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Dietiker

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(54) **SYSTEM AND METHOD FOR A
NON-INVASIVE MEDICAL SENSOR**

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

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Primary Examiner — Eric F Winakur

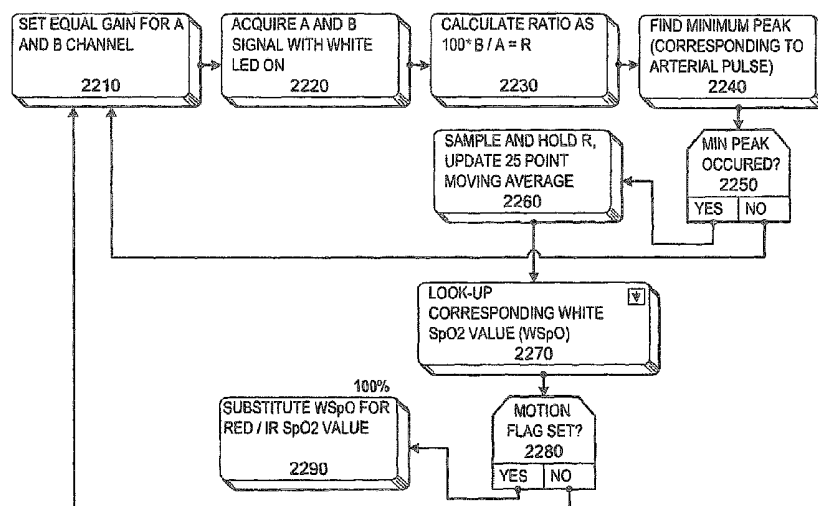
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(74) *Attorney, Agent, or Firm* — Adsero IP

(57) **ABSTRACT**

A method and system for measuring oxygen levels and various blood constituents utilizing a sensor having one or more light sources, and one or more light detectors is disclosed. The system is capable of using data collected by the one or more detectors from a non-monochromatic light source to provide accurate information during motion events occurring with an extremity the sensor. The system is also capable of detecting and providing an alert if the sensor is not properly placed on a patient or becomes disengaged therefrom.

6 Claims, 21 Drawing Sheets



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

CPC *A61B 5/6832* (2013.01); *A61B 5/6844*
 (2013.01); *A61B 5/7221* (2013.01); *A61B*
5/14552 (2013.01); *A61B 5/6838* (2013.01);
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 (2013.01)

(58) **Field of Classification Search**

CPC ... A61B 5/6838; A61B 5/6844; A61B 5/7207;
 A61B 5/7221; G01N 2201/121

See application file for complete search history.

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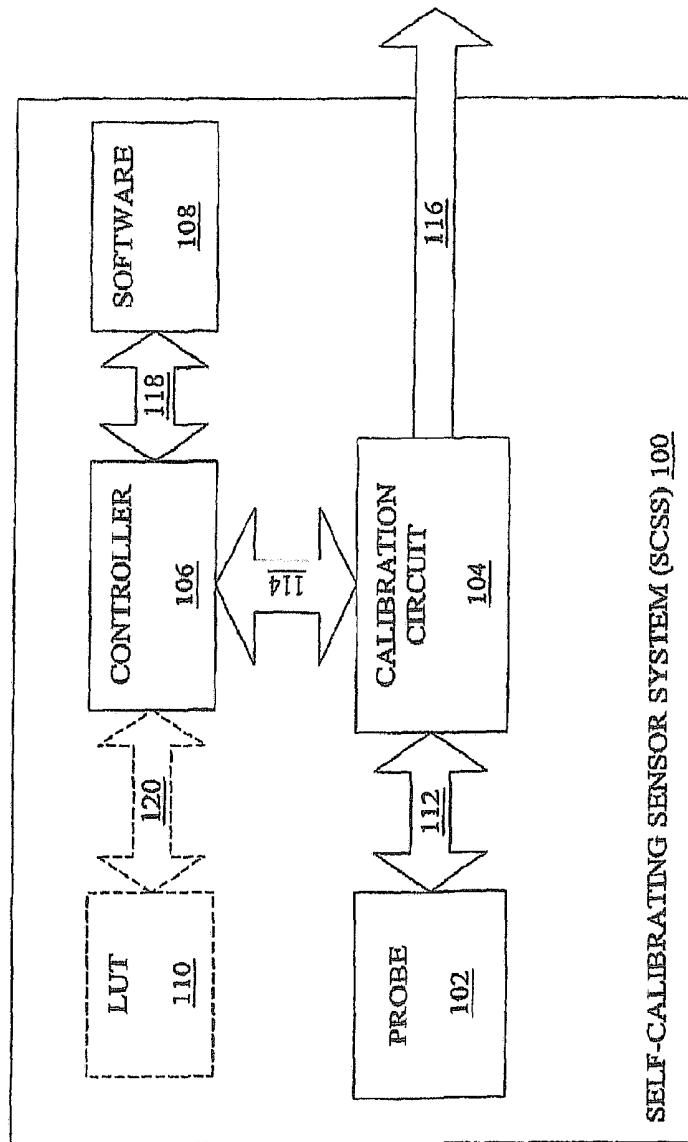


FIG. 1

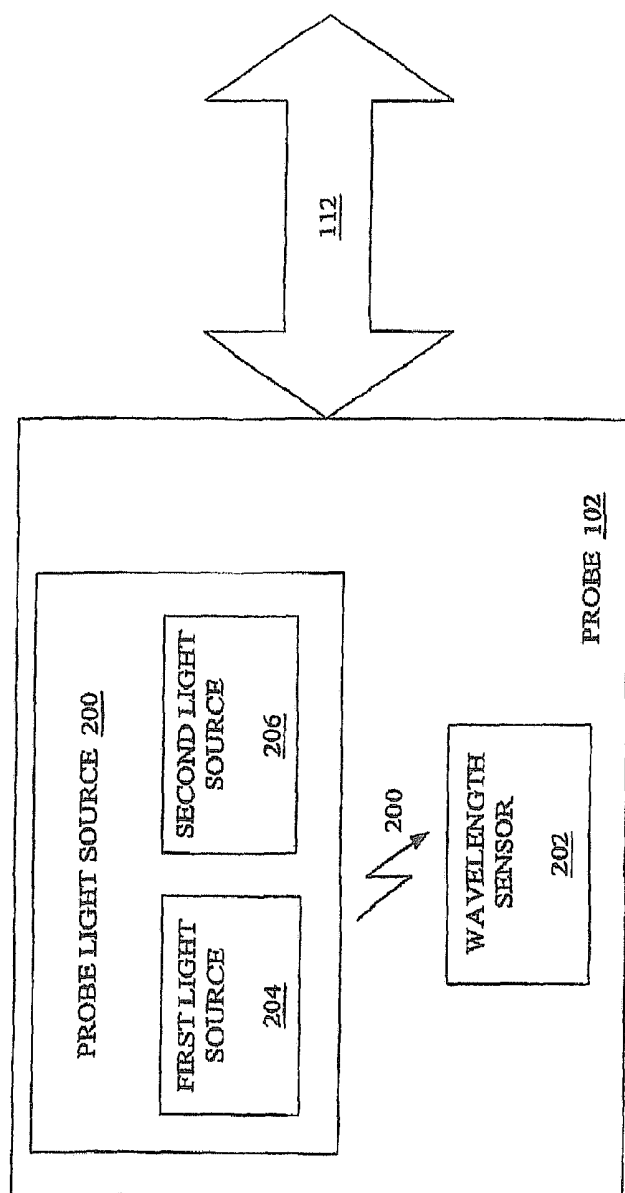


FIG. 2

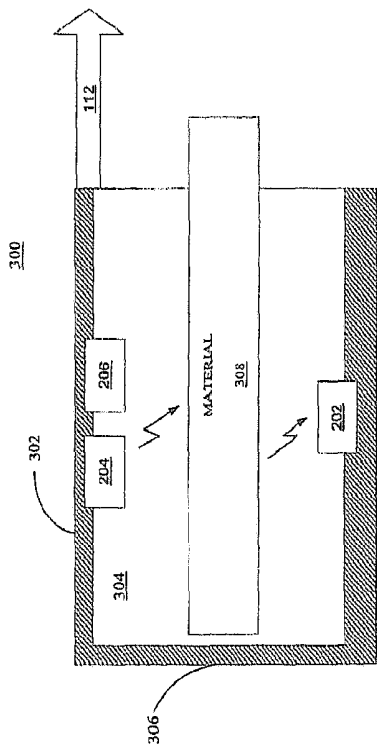


FIG. 3

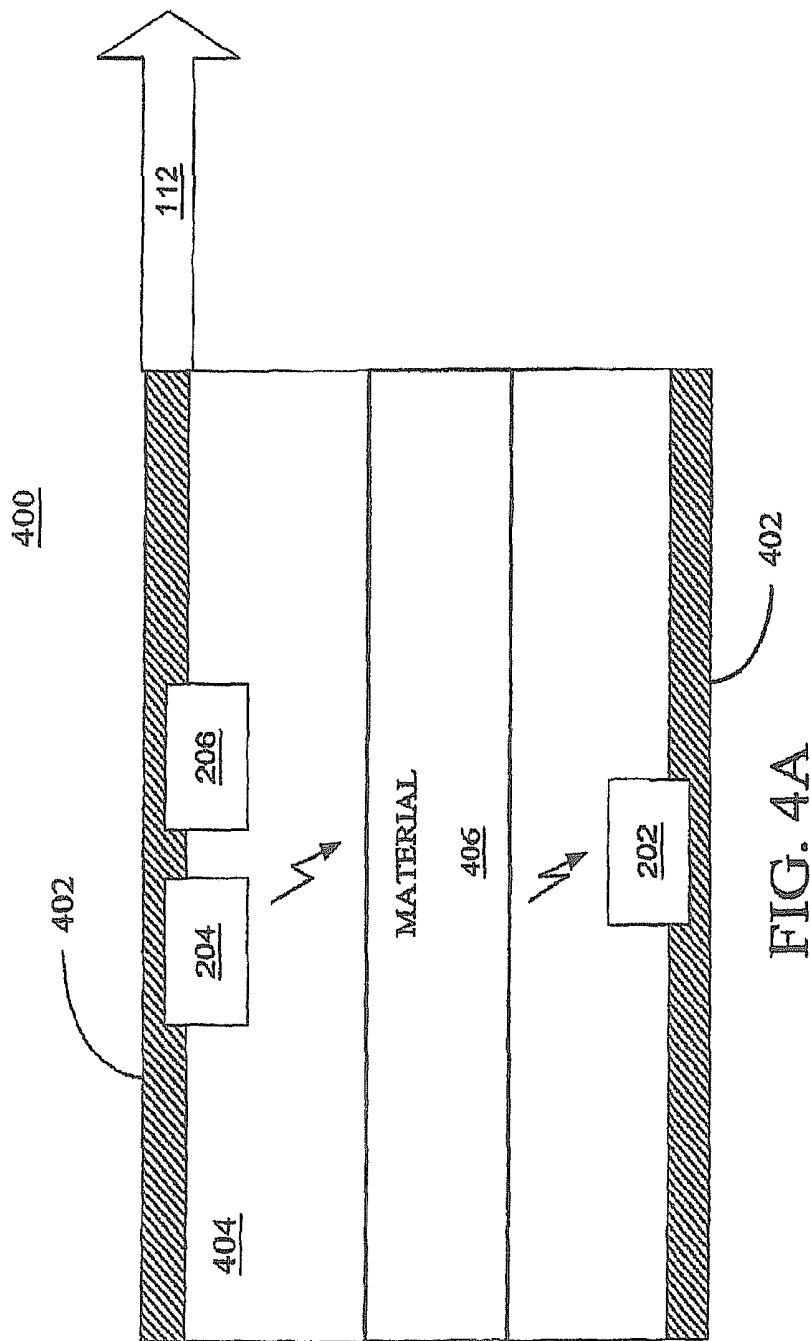
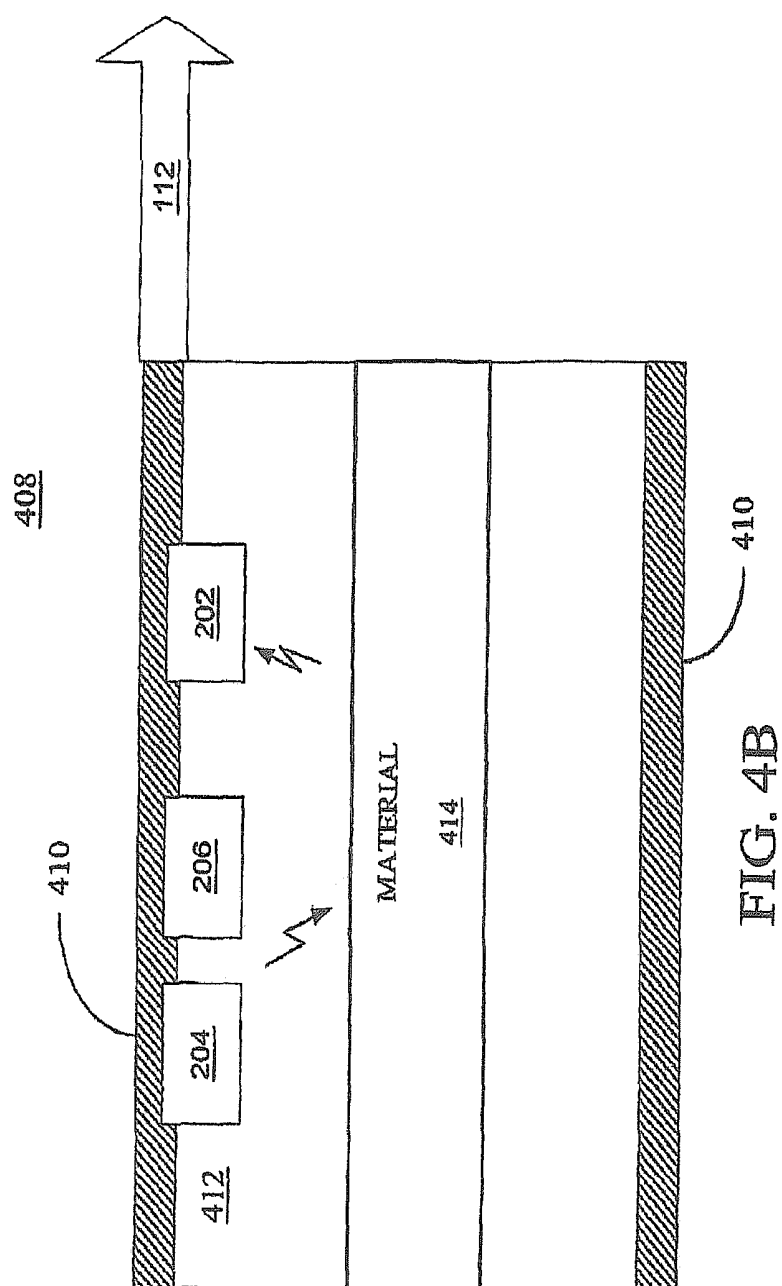


FIG. 4A



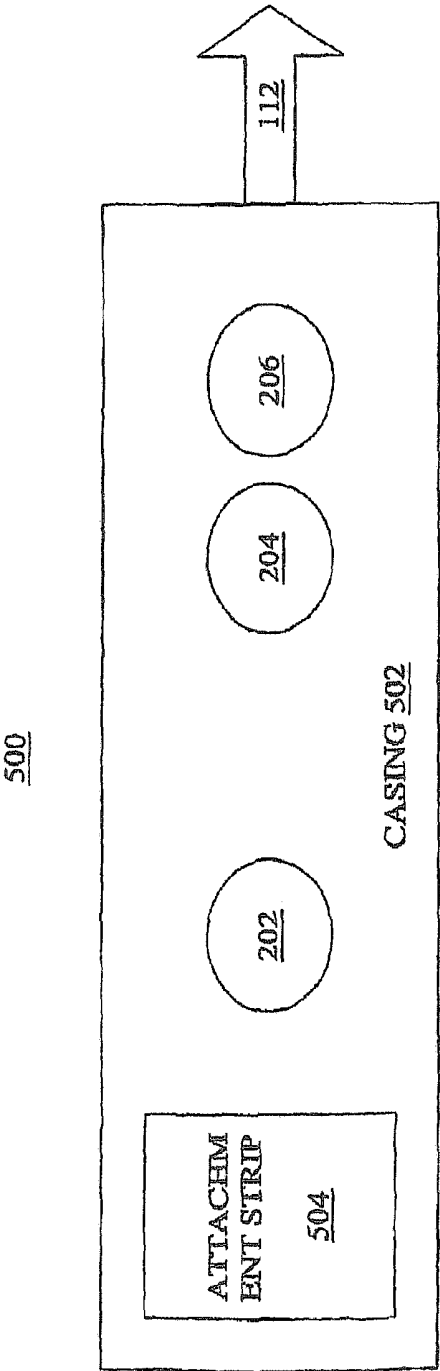


FIG. 5

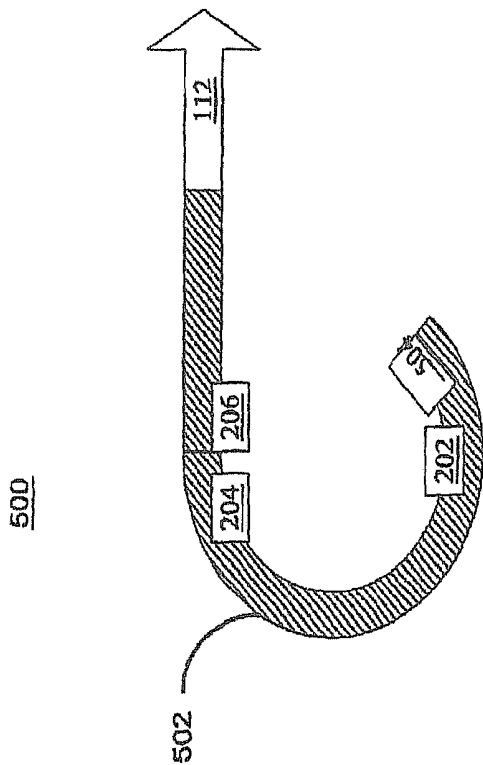


FIG. 6

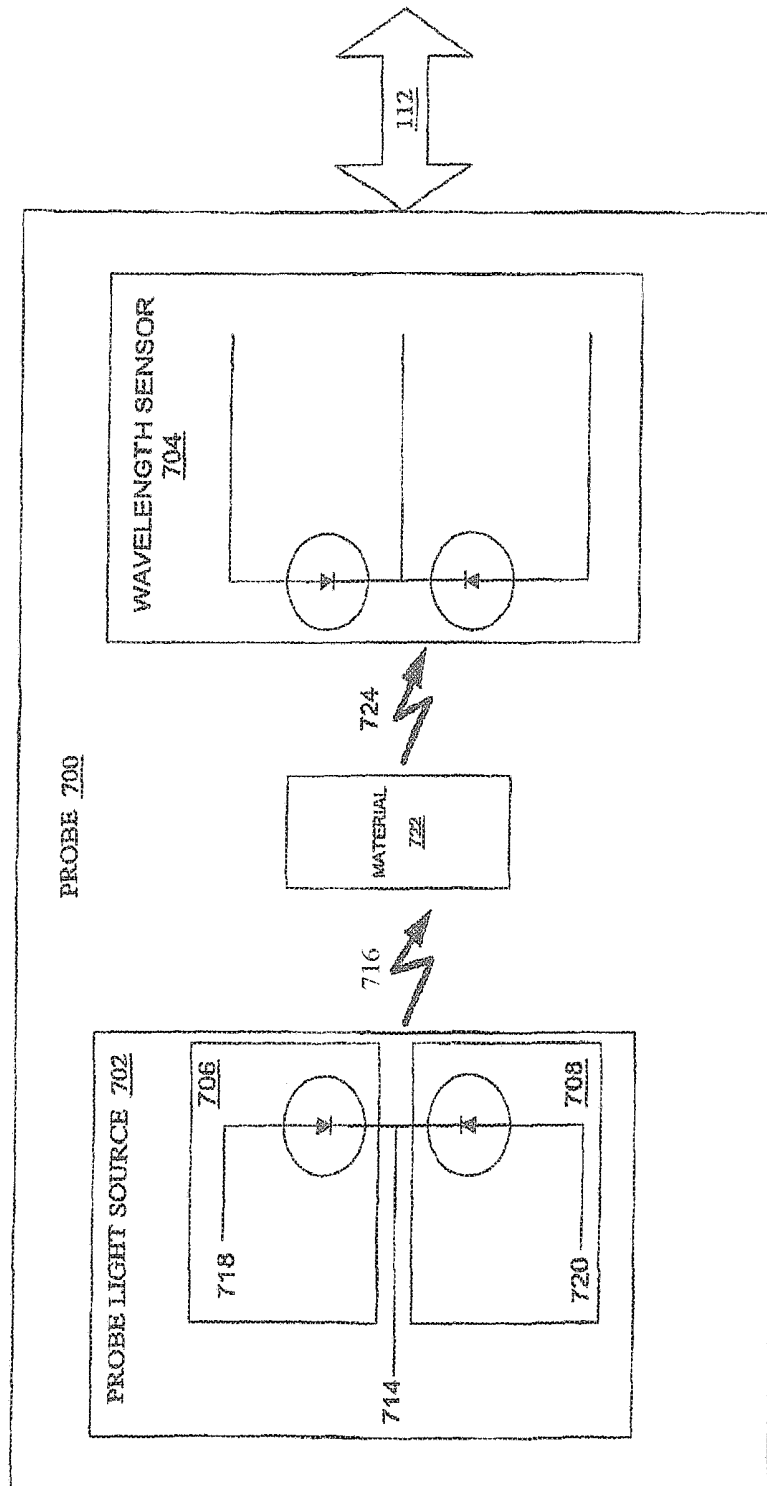


FIG. 7

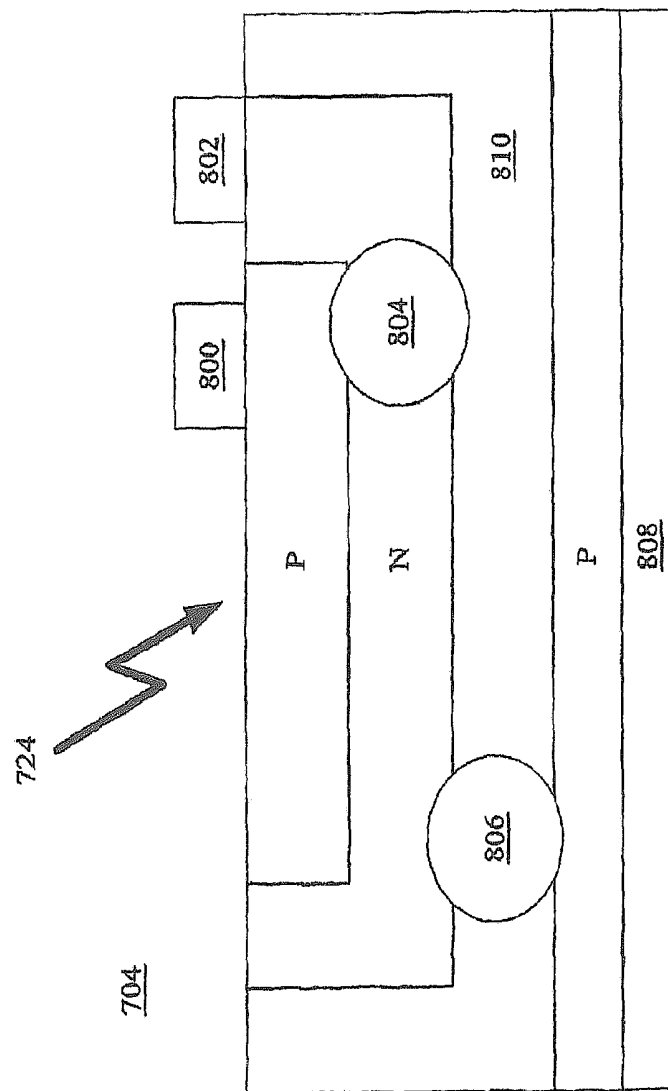


FIG. 8

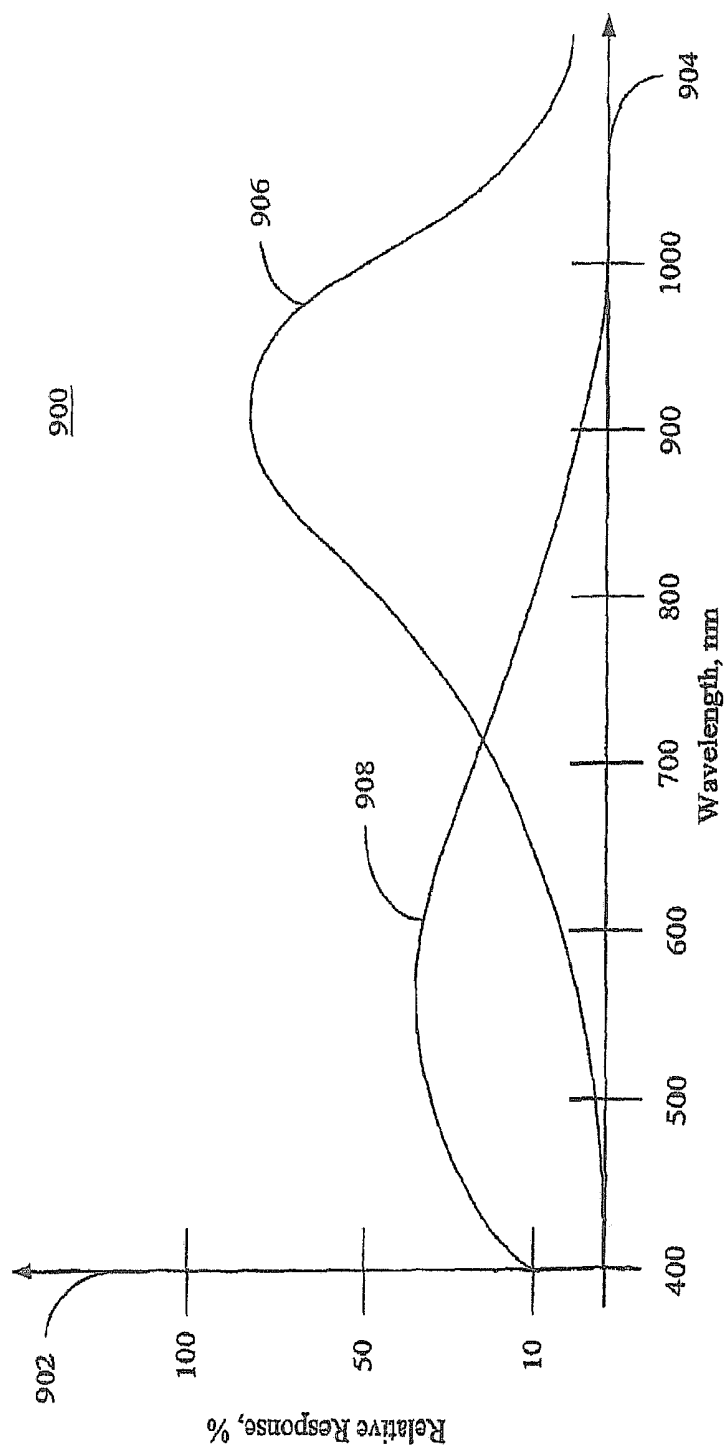


FIG. 9

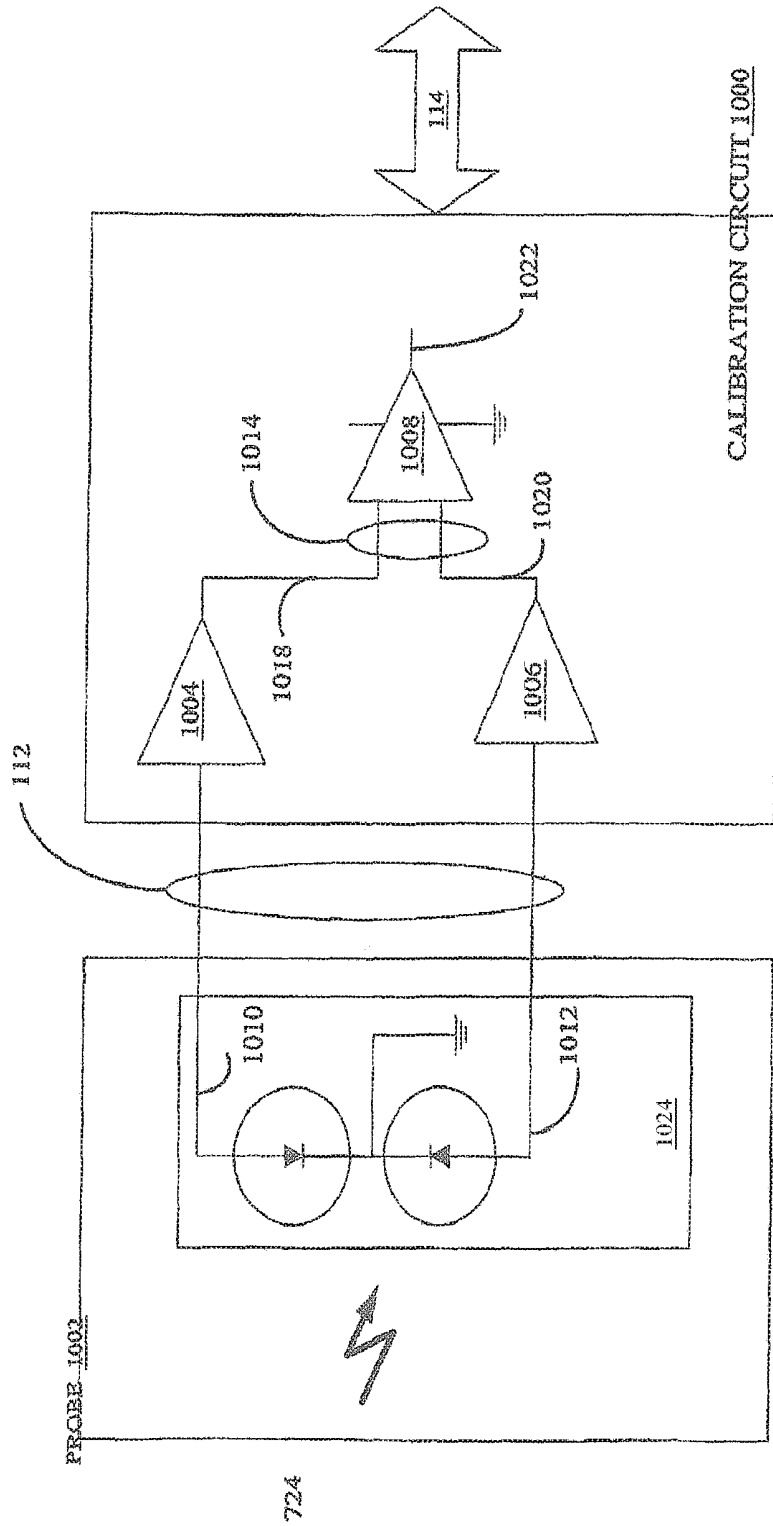


FIG. 10

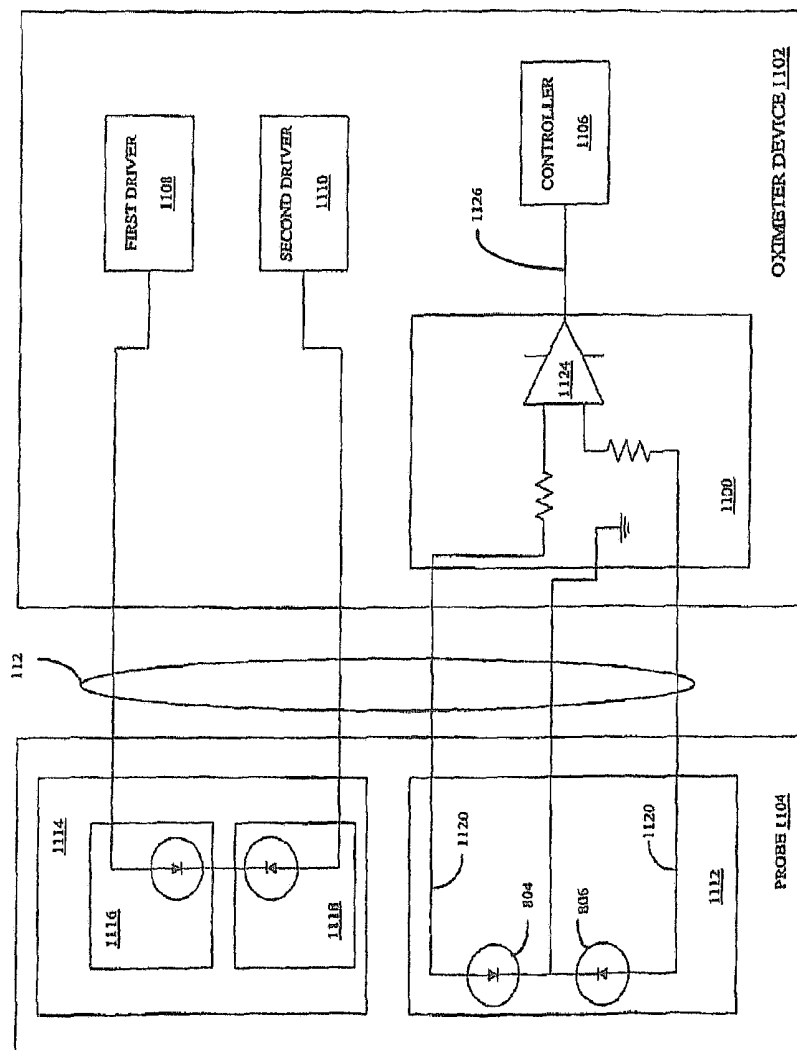


FIG. 11

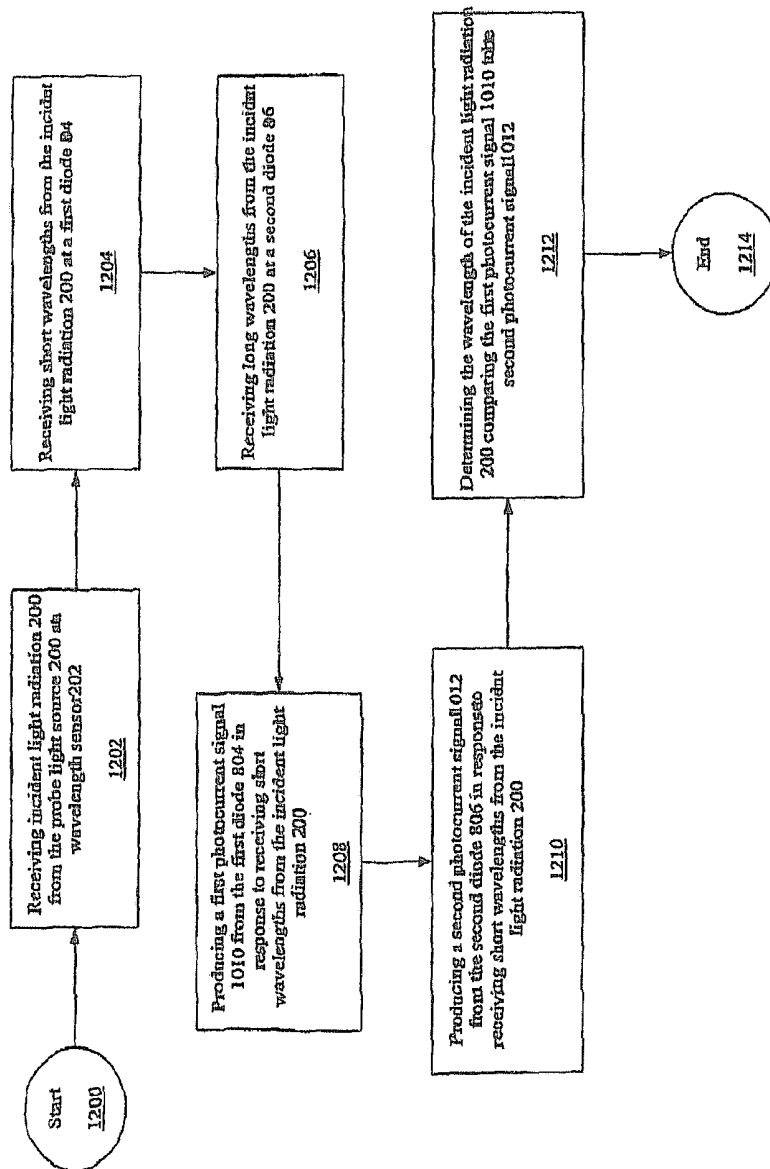


FIG. 12

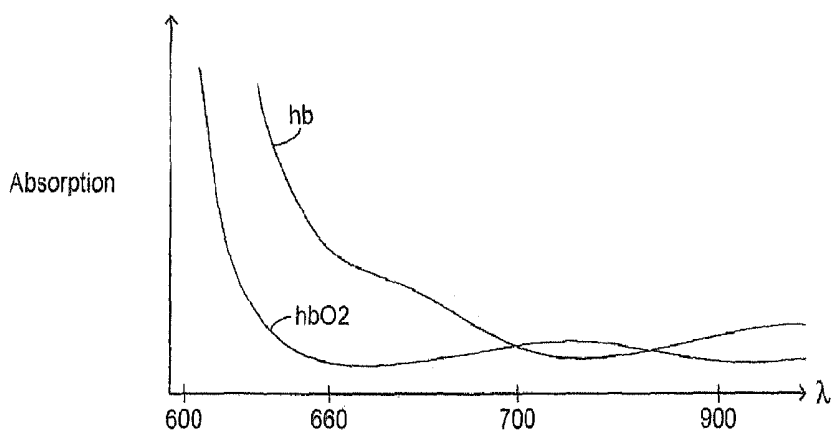
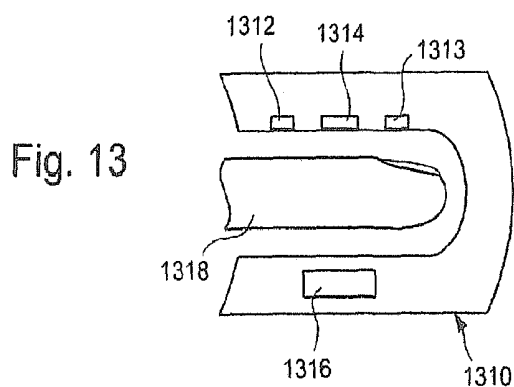


Fig. 14

Fig. 15

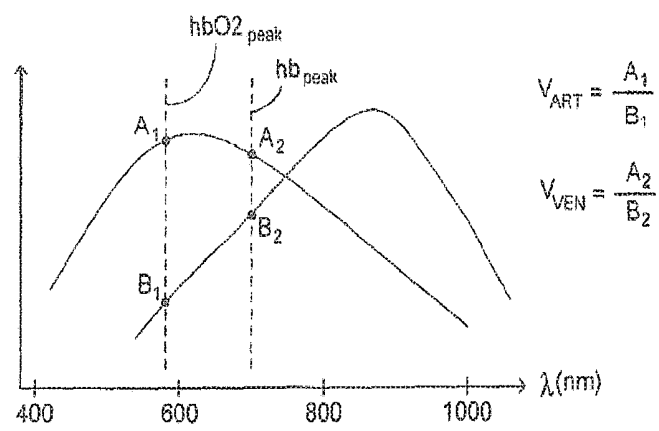
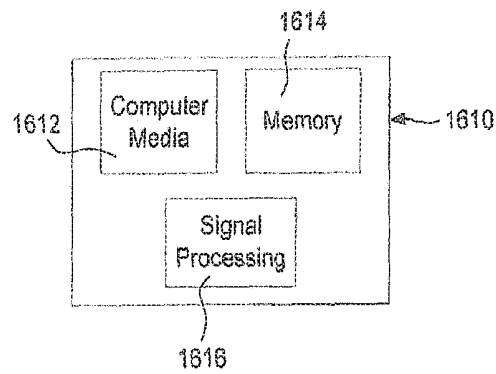


Fig. 16



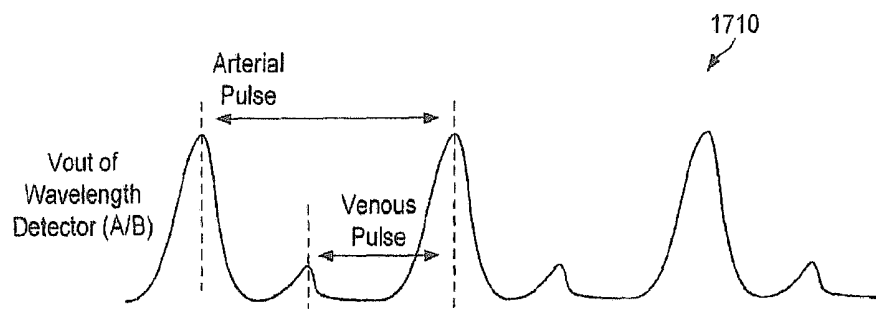


Fig. 17

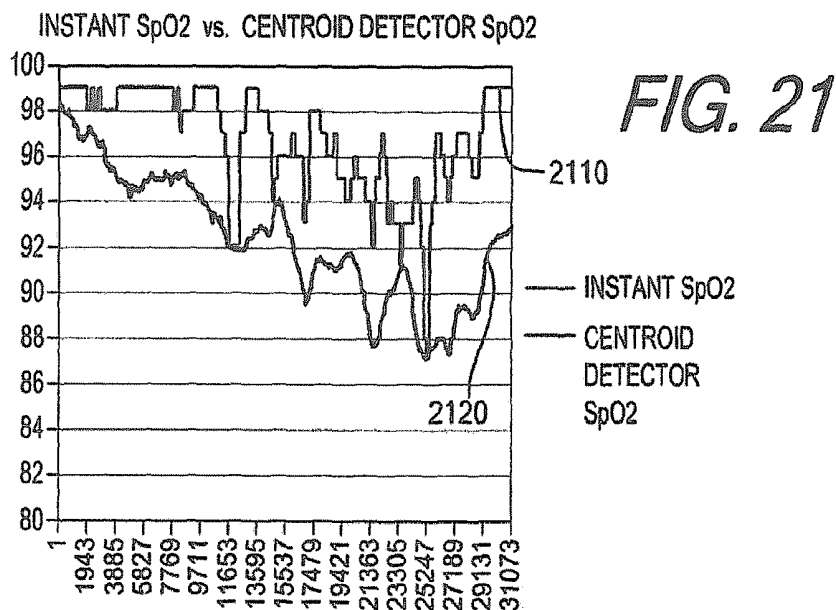
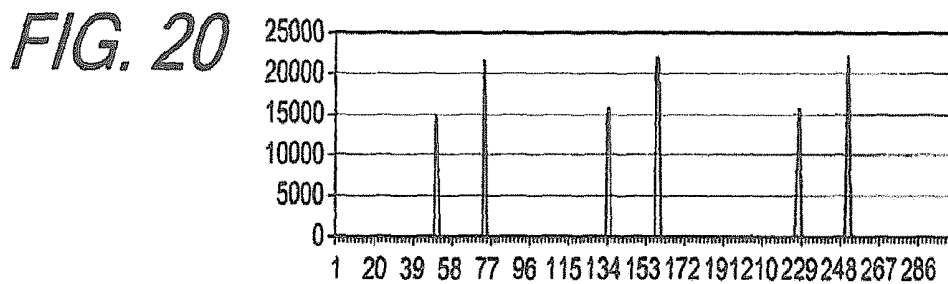
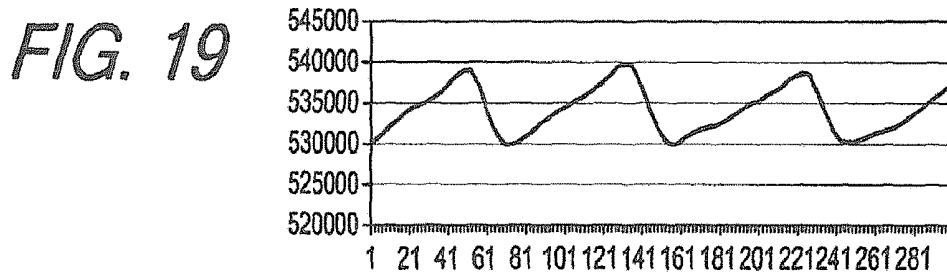
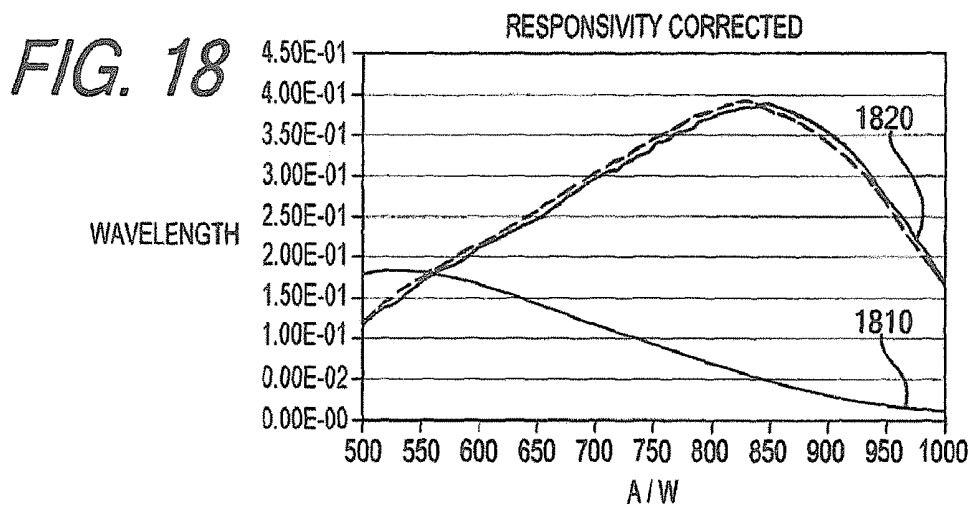


FIG. 22A

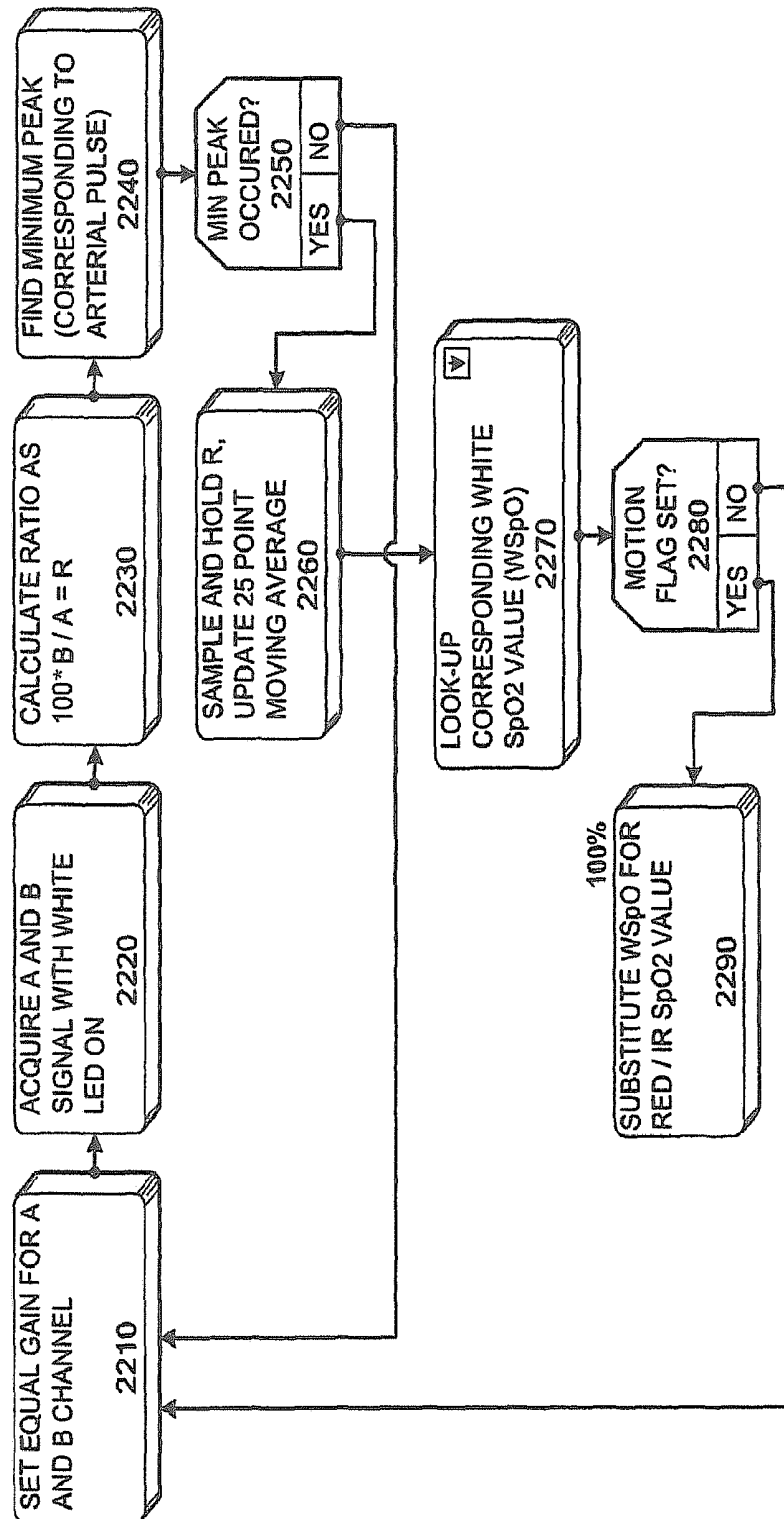


FIG. 22B

50%	
WHITE SPO2 LOOKUP TABLE	
87	
88	
89	
90	
91	
92	
93	
94	
95	
96	
97	
98	
99	

50%	
VAR SLOPE	
81.24750363	
81.56750363	
81.88750363	
82.20750363	
82.52750363	
82.84750363	
83.16750363	
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84.12750363	
84.44750363	
84.76750363	
85.08750363	

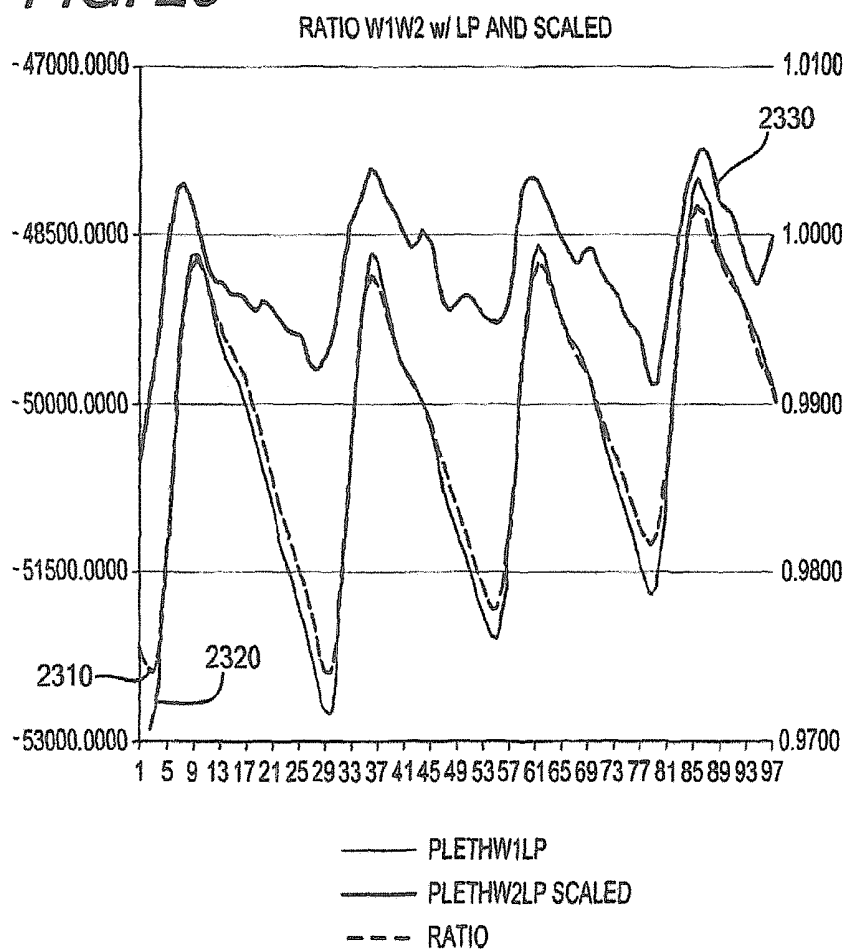
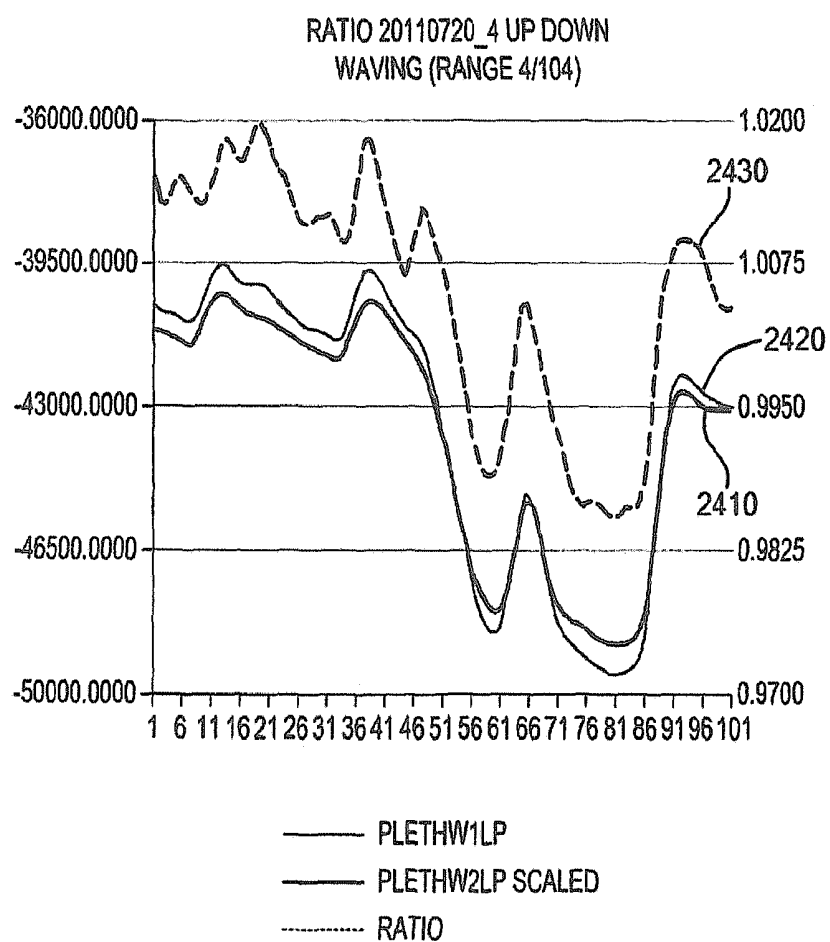
FIG. 23

FIG. 24

**Non-Perfused Object Signal from
Sprectral Detector with Broadband
Illumination**

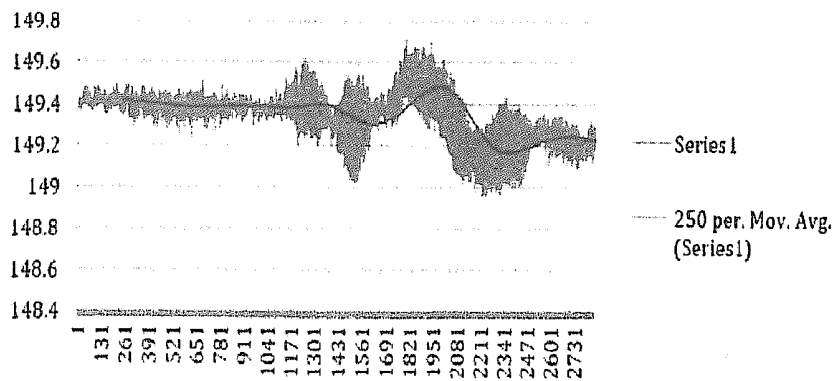


FIG. 25

**Perfused Subject Signal from Spectral
Detector with Broadband Illumination**

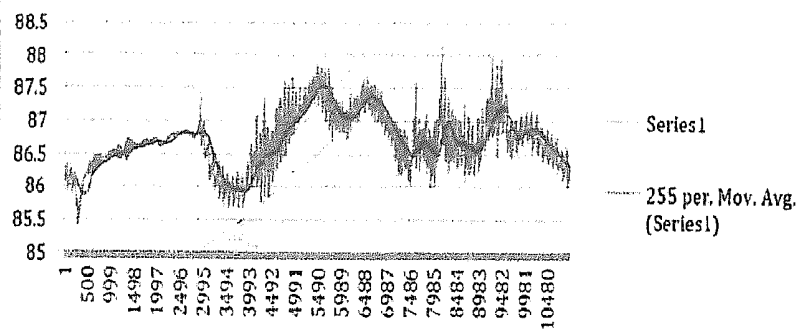


FIG. 26

SYSTEM AND METHOD FOR A NON-INVASIVE MEDICAL SENSOR

CROSS REFERENCE TO RELATED APPLICATIONS

This application is a national stage entry, under 35 USC § 371, of PCT PCT/US14/60004, filed Oct. 10, 2014 by Thomas Dietiker and titled, "System and Method for a Non-Invasive Medical Sensor", which is hereby incorporated by reference in its entirety.

RELATED APPLICATIONS

None.

FIELD OF THE INVENTION

The present invention relates generally to non-invasive medical sensing devices, and in particular to error correction for a medical sensor.

FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

None.

DESCRIPTION OF RELATED ART

Monochromatic light sources are utilized in a broad range of applications in many distinct fields of technology including the consumer, industrial, medical, defense and scientific fields. In the medical field an emitter-receiver pair of monochromatic light sources in form of light-emitting diodes (LEDs) or similar devices are often utilized in medical sensing devices to obtain accurate non-invasive measurements. An example application of such a medical sensing device may include a blood constituent monitoring system and/or a non-invasive oximeter that may be utilized to monitor arterial oxygen saturation.

In non-invasive oximetry, monochromatic light having a known specific wavelength is typically transmitted from an emitter LED through a target, such as biological tissue carrying blood, to a photodetector. The photodetector receives and measures a portion of transmitted coherent light that is neither absorbed nor reflected from the blood in the biological tissue in order to determine the oxygen saturation (SP02) within the blood. Similarly, an example of an industrial application may include a non-invasive sensor system having a monochromatic light of a known specific wavelength transmitted from a coherent light source (such as an LED emitter) through a target, such as a fluid or material, to a photodetector.

Unfortunately, these types of non-invasive sensor systems utilizing a monochromatic light source require accurate prior knowledge of the wavelength of the monochromatic light source in order to determine the amount of monochromatic light that is absorbed or reflected through the target. One way of having the prior knowledge of the wavelength is to select monochromatic light source emitters that have wavelengths within a certain range of tolerance. As such, attempts at determining the wavelength have included a binning process of selecting LEDs within the required nominal wavelength specifications and/or encoding data corresponding to the wavelength employed in a sensor as part of the sensor by means of a resistor, memory device or similar component.

However, it is appreciated by those skilled in the art and familiar with the production of emitter-photodiode sensing devices that there is a need to be able to select from a wider variation of emitter output wavelengths in reducing the production costs and defect rates of the sensing devices. As an example, typical production techniques require selection of an emitter within 2 nm of a target wavelength, which may lead to rejection of 40-60% of the component emitters. Moreover, an additional problem is that a selected emitter, which was within the target wavelength at time of production, will typically degrade over time, vary with temperature and drive current, and further the drive circuit may become unstable and cause a wavelength shift. Attempts to solve the wavelength shift problem have included systems that correlate the wavelength shift to a change in drive circuit current. The change in drive circuit current drives the LED to a specific wavelength. Typically, these systems include a scheme for determining the wavelength shift of the photodiodes via a series of filters, diffusers and a plurality of photodetectors. Unfortunately, this approach is too complex and expensive for practical manufacturing techniques and cannot be employed in each sensor being used with monitoring devices.

There is also a significant need to be able to reduce or eliminate variations resulting from patient movement, also commonly referred to as 'Motion Artefacts', differences in skin pigmentation and to detect fault situations, like for example, when a sensor becomes disengaged from a patient and is no longer able to deliver accurate information. Using current sensors and monitoring systems, such situations are generally difficult to detect and can lead to false readings and risk to the patient.

Therefore, there is a need for a non-invasive sensor system that is capable of measuring the wavelength of a light source without requiring prior knowledge of the wavelength of the light source and is not complex or expensive to manufacture. Moreover, there is a need for error reduction in such sensor systems.

There is also a need for a non-invasive sensor system which is capable of providing reliable data during motion events, or when the subject is engaged in movement or other physical activity or to provide a satisfactory notification or warning resulting from a patient disengaging the non-invasive sensor.

SUMMARY OF THE INVENTION

The present invention is directed to a non-invasive medical sensing device and system capable of providing correct data in response to a patient's movement and providing an alert should the device become disengage from the patient.

According to one aspect of the invention the sensor has one or more light sources and one or more light detectors. At least one of the detectors produces an output in response to the received light. Further, electrical circuitry responsive to the output generates a pulse waveform proportional to the arterial and venous pulse of a human body, wherein the pulse waveform is used to synchronize an arterial-pulse measurement of the absorption or reflectance of one or more monochromatic or non-monochromatic light sources.

According to one aspect of the invention, a method is provided.

The method includes providing a first detector and second detector in a non-invasive optical sensor and emitting light from a non-monochromatic light source in the non-invasive optical sensor periodically. A first signal from the first detector and a second signal from the second detector are

generated, resulting from light periodically emitted by the non-monochromatic light source sampled by the first detector and the second detector, both detectors having well defined and differing spectral responses. A third signal is generated using a ratio of the first signal and the second signal to determine an average spectral absorption and any changes in spectral absorption generated by the occurrence of an arterial pulse and base line spectral absorption of tissue located within the non-invasive optical sensor. According to another aspect of the invention, at least one light emitter emitting red light and at least one light emitter emitting infrared light is provided in the non-invasive optical sensor. A red light signal level from emitted red light received by the first detector and the second detector may be generated and infrared signal level received from emitted infrared light received by the first detector and the second detector may be generated. A ratio of the red light signal level to the infrared signal level may be correlated to a corresponding value sampled at an arterial peak while the non-monochromatic light source is emitting light in order to generate oxygen saturation data in the tissue.

Oxygen saturation data may be generated using the first signal and the second signal if a motion event occurs with the tissue located within the non-invasive optical sensor.

According to another aspect of the invention, a ratio of non-monochromatic light received by the first detector and the second detector may be measured. If the ratio of received non-monochromatic light is above or below a threshold, a notification may be generated.

According to another aspect of the invention, a centroid wavelength of emitted red light may be measured. If the measured centroid wavelength is determined to be outside an acceptable centroid range, and alert may be generated. According to another aspect of the invention, the first detector and second detector may have different spectral response curves, and. The different spectral response curves may be utilized to derive a fourth signal proportional to the centroid wavelength of emitted red light.

According to another aspect of the invention, a first detector and second detector are provided in a non-invasive optical sensor. Light may be emitted from a non-monochromatic light source in the non-invasive optical sensor periodically, and an amplitude of non-monochromatic light received by the first detector and the second detector may be measured. A notification may be generated if the amplitude of received non-monochromatic light is above or below a threshold.

Other embodiments, systems, methods, features, and advantages of the present invention will be, or will become, apparent to one having ordinary skill in the art upon examination of the following drawings and detailed description. It is intended that all such additional systems, methods, features, and advantages included within this description be within the scope of the present invention, and be protected by the accompanying claims.

BRIEF DESCRIPTION OF THE DRAWINGS

The invention may be better understood with reference to the following figures. The components in the figures are not necessarily to scale, emphasis instead being placed upon clearly illustrating the principles of the invention. Moreover, in the figures, like reference numerals designate corresponding parts throughout the several views:

FIG. 1 illustrates a block diagram of an example implementation of a self-calibrating sensor system (SCSS);

FIG. 2 illustrates a block diagram of an example implementation of the probe block of the SCSS shown FIG. 1;

FIG. 3 illustrates a cross-sectional view of an example implementation of the probe shown in FIG. 2;

FIG. 4A illustrates a cross-sectional view of another example implementation of the probe shown in FIG. 2;

FIG. 4B illustrates a cross-sectional view of example reflective implementation of the probe shown in FIG. 2;

FIG. 5 is a top view of an example implementation of the probe shown in FIG. 4;

FIG. 6 is a cross-sectional view of the probe implementation of FIG. 5;

FIG. 7 illustrates an example implementation of the probe block shown in FIG. 2 utilizing photodiodes;

FIG. 8 illustrates a cross-sectional view of an example implementation of the wavelength sensor block shown in FIG. 7 utilizing a double diffusion photodiode;

FIG. 9 is a graph of the response curve of the wavelength sensor shown in FIG. 8;

FIG. 10 is schematic diagram depicting an exemplary implementation of the calibration circuit block shown in FIG. 1;

FIG. 11 is schematic diagram depicting another exemplary implementation of the calibration circuit block shown in FIG. 1;

FIG. 12 is a flow chart illustrating the process performed by the SCSS shown in FIG. 1;

FIG. 13 is a simplified cross-sectional view of a sensor in accordance with the present invention;

FIG. 14 is a chart depicting the typical spectral absorption characteristics of an arterial and venous pulse;

FIG. 15 is a chart depicting response of the wavelength sensor to both arterial and venous pulses;

FIG. 16 is a simplified block diagram of a processing system for use with the sensor of FIG. 13;

FIG. 17 is a chart depicting a pulse waveform proportional to the arterial and venous pulse of the human body as produced after processing the signals from the wavelength sensor;

FIG. 18 is a chart depicting the response curves of the detectors of a dual detector module contemplated by the invention;

FIG. 19 is a chart depicting the change in spectral absorption generated by the arterial pulse blood flow with a broad emission illumination emitter as detected by a centroid wavelength detector as contemplated by the invention;

FIG. 20 is a chart depicting the peak and valley of the spectral absorption in the time domain;

FIG. 21 is a chart which depicts a signal proportional to the blood oxygen saturation level;

FIG. 22A is a flow chart showing the process through which pulse detection may be read and provided by the system through a motion event;

FIG. 22B shows an exemplary look up table utilized in the process shown in FIG. 22A;

FIG. 23 is a chart depicting saturation levels of a patient at rest;

FIG. 24 is a chart depicting saturation levels detected during a motion event;

FIG. 25 is a chart depicting raw data provided by a wavelength detector resulting from a non-perfused signal; and

FIG. 26 is a chart depicting raw data provided by a wavelength detector resulting from a perfused signal.

DESCRIPTION OF PREFERRED EMBODIMENTS

The following descriptions of detailed embodiments are for exemplifying the principles and advantages of the inven-

tions claimed herein. They are not to be taken in any way as limitations on the scope of the inventions.

FIG. 1 illustrates a block diagram of a self-calibrating sensor system (SCSS) 100. The SCSS 100 may include a probe 102, a calibration circuit 104, a controller 106, software 108 located in memory (not shown) and optional lookup table ("LUT") 110. The probe 102 is in signal communication, via the signal path 112, to the calibration circuit 104. The calibration circuit 104 may be a divider and/or comparator circuit.

The calibration circuit 104 is in signal communication to the controller 106 and an external output device (not shown) via signal paths 114 and 116, respectively. The controller 106 is in signal communication to software 108 and optional LUT 110 via signal paths 118 and 120, respectively.

The controller 106 may be any general-purpose processor such as an Intel XXX86, Motorola 68XXX or PowerPC, DEC Alpha or other equivalent processor. Alternatively, a specific circuit or oriented device may selectively be utilized as the controller 106. Additionally, the controller 106 may also be integrated into a signal semiconductor chip such as an Application Specific Integrated Chip (ASIC) or Reduced Instruction Set Computer (RISC), or may be implemented via a Digital Signal Processor (DSP) chip. Examples of a specific circuit or oriented device for the controller 106 may also be a mixed sionac ASIC.

The software 108 may be resident in memory (not shown) located either internally or externally to the controller 106. The software 108 includes both logic enabling the controller 106 to operate and also logic for self-calibrating the SCSS 100.

An example of the external output device may be an oximeter such as a NPB40 manufactured by Nellcor of Pleasanton, Calif., a 9840 Series pulse oximeter manufactured by Nonin Medical, Inc. of Plymouth, Minn., or an equivalent device.

FIG. 2 shows an example implementation of probe 102. Probe 102 may include a probe light source 200 and wavelength sensor 202. Probe light source 200 may include a first light source 204 and second light source 206. First light source 204 and second light source 206 may be implemented utilizing light-emitting diodes (LEDs). As an example implementation in oximeter application, first light source 204 may be an LED emitting light radiation at a wavelength of approximately 660 nm and second light source 206 may be an LED emitting light radiation at a wavelength of approximately 880 nm. Wavelength sensor 202 may be implemented utilizing a double diffusion photodiode. It is appreciated by those of skill in the art that probe light source 200 may also include multiple light sources in the order of three or more.

In FIG. 3, a cross-sectional view of an example implementation of the probe 300 is shown. In this example, probe 300 may be a medical device such as a transmissive blood oxygen saturation and pulse rate sensor. However, it would be appreciated by one skilled in the art that probe 300 may also be a reflective sensor. Additionally, probe 300 may also be utilized for measuring other blood constituents including, but not limited to, oxyhemoglobin, bilirubin, carboxy-hemoglobin, and glucose. Probe 300 may include a rigid casing 302 having a cavity 304 and casing butt 306, first light source 204, second light source 206 and wavelength sensor 202. Probe 300 is connected to calibration circuit 104, FIG. 1, via signal path 112. A material 308, FIG. 3, such as a finger may be inserted into the cavity 304.

As an example, first light source 204 and second light source 206 may be two LED emitters that produce light

radiation at a wavelength of approximately 660 nm and 880 nm, respectively. Wavelength sensor 202 is supported within the rigid casing 302 opposite first light source 204 and second light source 206. First light source 204 and second light source 206 and wavelength sensor 202 may be in signal communication with a control cable (not shown). The control cable is in signal communication with an oximeter (not shown) via signal path 112. The oximeter determines the oxygen saturation of the blood in the material 308 (in this example a finger) by measuring and processing the amount of incident light radiation reaching wavelength sensor 202 from a pulse of light radiation from first light source 204.

In operation, the SCSS 100, FIG. 1, performs a self-calibration procedure prior to and while measuring any of the properties of the material 308, FIG. 3. This self-calibration procedure includes emitting a pulse of light radiation from the first light source 204 that is received as incident light radiation by wavelength sensor 202 prior to inserting material 308 into the cavity 304. The oximeter utilizes the measured incident light radiation received by wavelength sensor 202 to determine the operating wavelength of the first light source 204. Once the operating wavelength of the first light source 203 is known, the SCSS 100, FIG. 1, is utilized in combination with the oximeter to accurately determine blood oxygen saturation of the material 308.

The continuous self-calibration procedure is beneficial because it is appreciated by those skilled in the art that light radiation output by first light source 204 of 660 nm in this example implementation is in the red spectral region. It is the absorption of this red light radiation that the oximeter utilizes to determine the oxygen saturation of the blood. As such, a relatively small variation in operating wavelength may results in inaccurate readings at the oximeter. As an example, without the self-calibration procedure, if the light radiation output by first light source 204 varied in excess of ± 2 nm from an operating wavelength required by the oximeter, the results would be inaccurate.

FIG. 4A illustrates a cross-sectional view of another example implementation of probe 400. In this example, probe 400 may include a rigid or flexible casing 402 having a cavity 404, first light source 204, second light source 206 and wavelength sensor 202. Similar to the previous example implementation, probe 400 is connected to calibration circuit 104, FIG. 1, via signal path 112, however, probe 400, FIG. 4A, does not have a cavity butt. A material 406 may be inserted into the cavity 404.

Similar to the previous example, first light source 204 and second light source 206 may be two LED emitters that produce light radiation at different wavelengths. Wavelength sensor 202 is supported within the rigid casing 402 opposite first light source 204 and second light source 206. First light source 204 and second light source 206 and wavelength sensor 202 may be in signal communication with a control cable (not shown). The control cable is in signal communication with a measuring device (not shown) via signal path 112. The measuring device determines the properties in the material 406 by measuring and processing the amount of incident light radiation reaching wavelength sensor 202 from a pulse of light radiation from first light source 204.

As an industrial example, the material 406 may be a fluid, liquid or solid material that exhibits optical transmissive characteristics that may be measured and utilized to determine the properties of the material. An example implementation would include measuring the properties of the material for process or quality control purposes.

Again in operation, the SCSS 100, FIG. 1, performs a self-calibration procedure prior to measuring any of the

properties of the material **406**, FIG. 4A. This self-calibration procedure includes emitting a pulse of light radiation from the first light source **204** that is received as incident light radiation by wavelength sensor **202** prior to inserting material **406** into the cavity **404**. The measuring device utilizes the measured incident light radiation received by wavelength sensor **202** to determine the operating wavelength of the first light source **204**. Once the operating wavelength of the first light source **204** is known, the SCSS **100**, FIG. 1, is utilized in combination with the measuring device to accurately determine the properties of the material **406**.

FIG. 4B illustrates a cross-sectional view of an example reflective implementation of probe **408**. In this example, probe **408** may include a rigid or flexible casing **410** having a cavity **412**, first light source **204**, second light source **206** and wavelength sensor **202**. Similar to the previous example implementation, probe **408** is connected to calibration circuit **104**, FIG. 1, via signal path **112**, however, probe **408**, FIG. 4B, does not have a cavity butt. A material **414** may be inserted into the cavity **412**.

Similar to the previous example, first light source **204** and second light source **206** may be two LED emitters that produce light radiation at different wavelengths. However, in this example, wavelength sensor **202** is supported within the rigid casing **410** adjacent to first light source **204** and second light source **206**. First light source **204** and second light source **206** and wavelength sensor **202** may be in signal communication with a control cable (not shown). The control cable is in signal communication with a measuring device (not shown) via signal path **112**. The measuring device determines the properties in the material **412** by measuring and processing the amount of incident light radiation reflected by material **412** and reaching wavelength sensor **202** from a pulse of light radiation from first light source **204**.

Again, as an industrial example, the material **412** may be a fluid, liquid or solid material that exhibits optical transmissive characteristics that may be measured and utilized to determine the properties of the material. An example implementation would include measuring the properties of the material for process or quality control purposes.

Again in operation, the SCSS **100**, FIG. 1, performs a self-calibration procedure prior to measuring any of the properties of the material **412**, FIG. 4B. This self-calibration procedure includes emitting a pulse of light radiation from the first light source **204** that is reflected by flexible casing **410** and later received as incident light radiation by wavelength sensor **202** prior to inserting material **412** into the cavity **410**. The measuring device utilizes the measured incident light radiation received by wavelength sensor **202** to determine the operating wavelength of the first light source **204**. Once the operating wavelength of the first light source **204** is known, the SCSS **100**, FIG. 1, is utilized in combination with the measuring device to accurately determine the properties of the material **412**.

It is appreciated by of skill in the art that it is possible to generate signals from the wavelength sensor **202**, FIG. 2 during operation of the light sources **204** and **206** through the medium (i.e., material **308**, FIG. 3, **406**, FIG. 4A, and/or **414**, FIG. 4B) being inspected. It is also possible to generate the same signals using light reflected off the medium. Therefore, it is not necessary to couple the light sources **204**, FIGS. 2 and **206** directly to the wavelength sensor **202** as long as the medium either transmits or reflects enough light to generate processable signals from the wavelength sensor **202**.

In FIG. 5, a top view of an example medical implementation of probe **500** having a flexible casing (i.e., flexible strip) **502** is shown. Probe **500** may include first light source **204**, second light source **206** and wavelength sensor **202**. In this example implementation, probe **500** is a blood oxygen saturation and pulse rate sensor that utilizes the flexible strip **502** to attach to a material, such as a body part (not shown). The probe **500** is connected to an oximeter (not shown) via signal path **112**. The flexible strip **502** may be wrapped around the body part and affixed to itself via an attachment strip (such as an adhesive strip) **504**. Example body parts would include a finger, toe, ear-lobe, arm, leg or other similar parts.

As an example, first light source **204** and second light source **206** may be two LED emitters that produce light radiation at a wavelength of approximately 660 nm and 880 nm, respectively. Wavelength sensor **202** is supported within the flexible strip **502** and placed opposite first light source **204** and second light source **206** when the flexible strip **502** is wrapped around a body part. First light source **204** and second light source **206** and wavelength sensor **202** may be in signal communication with a control cable (not shown). The control cable is in signal communication with an oximeter (not shown) via signal path **112**. The oximeter determines the oxygen saturation of the blood in the body part by measuring and processing the amount of incident light radiation reaching wavelength sensor **202** from a pulse of light radiation from first light source **204**.

As before, in operation, the SCSS **100**, FIG. 1, performs a self-calibration procedure prior to measuring any of the properties of the body part. This self-calibration procedure includes, prior to wrapping flexible strip **502** around the body part, bending the flexible strip **502** so that the first light source **204** and second light source **206** are opposite in special orientation to wavelength sensor **202** and then emitting a pulse of light radiation from the first light source **204** that is received as incident light radiation by wavelength sensor **202**. The oximeter utilizes the measured incident light radiation received by wavelength sensor **202** to determine the operating wavelength of the first light source **204**. Once the operating wavelength of the first light source **204** is known, placed around a body part and the wavelength sensor **202** measures the incident light radiation emitted by the first light source **204** and passing through the blood flowing within the body part. The SCSS **100**, FIG. 1, is then utilized in combination with the oximeter to accurately determine blood oxygen saturation of the body part. In FIG. 6, a cross-sectional view of the probe **500** is shown in a wrap type position.

In FIG. 7, an example implementation of the probe **700** is shown utilizing photodiodes. Similar to FIG. 2, Probe **700**, FIG. 7, includes probe light source **702** and wavelength sensor **704**. Probe light source **702** includes first light source **706** and second light source **708**. First light source **706** may include LED **710** and second light source may include LED **712**. Wavelength sensor **704** is a double diffusion photodiode.

As an example of operation, LED **710** and LED **712** may have their cathodes grounded in common at signal path **714** and may emit light radiation **716** at wavelengths 660 nm and 880 nm, respectively, when a voltage is applied at anodes **718** and **720**, respectively. The emitted light radiation **716** is incident on material **722**. A part of the emitted light radiation **716** is transmitted through material **722** and is received as incident light radiation **724** by wavelength sensor **704**. As before, in order to properly measure the properties of the

material **722** from the received incident light radiation **724**, the SCSS **100**, FIG. 1 performs a self-calibration procedure.

The SCSS **100**, FIG. 1, performs a self-calibration procedure prior to measuring any of the properties of the material **722**. This self-calibration procedure includes emitting a pulse of light radiation **716** from LED **710** that is received as incident light radiation **724** by wavelength sensor **704** prior to inserting material **722** between the probe light source **702** and wavelength sensor **704**. The oximeter utilizes the measured incident light radiation **724** received by wavelength sensor **704** to determine the operating wavelength of LED **710**. Once the operating wavelength of LED **710** is known, the SCSS **100**, FIG. 1, is utilized in combination with the oximeter to accurately determine blood oxygen saturation of the material **722**.

FIG. 8 illustrates a cross-sectional view of the wavelength sensor **704** receiving incident light radiation **724** utilizing a double diffusion photodiode (also known as a double junction photodiode). Photodiodes with double diffusion are typically utilized to accurately measure the centroid wavelength of light sources such as LEDs **710** and **712**. Double diffusion photodiodes are processed with two junctions, one on the top surface and one on the back surface of a semiconductor photodiode (such as a Si-photodiode), each junction typically exhibits a different and well-defined spectral response. As result, by measuring the quotient of signals generated by the two junctions, the centroid wavelength of any given monochromatic light source may be determined.

The wavelength sensor **704** has two p-n junctions constructed vertically on a common silicon substrate. The wavelength sensor **704** includes a first anode **800**, common cathode **802**, first diode **804** (also known as an upper diode), second diode **806** (also known as a lower diode), second anode **808**, and a thin active region **810**. The first anode **800** is positioned on the top surface above the common cathode **802** forming the first diode **804**. The thickness of the first diode **804** is chosen so that the energy of the shortest wavelength being measured from the incident light radiation **724** is absorbed entirely therein. The second diode **806** is formed between the common cathode **802** and the second anode **808** that placed on the bottom surface with the thin active region **810** between the common cathode **802** and the second anode **808**. The thickness of the thin active region **810** is selected to allow for absorption of substantially all of the longest measured wavelength of incident light radiation **724**.

FIG. 9 illustrates a typical plot **900** of the spectral response of the wavelength sensor **704**, FIG. 8. The plot **900**, FIG. 9, has a vertical axis **902** representing relative response, in percentage, of the wavelength sensor **704**, FIG. 8, and a horizontal axis **904**, FIG. 9, representing the wavelength of the incident light radiation **724**, FIG. 8. The plot **900**, FIG. 9, shows two response curves **906** and **908** representing the relative response versus wavelength for the first diode **804**, FIG. 8, and the second diode **806**, respectively.

As an example of operation of the wavelength sensor **704**, the first diode **804** may have an enhanced blue response and the second diode **806** may have an enhanced red response. In this example, the absorbed radiation of the incident light radiation **724** between the red and blue responses (such as between 450 and 900 nm) generates two photocurrent signals proportional to the wavelength of the incident light radiation **724**. The quotient of these photocurrent signals is independent of the light level up to the saturation point of the wavelength sensor **704**. Utilizing this example, the wavelength of either monochromatic incident light radiation **724** or the spectral density peak of polychromatic incident light

radiation **724** may be determined. An example of the wavelength sensor **704** may be a PSS WS-7.56 wavelength sensor produced by Pacific Silicon Sensor, Inc. of Westlake Village, Calif.

In FIG. 10, a schematic diagram depicting an exemplary implementation of calibration circuit **1000** is shown. The calibration circuit **1000** is in signal communication with the probe **1002** and controller **106**, FIG. 1, via signal paths **112** and **114**, respectively. The calibration circuit **1000** may include a pair of amplifiers **1004** and **1006** (such as log amplifiers) in signal communication with first anode **800**, FIG. 8 and second anode **808** of wavelength sensor **1024**, FIG. 10, and a differential amplifier **1008**, via signal paths **1010**, **1012**, and **1014**, respectively. The differential amplifier **1008** is in signal communication with the controller **106**, FIG. 1, via signal path **112**.

In operation, the wavelength sensor **1024** produces two photocurrent signals from the two junctions (i.e., photodiodes **804** and **806**) in the double diffusion photodiode. Each junction in the wavelength sensor **1024** exhibits a different and well-defined spectral response, which is known to the controller **106**, FIG. 1, and the magnitude of these two resulting photocurrent signals are proportional to the wavelength of the measured incident light radiation **724**, which correspond to one of the light sources (either **204** or **206**, FIG. 2) in probe **1002**, FIG. 10. The photocurrent signals are amplified by amplifiers **1004** and **1006** via signal paths **1010** and **1012**, respectively, and input into the differential amplifier **1008** via signal path **1018** and **1020**. If the amplified photocurrent signals **1018** and **1020** are approximately equal the corresponding differential output signal **1022** of the differential amplifier **1008** is almost equal to zero. Once the differential output signal **1022** is almost equal to zero the wavelength of the incident light radiation is determined and the SCSS **100**, FIG. 1, is calibrated.

When the amplified photocurrent signals **1018** and **1020** are not approximately equal the corresponding differential output signal **1022** will vary according to the difference in magnitude value between the amplified photocurrent signals **1018** and **1020**. The differential output signal **1022** is the utilized as a reference by the controller **106**, FIG. 1. The controller **106** determines the wavelength of the incident light radiation **724** by knowing the spectral response of the photodiodes **804** and **806**, FIG. 8. The controller **106** either determines the wavelength of the incident light radiation **724** utilizing software **108** or other hardware (not shown) located in the SCSS **100**. The software **108** may include logic that allows the controller **106** to calculate the wavelength values in real-time from the measure values received from the wavelength sensor **1004**.

Alternatively, the controller **106** may determine the wavelength of the incident light radiation **724** utilizing the lookup ("LUT") table **110**. The LUT **110** may be resident in memory (not shown) resident either internally or externally to the controller **106**. The LUT **110** includes a tabulation of known spectral response in voltage versus wavelength for each photodiode **804** and **806**, FIG. 8. Once the controller **106** measures the differential output signal **1022**, FIG. 10, the software **108**, FIG. 1, compares the value of the differential output signal **1022**, FIG. 10, against values stored in the LUT **110**, FIG. 1, and then retrieves a corresponding wavelength value. The controller **106** then utilizes the retrieved wavelength wave to self-calibrate the SCSS **100**.

Besides self-calibration, the SCSS **100** is also capable of temperature compensating for variation in the wavelength of the incident light radiation **724** due to temperature varia-

tions. The SCSS 100 may compensate for temperature variations by the same process utilized to self-calibrate.

FIG. 11 is another exemplary implementation of the SCSS 100, FIG. 1, with the calibration circuit 1100, FIG. 1, in an oximeter device 1102. The oximeter device 1102 is in signal communication with probe 1104, via signal path 112, and includes calibration circuit 1100, controller 1106, first driver 1108 and second driver 1110. The probe 1104 includes wavelength sensor 1112 and probe light source 1114 having first light source 1116 and second light source 1118.

In operation, the first driver 1108 drives the first light source 1116 and the second driver 1110 drives the second light source 1118. First light source 1116 and the second light source 1118 may individually produce light radiation which is incident of the wavelength sensor 1112. The wavelength sensor 1112 produces two photocurrent signals from the two junctions (i.e., photodiodes 804 and 806) in the double diffusion photodiode. Again, each junction in the wavelength sensor 1112 exhibits a different and well-defined spectral response, which is known to the controller 1106 and the magnitude of these two resulting photocurrent signals are proportional to the wavelength of the measured incident light radiation, which corresponds to one of the light sources (either 1116 or 1118) in probe 1104. The photocurrent signals 1120 and 1122 processed and input into the differential amplifier 1224. If the photocurrent signals 1120 and 1122 are approximately equal the corresponding differential output signal 1126 of the differential amplifier 1124 is almost equal to zero. Once the differential output signal 1126 is almost equal to zero the wavelength of the incident light radiation is determined and the SCSS 100, FIG. 1, is calibrated.

When the photocurrent signals 1120 and 1122 are not approximately equal the corresponding differential output signal 1126 will vary according to the difference in magnitude value between the photocurrent signals 1120 and 1122. The differential output signal 1126 is the utilized as a reference by the controller 1106. The controller 1106 determines the wavelength of the incident light radiation by knowing the spectral response of the photodiodes 804 and 806.

The controller 1106 either determines the wavelength of the incident light radiation utilizing software 108, FIG. 1, or other hardware (not shown) located in the SCSS 100. The software 108 may include logic that allows the controller 1106, FIG. 11, to calculate the wavelength values in real-time from the measure values received from the wavelength sensor 1112.

Alternatively, the controller 1106 may determine the wavelength of the incident light radiation utilizing the lookup LUT 110, FIG. 1. The LUT 110 may be resident in memory (not shown) resident either internally or externally to the controller 1106, FIG. 11. The LUT 110, FIG. 1, includes the tabulation of known spectral response in voltage versus wavelength for each photodiode 804 and 806. Once the controller 1106 measures the differential output signal 1126, FIG. 11, the software 108, FIG. 1, compares the value of the differential output signal 1126, FIG. 11, against values stored in the LUT 110, FIG. 1, and then retrieves a corresponding wavelength value. The controller 1106, FIG. 11, then utilizes the retrieved wavelength wave to self-calibrate the SCSS 100, FIG. 1.

FIG. 12 illustrates the process performed by the SCSS 100, FIG. 1. The process begins in step 1200, FIG. 12. In step 1202, the wavelength sensor 202, FIG. 2, receives incident light radiation 200 from the probe light source 200. Within the wavelength sensor 202, the first diode 804, FIG.

8, receives short wavelengths from the incident light radiation 200, in step 1204, FIG. 12, and the second diode 806, FIG. 8, receives long wavelengths from the incident light radiation 200 in step 1206, FIG. 12. In step 1208, the first diode 804 produces a first photocurrent signal 1010, FIG. 10, in response to receiving short wavelengths from the incident light radiation 200 and the second diode 806 produces a second photocurrent signal 1012, FIG. 10, in response to receiving short wavelengths from the incident light radiation 200 in step 1210, FIG. 12. Finally, in step 1212, the calibration circuit 104 and/or controller 106, FIG. 1, determine the wavelength of the incident light radiation 200 by comparing the first photocurrent signal 1010 to the second photocurrent signal 1012. The process then ends in step 1214.

The SCSS 100 may be selectively implemented in software, hardware, or a combination of hardware and software. For example, the elements of the SCSS 100 may be implemented in software 108 stored in a memory (not shown) located in a controller 106. The controller 106 may be in signal communication with a DSP or ASIC chip via communication link 112 (which may selectively be a system bus). The software 108 configures and drives the DSP or ASIC chip and performs the steps illustrated in FIG. 12.

The software 108 comprises an ordered listing of executable instructions for implementing logical functions. The software 108 may be embodied in any computer-readable medium for use by or in connection with an instruction execution system, apparatus, or device, such as a computer-based system, processor-containing system, or other system that may selectively fetch the instructions from the instruction execution system, apparatus, or device and execute the instructions. In the context of this document, a "computer-readable medium" is any means that may contain, store, communicate, propagate, or transport the program for use by or in connection with the instruction execution system, apparatus, or device. The computer readable medium may be for example, but not limited to, an electronic, magnetic, optical, electromagnetic, infrared, or semiconductor system, apparatus, device, or propagation medium. More specific examples (a non-exhaustive list) of the computer-readable medium would include the following: an electrical connection (electronic) having one or more wires, a portable computer diskette (magnetic), a RAM (electronic), a read-only memory (ROM) (electronic), an erasable programmable read-only memory (EPROM or Flash memory) (electronic), an optical fiber (optical), and a portable compact disc read-only memory (CDROM) (optical). Note that the computer-readable medium may even be paper or another suitable medium upon which the program is printed, as the program can be electronically captured, via for instance optical scanning of the paper or other medium, then compiled, interpreted or otherwise processed in a suitable manner if necessary, and then stored in a computer memory.

Turning to FIG. 13, in an embodiment, a non-invasive medical sensor is provided for detecting blood constituents of any kind. For instance, the sensor 1310 can be for detecting oxygen saturation of hemoglobin or other parameters.

The sensor 1310 can, but not necessarily, have one or more monochromatic light sources 1312, 1313, one or more broadband, non-monochromatic light source 1314 and one or more light detectors 1316. In an embodiment, the light sources and the detectors are mounted oppose to each other as shown in FIG. 13.

In an embodiment, the sensor 1310 is placed on a finger 1318 or other extremity of the human body and shines light

from the one or more light sources through the extremity. FIG. 14 provides a depiction of a typical spectral transmission of both oxygenated (arterial) hbO₂ and deoxygenated (venous) hb haemoglobin molecules (hg). Alternatively, the light is reflected off the skin of a part of the human body, instead of being shined through the extremity.

As indicated previously, the sensor 1310 includes a wavelength detector 1316, such as a dual photodiode or any other type, which produces an output indicating the centroid wavelength of the received light. FIG. 15 depicts responsiveness of a dual junction wavelength detector for both hbO₂ (arterial) and hb (venous) blood flow using a broad spectrum light source.

The sensor 1310 can further include a light source that emits light of a wide light spectrum from, for instance, 200 nm to 2000 nm or any portion thereof, such as a white LED or any other broad emission device. The sensor 1310 can be connected to a processing system 1610 (FIG. 16) that drives the light sources, including the broad emission device, and also receives signals from the photodiodes and the wavelength detector.

In an embodiment, the processing system 1610 contains, among other things, computer media 1612, memory 1614, and signal processing circuitry 1616 used to calculate various parameters of the blood stream through comparison of the signals received by the photodiodes of the sensor 1310. Moreover, the processing system 1610 can be used to illuminate the tissue with light generated from the broad emission light source within the sensor.

Moreover, in an embodiment, the wavelength detector 1316 generates a signal proportional to the strongest transmission wavelength of the light emitted from the broad emission light source 1314 after the light either passes through the body part 1318 or is reflected from the skin of a human. Turning to FIG. 17, a signal derived from the wavelength detector 1316 can be further processed by circuitry and computer processing, via the processing system 1610, to obtain a pulse waveform 1710 proportional to the arterial and venous pulse of the human body.

Via the processing system 1610, the time domain and amplitude properties of both arterial and venous pulse can be stored and further used to synchronize an arterial-pulse synchronized measurement of the absorption or reflectance of one or more monochromatic light-sources 1312, 1313 through the body's extremity 1318. These signals can then be used in calculating oxygen saturation, carbon-monoxide concentration, haemoglobin or any other parameter of the blood.

In an embodiment, the arterial and venous pulse coinciding wavelength detector signals can be used to eliminate or reduce false pulses commonly caused by motion of the human body or extremity, commonly labeled motion artifact, that could cause erroneous measurements.

Utilizing the non-invasive sensor shown in FIG. 13 may allow for the system to make additional measurements and calculations as well.

As discussed herein, the sensor may contain at least one monochromatic light source, at least one non-monochromatic broad spectrum emission light source, at least one light detector, and at least one wavelength detector. The light sources and detectors may be mounted opposing each other, or may be mounted adjacent each other. The sensor may be placed on tissue like a finger (as shown in FIG. 13) or other extremity, or any other position that provides sufficient blood perfusion for measurement for measurement. When the light sources are mounted opposite the detectors, perfusion may result from emitted light passing through the finger

or other body part. When mounted adjacent, the emitted light may be reflected off the skin or other body part, which may provide adequate blood perfusion for measurement.

As already described herein, the detector module may be a dual element light receiver or a dual junction detector element. The dual detector module utilized in the system may, for example, include two photodiodes (detector A and detector B) with different and tightly controlled spectral response curves. The photodiodes may be constructed, for example, on a single chip epitaxial silicon die. The system relies on the characteristics of the detector module to derive a signal proportional to the centroid wavelength of impinging light. A typical response curve of the detector can be seen, for example, in FIG. 18 where curve 1810 shows the responsive curve of detector A, while curve 1820 shows the response curve of detector B.

The sensor and system may include at least one or more light sources which emit non-monochromatic light energy of a broad spectrum, like for example, a white LED. The broad spectrum may cover the range of 200 nm to 1300 nm or any portion thereof. The sensor and system may further include one or more monochromatic light elements. For example, the sensor and system may include a red LED and an IR emitter which emit light to be received by the detector module through or across an extremity or tissue.

The sensor may be connected to a processing system which drives the light sources in a sequential, pulsed mode. The processing system may receive signals from the light receiver elements and the centroid wavelength detector. The processing system may include computer media, memory, and signal processing circuitry which is used to process the signals derived from light detectors contained in the sensor and derive various physiological parameters from these signals. The processing system may also provide control signals to the sensor. The signals may be used to illuminate the light sources which in turn illuminate the tissue or extremity in contact with the sensor. Illumination may occur from the at least one monochromatic light source as well as any one non-monochromatic light sources.

The photodiodes and centroid wavelength detector module may generate a signal proportional to the highest light intensity wavelength as filtered and either passed through an extremity or tissue, or reflected through the skin or tissue. The signal may be further processed to determine modulation in the signal proportional to the arterial pulse and spectral absorption changes between the peak and trough of the arterial pulse. The signal received from the sensor may be further used to correlate changes in spectral absorption to the pulse by pulse blood oxygen concentration as well as other blood constituents.

FIG. 19 represents the change in spectral absorption generated by the arterial pulse blood flow with a broad emission illumination emitter as detected by a centroid wavelength detector and processed by the system described herein. FIG. 20 depicts the peak and valley of such signal in the time domain, the valley coinciding with the peak of the arterial pulse of a patient.

The benefit to using the present system and method for determining blood oxygenation, for example, can be seen in FIG. 21 which shows the signal proportional to blood oxygen saturation derived and processed from centroid wavelength detector ratios (2110) and the pulse by pulse blood oxygen saturation derived from monochromatic light sources used in traditional oximeters (2120). As seen in FIG. 21, utilizing the centroid wavelength detector ratios provides a reasonably accurate and dynamically updated oxygen saturation measurement.

Utilizing the pulse blood oxygen saturation data from the non-monochromatic light element, a reliable data stream for blood oxygen concentrations or other blood constituent parameters during motion events occurring at the sensor site may be obtained and calculated. Events that normally would mask or invalidate data derived from only monochromatic illumination of tissue can be corrected or replaced. The substitute data derived from the changes in spectral absorption of the non-monochromatic illuminated tissue derived from a sensor for data normally derived from monochromatic light sources such as found in traditional pulse-oximeter systems may be used to correct or replace inaccurate data from the traditional monochromatic light sources.

FIGS. 22-24 show the process by which pulse detection may be read and provided through a motion event occurring with the patient. The non-monochromatic broad spectrum light emitting element may be illuminated and the emitted light pass through a patient's tissue before being detected and received by the dual detector element. The arterial pulse will change the spectral absorption enough to allow the system to detect the pulse occurring. After an equal gain is set for detectors A and B (2210), the needed analogue data generated by turning on the broad spectrum element periodically, like for example 30 times per second for a brief duration and sampling of the signals at both the detector A and detector B at the same gain (2220). The ratio of these two signals may be calculated (2230) and a result in a generated signal that reflects the average and changes in spectral absorption generated by both the occurrence of the arterial pulse and base line spectral absorption of the tissue. The system will then find the minimum peak signal which corresponds to the arterial pulse (2240) and determine whether or not the minimum peak occurred (2250). If the minimum peak did not occur, the system will reset the equal gain for detectors A and B and resample and recalculate the ratio.

If the minimum peak has occurred, it will sample and hold the ratio (2260). Using the sampled and held Rs, an average ratio of the previous 25 samplings will be continuously updated to create a moving point average. The system will then take the ratio and look up the corresponding SpO2 value from a stored table (2270), shown in FIG. 22B, and determine if a motion flag should have been set (2280). If the currently measured ratio is within the acceptable range in the system, the process will repeat for the next sample. If the measured ratio is not within the acceptable range in the system, the system will substitute the SpO2 value obtained based on the non-monochromatic LED for any monochromatic Red/IR SpO2 value (2290).

The ratio signal is used to establish a fall-back data stream in the event of a motion event occurring that makes it impossible to accurately calculate the oxygen saturation from the AC component ratio of red LED and IR emitter. During normal operation the oxygen saturation data from the red to IR ratio needs to be correlated to a corresponding value (preferably sampled at the arterial peak) of signal level derived from the ratio of the two elements while the broad spectrum light source is on. If a motion even occurs, the system will revert back to that broad spectrum light source derived data stream to generate accurate oxygen saturation data during the motion event by substituting that data for the saturation data normally calculated from the red to IR ratio. For this reason, the non-monochromatic broad spectrum light source must be turned on in sequence every time the red LED and IR emitter are activated.

The plot typically derived from tissue illuminated by the non-monochromatic or white LED or on a patient at rest and

normal saturation levels can be seen in FIG. 23, where curve 2310 shows the pleth of detector A and curve 2320 shows the pleth of detector B. The "Ratio" of detected light by photodiode A and photodiode B is also calculated as seen by ratio curve 2330. FIG. 24 shows the acquired pleth and A/B ratio during a rapid extremity wave or movement motion event on a normally saturated extremity. As seen in FIG. 24, the excursions of the pleth amplitude, which a traditional oximeter relies on, are in excess of 21%, the A/B ratio which represents only spectral absorption, changes only 1.5%.

The system may also be configured to measure the centroid wavelength of light produced by the non-monochromatic light source. Measuring the centroid wavelength allows for the system to detect when a probe or device is operating outside a centroid wavelength range in order to insure accuracy. The measurement may occur while the probe is applied to the patient and when the red light source is energized. The centroid wavelength data may be collected periodically, like for example every five (5) seconds or every 150 pulses based on an emitter pulse rate of 30 per second. The centroid data may be calculated by sampling both detector A and B in the dual detector element while the red LED is energized. The ratio of A to B corresponds to the centroid wavelength of the light emitted by the red LED and can be compared to limits embedded in the system and compared to acceptable values that assure accurate saturation measurements. It is not necessary to collect data for the IR emitter as the centroid wavelength of the IR emitter does not have a direct bearing on the accuracy of the saturation calculation. Since the measurement is taken on the patient, the system will detect shifts in the centroid wavelength induced by, for example, substances on the patient's extremity or skin pigmentation, skin defects, or the like. A system message can be generated if the centroid falls outside the limits within the system, alerting the user to replace the probe as readings may be inaccurate. Such readings may result from a motion event, as discussed above, or may be the result of the sensor becoming disengaged from the patient.

FIGS. 25 and 26 show the raw data provided from the wavelength detector using non-monochromatic broad-band (white) light source and non-perfused object and a perfused object. Ratio based amplitudes of 110 or higher indicate a non-perfused object, while amplitudes of 90 or lower indicate the presence of a perfused subject. By setting limits and detecting the raw data in the wavelength detector, the detector can accurately determine whether or not an extremity or tissue is located within the sensor, and whether or not the light is passing through the tissue or being directly provided to the sensor.

While various implementations of the application have been described, it will be apparent to those of ordinary skill in the art that many more embodiments and implementations are possible that are within the scope of the invention. Accordingly, the invention is not to be restricted except in light of the attached claims and their equivalents.

What is claimed is:

1. A method for eliminating error when using a non-invasive optical sensing system comprising:

- providing a first detector and a second detector in a non-invasive optical sensor, the first detector and the second detector having different respective spectral response curves;
- emitting light from a non-monochromatic light source in the non-invasive optical sensor periodically;
- emitting red light and infrared light from one or more monochromatic light sources;

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generating a first oxygen saturation value, based on a ratio of the red light detected by the first and second detectors, and the infrared light detected by the first and second detectors;

generating a second oxygen saturation value, based on the non-monochromatic light detected by the first and second detectors; and

substituting the first oxygen saturation value with the second oxygen saturation value in response to a determination that a ratio of a first measured centroid wavelength of the red light by the first detector and a second measured centroid wavelength of the red light by the second detector is outside of a threshold.

2. The method claim 1 further comprising the step of generating arterial pulse rate data using the first signal and the second signal if a motion event occurs with the tissue located within the non-invasive optical sensor.

3. The method of claim 1 further comprising the steps of: generating a notification if the ratio of the first signal to the second signal is above or below a threshold.

4. The method of claim 1 further comprising the steps of: measuring a centroid wavelength of emitted red light; determining if the measured centroid wavelength is within a centroid range configured to ensure accuracy of the first oxygen saturation measurement;

generating an alert if the measured centroid wavelength is outside the acceptable centroid range.

5. The method claim 1 further wherein the first detector and second detector have different spectral response curves, and the different spectral response curves are utilized to derive a fourth signal proportional to the centroid wavelength of emitted red light.

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6. A system comprising:

a non-monochromatic light source configured to provide non-monochromatic light periodically;

one or more monochromatic light sources configured to provide red light and infrared light;

a wavelength sensor including a first detector with a first spectral response and a second detector with a second spectral response different from the first;

a processor;

non-transitory computer readable media having encoded thereon computer software comprising a set of instructions that when executed by the at least one processor, causes the at least one processor to:

generate a first signal from the first detector and a second signal from the second detector resulting from light periodically emitted by the non-monochromatic light source sampled by the first detector and the second detector;

generate a first oxygen saturation value, based on a ratio of the red light detected by the first and second detectors, and the infrared light detected by the first and second detectors;

generate a second oxygen saturation value, based on the first signal and the second signal; and

substitute the first oxygen saturation value with the second oxygen saturation value in response to a determination that a motion event has occurred, wherein the determination that the motion event has occurred is based, at least in part, on a ratio of the first signal and the second signal.

* * * * *

专利名称(译)	用于非侵入式医学传感器的系统和方法		
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[标]申请(专利权)人(译)	MEDTOR		
申请(专利权)人(译)	MEDTOR LLC		
当前申请(专利权)人(译)	MEDTOR LLC		
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其他公开文献	US20170311855A1		
外部链接	Espacenet		

摘要(译)

公开了一种利用具有一个或多个光源的传感器以及一个或多个光检测器来测量氧气水平和各种血液成分的方法和系统。该系统能够使用由一个或多个检测器从非单色光源收集的数据在传感器末端发生运动事件期间提供准确的信息。该系统也能够检测，并且如果传感器没有正确放置在患者或变得与其脱开提供警报。

