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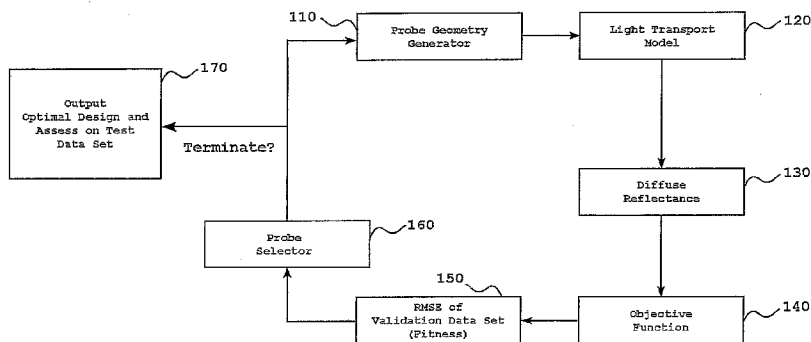
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(54) Title: METHODS, SYSTEMS, AND COMPUTER PROGRAM PRODUCTS FOR OPTIMIZATION OF PROBES FOR SPECTROSCOPIC MEASUREMENT IN TURBID MEDIA



(57) Abstract: The presently disclosed subject matter provides methods, systems, and computer program products for optimizing a probe geometry for spectroscopic measurement in a turbid medium. According to one method, a probe geometry comprising one emitting entity for emitting electromagnetic radiation into a turbid medium and at least one collecting entity for collecting the electromagnetic radiation that has interacted with the turbid medium is selected. A simulation is performed with inputs of the probe geometry and a plurality of sets of optical property values associated with the turbid medium to generate output comprising optical parameter values measured by the probe geometry for each set of input optical property values. The measured optical parameter values are input to an inversion algorithm to produce corresponding optical properties as output. The produced optical properties are compared with optical properties known to correspond to the measured optical parameter values and a degree of matching between the produced optical properties and the known optical properties is determined. The simulation and inversion steps are repeated for a plurality of additional probe geometries. Each additional probe geometry differs from the previously tested probe geometry in at least one property. The property may be a quantity of collecting entities, a diameter of at least one emitting or collecting entity, a linear between the emitting and collecting entities, or combinations thereof. An optimization algorithm is applied at each iteration to select a probe geometry such that the resulting degree of matching will converge to an optimum value. An optimal geometry is selected based on the degree of matching determined for each geometry.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

DESCRIPTION

METHODS, SYSTEMS, AND COMPUTER PROGRAM PRODUCTS FOR OPTIMIZATION OF PROBES FOR SPECTROSCOPIC MEASUREMENT IN TURBID MEDIA

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CROSS REFERENCE TO RELATED APPLICATIONS

This application claims the benefit of U.S. Provisional Patent Application Serial No. 60/702,228, filed July 25, 2005, the disclosure of which is incorporated herein by reference in its entirety.

10

GRANT STATEMENT

The presently disclosed subject matter was made with United States Government support under Grant No. CA100599 awarded by the National Institutes of Health. Thus, the United States Government has certain rights in the presently disclosed subject matter.

15

TECHNICAL FIELD

The presently disclosed subject matter relates to the field of fiber optics. More particularly, the presently disclosed subject matter relates to methods, systems, and computer program products for optimization of probes for spectroscopic measurement in turbid media.

20

BACKGROUND

Diffuse reflectance spectra, which are indicative of the absorption and scattering properties of cells and/or tissues, are sensitive to a number of important biological molecules. In cells and/or tissues, absorption is due at least in part to the presence of various biological molecules, such as proteins, carotenoids, and hemoglobin, and scattering is attributed *inter alia* to the size and density of intracellular and extracellular structures. Diffuse reflectance spectroscopy has therefore been investigated as a possible approach to diagnosing early pre-cancerous and cancerous changes in such cells and/or tissues (Thueller *et al.* (2003) 8 *J Biomed Opt* 495-503; Muller *et al.* (2001) 40 *Appl Opt* 4633-46; Palmer *et al.* (2003) 50 *IEEE Trans Biomed Eng* 1233-42;

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30

Finlay & Foster (2004) 31 *Med Phys* 1949-59; Georgakoudi *et al.* (2002) 62 *Cancer Res* 682-687, 2002). However, due to the complex interplay between absorbers and scatterers in cells and/or tissues, it can be difficult to relate a measured diffuse reflectance spectrum to the underlying physical features of the cells and/or tissues.

The illumination/collection geometry of the probe that is employed can be an important aspect of cell/tissue optical spectroscopic measurements in that it can affect sensitivity to the optical properties (absorption and scattering coefficients), sensing volume, and signal to noise (Mourant *et al.* (1997) 36 *Appl Opt* 5655-5661; Zhu *et al.* (2003) 8 *J Biomed Opt* 237-247; Pogue & Burke (1998) 37 *Appl Opt* 7429-36). There are numerous possible probe designs to select from for a given biomedical application.

Specialized probe designs have been previously shown to be useful in characterizing tissue properties from fluorescence (Pogue & Burke (1998) 37 *Appl Opt* 7429-36; Pfefer *et al.* (2004) 42 *Med Biol Eng Comput* 669-73; Pfefer *et al.* (2005) 10 *J Biomed Opt* 44016; Zhu *et al.* (2005) 10 *J Biomed Opt* 024032; Quan & Ramanujam (2004) 29 *Opt Lett* 2034-2036) and diffuse reflectance measurements (Mourant *et al.* (1997) 36 *Appl Opt* 5655-5661; Amelink *et al.* (2004) 29 *Opt Lett* 1087-1089). For example, Mourant *et al.* (36 *Appl Opt* 5655-5661, 1997; hereinafter, "Mourant") discloses that at a source-detector separation of approximately 1.7 mm, the diffuse reflectance collected was insensitive to the scattering coefficient.

Thus, the measured diffuse reflectance could be directly related to the absorption coefficient. Mourant further discloses that for a source-detector separation of 1.7 mm, this relationship is valid for absorption coefficients in the range of 0-0.86 cm^{-1} and reduced scattering coefficients in the range of 7-21 cm^{-1} . Using this relationship, the authors were able to extract the concentration of Direct Blue dye from a phantom with errors of 20% or less. This method furthermore required no *a priori* information about the absorbers and scatterers present in the medium.

However, the error for the reported probe is potentially too great to allow the disclosed probe to be employed for sensitive medical applications, and it is not valid for optical properties typical of tissue in the UV-visible wavelength

range. Additionally, Mourant does not optimize the geometry of the fiber optic probe, instead simply testing only the operation of probes with a different separation between source and detector fibers. What are needed, then, are methods for testing various parameters of fiber optic probes for spectroscopic measurements that can be used to optimize probe geometries for applications for which enhanced accuracy is important.

To address this need, the presently disclosed subject matter provides methods for optimizing a fiber optic probe geometry for spectroscopic measurement. Such methods are useful for identifying probe geometries that can be employed for measuring optical properties of cells, tissues, or other turbid media.

SUMMARY

The presently disclosed subject matter provides methods, systems, and computer program products for optimizing a probe geometry for spectroscopic measurement in a turbid medium. According to one method, a probe geometry comprising one emitting entity for emitting electromagnetic radiation into a turbid medium and at least one collecting entity for collecting electromagnetic radiation that interacted with the turbid medium is selected. A simulation is performed with inputs of the probe geometry and a plurality of sets of optical property values associated with the turbid medium to generate output comprising optical parameter values measured by the probe geometry for each set of input optical property values. The measured optical parameter values are input to an inversion algorithm to produce corresponding optical properties as output. The produced optical properties are compared with optical properties known to correspond to the measured optical parameter values and a degree of matching between the produced optical properties and the known optical properties is determined. The simulation and inversion steps are repeated for a plurality of additional probe geometries. Each additional probe geometry differs from the previously tested probe geometry in at least one property. The property may be a quantity of collecting entities, a diameter of at least one emitting or collecting entities, a linear distance between the emitting and collecting entities, or combinations thereof. An optimization algorithm is

applied at each iteration to select a probe geometry such that the resulting degree of matching will converge to an optimum value. An optimal geometry is selected based on the degree of matching determined for each geometry.

5 The terms "emitting entity" and "collecting entity" refer to any structures capable of respectively emitting and collecting electromagnetic radiation at wavelengths of interest. Examples of structures suitable for use as the emitting and collecting entities include optical fibers capable of emitting and collecting light. However, even though the examples below relate to a probe that includes optical fibers, the subject matter described and claimed herein is not limited to
10 optimizing a probe that emits and collects visible wavelengths of electromagnetic radiation. Optimizing a probe that emits and collects any wavelengths of electromagnetic radiation suitable for determining properties of turbid media is intended to be within the scope of the subject matter described herein.

15 The subject matter described herein for optimization of probes for spectroscopic measurement in turbid media may be implemented using a computer program product comprising computer executable instructions embodied in a computer readable medium. Exemplary computer readable media suitable for implementing the subject matter described herein include
20 chip memory devices, disk memory devices, programmable logic devices, application specific integrated circuits, and downloadable electrical signals. In addition, a computer program product that implements the subject matter described herein may be located on a single device or computing platform or may be distributed across multiple devices or computing platforms.

25

BRIEF DESCRIPTION OF THE DRAWINGS

Preferred embodiments of the subject matter described herein will now be explained with reference to the accompanying drawings of which:

30 Figure 1 is a block diagram illustrating exemplary components of a system for optimization of fprobes for spectroscopic measurement in turbid media according to an embodiment of the subject matter described herein;

Figure 2 is a flow chart illustrating an exemplary process for optimization of a probe for spectroscopic measurement in turbid media according to an embodiment of the subject matter described herein;

Figure 3 depicts an optimized fiber design showing illumination (gray) and collection fibers 1 and 2;

Figures 4A and 4B are scatter plots of the extracted vs. expected absorption coefficient (Figure 4A) and reduced scattering coefficient (Figure 4B);

Figures 5A and 5B are log contour plots of the collected diffuse reflectance from fiber 1 (Figure 5A) and fiber 2 (Figure 5B);

Figures 6A and 6B are scatter plots of the extracted vs. true optical properties for experimental studies over the wavelength range 400-616 nm for the absorption coefficient (Figure 6A) and for the reduced scattering coefficient (Figure 6B);

Figures 7A and 7B are plots depicting the ratio of the measured to simulated diffuse reflectance spectra at the smaller (Figure 7A) and larger (Figure 7B) source detector separations for phantoms with scattering level 2; and

Figures 8A and 8B are scatter plots of the extracted vs. true absorption coefficient (Figure 8A) and for the reduced scattering coefficient (Figure 8B) for experimental studies over the wavelength range 450-616.

DETAILED DESCRIPTION

All references cited herein are incorporated herein by reference in their entireties to the extent that they supplement, explain, provide a background for, or teach methodology, techniques, and/or compositions employed herein.

Figure 1 is a block diagram illustrating exemplary components of a system for optimization of probes for spectroscopic measurement in turbid media according to an embodiment of the subject matter described herein. Referring to Figure 1, the system includes a probe geometry generator **110** for generating a probe geometry to be tested. Probe geometry generator **110** may generate an initial probe geometry including sizes, spacings, and numbers of illumination and collection fibers. The sizes of the fibers may be selected from

a group of available fibers sizes (e.g., fibers sizes that are commercially available, including, but not limited to 50 μm , 100 μm , 200 μm , 400 μm , and/or 500 μm). Initial spacings may be selected randomly from increments within bounds defined by the spectroscopic measurement environment. In some
5 embodiments, the initial spacings are limited to a maximum value, and in one implementation, the maximum value is 1.5 mm.

A light transport model **120** may be used to simulate diffuse reflectance properties measured by the probe geometry. In one implementation, light transport model **120** may be implemented using a Monte Carlo simulation
10 model. The output of light transport model **120** for the given probe geometry is diffuse reflectance **130** that would be measured by the probe geometry. In one implementation, ranges of optical properties are input and corresponding diffuse reflectance values are output.

Diffuse reflectance **130** is input to an objective function **140**, which takes
15 as input the diffuse reflectance values computed for the different optical properties generated by light transport model **120**. The objective function implements an inversion algorithm that determines optical properties corresponding to the diffuse reflectance values produced by light transport model **120** and compares the produced optical properties with optical
20 properties known to correspond to the input diffuse reflectance values. Objective function **140** computes a degree of closeness or matching between the produced optical properties and the known optical properties. In one implementation, the degree of closeness or matching is the root mean squared error (RMSE) **150** between the produced and known optical properties. A
25 probe selector **160** receives the RMSE value for each probe design and selects a probe design that minimizes the RMSE value. Probe selector **160** may apply an optimization algorithm at each iteration to select a probe geometry such that the resulting degree of matching will converge to an optimum value. In one implementation, the optimization algorithm may be a genetic algorithm.

30 Figure 2 is a flow chart illustrating an exemplary process for optimization of a probe for spectroscopic measurement in turbid media according to an embodiment of the subject matter described herein. Referring to Figure 2, the

fiber optic probe optimization method will be described hereinafter. First, at step **S110**, an initial probe geometry is generated. The initial probe geometry can comprise at least one illumination fiber and at least one collection fiber. In some embodiments, the illumination fiber and the collection fiber are the same fiber, and in some embodiments, the illumination fiber is not a collection fiber.

At step **S120**, diffuse reflectance is modeled. In order to model diffuse reflectance, a model of light transport that has the ability to quickly and efficiently calculate the diffuse reflectance measured by a given probe geometry for a wide range of optical properties is employed. In some embodiments, a Monte Carlo model of light transport can be used for this purpose.

Next, an appropriate objective function is required to quantitatively evaluate the effectiveness or fitness of a probe having a particular geometry in extracting the optical properties from a turbid medium. In some embodiments, the objective function can employ a neural network algorithm to relate the measured diffuse reflectance to the optical properties of the medium. The neural network can be optimized on a training set and then evaluated on an independent validation set.

At step **S130**, the root mean square error (RMSE) between the extracted (from the neural network algorithm) and actual optical properties (input into the Monte Carlo simulation) can be output from the objective function to determine the fitness of that particular probe.

At step **S140**, it is determined whether or not the optimal geometry condition is satisfied. If so, the optimal geometry is outputted, and if not, a new probe geometry is generated at step **S150**, and steps **S110-S140** are iteratively repeated. A genetic optimization algorithm can be used to find the most fit fiber optic probe design, *i.e.*, one that minimizes the RMSE of the optical properties calculated by the objective function, through an adaptive process. The RMSE of the optical properties generated by the objective function using the optimal probe geometry can be evaluated on an independent testing data set. A commercially available genetic algorithm suitable for selecting a probe design that minimizes the RMSE is the GAlib algorithm.

EXAMPLES

The following Examples have been included to illustrate modes of the presently disclosed subject matter. In light of the present disclosure and the general level of skill in the art, those of skill will appreciate that the following

5 Examples are intended to be exemplary only and that numerous changes, modifications, and alterations can be employed without departing from the scope of the presently disclosed subject matter.

EXAMPLE 1

10 Reflectance Calculation Using a Model of Light Transport

One of the aspects of the optimization process outlined in Figure 2 is a model of light transport that can generate a diffuse reflectance spectrum for a wide range of optical properties and probe geometries. In some optical property and fiber geometry regimes, the diffusion equation could be used.

15 However, since with regard to reflectance in the UV-VIS range, where tissue is highly absorbing, this method is unsuitable. Therefore, Monte Carlo modeling was used to compute the diffuse reflectance for a given probe geometry. Monte Carlo simulations with point source geometries were used, so that any arbitrary probe geometry could be modeled using convolution.

20 In addition, the scaling procedures described by Graaff *et al.* (32 *Appl Opt* 426-34, 1993) were used to scale the output of a single Monte Carlo simulation for any set of optical properties as disclosed in Palmer & Ramanujam (2006) 45 *Appl Opt* 1062-1071. Briefly, the method comprised running a single simulation for a given set of absorption ($\mu_{a,sim}$) and scattering

25 ($\mu_{s,sim}$) coefficients and recording the exit weight ($W_{exit,sim}$), net distance traveled ($r_{t,sim}$), and total number of interactions for each photon (N) that exits the tissue surface. The scaling method then used these stored parameters to calculate the new exit weight ($W_{exit,new}$) [Eq. (1)] and net distance traveled ($r_{t,new}$) [Eq. (2)] for a given photon that had a different absorption ($\mu_{a,new}$) and scattering

30 coefficient ($\mu_{s,new}$) used in the same simulation. The scaling relationships were:

$$W_{exit,new} = W_{exit,sim} \left(\frac{\mu_{s,new}}{\mu_{s,new} + \mu_{a,new}} \frac{\mu_{s,sim} + \mu_{a,sim}}{\mu_{s,sim}} \right) \quad (1)$$

$$r_{t,new} = r_{t,sim} \left(\frac{\mu_{s,sim} + \mu_{a,sim}}{\mu_{s,new} + \mu_{a,new}} \right) \quad (2)$$

To further simplify the scaling process, it was assumed that, for a given
 5 value of the reduced scattering coefficient, $\mu_s' = \mu_s(1-g)$, the diffuse reflectance
 would be the same for any values of μ_s and anisotropy factor (g) that generate
 the same μ_s' . This has been shown to be valid over the range of g values
 present in human tissue (*i.e.*, for g values greater than 0.8; Graaff *et al.* (32
Appl Opt 426-34, 1993); Kienle & Patterson (41 *Phys Med Biol* 2221-2227,
 10 1996)). By use of this similarity relation and the scaling procedure outlined
 hereinabove, only a single Monte Carlo simulation needed to be run to
 determine the output of a Monte Carlo simulation for any set of optical
 properties. The Henyey-Greenstein phase function was used in the single
 Monte Carlo simulation as described hereinbelow.

15 Convolution was used to integrate over the illumination and collection
 fibers to determine the probability that a photon, traveling a fixed distance,
 would be collected for a given probe geometry. This takes advantage of the
 spatial invariance and rotational symmetry present in a homogeneous medium.

For a pair of illumination and collection fibers, the probability of collection of a
 20 photon traveling a net distance r_t between the points of entering and exiting the
 medium is given by:

$$\frac{1}{\pi^2 r_i^2} \int_{\max(-r_i, s-r_i-r_c)}^{\min(r_i, s-r_i+r_c)} (s-x) \cos^{-1} \left[\frac{s^2 + (s-x)^2 - r_i^2}{2(s-x)s} \right] \\ \times \cos^{-1} \left[\frac{r_t^2 + (s-x)^2 - r_i^2}{2(s-x)r_t} \right] dx \quad (3)$$

where r_i is the radius of the illumination fiber, r_c is the radius of the collection
 25 fiber, s is the separation between the centers of the illumination and the
 collection fibers, and x is the spatial variable over which the integral is taken
 (see appendix for derivation). This equation was numerically integrated. To
 adapt this to the fiber bundle used in this study (the geometry of the fiber

bundle is described below), the common end of the fiber bundle was imaged, and the centers of each illumination and collection fiber in the bundle were determined. Then the probe geometry was integrated pairwise (for each illumination- collection fiber pair) to determine the total probability of collection.

5 It was found that imaging the fiber bundle to obtain the exact location of each illumination and collection fiber was necessary. Approximating the fiber bundle as solid rings of illumination and collection fibers produced significant errors in the model, likely because of imperfect physical placement of the fibers within the bundle.

10 It was found that the scaling process and subsequent numerical integration for the probe geometry of a large number of photons required approximately 1 s to complete, which, although much faster than running an independent simulation, was still rather slow for performing an inversion procedure. Therefore, the diffuse reflectance values for a range of optical
15 properties (μ_s , 5-500 cm^{-1} ; μ_a , 0-200 cm^{-1} ; g , 0.8) were determined ahead of time to form a lookup table, and cubic splines were used to interpolate between table values. The smallest increment used in the lookup table was 0.1 cm^{-1} for μ_a and 2.5 cm^{-1} for μ_s . This allowed for rapid determination of the diffuse reflectance for a given fiber probe geometry and a wide range of optical
20 properties, without requiring independent Monte Carlo simulations.

Three simulated data sets were generated using Monte Carlo simulations: training, validation, and testing data sets. The training data set was used to train the neural network algorithm to extract optical properties from diffuse reflectance measurements with a particular probe geometry over a wide
25 range of optical properties. The validation and testing data sets were used in two different stages of the optimization process. The validation data set was used in each iteration of the optimization loop shown in Figure 2, to evaluate the fitness of a given probe geometry with an independent set of optical properties (which were not used in training the algorithm). The RMSE
30 calculated from the results of the validation data set was used as the measure of probe fitness. The optimal probe design selected by the genetic algorithm at the end of the iterative process was applied to the testing data set. The output of the testing data set was used to determine the accuracy with which the

optimal probe design could extract optical properties from an independent data set in an unbiased manner.

Training Data Set. A single Monte Carlo simulation, consisting of 55×10^6 photons, having the following properties: absorption coefficient (μ_a): 0, scattering coefficient (μ_s): 150 cm^{-1} , anisotropy factor (g): 0.8, was first simulated. The refractive indexes were chosen to be 1.4 for the medium, which is representative of a number of tissue types (Bolin *et al.* (1989) 28 *Appl Opt* 2297-2303), and 1.47 for the fiber (representative of fused silica) above this medium. The medium was semi-infinite and homogeneous. The simulation was run using a point-source geometry and photons were collected at the surface. The scaling procedures described by Graaff *et al.* (32 *Appl Opt* 426-34, 1993) were used to adjust the simulation to any desired set of optical properties. Convolution was used to account for the specific fiber probe geometries evaluated in the optimization process. The Quasi-Discrete Hankel Transform, described by Li *et al.* (23 *Opt Lett* 409-11, 1998) was used to perform the convolution.

The training data set consisted of 144 sets of optical properties. The total attenuation coefficient μ_t (defined as $\mu_a + \mu_s$) ranged from 25 to 200 cm^{-1} . At each μ_t , the albedo (defined as μ_s / μ_t) ranged from 0.6 to 1, with g fixed at 0.8. This corresponds to a range of μ_a from 0 to 80 cm^{-1} and reduced scattering coefficient μ_s' (defined as $\mu_s \times (1-g)$) from 3 to 40 cm^{-1} . This set of optical properties encompasses that found in a number of tissue types in the UV-VIS spectral range (see Cheong (1995) in Optical-Thermal Response of Laser-Irradiated Tissue, Lasers, Photonics, and Electro-optics (Welch & Gemert (eds)), Plenum Press, New York, New York, United States of America, pp. 275-303), and is consistent with findings of the optical properties in human breast tissue reported by the co-inventors (Palmer *et al.* (2006) 45 *Appl Opt* 1072-1078). μ_t was assigned because for a given fixed μ_t , the spatial scaling of the Monte Carlo simulation to accommodate the desired set of optical properties is fixed. This leads to an increase in the efficiency of the Hankel transform by allowing much of the computational load involved in calculating the Hankel transform to be conserved (Li *et al.* (23 *Opt Lett* 409-11, 1998)

refers to this as the "C" matrix, which needs only be calculated once in this case). The output is the modeled diffuse reflectance for a wide range of optical properties, calculated for the specific fiber geometry being evaluated.

Validation/Testing Data Set. The validation and testing data sets both
5 consisted of 25 randomly assigned sets of optical properties (chosen from within the range specified for the training data set). The anisotropy factor, g , was also randomly chosen within the range of 0.8 to 0.95, while scaling μ_s to keep μ_s' within the same range used in the training data set. Independent Monte Carlo simulations were run for the 25 sets of optical properties with 10^6
10 photons used in the validation and testing data sets and convolution was again used to model a specific probe geometry. As stated above, the validation data set was used in each iteration of the optimization loop to evaluate fitness of a probe geometry to a set of optical properties. The testing data set was used to test the probe geometry selected using the genetic algorithm for its ability to
15 extract optical properties.

EXAMPLE 2

Objective Function

Once the diffuse reflectance for a probe geometry has been modeled
20 using the light transport model, the next phase of the optimization flow chart shown in Figure 2 is the objective function. In order to extract the optical properties of the medium from its diffuse reflectance, a neural network objective function was used to determine the optical properties based on the diffuse reflectance collected for a given probe geometry. A neural network
25 objective function was chosen because it is useful in approximating complex non-linear functions and has previously been shown effective in extracting optical properties (Pfefer *et al.* (2003) 8 *J Biomed Opt* 206-215). This works on the principle that the spatial distribution and intensity of the diffusely reflected light can be determined by the optical properties of the medium, under the
30 assumption that given enough sampling points (in this case collection fibers), these optical properties can be uniquely determined.

The neural network employed consisted of 10 neurons having hyperbolic tangent activation functions, and two output neurons (corresponding to μ_a and

μ_s') having linear activation functions. The neural network was trained using the Monte Carlo generated training data set described hereinabove using a Levenberg-Marquardt algorithm in MATLAB® (Mathworks Inc., Natick, Massachusetts, United States of America).

5 Briefly, the neural network works as an interconnected assembly of simple processing units. The neurons themselves are fixed as simple functions (in this case hyperbolic tangent or linear functions). The network is adaptable by weighting the connections from one neuron to the next: *i.e.*, the output of one neuron is the input to another neuron, modulated by a particular weight.

10 The universal approximation theorem states that in using a neural network with only a single hidden layer (such as the one employed herein – having one layer of neurons between the input and output neuron layers), any continuous function can be approximated with arbitrary precision (Chen & Chen (1993) 4 *IEEE Trans Neural Networks* 910-918). This thus represents a powerful tool for

15 approximating complex, non-linear relationships where analytical solutions are not possible.

Upon training the neural network for a given fiber geometry, the fitness of that individual fiber geometry was quantified by the RMSE of the extracted optical properties from the 25 independent simulations in the validation set.

20 The RMSE was calculated separately for absorption and reduced scattering coefficients (μ_a and μ_s' , respectively). The RMSE for the absorption coefficient and the RMSE for the reduced scattering coefficient were then summed, and this value was used to characterize the fitness of the particular probe fiber geometry. This procedure allowed for selection of a fiber geometry that

25 performs well not only on the training data set, but which also generalizes well to a set of independent simulations.

EXAMPLE 3

Optimization with Genetic Algorithm

30 The goal of the genetic algorithm was to minimize the returned fitness score of a given fiber geometry, thereby reducing the error with which the optical properties can be extracted. Genetic algorithms represent a robust means of optimization and have advantages over commonly used gradient

techniques in that they are less sensitive to the initial guess and perform well in regions of small gradients or discontinuities (see e.g., Eiben & Smith (2003) Introduction to Evolutionary Computing, New York, New York, United States of America, Springer-Verlag).

5 Genetic algorithms have been effectively employed in optimization applications. For example, they have been used successfully in optimizing the microstructure of communications fibers to maximize throughput (Manos & Poladian (2004) *21 Eng Comput (Swansea, Wales)* 564-576). As opposed to gradient methods, which take an initial guess and attempt to move it in a
10 direction that leads to a better solution, genetic algorithms work with a population of solutions. Operators similar to those of natural evolution, such as crossover, mutation, and selection are employed to produce offspring and evolve the population of solutions toward an optimal solution.

In generating a new solution, two "parent" solutions were selected from
15 which a new "child" solution was derived. "Crossover" refers to the blending of the two parent solutions to form a child solution (e.g., the illumination fiber diameter of the child solution might be chosen to be the same as the diameter of one of the parent solutions). Random mutation introduced random variability into the child solution (e.g., by randomly altering the source-detector
20 separation), to introduce greater diversity into the resulting solution set. The specific guidelines by which these operators were applied are described in the following section.

The population of solutions was updated in steps known as generations. At each generation a population of child solutions was generated from the
25 parent population. Parent individuals for the next generation of child solutions were chosen based on the process of selection. Selection was weighted towards the fittest individuals (i.e., fitter individuals are more likely to be chosen as parents), thereby introducing selective pressure on the population to evolve towards an optimal solution.

30 Variable Fiber Parameters. The GAlib genetic optimization library (Wall (2005) "GAlib: Matthew Wall's Genetic Algorithm Library," vol. 2005) was used to optimize the fiber design parameters. A number of adjustable fiber parameters are possible. These include the numerical aperture (NA), fiber

diameter, and number and location of the illumination and collection fibers. It is also possible to use angled fibers to alter the direction of photon propagation, and so affect the probing volume (Quan & Ramanujam (2004) 29 *Opt Lett* 2034-2036). In order to simplify the modeling, only the diameter and source-detector separations were included as free parameters.

Two basic probe designs were tested. In the first case, the probe consisted of a single illumination fiber, and between two and six collection fibers (a total of five possible configurations). In the second case, there was similarly a single illumination fiber, but it was also used for collection of light. There were additional fibers (between one and five) that served to collect light only (another 5 possible configurations). Thus, each of the basic probe designs had between two and six independent channels of light collection, for a total of ten configurations.

A series of optimizations employing the methodology shown in Figure 2 was run to consider each of the ten probe configurations separately. This approach allowed for a comparison of the cases where overlapping and non-overlapping source-detector geometries were used and also permitted an evaluation of additional design complexity by adding additional channels of light collection.

The illumination fiber had a variable diameter, having possible values of 50, 100, 150, 200, 300, 400, and 500 μm . These properties corresponded to commercially available fibers. The remaining collection fibers also had a variable fiber diameter, as well as a variable center-to-center distance from the source fiber. The possible diameters consisted of the same set of diameters used for the source fiber, while the source-detector separation was limited to less than 1.5 mm, and greater than the sum of the source and collection fiber radii (cladding included). All fibers had a numerical aperture (NA) of 0.22 and specular reflection was not collected.

Genetic optimization parameters. First, a set of 25 fiber optic probe solutions was randomly initialized within the set of possible solutions outlined above, to form the initial population of solutions. From this set pairs of two at a time were randomly selected. Next, a child solution was generated, which exhibited a mixture of the characteristics of the selected parent solutions. For

the fiber diameter, the diameter of the offspring was selected randomly from either of the two parents. For the source-detector separation distance, a blending algorithm (Wall (2005) "GAlib: Matthew Wall's Genetic Algorithm Library," vol. 2005) was chosen such that the separation distance was
5 randomly selected from a region centered at the mean of the two parent separation distances, and having a range of twice the difference between the two parent separations. Crossover was performed with a probability of 0.9 for each new individual. In cases where it was not performed, the child solution was identical to one of the parents.

10 In addition to the crossover operator, a random mutation operator was also employed to introduce greater diversity into the population. This was done after a child solution was generated, and when mutation occurred, the mutated solution took the place of the original child solution. For the fiber size, a new size was randomly selected from the allowed set (commercially available fiber
15 sizes). For the separation distance, a random Gaussian number with a standard deviation of 0.3 mm was added to the separation of the fiber. The random mutation was applied with a probability of 0.005 for each variable. For all operators, the bounds of the variables, as described above, were respected. The probabilities at which the crossover and mutation operators were applied
20 were chosen based on commonly used values in the literature.

Finally, the roulette method of selection (Eiben & Smith (2003) Introduction to Evolutionary Computing, Springer-Verlag, New York, New York, United States of America) was employed to select offspring that make up the next generation, weighted towards the most fit individuals. This method was
25 analogous to spinning a roulette wheel, where larger sections of the wheel were devoted to the fittest individuals. In other words, the fittest individuals had a higher probability of being selected as parents for the next generation. This was the mechanism that drove the population of solutions towards the optimum. A total of 500 generations were run with the best individual taken as
30 the final solution.

After termination of the optimization loop, the optimal fiber geometry was output by the algorithm. The neural network algorithm incorporating the optimal probe geometry was applied to a testing data set in order to provide an

unbiased estimate of the effectiveness of a given fiber geometry/neural network algorithm for extracting optical properties from the diffuse reflectance signal.

EXAMPLE 4

5

Phantom Validation

The approach disclosed herein was experimentally validated by constructing a fiber optic probe according to the specifications for the optimal probe provided by the optimization algorithm and using it to acquire diffuse reflectance measurements from tissue phantoms that have optical properties that fall within the range used in the simulation studies. The constructed probe consisted of a single source fiber and two separate collection fibers (which was found to have the best performance in the simulation studies). The tip of the probe was imaged to determine the fiber sizes and their positions relative to each other, and the neural network algorithm was trained using a training data set of Monte Carlo simulations for the exact probe geometry at the tip of the constructed fiber probe.

The probe was coupled to a custom built spectrometer, consisting of a 450-W xenon lamp (FL-1039, HORIBA Jobin Yvon Inc., Edison, New Jersey, United States of America), a scanning double-excitation monochromator (Gemini 180, HORIBA Jobin Yvon Inc.), an imaging spectrograph (IHR320, HORIBA Jobin Yvon Inc.), and a CCD camera (Symphony, HORIBA Jobin Yvon Inc.). All measurements were conducted with the excitation monochromator passing zero order white light, with slits at 1 mm. The imaging spectrograph was set to have a center wavelength of 485 nm, corresponding to a range of 353-616 nm. The entrance slit to the imaging spectrograph was set to 0.4 mm, corresponding to a full-width half maximum bandpass of 3.76 nm. Integration times ranged from 40 ms to 3 s.

A set of liquid homogeneous phantoms was then created. The phantoms contained variable concentrations of hemoglobin (absorber) and polystyrene spheres (scatterer). First, three solutions with variable volume densities of polystyrene spheres (07310-15, Polysciences, Inc., Warrington, Pennsylvania, United States of America) suspended in water were made to produce three phantoms with different scattering coefficients. To each of these

base phantoms, 3 titrations of hemoglobin (H7379-16, Sigma Co., St. Louis, Missouri, United States of America) solution were added to produce three different absorption coefficients, with a diffuse reflectance measurement made after each addition. This produced a set of 9 phantoms with a range of absorption and scattering properties. All measurements were conducted the day the phantoms were made.

The wavelength dependent extinction coefficients for hemoglobin were measured using an absorption spectrophotometer (Cary 300, Varian, Inc., Palo Alto, California, United States of America). It was assumed that the oxygenation of hemoglobin was constant through the course of the experiment.

The reduced scattering coefficient was determined from Mie theory using freely available software (Prahl (2003) "Mie Scattering Program", vol. 2003: Oregon Medical Laser Center), given the known size (1 μm), density, and refractive index of the spheres (1.60) and the surrounding medium, water (1.33). The refractive index of polystyrene spheres has been reported to be constant to within approximately 1% of this value over the wavelength range used (Xiaoyan *et al.* (2003) 48 *Phys Med Bio* 4165-4172). Tables 1 and 2 show the means and ranges of μ_s' and μ_a , respectively, for each absorption and scattering level over the wavelength range of 353-617 nm. All combinations of these absorption and scattering levels were measured with the fiber-optic probe and spectrometer. The range of optical properties represents a subset of those used in the simulation studies, but still cover a range of optical properties representative of those found in the human breast and other tissues (Palmer *et al.* (2006) 45 *Appl Opt* 1072-1078).

Table 1

Reduced Scattering Coefficients for Each of Three Concentrations of Polystyrene Spheres Employed in Phantom Experiments

Phantom	Mean μ_s' (cm^{-1})	μ_s' range (cm^{-1})
μ_s' level 1	6.1	5.5-6.9
μ_s' level 2	16	14-18
μ_s' level 3	24	22-28

Table 2
Absorption Coefficients for Each of Three Concentrations of Hemoglobin
Employed in Phantom Experiments

Phantom	Mean μ_a (cm^{-1})	μ_a range (cm^{-1})
μ_a level 1	1.3	0.35-6.0
μ_a level 2	2.6	0.69-12
μ_a level 3	5.0	1.3-23

5

To correct for the instrument response, and the difference in magnitude between the Monte Carlo simulations (which are on an absolute scale) and those of the phantoms (which are on a relative scale) a single phantom with known optical properties was used as a reference. The ratio of the simulated spectrum, given the known optical properties of this phantom, and its measured spectrum was taken. All experimental data was multiplied wavelength by wavelength by this calibration ratio.

Table 3 shows the RMSE in μ_a and μ_s' selected from the testing data sets for the optimal probe configuration from each of the ten basic fiber probe configurations. The optimal probe geometries are also listed. It can be seen that as the number of fibers increased beyond two collection fibers, the RMSE of the testing data set did not decrease. This indicated that two collection fibers were sufficient for extraction of the optical properties, and the use of additional fibers led to over-training of the algorithm. Fiber probe configurations with additional fibers (greater than 5 collection channels) were not considered for this reason.

20

Table 3
Root Mean Square Errors (RMSE) in μ_a and μ_s from Testing Data Sets for the Optimal Probe Configuration
Selected from Each of Ten Basic Fiber Probe Configurations*

Fiber design	RMSE – testing (cm^{-1})		Optimal Geometry	
	μ_a	μ_s	Illumination Diameter (μm)	Diameter – Source Detector Separation Pairs fo Each Collection Fiber (μm)
3 total fibers, no collection by source fiber	0.41	0.30	500	(200-380) (400-1360)
4 total fibers, no collection by source fiber	0.47	0.32	200	(200-610) (500-1330) (500-440)
5 total fibers, no collection by source fiber	0.46	0.34	400	(50-550) (400-530) (50-1180) (300-1230)
6 total fibers, no collection by source fiber	0.40	0.29	400	(200-640) (300-1120) (400-490) (500-1190) (50-700)
2 total fibers, collection by source fiber	0.48	0.81	500	(500-1490)

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Fiber design	RMSE – testing (cm ⁻¹)	Optimal Geometry
3 total fibers, collection by source fiber	0.49 2.30 500	(150-1430) (150-1370)
4 total fibers, collection by source fiber	1.07 1.09 500	(100-1390) (50-1270) (100-1250)
5 total fibers, collection by source fiber	0.50 0.89 500	(100-1200) (50-1220) (400-1130) (500-1320)

* The optimal probe geometry is also provided, showing the diameter of the illumination fiber, with the collection fiber diameter and source detector separation pairs for each of the collection fibers – listed as (diameter – separation) for each fiber.

In addition, the probe configurations where the source fiber was not used for collection outperformed the case where it was. The fiber design that performed the best was the case where there were 3 total fibers, and the source fiber was not used for collection. This fiber design provided the best balance between maximizing performance and minimizing probe complexity.

Figure 3 shows the optimal fiber probe geometry. It consisted of a single 500 μm diameter illumination fiber, a 200 μm diameter collection fiber at a center-to-center distance of 380 μm , and a second 400 μm diameter collection fiber at a center-to-center distance of 1360 μm . In Figure 3, solid areas indicate fiber dimensions to scale. This design comprises 3 total fibers where the illumination fiber is not used for collection.

Figure 4 shows a scatter plot of the extracted vs. expected optical properties obtained from the testing data set for the probe design shown in Figure 3. The solid lines in Figures 4A and 4B depict the line of perfect agreement. These results are for the fiber design with 3 total fibers where the illumination fiber is not used for collection. It can be seen that the extracted optical properties showed minimal deviation from the expected optical properties over the entire range tested for both μ_a (Figure 4A) and μ_s' (Figure 4B). The optical properties were extracted with an RMSE of 0.41 cm^{-1} for μ_a (tested range of 0-80 cm^{-1}) and 0.30 cm^{-1} for μ_s' (tested range of 3-40 cm^{-1}) using the simulated data.

Next, because practical implementation of this probe design would introduce some variability into the positioning of the fibers, the sensitivity of the RMSE to positioning error was evaluated. To evaluate the effects of a wide range of positioning errors, two random numbers were generated following a Gaussian distribution with a mean of zero and a variable standard deviation to evaluate a range of design tolerances. The optimal fiber design described above was altered by adding these numbers to the two source detector separations (both separations were thus randomly altered at the same time). The minimum separation was fixed such that the fibers would not overlap. The neural network was then trained on diffuse reflectance generated for the resulting probe

geometry (that which includes the positioning errors), and evaluated with an independent testing data set for the same probe geometry. The positional errors were incorporated into both the training and testing data sets because in practical implementation, the fiber probe would be in direct contact with the tissue surface, with no shield or casing covering the fibers themselves, thus allowing the tip of each fiber to easily be imaged using a reflected light microscope. Any positional errors in the practical probe geometry can thus be easily incorporated into the model, as was done here.

The process was repeated 5 times (using 5 different sets of random numbers selected from the Gaussian distribution) to evaluate the variability of the error with different random perturbations of the fiber positions. Next, the mean and standard deviations of the RMSE were calculated.

Table 4 shows the results of the sensitivity analysis on fiber positioning error. It can be seen that for the case where the positioning error had a standard deviation of 50 μm , the RMSE was identical to those obtained when the source detector separations was exactly defined. As the positioning error increased, the RMSE also increased, with the error in μ_s' increasing more rapidly than that of μ_a . It also was determined that small changes in the fiber diameters did not greatly impact the accuracy with which the optical properties could be extracted, although this would likely not be as much of a concern in practice since the fiber diameters are well defined.

Table 4
RMSE of Extracted Optical Properties vs. Positioning Errors

Fiber Positioning Standard Deviation (μm)	Mean RMSE (cm^{-1})	
	μ_a	μ_s'
0	0.41	0.30
50	0.41 ± 0.01	0.30 ± 0.02
100	0.43 ± 0.01	0.36 ± 0.06
200	0.46 ± 0.07	0.55 ± 0.35
500	0.51 ± 0.11	0.94 ± 0.70

It is desirable to gain some understanding as to why this particular fiber design performs well. Figure 5 shows a log scale contour plot of the collected diffuse reflectance as a function of μ_t and albedo for each of the two collection fibers (illustrated in Figure 3). The plots show that the contour lines in Figure 5A roughly follow the vertical lines of equal albedo, while the contour lines in Figure 5B roughly follow the lines of equal μ_a (bold lines). It can be seen in Figure 5A that the contour plot of the diffuse reflectance collected from fiber 1 had roughly vertical contour lines. This indicated that for a given albedo (or fixed proportion of μ_s and μ_t), the reflectance collected by this fiber was relatively insensitive to changes in μ_t .

On the other hand, for collection fiber 2 (Figure 5B), the contour lines roughly followed the lines of constant μ_a , which are shown as thick black lines. This indicated that for a given fixed absorption coefficient, the diffuse reflectance was relatively insensitive to changes in scattering. As a result, the diffuse reflectance collected by fiber 1 gave a direct measure of the albedo and the diffuse reflectance collected by fiber 2 gave a direct measure of the absorption coefficient. The scattering coefficient could be determined directly from these two parameters. Note that these relationships appeared to break down somewhat at low values of μ_t , and in this case the extraction of optical properties would become on a more non-linear function of the collected reflectance from each fiber, all of which was handled by the neural network function.

Next, this approach was experimentally validated. A probe was constructed meeting the design specifications shown in Figure 3 and it was used to measure diffuse reflectance from a series of phantoms with a wide range of optical properties. The neural network algorithm was trained using a Monte Carlo training data set as before using the exact fiber configuration in the actual probe, but was tested on experimental measurements made with that probe. Figure 6 shows the extracted vs. true μ_a (Figure 6A) and μ_s' (Figure 6B) for the set of phantoms with optical property ranges in Tables 1-2. These data were obtained using the phantom with scattering level 2 and absorption level 2 as the reference phantom, with all other phantoms used to test the algorithm.

The data from 400-616 nm are shown. Wavelengths shorter than 400 nm were excluded due to poor signal to noise owing to the poor efficiency of the charge coupled device (CCD) and lower lamp output at these wavelengths. It can be seen that the accuracy of the method was somewhat less than that
5 obtained with simulation, which was not unexpected, but the performance degraded particularly for high μ_a . The RMSE for the data shown in Figure 5 was 2.0 cm^{-1} and 2.3 cm^{-1} for μ_a and μ_s' , respectively. Repeating this for all possible reference phantoms, and averaging the RMSE yields mean RMSEs of $2.4 \pm 0.5 \text{ cm}^{-1}$ and $3.7 \pm 1.4 \text{ cm}^{-1}$ for μ_a and μ_s' , respectively. These phantoms had a range
10 of $0.35\text{-}23 \text{ cm}^{-1}$ and $5.2\text{-}26 \text{ cm}^{-1}$ for μ_a and μ_s' , respectively, over this wavelength range.

The sources of error in the experimental results were evaluated by comparing the simulated and diffuse reflectance measurements at the two different source-detector separations in the probe geometry. Figure 7 shows the
15 ratio of the measured and simulated diffuse reflectance for the shorter (Figure 7A) and larger (Figure 7B) source-detector separations for the three phantoms with scattering level two. Only a subset of the phantoms is shown to make the plot easier to interpret, however similar trends are seen in phantoms at the other two scattering levels.

20 For this plot a solution of polystyrene spheres of the same concentration was used as a calibration standard to correct for the wavelength dependent response of the system (*i.e.*, the ratios were normalized wavelength by wavelength to that of the calibration standard). A phantom with no absorber added was chosen as the reference in this case to exclude errors associated
25 with absorption effects from the reference phantom, which tends to cancel out errors seen in the highly absorbing phantoms, and introduce artifacts into the minimally absorbing phantoms. Were the experimental and simulated data in perfect agreement, there would be a ratio of 1 across all wavelengths. It can be seen that there is some deviation for the larger source-detector separation
30 (Figure 7B) around the Soret band of hemoglobin absorption (420 nm).

Finally, the region of discrepancy was excluded by limiting the wavelength range to be from 450-616 nm. Figure 8 shows the extracted vs. true μ_a (Figure 8A) and μ_s' (Figure 8B), again using the phantom with scattering level 2 and absorption level 2 as the reference phantom. It can be seen that there is substantial agreement between the extracted and true optical properties over this limited wavelength range, resulting in a substantial improvement in the accuracy, albeit over a smaller range of absorption coefficients. The RMSE for the data shown in Figure 8 is 0.2 cm^{-1} and 1.4 cm^{-1} for μ_a and μ_s' , respectively. Repeating this for all possible reference phantoms, and averaging the RMSE yielded mean RMSEs of $0.4 \pm 0.1 \text{ cm}^{-1}$ and $2.2 \pm 0.6 \text{ cm}^{-1}$ for μ_a and μ_s' , respectively. The phantoms had a range of $0.35\text{-}3.2 \text{ cm}^{-1}$ and $5.2\text{-}25 \text{ cm}^{-1}$ for μ_a and μ_s' , respectively, over this narrower wavelength range.

EXAMPLE 5

Derivation of Equation (3)

Equation (3) can be used to derive the probability that a photon launched into a circular illumination fiber of radius r_i , which travels a given net distance r_t , will be collected by a separate circular fiber of radius r_c at a fixed center-to-center distance s from the illumination fiber. This probability can be derived for uniform fiber illumination and collection efficiencies. Both fibers are normal to the medium, which produces a circularly symmetric and translationally invariant system, assuming a homogeneous or homogeneous layered medium. Let the illumination fiber be centered at the origin, and the collection fiber be centered at $(s, 0)$. First, the case in which photons are launched only at the center of the illumination fiber is considered. Because the system is circularly symmetric, the photon can exit the surface anywhere along the circle centered at the origin, with radius r_i , with equal probability. The probability that the photon will exit within the region contained by the collection fiber, and thus be collected, is given by $p = r_t\theta/2\pi r^2$, which corresponds to the arc length contained within the collection fiber, divided by the total circumference of the circle that defines all possible exit

locations. This can be shown to be

$$p = \frac{1}{\pi} \cos^{-1} \left(\frac{r_t^2 + s^2 + r_c^2}{2sr_t} \right), \quad s - r_c < r_t < s + r_c, \quad (4)$$

$p = 0$, otherwise.

5

This can then be extended to a line source located at $y = 0$, and $-r_i \leq x \leq r_i$, by noting that a displacement in x in the source effectively changes the source-detector separation s and then integrating. This is normalized to the length of the source line to produce the average probability for all source locations from which the photon could originate. For the following derivations it was assumed that the mean probability of collection is nonzero: *i.e.*, $s - r_i - r_c < r_t < s + r_i + r_c$. In this case the probability of collection is given by:

10

$$p = \frac{1}{2r_i} \frac{1}{\pi} \int_{lb}^{ub} \cos^{-1} \left[\frac{r_t^2 + (s-x)^2 - r_c^2}{2(s-x)r_t} \right] dx, \quad (5)$$

15

where $ub = \min(r_i, s - r_t + r_c)$ and $lb = \max(-r_i, s - r_t - r_c)$. These bounds correspond to the launch locations for which the probability of collection is nonzero.

20

Finally, this system can be extended to a fiber source by noting that the probability of collection is the same for any source location at a given distance from the center of the collection fiber. Thus each point in the integral given in Eq. (5) is weighted by the arc length of all source locations occurring within the source fiber, equidistant to the collection fiber center. The integral is then

normalized to the area of the source fiber to produce the average probability of collection for all possible source locations. This gives

$$P = \frac{1}{\pi r_i^2} \frac{1}{\pi} \int_{lb}^{ub} (s-x) \cos^{-1} \left[\frac{s^2 + (s-x)^2 - r_i^2}{2(s-x)s} \right] \times \cos^{-1} \left[\frac{r_i^2 + (s-x)^2 - r_c^2}{2(s-x)r_i} \right] dx, \quad (6)$$

with the bounds of the integral being the same as those given hereinabove.

5

Discussion of the EXAMPLES

The outcome of the fiber-optic probe design strategy described herein comprises a fairly simple illumination and collection geometry that is capable of extracting the optical properties of a medium from the diffuse reflectance spectra with RMSEs of 0.41 cm⁻¹ (tested range of 0-80 cm⁻¹) and 0.30 cm⁻¹ (tested range of 3-40 cm⁻¹), for μ_a and μ_s' , respectively. Upon experimental validation of this algorithm using phantom studies, it was found that the algorithm did not perform as expected at high μ_a , with RMSEs of 2.4 cm⁻¹ (tested range of 0.35-23) and 3.7 cm⁻¹ (tested range of 5.2-26), for μ_a and μ_s' , respectively. It also was found that when the wavelength range was restricted to wavelengths greater than 450 nm, the experimental accuracy was similar to that of the simulation studies, with RMSEs of 0.4 cm⁻¹ (tested range of 0.35-3.2) and 2.2 cm⁻¹ (tested range of 5.2-25), for μ_a and μ_s' , respectively.

It was noted that the case where the illumination fiber was also used for collection produced somewhat lower accuracy than the case where it was not. This could be due to the fact that the Henyey-Greenstein phase function was used to describe light transport for a very small volume in the case of the overlapping probe geometry. This phase function allows one to fix only the first moment of anisotropy, however, it has been shown that higher order moments of anisotropy must be accounted for when scaling Monte Carlo simulations at short source-detector separations (Thueller *et al.* (2003) 8 *J Biomed Opt* 495-503), as

25

is the case when the source and collection fibers are overlapping. Incorporation of a more complex phase function could potentially improve this result. However, given that the separate collection fiber designs perform well, and the technical difficulties in eliminating specular reflection with a common source-
5 collection fiber, the separate collection fiber design might be a preferred solution.

The success of this method is dependent on the ability to construct a fiber probe meeting the design specifications produced by the optimization algorithm.

Errors in the positioning of the individual fibers introduced in the construction of this probe are thus a potential source of error. To minimize this, two design
10 constraints were introduced. First the probe geometry was limited to a simple design consisting of a single illumination fiber and a variable number of collection fibers. This ensured that the resulting probe would be easily manufactured. Second, the fibers were placed in direct contact with the tissue, with no shield or casing to allow easy access for visualizing the fiber tips. This enables one to
15 image the tip of the probe, and determine the exact positions of the individual fibers in the probe. In addition, it was found that for a reasonable range of positioning errors, the accuracy of this method in extracting optical properties was not adversely affected, provided the positioning errors were accounted for in this way.

20 In the optimal probe geometry, it was noted that one of the collection fibers (at 1.36 mm separation) was relatively insensitive to changes in the scattering properties of the medium. This fiber is similar in separation to that reported by Mourant *et al.* (1.7 mm; 36 *Appl Opt* 5655-5661, 1997), who found that at similar source-detector separations, the reflectance measured had a path
25 length that was insensitive to changes in scattering. This fiber thus has the useful property of providing a direct reflection of the absorption properties of the medium. In the optimal probe geometry reported here, this was combined with a shorter source-detector separation (380 μm). At this shorter source-detector separation, the reflectance collected by the probe was insensitive to changes in
30 μ_t . This fiber thus provides a direct reflection of the albedo of the medium.

Combining these two measurements enables a complete description of the optical properties of the medium.

Experimental validation of this approach yielded results that did not quite perform as well as the simulations had suggested. The primary source of this
5 discrepancy was identified as the large deviation of the Monte Carlo simulations from the diffuse reflectance measured at the larger source detector separation, having a maximum error of 39 fold for the larger wavelength range (400-616 nm), and 87% for the smaller wavelength range (450-616 nm).

There are two likely explanations for this effect. First, the signal collected
10 at high μ_a was extremely weak, particularly for the larger source detector separation, suggesting a limitation due to signal to noise issues. As a result, the signal collected may have been outside the linear dynamic range of the CCD detector. A wide range of integration times were used in an attempt to compensate for this, however there was still difficulty in obtaining sufficient signal
15 at the Soret band of hemoglobin absorption without saturating adjacent portions of the spectra for the highest concentrations of hemoglobin.

A second potential source of error is in the construction of the fiber optic probe. At the common end of this probe, a metal spacer was used between the collection fibers at small and large separations. This has the effect of making
20 the surface of the medium more reflective once photons extend beyond shorter source detector separation. Simonson *et al.* have reported that increasing the reflectivity of the probe between the source and collection fibers has the effect of reducing the mean probing depth of the collected photons (Simonson *et al.* (2006) 11 *J Biomed Opt* 014023). This would have the effect of shortening the
25 mean path length and reducing the effect of absorption. In the Monte Carlo simulations, the entire surface of the medium above the phantom was modeled as an optical fiber in order to enable the use of scaling relationships. Thus, the effect reported by Simonson *et al.* could influence the deviation observed between the measured and Monte Carlo simulation for the larger source detector
30 separation.

Certain experimental problems might need to be addressed before accurate retrieval of optical properties over the 400-450 nm wavelength range will be possible in tissue. However and as disclosed herein, the presently disclosed method showed reasonable accuracy for wavelengths longer than 450 nm, which has previously been demonstrated to be sufficient for the retrieval of hemoglobin concentrations (Finlay & Foster (2004) 31 *Med. Phys* 1949-59), and can also be used to characterize carotene absorption (Palmer *et al.* (2006) 45 *Appl Opt* 1072-1078). Thus, any possible inaccuracies in the range of 400-450 nm would not be expected to compromise the capability of this method for many biomedical applications where biological absorber concentrations need to be obtained.

A number of other studies have also investigated the extraction of optical properties from diffuse reflectance spectra (Thueler *et al.* (2003) 8 *J Biomed Opt* 495-503; Amelink *et al.* (2004) 29 *Opt Lett* 1087-1089; Palmer & Ramanujam (2006) 45 *Appl Opt* 1062-1071; Finlay & Foster (2004) 31 *Med Phys* 1949-1959; Zonios *et al.* (1999) 38 *Appl Opt* 6628-6637; Ghosh *et al.* (2001) 40 *Appl Opt* 176-184; Pfefer *et al.* (2003) 8 *J Biomed Opt* 206-215). These studies can be broken down into two general categories: (1) those using multiple fibers in the probe (six or more collection channels) to take advantage of the spatially resolved diffuse reflectance of a turbid medium to extract the optical properties (e.g., (Thueler *et al.* (2003) 8 *J Biomed Opt* 495-503; Ghosh *et al.* (2001) 40 *Appl Opt* 176-184; Pfefer *et al.* (2003) 8 *J Biomed Opt* 206-215), and (2) those studies that use relatively simple probe geometries (two or fewer independent collection channels), to extract the optical properties from the diffuse reflectance by making assumptions concerning the tissue constituents, including the absorbers and scatterers present in the tissue (e.g., Amelink *et al.* (2004) 29 *Opt Lett* 1087-1089; Palmer & Ramanujam (2006) 45 *Appl Opt* 1062-1071; Finlay & Foster (2004) 31 *Med Phys* 1949-1959; Zonios *et al.* (1999) 38 *Appl Opt* 6628-6637).

It can be desirable to use a diffuse reflectance model that minimizes the complexity of the probe geometry, in order to maintain cost-effectiveness, and minimize the overall dimensions of the fiber-optic probe such that it can be easily

adaptable to endoscopic measurements (as in case (2)). However, as demonstrated in Palmer *et al.* (2006) 45 *Appl Opt* 1072-1078, the requirement for *a priori* information about the tissue constituents can negatively impact the effectiveness of such diffuse reflectance models. In particular, an inaccurate
5 description of the absorption spectrum of beta-carotene was shown to have a negative impact on the quality of the fits to the breast tissue diffuse reflectance spectra obtained using an inverse Monte Carlo based model of diffuse reflectance (Palmer *et al.* (2006) 45 *Appl Opt* 1072-1078). The optimized approach employed in this study allows one to obtain the advantages of each
10 approach by obtaining accurate optical property information using a simple probe geometry while requiring no *a priori* information about the tissue constituents.

This paper focused on optimizing the probe geometry for a specific application: *i.e.*, reflectance-based extraction of optical properties for a semi-infinite medium. However, the methodology presented could be easily adapted
15 to a number of other applications. By modifying the objective function appropriately, this approach could be employed to optimize a probe to meet any quantifiable goal, such as limiting the probing depth, or extracting the optical properties from the top layer of a layered medium. Furthermore, alternative methods of extracting optical properties could be used, such as the diffusion
20 equation or Monte Carlo models instead of the neural network algorithm. These approaches would generally require that a nonlinear least squares fit be applied to fit the reflectance spectrum to the modeled spectrum given a particular set of optical properties (Palmer & Ramanujam (2006) 45 *Appl Opt* 1062-1071). This would have the advantage of using an established light transport model to relate
25 the diffuse reflectance to the underlying optical properties.

However, these alternative methods are iterative in nature and thus would be much slower (likely several orders of magnitude) in both the optimization and extraction phases. In addition, while the EXAMPLES focus on the UV-VIS spectral range, these methods disclosed herein would also be applicable to the
30 near infrared (NIR). In fact, the application to the UV-VIS is likely the more difficult task due to the wider range of optical properties present in biological

samples. Finally, this approach could also be applied to fluorescence spectroscopy for the extraction of intrinsic fluorescence properties. This approach could be used to determine the optimal fiber probe design for essentially any spectroscopic application.

5 Although degree of matching between produced optical properties with known optical properties for given diffuse reflectance was used as the optimization metric in the experiments described above, the subject matter described herein is not limited to using this optimization criterion. Other optical properties that can be used for optimization of probe design include sensing
10 depth of a probe geometry, sensing volume or spatial resolution of a probe geometry, measurement of physical properties of tissue, measurement of fluorescence properties of tissue, or measurement of any optical signal, such as but not limited to Raman scattering. In addition, the subject matter described herein is not limited to using a genetic algorithm to select the optimal probe
15 geometry. Any selection algorithm, such as the gradient methods described above, may be used to select the optimal probe geometry.

The disclosure of each of the publications referenced herein is hereby incorporated by reference in its entirety.

20 It will be understood that various details of the presently disclosed subject matter can be changed without departing from the scope of the presently disclosed subject matter. Furthermore, the foregoing description is for the purpose of illustration only, and not for the purpose of limitation.

CLAIMS

What is claimed is:

1. A method for optimizing a probe geometry for spectroscopic measurement in a turbid medium, the method comprising:
 - 5 (a) selecting a probe geometry comprising at least one emitting entity for emitting electromagnetic radiation into a turbid medium and at least one collecting entity for collecting the electromagnetic radiation that has interacted with the turbid medium;
 - 10 (b) performing a simulation with inputs of the probe geometry and a plurality of sets of optical property values associated with the turbid medium to generate output comprising optical parameter values measured by the probe geometry for each set of input optical property values;
 - 15 (c) providing the measured optical parameter values as input to an inversion algorithm and thereby producing corresponding optical properties as output;
 - 20 (d) comparing the produced optical properties with optical properties known to correspond to the measured optical parameter values and determining a degree of matching between the produced and known optical properties;
 - 25 (e) repeating steps (b)-(d) for a plurality of additional probe geometries, wherein each additional probe geometry differs from the probe geometry of step (a) in at least one property selected from the group consisting of a quantity of collecting entities, a diameter of at least one collecting entity, a linear distance between the emitting entity and the collecting entity, and combinations thereof, wherein repeating steps (b)-(d) comprises, at each iteration, applying an optimization algorithm to select a probe geometry such that the resulting degree of matching will converge
30 to an optimum value; and

- (f) selecting from among the different probe geometries, an optimal geometry based on the degree of matching determined for each geometry in step (d).
- 5 2. The method of claim 1, wherein the emitting entity is not a collecting entity.
3. The method of claim 1 wherein the emitting and collecting entities comprise optical fibers.
4. The method of claim 1, wherein the simulation employs a Monte Carlo model.
- 10 5. The method of claim 1, wherein the sets of optical properties include scattering coefficients (μ_s), absorption coefficients (μ_a), and anisotropy factors (g).
6. The method of claim 1, wherein the optical parameter values are selected from the group consisting of diffuse reflectance values, sensing depths of a probe geometry, sensing volume or spatial resolution of a probe geometry, measurements of a physical property of a tissue, measurements of a fluorescent property of a tissue, measurement of Raman scattering, and combinations thereof.
- 15 7. The method of claim 6, wherein the optical parameter values are diffuse reflectance values.
- 20 8. The method of claim 1, wherein each emitting and collecting entity of the probe has a diameter selected from the group consisting of 50, 100, 150, 200, 300, 400, and 500 μm .
9. The method of claim 1, wherein the turbid medium comprises a group of cells or a tissue in a subject or isolated from a subject.
- 25 10. The method of claim 9, wherein the group of cells or the tissue comprises a tumor or a tumor biopsy.
11. The method of claim 1, wherein the method simultaneously and/or consecutively optimizes the probe in terms of fiber diameter and distance from the emitting entity to each collecting entity.
- 30

12. The method of claim 1, wherein the inversion algorithm is executed by a neural network.
13. The method of claim 12 comprising training the neural network using a training set of optical parameter values and the corresponding known optical properties
- 5 14. The method of claim 1, wherein the optimization algorithm comprises a genetic algorithm.
15. The method of claim 14, wherein the genetic algorithm comprises a component selected from the group consisting of a crossover operator, a mutation operator, and combinations thereof.
- 10 16. The method of claim 1, wherein the probe geometry comprises from 1 to 5 collecting entities, inclusive.
17. The method of claim 16, wherein each of the plurality of additional probe geometries differs from the probe geometry of step (a) in at least one property selected from the group consisting of a quantity of collecting entities, a diameter of at least one emitting or collecting entity, a linear distance between the emitting and collecting entities, and combinations thereof.
- 15 18. The method of claim 1, wherein determining a degree of matching between the produced and known optical properties comprises:
- 20 (a) determining a root mean square error (RMSE) for a parameter selected from the group consisting of absorption coefficient (μ_a) and reduced scattering coefficient (μ_s' ; $= \mu_s \times (1-g)$, wherein μ_s is a scattering coefficient and g is an anisotropy factor) for each probe geometry; and
- 25 (b) choosing a probe geometry for which the RMSE for each of μ_a and μ_s' is less than or equal to 0.5 cm^{-1} .
19. The method of claim 18, wherein choosing a probe geometry includes choosing a probe geometry for which the RMSE for each of μ_a and μ_s' is less than or equal to 0.45 cm^{-1}
- 30

20. A probe comprising a probe geometry selected using the method of claim 1.
21. A probe for generating and collecting electromagnetic radiation that interacts with a turbid medium, the probe comprising:
- 5 (a) at least one emitting entity for emitting electromagnetic radiation into a turbid medium; and
- (b) at least two collecting the electromagnetic radiation emitted by the emitting entity that has interacted with the turbid medium,
- 10 wherein a linear distance between the emitting entity and each collecting entity does not exceed 1.5 millimeters and wherein the linear distance is determined using an optimization algorithm that optimizes measurement accuracy of the probe with regard to at least one optical property of a turbid medium.
22. The probe of claim 21, wherein the probe comprises between 2 and 6
15 collecting entities, inclusive.
23. The probe of claim 22, wherein the probe comprises between 2 and 5 collecting entities, inclusive, and the emitting entity is not also a collecting entity.
24. The probe of claim 21 wherein the emitting and collecting entities
20 comprise optical fibers.
25. The probe of claim 21, wherein:
- (i) the collecting entity comprises a first collection fiber and a second collection fiber;
- (ii) the emitting entity comprises an illumination fiber having a
25 diameter of about 500 μm , the first collection fiber has a diameter of about 200 μm , and the second collection fiber has a diameter of about 400 μm ; and
- (iii) a linear distance from the center of the illumination fiber to the center of the first collection fiber is about 380 μm and a
30 linear distance from the center of the illumination fiber to the center of the second collection fiber is about 1360 μm .

26. The probe of claim 25, wherein a linear distance from the center of the first collection fiber to the center of the second collection fiber is between about 980 μm and about 1740 μm .
27. The probe of claim 21, wherein the optimization algorithm includes a genetic algorithm usable to select a probe geometry such that a degree of matching between a value of the optical property measured by the fiber optic probe and a known value for the optical property will converge to an optimum value.
28. A system for selecting an optimal geometry for a probe for spectroscopic measurement in turbid media, the system comprising:
- (a) a light transport model for receiving as inputs a probe geometry and a plurality of sets of optical properties of a turbid medium and for producing as output optical parameter values that would be measured by the probe geometry for each set of input optical properties;
 - (b) an objective function for implementing an inversion algorithm for receiving as input the measured optical parameter values, for producing corresponding optical properties, for comparing the produced optical properties with optical properties known to correspond to the measured optical parameter values, and for determining a degree of matching between the produced and known optical properties for the given probe geometry, wherein the light transport model and the inversion algorithm are adapted to test a plurality of different probe geometries and wherein the inversion algorithm is adapted to determine a degree of matching between the produced and known optical properties for each geometry; and
 - (c) a probe selector for selecting one of the geometries as an optimal geometry based the degree of matching associated with the selected geometry.

29. The system of claim 28 wherein the probe selector is adapted to iteratively select new probe geometries to be tested by the light transport model and the inversion algorithm and, at each iteration, to apply an optimization algorithm to select a probe geometry such that the resulting degree of matching will converge to an optimum value.
- 5
30. The system of claim 29 wherein the optimization algorithm comprises a genetic algorithm.
31. A computer program product comprising computer executable instructions embodied in computer readable medium for performing steps comprising:
- 10
- (a) selecting a probe geometry comprising at least one emitting entity for emitting electromagnetic radiation into a turbid medium and at least one collecting entity for collecting the electromagnetic radiation that has interacted with the turbid medium;;
- 15
- (b) performing a simulation with inputs of the probe geometry and a plurality of sets of optical property values associated with the turbid medium to generate output comprising optical parameter values measured by the probe geometry for each set of input optical property values;
- 20
- (c) providing the measured optical parameter values as input to an inversion algorithm and thereby producing corresponding optical properties as output;
- (d) comparing the produced optical properties with optical properties known to correspond to the measured optical parameter values and determining a degree of matching between the produced and known optical properties;
- 25
- (e) repeating steps (b)-(d) for a plurality of additional probe geometries, wherein each additional probe geometry differs from the probe geometry of step (a) in at least one property selected from the group consisting of a quantity of collecting entities, a diameter of at least one emitting or collecting entity, a linear
- 30

distance between the emitting and collecting entities, and combinations thereof, wherein repeating steps (b)-(d) comprises, at each iteration, applying an optimization algorithm to select a probe geometry such that the resulting degree of matching will converge to an optimum value; and

5

- (f) selecting, from among the different probe geometries, an optimal geometry based on the degree of matching determined for each geometry in step (d).

10

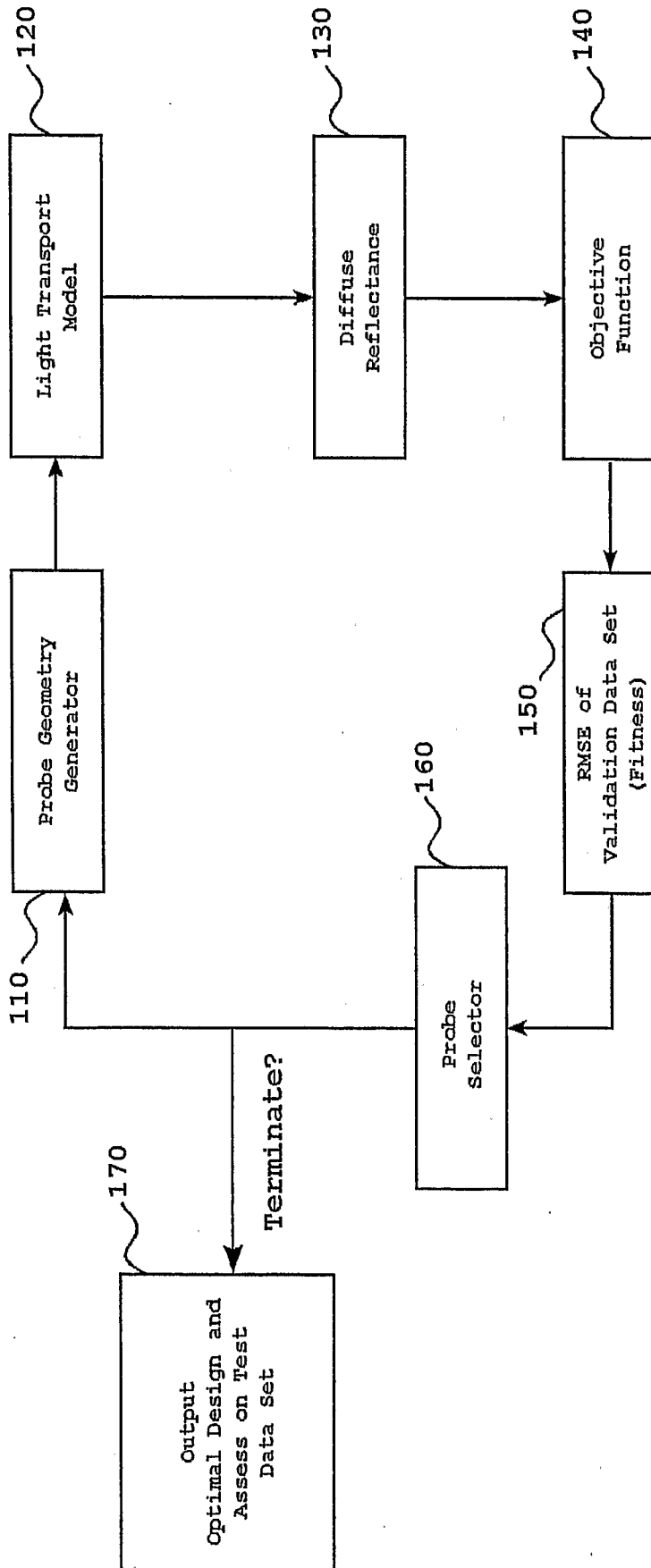


FIGURE 1

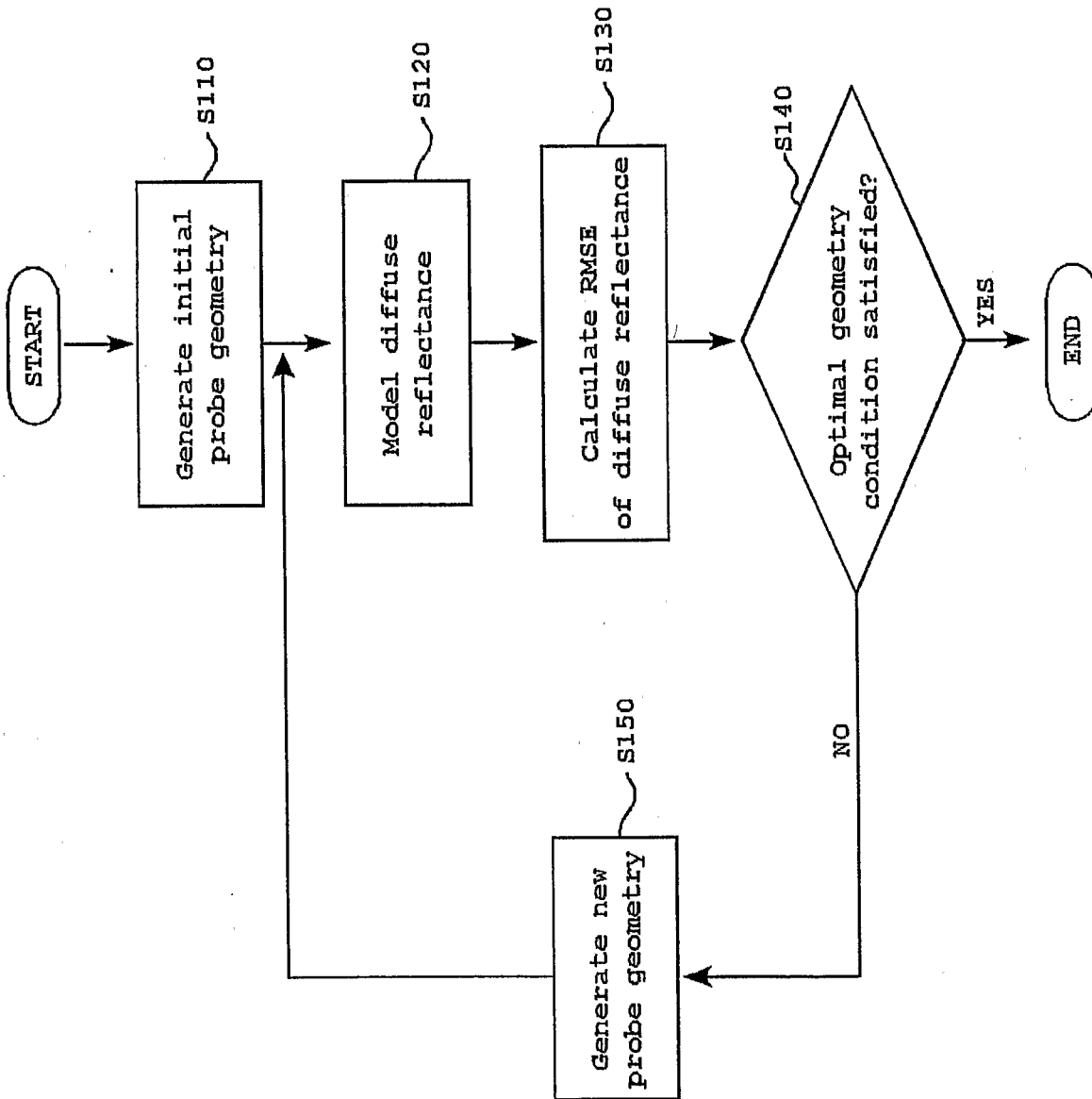


FIGURE 2

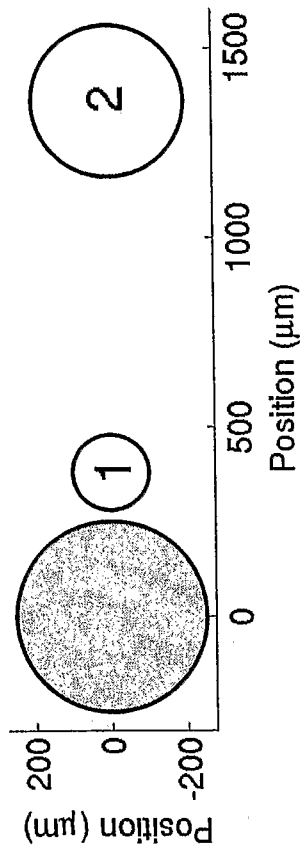


FIGURE 3

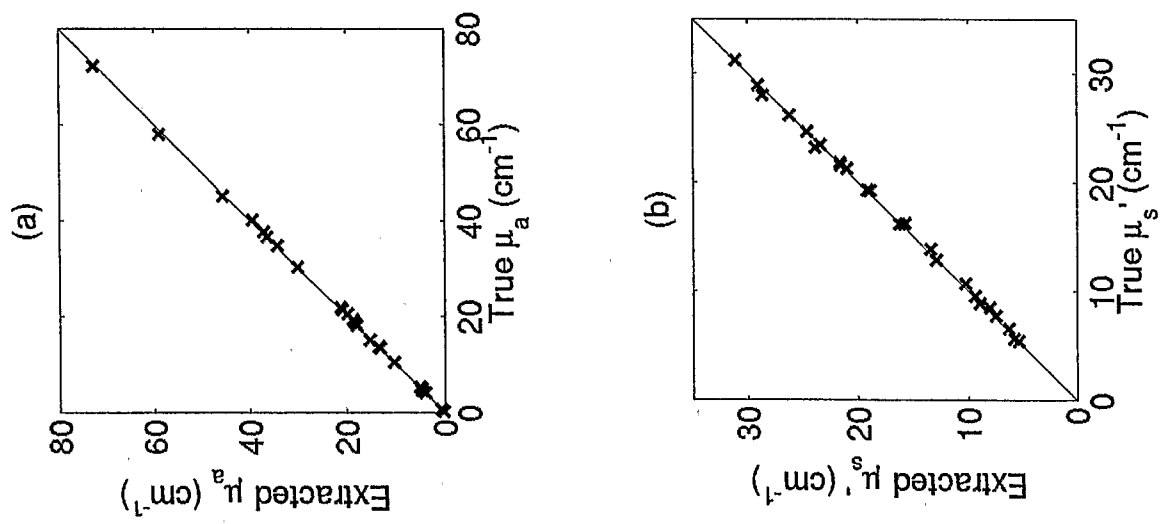


FIGURE 4

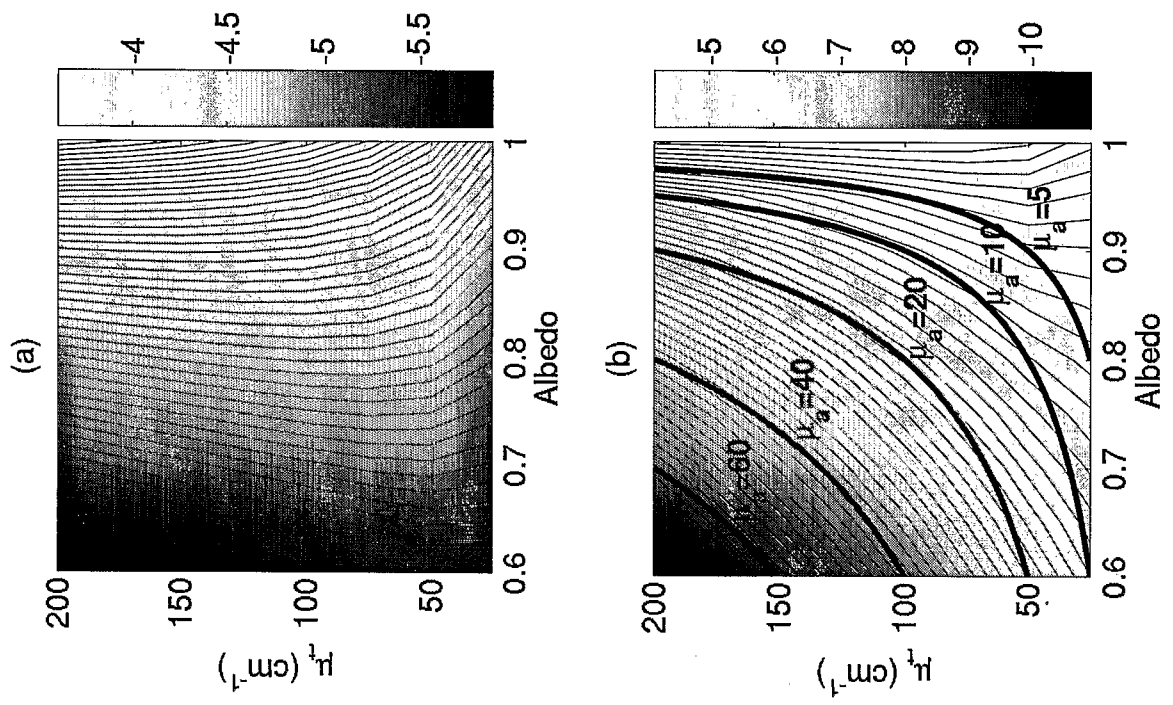


FIGURE 5

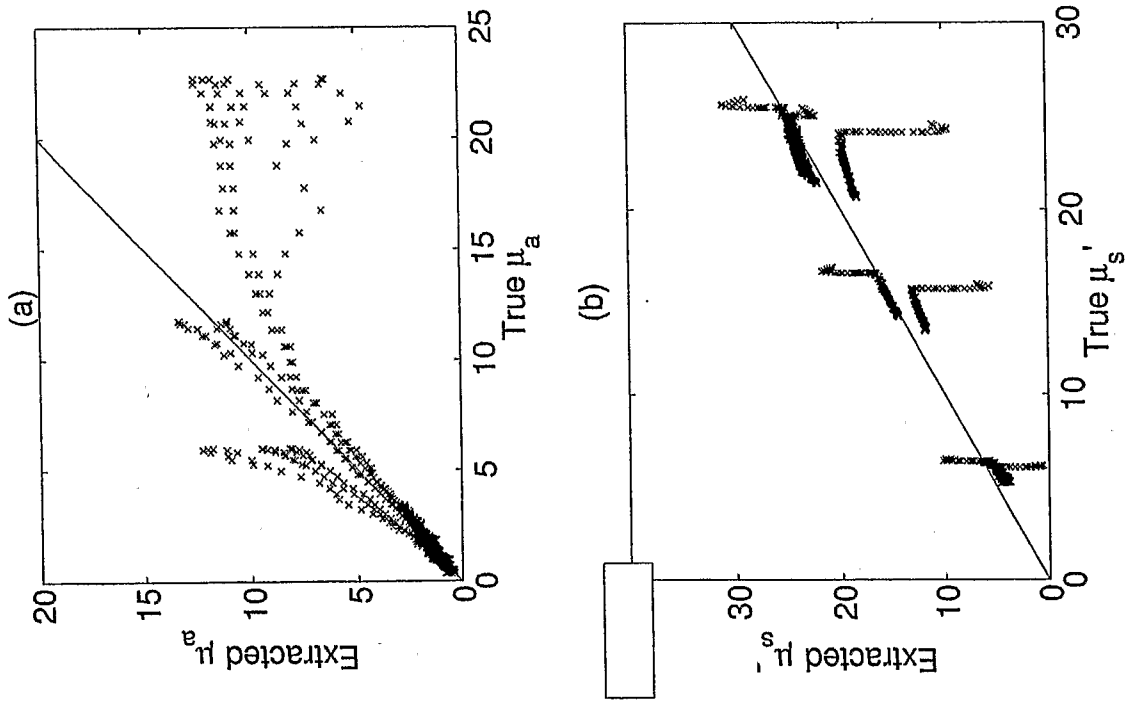


FIGURE 6

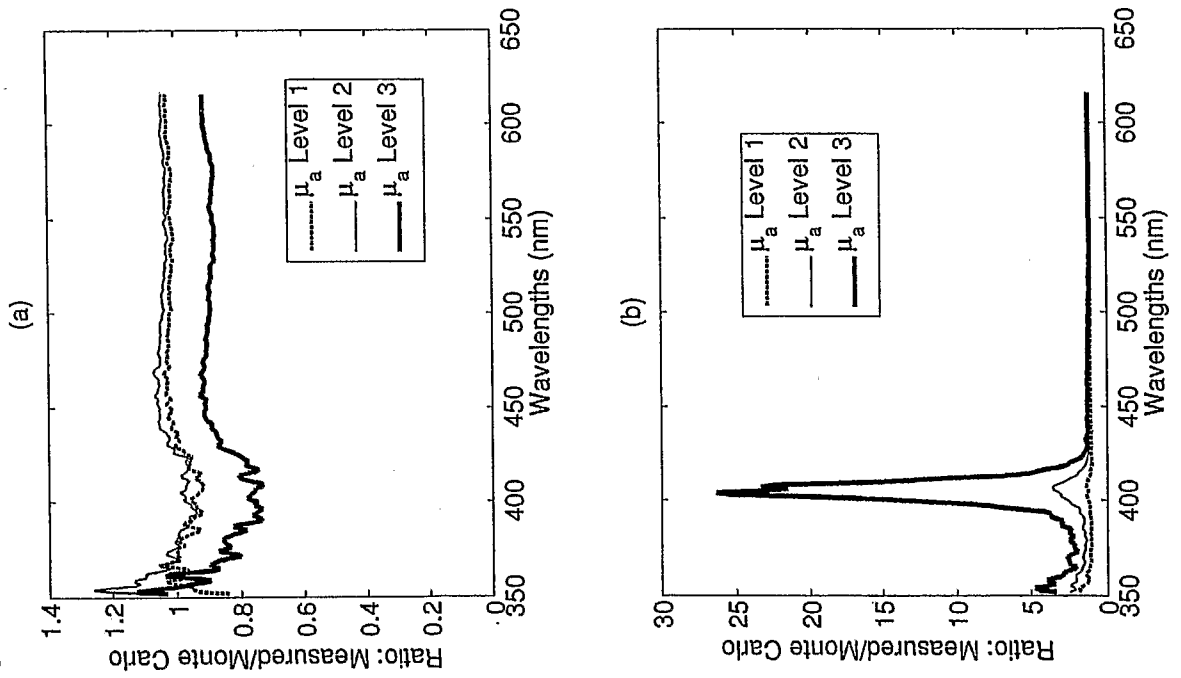


FIGURE 7

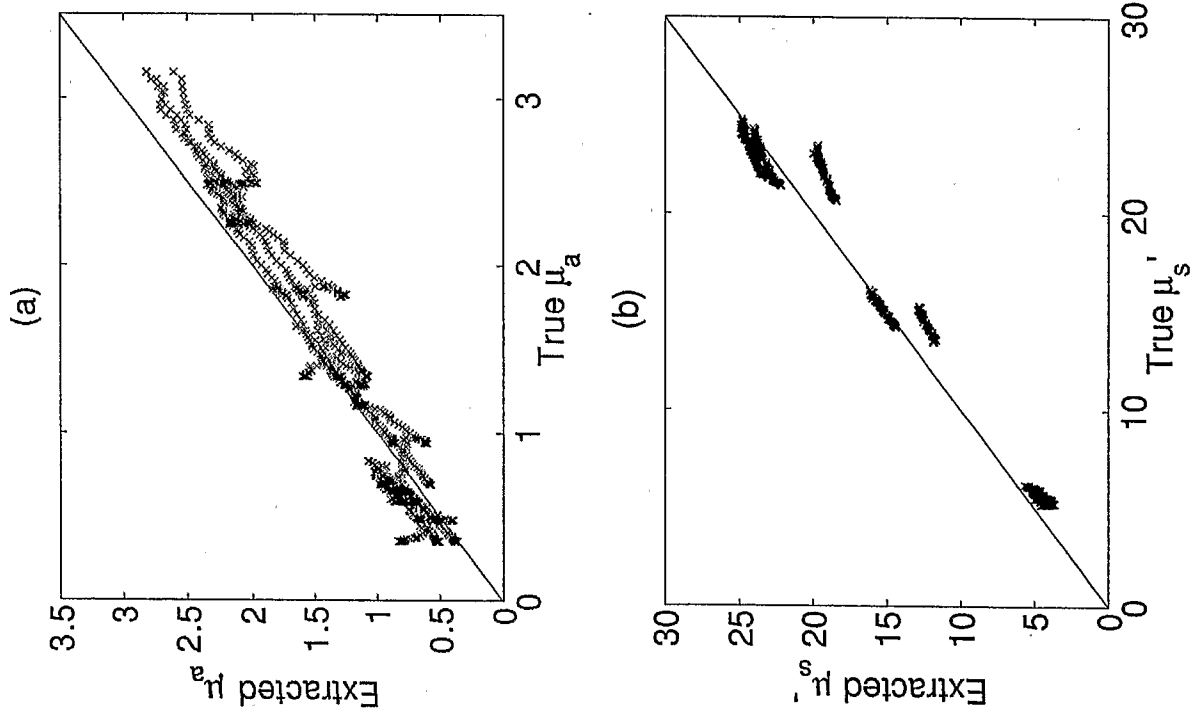


FIGURE 8