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# Noise and determinism in cardiovascular dynamics

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#### Abstract

Signals derived from the human cardiovascular system are well known to exhibit highly complex, nearly periodic, oscillatory behaviour whose nature is something of an enigma. It has, for example, been variously described as chaotic, fractal, stochastic, and subject to 1/f fluctuations—and its true nature is still the subject of vigorous debate. We review and describe some recent experiments that illuminate the problem and discuss a combination of noise and almost periodic frequency modulation as a signature of the system dynamics.

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#### 1. Introduction

Physiological signals derived from humans are extraordinarily complex—faithfully reflecting their origin in what is arguably the most complicated mechanism ever to have existed. Because they must reflect ongoing processes that normally occur unseen, within the interior of the body, such signals repay close attention. In particular, they can be used to diagnose incipient pathophysiological conditions before symptoms become obvious. A well-known example is the electrocardiogramme (ECG) signal, representing the electrical activity of the heart. ECG measurements have been used for diagnostic purposes for almost a century. For the first several decades of such measurements, attention was focussed mainly on the detailed *shape* of the approximately periodic

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pulses seen in the signal. More recently, however, attention has shifted to the *separation* of the pulses because it has become apparent that the heart of a healthy human in repose does not beat at a constant rate. Rather, the cardiac frequency varies in time, a phenomenon known as heart-rate variability (HRV).

Gene Stanley and his group have probably done more than anyone to try to characterise and analyse cardiovascular signals, so it is a particular pleasure to have been invited to present a paper on this topic in Messina. Approaches to the problem introduced by different authors have included, for example, studies of: Fourier spectra [1]; chaotic behaviour [2,3]; wavelet spectra [4–6]; Karhunen–Loève decomposition [7]; scaling properties [8–13]; Lyapunov exponents [14]; multifractal properties [15,16]; correlation integrals [17]; 1/f spectra [18–20]; and synchronization properties [21–25]. The two extreme perceptions of HRV consider it to be deterministic in origin, e.g. sometimes resulting in deterministic chaos [2,3], or stochastic [26]. It is not immediately obvious which of these several diverse approaches is the most promising: the criteria to be met, presumably, are that the technique of analysis should (i) be able to deal successfully with time variations of characteristic frequencies and amplitudes, and that (ii) it should yield some insight into the physiological processes responsible for the HRV, so that (iii) their individual function—normal or pathological—can be evaluated separately for diagnostic purposes.

In this paper we focus on the results obtained through use of one of the techniques mentioned above, wavelet analysis [4,27], and we point out that it apparently meets all three criteria. In Section 2 we summarise the experimental information obtained by this method and, in Section 3 we show how it can be accounted for in terms of mutually interacting oscillatory processes each of which appears to relate to a distinct physiological mechanism. In Section 4 we report the present status of attempts to model the cardiovascular system as a set of coupled oscillators, showing that important features of the measured signals can be reproduced by the model. In Section 5 we draw conclusions and try to point the way forward.

## 2. Cardiovascular signals

Many different protocols have been used for acquiring cardiovascular data, one of the most thorough being that described in Ref. [27] where ECG, blood pressure, and blood flow rate are recorded simultaneously over a period of about 20 min. Interest centres on the bloodflow circulatory control mechanisms, and so any processes occurring on timescales longer than about 1 min (the average circulation period) are ignored. For healthy subjects in repose, the results are typically as in Fig. 1(a); Fig. 1(b) shows the averaged wavelet transform of the same data, but calculated over the full 20 min. A detailed discussion of such results is presented in Ref. [27], but it is immediately evident that

- There are (at least) five characteristic spectral peaks.
- Remarkably, the *same*, or almost the same, peaks appear in all the spectra, regardless of where or how the corresponding signals were recorded, though there are considerable differences in amplitude.
- All the peaks are broadened.

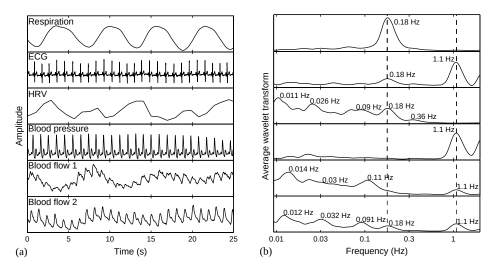


Fig. 1. Samples of cardiovascular signals (a) and time averages of their wavelet transforms (b). The positions of the peaks are almost the same for all signals, while the corresponding amplitudes may be considerably different. After Ref. [27].

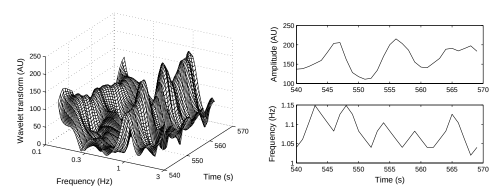


Fig. 2. A segment of the wavelet transform of the blood flow signal in the time-frequency plane (left). Peaks at the heart ( $\sim 1.1 \text{ Hz}$ ) and respiration rates ( $\sim 0.18 \text{ Hz}$ ) are visible. On the right are plotted the amplitude and frequency variations of the heartrate peak. After Ref. [5].

A clue to the origin of some of the broadening can be found by inspection of the full time-frequency wavelet spectrum, part of which is shown in Fig. 2. It can be seen that both the instantaneous amplitudes and central frequencies vary in time, giving rise to a considerable part of the broadening observed in the averaged spectra of Fig. 1(b).

What is the origin of this frequency and amplitude wandering? It appears that the instantaneous frequency of any given spectral peak oscillates. It does so at the frequency of the spectral peak next-lowest in frequency, with contributions coming from all of the other oscillatory processes too. HRV, for example, is a signal representing the variations of the cardiac frequency, and from Fig. 1(b) we can see (third spectrum

from the top) that its oscillations are also modulated by the processes with frequencies near 0.011, 0.026, 0.10, and 0.18 Hz (as well as its own second harmonic at 0.036 Hz). Such observations can be construed as evidence that the five oscillatory processes are mutually coupled. They seem to influence each other via couplings at least some of which are parametric, thus giving rise to the observed frequency modulation.

# 3. Interacting oscillatory processes

With varying degrees of confidence, all of the oscillatory processes can now be related to underlying physiological mechanisms. Their characteristic average frequencies vary slightly between individuals, and, in some cases, between different measurement locations, but in each case they lie within definite limits. The physiological origins of the peaks at 1 Hz (heart beat), and 0.2 Hz (respiration) are obvious. It is reasonably well established that the peak at 0.1 Hz is attributable to intrinsic myogenic activity of smooth muscles [28–30]. The 0.03 Hz peak is apparently connected to autonomous nervous control [31,32] (neurogenic), and there is strong evidence that the 0.01 Hz peak arises from metabolically related endothelial activity [33,34].

Thus wavelet analysis of signals from the cardiovascular system provides a noninvasive technique for acquiring information about these distinct physiological processes. Additional information, yielding deeper insight into the dynamics, can also be inferred from the *couplings* between the processes. These can manifest themselves either through mutual modulation (Fig. 2) or through synchronization between two or more of the processes. Which of these phenomena occurs in any given case probably depends partly on the strength of the coupling and partly on how close in frequency low harmonics of the processes would be to each other in the absence of coupling. In practice, measurements on healthy subjects show transient episodes of cardio-respiratory synchronization lasting typically a few tens of seconds. The average length of these episodes depends on the physiological state, and differs between, e.g. athletes, sedentary individuals, and patients with pathological conditions such as diabetes.

Synchronization effects are potentially of particular interest because of the information they carry about the couplings. But, with the exception of the cardio-respiratory coupling, they have not yet been closely studied. A major difficulty is that, unlike the heart and respiration, the lower-frequency processes do not give rise to separate signals that can be acquired noninvasively. In practice, therefore, it is necessary to infer the existence or absence of synchronization from univariate time series. This challenging problem is already being approached in three distinct ways: (i) filtration of the univariate data to create two "separate" signals that can then be tested for mutual synchronization [23] using established techniques such as synchrogrammes [21] or synchronization indices [35]; (ii) the use of angles-of-return-time maps [25,36]; and (iii) bispectral analysis [37]. In each case, it has been possible to derive information about the presence or absence of synchronization between the 0.1 Hz (myogenic) process and respiration or heartbeat. Problems of resolution—in time, frequency and phase—have so far made it impossible to obtain synchronization information for the 0.03 Hz (neurogenic) and 0.01 Hz (endothelial) processes.

Note that the three lower frequencies are derived in each case from spatially distributed processes. Within each of them, there must be significant global synchronization because, if this were not the case, low frequency oscillations would be undetectable in centrally measured quantities such as HRV (Fig. 1(b)) because they would all have averaged out.

To make further progress, it is necessary to develop a model to which the cardiovascular data can be fitted. One may then hope to characterise different physiological and pathophysiological conditions quantitatively in terms of the model parameters.

# 4. A coupled oscillator model of the cardiovascular system

Given the clear evidence of (a) well-defined spectral peaks (implying the presence of oscillatory processes) and (b) amplitude and frequency modulation, and synchronization effects (all indicating the existence of inter-oscillator interactions), it is natural to try to model the system with a set of oscillators [38] whose couplings [39] can be adjusted to try to reproduce the observed phenomena.

Little experimental information exists yet about either the nature of the couplings or the details of the oscillators. So we have used [38] the Poincaré oscillator

$$\dot{x}_i = -x_i q_i - \omega_i y_i + g_{x_i}(\mathbf{x}),$$

$$\dot{y}_i = -y_i q_i + \omega_i x_i + g_{y_i}(\mathbf{y}), \quad q_i = \alpha_i (\sqrt{x_i^2 + y_i^2} - a_i),$$
(1)

where,  $\mathbf{x}$ ,  $\mathbf{y}$  are vectors of the oscillator state variables,  $\alpha_i$ ,  $a_i$ , and  $\omega_i$  are constants and  $g_{yi}(\mathbf{y})$  and  $g_{xi}(\mathbf{x})$  are linear coupling vectors. Although to some extent this choice is arbitrary, (1) possesses properties of structural stability, robustness and symmetry consistent with physiological understanding and the analyses of measured time series.

Using a numerical simulation [39] of five coupled oscillators (1), with characteristic frequencies chosen to accord with measured data (Fig. 1), and amplitudes set initially to unity, we find that we can generate signals that to the eye seem in many respects to resemble those from the cardiovascular system. We were especially interested to establish whether frequency and amplitude modulation would occur with short episodes of synchronization at random intervals, as observed in the experiments. The investigations are still at an early stage, but we have already established: (a) that the observed depth of modulation requires parametric couplings (as expected); (b) that with appropriately chosen parameter values parametric modulation indeed gives rise to episodes of synchronization, but of course in a totally deterministic fashion with equal intervals between the episodes; and (c) with purely linear couplings, and added noise (random fluctuations), the synchronization episodes occur briefly and randomly just as observed in reality. An example showing the modelling of cardiorespiratory synchronization is shown in Fig. 3. It appears, therefore, that both linear and parametric couplings exist and that it is essential to take into account the influence of stochastic effects resulting from the (unmodelled) rest of the system.

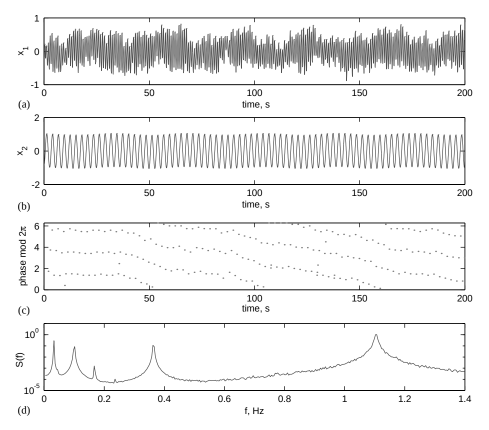


Fig. 3. Results of modelling with linear couplings, in the presence of fluctuations. (a) and (b) The time series showing the rhythmic activities of the cardiac and respiratory flow components. (c) The corresponding cardio-respiratory synchrogram. (d) Power spectrum of oscillation in the blood flow generated by the cardiac activity. After Ref. [39].

#### 5. Conclusions

Information derived from analysis of cardiovascular signals in the time-frequency and time-phase domains has led to a coupled oscillator model able to reproduce many features seen in the data. From the model, we conclude that cardiovascular signals have a strong deterministic element, but that random noise (i.e. external influences and all effects not explicitly considered) also plays a crucially important role. Our approach meets all three of the criteria mentioned in Section 1. In particular, it relates the underlying physiological processes to particular spectral peaks and thus allows them to be studied individually. Furthermore, it promises quantitative evaluation of the couplings between them. The latter feature is potentially of particular interest for diagnosis and treatment because it enables the function and health of the cardiovascular system as a whole to be evaluated.

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## References

- [1] S. Akselrod, D. Gordon, F.A. Ubel, D.C. Shannon, A.C. Barger, R.J. Cohen, Science 213 (1981) 220.
- [2] A. Babloyantz, A. Destexhe, Biol. Cybernet 58 (1988) 203.
- [3] C.S. Poon, C.K. Merrill, Nature 389 (1997) 492.
- [4] M. Bračič, A. Stefanovska, Bull. Math. Biol. 60 (1998) 919.
- [5] M. Bračič, P.V.E. McClintock, A. Stefanovska, Characteristic frequencies of the human blood distribution system, in: D.S. Broomhead, E.A. Luchinskaya, P.V.E. McClintock, T. Mullin (Eds.), Stochastic and Chaotic Dynamics in the Lakes, American Institute of Physics, Melville, New York, 2000, pp. 146–153.
- [6] M. Bračič-Lotrič, A. Stefanovska, D. Štajer, V. Urbančič-Rovan, Physiol. Meas. 21 (2000) 441-457.
- [7] M. Hožič, A. Stefanovska, Physica A 280 (2000) 587.
- [8] P.C. Ivanov, M.G. Rosenblum, C.K. Peng, J.E. Mietus, H.E. Stanley, A.L. Goldberger, Physica A 249 (1998) 587.
- [9] L.A.N. Amaral, A.L. Goldberger, P.C. Ivanov, H.E. Stanley, Phys. Rev. Lett. 81 (1998) 2388.
- [10] S. Havlin, S.V. Buldyrev, A.L. Goldberger, P.C. Ivanov, C.K. Peng, H.E. Stanley, Physica A 273 (1999) 46.
- [11] S. Havlin, L.A.N. Amaral, Y. Ashkenazy, A.L. Goldberger, P.C. Ivanov, C.K. Peng, H.E. Stanley, Physica A 274 (1999) 274.
- [12] Y. Ashkenazy, P.C. Ivanov, S. Havlin, C.K. Peng, A.L. Goldberger, H.E. Stanley, Phys. Rev. Lett. 86 (2001) 1900.
- [13] P. Bernaola-Galván, P.C. Ivanov, L.A.N. Amaral, H.E. Stanley, Phys. Rev. Lett. 87 (2001) 168105.
- [14] M. Bračič, A. Stefanovska, Bull. Math. Biol. 60 (1998) 417.
- [15] P.C. Ivanov, L.A.N. Amaral, A.L. Goldberger, S. Havlin, M.G. Rosenblum, Z.R. Struzik, H.E. Stanley, Nature 399 (1999) 461.
- [16] L.A.N. Amaral, P.C. Ivanov, N. Aoyagi, I. Hidaka, S. Tomono, A.L. Goldberger, H.E. Stanley, Y. Yamamoto, Phys. Rev. Lett. 86 (2001) 6026.
- [17] A. Stefanovska, P. Krošelj, Open Systems Inform. Dyn. 4 (1997) 457.
- [18] M. Kobayashi, T. Musha, IEEE Trans. Biomed. Eng. 29 (1982) 456.
- [19] C.K. Peng, J. Mietus, J.M. Hausdorff, S. Havlin, H.E. Stanley, A.R. Goldberger, Phys. Rev. Lett. 70 (1993) 1343.
- [20] P.C. Ivanov, L.A.N. Amaral, A.L. Goldberger, S. Havlin, M.G. Rosenblum, H.E. Stanley, Z.R. Struzik, Chaos 11 (2001) 641.
- [21] C. Schäfer, M.G. Rosenblum, J. Kurths, Nature 392 (1998) 239.
- [22] A. Stefanovska, H. Haken, P.V.E. McClintock, M. Hožič, F. Bajrović, S. Ribarič, Phys. Rev. Lett. 85 (2000) 4831.
- [23] A. Stefanovska, M. Hožič, Prog. of Theor. Phys. (Suppl.) 139 (2000) 270.
- [24] M. Bračič Lotrič, A. Stefanovska, Physica A 283 (2000) 451.
- [25] N.B. Janson, A.G. Balanov, V.S. Anishchenko, P.V.E. McClintock, Phys. Rev. Lett. 86 (2001) 1749.
- [26] P.C. Ivanov, L.A.N. Amaral, A.L. Goldberger, H.E. Stanley, Europhys. Lett. 43 (1998) 363.

- [27] A. Stefanovska, M. Bračič, Contemporary Phys. 40 (1999) 31.
- [28] P.C. Johnson, News Physiol. Sci. 6 (1991) 41.
- [29] S. Bertuglia, A. Colantuoni, M. Intaglietta, Microcirc. Res. 48 (1994) 68.
- [30] H.D. Kvernmo, A. Stefanovska, M. Bračič, K.A. Kirkebøen, K. Kvernebo, Microvasc. Res. 56 (1998) 173.
- [31] J. Karstrup, J. Bühlow, N.A. Lassen, Int. J. Microcirc.: Clin. Exp. 8 (1989) 205.
- [32] T. Söderström, A. Stefanovska, M. Veber, H. Svenson, Am. J. Physiol., in press.
- [33] H.D. Kvernmo, A. Stefanovska, K.A. Kirkebøen, K. Kvernebo, Microvasc. Res. 57 (1999) 298.
- [34] A. Stefanovska, M. Bračič, H.D. Kvernmo, IEEE Trans. Biomed. Eng. 46 (1999) 1230.
- [35] P. Tass, M.G. Rosenblum, J. Weule, J. Kurths, A.S. Pikovsky, J. Volkmann, A. Schnitzler, H.-J. Freund, Phys. Rev. Lett. 81 (1998) 3291.
- [36] N.B. Janson, A.G. Balanov, V.S. Anishchenko, P.V.E. McClintock, Phys. Rev. E 65 (2002) 036211 and 036212.
- [37] J. Jamšek, A. Stefanovska, Bispectral analysis of cardiovascular oscillations, Phys. Rev. E., submitted for publication.
- [38] A. Stefanovska, M. Bračič Lotrič, S. Strle, H. Haken, Physiol. Meas. 22 (2001) 535.
- [39] A. Stefanovska, D.G. Luchinsky, P.V.E. McClintock, Physiol. Meas. 22 (2001) 551.