EQUILIBRIUM NON-CONSUMING FLUORESCENCE SENSOR FOR REAL TIME INTRAVASCULAR GLUCOSE MEASUREMENT

Abstract: Embodiments of the present invention relate to analyte sensors. In particular, the preferred embodiments of the present invention relate to non-consuming intravascular glucose sensors based on fluorescence chemistry.
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EQUILIBRIUM NON-CONSUMING FLUORESCENCE SENSOR FOR REAL TIME INTRAVASCULAR GLUCOSE MEASUREMENT

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] This application claims the priority benefit to U.S. Provisional No. 60/917,307 filed May 10, 2007, the entirety of which is hereby incorporated by reference herein.

BACKGROUND OF THE INVENTION

Field of the Invention

[0002] Embodiments of the present invention relates to analyte sensors. In particular, the preferred embodiments of the present invention relate to non-consuming intravascular glucose sensors based on fluorescence chemistry.

Description of the Related Art

[0003] There has been an on-going effort over many years to use fluorescence techniques to measure polyhydroxyl compound (e.g., glucose) concentration in bodily fluids. But despite the effort, no practical system has been developed and commercialized for in vivo monitoring. Several attempts have been made to detect glucose by fluorescence using dyes associated with boronic acid groups. Boronate moieties bind glucose reversibly. When boronic acid functionalized fluorescent dyes bind glucose, the properties of the dye are affected, such that a signal related to the concentration of glucose may be generated and detected. These changes have been used in the past to measure glucose concentration.

[0004] Russell (U.S. Pat. Nos. 5,137,833 and 5,512,246) used a boronic acid functionalized dye that bound glucose and generated a signal related to the glucose concentration. James et al. (U.S. Pat. No. 5,503,770) employed a similar principle, but combined a fluorescent dye, an amine quenching functionality, and boronic acid in a single complex. The fluorescence emission from the complex varied with the amount of glucose binding. Van Antwerp et al. (U.S. Pat. Nos. 6,002,954 and 6,01 1,984) combined features of the previously cited references and also disclosed a device purported to be implantable. A. E.
Colvin, Jr. (U.S. Pat. No. 6,304,766) also disclosed optical-based sensing devices for *in situ* sensing in humans that utilize boronate-functionalized dyes.

[0005] Certain measurable parameters using blood or bodily fluid, such as pH and concentrations of $O_2$, $CO_2$, $Na^+$, $K^+$, and polyhydroxyl compounds, like glucose, have been determined *in vivo*. The ability to do these measurements *in vivo* is important because it is necessary to make frequent determinations of such analytes when monitoring a patient. Typically, one sensor for each analyte has been placed in a patient’s blood vessel(s). If it is desired to measure several analytes, a plurality of sensors is often required, which can cause attendant discomfort to the patient and complexity of the electronic monitoring equipment.

[0006] In an effort to solve the design problems posed by the limitation in physical dimension for *in vivo* monitoring, others have incorporated different dyes into one device to get simultaneous readings of two parameters. For example, Alder et al. (U.S. Pat. No. 5,922,612) disclosed a method for optical determination of pH and ionic strength of an aqueous sample using two different dyes on one sensor. Gray et al. (U.S. Pat. No. 5,176,882) taught the use of a fiber optic device incorporating a hydrophilic polymer with immobilized pH sensitive dye and potassium or calcium sensitive fluorescent dyes to measure the analyte concentration in conjunction with pH. In U.S. Pat. No. 4,785,814. Kane also disclosed the use of two dyes embedded in a composite membrane for the simultaneous measurements of pH and oxygen content in blood. However, incorporation of multiple dyes into a single sensor complicates the manufacture of such sensors.

[0007] Besides the foregoing problems associated with separate indwelling sensors for each analyte being monitored, particularly in the intensive care setting, and multiple dye sensors, another problem associated with many dye-based analyte sensors is pH sensitivity. A slight change in pH may modify or attenuate fluorescence emissions, and cause inaccurate readings. This problem is particularly acute for monitoring blood glucose levels in diabetic patients, whose blood pH may fluctuate rapidly. Since accurate blood glucose level measurements are essential for treating these patients, there is a significant need for a glucose sensor that facilitates real-time correction of the pH effect without requiring separate indwelling pH and analyte sensors, or sensors having multiple dyes.
[0008] Ratjometric pH determination using fluorescent dye(s) is known. Given a fluorophore that has an acid and base form, the ratio of the emission intensity of the two forms can be used as a measure of the pH that is insensitive to fluorophore concentration. See e.g., U.S. Patent Publication No. 2005/0090014 which describes an HPTS-derived pH sensitive dye (incorporated herein in its entirety by reference); Niu CG. et al. 2005 Anal. Bioanal. Chem. 383(2):349-357, which describes a pH-sensitive dye meso-5,10,15,20-tetra-(4-allyloxyphenyl)porphyrin (TAPP) as an indicator, and a pH-insensitive benzothioxanthene derivative as a reference, for fluorescence ratiometric measurement; Turner N.G. et al. 1998 J. Investig. Dermatol. Symp. Proc. Aug 3(2):110-3, which discloses dual-emission ratiometric imaging using the fluorophore, carboxy seminaphthorhodafluor-1, which displays a pH-dependent shift in its emission spectrum; and Badugu R. et al. 2005 Talanta 66:569-574, which describes the use of 6-aminoquinolinium boronic acid dyes that show spectral shifts and intensity changes with pH in a wavelength-ratiometric manner.

[0009] However, despite the inventor’s recognition of a substantial unmet need for a sensor adapted to provide continuous intravascular monitoring of pH and glucose, wherein the glucose measurement may be corrected for pH effects, no one has disclosed or even suggested using a sensor comprising a single fluorophore that exhibits properties suitable to make a ratiometric pH measurement that is independent of the fluorophore concentration, where the same fluorophore is functionalized to bind glucose and generate a signal the intensity of which is related to the glucose concentration.

SUMMARY OF THE INVENTION

[0010] An intravascular sensor is disclosed in accordance with preferred embodiments of the present invention for determining the analyte concentration in blood. The sensor comprises an analyte binding molecule, wherein the analyte binding molecule is associated with a fluorophore; an analyte analog, wherein the analyte analog is associated with an acceptor, wherein the fluorophore is capable of emitting radiation at a wavelength that is absorbed at least in part by the acceptor, wherein the fluorophore is capable of transferring energy to the acceptor via fluorescence resonance energy transfer when the fluorophore is in close proximity to the acceptor; a light source; and a detector.
In some embodiments, the sensor further comprises a hydrogel, wherein the analyte binding molecule and the analyte analog are substantially immobilized in the hydrogel, wherein the hydrogel is permeable to the analyte.

In some embodiments, the light source is a laser.

In some embodiments, the laser is capable of delivering a frequency modulated excitation signal.

In some embodiments, the laser is capable of delivering a pulse of light.

In some embodiments, the sensor further comprises an optical fiber.

In some embodiments, the analyte is glucose.

A method is disclosed in accordance with preferred embodiments of the present invention for determining the glucose concentration in blood. The method comprises providing the sensor as described above; inserting the sensor into a blood vessel; irradiating the fluorophore with an excitation signal; detecting a fluorescence emission from the fluorophore; and determining the concentration of glucose.

In some embodiments, the excitation signal is a pulse of light.

In some embodiments, the method further comprises measuring the decay of the fluorescence emission over time and calculating a fluorescence lifetime.

In some embodiments, the excitation signal is frequency modulated.

In some embodiments, the method further comprises measuring the phase shift between the emission and the excitation signal.

In some embodiments, the method further comprises calculating a fluorescence lifetime.

In some embodiments, the method further comprises measuring a modulation ratio and calculating a fluorescence lifetime.

In one preferred embodiment, an intravascular sensor is disclosed for determining an analyte concentration in blood. The sensor comprises an optical fiber comprising a sensor chemistry portion disposed along a distal region of the optical fiber, wherein the sensor chemistry portion is sized and physiologically compatible with residing in a blood vessel, wherein the sensor chemistry portion is selected from equilibrium fluorescence chemistry or lifetime chemistry.
BRIEF DESCRIPTION OF THE DRAWINGS

[0025] FIG. 1 is a flow chart showing the sensing mechanism of one embodiment of the present invention.

[0026] FIG. 2 shows a glucose and pH sensor and optical system comprising two excitation light sources and two detectors in accordance with one preferred embodiment of the present invention.

[0027] FIG. 3 shows the absorption spectra of HPTS at different pHs.

[0028] FIG. 4 shows independence of ratiometric pH sensing using HPTS/MABP4 using the $I_{(base)} / I_{(iso)}$ ratio from glucose concentration. The data are plotted as a ratio of the fluorescence emission for corresponding to excitation at 454 nm (base) and 422 nm (isobestic point) vs. pH in various glucose concentrations.

[0029] FIG. 5 shows glucose response curves for HPTS/MABP4 excited at 422 nm (isobestic point) at different pHs.

[0030] FIG. 6 shows the absorption spectra of SNARF-1 at different pHs in solution.

[0031] FIG. 7 shows glucose response curves for SNARF-1/3,3'-oBBV in solution at different pHs excited at 514 nm/emission at 587 nm.

[0032] FIG. 8 shows glucose response curves for SNARF-1/3,3'-oBBV in solution at different pHs excited at 514 nm/emission at 625 nm.

[0033] FIG. 9 shows ratiometric sensing of pH at different glucose concentrations with SNARF-1/3,3'-oBBV in solution using the $I_{(base)} / I_{(acid)}$ ratio.

[0034] FIG. 10 shows glucose response curves for HPTS-triLysMA/3,3'-oBBV/DMMA at different pHs.

[0035] FIG. 11 shows ratiometric sensing of pH at different glucose concentrations using HPTS-triLysMA/3,3'-oBBV/DMMA, using the $I_{(base)} / I_{(acid)}$ ratio.

[0036] FIG. 12 shows ratiometric sensing of pH at different glucose concentrations using HPTS-triCysMA/3,3'-oBBV/DMMA wherein the indicator system is immobilized on the end of an optical fiber, using the $I_{(base)} / I_{(acid)}$ ratio.
[0037] FIG. 13 shows a graph of the decay of fluorescence intensity over time after a fluorophore is subjected to a pulse of excitation light.

[0038] FIG. 14 shows a graph of the emission signal resulting from a frequency modulated excitation signal.

[0039] FIGS. 15A-C show the interaction between a glucose binding molecule linked to a fluorophore, a glucose analog linked to an acceptor and a glucose molecule.
DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

[0040] In a preferred embodiment, the present invention is directed to an optical sensor capable of measuring two analytes with a single indicator system. More particularly, the preferred sensor employs a single fluorophore (e.g., a fluorescent dye) to: (1) determine the concentration of a first analyte, e.g., H⁺ (pH), by a ratiometric method, wherein such determination is independent of the concentration of the fluorophore; and (2) determine the concentration of a second analyte, e.g., a polyhydroxyl compounds (e.g., preferably glucose) by measuring the apparent fluorophore concentration (e.g., emission intensity of the fluorophore upon excitation), wherein the apparent fluorophore concentration is dependent on the concentration of the second analyte. Further, where measurement of the second analyte concentration is dependent on the first analyte concentration (e.g., in optical systems in which glucose measurement varies with pH—a common problem in this field), then in accordance with a preferred embodiment of the present invention, the measured second analyte concentration may be corrected for the contribution of the first analyte concentration.

The sensor is preferably stable in aqueous media (e.g., physiological media, blood, interstitial fluid, etc.), and more preferably, the sensor is configured to be inserted into a blood vessel where it can remain indwelling for a period of time. Thus, in accordance with a preferred embodiment of the present invention, an optical sensor configured for intravascular placement is disclosed, which sensor is capable of measuring two analytes (preferably pH and glucose) with a single indicator system and correcting the glucose measurement for any contributions of pH.

[0041] Although preferred embodiments of the sensor are directed inter alia to ratiometric pH sensing, other first analyte concentrations may be determined in accordance with the broader scope of the present invention, as long as the indicator system comprises a fluorophore that exists in at least two forms the concentration of which are associated with the concentration of the first analyte and the emission ratio of which is independent of the fluorophore concentration. Likewise, although glucose is used as a second analyte example herein, it is understood that the concentration of other polyhydroxyl-containing organic compounds (carbohydrates, 1,2-diols, 1,3-diols and the like) in a solution may be determined using embodiments of this invention, as long as the indicator system comprises a fluorophore.
that is operably coupled to a binding moiety that binds the second analyte, wherein the signal intensity of the fluorophore varies with the concentration of second analyte. In some embodiments, the concentration of second analytes may including non-carbohydrates.

**Indicator System**

[0042] The indicator systems used in accordance with preferred embodiments of the present invention comprise a fluorophore operably coupled to an analyte binding moiety, wherein analyte binding causes an apparent optical change in the fluorophore concentration (e.g., emission intensity). It is further desired that the fluorophore has different acid and base forms that exhibit a detectable difference in spectral properties such that ratiometric pH sensing may be enabled. For example, a glucose binding moiety such as 3,3'-oBBV (described in detail below) that is operably coupled to a fluorescent dye such as HPTS-triLysMA (described in detail below) will quench the emission intensity of the fluorescent dye, wherein the extent of quenching is reduced upon glucose binding resulting in an increase in emission intensity related to glucose concentration. In preferred embodiments, the indicator systems comprise a dye having at least two anionic groups and a quencher having at least two boronic acids. In further preferred embodiments, the indicator systems also comprise a means for immobilizing the sensing moieties (e.g., dye-quencher) such that they remain physically close enough to one another to react (quenching). Where in vivo sensing is desired, such immobilizing means are preferably insoluble in an aqueous environment (e.g., intravascular), permeable to the target analytes, and impermeable to the sensing moieties. Typically, the immobilizing means comprises a water-insoluble organic polymer matrix. For example, the HPTS-triLysMA dye and 3,3'-oBBV quencher may be effectively immobilized within a DMAA (N,M-dimethylacrylamide) hydrogel matrix (described in detail below), which allows pH and glucose sensing in vivo.

[0043] Some exemplary and preferred fluorophores, analyte binding moieties and immobilizing means are set forth in greater detail below.

**Fluorophores**

[0044] "Fluorophore" refers to a substance that when illuminated by light at a particular wavelength emits light at a longer wavelength; i.e. it fluoresces. Fluorophores include but are not limited to organic dyes, organometallic compounds, metal chelates,
fluorescent conjugated polymers, quantum dots or nanoparticles and combinations of the above. Fluorophores may be discrete moieties or substituents attached to a polymer.

[0045] Fluorophores that may be used in preferred embodiments are capable of being excited by light of wavelength at or greater than about 400 nm, with a Stokes shift large enough that the excitation and emission wavelengths are separable by at least 10 nm. In some embodiments, the separation between the excitation and emission wavelengths may be equal to or greater than about 30 nm. These fluorophores are preferably susceptible to quenching by electron acceptor molecules, such as viologens, and are resistant to photo-bleaching. They are also preferably stable against photo-oxidation, hydrolysis and biodegradation.

[0046] In some embodiments, the fluorophore may be a discrete compound.

[0047] In some embodiments, the fluorophore may be a pendant group or a chain unit in a water-soluble or water-dispersible polymer having molecular weight of about 10,000 daltons or greater, forming a dye-polymer unit. In one embodiment, such dye-polymer unit may also be non-covalently associated with a water-insoluble polymer matrix M1 and is physically immobilized within the polymer matrix M1, wherein M1 is permeable to or in contact with analyte solution. In another embodiment, the dye on the dye-polymer unit may be negatively charged, and the dye-polymer unit may be immobilized as a complex with a cationic water-soluble polymer, wherein said complex is permeable to or in contact with the analyte solution. In one embodiment, the dye may be one of the polymeric derivatives of hydroxypyrene trisulfonic acid. The polymeric dyes may be water-soluble, water-swellable or dispersible in water. In some embodiments, the polymeric dyes may also be cross-linked. In preferred embodiments, the dye has a negative charge.

[0048] In other embodiments, the dye molecule may be covalently bonded to the water-insoluble polymer matrix M1, wherein said M1 is permeable to or in contact with the analyte solution. The dye molecule bonded to M1 may form a structure M1-L1-Dye. L1 is a hydrolytically stable covalent linker that covalently connects the sensing moiety to the polymer or matrix. Examples of L1 include lower alkylene (e.g., C1-C8 alkylene), optionally terminated with or interrupted by one or more divalent connecting groups selected from sulfonamide (--SO2NH--), amide --(C=O)N--, ester --(C=O)--O--, ether.--O--, sulfide --S--,
sulfone (--SO$_2$--), phenylene --C$_6$H$_4$--, urethane --NH(C=O)--O--, urea --NH(C=O)NH--, thiourea --NH(C=S)--NH--, amide --(C=O)NH--, amine --NR-- (where R is defined as alkyl having 1 to 6 carbon atoms) and the like, or a combination thereof. In one embodiment, the dye is bonded to a polymer matrix through the sulfonamide functional groups.

[0049] In some embodiments, useful dyes include pyranine derivatives (e.g. hydroxypyrene trisulfonic acid derivatives and the like), which have the following formula:

![Pyranine derivative structure]

wherein R$^1$, R$^2$, R$^3$ are each -NHR$^4$, R$^4$ is -CH$_2$CH$_2$(-OCH$_2$CH$_2$)$_n$X$^1$; wherein X$^1$ is -OH, -OCH$_3$COOH, -CONH$_2$, -SO$_3$H, -NH$_2$, or OMe; and n is between about 70 and 10,000. In one embodiment, the dyes may be bonded to a polymer through the sulfonamide functional groups. In other embodiments, the dye may be one of the polymeric derivatives of hydroxypyrene trisulfonic acid.

[0050] In some embodiments, the fluorescent dye may be 8-hydroxypyrene-1,3,6-trisulfonate (HPTS). The counterions can be H$^+$ or any other cation. HPTS exhibits two excitation wavelengths at around 450 nm and around 405 nm, which correspond to the absorption wavelengths of the acid and its conjugate base. The shift in excitation wavelength is due to the pH-dependent ionization of the hydroxyl group on HPTS. As the pH increases, HPTS shows an increase in absorbance at about 450 nm, and a decrease in absorbance below about 420 nm. The pH-dependent shift in the absorption maximum enables dual-excitation ratiometric detection in the physiological range. This dye has a molecular weight of less than 500 daltons, so it will not stay within the polymer matrix, but it can be used with an anion exclusion membrane.
[0051] In another embodiment, the fluorescent dye may be polymers of 8-acetoxy-pyrene-1,3,6-N, N',N"-tris-(methacrylpropylamidosulfonamide) (acetoxy-HPTS-MA):

[0052] It is noted that dyes such as acetoxy-HPTS-MA (above) having no anionic groups, may not give very strong glucose response when operably coupled to a viologen quencher, particularly a viologen quencher having only a single boronic acid moiety.

[0053] In another embodiment, the fluorescent dye may be 8-hydroxy-pyrene-1,3,6-N, N',N"-tris-(carboxypropylsulfonamide) (HPTS-CO$_2$):

[0054] In another embodiment, the fluorescent dye may be 8-hydroxy-pyrene-1,3,6-N, N',N"-tris-(methoxypolyethoxyethyl (-125) sulfonamide) (HPTS-PEG):
It is noted that dyes such as HPTS-PEG (above) having no anionic groups, may not provide a very strong glucose response when operably coupled to a viologen quencher, particularly a viologen quencher having only a single boronic acid moiety.

Representative dyes as discrete compounds are the tris adducts formed by reacting 8-acetoxypyrene-1,3,6-trisulfonylchloride (HPTS-Cl) with an amino acid, such as amino butyric acid. Hydroxypyrene trisulfonamide dyes bonded to a polymer and bearing one or more anionic groups are most preferred, such as copolymers of 8-hydroxypyrene-1-N-(methacrylamidopropylsulfonamido)-N,N"-3,6-bis(carboxypropylsulfonamide) HPTS-CO$_2$-MA with HEMA, PEGMA, and the like.

In another embodiment, the fluorescent dye may be HPTS-TriCys-MA:

This dye may be used with a quencher comprising boronic acid, such as 3,3'-oBBV.

Of course, in some embodiments, substitutions other than Cys-MA on the HPTS core are consistent with aspects of the present invention, as long as the substitutions are negatively charged and have a polymerizable group. Either L or D stereoisomers of
cysteine may be used. In some embodiments, only one or two of the sulfonic acids may be substituted. Likewise, in variations to HPTS-CysMA shown above, other counterions besides NBu₄⁺ may be used, including positively charged metals, e.g., Na⁺. In other variations, the sulfonic acid groups may be replaced with e.g., phosphoric, carboxylic, etc. functional groups.

[0060] Another suitable dye is HPTS-LysMA, which is pictured below as follows:

![HPTS-LysMA structure](image)

Other examples include soluble copolymers of 8-acetoxypyrene-1,3,6-N, N', N''-tris(methacrylamidopropylsulfonamide) with HEMA, PEGMA, or other hydrophilic comonomers. The phenolic substituent in the dye is protected during polymerization by a blocking group that can be removed by hydrolysis after completion of polymerization. Such suitable blocking groups, as for example, acetoxy, trifluoroacetoxy, and the like, are well known in the art.

[0061] Fluorescent dyes, including HPTS and its derivatives are known and many have been used in analyte detection. See e.g., U.S. Pat. Nos. 6,653,141, 6,627,177, 5,512,246, 5,137,833, 6,800,451, 6,794,195, 6,804,544, 6,002,954, 6,319,540, 6,766,183, 5,503,770, and 5,763,238; and co-pending U.S. Patent Application Nos. 11/296,898 and 60/833,081; each of which is incorporated herein in its entirety by reference thereto.

[0062] The SNARF and SNAFL dyes from Molecular Probes may also be useful fluorophores in accordance with aspects of the present invention. The structures of SNARF-1 and SNAFL-1 are shown below.
Additionally, a set of isomeric water-soluble fluorescent probes based on both the 6-aminoquinolinium and boronic acid moieties which show spectral shifts and intensity changes with pH, in a wavelength-ratiometric and colorimetric mariner may be useful in accordance with some embodiments of the present invention (See e.g., Badug, R. et al. 2005 Talanta 65 (3):762-768; and Badug, R. et al. 2005 Bioorg. Med. Chem. 13 (1):113-119); incorporated herein in its entirety by reference.

Another example of a fluorescence dye that may be pH and saccharide sensitive is tetrakis(4-sulfophenyl)porphine (TSPP)—shown below. TSPP may not work optimally in blood, where the porphyrin ring may react with certain metal ions, like ferric, and become non-fluorescent.

Additional examples of pH sensitive fluorescent indicators that may be useful for simultaneous determination of pH and glucose in the sensor of the present
invention are described in US 2005/0233465 and US 2005/0090014; each of which is incorporated herein by reference in its entirety.

**Analyte Binding Moieties—Quenchers**

[0067] In accordance with broad aspects of the present invention, the analyte binding moiety provides the at least dual functionality of being able to bind analyte and being able to modulate the apparent concentration of the fluorophore (e.g., detected as a change in emission signal intensity) in a manner related to the amount of analyte binding. In preferred embodiments, the analyte binding moiety is associated with a quencher. "Quencher" refers to a compound that reduces the emission of a fluorophore when in its presence. Quencher (Q) is selected from a discrete compound, a reactive intermediate which is convertible to a second discrete compound or to a polymerizable compound or Q is a pendant group or chain unit in a polymer prepared from said reactive intermediate or polymerizable compound, which polymer is water-soluble or dispersible or is an insoluble polymer, said polymer is optionally crosslinked.

[0068] In one example, the moiety that provides glucose recognition in the embodiments is an aromatic boronic acid. The boronic acid is covalently bonded to a conjugated nitrogen-containing heterocyclic aromatic bis-onium structure (e.g., a viologen). "Viologen" refers generally to compounds having the basic structure of a nitrogen containing conjugated N-substituted heterocyclic aromatic bis-onium salt, such as 2,2'-, 3,3'- or 4,4'-N,N' bis-(benzyl) bipyridium dihalide (i.e., dichloride, bromide chloride), etc. Viologen also includes the substituted phenanthroline compounds. The boronic acid substituted quencher preferably has a pKa of between about 4 and 9, and reacts reversibly with glucose in aqueous media at a pH from about 6.8 to 7.8 to form boronate esters. The extent of reaction is related to glucose concentration in the medium. Formation of a boronate ester diminishes quenching of the fluorophore by the viologen resulting in an increase in fluorescence dependent on glucose concentration. A useful bis-onium salt is compatible with the analyte solution and capable of producing a detectable change in the fluorescent emission of the dye in the presence of the analyte to be detected.

[0069] Bis-onium salts in the embodiments of this invention are prepared from conjugated heterocyclic aromatic di-nitrogen compounds. The conjugated heterocyclic
aromatic di-nitrogen compounds are selected from dipyridyls, dipyridyl ethylenes, dipyridyl phenylenes, phenanthrolines, and diazafluorenes, wherein the nitrogen atoms are in a different aromatic ring and are able to form anonium salt. It is understood that all isomers of said conjugated heterocyclic aromatic di-nitrogen compounds in which both nitrogens can be substituted are useful in this invention. In one embodiment, the quencher may be one of the bis-onium salts derived from 3,3'-dipyridyl, 4,4'-dipyridyl and 4,7-phenanthroline.

In some embodiments, the viologen-boronic acid adduct may be a discrete compound having a molecular weight of about 400 daltons or greater. In other embodiments, it may also be a pendant group or a chain unit of a water-soluble or water-dispersible polymer with a molecular weight greater than about 10,000 daltons. In one embodiment, the quencher-polymer unit may be non-covalently associated with a polymer matrix and is physically immobilized therein. In yet another embodiment, the quencher-polymer unit may be immobilized as a complex with a negatively charge water-soluble polymer.

In other embodiments, the viologen-boronic acid moiety may be a pendant group or a chain unit in a crosslinked, hydrophilic polymer or hydrogel sufficiently permeable to the analyte (e.g., glucose) to allow equilibrium to be established.

In other embodiments, the quencher may be covalently bonded to a second water-insoluble polymer matrix M^2, which can be represented by the structure M^2-L^2-Q. L^2 is a linker selected from the group consisting of a lower alkylene (e.g., C_1-C_8 alkylene), sulfonamide, amide, quaternary ammonium, pyridinium, ester, ether, sulfide, sulfone, phenylene, urea, thiourea, urethane, amine, and a combination thereof. The quencher may be linked to M^2 at one or two sites in some embodiments.

For the polymeric quencher precursors, multiple options are available for attaching the boronic acid moiety and a reactive group which may be a polymerizable group or a coupling group to two different nitrogens in the heteroaromatic centrally located group. These are:

a) a reactive group on a first aromatic moiety is attached to one nitrogen and a second aromatic group containing at least one -B(OH)2 group is attached to the second nitrogen;
b) one or more boronic acid groups are attached to a first aromatic moiety which is attached to one nitrogen and one boronic acid and a reactive group are attached to a second aromatic group which second aromatic group is attached to the second nitrogen; and

c) one boronic acid group and a reactive group are attached to a first aromatic moiety which first aromatic group is attached to one nitrogen, and a boronic acid group and a reactive group are attached to a second aromatic moiety which is attached to the second nitrogen; and

d) one boronic acid is attached to each nitrogen and a reactive group is attached to the heteroaromatic ring.

[0074] Preferred embodiments comprise two boronic acid moieties and one polymerizable group or coupling group wherein the aromatic group is a benzyl substituent bonded to the nitrogen and the boronic acid groups are attached to the benzyl ring and may be in the ortho-meta or para-positions.

[0075] In some embodiments, the boronic acid substituted viologen as a discrete compound useful for in vitro sensing may be represented by one of the following formulas:

\[
\begin{align*}
\text{Y}^1\text{-(CH}_2\text{)}h & \quad \text{Y}^2 \quad 2\text{X}^2 \\
\text{Y}^1\text{-(CH}_2\text{)}h & \quad \text{Y}^2 \quad 2\text{X}^2 \\
\text{Y}^1\text{-(CH}_2\text{)}h & \quad \text{Y}^2 \quad 2\text{X}^2
\end{align*}
\]

[0076] where \( n = 1-3 \), X is halogen, and \( Y^1 \) and \( Y^2 \) are independently selected from phenyl boronic acid (o- m- or p-isomers) and naphthyl boronic acid. In other embodiments, the quencher may comprise a boronic acid group as a substituent on the heterocyclic ring of a viologen.

[0077] A specific example used with TSPP is m-BBV:
The quencher precursors suitable for making sensors may be selected from the following:

- m-BBV

[0078]

- Other compounds with different structures are also possible.
The quencher precursor 3,3'-oBBV may be used with HPTS-LysMA or HPTS-CysMA to make hydrogels in accordance with preferred aspects of the invention.

Preferred quenchers are prepared from precursors comprising viologens derived from 3,3'-dipyridyl substituted on the nitrogens with benzylboronic acid groups and at other positions on the dipyridyl rings with a polymerizable group or a coupling group. Representative viologens include:

![Chemical structures of quenchers and viologens](image)
where \( L \) is \( L_1 \) or \( L_2 \) and is a linking group

\( Z \) is a reactive group; and

\( R_1 \) is \(-\text{B(OH)}_2\) in the ortho- meta- or para- positions on the benzyl ring and \( R'' \) is \( \text{H}^- \); or optionally \( R'' \) is a coupling group as is defined herein or a substituent specifically used to modify the acidity of the boronic acid such as fluoro- or methoxy-

\( L \) is a divalent moiety that covalently connects the sensing moiety to a reactive group that is used to bind the viologen to a polymer or matrix. Examples of \( L \) include those which are each independently selected from a direct bond or, a lower alkylene having 1 to 8 carbon atoms, optionally terminated with or interrupted by one or more divalent connecting groups selected from sulfonamide (-SO_2NH-), amide -(C=O)N-, ester -(C=O)-O-, ether -O-, sulfide -S-, sulfone (-SO_2-), phenylene \(-\text{C}_6\text{H}_4\)-, urethane \(-\text{NH(C=O)}\text{O-}\), urea \(-\text{NH(C=O)}\text{NH-}\), thiourea \(-\text{NH(C=S)}\text{NH-}\), amide \(-\text{NH(C=O)}\text{NH-}\), amine \(-\text{NR-}\) (where \( R \) is defined as alkyl having 1 to 6 carbon atoms) and the like.

\( Z \) is either a polymerizable ethylenically unsaturated group selected from but not limited to methacrylamido-, acrylamido-, methacryloyl-, acryloyl-, or styryl- or optionally \( Z \) is a reactive functional group, capable of forming a covalent bond with a polymer or matrix. Such groups include but are not limited to \(-\text{Br}, -\text{OH}, -\text{SH}, -\text{CO}_2\text{H}, \) and \(-\text{NH}_2\).

Boronic add substituted polyviologens are another class of preferred quenchers. The term polyviologen includes: a discrete compound comprised of two or more viologens covalently bonded together by a linking group, a polymer comprised of viologen repeat units in the chain, a polymer with viologen groups pendant to the chain, a dendrimer comprised of viologen units, preferably including viologen terminal groups, an oligomer comprised of viologen units, preferably including viologen endgroups, and combinations thereof. Polymers in which mono-viologen groups form a minor component are not included.
The preferred quenchers are water soluble or dispersible polymers, or crosslinked, hydrophilic polymers or hydrogels sufficiently permeable to glucose to function as part of a sensor. Alternatively the polyviologen boronic acid may be directly bonded to an inert substrate.

[0087] A polyviologen quencher as a polymer comprised of viologen repeat units has the formula:

![Chemical Structure]

[0088] In another embodiment, the polyviologen boronic acid adducts are formed by covalently linking two or more viologen/boronic acid intermediates. The bridging group is typically a small divalent radical bonded to one nitrogen in each viologen, or to a carbon in the aromatic ring of each viologen, or one bond may be to a ring carbon in one viologen and to a nitrogen in the other. Two or more boronic acid groups are attached to the polyviologen. Optionally, the polyviologen boronic adduct is substituted with a polymerizable group or coupling group attached directly to the viologen or to the bridging group. Preferably the polyviologen moiety includes only one such group. Preferably, the bridging group is selected to enhance cooperative binding of the boronic acids to glucose.

[0089] The coupling moiety is a linking group as defined previously with the proviso that the linking group is optionally further substituted with a boronic acid, a polymerizable group, an additional coupling group, or is a segment in a polymer chain in which the viologen is a chain unit, a pendant group, or any combination thereof.

**Immobilizing Means**

[0090] In some embodiments, for use *in vitro* not involving a moving stream, the sensing components are used as individual (discrete) components. The dye and quencher are mixed together in liquid solution, analyte is added, the change in fluorescence intensity is measured, and the components are discarded. Polymeric matrices that can be used to trap the
sensing components to prevent leaching need not be present. Optionally, the sensing components are immobilized which allows their use to measure analytes in a moving stream.

For in vivo applications, the sensor is used in a moving stream of physiological fluid which contains one or more polyhydroxyl organic compounds or is implanted in tissue such as muscle which contains said compounds. Therefore, it is preferred that none of the sensing moieties escape from the sensor assembly. Thus, for use in vivo, the sensing components are preferably part of an organic polymer sensing assembly. Soluble dyes and quenchers can be confined by a semi-permeable membrane that allows passage of the analyte but blocks passage of the sensing moieties. This can be realized by using as sensing moieties soluble molecules that are substantially larger than the analyte molecules (molecular weight of at least twice that of the analyte or greater than 1000 preferably greater than 5000); and employing a selective semipermeable membrane such as a dialysis or an ultrafiltration membrane with a specific molecular weight cutoff between the two so that the sensing moieties are quantitatively retained.

Preferably the sensing moieties are immobilized in an insoluble polymer matrix, which is freely permeable to glucose. The polymer matrix is comprised of organic, inorganic or combinations of polymers thereof. The matrix may be composed of biocompatible materials. Alternatively, the matrix is coated with a second biocompatible polymer that is permeable to the analytes of interest.

The function of the polymer matrix is to hold together and immobilize the fluorophore and quencher moieties while at the same time allowing contact with the analyte, and binding of the analyte to the boronic acid. To achieve this effect, the matrix must be insoluble in the medium, and in close association with it by establishing a high surface area interface between matrix and analyte solution. For example, an ultra-thin film or microporous support matrix is used. Alternatively, the matrix is swellable in the analyte solution, e.g. a hydrogel matrix is used for aqueous systems. In some instances, the sensing polymers are bonded to a surface such as the surface of a light conduit, or impregnated in a microporous membrane. In all cases, the matrix must not interfere with transport of the analyte to the binding sites so that equilibrium can be established between the two phases. Techniques for preparing ultra-thin films, microporous polymers, microporous sol-gels, and
hydrogels are established in the art. All useful matrices are defined as being analyte permeable.

[0094] Hydrogel polymers are used in some embodiments. The term, hydrogel, as used herein refers to a polymer that swells substantially, but does not dissolve in water. Such hydrogels may be linear, branched, or network polymers, or polyelectrolyte complexes, with the proviso that they contain no soluble or leachable fractions. Typically, hydrogel networks are prepared by a crosslinking step, which is performed on water-soluble polymers so that they swell but do not dissolve in aqueous media. Alternatively, the hydrogel polymers are prepared by copolymerizing a mixture of hydrophilic and crosslinking monomers to obtain a water swellable network polymer. Such polymers are formed either by addition or condensation polymerization, or by combination process. In these cases, the sensing moieties are incorporated into the polymer by copolymerization using monomeric derivatives in combination with network-forming monomers. Alternatively, reactive moieties are coupled to an already prepared matrix using a post polymerization reaction. Said sensing moieties are units in the polymer chain or pendant groups attached to the chain.

[0095] The hydrogels useful in this invention are also monolithic polymers, such as a single network to which both dye and quencher are covalently bonded, or multi-component hydrogels. Multi-component hydrogels include interpenetrating networks, polyelectrolyte complexes, and various other blends of two or more polymers to obtain a water swellable composite, which includes dispersions of a second polymer in a hydrogel matrix and alternating microlayer assemblies.

[0096] Monolithic hydrogels are typically formed by free radical copolymerization of a mixture of hydrophilic monomers, including but not limited to HEMA, PEGMA, methacrylic acid, hydroxyethyl acrylate, N-vinyl pyrrolidone, acrylamide, N,N'-dimethyl acrylamide, and the like; ionic monomers include methacryloylammonopropyl trimethylammonium chloride, diallyl dimethyl ammonium, chloride, vinyl benzyl trimethyl ammonium chloride, sodium sulfopropyl methacrylate, and the like; crosslinkers include ethylene dimethacrylate, PEGDMA, trimethylolpropane triacrylate, and the like. The ratios of monomers are chosen to optimize network properties including permeability, swelling index, and gel strength using principles well established in the art. In one embodiment, the
dye moiety is derived from an ethylenically unsaturated derivative of a dye molecule, such as 8-acetoxy pyrene-1,3,6-N, N', N''-tris(methacrylamidopropylsulfonamide). the quencher moiety is derived from an ethylenically unsaturated viologen such as 4-N-(benzyl-3-boronic acid)-4'-N'-(benzyl-4ethenyl)-dipyridinium dihalide (m-SBBV) and the matrix is made from HEMA and PEGDMA. The concentration of dye is chosen to optimize emission intensity. The ratio of quencher to dye is adjusted to provide sufficient quenching to produce the desired measurable signal.

[0097] In some embodiments, a monolithic hydrogel is formed by a condensation polymerization. For example, acetoxy pyrene trisulfonyl chloride is reacted with an excess of PEG diamine to obtain a tris-(amino PEG) adduct dissolved in the unreacted diamine. A solution of excess trimesoyl chloride and an acid acceptor is reacted with 4-N-(benzyl-3-boronic acid)-4'-N'-(2 hydroxyethyl) bipyridinium dihalide to obtain an acid chloride functional ester of the viologen. The two reactive mixtures are brought into contact with each other and allowed to react to form the hydrogel, e.g. by casting a thin film of one mixture and dipping it into the other.

[0098] In other embodiments, multi-component hydrogels wherein the dye is incorporated in one component and the quencher in another are preferred for making the sensor of this invention. Further, these systems are optionally molecularly imprinted to enhance interaction between components and to provide selectivity for glucose over other polyhydroxy analytes. Preferably, the multicomponent system is an interpenetrating polymer network (IPN) or a semi-interpenetrating polymer network (semi-IPN).

[0099] The IPN polymers are typically made by sequential polymerization. First, a network comprising the quencher is formed. The network is then swollen with a mixture of monomers including the dye monomer and a second polymerization is carried out to obtain the IPN hydrogel.

[0100] The semi-IPN hydrogel is formed by dissolving a soluble polymer containing dye moieties in a mixture of monomers including a quencher monomer and polymerizing the mixture. In some embodiments, the sensing moieties are immobilized by an insoluble polymer matrix which is freely permeable to polyhydroxyl compounds. Additional details on hydrogel systems have been disclosed in US Patent Publications Nos.
US2004/0028612, and 2006/0083688 which are hereby incorporated by reference in their entireties.

[0101] The polymer matrix is comprised of organic, inorganic or combinations of polymers thereof. The matrix may be composed of biocompatible materials. Alternatively, the matrix is coated with a second biocompatible polymer that is permeable to the analytes of interest. The function of the polymer matrix is to hold together and immobilize the fluorescent dye and quencher moieties while at the same time allowing contact with the analytes (e.g., polyhydroxyl compounds, H⁻ and OH⁻), and binding of the polyhydroxyl compounds to the boronic acid. Therefore, the matrix is insoluble in the medium and in close association with it by establishing a high surface area interface between matrix and analyte solution. The matrix also does not interfere with transport of the analyte to the binding sites so that equilibrium can be established between the two phases. In one embodiment, an ultra-thin film or microporous support matrix may be used. In another embodiment, the matrix that is swellable in the analyte solution (e.g. a hydrogel matrix) can be used for aqueous systems. In some embodiments, the sensing polymers are bonded to a surface such as the surface of a light conduit, or impregnated in a microporous membrane. Techniques for preparing ultra-thin films, microporous polymers, microporous sol-gels, and hydrogels have been established in the prior art.

[0102] In one preferred embodiment, the boronic acid substituted viologen may be covalently bonded to a fluorescent dye. The adduct may be a polymerizable compound or a unit in a polymer. One such adduct for example may be prepared by first forming an unsymmetrical viologen from 4,4'-dipyridyl by attaching a benzyl-3-boronic acid group to one nitrogen and an aminoethyl group to the other nitrogen atom. The viologen is condensed sequentially first with 8-acetoxy-pyrene-1,3,6-trisulfonyl chloride in a 1:1 mole ratio followed by reaction with excess PEG diamine to obtain a prepolymer mixture. An acid acceptor is included in both steps to scavange the byproduct acid. The prepolymer mixture is crosslinked by reaction with a polyisocyanate to obtain a hydrogel. The product is treated with base to remove the acetoxy blocking group. Incomplete reaction products and unreacted starting materials are leached out of the hydrogel by exhaustive extraction with deionized
water before further use. The product is responsive to glucose when used as the sensing component as described herein.

[0103] Alternatively, such adducts are ethylenically unsaturated monomer derivatives. For example, dimethyl bis-bromomethyl benzene boronate is reacted with excess 4,4'-dipyridyl to form a half viologen adduct. After removing the excess dipyridyl, the adduct is further reacted with an excess of bromoethylamine hydrochloride to form the bis-viologen adduct. This adduct is coupled to a pyranine dye by reaction with the 8-acetoxypyrene-tris sulfonyl chloride in a 1:1 mole ratio in the presence of an acid acceptor followed by reaction with excess aminopropylmethacrylamide. Finally, any residual amino groups may be reacted with methacrylol chloride. After purification, the dye/viologen monomer may be copolymerized with HEMA and PEGDMA to obtain a hydrogel.

Ratiometric pH Sensing

[0104] Ratiometric pH sensing is known. See e.g., US Pat. Publication Nos. 2006/0105174; 2005/0090014; incorporated herein in their entirety by reference. Given an indicator system comprising a fluorophore (e.g., a fluorescent indicator dye) that exists in two forms (an acid form and a base form) the ratio of the emission intensity at the two wavelengths can be used to measure pH independent of the fluorophore concentration. The fluorescent indicator dyes suitable for ratiometric pH sensing may be: (1) dyes that exhibit dual excitation wavelengths (corresponding to acid and conjugate base forms) and single emission wavelengths (e.g., HPTS dyes); (2) single excitation wavelengths and dual emission wavelengths (acid and base forms); or (3) dual excitation - dual emission dyes. Some dyes, such as the SNARF or SNAFL dyes may have both dual-emission and dual-excitation properties. However a dual-dual dye, e.g., SNARF can be used as a single-dual or a dual-single.

using a dual-emission sensitive dye" Analytical Chemistry 69: 863-867. The extensive photobleaching observed for these dyes may be accounted for by the ratiometric approach, but it would still limit the useful lifetime of the sensor.


[0107] The presence of the three strongly anionic sulphonic acid groups allows for HPTS to be immobilized by ionic binding to cationic supports. To date, covalent attachment of HPTS has been via sulfonamide coupling (U.S. Pat. No. 4,798,738). While effective in immobilizing the dye and preserving pH sensitivity, polymer substrates are limited to those that contain primary amines. In addition, amine groups which remain on the substrate after coupling will affect the local pH inside the polymer matrix. The dye has been covalently attached to controlled pore glass (Offenbacher, H., O. S. Wolfbeis, et al. (1986). "Fluorescence optical sensors for continuous determination of near-neutral pH values." Sensor Actuator 9: 73-84) and aminoethyl cellulose (Schulman, S. G., S. Chen, et al. (1995).

[0108] For example U.S. Pat. No. 5,114,676 (incorporated by reference herein in its entirety) provides a pH sensor with a fluorescent indicator which may be covalently attached to a particle or to a microcrystalline cellulose fiber. The sensor comprises an optically transparent substrate, a thermoplastic layer and a hydrogel. Part of the particle with the indicator attached thereto is imbedded in a thermoplastic layer that is coated on the substrate and mechanically adhered using heat and pressure. The majority of the particle/indicator is imbedded within a hydrogel layer that is applied over the thermoplastic layer. The pH sensor is applied to the tip of an optical waveguide.

[0109] Furthermore, with the recent availability of low cost UV LEDs, the dye can be measured with relatively inexpensive instrumentation that combines UV and blue LEDs and a photodiode module. Such a setup has been described (Kostov, Y., P. Harms, et al. (2001). "Low-cost microbioreactor for high-throughput bioprocessing." Biotechnol Bioeng 72: 346-352) to detect the pH of a high throughput microbioreactor system via HPTS directly dissolved in the fermentation media.

[0110] In one embodiment of the present invention, the preferred sensing device comprises at least one light source, a detector, and a sensor comprising a fluorescent reporter dye system. In one embodiment, the fluorescent reporter dye system comprises a fluorescent dye operably coupled to an analyte-binding quencher. The dye may be covalently bound to the quencher or merely associated with the quencher. The dye and quencher are preferably
operably coupled, which means that in operation, the quencher is in close enough proximity to the dye to interact with and modulate its fluorescence. In one embodiment, the dye and quencher may be constrained together within an analyte-permeable hydrogel or other polymeric matrix. When excited by light of appropriate wavelength, the fluorescent dye emits light (e.g., fluoresces). The intensity of the light is dependent on the extent of quenching which varies with the amount of analyte binding. In other embodiments, the fluorescent dye and the quencher may be covalently attached to hydrogel or other polymeric matrix, instead of to one another.

[0111] In one embodiment, a separate pH indicator dye is combined with a different dye that is functionalized with an analyte-binding moiety, such that the two dye system are immobilized together (e.g., in a hydrogel) in the sensor.

[0112] Some fluorescent pH indicator molecules absorb light at a particular wavelength and emit light at a second, longer wavelength. Their pH indicating function typically involves protonation and deprotonation. This means that these fluorescent pH indicators include a hydrogen atom (proton, H+) which forms part of the molecule (is bound to the molecule) in one pH range, but within another pH range the proton is dissociated from the molecule. When the proton is disassociated from the molecule, the molecule takes on a negative charge, which is balanced by a positively-charged ion (e.g., Na+) in solution with the indicator. This arrangement is illustrated by Equation 1. \[ R - H \leftrightarrow R^- + H^+ \]

[0113] Where R represents a fluorescent molecule, it generally will exhibit fluorescence at a different wavelength (will be visible as a very different color) based upon whether it is in the R—H form or in the R− form. For most molecules represented by R, this change will occur generally quite abruptly within a very narrow pH range, allowing R to serve as a very simple and reliable pH indicator. When placed in solution, it will exhibit one very distinct color (a color associated with its R—H form), and another very distinct color associated with its R−.

[0114] For example, 8-Hydroxyl-1,3,6-pyrenetrisulphonate (HPTS) has been considered one of the best potential indicators for pH determination because of its excellent photo-stability, high quantum yield, dual excitation, large Stokes' shift and long fluorescence emission. A desirable feature of this indicator is that the acidic (associated HPTS form) and
basic (dissociated PTS) forms have different excitation wavelengths at 406 and 460 nm, with an isosbestic point at 418 nm, but exhibit a similar fluorescence emission maximum at 515 nm. The dual excitation and single emission make HPTS suitable for ratiometric detection of pH. The fluorescence intensity at 406 nm for the acid form decreases but the intensity at 460 nm for the base form increases as the pH is raised accompanying the conversion of the acidic into basic forms of the dye.

[0115] Due to the hydroxyl (-OH) group on dyes such as HPTS and its derivatives, these dyes are sensitive to the pH changes in the environment. The pH-dependent ionization of the hydroxyl group causes these pyranine derivatives to have a pH-dependent absorption spectra with different absorption maxima in its acidic form and basic form. The first absorption maximum is the first excitation wavelength and the second absorption maximum is the second excitation wavelength. The amounts of light absorbed by the fluorescent dye at the first excitation wavelength and the second excitation wavelength depend on or relate to the pH of the medium the fluorescent dye is in contact with. The amount of light emitted by the dye (e.g., the fluorescent emission) at the emission wavelength depends on the amount of light absorption when the dye is irradiated at the excitation wavelength. Since the absorption is affected by the pH of the medium, the fluorescent emission is also affected by the pH. This provides the basis for the pH determination while being able to measure the polyhydroxyl compound concentration.

[0116] In one preferred embodiment of the present invention, ratiometric pH sensing is accomplished using an optical sensor comprising at least one excitation light source operably coupled to the proximal end region of an optical fiber, wherein the fiber has disposed along its distal end region within the light path of the fiber, an indicator system configured to generate a detectable emission signal in response to the excitation light. Preferred embodiments of the sensor further comprise optical means for sending the emission signal to a detector. Such optical means are well known in the art, and may involve e.g., a mirror to return light, filters, lens, beam splitters, and optical fiber bundles and split configurations.

[0117] In preferred embodiments, the indicator system comprises a fluorophore that exhibits at least two different forms and a pH-dependent shift between these different
forms, wherein this shift can be detected as a change in the emission intensity at a single wavelength or at two different wavelengths. For example, one indicator system for ratiometric pH sensing comprises an fluorescent dye (e.g., HPTS) that absorbs light at two different wavelength maxima’s (\( \lambda_{\text{acid}} \) and \( \lambda_{\text{base}} \)) depending on whether the dye is in its acid or base forms, and it emits light at a single longer emission wavelength. More particularly, as pH is increased, HPTS shows an increase in absorbance corresponding to the \( \lambda_{\text{base}} \) and a decrease in absorbance corresponding to the \( \lambda_{\text{acid}} \). These changes are due to the pH-dependent ionization of the hydroxyl group. The emission spectrum for HPTS is independent of pH, with a peak emission wavelength of about 511 nm, but the intensity of the emitted light depends on the amount of light absorbed (which varies with pH and the excitation wavelength). So for example, if one excites HPTS at a given pH with light of a first wavelength (e.g., \( \lambda_{\text{acid}} \)), one can measure the emission intensity at the single emission wavelength; the intensity will depend on the form of the dye (i.e., degree of ionization - which depends on the pH). One can also excite at a second wavelength (e.g., \( \lambda_{\text{base}} \)) and measure the emission intensity at the same given pH. The ratio of the emission intensities relates to the pH and is independent on the amount of the dye as well as certain optical artifacts in the system. It is noted that any excitation wavelengths may be used for the ratiometric sensing, but the \( \lambda_{\text{acid}} \) and \( \lambda_{\text{base}} \) are preferred in accordance with one embodiment of the invention. The wavelength at which the absorption is the same for the acid and base forms of the dye is called the isobestic point—excitation at this wavelength (\( \lambda_{\text{iso}} \)) may also be used in ratiometric sensing in accordance with other preferred variations to the invention.

When a ratio of emission intensities (e.g., \( I_{\text{base}}/I_{\text{iso}} \) or \( I_{\text{base}}/I_{\text{acid}} \)) is plotted against pH, a standard or calibration curve is generated (See e.g., FIGS. 3, 5 and 9). The ratiometric method is similar regardless of whether the dye used is a dual exciter—single emitter (like HPTS), or a single exciter—dual emitter, or a dual exciter—dual emitter, as long as the dye undergoes a pH sensitive shift in form that yields a detectable change in spectral property.

**Optical Glucose Sensing**

[0118] Indicator systems comprising fluorescent dyes, including HPTS and its derivatives, have been used in analyte detection. See e.g., U.S. Pat. Nos. 6,653,141, 6,627,177, 5,512,246, 5,137,833, 6,800,451, 6,794,195, 6,804,544, 6,002,954, 6,319,540,
6,766,183, 5,503,770, and 5,763,238; and co-pending U.S. Patent Application Nos. 11/296,898 and 60/833,081: each of which is incorporated herein in its entirety by reference thereto. In particular, details related to some preferred fluorescent dyes, quenchers/analyte binding moieties, and methods for optically determining polyhydroxyl compound concentrations are disclosed in U.S. Pat. Nos. 6,653,141 and 6,627,177, and U.S. Patent Application Nos. 11/296,898 and 60/833,081.

**Device for Intravascular Determination of pH and Glucose**

[0119] In one embodiment, the method and sensor monitor the pH of the media and the concentration of analyte *in vitro*. In another embodiment, the method and sensor monitor pH and analyte concentration *in vivo*. In another embodiment, the measured pH value can also be used to more correctly determine glucose concentration *in vitro or in vivo*. Specifically, the simultaneous measurement of the pH value and the glucose concentration would enable real-time correction of the signal of glucose response. Although it will be appreciated that the device in accordance with some embodiments comprise a sensor that may be used only to determine pH or analyte (correction of which for pH may be done by conventional two-sensor technologies or by testing the blood pH *in vitro*).

[0120] One embodiment provides a device for determining pH and the concentration of a polyhydroxyl compound simultaneously, comprising a sensor comprising a fluorescent dye operably coupled to a quencher; means for delivering one or more excitation wavelengths to said sensor, and means for detecting fluorescence emission from said sensor.

[0121] Another embodiment provides a device for determining the pH and die polyhydroxyl compound concentration in a physiological fluid, comprising a water-insoluble polymer matrix, wherein said polymer matrix is permeable to polyhydroxyl compound; a fluorescent dye associated with said polymer matrix, wherein the fluorescent dye is configured to absorb light at a first excitation wavelength and a second excitation wavelength, and to emit light at an emission wavelength; a quencher comprising an aromatic boronic acid substituted viologen, adapted to reversibly bind an amount of polyhydroxyl compound dependent on the polyhydroxyl compound concentration, wherein said quencher is associated with said polymer matrix and operably coupled to the fluorescent dye, and wherein the quencher is configured to reduce the light intensity emitted by said fluorescent dye related
to the amount of bound polyhydroxyl compound; at least one excitation light source; and an emission light detector.

[0122] In one aspect, the present invention comprises a class of fluorescence quenching compounds that are responsive to the presence of polyhydroxyl compounds such as glucose in aqueous media at or near physiological pH. In other words, the quenching efficiency is controlled by the concentration of these compounds in the medium. Preferred quenchers comprise a viologen substituted with at least one boronic acid group wherein the adduct is immobilized in or covalently bonded to a polymer. The quencher, dye and polymer may also be covalently bonded to each other. In another aspect, the present invention comprises a class of fluorescent dyes which are susceptible to quenching by the viologen/boronic acid adduct.

[0123] The fluorescent dye and quencher are operably coupled to each other for polyhydroxyl compound sensing. The dye and quencher may be linked through a polymer backbone in some embodiments. In other embodiments, the dye and quencher could be in close proximity to each other for the quenching of the fluorescent dye to occur, thereby reducing the fluorescent emission of the dye. When polyhydroxyl compound (e.g., glucose) binds to the boronic acid to form boronate ester, the boronate ester interacts with the viologen and alters its quenching efficacy according to the extent of polyhydroxyl compound binding. As a result, the intensity of fluorescent emission increases as more polyhydroxyl compounds are bonded to the quenchers.

[0124] In one preferred embodiment, the device comprises an optical fiber comprising a cavity disposed therein and having immobilized within the cavity an indicator system as described above (e.g., a fluorophore operably coupled to a glucose binding moiety/quencher and an immobilizing polymeric matrix). The device further comprises a light source and a detector.

Methods for Simultaneous Determination of pH and Glucose

[0125] One embodiment provides a method for determining the pH and the polyhydroxyl compound concentration with one fluorescent dye, comprising providing a sensor comprising a fluorescent dye operably coupled to a quencher, contacting said sensor with a sample; irradiating said sensor at the first excitation wavelength; detecting a first
fluorescence emission of said sensor at an emission wavelength: irradiating said sensor at the
temperature wavelength; measuring a second fluorescence emission of said sensor at
at said emission wavelength; comparing the ratio of the first and second emissions with a pH
calibration curve to determine the pH of the sample: correlating the emission quenching with
a standard curve at the known pH to determine the polyhydroxyl compound concentration in
said sample. Of course other algorithms are known for ratiometric pH sensing and may be
used in accordance with embodiments of the present invention. A controller, such as a
computer or dedicated device, may be used in some embodiments to control the operations,
including application of the excitation light, monitoring of detector signals, determining
ratios, correlating ratios with calibration curves, correlating glucose signals with standard
curves, correcting for pH changes, running routine sensor calibration operations, prompting
operator actions, integrating user data input (e.g., finger stick glucose measurements) as
programmed to maintain accuracy, etc.

[0126] With respect to Figure I, a sensing device 100 in accordance with one
embodiment of the present invention comprises at least one light source 11 (e.g., an
excitation light source), a detector 15 (e.g., an emission light detector), and a sensor 13
comprising a fluorescent dye operably coupled to a quencher and an optional polymer matrix.
In some embodiments, the light source 11 may be adapted to selectivity deliver two or more
different wavelength for the excitations of fluorescent dyes. This type of light source may be
a tunable light source. In other embodiments, one or more light sources may be used in
conjunction with an optical filter 12 for attenuating the wavelengths. In other embodiments,
more than one light source 11 may be used to deliver different excitation wavelengths. Such
light source is also a means for delivering a first and a second excitation wavelengths to the
sensor.

[0127] The sensor 13 comprises a fluorescent dye that is sensitive to both the pH
and the polyhydroxyl compound (e.g., sugar or glucose) concentration of the medium when
the dye is operably coupled to a quencher. Such fluorescent dye exhibits a shift in excitation
wavelength maximum with a corresponding shift in pH of the local environment of the
fluorescent dye. As the pH of the local environment changes, the absorption at a first
excitation wavelength may increase, while the absorption at a second excitation wavelength
decreases, or vice versa. The change in absorption at a selected wavelength can affect the level of fluorescence emission, therefore ultimately permitting pH detection. The pH detection is independent of the concentration of the polyhydroxyl compound in the environment. A suitable fluorescent dye is also susceptible to quenching by molecules such as viologens. When the fluorescent dye is operably coupled to a quencher (e.g., a viologen), the fluorescence emission is attenuated. The quencher may have an aromatic boronic acid moiety that is capable of providing glucose recognition. The boronic acid reacts reversibly with glucose in aqueous media to form boronate ester, and the extent of such reaction is related to the glucose concentration in the medium. As more glucose is available to react with the quencher, the quencher’s ability to accept electron from the dye decreases. As a result, the attenuation of fluorescence emission by the quencher is dependent on the concentration of the polyhydroxyl compound (e.g., glucose) to be detected.

[0128] A detector 15 is used to detect the fluorescent emission and in preferred embodiments, may be linked to the electronic control 20 for analysis. Optical filters, e.g., 14, can be placed between the sensor 13 and the detector 15 for wavelength selection. Other optical components may also be utilized, e.g., mirrors, collimating and/or focusing lenses, beam splitters, etc. Optical fibers can be used to deliver selected wavelengths to the sensor and to deliver the fluorescence emission from the sensor to the detector. The light source and the detector may be controlled by electronic control 20 such as a computer.

[0129] One embodiment of this invention provides a method for measuring pH and polyhydroxy] compound concentration with a single fluorescent dye. Measurements can be carried out in vitro or in vivo. It may be necessary to calibrate the sensor prior to performing the first measurement. This may be done by first acquiring the absorbance spectra of the sensor at various pHs to determine the wavelengths where isobestic point and absorption maxima for acid and base forms occur and then acquiring the emission signals from at least two of these wavelengths at at least one known pH and glucose concentration.

[0130] For the pH and polyhydroxyl concentration measurements, the sensor 13 is first placed in contact with a sample. The sensor 13 is then irradiated at the first excitation wavelength followed by the second excitation wavelength. The first and second excitation wavelengths are typically chosen near the wavelength of the absorption maximum for the
acidic form of the fluorescent dye \( \lambda_{\text{acid}} \), the wavelength of the absorption maximum for the basic form of the fluorescent dye \( \lambda_{\text{base}} \), or the wavelength of the isobestic point \( \lambda_{\text{iso}} \), or other selected wavelength. The ratio of the emissions from the first and second excitation wavelengths are used to determine the sample pH. Either the first or second emission, once corrected for pH, can be used to determine the sample glucose concentration.

[0131] In variations to the sensing device shown in Figure 1, the detector may be a standard photodiode detector. There may be two diode detectors, one for a reference and one for the emission signal. Instead of diode detectors, the optical fiber carrying sensor output (fluorescent emission and/or reflected excitation light) may provide input directly to a spectrophotometer or microspectrometer. In a preferred embodiment, the detector comprises a microspectrometer such as the UV/VIS Microspectrometer Module manufactured by Boehringer Ingelheim.

[0132] Figure 2 shows one embodiment of an optical system that may be used in accordance with preferred aspects of the present invention. With reference to FIG. 2, certain embodiments comprise at least two light sources, 301A and 301B. The light sources generate excitation light that may be transmitted (as illustrated) through collimator lenses 302A and 302B. In certain embodiments, the resulting light from collimator lenses may be transmitted (as illustrated) to interference filters 303A and 303B. In certain embodiments, the resulting light from interference filters may be focused (as illustrated) by focusing lenses 304A and 304B into fiber optic lines 305A and 305B. In certain embodiments, fiber optic lines merge into a single fiber 306 that is continuous with the sensor 307, having the imbedded indicator system 307A. The cross-sections of the fibers may vary (as illustrated) from a bundle of fibers surrounding a central optical fiber 306A to a single fiber 307A.

[0133] In certain embodiments (as illustrated), the emission light signals generated by the indicator system 307A as well as the excitation light signals are reflected by mirror 308 and transmitted back out of the sensor into the fiber optic outlet lines 309 and 309A. In the illustrated system, the outlet lines are augmented by including two interference filters 312A, 312B and two detectors 313A, 313B. In preferred embodiments, the interference filter 312A is configured to block the excitation light and allow the emission light to pass to detector 313A where the emission light is detected. In certain embodiments,
the signal produced by the detector 313A is amplified by the amplifier 314A and converted into a digital signal by analog-to-digital converter 315A and transmitted to computer 316. In certain embodiments, the interference filter 312B is configured to block the emission light and allow the excitation lights to pass to detector 313B where the excitation light is measured. In certain embodiments, the signal produced by the detector 313B is amplified by the amplifier 314B and converted into a digital signal by analog-to-digital converter 315B and transmitted to computer 316. Ratiometric calculations may be employed to substantially eliminate or reduce non-glucose related factors affecting the intensity of the emission light; these methods are disclosed in detail in co-pending US Provisional Application No. 60/888,477, entitled "Optical systems and methods for ratiometric measurement of glucose using intravascular fluorophore sensors," filed herewith on the same day, and incorporated herein in its entirety by reference thereto.

EXAMPLES

[0134] Example 1 - Figure 3 shows an example of the excitation/absorption spectrum of a fluorescent dye, in this case HPTS. From the absorption spectra of the fluorescent dye acquired at different pHs, $\lambda_{\text{acid}}, \lambda_{\text{base}}$ and $\lambda_{\text{iso}}$ can be determined. At a lower pH (e.g., more acidic condition), the peak at around 405 nm is higher than the peak at around 460 nm, and is therefore the absorption maximum for the acidic form of the fluorescent dye. At a higher pH (e.g., more basic condition), the peak at round 460 nm is higher than the peak at around 405 nm, therefore is the absorption maximum for the basic form of the fluorescent dye. The $\lambda_{\text{iso}}$ would be the wavelength where the absorption is independent of the pH, and it would be, for example, around 422 nm for HPTS.

[0135] The first fluorescence emission intensity ($I_x$, which could be $I_{\text{acid}}, I_{\text{base}}$ or $I_{\text{iso}}$) at a emission wavelength, resulting from the irradiation at the first excitation wavelength (e.g., $\lambda_{\text{acid}}, \lambda_{\text{base}}$ or $\lambda_{\text{iso}}$), is then measured by the detector and the result is stored in the electronic control. Then the sensor is again irradiated at the second excitation wavelength. The second excitation wavelength is different from the first excitation wavelength and can also be selected from $\lambda_{\text{acid}}, \lambda_{\text{base}}$ or $\lambda_{\text{iso}}$. The detector will then detect/measure the second fluorescence emission intensity ($I_y$, which could be $I_{\text{acid}}, I_{\text{base}}$ or $I_{\text{iso}}$) resulting from the irradiation at the second excitation wavelength (e.g., $\lambda_{\text{acid}}, \lambda_{\text{base}}$ or $\lambda_{\text{iso}}$). The ratio of the first
and the second fluorescence emissions ($I_x/I_y$) can then be computed. Since the $I_x/I_y$ is independent from the polyhydroxyl concentration, a pH standard curve ($I_x/I_y$ vs. pH) can be plotted without considering the effect of polyhydroxyl concentration.

Example 2 (HPTS/MABP4) - Figure 4 shows independence of ratiometric pH sensing using HPTS/MABP4 using the $I_{(base)}/I_{(iso)}$ ratio from glucose concentration. The structure of MABP4 is:

![MABP4 Structure](image)

The data are plotted as a ratio of the fluorescence emission for corresponding to excitation at 454 nm (base) and 422 nm (isobestic point) vs. pH in various glucose concentrations. The changes in glucose concentrations have no discernible effects on the value of $I_{base}/I_{iso}$ at each specific pH. Thus the pH of the sample can be measured using a standard curve of $I_x/I_y$ vs. pH, regardless of the polyhydroxyl compound concentration in the sample. By correlating or comparing the measured $I_x/I_y$ to the standard curve, one may determine the pH of the sample being measured.

Figure 5 shows glucose response curves for HPTS/MABP4 excited at 422 nm (isobestic point) at different pHs. By plotting the ratio of $I_x/I_y$ at various glucose levels (1) to $I_x/I_y$ at zero glucose concentration (I0) vs. glucose concentration, a standard polyhydroxyl response curve can be used to determine the glucose concentration in a sample from measured I/I0 values. However, since I/I0 value is dependent on the pH of the sample, the standard glucose response curve can be affected by the different pH. To circumvent this, several standard glucose response curves at different pHs within the physiological range can
be plotted and available for selection by either the electronic control or the operator of the sensor device. When the $I_x/I_y$ measurement of the sample is available, the electronic control or the operator would know the pH of the sample from the standard $I_x/I_y$ vs. pH curve, and the correct standard polyhydroxyl response curve (e.g., glucose response curve) may be used for determining the accurate glucose concentration. Although the examples shown in the figures concern determination of glucose concentration, the application of the method and device of the present invention is not limited to detecting glucose concentration. Since the fluorescent system responds to polyhydroxyl compounds the same way it responds to glucose, the sensor device can be used to detect any polyhydroxyl compound concentration and the pH at the same time.

[0139] Example 3 (SNARF-1) - Figure 6 shows the absorption spectra of SNARF-1 at different pHs in solution. SNARF is a tradename for a class of commercial dyes from Molecular Probes, Inc. These experiments were carried out using SNARF-1. Figures 7 and 8 show glucose response curves for SNARF-1/3,3'-oBBV in solution at different pHs determined at 514 nm excitation/587 nm emission (Figure 7), or at 514 nm excitation/625 nm emission (Figure 8). Figure 9 shows ratiometric sensing of pH at different glucose concentrations with SNARF-1/3,3'-oBBV in solution using the $I_{(base)}/I_{(acid)}$ ratio determined at a single excitation wavelength of 514 nm and emission wavelengths of 587 and 625 nm. Thus, the dual-dual dye SNARF-1 may be used operably coupled to the quencher 3,3'-oBBV (in solution) as a single exciter-dual emitter fluorophore to determine both pH ratiometrically and glucose.

[0140] Example 4 (HPTS-triLysMA/3,3'-oBBV/DMMA) - Figure 10 shows the glucose response of HPTS-triLysMA/3,3'-oBBV/DMMA indicator system at different pHs. Figure 11 shows ratiometric sensing of pH at different glucose concentrations with the HPTS-triLysMA/3,3'-oBBV/DMMA indicator system, using the $I_{(base)}/I_{(acid)}$ ratio. It can be seen that this indicator system provides a linear pH curve over the physiologic pH range.

[0141] Example 5 (HFTS-tri CysMA/3,3'-oBBV/DMMA) - Figure 12 shows ratiometric sensing of pH at different glucose concentrations with the HFTS-triCysMA/3,3'-oBBV/DMMA indicator system, using the $I_{(base)}/I_{(acid)}$ ratio. It can be seen that this indicator system provides a linear pH curve over the physiologic pH range. For this example, the
indicator system was immobilized in a hydrogel embedded at the end of an optical fiber. The acid and base emission signals were measured using a hand-held detector.

[0142] **Lifetime Chemistry**

[0143] In another preferred embodiment, glucose concentrations can be determined by exploiting the phenomena of fluorescence resonance energy transfer (FRET). FRET is the transfer of energy from a donor fluorophore to an acceptor molecule. FRET occurs when the donor fluorophore, which fluoresces at a wavelength absorbed at least in part by the acceptor molecule, is in close proximity to the acceptor such that the donor fluorophore can transfer energy to the acceptor through molecular interactions. The fluorescence lifetime of the fluorophore, where the fluorescence lifetime is the time the fluorophore remains in the excited state, is altered by FRET. Thus, measuring the fluorescence lifetime of the fluorophore allows one to determine whether the fluorophore is bound to the acceptor.

[0144] Lifetime can be measured by using a time-domain method where the fluorophore is excited by a brief pulse of excitation light and the fluorescence intensity is measured over time. The excitation pulse can be a pulse from a laser with a duration in the picoseconds range up to a duration of about a few nanoseconds. In other embodiments, the pulse duration can be greater than about a few nanoseconds. The fluorescence intensity of the fluorophore as a function of time is given by the equation:

\[ I(t) = I_0 \times \exp(-t/\tau) \]  

**Equation 1**

[0145] \( I(t) \) is the fluorescence intensity at time (t), \( I_0 \) is the initial intensity after excitation and \( \tau \) is the fluorescence lifetime which is defined as the time required for \( I(t) \) to decay to \( I_0/e \). **Equation 1** is applicable to a fluorophore with a single exponential decay of fluorescence and a lifetime that is substantially longer than the excitation pulse. Figure 13 shows a graph of the decay of the fluorescent emission 400 over time after a pulse of excitation light 402. The time it takes the initial intensity, \( I_0 \), to drop to \( I_0/e \) is equal to the lifetime, \( \tau \).
An alternative method of measuring lifetime is by a frequency-domain method where the fluorophore is excited by a frequency modulated excitation light. The fluorescence lifetime, $\tau$, can be determined by measuring the phase shift of the emission from the fluorophore relative to the excitation light, or by measuring the modulation ratio, using the following equations:

$$\tau_\phi = \omega^{-1} \cdot \tan(\phi) \quad \text{Equation 2}$$

$$\omega = 2\pi f \quad \text{Equation 3}$$

$$\tau_M = \omega^{-1} \cdot (M^{-2} - 1)^{\frac{1}{2}} \quad \text{Equation 4}$$

$$M = \frac{(AC/DC)_EM}{(AC/DC)_EM} \quad \text{Equation 5}$$

$\tau_\phi$ is the lifetime determined by measuring the phase shift, $\phi$. $\omega$ is the angular frequency of the frequency modulated excitation light and $f$ is the linear frequency. $\tau_M$ is the lifetime determined by measuring the modulation ratio, $M$. AC is the magnitude of the alternating portion of the signal, or the amplitude of the wave, while DC is the amplitude of the DC portion of the signal. EM refers to the emission signal, and EX refers to the excitation signal. Figure 14 is a graph showing the relationship between the emission signal 500 and the excitation signal 502 and the variables described in Equations 2-5.

Preferred binding assay configurations for use in the sensor include a reversible competitive, reagent limited, binding assay, the components of which include an analyte analog and an analyte binding agent capable of reversibly binding both the analyte of interest and the analyte analog. The analyte of interest and the analyte analog compete for binding to the same binding site on the analyte binding agent. Such competitive binding assay configurations are well known in the art of clinical diagnostics and are described, by way of example, in The Immunoassay Handbook, ed. David Wild, Macmillan Press 1994. Suitable analyte binding agents for use in the assay would include antibodies or antibody fragments which retain an analyte binding site (e.g. Fab fragments), lectins (e.g. concanavalin A), hormone receptors, drug receptors, aptamers and molecularly-imprinted polymers. Preferably the analyte analog should be a substance of higher molecular weight than the analyte such
that it cannot freely diffuse out of the sensor. For example, an assay for glucose might employ a high molecular weight glucose polymer such as dextran as the analyte analog.

[0148] Suitable optical signals which can be used as an assay readout in accordance with the invention include any optical signal which can be generated by a proximity assay, such as those generated by fluorescence resonance energy transfer, fluorescence polarisation, fluorescence quenching, phosphorescence technique, luminescence enhancement, luminescence quenching, diffraction or plasmon resonance.

[0149] In some preferred embodiments of the sensor of the invention incorporates a competitive, reagent limited binding assay which generates an optical readout using the technique of fluorescence resonance energy transfer. In this assay format the analyte analog is labelled with a first chromophore and the analyte binding agent is labelled with a second chromophore. One of the first and second chromophores acts as a donor chromophore and the other acts as an acceptor chromophore. It is an important feature of the assay that the fluorescence emission spectrum of the donor chromophore overlaps with the absorption spectrum of the acceptor chromophore, such that when the donor and acceptor chromophores are brought into close proximity by the binding agent a proportion of the energy which normally would produce fluorescence emitted by the donor chromophore (following irradiation with incident radiation of a wavelength absorbed by the donor chromophore) will be non radiatively transferred to the adjacent acceptor chromophore, a process known in the art as FRET, with the result that a proportion of the fluorescent signal emitted by the donor chromophore is quenched and, in some instances, that the acceptor chromophore emits fluorescence. Fluorescence resonance energy transfer will generally only occur when the donor and acceptor chromophores are brought into close proximity by the binding of analyte analog to analyte binding agent. Thus, in the presence of analyte, which competes with the analyte analog for binding to the analyte binding agent, the amount of quenching is reduced (resulting in a measurable increase in the intensity of the fluorescent signal emitted by the donor chromophore or a fall in the intensity of the signal emitted by the acceptor chromophore) as labelled analyte analog is displaced from binding to the analyte binding agent. The intensity or lifetime of the fluorescent signal emitted from the donor chromophore thus correlates with the concentration of analyte in the fluid bathing the sensor.
An additional advantageous feature of the fluorescence resonance energy transfer assay format arises from the fact that any fluorescent signal emitted by the acceptor chromophore following excitation with a beam of incident radiation at a wavelength within the absorption spectrum of the acceptor chromophore is unaffected by the fluorescence resonance energy transfer process. It is therefore possible to use the intensity of the fluorescent signal emitted by the acceptor chromophore as an internal reference signal, for example in continuous calibration of the sensor or to monitor the extent to which the sensor has degraded and thus indicate the need to replace the sensor. As the sensor degrades, the amount of acceptor chromophore present in the sensor will decrease and hence the intensity of fluorescent signal detected upon excitation of the acceptor chromophore will also decrease. The fall of this signal below an acceptable baseline level would indicate the need to implant or inject a fresh sensor. Competitive binding assays using the fluorescence resonance energy transfer technique which are capable of being adapted for use in the sensor of the invention are known in the art. U.S. Pat. No. 3,996,345 describes immunoassays employing antibodies and fluorescence resonance energy transfer between a fluorescer-quencher chromophoric pair. Meadows and Schultz (Anal. Chim. Acta (1993 250: pp21-30) describe a homogeneous assay method for the measurement of glucose based on fluorescence resonance energy transfer between a labelled glucose analog (FITC labelled dextran) and a labelled glucose binding agent (rhodamine labelled concanavalin A). In all of these configurations the acceptor and donor chromophores/quenchers can be linked to either the binding agent or the analyte analog.

Fluorescence lifetime or fluorescence intensity measurements may be made. As described in Lakowitz et al, Analytica Chimica Acta, 271, (1993), 155-164, fluorescence lifetime may be measured by phase modulation techniques.

In some preferred embodiments as shown in Figures 15A, 15B and 15C, a competitive binding system to measure glucose using FRET comprises a glucose binding molecule 600 linked to a donor fluorophore 602 and a glucose analog 604 linked to an acceptor molecule 606. The glucose binding molecule 600 is capable of binding with both glucose 608 and the glucose analog 604. As shown in Figure 15A, when the glucose analog 604 is bound to the glucose binding molecule 600, the fluorescent emission 500 from the
fluorophore 602 is reduced in magnitude and shifted in phase and lifetime by FRET 610 because the fluorophore 502 is in close proximity to the acceptor 606. In other embodiments, the fluorophore 602 is linked to the glucose analog 604 and the acceptor 606 is linked to the glucose binding molecule 600.

[0153] As shown in Figure 15B, glucose 608 competes with the glucose analog 604 for the binding site on the glucose binding molecule 600. As shown in Figure 15C, the glucose molecule 608 can displace the glucose analog 604 from the glucose binding molecule 600 so that the acceptor 606 does not alter the emission lifetime S00 of the fluorophore 602 via FRET 610.

[0154] In a system where there are a certain concentration of glucose binding molecules, glucose analogs and glucose molecules, an equilibrium will exist between the number of bound glucose molecules to the number of bound glucose analogs. A change in the number of glucose molecules in the system, changes the equilibrium between bound glucose molecules to bound glucose analogs. This in turn changes the mean lifetime of the fluorophore emission.

[0155] In some preferred embodiments, the system is excited by a frequency modulated excitation light less than approximately 1 MHz, between approximately 1 to 200 MHz, or greater than approximately 200 MHz. In some embodiments, the frequency is approximately 0.05, 0.1, 1, 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190 or 200 MHz. By measuring the degree of the phase shift of the system, an average FRET induced phase shift for the system can be determined which corresponds to an average lifetime value for the system as defined by Equations 2 and 3 described above. Both the phase shift and the lifetime values can be correlated to the glucose concentration. The magnitude of the phase shift is independent of the amplitude of the emission.

[0156] In other preferred embodiments, the system is excited by a pulse and the decay of the fluorescence is measured over time. The lifetime can be determined using Equation 1 described above, and glucose concentration can be correlated to the lifetime value.

[0157] In preferred embodiments, the glucose binding molecule with a donor fluorophore and the glucose analog with an acceptor can be substantially immobilized in the
hydrogel described above such that diffusion of the glucose binding molecule and the glucose analog out of the hydrogel is substantially reduced. In addition, the sensor is configured to provide excitation light at a wavelength absorbed by the donor fluorophore as described above. In some embodiments, the excitation light is provided as a short pulse from a laser or a light emitting diode (LED). In other embodiments, the excitation light is frequency modulated. In some embodiments, the frequency modulated excitation light is provided by a laser. In some embodiments the frequency modulated excitation light is provided by a LED. The sensor also has a detector that detects the amplitude of the emission over time and/or the phase shift of the emission and/or the amplitudes of the AC and DC portions of the emission and excitation light. The detector can be a photodetector or multiple photodetectors. The excitation and emission light can be transmitted throughout the sensor via optical fibers.

[0158] In some embodiments, the sensor can be introduced into a patient's blood vessel, such as a vein, artery or capillary, for measuring the intravascular concentration of an analyte in the patient's blood. In some embodiments, the chemistry used to measure the concentration of the analyte is based on a correlation of fluorescence intensity of a fluorophore to analyte concentration, as described above in more detail. In some embodiments, the chemistry used to measure the concentration of the analyte is based on a correlation of fluorescence lifetime of a fluorophore to analyte concentration, as described above in more detail. In some embodiments, the sensor is used to measure the concentration of glucose.

[0159] While a number of preferred embodiments of the invention and variations thereof have been described in detail, other modifications and methods of using and medical applications for the same will be apparent to those of skill in the art. Accordingly, it should be understood that various applications, modifications, and substitutions may be made of equivalents without departing from the spirit of the invention or the scope of the claims.
WHAT IS CLAIMED IS:

1. An intravascular sensor for determining an analyte concentration in blood comprising:
   an optical fiber comprising a sensor chemistry portion capable of insertion into a blood vessel, the sensor chemistry portion comprising:
   an analyte binding molecule capable of binding an analyte, and
   a fluorophore associated with the analyte binding molecule, the fluorophore having a first fluorescence when the analyte binding molecule is not bound to the analyte and a second fluorescence when the analyte binding molecule is bound to the analyte;
   a light source; and
   a detector.

2. The sensor of Claim 1, wherein the first fluorescence is a first fluorescence intensity and the second fluorescence is a second fluorescence intensity.

3. The sensor of Claim 1, wherein the first fluorescence is a first fluorescence lifetime and the second fluorescence is a second fluorescence lifetime.

4. The sensor of Claim 1, wherein the analyte is glucose.

5. The sensor of Claim 1, further comprising an optical fiber.

6. The sensor of Claim 1, wherein the light source is a laser.

7. The sensor of Claim 1, wherein the light source is a light emitting diode.

8. The sensor of Claim 1, wherein the sensor chemistry portion further comprises a hydrogel that substantially immobilizes the analyte binding molecule and the fluorophore and is permeable to the analyte.

9. The sensor of Claim 1, wherein the sensor chemistry portion further comprises a membrane wherein the analyte binding molecule and the fluorophore are substantially retained within a volume at least partially enclosed by the membrane.

10. An intravascular sensor for determining an analyte concentration in blood comprising:
   a sensor chemistry portion capable of insertion into a blood vessel, the sensor chemistry portion comprising:
an analyte binding molecule capable of binding an analyte; and
a fluorophore associated with the analyte binding molecule, the fluorophore having a first fluorescence intensity when the analyte binding molecule is not bound to the analyte and a second fluorescence intensity when the analyte binding molecule is bound to the analyte;
a light source; and
a detector.
11. An intravascular sensor for determining an analyte concentration in blood comprising:
a sensor chemistry portion capable of insertion into a blood vessel, the sensor chemistry portion comprising:
an analyte binding molecule capable of binding an analyte; and
a fluorophore associated with the analyte binding molecule, the fluorophore having a first fluorescence lifetime when the analyte binding molecule is not bound to the analyte and a second fluorescence lifetime when the analyte binding molecule is bound to the analyte;
a light source; and
a detector.
12. A method for determining the glucose concentration in blood comprising:
providing the sensor of any of Claims 1-11;
inserting the sensor into a blood vessel;
irradiating the fluorophore with an excitation signal;
detecting a fluorescence emission from the fluorophore; and
determining the concentration of glucose.
13. The method of Claim 12, wherein the excitation signal is a pulse of light.
14. The method of Claim 13, further comprising measuring the decay of the fluorescence emission over time; and
calculating a fluorescence lifetime.
15. The method of Claim 12, wherein the excitation signal is frequency modulated.
16. The method of Claim 15, further comprising measuring the phase shift between the fluorescence emission and the excitation signal.

17. The method of Claim 16, further comprising calculating a fluorescence lifetime.

18. The method of Claim 15, further comprising measuring a modulation ratio; and calculating a fluorescence lifetime.

19. The method of Claim 12, further comprising measuring the fluorescence intensity of the fluorescence emission.

20. An intravascular sensor for determining an analyte concentration in blood, comprising an optical fiber comprising a sensor chemistry portion disposed along a distal region of the optical fiber, wherein the sensor chemistry portion is sized and physiologically compatible with residing in a blood vessel, wherein the sensor chemistry portion is selected from equilibrium fluorescence chemistry or lifetime chemistry.
FIG 1

11 Light Source
12 Optical Filter
13 Sensor
14 Optical Filter
15 Detector
20 Electronic Control
FIG. 4
Glucose Response @ Several pH's (Excitation at the Dye's Isobestic Point)

FIG. 5
Glucose Response of SNARF/3,3'-dBBV at Different pHs

\[ \text{Ex}=514 \text{nm}, \text{Em}=625 \text{nm} \]
Ratiometric Sensing of pH at Different Glucose Levels using SNARF/3,3'-oBBV

**FIG. 9**
FIG. 11

Glucose Response of HPMS-\((\text{r})\text{lysMA/3,3'~oBBV at different pHs}

5 6 7 8

5 6 7 8

pH

50 mg/dL
100 mg/dL
200 mg/dL
400 mg/dL
Base Emission Intensity/Acid Emission Intensity vs pH at Four Glucose Concentrations (Dye=HPTS Cystine, Quencher=3-3'-obvb, Polymer+DMMA)
(Data taken with chemistry immobilized on the end of an optical fiber)

FIG. 12
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

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

INV. A61B5/00

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

A61B

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 practical, search terms used)

EPO-Internal, WPI Data

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>
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Further documents are listed in the continuation of Box C.

* * * Special categories of cited documents :

'A' document defining the general state of the art which is not considered to be of particular relevance

'B' earlier document but published on or after the international filing date

'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

'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

'T' 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

'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

'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

'S' document member of the same patent family

Date of the actual completion of the international search

26 August 2008

Date of mailing of the international search report

03/09/2008

Name and mailing address of the ISA:
European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk
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Authorized officer

Abraham, Volkhard
<table>
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<tr>
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<th>Citation of document with indication where appropriate of the relevant passages</th>
<th>Relevant to claim No</th>
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INTERNATIONAL SEARCH REPORT

Box No. II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. [✓] Claims Nos.: 12-19
   because they relate to subject matter not required to be searched by this Authority, namely:
   Rule 39.1(iv) PCT - Method for treatment of the human or animal body by surgery

2. [ ] Claims Nos.:
   because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. [ ] Claims Nos.:
   because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1. [ ] As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. [ ] As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. [ ] As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically Claims Nos.:

4. [ ] No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the Invention first mentioned in the claims. It is covered by claims Nos.:

Remark on Protest

[ ] The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.

[ ] The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.

[ ] No protest accompanied the payment of additional search fees.
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