Globally clustered chimera states in delay–coupled populations

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We have identified the existence of globally clustered chimera states in delay coupled oscillator populations and find that these states can breathe periodically, aperiodically and become unstable depending upon the value of coupling delay. We also find that the coupling delay induces frequency suppression in the desynchronized group. We provide numerical evidence and theoretical explanations for the above results and discuss possible applications of the observed phenomena.

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The existence of chimera states (states characterized by the separation of identical oscillator groups into synchronized and desynchronized subgroups) in coupled oscillator populations came as a surprise in the study of synchronization phenomenon in complex systems. Since its discovery [1, 2], various theoretical and numerical developments have been reported on the stability of chimera states and their existence in systems with varied structures [2, 3], including time delay [4]. By and large, synchronization in coupled oscillator systems has been analytically and numerically investigated in a rigorous manner over the past years [5, 6]. Possible routes to global synchronization and methods to control synchronization have also been proposed [6, 7]. However, complete understanding of the effects induced by coupling delay in synchronization of coupled oscillator systems is still an open problem. The consideration of delayed coupling is vital for modeling real life systems. For example, in a network of neuronal populations, there is certainly a significant delay in propagation of signals. In addition there can also be synaptic and dendritic delays. Other examples include, finite reaction time of chemicals, finite transfer time associated with the basic mechanisms that regulate gene transcription and mRNA translation.

In this paper, we demonstrate that coupling delay can induce globally clustered chimera (GCC) states in systems having more than one coupled identical oscillator (sub) populations. By a GCC state, here we mean a state where the system, which has more than one (sub) population, splits into two different groups, one synchronized and the other desynchronized, each group comprising of oscillators from both the populations (note that this is in contrast to the chimera state where one of the populations is synchronized while the other is desynchronized). The system under study is a system of two populations of identical oscillators coupled through a finite delay, represented by the following equation of motion

$$\dot{\theta}_{i}^{(1,2)} = \omega - \frac{A}{N} \sum_{j=1}^{N} f(\theta_{i}^{(1,2)}(t) - \theta_{j}^{(1,2)}(t - \tau_{1})) + \frac{B}{N} \sum_{j=1}^{N} h(\theta_{i}^{(1,2)}(t) - \theta_{j}^{(2,1)}(t - \tau_{2})).$$

A typical example of such a system is the two groups of interacting neurons in the brain such as those in the cortex and the thalamus [6]. Here $\omega$ is the natural frequency of the oscillators in the populations and it is the same for all oscillators in both the populations making all of them identical. However, the two populations are distinguished by the initial distribution of their phases; the phases are uniformly distributed between 0 and $\pi$ for the first population and between $\pi$ and $2\pi$ for the second population. $A$ and $B$ refer to coupling strengths within and between populations, respectively. The functions $f$ and $h$ are $2\pi$–periodic that describe the coupling. $N$ refers to the size of the populations. The complex mean field parameters $X^{(1,2)} + iY^{(1,2)} = r^{(1,2)}e^{i\theta^{(1,2)}} = \frac{1}{N} \sum_{j=1}^{N} e^{i\theta^{(1,2)}_{j}},$ characterize synchronization within a population but not global clustering. $\tau_{1}$ and $\tau_{2}$ quantify coupling delay within and

![FIG. 1: (Color online) Occurrence of (stable) GCC in system (1) as explained in the text. Top panel: Global clustering phenomenon - synchronized and desynchronized (frequency suppressed) groups have oscillators from both the populations. Bottom panel: One of the populations is synchronized and the other desynchronized (frequency suppressed). Green (light gray) and red (dark gray) lines represent oscillators in the first and the second populations, respectively. Here \( \{f, h\} = \{\sin(\theta), \cos(\theta)\} \), \( \tau_{1} = n\tau_{2} = n\pi \) with \( n = 1 \) (top panel) \( A = 1.2, B = 1 \) and \( \tau = 2 \), (bottom panel) \( A = 1.6, B = 1 \) and \( \tau = 1 \).}
between populations, respectively.

The investigation is motivated by the numerical discovery of the existence of GCC states in a system of two identical populations that are delay–coupled and are given by Eq. (1) (see Fig. 1). We found that coupling delay can induce splitting of identical delay–coupled populations into desynchronized frequency suppressed (vanishing oscillating frequencies) clusters and synchronized clusters. This splitting can occur either within the populations or between the populations. The former represents the chimera and the latter is the GCC, as noted earlier. Further, the GCC state need not be stable but it can either breathe or can be unstable as will be discussed later.

For illustrative purpose, we simulate system (1) using Runge–Kutta fourth order routine with a time step of 0.01 (the results are not affected by decreasing the time step below 0.01). For all the numerical plots shown, we allow a transient time of 2000 units and take $N = 32$ (the results have been verified to be independent of the size of the system) and $\tau_1 = n \tau_2 = n \tau$, where $n$ is an arbitrary constant. We further found that the GCC need not be stable but can breathe depending upon the value of the coupling delay. Since the coherence parameter $r$ quantifies synchronization within a population, it can also be used to quantify a breathing or unstable chimera. However, as mentioned earlier, global clustering cannot be quantified using this order parameter. Therefore, in order to quantify a breathing GCC numerically, after allowing the transients, we identify those oscillators whose $\theta_{DS}$ are equal for all times and neglect them so as to end up with the desynchronized group (that comprises oscillators from both the populations, whose size is $N_{DS}$) and calculate its order parameter as

$$r_{DS} e^{i \psi_{DS}} = \frac{1}{N_{DS}} \sum_{j=1}^{N_{DS}} e^{i \theta_{DS}}, \tag{2}$$

where $N_{DS} = 2N - N^s$. This order parameter $r_{DS}$ can be used to quantify both the chimera and GCC states and also valid for cases where there exists more than one synchronized cluster. Such multi-cluster states also occur for model (1), the details of which will be published elsewhere. While a GCC is breathing, one of the groups is completely synchronized while the desynchronized group continuously fluctuates. Fig. 2 illustrates breathing GCC where we plot the order parameters $r_{DS}$ for two different values of $\tau$ in the top panel. The grey line represents a long period breather for $\tau = 3.6$ where switching occurs between frequency suppressed synchronized state and the desynchronized state. Increasing $\tau$ further to 4 results in a short period breather (the black line) where the desynchronized state oscillates similar to the previous case but in a faster manner.

The GCC can also be unstable where the oscillators in the desynchronized group remain desynchronized for a while after which this state loses its stability and all the oscillators lock to one phase. Thus at this stage the GCC loses stability and a two clustered synchronized state becomes stable. Therefore, finally the system goes from a GCC to a state with two separate synchronized clusters. This phenomenon is depicted in Fig. 3 where for a sufficiently large value of $\tau$ the GCC breathes in an aperiodic manner (top panel). On decreasing $\tau$, this breather loses stability and the desynchronized group entrains itself to a synchronized state (bottom panel). The regions of occurrence of these phenomena in the phase plane, obtained numerically corresponding to Fig. 3 is shown in Fig. 4. The black line is the stable limit cycle attractor of the synchronized group (which is always the same whatever the value of the entrainment frequency of the synchronized group is). The grey region represents the aperiodic

FIG. 2: Illustration of a breathing GCC state with $A = 0.3$, $B = 0.2$, $n = 1$, $\{f, h\} = \{\sin(\theta), \sin(\theta)\}$ and initial condition close to the GCC state. Grey and black lines represent the long and short–periodic breather with $\tau = 3.6$ and $\tau = 4$, respectively. Order parameter $r_{DS}$ and the corresponding phases $\theta_{DS}$ (see text) are plotted against time in the top and bottom panels, respectively.

FIG. 3: Top panel: Aperiodic breathing GCC with $\tau = 5$ and $N_{DS} = 17$; Bottom panel: Unstable breathing GCC with $\tau = 4$ and $N_{DS} = 12$, (left) order parameter $r_{DS}$ and (right) the corresponding phases $\theta_{DS}$ of the desynchronized group. Here $A = 0.6$, $B = 0.3$, $n = 1$, $N = 32$ and $\{f, h\} = \{\sin(\theta), \cos(\theta)\}$. 

FIG. 4: 

\[ e^{i \psi_{DS}} = \frac{1}{N_{DS}} \sum_{j=1}^{N_{DS}} e^{i \theta_{DS}}, \tag{2} \]
synchronized state is stable. This provides the stability
region in a parametric form as

\[ \int_{t_0}^{\infty} [A f'(\beta) - |A f'(\beta) + B h'(\beta)|] dt = \infty. \tag{5} \]

The stability of the global synchronization state is determined by the integrand in this condition. From equations (3–5) it becomes evident that the stability of the synchronized/desynchronized state switches periodically between stable and unstable states depending on the signs of \( A \) and \( B \), since \( h \) and \( f \) are \( 2\pi \) periodic. This is obvious from Fig. 5, where on increasing \( \tau \), regions of synchronization and desynchronization alternate each other. This is in agreement with the numerical analysis as pointed out earlier and hence forms a theoretical basis. The GCC state can be expected near the stability boundaries shown in Fig. 5. This is evident from the numerical results depicted in Figs. 1 and 3.

The knowledge about synchrony control methods is very important because synchronization is desirable sometimes as in neuronal networks while they support cognition via temporal coding \( [9, 11] \) and in the case of lasers and Josephson junction arrays \( [12] \). However, synchronization can also be dangerous in cases like epileptic seizures \( [13] \), Parkinson’s tremor \( [14] \), or pedestrians on the Millennium Bridge \( [6] \). For example, in \( [3] \) a thalamocortical model of asymmetrically interacting neuronal populations has been proposed to simulate the state of emergence from deep to light anaesthesia. The model results revealed the fact that successful coding of information and consciousness is achieved by the occurrence of global synchronization between the thalamus and the

breather. The GCC is unstable while in the white region between these two; a GCC in this white region is attracted to the limit cycle and a stable synchronized state is established (as shown in Fig. 3 (bottom panel)). A GCC in the innermost white region is always stable. The sizes of all these regions change with respect to system parameters.

Thus we find that, for a given set of system parameters, increasing/decreasing (depending on the values of the parameters \( A, B \) and \( \tau \), since the behaviour repeats itself periodically as will be discussed later under Fig. 3) the coupling delay parameter \( \tau \) results in the following sequence of GCC dynamics: stable GCC, long–period breather, short–period breather, aperiodic breather and unstable GCC leading to global synchronization. Further increase in \( \tau \) from the global synchronization state leads to a stable GCC by following the above mentioned route in the reverse order.

If we are able to discriminate the regions of stability of the synchronized and the desynchronized states, we will be able to expect the occurrence of GCC near these boundaries with reference to the numerical observations. This expectation also depends on the fact that the stability of the GCC state changes periodically with respect to \( \tau \) incorporating regions of synchronization and desynchronization. In order to gain a better understanding of the numerically observed phenomena, we analyze system (1) in the continuum limit \( N \to \infty \). We write down the continuity equation [5]

\[ \rho_{f, h} = \Omega \rho_{f, h}, \]

for the density of phases \( \rho \) and then express \( \rho \) and \( \{f, h\} \) as Fourier expansions,

\[ \rho = \sum_{k=-\infty}^{\infty} \rho_k e^{i k \theta} \quad \text{and} \quad \{f, h\} = \sum_{k=-\infty}^{\infty} \{f, h\}_k e^{i k \theta}. \]

Now by considering only the non-trivial \( k \)th Fourier mode we arrive at the eigen value of that mode \( \lambda_k = A e^{-\lambda_k \tau} \pm B e^{-\lambda_k \tau} - i \omega_0 \) which characterizes the stability of the desynchronized state. Here \( A = ik f_k A, B = ik h_k B \) and \( \omega_0 = k(\omega - Af_0 \mp Bh_0) \). Assuming \( \lambda_k = -i \beta / \tau \), one can find the \( k \)th stability region in a parametric form as

\[ B = \pm k A \frac{|f_k| \cos(n \beta - \alpha_f)}{|h_k| \cos(\beta - \alpha_h)}; \quad \tau = \beta / |k(\omega_0 + Af_0 n \tau) + Bh_0 n \tau| \pm B h_k |sin(\beta - \alpha_h)|^{-1}. \tag{3} \]

where \( \{f, h\}_k = -i [\{f, h\}_k e^{i \theta(f, h)} \text{ and } \tau_1 = n \tau_2 = n \tau. \)

The overall stability of the desynchronized state is determined by the overlap of these domains for all the modes.

Now it is also of importance to investigate the stability of the synchronized state for which we consider the solution to the synchronization state \( \theta^{[1,2]} = \Omega t \). With this solution, system (1) becomes \( \Omega = \omega - Af(n \Omega \tau) \mp Bh(n \Omega \tau) \). Along with this relation, the condition \( Af'(n \Omega \tau) + Bh'(n \Omega \tau) > 0 \) should also be satisfied in order that the synchronized state is stable. This provides the stability regime

\[ B = \mp Af'(n \beta) / h'(\beta); \quad \tau = \beta / \omega - Af(n \beta) \mp Bh(\beta), \tag{4} \]

where \( \beta = \Omega \tau \). The parametric forms (3) and (4) separate the desynchronization and synchronization regimes.
Global synchronization regions.

II. Synchronization of the populations individually and III. theory with

FIG. 5: (Color online) Stability regions as obtained from

theory with the stability analysis on the incoherent ( in equations (3)-(5). Note that the boundaries obtained by

and grey symbols correspond respectively to the + signs and grey symbols are the locations of the GCC as from the numerical

examples in Figs. 1 and 3. The red

and black symbols are the stable, unstable and the breathing GCC respectively.

cortex. Further, it was elucidated that consciousness and cognition are kept away during deep anaesthesia because of the lack of phase locking between the cortex and the thalamus. This is one example of a situation where
global clustering/synchronization and hence controlling the same prove to be very crucial. There are various methods to control synchronization (even its rate and velocity). However if we could handle it all with one parameter, it makes life much easier.

In summary, a new type of globally clustered chimera states have been identified in delay coupled populations – a system of two identical, delay-coupled populations split into two groups, one synchronized and the other desynchronized, each group having a fraction of oscillators from both the populations. We have found that this state need not be stable always but can breathe periodically, aperiodically or become unstable, depending upon the value of coupling delay. A modified version of the order parameter is introduced in order to capture these phenomena. In the presence of coupling delay, frequency suppression is induced in the desynchronized group. We have also provided analytical explanations of the observed effects on the basis of linear stability theory. The illustrative model presented here can be considered as a phenomenological model of oscillatory neural networks. Since coupling delay induces globally clustered chimera, this can prove to be a mechanism for temporal coding of information and cognition and also for memory storage in the nervous system.

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[1] Y. Kuramoto and D. Battogtokh, Nonlinear Phenom. Complex Syst. 5, 380 (2002); in Nonlinear Dynamics and Chaos: Where Do We Go From Here?, edited by S. J. Hogan et al. (Institute of Physics, Bristol, England, 2003), p. 209; S.I. Shima and Y. Kuramoto, Phys. Rev. E. 69, 036213 (2004).
[2] D. M. Abrams and S. H. Strogatz, Phys. Rev. Lett. 93, 174102 (2004).
[3] D. M. Abrams, R. Mirollo, S. H. Strogatz and D. A. Wiley, Phys. Rev. Lett. 101, 084103 (2008).
[4] G. C. Sethia, A. Sen and F. M. Atay, Phys. Rev. Lett. 100, 144102 (2008); O. E. Omelchenko, Y. L. Maistrenko and P. A. Tass, Phys. Rev. Lett. 100, 044105 (2008).
[5] A. T. Winfree, J. Theor. Biol. 16, 15 (1967); Y. Kuramoto, Chemical Oscillations, Waves, and Turbulence (Springer-Verlag, Berlin, 1984); A. Pikovsky, M. Rosenblum, and J. Kurths, Synchronization – A Universal Concept in Nonlinear Sciences (Cambridge University Press, Cambridge, 2001).
[6] S. H. Strogatz, Nature 410, 268 (2001);
[7] A. Sherman, J. Rinzel, Proc. Natl. Acad. Sci. U.S.A. 89, 2471 (1992); P. R. Roelfsema, A. K. Engel, P. Konig and W. Singer, Nature 385, 6612 (1997); W. Singer, Nature 397, 6718 (1999); I. Z. Kiss, Y. M. Zhai, J. L. Hudson, Science 296, 1676 (2002).
[8] M. G. Rosenblum and A. S. Pikovsky, Phys. Rev. Lett. 92, 114102 (2004); H. Daido, K. Nakanishi, Phys. Rev. Lett. 96, 054101 (2006); Jane H. Sheeba, V. K. Chandrasekar, A. Stefanovska, and P. V. E. McClintock, Phys. Rev. E 78, 025201(R) (2008).
[9] I. Z. Kiss, M. Quigg, S. H. C. Chun, H. Kori and J. L. Hudson, Biophys. J. 94, 1121 (2008); Jane H. Sheeba, A. Stefanovska, and P. V. E. McClintock, Biophys. J. 95, 2722 (2008).
[10] D. V. Senthilkumar and M. Lakshmanan, Phys. Rev. E 76, 066210 (2007); M. Chen and J. Kurths, Phys. Rev. E 76, 036212 (2007).
[11] W. Singer, Neuron 24 49 (1999); P. Fries, Trends. Cogn. Sci. 9 474 (2005); Y. Yamaguchi, N. Sato, H. Wagatsuma, Z. Wu, C. Molter and Y. Aota, Curr. Opin. Neurobiol. 17, 197 (2007).
[12] B. R. Trees, V. Saranathan, D. Stroud, Phys. Rev. E 71, 016215 (2005); F. Rogister, R. Roy, Phys. Rev. Lett. 98 104101 (2007).
[13] L. Timmermann, J. Gross, M. Dirks, J. Volkman, H. Freund and A. Schnitzler, Brain 126, 199 (2003); J. A. Goldberg, T. Boraud, S. Maraton, S. N. Haber, E. Vassia, and H. Bergman, J. Neurosci. 22, 4639 (2002).
[14] B. Percha, R. Dzakpasu, M. Zochowski and J. Parent, Phys. Rev. E 72, 031909 (2005); M. Zucconi, M. Manconi, D. Bizzozero, F. Rundo, C. J. Stam, L. Fermini-Strambi, R. Ferri, Neural. Sci. 26 199 (2005).