**Intestinal persistence of *Bifidobacterium infantis* is determined by interaction of host genetics and antibiotic exposure**

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**ABSTRACT**

Probiotics have gained significant attention as a potential strategy to improve health by modulating host-microbe interactions, particularly in situations where the normal microbiota has been disrupted. However, evidence regarding their efficacy has been inconsistent, with considerable inter-individual variability in response. We aimed to explore whether a common genetic variant that affects the production of mucosal α(1,2)-fucosylated glycans, present in around 20% of the population, could explain the observed interpersonal differences in the persistence of commonly used probiotics. Using a mouse model with varying α(1,2)-fucosylated glycans secretion (*Fut2*WTor *Fut2*KO), we examined the abundance and persistence of *Bifidobacterium* strains (*infantis, breve* and *bifidum*). We observed significant differences in baseline gut microbiota characteristics between *Fut2*WT and *Fut2*KOlittermates, with *Fut2*WT mice exhibiting enrichment of species able to utilise α(1,2)-fucosylated glycans*.* Following antibiotic exposure, only *Fut2*WT animals showed persistent engraftment of *Bifidobacterium infantis*, a strain able to internalise α(1,2)-fucosylated glycans, whereas *B. breve* and *B. bifidum*, which cannot internalise α(1,2)-fucosylated glycans, did not exhibit this difference. In mice with an intact commensal microbiota, the relationship between secretor status and *B. infantis* persistence was reversed, with *Fut2*KO animals showing greater persistence compared to *Fut2*WT. Our findings suggest that the interplay between a common genetic variation and antibiotic exposure plays a crucial role in determining the dynamics of *B. infantis* in the recipient gut, which could potentially contribute to the observed variation in response to this commonly used probiotic species.

**Keywords**

Microbiome, secretor status, α(1,2)-fucosylated glycans, *Bifidobacterium*, *Bifidobacterium infantis*

**INTRODUCTION**

Host-microbiome interactions play a pivotal role in shaping human physiology. The intestinal microbiome in particular is an important regulator of innate and adaptive immunity (1), metabolic control (2), the central nervous system (3), as well as contributing to energy and nutrient harvest (4), and suppressing pathogen proliferation (5). Given the association between disruption of the commensal gut microbiota and adverse outcomes, there is significant interest in approaches that facilitate its restoration following perturbation. Among the most well-established of these approaches is the ingestion of viable commensal bacteria in the form of probiotics.

Probiotics can be defined as “live microorganisms, which when administered in adequate amounts, confer a health benefit on the host” (6). Most commonly, these take the form of individual strains or multi-strain consortia, of well-characterised commensal bacteria, prepared either as liquid suspensions or in freeze-dried capsules. The principal concept underlying the use of probiotics is that the introduction of live bacteria can re-establish physiological homeostasis by modifying the composition or behaviour of the gut microbiota, or by directly providing regulatory cues to the host. Despite substantial evidence supporting the efficacy of probiotics in principle, their use remains poorly supported by empirical data in many physiological or health contexts (7, 8). Further, substantial inter-individual variance in probiotic persistence has been noted, in part explained by variation in colonisation resistance by the microbiome (9, 10). Consequently, the global probiotics market, which is projected to reach USD 73.9 billion by 2030 (11), is dominated by direct-to-consumer sales, with little or no consideration is given to recipient traits that might substantially influence probiotic efficacy.

Various mechanisms enable the human gut to regulate commensal microbiota composition. One of the principal mechanisms involves the secretion of specific types of sugars that are utilised by beneficial microbial species. Many mucosal constituents and secreted factors are decorated with glycans (oligosaccharides), that are added by a diverse family of glycosyltransferase enzymes (12). Of these, the *FUT2* gene encodes a galactoside α(1,2)-fucosyltransferase, which adds a L-fucose monosaccharide to non-reducing end Gal residues to form Fucα1-2Gal-O-R glycans, termed the H antigens (13, 14). Expressed by multiple mucosal epithelial cell types, this H antigen, is a highly versatile structure that can be further modified to form many other important glycans, including the AB blood group glycans. Because *FUT2* controls the nature of the various α(1,2)-fucosylated glycans secreted by mucosal surfaces, it is commonly referred to as the “secretor” gene (13).

Across the human population, multiple nonsense single nucleotide polymorphisms (SNPs) are found within the *FUT2* gene (15), leading to a “non-secretor” phenotype. The non-secretor phenotype, like the AB blood groups, is one of the more common functional mutations maintained in the population, with approximately one-fifth of people carrying homozygous loss-of-function *FUT2* genes (15). This high carriage of loss-of-function mutations is likely a result of positive selection from altered susceptibility to infections by certain bacterial and viral pathogens (16). However, as fucosylated glycans are an important nutrient source for gut microbes, their absence in non-secretors has been shown to influence commensal microbiota composition (16, 17).

The intact commensal microbiota of secretor individuals is likely to be enriched for glycan utilising bacteria. In contrast, depletion of commensal taxa, for example, through antibiotic exposure, can provide a selective advantage to exogenous glycan utilisers that are absent in non-secretors (9, 18). Probiotic preparations typically contain *Bifidobacteria* (*Bifidobacterium* *adolescentis*, *animalis*, *bifidum*, *breve* and *longum*) and/or Lactobacilli (*Lactobacillus acidophilus*, *casei*, *fermentum*, *gasseri*, *johnsonii*, *paracasei*, *plantarum*, *rhamnosus* and *salivarius*). Both genera include species that encode the specific glycoside hydrolases (GHs), GH29, GH95, and GH151, which can utilise the H antigen. However, both genera also include species without this glycoside hydrolase capacity. Therefore, the ability of a probiotic to colonise and persist in an individual may depend on the presence of secreted glycans and the ability of the introduced bacterial strain to utilize them. This is supported by studies identifying increased persistence of such glycan-utilising species when supplemented with exogenous oligosaccharides (19, 20).

We hypothesised that the interplay between secretor status, the glycan utilisation ability of the probiotic strain, and the presence of a disrupted commensal microbiota due to antibiotic exposure, would collectively influence the abundance and persistence of probiotic populations in the gut. To test this hypothesis, we introduced probiotic *Bifidobacterium* strains into a murine model of secretor/non-secretor status, with or without prior antibiotic depletion of commensal microbiota.

**MATERIALS AND METHODS**

Details of reagent catalogue numbers and resource links are provided in **Supplementary Table S1**.

**Mouse model**

***Establishment of a Fut2*** ***knockout mice***

A *Fut2*KO mouse line was developed using CRISPR/Cas9 technology in C57BL/6 mice (IMSR\_JAX:000664) by South Australian Genome Editing (SAGE). Briefly, a 1230 bp region of the *Fut2* exon region was excised using targeted CRISPR guide sequences. Gene knock out was confirmed by Sanger sequencing and phenotype confirmed by α(1,2)-fucosylated glycan staining of intestinal biopsies using Ulex Europaeus lectin 1 (**Supplementary Fig. S1)**, as described previously (21). Littermate *Fut2*WT and *Fut2*KOmice (six weeks of age, gender-matched) were obtained by mating heterozygous male and female mice originating from F1 heterozygotes.

***Breeding and housing***

All mice were bred and maintained under speciﬁc and opportunistic pathogen free (SPF) conditions at 22°C ± 2°C, under a 12 h light-dark cycle, at the South Australian Health and Medical Research Institute (SAHMRI). All mice were housed in individually ventilated cages, fed an identical diet (Teklad Global 18% Rodent Protein Diet, Envigo, Huntingdon, UK), maintained under the Federation of European Laboratory Animal Science Associations (FELASA) standards and routinely screened using a SNP genotyping panel.

Heterozygous x heterozygous breeding was performed to allow for *Fut2*KO and *Fut2*WT littermates, while also standardising effects of *Fut2* that occur through vertical transmission. *Fut2*KO, *Fut2*HET and *Fut2*WT littermates were co-housed from birth until weaning (~3 weeks), where they were genotyped by PCR amplicon melt curve using primers targeting the outer and inner regions of the *Fut2* gene. *Fut2*KO and *Fut2*WT mice separated into cages after weaning based on sex and *Fut2* genotype (**Figure 1A**). No experiments were performed on *Fut2*HET mice. In all experiments, 6-week old, age- and sex-matched mice were used. Each experimental group consisted of at least 4 cages to control for cage effects.

***Antibiotic treatment***

A cocktail of ampicillin (1 g/L, Sigma-Aldrich) and neomycin (0.5 g/L, Sigma-Aldrich) was provided to mice via drinking water for 7 days. Water bottle volume and mouse weight were monitored to assess water intake and antibiotic activity was confirmed by 16S qPCR (**Supplementary Table S2**) of faecal samples at day 7.

***Probiotic supplementation***

Mice received 5 × 107 colony forming units (CFU)/g of mouse of either *B. infantis, B. bifidum*, or *B. breve* daily for 5 days via oral gavage. A starting gavage concentration of 5 × 109 CFU/mL in PBS was prepared daily from fresh overnight cultures. The dose was selected based on previous reports of safety, persistence, and immune modulation capability (22, 23).

**Bacterial strains**

*B. bifidum* JCM 1255 (ATCC equivalent: 29521), *B. longum subspecies infantis* JCM 1222 (ATCC equivalent: 15697), *B. breve* JCM 1192 (ATCC equivalent: 15700) were obtained from the Japan Collection of Microorganisms (JCM; RIKEN, Saitama, Japan). Utilisation of α(1,2)-fucosylated glycans was assessed by *in vitro* culture in mBasal media with/without supplementation of 5% w/v 2′-fucosyllactose (Layer Origin, Ithaca, New York, USA) (**Supplementary Fig. S2)**. This was confirmed by existing literature (24, 25) and the Carbohydrate-Active enZYmes (CAZy) database (26) showing that *B. bifidum* JCM 1255 encodes external GH29 and GH95 α-1,2-L-fucosidases, but does not consume fucose as a carbon source (27), *B. infantis* JCM 1222 encodes intracellular GH29, GH95 and GH151 α-1,2-L-fucosidases along with glycan transporters, and *B. breve* JCM 1192 encodes only the GH95 family.

**Faecal and tissue collection**

At least two faecal pellets were collected from separated mice at the beginning of the light phase, unless specified otherwise. For intestinal tissue collection, mice were sacriﬁced by CO2 asphyxiation and laparotomy was immediately performed using a vertical midline incision. Once the digestive tract was exposed, separate dissection tools were used to dissect tissue into four parts: the proximal small intestine; distal small intestine; caecum and large intestine. For small and large intestine tissue segments, the luminal content was collected by instilling sterile PBS using a syringe barrel and the flushed mucosal tissue was collected into separate tubes. All collected faecal samples, organs, and luminal contents were immediately frozen on dry ice and stored at -80°C until further processing.

**DNA extraction**

***DNA extraction of faecal samples***

Faecal pellets were weighed, and 25 mg samples (±10 mg) were resuspended in 300 µL of cold phosphate buffered saline (PBS) (pH 7.2) by vortexing and pelleted by centrifugation at 10,000 × *g* for 10 min at 4°C. Microbial DNA was extracted from faecal samples using the PowerLyzer PowerSoil DNA Isolation Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions as described previously (28).

***DNA extraction on mucosal tissue samples***

Mucosal tissue from the proximal small intestine, distal small intestine, and large intestine were semi defrosted and 3 cm was removed from the tissue centre using sterile scalpel. The dissected tissues were cut open longitudinally and mixed with 750 µL PowerSoil bead solution and 60 µL solution C1 in a PowerSoil bead tube. The bead tube was then incubated at 65°C for 10 mins prior to bead beating. The subsequent DNA isolation was performed using the PowerLyzer PowerSoil DNA Isolation Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions as described previously (28).

**16S rRNA gene amplicon sequencing and bioinformatic processing**

Amplicon libraries of the V4 hypervariable region for 16S rRNA gene were prepared from DNA extracts using modified universal bacterial primer pairs 515F and 806R (29). Amplicon libraries were indexed, cleaned, and sequenced according to the Illumina MiSeq 16S Metagenomic Sequencing Library Preparation protocol on a 2 × 300 bp Miseq reagent kit v3 at the South Australian Genomics Centre (SAGC), South Australian Health and Medical Research Institute. Paired-end 16S rRNA gene sequence reads were analysed using QIIME2 version 2021.11.0 (30). Briefly, de-noising was performed on de-multiplexed sequences using the DADA2 plugin (31), resulting in a mean read depth of 15,563 ± 2,719 for stool and 6,941 ± 4,503 for tissue. Taxonomic classification of amplicon sequence variants (ASVs) was performed based on the V4 hypervariable region of the SILVA 16S rRNA gene reference database (version 138) at 99% similarity (32). Sufficient coverage at this depth is confirmed by the rarefaction curve, which reached an asymptote. Sequence data has been deposited in the NCBI SRA under accession number PRJNA1011386.

**Microbiota characterisation**

The taxonomic relative abundance at the genus level was used to generate alpha diversity (within-group) and beta diversity (between-group) measures. Alpha diversity metrics (observed ASVs, Pielou’s evenness, Shannon diversity, and Faith’s phylogenetic diversity) were obtained from QIIME2 at sampling depth of 9,883 reads (faecal samples) and 662 reads (mucosal tissue samples). The Bray-Curtis dissimilarity index was calculated to compare microbiome similarity between groups (beta diversity), using square-root transformed species relative abundance data using the ‘vegan’ package in R. Non-metric multidimensional scaling (nMDS) for all beta diversity measures were generated using the ‘vegan’ package in R. Core taxa were defined as those present in more than 95% of samples, with a mean relative abundance of >0.01%. Identification of taxa with α-1,2-L-fucosidases capability was determined by comparing the genus-level taxonomic classification to genomes identified by CAZy (26) as carrying either the GH29, GH95, or GH151 enzyme families.

**Quantification of *Bifidobacterium* speciesand total bacterial load**

Quantification of total bacterial load, *B. breve*, *B. infantis* and *B. bifidum* was performed by SYBR Green-based qPCR assays (**Supplementary Table S2)**. For all qPCR assays, 1 µL of DNA template was combined with 0.7 µL of 10 µM forward primer, 0.7 µL of 10 µM reverse primer, 17.5 µL of 2×SYBR Green (Applied Biosystems, Waltham, Massachusetts, USA) and 15.1 µL nuclease-free water. All samples were run in triplicate (10 μL each replicate). Gene copy quantification was performed using a standard curve generated from a known concentration of a pure colony control. Any sample with a cycle threshold (CT) ≥40 cycles was deﬁned as 40 (limit of detection).

**Culture of *Bifidobacterium* strains**

All *Bifidobacterium* strains used in this study were cultured in De Man, Rogosa and Sharpe (Becton Dickinson, Franklin Lakes, NJ, USA) broth or agar supplemented with 0.34% (w/v) sodium ascorbate and 0.02% (w/v) cysteine-HCl (MRS-CS) and were grown under anaerobic conditions (75% N2, 20% CO2, 5% H2, Coy Laboratory Products, Grass Lake, Michigan, USA) at 37 °C. The growth of *Bifidobacterium* was measured by optical density (600 nm) using multimode plate reader (PerkinElmer, Waltham, Massachusetts, USA).

***In vitro* glycan utilisation assay**

Faecal pellets from untreated WT mice were incubated in mBasal media (mBasal; 10 g/L Trypton, 2 g/L yeast extract, 5 g/L NaCl, 0.2 g/L magnesium sulfate, 2 g/L dipotassium hydrogen phosphate, pH 6.4) with/without supplementation of 5% w/v of 2′-Fucosyllactose (Layer Origin, Ithaca, New York, USA) at 37°C under strict anaerobic conditions (75% N2, 20% CO2, 5% H2, Coy Laboratory Products, Grass Lake, Michigan, USA). Bacterial biomass was measured by optical density (600 nm, OD600) using multimode plate reader (PerkinElmer, Waltham, Massachusetts, USA). Bacterial colonies cultured from faecal samples were identified by matrix assisted laser desorption/ionization time-of-flight (MALDI-TOF) (Billerica, Massachusetts, United States), as described previously (33).

**Statistical analysis**

Experimental mice were randomly assigned to different treatment groups. The investigators were not blinded to the experimental groups. No outliers have been removed from any of the data presented. All data analyses were performed using either R (R Foundation for Statistical Computing; version 4.1.0) or GraphPad Prism software (GraphPad Software, Inc.; version 9.00). For parametric data, unpaired Student’s t test was used to compare data between two unpaired groups; One-way ANOVA was used to compare data among three or more unpaired groups. For non-parametric data, Mann-Whitney *U* test was used to compare data between two unpaired groups; Kruskal-Wallis test was used to compared data among three or more unpaired groups. Differences in Bray-Curtis dissimilarity between groups was performed by permutational multivariate ANOVA (PERMANOVA) and pairwise PERMANOVA, using the “adonis” package in R, with 9,999. Linear discriminant analysis Effect Size (LEfSe) was applied to identify the abundant taxa in each site, using default parameters (34). Area under the curve (AUC) was calculated for *in vitro* growth experiments (using OD600 values) and bifidobacterial persistence in mice (using copies/ng faecal DNA). Log-rank test was employed to compare survival time differences based on bacterial qPCR detection. One-tailed tests were used where differences between groups were hypothesised to be in a single direction. Statistical outcomes with *P* value <0.05 were considered statistically significant. Core taxa plot was generated using GraphPad Prism, other data were visualised using R.

**RESULTS**

***Fut2* shapes the faecal microbiota**

Assessment of the faecal microbiota between SPF *Fut2*WT and *Fut2*KO littermates was performed at 6 weeks of age in both male and female mice (**Fig. 1A**). Faecal microbiota composition (**Fig. 1B**) differed significantly between *Fut2*WT and *Fut2*KO littermates (PERMANOVA: *R*2=0.028; *P=*0.028, **Fig. 1C**) when male and female mice were assessed together. However, stratification according to sex identified a greater divergence according to genotype in male mice (PERMANOVA: *R*2=0.12; *P=*0.021) compared to female mice (PERMANOVA: *R*2=0.037; *P=*0.38, **Fig. 1C**). Exploration of this sex effect identified a significant interaction between sex and *Fut2* genotype (PERMANOVA: R2=0.11; *P=*0.0068, **Supplementary Table S3**). These findings were unchanged after adjustment for cage effects (**Supplementary Table S3**). Given the interaction between sex and genotype, all subsequent experiments involved male mice only.

Potential relationships between secretor status and microbiota characteristics were then explored. While bacterial alpha-diversity measures did not differ substantially between differed between *Fut2*WT and *Fut2*KO groups (**Supplementary Fig. S3**), the membership of core microbiota (taxa present in ≥95% of samples at ≥0.01%) did (**Fig. 1D**). Specifically, *Oscillospiraceae* (unassigned) were exclusively core in *Fut2*WT, whereas *Bifidobacterium*, Clostridia (UCG014), *Lachnoclostridium*, and *Parabacteroides* were exclusively core in *Fut2*KO mice (**Fig. 1E**). Three bacterial genera had a significantly higher relative abundance in *Fut2*WTmice, whereas 13 were more prevalent in *Fut2*KO mice (**Fig. 1F**). Of these, *Lactobacillus,* a genus with the α-1,2-L-fucosidases GH29 and GH151 (**Supplementary Table S4)**,was more abundant in *Fut2*WT (LDA score [log10]=4.88). Together, these findings support previous reports of a relationship between *Fut2* and faecal microbiome characteristics.

**The distal small intestine microbiota is influenced by *Fut2***

As *Fut2* is differentially expressed throughout the gastrointestinal tract (35), we then compared microbiota characteristics in tissue from the proximal small intestine, the distal small intestine, and the large intestine between *Fut2*WT and *Fut2*KO mice (**Fig. 2A**). Microbiota composition differed significantly between *Fut2*WT and *Fut2*KO groups in the distal small intestine (PERMANOVA: *R*2=0.54; *P=*0.0071, **Fig. 2B**), coinciding with inducible *Fut2* expression, but not in the proximal small intestine (PERMANOVA: *R*2=0.19; *P=*0.13, **Fig. 2B**) or the large intestine (PERMANOVA: *R*2=0.16; *P=*0.18, **Fig. 2B**). Taxonomic assessment identified that *Candidatus* Arthromitus (LDA score [log10]=5.43) was more abundant in the distal small intestine in *Fut2*WT compared to *Fut2*KOmice, whereas *Lachnospiraceae* (unassigned) and *Acetatifactor* were more abundant in in the large intestine of *Fut2*WT compared to *Fut2*KOmice(LDA score [log10]=4.35, and LDA score [log10]=3.93, respectively) (**Fig. 2C**).

***Fut2*-microbiota relationships can be recapitulated *in vitro* through glycan exposure**

To further investigate the relationship between α(1,2)-fucosylated glycans and intestinal microbiology, faecal homogenate from *Fut2*WT mice was grown in a mBasal media with or without the α(1,2)-fucosylated glycan, 2ˈ-FL. Microbiota assessment following *in vitro* culture (**Fig. 3A**) confirmed the findings of the *in vivo* faecal microbiota analysis, with a significant difference in Bray-Curtis similarity between faecal cultures with and without 2ˈ-FL (PERMANOVA: *R*2=0.90; *P<*0.0001, **Fig. 3B**). This difference was marked by an enrichment of glycan-utilising genera (*Bacteroides*, *Enterococcus*, *Lactobacillus*), with compensatory decreases in the relative abundance of other taxa (**Fig. 3C**).

Analysis involving inoculation of solid basal media, either alone or supplemented with 2ˈ-FL, with faecal homogenate from *Fut2WT* mice further supported these findings. Specifically, proportional colony counts (**Supplementary Fig. S4, Supplementary Table S5**) showed 2ˈ-FL led to enrichment of *Enterococcus faecalis* (66% with 2ˈ-FL vs 4% without 2ˈ-FL) and *Lactobacillus murinus/reuteri* (18% with 2ˈ-FL vs 1% without 2ˈ-FL), with a corresponding decrease in *E. coli* (16% with 2ˈ-FL vs 95% without 2ˈ-FL) (**Fig. 3D**). Enrichment of bacteria with 2ˈ-FL supplementation was further confirmed by increased growth rate and bacterial density (median AUC[mBasal]=15.1 [IQR=14.3, 16.5]; AUC[mBasal+2ˈ-FL]=35.6 [33.3, 38.7]; *P<*0.0001, **Fig. 3E**).

We next investigated whether differences in microbiota composition between *Fut2*WT and *Fut2*KO mice reflected selection for bacterial populations able to utilise α(1,2)-fucosylated glycans for growth in *Fut2WT* animals. Faecal homogenates from *Fut2*WT and *Fut2*KO mice were used to inoculate mBasal media with or without 2ˈ-FL supplementation. The increase in bacterial density between 2ˈ-FL supplemented media and media alone was significantly greater when faecal homogenates were derived from *Fut2*WT compared to *Fut2*KO mice (median AUC[WT]=20.5 [IQR=16.6, 23.5]; AUC[KO]=16.7 [13.1, 19.7]; *P=*0.0051, **Fig. 3F**), consistent with a greater abundance of glycan-utilizing bacteria.

**Probiotic bifidobacterial species differentially colonise *Fut2*WT and *Fut2*KO mice**

Based on the relationship between host glycan production and gut microbiota composition, we hypothesised that the colonisation dynamics (abundance and persistence) of bifidobacterial probiotic species, introduced following antibiotic depletion, would differ between *Fut2*WT and *Fut2*KO recipients (**Fig. 4A**). We utilised strains of three *Bifidobacterium* species that interact with glycan in different ways; *B. infantis*, an intracellular α(1,2)-fucosylated glycan-utiliser, *B. bifidum*, an extra-cellular α(1,2)-fucosylated glycan utiliser, and *B. breve*, a species that does not utilise α(1,2)-fucosylated glycans (characteristics that were confirmed by *in vitro* growth assays, **Supplementary Fig. S2**).

The persistence of species that do not internalise and degrade α(1,2)-fucosylated glycans (*B. bifidum* and *B. breve*) did not differ substantially when introduced to *Fut2*WT or *Fut2*KO mice (**Fig. 4B-4C**). In contrast, the abundance and persistence of *B. infantis*, an intracellular glycan utiliser, differed significantly according to the genotype of the recipient (**Fig. 4B-4C**). Ten days post-gavage, *B. infantis* was detected in 7 out of 8 *Fut2*WT mice but 0 out of 8 *Fut2*KO mice (Log-rank test: chi-q=11.4; *P=*0.00074, **Fig. 4B**). The overall abundance of *B. infantis* was also significantly higher in *Fut2*WT compared to *Fut2*KO mice in the following 14 days post-gavage (mean AUC[WT]=31.1 [SD=4.1]; AUC[KO]=26.2 [5.2]; *P=*0.028, **Fig. 4C**).

A difference in *B. infantis* persistence was also evident in intestinal tissue assessed five days post-gavage. *B. infantis* was significantly more abundant in *Fut2*WT large intestine (median *Fut2*WT =log10 6.7 copies/g tissue [IQR=log10 6.0, log10 6.8]; *Fut2*KO =log10 2.6 [IQR=log10 2.6, log10 3.9]; *P=*0.011), and numerically, though not significantly, more abundant in *Fut2*WT proximal (*P=*0.085) and distal small intestine (*P=*0.094) (**Fig. 4D**).

**Prior antibiotic exposure profoundly affects the *Fut2*-probiotic relationship**

To test whether the relationship between *Fut2* genotype and probiotic strain characteristics were independent of antibiotic exposure, we supplemented non-antibiotic exposed mice with *B. infantis* (**Fig. 5A**). No significant difference in *B. infantis* persistence post-gavage was observed between *Fut2*WT and *Fut2*KO mice (**Fig. 5B**) and *B. infantis* was not detectable in intestinal tissue from either *Fut2*WT or *Fut2*KO mice at day 5 post-gavage (**Supplementary Fig. S5A**). However, analysis of *B. infantis* abundance following gavage (based on AUC) revealed an effect that was opposite to that observed in antibiotic exposed mice, with *B. infantis* significantly higher in *Fut2*KO mice compared to *Fut2*WT (mean AUC[WT]=3.8 [SD=0.6]; AUC[KO]=5.6 [0.90]; *P=*0.00046; **Fig. 5C**).

Faecal levels of *Bifidobacterium* probiotics were assessed during the instillation period (samples collected two hours prior to gavage and six hours after gavage). Levels in post-gavage samples (6 hours) did not differ between *Fut2*WT and *Fut2*KO mice, consistent with the instillation of equal probiotic loads. However, at 22 hours post-gavage (2 hours prior to gavage), levels in *Fut2*KO mice were significantly higher than in *Fut2*WT mice (*P<*0.0001; **Supplementary Fig. S5B**). No cumulative effect was observed with repeat gavage and the decline in probiotic levels after 22 hours was comparable to that observed in the 24 hours post cessation of installation.

**DISCUSSION**

Although probiotics have shown great potential in modifying host-microbiome interactions (36), their actual performance has been disappointing in many clinical contexts (37). Previous studies have investigated the reasons for this underperformance, relating to cohort-level effects and variation in response between individuals. Mode of delivery and dose have both been shown to contribute to overall efficacy (38, 39), whereas the habitual diet of the recipient, particularly fibre intake, is an important determinant of probiotic response (40). The influence of factors that shape the gut microbiome on the abundance and persistence of probiotics is unsurprising (41, 42), given the ability of resident gut microbiota to competitively exclude introduced populations (10). Indeed, exposure to antibiotics, a factor that greatly impacts the gut microbiome, has been shown to considerably influence probiotic effects at a microbiological level (9, 43). However, while common *FUT2* genetic variants are known to help shape intestinal microbiology (16, 17), the effect of secretor status on probiotics had not been described.

Our study highlights several important points in relation to inter-individual variance in intestinal microbiology and probiotic efficacy. Firstly, with 20% of the global population also homozygous for a non-functional *FUT2* gene (15), our findings suggest that these “non-secretor” individuals will also experience different probiotic population dynamics compared to “secretor” individuals, if the probiotic taken contains one of the many bacterial species able to utilise α(1,2)-fucosylated glycans (H antigens). In our study, this was reflected in the significantly greater transience of *B. infantis* in the faecal and intestinal microbiome of non-secretor (*Fut2*KO) mice compared with secretor mice following antibiotic exposure.

Bifidobacterial species that are commonly used as probiotics are relatively close phylogenetically but differ in their ability to use glycans, even at a strain level (44). We showed that neither *B. breve* (JCM 1192) nor *B. bifidum* (JCM 1255) differed in their abundance or persistence between secretor and non-secretor animals. In contrast, *B. infantis* (JCM 1222) persisted for significantly longer and showed a significantly higher abundance in secretor mice compared to non-secretor mice. This finding likely reflects differences in H antigen hydrolysis and catabolism capacities between species when administered as a probiotic. For example, *B. infantis* encodes GH29, GH95, and GH151 family intracellular α-1,2-L-fucosidases, along with fucose transporters to facilitate internalisation (44). While independent hydrolysis and catabolism of mucin-bound H antigens by *B. infantis* is not hypothesised (25), cross-feeding by organisms with extracellular α-1,2-L-fucosidases is likely, even following antibiotic supplementation (45). In contrast, *B. bifidum*, while expressing extracellular GH29 and GH95 α-1,2-L-fucosidases, does not consume fucose to facilitate growth (27). Finally, *B. breve*, encodes a separate GH95 intracellular α-1,2-L-fucosidase along with fucose transporters. While this species is capable of utilising the H antigen with support from cross-feeding (46), these findings suggest reduced persistence compared with *B. infantis*, when administered as a probiotic.

Secondly, we found that antibiotic exposure influenced the persistence of probiotics in a secretor status-dependent manner. In the absence of microbiota depletion through antibiotic exposure, it would be expected that other commensal bacteria would utilise available glycans within the secretor gut. Moreover, such strains are highly adapted to an individual’s gut environment, making them likely to outcompete any exogenous glycan-utilisers that are introduced. When we explored this directly, we found that in the absence of a prior period of antibiotic exposure, the higher levels and greater persistence of *B. infantis* in secretors was inverted, with these *B. infantis* being significantly higher in non-secretor mice. These findings likely reflect the competitive exclusion of H antigen-utilising probiotics in the secretor gut and highlight the importance of considering the ecological context in relation to probiotic impact.

While this study was performed in mice, the effect of secretor status on bifidobacterium supplementation has important implications for probiotic strategies in humans. It is crucial to consider individual host traits and recent antibiotic exposure when designing a probiotic intervention (47). The findings here suggest that the 20% of the population who are non-secretors may have poorer persistence of H antigen utilising probiotics, such as *B. infantis,* compared to secretorsfollowing antibiotic exposure. Conversely, in the absence of recent antibiotic exposure, higher levels of microbial niche occupancy in secretors may hamper *B. infantis* persistence compared to non-secretors. An individualised supplementation with prebiotics may have potential as a means to optimise probiotic uptake in non-secretors. For example, previous studies have shown that supplementation with human milk oligosaccharides can enhance *B. infantis* engraftment (20), with successful supplementation shown to reduce intestinal inflammation in infants (48). Investigating additional α(1,2)-fucosylated glycans, given as prebiotics, may lead to improved outcomes of *B. infantis* supplementation in non-secretor individuals.

Determining the impact of secretor status on other species commonly considered beneficial and marketed as probiotics is challenging due to their broad range of carbohydrate utilisation capabilities (49). For instance, *Akkermansia muciniphila*, a mucin-degrading species, has been associated with a reduced risk of chronic inflammatory diseases in humans and mice (50), and is a potential target for probiotic development (47). However, its utilisation of mucin glycoproteins, including the α(1,2)-fucosylated glycan, 2’-fucosyllactose (49), suggests that it may also be affected by secretor status. While *Akkermansia* was not detected in the mice of this study, we found that *Candidatus* Arthromitus, another genus associated with immune modulation (51),was enriched in the distal small intestinal mucosal tissue of secretor mice. Genome annotation of *Candidatus* Arthromitus has indicated other fucose utilisation capabilities (52), indicating that the functional *FUT2* gene may promote colonisation by this species.

Our experiments involved SPF mice that were obtained through heterozygous mating. Such breeding was essential to allow comparison of *Fut2*WT and *Fut2*KO littermates from a maternal secretor lineage. The findings from this study are therefore independent of vertical transmission effects, which are known to influence the microbiome of the offspring (53, 54), and indicate that a change in gut microbiology occurred post-weaning. This difference in baseline gut microbiota composition between secretors and non-secretors was also only evident in male mice. The effect of sex on the relationship between secretor status and the gut microbiome is difficult to explain but may relate to variable intestinal expression of *Fut2*, which can be altered factors such as stress (55). In addition, independent interactions between sex hormones and the gut microbiome (56) may affect the relationship between *Fut2* and the gut microbiome.

Our study demonstrates a *Fut2*-dependent genetic determinant for inter-individual response to probiotic supplementation, which is affected by antibiotic exposure and glycan utilisation capabilities of the probiotic strain. With prior antibiotic exposure, *Fut2* functionality was associated with increased persistence of *B. infantis*, consistent with its ability to utilise the H antigen. However, without antibiotic exposure, *Fut2* functionality was associated with lower abundance of *B. infantis,* relating to difference in baseline microbiology and niche space occupation.

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**AUTHOR CONTRIBUTIONS**

The study was conceived by SW, SLT and GBR. SLT, GBR, JMC, and YW designed the experiments. YW and SLT performed the experiments; YW and SLT undertook the data generation and statistical analysis. GBR, JMC, AR, and LEP provided technical support and guidance. YW, JMC, SLT, and GBR drafted the manuscript. All authors approved the final version of the manuscript.

**DECLARATION OF INTERESTS**

The authors declare that they have no competing interests, nor any financial or personal relationships with other people or organisations that could bias this study.

**ETHICS APPROVAL AND CONSENT TO PARTICIPATE**

All mice experiments were approved by the South Australia Health and Medical Research Institute Animal Ethics Committee (application numbers SAM-21-07 and SAM-21-036).

**DATA AVAILABILITY STATEMENT**

All 16S rRNA gene sequencing data have been deposited to the NCBI SRA and are available under accession number PRJNA1011386. All qPCR data is available through FigShare (<https://figshare.com/s/9fa55ea8b65304d9f722>). Data processing, statistical analysis and visualisation is available through GitHub (<https://github.com/Yiming-Wang-1992>).

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**FIGURES AND TABLES**

**Fig. 1: *Fut2* affects the faecal microbiota of male mice.** (**A**) Breeding and co-housing design. *Fut2*HET mice were bred to produce *Fut2*WT and *Fut2*KOlittermates. Genotypes were co-housed until weaning (~3 weeks), where mice were separated by sex and *Fut2* genotype until 6 weeks of age. (**B**) Taxa bar plot of the faecal microbiota of *Fut2*WT and *Fut2*KOmice, also stratified by sex. (**C**) Non-metric multidimensional scaling (NMDS) plot of faecal microbiota showing significant separation between *Fut2*WT and *Fut2*KOmale mice but not *Fut2*WT and *Fut2*KOfemale mice. Significance: Permutational multivariate ANOVA. (**D**) The prevalence and mean relative abundance of faecal taxa in *Fut2*WT and *Fut2*KOmale mice, highlighting the “core” taxa present in ≥95% of mice and at a mean relative abundance of ≥0.01%. (**E**) Venn diagram of core taxa shared or different between *Fut2*WT and *Fut2*KOmale mice. (**F**) Faecal taxa significantly different between *Fut2*WT and *Fut2*KOmale mice. Significance: Linear discriminant analysis (LDA) Effect Size (LEfSe).

**Fig. 2: *Fut2* affects the intestinal microbiota.** (**A**) Taxa bar plot of the intestinal mucosal microbiota of *Fut2*WT and *Fut2*KO mice. (**B**) Non-metric multidimensional scaling (NMDS) plots of *Fut2*WT and *Fut2*KO intestinal mucosal microbiota. Significance: Permutational multivariate ANOVA. (**C**) Taxa significantly different between *Fut2*WT and *Fut2*KO intestinal mucosal tissue. Significance: Linear discriminant analysis (LDA) Effect Size (LEfSe). No taxa differed between *Fut2*WT and *Fut2*KO proximal small intestine.

**Fig. 3: 2′-Fucosyllactose** **modifies the faecal microbiota *in vitro* and enhances growth of α(1,2)-fucosylated glycan utilising bacteria.** (**A**) Taxa bar plot of the faecal microbiota of *Fut2*WT mice following anaerobic growth either with or without 2′-Fucosyllactose (2′-FL). (**B**) Non-metric multidimensional scaling (NMDS) plot of faecal microbiota following anaerobic growth with or without 2′-FL. Significance: Permutational multivariate ANOVA. (**C**) Taxa significantly different between *Fut2*WT faecal samples following anaerobic growth with or without 2′-FL. Significance: Linear discriminant analysis (LDA) Effect Size (LEfSe). (**D**) Comparison of colonies relative abundance of identified colonies following anaerobic growth with or without 2′-FL. \**E. coli* or *Shigella.* (**E**) Optical density (OD) following *Fut2*WT faecal bacteria cultured with or without 2′-FL. (**F**) OD of *Fut2*WT or *Fut2*KO faecal bacteria following growth with 2ˈ-FL. OD normalised to growth in media without 2ˈ-FL.

**Fig. 4: *Bifidobacterium infantis*, but not *B. breve* or *B. bifidum,* persists longer in *Fut2*WT mice following antibiotic pre-exposure.** (**A**) Experimental design. (**B**) Persistence of detectable *Bifidobacterium* species in stool following antibiotic pre-exposure. Significance: log-rank test. (**C**) Bacterial copies of *Bifidobacterium* species in stool following antibiotic pre-exposure. Significance: T-test of area under the curve. (**D**) Bacterial copies of *B. infantis* in intestinal tissue mucosa following antibiotic pre-exposure. Significance: Mann-Whitney U test.

**Fig. 5: *Bifidobacterium infantis* persists longer in *Fut2*KO mice without antibiotic pre-treatment.** (**A**) Experimental design. (**B**) Persistence of gavaged *Bifidobacterium* *infantis* in stool without antibiotic pre-treatment. Significance: log-rank test. (**C**) Bacterial copies of *B.* *infantis* in stool without antibiotic pre-treatment. Significance: T-test of area under the curve.