

## Article

# A Non-Invasive Optical Multimodal Photoplethysmography-Near Infrared Spectroscopy Sensor for Measuring Intracranial Pressure and Cerebral Oxygenation in Traumatic Brain Injury

Maria Roldan \*  and Panicos A. Kyriacou 

Research Centre for Biomedical Engineering, City, University of London, London EC1V 0HB, UK

\* Correspondence: maria.roldan@city.ac.uk

**Abstract:** (1) Background: Traumatic brain injuries (TBI) result in high fatality and lifelong disability rates. Two of the primary biomarkers in assessing TBI are intracranial pressure (ICP) and brain oxygenation. Both are assessed using standalone techniques, out of which ICP can only be assessed utilizing invasive techniques. The motivation of this research is the development of a non-invasive optical multimodal monitoring technology for ICP and brain oxygenation which will enable the effective management of TBI patients. (2) Methods: a multiwavelength optical sensor was designed and manufactured so as to assess both parameters based on the pulsatile and non-pulsatile signals detected from cerebral backscatter light. The probe consists of four LEDs and three photodetectors that measure photoplethysmography (PPG) and near-infrared spectroscopy (NIRS) signals from cerebral tissue. (3) Results: The instrumentation system designed to acquire these optical signals is described in detail along with a rigorous technical evaluation of both the sensor and instrumentation. Bench testing demonstrated the right performance of the electronic circuits while a signal quality assessment showed good indices across all wavelengths, with the signals from the distal photodetector being of highest quality. The system performed well within specifications and recorded good-quality pulsations from a head phantom and provided non-pulsatile signals as expected. (4) Conclusions: This development paves the way for a multimodal non-invasive tool for the effective assessment of TBI patients.

**Keywords:** photoplethysmography (PPG); near infrared spectroscopy (NIRS); instrumentation system; cerebral oxygenation; intracranial pressure; non-invasive brain monitoring; optical sensors



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## 1. Introduction

Traumatic brain injury (TBI) is caused by an external force that disturbs the brain's function [1,2]. Every year 50 to 60 million new cases are reported worldwide, including mild, moderate, and severe cases of TBI with mortality rates as high as 40% for the latter [3]. Survivors also present disabling outcomes, reducing their life expectancy by six years [3]. After a traumatic brain injury, cerebral hemodynamics are disrupted, with intracranial pressure (ICP) and cerebral oxygenation as two of the most relevant impaired parameters [4]. These variables' alterations occur in the first hours after injury and can usually lead to secondary injuries such as ischemia or hyperemia [5]. Unfortunately, during this "golden period" of around two hours, doctors know almost nothing of what is happening inside the skull because current techniques to assess TBI are intermittent and invasive. In neurocritical care, TBI monitoring involves invasive measurements of ICP and cerebral oxygenation (PbtO<sub>2</sub>, StO<sub>2</sub>) by placing a transducer into the brain tissue through a small hole drilled in the skull, also known as a bolt [4]. Although guidelines recommend prompt and continuous monitoring of physiological variables soon after TBI, current intracranial monitoring methods are invasive, require specialized medical staff to place the bolt inside the skull, and can only be placed 3 to 4 h after injury [6]. Moreover, these methods are

used only in severe cases due to their association with infection risks. Evidence has shown that access to early monitoring and treatment can decrease the risk of secondary injury, resulting in better outcomes and reduced mortality. Similarly, the integration of multimodal monitoring in patient management has shown better results than treatment guided by one single parameter [4,7].

Non-invasive cerebral oxygenation and ICP monitoring would also drastically reduce financial costs and patient risks, including risks from patients being left unmonitored. Even in developed nations and among people with 50% chance of developing raised ICP (people with severe TBI and 2+ clinical risk factors), the ICP monitoring rate is only ~43% [8]. Non-invasive monitoring could not only dramatically increase monitoring rates in TBI; it could open up monitoring to new patient populations e.g., hydrocephalus (CSF shunt assessment), meningitis, brain tumor, and stroke [9].

This research proposes to develop a non-invasive, continuous multimodal monitoring system to assess ICP and cerebral oxygenation in TBI patients, with a primary aim to facilitate a timely treatment to decrease the severity of secondary injuries. This non-invasive and portable technology will comprise a multimodal probe and an embedded system to acquire, process, control, and visualize the signals and vital signs (absolute measurements of ICP and cerebral oxygenation), and it will allow a preliminary assessment of TBI-related physiology during the early stages of the injury. The technology will be a tool to be used by paramedics and physicians in and outside the hospital, and, as it is low-risk, it will allow the assessment of every type of TBI severity, hence improving the treatment of patients with mild or moderate TBI.

There is no evidence of any other multimodal, non-invasive technology that uses pulsatile and non-pulsatile optical brain signals to assess ICP and cerebral oxygenation quantitatively and simultaneously in TBI patients. Near-infrared spectroscopy (NIRS) has been broadly used to assess cerebral oxygenation non-invasively [10–15]. However, current NIRS devices only provide relative changes of oxy- and deoxyhemoglobin (Hb, HbO<sub>2</sub>, TOI), utilizing only the non-pulsatile brain signals [16]. A broad description of arterial and cerebral oxygenation techniques is presented in the background section. However, current NIRS technologies do not measure ICP; hence, they are not routinely used by neurosurgeons.

This research will enable the creation of a new multimodal optical sensor technology by utilizing both the pulsatile (Photoplethysmogram (PPG)) and non-pulsatile components of the backscattered light at visible and infrared wavelengths to assess absolute measurements of ICP and cerebral oxygenation. The system presented in this manuscript includes a probe designed specifically for monitoring the main physiological parameters of interest in TBI patients. In addition to the probe, the researchers developed an instrumentation system for the acquisition and pre-processing of proximal and distal PPG signals at multiple wavelengths. Different source–detector separations and multiple wavelengths are key for the assessment of superficial and deeper tissue layers and provide information to calculate oxygenation parameters such as SpO<sub>2</sub> and TOI. The sensor has the capacity to acquire good quality PPG signals (AC and DC) from the photodiodes and thus enable their implementation in further computational analysis. The functioning of the proposed TBI multimodal sensor is based on the novel analysis of the NIR pulsatile signals acquired from cerebral tissue. Changes in intracranial pressure will affect the morphology of the optical pulse, meaning that rigorous analysis of quantifiable features of the pulsatile signals could enable the development of a computational model to estimate absolute values of intracranial pressure [17].

## 2. Background

Pulse oximetry principles rely on light absorbance by different chromophores in blood. For instance, at high blood oxygen saturations (>80%), the red light (600 to 750 nm) is absorbed more than the infrared light (750 to 1000 nm), while at low saturations (<80%), the absorption is inverted. The pulse oximetry technique utilizes the pulsatile (AC) PPG components at both wavelengths to estimate the ratio of absorbances, limiting the use of

the DC component for normalization only. The ratio of ratios ( $R$ ) is then related to arterial oxygen saturation by empirical curves obtained from controlled hypoxia studies on healthy volunteers. A typical relation of  $R$  and  $SpO_2$  is shown in Equation (1) [18,19].

$$SpO_2 = 110 - 25 \cdot R \tag{1}$$

$$\text{where, } R = \frac{AC_R/DC_R}{AC_{IR}/DC_{IR}}$$

Equation (1): Oxygen saturation ( $SpO_2$ ), Ratio of ratios ( $R$ ), pulsatile component of the red wavelength ( $AC_R$ ), non-pulsatile component of the red wavelength ( $DC_R$ ), pulsatile component of the infrared wavelength ( $AC_{IR}$ ), and non-pulsatile component of the infrared wavelength ( $DC_{IR}$ ).

NIRS does not include the pulsatile component of the signal and focuses on the analysis of the slow-changing DC components. Following the modified Beer–Lambert law ( $\Delta A = \Delta \mu_a * l$ ), changes in chromophores absorption coefficients ( $\Delta \mu_a$ ) can be calculated from changes in light attenuation ( $\Delta A$ ) and the differential path length ( $l$ ). Due to  $\mu_a = \epsilon \cdot c$ , change in light attenuation allows the calculation of changes in chromophores concentrations once the light travelling path ( $l$ ) is known.

However, the modified Beer–Lambert law method measures only the changes in concentrations from an arbitrary point. Therefore, spatially resolved spectroscopy (SRS) is implemented to measure an absolute tissue oxygenation index. The Tissue Oxygenation Index (TOI) is expressed in percentage and represents the hemoglobin oxygenation saturation of the sampled volume of tissue. This method calculates the relative chromophores concentrations by measuring the relative change in light attenuation ( $A$ ) along the distance ( $\rho$ ). In NIRS, two distal photodetectors are employed—close together (8 mm), and far from the light source, so that the measurement is less affected by the head geometry or surface non-uniformity. Moreover, due to the small distance between the photodetectors (<10 mm), the change of  $A$  along  $\rho$  can be assumed linear, which is a prerequisite for SRS. Furthermore, the multi-distance approach makes SRS less susceptible to the superficial structures within the light path.

The value of  $\frac{\partial A}{\partial \rho}$  is measured by the probe as the change of light attenuation along the photodetectors’ distance. Light attenuation is related to light intensity (voltage measurements) by applying the Beer–Lambert Law as follows:

$$\frac{\partial A(\lambda_j)}{\partial \rho} = \frac{\log\left(\frac{I_{\text{photodiode2}}(\lambda_j)}{I_{\text{photodiode1}}(\lambda_j)}\right)}{\rho_{\text{photodiode2}} - \rho_{\text{photodiode1}}} \tag{2}$$

Equation (2): Specific wavelength ( $\lambda_j$ ), change of light attenuation along the photodetectors’ distance ( $\partial A/\partial \rho$ ), light intensity at each photodetector ( $I_{\text{photodiode}}$ ), and source-detector distance of each photodetector ( $\rho_{\text{photodiode}}$ ).

By solving the diffusion approximation for a highly scattering medium ( $\mu_a \ll \mu_s$ ), the value of  $\frac{\partial A}{\partial \rho}$  is related to the relative absorption coefficient of the specific wavelength as follows:

$$K * \mu_a(\lambda_j) = \frac{1}{3 * (1 - h * \lambda_j)} * \left[ \ln * \left( 10 * \frac{\partial A(\lambda_j)}{\partial \rho} \right) - \frac{2}{\rho} \right]^2 \tag{3}$$

Equation (3): Unknown constant for scattering in the near-infrared region ( $K$ ), absorption coefficient ( $\mu_a$ ), specific wavelength ( $\lambda_j$ ), normalized slope of  $\mu_s$  along  $\lambda$  ( $h$ ), change of light attenuation along the photodetectors’ distance ( $\partial A/\partial \rho$ ), and mean emitter-detectors distance ( $\rho$ ).

Where  $h$  is invariable among tissues and subjects,  $h = 6.3 \times 10^{-4}$  (mm<sup>-1</sup>/nm) is used [20]. Furthermore, it is possible to calculate the relative concentrations of both chromophores by solving the following equations system for at least two wavelengths

(770 nm and 880 nm). Where  $\varepsilon_{ij}$  are the extinction coefficients of  $i = [HbO_2], [HHb]$  and  $j = \lambda_1, \lambda_2$ .

$$\begin{bmatrix} K * HbO_2 \\ K * HHb \end{bmatrix} = [\varepsilon_{ij}]^{-1} * \begin{bmatrix} K * \mu_a(\lambda_1) \\ K * \mu_a(\lambda_2) \end{bmatrix} \quad (4)$$

Equation (4): Unknown constant for scattering in the near-infrared region ( $K$ ), oxy-hemoglobin concentration ( $[HbO_2]$ ), deoxyhemoglobin concentration ( $[HHb]$ ), absorption coefficient ( $\mu_a$ ), and specific wavelength ( $\lambda$ ).

These concentrations are used to find the total oxygenation index ( $TOI$ ).

$$TOI = \frac{K * [HbO_2]}{K * [HbO_2] + K * [HHb]} \quad (5)$$

Equation (5): Total oxygenation index ( $TOI$ ), unknown constant for scattering in the near-infrared region ( $K$ ), oxyhemoglobin concentration ( $[HbO_2]$ ), and deoxyhemoglobin concentration ( $[HHb]$ ).

### 3. Materials and Methods

This section presents the description of the probe design and development, the instrumentation and acquisition system, and the circuits implemented in each of them. Finally, the performance tests, using both the probe and the system to detect PPG and non-pulsatile signals from the cerebral arteries of a head phantom, are presented.

#### 3.1. Probe

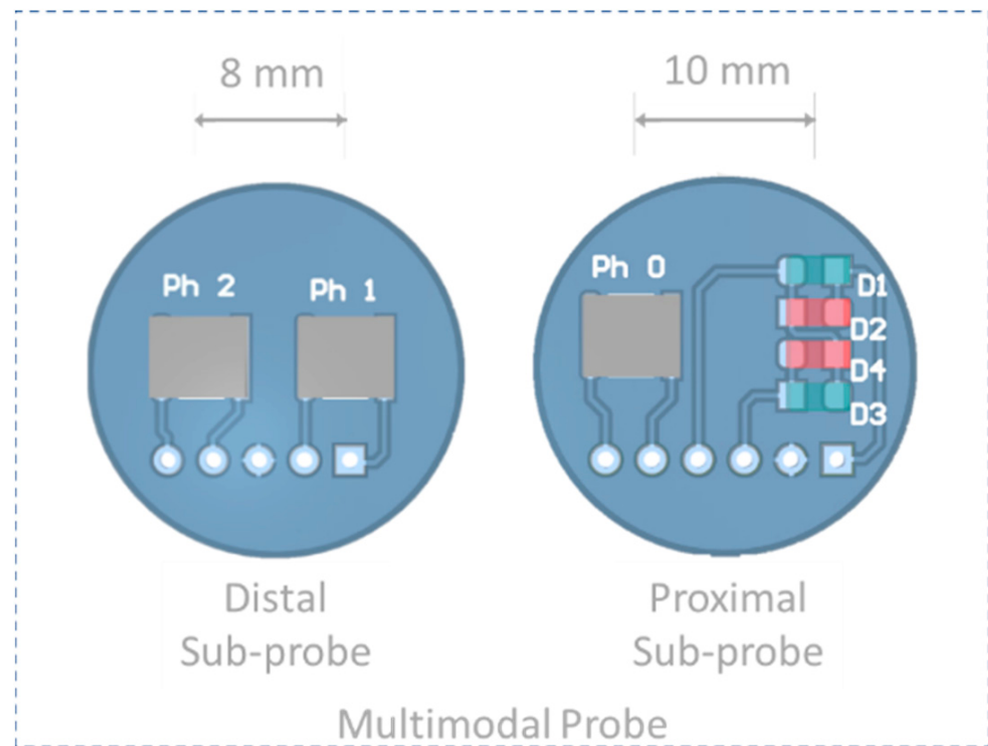
The sensor's probe design considered, as many other optical modalities (Photoplethysmography (PPG), NIRS, among others), light emitters to shine into the tissue and photodetectors to acquire the backscattered light. The quantity of these optical components, their topology, and their placement, were defined based on the sensor's application and the anatomy location. Firstly, the sensor's application is to interrogate non-invasively the brain using light and to estimate changes in ICP and cerebral oxygenation. Secondly, the sensor's location is on the right or left side of the forehead, avoiding the midline sinuses. Implementing NIRS principles to PPG measurements opens up great opportunities for extending the capabilities of both techniques. By doing so, the same sensors and technology can be used to obtain parameters relating to tissue oxygenation and intracranial pressure. Therefore, in this research, four wavelength emitters and three photodetectors were used in a reflectance mode probe to apply photoplethysmography and NIRS physical principles.

##### 3.1.1. Optical Parts

Four light-emitting diodes (LEDs), also known as sources, were carefully chosen to interrogate specific information from the brain. The wavelengths used are 660 nm, 770 nm, 810 nm, and 880 nm. The 810 nm wavelength, also known as the isosbestic wavelength, has the unique characteristic of being absorbed to the same extent by both oxy- and deoxy-hemoglobin [18]. Therefore, it is an optimal wavelength to record an optical signal which is independent of blood oxygenation and could be correlated to intracranial volumetric changes.

Wavelengths at 660 nm and 880 nm were also included in order to assess extracerebral oxygen saturation. The utilization of these wavelengths in pulse oximetry ( $SpO_2$ ) is widely reported [18]. Likewise, NIRS commercial devices generally use at least two infrared wavelengths (750–1000 nm) to calculate cerebral oxygenation parameters. These two wavelengths are usually selected on opposite sides with respect to the isosbestic point of the hemoglobin absorption spectra (810 nm). Accordingly, LEDs of 770 nm and 880 nm were included in the probe to implement spatially resolved spectroscopy (SRS) measurements.

Previous evaluation of infrared light and cerebral tissue interaction at different source-detector (S-D) separations showed that higher S-D separation increases the penetration depth, but it also causes a decrease in the overall signal quality due to high absorption [21]. Accordingly, the probe design was subdivided into two individual sub-probes called proximal probe and distal probe (Figure 1). As this is a research probe, the proposed configuration will enable the flexibility to investigate the effect of S-D on optical signals on demand.



**Figure 1.** PCB design of the multimodal probe showing both proximal and distal sub-probes, with their respective optical components.

The proximal sub-probe consists of the four LEDs (D1–D4), and one photodiode (Ph 0) placed 10 mm from the sources to detect the backscattered light from superficial tissues, which will allow arterial oxygen saturation ( $SpO_2$ ) measurements. The distal sub-probe includes two photodiodes (Ph 1 and Ph 2) to detect the backscattered light from cerebral tissues, which can be separated from the proximal probe between 25 and 50 mm. The sources and distal photodiodes will allow the evaluation of cerebral oxygenation parameters through SRS.

Photoplethysmograms (PPG) from one of the distal photodiodes are acquired to obtain volumetric information from the cerebral tissue. However, due to the spatial sensitivity profile of a reflectance probe, it is likely that the signals are contaminated by extracerebral information. Therefore, extracerebral volumetric signals are also acquired by the proximal photodiode in order to subtract them from the contaminated brain signals. The latter should result in the acquisition of only cerebral pulsatile signals for the further estimation of ICP from changes in their morphological features.

Table 1 describes the optical and electrical characteristics of the optical components selected to build the proposed multimodal probe.

**Table 1.** Optical characteristics of the light source and detectors.

LEDs					
Peak wavelength (nm)	660	770	810	880	
FWHM (nm)	22	24	28	30	OV < 20
Forward voltage (V)	1.7	1.6	1.5	1.3	2 < OV < 4
Radiant intensity (mW/sr)	3.5	3.5	12	28	OV ≈ 30
Viewing angle (deg)	40	40	40	40	OV ≈ 30
Forward current (mA)	30	50	30	50	OV ≈ 20
Photodetectors					
Spectral bandwidth (nm)	430–1100				
Forward voltage (V)	1				
Reverse dark current (nA)	2				
Capacitance (pF)	25				1 < OV < 100
Angle half sensitivity (deg)	±65				
Rise and fall time (ns)	100 each				
Material and housing	Silicone/planar				

OV: optimal value for PPG applications [18].

### 3.1.2. Printed Circuit Board (PCB) Design

The PCBs for the probe and the processing system were designed using an electronic design automation software package known as Altium Designer (Altium Limited, Sydney, Australia) and were manufactured using a Computer Numeric Control machine (Elecrow, Shenzhen, China). Each sub-probe PCB has a circular shape, with a diameter  $d = 21$  mm and boards of 1.6 mm thickness, in order to facilitate good contact with the forehead.

The four LEDs were placed in the proximal probe in a two-pairs configuration. Each pair of LEDs was connected antiparallel; this configuration permits the intermittent switching of the four LEDs when connected to the processing unit. The photodiodes were positioned over the middle horizontal axis of the sub-probes, aligned with each other.

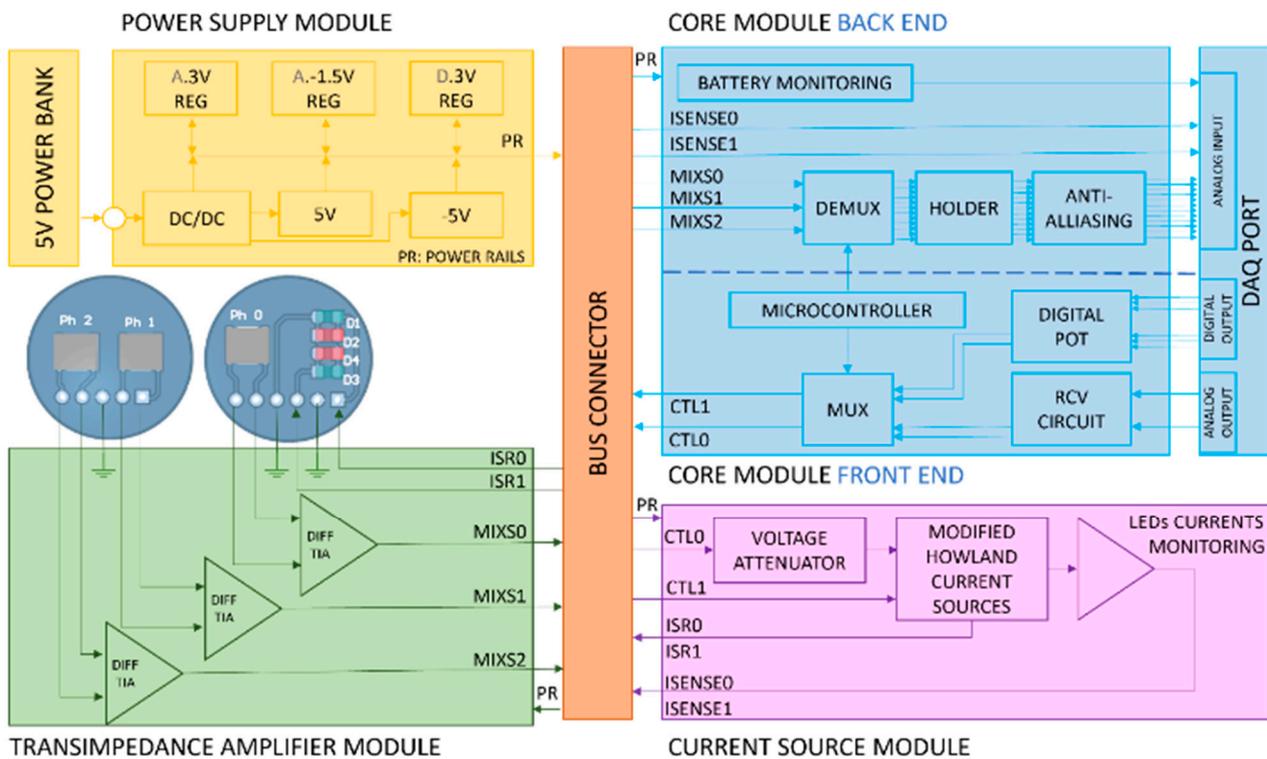
### 3.1.3. Probe Case Design

Since the probe needs to be in contact with the skin, the optical components and PCBs are required to be isolated and protected to avoid short circuits, shunting (i.e., light reaching the photodiode without passing through the tissue), skin burns, or marks. Additionally, without an appropriate case, the photodiodes can be reached by ambient light, increasing dark currents and interference noise. Therefore, the following mechanical structure aimed to shield the optical components and facilitate the contact of the sensor on the skin. The probe case was designed in a 3D CAD design software (Solidworks 2021, Cambridge, UK) and 3D printed using a Formlabs black resin (Somerville, MA, USA). The design of each sub-probe case comprises a compartment and a cover. The two parts were designed and separately manufactured in order to enclose the PCB. Moreover, the cover of the proximal and the distal probes have different designs according to the optical components involved in them. A clip system at the back of each sub-probe case allows the attachment of the sub-probes to a snug-fitting headband. The band fits a range of sizes and has multiple marks to slide the sub-probes to different S-D distances. Both cases are detachable.

## 3.2. Instrumentation and Acquisition System

The processing system described in this section is called ZenTBI and consists of a four-wavelength, three-channel research PPG system composed of five independent and interchangeable subsystems. Moreover, three external devices are connected to the ZenTBI, namely, a 5 V power bank (RavPower, Shenzhen, China), a NI data acquisition card (DAQ) (National Instruments, Austin, TX, USA), and the probe. Figure 2 shows the architecture of the ZenTBI. The dimensions of all the modules are  $100 \times 80 \times 1.6$  mm, except for the current source module and the bus connector. The dimensions of these two boards are  $100 \times 60 \times 1.6$  mm and

100 × 48 × 1.6 mm, respectively. All modules are double-sided copper-clad boards with a HASL surface finish that is generally preferred for hand-soldering.



**Figure 2.** Detailed block diagram showing the ZenTBI architecture, considering the power supply module, core module, current source module, transimpedance amplifier module, and bus board.

### 3.2.1. Bus Connector

This allows the connection between a maximum of six modules, three at each layer. The modules are connected to receptacle 50-way surface mount connectors in which every pin at each connector corresponds to the same pin in all other connectors. Every pin has been assigned an unalterable net to ensure the transmission of the voltages and signals between boards.

### 3.2.2. Power Supply Module

This module is connected through a micro-USB port to a 5 V power bank with a capacity of 26,800 mAh (RavPower, Shenzhen, China). The board consists of the circuitry required to regulate the USB power supply into multiple DC power rails. The initial power bank supply (PS) passes through an isolated point-of-load DC–DC converter (ITX0505S), regulating the initial voltage to IPOS (5 V) and INEG (−5 V). This dual source powers the microchips on the Current source module and the multiplexers and demultiplexers included in the Core module. The DC power supply from the dual source is regulated down to APOS (3.3 V) and ANEG (−1.5 V) in order to power the remaining microchips on the processing system. However, from the DC-DC converter output, a separate digital power line is regulated down to DPOS (3.3 V) and has its own digital ground (DGND). The later power line supplies the microcontroller to avoid additional digital noise in the analogue signals.

### 3.2.3. Core Module Front End

The front end is responsible for the intermittent light switching control that allows the sampling of four wavelengths simultaneously. This part of the core board hosts four independent controls to set the emitters' currents and a multiplexing circuit. Previous

studies on reflectance PPG demonstrated that acceptable signal quality could be guaranteed without exceeding 50 mA LEDs driving currents [22]. Accordingly, each current control was designed to allow a maximum drive current of 100 mA. The first and second controls are the DAQ analogue outputs (A0 and A1) connected to voltage attenuator circuits (RCV). Thus, from LabVIEW, A0 is set between 0 V and 5 V, while A1 can be defined between −5 V and 0 V. Both outputs were then attenuated down to 0 V to 1 V and −1 V to 0 V, respectively, using voltage dividers and op-amp buffers. These controls pass through a second attenuation phase at the beginning of the Current source module to finally obtain 0 to 100 mV and −100 mV to 0 V, respectively. Due to the limited number of analogue output channels on the DAQ card, the third and fourth controls consist of 10 K digital potentiometers (X9C103SIZT1) configured into two independent voltage dividers. One of them is supplied with 3.3 V and configured with a fixed resistor of 23 K, allowing changes between 0 and 1 V. The other is supplied with −1.5 V and configured with a fixed resistor of 5 K, creating the last output of −1 to 0 mV. Each digital potentiometer is controlled from LabVIEW, where three digital signals are generated to activate (CS) the chip, toggle the INC pin to move the wiper, and either increment or decrement the counter in the direction indicated by the logic level on pin U/D.

The multiplexing circuit generates timed switching signals used to turn on one LED at a time, which allows the independent sampling of light at each wavelength by photodetectors. All four current controls and the analogue ground (0 V) are connected to a double-pole four-throw multiplexer (MC14052BD). Each multiplexor pole selects one of the three input voltage signals (either positive or negative or ground) and outputs the selected input into a single line, producing two control lines called CTL0 and CLT1. Each control line switches a pair of LEDs. When CTL0 is either positive or negative, CTL1 output is ground (0 V), and vice versa. Due to the antiparallel position of the LEDs in each pair, when one of the control lines is positive, only one LED of the pair, controlled by that control line, is switched “ON,” and when negative, the other LED of the same pair is switched “ON.” The multiplexer switching time is controlled by an 8-bit Atmel ATtiny 2313–20SU microcontroller unit (MCU) (Atmel Corp, San Jose, CA, USA) at a frequency of 1 kHz. A byte data sequence is created using the multiplexer’s truth table and is written on port-D of the MCU.

The two control lines of the multiplexer are passed through the bus connector into the current source module, where the voltage is converted into the current required to power the LEDs. The duration time of each LED “ON” is calculated by Equation (6). Having a sampling frequency of 1000 Hz, three photodiodes, and four wavelengths, each emitter turns “ON” every 2.5 ms.

$$t = \frac{1/F_s}{n_o.photodiodes \times n_o.wavelengths} \quad (6)$$

Equation (6): Duration time of each LED “ON” ( $t$ ), sampling frequency ( $F_s$ ), number of photodiodes ( $n_o.photodiodes$ ), and number of wavelengths ( $n_o.wavelengths$ ).

### 3.2.4. Current Source Module

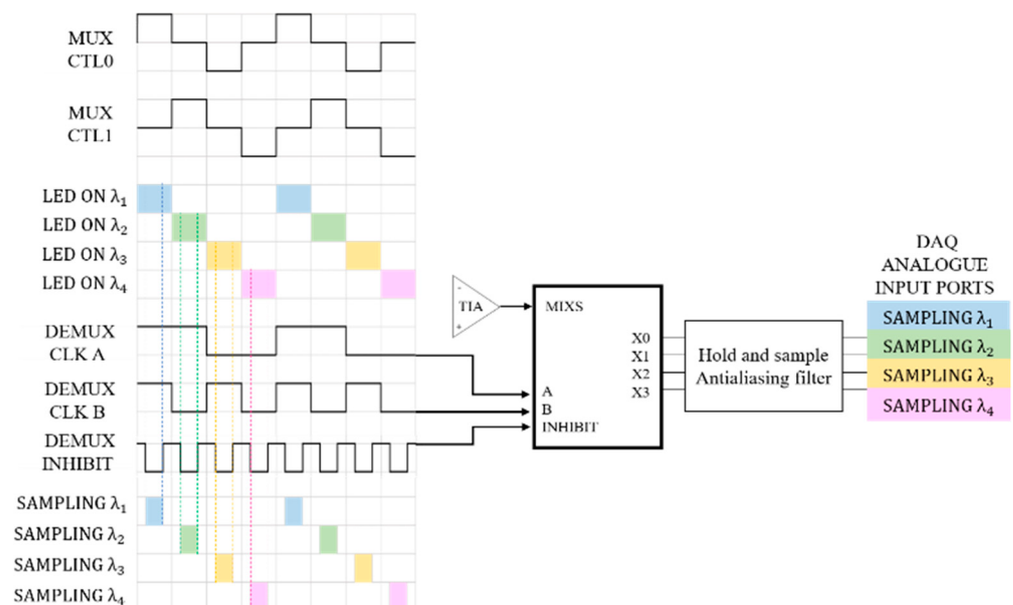
This module has two main functions: (1) to limit the current driven through the LEDs to a maximum of 100 mA, avoiding damages in the probe and causing potential skin burns for overheating the emitters; and (2) To measure the current across each LED, providing feedback to control the LED currents. In order to accomplish the first function, the module involves two identical and improved Howland current sources followed by a push–pull (npn–pnp) transistor switch. The Howland is unique and interesting because it has both positive and negative feedback paths, as is shown in Figure 3. When the array of resistance balances the feedback paths ( $R_2/R_1 = R_4/R_3$ ), the operational amplifier behaves as a linear voltage-controlled current source [23].



DC outputs of both photodiodes were compared using a homogeneous phantom to set equal gains.

### 3.2.6. Core Module Back End

The core module back end is mainly responsible for separating the mixed signals from each TIAs output into four independent raw signals containing information at specified wavelengths. As mentioned earlier, the port-D of the microcontroller is used to generate four clocks for multiplexing the LEDs. However, port-B of the MCU is programmed to output another three clocks used for synchronizing the demultiplexer, which separates the mixed signals into independent wavelength components. Clocks A and B control the internal switches of the demultiplexer to separate the mixed signal into four outputs. As the clocks of both ports D and B are synchronized, it is possible to sample a specific wavelength at each demux output. However, when an LED is switched from one state to the other (ON/OFF), there is a slight delay between the transitions, called the transient response of a diode. Therefore, by sampling the entire LED activation period, the switching noise is also sampled. To overcome this problem, the demultiplexer inhibits control and is activated during these transitions, limiting the sampling period only to the middle of the signal, where it is expected that the signal has enough time to settle before sampling (Figure 4).



**Figure 4.** LEDs multiplexing and photodetectors' signals demultiplexing. Each color box represents a wavelength.

Moreover, it is necessary to 'hold' the sampled voltage until the next clock pulse triggers a new acquisition. Accordingly, a hold capacitor is placed at the output of the demux ports, followed by a buffer circuit that offers a high impedance to the hold capacitor to keep the held voltage from discharging prematurely. Previous studies have tested different hold capacitors for PPG signal sampling, establishing that a 220-nF capacitor provides an optimal performance [18].

After the hold buffer, individual antialiasing filters with a cut-off frequency of 80 Hz were implemented. These passive RC filters limit the bandwidth of the signals and stop the high-frequency switching noise from corrupting them. Every low-pass filter output is the input of an op-amp buffer before the signals are passed onto the NI DAQ card for digitization and further analysis. The raw signals are sampled at 2 kHz and contain both AC and DC components, which are later separated digitally using LabVIEW digital filters.

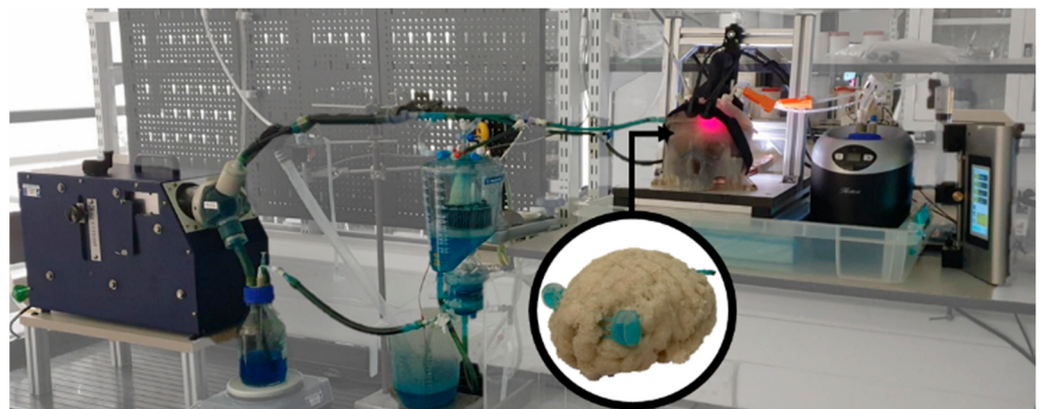
Finally, the core module back end also allows the digitalization of the battery's state and LED's current monitoring, which are displayed on the LabVIEW interface to give

the user an indication of the battery's state of charge and to adjust the LED's intensity on demand.

### 3.3. Technical Evaluation

The technical evaluation consisted of three sections. Firstly, the input and output of each of the ZenTBI modules were evaluated using an Oscilloscope. Secondly, the power consumption was calculated by measuring the voltage over a 1-ohm resistor placed in series between the battery and the ZenTBI supply input. According to Ohm's law, the voltage measured with a digital multimeter corresponded to the current consumption ( $I = V/R$ ).

Finally, a quality evaluation of the optical signals was conducted in MATLAB to demonstrate the functionality of all the components of the system. A head phantom was used to simulate pulsations on the cerebral arteries that could be measured by the optical sensor [24], as is shown in Figure 5. The use of a head phantom rather than a human head allows the control of multiple parameters such as the oxygen levels and the intracranial pressure. Moreover, the lack of scalp or extracerebral circulation ensures the acquisition of signals only from the cerebral arteries, reducing this confounding factor. Moreover, Table 2 presents the parameters of the setup.



**Figure 5.** The head phantom consists of a brain, skull, cerebrospinal fluid, and blood circulation. The sensor on top of the forehead shines light to reach the pulsatile vessels of the brain and receives light that correlates to oxygenation and ICP.

**Table 2.** Quality evaluation setup parameters.

Protocol	Parameter	Value	Units
Sensor	Recording time	60	Sec
	LEDs currents	30	mA
	Sample frequency	2000	Hz
	Probes distance	5	mm
	Location	Phantom's forehead	
Phantom	Circulating fluid	Artificial blood	65%
	Heart rate	60	bpm
	Stroke	20	cc/stroke
	Intracranial pressure	15	mmHg

The analysis included a pre-processing step, where the signals from all wavelengths and both proximal and distal photodetectors were filtered using a Butterworth bandpass filter of order 3 ( $f_c$ : 0.5 to 8 Hz). Then, a 60-s recording was segmented into 5-s windows, where nine different signal quality indices were calculated based on the literature [25,26]. The list of indices is perfusion, kurtosis, skewness, kurtosis in the frequency domain, skewness in the frequency domain, entropy, zero-crossing rate, signal-to-noise-ratio (SNR),

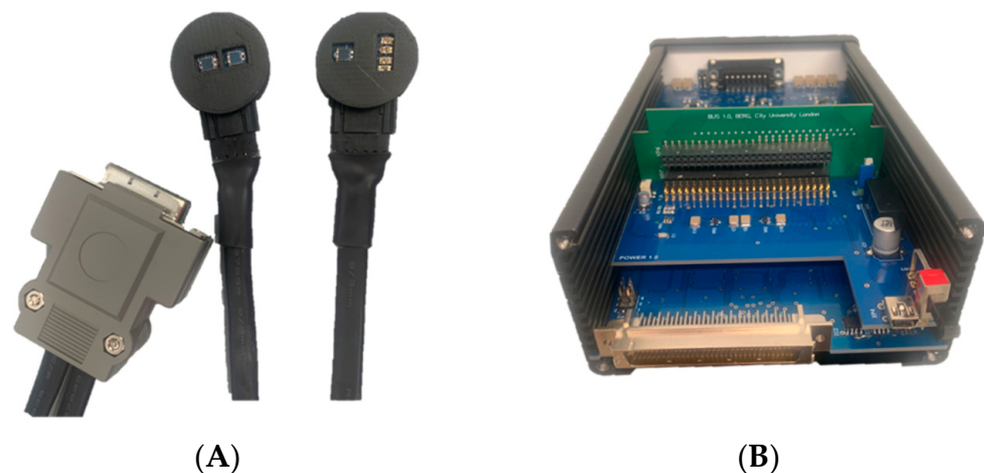
and relative power. Outliers were replaced by the median in all the indices. Finally, the comparison of proximal and distal signal quality at all wavelengths was made by a non-parametric ANOVA (Kruskal–Wallis), considering a confidence level of 95%.

#### 4. Results and Discussion

This section describes the ZenTBI system and its performance in acquiring PPG signals from both photodetectors.

##### 4.1. Probe and Instrumentation System

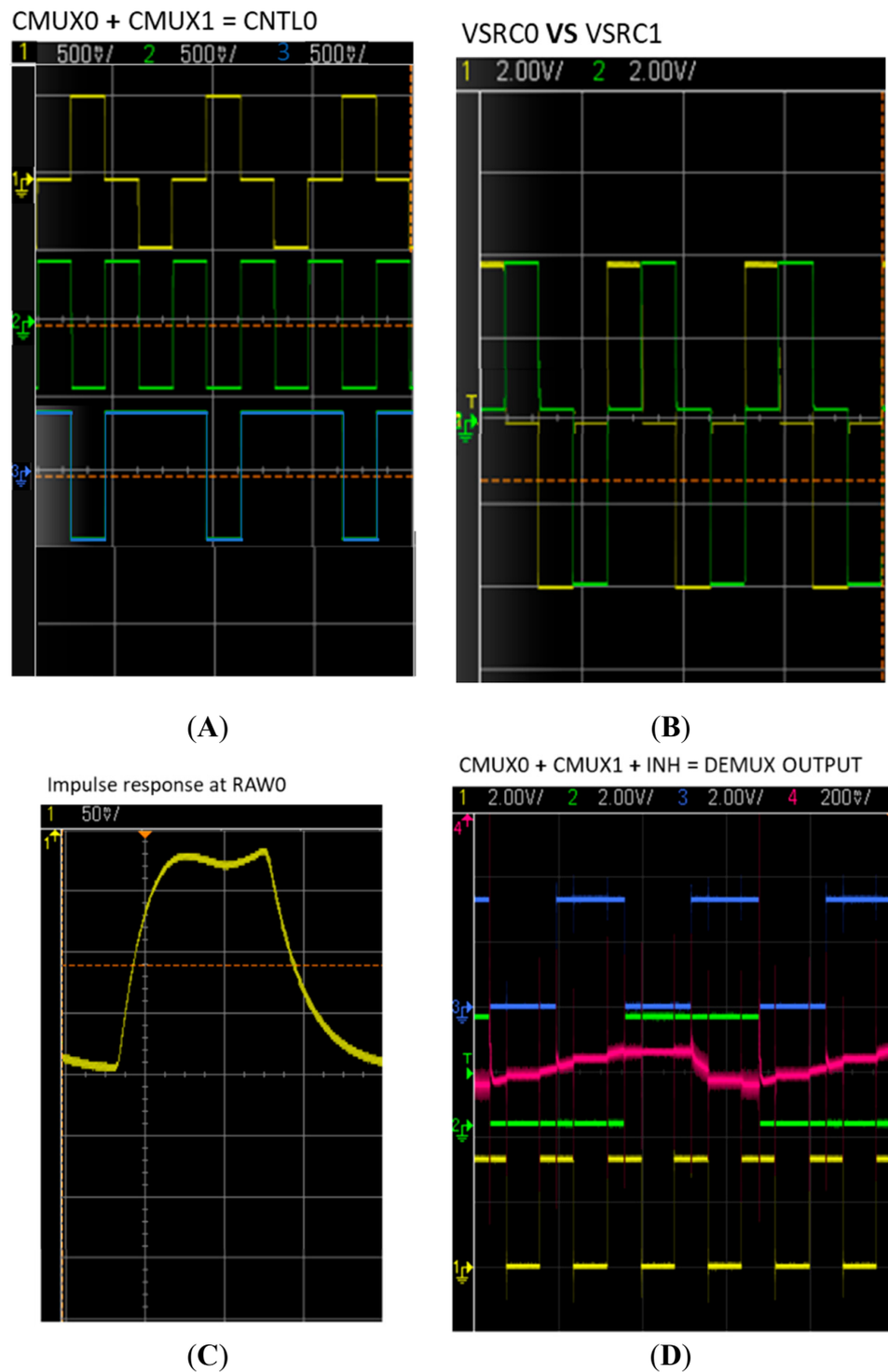
Figure 6 displays the probe prototype with its instrumentation system which allows the multiplexing of four different light sources and the acquisition of the reflected light into three photodiodes. The ZenTBI has to be connected to a DAQ for the visualization and recording of the signals in LabVIEW 2021, where the current of the LEDs can be adjusted. An HP EliteDesk computer with Intel® i7, ×64 and 32 GB RAM allowed the successful implementation of LabVIEW for the current application.



**Figure 6.** (A) Probe prototype with its 4 LEDs, 3 photodiodes and 15D connector. (B) Open ZenTBI view of the instrumentation system.

##### 4.2. Technical Evaluation

In order to demonstrate the correct performance of the developed system, this section presents the results of the technical evaluation. Firstly, the debugging of the modules using the oscilloscope upholds the expected behaviour of the printed boards. For instance, Figure 7A shows the outputs of the microcontroller CMUX0 in blue and CMUX1 in green. In accordance with the multiplexor's truth table, CMUX0 and CMUX1 generate CNTL0, shown in yellow. The positive and negative amplitudes of the yellow signal are defined by the DAQ analogue outputs, and the digital potentiometers previously described. The amplitude of these pulses controls the LED currents using a Howland pump, the output of which is shown in Figure 7B. Besides the LEDs' supply, the response of each photodiode was tested at the output of the transimpedance amplifier. An impulse response is shown in Figure 7C, where the photodiode was covered from light and exposed very quickly.



**Figure 7.** ZenTBI debug tests on the oscilloscope. (A) shows the outputs of the microcontroller CMUX0 in blue and CMUX1 in green. In accordance with the multiplexor’s truth table, CMUX0 and CMUX1 generate CNTL0, shown in yellow. (B) shows the output of the Howland pump circuits that control the LEDs’ currents. Two LEDs are controlled by the yellow pulses and other two by the green pulses. (C) shows the impulse response of a photodetector. (D) shows the demux controls generated by the microcontroller, CMUX0 as the green line, CMUX1 as the purple line, and the INHIBIT as the yellow line.

Finally, the demultiplexer was evaluated to ensure the effective decomposition of the mixed signal (from all four wavelengths) into four independent signals. Figure 7D displays the demux controls generated by the microcontroller, CMUX0 as the green line, CMUX1 as the purple line, and the INHIBIT as the yellow line. The output of the demultiplexer is the pink signal that becomes erratic every time the INHIBIT is ON; a constant voltage can then be seen at the different demux combinations according to its true table. The presented debug sequence demonstrates the correct operation of the main ZenTBI stages.

During the power consumption assessment, two measurements took place: one without the probe and the second one with the probe connected to the system. The LEDs in the second measurement were driven by 40 mA of current. The load current calculated using Ohms Law was 360 mA without the probe and 420 mA with the system complete (probe connected). From the second measured load current, the battery life of the entire system can be calculated (Equation (7)), and is given as:

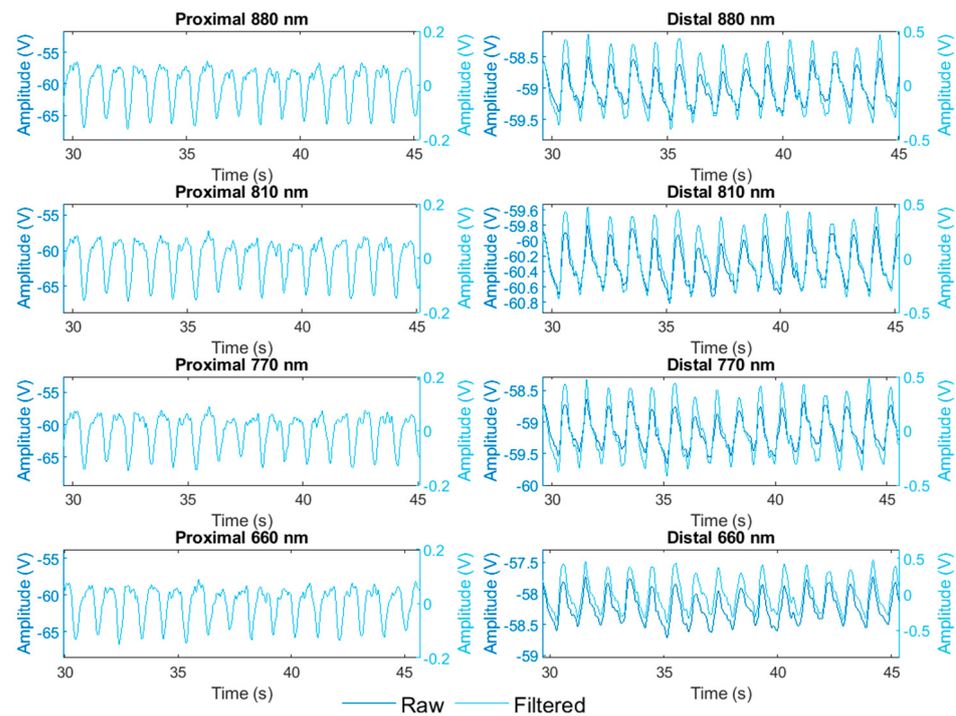
$$\begin{aligned} \text{Battery life} &= \frac{\text{Battery capacity}}{\text{Load current}} \times 0.7 \\ \text{Battery life} &= \frac{26,800 \text{ mAh}}{420 \text{ mA}} \times 0.7 \\ \text{Battery life} &= 45 \text{ hours} \end{aligned} \quad (7)$$

The system aims to assess cerebral oxygenation parameters by computing the DC component of the photodiodes placed at the distal probe. However, arterial saturation from the scalp could also be calculated from both AC and DC PPG components of the photodiode at the proximal probe. Moreover, the system also aims to utilize the NIRS-PPG signals of the proximal photodiode at the distal probe for the estimation of intracranial pressure. Therefore, the consistent acquisition of the DC and AC components from the respective photodiodes is crucial for the proposed applications of the developed probe. The system's performance was validated by the quality assessment of the PPG signals acquired from the proximal detector (proximal probe) and the distal detector (proximal photodiode of the distal probe).

Figure 8 displays the raw and filtered PPGs signals from the detectors of interest at all wavelengths. From this Figure, it can be concluded that the acquisition of PPG signals from both detectors at all wavelengths (all LEDs currents were equal) was successful.

Table 3 presents the median and interquartile ranges of the quality indices calculated from the signals displayed in Figure 8. The perfusion index is the gold standard for assessing PPG signal quality, and it is calculated as the ratio of the pulsatile blood flow to the non-pulsatile (DC) [25]. Perfusion indices below 0.4% show weak pulse strength, which is the case for the proximal signals, as it is expected due to the lack of extracerebral perfusion (phantom). The perfusion index was significantly different between detectors as well as it differs between wavelengths.

Skewness measures how symmetric the signal's distribution is and kurtosis measures how sharp the peak of the signal's distribution is. A distribution with many outliers will have low values of skewness and kurtosis since its probability distribution will be asymmetric, flatter and will approach zero slower [27]. Hence, there are more erratic peaks on the distal signals than the proximal, with a statistically significant difference. However, the skew and kurtosis on the frequency domain were the same at all levels. Entropy provides a quantitative measure of the uncertainty present in a signal [25]. The lower values ( $E < 5$ ) correlate to low noise which is the case for both proximal and distal signals despite the significant difference between the detectors and wavelengths. Another important index on pulsatile signals quality is the zero-crossing rate, which, on PPG signals, is calculated as the rate of sign-changes with the signal mean as the reference. On a window of 5 s (10,000 samples) and simulating a heart rate of 60 bpm, the results showed a zero-crossing rate of 0.0010, as was expected. The latter demonstrated the absence of erratic peaks crossing the mean signal.



**Figure 8.** Raw and filtered PPG signals from two photodiodes using the head phantom at a normal ICP of 15 mmHg.

**Table 3.** Signal quality indices comparison of the detectors and wavelengths.

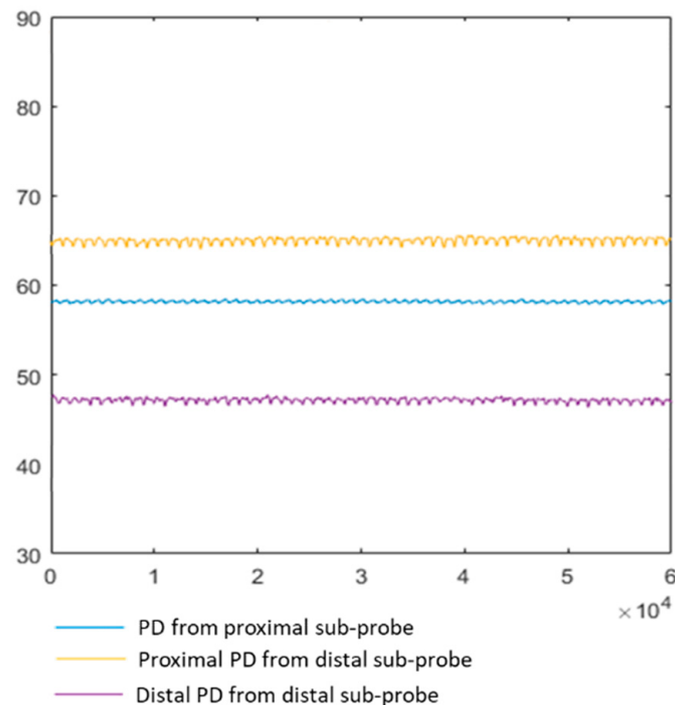
Quality Index	Detector	880 nm		810 nm		770 nm		660 nm		Wavelength		Detector	
		Me	RI	Me	RI	Me	RI	Me	RI	<i>p</i> -Value	*	<i>p</i> -Value	*
Perfusion index	Proximal	0.31	0.03	0.29	0.03	0.28	0.02	0.31	0.02	0.00	*	0.00	*
	Distal	1.41	0.08	1.40	0.05	1.45	0.04	1.41	0.08	0.01	*	0.00	*
Skewness	Proximal	0.84	0.04	0.85	0.05	0.84	0.06	0.83	0.05	0.51		0.00	*
	Distal	0.23	0.15	0.23	0.13	0.22	0.14	0.22	0.13	0.95		0.00	*
Kurtosis	Proximal	2.33	0.19	2.36	0.22	2.39	0.26	2.38	0.16	0.69		0.00	*
	Distal	1.89	0.06	1.90	0.09	1.90	0.11	1.90	0.13	0.79		0.00	*
Skewness freq. domain	Proximal	2.47	0.00	2.47	0.00	2.47	0.00	2.47	0.00	0.92		0.51	
	Distal	2.47	0.00	2.47	0.00	2.47	0.00	2.47	0.00	0.11		0.51	
Kurtosis freq. domain	Proximal	7.12	0.00	7.12	0.00	7.12	0.00	7.12	0.00	0.92		0.51	
	Distal	7.12	0.00	7.12	0.00	7.12	0.00	7.12	0.00	0.11		0.51	
Entropy	Proximal	2.09	0.09	2.08	0.07	2.09	0.08	2.11	0.11	0.98		0.00	*
	Distal	2.24	0.03	2.24	0.03	2.24	0.03	2.24	0.04	0.96		0.00	*
Zero crossing rate	Proximal	0.0010	0.0000	0.0010	0.0000	0.0010	0.0002	0.0010	0.0000	0.00	*	0.00	*
	Distal	0.0010	0.0002	0.0010	0.0002	0.0010	0.0002	0.0010	0.0000	0.08		0.00	*
Signal-to-noise ratio	Proximal	23.82	2.71	22.76	9.23	22.33	7.79	22.05	1.99	0.61		0.00	*
	Distal	24.22	4.37	24.86	6.92	27.56	7.59	28.12	8.76	0.12		0.00	*
Relative power	Proximal	0.76	0.02	0.76	0.02	0.76	0.02	0.75	0.02	0.97		0.01	*
	Distal	0.74	0.02	0.74	0.03	0.75	0.03	0.74	0.03	0.97		0.01	*

\* Significant difference defined by a *p*-value < 0.05.

The signal-to-noise ratio (SNR) is a well-used measure that compares the level of the desired signal to the level of background noise [25]. The higher the ratio, the lower the noise. The resultant median ratios over 22 dB indicate the low proportion of the noise against the PPGs signals. Better ratios were found on the distal signals with a significant difference on at least one of the wavelengths compared to the proximal measurements. Finally, it has been found that most of the energy of the systolic and diastolic waves is concentrated within the 1–2.5 Hz frequency band. The ratio of the power spectral density

(PSD) in this band compared to the PSD in the overall signal 0–8 Hz, provides a measure of the signal quality (Expected value > 0.85) [28]. This index was around 0.75 for all the signals included in the analysis, which is good enough, though it is not sufficient to do signal comparison.

In addition to the analysis of the pulsatile signals from the proximal and distal detectors, the acquisition of the DC components from all three photodiodes is needed for the sensor's applications. For instance, Figure 9 shows the DC level recorded from all photodetectors at 810 nm. Further evaluation, changing the phantom's fluid oxygen level, will assess the response of the DC signals for the subsequent calculation of oxygenation parameters.



**Figure 9.** Light intensity DC from all three photodetectors at 810 nm.

The ZenTBI system fulfilled the design requirements for acquiring PPG and NIRS signals, which were methodically evaluated in an in vitro setup. This study showed the morphology of the pulsations and calculated some quality indices to evaluate the signals. Even though the signals' quality depends on the subject (in vitro/in vivo), the PPG signals acquired from the phantom using artificial blood successfully led to the extraction of quality measurements. Still, there is always room for improvement in the acquisition system design. Other authors have reported techniques to reduce the impact of ambient light, signal saturation, and movement artefacts in pulsatile signals [29,30], which could be added to the current ZenTBI system to reduce common noise sources. For instance, an ambient light cancellation circuit would avoid signal saturation due to high levels of ambient light or small gaps between the sensor and the measuring surface. Moreover, the separation of the AC and DC components using analogue filters could allow higher amplification of the pulsatile signals to improve the quality of these signals, as well as the implementation of an offset cancelling circuit to reduce the movement artefacts in the pulsatile signals.

## 5. Conclusions

This research has presented the design and manufacture of an optical sensor that would allow the identification of changes in cerebral oxygenation and the estimation of intracranial pressure by the analysis of the pulsatile and non-pulsatile NIRS signals acquired from cerebral tissue. The system includes a probe designed specifically for monitoring the main physiological parameters of interest in TBI patients. This manuscript presents the development and technical evaluation of an instrumentation system for the acquisition of proximal and distal PPG signals at multiple wavelengths. Different source-detector separations and multiple wavelengths are key for the assessment of superficial and deeper tissue layers and provide information to calculate oxygenation parameters and SpO<sub>2</sub> and TOI. The sensor has the capacity to acquire good quality signals (AC and DC) from the photodiodes and hence enable their implementation in further computational analysis. Although NIRS has been implemented before for cerebral oxygenation, future work will include the analyses of the cerebral PPG signals' features, using this sensor and the phantom, which could open the possibility for a multimodal non-invasive tool to assess TBI patients.

## 6. Patents

Device and method for non-invasive estimation of intracranial pressure based on cerebral PPG signals from the pulsatile component of NIRS: European office (EP4061204A1) and United States office (US 2022/0409080 A1).

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## References

1. Brain Injury Association of America of ABI vs. TBI: What Is the Difference? Available online: <https://www.biausa.org/brain-injury/about-brain-injury/nbiic/what-is-the-difference-between-an-acquired-brain-injury-and-a-traumatic-brain-injury> (accessed on 8 January 2021).
2. Headway Types of Brain Injury | Headway. Available online: <https://www.headway.org.uk/about-brain-injury/individuals/types-of-brain-injury/> (accessed on 8 January 2021).
3. Maas, A.I.R.; Menon, D.K.; Adelson, P.D.; Andelic, N.; Bell, M.J.; Belli, A.; Bragge, P.; Brazinova, A.; Büki, A.; Chesnut, R.M.; et al. Traumatic brain injury: Integrated approaches to improve prevention, clinical care, and research. *Lancet Neurol.* **2017**, *16*, 987–1048. [[CrossRef](#)] [[PubMed](#)]
4. Roldan, M.; Abay, T.Y.; Kyriacou, P.A. Non-invasive techniques for multimodal monitoring in Traumatic Brain Injury (TBI): Systematic review and meta-analysis. *J. Neurotrauma* **2020**, *37*, 2445–2453. [[CrossRef](#)]
5. Mazzeo, A.T.; Gupta, D. Monitoring the injured brain. *J. Neurosurg. Sci.* **2018**, *62*, 549–562. [[CrossRef](#)] [[PubMed](#)]
6. Carney, N.; Totten, A.M.; O'Reilly, C.; Ullman, J.S.; Hawryluk, G.W.J.; Bell, M.J.; Bratton, S.L.; Chesnut, R.; Harris, O.A.; Kisson, N.; et al. *Guidelines for the Management of Severe Traumatic Brain Injury*, 4th ed.; Brain Trauma Foundation: New York, NY, USA, 2017; Volume 80.
7. Ko, S.-B. Multimodality Monitoring in the Neurointensive Care Unit: A Special Perspective for Patients with Stroke. *J. Stroke* **2013**, *15*, 99. [[CrossRef](#)]
8. Zhang, X.; Medow, J.E.; Iskandar, B.J.; Wang, F.; Shokouejad, M.; Koueik, J.; Webster, J.G. Invasive and noninvasive means of measuring intracranial pressure: A review. *Physiol. Meas.* **2017**, *38*, R143–R182. [[CrossRef](#)] [[PubMed](#)]
9. Harary, M.; Dolmans, R.G.F.; Gormley, W.B. Intracranial pressure monitoring—Review and avenues for development. *Sensors* **2018**, *18*, 465. [[CrossRef](#)]

10. Benni, P.B.; MacLeod, D.; Ikeda, K.; Lin, H.M. A validation method for near-infrared spectroscopy based tissue oximeters for cerebral and somatic tissue oxygen saturation measurements. *J. Clin. Monit. Comput.* **2018**, *32*, 269–284. [[CrossRef](#)]
11. Davie, S.N.; Grocott, H.P. Impact of Extracranial Contamination on Regional. *Anesthesiology* **2012**, *116*, 834–840. [[CrossRef](#)]
12. Patil, A.V.; Safaie, J.; Moghaddam, H.A.; Wallois, F.; Grebe, R. Experimental investigation of NIRS spatial sensitivity. *Biomed. Opt. Express* **2011**, *2*, 1478. [[CrossRef](#)]
13. Fischer, G. EQUANOX Advance™ Sensor. New York. 2011. Available online: [www.noninequanox.com](http://www.noninequanox.com) (accessed on 6 January 2023).
14. Corporation Near Infrared Oxygenation Monitor NIRO-200NX. Available online: [www.hamamatsu.com](http://www.hamamatsu.com) (accessed on 6 January 2023).
15. Redford, D.; Paidy, S.; Kashif, F. Absolute and Trend Accuracy of a New Regional Oximeter in Healthy Volunteers During Controlled Hypoxia. *Anesth. Analg.* **2014**, *119*, 1315–1319. [[CrossRef](#)]
16. Roldan, M.; Kyriacou, P.A. Near-Infrared Spectroscopy (NIRS) in Traumatic Brain Injury (TBI). *Sensors* **2021**, *21*, 1586. [[CrossRef](#)]
17. Abay, T.Y.; Phillips, J.P.; Uff, C.; Roldan, M.; Kyriacou, P.A. In Vitro Evaluation of a Non-Invasive Photoplethysmography Based Intracranial Pressure Sensor. *Appl. Sci.* **2022**, *13*, 534. [[CrossRef](#)]
18. Kyriacou, P.A.; Allen, J. *Photoplethysmography Technology, Signal Analysis and Applications*; Kyriacou, P.A., Allen, J., Eds.; Elsevier: London, UK, 2021; ISBN 9780128233740.
19. Ysehak Abay, T. Reflectance photoplethysmography for non-invasive monitoring of tissue perfusion. *IEEE Trans. Biomed. Eng.* **2016**, *62*, 2187–2195. [[CrossRef](#)]
20. Suzuki, S.; Takasaki, S.; Ozaki, T.; Yukio, K. A tissue oxygenation monitor using NIR spatially resolved spectroscopy. In Proceedings of the SPIE Conference on Optical Tomography and Spectroscopy of Tissue III, San Jose, CA, USA, 15 July 1999.
21. Roldan, M.; Chatterjee, S.; Kyriacou, P.A. Light-Tissue Interaction Modelling of Human Brain towards the Optical Sensing of Traumatic Brain Injury. In Proceedings of the 43rd Annual International Conference of the IEEE Engineering in Medicine & Biology Society (EMBC), Jalisco, Mexico, 26 July 2021; pp. 1–4.
22. Mendelson, Y.; Ochs, B.D. Noninvasive Pulse Oximetry Utilizing Skin Reflectance Photoplethysmography. *IEEE Trans. Biomed. Eng.* **1988**, *35*, 798–805. [[CrossRef](#)] [[PubMed](#)]
23. Tucker, A.S.; Fox, R.M.; Sadle, R.J. Biocompatible, High Precision, Wideband, Improved Howland Current Source With Lead-Lag Compensation. *IEEE Trans. Biomed. Circuits Syst.* **2013**, *7*, 63–70. [[CrossRef](#)]
24. Roldan, M.; Kyriacou, P.A. Development of a head phantom for the acquisition of pulsatile optical measurements: Mimicking Blood and CSF circulation. *Photonics*, 2023; *in press*.
25. Elgendi, M. Optimal signal quality index for photoplethysmogram signals. *Bioengineering* **2016**, *3*, 21. [[CrossRef](#)] [[PubMed](#)]
26. Orphanidou, C. *Signal Quality Assessment in Physiological Monitoring*; SpringerBriefs in Bioengineering; Springer International Publishing: Cham, Switzerland, 2018; ISBN 978-3-319-68414-7.
27. Krishnan, R.; Natarajan, B.; Warren, S. Two-stage approach for detection and reduction of motion artifacts in photoplethysmographic data. *IEEE Trans. Biomed. Eng.* **2010**, *57*, 1867–1876. [[CrossRef](#)]
28. Elgendi, M.; Norton, I.; Brearley, M.; Abbott, D.; Schuurmans, D. Systolic Peak Detection in Acceleration Photoplethysmograms Measured from Emergency Responders in Tropical Conditions. *PLoS ONE* **2013**, *8*, 76585. [[CrossRef](#)]
29. Kim, J.; Kim, J.; Ko, H. Low-power photoplethysmogram acquisition integrated circuit with robust light interference compensation. *Sensors* **2016**, *16*, 46. [[CrossRef](#)] [[PubMed](#)]
30. Park, J.; Seok, H.S.; Kim, S.-S.; Shin, H. Photoplethysmogram Analysis and Applications: An Integrative Review. *Front. Physiol.* **2022**, *12*, 2511. [[CrossRef](#)] [[PubMed](#)]

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