

Learning to Cooperate: The Evolution of Social Rewards in Repeated Interactions

Slimane Dridi* and Erol Akçay

Department of Biology, University of Pennsylvania, 433 South University Avenue, Philadelphia, Pennsylvania 19104

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ABSTRACT: Understanding the behavioral and psychological mechanisms underlying social behaviors is one of the major goals of social evolutionary theory. In particular, a persistent question about animal cooperation is to what extent it is supported by other-regarding preferences—the motivation to increase the welfare of others. In many situations, animals adjust their behaviors through learning by responding to the rewards they experience as a consequence of their actions. Therefore, we may ask whether learning in social situations can be driven by evolved other-regarding rewards. Here we develop a mathematical model in order to ask whether the mere act of cooperating with a social partner will evolve to be inherently rewarding. Individuals interact repeatedly in pairs and adjust their behaviors through reinforcement learning. We assume that individuals associate with each game outcome an internal reward value. These perceived rewards are genetically evolving traits. We find that conditionally cooperative rewards that value mutual cooperation positively but the sucker's outcome negatively tend to be evolutionarily stable. Purely other-regarding rewards can evolve only under special parameter combinations. On the other hand, selfish rewards that always lead to pure defection are also evolutionarily successful. These findings are consistent with empirical observations showing that humans tend to display conditionally cooperative behavior and also exhibit a diversity of preferences. Our model also demonstrates the need to further integrate multiple levels of biological causation of behavior.

Keywords: reinforcement learning, natural selection, other-regarding preferences, altruism.

Introduction

In animals, repeated interactions often lead to mutual cooperation (Trivers 1971; Axelrod and Hamilton 1981; Wilkinson 1988; Lehmann and Keller 2006; Schneeberger et al. 2012; Stewart and Plotkin 2013). Because repeated interactions offer the opportunity for learning, there is growing interest in characterizing the learning mechanisms and in-

ternal social motivations that lead to cooperation. Recognizing that natural selection acts on those behavioral mechanisms (McNamara and Houston 2009; Hammerstein and Stevens 2012; Fawcett et al. 2013; Dridi and Lehmann 2014) rather than directly on the cooperative phenotypes themselves generates a new perspective on questions about the evolution of cooperation. In particular, an important question at the interface of psychological mechanisms and evolutionary theory is whether biological altruism requires or necessarily leads to other-regarding preferences. In other words, when we observe cooperation, is it because the individuals performing the cooperative act have other-regarding preferences, that is, they evolved motivations to provide a positive outcome for their social partners? This question about the proximate mechanisms underlying cooperation is important to understand both how individuals will behave in novel social and environmental contexts and how natural selection will shape the evolution of social traits (Akçay et al. 2009; Akçay and Van Cleve 2012; Van Cleve and Akçay 2014).

Several studies of cooperation in animals suggest that individuals may have other-regarding preferences (mostly in primates [Lakshminarayanan and Santos 2008; Brosnan et al. 2010; Chang et al. 2011; Claidière et al. 2015] but recently also in rats [Hernandez-Lallement et al. 2015]). However, other studies found that animals seem to pursue only their own personal gain (Silk et al. 2005; Jensen et al. 2006). In these experiments, animals are generally presented with the choice between a selfish option (obtaining a reward only for oneself) and a social option (providing a reward for both oneself and a partner), and a preference for the social option may be interpreted as an other-regarding preference. Evidence for such prosocial tendencies is also abundant in humans (Fehr and Gächter 2000; Henrich et al. 2001; Camerer 2003; Fehr and Fischbacher 2003; Chaudhuri 2010). Nonetheless, other researchers (Binmore 2005; Burton-Chellew et al. 2015, 2016) argue that apparently other-regarding behavior may be explained by the participants' not fully understanding the experiment's setup,

* Corresponding author; e-mail: slimane.dridi@gmail.com.

ORCID: Dridi, <http://orcid.org/0000-0002-0159-9911>; Akçay, <http://orcid.org/0000-0001-8149-7124>.

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combined with payoff-based learning during the course of the game.

A much greater number of empirical studies (Taborsky et al. 2016) indicate that reciprocal cooperation occurs in species as diverse as fish (Dugatkin and Alfieri 1991), birds (Voelkl et al. 2015), bats (Wilkinson et al. 2016), and primates (Schino and Aureli 2010). However, our understanding of the psychological motives underlying reciprocation remains limited. In particular, reciprocal cooperation can come about from simple reaction norms (McNamara et al. 1999), other-regarding preferences (Akçay et al. 2009), or learning from past rewards (Dridi and Lehmann 2015). In short, there is still disagreement about whether prosocial preferences combined with learning explain the cooperation that we observe in human and other animal societies. Answering this question requires explicit models that combine behavioral dynamics of learning with evolution of rewards, which is what motivates this article.

Much of standard economics and decision theory is built on the idea that individuals strive to maximize a quantity called “utility.” Likewise, one can show that in the long run, natural selection will cause agents to behave as if they are maximizing an appropriately constructed fitness function (Lehmann et al. 2015). These results, however, offer little guidance on what (if anything) individuals maximize proximately; that is, they do not provide specific psychological mechanisms that generate fitness-maximizing behavior. In particular, individuals might be selected to maximize individual fitness but do this through other-regarding preferences (Akçay et al. 2009). This idea has led to models of preference evolution, where individuals play a given game that has fitness consequences (material payoffs) but where each individual possesses an arbitrary utility function that is genetically determined (Ockenfels 1993; Güth 1995; Akçay et al. 2009; Akçay and Van Cleve 2012; Alger and Weibull 2013). This utility function itself then evolves according to the fitness consequences of the behaviors it generates. Importantly, both the fitness function and the utility function order the outcomes of the social interaction, but these two orderings may be different from each other.

The main result from preference evolution models is that if players can observe each other’s utility functions before choosing an action, then other-regarding preferences may be evolutionarily stable; otherwise, natural selection leads to a utility function that corresponds exactly to the fitness function (Ok and Vega-Redondo 2001; Dekel et al. 2007). This is the same principle that explains the evolution of green-beard genes, where cooperators recognize each other (Robson 1990). A common way for animals to achieve such recognition is repeated interactions where individuals’ behaviors and responses to each other’s behavior are informative of their preferences (Akçay et al. 2009; Akçay and Van Cleve 2012; Jordan et al. 2016). In-

teractions between relatives also have been shown to promote other-regarding preferences by interacting with such behavioral responses (Akçay and Van Cleve 2012) or recognition of partners (Alger and Weibull 2013).

At the same time, most previous theories that model preference evolution or try to explain cooperation in the laboratory do not take into account that the behavior of humans and other animals is modified by learning based on experienced rewards. Indeed, learning (or initial lack thereof) is usually presented as an alternative to prosocial preferences for explaining behavior in experiments (Binmore 2005). However, as with many social and nonsocial behaviors consistently produced by a species with a neural system, cooperative behavior must generate positive rewards (in the proximate sense, see below) for an individual (Pearce 2008; Shettleworth 2009; Dugatkin 2010; Schultz 2015). If cooperation is to be observed in those species, then the temporal sequence of cooperation must be consistent with known principles of learning (Sutton and Barto 1998). Moreover, very often in social settings there is uncertainty and variability regarding who is going to be one’s social partner (because of the frequency of types changes between generations and because of randomness in the matching process), in which case learning can allow an individual to adapt to its social partners on the timescale of its lifetime. In sum, a theory for the proximate mechanisms of human and animal cooperation is incomplete without accounting for learning at the same time. In a learning context, the question of whether animals have other-regarding preferences thus becomes, can the cooperative act in itself be rewarding?

One may define a reward as an event that generates a particular pattern of activation of neural circuits that induces positive feedback on behavior (Dickinson and Balleine 1994; Pearce 2008; Schultz 2015). Essentially, animals tend to repeat actions that are followed by rewards; this phenomenon constitutes the core of associative learning. Punishments, on the other hand, are stimuli that generate a negative feedback on behavior, whereby actions followed by punishments tend to be avoided in the future. Certain stimuli act as intrinsic rewards (also called primary rewards), thus allowing an animal to build associations between these intrinsic rewards and new actions or stimuli. Glucose is such an intrinsic reward in many animals: an animal can learn to associate glucose with another stimulus (e.g., a particular fruit) or with an action (e.g., in the laboratory, pulling a lever). Once learning has taken place, the associated stimulus (e.g., the fruit) or the associated action (e.g., pulling a lever) becomes a reward predictor (Niv 2009). Because brain regions involved in decision making and social cognition project to the mesolimbic reward system (Declerck et al. 2013), it is possible that the part of the brain responsible for social cognition activates this innate reward system. In other words, cooperation may be intrinsically rewarding in the brain.

There is evidence that this may be true in humans and other primates (Chang et al. 2015). Given the prevalence of cooperation (especially reciprocal cooperation; Taborsky et al. 2016) in many other species, the fact that cooperation is rewarding in the brain is likely to be widespread, although direct neurobiological evidence in other species is scarce. Thus, this basic reward system can be thought of as the proximate basis of learning in social interactions.

From an evolutionary perspective, intrinsic rewards can be viewed as proximate mechanisms that natural selection shapes to make individuals behave in ways that increase fitness. In many cases, intrinsic rewards could be direct proxies of material benefits, as in many models of the evolution of learning (Boyd and Richerson 1988; Josephson 2008; Hamblin and Giraldeau 2009; Arbilly et al. 2010; Katsnelson et al. 2011; Dridi and Lehmann 2014, where the reinforcement term in the equation describing learning is equated to incremental fitness effects). However, in social interactions, intrinsic rewards that are systematically different from one's own material gains can be evolutionarily stable (Ok and Vega-Redondo 2001; Dekel et al. 2007; Akçay et al. 2009; Akçay and Van Cleve 2012; Alger and Weibull 2013). Natural selection can shape the way social cognition can activate the mesolimbic reward system to take into account stimuli other than one's own material gain, such as social partners' payoffs or even abstract social concepts such as fairness and honor, if the resultant behavior is fitness enhancing. Such deviations from a direct mapping from material payoff to intrinsic rewards can evolve either through direct fitness benefits (e.g., because they generate behavioral feedbacks that benefit their carriers; Akçay et al. 2009) or through indirect fitness benefits (Akçay and Van Cleve 2012). These observations raise the question of how intrinsic rewards that drive learning in social interactions evolve.

In this article, we present a model of the evolution of such intrinsic rewards when individuals interact in the Prisoner's Dilemma game, where they have the choice between cooperation and defection. To capture general learning processes in humans and other animals, we model learning as a basic trial-and-error process where individuals repeat actions followed by rewards and avoid actions followed by punishments. In our model, individuals interact in games whose material payoffs determine fitness. Instead of learning according to the real material payoffs, an individual associates with each game outcome a genetically determined utility, which is used as the intrinsic reward/punishment for that particular outcome. For example, other-regarding individuals may associate positive utilities with outcomes where their partner obtains a positive material payoff and thus might learn to cooperate as an intrinsically rewarding action. This decoupling of material payoffs and rewards allows us to address the question of how rewards evolve in social interactions. We look for the evolutionarily stable utility functions

when individuals interact repeatedly in a game whose material one-shot payoffs determine the two-person Prisoner's Dilemma game.

Model

Social Interactions and Rewards

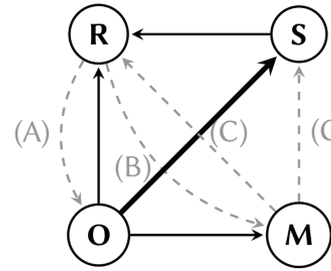
We consider an evolutionary model of repeated pairwise games in a large, well-mixed population of learners with nonoverlapping generations. Every generation of the evolutionary process consists of a sequence of interaction rounds, $t = 1, 2, \dots, T$. At each generation just before $t = 1$, individuals in the population are randomly matched in pairs, and each pair remains together for the entire duration of the game (until $t = T$). Hence, individuals are playing a repeated game with their partner. The one-shot game, played at every time t , is a Prisoner's Dilemma game with two possible actions: cooperate, C (or action 1), or defect, D (or action 2). The one-shot material payoffs for individual i are denoted $\pi_i(C, C) = b - c$, $\pi_i(C, D) = -c$, $\pi_i(D, C) = b$, $\pi_i(D, D) = 0$, where the first element in parentheses of $\pi_i(a_i, a_{-i})$ denotes player i 's action (a_i) and the second element denotes its opponent's action (a_{-i}). We assume also that $b > c > 0$. The sequence of material payoffs ultimately determines fitness (see below for details on how fitness is evaluated).

At every interaction round t , each individual in every pair chooses an action. Individual i 's action at time t is denoted $a_{i,t}$ and its opponent's action is $a_{-i,t}$. After actions are chosen, both players observe the outcome $(a_{i,t}, a_{-i,t})$ and subjectively evaluate how good the outcome was, which is genetically determined. We call this subjective evaluation the utility function of a player, which may be different from the actual material payoff, $\pi_i(a_{i,t}, a_{-i,t})$, obtained at time t . This utility (rather than the material payoff) determines the reward sensation of a game outcome, and this reward is used by an individual to learn its strategy in the repeated game (see below for details about the learning process). Specifically, the genotype of each individual i associates with each outcome (a_i, a_{-i}) a utility $u_i(a_i, a_{-i})$ that can take any negative or positive real value. We say that the utility is a reward if it is positive, $u_i(a_i, a_{-i}) > 0$, while we call it a punishment if $u_i(a_i, a_{-i}) < 0$. Hence, a genotype consists of the four utilities $u_i(C, C)$, $u_i(C, D)$, $u_i(D, C)$, and $u_i(D, D)$. We can arrange these four utilities in a matrix according to the game outcomes, which we call the utility matrix of individual i (fig. 1A). However, evolutionarily speaking, it is easier to think of these utilities as the vector $\mathbf{u}_i = (u_i(C, C), u_i(C, D), u_i(D, C), u_i(D, D))$; below we also use the more compact notation $\mathbf{u} = (u_{11}, u_{12}, u_{21}, u_{22})$, dropping the player's index. The state space is thus \mathbb{R}^4 . Our interest in this article is to find the evolutionarily stable utility vector \mathbf{u}^* .

A

	C	D		C	D
C	+	-	Realistic	-	-
D	+	0		+	+
	C	D		C	D
C	+	+	Other-regard	+	-
D	-	-		+	-
	C	D		C	D
C	+	-	Manipulator	+	-
D	+	-		+	-

B

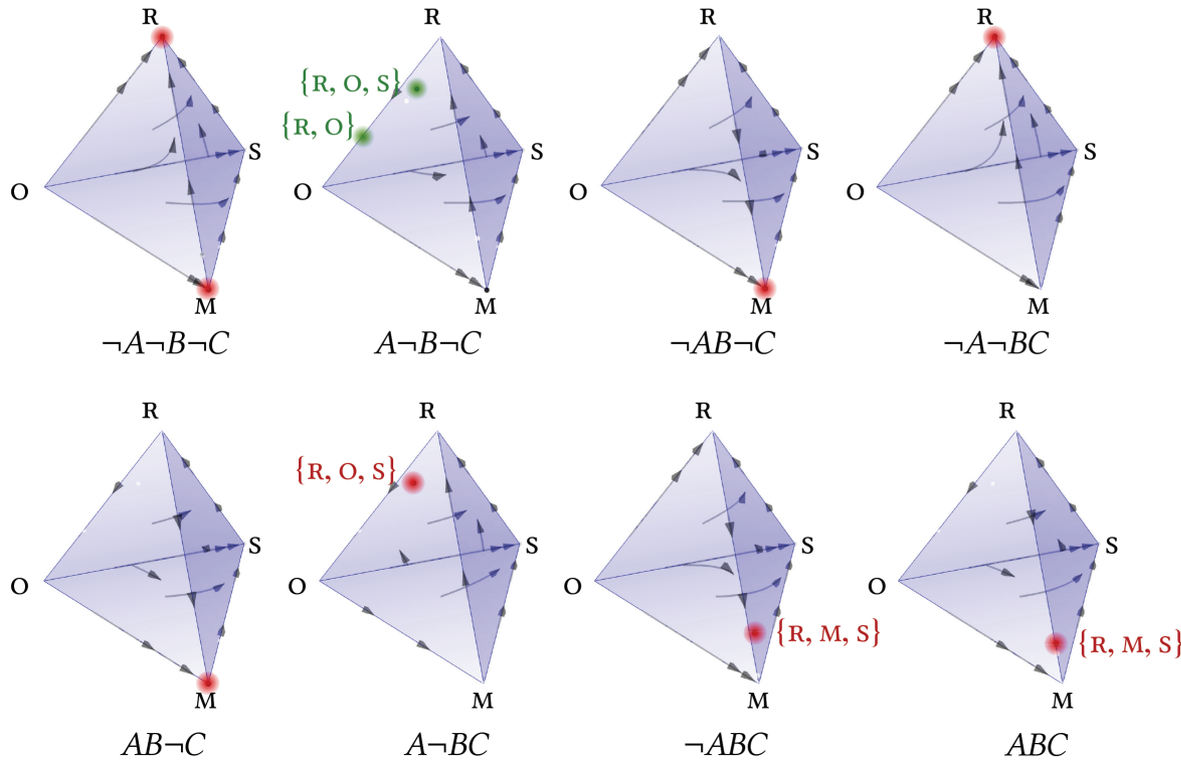


(A): $q_{RR} < q_{RO}$ and $\frac{b}{c} > \frac{1-q_{RR}}{q_{RO}-q_{RR}}$

(B): $q_{RR} < q_{RM}$ and $\left(\left(\frac{b}{c} \geq \frac{1-q_{RR}}{q_{RM}-q_{RR}} \right) \text{ or } \left(\frac{b}{c} < \frac{1-q_{RR}}{q_{RM}-q_{RR}} \text{ and } \frac{b}{c} + \frac{u_{12}}{u_{22}} > \frac{1-q_{RR}}{q_{RM}-q_{RR}} \right) \right)$

(C): $\frac{u_{22}}{u_{12}+u_{22}} b > b - c$

C



To do so, we need to know the fitness $f(\mathbf{u}_i)$ of an individual with utility \mathbf{u}_i . To arrive there, we first need to specify how the utility vectors of a pair of players determine behavior in the repeated game.

Learning

We assume that individuals learn to play the game according to a simple trial-and-error procedure. We use a standard model of learning dynamics (Sutton and Barto 1998; Dridi and Lehmann 2014), except that actions are reinforced according to the subjective utilities of a game outcome $u_i(\cdot)$, rather than the objective material payoff $\pi_i(\cdot)$. At every time t , an individual i holds in memory action values $V_{i,t}(a_i)$ for both actions $a_i \in \{C, D\}$ and chooses to cooperate at time t with a probability $p_{i,t}$ that depends on its action values $\{V_{i,t}(C), V_{i,t}(D)\}$. We adapted an existing model of the evolution of learning rules (Dridi and Lehmann 2015) so that the learning rule of individual i in our model is to update action values according to

$$V_{i,t+1}(a_i) = V_{i,t}(a_i) + \gamma_t \mathbf{1}(a_i, a_{i,t}) u_i(a_i, a_{-i,t}), \quad (1)$$

where $\mathbf{1}(a_i, a_{i,t})$ is an indicator variable that equals 1 if $a_i = a_{i,t}$ and 0 otherwise and $\gamma_t \in (0, 1)$ is a dynamic learning rate. This learning rate is decreasing as the game proceeds, which implies that the initial rounds of interaction are critical in determining the stable outcome of the learning process. Such a condition ensures that learning converges during an individual's lifetime (Benaïm 1999; Dridi and Lehmann 2014) and can be justified by the fact that the game faced by the individuals is constant. Finally, $u_i(a_i, a_{-i,t})$ is the utility to i if the individual plays a_i given that its opponent plays $a_{-i,t}$ at time t .

We then assume that individuals want to choose the action with the highest value, $V_{i,t}(a_i)$, but also have some tendency to explore the action with a smaller value. A widely

used choice rule to capture this principle is the logit-choice function,

$$p_{i,t}(a_i) = \frac{\exp[\xi V_{i,t}(a_i)]}{\sum_{b_i \in A_i} \exp[\xi V_{i,t}(b_i)]}, \quad (2)$$

where $\xi > 0$ is the exploration parameter (the inverse, $1/\xi$, can be seen as the noise level if we interpret this model as a perturbed maximization of action values; Hofbauer and Sandholm 2002) in choosing actions. In our case, there are two actions, C and D ; hence, equation (2) is a sigmoid function that can be thought of as a generalization of the threshold rule that chooses the action with greater value, $V_{i,t}(a_i)$. Equation (2) approaches such a threshold function when ξ becomes larger.

The behavioral interaction between a reinforcement learner with utilities $\mathbf{u} = (u_{11}, u_{12}, u_{21}, u_{22})$ and another reinforcement learner with utilities $\mathbf{v} = (v_{11}, v_{12}, v_{21}, v_{22})$ is what we need to analyze in order to compute fitness. By a slight abuse of notation, we denote these two players u and v and their probabilities to cooperate p_u and p_v , respectively. Stochastic approximation theory can be used (see, e.g., Dridi and Lehmann 2014) to show that the long-run learning dynamics (eqq. [1], [2]) for a pair of learners can be described as

$$\begin{aligned} \dot{p}_u &= p_u(1 - p_u) \xi [p_v \{p_u u_{11} + (1 - p_v) u_{12}\} \\ &\quad - (1 - p_u) \{p_v u_{21} + (1 - p_v) u_{22}\}], \\ \dot{p}_v &= p_v(1 - p_v) \xi [p_u \{p_v v_{11} + (1 - p_u) v_{12}\} \\ &\quad - (1 - p_v) \{p_u v_{21} + (1 - p_u) v_{22}\}]. \end{aligned} \quad (3)$$

Equation (3) displays 10 generic behavioral equilibria (fig. B1; figs. B1–B10 available online). Depending on the specific values of \mathbf{u} and \mathbf{v} , one or more of these equilibria may exist. Note that because the original dynamic is stochastic, when the corresponding deterministic system admits several locally

Figure 1: Replicator dynamics for the competition between Realistic, Other-regard, Manipulator, and Selfish. *A*, The four strategies considered in the analytical model. A strategy is defined by the outcomes to which it associates a positive or negative utility. The first row/column corresponds to cooperate and the second row/column to defect. Utilities are to row-player. *B*, Pairwise invasion diagram between the four strategies Realistic, Other-regard, Manipulator, and Selfish and associated invasion conditions. A plain directed edge from node X to node Y means that strategy Y always invades a monomorphic population of X (but does not necessarily reach fixation). A dashed directed edge from node X to node Y means that Y can invade X under certain conditions (A , B , and C) on the model parameters. When a given strategy can be invaded by more than one other strategy, a thick edge designates the best response. Note that all combinations of these three conditions are possible. *C*, Classification of phase portraits for the replicator dynamics in the four-strategy game defined by the competition between Realistic, Other-regard, Manipulator, and Selfish. Each subfigure is a drawing of the four-simplex (produced using Dynamio; Sandholm et al. 2012). At each vertex, one of the four strategies is at frequency 1: Realistic at the top, Manipulator at the bottom-front, Other-regard at the back left, and Selfish at the back right. The letters A , B , and C refer to the conditions in *B*, where the \neg sign denotes logical negation. For instance, the subfigure labeled $\neg ABC$ is drawn for parameter values such that condition A is not true but conditions B and C are true. Red dots denote locally stable equilibria, that is, possible outcomes of natural selection. To disambiguate the three-dimensional view, red labels in curly braces indicate the set of strategies present at an equilibrium. In the subfigure for the case $A \neg B \neg C$, the green dots are two alternative outcomes: $\{R, O\}$ occurs when Selfish cannot invade this polymorphism; if Selfish does invade this polymorphism, $\{R, O\}$ becomes unstable and $\{R, O, S\}$ stable. The condition for this to happen is given by equation (A11) in appendix A, available online.

stable equilibria, the stochastic dynamics may reach any of these equilibria. It turns out that the theory of stochastic approximations is almost silent about which particular equilibrium will be reached. These lock-in probabilities will, however, play an important role for the evolutionary stability of the different utility functions we will study below.

Another important fact about the behavioral dynamics is that the stability of the possible behavioral equilibria is very much dependent on the signs of utilities of the individuals involved in an interaction. In particular, a pure equilibrium is locally stable if and only if both players have a positive utility (making it a reward) for this outcome. The implication of this is that if at least one player has a negative utility for the outcome, then this outcome is unstable. In other words, if players u and v do not “agree” on preferred outcomes, then a pure behavioral equilibrium cannot be stable. This intuitive result is mathematically true because the eigenvalues of the Jacobian matrix associated with equation (3) evaluated at a pure outcome (i, j) are simply

$$\lambda_1 = -\xi u_{ij}, \lambda_2 = -\xi v_{ji}. \quad (4)$$

This fact has important evolutionary consequences, as will be detailed below when we analyze interactions between individuals with particular utility functions. In particular, it allows us to classify different utility functions by their sign for each of the four outcomes.

Fecundity

Assuming that interactions last long enough ($T \rightarrow \infty$), we define the fecundity of individual i as being proportional to the average material payoff obtained at equilibrium of the learning process, that is,

$$f_i = f(u_i) = \sum_{\mathbf{a} \in \mathcal{A}} \hat{\mathbf{p}}(\mathbf{a}) \pi_i(\mathbf{a}), \quad (5)$$

where $\hat{\mathbf{p}}(\mathbf{a}) = \hat{p}_i(a_i) \hat{p}_{-i}(a_{-i})$ is the equilibrium probability of outcome $\mathbf{a} = (a_i, a_{-i})$. The sum in equation (5) is taken over the set of possible game outcomes, $\mathcal{A} = \{(C, C), (C, D), (D, C), (D, D)\}$. We call $\hat{\mathbf{p}}$ the behavioral equilibrium. Importantly, while the utility function does not appear on the right-hand side of equation (5), we still define it as $f(u_i)$ because the equilibrium choice probabilities of a player, $\hat{p}_i(a_i)$, implicitly depend on the utility function of player i , as will become clearer when we derive expressions for the behavioral equilibria below.

The fecundity $f(\mathbf{u})$ depends on the outcome of the learning dynamics and is therefore not continuous in \mathbf{u} , which renders difficult a full analytic treatment of the model. To overcome this problem we adopt two complementary approaches. First, we focus on a smaller number of utility functions that are

relevant to our original question of the evolution of other-regarding preferences. Second, we run evolutionary simulations of the full model to have a more comprehensive view of our model.

Results

Analytical Results for Four-Strategy Competition

We first consider the evolutionary dynamics for the competition between the following four possible utility functions represented in figure 1A using the replicator dynamics (for details of the analysis, see app. A; apps. A, B available online).

The Realistic function associates with outcomes a utility of the same sign as the real material payoff. This type of utility function is the default utility function, used in virtually all models of the evolution of learning (Boyd and Richerson 1988; Josephson 2008; Hamblin and Giraldeau 2009; Arbilly et al. 2010; Katsnelson et al. 2011; Dridi and Lehmann 2014). It takes as a special case the material payoff function, that is, $u_i = \pi_i$. It is the function that evolves when interactions between players are completely anonymous, one shot, and there is no assortment in the matching process (Ok and Vega-Redondo 2001; Dekel et al. 2007).

The Other-regard function associates positive utility with the outcomes where the opponent obtains a strictly positive payoff. In other words, this strategy associates positive utilities only with the outcomes where it cooperates.

The Selfish function associates positive utility with the outcomes where it defects.

The Manipulator function associates positive utility only with the outcomes where its opponent cooperates. The name of this utility function stems from the fact that it will drive a compliant opponent (that associates positive utility with all outcomes) to cooperate.

We first construct the fitness matrix for the evolutionary game in table 1 by considering the stable equilibria of learning dynamics for all possible pairwise matchings between the four strategies (described in detail in app. A; see also figs. 2, B2). For the four strategies we consider in this section, no more than two behavioral equilibria are locally stable at the same time. It turns out that in all cases where two equilibria are locally stable, one of them is mutual cooperation, (1, 1) (figs. 2, B2). Because the underlying learning model is stochastic (eqq. [1], [2]), the lock-in probability in the cooperative equilibrium (1, 1) will affect the fitness and hence the evolutionary competition between the four strategies we are considering. However, there is no general technique to obtain an expression of the lock-in probability. At this point, we leave these probabilities unspecified and denote by q_{uv} the probability that an interaction between strategy u and strategy v leads to the cooperative

Table 1: Evolutionary fitness matrix among the four strategies considered in the analytical model

	R	O	M	S
R	$q_{RR}(b - c)$	$q_{RO}(b - c) + (1 - q_{RO})b$	$q_{RM}(b - c) + (1 - q_{RM})\left(b\left(\frac{u_{22}^M}{u_{11}^M + u_{22}^M}\right)\right)$	0
O	$q_{RO}(b - c) + (1 - q_{RO})(-c)$	$b - c$	$q_{OM}(b - c) + (1 - q_{OM})(-c)$	$-c$
M	$q_{RM}(b - c) + (1 - q_{RM})\left((-c)\left(\frac{u_{22}^M}{u_{11}^M + u_{22}^M}\right)\right)$	$q_{OM}(b - c) + (1 - q_{OM})b$	$b - c$	$-c\frac{u_{22}^M}{u_{11}^M + u_{22}^M}$
S	0	b	$b\frac{u_{22}^M}{u_{11}^M + u_{22}^M}$	0

Note: q_{ij} denotes the probability that a behavioral interaction between strategy i and strategy j leads to mutual cooperation at a behavioral equilibrium. The expression u_{ab}^i denotes the utility of strategy i for the game outcome where it chooses action a and its opponent chooses b . M = Manipulator; O = Other-regard; R = Realistic; S = Selfish.

equilibrium (1, 1). For instance, two Realistic individuals can learn mutual cooperation, (1, 1), or mutual defection, (0, 0). The probability that an interaction between two Realistic individuals leads to mutual cooperation is thus denoted q_{RR} ; the probability of locking in the defective equilibrium is $1 - q_{RR}$.

Evolutionary Dynamics for the Prisoner's Dilemma

We use the replicator dynamics (Taylor and Jonker 1978, eq. [A6] in app. A) to describe the competition between Realistic, Other-regard, Manipulator, and Selfish, with the evolutionary game given in table 1. Determining the outcome of the replicator dynamics is dependent on the parameters b (benefit to a receiver of a cooperative act) and c (cost of cooperating) and the lock-in probabilities in the cooperative equilibrium q_{RR} , q_{RO} , q_{RM} , q_{OM} , for the different behavioral interactions where several equilibria are locally stable.

We first find that although (Selfish, Selfish) is always a weak Nash equilibrium of the evolutionary game between the four strategies, regardless of parameter values, it is never evolutionarily stable (fig. 1; table 1). This is because Selfish gets invaded by Realistic, which learns to defect against Selfish but cooperates with itself. On the other hand, the strategy Other-regard is also always invaded by every other strategy in pairwise competitions, although it can be part of a mixed equilibrium, as we will see below. All other important results depend on the parameters of the model, and three basic conditions of the parameters help classify the possible evolutionary outcomes (conditions A, B, and C in fig. 1B). For certain parameter values, Realistic can be an evolutionarily stable strategy when the benefit-to-cost ratio is sufficiently low (conditions A and B in fig. 1B). Also, for other parameter values, Manipulator can be evolutionarily stable (condition C in fig. 1B). Note that these conditions are not mutually exclusive, so both Realistic and Manipulator can be evolutionarily stable at the same time (fig. 1C). When at least one Realistic or Manipulator is not evolutionarily stable, then we obtain poly-

morphic equilibria. In such polymorphic equilibria, we have either three strategies (there is an equilibrium with Realistic, Other-regard, and Selfish and an equilibrium with Realistic, Manipulator, and Selfish) or two strategies (Realistic and Other-regard), the common feature of these being that Realistic is always present (fig. 1C). We note here that Other-regard can be present only when Realistic is present. Moreover, according to condition A in figure 1B, Realistic should

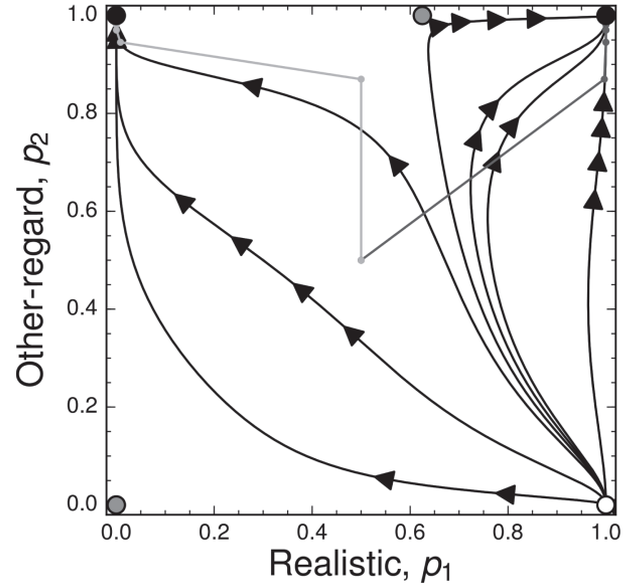


Figure 2: Solution trajectories (black lines) and stochastic trajectories (gray lines) for the behavioral interaction between Realistic and Other-regard. On the X-axis is represented the probability that Realistic cooperates (p_1), while on the Y-axis, this is the probability that Other-regard cooperates (p_2). The stochastic trajectories are started from the center of the state space $(p_1, p_2) = (1/2, 1/2)$, and dots on it represent interaction rounds between the players. Circles represent equilibria: a white-filled circle is a source (both associated eigenvalues are positive); a gray-filled circle is a saddle (one positive and one negative associated eigenvalue); a black circle is a sink (both associated eigenvalues are negative).

cooperate more often with Other-regard than with itself for the latter to make part of an equilibrium.

Simulations

Freely Evolving Utilities. The above analysis pertains to “internal stability” (see Eshel 1996) among a restricted class of utilities in the short term. However, utilities in our model comprise a four-dimensional vector of continuous evolutionary strategies. Therefore, it is natural to ask how populations, impelled by natural selection, move through such a strategy space in the long term. To do that, we conducted stochastic evolutionary simulations, in which we introduce new mutations from a much less constrained strategy space to a monomorphic population and use results from well-established population genetic theory (Van Cleve 2015) to calculate the invasion success of this mutant. This allows us to explore the strategy space in a computationally efficient manner.

In our simulations, the lock-in probabilities in behavioral equilibria, which played a critical role in determining the evolutionary outcome in the above four-strategy model, will no longer be parameters but will have a value that depends on the utilities of the particular strategies involved in behavioral interactions. Our evolutionary simulations consist of the trait substitution sequence of adaptive dynamics. Namely, we assume that the genotype of an individual, $\mathbf{u} = (u_{11}, u_{12}, u_{21}, u_{22})$, is supported by one locus and that the population is always monomorphic. At each iteration, we propose a mutation and determine whether the mutant invades the resident population using equations [41], [42] of Van Cleve (2015), calculated for Wright’s island model (in our case, the population is panmictic, or there is only one deme). We performed our evolutionary simulations for various values of the benefit-to-cost ratio, b/c , as well as different values of game duration, T .

To describe the results and in order to represent the four utilities at the same time, we classified all strategies according to the sign of their utilities (as we demonstrated above, these signs provide necessary conditions on the possible behavioral equilibria), which results in $2^4 = 16$ classes of strategies (because each of the four utilities has two possible signs). We can first look at the proportion of time a simulation run spends in each of the 16 strategy classes, which is an approximation of the stationary distribution of the evolutionary dynamics. We find that six strategies are consistently represented more than 10% of the time in the stationary distribution: Selfish, Avoid Sucker’s Payoff, Manipulator, Matcher, Pareto, and Anti-Cooperation. Avoid Sucker’s Payoff (AS) is similar to Realistic except that it has a positive utility for mutual defection instead of a 0; AS produces the same behavioral equilibria as Realistic when paired with other strategies (fig. B3). Matcher has positive

utilities only for outcomes where its own action matches that of its opponent; thus, the pure outcomes it can learn are mutual cooperation or mutual defection. Pareto has positive utility only for the outcome of mutual cooperation; thus, it will never learn full defection and will learn mutual cooperation against any opponent who is also willing to do so. Finally, Anti-Cooperation is the exact opposite of Pareto, as it has positive utilities for all outcomes except for mutual cooperation; this utility matrix cannot learn mutual cooperation and generally learns to defect but may be exploited by exploiting strategies such as Manipulator (fig. B3). In figure 3, we show results for various benefit-to-cost ratios, b/c , which lead to two main observations.

The first observation is that our simulations confirm the overall pattern in the analysis of the replicator dynamics, where at low values of b/c the strategy AS (corresponding to the Realistic strategy in the analytical model) experiences few invasions. As b/c increases, more strategies are able to invade AS, and, consequently, the frequency of AS declines (figs. 3, 4). In particular, if we analyze the invasions between the six dominant strategies in our simulations (fig. B4), we find that Manipulator, Matcher, and Pareto invade AS only for sufficiently high b/c . All these strategies have a positive utility for mutual cooperation; they also have a negative utility for the sucker’s outcome ($u_{12} < 0$). The success of AS and of cooperative strategies more generally yields an average utility matrix of the AS type (fig. 3), where average utilities are ordered as $\bar{u}(D, D) > \bar{u}(D, C) > \bar{u}(C, C) > \bar{u}(C, D)$, which is different from the ordering of the material payoffs, $\pi(D, C) > \pi(C, C) > \pi(D, D) > \pi(C, D)$. The strategy Pareto increases in frequency in the stationary distribution for increasing b/c (fig. 4A), as the analysis shows that it invades AS for high enough b/c (fig. B4). Strategies that are able to invade AS (Manipulator, Matcher, Pareto) can mutually invade one another, and we indeed observe that an important number of invasions occur between AS, Manipulator, Matcher, and Pareto (fig. 3). As a consequence of the increasing success of strategies that positively value cooperation as a function of b/c , we observe that the overall cooperation frequency in the population increases for increasing b/c (fig. 4B). Even though previous work has shown that cooperative strategies in the iterated Prisoner’s Dilemma can be evolutionarily robust (Stewart and Plotkin 2013), we could not expect this for the particular type of learning strategies that we have decided to study.

When we contrast the apparent success of conditionally cooperative strategies, a second major feature of our simulations becomes the success of the Selfish strategy. For all b/c , the simulation spends approximately 15%–20% of the time in this strategy class, and for high b/c , this makes Selfish the most represented strategy class in the stationary distribution (because of the decline of AS; figs. 3, 4A). Although this result could not be anticipated from our analysis of the

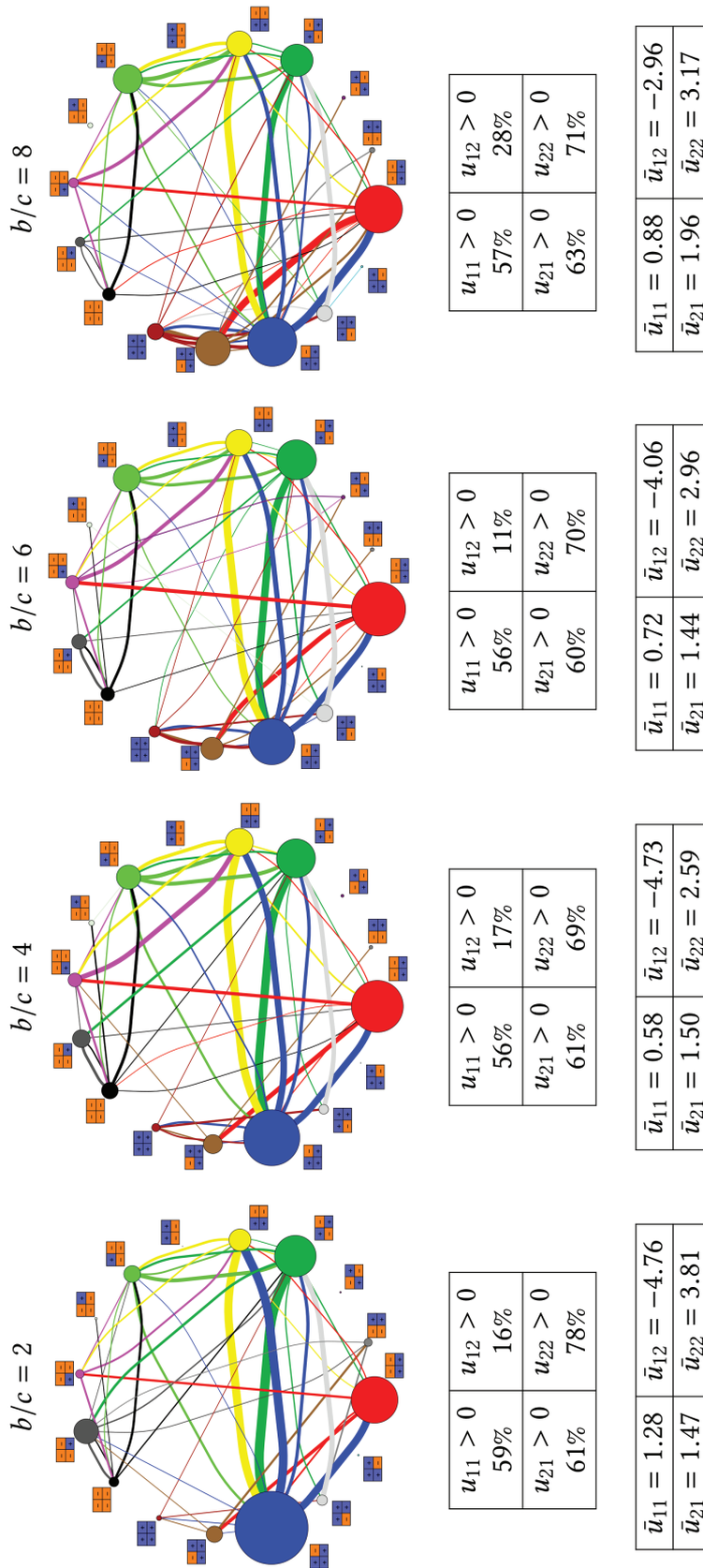


Figure 3: Invasion analysis and utilities in simulations for various benefit-to-cost ratios (b/c). The top row shows the invasion graph between the 16 classes of strategies defined by their utilities' sign (see main text), the second row shows the proportion of time each utility was positive in a simulation run, and the third row shows the time average of utilities. In the invasion graph, near each node we show the utility matrix of the corresponding strategy, with blue cells indicating a positive utility and orange cells indicating a negative utility. The utility matrix is oriented as in figure 1. The size of the nodes is proportional to the amount of time a simulation run spends in the corresponding strategy class. The edges are colored according to the invader strategy and thus indicate the direction of the edges. Edge thickness is proportional to the number of invasions that occurred between a pair of strategies (and we do not show edges between pairs of strategies for which the number of invasions was less than 10). Parameter values: min $u = -10$; max $u = 10$; $\xi = 2$; $c = 1$; $T = 150$; population size $N = 2,000$.

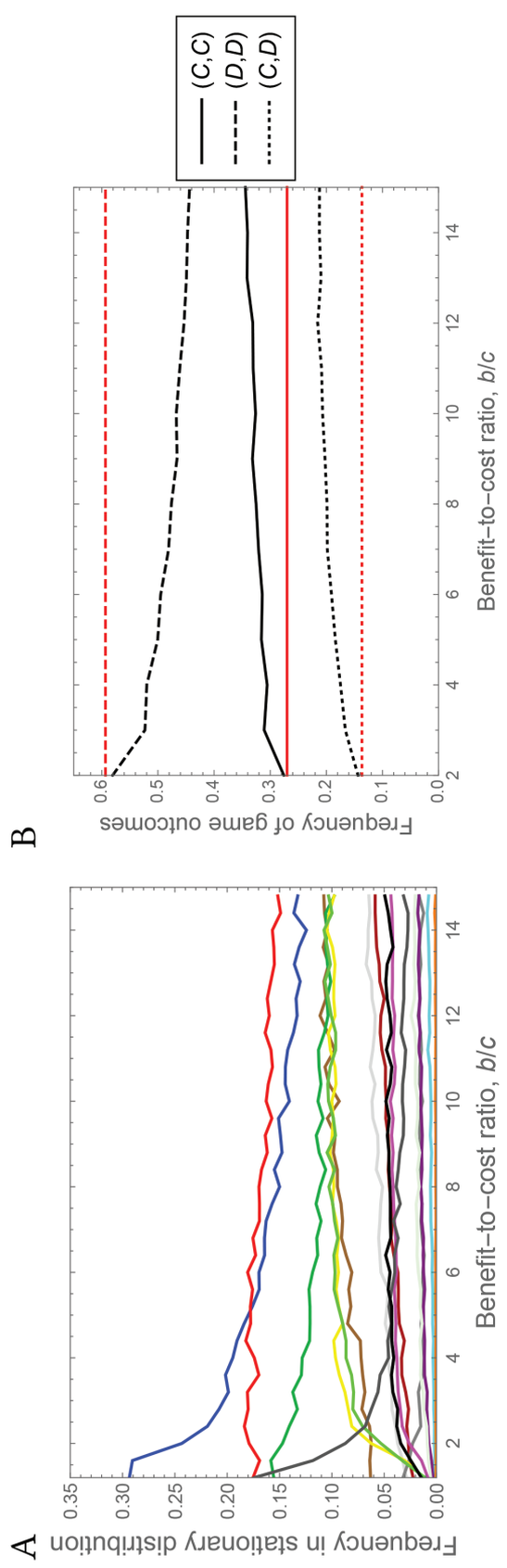


Figure 4: Effect of the benefit-to-cost ratio, b/c , on the stationary distribution of strategies and effective cooperation. *A*, Proportion of time the simulation spends in each of the 16 strategy classes as a function of b/c , which is a measure of the stationary distribution. Colors of strategies are as in figure 3 (see also fig. B5, available online). *B*, Time average of the frequency of game outcomes in a simulation run as a function of b/c (black lines; note that (C, D) and (D, C) are the same outcome). The red lines show the expected frequency of the corresponding game outcomes in the population if we draw strategies randomly from a uniform distribution. Parameter values are as in figure 3.

replicator dynamics above, it is still consistent with the fact that Selfish was relatively stable (only Realistic could invade it). Analytically considering the invasion conditions between the six dominant strategy classes in the simulations (fig. B4) reveals that Selfish is also relatively stable in this set, with only AS and Matcher being able to invade it. In our simulations, AS invades more frequently Selfish than Matcher does because, with our mutation scheme, an AS mutant is much more likely than a Matcher mutant to occur in a Selfish population, given that we draw mutations from a doubly exponential distribution centered at the resident phenotype.

A final observation regarding strategy classes is that Anti-Cooperation is relatively successful for high b/c (fig. 4A). Our invasion analysis in figure B4 shows that this strategy, despite having a positive utility for the sucker's outcome (where the individual cooperates and its opponent defects), compensates by exploiting certain cooperative strategies, such as Pareto. At low b/c , Anti-Cooperation gets exploited by strategies that have positive utilities for defection (such as AS or Selfish; fig. B3), but as b/c increases, Anti-Cooperation becomes more stable against these strategies, which explains why it makes part of an important proportion of the stationary distribution of the evolutionary dynamics.

In order to verify whether these results were sensitive to the length of the repeated game, we ran additional simulations for lower values of T . Our analytical results were obtained under the assumption that T is large enough so that learning reaches an equilibrium during individuals' lifetimes, and our standard simulations were run for $T = 150$. When using $T = 50$, we essentially obtained the same results in terms of the stationary distribution of strategies (fig. B9A, B9B). We needed to decrease the duration of the game to $T = 10$ to obtain different results (fig. B9C, B9D). Namely, in this case we find that Selfish is the mode of the stationary distribution for all benefit-to-cost ratios. Otherwise, we observe a similar pattern as for higher T values, with AS being represented more than other strategies for low b/c but slowly decreasing as b/c increases (fig. B9C). For this low T value, we also observe that the strategies Manipulator, Matcher, Pareto, and Anti-Cooperation, which previously grew in frequency for increasing b/c , still do so. The apparent success of Selfish for low T is due to the fact that for this duration of the game, learning cannot reach an equilibrium for all strategies and strategies that can learn multiple stable behavioral outcomes may wander between equilibria. In contrast, Selfish can only learn defection, irrespective of the opponent, and its convergence to the equilibrium occurs faster. Hence, the strategies that previously (i.e., for $T = 150$ or $T = 50$) succeeded in cooperating with cooperators but defecting with Selfish now fail to learn fast defection against Selfish. In appendix B we illustrate this phenomenon for interactions between the AS strategy and Selfish (fig. B10).

Utilities That Explicitly Depend on Material Payoffs. In this subsection we perform additional simulations by constraining the utility function to be dependent on the material payoffs of the focal player and its opponent. This allows us to address more directly the question of whether (and, if any, what type of) other-regarding preferences evolve in our model. Specifically, for any game outcome $\mathbf{a} = (a_i, a_{-i})$, we consider utility functions of the form

$$u_i(\mathbf{a}) = \pi_i(\mathbf{a}) + \beta(\pi_i(\mathbf{a}) + c + k)(\pi_{-i}(\mathbf{a}) + c + k) + \alpha\pi_{-i}(\mathbf{a}) + \gamma|\pi_i(\mathbf{a}) - \pi_{-i}(\mathbf{a})|, \quad (6)$$

where (α, β, γ) are player i 's genetically determined parameters. Here we will be interested in the evolution of these three parameters. In equation (6), c is the negative of the sucker's payoff ($-c$) and is added to the realized payoff to ensure that the term multiplied by β is always positive. The parameter k is here to allow the utility to vary as a function of β . Our utility function then measures the extent to which an individual is "additively" other-regarding ($\alpha \in [-1, 1]$), the extent to which the individual is "multiplicatively" other-regarding ($\beta \in [-1, 1]$), and inequity aversion ($\gamma \in [-1, 1]$). Even though this utility function can realize all of the 16 possible utility matrices discussed above, the structure of the phenotype space changes as compared to the above simulations where we let the utility matrix evolve in an unconstrained way (fig. B5).

Our simulations with the utility function in equation (6) show that the selection pressure on other-regarding preferences increases with b/c (figs. 5A, B8). The average value of β is close to 0 for low enough b/c but suddenly increases at a threshold value of b/c . For these higher b/c values, the average β is approximately 0.5, indicating the evolution of multiplicative Other-regard. The average values of α and γ are negative for low b/c , indicating, respectively, a combination of competitive preferences (valuing negatively others' success) and inequity aversion. Both α and γ decrease in magnitude as b/c increases but remain negative. This is a consequence of the fact that the selection pressure on α and γ decreases with increasing b/c , because the absolute difference between the temptation to defect, b , and the sucker's payoff, $-c$, decreases. This pattern is accompanied by a general increase in the utility for mutual cooperation as a function of b/c (fig. B7A). For high b/c , mutual cooperation becomes the preferred outcome of the evolutionarily stable utility function and mutual defection the least preferred outcome. In agreement with the above simulations for freely evolving utilities, AS is the dominant utility matrix for low b/c . The Compliant utility matrix (with all four utilities positive) becomes the dominant one for high b/c (fig. B7B).

Even though other-regarding preferences evolve for sufficiently high b/c , this is not accompanied by the evolution of increased effective mutual cooperation, even though the

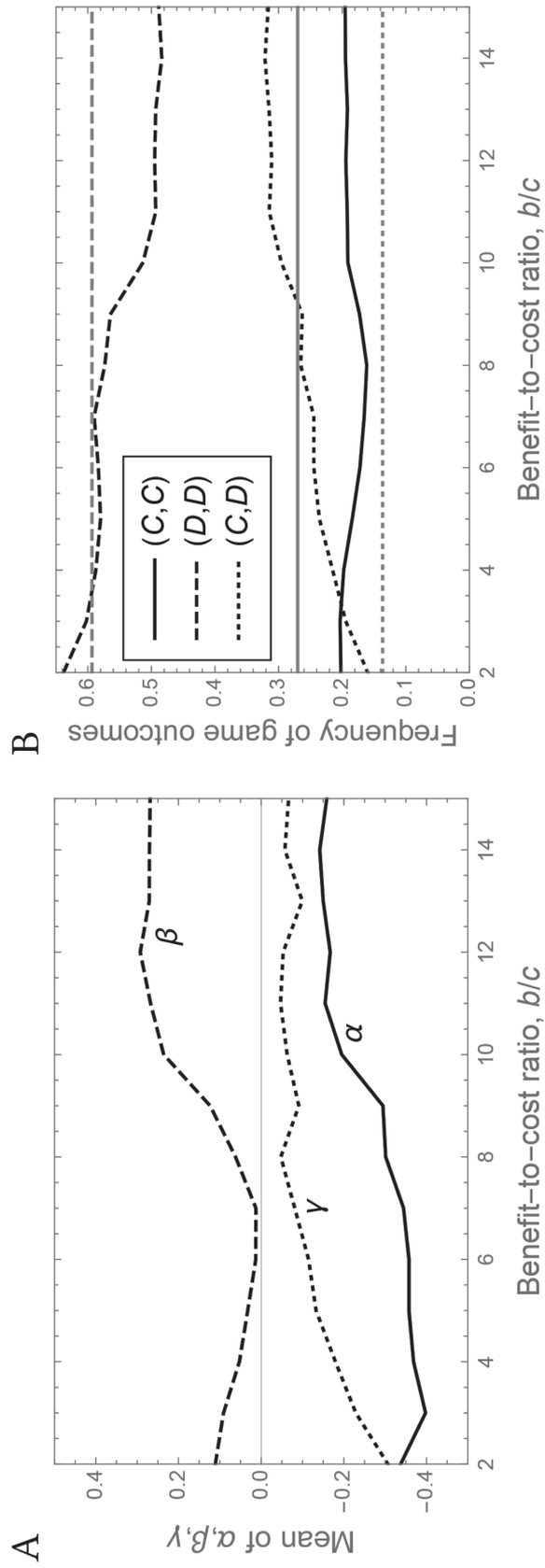


Figure 5: Results for the model where the utility depends explicitly on material payoffs (eq. [6] with $k = 2$) as a function of the benefit-to-cost ratio, b/c . A, Time average of α , β , γ in a simulation run. B, Time average of the frequency of game outcomes in a simulation run (similar to fig. 4).

frequency of mutual defection decreases. This decrease in mutual defection is due to an increase in the asymmetric (C, D) outcome (fig. 5B). Overall cooperation is thus increasing, but individuals do not coordinate on cooperating at the same time. This can be explained by the fact that the Compliant utility matrix that evolves for high b/c can learn any outcome (all pure equilibria are stable when all utilities are positive). However, the fact that mutual defection is the least preferred outcome implies that the probability to learn this equilibrium will be the lowest of the four outcomes.

Discussion

Evolution of Rewards for Prosocial Learning

We presented a model of how intrinsic rewards that drive learning in social interactions evolve. Rewards capture the intrinsic preferences of individuals over states of the world and constitute the fundamental building block of reinforcement learning. Because all behaviors are in part influenced by learning, modeling the evolution of social behaviors in animals requires that we take into account how behavior is generated through learning within an individual's life span. Within this framework, we developed a model to provide insight into the question of whether other-regarding preferences support the evolution of cooperation, under the constraint of reinforcement learning. While previous theoretical work tended to either ignore or oversimplify learning mechanisms (Ok and Vega-Redondo 2001; Dekel et al. 2007; Akçay et al. 2009; Akçay and Van Cleve 2012; Alger and Weibull 2013), our model tries to account for the increasing empirical evidence that reward processing and learning are critical aspects of prosocial preferences (Fehr and Camerer 2007; Declerck et al. 2013; Ruff and Fehr 2014). Overall, our results indicate that multiple preference functions can be evolutionarily stable when individuals interact repeatedly in the Prisoner's Dilemma. In particular, we find that evolutionarily successful preferences are of two general types: (1) those that have a positive utility for mutual cooperation but a negative utility for being exploited and (2) selfish preferences that associate positive utilities with outcomes where their carriers defect and that have negative utility for cooperation. This is true in both our analytical results in replicator dynamics and the numerical simulations in the whole strategy set. Further simulations show that other-regarding preferences evolve for a sufficiently high benefit-to-cost ratio.

A majority of the empirical evidence for the existence of other-regarding preferences comes from experiments performed by economists with human participants. Economic theory relies on the concept of utility to capture behavior, but the utility function of an individual is, by definition, an internal construct that is difficult to access (Fehr and Camerer 2007). In the context of learning, utility can be equated to re-

ward, because rewards are at the core of repeated behaviors (Schultz 2015). Empirically, one way to try to access the utility or reward function is to observe the pattern of activation in the brain when individuals make decisions. One of our main findings is that positive preferences for cooperation are evolutionarily prevalent. This finding is interesting when paralleled with neurobehavioral studies of social decision making that reveal that cooperation can generate rewards in the human brain, which seems consistent with the positive utility of winning strategies for mutual cooperation found in our model (Fehr and Camerer 2007; Declerck et al. 2013; Ruff and Fehr 2014). Additional empirical and theoretical work focusing on the cooperative behavior, but not on the preferences generating it, has already been conducted based on the premise that individuals may act prosocially in order for others to recognize their willingness to cooperate (Gintis et al. 2001; Jordan et al. 2016) or that cooperation could rely on fast decision making, implying a possible intrinsic preference for cooperation (Rand et al. 2012; Tinghög et al. 2013). Although these studies do not directly measure or model social preferences, they are, in principle, compatible with evolved preferences that find cooperation intrinsically rewarding. More generally, our results suggest a possible psychological mechanism for reciprocal cooperation in other animal species (Taborisky et al. 2016). Indeed, the evolutionarily stable utility functions in our model that positively value mutual cooperation produce behavioral dynamics that resemble the dynamics of reciprocal strategies such as tit for tat. It will be interesting in future empirical research to test whether these many examples of reciprocal cooperation may be based on learning combined with psychological preferences that value cooperation.

While our model shows that evolution can lead to intrinsically rewarding mutual cooperation, such utilities do not necessarily correspond to pure other-regarding preferences. For low benefit-to-cost ratios, competitive preferences that value others' payoff negatively tend to evolve. In contrast, for a sufficiently high benefit-to-cost ratio, we see the evolution of conditional (multiplicative) other-regarding preferences, in agreement with previous results that found these preferences to be evolutionarily stable in continuous social dilemmas (Akçay et al. 2009). On the other hand, one could interpret our results for the freely evolving utilities as reflecting the evolution of the correct representation of the real fitness effect of mutual cooperation, because mutual cooperation generates a positive effect on fitness. However, the Realistic utility function is not the only evolutionarily successful one in our model. For example, some evolutionarily successful preference functions value positively both mutual cooperation and mutual defection. These signs, together with a negative utility for the sucker's outcome, guarantee uninvadability by the Selfish preference function, because individuals with such preferences will learn to defect against Selfish. Moreover, on average, the utilities for the four different outcomes are or-

dered differently from the real material payoffs (e.g., mutual defection is the outcome with the highest utility, on average, while the real material payoff for this outcome is only the third material payoff). Therefore, our results do not show that natural selection leads to the correct representation of fitness effects in the brain in the context of learning. Another important distinction is that even though the utility for the temptation to defect is the highest in the model with freely evolving utilities, this does not necessarily mean that there are no other-regarding preferences: in our simulations where the utility is a function of payoffs, Other-regard (e.g., a positive β) can evolve even if the values of other evolutionary parameters make the temptation outcome more rewarding than mutual cooperation.

Our finding that reward representations in the brain do not necessarily correspond to real fitness effects adds to a growing realization that natural selection can shape decision-making mechanisms to have specific biases in different ecological situations (e.g., McNamara et al. 2013). Generally, perceptual systems that represent the world accurately may not be evolutionarily stable, contrary to a naive understanding of the workings of natural selection (Mark et al. 2010). In our scenario, the mismatch occurs because the game is repeated while an individual represents in its mind only the one-shot version of the game. It will be interesting in the future to examine whether our results obtained for the Prisoner's Dilemma extend to other repeated games.

Diversity in Preferences

Another main finding from our model is that a diversity of utility functions can be evolutionarily favored. This result is consistent with empirical findings that humans in behavioral experiments show behavioral diversity. In particular, strategies that value cooperation positively can produce behavior similar to that of reciprocating strategies (repeating the action of the partner in the previous round), and Selfish can produce the behavior of noncooperators; these two behavioral types have recently been found to represent the action sequence of many human participants in laboratory experiments (Fischbacher et al. 2001; Burton-Chellew et al. 2016) and have been considered as plausible evolutionarily significant behavioral rules in theoretical models (Trivers 1971; Axelrod and Hamilton 1981; Lehmann and Keller 2006; Stewart and Plotkin 2013). Moreover, in addition to a diversity of preference types, our model also shows the potential for multiple behavioral outcomes in a population monomorphic for a given preference function. This is because of the fact that stochastic learning processes can converge to different equilibrium profiles, which provides another potential explanation for the behavioral variation observed in learning experiments (Chmura et al. 2012). That a single decision rule can produce behavioral polymorphism is a result that

has been previously obtained in other models focusing on the evolution of cognitive mechanisms (Dridi and Lehmann 2015). This result illustrates that by modeling the evolution of the decision rules rather than the behaviors themselves (McNamara and Houston 2009; Hammerstein and Stevens 2012; Fawcett et al. 2013; Dridi and Lehmann 2014), one can account for richer behavioral patterns and potentially provide insights into the psychological underpinnings of social behavior. Our model can indeed be viewed as capturing variations in the decision rules because by changing the utility function of an individual, the updating rule for action values also changes (see eq. [1]), which subsequently produces different behavioral dynamics. These types of models require a detailed integration of two timescales (behavioral and evolutionary dynamics) and are consequently more difficult to analyze, but this difficulty cannot be avoided in trying to represent animal behavior more realistically.

In conclusion, our model articulates four levels of determinants of behavior: (1) the biological rewards at the core of brain functioning, (2) the psychological preferences that determine which states of the world are rewarding, (3) the social interactions that affect changes in the states of the world, and (4) the biological process of natural selection determining which behavioral mechanisms prevail in an evolving population. We find that evolution of rewards for learning captures both the possibility of cooperation and a diversity of individual preferences that can be evolutionarily successful. These results show the promise of integrating learning based on evolving intrinsic rewards from social interactions as a proximate mechanism for understanding the nature of cooperation in humans and other animals.

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Literature Cited

- Akçay, E., and J. Van Cleve. 2012. Behavioral responses in structured populations pave the way to group optimality. *American Naturalist* 179:257–269.
- Akçay, E., J. Van Cleve, M. W. Feldman, and J. Roughgarden. 2009. A theory for the evolution of other-regard integrating proximate and ultimate perspectives. *Proceedings of the National Academy of Sciences of the USA* 106:19061–19066.
- Alger, I., and J. W. Weibull. 2013. *Homo moralis*—preference evolution under incomplete information and assortative matching. *Econometrica* 81:2269–2302.

- Arbilly, M., U. Motro, M. W. Feldman, and A. Lotem. 2010. Coevolution of learning complexity and social foraging strategies. *Journal of Theoretical Biology* 267:573–581.
- Axelrod, R., and W. D. Hamilton. 1981. The evolution of cooperation. *Science* 211:1390–1396.
- Benaïm, M. 1999. Dynamics of stochastic approximation algorithms. Pages 1–68 in *Séminaire de probabilités XXXIII*. Vol. 1709. J. Azéma, M. Emery, P.-A. Meyer, and M. Yor, eds. Springer, Berlin.
- Binmore, K. 2005. Economic man—or straw man? *Behavioral and Brain Sciences* 28:817–818.
- Boyd, R., and P. J. Richerson. 1988. *Culture and the evolutionary process*. University of Chicago Press, Chicago.
- Brosnan, S. F., C. Talbot, M. Ahlgren, S. P. Lambeth, and S. J. Schapiro. 2010. Mechanisms underlying responses to inequitable outcomes in chimpanzees, *Pan troglodytes*. *Animal Behaviour* 79:1229–1237.
- Burton-Chellew, M. N., C. E. Mouden, and S. A. West. 2016. Conditional cooperation and confusion in public-goods experiments. *Proceedings of the National Academy of Sciences of the USA* 113:1291–1296.
- Burton-Chellew, M. N., H. H. Nax, and S. A. West. 2015. Payoff-based learning explains the decline in cooperation in public goods games. *Proceedings of the Royal Society B* 282:20142678.
- Camerer, C. F. 2003. *Behavioral game theory: experiments in strategic interaction*. Princeton University Press, Princeton, NJ.
- Chang, S. W. C., N. A. Fagan, K. Toda, A. V. Utevsy, J. M. Pearson, and M. L. Platt. 2015. Neural mechanisms of social decision-making in the primate amygdala. *Proceedings of the National Academy of Sciences of the USA* 112:16012–16017.
- Chang, S. W. C., A. A. Winecoff, and M. L. Platt. 2011. Vicarious reinforcement in rhesus macaques (*Macaca mulatta*). *Decision Neuroscience* 5:27.
- Chaudhuri, A. 2010. Sustaining cooperation in laboratory public goods experiments: a selective survey of the literature. *Experimental Economics* 14:47–83.
- Chmura, T., S. J. Goerg, and R. Selten. 2012. Learning in experimental 2 by 2 games. *Games and Economic Behavior* 76:44–73.
- Claidière, N., A. Whiten, M. C. Mareno, E. J. E. Messer, S. F. Brosnan, L. M. Hopper, S. P. Lambeth, S. J. Schapiro, and N. McGuigan. 2015. Selective and contagious prosocial resource donation in capuchin monkeys, chimpanzees and humans. *Scientific Reports* 5:7631.
- Declerck, C. H., C. Boone, and G. Emonds. 2013. When do people cooperate? the neuroeconomics of prosocial decision making. *Brain and Cognition* 81:95–117.
- Dekel, E., J. C. Ely, and O. Yilankaya. 2007. Evolution of preferences. *Review of Economic Studies* 74:685–704.
- Dickinson, A., and B. Balleine. 1994. Motivational control of goal-directed action. *Animal Learning and Behavior* 22:1–18.
- Dridi, S., and L. Lehmann. 2014. On learning dynamics underlying the evolution of learning rules. *Theoretical Population Biology* 91:20–36.
- . 2015. A model for the evolution of reinforcement learning in fluctuating games. *Animal Behaviour* 104:87–114.
- Dugatkin, L. A. 2010. *Principles of animal behavior*. 2nd ed. Norton, New York.
- Dugatkin, L. A., and M. Alfieri. 1991. Tit-for-tat in guppies (*Poecilia reticulata*): the relative nature of cooperation and defection during predator inspection. *Evolutionary Ecology* 5:300–309.
- Eshel, I. 1996. On the changing concept of evolutionary population stability as a reflection of a changing point of view in the quantitative theory of evolution. *Journal of Mathematical Biology* 34:485–510.
- Fawcett, T. W., S. Hamblin, and L.-A. Giraldeau. 2013. Exposing the behavioral gambit: the evolution of learning and decision rules. *Behavioral Ecology* 24:2–11.
- Fehr, E., and C. F. Camerer. 2007. Social neuroeconomics: the neural circuitry of social preferences. *Trends in Cognitive Sciences* 11:419–427.
- Fehr, E., and U. Fischbacher. 2003. The nature of human altruism. *Nature* 425:785–791.
- Fehr, E., and S. Gächter. 2000. Cooperation and punishment in public goods experiments. *American Economic Review* 90:980–994.
- Fischbacher, U., S. Gächter, and E. Fehr. 2001. Are people conditionally cooperative? evidence from a public goods experiment. *Economics Letters* 71:397–404.
- Gintis, H., E. A. Smith, and S. Bowles. 2001. Costly signaling and cooperation. *Journal of Theoretical Biology* 213:103–119.
- Güth, W. 1995. An evolutionary approach to explaining cooperative behavior by reciprocal incentives. *International Journal of Game Theory* 24:323–344.
- Hamblin, S., and L.-A. Giraldeau. 2009. Finding the evolutionarily stable learning rule for frequency-dependent foraging. *Animal Behaviour* 78:1343–1350.
- Hammerstein, P., and J. R. Stevens, eds. 2012. *Evolution and the mechanisms of decision making*. MIT Press, Cambridge, MA.
- Henrich, J., R. Boyd, S. Bowles, C. Camerer, E. Fehr, H. Gintis, and R. McElreath. 2001. In search of *Homo economicus*: behavioral experiments in 15 small-scale societies. *American Economic Review* 91:73–78.
- Hernandez-Lllement, J., M. van Wingerden, C. Marx, M. Srejjic, and T. Kalenscher. 2015. Rats prefer mutual rewards in a prosocial choice task. *Frontiers in Neuroscience* 8:443.
- Hofbauer, J., and W. H. Sandholm. 2002. On the global convergence of stochastic fictitious play. *Econometrica* 70:2265–2294.
- Jensen, K., B. Hare, J. Call, and M. Tomasello. 2006. What's in it for me? self-regard precludes altruism and spite in chimpanzees. *Proceedings of the Royal Society B* 273:1013–1021.
- Jordan, J. J., M. Hoffman, M. A. Nowak, and D. G. Rand. 2016. Uncalculating cooperation is used to signal trustworthiness. *Proceedings of the National Academy of Sciences of the USA* 113:8658–8663.
- Josephson, J. 2008. A numerical analysis of the evolutionary stability of learning rules. *Journal of Economic Dynamics and Control* 32:1569–1599.
- Katsnelson, E., U. Motro, M. W. Feldman, and A. Lotem. 2011. Evolution of learned strategy choice in a frequency-dependent game. *Proceedings of the Royal Society B* 279:1176–1184.
- Lakshminarayanan, V. R., and L. R. Santos. 2008. Capuchin monkeys are sensitive to others' welfare. *Current Biology* 18:R999–R1000.
- Lehmann, L., I. Alger, and J. Weibull. 2015. Does evolution lead to maximizing behavior? *Evolution* 69:1858–1873.
- Lehmann, L., and L. Keller. 2006. The evolution of cooperation and altruism—a general framework and a classification of models. *Journal of Evolutionary Biology* 19:1365–1376.
- Mark, J. T., B. B. Marion, and D. D. Hoffman. 2010. Natural selection and veridical perceptions. *Journal of Theoretical Biology* 266:504–515.
- McNamara, J. M., T. W. Fawcett, and A. I. Houston. 2013. An adaptive response to uncertainty generates positive and negative contrast effects. *Science* 340:1084–1086.
- McNamara, J. M., C. E. Gasson, and A. I. Houston. 1999. Incorporating rules for responding into evolutionary games. *Nature* 401:368–371.

- McNamara, J. M., and A. I. Houston. 2009. Integrating function and mechanism. *Trends in Ecology and Evolution* 24:670–675.
- Niv, Y. 2009. Reinforcement learning in the brain. *Journal of Mathematical Psychology* 53:139–154.
- Ockenfels, P. 1993. Cooperation in prisoners' dilemma. *European Journal of Political Economy* 9:567–579.
- Ok, E. A., and F. Vega-Redondo. 2001. On the evolution of individualistic preferences: an incomplete information scenario. *Journal of Economic Theory* 97:231–254.
- Pearce, J. M. 2008. *Animal learning and cognition: an introduction*. 3rd ed. Psychology, Hove, East Sussex.
- Rand, D. G., J. D. Greene, and M. A. Nowak. 2012. Spontaneous giving and calculated greed. *Nature* 489:427–430.
- Robson, A. J. 1990. Efficiency in evolutionary games: Darwin, Nash and the secret handshake. *Journal of Theoretical Biology* 144:379–396.
- Ruff, C. C., and E. Fehr. 2014. The neurobiology of rewards and values in social decision making. *Nature Reviews Neuroscience* 15: 549–562.
- Sandholm, W. H., E. Dokumaci, and F. Franchetti. 2012. Dynamo: diagrams for evolutionary game dynamics. <http://www.ssc.wisc.edu/~whs/dynamo>.
- Schino, G., and F. Aureli. 2010. The relative roles of kinship and reciprocity in explaining primate altruism. *Ecology Letters* 13:45–50.
- Schneeberger, K., M. Dietz, and M. Taborsky. 2012. Reciprocal cooperation between unrelated rats depends on cost to donor and benefit to recipient. *BMC Evolutionary Biology* 12:41.
- Schultz, W. 2015. Neuronal reward and decision signals: from theories to data. *Physiological Reviews* 95:853–951.
- Shettleworth, S. J. 2009. *Cognition, evolution, and behavior*. Oxford University Press, New York.
- Silk, J. B., S. F. Brosnan, J. Vonk, J. Henrich, D. J. Povinelli, A. S. Richardson, S. P. Lambeth, J. Mascaró, and S. J. Schapiro. 2005. Chimpanzees are indifferent to the welfare of unrelated group members. *Nature* 437:1357–1359.
- Stewart, A. J., and J. B. Plotkin. 2013. From extortion to generosity, evolution in the Iterated Prisoner's Dilemma. *Proceedings of the National Academy of Sciences of the USA* 110:15348–15353.
- Sutton, R. S., and A. G. Barto. 1998. *Reinforcement learning: an introduction*. MIT Press, Cambridge, MA.
- Taborsky, M., J. G. Frommen, and C. Riehl. 2016. Correlated pay-offs are key to cooperation. *Philosophical Transactions of the Royal Society B* 371:20150084.
- Taylor, P. D., and L. B. Jonker. 1978. Evolutionary stable strategies and game dynamics. *Mathematical Biosciences* 40:145–156.
- Tinghög, G., D. Andersson, C. Bonn, H. Böttiger, C. Josephson, G. Lundgren, D. Västfjäll, M. Kirchler, and M. Johannesson. 2013. Intuition and cooperation reconsidered. *Nature* 498:E1–E2.
- Trivers, R. L. 1971. The evolution of reciprocal altruism. *Quarterly Review of Biology* 46:35–57.
- Van Cleve, J. 2015. Social evolution and genetic interactions in the short and long term. *Theoretical Population Biology* 103:2–26.
- Van Cleve, J., and E. Akçay. 2014. Pathways to social evolution: reciprocity, relatedness, and synergy. *Evolution* 68:2245–2258.
- Voelkl, B., S. J. Portugal, M. Unsöld, J. R. Usherwood, A. M. Wilson, and J. Fritz. 2015. Matching times of leading and following suggest cooperation through direct reciprocity during V-formation flight in ibis. *Proceedings of the National Academy of Sciences of the USA* 112: 2115–2120.
- Wilkinson, G. S. 1988. Reciprocal altruism in bats and other mammals. *Ethology and Sociobiology* 9:85–100.
- Wilkinson, G. S., G. G. Carter, K. M. Bohn, and D. M. Adams. 2016. Non-kin cooperation in bats. *Philosophical Transactions of the Royal Society B* 371:20150095.

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