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Evolving learning rules and emergence of cooperation in spatial Prisoner’s Dilemma

Luis G. Moyano\textsuperscript{a} and Angel Sánchez\textsuperscript{a,b,c}

\textsuperscript{a}Grupo Interdisciplinar de Sistemas Complejos (GISC), Departamento de Matemáticas, Universidad Carlos III de Madrid, 28911 Leganés, Madrid, Spain
\textsuperscript{b}Instituto de Ciencias Matemáticas CSIC-UAM-UC3M-UCM, 28006 Madrid, Spain
\textsuperscript{c}Instituto de Biocomputación y Física de Sistemas Complejos (BIFI), Universidad de Zaragoza, 50009 Zaragoza, Spain

Abstract

In the evolutionary Prisoner’s Dilemma (PD) game, agents play with each other and update their strategies in every generation according to some microscopic dynamical rule. In its spatial version, agents do not play with every other but, instead, interact only with their neighbors, thus mimicking the existing of a social or contact network that defines who interacts with whom. In this work, we explore evolutionary, spatial PD systems consisting of two types of agents, each with a certain update (reproduction, learning) rule. We investigate two different scenarios: in the first case, update rules remain fixed for the entire evolution of the system; in the second case, agents update both strategy and update rule in every generation. We show that in a well-mixed population the evolutionary outcome is always full defection. We subsequently focus on two-strategy competition with nearest-neighbor interactions on the contact network and synchronized update of strategies. Our results show that, for an important range of the parameters of the game, the final state of the system is largely different from that arising from the usual setup of a single, fixed dynamical rule. Furthermore, the results are also very different if update rules are fixed or evolve with the strategies. In these respect, we have studied representative update rules, finding that some of them may become extinct while others prevail. We describe the new and rich variety of final outcomes that arise from this co-evolutionary dynamics. We include examples of other neighborhoods and asynchronous updating that confirm the robustness of our conclusions. Our results pave the way to an evolutionary rationale for modelling social interactions through game theory with a preferred set of update rules.

Key words: Game theory, evolution, prisoner’s dilemma, learning, emergence of cooperation.

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1 Introduction

The quest for the origins of the cooperative behavior observed in nature, in many different species or at different biological levels, from molecules to individuals, is an exciting project that has received much attention in the last decades [Darwin, 1871; Axelrod and Hamilton, 1981; Maynard-Smith and Szathmáry, 1995; Hammerstein, 2003]. Evolutionary game theory has been one of the most successful frameworks to address this issue in a quantitative manner [Gintis, 2000; Nowak, 2006b; Pennisi, 2005] and by allowing a stylization of the main strategic interactions and social dilemmas, has provided many insights into the reasons of the emergence of cooperation.

A particularly fruitful line of research has focused on the interactions between two individuals, trying to explain cooperative behaviors in a population with a bottom-up approach. In this context, interactions are modelled by means of 2x2 games, such as the Prisoner’s Dilemma [Rapoport and Chammah, 1965] or the Hawk-Dove game [Maynard-Smith, 1982]. These games have proven themselves relevant in situations arising in biochemistry [Frick and Schuster, 2003], cooperation between bacteria [Crespi, 2001], mutualistic interactions [Kiers et al., 2003], fish [Dugatkin and Mesterton-Gibbons, 1996] and, of course, human societies [Kollock, 1998].

Within the framework of evolutionary game theory, a number of explanations have been proposed to understand the origin of cooperation [Nowak, 2006a]. In this work, we focus on one of them, namely the existence of a spatial structure, as such or as a representation of a social network. Indeed, many studies have pointed out that certain types of spatial structure foster cooperation in simple two-player symmetric games, beginning with the pioneering work by Nowak and May [1992]. Subsequent papers [Nowak and May, 1993; Hauert, 2002, 2006; Santos et al., 2006a,b; Jiménez et al., 2008; Lozano et al., 2008; Gardeñes et al., 2008; Langer et al., 2008] analysed different aspects of the emergence of cooperation in spatial games and as a conclusion of this work the general idea that spatial structures supported cooperative behaviors began to shape up. For a comprehensive summary of all the recent work on evolutionary game theory on graphs, see the review by Szabó and Fáth [2007].

Recently, some authors have questioned the generality of the above assertion, at least as far as other games are concerned. Thus, Hauert and Doebeli [2004] and Sysi-Aho et al. [2005] have shown that spatial structure may decrease the cooperation level attained in the Hawk-Dove game as compared to that observed in a well-mixed population. Following this result, some researchers have looked in detail into the different reports and found that the phenomenon of the emergence of cooperation, when truly existing, turns out to be dependent on the microscopic update rules used in the simulation. In other words, it
is possible that within the same game, played in the same spatial structure, cooperation arises or not depending on the way the players change their strategy during evolution. This is the case, for instance, when playing Prisoner’s Dilemma on a square lattice: whereas unconditional imitation (see below for a description of this and the rest of update rules studied in this paper) gives rise to cooperation [Nowak and May, 1992], replicator dynamics leads to full defection [Roca et al., 2008]. Similar dependences of the results for other update dynamics have been reported by Ohtsuki and Nowak [2006] for death-birth, birth-death and imitation.

In view of this situation, in this paper we aim to going beyond the approach that has been traditionally used in the study of spatial games. Specifically, we intend to provide an evolutionary rationale for choosing a particular update rule in the implementation of spatial models of cooperation. To this end, we will allow agents to update not only their strategies but also the update rule itself. The outcome of these simulations will show whether or not all update rules are equally likely to appear in a structured population and, if not, which ones are evolutionarily selected. One can then compare this conclusion to the scenarios of emergence of cooperation on networks and discuss the applicability of the different results already known. Indeed, the fact that rules favoring cooperation were evolutionarily favored would support the mechanism of network reciprocity as one of the most important ones for the emergence of cooperation. On the contrary, if the competition among update rules leads to the survival only of those that do not support cooperation, it would be difficult to argue that networks promote cooperation.

For the present study, we will be concerned only with the problem of one-shot or memoryless Prisoner’s Dilemma, as has been generally studied in the context of evolutionary game theory on graphs; rules for deciding the action to be taken next on the basis of previous ones, such as tit-for-tat, Pavlovian strategies, stochastic reactive strategies, etc. Although it is possible to think of these strategies as C or D strategists with a learning rule, their use of memory place them in a different class that and will not be considered here. On the other hand, we here focus on a typical set of local rules, as considered, for instance in Hauert [2002]; it is clear that this kind of evolutionary competition may extend to many other update dynamics but an exhaustive analysis of every possible rule is beyond the scope of the present research, that intends only to assess the relevance of such an evolutionary process.

In this paper we address the co-evolution of strategies and update rules in a three-step process. As a preliminary result, we discuss the case of well-mixed populations and show by an example that in this situation including evolving update rules does not change the well-known outcome, namely that defection prevails. Then we move to the case in which the population interaction is governed by a lattice, as a specific example of social network in which it is
easier to interpret the results. In order to have a reference, a first step in our approach is the comparison of mixed systems, consisting of agents with different (but fixed during evolution) update rules, with the emergence of cooperation in pure single-dynamics systems. Already at this stage we find differences between the two scenarios which are worth describing; on the other hand, this is a necessary reference frame to understand the subsequent results of co-evolution. Indeed, after this first step, we proceed to let update rules evolve along with strategies. In this situation, we find new results that differ both from the single-rule case and from the mixed-rule case. We will describe the results of our simulations and in the conclusions we summarize our findings and discuss their implications for the emergence of cooperation.

2 Model

Our model is based on the well-known Prisoner’s Dilemma (PD) game [Rapoport and Chammah, 1965]. An archetype in game theory, the PD game belongs to a general class of symmetric games that consist of two players or agents that confront, each with a definite strategy: either to cooperate or to defect with the opposite player. Each combination of strategies between the players has an associated payoff, and hence there are four possible outcomes: if the player cooperates, the associated payoff she gets is $S$ if the other player chooses to defect, or $R > S$, if the other player reciprocates the cooperation. On the other hand, if the first agent defects, her payoff is $P$, if the other also defects, or $T > P$, if the other cooperates. The PD corresponds to any choice of payoffs ordered according to $T > R > P > S$. It is customary to assume that $2R > T + S$ to avoid that players take turns at defecting and win a larger payoff than the one they would have just cooperating. We will stick to this constraint although in our case we deal with one-shot, non-repeated games because strategies and payoffs are updated after every single game, and hence strictly speaking we need not impose this condition. In the rest of this work, we will adopt the commonly-used rescaled payoff $T = b$, $R = 1$, and $P = S = 0$ [Nowak and May, 1992]. This does not affect the general essence of the game, and reduces the study to just one free parameter, usually called the temptation parameter. Nevertheless, we checked the robustness of our simulations by comparing some cases with the choice $S = -0.2$, finding very approximately the same results.

In the evolutionary version of the PD game, $N$ agents play between them, and after every instance of the game they decide whether to change strategies or not according to some microscopic update rule. In a well-mixed situation, each player plays every other once and afterwards they proceed to the update stage. In our case, as we are interested in the rules that can promote cooperation in a spatially structured population, the players are located at the nodes of a square...
lattice, where each agent is connected with her four closest neighbors and plays the game only with those neighbors. The reason we have chosen a square lattice is that it is a well studied model [Nowak and May, 1992; Hauert, 2002; Roca et al., 2008; Langer et al., 2008] in the single-update rule framework, and therefore we can compare our results to those previous works. The sequence of steps for the simulation is as follows: Each player is assigned, randomly and with equal probability, an initial strategy of cooperation (C) or defection (D), and a payoff, initially set to zero. In each generation, all agents play PD once with each neighbour, with an associated payoff for each game, collecting a total final payoff for each player. After each generation, all agents update their strategies simultaneously, according to a certain update rule (defined separately), that may depend on the agent’s payoff and her neighbour’s, and that defines the dynamics of the game. Once defined, for all players, what the strategy for the next generation will be, all payoffs are reset to zero and the cycle starts over.

As we have said, the new ingredient of our model is twofold: two different update rules and the possibility that the update rules themselves evolve. Therefore, in our model players may be set to have rule A or rule B, where A and B stand for specific dynamical rules that will be explained below. In this way, each player has its own individual dynamical rule. Within this framework, we devised two possible alternatives. In the first case, agents are assigned their dynamical rule, that remains fixed for the rest of the simulation. In the second case, agents may change their dynamical rule during the game according to a simple criterion: an agent copies the dynamical rule from its neighbour whenever it copies its strategy. This can be interpreted as a complete replacement of one agent for another agent’s offspring, which may be convenient or useful in certain descriptions. This interpretation is suitable for both biological and sociological situations, in this last case in terms of culture transmission and learning. Furthermore, the possibility of variable update rules implies an evolution (and therefore a competition) of the rules themselves. If as a consequence of this evolution one rule, or a restricted set of rules, are selected, the results on the emergence of cooperation on lattices will have to be revisited again in the light of our findings.

We implement three of the most representative, and most commonly considered, dynamical rules: the unconditional imitation (UI, a.k.a. as imitate-the-best) rule, the Moran (MOR) rule and the replicator (REP) rule. Unconditional imitation is a completely deterministic rule: at the end of each generation, an agent simply adopts the strategy of her neighbour with the highest payoff (i.e., the most successful one), given that this neighbour has a greater payoff. Note also that this rule checks the payoff of all the neighbors of the agent whose strategy is to be updated. In this sense we will refer to this strategy as global (not to be confused with global in terms of the whole lattice). In the replicator dynamics, an agent randomly chooses another agent
(in our case, one of its four neighbours) and, if the chosen agent has a higher payoff, the original one adopts that neighbour’s strategy with a probability proportional to the difference of payoffs between the two. In this case, we are faced with a local update rule, that does not look at all the updating agent’s neighbourhood. Another important remark is that, in our model, an agent having an imitation or replicator rule will never adopt another strategy (or rule) that performed worse. Finally, in a Moran process, the agent, at the end of each generation, evaluates a set of probabilities, one for each neighbour and proportional to that neighbour’s payoff. Then the agent randomly selects a neighbour’s strategy according to that set of probabilities. In this case, there is a possibility that an agent adopts a strategy that performed worse in a previous round. On the other hand, this is again a global rule, in the same sense as we referred above to the imitation one.

3 Well-mixed populations

Before proceeding with the study of the evolutionary competition of learning rules on lattices, it is important to analyze the case in which every player plays against every other one, i.e., a well-mixed population. The reason for the need of such a study is twofold: on one hand, knowing the behavior of a well-mixed population is necessary in order to assess whether or not changing the scenario to a lattice has any new effect; on the other hand, the well-mixed population can be used to understand at least the initial stages of the evolution on a lattice, when correlations arising because of evolution are not yet important and the assumption that agents meet an "average" agent can be made. We note that this assumption is very common in the statistical physics literature where it is referred to as the mean-field approach (see, e.g., Szabó and Fáth [2007] and references therein).

For the sake of simplicity, in what follows we will consider the case in which an initial fraction $x$ of imitator agents, i.e., agents that learn through UI, compete with an initial fraction $1 - x$ of replicator agents, agents using REP. The other possible competitions can be treated in much the same way with similar results, therefore we use this particular choice as an illustration. If initially a fraction $y$ of agents are cooperators (equally distributed among UI and REP players), the four types of agents present in the population at time $t = 0$ are given by

\begin{align}
    f_{CI}^0 &= xy, \\
    f_{DI}^0 &= x(1 - y), \\
    f_{Cr}^0 &= (1 - x)y, \\
    f_{Dr}^0 &= (1 - x)(1 - y),
\end{align}
where the subindices \( C \) or \( D \) represent cooperators and defectors, respectively, and \( i \) and \( r \) refer to imitators and replicators, also respectively.

Evolution begins by all agents playing the game with all the population. With our choice of payoffs, cooperators receive a payoff \( w_C = x \) and defectors receive a payoff \( w_D = bx \). Let us now examine the evolutionary process at the first step, beginning with cooperators:

**Imitator agents:** As \( b > 1 \), we have \( w_D > w_C \) and therefore all imitator agents switch to defection at time \( t = 1 \), and hence \( f^1_{Ci} = 0 \); the newly created defector will be an imitator with probability \( x \) and a replicator with probability \( (1 - x) \).

**Replicator agents:** The fate of replicator agents is more complicated and in particular the evolution of their number in time depends on the form chosen for the probability to copy the other agent’s strategy. If \( p \) is the probability of a cooperator replicator to switch to defection (the value of \( p \) depends on the payoff difference, which is \( b(1 - x) \), and on the normalization), we have the following scenario: The probability that a replicator chooses a cooperator to compare her strategy is \( y \) (the fraction of cooperators at \( t = 0 \)); in that case, the payoffs are the same and she will not change her strategy and update rule. On the contrary, with probability \( 1 - y \) she will compare with a defector, and will turn into a defector herself with probability \( p \). However, this defector will be an imitator with probability \( x \) and a replicator with probability \( 1 - x \).

Finally, in view of the payoffs above, defectors never change, irrespective of their update rule. Collecting all the different contributions, we have at \( t = 1 \)

\[
\begin{align*}
    f^1_{Ci} &= 0, \\
    f^1_{Di} &= f^0_{Di} + x f^0_{Ci} + px(1 - y) f^0_{Cr}, \\
    f^1_{Cr} &= [y + (1 - p)(1 - y)] f^0_{Cr} = [1 - p(1 - y)] f^0_{Cr}, \\
    f^1_{Dr} &= f^0_{Dr} + (1 - x) f^0_{Ci} + p(1 - x)(1 - y) f^0_{Cr}. 
\end{align*}
\]

From Eq. (7) it is clear that the fraction of cooperator agents will steadily decrease; imitators disappear at the first step and replicators will decrease exponentially [note that the coefficient in Eq. (7) is smaller than 1]. As a consequence, asymptotically the population will evolve to full defection. On the other hand, when all cooperator replicators disappear, it can be shown straightforwardly that they would have contributed to the two types of defectors simply proportionally to their initial fraction, i.e.,

\[
\begin{align*}
    f^\infty_{Di} &= f^0_{Di} + x(f^0_{Ci} + f^0_{Cr}) = x(1 - y) + xy = x, \\
    f^\infty_{Dr} &= f^0_{Dr} + (1 - x)(f^0_{Ci} + f^0_{Cr}) = (1 - x)(1 - y) + (1 - x)y = (1 - x). 
\end{align*}
\]
This means that the effect of introducing a lattice as support of the game has indeed non-trivial consequences because, as will be shown below, there is cooperation in a wide range of parameters, and the level of cooperation and the fractions of the different types of strategists and update rules can not be predicted from the initial fractions in such an straightforward manner.

4 Results on lattices

To study our model, we performed a series of numerical experiments for both cases, namely with fixed and with variable update rules, for the same set of game parameters. Each experiment consists in a population of \( N = 10^4 \) agents, spatially arranged in a square lattice. Generally speaking, each agent is endowed initially with one of two available dynamical rules, A or B (where A and B stand for UI, MOR or REP), and one of two possible strategies, C or D. To monitor the evolution of the system we will observe the frequency or density of a certain type of player, for example those that are cooperators or those with a certain update rule. Thus, we define \( f_A \) to be the number of agents with rule A divided by \( N \), and \( f^C \) the number of cooperator agents divided by \( N \). Every experiment is characterised by an initial density of cooperators \( f^C(t = 0) \) (or in a simpler notation, \( f^C(0) \)) and by an initial fraction of agents with rule A, \( f_A(0) \). In this way, \( f^C \) will generally change. The homogeneous case (all agents with the same update rule) is recovered by choosing \( f_A(0) = 1 \). To simplify the parameter space of our simulations, we only considered initial strategies assigned randomly and with equal probability, i.e. \( f^C(0) = 0.5 \) in all experiments, independent of the update rule considered. All numerical simulations run for \( T = 10^4 \) generations or time steps, for a given value of \( b \). Within this duration equilibration was achieved (often much earlier), i.e., the densities reached their asymptotic values and remain there within small fluctuations.

4.1 Fixed update rules

We begin by looking at the simplest case of agents that can have two different update rules but they cannot change them during evolution. In this case, \( f_A = f_A(0) \) and remains constant for all times. It is useful to analyse the normalised fraction of cooperators of, say, rule A, which we will refer to as \( \mu_A = f^C/f_A \), where \( 0 \leq \mu_A \leq 1 \), so we can better compare systems with different values if \( f_A(0) \).

A first finding arising from our experiments is that the initial fraction of the populations, \( f_A(0) \) may affect greatly the final cooperator outcome. Consider
for instance the left panel of Fig. 1, where we show an example of a system with fixed rules, plotting, for a population of UI and REP, the final (equilibrium) value of the relative cooperator density $\mu^T_{UI}$ and $\mu^T_{REP}$ as a function of $f_{UI}(0)$. We indeed observe that the initial ratio of agent types has a significant effect on the final outcome of the simulations, although this is not so in the case of UI (solid line): up to a value $f_{UI}(0) \sim 0.75$ of the total initial fraction $f_{UI}(0)$ the ratio $\mu^T_{UI}$ of UI cooperators is more or less constant, increasing slightly for larger values of $f_{UI}(0) \sim 0.75$. In the case of REP agents (dashed line), we get another interesting result: The initial fraction of replicator agents $f_{REP}(0) = 1 - f_{UI}(0)$ is approximately proportional to the amount of replicator cooperators within the replicator population, $\mu_{REP}$ being smaller (larger) than 0.5 whenever $f_{REP}(0)$ is smaller (larger) than 0.5. While this may look intuitive, it must be recalled that when the population consists only of REP agents, the cooperators die out for any value of the temptation parameter $b$. Regarding the time evolution of the experiment, the right panel of Fig. 1 shows two examples of the evolution of the normalized fraction $\mu_{UI}$ for different values of the initial fraction of imitators $f_{UI}(0)$. There is always an initial drop of $\mu_{UI}$ and a final relaxation to a value $\mu^T$ that, as stated before, depends on $f_{UI}(0)$.

![Fig. 1. Left: Value of $\mu^T_{UI}$ (solid) and $\mu^T_{REP}$ (dashed) at $T = 10^4$ as a function of the initial frequency of imitators agents $f_{UI}(0)$. Rules are fixed throughout the simulation. Right: Time evolution of the normalized cooperator frequency $\mu_{UI}$ in the UI vs. REP game, for $b = 1.05$ and two values of the initial frequency of imitators $f_{UI}(0)$.]

Having considered the effect of different populations of update rules, let us now discuss the dependence on the temptation parameter $b$, and let us compare the results for the homogeneous case (all agents equal) with the case where two different rules are present. As a specific example, in Fig. 2 we plot, for the same game as before, the dependence of the total cooperator density as a function of the parameter $b$ for the homogeneous case $f_{UI}(0) = 1$ and for a mixed case with fixed $f_{UI}(0) = 0.75$. For the homogeneous case, using the fact that unconditional imitation is a deterministic rule, one can show that $f_{UI}$ depends on $b$ in a step-like fashion, changing when $b$ is $\frac{4}{3}$, $\frac{4}{5}$, and $\frac{3}{2}$.
perfect agreement with the results of our simulations. As we may see from the plot, this functional dependence changes when there are replicators among the imitators. We find that the effect of the replicators is to lower the amount of total cooperators for a the whole range of values of $b$ corresponding to the PD, e.g. $b \geq 1$. In this respect, it is interesting to look at how the fraction of cooperators changes relative to the population with the same rule. This is also plotted in Fig. 2, by showing both $\mu_{UI}$ and $\mu_{REP}$ as a function of $b$. We see that, for $1 < b < \frac{3}{2}$, even though the total cooperator density is lower than in the pure UI case, the cooperative imitators are a fraction larger than the total average, whereas the replicators are less than the average. On the other hand, for $b > \frac{3}{2}$, defectors take over the entire population, so $\mu_{UI} = \mu_{REP} = 0$. This means that the level of cooperation attained for a pure UI population in the PD suffers a considerable decrease or are totally suppressed by the presence of a minority of REP agents (we have checked with populations as small as $f_{REP} = 0.01$ obtaining similar results). We will come back to this issue when considering different neighborhoods below.

![Fig. 2. Density of cooperators for the UI vs. REP game with fixed dynamical rules. In thick solid lines, we present the case of a pure imitator population. In thick dashed lines, same settings but with $f_{UI}(0) = 0.75$. In thin solid (dashed) line we show the normalised density of cooperator imitators (replicators) $\mu$. All quantities are averages of over 30 realisations.](image)

4.2 Variable dynamical rule

In the previous subsection, we have reported that the presence of two different update rules in the population may considerably change the behavior of the PD on a lattice, the general conclusion being that the level of cooperation is lower than for the pure UI population. This result must be kept in mind when
analysing the outcome of allowing the update rules themselves to evolve, the issue which we focus upon in the following. When agents are allowed to switch their update rule when updating their strategies, then the fraction of agents of rules A and B, \( f_A, f_B \), will generally change. Indeed, in this case, we will see that, for certain values of the game parameters, a rule can completely overtake the system as the other one disappears, yielding different outcomes than the ones obtained with fixed update rules, or with just one update rule.

\[\begin{align*}
\text{Fig. 3. Top panel: The UI vs. MOR game with fixed rules for } f_{UI}(0) &= 0.12 \text{ and } b = 1.20. \text{ In solid line, the total cooperator density } f_C. \text{ In thin solid (dashed) line the imitator (Moran) cooperator density. Bottom panel: same simulation as in top panel with update rule changing permitted. Lines have the same meaning as in top panel.}
\end{align*}\]
To begin with, let us present and discuss a specific example: In Fig. 3 we collect the results for the competition between UI agents versus MOR agents. On the top panel we show the cooperator frequency evolution with fixed update rules, i.e., when agents cannot change their update rules. We clearly observe that the total cooperator fraction \( f^C \) disappears at about \( t \sim 300 \). Note that both UI and MOR cooperators disappear at about the same time as defectors take over the entire population, in agreement with our conclusion of the previous subsection that inhomogeneous agents lead to lower levels of cooperation. Subsequently, let us consider the bottom panel of Fig. 3, where we show a simulation under the same conditions with the only difference that in this case agents also copy their neighbour’s update rule if they copy their (C or D) strategy. Opposite to the situations with fixed update rules, not only the total fraction of cooperators does not disappear, but also we find that cooperators end up forming about three quarters of the total population. Interestingly, there is an initial decrease in the cooperation fraction, in agreement with the predictions of the well-mixed/mean field calculation summarized in Sec. 3 (here REP is replaced by MOR, but the argument is very similar and applies as well). It’s only at a later stage when the effect of spatial correlations, namely the formation of clusters of cooperators, enters into play and leads to an increase of cooperation [Hauert, 2002; Nowak and May, 1992; Roca et al., 2008]. Analysing the individual rules, we see that MOR cooperators disappear roughly at the same pace (actually, a little faster than in the fixed rule case), so all cooperators remaining are of the UI type. Remarkably, UI agents completely replace MOR agents in spite of the fact that the initial fraction of imitators in this simulation is \( f_{UI}(0) = 0.12 \).

We could continue our discussion of specific examples but, to avoid a very lengthy description, let us only mention that other interesting outcomes may appear depending on the game parameters and update rule combinations. For instance, it is possible to see a coexistence of cooperators of one rule with defectors of the other rule, as occurs for imitators versus Moran agents with \( f_{UI}(t = 0) < 0.5 \) and \( b = 1.05 \). However, as we are more interested in general conclusions than on a detailed classification of all the possible outcomes, we will switch to a more general viewpoint in what follows.

We will now discuss the dependence of the results on the temptation parameter \( b \). In fact, as agents are now allowed to change the dynamical rules, both the fraction of cooperators, \( f^C \), and the fraction of agents with rule A, \( f_A \), may change, so it is relevant to ask what is the dependence on the temptation parameter \( b \) for both of these quantities. In Fig. 4 we show, as an example, our results for the case of UI and MOR agents as a function of \( b \). We see that depending on the value of \( b \), the interaction affects the final fraction of cooperators to different extents. Comparing with the homogeneous UI case, we observe that there is a lower fraction of cooperators for the mixed case when \( 1 < b < 1.25 \) (the second bound is approximate) and for \( b > \frac{5}{2} \). On the
other hand, the mixed case shows more cooperation when $b < 1$ and in the interval $1.25 < b < \frac{4}{3}$. Finally, the two curves coincide for $\frac{4}{3} < b < \frac{3}{2}$. The plot also depicts the final fraction of imitator agents $f_{UI}$ (cf. right axis) for the mixed case. Indeed, for most values of $b$, i.e. $b < \frac{3}{2}$, UI agents have increased their frequency from the initial value $f_{UI}(0) = 0.12$, even becoming the only rule ($f_{UI} = 1$) for certain regions ($b < 1$ and $1.25 < b < \frac{4}{3}$). Note that, for the cases $b < 1$ and $1.25 < b < \frac{4}{3}$, the presence of Moran agents at early stages of the evolution makes the equilibrium state to have more cooperation than the homogeneous UI case, even though Moran agents finally disappear completely. This contrasts with the case $\frac{4}{3} < b < \frac{3}{2}$, where the cooperator frequency is the same as in the homogeneous case.

![Fig. 4. Density of cooperators $f_C$ and density of imitators $f_{UI}$ as a function of the temptation parameter $b$, for the UI vs. MOR game. Simulation time is $T = 10^4$. In thick solid line, we show $f_C$ for the case of a population composed by imitators only. In dashed line, we present the same quantity for a mixed system with $f_{UI}(0) = 0.12$. In thin solid line, the final fraction of imitators (both cooperators and defectors). All quantities are averages of over 30 realisations.](image)

5 Extensions

5.1 Moore neighborhood

The results in Subsec. 4.1 are an indication that conclusions such as the promotion of cooperation on the PD on lattices found by Nowak and May [1992] may not be robust against the presence of other types of update strategists, and therefore that their applicability must be studied depending on the way individual agents behave. However, those results have been obtained under the
restriction that players interact only with their nearest-neighbors (von Neumann neighborhood), whereas Nowak and May included next-nearest neighbor interactions (Moore neighborhood). To check that the size of the neighborhood does not change our conclusions, we have carried out simulations with the Moore neighborhood, the results being depicted in Fig. 5. In fact, we have reproduced the simulations reported by Nowak and May [1992], which do not correspond exactly to a Moore neighborhood in so far as every player plays also with herself, i.e., it is an 8+1 neighborhood. As we may see from the example in the figure, we again find much lower levels of cooperation than in the pure imitator case, irrespective of whether the learning rules themselves evolve (right panels) or not (left panels). We note that this is not a general feature, as striking differences have been found between the presence and the absence of next-nearest-neighbor interactions, which makes our conclusion even more relevant as it is not trivial. We thus see that indeed the existence of different update rules in the population hinders cooperation on the square lattice.

5.2 Asynchronous updating

Although it is beyond our purposes to go into an exhaustive study of asynchronicity, we think that providing at least some examples of that case will increase the value and relevance of our results. To this end, we have run some simulations in which every time step a single agent is chosen at random and updates her strategy according to her current learning rule. Our results are summarized in Fig. 6 for fixed update rules (left) and changing update rules (right). We see from the plot that although there are a few quantitative differences, particularly relevant for the competition between UI and MOR with fixed update rules, the behavior is qualitatively the same in the synchronous and the asynchronous updating schemes. It is thus clear that our conclusions regarding the replacement of one rule by other are not an artifact of the synchronous update.

6 Discussion

As we have already said, our goal in this work is to extract general conclusions, and therefore we will now proceed to summarize the main results of our experiments. We simulated three possible competitions, namely UI vs MOR, UI vs REP, and REP vs. MOR, for both fixed and variable update rules, exploring the whole interval of temptation values as well as initial conditions with different composition for each case. Our results are summarized in Fig. 7, where we can see that, generally speaking, the dominant update rule is REP, in the sense that in most situations it dominates over the other rules.
Fig. 5. Mixed games with Moore neighbourhood and self-interaction (8+1 neighbours), for fixed and variable rules and $f^C(0) = 0.50$. Initial rule composition is half and half for all cases.

even if its initial population is small. This is particularly so in the case of REP vs MOR, for which the presence of MOR agents in the final population is almost negligible even if the initial population contains only a 25% of REP agents (see Fig. 7 bottom). The REP rule also dominates over the UI rule, although in this case UI does not go fully extinct except for very small ranges
Fig. 6. Mixed games with synchronous (thick, dashed line) and asynchronous (thin, solid line) update, for fixed and variable rules and $f^C(0) = 0.50$. Initial rule composition is half and half for all cases.

of parameters, and, in turn UI prevails over MOR, again without driving it to extinction. This is also shown in the asymptotic cooperation levels (Fig. 7, left panels): when the population is a mixture of REP with either UI or MOR, the dependence of the cooperation level on $b$ is similar to that of a full REP population, with some influence of the other rule (e.g., the abrupt drops
in cooperation at certain values of $b$ when the mixed population consists of REP and UI agents). A remarkable feature is that, when the population is a combination of UI and MOR agents, there is a range of values of $b$, between 1.25 and 1.5 (and up to 2, depending on the initial concentration of imitators), for which the cooperation level is larger than for imitators, but for parameter values and populations other than this case, the cooperation level is smaller than that reached in a full UI population. This result suggests that the good behavior observed on lattices and other networks homogeneous in degree when the agents are of the UI type may not be very robust to perturbations arising from agents learning with other rules. In any event, it is clear that the lattices with mixed populations do support cooperative states, which is a most important difference with the well-mixed population case discussed in Sec. 3.

Regarding the prevalence of REP, we believe that it is a consequence of its own dynamic mechanism, that has a built-in tendency to avoid changing its rule relative to the other dynamics tested in this work. Indeed, if we compare the definitions of the three dynamics, we notice that UI agents always updates its strategy (and consequently its update rule), excepting when the other agent has less or equal payoff; MOR agents have a set of probabilities that will always trigger the rule updating, unless all probabilities are zero, as in the very particular case of zero payoff for all neighbours (for example, in a defector-only population). Otherwise also the MOR agents update their rule with large probability, even adopting another rule with less payoff, leading to the preservation of a rule that performed poorer; and, finally, a REP agent randomly chooses a neighbour, and then assigns a probability to adopt its rule proportional to the payoff difference between both agents. For this rule there is only one possibility for having certainty of the rule update, and this is the case where all neighbours are defectors fully surrounded by cooperators, and the updating agent is a cooperator, in turn fully surrounded by defectors (in a chessboard-like fashion). In this case we have certainty because of our normalization factor for the probability, but it could even be the case that larger normalisation factors were used, which would lead to a large but smaller than 1 probability to change.

With this in mind, we monitored the fraction of agents of rule A (B) that update their rule relative to the total population having rule A (B), in each generation, called $\xi_A (\xi_B)$. As an illustration of the observed behaviours, in Fig. 8 we plot this quantity for the same case as Fig. 3 (bottom panel), i.e., UI vs MOR agents. The fraction of changing Moran agents rapidly becomes higher than that of imitators, and before 100 time steps is (and remains) much higher. We repeated this observation for many other cases as well and, in general, our results confirm that, in most cases, REP agents change their strategy and update rule much less frequently in comparison to the other dynamics. In some cases, this difference may be an order of magnitude smaller for the replicator rule. On the other hand the rule that updates the most is the
Fig. 7. Asymptotic fraction of cooperators (left) and of players with a given learning rule (right) for different initial concentrations of that rule for the cases UI vs REP (top), UI vs MOR (middle) and REP vs MOR (bottom). Thick solid lines, initial condition with a 100% of players of the first rule; thin lines correspond to mixed populations: solid, 75%, dotted, 50% and dashed, 25%, always of the first rule. Finally, thick dashed lines correspond to pure populations with a 100% of the second rule. All quantities are averages of over 5 realisations.
Fig. 8. Rate of change of imitator $\xi_{UI}$ and Moran $\xi_{MOR}$ agents, relative to their own population. Parameters are the same as Fig. 3 (bottom panel). Moran agents change most of the time at a higher rate than imitators (note the logarithmic scale in time).

Moran dynamics. Once again, this is consistent with our results: Most times, MOR agents disappear completely when confronting REP agents, and they survive in a restricted interval $1 < b < 1.1$ against UI, never being more than half of the final population (see Fig. 4, thin solid line), whereas for $\frac{4}{3} < b < \frac{3}{2}$ their presence is solely due to finite size effects. For UI vs REP, imitators change their rule more frequently, and replicators appear to be systematically the major part of the population.

In order to further investigate the hypothesis that the fraction of changing agents determine their prevalence, we devised the following experiment: We performed simulations with a similar setup as before, but in this case one of the populations updates their strategies and rules only at times $t = n\tau$, with $n = 0, 1, 2 \ldots$, being $\tau$ fixed (these would be “slow” agents). In this respect, it is interesting to recall that a few recent works have proposed that the existence of two types of agents, one of them with a smaller capability to transfer its strategy, may promote cooperation [Szolnoki et al., 2007, 2008; Masuda, 2008]. Results of applying this two-time scale setup to our simulation are collected in Fig. 9, where we see an example for normal REP agents against slow MOR agents. These results confirm our hypothesis: With $\tau = 10^1$, replicators prevail as usual (top panel), whereas for a sufficiently high update time, $\tau = 10^2$, MOR agents prosper, and replicators tend to become extinct (bottom panel).
Fig. 9. Normal replicator versus slow Moran agents, for different values of the parameter $\tau$. Temptation parameter is $b = 1.2$ and $f_{REP}(0) = 0.5$. Top panel: For $\tau = 10$ evolution of the cooperator density. Replicators (solid line) outperform Moran agents (dashed line), as in the normal ($\tau = 1$) case. Bottom panel: For $\tau = 10^2$ the opposite happens.

However, while this is an appealing mechanism, it cannot be the only responsible for the prevalence of strategies. To show it, we considered the situation where agents of the same kind are put together, the inhomogeneity being only that they can be “normal” or “slow”, i.e., that there is one fraction with a different $\tau$. In Fig. 10 we show the result of normal UI vs slow UI (see parameters in caption), where we observe that for all values of the temptation
parameter $b$, there is still a fraction of normal agents. If our hypothesis above were the only reason for the survival of one strategy, the result should be that slow UI would always take over the entire population, but this is clearly not the case. Indeed, normal agents do not disappear, and they only decrease in frequency noticeably (to $f_{UI} \approx 0.2$) in the interval $\frac{4}{3} < b < \frac{3}{2}$. We obtained similar results for a wide range of $\tau$, up to $\tau = 10^3$. This indicates that, at least for agents of the same type, the overall rule selection mechanism is not straightforward and should be further studied, and we believe that the conclusions would also carry over to the two update rule case. In addition, this makes clear that the two-learning rule model is a scenario that goes beyond that proposed by Szolnoki et al. [2007, 2008] and Masuda [2008] and therefore deserves further attention to be completely understood.

Fig. 10. Fraction of slow imitator agents $f_{UI^*}$ (thick solid line) as a function of the temptation parameter $b$. For comparison, we show also the total cooperation fraction $f^C$ (thin solid line) and the homogeneous (i.e., all agents with $\tau = 1$) cooperator frequency (thin dashed line). Initial fraction of slow agents is $f_{UI}(0) = 0.50$ and $\tau_{UI^*} = 10^2$.

A last point we want to remark from a general viewpoint is the following. Within our simulation procedure, a population composed only by defectors do not interact. Indeed, in this case the only available payoff is $P = 0$ independent of the agent’s update rule. REP or UI will not update their rule (they do so only with more successful neighbours), and MOR agents build a set of zero probabilities, thus keeping their rule as well. Thus, irrespective of the rules, the system remains in a “frozen” state, with no further evolution. A similar situation is found with a population composed only by cooperative imitators and replicators. In this case, all agents receive $R = 1$ and thus have the same payoff. Therefore, there is no rule update, and the system is, too, in a frozen state. On the other hand, this is not the case when any of these rules are set
against the Moran rule, where evolution does occur, and in most cases Moran players disappear. While all these are direct consequences of the design of our simulation, we have also found an interesting result regarding these locked situations, where no evolution is possible, namely that the presence of a small amount of a third type of agent may lead to evolution and, moreover, to a completely different state. Indeed, when these few agents can interact with at least one of the other type of (otherwise frozen) players, the system as a whole may evolve, even with the possibility of the extinction of one or more species, that otherwise would be present. We have not studied in detail this three update rule scenarios, but we envisage that the dynamics will be much richer and therefore deserves an in-depth analysis which is beyond the scope of the present paper.

7 Conclusions

In this paper we have presented a first attempt to provide an evolutionary rationale for the update rules used in spatially distributed models of emergence of cooperation. This is a key aspect of these models in so far as the choice of update rule largely influences the appearance and stabilisation of cooperation. In this context, our work must be viewed as a proposal for a general mechanism that would allow modellers to decide upon the rule of application in specific contexts. The main ingredient of this mechanism is the evolvability of update rules according to the same scheme as the strategies themselves, i.e., when an agent looks at her neighborhood and decides to copy the strategy of another agent, she also copies the agent’s update rule. In this respect, we want to stress that a related approach was proposed by Harley [1981] as a rationale to explain how populations can learn the evolutionary stable strategy. His results, which involve strategies with memory and referred to accumulated payoff, relate the stability of learning rules to their ability to take the population to the evolutionary stable strategy. Although our proposal here is quite different, in particular because our focus is not reaching an equilibrium but rather letting the system evolve at will, it is only fair to acknowledge Harley’s pioneering work in proposing the evolvability of strategies. On the other hand, our model, much as Harley’s procedure, can be interpreted as learning in social contexts, and hence it has in turn a much more biological character than endogenous learning rules such as those introduced by Kirchkamp [1999], that, to our knowledge, is the only other study where a evolutionary origin of learning rules has been explored. Note, however, that Kirchkamp [1999] considered a variety of games, while here we focus on the Prisoner’s Dilemma in view of its applicability in a number of social, biological and even biochemical systems. Finally, it is worth mentioning recent work by Szabó et al. [2008], where Darwinian selection is applied to a one-parameter stochastic update rule (similar
to those used by Szolnoki et al. [2007, 2008]), leading to the selection of a specific value of the parameter. This work is not related to ours in the update rule they use but the spirit is quite the same.

Beyond this general statement of the importance of the idea of evolvability of learning rules and its role to decide which ones should be used, we have reached several important conclusions of our work, that to our knowledge have never been reported elsewhere. In fact, their relevance arises not only for their own sake but also because they affect to two of the most often employed learning rules (UI and REP, see Szabó and Fátó [2007]) and because among the three rules we consider we cover the options local-global and deterministic-stochastic. Our main findings can thus be summarized as follows:

- A well-mixed population playing the Prisoner’s Dilemma evolves to full defection when individuals have two learning strategies, even if these learning strategies can in turn evolve. Section 3 provides an example for replicators vs imitators but other combinations can be worked out in a similar manner.
- When the population is placed on a square lattice, the existence of individuals with two different, permanent learning rules leads to dramatic changes with respect to the separate cases of the two strategies. Using again the example of replicators vs imitators (Subsection 4.1), a small proportion of replicators may lead to the breakdown of cooperation generically observed for imitators [Nowak and May, 1992].
- Evolvability of learning rules has crucial implications on the outcome of evolutionary game dynamics. Thus, Subsection 4.2 shows that replicator displaces imitators leading to a rapid decreasing of cooperation. In turn, imitators take over the fraction of global but stochastic imitators (MOR) and lead to a promotion of the cooperative behavior. The phenomenon, however, is not trivial, and regions where evolution leads to an outcome opposite to what is expected in general are also observed (e.g., Fig. 4, $1.25 < b < 4/3$).
- This competition process takes place on lattices under different environments. Section 5 shows that our results are robust against changes on the neighborhood considered (Moore instead of von Neumann) and against asynchronous dynamics. This indicates that our conclusions have a large degree of generality.
- We have shown that, in general, rules that lead to less frequent strategy changes (and consequently to their own update in our model) tend to be selected, although our simulations also prove that this is not the whole story (Section 6).
- Finally, as a general conclusion, our work makes it clear that modelling the emergence of cooperation in the Prisoner’s Dilemma on networks must go beyond the work done so far in terms of populations with a single, constant in time learning strategy. While by no means have we considered all possible rules or all possible networks, our simulations provide well-established evi-
idence that the presence of different learning rules and their own evolution may lead to unexpected phenomena, sometimes opposite to the observations available so far on unique-learning dynamics. Games with even more delicate equilibria structure such as Snowdrift or Stag-Hunt are likely to be affected in yet a stronger manner.

As a closing remark, we want to stress that, while we acknowledge the limited scope of the present report, we believe that our study opens the way to a much more complete analysis of the evolution of the update rules. It is clear that the present work asks for further research, regarding, e.g., the case in which three different update rules are present simultaneously. As we have advanced above, in this scenario much more complicated process may appear as the third rule helps other resist invasion by a dominant one. On the other hand, we have by no means exhausted the possible update rules, and a more thorough simulation program which would include more deterministic local rules is needed, in order to determine whether or not local rules invade global rules, or whether stochastic is better than deterministic. A specific case of relevance in social networks is that of reinforcement learning [Wang et al., 2008], which is much more complex than the rules considered here and might provide an interesting first step to extend our results to more realistic situations. It would also be interesting to consider different time scales for the update of strategies and the update of learning rules. Thus, it is conceivable that an agent only changes her update rule after copying the strategy of the same neighbour a certain number of times. Mutations in the learning rule can also be included, and even heterogeneity in the details of the rule (for instance, different proportionality factors in the REP rule). These and other extensions of the present work should include in addition the comparison of the results of different games beyond the PD. We envisage that such a programme would be extremely useful for clarifying the big picture of evolutionary game theory on graphs and its applications.

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References

Axelrod, R., Hamilton, W. D., 1981. The evolution of cooperation. Science 211, 1390–1396.
Crespi, B.J., 2001. The evolution of social behavior in microorganisms. Trends Ecol. Evol. 16, 178-183.
Darwin, C., 1871. The Descent of Man, and Selection in Relation to Sex. Murray, London.
Dugatkin, L.A., Mesterton-Gibbons, M., 1996. Cooperation among unrelated individuals: reciprocal altruism, by-product mutualism and group selection in fishes. BioSystems 37, 19-30.
Frick, T., Schuster, S., 2003. An example of the prisoners dilemma in biochemistry. Naturwissenschafte 90, 327-331.
Gardênes, G., Poncela, J., Floria, L.M., Moreno, Y., 2008. Natural Selection of cooperation and degree hierarchy in heterogeneous populations. J. Theor. Biol., in press (2008).
Gintis, H., 2000. Game Theory Evolving. Princeton University Press, Princeton.
Hammerstein, P., ed., 2003. Genetic and Cultural Evolution of Cooperation. Dahlem Workshop Report 90. MIT Press, Cambridge, MA.
Harley, C. B., 1981. Learning the evolutionarily stable strategy. J. Theor. Biol. 89, 611-633.
Hauert, C., 2002. Effects of space in 2×2 games. Int. J. Bif. Chaos 12, 1531-1548.
Hauert, C., 2006. Spatial effects in social dilemmas. J. Theor. Biol. 240, 627-636.
Hauert, C., Doebeli, M., 2004. Spatial structure often inhibits the evolution of cooperation in the Snowdrift game. Nature, 428, 643-646
Jiménez, R., Lugo, H., Cuesta, J. A., Sánchez, A., 2008. Emergence and resilience of cooperation in the spatial Prisoner’s dilemma via a reward mechanism. J. Theor. Biol. 250, 475-483.
Kiers, E.T., Rousseau, R.A., West, S.A., Denison, R.F., 2003. Host sanctions and the legume-rhizobium mutualism. Nature 425, 78-81.
Kirchkamp, O., 1999. Simultaneous evolution of learning rules and strategies. J. Econ. Behav. Org. 40, 295-312.
Kollock, P., 1998. Social dilemmas: The anatomy of cooperation. Annu. Rev. Sociol. 24, 183-214.
Langer, P., Nowak, M. A., Hauert, C., 2008. Spatial invasion of cooperation. J. Theor. Biol. 250, 634-641.
Lozano, S., Arenas, A., Sánchez, A., 2008. Mesoscopic Structure Conditions the Emergence of Cooperation on Social Networks. PLoS ONE 3(4): e1892.
Masuda, N., 2008. Oscillatory dynamics in evolutionary games are suppressed by heterogeneous adaptation rates of players. J. Theor. Biol. 251, 181-189.
Maynard-Smith, J., 1982. Evolution and the Theory of Games. Cambridge University Press, Cambridge.
Maynard-Smith, J., Szathmáry, E., 1995. The Major Transitions in Evolution. Oxford University Press, Oxford.

Nowak, M.A., 2006a. Five rules for the evolution of cooperation. Science 314, 1560-1563.

Nowak, M.A., 2006b. Evolutionary Dynamics. Harvard University Press, Cambridge, MA.

Nowak, M.A., May, R.M., 1992. Evolutionary games and spatial chaos. Nature 415, 424-426.

Nowak, M.A., May, R.M., 1993. The spatial dilemmas of evolution. Int. J. Bifurcat. Chaos 3, 35-78.

Ohtsuki, H., Nowak, M. A., 2006. Evolutionary games on cycles. Proc. R. Soc. B 273, 2249-2256.

Pennisi, E., 2005. Science 309, 93.

Rapoport, A., Chammah, A. M., 1965. Prisoner’s Dilemma. University of Michigan Press, Ann Arbor.

Roca C.P., Cuesta, J. A., Sánchez, A., 2008. Work in progress.

Santos FC, Pacheco JM, Lenaerts T (2006) Evolutionary dynamics of social dilemmas in structured heterogeneous populations. Proc. Natl. Acad. Sci. 103, 3490-3494.

Santos FC, Rodrigues JF, Pacheco JM (2006) Graph topology plays a determinant role in the evolution of cooperation. Proc, Roy, Soc, B 273, 51-55.

Sysi-Aho, M., Saramäki, J., Kertész, J., Kaski, K., 2005. Spatial snowdrift game with myopic agents. Eur. Phys. J. B 44, 129-135.

Szabó, G., Fáth, G., 2007. Evolutionary games on graphs. Phys. Rep. 446, 97-216.

Szabó, G., Szolnoki, A., Vukov, J., 2008. Application of Darwinian selection to dynamical rules in spatial Prisoners Dilemma games. Preprint (private communication from G. Szabó, available at http://www.mfa.kfki.hu/~szabo/szaborms.html).

Szolnoki, A., Szabó, G., 2007. Cooperation enhanced by inhomogeneous activity of teaching for evolutionary Prisoner’s dilemma games. Europhys. Lett. 77, 30004.

Szolnoki, A., Perc, M., Szabó, G., 2008. Diversity of reproduction rate supports cooperation in the prisoner’s dilemma game on complex networks. Eur. Phys. J. B 61, 505-509.

Wang, S., Szalay, M. S., Zhang, C., Csermely, P., 2008. Learning and innovative elements of strategy adoption rules expand cooperative network topologies. PLoS ONE 3, e1917.