Spatial Population Expansion Promotes the Evolution of Cooperation in an Experimental Prisoner’s Dilemma

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Summary

Cooperation is ubiquitous in nature, but explaining its existence remains a central interdisciplinary challenge [1–3]. Cooperation is most difficult to explain in the Prisoner’s Dilemma game, where cooperators always lose in direct competition with defectors despite increasing mean fitness [1, 4, 5]. Here we demonstrate how spatial population expansion, a widespread natural phenomenon [6–11], promotes the evolution of cooperation. We engineer an experimental Prisoner’s Dilemma game in the budding yeast Saccharomyces cerevisiae to show that, despite losing to defectors in nonexpanding conditions, cooperators increase in frequency in spatially expanding populations. Fluorescently labeled colonies show genetic demixing [8] of cooperators and defectors, followed by increase in cooperators frequency as cooperator sectors overtake neighboring defector sectors. Together with lattice-based spatial simulations, our results suggest that spatial population expansion drives the evolution of cooperation by (1) increasing positive genetic assortment at population frontiers and (2) selecting for phenotypes maximizing local deme productivity. Spatial expansion thus creates a selective force whereby defector-enriched demes overtake neighboring defector-enriched demes in a “survival of the fastest.” We conclude that colony growth alone can promote cooperation and prevent defection in microbes. Our results extend to other species with spatially restricted dispersal undergoing range expansion, including pathogens, invasive species, and humans.

Results and Discussion

Cooperation forms the basis for numerous complex phenotypes, from cell-cell communication and biofilm formation in microbes to nest construction in multicellular species [12, 13]. Explaining how cooperation evolves despite the direct fitness advantage gained by “free riding” remains a central challenge in biology and the social sciences [1–3]. This is particularly true in the Prisoner’s Dilemma game [1, 4, 5], where cooperators always lose in direct competition with defectors, leading to a “tragedy of the commons” [14]. Nearly half a century of research on social evolution has offered insight into this dilemma, with inclusive fitness theory focusing attention on the importance of high genetic relatedness [15], multilevel selection theory highlighting the role played by competition among social groups [16, 17], and spatial games showing the importance of population structure [18–21]. Here we bring these three frameworks together, demonstrating that spatial population expansion creates a setting where both relatedness and intergroup competition are amplified, promoting the evolution of cooperation.

Spatial population expansion occurs when a species spreads outward to fill vacant ecospace or to supplant resident species, resulting in increased geographic range. Human migration out of Africa [7] is one salient example; more generic examples include ecological invasions, epidemics, growth of microbial colonies, and expansion due to habitat modification caused by environmental disturbance or climate change [8–11]. Spatially expanding populations form a propagating density wave called a “Fisher wave” with a constant speed proportional to the square root of the mean growth rate (i.e., Malthusian fitness) of subpopulations at the front [22]. Genetically heterogeneous populations may also form an “allele frequency” wave representing the spatial spread of alleles [22] (see Figure S1 available online).

A small number of colonists initiate new subpopulations at the leading edge of the propagating Fisher wave, creating a repeated series of genetic bottlenecks or “founder effects” that cause stochastic loss of genetic diversity at frontiers [23, 24]. Because cooperation is favored under conditions of high genetic relatedness [15], range expansions could thus in principle favor the evolution of cooperation. However, there are at least three complicating factors. First, within-subpopulation selection favoring defection opposes genetic demixing, potentially preventing high defector-relatedness from ever arising. Second, even with high relatedness it is not clear what selective force, if any, favors cooperation over defection in expanding populations. Finally, almost any genotype that stochastically fixes at the front can increase in frequency via “surfing” [6, 10, 24, 25]. Other genotypes that stochastically fall behind this front cannot typically catch up, even if they are more fit. This is because they expand outward in a trailing allele frequency wave traveling at a speed determined by the difference in fitness between defector and cooperator genotypes, WD − WC, which will often be much smaller than mean absolute fitness. For social traits, a genotype fixed at the frontier will outrun genotypes in the population interior and increase in global frequency provided that (1 + b)/2 > c, where b is the social effect and c the direct fitness effect of the leading genotype (see section “Heuristic Description of Model and Derivation of Analytical Results” in Supplemental Information). The social effect here refers to the fitness increment or decrement received by an individual from social partners (e.g., the benefit of the public good), whereas the direct effect is the fitness increment or decrement accrued to an individual for engaging in a social behavior (e.g., the cost of producing the public good). Note that this condition (1 + b)/2 > c can be satisfied even when b < 0, and thus in principle surfing may promote cooperation’s opposites, selfishness and spite [26], including spite against relatives. Given these complications, it is not clear whether spatial expansion will in fact promote the evolution of cooperation.
To test the effect of spatial expansion on defector/cooperator dynamics, we engineered an experimental Prisoner’s Dilemma game using cooperative sucrose metabolism in haploid, vegetatively growing strains of the budding yeast *Saccharomyces cerevisiae* [27]. Yeast secrete the exoenzyme invertase in order to digest the disaccharide sucrose, which cannot easily be imported into the cell, forming monosaccharides that are readily imported. In our strains, sucrose cannot be imported at all due to disruption of the genes *MAL12* and *MAL22* [28]. Because digestion occurs externally, invertase producers (“cooperators”) create a public good that is exploitable by nonproducers (“defectors”), who gain a relative fitness advantage by not paying the fitness cost of production [27, 29].

We engineered a fluorescently marked defector strain by deleting the invertase gene *SUC2*.

We note that in minimal sucrose medium (YNB + 2% sucrose), competitions between Suc2⁺ and Suc2⁻ strains in shaken liquid culture were previously found to follow Snowdrift game dynamics [29]. In a Snowdrift game, the rarer type (regardless of whether it is a cooperator or a cheater) has a fitness advantage, leading to stable maintenance of both cooperators and defectors [1, 30]. The maintenance of cooperation is therefore easily ensured, in contrast to the Prisoner’s Dilemma game, where the maintenance of cooperation is much more difficult to explain [1, 30]. In addition, cooperators in a Snowdrift game have a colonization advantage over defectors because defectors cannot colonize habitat unoccupied by cooperators [28]. This facilitates colonization ability and cooperation by linking both to a single genotype. Because spatial expansion is already known to select for colonization ability [31, 32], such linkage would prevent us from concluding that spatial expansion favors cooperation per se rather than superior colonization ability. By contrast, defectors in a Prisoner’s Dilemma game do not require the presence of cooperators to colonize new habitat, making it possible to disentangle selection for cooperation from colonization ability.

We therefore used two approaches to construct a Prisoner’s Dilemma from this system. First, we eliminated the rare advantage of cooperators that is necessary for Snowdrift dynamics by conducting competitions in medium (YEP + 2% sucrose) in which our defector strains could grow in the absence of cooperators (most likely by consuming amino acids available in YEP, although growth is slower than for cooperators; Figure 1A). This environment also eliminates the difference in colonization ability between cooperator and cheater strains, in that cheaters no longer require the presence of cooperators to colonize the frontier (green line in Figure 1A). Next, we engineered a defector strain resistant to cycloheximide, a translation-inhibiting drug that limits growth by binding to ribosomal subunit Cyh2. This creates a system in which we can experimentally impose a tunable “cost of cooperation” by varying the level of cycloheximide in the growth medium. Specifically, increasing the cycloheximide concentration slows the growth of cooperators but not the resistant defectors, leading to an increased “cost of cooperation.”

When mixed with our defector strain in an unstructured environment (shaken liquid culture), our cooperator strain declines at all frequencies when a cost of cooperation is imposed, despite having a superior growth rate over defectors in pure culture (Figures 1 and S3). These results are consistent with Prisoner’s Dilemma evolutionary dynamics. Unlike in a Snowdrift game, any increase in frequency of cooperators in our experiments is not due to rare cooperator advantage.

To determine whether spatial expansion can promote cooperation in our experimental Prisoner’s Dilemma, we initiated spatial expansions by spotting a droplet of mixed cooperator/defector cultures onto solid medium (YEP + 2% sucrose + 2% agar) for a range of imposed costs (see Supplemental Experimental Procedures). Spatial diffusion of cells in *S. cerevisiae* is caused when cellular growth generates an outward force leading to radial spatial expansions of colonies [8]. Note that expansion is not caused by active cell motility in this system, as yeast lack motility. Relative frequency measurements taken using flow cytometry show that cooperators initially decline in frequency at a rate consistent with that of well-mixed liquid competitions but then increase in frequency as expansion proceeds (Figure 2C). Likewise, image analysis of fluorescently labeled colonies shows low cooperator frequency near the initial site of inoculation (the “homeland”) but increasing frequency with increasing distance from the homeland (Figure 2E). Lattice-based spatial simulations of a Prisoner’s Dilemma show the same spatiotemporal dynamic of initial decline in cooperator frequency followed by increase as expansion proceeds (Figure 2D). Cooperators invade when rare over a range of imposed costs (Figure 3). Furthermore, when the benefit of cooperation is removed by competing strains on glucose medium, the cooperator strain no longer increases in frequency upon spatial expansion (Figure S4).

These data clearly demonstrate that spatial expansion can promote the evolution of cooperation.

How does spatial expansion promote cooperation? Fluorescent colony images reveal the formation of discrete sectors of fixed genotypes (Figures 2A, 2B, and 3), which is the colony-level signature of genetic demixing [8]. Thus, spatial
expansion can lead to high positive assortment of cooperators via genetic demixing, even though this assortment is opposed by selection favoring defectors within demes (Figure 1B; Figure S1). Local fixation of cooperator sectors despite counterselection within demes is analogous to surfing by deleterious mutations [6, 10, 24, 25] and requires similar conditions to obtain. In other words, spatial expansion leads to the formation of uniform sectors of cooperators or defectors, increasing genetic relatedness of nearby individuals. This diminished local genetic diversity reduces the direct competition between cooperators and defectors (see also [33]), thereby mitigating the principle selective advantage of defection.

We note that genetic demixing (i.e., “sectoring”) is particularly clear in our experimental yeast system because yeast lack motility and “dispersal” of offspring is local. In other systems, movement of individuals and dispersal of offspring can in principle blur sector boundaries and oppose demixing at the frontier. In the extreme case where movement and dispersal are very long range, the spatial sectoring we describe here will not occur, and neither our analysis nor our results will generalize to this situation. In real populations, however, movement and dispersal are usually spatially restricted: a migrant is more likely to disperse nearby than far away. In this case, genetic demixing will occur provided that outward range expansion is sufficiently rapid compared to the rate of dispersal between occupied demes across sector boundaries (i.e., perpendicular to the expansion direction) [24, 25, 34]. In nature, species as diverse as rabies virus [35] and humans [36] show genetic signatures of expansion-associated demixing and sectoring, suggesting that the phenomenon we describe here may apply more generally. To the extent that spatial expansion-associated genetic demixing is possible in a species, the mechanism we describe here promoting the evolution of cooperation will also be possible.

Once cooperators establish habitat, their overall productivity will be higher than that of defector sectors provided the fitness benefit of cooperation exceeds the cost of cooperation, b > c (Figure S2; Heuristic Description of Model and Derivation of Analytical Results). When this is true, cooperators will expand radially faster than neighboring defector sectors, leading to a corresponding expansion of the boundaries of the cooperators at the expense of neighboring defector sectors as we see in our experiments (Figures 2B, 2E and 3). This leads to an overall increase in cooperator frequency and suggests that range expansion creates a force of natural selection favoring the “survival of the fastest.” This force acts to promote genotypes supporting maximal group productivity, because high-productivity sectors expand at a faster rate, allowing them to outcompete lower-productivity sectors.

We turn to stochastic, lattice-based spatial simulations to further test the “survival of the fastest” hypothesis. To test whether “survival of the fastest” is indeed necessary for range expansion to promote cooperation, we eliminate this force by restricting expansion to one dimension in our spatial simulations. In one dimension (e.g., a linear stepping-stone model), an expanding subpopulation has no neighboring subpopulations to compete with, so that intergroup competition is absent. In this case, we find that the probability of cooperator establishment at the front is never greater than the neutral probability of establishment (which is equal to the initial frequency of the allele, \( p_0 \)) and declines with increasing cost (Figure 4). Put differently, cooperators can only outrun defectors in a one-dimensional Prisoner’s Dilemma if they randomly take over the frontier, an outcome uniformly opposed by selection (Figure S2; Heuristic Description of Model and Derivation of Analytical Results).
In contrast to the one-dimensional case, the probability of cooperators fixing at the frontier is substantially higher in two dimensions, where subpopulations compete with neighbors for occupancy of uncolonized habitat (Figure 4). Two-dimensional expansions are also more efficient at purging deleterious alleles from frontiers, as seen in the comparison of the black and gray dashed lines in Figure 4. Our data support the conclusion that two-dimensional spatial expansions generate selection at the frontier for genotypes that maximize group productivity, because these genotypes lead to the greatest expansion velocity of the front, allowing cooperator-enriched demes to overtake defector-enriched demes. Spatial expansion generates both conditions necessary for natural selection: heritability (positive assortment of social strategies, making one's social environment heritable) and differential success (“survival of the fastest”).

Microbes possess a multiplicity of cooperative phenotypes [12], and rapid cell division in conjunction with large colony sizes makes the repeated emergence of defector mutants cooperation more generally and may allow already cooperative species to shed social parasites, so long as the pattern and rate of dispersal and reproduction allow for genetic demixing upon expansion. It is possible that reduced cheater load upon expansion may accelerate biological invasion by cooperative species, with potential implications for biological control. Stochastic demixing may also occur with culturally transmitted phenotypes, such that range expansion may have been important in the spread of cultural norms facilitating cooperation in humans. Yet this force persists only as long as expansion continues. Repeated cycles of expansion and contraction, possibly due to frequent disturbance, may be necessary to maintain persistent selection for cooperation by this mechanism.

Supplemental Information

Supplemental Information includes a heuristic description of the model and derivation of analytical results, four figures, and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.cub.2013.04.026.

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