Modeling approaches for probing cross-feeding interactions in the human gut microbiome

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ABSTRACT

Microbial communities perform emergent activities that are essentially different from those carried by their individual members. The gut microbiome and its metabolites have a significant impact on the host, contributing to homeostasis or disease. Food molecules shape this community, being fermented through cross-feeding interactions of metabolites such as lactate, acetate, and amino acids, or products derived from macromolecule degradation. Mathematical and experimental approaches have been applied to understand and predict the interactions between microorganisms in complex communities such as the gut microbiota. Rational and mechanistic understanding of microbial interactions is essential to exploit their metabolic activities and identify keystone taxa and metabolites. The latter could be used in turn to modulate or replicate the metabolic behavior of the community in different contexts. This review aims to highlight recent experimental and modeling approaches for studying cross-feeding interactions within the gut microbiome. We focus on short-chain fatty acid production and fiber fermentation, which are fundamental processes in human health and disease. Special attention is paid to modeling approaches, particularly kinetic and genome-scale stoichiometric models of metabolism, to integrate experimental data under different diet and health conditions. Finally, we discuss limitations and challenges for the broad application of these modeling approaches and their experimental verification for improving our understanding of the mechanisms of microbial interactions.

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1. Introduction

The gut microbiota harbors hundreds of different microbial species that actively metabolize dietary and host-derived substrates, with an evident influence on host health [1-3]. Next-generation sequencing has greatly increased our understanding of the composition of the microbiota. 16S rRNA and metagenomic studies have allowed scientists to evaluate the structure of this community in terms of relative abundance of species (richness), intra and inter-diversity, and metabolic functions [4]. Advances have been made to understand the assembly process in the infant gut microbiota [5], characterized by a succession of microorganisms, the wide diversity in microbiome compositions in the adult and elderly, the impact of antibiotics on this community, and its resilience [6,7]. These advances have contributed to new hypotheses or validations of the role of the gut microbiota in disease.

Unfortunately, this static view does not take into account all the interactions that shape the composition of a microbial community so complex such as the microbiota. With hundreds of gut microorganisms inhabiting (or transiting) this ecosystem, active at disparate rates, present at different abundances and colonizing different niches, the extent of microbial interactions is enormous and a challenging task to analyze or quantify. At least 30,000 interactions could be counted at a specific time in the gut microbiota [8]. Conversely, current experimental set-ups to study microbial interactions have analyzed only small synthetic consortia up to 25 gut species [9,10]. There is evidently a gap in our understanding of microbial interactions, which has raised the interest in top-down systems biology approaches to simplify this tremendous complexity.

Microbial interactions vary in their specificity and cost resulting in different ecological outcomes [11]. Microbe-microbe interactions in the gut include quorum-sensing [12], a process little studied in gut microbes. Genomic analysis of gut microbes suggest the intestine is dominated by interference competition [13], with a high abundance of genes producing antimicrobials such as bacteriocins and Type 6 Secretion Systems (T6SS) [14]. The human host has several ways to modulate its microbiome, such as immunoglobulin A coating, secretions, or antimicrobial peptides [15,16]. The forces influencing microbiome profiles are largely unknown. However, physiological factors such as intestinal oxygen concentration, peristaltic movements, and innate immunity effectors are evident [17]. Moreover, microbes are not homogeneously distributed in the intestine, displaying a concentration gradient, and they usually prefer a luminal lifestyle or residing near the mucus layer [18].

Diet is one of the main factors dictating which species will be dominant in the microbiota. Major bacterial groups and patterns regarding the fermentation of complex dietary polysaccharides have been described [19]. These include microbes able to access complex glycans and other species that can capture degradation products (monosaccharides, amino acids) or fermentation byproducts. Then cross-feeding takes place, which can be understood as the metabolic exchange between microorganisms. The exchange of short-chain fatty acids (SCFAs) such as acetate, and organic acids such as lactate and succinate, is a common metabolic interaction [20,21]. Other molecules such as amino acids, vitamins, or branched SCFAs also participate in metabolic cross-feeding [22].

While cross-feeding interactions are at the core of the gut microbiome network [8,9,23], there is a lack of quantitative studies revealing their actual prevalence in the gut. Among other approaches, kinetic models based on ordinary differential equations (ODE) have been used to quantify the extent of interactions and influence of certain gut microbes, among others [24]. An approach of preference are genome-scale metabolic models (GSMMs). These are mathematical structures derived from genome-scale network reconstructions [25], described by all the reactions associated with metabolic genes in the organism’s genome. GSMMs are complemented with experimental data about cellular composition, growth capabilities, and consumption/production profiles to represent the metabolic phenotype [26]. By using appropriate optimization procedures [27,28], cellular growth and the exchange rate of metabolites can be quantified, which can be later validated using co-culture studies [29].

The cross-talk between top-down and bottom-up systems biology approaches, modeling, and experimental approaches is currently being used to provide a rationale for complex diseases. They have been applied to determine the alteration of cross-feeding patterns in inflammatory bowel disease (IBD) [30] and Parkinson’s disease (PD) [31,32] and to design therapeutic approaches towards health improvement through microbiota modulation [9]. Ultimately, through the integration of increasingly broader and deeper omics datasets, modeling approaches enable the exploration of explanatory hypotheses with varying degrees of complexity that can be later verified, significantly reducing the amount of experimentation, and yielding novel insights about the interaction mechanisms. However, there are several challenges and opportunities in this field. This review summarizes recent advances in the ecology and modeling approaches to study cross-feeding in the gut microbiota.

2. Experimental studies of cross-feeding

The study of cross-feeding interactions has been supported by improvements in microbial isolation techniques and deposition in culture banks [33]. Most gut microbes are highly oxygen-sensitive and present unpredictable metabolic requirements. One example is KLE1738, a gut isolate that solely uses Bacteroides-derived GABA as a carbon and energy source [34]. Cross-feeding experimental set-ups often use a catalog of potentially representative gut microbes that should be simple to culture, traceable, and representative or relevant to their ecosystem [35]. Current experimental models to study cross-feeding include microwell assays, batch fermentations with a small number of microorganisms, and fecal inocula in bioreactors [35]. Advanced systems such as SHIME (Simulator of Human Intestinal Microbial Ecosystem) provide global information and general deductions regarding dietary alterations [36]. Continuous bioreactors have also been studied, with the advantage of reaching steady-state and inflow and outflow rates similar to human digestion [37]. Advances in multi-layer microfluidics in vitro systems have enabled satisfactory mimicking microbial processes along the gastrointestinal tract [38-41]. There are still limitations with the accurate simulation of host-related responses, e.g., continuous removal of cross-feeding metabolites, enzyme and acid secretions, and immune surveillance. However, current systems have shown promise for a representative simulation of human-microbe interactions under relevant conditions [41].

Metabolic interactions in the gut microbiota usually involve a complex carbohydrate and an SCFA, although amino acids have also been shown to participate [42,43]. For example, human milk oligosaccharides (HMOs) promote cross-feeding of glycan degradation products between Bifidobacterium species, thereby promoting the colonization of butyrate-producing bacteria in the newborn (Fig. 1) [5]. Host-derived molecules, such as mucin glycans and bile-salts, can also support metabolic exchanges [22,44-48]. Butyrate production derived from microbial fermentation networks balances pH, provides immune system stimulation and reduces the risk of infections [2]. Bacteroides spp. are among the most studied bacteria for their broad preference for dietary glycans such as...
xylan, pectins, and starch [49]. Their degradation process usually results in cross-feeding interactions. Inulin is a prebiotic, a fructose polymer of up to 60 molecules. Several members of the gut microbiota have the machinery for its consumption, including *Bifidobacterium*, *Roseburia*, *Bacteroides*, and *Lactobacillus* species [50]. Inulin-derived fructans, as well as SCFAs derived from inulin fermentation, are well known for promoting the growth of butyrate-producing bacteria such as *Faecalibacterium prausnitzii* [51]. This stimulation is mediated by acetate and lactate. Other important cross-feeding currencies are formate [52] and H₂ [53], which could be used by methanogenic gut archaea (*Blautia hydrogenotrophica*). Notably, there are indications of complementary functions as evidenced by pathway parts (e.g., related to nitrogen metabolism) expressed by different human gut bacteria [54]. Fig. 1 provides examples of cross-feeding interactions in the gut microbiota.

3. Metabolic interactions and ecology based-modeling

Metabolic cross-feeding can be understood as the exchange of metabolites between microorganisms. It differs from true cooperation in that the production of metabolites does not represent a fitness cost for the producer, i.e., they are byproducts or the result of macromolecule degradation, and therefore their production does not need additional energy [55]. Cooperation is associated with more dependent interactions, where one microorganism invests part of its energy in synthesizing a metabolite that is necessary for another, concomitantly receiving a benefit for this production. An interesting example of true cooperation is presented by *Bacteroides ovatus*, a gut commensal that degrades several polysaccharides such as inulin, amylopectin, and xylan [56]. Inulin degradation by this microorganism requires two putative extracellular xylanases (BACOVA_04502/04503). Mutant analyses revealed that these enzymes do not provide a direct benefit for *B. ovatus* in vivo [56]. The expression of these enzymes represents a cost for the producer *B. ovatus* [56]. Interestingly, mutant analyses also showed that *B. ovatus* receives a benefit in a more complex microbiota mediated by these enzymes. However, most *Bacteroides* species do not engage in cooperative metabolic interactions and maximize intracellular glycan degradation instead [57,58]. A similar observation is found in the infant gut microbiota, where two glycan degradation strategies have been characterized between *Bifidobacterium longum* subsp. *infantis* and *Bifidobacterium bifidum* [59]. These interspecies interactions could be classified as true cooperation involving the production of costly proteins that benefit the community. Some studies predict cooperation to be unstable in steady-state and probably not dominant in complex ecosystems [4]. However, this view has been challenged by other studies [55,60,61].

Exploitative competition is characterized by limited resources that reduce the growth of two or more microorganisms.
Exploitative competition and metabolic commensalism were predicted as the most prevalent interactions among 773 gut microbes [62]. Competition for cross-feeding substrates such as monosaccharides, oligosaccharides, and SCFAs is common in the gut microbiota, even for taxonomically unrelated microorganisms [62]. Competition usually results in two possible outcomes: niche partitioning or competitive exclusion [63].

A dense cross-feeding network is predicted in the gut microbiome [62]. Exchanged metabolites derive from the fermentation of complex dietary carbohydrates, proteins, or host secretions. Other released metabolites are fermentation end-products that report no cost for the producer, but a clear benefit for the user. Anaerostipes cacaee, a prominent butyrate-producing bacterium in the gut microbiota, presents a product yield five-fold higher in butyrate from lactate, a cross-feeding metabolite, compared to glucose [9,64]. Sometimes this metabolic interaction could be bidirectional (both microorganisms receiving a benefit), or unidirectional instead [21]. Indirect benefits for the release of cross-feeding metabolites could be the removal of potentially toxic byproducts or continuing microbial fermentation.

While cross-feeding interactions are widespread in the gut microbiota, in vivo they are limited by distance [65]. Four Lactococcus lactis strains with different metabolic capabilities were cultured in a 3D system allowing unidirectional cross-feeding (a galactose consumer-glucose producer, a lactose consumer, a glucose consumer, and a competitive glucose consumer). All strains were embedded in agarose beads to monitor glucose exchange by flow cytometry [66]. Combining the experimental results with mathematical modeling of reaction-diffusion processes, a maximum of 15 μm between beads was established as the limit for cross-feeding in an aqueous medium. This study highlights the importance of spatial limitations in metabolite diffusion in cross-feeding. It would be interesting to contrast these findings with co-localization data in biofilms of microcolonies in the gut microbiota. These communities are exposed to other influences such as shear forces, oxygen gradients, and host-derived antimicrobials.

In a more complex set up, intestinal contractions have been shown to influence microbiota composition through mixing [67]. The minigut is a fluid channel mimicking the gut including important variables such as flow rate and mixing. Simple diffusion models were able to capture the microbial behavior in time in this system [67]. Interestingly, the gut microbiota is dependent on feeding/fasting regimes and the host’s circadian rhythm [68]. Therefore, the gut microbiota shows an oscillatory nature in its composition with diurnal cycles [69,70]. Interestingly, jet-lag induces microbiome dysbiosis in mice, characterized by glucose resistance [71]. Microbial metabolites such as butyrate also show daily variations [72]. Some studies have proposed mathematical models to understand the influence of time and space in the gut microbiota [37,73,74].

How do cross-feeding interactions emerge? McNally combined an in silico reductive gene-loss model with GSMMs between two generalist E. coli strains and identified that metabolic interdependencies commonly appear as evolved cross-feeding interactions [75]. Some strains could not develop the ability to cross-feed on certain metabolites being released (missed opportunities). From thousands of trajectories, 35% resulted in a commensal community, 10% collapsed, and 51% were independent. Metabolic interdependencies were found between tyrosine and thymidine. Formate and tyrosine were the metabolites that created the largest dependency from the other partner, while acetate and succinate most usually never created interdependency. In the long term, metabolic cross-feeding permits the co-occurrence of species in certain communities by the emergence of metabolically interdependent groups [76]. This co-occurrence is reinforced by gene loss as an adaptation where one microorganism obtains all its carbon and energy sources from the enzymatic action of others [63].

Are some gut species essential, or in contrast functionally replaceable, in terms of ecosystem stability? The gut microbiome is characterized by functional redundancy [77], supporting stability and resilience to particular interventions. The keystone species concept derives from the ecological theory of Paine [78], which are species so essential for the stability of an ecosystem that their loss or removal results in a collapse of the community. A synthetic microbiome with 14 species, including one butyrate producer, was recently cultured in batch bioreactors, with inulin as a dietary source. Experiments were repeated leaving one species out at a time, to study the impact of individual deletions [10]. All combinations followed similar trajectories in terms of their biomass production and inulin substrate consumption. The community however did not collapse in any of the deletions [10]. SCFAs profiles were however different in each deletion experiment. For example, deletion of Bacteroides dorei incremented acetate amounts, and deletions of Ruminococcus gnavus and Bacteroides vulgatus resulted in higher butyrate concentrations. An increment in a SCFA suggested that the deleted species contributed negatively to SCFA production. Consequently, only three species contributed significantly to butyrate production: E. coli, B. dorei, and L. symbiosum. Besides, negative interactions between bacterial pairs were also identified: E. coli absence allowed a higher growth of Bifidobacterium adolescentis, and the deletion of B. dorei and L. symbiosum resulted in a community dominated by Lactobacillus plantarum [10].

Another recurrent question when analyzing cross-feeding interactions is whether they are organized in modules or form higher-level hierarchies. Functional studies indicate a multi-level trophic organization in the microbiome, with at least three and up to 10 functional groups [19]. Using ecology-based modeling, Wang et al. predicted four iterative, trophic levels of metabolite production and consumption in the gut microbiota [79]. Their analysis started with a manually-curated metabolic reconstruction of microbial interactions and nutrient dependencies [46], containing 567 gut microbes and 235 metabolites consumed and secreted by these microorganisms. Combined with metagenomics data, the number of substrates converted into metabolites feeding other microbes potentially being carried to lower trophic levels was determined. A part of these substrates was also converted into bacterial biomass, and certain byproducts are used by the next trophic level [79]. Their approach also includes intestinal nutrient intake and byproduct release. Intra and inter-person diversity were shown to be unevenly distributed across four trophic levels. As expected, the diversity of metabolites was higher in upper trophic groups and narrower in the lower groups. One conclusion obtained from this analysis was that genus-level competition could explain taxonomic diversity of the microbiome [79].

This ecological platform has been recently combined with machine-learning methods to predict novel, unexplored metabolic interactions in the gut microbiome. The gut cross-feeding predictor (GutCP) builds from the multi-level trophic organization and predicts a theoretical metabolome from metagenomics data and known consumption links [8]. The GutCP algorithm improved the original network, reducing the error between prediction and actual data by adding new consumption links. Undiscovered interactions that result in more accurate estimations would possibly be true cross-feeding interactions. The main features of ecological as well as other modeling approaches are shown in Fig. 2.

4. ODE-based mechanistic models of microbial communities

Computational modeling approaches of microbial interactions are at the forefront of microbial community research [80-82]. They
are helpful to identify interactions and mechanisms that are likely to occur, reducing the burden of extensive experimentation (Fig. 2). They could be classified as static or dynamic models, depending on if they explicitly include information on how the components of a community interact over time. Dynamic modeling approaches describe how a microbial community and its components change over time, enabling the study of variations in the properties included in the model. The first classification of dynamic models is based on the level of detail, from the coarseness of the population-level strain composition of a consortium to detailed models of all metabolic reactions in each cell. A second classification uses the type of algorithm employed to simulate the model. Two of the main types are deterministic models based on differential equations and stochastic models whose behavior depends on randomly selected events. A third criterion serves to classify what is represented in the model. Metabolic models represent the metabolism of the bacteria in the community, often at a molecular level, while in ecological models, strains or species interact, reproduce and change their numbers and behaviors at what can be called cellular level [8,83].

Models based on Ordinary Differential Equations (ODEs) employ a set of mathematical functions to represent microbial communities at different levels of detail. These deterministic mathematical functions describe the changes in quantity or concentration of different species in time. One of the simplest microbial community ODE-based models is the generalized Lotka-Volterra equation (gLV) [84]. Using a small number of parameters such as growth rate and abundances in single and co-culture, it evaluates the effect of one microorganism on another’s growth, usually by estimating an interaction coefficient $\alpha$. Its combination with Bayesian methods has been used to design microbiome consortia. C: ODE-based models simulate mechanistic processes as equations and require extensive parameterization. D: GSMMs combined with community design algorithms enable the identification of cross-feeding metabolites in simple and large networks. The main features and requirements of each approach are highlighted in the blue and orange boxes, respectively.

Fig. 2. Representation of computational biology approaches to interrogate metabolic interactions in the gut microbiota. A: Species deletions simulate and study the impact of the absence of one species in a community, while combinations of literature-based metabolic reconstructions of the gut microbiota and machine learning allow evaluating the hierarchy in metabolic interactions. B: gLV-based models evaluate the impact of one microorganism in another’s growth, usually by estimating an interaction coefficient $\alpha$. Its combination with Bayesian methods has been used to design microbiome consortia. C: ODE-based models simulate mechanistic processes as equations and require extensive parameterization. D: GSMMs combined with community design algorithms enable the identification of cross-feeding metabolites in simple and large networks. The main features and requirements of each approach are highlighted in the blue and orange boxes, respectively.
concentration to zero, and deletion of Desulfovibrio piger, a H₂S producing microbe, significantly increased butyrate concentration. It was later shown that butyrate production by A. caccae is highly sensitive to H₂S and pH. This modeling framework identified how much one single species contributed to growth and butyrate production in the synthetic consortium [9].

Other approaches use ODE-based models where the equations aim to predict microbial interactions based on mechanistic knowledge. For example, in Pinto et al. [87], four equations with 17 parameters were employed to build an ODE system that simulates microbial abundance, substrate consumption, and production of SCFA in a four infant gut species consortium. This approach used modified Monod expressions [88] to model cell growth under various conditions including product inhibition and cooperation. Adding more bacteria to the simulation means the number of equations and parameters increases, giving rise to the so-called combinatorial expression problem when it comes to fixing the values of the parameters that reproduce known experimental data [89]. For relatively small communities with few different types of microorganisms, this problem is manageable with current computational infrastructures and time-resolved measurements.

5. Genome-scale stoichiometric models of metabolism

5.1. Genome-scale network reconstructions

Genome-scale network reconstructions (GENREs) encompass the entire collection of enzyme-catalyzed reactions encoded by genes of known function found in an organism’s genome [90]. As such, they describe its metabolic potential. Starting from the annotation of the organism’s genome, Gene-Protein-Reaction (GPR) relationships are identified, which link the genotype with the metabolic phenotype. Detailed protocols have been developed to standardize and ensure high quality for the reconstruction [90]. Several computational platforms and tools have been developed to leverage the current genomic information and accelerate the reconstruction of these networks for single and multiple organisms in a (semi)automatic fashion [91-93]. These top-down tools integrate functional gene annotation – either performed by these or external tools – using databases like KEGG [94] and BioCyc [95]. Potential gaps in the network, i.e., missing reactions due to unidentified metabolic enzymes, can be resolved using appropriate gap-filling tools [96,97]. There are also bottom-up tools for GENRE construction that employ high-quality template metabolic network(s) for the reconstruction process [97-99]. These tools typically yield higher quality reconstructions, although their performance ultimately depends on the initial network template(s). CarveMe is an automatic tool for the development of metabolic models for microbial communities [98]. It has demonstrated high efficiency and performance for generating a community network of 74 members of the human gut microbiota, and the assembly of a comprehensive GENREs database with over 5,500 bacterial metabolic reconstructions [98]. More recently, novel reconstruction pipelines tailored to use metagenomic data have been proposed for building community reaction networks. Metage2Metabo [100] has shown to yield meaningful community networks of the human gut microbiota capable of respectively enabling community-level flux balance computations and identifying keystone species in terms of metabolic cooperation [100].

The continuous and rapid reconstruction of GENREs of greater quality has led to the emergence of various reconstruction catalogs. AGORA is a comprehensive resource for community modeling, particularly human gut microbiota. It contains over 773 literature-based GENREs for various microbial species [62]. An extension to this resource with over 7200 GENREs of members of the human microbiome, called AGORA2, has been recently developed, and it is under review at the time of preparing this article [101,102]. GENREs from AGORA have been employed to simulate bile acids metabolism in the human gut [103], identify microbial gut alterations associated with Parkinson’s disease [31], characterize host-microbe symbiosis in the mammalian gut [104] and co-metabolism in the human intestine [105]. NJC19 [106] is another reconstruction resource that contains human and mouse microbiome data (6 mouse and human cell types and 883 microbial species) that can be leveraged to construct more comprehensive models of human molecular physiology [107].

5.2. Genome-scale metabolic models and constraint-based methods

GSMMs encode the information contained in GENREs into a mathematical structure – the stoichiometric matrix –, which contains the mass balances for each intracellular metabolite and enables flux calculations. Provided that sufficient external substrate sources (i.e., exchange reactions) and a growth equation are defined and included in this matrix, a feasible flux distribution or metabolic phenotype can be computed. It assumes no accumulation of intracellular metabolites (i.e., steady-state) and requires solving the resulting system of linear equations. As this system is almost always undetermined, i.e., there are infinite solutions that can explain the set of observed exchange reactions, particular flux solutions are computed using optimization or constraint-based modeling (CBM) methods [108]. A popular tool for CBM is the COBRA Toolbox (COntest-Based Reconstruction and Analysis), which includes a diversity of computational methods for genome-scale metabolic modeling and reconstruction and analysis [28].

CBM methods require the formulation of an objective function and definition of capacity constraints (reaction bounds) to compute a feasible steady-state flux solution. A typical objective function is the maximization of cellular growth as a proxy for biological fitness. However, other alternatives have also been proposed when the latter objective does not yield accurate predictions [109]. This method is known as Flux Balance Analysis (FBA), and it has been successfully applied in numerous studies for modeling interactions among multiple cells or organisms [110], understanding human physiology [43,111], disease [112], and studying evolutionary processes [113]. Another commonly used FBA-derived method is Flux Variability Analysis (FVA) [114], which enables the calculation of alternative optima under the assumption of FBA (sub)optimality. FBA-based methods have also been extended for modeling dynamic processes. For instance, dynamic FBA (d-FBA) [115] describes the accumulation of biomass and external metabolite concentrations by coupling the evolution of a set of ODEs for the latter species – parameterized with kinetic expressions (e.g., Michaelis-Menten) – with the iterative solution of FBA problems at each time step. As in FBA, no net accumulation of intracellular metabolites is enforced. Importantly, d-FBA enables the description of the evolution of cellular growth with the corresponding changes in metabolic states and can be naturally extended to model communities.

The modeling capabilities of FBA-based methods are ideal for modeling complex communities where metabolic parameters are scarce. Applications include modeling microbe-microbe and host-microbe interactions such as metabolic cross-feeding (metabolite exchange between species) and competition (uptake of common substrates). For example, FBA shows how fructoselysine is a key cross-feeding metabolite in dual gastrointestinal infections [116]. Nevertheless, FBA-based methods for modeling microbial communities require special considerations, and it has not been until recently that robust methods have emerged. Challenges related to the accurate description of flux exchanges between taxa and
the selection of a biologically relevant objective function must be addressed. One of the earliest CBM methods for modeling community growth was OptCom, which was later followed by its dynamic extension d-OptCom [27], based on the same assumptions of d-FBA. These methods use a multi-level multiobjective optimization approach to optimize the total community biomass while maximizing individual-specific growth rates. Inspection of the directionalities of the metabolic exchange profile enables the determination of possible interactions (e.g., syntrophy, cross-talk, commensalism, and competition). Community FBA (cFBA) [117] and SteadyCom [118] are more recent methods where the community growth rate is maximized, which is the same for all the community members under the assumption of a steady-state balanced growth. As a byproduct of this calculation, the microbial composition that supports this optimal growth can also be found. For instance, the application of SteadyCom recapitulated the known dominance of the bacterial phyla Bacteroides and Firmicutes in the gut microbiota under the average American diet [118]. Notably, SteadyCom can be readily combined with other CBM methods like FVA to determine the abundance and metabolic flexibility of a community under suboptimal conditions. The aforementioned methods assume perfect mixing in the culture, i.e., there are no concentration gradients. This assumption may not be reasonable in some scenarios, for example, in some regions of the human gut [119]. Furthermore, it may hinder the analysis of emergent metabolic interactions due to spatial distribution and concentration gradients [25,120,121].

5.3. Probing microbe-diet-host interactions in health and disease using GSMMs

A key indicator of a healthy gut microbiota health is its composition, which remains fairly stable over long periods even under short-term perturbations [122]. Alterations in the human gut microbiota are collectively referred to as dysbiosis and can be typically linked to certain health conditions such as metabolic, gastrointestinal, and mental disorders [123-125]. A recent study illustrated the predictive power of GSMMs and CBM methods for probing the metabolic behavior of the microbiota of healthy and Parkinson's disease patients [31]. By constraining the relative abundance of the microbiota members derived from 16S rRNA sequencing data of fecal samples, personalized community models were evaluated for their production potential of 129 metabolites under a simulated average European diet. FBA showed that nine metabolites were differentially affected by Parkinson's disease, including methionine and cysteinylglycine. These results were later verified in a different study where four microbial reactions involved in homoserine metabolism (precursor of methionine) were identified to be altered in Parkinson's disease patients [32]. In a recent study, GSMMs were used to build and interrogate personalized brain region models of diseased Alzheimer's and healthy patients [112]. Integration of large transcriptomic datasets and targeted metabolomics data into the GSMMs suggested that some of the bile acids found in brain samples from Alzheimer's subjects originate in the gut microbiota and are then transported to the brain [112].

Another application of GSMMs for probing the consequences of dysbiosis comes from Crohn's disease. The latter condition is characterized by altered blood and fecal metabolome and gut microbiota composition [126]. By using metagenomics data and fecal metabolites from a group of control and Crohn's disease patients, personalized microbiota models were built and analyzed for significant differences in the predicted metabolite secretion profile in the feces and the species contribution to each metabolite under an average European diet [30]. Simulation results pointed to reduced diversity in the number of secreted metabolites due to the reduced microbial diversity in Crohn's disease patients. Proteobacteria were found to increase the potential for amino acids synthesis, which was in line with experimental observations [30].

GSMMs have also been employed to understand the interactions present in a healthy gut microbiome. A collection of metabolic models was built from metagenomic samples from individuals fed on a Western diet [127,128], to simulate cross-feeding interactions potentially involved in SCFA production in the gut microbiota. Analysis of possible metabolic interactions revealed competition as the most common interaction, agreeing with previous reports [129]. A closer examination of the microbiota behavior also suggested potential niche partitioning; either a group of bacteria consumes fibers and starches (e.g., inulin, xylan, and pectin), or another consumes branched-chain amino acids [127]. In another study, a tool called CASINO [42] was developed to quantify the changes induced in the human gut microbiota following a dietary intervention in obese and overweight individuals. Again, 16S rRNA data was employed to personalized the metabolic models [43] and predict metabolic production profiles in plasma and feces. Amino acids and SCFA levels were correctly predicted after the dietary change. These predictions were underpinned by acetate cross-feeding between B. thetaiotaomicron, K. bromii, and B. adolescentis to E. rectale and F. prausnitzii, which in turn supported butyrate production by the latter [43].

6. Other approaches for community modeling

Other dynamic modeling approaches such as stochastic modeling [130,131] or agent-based modeling [132,133] have been applied to interrogate microbial interactions. They use experimental time series of microbiome compositions. Unfortunately, data on microbial interactions at a large scale is usually scarce, and this information is indispensable for this type of modeling [134]. Other approaches help to understand the composition and functions of different communities but do not simulate their dynamical properties. This type of static modeling relies on a wide array of algorithms and computational methods to determine and study the interaction between different organisms, even if most of the time they only evidence the existence of a relationship, not its type. For example, using Bayesian modeling, it is possible to determine the structure/topology of the gene regulatory network of several bacteria living together in the same environment [135]. Other approaches use machine learning algorithms, for example, to determine a network of cross-feeding interactions [136]; classification of host-disease phenotypes from metagenomic data [137,138]; or which bacteria are part of the normal community in certain gut regions [139]. Machine learning has been recently combined with GSMMs, expanding the potential of both approaches [140]. An inherent difficulty of machine learning is that this approach requires relatively large amounts of data to parameterize the algorithm and estimate its performance [141], thus reducing the applicability of machine learning in this field.

Finally, co-occurrence networks could provide important information regarding microbial interactions [142]. They rely on the identification of the co-occurrence of two or more species in multiple samples. In this way, if two species appear in the same sample and are absent in others, they are related. This approach also requires numerous experiments that help determine the species found in a sample, taking advantage of extensive collections available in databases such as IMG/M [143] or datasets like the Human Microbiome Project [HMP] [144]. Co-occurrence has been applied to different conditions, for example, to identify relationships between different taxa [145], predict microbial relationships within and between body sites [142], microbial communities
linked to colorectal tumors [146], obesity, and diabetes type 2 [147], inflammatory bowel disease [148], and asthma [149].

7. Challenges and outlook

Microbial interactions stand out at the inner core of the composition of the human gut microbiota. Their study is hampered by the inherent complexity of this community in terms of functional diversity and forces shaping these cross-talks. For this task, experimental and modeling approaches have shown relevant patterns and highlighted how microbial interactions influence host health and disease.

A significant experimental gap in this field is how little we know regarding the functions and metabolic potential of gut microbes. Their high numbers, differences in spatial and time distributions as well as the large diversity of species, sets a high bar for their modeling. The metabolism of only a few gut species is well known, but for most gut microbes, only general aspects are understood. Moreover, sometimes only one amino acid different is necessary for a gut microbe to inactivate a drug [150]. Besides culturing techniques and microbial collections, high-throughput experimental studies for pairwise interactions are highly demanded, as well as reliable gut simulation platforms closely simulating intestinal processes.

Another experimental gap is how to reduce the breach between synthetic in vitro studies and the actual relevance of microbial interactions in the gut. It is difficult to comprehensively quantify the extent of microbial interactions in the gut or how changes in metabolic interactions contribute to disease, such as IBDs. Besides, it is unclear if the extent of experimentally validated cross-feeding interactions is biologically meaningful. Importantly, cross-feeding is only one of many microbial interactions that could be found in the gut microbiota. The host or other competitive or cooperative interactions would challenge in vitro results. In any case, experimental studies of cross-feeding have been instrumental in deriving general conclusions on which metabolites, groups of microorganisms, or pathways are involved in this process. These validations are the foundations of several modeling approaches, and several of them might not have been prevalent in vivo.

Modeling approaches can significantly help reduce the hypothesis space and offer testable experiments to validate potential metabolic interactions in the human gut. Amongst the various modeling approaches employed for this purpose, we have highlighted ecological, mechanistic ODE-based (kinetic), and genome-scale stoichiometric modeling as valuable tools for probing the interaction space. While ecological and kinetic models are relatively simple to interpret, they are often limited by the amount of data required for their construction (i.e., identifying the relevant model structure) and parameter fitting [151]. It is expected that increasing accessibility to next-generation sequencing, especially metagenomics, contribute positively to this gap. In contrast to ecological and kinetic models, GSMMs are mathematical structures that can be readily reconstructed using high-throughput sequencing data fed to various computational pipelines [91–93]. These model structures are often referred to as “parameter-free”, meaning they do not require to fit parameters but rely on the imposition of constraints derived from the data. The larger the amount of data available, the more reduced the solution space, leading to tighter predictions. While expression, protein, and fluxomic data can be readily used as constraints [109], the use of metabolomic data is not so direct. However, recent approaches have shown promise (see for example, MAMBO [152]). More importantly, GSMMs describe the metabolic potential of the cell and thus, offer unparalleled capabilities for understanding the mechanisms of interaction in great detail. The latter helps explain the surge in the GSMMs use to analyze the gut microbiota [110]. Still, reconstruction of accurate GENRES remains a challenging task, as our knowledge of the diversity in metabolic functions is limited to well-studied organisms. Consequently, the metabolic capabilities of GSMMs are inevitably restricted, which raises questions about the relevance and overall validity of GSMMs for less studied organisms – particularly prevalent in the gut microbiome [153]. Overall, careful consideration and use of GSMMs has to be exercised when modeling the gut microbiota. Recent tools like MEMOTE [154] can help users better understand the quality and coverage of GSMMs.

A critical challenge for modeling the host-gut interactions is the temporal and spatial resolution required to identify critical cross-feeding interactions. The spatial distribution of the gut microbiota is highly heterogeneous as its diversity. Different types of Inflammatory Bowel Diseases (IBDs) can be promoted by an abnormal spatial distribution of the microbial gut members due to diet and inherent host responses, among others [155]. Scenarios resembling some of these conditions can be modeled with GSMMs [25], although they require more sophisticated computational methods that are more far more difficult to implement [120,156,157]. More recently, the COMETS framework has been released as an opensource package enabling the temporal and spatial simulation of microbial consortia using GSMMs in different environments [158]. As more data under experimentally relevant conditions becomes available (e.g., using microfluidics in vitro systems [41]), we foresee that modeling tools capturing temporal and spatial variations in microbial consortia will become more relevant for contextualizing cross-feeding in the human gut.

CRediT authorship contribution statement

Pedro Saa: Conceptualization, Writing – original draft, Writing – review & editing, Funding acquisition. Aries Urrutia: Writing – original draft. Claudia Silva-Andrade: Writing – original draft. Alberto J. Martin: Writing – original draft, Funding acquisition. Daniel Garrido: Conceptualization, Writing – original draft, Writing – review & editing, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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