Cooperation in microbial communities and their biotechnological applications

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Statement of significance:
In this review we summarise advances in the field of Evolutionary Dynamics applied to microbial communities and their applications in biotechnology. We discuss different kinds of cooperative interactions, their potential mechanistic origins and the factors that contribute to their stability. We also analyse the advantages of cooperative behaviours in microbial populations and evaluate their possible use to develop robust biotechnological applications.
Abstract
Microbial communities are increasingly utilised in biotechnology. Efficiency and productivity in many of these applications depends on the presence of cooperative interactions between members of the community. Two key processes underlying these interactions are the production of public goods and metabolic crossfeeding, which can be understood in the general framework of ecological and evolutionary (eco-evo) dynamics. In this review we illustrate the relevance of cooperative interactions in microbial biotechnological processes, discuss their mechanistic origins, and analyse their evolutionary resilience. Cooperative behaviours can be damaged by the emergence of ‘cheating’ cells that benefit from the cooperative interactions but do not contribute to them. Despite this, cooperative interactions can be stabilized by spatial segregation, by the presence of feedbacks between the evolutionary dynamics and the ecology of the community, by the role of regulatory systems coupled to the environmental conditions and by the action of horizontal gene transfer. Cooperative interactions enrich microbial communities with a higher degree of robustness against environmental stress and can facilitate the evolution of more complex traits. Therefore, the evolutionary resilience of microbial communities and their ability to constraint detrimental mutants should be considered in order to design robust biotechnological applications.
Evolutionary dynamics and cooperation in microbial populations

The design and optimization of microorganisms for biotechnological purposes often considers cells in isolation. While this reductionist approach aims to thrive for simplicity in the process, it creates a situation that rarely takes place in Nature. In their natural environment microorganisms thrive in complex communities in which the fitness of a single cell depends on the interactions with other cells in the population (West et al., 2006). This scenario also applies to bioproceses in which the efficiency of the process is coupled to the production of shared (public) goods that allow cells to perform tasks in a ‘cooperative’ manner (Lindemann et al., 2016): a good example of shared goods are the cellulases secreted in the production of cellulosic ethanol (Zomorrodi and Segrè, 2016).

The presence of cooperative interactions has a significant impact on the evolutionary dynamics of microbial communities, represented by the change in the frequencies of cells and species that implement different physiological strategies (such as production of public goods vs. not). Thus, cooperative traits need to be taken into account when using an evolutionary approach for optimising a given bioprocess. It is possible that simple selection schemes targeting a bioprocess-related trait (e.g. growth rate) will not align with the selection for the cooperative trait (e.g. production of costly extracellular enzymes) ultimately resulting in the loss of the trait. Indeed, tradeoffs between the optimization of so-called high-rate and high-yield are frequently observed in controlled evolutionary experiments (Bachmann et al., 2013). Thus, we advocate considering the interactions between the cells and the functioning of cooperative traits when designing evolutionary optimisation and stabilisation of bioprocesses. Achieving this would require considering how ‘social’ interactions shape microbial processes, rather than simply focusing solely on individualistic traits such as growth rate.

This situation may confront the intuitive idea that ‘evolution implies improvement’ (i.e. the average fitness of the community is expected to increase over generations as it would be expected for monocultures). The key point is that the presence of interactions between the species gives rise to a more complicated evolutionary picture in which the fitness of a cell depends not only on its phenotype but also on the overall composition of the population. The spreading of a given phenotypic trait may thus change the fitness of other members of the community and these changes may in turn feedback on the fitness of the individual cells (West et al., 2006). These intertwined selection mechanisms are expected to operate in any microbial population where there is possibility of different cells implementing different strategies with respect to their physiology, as is the case of phenotypic heterogeneity.

Phenotypic heterogeneity arises even in monocultures and simple bioproceses due to different reasons, such as the use of non-homogenous culture conditions, stochasticity in gene
expression and differential epigenetic control (Enfors et al., 2001; Avery, 2006; Müller et al., 2010). Such heterogeneity does not represent a static picture – cells communicate, compete and cooperate and the success of a trait may be consequence of the interaction with the other traits and of the specific ecological context (Carlquist et al., 2012). Therefore, it is not sufficient for a trait to be successful in one specific setting but rather, it needs to be successful given the presence of other traits and the associated ecological context. Moreover, the dilution of a trait may lead to changes in the community (both ecological and/or in the frequency of other traits) that could feedback on the evolutionary dynamics of the trait itself. For instance, a trait may be favoured by natural selection only when rare in a complex population, becoming disfavoured when it is more frequent. These complex evolutionary and ecological dynamics, where essentially the success of a trait depends on the composition of the community, can be mathematically analysed with evolutionary game theory (Nowak and Sigmund, 2004; Frey, 2010).

Evolutionary game theory is a mathematical framework that comes from classical game theory used to describe the behaviour of rational players. Classical game theory tries to analyse the behaviour in conflicts in economic and social settings in which the success of an individual strategy depends on the strategies employed by the other players. A well-studied example in game theory is the prisoner’s dilemma in which the choices to either confess or remain silent determine whether two suspects are considered guilty (Axelrod, 1990). In evolutionary game theory, the strategies are not associated to rational and cognitive choices, but are traits encoded into inherited programs that can be passed to the offspring (for this reason, the terms trait and strategies are used in an indistinguishable manner). Traits such as the usage of metabolic pathways or the expression of certain enzymes can be then regarded as strategies and a successful strategy is then selected for.

In a microbial community composed of species that compete using different strategies, each of the individual cells possesses a fitness that depends on its strategy and on the strategy of the individuals with whom it interacts. Individuals that use more successful strategies have higher chances to propagate and their frequency in the community will increase. Although the dynamics of an evolutionary game theory model can be studied analytically when the set of strategies is small, due to the large number of interactions taking place in microbial communities many authors prefer to simulate the dynamics of the community using agent-based modelling. In these models, the replication and death of individual cells (agents) are explicitly simulated using a system updated by a series of discrete events (Adami et al., 2016). These types of models also include the possibility of adding mutations that can introduce novel strategies not yet present in the species, which can be used to simulate random evolution of members of the community (Eriksson and Lindgren, 2005).
In cellular populations, a cooperative trait is often characterised by the presence of a shared public good, which is a finite resource, produced by cooperative cells and that is freely available to all other cells. The presence of a public good is always associated with the risk of cheating cells, which exploit the public good without providing any contribution to it and which can spread in the population – due to their improved fitness arising from not investing the costs associated with public good production. Although in this review we focus on microbial populations, this is a very general issue in the sustainability of many organisms at different scales including humans, justifying why the evolution (and resilience) of cooperation is considered one of the major open questions in biology (Pennisi, 2009).

Evolutionary conflicts between cooperative and cheating cells have been studied in a variety of microbial scenarios, including the conversion of sucrose into glucose by the yeast Saccharomyces cerevisiae (Gore et al., 2009), the production of the shareable iron-scavenging siderophore pyoverdine in Pseudomonas aeruginosa (Kümmerli et al., 2009) and the formation of fruiting bodies in Myxobacteria (Velicer and Vos, 2009). Given the potential similarities with cellulose and other polymers biodegradation, the example from yeast is worth explaining further. In this case, cooperative and cheating cells only differ by the production of the enzyme invertase that converts sucrose into glucose and fructose. Both monosaccharides can eventually diffuse away from the producing cell and become available to neighbouring cells. In other words, they become public goods: cooperators — the cells that ‘feed’ themselves and their neighbours at the expense of expressing the enzyme — can be exploited by cheaters, cells that do not express the enzyme and rely on cooperators to make food (Fig. 1A). In a scenario like this, it would be expected that cheaters could take over the population. However, the fitness of the cells is a non-linear function of the glucose concentration and, for certain values of glucose uptake and metabolic cost of enzyme production, it is possible to observe the co-existence of the two species as anticipated by an evolutionary game theory model (Gore et al., 2009). In fact, in a complex community composed of multitude of species it is likely that such mechanistic properties relating to the implementation of the different strategies, such as regulatory mechanisms controlling the production of a public good, will affect the evolutionary and ecological dynamics of the strategies and thus the whole community.

Before discussing further these potential mechanisms that can stabilise cooperative interactions, we will first describe types of cooperative interactions in microbial populations.

Microbial cooperations based on public goods

Shared (public) goods are molecules produced by certain individuals and can benefit the entire population (West et al., 2007). As explained above, these molecules are synthesised at a cost and,
therefore, are susceptible to be exploited by cheater cells that can benefit from them but do not contribute to their production – hence acquiring a fitness advantage over cooperators. This type of cooperation is based on a large variety of shared molecules: siderophores, enzymes, biosurfactants, components of biofilm matrix, quorum sensing molecules, bacteriocins (proteins secreted by one strain to inhibit the growth of a closely related strain) and toxins as summarized in (West et al., 2007). Given their interest in microbial biotechnology, in this review we will focus on secretion of degradatory enzymes.

Microorganisms digest large macromolecules, which are poorly soluble, through the secretion of extracellular enzymes. The macromolecules are typically polymers of biological or synthetic origin, such as starch, cellulose and polyesters, which constitute an abundant source of nutrients for bacteria, fungi and other eukaryotic microorganisms (Allison, 2005; Richards and Talbot, 2013). These polymers also constitute a very interesting substrate for industrial bioprocesses, as they are inexpensive, biodegradable at some extent and often obtained from renewable sources (Gross and Kalra, 2002). The enzymes secreted by microorganisms act by degrading the macromolecules into simpler and smaller components that can then be assimilated by the microbial community (Burns, 2010). In this scenario, the dynamics of the cooperating and cheating populations depend on parameters such as the cost of producing the enzymes and their diffusibility (Allison, 2005).

Cellulases and oxidative enzymes secreted to cleave cellulose such as cellobiase dehydrogenases can be considered as instances of ‘public goods’ (Dimarogona et al., 2012) and are found in the genome of most wood-degrading microbial communities (Zamocky et al., 2006). Similar to cellulases, amylases capable of degrading the glycosidic linkages of starches also play an important role as public goods and have been identified in many bacteria and fungi, such as Bacillus subtilis (Coleman and Elliott, 1962), Thermomyces lanuginosus (Arnesen et al., 1998), Penicillium expansum (Doyle et al., 1998), and several species of Streptomyces (El-Fallal et al., 2012). Similarly, enzymes responsible for the digestion of other macromolecules such as extracellular lipases and proteases are also examples of public goods, and their production in a complex microbial community is influenced by the interactions between its members (Willsey and Wargo, 2015). Collectively produced enzymes are also responsible for the degradation of oil-derived plastic polymers such as poly-ethylene terephthalate (PET). The identification of bacterial species producing enzymes capable of PET depolymerisation, therefore generating molecules that can then be assimilated by the microbial community in that niche (Chen et al., 2010; Yoshida et al., 2016) paves the way for the remediation of PET waste and its use as a bioprocessing substrate (Wierckx et al., 2015).
Microbial cooperations based on metabolic interactions

Metabolic exchange is another way in which microorganisms can interact cooperatively. Metabolic interactions are widespread in natural microbial communities and arise from metabolites from one species being used as energy sources or building blocks by other species (Paczia et al., 2012; Cooper and Smith, 2015; Fiore et al., 2015). The former scenario leads to cross-feeding, whereas the latter can lead to emergence of auxotrophies (an organism fully relying on the environmental provision of certain compounds required for its growth) (Fig 1B). The metabolites released into the environment can be explained by either passive or active means, i.e. organisms not being able to maintain certain compounds due to leakage issues or actively secreting those compounds due to some functional benefits. While the former explanation could arise due to some fundamental biophysical limitations on biological membranes, the second (functional) explanation is difficult to ration within a simplistic view of organismal fitness. One could naively argue that since other organisms use the secreted metabolites as a resource, evolution should have allowed the ‘secreting organism’ also to innovate that capacity of using this metabolite (as an energy source or building block) rather than secreting it. This naive view, however, ignores limitations arising from cellular tradeoffs and thermodynamics.

Metabolic interactions emerging from thermodynamic limitations

In principle, cross-feeding and auxotrophic interactions could be seen as an extreme form of cooperation (i.e., ‘altruism’) as they benefit only the receiving organisms. Under certain conditions, however, secretion of internal metabolites can also benefit the producer leading to a mutually-beneficial interaction: if the products released have an inhibitory effect on the producer, the presence of an additional species that would assimilate these products would lead to more mild forms of cooperative interaction rather than a straight ‘altruistic’ act on behalf of the producer (Lilja and Johnson, 2016). More specifically, this type cross-feeding interaction, involving release of inhibition arising from byproducts of metabolism of one organism by another is often referred to as syntrophy (Fig 2A). The most-well known example is the H₂-mediated syntrophic interactions between secondary degraders and methanogens (Schink, 1997). In these interactions, the inhibition of the degrading species arises due to its growth-supporting metabolic reaction reaching towards thermodynamic equilibrium as H₂ accumulates (Schink, 1997; Großkopf and Soyer, 2016). This ‘thermodynamic inhibition’ is relieved by the consuming of H₂ by the syntrophic partners (McInerney and Bryant, 1981; Seitz et al., 1988; Scholten and Conrad, 2000), creating a situation in which continued growth is only possible when the two partners co-exist. Many of the biodegradation processes consist of individual syntrophic and cross-feeding interactions among different species (Schink, 1997), with examples including the degradation of monoaromatic and polyaromatic
compounds in syntrophy with methanogens (Knoll and Winter, 1989; Berdugo-Clavijo et al., 2012; Morris et al., 2013). Syntrophic interactions are also important in oil-degrading microbial communities, although the exact roles of many individual members in these communities are less clear. It has been reported, for instance, that syntrophic interactions between Desulfatibacillum alkenivorans and Methanospirillum hungatei are necessary to degrade refractory hydrocarbons (Westerholm et al., 2011; Callaghan et al., 2012).

These examples illustrate how ubiquitous and essential syntrophic interactions are for complete degradation of organic compounds. Therefore, for fully being able to optimize bioprocesses and biotechnologies around organic degradation and transformations we need a better understanding of the emergence and maintenance of metabolic cooperations. It is important to note that syntrophic and cross-feeding interactions are shown to alter cellular metabolic fluxes within individual species, as well as in simple communities such that the presence of a downstream syntrophic partner can result in changes in the metabolic by-products and yields from upstream producer microorganisms (McInerney and Bryant, 1981; Seitz et al., 1988; Schink, 1997; Scholten and Conrad, 2000). In other words, organisms’ preferred metabolic routes (or ‘strategies’) would change with local substrate/product availabilities (as well as internal constraints such as on uptake rates or cofactor availabilities), but these in turn would depend on what other organisms would choose to do metabolically. From a theoretical perspective, this situation cannot be analysed assuming a simple individual fitness optimization under constant selection pressure, but would require instead the combination of evolutionary game theory and ecology in order to develop theoretical frameworks and experimental model systems accounting for the described complex interplays.

The inclusion of thermodynamics in models of microbial growth and metabolism could contribute to unravel the emergence of metabolic interactions. Taking into account the thermodynamic constraints of growth-supporting microbial biochemical reactions would enable better capturing changes in the concentrations of different compounds in the environment and thus allow direct linkage between ecology and individual growth rates. There have been several recent attempts in this direction, and models including the thermodynamics of metabolic reactions have been successfully employed to describe the dynamics of some biodegradation processes, such as the fermentation of glucose and the reduction of nitrate (González-Cabaleiro et al., 2013, 2015; Cueto-Rojas et al., 2015), to explain microbial diversity (Großkopf and Soyer, 2016), as well as to model individual species growth (Hoh and Cord-Ruwisch, 1996; Jin and Bethke, 2007). Additional works in this direction will allow better predictive models to explain evolutionary and ecological dynamics of
microbial communities under conditions where thermodynamics-driven metabolic interactions dominate.

**Metabolic interactions emerging from cellular tradeoffs**

As discussed above, fitness optimization is a complex function of multiple traits and it is subject to intrinsic tradeoffs that could readily explain metabolic secretions. In particular, the optimization of ATP-generating pathways under limitations on enzyme investment and internal metabolic concentrations is shown to lead to the evolution of impartial pathways and metabolite excretion (Pfeiffer and Bonhoeffer, 2004). Similarly, limitations on membrane space and internal resources such as enzymes and conserved moieties can cause tradeoffs in substrate uptake rates and internal metabolic fluxes, resulting in different genotypes that differentially utilize respiratory (i.e. pathways ending with inorganic terminal electron acceptors) and fermentation (i.e. pathways ending with organic terminal electron acceptors) pathways (Majewski and Domach, 1990; Vemuri et al., 2006; Molenaar et al., 2009; Zhuang et al., 2011; van Hoek and Merks, 2012; Flamholz et al., 2013; Basan et al., 2015). Since the end products of fermentative pathways are usually still able to sustain further microbial growth, this could again explain the first stage of formation of metabolic interactions through metabolic excretions. Subsequently, limitations on substrate uptake are predicted to act as a force to drive metabolic specialization on such excreted compounds (Doebeli, 2002; Spencer et al., 2007).

The idea of cellular tradeoffs driving the emergence of metabolic cross-feeding has recently been evaluated in a combined *in silico* and experimental evolution study (Großkopf et al., 2016). In that study, the authors have incorporated tradeoffs in a stoichiometric metabolic model of *E. coli* by imposing global constraints on the total uptake rates. This model was then simulated using dynamical flux balance analysis, which allows modelling of both microbial growth and environmental substrate concentrations, and mutations, which can alter the distribution of total uptake flux among different substrates. In other words, this approach combined simulation of ecological and evolutionary dynamics at the same time; starting from a single model, the *in silico* simulations can lead to alterations both in the environmental conditions and mutant models (Fig. 2B). The application of this approach to the modelling of the experimental long-term evolution of *Escherichia coli* revealed that the combination of tradeoffs and ecological/evolutionary dynamics results in the emergence of two dominant models (Fig. 2C). These two models have distinct uptake fluxes suggestive of a cross-feeding interaction; one model had increased glucose uptake and acetate excretion rate and the other had increased acetate uptake rate (Großkopf et al., 2016). Further experimental analyses revealed that the two models show metabolic flux patterns that qualitatively
match experimentally observed genotypes in one lineage of the long-term experiments, indicating that this approach might provide useful insights into how ecological and evolutionary dynamics can shape metabolic systems. Indeed, an emerging trend in the analysis of community dynamics is to increasingly combine multi-species ecological simulations with stoichiometric models describing the metabolism of those interacting species in an attempt to generate insights into ecology – evolutionary interplay (Louca and Doebeli, 2015; Widder et al., 2016; Zomorrodi and Segrè, 2016).

Factors contributing to the stabilization of cooperative interactions in microbial populations

Structured environments

One of the basic mechanisms that affect the resilience of cooperation is the presence of spatial structure. Structure would ultimately facilitate the resilience of cooperation as it allows the ‘segregation’ of cooperative from cheating cells (Nowak, 2006) (i.e., cooperative cells can then share the produced public good with the similar trait, excluding cheating cells) (Fig 3A).

There are several theoretical studies and experimental evidences of spatial segregation in cellular populations (Van Dyken et al., 2013), with biofilms being a paradigmatic example of bacterial communities exhibiting stable cooperation due to the segregation in structured environments (Nadell et al., 2009). The structure and composition of biofilms can feedback on the highly dynamic competition between sub-populations of cooperators (i.e., contributing to the biofilm assembly) and cheaters. In these circumstances, the spatial arrangements of the distinct genotypes crucially affect the degree of cooperation and competition present in the biofilm (Nadell et al., 2016).

A broader notion of structure can also refer to the case of having a population distributed into different heterogeneous sub-populations that may be spatially segregated (e.g. forming colonies). In this case the structure of the population can lead to a characteristic issue of multi-level selection known as Simpson’s paradox. Simpson’s paradox is a statistical phenomenon that can emerge when comparing groups of data; groups can display a trend when analysing them individually, but this trend is reversed when the groups are combined. A famous example of Simpson’s paradox is the one behind the gender discrimination accusation against the University of Berkeley in early 1970s. In that case, 44% of the total male applications to the graduate school were accepted against the 35% of the female applicants suggesting a bias against female applicants. Looking into how the applications were distributed among the different departments, however, it became clear that there was no bias, and the differences in the rates were the result of a majority of women having applied to the most competitive departments, which decreased the success rate of the female applicants. In other words, the apparent bias is only the result of the ways the
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1. applications are aggregated together (Bickel et al., 1975). In the context of microbial communities, Simpson’s paradox is shown to emerge when the different sub-groups are sufficiently heterogeneous in their composition to guarantee that in the aggregate population the cooperative individuals have an advantage over the cheating cells (despite in each of the colonies – the disaggregated population – cheaters are favoured) (Chuang et al., 2009). This finding suggests that the opportune design of the organization of a microbial community in sub-populations (and subsequent coalescence of those sub-populations) may be useful to improve its resilience to detrimental mutants. In general, other more complex notions of structured populations from ecology (e.g., meta-population dynamics) could also be relevant to understand and control the evolutionary dynamics of cooperative interactions (Datta et al., 2013).

Interplay between ecological and evolutionary dynamics

Another stabilising and driving factor beyond cooperative interactions in microbial communities is the interplay between ecological and evolutionary dynamics that results in changes in the composition of the community over time. This happens when, due to the interactions in a community, certain traits (such as cheating and cooperation) are selected for or against, resulting in rapid changes in the frequency of the individuals carrying the trait that affect the ecology of the global community. The changes in the ecology can then feed-back on the selective advantage of the different traits (as discussed above), leading to an eco-evolutionary feedback (Fig. 3B) (Lennon and Denef, 2015). This aspect has become of recent interest due to several theoretical and experimental studies showing the non-trivial effects of the time-scales overlap between ecology and evolution in what are called eco-evo feedbacks (Schoener, 2011). There are several examples of eco-evo feedbacks in microbial populations investigated experimentally (Fiegnana and Velicer, 2003; Ross-Gillespie et al., 2009; Moreno-Fenoll et al., 2017) with the most known example being the interplay between population density and fitness (Sanchez and Gore, 2013). For instance, in the yeast communities discussed above, cooperative cells have higher fitness than cheating cells only at lower population density. This, coupled to the fact that cheaters lead to lower population growth, facilitates the observed co-existence between the two traits, i.e. the stabilisation of cooperation (Sanchez and Gore, 2013). Eco-evo feedbacks can be modelled by adding notions of population dynamics to evolutionary game theory, leading to the framework of ecological public good games (Hauert et al., 2008) that extend the standard evolutionary game theory (in which, usually, the focus of the analysis is the change in frequency of a certain trait). Combination of population dynamics with metabolic models at the level of individual species or genotypes (Harcombe et al., 2014) with
evolutionary dynamics (Großkopf et al., 2016) is another promising route towards capturing eco-
evolutionary dynamics, especially when cooperative interactions involve metabolite secretions.

Regulatory mechanisms

Another potential factor for the stabilisation of cooperation that has recently attracted attention is
cellular regulatory mechanisms. Animals, including humans, have developed complex social
strategies to control cheaters, and there is great interest in determining to which extent single cell
organisms could employ similar mechanisms to fight detrimental mutants (Travisano and Velicer,
2004).

One of these regulatory mechanisms is known as ‘reciprocity’. In this case the amount
contributed of a public good depends on the environmental conditions, which in turn may depend
on the contributions made by others. This is for instance the case of iron uptake in *P. aeruginosa*
where iron scavenging siderophores (the public good) are released in greater or smaller quantities
depending on the amount of iron in the environment (Kümmerli et al., 2009). Recent experiments
using this system have confirmed that cells use a type of ‘reciprocity’ that facilitates the control of
cheaters: the cellular decision of producing public good is made only in an environment with many
producers. In other words, the cells seem to implement a rule stating ‘cooperate when surrounded
by mostly cooperators’. Coupled to quorum sensing, this rule allows bacteria to match their
investment at lower levels of population structuring and it is an effective way to repress cheaters
(Allen et al., 2016). In yeast, a similar mechanism happens in the production of invertase. Another
regulatory mechanism that could be interpreted as a functional ‘decision’ to limit the spread of
cheaters is to increase the noise in the expression of genes encoding for public goods (Gore et al.,
2009). This is the case of self-destructive cooperation, in which cooperative cells die while helping
others, for example, as it happens during the secretion of toxins that enhance the colonization of
tissues by certain bacterial pathogens (Ackermann et al., 2008). Since the toxin is genetically
encoded, it is only expressed by a fraction of the population or the whole microbial population
would die. The ‘decision’ on which cells make the ultimate sacrifice is given by the stochastic
expression of the gene encoding the toxin. Similarly, cell-cell variability in the production of other
kinds of public goods may allow cooperative cells to temporarily switch off the production of a public
good, therefore limiting its cost and allowing for enhanced competition against the cheating cells.

These types of cellular decision-making mechanisms can interplay with an underlying eco-
evo dynamics (Harrington and Sanchez, 2014) and crucially affect the resilience of cooperation, as
shown in theoretical models (Cavaliere and Poyatos, 2013) (Fig. 3C). Thus, it is plausible to propose
the control of public good production for successful bioprocesses (such as the described cellulose
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1. Cooperation in microbial biotechnology

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4. Horizontal gene transfer of cooperative traits

5. Mobile genetic elements (plasmids, bacteriophages, transposons, etc.) transmitted via horizontal gene transfer are one of the main factors contributing to shaping microbial evolution. Apart from the genes essential for the replication and transmission essential for the mobile elements, they often carry multiple traits the enable social interactions in microbial communities and make them active agents defining the evolutionary dynamics of these communities (Rankin et al., 2011).

6. Cooperative traits such as public good producing exoenzymes are commonly acquired due to the transference of mobile elements. In fact, a genomic analysis in some bacterial species show that the frequency of genes encoding extracellular proteins is significantly higher in chromosomal locations known to be transferred (e.g. transposons) compared to regions that are not, and the frequency is even higher in plasmids, which were the most mobile elements present in the analysis (Nogueira et al., 2009). Horizontal gene transfer is also responsible for the transmission of exoenzymes in eukaryotic microorganisms, as revealed by a similar analysis carried out in osmotrophic fungi, in which it became evident that not only the enzymes, but also the transporters required for the uptake of the products resulting from the activity of the enzymes on large polymers, were encoded in mobile genetic elements (Richards and Talbot, 2013).

7. These observations are consistent with the idea of mobile elements enabling cooperation in a community owing to the invasion of mobile elements transmitting cooperative traits. However, the mobile elements also generate a cost to the cells harbouring them and, therefore, can potentially be lost or outcompeted by ‘cheat’ genetic elements (Rankin et al., 2011). Recent experimental evidences show nevertheless that horizontal gene transfer helps to maintain the production of public goods despite the potential presence of non-cooperative organisms and non-cooperative mobile elements (Dimitriu et al., 2015) owing, among other factors, to the increase in genetic relatedness due to the presence of the mobile elements (Mc Ginty et al., 2013). In other words, transmissible mobile elements allow for the local enrichment in cooperative interactions, which may, in the long term, lead to the specialization of sub-populations in cooperative niches specially in the presence of strong structure (Niehus et al., 2015).

8. The relevance of cooperation for biotechnological applications

9. The presence of cooperative interactions facilitates the development of complex functions that would be otherwise difficult or impossible (Nowak, 2006).
Cooperative microorganisms can exhibit distribution of labour: a large collection of distinct phenotypic behaviours, organized in subpopulations, can coordinate to fulfil some complex tasks in a collective way (Fig 4A). Shared diffusible molecules allow cells to communicate and spatially distribute the labour. Examples of complex tasks range from the controlled growth of biofilms depending on environmental conditions (Liu et al., 2015; Kim et al., 2016) to the distributed computation of Boolean functions (Regot et al., 2011).

This type of interaction is commonly observed in biodegradative processes carried out by interspecies biofilms. For instance, the presence of an algae in a microbial consortium with more than nine bacterial species enhances the degradation of the pesticide diclofop methyl (Wolfaardt et al., 1994). Another interesting example is the syntrophic interaction between the non-cellulolytic species *Treponema bryantii*, and the cellulolytic species *Ruminococcus flavefaciens*, to enhance the rate of cellulose degradation. The slowly growing cultures of *R. flavefaciens* benefits from *T. bryantii* removing the cellulolytic product, which results in higher population density and degradation rates (James et al., 1995).

Distribution of labour is, however, not restricted to spatially structured populations or populations composed by more than one species, but can also apply to other biological processes like the biochemical pathways for the degradation of aromatics in populations composed of one strain (Nikel et al., 2014). These pathways are sometimes organised into two distinct gene operons, one encoding for the activities required to funnel the aromatic substrate into a more affordable aromatic carbon source and a second required to transform this aromatic compound into central metabolites. For instance, the TOL pathway of *Pseudomonas putida* responsible for toluene and xylene degradation contains an ‘upper’ part that converts toluene into benzoate, and a ‘lower’ segment responsible for the degradation of benzoate (Franklin et al., 1981). In principle, it would be expected that all cells express both operons when a clonal population of *P. putida* is cultured in the presence of toluene but, surprisingly, many of the cells display a near bimodal distribution expressing either one operon or the other (Nikel et al., 2014). The mechanistic explanation of this behaviour is unknown although a plausible explanation of the phenotypic distribution may arise from the intricate transcriptional control of the operons (Silva-Rocha and de Lorenzo, 2012).

Distribution of labour also appears in the anaerobic metabolism of aromatic compounds in *Rhodopseudomonas palustris*. Monocultures of this species organise in three different subpopulations when using *p*-coumarate or benzoate as the carbon source. Each of these subpopulations is responsible for the utilization of either the aromatic compound, CO₂ and H₂ or, when growing on benzoate, N₂ and formate, forming a syntrophic consortia *de facto* composed of a single species (Karpinets et al., 2009). However, whether this particular type of cooperative cross-
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Feeding interaction is advantageous to prevent waste of resources or accumulation of toxic intermediates is an open question.

Distribution of labour can also be engineered together with cooperative traits in ‘synthetic’ communities (Fig. 4B). This is the case of co-culturing engineered strains of the bacterium *E. coli* and the yeast *S. cerevisiae* that are artificially mutualistic. Each of these strains is modified to express one module of the biosynthetic pathway of an antitumoral compound of interest (the acetylated diol paclitaxel precursor). The cooperation between these species allows production of taxanes with higher yields than using *E. coli* alone. The mixed culture combines the capabilities of *E. coli* for producing the intermediate taxadiene with the superior properties of *S. cerevisiae* compared to *E. coli* to catalyse the oxygenation reactions required to render the final compound (Zhou et al., 2015).

Synthetic consortia can be used in bioprocesses even in the absence of mutualism as explained in the previous sections (e.g. if eco-evo feedbacks take place). This is the case of an artificial community designed to produce isobutanol from cellulosic biomass composed by the fungus *Trichoderma reesei* and an engineered strain of *E. coli*. In this consortium *T. reesei* acts as a cooperator secreting cellulases required to degrade lignocellulosic polymers and the resulting saccharides are used to feed the *E. coli* strain that delivers the final product (Minty et al., 2013).

Synthetic communities can also improve biodegradation processes compared to monocultures. Degradation of crude oil is a good example in which microbial communities can exhibit cooperative interactions in Nature including metabolic cross-talk and shared goods that may contribute to the formation of interspecies biofilms (McGenity et al., 2012). Moreover, these interactions can be harnessed to produce artificial communities with enhanced degradation capabilities suitable for oil removal (Gallego et al., 2007). Another example is the desulphurization of dibenzo thiophene (DBT) to form sulphur-free 2-hydroxybiphenyl. In a recent work, DBT desulphurization was carried out using either an engineered *P. putida* strain expressing all the *dszABCD* genes required in the process, or a mixed culture of the same strain expressing only some of the genes. In this experiment, desulphuration of DBT was higher when combining multiple cells ‘specialising’ in one step of the biochemical pathway compared to the case of having all reactions taking place in the same organism (Martínez et al., 2016).

Cooperative interactions in microbial communities can also lead to higher resistance to environmental and ecological stress. Empirical observations using artificial communities of yeast show that this resistance takes place over a wide range of conditions (Gore et al., 2009). In addition, experiments carried out with engineered populations of *Bacillus subtilis* lacking the ability to form biofilms show that they nevertheless tend to form clusters that, although can have reduced growth due to limited mobility, allow the cells to endure harsh environmental conditions (Ratzke and Gore, 2016).
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In this case, cooperative individuals tend to aggregate leading to the ‘privatization’ of public goods and to the exclusion of cheating individuals (Pande et al., 2016). On the other hand, the loss of cooperation makes cellular communities more fragile (Sanchez and Gore, 2013) and more vulnerable to compositional shifts arising, for example, from antibiotic treatments (Liu et al., 2015). The fact that these behaviours are observed in experiments with different manipulated species suggests that these mechanisms are general and could be commonplace in Nature.

The presence of mechanisms that facilitate cooperation can also lead to complex co-evolutionary dynamics with the consequent emergence of novel social interactions. The most significant example in this respect is the mechanism of quorum sensing (QS) that is involved in controlling the investment in ‘public goods’ (Allen et al., 2016). Although the original role of QS is unknown, its ability to facilitate the (beneficial) presence of cooperative interactions may have led to the selection of complex functionalities, e.g., coordinating the expression of genes involved in multiple cooperative strategies, often co-evolving with them (Popat et al., 2015). This example suggests the possibility of using the presence of cooperative interactions to direct the evolution of the communities towards other properties of interest.

**Conclusion**

The key point of evolutionary game theory is that the fitness of individuals depends not only on the environment but also on other members in the population. This theory provides a framework to understand the dynamics of many bioprocesses involving complex microbial populations (natural and synthetic) in which the fitness of an individual cell is in fact affected by the environment and by the presence of other cells. A particular case of this scenario concerns the presence of cooperative interactions based on public goods and metabolic interactions and that have been the main focus of this review. We have also discussed some of the factors shaping these interactions such as cellular and thermodynamic constraints, as well as factors stabilising them such as structured environments, feedbacks arising from the ecology of the population, cellular regulatory mechanisms implementing certain behavioural strategies and the role of mobile genetic elements. These properties endow cooperative microbial populations with the possibility to resist cheaters invasions and the capability of performing more sophisticated tasks.

Despite its growing use to study the evolution of cooperation, evolutionary game theory has had so far a very limited impact in field or industrial biotechnological applications in which the environmental conditions are generally not well-defined and may affect the microbial communities (Bouchez et al., 2000; Sayler and Ripp, 2000; Cases and de Lorenzo, 2005). In fact, we have presented several examples suggesting that cooperative interactions based on cross-feeding and
public goods are at the core of many processes relevant for industrial biotechnology including food, energy and environmental applications of microorganisms.

Therefore, they are suitable of improvement by incorporating the mechanisms investigated in the large literature of the evolution of cooperation. As we have discussed, populations could be manipulated based on thermodynamic constrains to promote certain metabolic (cooperative) interactions. Similarly, bioprocesses, including bioreactor design, could be engineered to account (and exploit) for eco-evo feedbacks and spatial organizations.

Understanding how syntrophy and cooperation endow the microbial populations with resistance and resilience against ecological and environmental disturbances like compositional shifts in the environment or antibiotic shocks could be used to engineer robust microbial communities with enhanced performance and predictable dynamics (Briones and Raskin, 2003; Allison and Martiny, 2008; Sözen et al., 2014). Overall, we believe that the migration of results and methodologies from the area of evolutionary game theory into the design of microbial consortia would facilitate the engineering of evolutionary resilient communities with a better performance in a wide range of biotechnological applications.

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Figure 1. (A) Interactions based on shared public goods. Some cells (cooperators, shown in black) produce an enzyme required to split a substrate into digestible products. Other cells (cheats, shown in grey), do not produce the enzyme but take advantage of the public goods produced by the others. (B) Interactions based on cross-feeding. Some cells in the community excrete metabolites that can be taken up by other cells giving rise to a web of interactions.
Figure 2. (A) Metabolic interactions that can take place in a population. Cells can exchange metabolites that are required to support each other’s growth in a mutualistic interaction (left). One of the cells can use a metabolite excreted by another cell, favouring in this way the metabolism of the producer through the pathways leading to the excretion (centre). When the metabolites excreted have an inhibitory effect on the producer (e.g. because they lead to thermodynamic equilibrium), the relationship with a degrader cell of the inhibitory metabolite is mutually beneficial and known as syntrophy (right). (B) Dynamic modelling of the evolution of FBA models. Cells can be modelled as metabolic networks exchanging metabolites with other cells in the population. In this abstraction each cell is represented by a Flux Balance Analysis model. These models can replicate over time and also evolve, producing populations composed by models with different constrains for uptake and secretion of metabolites. (C) Dynamic analysis of model genealogy. The frequency of each model in the population changes over time being the darkest bars the most abundant models. Due to mutations, new models arise and they are represented as new branches in the phylogeny. Plot redrawn from (Großkopf et al., 2016).
Figure 3. Mechanisms to Preserve Cooperation in Cellular Communities. (A): A structured environment can facilitate cooperation. The figure shows the growth of fluorescently labeled colonies (cooperators in red, cheaters in green) of *S. cerevisiae* (Figure from (Van Dyken et al., 2013)). Cooperative cells produce invertase that breaks down sucrose into digestible glucose and fructose. Non-producers cells (cheaters) have a fitness advantage because they do not produce invertase but can access glucose. In unstructured environment (liquid culture) cooperators decline. However, in a spatial environment (obtained by spotting a droplet of mixed cooperator/cheater cultures onto solid medium) cooperators can spread over cheaters. The diffusion of cells lead to the formation of discrete sectors - cooperator sectors are more productive than cheater sectors and will expand radially faster. (B): Eco-evo dynamics can preserve cooperation in communities of *S. cerevisiae* (redrawn from (Sanchez and Gore, 2013)). Red circles represent cooperative cells (invertase producers), green circles represent cheaters (non-producers). Below a certain cooperator density, there is little glucose available. Cooperative cells grow at a slow rate on the little amount of glucose they can retain, while cheater cells grow more slowly (it is crucial that cooperators have preferential access to the glucose). Above a certain cooperator density, both cooperators and cheaters grow at a fast rate because of the large pool of available glucose, but cheaters grow faster as they do not have the burden of producing invertase. Such density-dependent selection favors cooperators at low densities and cheaters at high densities, which leads to the stable coexistence of cooperative and cheating yeast cells. (C): Regulation of public good production can preserve cooperation in a meta-population model in which the population is transiently divided in sub-populations.
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(figure from Cavaliere and Poyatos, 2013). In-silico simulations present two possible successful types of regulation against cheaters: positive plasticity (top row) in which cooperators constraint cheaters by stopping the production of public good when cheaters appear (a) and fully restarting only when cheaters have disappeared (b) and negative plasticity (bottom row) in which cooperators produce permanently low amounts of public good which helps controlling cheaters invasion (c). Thick arrows denote the cellular decision to produce (P) or not produce (nP) the public good. The success of the regulation is coupled to the heterogeneity (variance) of the sub-populations, i.e., positive plasticity transiently modifies the variance while negative plasticity keeps a relatively constant heterogeneity (variance shown in (d) correspond to trajectories (b) and (c), respectively).
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Figure 4. (A) Division of labour in microbial populations. Colonies of Pseudomonas fluorescens P0-1 are composed by cells with two different morphologies known as mucoid and dry that can evolve from each other due to a single mutation (left picture). Colonies composed by a mixture of the two phenotypes expand faster allowing cells to colonise larger regions in shorter periods of time compared to colonies composed by each of the individual phenotypes. The two morphotypes occupy different regions of the colony as shown when labelled with fluorescent reporters (centre). Dry cells (in red) exhibit a radial distribution growing on top of the mucoid (in green). Confocal microscopy reveals that the edge of the colony (right picture) displays a distinct spatial organization in which mucoid cells form a thin strip at the very edge. The differentiation and spatial segregation allows the distribution of labour in the population: Mucoid cells produce a lubricant polymer at the edge, whereas dry cells sit behind and push both of them along. The cooperation of these two phenotypes results in a fast growing colony. Pictures have been reproduced from (Kim et al., 2016). (B) Engineered populations can improve bioprocesses. Two strains are combined to carry out the synthesis of a product of interest (red pentagons) that can not be produced using each of the strains individually. The process involves that one of produces an intermediate (the yellow pentagon) that is used by the other to synthesize the final product. If the two cells compete for the same resources (e.g. carbon source shown by the blue hexagon; left panel) the population with the lower fitness under those conditions will eventually collapse. However, when the two populations are engineered so that one grows at the expenses of the other (e.g. through cross-feed or syntrophy shown by the purple triangle), the two populations cooperate (centre panel) and the synthesis of the product of interest takes place for a longer period of time resulting in higher yields (right panel). Panels inspired by (Zhou et al., 2015).
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