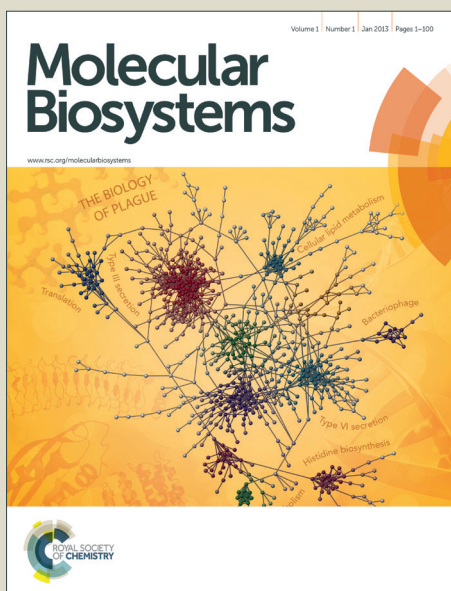


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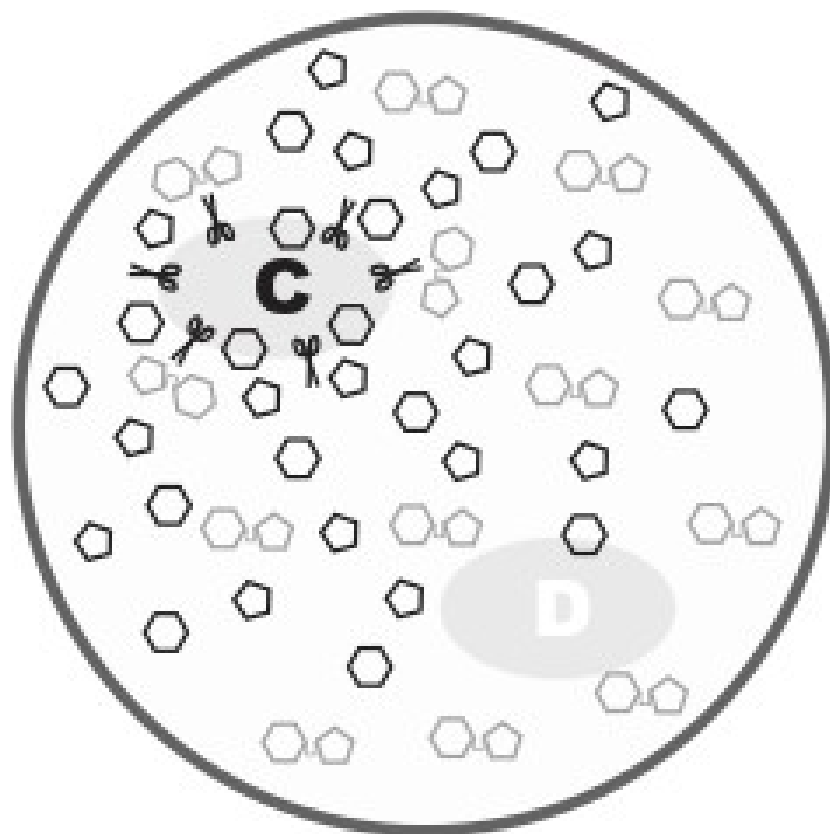
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The interaction of living cells can often be described by game theory, for example, in the case of secretion of extracellular enzymes. In this review we discuss numerous examples of applying game theory to cellular processes.

Evolutionary game theory: cells as players

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In two papers we review game theory applications in biology below the level of cognitive living beings. It can be seen that evolution and natural selection replace the rationality of the actors appropriately. Even in these micro worlds, competing situations and cooperative relationships can be found and modeled by evolutionary game theory. Also those units of the lowest levels of life show different strategies for different environmental situations or different partners. We give a wide overview of evolutionary game theory applications to microscopic units. In this first review situations on the cellular level are tackled. In particular metabolic problems are discussed, such as ATP-producing pathways, secretion of public goods and cross-feeding. Further topics are cyclic competition among more than two partners, intra- and inter-cellular signalling, the struggle between pathogens and the immune system, and the interactions of cancer cells. Moreover, we introduce the theoretical basics to encourage scientists to investigate problems in cell biology and molecular biology by evolutionary game theory.

1 Introduction

Replicating units such as living organisms practically never exist alone. Rather, they interact with each other in various ways, in cooperative or competing relationships. Not least, this applies to cells within multicellular organisms (where usually cooperative types of interaction prevail). Evolutionary theory often tries to understand heritable changes of organisms and cells in terms of better adaptation to a well-defined or even constant environment. This reasoning regarding optimization is, however, often insufficient for understanding biological evolution. It needs to be acknowledged that evolution is almost always co-evolution^{1,2}. That is, the properties of interacting macromolecules or cells evolve in dependence on each other. This may affect the optimum on all sides, and leads to an “open ended” dynamics of evolution rather than to the approximation to a fitness optimum³.

An appropriate framework for analysing evolution in dynamic fitness landscapes is evolutionary game theory because it allows one to describe the maximization of fitness of several interacting partners^{4–7}. Game theory had originally been developed mainly for applications in economics⁸, besides inspirations from parlour games such as poker. Later, it was also applied and adapted to applications in biology, resulting in the field of evolutionary game theory.

The metaphor of games is wide-spread in human society. Companies describe themselves as “global players”, the sentence “The game is over” occurs in many thriller movies, and the German word “Beispiel” (example) refers to “Spiel” (game). Especially, the “Games people play”⁹ have always been of strong interest. Thus, thinking in terms of games helps us to understand complex phenomena in an intuitive way, even if these phenomena are very serious things like conflicts among animals, companies or nations. The book “Game Theory” by Nobel prize winner Roger Myerson¹⁰ is thus aptly subtitled “Analysis of Conflict”.

Independently of evolutionary game theory, Eigen and Winkler¹¹ considered many biological processes including prebiotic evolution to have properties of games. In this and an accompanying review¹², however, we use the term “game” in the sense of classical game theory as introduced by Neumann and Morgenstern⁸ and developed further by Nash¹³ and Maynard-Smith⁴. A game describes the interaction of two or more agents (players) in which the outcome of each player depends not only on the own decision, but also on the strategies adopted by the coplayers.

Since the pioneering work by Maynard-Smith and Price¹⁴, Hamilton¹⁵ and others, evolutionary game theory has been

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applied for decades to the interaction among higher organisms on a macroscopic scale. Examples concern animal behaviour^{16–21}, competition among plants^{22,23} and interactions between animals and plants²⁴. Its application to the microscopic scale - the cellular and subcellular levels of life - is much less common, but of increasing interest and importance.

Not only multicellular organisms as a whole, but also cells and some macromolecules within cells can be considered as players in the sense of game theory. Many of their traits can then be regarded as strategies, such as usage of metabolic pathways, expression of virulence factors or generation of different splice variants. The change between strategies can occur by mutations or epigenetic modifications and the succeeding strategy is chosen by natural selection. Therefore, cognitive and rational capabilities are no prerequisites for being considered as a player in the sense of game theory. Merely the property of having a trait (strategy) that is subject to selection and, thus, influences the replicational success (the payoff of the game) makes them potential players.

In this and an accompanying review¹², we provide a broad overview of applications of evolutionary game theory in molecular and cell biology. While the accompanying review¹² focuses on the potential of the application of game theory to interactions on the subcellular level, the present one is devoted to interactions between cells. Examples include the formation of biofilms²⁵, division of labour in consortium pathways^{26,27}, intercellular signalling^{28,29}, and tumour biology^{30,31}. An outline of the mathematical fundamentals of game theory is given in the Appendix.

The sections of this review are ordered according to different biological applications rather than different theoretical game types. Thus, it is structured with respect to biological players rather than games. It is worth noting that one of the same game type such as the hawk-dove game occurs in different applications.

In our view, the potential of game theory is largely underestimated in molecular and cell biology. One intention of our two reviews is certainly to motivate more researchers to employ the powerful tools of game theory.

2 Respiration versus fermentation

During evolution of metabolic pathways, several alternatives have evolved. Examples are provided by glycolysis and the Entner-Doudoroff pathway, both of which degrade glucose to trioses. While most present-day organisms use glycolysis, the bacterium *Zymomonas mobilis* (used technologically for producing Tequila) relies on the Entner-Doudoroff pathway, and *Escherichia coli* and several other bacteria harbour both pathways. Alternatively or in addition, many organisms can use the pathway of respiration (oxidative phosphorylation). Path-

way usage can be changed by regulation even during the lifespan of one individual.

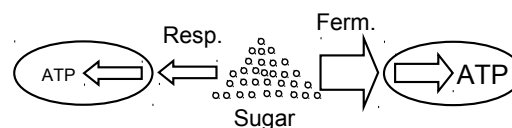


Fig. 1 Two cells feeding on the same resource (sugar, e.g. glucose), using two different metabolic strategies. The left cell uses respiration, with a low rate (thin arrows) but high yield (high ratio of arrow thickness). The right cell uses fermentation, with a high rate (thick arrows) but low yield (lower ratio of arrow thickness). Thus, the right cell can grow faster but partly wastes the resource and uses it up much faster. This can be compared to two factories utilizing a common resource such as ore, water or wood with different efficiency and speed.

Over the last decade, metabolic games have attracted more and more interest. Different metabolic pathways can be regarded as different strategies in the sense of game theory. Pathways can be restricted to the interior of the cell as well as comprise reactions outside of the cell, as in the case of degradation of substances by means of extracellular enzymes.

The publication of Pfeiffer and coworkers³² is a pioneering paper on game-theoretical approaches in biochemistry, dealing with cooperation and competition in ATP-producing pathways. They analyse the two ATP-producing pathways of respiration and fermentation. Both pathways start with glycolysis and differ in that respiration involves the tricarboxylic acid cycle and respiratory chain, while fermentation includes one or several steps in which pyruvate is converted to an excreted product such as lactate, acetate or ethanol. The former pathway has an ATP-over-glucose yield of about 20–30³³ and a relatively low rate, while fermentation only has a yield of two but its rate of ATP production is usually higher than that of respiration. For example, in striated muscle cells, where both pathways exist, fermentation is up to 100 times faster (c.f. Voet and Voet³³). Mixed strategies in the sense of simultaneous usage of two or more pathways occur as well: For example, baker's yeast uses respiro-fermentation, meaning that, when sufficient glucose is available, respiration proceeds at maximum rate and fermentation is used to a large extent in addition. While respiration utilizes the common pool of the resource efficiently, it has the drawback that it leads to a lower growth rate for pure respirators. The proportions in mixed strategies may differ. For example, in baker's yeast, so-called "ants" have lower glycolytic gene dosage and have a small cell size but reach a high carrying capacity, while the "grasshoppers" have higher glycolytic gene dosage, consume glucose more rapidly, and reach a larger cell size and lower carrying capacity³⁴.

In a spatially homogeneous environment, a model of or-

dinary differential equations leads to a “tragedy of the commons”, where all cells use the shared resource inefficiently by additionally fermenting sugars (defector strategy)³². The “tragedy of the commons” is a situation widely considered in evolutionary game theory to describe evolution towards the inefficient use of a common resource^{32,35–37}. It also occurs frequently in an economic context, for example, in the overfishing of oceans or over-extraction of groundwater³⁸. The short-term “advantage” of inefficient resource usage may explain why many micro-organisms such as *Saccharomyces cerevisiae* (baker’s yeast) and *Dekkera bruxellensis* (used technologically for brewing Belgian beer) mainly rely on fermentation even under aerobic conditions. In contrast, other micro-organisms such as *Kluyveromyces marxianus* and *Pichia fermentans* use respiration under aerobic conditions and fermentation under anaerobic conditions. They can be considered as cooperators. A compilation of data on the trade-off between rate and yield in several micro-organisms can be found in MacLean³⁷. The question arises how they could escape the “tragedy of the commons” during evolution.

A transition to cooperation can be achieved by spatial structure³⁹. Implementing spatial structure in a model consisting of partial differential equations (PDEs) where cells as well as nutrients are allowed to diffuse can promote respirators. For low resource influx rates as well as low diffusion rates, respirators coexist with respiro-fermenters or even take over the population³².

Frick and Schuster⁴⁰ study the non-spatial version of the Pfeiffer model³² at the level of competing populations rather than cells. The fermentation-respiration game can then be interpreted as a Prisoner’s Dilemma, the two-player analog of the “tragedy of the commons”⁴¹. The cooperative strategy is to use respiration because it uses the common resource more economically. Fermentation is the defector strategy because it provides a short-term advantage but wastes the common resource. In the Prisoner’s Dilemma, the only Nash equilibrium is where both players defect, although they would obtain higher payoffs if they cooperated, see Appendix B. This is because the highest payoff is obtained by a defector playing against a cooperator, so that there is an incentive to defect. In fact, this applies to the respiration-fermentation game. As shown by Aledo *et al.*³⁵, a similar game can arise when only the glycolytic pathway is considered, but operates in a more efficient and a less efficient mode. Also in this situation, rapid and inefficient use of glucose is favored by natural selection when the environment is spatially homogeneous.

MacLean and Gudelj⁴² extend the game-theoretical problem of yield versus rate in ATP production by including temporal structure. Their ordinary differential equation model introduces seasonality, resembling a batch culture. Their results are in line with results from ecology saying that oscillations in the environment promote coexistence.

The respiration-fermentation game is also relevant in the competition between cancer cells and healthy cells in humans and higher animals (see Section on Malignant tumour growth). This is one possible explanation for the Warburg effect⁴³, saying that tumour cells mainly rely on glycolysis. As shown by a simple, analytically tractable model, the choice of optimal pathway depends on protein costs as well⁴⁴. If the enzymes of the high-yield pathway are costly in their synthesis, which is the case for the enzymes of the respiratory chain, the low-yield pathway is favoured. If not, then the high-yield, low-rate pathway is favoured.

The secretion of intermediates that are toxic to other cells (ethanol, acetate and lactate in the cases of *Saccharomyces cerevisiae*, *E. coli* and tumours, respectively) is a further effect in the competition^{42,45}. MacLean and Gudelj⁴² include the ethanol excreted by respiro-fermenters into their model, thus imposing a cost in terms of reduced reproductive yield to those cells. This effect also promotes coexistence of respirators and respiro-fermenters.

Using ordinary differential equation models and calculating ESSs, Frank^{46,47} analyses further factors relevant for the trade-off between rate and yield: kin selection, mutation rates and colony demography. Lower genetic relatedness as well as higher mutation rates favour rate over yield. Yield is favoured over rate for long colony survival times.

3 Public goods

Besides the trade-off between rate and yield, public goods games play a major role in metabolism. The production and secretion of, for example, extracellular enzymes is costly to the cooperating cell and benefits surrounding cells (even if the latter do not contribute to the public good). Productivity is often reduced by “cheater” mutants. This is known since the early 20th century as for the secretion of proteases by *Streptococcus lactis* in milk and represents a major problem in the dairy industry⁴⁸.

Besides general models on exoenzyme games, models tailored to specific extracellular enzymes have been proposed. Greig and Travisano⁴⁹ published an experiment on the secretion of invertase in baker’s yeast. They find that the individual advantage of cheaters increases with total cell density. Their interpretation that the invertase game would always have the structure of a Prisoner’s Dilemma is, however, doubted by several groups, among them Schuster *et al.*⁴¹. Gore *et al.*⁵⁰ presented experimental results supported by a phenomenological model. Both their experiments and model show that the secretion of invertase is a harmony game when the cost-to-benefit ratio of secretion is low, hawk-dove game (medium cost-to-benefit ratio) or Prisoner’s Dilemma (high cost-to-benefit ratio). The different game types are explained in Appendix B and the transition between them upon parameter change can

be seen in Fig. B1. The same conclusion is derived by Schuster *et al.*⁵¹, based on a model that is closer to biological mechanisms and simulating the effect of total cell density. Coexistence can arise because of spatial gradients of the nutrient – the glucose concentration is lower around defector cells than around cooperators. This is consistent with numerical simulations based on a grid model⁵² and with experimental studies and a model of protease secretion by *Lactococcus lactis*⁵³. These results can be interpreted in that secreting cells do not cooperate due to altruism but because it is favourable to increase glucose concentration locally. Both cell types can coexist when this advantage outweighs the higher costs of production and secretion of the extracellular enzyme.

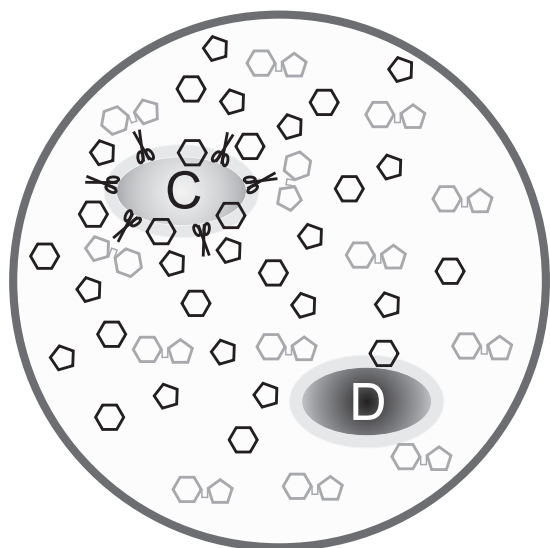
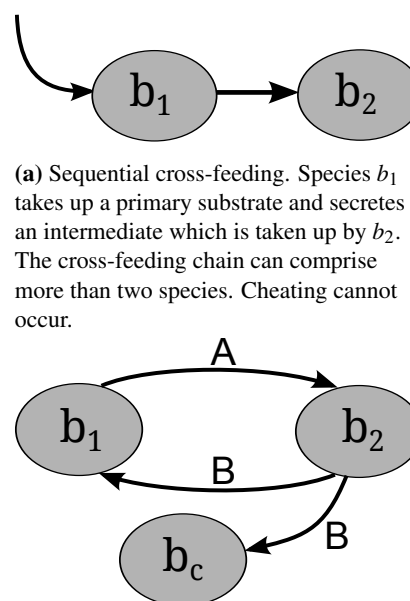


Fig. 2 Scheme of invertase secretion by baker's yeast. Defector cells (marked D) do not produce invertase and benefit from the glucose (hexagons) released by the enzyme (indicated by scissors) secreted by cooperating cells (C). The glucose concentration is higher near cooperating cells unless the system is well mixed, providing an advantage to these cells, while the cost of secretion is a disadvantage. Fructose, the second constituent of sucrose, is indicated by pentagons.

In the case of a Prisoner's Dilemma, the defectors may even drive the population to extinction^{51,53}. An even more detailed model of invertase secretion by yeast (including the metabolism of fructose in addition to that of glucose) is proposed by MacLean *et al.*⁵⁴. They derive the interesting conclusion that, under certain conditions, a mixture of cheaters and cooperators can be better for the population than the absence of cheaters. This may be due to a nonlinear upregulation of invertase in cooperating cells when glucose goes down due to cheater cells.

4 Cross-feeding

Cross-feeding occurs whenever an organism uses metabolites that are produced by another organism for its own growth and maintenance⁵⁵. "Cross-feeding" and "syntrophy" are synonymous terms^{56,57}. Two different types of cross-feeding can be distinguished. First, sequential cross-feeding in which one species b_1 partially degrades the primary resource and excretes an intermediate or waste product, which is then taken up by another species b_2 (or another strain of the same species) and used as a secondary resource⁵⁸ (see Figure 3). Second, cooperative or reciprocal cross-feeding is the mutually beneficial exchange of nutrients among species⁵⁹.



(a) Sequential cross-feeding. Species b_1 takes up a primary substrate and secretes an intermediate which is taken up by b_2 . The cross-feeding chain can comprise more than two species. Cheating cannot occur.

(b) Reciprocal cross-feeding. Two different metabolites A and B (essential for the growth of the respective organism) are exchanged between bacteria b_1 and b_2 . Cheating occurs for bacterium b_c which does not return any metabolites.

Fig. 3 Two different types of cross-feeding as described in the text.

Examples for cross-feeding in nature involve anaerobic methane oxidation by methanogenic and sulfate-reducing bacteria⁶⁰, which leads to microbially mediated pyrite formation, degradation of xenobiotic compounds^{58,61} and the partial degradation of glucose to acetate⁶². It has been known since the end of the 19th century that nitrification is performed in two consecutive steps by two distinct groups of bacteria: ammonia-oxidizing and nitrite-oxidizing bacteria²⁷.

The question arising here is why a homogeneous population that degrades the primary resource completely does not perform better than a population of cross-feeders⁵⁸. Some

possible explanations are outlined below. Moreover, an intriguing problem is how mutualistic cross-feeding can be robust against cheaters. If species b1 and b2 overproduce and secrete substances A and B respectively, and these substances are taken up by the respective counterpart, a cheater mutant of b1 that does not overproduce nor secrete A would, in a spatially homogeneous habitat, always have an advantage. It saves the cost for overproducing A while still benefitting from taking up B (see Figure 3).

Several theoretical studies examine the conditions under which cross-feeding can evolve. Doebeli⁵⁸ uses adaptive dynamics (see Appendix A or Hofbauer and Sigmund⁶) to investigate when an evolutionary branching can occur so that the population splits into two diverging phenotypic clusters representing the two resource specialists coexisting in a (sequential) cross-feeding polymorphism. Doebeli⁵⁸ compares the situation in a chemostat with that in a serial batch culture. He concluded that cross-feeding is less likely in serial batch cultures, because the secondary nutrient is available at appreciable levels only for a relatively short period of time, thus making conditions harsher for specialists on the secondary resource, which is in agreement with experimental results⁶².

A different approach to study the evolution of sequential cross-feeding is proposed by Pfeiffer and Bonhoeffer²⁶. They search for so called evolutionary endpoints, which are analogous to the concept of an evolutionarily stable strategy (ESS), see Appendix A, by simulating competing strains that differ in the expression of the enzymes of a certain metabolic degradation pathway. They assume that the rate of ATP production is maximized and that the total concentrations of enzymes and of intermediates of the pathway are minimized. These two assumptions provide a straightforward explanation for the advantage of sequential cross-feeding: Both species or strains may thus carry less enzymes and also less intermediates. They predict that partial resource degradation is more advantageous than complete degradation if the resource concentration is high or if the concentration of intermediates is restricted to a low value. They find that for intermediate dilution rates of the chemostat the coexistence of complete and partial degradation and coexistence between two partial degraders is possible. At very high or very low dilution rates a coexistence of these strains is very unlikely, which is in agreement with the results from Doebeli⁵⁸. It is concluded that cross-feeding emerges naturally under the above-mentioned extremality principles. On the basis of similar principles, Costa and coworkers²⁷ explained division of metabolic labour by bacteria in nitrification.

A consortium degrading xenobiotics is studied by Katayama *et al.*⁶¹. They develop a mathematical model based on *Sphingomonas sp.* TFEE and *Burkholderia sp.* MN1 degrading fenitrothion, both lacking the complete pathway. Therefore, both bacteria partially degrade fenitrothion. *Burkholde-*

ria lives on the intermediate excreted by *Sphingomonas*. Their findings are in accordance with Pfeiffer and Bonhoeffer²⁶: The syntrophic association observed in this system possesses the minimum number of pathways along which fenitrothion is completely degraded and the total concentration of the intermediates is minimized.

Another, rather different approach is chosen by Gerlee and Lundh⁶³. They investigate the evolution of cross-feeding in a population of artificial individuals that are capable of successively degrading artificial metabolites from which they extract energy for self-maintenance and reproduction. By computer simulations, Gerlee and Lundh⁶³ find out that cross-feeding is the more favoured the more the flow rate of new metabolites into the system is decreased. This is explained by arguing that with low influx rates of new metabolites each metabolite stays longer in the system, such that its probability to be degraded by several organisms is increased. Thus, interactions between organisms are strengthened. That is in accordance with the result in reference Pfeiffer and Bonhoeffer²⁶ that a coexistence of bacterial strains is more likely to evolve if the dilution rate of the chemostat is low.

In a theoretical study by Bull and Harcombe⁵⁹, a model for the growth of two populations interacting by reciprocal cross-feeding is developed. Therein, ordinary differential equations are used to model a system that resembles Maynard Smith's haystack model⁶⁴. Three general factors are found to affect the evolution of cooperative cross-feeding. First, the population needs to be spatially structured in order to ensure that a returned benefit can be directed to individuals or clones, not entire populations. Second, cross-feeding must pre-exist in that, in the beginning, one partner already provides a benefit to the partner by overproduction of, for example, a byproduct. Third, cross-feeding is more easily selected when its cost to the donor is low in comparison to the benefit of the recipient. A further important finding of that paper is that cooperative cross-feeding is favoured during growth at intermediate population densities.

The model by Bull and Harcombe is subsequently revisited by Estrela and Gudelj⁶⁵, using non-linear dynamics. The following outcomes contradict those found by Bull and Harcombe: First, instead of cross-feeding being favoured at intermediate densities, cross-feeders may never be able to outgrow non-cross-feeders at intermediate densities. Second, the shape of the trade-off functions between the cost and benefit of cooperation has a profound effect on the success of cross-feeders in comparison to non-cross-feeders. Third, the result that a small population of cheaters cannot invade an already established population of cooperating cross-feeders requires the same carrying capacity for all interacting populations. They come to the conclusion that cooperative cross-feeding is a robust interaction.

Estrela *et al.*⁵⁵ study a special interaction: a one-way

byproduct cross-feeding in which a cross-feeder species takes up a by-product from a producer species that is toxic to the producer. This cross-feeding interaction is favoured by intermediate toxicity degrees, an increasing by-product production and an increasing consumption of the by-product. Furthermore, the consequences of environmental fluctuations are discussed. The authors speculate that such a one-way by-product interaction could evolve to an obligate interaction.

The evolution of cross-feeding can be explained by the concept of an evolutionarily stable strategy or by adaptive dynamics. Interestingly, in the studies cited above, payoff matrices were not used. Especially for the case of sequential cross-feeding on glucose and acetate we suggest a payoff matrix of the form:

$$\begin{array}{cc} & \text{Ac} & \text{Glc} \\ \text{Ac} & \left(\begin{array}{cc} 0 & 2 \end{array} \right) & \\ \text{Glc} & \left(\begin{array}{cc} 3 & 1 \end{array} \right) & \end{array} \quad (4.1)$$

where the strategies Ac and Glc stand for cross-feeding on acetate or glucose, respectively. This would be the game type “Leader III” (see Appendix B). The strain degrading glucose to acetate would be the “leader”. The second strain can either accept the offered acetate and, thus, be the “follower” or try to use glucose as well. It is impossible that both strains use acetate (payoffs of 0) because this must be produced by one of them. If both use glucose, they have to share the resource and, thus, get a comparably low payoff of 1. The highest payoff is obtained by the “leader” in the above sense, while the “follower” gets a somewhat lower payoff (acetate is the less rewarding nutrient), which is nevertheless higher than 1. Also, the Prisoner’s Dilemma and the hawk-dove game are discussed as potential games in this context^{66,67}.

However, for reciprocal cross-feeding the concepts of two individuals playing against each other or one individual playing against a population are not sufficient⁶⁸. Here, the game is running on the level of two (or more) populations against each other. Usually, cross-feeding are types of N-player games⁶⁸. Furthermore, we can have four types of players: two partners each of which can cooperate or cheat. Also, the game does not necessarily have to be symmetric. Thus, existing concepts need to be refined or new ones need to be developed. Archetti and Scheuring⁶⁷ discuss the importance of non-linear public good’s games and propose for the problem of cooperative cross-feeding between two species a game of two simultaneous public good’s games. Assuming that the fitness function of the public goods are non-linear, they show that, without the ability to discriminate between cheater and cooperator and in the absence of sanctions, a stable coexistence between cooperators and cheaters is possible.

5 Cyclic competition

A central question in evolutionary biology is to identify mechanisms maintaining and even increasing biodiversity. How can different phenotypes (strategies) of the same species coexist?

In several ecosystems a non-transitive, cyclic competition of three species has been observed, which resembles the famous rock-paper-scissors (RPS) game, where rock is wrapped by paper, paper is cut by scissors and scissors are crushed by rock. Jackson and Buss⁶⁹ describe such cyclical dynamics of coral reef invertebrates and Sinervo and Lively⁷⁰ observe the rock-paper-scissors structure of mating strategies of the side-blotched lizard *Uta stansburiana*.

An interesting aspect of the RSP game is that it arises from the Prisoners Dilemma game by adding a strategy called “loner”⁷¹. This additional strategy implies that the players are no longer stuck in the situation “defect, defect” (see Appendix C).

The most famous and frequently investigated example in microbiology is given by colicin producers and non-producers in *Escherichia coli* populations^{72,73}. An early work describing the difference between two competing strains and three competing strains is that of Durrett and Levin⁷⁴.

Strains that produce colicin (P) also have to be resistant against it. Non-producing strains can either be resistant (R) or sensitive (S). Producing toxins against competitors is called allelopathy. Pairwise competitions show that P strains poison S strains, S beats R because of R having the costs of being resistant and R outcompetes P because of P bearing additionally the costs of producing colicin. Neumann and Schuster^{75,76} investigate a model with intrinsic growth rates and self limitation terms. Due to the competition for nutrients among the three strains and the poisoning effect caused by the colicin producers all interaction coefficients are assumed to be negative. This model is structurally stable. Depending on parameter values a stable steady state, stable limit cycle or heteroclinic cycle is obtained. The latter describes an oscillatory dynamics converging to one of the three pure states. Such a dynamics is indeed found in experiments⁷². In Neumann and Jetschke⁷⁷ this model is developed further by the use of adaptive dynamics. This leads to the interesting prediction that the toxicity should decrease during evolution.

In many situations of cyclic dominance in biology, spatial structure is of importance. To include this, partial differential equation (PDE) systems, here reaction-diffusion models, can appropriately be adapted. Nakamaru and Iwasa⁷⁸ consider the boundaries (traveling waves) between areas dominated by different strains. Adamson and Morozov⁷⁹ use the spatiotemporal reaction-diffusion scheme based on the three-species Lotka-Volterra competition approach, as considered by May and Leonard⁸⁰. Focussing on a variable mobility of the species results in a wide range of spatiotemporal patterns.

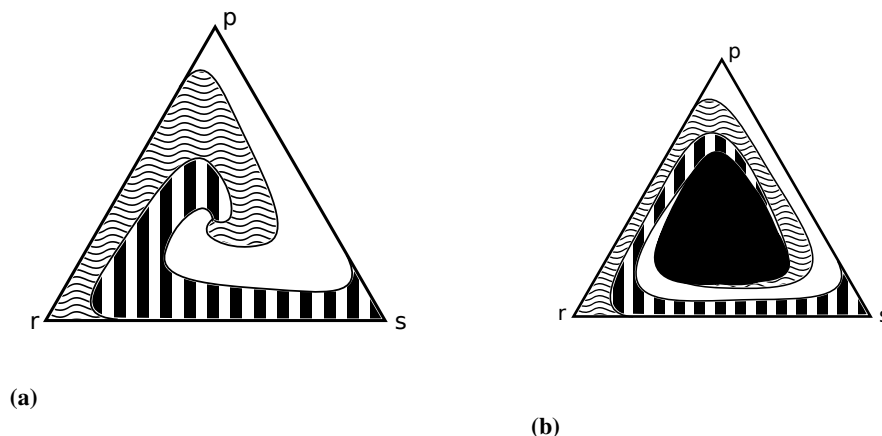


Fig. 4 Simplex representation of the different outcomes of the rock-scissors-paper game⁸¹. Each point of the simplex represents one initial three species population and the pattern shows the final state (white, black and white, curled region, respectively). (a) high mobility (parameter) ends up in extinction - only one of the three strains can survive, which one depends on the initial densities (b) decreasing the mobility parameter leads to an area (black region) in the middle where coexistence is possible.

Several spatial models using cellular automata that represent structured environments and finite populations have been established, often hand in hand with and in comparison to a differential equation model. In Durrett and Levin⁷⁴ the spatial model shows coexistence for a long simulation time, in contrast to the mean-field ordinary differential equation model. The ordinary differential equation model of Frean and Abraham⁸² shows that the populations are not controlled by their own invasion rates, but by the rates of the type they invade. Adapting the spatial model, it still holds that the most aggressive species does not have the largest population (“survival of the weakest”).

An important experimental work is that of Kerr *et al.*⁷². They do experiments with three *E. coli* strains that satisfy a rock-paper-scissors competitive relationship and modify the simulation of Durrett and Levin⁷⁴. When dispersal and interaction are local, coexistence occurs over a substantial range of model parameter values. However, regarding a global neighborhood coexistence never occurs and only the resistant strain survives (contrary to ordinary differential equation models). Károlyi *et al.*⁸³ introduce a mixing step to simulate chaotic flow.

Berr *et al.*⁸⁴ investigate asymmetric interaction rates and compare small and large population sizes with the help of ordinary differential equations. Wang *et al.*⁸⁵ consider mobile species that can carry a virus. The authors study the epidemic spreading (intra- and interspecific). Ni *et al.*⁸⁶ find that the structures of the basins of attraction yielded by a PDE model and by a lattice model are in good agreement.

Müller and Gallas⁸⁷ try to find an explanation for the different outcomes of the model in Kerr *et al.*⁷² (only the resis-

tant strain survives in long-range interactions) and the model in Károlyi *et al.*⁸³ (predicts the winner to be the sensitive strain). They discover that cyclic competition has a characteristic quasiextinction period, during which the sensitive population remains smaller than the other strains, so that the critical factor controlling the final strain surviving is actually the size of the community.

Further cellular automaton models are used to explain the promotion of biodiversity^{88,89} and to investigate the influence of mutation rates^{36,90} or of mobility^{91,92}. Shi *et al.*⁸¹ emphasise the dependence of dynamical properties on initial conditions (see Fig. 4). Venkat and Pleimling⁹³ study mobility and asymmetry effects in one-dimensional rock-paper-scissors games.

Szolnoki and Szabó⁹⁴ and Szabó *et al.*⁹⁵ compare the behaviour of the rock-scissors-paper game on different networks (for example square lattice, honeycomb lattice, random networks).

Well-mixed and large but finite populations can be modeled with the help of stochastic processes. This approach also emphasises the underlying stochastic character. It was used, for example, by Tainaka⁹⁶, Ifti and Bergersen⁹⁷ (introducing mutations and migrations), and Traulsen *et al.*⁹⁸ (including mutation rates). Claussen and Traulsen⁹⁹ recall the replicator dynamics for large populations. With the help of microscopic stochastic processes it is shown, that also the population size determines whether extinction or coexistence occurs. A critical population size is calculated, above which coexistence is likely.

Last, but not least, there are studies that consider more than three strains^{100–103}. Different modeling techniques and inter-

action topologies are investigated.

There is a lot of literature describing the dynamics of the rock-paper-scissors game. Different modeling techniques depending on the investigated population (finite or infinite) and on the space (homogeneous or structured) can be found as well as different parameters of interest. There are studies concerning the invading probability (speed / interaction rate)^{36,82,84,104}, mutation rates^{36,90,97–99,105,106}, the toxicity of the bacteriocin^{74,75,77,89}, the death rate^{74,89}, mobility of the species^{79,81,85,86,91–93,97}, population size^{84,87,99}, intraspecific competition⁸⁶ and even one- or two-dimensional space^{93,107} and the initial values.

Not all of the literature cited in this section uses game theory in a strict sense. However, as we are interested in the results of the rock-paper-scissors dynamics, we find it worth mentioning all the modeling approaches inspired by this game.

6 Signalling

Signalling is a concept that can be found in many fields like sociology, economics, philosophy and of course in biology¹⁰⁸. The term of signalling is closely related with the concepts of communication and information. Signalling games are two-player games between a sender S and a receiver R¹⁰⁸. Both players have different strategies. The sender has a potentially infinite set of messages and the receiver can choose from a set of actions¹⁰⁸. Furthermore, the sender has a certain type given by nature, which is not known by the receiver. The type could be for example male or female, venomous or non-venomous insects. For example, a non-venomous insect could send the message that it is non-venomous or pretend that it is venomous. In contrast to some classical two-player games, like the Prisoner's Dilemma or the hawk-dove-game (see Appendix B), the players in a signalling game always act sequentially rather than simultaneously. R responds to the message of S. The payoffs for the two players depend on the type of S, the message sent by S and the action of R^{109,110}. In the signalling games often the term quality is used. It describes a subset of characteristics which are taken into account for the fitness of an individual.

Signalling games in biology deal with the following aspects: how costly is the signal for the sender and how to differentiate between honest and dishonest (cheating) signalling. Often not the real payoff values are calculated but only the ESSs are identified on the basis of the payoff relations. Most of these game-theoretical studies are focused on animals like mammals and birds. John Maynard Smith, the developer of the concept of ESS, and David Harper concentrate on animal signals in their homonymous book¹¹¹. However, game-theoretical analysis of signalling is not only possible for animals, but also for other multicellular organisms such as plants,

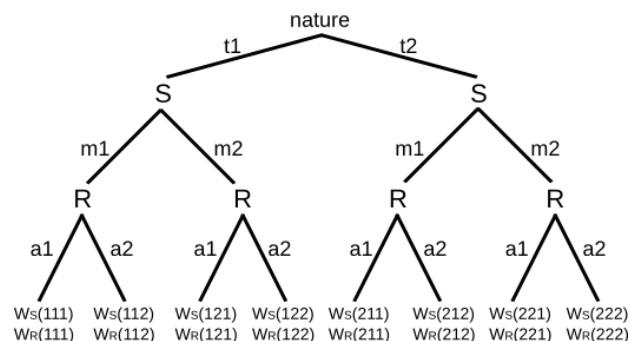


Fig. 5 Tree diagram of a signaling game for the determination of the payoff, which depends on the type (t_1 or t_2) given by nature. The messages (m_1 or m_2) of the sender (S) and the actions (a_1 or a_2) of the receiver (R) can be seen as strategies. W_S and W_R denote the payoffs for the sender and the receiver, respectively, in dependence on the type, message and the action.

and on the cell level.

An important question arising in signalling games is, how evolution produced “honest” or reliable signals between organisms despite the incentive to betray the receiver of the signal. Zahavi¹¹² provides an attempt to explain this with a hypothesis called “Zahavi’s handicap principle”. It suggests that reliable signals have also to be costly to the sender, and therefore implying that other individuals without a specific trait could not afford the cost for this honest signal. An ESS model of the handicap principle is derived by Grafen¹¹³. But the handicap principle is not without controversy, e.g. Bergstrom *et al.*¹¹⁴ find that reliable, honest signals need not be costly.

In a publication by Page²⁹ the ESS of signalling between gametes in lower eukaryotic organisms is analysed. The gametes emit pheromones and affect movement or growth of other gametes in the direction of the pheromone. The result is that with increasing quality of the signalling gamete, also the level how strong the signaller displays its fitness and the probability that the receivers prefer that signaller increase.

In bacteria, signalling is often mentioned in the context of quorum sensing. For example in biofilms quorum sensing in combination with cheating plays a very important role¹¹⁵.

Brown and Johnstone²⁸ develop a two-trait ESS function describing the optimal level of cooperation and of signalling depending on the costs to the individual, benefit to the group, colony size and relatedness of the bacteria. For their ESS calculations, they fixed the costs and passive fitness (fitness of a non-cooperating bacterium in a group of non-cooperators). There is only a stable cooperation effort if the relatedness and colony size are higher than certain threshold values. The highest stable signalling effort is at a relatedness near 0.35 and large colony size.

Quorum sensing with a two-dimensional cellular automaton

of toroidal lattice topology is modeled by Czárán and Hoekstra¹¹⁶. They assume three genetic loci: C for cooperation, S for production of signal molecules and R for signal response. All loci can have an active allele or inactive allele, thus there are eight different genotypes. Different scenarios, with different initial combinations of genotypes and different costs for cooperation are tested. In environments which promote cooperation, the C allele is almost fixed. Simulation of the scenario, in which cooperation is costly, results in a cyclic interaction pattern of the genotypes “Honest” (CSR), “Vain” (CSr) and “Blunt” (Csr) comparable to a rock-scissors-paper game (see section 5). The “Vain” can invade a population of “Honest”, because it saves the costs for response. The “Blunt” has no costs for the QS, that is why it invades the “Vain”. This unconditional cooperater can be invaded by the “Honest”, because the last one only cooperate if there are QS-signals, so if it is surrounded by “Blunts” it has not to pay the costs for cooperation but benefits from the public good. If cooperation is relatively cheap, the simulation results in a dynamical coexistence of “Honest”, “Vain” and “Blunt” genotypes reminiscent of the rock-scissors-paper game (see Section 5). The authors also investigate other environments with other fixed alleles or no fixed alleles.

The work of Lotem *et al.*¹¹⁷ deals with unconditional altruism through signalling benefits. Three different strategies are assumed: Unconditional altruism, helping all other players; defection by soliciting but never donating help; and conditional altruism or tit-for-tat strategy¹¹⁸. The latter is a strategy in the iterated Prisoner’s Dilemma implying to replicate an opponent’s previous action. There are a total of six ESSs. Provided that the signalling benefit of donating help only exceeds the costs for some individuals (high-quality players) and not for others (low-quality players), the ESS is that the high-quality players are unconditional altruists and the low-quality players are defectors or play tit-for-tat.

Traulsen and Nowak¹¹⁹ develop a tag-based model for cooperation. The basic idea is that individuals recognise each other via arbitrary signals, so-called tags. A population of cooperators (or a fraction of this population) is able to change their secret tag when a defector misuses the tag. In the theoretical study of Traulsen and Nowak¹¹⁹ the authors find that there are more cooperators than defectors if $b/c > 1 + (2u/v)$ (with b benefit; c costs; u mutation rate changing only the strategy; v mutation rate changing both strategy and tag). If there are K different tags and the mutation rate between all phenotypes is constant, the constraint is $b/c > (K + 1)/(K - 1)$.

An example for a tag-based system is the *csA* gene in *Dicystostelium discoideum*, encoding a homophilic cell adhesion protein¹²⁰. In a mixed culture of wild-type and *csA*-knockout cells, the wild-type cells are more altruistic but they can distinguish between their neighbours and direct the benefit pref-

erentially to other wild-type cells. This effect is called “green-beard” mechanism¹²⁰, meaning that cooperators can recognize each other by a striking trait.

7 Host cells meet parasitic cells

Several papers analyse two-player games between a host and its pathogen^{121–123}. In Renaud and de Meeüs¹²¹ a two-player game between one host and one pathogen is analysed. The two players can choose between an aggressive strategy that seeks to eliminate the adversary (killer strategy) and a less aggressive strategy (diplomat strategy). The killer strategy is always an ESS, whereas the diplomat strategy is a second solution only under certain parameter settings (costs of virulence and resistance).

In Frean and Abraham¹²² an endosymbiont-host association is studied additionally by a two-populations simulation (intra- and inter-specific competition). Huang *et al.*¹²³ uses focal-point theory to investigate three host-microbe game models (cooperation, dilemma, conflict).

A spatial model on the basis of a cellular automaton is given by Rauch and Bar-Yam¹²⁴. The effect of long-range mixing of species is investigated, which occurs for example due to human intervention and to global transport of an infectious disease.

A third general host-pathogen approach (considering infectiousness) is based on epidemiological modeling and adaptive dynamics: Ordinary differential equations for susceptible and infected subpopulations are established^{125,126}, subsequently some fitness functions are found (function A in the Appendix A) with the help of the solution of these ordinary differential equations and final investigations on evolutionary stability.

Studies considering parasites competing within a “constant” host are for example the work of Bremermann and Pickering¹²⁷, van Baalen¹²⁸ and Day *et al.*¹²⁹. There, individual parasites are considered as players. In Bremermann and Pickering¹²⁷ it is assumed that the disease-induced mortality rate of the host grows with the increase of a parasite’s reproductive rate. In order to extend the length of reproduction time, competing parasites within a host may prefer to reproduce at rates below their maximum reproductive rates. The work by van Baalen¹²⁸ shows that different evolutionary outcomes are possible if the parasites coevolve with the host (a common but quite avirulent parasite meets a host investing little in defence or a rare but virulent parasite in a heavily defending host). In Day *et al.*¹²⁹ parasite’s virulence and host clearance rate are considered as strategies. It is found that the total infection-induced mortality can be a by-product of host immune response.

Several studies investigate the evolution of host defence against a fixed parasite strain. While Boots and Haraguchi¹³⁰ examine resistance strategies of the host, Bonds *et al.*¹³¹ focus

on the evolution of host sociality with respect to an infectious disease and Miller *et al.*¹³² study the influence of the host life span on different host resistance mechanisms. In Best *et al.*¹³³ the question why hosts are differently tolerant is analysed.

The coevolution of host and pathogen is modeled for example by Taylor *et al.*¹³⁴. The level of virulence that the pathogen exhibits and the level of immune clearance that the host exhibits are used as continuous strategies. Further coevolutionary population models are reviewed in Boots *et al.*¹³⁵.

The interesting effect of host infection on the evolution of biodiversity of pathogen strains is studied by Morozov and Best¹³⁶. It is argued that the role of host-parasite interaction can be important in explaining polymorphism. The model is a combination of an epidemiological model and a predator-prey model and is analysed by adaptive dynamics and evolutionary game theory.

Alonzo and Calsbeek¹³⁷ build a game with three strategies: a reproductive parasite and two different resource defenders. The model shows continuous sequential invasion and no stable coexistence.

In order to investigate age-dependent behaviour variation¹³⁸ and heterogeneous host populations¹³⁹ epidemic modelling is combined with game theory. Williams¹³⁹ argues that individuals are differently susceptible and vulnerable to a pathogen and these differences may also influence the pathogen evolution.

In the following, we present some interaction models with specific biological application.

Wolf and coworkers^{140,141} explore the origin and utility of random phase variation (RPV). This strategy can be described as a random switching of phenotypes in order to adapt to rapidly varying environments without requiring mutation. Bacteria as players have sensors and a number of options to react (e.g. cell states or phenotypes). A strategy is defined as a map from sensor information onto behaviour. Certain sensor defects (unobservable, incorrect, delays, noise) and different environments (time-invariant, time-varying, stochastic) are analysed for combinations that give rise to RPV as an ESS. As RPV can be seen as random alternations between losing strategies, that form a winning strategy when sensing is absent, an example for the Parrondo paradox is found. The Parrondo paradox describes the effect that mixing losing strategies can produce a winning outcome. The problem is solved with a matrix population model, that is based on Markov chains.

Endosymbiosis of proto-mitochondrial prokaryotes into proto-eukaryotic host-cells is investigated in Bivort *et al.*¹⁴². A game-theoretical two-player game analysis suggests that active symbiosis would not have been favourable to both species as an asymmetric Prisoner's Dilemma arises from the payoff matrix, indicating that some assumptions are not adequate. Agent-based model simulations exhibited potential conditions that could lead to a stable symbiosis.

Eswarappa¹⁴³ studies persistent infections. Bacteria stay in both intracellular (mostly phagocytes) and extracellular compartments of the host. Bacteria are safe inside phagocytes if they can avoid lysis. Here, the bacteria are protected from immune recognition and serum mediated extracellular killing by the host system. Furthermore, bacteria can access the external environment for dissemination. Examples for such bacteria are *Helicobacter pylori*, *Mycobacterium tuberculosis* and *Salmonella spp.* On the other hand, the host can use extracellular or intracellular defense mechanisms. A very basic and heuristic model shows three Nash equilibria: two pure equilibria that are not biologically plausible and a mixed Nash equilibrium that exists independently of the payoff relations and is biologically relevant. Bacteria remain in extra- and intracellular compartments of the host and the host uses extra- and intracellular defense mechanisms.

The polymorphic fungus *Candida albicans* has two strategies when engulfed by macrophages: avoiding lysis transiently (silencing) or forming hyphae and escaping (piercing). Two different games among *C. albicans* yeast cells inside the macrophage (macrophage is considered as the environment) are developed by Hummert *et al.*¹⁴⁴. In the two-player game it is assumed that each macrophage engulfs exactly two yeast cells. If costs for silencing are not too high, the game is a hawk-dove game with a single mixed ESS.

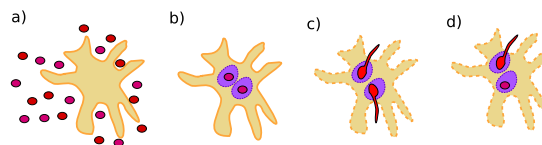


Fig. 6 Possible interaction scenarios: (a) *C. albicans* yeast cell population and a macrophage before ingestion, (b) both ingested cells silence, (c) both ingested cells pierce (d) ingested cells adopt different strategies. Figure taken from Hummert *et al.*¹⁴⁴.

$$\begin{array}{c} p \\ s \end{array} \begin{array}{cc} p & s \\ \left(\begin{array}{cc} l-c & l-c \\ l & 0 \end{array} \right) \end{array} \quad \begin{array}{c} p \\ s \end{array} \begin{array}{cc} p & s \\ \left(\begin{array}{cc} l-c & l-c \\ (1-e^{-\lambda})l & 0 \end{array} \right) \end{array} \quad (7.1)$$

The left Matrix of (7.1) shows the payoffs, where p stands for the strategy piercing, s stands for the strategy silencing, l stands for having survived, and c having payed the costs for piercing. In the second game, which is a playing the field game (see Appendix A), a Poisson distribution for the number of ingested *C. albicans* yeast cells is assumed. The fitness matrix is shown in the right part of equation 7.1. An entry gives the payoff of an individual playing a row strategy against a pure population playing the column strategy. The ESS in a mixed population (under certain parameter conditions) can be calculated, but also a pure piercing population can exist (under contrary parameter conditions). Additionally, survival rates

of phagocytosed yeast cells can be calculated. A comparison with experimental studies showed that there exist two different karyotypes, a pure piercing population and a mixed population, the difference is explained in the difference of the costs these two karyotypes have, or in other words, in the ability to secrete proteins that enhance producing hyphae.

Bewick *et al.*¹⁴⁵ investigate two different HIV variants and the corresponding adaptive immune responses. Two different scenarios are analysed and the Nash-equilibria are calculated. In another work Bewick *et al.*¹⁴⁶ model the immune defense against hepatitis B in the liver by comparing the immune defense strategy with military defense strategies.

8 Malignant tumour growth

Cancer is known to be a genetic disease where accumulation of mutations can transform cells from a homeostatic tissue into a complex ecosystem of individualistic cells that compete for limited space and resources^{147–149}. Cancers are also very heterogeneous, both genotypically and phenotypically. A single tumour may contain several different types of tumour cells¹⁵⁰, with each type potentially having different traits and ways to interact with other cells and the local microenvironment. This heterogeneity is passed on from parent to daughter cells and affects their probability of proliferation and survival. Furthermore, this heterogeneity is key to explain important aspects of cancer such as the emergence of increasingly more aggressive cell types and resistance to treatments.

Tumour cells are thus part of an ecosystem where interactions between different cells and the environment determine the somatic evolution^{147,148,151}. Any given tumour can be made of a number of different highly dynamic microenvironments^{150,152}. Studying cancer progression requires access to mathematical tools where the competition of different cancer phenotypes and strategies can be taken into consideration¹⁵³. The use of evolutionary game theory to model and analyse cancer progression is still a nascent field with only about 15 years of history and just a few papers¹⁵⁴. In that time it has been used to study cancer progression both *in vitro* (as a loose collection of tumour cell populations growing out of control) and also *in vivo* in organs such as the brain, bone, prostate or the lung. Indeed, most uses of evolutionary game theory in cancer so far involved studying how certain tumour phenotypes are selected for. This is a particularly important part of cancer progression as new phenotypes emerge in the tumour population and, provided they are fitter than the current mix of phenotypes, grow and become an important (but rarely the only) subpopulation in the tumour. The emphasis on phenotypic transitions as hallmarks of cancer has been previously highlighted^{155–157} and considered therapeutically.

The evolution of increasingly more aggressive phenotypes is studied in the context of the competition between differ-

ent cell lines by Anderson and colleagues¹⁵⁸. The study focuses on the emergence of phenotypes with ever decreasing dependence on the local microenvironment, one of the hallmarks of cancer aggressiveness¹⁵⁷. The results of the game between cells that are microenvironmentally independent and those that can rely on the microenvironment, shows that the two types are likely to be found coexisting under most circumstances. Other studies on the emergence of aggressiveness focus on glioblastomas, malignant brain cancers characterised especially by high invasiveness and motility of the constituting cancerous glia cells. In particular, the emergence of invasiveness is studied in the context of glioblastomas using a cellular automaton model whose rules are based on game theory but where fitness is not defined explicitly but implicitly via the phenotype¹⁵⁹. A more conventional evolutionary game theory approach^{31,160} considering invasive and proliferative subpopulations, highlights the importance of the microenvironment in the progression to malignancy. The model results are later corroborated via an agent-based model³¹. Both models consider a population of rapidly proliferating cells where a mutation might confer a higher degree of motility to certain cells at the cost that cells that move cannot divide at the same time. In essence, both models present a game that is similar to the standard hawk-dove game¹⁶¹ so that the local population of proliferative cells is better off interacting with the new motile cells than with other proliferative cells but a population composed of proliferative cells is still susceptible to be invaded by invasive ones. This is akin to hawks receiving a higher fitness payoff from their interactions with doves but still being susceptible of being invaded by a small number of doves (under the assumption that injuries can be costly). As in most evolutionary game theory, a payoff table completely describes the model:

	Motile	Autonomous Growth
Motile	$(1 - c/2)$	$1 - c$
Autonomous Growth	1	$1/2$

This simple game leads to an equilibrium described by this equation:

$$p = \frac{1 - 2c}{1 - c},$$

where p is the proportion of motile cells in the population, c is the cost of motility and the ESS is calculated with the Bishop-Cannings theorem. This model has been extended by including glycolytic cancer cells. Emergence of glycolytic cells with a less efficient metabolism that does not require oxygen and produces lactic acid as a byproduct, is considered one of the hallmarks of cancer¹⁶² and known as the Warburg effect⁴³ (see also Section 2). The Warburg effect has been previously studied with the help of mathematical models that show that there are circumstances under which glycolysis can be beneficial to a tumour cell. This is important because traditionally

glycolysis has been viewed as a low yield alternative to regular metabolism. However, cells with a glycolytic metabolism are known to emerge in hypoxic regions of the tumour. It has been hypothesised that, in rare occasions, glycolysis can lead to a Prisoner's Dilemma game in which cells with glycolytic metabolism could be considered cooperators whereas the traditional aerobic cells would be defectors of the game⁴⁵. Lactic acid being produced by glycolytic cells can be viewed as a common good for these cells. Alternatively, as done in Section 2, one may argue the other way round and consider the glycolytic cells as cheaters.

Meanwhile, the invasion/proliferation game mentioned before has been enriched with the addition of cells with a glycolytic phenotype and interpreted in a glioblastoma context¹⁶³. The results of this game suggest that invasiveness is more likely to emerge in tumours where there are glycolytic cells present, strengthening the notion that non-competitive interactions between tumour cells play an important role in glioblastoma progression. Further work studying secondary glioblastomas characterised by the IDH1 (isocitrate dehydrogenase) mutation, suggest the following mechanism: rapidly proliferating cells grow beyond oxygen reach leading to glycolytic phenotypes that incentivise the emergence of the invasive phenotypes typical in glioblastomas¹⁶⁴. Building on the research by Basanta¹⁶⁵ and Kareva⁴⁵, Archetti¹⁶⁶ has recently modelled the Warburg effect following the idea of lactic acid viewed as a public good (it helps the tumour in small doses but it harms if there is too much of it). A modelling approach that would allow for a nonlinear dependence of the benefit on lactic acid concentration could yield new insights on its role in cancer progression. Archetti uses a non-linear public goods game based on the properties of Bernstein polynomials and suggests that reducing the acidity of the environment might not significantly change the dynamics of the game and thus should not be a useful therapeutical approach.

Evolutionary game theory is thus a very powerful mathematical modeling tool in which to study how the interactions between different cellular populations could give rise to the different hallmarks of cancer¹⁵⁶. Work by Axelrod and collaborators¹⁵¹ nonetheless suggests that not all tumour cells have to acquire all the traits in the hallmarks for the tumour to develop a cancer. The interactions between tumour cells or even stromal cells are not always exclusively competitive. There is then the potential that tumour cells with different capabilities could cooperate so that while no single cell would have all the hallmarks of cancer, a group of cells would collectively have them. Similar types of interactions between individuals in a population have been studied by Nowak before¹⁶⁷ and it is clear now that these interactions play an important role in cancer progression¹⁶⁸ (see Figure 7). They can be mutualistic (if both cells interacting benefit from the interaction), competitive (if both cells can be harmed as a result of the interaction),

predatory (if the benefit of one of the cells comes at a huge cost to the other cell), parasitic (if the benefit to one cell comes at a small cost to the other cell) or commensalistic (if the benefit to one cell comes at no cost to the other cell).

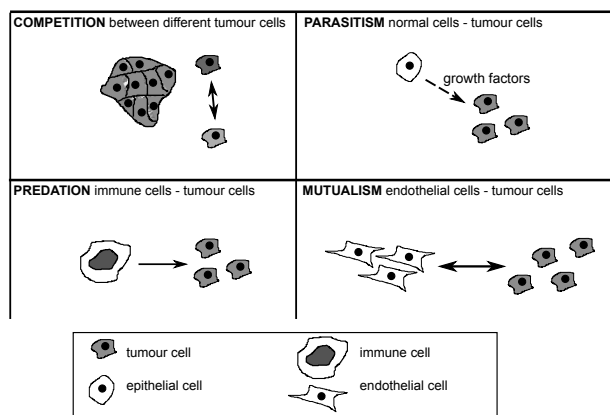


Fig. 7 The four types of interaction among tumour cells and among tumour and healthy cells. For further explanation see text.

Interestingly, the first known application of evolutionary game theory in cancer investigates how angiogenesis can emerge if not all tumour cells contribute¹⁶⁹, thus constituting a model of commensalism. In their model, Tomlinson and colleagues consider two subpopulations of tumour cells, one producing VEGF (an angiogenic factor) whereas the other one devotes all its energy to proliferation. The coexistence shown by their game theoretical model is also later confirmed in a more sophisticated threshold model by Bach and colleagues¹⁷⁰ where the benefits of angiogenesis can only be realised when a majority of interacting cells are producing VEGF. The general mechanism describing the dynamics of cancers with cooperating and cheating subpopulations (where cooperation is defined as the production of growth factors such as VEGF) is also studied using a spatial implementation of evolutionary game theory in the paper by Bach *et al.*¹⁷¹.

The interactions between tumour cells and non tumour cells are equally important. It is well known that tumour cells interact with stromal cells such as fibroblasts, endothelial cells and macrophages, in ways that could be considered co-option¹⁷². Dingli *et al.*¹⁷³ study interactions between different types of tumour cells and stromal cells in the bone (osteoclasts and osteoblasts) in multiple myeloma. The model shows that targeting the cancer cells directly is unlikely to be an effective long term cure of myeloma but targeting the interactions between the tumour and the stromal cells so that those can effectively outcompete the tumour is a more promising approach. In a similar line, game theory is used to study interactions between a heterogeneous tumour with fibroblasts in prostate cancer¹⁷⁴. In this game tumour cells can co-opt the fibroblasts to obtain

certain growth factors or ignore the microenvironment at the cost of having to produce those factors. The study of different treatment options demonstrates the role they have as selection forces. In some occasions, these forces can potentially eliminate the bulk of the tumour but only at the expense of increasing the relative fitness of more aggressive and less treatable phenotypes.

This last study highlights not only the role of evolutionary game theory to understand the role of cell interactions in cancer progression but also how therapies can influence the evolutionary dynamics of a cancer. One example is the use of evolutionarily enlightened therapies such as the evolutionary double bind as advocated by Gatenby and colleagues¹⁷⁵. The evolutionary double bind suggests that combination therapies, where treatments can be considered as tumour cell predators, can be applied so that tumour cells that are resistant to one of the treatments are so at the expense of being even more susceptible to the second treatment. Mathematical models are used to investigate this intriguing hypothesis¹⁷⁶ but more recently a game theoretical approach is used to study a specific case where researchers at the Moffitt Cancer Center find that chemotherapy and a p53 vaccine could work synergistically in lung cancer^{177,178}. The model assumes that, given the two treatment options, a heterogeneous tumour can be composed of three subpopulations: susceptible to both treatments, resistant to the p53 vaccine and resistant to chemotherapy. The interactions between these populations are important since the clinical work could be explained by the parasitism of the p53 resistant populations (providing protection from the immune system to their neighbours) by the subpopulations that are sensitive. The study also suggests that the key to the synergistic effect is the sequence in which the treatments are applied which results in an increasing reduction of tumour heterogeneity.

In summary, cancer is an evolutionary disease where cells that can adapt to the tumour environment will thrive whereas those that cannot become extinct. The evolutionary dynamics of cancer can be understood with the help of mathematical models based on evolutionary game theory, especially regarding two key and intertwined aspects of evolution: the role of interactions between individuals and the emergence of new traits in a population.

Current evolutionary game theory approaches have a number of limitations: they assume homogeneously mixed populations where all the relevant phenotypic strategies are known from the start. Although not a limitation of evolutionary game theory, current game theoretical models of cancer usually disregard the role of space and usually consider the proportions of populations as opposed to the total numbers of a particular phenotype in a tumour. This can obscure the fact that in many situations the proportion of a phenotype could be going down as the number of cells with that phenotype keeps increasing

during tumour growth. Despite the limitations, evolutionary game theory is a very useful tool in which to study the emergence of new, increasingly more aggressive phenotypes that characterise tumour evolution and progression. It is also an ideal framework in which to investigate how treatments can affect selection and alter the evolutionary landscape.

9 Application in medicine and biotechnology

As shown in the previous section by the example of cancer treatment, evolutionary game theory is a promising tool also for applications in medicine, producing counter-intuitive results where conventional approaches fail. The same holds for applications in biotechnology. Both directions are discussed in the present section.

In medicine, conventional treatments of tumours as well as life-threatening infections by viruses like HIV or bacteria often tend to fight tumour cells or pathogens by (single) drug usage at high dosage. What seems to be a success on short-sight often leads to resistance to the given drug and relapse on the long run.

Optimal drug dosage over time is treated generally by Bewick *et al.*¹⁷⁹ in a model that combines evolutionary game theory with optimal control theory¹⁷⁹. Based on the assumption that viruses have to make a trade-off between replication capacity and drug resistance, their results give advice for an optimal treatment scheme in a rather non-intuitive way: When drug resistance increases faster with mutations, leading to resistance, than the replication rate decreases, the optimal strategy of drug dosage is to give high dosages during the early phase of the treatment regime, and then to lower dosages as the treatment process proceeds. In the other case, if drug resistance increases slower than the replication rate lowers during mutational processes, leading to resistance, the optimal drug strategy is to limit drug application during the first half of the treatment phase, and then to increase the drug load over the second half of the treatment phase.

In a similar manner, Wu and Zhang¹⁸⁰ use the combined approach of evolutionary game theory and optimal control theory to propose optimal medication schemes for HIV patients. They come to the conclusion that it is optimal to dose the medication only moderately at later stages of the treatment, so that T cells and HI-viruses attain a stable coexistence rather than maximize the drug load which leads to a rapid resistance mutation of HI-viruses.

Also in cancer treatment, evolutionary enlightened therapies are promising (cf. Section 8). Tumours are now seen as heterogeneous formations consisting of interacting subpopulations, each bearing different properties that can be seen as strategies. Combination therapies successively weaken the cancerous phenotypes, which may be susceptible to all treatments or susceptible to one respective agent of the combina-

tion therapy¹⁷⁸. The key to the synergistic effect is the sequence in which the treatments are applied, which results in a stepwise reduction of tumour heterogeneity.

Interactions not only among tumour subpopulations but also with surrounding cells in the microenvironment attracted more and more interest as therapeutical targets in the fight against cancer¹⁸¹ (cf. Section 8). Game-theoretic models illustrated the role of different treatments as selection forces^{173,174}. Targeting tumour cells directly often merely leads to short-term success, clearing the way for more aggressive and less treatable phenotypes. Instead, targeting interactions between the tumour and surrounding cells, enabling the latter to outcompete the tumour cells, is a promising approach to long-term survival.

Metastasis is one of the clinically most important transitions in tumour progression, leading to more than 90% of cancer deaths¹⁸². Intuitively, “starving out” the primary tumour by limiting its resources to prevent metastasis seems to be a suitable therapeutical approach. However, game theoretic models show that the contrary is the case. Providing a nutrient rich environment to the tumour increases the fitness of proliferative cells over motile cells, acting as a disincentive for a tumour to become invasive and metastasising³¹. Improving tissue oxygenation, which increases the relative fitness costs of tumour cells for switching to anaerobic glycolysis, has a similar effect¹⁶⁵.

As mentioned in the title of their paper, “An analogy between the evolution of drug resistance in bacterial communities and malignant tissues” is seen by Lambert *et al.*¹⁸³. They suggest a fundamental mechanistic similarity between the rapid evolution of resistance to drugs in communities of cells in malignant tissues and in bacterial communities. As a cause of the evolution, they propose a programmed and collective stress response performed by interacting cells and suggest bacterial communities as a model system for tumour cells. From this perspective, a united view based on game theoretical models seems to be reasonable.

Bacteriocins are of current interest as probiotic and bio-protective agents¹⁸⁴ as well as a potential viable alternative to conventional antibiotics^{185–187}, showing promising success in experimental studies on the treatment of chronic biofilm-mediated infections¹⁸⁸. Kirkup and Riley⁷³ emphasise the role of bacteriocins in a game of rock-paper-scissors as potential promoters rather than inhibitors of microbial diversity (in the mammalian gut). Indeed, Brown *et al.*¹⁸⁸ identify bacteriocins as potential selective correctors of imbalances in the microbiome that are associated with chronic conditions such as Crohn’s disease.

Evolutionary game theory is also a promising tool for biotechnology. Game-theoretical analyses of the production of extracellular enzymes bear a great biotechnological potential, for example, in the context of renewable fuels and

biodegradation of xenobiotics. Biotechnological processes are optimized regarding their cost-benefit ratio. High density cell cultivation is thus favourable^{189,190}. However, if production of extracellular enzymes is involved in the biotechnological process, “cheating” mutants deficient in exoenzyme production can arise, which benefit from the exoenzymes produced by “cooperating” cells. Game-theoretical models predict a high percentage of “cheating” mutants at high cell densities and suggest optimal densities for maximised volumetric productivity to be lower⁵¹.

10 Challenges and future prospects

In this and an accompanying review¹², we discuss the use of evolutionary game theory in cell and molecular biology including biochemistry. In spite of its “playful” name, game theory has manifold practical applications, as witnessed by several Nobel prizes in economics. As outlined above (especially in Sections 7 - 9), evolutionary game theory can be applied beneficially in medicine and biotechnology.

While evolutionary game theory has been applied in ecology and behavioral biology since the mid-20th century, applying it to “lower” levels in biology is more recent. As mentioned in the Introduction, an important aspect in this development is that, earlier, it was assumed that the players should have a certain degree of cognitive and rational capabilities, so that only humans and sufficiently complex animals could be considered as players. However, applying the conceptual toolkit of game theory does clearly not require that the players are conscious, rational beings (cf. Maynard-Smith and Price¹⁴). This becomes even clearer from the numerous examples reviewed above. By selection, certain behaviours or features outcompete others, so that, in retrospect, it may look as if a certain appropriate strategy had been chosen deliberately. Interactions between microorganisms can obviously be described by game theory very successfully. Importantly, human behaviour is often superimposed by moral and social factors. This enables more complex solutions such as mutualism based on trust leading to non-Nash states, equilibria based on a rationality in favour of the community or apparently irrational responses like defiance or spite. Therefore, it might be argued that the concept of Nash equilibrium is often much better suited to micro-organisms than to human behaviour.

This raises the general question as to what can be gained by applying game theory to non-cognitive systems that is not provided by more conventional approaches such as optimization theory or classical evolutionary theory. The arguably most important aspect is a better understanding of seemingly counter-intuitive findings such as “survival of the weakest”⁸², as it can occur in the rock-paper-scissors game. As explained in more detail above, the probably most intriguing example is provided by a bacterial strain that is sensitive to a toxin but

can eventually be the only survivor in an oscillatory dynamics because it grows with a higher intrinsic rate than the toxin producer and resistant strains, although it is the least competitive strain. Phenomena like this apparently contradict Darwin's notion of the "survival of the fittest". Whether they actually do so, however, depends on how fitness is defined. The attribute "weakest" strongly depends on definition. In terms of growth rate, the sensitive strain is actually the fittest. In any case, the rock-paper-scissors game is more complex than traditional Darwinian concepts because the transitivity relation of fitness is violated. That is, if A is fitter than B and B is fitter than C, (counter-intuitively) C needs not be fitter than A. This non-transitivity actually implies that there is no weakest, so that "survival of the weakest" is just a provocative notion to make the point.

When analysing multi-cellular organisms, the question arises as to what entities can be considered to be players. One obvious option is to consider entire organisms as players. However, in the case of cancer, the tumours or even the particular tumour cells act (more or less selfishly) as independent agents. This can be considered as an evolutionary regression to unicellular forms of life¹⁹¹. Interestingly, also adipocytes can act independently to a certain extent. These cells excrete signalling substances, so-called adipokines, regulating glucose metabolism. In obese individuals, that signalling is perturbed so that adipocytes become, in a sense, selfish¹⁹².

Game-theoretical approaches can be extended in several directions in the future. They can be extended to involve spatial dimensions, for example, using individual-based models. They can involve a temporal dimension (for example, using differential equations such as in several rock-scissors-paper models, Section 5) and stochastic components. Many dynamical processes can be regarded as iterated games, such as the iterated Prisoner's Dilemma¹¹⁸. These often show very complicated dynamics or it may even occur that no optimal time pattern exists¹¹⁸. This is one possible explanation for the open-endedness of evolution.

Here, we have concentrated on spatially homogeneous situations. Of course, many processes of cooperation and competition occur in heterogeneous setups. Accordingly, many game-theoretical analyses have been performed by spatial models, either continuous models^{32,39} or discrete models on lattices (grids) (e.g. Reichenbach *et al.*⁹¹, Szolnoki and Perc¹⁹³, Allison⁵²). The boundary between game theory and other modelling methods such as lattice models is not clear-cut then. Also in other extensions of game theory, the boundary is not clear-cut, for example, with respect to ordinary differential equation models in population dynamics⁷⁵.

11 Acknowledgements

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Appendix A Mathematical background

Typical tools and concepts in Evolutionary Game Theory are payoff matrices, Nash equilibria and evolutionarily stable strategies (ESS). In a Nash equilibrium, no player would gain an advantage (e.g. in terms of growth rate) when changing strategy unilaterally, that is, without coordinated changes by other players. The related concept of ESS describes a strategy that, if adopted by a population, cannot be invaded by any mutant strategy.

Appendix A.1 Two-player games

Each player can choose a particular strategy from a predefined strategy set. For each strategy combination there are defined payoffs for each player, written down in the **payoff matrix** (outcome of the game).

Example: A symmetric two-player game with strategy set $\{A, B\}$. The payoff for strategy A versus B is given by $W(A, B)$. Because of symmetry it is sufficient to fill the payoff matrix with the outcomes of the row-player (player 1):

$$\begin{array}{cc} & \begin{array}{cc} \text{player 2} \\ A & B \end{array} \\ \begin{array}{c} \text{player 1} \\ A \\ B \end{array} & \begin{pmatrix} W(A, A) & W(A, B) \\ W(B, A) & W(B, B) \end{pmatrix} \end{array} \quad (\text{A1})$$

A strategy is said to be a **dominant strategy** if for every choice of strategies of the other players the payoff from choosing this strategy is greater than the payoffs from choosing any other strategy. Strategy A is dominant if $W(A, A) \geq W(B, A)$ and $W(A, B) \geq W(B, B)$, for all strategies $B \neq A$.

Given a certain amount of pure strategies, a further possibility is to adopt each (pure) strategy with a certain probability. Such a set of probabilities is said to be a **mixed strategy**. Of course, the probabilities have to sum up to one. In the symmetric two-player game with pure strategies A and B, we write x for the mixed strategy to adopt A with probability x and B with probability $(1 - x)$.

An important notion in game theory is the **Nash equilibrium**. It is a situation (a certain strategy combination of all players) where no player has an incentive to choose another

strategy given that all other players stay at their chosen strategy. In a Nash equilibrium, the strategy of each player is the best response to the strategies chosen by all other players¹⁹⁴.

Example: Given the symmetric game formalized in matrix (A1), possible Nash equilibria in pure strategies are:

- (A,A) is Nash equilibrium, if $W(A,A) \geq W(B,A)$
- (B,B) is Nash equilibrium, if $W(B,B) \geq W(A,B)$
- (A,B) and (B,A) are Nash equilibria, if $W(A,B) \geq W(B,B)$ and $W(B,A) \geq W(A,A)$

Note that in the cases (A,A) and (B,B) one can also say that the strategies A and B , respectively, are Nash equilibria. In general, in a Nash equilibrium where all players adopt the same strategy, also the strategy itself is called a Nash equilibrium. Further, if the above inequalities hold as strict inequalities, then the Nash equilibria are called **strict**.

According to the **Bishop-Cannings Theorem**¹⁹⁵, the idea to find a mixed strategy that is a Nash equilibrium in a symmetric game is the following. Each single pure strategy adopted against this mixed strategy does equally well, i.e. the **expected payoff** to a player is independent of its adopted strategy: $W(A,x) = W(B,x)$, where $W(A,x) = xW(A,A) + (1-x)W(A,B)$ and $W(B,x) = xW(B,A) + (1-x)W(B,B)$. Solving this equation for x gives

$$x = \frac{W(B,B) - W(A,B)}{W(A,A) - W(A,B) - W(B,A) + W(B,B)} \quad (\text{A2})$$

Check if the solution is between 0 and 1, because it represents a fraction or probability. If the solution is outside of that range, only the pure Nash equilibria are relevant.

Appendix A.2 Evolutionary game theory

If applying the concepts of game theory to animals, plants, microorganisms, cells or even molecules the prerequisite of rational choices made by the players is not required. Without rational players a Nash equilibrium can be found by an evolutionary process in a population. Individuals using a strategy that is not optimal in the population will reproduce at a lower rate than individuals with an optimal strategy, which will thrive and pass this strategy on to the next generation. New strategies can appear by (epi-)mutation or by immigration into the population.

Among an infinite population we consider repeated random encounters of opponents. In terms of a population, a mixed strategy can also define the state of the population, where certain fractions of the population play one of the strategies. Again, the fractions have to sum up to unity.

A strategy is **evolutionarily stable** if a whole population using that strategy cannot be invaded by a small group with

a mutant genotype. Strategy x is evolutionarily stable if and only if for any mutant type p we have: $W(x,x) \geq W(p,x)$ and if $W(x,x) = W(p,x)$, then $W(x,p) > W(p,p)$. The first condition is the Nash equilibrium condition and the latter one is the evolutionary stability condition, meaning that x can invade any p population.

Appendix A.2.1 Theorems.

1. Every game (given by an $n \times n$ payoff matrix) has at least one Nash equilibrium¹⁹⁶.
2. If all players have a dominant strategy, then the situation where all players adopting it is a Nash equilibrium (NE in dominant strategies)¹⁹⁴.
3. Every evolutionarily stable strategy is a Nash equilibrium, but not every Nash equilibrium is evolutionarily stable¹⁹⁶.
4. Every strict Nash equilibrium is evolutionarily stable^{194,196}.
5. In asymmetric games (asymmetric in payoffs) only strict Nash equilibria are evolutionarily stable¹⁹⁴.

Appendix A.2.2 Playing the field. Several situations are imaginable, where the success of a strategy does not depend on the outcome of pairwise encounters, but on the average strategy used in the population, for example the success of a sex ratio. Another example is the production of exoenzymes by yeast cells^{50,51} (see Section 3). Each yeast cell interacts with many cells rather than only in pairwise encounters. As a first step, we can draw up the **fitness matrix**, where the fitness of a single A strategist in a population of B strategists $W(A,B)$ are written down. Clearly, A will be an ESS if, $W(A,A) > W(B,A)$ and B will be an ESS if $W(B,B) > W(A,B)$. If neither of these inequalities hold, then the ESS is a mixture of A and B . To find the mixed ESS, again $W(A,s) = W(B,s)$ must hold, but the payoff for A and B in a mixed population s is often hard to find and may be a nonlinear function of frequency.

Because a Nash equilibrium does not necessarily have to be evolutionarily stable, one has to check for the evolutionary stability of the (mixed) Nash equilibrium.

Evolutionary stability is particularly useful because it says something about the dynamic properties of a system without being committed to any particular dynamic model.

Appendix A.3 Dynamic game theory

Dynamic game theory is important if one wants to study the behaviour of a system beyond an “equilibrium based viewpoint”¹⁹⁷. Investigating the long-term change of the strategies’ frequencies is only possible when considering dynamics.

It can also be used for studying frequency dependent selection⁷. Further, the notion of evolutionary stability relies upon implicit dynamical consideration. In certain situations, the underlying dynamics can be modelled by differential equations⁶.

Another point of dynamic game theory is the question whether games do reach a Nash equilibrium, and if so, by what process¹⁹⁴? Further questions include: If there are several Nash equilibria, to which one does the game converge? The answers can be found by studying the behaviour of dynamical systems.

The dynamical behaviour of a game is interesting especially if the long-term outcome is not a Nash equilibrium (this is the case for *e.g.* variants of the rock-scissors-paper game). In such a case, the behaviour over time are endless regular or irregular oscillations.⁷ Hence, the static approach often is not sufficient for a full analysis. Only a dynamical theory can describe *e.g.* the “Red Queen” aspect of “Alice’s Wonderland”. (There, in the kingdom of the Red Queen one has to run to remain on the same spot.) Such phenomena reign in evolution as in the arms races between hosts and parasites or predators and prey.

Appendix A.4 Spatial games

Individuals are arranged on a **regular lattice** (with arbitrary dimension, but usually two or three dimensional) playing the game (a two-player game) with its immediate **neighbours**^{7,198}. Different neighbourhoods can be defined: the *von Neumann* or the *Moore* neighborhood¹⁹⁶, both using different distances. The **update** can be either synchronous or asynchronous⁷.

An example for one of the first spatial games is a spatial Prisoner’s Dilemma by Nowak and May¹⁹⁹. There, the payoffs from these interactions are added up for each individual. In the following round (*i.e.* generation) each player adopts the strategy with the highest payoff in its immediate neighborhood. All cells are updated in synchrony. This and other simple spatial games investigate deterministic evolutionary dynamics without random mutations.

Often, the **boundary conditions** are defined as periodic boundaries, but also other boundary conditions are possible (*e.g.* fixed boundaries). Different **initial settings** (*e.g.* initial random distribution of strategies) can be chosen. Furthermore, different lattice geometries can be used (*e.g.* triangular or hexagonal).

To analyze **invasion conditions**, it is useful to start with one mutant strategy among a homogeneous array. Another aim of exploring spatial games is to find interesting sequences of **patterns** (*e.g.* dynamic fractals, evolutionary kaleidoscopes, chaos)¹⁹⁹. Empty sites or more than two competing strategies can lead to spiral waves. For example, in the spatial Prisoner’s Dilemma, **coexistence** between cooperators and defectors can be observed¹⁹⁹. Cooperators survive in clusters and under cer-

tain conditions, cooperators can invade defectors. Often, spatial structure tends to allow more diversity than well-mixed populations do^{7,199}.

Appendix A.5 Evolutionary game dynamics

Appendix A.5.1 Replicator Dynamics. The replicator dynamics was described *e.g.* by Hofbauer and Sigmund⁶ and Page and Nowak²⁰⁰. It describes the evolution (dynamical changes) of the frequencies of **finite strategies** in a **heterogeneous population** by mimicking the effect of natural selection. Such dynamical changes of frequencies of **discrete** phenotypes in evolutionary game dynamics are described by the replicator equation:

$$\frac{dx_i}{dt} = x_i(f_i(\mathbf{x}) - F(\mathbf{x})) \quad (\text{A3})$$

where x_i denotes the frequency of strategy i , $f_i(\mathbf{x})$ denotes the (expected) payoff (fitness) for playing strategy i in a population with a distribution given by the vector $\mathbf{x} = (x_1, \dots, x_n)$ and $F(\mathbf{x})$ denotes the average (expected) payoff in an \mathbf{x} -population. The replicator equation is equivalent to the Lotka-Volterra equation^{200,201}. The replicator equation is directly computable from a two player matrix game⁶.

- Every Nash equilibrium of a matrix game is a rest point (=fixed point) of the replicator equation, but not every fixed point of the replicator equation is a Nash equilibrium.
- Every evolutionary stable state of a matrix game is an asymptotically stable fixed point of the replicator equation.

Appendix A.5.2 Adaptive Dynamics. Adaptive dynamics is based on calculations of the invasion fitness function for a minority mutant phenotype in the ecological environment generated by a resident phenotype^{197,202}. The approach of adaptive dynamics seeks to answer the question whether this mutant, which has a strategy close to the resident population, can invade. Therefore, one can compare this approach with the playing-the-field scenario proposed by Hamilton¹⁵. In contrast to the replicator equation one calculates the dynamical change of **continuous strategies** or traits^{6,200} under frequency dependent selection and mutation in a **homogeneous population**. Mutations are assumed to happen rarely⁶.

The approach considers a resident population which is invaded by mutants. The most fit mutant (in comparison to the resident population) is chosen by natural selection²⁰⁰. We assume that the whole population uses strategy \mathbf{x} and a mutant uses strategy $\mathbf{y} = \mathbf{x} + \mathbf{h}$ which is close to \mathbf{x} . $A(\mathbf{y}, \mathbf{x})$ denotes the total payoff obtained by an \mathbf{y} strategist against an \mathbf{x} -opponent. The relative advantage $W(\mathbf{h}, \mathbf{x})$ is calculated by

$A(\mathbf{y}, \mathbf{x}) - A(\mathbf{x}, \mathbf{x})$. The adaptive dynamics is then calculated by the differential equation

$$\frac{dx_i}{dt} = \frac{\partial}{\partial y_i} A(\mathbf{y}, \mathbf{x}) \quad (\text{A4})$$

evaluated at $\mathbf{y} = \mathbf{x}$ and $i = 1, \dots, n$. This vector (a gradient of $\mathbf{y} \rightarrow A(\mathbf{y}, \mathbf{x})$) points in the direction of the maximal increase of the fitness advantage of the mutant⁶. This is mimicking an evolutionary process.

Appendix B Types of games

Here we present an overview over all possible symmetric games with two players and two strategies. Similar classifications are proposed by Hauert²⁰³ and Stark²⁰⁴.

The payoff matrix of any symmetrical two-player game can be written as:

$$\begin{array}{c|cc} & c & d \\ \hline c & \begin{pmatrix} R & S \end{pmatrix} \\ d & \begin{pmatrix} T & P \end{pmatrix} \end{array} \quad (\text{B1})$$

The payoffs are termed according to the Prisoner's Dilemma, where they mean R -reward for mutual cooperation (strategy c), T -temptation to defect (strategy d), S -sucker's payoff and P -punishment for mutual defection. If we set $R > P$, the remaining two values (S and T) span a plane, where 12 different regions can be distinguished. This plane is depicted in figure B1. The character of a game does not change if rows and columns are permuted simultaneously. We now characterise the various games with an exemplary payoff.

1. Prisoner's Dilemma

$$\begin{pmatrix} 2 & 0 \\ 3 & 1 \end{pmatrix}$$

Cover story: Two suspects in a crime are put into separate cells. If they both confess, each will be sentenced to three years in prison. If only one of them confesses, he will be set free and used as a witness against the other, who will receive a sentence of five years. If neither confesses, they will both be convicted of a minor offence and spend one year in prison. The highest payoff represents the best output of this situation (to be set free) and 0 stands for the worst case (five years in prison).

Game properties: The dominant strategy is to defect. It is the only Nash equilibrium and evolutionarily stable strategy (ESS). This solution is Pareto inefficient.

2. Hawk-dove game (also: game of chicken, snowdrift game) (Leader II)

$$\begin{pmatrix} 2 & 1 \\ 3 & 0 \end{pmatrix}$$

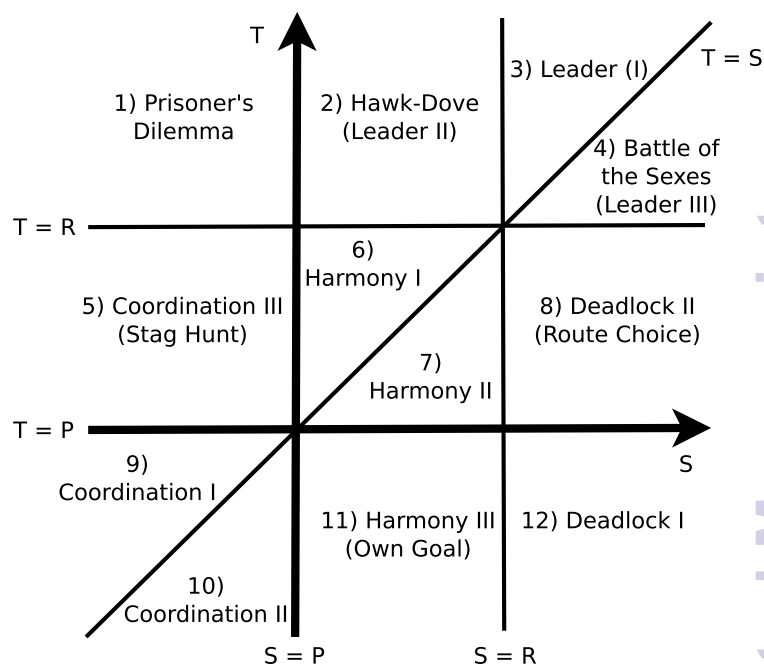


Fig. B1 Classification diagram for two-player games. A point in the diagram represents the complete relation of R , S , T and P . The diagram is read as the following example: Region 2 (cf. hawk-dove game) lies in the area above the diagonal $T = S$, therefore $T > S$ holds true there. Also, it is above the line $T = R$, therefore $T > R$ holds true. In addition, it is below the line $S = R$, so that $S < R$ holds true. Since $P < R$ holds true for the whole diagram, we can follow the order $T > R > S > P$ for region 2 (cf. hawk-dove game).

Cover stories:

Hawk-dove game The opponents are fighting for food, a territory or a potential mate. To be a hawk is an aggressive strategy (defect), while dove is to denote a peaceful (cooperate), surrendering strategy. Two hawks damage each other, two doves share the good (or one dove gets the whole good with probability 0.5). A dove meeting a hawk flees, no fitness gain nor loss, while the hawk gets the good.

Game of chicken, also Chicken-game Two cars are speeding towards each other on a collision. The loser must chicken out, or both suffer from a bad accident. If none of them stay on track, they stay unhurt but share the disgrace.

Snowdrift Two car drivers are stuck in a snowdrift. If both refuse to shovel, none of them will get home. If one of them shovels, while the other refuses to do so, both will get home. However, the shoveling player cooperates and pays a cost. If both shovel, both will get home, and share the cost of shoveling.

Game properties: The two pure Nash equilibria are on the secondary diagonal (to choose different strategies). The defector has the greater payoff. One mixed Nash equilibrium can be found, which is also an ESS.

3. Leader (I)

$$\begin{pmatrix} 1 & 2 \\ 3 & 0 \end{pmatrix}$$

Cover story: A strategic game in economics is played by the leader firm that moves first and by the follower firm that moves sequentially, while competing for quantity. The leader always has an advantage, similar to the hawk-dove game. However, here the cooperator has a lower disadvantage.

Game properties: The two pure Nash equilibria are on the secondary diagonal (to choose different strategies). The defector has the greater payoff. One mixed Nash equilibrium can be found, which is also an ESS. (cf. hawk-dove game)

4. Battle of the sexes (Leader III)

$$\begin{pmatrix} 1 & 3 \\ 2 & 0 \end{pmatrix}$$

Cover story: A couple wants to meet this evening. The husband prefers to visit the football game, while his wife would like to go to the opera. Both prefer to go out together rather than to different places. They are not able to communicate.

Originally this game is asymmetric (i.e. the payoffs are different, though both chose the same strategy), but it can be transformed in a symmetric one by renaming the strategies in e.g. “my preference” and “his/her preference”.

Game properties: The battle of the sexes has the same properties as the hawk-dove game, but the cooperator (in terms of the Prisoner’s Dilemma), the player applying “my preference”, has the greater payoff. The two pure Nash equilibria are on the secondary diagonal. One mixed Nash equilibrium can be found, which is also an ESS. (cf. hawk-dove game)

5. Stag Hunt (Coordination III)

$$\begin{pmatrix} 3 & 0 \\ 2 & 2(1) \end{pmatrix}$$

Cover story: Two huntsmen have the choice to hunt a stag or a hare. They can only hunt the stag cooperatively, and share the stag afterwards. Also, each one can hunt a hare on its own, but half a stag is worth more than a hare. If they are not able to communicate, which is the best strategy?

Game properties: The two pure Nash equilibria are on the main diagonal. One mixed Nash equilibrium can be found, which is *not* an ESS.

6. Harmony games

(a) Harmony I

$$\begin{pmatrix} 3 & 1 \\ 2 & 0 \end{pmatrix}$$

(b) Harmony II

$$\begin{pmatrix} 3 & 2 \\ 1 & 0 \end{pmatrix}$$

Game properties: Cooperation is the dominant strategy that leads to a pure Nash equilibrium where both players receive the highest possible payoff. Thus, they are “in harmony”. The difference between these two versions lies in the “strength” of the dominance of cooperation. In version II cooperation has the stronger dominance.

7. Route Choice (Deadlock II)

$$\begin{pmatrix} 2 & 3 \\ 1 & 0 \end{pmatrix}$$

Cover story: There are two routes with different capacities: a freeway and a side route. If both players decide to drive the freeway (cooperate), they can drive fast, but have to consider the other car, while a single car on the freeway can speed up. A single car on the side route is still quick, while two cars on the side route hinder each other.

Game properties: Cooperation is the dominant strategy. It exists only one Nash equilibrium with the second highest possible payoff. A player has the interest to encourage the opponent to play the dominated strategy.

8. Coordination games

(a) Coordination game I

$$\begin{pmatrix} 3 & 0 \\ 1 & 2 \end{pmatrix}$$

(b) Coordination game II

$$\begin{pmatrix} 3 & 1 \\ 0 & 2 \end{pmatrix}$$

Game properties: The two players have to coordinate their actions (i.e. play the same strategies) to receive a high payoff. The two pure Nash equilibria are on the main diagonal. Thus, the system shows bistability. One mixed Nash equilibrium can be found, which is NOT an ESS.

9. Own goal (Harmony III)

$$\begin{pmatrix} 3 & 2 \\ 0 & 1 \end{pmatrix}$$

Game properties: Cooperation is the dominant strategy. It exists only one Nash equilibrium with the highest possible payoff. An unilateral deviation of one player from the dominant strategy would be an “own goal” for them because the loss is very high.

10. Deadlock (I)

$$\begin{pmatrix} 2 & 3 \\ 0 & 1 \end{pmatrix}$$

In terms of the cover story of the route-choice game, one may say that when both drivers choose the side road, they are better off (for example, because they can help each other in case if an accident) than a driver that goes alone on the side road.

Game properties: Cooperation is the dominant strategy. It exists only one Nash equilibrium with the second highest possible payoff. A player has some interest to encourage the opponent to play the dominated strategy.

The game properties are summarised with the classification tree in figure B2.

Appendix C Rock-Scissors-Paper Game

This popular children’s game, where rock is wrapped by paper, paper is cut by scissors and scissors are crushed by rock can also be described by a game theoretical model. In a biological context the strategies are renamed to producer (P), resistant (R) and sensitive (S), see chapter 5. Consider the following payoff matrix:

$$\begin{array}{c} \begin{array}{ccc} & \text{P} & \text{R} & \text{S} \\ \text{P} & \begin{pmatrix} 0 & -a & b \\ a & 0 & -c \\ -b & c & 0 \end{pmatrix} & & \\ \text{R} & & & \\ \text{S} & & & \end{array} \end{array}, \quad (\text{C1})$$

where a, b and c represent the payoffs.

There is no Nash equilibrium in pure strategies, because for each entry of the matrix one of the players can increase payoff by changing strategy. For example, if a producer plays against a resistant strain, the producer can improve by switching to any other strategy. A mixed Nash equilibrium can be found by applying the Bishop-Cannings-Theorem (see Appendix A) with the following distribution of strategies $p = a/(a+b+c), r = b/(a+b+c), s = c/(a+b+c)$. In this biological context, the distribution can be interpreted as the corresponding fractions of the population. In the special case of $a = b = c$, an equal distribution of one third is obtained as is known from the hand game of the same name. However, this

mixed Nash equilibrium is not evolutionarily stable. That is why it is interesting to study the dynamics of this game. Different dynamical approaches are reviewed in Appendix A.

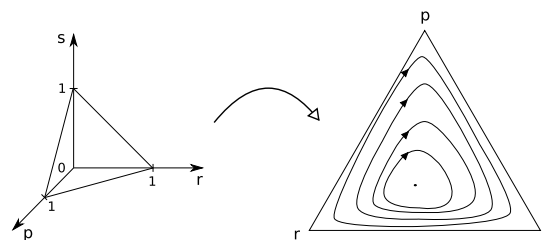


Fig. C Simplex representation. (a) Simplex in the three dimensional Cartesian coordinate system, (b) It is sufficient to visualise the simplex without the coordinate system. Here, periodic orbits around a fixed point are shown, similar to⁸². Such cyclic orbits correspond to oscillating sub-populations.

To illustrate the dynamics of a three species population, often the simplex, an equilateral triangle, is used. It represents the plane from the three dimensional Cartesian coordinate system with $p + r + s = 1$, where each fraction stands for one subpopulation, see Figure C.

References

- 1 Y. K. Lee and S. K. Mazmanian, *Science*, 2010, **330**, 1768–1773.
- 2 D. J. Schneider and A. Collmer, *Annu. Rev. Phytopathol.*, 2010, **48**, 457–479.
- 3 M. Ebner, R. Watson and J. Alexander, In: Proceedings of the 2000 Congress on Evolutionary Computation, San Diego, IEEE Press, 2000, pp. 1284–1291.
- 4 J. Maynard-Smith, *Evolution and the Theory of Games*, Cambridge University Press, Cambridge, 1982.
- 5 R. Axelrod, *The Evolution of Cooperation*, Basic Books, New York, 1984.
- 6 J. Hofbauer and K. Sigmund, *Evolutionary Games and Replicator Dynamics*, Cambridge University Press, Cambridge, 1998.
- 7 M. A. Nowak and K. Sigmund, *Science*, 2004, **303**, 793–799.
- 8 J. V. Neumann and O. Morgenstern, *Theory of Games and Economic Behaviour*, Princeton University Press, 1944.
- 9 E. Berne, *Games People Play: The Psychology of Human Relationships*, Penguin, 1973.
- 10 R. B. Myerson, *Game Theory: Analysis of Conflict*, Harvard University Press, 1991.
- 11 M. Eigen and R. Winkler, *Das Spiel. Naturgesetze steuern den Zufall*, Pieper, München, 1975.
- 12 K. Bohl, S. Hummert, S. Werner, D. Basanta, A. Deutsch, S. Schuster, G. Theißen and A. Schröter, *Mol. BioSyst. submitted*, 2013.
- 13 J. F. Nash, *Proc. Natl. Acad. Sci. U.S.A.*, 1950, **36**, 48–49.
- 14 J. Maynard-Smith and G. R. Price, *Nature*, 1973, **246**, 15–18.
- 15 W. D. Hamilton, *Science*, 1967, **156**, 477–488.
- 16 J. M. Smith and G. R. Price, *Nature*, 1973, **246**, 15–18.
- 17 K. C. Clements and D. W. Stephens, *Anim. Behav.*, 1995, **50**, 527–535.
- 18 M. Broom and G. Ruxton, *Behav. Ecol.*, 1998, **9**, 397–403.
- 19 J. Brown and T. Vincent, *Evolution*, 1992, **46**, 1269–1283.
- 20 D. W. Dickens and R. A. Clark, *J. Theor. Biol.*, 1987, **125**, 301–305.

- 21 Poulin and Vickery, *J. Theor. Biol.*, 1995, **175**, 63–70.
- 22 D. King, *Am. Nat.*, 1990, **135**, 809–828.
- 23 M. Augner, T. Fagerstrom and J. Tuomi, *Behav. Ecol. Sociobiol.*, 1991, **29**, 231–234.
- 24 M. Archetti, *J. Theor. Biol.*, 2000, **205**, 625–630.
- 25 J.-U. Kreft, *Microbiology*, 2004, **150**, 2751–2760.
- 26 T. Pfeiffer and S. Bonhoeffer, *Am. Nat.*, 2004, **163**, E126–35.
- 27 E. Costa, J. Pérez and J. U. Kreft, *Trends Microbiol.*, 2006, **14**, 213–219.
- 28 S. P. Brown and R. A. Johnstone, *Proc. Biol. Sci.*, 2001, **268**, 961–965.
- 29 M. Pagel, *Nature*, 1993, **363**, 539–541.
- 30 R. A. Gatenby and T. L. Vincent, *Cancer Res.*, 2003, **63**, 6212–6220.
- 31 D. Basanta, H. Hatzikirou and A. Deutsch, *Eur. Phys. J B*, 2008, **63**, 393–397.
- 32 T. Pfeiffer, S. Schuster and S. Bonhoeffer, *Science*, 2001, **292**, 504–507.
- 33 D. Voet and J. Voet, *Biochemistry*, John Wiley & Sons, 2011.
- 34 Z. Wang and N. Goldenfeld, *Phys. Rev. E*, 2011, **84**, 020902.
- 35 J. C. Aledo, J. A. Perez-Claros and et al., *J. Mol. Evol.*, 2007, **65**, 328–339.
- 36 J. Juul, K. Sneppen and J. Mathiesen, *Phys. Rev. E*, 2012, **85**, 061924.
- 37 R. C. MacLean, *Heredity*, 2007, **100**, 471–477.
- 38 I. A. Shiklomanov, *Water Int.*, 2000, **25**, 11–32.
- 39 J. Y. Wakano, M. A. Nowak and et al., *Proc. Natl. Acad. Sci. U.S.A.*, 2009, **106**, 7910–7914.
- 40 T. Frick and S. Schuster, *Naturwissenschaften*, 2003, **90**, 327–331.
- 41 S. Schuster, J. U. Kreft, A. Schroeter and T. Pfeiffer, *J. Biol. Phys.*, 2008, **34**, 1–17.
- 42 R. C. MacLean and I. Gudelj, *Nature*, 2006, **441**, 498–501.
- 43 O. Warburg, *Science*, 1956, **123**, 309–314.
- 44 S. Schuster, L. F. de Figueiredo, A. Schroeter and C. Kaleta, *BioSystems*, 2011, **105**, 147–153.
- 45 I. Kareva, *PLoS One*, 2011, **6**, e28576.
- 46 S. A. Frank, *J. Evol. Biol.*, 2010, **23**, 609–613.
- 47 S. A. Frank, *Phil. Trans. R. Soc. B*, 2010, **365**, 2515–2522.
- 48 L. A. Harriman and B. W. Hammer, *J. Dairy Sci.*, 1931, **14**, 40–49.
- 49 D. Greig and M. Travisano, *Proc. Biol. Sci.*, 2004, **271**, S25–S26.
- 50 J. Gore, H. Youk and et al., *Nature*, 2009, **459**, 253–256.
- 51 S. Schuster, J. U. Kreft and et al., *Biotechnol. J.*, 2008, **5**, 751–758.
- 52 S. D. Allison, *Ecol. Lett.*, 2005, **8**, 626–635.
- 53 H. Bachmann, D. Molenaar, M. Kleerebezem and J. E. van Hylckama Vlieg, *ISME J.*, 2011, **5**, 929–932.
- 54 R. C. MacLean, A. Fuentes-Hernandez, D. Greig, L. D. Hurst and I. Gudelj, *PLoS Biol.*, 2010, **8**, e1000486.
- 55 S. Estrela, C. H. Trisos and S. P. Brown, *Am. Nat.*, 2012, **180**, 566–576.
- 56 H. C. Bernstein, S. D. Paulson and R. P. Carlson, *J. Biotechnol.*, 2012, **157**, 159–166.
- 57 B. E. Morris, R. Henneberger, H. Huber and C. Moissl-Eichinger, *FEMS Microbiol. Rev.*, 2013, **37**, 384–406.
- 58 M. Doebeli, *Population Ecol.*, 2002, **44**, 59–70.
- 59 J. J. Bull and W. R. Harcombe, *PLoS One*, 2009, **4**, e4115.
- 60 B. Schink, *Antonie van Leeuwenhoek*, 2002, **81**, 257–261.
- 61 C. Katsuyama, S. Nakaoka, Y. Takeuchi, K. Tago, M. Hayatsu and K. Kato, *J. Theor. Biol.*, 2009, **256**, 644–654.
- 62 D. E. Rozen and R. E. Lenski, *Am. Nat.*, 2000, **155**, 24–35.
- 63 P. Gerlee and T. Lundh, *Evolution*, 2010, **64**, 2716–2730.
- 64 J. M. Smith, *Nature*, 1964, **201**, 1145–1147.
- 65 S. Estrela and I. Gudelj, *PLoS One*, 2010, **5**, e14121.
- 66 M. L. Friesen and E. I. Jones, *Methods in Molecular Biology*, Springer New York, 2012, vol. 804, pp. 481–499.
- 67 M. Archetti and I. Scheuring, *J. Theor. Biol.*, 2012, **318C**, 58–67.
- 68 M. Archetti, I. Scheuring, M. Hoffman, M. E. Frederickson, N. E. Pierce and D. W. Yu, *Ecol. Lett.*, 2011, **14**, 1300–1312.
- 69 J. B. Jackson and L. Buss, *Proc. Natl. Acad. Sci. U.S.A.*, 1975, **72**, 5160–5163.
- 70 B. Sinervo and C. M. Lively, *Nature*, 1996, **380**, 240–243.
- 71 F. Michor and M. A. Nowak, *Nature*, 2002, **419**, 677, 679.
- 72 B. Kerr, M. A. Riley, M. W. Feldman and B. M. Bohannan, *Nature*, 2002, **418**, 171–174.
- 73 B. C. Kirkup and M. A. Riley, *Nature*, 2004, **428**, 412–414.
- 74 R. Durrett and S. Levin, *J. Theor. Biol.*, 1997, **185**, 165–171.
- 75 G. Neumann and S. Schuster, *J. Math. Biol.*, 2007, **54**, 815–846.
- 76 G. Neumann and S. Schuster, *Discrete Cont. Dyn.-B*, 2007, **8**, 207–228.
- 77 G. F. Neumann and G. Jetschke, *BioSystems*, 2010, **99**, 155–166.
- 78 M. Nakamaru and Y. Iwasa, *Theor. Popul. Biol.*, 2000, **57**, 131–144.
- 79 M. W. Adamson and A. Y. Morozov, *Bull. Math. Biol.*, 2012, **74**, 2004–2031.
- 80 R. M. May and W. J. Leonard, *Siam J. Appl. Math.*, 1975, **29**, 243–253.
- 81 H. Shi, W.-X. Wang, R. Yang and Y.-C. Lai, *Phys. Rev. E*, 2010, **81**, 030901.
- 82 M. Frean and E. R. Abraham, *Proc. Biol. Sci.*, 2001, **268**, 1323–1327.
- 83 G. Károlyi, Z. Neufeld and I. Scheuring, *J. Theor. Biol.*, 2005, **236**, 12–20.
- 84 M. Berr, T. Reichenbach, M. Schottenloher and E. Frey, *Phys. Rev. Lett.*, 2009, **102**, 048102.
- 85 W.-X. Wang, Y.-C. Lai and C. Grebogi, *Phys. Rev. E*, 2010, **81**, 046113.
- 86 X. Ni, R. Yang, W.-X. Wang, Y.-C. Lai and C. Grebogi, *Chaos*, 2010, **20**, 045116.
- 87 A. P. O. Müller and J. A. C. Gallas, *Phys. Rev. E*, 2010, **82**, 052901.
- 88 T. L. Czárán, R. F. Hoekstra and L. Pagie, *Proc. Natl. Acad. Sci. U.S.A.*, 2002, **99**, 786–790.
- 89 F. Prado and B. Kerr, *Evolution*, 2008, **62**, 538–548.
- 90 A. A. Winkler, T. Reichenbach and E. Frey, *Phys. Rev. E*, 2010, **81**, 060901.
- 91 T. Reichenbach, M. Mobilia and E. Frey, *Nature*, 2007, **448**, 1046–1049.
- 92 T. Reichenbach, M. Mobilia and E. Frey, *J. Theor. Biol.*, 2008, **254**, 368–383.
- 93 S. Venkat and M. Pleimling, *Phys. Rev. E*, 2010, **81**, 021917.
- 94 A. Szolnoki and G. Szabó, *Phys. Rev. E*, 2004, **70**, 037102.
- 95 G. Szabó, A. Szolnoki and R. Izsák, *J. Phys. A: Math. Gen.*, 2004, **380**, 2599–2609.
- 96 K.-i. Tainaka, *Phys. Rev. E*, 1994, **50**, 3401–3409.
- 97 M. Ifiti and B. Bergersen, *Eur. Phys. J. B*, 2004, **37**, 101–107.
- 98 A. Traulsen, J. C. Claussen and C. Hauert, *Phys. Rev. E*, 2012, **85**, 041901.
- 99 J. C. Claussen and A. Traulsen, *Phys. Rev. Lett.*, 2008, **100**, 058104.
- 100 Frachebourg, Krapivsky and Ben-Naim, *Phys. Rev. E*, 1996, **54**, 6186–6200.
- 101 P. P. Avelino, D. Bazeia, L. Losano and J. Menezes, *Phys. Rev. E*, 2012, **86**, 031119.
- 102 P. P. Avelino, D. Bazeia, L. Losano, J. Menezes and B. F. Oliveira, *Phys. Rev. E*, 2012, **86**, 036112.
- 103 A. F. Lütz, S. Risau-Gusman and J. J. Arenzon, *J. Theor. Biol.*, 2012, **317C**, 286–292.
- 104 M. Peltomäki and M. Alava, *Phys. Rev. E*, 2008, **78**, 031906.
- 105 G. Szabó and T. Czárán, *Phys. Rev. E*, 2001, **64**, 042902.
- 106 B. Allen and D. I. S. Rosenbloom, *Bull. Math. Biol.*, 2012, **74**, 2650–2675.
- 107 Q. He, M. Mobilia and U. C. Täuber, *Phys. Rev. E*, 2010, **82**, 051909.
- 108 G. Jäger, *J. Theor. Biol.*, 2008, **253**, 131–141.
- 109 R. Gibbons, *A primer in game theory*, Financial Times Prentice Hall, 1992.
- 110 E. Rasmusen, *Games and information: an introduction to game theory*, Wiley-Blackwell, 1995.

- 111 J. Maynard-Smith and D. Harper, *Animal Signals*, Oxford University Press, 2003.
- 112 A. Zahavi, *J. Theor. Biol.*, 1975, **53**, 205–214.
- 113 A. Grafen, *J. Theor. Biol.*, 1990, **144**, 517–546.
- 114 C. T. Bergstrom, S. Számádó and M. Lachmann, *Phil. Trans. R. Soc. B*, 2002, **357**, 1595–1606.
- 115 R. Popat, S. A. Crusz, M. Messina, P. Williams, S. A. West and S. P. Diggle, *Proc. Biol. Sci.*, 2012, **279**, 4765–4771.
- 116 T. Czárán and R. F. Hoekstra, *PLoS One*, 2009, **4**, e6655.
- 117 A. Lotem, M. A. Fishman and L. Stone, *Proc. Biol. Sci.*, 2003, **270**, 199–205.
- 118 R. Axelrod and W. D. Hamilton, *Science*, 1981, **211**, 1390–1396.
- 119 A. Traulsen and M. A. Nowak, *PLoS One*, 2007, **2**, e270.
- 120 D. C. Queller, E. Ponte, S. Bozzaro and J. E. Strassmann, *Science*, 2003, **299**, 105–106.
- 121 D. Renaud and T. de Meeüs, *J. Theor. Biol.*, 1991, **152**, 319–327.
- 122 M. R. Frean and E. R. Abraham, *Phys. Rev. E*, 2004, **69**, 051913.
- 123 S.-H. Huang, W. Zhou and A. Jong, *J. Biomed. Biotechnol.*, 2008, 856314.
- 124 E. M. Rauch and Y. Bar-Yam, *Phys. Rev. E*, 2006, **73**, 020903.
- 125 R. M. Anderson and R. M. May, *Nature*, 1979, **280**, 361–367.
- 126 R. M. Anderson and R. M. May, *Parasitology*, 1982, **85**, 411–426.
- 127 H. J. Bremermann and J. Pickering, *J. Theor. Biol.*, 1983, **100**, 411–426.
- 128 M. van Baalen, *Proc. Biol. Sci.*, 1998, **265**, 317–325.
- 129 T. Day, A. L. Graham and A. F. Read, *Proc. Biol. Sci.*, 2007, **274**, 2685–2692.
- 130 M. Boots and Y. Haraguchi, *Am. Nat.*, 1999, **153**, 359–370.
- 131 M. H. Bonds, D. C. Keenan, A. J. Leidner and P. Rohani, *Evolution*, 2005, **59**, 1859–1866.
- 132 M. R. Miller, A. White and M. Boots, *Evolution*, 2007, **61**, 2–14.
- 133 A. Best, A. White and M. Boots, *Proc. Natl. Acad. Sci. U.S.A.*, 2008, **105**, 20786–20791.
- 134 P. D. Taylor, T. Day, D. Nagy, J.-B. Wild, Geoff and André and A. Gardner, *Theor. Popul. Biol.*, 2006, **69**, 323–331.
- 135 M. Boots, A. Best, M. M. R. and A. White, *Phil. Trans. R. Soc. B*, 2009, **364**, 27–36.
- 136 A. Morozov and A. Best, *J. Theor. Biol.*, 2012, **307**, 29–36.
- 137 S. H. Alonzo and R. Calsbeek, *J. Evol. Biol.*, 2010, **23**, 2614–2624.
- 138 T. Reluga and J. Li, *J. Math. Biol.*, 2012, **66**, 1–27.
- 139 P. D. Williams, *Am. Nat.*, 2012, **179**, 228–239.
- 140 D. M. Wolf, V. V. Vazirani and A. P. Arkin, *J. Theor. Biol.*, 2005, **234**, 227–253.
- 141 D. M. Wolf, V. V. Vazirani and A. P. Arkin, *J. Theor. Biol.*, 2005, **234**, 255–262.
- 142 B. L. d. Bivort, C.-C. Chen, F. Perretti, G. Negro, T. M. Philip and Y. Bar-Yam, *J. Theor. Biol.*, 2007, **248**, 26–36.
- 143 S. M. Eswarappa, *PLoS One*, 2009, **4**, e5383.
- 144 S. Hummert, C. Hummert, A. Schröter, B. Hube and S. Schuster, *J. Theor. Biol.*, 2010, **264**, 312–318.
- 145 S. Bewick, J. Wu, S. C. Lenaghan, R. Yang, M. Zhang and W. Hamel, *Bull. Math. Biol.*, 2011, **73**, 2339–2356.
- 146 S. Bewick, M. Zhang and W. Hamel, *IEEE Trans. Syst. Man Cybernetics A: Syst. Humans*, 2011, **41**, 583–588.
- 147 B. Crespi and K. Summers, *Trends Ecol. Evol.*, 2005, **20**, 545–552.
- 148 L. M. F. Merlo, J. W. Pepper, B. J. Reid and C. C. Maley, *Nat. Rev. Cancer*, 2006, **6**, 924–935.
- 149 P. C. Nowell, *Science*, 1976, **194**, 23–28.
- 150 M. Gerlinger, A. J. Rowan, S. Horswell, J. Larkin, D. Endesfelder, E. Gronroos, P. Martinez, N. Matthews, A. Stewart, P. Tarpey, I. Varela, B. Phillimore, S. Begum, N. Q. McDonald, A. Butler, D. Jones, K. Raine, C. Latimer, C. R. Santos, M. Nohadani, A. C. Eklund, B. Spencer-Dene, G. Clark, L. Pickering, G. Stamp, M. Gore, Z. Szalasi, J. Downward, P. A. Futreal and C. Swanton, *N. Engl. J. Med.*, 2012, **366**, 883–892.
- 151 R. Axelrod, D. E. Axelrod and K. J. Pienta, *Proc. Natl. Acad. Sci. U.S.A.*, 2006, **103**, 13474–13479.
- 152 R. J. Gillies, D. Verduzco and R. A. Gatenby, *Nat. Rev. Cancer*, 2012, **12**, 487–493.
- 153 *Selected Topics in Cancer Modeling: Genesis, Evolution, Immune Competition, and Therapy*, ed. N. Bellomo, and E. de Angelis, Springer, Birkhäuser Boston, 2008, ch. 5.
- 154 C. Cleveland, D. Liao and R. Austin, *AIP Adv.*, 2012, **2**, 011202.
- 155 W. C. Hahn and R. A. Weinberg, *N. Engl. J. Med.*, 2002, **347**, 1593–1603.
- 156 D. Hanahan and R. A. Weinberg, *Cell*, 2000, **100**, 57–70.
- 157 D. Hanahan and R. A. Weinberg, *Cell*, 2011, **144**, 646–674.
- 158 A. R. A. Anderson, M. Hassanein, K. M. Branch, J. Lu, N. A. Lobdell, J. Maier, D. Basanta, B. Weidow, A. Narasanna, C. L. Arteaga, A. B. Reynolds, V. Quaranta, L. Estrada and A. M. Weaver, *Cancer Res.*, 2009, **69**, 8797–8806.
- 159 Y. Mansury, M. Diggory and T. Deisboeck, *J. Theor. Biol.*, 2006, **238**, 146–156.
- 160 M. Martin, *J. Natl. Cancer Inst.*, 2009, **101**, 220–222.
- 161 J. M. Smith, *Evolution and the Theory of Games*, Cambridge University Press, Cambridge, 1982.
- 162 R. A. Gatenby and R. J. Gillies, *Nat. Rev. Cancer*, 2004, **4**, 891–899.
- 163 D. Basanta and A. Deutsch, *Selected Topics in Cancer Modeling*, Birkhäuser Boston, 2008, pp. 1–16.
- 164 D. Basanta, J. G. Scott, R. Rockne, K. R. Swanson and A. R. A. Anderson, *Phys. Biol.*, 2011, **8**, 015016.
- 165 D. Basanta, M. Simon, H. Hatzikirou and A. Deutsch, *Cell Prolif.*, 2008, **41**, 980–987.
- 166 M. Archetti, *J. Theor. Biol.*, 2014, **341**, 1–8.
- 167 M. A. Nowak, *Science*, 2006, **314**, 1560–1563.
- 168 D. W. Strand, O. E. Franco, D. Basanta, A. R. A. Anderson and S. W. Hayward, *Curr. Mol. Med.*, 2010, **10**, 95–112.
- 169 I. P. Tomlinson and W. F. Bodmer, *Br. J. Cancer*, 1997, **75**, 157–160.
- 170 L. A. Bach, S. M. Bentzen, J. Alsner and F. B. Christiansen, *Eur. J. Cancer*, 2001, **37**, 2116–2120.
- 171 L. A. Bach, D. J. T. Sumpter, J. Alsner and V. Loeschcke, *J. Theor. Med.*, 2003, **5**, 47–58.
- 172 M. Billaud and M. Santoro, *Cancer Res.*, 2011, **71**, 6572–6575.
- 173 D. Dingli, F. A. C. C. Chalub, F. C. Santos, S. V. Segbroeck and J. M. Pacheco, *Br. J. Cancer*, 2009, **101**, 1130–1136.
- 174 D. Basanta, J. G. Scott, M. N. Fishman, G. Ayala, S. W. Hayward and A. R. A. Anderson, *Br. J. Cancer*, 2012, **106**, 174–181.
- 175 R. A. Gatenby, J. Brown and T. Vincent, *Cancer Res.*, 2009, **69**, 7499–7502.
- 176 A. S. Silva, R. A. Gatenby, R. J. Gillies and J. A. Yunes, *J. Theor. Biol.*, 2010, **262**, 601–613.
- 177 S. J. Antonia, N. Mirza, I. Fricke, A. Chiappori, P. Thompson, N. Williams, G. Bepler, G. Simon, W. Janssen, J.-H. Lee, K. Menander, S. Chada and D. I. Gabrilovich, *Clin. Cancer Res.*, 2006, **12**, 878–887.
- 178 D. Basanta, R. A. Gatenby and A. R. A. Anderson, *Mol. Pharmacol.*, 2012, **9**, 914–921.
- 179 S. Bewick, R. Yang and M. Zhang, Engineering in Medicine and Biology Society, 2009. EMBC 2009. Annual International Conference of the IEEE, 2009, pp. 6026–9.
- 180 J. Wu and M. J. Zhang, *IEEE Trans. Syst. Man Cybernetics B: Cybernetics*, 2010, **40**, 694–702.
- 181 A. Ostman, *Nat. Med.*, 2012, **18**, 1332–1334.
- 182 V. Marx, *Nature*, 2013, **494**, 133–136.
- 183 G. Lambert, L. Estévez-Salmeron, S. Oh, D. Liao, B. M. Emerson, T. D. Tlsty and R. H. Austin, *Nat. Rev. Cancer*, 2011, **11**, 375–382.

- 184 O. Gillor, A. Etzion and M. A. Riley, *Appl. Microbiol. Biotechnol.*, 2008, **81**, 591–606.
- 185 P. D. Cotter, R. P. Ross and C. Hill, *Nat. Rev. Microbiol.*, 2013, **11**, 95–105.
- 186 C. T. Lohans and J. C. Vederas, *Int. J. Microbiol.*, 2012, **2012**, 386410.
- 187 M. Montalbán-López, M. Sánchez-Hidalgo, E. Valdivia, M. Martínez-Bueno and M. Maqueda, *Curr. Pharm. Biotechnol.*, 2011, **12**, 1205–1220.
- 188 C. Brown, M. P. Garwood and J. E. Williamson, *Biol. Lett.*, 2012, **8**, 729–732.
- 189 M. R. Silva, M. G. Severo, P. S. Delabona, R. Ruller, J. G. C. Pradella, V. M. Goncalves and S. Freitas, *Biochem. Eng. J.*, 2013, **71**, 38–46.
- 190 G. Potvin, A. Ahmad and Z. S. Zhang, *Biochem. Eng. J.*, 2012, **64**, 91–105.
- 191 T. Pfeiffer and S. Bonhoeffer, *Proc. Natl. Acad. Sci. U.S.A.*, 2003, **100**, 1095–1098.
- 192 H. Cao, M. Sekiya, M. Ertunc, M. Burak, J. Mayers, A. White, K. Inouye, L. Rickey, B. Ercal, M. Furuhashi, G. Tuncman and G. Hotamisligil, *Cell Metab.*, 2013, **17**, 768–778.
- 193 A. Szolnoki and M. Perc, *Phys. Rev. E*, 2012, **85**, 026104.
- 194 H. Gintis, *Game theory evolving*, Princeton University Press, Princeton, 2000.
- 195 D. T. Bishop and C. Cannings, *J. Theor. Biol.*, 1978, **70**, 85–124.
- 196 M. A. Nowak, *Evolutionary dynamics - Exploring the equations of life*, Harvard University Press, 2006.
- 197 J. Hofbauer and K. Sigmund, *Bull. Am. Math. Soc.*, 2003, **40**, 479–519.
- 198 M. A. Nowak, C. E. Tarnita and T. Antal, *Phil. Trans. R. Soc. B*, 2010, **365**, 19–30.
- 199 M. A. Nowak and R. M. May, *Nature*, 1992, **359**, 826–829.
- 200 K. M. Page and M. A. Nowak, *J. Theor. Biol.*, 2002, **219**, 93–98.
- 201 J. Hofbauer, *Nonlinear Anal. Theor. Meth. App.*, 1981, **5**, 1003–1007.
- 202 O. Diekmann, *Banach Center Publ.*, 2004, **63**, 47–86.
- 203 C. Hauert, *Int. J. Bifurcation Chaos*, 2002, **12**, 1531–1548.
- 204 H.-U. Stark, *Evolution*, 2010, **64**, 2458–2465.

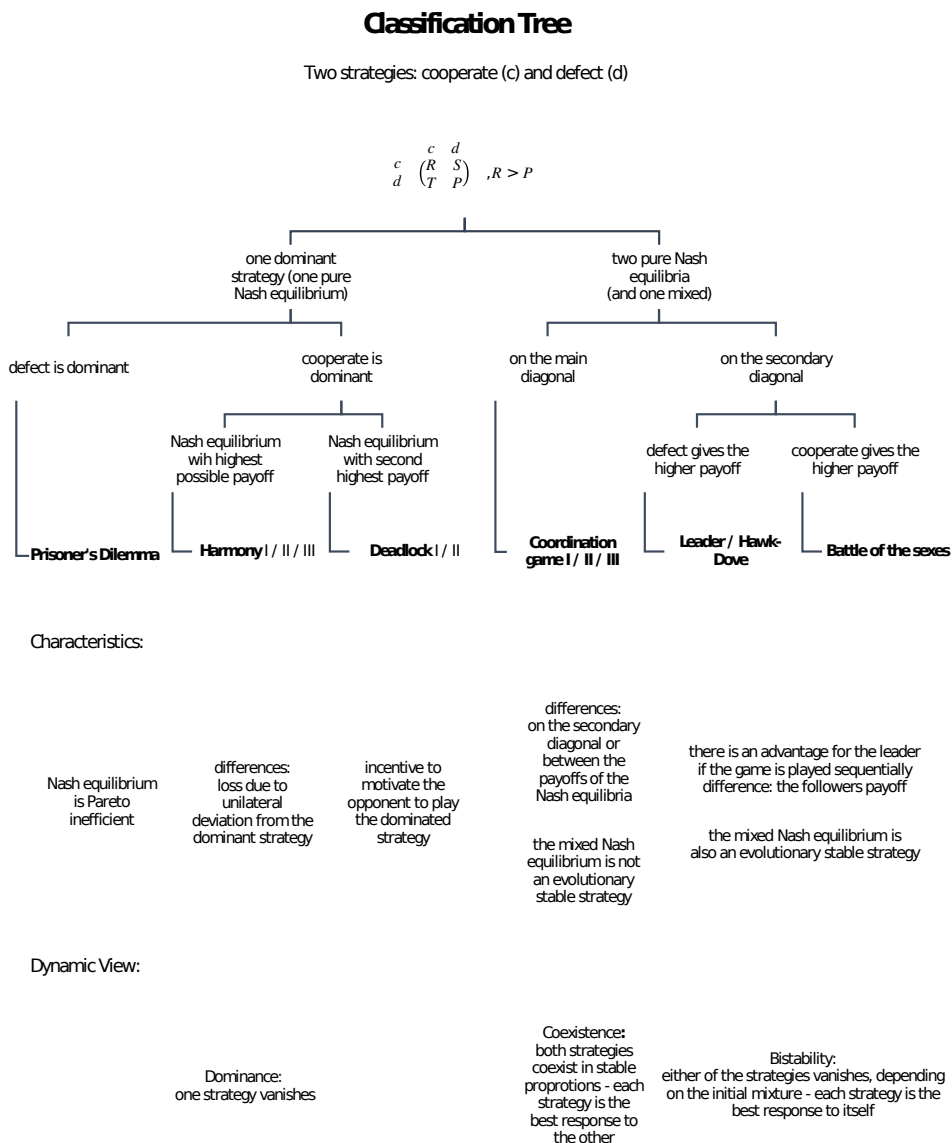


Fig. B2 Classification tree for two-player games summarising game properties. The dynamic is characterized as in Nowak and Sigmund⁷.