frequency (RF) device operating in one of the industrial, scientific, and medical (ISM) radio bands. One ISM band in the United States (US) is, for example, centered at 915 MHz. An example of an equivalent band in the European Union (EU) is centered at 868 MHz. Other frequencies of operation may be possible dependent upon local regulations and interference from other wireless devices.

Surveillance server 60 may be a standard computer server connected to the hospital communication network and preferably located in the hospital data center or computer room, although other locations may be employed. The server 60 stores and processes signals related to the operation of the patient monitoring system 112 disclosed herein, including the association of individual monitor patches 20 with patients and measurement signals received from multiple monitor patches 20. Hence, although only a single patient and monitor patch 20 are depicted in FIG. 1, the server 60 is able to monitor the monitor patches 20 for multiple patients.

Bridge 40 is a device that connects, or "bridges", between monitor patch 20 and server 60. Bridge 40 communicates with monitor patch 20 over communication link 30 and server 60. Bridge 40 communicates with monitor patch 20 over communication link 30 operating, in these exemplary embodiments, at approximately 915 MHz and at a power level operating, in these exemplary embodiments, at approximately 915 MHz and at a power level that enables communication link 30 to function up to a distance of approximately 3 meters. It is preferable to place a bridge 40 in each room and at regular intervals along hallways of the healthcare facility where it is desired to provide the ability to communicate with monitor patches 20. Bridge 40 also is able to communicate with server 60 over network link 50 using any of a variety of computer communication systems including "hardwired" and "wireless" Ethernet using protocols such as 802.11(a/b/g/i/n) or 802.3af. As the communication protocols of Ethernet using protocols such as 802.11(a/b/g) or 802.3af. As the communication protocols of
communication link 30 and network link 50 may be very different, bridge 40 provides data
buffering and, therefore, conversion to enable bidirectional signal transmission between
monitor patch 20 and server 60.

[0038] While the embodiments illustrated by FIG. 1 employ a bridge 20 to provide
communication link between the monitor patch 20 and the server 60, in certain alternative
embodiments, the monitor patch 20 may engage in direct wireless communication with the
bridge 40. In such alternative embodiments, the server 60 itself or a wireless modem
can be connected to the server 60 to receive data from the monitor patch 20.

[0039] In use, a monitor patch 20 is applied to a patient 10 by a clinician when it is desirable
to continuously monitor basic vital signs of patient 10 while patient 10 is in this
embodiment in a hospital. Monitor patch 20 is intended to remain attached to patient 10 for
an extended period of time—such as, for example, up to 5 days in certain embodiments, limited by the
battery life of the monitor patch 20. In some embodiments, monitor patch 20 is disposable when
removed from patient 10.

[0040] Server 60 executes analytical protocols on the measurement data that it receives from
monitor patch 20 and provides this information to clinicians through external workstations
11, preferably personal computers (PCs), over the hospital network 70. Server 60 may also
send messages to mobile devices 90, such as cell phones or pagers, over a mobile device link
80 if a measurement signal exceeds specified parameters. Mobile device link 80 may include
60 if a measurement signal exceeds specified parameters. Mobile device link 80 may include
the hospital network 70 and internal or external wireless communication systems that are
the hospital network 70 and internal or external wireless communication systems that are
capable of receiving mobile communications that can be received by mobile devices 90.

[0041] FIG. 2A is a perspective view of the vital-signs monitor patch 20 shown in FIG. 1 according to certain aspects of the present disclosure. In the illustrated embodiment, the monitor patch 20 includes component carrier 23 comprising a central segment 21 and side
segments 22 on opposing sides of the central segment 21. In certain embodiments, the
central segment 21 is substantially rigid and includes a circuit assembly (24, FIG. 2B) having
electronic components and battery mounted to a rigid printed circuit board (PCB). The side
segments 22 are flexible and include a flexible conductive circuit (26, FIG. 2B) that connects
connections.
the circuit assembly 24 to electrodes 28 disposed at each end of the monitor patch 20, with the circuit assembly 24, the electrodes 28 disposed at each end of the monitor patch 20, with side segment 22 on the right shown as being bent upwards for purposes of illustration to make one of the electrodes 28 visible in this view.

[0042] FIG. 2B is a cross-sectional view of the vital-signs patch 20 shown in FIGS. 1 and 2A according to certain aspects of the present disclosure. The circuit assembly 24 and flexible conductive circuit 26 described above can be seen herein. The flexible conductive circuit 26 operably connects the circuit assembly 24 to the electrodes 28. Top and bottom layers 23 and 27 form a housing 25 that encapsulate circuit assembly 28 to provide a water and particulate barrier as well as mechanical protection. There are sealing areas on layers 23 and 27 that encircles circuit assembly 28 and is visible in the cross-section view of FIG. 2B. Layers 23 and 27 are sealed to each other in this area to form a substantially hermetic seal. Within the context of certain aspects of the present disclosure, the term "hermetic" implies that the rate of transmission of moisture through the seal is substantially the same as through the material of the layers that are sealed to each other, and further implies that the size of particulates that can pass through the seal are below the size that can have a significant effect on circuit assembly 24. Flexible conductive circuit 26 passes through portions of sealing areas 29 and the seal between layers 23 and 27 is maintained by sealing of layers 23 and 27 to flexible circuit assembly 28. The layers 23 and 27 are thin and flexible, as is the flexible conductive circuit 26, allowing the side segment 22 of the monitor patch 20 between the electrodes 28 and the circuit assembly 24 to bend as shown in FIG. 2A.

[0043] FIG. 2C is a functional block diagram 200 illustrating exemplary electronic and sensor components of the monitor patch 20 of FIG. 1 according to certain aspects of the present disclosure. The block diagram 200 shows a processing and sensor interface module present disclosure. The block diagram 200 shows a processing and sensor interface module 201 and external sensors 232, 234 connected to the module 201. In the illustrated example, the module 201 includes a processor 202, a wireless transceiver 207 having a receiver 206 and a transmitter 209, a memory 210, a first sensor interface 211, a second sensor interface 213, a third sensor interface 215, and an internal sensor 236 connected to the third sensor interface 215. The first and second sensor interfaces 211 and 213 are connected to the first and second external sensors 232, 234 via first and second connection ports 222, 224, and second external sensors 232, 234 via first and second connection ports 222, 224.
respectively. In certain embodiments, some or all of the aforementioned components of the module 201 and other components are mounted on a PCB.

[0044] Each of the sensor interfaces 212, 214, 216 can include one or more electronic components that are configured to generate an excitation signal or provide DC power for the sensor that the interface is connected to and/or to condition and digitize a sensor signal from the sensor. For example, the sensor interface can include a signal generator for generating an excitation signal or a voltage regulator for providing DC power to the sensor. The sensor excitation signal or a voltage regulator for providing DC power to the sensor. The sensor interface can further include an amplifier for amplifying a sensor signal from the sensor and/or an analog-to-digital converter for digitizing the amplified sensor signal. The sensor interface can further include an analog-to-digital converter for digitizing the amplified sensor signal. The sensor interface can further include a filter (e.g., a low-pass or bandpass filter) for filtering out spurious noises (e.g., 60Hz noise pickup).

[0045] The processor 202 is configured to send and receive data (e.g., digitized signal or control data) to and from the sensor interfaces 212, 214, 216 via a bus 204, which can include one or more wires or traces on the PCB. Although a bus communication topology is used in this embodiment, some or all communication between discrete components can also be implemented as direct links without departing from the scope of the present disclosure. For example, the processor 202 may send data representative of an excitation signal to the sensor excitation signal generator, inside the sensor interface and receive data representative of the sensor signal from the sensor interface over either a bus or direct data links between processor 202 and each of sensor interface 212, 214, and 216.

[0046] The processor 202 is also capable of communication with the receiver 206 and the transmitter 209 of the wireless transceiver 207 via the bus 204. For example, the processor transmitter 209 of the wireless transceiver 207 via the bus 204. For example, the processor transmitter 202 using the transmitter and receiver 209, 206 can transmit and receive data to and from the 202 using the transmitter and receiver 209, 206 can transmit and receive data to and from the bridge 40. In certain embodiments, the transmitter 209 includes one or more of an RF signal generator (e.g., an oscillator), a modulator (a mixer), and a transmitting antenna; and the receiver 206 includes a demodulator (a mixer) and a receiving antenna which may or may not be the same as the transmitting antenna. In some embodiments, the transmitter 209 may include a digital-to-analog converter configured to receive data from the processor 202 and to generate a base signal; and/or the receiver 206 may include an analog-to-digital converter to generate a base signal; and/or the receiver 206 may include an analog-to-digital converter.
configured to digitize a demodulated base signal and output a stream of digitized data to the processor 202.

[0047] The processor 202 may include a general-purpose processor or a specific-purpose processor for executing instructions and may further include a memory 219, such as a random or non-volatile memory, for storing data and/or instructions for software programs, volatile or non-volatile memory, for storing data and/or instructions for software programs. The instructions, which may be stored in a memory 219 and/or 210, may be executed by the processor 202 to control and manage the wireless transceiver 207, the sensor interfaces 212, 214, 216, as well as provide other communication and processing functions.

[0048] The processor 202 may be a general-purpose microprocessor, a microcontroller, a Digital Signal Processor (DSP), an Application Specific Integrated Circuit (ASIC), a Field Programmable Gate Array (FPGA), a Programmable Logic Device (PLD), a state machine, gated logic, discrete hardware components, or any other suitable device or a combination of devices that can perform calculations or other manipulations of information.

[0049] Information, such as program instructions, data representative of sensor readings, preset/ alarm conditions, thresholds, limits, may be stored in a computer or processor-readable medium, such as memory, internal to the processor 202 (e.g., the memory 219) or a memory external to the processor 202 (e.g., the memory 210), such as a Random Access Memory (RAM), a flash memory, a Read-Only Memory (ROM), a Programmable Read-Only Memory (PROM), an Erasable PROM (EPROM), registers, a hard disk, a removable disk, or any other suitable storage device.

[0050] In certain embodiments, the internal sensor 236 can be one or more sensors configured to measure certain properties of the processing and sensor interface module 201, such as a board temperature sensor thermally coupled to a PCB. In other embodiments, the internal sensor 236 can be one or more sensors configured to measure certain properties of the patient 10, such as a motion sensor (e.g., an accelerometer) for measuring the patient’s motion.

[0051] The external sensors 232, 234 can include sensors and sensing arrangements that are configured to produce a signal representative of one or more vital signs of the patient to
which the monitor patch 20 is attached. For example, the first external sensor 232 can be a set of sensing electrodes that are affixed to an exterior surface of the monitor patch 20 and configured to be in contact with the patient for measuring the patient's respiratory rate, and the second external sensor 234 can include a temperature sensing element (e.g., a thermocouple or a thermometer) affixed, either directly or via an interposing layer, to skin of the patient 10 for measuring the patient's body temperature. In other embodiments, one or more of the external sensors 232, 234 and one or more additional external sensors can measure more of the external sensors 232, 234 or one or more additional external sensors can measure other vital signs of the patient, such as blood pressure and pulse rate. Other vital signs of the patient, such as blood pressure and pulse rate.

[0052] FIG. 3A is a functional block diagram illustrating exemplary electronic components of bridge 40 of FIG. 1 according to one aspect of the subject disclosure. Bridge 40 includes a processor 310, a transmitter 324, a receiver 322, and a wireless interface 352 connected to the processor 310 via a bus 314. In some embodiments, the processor 310 is configured to send data to and receive data from the transmitter 324 and the receiver 322 and the wireless interface 352 and the wired interface 354 of network interface 350 via bus 314. In certain embodiments, transmitters and receivers may include a radio frequency signal generator (oscillator), a modulator, and a transmitting antenna, and the receivers may include a demodulator and an antenna which may or may not be the same as the transmitting antenna of the radio. In some embodiments, transmitters and receivers may include a digital-to-analog converter configured to convert data received from processor 310 to generate a base signal, which may include analog-to-digital converters configured to convert a demodulated base signal and sent a digitized data stream to processor 310.

[0053] Processor 310 may include a general-purpose processor or a specific-purpose processor for executing instructions and may further include a memory 312, such as a volatile or non-volatile memory, for storing data and/or instructions for software programs. The instructions, which may be stored in memories 312 or 340, may be executed by the processor 310.
processor 310 to control and manage the transceivers 320, 330, and 350 as well as provide other communication and processing functions.

[0055] Processor 310 may be a general-purpose microprocessor, a microcontroller, a Digital Signal Processor (DSP), an Application Specific Integrated Circuit (ASIC), a Field-Programmable Gate Array (FPGA), a Programmable Logic Device (PLD), a controller, a state machine, a gated logic, discrete hardware components, or any other suitable device or a combination of devices that can perform calculations or other manipulations of information.

[0056] Information such as data representative of sensor readings may be stored in memory such as data representative of sensor readings may be stored in memory 312, 320, 330, 340, or 350. Memory 312, 320, 330, 340, or 350 may be a Random Access Memory (RAM), a Read Memory, Read Only Memory (ROM), a Programmable Read-Only Memory (PROM), a Programmable Read-Only Memory (EPROM), a Flash Memory, a Read-Only Memory (ROM), a Hard Disk, a Removable Disk, a Solid State Memory (SSD), a suitable storage device.

[0057] Memory 312, 320, 330, 340 can also store a list or a database of established communication links and their corresponding characteristics (e.g., signal levels) between the bridge(s) and its related monitor patches. In the illustrated example of FIG. 3 A, the memory 340, external to the processor 310, includes such a database 342; alternatively, the memory 320, internal to the processor 310, may include such a database.

[0058] FIG. 3 B is a functional block diagram illustrating exemplary electronic components of server 60 of FIG. 1 according to one aspect of the subject disclosure. Server 60 includes a processor 360, memory 370, display 380, and network interface 390 having a wireless interface 392, and a wired interface 394. Processor 360 may include a general-purpose processor or a specific-purpose processor for executing instructions and may further include a memory 362, such as a volatile or non-volatile memory, for storing data and/or instructions. Memory 362 may include a volatile or non-volatile memory, for storing data and/or instructions for software programs. The instructions, which may be stored in memory 362 or other memory, may be executed by the processor 360 to control and manage the wireless and wired network interfaces 392, 394 as well as provide other communication and processing functions.
Processor 360 may be a general-purpose microprocessor, a microcontroller, a Digital Signal Processor (DSP), an Application Specific Integrated Circuit (ASIC), a Field Programmable Gate Array (FPGA), a Programmable Logic Device (PLD), a controller, a state machine, gated logic, discrete hardware components, or any other suitable device or a combination of devices that can perform calculations or other manipulations of information.

Information such as data representitive of sensor readings may be stored in memory. Information such as data representative of sensor readings may be stored in memory 362 or 370 can also store a database of communication links and their corresponding characteristics (e.g., signal levels) between monitor patches 20 and bridges 40. In the illustrated example of FIG. 3B, the memory 370 external to the processor 360 includes such a database 372; alternatively, the memory 362 internal to the processor 360 may include such a database.

As indicated above, with respect to FIG. 2C, certain embodiments of the monitor patch 20 are configured to operate, with external sensors that are in turn configured to produce a signal representative of one or more vital signs of the patient to whom the monitor patch 20 is attached. For example, the second external sensor 234 can be a temperature probe that includes a temperature sensing element (e.g., a thermocouple or thermistor) that is configured to measure the temperature of the patient's body temperature. FIG. 4A is a diagram depicting a patient 10 wearing a temperature monitoring system 20A comprising a monitor patch 20 and a temperature probe 400 that is configured to measure body temperature of the patient 10. In the illustrated example, the temperature probe 400 is configured for auxiliary temperature sensing of the patient 10. The temperature probe 400 can be configured for auxiliary temperature sensing of the patient 10 to whom the monitor patch 20 is attached. The monitor patch 20 is configured to attach on the chest of the patient 10, with a sensing portion of the temperature probe 400 retained in the axilla 12 of the patient 10 during body temperature monitoring.
FIG. 4B is a diagram providing an enlarged view of the monitoring system 20A depicted in FIG. 4A according to certain aspects of the present disclosure. As indicated above, the monitor patch 20 is attached to the chest 1 of the patient 10, via, e.g., an adhesive backing (not shown). The temperature probe 400 has a proximal end 401 and a distal end 402 and includes a wiring portion 410, a body connection portion 430, and a sensing portion 420 disposed between the wiring and body connection portions 410, 430. The proximal end 401 of the temperature probe 400 is connected to the monitor patch 20 at its connection port 410 of the temperature probe 400 is connected to the monitor patch 20 at its connection port 410. In certain embodiments, the proximal end 401 of the temperature probe 400 is removably attached (e.g., plugged) to the monitor patch 20. In other embodiments, the proximal end 401 is fixedly attached (e.g., epoxied or fused) to the monitor patch 20.

The sensing portion 420 of the temperature probe 400 is configured for placement between the axilla 12 of the patient 10 and includes a temperature sensing element (e.g., 234A-D of FIGS. 6A-B) and 234E-F of FIGS. 6A-B). The wiring portion 410 of the temperature probe 400 includes one or more electrical conductors (512, 514 of FIGS. 5A-D and of FIGS. 6A-B) for carrying a signal responsive to a change in body temperature of the patient 10 between the temperature sensing element 234 and the monitor patch 20. In the illustrated example, the wiring portion 410 includes a flexible cable comprising a tubing and electrical conductors (e.g., a pair of twisted copper wires) placed within the tubing. The wiring portion 410 includes a coiled section 414 acting as a spring to take up any slack in the cable so as to accommodate patients of different sizes. In the illustrated example, the monitoring system further includes an adhesive element 416 (e.g., a tape) configured to attach the wiring portion 410 of the cable to the patient's body, e.g., at a point between the chest and the arm pit 2 of the patient.

The body connection portion 430 has one end connected to the sensing portion 420 and is configured to be attached to another body portion of the patient 10 such that the proximal end 431 of the body connection portion 430 can be retained within the axilla 12 of the patient 10. In the illustrated example, such attachment is achieved via an adhesive element 426 (e.g., a tape) coupled to the distal end of the body connection portion 430. The coupled adhesive element 426 is then attached to a second body portion 13 (e.g., the back of the patient's arm) of the patient 10, attached to a second body portion 13 (e.g., the back of the patient's arm) of the patient 10.
A multitude of modifications and additions to the illustrated embodiment of FIG. 4B are possible without departing from the scope of the disclosure. For example, the body connection portion 430 of the temperature probe 400 can include one or more coiled sections so that both the distal end of the wiring portion 416 can act as a spring similar to the coiled section 411 of the wiring portion 412. The adhesive element 416 may be coupled to the body connection portion 430 at a point different than the distal end of the body connection portion 430. In certain embodiments, entirely different means of attaching the body connection portion 430 to the patient's body may be used. For example, the body connection portion 430 may itself be in the form of an adhesive tape that can stick to the body of the patient 10 or may include an elastic loop (e.g., a rubber band) to be placed around the patient's arm.

While the temperature probe 400 in the illustrated embodiments of FIGS. 4A-B is shown to be operatively coupled to a vital sign monitor patch worn by the patient 10, the temperature probe 400 may be alternatively operatively coupled to other types of monitoring devices such as a stationary monitoring unit located near the patient's hospital bed. Such a stationary monitoring unit can take readings of the patient's body temperature based on a signal from the temperature probe 400 and send the temperature readings to a surveillance server via a wired or wireless connection and make other decisions such as providing an indication of an alarm condition (e.g., a high body temperature condition or a loss of thermal contact between the temperature probe and the patient).

An important concept in the vital-sign monitoring system of the present disclosure is the "measurement interval," or "sampling rate," which is an inverse of the measurement interval. In certain embodiments, the monitor patch generates data samples of a vital sign or vital signs of a patient at a predetermined measurement interval or sampling rate and transmits the data samples to a surveillance server, either directly or via a bridge. In some embodiments, the default measurement interval is 10 minutes, providing, for example, about 5 days of operation for a disposable monitor patch. Measurement intervals can be set at the hospital or care unit level and can be set to a value in a range of between about 2 and 30 minutes. Because vital-sign measurements and transmissions account for a significant portion of power consumption for a monitor patch, a measurement-interval setting directly impacts the life expectancy of the monitor patch, along with the power consumption for the monitor patch.
In certain embodiments, the monitoring system can dynamically adjust the measurement interval (hence, the sampling rate) depending on values of the vital-sign or vital signs of interest. For example, under normal patient conditions, the monitor patch generates data samples at a first (normal) sampling rate. When an alert condition is detected (e.g., a vital-sign value exceeding and staying above a threshold limit), the monitor patch can automatically change from the first (normal) sampling rate (e.g., every 10 minutes) to a second (alert) sampling rate (e.g., every 2 minutes). The monitor patch returns to the first (normal) sampling rate when the vital-sign value returns to a value below the threshold limit.

FIG. 5 depicts a graph 500 representing a series of data samples (solid dots) corresponding to vital-sign values of a patient generated by a monitor patch in a certain region 214 of FIG. 2C. FIGS. 4A and 4B according to certain aspects of the present disclosure. The vital-sign values can represent, for example, body temperature or heart rate of the patient. The graph 500 shows two threshold limits: a first threshold limit 501 (the "yellow limit") and a second threshold limit 502 (the "red limit"). The graph 500 is divided into three regions: a first region 504 corresponding to the monitor patch 20 operating in a normal mode, a second region 506 corresponding to the monitor patch 20 operating in an alert mode, and a third region 508 corresponding to the monitor patch 20 reverting to its normal mode of operation.

FIG. 6 is a flowchart illustrating an exemplary vital-sign monitoring process 600 from the perspective of a vital-sign monitor patch according to certain aspects of the present disclosure. For the purposes of illustration only, without any intent to limit the scope of the disclosure, the process 600 will be described with reference to the graph 500 of FIG. 5. The process 600 begins at start state 601 and proceeds to operation 610 in which the monitor patch 20 generates a data sample of a vital sign of a patient at a first sampling rate (e.g., after a first measurement interval from a previous data sample). At this stage, the 'monitor patch 20 is assumed to be in a normal mode of operation 606 corresponding to the first region 504 of FIG. 5. In the illustrated example, the first sampling rate is one data sample per every 8 minutes 'corresponding to a first (normal) measurement interval of 8 minutes.' In certain embodiments, an analog-to-digital converter in a sensor interface (e.g., FIG. 2C) of the monitor patch 20 converts a sensor signal.
indicative of a vital sign (e.g., a patient's respiration, heart beat, or body temperature) into digital representations of the sensor signal. A processor (e.g., 202 of FIG. 2) in the monitor patch 20 reads the data samples 511 and processes them and generates data samples (e.g., vital-sign values) of the sensor signal based on, e.g., an equation or a lookup table stored in a memory (e.g., 219, 210 of FIG. 2C).

[0072] In certain embodiments, the monitor patch 20 captures multiple readings of the sensor signal during the normal measurement interval (e.g., 8 minutes) before generating a data sample. For example, the monitor patch 20 can read the sensor signal enough times for a sample. For example, the monitor patch 20 can read the sensor signal enough times for a signal processing algorithm coded into the patch firmware to identify certain relevant features of the waveform and to calculate the values. In one exemplary embodiment, the features of the waveform and to calculate the values. In one exemplary embodiment, the sensor signal is read at 500 samples per second and the processor converts (e.g., calculates and averages) the multiple readings (e.g., 2000 readings) into one data sample at the end of the normal measurement interval.

[0073] The process 600 proceeds to operation 620 in which each of the data samples 511-514 is transmitted to a surveillance server 60 either directly or via a bridge. The monitor patch transmits the data samples to either a bridge (e.g., 401 of FIG. 11), which then transmits the data sample to the server 60 either directly or via a bridge 401. Such data samples representing vital-sign values of a patient may be displayed on a display terminal or logged in a database. The monitor patch is still considered to be in the normal mode of operation 606 at this stage.

[0074] The process 600 proceeds to decision state 630 in which it is determined whether the generated data sample satisfies an alert condition. In the illustrated example of FIG. 5, the generated data sample satisfies an alert condition. In the illustrated example of FIG. 5, the alert condition includes one data sample being at or above the first threshold limit 501. In the alert condition includes one data sample being at or above the first threshold limit 501. In the illustrated example of FIG. 5, data sample 514 has reached the first threshold limit 514, thereby satisfying the alert condition. In other embodiments, the alert condition may include one or more consecutive samples being at or above the first threshold limit 501.

[0075] If it is determined at the decision state 630 that the alert condition is not satisfied (No), the process 600 loops back to the operation 610 where another data sample is generated. On the other hand, if it is determined at the decision state 630 that an alert is generated. On the other hand, if it is determined at the decision state 630 that the alert
condition is satisfied (Yes) as a result of, for example, one or more data samples being at or above the first threshold limit, the process 600 proceeds to operation 640, in which a message indicative of the alert condition is transmitted to the surveillance server 60 either directly or via a bridge 40.

[0076] The process 600 then proceeds to operation 650 in which a data sample is generated at a second (alert) sampling rate (i.e., after a second measurement interval from a previous data sample), where the second sampling rate is higher than the first sampling rate. In the illustrated example of FIG. 5, the second sampling rate is one data sample per every 2 minutes corresponding to a second (alert) measurement interval of 2 minutes. At this stage, the monitor patch 20 is considered to be in an alert mode of operation 608 corresponding to, e.g., the second region 506 of FIG. 5. In certain embodiments, the monitor patch 20 captures multiple readings of the sensor signal during the second measurement interval (e.g., 2 minutes) and converts the multiple readings into one data sample at the end of the measurement interval. The process 600 proceeds to operation 660 in which each of data samples 535-545, thus generated, is transmitted to the surveillance server 60 either directly or via a bridge 40 at the second sampling rate.

[0077] If the process 600 then proceeds to decision state 670 in which it is determined whether the alert condition that caused the monitor patch to enter the alert mode 608 has ceased to exist. In certain embodiments, the alert condition is considered to have ceased to exist when the most recent data samples of the vital sign are less than the first threshold limit 501 consecutively for a predetermined number (e.g., 1-20) of times. For example, in the illustrated example of FIG. 5, data samples 536-545 have fallen below the first threshold limit 501 and remain consecutively so for 10 times. Assuming that the predetermined number is 10, the monitor patch determines that the alert condition has ceased to exist. If it is determined at the decision state 670 that the alert condition still exists (No), the process 600 proceeds to the operation 650 where a next data sample is generated and then to the operation 660 where the next data sample is transmitted to the surveillance server. On the other hand, if it is determined at the decision state 670 that the alert condition has ceased to exist (Yes), the process 600 exits the alert mode 608 and proceeds to operation 680 in which the process 600 exits the alert mode 608 and proceeds to operation 680 in
which a message indicative of the cessation of the alert condition is transmitted to the surveillance server either directly or via a bridge and then back to the operation 610 in the normal mode 606 (corresponding to the third region 508 of FIG. 5). Where data samples are again generated at the first sampling rate starting with data sample 546.

[0079] FIG. 7 is a flowchart illustrating an exemplary vital-sign monitoring process 700 from the perspective of a surveillance server (e.g., 610 of FIG. 1) according to certain aspects of the present disclosure. For the purposes of illustration only, without any intent to limit the scope of the present disclosure in any way, the process 700 will be described with references to FIG. 6. The process 700 begins at start state 701 and proceeds to operation 710 in which a data sample from the vital-sign monitor patch 20 is received. Without loss of generality, it is assumed that the data sample was sent while the monitor patch 20 was in the normal mode 606. Therefore, at this point in the process, such data samples are received at the first (normal) sampling rate. The surveillance server 605, in some embodiments, sends the data samples to a hospital system (e.g., 100 of FIG. 1) to be displayed on a display terminal or logged in a database.

[0080] The process 700 proceeds to decision state 710 in which it is determined whether the surveillance server has received a message indicative of the alert condition sent from the monitor patch 20, as discussed above with respect to operation 640 of FIG. 6. If it is determined at the decision state 710 that such a message has not been received by the surveillance server (No), the process 700 loops back to the operation 710 where the surveillance server 605 waits for and receives another data sample from the monitor patch 20. If the surveillance server 605 waits for and receives another data sample from the monitor patch 20 in which the surveillance server 60 waits for and receives a next data sample from the monitor patch 20 at the second (alert) sampling rate and then to decision state 740 in which it is determined whether one or more recently received data samples including the present data sample satisfy an alarm condition. In certain embodiments, the alarm condition includes the set of data samples having been at or above a threshold limit (e.g., the first threshold limit).
501 of FIG. 5) and rising consecutively for a predetermined number of times. For example, a processor in the surveillance server 60 can perform such an alarm condition determination.

[0082] If it is determined at the decision state 740 that the alarm condition has been satisfied, the process 700 proceeds to operation 745 in which an alarm notification is sent to the hospital system (e.g., 100 of FIG. 1). In the illustrated example of FIG. 5, a first alarm notification is sent to the hospital system upon receiving data sample 517, which corresponds to a first consecutively rising data sample after the vital sign has reached the first threshold limit 501. Subsequently, a second alarm notification is sent to the hospital system upon receiving data sample 523, which corresponds to a second consecutively rising data sample after the second (e.g., danger) threshold limit 502 has been reached. In certain embodiments, an alarm notification is sent after reaching a threshold limit. A current data sample is compared to a previously notified data sample (data sample at which an alarm notification was sent), and a new alarm notification is sent if a difference between the current data sample and the previously notified data sample exceeds a certain threshold difference value. In the illustrated example, based on this algorithm, a third alarm notification is sent to the hospital system upon receiving data sample 526 because the difference between the data sample 526 and the previously notified data sample 523 has exceeded a threshold difference value (e.g., 12).

[0083] Therefore, in certain algorithmic embodiments of the present disclosure, an alarm notification is sent when a current data sample exceeds a threshold limit (e.g., 501, 502) and satisfies one of the following "high" conditions:

1) The current data sample corresponds to X\textsuperscript{th} consecutively rising data sample after initially crossing the threshold high limit.

2) A difference between the current data sample and a previously notified data sample has exceeded a certain threshold difference value.

The first high condition can serve the purpose of providing an alarm notification indicative of a crossing of the threshold high limit (warning or danger). The second high condition can serve the purpose of providing an alarm notification indicative of a crossing of the threshold high limit (warning or danger).
serve the purpose of providing an alarm notification indicative of a progressively worsening
condition (e.g., rapidly rising temperature).

[0084] The hospital system may use the received notification, for example, to cause a nurse
[0084] The hospital system may use the received notification, for example, to cause a nurse
or doctor responsible for the patient to either monitor the progress of the condition in case of
or doctor responsible for the patient to either monitor the progress of the condition in case of
the first and second alarm notifications or to take certain actions (e.g., administering of a
the first and second alarm notifications or to take certain actions (e.g., administering of a
fever-reducing medication) in case of the third alarm notification.

[0085] The above-described algorithmic embodiments related to sending of an alarm
[0085] The above-described algorithmic embodiments related to sending of an alarm
notification when the current data sample exceeds a threshold high limit (e.g., 501, 502). In
notification when the current data sample exceeds a threshold high limit (e.g., 501, 502). In
other embodiments, an alarm notification can be sent when the current data sample is below a
other embodiments, an alarm notification can be sent when the current data sample is below a
low threshold limit (e.g., a first or second threshold limit) and satisfied one or more
low threshold limit (e.g., a first or second threshold limit) and satisfied one or more
additional low conditions, examples of which include:
additional low conditions, examples of which include:

1) The current data sample corresponds to \( Y \) consecutively falling data samples after
initially crossing the low threshold limit.
2) A difference between a previously notified data sample and the current data sample
has exceeded a certain threshold difference value.

Similar to the high conditions described above, the first low condition can serve the purpose
of providing an alarm notification indicative of crossing of a threshold limit (warning or
danger). The second low condition can serve the purpose of providing an alarm notification
indicative of a progressively worsening condition (e.g., rapidly falling heart rate).

[0086] While the alarm notification algorithms have been described with respect to a
[0086] While the alarm notification algorithms have been described with respect to a
particular vital-sign measuring, either an increase or decrease of any or all of multiple vital
particular vital-sign measuring, either an increase or decrease of any or all of multiple vital
-signs that the monitor patch 20 is configured to monitor can cause an alarm notification to be
-signs that the monitor patch 20 is configured to monitor can cause an alarm notification to be
sent. For example, assuming that a monitor patch 20 is configured to monitor three vital
sent. For example, assuming that a monitor patch 20 is configured to monitor three vital
-signs (e.g., heart rate, respiratory rate, and temperature), there can be 12 threshold limits (two
-signs (e.g., heart rate, respiratory rate, and temperature), there can be 12 threshold limits (two
high limits and two low limits for each of the three vital signs).
high limits and two low limits for each of the three vital signs).

[0087] On the other hand, if it is determined at the decision state 740 that the one or more
[0087] On the other hand, if it is determined at the decision state 740 that the one or more
recently received data samples do not satisfy an alarm condition (No), the process 700
recently received data samples do not satisfy an alarm condition (No), the process 700
proceeds to decision state 750 in which it is determined whether the monitor patch 20 is still in the alert mode. In certain embodiments, this determination involves checking to determine if a message indicative of the cessation of the alert condition as such as the one, discussed above with respect to the operation 680 of FIG. 6 was received from the monitor patch 20. In other embodiments, this determination involves the surveillance server 60 applying the same criteria as the monitor patch 20 to the received data samples to determine whether the alert condition has ceased to exist. For example, the surveillance server can independently determine that the received data samples 536-545 have been below the first threshold limit 501 for 10 consecutive times, or 501 for 0 consecutive times.

If it is determined at the decision state 750 that the monitor patch has ceased to be in the alert mode (No), the process 700 loops back to the operation 710, where the operation 710 determines whether the alert condition has been satisfied. The processes 600 and 700 described above are for illustration only, and a multitude of additions, deletions, and modifications thereto are possible without departing from the scope of the present disclosure. For example, in certain alternative embodiments, a separate message indicative of the alert condition is not sent to the surveillance server as in the operation 640 of FIG. 6. Instead, the alert condition is indicated in the transmission of the data sample in the alert mode 608 at the operation 660. Similarly, a separate message data sample in the alert mode 608 at the operation 660. Similarly, a separate message indicative of the cessation of the alert condition may not be sent to the surveillance server as in the operation 680 of FIG. 6. Instead, the surveillance server can determine the cessation of the alert condition from applying the criteria of the decision state 670 to the received data samples as discussed above or from an indication of the 'cessation' included in the transmission of the received data samples. In some alternative embodiments, the surveillance server does not require a series of received samples to be consecutively rising as well as at or above a
threshold limit for a predetermined number of times before sending the alert notification to the hospital system. As applied to the illustrated example of FIG. 15, in such an alternative embodiment, a fourth alarm notification would have been sent to the hospital system upon receiving data sample 530.

[0090] One skilled in the art would understand in view of the present disclosure that various systems and methods described above with respect to FIGS. 5-7 provide a number of important benefits to the vital-sign monitoring system of the present disclosure. For example, because a monitor patch worn by a patient performs the highly power-consumptive data acquisition and transmission operations relatively infrequently while vital-sign values of the patient are normal (e.g., in the normal mode 606) and frequently only when sign values of the patient are abnormal (e.g., in the abnormal mode 608), the monitor patch 20 can exhibit significantly higher life expectancy. Furthermore, from the perspective of a surveillance server 60, because data samples are received from the monitor patch 20 relatively infrequently while the patient’s vital signs are normal, a significant reduction in the server’s resources (e.g., processing time and memory space) can accrue.

[0091] In addition, the systems and methods provide for filtering out medically insignificant events that can otherwise trigger alarm notifications to the healthcare provider, thereby causing the provider to make unnecessary trips to the patient, for example. For instance, an electrical noise or movement of the patient may produce a short-lived spike (1 or 2 samples) in the measurements, but an influence of such a spike would be ignored by the alarm condition determination algorithm in the surveillance server 60. The aforementioned benefits of this condition determination algorithm in the surveillance server 60 can be obtained without sacrificing the response time of the monitoring system. For instance, in the illustrated example of FIG. 5, whenever vital sign values exceed a threshold limit (e.g., 501, 502), the vital-sign values can be displayed on a display terminal of the hospital system 501, 502, the vital-sign values can be displayed on a display terminal of the hospital system 501, 502, the vital-sign values can be displayed on a display terminal of the hospital system 501, 502, the vital-sign values can be displayed on a display terminal of the hospital system 501, 502, the vital-sign values can be displayed on a display terminal of the hospital system (e.g., 100 of FIG. 1) with a delay no longer than 2 minutes. Further, the time necessary for the values to be displayed on the display terminal is independent of the time necessary to transmit alarm notifications to the hospital system.

[0092] Certain aspects of the alarm condition determination and the dynamic measurement interval adjustment described herein can be performed by the processor 202 (FIG. 2C)
executing one or more sequences of one or more instructions using threshold limits and/or
alert conditions contained in an internal machine-readable medium, such as, the internal
memory 219 or the memory 210. For example, the processor 202 can determine that one or
more vital-sign readings exceed the first threshold limit 501 and switch from a normal mode
to an alert mode and change the sampling rate from a first rate to a second rate higher than
the first rate. The processor 202 can also revert to the normal mode operation of the vital-
sign readings return below the first threshold limit 501 and change the sampling rate from the
second rate to the first rate.

The processor 202 may be a microprocessor, a microcontroller, a digital signal
processor (DSP), an application specific integrated circuit (ASIC), or an
microcontroller, a digital signal processor (DSP), or an application specific integrated circuit (ASIC) capable of executing
processor (DSP), or an application specific integrated circuit (ASIC) capable of executing
instructions, computer instructions, such instructions, threshold limits and alert conditions may be read
from an internal memory and/or transmitted. Execution of the sequences of instructions
contained in the memory 210 causes the processor 202 to perform the process steps (e.g., of FIG. 6)
described herein. One or more processors in a multi-processing arrangement may also be employed to execute the sequences of instructions contained in the memory 219. In [alternative embodiments,] hard-wired circuitry may be used in place of or in combination with, software
instructions to implement various embodiments. Thus, embodiments are not limited to any
specific combination of hardware, circuitry and software.

Certain aspects of the alarm condition determination described herein can be performed by the processor 310 (FIG. 3) executing one or more sequences of one or more
instructions using threshold limits and alert conditions contained in an internal machine-readable
medium such as the internal memory 312 or the memory 340. For example, the processor 310 can determine that one or more vital-sign readings exceed the first threshold
limit 501 consecutively for a preset number of times and send an alarm notification. The
processor 310 can also send the alarm notification more frequently if one or more vital-
sign readings exceed the second threshold limit 502.

The processor 310 may be a general-purpose microprocessor, a microcontroller, a
Digital Signal Processor (DSP), or an Application Specific Integrated Circuit (ASIC).
Programmable Gate Array (FPGA), a Programmable Logic Device (PLD), a controller, a Programmable Gate Array (FPGA), a Programmable Logic Device (PLD), a controller, a state machine, gated logic, discrete hardware components, or any other suitable device, or a combination of devices that can perform calculations or other manipulations of information.

[0096] Instructions, threshold limits, and alarm conditions may be read into the memory 312.

[0096] Instructions, threshold limits, and alarm conditions may be read into the memory 312, and/or from another machine-readable medium, such as a CD, flash memory, or a wireless transmission. Execution of the sequence of instructions contained in the memory 312, or the memory 313, causes the processor 310 to perform the process steps (e.g., FIG. 7) described herein. One or more processors in a multi-processing arrangement may also be employed to execute the or more processors in a multi-processing arrangement may also be employed to execute the sequences of instructions contained in memory 312. In alternative embodiments, hard-wired sequences of instructions contained in memory 312. In alternative embodiments, hard-wired circuitry may be used in place of or in combination with software instructions to implement circuitry may be used in place of or in combination with software instructions to implement various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments. Thus, embodiments are not limited to any specific combination of various embodiments.

[0097] The term "machine-readable medium" as used herein refers to any medium that participates in providing instructions to processor 202, 310, for execution or storing results of or parameters (e.g., variables or constants) for computations such as for the alert condition determination and the dynamic measurement interval adjustment by the processor 202 and the alarm condition determination, by the processor 310. Such a medium may take many forms, including, but not limited to, non-volatile media, volatile media, and transmission media. Non-volatile media include, for example, optical or magnetic disks, such as data storage device. Volatile media include dynamic memory, such as the memory 210. Transmission media include coaxial cables, copper wire, and fiber optics, including the wires that comprise bus 204. Common forms of machine-readable media include, for example, floppy disk, a flexible disk, hard disk, magnetic tape, any other magnetic medium, a CD-ROM, DVD, any other optical medium, punch cards, paper tape, any other physical medium with patterns of holes, a RAM, a PROM, an EPROM, a FLASH EPROM, any other medium with patterns of holes, a RAM, a PROM, an EPROM, a FLASH EPROM, any other memory chip or cartridge, a carrier wave, or any other medium from which a computer can read.

[0098] The foregoing description is provided to enable any person skilled in the art to practice the various embodiments described herein. While the foregoing embodiments have been particularly described with reference to the various figures and embodiments, it should be understood that the various figures and embodiments, it should
be understood that these are for illustration purposes only and should not be taken as limiting the scope of the claims.

[0099] The word "exemplary" is used herein to mean "serving as an example or illustration."

[0099] The word "exemplary" is used herein to mean "serving as an example or illustration."

Any aspect or design described herein as "exemplary" is not necessarily to be construed as preferred or advantageous over other aspects or designs.

A reference to an element in the singular is not intended to mean "one and only one" unless specifically stated, but rather "one or more." The term "some," refers to one only one" unless specifically stated, but rather "one or more." The term "some," refers to one or more. Underlined and/or italicized headings and subheadings are used for convenience or more. Underlined and/or italicized headings and subheadings are used for convenience only, do not limit the invention, and are not referred to in connection with the interpretation only, do not limit the invention, and are not referred to in connection with the interpretation of the description of the invention. All structural and functional equivalents to the elements of the various embodiments of the invention described throughout this disclosure that are known to or later come to be known to those of ordinary skill in the art are expressly incorporated herein by reference and intended to be encompassed by the invention.

Moreover, nothing disclosed herein is intended to be dedicated to the public regardless of whether such disclosure is explicitly recited in the above description.

All elements, parts, and steps described herein are preferably included. It is to be understood that any of these elements, parts and steps may be replaced by other elements, parts and steps or deleted altogether as will be obvious to those skilled in the art.

The person skilled in the art will understand that the method steps mentioned in this description may be carried out by hardware including but not limited to processors; input devices comprising at least keyboards, mouse, scanners, cameras; output devices comprising at least monitors, printers. The method steps are to be carried out with the appropriate devices when needed. For example, a decision step could be carried out by a decision-making unit in a processor by implementing a decision algorithm. The person skilled in the art will understand that this decision-making unit can exist physically or effectively, for example in a computer's processor when carrying out the aforesaid decision algorithm. The above analysis is to be applied to other steps described herein.
CONCEPTS

This writing has disclosed at least the following concepts.

Concept 1. A method of reducing false alarms associated with vital-signal monitoring, the method comprising:

- generating one or more data samples of a vital signal of a patient at a first sampling rate in a normal mode of operation;
- determining whether the one or more data samples satisfy an alert condition;
- entering an alert mode of operation if the alert condition is satisfied; and
- generating one or more additional data samples of the vital signal at a second sampling rate higher than the first sampling rate in the alert mode.

Concept 2. The method of Concept 1, wherein the alert condition comprises one or more data samples having a parameter value being equal to or greater than a first threshold limit of that parameter.

Concept 3. The method of Concept 2, wherein the entering comprises switching from the normal mode to the alert mode when the one or more data samples are equal to or greater than the first threshold limit for a predetermined number of times.

Concept 4. The method of Concept 2 further comprising switching from the alert mode to the normal mode when the one or more data samples are less than the first threshold limit for a predetermined number of times.

Concept 5. The method of Concept 1 further comprising sending the one or more data samples to a surveillance server at the first sampling rate in the normal mode.

Concept 6. The method of Concept 1 further comprising sending the one or more additional data samples to a surveillance server at the second sampling rate in the alert mode.

Concept 7. The method of Concept 1, wherein the steps of Concept 1 are performed by a processor in a vital-signal monitor patch.
Concept 8. A vital sign monitoring system, comprising:

- a vital-sign monitor configured to monitor one or more vital signs of a patient; and
- a surveillance server configured to gather data relating to the one or more vital signs of the patient from the vital-sign monitor.

Concept 8. The system of Concept 8, wherein the vital-sign monitor is further configured to:

- generate one or more data samples of a vital sign of a patient at a first sampling rate in a normal mode of operation;
- determine whether the one or more data samples satisfy an alert condition; and
- generate one or more additional data samples of the vital sign at a second sampling rate higher than the first sampling rate in the alert mode.

Concept 8. The system of Concept 8, wherein the vital-sign monitor is configured to monitor one or more vital signs of a patient; and

- a surveillance server configured to gather data relating to the one or more vital signs of the patient from the vital-sign monitor.

Concept 8. The system of Concept 8, wherein the vital-sign monitor is further configured to:

- generate one or more data samples of a vital sign of a patient at a first sampling rate in a normal mode of operation;
- determine whether the one or more data samples satisfy an alert condition; and
- generate one or more additional data samples of the vital sign at a second sampling rate higher than the first sampling rate in the alert mode.

Concept 9. The system of Concept 8, wherein the vital-sign monitor is a vital-sign monitor patch.

Concept 10. The system of Concept 8, wherein the one or more vital signs include at least one of body temperature, pulse rate, blood pressure, and respiratory rate.

Concept 11. The system of Concept 8, wherein the alert condition comprises the one or more data samples having a parameter value being equal to or greater than a first threshold limit of that parameter value.

Concept 12. The system of Concept 11, wherein the vital-sign monitor is configured to switch from the normal mode to the alert mode when the one or more data samples are equal to or greater than the first threshold limit consecutively for a first number of times.

Concept 13. The system of Concept 11, wherein the vital-sign monitor is configured to switch from the normal mode to the alert mode when the one or more data samples are less than the first threshold limit consecutively for a second number of times.

Concept 14. The system of Concept 8, wherein the vital-sign monitor is further configured to provide the one or more data samples to the surveillance server at the first sampling rate in the normal mode and at the second sampling rate in the alert mode.
Concept 15. The system of Concept 14 further comprising a bridge configured to provide a communication connection between the vital-sign monitor and the surveillance server.

Concept 16. The system of Concept 14, wherein the surveillance server is further configured:

- determine whether a series of data samples of the vital sign sent from the vital-sign monitor satisfy an alarm condition; and
- provide an alarm notification to a hospital system if the alarm condition is satisfied.

Concept 17. The system of Concept 16, wherein the alarm condition comprises the series of data samples sent from the vital-sign monitor being equal to or greater than a first threshold limit consecutively for a third number of times.

Concept 18. The system of Concept 16, wherein the alarm condition comprises:

- the series of data samples sent from the vital-sign monitor being equal to or greater than a second threshold limit higher than the first threshold limit consecutively for a fourth number of times;
- the series of data samples sent from the vital-sign monitor at the second sampling rate rising at a rate higher than a threshold rate.

Concept 19. The system of Concept 16, wherein the alarm condition comprises a difference between a current data sample and a data sample at which a previous alarm notification was sent exceeding a threshold difference value.

Concept 20. The system of Concept 16, wherein the surveillance server is further configured:

- to send at least some of the series of data samples to the hospital system to be displayed at a display terminal.

Concept 21. The system of Concept 16, wherein the alarm condition comprises a difference between a current data sample and a data sample at which a previous alarm notification was sent exceeding a threshold difference value.

Concept 22. The system of Concept 16, wherein the alarm condition comprises:

- the series of data samples sent from the vital-sign monitor being equal to or less than a third threshold limit consecutively for a fourth number of times; and
- the series of data samples being equal to or less than a third threshold limit consecutively for a fourth number of times.
WHAT IS CLAIMED IS:

1. A method of reducing false alarms associated with vital-sign monitoring, the method comprising:
   generating one or more data samples of a vital sign of a patient at a first sampling rate in a normal mode of operation;
   determining whether the one or more data samples satisfy an alert condition;
   entering an alert mode of operation if the alert condition is satisfied; and
   generating one or more additional data samples of the vital sign at a second sampling rate higher than the first sampling rate in the alert mode.

2. The method of claim 1, wherein the alert condition comprises the one or more data samples having a parameter value being equal to or greater than a first threshold limit of that parameter.

3. The method of claim 2, wherein the entering comprises switching from the normal mode to the alert mode when the one or more data samples are equal to or greater than the first threshold limit a predetermined number of times.

4. The method of claim 2, further comprising switching from the alert mode to the normal mode when the one or more data samples are less than the first threshold limit for a predetermined number of times.

5. The method of claim 1 further comprising sending the one or more data samples to a surveillance server at the first sampling rate in the normal mode.

6. The method of claim 1 further comprising sending the one or more additional data samples to a surveillance server at the second sampling rate in the alert mode.

7. The method of claim 1, wherein the steps of claim 1 are performed by a processor in a vital-sign monitor patch.

8. The method of claim 1, wherein the steps of claim 1 are performed by a processor in a vital-sign monitor patch.
8. A vital sign monitoring system, comprising:

a vital-sign monitor configured to monitor one or more vital signs of a patient; and

a surveillance server configured to gather data relating to the one or more vital signs of the patient from the vital-sign monitor, wherein the vital-sign monitor is further configured to:

- generate one or more data samples of a vital sign of a patient at a first sampling rate in a normal mode of operation;
- determine whether the one or more data samples satisfy an alert condition;
- enter an alert mode if the alert condition is satisfied; and
- generate one or more additional data samples of the vital sign at a second sampling rate higher than the first sampling rate in the alert mode.

9. The system of Claim 8, wherein the vital-sign monitor is a vital-sign monitor patch.

10. The system of Claim 8, wherein the one or more vital-signs include at least one of body temperature, pulse rate, blood pressure, and respiratory rate.

11. The system of Claim 8, wherein the alert condition comprises the one or more data samples having a parameter value being equal to or greater than a first threshold limit of that parameter value.

12. The system of Claim 11, wherein the vital-sign monitor is configured to switch from the normal mode to the alert mode when the one or more data samples are equal to or greater than the first threshold limit consecutively for a first number of times.

13. The system of Claim 11, wherein the vital-sign monitor is configured to switch from the alert mode to the normal mode when the one or more additional data samples are less than the first threshold limit consecutively for a second number of times.

14. The system of Claim 8, wherein the vital-sign monitor is further configured to provide the one or more data samples to the surveillance server at the first sampling rate in the normal mode and at the second sampling rate in the alert mode.
15. The system of Claim 14 further comprising a bridge configured to provide a
communication connection between the vital-sign monitor and the surveillance server.

16. The system of Claim 14, wherein the surveillance server is further configured to:
   determine whether a series of data samples of the vital sign sent from the vital-sign
   monitor at the second sampling rate satisfy an alarm condition; and
   provide an alarm notification to a hospital system if the alarm condition is satisfied.

17. The system of Claim 16, wherein the alarm condition comprises the series of data
   samples sent from the vital-sign monitor being equal to or greater than a first threshold limit
   consecutively for a third number of times.

18. The system of Claim 16, wherein the alarm condition comprises the series of data
   samples sent from the vital-sign monitor at the second sampling rate being equal to or greater
   than a second threshold limit higher than the first threshold limit consecutively for a fourth
   number of times.

19. The system of Claim 16, wherein the alarm condition comprises the series of data
   samples sent from the vital-sign monitor at the second sampling rate rising at a rate higher
   than a threshold rate.

20. The system of Claim 16, wherein the surveillance server is further configured to send
   at least some of the series of data samples to the hospital system to be displayed at a display
   terminal.

21. The system of Claim 16, wherein the alarm condition comprises a difference between
   a current data sample and a data sample at which a previous alarm notification was sent
   exceeding a threshold difference value.

22. The system of Claim 16, wherein the alarm condition comprises a difference between
   the series of data samples sent from the vital-sign monitor being equal to or less than a third threshold limit
   consecutively for a fourth number of times.
FIG. 2C
FIG. 3B
START

- GENERATE A DATA SAMPLE AT A FIRST SAMPLING RATE
  - TRANSMIT THE DATA SAMPLE AT THE FIRST SAMPLING RATE
  - ALERT CONDITION SATISFIED?
    - NO
    - YES
      - TRANSMIT A MESSAGE INDICATIVE OF THE ALERT CONDITION TO SERVER

- NORMAL MODE 606

- ALERT MODE 608
  - GENERATE A DATA SAMPLE AT A SECOND SAMPLING RATE
  - TRANSMIT THE DATA SAMPLE AT THE SECOND SAMPLING RATE
  - ALERT CONDITION CEASES TO EXIST?
    - NO
    - YES

TRANSMIT A MESSAGE INDICATIVE OF THE CESSATION OF THE ALERT CONDITION TO THE SERVER

FIG. 6
START

RECEIVE A DATA SAMPLE FROM PATCH

RECEIVED A MESSAGE INDICATIVE OF AN ALERT CONDITION FROM PATCH?

YES

RECEIVE ANOTHER DATA SAMPLE FROM PATCH

ALARM CONDITION?

YES

SEND HIGH ALARM NOTIFICATION TO HOSPITAL SYSTEM

NO

STILL IN THE ALERT MODE?

YES

NO

FIG. 7