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Altruism Displays a Harmonic Signature in Structured Societies

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Abstract

Several parameters combine to govern the nature of agents interactions in evolutionary social simulations. Previous work has suggested that these parameters may have complex interplay that is obscured when they are not analyzed separately. Here we focus specifically on how three population-level parameters, which govern agent interactions, affect levels of altruism in a population. Specifically we vary how frequently agents interact in a generation, how far along a network they may interact, and the size of the population. We show that the frequency with which an agent interacts with its neighbors during a generation has a strong effect on levels of evolved altruism – provided that those pairings are stochastic. When agents interact equally with all of their neighbors, regardless of how often, minimal levels of altruism emerged. However, a spurious harmonic signature in the level of altruism resulting from the interplay of the benefit-cost ratio of an altruistic act and the number of agent interactions per generation. While the level of altruism is generally an increasing function of the number of pairings per generation, at each instance where pairings equals a multiple of the benefit-cost ratio a sharp discontinuity occurs, precipitating a drop onto a lower-value function. We explore the nature of these discontinuities by examining the temporal dynamics and spatial configuration of agents. Finally, we show that rules for the evolution of cooperation that are based on network density may be inadvertently missing effects that are due to the frequency of interactions and whether those interactions are symmetrical among neighbors.

Keywords: Tags, Thresholds, Altruism, Evolution, Cooperation, Social Harmonics

Introduction & Background

1.1 Computer simulated societies have long been used to understand the emergence of group-level phenomena such as cooperation, altruism, segregation, and coordination. It has been well established that embedding simulated agents in a structured society, either spatially or more generally in a network, can lead to very different outcomes than when the population is well mixed (Broid & Richardson 2002; Hauert 2001; Jun & Sethi 2007; Németh & Takács 2007; Nowak & May 1992; Ohtsuki et al. 2006; Perc et al. 2013). This added structure has proven to be an important addition to earlier social simulations and has led many authors to conclude that the localization of interactions resulting from structured societies is a key to the emergence of cooperation (Flache 2002; Ohtsuki et al. 2006; Shuttles 2012; Takács et al. 2008; Wilson & Wilson 2007).

1.2 A small number of population-level parameters often govern how agents interact in simulations with structured societies. Some of these parameters include population size, neighborhood size for both interactions, and number of agent interactions per generation. Yet the complex interplay between these structure-related parameters is poorly understood. Indeed there continues to be disagreement regarding the effect that network density has on cooperative outcomes. One body of research finds that when a population is more densely connected the likelihood of cooperation increases (Marwell & Oliver 1993; Opp & Gem 1993). As Jun and Sethi (2007, p. 625) assert, “dense networks are more conducive to the evolution of cooperation.” On the other hand, recent research suggests the opposite, demonstrating that denser networks inhibit cooperation in structured populations (Flache 2002; Hauert & Doebeli 2004; Shuttles 2012; Takács et al. 2008). This ability of network density to both facilitate and inhibit cooperative outcomes has been noted in informal networks, dynamic networks, and especially in networks in which “deviant” cliques can emerge, which are otherwise largely isolated from the larger population (Flache 1996; Flache & Macy 1996; Heckathorn 1996).

1.3 In a recent study Shuttles and Hales (2013) showed that parameters governing how agents interact with one another had significant effects on rates of altruism. These parameters included population size N, neighborhood radius R, and pairings (or interactions) per agent per generation P. Results of their study showed that decreasing neighbor radius or increasing population size led to higher donation rates. These parameters are related in that they reduce the proportion of agents within the total population with which a single agent is likely to interact. Thus, restricting the number of an agent’s potential partners had a positive effect on levels of altruism, confirming the predicted effects of localization of interactions discussed in the previous paragraph. This result also reinforces previous findings that increasing neighborhood radius or group size can inhibit cooperative outcomes (Flache et al. 2004; Olson 1965).

1.4 Decreasing the third parameter in Shuttles and Hales study, pairings, also decreases the number of potential interaction partners. However, the authors found that decreasing the number of pairings per generation had a negative effect on altruism and thus appeared to contradict the predictions of localized interactions. While the authors speculated on possible reasons for this seeming contradiction, they did not elaborate as the focus of their study was on changing altruistic behaviors.

1.5 Here we implement the model of Shuttles and Hales (2013) to perform a detailed parameter sweep of the three interaction-governing parameters N, R, and P. This simulation protocol, developed to study the evolution of altruistic behavior in structured societies, was based on an earlier simulation of Riolo et al. (2001) that was subsequently modified by Roberts and Sherratt (2002) and Specter and Klein (2006). In each of these studies, a ring was used to structure societies and a parameter determined how far along that ring agents were able to interact. In the case of Riolo et al, agents could interact along the entirety of the ring making their system equivalent to a well-mixed society.

The simulation model

2.1 The present model consists of a population of N agents arrayed in a ring. Every agent i is surrounded by a neighborhood consisting of all other agents within R links of i. Thus, every agent has 2R neighbors. Every agent i has a neighborhood consisting of all other agents within a distance of 2R.

2.2 Each agent is represented by three strategy components. Two of these components are real-valued numbers: a tag T (0, 1] and a behavioral trait called a “tolerance threshold” T (10^-5, 1]. Both tag and tolerance are initially drawn randomly, with uniform distribution, from [0, 1]. Note that the tolerance T may subsequently evolve beyond its initialization range. The third component is a binary trait which determines whether the agent is a cheater or not, the ramifications of which are explained below.

2.3 Prior to a new generation, the simulation’s pseudo-random number generator is initialized with the system time as a seed. In each generation each agent i is paired with P other agents chosen randomly with replacement from its neighborhood. Thus, every generation has a total of P*N pairings. During each pairing the agent determines whether to make a donation to a recipient agent j (donations to j if their tags are sufficiently similar. Agent i assesses similarity by comparing its own tag (i) with the tag of agent j). If the difference between the tags is less than or equal to its tolerance threshold T, then a donation takes place from i to j. Hence a donation occurs when |T - T|<T. If a donation does occur j accumulated payoff increases by the benefit of d and i s payoff decreases by the cost of c. In results presented here b > c. It is important to note that since partners are chosen randomly with replacement, when neighborhoods are small relative to N it becomes increasingly likely that a given neighbor will be chosen by an agent multiple times during a generation.

2.4 Donation therefore represents altruistic behavior since it incurs a cost to benefit ratio. An agent with a high T will donate to others holding a wide range of tags but an agent with a low T will only donate to those with very similar tag values. In the extreme cases, an agent with T=0 will donate to all other agents whereas an agent with T=1 will only donate to those with tag values identical to its own. In cases where an agent’s T<<10^-5, the agent will not donate to any other agent.

2.5 After each generation the accumulated costs and benefits (net payoffs) of each agent is calculated and serves as the agent’s fitness. Evolution is then implemented in the form of a tournament selection process in which each agent compares its value of (d - c) to a randomly chosen neighbor j. If j > i, then i copies j’s trait; tag, tolerance, and cheater status. But if j < i, then i s strategy remains unchanged. j and i are effectively competing to occupy i’s space in the next generation. If j = i, the coin toss is used to determine which agent will have its traits copied into i’s position in the next generation. Thus, fitness is compared locally and not globally, in accordance with the Darwinian notion that evolution is driven by relative (local) fitness and not absolute (global) fitness.

2.6 Mutation is applied to newly copied values with probability m. Mutation of the tag involves its replacement with a new random tag uniformly chosen from [0, 1]. Mutation of the tolerance T is achieved by adding Gaussian noise to the old value with mean 0, standard deviation 0.01. If mutation results in T < 10^-6, T is reset to 10^-6.

2.7 The binary trait third trait noted above determines whether the agent is a normal donating member of the society (trait = true) or a cheater (trait = false). When an agent’s trait is false (cheater) its ability to donate is deactivated and its tolerance trait is ignored. The motivation for this additional trait is described in (Shuttles & Hales 2013) and permits the presence of an important real-world participant in social systems – the social parasite, which simply displays a suitable tag to altruistic donors and only collects benefits. This binary trait can be reversed through mutation, at the mutation rate m, allowing a resumption of altruistic behavior (assuming the agent’s tolerance is sufficiently high). Every simulation is initialized so that all agents have this binary trait turned on at the start. Hence, there are no cheaters at the beginning of a simulation and cheaters can only arise through mutation. Once present, cheaters may proliferate through mutation and may be selected for during the reproduction routine.
2.8 Unless otherwise stated, parameters are fixed in all simulations such that the benefit conferred during an altruistic act $b = 1.0$, cost to the altruistic donor $c = 0.1$, mutation rate $m = 0.01$, number of generations per simulation run $g = 30,000$, and number of replications (independent simulation runs with a different initial pseudo-random number seed) performed with each combination of parameters $= 100$. During the final 1,000 generations of each simulation, data were collected tabulating the percentage of interactions that resulted in an altruistic act (donation rate). These values were then averaged across all 100 replications of a parameter set. Standard deviations were relatively small and are not presented here.

2.9 Three independent replications were implemented by different researchers using only pseudo code as a guide. The initial model was implemented in Java and subsequently replicated in NetLogo, the latter of which is freely available online at the OpenABM repository (see: https://www.openabm.org/model/4510/version/2/).

Results and discussion 1: the complex interplay between $N$, $R$, and $P$

3.1 A sweep of parameters was conducted of the three variables $N$, $R$, and $P$ to test for their effects on mean donation levels. Results are discussed for each of the variables individually, keeping one of the other variables fixed and varying the third variable.

Effects of population size ($N$)

3.2 As population size increased, simulations produced higher mean contribution rates. However, the effect diminished with increasing population and, beyond a critical population size, there was little change in donation rates ceteris paribus (Figure 1).

![Figure 1. Effect of population size $N$ on altruistic donation rates. Donation rates vs. population size are presented using a number of different pairing values $P$ with (a) radius $R = 1$ and (b) radius $R = 12$. In general, donation rates increase with increasing population size but this effect diminishes as population continues to increase.](http://jasss.soc.surrey.ac.uk/18/3/2.html)

Effects of neighborhood radius ($R$)

3.3 At low values of $P$, increasing the neighborhood radius $R$ above 1 resulted in lower donation rates. This is consistent with the findings of Ifti et al (2004) who showed that cooperation in a spatial prisoners dilemma diminished with increasing neighborhood radius. However, with increasing $P$, maximum donation rates begin to occur at values of $R$ between 1 and 5 (Figure 2). The value of $P$ at which this movement to an internal maximum occurs depends on the population size, with larger populations shifting to an internal maximum at lower values of $P$. 

http://jasss.soc.surrey.ac.uk/18/3/2.html
Effect of pairings per generation ($P$)

In contrast to the response of increasing population, the response of donation rates to changes in pairings is richer and quite perplexing. Though donation rates initially increase with increasing pairings, rates make a sharp drop between $P = 9$ and $P = 10$ (Figure 3). Donation rates then increase slightly before another drop in the neighborhood of $P = 20$. 

Figure 2: Effect of neighbor radius ($R$) on altruistic donation rates. Donation rates vs. neighborhood radius $R$ are presented using a number of different population sizes ($N$) with pairings $P$ fixed at four different values. At low values of $P$, donations are maximized when $R = 1$. At higher values of $P$, the optimal radius is greater than 1.
Figure 3. Effect of pairings per generation on altruistic donation rates. Results are presented for 4 different population sizes and at 4 different neighbor radii for each of those population sizes. The sharp drop in donations above $P = 9$ is more pronounced (though diminishing) with increasing population size. Points A and B in figure (b) are explored further below in Figure 4.

3.5 To better understand the temporal dynamics underlying the results in Figure 3, donation rate vs. generation is presented for two cases. Fixing parameters $N = 90$ and $R = 1$, simulations were run at two pairings values, $P = 9$ and $P = 12$, corresponding to points A and B in Figure 3b. Results of these two simulations are presented in Figure 4 and give rise to several interesting points. First, donation rates do not converge over time in either simulation but appear to drift stochastically about a mean that is dictated by the simulation's pairing parameter $P$. Second, when $P = 9$, no generation had a donation rate of 0, meaning that some altruism existed in every generation. When $P = 12$, however, 118 of 1,000 generations had a donation rate of 0. This finding contributes to an explanation of why simulations with $P = 9$ evolved donation rates about twice as high as those when $P = 12$. Yet it remains to explain why lower donation rates are so much more prevalent at $P = 12$, especially as donation rates increase as $P$ increases from 1 to 9.

Figure 4. Donation rates observed across final 1,000 generations for two values of $P$. In both cases $N = 90$ and $R = 1$. The mean of each curve corresponds to a single simulation run of the 100 represented in points A and B in Figure 3b. Points C and D, both on the curve for $P = 9$, are explored further below in Figure 5.

3.6 To help explain the temporal differences in donation rates shown in Figure 4, the tag value of each agent in the two simulations presented in Figure 4 were tracked across all 1,000 observed generations. This allows us to plot an agent's tag versus the agent's position in the network so that we may visually inspect the clustering of trait values in a networked population. Results are presented in Figure 5 for two specific generations, selected because they were representative of generations with high and low donation rates. These graphs correspond to points C (generation 20,091) and D (generation 20,886) in Figure 4, which had donation rates of 0.04 and 0.77 respectively. Both points lie on the curve for $P = 9$. Results reveal much higher clustering and uniformity of tag values in the generation that experienced a high donation rate (Figure 5b). These results suggest that the clustering of trait values is an important driver of altruism in these simulations.
3.7 Two potential indicators of the degree of trait clustering within the population are the number of unique trait values and the largest group size of agents holding the same trait value. Accordingly, the mean number of unique tag values and the size of the largest group of any tag value were calculated across the 1,000 generations presented in Figure 4. For each of the curves ($P = 9$ and $P = 12$) the two measures of trait clustering were regressed against donation rates to understand their effects on altruistic outcomes (Table 1). While the size of the largest tag group appeared to have no relation to donation rates, the number of unique tags showed a discernible relationship to the donation rate within a generation. In addition, this relationship differed significantly between treatment with $P = 9$ and that with $P = 12$.

3.8 In general, both simulations had higher donation rates during generations containing a low number of unique tags. However, the strength of this effect was nearly twice as strong when $P = 9$ compared to $P = 12$ (see regression coefficients, Table 1). This is intriguing since the only difference between the two simulations was the number of pairings an agent initiated during each generation. And while this helps to further explain why mean donation rates overall were so much higher when $P = 9$ versus $P = 12$, it still does not explain why clustering was so much more prevalent under $P = 9$ or why the effect of this clustering was stronger.

Table 1: Comparison of two individual simulation runs, each using a different value for pairings ($P$), all other parameters equal. Mean results over the final 1,000 generations are shown (shown graphically in Figure 4).

|                       | $P = 9$ | $P = 12$ |
|-----------------------|---------|----------|
| Percentage of generations having 0 donations | 0.0%    | 11.8%    |
| Mean values over final 1,000 generations |         |          |
| Donation rate          | 0.32    | 0.11     |
| Number of unique tags  | 9.7     | 13.1     |
| Size of largest tag group | 34.7   | 26.4     |
| Simple linear regression – donation rate vs. number of unique tags | Coefficient | $R^2$ |
|                       | -0.0253 | 0.224    |
|                       | -0.0149 | 0.234    |

Note: In both cases $R = 1$, $N = 90$, and $b/c = 10$.

Results and Discussion 2: The benefit to cost ratio and harmonic signature

4.1 In all results thus far discussed the benefit and cost of an altruistic act were kept fixed at $b = 1$ and $c = 0.1$. Thus, a fixed benefit-cost ratio $b/c = 10$ has been used throughout. However, we note that in Figure 3, when $R = 1$ the sharp drops, or discontinuities, in donation rates with increasing pairings occur consistently at $P = 10$ and again at $P = 20$. This was true regardless of the population size or neighborhood radius. Realizing these values are multiples of $b/c$, we wished to further explore the effects of $b/c$ and to see if varying the ratio would change the points of discontinuity in donation rates. Therefore, keeping population size fixed at $N = 400$ and neighborhood radius fixed at $R = 1$, we ran a number of further simulations varying $P$ against different values for $b/c$. In all cases, $c = 0.1$ and $b$ was varied to generate different values of $b/c$.

4.2 Results shown in Figure 4a are striking. Not only do discontinuities appear precisely when the value of $P$ is a multiple of $b/c$, regardless of the value of $b/c$, the curves for differing values of $b/c$ appear to overlay one another so that a fundamental set of curves, consisting of discrete steps, would seem to predict all simulation outcomes. This second effect is more pronounced when lines corresponding to discontinuities have been removed (Figure 4b).
4.4 In effect, these different curves are "activated" depending on the value of simulation parameters $P$, $b$, and $c$, which determines which active range, and thus active is determined as follows:

- Curve A: $P^1(0, b/c)$
- Curve B: $P^1(b/c, 2b/c)$
- Curve C: $P^1(2b/c, 3b/c)$
- Curve D: $P^1(3b/c, 4b/c)$
- Curve E: $P^1(4b/c, 5b/c)$

4.5 Actual simulation results using two different values of $b/c$ are superimposed on this set of curves to present examples of their predictive power (Figure 8).

4.6 To assess the sensitivity of the transition point $P = b/c$ we analyzed points just before and after the first transition point for $P = 10$. Using parameters $P = 10$, $R = 1$, and $N = 400$, an additional 100 simulations were run at $b/c = 10 \pm \epsilon$ where $\epsilon = 10^{-5}$. Even with these minute deviations from the transition point, results corresponded to the different curves as predicted in Figure 8b, indicating that the transitions described above occur precisely at multiples of $b/c$. Specifically, with $P = 10$, mean donation rates were 0.240 when $b/c = 10, 0.0000006$, 0.126 when $b/c = 9.999999$, and 0.182 when $b/c = 10$ exactly.

4.7 However, the fact that a donation rate does occur precisely when $b/c = 10$, and the fact that its value is between the donation rates resulting from small deviations above and below $b/c = 10$ (Figure 3b), indicates that it is possible, and even probable, that what we have termed a discontinuity is in fact a very sharp, yet continuous, transition between curves shown in Figure 7, with an inflection point at $P = 4b/c$. The resolution of this question requires simulations with even smaller deviations from the transition point that were beyond the scope of our simulation, and thus this question remains an important point to address in future work.
To remain consistent with the models on which this study is based (see Introduction), we have used a ring to structure the societies in our simulations. However, it is important to consider how our results with a ring structure may be manifest for other network types, particularly the more complex networks found in real societies. Thus, in addition to the ring configuration used thus far in this study, we also implemented our simulation protocol on other commonly studied configurations. With $R = 1$ and $k$ representing the network’s average nodal degree, these other structures include a regular lattice ($k = 4$), regular lattice ($k = 8$), random regular network ($k = 4$), and scale-free network ($k = 4$). Scale-free networks were generated through growth by preferential attachment (Barabási & Albert 1999), starting with a 3-node complete network and adding new nodes via two new links. Though we did not analyze results from these additional configurations to the extent that we did with rings ($k = 2$), our limited results show that the discontinuities in donation rates when $P$ equals a multiple of $b/c$ are robust to configuration type. Results for various social structures, with parameters $N = 400$, $R = 1$, $b/c = 8$, are shown in Figure 9. On all population structures, donation rates rise with increasing $P$ until just before $P = b/c$ drop, rise again until just before $P = 2b/c$, and then drop once more. These results suggest that there exists a fundamental set of predictive curves, similar to those in Figure 7, for many social structures, including those with variance in individual nodal degrees (i.e. scale-free networks). The study of this phenomenon on more complex networks thus presents a promising future direction for this work.

**Relation to similar findings**

5.1 The sharp drop in donations with increasing pairings that occurs when $P = b/c$ is related to a previously suggested rule for the evolution of cooperation. Ohtsuki et al. (2006) proposed that in structured societies, cooperation will evolve when $k < b/c$, where $k$ is the average nodal degree of agents embedded in a network. Here, we show that maximum levels of altruism occur when $P < b/c$ as $P \rightarrow b/c$.

However, if a simulation is constructed in such a way that each agent interacts with every one of its neighbors exactly once, then our parameter pairings and the average nodal degree are equivalent so that $P = k$. In such cases, it is impossible to determine whether the condition $k < b/c$ or the condition $P < b/c$ is the primary rule for the evolution of cooperative behavior. Thus simulations or experiments (e.g. Rand et al. 2014) that are structured so that every agent interacts exactly $n$ times per generation with every one of its neighbors – what we will refer to here as symmetrical pairing – may inadvertently obscure the effects of pairings. It is therefore important to disentangle the relative importance of these conditions by parsing the individual effects of $k$ and $P$ on altruistic outcomes.

5.2 In attempting to separate these effects, note that the degree of an agent in our ring structure is determined solely by the neighborhood radius such that $k = 2R$. Thus, results presented in Figure 2 on the effects of varying $R$ with other parameters held constant, also show the effects of varying $k$. In Figure 2, $b/c = 10$ and therefore if Ohtsuki’s rule is manifest, there should be an impact when $k = 10(R = 5)$. As Figure 2 reveals, the role of $k$ is quite complicated. When pairings $P$ are low or population size $N$ is low, altruism is maximized when $k = 2$ (and $R = 1$), its minimum possible value. In these cases altruism decreases monotonically with increasing $k$ and there is no notable effect at the point where $k = b/c$. However, at higher values of $P$ (or both), results are more complicated and not easily interpreted. Altruism does appear to reach maximal levels in the vicinity of $k = b/c$ when $N = 400$ and $P = 10$.

5.3 For a possible explanation of this finding, recall that our pairings are stochastic and thus interactions are heterogeneous. Yet this heterogeneity diminishes as the number of pairings increases or when the population becomes larger – the same conditions under which $k$ begins to affect rates of altruism. Thus parameters that shift the nature of interactions from highly heterogeneous to highly symmetrical appear to moderate the efficacy of the $k < b/c$ rule. As Figure 6 shows, high values of $P$ tend to reduce donation rates in the non-monotonic form related to $b/c$. Since agents accumulate net fitness and interactions are stochastic within the agent radius we hypothesize that very high $P$ will result in altruism levels that equate to deterministic and symmetric interactions. This is true because as $P \rightarrow \infty$, stochastic pairing approaches symmetrical pairing.

5.4 Thus, to more effectively parse the differential effects of $k$ and $P$ we also examine the differential effects of stochastic vs. symmetrical pairing. We do so by implementing supplemental simulations on a ring network in which each agent interacts with each of its two neighbors exactly $n$ times, where $n = (1, 2, 3, 4, \ldots)$. Thus $P = 2n$ in these simulations with symmetrical pairing. Other parameters were fixed such that $R = 1, N = 90$, and $b/c = 10$. Because the neighbor radius is fixed, the average nodal degree is also fixed at $k = 2$. These runs produced mean donation rates of approximately 0.05 for all $P$(Figure 10). Thus, when pairings are symmetrical, the value of $P$ has no discernible effect on levels of altruism, which remained at minimal levels.

5.5 Yet when $k$ and $b/c$ remain fixed but interactions are stochastic, donation rates vary with $P$ and are maximized when $P = b/c$ as $P \rightarrow b/c$. Because the condition $k < b/c$ is fixed here, the variation in donations, and particularly its peak when $P$ approaches $b/c$, is an effect of $P$ alone. This suggests that the stochastic nature of pairings in our simulation and its effect on cumulative fitness is highly significant in producing the donation rates presented previously.
This finding raises interesting issues concerning interaction contingency and fitness calculation in evolutionary systems. By accumulating fitness, rather than averaging over interactions, the contingent aspects of those interactions with respect to reproduction are magnified. One might expect this to produce noise. Yet in the model presented here we see a robust and interesting donation function emerge.

Note also that the heterogeneity in interactions may be more generally represented in a network by weighting the network’s links. Thus, the use of symmetrical pairings in simulations and experiments is analogous to using unweighted networks in those cases, while this study’s stochastic interaction is analogous to using a weighted network, where the weights are stochastic. It is obvious that in human social networks interactions are not symmetrical. Yet, neither are they strictly random in their frequency. It is likely that a more realistic implementation than either of these cases is the use of a heterogeneously weighted network where the distribution of weights is empirically informed. This presents a fruitful future direction for work on the evolution of cooperative behavior in networked populations.

A note on mutation

Because the effect of mutation rates in similar simulations has been previously examined (Hales 2005; Shutters & Hales 2013; Spector & Klein 2008), and because our goal was to isolate the effects of parameters that govern interactions, this study used a fixed value for the probability of mutation of $m = 0.01$. However, to determine whether our findings regarding a harmonic signature are robust to changes in mutation rate we ran a further set of simulations using our study’s mutation rate along with two additional values of mutation rate $m = 0.1, 0.01, 0.001$. In all cases $N = 90$ and $bc = 10$. Results presented in Figure 11 show that when mutation rate was relatively low ($m = 0.001$) varying $P$ had no effect on donation rates and only minimal donation rates evolved. Relatively high mutation rates ($m = 0.1$) did lead to a noticeable drop in donation rate when $P > bc$; especially when neighborhood radius was larger. However, this drop in donations when $m = 0.1$ was relatively small compared to treatments with $m = 0.01$.

Conclusion

Here we have used a simple simulation to test several parameters that govern interactions between agents in a structured society. We have shown that the important parameters, population size, number of neighbors, and number of interactions per generation exhibit a complex interplay and disentangling the effects of these different attributes thus presents a very promising direction for future research. Importantly, we show that when the structure of potential interactions (the static network) and the structure of actual interactions are equivalent, as is often the case in these types of simulations, important effects are hidden. When interactions are heterogeneous, as they are through the stochastic application of pairings used in our simulation, significantly higher levels of altruism emerge compared to when pairings are symmetrical.

To state this in a more general way: where static networks structure a population of agents by constraining interactions between agents it is important to contrast this with the actual realized interactions. The actual interactions in any given time period constitute a different network from the static one that is dynamically realised through agent behaviour combined with the static network. It is consequently important for the interpretation of such models to be clear about the difference between the static "structural network" and the dynamic "interaction network". The latter will often be implicit and emergent from agent behaviour. In much work these two networks are conflated and this can make results hard to interpret and understand.

Finally, we reported a curious harmonic signature in simulation outcomes resulting from the interaction of pairings and benefit-cost ratios. This last phenomenon, in particular, is in need of further investigation to understand its underlying cause and we welcome the chance to explore this further with our colleagues in the social simulation community.

Figure 10: Random vs. Symmetrical Pairings. In both cases $N = 90$, $R = 1$, and $bc = 10$. Under symmetrical pairings, varying the value of $P$ appears to have little effect.

Figure 11: Mutation rate vs. Pairings. Results are presented with neighborhood radius (a) $R = 1$ and (b) $R = 5$. In both cases $N = 90$ and $bc = 10$. The lower mutation rate led to no changes in aggregate behavior while the higher mutation rate led to a very slight drop in donations when $bc = P$.

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Notes

1 Tuning parameters z used for curves (A, B, C, D, E) were (16.6, 8.5, 12.5, 15.6, 18.5) respectively. These were determined by simple visual estimation since our intent was only to demonstrate robustness of a set of curves across different parameter sets.

Appendix

Algorithm pseudo code and parameter details

7.4 The simulation model used in this article is described below. The simulation replicates several existing models under certain parameter settings and provides novel results in other cases. Under the parameters we list the range of values used for each parameter. To find the specific values for a given result see the text of the relevant section of the article.

Parameters:

- G = Number of generations = 30,000
  - The number of generations to run the simulation for.
- N = Population size = [10..1000]
  - The number of agents in the population.
- b = Benefit of donation = 1
  - The amount of fitness benefit an agent receiving a donation obtains.
- c = Cost of donation ? [0..1]
  - The amount of fitness cost applied to an agent making a donation.
- P = Pairings ? [1..100]
  - The number of random pairings made for each agent in each generation.
- m = Mutation rate ? (0.001, 0.01, 0.1)
  - The probability that agent traits will be mutated during reproduction to the next generation.
- R = Radius of interaction ? [1..1000]
  - The radius around the agent from which random pairings with other agents are drawn. Constraint: 0 ≤ R ≤ N/2. When R = N/2 = 50, radius is entire population (as in the RCA model). When R = 1, radius is only immediate two neighbors (as in SK model).

Agent state variables:

- t = tag value of agent i. t ? [0..1]
  - The tag value is a real number that is attached to each agent and is observable by other agents.
- T = tolerance value of agent i. T ? [-10^6, 8].
  - The tolerance value is a real number stored by each agent, used, in combination with tag values, to determine if to make a donation to another agent.
- BT = binary trait of agent i. BT ? {0, 1}.
  - The binary trait, stored by each agent, determines if to make donations or not: 0 = turn off all donation, 1 = donate using tag / tolerance mechanism.
- f = fitness of agent i. f ? R.
  - Agents accumulate a fitness value in each generation through interactions involving costs and benefits.

Main Outline Algorithm:

// Initialize population of agents
Initialize population of agents () // see function below
Loop for G generations
  Loop for each agent A in population N
    Loop for P pairings
      Select a random agent B within agent A’s radius R
      Agents Interact (A, B) // see function below
    End loop
  End loop
End loop
Reproduce Population () // see function below
End loop

Functions used in main outline algorithm:

// Initialize population of agents
Initialize population of agents ()
  Create a ring of size N
  Loop for each agent A in population N
    Place agent A at next free position on ring
    Set tag (tA) and tolerance (TA) to uniform random values [0..1]
    Set binary trait to 1 (BT A = 1)
    Set fitness of agent to zero
  End loop
End Function

// Agents A and B interact. A decides if to make a donation to B
Agents Interact (agent A, agent B)
  If binary trait of A is turned on (BT A = 1) then
    If tags of A and B are within tolerance of A (|T A - tA| ≤ TA) then
      Fitness of B is increased by benefit (fB = fB + b)
    Fitness of A is decreased by cost (fA = fA - c)
  End if
End function

// Reproduce a new population based on fitness of agent in current population
Reproduce Population ()
  Create a new ring of size N for the new population
  Loop for each agent A in population N
    Select a random agent B within agent A’s radius R
    Reproduce agent (A or B) with higher fitnessB/f as agent C

http://jasss.soc.surrey.ac.uk/18/3/2.html
21/10/2015
Agent C is placed in the same location on the ring as agent A. With probability \( m \) mutator* tag (\( t_C \)) and tolerance (\( T_C \)) of agent C with probability \( m \) mutator* binary trait (\( B(t_C) \)).

End loop.

New population becomes the current population.

All agents in new population have fitness initialize to zero.

End function.

Notes:

#Note: if the fitness of A and B are the same a random selection is made.

*Note: mutation takes the following form:

A tag (\( t \)) is replaced with a uniformly random value \([0, 1]\).

Tolerance (\( T \)) is changed by adding Gaussian noise with mean 0 and standard deviation 0.01. If \( T < -10^{-5} \) then \( T \) is reset to \(-10^{-5}\).

A binary trait (\( B(t) \)) is flipped (such that 1 becomes 0 and vice versa).

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