The 4/3 exponent is sharp. Consider the Hopf link of two tori in its natural geometrical position. Fill each torus with N loops parallel to the centre curve, each loop a strand of radius 1 (Fig. 1a). Then with any tight packing of the loops, the minor radii of the tori is of the order of \( L \). The conformations inside a sphere of radius \( L N \), so the total rope length is about \( N^{3/2} \). Each loop is linked with N loops in the perpendicular torus, so the crossing number is about \( N^2 \). Therefore the rope length is of the order of \( C(K)^{3/2} \). Because \( 11L(K)^{3/2} \geq 4\pi A(K) \geq C(K) \), this example has \( A \) in the order of \( L(K)^{3/2} \).

The minimum rope length for a knot is bounded by \( 3C(K)^{2} \). This can be seen by arranging the knot so that the crossings are evenly spaced along a line (Fig. 1d). For the simpler knot types, \( L(K) \), \( S(K) \) and \( A \) in minimized conformations all ‘appear’ to be linearly related. An explanation is that the simpler conformations are ‘planar’: from most perspectives a unit arc of the knot crosses only a few other unit arcs.

As complexity increases, there are many families of knots and links with three-dimensional growth, exhibiting the 4/3 power law. Families with single-dimensional growth, e.g. (Fig. 1b,c) have a linear relationship among the measures. With planar growth, we expect \( A \) to be linear with \( C(K) \) and \( S(K) \) to be of the order of \( L(K) \log L(K) \).

We propose that the rope length required (Fig. 1e–i) to tie an \( N \)-crossing knot or link varies only between \( kN^{3/2} \) and \( kN \). Other investigators have also recently observed the 4/3 law in knots on the cubic lattice and in vector fields.

A good knot energy has only a finite number of knot types realized below any given energy level. Our theorem gives us this property for \( L(K) \) and \( S(K) \), proving that there is a finite number of knots that can be tied with a finite length of mathematical rope.

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Heartbeat synchronized with ventilation

It is widely accepted that cardiac and respiratory rhythms in humans are unsynchronised. However, a newly developed data analysis technique allows any interaction that does occur in even weakly coupled complex systems to be observed. Using this technique, we found long periods of hidden cardiorespiratory synchronization, lasting up to 20 minutes, during spontaneous breathing at rest.

Synchronization is a universal phenomenon that occurs when two or more non-linear oscillators are coupled. It is observed in many fields of science and is widely applied in engineering. The case of synchronization in periodic, or even noisy, oscillators is well understood. The notion of synchronization has often been used to analyse the interaction between physiological (sub)systems, but these studies have been restricted to almost periodic rhythms. No approach has been suggested to probe the weak interactions between such irregular and non-stationary oscillators as the human heart and respiratory system.

These two physiological systems are known to be coupled by several mechanisms, but apart from respiratory modulation of heart rate, first observed in 1847 and known as ‘respiratory sinus arrhythmia’ (RSA), no other effects have been reported. Moreover, in spite of some early communications, it has been concluded that “there is comparatively weak coupling between respiration and the cardiac rhythm, and the resulting rhythms are generally not phase locked”

We used the concept of phase synchronization of chaotic oscillators to develop a technique to analyse irregular non-stationary bivariate data. We analysed data obtained in non-invasive examinations of eight healthy volunteers (14–17-year-old, high-performance swimmers; four of them male and four female). While subjects lay at rest, electrocardiograms (ECGs) were recorded while respiratory
flow was simultaneously measured with a thermistor at the nose. Both signals were digitized with a 1,000-hertz sampling rate and 12-bit resolution. Each record lasted 30 minutes.

The resulting time series were irregular, strongly non-stationary and noisy. These characteristics made it inappropiate, in analyzing the mutual dependencies involved, to use even sliding versions of traditional spectral and correlations techniques, or direct computation of instantaneous phase differences. So instead of these techniques, we used a new kind of data presentation which we call a cardiorespiratory synchronogram (CRS), to detect different synchronous states and transitions between them.

We first used the Hilbert transform 9 to calculate the instantaneous phase \( \phi_t(t) \) of the respiratory signal. \( \phi_t(t) \) is defined on the real line (not restricted to the \([0,2\pi]\) interval). Next, we regarded the respiratory phase stroboscopically at times \( t_k \), where the R-peak in the \( k \)th heartbeat occurs and hence the phase of the heart rhythm increases by \( 2\pi \). In the simplest case of \( m:1 \) synchronization, there are \( n \) heartbeats in each respiratory cycle; these beats appear at \( n \) certain values of respiratory phases, which are constant over all cycles.

Plotting these relative phases \( \psi \) as a function of time shows \( n \) horizontal stripes. In the general case of \( mn \) synchronization, such a structure appears if we relate the phases of the heart beats to the beginning of \( m \) adjacent respiratory cycles, \( \psi(t_k) = (\phi_t(t_k) \mod 2\pi)/2\pi \); we have \( n \) horizontal stripes within \( m \) respiratory cycles.

This technique allows us to distinguish between different periods of synchronization, even for noisy and non-stationary data. For example, we observe 5:2 locking between the respiratory frequency \( \omega_R \) and the heart rate \( \omega_{HR} (3\omega_R=2\omega_{HR}) \) during a period of about 300 seconds, then after a transition period, a regime of 3:1 phase locking is found for about 20 minutes (Fig. 1). These two kinds of locking can be recognized using histograms (Fig. 1c) and the autocorrelation function of phases (Fig. 1d).

Our analysis showed cardiorespiratory synchronization of several locking ratios occurring in six out of eight subjects (Table 1). Subjects with the strongest synchronization had no remarkable RSA, whereas both subjects with the highest RSA exhibited no synchronization.

We conclude that phase locking of respiratory and the cardiac rhythms, and respiratory modulation of heart rate, are two competing aspects of cardiorespiratory interaction. From a physical viewpoint, synchronization and modulation are different phenomena and are related to different coupling. RSA generation is associated mainly with the baroreflex feedback loop and its respiratory phase-dependent information processing.

Perhaps cardiorespiratory synchronization is an expression of another kind of cardiorespiratory interaction, such as a central coupling between cardiovascular and respiratory neuronal activities.

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