

SHORT COMMUNICATION

The networks of human gut microbe–metabolite associations are different between health and irritable bowel syndrome

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The goal of this study was to determine if fecal metabolite and microbiota profiles can serve as biomarkers of human intestinal diseases, and to uncover possible gut microbe–metabolite associations. We employed proton nuclear magnetic resonance to measure fecal metabolites of healthy children and those diagnosed with diarrhea-predominant irritable bowel syndrome (IBS-D). Metabolite levels were associated with fecal microbial abundances. Using several ordination techniques, healthy and irritable bowel syndrome (IBS) samples could be distinguished based on the metabolite profiles of fecal samples, and such partitioning was congruent with the microbiota-based sample separation. Measurements of individual metabolites indicated that the intestinal environment in IBS-D was characterized by increased proteolysis, incomplete anaerobic fermentation and possible change in methane production. By correlating metabolite levels with abundances of microbial genera, a number of statistically significant metabolite–genus associations were detected in stools of healthy children. No such associations were evident for IBS children. This finding complemented the previously observed reduction in the number of microbe–microbe associations in the distal gut of the same cohort of IBS-D children.

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Introduction

Gastrointestinal (GI) tract of humans is a complex environment characterized by a high diversity of inhabiting microorganisms, complex metabolic and spatial interactions among community members, and intricate interplay between microbes and the host. Changes in the human gut microbiota have now been associated with a number of GI diseases, including Crohn's disease and ulcerative colitis, colorectal cancer and obesity (Sekirov *et al.*, 2010). Many of these microbial effects are exerted through the metabolic interactions between microbes and host tissues. Complex carbohydrates from the diet are degraded by gut microbes in anaerobic fermentation to formate, lactate, pyruvate, succinate (intermediates) and short-chain fatty acids (SCFAs, end products; Karasov and Carey, 2009). SCFAs have a major role in the microbiota–host interactions, for example, by inhibiting inflammation and malignant growth (Guilloteau *et al.*, 2010).

Microbial degradation of proteins produces branched-chain fatty acids, amino acids, amines, as well as harmful phenolic compounds, ammonia and hydrogen disulfide (Bernalier-Donadille, 2010).

Thus, simultaneous interrogation of microbes and metabolites of the intestinal environment can provide valuable insight into possible gut microbial interactions and allow better understanding of how microbiota alterations in diseases such as irritable bowel syndrome (IBS) lead to detrimental effects on patient's health. IBS is one of the most common disorders of the human GI system affecting 10–20% of the population with prevalence among adolescents (Quigley *et al.*, 2012). In this study, we obtained quantitative metabolite signatures and carried out correlative microbe–metabolite analysis of fecal samples from healthy children and those diagnosed with diarrhea-predominant IBS (IBS-D).

Results

Metabolite profiling of fecal extracts

We extracted water-soluble metabolites from 44 fecal samples obtained from healthy and IBS-D children (designated kHLT and kIBS, respectively, 22 samples

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per group) and subsequently subjected these fecal extracts to proton nuclear magnetic resonance-based metabolomics analysis (see Supplementary Figure S1). All spectra were divided into ~300 bins (see detailed description of all methods in Supplementary Materials), and these binned data were provided as input to several ordination analyses to reduce data set dimensionality and

visualize sample dissimilarity (Figure 1). Using a non-supervised principal components analysis, a partial, but statistically significant, separation of samples from kIBS and kHLT groups was achieved ($P=0.01$, Figure 1a). We previously revealed a similar partial separation of the same set of fecal samples based on the microbiota abundances obtained with phylogenetic Microbiota Array

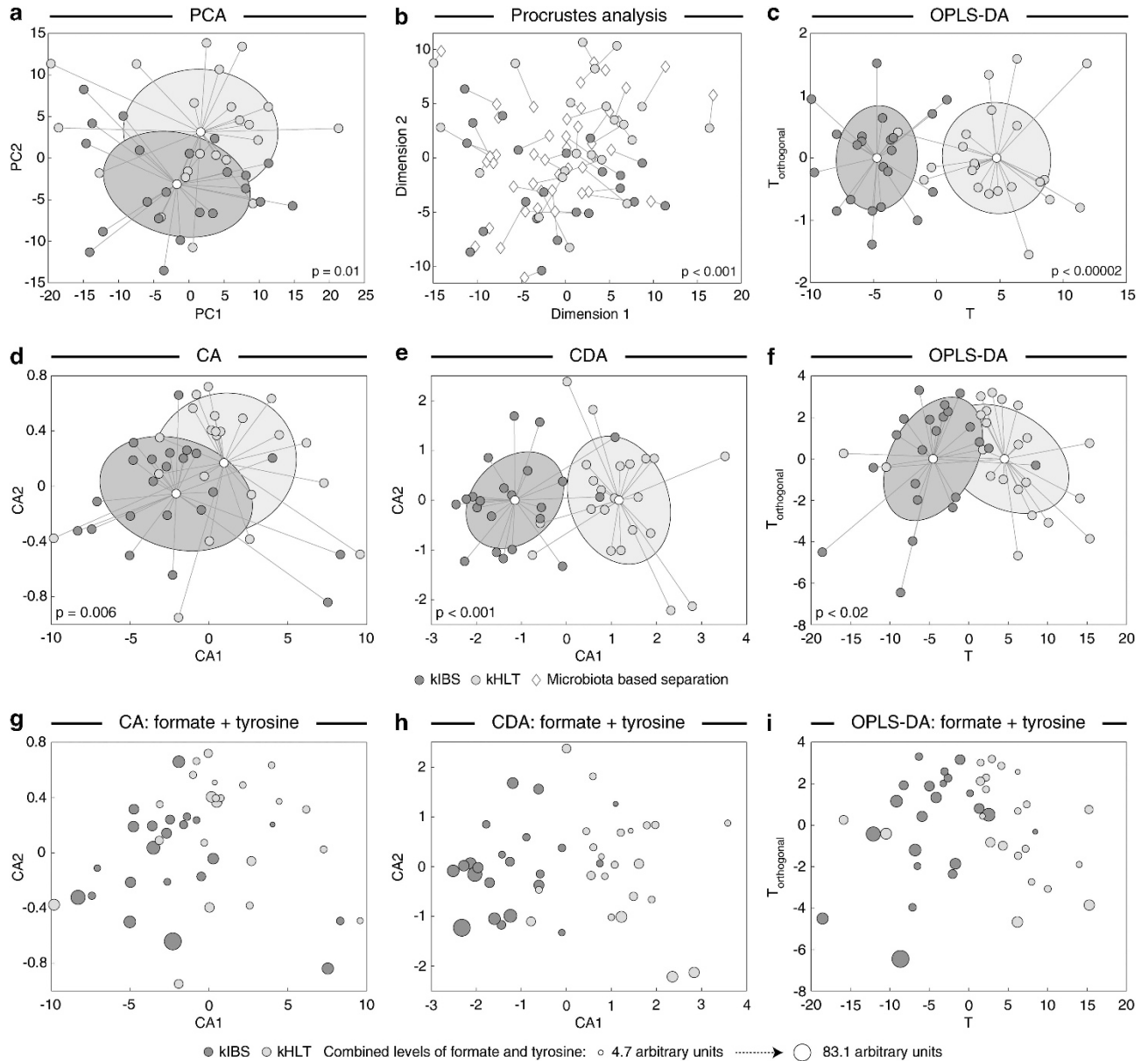


Figure 1 Separation of healthy and IBS fecal samples based on fecal metabolite profiling. Sample separation was assessed by the following ordination techniques: principal components analysis (PCA, **a** and **b**), correspondence analysis (CA, **d** and **g**), canonical discriminant analysis (CDA, **e** and **h**) and orthogonal projection to latent structures discriminant analysis (OPLS-DA, **c**, **f** and **i**). Sample separations in **a–c** are based on binned spectral data; sample separations in **d–i** are based on the levels of quantified metabolites. Panels **g–i** display CA, CDA and OPLS-DA sample distributions with the area of each sample spot proportional to the combined level of tyrosine and formate metabolites in that sample. P -values indicate the statistical significance of group separation. Where shown, group clouds represent areas of three standard errors around the centroid for each group. Panel **b** shows the results of Procrustes comparison of PCA separation of healthy and IBS samples based on binned metabolomic and microbial phylotype data sets. The distance between each sample position on two PCA plots is indicated by connecting line. Shorter lines represent more similar sample positioning on both plots. P -value indicates the statistical significance of sample separation congruency between two PCAs. kIBS, children diagnosed with IBS-D; kHLT, healthy children.

(Shankar *et al.*, 2013). To evaluate if both principal components analyses produced congruent sample distributions, Procrustes analysis, which seeks to superimpose, align and scale multiple graphical structures (Rohlf and Slice, 1990), was performed and revealed a strong congruency of sample separations based on metabolite and microbial signatures, respectively ($P < 0.001$, Figure 1b). As expected, because supervised orthogonal projections to latent structures discriminant analysis (OPLS-DA) takes into consideration the known group assignment (for example, kIBS versus kHLT) for each sample, OPLS-DA analysis of the binned spectral data revealed a considerably better sample separation between groups (Figure 1c, model significance $P < 0.00002$).

Focusing our attention on SCFAs, amino acids and compounds involved in energy metabolism, we quantified relative levels of 19 discrete fecal metabolites (see Supplementary Table 1 in Supplementary Materials). No significant difference in average total metabolite levels was evident between kIBS and kHLT groups. Statistically significant differences were observed for individual metabolites including formate, glucose, lactate, pyruvate and several amino acids (tyrosine, lysine and leucine). With an exception of pyruvate, kIBS samples contained higher levels of these metabolites. Total bile acids, one of the proposed causative agents of IBS development (Duboc *et al.*, 2012), were also higher in IBS, but the differences were not wide enough to overcome sample-to-sample variability in a significance test. Applying ordination techniques to the data set of measured metabolites revealed a statistically significant separation of samples between two groups (Figure 1d–i). OPLS-DA model as well as Random Forest analysis indicated that formate and tyrosine were the top two discriminating compounds.

Putative associations between intestinal metabolites and microbes

We computed the Spearman rank-based correlation matrix between metabolites measured here and microbial genus abundances assessed in the same set of samples in our previous report (Rigsbee *et al.*, 2012). We previously reported that microbe–microbe interactions were severely limited in IBS samples (Shankar *et al.*, 2013), therefore, metabolite–microbe associations were calculated separately for the kIBS and kHLT sample sets. Although many strong associations were discovered in the kHLT data set (Figure 2a), associations were weak in the kIBS sample cohort (Figure 2b). Simultaneous microbe- and metabolite-based bi-clustering of the correlation matrix revealed several local clusters of similar relationships in the kHLT data set. For example, all amino acids were clustered together, which can be attributed to their concurrent release through microbial proteolysis. Several groupings of microbial genera were also evident.

In contrast, little noticeable clustering was observed in the kIBS correlation matrix.

By employing a false discovery rate of 10% to control for multiple hypothesis testing, 21 significant metabolite–microbe relationships were identified in healthy teenagers (Figure 2c); none were found in the kIBS sample set. Among the statistically significant relationships revealed in the kHLT data set, several have been previously uncovered. These included strong positive association between glucose level and *Ruminococcus* abundance (glucose is released by ruminococci during fermentation), negative relationship between glucose and *Coprococcus* (some coprococci use glucose under anaerobic conditions), and positive associations of *Acidaminobacter*, *Coprococcus* and *Prevotella* with acetate, valerate and fumarate, respectively, likely explained by the observed production of these metabolites by the members of these genera (Holdeman and Moore, 1974, Bergey and Holt, 1994, Takahashi and Yamada, 2000, Duncan *et al.*, 2002, Iakiviak *et al.*, 2011). Abundances of genera *Bacteroides* and *Bifidobacterium*, which contain many members with large repertoires of carbohydrate-degrading enzymes, were not associated significantly with any of the measured fermentation-derived metabolites, likely due to the ability of these organisms to switch from one carbohydrate to another based on luminal availability and thus produce a variety of fermentation products (Martens *et al.*, 2011). The revealed associations of other genera with specific metabolites shown in Figure 2c have not yet been reported and represent putative novel interactions.

Discussion

The present study was undertaken to obtain quantitative metabolite signatures of fecal samples from healthy and IBS children, and to evaluate possible microbe–metabolite associations in the human gut. Overall, fecal samples from healthy and IBS adolescents could be distinguished based on the metabolite measurements of fecal extracts. Levels of formate, most of the identified amino acids, glucose and lactate differed between IBS and healthy samples. The presence of the higher levels of amino acids might be an indication of increased luminal proteolysis by IBS gut microbiota utilizing either diet-derived proteins or host mucins (Steck *et al.*, 2011). Elevated levels of fermentation intermediates in IBS stools might point to a disbalance of the gut environment and intestinal microbiota in this disease, leading to reduced metabolite cross-feeding among microbes and a resulting incomplete fermentation process. Because formate is utilized in the gut by methanogenic microbes, increased fecal formate is consistent with a lower methanogenic activity in the IBS-D lumen (Chatterjee *et al.*, 2007). And as intestinal methane adversely affects gut transit time

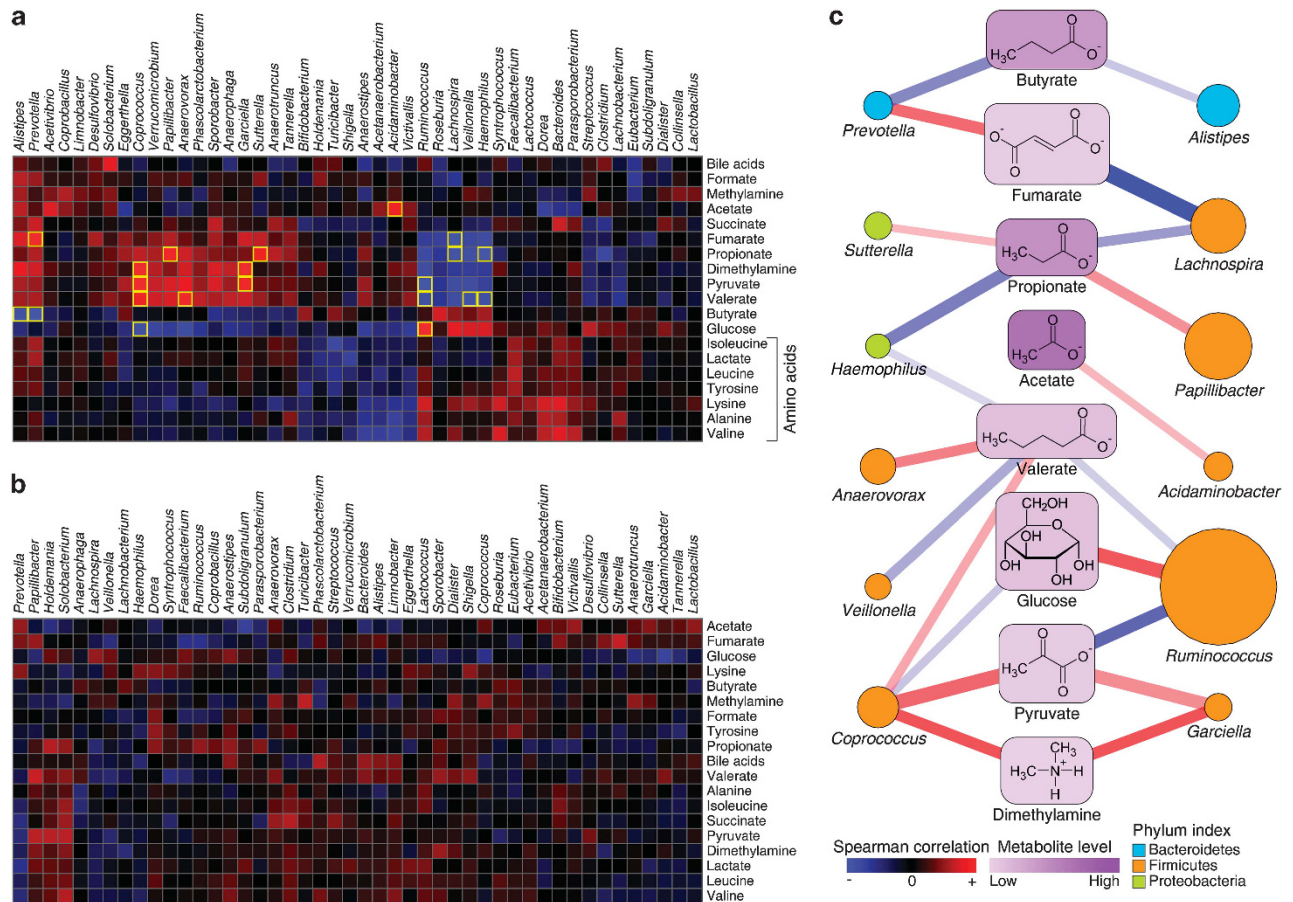


Figure 2 Correlations of genus and metabolite abundances among healthy and IBS samples. (**a** and **b**) Euclidean distance-based bi-clustering of Spearman rank correlation matrices between microbial genus abundances (46 genera with average abundance among all samples above 0.1%) and fecal metabolites in kHLT (**a**) and kIBS (**b**) samples. Each square represents a correlation value between specific genus and specific metabolite based on the color gradient as shown in the legend. Statistically significant correlations are highlighted with a yellow border. (**c**) A bipartite graph of the statistically significant metabolite–genus associations identified in the kHLT sample set. Positive and negative relationships are designated by red and blue colors, respectively; the line width and transparency are proportional to the absolute value of the Spearman rank correlation. Metabolite level is indicated by a node background color gradient as shown in the legend. The area of each genus node is proportional to the square root of the average genus abundance among all healthy samples. Genus assignment to a particular phylum is represented by the color of the node (see legend).

(Ghoshal *et al.*, 2011), lower methane production in IBS-D would facilitate faster gut transit resulting in a more frequent bowel movement.

A case for the loss of microbial interactions in IBS was also supported by our analysis of microbe–metabolite interactions. Although many strong associations were observed in healthy samples, such interactions were significantly weaker in IBS, with none reaching the level of statistical significance (see Figure 2). A concordant loss of microbe–microbe relative abundance associations in IBS was previously revealed (Shankar *et al.*, 2013). We offer several possible explanations of these findings. If IBS is associated with a disturbance of the gut environment, the resulting shifts in relative abundances of microbial taxa (Rigsbee *et al.*, 2012) may lead to a loss of the ‘steady-state’ microbial interactions within the healthy gut. The disruption of microbial metabolite exchanges would result in altered luminal and fecal metabolite levels we observed (Watanabe *et al.*, 2012). Alternatively, the

lack of strong metabolite–microbe interactions in IBS cohort might be an effect of the broad etiology of IBS. Different pathophysiological conditions that lead to IBS symptoms may result in unique shifts of microbiota community structure and/or functional capacity in different individuals. This, in turn, will either produce differentially altered luminal metabolite profiles, or a similar metabolic environment will be created by microbial communities of dissimilar compositions. Both cases will lead to an overall loss of observable cohort-wide interactions. Finally, although it is tempting to infer that the observed metabolite and microbial differences between healthy and IBS-D fecal samples are indicative of IBS pathophysiology, this syndrome is associated with many confounding factors such as change of eating habits or consumed diets, and alteration in stool consistency and frequency of defecation. Even though none of the children in IBS-D cohort were on any restricted diet or reported altered eating patterns at the time of sample

collection, and there was only a small difference in the average water content of the collected stool samples between healthy and IBS-D groups, both diet changes and diarrhea have been shown to cause alterations in human gut microbiota (David *et al.*, 2014, Gorkiewicz *et al.*, 2013). Thus, some of the observed IBS-associated effects on the gut environment might be manifestations of the concurrent changes in eating and bowel evacuation patterns in individuals with IBS-D.

Conflict of Interest

The authors declare no conflict of interest.

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Supplementary Information accompanies this paper on The ISME Journal website (<http://www.nature.com/ismej>)