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
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## Article

# Open-Access fNIRS Dataset for Classification of Unilateral Finger- and Foot-Tapping

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**Abstract:** Numerous open-access electroencephalography (EEG) datasets have been released and widely employed by EEG researchers. However, not many functional near-infrared spectroscopy (fNIRS) datasets are publicly available. More fNIRS datasets need to be freely accessible in order to facilitate fNIRS studies. Toward this end, we introduce an open-access fNIRS dataset for three-class classification. The concentration changes of oxygenated and reduced hemoglobin were measured, while 30 volunteers repeated each of the three types of overt movements (i.e., left- and right-hand unilateral complex finger-tapping, foot-tapping) for 25 times. The ternary support vector machine (SVM) classification accuracy obtained using leave-one-out cross-validation was estimated at 70.4%  $\pm$  18.4% on average. A total of 21 out of 30 volunteers scored a superior binary SVM classification accuracy (left-hand vs. right-hand finger-tapping) of over 80.0%. We believe that the introduced fNIRS dataset can facilitate future fNIRS studies.

**Keywords:** brain–computer interfaces; functional near-infrared spectroscopy; open-access dataset; finger-tapping; foot-tapping; three-class

## 1. Introduction

Owing to its many advantages, including portability, convenience in use, and scalability, when compared to functional magnetic resonance imaging, hemodynamic responses can be captured more easily than before using functional near-infrared spectroscopy (fNIRS) [1,2]. fNIRS is less vulnerable to electric noises and less sensitive to motion artifacts than electroencephalography (EEG) [3]. Especially, fNIRS is strongly robust to ocular artifacts; thus, frontal and prefrontal hemodynamic responses are not contaminated by the electrooculogram. These facts have turned the attention of those in the field of brain–computer interfaces (BCIs) towards fNIRS-BCIs as an alternative of EEG-BCIs (EEG-BCIs). In the field of fNIRS-BCIs, a variety of mental tasks (e.g., motor imagery, mental calculation, 3D rotation, and word association) have been adopted to induce task-related hemodynamic responses [4–24]. However, the hemodynamic responses induced by the visual stimulus are usually much slower than the steady-state visually evoked potentials, and do not provide unique advantages. Therefore, these hemodynamic responses have not been used for fNIRS studies actively [25,26].

Since hemodynamic responses are inevitably contaminated by physiological noises such as respiration and cardiac pulse, it is essential to separate the task-relevant hemodynamic responses from irrelevant components. Toward this end, researchers have so far conducted interesting studies concerning low-pass and band-pass filters [27–31], adaptive filters [32,33], wavelet filters [34,35], short channel separation [36–38], and principal component analysis filters [39,40]. Different types of machine learning methods, such as support vector machine (SVM) [41], linear discriminant analysis [42–44], hidden Markov model [45,46], and neural networks, have been considered to

discriminate the task-related hemodynamic responses induced by different types of mental tasks. Furthermore, feature selection techniques such as the Fisher score [47], sequential forward selection [48], and genetic algorithm [49] have been applied to enhance the performance improvement of fNIRS-BCI systems. Although fNIRS-BCIs have mainly focused on the implementation of binary BCI systems, multi-class fNIRS-BCI systems have recently been proposed [50,51]. In addition to BCIs, fNIRS has been actively used in clinical and neuroscience areas [52–54].

Data acquisition through fNIRS is normally a complicated and laborious task due to the hemodynamic delay on the order of several seconds [55,56]. For this reason, those interested in fNIRS have utilized their time for less essential works rather than the enhancement of fNIRS systems and techniques [57]. These laborious tasks have been redundant in many studies, and such repetitive works should be lessened. Therefore, we present an open-access fNIRS dataset containing 75 trials of three-class overt movements. We provide the temporal hemodynamic responses and ternary classification accuracy of task-related hemodynamic responses as references. The reminder of this paper is organized as follows: Section 2 describes the experiment and fNIRS dataset in detail. In Section 3, we overview the fNIRS system configuration and experimental paradigm. Temporal hemodynamic responses and classification results are presented in Section 4. A brief conclusion is provided in Section 5.

## 2. fNIRS Dataset

### 2.1. Participants

A total of 30 volunteers (29 right-handed; 17 males;  $23.4 \pm 2.5$  years old (mean  $\pm$  standard deviation)) participated in this study. No volunteers reported a history of psychiatric or neurological disorders that could affect the experiment. All the volunteers were provided complete information about the experiment along with the required instructions. Written consent from the participants approved by the Korea University Institutional Review Board (KUIRB-2019-0254-01) were submitted prior to data recording. Experiments were conducted according to the criteria set by the declaration of Helsinki. All volunteers were monetarily reimbursed for their participation.

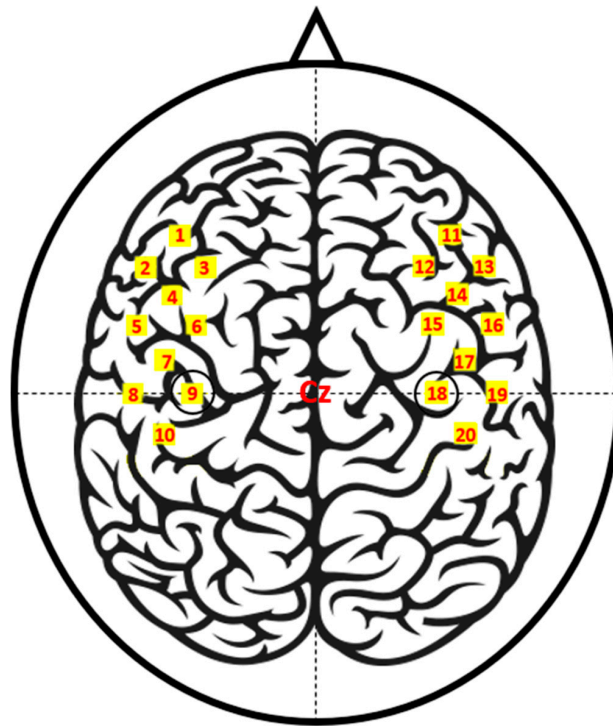
### 2.2. Apparatus

The fNIRS data were recorded by a three-wavelength continuous-time multi-channel fNIRS system (LIGHTNIRS, Shimadzu, Kyoto, Japan) consisting of eight light sources (Tx) and eight detectors (Rx). Four each of Tx and Rx were placed around C3 on the left hemisphere, and the rest were placed around C4 on the right hemisphere. Figure 1 depicts the placement of the fNIRS optodes.

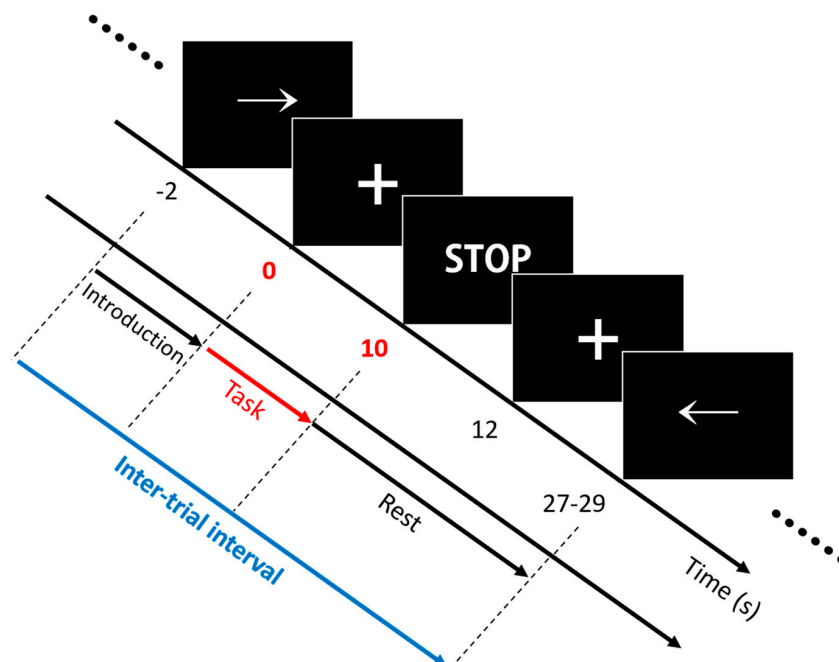
### 2.3. Experimental Paradigm

The experiment included three separate sessions comprising 25 trials with respect to each task. A single trial included an introduction period (2 s) and a task period (10 s), followed by an inter-trial break (17–19 s). Note that triggers were transmitted and marked in the data file at the start of the task periods. The inter-trial interval (i.e., the time interval between adjacent triggers) was 30 s on average. The volunteers were seated on a chair in front of a 27-inch LED monitor. The complete information and instructions regarding the experiment were displayed on the monitor. Out of right-hand finger-tapping (RHT), left-hand finger-tapping (LHT), and foot-tapping (FT), a specific task type was displayed at random, which volunteers were required to perform. The assigned task was initiated with a short beep, and the volunteers performed this task continuously during the task period. Then, an “inter-trial break period” was initiated with both a short beep and the display of a “STOP” sign on the monitor. Figure 2 presents the experimental paradigm. For RHT/LHT, the volunteers performed unilateral complex finger-tapping. They tapped their thumbs with other fingers one by one in the direction from the index finger to little finger and repeated it in the reverse order. The tapping continued at a steady rate of 2

Hz. For FT, the volunteers tapped their foot on the same side of their dominant hand constantly at a 1 Hz rate.



**Figure 1.** Functional near-infrared spectroscopy (fNIRS) channel locations. Ch01–10 and Ch11–20 are located around C3 (Ch09) and C4 (Ch18), respectively. The channels are created by a pair of adjacent light sources (Tx) and detectors (Rx) placed 30 mm away from each other.



**Figure 2.** Experimental paradigm. A single trial comprised an introduction period (−2 to 0 s) and a task period (0 to 10 s), followed by an inter-trial break period (10 to 27–29 s). Among right-hand finger-tapping (RHT) (→), left-hand finger-tapping (LHT) (←), and foot-tapping (FT) (↓), a random task type was displayed during the introduction period, which the volunteers were required to perform.

## 2.4. Dataset Description

Data with respect to a total of 30 volunteers were stored in MATLAB (Mathworks, Natick, USA) structure array format. Each volunteer's data comprised concentration changes of oxygenated/reduced hemoglobin  $\Delta\text{HbO}/\text{HbR}$  (cntHb), trigger (mrk), and fNIRS channel information (mnt). Each MATLAB structure array includes several fields, as listed in Table 1. The fNIRS dataset can be conveniently processed through the BCI toolbox ([https://github.com/bbci/bbci\\_public](https://github.com/bbci/bbci_public)) implemented by the Berlin Brain–Computer Interface group [58]. The fNIRS dataset is freely downloadable via [59] along with the hands-on tutorials from the GitHub repository [60].

**Table 1.** Functional near-infrared spectroscopy dataset description.

Structure	Field	Description
cntHb	.fs	Sampling rate (Hz)
	.clab	Channel labels
	.xUnit	X-axis unit
	.yUnit	Y-axis unit
	.snr	Signal-to-noise ratio
	.x	Concentration changes of oxygenated/reduced hemoglobin ( $\Delta\text{HbO}/\text{R}$ )
Mrk	.event.desc	Class labels' descriptions
	.time	Event occurrence times <sup>1</sup>
	.y	Class labels in vector form
mnt	.clab	Channel labels
	.box	Channel arrangement in Figures 3 and 4

<sup>1</sup> A trigger is marked where each task period starts.

## 3. Signal Processing

This section presents the signal processing methods employed to obtain the analysis results. All of the signal processing was conducted using MATLAB 2019a.

### 3.1. Preprocessing and Segmentation

The concentration changes with respect to the oxygenated and reduced hemoglobin ( $\Delta\text{HbO}/\text{R}$ ) were band-pass filtered through a zero-order filter implemented by the third-order Butterworth filter with a passband of 0.01–0.1 Hz to remove the physiological noises and DC offset. The  $\Delta\text{HbO}/\text{R}$  values were segmented into epochs ranging from −2 to 28 s relative to the task onset (i.e., 0 s). The epochs were subjected to a baseline correction to subtract the average value within the reference interval ranging from −1 to 0 s. The epochs included three-class (i.e., RHT, LFT, and FT)  $\Delta\text{HbO}/\text{R}$  data with respect to a total of 25 trials. We computed the signal-to-noise ratio (SNR) as follows:

$$\text{SNR} = 10 \log \frac{P_{\bar{s}}}{P_{\bar{n}}}, \quad (1)$$

where  $P_{\bar{s}}$  and  $P_{\bar{n}}$  are the powers of filtered data (signal estimate) and “unfiltered data – filtered data” (noise estimate), respectively. The SNRs of each of the channels were stored in the dataset; however, we did not discard the channels with low SNR values to ensure that the low-quality channel rejection according to arbitrary SNR threshold did not affect the classification results. Low-quality channel rejection can be a good practice in some cases to enhance the classification accuracy.

### 3.2. Classification

To extract the  $\Delta\text{HbO}/\text{R}$  features, three time windows in the ranges of 0–5 s, 5–10 s, and 10–15 s within epochs were employed to compute the average  $\Delta\text{HbO}/\text{R}$  for each of the 20 channels (the left and right hemispheres with 10 channels each). No feature/channel selection method was applied; therefore,

the feature vector comprised three features extracted from 20 channels, and the dimensionality of the feature vector was computed as 120 (3 time windows  $\times$  20 channels  $\times$  2 chromophores). A linear SVM was used to calculate the binary and ternary classification accuracies. Leave-one-out cross-validation (LOOCV) was applied to validate the fNIRS dataset. LOOCV allocates a single trial data to a test set and the rest of the trial data to a training set in order to train and validate the classifier. This method increases the size of the training set to alleviate problems which could result from the high-dimensional feature vector, and ensure the reproducibility of the cross-validation results. Feature vectors were standardized using the mean and standard deviation of the features included in the training set in order to render the performance of the linear SVM consistent.

#### 4. Results

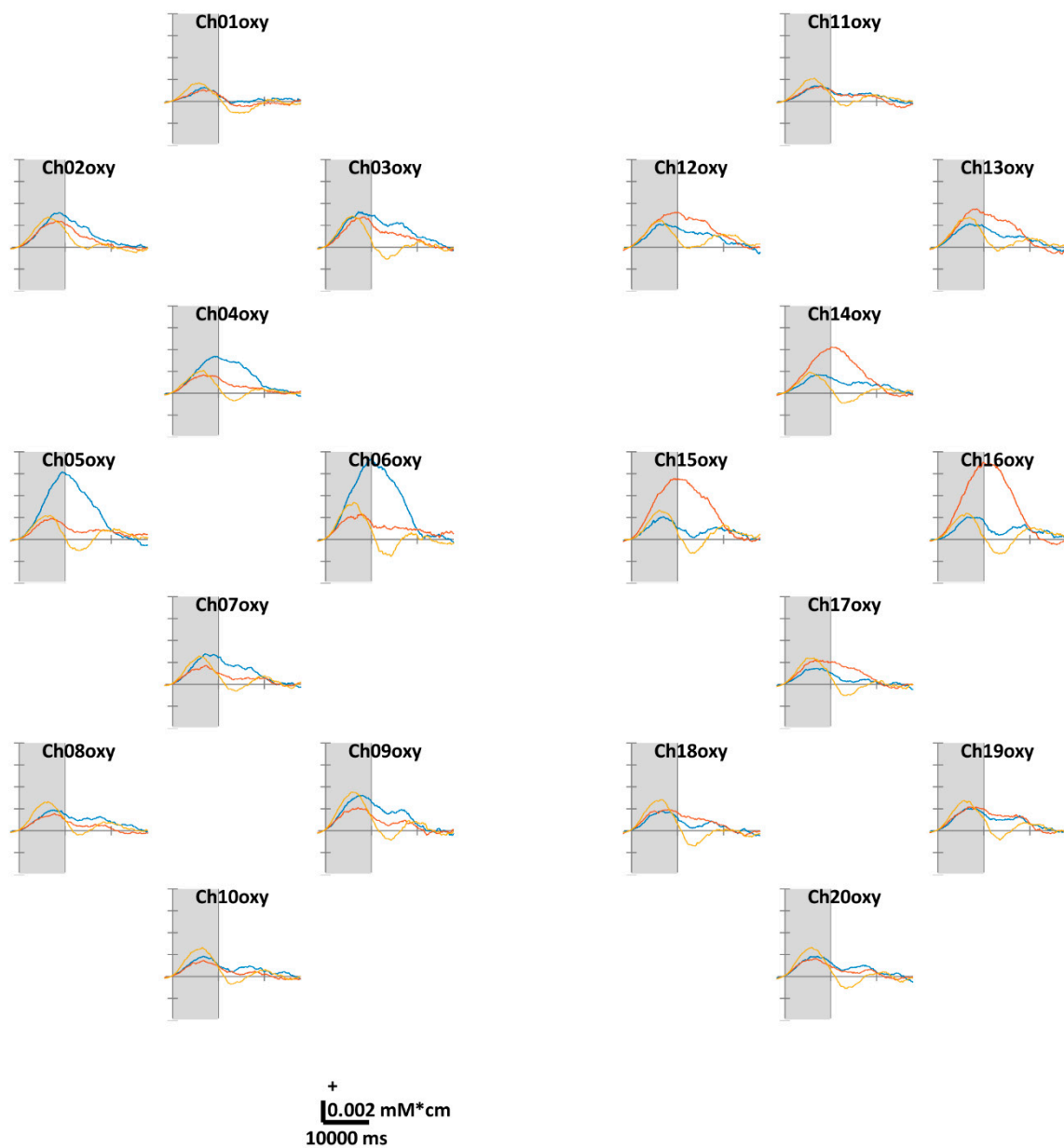
In this section, we present the primary analysis results, such as the grand averages of temporal  $\Delta\text{HbO/R}$  across all volunteers as well as the average/individual binary and ternary classification accuracies.

##### 4.1. Temporal $\Delta\text{HbO}$ and $\Delta\text{HbR}$

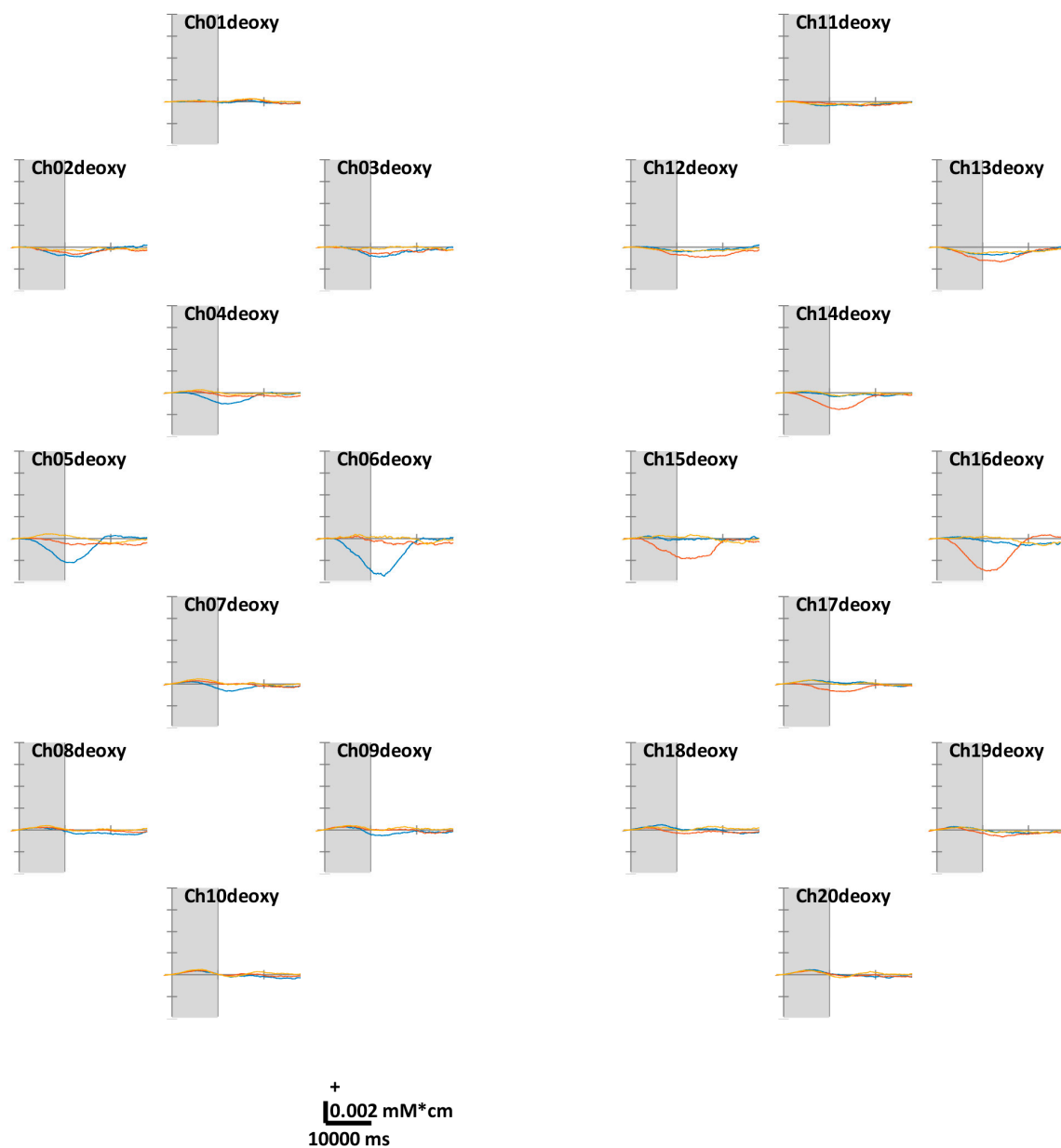
Figure 3 illustrates the task-related  $\Delta\text{HbO}$ . According to the theoretical fact empirically established by numerous studies, the motor cortex regions in contralateral hemispheres were well-activated while the volunteers were performing unilateral finger-tapping. Distinct  $\Delta\text{HbR}$  values were observed at Ch05, 06, 15, and 16 located in the anterior areas of C3 and C4. Unlike unilateral finger-tapping, while performing FT, the HbO responses were completely activated before (at 5–7 s) the end of the task period (at 10 s). At the end of the task period, the HbO level returns close to the baseline. We observed similar trends across all channels. For the  $\Delta\text{HbR}$  values presented in Figure 4, as in the case of HbO, distinct  $\Delta\text{HbR}$  in the direction opposite to that of  $\Delta\text{HbO}$  were observed in (pre)motor cortices in contralateral hemispheres. Distinct  $\Delta\text{HbR}$  was observed in the direction opposite to that of  $\Delta\text{HbO}$  corresponding to the same location, particularly at Ch05, 06, 15, and 16.

##### 4.2. Classification Accuracy

The LOOCV results are presented in Figure 5. The grand averages of binary classification accuracies were estimated at  $83.4\% \pm 17.0\%$ ,  $77.4\% \pm 14.6\%$ , and  $80.6\% \pm 14.3\%$  for RFT vs. LFT, RFT vs. FT, and LFT vs. FT, respectively. A significant difference in the classification accuracy with respect to the three binary classification cases was observed (Friedman test,  $p < 0.01$ ). Using post-hoc analysis, we revealed that RFT vs. LFT classification accuracy was significantly higher than RFT vs. FT classification accuracy (Wilcoxon signed-rank test with Bonferroni correction; corrected  $p < 0.01$ ). Differences in the classification accuracies of the rest of the binary classification cases were not statistically significant. The grand average of the ternary classification accuracy was estimated at  $70.4\% \pm 18.4\%$ . Note that the results with respect to 27 out of 30 volunteers exceeded the theoretical chance level of ternary classification of 42.7% ( $p < 0.05$ ) [61].

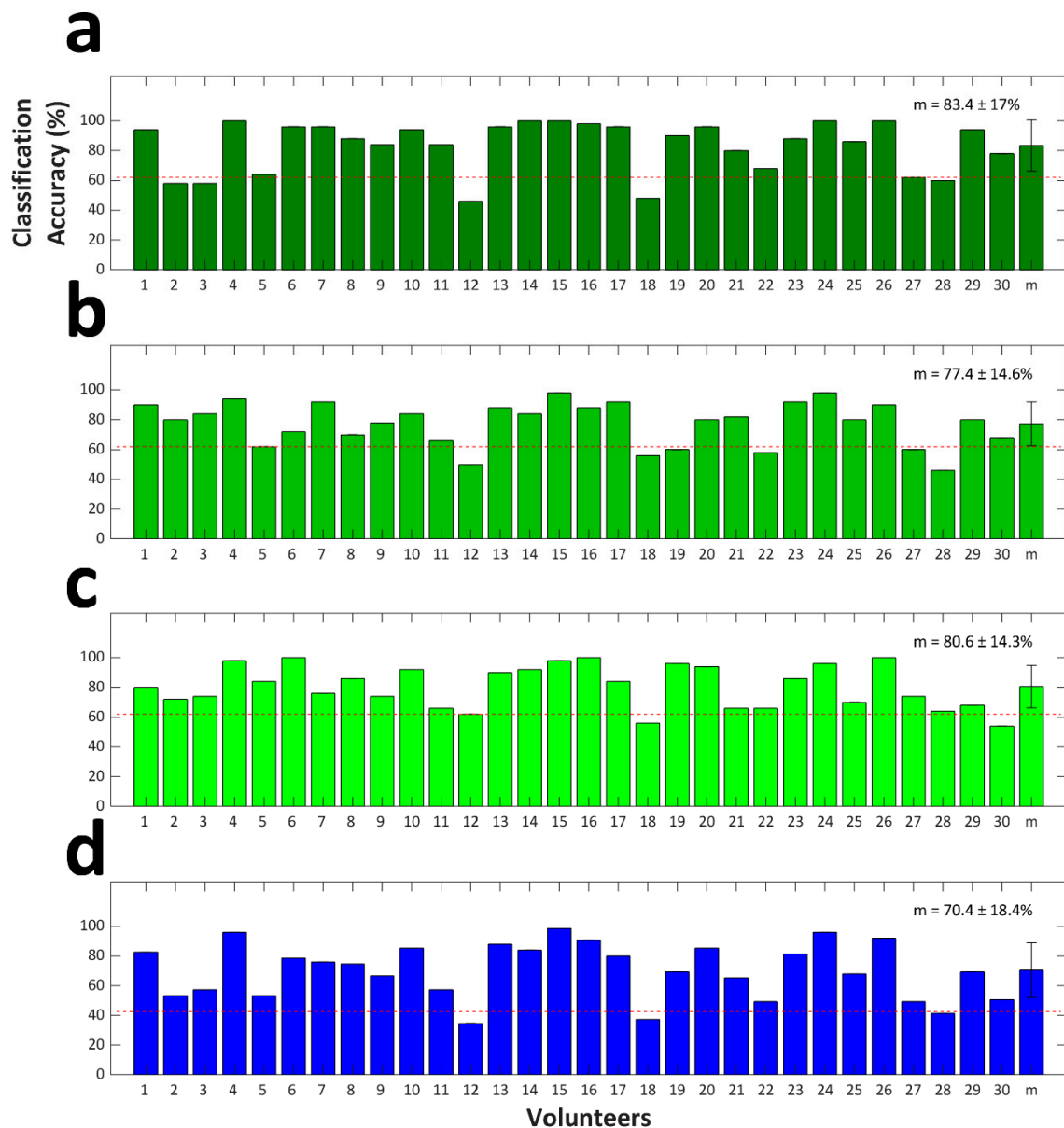


**Figure 3.** Grand average (across all volunteers) of temporal concentration changes of oxygenated hemoglobin ( $\Delta\text{HbO}$ ) within epochs' interval. The gray shade indicates the task period (0–10 s). The X- and Y-axes units are ms and  $\text{mM}\cdot\text{cm}$ , respectively. The blue, red, and yellow solid lines correspond to RHT, LHT, and FT, respectively.



**Figure 4.** Grand average (across all volunteers) of temporal concentration changes of reduced hemoglobin ( $\Delta\text{HbR}$ ) within epochs' interval. The gray shade indicates the task period (0–10 s). The X- and Y-axes units are ms and  $\text{mM}\cdot\text{cm}$ , respectively. The blue, red, and yellow solid lines correspond to RHT, LHT, and FT, respectively.





**Figure 5.** (a) RFT vs. LFT, (b) RFT vs. FT, and (c) LFT vs. FT binary classification accuracies. (d) Ternary classification accuracies. The error bars indicate the standard deviation. The X-label denoted by “m” indicates the grand average classification accuracy across the 30 volunteers. The grand average classification accuracy is provided in each figure. Red dashed lines indicate the theoretical chance levels (62.0% and 42.7% for (a–c) and (d), respectively).

## 5. Discussion and Conclusions

As shown in Figures 3 and 4, the  $\Delta\text{HbO/R}$  values with respect to the three tasks demonstrated consistent differences in specific channels and time intervals. Therefore, it was observed that the temporal averages of  $\Delta\text{HbO/R}$  within specific time intervals could be considered as relevant features to discriminate the task-related fNIRS data. As the feature extraction procedure employed for the classification was not optimized, more relevant feature extraction and selection could improve the binary and ternary classification accuracies. We provide brief tutorials along with the fNIRS dataset in order to enable an efficient handling of the dataset. We anticipate more interesting fNIRS studies and consider that the provided fNIRS dataset could contribute to the advancement of fNIRS technologies in the future.

**Author Contributions:** J.S. conceptualized the study and J.J. supervised the study. S.B. and J.P. implemented software for experiments and conducted experiments. S.B. and J.S. analyzed the data and wrote the manuscript. All authors reviewed the final manuscript.

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