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**Characterizing Fluctuations of Arterial and Cerebral Tissue Oxygenation in Preterm Neonates by Means of Data Analysis Techniques for Nonlinear Dynamical Systems**

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**Abstract:** The cerebral autoregulatory state as well as fluctuations in arterial (SpO₂) and cerebral tissue oxygen saturation (StO₂) are potentially new relevant clinical parameters in preterm neonates. The aim of the present study was to test the investigative capabilities of data analysis techniques for nonlinear dynamical systems, looking at fluctuations and their interdependence. StO₂, SpO₂ and the heart rate (HR) were measured on four preterm neonates for several hours. The fractional tissue oxygenation extraction (FTOE) was calculated. To characterize the fluctuations in StO₂, SpO₂, FTOE and HR, two methods were employed: (1) phase-space modeling and application of the recurrence quantification analysis (RQA), and (2) maximum entropy spectral analysis (MESA). The correlation between StO₂ and SpO₂ as well as FTOE and HR was quantified by (1) nonparametric nonlinear regression based on the alternating conditional expectation (ACE) algorithm, and (2) the maximal information-based nonparametric exploration (MINE) technique. We found that (1) each neonate showed individual characteristics, (2) a ~60 min oscillation was observed in all of the signals, (3) the nonlinear correlation strength between StO₂ and SpO₂ as well as FTOE and HR was specific for each neonate and showed a high value for a neonate with a reduced health status, possibly indicating an impaired cerebral autoregulation. In conclusion, our data analysis framework enabled novel insights into the characteristics of hemodynamic and oxygenation changes in preterm infants. To the best of our knowledge, this is the first application of RQA, MESA, ACE and MINE to human StO₂ data measured with near-infrared spectroscopy (NIRS).

**Keywords:** Long term measurements • Autoregulation • Near infrared spectroscopy • Correlation analysis • Spontaneous fluctuations

1. **Introduction**

Preterm infants exhibit an immature regulation of respiration as well as systemic and cerebral blood circulation (i.e. cerebral autoregulation, CO₂ vasoreactivity), leading to an increased incidence of hypoxic and hyperoxic episodes due to (1) large fluctuations in cerebral hemodynamics, and (2) impaired coupling between cerebral blood flow (CBF) and metabolic demand [1]. Episodes of intermittent hypoxemia occur in 74% of preterm infants, compared to 62% of term infants [2]. Hypoxemia or hyperoxemia may lead to an increase in mortality and neurological morbidity with long-term effects in later adult life. Greater variability in arterial oxygen saturation (SpO₂) [3] correlates with an increased incidence of retinopathy of prematurity (ROP). Thus, the assessment of the dynamics of SpO₂ and cerebral tissue oxygen saturation (StO₂) in preterm neonates may be of high clinical relevance. Due to continuous advancement in biomedical optics [4, 5], a reliable noninvasive long-term measurement of StO₂ in preterm neonates is in principle feasible [6, 7].

The aim of the present study was to analyze long-term measurements of StO₂ (conducted by multi-distance near-infrared spectroscopy, MD-NIRS) and SpO₂ heart rate (HR) and the fractional tissue extraction (FTOE) in preterm infants by means of data analysis techniques for nonlinear dynamical systems in order to investigate the characteristics of cerebral and systemic hemodynamic fluctuations and their interdependence.

2. **Material and Methods**

2.1 **Subjects, Instrumentation and Experimental Protocol**

A total of 20 clinically stable preterm neonates were enrolled. The study was approved by the ethics committee, and written informed consent was obtained from the parents before the study. Four neonates were selected for the present analysis, namely those with long continuous signals and the highest signal-to-noise ratio (SNR) (Table 64.1). SpO₂ and HR were determined by a standard patient monitor (Infinity Delta XL, Dräger, Germany) and StO₂ by an internally developed MD-NIRS device (OxyPrem, 4 × 3 [760, 805, 870 nm] light sources, two source-detector distances, i.e. 1.5, 2.5 cm [8]). OxyPrem uses the self-calibrating approach [9] which ensures a robust and high-precision measurement of absolute StO₂ values [10]. The NIRS optode was positioned over the left prefrontal cortex (PFC).

Measurements were performed continuously during the night (from ~10 pm till ~6 am), i.e. NIRS measured the resting-state activity of cerebral hemodynamics.
Table 64.1 Description of the study sample

2.2 Signal Processing and Data Analysis

From the SpO₂ and StO₂, we calculated the fractional tissue oxygenation extraction (FTOE = (SpO₂ – StO₂)/SpO₂ × 100 [%]). FTOE quantifies the balance between oxygen delivery and oxygen consumption and correlates significantly with the invasively measured oxygen extraction fraction [11]. All signals (SpO₂, StO₂, FTOE and HR) were downsampled to 0.05 Hz to increase the SNR and since only low frequencies were of interest. For each of the four datasets, an interval was chosen for the subsequent analysis which contains data without any signal distortion. The lengths of the data are given in Table 64.1. To characterize the fluctuations in StO₂, SpO₂, FTOE and HR, two different methods were applied:

- **Phase-space modeling and application of the recurrence quantification analysis (RQA)** [12, 13]. Each signal (StO₂, SpO₂, FTOE and HR) was embedded into a phase space with the dimension m and time delay τ. The optimal values for m and τ were determined by finding the first minimum of the false nearest neighbors function depending on m, and the autocorrelation function depending on τ, respectively. In a subsequent step, the phase space trajectories were characterized by the RQA. In particular, the determinism (DET, i.e. the predictability of the system), entropy of the diagonal length (ENT, i.e. the complexity of the system’s deterministic dynamics), and laminarity (LAM, i.e. the amount of intermittency of the system’s dynamics) were calculated.

- **Maximum entropy spectral analysis (MESA)** [14]. This method enables a high-precision spectral analysis based on the principle of maximum entropy. To prevent spurious peaks, the order of the MESA-based periodogram was set at one third of the number of samples [15].

The correlation between StO₂ and SpO₂ as well as FTOE and HR were quantified by two nonparametric methods:

- **Nonparametric nonlinear regression based on the alternating conditional expectation (ACE) algorithm** [16]. This technique finds the optimal transformations for the dependent and independent variables in order to maximize the correlation. The correlation strength is quantified by the maximal correlation coefficient, rACE.

- **Maximal information-based nonparametric exploration (MINE) technique** [17]. MINE enables the characterization of dependencies between variables. We calculated the maximal information coefficient (MIC) (relationship strength) and maximum asymmetry score (MAS) (departure from monotonicity).

In addition, each signal was characterized by calculating the median, and variability index 1 (V₁), quantified as the mean of the modulus of the first derivative. In addition, the relationship of the fluctuation strength of StO₂ vs. SpO₂ was determined by the ratio of their standard deviations (variability index 2, V₂).

### 3. Results, Discussion, Conclusion and Outlook

Figure 64.1a–d shows the time courses of SpO₂, StO₂, HR and FTOE. In Fig. 64.1b the normalized (i.e. subtraction of the mean value) SpO₂ and StO₂ were plotted to increase the visibility of similar dynamics. Figure 64.1d shows FTOE and HR after normalization (z-score) and smoothing (Kolmogorov-Zurbenko filter, window length: 180 s, iterations: 2) which increases the visibility of the similar long-term variability of both signals. The ACE correlation plots as well as the RQA, ACE and MINE results are visualized in Fig. 64.2. All signals show subject-specific dynamics:

- **RQA:** Noticeable low values for DET and LAM in neonate #4, high values for neonate #1. ENT (for StO₂ and SpO₂) has low values for neonate #3.

- **MESA:** (1) Neonate #1 exhibited a large oscillation with a period length (T) of 60 min in StO₂ and SpO₂ as well as HR and FTOE.

- **(2)** A large oscillation with T ≈ 60 min is present in neonate #2 for StO₂ and SpO₂, followed by a second strongest oscillation with T ≈ 30 min. In FTOE the predominant oscillation was at T ≈ 60 min. The spectra of StO₂ and SpO₂ have a remarkably similar fine structure of oscillatory peaks indicating a large degree of similarity in the dynamics. In addition, an oscillation with T ≈ 15 min can be seen in StO₂, SpO₂, HR and FTOE.

- **(3)** Neonate #3 shows an oscillation peak with T ≈ 60 min in StO₂, SpO₂, HR and FTOE, whereas for FTOE a larger oscillation with T = 30 min is present. The spectra of StO₂ and SpO₂ have different fine structures, which is also true for the spectra of HR and FTOE.

- **(4)** Neonate #4 shows a strong peak with T ≈ 60 and 30 min in StO₂ and SpO₂.

- **ACE and MINE:** Concerning the relationship between StO₂ and SpO₂, the largest rACE and MIC value was found for neonate #2, the lowest for neonate #3; MAS was highest in neonate #3. Concerning HR vs. FTOE, neonate #2 had the largest rACE, MIC and MAS values. Neonate #4 showed significantly low MAS values for all four conditions (StO₂ vs. SpO₂ or HR, HR vs. FTOE or SpO₂).

To interpret the results it is helpful to discuss the similarities and differences of the signal characteristics with respects to the four neonates:

- **Similarities:** (1) The values for DET, ENT and LAM were all higher for StO₂ compared to SpO₂ (except for ENT of neonate #4), indicating more complex signal characteristics of StO₂ than of SpO₂.

- **Differences:** (1) DET, LAM and ENT of SpO₂ were highest for neonate #3.

- **(2) The correlation (rACE, MIC) between StO₂ and SpO₂ as well as HR and FTOE was highest for neonate #2; neonate #3 showed the lowest StO₂/SpO₂ correlation.**

- **(3) The smallest MAS values for all four correlations**
Fig. 64.1 (a–d) Visualization of the analyzed signals (StO$_2$, SpO$_2$, FTOE and HR). (e–l) Frequency spectra obtained by MESA.
Fig. 64.2 (a, b) Correlation diagrams based on ACE nonparametric nonlinear regression. (c–j) Parameters obtained by RQA, MESA, ACE, MINE as well as the values for the median and variability
were found for neonate #4 (except for StO2 vs. HR, neonate #2).
(3) The smallest MAS values for all four correlations were found for neonate #4 (except for StO2 vs. HR, neonate #2).
(4) In neonate #3 we observed the highest median values for StO2 and SpO2 as well as the lowest ones for HR and FTOE. In neonate #4 the lowest StO2 and the highest FTOE value were measured.
(5) High Vf values in StO2 and SpO2 were present in neonates #2 and #4. Neonate #3 showed the largest Vf value for StO2/SpO2.
(6) The mean StO2 values correlated inversely with the individual Hct and GA (at birth) ($r = -0.927$ and $r = -0.982$, respectively; $p < 0.05$).

The physiological interpretation of these findings is not straightforward since all patient-specific characteristics have an influence on the analyzed parameters.

In particular, the general health state (e.g. PDA, microbleeds, ischemia: yes/no), the type of respiration (ventilatory support: yes/no, type of support), and the GA (at birth/measurement) could potentially have a strong impact on the parameters.

The following observations were made based on our analysis:

(1) The general inverse correlation observed between StO2 and Hct was also noticed by other studies (e.g. [18]).
(2) Neonate #3 exhibited a large Vf, e.g. the fluctuations in StO2 were much stronger than in SpO2 (especially the decreases), a pattern that is observed by neonates with a PDA – indeed, neonate #3 had a PDA (which was however classified as not hemodynamically relevant).

The oscillations in the data with $T \approx 60$ and 30 min could originate from sleep phases. A sleep-wake cycling (with a quiet sleep phase with $T \approx 20$ min) is known [20] in term newborns with $T \approx 50–60$ min and an increase in total hemoglobin and HR during active sleep (compared to quite sleep) has previously been observed [21, 22]. (4) The two neonates with the lowest GA at birth (#2, #4) had the largest variability of StO2, SpO2 and FTOE which could indicate an immature functioning of cerebral hemodynamic regulation.

In conclusion, using four case studies, we demonstrated the possibility of realizing long-term measurements in preterm neonates with MD-NIRS and we presented a novel framework for investigating the characteristics of cerebral and systemic hemodynamic fluctuations and their interdependence. A follow-up study, investigating the signal characteristics in healthy and ill preterm neonates using the same framework would be the next step. Focusing on the fluctuation characteristics of the signals may offer novel insights into systemic and cerebral hemodynamics that are not assessed routinely only using traditional analyses (i.e. based on moments and linear correlations). To the best of our knowledge, this is the first application of RQA, MESA, ACE and MINE to human NIRS data.

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