

The computational complexity of ecological and evolutionary spatial dynamics

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Abstract

There are deep, yet largely unexplored connections between computer science and biology. Both disciplines examine how information proliferates in time and space. Central results in computer science describe the complexity of algorithms that solve certain classes of problems. An algorithm is deemed efficient if it can solve a problem in polynomial time, which means the running time of the algorithm is a polynomial function of the length of the input. There are classes of harder problems for which the fastest possible algorithm requires exponential time. Another criterion is the space requirement of the algorithm. There is a crucial distinction between algorithms that can find a solution, verify a solution, or list several distinct solutions in given time and space. The complexity hierarchy that is generated in this way is the foundation of theoretical computer science. Precise complexity results can be notoriously difficult. The famous P=NP question is one of the hardest open problems in computer science and all of mathematics. Here we consider simple processes of ecological and evolutionary spatial dynamics. The basic question is: what is the probability that a new invader (or a new mutant) takes over a resident population? We derive precise complexity results for a variety of scenarios. We therefore show that some fundamental questions in this area cannot be answered by simple equations.

1 Significance

There is a deep connection between computer science and biology, as both fields study how information proliferates in time and space. In computer science, the space and time requirements of algorithms to solve certain problems generate complexity classes, which represent the foundation of theoretical computer science. The theory of evolution in structured population has provided an impressive range of results, but an understanding of the computational complexity of even simple questions is still missing. In this work we prove – unexpectedly – that some fundamental problems in ecological and evolutionary spatial dynamics can be precisely characterized by well-established complexity classes of the theory of computation. Since we show computational hardness for several basic problems, our results imply that the corresponding questions cannot be answered by simple equations. For example, there cannot be a simple formula for the fixation probability of a new mutant given frequency dependent selection in a structured population. We also show that some problems, such as calculating the molecular clock of neutral evolution in structured populations, admit efficient algorithmic solutions.

Evolutionary games on graphs — Fixation probability — Complexity classes

2 Introduction

Evolution occurs in populations of reproducing individuals. Mutation generates distinct types. Selection favors some types over others. The mathematical formalism of evolution describes how populations change in their genetic (or phenotypic) composition over time. Many papers study evolutionary dynamics in structured

Table 1: Complexity results for various models and computational questions

	Qualitative	Quantitative
Ecological Scenario	NP-complete	#P-complete
Linear fitness	PSPACE-complete	PSPACE-complete
Exponential fitness	P	PSPACE-complete

populations [1, 2, 3, 4, 5, 6, 7, 8]. Spatial structure can affect the rate of neutral evolution [9]. There are results that describe which spatial structures do or do not affect the outcome of constant selection [10, 11, 12]. Constant selection refers to a situation where the competing types have constant reproductive rates independent of the composition of the population. Some population structures can be amplifiers or suppressors of constant selection [13, 6, 14] meaning that they modify the intensity of selective differences. A large literature deals with evolutionary games [15, 16, 17, 18, 19] in structured populations [1, 20, 21, 22, 23, 24, 25, 26, 27, 28]. In evolutionary games the reproductive success of an individual depends on interactions with others. Many population structures and update rules can affect the outcome of evolutionary games. For example, spatial structure can favor evolution of cooperation [1, 29].

In this paper we are interested in stochastic evolutionary dynamics in populations of finite size. A typical setting is provided by evolutionary graph theory [6, 30, 31, 32, 33, 34]. The individuals of a population occupy the vertices of a graph. The links determine who interacts with whom for receiving payoff and for reproduction. There can be a single graph for game dynamical interaction and evolutionary replacement, or the interaction and replacement graphs can be distinct [35]. Often the graph is held constant during evolutionary updating, but it is also possible to study dynamically changing graphs [36, 37, 38, 39, 40, 41, 42, 43, 44].

The study of spatial dynamics also has a long tradition in ecology [45, 46, 47, 48, 49]. Here the typical setting is that different species compete for ecological niches. Many evolutionary models are formally equivalent to ecological ones - especially if we consider only selection and not mutation. Then we can interpret the different types as individuals of different species.

This paper is structured as follows. First we give an intuitive account of the foundation of theoretical computer science. We describe classes of problems that can be solved by algorithms in certain time and space constraints. Subsequently we present two simple problems of evolutionary dynamics in spatial settings. The first problem is motivated by a very simple ecological dynamic; the second problem is the general setting of evolutionary games on graphs. In both cases, the basic question is to calculate the take over probability (or fixation probability) of a new type. That is we introduce a new type in a random position in the population and we ask what is the complexity of an algorithm that can characterize the probability that the new type takes over the population (becomes fixed). Unexpectedly we are able to prove exact complexity results (see Table 1).

The class PTIME (denoted as P) consists of problems whose solutions can be computed by an algorithm that uses polynomial time. This means the running time of the algorithm grows as a polynomial function of the size of the input. In computer science, PTIME represents the class of problems which can be solved efficiently.

The class NP (non-deterministic polynomial time) consists of problems, for which solutions exist that are of polynomial length, and given a candidate for a solution of polynomial length, whether the candidate is indeed a solution can be checked in polynomial time. Therefore, an NP algorithm can verify a solution in polynomial time.

In order to proceed further, we need the notion of ‘reduction’ between classes of problems. A reduction, from a given problem P_1 to a problem P_2 , is a translation such that a solution for P_2 can provide a solution for P_1 . More precisely, if there is a polynomial-time reduction from P_1 to P_2 , then a polynomial-time algorithm for P_2 implies a polynomial-time algorithm for P_1 .

A given problem is NP-hard if for every problem in NP there is a polynomial reduction to the given problem. A problem is NP-complete, if it is both NP-hard, as well as there is an NP algorithm for the problem.

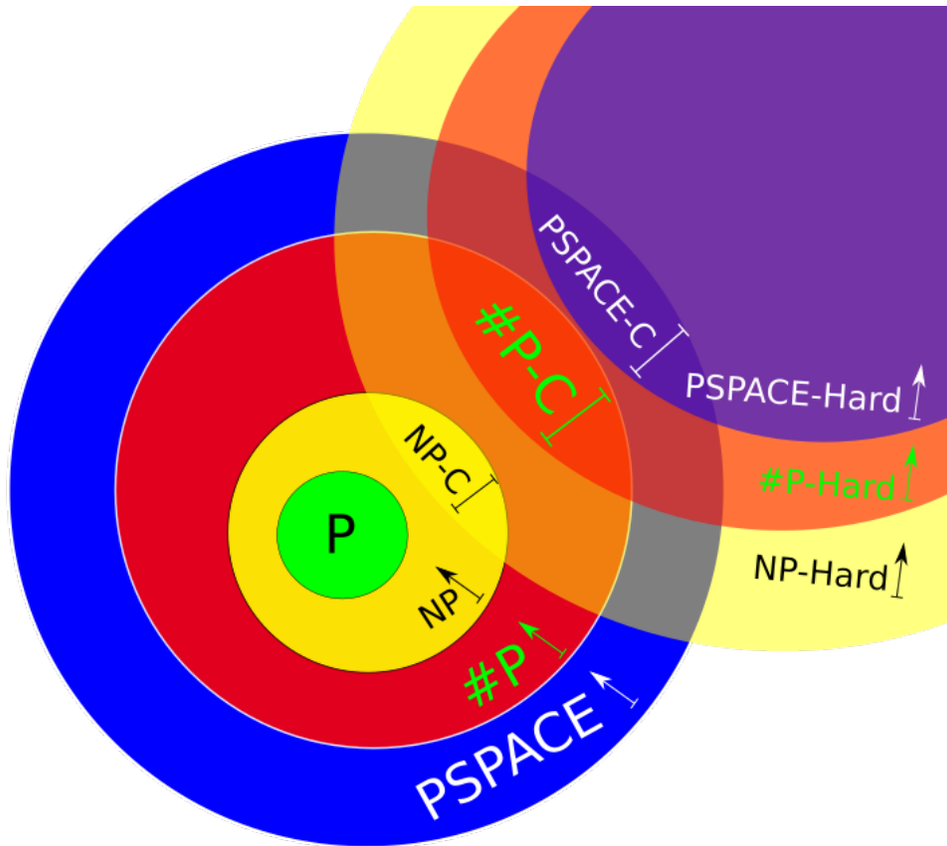


Figure 1: A pictorial illustration of the complexity classes P , NP , $\#P$, and $PSPACE$. The complexity class P is contained in NP , NP is contained in $\#P$, and $\#P$ is contained in $PSPACE$. The widely believed conjecture is that the complexity classes are different. A problem is NP -hard if it is at least as hard as each problem in NP ; and similar for $\#P$ -hardness and $PSPACE$ -hardness. The intersection of NP and NP -hard gives the NP -complete problems, and similarly for $\#P$ -complete and $PSPACE$ -complete problems. Hence a polynomial-time solution for a NP -hard or NP -complete problem would imply $P=NP$.

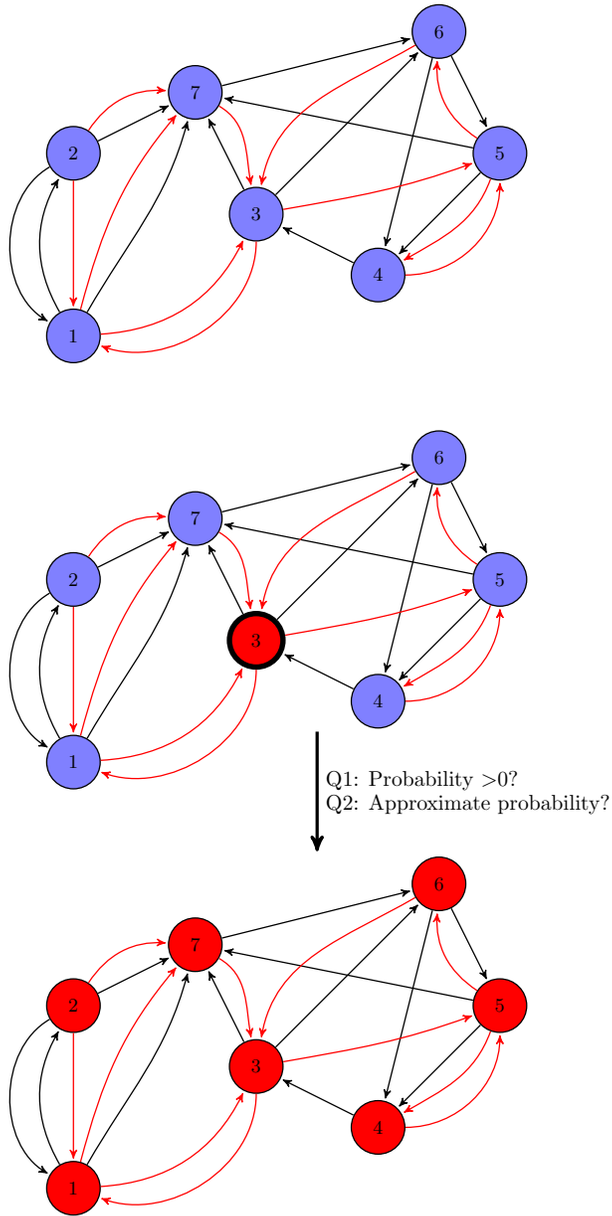


Figure 2: Illustration of mutant introduction. The residents (type A) are colored blue and the mutants (type B) are colored red. The black edges are the edges of the interaction graph and the red are the edges of the reproduction graph. The probability to introduce a mutant in a specific vertex is always one over the number of vertices. The computational questions of interest regarding the take over probability are as follows: whether the probability is positive (qualitative question), and compute an approximation of the probability (quantitative question).

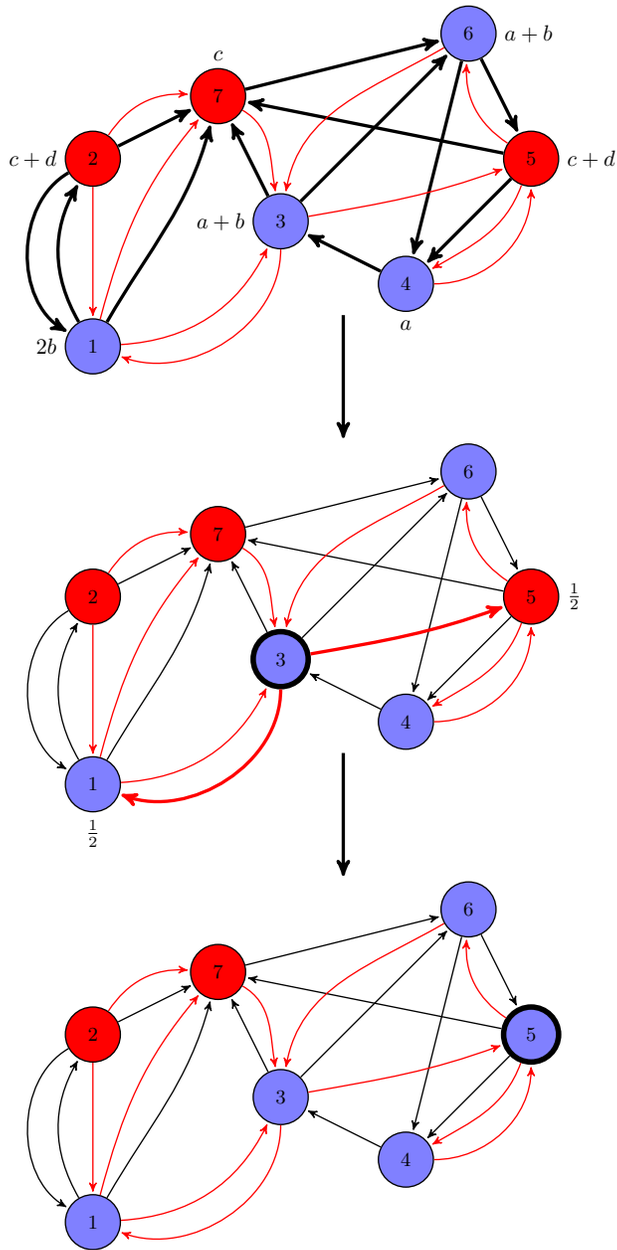


Figure 3: Illustration of reproduction with matrix $B \begin{pmatrix} a & b \\ c & d \end{pmatrix}$. The residents (type A) are colored blue and the mutants (type B) are colored red. The black edges are the edges of the interaction graph and the red are the edges of the reproduction graph. In the first figure beside each vertex the payoff of the vertex (which is the sum of the payoff of the interactions) is shown. Since the first figure shows the payoff computation, the interaction edges that are responsible for payoff calculation are boldfaced. In the second figure the vertex labeled 3 is selected for reproduction. The reproduction edges from vertex 3 are boldfaced, and each edge has probability $1/2$. Finally, the successor 5 is chosen for replacement, i.e., vertex 3 reproduces to vertex 5.

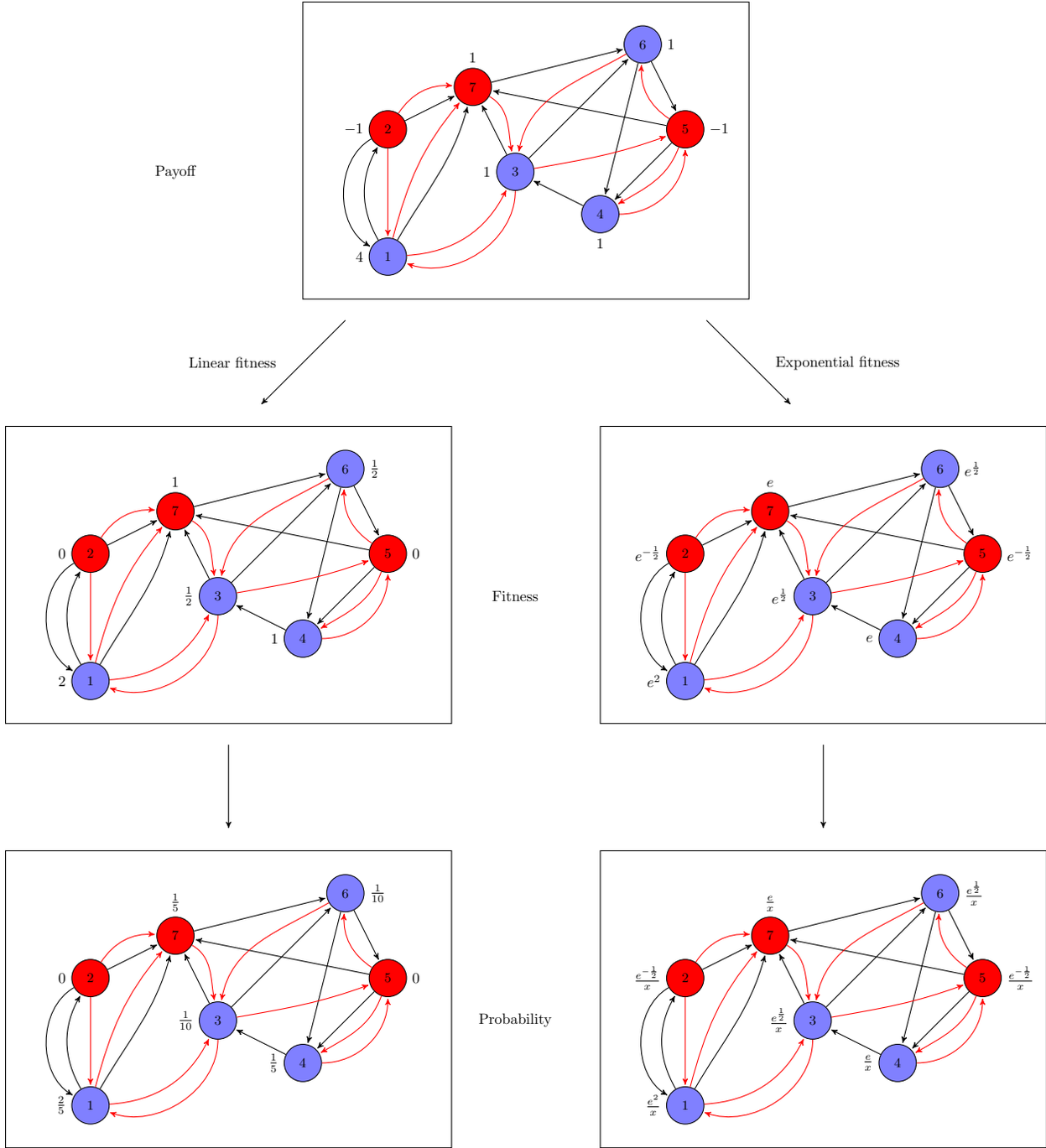


Figure 4: Illustration of different payoffs to fitness with $\begin{matrix} A & B \\ \begin{pmatrix} -1 & 2 \\ 1 & -2 \end{pmatrix} \end{matrix}$. The residents (type A) are blue and the mutants (type B) red. The black edges are the edges of the interaction graph and the red are the edges of the reproduction graph. In the figure of the first row we show the payoff for every vertex. In the next row we show the fitness which is either a linear function of the payoff but at least 0; or an exponential function of the payoff. Finally, in the third row, with each vertex we show the probability, which is the normalized fitness, that the vertex is selected for reproduction (in the last figure, the number x is the sum of the fitness, i.e., $x = e^2 + 2e + 2e^{\frac{1}{2}} + 2e^{-\frac{1}{2}}$).

For example, consider a Boolean formula over variables, and the question whether there exists an assignment to the variables such that the formula is true. A polynomial candidate solution is an assignment of truth values to variables, and given a candidate assignment the formula can be evaluated in polynomial time. This is the famous satisfiability, SAT, problem in computer science. The SAT problem is NP-complete.

The class P is contained in NP, and a major, long-standing open question in computer science is whether $P=NP$? A polynomial-time algorithm for an NP-complete (or an NP-hard) problem would imply that $P=NP$, resolving the long-standing open problem.

The class #P (sharpP) intuitively corresponds to counting the number of solutions. A problem is in #P if it counts the number of distinct solutions such that (i) every possible candidate for a solution is of polynomial length, and (ii) given a candidate for a solution, it can be checked in polynomial time whether the candidate is a solution. For example, given a Boolean formula, the problem whether there are at least k distinct satisfying assignments to the formula is a #P-problem. A given problem is #P-hard, if for every #P-problem there is a polynomial-time reduction to the given problem. A #P-complete problem is a problem that is both #P-hard, and there is a #P-solution. For example, counting the number of solutions in SAT is #P-complete.

The class NP is contained in #P because given the enumeration of solutions for #P, it is easy to check if there exists at least one solution. Intuitively, an NP problem asks whether there is at least one solution, whereas #P is the counting version which asks if there are least k distinct solutions (and the special case of $k = 1$ gives NP). Again a major open question is whether $NP=#P$? Note that a polynomial-time algorithm for a #P-complete problem would be an even bigger result as it would imply both $P=NP$ and $P=#P$.

The class PSPACE consists of problems which can be solved with polynomial space. Note that a polynomial space algorithm can reuse space and can in general require exponential time. Every #P problem can be solved in PSPACE by simply enumerating each candidate for a solution and checking if it is a solution. Since we can reuse space to enumerate the candidates for solutions, the enumeration can be achieved in polynomial space. Moreover, every polynomial-time algorithm uses at most polynomial space. Hence it follows that #P is contained in PSPACE. The notion of PSPACE-hardness and PSPACE-completeness is similar to the notion of NP-hardness and NP-completeness, but with respect to the problems in PSPACE. Again a long-standing open question in computer science is whether $#P=PSPACE$, and a polynomial-time algorithm for a PSPACE-complete (or PSPACE-hard) problem would imply $P=NP=#P=PSPACE$.

We have mentioned that the major questions about the equality of the complexity classes are open problems, but the widely believed conjecture is that P is strictly contained in NP, NP is strictly contained in #P, and #P is strictly contained in PSPACE. In other words, it is widely believed that NP-complete problems cannot be solved in polynomial time, #P-complete problems are harder than NP-complete problems, and PSPACE-complete problems are harder than #P-complete problems. A pictorial illustration of the complexity classes is shown in Figure 1.

3 Results

The first problem is motivated by ecological dynamics. There is an ecosystem occupied by resident species. The spatial structure of the ecosystem is given by a graph. An invading species is introduced (see Figure 2 for an illustration). We assume the invading species has a competitive advantage in the sense that once a position is occupied by the invading species the resident cannot get it back. The invading species, however, has a density constraint: if the number of invaders around a focal invader is above a threshold, h , then the invader in the focal vertex can not colonize another vertex.

We are interested in the probability that the invader starting from a random initial position will take over the entire ecosystem (and therefore drive the resident to extinction). There are two types of questions. The ‘qualitative question’ is whether the take over probability is greater than zero. The ‘quantitative question’ is concerned with computing the take over probability subject to a small error. Figure 2 gives a pictorial illustration. We prove the following results. The qualitative question is NP-complete (See SI Appendix Theorem 4). The quantitative question is #P-complete (See SI Appendix Theorem 8).

The second problem is concerned with evolutionary games in structured populations. There are two

types, A and B , whose reproductive rates depend on local interactions. We consider the setting of games on graphs. Each vertex is occupied by one individual, which is either A or B . Interactions occur pairwise with all neighbors. The payoff matrix is given by

$$\begin{array}{cc} & \begin{array}{cc} A & B \end{array} \\ \begin{array}{c} A \\ B \end{array} & \begin{pmatrix} a & b \\ c & d \end{pmatrix} \end{array} \quad (1)$$

The entries of the payoff matrix can be positive or negative (or zero). Each individual interacts with all of its neighbors on the graph to derive a payoff sum. The payoff sum is translated into reproductive success as follows. If the payoff sum is positive, then the fecundity equals the payoff sum. If the payoff sum is negative, then the fecundity is zero. We refer to this translation as linear fitness. In any one time step, a random individual is chosen for reproduction proportional to its fecundity. The offspring, which is of the same type as the parent, is placed into an adjacent position on the graph (see Figure 3 and Figure 4 for an illustration).

We are interested in the probability that a single A individual starting in a random position on the graph generates a lineage which will take over the entire population; this probability is generally called fixation probability. As before, there are two types of questions. The ‘qualitative question’ is whether the fixation probability is positive. The ‘quantitative question’ is concerned with computing the fixation probability subject to a small error. We prove the following results. The qualitative question is NP-hard and in PSPACE. The quantitative question is #P-hard and in PSPACE. The results follow from SI Appendix Theorem 4, Theorem 8, and Theorem 15.

Note that the first problem can also be obtained as a special case of the second problem. In the payoff matrix (1) we can set, for example, $a = -1$, $b = 1$, $c = d = 0$. This ‘game’ has the property that type B never reproduces and type A reproduces until half its neighbors are also of type A . This parameter choice leads to the same qualitative behavior and the same complexity bounds as described in the first problem.

A generalization of games on graphs is the setting where the interaction graph and the replacement graph are distinct [35]. Thus each individual interacts with all of its neighbors on the interaction graph to receive payoff. Subsequently an individual is chosen for reproduction proportional to its fecundity. The offspring is placed randomly among all neighbors of the focal individual on the replacement graph. In this case, both the qualitative and quantitative questions become PSPACE-complete (See SI Appendix Theorem 15)

We also consider a variation of the second problem. In particular we change the mapping from payoff to fecundity. We now assume that fecundity is an exponential function of payoff, and refer to it as exponential fitness (see Figure 4 for an illustration). Therefore the fecundity of an individual is always positive (even if its payoff sum is negative). In this setting the qualitative question can be decided in polynomial time. The reason is that the fixation probability is positive if the graph is connected. Thus, in order to answer the qualitative question the algorithm only needs to check whether the graph is connected; this problem is in P. However, the quantitative question has the same complexity as the previous problem (See SI Appendix Theorem 16 and Theorem 17).

A very special case of games on graphs is constant selection. Type A has constant fecundity a and type B has constant fecundity b independent of any interactions, i.e., fecundity is independent of the population structure. The qualitative question concerning the fixation probability of A is in P. The quantitative question is in PSPACE, but any non-trivial lower bound is an open question.

Finally, while we establish computational hardness for several problems, we also show that two classic problems can be solved in polynomial time (See SI Appendix Section 7). First, we consider the molecular clock, which is the rate at which neutral mutations accumulate over time. The molecular clock is affected by population structure [35]. We show that the molecular clock can be computed in polynomial time because the problem reduces to solving a set of linear equalities, which can be achieved in polynomial time using Gaussian elimination. Second, we consider evolutionary games in a well-mixed population structure, where the underlying structure is the complete graph [51]. We show that the exact fixation probability can be computed in polynomial time. In this case the problem can be reduced to computing absorption probabilities in Markov chains, where each state represent the number of mutants. Hence the Markov chain is linear in the number of vertices of the graphs, and since absorption probabilities in Markov chains can be

computed in polynomial time (by solving a set of linear equalities) we obtain the desired result.

4 Methods: Proof Ideas

We now present the key intuition and main ideas of our results. The most interesting and technically insightful results are the lower bounds (i.e., the hardness proofs), and we present the key ideas only for them.

1. *NP-hardness of the qualitative ecological problem.* One of the most classic NP-complete problems is the 3SAT-problem, which is the SAT problem where every clause has exactly three literals (a literal is a Boolean variable x or a negation of a variable \bar{x}). Given instances of the 3SAT problem we construct instances of the ecological problem, where we have a start vertex where the mutant arises, followed by a sequence of vertices (i.e., each vertex can reproduce a mutant to the next), one for each clause. By our construction, a vertex in this sequence can reproduce at most three times, one of which must be the next vertex of the sequence, and the others correspond to at most two literals of the clause (intuitively these two literals represent the ones that are not set to true by a candidate satisfying assignment). The last vertex of the sequence reproduces to a new sequence of vertices that corresponds to an assignment of truth values to the variables. Each vertex in this new sequence can reproduce twice, one to the next vertex of the sequence, and other to a variable or its negation. The variables or the corresponding negation can then reproduce mutants to the corresponding literals of the clauses. After this sequence, all vertices that do not correspond to a literal in a clause become mutants. In essence our construction ensures if there is a satisfying assignment, then with positive probability all vertices can become mutants, and conversely, if there is no satisfying assignment, then the probability that all vertices become mutants is zero.
2. *#P-hardness of the quantitative ecological problem.* A #P-complete problem is counting the number of *perfect matchings* in a bipartite graph (which also corresponds to computing permanent of Boolean matrices). A bipartite graph consists of two set of vertices, a set on the left side and a set on the right side, with edges from the left side to right side. A perfect matching is a one-to-one mapping of each vertex of the left side to a vertex on the right side such that there is an edge between them. First we argue that for the hardness proof it suffices to consider bipartite graphs in which each vertex in the left side has outdegree 2^k for some integer k . A key idea in our construction is that in a full binary tree, if the root becomes a mutant, and every vertex can reproduce exactly once, then the set of mutants will eventually consists of a path from the root to a leaf, chosen uniformly at random. Our construction is then as follows: We have a start vertex where the mutant arises, which reproduces to turn each of the vertices in the left side of the bipartite graph to mutants. Each of the vertices in the left side is the root of a full binary tree, where the leaves correspond to the right side of the bipartite graph. We show that the fixation process corresponds to a perfect matching (defined from the path in the full binary trees), and given an approximation of the fixation probability, the exact number of perfect matchings of the bipartite graph can be computed.
3. *PSPACE-hardness for the game on evolutionary graph problem.* Our PSPACE-hardness proof shows that the evolutionary process can solve the following *concurrent-if problem*, which we show is PSPACE-hard. The concurrent-if problem consists of a set of Boolean variables x_1, x_2, \dots, x_n , with a given initial truth assignment to the variables, and a set of if-statements. Each if-statement s_i is of the following form: If a conjunctive clause C_i over the variables is true, then assign a truth value to a variable (e.g., if $(x_2 \wedge x_4 \wedge \bar{x}_5)$, then x_3 is assigned false). The problem is to decide whether the first variable (which is the accepting variable) eventually becomes true. We show that each variable can be represented as four vertices, and each if statement as a single vertex, in the evolutionary graph, and the evolutionary process can mimic the execution of the concurrent-if problem. Finally, if the accepting variable becomes true, then it corresponds to making a special vertex in the evolutionary graph as mutant. There exists a part of the evolutionary graph that can only become mutants after the special vertex has become a

mutant. Using this construction we show that for evolutionary games on graphs both the qualitative and quantitative problems are PSPACE-hard.

5 Discussion

In summary, we have established computational complexity results for some fundamental problems of ecological and evolutionary dynamics in structured populations. Our main results are summarized in Table 1. We now discuss the significance of our findings.

1. *Interdisciplinary connection.* While both computer science and biology examine the proliferation of information in time and space, the deep connection between them has been largely unexplored. Our work provides precise computational complexity results for several well-studied problems in biology and can be viewed as a step to establish a connection between the two disciplines.
2. *Well-studied open problem.* The problems we have considered are basic aspects of well-studied questions for ecological and evolutionary dynamics in structured populations [1, 5, 6, 14, 28, 29]. Several reviews have been written on this topic [7, 30, 33, 50]. We first discuss the significance of an algorithmic approach in evolutionary graph theory. An efficient algorithm, which considers all (even worst-case) graphs for evolutionary processes, is important for the following reasons: (i) It has been shown that some population structures (called amplifiers) can increase the effect of natural selection [6, 29]; but amplifiers are rare and constructing them is difficult [6, 29, 14, 52, 53, 33]. If there was an efficient algorithmic approach that worked for all graphs, then one could design candidates for amplifiers and efficiently check their fixation probabilities. Since there exists no algorithmic approach, research has to focus on special classes of graphs to identify simple formulas, such as calculating the fixation probabilities on star like graphs [52]. (ii) It is known that some population structures and evolutionary dynamics promote evolution of cooperation, but others do not [29]. An important open problem is to characterize the set of graphs that promote cooperation. An efficient algorithmic approach would be useful to check candidate structures. Since no efficient algorithm exists, one has to study special cases, for example by considering nearly regular graphs [29].

Thus a general algorithmic approach is a very important problem for the well-studied question concerning the effect of population structures on evolutionary dynamics. An algorithmic approach has been studied for important special cases such as for complete graphs [54], and NP-hardness was stated for the quantitative problem [6]. The review [50] identifies the complexity of computing fixation probabilities on evolutionary graphs as important open question in the area. In [50], two open problems (2.1 and 2.2) are identified that ask for the complexity of computing the exact fixation probabilities for graphs and for games on graphs. Our results not only present answers to those crucial questions, but also show that both the approximation problem and the qualitative question are computationally hard. The most interesting aspects of our results are the lower bounds, which show that in most cases there exists no efficient algorithm, under the widely believed conjecture that P is different from NP. A simple equation based solution would give an efficient algorithm, and thus our result formally shows that for evaluating the fixation probability in spatial settings there does not exist a simple equation based solution in general. Our results are significant for the following reasons: (a) it establishes the computational complexity for fundamental problems of ecological and evolutionary dynamics in structured populations considered in for example [1, 5, 6, 14, 28, 29, 7, 30, 33, 50]; and (b) it significantly improves the complexity result of [6] and solves the computational complexity questions of the area as identified in [50].

3. *Methodological insight.* Our proof ideas also reveal some important points. We show how evolutionary processes in structured populations can mimic aspects of computation. This insight could be useful for future research on understanding the computational complexities of other stochastic processes on population structures.

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