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Heavy metal exposure causes changes in the metabolic health-associated gut microbiome and metabolites

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\textbf{ABSTRACT}

\textbf{Background:} Exposure to arsenic and cadmium is common. Epidemiological and animal studies have suggested that exposure to these two heavy metals can cause metabolic health problems, including type 2 diabetes (T2DM). It has been hypothesized that T2DM could be mediated through the gut microbiome and the metabolites it produces. Although many studies have investigated the association between the gut microbiome and T2DM, few have focused on the connection to arsenic and cadmium.

\textbf{Results:} We applied 16S rRNA gene amplicon sequencing and untargeted LC-MS/MS metabolomics to examine the changes in the gut microbiome and metabolite profiles of exposed mice to relevant levels of cadmium and arsenic in the drinking water over two weeks. Cadmium chloride (Cd) exposure significantly changed the mice gut microbiome and resulted in a significantly lower microbial diversity whereas sodium arsenite (As) caused a non-significant decrease in microbial diversity. For Cd and As treatment respectively, we identified 5 and 2 phyla with significant changes and 42 and 24 genera. Bacterial genera that were observed to decline upon both treatments, included several butyrate-producers. Both As and Cd treatment perturbed the metabolome significantly, with 50 ppm Cd compound exposure having the greatest effect when compared to 50 ppm As compound exposure. Two unidentified features were differentially abundant in the As group, while 33 features changed in the Cd group. Differential abundance analysis of all bile acid associated molecular components showed differences under both treatments. Finally, integrative network analysis via bipartite correlation networks suggested that several genera, including the metabolically important \textit{Blautia}, \textit{Eisenbergiella}, \textit{Clostridium_XIVa}, etc. declined in numbers of metabolite interactions.

\textbf{Conclusions:} These results demonstrated that As and Cd exposure caused significant changes to the gut microbiome and metabolome by affecting bile acids, amino acids and taxa associated with metabolic health.

\section{1. Introduction}

Diabetes mellitus is a worldwide metabolic syndrome, which is characterized by fasting hyperglycemia, insulin secretory dysfunction or insulin resistance (Kim et al., 2001). According to a report from the International Diabetes Federation, the diabetic population now stands at about 382 million (8.3% of adults) globally and is estimated to rise to 592 million people (10% of adults) by 2035 (Guariguata et al., 2014).

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Among the different types of diabetes, type 2 diabetes (T2DM) is the most prevalent, accounting for 90–95% of all cases (Wild et al., 2004). Traditional T2DM risk factors include age, obesity, lifestyle and, in some rare instances, genetic predisposition (Jääskeläinen et al., 2013). However, these factors seem insufficient to explain the mounting worldwide T2DM epidemic. Some epidemiological studies have shown that environmental chemicals may play an etiological role in the progression of T2DM (Jeon et al., 2015). As exposure has been linked to the impairment of glucose metabolism (Jeon et al., 2015). Hundreds of millions of people around the world live in regions where natural As concentration in drinking water far exceeds the safety standard of 10 μg/L accepted by the World Health Organization and the U.S. Environmental Protection Agency (EPA) (Chen et al., 2009; Hughes et al., 2011). The occurrence of As contamination in soil and ground water caused by human activities is also very common (Jeon et al., 2015). Cadmium (Cd) is also known as a highly toxic heavy metal that poses increasing risk to populations in many parts of the world (Diarra et al., 2006; Edwards and Prozialek, 2009). Studies suggest that low but chronic levels of Cd exposure can impair the function of insulin-producing β-cells and may be associated with T2DM (El Muayed et al., 2012). Cigarette smoke and diet are among the main routes of Cd for nonoccupational Cd exposure (Madeddu et al., 2009; El Muayed et al., 2012). Despite high toxicity and easy accessibility, so far, only a handful of studies mainly from epidemic and animal physiological research provide proofs for the adverse effects of the two metals on health.

Accumulating evidence from animal and human studies suggest that alteration of the gut microbiome through dietary manipulation and the presence of environmental pollutants may disturb physiological and metabolic homeostasis, contributing in part to the development of various diseases (Everard and Cani, 2013; He et al., 2015; Larsen et al., 2010). As and Cd exposure have been reported to result in changes of gut microbiota and metabolic profiles in some exploratory mice studies (Liu et al., 2014; Lu et al., 2014), however T2DM was not the study target. T2DM has also been reported to be closely associated with the gut microbiome (He et al., 2015; Larsen et al., 2010; Qin et al., 2012). For example, gram-negative bacteria in the gut have received much attention for their role in the development of diabetes (Amar et al., 2011; Cani et al., 2007). A major compound in the outer membrane of gram-negative bacteria is lipopolysaccharides (LPS), which are potent stimulators of the immune system causing inflammations and endotoxemia (Nielsen et al., 2003). Consequently, the continuous LPS production within the gut might trigger an inflammatory response and further promote the development of diabetes. Furthermore, some specific groups of gram-negative bacteria have received more attention recently, as they have been linked to gut microbiome dysbiosis. Akkermansia muciniphila, an important intestinal mucus-degrading gram-negative bacterium that accounts for densities of up to 3% of the total number of bacteria in adult feces, was linked to diabetes and obesity (Everard et al., 2013; Plovier et al., 2017). In this study, we specifically focused on gram-negative bacteria by treatment with vancomycin to enrich for this type of bacteria in a murine model. This treatment is known to induce a dysbiotic state in both humans (Vrieze et al., 2014; Zarrinpar et al., 2018) and mice (Hansen et al., 2012). The result of this treatment impacts a wide range of metabolic outputs from the microbiome such as bile acids and short chain fatty acids. To observe the effect of the intervention, control mice were also treated with vancomycin. We used a combination of 16S rRNA gene sequencing and liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS) based metabolomics to analyze the alterations in the murine gut microbiome and its metabolic profile induced by exposure to As and Cd.

2. Material and methods

2.1. Chemicals, animals, and experimental protocol

Five-week-old C57BL/6 mice were purchased from the China National Laboratory Animal Resource Center. All the operations in the animal experiment were followed by the 3R principle (Russell and Burch, 1959). A total of 48 mice (body weight = 20 ± 3 g) were housed in static microisolator cages (4 mice/cage) under environmental conditions of 22 °C, 40–70% humidity, and a 12:12 h light:dark cycle. Water and food was available ad libitum during the whole experiment. Mice were treated with 500 mg/L vancomycin in the drinking water for one week after one-week initial adaption to cage-food and laboratory conditions. Afterwards, the mice were randomly divided into 3 experimental groups containing 16 animals each (4 mice/cage). One group, serving as a control drank water free of contaminants, and the other two groups were separately treated with 50 ppm cadmium chloride (Cd) (equals to 0.273 mmol/L Cd2+) or 50 ppm sodium arsenite (As) (equals to 0.385 mmol/L As3+) supplied in the drinking water for 2 weeks. Mice were assessed every day for signs of diarrhea, dehydration, mortality and deteriorating body conditions. Body weight, food and water intake were assessed twice per week during the Cd/As treatment. All mice were sacrificed after anesthetization with ether on the last day of the treatment. Colon and caecum content were collected and stored at −80 °C before use. During the experiment, three animals in the As group died from fight trauma.

2.2. 16S rRNA gene sequencing and analysis

Genomic DNA was extracted from colonic content with the QIAamp PowerFecal DNA Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instruction. All operations were performed under aseptic conditions. The NanoDrop spectrophotometer (Thermo Scientific NanoDrop Products) was used to check DNA concentration and a 1% agarose gel was used to test DNA quality. The 16S rRNA gene hypervariable V3-V4 region was amplified with 1 μL template DNA (10–50 ng/μL), using 12.5 μL KAPA HiFi HotStart Ready Mix (Anachem, Dublin, Ireland), 0.25 μL 25 μM of each primer were PCR amplified using 25 cycles with the primers Bakt_341F (CCTACGGGNGGCWGCAG) and Bakt_805R (GACTACHVGGGTATCTAATCC) (Klindworth et al., 2013) in a 25 μL PCR reaction volume. The first PCR program included 3 min at 95 °C, 25 cycles of 30 s at 95 °C, 30 s at 55 °C, and 72 °C for 30 s, and then 5 min at 72 °C. In the second PCR, the sequencing primers and adaptors were attached to the amplicon library following the first PCR conditions with only 8 cycles. The size of the PCR product was evaluated using gel electrophoresis. The amplicon products were purified by use of Agencourt AMPure XP Beads (Beckman Coulter Genomics, MA, USA) and were quantified by densitometry using Quantity one software (Bio-Rad, Hercules, CA, USA), then pooled in equimolar concentrations. The purification of DNA library was done by gel extraction with the Qiagen gel extraction kit (Qiagen, Hilden, Germany). The concentration and length distribution of DNA library was checked by Qubit Fluorometer (Invitrogen, Carlsbad, CA, USA) and Qseq100 (BioOptic Inc., Taiwan, China). Sequencing of V4 region of bacterial 16S rRNA genes was performed with Miseq PE300 platform as previously described (Lu et al., 2015). The obtained paired-end raw data was assembled by Flash (Magoc and Saltberg, 2011). Quality filtered, Length trimming, Homopolymer truncation were processed by mother (Schloss et al., 2009). OTU clustering and Taxonomy classification was carried out by UPARSE, v9.0.2132 (Edgar, 2013), sorting 1,937,439 reads into 662 OTUs at 97% sequence homology. The open source statistical program “R” was used for data treatment and analysis (R Development Core Team, 2011), predominantly the R-package “phyloseq” (McMurdie and Holmes, 2013). Alpha diversity between the groups was tested by analysis of variance, using the R function “aov”. UniFrac distance (Lozupone and Knight, 2005), ordered by principal
coordinates analysis (PCoA, R function “ordination”, R-package “phyloseq”), was examined by permutational multivariate analysis (PERM-ANOVA, R function “adonis”, R-package “vegan”) (Oksanen et al., 2008). Identification of features differentially abundant between samples was performed by random forest classifier and differential expression analysis based on the Negative Binomial (a.k.a. Gamma-Poisson) distribution (R function “differential_abundance” in R-package “MicrobiomeSeq”, https://github.com/umerijaz/microbiomeSeq). Mean decrease in accuracy (the decrease of accuracy due to the exclusion of a specific feature) was used to evaluate the importance of the feature.

2.3. Sample preparation and mass spectrometric analysis

Approximately 180 mg fecal samples were suspended in water, a subsample of each was mixed with acetonitrile to a final concentration of 1% and acidified with formic acid to a final concentration of 0.5% to counteract the buffer capacity of the samples. Samples were filtered on 3000 Da Amicon Spinfilters to remove debris. The samples were analyzed by liquid chromatography tandem mass spectrometry (LC-MS/MS) and data was recorded in a data dependent manner, on a Q-Exactive (Thermo Scientific, Bremen, Germany). An EASY nLC-1000 liquid chromatography system (Thermo scientific, Odense, Denmark) was coupled to the mass spectrometer through a 2 cm C18 pre-column (300 μm inner diameter, 1.9 μm particle size) and a self-packed 15 cm C18 column (Pico Frit columns from New Objective) with a 75 μm inner diameter, packed with (1.9 μm C18 Dr. Maisch Mat. No. r119.a). Metabolites were eluted with a mobile phase consisting of solvent A (0.1% formic acid) and B (80% acetonitrile in 0.1% formic acid). The concentration of solvent B was linearly increased from 2 to 10% over 3 min and from 10 to 95% solution B over 15 min. Solvent B was maintained at 95% for 2 min. Full scans were acquired in the Orbitrap with a resolution of 70,000, maximum injection time of 20 ms and a scan range of 80-1200 m/z using a Top 10 method with an isolation window of 1.6 Da and a dynamic exclusion time of 15 s. For the MS/MS scans the resolution was adjusted to 17,500 and maximum injection time of 60 ms. Ions were fragmented with a normalised collision energy (NCE) of 30%.

2.4. Metabolomics processing and analysis

LC-MS/MS raw data files were converted to .mzXML using msconvert from ProteoWizard (v. 3.0) and preprocessed using MZmine 2.3 (Pluskal et al., 2010) with parameters set to Peak detection, Mass detection, Mass detector: Centroid, MS1 noise level 2.0E5, MS2 noise level 100; Chromatogram builder: MS1 level, Min time span (min) 0.01, Min height 6.0E5, m/z tolerance 0.01 Da or 10 ppm; Chromatogram de-convolution: Algorithm Local minimum search, Chromatographic threshold 90%, Search minimum in RT range (min) 0.01, Minimum relative height 10%, Minimum absolute height 1000, Min ratio of peak top/edge 1.3, Peak duration range (min) 0.01–10.00, m/z range for MS2 scan pairing (Da) Off, RT range for MS2 scan pairing (min) Off; Isotopic peak group: m/z tolerance 0.01 Da or 10 ppm, Retention time tolerance 0.5 (min), Maximum charge 4, Representative isotopes Most intense; Feature alignment: m/z tolerance 0.01 Da or 10 ppm, Weight for m/z 75, Retention time tolerance 0.5, Weight for RT 25; Gap filling: Intensity tolerance 10%, m/z tolerance 0.01 m/z or 10 ppm, Retention time tolerance 0.5 (min), RT correction Off. Mass spectral molecular networks (Watrous et al., 2012) were created by uploading the MZmine preprocessed .mzXML file to the Global Natural Products Social Molecular Newtorking platform (GNPS) (Wang et al., 2016) with parameters described at https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=806fa2cdec63b41f59b17b59c2a5da0ee.

Before statistical analysis, features not found in a minimum of 20 samples were filtered out, reducing dimensions from 8562 to 2130. Upon visual inspection of a PCA plot of all samples, Sample36 (Cd group) were discarded as an outlier. The best performing univariate statistical test were determined using the R-package “DAtest” (https://github.com/Russell88/DAtest) with default settings. The t-test performed the best in terms of controlling the false positive rate at 0.05, while having the best ROC of the methods tested. The tests were then performed using the R function “DAtest.tte”, and evaluated by the FDR corrected p-values, with 0.05 deemed significant. PCA analysis was done using the R function “prcomp” on square root transformed features. Multivariate hypothesis testing was performed using the R function ”adonis” in R-package “vegan” (2.4-4) (Oksanen et al., 2016) with the Euclidian distances of square root transformed features.

To putatively identify molecular features, we used both GNPS library matching as well as in silico structure annotation by the Network Annotation Propagation (NAP) tool (da Silva et al., 2018). Due to the aforementioned interest in bile acids, we submitted a user provided database to NAP. This database was compiled by downloading all metabolites in the “Bile acids and derivatives [ST04]” category of LIPID-MAPS and is publically accessible at Additional file C. NAP parameters are described and can be accessed at: https://proteomics2.ucsd.edu/ProteoSAFe/status.jsp?task=4844738f1ed8b4c49a836badbc9c5b280 https://proteomics2.ucsd.edu/ProteoSAFe/status.jsp?task=232cf5a822864a31aeb6063b957c11c2

2.5. Concordance analysis

Correlations between OTUs and metabolites were calculated using the R function “protest” in the R-package “vegan” (2.4-4) (Oksanen et al., 2016).

2.6. Integrative network analysis

Integrative network analyses were performed in R. Pearson correlations were calculated for a bipartite network of OTUs and metabolites using a Fisher transformation, and correlations at the 0.05 FDR level were included (Kolaczyk et al., 2014). Both OTU and metabolite tables were pruned for near zero variance features, using the NearZeroVariance function from the “caret” package (Kuhn, 2008). Networks were managed and visualized in the R-package “igraph” (1.2-2) (Csardi, 2015). Differential networks were calculated using fishers transformation of correlation coefficients as shown in the R-package “Diffcorr” (Fukushima, 2013). Briefly, a network of Pearson correlations was calculated for each condition (control, As and Cd treatment), where links were defined by significant correlations after FDR adjustment. The difference in correlations (r) between samples A and B are then calculated by transforming each coefficient into a Z-score (Z) by $Z = 0.5 \log \left( \frac{1 + r_{A}}{1 - r_{A}} \right) \frac{1}{\sqrt{1 - r_{A}^2}}$. $p$-values were calculated assuming a normal distribution. Taxa enriched in differential links were identified by permuting the taxa label 100 times and performing a one-sided evaluation. Adjustment for permutation p-value misestimation was applied (Phipson and Smyth, 2010). To further avoid false positives all taxa with prevalence lower than 1/3 presence in the condition were set to 0.

3. Results

3.1. Sequencing summary and phenotype assessment

The experimental workflow combining 16S rRNA gene sequencing and metabolite profiling to test the impact of As and Cd exposure on the gut microbiome and its metabolism, is shown in Fig. 1. The V3-V4 region of 16S rRNA gene amplicons from fecal samples from a total 31 subjects (control group (n = 13), As group (n = 7), Cd group (n = 11))
(14 samples missing because of no amplification) were sequenced using the Miseq PE300 platform. The average sequencing depth obtained for the 31 subjects was 142,807 reads per sample, which was reduced to 62,498 (SD = 24,455) high quality sequences after QC. Body weight, food and water consumption during the experiment for all the groups did not differ significantly, as shown by Wilcoxon Rank Sum test (see Tables S1 and S2).

3.2. Exposure to cadmium significantly altered the gut microbial diversity

In order to investigate the changes in gut microbial community triggered by the different treatments, we compared the microbial diversity of the different treatment groups. As shown in Fig. 2A, we found that both treatments caused a decrease in diversity of the gut microbiome, but only to a significant degree in the Cd treated mice ($p = 0.028$). A clear distinction was also observed between the microbial communities of the control group and the Cd group, as shown in the PCoA plot (Fig. 2B). The control and treated mice were significantly separated, with 42.6% and 22% variation explained by principle component PC1 and PC2, respectively. The Cd treatment explained 25% of the variance ($p = 0.003$), while the As treatment explained 2% of the variance ($p = 0.8$). No significant differences in microbial diversity were observed between the As group and the control group (Fig. 2B).

3.3. Heavy metals induced significant changes in the gut microbiome

The four major phyla composing the microbial community are listed in Fig. 3A. The phylum Verrucomicrobia mostly composed of the genus Akkermansia, was the second most abundant phylum, verifying the enrichment effect of vancomycin on gram-negative bacteria.

A random forest classifier approach was used to explore the specific changes in the gut microbiome related to the exposure to heavy metals.
Changes in the gut microbiome caused by the different treatments. Two phyla in the As group and five phyla in the Cd group were differentially abundant compared to the control group (Fig. 3B and C). In both groups, Bacteroidetes rank first in importance by the random forest classifier analysis. The abundance of Bacteroidetes was significantly higher in the control group as compared to both As ($p = 0.002^{**}$) and Cd ($p = 1.6e-16^{***}$). Proteobacteria in the Cd group and Tenericutes in the As group were less abundant than in the control group ($p = 2.03e-06^{***}$ and $p = 0.006^{**}$ respectively). In addition, the abundance of Deferribacteres, Cyanobacteria and Candidatus_Saccharibacteria in the Cd group showed significant differences from the control group. Further information about these significantly changed phyla has been listed in Tables S3 and S4. The arrows inside the middle section of the plot indicate the upregulation (blue arrow) or downregulation (red arrow) of the corresponding phylum abundance (Cd/As vs Control). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

![Figure 3](image_url)

Fig. 3. (A) Distribution of phyla between experimental groups. (B,C) Significantly differentially abundant phyla between Cd (B)/As (C) and control group: The boxplots are the log relative normalised abundances for the treatment groups. The bar plot correspond to importance of a corresponding feature based on mean decrease accuracy and on bottom of which the ranks are indicated. The middle section of the plot shows phyla description and corresponding adjusted $p$-value. The arrows inside the middle section of the plot indicate the upregulation (blue arrow) or downregulation (red arrow) of the corresponding phylum abundance (Cd/As vs Control). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
42 different genera in the Cd group. Further, 9 of the downregulated genera (Clostridium_XIVb, Syntrophococcus, Cellulosilyticum, Blautia, Anaeroplasma, Gemmiger, Anaerovibrio, Intestinimonas, Parasutterella) and 2 of the upregulated genera in both groups (Alistipes and Alkalitalea) showed overlap between the As and Cd groups. Additional information for the significantly changed genera has been listed in Tables S3 and S4. Fold changes for all the genera in both groups have been shown in Fig. S2.

Due to the vancomycin treatment, which resulted in accumulation of gram-negative gut bacteria, Akkermansia was the most abundant genus in both groups (Fig. S3). Despite detected high sensitivity to As and Cd in vitro (Fig. S4), only the As group showed a decrease in abundance of Akkermansia (1.8 fold) whereas the Cd treatment showed a surprising 1.25 fold increase.
3.4. Heavy metals induced global changes in the gut metabolome

Metabolic profiles of the fecal contents were generated using liquid chromatography coupled to tandem mass spectrometry (LC–MS/MS), including 16 control samples, 13 As group samples and 15 Cd group samples. Mass spectral molecular networks (Watrous et al., 2012) were created using the online Global Natural Products Social Molecular Networking platform (GNPS) (Wang et al., 2016). Processing of spectra resulted in 26,978 features for all the samples. Rarefaction curves of feature richness showed little difference between the As and control groups, but there were fewer observed features in the Cd group (Fig. S5-A). Multivariate differences between treatment and control were assessed using PERMANOVA and PCA. The As treatment explained 5.1% of the variance in the metabolome (p = 0.0456*), while the Cd treatment explained 7% of the variance (p < 0.001***). PCA analysis (Fig. 6) visually separated the different groups. Both treatments induced the presence of features not found in the controls. Cd treatment showed 111 uniquely present features while As treatment showed 11.
molecular relatedness. The network was quite homogenous, providing additional evidence for the hypothesis testing. This resulted in 2 and 33 features identified for every feature in each group and families (Fig. 7A). The recorded Closer inspection of the molecular network showed that many unique associated with bile acids according to a GNPS library search, implying they were structurally related to bile acids. (B) An Fig. 7. Bile acid analysis of the metabolome. (A) Investigation of features uniquely found in As treated mice revealed that most were found in molecular families (ST04) category of LIPID-MAps were downloaded and fragmented in silico to compare with the acquired fragmentation spectra. This identified 633 features as bile acid related. In the previously constructed molecular network they were hydroxycinnamic acid, and manual inspection of the spectra confirmed this. The latter being strongly decreased in the treated mice with a log fold change of \(-15\).

3.5. Heavy metals induced changes in the bile acid fraction of the metabolome

Due to the bile acids associated molecular networks unique to As treatment, we putatively annotated the bile acid fraction of the fecal metabolome using in silico structure annotation through the recently available Network Annotation Propagation algorithm (NAP). All metabolites in the "Bile acids and derivatives [ST04]" category of LIPID-MAps were downloaded and fragmented in silico to compare with the acquired fragmentation spectra. This identified 633 features as bile acid related. In the previously constructed molecular network they were...
distributed in 181 different molecular families (two or more connected molecular signatures). Aggregating the intensities of these components, we created a boxplot to show differentially abundant (DA) molecular families of putative bile acid molecules in the As group (Fig. 7B). We identified 6 DA molecular families in the As treatment and 2 DA molecular families in the Cd treatment. The intensities of 5 DA bile acid molecular families in the As group elevated but one DA bile acid molecular family decreased.

3.6. Bipartite interaction network identified bacteria-metabolite interactions in Cd associated bacteria

We examined the concordance between metabolite and microbiome data by Procrustes correlation which was significant (protest, correlation $= 0.65, p = 0.006$), suggesting that a fraction of the fecal metabolites are substrates or products of the microbiome.

To further investigate how this connection might change upon heavy metal treatment, bipartite correlation networks were created from the paired microbiome-metabolome tables. This resulted in three networks (not listed), of which the control treatment had the fewest interactions and the Cd treatment had by far the most interactions, despite the Cd-treated mice metabolomes having the fewest detected features. To identify the effects of the treatment on these networks, differential networks in which the difference in correlations between control and treatment were calculated. At the same time, a permutation test was applied to compare the number of changed links to the null hypothesis of random changes to identify genera with potential changed metabolic output. Correspondingly, we identified a number of significant metabolite interactions within each genera from these networks.

Several genera with significant changes in the number of metabolite interactions were identified. We found 7 genera which presented significant changes to their differential networks in the Cd treatment and 3 under the As treatment. Clostridium_XIVa had a large amount of changes in both treatments. All differential correlation networks are presented in Fig. 8. While the genera had overall significantly different interaction patterns, there were clear within-genus differences at the OTU level. Fig. 8 illustrates Blautia metabolic interaction changes, while OTU_7 primarily increased interactions, OTU_356 clearly decreased its interaction strengths with many metabolites. In other cases such as Eisenbergiella, several OTUs regulated their interaction with several metabolites in both treatments, suggesting a shared response to the treatment. The differential network for the two Anaerostipes OTUs under cadmium treatment is entirely disconnected.

4. Discussion

The current experiment was set up to connect the associations from the epidemiology literature concerning induction of type II diabetes with the observations from the microbiome literature, and hopefully narrow the role of the metabolome. In this study we hypothesized that Cd and As treatment would change the gut microbiome in mice. The tendency of significantly decreased microbial diversity in the Cd group and reduced microbial diversity in the As group observed in this study (Fig. 2), indicated that the response to toxic metals pose an adverse challenge to many gut bacteria. The relative abundances of Bacteroides in the Cd and As groups were significantly higher compared to the control group (Fig. 3), and the relative abundance of Proteobacteria in the Cd group was significantly lower. The trends regarding change in abundance for these two phyla are supported by current studies relevant to T2DM. Yet, there were still some overlapping findings for taxa in Clostridiales, but overall this strategy did not induce a difference in the microbiome as reflected by multivariate analysis. Similarly there was a small overlap in the metabolome, and more fatty acids were detected in Lu et al., possibly due to methanol extraction rather than water as in the present study.

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Fig. 8. Integrative bipartite networks. Networks of Pearson’s correlations were calculated to investigate interactions between OTUs and metabolites. (A) Genera with significantly changed numbers of interactions under the Cd treatment (adjusted $p < 0.01$, the connections with the absolute value of weights below 5 in the genus Clostridium XIVa were filtered out). (B) Genera with significantly changed numbers of interactions under the As treatment (adjusted $p < 0.01$, the connections with the absolute value of weights below 5 in the genus Clostridium XIVa were filtered out). Eisenbergiella, Blautia and Anaerostipes have several OTUs, some of which display a finer scale structure.
Fig. 8. (continued)
further improved oral glucose tolerance and reduced food consumption (Pichette et al., 2017). We also found that a series of genera belonging to the butyrate-producing superfamily Lachnospiraceae within the order Clostridiales that were downregulated in both groups, which included Fusobacterium, Eisenbergilla, Syntrophoccus, Blautia, Clostridium_Clostridium_XIVB, Cellulosilycitum, Orbaciterium, Coprococcus, Anaerostipes, Hespellia, Cellulosilycitum and Lachnospiraceae_Incertae_Sedis. Besides, the relative abundances of some genera belonging to family Erysipelotrichaceae, which is distantly related to Lachnospiraceae, Clostridium_Clostridium_XVIII, Kandleria and Erysipelotrichaceae_Incertae_Sedis, were decreased in the treatment group. Butyrate produced by gut bacteria is a major energy source for the colon epithelium (Roediger, 1980) and could improve insulin sensitivity and increase energy expenditure (Gao et al., 2010). Besides, genera within butyrate-producing families, the other downregulated genera shared by the As and Cd groups, such as Parasutterella and Gemmiger, have also been associated with alleviation of T2DM (Xu et al., 2015).

Akkermansia was not as sensitive to Cd and As in vivo as in vitro experiments. Although the relative abundance of Akkermansia was higher in the Cd group and lower in the As group, the changes were not substantial and not statistically significant. We assumed that the intestinal microbial consortia tend to be cooperative (Bäckhed et al., 2005) and this could further weaken the adverse effects from exogenous metals on a single species, which may explain the different results in vivo and in vitro. Some bacteria with probiotic properties such as Faecalibacterium and Bifidobacterium which have displayed beneficial effects on glucose metabolism in humans (Cani et al., 2009; Hill et al., 2014; Miquel et al., 2014) were also detected in our study. In accordance with these outcomes, both Faecalibacterium and Bifidobacterium decreased slightly in the Cd and As groups.

Despite large improvements in LC-MS/MS library searching technology and expansion of the libraries, identifying features is still not perfect, and overall in this study we could identify 3770 library hits out of 26,978 features (13.9%) using the GNPS platform. Previous summaries on the studies of metabolomics have estimated an average of 2% identification rates of features (Aktenov et al., 2017). One advantage of using the GNPS platform is that the data is continuously being updated with improved annotations (Wang et al., 2016). Of the differentially abundant features identified from the Cd group, several of them could be annotated as amino acids through library matching. It is well-established by serum metabolomics that the concentrations of amino acids change in T2DM subjects. This applies to amino acids in general (Guasch-Ferré et al., 2016), but branched chain amino acids (BCAAs) have, in particular, been associated with insulin resistance, and the microbiome has been demonstrated to contribute to the increased concentrations of amino acids in serum (Pedersen et al., 2016). At present, the causal path to insulin resistance remains unclear with Mendelian randomization studies suggesting that insulin resistance preceded increased BCAA levels (Mahendran et al., 2017). However, this accounts only for the variation caused by genetics. Additionally, it is unclear how amino acid levels in feces translate to amino acid levels in serum. A previous study observed that the main site of murine amino acid metabolism was higher in the gastro-intestinal tract, peaking in the jejunum (Heinzmann and Schmitt-Kopplin, 2015), possibly blurring the observations of caecal amino acids.

There are a limited number of feces metabolomic studies in T2DM but high-fat diet fed mice are known to have higher levels of tyrosine in fecal metabolomic phenotyping using an untargeted NMR investigation (Lin et al., 2016), which are consistent with our observations in the present study. In addition, in the present study, the abundance of 4-hydroxyphenylacetic acid which is an intermediate of several pathways, including the degradation of tyrosine deceased significantly by 15 fold in the Cd group. Tyrosine in blood has previously been identified as increasing the T2DM risk in a meta-analysis of 8000 metabolomic samples (Guasch-Ferré et al., 2016). In the present study, there was an increase in valine abundance after Cd exposure. Serum valine in humans has been linked to the visceral deposition of fat which has strongly been suggested to drive insulin resistance (Lebovitz and Banerji, 2005; Schlecht et al., 2017). Previous studies on insulin resistant rats using NMR showed consistent longitudinal increases in isoleucine (Yang et al., 2015), a degradation product of valine (Zarling and Ruchim, 1987). A metabolomics study of beta-cell function in insulin secretion had previously identified intracellular norleucine levels as statistically associated with insulin secretion (Huang and Joseph, 2014). However, connecting this finding to the fecal content would require a further investigation of the norleucine homeostasis.

We specifically chose to focus on the fraction of the metabolome that could be assigned to bile acids, as changes in this fraction has previously been associated with metabolic problems in humans. Additionally, it has been shown that microbes can actively manipulate this fraction of the metabolome by making an array of chemical modifications that alter the potency of the impact of a bile acid. Bile acid concentrations have been correlated with the microbiome in germ free mice (Claus et al., 2008) and shown implications in obesity in feeding studies (Lin et al., 2016; Walker et al., 2014). Our results displayed a change in features of the bile acid fraction of the metabolome. These findings should be followed up with more targeted metabolomics to identify the exact molecular identities of the bile acids.

We built the integrative networks to identify exactly which OTUs had altered interactions with the metabolites, as it has been hypothesized that deteriorating metabolic health is mediated through microbiome interaction with the metabolites, either through production or through consumption. We demonstrated that certain genera seem to have enriched or lowered metabolic interactions. This is particular noteworthy for the genera, Blautia, which had also been highlighted in previous literature for their metabolic capabilities, that showed a large number of interactions in the control network, but had a decreased number of interactions in both treatments. Similarly, Anaerostipes species had been shown to aggravate DSS induced colitis, presumably through its metabolic output (Zhang et al., 2016). These genera have been implicated in several types of metabolically related human health issues, presumably because of their butyrate producing capacity. In addition, the relative recently isolated genus Eisenbergilla displayed significant interactions after both treatments. The interaction networks at the OTU level made it clear that there were finer scales of interactions, with certain OTUs losing many interactions and some showing no difference under treatment conditions. This will be a starting point for a more targeted investigation at the species/strain level and for the metabolites that could reveal the exact interactions.

5. Conclusion

We have conducted an exploratory feeding study in mice to investigate the impact of heavy metals mainly on the fecal gram-negative bacteria enriched by Vancomycin and its metabolome to validate the hypothesis generated from epidemiology and cell culture studies that heavy metals can impact host metabolic health. Heavy metals do indeed impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice. Cadmium exposure significantly changed the mice gut microbiome as well as resulting in a significant impact both microbiome and metabolome, without substantially affecting the phenotypes of the mice.
their metabolites. More research is needed to clarify the exact nature of these metabolites and their impact. 

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Ethics approval and consent to participate

The animal research was approved by China National Accreditation Board for Laboratories and China Food and Drug Administration (Ethical approval number: BLARC2015A1203).

Consent for publication

Not applicable.

Availability of data and materials

The 16s rRNA gene sequencing dataset analyzed during the current study are available, upon publication, in the Sequence Read Archive (SRA) repository, https://www.ncbi.nlm.nih.gov/bioproject/PRJNA476308/. LC-MS/MS raw data files are publicly accessible under the MassIVE https://massive.ucsd.edu/

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Competing interests

The authors declare that they have no competing interests.

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References


