Reanalysis of the Mars500 experiment reveals common gut microbiome alterations in astronauts induced by long-duration confinement

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Abstract

Maintaining astronaut health throughout long-duration spaceflight is essential to the feasibility of a manned mission to Mars. The ground-based Mars500 experiment investigated long-duration health by isolating six astronauts for 520 days, the longest controlled human confinement study conducted to date. After 520 days, astronauts had uniform strength and lean body mass losses, and increased fasting plasma glucose, calprotectin, and neutrophil levels characteristic of intestinal inflammation but previous analyses revealed no common significant changes in gut microbiota. This study reanalysed data from early (days 7–45) and late (days 420–520) faecal samples and identified 408 exact sequence variants (ESVs) including 213 shared by all astronauts. Thirty-two ESVs were significantly differentially abundant over time, including depletion of keystone resistant starch degrading, anti-inflammatory and insulin sensitivity-associated species, such as Faecalibacterium prausnitzii, Ruminococcus bromii, Blautia luti, Anaerostipes hadraus, Roseibacteria faces, and Lactobacillus rogosae, and enrichment of yet-to-be-cultured bacteria. Additionally, the extraordinary experimental confinement allowed observation of microbiota potentially shared between astronauts and their habitat. Forty-nine species were shared, representing 49% and 12% of the human and environmental microbiome diversity, respectively. These findings reveal the microbiota which significantly altered in relative abundance throughout confinement, including species known to influence inflammation and host glucose homeostasis consistent with astronaut symptoms. Identification of microbiome alterations after 520 days of isolation represents a missing piece connecting Mars500 astronaut physiological studies. Knowledge of the impact of long-term confinement upon the human microbiome helps to improve our understanding of how humans interact with their habitats and is a valuable step forward towards enabling long-duration spaceflight.

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1. Introduction

The International Space Exploration Coordination Group, which now comprises 20 national space agencies, recently published the third edition of the Global Exploration Roadmap [1] outlining a strategy to expand human presence in the solar system and setting the surface of Mars as a collaborative target for 2030. Safeguarding astronaut health is a critical factor in meeting this ambitious target. The Canadian Space Agency currently considers the major human health risk categories associated to space flight as: musculoskeletal (reduced muscle strength, aerobic capacity and bone fragility), sensorimotor (visual impairment), metabolic (nutritional status and sickness), behavioural health and performance (stress, fatigue, cognitive deterioration and wellbeing), radiation (ultraviolet and ionising), autonomous medical care (medical access), and physiological adaptation to variable gravity and environmental stressors (including human-associated microbial communities) [2]. These challenges are magnified in long-duration space flight such as during a two–three year mission to Mars, where no resupply of air, water, food or medical supplies is possible, radiation and microgravity exposure will be for longer than has yet been tested, and abotion of a mission due to crew illness will not be possible [3,4].

While the human gastrointestinal track (GIT) microbiome is still being explored, it is considered essential for long-term maintenance of many aspects of human health, including healthy

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nutritional, immune, metabolic and psychophysiological function [5–10]. Long-duration habitat confinement, such as that faced by astronauts during spaceflight, could interrupt the dialog between environmental microbial ecosystems and the human microbiome [4]. Previous research has established that confined habitats subject to strict cleaning procedures on Earth, such as intensive care units and surgical theatres, as well as those in space, can produce a unique surface environment dominated by human GIT microorganisms [11–14]. The influence of confinement upon the microbiome has been studied recently in four astronauts over 60 days by Chen et al [15] using bioregenerative life support systems (confined and self-sufficient ecosystems). Although no common microbiome composition was observed over time they did report a potential non-significant trend of depletion of Faecalibacterium prausnitzii as well as large individual differences between individuals (expected in microbiome studies with very low participant numbers). Whether confinement can influence the GIT microbiome of long-term inhabitants is an important priority for future space missions [16] and was directly addressed in the longest-running human confined habitat experiment conducted to date, the Mars500 experiment [17].

The Mars500 experiment isolated six astronauts within an analogue Mars-surface habitat. After 520 days, the six astronauts emerged as largely healthy, but the long-term confinement had significantly altered some of their physiology. Research conducted by Strollo et al [18] measured significant decreases in total body mass (−9.3%), BMI (−5.4%) and lean (non-fat) mass (−11.8%) by day 417 of isolation. When considered alongside significant increases in fasting, plasma glucose from a healthy median of 4.65 mmol/L at the beginning of the experiment to prediabetic levels of 6.02 mmol/L (impaired fasting glucose) [19], these findings are indicative of potential disruption in glucose metabolism and insulin sensitivity in astronauts after extensive confinement. This aligned with significant losses of up to 22% of quadriceps/hamstring strength in astronauts, observed by Gaffney et al. [20]. In parallel, Roda et al. [21] observed that the loss of body mass was accompanied by a significant increase in the faecal protein calprotectin, the intestinal inflammation biomarker making up 60% of neutrophils (the cells which characterise intestinal inflammation [22]); from calprotectin negative in astronauts at the beginning of the experiment, towards varying degrees of positive by day 475. Further supporting this general trend towards an onset of intestinal inflammation, Yi et al. [23] directly observed a significant increase in proinflammatory cytokines and neutrophils in the Mars500 astronauts after the 520 days of isolation, symptoms which are commonly observed in astronauts during space flight [24]. While Strollo et al. [18] hypothesised that the observed symptoms accompanied by an increase in calprotectin in later faecal samples could be indicative of common metabolic alterations associated with intestinal inflammation, they discounted any microbiome interactions due to prior research conducted by Turroni et al. [17], which did not identify significant microbiome changes over time and found only 14 OTUs (operational taxonomic units, representing putative microbiome species) were shared between the six astronauts. Here, the potential for common microbiome change in the Mars500 astronauts over time is reassessed using improved 16S rRNA gene amplicon bioinformatics technology [12] and a common data normalisation strategy for all six astronauts.

## 2. Materials and methods

### 2.1. Sample collection and microbiome sequencing

The Mars500 experiment was a ground-based human isolation study conducted over 520 days which aimed to simulate a manned mission to Mars [17,20,25]. The study participants were all male and their ages ranged from 29 to 40. Faecal samples used for analysis here were taken on the days 7, 14, 21, 30 and 45 (early) and days 420, 450, 490, 504, 520 of the Mars500 experiment (Supplementary file 1). Sampling, DNA extraction, amplification and sequencing approaches were not conducted in this study but are reported in [17]. Briefly, the authors report that total DNA was isolated from faeces using a modified bead-beating and column extraction with QIAamp DNA Stool miniKit. Amplification used 341F (‘5’-CCTACGGGNGGCWGCAG-3’) and 805R (‘5’-GACTACHVGGGTATCTAATCC-3’) primers targeting the V3-V4 hypervariable region and amplicons were sequenced using an Illumina MiSeq platform (2 × 300 bp paired-end). Raw sequences available in NCBI SRA BioProject PRJNA358005 or from [https://genelab.nas.gov/](https://genelab.nas.gov/).

Mars500 habitat environmental samples used for analysis were taken from surfaces using swabs on the days 14, 44, 436, 467, 496 to correspond to faecal microbiome sampling and were from the utility, medical or habitat modules (Supplementary file 1). Sampling, DNA extraction, amplification and sequencing approaches were not conducted in this study but are reported in [25]. Briefly, the authors report that total DNA was isolated from surfaces using a 552C regular ethylene oxide sterilised swab moistened with PCR grade H2O. Total DNA was extracted using a FastDNA SPIN Kit. Amplification used 515F (‘5’-GTGCCAGCMGCCGCGGTAA-3’) and 926R (‘5’-GGTYCTAAATATTGATATGG-3’) primers targeting the V4 hypervariable region and amplicons were sequenced using an Illumina MiSeq platform (2 × 300 bp paired-end). Raw sequences were deposited in the European Nucleotide Archive accession number: PRJEB21072.

### 2.2. Data processing

Amplicon sequence reads were processed and annotated using Anchor [12,26–28]. The amplicon terminology of exact sequence variants (ESV) are used in place of operational taxonomic unit as no 97% similarity clustering is used [29–32]. Sequences were aligned and dereplicated using Mothur [33] and an ESV count threshold of 18 for the astronaut faecal microbiome dataset and 30 for the Mars500 habitat dataset. Annotation at family, genera or species-level used BLASTn criteria of BLASTn criteria of >99% identity and coverage to the NCBI curated bacterial and Archaea RefSeq and NCBI nr/nt databases (January 2020 versions). All annotation calls are putative despite >99% similarity, as databases are subject to change as discoveries are made and phylogenies revised. Reporting of 100% similar sequence matches to unique species is more accurate than reporting such hits at higher taxon levels [12]; however, all annotation should be considered as putative and interpreted with care. Differentially abundant ESVs annotated from the NCBI nr/nt database were manually assessed for quality. Annotation improvement is possible as databases improve/expand using the sequences provided with each ESV in Supplementary file 1. When the highest identity/coverage is shared amongst multiple different putative annotation, all annotation is retained and reported. Amplicons with low-counts (<30 or <18) are binned to high-count sequences in a second BLASTn, using a lower threshold of >98% identity/coverage.

### 2.3. Statistical analysis

The normalisation strategy conducted by Turroni et al. [17] was distinct for each astronaut to allow observation of individual microbiome change in steady states [34] over time. While this described some unique elements of each astronaut, the strategy made observation of common OTUs challenging. Here, while the limited biological replication of 6n (astronauts) could not be
altered, the variation introduced from single replicate stool sampling [35–37] could be reduced by pooling libraries from the first five samples and from the final five samples to improve representation of early (7–45 days) and late (420–520 days) microbiome states of the each astronaut. Samples selected to represent early and late periods of confinement were defined as the first and last set of five samples taken from all six astronauts and the libraries were pooled in silico per astronaut and period. These timepoints also represent the samples available for all six astronauts which best aligned with the Mars500 physiology studies measuring astronaut body mass [18], strength [20], faecal protein calprotectin [21] and immune responses [23]. Differential abundance analysis was performed using DESeq2 [38,39], which performs well with 16S rRNA gene amplicon data and uneven library sizes [40]. A false discovery rate (FDR; Benjamini-Hochberg procedure) < 0.1 was applied [41]. Sparsity and count thresholds were applied whereby an ESV count in a single sample is <90% of the count in all samples and ESV counts must be >0 in at least 3 samples from the same group (early or late) [12]. Constrained ordination analysis was performed using canonical analysis of principal coordinates (CAP; Fig. 2) with the 1-dimensional constraint (early vs late) represented on the x-axis and the first axis of non-constrained ordination (multidimensional scaling) on the y-axis. This analysis was performed with the Vegan library [42] on R via the capscale function.

2.4. Data and code availability

ANCHOR code is available at https://github.com/gonzalezem/ANCHOR. ESV tables including relative abundance, annotation, count distribution, blast statistics, alternative database hits, and sequences are provided in Supplementary file 1.

3. Results

3.1. Mars500 astronaut gastrointestinal track microbiome community overview

A total of 408 exact sequence variants (ESVs) were assembled and captured 71.78% of the total 4,837,317 amplicons sequenced, including 213 ESVs present in all six astronauts in the early (7–45 days) and late (420–520 days) samples alone (Supplementary file 1). Amplicon lengths ranged between 439 and 466 nt. All ESVs were annotated as >99% similar sequence identity: 162 ESVs as putative species (including 14 which were ambiguous sequences to multiple species), 47 ESVs at genera-level, 34 at family-level, while 165 were poorly classified or unknown (<99% similarity to any well-characterised taxa) (Fig. 1). ESVs which could be annotated at species-level had an average identity of 99.92% (135 ESVs had 100% sequence similarity) and captured 67% of counts, those annotated at genera-level averaged 99.83% identity and captured 7% of counts, whereas those annotated at family-level averaged 99.92% identity and captured 11% of counts. ESVs identified from the phylum Firmicutes were the most prevalent, making up 64% of counts, while 7% were from Bacteroidetes, 6% from Actinobacteria, 6% from Verrucomicrobia, 2% from Proteobacteria and 15% were from unknown organisms.

The ESVs which could be confidently identified as putative species with the highest relative abundance across all astronauts in the phylum Firmicutes were all Clostridia, including Agathobacter_rectalis_1 with 10.1% of total counts, Gemmiger_formicilis_1 with 3.5%, Faecalibacterium_prausnitzii_3 with 3.3%, Fusciatenibacter_saccharivorans_1 with 2.5% and Blautia_wexlerae_1 with 2.1% (Supplementary file 1). In Verrucomicrobia, two distinct ESVs had high relative abundance, Akkermansia_muciniphila_1 with 3.5% of total counts and Akkermansia_muciniphila_2 with 2.3% (both having 100% similarity to distinct strain groups, ie. JCM30893 and BSH01), whereas Bacteroides_dorei_1 had the highest relative abundance in the phylum Bacteroidetes with 1.2% of total counts. The ESVs with the highest relative abundance from the phyla Proteobacteria and Actinobacteria were both ambiguous, with the amplified sequence identical between multiple species (ESVs labelled as _MS), Enterobacteriales_MS_1 (sequence common to Bacteroides_subtilis, Escherichia coli, Escherichia fergusonii, Shigella flexneri or Shigella sonnei) and Bifidobacterium_MS_2 (sequence common to Bifidobacterium adolescentis, Bifidobacterium faecale or Bifidobacterium stercoris).

3.2. Significant microbiome changes identified across astronauts over time

To determine whether there were common microbiome changes over time, a simplified experimental design was used to compare the first five samples (7–45 days) and the last five samples (420–520 days) taken from all six astronauts (Fig. 2). When comparing early with late GIT microbiome samples taken from all six astronauts, alpha-diversity (Shannon and inverse Simpson) indices were not significantly different in estimated diversity between groups (t-tests p > 0.05). Canonical analysis of principal coordinates suggest samples separated by early and late groups (Fig. 2) and multivariate analysis identified significant variance between the groups (PERMANOVA, p < 0.001). To identify which ESVs were significantly altered in relative abundance over time, differential abundance analysis was performed. Thirty-one ESVs were significantly different between timepoints, including 16 significantly enriched and 15 significantly depleted in late samples compared to early samples.

3.2.1. ESVs higher in relative abundance in late astronaut samples

The 16 ESVs in significantly higher relative abundance in the astronauts after 420–520 days of confinement were all poorly characterised with the exception of Streptococcus_thermophilus_1 (Streptococcus salivarius subsp. thermophilus). Five ESVs have been putatively placed at genera or family levels (Fig. 2): Kineothrix_1, Lachnospiraceae_10, Ruminococcus_3, Ruminococcaceae_1 and Christensenellaceae_1. The remaining ten differentially abundant ESV are classified here as unknown (<99% similarity to previously identified taxa), although all have previously been observed as unknown bacteria at >99% sequence similarity.

Kineothrix_1 and Lachnospiraceae_10 were both most similar to the Kineothrix alysoides strain KNHS209, with 97.50% and 98.41% similar sequence identity, respectively. The ESV Kineothrix_1 did share 100% similarity to a genome assembled from work sequencing difficult to culture (yet-to-be-cultured) microbiome species from the human GIT (uncultured Clostridium sp. isolate 27895TDY5608883) [43]. Ruminococcus_3 is commonly observed at 100% sequence similarity (as an unknown bacteria) in human GIT microbiome studies (for example NCBI:AM275462 [44]) and has previously been classified as Ruminococcus sp. strain 95 (unpublished, but most similar to Ruminococcus bromii YE282 genome [45]) although the most similar well-characterised species was the type strain Ruminococcus bromii ATCC 27255 (97.97% similar), Ruminococcaceae_1 and Christensenellaceae_1, although also commonly observed as unknown bacteria in human faecal samples at 100% sequence similarity, were both highly dissimilar to any known bacterial species (<94% 16S rRNA sequence similarity).

The ten unknown sequences identified as significantly differentially abundant and enriched in astronauts after long-term confinement included the three ESVs, Unknown_254, Unknown_268 and Unknown_354, sharing 96–97% similarity to the known species, Anaerococcus cellulolyticus, Monoglobus pectinilyticus and Intes-
tinimonas butyriciproducens, respectively. The seven remaining unknown ESVs had <90% sequence similarity to any known species but have all been previously observed as unknown bacteria (at >99% sequence similarity) in GIT samples of humans as well as other animals [46–52]. Two of these ESVs, Unknown_309 (the highest in relative abundance) and Unknown_49, shared 100% sequence similarity with genomes recently assembled by Almeida et al. [53] in breakthrough research targeting yet-to-be-uncultured species in humans with de novo whole genome shotgun techniques (uncultured Clostridium sp. isolates UMGS1238 and UMGS1543).
The Unknown_309 sequence has also recently been observed (100% sequence similarity) and classified by Hynönen et al. [54] as the "Ct85 cluster" of uncultured GIT bacteria in mammals (AP10s.319, LR595742.1 Ct85 type-f), alongside other ESVs present in astronauts but which did not significantly vary over time: Unknown_40 (AP07s.345 Ct85 type-d), Unknown_104 (AP07s.302 Ct85 untyped) and Unknown_38 (AP07s.190 Ct85 type-b).

3.2.2. ESVs lower in relative abundance in late astronaut samples

After 420–520 days in isolation, 15 ESVs were significantly lower in relative abundance in astronauts. These were better characterised than those enriched over time, including ten ESVs which could be annotated at species level (Fig. 2). These putative species included four distinct Faecalibacterium prausnitzii ESVs (labelled here as Faecalibacterium_prausnitzii_1, 3, 4 and 5), Ruminococcus_bromii_2, Blautia_luti_2, Anaerostipes_hadrus_1, Roseburia_faecis_1 and Lactobacillus_rogosae_1, as well as the ambiguous ESV Clostridium_MS_1, which shared 100% sequence similarity to both Clostridium_disporicum and Clostridium_saudiense. Additionally, two ESVs were poorly characterised at genera level, Roseburia_1 and Lachnospira_1, and three were unknown sequences, Unknown_181, Unknown_220 and Unknown_83.

Six F. prausnitzii ESVs in total were identified in astronauts, with 4/6 significantly changing in relative abundance, reducing over time in all four cases. All of the ESVs corresponded to known strain groups [55–57] with 100% similarity (Fig. 3); depleted ESVs were Faecalibacterium_prausnitzii_1 (identical to the type strain F. prausnitzii ATCC27768 in phylogroup 1 subgroup-A), Faecalibacterium_prausnitzii_3 (identical to F. prausnitzii CNCM 4546; 4573; 4644;
Fig. 3. Phylogenetic tree of differentially abundant ESVs and Faecalibacterium prausnitzii ESV characterisation by alignment. A) Phylogenetic tree of differentially abundant ESVs and their most similar well-characterised species (if >90% similarity) from the curated 16S rRNA gene NCBI RefSeq database. Genomes from uncultured candidate bacterial species assembled within Almeida et al. [53] were compared to unknown ESVs. Jukes–Cantor model with neighbour-joining with 10,000 replications and branch support is shown. The scale bar represents 1 substitution in 100 bp. B) Distinct Faecalibacterium prausnitzii ESVs were aligned against sequences from representative strains with genome assemblies. Phylogroup assignment derives from Benevides et al. [55] and Fitzgerald et al. [57].
M21/2 in phylgroup I/subgroup-A, Faecalibacterium_prausnitzii_4 (identical to F. prausnitzii NCIM 4544; 4542 in phylgroup Ila/subgroup-C) and Faecalibacterium_prausnitzii_5 (identical to F. prausnitzii ACP942/32 in phylgroup Iib/outgroup). The two F. prausnitzii ESVs which didn’t significantly vary between confinement time were Faecalibacterium_prausnitzii_2 (identical to F. prausnitzii NCIM 4543; 4574; A2-165 PacBio phylgroup Iib/subgroup-B) and Faecalibacterium_prausnitzii_6 (identical to F. prausnitzii NCIM 4575 phylgroup Iib/outgroup).

3.3. A direct comparison between the Mars500 astronaut and habitat microbiome communities

To assess the degree to which the astronauts’ microbiomes was associated with their habitat, the surface microbiome of the Mars500 modules, reported by Schwendner et al. [25], was directly compared after reanalysis. A total of 1494 ESVs were assembled from 135 habitat samples, capturing 88.02% of the total 23,417,169 amplicons sequenced (Fig. 4). Amplicon lengths (>0.1% of counts) ranging between 403 and 426 nt. Seven hundred and ninety ESVs could be annotated as putative species (including 341 which were ambiguous sequences to multiple species), 167 ESVs at genera-level, 21 at family-level, while 305 were poorly classified or unknown (<99% similarity to any well-characterised taxa). ESVs which could be annotated at species-level had an average identity of 99.95% (790 ESVs had 100% sequence similarity) and captured 88.39% of counts, those annotated at genera-level averaged 99.85% identity and captured 3.20% of counts, whereas those annotated at family-level averaged 99.88% identity and captured 0.40% of counts.

The V3-V4 16S rRNA gene primer pair used to assess the astronaut GIT microbiome, 341F and 805R, was distinct to the V4 16S rRNA gene primer pair used to assess the Mars500 habitat microbiome, 515F and 926R [17,25]. To directly compare the astronaut and habitat microbiome datasets, ESVs annotated at species level were aligned directly. Alignment of each library revealed 79/162 astronaut ESVs had an identical overlapping V4 hypervariable sequence region (252–253 nts) with one or more of 98/790 habitat ESVs (12.4% of total species), 42 of which were single astronaut-habitat ESV pairs unique to a single species. When gene sequences from genome assemblies were used as scaffolds for astronaut-habitat (gut-environment) ESV pairs, a total of 49 Scaffolds (SCamps) aligned as 100% identical to reference genomes (Fig. 5). Astronaut-habitat SCamps allowed for improved resolution of distinct 16S rRNA gene sequences and taxonomic annotation. Examples where two distinct taxa could be distinguished using SCamps but could not from a single primer pair included distinct Anaerostipes hudson. Bacteroides uniformis, Bacteroides caccae and Eggerthella lenta strain groups. The increased length and hypervariable region coverage of SCamps, between 557 and 590 nt, resolved ESV annotation ambiguity in 8 cases, identifying two distinct Veillonella parvula strain groups (NCTC 18110 and UTD81-3) in the astronaut microbiome, Bifidobacterium longum subsp. longum, Bacteroides thetaetotamiron, Streptococcus thermophilus and Lactococcus lactis in the habitat microbiome, and Bifidobacterium pseudocatenulatum and Bifidobacterium adolescens in both.

4. Discussion

4.1. Mars500 astronaut gastrointestinal track microbiome community overview

Recent metagenomic studies indicate that the majority of gastrointestinal tract (GIT) microbiome members are still unknown or uncharacterised species [53]. It is therefore not surprising that over half of the ESVs identified across the astronauts were uncharacterised (Fig. 1). This complexity is further complicated as substantial proportions of the human microbiome are often highly unique to individuals. The highly distinct microbiome compositions of each individual astronaut were well described by the initial Mars500 GIT microbiome research conducted by Turroni et al. [17] and make general patterns of microbiome change challenging to observe. The original analysis did, however, report similar phyla present across astronauts, including: Firmicutes, Bacteroidetes, Actinobacteria, Verrucomicrobia (although present in all astronauts, bacteria from within this group were originally not detected in two astronauts) and Proteobacteria. Instead of using a distinct normalisation strategy for each astronaut to observe change in steady states [17,34], here, libraries were pooled from the first five samples and the last five samples to improve representation of early (7–45 days) and late (420–520 days) microbiome states (reducing technical noise common to single replicate stool sampling [35–37]) before all astronauts were compared directly to make statistical analysis of the common change over time more straightforward.

4.2. Enriched taxa in astronauts after long-term confinement

The presence of Streptococcus thermophilus (Streptococcus salivarius subspecies thermophilus) (Fig. 2) within the Mars500 habitat is not unexpected as the species is commonly used in the dairy industry and is added to yogurts as live culture to market as bestowing health benefits upon consumption (as a probiotic). Subsequent assessment of astronaut dietary reports during the experiment revealed that the Italian company Granarolo provided lyophilised ‘Yomo’ yogurts to the Mars500 project, which do indeed contain live Streptococcus thermophilus when sold commercially [58]. Interestingly, S. thermophilus is known to be particularly resilient to the freeze-drying process [59], so would likely have been present for consumption as live culture in the habitat. The observed significant increase in relative abundance between early and late confinement times is also coherent as the Italian astronaut reported that the team only ate the lyophilised yogurt in the final 8 months of the experiment (during the simulated “return journey”, reported in a July 2014 interview with astronaut Diego Urbina for the Italian Space Agency’s Outpost 42: Space food in training for Mars). Research suggests that culturing can sometimes capture as low as <1% of a microbial community [60]; however, culture-dependent approaches could have been attempted to confirm viability here. Although bacterial viability was not established, these findings do suggest the introduction of these species to the astronauts has been successfully tracked through to their respective microbiomes.

Three ESVs enriched in astronauts after isolation were identified as unknown but shared 97% sequence similarity to well-characterised bacteria A. cellulolytica, M. pectinilyticus and I. butyriciproducens. A. cellulolytica and M. pectinilyticus are plant cell wall degrading bacteria with carbohydrate active enzymes (CAzY) suites allowing degradation of cellulose and hemicellulose [61], and pectin [62], respectively, alongside corresponding fermentation of xylose, arabinose and galacturonic acid. I. butyriciproducens is a butyrate producing species [63] previously identified as increasing in relative abundance alongside a decrease in other common butyrate producers (such as Faecalibacterium prausnitzii) in patients with chronic pain [20]. While changes in bacteria similar to M. pectinilyticus and A. cellulolytica could be associated with dietary modifications involving fruit or vegetables, alterations to the butyrate producing community could have important implications for GIT homeostasis when considered in the context of increased muscle loss and intestinal inflammation markers over
astronaut confinement. Seven ESVs were dissimilar to any well-characterised species. In complex soil or rhizosphere (plant-soil interface) samples where the majority of microbial species are often yet-to-be-characterised [64,65], difficult to culture groups of bacteria such as the TM7 (group 1) bacteria have previously been observed as differentially abundant [28]. In humans, uncharacterised species often share the most sequence similarity to the TM7 (group 3) bacteria, which are typified by recalcitrance to standard culturing techniques [66] currently thought to stem from their epibiont lifestyle (living on the surface of other microbes) [67]. However, the ubiquitous presence of unknown or poorly characterised microbes in human GIT samples is often not
reported, hindering their study. The scale of uncharacterised human microbiome members was recently well-captured by Almeida et al.[53] using de novo whole genome shotgun metagenomics to assemble 1952 uncultured candidate bacterial species genomes from 11,850 human GIT microbiome samples, some of which shared 100% sequence identity with ESVs observed here (Fig. 3). While reporting unknown or poorly characterised organisms can be challenging, these ESVs are significantly associated with long-term isolation in astronauts. Although little is known as to the role that these microbes might play in health, they do also seem to be consistently present in humans and are here implicated as dynamic members of the human microbiome meriting further study as potentially having novel biological importance.

4.3. Depleted taxa in astronauts after long-term confinement

Although the microbiome species enriched after long-term confinement were largely uncharacterised organisms, the majority of the significantly depleted species were well known. *F. prausnitzii* is one of the best characterised inhabitants of a healthy human GIT and is a butyrate producer with well-documented anti-inflammatory properties whose reduction in relative abundance is consistently associated with intestinal inflammation [68,69] and an increase in calprotectin [70,71]. *F. prausnitzii* also almost always negatively associates with type 2 diabetes (4/5 controlled studies) [72], implicating a putative role in insulin resistance. The significant reduction in *F. prausnitzii* in astronauts after long-term confinement (Figs. 2 and 3) is therefore consistent with the intestinal inflammation, increased faecal calprotectin levels, moderate insulin resistance and lean body mass loss reported by Roda et al. [21], Yi et al. [23], Strollo et al. [18] and Gaffney et al. [20] in Mars500 astronauts over the 520 days of confinement. Interestingly, *F. prausnitzii* phylogroup II is considered to be the less sensitive marker of intestinal inflammation when depleted [57,73]. This aligns to the pattern of depletion in only four putative strain-types observed here, as the two out of six *F. prausnitzii* ESVs which were not significantly depleted over time shared 100% similarity to members of phylogroup IIb [55–57]. Turroni et al. [17] reported a decreasing trend in OTUs associated to *F. prausnitzii* over time when first assessing the Mars500 astronaut microbiomes, but were limited in capability to statistically compare OTUs across astronauts owing to tailoring normalisation to each astronaut as well as because low taxonomic resolution would confute the distinct patterns of relative abundance in these *F. prausnitzii* ESVs.

The five other putative species identified as significantly depleted in relative abundance after long-term confinement, *Ruminococcus bromii*, *Blautia luti*, *Anaerostipes hadrus*, *Roseburia faecis*, and *Lactobacillus roseiscis* (Fig. 2), were each consistent with this association with intestinal inflammation and disruption of glucose metabolism. The *Ruminococcus bromii* ESV was only 99.10% similar to the type strain, *Ruminococcus bromii* strain ATCC 27255, but was 100% similar to the strain L2-36 (NCBI:GCA_002834165.1) (Fig. 3) which is considered a ‘keystone’ degrader of resistant starch in the human GIT [74,75]. Robinson et al. [76,77] first reported a direct relationship between dietary resistant starch and insulin sensitivity improvement but could not explain the association through investigation of host metabolism alone. Although a study in germ-free mice has suggested the impact of resistant starch on insulin resistance could be independent of intestinal bacteria [78], members of the GIT microbiome have largely been considered to underlie the relationship [79–81]. *R. bromii* has been more directly associated to insulin resistance through increased relative abundance, alongside *F. prausnitzii*, in patients with metabolic syndrome after faecal matter transplant from healthy donors which improved insulin sensitivity [82]. Similarly, Benitez-paez et al. [83] recently reported decreases in the relative abundance of *B. luti* (reclassified from *Ruminococcus luti*) associated with insulin resistance of obese children, alongside significant increases in proinflammatory cytokines similar to those observed by Yi et al. [23] in Mars500 astronauts after 520 days isolation. Benitez-paez

![Fig. 5. Astronaut gut-habitat environment connection using sequences from reference genomes. A) Illustration of scaffolded amplicons (ScAmps) construction from paired ESVs and sequences from reference genomes, B) ScAmps shared between Mars500 astronauts and their habitat; gut and environment ESVs aligned as 100% identical to reference genomes.](image-url)
| Name                          | Accession   |
|-------------------------------|-------------|
| A. rectilis strain ATCC 33656 |             |
| Agrobacter_validis            |             |
| Agrobacterium_rectilis        |             |
| A. muciniphila isolate MGYG-HGUT-02454 |         |
| Akkermansia muciniphila_1     |             |
| Akkermansia muciniphila_1     |             |
| Akkermansia muciniphila_2     |             |
| Akkermansia muciniphila_2     |             |
| A. endoderini strain 5BB96/5SCP2FAH4/SNY2CFHA         |     |
| Allistipes endoderini_1       |             |
| A. hadaricus strain BP05/ MGYG-HGUT-02528 |         |
| Anaerostipulales hadaricus_1   |             |
| A. hadaricus strain BP06      |             |
| Anaerostipulales hadaricus_2   |             |
| B. dorei strain CLGST12C01    |             |
| Bacteroides dorei_1           |             |
| B. stercoris strain AM40-34/AM47-5 |         |
| Bacteroides stercoris_1       |             |
| Bacteroides stercoris_1       |             |
| B. uniformis strain 3978 E 13/31/AH-09/31/32 |         |
| Bacteroides uniformis_2       |             |
| Bacteroides_uniformis_1       |             |
| B. uniformis strain ATCC 8492/BIO111-92 |         |
| Bacteroides_uniformis_1       |             |
| B. coccoe strain AM31-16AC/AM16-49B/BIO112-79 |         |
| Bacteroides coccoe_2          |             |
| Bacteroides coccoe_2          |             |
| B. coccoe strain CLD3112C61   |             |
| Bacteroides coccoe_1          |             |
| B. thetaiotaomicron strain 7210 |         |
| Bacteroides, thetaiotaomicron_1 |         |
| Bacteroides, M5_1             |             |
| B. fragilis strain BF98       |             |
| Bacteroides, fragilis_2       |             |
| Bacteroides, fragilis_1       |             |
| B. fragilis strain NCCF343     |             |
| Bacteroides, fragilis_1       |             |
| B. predictabilis strain DSM20435/12140/LMG10359        |     |
| Bifidobacterium, M5_1         |             |
| Bifidobacterium, M5_1         |             |
| B. adolescentis strain NCTC11814/BIOM1 A1-248        |         |
| Bifidobacterium, M5_2         |             |
| Bifidobacterium, M5_3         |             |
| B. longum subsp. longum strain ATCC 55813             |             |
| Bifidobacterium, longum_1     |             |
| Bifidobacterium, M5_3         |             |
| B. gasseri strain ATCC 29149   |             |
| Blautia gasseri_1             |             |
| Blautia gasseri_1             |             |
| B. faecis strain M5S           |             |
| Blautia, M5S                  |             |
| B. faecis strain M5S           |             |
| Blautia, M5S                  |             |
| B. eubacterium strain ATCC 29714 |         |
| Blautia eubacterium_1         |             |
| Blautia, eubacterium_1         |             |
| C. aerofaciens strain ATCC 25986 |         |
| Collinsella aerofaciens_1     |             |
| Collinsella aerofaciens_1     |             |
| C. coccoides strain ATCC 27758 |         |
| Coprococcus, coccoides_1      |             |
| Coprococcus, coccoides_1      |             |
| D. fermentigenes strain ATCC 27755 |         |
| Dorea fermentigenes_1          |             |
| Dorea fermentigenes_1          |             |
| D. longiseta strain DSM 13814  |             |
| Dorea longiseta_1              |             |
| Dorea longiseta_1              |             |
| D. ivoroensis strain DSM 15470 |         |
| Dialestera, ivoroensis_1       |             |
| D. ivoroensis strain DSM 15470 |         |
| D. lenta strain DSM 11747      |             |
| Eggerthella lenta_2            |             |
| Eggerthella lenta_2            |             |
| E. lentus strain DSM 2233      |             |
| Eggerthella lenta_1            |             |
| E. ramulosis strain ATCC 29099 |         |
| Eubacterium ramulosis_1        |             |
| Eubacterium ramulosis_1        |             |
| F. saccharivorans isolate MGYG-HGUT-0215 |         |
| Fusificatenbacter, saccharivorans_1 |       |
| Fusificatenbacter, saccharivorans_1 |       |
| I. bartlettii DSM 16795        |             |
| Intestinibacter bartlettii_1   |             |
| Intestinibacter bartlettii_1   |             |
| Lachnospiraceae strain MGYG-HGUT-02455 |         |
| Lachnospiraceae, L. edwardii_1 |             |
| L. reaper strain ATCC 27753    |             |
| Lactobacillus reaper_1         |             |
| Lactobacillus reaper_2         |             |
| L. lactis subsp. lactiscremonis |         |
| Lactococcus lactis_1           |             |
| Lactococcus lactis_M5_1        |             |
| P. merdae ATCC 4932             |             |
| Parabacteroides merdae_1       |             |
| Parabacteroides mercury_1      |             |
| P. micro strain NCTC11808      |             |
| Parvimonas micro_1             |             |
| Parvimonas micro_1             |             |
| P. farcinum ICM 30044          |             |
| Phaceloscellsia farcinum_1     |             |
| Phaceloscellsia, farcinum_1    |             |
| P. ciprip strain AF79-19       |             |
| Prevotella ciprip_1            |             |
| Prevotella ciprip_1            |             |
| R. timonensis strain Marseille-P326 |       |
| Rombustia timonensis_1         |             |
| Rombustia timonensis_1         |             |
| R. intestinalis 1.1-42          |             |
| Roseburia intestinalis_1       |             |
| Roseburia, intestinalis_1      |             |
| R. faecis strain BIOM1 A1_1    |             |
| Roseburia, faecis_1            |             |
| Roseburia, faecis_1            |             |
| R. faecis strain BIOM1 A1_1    |             |
| Ruminococcus, faecis_1         |             |
| Ruminococcus, faecis_1         |             |
| R. bromii strain ATCC 27259    |             |
| Ruminococcus, bromii_1         |             |
| Ruminococcus, bromii_1         |             |
| R. biformis strain 80-2        |             |
| Ruminococcus, bififormis_1     |             |
| Ruminococcus, bififormis_1     |             |
| R. callidus ATCC 27760         |             |
| Ruminococcus, callidus_1       |             |
| Ruminococcus, callidus_1       |             |
| S. thermophilus strain ATCC 19725 |         |
| Streptococcus, thermophilus_1   |             |
| Streptococcus, M5_11           |             |
| V. parvula strain NCTC11810    |             |
| Veillonella parvula_2           |             |
| Veillonella, M5_2               |             |
| V. dispar strain NCTC11831     |             |
| Veillonella dispar_1            |             |
| Veillonella, dispar_1           |             |
| V. parvula strain U7031-1      |             |
| Veillonella parvula_2           |             |
| Veillonella, M5_5               |             |
et al. [83] also demonstrated B. _luti_ had anti-inflammatory activity using _in vitro_ cell cultures and speculated that a reduction of _B. luti_ within the GIT may contribute to insulin resistance due to increase intestinal inflammation. Both _A. hadrus_ and _R. faecis_ are butyrate producers with the former associated to insulin metabolism, significantly increasing in relative abundance (3-fold) in patients responding to FMT treatment with improved insulin sensitivity and decreased proinflammatory markers [84], and the latter whose reduction has been associated to increased inflammation alongside _F. prausnitzii_ [85]. The _L. rogosae_ ESV shares 100% sequence identity to the type-strain _L. rogosae_ strain ATCC 27753. Although less well-studied and potentially misplaced taxonomically [86], _L. rogosae_ was found to be significantly reduced in relative abundance in patients with systemic inflammatory response syndrome (alongside _R. faecis_ and _Faecalibacteria_ sp.) [87] and in MRSA positive patients with increased serum inflammatory markers (alongside _F. prausnitzii_) [88] when compared to healthy controls in both cases.

Despite the highly unique nature of each astronaut’s microbiome, these findings indicate common changes in specific microbiome species over the 520 days of isolation which are consistent with the reported common significant increases in proinflammatory markers and losses in strength and leanbody mass. While the potential involvement of these observed microbiome species is discussed (alongside appropriate evidence), no causal relationship between their relative abundance and astronaut health is established in this work.

4.4. Commonalities between the Mars500 astronaut and habitat microbiome communities

Enrichment of in-built environments with human-associated bacteria is thought to occur rapidly [89] and surface microbiome research conducted in hospital environments (intensive care units and operating theatres), clean rooms and the international space station (ISS) revealed that the selective pressure of confinement and cleaning procedures can promote acquisition of broad resistance mechanisms in a microbial community adapted to human habitation [11,90]. Other studies into the environmental microbiome of MIR [91] and the ISS [14,92–94] have also revealed enrichment of human-associated microbes, in particular within compartments with the highest human inhabitedness (such as the crew quarters [12]). This phenomenon was also reported in the Mars500 confinement experiment, where the environmental microbiome was explored in samples taken from the habitation, utility and medical modules throughout the course of the 520-day experiment. Schwendner et al. [25] reported that the only bacteria detected in all three Mars500 modules were some of the most common human microbiome species: _Staphyloococcus aureus_, _Staphylococcus epidermidis_, _Staphylococcus haemolyticus_ and _Staphylococcus hominis_. The potential for opportunistic pathogen selection is of particular concern for long-duration space travel where astronaut health is at risk and medical facilities are limited.

Comparisons of 16S rRNA gene OTUs (97% clustered) have previously been performed comparing faecal matter and local water samples to investigate environmental contamination in Kenya [95] using SourceTracker [96]. Improved 16S rRNA gene bioinformatics approaches (ANCHOR [12], UPARSE [97], DADA2 [98] or QIME2 [99]) are able to discern amplicon sequences with more confidence in complex metagenomic samples (Fig. 4), allowing for sequences to be compared directly between experiments (Fig. 5). Different primer pairs can have variable 16S rRNA gene amplification rates for different bacteria, potentially confounding direct quantitative integration of amplicon counts across two primer pairs; however, the Mars500 experiment, specifically the high quality datasets shared by Turroni et al. [17] and Schwendner et al. [25], provided a one-off opportunity for qualitative comparison of the microbiota commonality between a closed system environment and the gut of its entire inhabiting population over 520 days as both of these primer pairs have over 95% coverage of known bacterial species [100]. Direct sequence comparison indicated that 49% and 12% of species were shared between the Mars500 astronaut GIT and environmental habitat microbiomes, respectively, suggesting extensive interactions between the communities.

5. Conclusions

Significant changes in the relative abundance of astronaut microbiome species occurred over the 520 days of isolation. Some enrichment of known bacteria corresponded to changes in the crew’s diet, although significant enrichment of unknown species highlights the ongoing challenge faced in understanding how complex microbiome interactions influence human health. Depletion of major anti-inflammatory gastrointestinal tract bacteria over time is consistent with symptoms associated with the intestinal inflammation and insulin resistance measured during the Mars500 experiment and sometimes observed in astronauts during long-duration space flight. Sequence comparison of the astronauts’ faecal microbiome to the Mars500 environmental microbiome revealed direct large-scale associations between the gut and environmental microbiomes, an observation made possible due to the extreme experimental confinement. Identification of significant species-level microbiome alterations which align with astronaut symptoms connects the Russian, European, Chinese and now Canadian Space Agency Mars500 studies. This initial evidence of a potentially deleterious impact of long-term confinement upon microbiome health is an important step forward in the current drive to enable long-duration spaceflight.

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Author contributions

NJBB, FEP and EG all analysed and interpreted the data and wrote the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Appendix A. Supplementary data**

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