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Highlights

- Larger group sizes can both hinder and promote the evolution of cooperation.
- Increasing the group size decreases the proportion of cooperators at equilibrium.
- Increasing the group size increases the basin of attraction of more cooperative outcomes.

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Group size effects in social evolution

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Abstract

How the size of social groups affects the evolution of cooperative behaviors is a classic question in evolutionary biology. Here we investigate group size effects in the evolutionary dynamics of games in which individuals choose whether to cooperate or defect and payoffs do not depend directly on the size of the group. We find that increasing the group size decreases the proportion of cooperators at both stable and unstable rest points of the replicator dynamics. This implies that larger group sizes can have negative effects (by reducing the amount of cooperation at stable polymorphisms) and positive effects (by enlarging the basin of attraction of more cooperative outcomes) on the evolution of cooperation. These two effects can be simultaneously present in games whose evolutionary dynamics feature both stable and unstable rest points, such as public goods games with participation thresholds. Our theory recovers and generalizes previous results and is applicable to a broad variety of social interactions that have been studied in the literature.

Keywords: evolution of cooperation, evolutionary game theory, replicator dynamics, public goods games.

JEL classification: C73; H41

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1 Introduction

2 Cooperative behaviors increase the fitness of other individuals, possibly at the expense
3 of a personal fitness cost (Sachs et al., 2004). Biological examples include the produc-
4 tion of extracellular public goods in microbes (e.g., iron-scavenging molecules, West
5 and Buckling 2003, bacteriocins that eliminate competition, Bucci et al. 2011, and
6 factors that contribute to biofilm formation, Rainey and Rainey 2003), vigilance and
7 sentinel behavior in meerkats (Clutton-Brock et al., 1999), group hunting in social
8 carnivores (Packer and Ruttan, 1988), and the costly punishment of free-riders in hu-
9 mans (Raihani and Bshary, 2011). Identifying the different pathways that allow coop-
10 erative behavior to be favored by natural selection (Lehmann and Keller, 2006; Nowak,
11 2006; West et al., 2007; Van Cleve and Akçay, 2014) is important for understanding
12 the origin of social groups (Krause and Ruxton, 2002) and the major transitions in
13 evolution (Maynard Smith and Szathmáry, 1995; Bourke, 2011).

14 Group size is a crucial variable of social life. Therefore, how an increase or decrease
15 in group size affects individual incentives to cooperate is a recurrent question across
16 the behavioral sciences. In economics and political science, the “group-size paradox”
17 refers to cases where larger groups are less successful than smaller groups in pursuing
18 their common goals because individuals have a greater incentive to shirk when group
19 size is large (Olson, 1965; Esteban and Ray, 2001). In behavioral ecology, one of the
20 most replicated findings is the negative relationship between group size and level of
21 vigilance in social foragers due to increased predator detection and dilution of preda-
22 tor risk (Elgar, 1989; Roberts, 1996; Beauchamp, 2008). Increasing group size has also
23 been shown to reduce voluntary contributions to public goods (Isaac and Walker, 1988)
24 and reciprocity-based cooperation in multi-person interactions (Boyd and Richerson,
25 1988). More generally, however, whether or not larger groups are less conducive to
26 cooperation might depend on specific assumptions about group interactions. In par-
27 ticular, instances of positive group size effects have also been reported in the empirical
28 literature (Isaac et al., 1994; Yip et al., 2008; Powers and Lehmann, 2017) and are of
29 significant theoretical interest (Dugatkin, 1990; Shen et al., 2014; Powers and Lehmann,
30 2017; Cheikbossian and Fayat, 2018).

31 To study how the size of social groups affects the evolution of cooperation we fol-
32 low the standard approach of modelling social interactions as symmetric games with
33 two strategies (“cooperate” and “defect”) between several players, i.e., as symmetric
34 multiplayer matrix games (Broom et al., 1997; Gokhale and Traulsen, 2014). Payoffs
35 depend on the own strategy and on the number of co-players choosing to cooperate,
36 possibly in a nonlinear way. Strategies are genetically or culturally transmitted, and
37 populations are large enough that the replicator dynamic (Weibull, 1995; Hofbauer and
38 Sigmund, 1998) provides a reasonable model of evolution. Within this framework, the
39 stable rest points of the replicator dynamic correspond to evolutionary endpoints, while
40 the unstable rest points signpost the basins of attraction of such evolutionary attrac-
41 tors. Many social dilemmas for which cooperation can be maintained without repeated
42 interactions or genetic assortment have been theoretically studied using this or related
43 formalisms during the last decades (Taylor and Ward, 1982; Palfrey and Rosenthal,

44 1984; Diekmann, 1985; Boyd and Richerson, 1988; Motro and Eshel, 1988; Dugatkin,
 45 1990; Dixit and Olson, 2000; Goeree and Holt, 2005; Bach et al., 2006; Hauert et al.,
 46 2006; Archetti, 2009; Pacheco et al., 2009; Souza et al., 2009; Archetti and Scheuring,
 47 2011; Chen et al., 2013; Van Cleve and Lehmann, 2013; Sasaki and Uchida, 2014; Chen
 48 et al., 2015; Peña et al., 2015; Chen et al., 2017; De Jaegher, 2017; dos Santos and
 49 Peña, 2017; Kaznatcheev et al., 2017).

50 To obtain our results, we make use of the fact that the gain function determin-
 51 ing the direction of selection in the replicator dynamic is a polynomial in Bernstein
 52 form (Farouki, 2012). The coefficients of this polynomial are given by the gains from
 53 switching (Peña et al., 2014), i.e., the differences in payoff a focal player obtains by
 54 switching from defection to cooperation as a function of the number of other coop-
 55 erators in the group. Our analysis makes essential use of the structure of the gain
 56 sequence of the game, which collects such gains from switching. We illustrate our re-
 57 sults with examples and discuss how previous results in the literature (either proven
 58 using alternative arguments or hinted at by numerical analysis) can be recovered using
 59 our approach.

60 Under the conditions that payoffs do not depend directly on the size of the group
 61 and that the number of interior rest points of the replicator dynamics do not change
 62 as group size increases, we establish that the proportion of cooperators at both stable
 63 and unstable interior rest points decreases with group size. This finding, summarized
 64 in Proposition 1 in Section 3.1, is our main result. Proposition 1 implies that two kinds
 65 of group size effects are possible in the games we analyze. First, a negative group size
 66 effect, as the levels of cooperation at stable polymorphisms decrease with increasing
 67 group size. Second, a positive group size effect, as the size of the basin of attraction of
 68 the stable rest point with the largest level of cooperation increases as well. Proposition 2
 69 identifies general conditions under which the number of rest points is independent of
 70 group size.

71 Sections 3.2 and 3.3 explore the consequences of these general results for two im-
 72 portant particular cases subsuming many of the multiplayer matrix games appearing
 73 in the literature studying the evolution of cooperation (e.g., Dugatkin 1990; Weesie
 74 and Franzen 1998; Bach et al. 2006; Pacheco et al. 2009; Souza et al. 2009; Archetti
 75 and Scheuring 2011). Section 3.2 considers games with gain sequences having a single
 76 sign change. For such games the replicator dynamics have a unique interior rest point
 77 that is decreasing in group size (Proposition 3). If the sign change is from positive to
 78 negative, the interior rest point is stable and the group size effect is negative, as the
 79 proportion of cooperators at the interior rest point decreases. Conversely, if the sign
 80 change is from negative to positive, the interior rest point is unstable and the group size
 81 effect is positive, as the basin of attraction of full defection decreases while the basin
 82 of attraction of full cooperation increases. In Section 3.3, we focus on games charac-
 83 terized by “bistable coexistence” (Gokhale and Traulsen, 2014; Peña et al., 2015), i.e.,
 84 a phase portrait where the unstable interior rest point divides the basins of attraction
 85 of the stable interior rest point and full defection. Such a phase portrait is typical of
 86 many nonlinear social dilemmas, including those featuring participation thresholds or
 87 public goods games with sigmoid production functions (Dugatkin, 1990; Bach et al.,

88 2006; Pacheco et al., 2009; Souza et al., 2009; Archetti and Scheuring, 2011; Peña et al.,
 89 2014; Archetti, 2018). For these games there is both a negative group size effect (as
 90 the proportion of cooperators at the stable interior rest point decreases) and a positive
 91 group size effect (as the basin of attraction of full defection decreases). This result
 92 is stated in Proposition 4. Alternatively, an increase in group size can lead to a loss
 93 of both interior rest points. This makes the group size effect negative as an increase
 94 in group size results in full defection being the only attracting point of the replicator
 95 dynamics.

96 Several models in the literature consider a more complicated dependence of payoffs
 97 on group size than the one we consider in our main result. For instance, if the total
 98 benefit from cooperating has to be shared among group members (as in standard for-
 99 mulations of the linear public goods game, see, e.g., Boyd and Richerson 1988), then
 100 the gains from switching themselves depend on group size. This introduces an addi-
 101 tional effect, which might either reinforce or countervail the fundamental group size
 102 effect investigated in Section 3. We investigate this additional effect in Section 4 and
 103 state counterparts of Propositions 3 and 4 as Propositions 5 and 6. Section 5 discusses
 104 and concludes.

105 2 Model

106 2.1 Social interactions

107 Social interactions take place in groups of equal size n . Throughout the paper, n is
 108 treated as a parameter that satisfies $\underline{n} \leq n \leq \bar{n}$ for some given numbers $\underline{n} < \bar{n}$ and we
 109 use N to denote the set of all such group sizes. Individuals within each group participate
 110 in a symmetric n -player game, playing either strategy A (“cooperate”) or strategy B
 111 (“defect”). The payoff for an individual is determined by its own strategy and the
 112 number of other individuals in the group who cooperate but is otherwise independent
 113 of group size. Let a_k denote the payoff to an A -player (“cooperator”) and b_k denote the
 114 payoff to a B -player (“defector”) when $k = 0, 1, \dots, n-1$ co-players play A (and hence
 115 $n-1-k$ co-players play B). Irrespective of their own strategy, players prefer other
 116 group members to cooperate, i.e., $a_{k+1} \geq a_k$ and $b_{k+1} \geq b_k$ hold for all $k = 0, 1, \dots, n-2$
 117 (Uyenoyama and Feldman, 1980; Kerr et al., 2004). We begin our analysis by assuming
 118 that the payoffs a_k , and b_k do not depend explicitly on group size n . In Section 4 we
 119 relax this assumption.

120 The gain in payoff an individual makes from cooperating rather than defecting when
 121 k co-players cooperate is $d_k = a_k - b_k$. We refer to this as the k -th gain from switching
 122 (to cooperation). In all of the games we consider in the following, d_k will be negative
 123 for some k , indicating the presence of a social dilemma in which individuals increase
 124 their own payoff by defecting but thereby lower the payoffs of all other group members
 125 (Matessi and Karlin, 1984; Kerr et al., 2004).

126 While our results apply more generally, we will consider a variety of public goods
 127 games to motivate and illustrate our results. In these games, cooperators make a
 128 costly contribution to the provision of a public good, whereas defectors free ride on

129 the contribution of cooperators. Unless indicated otherwise, we will suppose that the
 130 cost $c > 0$ incurred by each contributor is independent of the the number of other
 131 contributors and that all group members obtain the same benefit u_j , which is increasing
 132 in the number of cooperators j . As the number of contributors includes the focal
 133 player, we have $j = k$ if the focal player defects, but $j = k + 1$ if the focal player
 134 cooperates. Therefore, in such a public goods game payoffs are given by $a_k = u_{k+1} - c$
 135 and $b_k = u_k$, and the k -th gain from switching is $d_k = \Delta u_k - c$, where $\Delta u_k = u_{k+1} -$
 136 $u_k \geq 0$. Perhaps the simplest example of such a public goods game is the volunteer's
 137 dilemma (Diekmann, 1985) in which at least one cooperator is required for a benefit
 138 $v > c$ to be enjoyed by all group members. This corresponds to the case $\theta = 1$ of a
 139 threshold public goods game, in which a minimum number θ of cooperators is required
 140 for a benefit $v > c$ to be enjoyed by all group members, so that $u_j = v$ if $j \geq \theta$ and
 141 $u_j = 0$ otherwise (Taylor and Ward, 1982; Palfrey and Rosenthal, 1984; Bach et al.,
 142 2006; Archetti, 2009).

143 2.2 Evolutionary dynamics

144 Evolution occurs in a large, well-mixed population with groups of identical size n ran-
 145 domly formed by binomial sampling. Hence, if there is a proportion x of A -players
 146 and a proportion $1 - x$ of B -players in the population, then the expected payoffs to
 147 A -players and B -players are respectively given by

$$\pi_n^A(x) = \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} a_k,$$

148 and

$$\pi_n^B(x) = \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} b_k.$$

149 We assume that the change in the proportion of A -players over evolutionary time
 150 is given by the continuous-time replicator dynamic (Weibull, 1995; Hofbauer and Sig-
 151 mund, 1998)

$$\dot{x} = x(1-x)g_n(x), \quad (1)$$

152 where

$$g_n(x) = \pi_n^A(x) - \pi_n^B(x) = \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} d_k, \quad (2)$$

153 i.e., the difference in expected payoffs between the two strategies, is the “gain function”
 154 (Bach et al., 2006), which can also be interpreted as the selection gradient on a contin-
 155 uously varying mixed strategy x (Peña et al., 2015). Since the factor $x(1-x)$ is always
 156 nonnegative, the sign of the gain function $g_n(x)$ indicates the sign of \dot{x} in Eq. (1) and
 157 hence the direction of selection, that is, whether or not the proportion of A -players will
 158 increase for a given population composition x and group size n .

159 The replicator dynamic has two trivial (or “pure”) rest points at $x = 0$ (where
 160 the whole population consists of defectors) and at $x = 1$ (where the whole population

161 consists of cooperators). Interior (or “mixed”) rest points are given by the values
 162 $x^* \in (0, 1)$ satisfying

$$g_n(x^*) = 0.$$

163 To simplify the exposition, we impose the regularity condition that $dg_n(x^*)/dx \neq 0$
 164 holds at all interior rest points. An interior rest point is then stable (i.e., evolution-
 165 arily attracting) if and only if $dg_n(x^*)/dx < 0$ holds, and unstable (i.e., evolutionarily
 166 repelling) otherwise. We further suppose that for $n \in N$ the number of sign changes
 167 s of the gain sequences $(d_0, d_1, \dots, d_{n-1})$ is independent of group size n (i.e., the gain
 168 sequence $(d_0, d_1, \dots, d_{\bar{n}-1})$ has no sign changes between $\underline{n} - 1$ and $\bar{n} - 1$), indicating
 169 that the fundamental structure of the social dilemma under consideration is the same
 170 for all group sizes in the range under consideration. Moreover, we suppose that $s \geq 1$
 171 holds as otherwise either full defection ($x = 0$) or full cooperation ($x = 1$) is the unique
 172 stable rest point of the replicator dynamics for all group sizes $n \in N$.

173 3 Results

174 3.1 General results

175 Our first result shows that if the number of interior rest points is independent of group
 176 size, then the proportion of cooperators at all interior rest points decreases when group
 177 size increases.

178 **Proposition 1.** *Suppose that the replicator dynamics (1)–(2) have the same number*
 179 *of interior rest points $\ell \geq 1$ for all group sizes $n \in N$, and denote these rest points*
 180 *by $0 < x_{n,1}^* < \dots < x_{n,\ell}^* < 1$ for group size n . Then $x_{n+1,r}^* < x_{n,r}^*$ holds for all*
 181 *$n = \underline{n}, \dots, \bar{n} - 1$ and $r = 1, \dots, \ell$.*

182 The full proof of Proposition 1 is in Appendix A.1. The key step towards obtain-
 183 ing this result is the following identity, which links the gain functions (and thus the
 184 replicator dynamics) for adjacent group sizes:

$$g_n(x) = g_{n+1}(x) - \frac{x}{n} \frac{dg_{n+1}}{dx}(x). \quad (3)$$

185 Eq. (3) is a simple consequence of properties of the gain functions $g_n(x)$, previously
 186 observed by Motro (1991), which stem from the fact that the gain functions are poly-
 187 nomials in Bernstein form (Peña et al., 2014) with coefficients (given by the gains from
 188 switching d_k) that do not depend on group size.

189 To see how Eq. (3) yields Proposition 1, observe that this equation implies that at
 190 the interior rest points of the dynamic with group size $n + 1$ (where the gain function
 191 $g_{n+1}(x)$ vanishes), the gain function $g_n(x)$ will have the opposite sign of the derivative
 192 $dg_{n+1}(x)/dx$. This ensures that between any two interior rest points of the replicator
 193 dynamic for group size $n + 1$ the replicator dynamic for group size n has exactly one
 194 rest point. The result then follows upon establishing that the remaining interior rest
 195 point for the replicator dynamic with group size n must have a higher proportion of
 196 cooperators than the largest interior rest point $x_{n+1,\ell}^*$ for group size $n + 1$.

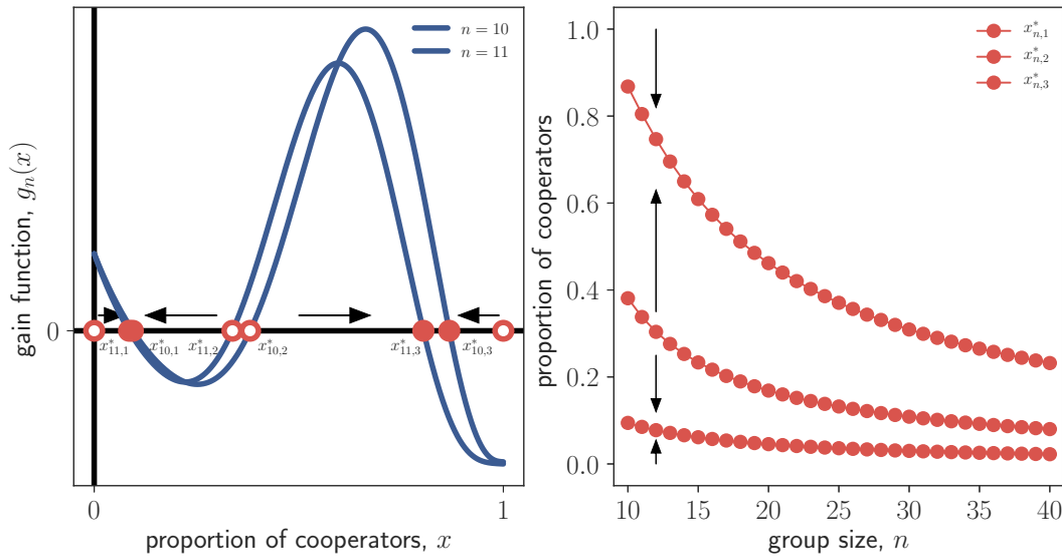


Figure 1: Group size effects in the threshold public good game with an additional reward $\delta > 0$ shared among cooperators considered by Chen et al. (2013). Payoffs are given by $a_k = u_{k+1} + \delta/(k+1) - c$ and $b_k = u_k$, where $u_j = v$ if $j \geq \theta$ and $u_j = 0$ otherwise. In all panels, $c = 1$, $v = 5$, $\theta = 7$, $\delta = 1.5$. *Left panel:* Gain functions (blue lines) with corresponding rest points (red symbols), and direction of selection (black arrows) for two group sizes: $n = 10$, and $n = 11$. Full circles represent stable rest points and empty circles represent unstable rest points. *Right panel:* Proportion of cooperators at the interior rest points as function of group size for $10 \leq n \leq 40$. The direction of selection (black arrows) is also shown. As group size increases, the proportion of cooperators at interior rest points decreases.

197 The decrease in the proportion of cooperators at all interior rest points as group
 198 size increases asserted in Proposition 1 leads to contrasting effects of group size on the
 199 evolution of cooperation. First, there is an obvious negative group size effect, as the
 200 proportion of cooperators at stable polymorphisms decreases with group size. Second,
 201 the proportion of cooperators at unstable rest points decreases as well. As the rest
 202 points of the replicator dynamics alternate between being stable and unstable, this
 203 implies an increase in the size of the basin of attraction of the stable rest point with
 204 the largest proportion of cooperators. Hence, there is also a positive group size effect.
 205 These two effects are illustrated in Fig. 1 for the relatively complex case of a game with
 206 three interior rest points: $x_{n,1}^*$ (stable), $x_{n,2}^*$ (unstable), and $x_{n,3}^*$ (stable). In line with
 207 Proposition 1, larger group sizes lead to smaller proportions of cooperators at the two
 208 stable interior rest points $x_{n,1}^*$ and $x_{n,3}^*$ but also, via a decrease in the value of $x_{n,2}^*$,
 209 to a larger basin of attraction for $x_{n,3}^*$ and a smaller basin of attraction for $x_{n,1}^*$. As
 210 $x_{n,3}^*$ sustains a higher level of cooperation than $x_{n,1}^*$, this latter effect can be said to
 211 promote the evolution of cooperation.

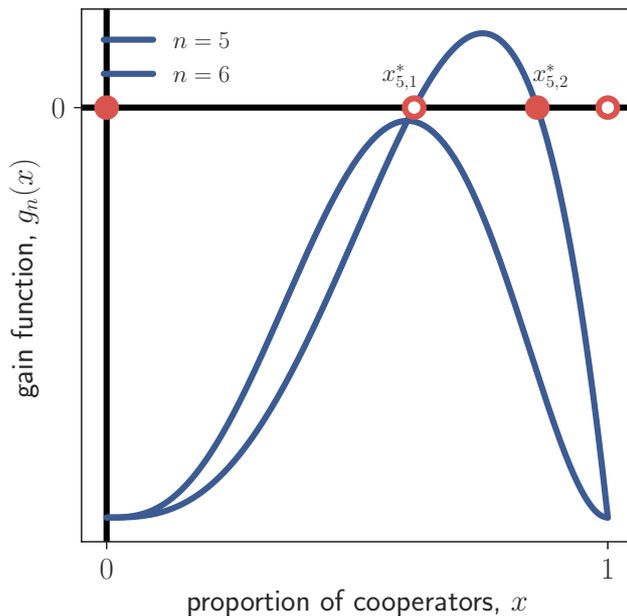


Figure 2: An increase in group size can lead to a reduction in the number of rest points. Here we illustrate this effect for a threshold public goods game with $c = 1$, $v = 2.8$, $\theta = 4$, which has two interior rest points for group size $n = 5$ but no interior rest points for group size $n = 6$.

212 Proposition 1 is predicated on the assumption that the number of interior rest points
 213 for the different group sizes under consideration is the same. This does not have to be
 214 the case. In particular, it is possible that an increase in group size leads to a decrease
 215 in the number of rest points. Fig. 2 illustrates this possibility for the case of a threshold
 216 public goods game. On the other hand, the arguments establishing Proposition 1 show
 217 that an increase in group size can never lead to an increase in the number of rest points.
 218 Further, it is known that the number of interior rest points of the replicator dynamics
 219 cannot exceed the number of sign changes s in the gain sequences (Peña et al., 2014,
 220 Property 2). Therefore, if the number of interior rest points of the replicator dynamic
 221 for the maximal group size \bar{n} is equal to s , then the number of interior rest points of
 222 the replicator dynamics is independent of group size. The proof of the following result
 223 in Appendix A.2 demonstrates that, in addition, if an increase in group size causes
 224 a reduction in the number of interior rest points, then the number of rest points is
 225 reduced by an even amount.

226 **Proposition 2.** *Suppose that the number of interior rest points of the replicator dy-*
 227 *namic (1)–(2) for group size \bar{n} is equal to the number of sign changes s of the gain*
 228 *sequences. Then for all group sizes $n \in N$ the number of interior rest points of the*
 229 *replicator dynamics is equal to s . More generally, if $\bar{n} \geq n > m \geq \underline{n}$, then the number*
 230 *of interior rest points of the replicator dynamic with group size n is either equal to the*

231 *number of interior rest points of the replicator dynamic with group size m or lower by*
 232 *an even amount.*

233 3.2 Games with a unique interior rest point

234 Suppose that for all group sizes $n \in N$ the replicator dynamics have a unique interior
 235 rest point that, for simplicity, we denote by x_n^* . It is then immediate from Proposition
 236 1 that the proportion of cooperators at this rest point is decreasing in group size.
 237 Combining this observation with the sufficient condition for the existence of a unique
 238 interior rest point from Result 3 in Peña et al. (2014) immediately yields:

239 **Proposition 3.** *Suppose that for all $n \in N$ the gain sequences have a single sign change*
 240 *($s = 1$). Then the replicator dynamics (1)–(2) have a unique interior rest point for all*
 241 *$n \in N$, and the proportion of cooperators x_n^* at this interior rest point is decreasing in*
 242 *group size n .*

243 Proposition 3 encompasses two cases. First, the gains from switching can be positive
 244 for a small number of cooperators (up to some threshold $\hat{k} < \underline{n}$) and negative for a
 245 large number of cooperators (beyond the threshold \hat{k}). In this case there exists a
 246 unique interior rest point x_n^* that is also the unique stable rest point of the replicator
 247 dynamics (Peña et al., 2014, Result 3.2). For this case, Proposition 3 indicates that the
 248 group size effect is negative in the sense that an increase in group size causes a decrease
 249 in the proportion of cooperators at equilibrium. This finding generalizes a result due
 250 to Motro (1991), who showed that there is a unique stable interior rest point and a
 251 negative group size effect for public goods games with concave benefits and intermediate
 252 costs (for which Δu_k , and therefore d_k , is decreasing in k , and $\Delta u_0 > c > \Delta u_{\underline{n}}$ holds).
 253 It also generalizes the well-known result that the proportion of cooperators at the
 254 unique stable rest point of the volunteer's dilemma is decreasing in group size (cf.,
 255 e.g., Archetti 2009) and corresponding results for the volunteer's dilemma with cost
 256 sharing (Dugatkin, 1990; Weesie and Franzen, 1998). This last example, which differs
 257 from the other two in that the gains from switching are not monotonically decreasing
 258 in k , is illustrated in Fig. 3.

259 The second case encompassed by Proposition 3 is the one in which the gains from
 260 switching are negative for a small number of cooperators (up to some threshold $\hat{k} < \underline{n}$)
 261 and positive for a large number of cooperators (beyond the threshold \hat{k}). In this case
 262 the two trivial rest points $x = 0$ and $x = 1$ are stable and the unique interior rest
 263 point x_n^* , which separates the basins of attraction of the two stable rest points, is
 264 unstable (Peña et al., 2014, Result 3.2). For this case, Proposition 3 indicates that
 265 the group size effect is positive in the sense that with an increase in group size the
 266 basin of attraction of full defection ($x = 0$) shrinks while the basin of attraction of
 267 full cooperation ($x = 1$) increases. For public goods games with convex benefits and
 268 intermediate cost (for which Δu_k , and therefore d_k , is increasing with $\Delta u_0 < c < \Delta u_{\underline{n}}$)
 269 this effect has been previously noted in Motro (1991).

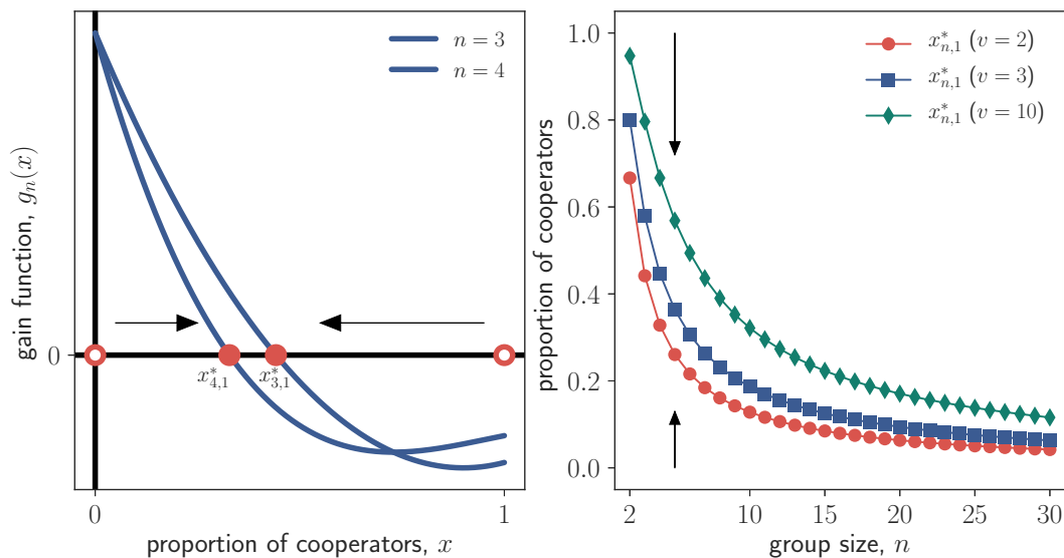


Figure 3: Group size effects in the volunteer's dilemma with cost sharing considered by Weesie and Franzen (1998). Payoffs are given by $a_k = v - c/(k + 1)$, $b_0 = 0$, and $b_k = v$ for $k \geq 1$. In all panels, $c = 1$. *Left panel:* Gain functions (blue lines) with corresponding rest points (red symbols), and direction of selection (black arrows) for $v = 2$, and two group sizes: $n = 3$, and $n = 4$. Full circles represent stable rest points and empty circles represent unstable rest points. *Right panel:* Proportion of cooperators at the interior rest point as function of group size for different parameter values. The direction of selection (black arrows) is also shown. As group size increases, the proportion of cooperators at the unique stable interior rest point decreases, i.e., the group size effect is negative.

270 3.3 Games with two interior rest points

271 Many social dilemmas are such that defection is individually advantageous if the number
 272 of cooperating co-players is either sufficiently small or sufficiently high, while cooper-
 273 ation is individually advantageous in between, i.e., the gains from switching satisfy
 274 $d_0 < 0$ and the gain sequences $(d_0, d_1, \dots, d_{n-1})$ have two sign changes for all group
 275 sizes $n \in N$. This scenario arises in the threshold public goods game when the mini-
 276 mum number θ of cooperators required for the benefit $v > c$ to be enjoyed by all group
 277 members satisfies $2 \leq \theta < \underline{n}$. More generally, public goods games in which the benefits
 278 from the provision of the public good are sigmoid in the number of contributors and
 279 the costs of provision are intermediate (Archetti and Scheuring, 2011; Archetti, 2018)
 280 have this structure. Peña et al. (2014) provide further examples and discussion.

281 Assuming that the gains from switching have the structure described above ensures
 282 that the rest point at $x = 0$ is stable and the rest point at $x = 1$ is unstable for
 283 all $n \in N$ (Peña et al., 2014, Result 1). Further, there are at most two interior rest
 284 points satisfying $0 < x_{n,1}^* < x_{n,2}^* < 1$, with the smaller of these rest points ($x_{n,1}^*$) being
 285 unstable and the larger interior rest point ($x_{n,2}^*$) being stable. The existence of these
 286 rest points is guaranteed if $\bar{g}_n = \max_{0 \leq x \leq 1} g_n(x) > 0$ holds (Peña et al., 2014, Result
 287 4.1). Combining these observations with the arguments yielding the results in Section
 288 3.1, Appendix A.3 proves:

289 **Proposition 4.** *Suppose that for all $n \in N$ the gain sequences have two sign changes*
 290 *($s = 2$), their initial signs are negative, and that $\bar{g}_n > 0$ holds. Then, the replicator*
 291 *dynamics (1)–(2) have two interior rest points for all group sizes $n \in N$. Further,*
 292 *at both the unstable rest point $x_{n,1}^*$ and the stable rest point $x_{n,2}^*$ the proportion of*
 293 *cooperators is decreasing in group size and we have*

$$x_{n+1,1}^* < x_{n,1}^* < x_{n+1,2}^* < x_{n,2}^* \quad (4)$$

294 for all n satisfying $\underline{n} \leq n < \bar{n}$.

295 Proposition 4 indicates that there are two different effects of group size on cooper-
 296 ation in games with two interior rest points. First, there is a negative group size effect,
 297 as the proportion of cooperators at the stable interior rest point decreases as group
 298 size increases, i.e., $x_{n+1,2}^* < x_{n,2}^*$ holds. Second, there is a positive group size effect,
 299 as the proportion of cooperators at the unstable interior rest point also decreases as
 300 group size increases, i.e., $x_{n+1,1}^* < x_{n,1}^*$ holds, implying that the basin of attraction of
 301 full defection ($x = 0$) shrinks while the basin of attraction of the stable rightmost rest
 302 point increases. These effects are in line with what happens in games with a unique
 303 interior rest point that we have discussed in Section 3.2. The additional twist is that
 304 rather than having the group size effect being negative or positive depending on the
 305 structure of the game, both the negative and the positive group size effects co-occur in
 306 the same game.

307 Fig. 4 illustrates Proposition 4 for the case of a threshold public goods game. As
 308 noted above, the result is applicable more generally. For instance, the observations
 309 (obtained via numerical calculations) that both interior rest points decrease with group

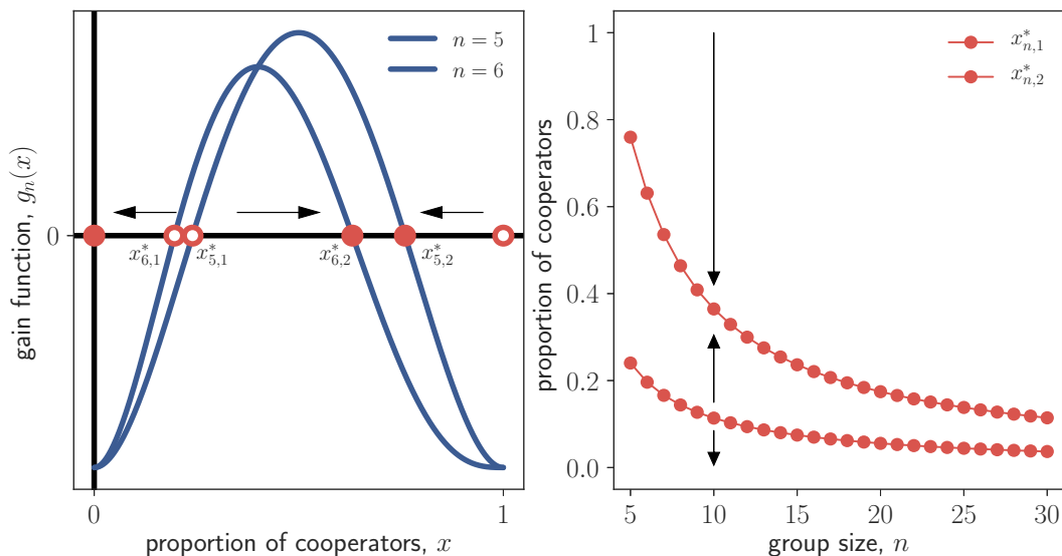


Figure 4: Group size effects in a threshold public good game. Payoffs are given by $a_k = u_{k+1} - c$ and $b_k = u_k$, where $u_j = v$ if $j \geq \theta$ and $u_j = 0$ otherwise. In all panels, $c = 1$, $v = 5$, and $\theta = 3$. *Left panel:* Gain functions (blue lines) with corresponding rest points (red symbols), and direction of selection (black arrows) for two group sizes: $n = 5$, and $n = 6$. Full circles represent stable rest points and empty circles represent unstable rest points. *Right panel:* Proportion of cooperators at the interior rest points as function of group size for $5 \leq n \leq 30$. The direction of selection (black arrows) is also shown. As group size increases, the proportion of cooperators at both interior rest points decreases. This leads to both a negative group size effect (the proportion of cooperators at the interior stable rest point decreases) and a positive group size effect (the basin of attraction of the interior stable rest point increases).

size for the n -person tit-for-tat model of Dugatkin (1990) (his “Model II”) and the n -person snowdrift game discussed by Souza et al. (2009), are implied by Proposition 4.

The role of the condition $\bar{g}_{\bar{n}} > 0$ in the statement of Proposition 4 is to ensure that the replicator dynamic has two interior rest points for group size \bar{n} and, therefore, has these two rest points for all group sizes (Proposition 2). If the reverse inequality $\bar{g}_{\bar{n}} < 0$ holds, then for large groups there are no interior rest points, whereas (provided that the inequality $\bar{g}_{\bar{n}} > 0$ holds) for small group sizes there exists two interior rest points. In such a situation there is (as illustrated in Fig. 2) a critical group size such that for smaller group sizes the rest point $x = 0$ is the only stable rest point, whereas for larger group sizes there is a stable polymorphism at which some proportion of the population cooperates. Hence, this describes a case in which the group size effect is unambiguously negative.

4 Extension: games with gain sequences depending on group size

So far our analysis has assumed that the payoffs a_k and b_k , and therefore the gains from switching d_k , depend only on the number of other cooperators a focal player interacts with and not directly on the size of the group. This assumption is not always warranted. For instance, Hauert et al. (2006) and Pacheco et al. (2009) consider variants of a public goods game in which the benefits u_k from cooperation are shared among all group members rather than accruing to each individual. The payoffs to cooperators and defectors are then $a_k^n = u_{k+1}/n - c$ and $b_k^n = u_k/n$. The resulting gains from switching

$$d_k^n = \frac{\Delta u_k}{n} - c, \quad (5)$$

depend not only on k but also on group size n .

If the gains from switching are, as in Eq. (5), decreasing in group size, then the proportion of cooperators at an unstable interior rest point may increase with group size. In particular, as illustrated for the case of a threshold public goods game in Fig. 6 below, Propositions 3 and 4 are no longer applicable to describe the group size effect on unstable interior rest points. Further, there is no hope to obtain a counterpart to Proposition 1. In the following we therefore focus on stable interior rest points and show that for these the conclusions from Propositions 3 and 4 remain intact.

Consider, first, the case in which the gain sequences $(d_0^n, d_1^n, \dots, d_{n-1}^n)$ have a single sign change from positive to negative for all group sizes $n \in N$. This ensures that the replicator dynamics, given by Eq. (1) with

$$g_n(x) = \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} d_k^n, \quad (6)$$

have a unique interior rest point x_n^* for all group sizes $n \in N$ and that this rest point is

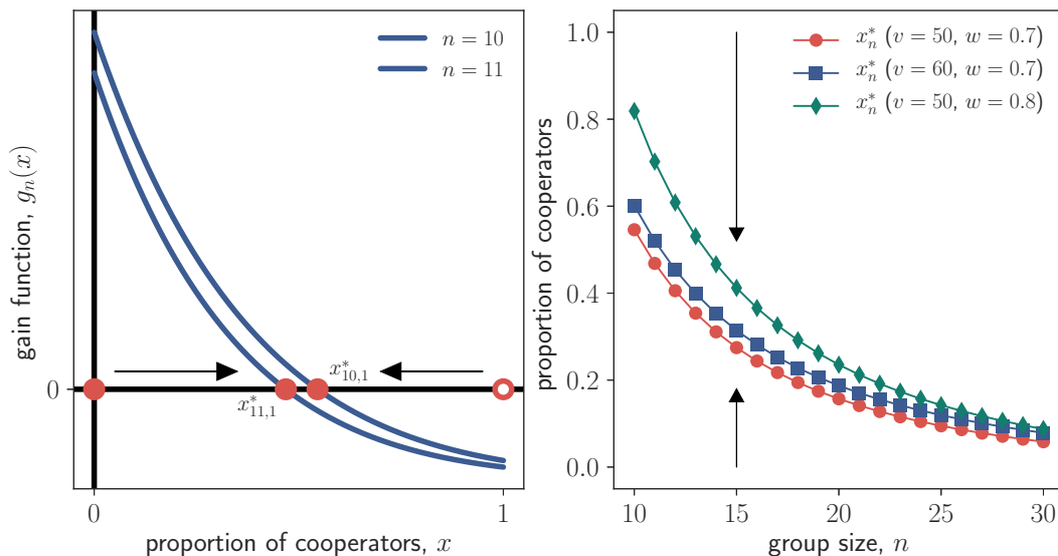


Figure 5: Group size effects in the model with discounted benefits from Hauert et al. (2006). Payoffs are given by $a_k = u_{k+1}/n - c$ and $b_k = u_k/n$, where $u_k = v(1 - w^k)/(1 - w)$ with $0 < w < 1$. For intermediate values of c ($w^{n-1}/n < c/v < 1/n$) the gains from switching $d_k^n = vw^k/n - c$ satisfy the assumptions in the statement of Proposition 5. In all panels, $c = 1$. *Left panel:* Gain functions (blue lines) with corresponding rest points (red symbols), and direction of selection (black arrows) for $v = 50$, $w = 0.7$, and two group sizes: $n = 10$, and $n = 11$. Full circles represent stable rest points and empty circles represent unstable rest points. *Right panel:* Proportion of cooperators at the interior rest point as function of group size for different parameter combinations. The direction of selection (black arrows) is also shown.

345 stable. Appendix A.4 shows the following result, and Fig. 5 illustrates it for the model
 346 with discounted benefits proposed by Hauert et al. (2006).

347 **Proposition 5.** *Suppose that the gain sequences $(d_0^n, d_1^n, \dots, d_{n-1}^n)$ have a single sign
 348 change from positive to negative for all $n \in N$. Then the replicator dynamics defined by
 349 Eq. (1) and Eq. (6) have a unique stable interior rest point x_n^* for all $n \in N$. Further,
 350 if $d_k^{n+1} \leq d_k^n$ holds for all $k = 0, 1, \dots, n-1$ and all n satisfying $\underline{n} \leq n < \bar{n}$, then the
 351 proportion of cooperators x_n^* at this interior rest point is decreasing in group size n .*

352 The intuition for Proposition 5 is that a decrease in the gains from switching de-
 353 creases the gain function and that such a decrease in the gain function reduces the
 354 proportion of cooperators at the stable interior rest point. Therefore, the negative de-
 355 pendence of the gains from switching on group size considered here reinforces the group
 356 size effect observed in Proposition 3 by further reducing the proportion of cooperators
 357 at the stable interior rest point. The same intuition applies to the following counterpart
 358 to Proposition 4 that we prove in Appendix A.5:

359 **Proposition 6.** *Suppose that the gain sequences $(d_0^n, d_1^n, \dots, d_{n-1}^n)$ have two sign changes,
 360 their initial signs are negative, and that $\bar{g}_n > 0$ holds for all $n \in N$. Then the replicator
 361 dynamics defined by Eq. (1) and Eq. (6) have two interior rest points $x_{n,1}^* < x_{n,2}^*$ for
 362 all group sizes $n \in N$ with the first of these unstable and the second stable. Further,
 363 if $d_k^{n+1} \leq d_k^n$ holds for all $k = 0, 1, \dots, n-1$ and all n satisfying $\underline{n} \leq n < \bar{n}$, then the
 364 proportion of cooperators at the stable rest point $x_{n,2}^*$ is decreasing in group size n .*

365 Fig. 6 illustrates the conclusions from Proposition 6 for a game having the same
 366 structure as a threshold public goods game, except that the benefits u_k are shared
 367 among all group members as in Eq. 5. Fig. 6 also illustrates that sharing the benefits
 368 among more group members decreases the gain function and thereby increases the
 369 proportion of cooperators at the unstable interior rest point compared to the benchmark
 370 case considered in Proposition 4. Depending on parameter values, this effect may or
 371 may not be large enough to overturn the conclusion from Proposition 4.

372 5 Discussion

373 We have investigated how group size affects the evolutionary dynamics of multiplayer
 374 cooperation. More specifically, we have shown that an increase in group size can have
 375 a negative effect (a decrease in the proportion of cooperators at equilibrium) and a
 376 positive effect (an increase in the basin of attraction of the stable rest point sustaining
 377 the largest proportion of cooperators) on social evolution. Depending on the payoff
 378 structure of the social interactions one effect can be present and the other absent (as in
 379 games featuring a single interior rest point), or both effects can be present at the same
 380 time (as in games featuring two interior rest points). For threshold public goods games
 381 and other games characterized by bistable coexistence both the invasion barrier needed
 382 for cooperators to invade a population of defectors and the proportion of cooperators
 383 expected at the stable interior rest point decrease as group size increases. We have

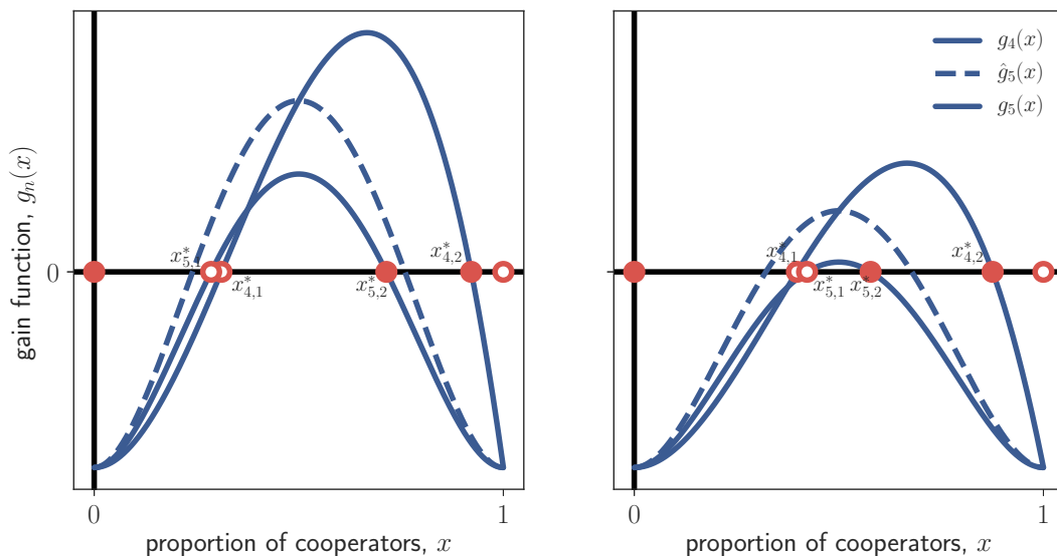


Figure 6: Illustration of Proposition 6 for the case of a threshold game with shared benefits. Payoffs are given by $a_k^n = u_{k+1}/n - c$ and $b_k^n = u_k/n$, where $u_j = v$ if $j \geq 3$ and $u_j = 0$ otherwise. Both panels show the gain functions $g_n(x)$ for group sizes $n = 4$ and $n = 5$ (solid lines) and the gain function $\hat{g}_5(x)$ for the larger group size (dashed lines) corresponding to the benchmark of a threshold public goods game in which payoffs for group size 5 are the same as for group size 4. In both panels the proportion of cooperators at the stable interior rest point decreases as group size increases. *Left panel:* the proportion of cooperators at the unstable interior rest point decreases as group size changes from $n = 4$ to $n = 5$ ($v = 20$, $c = 1$). *Right panel:* the proportion of cooperators at the unstable interior rest point increases as group size changes from $n = 4$ to $n = 5$ ($v = 14$, $c = 1$).

384 also shown that if payoffs depend explicitly on group size and such dependence is
385 negative, the negative group size effect is reinforced, while the positive group size effect
386 is attenuated or, depending on the particular payoff structure of the game, reversed.

387 The negative group size effect we identify is in line with the common expecta-
388 tion that the selection pressure on certain types of cooperation decreases as group size
389 rises. Such a negative group size effect requires that the gain sequence is sometimes
390 decreasing, meaning that individual incentives to cooperate are (at least for some social
391 contexts) decreasing in the number of cooperators in the group. When this is the case,
392 the decisions to cooperate are strategic substitutes (Bulow et al., 1985); equivalently,
393 cooperation is discounted or subject to diminishing returns. Anti-predator vigilance
394 often follows this payoff structure, as the presence of other vigilant individuals usually
395 disincentivizes individual investment in vigilance, i.e., there is a “many eyes” effect (Pul-
396 liam, 1973; McNamara and Houston, 1992); in extreme cases one vigilant individual
397 is enough for the group to be protected (Bednekoff, 1997; Clutton-Brock et al., 1999).
398 In agreement with our results, empirical and theoretical studies indicate that vigilant
399 behavior often decreases with group size (Elgar, 1989; McNamara and Houston, 1992;
400 Beauchamp, 2008).

401 Contrastingly, the positive group size effect we identify has been less emphasized in
402 evolutionary game theory (but see Sumpter and Brännström 2008 and Cornforth et al.
403 2012, who demonstrate this effect in models with continuous strategies). In an early
404 paper, Dugatkin (1990) noted that, in his model of n -person reciprocity, the threshold
405 frequency of cooperators needed to invade a population of defectors decreased as group
406 size increased. Our analysis reveals that such a positive group size effect is not specific
407 to the payoff structure assumed in Dugatkin (1990), but that it holds more generally
408 for any matrix game featuring unstable interior rest points. As a necessary condition
409 for the existence of unstable interior rest points is that the gain sequence is sometimes
410 increasing, the group size effect can be positive only when the individual incentives to
411 cooperate are (for at least some social contexts) increasing in the number of cooperators
412 in the group. In this case, the decisions to cooperate are strategic complements (Bulow
413 et al., 1985); equivalently, cooperation is synergistic or subject to increasing returns. A
414 common form of synergistic cooperation occurs when a critical number of cooperators
415 is required for cooperation to be individually worthwhile. Examples of such threshold
416 effects have been documented in empirical studies, and hypothesized to be a causal
417 factor behind inverse density dependence or Allee effects (Courchamp et al., 1999). For
418 instance, a large critical number of bark beetles is needed to overcome the defenses of
419 the tree they attack (Franceschi et al., 2005), and cooperative hunting often requires a
420 critical number of hunters to be energetically efficient (Creel and Creel, 1995; Alvard and
421 Nolin, 2002; MacNulty et al., 2014). Also, in group-hunting sailfish, a larger number of
422 hunters improves the hunting success of the group by allowing individuals to alternate
423 their attacks (Herbert-Read et al., 2016), and by keeping group-level unpredictability
424 high in the face of individual lateralization (Kurvers et al., 2017). In all of these
425 cases of synergistic cooperation, our theory suggests that larger groups can be more
426 favorable to cooperation and less favorable to free riding. Indeed, this general prediction
427 is in agreement with both general models of synergistic cooperation with continuous

428 cooperative investments (Cornforth et al., 2012), and a recent mechanistic model of
 429 free riding in group-hunting sailfish (Herbert-Read et al., 2016).

430 We used a variety of public goods games to illustrate our results. In such games,
 431 both cooperators and defectors gain equal access to the collective good produced by
 432 cooperators, i.e., the collective good is public. Notwithstanding the importance of these
 433 models, there are other social dilemmas for which public goods games are not a natural
 434 description of the relevant strategic trade-offs. For instance, social interactions can take
 435 the form of a collective action problem where the produced good can be accessed only
 436 by cooperators or only by defectors, i.e., the collective good is in some sense excludable.
 437 Group size effects in such “club” and “charity” goods games (Peña et al., 2015) are
 438 readily amenable to analysis by applying our results.

439 We conclude by noting that our analysis assumed populations were well-mixed and
 440 hence without genetic structure. This assumption is not always justified, as many
 441 social interactions take place in spatially structured populations characterized by non-
 442 negligible amounts of genetic structure (Rousset, 2004; Lehmann and Rousset, 2010;
 443 Van Cleve, 2015). A simple way of modeling social evolution in these populations
 444 is to focus on a continuously varying mixed strategy and to identify the convergence
 445 stable strategies of the resulting adaptive dynamics (e.g., Rousset 2004; Van Cleve and
 446 Lehmann 2013; Peña et al. 2015). In this case, the counterpart to the gain function we
 447 have analyzed in this paper is also a polynomial in Bernstein form, now with coefficients
 448 given by “inclusive gains from switching” depending on the payoffs of the game, the
 449 group size, and demographic parameters of the particular spatial model determining
 450 the degree of genetic relatedness and the amount of local competition (Peña et al.,
 451 2015). In this light, the analysis conducted here is also relevant to investigate group
 452 size effects in genetically structured populations, provided that the likely dependence
 453 of the inclusive gains from switching on group size is taken into account. Investigating
 454 the effects of group size on the evolution of cooperative behaviors under nontrivial
 455 population structure with the tools developed here would complement recent efforts in
 456 this area (Shen et al., 2014; Powers and Lehmann, 2017; Van Cleve, 2017).

457 Acknowledgements

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459 Appendix

460 A.1 Proof of Proposition 1

461 We first obtain Eq. (3). To do so, we make use of two identities established in the
 462 appendix of Motro (1991). Using our notation for the gain function and the gains from
 463 switching, these identities are

$$\frac{dg_n}{dx}(x) = (n-1) \sum_{k=0}^{n-2} \binom{n-2}{k} x^k (1-x)^{n-2-k} (d_{k+1} - d_k), \quad (7)$$

464 and

$$g_{n+1}(x) - g_n(x) = x \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} (d_{k+1} - d_k). \quad (8)$$

465 Applying Eq. (7) (which is nothing but the derivative property of polynomials in
466 Bernstein form; see, e.g., Peña et al. 2014) to group size $n+1$ and dividing both sides
467 of the resulting equation by n yields

$$\frac{1}{n} \frac{dg_{n+1}}{dx}(x) = \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} (d_{k+1} - d_k). \quad (9)$$

468 Substituting Eq. (9) into Eq. (8) we obtain

$$g_{n+1}(x) - g_n(x) = \frac{x}{n} \frac{dg_{n+1}}{dx}(x),$$

469 from which Eq. (3) is immediate.

470 Consider any n satisfying $\underline{n} \leq n < \bar{n}$. The following establishes that the replica-
471 tor dynamic for group size n must have a rest point in the interval $(x_{n+1,\ell}^*, 1)$: Be-
472 cause $g_{n+1}(x)$ has no root in $(x_{n+1,\ell}^*, 1)$, $g_{n+1}(x)$ has the same sign as the derivative
473 $dg_{n+1}(x_{n+1,\ell}^*)/dx$ for all $x \in (x_{n+1,\ell}^*, 1)$. As the gain sequences (d_0, \dots, d_{n-1}) and
474 (d_0, \dots, d_n) have the same initial sign (given by the sign of the first non-zero gain from
475 switching d_k) and the same number of sign changes s , they also have the same final
476 sign. Hence, the final sign of the gain sequence (d_0, \dots, d_n) is the same as the sign of
477 $dg_{n+1}(x_{n+1,\ell}^*)/dx$, too (Peña et al., 2014, Property 1). Therefore, for sufficiently large
478 $\hat{x} \in (x_{n+1,\ell}^*, 1)$ the sign of $g_n(\hat{x})$ coincides with the sign of $dg_{n+1}(x_{n+1,\ell}^*)/dx$. From
479 Eq. (3) and $g_{n+1}(x_{n+1,\ell}^*) = 0$ we then have that $g_n(x_{n+1,\ell}^*)$ and $g_n(\hat{x})$ have opposite
480 signs, so that $g_n(x)$ has a root in the interval $(x_{n+1,\ell}^*, \hat{x})$. Consequently, the replicator
481 dynamic for group size n has a rest point in the interval $(x_{n+1,\ell}^*, \hat{x})$. For the case $\ell = 1$
482 this finishes the proof of the proposition.

483 Suppose $\ell \geq 2$ and let n again satisfy $\underline{n} \leq n < \bar{n}$. Consider (with $r = 1, \dots, \ell - 1$)
484 any adjacent interior rest points $x_{n+1,r}^* < x_{n+1,r+1}^*$ of the replicator dynamic for group
485 size $n+1$. As stable and unstable rest points alternate, the derivatives $dg_{n+1}(x_{n+1,r}^*)/dx$
486 and $dg_{n+1}(x_{n+1,r+1}^*)/dx$ have opposite signs. As $g_{n+1}(x_{n+1,r}^*) = g_{n+1}(x_{n+1,r+1}^*) = 0$
487 holds, it follows from Eq. (3) that $g_n(x_{n+1,r}^*)$ and $g_n(x_{n+1,r+1}^*)$ have opposite signs, too.
488 Therefore, $g_n(x)$ has at least one root in the interval $(x_{n+1,r}^*, x_{n+1,r+1}^*)$, with each such
489 root corresponding to an interior rest point of the replicator dynamic for group size n .
490 As there are $\ell - 1$ intervals of the form $(x_{n+1,r}^*, x_{n+1,r+1}^*)$ and the replicator dynamic for
491 group size n has an interior rest point in the interval $(x_{n+1,\ell}^*, 1)$, this implies that there
492 is exactly one interior rest point of the replicator dynamic for group size n in each of
493 the intervals $(x_{n+1,r}^*, x_{n+1,r+1}^*)$ for $r = 1, \dots, \ell - 1$. Therefore, for all $n = \underline{n}, \dots, \bar{n} - 1$
494 and $r = 1, \dots, \ell - 1$, we have

$$x_{n+1,r}^* < x_{n,r}^* < x_{n+1,r+1}^*. \quad (10)$$

495 In conjunction with the inequality $x_{n+1,\ell}^* < x_{n,\ell}^*$ established in the preceding paragraph,
496 Eq. (10) finishes the proof.

497 A.2 Proof of Proposition 2

498 Let ℓ denote the number of interior rest points of the replicator dynamic for a given
 499 group size $n \in N$. We begin by showing that the number of interior rest points of the
 500 replicator dynamic for group size $m \in N$ satisfying $m < n$ must be at least ℓ . This is
 501 trivially true for $\ell = 0$, so consider $\ell \geq 1$. By the same arguments as in the proof of
 502 Proposition 1, the replicator dynamic for group size $n - 1$ has at least one rest point in
 503 the interval $(x_{n,\ell}^*, 1)$ and, in case $\ell > 1$, at least one rest point in each of the intervals
 504 $(x_{n,r}^*, x_{n,r+1}^*)$ for $r = 1, \dots, \ell - 1$. As there are $\ell - 1$ such intervals, the replicator
 505 dynamic for group size $n - 1$ has at least as many rest points as the replicator dynamic
 506 for group size n . By a straightforward induction argument, it follows that the same
 507 conclusion obtains not only for group size $n - 1$ but for all group sizes $m \in N$ satisfying
 508 $m < n$.

509 Suppose that the number of interior rest points for group size \bar{n} is equal to the
 510 number of sign changes s of the gain sequences. It then follows from the argument in
 511 the previous paragraph that, for all group sizes $n \in N$, the number of interior rest points
 512 is at least s . On the other hand, the number of interior rest points of the replicator
 513 dynamic for group size n cannot be larger than the number of sign changes s of the
 514 gain sequence (Peña et al., 2014, Property 2). Hence, independently of group size the
 515 number of interior rest points is s .

516 The assumption that the regularity condition $dg_n(x^*)/dx \neq 0$ holds for all interior
 517 rest points implies that all roots of the polynomials $g_n(x)$ are simple. Therefore, for all
 518 group sizes $n \in N$ the number of interior rest points is either equal to the number of sign
 519 changes s of the gain sequences or less by an even amount Peña et al. (2014, Property
 520 2). It follows that the number of interior rest points for the replicator dynamics for
 521 two different group sizes either are equal or differ by an even amount. As it has been
 522 established above that the number of interior rest points cannot increase with group
 523 size, this observation finishes the proof.

524 A.3 Proof of Proposition 4

525 From Result 4.1 in Peña et al. (2014) the condition $\bar{g}_{\bar{n}} > 0$ (in conjunction with the
 526 assumption on the sign pattern of the gain sequences) is sufficient to imply that the
 527 replicator dynamic for group size \bar{n} has two interior rest points $x_{\bar{n},1}^* < x_{\bar{n},2}^*$ with the
 528 first of these being unstable and the second stable. As the gain sequences have two sign
 529 changes for all $n \in N$, Proposition 2 then implies that the replicator dynamic for any
 530 group size $n \in N$ has two interior rest points with the same stability pattern. From
 531 Proposition 1, the inequalities $x_{n+1,1}^* < x_{n,1}^*$ and $x_{n+1,2}^* < x_{n,2}^*$ hold for all n satisfying
 532 $\underline{n} \leq n < \bar{n}$. The remaining inequality in Eq. (4) follows from Eq. (10) in the proof of
 533 Proposition 1 in Appendix A.1.

534 A.4 Proof of Proposition 5

535 The existence of a unique interior rest point x_n^* and its stability for all group sizes is
 536 immediate from Result 3 in Peña et al. (2014).

537 Fix n satisfying $\underline{n} \leq n < \bar{n}$ and let

$$h_n(x) = \sum_{k=0}^{n-1} \binom{n-1}{k} x^k (1-x)^{n-1-k} d_k^{n+1}. \quad (11)$$

538 Observe that the assumption $d_k^{n+1} \leq d_k^n$ for all $k = 0, 1, \dots, n-1$ implies $h_n(x) \leq g_n(x)$
 539 for all $x \in [0, 1]$, where $g_n(x)$ has been defined in (6).

540 An argument identical to the one that we have used to obtain Eq. (3) in Appendix
 541 A.1, yields

$$h_n(x) = g_{n+1}(x) - \frac{x}{n} \frac{dg_{n+1}}{dx}(x). \quad (12)$$

542 As the rest point x_{n+1}^* is stable, Eq. (12) implies $h_n(x_{n+1}^*) > 0$ and therefore $g_n(x_{n+1}^*) >$
 543 0 . By the stability of the rest point x_n^* , we have $g_n(x) < 0$ for all $x \in (x_n^*, 1)$. Therefore,
 544 the inequality $g_n(x_{n+1}^*) > 0$ implies $x_{n+1}^* < x_n^*$, which is the desired result.

545 A.5 Proof of Proposition 6

546 From Result 4.1 in Peña et al. (2014) the condition $\bar{g}_n > 0$ (in conjunction with the
 547 assumption on the sign pattern of the gain sequences) is sufficient to imply that for all
 548 group sizes $n \in N$, two interior rest points $x_{n,1}^* < x_{n,2}^*$ exist with $x_{n,1}^*$ being unstable
 549 and $x_{n,2}^*$ being stable. The proof is then finished by observing that the same argument
 550 as in the proof of Proposition A.4 implies the inequality $x_{n+1,2}^* < x_{n,2}^*$ for all n satisfying
 551 $\underline{n} \leq n < \bar{n}$.

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