

RISK APPRAISAL BY NOVEL CHAOTIC GLOBALS TO HRV IN SUBJECTS WITH MALNUTRITION

Gláucia Siqueira Barreto¹, Franciele Marques Vanderlei², Luiz Carlos M. Vanderlei², David M. Garner³

DOI: dx.doi.org/10.7322/jhdg.88900

ABSTRACT

The aim of this study is to assess the risk of dynamical diseases in malnourished children. This is achieved by the application of novel chaotic global techniques to the RR-intervals of the electrocardiogram (ECG) in the cohort. Heart Rate Variability (HRV) is an inexpensive and non-invasive tool to measure the autonomic impulses. Here there has been a decrease in chaotic response of HRV. Seventy children were divided into equal groups and the HRV monitored for 20-25 minutes. The Chaos Forward Parameter (CFP) which applies all three chaotic global parameters is suggested to be the most robust algorithm. These three parameters are high spectral entropy (hsEntropy), high spectral detrended fluctuation analysis (hsDFA) and spectral multi-taper method (sMTM). hsEntropy is a function of the irregularity of amplitude and frequency of the power spectrum peaks. It is derived by applying Shannon entropy to the multi-taper method power spectrum. To derive hsDFA we calculate the spectral adaptation in exactly the same way as for hsEntropy using an adaptive multi-taper method power spectrum with the same settings; but DFA rather than Shannon entropy is the algorithm applied. sMTM is the area between the multi-taper method power spectrum and the baseline. After Anderson-Darling and Lilliefors tests of normality; Kruskal-Wallis was used for the statistical analysis, with the level of significance set at ($p < 0.01$). Principal Component Analysis (PCA) identified two components representing 100% of total variance. Autonomic imbalance measured as HRV and an increased cardiovascular risk are described for overweight children as well as for malnourished and those with anorexia nervosa. The relationship between malnourishment and complexity measures is useful in the risk assessment of dynamical diseases associated with the condition. This is supportive in treatments, assisting the determination of the level of dietary or pharmacological intervention especially in related dynamical diseases.

Keywords: malnutrition, non-linear dynamics, heart rate variability, multi-taper method.

INTRODUCTION

Heart rate variability (HRV) can oscillate in a chaotic and complex way¹⁻⁴. In the past, methods derived from statistical physics and thermodynamics have allowed researchers to study such systems⁵. HRV is a cheap and non-invasive method of monitoring the sympathetic and parasympathetic balance. Recently, study of HRV in this way has become important; and often termed 'dynamical disease study'⁶. High HRV is a signal of good adaptation and characterize a healthy person with efficient autonomic mechanisms. Whilst lower HRV is frequently an indicator of abnormal and insufficient adaptation of the autonomic nervous system (ANS); causing the subject low physiological function. This decrease is consistent with a dysfunctional vagus.

Typically spectral entropy⁷ and techniques termed spectral detrended fluctuation analysis (sDFA) and spectral multi-taper method (sMTM) are based on 'chaotic globals'⁸. Spectral entropy applies the standard Shannon entropy^{9,10} algorithm to a Welch power spectrum¹¹. Whereas, sDFA applies the DFA algorithm in the same manner to the same power spectrum. This attempts to overcome the disadvantage of sparse data vulnerability – only phase information is lost. sMTM applies the responsive and adaptive multi-taper method (MTM)^{12,13} to the data. sMTM is the value of the

area between the MTM spectrum and the baseline. There has been speculation^{14,15} that if spectral entropy and sDFA previously applied to Welch power spectra were applied to multi-taper spectra which are adaptive and more sensitive; results may have greater chaotic parametric response.

These computations are useful clinically in patients under anaesthesia^{16,17}; or unable to communicate distress as in sleep apnea¹⁸ or dyspnea¹⁹⁻²¹ and the potential risk associated with diabetes mellitus¹⁴ and obesity¹⁵ has been studied previously using these methods. This study aimed to assess the risk of dynamical disease by novel chaotic globals to HRV in subjects with malnutrition. Autonomic imbalance measured as HRV and an increased cardiovascular risk are described for overweight children as well as for malnourished and those with anorexia nervosa²². Whether those with mild to moderate malnutrition compromise the ANS and therefore HRV is less certain²³.

METHODS

Population and Sample

A total of 70 volunteers of both sexes between three and five years of age were divided into two equal groups: malnourished (23 girls; 3.71 ± 0.75 years; 13.02 ± 1.71 Kg; 91.53 ± 5.47 cm; Z-score = -2.80 ± 0.59) or eutrophic (20 girls; 4.09 ± 0.85

¹ Faculdade de Tecnologia Intensiva. FATECI – Fortaleza, Ceará, Brazil.

² Department of Physiotherapy, UNESP - Univ Estadual Paulista - Presidente Prudente, São Paulo, Brazil.

³ Cardiorespiratory Research Group, Department of Biological and Medical Sciences, Faculty of Health and Life Sciences, Oxford Brookes University, Gypsy Lane, Oxford OX3 0BP, United Kingdom.

Corresponding author: dgarner@brookes.ac.uk

Suggested citation: Barreto GS, et al. Risk appraisal by novel chaotic globals to HRV in subjects with malnutrition. 24(3): 243-248

Manuscript submitted Dec 10 2013, accepted for publication Oct 10 2014.

years; $17.89 \pm 3.04\text{Kg}$; $106.83 \pm 8.15\text{ cm}$; Z-score = 0.191 ± 1.28). The malnourished group consisted of children less than -2 in Z score in relative to the height for the age, according with the criteria for age and sex by the World Health Organization (WHO)²⁴. The eutrophic group consisted of children with Z scores greater than or equal to -2 and less than +3, also according to WHO criteria. Excluded from the study were obese children (Z-score greater than +3) or who had at least one of the following features: children who were taking medications that would influence autonomic activity of the heart, such as propranolol and atropine. In addition to children who presented infections, metabolic diseases or known cardiorespiratory system diseases, which could affect the cardiac autonomic control.

The volunteers and parents/guardians were duly informed as to the procedures and objectives of the study and, after agreeing to participate, the parents/guardians signed terms of informed consent. All procedures received approval from the ethics committee of the Institution (Process n° 275.310).

Experimental Protocol

Prior to beginning the experimental procedure, information was recorded on age, gender, weight and height. The anthropometric measurements were undertaken following the recommendations of Lohman *et al.*²⁵ Weight was determined using a digital scale (Filizzola PL 150, Filizzola Ltda., Brazil) with a precision of 0.1 kg, with the children barefoot and wearing light-weight clothing. Height was determined using a stadiometer with a precision of 0.1 cm. The data collection was performed in a room with the temperature between 21° C and 23° C and relative humidity between 40 and 60%. Data were collected between 14:00 and 17:00 to minimize the interference of circadian rhythm. After the initial evaluation, all procedures necessary for the data collection were explained on an individual basis and the children were instructed to remain at rest and avoid talking during the collection.

The heart monitor belt was then placed over the thorax, aligned with the distal third of the sternum and the Polar S810i heart rate receiver (Polar Electro, Finland) was placed on the wrist. The equipment was previously validated for monitoring beat-by-beat heart rate and the use of

these data for HRV analysis in children and adults²⁶. The children were placed in the dorsal decubitus position on a cushion and remained at rest with spontaneous breathing for 20 minutes. After the collection, the child was discharged. The HRV behavior pattern was recorded beat-by-beat throughout the monitoring process at a sampling rate of 1000 Hz. Following digital filtering complemented with manual filtering for the elimination of premature ectopic beats and artifacts, 1000 consecutive R-R intervals were used for the data analysis. Only series with more than 95% sinus rhythm were included in the study²⁷.

CHAOTIC GLOBAL PARAMETERS

Multi-Taper Method: Power Spectrum

As mentioned in the introduction there is criticism in previous studies on diabetes and obesity with respect to chaotic global parameters in that the spectral entropy and sDFA analysis may be more sensitive if we applied the Shannon entropy and DFA algorithms to the multi-taper spectrum rather than the Welch power spectrum. Thus the spectra applied in all three chaotic global parameters would be the same.

MTM provides a useful tool for spectral estimation and signal reconstruction, of a time series of a spectrum that may contain broadband and line components. MTM is non-parametric since it does not apply an *a priori*, parameter dependent model of the process that generated the time series under analysis. In this sense it is similar to the Maximum Entropy Measure^{28,29}. MTM reduces the variances of spectral estimates by using a small set of tapers. Data is pre-multiplied by orthogonal tapers created to minimize the spectral leakage owing to the finite length of the time series. A set of independent approximations of the power spectrum is calculated. Functions known as Discrete Prolate Spheroidal Sequences (DPSS) are a set of functions which optimize the tapers. They are defined as eigenvectors of a Rayleigh-Ritz minimization problem.

MTM has the following features (1) Efficient in detecting periodic components; (2) A random signal may generate many false peaks which may or may not be significant; (3) There are two ways of testing the spectrum (red-noise³⁰ and harmonic tests).

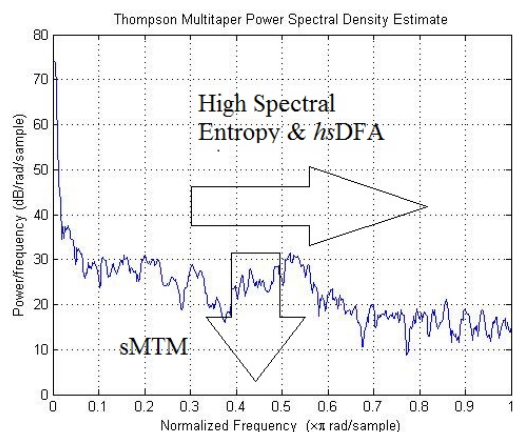


Figure 1: A Multi-Taper Method power spectrum of 1000 ECG RR intervals of a Malnourished subject. Illustrated by the arrows are high spectral entropy, high spectral detrended fluctuation analysis and spectral multi-taper method.

Chaotic Globals

High spectral entropy (*hs* Entropy) is a function of the irregularity of amplitude and frequency of the power spectrums peaks. It is derived by applying Shannon entropy^{9,10} to the MTM power spectrum. The parameters for MTM are: (1) sampling frequency of 1Hz; (2) time bandwidth for the DPSS is 3; (3) FFT length of 256; (4) Thomson’s adaptive nonlinear combination method to combine individual spectral estimates.

This output is then normalized so that the sum of the magnitude is equal to unity; giving a normalized power spectrum. We then calculate an intermediate parameter which is the median Shannon entropy of the value obtained from three different power spectra using the MTM power spectra under three test conditions: a perfect sine wave, uniformly distributed random variables, and finally the experimental oscillating signal.

These values are then again normalized mathematically so that the sine wave gives a value of zero, uniformly random variables give unity, and the experimental signal between zero and unity. It is this final value that corresponds to high spectral entropy.

The standard DFA algorithm^{31,32} can be applied to datasets where statistics such as mean, variance and autocorrelation vary with time. The difference with the high spectral detrended fluctuation analysis (*hsDFA*) algorithm is that the DFA is applied to the frequency rather than time on the horizontal axis. So, once more, the x-axis is frequency and the y-axis is amplitude. To obtain *hsDFA* we calculate the spectral adaptation in exactly the same way as for high spectral entropy using a MTM power spectrum with the same settings; but DFA rather than Shannon entropy is the algorithm applied. Spectral Multi-Taper Method (*sMTM*) is founded on the increased intensity of broadband noise in power spectra generated by irregular and chaotic signals. *sMTM* is the area between the MTM power spectrum and the baseline.

Chaotic Forward Parameter

The parameter [CFPx 1-7] is referred to as Chaotic Forward Parameter for the functions 1 to 7 below where it is applied to normal and malnourished subject datasets. Since *hsDFA* responds to chaos in the opposite way to the others we subtract its value from unity when applying here. All three chaotic global values have equal weightings.

$$\begin{aligned}
 1 [CFPx] &= \left[\left(\left[\frac{hsEntropy}{\max(hsEntropy)} \right] \right)^2 + \left(\left[\frac{sMTM}{\max(sMTM)} \right] \right)^2 + \left(1 - \left[\frac{hsDFA}{\max(hsDFA)} \right] \right)^2 \right]^{\frac{1}{2}} \\
 2 [CFPx] &= \left[\left(\left[\frac{hsEntropy}{\max(hsEntropy)} \right] \right)^2 + \left(1 - \left[\frac{hsDFA}{\max(hsDFA)} \right] \right)^2 \right]^{\frac{1}{2}} \\
 3 [CFPx] &= \left[\left(\left[\frac{hsEntropy}{\max(hsEntropy)} \right] \right)^2 + \left(\left[\frac{sMTM}{\max(sMTM)} \right] \right)^2 \right]^{\frac{1}{2}} \\
 4 [CFPx] &= \left[\left(\left[\frac{sMTM}{\max(sMTM)} \right] \right)^2 + \left(1 - \left[\frac{hsDFA}{\max(hsDFA)} \right] \right)^2 \right]^{\frac{1}{2}} \\
 5 [CFPx] &= \left[\left(1 - \left[\frac{hsDFA}{\max(hsDFA)} \right] \right)^2 \right]^{\frac{1}{2}} \\
 6 [CFPx] &= \left[\left(\left[\frac{sMTM}{\max(sMTM)} \right] \right)^2 \right]^{\frac{1}{2}} \\
 7 [CFPx] &= \left[\left(\left[\frac{hsEntropy}{\max(hsEntropy)} \right] \right)^2 \right]^{\frac{1}{2}}
 \end{aligned}$$

RESULTS

Mean Variation & Significances

Parametric statistics generally assume the data are normally distributed, hence the use of the mean as a measure of central tendency. If we cannot normalize the data we should not compare means. To test our assumptions of normality we apply the Anderson-Darling³³ test and the Lilliefors test³⁴. The Anderson-Darling test for normality applies an empirical cumulative distribution function. The Lilliefors test is useful in studies with small sample sizes. In the majority of cases the $p < 0.05$; for both tests so we cannot pronounce that the observations follow a normal distribution.

Therefore we have a probability plot of mainly non-normal data and so we must apply the Kruskal-Wallis³⁵ test of significance (non-parametric) as the ANOVA¹³⁶ (parametric) is unreliable for strong departures from normality. The results illustrate that there is a wide variation in both the mean values for both normal and malnourished (See Table 1). The test of significance applied is the Kruskal-Wallis test. The algorithm computes a significant statistical result for two of the seven combinations ($p < 0.01$). These are combinations [CFPx 1 & 3]. In both cases there is a decrease in chaotic response when going from normal to malnourished subjects. Standard deviation is also reduced.

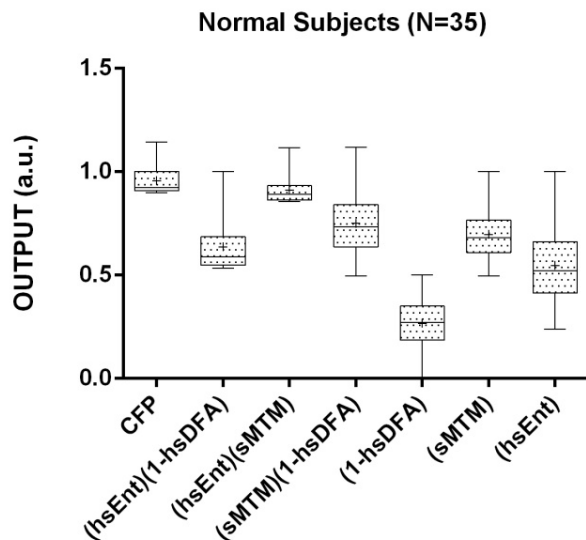


Figure 2: The boxplots illustrate the mean values and standard deviation of CFP for the RR intervals of normal subjects. Mean values are indicated by the (+) symbol. The number of RR intervals is 1000 and number of subjects is 35.

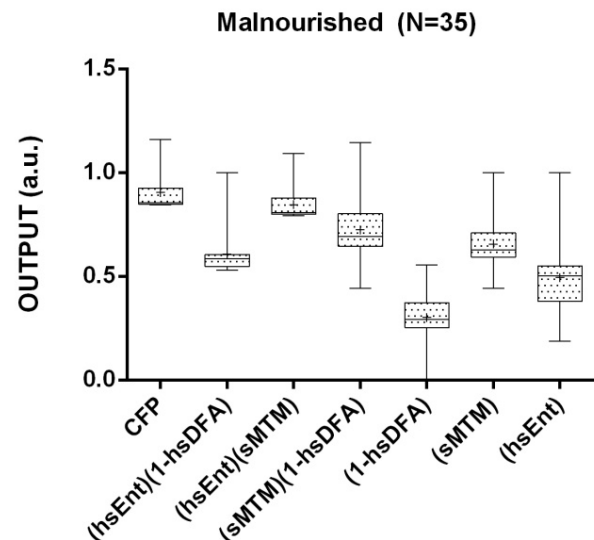


Figure 3: The boxplots illustrate the mean values and standard deviation of CFP for the RR intervals of malnourished subjects. The number of RR intervals is 1000 and number of subjects is 35. Mean values are indicated by the (+) symbol.

Table 1: The table below shows the mean values and standard deviation of the Chaos Forward Parameters [1 to 7] for the normal and malnourished subjects RR intervals. Kruskal-Wallis tests of significance was applied to results.

Combination of Chaotic Globals [CFPx 1 to 7]	Mean Normal (n = 35)	Standard Deviation Normal	Mean Malnourished (n = 35)	Standard Deviation Malnourished	Kruskal-Wallis P-value
Chaos Forward Parameter	0.9564	0.0678	0.9046	0.0858	<0.0001
(hsEntropy)(1-hsDFA)	0.6367	0.1142	0.6085	0.1044	0.3296
(hsEntropy)(sMTM)	0.9109	0.0618	0.8451	0.0758	<0.0001
(sMTM)(1-hsDFA)	0.7503	0.1567	0.7248	0.1607	0.5689
(1-hsDFA)	0.2667	0.1226	0.3015	0.1236	0.2219
(sMTM)	0.6972	0.1237	0.6554	0.1250	0.0885
(hsEntropy)	0.5473	0.1838	0.4942	0.1782	0.2131

Principal Component Analysis

Principal Component Analysis (PCA) ³⁷; a multivariate technique is applied here. We have the values of [CFPx] for two groups for 35 subjects who are malnourished; hence a grid of 2 by 35 to be assessed. The First Principal Component (PC1) has a variance (eigenvalue) of 1.9 and accounts for 95% of the total variance. The Second Principal Component (PC2) has an eigenvalue of 0.1 accounting for 100% of cumulative total variance. PC2 accounting for 5% of its proportion of the variance. Therefore we can assume that most variance is achieved in the first two components. Only [CFPx 1 & 3] are significantly different when tested by Kruskal-Wallis test (p < 0.01).

[CFPx1] has the First Principal Component (0.707) and the Second Principal Component (-0.707). [CFPx3] has the First Principal Component (0.707) and the Second Principal Component (0.707). Only the first two components need be considered due to the steep scree plot. Only [CFPx 1 & 3] need to be considered due to Kruskal-Wallis statistical significance at the level p < 0.01 (Table 1).

So, both are suitable functions as deduced by the three assessments (Kruskal-Wallis, Standard Deviation and PCA). There is evidence to apply [CFPx 1] as the most robust function, as in the study which analyzes the ‘inverse problem’ posed by Garner and Ling⁸. This in addition to forward problems in obesity¹⁵, diabetes mellitus¹⁴ and chronic obstructive pulmonary disease ^{38,39}.

DISCUSSION

The first algorithm which applies all three chaotic global parameters is suggested as the most robust algorithm. Referring to Garner and Ling ⁸; which uses three models — Duffing, Brusselator and Lorenz for the purposes of optimization; [CTF] a variant of [CFPx] is the most reliable objective function when tested by PCA. This is reinforced here by PCA applied to the two adaptations of [CFPx] for malnourished subjects. Here 100% of influence is achieved by the first two Principal Components; with the [CFPx] with all three chaotic globals applied testing as most influential algorithm.

Future development could involve DPSS of the MTM being adjusted to optimize the final level of significance. In addition the weighting of the three chaotic global parameters could be attuned since here they have only equal weightings of unity. It would also be statistically favourable to have larger, but equal datasets for both normal and malnourished subjects.

We have developed two robust functions which can take short time-series of HRV and discriminate between the control and experimental groups. There is a very high level of significance for these algorithms ($p < 0.0001$). By applying either of these novel functions to the shorter time-series via spectrally determined parameters it should be possible to determine which are malnourished or normal. This achieved more rapidly and efficiently with regards to time and data length. There has been a decrease in chaotic response of HRV in malnourishment. The relationship between malnourishment and complexity measures is useful in the risk assessment of dynamical diseases associated with

the condition. It identifies severity of the situation from a cheap and reliable method of monitoring the ANS. This is helpful in treatments, assisting the determination of the level of dietary or pharmacological intervention especially in related dynamical diseases. For example, autonomic neuropathy measured by HRV in Vitamin B₁₂ deficiency appears to precede other neurological tests⁴⁰. Nevertheless, proceeding cautiously, since the subject's autonomic modulation may be critical. Moreover, it is noted that there are other conditions which could cause the complexities correlation.

In conclusion, there has been a decrease in chaotic response of HRV in malnutrition children. The parameter which applies all three parameters is the most influential and statistically more significant. This method is useful in the risk assessment of dynamical diseases associated with malnourishment.

Competing interests

The authors declare that there is no conflict of interests regarding the publication of this article.

REFERENCES

- Seely AJ, Macklem P. Fractal variability: an emergent property of complex dissipative systems. *Chaos* 2012;22(1):013108. DOI: 10.1063/1.3675622.
- Goldberger AL, West BJ. Chaos and order in the human body. *MD Comput* 1992;9(1): 25-34.
- Goldberger AL. Cardiac chaos. *Science* 1989;243(4897):1419. DOI: 10.1126/science.2928773.
- Elbert T, Ray WJ, Kowalik ZJ, Skinner JE, Graf KE, Birbaumer N. Chaos and physiology: deterministic chaos in excitable cell assemblies. *Physiol Rev* 1994;74(1):1-47. DOI: MD5:764bd488f011be86fbf44bdeb3c29845.
- Ho M-W. The rainbow and the worm: The physics of organisms. Singapore: World scientific; 2008.
- Mackey MC, Milton JG. Dynamical diseases. *Ann NY Acad Sci* 1987;504(1):16-32. DOI: 10.1111/j.1749-6632.1987.tb48723.x
- Johnson R, Shore J. Which is the better entropy expression for speech processing: $-S \log S$ or $\log S$? *IEEE Trans Acoust, Speech, Signal Processing* 1984;32(1):129-37. DOI: 10.1109/TASSP.1984.1164296.
- Garner DM, Ling BWK. Measuring and locating zones of chaos and irregularity. *J Syst Sci Complex* 2014;27(3):494-506. DOI: 10.1007/s11424-014-2197-7.
- Shannon CE. A mathematical theory of communication. *ACM Mob Comput Commun Rev* 2001;5(1):3-55. DOI: 10.1145/584091.584093.
- Zyczkowski K. Renyi extrapolation of Shannon entropy. *Open Sys & Information Dyn* 2003;3(10):297-310. DOI: 10.1023/A:1025128024427.
- Alkan A, Kiyimik MK. Comparison of AR and Welch methods in epileptic seizure detection. *J Med Syst* 2006;30(6):413-9. DOI: 10.1007/s10916-005-9001-0.
- Ghil M. The SSA-MTM Toolkit: Applications to analysis and prediction of time series. *Appl Soft Comput* 1997;3165:216-30. DOI: 10.1117/12.279594.
- Percival DB, Walden AT. Spectral Analysis for Physical Applications: Multitaper and Conventional Univariate Techniques. New York: Cambridge University Press; 1993. DOI: dx.doi.org/10.1017/CBO9780511622762.
- Souza NM, Vanderlei LC, Garner DM. Risk evaluation of diabetes mellitus by relation of chaotic globals to HRV. *Complexity* 2014. DOI: 10.1002/cplx.21508.
- Vanderlei FM, Vanderlei LC, Garner DM. Chaotic global parameters correlation with heart rate variability in obese children. *J Hum Growth Dev* 2014;24(1):24-30.
- Kawaguchi M, Takamatsu I, Kazama T. Rocuronium dose-dependently suppresses the spectral entropy response to tracheal intubation during propofol anaesthesia. *Br J Anaesth* 2009;102(5):667-72. DOI: 10.1093/bja/aep040.
- Morgaz J, Granados MD, Dominguez JM, Navarrete R, Fernandez A, Galan A, et al. Evaluation of spectral entropy to measure anaesthetic depth and antinociception in sevoflurane-anaesthetised Beagle dogs. *Vet J* 2010; 188(3):352-5. DOI: 10.1016/j.tvjl.2010.06.001.
- Alvarez D, Hornero R, Marcos J, Del Campo F, Lopez M. Spectral analysis of electroencephalogram and oximetric signals in obstructive sleep apnea diagnosis. *Conf Proc IEEE Eng Med Biol Soc* 2009;2009:400-3. DOI: 10.1109/IEMBS.2009.5334905.
- Moosavi SH, Topulos GP, Hafer A, Lansing RW, Adams L, Brown R, et al. Acute partial paralysis alters perceptions of air hunger, work and effort at constant P(CO₂) and V(E). *Respir Physiol* 2000;122(1):45-60. DOI: 10.1016/S0034-5687(00)00135-3.
- Banzett RB, Lansing RW, Reid MB, Adams L, Brown R. 'Air hunger' arising from increased PCO₂ in mechanically ventilated quadriplegics. *Respir Physiol* 1989;76(1):53-67. DOI: 10.1016/0034-5687(89)90017-0.

21. Binks AP, Vovk A, Ferrigno M, Banzett RB. The air hunger response of four elite breath-hold divers. *Respir Physiol Neurobiol* 200;159(2):171-7. DOI: 10.1016/j.resp.2007.06.014.
22. Dippacher S, Willaschek C, Buchhorn R. Different nutritional states and autonomic imbalance in childhood. *Eur J Clin Nutr* 2014; 1-3. DOI: 10.1038/ejcn.2014.198.
23. Srivastava D, Singh K, Misra R, Prakash A. Comparison of cardiac autonomic activity between malnourished and healthy children. *Indian J Physiol Pharmacol* 2012; 56(3):222-8.
24. World Health Organization, UNICEF. WHO child growth standards and the identification of severe acute malnutrition in infants and children: joint statement by the World Health Organization and the United Nations Children's Fund. 2009.
25. Lohman TG, Roche AF, Martorell R. Anthropometric standardization reference manual. 1988.
26. Vanderlei LC, Silva RA, Pastre CM, Azevedo FM, Godoy MF. Comparison of the Polar S810i monitor and the ECG for the analysis of heart rate variability in the time and frequency domains. *Braz J Med Biol Res* 2008;41(10):854-9. DOI: <http://dx.doi.org/10.1590/S0100-879X2008005000039>.
27. Godoy MF, Takakura IT, Correa PR. Relevância da análise do comportamento dinâmico não linear (Teoria do Caos) como elemento prognóstico de morbidade e mortalidade em pacientes submetidos à cirurgia de revascularização miocárdica. *Arq Ciênc Saúde* 2005;12(4):167-71. DOI: <http://dx.doi.org/10.1590/S0102-76382009000200018>.
28. Grassberger P. Information flow and maximum entropy measures for 1-D maps. *Physica D* 1985;14(3):365-73. DOI: 10.1016/0167-2789(85)90095-8.
29. Landau HJ. Maximum entropy and maximum likelihood in spectral estimation. *IEEE Trans Inform Theory*, 1998;44(3):1332-6. DOI: 10.1109/18.669428.
30. Mann ME, Lees JM. Robust estimation of background noise and signal detection in climatic time series. *Clim Chang* 1996;33(3):409-45. DOI: 10.1007/BF00142586.
31. Donaldson GC, Seemungal TA, Hurst JR, Wedzicha JA. Detrended fluctuation analysis of peak expiratory flow and exacerbation frequency in COPD. *Eur Respir J* 2012;40(5):1123-9. DOI: 10.1183/09031936.00180811.
32. Peng CK, Havlin S, Stanley HE, Goldberger AL. Quantification of scaling exponents and crossover phenomena in nonstationary heartbeat time series. *Chaos* 1995;5(1):82-7. DOI: <http://dx.doi.org/10.1063/1.166141>.
33. Anderson TW, Darling DA. A test of goodness of fit. *J Am Statist Assoc* 1954;49(268):765-9. DOI: 10.1080/01621459.1954.10501232.
34. Razali NM, Wah YB. Power comparisons of shapiro-wilk, kolmogorov-smirnov, lilliefors and anderson-darling tests. *J Statist Model Analyt* 2011;2(1):21-33.
35. Kruskal WH, Wallis WA. Use of ranks in one-criterion variance analysis. *J Am Statist Assoc* 1952;47(260):583-621. DOI: 10.2307/2280779.
36. Hsu JC. Multiple comparisons in the general linear model. *Multiple Comparisons*. Springer; 1996. p. 181-226. DOI: 10.2307/1390767.
37. Jolliffe IT. *Principal Component Analysis*, Series: Springer Series in Statistics. New York: Springer; 2002. DOI: 10.1007/b98835.
38. Bernardo AFB, Vanderlei LCM, Garner DM. HRV Analysis: A clinical and diagnostic tool in Chronic Obstructive Pulmonary Disease. *International Scholarly Research Notices* 2014; 673232:6. DOI: <http://dx.doi.org/10.1155/2014/673232>.
39. Roque AL, Valenti VE, Massetti T, da Silva TD, Monteiro CB, Oliveira FR, et al. Chronic obstructive pulmonary disease and heart rate variability: a literature update. *Int Arch Med* 2014;7(1):43. DOI: 10.1186/1755-7682-7-43.
40. Sucharita S, Sowmya S, Thomas T, Kurpad AV, Vaz M. Plasma Vitamin B 12, Methylmalonic Acid and Heart Rate Variability in Healthy Young Indian Adults. *Int J Vitam Nutr Res* 2013;83(3):147-53. DOI: 10.1024/0300-9831/a000155.

RESUMO

O objetivo deste estudo é avaliar o risco de doenças dinâmicas em crianças desnutridas. Isto é feito por meio da aplicação de novas técnicas caóticas globais para os intervalos R-R do ECG. A variabilidade da frequência cardíaca (VFC) é uma ferramenta de baixo custo e não-invasiva que mensura os impulsos autonômicos. Na desnutrição ocorreu uma diminuição na resposta caótica da VFC. Setenta crianças foram divididas em grupos iguais e a VFC foi monitorada por 20 a 25 minutos. O parâmetro caótico avançado (CFP), que utiliza os três parâmetros caóticos globais é o algoritmo mais robusto. Estes três parâmetros são: alta entropia espectral (*hsEntropia*), análise da alta flutuação depurada espectral (*hsDFA*) e método espectral multi-taper (sMTM). *hsEntropia* é uma função da irregularidade da amplitude e frequência dos picos dos espectros de potência. É obtida pela aplicação da entropia de Shannon ao método multi-taper do espectro de potência. Para derivar *hsDFA* calculamos a adaptação espectral exatamente da mesma forma que para o *hsEntropia* utilizando um método adaptativo multi-taper do espectro de potência com as mesmas configurações; mas, em vez da entropia de Shannon, o DFA é o algoritmo aplicado. sMTM é a área entre o método multi-taper do espectro de potência e a linha de base. Foi utilizado o teste de normalidade de *Anderson-Darling* e *Lilliefors*. Para análise estatística foi utilizado o teste de *Kruskal-Wallis* com nível de significância de $p < 0.01$. A análise de Componentes Principais (PCA) identificou dois componentes que representam 100% da variância total. Desequilíbrio autonômico mensurado pela VFC e um aumento do risco cardiovascular são descritos em crianças com excesso de peso, bem como para desnutridos e indivíduos com anorexia nervosa. A relação entre a desnutrição e as medidas de complexidade é útil na avaliação do risco de doenças dinâmicas associadas com a condição. Isto fornece suporte no tratamento, auxiliando na determinação do nível de intervenção dietética ou farmacológica especialmente em relação às doenças dinâmicas.

Palavras-chave: desnutrição; dinâmica não linear; método multi-taper; variabilidade da frequência cardíaca.