Time-resolved optical imager for assessment of cerebral oxygenation

Michal Kacprzak
Adam Liebert
Piotr Sawosz
Norbert Zolek
Roman Maniewski
Institute of Biocybernetics and Biomedical Engineering
Trojdena 4
02-109 Warsaw, Poland
E-mail: michal.kacprzak@ibib.waw.pl

Abstract. A time-resolved optical instrument allowing for noninvasive assessment of cerebral oxygenation is presented. The instrument is equipped with picosecond diode lasers, fast photodetectors, and time-correlated single photon counting electronics. This technology enables depth-resolved estimation of changes in absorption and, in consequence, assessment of changes in hemoglobin concentrations in the brain cortex. Changes in oxyhemoglobin (HbO2) and deoxyhemoglobin (Hb) can be evaluated selectively in extra- and intracerebral tissue compartments using the moments of distributions of times of flight of photons measured at two wavelengths in the near-infrared region. The combination of the data acquired from multiple sources and detectors located on the surface of the head with the depth-resolved analysis, based on the moments, enables imaging of cortex oxygenation. Results of the tests on physical phantoms as well as in vivo validation of the instrument during the motor stimulation experiment are presented. © 2007 Society of Photo-Optical Instrumentation Engineers.

Keywords: imaging systems; time-resolved imaging; tissue light propagation.

1 Introduction

Absorption of light in human tissues is relatively low in the near-infrared wavelength region. Thus, light from this so-called “biological optical window” can penetrate even several centimeters into the tissue. However, high scattering at these wavelengths causes optical transmission/reflectance measurements to have a limited spatial resolution. On the other hand, different spectral properties of oxy- and deoxyhemoglobin measurements of the tissue absorption at several wavelengths allow for estimation of tissue oxygenation.1,2 This functional information may be important for clinical assessment of oxygenation of the brain,3,4 muscles,5,6 or for detection of breast tumors.7

Recently it has been shown that the spatial resolution of near-infrared spectroscopy (NIRS) can be improved by application of modern measurement techniques. Frequency- and time-domain techniques were developed and applied in many NIRS studies. These methods are based on the emission of intensity modulated light or picosecond light pulses into the tissue. The specific advantage offered by these techniques is that they allow for direct measurement of mean photon pathlength and the estimation of absolute concentrations of chromophores. In the frequency-domain technique, analysis of the phase shift between the initial wave and the light detected on the surface of the tissue is carried out.8 The phase shift is directly related to the mean time of flight of photons. In time-resolved near infrared spectroscopy (TRNIRS), emission of short (picosecond) light pulses and the analysis of the broadening of the pulse during its travel between the points of emission and detection is performed.9,10

NIRS has been successfully applied in the analysis of brain oxygenation in neonates and infants.11,12 In such cases the influence of the extracerebral tissue (skin, skull) on the signals measured is small, and it can be neglected. Because of the small size of the head, even the tomographic reconstruction of absorption and the reduced scattering coefficient in the brain tissue can be obtained.13–16 Application of frequency- or time-resolved NIRS is beneficial, especially when the head of an adult is examined and a depth-resolved analysis of the hemoglobin concentration changes must be applied to estimate oxygenation of the brain tissue with higher accuracy. Recently, the time-domain method has been shown to allow for evaluation of changes of the optical properties, as well as changes of oxy- and deoxyhemoglobin concentrations with depth discrimination.17,18

Measurements of brain oxygenation changes caused by neuronal stimulus are often challenging because of the difficulties in determining the location of the focal point of stimulation on the surface of the cortex. In particular, proper positioning of the optode on the surface of the head is important when oxygenation changes are observed in neurophysiological experiments, i.e., during motor or visual stimulations. The NIRS technique has been extensively tested in monitoring changes of brain oxygenation in acute stroke,19 during hypothermic heart surgery,20 and in carotid artery endarterectomy.21 Moreover, NIRS has been applied to estimate brain perfusion by analysis of the inflow and the washout of the optical contrast agent.22–24 In all these applications,

a large number of the emission and detection points located on the surface of the head can be applied. In consequence, the changes of brain oxygenation and/or perfusion can be imaged, and the ischemic areas can be localized. Several optical systems allowing image changes of brain oxygenation have been proposed. Continuous wave,25–27 frequency-domain,28,29 and time-domain30–34 systems have been reported.

We present a time-resolved optical instrument that allows for noninvasive assessment of oxygenation of the brain cortex. The depth-resolved changes in oxy- and deoxyhemoglobin levels can be calculated using the moments of distributions of times of flight of photons (DTOFs) measured at two wavelengths in the near-infrared region. The instrument is equipped with picosecond diode lasers, fast photodetectors, and time-correlated single photon counting electronics. The combination of data from multiple sources that are switched sequentially and detectors distributed on the surface of the head, with the depth-resolved analysis based on the moments of DTOFs, enables imaging of cortex oxygenation.

2 Methods

2.1 Instrumentation

The TRNIRS system we developed is shown in Fig. 1. It consists of two semiconductor diode lasers and eight detection channels with time-correlated single photon counting electronics.

A laser diode driver (PDL 808 Sepia, PicoQuant, Berlin, Germany) and two semiconductor laser heads operating at wavelengths of 832 and 687 nm were used. Light pulses from both diodes were generated with the repetition rate of 80 MHz. The light pulse trains from both diodes were shifted in time by half of the repetition period. This temporal shift allowed us to analyze distributions of times of flight of photons at both wavelengths. To introduce laser light from the two laser heads into two optical fibers, an adjustable optical setup with a 50:50 glass beam splitter (Thin Film Imaging Technologies, Incorporated, Greenfield, Massachusetts) was constructed. After splitting the beams from the two lasers, the light pulses are delivered sequentially to 18 locations on the head (nine per each hemisphere) by using two optomechanical switches (Piezosystem jena, GmbH, Jena, Germany). The switches are controlled with the PCI board NI-DAQ 6221 (National Instruments, Austin, Texas) at a switching frequency of 10 Hz. The switching time is less than 5 ms. The tips of the output fibers from the switches are positioned in a 3×3 array on each hemisphere, and they are located directly on the surface of the head, together with the detecting fiber bundles. An optode holder was constructed using a thermo-plastic material (WFR/Aquaplast Corporation, Wyckoff, New Jersey), soft rubber, and Velcro strips [Figs. 2(a) and 2(b)] to fix the source fibers and the detecting bundles. For every source-detector pair, the interoptode distance is 3 cm. Different sizes of the optode setups allowed us to fix the fibers and bundles on heads of different sizes.

The diffusely reflected light was delivered to the detectors using 1.5-m-long fiber bundles with a diameter of 4 mm and NA=0.54 (Loptek, Berlin, Germany). Eight photomultiplier tubes (R7400U-02, Hamamatsu Photonics, Japan) were used for detection of the diffusely reflected light. Each photomultiplier was located separately in the electrically shielded box, together with the high voltage unit and preamplifier. The power of the light delivered to photomultipliers was adjusted by the neutral density filters located between the fiber bundles and the detectors. Independent PCI boards with time-correlated single photon counting electronics (SPC-134, Becker and Hickl, Berlin, Germany) were used for acquisition of distribution of times of flight of photons. Recording of DTOFs was triggered by the 10-Hz generator, which synchronously controlled the optomechanical switches. To avoid data

Fig. 1 Time-resolved NIR topography system for cerebral oxygenation imaging. SEPIA is the laser diode driver, LH1 and LH2 are lasers heads, SW1 and SW2 are 1:9 optical switches, PMT is the photomultiplier tubes, AMP is the preamplifiers and TCSPC is the PCI boards for time-correlated single photon counting.
acquisition during the switching period, the trigger signal 
from the generator was delayed by 5 ms.

The software for acquisition of distribution of times of 
flight of photons was developed in the LabView v.7 environ-
ment (National Instruments).

The instrumental response function was measured by pos-
tioning the source fibers in front of the detecting bundles. 
The detecting bundles were covered with a sheet of white 
paper to fill up the whole numerical aperture of the bundle.  
The instrumental response function as measured for every 
source-detector pair at both wavelengths was not longer than 
800 ps [full width at half maximum (FWHM)]. The measure-
ment system in the presented configuration allows us to 
record distributions of times of flight of photons for 32 
source-detector pairs corresponding to 32 “banana” volumes, 
forming a 4 × 4 array on each hemisphere. The acquisition of 
DTOFs starts after the switching period is finished, and for 
every location of the source it is completed within 95 ms. 
Parallel acquisition of the two 4 × 4 spot maps takes about 
0.9 s.

2.2 Data Analysis

For analysis of the measured data, algorithms based on a 
mathematical model of the light transport in the turbid media  
and a method of depth discrimination using moments of the 
DTOFs 18 were implemented. To separate information 
from the extra- and intracerebral layers of the head, so-called 
sensitivity factors were computed from the model of light 
propagation in the tissue. These sensitivity factors bind the 
calculated statistical moments of DTOFs with the changes in 
the absorption coefficient at different depths in the tissue un-
der investigation. 17,18,37

The phenomena of light propagation in the turbid medium 
can be described by the time-dependent diffusion 
equation. 38,39 The photon fluence rate \( \Phi(r, t) \) as a function of 
time \( t \) and position in the tissue described by vector \( r \) depends 
on the optical properties of the medium:

\[
\frac{1}{c} \frac{\partial}{\partial t} \Phi(r, t) - D \nabla^2 \Phi(r, t) + \mu_a \Phi(r, t) = S(r, t),
\]

where \( S(r, t) \) describes the photon source, \( c \) is the velocity of 
the light in the medium, \( D \) is the diffusion coefficient \( D = (3 \mu_r^s)^{-1} \), \( \mu_a \) is the absorption coefficient of the medium, and 
\( \mu_r^s \) is the reduced scattering coefficient. 40,41

By solving this equation for the semi-infinite, homogenous 
medium, assuming a delta Dirac light source pulse, diffuse 
reflectance \( R_d(p, t) \) can be derived as a function of time \( t \) and 
distance \( \rho \) between the source and the detector 38:

\[
R_d(p, t) = (\mu_r^s)^{-1}(4\pi Dc)^{-3/2}r^{-5/2} \exp\left(\frac{-\rho^2}{4Dc - \mu_dr^2}\right).
\]

Equation (2) describes light propagation in the medium with 
absorption and the diffusion coefficient distributed homog-
enuously over the whole volume. When a change of absorption 
coefficient \( \Delta \mu_a \) appears in a small inclusion of a sampling 
volume \( dV_s \) inside the investigated medium in a location 
defined by vector \( r \), the time-dependent change of diffuse 
reflectance \( \Delta R(r, p, t) \) on the surface of the medium at source-
detector separation \( \rho \) can be calculated:

\[
\Delta R(r, p, t) = -\Delta \mu_a dV_s[\Phi(r, t) \otimes E(r, p, t)],
\]

where \( \otimes \) means operation of convolution. Function \( \Phi \) 
describes the time-dependent fluence rate of photons in the 
medium at the location \( r \). \( E \) is the escape function, which 
describes the probability that the photon emitted from location \( r \) 
will reach the detector at distance \( \rho \) from the source position. 36

Distributions of times of flight of photons \( R \) diffusely re-
lected from the medium with the inclusion located at \( r \) can be 
then calculated using the formula:

\[
R(r, p, t) = R_d(p, t) + \Delta R(r, p, t).
\]

Normalized moment of order \( k \) of the distribution \( R(t) \) is 
defined by:

\[
m_k = \int_0^\infty t^k R(t) dt / \int_0^\infty R(t) dt.
\]

Changes in these moments of theoretical DTOFs caused by 
local changes of absorption \( \Delta \mu_a \) in a sampling volume \( dV_s \) 
located at \( r \) can be calculated using the formulas 18:

\[
\Delta A = -\log \left( \frac{N_{tot}}{N_{toth}} \right),
\]

\[
\Delta(t) = \langle t \rangle - \langle t \rangle_{0},
\]
\[ \Delta V = V - V_b, \]  
(8)

where \( \Delta A \) is the change in attenuation, \( N_{tot} = \int_0^\infty R(t)dt \) is the total number of photons, \( \langle t \rangle = m_1 \) is the mean time of flight of photons, and \( V = m_2 - m_1^2 \) is the variance of the DTOF. The quantities with index \( h \) refer to a homogeneous medium.

Furthermore, the obtained changes in the moments can be used in the calculation of sensitivity factors describing changes of the moments (attenuation, the mean time of flight, and the variance of the DTOF) caused by changes of absorption in defined small volume \( dV_i \) indexed by \( i \). These sensitivity factors, called the mean partial pathlength (MPP), the mean time sensitivity factor (MTSF), and the variance sensitivity factor (VSF)\(^{19} \)

\[
\text{MPP}_i = \frac{\Delta A}{\Delta \mu_{a,i}}, 
\]
(9)

\[
\text{MTSF}_j = \frac{\Delta \langle t \rangle}{\Delta \mu_{a,j}}, 
\]
(10)

\[
\text{VSF}_j = \frac{\Delta V}{\Delta \mu_{a,j}}. 
\]
(11)

For the analysis of the \textit{in vivo} data, it has been assumed that the medium can be simulated by a layered structure with the sensitivity factors for the layers obtained by summing up MPP\(_i\), MTSF\(_j\), and VSF\(_j\) in volumes corresponding to extracerebral and brain tissues.

Particularly, the medium was divided into voxels of size 0.2 \( \times \) 0.2 \( \times \) 0.2 cm. The sensitivity factor of the extracerebral tissue was obtained by the integration of five layers of voxels forming the tissue compartment of thickness 1 cm. The next 15 layers were assumed to correspond to intracerebral tissue, and the sensitivity factors for this layer were calculated by integration of these layers.

The proposed method of calculation of the sensitivity factors by using the diffusion approximation is much faster than the Monte Carlo algorithm described in Ref. 18. Unfortunately, this method can only be used to calculate sensitivity factors with the assumption that initially the medium is homogeneous, and it gives significant errors in the vicinity of the source and the detector positions.

To provide the background optical properties for calculations of the sensitivity factors \( a \) used for the analysis of the \textit{in vivo} data, \( \mu_a \) and \( \mu'_s \) were evaluated from the measurement of the DTOFs on the head of the subject. The method of moments\(^{42} \) was used to obtain the optical properties of the medium with the assumption of homogeneity of the tissue under investigation.

In Ref. 18 the authors used changes in the moments measured at four interoptode separations for estimation of changes of the absorption coefficient in ten layers of the tissue. Unfortunately, our imaging system does not allow for a multidistance time-resolved measurement. Thus, we used the calculated sensitivity factors together with changes in three moments measured at a single interoptode distance to estimate changes of the absorption coefficient in two layers of the medium (corresponding to extra- and intracranial tissue) according to the algorithm proposed in Ref. 18.

A system of equations with two unknowns: \( \Delta \mu_{a,1} \) in the superficial layer and \( \Delta \mu_{a,2} \) in the deeper layer, and three knowns represented by changes in three moments of DTOFs, together with their sensitivity factors for each layer, was solved using the least squares method.

Changes of absorption coefficient \( \Delta \mu_{a,j} \) in the two layers (indexed by \( j \)) were estimated at two wavelengths. These changes are related to the changes in concentrations of oxy- and deoxyhemoglobin (\( \Delta c_{HbO_2} \) and \( \Delta c_{Hb} \), respectively):

\[
\Delta \mu_{a,j}(\lambda) = e_{HbO_2}(\lambda) \cdot \Delta c_{HbO_2,j} + e_{Hb}(\lambda) \cdot \Delta c_{Hb,j}, 
\]
(12)

where \( e_{HbO_2}(\lambda) \) and \( e_{Hb}(\lambda) \) are the molar extinction coefficients of oxy- and deoxyhemoglobin, respectively.

For every measured DTOF, background subtraction and correction for differential nonlinearity of the TCSPC electronics were performed. In calculation of the moments, integration was carried out for that part of the DTOF in which the number of counts dropped below 1% of its maximum value.

### 3 Results of the Phantom Experiments

#### 3.1 Liquid Phantom

Stability, accuracy, and linearity of the imager were tested. The protocols used were similar to those proposed by the European Network MEDPHOT (optical methods for medical diagnosis and monitoring of diseases).\(^{44} \) For this purpose, the physical, liquid phantom was constructed allowing for simultaneous data acquisition from 32 source-detector pairs. A fish tank was filled with a solution of milk (3.5% fat) and water (3:1), with black ink (Rotring, Germany) added to obtain optical properties in the range of the optical properties typical of living tissue. All the optodes at the two source-detector arrays were placed on the same surface of the phantom to provide measurements in semi-infinite geometry and to obtain the same homogeneous conditions for all the measured source-detector pairs. The optodes were located not closer than 10 cm apart from the walls of the fish tank, which minimized influence of ambient light or interactions of photons with the walls.

#### 3.2 Stability of the Apparatus

32 DTOFs corresponding to all source-detector pairs were measured during 90 min just after switching on all the subunits of the system. This test was repeated five times on consecutive days, on the liquid phantom described before. Typical time courses of the moments of DTOFs recorded during these experiments are presented in Fig. 3.

For the total number of detected photons \( N_{tot} \) and for variance \( V \) in Fig. 3, a warm-up time necessary to avoid the low frequency drift was about 40 min. After this time, \( N_{tot} \) and \( V \) were stable, and the fluctuations of these signals were not larger than \( \pm 2\% \). The mean time of flight of photons presented in line (b) in Fig. 3 demonstrates a drift even after 1.5 h. This effect was caused by the time drift of the laser pulses,\(^{44} \) and it had a relatively low influence on the measured...
changes in the mean time of flight and the changes of the absorption coefficient in shorter time periods. Unfortunately, this drift caused errors when the optical properties in absolute values were calculated. To minimize this error, the measurements of the absolute values of the absorption and scattering coefficients should be performed after a very long warm-up time (3 h) when the drift of the mean time of flight did not exceed 10% per 1 h.

The noise level of the measurement was also evaluated by calculation of the coefficient of variation of the moments of DTOFs for all the source-detector pairs. Coefficients of variations (CV) of the moments of DTOFs (see Fig. 4) were defined as the ratios of standard deviation and the mean value:

\[
CV(x) = \frac{\sigma(x)}{\langle x \rangle} \times 100\%.
\]

The calculated coefficients of variations of the moments showed that, for a typical range of number of counted photons per second (from 30,000 to 2,000,000), the noise level was lower than 1%. For the variance, the noise level was higher, and it reached a few percent for a low number of the counted photons.

To prove the accuracy and linearity of the imager, phantom experiments were carried out with different absorption coefficients of the liquid. This experiment was carried out after a very long warm-up time period (3.5 h) to ensure small errors resulting from the drift of the laser pulses. Statistics of the results of absorption and the reduced scattering coefficients (\(\mu_a\) and \(\mu'_s\)) measured in 32 source-detector pairs are presented in Fig. 5. Both coefficients were calculated from the moments of DTOFs with the method described by Liebert et al.\(^{42}\) The inaccuracy of the measurements of \(\mu_a\) and \(\mu'_s\) between all 32 measured spots did not exceed 18% for both coefficients in the worst cases. The error of the measurement increased with the ink concentration in the phantom because of the lower number of detected photons during the 60-s-long data acquisition period.

3.3 Depth Sensitivity

To validate depth sensitivity of the constructed imager, the local change of the optical properties in the phantom was introduced using a 2-mm-diam plastic black ball immersed into the liquid and fixed using a transparent, thin fishing line (diameter of 0.2 mm). It was assumed that the absorption coefficient of this small inclusion was very high and that the ball did not scatter the light. The surface of the black ball was rough, but it was possible that some light could be reflected from that surface. The depth of the ball immersion in the liquid was changed from 0.5 up to 3.5 cm with a step of 0.5 cm. The reduced scattering coefficient (\(\mu'_s=7.0\ \text{cm}^{-1}\)) and the absorption coefficient (\(\mu_a=0.12\ \text{cm}^{-1}\)) of the medium were estimated by calculating the moments of the measured DTOFs as it was proposed in Ref. 42. The source-detector distance was 3 cm, thus the whole measurement area was approximately \(8.5 \times 8.5\ \text{cm}^2\) with a \(4 \times 4\) pixel resolution. Changes in the moments of DTOFs obtained after immersion of the black ball, at various depths, were related to...
the measurement in homogenous medium without this inclusion. The results of $N_{tot}/N_{tot0}$, $\Delta(t)$, $\Delta V$ for $4 \times 4$ pixels for six depths are presented in Fig. 6.

Changes in the moments of DTOFs were visualized for various locations of the ball in respect to the array of the optodes. In a series of experiments, the ball was located directly below the detecting bundle, directly below the source fiber, in a middle position between the two sources and two detectors, and in the middle between the source fiber and the detecting bundle.

It should be noted that the contrast in the $N_{tot}/N_{tot0}$ images rapidly decreases with the depth of the location of the inclusion, whereas the contrast decrease for $\Delta(t)$ and $\Delta V$ is much slower. It can be also pointed out that an inclusion located superficially causes an increase of the mean time of flight and, when the inclusion is located deeper, the decrease of the mean time of flight occurs. A similar effect can be noted on the maps obtained by the analysis of variance of the DTOFs. Additionally, a difference in the size of the inclusion positioned at locations different in respect to the array optodes can be noted.

The noted differences in the size and contrast of the inclusion, as provided by the images of the different moments, are related to a higher depth and longitudinal sensitivity of the mean time of flight and variance of DTOF to the changes of the absorption coefficient, as compared to the sensitivity of $N_{tot}$. The changes in the moments as a function of the depth, obtained for inclusions located between the source and the detector case presented in Fig. 6(a), are presented in Fig. 7(a). The total number of photons showed the highest sensitivity to the absorbing inclusion located superficially, but it decreased rapidly with the depth. On the contrary, the mean time of flight and, to an even higher extent, the variance, shows a high sensitivity to absorption of an inclusion positioned deeply. Moreover, changes in variance are positive when a change of absorption appears in superficial layers, and
Fig. 6 Maps of the moments of recorded DTOFs from the liquid phantom with the absorbing inclusion located at various positions in respect to the array of optodes.
they become negative when the absorbing inclusion is located deeper. A similar phenomenon can be observed for the sensitivity factor curves as a function of depth [see Fig. 7(b)].

4 Results of the In Vivo Experiment

To check the features of the system in the in vivo measurements on the human head, several tests with volunteers during functional motor stimulation were performed. 15 healthy subjects, the mean age of 30 years (nine females and six males) were examined. Before the experiment, all the volunteers gave informed consents.

The motor cortex stimulation was carried out by a voluntary middle and index finger tapping with the thumb in a self-paced mode at a frequency of about 3 to 4 Hz. The volunteer was examined in a comfortable sitting position, and the optode arrays were fixed on the head using flexible Velcro strips. For every source fiber and detecting bundle, the hair was removed carefully from the space between the skin and fiber/fiber bundle tip. The source fibers and fiber bundles were fixed in a stand that was positioned over the subject’s head to avoid bending in the holder and movement of the fibers tips. The optode arrays were positioned on the head in such a way that the source detector, in pair 10, on the left hemisphere and respectively in pair 27 on the right hemisphere (see Fig. 8) matched the C3 and C4 locations (according to the 10-20 EEG system) corresponding to the position of the motor area.

The finger tapping experiment lasted up to ten minutes and was performed in a dark room. For all 15 subjects, two experiments were carried out. 20 cycles of 15-s long left-hand finger tapping was followed by 15 s of rest, and then the same experiment was repeated after 5 min of rest with the right-hand finger tapping.
hand finger tapping. The subjects were instructed visually and by sound commands to start and finish the task.

The recorded DTOFs were analyzed by calculation of their moments for both wavelengths. For every subject, the signals of the moments were averaged synchronously with the cycles of the finger tapping.

As an example, results of the motor stimulation experiment for one selected subject. Time courses of changes of moments of the DTOFs averaged over all cycles of finger tapping: (a) $N_{tot}/N_{tot0}$ are the changes in total number of photons, (b) $Δτ$ is the changes in mean time of flight of photons, and (c) $ΔV$ is the changes in variance of the DTOF. Changes in moments are monitored at two wavelengths 687 nm (gray line) and 832 nm (black line). Vertical lines mark the beginning and end of the finger tapping task.

Fig. 9 Results of the motor stimulation experiment for one selected subject. Time courses of changes of moments of the DTOFs averaged over all cycles of finger tapping: (a) $N_{tot}/N_{tot0}$ are the changes in total number of photons, (b) $Δτ$ is the changes in mean time of flight of photons, and (c) $ΔV$ is the changes in variance of the DTOF. Changes in moments are monitored at two wavelengths 687 nm (gray line) and 832 nm (black line). Vertical lines mark the beginning and end of the finger tapping task.

The concentration of deoxyhemoglobin decreases. The total absorption coefficient at 832-nm wavelength increases and the mean time of flight and variance decreases, respectively. On the other side of the isobestic point at 687 nm, the opposite phenomenon was observed. The trend in the $N_{tot}$ courses was probably caused by the systemic changes in the blood flow during the finger tapping or by movement artifacts. Similar trends of changes of moments were observed in most of the investigated subjects. The optical studies of motor-evoked response are often characterized by a change in blood volume rather than a significant change in oxygenation. If the blood volume increases due to a sudden inflow of oxygenated blood to the stimulated cortex area, then the absorption will increase at wavelengths located on both sides of the isobestic point. This effect may cause the observed drop of total number of photons at both wavelengths used in our investigations [Fig. 9(a)].

Using the data analysis algorithms and theory described, a separation of information originating from the intra- and extracerebral layers was performed. Reconstructed signals of the changes in concentration of oxy- ($Δc_{HBO2}$) and deoxyhemoglobin ($Δc_{Hb}$) in the intracerebral layer during the right-hand finger tapping task are presented in Fig. 10. Positioning of subplots presented in Fig. 10 correspond to the positioning of source-detector pairs according to the array shown in Fig. 8(a). In the left hemisphere, a distinct increase of $Δc_{HBO2}$, and a decrease of $Δc_{Hb}$ can be observed. A minor contralateral response of the motor cortex can also be observed.

Minor responses to the motor stimulation can be also noted in the whole image. These responses can be explained by the systemic responses or by cross talk effects, which cannot be avoided in the data analysis process. Inaccuracy of the results can appear because of the assumption on the thickness of the extracerebral tissue layer. Another source of errors is the assumption on homogeneity of the tissue used in calculation of the sensitivity factors.

Time courses of $Δc_{HBO2}$ and $Δc_{Hb}$ presented in Fig. 10 were obtained in one selected subject. In most of the investigated volunteers, much of the same trends were observed. A sequence of images of $Δc_{HBO2}$ and $Δc_{Hb}$ during the motor cortex activation for both hemispheres is presented in Fig. 11.

The response of the motor cortex for the finger tapping task is clearly visible on the contralateral side. An increase of oxyhemoglobin is combined with a decrease of deoxyhemoglobin. A minor response to the motor stimulation can be observed in the whole monitored region of the tissue and also in the ipsilateral site of the brain.

5 Discussion and Conclusions

Near-infrared spectroscopy enables us to assess the brain oxygenation level in clinical conditions at bedside. In recent years, this method has been intensively explored by many research groups. Continuous wave, frequency domain, and most advanced time domain systems were used with various values of wavelengths in the near-infrared region. For evaluation of hemodynamics of the whole cortex regions (i.e., motor and visual cortex), multidetector systems were developed. A review of the NIRS imaging systems used in the experiments on adult heads is presented in Table 1.
Optical imaging systems developed in recent years have been used in many phantom studies and in vivo applications. In several reports, the authors used optical imaging systems for tissue phantom studies, measurements on neonates, and on animals. Comparison and review of these instruments can be found in Refs. 61 and 62. The systems presented in Table 1 were successfully applied to evaluate human brain hemodynamics in adults, particularly during the functional activation.

In the last few years, most up-to-date systems have been equipped with time-resolved data acquisition. Most of the advanced instruments based on time-resolved spectroscopy are equipped with a large number of source-detector pairs, and 1 to 3 wavelengths are used.

The original construction based on the intensified charge-coupled device (CCD) is limited only to one wavelength and analysis of the time windows. Most of the time-domain systems that have recently been constructed are based on the TCSPC technique for recording distributions of time of flight of photons. An interesting solution to increase the number of detecting channels by providing four-anode photomultipliers and routing signals from them was proposed by the group from Politecnico di Milano, which applied only four TCSPC boards. However, such a procedure leads to a decrease of the number of photons collected in individual channels in proportion to their number, which decreases the signal-to-noise ratio.

The design of the instrument reported here is based on the idea realized in Physikalisch-Technische Bundesanstalt in Berlin, in which four independent TCSPC channels were used with four photomultipliers and with a $1 \times 9$ fiber switch to deliver the light pulses into nine points on the tissue under investigation. In such a solution, one map consists of 16 source-detector pairs for each of three wavelengths, and it can be recorded in less than 1 s. The time-domain imager described in the presented study is an extension of this design. We construct a system with eight independent TCSPC channels, which enable us to measure optical signals with very high photon count rates compared to the solutions where signal routing is applied. The time-resolved multichannel system

![Fig. 10 Time courses of reconstructed changes in oxy- (black line) and deoxy- (gray line) hemoglobin in the (a) left and (b) right hemisphere. A vertical line marks the beginning and end of the right-hand finger tapping task.](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics)

![Fig. 11 Maps of oxy- and deoxyhemoglobin during the motor cortex stimulation experiment (LH being the left hemisphere and RH the right hemisphere). During the right-hand finger tapping task, the changes in chromophore concentration appear on the opposite hemisphere.](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics)
Table 1 Basic technical data of the optical systems used for brain imaging in adults.

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Group</th>
<th>Sources/detectors</th>
<th>Continuous wave/frequency-domain</th>
<th>Wavelengths</th>
</tr>
</thead>
<tbody>
<tr>
<td>46 to 49</td>
<td>Hitachi, Japan</td>
<td>8/8</td>
<td>CW</td>
<td>780 and 830 nm</td>
</tr>
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<td>25 and 26</td>
<td>Martinos Center for Biomedical Imaging, Boston</td>
<td>8/16</td>
<td>CW</td>
<td>690, 830, and 780 nm</td>
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<td>50 and 51</td>
<td>Charite, Berlin, Germany</td>
<td>8/7</td>
<td>CW</td>
<td>760 and 850 nm</td>
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<td>28</td>
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<td>9/4</td>
<td>FD</td>
<td>780 nm</td>
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<td>52</td>
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<td>8/4</td>
<td>CW</td>
<td>785 and 850 nm</td>
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<tr>
<td>53</td>
<td>Physikalisch-Technische Bundesanstalt, Berlin, Germany</td>
<td>9/4</td>
<td>TD</td>
<td>687, 803, 826 nm</td>
</tr>
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<td>34 and 54</td>
<td>Politecnico di Milano, Milan, Italy</td>
<td>18/16</td>
<td>TD</td>
<td>690 and 820 nm</td>
</tr>
<tr>
<td>33 and 55</td>
<td>Martinos Center for Biomedical Imaging, Boston</td>
<td>32/18</td>
<td>TD-gating</td>
<td>One wavelength tuned in range 750 to 850 nm</td>
</tr>
<tr>
<td>Present study</td>
<td>Institute of Biocybernetics and Biomedical Engineering, Warsaw, Poland</td>
<td>18/8</td>
<td>TD</td>
<td>687 and 826 nm</td>
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is developed for diffuse optical topography of the adult human brain cortex oxygenation. The instrument allows for time-resolved measurements in a set of 32 source-detector pairs on the human head at two wavelengths. The instrument is mounted in a trolley (h=140 cm, w=120 cm, d=60 cm), and it can be easily transported within the hospital and used in clinical settings for monitoring of the brain function at bedside.

In the choice of wavelengths, the spectral properties of oxy- and deoxyhemoglobin are considered. In recent studies it has been shown that two wavelengths are sufficient for an accurate estimation of the changes in hemoglobin concentrations. We use wavelengths located on both sides of the isosbestic point, at which the extinction coefficients of both hemoglobins are equal. A compromise is needed to choose proper wavelengths, considering the properties of photodetectors available for detection of diffusely reflected light with/over large interoptode distances. Above 850 nm, the sensitivity of the photomultiplier tubes becomes a problem, and for short wavelengths (below 650 nm) absorption of light in tissue caused by hemoglobin is very high. These technical drawbacks and limitations, together with the analysis of recent studies on the cross talk of hemoglobin concentration calculation using attenuation of the optical signals and availability of wavelengths of picosecond laser diodes, prompted us to choose these wavelengths. The wavelengths that we adapt to our system—687 and 832 nm—fit with the sensitivity characteristics of the photomultipliers tubes that we used, and are optimal to assess the changes in concentrations of oxy- and deoxyhemoglobin in the tissue. The measurements at two widely spaced wavelengths (687 and 832 nm) cause the volume of sampled tissues to be different due to differences in optical properties at these wavelengths. This difference in sampling volume may lead to uncertainty of calculated changes of oxy- and deoxyhemoglobin concentration for inhomogeneous tissue structures.

The analysis of the recorded curves, using the calculation of moments of DTOFs, allows us to evaluate the changes in oxy- and deoxyhemoglobin with depth discrimination, and it can be used for monitoring of the patient’s cerebral hemodynamics in clinical settings.

The theoretical analysis of light propagation in tissue is carried out with the use of diffusion approximation, and the moments of distributions of times of flight of photons. This analysis allows for a depth-resolved estimation of changes of the absorption coefficient, and finally, estimation of changes in oxygenation in intracerebral tissues.

The accuracy, stability, and linearity of the developed system are verified using the technical test based on the protocols provided by MEDPHOT. Standard deviation of the absorption coefficient measured at 32 source-detector pairs is about 5%. The signal-to-noise ratio analysis shows that for count rates used typically in in vivo experiments, the coefficient of variation is below 1% in all the calculated moments of DTOFs.

The signal-to-noise ratio of the calculated moments decreases with the order of the moment and is the poorest in...
signals representing the changes in variance. This phenomenon is related to the small number of late photons detected. The signal-to-noise ratio can be improved by the increase of the number of photons that arrive to the photodetector, which can be potentially increased by increase of the power of the laser source, but such a solution needs an application of high power lasers and is restricted by ethical issues. Unfortunately, the number of photons escaping from the tissue at large source-detector separations is very limited. Additionally, the high frequency pulses from the photomultiplier cannot be processed by time-correlated single photon counting electronics because of the appearance of the pile-up effect. Considering these limitations, in the depth-resolved reconstruction of oxy- and deoxyhemoglobin, a low noise level of moments (in particular $\langle t \rangle$ and $V$) is needed, and the count rates should be as high as possible.

For validation of the depth sensitivity of the instrument and the data analysis procedures, a liquid physical phantom is constructed in which the local absorption inclusion is located at various depths. This local change of absorption is simulated by the black plastic ball. Analysis of the recorded data confirms that the time-resolved measurement allows us to differentiate information from intra- and extracerebral tissues. The measured distributions of times of flight of photons are analyzed using their moments (attenuation, the mean time of flight, and variance). It has experimentally been confirmed that these moments of the DTOFs are differentially sensitive to the changes appearing in the superficial and deeper tissue layers. In particular, it is noted that an increase in absorption of the superficial layer leads to an increase of $\langle t \rangle$ and $V$, whereas an increase of absorption in the deeper layer causes a decrease of both these moments. These phenomena may be very useful in analysis of the responses caused by the cerebral functional stimulation. During such stimulations, a certain systemic increase of oxygenation can be expected in both intra- and extracerebral tissues. This increase should be discriminated from the increase of oxygenation caused by the neuronal activation originating from the stimulated cortical area. The systemic increase of absorption may potentially cause an increase of $\langle t \rangle$ and $V$ originating from superficial tissues, which is superimposed by the corresponding decrease of the moments originating from the brain cortex. This superposition could lead to a suppression of the influence of the systemic stimulation of signals of the moments ($\langle t \rangle$ and $V$) measured, and thus to a more adequate contrast of changes of absorption caused by the neuronal stimulation reflected in these moments. The results of the stimulation of the motor cortex by the finger tapping task shows that the higher moments ($\langle t \rangle$ and $V$) allow us to image changes in the motor cortex area caused by the functional stimulation with better contrast than the attenuation data. In attenuation signals, nonphysiological responses are present, which are manifested as synchronous drops of the number of photons at both wavelengths used. These responses are present in all the optode positions on the head and can be connected with the movement artifacts, which are difficult to avoid during finger tapping. Such problems tend to disappear when the mean time of flight or variance of DTOF is considered.

The constructed instrument demonstrates relatively good properties when compared with other time-resolved spectroscopy systems used in experiments on adults. The time-correlated single photon counting method with the independent acquisition channels allows us to measure DTOFs with high count rates of the detected photons (more than 1.5 million per second at one wavelength). This characteristic of the system allows us to switch the source position without a significant decrease in the signal-to-noise ratio.

Our future work will focus on further testing of the system on more realistic tissue phantoms as well as on comparison of the time-resolved optical imager with other imaging modalities, i.e., single photon emission computed tomography (SPECT) and functional magnetic resonance imaging (fMRI) in clinical settings. We also will consider an application of the presented imager in measurements with exogenous dye.

Acknowledgments

This study has been supported by research project number 3T11E006227 financed by the State Committee for Scientific Research of the Republic of Poland.

References


