The Integrative Human Microbiome Project: Dynamic Analysis of Microbiome-Host Omics Profiles during Periods of Human Health and Disease

The Integrative HMP (iHMP) Research Network Consortium*

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Much has been learned about the diversity and distribution of human-associated microbial communities, but we still know little about the biology of the microbiome, how it interacts with the host, and how the host responds to its resident microbiota. The Integrative Human Microbiome Project (iHMP, http://hmp2.org), the second phase of the NIH Human Microbiome Project, will study these interactions by analyzing microbiome and host activities in longitudinal studies of disease-specific cohorts and by creating integrated data sets of microbiome and host functional properties. These data sets will serve as experimental test beds to evaluate new models, methods, and analyses on the interactions of host and microbiome. Here we describe the three models of microbiome-associated human conditions, on the dynamics of preterm birth, inflammatory bowel disease, and type 2 diabetes, and their underlying hypotheses, as well as the multi-omic data types to be collected, integrated, and distributed through public repositories as a community resource.

Introduction

The human microbiome is important for human health, behavior, and disease, yet its function and dynamics during healthy and disease states are only partially understood. Studies of the microbiome to date have indicated impacts on personalized medicine ranging from inactivation of pharmaceuticals in some individuals (Haiser et al., 2013) to increased risk of cardiovascular disease due to microbial metabolic byproducts (Koeth et al., 2013). These findings show particular promise given that they have already come to light early in the field’s transition from high-throughput discovery to detailed mechanistic studies, a process still ongoing after more than a decade of human omics. The first phase of the NIH Human Microbiome Project (HMP, fiscal years 2008–2012, http://www.commonfund.nih.gov/hmp) examined the diversity and composition of the human microbiome to evaluate (1) common patterns of microbial diversity associated with health and (2) whether taxonomic or functional features of the microbiome correlated with diseases by analyzing a large healthy cohort and a set of demonstration projects. These efforts revealed the vast microbial diversity associated with humans and provided new insights into the ecology of the host-microbiome superorganism (Human Microbiome Project Consortium, 2012a, 2012b).

Results from previous human microbiome studies (Dethlefsen et al., 2006; Hooper and Gordon, 2001; Qin et al., 2010; Savage, 1977) and from the first phase of the HMP (Human Microbiome Project Consortium, 2012a, 2012b) suggested that taxonomic composition of the microbiomes between subjects can differ significantly. This indicated that taxonomic characterization alone may not be sufficient to reveal relationships between the microbiome and specific health or disease states. Interestingly, metabolic pathway reconstructions of the metagenomic data from the HMP healthy cohort study suggested that the microbiomes of healthy subjects may share similarities in their metabolic pathways; this feature may be a key common property of the microbiome within any given cohort (Human Microbiome Project Consortium, 2012b). Further, in some of the HMP Demonstration Projects (http://commonfund.nih.gov/hmp/fundedresearch), analysis of microbiome biological properties, such as community transcripts, proteins, or metabolites, hinted at losses or gains of key microbiome functions associated with particular diseases.

The second phase of the HMP (or iHMP, Integrative Human Microbiome Project Consortium, fiscal years 2013–2016, http://hmp2.org) will further examine the role of the microbiome in human health and disease through a study of three models of microbiome-related human conditions. These three longitudinal studies focus on (1) pregnancy, including those resulting in preterm birth; (2) gut disease onset, using inflammatory bowel disease (IBD) as a model; and (3) respiratory viral infection and onset of type 2 diabetes (T2D). In each case, subjects will be followed for up to 3 years. Multiple properties of the microbiome, such as phylogenetic composition and associated functional omic data (e.g., transcriptome, proteome, metabolites, etc.) will be analyzed. Importantly, host functional changes will also be monitored as well as the metabolites originating from both the host and the microbiome so that microbiome and host changes can be simultaneously followed to provide a more comprehensive picture of the dynamic changes that occur during these periods. The studies were selected as exemplars of microbiome-associated human conditions and therefore should be of broad interest to the research community. These resulting data sets will provide opportunities for new hypotheses to be tested and for evaluating the utility of simultaneously assessing the microbiome and the host biological properties for understanding host-microbiome interactions.

Data Deposition, Access, and Sharing

Primary data (e.g., sequence reads or mass spectrometry profiles) and derived properties (e.g., microbial composition, gene expression and protein levels, and metabolite profiles) will be generated by each project. Table 1 lists the biospecimens and
The Multi-Omic Microbiome Study: Pregnancy Initiative (http://vmc.vcu.edu/momspi) is a study designed to understand the role of the microbiome in the health of the pregnant woman and her fetus. The goal of this study is to promote women’s health during pregnancy and to explore the hypotheses that the microbiota contributes to preterm birth and stillbirth. To explore these hypotheses, women will be enrolled early in pregnancy and followed longitudinally at multiple prenatal visits, at admission to and discharge from labor and delivery, and at postpartum visits. Enrollment will be skewed toward populations at higher risk for preterm birth. Microbiome compositions will be measured by 16S rRNA gene surveys, whole metagenome shotgun sequencing, and bacteriological cloning and sequencing. Microbial and host gene expression profiles will be examined by whole metatranscriptome sequencing, and cytokine and lipid profiles will be explored. Protein-protein interactomic maps of the microbiomes and the host will also be generated (Table 1).

Specifically, this project will:

1. Enroll up to 2,000 pregnant women and their neonates from clinics associated with the VCU Health Center and the Global Alliance to Prevent Prematurity and Stillbirth (GAPPS). Women will be sampled at multiple body sites (vagina, rectum, skin, nares, mouth, blood) longitudinally throughout pregnancy, at labor and delivery, and at follow-up visits. Neonates will be sampled at multiple body sites (rectum, mouth, skin, nares, meconium, stool) at birth and discharge visits. Placenta, cord blood, and choioamniotic membranes will be collected, and amniotic fluid samples will be collected during cesarean deliveries.

2. Perform multi-omics analyses on maternal and neonatal samples. All samples will be subjected to 16S rRNA gene surveys and all vaginal and serum samples will be subjected to lipidomic analysis. Selected samples will be processed for whole metagenomic and whole metatranscriptomic sequence analysis, host transcriptome profiling, cytokine profiling, bacteriological culture, and protein–protein interaction analyses.

3. Integrate the data from these multi-omic studies into existing and local databases available to vaginal microbiome investigators. The results will reveal associations between the microbiome composition, microbial metabolic potential, gene expression profiles of the host and microbiota, and the host cytokine and metabolome/lipidome expression profiles. Dynamic changes in the microbiomes and in gene expression profiles will be monitored longitudinally during and after pregnancy. Associations with preterm birth, stillbirth, and other adverse outcomes of pregnancy will be identified.

Overall, this research will reveal the impact of the microbiome on the gene expression, cytokine, and metabolome/lipidome profiles of pregnant women and their relationship to preterm birth or other adverse outcomes of pregnancy. The long-term objective of this study is to promote women’s health during pregnancy and prevent adverse outcomes.

**Cohort Description**

This study will longitudinally sample about 2,000 women, at least 15 years old, during and after their pregnancies. The neonates of these women will also be sampled. The predicted racial/ethnic composition of our cohort, based on the demographics of visitors to the VCU and GAPPS collection sites, is highly diverse. African Americans, who are at higher risk for preterm birth, represent about one-third of our anticipated cohort. Of the remainder,
| Source of Property | Property Derived from Primary Data | Primary Data from Biospecimen | Biospecimen from Preterm Birth Cohort | Biospecimen from IBD Cohort | Biospecimen from Prediabetic Cohort | Repository for Primary Data |
|-------------------|-----------------------------------|-------------------------------|--------------------------------------|----------------------------|-----------------------------------|----------------------------|
| Microbiome        | microbial community composition   | 16S rRNA gene survey          | cervical⁸, vaginal⁸, rectal, buccal, fetal membranes, placenta, amniotic fluid from women; buccal, rectal, stool, meconium, respiratory secretions from neonate | stool                      | anterior nares, stool              | SRA                        |
|                   |                                   |                               |                                       |                            |                                   |                            |
|                   | microbial community composition   | whole metagenome shotgun sequences | vaginal⁸                          | stool                      | anterior nares, stool              | SRA                        |
|                   | predictions of microbial genes, metabolic pathways | whole metagenome shotgun sequences | vaginal⁸                          | stool                      | anterior nares, stool              | SRA                        |
| RNA transcript profiles | whole metatranscriptome shotgun sequences | vaginal⁸                        | stool                      | anterior nares, stool              | SRA                        |
| microbiome metaproteome profiles | LC-MS/MS peptide profiles | –                             | stool                      | –                           | EBI PRIDE and/or Peptide Atlas |
| viral community composition | whole virome shotgun sequences | –                             | stool                      | –                           | SRA                        |
| bacterial cultures | bacterial isolates               | cervical⁸, vaginal⁸, rectal, buccal, stool from mothers or neonates | –                           | –                           | ATCC/BEI                     |
| bacterial whole genome sequences | bacterial isolates               | cervical⁸, vaginal⁸, rectal, buccal, stool from mothers or neonates | –                           | –                           | SRA                        |
| bacterial whole genome sequences | bacterial single-cell sequences | –                             | stool                      | –                           | SRA                        |
| single-cell bacterial RNA transcript profiles | single-cell bacterial transcript sequences | –                             | stool                      | –                           | SRA                        |
| Host              | subject exome/whole genome       | subject genome sequences      | blood (future⁹) from mothers and neonates | blood                     | blood                           | dbGaP/SRA                   |
| RNA transcript profiles | whole transcriptome sequences | vaginal⁸                        | colon biopsy | PBMCs                     | dbGaP/SRA and GEO             |
| subject protein profiles | LC-MS/MS peptide profiles | –                             | stool                      | PBMCs, serum (future)       | EBI PRIDE and/or Peptide Atlas |
| systemic inflammation levels | cytokine profiles               | vaginal⁸, buccal from mothers or neonates | blood                     | plasma                      | Study DB                     |
| intestinal epithelial cell cultures | intestinal epithelial cell isolates | –                             | colon biopsy | –                         | –                             |
| subject DNA methylation profiles | reduced representation bisulfite sequencing (RRBS) profiles | –                             | blood                     | PBMC (future) | SRA                           |

(Continued on next page)
Table 1. Continued

| Source of Property | Property Derived from Primary Data | Primary Data from Biospecimen | Biospecimen from Preterm Birth Cohort | Biospecimen from IBD Cohort | Biospecimen from Prediabetic Cohort | Repository for Primary Data |
|--------------------|-----------------------------------|-------------------------------|----------------------------------------|----------------------------|------------------------------------|-----------------------------|
| intestinal inflammation levels | fecal calprotectin protein concentrations | – | stool | – | | Study DB |
| serum antibody composition and levels | serology | – | blood | – | | Study DB |
| subject contaminating genome sequences | human sequence from unfiltered total microbial community sequences | vaginal<sup>a</sup> | stool | stool | | dbGaP/SRA |
| subject profiles for cohort | subject phenotypes, clinical metadata, medical panels | collected on each subject in the study | collected on each subject in the study | collected on each subject in the study | | dbGaP |
| Global (host and microbiome) | protein-protein interaction network between host and microbiome | vaginal<sup>a</sup> | – | – | | EBI IntAct |
| pathway-level crosstalk between host and gut microbiome | intestinal epithelial cell profiling response to bacterial metabolites | – | colon biopsy | – | | dbGaP/SRA and GEO |
| global metabolite profiles | untargeted and targeted LC-MS metabolomic profiles | – | stool | urine, plasma | | Metabolomics Workbench |
| global lipid profiles | untargeted and targeted LC-MS metabolomic profiles | vaginal<sup>a</sup> | stool | urine, serum | | Metabolomics Workbench |

<sup>a</sup>From mothers only

<sup>b</sup>Samples collected for possible future analysis
Table 2. Glossary of Terms Used to Describe Primary Data in the Paper, in the Figures, and in Table 1.

| Primary Data Type                      | Definition                                                                                                                                 |
|----------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------|
| 16S rRNA gene survey                   | Sequence-based analysis of 16S ribosomal RNA gene in total DNA extracts; the data are used to develop microbial community compositional profiles. |
| Whole metagenome shotgun sequences     | Sequence-based analysis of all genes in total DNA extracts; the data are used to develop microbial community compositional, functional, and genomic profiles. |
| Whole metatranscriptome shotgun sequences | Sequence-based analysis of RNA transcripts in a microbial community by converting RNA in total RNA extracts to complementary DNA and sequencing the cDNA. |
| Whole virome shotgun sequences         | Sequence-based analysis of all virus genes and genomes used to develop viral community profiles. These sequence data are derived from two approaches, either by first isolating viral-sized particles from a sample and then sequencing this fraction, or by sequencing the DNA (or cDNA from RNA) extract and computationally determining DNA and RNA viral sequences in the metagenome. |
| Bacterial isolates                     | Isolation and cultivation of a specific bacterium in a mixed community through the use of selective media or enrichment techniques to preferentially grow one bacterium of interest. |
| Single-cell genome sequences           | Physical separation or physical enrichment of a single microbial cell from a mixed population of cells. Sequence-based analysis of the single cell's genome is conducted using a DNA random priming method to first increase the total DNA concentration in the cell and subsequently sequence this amplified DNA. |
| Single-cell RNA sequences              | Sequence-based analysis of RNA transcripts in a cell by converting RNA in total RNA extracts to complementary DNA and sequencing the cDNA. |
| LC-MS/MS peptide profiles              | Analysis of individual peptides in a mixture of proteins by a combination of liquid chromatography and mass spectrometry after enzymatic digestion of the protein mixture. To derive the microbiome metaproteome, data from the companion metagenome is used along with KEGG and other protein databases to verify the microbial proteins in the total protein mixture. To derive the host proteins, a similar approach is used based on search against the human genome. |
| Human subject whole genome sequences   | Analysis of the sequence of the host genome using high-throughput DNA sequencing. |
| Whole exome sequences                  | Analysis of the protein coding regions of the host genome, which is generally done through sequence analysis. |
| Whole transcriptome sequences          | Sequence-based analysis of RNA transcripts in host tissue using polyA tail separation of eukaryotic transcripts from a mixture of transcripts, converting the RNA to complementary DNA and sequencing the cDNA. |
| Cytokines                              | Circulating immune system glycoproteins in blood, plasma, or other bodily fluids are measured through an ELISA assay; data are used as a marker for systemic inflammation. |
| DNA methylation profiles               | Measurement of methylated regions of the host genome. This method uses a combination of restriction enzymes and a reduced representation bisulfite sequencing (RRBS) method to enrich for regions of the genome with cytosine-guanine pairs; these regions are then screened for methylated bases. |
| Fecal calprotectin proteins            | Fecal calprotectin is a protein found in stool, the concentration of which is measured through a standardized immunoassay method and is used to evaluate levels of intestinal inflammation. |
| Serology                               | Measurement of antibodies for specific pathogens or protein markers conducted with serum through an ELISA assay. |
| Human subject contaminating genome sequences | Sequence from total nucleotide extracts from most microbially focused sample types yields a combination of host and microbial sequence data. A computational filtering step, using human genome reference sequence, will separate microbiome from human sequences. |
| Interactomes                           | Analysis of protein-protein interactions between members of the microbiota or between the microbiota and the host. |
| Intestinal epithelial cell profiles    | Functional or sequence-based readouts of phenotypes for cell lines derived from primary epithelial cells from individual hosts. |
| Metabolomes                            | Measure of metabolomic profiles using untargeted and targeted LC-MS methods. |
| Lipidomes                              | Measure of lipid profiles using 2D UPLC-ESI-MS/MS (ultra performance liquid chromatography combined with electrospray ionization tandem mass spectrometry) with an initial focus on AA (arachidonic acid) and eicosanoids. |

About 37% will be Hispanic, ~20% Caucasian, ~5% Asian, and ~4% Native American. Because high-risk populations attending high-risk clinics will be sampled, it is anticipated that approximately 15%–20% of the study participants will deliver preterm.

**Sampling**

Samples will be collected from pregnant women at 3–5 antepartum visits, at delivery, at discharge, and at subsequent follow-up visits. Neonate samples will be collected at delivery and at discharge. Placenta, cord blood, and amniotic fluid (C-section only) will be collected at delivery. Participants will complete comprehensive electronic surveys that detail health history, background, habits, and diet. These data will be incorporated into a secure database at VCU and a subset will be uploaded to dbGaP. Vaginal, cervical, rectal, buccal, skin, and nares samples will be collected with soft swabs. Cervical samples will be collected during a speculum exam. Samples from the vagina and rectum and the vaginal pH will be taken at each visit, except discharge, and again at follow-up visits, either by a clinician or by
self-sampling. Buccal samples will be collected at each visit, and skin (chest and antecubital fossa) and nares samples at each visit except triage and delivery. Additional skin samples (chest and dominant palm) will be collected at discharge. Blood will be collected from participants early in pregnancy and again at triage. Amniotic fluid will be collected with a sterile syringe, and placenta and membrane samples will be collected with sterile scissors. Cord blood will also be collected.

Buccal, rectal, meconium, stool, skin, and nares samples from term and preterm neonates will be collected with soft swabs at delivery and discharge. Tracheal aspirates will be collected from neonates who are intubated. Over 100,000 samples will be deposited into the RAMS Registry/Repository at VCU (http://www.ramsregistry.vcu.edu) for future research.

**Multi-Omic Assays**

Samples will be subjected to a panel of technologies, including 16S rRNA gene surveys, whole metagenome and metatranscriptome shotgun sequencing, bacterial genome sequencing, cytokine and lipid profiling, and generation of protein-protein interaction networks of the microbiome and the host (Figure 1).

**16S rRNA Gene Surveys for Microbiome Profiles: All Sample Types.** For 16S rRNA gene survey analysis, DNA will be sequenced in barcoded pools using 2 × 250 bp PE technology on an Illumina MiSeq sequencer in a standard pipeline that generates more than 30,000 reads per sample. Reads will be processed using the RDP Classifier (Wang et al., 2007) and assigned to species using STIRRUPS (Fettweis et al., 2012) or by OTU-based methods to generate microbiome profiles.

**Whole Metagenome Shotgun Sequencing and Whole Metatranscriptome Sequencing: Vaginal Samples.** To clarify the contribution of the metabolic potential and gene expression profiles of the vaginal microbiome to preterm birth, case-matched samples from 60 women who experience preterm birth and 60 women who experience normal term birth will be subjected to in-depth whole metagenome shotgun sequencing and whole metatranscriptome sequencing. Reads will be taxonomically classified and quantified, and metabolic reconstruction and analysis will be performed using the HUMAnN pipeline (Abubucker et al., 2012), ASGARD (Alves and Buck, 2007), or similar tools.

**Cytokine Profiles: Selected Vaginal and Buccal, Mother and Infant.** The immune response will likely undergo dynamic changes during pregnancy, and these may be particularly evident in cases associated with adverse events. To monitor these changes, we will analyze the profiles of a panel of cytokines, including some
that were previously linked to preterm birth and/or inflammation (IL-1α, IL-1β, IL-1ra, IL-2, IL-4, IL-6, IL-8, IL-10, IL-12 [p40], IL-12 [p70], IL-17, sCD40L, TNF-α, TNF-β, IFN-γ) in vaginal (mother) and buccal (mother and infant) samples. Representative vaginal and buccal samples from the ~2,000 mothers and infants will be analyzed. Analyses will be performed in a high-throughput format using a multiplex magnetic bead-based human cytokine ELISA assay.

Lipidomics: All Vaginal Samples. Bioactive lipids play an important role in the maintenance of pregnancy and the induction of labor. These lipids are likely associated with the local microbiome and pregnancy outcomes. Thus, we plan to analyze levels of more than 150 lipids (e.g., eicosanoids, sphingolipids, and steroid hormones) in vaginal samples collected longitudinally throughout pregnancy using methods previously described (Wijesinghe et al., 2010, 2011). Assays will employ liquid chromatography-tandem mass spectrometry (LC-MS/MS) with a focus on the metabolites that play a role in pregnancy and parturition.

Bacterial Genome Sequencing: Selected Samples from All Sites. Many bacteria identified in 16S rRNA surveys of vaginal samples, some of which are associated with women’s urogenital health, have never been cultured or characterized. Several isolates associated with preterm birth (Harwich et al., 2012) have been cultured and characterized, and their pathogenic potential is now being studied. Select isolates cultured from samples with previously unidentified taxa associated with preterm risk will be assayed for pathogenic potential and subjected to whole genome sequence analysis. The assemblies will be deposited into GenBank and the cultures at ATCC/BEI.

Interactomics: Protein-Protein Interaction Maps: Selected Vaginal Samples. Microbes colonizing the vagina likely impact vaginal health by secreting proteins that bind to or interact with proteins of cocolonizers or the host. A pilot study will be performed that will apply a novel yeast two-hybrid strategy that invokes high-throughput in silico identification of interacting partners using targeted next-generation sequencing of an amplicom including both bait and prey coding sequences from a specific hybrid construct following selection. If the pilot is successful, the technology can be applied to interactions among complex bacterial communities or between bacteria and the host.

iHMP Study: Characterizing the Gut Microbial Ecosystem for Diagnosis and Therapy in IBD

Over the past decade, IBDs have emerged as one of the most important human conditions linked to the gut microbiota (Abraham and Cho, 2009; Danese and Fiocchi, 2011; Khor et al., 2011; Xavier and Podolsky, 2007). IBD comprises both Crohn’s disease (CD) and ulcerative colitis (UC), which together affect nearly 1.5 million Americans (Loftus, 2004). Studies of environmental (Baumgart and Sandborn, 2012; Molodecky et al., 2011; Ordás et al., 2012; Renz et al., 2011) and microbial (Packey and Sartor, 2009; Sokol et al., 2008) associations with IBD have resulted in neither simple diagnostic markers nor targetable points of intervention. Instead, IBD has been repeatedly linked to the overall ecology of the human gut microbial ecosystem, including community diversity (Frank et al., 2007; Manichanh et al., 2006; Ott et al., 2004), a range of microbial over- and underabundances (Joossens et al., 2011; Scanlan et al., 2006; Sokol et al., 2009; Willing et al., 2009), and the depletion and enrichment of microbial metabolic activities (Erickson et al., 2012; Meyer et al., 2012; Morgan et al., 2012). To date, however, no one aspect of the gut microbiome has yielded data that identify causal biomolecular mechanisms in IBD or provide a translationally actionable target for IBD.

Project Description

An IBD multi-omic database, the IBDMDB (http://ibdmdb.org), comprising an integrated data resource, will be developed to enable the gut microbial ecosystem as a target for diagnosis, therapy, and mechanistic understanding of IBD. This study will leverage well-phenotyped patient cohorts in both pediatric and adult populations to provide longitudinal profiling of the biological properties of the human gut microbiome in IBD. To further characterize the microbiota’s mechanisms of host interaction, profiles of host genetic and functional properties from a subset of longitudinal time points will also be collected. In addition to data being made rapidly accessible to the community by building on this research group’s current computational infrastructure for multi-omic data processing and distribution, this study will validate and distribute its sample collection and bioinformatic protocols, including platforms for single-cell- and host-cell-focused multi-omic assays.

The biological questions of interest to be addressed by the multi-omic data generated by this project include (1) identifying the molecular mechanisms by which the intestinal microbiome may trigger disease activity in patients with IBD, (2) determining if microbial composition predicts subsequent risk of flares in disease activity, and (3) testing whether successful response to therapy can be predicted from the stool microbiota. These hypotheses are testable by the dense longitudinal multi-omic surveys of the IBD patients and controls in this study design. Specifically, this project will:

1. Create an integrated data repository, the IBDMDB, for standardized community access to multi-omic data, building on existing resources and supporting rapid future algorithm development. This will begin with a pilot data set of existing multi-omic data from two cohorts, including cross-sectional and longitudinal 16S, metagenomic, metabolomic, and proteomic profiles.

2. Implement and validate standardized stool and biopsy sample collection methods among three established IBD patient cohorts.

3. Recruit well-characterized subjects within these three cohorts targeting, respectively, new-onset pediatric subjects, new-onset adult subjects, and established Crohn’s disease and ulcerative colitis, in addition to non-IBD controls.

4. Longitudinally collect stool samples and, for a subset of collection points, rectal and colon biopsies from which microbial 16S surveys, metagenomic and metatranscriptomic sequences, viromes, metabolites, and proteins will be surveyed. A subset of stool and biopsy samples will additionally provide 16S and metagenomic and metatranscriptomic population analyses through single-cell techniques (Table 1).

5. Acquire host information, including rich clinical and phenotypic metadata for all collection points and, for a subset, host genetics, gene expression data, DNA methylation profiles, serology, and primary intestinal epithelial cells for profiling response to microbial products.
**Cohort Description**

The study will collect stool samples, biopsies, and blood samples from three clinical cohorts. The primary sample will be serial stools every 2 weeks for 1 year, which will be analyzed using 16S rRNA gene surveys, whole metagenome and metatranscriptome shotgun sequencing, metabolomics, whole virome shotgun sequencing, proteomics, and a fecal calprotectin assay. Intestinal biopsies will also be collected at a standardized rectal location for all subjects, plus at the site of active inflammation for IBD patients, at time zero and as clinically indicated over the following year (approximately one additional time point per subject). 16S surveys will be carried out on these biopsies, in addition to host whole transcriptome sequencing and host cell profiling by culture. A subset of biopsies and stools will be additionally assessed by single-cell 16S and whole metagenome and metatranscriptome shotgun sequencing. Finally, blood samples will be drawn at recruitment and 6 month intervals for host genetics, serology, and epigenetics by reduced representation bisulfite sequencing (RRBS).

This study includes three IBD cohorts providing these samples and metadata, coupled with two additional cohorts from which early “proof-of-principle” data are already available. The samples described above will be collected from (1) PRISM (the Prospective Registry in IBD Study at MGH), recruiting from Massachusetts General Hospital (Ananthakrishnan et al., 2014; Morgan et al., 2012); (2) the MLI (Mucosal Luminal Interface) cohort recruiting from Cedars-Sinai Medical Center (McHardy et al., 2013; Tong et al., 2013); and (3) a pediatric IBD cohort recruiting from Emory University and Cincinnati Children’s Hospital (Gevers et al., 2014; Walters et al., 2014). Each cohort will provide 30 subjects (12 CD, 12 UC, and 6 non-IBD controls), with MGH focusing on new-onset adult patients, CSMC on adult patients with established disease, and Emory/Cincinnati on new-onset pediatric patients. Together, these cohorts have previously enrolled a total of over 3,600 subjects and collected more than 1,500 samples. A smaller, existing multi-omic data set is included in this study as a proof-of-principle, derived from two Swedish cohorts, the Swedish Twin and Swedish Longitudinal studies of IBD (Dicksved et al., 2008; Erickson et al., 2012; Willing et al., 2010). These include up to 40 twin pairs and 119 individuals providing up to ten time points, respectively, and represent one of the only existing multi-omic data sets in IBD around which the computational infrastructure will be organized and prototyped.

**Sampling**

This study has already established a convenient method for self-collected fecal samples compatible with all of the multi-omic assays, enabling longitudinal profiling in smaller collection tubes tolerant of over 24 hr at ambient temperature (Franzosa et al., 2014). Two tubes are provided per subject, one dry (for proteomics and calprotectin) and the other containing an ethanol fixative (for all other assays). Validation studies indicate that there is close agreement between freshly frozen fecal sample assays and samples in preservative. Each fecal collection tube cap includes a spoon for easier collection and can be shipped under ambient conditions, and clear instructions are provided to the subjects (publicly available at http://ibdmdb.org/protocols). Each sample is shipped by subjects in a self-addressed, prepaid FedEx box for same-day delivery, with aliquots distributed in subsequent batches for off-site assays. Samples are accompanied by short metadata questionnaires, and clinical data at first visit includes demographics, disease characterization (including medication history), short- and long-term food frequency questionnaires, and an environmental factors questionnaire.

**Multi-Omic Assays**

**Microbial Community Composition, Functional Potential and Activity: All Samples.** The first tier of multi-omic data in IBD (Figure 2) will include 16S rRNA surveys, whole metagenomic shotgun sequencing, and whole metatranscriptome shotgun sequencing from stool samples (all assays) and biopsies (16S only). Improving our understanding of complex human-associated microbial communities will require at a minimum the aspects of microbial community structure, its functional potential, and realized gene expression activity. This dictates the inclusion of 16S rRNA gene-based, whole metagenome DNA shotgun-based, and community mRNA-based sequencing (Gevers et al., 2012). This study will also survey the nonbacterial members of the gut microbiota, including archaeal, viral, and eukaryotic components. This study includes a viral enrichment and sequencing protocol for each stool sample.

**Metabolomic and Protein Profiles: Stool Samples.** This study will measure metabolomic and protein profiles of stool samples from participants in the project cohorts using a combination of mass spectrometry (MS) methods. Four LC-MS/MS approaches will measure complementary metabolite classes using sensitive, high-resolution mass spectrometers enabling nontargeted measurements of known metabolites (several hundred in total) and unidentified metabolites in the same run. For each stool sample, metabolites will be identified using reference data generated from an in-house Broad Institute library of > 500 compounds, and novel metabolites will be annotated using a database of > 8,000 metabolites and confirmed using references when possible.

Protein profiles will be measured by two-dimensional nano liquid chromatography (2D-LC) and tandem mass spectrometry (MS/MS) (Cantarel et al., 2011; Presley et al., 2012), with proteins identified by searching custom databases consisting of the human genome, matched metagenomes from the same samples, and a defined set of reference genomes, including many that are in the HMP reference genome database. Based on preliminary results, it is anticipated that > 3,000 proteins will be identified per sample and that approximately half of the proteins will be human proteins based on a pretested method for preparation of the protein extracts. The combination of human and microbial protein data will provide a window into the types of host-microbe interactions that occur in the gut and how these functions are correlated to IBD.

**Host Biomolecular Activities in the Gut: Colon Biopsy Samples and Cultured Host Cells.** To begin linking gut microbial community biology to that of the host, this study will assess changes in host biomolecular activities in the gut as part of multi-omic profiling. Specifically, host gene expression will be assayed by RNA-seq from colon biopsies (from a standardized rectal location in all subjects and from the active site of IBD in patients) and the DNA methylation state of the host genome in circulating blood by RRBS. This will allow the discovery of links between epigenetic content and disease states relating to microbial communities. Finally, to detail pathway-level crosstalk between gut
microbes and the intestinal epithelium, host cells isolated from primary biopsy samples will be cultured and perturbed in vitro (Gallo and Hooper, 2012; Maynard et al., 2012; Stappenbeck et al., 1998) to determine the relative contribution of metabolites and host epithelial cells in this crosstalk.

**iHMP Study: Microbiome and Host Changes during Respiratory and Other Stress Conditions in Individuals at Risk for T2D**

Type 2 diabetes mellitus (T2D) is a significant health problem facing our nation. Close to 20 million individuals in the United States have T2D, costing hundreds of billions of dollars annually, and the incidence is projected to double by 2050 (American Diabetes Association, 2008). Differences in the gut microbiome have been noted between diabetics and healthy individuals (Qin et al., 2012), and direct alteration of the microbiome in mice has been shown to lower glucose levels (Turnbaugh et al., 2006). The hypotheses to be tested in this study include the following: (1) environmental stress causes dynamic changes in specific biological pathways in the human body, and these changes lead to alteration of the human microbiome and the metabolome, including glucose; (2) some of the changes may affect the

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**Figure 2. Characterizing the Gut Microbial Ecosystem for Diagnosis and Therapy in Inflammatory Bowel Disease: Sample Collection, Assay, and Data Generation Workflow**

Samples from Crohn’s disease patients, ulcerative colitis patients, and non-IBD controls are collected at Massachusetts General Hospital (adult new onset), Emory University (pediatric new onset), Cincinnati Children’s Hospital (pediatric new onset), and Cedars-Sinai Medical Center (adult established). From each patient, three different types of samples are collected: longitudinal stool samples, periodic biopsies, and regularly scheduled blood samples. Biopsies are collected as clinically indicated, blood during clinical visits, and stool samples are self-collected by participants at home and shipped directly to a centralized handling and aliquoting pipeline. Multi-omic data generation (primarily, but not entirely, nucleotide sequence- and mass spectroscopy-based) provides microbial, host, and mixed profiles including 16S rRNA gene surveys, whole metagenome and metatranscriptome shotgun sequences, metabolite and protein profiles, single-cell assays, whole virome shotgun sequences, and serological profiles. Each sample is further accompanied by clinical (bloods/biopsies) or self-reported environmental and dietary (stools) metadata.
epigenome, leading to molecular and biological alterations that extend well beyond the time of the stress period; and (3) different physiological stresses, such as respiratory viral infections and diet changes, may have common effects in both the host and microbiome.

**Project Description**

To better understand the biological changes that occur during the onset and progression of T2D, this study ([http://med.stanford.edu/ipop/](http://med.stanford.edu/ipop/)) will perform a detailed analysis of the biological processes that occur in the human microbiome and host by longitudinal profiling of patients at risk for T2D during healthy and stress periods. Multiple sample types will be collected from the study participants every 2–3 months during their healthy periods, with more frequent sampling during periods of respiratory illness and other environmental stress. Multiple microbiome biological properties will be analyzed. Changes in host functional profiles will also be revealed.

This study is designed to develop a detailed understanding of the physiological changes that occur in the microbiome and host during viral infection and during changes in glucose levels and insulin resistance. This longitudinal study is expected to reveal changes in the microbiome and host at an unprecedented level and frequency and to identify the molecules and pathways that change during viral infections and diabetes onset and progression. Specifically, this project will:

1. Establish a cohort of approximately 60 consented individuals at risk for T2D. Dense sampling will occur during respiratory infection and other stress states, with less frequent sampling every 2–3 months during healthy periods.
2. Perform multi-omic analysis of longitudinal microbiome and host samples (Table 1). Microbiome genomic, transcriptome, and proteome content will be determined in fecal and nasal samples and in peripheral blood mononuclear cell (PBMC) samples for viral and endogenous microbiome populations. Host genomic, transcriptomic, and proteomic data will be concurrently analyzed in blood components (PBMCs for genome and transcriptome; plasma for proteome). Combined microbiome and host metabolites will be analyzed in serum and urine. Antiviral antibodies will also be analyzed.
3. Analyze and integrate omic information from the microbiome and host. The different omics information (genome, transcriptome, and proteome) from nasal and fecal samples will be analyzed individually and in a combined fashion to fashion dynamic pathways that change during viral infection. In addition this study will correlate microbiome omic temporal profiles with those of the host to obtain an unprecedented view of the global microbiome-host changes that occur during viral infections.

**Cohort Description**

This study will execute a 3 year study in 60 males and females at high risk for T2D. Those individuals with either impaired fasting glucose (fasting plasma glucose > 100 mg/dL) or impaired glucose tolerance (2 hr glucose > 140 mg/dL in oral glucose tolerance test) will be enrolled. In order to further increase the number of study participants with chances of conversion to diabetes, but to maintain some degree of homogeneity in study subjects, the study will include only those volunteers with body mass index 25–40 kg/m² and age 35–65 years. Given estimates of 11% conversion to frank diabetes per year ([Ratner and Diabetes Prevention Program Research, 2006](http://med.stanford.edu/ipop/)), the study anticipates up to ten cases of diabetes to occur over the 3 year study period. All participants will receive genetic counseling prior to enrollment.

**Sample and Data Collection**

Qualified and consented participants will proceed to baseline testing, which includes a blood draw for molecular omics analysis, as well as measurement of fasting lipid panel markers HbA1c and hsCRP. Subjects will return every 3 months for blood draws and microbiome samples and will complete online surveys that are automatically sent to them to document changes in medications, activity, body weight, blood pressure, diet, menstrual cycle, stresses, etc. Subjects will undergo frequent sampling during periods of environmental or medical stress. In this study, stresses are defined as medical illness, physical injury/pain, major or minor operation, major life changes (birth, death, divorce, marriage, and change of home or job). The study will reinforce different collection schemes depending on types of stress events; for example, in cases of viral infection, the study will collect samples on days 1, 2, 4, 7, 21, and 35 after the onset of infection symptoms.

For each visit, either quarterly healthy visit or during stress periods, the study will collect peripheral blood to prepare PBMCs, plasma and serum, as well as urine and microbiome samples (nasal, fecal, skin, and tongue) using standard HMP protocols.

**Multi-Omic Assays**

To better understand the relationship between healthy and stressed states in this cohort, the study will be analyzing a suite of omic properties from both the host and microbiome. These include whole metagenome shotgun and metatranscriptome sequencing, host whole genome/transcriptome sequencing, cytokine and autoantibody profiles, metabolomics profiles, and standard clinical tests (Figure 3).

**Whole Metagenome Shotgun and Metatranscriptome Sequencing: Stool and Nasal Samples.** To follow the diversity of the microbiome in prediabetics and how it changes during stress periods, both 16S and whole metagenome shotgun sequencing of nucleic acids isolated from nasal and stool samples will be performed. Previously established HMP methods for alignment of whole metagenomic shotgun sequences to reference genomes will be used ([Human Microbiome Project Consortium, 2012a, 2012b; Martin et al., 2012](http://med.stanford.edu/ipop/)) for computation of taxonomic representation and breadth and depth of coverage, for variant calling in metagenomic data ([Schloissnig et al., 2013](http://med.stanford.edu/ipop/)), for pathway analysis ([Abubucker et al., 2012](http://med.stanford.edu/ipop/)), and for virome analysis ([Wylie et al., 2012a, 2012b](http://med.stanford.edu/ipop/)).

To follow microbiome activity, microbial gene expression will be analyzed. Bacterial RNA will be isolated and enriched for mRNA using procedures that remove > 95% of 16S and 23S rRNAs ([Maurice et al., 2013; Turnbaugh et al., 2009](http://med.stanford.edu/ipop/)). The resulting RNA is converted into cDNA using random priming and sequenced using the Illumina platform. Analysis of gene expression for each gene will follow previously published computational pipelines and algorithms ([Maurice et al., 2013; Turnbaugh et al., 2009](http://med.stanford.edu/ipop/)).

**Host Whole Genome/Transcriptome Sequencing: PBMCs.** To better understand how each subject responds to dynamic
changes as well as to facilitate mapping for downstream transcriptome and proteome analyses, participants will have their genome sequenced (from PBMCs) using Illumina technology. Single nucleotide variants, short insertions and deletions (<100 bp Indels), and structural variants will be scored using the HugeSeq pipeline (Lam et al., 2012). For each PBMC sample, the transcriptome will be analyzed using RNA-seq, involving the random primed synthesis of cDNA from RNA depleted of rRNA. The reads will be mapped, and gene expression, isoform expression, allele expression, and RNA editing will be mapped using a previously published pipeline (Chen et al., 2012).

| Subjects | Sample Collection | Assays |
|----------|------------------|--------|
| Pre-diabetes | Stool samples | Microbes: Metagenome sequencing |
| Diabetes at risk | Nasal Swabs | Microbes: Metatranscriptome seq |
| RSV and/or other stress | Urine samples | Microbes: Proteomic profiling |
| Other perturbations | Blood samples | Microbes: Metabolite profiling |
| Host characteristics | Plasma | Host: Whole genome sequencing |
| Activity level | Serum | Host: Transcriptome sequencing |
| Stress questionnaires | PBMC | Host: Proteomic profiling |
| Diet info | | Host: Clinical lab tests |
| Environment info | | | |

**Figure 3. Microbiome and Host Changes during Respiratory and Other Stress Conditions in Individuals at Risk for Type 2 Diabetes: Sample Collection, Assay, and Data Generation Workflow**

All samples are collected at the Stanford Clinical & Translational Research Unit. From each patient in every visit, blood sample and microbiome sample (including nasal swabs and stool and urine samples) are collected. Multi-omic data generation (primarily, but not entirely, nucleotide sequence- and mass spectrometry-based) will provide profiles of microbial phylogenetic composition, metagenomes, metatranscriptomes, and metaproteomes; host protein profiles, cytokines, and autoantibodies; and global metabolome profiles. Each sample is further accompanied by clinical (blood) or self-reported stress level, environmental, and dietary (stool and urine) metadata.

**Microbiome and Host Protein Expression Profiles Using LC-MS/MS:** Most studies analyze RNA expression; however, protein levels often do not correlate well with RNA expression levels (Maier et al., 2009). Therefore, microbial and host protein expression will also be analyzed from fecal samples and host PBMC cells, respectively. Proteins will be extracted, trypsin digested, and analyzed by 2D-LC and MS/MS (Chen et al., 2012). MS/MS spectra will be searched using the Sequest algorithm against human and microbiome protein databases and a specialized database of each individual based on the microbiome metagenome and host genome sequence.
Protein false discovery rates of 1% and only proteins with at least two peptides will be used. Relative protein levels using TMT isobaric tags and a reference sample prepared from multiple individuals or time points will be quantified (Chen et al., 2012). The dynamic changes in protein levels will be scored during healthy and stress periods.

**Cytokines and Autoantibody Profiles: Serum.** Since inflammation and immune responses have been implicated in T2D (Esser et al., 2014; Shu et al., 2012), and viral infections dramatically affected the immune response, the levels of 50 diverse cytokines (e.g., inflammatory proteins such as TNF-α, IFN-α, CRP, and insulin peptides) in host serum will be followed using throughput ELISA assays (Chen et al., 2012). Autoantibodies and antiviral antibodies will be profiled from selected sera using the Invitrogen ProtoArray Protein Microarray v5.0, which contains 9,483 unique human proteins (Hudson et al., 2007), and other comprehensive human and virome arrays.

**Metabolomes: Urine and Plasma.** Since diabetes is a metabolic disease (Akash et al., 2013), it is of particular importance to profile as many metabolites as possible (Friedrich, 2012), many of which are likely synthesized or influenced by the microbiome. To gain insight into the breadth of metabolic changes that occur, untargeted metabolomics profiling of urine and plasma samples will be performed using an optimized LC-MS procedure (liquid chromatography coupled to mass spectrometry). Two complementary separation methods will be combined and run in ESI (electrospray ionization) positive and negative modes to maximize the metabolome coverage from polar to nonpolar metabolites: HILIC (hydrophilic interaction liquid chromatography) and RPLC (reverse-phase liquid chromatography). The MS data will be acquired using high-sensitivity, high-resolution mass spectrometers. The optimized analytical procedure will monitor 18,000 and 14,000 metabolic features, characterized by a unique retention time and exact mass, for each urine and plasma sample, respectively. Metabolites will then be annotated using public databases of > 40,000 metabolites, and their identity will be confirmed with authentic standards when possible.

**Standard Clinical Tests: Plasma and Serum.** Standard clinical tests will be performed to follow typical plasma and serum markers. We will quantify lipids, lipoproteins, hsCRP, hemaglobin A1c, insulin, and glucose. The degree of hyperglycemia and insulin resistance of subjects will be measured. When patients are ill, standard nasal swabs will be used to identify the pathogen, which will complement the microbiome analyses. The clinical data may be redundant with the omics profiles presented above but will both serve as useful controls and ensure that no key metabolite is missed. In order to document behavior changes of patients, emotional and psychological stress will be quantified using the Perceived Stress Scale instrument, which measures an individual’s perception of stress (Cohen et al., 1983). Physical activity will be scored using the International Physical Activity Questionnaire (Hallal and Victoria, 2004), and food intake will be recorded for the last 24 hr before each visit.

**Conclusion**

These three examples of microbiome-associated human conditions will provide innovative, freely available, multi-omic data of properties analyzed in both the microbiome and host with which to explore the behavior of the human microbiome over the course of complex conditions using preterm birth, onset of IBD, and onset of T2D as dynamic model systems. In addition, they will help lay the foundation for shared protocols and analysis methods for highly detailed studies of the microbiome’s basic biology and translational potential. Data produced by the iHMP Consortium will be routinely released, and ongoing feedback is encouraged on how best to leverage these resources. For those interested in participating in the iHMP Consortium or learning more about the iHMP data sets or tools, please either contact the corresponding author or send an inquiry via the iHMP portal (http://ihmp2.org).

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes the complete list of co-authors of the iHMP Research Network Consortium, along with their affiliations and contact information, and can be found with this article online at http://dx.doi.org/10.1016/j.chom.2014.08.014.

**AUTHOR CONTRIBUTIONS**

This paper was co-authored by the iHMP Research Network Consortium, which includes the NIH iHMP program management, the iHMP project teams, and the iHMP DCC. The corresponding author is Dr. Lita M. Proctor, National Human Genome Research Institute, NIH, lita.proctor@nih.gov, (301) 496-4550. Please see Supplemental Information for a full list of the co-authors and their contact information.

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