Dual-wavelength laser speckle imaging for monitoring brain metabolic and hemodynamic response to closed head traumatic brain injury in mice

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Abstract. The measurement of dynamic changes in brain hemodynamic and metabolism events following head trauma could be valuable for injury prognosis and for planning of optimal medical treatment. Specifically, variations in blood flow and oxygenation levels serve as important biomarkers of numerous pathophysiological processes. We employed the dual-wavelength laser speckle imaging (DW-LSI) technique for simultaneous monitoring of changes in brain hemodynamics and cerebral blood flow (CBF) at early stages of head trauma in a mouse model of intact head injury \( n = 10 \). For induction of head injury, we used a weight-drop device involving a metal mass \( \sim 50 \) g striking the mouse’s head in a regulated manner from a height of \( \sim 90 \) cm. In comparison to baseline measurements, noticeable dynamic variations were revealed immediately and up to 1 h postinjury, which indicate the severity of brain damage and highlight the ability of the DW-LSI arrangement to track brain pathophysiology induced by injury. To validate the monitoring of CBF by DW-LSI, measurements with laser Doppler flowmetry (LDF) were also performed \( n = 5 \), which confirmed reduction in CBF following injury. A secondary focus of the study was to investigate the effectiveness of hypertonic saline as a neuroprotective agent, inhibiting the development of complications after brain injury in a subgroup of injured mice \( n = 5 \), further demonstrating the ability of DW-LSI to monitor the effects upon brain dynamics of drug treatment. Overall, our findings further support the use of DW-LSI as a noninvasive, cost-effective tool to assess changes in hemodynamics under a variety of pathological conditions, suggesting its potential contribution to the biomedical field. To the best of our knowledge, this work is the first to make use of the DW-LSI modality in a small animal model to (1) investigate brain function during the critical first hour of closed head injury trauma, (2) correlate between injury parameters of LDF measurements, and (3) monitor brain hemodynamic and metabolic response to neuroprotective drug treatment. © 2015 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.20.10.106009]

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1 Introduction

Traumatic brain injury (TBI), a global health problem, is a highly complex disorder involving a large number of biochemical and metabolic changes leading to a cascade of pathophysiological events such as inflammation, necrosis, and excitotoxicity, resulting in tissue damage and neuronal cell death. Over the years, substantial progress in both hardware and software has been made in conventional neuroimaging modalities, such as computed tomography (CT), magnetic resonance imaging (MRI), and positron emission tomography, to monitor and evaluate these pathological changes, each with its specific advantages and limitations. These techniques have revolutionized medical research and greatly advanced our understanding of the pathophysiological response to head injury; however, their routine use remains complex and expensive. In contrast to these modalities, optical tissue imaging techniques circumvent several limitations of conventional neuroimaging and possess specific advantages: they are relatively inexpensive, easy to operate, portable, and are free from the adverse effects of ionizing radiation. The advantages of optical imaging techniques have garnered significant attention to their suitability to biomedical applications. These optical imaging platforms work on the principle that hemodynamic parameters, such as blood volume, oxygenation, concentration of hemoglobin, lipids, and water, as well as structural changes, can be measured indirectly by tissue’s absorption and scattering coefficients. Thus, dynamic changes in brain tissue during the development and treatment of head injury can be detected and analyzed via changes in these primary coefficients. With that, many researchers were inspired to develop a variety of optical diagnostic methods for functional brain monitoring.

Prominent among these methods is dual-wavelength laser speckle imaging (DW-LSI) modality, first introduced in the pioneering study of Dunn et al., who were the first to utilize the combination of LSI and spectroscopic imaging in a single optical imaging instrumentation to characterize cerebral hemodynamic, oxygen metabolic, and blood flow changes in real time during cortical spreading depression and whisker stimulation in rats. DW-LSI is relatively inexpensive, simple, wide-field, and noncontact optical biomedical imaging modality that integrates the principles of laser flowmetry and oximetry to
obtain macroscopic information such as hemoglobin concentration and blood flow. These data are obtained by a single standard CCD camera collecting the diffused speckled light reflected from a sample illuminated by laser sources of two different light wavelengths. Thus, using the modified Beer-Lambert law, one can extract a range of tissue hemodynamic parameters from the collected light reflection data. Since the initial development of DW-LSI, several modifications and applications have been reported using this principle adopted by the biomedical optics community. As suggested recently, simultaneous changes in blood flow and metabolic oxygenation under physiological and pathological conditions (TBI, stroke, epilepsy, etc.) can be monitored optically with high spatiotemporal resolution utilizing DW-LSI.

The present work implements DW-LSI for investigation of brain pathophysiological changes during the first critical hour following closed head injury (CHI) in intact mouse head model. CHI is a type of TBI in which the brain is injured as a result of a blow to the head or a sudden, violent motion that causes the brain to move inside the skull. It is different from open head injury in that no object penetrates the intact skull. In this work, CHI was induced by the regulated impact of a 48 g cylindrical metallic rod upon the intact mouse scalp. The mouse CHI model is used to study the cascade of neurophysiological deficits caused by CHI, leading to cell death. Since the layers of the mouse skull are optically thin and highly transparent to light, we consider the diffuse reflected signals to originate from the brain itself. During experiments, the anesthetized mouse head was irradiated continuously by two laser wavelengths before, during, and after the onset of injury. Captured images were analyzed to examine the spatiotemporal characteristics of a selected, clinically relevant battery of volume-averaged hemodynamic and metabolic parameters. Results indicate dynamic changes in brain physiological properties relative to baseline measurements, observed up to 1 h post-injury. Additionally, DW-LSI was applied to test the effect of the neuroprotective drug therapy with hypertonic saline (HTS) on brain function after injury. Furthermore, the variations in cerebral blood flow detected using DW-LSI were found to be similar to measurements obtained using a commercially available laser Doppler flowmetry device. Overall, our findings indicate that DW-LSI is a suitable platform for study of both normal and pathophysiologic brain conditions, such that it holds promise for neuroscience research applications. The present work is part of a series of studies aimed to assess brain hemodynamic and metabolic markers during CHI in anesthetized rodents, using different optical platforms.

The continuation of this paper is organized as follows. Section 2 provides the experimental protocol, instrumentation setup, and data processing, including a brief review of the methods by which hemodynamic and metabolic parameters were estimated. Experimental results and discussion are presented in Sec. 3. Finally, in Sec. 4, we present a brief summary of results reported in this paper and consider the potential use of DW-LSI technique as a clinical diagnostic tool.

2 Materials and Methods

2.1 Animals and Injury Protocol

Animal procedure was performed in accordance with the regulations of the Institutional Animal Care and Use Committee of Ariel University, in accordance with the guidelines of the National Institute of Health. Twenty healthy male Sabra mice with an average body mass of 48 ± 4 g weight at 25 ± 1 weeks of age were used in the following experiments and housed in groups of four to five per cage in a 12 h:12 h light:dark cycle with a room at a controlled temperature of 22 ± 1°C. Food and water were provided ad libitum. After anesthesia by intraperitoneal injection with a cocktail of ketamine (80 mg/kg), xylazine (20 mg/kg), and saline (NaCl, 0.9%), each mouse was restrained on an in-house-made stand, the head was immobilized, scalp hair carefully removed using a commercial hair removing lotion, and a sponge was inserted underneath the chin. Core body temperature was monitored continuously during the experiment using a thermocouple rectal probe and body temperature was maintained at ~34°C throughout experimentation. Respiratory activity and blood oxygen saturation (SpO₂) were monitored at regular intervals using a commercial pulse oximeter device (Nonin, 8600) attached to the forelimb. CHI was induced under anesthesia using a weight-drop apparatus including a vertical 1-m-long metal tube (inner diameter 13 mm) with a cylindrical metallic weight (48 g, length = 9.5 cm, Φ = 10 mm) dropped through the tube from 87 cm directly above the intact mouse scalp, producing an impact of 4176 g cm between the anterior coronal suture (Bregma) and posterior coronal suture (Lambda). This model of injury closely mimics real-life focal head trauma and was shown to not produce structural brain damage. To validate that no physical damage occurred following weight impact, mouse heads were imaged using an x-ray system (Mobielt, Siemens) combined with a computerized radiography system (Kodak, CR360), before and after injury. Representative head photographs can be seen in Fig. 1. No contusion or hemorrhage is observed. Baseline reflectance measurements (15 min) were obtained prior to induction of injury, after which the mouse was placed under the weight-drop device orthogonally to the point of impact.
impact. Immediately postinjury, the mouse was returned to the optical setup and the effects of injury were studied for 100 min. Prior to data analysis, the collected diffuse images were first normalized to overcome the nonlinearity of the camera quantum efficiency at the above wavelengths, after which the images were digitally filtered (averaging filter using fspecial function in MATLAB®) to (1) remove periodic noise due to heart beat and respiration, (2) suppress speckle noise resulting from coherent illumination, and (3) to eliminate the high-frequency noise originating from the camera itself during recording. No other processing was applied to the images. It should be emphasized that the above filtering process was not applied for the flow analysis since it would reduce the information content in speckle patterns, and was used only as a preprocessing step before hemoglobin concentration calculation. A selected region of interest (ROI) with the impact site at its center was selected by the investigators for data processing.

2.3 Imaging Sampling

The intact mouse scalp was sequentially illuminated at 530 and 660 nm wavelengths and the remitted reflectance was captured by a single CCD camera (raw speckle images) and saved for offline processing and analysis with in-house scripts written in MATLAB® software. Two images at two different wavelengths were acquired in order to assess variation in tissue composition levels (diffuse reflectance spectroscopy concept). Throughout the experiments, the time interval between the images acquired from each wavelength was ∼1 s. Imaging commenced before induction of CHI to establish baseline physiological parameters and continued during and following experimental intervention. Baseline images were obtained 15 min before injury induction, such that each mouse served as its own control. Imaging commenced ∼3 min after CHI and continued for 100 min. Prior to data analysis, the collected diffuse images were first normalized to overcome the nonlinearity of the camera quantum efficiency at the above wavelengths, after which the images were digitally filtered (averaging filter using fspecial function in MATLAB®) to (1) remove periodic noise due to heart beat and respiration, (2) suppress speckle noise resulting from coherent illumination, and (3) to eliminate the high-frequency noise originating from the camera itself during recording. No other processing was applied to the images. It should be emphasized that the above filtering process was not applied for the flow analysis since it would reduce the information content in speckle patterns, and was used only as a preprocessing step before hemoglobin concentration calculation. A selected region of interest (ROI) with the impact site at its center was selected by the investigators for data processing.

2.4 Calculation of Hemodynamic and Metabolic Parameters

As noted before, DW-LSI integrates two wavelengths in an LSI setup to simultaneously image both the relative changes in blood flow (with 660 nm) and changes in hemodynamic parameters (with both 530 and 660 nm) over both space and time. After data collection and preprocessing as described in Sec. 2.2, a standard sliding-window algorithm (7 × 7 pixel
dimension) was applied to convert each 660 nm raw speckle image to a speckle contrast \((K_s)\) image calculated by the ratio of the image’s standard deviation \((\sigma_s)\) to its mean intensity \((\langle I \rangle)\):

\[
K_s = \sigma_s \langle I \rangle^{-1} \times \frac{1}{\sqrt{V}}.
\]

(1)

where \(V\) is the mean cerebral blood flow \((\text{CBF})\) speed, such that lower speed results in higher \(K_s\) and vice versa. This \(K_s\) \(V\) relationship is true when the camera exposure time is higher than speckle decorrelation time. Thus, blood flow dynamics were extracted from the reflectance images. For the reader’s knowledge, transcranial Doppler ultrasonography is the most common noninvasive tool used clinically to estimate CBF. Other clinical flow measurement modalities, such as xenon CT and arterial spin labeling MRI, are also in use.

Simultaneously to flow measurements, temporal changes in concentrations of oxyhemoglobin \((\text{HbO}_2)\) and deoxyhemoglobin \((\text{Hbr})\) were extracted from the diffuse reflectance data based on the Beer-Lambert law described below:

\[
\begin{align*}
\Delta \text{HbO}_2(t) &= \left\{ \frac{\varepsilon_{530} K_\text{ffiffiffi} \sqrt[3]{R_{530}^{ \text{HbO}_2(\text{t})} - R_{530}^{ \text{HbO}_2(\text{o})}}}{\varepsilon_{660} K_\text{ffiffiffi} \sqrt[3]{R_{660}^{ \text{HbO}_2(\text{t})} - R_{660}^{ \text{HbO}_2(\text{o})}}} \right\}^{-1} \left\{ \frac{\ln[R_{660}^{ \text{HbO}_2(\text{t})}/R_{660}^{ \text{HbO}_2(\text{t})}]}{D_{530}} - \frac{\ln[R_{530}^{ \text{HbO}_2(\text{t})}/R_{530}^{ \text{HbO}_2(\text{t})}]}{D_{660}} \right\},
\end{align*}
\]

(2)

where \(\varepsilon\) is the molar spectral absorbance coefficient of an absorbing chromophore, \(R_{530}(t)\) and \(R_{660}(t)\) are the measured diffuse reflectance values at time \(t\) for the two wavelengths after injury, and \(R_{530}(0)\) and \(R_{660}(0)\) are the baseline values prior to injury. \(D_{530}\) and \(D_{660}\) are the differential path length factors of the light inside the tissue. We assume that at the two wavelengths used in the present setup, (1) scattering is nearly constant and (2) other chromophores such as lipids, cytochrome-\(c\) oxidase, collagen, etc. contribute insignificantly to the absorption spectrum, relative to hemoglobin. The total hemoglobin concentration change can also be quantified by \(\Delta \text{THC} = \Delta [\text{HbO}_2] + \Delta [\text{Hbr}]\). For the reader’s knowledge, \(\Delta \text{THC}\) can give information about changes in cerebral blood volume \((\Delta \text{CBV})\) and it is assumed to be linearly proportional to CBV, thus \(\Delta \text{THC} \sim \Delta \text{CBV}\). We chose to record at 530 and 660 nm based on their sensitivity to THC and Hbr, respectively. At 530 nm (isosbestic point), \(\text{HbO}_2\) and \(\text{Hbr}\) absorb light equally and the change in reflection is proportional to THC; THC can be readily obtained at this wavelength. At 660 nm, the absorption of \(\text{HbO}_2\) is threefold less compared with the absorption of \(\text{Hbr}\) such that both can be easily calculated.

In addition to hemoglobin concentration changes and based on Eqs. (1) and (2), the relative changes in oxygen metabolism, known as cerebral metabolic rate of oxygen \((\text{CMRO}_2)\), can be calculated:

\[
\text{rCMRO}_2 = \frac{\Delta \text{CMRO}_2(t) - \Delta \text{CMRO}_2(o)}{\text{CMRO}_2(o)} \propto 1 + \gamma_R \frac{\Delta \text{Hbr}(t)}{\text{Hbr}(o)} - 1.
\]

(3)

\(\gamma_R\) and \(\gamma_T\) are vascular weighting constants that are assumed to be 1 for simplification, which is within a physiologically plausible range of 0.75 to 1.25. \(\text{rCBF}\) is the fractional change of CBF compared with baseline and is approximated by \[\Delta V(t)/V(o)\] calculated with Eq. (1). It has been shown that \(\gamma_R\) and \(\gamma_T\) do not influence the calculated \(\text{rCMRO}_2\). \(\text{CMRO}_2\) (i.e., oxygen consumption) measures the difference between the rate of oxygen flowing into and out of a region of investigation. \(\text{CMRO}_2\) is a valuable tool in the clinical setting, as it helps to study brain function. It is considered a function of mitochondrial activity and can be calculated from CBF and the arteriovenous oxygen content difference. To highlight relative changes over time, each individual image was averaged using mean2 function in MATLAB® providing a single mean value of individual pixel intensities within the entire ROI. Once the CBF and \(\text{CMRO}_2\) values are available, cerebral

![Fig. 4](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics attachment)

**Fig. 3** Flow chart describing the estimation of different brain tissue properties. First, raw consecutive images at two wavelengths of illumination are acquired. Next, the flow image is obtained with the 660 nm image via Eq. (1). Concurrently, hemoglobin features are calculated through Eq. (2) using both images of 530 and 660 nm. Next, the combination of measured hemoglobin and blood flow was utilized to derive relative cerebral metabolic rate of oxygen \((\text{CMRO}_2)\) by Eq. (3). Finally, based upon the cerebral blood flow \((\text{CBF})\) and \(\text{CMRO}_2\) values, cerebral arteriovenous difference of oxygen \((\text{AVDO}_2)\) is obtained by Eq. (4).

**Fig. 4** Representative experimental raw speckle images at 530 and 660 nm (left) recorded from an intact mouse scalp, respectively, and the corresponding speckle contrast \((K_s)\) (middle) and velocity (right) maps at representative time points. These maps were computed directly from raw speckle image (660 nm) using Eq. 4 with 7 × 7 areas of pixels.
arteriovenous difference of oxygen (AVDO₂) can be obtained. AVDO₂ represents the amount of oxygen extracted by the brain, and reflects the balance between CMRO₂ and CBF calculated by:

\[ \Delta AVDO₂ = \frac{\Delta CMRO₂}{\Delta CBF} \]  

Overall, in the present study, a total of six hemodynamic and metabolic parameters can be derived simultaneously within specific areas of interest by the DW-LSI system: V (or \( K^{-2} \)), \( \Delta HbO₂ \), \( \Delta Hbr \), \( \Delta THC \) (or \( \Delta CBV \)), \( rCMRO₂ \), and \( \Delta AVDO₂ \). It should be pointed out that because of the partial volume effect, we were able to obtain only relative changes in the above parameters over time, relative to baseline values, rather than their absolute values. The above steps are summarized in the block diagram illustrated in Fig. 3. In Fig. 4, representative raw images at two wavelengths used and speckle contrast and velocity maps at specific time point of baseline, postinjury, and euthanasia are presented, highlighting flow changes occur over time.

### 2.5 Statistical Analysis

All the data are expressed as mean ± standard error, and two-way analysis of variation was performed to determine...
significant differences in means. Results were judged to be statistically significant if the \(p\) value was < 0.05. Calculations were executed using MATLAB®’S Statistic Toolbox (Version 2010a, The MathWorks, Inc.).

3 Results and Discussion

Figures 5(a)–5(d) summarize the statistical analysis of changes in chromophore concentrations and metabolism properties over time in 10 animals at normal state (baseline), following brain injury challenge, and after overdose administration. Please note that the error bars (standard error) in all panels may represent the variation in the calculated mean reflectance for each ROI. These variations expressed as error bars can be explained by variability among individual tissue sample properties, as well as unavoidable variances due to breathing, involuntary muscle contraction, etc. Relative to baseline, a salient difference in a range of brain physiological properties was demonstrated. Viewing the outcomes revealed that progress from baseline to brain injury induces an abrupt increase in [Hbr] and decrease in [HbO2] [Fig. 5(a)], concurrently with slightly increased \(\Delta THC\) (consequently, \(\Delta CBV\)). The increase in \(\Delta THC\) following injury appeared to be primarily due to the relative rise in deoxyhemoglobin concentration. It can be seen that the chromophore concentrations before and after injury are distinct across the experiment duration. These results appear to reflect the pathophysiology of the brain following trauma, which can lead to permanent brain damage, neurological disorders, or even death. The higher level of [Hbr] due to ongoing metabolic activity promotes the development of secondary brain damage stemming from ischemia, a common occurrence postinjury. The difference between \(\Delta[HbO2]\) and \(\Delta[HbO2]\), expressed as \(\Delta[HbD]\) representing the mismatch between the two parameters, can serve as a marker of oxygen delivery. Indeed, \(\Delta[HbD]\) was observed to decrease following injury. The slower rate of change in THC may have corresponded to decreased blood flow resistance in the injured area, as confirmed by Figs. 5(b) and 5(c).

Figure 5(b) represents the variation in CBF, derived as in Eq. (1) above. Three phases of spatial changes in CBF, which highlight the cerebral autoregulation (CAR) function, were observed: following the first few minutes after injury, blood flow increased and subsequently decreased toward its baseline level. In the homeostatic CAR process, arterioles undergo ongoing dilation and constriction, yielding cyclical changes in cerebral vascular resistance, in turn changing arterial blood pressure and cerebral perfusion to maintain a constant blood flow. Disturbed CAR can reduce the ability of the injured brain to preserve adequate blood flow, as observed in patients with mild to severe brain injuries. Generally speaking, impaired CAR adversely impacts clinical outcome and is associated with cerebrovascular abnormalities and increased mortality. The increases in blood flow we observed may be compensatory mechanisms, which serve to overcome the effect of the injury or as a result of the euthanasia injection.

Under decreased CBF, brain cells undergo a series of pathophysiologic changes further complicated by disrupted oxygen balance, leading to disrupted extracellular ion concentrations, increased lactate levels, decreased extracellular pH, abnormal accumulation of glutamate, etc. Studies of CBF in severe head-injured patients show that, among other factors, altered CBF levels are related to poor clinical outcome. The variations in CBF currently observed with the DW-LSI system were

![Fig. 6 Scatter plot showing correlation between \(rCMR_{O2}\) and \(\Delta CBF\) for entire experiments. To facilitate comparison, both \(\Delta CBF\) and \(rCMR_{O2}\) were normalized between 0 and 1 \((n = 10)\).](https://example.com/fig6)

![Fig. 7 Change in AVDO\(_2\) during postinjury period: (a) dynamics of AVDO\(_2\) postinjury and (b) \(\Delta AVDO_2\) versus \(\Delta CBF\) following injury. The line is the best-fit linear regression (circles), with correlation \(R^2 = 0.8\) \((n = 10)\). The slope of the linear fit was \(-1.14\) and the overall mean difference between the two was \(\sim8\%\).](https://example.com/fig7)
confirmed using a laser Doppler flowmetry (LDF) probe (Periflux, PF 2B), as presented by Fig. 5d. As demonstrated by both methodologies, CBF dropped after injury, indicating decreased oxygen delivery to the brain, which may result in permanent hypoxic brain damage. Similar changes in CBF following CHI in piglet brains were observed by Yodh’s group during axonal injury [Ref. 47, Fig. 4c] and others reporting alterations in CBF following traumatic brain injury in both animal models and clinical findings.

In addition to the above panels, time-course assessment of the changes in oxygen metabolism (relative \(rCMRO_2\)) is presented in Fig. 5c. \(rCMRO_2\), combining Figs. 5a and 5b according to Eq. (3), is considered an indicator of brain tissue viability. Figure 5c demonstrates an ~35% reduction in \(rCMRO_2\) relative to baseline, suggesting metabolic suppression, mitochondrial dysfunction, and reduced oxygen consumption following injury. The average normal \(CMRO_2\) is 1.5 \(\mu\text{mol/g/min}\), and after head injury, it often falls to an average of 0.9 \(\mu\text{mol/g/min}\), a reduction of 40%, close to that observed presently. Several groups have reported a similar decrease in \(CMRO_2\) in brain injury models during cerebral ischemia. Decreased \(CMRO_2\) postinjury also indicates a reduction in electrocortical activity. Changes in metabolism and CBF tend to correlate tightly, such that increased metabolic demand is met rapidly by increased CBF and substrate delivery. These changes are thought to be controlled by several vasoactive metabolic mediators including hydrogen ions, potassium, CO\(_2\), adenosine, glycolytic intermediates, phospholipid metabolites, nitric oxide, etc. However, as observed here, the timecourse of \(CMRO_2\) changes in head injury differs substantially from those observed in \(\Delta\text{CBF}\), such that \(\Delta\text{CBF} > rCMRO_2\). This condition, known as cerebral hyperemia, is frequently seen in severe head injury patients and contributes to brain swelling and high intracranial pressure (ICP). Although CBF and \(CMRO_2\) share a common variable, they have been noted often not to be correlated in patients with head injuries. The uncoupling between these two parameters induced by head injury is believed to contribute to cerebral ischemia and cerebral hyperemia. Muizelaar et al. observed this uncoupling also among children suffering from head injury, while Robertson et al. show that when coupled with elevated CBF, a higher \(CMRO_2\) is associated with a better outcome (Ref. 56, Fig. 9). To facilitate comparison, both \(\Delta\text{CBF}\) and relative \(rCMRO_2\) were first normalized between 0 and 1 and presented relative to each other in Fig. 6. Since \(CMRO_2\) possesses temporal dynamics different from those of hemodynamic response, both parameters together provide a more complete view on the changes. 

Fig. 8 Box plot graph displays the average values of (a) arterial oxygen saturation (SpO\(_2\)) and (b) heart rate (HR) at representative time points \((n = 10)\). 

Fig. 9 Example plot of a representative mouse experiment. (a) Traces of hemodynamic and metabolic response following CHI and (b) change in CBF over the entire time course as measured by the LDF probe. Note similarity to DW-LSI results (Fig. 5d).
brain function following injury. The discrepancy between the two, presented in a scatter plot, documents the uncoupling between blood flow and cellular metabolism, which stems from variations in the cerebral oxygen extraction fraction. The change in \( \Delta AVDO_2 \) during the postinjury period was analyzed and shown in Fig. 8. \( \Delta AVDO_2 \) time course and its negative correlation to \( \Delta CBF \) are presented in Figs. 7(a) and 7(b), respectively. Variations in \( \Delta AVDO_2 \) following injury are believed to result from the changes observed in CBF. The correlation between \( \Delta AVDO_2 \) and CBF has been reported elsewhere.

As expected and shown in all figures, euthanasia (HIBI model) worsens both hemodynamic and metabolic parameters over time, causing a further decrease in the already limited blood supply to the injured zone, leading to permanent neuronal damage and finally death. HIBI most often results from insults such as cardiac arrest, vascular catastrophe, poisoning (i.e., drug overdose), or head trauma. In our experiments, decreased oxygen delivery to the brain accompanied decreased cerebral perfusion, resulting in failure of autoregulation mechanisms and a severe decrease in cerebral blood flow over time.

The early changes reported during the injury and euthanasia periods delineate: (1) the pathophysiological processes of the brain in response to CHI and overdose injection and (2) changes in brain hemodynamics and metabolism, in agreement with the results available in the literature and (3) demonstrate the capability of DW-LSI to monitor dynamics of hemodynamics and metabolism during brain injury over time through the intact scalp. In addition to hemodynamics and metabolic activities, Fig. 3 presents box plots at representative time points of the heart rate (HR) and arterial oxygen saturation (SpO\(_2\)) values during the experiments. The HR was found to significantly decrease with time postinjury, which reflects the severity of the injury model used, indicating that the damage reached the brainstem. SpO\(_2\) fluctuated near its baseline level following injury and dropped following euthanasia, without any significant changes relative to baseline.

Brain parameters over time obtained from a representative mouse are illustrated in Fig. 9(a), alongside changes in CBF measured by the LDF setup [Fig. 9(b)]. The effects of injury and euthanasia are clearly visible and reflect the experiments outlined in Fig. 8. Figure 10 displays the speckled contrast map and its corresponded pixel histogram profile for baseline [Fig. 10(a)], following brain injury [Fig. 10(b)], and after overdose injection [Fig. 10(c)]. Each contrast map covers an area of \( \sim 10 \times 10 \text{ mm}^2 \), corresponding to 656 x 492 pixels, resulting in an average spatial resolution of \( \sim 0.015 \text{ mm/pixel} \). The color bar to the right of each map indicates the \( K_s \) [Eq. (1)] value. As expected, spatiotemporal modification of the speckle patterns is apparent: as time elapses from the baseline, increased map contrast indicates decreased blood flow (\( K_s \sim 1/\sqrt{V} \)). In addition, we can observe from the histogram profile that each state is characterized with a characteristic peak value and distribution width (at half of peak height). Both parameters together may serve as an indicator of the brain’s pathophysiological state.

Finally, the effects of the neuroprotective agent frequently used in intensive care and operating units for treatment of head injury, HTS, was tested on injured mice (\( n = 5 \)). HTS was arbitrarily selected among drugs in clinical use for treatment of head trauma. HTS is an osmotic agent administered intravenously to patients in the acute phase of severe TBI. It acts to reduce brain swelling and ICP by increasing the sodium levels in the bloodstream, thereby inducing a shift of fluid across the osmotic gradient it generates from the intracellular to the extracellular space. Shortly after administration, HTS reduces blood viscosity, which improves CBF and cerebral oxygenation.

**Fig. 10** Two-dimensional speckle contrast map and pixel histograms: (a) baseline, (b) injury period, and (c) following euthanasia. The scale bar on the right side of each map panel represents the speckle contrast value, \( K_s \), of each pixel in the map. Map size is \( \sim 10 \times 10 \text{ mm} \), 656 x 492 pixels. As time elapses, the map contrast increase indicates decreased blood flow. Histograms showing the distribution of the \( K_s \) parameter value for the three states indicate the degree of spatial variation in \( K_s \). The solid curve is a Gaussian fit with appropriate mean and standard deviation. In each state, peak value and distribution width of the histogram are modified, respectively.
causing autoregulatory vasoconstriction, thereby further reducing ICP. The protocol described in Sec. 2.1 was slightly modified for the HTS study: baseline measurements were obtained prior to induction of injury for 15 min, after which the mouse was removed and placed under the weight-drop device, and underwent focal TBI. Immediately post-injury, the mouse was returned to the optical setup and the effects of injury were studied 20 min post-trauma. At the end of the 20 min, a single injection of 5 μL/g of 23.4% NaCl HTS was administered intraperitoneally and measurements were acquired for another 30 min. After 30 min, the mouse was sacrificed. This modified protocol is outlined in Fig. 11. Figure 12 presents the time course of hemodynamic and metabolic parameters of mice treated with HTS; an ∼20% decrease in ΔHbr and an ∼25% increase in ΔHbO2 follow HTS injection. At the same time, both ΔCBF and rCMRO2 increased gradually (Fig. 13) in contrast to mice not treated with HTS [Figs. 5(b) and 5(c)], suggesting a possible mechanism for the clinical neuroprotective properties of HTS. Increased CBF may be attributed to the decreased ICP and increased systematic blood pressure caused by HTS. The correlation between ΔAVDO2 and ΔCBF over time is presented in Fig. 14, and the differences pre- and post-HTS injection are demonstrated. As shown in the graph, HTS increases the linearity of the slope between the two states from 0.5 to 0.97, coupled with a correlation of $R = 0.78$ ($p < 0.05$). Increase in slope may highlight decrease in cell swelling and brain edema evolution. In addition, HR and SpO2 measurements in Fig. 15 reveal that HTS treatment stabilizes both parameters in comparison to untreated mice (Fig. 8). In light of these results, we conclude that HTS successfully mitigated brain tissue damage in early stages post-injury. Nevertheless, further studies with

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**Fig. 11** Modified experimental protocol including hypertonic saline (HTS) injection.

**Fig. 12** Time series of hemodynamic and metabolic response following HTS injection. ΔHbr decreased by ∼20% while ΔHbO2 increased by ∼25% following HTS injection ($n = 5$).

**Fig. 13** Time series of the changes in (a) CBF and (b) CMRO2 following the HTS protocol. Both metrics increased following HTS administration ($n = 5$).

**Fig. 14** Changes in AVDO2 as a function of CBF post-injury and post-HTS injection. Data points represent measurements and lines represent linear fits to this data. HTS increases the linearity of the slope between the two from 0.5 to 0.97. Both parameters are in correlation: $R = 0.78$. 

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larger animal populations are needed in order to further elucidate the effects of HTS treatment, which will be elaborated in our ongoing research.

4 Conclusion

In the present study, we demonstrated the feasibility of a DW-LSI approach to simultaneously and continuously measure brain hemodynamic and metabolic parameters in a mouse model of intact TBI. In addition to application of DW-LSI to head injury treatment, the ability to continuously monitor the dynamics of hemodynamic and metabolic parameters may also aid the optimization treatment of cerebral hemorrhage or aneurism. To this end, we also demonstrated that DW-LSI can be utilized to study brain pathophysiology in a mouse model of HIBI. In addition, the effectiveness of HTS as a neuroprotective drug to reduce brain tissue damage was evaluated with DW-LSI. The ability to continuously monitor such neuro-pathophysiological parameters immediately after injury and following therapeutic strategies serves a critical clinical need, enabling the management of proper treatment. Therefore, DW-LSI may be further developed as a low-cost, noninvasive, and noncontact imaging modality for monitoring neuropathology in clinical settings. In the near future, larger-scale experiments are warranted to further elucidate our findings. Additionally, our ongoing research is aimed to test the capability of DW-LSI to monitor other pathological parameters of the brain, in concert with other optical and/or conventional approaches.

References


Biographies for the authors are not available.