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Genomic Signatures of Cooperation and Conflict in the Social Amoeba

Highlights

- Molecular evolution analyses reveal the history of social conflict
- Genes that mediate social conflict show signatures of frequency-dependent selection
- Balanced polymorphisms suggest that cheating may be stable and endemic

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In Brief

Cooperative systems are susceptible to cheating, but little is known about whether cheaters are successful in the long term. Ostrowski et al. examine the evolution of genes that mediate social cheating in *Dictyostelium discoideum* and find patterns of balancing selection, suggesting that cheating may be stable and endemic in natural populations.

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Genomic Signatures of Cooperation and Conflict in the Social Amoeba

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SUMMARY

Cooperative systems are susceptible to invasion by selfish individuals that profit from receiving the social benefits but fail to contribute. These so-called “cheaters” can have a fitness advantage in the laboratory, but it is unclear whether cheating provides an important selective advantage in nature. We used a population genomic approach to examine the history of genes involved in cheating behaviors in the social amoeba *Dictyostelium discoideum*, testing whether these genes experience rapid evolutionary change as a result of conflict over spore-stalk fate. Candidate genes and surrounding regions showed elevated polymorphism, unusual patterns of linkage disequilibrium, and lower levels of population differentiation, but they did not show greater between-species divergence. The signatures were most consistent with frequency-dependent selection acting to maintain multiple alleles, suggesting that conflict may lead to stalemate rather than an escalating arms race. Our results reveal the evolutionary dynamics of cooperation and cheating and underscore how sequence-based approaches can be used to elucidate the history of conflicts that are difficult to observe directly.

RESULTS

The social amoeba *Dictyostelium discoideum* is a model system for cooperation and conflict [1, 2]. Upon starvation, up to hundreds of thousands of amoebae converge, forming a multicellular slug and eventually a fruiting body. Some cells form the stalk of the fruiting body and die, while others rise to the top, form resistant spores, and disperse. Stalk formation is altruistic

because death of the stalk cells enhances the survival and dispersal of the spores. However, because fruiting bodies can contain multiple, genetically different clones, selection can favor cheaters—individuals that avoid forming the stalk themselves, yet benefit from its production by others [2].

Consistent with the prediction of cheating, cheaters can be readily found in natural populations of *D. discoideum* [1–3]. However, whether individuals that cheat are evolutionarily successful is unclear, and several hypotheses have been proposed. One hypothesis is that cheating selects for resistance, and resistance in turn selects for greater cheating. Social conflict could thus drive an escalating arms race of adaptations and counter-adaptations, reminiscent of the arms races between hosts and pathogens or predators and prey (Figure 1A) [4–7]. An alternative possibility is that cheaters have a selective advantage only when rare. For example, as cheaters increase in frequency in a population, they potentially displace the very victims on which they depend, or face other trade-offs [8]. This negative frequency dependence predicts that cheaters and cooperators can be maintained as a balanced polymorphism, effectively leading to a stalemate (Figure 1B) [9, 10]. Finally, some have suggested that there is no selective advantage to cheating [11]. Cheating might be selected against if relatedness among the strains in a fruiting body is high, such that cheaters primarily cheat their own relatives [12]. In this case, cheating might persist in populations as a “cheating load,” analogous to a genetic load for deleterious mutations (Figure 1C). Alternatively, cheating might also not be favored if the multicellular stage occurs only rarely in nature [13], such that there is little selection for or against these phenotypes.

Crucially, these different hypotheses about the long-term success of social cheating make unique, testable predictions about variation in the genes that mediate these conflicts (Table 1). To distinguish among these different possibilities, we took advantage of a previous screen that identified over 150 loci in *D. discoideum* that impact cheating behaviors [5]. We used whole-genome sequencing and molecular evolution to ask whether genes that mediate cheating behaviors show

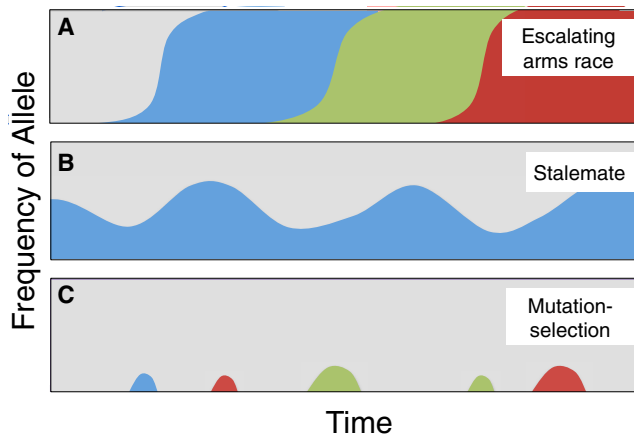


Figure 1. Example Scenarios for the Evolutionary Dynamics of Cheating Behaviors

Shaded areas are proportional to the frequencies of different alleles (colors) in a population.

(A) Escalating arms races, where epidemics of cheating and/or resistance sweep through populations.

(B) Stalemates, where cheaters invade but neither fix nor become lost from the population, resulting in endemic cheating.

(C) Mutation-selection balance, where new mutations that produce cheating behaviors are continually introduced into a population, but selection removes them.

distinctive signatures of molecular evolution that differ from the rest of the genome, distinguishing among the different hypotheses described in Table 1.

Candidate Genes Show Elevated Polymorphism

The different evolutionary scenarios for cheating alleles make unique predictions about the levels of polymorphism versus divergence (Table 1). For example, an escalating arms race driven by repeated selective sweeps of cheating alleles should reduce variation within species while elevating the sequence divergence between species, whereas the stalemate model

makes the opposite prediction. To test these possibilities, we first examined levels of polymorphism in regions surrounding candidate genes, comparing these values to a null hypothesis based on other regions of the genome. We observed higher polymorphism in candidate genes as a group compared to randomly chosen regions, which was significant for both mean and median levels at 20 kb (Figure 2). Compared to other genes, sequence variation was also disproportionately non-synonymous (higher p_N/p_S ; Table S1). Higher levels of polymorphism might occur if genes important for cheating behaviors show limited expression (e.g., if they are expressed in an infrequent portion of the life cycle [13]), but analysis of the published transcriptome of the laboratory strain [15] indicated no difference in their timing or levels of expression compared to other genes in the genome (Table S2).

No Evidence of Elevated Sequence Divergence between Species

The arms race hypothesis also predicts elevated rates of divergence between species, so we compared *D. discoideum* to its sister species *D. citrinum* at all identified orthologs. These analyses revealed lower rates of non-synonymous to synonymous substitution (d_N/d_S) for candidate genes compared to other genes in the genome (Table S1), which was inconsistent with the predictions of an arms race (Table 1). We reached a similar conclusion using McDonald-Kreitman tests [16], which compare P_N/P_S to D_N/D_S for each gene individually. Given very low levels of sequence polymorphism within *D. discoideum*, there was little resolution to detect significant deviations in this ratio for each locus individually. Nevertheless, two candidate genes (DDB_G0285541 and *chtC*) had strongly significant McDonald-Kreitman tests that were also in the extreme tail of the genome-wide distribution. These genes showed elevations in D_N/D_S relative to P_N/P_S , indicative of directional selection driving sequence divergence. In the remainder of candidate genes (that is, removing these two genes), the ratio was strongly significant and opposite in direction, indicating an excess of non-synonymous polymorphism to non-synonymous divergence compared to other genes (Figure 3). Taken together, genes mediating

Table 1. Predicted Sequence Patterns for Cooperation and Cheating Genes under Different Evolutionary Scenarios

Scenario	Description	Within-Species Polymorphism	Between-Species Divergence	Additional Signatures
Escalating arms race (directional selection)	repeated selective sweeps of cheating alleles through populations remove variation within populations and drive rapid divergence between species	decreased	increased	elevated population structure (higher F_{ST}), excess of high-frequency derived alleles (negative Fay and Wu's H)
Stalemate (balancing selection)	negative frequency dependence maintains both cheaters and cooperators within populations	increased	decreased	reduced population structure (lower F_{ST}), excess of intermediate frequency alleles (positive Tajima's D), elevated linkage disequilibrium (lower C), haplotype structure (higher Wall's B and Wall's Q)
Relaxed selection	cheating behaviors do not experience strong selection, possibly because the multicellular (social) stage is rare in nature	increased	increased	allele frequency skew closer to zero (Tajima's $D = 0$)
Purifying selection	cheaters are selected against, for example, if relatedness is high	decreased	decreased	excess of low frequency alleles (negative Tajima's D)

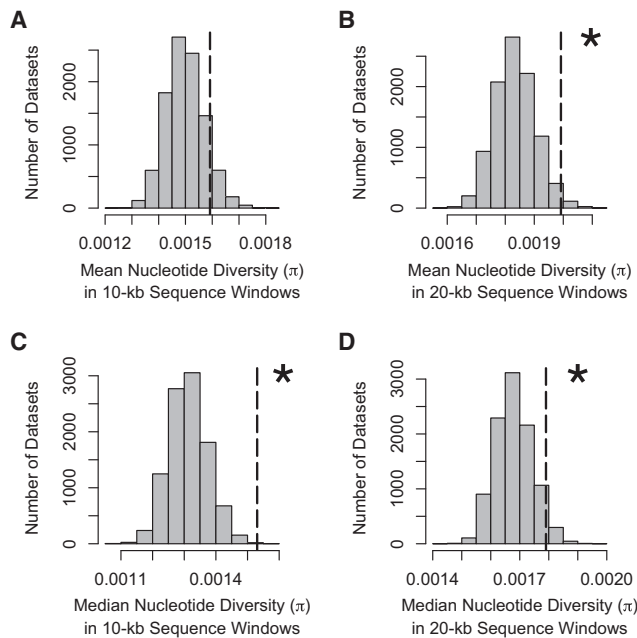


Figure 2. Comparison of Polymorphism in Regions Surrounding Candidate Loci Compared to the Rest of the Genome

Dotted lines show the mean or median nucleotide diversity in sequence windows of 10 kb or 20 kb compared to the null distribution based on 10,000 data sets of the same size where sequence windows were chosen randomly. Asterisks indicate statistically significant results.

(A) Mean nucleotide diversity in 10-kb sequence windows.

(B) Mean nucleotide diversity in 20-kb sequence windows.

(C) Median nucleotide diversity in 10-kb sequence windows.

(D) Median nucleotide diversity in 20-kb sequence windows.

social cheating did not show the elevated rates of amino acid substitution predicted under an escalating arms race or relaxed selection scenario.

Additional Signatures of Selection Support Stalemates

At first glance, elevated non-synonymous polymorphism, combined with low amino acid divergence, is potentially consistent with the stalemate model of frequency-dependent selection, where novel alleles can invade and establish in populations but ultimately fail to take over. Several additional tests support this interpretation. First, scaled to gene length, we observed significantly fewer haplotypes than expected and high levels of intra-genic linkage disequilibrium (lower C ; Figure S1), indicating divergent alleles, a signature of balancing selection. In addition, two metrics of balancing selection, Wall's B and Wall's Q [17], were significantly elevated in sequence windows surrounding candidate loci (Table S3).

A common test for balancing selection is to examine the distribution of allele frequencies. Whereas positive or purifying selection produces a strongly skewed distribution, balancing selection can maintain multiple alleles at intermediate frequencies. Surprisingly, given our results above supporting balancing selection, candidate loci showed greater skew, indicated by more negative values for two metrics of the site frequency spectrum (Table S4). Candidate genes as a

group also showed a significant excess of high-frequency derived alleles (Fay and Wu's H : -0.002 ; $p = 0.03$), which can indicate incomplete selective sweeps. The significant, negative Fay and Wu's H test for candidate genes suggests that variants rise to high frequency quickly and that these genes experience stronger-than-expected selection for sequence changes.

Finally, we calculated the index of population structure (F_{ST}) at each segregating site in the genome, comparing SNPs in close proximity to candidate genes against the rest of the genome. Unusually high F_{ST} can indicate that different alleles predominate in different geographic locations (in this case, Texas and Virginia) and can be caused by geographically restricted selective sweeps. On the other hand, unusually low F_{ST} values indicate alleles attain similar frequencies across geographically distinct populations, with each subpopulation maintaining multiple divergent alleles—a signature of negative frequency-dependent selection [18]. These results revealed lower-than-expected F_{ST} at candidate compared to non-candidate loci (Figure S2). The elevation in polymorphism in these genes, combined with significant reductions in population structure, argues against divergent alleles caused by local sweeps and suggests a role for negative frequency-dependent selection, with selection maintaining the same variants across subpopulations.

Evidence for Multiple Modes of Selection

By examining the molecular evolution patterns of candidate genes as a group, we could determine whether genes that mediate social conflict have general, recognizable patterns indicating unique forms of selection. Analyzing these genes as a group also provided greater sensitivity, which was helpful given the low levels of polymorphism in this species. However, these results reflect only average differences between candidate and non-candidate genes, and in fact, a diversity of dynamics is possible. Moreover, for many metrics, candidate genes might be extreme in opposite directions, such that we may fail to observe a strong signal of selection because these effects average out. We addressed this possibility in two ways. First, for each evolutionary metric, we asked whether the variance was higher for candidate genes compared to groups where genes are chosen randomly, indicating extremes in opposite directions. However, we did not observe elevated variance for any metric (Table S5). We also asked whether there was overrepresentation of candidate genes in both tails of the genome-wide distribution for each metric, but we observed no such cases (Table S6). While there was no overrepresentation in either tail of the genome-wide distribution for the McDonald-Kreitman test, two genes (DDB_G0285541 and *chtC*) showed extreme signatures of positive (directional) selection. The remaining genes showed the opposite pattern, an average excess of within-species non-synonymous polymorphism compared to non-synonymous divergence. Thus, while the average signature in conflict-related genes was elevated polymorphism and other patterns suggesting the selective maintenance of multiple alleles, other signatures were evident as well, including signatures of strong directional selection that could reflect escalating arms races at these particular loci.

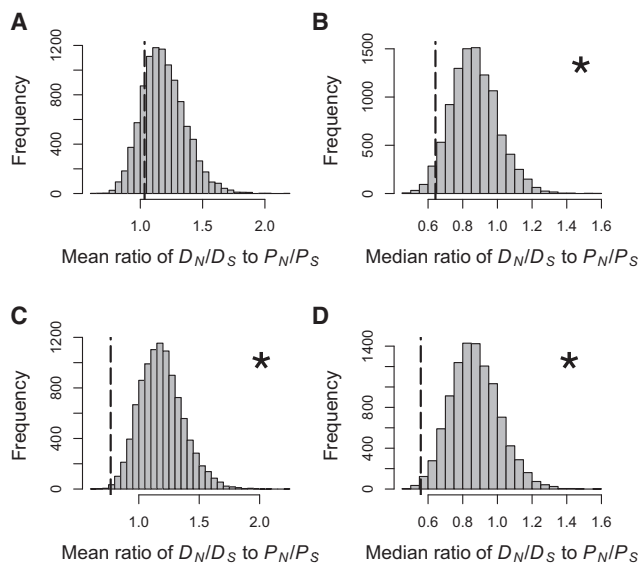


Figure 3. McDonald-Kreitman Tests

Histograms show the mean or median ratio of D_N/D_S to P_N/P_S for 10,000 randomly generated gene sets, and the dotted line shows the observed value of this ratio for candidate genes. Asterisks indicate statistically significant results.

(A) Mean ratio.

(B) Median ratio.

(C and D) Mean and median ratios after removing two candidate genes with extreme values.

See text for details.

DISCUSSION

In *D. discoideum*, a model system for the study of social conflict, it has long been noted that cheating behaviors are present among natural isolates [2, 3], but there is little understanding of why selfish behaviors arise, whether they persist, and whether the prevalence of cheaters in natural populations signifies long-term evolutionary success of this social strategy. We have indications of three main signatures of balancing selection on these genes as a class compared to other genes. First, they show higher levels of polymorphism, as expected when at least one SNP is under balancing selection and increases diversity at linked neutral loci. However, they did not show the concomitant elevation in amino acid substitution expected if polymorphism simply reflected relaxed selection. Second, they showed lower F_{ST} values, which can occur if selection is maintaining the same balanced polymorphisms across geographically different subpopulations. Third, other indicators of the deeper coalescence time characteristic of balancing selection are the significantly fewer haplotypes and higher linkage disequilibrium values, including significantly elevated Wall's B and Wall's Q .

The observation of elevated levels of polymorphism surrounding genes implicated in social cheating combined with other signatures of recent selection argue that *D. discoideum* experiences ongoing selection at these loci and is consistent with frequency-dependent selection allowing multiple types (or alleles) to coexist. Notably, we failed to observe the molecular signatures of a simple arms race—these dynamics are

expected to reduce genetic diversity and drive long-term sequence divergence between populations and/or species. Rather, our results are reminiscent of “trench warfare,” an alternative arms race scenario where alternative alleles do not rapidly displace one another, resulting in a prolonged stalemate [19]. Negative frequency dependence is also a hallmark of Red Queen dynamics, a form of coevolutionary arms race where alleles continually cycle but rarely fix [20].

The finding of balancing selection is also consistent with evolutionary theory about the role of frequency dependence in social interactions [21]. Many social behaviors are inherently frequency dependent, where the fitness of a given strategy (e.g., cheat or cooperate) is dependent on whether an individual's social partners employ the same strategy or not. Experimental studies of bacterial mutants that exhibit cheating behaviors suggest frequency dependence might be common [22–26]. In *Dictyostelium*, frequency-dependent fitness was shown for the *fbxA*[−] strain, a mutant that allocates fewer cells to the stalk but produces disproportionately many spores when co-developed with another strain, which it cheats [12].

Like social conflict, conflicts between the sexes over optimal levels of mating and between parents and offspring over optimal provisioning are also hypothesized to result in antagonistic coevolution [20, 27–29]. While some studies have shown directional selection on genes underlying these other forms of intraspecific conflict, many others have found signatures of balancing selection, diversifying selection, or a combination of both [30–33]. Thus, taken together, our results not only indicate stalemate as a possible outcome of social conflict but add to a growing body of evidence that stalemates may be a common outcome in conflict-driven systems more generally. While identification of these polymorphisms should open the door to investigation into the functional consequences of this variation for cheating and resistance behaviors, the population genomic approach used here provides insight into the long-term consequences of social conflict and highlights the possibility of an ongoing, dynamic interaction at these loci.

EXPERIMENTAL PROCEDURES

Strains

We re-sequenced the genomes of 20 natural isolates of *D. discoideum*, primarily from two locations: Houston, Texas (six strains) and Mountain Lake, Virginia (nine strains), as well as five additional strains from different geographic locations (two sites in Texas, and one site in each of Massachusetts, Kentucky, and Illinois) using 454 or Illumina. Sequencing reads were aligned to the AX4 reference genome (Assembly/GFF3 file generated June 9, 2010, available at <http://www.dictybase.org>) using MAQ for Illumina sequencing reads and ATLAS-SNP for 454 data. Detailed mapping and SNP calling procedures are available in the [Supplemental Information](#).

Molecular Evolution Analyses

Nucleotide diversity, Tajima's D , Fu and Li's D^* , Hudson's C (recombination, or ρ), haplotype diversity, Fay and Wu's H , and haplotype number were determined for all genes in the genome and in sequence windows using “compute” (available at <http://www.molpopgen.org>). Levels of non-synonymous (p_N) and synonymous (p_S) diversity were calculated using the program “gestimator,” and the McDonald-Kreitman tests were obtained using “MKtest” (both available at <http://www.molpopgen.org>). F_{ST} was calculated for all segregating sites using scripts written in Ruby and Python. Resampling analyses were performed using R. More details are available in the [Supplemental Information](#).

ACCESSION NUMBERS

The accession number for the *D. discoideum* sequencing reads reported in this paper is NCBI BioProject: PRJNA45873. The accession number for the *D. citrinum* assembly is GenBank: AJWG00000000.1.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, two figures, and six tables and can be found with this article online at <http://dx.doi.org/10.1016/j.cub.2015.04.059>.

AUTHOR CONTRIBUTIONS

E.A.O., C.D., M.K.-K., and D.A.B. handled cell culture and DNA extraction. E.A.O., J.E.S., D.C.Q., S.R., K.C.W., R.A.G., and R.S. conceived and designed the experiments. F.L.-G., S.L.L., C.L.K., H.H.D., V.K., L.J., S.P., Y.H., L.C., D.M.M., S.R., R.A.G., and K.C.W. carried out library preparation and sequencing. E.A.O., Y.S., X.T., R.S., and S.R. analyzed the data. J.Q., H.J., and K.C.W. carried out assembly and annotation of *D. citrinum*. E.A.O. and X.T. handled analysis of *D. citrinum* and *D. discoideum*. E.A.O., J.E.S., A.K., and D.C.Q. wrote the paper.

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