Network Reconstruction from Random Phase Resetting

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(Received 16 December 2010; published 11 July 2011)

We propose a novel method of reconstructing the topology and interaction functions for a general oscillator network. An ensemble of initial phases and the corresponding instantaneous frequencies is constructed by repeating random phase resets of the system dynamics. The desired details of network structure are then revealed by appropriately averaging over the ensemble. The method is applicable for a wide class of networks with arbitrary emergent dynamics, including full synchrony.

DOI: 10.1103/PhysRevLett.107.034101 PACS numbers: 05.45.Xt, 87.19.lj

Complex networks of many interacting units found at all scales in nature are the subject of intense research in many scientific areas [1]. Among the central issues in this field are the exploration and development of methods for determining the architecture of a network based on the observable data. Knowing the network structure helps in understanding its collective behavior, and indicates ways to engineer networks with the desired properties. For instance, it has been realized that inferring the topology of gene regulatory networks is crucial for completing our knowledge about the inner workings of cells [2]. Many real networks display modular and community structure that is essential for their functioning [3] and can be extracted using a variety of methods [4], as done for yeast metabolic network [5]. Reconstruction techniques often rely on examining the time series of network dynamics that can reveal its interaction functions [6]. The network topology can be detected by studying the interchanges among its collective behaviors or investigating its response dynamics [7]. Structural properties can be determined from various time scales in the emergence of synchronization [8] or by employing specific control theory methods [9]. Recently proposed techniques involve noisy dynamical correlations between the nodes [10], and even tackle models with nonequilibrium dynamics [11].

However, existing reconstruction methods, that often use network models with single-node dynamics represented by different types of oscillators [12], typically require long time series of dynamical data or a certain level of complexity in the emergent dynamics [6,7]. Since synchronization destroys the initial node-related information, detecting network topology in such cases is extremely difficult. Some methods are applicable only to sparse or nondirected networks, often providing results with only a limited precision [10].

In this Letter, we propose a novel method of reconstructing the topology and interaction functions of a general oscillator network. Our idea relies on repeatedly reinitializing the network dynamics (e.g., by performing random phase resets), in order to produce an ensemble of the initial dynamical data. We design the quantities obtained by averaging this ensemble, whose values reveal the desired details of the network structure. Our method is applicable to any directed and weighted network, with general interaction functions and oscillator frequencies, and with arbitrary emergent dynamics, while avoiding the need for long time series.

In the context of phase resets, one is typically interested in the phase-resetting curves, which specify the system’s response to weak external perturbations [13]. They are investigated both experimentally [14] and theoretically [15,16], and known to contain properties relevant for determining network details such as clustering [17]. An algorithm for the estimation of neuron interaction and its stability based on phase resets has been proposed [14]. We here employ phase resetting somewhat differently, since our interest lies in the internal network interactions, rather than its response to stimuli. Contrary to [14], we use phase resets only as a natural way to reinitialize the dynamics of an oscillator network, without measuring the phase shifts occurring due to resetting.

Our model consists of $N$ oscillators (nodes), characterized by their phases $\varphi_i \in [0, 2\pi)$ and natural frequencies $\omega_i$. They are coupled pairwise, via general $2\pi$-periodic interaction functions $f_{ij}$ with zero mean:

$$\varphi_i = \omega_i + \sum_{j=1}^{N} f_{ij}(\varphi_j - \varphi_i).$$

Models of this type include the famous Kuramoto model and its generalizations, widely used in theoretical studies, as well as for describing specific experimental situations [8,12,14]. The functions $f_{ij}(\phi)$ are generally nonsymmetric with respect to exchange of indices, and thus fully define the dynamical network (order of indices determines the direction of interaction). The network adjacency matrix given as $A_{ij} = \text{sgn}[f_{ij}]$ specifies its topology. Dynamics starts from a set of initial phases (IP) which we denote as $\varphi = (\varphi_1, \ldots, \varphi_N)(t = 0)$, chosen from a distribution $p(\varphi) > 0$ normalized to $(2\pi)^N$. The method is based on two assumptions. (i) We are able to arbitrarily reinitialize the network dynamics $I$ times, by independently resetting
the corresponding interaction parameters \(\phi_i\) in each of the dynamical networks defined as

\[
S_{ij}[\phi] = (2\pi)^{-N} \int_{[0,2\pi]^N} d\phi g(\phi_i - \phi_j) \phi_j(\phi).
\]

Taking the functions \(f_{ij}\) in Eq. (1) to be generally given by the Fourier series \(f_{ij}(\phi) = \sum a_{ij}^{(n)} \sin(n\phi) + b_{ij}^{(n)} \cos(n\phi)\), we obtain the following expression for \(S_{ij}\):

\[
S_{ij} = (2\pi)^{-N} \sum_{k=1}^{N} \sum_{n=1}^{\infty} \int_{[0,2\pi]^N} d\phi g(\phi_i - \phi_j) \times \left[a_{ij}^{(n)} \sin(n(\phi_i - \phi_j)) + b_{ij}^{(n)} \cos(n(\phi_i - \phi_j))\right],
\]

which is independent of the frequencies \(\omega\). The integral over \(\phi\) vanishes unless \(i = k\). This implies that if \(A_{ij} = 0\), the corresponding \(S_{ij} = 0\), independently of the choice of \(g\). The nonzero entries of \(S_{ij}\) directly reveal the presence of network links. In addition, matrix \(S_{ij}\) detects the desired properties of the interaction functions for appropriately selected test function \(g\). In particular, using \(g(\phi) = 2e^{i\phi}\) we obtain the Fourier harmonics of \(f_{ij}\), which are the interaction parameters \(a_{ij}^{(n)}\) and \(b_{ij}^{(n)}\):

\[
S_{ij}[2e^{i\phi}] = b_{ij}^{(n)} + ia_{ij}^{(n)} = \frac{1}{\pi} \int_0^{2\pi} f_{ij}(\phi)e^{i\phi} d\phi.
\]

Any dynamical network described by Eq. (1) can be reconstructed by computing \(S_{ij}\) for adequate \(g\). Depending on the properties of \(f_{ij}\) that are to be examined, other choices of \(g\) are also possible. When dealing with the empirical interaction functions involving an unknown number of Fourier harmonics, a specifically designed \(g\) based on the experimental assumptions about \(f_{ij}\) might be useful. This result is largely independent of the frequencies \(\omega\), network’s directedness, distribution \(\rho\), and the network’s final dynamical state. However, a constant component in the case of \(f_{ij}\) with nonzero mean cannot be detected, since its presence is indistinguishable from the natural frequency \(\omega\).

To practically implement our method, we need to convert the integral from Eq. (2) into an average involving discrete nonuniformly distributed empirical data \(\{\phi_i\}_{i=1}^{N}\) and \(\{\phi_i\}_{i=1}^{N}\). To that end, we represent the function \(\phi_j(\phi)\) using the kernel smoother \(Q(\phi - \phi_i)\) [18] as

\[
\phi_j(\phi) = \sum_{i=1}^{N} Q(\phi - \phi_i) \phi_j(\phi_i).
\]

The denominator is just the empirical density \(\rho(\phi) = \sum_{i=1}^{N} Q(\phi - \phi_i)\) obtained via the kernel distribution estimate [18]. Since the integration over \(\phi\) already provides smoothing, we take \(Q(\phi - \phi_i) \rightarrow \delta(\phi - \phi_i)\) and replace Eq. (2) with a practical formula for \(S_{ij}\):

\[
S_{ij}[g] = \left\langle \phi_j g(\phi_i - \phi) \rho(\phi) \right\rangle = \frac{1}{\rho(\phi)} \frac{1}{I} \sum_{i=1}^{I} \phi_j(\phi_i) g(\phi_i - \phi),
\]

which is the average of empirical \(\phi_j g\) weighted by \(1/\rho(\phi)\).

The most trivial way to obtain the ensemble \(\{\phi_i\}_{i=1}^{N}\) would be to pick the values from a fixed distribution \(\rho(\phi)\). Instead, we seek to mimic an experimentally feasible situation by performing \(I\) random phase resets of the network dynamics, separated by the time interval \(\tau\). Mathematically, this amounts to adding the term \(\sum_{i=1}^{I} K_{ij} \sin(\phi_i + \alpha_{ij}) \delta(t - l\tau)\) to the right-hand side of Eq. (1) [16]. For each reset \(l\) and each oscillator \(i\), we independently pick the kicking strength \(K_{ij}\) from a zero mean Gaussian distribution with standard deviation \(K = 1\) and the phase shift \(\alpha_{ij}\) uniformly from \([0,2\pi]\). The ensemble is constructed by storing the phase values immediately after resets. The resulting artificially created ensemble has little in common with the natural distribution of phases, and can be considered as approximately independent. This is expressed by separability of \(\rho(\phi)\) into a product of \(N\) one-dimensional distributions \(\rho_i(\phi_i)\):

\[
\rho(\phi) = \prod_{i=1}^{N} \rho_i(\phi_i),
\]

each of which we determine from generated data using the kernel estimation method [19]. After each reset, the ensemble of \(\phi\) is computed using a small time interval. The phase value prior to reset is of no importance, since our interest is solely in modeling a realistic way to create the ensemble \(\phi\). This procedure resembles recent experiments where the random stimulation was used to induce transient desynchronization [20] in epileptic neuronal populations [21]. In these experiments, however, it is still not verified that the random independent phase resetting is achieved.

We now illustrate our theoretical findings through numerical simulations on simple network examples, computing the reconstruction index \(S_{ij}\) as described above. Consider a simple network with \(N = 4\) oscillators shown in Fig. 1. We pick the natural frequencies at random from \(\omega_i \in [-1, 1]\). The interaction functions \(f_{ij}\) are defined for linked node pairs by randomly choosing \(a_{ij}^{(n)}, b_{ij}^{(n)} \in [-1, 1]\), while taking \(a_{ij}^{(0)} = b_{ij}^{(0)} = 0\) for \(n \geq 2\). Since
such a network typically does not synchronize, our approximation of independent IP after resetting is appropriate. We take \( g = 2e^{i\phi} \) and compute \( S_{ij} \) from an ensemble of \( I = 10^4 \) IP to obtain the numerical approximations of \( a_{ij}^{(1)} \) and \( b_{ij}^{(1)} \) via Eq. (4). In Fig. 2 we compare the numerical \( a_{ij}^{(1)} \) and \( b_{ij}^{(1)} \) (crosses) with the actual values (circles). All values display a very good agreement for both linked node pairs (different from zero) and nonlinked node pairs (zero). We have not only revealed the adjacency matrix \( A_{ij} \), but also found the interaction parameters \( a_{ij}^{(1)} \) and \( b_{ij}^{(1)} \), thus reconstructing the entire dynamical network.

Below we discuss the limitations of our method. If the available data ensemble \( I \) is too small, the statistics is poor and the obtained network characteristics have large uncertainties, which typically decrease as \( I^{-1/2} \). To illustrate this, in Fig. 3(a) we present the numerical values of parameter \( a_{ij}^{(1)} \), computed for network in Fig. 1 using the ensemble of IP \( \varphi \) of size \( I \). While the distinction between links and nonlinks can already be seen for \( I \approx 10^3 \), for good approximation one needs \( I \approx 10^4 \) (as done in Fig. 2). For higher Fourier harmonics, the convergence is gradually slower, but maintains the same properties.

Another limitation is related to the validity of our independence assumption for the ensemble of IP which is expressed by the separability of distribution \( \rho(\varphi) \) Eq. (5). This heavily depends on the network’s dynamical regime and the resetting strength. For a full synchrony and weak kicking, the reset state is expected to be strongly correlated, whereas for chaotic dynamics and strong resets, the independence assumption is essentially correct. To study this, we consider again the network from Fig. 1, but now we fix all frequencies to \( \omega_i = 1 \), and take all interactions to be attractive \( a_{ij}^{(1)} = 1, b_{ij}^{(1)} = 0 \) (Kuramoto-type model with identical oscillators). After allowing the network to synchronize, we apply kicks as described above (\( \tau \gg t_{\text{synch}} \)). This time, however, for each \( K \in [0, 10] \) we randomly choose kicking strengths \( K_{ij} \) from a Gaussian with standard deviation \( K \). We thus create an ensemble of \( I = 10^4 \) IP from which we compute \( a_{ij}^{(1)} \) as done above.

In Fig. 3(b) we show the reconstructed values of \( a_{ij}^{(1)} \) for links and nonlinks in relation to \( K \). Sufficiently strong kicking \((K \approx 5)\) succeeds in destroying the network’s synchrony and generating the independent IP, from which a good approximation of \( a_{ij}^{(1)} \) is computed. Moderate \( K \approx 1 \) applied previously is now insufficient. This furthermore depends on the relation between \( \tau \) and \( t_{\text{synch}} \); if \( \tau \ll t_{\text{synch}} \) (frequent resets), the separability of \( \rho \) is easier to achieve. Too strong kicking can also induce correlations in \( \varphi \), regardless of dynamical regime and \( \tau \). However, note that \( \rho \) can be estimated using more elaborate techniques than simple one-dimensional kernels [18], which can in principle yield a good estimate even in the nonseparable case. On the other hand, phase resetting is potentially not the only mechanism of obtaining the ensemble \( \varphi \); recall that our theory with a known \( \rho(\varphi) \) works equally well for any case, including full synchrony and inseparability.

Adding noise terms to the right-hand side of Eq. (1) does not formally change the derivation of our main result, rendering our theory valid in the presence of noise. However, in light of the discussion above, noise will have an effect on the performance of method: additional uncertainty due to larger fluctuations of the estimated \( \varphi \) require larger ensembles to achieve the desired precision. On the other hand, noise may play a constructive role by

![Diagram](https://example.com/diagram.png)
destroying the undesired correlations within $\varphi$, and thus facilitating the separability of $\rho$.

While the experimental techniques for measuring $\varphi$ are already in use [14], in a potential realistic application of our method a problem may arise in relation to the measurement of $\varphi$. The entire cycle of a real oscillator is often not accessible; instead, one can observe only a single event per period (e.g., a spike produced by a neuron). In such cases, one is forced to estimate the instantaneous frequencies relying solely on the time intervals between the spikes. To illustrate this, we consider again the system studied in Fig. 3(b), but now we replace $a_{ij}^{(l)}$ with $\varepsilon a_{ij}^{(l)}$. The parameter $\varepsilon$ (coupling strength) controls the ratio between the oscillation time scale (period) and the interaction time scale (synchronization). Rather than computing instantaneous $\varphi$ after each reset, we observe only the event of an oscillator passing through the phase value $\varphi = 0$ (spike), and estimate both $\varphi$ and $\varphi'$ from the first two spikes observed after resetting. We then reconstruct the values of $a_{ij}^{(l)}$ using the ensemble of $I = 10^5$ IP as done before (strong resetting is applied immediately after the spikes are recorded). The results shown in Fig. 3(c) have a clear physical interpretation: for too small coupling $\varepsilon \ll 0.03$ the links cannot be revealed since the interaction is too weak. For too large $\varepsilon \gg 0.4$ the two time scales are too close, and the detection is again impossible since the distribution of phases changes significantly over a period. However, between these extremes, there is a range of coupling around $\varepsilon \sim 0.1$ where the two time scales are well separated allowing a reliable reconstruction. This shows that with an adequately big ensemble our method works even if the entire oscillator cycle is not accessible: errors in the estimation of $\varphi$ and $\varphi'$ play a role similar to the noise. The method fails in the case of too strong coupling, similarly to the case of too weak resetting after synchronization [cf. Fig. 3(b)].

In conclusion, we proposed a method of reconstructing oscillator networks by repeating random phase resets, applicable to a general network irrespectively of the dynamical regimes. Our theory emphasizes the importance of the transient dynamics in the context of network reconstruction, thus complementing the available techniques that rely on time series recorded in final stationary state. Our theoretical model can be generalized to models beyond Eq. (1). If the couplings depend on two phases in a more general way, or depend on more than two phases, one should use more elaborate test functions. For high-dimensional oscillators only a single scalar might be observable: our method can still be applied through the appropriate transformation to phases [6]. Finally, a real experimental situation may involve a network whose dynamics cannot be reset for all nodes simultaneously, which renders the independence assumption invalid. This is a much more challenging and realistic case that requires additional study.

Support from DFG via project FOR868 is acknowledged. We thank A. Díaz-Guilera for useful discussions.

[19] Using the kernel $Q = e^{\sigma(i - j)}$, we find for each node $i$ the optimal value $\sigma_i$ by cross-validation via the maximal likelihood method [18]. This value defines the 1D distribution estimate $\rho_i(\varphi) = \sum Q(\varphi - \varphi_i) = \sum e^{\sigma_i(\varphi - \varphi_i)}$ for this node, based on the empirical ensemble $\{\varphi_i\}_{i=1}^N$.