Spread spectrum time-resolved diffuse optical measurement system for enhanced sensitivity in detecting human brain activity

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Abstract. Diffuse optical spectroscopy (DOS) and imaging methods have been widely applied to noninvasive detection of brain activity. We have designed and implemented a low cost, portable, real-time one-channel time-resolved DOS system for neuroscience studies. Phantom experiments were carried out to test the performance of the system. We further conducted preliminary human experiments and demonstrated that enhanced sensitivity in detecting neural activity in the cortex could be achieved by the use of late arriving photons.

Keywords: diffuse optical imaging; functional imaging; spread spectrum.

1 Introduction

Diffuse optical spectroscopy (DOS) and diffuse optical tomography (DOT) can probe hemodynamic changes in large human organs due to the large penetration depth of near-infrared photons in biological tissues. Recently, DOS has become increasingly popular in neurosciences as it can provide real-time information about neurovascular coupling in the human brain. Moreover, it can be used alone or in combination with other functional imaging modalities, such as electroencephalogram and functional magnetic resonance imaging to investigate brain function.

While a variety of DOS systems have been reported, most of them fall into three categories: continuous-wave (CW), frequency-domain (FD), and time-domain (TD) methods. Among all the implementations, the CW method is the simplest technique: it is low cost, fast in data acquisition, and portable. The major drawback with this technique is that it measures the combined contributions from photons diffusing through superficial tissues and deep cerebral tissues. The systemic physiological changes in superficial layers (e.g., scalp and skull), due to heart beats and/or blood pressure fluctuation, may vary the intensity of early arriving photons. Such a variation could confound the optical measurements as the early arriving photons usually overwhelm the late arriving photons, which carry information about the deep layers (cortical tissues). As a result, the sensitivity in detecting the signal from the deep layers is limited for the CW method. To improve the depth sensitivity, CW domain systems usually resort to multistance measurements with a number of source-detector pairs.

It has been reported that with a single source-detector pair, a TD system can discriminate between the photons coming from deep regions and those from the superficial layers. Several groups have demonstrated that a TD system provides enhanced sensitivity to cortical hemodynamic changes due to its capability to detect time-resolved arrival of photons. Along with the improvement in depth resolved sensitivity, a time domain system has enhanced localization in detecting hemodynamic changes.

In conventional TD DOT systems, an ultrashort pulsed laser is used to illuminate the tissue sample and the optical signal from the sample is detected using either a streak camera or a time correlated single photon counter (TCSPC). The conventional time domain system has several limitations, such as small dynamic range, temporal nonlinearity in the case of streak camera-based detection and long data acquisition time and sensitivity to ambient environment in the case of the TCSPC-based detection. The conventional TD systems are usually bulky, slow, expensive, complicated, and less scalable to increase the number of source detector pairs.

To overcome these issues, a time domain DOT system based on the spread spectrum method was proposed. Spread spectrum techniques are used in communication systems for secure and high fidelity communication and have many advantages such as low-error rate, interference rejection, and selective addressing capability. In a spread spectrum method, a communication signal with a particular bandwidth is deliberately spread in the frequency domain to transform it into a wide-band signal. These techniques require the use of pseudorandom number sequences (PRBS) to determine and control the spreading behavior of the signal along the bandwidth. Using this technique, it is possible to use low cost laser diodes as sources and avalanche photodiodes (APDs) as detectors. A variable delay line is combined with an electronic correlator in the detector side to demodulate the optoelectric signal and generate the time resolved profile of arriving photons. Previously, a spread spectrum method-based system was reported for mapping the optical properties of breast tissues.

In this paper, we report a single-channel DOS system for brain functional tests. It is a time-resolved instrument based on our spread-spectrum method. However, new design elements have been included to further reduce the system complexity and

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system cost (<$5000). We further demonstrate that such a low-cost system can provide the high performance needed for in vivo monitoring of human brain activity.

2 Materials and Methods

2.1 System Design

A previously reported spread-spectrum DOT system used a commercially available PRBS generator to generate the spread-spectrum signals. However, commercial PRBS devices are usually bulky and expensive (>$10,000) as they usually come with additional and unnecessary functions. A key improvement in our design is a field programmable gate array (FPGA)-based dual-channel PRBS generator. An FPGA evaluation board (SP605, Xilinx) is chosen to implement two PRBS generators in the same device. The two generators are structurally identical. Each of them includes a 10-stage linear feedback shift register, which abides by a particular sequence generating polynomial \( X^{10} + X^7 + 1 \). A combinational logic circuit is used to predict the subsequent 20 bits within one internal clock cycle (internal clock frequency: 125 MHz). These bits are serialized by the onboard serializer/deserializer (ser/des) to the I/O transceiver port. The serial output is a 2.5 Gbps PRBS (pattern 10) that repeats itself for every \( 2^{10} - 1 \) bits. The output can be seeded with different initial states to form two independent generators, so that their corresponding output sequences are identical but time shifted by an integer multiplication of 400 ps. The minimum delay step size is 400 ps that is limited by the clock rate of FPGA board (2.5 GHz). The dual-channel PRBS generator eliminates the need for a programmable delay line, which costs another few thousand dollars.

As shown in the system schematic (Fig. 1), the dual-channel PRBS generator is controlled by a National Instruments data acquisition (NI-DAQ) device for variable time delay. One of the output PRBS sequences is modulated by a 2-kHz square wave, which functions as the secondary modulation signal. Previously, we used an external Mach–Zehnder modulator to achieve modulated illumination. However, such a fiber-based external modulator is subject to the photorefractive effect, not to mention its high cost (around $6000). In this design, we chose an 850-nm high-speed vertical cavity surface emitting laser (VCSEL) diode (VIXAR Inc, MULTM-000-B002) as the light source, which could be direct intensity modulated with the modulation signal. The VCSEL is fiber coupled to a head mount for brain activity monitoring. A light guide (Edmund optics, 39-366) is also placed on the head mount with a fixed distance from the source fiber. It collects the diffuse reflectance from the brain or a tissue phantom. The collected light is coupled into a thermoelectrically cooled avalanche photodiode (TE-APD, based on Hamamatsu S2383), which converts the weak optical signal into an electrical signal. The optoelectrical signal is preamplified by low-noise amplifiers (not shown in Fig. 1) before demodulation. The first stage demodulation is done by mixing the preamplified optoelectrical signal with the time-delayed PRBS sequence. The down-converted signal is the time-resolved (for a specific time delay) diffuse light intensity carried by the 2 kHz square wave. It is digitized by the NI-DAQ device so that the second-stage demodulation can be carried out numerically on a PC. The sampling rate for the signal acquisition is 50 Ks/s. The integration time to record the raw signal for one specific time delay is set to 25 ms. A LABVIEW program on the PC is responsible for scanning the delay time, collecting the raw data, performing secondary demodulation, and recording the time-resolved diffuse light intensity, the so-called temporal point spread function (TPSF).

2.2 Tissue Phantom and In-Vivo Imaging

To characterize the performance of the system, a homogeneous tissue phantom was made by diluting 20% Intralipid in a water tank. The diluted solution was 1% in concentration. The true value of reduced scattering coefficient (\( \mu_s' \)) at this concentration was around 8.9 cm\(^{-1}\). Ink solution was added into the diluted Intralipid solution to control its absorption coefficient (\( \mu_a \)) values. A spectrometer (Ocean Optics, USB4000-VIS-NIR) was used to quantify the true value of the absorption coefficient (\( \mu_a \)) for various ink concentrations. For homogeneous phantom experiments, three concentrations of ink were used (0.01, 0.02, and 0.03 ml/l). The respective values of absorption coefficient for these concentrations were: 0.08, 0.15, and 0.31 cm\(^{-1}\). A Mie calculator was used to estimate the theoretical value of the reduced scattering coefficient of the Intralipid solution. The system was first tested for its accuracy in quantifying the absorption and scattering coefficients of the homogeneous sample with a source-detector distance of 2.5 cm. A background optical property fitting method was used to simultaneously estimate absorption and scattering coefficients of the homogeneous phantoms. Compared with continuous wave and frequency domain systems, a conventional time domain system provides enhanced sensitivity to inhomogeneities located at deeper regions using late arriving photons. To demonstrate that our low-cost system can provide similar sensitivity enhancement, it was characterized with experiments in which an absorbing target was embedded in a homogeneous Intralipid solution (\( \mu_s = 0.08 \) cm\(^{-1}\) and \( \mu_a = 8.9 \) cm\(^{-1}\)). In these experiments, the source and the detector were kept at a distance of 2.5 cm. The source fiber and the detection light guide gently touched the surface of the solution. The absorbing target was a cylindrical tube with a high absorption coefficient of around 0.64 cm\(^{-1}\) and a reduced scattering coefficient identical to that of the background medium. It was embedded in the tissue phantom with its center located at different depths (1 to 2.5 cm) and equally distant from both the source and the detector. The tube was 0.5 cm in diameter and 0.5 cm in length. To quantify the contrast enhancement, the perturbation as a function of time of flight was defined as.
Perturbation \((t, \text{time of flight}) = 100 \times \frac{\text{TPSF}_{\text{inhomo}}(\text{depth}, \text{time of flight}) - \text{TPSF}_{\text{homo}}(\text{time of flight})}{\text{TPSF}_{\text{homo}}(\text{time of flight})} \tag{1}\)

where \(\text{TPSF}_{\text{inhomo}}(\text{depth}, \text{time of flight})\) and \(\text{TPSF}_{\text{homo}}(\text{time of flight})\) were temporal profiles of diffuse photons measured with and without the embedded target, respectively.

In addition to the phantom experiments, we conducted a cognitive experiment on a healthy human subject to demonstrate applicability of our approach to brain functional studies. The research protocol was approved by the National University of Singapore Institutional Review Board. The experiment was designed to record the hemodynamic changes in the middle frontal cortex about 3.5 cm above the left pupil while the participant solved simple arithmetic problems in a short preset duration. A computer program was used to generate random four digit subtraction problems, e.g., “7654 – 3976 = ?.” Each cognitive test run was divided into 10 blocks. Each block was 3 min in total, consisting of 1 min of rest + 1 min of problem solving + 1 min of rest. The subject was instructed to solve as many problems as he could in 1 min. A timer was set in the background and the computer screen became blank with a white spot when 1 min of problem solving was over. A healthy right-handed male subject in his twenties was recruited for this experiment. The recorded raw data were averaged over blocks to increase the signal-to-noise ratio.

In the brain measurement experiment, the source-detector distance was 3 cm. The intensities of diffusive photons were measured at 11 time delay points (at a 400-ps interval). The photon intensities averaged over the rest period were considered as base values \([\text{TPSF}_{\text{m}}(\text{time of flight})]\). The changes in optical signal relative to the base values were computed as perturbations.

\[\text{Perturbation} \quad (t, \text{time of flight}) = 100 \times \frac{\text{TPSF}_{t}(\text{time of flight}) - \text{TPSF}_{m}(\text{time of flight})}{\text{TPSF}_{m}(\text{time of flight})}. \tag{2}\]

2.3 Numerical Simulation

To independently validate our time-resolved measurement results for perturbation quantification, finite element numerical simulations were carried out to theoretically predict the time domain responses. The simulation was based on a coefficient form partial differential equation

\[
\frac{n}{c} \frac{\partial \varnothing}{\partial t} + \nabla (-D \nabla \varnothing) + \mu_s \varnothing = P_0(r, t), \tag{3}\]

where \(\varnothing\) is the light fluence rate, \(D\) is the diffusion coefficient, \(\mu_s\) is the absorption coefficient, \(n\) is the refractive index, \(c\) is the speed of light in vacuum, and \(P_0(r, t)\) is the source distribution. The diffusion coefficient is defined as

\[D = \frac{1}{3(\mu_s' + \mu_a)}. \tag{4}\]

where \(\mu_s' = (1 - g)\mu_s\) is the reduced scattering coefficient, \(\mu_s\) and \(g\) are the scattering coefficient and anisotropy factor of the medium, respectively. A point source with a Gaussian temporal profile \([1 \text{ ns full width half maximum (FWHM)}]\) was used to excite the sample. The simulation geometrical configuration is shown in Fig. 2. Like the tissue phantom experiment, a source detector pair separated by 2.5 cm were used. The background optical properties used in simulations were identical to those in the phantom experiments \((\mu_s = 0.08 \text{ cm}^{-1} \text{ and } \mu_s' = 8.9 \text{ cm}^{-1})\). The point source was located at one mean free path distance \((1/\mu_s')\) inside the medium. A 15 x 15 x 15 cm³ cube was used to define the tissue phantom geometry. At the bottom and side surfaces, zero rate of change in fluence was used as the boundary condition to simulate a semi-infinite medium. On the top surface (Intralipid air interface) where the source and detector were located, the robin boundary condition was applied.

![Fig. 2](image-url) Two-dimensional cross section of finite element simulation geometry.

![Fig. 3](image-url) (a) Measured IRF from a white diffusive reflecting surface; (b) IRF peak intensity measured over a period of 1 h.
A cylindrical bead was included in the background medium as an inhomogeneity. The bead had a diameter of 0.5 cm and a length of 0.5 cm. The absorption coefficient for the bead was 0.64 cm$^{-1}$ and the reduced scattering coefficient of the bead was the same as that of the background medium. The bead was located at a depth varying from 1 to 2.5 cm.

3 Results and Discussion

To quantify the system stability and the system impulse response function (IRF), a TPSF was measured by placing a white reflecting surface at 30 cm away from the source-detector pair. Figure 3(a) shows an IRF measured from the reflecting surface. The FWHM of the IRF was around 1 ns. The theoretical limit of time resolution should be 400 ps given a PRBS bit rate of 2.5 Gbps. However, various components used in the system limited the overall system bandwidth. For example, the APD detector had a bandwidth of 600 MHz, which was the main bottleneck for a better time resolution. To demonstrate the long-term system stability, we repeated the above-mentioned IRF measurement at intervals of 1 min over a period of 1 h. The fluctuation in the peak value (normalized to the 1-h time average) of the IRF was <1% [Fig. 3(b)].

To evaluate the performance of our system in quantifying the optical properties of a homogeneous medium, the system was tested on homogeneous phantoms with different absorption coefficients (0.08, 0.15, and 0.31 cm$^{-1}$), but with the same reduced scattering coefficient (8.9 cm$^{-1}$). Figure 4(a) compares the IRF and TPSF’s. The IRF of the system was normalized to its peak value and the TPSF’s for different absorption coefficients were normalized to the maximum value of the TPSF measured from the phantom of a low absorption coefficient of 0.08 cm$^{-1}$. In Fig. 4(b), the true values of the absorption coefficients are compared with the absorption coefficients obtained by fitting the experimental data. The true values of the absorption coefficient were measured using the Ocean Optics spectrometer. Figure 4(c) shows the reduced scattering coefficient values obtained for all three homogeneous tissue phantoms.

The main advantage of time domain diffuse optical measurement is the enhanced sensitivity to deep perturbation in comparison with CW and FD methods. Our experimental results from inhomogeneous phantoms were validated against numerical simulation results. Figure 5(a) shows a simulated TPSF excited by a pulse 1 ns in duration (FWHM). Figure 5(b) shows an experimental TPSF measured from a homogeneous phantom of the same optical properties. The mean value (solid line) and standard deviations (error bars) were estimated from 20 repeated measurements. The TPSF was perturbed once the absorber was embedded in the otherwise homogeneous background. Figure 5(c) shows the perturbation predicted by numerical simulation and Fig. 5(d) shows the experimental perturbation levels for various times of flight of diffusive photons. For both simulations and experiments, the inhomogeneity was placed at various depths in the tissue phantom. The time of flight was mapped from the PRBS delay time after system calibration. Both experimental and simulation results show that, when the absorber was located in the shallow region (1 cm), the perturbation changed at a rather slow rate with an increasing time of flight. In contrast, the rate of change in perturbation increased significantly for late arriving photons when the absorber was moved to a deeper...
region (2.5 cm). Shown in Figs. 5(e) and 5(f) are perturbations as functions of bead depth based on simulation and experimental data, respectively. These results clearly demonstrated that late arriving photons captured by our system provided higher perturbation and therefore higher sensitivity when the absorber was located at a deep location. The phantom experimental results serve as direct evidence that our single-channel time resolved DOS system is capable of detecting heterogeneities deep inside a turbid medium with improved sensitivity.

Similarly, the human subject experiment also demonstrated the capability of our system in providing enhanced contrast using the late arriving photons. Figure 6(a) shows the measured TPSF from the human subject during the rest period. The Fig. 6(b) shows the block-averaged time traces of the diffuse light intensities, which reflects the underlying hemodynamic changes in the frontal cortex. Five traces are associated with five different times of flight, i.e., 2 ns (blue dot), 2.4 ns (black dot), 2.8 ns (black asterisk), 3.2 ns (red dot), and 3.6 ns (red asterisk), respectively. It is clear that late arriving photons show high perturbation values compared to the early arriving photons.

To conclude, we have successfully developed a low-cost, single-channel DOS system for brain functional monitoring and studies. We have demonstrated experimentally that this system can provide enhanced sensitivity in detecting human brain activities.
Fig. 6 (a) Measured TPSF (normalized) for a human subject during the rest period; (b) effect of time of flight on optical signals associated with hemodynamic changes. Time zeros corresponds to the onset of the problem solving phase for the cognitive task. Legend indicates the time of flight.

Disclosures

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