Breathing and switching cyclops states in Kuramoto networks with higher-mode coupling

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Cyclops states are intriguing cluster patterns observed in oscillator networks, including neuronal ensembles. The concept of cyclops states formed by two distinct, coherent clusters and a solitary oscillator was introduced by Munyaev *et al.* [Phys. Rev. Lett. **130**, 107021 (2023)], where we explored the surprising prevalence of such states in repulsive Kuramoto networks of rotators with higher-mode harmonics in the coupling. This paper extends our analysis to understand the mechanisms responsible for destroying the cyclops' states and inducing dynamical patterns called breathing and switching cyclops states. We first analytically study the existence and stability of cyclops states in the Kuramoto-Sakaguchi networks of two-dimensional oscillators with inertia as a function of the second coupling harmonic. We then describe two bifurcation scenarios that give birth to breathing and switching cyclops states and their hybrids are prevalent across a wide coupling range and are robust against a relatively large intrinsic frequency detuning. Beyond the Kuramoto networks, breathing and switching cyclops states promise to strongly manifest in other physical and biological networks, including coupled theta neurons.

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I. INTRODUCTION

Phase oscillator networks have emerged as a paradigmatic 22 reduced model for describing emergent cooperative properties 23 of diverse real-world systems, including neuronal networks 24 [1-3], laser arrays [4-6], and power grids [7-9]. The cele-25 brated Kuramoto model of one-dimensional (1D) oscillators 26 [10,11] and its extension to two-dimensional (2D) oscil-27 lators with inertia [12] captures the essence of the phase 28 networks and provides a concise framework to explore the 29 richness of their cooperative dynamics [13-19]. These dy-30 namics include full [20–24], partial [25,26], explosive [27–29] 31 and asymmetry-induced synchronization [30,31], chimeras 32 [32–39], solitary states [40–44], clusters [45–48], generalized 33 splay [49], and cyclops states [50]. The cooperative dynamics 34 of Kuramoto networks with attractive coupling have been 35 studied more extensively than their counterparts in repulsive 36 networks. While full synchronization is the simplest and most 37 dominant rhythm in attractive networks, splay [51,52] and 38 generalized and cluster splay states [48,49] are expected to 39 be the most probable patterns in repulsive networks. Yet a 40 complete understanding of rhythmogenesis in repulsive net-41 works is still lacking. Two repulsively coupled oscillators tend 42 to achieve antiphase synchronization; however, predicting an 43 outcome of such interactions in large repulsive networks is 44 often elusive. Notably, the role of repulsive connections can be 45 counterintuitive, especially in networks with mixed attractive and repulsive coupling [53–56]. For example, adding pairwise 47 repulsive inhibition to excitatory networks of bursting neurons 48

can synergistically promote synchronization by facilitating transitions between different types of bursting [55].

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In the context of Kuramoto-type networks, the prevailing approach is to model interactions by the first sinusoidal harmonic from a Fourier decomposition of a 2π -periodic coupling function. This simplest choice of the coupling form adequately describes many dynamical features of real-world networks and is analytically tractable. However, higher-order coupling harmonics have been observed to play a significant role in rhythmogenesis in various scenarios. These encompass Kuramoto-type models of neuronal plasticity [57,58], coupled electrochemical oscillators [59], and Josephson junctions [60]. In particular, previous research has demonstrated that augmenting the classical Kuramoto model with higher-mode coupling can result in the emergence of multiple phase-locked states [61,62] and facilitate switching between synchrony clusters [63].

In a recent work [50], we studied rhythmogenesis in re-66 pulsive Kuramoto networks of identical 2D phase oscillators 67 with phase-lagged first-mode and higher-mode coupling. We 68 introduced the concept of cyclops states formed by two dis-69 tinct, coherent clusters and a solitary oscillator reminiscent 70 of the Cyclops's eye. These cyclops states represent a partic-71 ular class of three-cluster generalized splay states [49] with 72 the solitary oscillator maintaining constant phase differences 73 with the coherent clusters. We reported a surprising finding 74 that adding the second or third harmonic to the Kuramoto 75 coupling makes the cyclops state global attractors in a wide 76 range of couplings' repulsion. Beyond Kuramoto networks, 77 we showed that the stabilization of cyclops states by the higher 78 coupling harmonics is also robustly present in theta neurons 79 with adaptive coupling. 80

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This paper extends our previous analysis to reveal higher-81 mode coupling-induced mechanisms for emerging dynamical 82 patterns termed breathing and switching cyclops states. 83 Toward this goal, we derive analytical conditions on the exis-84 tence and stability of cyclops states with constant intercluster 85 phases in the presence of the second coupling harmonic. 86 These conditions reveal two bifurcation scenarios for desta-8 bilizing the cyclops states. In the first scenario, the cyclops 88 states with constant intercluster phases between its three clus-89 ters undergo an Andronov-Hopf bifurcation, preserving their 90 intracluster formations but making the intercluster relative 91 phase differences oscillate periodically. Similarly to breathing 92 three-cluster patterns introduced in [47], we call these breath-93 ing cyclops states. These states can evolve into rotobreathers 94 with intercluster phase differences governed by mixed-mode, 95 oscillatory-rotatory phase difference dynamics. In the second 96 bifurcation scenario, the cyclops state with constant interclus-97 ter phases loses its structural stability but quickly reforms into 98 99 a new cyclops state with a reshuffled configuration. This repetitive switching process yields switching cyclops states. These 100 states are similar to blinking chimeras, also characterized by 101 a death-birth process in which the coherent cluster dissolves 102 and is quickly reborn in a new configuration [64]. 103

We show that breathing, rotobreathing, and switching 104 cyclops states are stable in a wide range of the second 105 harmonic coupling strength and phase lag parameter. Remark-106 ably, breathing and rotobreathing cyclops states are dominant 107 states, acting as the system's global attractors in a large 108 interval of the second harmonic's phase lag parameter, cor-109 responding to the overall repulsiveness of the combined first-110 111 and second-harmonic coupling. We also demonstrate that the cyclops states can robustly emerge in Kuramoto networks 112 of nonidentical oscillators. In [50], we proved that the 2D 113 Kuramoto model with the first and second-harmonic coupling 114 is dynamically equivalent to a network of canonical theta neu-115 rons with adaptive coupling. Therefore, we expect breathing 116 and switching cyclops states to manifest strongly in theta-117 neuron networks, pointing to the broader applicability of our 118 results. 119

The layout of this paper is as follows. In Sec. II we in-120 troduce the oscillator network model and state the problem 121 under consideration, and give formal definitions. In Sec. III 122 we study the existence of cyclops states with constant inter-123 cluster phase differences, called stationary cyclops states. We 124 derive an upper bound for the maximum number of station-125 ary cyclops states with distinct intercluster phase differences. 126 In Sec. IV we derive a four-dimensional (4D) system that 127 governs the dynamics of the intercluster phase differences. 128 We study the stability of the fixed point of the 4D system, 129 which corresponds to constant intercluster phase differences. 130 We derive the conditions under which the fixed point under-131 goes an Andronov-Hopf bifurcation, giving rise to a breathing 132 133 cyclops state. In Sec. V we analyze the variational equations for the transversal stability of stationary cyclops states 134 that determines the stability of their coherent clusters. In 135 Sec. VI we numerically study breathing and switching cyclops 136 states emerging from stationary cyclops states via two distinct 137 bifurcation routes. We demonstrate the emergence of more 138 complex, hybrid dynamical patterns that combine the prop-139 erties of both breathing and switching cyclops states. We also 140

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study the prevalence of different cyclops states and show that
they robustly appear from large sets of randomly chosen initial
conditions. In Sec. VII we show the persistence of cyclops
states against relatively large intrinsic frequency detuning.141
142Section VIII contains concluding remarks and discussions.
The Appendix contains the derivation of the upper bound for
the maximum number of stationary cyclops states.143

II. THE MODEL AND PROBLEM STATEMENT

We consider the Kuramoto-Sakaguchi network of 2D phase 149 oscillators 150

$$\mu \ddot{\Theta}_k + \dot{\Theta}_k = \omega + \sum_{n=1}^N \sum_{q=1}^2 \frac{\varepsilon_q}{N} \sin[q(\Theta_n - \Theta_k) - \alpha_q], \quad (1)$$

where the kth oscillator's phase Θ_k ranges from $-\pi$ to 151 π , and the second-order Kuramoto-Sakaguchi coupling [65] 152 represents a pairwise interaction function $H(\Theta_n - \Theta_k) =$ 153 $\sum_{q=1}^{2} \varepsilon_q \sin[q(\Theta_n - \Theta_k) - \alpha_q].$ The oscillators are assumed 154 to be identical with frequency ω , inertia μ , and phase lag 155 parameters α_1 and α_2 . We consider the phase lag $\alpha_1 \in$ 156 $(\pi/2, \pi)$, that makes the first-harmonic coupling repulsive 157 and fix $\varepsilon_1 = 1$ that corresponds to a strong first-harmonic 158 coupling. Throughout the paper, we also choose and fix a 159 relatively strong inertia $\mu = 1$ that is sufficient to make the 160 dynamics of the 2D system qualitatively distinct from the 1D 161 classical model and enable the emergence of breathing cluster 162 dynamics [46]. We will consider a broader range of $\alpha_2 \in$ 163 $(-\pi, \pi)$, so that the second harmonic may be pairwise attrac-164 tive or repulsive. As a result, the overall combined coupling 165 may be repulsive with H'(0) < 0 or attractive with H'(0) > 0. 166 The latter is possible when the second-harmonic coupling ε_2 167 is sufficiently strong to overcome the first-harmonic coupling 168 contribution. 169

Phase coherence and cluster synchrony in the system (1) 170 can be characterized via the *l*th-order complex Kuramoto 171 parameters [63,66]: 172

$$R_l(t) = \frac{1}{N} \sum_{k=1}^{N} e^{il\Theta_k} = r_l e^{i\psi_l},$$

where r_l and ψ_l define the magnitude and the phase of the *l*th 173 moment Kuramoto order parameter $R_l(t)$, respectively. The 174 first-order scalar parameter $r_1 = |R_1|$ characterizes the degree 175 of phase synchrony with $r_1 = 1$ corresponding to full phase 176 synchrony. Splay states or generalized splay states $\Theta_k = \omega t + \omega_k +$ 177 $\varphi_k, k = 1, \dots, N$ with constant nonuniform relative phases 178 $\varphi_k \in [-\pi, \pi]$ satisfy the condition $r_1 = 0$ in the 2D Kuramoto 179 model with the first-harmonic coupling ($\varepsilon_2 = 0$). The second-180 order scalar parameter $r_2 = |R_2|$ determines the degree of 181 cluster synchrony. In the case of the first-harmonic coupling 182 $(\varepsilon_2 = 0)$, r_2 controls the stability of generalized splay states 183 so that increasing r_2 enlarges their stability parameter regions 184 [49,50]. It was shown in [50] that generalized splay states with 185 a maximum r_2 are (i) two-cluster symmetric splay states (for 186 even N) and (ii) three-cluster splay states with the relative 187 phases (for odd N): 188

$$\varphi_1 = \varphi_2 = \dots = \varphi_{M-1} = \gamma, \quad \varphi_M = 0,$$

$$\varphi_{M+1} = \dots = \varphi_N = -\gamma, \quad (2)$$



FIG. 1. Snapshot of a breathing cyclops state in network (1) of 11 oscillators. Periodically oscillating x(t) and y(t) govern the phase difference between the synchronous clusters (blue and pink circles) and the solitary oscillator (gray circle). The solitary oscillator's phase is chosen at $\theta_M = 0$ as a reference. Parameters are $\alpha_1 = 1.7$, $\varepsilon_2 = 0.08$, $\alpha_2 = -0.3$.

where $\gamma = \arccos[1/(1-N)]$, M = (N+1)/2, and the 189 choice of the reference zero phase for φ_M is arbitrary. We 190 termed three-cluster splay states (2) cyclops states. Adding the 191 second-harmonic coupling with $\varepsilon_2 \neq 0$ breaks their symmetry 192 in γ and makes r_1 nonzero, albeit small. We demonstrated in 193 [50] that the second- or higher-harmonic coupling can make 194 these asymmetric patterns dominant states. In this paper, we 19 generalize the definition of cyclops states (2) for the system (1) with second-harmonic coupling and odd N by relaxing 197 the condition $r_1 = 0$. As a result, we refer to the following 198 three-cluster state determined by the three-dimensional cluster 199 manifold: 200

$$D(3) = \begin{cases} \Theta_1(t) = \dots = \Theta_{M-1}(t) = x + \Omega t, \\ \Theta_M(t) = \Omega t, \\ \Theta_{M+1}(t) = \dots = \Theta_N(t) = y + \Omega t \end{cases}$$
(3)

as to a stationary cyclops state in which two equal clusters 20 of M-1 oscillators rotate with the common frequency Ω , 202 preserving the stationary phase differences $x = \gamma_1$, and $y = \gamma_2$ 203 with the *M*th solitary oscillator. Note that due to the asym-204 metry of the relative phases $x \neq y$, the common rotational 205 frequency Ω differs from the oscillators' intrinsic frequency 206 ω and can be derived explicitly from (1) as shown in the next 207 section. 208

Similarly to (2), the stationary cyclops state defined by (3) can be conveniently represented within the reference frame $\theta_k = \Theta_k - \Omega t$, k = 1, ..., N where the relative constant phase of the *M*th oscillator, θ_M , is chosen at zero so that

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$$\theta_1 = \dots = \theta_{M-1} = x, \quad \theta_M = 0,$$

$$\theta_{M+1} = \dots = \theta_N = y. \tag{4}$$

0

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Due to the system's global coupling symmetry and equal clus-213 ter sizes M - 1, the existence of the stationary cyclops state 214 with intercluster phase differences $x = \gamma_1$ and $y = \gamma_2$ implies 215 the existence of its counterpart with $x = \gamma_2$ and $y = \gamma_1$. Thus, 216 cyclops states exist in symmetrical pairs. In the following, we 217 218 will analyze the existence and stability of stationary cyclops states in the system (1) with odd N. We will describe two 219 main scenarios for destabilizing a stationary cyclops state that 220 yield (i) a breathing cyclops state with periodically oscillating 221 x(t), y(t) so that the intercluster phase differences are bounded 222 as $|x(t)| < \pi$, $|y(t)| < \pi$ to produce no phase slips (Fig. 1) 223

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and (ii) *a switching cyclops state*, representing a repetitive death-birth process in which the clusters disintegrate to form a new cyclops state with a new reshuffled cluster configuration and a new solitary node. 227

We will also study how the breathing and switching cyclops states can (i) further evolve into more complex dynamical patterns, including hybrid switching-breathing states, (ii) become globally stable, and (iii) persist against intrinsic frequency detuning.

III. POSSIBLE CONSTANT INTERCLUSTER PHASE DIFFERENCES

We seek to find permissible stationary cyclops states as a function of the system's parameters. To determine the constant phase differences x, y, and the rotational frequency Ω , we substitute the stationary cyclops state solution (3) into (1) and obtain the system of nonlinear transcendental equations: 236

$$\omega - \Omega - \sum_{q=1}^{2} \frac{\varepsilon_q}{N} \left(\sin(qx + \alpha_q) + \frac{N-1}{2} \{ \sin \alpha_q + \sin[q(x-y) + \alpha_q] \} \right) = 0,$$

$$\omega - \Omega - \sum_{q=1}^{2} \frac{\varepsilon_q}{N} \left(\sin \alpha_q - \frac{N-1}{2} [\sin(qx - \alpha_q) + \sin(qy - \alpha_q)] \right) = 0,$$

$$\omega - \Omega - \sum_{q=1}^{2} \frac{\varepsilon_q}{N} \left(\sin(qy + \alpha_q) + \frac{N-1}{2} \{ \sin \alpha_q + \sin[q(y-x) + \alpha_q] \} \right) = 0.$$
 (5)

We subtract the second equation from the first and third equations of (5) to eliminate Ω and obtain the system of two equations for finding the unknown constants *x* and *y*: 242

$$\frac{N-3}{2}\sum_{q=1}^{2}\varepsilon_{q}\sin\alpha_{q} + \sum_{q=1}^{2}\varepsilon_{q}\sin(qx + \alpha_{q}) + \frac{N-1}{2}\left(\sum_{q=1}^{2}\varepsilon_{q}\sin(qx - \alpha_{q}) + \sum_{q=1}^{2}\varepsilon_{q}\sin(qy - \alpha_{q}) - \sum_{q=1}^{2}\varepsilon_{q}\sin[q(y - x) - \alpha_{q}]\right) = 0,$$

$$\frac{N-3}{2}\sum_{q=1}^{2}\varepsilon_{q}\sin\alpha_{q} + \sum_{q=1}^{2}\varepsilon_{q}\sin(qy + \alpha_{q}) + \frac{N-1}{2}\left(\sum_{q=1}^{2}\varepsilon_{q}\sin(qx - \alpha_{q}) + \sum_{q=1}^{2}\varepsilon_{q}\sin(qy - \alpha_{q}) - \sum_{q=1}^{2}\varepsilon_{q}\sin[q(x - y) - \alpha_{q}]\right) = 0.$$
(6)

In turn, we find the rotational frequency Ω from the second equation of (5):

$$\Omega = \omega - \frac{1}{N} \sum_{q=1}^{2} \varepsilon_q \sin \alpha_q + \frac{N-1}{2N} \left[\sum_{q=1}^{2} \varepsilon_q \sin(qx - \alpha_q) + \sum_{q=1}^{2} \varepsilon_q \sin(qy - \alpha_q) \right]$$
(7)

with x and y calculated from (6).

Due to the complexity of system (6), its solution for x and 246 y cannot be found in closed form. Yet we derive an upper 247 bound for the maximum number of stationary cyclops states 248 with distinct x and y. To do so, we transform the real-valued 249 system (6) into a system of complex polynomial equations and 250 apply the Bernshtein theorem [67], a practical tool in algebra 251 that bounds the number of nonzero complex solutions by the 252 mixed volume of their Newton polytopes. The details of this 253 analysis are quite technical and are delegated to the Appendix. 254 This analysis shows that the complex form of system (6) may 255 have up to 17 possible solutions (including some nonphysical) 256 corresponding up to 16 stationary cyclops states with distinct 257 ordered pairs of constant phase differences x, y. As stationary 258 cyclops states exist in pairs, there may be at most eight com-259 binations of x, y (up to the cluster permutation $x \leftrightarrow y$). It is 260 worth noticing that there is a continuum of stationary cyclops 26 states with a given pair (x, y) due to an arbitrary choice of the 262 reference solitary state's phase θ_M . 263

Figure 2 displays the number of different stationary cy-264 clops states calculated by solving the complex polynomial 265 equation (A1) using the NSolve function of Wolfram Math-266 ematica. Note that this number equals two for small values 267 of the second-harmonic amplitude ε_2 . This pair of stationary 268 cyclops states with $x = \gamma_1$ and $y = \gamma_2$ ($x = \gamma_2$ and $y = \gamma_1$) 269 emerges continuously from the symmetrical cyclops state (2) 270 that exists in the system (1) in the absence of the second-271 harmonic coupling ($\varepsilon_2 = 0$). As Figs. 2(a) and 2(b) indicate, 272 increasing ε_2 increases the number of coexisting stationary 273 cyclops states and induces richer dynamics. In the following, 274 we will derive general stability conditions for a permissible 275 stationary cyclops state and specify these conditions to the 276 cyclops state (4) with $x = \gamma_1$ and $y = \gamma_2$ from the dashed 277 parameter region in Fig. 2 where no other cyclops states with 278 distinct x and y (up to permutation of clusters $x \leftrightarrow y$) exist. 279

IV. STABILITY OF THE INTERCLUSTER PHASE DIFFERENCES

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We seek to obtain the conditions for the stability of the constant intercluster phase differences to small perturbations of x and y. The dynamics of the intercluster phase differences are governed by the system

$$\mu \ddot{x} + \dot{x} = \sum_{q=1}^{2} \frac{\varepsilon_q}{N} (\sin \alpha_q - \sin(qx + \alpha_q))$$
$$- \frac{N-1}{2} [\sin(qx - \alpha_q) + \sin(qy - \alpha_q))$$
$$+ \sin \alpha_q + \sin[q(x - y) + \alpha_q] \}),$$



FIG. 2. (a, b) The color shows the number of distinct stationary cyclops states in the network (1) as a function of the second-harmonic coupling and phase lag parameters (α_2 , ε_2). Other parameters are (a) N = 5, $\alpha_1 = 2.0$ and (b) N = 11, $\alpha_1 = 1.7$. The number of cyclops is calculated by numerically finding solutions of system (6). The arrow points to the dashed area corresponding to the stability diagram of Fig. 3. (c) Snapshots of three distinct stationary cyclops states (up to permutation of clusters $x \leftrightarrow y$) for the parameter set N = 11, $\alpha_1 = 1.7$, $\alpha_2 = 0.0$, $\varepsilon_2 = 6.0$ corresponding to the open diamond in the green area in panel (b). The oscillator coloring corresponds to the intercluster differences *x* and *y* according to positive and negative values depicted from the horizontal color bar.

$$\mu \ddot{y} + \dot{y} = \sum_{q=1}^{2} \frac{\varepsilon_q}{N} (\sin \alpha_q - \sin(qy + \alpha_q))$$
$$- \frac{N-1}{2} \{ \sin(qx - \alpha_q) + \sin(qy - \alpha_q) + \sin\alpha_q + \sin[q(y - x) + \alpha_q] \} \}. \tag{8}$$

The 4D dynamical system (8) may be viewed loosely as 286 a system of two nonlinearly coupled driven pendulum-287 like equations with the terms $\sin \alpha_q$ representing constant 288 torques and the sine terms with x and y corresponding to 289 pendulum-like nonlinearities and coupling. The presence of 290 the second-harmonic coupling prevents transforming the sys-291 tem (8) into a more explicit system of two coupled pendula as 292 was achieved for a three-cluster state in [47]. However, the 293 pendulum-like structure of the 4D system (8) points to the 294 possible existence of nontrivial dynamics related to oscillating 295 and even chaotically evolving intercluster phase differences 296 x(t) and y(t). 297

Fixed points of system (8) correspond to constant intercluster phase differences x, y calculated from (6). We aim to study 299

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(9)

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the local stability of the fixed points and derive bifurcation 300

conditions that induce oscillating phase differences x(t), y(t).

Toward this goal, we consider small deviations $\delta x(t)$ and 302 $\delta y(t)$ from a fixed point $x = \gamma_1$, $y = \gamma_2$ corresponding to a 303

stationary cyclops state. So $x(t) = \gamma_1 + \delta x(t)$, $y(t) = \gamma_2 + \delta x(t)$ 304 $\delta y(t)$. We linearize the system (8) in the vicinity of the fixed 305 point state and obtain the following equations that govern the 306 evolution of small deviations $\delta x(t)$ and $\delta y(t)$: 307

$$\mu \delta \ddot{x} + \delta \dot{x} = -\sum_{q=1}^{2} \frac{\varepsilon_q q}{N} \bigg[\cos(q\gamma_1 + \alpha_q) \delta x + \frac{N-1}{2} [\cos(q\gamma_1 - \alpha_q) \delta x + \cos(q\gamma_2 - \alpha_q) \delta y + \cos(q\sigma + \alpha_q) (\delta x - \delta y)] \bigg],$$

$$\mu \delta \ddot{y} + \delta \dot{y} = -\sum_{q=1}^{2} \frac{\varepsilon_q q}{N} \bigg[\cos(q\gamma_2 + \alpha_q) \delta y + \frac{N-1}{2} [\cos(q\gamma_1 - \alpha_q) \delta x + \cos(q\gamma_2 - \alpha_q) \delta y + \cos(q\sigma - \alpha_q) (\delta y - \delta x)] \bigg],$$

where $\sigma = \gamma_1 - \gamma_2$. 308

Following the standard stability approach, we seek solu-309 tions $\delta x(t) = A_1 e^{\lambda t}$, $\delta y(t) = A_2 e^{\lambda t}$ and derive a system of two 310 characteristic equations for finding constants λ , A_1 , and A_2 : 31

$$(\mu\lambda^2 + \lambda)A_1 = -(p_{11}A_1 + p_{12}A_2),$$

$$(\mu\lambda^2 + \lambda)A_2 = -(p_{21}A_1 + p_{22}A_2),$$
 (10)

where 312

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$$p_{11} = \sum_{q=1}^{2} \frac{\varepsilon_q q}{N} \left\{ \frac{N-1}{2} [\cos(q\gamma_1 - \alpha_q) + \cos(q\sigma + \alpha_q)] + \cos(q\sigma + \alpha_q)] \right\},$$

$$p_{12} = \sum_{q=1}^{2} \frac{\varepsilon_q q}{N} \left\{ \frac{N-1}{2} [\cos(q\gamma_2 - \alpha_q) - \cos(q\sigma + \alpha_q)] \right\},$$

$$p_{21} = \sum_{q=1}^{2} \frac{\varepsilon_q q}{N} \left\{ \frac{N-1}{2} [\cos(q\gamma_1 - \alpha_q) - \cos(q\sigma - \alpha_q)] \right\},$$

$$p_{22} = \sum_{q=1}^{2} \frac{\varepsilon_q q}{N} \left\{ \frac{N-1}{2} [\cos(q\gamma_2 - \alpha_q) + \cos(q\sigma - \alpha_q)] + \cos(q\sigma - \alpha_q)] \right\}.$$
(11)

Solving the characteristic system (10) of two coupled 313 quadratic equations to explicitly find λ is out of reach. Instead, 314 we introduce the variable $\Lambda = \mu \lambda^2 + \lambda$ and turn the system 315 (10) into the system of linear equations 316

$$\boldsymbol{P}(A_1, A_2)^T = \Lambda(A_1, A_2)^T,$$
(12)

317 where

$$\boldsymbol{P} = \begin{pmatrix} -p_{11} & -p_{12} \\ -p_{21} & -p_{22} \end{pmatrix}.$$
 (13)

Therefore, the stability of (9) can be assessed from (13) in 318 terms of its eigenvalues Λ . To do so, we aim to determine 319 the boundary of the stability region that is determined by $\lambda =$ 320 $i \text{ Im}\lambda$ and corresponds to an Andronov-Hopf bifurcation of 321

the fixed point that induces oscillating phase differences x(t)322 and y(t). Therefore, we can set $\text{Re}\Lambda + i \text{Im}\Lambda = -\mu(\text{Im}\lambda)^2 +$ 323 *i* Im λ so that the real part equality Re $\Lambda + \mu (Im\Lambda)^2 = 0$ de-324 fines the stability boundary $\lambda = i \text{Im}\lambda$. To ascertain what side 325 of the stability boundary corresponds to stability, we select the 326 test value $\mu = 0$. With this choice, we have $\Lambda = \lambda$, and the 327 inequality $\text{Re}\Lambda < 0$ ensures the stability of the fixed point. 328 Extending this argument to nonzero μ , we can conclude 329 that the inequality $\text{Re}\Lambda + \mu(\text{Im}\Lambda)^2 < 0$ makes $\text{Re}\lambda < 0$ and 330 determines the fixed point stability. Thus, we arrive at the 331 following assertion. 332

Statement 1. [Internal stability of stationary cyclops 333 states]. 334

1. Constant intercluster phase differences $x = \gamma_1$ and y =335 γ_2 of the stationary cyclops state (3) are locally stable iff

$$\operatorname{Re}\Lambda_{1,2} + \mu(\operatorname{Im}\Lambda_{1,2})^{2} < 0_{\star}\Lambda_{1,2}$$
$$= \frac{\operatorname{Tr}\boldsymbol{P}}{2} \pm \frac{\sqrt{(\operatorname{Tr}\boldsymbol{P})^{2} - 4\operatorname{det}\boldsymbol{P}}}{2}, \qquad (14)$$

where TrP and detP are, respectively, the trace and determi-337 nant of matrix P(13) whose coefficients are defined in (11). 338 2. The stability boundary 339

$$\operatorname{Re}\Lambda_{1,2} + \mu(\operatorname{Im}\Lambda_{1,2})^2 = 0 \tag{15}$$

corresponds to an Andronov-Hopf bifurcation that destabilizes 340 the stationary cyclops state, turning it into a breathing cyclops 341 state with oscillating intercluster phase differences x(t) and 342 y(t). 343

The stability of constant phase differences x and y de-344 fined via (14) can be interpreted as the internal (longitudinal) 345 stability of the stationary cyclops state within the invariant 346 three-cluster manifold D determined by (3) with arbitrary, 347 possibly time-varying x(t) and y(t). The stability boundary 348 (15) depicted by the blue solid curve in Fig. 3(a) corresponds 349 to emerging instability of constant phase differences x and 350 y that preserves the three-cluster cyclops formation on the 351 invariant three-cluster manifold D. 352

Having studied the stability of the constant intercluster 353 phase differences, we proceed with the stability analysis of 354 the synchronous clusters, each composed of M-1 oscilla-355 tors. These conditions, paired with the condition (14), shall 356 indicate what stationary cyclops states can stably emerge in 357 the network. 358



FIG. 3. (a) The stability diagram for cyclops states. Regions of stable stationary cyclops states (CS) are shown in blue, switching cyclops states (SCS) in yellow, breathing cyclops states (BCS) in green, and two-cluster regimes (5:6) in white. Analytical boundaries: the blue solid line corresponds to the stability boundary (15), the red dashed line to $\text{Re}\lambda_1^{\text{tran}} = 0$, and the green dash-dotted curve to $\operatorname{Re}\lambda_2^{\operatorname{tran}} = 0$. Two numerical curves marked by the solid circles separate the stability regions of the switching and breathing stationary cyclops states. The black dotted line Γ corresponds to H'(0) = 0. Values above the curve make the coupling attractive and full synchronization locally stable. Stationary cyclops states are found as a solution of system (6) and used as initial conditions. The round, diamond, and circled times correspond to the parameters used in Figs. 4-6. (b) The real part of the eigenvalues, associated with the stationary cyclops state, that determine the (internal) stability of the intercluster phase differences (blue solid line) and transversal (external) stability of the first (red dashed line) and second cluster (green dash-dotted line) for fixed $\varepsilon_2 = \varepsilon_2^*$ and varying α_2 [along the white dashed horizontal line in panel (a)]. The background color indicates the type of the emerged cyclops states as in panel (a). (c) The diagram is similar to panel (b), but for fixed $\alpha_2 = \alpha_2^*$ and varying ε_2 [along the black-white dashed vertical line panel (a)]. The shaded area indicates the bistability of switching and breathing cyclops states. Parameters: $N = 11, \varepsilon_1 = 1.0, \alpha_1 = 1.7, \varepsilon_2^* = 0.08, \alpha_2^* = 0.78.$

V. STABILITY OF SYNCHRONOUS CLUSTERS

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We aim to derive the conditions for transversal stability of the stationary cyclops state (3) that amounts to the stability of the two synchronous clusters composing the stationary cyclops state. We introduce small deviations from the oscillators' phases $\theta_n \longrightarrow \theta_n + \delta \theta_n$, composing the first cluster for n = 1, ..., M - 1 and the second cluster for $n = M + \delta \theta_n$

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1, ..., N. To study the local stability of each synchronous $_{366}$ cluster, we consider the difference variables $_{367}$

$$\xi_n = \delta \theta_{n+1} - \delta \theta_n, \quad n = 1, \dots, M - 2, \tag{16}$$

$$\zeta_n = \delta \theta_{n+1} - \delta \theta_n, \quad n = M + 1, \dots, N - 1$$
(17)

that describe the phase difference dynamics within the first and second clusters, respectively. Therefore, from (1), (3), and (16), (17), we obtain two uncoupled variational equations with time-invariant coefficients. Each of the equations determines the local stability of the corresponding cluster within the cyclops state (3):

$$\mu \ddot{\xi}_n + \dot{\xi}_n + \frac{1}{N} \sum_{q=1}^2 \varepsilon_q q \left\{ \cos(q\gamma_1 + \alpha_q) + \frac{N-1}{2} [\cos\alpha_q + \cos(q\sigma + \alpha_q)] \right\} \\ \xi_n = 0, \quad (18)$$

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where n = 1, 2, ..., M - 2, and

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$$\begin{aligned} u\ddot{\zeta}_n + \dot{\zeta}_n + \frac{1}{N}\sum_{q=1}^2 \varepsilon_q q \left\{ \cos(q\gamma_2 + \alpha_q) + \frac{N-1}{2} [\cos\alpha_q + \cos(q\sigma - \alpha_q)] \right\} \zeta_n &= 0, \end{aligned}$$
(19)

where n = M + 1, ..., N - 1. The variational equations (18) and (19) are stable iff the time-invariant coefficients of the terms ξ_n and ζ_n are positive. Therefore, we can formulate the stability conditions in the following assertion.

Statement 2. [Transversal stability of stationary cyclops379states]. Clusters of oscillators composing the stationary cyclops380clops state (3) are locally stable iff:381

$$\sum_{q=1}^{2} \varepsilon_{q} q \cos(q\gamma_{1} + \alpha_{q}) + \frac{N-1}{2} \left(\sum_{q=1}^{2} \varepsilon_{q} q \cos \alpha_{q} + \sum_{q=1}^{2} \varepsilon_{q} q \cos(q\sigma + \alpha_{q}) \right) > 0,$$

$$\sum_{q=1}^{2} \varepsilon_{q} q \cos(q\gamma_{2} + \alpha_{q}) + \frac{N-1}{2} \left(\sum_{q=1}^{2} \varepsilon_{q} q \cos \alpha_{q} + \sum_{q=1}^{2} \varepsilon_{q} q \cos(q\sigma - \alpha_{q}) \right) > 0,$$
(20)

where the left-hand sides of the inequalities (20) are the coefficients of the variational equations (18) and (19).

It is also straightforward to show that the stationary cyclops state is always stable to the shift of all phases by a constant value $\delta\theta_k = \delta\theta$ (k = 1, ..., N).

It is worth noticing that the eigenvalues $\lambda_{1,2}^{\text{tran}}$ associated with the variational equations (18) and (19) have multiplicity M - 2. Thus, the eigenvalues λ_1^{tran} and λ_2^{tran} define the transversal stability of the first (n = 1, 2, ..., M - 2)and second (n = M + 1, ..., N) clusters, respectively. Figure 3(a) displays their stability boundaries defined by the

BREATHING AND SWITCHING CYCLOPS STATES IN ...

conditions (20) with the left-hand sides set to 0 to correspond to $\text{Re}\lambda_1^{\text{tran}} = 0$ (the red dashed line) and $\text{Re}\lambda_2^{\text{tran}} = 0$ (the green dash-dotted line). To highlight the constructive role of the second-harmonic coupling with $\varepsilon_2 \neq 0$, we chose the parameter values that yield unstable stationary cyclops in the network with only first-harmonic coupling with $\varepsilon_2 = 0$ (see Fig. 3).

As Fig. 3(a) indicates, crossing the stability boundary (15) 400 (the lower border of the region CS) induces breathing cyclops 401 states in the region BCS (green) in accordance with State-402 ment 1. In turn, crossing the upper border of the region CS, 403 composed of the transveral stability boundaries $\text{Re}\lambda_1^{\text{tran}} = 0$ 404 (the red dashed line) and $\text{Re}\lambda_2^{\text{tran}} = 0$ (the green dash-dotted 405 line) can yield either switching cyclops states in the region 406 SCS (yellow) or asymmetrical, two-cluster states with five-407 and six-oscillator synchronous clusters (white region 5:6). In 408 the following, we will primarily focus on the properties of 409 410 emerging breathing and switching cyclops states.

VI. EMERGING BREATHING AND SWITCHING CYCLOPS STATES

We performed numerical calculations using a widely 413 adopted fifth-order Runge-Kutta scheme with a fixed time step 414 0.01 to further validate our analytical results and predictions. 415 Figure 3 confirmed the two main bifurcation scenarios for 416 destroying the stationary cyclops states and generating breath-417 ing and switching cyclops states described by Statements 1 418 and 2. In the first scenario, complex conjugate eigenvalues 419 $\lambda_{1,2}$, that determine the stability of constant intercluster phase 420 421 differences x and y via (10), become purely imaginary and induce oscillating x(t) and y(t) [Fig. 4(c)]. As a result, the 422 stationary cyclops state becomes internally unstable; however, 423 the stability of the clusters preserves and guarantees the emer-424 gence of a breathing cyclops state [see Figs. 4(a) and 4(d) and 425 Supplemental Movie 1 [68] for the animation of the breathing 426 cyclops state dynamics]. Periodic oscillations of the first two 427 order parameters r_1 and r_2 depicted in Fig. 4(b) are a signature 428 of such a breathing cyclops state. As the distance from the 429 stability boundary of the CS region [solid blue line in Fig. 3 430 a(a)] increases when changing the second-harmonic coupling 43 strength ε_2 and phase lag α_2 , the amplitudes of intercluster 432 difference oscillations x(t), y(t), and order parameters $r_1(t)$, 433 $r_2(t)$ increase. It is worth noticing that for the parameters 434 α_1 and ε_1 used in Fig. 3(a), the breathing cyclops state is 435 also stable in the absence of the second-harmonic coupling 436 $(\varepsilon_2 = 0).$ 437

In the second bifurcation scenario determined via State-438 ment 2, the stationary cyclops state loses its transversal 439 stability when one of the eigenvalues $\lambda_{1,2}^{tran}$ becomes positive 440 [Fig. 5(c)]. Note that the real parts of the other eigenvalues 441 controlling the internal stability of the intercluster differences 442 remain negative, thereby preserving the stable component 443 of the saddle dynamics. While the transversal instability 444 of the cyclops state may lead to its complete destruction, 445 it induces a switching cyclops state [Figs. 5(a) and 5(d)] 446 when the transversal instability is weak [note the slightly 447 positive eigenvalue, depicted by the red nabla in Fig. 5(c)]. 448 This nonstationary cyclops state represents a two-stage repet-449 itive process. During the first relatively long stage, the 450



FIG. 4. Breathing cyclops state. (a) The colors depict the phase differences $\theta_k(t) - \theta_6(t)$. The gray strip indicates the reference solitary oscillator. (b) The corresponding values of r_1 and r_2 . (c) The eigenvalues associated with the destabilized stationary cyclops state. Some eigenvalues are repeated. The round (triangular) labels correspond to the internal (transversal) stability. Note a pair of complex eigenvalues with a positive real part (red) that emerged due to an Andronov-Hopf bifurcation and yielded periodic oscillations of intercluster differences. (d) Phase distributions θ_k at several time instants. The arrows indicate the direction of periodic phase clusters' oscillations (see Supplemental Movie 1 [68] demonstrating this breathing cyclops state). The oscillators' coloring represents their relative phase difference with the solitary oscillator as in Fig. 2(c). Parameters N = 11, $\mu = 1.0$, $\varepsilon_1 = 1.0$, $\alpha_1 = 1.7$, $\varepsilon_2 = 0.08$, $\alpha_2 = -0.1$ correspond to the open circle label in Fig. 3(a).

intercluster differences x and y practically do not change, 451 and the synchronous clusters preserve their formation, i.e., 452 the dynamical pattern is similar to a stationary cyclops state 453 (Fig. 5). During the second short stage, one cluster reshuffles 454 so that one node leaves the unstable cluster to become a new 455 solitary oscillator, whereas the remaining oscillators from the 456 cluster merge with the old solitary node. Figure 5(d) and Sup-457 plemental Movie 2 [68] illustrate this process. Accordingly, 458 during the first stage, the magnitudes of the order parameters 459 r_1 and r_2 are practically constant. They undergo an abrupt 460 change during the second stage to return to a constant value 461 [Fig. 5(b)]. As the parameter α_2 increases, the duration of the 462 first stage decreases, and, hence, the period of oscillations in 463 r_1 and r_2 gradually decreases, causing the switching cyclops 464 state to eventually turn into a chaotically switching dynamical 465 pattern (not shown in Fig. 5). 466

We also observe a hybrid of the switching and breathing 467 cyclops states [Fig. 6(b)]. This hybrid state emerges when, 468 in addition to the external instability of one cluster, there 469 is an internal instability of the intercluster phase differences 470 x and y (Fig. 3). In terms of the eigenvalue spectrum, this 471 amounts to the presence of a pair of complex conjugate 472 eigenvalues $\lambda_{1,2}$ (corresponding to the internal instability) 473 and one real eigenvalue λ_1^{tran} lying to the right from the 474



FIG. 5. Switching cyclops state. (a) The colors depict the phase differences $\theta_k(t) - \theta_0(t)$. The strips with solid black borders indicate the reference solitary oscillator during the lifetime of a cyclops state configuration (the first stage). Note that clusters disintegrate to form a new cyclops state with a different solitary oscillator (the second stage). (b) The corresponding values of r_1 and r_2 . The gray fragments correspond to the zoomed-in insets (right panels). (c) The eigenvalues associated with the destabilized stationary cyclops state. Some eigenvalues are repeated. The round (triangular) labels correspond to the internal (transversal) stability. Note a positive real eigenvalue (red) corresponding to the loss of the transversal stability of the stationary cyclops state due to Statement 2. (d) Phase distributions θ_k corresponding to a death-birth process in which a cyclops state existing at $t = t_1$ disintegrates to form a new cyclops state at $t = t_4$ (see Supplemental Movie 2 [68] for the details of this dynamical evolution). Parameters N = 11, $\mu = 1.0$, $\varepsilon_1 = 1.0$, $\alpha_1 = 1.7$, $\varepsilon_2 = 0.08$, $\alpha_2 = 0.78$ correspond to the diamond label in Fig. 3(a).

imaginary axis [Fig. 6(c)]. We term this hybrid a switching-475 breathing cyclops state, which is effectively a switching 476 cyclops state, which, during its first stage, has oscillating inter-477 cluster phase differences x(t) and y(t). Accordingly, the order 478 parameter amplitudes r_1 and r_2 are time-periodic functions 479 [Fig. 6(b)]. Figure 6(d) and Supplemental Movie 3 [68] detail 480

the dynamical evolution of the switching-breathing cyclops 481 state.

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Breathing and switching cyclops states can also merge to 483 form another hybrid cyclops state, termed rotobreathing cy-484 clops states (Fig. 7) in the range of the second-harmonic phase 485 shift with $|\alpha_2| > \pi/2$ (Fig. 8, the pink regions). Rotobreathing 486



FIG. 6. Switching-breathing cyclops state. The notations are as in Fig. 5. One cluster of the breathing cyclops state [depicted in orange in panel (a)] eventually disintegrates, forming a reshuffled synchronous cluster and a new solitary oscillator. Note the weak internal and transversal instability of the destabilized stationary cyclops state due to the three eigenvalues with small positive real parts [red circles and nabla in panel (c)]. Supplemental Movie 3 [68] animates the sequence given in panel (d). Parameters N = 11, $\mu = 1.0$, $\varepsilon_1 = 1.0$, $\alpha_1 = 1.7$, $\varepsilon_2 = 0.0578$, $\alpha_2 = 0.78$ correspond to the circled times label in Fig. 3(a).



FIG. 7. Rotobreathing cyclops state. The notations are as in Fig. 5. From left to right: the relative phase between the first synchronous cluster and the sixth reference oscillator oscillates, whereas the phase of the second synchronous cluster passes zero and rotates until the clusters exchange their roles. Supplemental Movie 4 [68] details this process. Parameters are N = 11, $\mu = 1.0$, $\varepsilon_1 = 1.0$, $\alpha_1 = 1.7$, $\varepsilon_2 = 0.08$, $\alpha_2 = -2.0$.

cyclops states, or simply rotobreathers, are also characterized 487 by a two-stage repetitive process in which, during the first 488 stage, an intercluster phase difference between one cluster and 489 the solitary oscillator oscillates while the relative phase differ-490 ence of the other cluster rotates. The clusters exchange their 49 oscillatory and rotatory phase roles during the second stage. 492 Figure 7 and Supplemental Movie 4 [68] give the full details 493 of this two-stage process. Accordingly, the amplitudes of the 494 order parameters r_1 and r_2 exhibit large periodic oscillations 495 [Fig. 7(b)]. 496

Figure 8 demonstrates the prevalence of cyclops states of 497 various types. Remarkably, rotobreathers and breathing cy-498 clops states, induced by nonzero second-harmonic phase lag 499 α_2 in the region where full synchronization is unstable, act 500 as global attractors and emerge with a probability close to 501 1 [Figs. 8(b) and 8(c)]. Note that breathing and switching 502 cyclops states can also emerge with a relatively high proba-503 bility even when they coexist with presumably dominant full 504 synchronization when the overall coupling is attractive with 505 H'(0) > 0 [the region bounded by the black dashed vertical 506 lines in Fig. 8(b); these lines correspond to the solid circles on 507 the black dashed parabola in Fig. 8(a)]. 508

It is worth noting that the globally coupled network (1)509 with the first- and second-harmonic coupling admits any clus-510 ter partition. As a result, stationary cyclops states with a 511 solitary oscillator and nonequally sized coherent clusters exist 512 for even N. However, our extensive simulations suggest that 513 514 such stationary cyclops states are unstable for even N in the entire parameter range of ε_2 and α_2 considered in this paper. 515 Figure 9(a) supports this claim and demonstrates that, in the 516 attractive coupling case, two-cluster states, coexisting with 517 complete synchronization, are the only stable cluster patterns 518 in the network with N = 10. The same claim holds for the 519 repulsive coupling case, with the exception that complete syn-520 chronization is no longer stable so that the two-cluster states 521



FIG. 8. Stability and prevalence of cyclops states. (a) Stability diagram extending Fig. 3(a) to the full range of the phase lag parameter α_2 . The notations are similar to Fig. 3(a), with the addition of rotobreathers (pink). The shaded vertical strip corresponds to the parameter region of Fig. 3(a). Stationary cyclops states in the region CS are chosen as initial conditions and further continued by changing the parameter α_2 right and left from each point on the line $\alpha_2 = 0.0$ for each value of ε_2 . The initial conditions for the subsequent calculation are carried over from the final state of the preceding computation. The double-shaded areas (inclined stripes) indicate overlapping stability regions and correspond to the bistability of different cyclops state types. The two dash-dotted horizontal lines indicate the values of ε_2 used in panels (b) and (c). (b, c) Probability of cyclops states' emergence (all types). The number of trials is 1000. The initial phases are uniformly distributed in the segment $[-\pi, \pi]$, and the initial velocities are uniformly distributed in the segment [-1.0, 1.0]. The black dashed vertical lines in panel (b) indicate the stability boundary of full synchronization. In panel (c) full synchronization is unstable. Parameters are $N = 11, \mu = 1.0$, $\varepsilon_1 = 1.0, \alpha_1 = 1.7.$ (b) $\varepsilon_2 = 0.08,$ (c) $\varepsilon_2 = 0.05.$

become prevalent and coexist with four-cluster states with rotating intercluster differences for N = 10 [Fig. 9(c)]. Similarly, stationary cyclops states (4) emerge as the only stable patterns along with complete synchronization for N = 11 in the attractive coupling case [Fig. 9(b)], and breathing cyclops states with equally sized coherent clusters appear as global attractors in the repulsive coupling case [Fig. 9(d)].

Remarkably, this distinct behavior in the emergence of two-cluster and cyclops states in networks of even and odd sizes carries over to large networks. Figure 10(a) demonstrates 531



FIG. 9. The probability of occurrence of two-cluster (N = 10) and cyclops states (N = 11) under attractive and repulsive interactions. The histograms indicate the percentage occurrence of dynamical states in 1000 simulations with random initial conditions. The initial phases $\Theta_k(0)$ are uniformly distributed on $[-\pi, \pi]$; the initial velocities $\dot{\Theta}_k(0)$ are uniformly distributed on [-5.0, 5.0]. The snapshots correspond to the established dynamical states with the relative phases θ_k in the rotating frame as in (4). (a, b) Attractive coupling: complete synchronization is prevalent with a 72 % (N = 10) and 62% (N = 11) probability. It coexists for N = 10with a stationary two-cluster 5:5 state with an even partition of five oscillators in each cluster and a constant intercluster phase difference (a) and for N = 11 with a stationary cyclops state 5:1:5 (b). (c, d) Repulsive coupling. (c) N = 10: stationary two-cluster 5:5 states coexist with four-cluster states 4:4:1:1 with nonstationary intercluster phase differences. (d) N = 11: breathing cyclops states 5:1:5 with oscillating intercluster phase differences are global attractors emerging with a 100% probability. The black arrows indicate the directions of evolving intercluster differences. Parameters are $\mu = 1.0$, $\varepsilon_1 = 1.0$, $\alpha_1 = 1.7$; (a, b) $\varepsilon_2 = 0.12$, $\alpha_2 = 0.0$; (c, d) $\varepsilon_2 = 0.04$, $\alpha_2 = 0.6$.

that stationary cyclops states (4) are stable in the odd-sized 532 network with N = 101. Eliminating one oscillator from the 533 network turns the cyclops state into a two-cluster state with 534 equally sized clusters [Fig. 10(b)]. Removing an extra oscil-535 lator from this 100-node network transforms the two-cluster 536 state back to a stationary cyclops state [Fig. 10(c)]. We expect 537 this effect to persist in the thermodynamic limit of very large 538 Ν. 539

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VII. PERSISTENCE OF CYCLOPS STATES

In this section we demonstrate that cyclops states resist 541 intrinsic frequency detuning. We mismatch the intrinsic fre-542 quency ω by choosing the kth oscillator's frequency ω_k , k =543 $1, \ldots N$ from a uniform random distribution in the interval 544 $[\omega - \delta, \omega + \delta]$, where δ is a frequency detuning. We consider 545 the parameter region where stationary cyclops states are stable 546 [region CS in Fig. 3(a)]. Figure 11 demonstrates the persis-547 tence of three stationary cyclops states, each induced by a 548 549 particular intrinsic frequency distribution. Note that although the oscillators' phases within each synchronous cluster may 550 not perfectly align due to the frequency detuning, they remain 55 relatively close to each other (see Fig. 11). Additionally, the 552 established frequencies of all oscillators are the same. The sta-553 tionary cyclops state can lose the transversal stability similarly 554 to their counterparts from the identical oscillator case (note the 555 stationary cyclops state marked by the red labels in Fig. 11 that 556



FIG. 10. Odd vs even-sized large networks: the transition between cyclops and two-cluster states as N changes. The colors depict the relative phases θ_k , k = 1, ..., N. (a) A cyclops state with the three-cluster partition 50:1:50 in the 101-oscillator network. The cyclops state with small phase offsets, randomly chosen from a uniform distribution [-0.01; 0.01] was selected as the initial condition. The horizontal black stripe indicates the solitary oscillator with $\theta_{51} = 0$. (b) The emergence of a two-cluster state 50:50 in the 100-oscillator network. The cyclops state from (a) with one oscillator removed from the second cluster was selected as the initial condition. (c) Removing one oscillator from the two-cluster state in (b) induces a cyclops state 49:1:49 in the 99-oscillator network. (d) The corresponding snapshots of phases θ_k at several time instants: $t_1 = 1250, t_2 = 3750$, $t_3 = 5200, t_4 = 6250, t_5 = 11250$, and $t_6 = 13750$. The black arrow indicates the direction of the solitary oscillator's phase evolution. Parameters are $\mu = 1.0$, $\varepsilon_1 = 1.0$, $\alpha_1 = 3.1$, $\varepsilon_2 = 0.002$, $\alpha_2 = 0.2$.

turns into a switching cyclops state at $\delta = \delta_1$). Remarkably, the frequency detuning can also induce a bifurcation scenario for disintegrating stationary cyclops states via a saddle-node bifurcation at $\delta = \delta_2$ and $\delta = \delta_3$. We did not observe such a bifurcation route in our extensive simulations of system (1) with identical frequencies reported in Figs. 3–8.

VIII. CONCLUSIONS

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Building upon our recent study [50], this work has sig-564 nificantly advanced an understanding of rhythmogenesis in 565 Kuramoto networks of 2D phase oscillators with first-mode 566 and higher-mode coupling. A key focus of our work has been 567 on the constructive role of higher coupling modes in inducing 568 and stabilizing a unique class of dynamical states known as 569 cyclops states. These states, characterized by two coherent 570 clusters and a solitary oscillator resembling the Cyclops's eye, 571 represent a particular form of three-cluster generalized splay 572 states [49]. 573

Our initial findings in [50] revealed the unexpected result that adding the second or third harmonic to the Kuramoto coupling makes cyclops states global attractors, exhibiting remarkable stability over a substantial range of coupling's repulsion. This paper delved deeper into the dynamic repertoire of cyclops states, introducing and systematically analyzing 579



FIG. 11. Persistence of cyclops states in system (1) with mismatched frequencies ω_k distributed evenly over the interval [ω – $\delta, \omega + \delta$, where $\omega = 1.7$ and δ is a frequency detuning. Global maxima (circles) and minima (crosses) of order parameter r_2 for three cyclops states (red, orange, and cyan). The global maximum was determined in the time interval $5 \times 10^3 \le t \le 10^4$ following the transition process. A stationary cyclops state is indicated by overlapping circles and crosses of the same color, while separated circles and crosses denote a switching cyclops state. The first cyclops state (red) originates from the original intrinsic frequency distribution. Subsequently, modifying the distribution by exchanging the intrinsic frequency of the Mth solitary oscillator, ω_M , with ω_1 (or ω_{M+1}) induces the emergence of the second (or third) cyclops state, depicted by orange and cyan, respectively. As the parameter mismatch δ increases, the distribution of mismatched frequencies widens. To induce each of the three cyclops states as δ increases, a stationary cyclops state from the identical frequency case was used as the initial condition for $\delta = 0$. The final phase distribution at a given δ is then used as initial conditions for subsequent simulations at a higher value of δ . The values of $\delta < \delta_1$ preserve all three stable stationary cyclops states. Increasing $\delta > \delta_1$ destabilizes the first stationary cyclops state (red) and turns it into a switching cyclops state. Further increasing $\delta > \delta_2$ leads to disintegrating the second cyclops state (orange) at $\delta = \delta_2$. The third stationary cyclops state (cyan) persists to $\delta = \delta_3$. The cyclops states are found from direct numerical simulations of system (1) for three sets of natural frequency distributions ω_k with a continuous increase in δ from zero. The inset shows instantaneous phase distributions θ_k for the third cyclops state with nonidentical frequencies. Parameters are N = 11, $\mu = 1.0, \varepsilon_1 = 1.0, \alpha_1 = 1.8, \varepsilon_2 = 0.12$. The bifurcation parameter values are $\alpha_2 = 0.2$, $\delta_1 = 0.034$, $\delta_2 = 0.105$, $\delta_3 = 0.183$.

breathing and switching cyclops states and their hybrids, in-580 cluding switching-breathing cyclops states and rotobreathers. 58 Through rigorous analytical derivations and numerics, we 582 have identified conditions for the existence and stability of 583 stationary cyclops states, elucidating two distinct bifurcation 584 scenarios. In both scenarios, the second coupling harmonic 585 acts as a constructive agent, either inducing periodic oscil-586 lations in intercluster relative phase differences (breathing 587 cyclops states) or facilitating swift reconfigurations and tran-588 sitions (switching cyclops states). These dynamical patterns 589 can be viewed as nontrivial hybrids of solitary states [41-44], 590 generalized splay [49], clusters with breathing and rotatory 591 592 intercluster phase shifts [46,47], and intermittent [37] and switching chimeras [64]. In particular, switching cyclops 593 states unite the properties of blinking chimeras [64] and three-594 cluster states [47]. 595

⁵⁹⁶ Our extensive stability analysis has underscored the re-⁵⁹⁷ silience and dominance of breathing, rotobreathing, and ⁵⁹⁸ switching cyclops states across wide parameter ranges, in-⁵⁹⁹ cluding the case of the overall attractive, two-harmonic coupling. Importantly, we have showcased that the constructive influence of higher coupling harmonics is not limited to networks of identical oscillators, as cyclops states persist robustly in Kuramoto networks of nonidentical oscillators.

Importantly, our prior work [50] demonstrated the dynamic 604 equivalence of the 2D Kuramoto model with first- and second-605 harmonic coupling to a network of canonical theta neurons 606 with adaptive coupling. This equivalence also suggests the 607 widespread manifestation of breathing and switching cyclops 608 states in theta-neuron networks, underscoring our results' 609 broad applicability and significance in diverse physical and 610 biological networks. 611

While it is crucial to differentiate between the higher-order 612 harmonic coupling studied in this paper and the higher-613 order nonpairwise coupling [69-72], it is equally important 614 to recognize their possible interplay and the richness they 615 bring to the dynamics of networked systems. These con-616 cepts are not mutually exclusive; they can coexist, adding 617 layers of complexity and fostering a diverse range of emer-618 gent behaviors. Recent research [73,74] analyzed the intricate 619 interplay between pairwise first-order harmonic and nonpair-620 wise higher-order coupling in shaping collective dynamics in 621 Kuramoto networks. The incorporation of both higher-order 622 harmonics and nonpairwise interactions promises to induce 623 even richer emerging dynamics, including various forms of 624 cyclops states, and may pave the way for a more holistic 625 comprehension of complex networked systems. 626

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APPENDIX: MAXIMUM NUMBER OF STATIONARY CYCLOPS STATES

Here we provide the details for deriving an upper bound for the maximum number of stationary cyclops states with distinct x and y, given in Sec. III.

Finding all possible solutions of system (6) that determine the existence of stationary cyclops states is elusive due to its complexity, and the number of solutions can vary depending on the parameters. In particular, it prevents locating all solutions of the system (6) by their continuation with respect to the parameters. However, this computational problem can be simplified by the change of variables $u = e^{ix}$, $v = e^{iy}$ (|u| = 1, |v| = 1) that transforms the real-valued system (6) into the system of complex polynomial equations:

$$(1-u)\left\{uv\left[e^{i\alpha_1}\left(u+v+\frac{2uv}{N-1}\right)\right.\right.\right.\\\left.\left.\left.+e^{-i\alpha_1}v\left(u+v+\frac{2}{N-1}\right)\right]\right\}$$

$$+ \varepsilon_{2}(u+1) \left[e^{i\alpha_{2}} \left(u^{2} + v^{2} + \frac{2u^{2}v^{2}}{N-1} \right) + e^{-i\alpha_{2}}v^{2} \left(u^{2} + v^{2} + \frac{2}{N-1} \right) \right] \right\} = 0,$$

$$(1-v) \left\{ uv \left[e^{i\alpha_{1}} \left(u + v + \frac{2uv}{N-1} \right) + e^{-i\alpha_{1}}u \left(u + v + \frac{2}{N-1} \right) \right] + \varepsilon_{2}(v+1) \left[e^{i\alpha_{2}} \left(u^{2} + v^{2} + \frac{2u^{2}v^{2}}{N-1} \right) + e^{-i\alpha_{2}}u^{2} \left(u^{2} + v^{2} + \frac{2}{N-1} \right) \right] \right\} = 0.$$
(A1)

The analysis of system (A1) is more manageable, and the
maximum number of its solutions (the roots of the complex
polynomials) can be estimated by applying the classical Bernshtein theorem from algebra. To facilitate the reading, we list
this theorem below.

Theorem [Bernshtein, 1975] [67]. Let a system of n polynomials have a finite number of roots in $(C^*)^n$, where $C^* = C \setminus 0$. Then, the number of roots is bounded from above by the mixed volume P_k of their Newton polytopes (the convex hull of polynomial supports S_k).

Before applying the theorem to (A1), we get rid of the 659 factors (1 - u) and (1 - v) on the right-hand side of (A1) 660 since we are interested only in solutions $u, v \neq 1$. By do-661 ing so, we have excluded the solutions that correspond to 662 a one-cluster solution and two-cluster solutions of the form 663 (N-1)/2:(N+1)/2. It is worth mentioning that, in contrast 664 to its real-valued counterpart (6), the complex polynomials 665 may have either nonphysical solutions with $|u| \neq 1$ or $|v| \neq 1$ 666 1, or solutions that do not correspond to stationary cyclops 667 states. The latter solutions with |u| = 1 and |v| = 1, include 668

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FIG. 12. The supports S_1 , S_2 (black dots) and the corresponding Newton polytopes P_1 , P_2 (shaded regions) of (a) the first and (b) second polynomials of system (A1). (c) The Minkowski sum $P_1 \oplus P_2$.

a two-cluster N - 1: 1 solitary state, corresponding to u = v, i.e., x = y.

The support of a polynomial $\sum_{j} \sum_{l} a_{jl} u^{j} v^{l}$ is the set of exponents (j, l). Thus, the supports S_{1} and S_{2} of the resulting polynomials (degrees u and v in each of the first and second equations terms) have the form $S_{1} =$ $\{(0, 2); (0, 4); (1, 2); (1, 3); (1, 4); (2, 0); (2, 1); (2, 2); (3, 0);$ $(3, 2)\}, S_{2} = \{(0, 2); (0, 3); (1, 2); (2, 0); (2, 1); (2, 2); (2, 3);$ $(3, 1); (4, 0); (4, 1)\}.$

Consider the mixed volume of Newtonian polytopes P_1 and 678 $P_2: M(P_1, P_2) = \operatorname{vol}_2(P_1 \oplus P_2) - \operatorname{vol}_2(P_1) - \operatorname{vol}_2(P_2)$. As we 679 need to handle only two equations, determining the mixed 680 volume of the two Newtonian polytopes is straightforward 681 and amounts to computing the area of the shaded regions 682 in Fig. 12. This yields the following values: $vol_2(P_1) = 8$, 683 $vol_2(P_2) = 8$, $vol_2(P_1 \oplus P_2) = 33$ (see Fig. 12). Therefore, 684 $M(P_1, P_2) = 17$. Excluding the nonphysical solutions and so-685 lutions corresponding to noncyclops regimes from the sets of roots in (A1), we can always find the number of cyclops 687 modes in the system (1), which is limited to 16 cyclops states. 688

Our numerical search for the roots of polynomials (A1) 689 was performed using the NSolve function of Wolfram Math-690 ematica. This search found 17 roots almost everywhere in 691 the considered broad parameter regions, suggesting that our 692 analysis effectively identified all possible solutions of (A1) 693 and, therefore, all possible stationary states cyclops, identified 694 from the 17 solutions by excluding the nonphysical solutions 695 $(|u| \neq 1 \text{ or } |v| \neq 1)$ and noncyclops states (u = 1 or v = 1 or v = 1)696 u = v). 697

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