



# Reinforcement learning facilitates an optimal interaction intensity for cooperation



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## ABSTRACT

Our social interactions vary over time and they depend on various factors that determine our preferences and goals, both in personal and professional terms. Researches have shown that this plays an important role in promoting cooperation and prosocial behavior in general. Indeed, it is natural to assume that ties among cooperators would become stronger over time, while ties with defectors (non-cooperators) would eventually be severed. Here we introduce reinforcement learning as a determinant of adaptive interaction intensity in social dilemmas and study how this translates into the structure of the social network and its propensity to sustain cooperation. We merge the iterated prisoner's dilemma game with the Bush–Mosteller reinforcement learning model and show that there exists a moderate switching dynamics of the interaction intensity that is optimal for the evolution of cooperation. Besides, the results of Monte Carlo simulations are further supported by the calculations of dynamical pair approximation. These observations show that reinforcement learning is sufficient for the emergence of optimal social interaction patterns that facilitate cooperation. This in turn supports the social capital hypothesis with a minimal set of assumptions that guide the self-organization of our social fabric.

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## 1. Introduction

In nature and society, there are various kinds of interactions to support the stability of biological, social, and technological systems. On the one hand, through the interactions, agents could compete for resources and pursue the goal of maximizing their interests, which is consistent with the evolution theory in terms of the survival of the fittest. On the other hand, however, there extensively exists cooperative interaction behavior between agents, which could not be well explained by the evolution theory. Therefore, understanding the emergence of cooperation becomes a key to solving some social dilemmas [1,2]. Aim to this issue, the evolutionary game theory has provided an effective framework because it can well capture the relationship between interactions and the evolutionary dynamics of cooperation [3–5].

With the evolutionary game theory, the prisoner's dilemma game (PDG) is frequently adopted as a standard framework to reflect the interest conflicts between agents and groups [6–8]. In this basic model, two agents need to choose a strategy from cooperation and defection at the same time. Thus, there are four different strategy combinations, wherein mutual cooperation is the best case for the collective benefit, while defection is the best decision to maximize agent's interest regardless of the strategy of the opposite. In traditional cases, the iterated PDG is usually used in a well-mixed infinite population, where agents will interact with the rest in an equal way. However, it is difficult for cooperation to survive under such a scenario. Aim to resolve this issue, hundreds of scenarios have been proposed both theoretically and experimentally. Typically, Nowak summarizes five rules for the evolution of cooperation, including the kin selection, direct and indirect reciprocity, group selection, and network reciprocity [9–11]. Besides, many other mechanisms are demonstrated to promote the evolution of cooperation, for instance, coevolution of strategy and structure [12], and mobility [13]. In particular, network reciprocity indicates that cooperators will resist the invasion of defectors by forming

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spatial clusters in networks [14–16]. Along this line, one can study agents' communications and their relationship formed via daily routines on a network, in which vertexes represent agents and edges reflect the relationship between them [17–20]. Taking advantage of networks' property, the research of cooperation on various networks arises to figure out the topological effects on cooperation [21–27]. Besides, some other mechanisms nourishing cooperation in structured population have also been put forward [28–39].

In spite of great progress, the vast majority of posting works are based on a prior assumption that agents interact with all the nearest neighbors. In other words, agents are supposed to have extremely strong interaction intensities to interact with their nearest neighbors during a whole evolutionary process, which is inconsistent with realistic observations. In reality, agents may not necessarily interact with all their nearest neighbors, and the interaction with neighbors will also change under different conditions. In this sense, the absolute interaction relationship between agents can be replaced by a stochastic probability, namely, the interaction intensity, which leads to diverse interaction patterns for agents, and thus various evolutionary dynamics [40–43]. Although it has been demonstrated that adopting such interaction probability in the iterated PDG is able to improve cooperation in structured populations, these works mainly focused on a constant interaction probability. It is apparent that the interaction intensity of an agent will change according to the specific environment. Thus, the adaptive interaction intensity needs to be studied, with which agents could change their interactions adaptively.

In recent years, the reinforcement learning (RL) methods, rooted in psychology and neuroscience, have become another approach to understanding the cooperation in evolutionary processes [44,45]. Ordinary RL algorithms contain the Monte Carlo method [46], Sarsa [47], dynamic programming [48], temporal differences method [49], Q-learning [50]. The core idea of RL methods is that the action an agent takes in the next step is affected by the reward, which is the feedback of the environment to its current state. For example, positive feedback, like a high reward, will lead to the enhancement of its current action when encountering the same state the next time, and vice versa [51,52]. Motivated by this mechanism, works that combine reinforcement learning and evolutionary game for solving social dilemmas have attracted much research interest [53,54]. These studies have established a reasonable framework, under which the strategy (cooperation or defection), is similar to the action in RL, and the payoff can be regarded as the reward. The Bush-Mostelle (BM) model, another classic RL algorithm, enables agents to adaptively change their current actions according to their rewards from the current states, which is helpful to depict the self-regarding process [55,56]. Under the BM model, the larger the reward exceeds an agent's aspiration, the more possibly the agent will continue adopting its current action. Based on this, we could study the dynamical process of the evolutionary game with the adaptive interaction intensity. Specifically, agents could strengthen (or weaken) the current strategies (interaction intensities) when receiving a higher (or lower) payoff than aspiration.

In this paper, we focus on exploring the effect of the adaptive interaction intensity on cooperation in both structured and well-mixed populations. We also investigate the co-evolution progress where the interaction intensity of agents changes as their strategies update. The main contributions of this paper can be summarized as i) We propose an adaptive interaction intensity model to optimize population benefits and inspire agent behaviors. ii) We construct a valid dynamical pair approximation framework for adaptive interaction intensity, which fills the gap in this direction. iii) We demonstrate the self-organization behavior pattern in the interaction fabric, which reflects the internal factors of social har-

mony. In the following part, we first show that the introduction of adaptive interaction intensity improves the cooperative level of iterated PDG on square lattices. We analyze the influence of the adaptive interaction intensity on cooperation from the view of co-evolution processes and microscopic mechanisms. Furthermore, we use the pair approximation method to obtain an analytical result. Finally, we validate the robustness of the proposed mechanism on a well-mixed population.

## 2. Preliminaries

Game theory provides a framework to investigate human behaviors in interactions. In a two-agent two-strategy game, each agent has two optional strategies, cooperation (C) or defection (D). After the interaction, agents obtain payoffs according to the payoff matrix:

$$\begin{matrix} & C & D \\ \begin{matrix} C \\ D \end{matrix} & \begin{pmatrix} R & S \\ T & P \end{pmatrix} \end{matrix}, \quad (1)$$

where  $R, P, S$ , and  $T$  denote the reward for mutual cooperation, punishment for mutual defections, sucker's payoff, and the defective temptation when one agent chooses cooperation and the other chooses defection, respectively. These parameters decide the game type. If the parameters satisfy the condition  $T > R > P > S$ , it is a prisoner's dilemma game; if the parameters satisfy the condition  $T > R > S > P$ , it is a snowdrift game; if the parameters satisfy the condition  $R > T > P > S$ , it is a stag hunt game; if the parameters satisfy the condition  $R > T$  and  $S > P$ , it is a coordinate game.

Every agent in a game pursues the maximum of its own payoff. Nash equilibrium refers to a strategy combination that contains the optimal strategies of two agents. The Nash equilibrium is  $(D, D)$  in prisoner's dilemma game,  $(C, D)$  and  $(D, C)$  in snowdrift game,  $(C, C)$  and  $(D, D)$  in stag hunt game, and  $(C, C)$  in coordinate game.

The Bush-Mostelle (BM) method, one of the classic reinforcement learning algorithms, describes the self-regarding process based on the current reward and action. It describes the probability that an agent takes the specific action next time. Assume agent's action (C or D) and reward in time  $t$  as  $a_t$  and  $r_t$ , the probability  $p_{t+1}$  of taking action C in  $t + 1$  is described as:

$$p_{t+1} = \begin{cases} p_t + (1 - p_t)s_t, & a_t = C, s_t \geq 0 \\ p_t + p_t s_t, & a_t = C, s_t < 0 \\ p_t - p_t s_t, & a_t = D, s_t \geq 0 \\ p_t - (1 - p_t)s_t, & a_t = D, s_t < 0 \end{cases}, \quad (2)$$

where  $s_t$  ( $-1 < s_t < 1$ ) is the stimulus in  $t$  and is defined as:

$$s_t = \tanh[\beta(r_t - A)], \quad (3)$$

where  $A$  is the aspiration level and  $\beta$  ( $\beta > 0$ ) is the sensitivity to  $r_t - A$ . Positive  $s_t$  will increase the probability of taking action C and reduce the probability of taking action D, while negative  $s_t$  will reduce the probability of taking action C and increase the probability of taking action D. Specially, when  $t = 0$ ,  $p_t$  obeys the uniform density on  $[0, 1]$ , independently for different agents.

## 3. Model and method

We consider evolutionary games on square lattices with periodic boundaries and Von Neumann neighborhood, where each vertex denotes an agent who can interact with four nearest neighbors  $\Omega$  along edges. Agents play the pairwise prisoner's dilemma game with their neighbors. To simplify yet without losing generality, we

adopt the weak prisoner's dilemma game as our main model in the following discussions, where  $S = P = 0, R = 1$  and  $T = b$  ( $b \geq 1$ ).

Then, we define  $p_{x \rightarrow y}$  as the willingness that agent  $x$  would like to interact with neighbor  $y$ . In our model, we assume the willingness of an agent to interact with different neighbors is independent, and so does the willingness of different agents. Thus, the interaction intensity, i.e., the probability that agent  $x$  and her neighbor  $y$  would successfully interact, can be denoted as:

$$p_{xy} = p_{x \rightarrow y} \times p_{y \rightarrow x}. \quad (4)$$

Based on this definition,  $p_{xy}$ , the interaction intensity between  $x$  and  $y$ , depends on the interacting willingness of  $x$  towards  $y$ ,  $p_{x \rightarrow y}$ , and the interacting willingness of  $y$  towards  $x$ ,  $p_{y \rightarrow x}$ . Note  $p_{xy} = p_{yx}$  because of the symmetry. We assume that each pair of agents interacts following the interaction intensity with probability  $1 - \epsilon$ , where  $\epsilon$  denotes the probability that agents would randomly choose to interact or not. (In this paper, the parameter  $\epsilon$  will be set as 0.1 for all simulations.)

For convenience, we divide the four nearest neighbors into two sets based on whether the interaction happens, including the interactive neighbors  $\Omega_i$  and non-interactive neighbors  $\Omega_n$ . In each Monte Carlo step, agent  $x$  plays the game with her interactive neighbors and obtains an accumulated payoff  $\Pi_x$  at the step  $t$ . With all these definitions, we can then describe the dynamics of our model as two processes. First, agent  $x$  updates strategy according to the "imitation rule", where uses Fermi function is a well-known method to calculate the imitation probability:

$$f_{y \rightarrow x} = \frac{1}{1 + \exp[(\Pi_x - \Pi_y)/K]}, \quad (5)$$

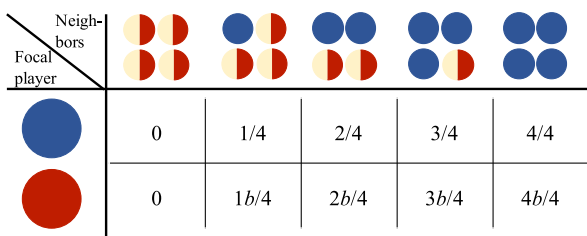
where  $\Pi_y$  is the accumulated payoff of the randomly selected neighbor  $y$  and the noise  $K$  is set to be 0.1. Then, the interactive willingness of agent  $x$  towards  $y$  will adaptively change according to the BM model:

$$p_{x \rightarrow y}(t+1) = \begin{cases} p_{x \rightarrow y}(t) + (1 - p_{x \rightarrow y}(t))s_{xy}(t), & a_{xy}(t) = 1, s_{xy}(t) \geq 0 \\ p_{x \rightarrow y}(t) + p_{x \rightarrow y}(t)s_{xy}(t), & a_{xy}(t) = 1, s_{xy}(t) < 0 \\ p_{x \rightarrow y}(t) - p_{x \rightarrow y}(t)s_{xy}(t), & a_{xy}(t) = 0, s_{xy}(t) \geq 0 \\ p_{x \rightarrow y}(t) - (1 - p_{x \rightarrow y}(t))s_{xy}(t), & a_{xy}(t) = 0, s_{xy}(t) < 0 \end{cases}, \quad (6)$$

where  $a_{xy} = 1$  and  $a_{xy} = 0$  represent interaction and non-interaction between agent  $x$  and  $y$  at the step  $t$ , respectively, and  $s_{xy}(t)$  denotes an stimulus that can be described as:

$$s_{xy}(t) = \tanh[\beta(r_{xy}(t) - A_x(t))], \quad (7)$$

where  $r_{xy}(t)$  is the payoff of agent  $x$  obtained from neighbor  $y$ ,  $A_x(t) = \Pi_x/4$  is the aspiration level averaged over the four nearest neighbors, and  $\beta$  represents the sensitivity of  $s_{xy}(t)$  to  $r_{xy}(t) - A_x(t)$ . Finally, we can give detailed aspirations and payoffs of focal cooperators and defectors in different configurations in Fig. 1. Note that



**Fig. 1.** Aspirations of cooperators and defectors. Aspiration is defined as the average payoffs of agents. Blue and red circles represent cooperators and defectors, respectively, and circles colored with both yellow and red represent non-interaction or defectors. Agents can only obtain payoffs by interaction with cooperators.

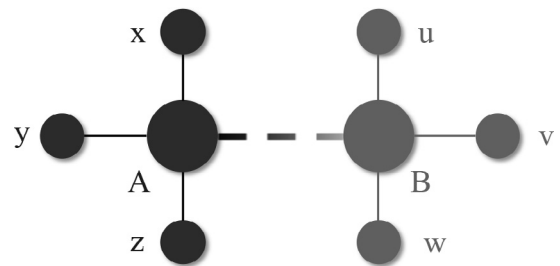
$r_{xy}(t) = 0$  when there is no interaction between two agents or the neighbor (opposite)  $y$  adopts the defection strategy, which means agents can only obtain a payoff from interaction with cooperative neighbors. In addition, when  $a_{xy}(t) = 0, r_{xy} = 0$  and  $s_{xy}$  is either less than or equal to zero. However, the BM model normally contains four scenarios. Thus, without loss of generality, we still describe the whole four scenarios in Eq. (7).

Initially, each agent randomly chooses cooperation or defection with the same probability and her willingness to interact with each nearest neighbor subjects to a uniform distribution  $[0, 1]$ . To avoid the willingness being too small or large during the evolutionary process (the evolution process may be frozen), we set the minimum of the willingness as 0.01 and the maximum as 1. We conduct all the Monte-Carlo simulations (MCS) on a square lattice with  $L = 300$  to obtain the evolution data. We take an average of the last 5,000 steps (with a total being  $3 \times 10^4$ ) to represent the steady states, and average over ten independent simulation runs for a fixed set of parameter values.

#### 4. Pair approximation

As shown in Fig. 2, a randomly selected 2-site configuration of a square lattice is the basic structure of the pair approximation method and it can contribute to the pair evolutionary dynamics, where the focal agent  $B$  is the randomly chosen nearest neighbor of the focal agent  $A$  in the strategy updating process. The payoff of  $A$  ( $\Pi_A$ ) and  $B$  ( $\Pi_B$ ) are determined by their nearest neighbors  $x, y, z, B$  and  $u, v, w, A$ , respectively. For example, if  $A$  adopts cooperation,  $B$  adopts defection, and  $A$  learns the strategy of  $B$ , as a result, the pair  $c - c$  and  $c - d$  will decrease while the pair  $d - c$  and  $d - d$  will increase.

In our model, the accumulated payoff of the focal agent depends on the actual interactive neighbors. In order to calculate the payoffs of focal agents, we need to know the interaction intensity of agents toward cooperators (interaction intensity towards defectors is omitted because it produces no payoffs). For a focal cooperator agent  $A$ , her neighbors are divided into interactive neighbors and non-interactive neighbors according to whether the interaction happens. Then, the overall  $c - c$  link denotes the link between  $A$  and all her cooperator neighbors, and the actual  $c - c$  link denotes the link between  $A$  and her interactive cooperator neighbors. Thus, for the whole population, we use the overall  $c - c$  link  $n_{c,c}(t)$  to denote links between the focal cooperator agents and their cooperator neighbors regardless of the interaction intensity, and the actual  $c - c$  link  $n_{c,c'}(t)$  to denote links between the focal cooperator agents and their interactive cooperator neighbors. In other words,  $n_{c,c'}(t)$  is the subset of  $n_{c,c}(t)$ . In this paper, these parameters are obtained from the Monte-Carlo simulation results. Specifically, the ratio of actual interacting  $c - c$  links  $n_{c,c'}(t)$  to the overall  $c - c$  links  $n_{c,c}(t)$  in the last  $m$  steps of the total  $M$  steps is used to repre-



**Fig. 2.** 2-site configuration. A randomly selected 2-site configuration in square lattice, where  $A$  and  $B$  are two focal agents with nearest neighbors  $x, y, z, B$  and  $u, v, w, A$ , respectively. Agent  $A$  learns the strategy of agent  $B$  according to the "imitation rule".

sent the average interaction intensity of cooperators towards cooperators, which can be denoted as  $\bar{I}_{c,c}$ :

$$\bar{I}_{c,c} = \frac{\sum_{t=M-m}^M n_{c,c}(t)}{\sum_{t=M-m}^M n_{c,c}(t)}. \quad (8)$$

In the same way, the average interaction intensity of defectors towards cooperators  $\bar{I}_{d,c}$  can be written as:

$$\bar{I}_{d,c} = \frac{\sum_{t=M-m}^M n_{d,c}(t)}{\sum_{t=M-m}^M n_{d,c}(t)}. \quad (9)$$

For a given 2-site configuration in which  $A$  is a cooperator and  $B$  is a defector, the focal agent  $A$  accumulates a payoff:

$$\Pi_c(x, y, z, \bar{I}_{c,c}) = \bar{I}_{c,c} \times n_c(x, y, z) \times 1, \quad (10)$$

and the focal agent  $B$  accumulates a payoff:

$$\Pi_d(u, v, w, \bar{I}_{d,c}) = \bar{I}_{d,c} \times n_c(u, v, w) \times b, \quad (11)$$

where  $n_c(x, y, z)$  and  $n_c(u, v, w)$  denote the number of cooperators among neighbors  $x, y, z$  and  $u, v, w$  of focal agent  $A$  and  $B$ , respectively. In a given configuration, when  $A$  is a defector and  $B$  is a cooperator, the focal agent  $A$ 's payoff can be calculated similarly.

The pair evolution with interaction intensity is determined by:

$$\begin{aligned} \dot{p}_{c,c} = & \sum_{x,y,z} [n_c(x, y, z) + 1] p_{d,x} p_{d,y} p_{d,z} \sum_{u,v,w} p_{c,u} p_{c,v} p_{c,w} f_d(x, y, z, \bar{I}_{d,c}) - c(u, v, w, \bar{I}_{c,c}) \\ & + \sum_{x,y,z} [-n_c(x, y, z)] p_{c,x} p_{c,y} p_{c,z} \sum_{u,v,w} p_{d,u} p_{d,v} p_{d,w} f_c(x, y, z, \bar{I}_{c,c}) - d(u, v, w, \bar{I}_{d,c}), \end{aligned} \quad (12)$$

$$\begin{aligned} \dot{p}_{c,d} = & \sum_{x,y,z} [1 - n_c(x, y, z) + 1] p_{d,x} p_{d,y} p_{d,z} \sum_{u,v,w} p_{c,u} p_{c,v} p_{c,w} f_d(x, y, z, \bar{I}_{d,c}) - c(u, v, w, \bar{I}_{c,c}) \\ & + \sum_{x,y,z} [n_c(x, y, z) - 2] p_{c,x} p_{c,y} p_{c,z} \sum_{u,v,w} p_{d,u} p_{d,v} p_{d,w} f_c(x, y, z, \bar{I}_{c,c}) - d(u, v, w, \bar{I}_{d,c}), \end{aligned} \quad (13)$$

where  $n_c(x, y, z)$  denotes the number of cooperators among neighbors  $x, y, z$  of the focal agent. Inserting Eq. (11) and (12),  $f_{* \rightarrow *}$  subjects to the Fermi function as described in Eq. (6)

$$\begin{aligned} f_d(x, y, z, \bar{I}_{d,c}) - c(u, v, w, \bar{I}_{c,c}) &= \frac{1}{1 + \exp\left[\left(\Pi_d(x, y, z, \bar{I}_{d,c}) - \Pi_c(u, v, w, \bar{I}_{c,c})\right)/K\right]} \\ f_c(x, y, z, \bar{I}_{c,c}) - d(u, v, w, \bar{I}_{d,c}) &= \frac{1}{1 + \exp\left[\left(\Pi_c(x, y, z, \bar{I}_{c,c}) - \Pi_d(u, v, w, \bar{I}_{d,c})\right)/K\right]} \end{aligned} \quad (14)$$

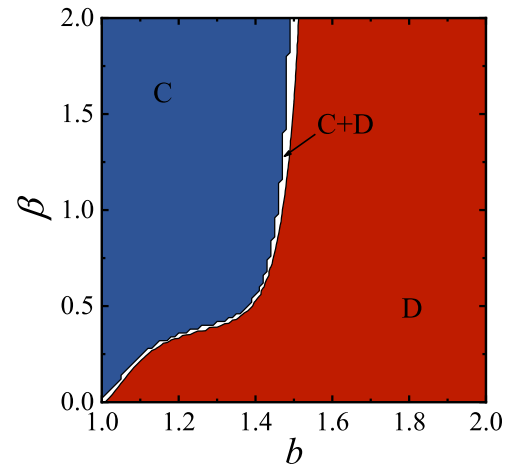
And  $p_{c,c}$  denotes the frequency of overall  $c-c$  links in the population,  $p_{c,d}$  denotes the frequency of overall  $c-d$  links in the population,  $p_{d,c}$  denotes the frequency of overall  $d-c$  links in the population, and  $p_{d,d}$  denotes the frequency of overall  $d-d$  links in the population. Note that we have omitted the multiplier factor  $\frac{2p_{c,d}}{p_c^2 p_d^2}$  because it has no influence on the equilibrium. One can also see that  $p_{c,d} = p_{d,c}$  and  $p_{c,c} + p_{c,d} + p_{d,c} + p_{d,d} = 1$  because of the symmetry and the natural constraint. Then, the frequency of cooperation can be acquired as  $f_c = p_{c,c} + p_{c,d}$ . Apparently, when the interaction intensity  $\bar{I}_{* \rightarrow *}$  equals to 1, it becomes the same case as the traditional pair approximation method. In other words, our model makes an extension of the previous work in terms of the evolutionary process and thus provides a more comprehensive framework [57].

## 5. Results

In weak prisoner's dilemma game, the defective temptation  $b$  is a pivotal parameter that influences the frequency of cooperation, while in the BM model,  $\beta$  plays an important role in deciding the stimulus, which is related to the dynamics of interaction intensity. So we start by exploring how the frequency of cooperation varies as a function of the parameter  $b$  and  $\beta$ . As shown in Fig. 3, obviously, cooperation is promoted by the introduction of the adaptive interaction intensity when compared to the traditional case (i.e.  $\beta = 0$ ). That is, the critical values at which cooperation becomes extinct are enlarged. In particular, cooperation is significantly enhanced when  $\beta$  is large. For instance, when  $\beta = 2$ , defectors cannot dominate the population until  $b$  is larger than 1.5. While for small  $\beta$ , the enhancement of cooperation is relatively slight. For example, when  $\beta = 0.1$ , cooperators can barely survive when  $b$  is around 1.05. In addition, it is worth noticing that there is a sharp transition from the full cooperation phase to the full defection phase in the population. Given a certain  $\beta$ , as  $b$  increases, cooperators first dominate the population (marked as the blue area in Fig. 3), then transiently co-exist with defectors (marked as the narrow white area), and finally die out with the dominance of defectors (marked as the red area).

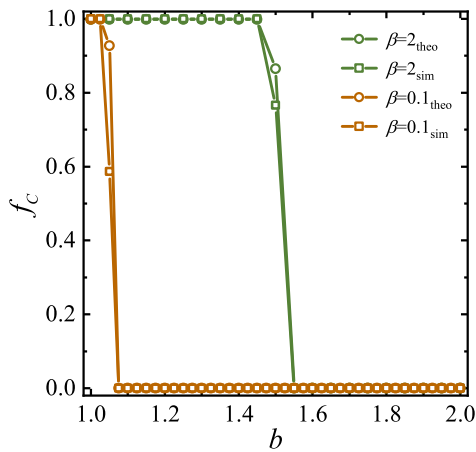
We then adopt the extended pair approximation method to theoretically demonstrate the promotion of cooperation and the sharp transition of the cooperation frequency. Take the cases of  $\beta = 0.1$  and  $\beta = 2$ , where cooperation is fragile and competitive in face of the invasion of defection, as an example. In Fig. 4, we plot the cooperation frequency  $f_c$  as a function of  $b$  based on both simulation and theoretical results. It shows the same transition trends in both simulation and theoretical analysis, although there exists little difference in the specific values since the evolutionary process on lattice networks is very complicated and often beyond the consideration of the pair approximation method. Comparison between simulation and theoretical results also confirms that the adaptive interaction intensity has a beneficial effect on cooperation. Namely, large  $\beta$  extends the range of  $b$  where cooperators survive.

To evaluate the above-observed phenomena, Figs. 5 ( $\beta=2$ ) and 6 ( $\beta=0.1$ ) show the distribution of interactive neighbors and strategies as the MCS step increases. Several characteristic spatial pat-



**Fig. 3.** Frequency of cooperation in  $b-\beta$  panel. Steady state frequency of cooperation is plotted as a function of  $b$  and  $\beta$ . The panel is separated into three phases. Blue, red and white represent full cooperation phase, full defection phase and the mixed strategy phase, respectively. We obtain the results with a random distribution for strategies.



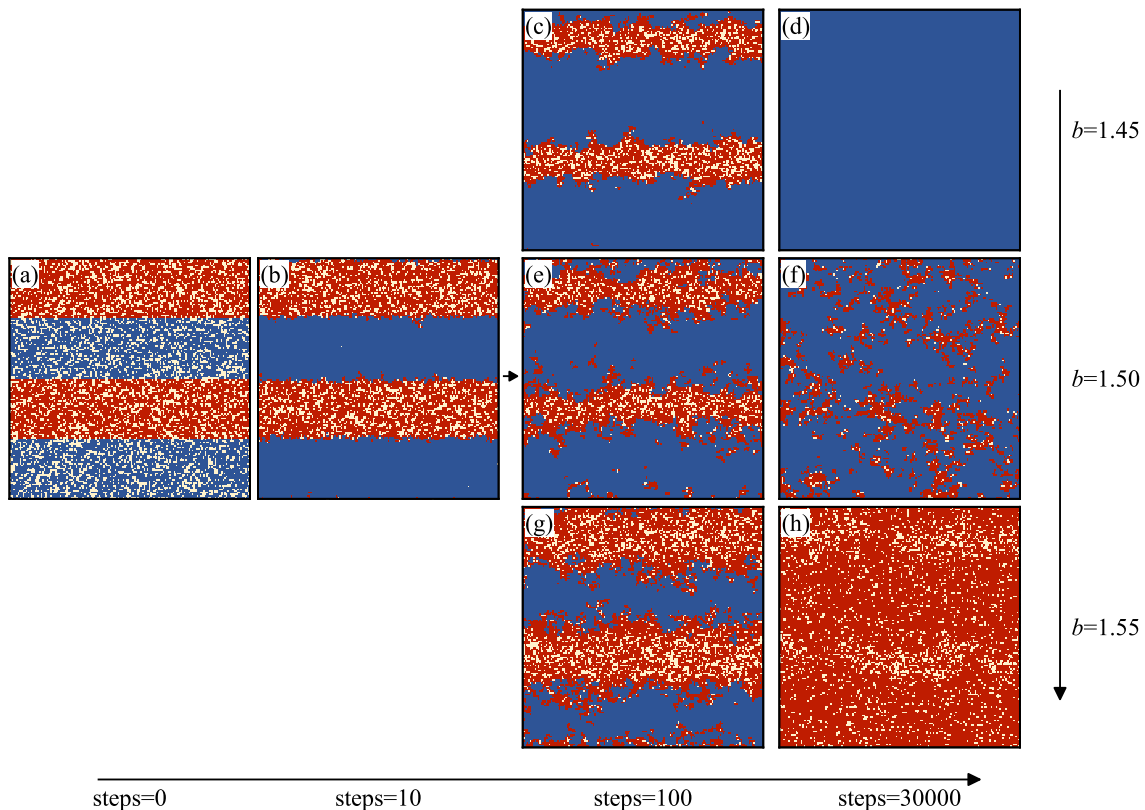


**Fig. 4.** Frequency of cooperation as a function of  $b$  with various  $\beta$ . Squares and circles represent the simulation and theoretical results, respectively. Orange and green represent results obtained with  $\beta = 0.1$  and  $\beta = 2$ , respectively. We obtain the results with a random distribution for the strategy in simulations.

terns of the entire evolutionary process with a prepared initial distribution are used to demonstrate the microscopic evolutionary dynamics under the influence of adaptive interaction intensity. There are three types of agents on the network, i) cooperator (blue) with at least one interactive neighbor; ii) defector (red) with at least one interactive neighbor; iii) isolated agent (yellow) with no interactive neighbors. As shown in Figs. 5(a) and 6(a), the initially prepared distribution divides the whole population into cooperative and defective areas. Then, because of the randomly ini-

tialized interaction willingness of agents, there are scattered yellow spots in blue and red stripes. In addition, since the early evolution (the first two columns) of different  $b$  seems to be similar, only some representative snapshots for all cases are plotted. In particular, we can observe the enduring (END) period where cooperation endures the invasion of defection and the expanding (EXD) period where cooperation spreads [58,59].

For the case of  $\beta = 2$ , isolated agents surrounded by cooperators disappear very quickly in the early stage, which means they have built at least one interaction with their cooperative neighbors. However, in defective areas, isolated agents hardly establish interaction with neighbors. In a word, agents are more willing to interact with cooperators than defectors. This leads to a strong interaction intensity within the former and a weak interaction intensity within the latter. As evolution proceeds, there will be different snapshots for different cases of  $b$ . When  $b = 1.45$ , the EXP period that features the expanding of cooperation shows up, and cooperation finally dominates the population. While the END period that features the endurance of cooperation is popular when  $b = 1.55$  [60].  $b = 1.5$  leads to the coexistence of cooperation and defection in the population. Of particular interest, the invasion of defection always brings isolated agents, which means that agents prefer to maintain interaction with cooperative neighbors instead of defective neighbors. In addition, agents are motivated to build new interactions with the cooperative neighbors and cut off the interaction with the defective neighbors. Therefore, during the evolution process, agents will dynamically adjust their interaction relationship with neighbors and gradually reach an optimal interaction intensity. Similar evolutionary patterns are reflected in the case of  $\beta = 0.1$ . But there exists some difference. Specifically, the evolutionary process is slower in the case of  $\beta = 0.1$ . For example,

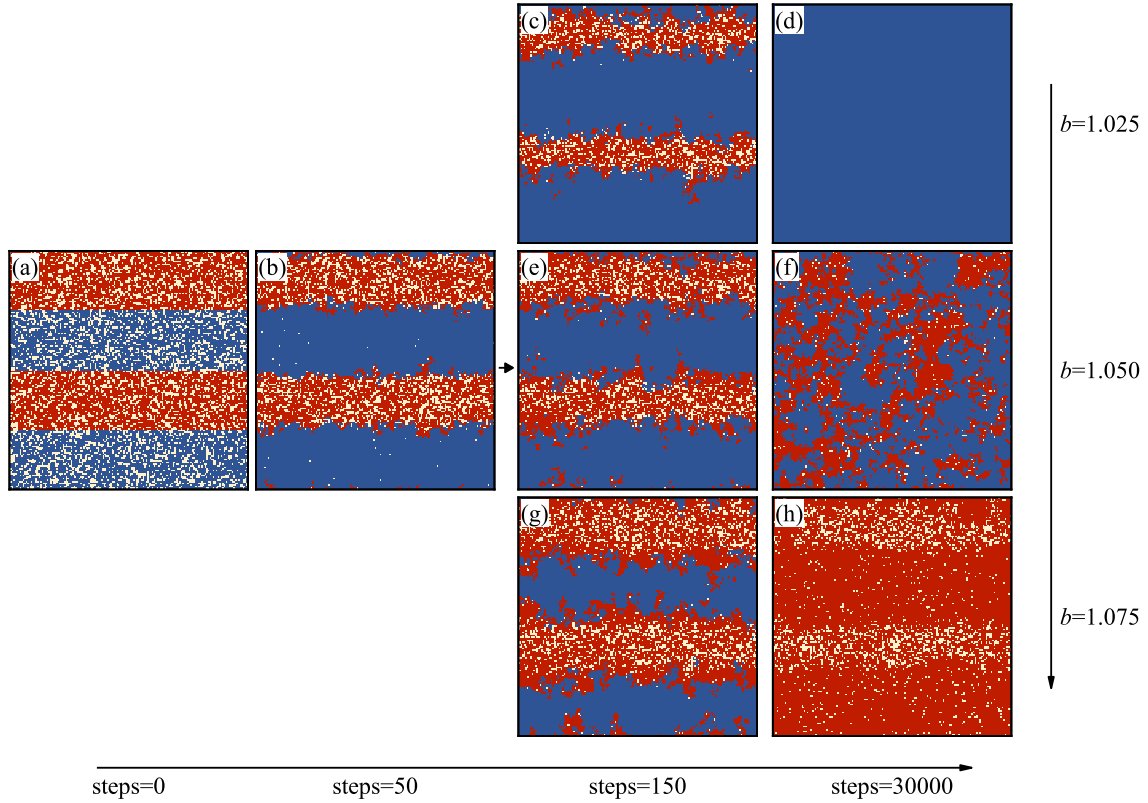


**Fig. 5.** Snapshots of the strategy distribution in evolutionary process. Snapshots are taken at step 0, 10, 100, and 30000, respectively with  $b = 1.45$ ,  $b = 1.5$  and  $b = 1.55$ . The layout follows the direction of arrows. First and second columns show the representative initial evolution process when  $\beta = 2$  for different  $b$ . Blue, red, and yellow represent cooperators, defectors, and isolated agents, respectively.

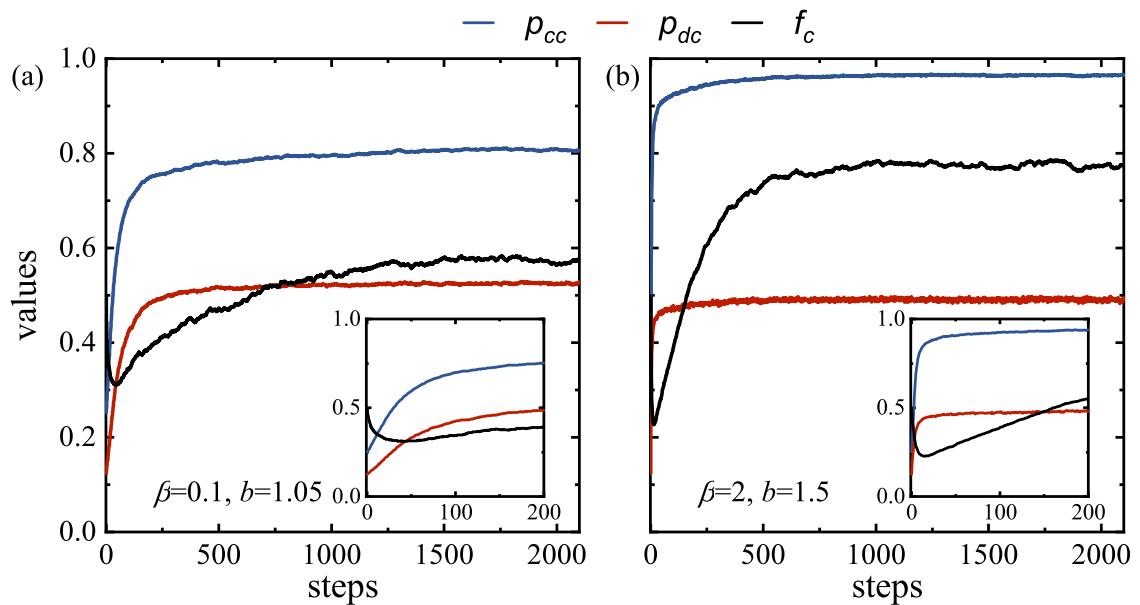
from Fig. 6 (a) to (b), isolated agents surrounded by cooperators need more time to build interaction with neighbors. Furthermore, the invasion of defection causes less emergence of isolated agents, which means the behavior of cutting off the interaction with defectors becomes slower, too. These results imply that a small  $\beta$  will

slow down the evolution process for agents to obtain the optimal interaction intensity.

To further explore the impact of the interaction intensity on the evolutionary process, Fig. 7 presents the evolution characteristic of cooperation and the interaction intensity. For different  $\beta$ , it is clear



**Fig. 6.** Snapshots of the strategy distribution in evolutionary process. Snapshots are taken at step 0, 10, 150, and 30000, respectively with  $b = 1.025$ ,  $b = 1.05$  and  $b = 1.075$ . The layout follows the direction of arrows. First and second columns show the representative initial evolution process when  $\beta = 0.1$  for different  $b$ . Blue, red, and yellow represent cooperators, defectors, and isolated agents, respectively.

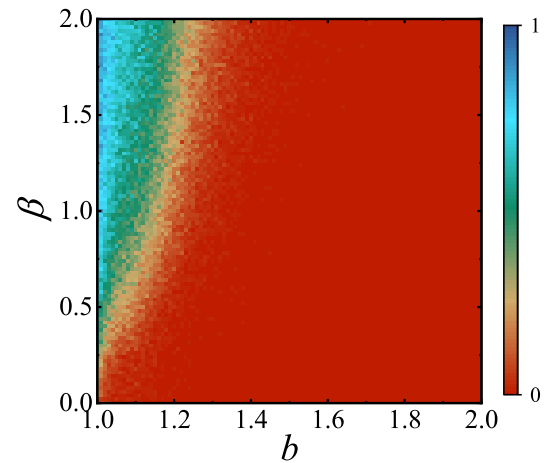


**Fig. 7.** Evolution of interaction intensity and cooperation frequency. The average values of the interaction intensity between cooperators  $p_{cc}$ , between defectors and cooperators  $p_{dc}$ , and the frequency of cooperation  $f_c$  evolve with time, which are represented by blue curve, red curve, and black curve, respectively. From left to the right, parameters are set as  $\beta = 0.1$ ,  $b = 1.05$  and  $\beta = 2$ ,  $b = 1.5$ , respectively. The insets are the amplifications of the first 200 steps of whole evolution.

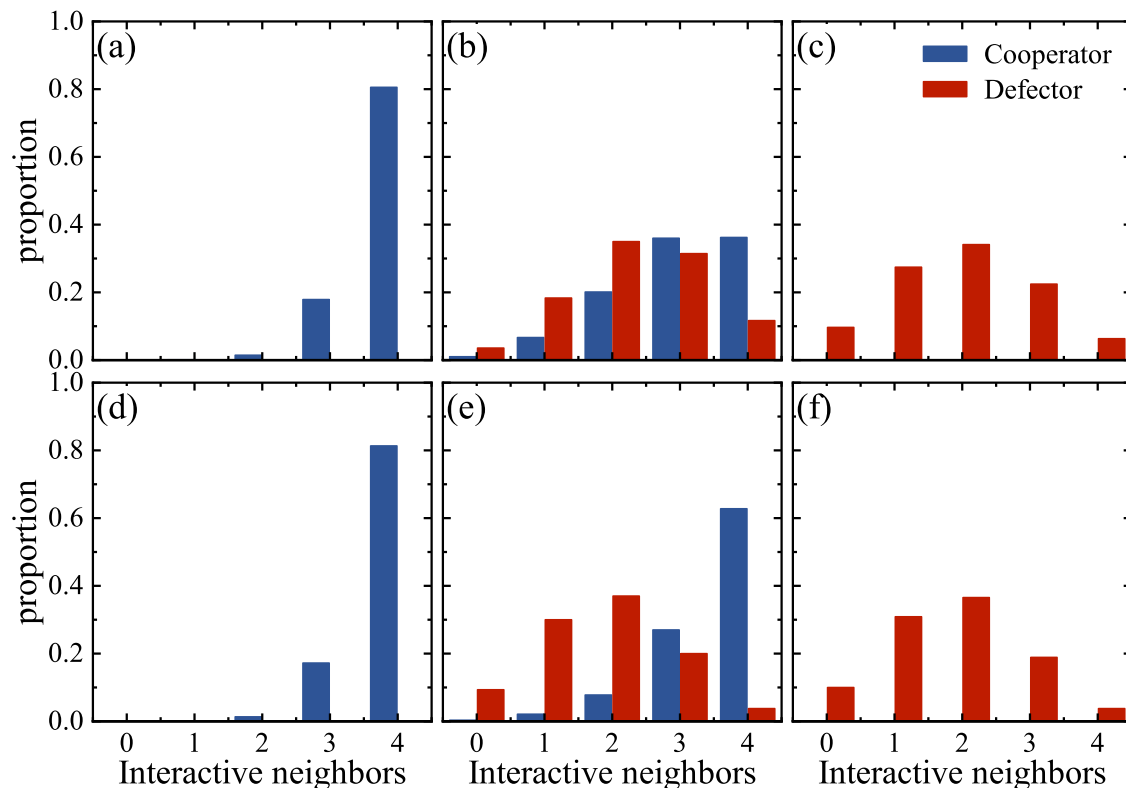
that the overall evolutionary patterns are similar, except for some notable differences. When  $\beta$  is small, the interaction intensity evolves relatively slowly. Consequently, it leads to the inability to provide the optimal interaction intensity for the evolution of cooperation, to effectively promote cooperation. However, when  $\beta$  is large, the interaction intensity evolves very fast, and it already reaches a steady state before the EXP period. This implies for large  $\beta$ , the population will efficiently establish an optimal interaction relationship. In addition, this optimal interaction relationship can benefit the cooperation of the population. These results are consistent with observations in Fig. 3. (See Fig. 8).

The number of interactive neighbors distribution for both cooperators and defectors is various for different cases of  $\beta$  and  $b$ . In the full cooperation phase (Fig. 8(a) and (d)), agents establish more interactions with their neighbors to obtain higher payoffs. While in the full defection phase (Fig. 8(c) and (f)), agents cannot receive payoffs whether they interact with neighbors or not, so they will finally form a random interaction relationship. It is worth noticing that in Fig. 8(b) and (e), where the cooperators and defectors coexist at a steady-state, the distributions vary for different  $\beta$ . The number of interactive neighbors of cooperators under  $\beta = 2$  is larger than the case of  $\beta = 0.1$ . On the contrary, the number of interactive neighbors of defectors under  $\beta = 2$  is smaller than the case of  $\beta = 0.1$ . This indicates that a relatively larger  $\beta$  is more beneficial for cooperation. More importantly, one can find when  $\beta$  is large, the distribution of both cooperation and defection is closer to that in the phases of full cooperation and defection, respectively. This means when  $\beta$  is large, the interaction intensity evolves faster, so agents will efficiently establish a steady interaction relationship that is closer to the optimal interaction intensity. Thus, it becomes more clear that the interaction intensity is enhanced by cooperation, which in return promotes cooperation in the population.

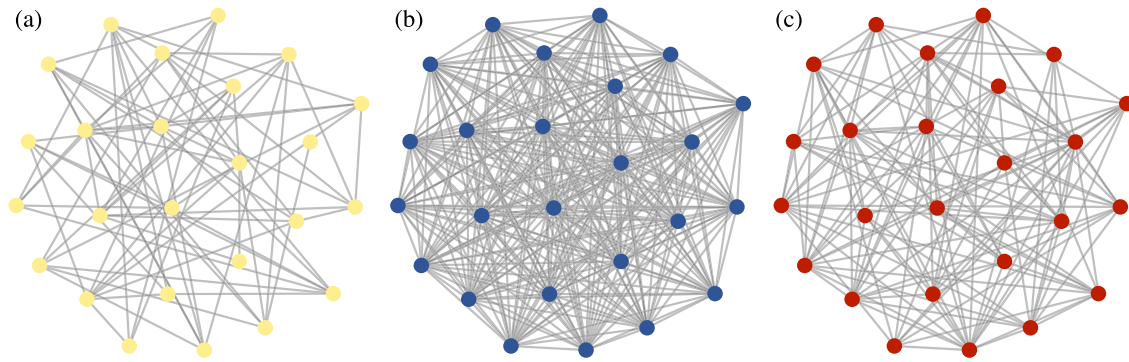
At last, we test whether the model presents a similar effect on the well-mixed network. Fig. 9 shows the cooperation frequency at the steady-state on a well-mixed network with 25 nodes. Initially, cooperators and defectors distribute randomly with the same probability, and the interaction willingness follows a uniform distribution. Since the final cooperation frequency will always be 0 or 1, the cooperation frequency is an average of 50 simulations for each combination of parameters  $\beta$  and  $b$ . The introduction of the



**Fig. 9.** Frequency of cooperation in  $b - \beta$  panel. The panel shows the average cooperation frequency of 50 independent simulations as a function as  $b$  and  $\beta$ . Red represents full defection phase in all 50 simulations, and blue represents full cooperation phase in all 50 simulations. The initial interaction willingness follows the uniform distribution in  $[0, 1]$ .



**Fig. 8.** Distribution of interactive neighbors of cooperators and defectors. Parameters are set as  $\beta = 0.1$ , and  $b = 1.025, 1.05, 1.075$  in (a)(b)(c), while  $\beta = 2$ , and  $b = 1.45, 1.5, 1.55$  in (d)(e)(f). Results of cooperators and defectors are denoted by blue and red columns, respectively.



**Fig. 10.** Illustrations of the well-mixed networks. (a), (b), and (c) represent the initial network, network in full cooperation phase, and network in full defection phase, respectively. Grey lines denote the interaction between agents. Yellow nodes denote agents at the initial state regardless of their strategy. Blue and red nodes denote cooperators and defectors, respectively. We obtain the results with parameters  $\beta = 2$ ,  $b = 1.2$  in (b), and  $b = 1.6$  in (c).

adaptive interaction intensity improves cooperation. Moreover, as  $\beta$  becomes large, the critical value at which cooperation goes extinct is also enlarged.

The interaction networks of the full cooperation phase and full defection phase are given in Fig. 10 (b) and (c), respectively. As the initial interaction willingness of agents follows a uniform distribution, there will only be a quarter of edges having successful interactions in the initial network, as shown in Fig. 10(a). In the full cooperation phase, nearly all pairs between two cooperators have successful interactions. While in the full defection phase, the interactions are heavily sparse. Therefore, the results on both lattice and well-mixed networks prove the interaction intensity is enhanced between cooperators, and the optimal interaction intensity will promote cooperation.

## 6. Discussion

We investigate the evolutionary process of the iterated prisoner's dilemma game with adaptive interaction intensity based on the reinforcement learning method. Different from the traditional evolution process, agents with adaptive interaction intensity can decide whether to interact with nearest neighbors or not. The results demonstrate that the adaptive interaction intensity enhances the cooperation in the population and is beneficial for solving social dilemmas. The evolution of the interaction intensity and cooperation is affected by the sensitivity parameter of the BM model. When this parameter is small, the interaction intensity and the strategy will mutually influence each other for a relatively long period. On the contrary, when the parameter is large, the interaction intensity will fast evolve to an optimal point, which enlarges the critical value of cooperation becoming extinct. In other words, with a relatively large sensitivity, the adaptive interaction intensity makes cooperation more competitive to survive. In addition, we obtain a universal conclusion in the well-mixed population, which excludes the influence of network structure. That is, cooperative behavior enhances interaction, which conversely improves cooperation in groups. The conclusions we obtained are also theoretically confirmed by the extended pair approximation.

Inspired by the reality that the interactions between agents change over time, we use the BM model to realize the adaptive interaction intensity. The simulation and theoretical results demonstrate adaptive interaction intensity can promote cooperation greatly. One may find that since this mechanism allows agents to timely terminate the interactions with poor-behaving neighbors, it leads to the appearance of isolated agents who have no interactive neighbors. However, we would stress that timely stopping the interaction with poor-behaving neighbors and establish-

ing new interactions with good-behaving neighbors still is a wise choice in the long term. Because it promotes cooperation in the population, which is beneficial for not only agents themselves but also the whole population [57].

Our work is a simple attempt to introduce adaptive behavior into evolutionary games. In addition to the adaptability of the agents' interacting behavior, it is also worth considering the environmental interventions and constraints in the further works. Along the line of our work, there could be some future endeavors. For example, agents might be able to switch the type of the game according to self-regarding skills and judgments (e.g., Q-learning process [46]). We hope this work will be an inspiring exploration that sheds light on solving social dilemmas.

## CRedit authorship contribution statement

**Zhao Song:** Software, Formal analysis, Investigation, Visualization, Writing - original draft. **Hao Guo:** Software, Writing - review & editing. **Danyang Jia:** Writing - review & editing. **Matjaž Perc:** Writing - review & editing. **Xuelong Li:** Methodology, Conceptualization, Writing - review & editing. **Zhen Wang:** Supervision, Writing - review & editing.

## Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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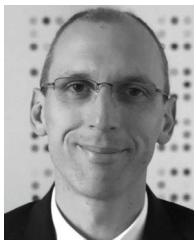
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