A method for converting a partial pressure of oxygen (pO₂) value in blood at a measurement temperature to a corresponding value at a reference temperature (37 °C). A pO₂ value is determined by measurements made in a patient's blood stream using a phosphorescent compound sensitive to oxygen concentration. The compound is illuminated with a short pulse of light, causing a phosphorescent emission having a rate of decay that varies as the function of the pO₂ in the blood. A detector (34) produces an electrical signal corresponding to the intensity of the emission, and the signal is converted to a corresponding digital value for input to a microcomputer (12). Supplied to the microcomputer in digital form is a signal indicative of the temperature at the measurement site where the compound is disposed. The microcomputer determines phosphorescent decay rate and from that determines pO₂ at the measurement site for measurement temperature.
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HEMOMETRIX TEMPERATURE COMPENSATION

Field of the Invention

The present invention generally relates to a method for determining a partial pressure of oxygen in blood, and more specifically, to a method for determining a partial pressure of oxygen at a reference temperature, as a function of a partial pressure of oxygen measured at a different temperature.

Background of the Invention

In conventional blood gas analysis machines, a sample of blood withdrawn from a patient is heated to a reference temperature of 37°C before the partial pressure of oxygen (pO₂) is determined. Over time, the medical community has thus developed a preference for reporting all pO₂ measurements referenced to 37°C. However, when pO₂ is measured in vivo instead of in a blood gas analyzer, the measurement is often made at a substantially different temperature even though 37°C is normal body temperature. For example, during certain surgical procedures, it is necessary to lower the patient's body temperature by as much as 20°C, thereby depressing metabolic activity. If the pO₂ measurement is made while the patient's body is chilled, the result is very different than a corresponding measurement made at 37°C. An anesthesiologist controlling the administration of oxygen and other gases to the patient typically prefers to record the pO₂ at the accepted reference temperature of 37°C rather than the lower measurement. Moreover, the anesthesiologist may prefer to base decisions concerning the patient's condition on pO₂ data referenced to 37°C. When making such decisions during a critical operation, the anesthesiologist may not have time to draw a sample of blood for analysis in a blood gas analyzer at the reference temperature. An in vivo, real time determination of pO₂ is sometimes essential, even if carried out at a different measurement temperature than the desired 37°C reference.
Conversion between a pO$_2$ measurement at one temperature to that at another temperature is not a trivial task. The solubility of oxygen in blood as a function of temperature is determined by a non-algebraic combination of transcendental functions. As a result, it is not possible to analytically solve a simple equation to convert a pO$_2$ measurement at a substantially different temperature to a corresponding value at the desired reference temperature of 37°C. In the past, medical personnel have been forced to manually convert a measured value for pO$_2$ to the 37°C reference temperature using a nomogram or by interpolating values from a look-up table. Neither of these techniques are particularly desirable when speed in determining the data is essential; furthermore, any human errors in the conversion process can have potentially life-threatening consequences.

The complex relationship between temperature and pO$_2$ is evident from the equation that has been developed in the prior art to convert from a pO$_2$ measurement made at 37°C to a different temperature. This equation is reported by R. A. Ashwood, G. Kost, and M. Kenny in "Temperature Correction of Blood-Gas and pH Measurements," Clinical Chemistry, Vol. 29, 11:1877-1885, 1983 and by J. W. Severinghaus in "Simple, Accurate Equations for Human Blood O$_2$ Dissociation Computations," American Journal of Physiology, Vol. 46, 3:599-602.

The equation in both of these references is as follows:

$$pO_{2PT} = pO_{2REF}^{10}$$

where $pO_{2PT}$ is the predicted partial pressure of oxygen at a temperature $T$ that is different than the reference temperature, and $pO_{2REF}$ is the measured partial pressure of oxygen at the reference temperature (37°C). One might assume that equation 1 could simply be rearranged to determine $pO_2$ at the reference temperature from the $pO_2$ measured at a temperature different than the reference temperature, i.e., as follows:

$$pO_{2REF} = pO_{2M}^{10}$$

where $pO_{2M}$ is the partial pressure of oxygen measured at the temperature $T$.

However, Equation 2 produces erroneous results due to the nature of the mathematical relationship between measurement temperature and $pO_2$. In the above-noted paper, Severinghaus recognized the difficulty of calculating $pO_2$ at
at 37°C based on measurements made at other temperatures. On page 600 of the Journal, he suggests that, "[t]o begin with some other temperature, one may estimate a trial 37°C pO₂ using the factors 6%/°C if pO₂ < 100, and 6 Torr/°C above 100 Torr, and proceed iteratively in Eq. (3)." (Equation 3 in the reference is equivalent to Equation 1, above.) What Severinghaus intended by this statement is not entirely clear, because he did not present any example of how the iterative process is carried out nor any support for its efficacy in producing an accurate result.

Accordingly, a method is required for quickly determining the pO₂ of a patient's blood at the reference temperature, based on a measurement made at another temperature. The method should be automatically carried out to avoid human error and must be implemented quickly (in real time) to make the results of an in vivo measurement of pO₂ at the patient's temperature immediately available as a corresponding value at the reference temperature during critical medical procedures.

Summary of the Invention

In accordance with the present invention, a method for determining a partial pressure of oxygen at a predefined reference temperature during an in vivo blood gas measurement begins with the step of measuring pO₂ at an arbitrary temperature (that is substantially different than the reference temperature), as a function of a physical parameter that changes to indicate the pO₂ at a measurement site. The measurement includes the steps of producing a first signal indicative of the pO₂ in response to changes in the physical parameter; producing a second signal indicative of the arbitrary temperature at the measurement site; and processing the first signal to determine the pO₂ at the arbitrary temperature. Next, an estimate of the pO₂ at the predefined reference temperature is determined as a function of the first signal and of the second signal. After the initial estimate is determined, a more accurate estimate of pO₂ at the predefined reference temperature is made as a function of the first signal, the second signal, and the estimate of the pO₂ at the predefined reference temperature. Subsequently, the step of determining the more accurate estimate is iteratively repeated, each iteration using the more accurate estimate of the pO₂ from a previous iteration as a value for the estimate. After a predefined number of such iterations have been completed, the resulting more accurate estimate approximates the pO₂ at the predefined reference temperature corresponding to the measurement of the pO₂ made at the arbitrary temperature.
Each iteration determines the more accurate estimate, as defined by the equation:

\[ X_{N+1} = X_0 10^{\left( \frac{(T-R)}{k_1} \right) f(X_N)} \]

where \( X_0 \) is the \( pO_2 \) measured at the arbitrary temperature; \( T \) is the arbitrary temperature; \( T_R \) is the predefined reference temperature; \( X_{N+1} \) is the more accurate estimate of the \( pO_2 \) for an \((N + 1)\)th iteration; \( X_N \) is the more accurate estimate of the \( pO_2 \) for an \( N \)th iteration; \( K_1 \) is a predetermined constant; and \( f(X_N) \) is a predefined function of \( X_N \). The function \( f(X_N) \) is represented by:

\[ f(X_N) = \frac{K_2}{K_3 \left( \frac{X_N}{K_4} \right)^{K_5}} + K_6 \]

where \( K_2 \) through \( K_6 \) are predefined constants.

If the arbitrary temperature is less than the reference temperature, for each iteration after the first computed iteration, an average of the previous two estimates of the \( pO_2 \) at the predefined reference temperature is determined and is used as the value for the estimate in the current iteration to determine the next more accurate estimate. The predefined number of iterations is selected so that the more accurate estimate of the \( pO_2 \) at the predefined reference temperature converges to a value within a predefined convergence limit. Preferably, the predefined number of iterations is less than ten.

The foregoing aspects and many of the attendant advantages of this invention will become more readily appreciated by reference to the following detailed description, when taken in conjunction with the accompanying drawings.

**Brief Description of the Drawings**

FIGURE 1 is a block diagram of a representative system for measuring \( pO_2 \) at an \textit{in vivo} measurement site;

FIGURE 2 is a flow chart showing the logical steps required in carrying out the present invention on a central processing unit (CPU);

FIGURE 3 is a graph showing the per cent error for the determination of \( pO_2 \) at a number of different measurement temperatures, when the present method is applied to determine a corresponding \( pO_2 \) at a reference temperature (37°C); and
FIGURE 4 is a three-dimensional graph showing the relationship between the number of iterations, probe (measurement) temperature, and pO₂ level with respect to the present invention.

**Detailed Description of the Preferred Embodiment**

A system for monitoring the concentration of oxygen (O₂) in blood is shown graphically in a block diagram generally reference 10 in FIGURE 1. As is conventional in the medical art, the concentration of O₂ in blood is measured in terms of the partial pressure of O₂ (pO₂). System 10 converts between a measurement of pO₂ at a temperature different than a desired reference temperature, which is normally 37°C, the temperature corresponding to normal human body temperature. System 10 determines pO₂ at the reference temperature in real time, for example, during an operation so that an anesthesiologist can adjust the relative proportion of oxygen to other gases being administered to the patient, and converts any measurement made at a temperature different than the reference temperature to a corresponding value at the reference temperature. This information is provided to medical personnel so they can make appropriate decisions concerning the patient. As noted above, it is important during such procedures that the conversion from pO₂ at a measurement temperature to the corresponding pO₂ at the reference temperature be carried out in real time, since medical decisions concerning patient's safety should not be delayed for determination of pO₂ in a blood gas analyzer. In certain medical procedures, the patient may be chilled to a temperature as much as 20°C below normal body temperature, or due to physiological changes affecting the patient's body temperature at which the temperature which the pO₂ measurement is made may be substantially higher than the desired reference temperature of 37°C. Accordingly, system 10 automatically displays pO₂ at the reference temperature, regardless of the actual temperature at which the pO₂ is measured in the patient's blood.

System 10 includes a microcomputer 12, which preferably comprises a laptop or other portable computer having a display 12a, a keyboard 12b, and a floppy disk or hard drive 12c for storing data and programs. Microcomputer 12 controls a light source 14 via a signal supplied to the source over a lead 16, the signal causing light source 14 to emit a short pulse of light at a time t₀. The light pulse produced by light source 14 passes through a filter 18 that allows only shorter wave length light, e.g., light having a wave length in the range of 480 to 600 nanometers (nm), to pass through into an optical fiber 20. Optical fiber 20 conveys the filtered light pulse to an optical coupler 22. Optical coupler 22
couples the filtered light pulse into an optical fiber 24, which conveys the filtered light pulse into the patient's body via an intravascular catheter 25 to a pO₂ sensor 26 that is disposed within the patient's blood stream 28. Sensor 26 comprises a temperature sensor 27, (for example, a thermistor, thermocouple, or other temperature sensitive element) and a phosphorescent compound 29. Preferably, the phosphorescent compound comprises a fluorinated derivative of platinum tetra(pentafluorophenyl)porphyrin, Pt(TFPP), which is excited into phosphorescence by light having a wave length in the range 480 to 600 nm, and the phosphorescent emission thereafter decays with a rate that is a function of the concentration of oxygen to which the phosphorescent compound is exposed. The oxygen in the blood stream in which sensor 26 is disposed quenches the phosphorescent emission of this compound, and the rate at which the phosphorescent emission decays is proportional to the pO₂ of the blood. The characteristic decay time of the phosphorescent emission produced by phosphorescent compound 29 is thus a measurable parameter indicative of pO₂ in blood stream 28. Details of this method for measuring oxygen concentration (i.e., pO₂) are fully disclosed in commonly assigned U.S. Patent No. 4,810,655, the specification and disclosure of which are specifically incorporated herein by reference.

Optical fiber 24 conveys the phosphorescent emission from phosphorescent compound 29 as a return light signal in optical fiber 24 back toward optical coupler 22, wherein it is coupled into an optical fiber 30. Optical fiber 30 conveys the phosphorescent emission from sensor 26 through a cut-off filter 32 that allows only light having a wave length longer than 620 nm to enter an adjacent detector 34. Optical filter 18 and cut-off filter 32 are thus selected to transmit complimentary wave lengths, thereby preventing light from light source 18 from directly reaching detector 34, but enabling the phosphorescent emission to be detected by it.

Detector 34 preferably comprises a photodiode that produces an electrical signal corresponding to the intensity of the phosphorescent emission returning from sensor 26, which is conveyed over leads 36 to a preamplifier 38. Preamplifier 38 amplifies the electrical signal and supplies the amplified signal through leads 40 to an analog-to-digital (A-D) converter 42. A-D converter 42 converts the amplified signal from preamplifier 38 into a corresponding digital signal that is conveyed through leads 46 into an input port (not separately shown) of microcomputer 12. The A-D converter is controlled by a signal output from microcomputer 12 over
lines 44 to select either the amplified signal or the analog temperature signal, which is produced by temperature sensor 27, for A-D conversion. This analog temperature signal is indicative of the temperature of blood stream 28, i.e., the temperature at the measurement site, and is conveyed over a lead 48 to A-D converter 42 from temperature sensor 27.

The intensity of the phosphorescent emission is determined by microcomputer 12 at two times, \( t_1 \) and \( t_2 \), after the light pulse from light source 14 has terminated. Using these two intensities, the microcomputer determines the simple exponential decay rate of the emission produced by phosphorescent compound 29. Based upon the decay rate, a value for \( pO_2 \) at the temperature at the measurement site is determined at the temperature sensed by temperature sensor 27. If the temperature at the measurement site is different than the reference temperature of 37°C, microcomputer 12 executes a program to convert the value for \( pO_2 \) at the measurement temperature into a corresponding value at the reference temperature as described below.

The logical steps implemented by microcomputer 12 in determining a value for \( pO_2 \) at the reference temperature based on a measured value at a different temperature are shown in a flow chart 50 (FIGURE 2). After a start block 52, a block 54 provides for the analog to digital conversion of the temperature, \( T \), at the measurement site. In a block 56, microcomputer 12 produces a signal causing light source 14 to produce the light pulse, to irradiate phosphorescent compound 29 with light for a short period of time. A block 58 provides for the A-D conversion of the phosphorescence emission produced by phosphorescent compound 29 at times \( t_1 \) and \( t_2 \). In a block 60, the decay rate of the phosphorescent emission is determined based upon the exponential relationship of the intensity of light at times \( t_1 \) and \( t_2 \). Using the decay rate from block 60, a partial pressure of oxygen, \( X_0 \), at the measurement temperature \( T \) is determined in a block 62. A decision block 63 then determines if \( T = 37^\circ C \), and if so, branches to a block 78 to display the \( pO_2 \) value to a user on display 12a (shown in FIGURE 1). If \( T \) is not equal to 37°C, the flow chart continues with a block 64.

Block 64 initializes a counter \( N \) to 1. In a block 66, an expression is evaluated to determine a first estimate, \( X_N \), for the \( pO_2 \) at the reference temperature, 37°C, as a function of the measured value for \( pO_2 \), \( X_0 \), and as a function of the temperature at the measurement site, \( T \).
\[ X_N = X_0 10 \] 

If the measurement temperature is less than 37°C, a decision block 68 proceeds to a decision block 69. Decision block 69 determines whether \( N \) is greater than 1, and if not, the flow chart proceeds to a block 70. However, for subsequent iterations through decision block 68 that occur as described below, an affirmative response to decision block 69 leads to a block 72, wherein an average of the previous two estimates of the \( pO_2 \) at the reference temperature, 37°C, is determined.

\[ X_N = \frac{X_N + X_{N-1}}{2} \]  

This average replaces the current estimate, \( X_N \), which is used for the next evaluation in block 70. In block 70, a more accurate estimate, \( X_{N+1} \), of the \( pO_2 \) at the reference temperature is determined as a function of the measured \( pO_2 \), the temperature at the measurement site, and the last estimate \( X_N \) of the \( pO_2 \) at the reference temperature (or its current value based on the average from block 72, for measurements made at a temperature less than the reference temperature).

\[ X_{N+1} = X_0 10 \]  

The flow chart then proceeds to a block 74 wherein \( N \) is incremented by one. A decision block 76 determines whether \( N \) exceeds a predetermined value representing the number of iterations desired. In the preferred embodiment, this predefined value is 9. If fewer than nine iterations have been made, decision block 76 loops back to decision block 68, iteratively repeating the evaluation to determine the more accurate estimate \( X_N \) of the \( pO_2 \) at the reference temperature. Once the number of iterations has reached the predefined number, block 78 displays the last more accurate estimate of the \( pO_2 \) at 37°C for use by medical personnel. Subsequently, the algorithm stops in a block 80 until the value for \( pO_2 \) at the measurement temperature is updated, which may occur as many as 100 times per second.

Referring now to FIGURE 3, the result of applying the iterative process to determine the \( pO_2 \) at the reference temperature is shown in terms of a percentage error for four different \( pO_2 \) levels and twelve measurement temperatures ranging from approximately 25°C to 40°C. The percentage errors shown are referenced to
measurements of pO₂ in blood samples at the reference temperature that were made on a Corning Model 178 blood gas analyzer.

Although a pO₂ at the reference temperature can be determined from a measurement at a temperature less than the reference temperature without the averaging step carried out in block 72 of FIGURE 2, a greater number of iterations are required to achieve the same acceptable convergence limit. In the preferred embodiment, an acceptable convergence limit is ± 0.5 Torr. A maximum of nine iterations are required to achieve this convergence limit, as reflected in decision block 76.

In FIGURE 4, the relationship between measurement temperature, pO₂ level, and the number of iterations required to achieve ± 0.5 Torr convergence limit are shown in a three-dimensional graph. As shown in this FIGURE, it is only when the measurement temperature is below 25°C that up to nine iterations are required to achieve the desired accuracy.

Although the method for converting a value for pO₂ measured at one temperature to a corresponding value at the reference temperature has been disclosed with respect to measurements made in a patient’s blood stream, it should be apparent that the same technique can be applied to other environments, for example, when the measurement is made in a specimen or sample of blood taken from a patient. Furthermore, the method for converting from pO₂ at an arbitrary temperature to a corresponding value for the pO₂ at the reference temperature of 37°C is also applicable to other techniques for measuring pO₂. While the preferred embodiment of the invention has been illustrated and described, it will be appreciated that various changes can be made therein without departing from the spirit and scope of the invention. Accordingly, it is not intended that the scope of the invention in any way be limited by the disclosure, but instead that it be determined entirely by reference to the claims that follow.
The embodiments of the invention in which an exclusive property or privilege is claimed are defined as follows:

1. A method for determining a partial pressure of oxygen at a predefined reference temperature during an in vivo blood gas measurement, comprising the steps of:
   (a) measuring a partial pressure of oxygen at an arbitrary temperature that is substantially different than the reference temperature, as a function of a physical parameter that changes to indicate the partial pressure of oxygen at a measurement site, said step of measuring including the steps of:
      (i) producing a first signal indicative of the partial pressure of oxygen, in response to changes in the physical parameter;
      (ii) producing a second signal indicative of the arbitrary temperature at the measurement site; and
      (iii) processing the first signal to determine the partial pressure of oxygen at the arbitrary temperature;
   (b) determining an estimate of a partial pressure of oxygen at the predefined reference temperature as a function of the partial pressure of oxygen at the arbitrary temperature, and of the second signal;
   (c) determining a more accurate estimate of the partial pressure of oxygen at the predefined reference temperature as a function of:
      (i) the partial pressure of oxygen at the arbitrary temperature;
      (ii) the second signal; and
      (iii) the estimate of the partial pressure of oxygen at the predefined reference temperature; and
   (d) iteratively repeating step (c), each iteration using said more accurate estimate of the partial pressure of oxygen from a previous iteration as a value for said estimate in (c)(iii), until a predefined number of such iterations have been completed, a more accurate estimate that results from a last of such iterations approximating the partial pressure of oxygen at the predefined reference temperature corresponding to the measurement of the partial pressure of oxygen made at the arbitrary temperature.

2. The method of Claim 1, wherein each iteration of step (c) determines said more accurate estimate as defined by:
\[ X_{N+1} = X_0 \cdot 10^{\left( \frac{3X_T - T}{K_1} \right) \cdot f(X_N)} \]

where:

- \( X_0 \) is the partial pressure of oxygen measured at the arbitrary temperature;
- \( T \) is the arbitrary temperature;
- \( T_R \) is the predefined reference temperature;
- \( X_{N+1} \) is the more accurate estimate of the partial pressure of oxygen for an \((N + 1)\)th iteration;
- \( X_N \) is the more accurate estimate of the partial pressure of oxygen for an \(N\)th iteration;
- \( K_1 \) is a predetermined constant; and
- \( f(X_N) \) is a predefined function of \( X_N \).

3. The method of Claim 2, wherein the predefined function \( f(X_N) \) is represented by:

\[
f(X_N) = \frac{K_2}{K_3 \left( \frac{X_N}{K_4} \right)} + K_5 + K_6
\]

where:

- \( K_2 \) through \( K_6 \) are predefined constants.

4. The method of Claim 1, wherein if the arbitrary temperature is less than the reference temperature, the method further comprises the steps of:

(a) for each iteration after a first iteration, determining an average of the previous two more accurate estimates of the partial pressure of oxygen; and

(b) using the average as the value for said estimate in a next iteration, in step (c)(iii).

5. The method of Claim 1, wherein the predefined number of iterations is selected so that the more accurate estimate of the partial pressure of oxygen at the predefined reference temperature converges to a value within a predefined convergence limit.
6. The method of Claim 1, wherein the predefined number of iterations is less than ten.

7. A method for determining a partial pressure of oxygen in a patient’s blood, at a predefined reference temperature, comprising the steps of:
   (a) transmitting a light signal to a measurement site in the patient’s body, said light signal causing a material sensitive to the light signal to emit a return light signal having a parameter that varies as a function of the partial pressure of oxygen that is present in the blood at the measurement site;
   (b) detecting the return light signal, producing an electrical signal corresponding thereto, said electrical signal being indicative of the partial pressure of oxygen at a temperature of the blood at the measurement site;
   (c) producing a temperature signal indicative of the temperature of the blood at the measurement site at the time the return light signal is produced, said temperature being substantially different that the reference temperature;
   (d) digitizing the temperature signal and the electrical signal;
   (e) determining an estimate of a partial pressure of oxygen at the predefined reference temperature as a function of the digitized electrical signal, which represents the partial pressure of oxygen at the temperature of blood at the measurement site, and as a function of the digitized temperature at the measurement site;
   (f) determining a more accurate estimate of the partial pressure of oxygen at the predefined reference temperature as a function of:
      (i) the digitized electrical signal;
      (ii) the digitized temperature at the measurement site; and
      (iii) the estimate of the partial pressure of oxygen at the predefined reference temperature; and
   (g) iteratively repeating step (f), each iteration using said more accurate estimate of the partial pressure of oxygen from a previous iteration as a value for said estimate in (f)(iii), until a predefined number of such iterations have been completed, a more accurate estimate that results from a last of such iterations approximating the partial pressure of oxygen at the predefined reference temperature corresponding to the measurement of the partial pressure of oxygen made at the temperature of blood at the measurement site.
8. The method of Claim 7, wherein the step of transmitting comprises the step of exciting the material to phosphorescence, causing it to emit the return light signal as phosphorescent light for a period of time dependent upon the partial pressure of oxygen at the measurement site.

9. The method of Claim 7, wherein each iteration of step (f) determines said more accurate estimate as defined by:

\[ X_{N+1} = X_0 10^{\left( \frac{T_{n+1} - T}{K_1} \right) f(X_n)} \]

where:

- \( X_0 \) is the partial pressure of oxygen measured at the temperature of the measurement site;
- \( T \) is the temperature of the measurement site;
- \( T_R \) is the predefined reference temperature;
- \( X_{N+1} \) is the more accurate estimate of the partial pressure of oxygen for an \((N + 1)\)th iteration;
- \( X_N \) is the more accurate estimate of the partial pressure of oxygen for an \(N\)th iteration;
- \( K_1 \) is a predetermined constant; and
- \( f(X_n) \) is a predefined function of \( X_N \).

10. The method of Claim 9, wherein the predefined function \( f(X_N) \) is represented by:

\[ X_{N+1} = X_0 10^{\left( \frac{T_{n+1} - T}{K_1} \right) f(X_n)} \]

where:

- \( K_2 \) through \( K_6 \) are predefined constants.

11. The method of Claim 9, wherein the predefined function \( f(X_N) \) is represented by:

\[ X_{N+1} = X_0 10^{\left( \frac{37 - T}{2303} \right) \left( \frac{0.058}{243 \left( \frac{X_n}{100} \right)^{1.1}} + 0.013 \right)} \]
12. The method of Claim 7, wherein if the temperature at the measurement site is less than the reference temperature, the method further includes the steps of:

(a) for each iteration after a first iteration, determining an average of the previous two more accurate estimates of the partial pressure of oxygen; and

(b) using the average as the value for said estimate in a next iteration, in step (f)(iii).

13. The method of Claim 7, wherein the predefined number of iterations is selected so that the more accurate estimate of the partial pressure of oxygen at the reference temperature converges to a value within a predefined convergence limit.

14. The method of Claim 7, wherein the predefined number of iterations is less than ten.

15. A method for determining a partial pressure of oxygen at a predefined reference temperature at a measurement site disposed on a distal end of an optical fiber within a patient’s cardiovascular system, a temperature at said measurement site being substantially different than the reference temperature, comprising the steps of:

(a) conveying a pulse of light through the optical fiber toward its distal end, said light exciting a phosphorescent material disposed on the distal end of the optical fiber into phosphorescence that is quenched at a rate dependent upon the partial pressure of oxygen at the measurement site, producing a return light signal that travels through the optical fiber;

(b) determining a decay time for the return light signal, and based upon the decay time, the partial pressure of oxygen at the measurement site at the temperature of the measurement site;

(c) sensing the temperature of the measurement site, producing a temperature signal indicative of said temperature;

(d) determining a first estimate of the partial pressure of oxygen at the predefined reference temperature as a function of the partial pressure of oxygen that was determined from the decay time and as a function of the temperature signal;
(e) determining a more accurate estimate of the partial pressure of oxygen at the predefined reference temperature as a function of:

(i) the partial pressure of oxygen that was determined from the decay time;

(ii) the temperature signal; and

(iii) the first estimate of the partial pressure of oxygen at the predefined reference temperature; and

(f) iteratively repeating step (e), a first such iteration substituting the more accurate estimate of the partial pressure of oxygen at the predefined reference temperature for the first estimate of step (e)(iii), and a subsequent iteration substituting the more accurate estimate of the partial pressure of oxygen determined from the previous iteration for the value previously used in step (e)(iii) in the iteration before, the more accurate estimate from the last of a predefined number of such iterations closely approximating the partial pressure of oxygen at the predefined reference temperature.

16. The method of Claim 15, wherein if the temperature at the measurement site is less than the reference temperature, after the first iteration, an average of the previous two more accurate estimates of the partial pressure of oxygen at the reference temperature is substituted for the last estimate in each further iteration.

17. The method of Claim 15, wherein the more accurate estimate is defined by:

\[
X_{N+1} = X_0 \times 10^{\left(\frac{37-T}{2.303} \times \left(\frac{0.58}{243 (X_{N+1}^{37^\circ C})^{0.100}} + 0.013\right)\right)}
\]

where:

- \(X_0\) is the partial pressure of oxygen determined as a function of the partial pressure of oxygen at the temperature of the measurement site;
- \(T\) is the temperature of the measurement site;
- \(X_{N+1}\) is the more accurate estimate of the partial pressure of oxygen for an \((N + 1)\)th iteration;
- the reference temperature equals 37°C; and
- \(X_N\) is the more accurate estimate of the partial pressure of oxygen for an \(N\)th iteration.
18. The method of Claim 15, wherein the predefined number of iterations is selected to determine a final approximation of the partial pressure of oxygen at the reference temperature that is within an accepted convergence limit.

19. The method of Claim 18, wherein the predefined number of iterations is less than ten and the accepted convergence limit is less than ±0.5 Torr.

20. The method of Claim 18, wherein fewer iterations are used to obtain a final approximation of the partial pressure of oxygen at the reference temperature that is within the accepted convergence limit when the temperature at the measurement site is greater than the reference temperature, as compared to when it is less.
START

A-D CONVERSION OF MEASURE TEMP.

INITIATE LIGHT PULSE TRANSMISSION THRU OPTICAL FIBER

A-D CONVERSION OF PHOSPHORESCENCE SIGNAL

DETERMINE DECAY RATE

USING DECAY RATE DETERMINE $p_{O_2} @ T$ (REPRESENTED BY $X_0$)

IS $T=37^\circ C$ ?

YES

NO

SET $N=1$

EVALUATE:

$$X_N = X_0 \cdot 10^{ \frac{(37-T)}{2.303} \left( \frac{0.58}{X_0} + \frac{0.013}{100} \right) \left( \frac{X_0}{100} \right) }$$

EVALUATE

$$X_N = \frac{X_N + X_{N-1}}{2}$$

IS $N > 1$ ?

YES

NO

IS $T < 37^\circ C$ ?

YES

NO

EVALUATE:

$$X_{N+1} = X_0 \cdot 10^{ \frac{(37-T)}{2.303} \left( \frac{0.58}{X_{N+1}} + \frac{0.013}{100} \right) \left( \frac{X_{N+1}}{100} \right) }$$

$N = N+1$

$N = 9$

DISPLAY $p_{O_2} @ 37^\circ C$

STOP
# INTERNATIONAL SEARCH REPORT

## A. CLASSIFICATION OF SUBJECT MATTER

- **IPC(5):** A61B 5/02 G01N 33/48; F21V 9/16
- **US CL.:** 128/666,356/41 250/459.1

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

- **Minimum documentation searched (classification system followed by classification symbols):**
  - U.S.: 128/633,634,634,664,665; 356/39,40
  - 250/458.1

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>US, A, 5,058,588 (KAESTLE) 22 OCTOBER 1991 Figures and abstract</td>
<td>1-20</td>
</tr>
<tr>
<td>A</td>
<td>US, A, 4,697,593 (EVANS ET AL) 06 OCTOBER 1987 Entire reference</td>
<td>1-20</td>
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<tr>
<td>A</td>
<td>US, A, 4,752,115 (MURAY, JR. ET AL) 21 JUNE 1988 Figures, claims</td>
<td>1-20</td>
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</table>

- **X** Further documents are listed in the continuation of Box C.  
  - **See patent family annex.**

- **"A"** Special categories of cited documents:
  - "I" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
  - "E" earlier document published on or after the international filing date
  - "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
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  - "O" document referring to an oral disclosure, use, exhibition or other means
  - "P" document published prior to the international filing date but later than the priority date claimed

**Form PCT/ISA/210 (second sheet)(July 1992)**

**Date of the actual completion of the international search:** 07 FEBRUARY 1993

**Date of mailing of the international search report:** 08 MAR 1993

**Name and mailing address of the ISA/US Commissioner of Patents and Trademarks:**
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  - Facsimile No. NOT APPLICABLE

**Authorized officer:**
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<td>US,A, 5,030,420 (BACON ET AL) 09 JULY 1991 Figures and abstract</td>
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