

Comparison of Methods for the Assessment of Nonlinearity in Short-Term Heart Rate Variability under different Physiopathological States

Luca Faes,¹ Manuel Gómez-Extremera,² Riccardo Pernice,¹ Pedro Carpena,^{2,3} Giandomenico Nollo,⁴ Alberto Porta,^{5,6} and Pedro Bernaola-Galván^{2,3}

¹*Department of Engineering, University of Palermo, 90128 Palermo, Italy*

²*Dpto. de Física Aplicada II, ETSI de Telecomunicación, University of Málaga, 29071 Málaga, Spain*

³*Instituto Carlos I de Física Teórica y Computacional, University of Málaga, 29071 Málaga, Spain*

⁴*Department of Industrial Engineering, University of Trento, 38123 Trento, Italy*

⁵*Department of Biomedical Sciences for Health, University of Milan, 20122 Milan, Italy*

⁶*Department of Cardiothoracic, Vascular Anesthesia and Intensive Care, IRCCS Policlinico San Donato, 20097 San Donato Milanese, Milan, Italy*

Despite the widespread diffusion of nonlinear methods for heart rate variability (HRV) analysis, the presence and the extent to which nonlinear dynamics contribute to short-term HRV is still controversial. This work aims at testing the hypothesis that different types of nonlinearity can be observed in HRV depending on the method adopted and on the physiopathological state. Two entropy-based measures of time series complexity (normalized complexity index, NCI) and regularity (information storage, IS), and a measure quantifying deviations from linear correlations in a time series (Gaussian linear contrast, GLC), are applied to short HRV recordings obtained in young (Y) and old (O) healthy subjects and in myocardial infarction (MI) patients monitored in the resting supine position and in the upright position reached through head-up tilt. The method of surrogate data is employed to detect the presence and quantify the contribution of nonlinear dynamics to HRV. We find that the three measures differ both in their variations across groups and conditions and in the percentage and strength of nonlinear HRV dynamics. NCI and IS displayed opposite variations suggesting more complex dynamics in O and MI compared to Y, and less complex dynamics during tilt. The strength of nonlinear dynamics is reduced by tilt using all measures in Y, while only GLC detects a significant strengthening of such dynamics in MI. A large percentage of detected nonlinear dynamics is revealed only by the IS measure in the Y group at rest, with a decrease in O and MI and during T, while NCI and GLC detect lower percentages in all groups and conditions. While these results suggest that distinct dynamic structures may lie beneath short-term HRV in different physiological states and pathological conditions, the strong dependence on the measure adopted and on their implementation suggests that physiological interpretations should be provided with cautiousness.

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Historically, the study of heart rate variability (HRV) has both received clinical attention, e.g. as a tool for risk stratification after myocardial infarction, and has attracted the interest of physicists who saw it as a particularly lucid example of chaos in physiology. Later on, after it was realized that a thorough evaluation of the chaotic nature of cardiac dynamics is precluded by difficulties inherent in the noisy nature of biological signals and in the restricted length of the data typically available, the field of HRV analysis underwent a shift in paradigm from chaos to complexity (intended as unpredictability) and detection of nonlinear dynamics in short-term HRV (up to a few minutes of recordings) analyzed in different pathophysiological states. The latter issue remains elusive, due to the difficulty of reliably assessing nonlinearity over short time series, to the proliferation of diverse nonlinear analysis methods each with its own strengths and limitations, and to the changing nature of nonlinear HRV dynamics across states and conditions. The present study contributes to settle this issue, implementing different of state-of-art nonlinear dynamic measures and comparing them as regards the detection of the presence and the contribution of nonlinear dynamics to short-term HRV. The comparison is

performed considering the progression across healthy and pathological states (i.e., aging and myocardial infarction) and investigating the effects on the cardiac dynamics of a specific physiological stressor (i.e., head-up tilt).

I. INTRODUCTION

Human heart rate variability (HRV), commonly assessed measuring the spontaneous beat-to-beat changes in the duration of the RR interval of the ECG, is the result of the activity of different physiological control systems which operate across multiple time scales to let the body functions adapt to environmental, physical and psychological challenges^{1,7}. RR interval fluctuations have been classically represented as a linear superposition of rhythms², leading to remarkable time- and frequency-domain descriptions of the factors contributing to the neuroautonomic modulation of the heart rhythm in healthy conditions, as well as of the alteration of these factors related to a variety of pathological states^{4,8,18,22,39,42}. Nevertheless, since the cardiac control is typically accomplished through the interaction among multiple complex regulatory

mechanisms, including self-sustained oscillators as well as control loops³¹, the linear description of the RR interval variability may be severely limited and disregard significant dynamical features.

As a consequence, a variety of nonlinear approaches to time series analysis have been devised to characterize RR interval fluctuations and extract additional physiological and clinical information from HRV^{60,64}. A class of these approaches, focused on long-term analyses spanning scales up to several hours, is mainly based on using methods able to assess scaling properties, long-range correlations, and multifractality of the RR time series^{5,24,30,46}. These nonlinear methods were often employed with the aim of identifying signatures typical of chaotic dynamics in long-term HRV recordings, leading to an animated discussion of this topic¹⁹. Besides the presence or absence of chaos^{14,21,28}, there is substantial consensus about the fact that long-term RR interval time series are nonlinear and multifractal, and that the scaling behavior of HRV is altered with aging or during physical exercise, and under pathological conditions such as myocardial infarction^{6,20,23,25,29}.

On the other hand, it is also widely accepted that the assessment of HRV over temporal scales ranging from seconds to a few minutes allows the indirect investigation of the mechanisms underlying the short-term cardiovascular control^{13,41} and this assessment might require nonlinear methods better suited for the evaluation of complex aspects of HRV dynamics. In fact, a number of nonlinear measures have been developed to this end, e.g. based on nonlinear prediction^{53,57}, entropy or mutual information^{50,66}, time irreversibility^{54,62}, or phase coupling^{3,12}. These and other studies have provided ample evidence that changes in nonlinear descriptors of short-term HRV such as complexity or regularity indexes, either induced by the modification of the experimental conditions or determined by spontaneous transitions among physiological states, can be reliably detected and associated to alterations of the autonomic control. Notwithstanding this, the presence and impact of nonlinear dynamics in short-term HRV is still a controversial issue. Some studies suggested that nonlinear components of HRV are of limited importance in resting conditions and are evoked by the presence of a dominant respiratory sinus arrhythmia^{53,57}, or in association with respiratory inputs to the cardiovascular system^{17,26}. Conversely, other studies assessing temporal asymmetries suggested that nonlinearities are relevant at rest and may be present even in conditions of small respiratory sinus arrhythmia⁵⁴. The contribution of the two branches of the autonomic nervous system to nonlinear HRV dynamics remains elusive and is likely linked to the time scales of their functioning⁵². Moreover, nonlinear dynamics might be sustained by the interaction between sympathetic and parasympathetic activities³.

Methodologically, it has been suggested that multiple nonlinear components, operating at different scales and possibly interacting with each other, may concur to the generation of short-term HRV^{3,27,52}. Since these different components of HRV nonlinear dynamics may be captured in a different way by different metrics, the aim of the present study is to test the hypothesis that distinct types of nonlinear dynamics underlie HRV during different physiopathological states. To this

end we apply three nonlinear dynamic measures to the RR interval time series measured in young and old healthy subjects, as well as in post acute myocardial infarction (AMI) patients, monitored at rest and during sympathetic activation induced by postural change. The measures were selected, among those allowing the assessment of nonlinear dynamics in short-term HRV (~ 300 points), to investigate such dynamics from different perspectives, exploiting different concepts and implementing different estimators. In particular, two of the measures quantify the common concepts of complexity and regularity previously defined in time series analysis with the indexes of Sample Entropy⁵⁸ and Information Storage³⁸, implementing them through refined estimation techniques devised recently^{51,66}. The third measure is defined building upon several previous works^{9,11,34-37} to quantify the deviation from linearity of the correlation structure of the observed time series according to a recently proposed Gaussian Linear Contrast method (GLC)¹⁰. The application of these approaches in conjunction with the method of surrogate data^{59,61} allows us to quantify the extent to which nonlinear dynamics impact on short-term HRV in different conditions of autonomic nervous system imbalance, also investigating the effects of age and pathology. The database used in the study is made publicly available to favor reproducibility and encourage the comparison with different nonlinear dynamic measures.

II. NONLINEAR DYNAMIC MEASURES

This section describes the methods used in the present work to quantify nonlinear dynamics in the temporal statistical structure of a system evolving in time. Our starting point is an experimental time series $\{s_n\}$, $n = 1, 2, \dots, N$, which is considered as a realization of a stochastic process S describing the evolution over time of an observed dynamical system \mathcal{S} . The process S is considered stationary, so that the random variables obtained sampling the process at the time n (i.e. S_n , $n = 1, 2, \dots, N$), are identically distributed with marginal probability density function $f(\cdot)$ and cumulative density function $F(\cdot)$. Moreover, without loss of generality, we assume that each S_n has zero mean and unit standard deviation.

To assess nonlinear dynamics in the stochastic process S we look at its temporal correlation structure: while for purely linear dynamics the dependence between s_n and $s_{n-\ell}$ is linear for any lag ℓ , in the case of nonlinear dynamics such dependence cannot be studied only in terms of linear correlations. In the first two methods considered, nonlinear correlations are investigated within an information-theoretic framework, separating the present state of the system S_n from its past states $S_n^- = [S_1, \dots, S_{n-2}, S_{n-1}]$ and quantifying their information content in terms of entropy measures⁶⁶. In fact, when the system transits from past states to a new state, new information is produced in addition to the information that is already carried by the past states. The rate of generation of new information is inversely related to the strength of nonlinear correlations in the process, while the information shared between the present and the past variables is directly related

to such correlations⁶⁶. On this basis, the measures of conditional entropy (Sect. II A) and information storage (Sect. II B) assess nonlinear correlations quantifying respectively the new information contained in S_n but not in S_n^- , and the amount of information carried by S_n that can be explained by S_n^- .

The third method takes its roots on the observation that a purely linear stochastic process is considered to have only linear correlations. However, the most feasible models for generation of pure linear processes give a Gaussian marginal distribution as output, so that linear processes with marginal Gaussian distributions are usually considered as reference for linearity. This can lead to false positive cases by nonlinearity methods when applied to linear processes with **non-Gaussian** marginal distribution. In the GLC (see Sect. II C) we test nonlinearity based only on the nature of correlations, which truly reflect the dynamics, and discard those effects induced by non Gaussian marginal distributions.

A. Complexity Index based on Local Sample Entropy

The information-theoretic assessment of nonlinear correlations in a dynamic process is based on applying the concepts of entropy and conditional entropy to the random variables representing the present and past states of the process. Given two generic continuous (possibly vector) random variables X and Y , the entropy of X and the conditional entropy of X given Y are defined as

$$H(X) = -\mathbb{E}[\log f(x)] = -\int_{\mathcal{D}_X} f(x) \log f(x) dx, \quad (1)$$

$$H(X|Y) = -\mathbb{E}[\log f(x|y)] = H(X, Y) - H(Y), \quad (2)$$

where \mathcal{D}_X is the domain of X , $f(x)$ and $f(x|y)$ are the probability density of X and the conditional probability of X given Y , and $\mathbb{E}[\cdot]$ is the expectation operator; the term $H(X, Y)$ in (2) is the joint entropy of X and Y , obtained generalizing (1) to the joint probability density $f(x, y) = f(x|y)f(y)$. Particularizing these definitions to the variables S_n and S_n^- describing the present and the past states of the process S , the conditional entropy becomes

$$H(S_n|S_n^-) = H(S_n^-, S_n) - H(S_n^-) \quad (3)$$

The conditional entropy quantifies the amount of information contained in the present of the process that cannot be explained by its past history: if the process is fully random, the system produces information at the maximum rate, yielding maximum conditional entropy; if, on the contrary, the process is fully predictable, the system does not produce new information and the conditional entropy is zero.

In the present work, practical computation of the conditional entropy is performed adopting kernel estimates of the probability density functions⁶⁶. In particular, we make use of the well known Sample Entropy index⁵⁸, improved through the implementation of a local version of the estimator⁵¹. The Sample Entropy (SampEn) estimates $H(S_n|S_n^-)$ in (3) first

truncating S_n^- to $S_n^m = [S_{n-1}, S_{n-2}, \dots, S_{n-m}]$, and then approximating $H(S_n^m, S_n)$ and $H(S_n^m)$ as the negative logarithm of the average joint probability of finding a pattern in the neighborhood of the reference pattern with a tolerance r in the $(m+1)$ dimensional and m -dimensional embedding space, namely

$$\text{SampEn}(m, r, n) = -\ln \langle p(S_n^m, S_n) \rangle + \ln \langle p(S_n^m) \rangle, \quad (4)$$

where $p(S_n^m, S_n)$ is the probability that the pattern $S_n^m \oplus S_n$ assumes the value $s_n^m \oplus s_n$, $p(S_n^m)$ is the probability that the pattern S_n^m takes the value s_n^m and $\langle \cdot \rangle$ performs the average over time (i.e., over all values $s_n^m \oplus s_n$). SampEn is a robust estimator of irregularity given that the log-of-zero situation is extremely unlikely because the logarithm is applied to the average of a quantity that has 0 as the lowest bound. However, as a consequence of computing an average over time, SampEn has the disadvantage to be a global marker of irregularity that might not represent reliably the local behavior in the neighborhood of a specific pattern and blur nonlinear features⁵¹. A local version of SampEn (LSampEn) was proposed in Ref.⁵¹ by directly approximating $H(S_n|S_n^m)$ instead of its constituents (i.e., $H(S_n^m, S_n)$ and $H(S_n^m)$) as

$$\text{LSampEn}(m, r, n) = -\ln \langle p(S_n|S_n^m) \rangle, \quad (5)$$

where $p(S_n|S_n^m)$ is the conditional probability that the current state S_n assumed the value s_n given that the past state S_n^m is s_n^m . The average operator makes the estimator robust against the log-of-zero situation and the estimation of $p(S_n|S_n^m)$ renders LSampEn a local estimator of irregularity given that the quantity being averaged referred specifically to the reference pattern S_n^m . To limit the consequence that, when solely S_n^m is found in the neighborhood of S_n^m , $p(S_n|S_n^m)$ is unreliably high⁴⁹, we applied the correction proposed by Porta et al.⁵¹, namely in this unfortunate case $p(S_n|S_n^m)$ is set to $(N - m + 1)^{-1}$ corresponding to the maximum uncertainty computable over the series. The resulting estimator, applied to the time series reduced to unit variance, is denoted as Normalized Complexity Index (NCI)⁵¹.

B. Regularity Index based on Information Storage

Information measures can be exploited also for evaluating in a direct way the strength of nonlinear correlations in the dynamical structure of a stochastic process, so that to assess its degree of regularity. To this end, a relevant entropy measure is the so-called information storage, which quantifies the amount of information shared between the present and the past observations of the considered process. The information storage of the process S is defined as:

$$I(S_n; S_n^-) = H(S_n) + H(S_n^-) - H(S_n^-, S_n), \quad (6)$$

where $I(\cdot; \cdot)$ denotes mutual information. The information storage reflects the degree to which information is preserved in a time-evolving system⁶⁵. As such, it measures how much of the uncertainty about the present can be resolved by knowing the past: if the process is fully random, the past gives no

knowledge about the present, so that the information storage is zero; if, on the contrary, the process is fully predictable, the present can be fully predicted from the past, which results in maximum information storage. Note that information storage and conditional entropy of a dynamic process are inversely related to each other, and depend on the entropy of the present state of the process through the equation $I(S_n; S_n^-) + H(S_n|S_n^-) = H(S_n)$.

In practical analysis, the information storage can be estimated from a time series of finite length following the same principles of conditional entropy estimation. These include the use of a finite number of samples in the past to approximate the history of the observed process (i.e., S_n^- is truncated to $S_n^m = [S_{n-1}, S_{n-2}, \dots, S_{n-m}]$), and the adoption of non-parametric estimators of the probability density functions involved in the computation of $I(S_n; S_n^m)$. However, since computation of the measure defined in (6) requires to estimate three entropy terms involving variables of different dimensions, and since the bias of entropy estimates depends strongly on the dimension, implementation of standard histogram or kernel-based methods typically results in inaccurate estimates of the information storage^{15,66}. Here, while the kernel-based estimation of information storage is implemented for comparative purposes in the Supplementary material, to overcome this limitation we resort to nearest neighbor entropy estimation³² and implement a strategy for bias compensation specific of mutual information estimates³³. The nearest neighbor entropy estimate of a generic d -dimensional random variable X can be obtained from a set of realizations $\{x_1, x_2, \dots, x_N\}$ of the variable as³²

$$H(X) = \psi(N) - \psi(k) + d \langle \ln \varepsilon_n \rangle, \quad (7)$$

where ψ is the digamma function, ε_n is twice the distance between the outcome x_n and its k^{th} nearest neighbor computed according to the maximum norm (i.e., taking the maximum distance of the scalar components), and $\langle \cdot \rangle$ stands for average over N outcomes. Then, the information storage could be computed applying (7) to the three terms in (6). However, doing so would result in different distance lengths when approximating the probability density in different dimensions, and this would introduce different estimation biases that cannot be compensated by taking the entropy differences. To keep the same distance length in all explored spaces, we perform a neighbor search only in the highest-dimensional space (the one spanned by the realizations of S_n^m, S_n) and then project the distances found in this space to the lower-dimensional spaces (those spanned by the realizations of S_n^m and S_n), keeping these distances as the range within which neighbors are counted. Specifically, the knn estimate of $H(S_n^m, S_n)$ is computed through a neighbor search, i.e. fixing the number of neighbors k and computing the distance to the k^{th} neighbor, as:

$$H(S_n, S_n^m) = \psi(N) - \psi(k) + (m+1) \langle \ln \varepsilon_n \rangle, \quad (8)$$

where ε_n is twice the distance from (S_n, S_n^m) to its k^{th} nearest neighbor, and then, given the distances ε_n , the entropies in the lower-dimensional spaces are estimated through a range

search, i.e. fixing the distance ε_n and counting for the neighbors falling within this distance, as:

$$H(S_n^m) = \psi(N) - \psi(N_{S_n^m}) + m \langle \ln \varepsilon_n \rangle, \quad (9)$$

$$H(S_n) = \psi(N) - \psi(N_{S_n}) + \langle \ln \varepsilon_n \rangle, \quad (10)$$

where N_{S_n} and $N_{S_n^m}$ are the number of points whose distance from S_n and S_n^m , respectively, is smaller than $\varepsilon_n/2$. Finally, our estimate of the information storage is obtained subtracting Eq. (8) from the sum of Eqs. (9) and (10)⁵⁵:

$$IS = \psi(N) + \psi(k) - \langle \psi(N_{S_n^m}) \rangle - \langle \psi(N_{S_n}) \rangle. \quad (11)$$

C. Nonlinearity Index based on Gaussian Linear Contrast

As we stated above, GLC assesses nonlinearities related only to the nature of the correlations and not to the non-Gaussianity of the data. Let us consider an experimental time series $\{s_n\}$ ($n = 1, 2, \dots, N$), with non-Gaussian marginal distribution. The observed autocorrelation function of $\{s_n\}$ is given by

$$C_{\text{obs}}(\ell) = \langle s_n s_{n+\ell} \rangle \quad (12)$$

Using $C_{\text{obs}}(\ell)$, GLC tries to determine if $\{s_n\}$ is originated from a Gaussian time series $\{z_{G,n}\}$ ($n = 1, 2, \dots, N$) with only *linear correlations*, which have been transformed to have the observed marginal distribution using an invertible transformation of the experimental time series. If this is the case, then GLC assumes that $\{s_n\}$ is linear, and is non-linear otherwise.

The theoretical background of the GLC method is the following. Let us consider a pair of correlated Gaussian variables x_G and y_G , both of $\mathcal{N}(0, 1)$ type, so that their corresponding probability density and cumulative distribution are the standard Gaussian $\varphi(x_G)$ and $\Phi(x_G)$. We assume that x_G and y_G are only linearly correlated, with a correlation value C_G , i.e.

$$C_G = \langle x_G y_G \rangle \quad (13)$$

This is equivalent to affirm that the joint distribution of x_G and y_G is the bivariate Gaussian distribution $\varphi_2(x_G, y_G, C_G)$. Then, we transform x_G and y_G to the variables x and y , which follow the marginal distribution of the experimental time series. This can be done with the usual method:

$$x = F^{-1}[\Phi(x_G)], \quad y = F^{-1}[\Phi(y_G)] \quad (14)$$

with $F^{-1}(\cdot)$ the inverse cumulative distribution of the experimental time series. Since F^{-1} is fixed by $\{s_i\}$, the linear correlation C between x and y , i.e. $C = \langle xy \rangle$ depend solely on the C_G value. Indeed, since x and y depend formally on x_G and y_G (Eq. (14)) with joint distribution $\varphi_2(x_G, y_G, C_G)$, C can be calculated as (see Refs.^{9-11,34-37}):

$$C(C_G) \equiv \langle xy \rangle = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} F^{-1}(\Phi(x_G)) F^{-1}(\Phi(y_G)) \varphi_2(x_G, y_G, C_G) dy_G dx_G \quad (15)$$

Solving numerically the previous integral for a dense set of C_G values in the interval $(-1, 1)$ we characterize the $C(C_G)$ function, which contains the information on how the Gaussian correlations are transformed when the distribution of the variables is transformed from Gaussian to the experimental distribution.

These results can be extrapolated straightforwardly to time series. Let us consider a Gaussian time series $\{z_{G,n}\}$, with autocorrelation function $C_G(\ell)$ given by $C_G(\ell) \equiv \langle z_{G,n} z_{G,n+\ell} \rangle$. Note that $z_{G,n}$ and $z_{G,n+\ell}$ are equivalent to x_G and y_G in Eq. (13). Then, let us transform $\{z_{G,n}\}$ into a time series $\{z_n\}$ with the same marginal distribution of the experimental time series using Eq. (14) for each $z_{G,n}$ value. The autocorrelation function $C(\ell)$ of $\{z_n\}$ can be then calculated using Eq. (15) simply by replacing x_G , y_G and C_G by $z_{G,n}$, $z_{G,n+\ell}$ and $C_G(\ell)$ respectively. In other words, once the $C(C_G)$ function is known by using Eq. (15) (which only requires the marginal distribution of the experimental time series), then $C(\ell) = C(C_G(\ell))$. This last equality holds if the non-Gaussian time series $\{z_n\}$ really comes via the transformation (14) from the Gaussian and linearly correlated series $\{z_{G,n}\}$ since this is the condition used in Eq. (15) to determine $C(C_G)$. This property is the key point in the GLC method.

With this theoretical background, the steps to apply the GLC method on an experimental time series $\{s_n\}$ are the following:

- (i) Determine the observed autocorrelation function $C_{\text{obs}}(\ell)$ of the experimental time series $\{s_n\}$.
- (ii) Transform $\{s_n\}$ to have Gaussian distribution using the inverse of the transformation in Eq.(14), and calculate its autocorrelation function $C_{G'}(\ell)$. Note that if $\{s_n\}$ has been obtained from a Gaussian time series using the transformation (14), simply by inverting the transformation the hypothetical original Gaussian time series is recovered, as long as the transformation from Gaussian to $\{s_n\}$ is invertible. The knowledge of $C_{\text{obs}}(\ell)$ and $C_{G'}(\ell)$ for each ℓ allows to obtain the function $C_{\text{obs}}(C_{G'})$.
- (iii) Obtain the real $C(C_G)$ function using Eq.(15) by giving to C_G a great number of values in the interval $(-1, 1)$. In practice, and specially for short experimental time series, the numerical solution of the integral might be a harsh task: due to finite size effects it can be difficult to correctly estimate F^{-1} . Then, to calculate $C(C_G)$ we adopt a different strategy: we use autoregressive time series of order 1 (AR1) with the same size as $\{s_n\}$. An AR1 process is defined as: $z_{G,n} = \varphi z_{G,n-1} + \eta_n$ where $\{\eta_n\}$ is a Gaussian $\mathcal{N}(0, 1)$ white noise and $\varphi \in (-1, 1)$ is a constant. AR1 processes are Gaussian with purely linear correlations, with generic autocorrelation $C_G(\ell) = \varphi^\ell$, so that changing the φ value we

can obtain any value of Gaussian correlation in the interval $(-1, 1)$. Thus, when generating a large set of AR1 time series for different φ values, and calculate the autocorrelation function of all of them we would obtain a huge amount of data points densely populating the $(-1, 1)$ Gaussian correlation interval. Nevertheless, in order to save computation time, in practice, after preliminar analysis of $C_{\text{obs}}(C_{G'})$ we restrict the φ values to those which will generate Gaussian correlation values in our region of interest. Then, we transform all AR1 time series using Eq.(14) to have the marginal distribution of $\{s_n\}$, and also calculate the autocorrelation function $C(\ell)$ for all series. Note that each $C(\ell)$ value is the image of a Gaussian autocorrelation value. Finally, we bin the Gaussian correlation interval $(-1, 1)$ into 0.01 length bins, and put in each one the images of all the Gaussian correlation values contained in the bin. The average of all the images in the respective bin gives the C value corresponding to the Gaussian correlation at the center of the bin, so that finally we have a numeric determination of the $C(C_G)$ function.

- (iv) If the experimental time series $\{s_n\}$ is really obtained by transforming a Gaussian time series, then the Gaussian series is the one determined in step (ii), with autocorrelation function $C_{G'}(\ell)$, and the observed autocorrelation are given by $C_{\text{obs}}(\ell) = C_{\text{obs}}(C_{G'}(\ell))$. However, the expected correlations in $\{s_n\}$ if the Gaussian series is purely linear, $C_{\text{lin}}(\ell)$, should be given by the $C(C_G)$ function determined in step (iii) evaluated at the $C_{G'}(\ell)$ values, i.e. $C_{\text{lin}}(\ell) = C(C_{G'}(\ell))$. The series $\{s_n\}$ is linear when $C_{\text{obs}}(\ell) = C_{\text{lin}}(\ell)$ and is not linear otherwise. In this way, to quantify the nonlinearity of $\{s_n\}$ we can define the GLC non-linearity index as

$$GLC \equiv \sum_{\ell=1}^{\ell_m} |C_{\text{obs}}(\ell) - C_{\text{lin}}(\ell)| \quad (16)$$

Actually, this GLC value measures the deviation of the observed dependence from that of a Gaussian process. Note that if the Gaussian process is transformed to a different marginal distribution (the one of the observed time series) using an invertible transformation, the dependence of the values is not modified since the transformation preserves the ordering of the values in the time series. In this case, GLC will be very close to zero and then large GLC values will indicate non-linearity. However, if the observed time series comes from a non-invertible transformation of a Gaussian process, GLC will be larger than zero despite the linear underlying dynamics, so that this situation could be considered as a false positive. Nevertheless, for non-invertible transformations the dependence of the time series will be different from that of a Gaussian process, since the transformation will change the ordering of

the values. In this sense, we could say that the observed time series presents a higher complexity than a Gaussian process, and will be reflected also in the GLC value.

In summary, GLC measures any deviation from the dependence in a Gaussian process, the paradigm of the simplest correlated process. According to this, GLC could be considered as a complexity measure, although strictly speaking GLC measures nonlinearity only when the signal is originated with an invertible transformation. This approach is similar to that used in the copula theory⁴³ when the dependence is studied avoiding the effects of the particular marginal distribution.

III. DETECTION AND QUANTIFICATION OF NONLINEARITY

A. Detection of Nonlinear Dynamics

For each individual analyzed HRV time series, we first assess the presence of nonlinear dynamics exploiting the method of surrogate data⁶¹. This approach is based on: (i) a null hypothesis to be rejected; (ii) a surrogate data set constructed in accordance with the null hypothesis; (iii) a discriminating statistic that has to be calculated on original and surrogate series; and (iv) a statistical test allowing to reject or confirm the null hypothesis.

The null hypothesis set in our case is that the investigated time series is a realization of a Gaussian stochastic process (fully described by linear temporal autocorrelations), eventually measured through a static and possibly nonlinear transformation distorting the Gaussian distribution.

The surrogate time series were generated in order to preserve the linear autocorrelation structure as well as the marginal distribution of the original time series. This was achieved through the iteratively refined amplitude adjusted Fourier Transform (IAAFT) method⁵⁹. The method is an improvement of the Fourier transform (FT) method⁶¹, which generates surrogate time series by computing the FT of the original series, substituting the Fourier phases with random numbers uniformly distributed between 0 and 2π , and finally performing the inverse FT. Since the FT method distorts the amplitude distribution of the original process when such a distribution is not Gaussian, the IAAFT method is followed implementing an iterative procedure that alternatively constrains the surrogate series to have the same power spectrum (by replacing the squared Fourier amplitudes of the candidate surrogate series with those of the original series) and to have the same amplitude distribution (by a rank ordering procedure) of the original series.

As discriminating statistic we employ each of the three nonlinear indexes presented in Sect. (II), i.e., the normalized complexity index (NCI) based on local Sample Entropy, the regularity index based on information storage (IS), and the nonlinear index based on Gaussian Linear Contrast (GLC).

As statistical test, we perform a nonparametric test based on percentiles. The test compares the selected nonlinear index, here denoted generically as NI , when calculated on the original time series (NI_o) and when calculated on n_s surrogate

time series ($n_s = 100$ in this work) generated under the null hypothesis. Specifically, NI_o was compared with a threshold for significance NI_α extracted from the empirical distribution of NI over the surrogates setting a prescribed confidence level α ($\alpha = 0.05$ in this work). In the case of the NCI index measuring the complexity of a time series, the index is expected to decrease in the presence of nonlinear dynamics compared to linear time series; therefore, NI_α was set at the $100 \cdot \alpha$ -percentile of the distribution of NI over the surrogates and the null hypothesis was rejected if $NI_o < NI_\alpha$. In the case of the IS and GLC indexes measuring the regularity of a time series or the amount of nonlinear correlations, the indexes are expected to increase in the presence of nonlinear dynamics compared to linear time series; therefore, NI_α was set at the $100 \cdot (1 - \alpha)$ -percentile of the distribution of NI over the surrogates and the null hypothesis was rejected if $NI_o > NI_\alpha$. In either case, rejection of the null hypothesis allows to determine, individually for each subject, if the original value of the considered measure stands outside the 95% confidence interval of the surrogate distribution, thus detecting the presence of nonlinear dynamics.

B. Quantification of Nonlinear Dynamics

Surrogate time series were also exploited to quantify the ‘extent’ of nonlinearity in each investigated HRV recording, intended as the extent to which the original value of a measure deviates from its average value obtained from linear time series sharing the autocorrelation and amplitude distribution of the original series. This was performed comparing the index NI_o computed on the original, possibly nonlinear time series, with the median NI_m of its values computed on the set of surrogate time series. The difference with the median, defined as $\Delta NI = NI_m - NI_o$ in the case of the complexity index (i.e., when $NI = NCI$), and defined as $\Delta NI = NI_o - NI_m$ in the case of the two other indexes (i.e., when $NI = IS$ or when $NI = GLC$), was taken as a measure of the amount of nonlinearity in the observed time series. **Moreover an alternative measure accounting for the spread of the distribution of the nonlinearity measure was defined by normalizing the difference with the median to the standard deviation of the distribution of the measure over the surrogates (i.e., computing the ratio $\Delta NI / SD(NI)$). Results for this standardized difference are reported in the Suppl. material.**

IV. PATIENTS, EXPERIMENTAL PROTOCOL AND DATA ANALYSIS

The time series analyzed in this study belong to an historical database collected to analyze the effects of aging and myocardial infarction on cardiovascular interactions⁴⁵. The database consists on heart rate variability measured in a group of 35 post-acute myocardial infarction (AMI, 4 female, 58.5 ± 10.2 years old) patients examined about 10 days after AMI, and in two control groups formed by 12 old healthy subjects (Old, 9 female, 63.1 ± 8.3 years) and by 19 young healthy

subjects (*Young*, 9 female, 25.0 ± 2.6 years). The *Old* subjects were matched in age with the post-AMI patients, and both groups were significantly older than the young subjects, (two sample t-test on age distributions: $p = 0.17$ *Old vs. AMI*; $p < 0.00001$ *Old vs. Young*, *AMI vs. Young*). Eight out of 35 post-AMI patients were initially under beta-blocker therapy, but they discontinued the treatment two half-lives before the recording session. Control subjects were normotensive and free from any known disease based on anamnesis and physical examination at the time of the study.

After a period of 15 min for subject stabilization, the electrocardiogram (Siemens Mingograph, hardware bandpass filter 0.3-1000 Hz, lead II ECG) was recorded for 10 min in the supine rest position, followed by 10 min of passive head-up tilt at 60 degrees. All ECG signals were digitized with a 12 bit resolution and 1-KHz sampling rate. After detecting the QRS complex on the ECG and locating the R apex through template matching, heart period variability was measured on a beat-to-beat basis calculating the sequence of the time intervals occurring between pairs of consecutive R peaks (RR intervals). The series were then cleaned up from artifacts, windowed to $N = 300$ points for each condition (*rest*, *tilt*), and detrended by a high-pass filter to fulfill stationarity criteria^{44,66}. Normalized time series were eventually obtained by subtracting the mean values and dividing by the standard deviation.

For each subject and condition, analysis of nonlinearity was performed using the three methods described in Sect. II and performing the tests described in Sect. III. NCI and IS indexes were computed using standard values for the free parameters of entropy estimators applied to short time series^{48,58}, namely using $m = 2$ values to approximate the past history of the process, setting a tolerance $r = 0.2\sigma$ to define similarity in Sample Entropy analysis (where σ is the standard deviation of the series equal to 1 after normalization), and employing $k = 10$ neighbors in the distance-based entropy estimations. Distances between patterns were obtained using the Euclidean norm in the kernel estimator used to compute NCI⁵¹, and the maximum norm in the nearest-neighbor estimator used in IS⁶⁶. Moreover, to investigate the dependence on the type of entropy estimator and on the embedding dimension, the IS measure was computed also using the kernel estimator and varying the dimension in the range $m = (2, 3, 4)$. In the computation of the GLC index, taking into account the short size of the time series ($N = 300$) and to align with the other measures, we choose $\ell_{\max} = m = 2$ to limit spurious results induced by the fact that the autocorrelation function tends to reach quickly the noise level.

For each assigned index (NCI, IS, GLC), the statistical significance of its changes across groups (*Young*, *Old*, *AMI*) and conditions (*rest*, *tilt*) was assessed by the two-way ANOVA, **introducing the gender of the subjects as a categorical confounder variable in the test design. The assumption of Gaussianity was checked using the Jarque-Bera normality test, suggesting Gaussianity in the large majority of the distributions tested (15 out of 18 tests).** Post-hoc tests were performed using the two sample t-test with unequal variances to detect pairwise differences between groups (limited to the comparisons

Young vs. Old and *Old vs. AMI*, with Bonferroni correction for this double comparison), and using the paired Student t-test to detect pairwise differences between the two conditions (*rest vs. tilt*). We computed also the percentage of subjects belonging to each group for which the null hypothesis of linear Gaussian dynamics was rejected in the two conditions; then, statistically significant variations between two groups in a given condition were assessed using the chi-square test for proportions, while significant variations between conditions for a given group were assessed using the McNemar test for paired proportions.

V. RESULTS

Figure 1 reports an illustrative example of the application of the three considered nonlinear dynamic measures to individual HRV time series measured from one *Young* subject, one *Old* subject, and one *AMI* patient in the two analyzed experimental conditions (*rest*, *tilt*). In each panel and for each of the two conditions, the individual value of the considered measure is plotted with a symbol and the distribution of values obtained from the same measure over 100 surrogate time series is represented with an error bar (median and 5th – 95th percentiles); significant nonlinear dynamics are detected when the original value lies outside the surrogate distribution, while the extent of the deviation from can be inferred comparing the original value and the surrogate median. Considering the two entropy measures, opposite response to the change in condition are observed consistently for the three cases, with lower values of NCI and higher values of IS measured during *tilt* compared to *rest*. On the contrary, moving from *rest* to *tilt* the nonlinear dynamic measure based on GLC decreases slightly for the *Young* subject (circles), decreased more consistently for the *Old* subject (squares), and increases for the *AMI* patient (triangles). Moreover, the comparison between the original value of a measure and its distribution on the surrogate time series reveals the different ability to detect nonlinear dynamics of the different measures. In particular, in both the experimental conditions nonlinear dynamics are detected only by the Information storage in the *Young* subject (Fig. 1a) and only by the Gaussian Linear Contrast method in the *AMI* patient (Fig. 1c), while all measures detect the presence of nonlinear dynamics in the *Old* subject (Fig. 1b, NCI and IS in both conditions and GLC only at *rest*).

Most of the trends observed for the representative subjects described above are reflected at the population level, as reported in Figure 2 showing the distributions across subjects and conditions of the three nonlinear dynamic measures. The indexes based on conditional entropy and mutual information display opposite trends in response to the change of posture: the transition from *rest* to *tilt* is associated with a statistically significant decrease of the complexity index (NCI, Fig. 2a) in both *Young* ($p < 0.0005$) and *AMI* ($p = 0.025$) and a statistically significant increase of the information storage (IS, Fig. 2b; $p < 0.0005$) in both *Young* ($p < 0.0005$) and *AMI* ($p < 0.0005$), while no significant changes are detected for both measures in the *Old* group. Moreover, during *tilt* NCI

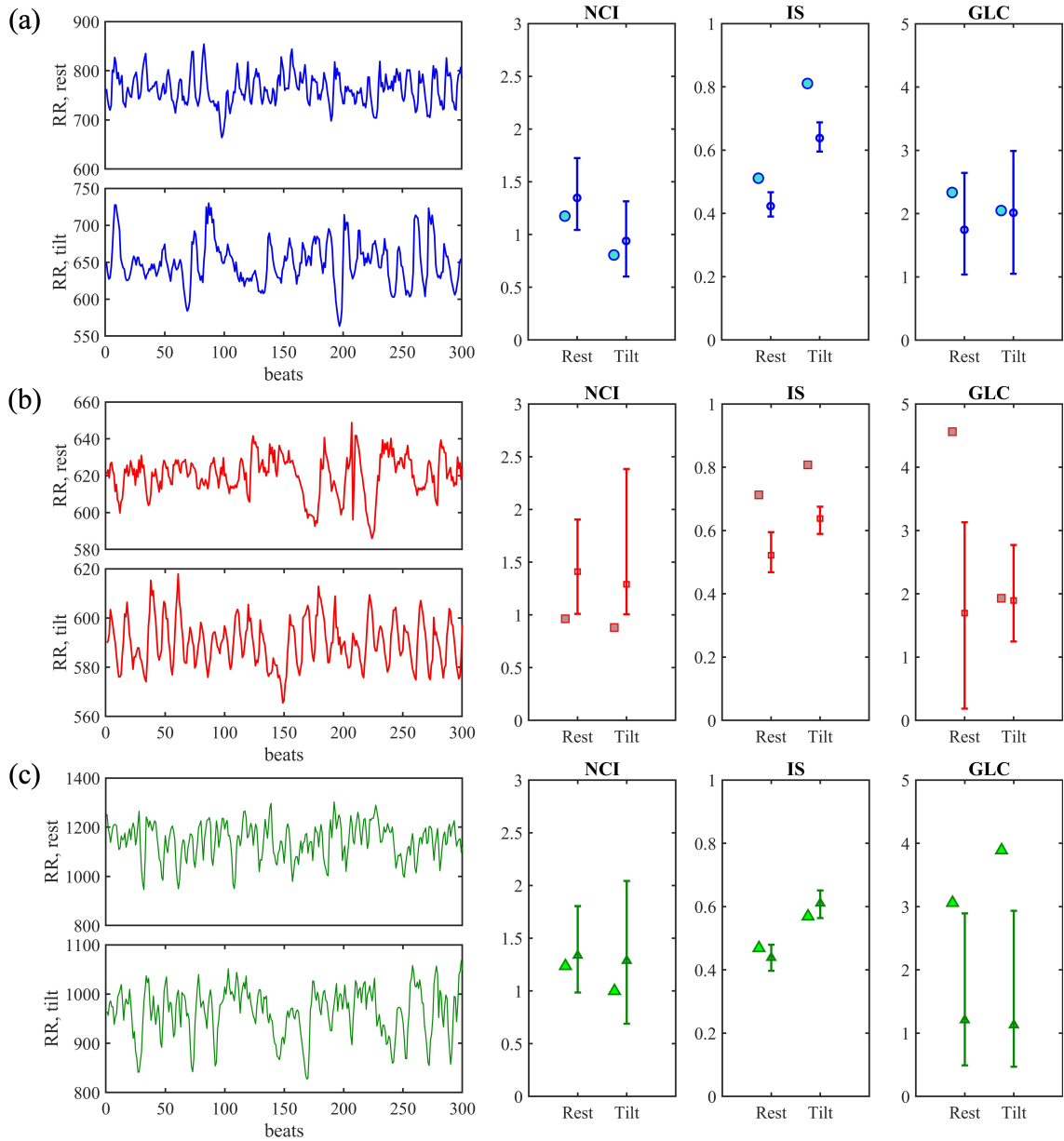


FIG. 1: Computation of nonlinear dynamic measures on heart period time series measured for a representative young subject (a, blue), old subject (b, red) and post AMI patient (c, green). For each subject, the time series of the RR interval measured in the two experimental conditions are reported on the left (above: *rest*; below: *tilt*), and the values of the nonlinear dynamic measures obtained with the three considered methods (NCI: Normalized Complexity Index; IS: Information Storage; GLC: Gaussian Linear Contrast) are reported on the right (markers: original values; error bars: 5^{th} , 50^{th} , 95^{th} percentiles of the distribution over 100 surrogates).

is significantly higher (Fig. 2a; $p < 0.0001$), and IS is significantly lower (Fig. 2b; $p < 0.0001$), in *Old* compared to *Young*. As to the GLC measure, it changes with the experimental condition in different ways for the different groups (Fig 2c): moving from *rest* to *tilt* the measure decreases significantly in the *Young* subjects ($p = 0.041$), does not change significantly in the *Old* subjects, and increases significantly in the *AMI* patients ($p = 0.023$).

Fig. 3 depicts the results of the analysis performed consid-

ering the deviation of each nonlinear dynamic measure from its median level assessed on linear Gaussian surrogates. We find that all three measures decrease significantly, in *Young* healthy subjects, with the transition from *rest* to *tilt* (ΔNCI : $p = 0.025$; ΔIS : $p = 0.024$; ΔGLC : $p = 0.01$), while no significant changes are observed for *Old* subjects and *AMI* patients (Fig. 3a,b). **It should be remarked that the assessment of the statistical significances reported for these cases may be influenced by deviations from normality, as the Jarque-Bera**

test did not reject the null hypothesis of joint Gaussianity for the rest and tilt distributions of the ΔIS and ΔGLC indexes evaluated in the *Young* subjects.

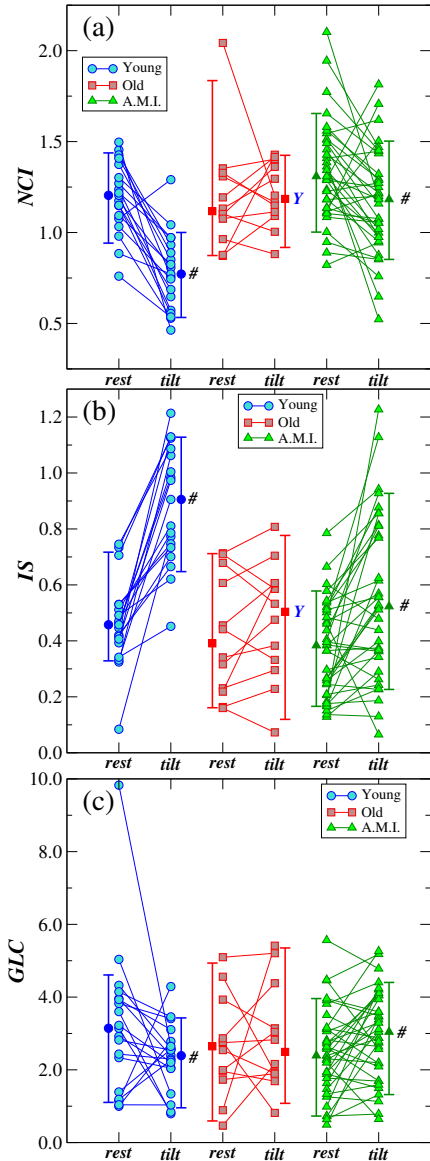


FIG. 2: Distribution of the three nonlinear dynamics measures (a, Normalized Complexity Index; b, Information Storage; c, Gaussian Linear Contrast) over the HRV time series of Young subjects (blue circles), Old subjects (red squares) and post AMI patients (green triangles) in the *rest* and *tilt* conditions, depicted as individual values (markers) and 5^{th} , 50^{th} , 95^{th} percentiles of the distributions across subjects (error bars). Symbols denote statistically significant changes between conditions ($\#$, *rest* vs. *tilt*) or groups (Y , *Old* vs. *Young*); see text for details.

Fig. 4 reports the the relevance of nonlinear dynamics in each group and experimental condition, measured as the percentage of subjects for which the value of the considered nonlinear dynamic measure computed for the original RR series

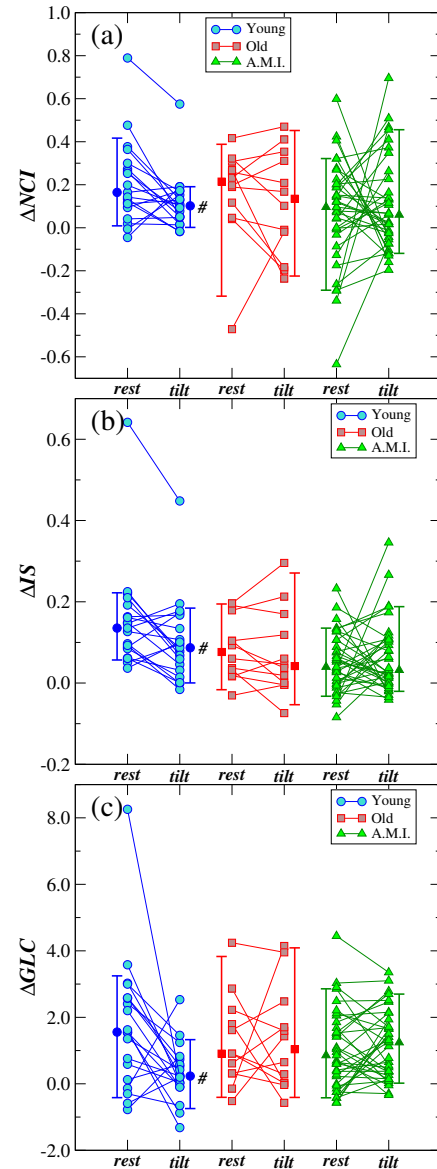


FIG. 3: Distribution of the difference with the median over surrogates of the three nonlinear dynamics measures (a, Normalized Complexity Index; b, Information Storage; c, Gaussian Linear Contrast) over the HRV time series of Young subjects (blue circles), Old subjects (red squares) and post AMI patients (green triangles) in the *rest* and *tilt* conditions, depicted as individual values (markers) and 5^{th} , 50^{th} , 95^{th} percentiles of the distributions across subjects (error bars). The symbol $\#$ denotes statistically significant changes between conditions (*rest* vs. *tilt*); see text for details.

is deemed (with 5 % significance) as not drawn from the distribution of the index derived from the surrogate RR series. The conditional entropy measure is associated with nonlinear dynamics in less than half of the subjects in each group, as the NCI index is found below the 5^{th} percentile of its surrogate distribution in $\sim 35\%$ of *Young* subjects, $\sim 45\%$ of *Old* subjects, and $\sim 25\%$ of *AMI* patients (with no substantial

differences between conditions, Fig. 4a). The mutual information measure detects a considerably higher percentage of nonlinear dynamics, as the IS index is found above the 95th percentile of its surrogate distribution in more than half of the subjects in all groups and conditions (Fig. 3b). In the *Young* group, the IS index is larger than the significance threshold in 95% of the subjects at rest and in $\sim 70\%$ of the subjects during tilt; in the *Old* and *AMI* groups the index is significantly lower during both conditions (Fig. 4b). The different detection rate of nonlinear dynamics exhibited by the NCI and IS measures is further investigated in the Supplementary material of this article implementing a kernel-based estimation of IS. The Gaussian Linear contrast approach detects nonlinear dynamics in $\approx 30 - 60\%$ of subjects in all groups and conditions (Fig. 4c). Moving from *rest* to *tilt*, the number of subjects with nonlinear dynamics detected by the GLC measure decreases in *Young*, while it increases in *Old* and *AMI*.

VI. DISCUSSION

The purpose of this study was to perform a comparative investigation of the aptitude of three recently proposed nonlinear dynamic measures (i.e., Normalized Complexity Index (NCI) recently defined as a refinement of the well known Sample Entropy measure⁵¹, Information Storage (IS) implemented through nearest neighbor estimation with bias compensation¹⁵, and Gaussian Linear Contrast (GLC)¹⁰) to quantify the presence and the extent of nonlinear dynamics in short-term recordings of HRV obtained under different physio-pathological states. In a time series observed as a realization of a stochastic process, nonlinear dynamics are typically described as nonlinear correlations between time-lagged variables taken from the process⁶¹. In our analysis, these correlations are detected directly in terms of mutual information between the present and the past samples of the process by the IS index, inversely in terms of conditional entropy of the present sample given the past by the NCI index, or in terms of deviation of the estimated correlation from the value that would be expected in case of linear correlations by the GLC index. Our results document that differences in the detection and quantification of nonlinearity emerge among the three measures, suggesting that a given nonlinear dynamic measure may be more or less sensitive to the detection of specific types of nonlinear dynamics depending on the properties of the measure and on the estimator adopted. Therefore, while some of the results are robust and suggest that distinct nonlinear dynamic structures may underlie the generation of HRV depending on the physio-pathological condition under analysis, physiological interpretations should be provided with cautiousness.

A first result is the opposite variation exhibited by the indexes of conditional entropy and information storage when moving from rest to tilt or while comparing two groups (Fig. 2a,b). This result, which is found consistently also varying some analysis parameters such as the type of entropy estimator and the embedding dimension (see Suppl. material), can be explained considering that NCI and IS are related to each

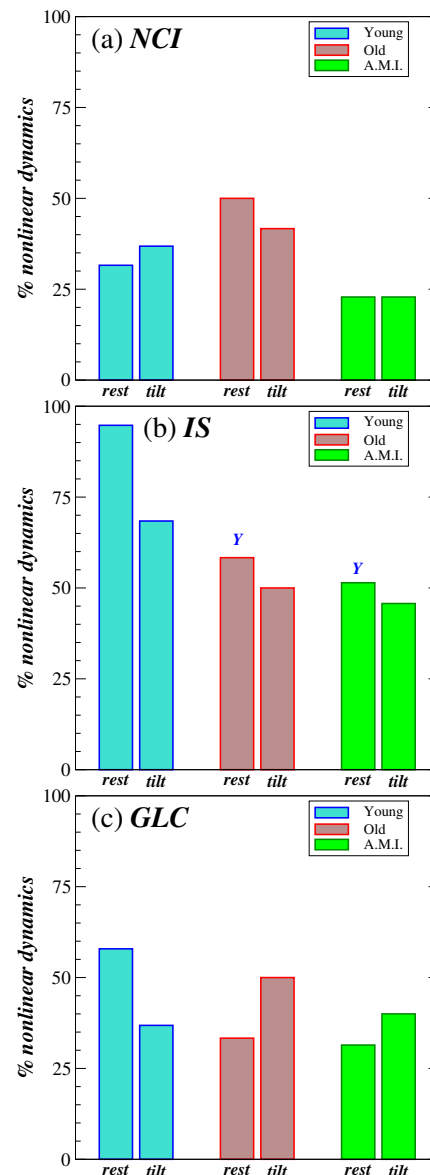


FIG. 4: Percentage of significant nonlinear dynamics obtained counting the subjects for which each nonlinear dynamic measure was significantly different for the original heart period than for the surrogate time series. Results are shown for each of the three proposed methods (a, Normalized Complexity Index; b, Information Storage; c, Gaussian Linear Contrast) applied to Young subjects (blue), Old subjects (red) and post AMI patients (green) in the *rest* and *tilt* conditions. The symbol *Y* denotes statistically significant changes between groups (*Young* vs. *Old*); see text for details.

other as they reflect respectively the unpredictability and the predictability of the dynamics⁶⁶. The lower NCI and higher IS measured in response to tilt indicate higher predictability of HRV, likely associated to the activation of the sympathetic nervous system induced by the postural challenge^{16,56}. Such an activation seems less important in the old and post-AMI

groups compared with the young subjects, as documented by the smaller variation of the indexes (though still statistically significant in AMI) and by the higher NCI/lower IS seen during tilt in Old and AMI compared to Young. Confirming previous studies^{39,45}, these results suggest that aging and myocardial infarction are associated with higher sympathetic tone and reduced capability to cope with the postural challenge with further sympathetic activation.

On the other hand, the trends displayed by the GLC measure (Fig. 2c) are in agreement with those of the conditional entropy in the young subjects (both GLC and NCI decrease with tilt), and with those of the information storage in the AMI patients (both GLC and IS increase with tilt). The different behavior of the GLC index can be explained by considering that this index reflects the extent to which the correlations of the time series deviate from those expected in the linear Gaussian case¹⁰, and thus it is not dependent on the extent of linear correlations within the observed time series. As such, GLC should be interpreted as a direct measure of nonlinearity rather than as a regularity index. This is confirmed by the consistent changes between conditions displayed by the absolute values of GLC and by the difference between the index and the median value of its surrogate counterparts (Fig. 2c vs. Fig. 3c). On the contrary, IS is a regularity measure which accounts for both linear and nonlinear correlations, and its increase with tilt is mainly driven by the enhancement of linear HRV correlations. In fact, when the effects of linear correlations are removed by subtracting the median on the surrogates the behavior of IS becomes more similar to that of GLC (Fig. 3b,c). This similar behavior is observed consistently for all three measures when nonlinear HRV dynamics were assessed computing the deviation of each measure from the median level of its surrogate distribution (Fig. 3a,b,c). The difference with the surrogate median reveals indeed that, in the young healthy subjects, the transition from rest to tilt is associated with a decreased degree of nonlinearity, and this result is in agreement with the observation that nonlinear dynamics are reduced in the presence of an increased sympathetic activity^{53,57}. Another peculiar result is the increased contribution of nonlinear dynamics to HRV measured during tilt in the post-AMI patients (Fig. 2c). This finding is novel and unexpected, and may reveal that a distinct type of nonlinearity takes place when the orthostatic stress is delivered in the presence of higher sympathetic tone.

In spite of the similar trends observed for the absolute values and for the deviation from the surrogate median value of NCI and IS (Fig. 3a,b), the two information measures exhibit different percentage of significant nonlinearity in the various conditions (Fig. 4a,b). In agreement with previous studies assessing complexity through prediction measures^{53,57}, the amount of nonlinear dynamics detected by the complexity measure based on local sample entropy was small in the young healthy subject at rest, and did not change significantly with the sympathetic activation induced by tilt or related to age and pathology (Fig. 4a). On the contrary, using a regularity measure based on information storage nonlinear dynamics were found consistently in the young subjects at rest, and their incidence decreased significantly with postural stress and in the

old and post-AMI groups (Fig. 4b). This finding may reflect the fact that the spontaneous cardiovascular regulation occurs through a variety of nonlinear mechanisms (e.g., saturation of receptors, effects of the respiratory centers at the brain stem level, interaction between sympathetic and parasympathetic nervous systems, etc.) in resting conditions³¹, and the rise of a specific oscillatory component (i.e., the low frequency one related to sympathetic activation) tends to simplify the dynamics reducing nonlinear components. The reduction of nonlinear dynamics with the tilt-induced sympathetic activation is confirmed (though to a lower extent) by the test using the GLC measure (Fig. 4c). The same test however indicates a tendency to increase the rate of detection of nonlinear dynamics with tilt in the old subjects and AMI patients. This could suggest that mechanisms more complex than a pure sympathetic activation are triggered by the orthostatic stress delivered in the elderly and pathological states⁴⁰.

However, more methodological factors might be responsible for the disparity of the conclusions drawn by the exploited markers. In a previous study⁵², different conclusions about HRV nonlinear dynamics were drawn using different nonlinearity measures (based on nonlinear prediction and time irreversibility) in fetal HRV recordings as well as in adults during graded head-up tilt. In particular, the different responses to tilt documented by Porta et. al⁵² using nonlinear prediction and time irreversibility are comparable to those observed here using the IS and GLC indexes. The different rates of detection of nonlinearity were explained by Porta et al.⁵² in terms of the different time scales spanned by the measures employed. However, this interpretation should not hold in our case since all measures work in the same low dimensional embedding space ($m = 2$ in this study). **The choice $m = 2$, which corresponds to a reconstructed state space defining vectors (S_n, S_{n-1}) , reflects a typical setting in short-term HRV analysis^{16,51,56,66}.** While this choice prevents the detection of higher dimensional dynamics which may occur in HRV, in short-term analysis it is recommended to keep reliability of the estimated entropy measures in the presence of short data sequences⁴⁷. Here, we observed that the adopted entropy measures bring to results which are robust against changes of this parameter in the range $m = (2, 3, 4)$ (see Suppl. material).

A difference between the information approach and the Gaussian Linear contrast method is that NCI and IS are obtained aggregating all time lags in the computation of the measure, while GLC results from analyses performed individually for each lag and then aggregated in the final measure. In addition, the performance of GLC might be affected by the comparison with surrogates that might have amplified eventual residual departures from gaussianity present in the surrogate data due to finite size effects. Moreover, the dependence of the cell size on the parameters set for the analysis (i.e., respectively, tolerance and number of nearest neighbors) might also have played a role. In order to better elucidate the nature of the observed differences and the capability of the various measures to detect different types of nonlinear dynamics, future studies should consider extension of these measures where longer temporal scales can be explored (e.g., analyzing longer stationary recordings and/or employing methods

for dimensionality reduction¹⁶), and deviations of the estimator specific parameters from their nominal typical value are investigated (e.g., for the information measures, the parameter setting the size of the cell used in the multidimensional space to estimate probabilities^{15,58}).

As to the differences observed between NCI and IS, our additional analysis comparing kernel and nearest neighbor estimates of the IS index (see Suppl. material) suggests that the type of estimator can be crucial in the detection of nonlinear dynamics. Indeed, when IS is computed via kernel estimates rather than via nearest neighbors, the percentage of significant nonlinear dynamics decreases, becoming comparable to that detected by the NCI index which is also grounded on kernel-based entropy estimation. This suggests that different coarse graining approaches using, respectively, equal versus different cell size⁵⁷, may result in a different sensitivity as regards the detection of nonlinear dynamics. **In particular, the nearest neighbor estimator turns out to be a better estimator for the information measures, as it shows higher sensitivity to the detection of nonlinear dynamics.** Another methodological aspect that can explain the difference is the fact that nearest neighbor estimation is accompanied by a distance-projection approach such that the cell size actually varies only while estimating entropies in the highest dimension, while it is kept constant (while allowing the number of neighbors to vary) in the lower dimensional spaces³³. Thus, we suggest that the bias of entropies computed in spaces of different dimension plays a role in the detection of nonlinear dynamics. This aspect should be investigated in more methodological studies involving simulations of linear and nonlinear dynamic systems.

Finally, we mention as possible limitations of the present study some characteristics of the analyzed database⁴⁵. The three analyzed groups are not homogeneous in the gender distribution (males are prevalent in *AMI*, females are prevalent in *Old*, and the gender is balanced in *Young*) and this may have an impact on the results as it is known that gender has a substantial effect on heart rate variability^{63,67}. Moreover, residual effects of the treatment with betablockers may be present in the *AMI* patients and have an effect on the dynamics analyzed. The results reported in this study should be confirmed on other databases with balanced gender distribution and absent of fully discontinued treatment after myocardial infarction.

SUPPLEMENTARY MATERIAL

See supplementary material for additional analysis illustrating the dependence of the measure of Information Storage on the embedding dimension and on the type of estimator used to compute entropies.

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