Investigating a holobiont: Microbiota perturbations and transkingdom networks

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ABSTRACT
The scientific community has recently come to appreciate that, rather than existing as independent organisms, multicellular hosts and their microbiota comprise a complex evolving superorganism or metaorganism, termed a holobiont. This point of view leads to a re-evaluation of our understanding of different physiological processes and diseases. In this paper we focus on experimental and computational approaches which, when combined in one study, allowed us to dissect mechanisms (traditionally named host-microbiota interactions) regulating holobiont physiology. Specifically, we discuss several approaches for microbiota perturbation, such as use of antibiotics and germ-free animals, including advantages and potential caveats of their usage. We briefly review computational approaches to characterize the microbiota and, more importantly, methods to infer specific components of microbiota (such as microbes or their genes) affecting host functions. One such approach called transkingdom network analysis has been recently developed and applied in our study. Finally, we also discuss common methods used to validate the computational predictions of host-microbiota interactions using in vitro and in vivo experimental systems.

Introduction
In addition to our own cells and genomes, humans are hosts to a vast community of microbes. This microbiota are not neutral neighbors, rather they are indispensable contributors to host physiology, acting in a symbiotic relationship with the host. The concept that we are holobionts, being comprised of hologenomes (our genomes and our microbial genomes) with complex interactions between our cells and our microbes, has been discussed for many years. More recently, the development of new tools to interrogate these hologenomic interactions has greatly expanded our ability to understand and define ourselves in terms of both our own human genome and that of our resident microbial partners. The holobiont view helps to explain the dramatic increase in the number of chronic inflammatory and autoimmune diseases with significant genetic component which occurred too rapidly to be attributed to the host genome alone. In contrast, our microbial genomes have a tremendous capacity for rapid adaptations which can influence health and disease. Therefore, in order to better understand the basis of many modern diseases a deeper insight into our interactions with our microbes is required.

Perturbing the microbiota: germ-free and antibiotic-treated mice
Microbiota contribution to specific host phenotypes, including diseases, is frequently determined through broad perturbation of the microbial community (Fig. 1). One extreme, but common method of microbiota perturbation is generation of animals, most commonly mice, devoid of microbes (i.e. germfree or axenic). Axenic and gnotobiotic insect and zebrafish models can also be informative in understanding fundamentals of host-microbe interactions; however, the mouse is the most widely used and powerful tool for understanding the connections between microbiota and disease. Study of germ free versus conventional organisms has revealed myriad roles for microbiota in host immunity and metabolism, among many other systems. For example, it has been long known that in the absence of microbiota, development

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of various immune cells is impaired and the ability to respond to infections is reduced. Germ free mice have also revealed that microbiota have profound influence on metabolism, for example, regulating bile acids pools and worsening glucose tolerance. Furthermore, it has been shown that the gut microbiota can mediate molecular cross-talk between host immune and metabolic functions. Additionally, derivation of specific genetically modified mouse models under germ free conditions has been used to investigate gene-specific interactions between host and microbiota. Although gnotobiotic technologies for germfree animals have existed since the 1950s, our ability to derive and work with germ-free mice is still a limiting factor for investigation. Therefore, an economically more feasible approach was introduced - using antibiotic treatments to alter microbiota composition ideally to the point of eliminating nearly all

![Diagram](image_url)

*Figure 1. An approach to predict and test for members of microbiota affecting specific host functions using transkingdom network analysis.*
microbes with cocktails of antibiotics. One protocol using a cocktail of 4 antibiotics for 4–5 weeks has been exceptionally popular among investigators and allowed to demonstrate the involvement of microbiota in host functions and disease pathogenesis in a variety of models. Furthermore, a few studies compared both germfree and antibiotic-treated animals and detected concordant results. However, concordant results are not always observed between these 2 models. For example, in an animal model of common variable immunodeficiency (CVID) associated enteropathy (in B lymphocyte deficient mice), we found that derivation of B cell knockout (BcKO) and control mice as germfree abolished differences in the host phenotype between the 2 genotypes observed in conventional mice (Fig. 2A). This result demonstrated the essential role of microbiota in BcKO phenotype. However, the antibiotics cocktail protocol that has successfully mimicked germfree mice in other studies did not revert host alterations in the intestine of B cell deficient mice (Fig. 2B).

Intrigued by this difference between germfree and antibiotic-treated mice, we sought to comprehensively evaluate the effects of antibiotics on the intestine. Surprisingly, we found that only about one third of gene expression changes in the gut of antibiotic-treated mice could be attributed to the depletion of normal microbiota verified by comparison to germfree mice. This effect was primarily manifested by decreased expression of genes related to many aspects of innate and adaptive immunity. Interestingly, antibiotics had much more pronounced effect on T cell numbers than on IgA-producing cells. This may be one of the reasons for the discrepancy between germfree and antibiotic-treated BcKO mice (Fig. 2) as their phenotype was IgA-dependent. An additional, non-mutually exclusive, explanation is that microbiota members insensitive to antibiotics (such as viruses) are responsible for alterations in intestines of B cell deficient mice. Indeed, this hypothesis is supported by other studies demonstrating outgrowth of diverse viruses in immunodeficient animals.

Thus, since each type of microbiota perturbation presents some drawbacks, the selection of appropriate models and interpretation of results must be performed with consideration to the potential of model-specific effects. The results of our study that evaluated global intestinal transcriptome after treatment with antibiotics cocktail can aid in these decisions by providing a thorough characterization of changes in host

Figure 2. Different effects of the absence of microbiota (A) and of antibiotics (B) on the B cell knockout (BcKO) gene expression phenotype in the intestine. Each dot on graphs represents ratios of gene expression between BcKO and control mice from BcKO signature (Shulzenko & Morgun et al., 2011). Similarity between gene expression alterations induced by B cell deficiency in conventional and germ-free (A) or conventional and antibiotic-treated (B) mice is estimated using correlation analysis and represented on the graphs.
gene expression that can be specifically attributed to 3 factors, namely, the lack of microbiota, the effect of antibiotics-resistant microbes, and direct effects of antibiotics.1

Microbiota characterization

Broad perturbations of microbial communities discussed above can provide evidence of causal roles of microbiota in host phenotypes. However, these experiments have to be followed by several questions to pinpoint the exact mechanisms: What microbes live in the particular host? Which of them are responsible for control of specific host phenotypes? What molecular mechanisms do microbiota use to influence the host? Advances in sequencing technologies over the past decade have greatly enhanced our ability to answer some of these questions. Herein, we briefly described culture-free methods, leaving out cultivation approaches reviewed elsewhere (Fig. 1).36,37

One approach to identify the taxonomic profile of a microbial community is with 16S rRNA amplicon sequencing, in which highly variable regions of 16S ribosomal rRNA from all microbes in a sample are amplified by PCR and subject to high throughput sequencing38 or hybridization to probes on a special microarray chip called Phylochip.1,39 In the case of sequencing, relative abundances of taxa are calculated from copy number of amplicons from the corresponding taxa. Abundance profiles for different taxonomic ranks are commonly generated (i.e., strain, species, genus, family etc.) and then can be used for further analyses.40 Taxonomic composition 16S amplicon sequencing can be also used to predict functional variables such as KEGG gene/pathway.41 In this case, gene composition of detected taxa is predicted from phylogenetic relationship of taxa with known annotated reference genome.

An alternative and/or complementary approach to identification of taxonomic composition by 16S is shotgun metagenomics sequencing, in which the whole genomes of all microbes (microbiome) in the sample are sequenced. With this method, gene/function abundance can be directly derived from shotgun metagenomics sequencing by assigning reads to protein sequence or protein families that have functional annotation in KEGG, SEED and COG database.42,43 This method can be very informative, as it could be not a single species that affects host phenotypes, but a bacterial function present across multiple species. In this case, the same function carried out by similar genes from different species is investigated as a single variable, providing potential molecular mechanisms regardless of taxonomy information. In addition, genome sequence reads can be compared to reference genomes in order to be assigned to specific taxa. Then the number of reads assigned to each taxon can be normalized to adjust for genome size and sequencing depth and generate abundance profiles for microbes.44

Systems approaches to infer microbial contributors to host physiology

Thanks to new sequencing technologies and new culturing techniques, we have made big advances in characterization of microbiota. Currently, the key challenge in the field is a transition from merely observational or descriptive studies to inference and testing of causal relationships between specific microbes and host biology. In other words, identification which specific taxa and/or bacterial functions are actually responsible for control of specific host function is now an area of active investigation. One recently proposed approach to this question is to systematically test libraries of randomly selected microbes through large scale colonization of germ free mice employed by Faith et al.45 Although the approach uses an analytical method that infers microbes specifically affecting host phenotype, this strategy is still highly labor and resource intensive, thus limiting its utilization to centers that have large gnotobiotic facilities.

Therefore, a more desirable approach would be to first computationally predict the most likely important players or causal factors (microbes and/or microbial genes) among hundreds of measured variables and then to experimentally test the most promising candidates. This type of question is not unique to the field of holobiont biology. For instance, cancer researchers have been facing a similar problem attempting identify driver mutations responsible for progression of tumors. Application of novel systems approaches to genomic and transcriptomic data from tumors has recently offered an efficient solution to infer a short list of candidate drivers. Indeed, reconstruction and analysis of gene regulatory networks has proven to be an excellent tool for causal inference, allowing
resistant to many antibiotics\(^1\) and therefore antibi-

- minor players in the healthy mouse microbiota, they
died in epithelium. Although these microbes are
candidates to drive mitochondrial depression and cell
death in epithelium. Indeed, we have faced the common problem that there is evidence of a microbiota effect on a host phenotype, but it was unclear which microbe was responsible. Building on our previous experience with gene networks,\(^{25,48,50}\) we developed a new approach that models host-micro-

- biota interaction that we called *transkingdom net-
w*

- work. Microbial genes (circle nodes) and host genes (triangle

- nodes). Bi-partite betweenness centrality is calculated for each
microbial gene based on the number of times it is present in the
shortest paths connecting microbial genes and human genes
(Dong et al., 2015). Microbial genes with high bipartite between-
ness centrality (red) are more likely to be key regulators of host
gene expression than genes with small values of bipartite
betweenness centrality (blue).

**Figure 3.** Bipartite betweenness centrality in transkingdom net-
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microbes predicted could out-compete *C. difficile* in mouse colonization experiments.

**Validating predictions: microbiota transplants and in vitro testing**

Similar to other fields of biology, the results of computational inferences of host microbiota interactions are much more valuable if confirmed by ‘wet-lab’ experiments (Fig. 1). Given our primary focus here in determining how microbes affect host biology, approaches taken in validation are reminiscent of widely used infection models using commensal microbes instead of pathogens. However, whereas in the infection models such microbes are normally absent in healthy hosts, commensal microorganisms can be present in normal microbiota. Therefore, monocolonization of germfree mice with a microbe of interest is seemingly the most straightforward solution to circumvent this problem and is still widely employed in experimental studies. Though it might look to be the “cleanest” way, concerns arise about the environment into which the selected microbes are introduced. For example, germfree mice have an undeveloped immune system and altered metabolism that could influence behavior of the tested species. Additionally, a given bacterium in monoassociation might not behave in the same way as it would in a community and might not induce same immune responses. For example, bacteria comprising a standard mix (Altered Schaedler Flora) were insufficient to induce colonic T regulatory cells in germfree mice when used individually. In another study, a common mouse pathogen *Citrobacter rodentium* did not trigger colitis when it was administered alone in contrast to when it was given in the presence of another bacterium. Another recently proposed approach would be to use “standardised” microbial communities which can be then supplemented with the microbe of interest. In addition, aforementioned antibiotic-treated mice can serve as another type of host. There are recent examples in the literature utilizing one of these or a combination of approaches to validate their predictions. For example, Iida and Dzutsev et al. predicted *Alistipes shahii* as a bacterium affecting the host’s ability to respond to chemotherapy. To test this, antibiotic-treated mice that otherwise have poor response were given *A. shahii*, which improved response by inducing expression of inflammatory cytokines. In another study, the genus *Sutterella* was identified by LEfSe to be associated with low fecal IgA levels. This prediction was tested not by monocolonization, but by administration of an enriched culture of *Sutterella* to mice, which converted high fecal IgA mice to a low IgA phenotype.

Finally, besides validation of effects of microbes predicted from animal models, there is a growing need for validating predictions generated from analyses of human microbiota associated with disease states. Therefore, humanized gnotobiotic mice (i.e. germfree mice colonized with human microbiota) have become a popular experimental system. Despite some concerns that this system does not fully recapitulate the effects human microbiota on immune system, some effects of microbiota on metabolism could be confirmed in humanized mice.

Classical in vitro approaches can also be used for evaluation of effects of microbes on host cells (Fig. 1). This strategy has been very useful for investigation of effects of pathogens and probiotics. Furthermore, if candidates are not just particular bacteria but specific bacterial genes, genetically modified microbes have to be tested alongside with wild-type bacteria, as was done in our study on the effects of antibiotics and antibiotic-insensitive bacteria on intestinal phenotypes. In this study, besides identifying *P. aeruginosa* as a candidate microbe as a regulator of a specific host phenotype (mitochondrial depression and cell death), we also identified LasR as a likely bacterial gene regulator of this phenotype. We anticipated that, as LasR is a primary regulator of quorum sensing, soluble factors secreted by bacteria should play in the effect on host. To validate these predictions we treated an intestinal cell line with wild-type and knockout *P. aeruginosa* conditioned growth medium. Indeed, our analytical predictions were confirmed as medium from wild type bacterium led to mitochondrial repression and cell death, while LasR-deficient bacteria were unable to produce this effect.

In conclusion, the field of host-microbiota interactions is rapidly transforming into a new discipline bringing biology and medicine into a new era. While hologenome theory makes peace between Darwin’s evolution and Lamarck’s theory of Inheritance of acquired characteristics, new systems biology approaches armed with metagenomics and gnotobiotic techniques revolutionize our understanding of health and disease. Transkingdom networks put in practice the ‘hologenome theory of evolution’ by
offering a robust framework for interrogation of hosts and their microbes as a whole. The insights from these networks provide a unique understanding of cellular and molecular mechanisms that govern social affairs between macro- and micro-species that altogether make up a holobiont.

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No potential conflicts of interest were disclosed.

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