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Scalable and exhaustive screening of metabolic functions carried out by microbial consortia

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Abstract

Motivation: The selection of species exhibiting metabolic behaviors of interest is a challenging step when switching from the investigation of a large microbiota to the study of functions effectiveness. Approaches based on a compartmentalized framework are not scalable. The output of scalable approaches based on a non-compartmentalized modeling may be so large that it has neither been explored nor handled so far.

Results: We present the Miscoto tool to facilitate the selection of a community optimizing a desired function in a microbiome by reporting several possibilities which can be then sorted according to biological criteria. Communities are exhaustively identified using logical programming and by combining the non-compartmentalized and the compartmentalized frameworks. The benchmarking of 4.9 million metabolic functions associated with the Human Microbiome Project, shows that Miscoto is suited to screen and classify metabolic producibility in terms of feasibility, functional redundancy and cooperation processes involved. As an illustration of a host-microbial system, screening the Recon 2.2 human metabolism highlights the role of different consortia within a family of 773 intestinal bacteria.

Availability and implementation: Miscoto source code, instructions for use and examples are available at: https://github.com/cfrioux/miscoto.

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1 Introduction

The rise of metagenomics through sequencing advances and efficient computational biology techniques has led to a broad range of data and perspectives for unraveling the complexity of ecosystems and elucidating the role of species within microbiomes (Marchesi et al., 2015). Altered traits displayed by some species in axenic cultures confirmed that phenotypes are linked to interactions occurring between an organism and its microbiota (Douglas, 1992; Ridley et al., 2012; Tapia et al., 2016). Therefore, more consideration tends to be given to organisms as parts of communities or interacting actors (Fuentes et al., 2016; KleinJan et al., 2017; Molloy et al., 2017) while exploration of organisms in isolation is acknowledged to be reductionist (Cavaliere et al., 2017). Bioinformatics methods are of great help in understanding, modeling, or suggesting interactions within microbiomes, at several scales and levels of resolution (Li et al., 2016). At a first rough scale, investigations relating to microbial co-occurrence apply to the full sequenced microbiotas (Faust et al., 2012). Then, metabolic models give insights into the precise mechanisms underlying the competition, cooperation or commensalism relationships. They have been widely exploited (van der Ark et al., 2017) using methods ranging from the modeling of meta-organisms with abstracted boundaries (Greenblum et al., 2012) to constraint-based modeling (Heinken et al., 2015b) of small groups of organisms, often called communities (Khandelwal et al., 2013; Stokyar et al., 2007).

Naturally, the resolution of the proposed interactions will be more precise as both the size of the community decreases and the degree of knowledge on its members increases (Johns et al., 2016). In large communities, graph-based analysis of metabolic models allows for a fast and efficient screening of organisms by scoring their competition (Kreimer et al., 2012) or cooperation (Levy et al., 2015) potential. When focusing on a smaller number of organisms, it becomes possible to quantitatively predict the behaviors of the system, which may be either the production of compounds of interest in synthetic biology studies in a controlled system (Großkopf et al., 2014), or simply growth with respect to environmental studies. For instance, (Greenblum et al., 2012) relied on a constraint-based framework to model each species of a community as a compartment that interacts with others. This allows growth to be simulated (Mendes-Soares et al., 2016) possibly in a dynamic way (Zomorodi et al., 2014), coupled with co-occurrence data (Zeleznik et al., 2015) or with the
integration of spatial/temporal constraints (Bauer et al., 2017; Harcombe et al., 2014). Budinich et al. (2017) studied microbial communities under Pareto optimality to describe all feasible growth rate solutions. Heinken et al. (2015a) studied eleven bacteria, along with the human host, to predict interactions under four dietary regimes.

The selection of species is a very challenging step when switching from the investigation of microbial co-occurrences within a large microbiota to the study of functions effectiveness within the community itself. This task entails identifying species of interest among all the microbiota in order to fit various criteria depicting the added value of the microbiome over an individual organism. Along these lines, Julien-Laferrière et al. (2016) proposed to extend the gap-filling concept used in metabolic network reconstruction frameworks (Pan et al., 2018) to create synthetic communities among a small number of bacterial candidates. They select species that own enzymes of interest, facultative exogenous genes and model transports between organisms, i.e. exchanged metabolites. The latter can be used either as energy sources or metabolic precursors i.e. building blocks by recipient organisms (Cavaliere et al., 2017). The very high combinatorics between all exchanges viewed as variables makes its impossible to scale their method either to the study of massive datasets involving hundreds or thousands of organisms or to the sorting of possibly hundreds of targeted functions. Eng et al. (2016) developed CoMiDA for the purpose of scaling to hundreds of species and targeted compounds using a non-compartmentalized, boundary-free level of modeling, called mixed-bag or gene-soup (Henry et al., 2016). The price to pay for scalability was to get rid of the number of exchanges metabolites involved in the selected community functioning. Although not applied in the same context in terms of the size of datasets considered or the optimization performed, both methods entail optimizing a parsimonious criterion. In one case, the MultiPus algorithm minimizes the total cost of the synthesis, mostly resulting from exchanges and exogenous reactions (Julien-Laferrière et al., 2016), whereas the second algorithm minimizes the size of the community (Eng et al., 2016) and provides one single solution that meets the desired objective. It does not enable sampling nor exploring the possibly broad range of feasible communities that meet the objective used for selection, which can be huge in large microbiotas. This is unfortunate considering that the number of equivalent feasible communities provides hints on redundancy in the microbiome regarding the targeted function, increasing its chances of being really effective. Having several equivalent solutions is an asset, as they can, a posteriori, be filtered by experimenters, based on biological knowledge or additional criteria, prior to experimentation.

In this study, we introduce a method based on the exhaustive enumeration of communities combining a total cost synthesis optimization criterion with a community-size optimization criterion. The tool uses logical programming and combines the non-compartmentalized and the compartmentalized frameworks in order to exhaustively identify and explore all families of species in a microbiota enabling targeted metabolic behaviors. As a first step, the method mimics the approach introduced in Eng et al. (2016) to classify a targeted biological function as (i) unfeasible, (ii) feasible with a single organism or (iii) feasible only with several organisms. In the second step, we used logical programming combined with SAT-based solvers (Answer Set Programming) to compute all communities with a minimal size allowing a biological function to be effective; this is a generalization of the mixed-bag approach used by Eng et al. (2016) in a combinatorial-optimization setting. In the third step, as transports between organisms are costly, we introduce a criterion based on the minimization of exchanges to discriminate between minimal-size community solutions. This entails defining exchangeable metabolites and applying an additional optimization, similar to the one used in Julien-Laferrière et al. (2016), to a pre-selected family of communities. This method can be used to facilitate the selection of a community optimizing a desired function in a microbiota by reporting several possibilities which can then be sorted according to biological criteria. The method can also bypass the pre-determination of desired function by being generalized to the screening of all single metabolites within a host-microbial system. An output of the workflow applied in this context is a classification of individual target metabolites in terms of feasibility, functional redundancy and cooperation processes involved.

We applied our workflow to the Human Microbiome Project (HMP) to study community selection for the 4.9 million metabolic functions corresponding to the production of a single metabolic compound (target) in the microbiome from a single metabolic input (seed). Our analysis focused on the distribution of community sizes, redundancy between communities equivalently enabling a function and the complementarity of the mixed-bag and the exchanged-based frameworks. Our approach shows that only 8% of the functions require a community to be enabled, with a maximum of six bacteria. We studied 10% of the latter seed/target functions and observed that in 36.7% of cases, the number of equivalent feasible communities ranges from 100 to 1000 per function, suggesting a significant redundancy of functionalities within the HMP. Using an exchanged-based minimization criterion reduces the family of relevant communities by 24% on average, confirming that both criteria deserve to be considered together. As a matter of application, we investigated the role of different consortia within 773 intestinal bacteria in the production of cytosolic compounds within Recon2.2 human metabolic network.

2 Methods and implementation

Given a set of organisms each described by a metabolic model, the goal of our paper is to find a minimal subset of the available organisms that can synthesize a set of target products using available substrates. Our main specificity is that we aim to select the organisms according either to a mixed-bag production of compounds or to a compartmentalized one. The mixed-bag criterion, introduced in (Henry et al., 2016), describes the theoretical capability of a community, considered as a meta-organism, to produce a selected set of compounds thanks to several reactions. However, the use of a reaction by such a consortium involves the exchange of several metabolites: e.g. host gives to the helper one or more reactants that it is able to produce, and the helper gives back the product(s) of this reaction. Thus, reasoning in terms of reactions can be very costly for the organisms when considering what is actually exchanged: the metabolites. This is the main reason for our second compartmentalized criterion, which aims at putting minimization constraints on the metabolites rather than on the number of species. Both criteria are modeled by combinatorial optimization problems, the latter being significantly more difficult than the former because it requires the introduction, as a variable, of all the possible exchanges between all species. We use logical programming to smartly encode a bipartite-based semantics of producibility and solve the combinatorial problems using SAT-based heuristics.

2.1 Metabolic models viewed as bipartite graphs

We represent a metabolic network as a labeled directed bipartite graph \( G = (M, R, E) \), where \( M \) and \( R \) are sets of nodes standing for
reactions and metabolites respectively and \( E \subset M \times E \cup E \times M \) depict input and output reaction relationships. When \((m, r) \in E\) or \((r, m) \in E\) for \(m \in M\) and \(r \in R\), the metabolite \(m\) is called a reactant or product of reaction \(r\) respectively. More formally, for any \(r \in R\), define \(rcts(r, G) = \{m \in M | (m, r) \in E\}\) and \(prds(r, G) = \{m \in M | (r, m) \in E\}\).

Seeds \(S \subset M\) are compounds that are available to initiate metabolic producibility and fuel metabolic reactions. They usually are the metabolites present in the growth medium of the considered organism, but can also include compounds known to be produced or cofactors in complex internal production cycles (Eng et al., 2016; Greenblum et al., 2012; Julien-Laferrière et al., 2016). Targets \(T \subset M\) are metabolites expected to be produced by the considered organism. They can be components (reactants) of the biomass reactions or other compounds, experimentally observed, e.g. in metabolomics studies.

The functionality of metabolism is assessed using a producibility criterion. This producibility is calculated using a recursive definition and starts from the seeds. A metabolite is considered producible in two cases: either it is a seed, or it is a product (\(prd\)) of a reaction \(r\) whose reactants (\(rct\)) are producible. Formally, given a metabolic network \(G = (M, R, E)\), and a set \(S \subset M\) of seed metabolites, a metabolite \(m \in M\) is reachable from the seeds \(S\) in the network \(G\) if \(m \in S\) or if \(m \in prds(r)\) for some reaction \(r \in R\) where all \(m' \in rcts(r, G)\) are reachable from \(S\). The scope of \(S\) according to \(G\), written \(mxdbag\(S\), \(G)\), is the closure of metabolites reachable from \(S\) (Handorf et al., 2005). It is formally defined by \(mxdbag(S, G) = \cup_{M_s} S\), where \(M_0 = S\) and \(M_{i+1} = M_i \cup prds(r \in R | rcts(r) \subseteq M_i)\).

The concept of scope has been compared with the flux-based producibility in Kruse et al., (2008). In particular, it is equivalent to the network flow modeling of Eng et al. (2016) and to flux-based modeling in general, providing metabolite stoichiometries are set to 1 and accumulation of metabolites is allowed.

2.2 Mixed-bag and compartmentalized target producibility

A community of species is formally defined as a family of metabolic networks \(\{G_1, \ldots, G_N\}\). Following the definition of Henry et al. (2016), the mixed-bag or gene-soup modeling of communities considers a boundary-free meta-organism. Organisms’ capabilities are no longer examined individually but collectively in a virtual compartment. Therefore, the associate metabolic network is defined as \(mxdbag(G_1, \ldots, G_N) = (\cup G_i, \cup R_i, \cup E_i)\). The scope of the mixed-bag community from a set of seeds \(S\) is naturally defined as the scope of \(S\) according to the non-compartmentalized metabolic network: \(mxdbag\(S\), \(G_1, \ldots, G_N) = \cup mxdbag(S, G_i)\).

Considering that all organisms in the community have boundaries and correspond to different compartments requires establishing the family of allowed exchanges between organisms. We denote by \(exchg(G_1, G_N) = \{(m, i, j) | m \in M_i \cap M_j, i = j\}\) the set of reactions enabling the exchange of each compound \(m\), which belong to pairs of metabolic networks \(G_i\) and \(G_j\). Consider a set of exchange reactions \(E \subset exchg(G_1, G_N)\). The compartmentalized metabolic network associated with the community \(G_1, \ldots, G_N\) and the family of exchange \(E\) is defined as \(cptModel(G_1, \ldots, G_N, E) = (\cup M_i, \cup R_i, \cup E_i)\), for which components are compartmentalized with an index: metabolites are denoted by \((m, i)\), reactions are denoted by \((r, j)\) and edges are denoted by \((e, i, j)\). More formally, we have \(M = \cup_{i=1}^{N} (M_i \times \{(i)\})\), \(R = \cup_{i=1}^{N} (R_i \times \{(i)\})\) and \(E = \{(m, i, (e, i, j)) | (e, i, j) \in E_i \cup \{(m, i, (e, i, j)) | (e, i, j) \in E_i \cup \{(e, i, j) | (e, i, j) \in E_i \cup \{(e, i, j) | (e, i, j) \in E_i \}}\))\).

Given a set of substrate metabolites \(S\), we allow each of the seed in \(S\) to be imported into each considered organisms by creating a set of compartmentalized seeds: \(cptSeed(G_1, G_N, S) = \cup_{i} S \cap M_i \times \{(i)\}\). As shown in Figure 1, the media compounds \(A\) and \(B\), which both belong to the cytoplasmic metabolic networks of host and symbiont \(1\), are considered their seeds. Symbiont \(2\) has only \(A\) as seed and symbiont \(3\) has no internal seeds. By extension, we say that a target \(t\) belongs to the scope of the compartmentalized metabolic model if there is a metabolic model \(G_i\) that contains \(t\) and has it in its scope: \(scope(G_1, G_N, E, S) = \{t \in \cup_{i} (M_i \times \{(i)\}) | (t, k) \in scope(cptSeed(G_1, G_N, S), cptModel(G_1, G_N, E))\}\). Based on this definition, in Figure 1, the compound \(E\) belongs to the scope of \([A, B]\) because it can be produced in symbiont \(1\). On the contrary, target \(F\) does not belong to the compartmentalized scope of \([A, B]\) even though it belongs to the mixed-bag scope of \([A, B]\). There is not way of producing the precursor \(E\) in the host species according to the compartmentalized metabolic network: a transport of \(E\) to the host is needed.

Given a set of compounds \(T\) considered as relevant targets, we define a set of exchanges \(E\) to be consistently associated with a mixed-bag community in agreement with \(T\) when it is possible to produce, in the compartmentalized model, all the compounds from \(T\) that used to be producible in the mixed-bag model, i.e.: \(T \cap scope(G_1, G_N, E, S) = T \cap mxdbag\(S\), \(G_1, G_N)\). Considering the producibility of metabolites for deciphering exchanges makes it possible to discriminate between models. In Figure 1, the three symbionts possess a reaction of interest that would enable the producibility of the target by the host when adding transports.

2.3 Mixed-bag selection of community

Selecting a sub-community following the mixed-bag model entails picking a minimal number of organisms, such that they can collectively meet an objective regardless of transport reactions and exchanges. The search space \(C\) is the set of the \(2^N\) sub-families of the whole community. We select organisms in the mixed-bag community to be added to an empty system (or a system with a host) such that a maximum of its targets are producible, by maximizing the size of \(T \cap mxdbag\(S\), \(G_1, G_N)\) among all possible sub-communities \(\{G_1, \ldots, G_N\} \subset \{G_1, \ldots, G_N\}\). This selection of reactions must occur in a minimal number of organisms, hence the minimization of the number of species \(L\) in a second step. Optimal community is defined by successively solving these two problems:

\[
mxdbag\text{Cnity}(S, T, G_1, G_N) = \arg \min_{\{G_1, \ldots, G_N\} \subset \{G_1, G_N\}} \left( \frac{\text{size}(T \cap mxdbag\(S\), \{G_1, \ldots, G_N\)))}{\text{size}(\{G_1, \ldots, G_N\})} \right).
\]

For instance, in Figure 1 (a, b), there are three different ways to produce target \(T\) from the medium \([A, B]\) according to a mixed-bag framework: the host can be combined with any of the three symbionts 1, 2 and 3, which all possess the reaction \(R_4\), producing \(E\) from \(C\) and \(D\) as the mixed-bag formalism is subjected to a small amount of constraints due to the absence of transport or exchanges modeling, it can be used as a first study of the community and to pinpoint members of interest for experimenters. Eng et al. (2016) implemented CoMiDA, an Integer Linear Programming (ILP) algorithm for solving this problem with a network-flow formalism. They tested it on 10,000 random pairs of seed and target singletons to identify a minimal community. However, CoMiDA, like the other
problem-solving techniques, proposes a single solution to the gene-soup sub-community problem, thus preventing the experimenter from catching the global and possibly complex combinatorics of the problem. Topological modeling with Answer Set Programming (ASP; Gebser et al., 2012) has solving assets, as it enables us to sample the whole space of the solutions and thus provide enumeration of optimal solutions or their intersection-union. The foundations of an ASP-encoding enabling the identification of optimal communities is depicted in Listing 1.

Listing 1 Selection of minimal-size communities in a mixed-bag framework

i. \{sel_orga(B): orga(B)\}.
ii. mixedbagScope(M):= seed(M).
iii. mixedbagScope(M):= sel_orga(B); product(M, R); reaction(R, B);
mixedbagScope(M2): reactant(M2, R).
iv. #minimize \{1@2, T: target(T), not mixedbagScope(T}\).
v. #minimize \{1@1, B: sel_orga(B)\}.

2.4 Exchange-based selection of community

The mixed-bag modeling should be considered as a method for globally studying the metabolic complementarity of community members. A limitation of the mixed-bag modeling is that it does not take into account the required exchanges needed when setting back the boundaries of the metabolic models. A natural motivation is that it is energetically demanding to export or import metabolites, hence a parsimonious hypothesis of exchange dependencies in organisms. Julien-Laferrière et al. (2016) have introduced an algorithm for the selection of synthetic community based on the minimization of exogenous reactions and transport reactions. Without much information on precise transportable mechanisms, as it is the case for many non-model organisms, the size of the search space consisting of all possible exchanges is \(2 \sum_{i<j} \text{size}(M_i,M_j)\). This makes the search for minimal exchanges an intractable problem for microbiomes with a high number of organisms, and particularly if their models contain a large set of shared metabolites. This occurs a lot in models that are built automatically from metagenomics as they cannot be individually curated and improved.

To solve this issue, we introduce a heuristic to approximate the exchange minimization problem, which entails filtering minimal size communities with criteria based on the number of exchange compounds. This is modeled using an optimization problem chaining three combinatorial optimizations: maximizing the number of produced targets in the community under mixed-bag assumption (as seen before), minimizing the size of the community (as seen before) and, then, minimizing the number of exchanges by considering organisms boundaries again. Therefore, the family of optimal communities in the compartmentalized setting is formally defined as follows:

\[
\text{cptCnity}(S, T, G_1, G_N) = \arg \min \left\{ \text{size}(T, \text{mixedbagScope}(G_{i}, \ldots, G_{n}, S)) \right\},
\text{size}(G_1, G_n),
\{E \subseteq \text{exchg}(G_1, \ldots, G_n) \}
\cap \{G_1, \ldots, G_N\}
\begin{cases} T \cap \text{cptScope}(G_1, G_n, E, S) \\ T \cap \text{mixedbagScope}(G_{i}, \ldots, G_{n}, S) \end{cases}
\]

Such optimal communities with their associated minimal sets of exchanges can be identified using the ASP programming paradigm with an extension of the previous encoding, and applied to communities pre-selected with the mixed-bag framework. The predicate escope is introduced to recursively compute the scope of the compartmentalized model associated with a selected family of organisms and exchange reactions (Listing 2).

Listing 2 Selection of minimal-size communities in a compartmentalized framework, together with a minimal size of exchange reactions

i. \{sel_orga(B): orga(B)\}.
ii. \{exchanged(M, O1, O2): metabolite(M, O1); metabolite(M, O2); escape(M, O1); sel_orga(O1); sel_orga(O2); O1!=O2\}.
iii. escope(M, O):= seed(M); sel_orga(O).
iv. escope(M, O):= exchanged(M, O), sel_orga(O).
v. eescape(M, O): - product(M, R); reaction(R, O); sel_orga(O);
escape(M2, O): reactant(M2, R).
vi. #minimize{1@3, M: target(M), not eescape(M, O)}.
vii. #minimize{1@2, B: sel_orga(B)}.
viii. #minimize{1@1, M, O1, O: - exchanged(M, O1, O)}.

This approach discriminates the three solutions obtained with the mixed-bag modeling in Figure 1. Cooperation between the host and symbiont 1 requires the latter to provide the compound E (that belongs to the individual scope of symbiont 1) to the host in order to activate the production of E. Alternatively, the host can be combined with symbiont 2 by providing it the C compound in order to produce E, which can be transferred to the host in return. Finally, the host can be combined with symbiont 3, by providing it with C and D compounds in order to produce E, which activates the production of E. The host-symbiont 1 community is the optimal one to restore productivity of T.

2.5 A workflow for the target-based screening of a microbial consortium associated with a host

The Miscoto Python tool (Microbiome Screening and COmmunity selection using TOpology) encapsulates the previous ASP encodings and takes as input a family of metabolic networks each corresponding to a symbiont, a metabolic network associated with a main species called host (optional), a set of seeds depicting the medium compounds and a set of targets depicting the expected products. For the sake of an exhaustive target study, the family of targets can be set to be equal to all cytosolic compounds of the host, or the whole set of compounds in the microbiome and host.

The first step of the analysis of the microbial consortium (Fig. 2) is a feasibility analysis. It entails identifying the added-value of the symbionts for the production of metabolic compounds of the system. Targets are classified according to three criteria: unsolvable target function (the target is not producible by the host associated with all its symbionts)—trivial target function (the target is producible by the host or, if no host is provided, a single species of the community)—community target function (the target is producible by the host only when it is combined with one or more symbionts in the community or, if no host is provided, two or more symbionts). To that end, a mixed-bag framework is sufficient.

As a second step, for each community function, the algorithm depicted in listing 1 allows for the computation of the minimal size of the community required to activate the function. The third step is an estimation of functional robustness. Intuitively, we can expect that the more complex (poor distribution of enzymes of interest among the bacteria, large set of targets) the metabolic objective is, the lower the number of minimal communities there should be to activate a function within a microbial community. In addition, enumerating solutions in synthetic community design can provide experimenters with alternatives for countering biological incompatibilities between bacteria (Julien-Laferrière et al., 2016). To that end, reasoning-modes of ASP are used to estimate the functional robustness associated with a family of selected functions: first, the union of species involved in at least one minimal community required to activate a function is easily computed with a brave enumeration mode to estimate the range of redundancy associated to the function. All minimal communities associated with the targeted function can then be enumerated if needed.

The last step is related to the identification of minimal exchange communities among the ones previously identified in step 2.

Minimal-exchange communities can also be enumerated for a biological-expert analysis. This allows us to select communities with a minimal number of required transports in order to decrease the expert-based analysis of exchanges needed to assess their biological feasibility.

Performances depend on the complexity and amount of species involved in the experiment. As an example, the first step combined with the enumeration of minimal-size communities (mixed-bag) lasted around 300 s for the study of Recon and the 773 bacteria described in the results on a personal computer. The combination of the second and third steps lasted 100 s on average for each individual target. As runs are independent they can easily be parallelized. Finally, the identification of minimal exchanges (and their union) lasted around 200 s per target, less than 4 h for all targets combined in one run.

Notice that a key point for the applicability of the workflow is that it relies on a three-level modeling strategy: feasibility is based on a scope computation within a global metabolic model; functional redundancy is addressed with a mixed-bag metabolism in which all metabolic enzymes are shared and exchanges are costless; cooperation processes rely on a compartmentalized framework. This strategy is motivated by computational issues: a direct computation of relevant communities by minimizing both the number of bacteria and the number of metabolic transports among them is not conceivable due to the high combinatorics of the problem. For instance, in the HMP benchmark studied in the Results Section, due to the large overlap of metabolites in the 2051 models integrated into the benchmark, examining exchanges during the community selection step without any a priori on the bacteria requires considering $2^{10^7}$ possible transports and makes it impossible to identify minimal communities. On the other hand, considering that exchanges can occur in communities without computing them as a first-line objective enables us to globally analyze the benchmark.
3 Results

3.1 Bacterial complementarity in the HMP

3.1.1 Design of the analysis workflow

The HMP Consortium stool sample (Human Microbiome Project, 2012) was the first large dataset to describe the abundance and variety of bacterial functions in the gut. It represents a valuable resource for exploring microbial cooperation in the gut metabolism. Eng et al. (2016) produced metabolic models for the HMP data. Each of the 2051 bacteria has its own metabolic model, the union of all consisting of 3606 unique reactions. The average size of each metabolic model is 1096 reactions. A benchmark was established to validate their flow-inspired algorithm by selecting a minimal community of bacteria to produce a target metabolite from a seed one. Ten thousand random pairs of two metabolites (one seed, one target) were randomly tested. The authors used a ILP algorithm and several formalisms, of which the most constrained was a bipartite graph modeling, equivalent to the mixed-bag framework introduced in Section 2.2. The goal of the algorithm was to enable the production of the targeted compound, assuming that the seed was considered to be the only available compound for the bacteria in the consortium to activate their reactions. The authors showed that species could be selected to produce the desired target from the unique seed in less than 5% of the cases in the benchmark, thus concluding that in most cases among the ones tested, no path exists between the seed and the target.

Miscoto allowed the analysis of Eng et al. (2016) to be pursued by designing a pipeline that enables an exhaustive exploration of the previously benchmarked dataset (available on demand to the authors). The pipeline is depicted in Figure 2. We first calculated the metabolic feasibility by a microbial community of all possible 4 948 400 seed/target pairs of metabolites, which we call functions. We then studied the functional redundancy by sampling functions associated with a community—i.e. requiring at least two bacteria to be met—and exhaustively enumerated all minimal solutions for each function that required at least three bacteria. Ultimately, we focused on cooperation processes and showed that exchange computation can discriminate between size-minimal communities by identifying minimal metabolic exchanges required within the selected bacteria.

3.1.2 Exhaustive feasibility study of the HMP seed/target functions

Figure 3 depicts the results of the exhaustive study of feasibility of all of the 4 948 400 functions in the HMP dataset, that is, all possible pairs combining any single seed from the microbiota compounds associated with any single target from the microbiota compounds. We classified functions in three categories: unsolvable functions that can never be reached regardless of the selected community, trivial functions that can be met by a single bacterium and community functions that can only be met through bacterial cooperation (two or more bacteria). Our analysis demonstrated that 76.8% of functions are unsolvable in the benchmark, which is expected, as we only allow one metabolite to initiate metabolic reactions, i.e. the unique seed. We notice however that feasible functions among the complete family of seed/target functions are five times more frequent than estimated in the random benchmark of Eng et al. (2016), militating in favor of exhaustive studies of feasibility with a complete view of the producibility capabilities of microbiomes. Our tool also evidenced that 15.0% of the functions are trivial i.e. intrinsically met in one metabolic model. The remaining 8.2%, or 405 473 functions, depend on two or more bacteria: they are community functions (Fig. 3a). This analysis illustrates that

![Fig. 3. (a) Exhaustive study of the feasibility for the 4 948 400 seed/target functions associated with the HMP dataset. (b) Community size computation for 10% of community functions (i.e. 40 000 seed/target pairs associated with a community of size 2 or more): all communities have size less than 6, the most frequent case being size 2](https://academic.oup.com/bioinformatics/article-abstract/34/17/i934/5093211/318x645 to 326x651)

Miscoto can efficiently classify functions by ensuring an exhaustive exploration of a large-scale microbiome and point out complex community functions.

To have a better insight of the size of the community associated with community functions, we randomly selected 10% of them (40 000 community functions) and used both our Miscoto tool and the network-flow CoMiDA algorithm (Eng et al., 2016) to identify a minimal community of bacteria that enables us to complete them. Both tools reported similar results, confirming that the scope-based semantics encoded in Miscoto and the bipartite-graph model implemented in CoMiDA are equivalent. We observed that 86.8% of community functions are simple: two bacteria cooperating are enough to restore the function (Fig. 3b). The maximal size of communities needed to restore a function is equal to 6. There are 5301 (13.2%) complex community functions of size three or more, with most of the functions depending on three bacteria (12.2% of the benchmark, i.e. 4881 functions), and 420 being of size 4 or more. Together, this analysis suggests that a very low percentage (1% provided that the sampling of complex community functions was representative, at most 7% in all cases) of the whole set of seed/target functions is made feasible with communities of three or more bacteria.

3.1.3 Exploring the whole space of solutions demonstrates functional redundancy of microbiota metabolism

We used the enumeration capability of ASP-based methods implemented in the Miscoto tool to enumerate minimal community solutions for all complex functions (associated with three or more bacteria) identified in the 40 000 functions sample, i.e. 5301 ones. The number of minimal communities per function ranged from 2 to 1 506 662. As shown in Figure 4a, 86.5% of functions generated more than 100 solutions, of which 49.8% generated more than 1000 solutions. The median is 977 solutions per function. This illustrates the high combinatorics of the community solving problem. In terms of biological interpretation, the number of minimal communities associated with a function can be regarded as a pointer to functional redundancy: a function associated with a large number of minimal community solutions in the microbiome is more likely to be effective than one with a very few number of solutions because several bacteria are able to play an equivalent role with respect to the function restoration (Moya et al., 2016). To confirm this hypothesis, for each seed/target function in our sample, we computed the number of bacteria involved in at least one minimal community restoring the function, i.e. the union of bacteria (Fig. 4b). Our analysis suggests that the number of solutions is linked with the number of species involved in a solution with a polynomial relation (Fig. 4c).
In this respect, computing the latter is much less time-consuming than the former and is easier to study than a large set of enumerated solutions (Fig. 4b) as the median size of the union is 68.5. This promotes the use of the quantity of bacteria involved in at least one minimal solution to the function restoration problem. When screening a large set of targeted functions, this criterion enables us to sort functions according to their redundancy and determine whether it is worth enumerating the whole set of minimal communities.

### 3.1.4 Identification of exchanges to discriminate minimal-size communities associated with a given function

In order to get a better insight into the variability of cooperation processes involved in minimal-size communities, we further developed the analysis by selecting a panel of 130 functions for which the minimal size of solution was three bacteria and which displayed a number of community solutions ranging from 100 to 1000, which is the main range identified in the enumeration study. We noticed that the number of minimal exchanges ranges from 2 to 8 for communities optimizing the producibility of one function (Fig. 4d). For 94% of all functions, there is at least one community solution with at most four exchanges. 38.7% of the functions can be met by at least one community associated with two exchanges only: they correspond to the simplest exchanged-based functioning for a community of three species, where two bacteria each provide a single precursor to a third one. The functioning of 60.9% of the remaining functions can be explained with three exchanges, suggesting that exchanges within several bacteria (e.g. the production of two precursors by a bacterium or a cycle system between two bacteria) are needed to make the function effective. The other cases correspond to more complex cases with multiple exchanges within the consortium.

As shown in Figure 4d, only 17% of the studied functions are associated with minimal-size communities that all depict the same number of exchanges (monocolor vertical bars in the picture). For the 83% non-homogeneous functions, focusing on communities with a minimal number of exchanges reduces the number of communities to be explored by 45%; these minimal-exchanges communities are depicted with the lowest color segment in each vertical bar. Focusing on these minimal-exchange communities allows us to reduce the average family of bacteria (union) involved in the possible communities from 43 to 30. This suggests that adding an exchanged-based criterion to the community-size optimization criterion may facilitate the selection of strains associated with a targeted function by reducing the number of relevant species to be investigated closely.

### 3.2 Recon 2.2 and gut microbiota complementarities

For the sake of application and illustration, the Miscoto tool was applied to study cooperation potential between the Recon 2.2 genome-scale metabolic model of the human metabolism (Swainston et al., 2016) and 773 gut microbial models [Magnúsdóttir et al., 2016], available on: vmh.uni.lu]. In order to globally analyze and screen the cooperation within these organisms under fixed nutritional conditions, nutrients were voluntarily restricted to 51 compounds of the Dulbecco’s Modified Eagle’s growth medium (DMEM). This medium mimics a type of cell culture conditions that can suit enterocytes (intestinal absorptive cells) and has been shown to enable some bacterial growth (Biedermann et al., 2014).

We set all cytosolic metabolites of Recon as metabolic targets, i.e. a set of 1920 compounds, having in mind that this target list could be refined in the future, following an intestinal version of the human metabolic network. We noticed that 831 compounds are producible from the DMEM growth medium, whereas 1451 metabolites are producible from native modeling conditions of Recon2.2 using as seeds all boundary compounds for which imports to the extracellular space and cytosol exist. A feasibility analysis with Miscoto highlighted that cooperation with the gut microbes may facilitate the producibility of 46 additional target metabolites with the
DMEM growth medium and 24 with the native modeling conditions of Recon. This confirms that the medium is very restrictive and that bacteria have an increased added value on the host in a limited growth medium rather than in the native modeling conditions.

The minimal-size community identification implemented in Miscoto evidenced that, in cooperation with Recon, 381 different communities of three bacteria are enough to enable the producibility of the whole set of 46 targets. Between 42 and 48 exchanges may be needed to make this producibility effective, mostly from the bacteria to the host. Only 11.5% of the gut microbiome, that is to say 89 bacteria, play a role in the at least one of these 381 communities with a cooperation potential. We identified a unique partition of the 89 bacteria in three disjoint clusters of 58, 15 and 16 species such that each of the cluster minimal communities comprises exactly one species from each cluster. We noticed however, that the bacteria are far from playing equivalent roles within each cluster: each species in a cluster is associated with only few species in the two others. This suggests that bacteria do not have equivalent roles regarding the individual producibility of targets.

In order to elucidate both the role of the three clusters of bacteria and the individual role of species in each group, we screened the impact of the 89 bacteria over the 46 individual targets with Miscoto. The feasibility analysis highlighted that the producibility of each individual target can be restored by at least a single bacterium. Figure 5 depicts the connection between each target and each bacterium. It highlights that some targeted compounds can be produced by only very few species, leading to further insights into the 381 optimal communities. Taken together, this analysis illustrates that the role of the different bacteria in the production of a multitarget function by an optimal community can be elucidated by using the screening of individual target feasibility with the Miscoto tool.

4 Discussion

Selecting communities in large microbiotas has a large spectrum of applications ranging from the understanding of complex eukaryote dependencies to their symbionts to the design of synthetic communities for industrial applications. In any case, having access to the whole of systems satisfying the desired objective is an asset, should additional criteria or biological incompatibilities be taken into account a posteriori. In this paper, we presented the Miscoto tool and its use to perform a wide and efficient screening of microbiomes in the context of community selection by combining size and exchange combinatorial optimizations. Miscoto enables the classification of metabolic producibility in terms of feasibility, functional redundancy and cooperation processes involved. By providing, as a first step, a feasibility analysis in the microbiota, it can assess the global added value of any microbial cooperation with regards to a target producibility objective. Identified community functions will require
cooperation to occur, either between a host and at least one symbiont, or within a group of two or more bacteria. These functions can be further analyzed to identify all minimal-size communities that can meet them. We showed that the union of symbionts that appear in at least one solution is helpful for determining the redundancy of the function in the microbiome. The following step of exchange minimization within minimal-size communities forms a means for classification of these communities. The application of this workflow to the HMP data constitutes, to the best of our knowledge, the first exhaustive and large scale analysis of cooperation capabilities in a microbiome dataset.

A first key feature of the method is to address scalability in large datasets by minimizing the size of the community prior to the exchanges. Let us emphasize that our approach is particularly suited for screening large-sets of bacteria and compounds. Once the family of bacteria has been drastically reduced, it becomes possible to skip the minimal-size criteria and directly run the exchange minimization-based criteria to have an exhaustive view of relevant exchanges within a small-scale community. A limitation of the method could occur in some cases, e.g. the selection by Miscoto of a 2-bacteria community with 3 minimal exchanges while there also exist a 3-bacteria community with 2 exchanges.

A second key feature of the method is that it entirely relies on a combinatorial modeling of metabolism based on bipartite graphs. The producibility is checked according to a recursive definition of the propagation (scope) which does not take into account the stoichiometry of reactions. This choice was motivated first for computational reasons, since the combinatorial framework makes the producibility criteria monotonous: adding a reaction to a network can only increases the scope (Prigent et al., 2017). The enumeration of all solutions is performed in a reasonable time with SAT-based techniques. Notice, however, that is is possible to filter a posteriori the family of reported communities with flux-based criteria.

As a last feature, our framework does not take into account the potential competition, incompatibilities in cultivating the proposed bacteria together as raised by Julien-Laferrière et al. (2016), nor the possibility that key metabolites could be degraded (Eng et al. (2016)). The ASP paradigm we use is flexible enough to include any knowledge about strain compatibility. However, we consider that not all experimental parameters can reasonably be formalized in such large scale datasets. Therefore, for the sake of genericity, we advocate for the identification of all optimal models without a priori considering experimental knowledge. Furthermore, metabolic transport identification is a challenge in metabolic models that highly depends on genome annotation quality. There usually are few metabolites for which transport between the cell and the extracellular space is assessed in non-model species without an extensive review of the literature (Sung et al., 2017) although efforts are made to compute them automatically (Dias et al., 2017). Community solutions that would require unexpected transports can be discarded a posteriori. For others, an additional optimization is an option, by prioritizing exchange of metabolites for which transport is identified in the metabolic models. More generally, the use of any criteria that are context-specific can be done a posteriori to rank the communities to be tested without leaving any optimal solution aside. In particular, such considerations have to be acknowledged when selecting a subset of a microbiome for experimental verification that a function performed by the host requires cooperation.

In addition to the selection of microbial communities, let us point out that the search for exchange interactions can also be of strong use in the process of metabolic network reconstruction and genome annotation, particularly when it is unclear whether a host can carry some metabolic functions by itself or if it relies on its symbionts to do so (Dittami et al., 2014). Gap-filling steps performed in model reconstructions risk overfitting when they do not systematically propose associated genes to the added reactions (Pan et al., 2018). This raises questions related the existence of some associated functions in the organism compared to its dependency on an obligate symbiont to execute it. Such reactions are included to meet a defined objective but in a context in which the organism is self-sufficient to meet it, which is a substantial hypothesis when facing difficulties with growing a non-model organism in axenic conditions. This shows the need to work with possibly incomplete models and, if possible, check the added-value of the microbiome before adding putative reactions with no genetic genetic support to restore functions (e.g. biomass production).

To conclude, we see our community selection workflow as an intermediary step in microbiome studies prior to quantitative constraint-based simulations (Hanemaaijer et al., 2017; Heinken et al., 2015a; van der Ark et al., 2017) in order to handle the combinatorics of community selection in large microbiotas.

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