



US010485462B2

(12) **United States Patent**
Benni

(10) **Patent No.:** **US 10,485,462 B2**
(45) **Date of Patent:** **Nov. 26, 2019**

(54) **METHOD FOR SPECTROPHOTOMETRIC BLOOD OXYGENATION MONITORING OF ORGANS IN THE BODY**

(71) Applicant: **CAS Medical Systems, Inc.**, Branford, CT (US)

(72) Inventor: **Paul B. Benni**, Acton, MA (US)

(73) Assignee: **EDWARDS LIFESCIENCES CORPORATION**, Branford, CT (US)

(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 604 days.

(21) Appl. No.: **15/172,992**

(22) Filed: **Jun. 3, 2016**

(65) **Prior Publication Data**
US 2016/0278675 A1 Sep. 29, 2016

Related U.S. Application Data
(62) Division of application No. 13/511,928, filed as application No. PCT/US2010/058059 on Nov. 24, 2010, now Pat. No. 9,364,175.
(Continued)

(51) **Int. Cl.**
A61B 5/1455 (2006.01)
A61B 5/00 (2006.01)

(52) **U.S. Cl.**
CPC **A61B 5/14551** (2013.01); **A61B 5/4836** (2013.01); **A61B 5/4848** (2013.01);
(Continued)

(58) **Field of Classification Search**
CPC A61B 5/1455; A61B 5/14551; A61B 5/14552; A61B 5/72; A61B 5/7221; A61B 2562/0233
See application file for complete search history.

(56) **References Cited**

U.S. PATENT DOCUMENTS

6,766,188 B2 7/2004 Soller
7,010,337 B2 3/2006 Furnary et al.
(Continued)

FOREIGN PATENT DOCUMENTS

EP 0352923 1/1990

OTHER PUBLICATIONS

Kawasuji et al. "Near-Infrared Monitoring of Myocardial Oxygenation During Intermittent Warm Blood Cardioplegia", European Journal of Cardio-Thoracic Surgery, vol. 12, No. 2, Aug. 1, 1997, pp. 236-241.

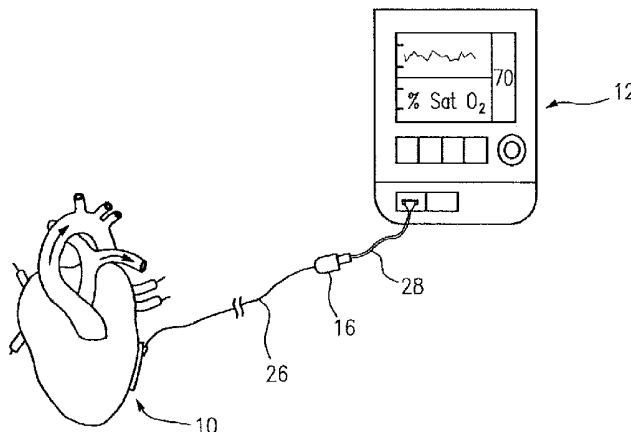
(Continued)

Primary Examiner — Eric F Winakur
(74) *Attorney, Agent, or Firm* — O'Shea Getz P.C.; Richard D. Getz

(57) **ABSTRACT**

A method and apparatus for non-invasively determining a blood oxygen saturation level within an organ of a subject using direct application of a near infrared spectrophotometric sensor is provided. The method includes the steps of: a) transmitting a light signal directly into the subject's organ using the sensor; b) sensing a first intensity of the light signal and a second intensity of the light signal, after the light signal travels a predetermined distance through the organ of the subject; c) determining an attenuation of the light signal along multiple different wavelengths using the sensed first intensity and sensed second intensity; d) determining a difference in attenuation of the light signal between wavelengths; and e) determining the blood oxygen saturation level within the subject's organ using the difference in attenuation between wavelengths.

6 Claims, 4 Drawing Sheets



Related U.S. Application Data

(60) Provisional application No. 61/264,080, filed on Nov. 24, 2009.

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

CPC *A61B 5/7246* (2013.01); *A61B 5/7271* (2013.01); *A61B 5/7278* (2013.01); *A61B 5/746* (2013.01); *A61B 2562/0233* (2013.01)

(56) **References Cited**

U.S. PATENT DOCUMENTS

7,123,950 B2* 10/2006 Mannheimer A61B 5/14551
600/323
2001/0047128 A1 11/2001 Benni
2004/0024297 A1 2/2004 Chen et al.
2004/0073278 A1 4/2004 Pachys
2004/0082841 A1 4/2004 Furnary et al.
2006/0034941 A1 2/2006 Dobson
2006/0167400 A1 7/2006 Ellingboe et al.
2007/0265513 A1 11/2007 Schenkman et al.
2008/0200780 A1 8/2008 Schenkman et al.
2009/0182209 A1 7/2009 Benni

OTHER PUBLICATIONS

Ho et al. "In Vivo Detection of Myocardial Ischemia in Pigs Using Visible Light Spectroscopy", *Anesthesia and Analgesia*, vol. 108, No. 4, Apr. 1, 2009, pp. 1185-1192.
EP office action for EP10833955.7 dated Oct. 25, 2018.

* cited by examiner

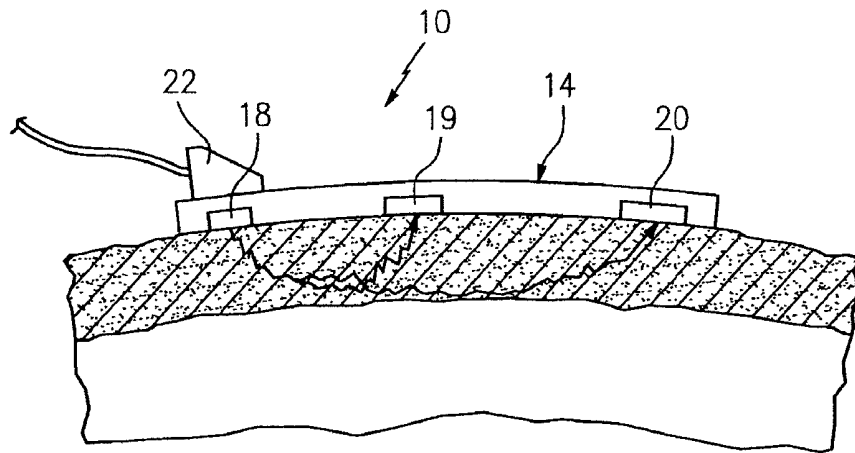


FIG. 1

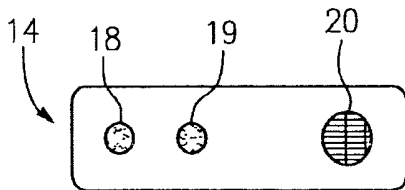


FIG. 2

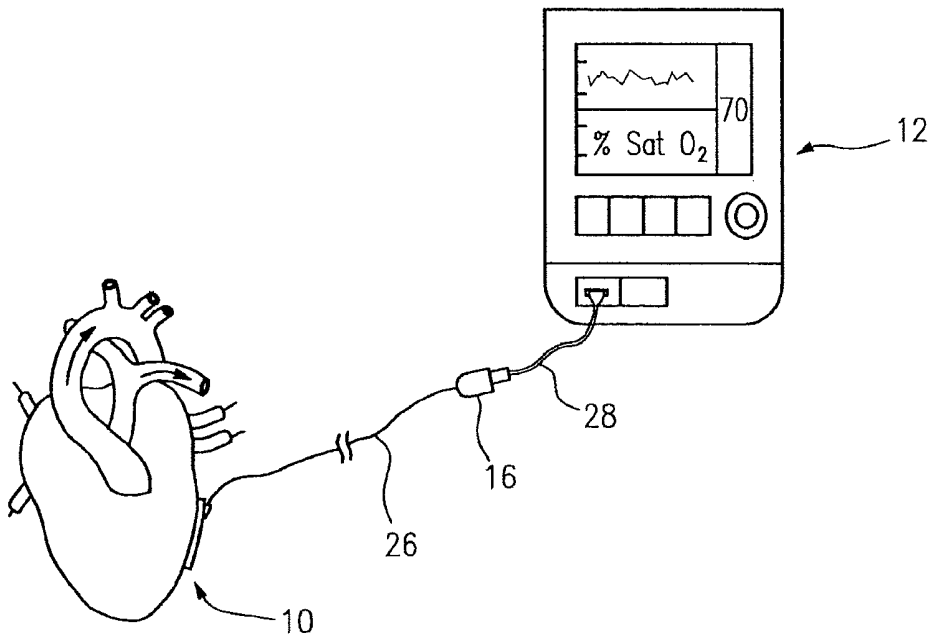


FIG. 3

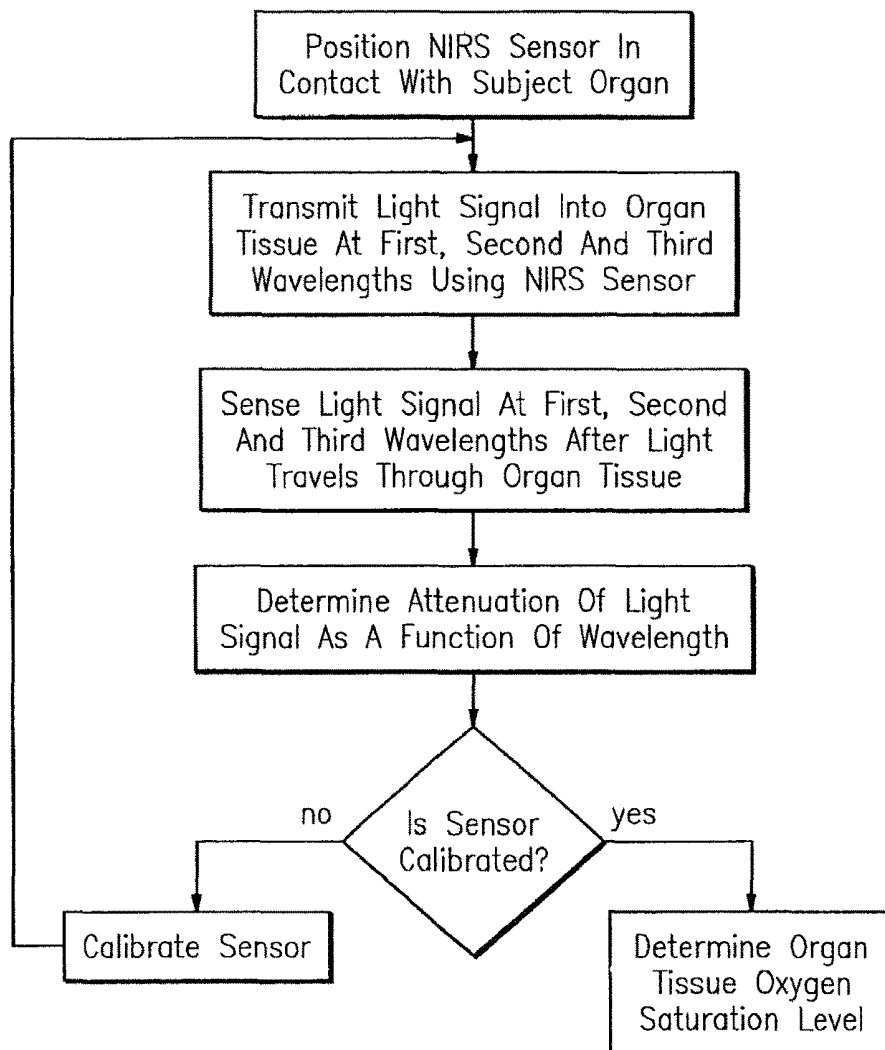


FIG. 4

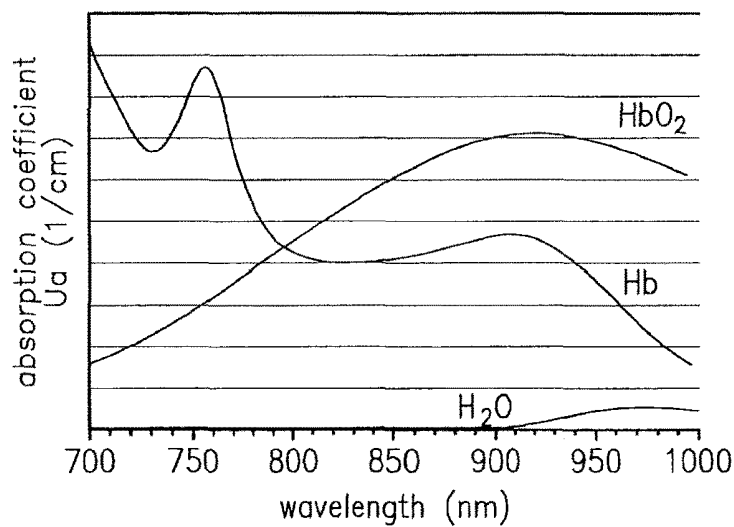


FIG. 5

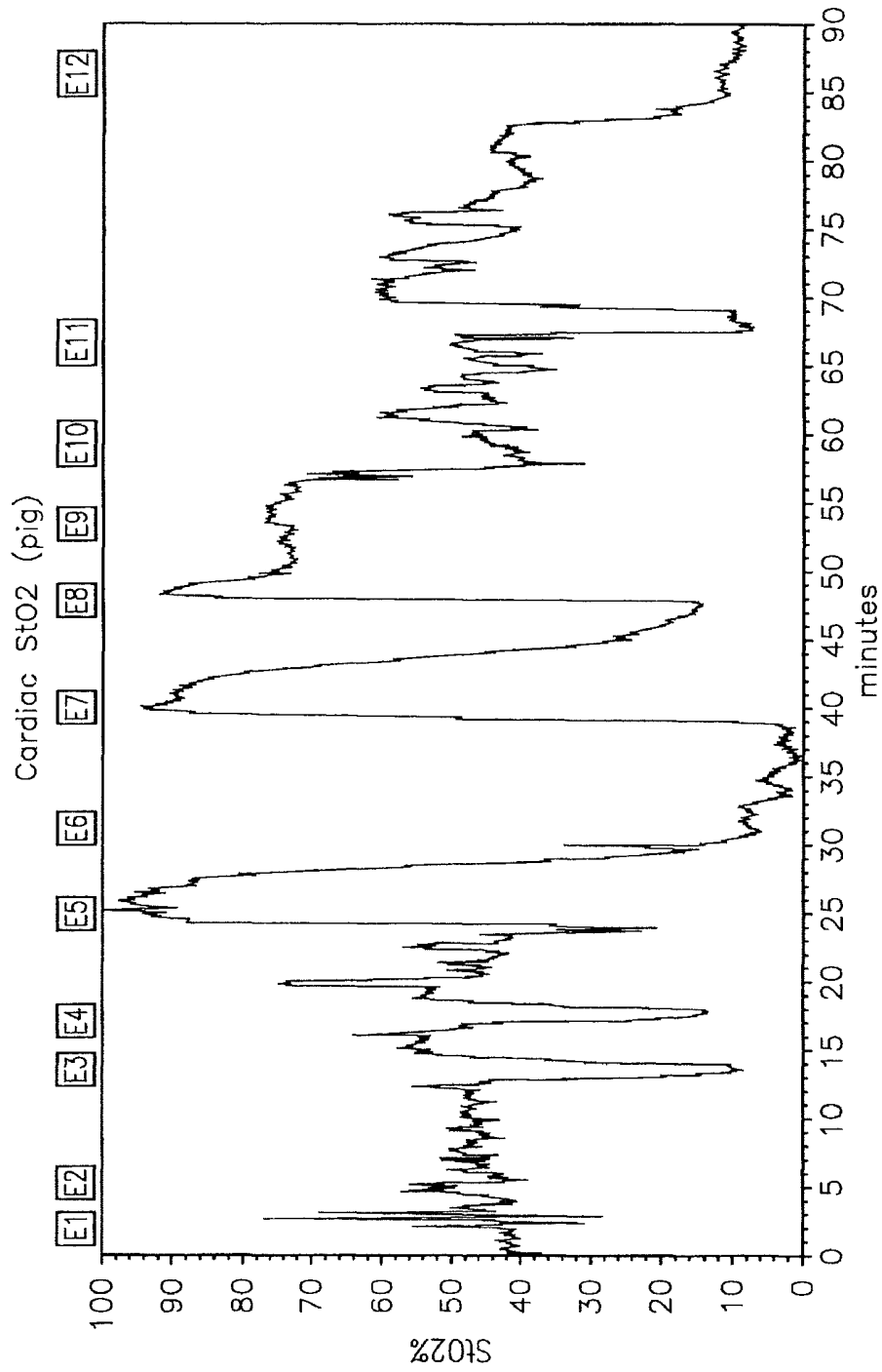


FIG. 6

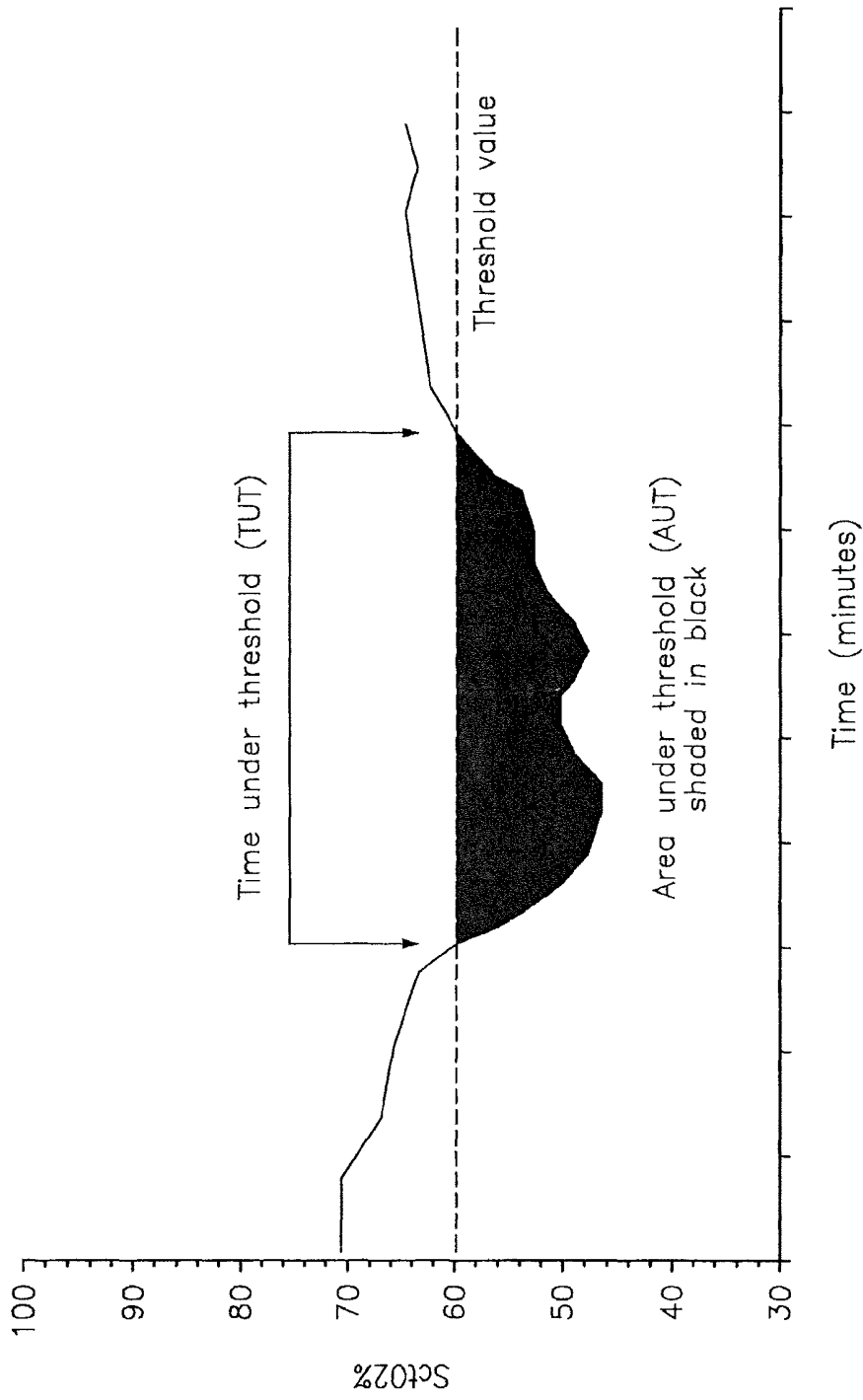


FIG. 7

METHOD FOR SPECTROPHOTOMETRIC BLOOD OXYGENATION MONITORING OF ORGANS IN THE BODY

This application is a divisional of U.S. patent application Ser. No. 13/511,928 filed May 24, 2012, which is a national stage application of PCT Patent Application No. PCT/US2010/058059 filed on Nov. 24, 2010, which claims priority to U.S. Provisional Patent Application No. 61/264,080 filed Nov. 24, 2009, the disclosures of which are herein incorporated by reference.

BACKGROUND OF THE INVENTION

1. Technical Field

This invention relates to methods for non-invasively determining biological tissue oxygenation in general, and to non-invasive methods utilizing near-infrared spectroscopy (NIRS) techniques in particular.

2. Background Information

Oxygen saturation in a mammalian subject can be defined as:

$$\text{O}_2 \text{ saturation } \% = \frac{\text{HbO}_2}{(\text{HbO}_2 + \text{Hb})} * 100\% \quad (\text{Eqn. 1})$$

where HbO₂ refers to oxygenated hemoglobin (i.e., “oxy-hemoglobin”) and Hb refers to deoxygenated hemoglobin (i.e., “deoxyhemoglobin”). In the arterial circulatory system under normal conditions, there is a high proportion of HbO₂ to Hb, resulting in an arterial oxygen saturation (defined as SaO₂%) of 95-100%. After delivery of oxygen to tissue via the capillaries, the proportion of HbO₂ to Hb decreases. Therefore, the measured oxygen saturation of venous blood (defined as SvO₂%) is lower and may be about 70%.

One spectrophotometric method, called pulse oximetry, determines arterial oxygen saturation (SaO₂) of peripheral tissue (i.e. finger, ear, nose) by monitoring pulsatile optical attenuation changes of detected light induced by pulsatile arterial blood volume changes in the arteriolar vascular system. The method of pulse oximetry requires pulsatile blood volume changes in order to make a measurement. Since venous blood is not pulsatile, pulse oximetry cannot provide any information about venous blood.

Near-infrared spectroscopy (NIRS) is an optical spectrophotometric method of continually monitoring tissue oxygenation that does not require pulsatile blood volume to calculate parameters of clinical value. The NIRS method utilizes light in the near-infrared range (700 to 1,000 nm) that can pass easily through skin, bone and other tissues where it encounters hemoglobin located mainly within micro-circulation passages; e.g., capillaries, arterioles, and venuoles. Hemoglobin exposed to light in the near infra-red range has specific absorption spectra that varies depending on its oxidation state; i.e., oxyhemoglobin (HbO₂) and deoxyhemoglobin (Hb) each act as a distinct chromophore. By using light sources that transmit near-infrared light at specific different wavelengths, and measuring changes in transmitted or reflected light attenuation, concentration changes of the oxyhemoglobin (HbO₂) and deoxyhemoglobin (Hb) can be monitored.

The apparatus used in NIRS analysis typically includes a plurality of light sources, one or more light detectors for detecting reflected or transmitted light, and a processor for processing signals that represent the light emanating from

the light source and the light detected by the light detector. Light sources such as light emitting diodes (LEDs) or laser diodes that produce light emissions in the wavelength range of 700-1000 nm at an intensity below that which would damage the biological tissue being examined are typically used. A photodiode or other light source detector is used to detect light reflected from or passed through the tissue being examined. The processor takes the signals from the light sources and the light detector and analyzes those signals in terms of their intensity and wave properties.

It is known that relative changes of the concentrations of HbO₂ and Hb can be evaluated using apparatus similar to that described above, including a processor programmed to utilize a variant of the Beer-Lambert Law, which accounts for optical attenuation in a highly scattering medium like biological tissue. The modified Beer-Lambert Law can be expressed as:

$$A_\lambda = -\log(I/I_0)_\lambda = \alpha_\lambda C * d * B_\lambda + G \quad (\text{Eqn.2})$$

wherein “A_λ” represents the optical attenuation in tissue at a particular wavelength λ (units: optical density or OD); “I₀” represents the incident light intensity (units: W/cm²); “I” represents the detected light intensity; “α_λ” represents the wavelength dependent absorption coefficient of the chromophore (units: OD*cm⁻¹*μM⁻¹); “C” represents the concentration of chromophore (units: μM); “d” represents the light source to detector (optode) separation distance (units: cm); “B_λ” represents the wavelength dependent light scattering differential pathlength factor (unitless); and “G” represents light attenuation due to scattering within tissue (units: OD). The product of “d*B_λ” represents the effective pathlength of photon traveling through the tissue.

Absolute measurement of chromophore concentration (C) is very difficult because G is unknown or difficult to ascertain. However, over a reasonable measuring period of several hours to days, G can be considered to remain constant, thereby allowing for the measurement of relative changes of chromophore from a zero reference baseline. Thus, if time t₁ marks the start of an optical measurement (i.e., a base line) and time t₂ is an arbitrary point in time after t₁, a change in attenuation (ΔA) between t₁ and t₂ can be calculated, and variables G and I₀ will cancel out providing that they remain constant.

The change in chromophore concentration (ΔC=C(t₂)-C(t₁)) can be determined from the change in attenuation ΔA, for example using the following equation derived from the modified Beer-Lambert Law:

$$\Delta A_\lambda = -\log(I_{t2}/I_{t1})_\lambda = \alpha_\lambda * \Delta C * d * B_\lambda \quad (\text{Eqn.3})$$

Presently known NIRS algorithms that are designed to calculate the relative change in concentration of more than one chromophore use a multivariate form of Equation 2 or 3. To distinguish between, and to compute relative concentration changes in, oxyhemoglobin (ΔHbO₂) and deoxyhemoglobin (ΔHb), a minimum of two different wavelengths are typically used. The concentration of the HbO₂ and Hb within the examined tissue is determined in μmoles per liter of tissue (μM).

The above-described NIRS approach to determining oxygenation levels is useful, but it is limited in that it only provides information regarding a change in the level of oxygenation within the tissue. It does not provide a means for determining the absolute value of oxygen saturation within the biological tissue.

At present, information regarding the relative contributions of venous and arterial blood within tissue examined by NIRS is either arbitrarily chosen or is determined by inva-

sive sampling of the blood as a process independent from the NIRS examination. For example, it has been estimated that NIRS examined brain tissue comprising about 60 to 80% blood venous and about 20 to 40% arterial blood. Blood samples from catheters placed in venous drainage sites such as the internal jugular vein, jugular bulb, or sagittal sinus have been used to evaluate NIRS measurements. Results from animal studies have shown that NIRS interrogated tissue consists of a mixed vascular bed with a venous-to-arterial ratio of about 2:1 as determined from multiple linear regression analysis of sagittal sinus oxygen saturation (SssO₂) and arterial oxygen saturation (SaO₂). An expression representing the mixed venous/arterial oxygen saturation (SmvO₂) in NIRS examined tissue is shown by the equation:

$$SmvO_2 = K_v * SvO_2 + K_a * SaO_2 \quad (\text{Eqn.4})$$

where "SvO₂" represents venous oxygen saturation; "SaO₂" represents arterial oxygen saturation; and K_v and K_a are the weighted venous and arterial contributions respectively, with K_v+K_a=1. The parameters K_v and K_a may have constant values, or they may be a function of SvO₂ and SaO₂. Determined oxygen saturation from the internal jugular vein (SijvO₂), jugular bulb (SjbO₂), or sagittal sinus (SssO₂) can be used to represent SvO₂. Therefore, the value of each term in Equation 4 is empirically determined, typically by discretely sampling or continuously monitoring and subsequently evaluating patient arterial and venous blood from tissue that the NIRS sensor is examining, and using regression analysis to determine the relative contributions of venous and arterial blood independent of the MRS examination.

Some medical procedures involve limiting or completely stopping the flow of blood to an organ. For example, some procedures involve isolating the heart from the rest of the body by means of a cross clamp on the aorta and then cold cardioplegia is given into the heart through the aortic root. The cold fluid (usually in the range of about 4-10° C.) ensures that the heart cools down to an approximate temperature of around 15-20° C. thus slowing down the metabolism of the heart and thereby preventing damage to the heart muscle. The process may be further augmented by a cardioplegic component which is high in potassium and magnesium. The potassium helps by arresting the heart in diastole thus ensuring that the heart does not use up the valuable energy stores during this period of heart isolation. Blood can be added to this solution especially for long procedures requiring more than half an hour of ascending aorta cross-clamp time. Blood acts as a buffer and also supplies nutrients to the heart during ischemia. Once the procedure on the heart vessels (e.g., coronary artery bypass grafting, or heart valve replacement, or correction of congenital heart defect, etc.) is over, the cross-clamp is removed and the isolation of the heart is terminated so that normal blood supply to the heart is restored and the heart starts beating again.

During the isolation period, it would be of great value to monitor the oxygen saturation level of the heart to ensure that the heart has a sufficient oxygen level to avoid damage.

The cessation of blood flow to an organ is not limited to hearts, however. For example, livers and kidneys are often removed from a donor for transplant into a recipient. In such cases, there would be value in knowing the oxygen saturation level in the organ before, during, and after the transplant.

What is needed, therefore, is a method for non-invasively determining the level of oxygen saturation within biological

tissue that can determine the absolute oxygen saturation value rather than a change in level, and one that can be used to directly determine the oxygen saturation levels of a body organ.

DISCLOSURE OF THE INVENTION

According to an aspect of the present invention, a method and apparatus for non-invasively determining a blood oxygen saturation level within an organ of a subject using direct application of a near infrared spectrophotometric sensor is provided. The method includes the steps of: a) transmitting a light signal directly into the subject's organ using the sensor along multiple wavelengths; b) sensing a first intensity of the light signal and a second intensity of the light signal, after the light signal travels a predetermined distance through the organ of the subject; c) determining an attenuation of the light signal along multiple third wavelengths using the sensed first intensity and sensed second intensity of wavelengths; d) determining a difference in attenuation of the light signal between multiple wavelengths; and e) determining the blood oxygen saturation level within the subject's organ using the difference in attenuation between the multiple wavelengths.

According to another aspect of the present invention, an apparatus for non-invasively determining a blood oxygen saturation level within an organ of a subject is provided. The apparatus includes at least one near infrared spectrophotometric sensor operable to be placed in contact with the organ, and a processor. The sensor has at least one light source and at least one light detector. The light source is operable to transmit a light signal that includes a first wavelength, a second wavelength, and a third wavelength. The light detector is operable to sense the light signal along the first, second, and third wavelengths after the light signal travels a predetermined distance through the organ tissue. The sensor is operable to produce data signals representative of the sensed light signal. The processor is adapted to receive the data signals from the sensor, and is adapted to determine an attenuation of the light signal for each of the first, second, and third wavelengths. The processor is further adapted to determine the blood oxygen saturation level within the organ tissue using the difference in attenuation.

According to another aspect of the present invention, a method for regulating a tissue oxygenation level within a mammalian organ is provided. The method includes the steps of: a) providing an organ tissue oxygenation level threshold; b) applying a near infrared spectrophotometric sensor onto a region of the organ, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within the organ tissue using the data signals; c) sensing the oxygen saturation level within the organ tissue using the spectrophotometric sensor, and producing a sensed organ tissue oxygenation level value using the processor; d) comparing the organ tissue oxygenation level threshold and the sensed organ tissue oxygenation level value; and e) selectively treating the organ to increase the tissue oxygenation level within the organ based on the comparison.

According to another aspect of the present invention, a method for monitoring the effects of cardioplegia administered to a heart is provided. The method includes the steps of: a) applying a near infrared spectrophotometric sensor onto a region of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation

level within heart tissue using the data signals; b) sensing the oxygen saturation level within the heart tissue using the spectrophotometric sensor before an administration of a cardioplegia to the heart, and producing a pre-administration sensed heart tissue oxygenation level value using the processor; c) administering a cardioplegia to the heart; and d) sensing the oxygen saturation level within the heart tissue using the spectrophotometric sensor after the administration of the cardioplegia to the heart, and producing a post-administration sensed heart tissue oxygenation level value using the processor.

These and other objects, features, and advantages of the present invention method and apparatus will become apparent in light of the detailed description of the invention provided below and the accompanying drawings. The methodology and apparatus described below constitute a preferred embodiment of the underlying invention and do not, therefore, constitute all aspects of the invention that will or may become apparent by one of skill in the art after consideration of the invention disclosed overall herein.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a diagrammatic side representation of a NIRS sensor disposed on the surface of an organ.

FIG. 2 is a diagrammatic planar view of a NIRS sensor.

FIG. 3 is a diagrammatic representation of a NIRS system, including a sensor placed on an organ of a subject.

FIG. 4 is a block diagram of the present methodology.

FIG. 5 is a graph showing an exemplary plot of absorption coefficient vs. wavelength.

FIG. 6 graphically illustrates case study data in the form of StO₂ value versus time.

FIG. 7 is a graphical illustration depicting TUT and AUT regions.

DETAILED DESCRIPTION THE INVENTION

The present method of and apparatus for non-invasively determining the blood oxygen saturation level within a subject's tissue is provided that utilizes a near infrared spectrophotometric (NIRS) sensor that includes a transducer capable of transmitting a light signal into the tissue of a subject and sensing the light signal once it has passed through the tissue via transmittance or reflectance. The present method and apparatus can be used with a variety of NIRS sensors. The present method is not limited to use with this preferred NIRS sensor, however.

Referring to FIGS. 1-5, the preferred NIRS sensor includes a transducer portion 10 and processor portion 12. The transducer portion 10 includes an assembly housing 14 and a connector housing 16. The assembly housing 14, which is a flexible structure that can be attached directly to a subject's body, includes one or more light sources 18 and one or more light detectors 19, 20. A sterile disposable envelope or pad may be used to mount the assembly housing 14 to the subject's skin or organ. Light signals of known but different wavelengths from the light sources 18 emit through a prism assembly 22. The light sources 18 are preferably laser diodes that emit light at a narrow spectral bandwidth at predetermined wavelengths. Alternative light emitting diodes (LEDs) can be used if the algorithm is drafted to consider the LEDs characteristic broad spectral bandwidth emitted light. In one embodiment, the laser diodes are mounted within the connector housing 16. The laser diodes are optically interfaced with a fiber optic light guide to the prism assembly 22 that is disposed within the assembly

housing 14. In a second embodiment, the light sources 18 are mounted within the assembly housing 14. A first connector cable 26 connects the assembly housing 14 to the connector housing 16 and a second connector cable 28 connects the connector housing 16 to the processor portion 12. The light detectors 19, 20 each include one or more photodiodes, which are operably connected to the processor portion 12 via the first and second connector cables 26, 28. The processor portion 12 includes a processor for processing light intensity signals from the light sources 18 and one or both light detectors 19, 20.

The processor 12 includes a central processing unit (CPU) and is adapted (e.g., programmed) to selectively perform the functions necessary to perform the present method. It should be noted that the functionality of processor 12 may be implemented using hardware, software, firmware, or a combination thereof. A person skilled in the art would be able to program the processor 12 to perform the functionality described herein without undue experimentation. The processor 12 utilizes an algorithm that characterizes a change in attenuation as a function of the difference in attenuation between different wavelengths. In some embodiments, the present method accounts for but minimizes the effects of pathlength and parameter "E", that latter of which represents energy losses (i.e. light attenuation) due to light scattering within tissue (G), other background absorption losses from biological compounds (F), and other unknown losses including measuring apparatus variability (N). E=G+F+N. Multi-tissue layer organs such as the kidney benefit from multi-detector NIRS sensors and algorithms. In some embodiments, an alternative algorithm may be used that considers fewer or different energy loss components; e.g., if a subject heart is being directly sensed for oxygen saturation, an alternative algorithm may be used that recognizes the absence of certain types of tissue (e.g., skin, bone, etc.) and the signal energy losses associated therewith.

Refer to FIG. 1, the absorption A_{bλ}, detected from the deep light detector 20 comprises of attenuation and energy loss from both the deep and shallow tissue, while the absorption A_{sλ}, detected from the shallow light detector 19 comprises of attenuation and energy loss from shallow tissue only. Absorptions A_{bλ} and A_{sλ} can be expressed in the form of Equation 5 and Equation 6 below which is a modified version of Equation 2 that accounts for energy losses due to "E":

$$A_{b\lambda} = -\log(I_b/I_o)_\lambda = \alpha_\lambda * C_b * L_b + \alpha_\lambda * C_s * L_s + E_\lambda \tag{Eqn.5}$$

$$A_{s\lambda} = -\log(I_s/I_o)_\lambda = \alpha_\lambda * C_s * L_s + E_\lambda \tag{Eqn.6}$$

As indicated above, in some applications (e.g., direct organ sensing, where the signal does not traverse skin or bone) it may be possible to sense using a sensor with a single detector. In those instances, the following equation can be used to represent absorption:

$$A_{b\lambda} = -\log(I_b/I_o)_\lambda = \alpha_\lambda * C_b * L_b + E_\lambda \tag{Eqn. 7}$$

Substituting Equation 6 into Equation 5 yields A'_λ, which represents attenuation and energy loss from deep tissue only:

$$A'_\lambda = A_{b\lambda} - A_{s\lambda} = \alpha_\lambda * C_b * L_b + (E_\lambda - E_{s\lambda}) = -\log\left(\frac{I_b}{I_s}\right)_\lambda \tag{Eqn. 8}$$

Where L is the effective pathlength of the photon traveling through the deep tissue and A'₁ and A'₂ are the absorptions of two different wavelengths. Let E'_λ=E_λ-E_{sλ}, therefore:

$$A'_1 - A'_2 = \Delta A'_{12} \quad (\text{Eqn. 9})$$

Substituting Equation 7 or 8 into Equation 9 for A'_1 and A'_2 , $\Delta A'_{12}$ can be expressed as:

$$\Delta A'_{12} = \alpha_{r,12} * C_b * L_b + \Delta E'_{12} \quad (\text{Eqn. 10})$$

and rewritten Equation 10 in expanded form:

$$\begin{aligned} \Delta A'_{12} = & (\alpha_{r1} - \alpha_{r2})[Hb]_b + (\alpha_{o1} - \alpha_{o2})[HbO_2]_b * L_b + (E'_1 - E'_2) = \\ & (\Delta \alpha_{r12} * [Hb]_b * L_b) + (\Delta \alpha_{o12} * [HbO_2]_b * L_b) + \Delta E'_{12} \end{aligned} \quad (\text{Eqn. 11})$$

where:

$(\Delta \alpha_{r12} * [Hb]_b * L_b)$ represents the attenuation attributable to Hb;

$(\Delta \alpha_{o12} * [HbO_2]_b * L_b)$ represents the attenuation attributable to HbO₂; and

$\Delta E'_{12}$ represents energy losses (i.e. light attenuation) due to light scattering within tissue, other background absorption losses from biological compounds, and other unknown losses including measuring apparatus variability. As indicated above, the specific energy losses accounted for can be varied to suit the application at hand; e.g., inclusion or exclusion of energy losses attributable to skin and bone, use of a single detector versus a pair of detectors, etc.

The multivariate form of Equation 10 is used to determine $[HbO_2]_b$ and $[Hb]_b$ with three different wavelengths:

$$\begin{bmatrix} \Delta A'_{12} - \Delta E'_{12} \\ \Delta A'_{13} - \Delta E'_{13} \end{bmatrix} (L_b)^{-1} = \begin{bmatrix} \Delta \alpha_{r12} & \Delta \alpha_{o12} \\ \Delta \alpha_{r13} & \Delta \alpha_{o13} \end{bmatrix} \begin{bmatrix} [Hb]_b \\ [HbO_2]_b \end{bmatrix} \quad (\text{Eqn. 12})$$

(Eqn. 12)

Rearranging and solving for $[HbO_2]_b$ and $[Hb]_b$, simplifying the $\Delta \alpha$ matrix into $[\Delta \alpha']$:

$$\begin{bmatrix} \Delta A'_{12} \\ \Delta A'_{13} \end{bmatrix} [\Delta \alpha']^{-1} (L_b)^{-1} = \begin{bmatrix} \Delta E'_{12} \\ \Delta E'_{13} \end{bmatrix} [\Delta \alpha']^{-1} (L_b)^{-1} = \begin{bmatrix} [Hb]_b \\ [HbO_2]_b \end{bmatrix} \quad (\text{Eqn. 13})$$

(Eqn. 13)

Then combined matrices $[\Delta A'] [\Delta \alpha']^{-1} = [A_c]$ and $[\Delta E'] [\Delta \alpha']^{-1} = [\Psi_c]$:

$$\begin{bmatrix} A_{Hb} \\ A_{HbO_2} \end{bmatrix} (L_b)^{-1} = \begin{bmatrix} \Psi_{Hb} \\ \Psi_{HbO_2} \end{bmatrix} (L_b)^{-1} = \begin{bmatrix} [Hb]_b \\ [HbO_2]_b \end{bmatrix} \quad (\text{Eqn. 14})$$

(Eqn. 14)

The parameters A_{Hb} and A_{HbO_2} represent the product of the matrices $[\Delta A_c]$ and $[\Delta \alpha']^{-1}$ and the parameters Ψ_{Hb} and Ψ_{HbO_2} represent the product of the matrices $[\Delta E'_c]$ and $[\Delta \alpha']^{-1}$. To determine the level of organ blood oxygen saturation (StO₂), Equation 14 is rearranged using the form of Equation 1 and is expressed as follows:

$$\text{StO}_2 \% = \frac{(A_{HbO_2} - \Psi_{HbO_2})}{(A_{HbO_2} - \Psi_{HbO_2} + A_{Hb} - \Psi_{Hb})} * 100\% \quad (\text{Eqn. 15})$$

Note that the effective pathlength L_b cancels out in the manipulation from Equation 14 to Equation 15.

The value for StO₂ is initially determined from SmvO₂ using Equation 4 and the empirically determined values for SvO₂ and SaO₂, determined from the organ to be monitored. The empirically determined values for SvO₂ and SaO₂ are based on data developed by discrete sampling (e.g., collections of blood sample at discrete points in time) or continuous monitoring (e.g., sensing blood flow through a catheter) of the subject's blood from the arterial input and venous output of the organ to be monitored, performed at or about the same time as the sensing of the organ tissue with the sensor. The temporal and physical proximity of the NIRS sensing and the development of the empirical data helps assure accuracy. For example in calibrating the NIRS sensor to the heart, SaO₂ can be determined from blood in the aorta (arterial blood enters to coronary arteries of the heart via the aorta) or any systemic artery. Heart SvO₂ will be determined from blood drawn from the coronary venous drainage of the heart. The initial values for Kv and Ka within Equation 4 are clinically reasonable values for the circumstances at hand. Alternatively, only SvO₂ may be used for empirical calibration (i.e., Ka=0, Kv=1). The values for A_{HbO_2} and A_{Hb} are determined mathematically using the values for $I_{b\lambda}$ and $I_{x\lambda}$ for each wavelength sensed with the NIRS sensor (e.g., using Equation 5 and 6). The calibration parameters Ψ_{Hb} and Ψ_{HbO_2} , which account for energy losses due to scattering as well as other background absorption from biological compounds, are then determined using Equation 14 and non-linear regression techniques by correlation to different weighted values of SvO₂ and SaO₂; i.e., different values of Ka and Kv. Statistically acceptable values of Kv and Ka and Ψ_{Hb} and Ψ_{HbO_2} are converged upon using the non-linear regression techniques. Experimental findings show that after proper selection of Ka and Kv, the calibration parameters Ψ_{Hb} and Ψ_{HbO_2} are constant within a statistically acceptable margin of error for an individual NIRS sensor used to monitor direct organ oxygenation on different human subjects. In other words, once the sensor is calibrated it can be used on various human subjects and produce accurate information for each human subject for each organ type. The same is true for animal subjects.

In the determination of the StO₂ percentage, the effective photon pathlength L_b cancels out. If, however, the photon pathlength is known or estimated, then the determination of the total value of Hb and/or HbO₂ is possible. For example, if a value for pathlength L_b is input into Equation 14 along with the calibration values Ψ_{Hb} and Ψ_{HbO_2} , then the total value of Hb and/or HbO₂ can be calculated. According to Equation 2, pathlength L can be estimated from the product of "B*d". The light source to detector separation (optode) distance parameter "d" in the pathlength calculation is a measurable value and can be made constant by setting a fixed distance between light source to detector in the NIRS sensor design. Alternatively, the parameter "d" can be measured once the optodes are placed on the subject by use of calipers, ruler, or other distance measurement means. The pathlength differential factor "B" is more difficult to measure and requires more sophisticated equipment. An estimation of the value of "B" can be determined within a statistically acceptable margin of error for empirical data collected from subjects; e.g., a large data set of measured organ differential pathlength factor values. Substitution of these predetermined values of "B" into Equation 14 results in the determination of the total values of Hb and HbO₂.

After the calibration parameters Ψ_{Hb} and Ψ_{HbO_2} are determined using the above-described methodology for an

individual NIRS sensor, this particular sensor is said to be calibrated. A calibrated NIRS sensor affords accurate measurement of total tissue oxygen saturation, StO_2 , by non-invasive means for each particular organ type. The calibrated sensor can be used thereafter on any human patient, including adults and neonates. The same is true for an animal subject if the sensor was calibrated on animals.

Besides Hb and HbO_2 , other biological constituents of interest (e.g., cytochrome aa_3 , etc.) could be determined using the multivariate forms of equations 2, 3, 6 or 7. For each additional constituent to be determined, an additional measuring wavelength will be needed.

In an alternative embodiment, the above-described methodology can be combined with pulse oximetry techniques to provide an alternative non-invasive method of distinguishing between oxygen saturation attributable to venous blood and that attributable to arterial blood. As demonstrated by Equation 4, $SmvO_2$ is determined by the ratio of venous oxygen saturation SvO_2 and arterial oxygen saturation SaO_2 from each particular organ type. A calibrated NIRS sensor affords accurate measurement of total tissue oxygen saturation, StO_2 , by using regression techniques by correlation to mixed venous oxygen saturation $SmvO_2$. Therefore, the following expression will result:

$$StO_2 = SmvO_2 = K_v * SvO_2 + Ka * SaO_2 \quad (\text{Eqn. 16})$$

Non-invasive pulse oximetry techniques can be used to determine the arterial oxygen saturation (SaO_2) of peripheral tissue (i.e. finger, ear, nose) by monitoring pulsatile optical attenuation changes of detected light induced by pulsatile arterial blood volume changes in the arteriolar vascular system. Arterial blood oxygen saturation determined by pulse oximetry is clinically denoted as SpO_2 . If NIRS monitoring and pulse oximetry monitoring are done simultaneously and SpO_2 is set equal to SaO_2 in Equation 16, then venous oxygen saturation of a particular organ being monitored can be determined from the following expression:

$$SvO_2 = \frac{StO_2 - (Ka * SpO_2)}{K_v} \quad (\text{Eqn. 17})$$

For the heart, venous oxygen saturation SvO_2 would be determined from the coronary drainage SvO_2 and the parameters Ka and K_v would be empirically determined during the calibration of the NIRS sensor. Under most physiological conditions, SpO_2 is representative of organ arterial oxygen saturation SaO_2 . Therefore, depending on which venous saturation parameter was used to calibrate the NIRS sensor, this clinically important parameter (organ venous oxygen saturation) can be determined by Equation 23 by non-invasive means.

To perform a corrective procedure (e.g., coronary artery bypass grafting, or heart valve replacement, or correction of congenital heart defect, etc.) on a subject's heart, it is often necessary to arrest or stop the subject's heart so that surgical procedures can be done in a still and bloodless field. In some instances this is achieved by diverting deoxygenated blood that would otherwise enter the heart into a heart-lung machine. The heart-lung machine takes over the function of the lung by oxygenating the blood and removing carbon dioxide. After oxygenation, filtration and removal of carbon dioxide, the machine pumps the blood back into the body usually through the aorta.

At the same time, the subject's heart is isolated, for example, using an aortic cross-clamp on the distal aorta. Cold cardioplegia is subsequently given into the heart

through the aortic root. The cold fluid (usually in the range of about 4-10° C.) ensures that the heart cools down to an approximate temperature of around 15-20° C. thus slowing down the metabolism of the heart and thereby preventing damage to the heart muscle. The process may be further augmented by a cardioplegic component which is high in potassium and magnesium. The potassium helps by arresting the heart in diastole thus ensuring that the heart does not use up the valuable energy stores during this period of heart isolation. Blood can be added to this solution especially for long procedures requiring more than half an hour of cross-clamp time. Blood acts a buffer and also supplies nutrients to the heart during ischemia.

During the procedure (e.g., before, during, and after isolation), a sterile NIRS sensor is placed on a region of the heart. The sensor may have one or more detectors. The sensor is operated to transmit light signals into the heart tissue and receive light signals that have passed through the heart tissue. The oxygen saturation of the heart tissue is determined using the oximeter described above.

Using the present invention oximeter in the way described (or similar way) enables the physician to monitor the oxygen saturation level of the heart tissue and take corrective action (e.g., add a cardioplegic component) if the saturation reaches a predetermined level. The ability to place a sensor directly in contact with the heart during the procedure enhances the accuracy of the data, as there is no interfering tissue overlaying the heart, thereby providing the care giver with quick, concise information relation to the oxygenation state of the organ. In this manner, the operator can monitor in real-time the cardioplegia of the heart, including the rate of recovery from the cardioplegia.

Once the procedure on the heart (e.g., coronary artery bypass grafting, or heart valve replacement, or correction of congenital heart defect, etc.) is over, the cross-clamp is removed and the isolation of the heart is terminated so that normal blood supply to the heart is restored and the heart starts beating again. At this point, the oximeter can be used to confirm the oxygen saturation level within the heart tissue is within acceptable limits.

The utility of the present invention can be seen through the case study depicted in FIG. 6, which shows a recording of StO_2 from a present NIRS sensor placed directly on the heart of a pig. The events occurring during the study are identified to illustrate the relationship between a change in the level of StO_2 and the event encountered during the cardiac surgery. Event 1 ("E1") shows normal sinus rhythm. The StO_2 value for Event 1 is fairly low due to the high oxygen extraction of the heart muscle. Blood drawn from the venous return of the heart (carotid sinus) is typically highly oxygen de-saturated, with an oxygen saturation value around 40%. During Event 2 ("E2"), the pig was put on a cardiac bypass pump. During Event 3 ("E3"), a cross clamp was placed on the ascending aorta, proximal to the cardiopulmonary bypass ("CPB") inflow cannula. The cross clamp stopped blood flow to the heart; e.g., to allow a physician to perform surgery on the heart devoid of blood in the field. As a result, the StO_2 value decreased because the heart was no longer receiving oxygenated blood. Removal of the cross-clamp resulted in a resumption of blood perfusion to the heart, and the StO_2 value consequently increased back to pre-clamping StO_2 levels. Event 4 ("E4") was a repeat of Event 3. During Event 5 ("E5"), the cross-clamp was applied again, with a consequent decrease in the StO_2 level. Subsequently, cardioplegia was injected into the ascending aorta. The cardioplegia consisted of cooled (10 degrees C.) arterial blood and a saline-like solution, that included a

relatively high level of potassium and other components such as adenin and Sodium (Na) channel blockers solution. The only vessels that extend out from the ascending aorta are the coronaries, provided that the aortic valve is competent. The infusion of cardioplegia into the ascending aorta passed through the coronary arteries. Since arterial blood is nearly 100% oxygenated, the cardiac NIRS sensor sensed a sharp rise in StO₂ level as the highly oxygenated blood perfused through the coronary vessels of the heart. The StO₂ level remained elevated for a few minutes as the heart muscle metabolizes the oxygen from the injected blood solution. Eventually all (or nearly all) the oxygen was extracted from the injected blood solution, causing the StO₂ level to decrease steadily until a value near zero was approached, which is depicted at Event 6 (“E6”). The StO₂ level remained near zero until a second dose of cardioplegia was administered—depicted at Event 7 (“E7”). The StO₂ level subsequently rose again to elevated values as the heart received freshly oxygenated blood solution from the second dose of cardioplegia. As the second dose of cardioplegia wore off, the StO₂ level decreased. This form of cardioplegia is referred to as antegrade cardioplegia. Another type of cardioplegia is called retrograde cardioplegia, which is injected into the coronary sinus of the heart to induce the same effect.

At Event 8 (“E8”), the cross-clamp was removed and the heart reperfed from systemic flow from the ascending aorta. At Event 9 (“E9”), the heart underwent ventricular fibrillation and arrhythmia. Because the heart’s contraction was uncoordinated and the cardiac muscles were not fully contracting, the oxygen consumption was reduced, resulting in a higher than normal cardiac StO₂ level. At Event 10 (“E10”), the heart was defibrillated to induce normal sinus rhythm, at which point the StO₂ level returned to a normal sinus rhythm value. Event 11 (“E11”) illustrates a potential problem from blood pooling around the heart. The pooled blood negatively affected the ability of the NIRS sensor to accurately sense the StO₂ level within the heart. A NIRS cardiac sensor that is well adhered to the heart muscle can avoid this problem because it prevents outside blood leaking into the sensor’s optical interface with the heart muscle. In one embodiment, the sensor can include a mechanism for inducing suction between the sensor and the surface of the heart; e.g., a syringe positioned to create suction can be used to create and maintain a seal that prevents outside blood leakage. At Event 12 (“E12”), the animal expired and the StO₂ level decreased to near zero values.

It can be seen from the above that the present apparatus for, and method of, directly monitoring the level of StO₂ in an organ using an NIRS sensor, provides an accurate and effective tool for directly monitoring both the StO₂ level of the organ and the affects of cardioplegia on the StO₂ level. Furthermore, the present invention provides the ability to regulate the StO₂ level within the organ using information collected during such monitoring. For example, the StO₂ level within a subject’s heart can be directly monitored using the present invention during organ surgery (e.g., heart surgery). In one embodiment, the level of StO₂ is monitored and if the StO₂ level decreases beyond a predetermined minimum value threshold while the heart is cross-clamped, a predetermined dosage of cardioplegia is injected into the organ to cause the StO₂ level to increase above the threshold. It should be noted that the StO₂ threshold may be chosen based on the circumstances of the application; e.g., the threshold may be a value recorded during normal sinus rhythm and cardiac function, or some other clinically relevant value.

In another embodiment, the StO₂ level is again monitored and the value of the StO₂ level is compared against a predetermined StO₂ level threshold as a function of time. For example, if the StO₂ level within the organ drops below the StO₂ threshold (referred to as an “event”), the amount of time that the StO₂ value is below the threshold (i.e., the duration of the event) is recorded and summed to create a time under threshold (i.e., “TUT”) value (typically expressed in units of seconds or minutes). FIG. 7 graphically illustrates a TUT region. Anytime the StO₂ level exceeds the threshold, the TUT value remains constant and unchanged. The TUT value can then be compared to a threshold value for supplementary and/or corrective action if required.

The monitoring and collective summing relative to StO₂ values below the threshold can also be implemented (e.g., within the processor 12) graphically using a graph of StO₂ (e.g., Y-axis) versus time (e.g., X-axis). For example, if the comparison of the sensed StO₂ level with the predetermined StO₂ level reveals that the StO₂ level within the organ is below the threshold value, the area of the graph defined by the data line and the threshold value (i.e., from start of the event to the finish of the event) is calculated and summed to create an area under threshold (i.e., “AUT”) value. FIG. 7 graphically illustrates an AUT region. Anytime the StO₂ level exceeds the threshold, the AUT value remains constant and unchanged. The AUT value can then be compared to a threshold value for supplementary action if required.

The area under StO₂ threshold (“AUT”) value can also be calculated mathematically, where the AUT value (in units of StO₂%-time) is the accumulated oxygen saturation deficit below a threshold multiplied by time. AUT can be calculated, for example, using the following equation:

$$AUT = AUT_{past} + (StO_{2threshold} - StO_{2value}) \times \text{sample rate} \quad [\text{Eqn. 18}]$$

where AUT is the collective value, AUT_{past} is the previous collective value, $StO_{2threshold}$ is the threshold value, and StO_{2value} is the currently sensed StO_{2value} . If the currently sensed StO₂ value is above the threshold, then the collective AUT value remains constant. The AUT is calculated for the entire procedure or just for one portion of the procedure (e.g., during a period of circulatory arrest). Because the StO₂ level of an organ will likely fluctuate during the course of an operation (i.e., StO₂ values above and below the predetermined threshold value), the AUT value can be periodically determined. To illustrate, consider the following monitoring using an arbitrary sample rate of sensing every five minutes and a StO₂ threshold value of 60): At a first point in time (T1), the AUT value may be considered to be zero (e.g., current sensing not yet started, and no past value of AUT). Five minutes later at T2, the StO₂ of the organ is sensed and has a value of 58. Using Eqn. 1 above, the AUT value is calculated: $[(0 + (60 - 58) \times 5) = 10]$, because there is no AUT_{past} value, and it is therefore assigned a value of zero. Five minutes later at T2, the StO₂ of the organ is sensed and has a value of 57. Again using Eqn. 1: $[(60 - 57) \times 5 + 10 = 25]$, because the AUT_{past} value is equal to ten. Five minutes later at T3, the StO₂ of the organ is sensed and has a value of 63. In this instance, because the sensed StO₂ value is greater than the threshold, the difference is assigned a value of zero, and the AUT value remains at twenty-five, because the AUT_{past} value is equal to twenty-five.

It should be noted that the AUT value is not a percentage of time spent under a given threshold, but rather is an absolute value that can be used during or after the procedure as part of the subject’s care; e.g., depending upon the application, there may be little difference in clinical terms between a collective StO₂ duration of one hour underneath

a threshold during a 10 hour case (10%) as opposed to a collective one hour during a one hour case (100%).

It can be seen from the description above, including the various different embodiments, that the present invention provides a method and apparatus for monitoring cardioplegia. The present invention can be used to protect the subject's heart (or other organ), by alerting the physician to take a proactive step (e.g., inject the next round of cardioplegia) when certain predefined conditions are met. The present method can be utilized in varying degrees of automation (e.g., algorithmically within the processor 12), including an automated cardioplegia device that includes structure adapted to automatically inject cardioplegia based on NIRS StO₂ monitoring input.

As indicated above, the present apparatus and method can be used to directly monitor numerous organs within a subject's body and is not, therefore, limited to a cardiac application. For example, the present invention can be used to monitor oxygenation levels within a liver or kidney allograft before, during and after transplantation. Issues may arise during transplantation such as reconnect of blood vessels that feed the organ, as well as preventing hyperoxygenation and reperfusion injury. An NIRS organ sensor could be moved to different positions on the organ to make sure each region or compartment is perfused adequately. This is especially true with liver transplantation. To prevent hyperoxygenation and in part, reperfusion injury, the organ StO₂ can be controlled by observing the NIRS monitor StO₂ value and then regulating blood flow or the oxygenation level so that StO₂ is maintained within a predetermined range during transplant. Specifically, a sterile NIRS sensor is placed on a region of the liver or kidney allograft. Secure attachment of the sensor can be made by suction means or by suturing the sensor onto the organ. Alternatively a loose wrapping over the sensor to hold it on the organ, such as a stretchable net-like apparatus can be used. Although the sensor may have one or more detectors, the direct sensing of the organ typically permits the use of a single detector sensor since attenuation attributable to non-organ tissue layers is avoided. The sensor is operated to transmit light signals into the liver tissue and receive light signals that have passed therethrough. The oxygen saturation of the liver tissue is determined using the oximeter described above. As indicated above, the direct organ monitoring made possible using the present invention enables the physician to accurately monitor the organ and take corrective action as needed.

Since many changes and variations of the disclosed embodiment of the invention may be made without departing from the inventive concept, it is not intended to limit the invention otherwise than as required by the appended claims.

What is claimed is:

1. A method for monitoring the effects of cardioplegia administered to a heart, comprising:
 applying a near infrared spectrophotometric (NIRS) sensor onto an external surface of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within heart tissue using the data signals;
 sensing a pre-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor, the pre-cardioplegia sensing including emitting an incident first light from a light source portion of the NIRS sensor directly into the tissue of the heart at a first tissue location, and detecting the emitted first light at a second

tissue location, which second tissue location is separated from the first tissue location, and which detected emitted first light has traveled through the tissue between the first tissue location and the second tissue location, and producing a first set of the data signals representative of an intensity of the detected emitted first light;
 producing a pre-cardioplegia sensed heart tissue oxygenation level value using the processor based on the first set of the data signals;
 administering a cardioplegia to the heart;
 sensing a post-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor after the administration of the cardioplegia to the heart, the post-cardioplegia sensing including emitting an incident second light from the light source portion of the NIRS sensor directly into the tissue of the heart at the first tissue location, and detecting the emitted second light at the second tissue location, and which detected emitted second light has traveled through the tissue between the first tissue location and the second tissue location, and producing a second set of the data signals representative of an intensity of the detected emitted second light;
 producing a post-cardioplegia administration sensed heart tissue oxygenation level value using the processor based on the second set of the data signals;
 comparing one or both of the pre-cardioplegia sensed heart tissue oxygenation level value and the post-cardioplegia sensed heart tissue oxygenation level value to a heart oxygenation level threshold;
 alerting an operator if the post-cardioplegia sensed heart tissue oxygenation level reaches a predetermined threshold after administration of cardioplegia; and
 administering a secondary cardioplegia if the post-administration post-cardioplegia sensed heart tissue oxygenation level drops below a predetermined threshold.

2. A method for monitoring the effects of cardioplegia administered to a heart, comprising:
 applying a near infrared spectrophotometric (NIRS) sensor onto an external surface of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within heart tissue using the data signals;
 sensing a pre-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor, the pre-cardioplegia sensing including emitting an incident first light from a light source portion of the NIRS sensor directly into the tissue of the heart at a first tissue location, and detecting the emitted first light at a second tissue location, which second tissue location is separated from the first tissue location, and which detected emitted first light has traveled through the tissue between the first tissue location and the second tissue location, and producing a first set of the data signals representative of an intensity of the detected emitted first light;
 producing a pre-cardioplegia sensed heart tissue oxygenation level value using the processor based on the first set of the data signals;
 administering a cardioplegia to the heart;
 sensing a post-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor after the administration of the cardioplegia to the heart, the post-cardioplegia sensing including emitting an incident second light from the light source portion of the NIRS

sensor directly into the tissue of the heart at the first tissue location, and detecting the emitted second light at the second tissue location, and which detected emitted second light has traveled through the tissue between the first tissue location and the second tissue location, and producing a second set of the data signals representative of an intensity of the detected emitted second light; and

producing a post-cardioplegia administration sensed heart tissue oxygenation level value using the processor based on the second set of the data signals;

comparing one or both of the pre-cardioplegia sensed heart tissue oxygenation level value and the post-cardioplegia sensed heart tissue oxygenation level value to a heart oxygenation level threshold;

alerting an operator if the post-cardioplegia sensed heart tissue oxygenation level reaches a predetermined time under threshold (TUT) after administration of cardioplegia; and

administering a secondary cardioplegia if the post-cardioplegia sensed heart tissue oxygenation level reaches the predetermined time under threshold (TUT).

3. A method for monitoring the effects of cardioplegia administered to a heart, comprising:

applying a near infrared spectrophotometric (NIRS) sensor onto an external surface of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within heart tissue using the data signals;

sensing a pre-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor, the pre-cardioplegia sensing including emitting an incident first light from a light source portion of the NIRS sensor directly into the tissue of the heart at a first tissue location, and detecting the emitted first light at a second tissue location, which second tissue location is separated from the first tissue location, and which detected emitted first light has traveled through the tissue between the first tissue location and the second tissue location, and producing a first set of the data signals representative of an intensity of the detected emitted first light;

producing a pre-cardioplegia sensed heart tissue oxygenation level value using the processor based on the first set of the data signals;

administering a cardioplegia to the heart;

sensing a post-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor after the administration of the cardioplegia to the heart, the post-cardioplegia sensing including emitting an incident second light from the light source portion of the NIRS sensor directly into the tissue of the heart at the first tissue location, and detecting the emitted second light at the second tissue location, and which detected emitted second light has traveled through the tissue between the first tissue location and the second tissue location, and producing a second set of the data signals representative of an intensity of the detected emitted second light; and

producing a post-cardioplegia administration sensed heart tissue oxygenation level value using the processor based on the second set of the data signals;

comparing one or both of the pre-cardioplegia sensed heart tissue oxygenation level value and the post-cardioplegia sensed heart tissue oxygenation level value to a heart oxygenation level threshold;

alerting an operator if the post-cardioplegia sensed heart tissue oxygenation level reaches a predetermined area under threshold (AUT) after administration of cardioplegia; and

administering a secondary cardioplegia if the post-cardioplegia sensed heart tissue oxygenation level reaches the predetermined area under threshold (AUT).

4. A method for monitoring the effects of cardioplegia administered to a heart, comprising:

applying a near infrared spectrophotometric (NIRS) sensor onto an external surface of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within heart tissue using the data signals;

sensing a pre-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor, the pre-cardioplegia sensing including emitting an incident first light from a light source portion of the NIRS sensor directly into the tissue of the heart at a first tissue location, and detecting the emitted first light at a second tissue location, which second tissue location is separated from the first tissue location, and which detected emitted first light has traveled through the tissue between the first tissue location and the second tissue location, and producing a first set of the data signals representative of an intensity of the detected emitted first light;

producing a pre-cardioplegia sensed heart tissue oxygenation level value using the processor based on the first set of the data signals;

administering a cardioplegia to the heart;

sensing a post-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor after the administration of the cardioplegia to the heart, the post-cardioplegia sensing including emitting an incident second light from the light source portion of the NIRS sensor directly into the tissue of the heart at the first tissue location, and detecting the emitted second light at the second tissue location, and which detected emitted second light has traveled through the tissue between the first tissue location and the second tissue location, and producing a second set of the data signals representative of an intensity of the detected emitted second light;

producing a post-cardioplegia administration sensed heart tissue oxygenation level value using the processor based on the second set of the data signals;

comparing the post-cardioplegia sensed heart tissue oxygenation level value to a heart oxygenation level threshold; and

administering a secondary cardioplegia if the post-administration post-cardioplegia sensed heart tissue oxygenation level drops below the threshold.

5. A method for monitoring the effects of cardioplegia administered to a heart, comprising:

applying a near infrared spectrophotometric (NIRS) sensor onto an external surface of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within heart tissue using the data signals;

sensing a pre-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor, the pre-cardioplegia sensing including emitting an incident first light from a light source portion of the NIRS sensor directly into the tissue of the heart at a first tissue

location, and detecting the emitted first light at a second tissue location, which second tissue location is separated from the first tissue location, and which detected emitted first light has traveled through the tissue between the first tissue location and the second tissue location, and producing a first set of the data signals representative of an intensity of the detected emitted first light; 5

producing a pre-cardioplegia sensed heart tissue oxygenation level value using the processor based on the first set of the data signals; 10

administering a cardioplegia to the heart;

sensing a post-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor after the administration of the cardioplegia to the heart, the post-cardioplegia sensing including emitting an incident second light from the light source portion of the NIRS sensor directly into the tissue of the heart at the first tissue location, and detecting the emitted second light at the second tissue location, and which detected emitted second light has traveled through the tissue between the first tissue location and the second tissue location, and producing a second set of the data signals representative of an intensity of the detected emitted second light; and 20

producing a post-cardioplegia administration sensed heart tissue oxygenation level value using the processor based on the second set of the data signals; 25

comparing the post-cardioplegia sensed heart tissue oxygenation level value to a heart oxygenation level threshold; 30

determining a time under threshold (TUT) value based on a collective amount of time the post-cardioplegia sensed heart tissue oxygenation level is below the threshold; and 35

administering a secondary cardioplegia if the TUT value reaches a predetermined TUT threshold.

6. A method for monitoring the effects of cardioplegia administered to a heart, comprising: 40

applying a near infrared spectrophotometric (NIRS) sensor onto an external surface of the heart, which sensor is in communication with a processor adapted to receive data signals from the sensor, and is adapted to determine the oxygen saturation level within heart tissue using the data signals;

sensing a pre-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor, the pre-cardioplegia sensing including emitting an incident first light from a light source portion of the NIRS sensor directly into the tissue of the heart at a first tissue location, and detecting the emitted first light at a second tissue location, which second tissue location is separated from the first tissue location, and which detected emitted first light has traveled through the tissue between the first tissue location and the second tissue location, and producing a first set of the data signals representative of an intensity of the detected emitted first light;

producing a pre-cardioplegia sensed heart tissue oxygenation level value using the processor based on the first set of the data signals;

administering a cardioplegia to the heart;

sensing a post-cardioplegia oxygen saturation level within the heart tissue using the NIRS sensor after the administration of the cardioplegia to the heart, the post-cardioplegia sensing including emitting an incident second light from the light source portion of the NIRS sensor directly into the tissue of the heart at the first tissue location, and detecting the emitted second light at the second tissue location, and which detected emitted second light has traveled through the tissue between the first tissue location and the second tissue location, and producing a second set of the data signals representative of an intensity of the detected emitted second light; and

producing a post-cardioplegia administration sensed heart tissue oxygenation level value using the processor based on the second set of the data signals;

comparing the post-cardioplegia sensed heart tissue oxygenation level value to a heart oxygenation level threshold;

determining an area under threshold (AUT) value based on a collective amount of time the post-cardioplegia sensed heart tissue oxygenation level is below the threshold; and

administering a secondary cardioplegia if the post-cardioplegia sensed heart tissue oxygenation level reaches a predetermined AUT threshold.

* * * * *

专利名称(译)	人体器官的分光光度血液氧合监测方法		
公开(公告)号	US10485462	公开(公告)日	2019-11-26
申请号	US15/172992	申请日	2016-06-03
申请(专利权)人(译)	CAS医疗系统, INC.		
当前申请(专利权)人(译)	爱德华生命科学公司		
[标]发明人	BENNI PAUL B		
发明人	BENNI, PAUL B.		
IPC分类号	A61B5/1455 A61B5/00		
CPC分类号	A61B5/7246 A61B5/7278 A61B5/4836 A61B5/746 A61B5/7271 A61B5/14551 A61B5/4848 A61B2562/0233		
代理人(译)	GETZ, RICHARD D.		
优先权	PCT/US2010/058059 2010-11-24 WO 61/264080 2009-11-24 US		
其他公开文献	US20160278675A1		
外部链接	Espacenet		

摘要(译)

提供了一种用于直接使用近红外分光光度传感器来非侵入性地确定对象器官内的血氧饱和度水平的方法和设备。该方法包括以下步骤：a) 使用传感器将光信号直接传输到受试者的器官中；以及b) 在光信号行进经过预定距离的对象的器官之后，感测光信号的第一强度和光信号的第二强度；c) 使用感测到的第一强度和感测到的第二强度来确定光信号沿多个不同波长的衰减；d) 确定光信号在波长之间的衰减差异；e) 利用波长之间的衰减差确定受试者器官内的血氧饱和度。

