

Robust Estimation of the Scaling Exponent in Detrended Fluctuation Analysis of Beat Rate Variability

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Abstract

Detrended fluctuation analysis is a popular method for studying fractal scaling properties in time series. The method has been successfully employed in studying heart rate variability and discovering distinct scaling properties in different pathological conditions. Traditionally the analysis has been performed by extracting two scaling exponents from linear fits, for short- and long-range correlations respectively. The extent of these ranges is subjective and the linear two-range model potentially disregards additional information present in the data.

Here we present a method based on the Kalman smoother for obtaining a whole spectrum of scaling exponents as a function of the scale. Additionally, we present an optimization scheme to obtain data-adaptive segmentation of the fluctuation function into approximately linear regimes. The methods are parameter-free and resistant to statistical noise in the fluctuation function.

We employ the methods in the analysis of the heart rate variability of patients with different heart conditions. The methods enhance the classification of these conditions, revealing more complex structure in the scaling exponents beyond the two-range model.

1. Introduction

The relationship between the autonomic nervous system and cardiovascular health has long been recognized. Their compounded effect is manifested in heart rate variability (HRV), and alterations in HRV may be indicative of various cardiovascular diseases [1]. Detrended fluctuation analysis (DFA), originally developed for studying long range correlations in DNA sequences [2], has been a popular method for studying self-affinity in signals. By computing the mean fluctuations $F(s)$ around a trend at multiple scales s , DFA is applied to assess power law scaling $F(s) \propto s^\alpha$ described by the scaling exponent α .

In the context of HRV, DFA is traditionally performed on the signal consisting of interbeat intervals, and the scaling exponent is determined by linear regression on a log-

log plot of the fluctuation function $F(s)$. Conventionally the linear regression is applied separately to two regimes to obtain short- and long-range scaling exponents [3]. This approach has been criticized as over-simplification, and alternative methods have been proposed [4, 5].

We propose two methods to complement the analysis. We present a method for obtaining a smooth spectrum of scaling exponents as a function of the scale, and an optimization scheme for determining a segmentation of the fluctuation function into approximately linear regimes.

2. Detrended fluctuation analysis

In the standard DFA formulation [3], integrated time series is divided into N_s non-overlapping windows of size s , from which the local trend is calculated in each window w by a least squares fit of a low order polynomial. The squared fluctuations $F_{s,w}^2$ in each window are defined as the mean squared differences from the local trend. We compute an error estimate for the fluctuation function as follows.

Let μ_s and ϵ_s denote the mean and its standard error, respectively, of the squared fluctuations over all windows of size s . The fluctuation function $F(s)$ and its error estimate $\Delta F(s)$ are then given by

$$F(s) = \sqrt{\mu_s}; \quad \Delta F(s) \approx \frac{\epsilon_s}{2\sqrt{\mu_s}}. \quad (1)$$

We denote their logarithmic counterparts by tildes, $\tilde{F}(\tilde{s})$ and $\Delta\tilde{F}(\tilde{s})$, respectively.

3. Kalman filter and smoother

Let us consider a linear probabilistic state space model of a system, described by its hidden state $\mathbf{x}_k \in \mathbb{R}^n$ and yielding the measurement $\mathbf{y}_k \in \mathbb{R}^m$ at the step k , described by the equations

$$\mathbf{x}_k = \mathbf{A}_{k-1}\mathbf{x}_{k-1} + \mathbf{q}_{k-1} \quad (2)$$

$$\mathbf{y}_k = \mathbf{H}_k\mathbf{x}_k + \mathbf{r}_k. \quad (3)$$

where $\mathbf{A}_{k-1} \in \mathbb{R}^{n \times n}$ and $\mathbf{H}_k \in \mathbb{R}^{m \times n}$ are the state transition and measurement model matrices, respectively. The

system is disturbed by process noise $\mathbf{q}_{k-1} \sim \mathcal{N}(\mathbf{0}, \mathbf{Q}_{k-1})$ and measurement noise $\mathbf{r}_k \sim \mathcal{N}(\mathbf{0}, \mathbf{R}_k)$, modeled by zero mean Gaussian noise with covariance matrices $\mathbf{Q}_{k-1} \in \mathbb{R}^{n \times n}$ and $\mathbf{R}_k \in \mathbb{R}^{m \times m}$.

The Kalman filter [6] provides a recursive closed-form minimum-mean-squared error solution for the posterior filtering distributions $p(\mathbf{x}_k | \mathbf{y}_{1:k}) = \mathcal{N}(\mathbf{m}_k, \mathbf{P}_k)$, given all the measurements up to the current step k . Similarly, the Kalman smoother [7] allows the computation of the smoothing distributions $p(\mathbf{x}_k | \mathbf{y}_{1:T}) = \mathcal{N}(\mathbf{m}_k^s, \mathbf{P}_k^s)$, that take into account all the T measurements to provide an estimate for the state of the system at the step $k \leq T$. The smoothing distributions are calculated from the filtering distribution means \mathbf{m}_k and covariances \mathbf{P}_k by backwards recursion. For a modern exposition about the subject, see, e.g., Ref. [8].

The scaling exponent $\alpha(s)$ is estimated from the logarithmically transformed quantities. The dynamic model applied in this paper assumes that the exponent remains approximately constant between adjacent window sizes, except for tiny perturbations derived from the data. The hidden state \mathbf{x}_k to be estimated then consists of the observed fluctuation function values and its derivative, which corresponds to the scaling exponent: $\mathbf{x}_k = [\tilde{F}_k \quad \alpha(\tilde{s}_k)]^\top$. Here \tilde{s}_k are the utilized logarithmic DFA window sizes in ascending order and $\tilde{F}_k = \tilde{F}(\tilde{s}_k)$. The state transition and measurement model matrices are then

$$\mathbf{A}_k = \begin{bmatrix} 1 & h_{k+} \\ 0 & 1 \end{bmatrix}; \quad \mathbf{H} = [1 \quad 0], \quad (4)$$

where $h_{k+} = \tilde{s}_{k+1} - \tilde{s}_k$ is the forward difference of the window size at k . The error estimate of the fluctuation function is readily utilized as the measurement noise covariance by $R_k = [\Delta \tilde{F}(\tilde{s}_k)]^2$, which is now simply a scalar.

The scaling exponent is gently perturbed by the process noise, allowing it to vary as a function of the window size. The noise covariance is estimated by taking a weighted sample variance of the finite difference approximation [9] of the logarithmic fluctuation function derivative. The derivative is estimated at each window size \tilde{s} by

$$\tilde{F}'_k \approx \frac{h_{k-}^2 \tilde{F}_{k+1} + (h_{k+}^2 - h_{k-}^2) \tilde{F}_k - h_{k+}^2 \tilde{F}_{k-1}}{h_{k-} h_{k+} (h_{k+} + h_{k-})}, \quad (5)$$

where $h_{k-} = \tilde{s}_k - \tilde{s}_{k-1}$ are the backwards differences. The boundaries are considered by assuming that the function continues linearly. The uncertainty $\Delta \tilde{F}_k$ in the fluctuation function is propagated through this expression to obtain squared error estimates $\Delta \tilde{F}'_k{}^2$ for these derivatives.

The weighted sample variance $\hat{\sigma}^2$ of the derivatives is computed with the weights taken to be inversely proportional to these squared error estimates. This variance estimates the magnitude of the nudges that the scaling exponent

experiences between logarithmic window sizes, leading to the following process noise covariance:

$$\mathbf{Q}_k = \hat{\sigma}^2 \begin{bmatrix} \frac{1}{3} h_{k+}^3 & \frac{1}{2} h_{k+}^2 \\ \frac{1}{2} h_{k+}^2 & h_{k+} \end{bmatrix}. \quad (6)$$

The prior distribution $\mathcal{N}(\mathbf{m}_0, \mathbf{P}_0)$ is estimated from the first data points, with \tilde{F}_1 and \tilde{F}'_1 as the mean and the squares of their error estimates as the covariance.

4. Integer linear programming

Let us consider the problem of optimally segmenting the logarithmic fluctuation function \tilde{F}_k into approximately linear regimes. First we compute all the possible n linear regression fits to the fluctuation function that cover some desired minimum number of different window sizes. The optimal segmentation is then sought by solving the following linear programming problem

$$\arg \min_{\mathbf{x}} \mathbf{c}^\top \mathbf{x}; \quad \mathbf{c} \in \mathbb{R}^n; \quad \mathbf{x} \in \{0, 1\}^n, \quad (7)$$

where the components x_i of the binary vector \mathbf{x} indicate whether the i -th fit is utilized in the optimal segmentation. The components c_i of the coefficient vector \mathbf{c} are the residual sum of squares RSS_i of the linear regression fits.

For expressing the optimization constraints we introduce the auxiliary variables a_{ik} , defined to be equal to unity if the i -th fit covers the k -th window size and zero otherwise. The constraints are then given by the following equations

$$\sum_{i=1}^n a_{ik} = 1; \quad \sum_{i=1}^n x_i = N, \quad (8)$$

where the former relations ensure that each point in the fluctuation function is covered by exactly one linear fit, and the latter relation ensures that the solution consists of the desired number of linear fits N . The problem is then readily solved by any existing integer linear programming suite.

Choosing an optimal value for the number of linear segments, N , is a partial and subjective decision. The approach taken here proceeds as follows. Let $\text{RSS}(N)$ denote the total residual sum of squares of the optimal segmentation with N segments. This residual is trivially reduced by increasing the number of segments. We seek a solution that consists of as few segments as admissible for equitable segmentation. Therefore, we consider the desirability $D(N) \propto 1/[N \cdot \text{RSS}(N)]$ of a solution to be inversely proportional to the number of segments and the total residual sum of squares. The solution is optimal when it maximizes this quantity.

This optimization scheme may be performed separately for each fluctuation function or simultaneously to a group

of fluctuation functions, provided that they all have been calculated at compatible window sizes. In the latter case the residual sum of squares RSS_{ij} of the i -th fit in the j -th fluctuation function are summed over all the fluctuation functions when computing the coefficients c_i .

5. Data and methods

The analysis is performed on the publicly available resources of PhysioBank [10], utilizing the following databases: Normal Sinus Rhythm RR Interval Database, The Long-Term ST Database, The Long-Term AF Database, Congestive Heart Failure RR Interval Database, and The BIDMC Congestive Heart Failure Database.

Distinguishing intrinsic variations in the heart rate from those induced by extrinsic influences in long term recordings may be perplexing, resulting in unknown trends in the data. While DFA is successful in detrending simple long-term trends, its ability to cope with complex nonlinear trends is disputed [11]. Therefore the analysis is additionally performed by explicitly removing a trend determined by the local median in a moving window of 101 beats wide. In both cases DFA is performed with linear detrending in 45 logarithmically distributed window sizes of 5–200 beats.

The scaling exponent α is determined from the fluctuation functions with three methods: the traditional linear fits in short (5–16 beats) and long (16–64 beats) range regimes, the optimal linear segmentation applied to the whole dataset at once, and the alpha spectra obtained with the Kalman smoother. These scaling exponents are then utilized as features in the classification task of identifying the different pathological conditions. The classification is performed by various algorithms available in the scikit-learn Python module [12]. Hyperparameters are optimized with a simple grid search.

6. Results

The Kalman smoother is found to be capable of obtaining robust estimates for the scaling exponent, even in the presence of moderate noise in the fluctuation functions. This is demonstrated in Fig. 1, where we observe that the alpha spectrum from the noisy fluctuation function follows very closely the spectrum from the smoother fluctuation function.

Performing the optimal linear segmentation on the whole data set reveals four segments in the following ranges: 5–12, 13–32, 35–81, and 87–200 beats. The scaling exponents from the explicitly detrended data are illustrated in Fig. 2 for the different pathological conditions. The results suggest that two scaling exponents may be insufficient for adequately describing the scaling behavior. Yet even with the more detailed methods the average scaling

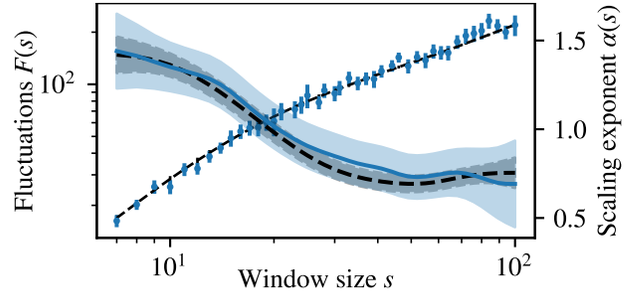


Figure 1. Performance of the Kalman smoother on noisy fluctuation function (blue dots with error bars). The smooth fluctuation function (dashed) is obtained by utilizing overlapping windows in its calculation. The corresponding alpha spectra are also shown with their error estimates.

behavior is very similar in healthy individuals and in those suffering from ST episodes.

Classification results with leave-one-out cross-validation are shown in Fig. 3. The more detailed picture depicted by the new methods results in moderately better classification performance over the traditional two-range model, for both multiclass and binary classification tasks. Particularly in the binary classification of healthy versus AF and CHF, the misclassification rates are reduced by approximately 40 %. However, ST episodes cannot be reliably diagnosed from the overall scaling behavior of long-term recordings. Results from the optimized linear segmentation are omitted, as they are already very similar to the results obtained with the smooth alpha spectra.

Curiously, the data detrended by the local median yields better classification results, despite the detrending removing long-range correlations. However, the classes become more distinct in the α_3 -regime, which may be caused by the detrending removing spurious correlations arising from extrinsic trends.

7. Conclusions

We have introduced methods for robustly determining the DFA scaling exponent, and a spectrum of exponents, from noisy fluctuation functions in a parameter-free manner. The methods have been tested by classifying different heart conditions according to the scaling exponents of interbeat intervals. The results support the hypothesis that the traditional two-range model for the scaling exponents can be insufficient. Furthermore, the significance of extrinsic trends in the heart rate variability may warrant further research. The presented methods enhance the classification of heart rate data covering different conditions, which could open up a path into predictive cardiological tools.

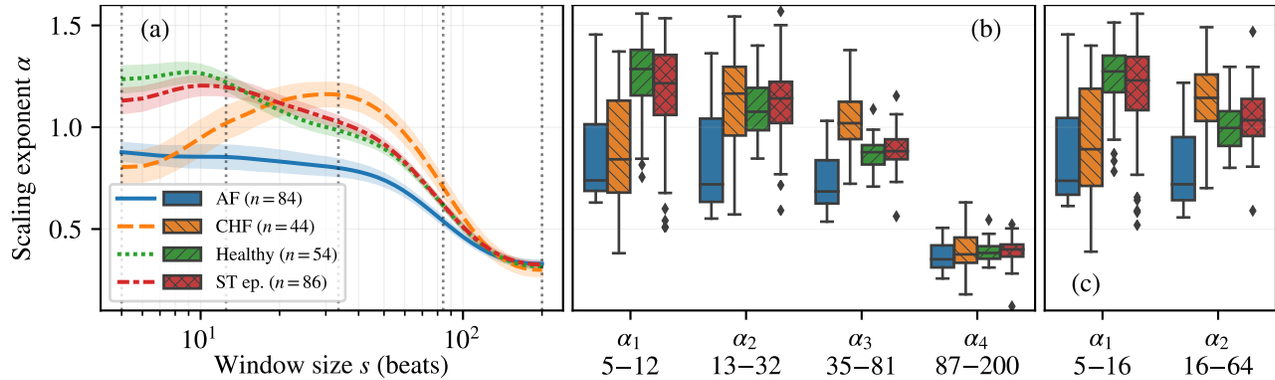


Figure 2. (a) The Kalman smoother estimates for the mean scaling exponents and their 95 % confidence intervals. Box plots of the scaling exponents for the optimal linear segmentation (b) and the traditional short- and long-range regimes (c). The numbers denote the fitting range in beats for the linear regression.

Healthy	0.76	0.24	Healthy	0.89	0.11	Healthy	0.19	0.81	AF	0.61	0.024	0.012	0.36
AF	0.2	0.8	CHF	0.25	0.75	ST ep.	0.12	0.88	CHF	0.16	0.57	0	0.27
Healthy	0.85	0.15	Healthy	0.96	0.037	Healthy	0.35	0.65	Healthy	0.15	0.019	0.13	0.7
AF	0.12	0.88	CHF	0.18	0.82	ST ep.	0.28	0.72	ST ep.	0.16	0.12	0.035	0.69
	Healthy	AF		Healthy	CHF		Healthy	ST ep.	AF	0.75	0	0.036	0.21
									CHF	0.091	0.61	0.023	0.27
									Healthy	0.019	0.019	0.39	0.57
									ST ep.	0.13	0.058	0.1	0.71
									AF				
									CHF				
									Healthy				
									ST ep.				

Figure 3. Confusion matrices of the classification task with support vector machines for the traditional short- and long-range alphas (upper) and for the Kalman smoother alpha spectra (lower) on the detrended data.

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