CASE REPORT

Case Report: The effect of intravenous and oral antibiotics on the gut microbiome and breath volatile organic compounds over one year [version 2; peer review: 2 approved with reservations]

Farah Shahi¹,², Sarah Forrester¹, Kelly Redeker¹, James P.J. Chong¹, Gavin Barlow²,³

¹Department of Biology, University of York, UK, York, YO10 5DD, UK
²Department of Infection, Hull University Teaching Hospitals NHS Trust, Hull, HU3 2JZ, UK
³Experimental Medicine and Biomedicine, Hull York Medical School, University of York, York, YO10 5DD, UK

Abstract
Background: Antimicrobial resistance (AMR) is a global concern and better understanding of the gut microbiome, a known ‘amplifier’ of AMR, may allow future clinicians to tailor therapy to minimise this risk and offer a personalised medicine approach. To examine the gut microbiome, patients are required to provide faecal samples; more convenient and cheaper solutions need to be found.

Methods: As part of a pilot study looking at how routes of administration affect the gut microbiome in NHS patients undergoing routine clinical management for infections, we hypothesised that effects on the gut microbiome varied with the route and metabolism of antibiotic used, and these changes may be reflected in breath metabolites. We present a case report of a patient with an unusual clinical history, alongside breath metabolite and gut microbiome data taken before, during and after antibiotic therapy over a period of one year.

Results: We noted a shift in the dominant Bacteroides strain in the patient’s gut microbiome between pre- and post-therapy samples, along with an alteration in the composition of breath metabolites.

Conclusions: This study provides a framework for similar future work and highlights the need for further research on the relationships between changes in microbial gut communities and antimicrobial exposure, patient clinical status, and the metabolites of human breath.
Keywords
microbiome, volatile organic compounds, antibiotics, breath metabolites, resistance, antimicrobials

Corresponding author: Farah Shahi (fshahi@doctors.org.uk)

Author roles: Shahi F: Data Curation, Funding Acquisition, Investigation, Project Administration, Resources, Validation, Writing – Original Draft Preparation, Writing – Review & Editing; Forrester S: Data Curation, Formal Analysis, Resources, Software, Validation, Visualization, Writing – Original Draft Preparation, Writing – Review & Editing; Redeker K: Conceptualization, Data Curation, Formal Analysis, Funding Acquisition, Investigation, Methodology, Resources, Supervision, Validation, Writing – Original Draft Preparation, Writing – Review & Editing; Chong JPJ: Conceptualization, Funding Acquisition, Methodology, Resources, Supervision, Writing – Review & Editing; Barlow G: Conceptualization, Data Curation, Funding Acquisition, Investigation, Methodology, Resources, Supervision, Visualization, Writing – Review & Editing

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Amendments from Version 1

The following updates have been made: i) clarified the rationale for the pilot study; ii) moved the patient's microbiology results table from supplementary files to main text; iii) clarified that no antimicrobials were received by the participant in the 12 months prior to starting the study; iv) clarified the methodology and assembly/analysis for the nanopore sequencing and sample pooling; v) added a reference to the legend for Figure 2A; vi) clarified that the patient did not use probiotics in the introduction; vii) removed an unnecessary sentence regarding the patient's enterotype to aid clarity of the text; viii) clarified that microbial abundance is relative rather than absolute; ix) added a figure to Figure 1 to provide a stack plot of abundances (now 1B); x) included data on resistance genes; xi) added an acknowledgement.

Any further responses from the reviewers can be found at the end of the article

Introduction

Scientific developments have enabled the possibility of personalised medicine at this time more than any other. In an age where antimicrobial resistance (AMR) is a global concern, insights into the gut microbiome, a known ‘amplifier’ of AMR, may in the future aid clinicians in tailoring therapy for their patients to reduce this and other health risks. However, for patients, providing faecal samples can be an uncomfortable experience, and the cost of sequencing individual patients’ gut microbiomes, both in terms of time and resource, still remains prohibitive. Increasingly, scientists and clinicians are searching for more convenient options for patients, including the use of rectal swabs.

Breath analysis, or breathomics, is an exciting but challenging field of science and medicine. Agreeable to the patient due to its non-invasive nature, it has great potential for individualized point-of-care diagnostics and screening. Volatilomics, the study of Volatile Organic Compounds present in breath, is a systemic analysis, combining metabolic signals from the entire body through the blood-lung interface as well as the gut microbiota.

In a pilot study looking at how routes of administration affect the gut microbiome in NHS patients undergoing routine clinical management for infections, we hypothesised that: i) the effects on the gut microbiome would vary with the spectrum of antibiotic used, the route of delivery, and the metabolism of the drug and ii) related effects may be identifiable in breath metabolites. Seven participants were recruited to assess the feasibility and suitability of such a study for NHS patients. Only patients who had not received antibiotics in the last 12 months were recruited.

Here we present a case report of one of the pilot study participants: a patient with an unusual clinical history, alongside breath metabolite and gut microbiome data taken before, during and after antibiotic therapy over a period of one year. There are few such case reports in the literature to date.

Case report

Patient A, a 57-year-old Caucasian male, presented to clinicians at Hull University Teaching Hospitals NHS Trust with an 11-year history of biliary system infection. In 2006 he had undergone a laparoscopic cholecystectomy for cholecystitis; this was complicated by a ‘dropped’ gallstone (a stone that remains in the abdominal cavity after surgery), subsequent recurrent hepatic infections and abdominal wall fistulation. Multiple surgical and radiological attempts at stone retrieval and drainage were eventually successful in 2016. However, six months later he continued to suffer recurrent infections and a CT scan in July 2017 revealed a persistent perihepatic abscess and hepatic-cutaneous fistula devoid of visible calculi. Consequently, he was referred to the Outpatient Parenteral Antimicrobial Therapy (OPAT) service for consideration of prolonged antimicrobial therapy.

On examination of the patient, a fistula was evident in the right upper quadrant, which correlated with a sinus tract on his CT. Baseline physiological observations were normal. Blood tests showed a C-reactive protein (CRP) of 17 mg/L (normal range 0 to 8 mg/L) and total white blood cell count of 7.9 × 10^9/L (normal range 4.0 to 11.0 × 10^9/L) but were otherwise unremarkable. Based on the patient’s relevant clinical microbiology results (Table 1), applicability to OPAT usage and longer-term tolerability, a decision was made to start intravenous (IV) Ertapenem 1g once daily. This was continued for 38 days, followed by oral Ciprofloxacin 750mg every 12 hours for 23 weeks.

Prior to starting therapy, Patient A was enrolled into the aforementioned pilot study following informed, written consent. Ethical approval for the study was granted by the HRA, Leicester REC 16/EM/0345 and the University of York Biology Ethics Committee. In total, seven faecal and seven breath samples were provided by Patient A between September 2017 and August 2018. The first samples were taken prior to starting antibiotic therapy; further sampling times are listed in Table 2. Patient A’s diet included a daily cod liver oil capsule and regular diets but he did not take probiotic drinks/supplements before, or for the duration of, this study.

Results

Sequencing and assembly of faecal specimens

Microbial DNA extracted from stool samples was subjected to long- and short-read DNA sequencing. The final polished assembly comprised of 9,554 contigs, N50 56,726bp and a total length of 159,132,373bp (see 6 for read statistics). 156,292 genes were identified using Prodigal version 2.6.3, within the anv’i’o pipeline. 71.7% of contigs were annotated, with 1,295 unique taxon names. CONCOCT annotation resulted in 81 bins from the data, and the manual tree-based approach resulted in 13 bins. The manually curated clusters were favoured over those generated by CONCOCT due to 48 of the 81 bins not being resolved into a genus annotation. This is because annotation is based on the most abundant annotation within the bin, and so the most abundant genera may be unknown, and account for less than 30% of the contigs within the bin. CONCOCT bin annotations are provided. All manually curated bins were annotated, with 8 of 13 being annotated as Bacteroides. The annotation and semi-quantitative abundance relative to the stage of treatment are provided.
Table 1. Relevant clinical microbiology results for Patient A.

<table>
<thead>
<tr>
<th>Date</th>
<th>Sample Type (site)</th>
<th>Organism</th>
<th>Sensitivities</th>
</tr>
</thead>
<tbody>
<tr>
<td>11/07/2016</td>
<td>Wound swab (drain site)</td>
<td>Moderate growth of MSSA</td>
<td>Resistant: Penicillin</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sensitive: Flucloxacin, Gentamicin, Ciprofloxacin, Clarithromycin, Clindamycin, Linezolid, Doxycycline, Tigecycline, Fusidic acid, Chloramphenicol, Rifampicin, Trimethoprim</td>
</tr>
<tr>
<td>05/10/2016</td>
<td>Wound swab (abdomen)</td>
<td>Moderate growth of MSSA</td>
<td>Resistant: Penicillin</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sensitive: Flucloxacin, Gentamicin, Ciprofloxacin, Clarithromycin, Clindamycin, Linezolid, Doxycycline, Fusidic acid, Rifampicin, Co-trimoxazole</td>
</tr>
<tr>
<td>12/10/2016</td>
<td>Wound site (drain)</td>
<td>Scanty growth of coliform bacilli</td>
<td>Not tested</td>
</tr>
<tr>
<td>08/09/2017</td>
<td>Wound swab (abdomen, pus from hepatic-cutaneous sinus tract from liver abscess- 'deep' sample taken in the OPAT service)</td>
<td>Scanty growth of E. coli</td>
<td>Resistant: Amoxicillin, Co-amoxiclav</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sensitive: Piperacillin/Tazobactam, Cefotaxime, Aztreonam, Ertapenem, Meropenem, Amikacin, Gentamicin, Tobramycin, Ciprofloxacin, Tigecycline, Ceftazidime, Cotrimoxazole</td>
</tr>
</tbody>
</table>

Table 2. Sampling Schedule for Patient A relative to antibiotic treatments. IV = intravenous, PO = per os (by mouth), OD= omni die (every day - implied to be once per day), BD = bis die (twice a day).

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>Antibiotic</th>
<th>Timing of Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-antibiotics</td>
<td>N/A</td>
<td>Baseline samples</td>
</tr>
<tr>
<td>Sample 1</td>
<td>Ertapenem 1g OD IV</td>
<td>8 days into therapy</td>
</tr>
<tr>
<td>Sample 2</td>
<td>Ertapenem 1g OD IV</td>
<td>14 days into therapy</td>
</tr>
<tr>
<td>Sample 3</td>
<td>Ertapenem 1g OD IV</td>
<td>21 days into therapy</td>
</tr>
<tr>
<td>Sample 4</td>
<td>Ciprofloxacin 750mg BD PO</td>
<td>5 days into oral therapy</td>
</tr>
<tr>
<td>Sample 5</td>
<td>Ciprofloxacin 750mg BD PO</td>
<td>78 days into oral therapy</td>
</tr>
<tr>
<td>Post-antibiotics</td>
<td>N/A</td>
<td>122 days without therapy</td>
</tr>
</tbody>
</table>

Neither of the previous strategies were able to resolve the data into single genome bins, and both resulted in bins that were either overfilled or highly contaminated with contigs from other genomes, preventing resolution of the data to species level. A custom analytical pipeline written in-house (CLUSTard, Cansdale et al., in prep) was able to identify 19 metagenome assembled genomes (MAGs) that we assumed represented different species as shown in Figure 1A/B, which constituted over 97% of the patient’s microbiome. The abundance of the additional annotated species is summarised within the “Other” bin within Figure 1A/B. These are cumulative abundances taken from all clusters containing the same species annotation, and so some of the species are explained through multiple bins. However, different MAGs with the same species identity have a similar trajectory for the majority of bases within that bin. Contigs that have the same annotation as any of the top 19 MAGs but are not included within Figure 1A/B are minimal, as shown in Figure 1C. These contigs had both a different trajectory and only represented a small percentage of the total bases of that bin. The full results showing the abundance per bin and per contig for the entire assembly are available as extended data.

The two most abundant species throughout the time course are Bacteroides coprophilus and Bacteroides dorei (Figure 1A). Bacteroides coprophilus is high in abundance during pre-treatment and the course of IV Ertapenem, whereas Bacteroides dorei is low in abundance during IV Ertapenem, but becomes highly abundant during oral Ciprofloxacin therapy, and following treatment.
Comparison of the MAGs used in the assembly for the species in Figure 1A/B compared to the National Center for Biotechnology Information (NCBI) reference genomes for the same species suggests that we do not have complete genomes for all of the top 19 most abundant species. The size of the assembled genomes relative to the NCBI reference for Oscillobacter (160%), Bacteroides dorei (124%), and Faecalibacterium prausnitzii (160%) suggest mis-assemblies that make the genomes larger than the expected size. However, the size of the assembled Bacteroides coprophilus genome is double that expected, which suggests that we could have two strains of Bacteroides coprophilus that have been assembled. There are two high quality MAGs that are singletons (one contig assembled genomes) that are annotated as Bacteroides coprophilus, Cluster Singleton2 and Cluster Singleton4, which are genome sized. However, they have the same trajectory suggesting that they respond to antibiotic treatment in the same way.

Three species, B. helcogenes, B. cellulosilyticus and Lactobacillus reuteri, are significantly increased in the post-treatment
samples (Figure 1A/B). Faecalibacterium is strongly suppressed during antibiotic treatment but recovers after treatment stops. Levels of Prevotella dentalis were higher prior to treatment with IV Ertapenem, and continued to reduce during oral Ciprofloxacin, remaining at low levels post-treatment. Parabacteroides is also more abundant during IV Ertapenem use and post-treatment but is suppressed during oral Ciprofloxacin treatment. There are three Alistipes species: two species are suppressed during oral Ciprofloxacin treatment, and displaced by Alistipes shahii, which is the most abundant during this phase. Following treatment there is a more equal mix of all three species, with Alistipes finegoldii and Alistipes sp. Marseille not recovering to pre-treatment levels but Alistipes shahii remaining higher (Figure 1A/B). Ruminococcus champanellensis significantly increased during treatment with IV Ertapenem but reduced during oral Ciprofloxacin therapy, remaining low post-treatment. Finally, three species were noted to become particularly abundant in the final sample taken on antibiotics: Lactobacillus reuteri, Akkermansia muciniphila and Bacteroides ovatus.

Resistance genes were identified in both the B. coprophilus and B. dorei reference genomes, and the most abundant MAGs in these two species. As suggested by the assembly size being double the size of the NCBI reference genome, the presence of two strains of B. coprophilus is also suggested by the identification of different resistance genes present within different B. coprophilus MAGs. 75% of all B. coprophilus annotated bases were split between four MAGs, Cluster_contig0015 (15%), Cluster_contig0024 (19%), Cluster_singleton2 (26%) and Cluster_singleton4 (14%). The remaining bases are split across 14 lower abundance and quality clusters. 67% of all B. dorei annotated bases are within the MAG Cluster_contig0017. The remaining bases are split across > 20 other lower abundance and quality clusters. Cluster_contig0017 was used as the representative MAG for B. dorei for gene resistance annotation and the four aforementioned MAGs were used for B. coprophilus.

In B. coprophilus (more abundant during pre-treatment and IV Ertapenem) MAGs Cluster_singleton4 and Cluster_contig0015 have lost/have absent all three resistance genes adeF, tetQ and ErmF present in the NCBI reference. However B. coprophilus MAGs Cluster_singleton2 and Cluster_contig0024 have lost tetQ and ErmF genes, but have maintained an adeF resistance gene. Reaffirming the suggestion that there are two strains of B. coprophilus strains present. In B. dorei (more abundant during oral Ciprofloxacin and post-treatment), we observed an Erm 23S ribosomal RNA methyltransferase and the loss of a tetracycline antibiotic resistance gene.

Blastp searches were performed to identify carbapenem resistance, the presence of gyrA and gyrB plasmids which have previously been identified within the Bacteroides fragilis group and shown to contribute to antibiotic resistance^8, qnr and parC resistance genes. We found no hits that were significant (had an e-value of equal or less than 0.05) in the QNR resistance genes for either B. dorei or B. coprophilus. We did identify carbapenem resistance in B. dorei, with two hits, one 75% of the length of the resistance gene in Cluster_contig0017 on contig3934, with 70.9% percent identity and an e-value of 1.00e-143. The other matched 51% of the resistance gene length in the same cluster, but on contig0116 with an e-value of 6.00e-25, but only a percentage identity of 38.64%. B. coprophilus had one valid GyrA match, with a 95.4% coverage of the resistance gene length, 48% percentage identity and e-value of 0. However all QNR, GyrB and ParC hits were insignificant, with a low percentage of the resistance gene matching to the contigs, and a high e-value. (Supplementary file added to OSF repository).

Three human gut enterotypes have been identified based on the relative abundance of Bacteroides, Prevotella and Ruminococcus in the gut^11. This patient identifies as a Type 1 enterotype in both pre and post samples due to the high abundance of Bacteroides. This is shown consistently between different clustering analyses. Despite no change in enterotype, the composition of the pre- and post-treatment microbiota is different. The Bacteroides genus remains the most abundant, however Bacteroides dorei displaces Bacteroides coprophilus and becomes the most abundant species.

Breath metabolites are atypical and vary in this patient

The breath samples from the patient showed substantial modification from ambient air masses, and for some volatiles the patient exhibited significantly different breath outcomes relative to other sampled and reported individuals. Most prominently different in the patient’s breath were the methyl halides and thiol, including methyl chloride, methyl bromide, methyl iodide and methane thiol (CH3X, where X = Cl, Br, I or SH and shown as MeCl, MeBr, MeI and MeSH, Figure 2).

In urban air masses methyl halides are found at concentrations of approximately 750, 15 and 5 pptv, respectively12. Humans metabolise these compounds (both in terms of uptake and production) leading to average exhaled breath containing 3500 ± 2400 parts-per-trillion-by-volume (pptv) MeCl, 6.5 ± 11 pptv MeBr and 1.1 ± 1.8 pptv MeI. In contrast, the patient’s average breath profile contained 13,000 ± 2000 pptv MeCl, 73 ± 27 pptv MeBr and 3.3 ± 0.8 pptv MeI. These breath concentrations were significantly different than the average male population for both MeCl and MeBr and consistently more concentrated for all methyl halides. Both methyl chloride and methyl bromide showed consistency from prior to antibiotics and throughout IV Ertapenem therapy (13400 ± 1200 pptv MeCl, 66 ± 9 pptv MeBr) but were significantly reduced during oral Ciprofloxacin (8900 pptv MeCl, 49 pptv MeBr at second sampling, 78 days into oral therapy). Methyl iodide concentrations in the patient’s breath did not notably change over time during antibiotic treatment.

Atmospheric data within urban environments is scarce but our previous breath results suggest that methanethiol (MeSH) is common in human breath at concentrations of 1.7 ± 8.6 pptv. Patient A's breath was elevated in MeSH relative to healthy human breath, 2.7 ± 2.8 pptv, but not significantly. MeSH results should be viewed with some caution however, due to challenges that derive from storage concerns^13. Despite these
considerations, MeCl, MeBr and MeSH were all significantly correlated ($p < 0.001$) with each other (MeCl v.s. MeSH - $r^2 = 0.49$; MeCl v.s. MeBr - $r^2 = 0.49$; MeBr v.s. MeSH - $r^2 = 0.67$). MeI is also significantly correlated ($p<0.001$) with MeCl, MeBr and MeSH, however correlations are limited ($r^2 \sim 0.15$) between MeI, MeCl and MeSH. Only in the correlation between MeI and MeBr does the correlation coefficient approach 0.5.

Other human-metabolised halogenated compounds showed interesting outcomes in Patient A, including dichloromethane (CH$_2$Cl$_2$) and chloroform (CHCl$_3$). Ambient urban air contains approximately 5600 and 1300 pptv, respectively, while average healthy human breath is generally less concentrated, although highly variable, at $1000 \pm 1500$ pptv for dichloromethane and $230 \pm 660$ pptv for chloroform. The patient exhibited

Figure 2. Plot showing the concentration of metabolites identified from breath of patient A. (A) shows a boxplot distribution of metabolite abundance of non-patient samples ($n=61$ except for MeCl where $n=68$), compared to the distribution of Patient A’s abundance over the time course. (B) shows the logged 10 abundance of each metabolite compared to the averaged value (unpublished, personal communication, Redecker et al. 2021) shown in blue, and the ambient value for the metabolite, shown in green. CH$_2$Cl$_2$ = Dichloromethane, CHCl$_3$ = Chloroform, DMS = Dimethyl sulfide, MeBr = Methyl bromide, MeCl = Methyl Chloride, MeI = methyl iodide, MeSH = Methanoethiol
uniform, and depleted, concentrations for dichloromethane throughout the study, 20 ± 6 pptv dichloromethane, while chloroform was uniformly low but highly variable, at 95 ± 71 pptv. Concentrations of chloroform showed a strong time, and possibly therefore antibiotic dependency, with starting concentrations near 105 pptv that fell throughout IV Ertapenem to a level below the detection limit (<5 pptv) by the start of oral Ciprofloxacin, returning to approximately the same concentrations as the earliest sample post-treatment (~105 pptv).

The sulfur-containing compound, dimethyl sulfide (DMS) was present in the patient’s breath, ranging from 900 to 16000 during the various stages of treatment. While the average breath concentration of DMS (4700 ± 4900 pptv) in patient A was similar to human breath averages (2300 ± 2300 pptv DMS), this was primarily due to the high variability of this data across individuals and populations. DMS concentrations appeared to be treatment dependent, with a substantial increase in DMS during IV Ertapenem therapy (from 2900 to 16000 pptv) before falling during oral Ciprofloxacin to near average healthy human concentrations (~2800 pptv).

Isoprene, the most abundant human metabolite in breath, was similar between the patient (125 ± 23 parts-per-billion-by-volume; ppbv) and the healthy human population (185 ± 205 ppbv). Baseline concentrations of isoprene can vary substantially between individuals (from 80 to 300 ppbv within our healthy controls), but within the established baseline it tends not to vary dramatically, which is what we observed in the patient post-antibiotic therapy when isoprene levels rose to 178 ppbv.

Clinical follow-up
Patient A symptomatically improved during treatment, which resulted in clinical and radiographical resolution of the hepatic-cutaneous fistula. A repeat CT scan seven weeks into antimicrobial therapy suggested the patient’s infection had not fully resolved, but a further CT scan at 16 weeks of therapy showed no evidence of ongoing infective changes and resolution of the previous peri-hepatic abscess (see Figure 3). He was discharged from clinical follow up in March 2019 with a CRP of 3.1 mg/L and has not suffered recrudescence since.

Discussion
Understanding and mitigating the consequences of antimicrobial exposure on the human gastrointestinal tract microbiome at an individual level is likely to become increasingly important in the era of personalised medicine, given our increasing understanding of the associations between the nature of an individual’s gastro-intestinal (GIT) microbiome and the risk of a disease state. The GIT microbiome is also recognised as an ‘amplifier’ of antimicrobial resistance and changes in an individual’s microbiome may contribute to this. In this pilot study, we sought to describe the changes occurring in patients’ gut microbiome during IV and oral antibiotic therapy, and assess whether these changes were reflected in breath metabolites.

In Patient A, B. coprophilus, a bacterium that has not been associated with human infection, was in relative abundance throughout IV Ertapenem exposure (suggesting the presence of a resistance mechanism), suppressed during oral ciprofloxacin, and abundant again post-therapy. Low GIT exposure is unlikely to be the cause of abundance during Ertapenem therapy as 10% of a 1g dose is excreted in human faeces (Ertapenem 1g Powder for Concentrate for Solution for Infusion, Electronic Medicines Compendium, 2020). Meropenem exposure, however, has previously been associated with the emergence of carbapenem resistance in Bacteroides fragilis, which is commonly associated with the cfiA metallo-beta-lactamase gene14. Fluoroquinolone susceptibility in B. fragilis in the presence of carbapenem resistance has been described in bloodstream infection15, although resistance specifically to Ciprofloxacin in B. fragilis strains is generally expected to be high16. However, in Patient A, the abundance of the adeF gene throughout, which has been associated with efflux-mediated resistance to fluoroquinolones and tetracyclines in clinical practice, suggests that this gene was not expressed by B. coprophilus during Ciprofloxacin exposure.

In contrast, B. dorei was more abundant during oral Ciprofloxacin exposure and post-therapy suggesting susceptibility to Ertapenem (despite the presence of a carbapenem resistance gene on blastp search) and resistance to Ciprofloxacin, in keeping with potential expression of the adeF gene or another

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**Figure 3.** CT imaging before (A) and at 16 weeks (B) of antibiotic therapy (labels added with Mac’s Preview App v11.0).
mechanism. Furthermore, the loss of ermF and tetQ from each of the most abundant Bacteroides spp., along with the subsequent gain of ermG in the prevailing most abundant species, B. dorei, highlights the impact of antimicrobial exposure on AMR within the gut microbiome, which may not always be intuitively related to the class of antimicrobial a patient is exposed to\textsuperscript{17,18}. The contrasting states of B. coprophilus and B. dorei with their respective genomic data may represent the important differences between phenotypic and genotypic expression by bacteria although without parallel transcriptomic data, it is not possible to comment further.

Faecalibacterium prausnitzii has been proposed as a biomarker of gut health. An important producer of butyrate, studies have demonstrated an association between gut disease and low levels of Faecalibacterium in patients with inflammatory bowel disease\textsuperscript{19,20}. Patient A's levels of Faecalibacterium remained suppressed during therapy, almost completely recovering to pre-therapy levels on discontinuation. A recent study in mice treated with Faecalibacterium supplementation and a prebiotic showed clearance of bacterial burden of Clostridium difficile and potential reduction in toxin production compared with untreated mice\textsuperscript{21}. Interestingly, Patient A's levels of Clostridium were low before and after antibiotic exposure, but were high at times during exposure to both antibiotics. The patient did not develop diarrhoea while on antibiotic therapy.

Bacteroides ovatus, Lactobacillus reuteri and Akkermansia muciniphila all showed marked increases at time point 5 of antimicrobial therapy (after 78 days of oral Ciprofloxacin). Akkermansia has been labelled a “sentinel of gut health” due to its apparent effect of decreasing gut permeability and its low levels in patients with diabetes and obesity\textsuperscript{22-24}. Similarly, studies of Lactobacillus reuteri have proposed possible anti-inflammatory effects\textsuperscript{25-28}. Bacteroides ovatus has been shown to reduce to LPS-mediated inflammation in animal studies and its use as a probiotic continues to be investigated\textsuperscript{29,30}. Concurrently there were appreciable decreases in the levels of methylated compounds in Patient A's breath sample. These findings may be associated with the recovery in Patient A's clinical status. Further research is warranted to investigate whether markers such as these in patients' breath and/or faecal samples have the potential, in real-time, to indicate whether a review of the need for ongoing antimicrobial therapy is required.

The use of breath samples in diagnosis and management is attractive because it is minimally invasive and acceptable to patients\textsuperscript{39}. Patient A demonstrated distinct differences in his breath samples compared to the average male in previous studies (unpublished; personal communication, Redeker et al. 2021). This highlights the individuality of each person’s results\textsuperscript{10} and the importance of large sample sizes of the population in understanding what may or may not be statistically generalisable in personalised medicine. However, the broad pattern of changes may in fact be more universal, as demonstrated through “electronic noses”: the use of breath metabolites in the diagnosis of human disease to assess patterns of volatile organic compounds (VOCs) associated with specific illnesses\textsuperscript{30,31}.

There is increasing evidence that exhaled breath and the gut microbiome are strongly correlated\textsuperscript{12-14}. The method of breath analysis we used for Patient A is more amenable to a personalised medicine approach but pattern changes over time may carry more significance than individual VOCs obtained at single timepoints during patient care. The inverse association of chloroform levels and antibiotic therapy may be of interest; if such relationships can be robustly established, it may be possible to monitor aspects of drug therapy at the bedside via the breath, for example, rather than current invasive techniques.

Most VOCs in Patient A's breath did not return to starting-point values on completion of therapy, suggesting that microbial/human metabolisms within the patient had been substantially modified during illness recovery and/or antimicrobial exposure. Combined with the shift demonstrated in the dominant Bacteroides strain within the bowel, as well as significant differences in the microbial community pre- and post-therapy, this suggests potentially measurable modification of microbial communities in patients receiving prolonged antimicrobial therapy. The relevance of these specific community and metabolism changes to health in the longer-term is yet to be understood, particularly as the shifts may represent a return to a pre- or post-illness healthier state or an adjustment to physiological niches held by certain strains that bear no impact on overall function. After all, Patient A clinically improved and became both subjectively and objectively healthier over time, but it is still possible that persisting changes in Patient A’s microbial community as a result of prolonged antimicrobial exposure may have longer-term health implications.

The complex interplay between different bowel flora \textit{in vivo} in humans remains an important driver for ongoing research of the gut microbiome. Limitations of this pilot project include the likelihood that breath metabolites may be affected by microbiomes or human metabolisms other than those within the gut; for example, the oral and lung microbial communities. However, this study provides a framework for similar future work and highlights the need for further research into the relationships between changes in microbial gut communities during antimicrobial exposure, AMR colonisation and subsequent infection risk, monitoring of patient clinical status, and the metabolites of human breath.

Methods

Sample collection

Faecal samples were aliquoted on the day of collection and stored at -80°C. Breath samples were collected in Tedlar bags, protected from light, and transferred into electropolished, stainless steel canisters (LabCommerce, Torrence, CA) via an Ascarite\textsuperscript{34} trap on the day of collection\textsuperscript{12}.

DNA extraction, library preparation and sequencing

Microbial DNA was extracted from each faecal sample within 1–18 days post-collection, using a QIAmp PowerFecal DNA Kit. For quantitation purposes, uniquely barcoded Illumina sequencing libraries were prepared from all samples and timepoints using the NEBNext Ultra II FS DNA library.
prep kit for Illumina (New England Biolabs) according to manufacturer’s guidelines. For each sample 150 ng starting DNA was subjected to a 13 minute fragmentation time, and 4 cycles of PCR amplification using NEBNext multiplex oligos for Illumina (unique dual index primers; NEB). Sample libraries were pooled at equimolar ratios and subjected to paired end 150 bp sequencing on an Illumina HiSeq 3000 (University of Leeds Next Generation Sequencing Facility). Illumina reads were adapter trimmed with cutadapt\(^3\) 1.18 with option -a AGATCGGAAGAG to match the Illumina universal adapter.

Long fragment DNA sequencing was performed using an Oxford Nanopore Technologies’ PromethION with all timepoints pooled to generate a long read assembly, subjected to bead cleaning using an equal volume of AMPure XP beads (Beckmann Coulter). 1.5 ug total DNA from each patient was used in library preparation using the ONT ligation sequencing kit (SQK-LSK109) with native barcode expansion pack (EXP-NB103). The DNA was subjected to DNA repair/ A-tailing using the NEBNext\(^8\) Ultra\(^TM\) II End Repair/DA-Tailing Module, with additional NEBNext FFPE repair enzyme (NEB), with sequential incubations for 30 minutes at 20 °C and then 65 °C. Following clean up with 0.9 volumes AMPure XP beads, a unique barcode adapter was ligated to the prepared DNA from each patient using NEB Blunt/TA ligase mastermix. An additional AMPure clean-up was performed before pooling approximately equimolar quantities of barcoded sample from each patient. The sequencing adapter from EXP-NBB103 was then ligated to the fragment ends using the NEBNext Quick ligation module. A final AMPure XP bead clean up, including two washes with ONT long fragment wash buffer (LFB) was performed prior to library quantitation using the Qubit fluorimeter (Thermo Fisher). The library was loaded onto a FLO-PR0002 R9.4.1 flow cell, following manufacturer’s guidelines, and sequenced for 56 hours.

Sequence assembly
Sequences from all timepoint samples were pooled for assembly, using long reads for the original assembly, and short reads for polishing the assembly. Sequencing data was basecalled with Guppy 1.8.3, and demultiplexed with Porechop 0.2.3. Nanopore reads were assembled with canu\(^6\) 1.8 using parameters genomeSize=100m, corMinCoverage=0, corOutCoverage=all, corMhapSensitivity=high. The canu assembly was polished with the nanopore reads using medaka 0.7.1 and with the Illumina reads using Pilon\(^7\) 1.23, running Pilon three times, aligning reads to the assembly with bwa\(^8\) mem 0.7.17 and indexing and sorting read alignments with SAMtools\(^9\) 1.9. All tools were run with default options unless otherwise stated.

Sequence analysis
The assembly was filtered to remove contigs less than 1000bp, (see extended data\(^3\) for contig length distribution) before being annotated and clustered using the anvi’o metagenomic workflow using the centrifuge nt database\(^7\). The short reads were used for clustering and taxonomic assignment and abundance was calculated based on the depth of coverage of the short reads mapped to the assembly. Clustering was performed using both an automated method using CONCOCT and manually using hierarchical clustering of contigs\(^3\). Taxonomy was generated based on the contigs, and the abundance of these taxon through the coverage of the short reads mapped to the assembly using bwa mem 0.7.17\(^8\).

The metagenomic clustering pipeline CLUSTard (public publication pending) was used to generate clusters of contigs based on mapped short read abundance, using a kraken2 database. Pearson coefficient correlation values were varied to determine percentage similarity in trajectory between contigs within a bin over time. Values between 0.997 and 0.8 were used to assess clustering, with 0.997 used in further analyses.

Contigs from both highly abundant clusters in Bacteroides coprophilus (Cluster Singleton2, Cluster Singleton4, Cluster contig0024 and Cluster contig0015) and the most abundant cluster in Bacteroides dorei (Cluster contig0017), were annotated using the CARD: RGI database\(^3\) to identify antimicrobial resistance genes that could be responsible for their trajectories throughout treatment. The NCBI reference genomes for Bacteroides salanitronis (CP002530.1) and Bacteroides dorei (CP008741.1) were used to identify expected resistance genes in the assembly and identify subsequent losses or gains in the contigs within these MAGs. Bacteroides salanitronis was used as a reference because it is closely related, and Bacteroides coprophilus is not present currently in this database. These MAGs were mapped to the NCBI references of both Bacteroides Coprophilus and Bacteroides Salanitronis using default options in minimap2\(^2\). There was no significant difference in the percentage of sequence mapping to these references, which we were satisfied meant that the genomes were similar enough that mapping to Bacteroides Salanitronis using the CARD RGI database would not impact on the resistance genes identified.

Seqkit version v0.11.0\(^3\) was used to convert contig nucleotide sequences to amino acid sequences in order to perform blastp searches with blast+ 2.2.31+ against additional resistance genes. The contigs from these highly abundant Bacteroides coprophilus and Bacteroides dorei were compared against gyrA, gyrB and parC sequences from biocyc accessions EG10423 and EG10424 and EG10686 respectively. Carbapenem resistance was identified using accession EEF178324.1 from NCBI as previously described by Goto et al.\(^4\). We also searched for QNR resistance, by using the QNRA alleles QNRA1-6 which have been identified in the Bacteroides fragilis group\(^4,4\). Hits were filtered so that only a e-value of less than 0.05 was used. The percentage identity and length of resistance gene covered were also used to determine significance.

Breath volatile metabolite analysis
Tedlar bags provide ease of sampling for patients but are unsuitable for long term storage of volatiles. To ensure stable long-term storage we transferred the breath sample from the
Tedlar bag to high-quality stainless steel canisters (electropolished prior to single welding) on the same day. Prior to sample transfer from the Tedlar bags the canisters were evacuated (<0.1mbar pressure). To ensure stability and sensitivity of analysis each sample passed through an Ascarite trap, driven by the pressure differential between the Tedlar bag and canister. The volatiles within this study have been shown previously\(^6\) to be stable in these canisters for over two weeks.

Canister-bound breath samples were concentrated onto a liquid nitrogen condensation trap through pressure differential, driven by a two-stage rotary pump, then run on a Gas Chromatograph-Mass Spectrometer (GC-MS) consisting of a Restek\textsuperscript{\textregistered} Porabond Q column (30m, 0.32mm ID, 0.5-\mu m thickness) within a HP 5972 MSD, running in selective ion mode, within six days of the date of collection\(^6\). We used CFC-11 as an internal standard since it is ubiquitous and constant in the atmosphere and there are no known metabolisms that produce or consume it within the human body.

Concentration calibrations were performed using purchased, low-concentration (ppbv) standards (BOC Specialty Gases; CFC-11, CFC-12, CFC-113, methyl halides, DMS) directly injected into the condensation system. Within our controlled temperature and volume trap, injection of a limited pressure of high concentration gas (the standards) mimics the same total amount of a greater pressure injection of a low concentration sample (breath samples). Accuracy of these standards for breath sample concentrations were ±10%. Alternatively, serial dilutions of headspace volatiles from neat standards were used (as in isoprene, methane thiol). These standards were less reliable than the prepared standards because of the small scale on which the system is operated as well as differences in reported vapor pressures for each compound. We calculated accuracy for quantification of samples based on these standards to be ±20%.

Data availability
Underlying data
Whole metagenome sequencing, using long and short reads for the assembly of a gut microbiome. Accession number PRJEB44880; https://identifiers.org/ena.emb|PRJEB44880

Extended data
Open Science Framework: The effect of intravenous and oral antibiotics on the gut microbiome and breath volatile organic compounds over one year: a case report. 10.17605/OSF.IO/TJW9X\(^6\) Raw data is archived under ‘Files.’

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Consent
Written informed consent for publication of their clinical details and images was obtained from the patient.

Author contributions
FS & SF are joint first authors. FS managed the study, sample processing and co-wrote the manuscript. SF completed the genomic data analysis, breath data synthesis and co-wrote the manuscript. KR completed the breath analysis and co-wrote the manuscript. JPJC provided oversight of part of the project, obtained additional funding and revised the manuscript. GB designed the study, was Principal Investigator for the project and revised the manuscript.

Acknowledgments
We would like to thank Patient A for taking part in this study and agreeing to the publication of these results, and Drs Sally James, Peter Ashton and John Davey within the Genomics Laboratory of the Biosciences Technology Facility at the University of York for nanopore sequencing support.

References

Publisher: Wellcome Open Research


Thank you for your clarifications to my specific comments. However, I don't believe that you have responded to my primary query regarding the association between the metabolome and the microbiome. I have copied my original query related to this below:

The overall rationale for this study was, as I understand, to assess (in a pilot study) whether exhaled breath metabolites could be used as a proxy for gut microbial profiles. However, no analysis explores the association between gut taxa and the measured metabolites. This is surely a missed opportunity, and reduces the impact of this work.

Also, some reflection on the metabolic activity of the gut taxa identified, and their role in producing or modifying the metabolites measured would surely be important. Since whole metagenome sequencing was done it should be possible to do functional analysis to interrogate this question.

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Clinical microbiology, microbiome, tuberculosis

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.
This work investigated both gut microbiome and breath metabolites of the patient who received antibiotic therapy following surgical complication of laparoscopic cholecystectomy. Authors demonstrated the change of gut microbiome together with breath analysis along the treatment. This work emphasized that antibiotic agents caused dysbiosis in the gut which results in metabolic disturbance, represented by different components of the breath.

Some points are needed to be clarified to improve this piece of work:

- In Figure 1A, are these percentage shown with respect to the total abundance of each species or to the total bacterial species at each time point?

- Stack plot showing the abundance of bacterial species at each time point should be shown to illustrate the dynamic of the species.

- Authors mentioned Ciprofloxacin resistance and investigated genes that are associated with the resistance. However, common fluoroquinolone resistance genes, such as $qnrA$, $gyrA$, $gyrB$, and $parC$, have not been mentioned. Please describe whether these genes were identified in the $B.\, dorei$.

- Also, gene associated with carbapenem resistance was not mentioned in $B.\, coprophilus$.

**Is the background of the case's history and progression described in sufficient detail?**

Yes

**Are enough details provided of any physical examination and diagnostic tests, treatment given and outcomes?**

Yes

**Is sufficient discussion included of the importance of the findings and their relevance to future understanding of disease processes, diagnosis or treatment?**

Yes

**Is the case presented with sufficient detail to be useful for other practitioners?**

Yes

**Competing Interests:** No competing interests were disclosed.
Reviewer Expertise: Microbiology, Antibiotic resistance

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 31 May 2022

Farah Shahi, University of York, UK, York, UK

Many thanks for your comments and for your time. Please see our responses below and within the article.

Some points are needed to be clarify to improve this piece of work:
  ○ In Figure 1A, are these percentage shown with respect to the total abundance of each species or to the total bacterial species at each time point?
  ○ It is based on the relative abundance per timepoint, so Bacteroides coprophilus at timepoint 3 is 50% of all of the bacteria sampled at time point 3.
  ○ Stack plot showing the abundance of bacterial species at each time point should be shown to illustrate the dynamic of the species.
  ○ We have added a new Fig 1B to display this, thanks.
  ○ Authors mentioned Ciprofloxacin resistance and investigated genes that are associated with the resistance. However, common fluoroquinolone resistance genes, such as qnrA, gyrA, gyrB, and parC, have not been mentioned. Please describe whether these genes were identified in the B. dorei.
  ○ Also, gene associated with carbapenem resistance was not mentioned in B. coprophilus.

Thank you. We have added to the article based on your comments.

Competing Interests: None.

Reviewer Report 21 February 2022

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Mark P. Nicol
Division of Infection and Immunity, School of Biomedical Sciences, The University of Western Australia, Perth, WA, Australia

This is an interesting report – more longitudinal analyses such as this are needed. The overall rationale for this study was, as I understand, to assess (in a pilot study) whether exhaled breath
metabolites could be used as a proxy for gut microbial profiles. However, no analysis explores the association between gut taxa and the measured metabolites. This is surely a missed opportunity, and reduces the impact of this work.

Also, some reflection on the metabolic activity of the gut taxa identified, and their role in producing or modifying the metabolites measured would surely be important. Since whole metagenome sequencing was done it should be possible to do functional analysis to interrogate this question.

My second major concern relates to conclusions about absolute abundance made on the basis of relative abundance measures, see below.

Other comments:

INTRODUCTION

○ It probably not possible to test the proposed hypotheses, viz. “i) the effects on the gut microbiome would vary with the spectrum of antibiotic used, the route of delivery, and the metabolism of the drug, ii) related effects may be identifiable in breath metabolites” with a single case report. It may be better to reframe the rationale for this study, which was presumably, as a pilot study?

○ It is stated that “Based on the patient's relevant clinical microbiology results, applicability to OPAT usage and longer-term tolerability, a decision was made to start intravenous (IV) Ertapenem 1g once daily.” It would be useful to include these results in the text.

○ Had no antibiotics been given in the period leading up to the baseline sample collection? This would seem unlikely, given the lengthy delay involved.

METHODS

○ For the nanopore sequencing, it is not clear what is referred to here “as part of a larger experiment where for each patient separately, equal quantities of DNA were pooled from each timepoint”. There is only one patient. Were all samples from this patient sequenced on the same flow cell?

○ Were sequences from all samples pooled for assembly, clustering and taxonomic assignment, or was this done separately for each sample?

○ Were ambient air samples collected as controls at the same time of breath collection? This would seem to be key. In figure 2A, what samples are used for the ‘averaged’ values?

○ Were breath samples collected in a hospital environment? Are any of the metabolites with high concentrations in this study likely to be found at higher concentration in hospital air vs ambient urban air (which is the comparator used)?

○ Were breath samples collected in the same environment throughout the study? Could this account for some of the differences over time?

RESULTS

○ Did the patient take probiotic therapy, which may account for the increase in *Lactobacillus reuteri* following treatment?
Given the likely mis-assemblies for several key taxa, do the authors think that a single sample-level assembly might assist in teasing this out?

Do the different MAGs for *Bacteroides coprophilus* differ in relative abundance by timepoint? If so, how does this relate to the prevalence of the different resistance genes over time? What about for *Bacteroides dorei*?

I found this sentence very confusing: “Prevotella is also identified but in low abundance, and low but higher levels, compared to Prevotella, of Ruminococcus is also observed.”

**DISCUSSION**

The increase in *B. coprophilus* relative abundance during ertapenem therapy is interesting. Of course, this is relative, not absolute abundance, which is an important difference. It would be very useful to have some measure of total bacterial load over the course of therapy. Would 16S qPCR help here (even if an imperfect measure)?

Given the point above, it is probably not correct to make statements such as “*B. coprophilus, a bacterium that has not been associated with human infection, was abundant throughout IV Ertapenem exposure*” – the persistence (or increase) was in relative abundance and may in fact reflect a decline in absolute abundance if all taxa decrease in abundance, as one might expect during broad spectrum antibiotic therapy.

**Is the background of the case’s history and progression described in sufficient detail?**

Partly

**Are enough details provided of any physical examination and diagnostic tests, treatment given and outcomes?**

Partly

**Is sufficient discussion included of the importance of the findings and their relevance to future understanding of disease processes, diagnosis or treatment?**

No

**Is the case presented with sufficient detail to be useful for other practitioners?**

Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Clinical microbiology, microbiome, tuberculosis

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.
Farah Shahi, University of York, UK, York, UK

Thank you for your time and comments. Please see our responses below.

INTRODUCTION

- It probably not possible to test the proposed hypotheses, viz. “i) the effects on the gut microbiome would vary with the spectrum of antibiotic used, the route of delivery, and the metabolism of the drug, ii) related effects may be identifiable in breath metabolites” with a single case report. It may be better to reframe the rationale for this study, which was presumably, as a pilot study?

The article text has been updated to clarify this.

- It is stated that “Based on the patient’s relevant clinical microbiology results, applicability to OPAT usage and longer-term tolerability, a decision was made to start intravenous (IV) Ertapenem 1g once daily.” It would be useful to include these results in the text.

The article text has been updated to clarify this in the OPAT usage and longer-term tolerability sections.

- Had no antibiotics been given in the period leading up to the baseline sample collection? This would seem unlikely, given the lengthy delay involved.

The article text has been updated to clarify this in the baseline sample collection section.

Not within 12 months of starting the study. The article text has been updated to clarify this, thanks.

METHODS

- For the nanopore sequencing, it is not clear what is referred to here “as part of a larger experiment where for each patient separately, equal quantities of DNA were pooled from each timepoint”. There is only one patient. Were all samples from this patient sequenced on the same flow cell?

The article text has been updated to clarify this in the DNA extraction section.

- Were sequences from all samples pooled for assembly, clustering and taxonomic assignment, or was this done separately for each sample?

The article text has been updated to clarify this in the Sequence Assembly and Analysis sections.

- Were ambient air samples collected as controls at the same time of breath collection? This would seem to be key. In figure 2A, what samples are used for the ‘averaged’ values?

- Were breath samples collected in a hospital environment? Are any of the metabolites with high concentrations in this study likely to be found at higher concentration in hospital air vs ambient urban air (which is the comparator used)?

The legend for Figure 2A has been updated with a reference. For your information though, we did not sample for ambient air at the time of the patient sampling, as sampling for all participants was done at different locations, not always in a hospital environment and often in different parts of the hospital. Also, these compounds change in air rapidly, so we can expect a 30% variability from one moment to the next anyway (Low et al., 2003). We, therefore, have a moving background, even if there had been an ambient air sample taken at the same place each time.

Variability within the local ambient environment, combined with the fact that the
individual's blood and body were equilibrating with lung volatiles for ~ the previous 15 minutes or so means that an ambient air sample is not particularly informative in this case, and the sampling strategy was constrained by care.

However, it is worth noting that methyl chloride, methyl bromide, methyl iodide, and dimethyl sulfide are EXTREMELY high (often over an order of magnitude more than average male concentrations), and not in a way that is matched through any ambient background, internal or external that have been reported. Where the patient is low (isoprene), this is known to be human metabolism (~1,000x ambient background anywhere), so this decrease is notable.

Specifically, there are no published records of ambient concentrations of methyl chloride in the parts per billion range. Likewise, dimethyl sulfide concentrations are not seen in the parts per billion range in the ambient atmosphere anywhere. Methyl bromide concentrations do not ever hit hundreds of parts per trillion either, and the key for all of this is that there are no known mechanisms by which they would be released or generated in a hospital or home environment. Normally, methyl iodide appears to be essentially completely utilized within the human body (as a strong methylating agent it reacts away while in transit), so the fact that it is observed here is indicative of VERY different processes in this individual’s metabolism and something similar can be said for methane thiol.

RESULTS

○ Did the patient take probiotic therapy, which may account for the increase in Lactobacillus reuteri following treatment?

Thanks - we have addressed this in the introduction now. For your information, participants were asked at baseline regarding their supplement consumption. This participant responded: Cod liver oil capsule every day, orange juice every morning, and regular yoghurts. He did not take probiotics before, or for the duration of, this study.

○ Given the likely mis-assemblies for several key taxa, do the authors think that a single sample-level assembly might assist in teasing this out?

Thank you. We are not convinced this would necessarily help. Metagenomes often contain quite a few misassemblies within the contigs, but given that we are looking at the taxonomy based on contig annotations, small errors such as SNPs caused by errors in the long reads, or chimeric reads, should not contribute to bad annotations because they will still match to the nearest taxonomy. Using all of the timepoints together in the assembly means that low abundance taxa are more likely to be represented

○ Do the different MAGs for Bacteroides coprophilus differ in relative abundance by timepoint? If so, how does this relate to the prevalence of the different resistance genes over time? What about for Bacteroides dorei?

Cluster_contig0024 and Cluster Singleton2 which have lost and maintained the same genes have a similar trajectory, with a low abundance pre-treatment, peak abundance between timepoints 2 and 3, and then the low abundance following the beginning of the second course of treatment (timepoint 4). This low abundance is maintained even post-treatment. Cluster singleton4 and Cluster_contig0951 both lost all resistance
genes. They do not have the same trajectory in relative abundance between timepoints pre- and up to timepoint 3. However, after timepoint 3, both MAGs become very low abundance and do which is maintained even in the post timepoint. Given that the transition between timepoint 3 and 4 represents the transition between different administration routes, it seems like the absence of these resistance genes is contributing to them becoming low abundance following the second treatment course.

I found this sentence very confusing: “Prevotella is also identified but in low abundance, and low but higher levels, compared to Prevotella, of Ruminococcus is also observed.”
Thanks, this has been removed as it didn't add to the point we were making overall regarding the patient’s enterotype.

DISCUSSION
- The increase in *B. coprophilus* relative abundance during ertapenem therapy is interesting. Of course, this is relative, not absolute abundance, which is an important difference. It would be very useful to have some measure of total bacterial load over the course of therapy. Would 16S qPCR help here (even if an imperfect measure)?
- It would, however, 16S abundance still isn't perfect, and we know that 16S sequencing often results in more skews. This would be less with qPCR versus sequencing, but we don't think it would add much more to the overall picture.
- Given the point above, it is probably not correct to make statements such as “*B. coprophilus, a bacterium that has not been associated with human infection, was abundant throughout IV Ertapenem exposure*” – the persistence (or increase) was in relative abundance and may in fact reflect a decline in absolute abundance if all taxa decrease in abundance, as one might expect during broad spectrum antibiotic therapy.

Thanks - we've edited this sentence to clarify it is relative abundance.

*Competing Interests:* None.