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Gram-scale chemical synthesis of galactosyllactoses and their impact on infant gut microbiota *in vitro*†

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Galactooligosaccharides (GOS) are widely used as a supplement in infant nutrition to mimic the beneficial effects found in prebiotic human milk oligosaccharides (HMOs). However, the complexity of the GOS mixture makes it challenging to ascertain which of the GOS components contribute most to their health benefits. Galactosyllactoses (GLs) are lactose-based trisaccharides containing a β -galactopyranosyl residue at the 3'-position (3'-galactosyllactose, 3'-GL), 4'-position (4'-galactosyllactose, 4'-GL), or the 6'-position (6'-galactosyllactose, 6'-GL). These GLs are of particular interest as they are present in both GOS mixtures and human milk at early stages of lactation. However, research on the potential health benefits of these individual GLs has been limited. Gram quantities are needed to assess their health benefits but these GLs are not readily available at this scale. In this study, we report the gram-scale chemical synthesis of 3'-GL, 4'-GL, and 6'-GL. All three galactosyllactoses were obtained on a gram scale in good purity from cheap and commercially available lactose. Furthermore, *in vitro* incubation of GLs with infant faecal microbiota demonstrates that the GLs were able to increase the abundance of Bifidobacterium and stimulate short chain fatty acid production.

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Introduction

The establishment of a healthy gut microbiome in early life contributes to healthy immune-development, metabolism and brain development of the infant.^{1,2} As a major component in breastmilk, human milk oligosaccharides (HMOs) have been found to drive the development of a healthy gut microbiota by specifically stimulating the growth of beneficial gut bacteria, such as bifidobacteria.³ The oligosaccharide content in standard infant formula is typically low, as it is based on bovine milk, which has a much lower oligosaccharide content ($\sim 0.05 \text{ g L}^{-1}$) compared to human milk ($\sim 20 \text{ g L}^{-1}$).^{4,5} Therefore, in the last decades, it has become common practice to supplement infant formula products with functional oligosaccharides, such as β -galactooligosaccharides (GOS). GOS has been shown to promote the growth of benign micro-organisms such as bifidobacteria.^{6,7} Upon fermentation by the colonic

microbiota, metabolites like short-chain fatty acids (SCFAs) are formed,^{7,8} which acidify the intestinal environment. This creates a favorable environment for bifidobacteria to grow and outcompete the growth of potentially harmful bacteria, such as *Enterobacteriaceae* or *Clostridium perfringens*.⁹ Next to the role of GOS in modulating the gut microbiome,¹⁰ GOS may also reduce the adhesion of pathogenic bacteria to the intestinal epithelium,¹¹ improve mineral absorption,^{12,13} enhance natural defenses,¹⁴ and improve stool consistency and frequency.^{6,15,16} Commercially available GOS are produced enzymatically using a β -galactosidase enzyme, which normally hydrolyses lactose to form galactose and glucose. However, at high lactose concentrations, kinetically controlled transglycosylation takes place resulting in the intermolecular transfer of the cleaved galactose moiety to another lactose molecule to provide a complex mixture GOS.^{7,17,18} The most widely used GOS in infant formula, Vivinal® GOS, is a heterogeneous mixture of >100 linear and branched oligosaccharides.^{19,20} The complexity of the GOS mixture makes it challenging to establish which of the GOS components contribute most to the development of a healthy infant microbiome. Nevertheless, three galactosyllactoses (GLs) present in GOS, 3'-galactosyllactose (3'-GL), 4'-galactosyllactose (4'-GL), and 6'-galactosyllactose (6'-GL), are of particular interest (Fig. 1) as they are also found in human milk and can hence be classified as a human milk

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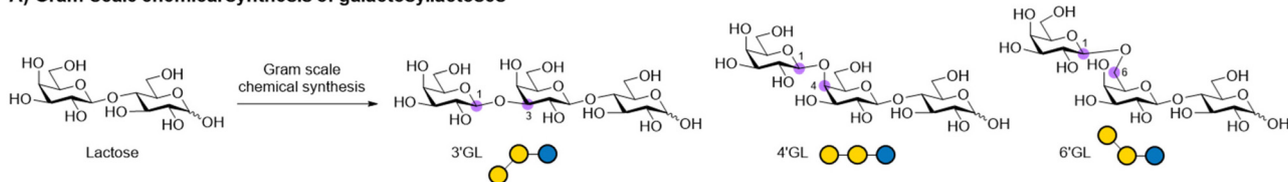
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A) Gram-scale chemical synthesis of galactosyllactoses



B) Experimental set-up to assay the effects of GLs on infant faecal microbiota

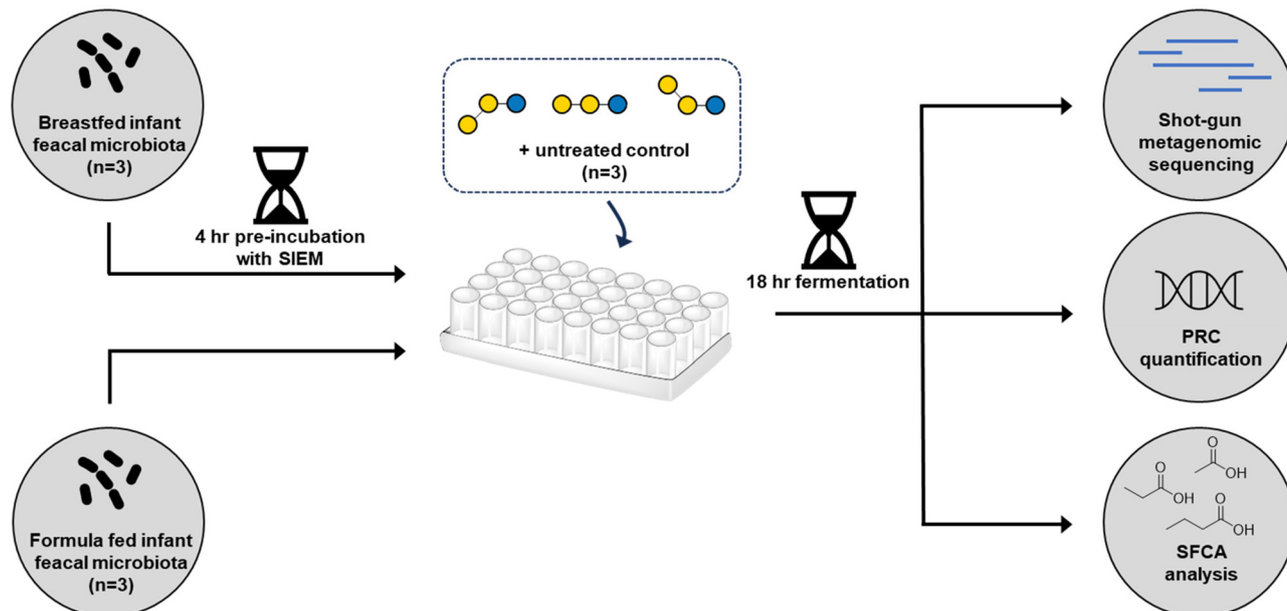


Fig. 1 (A) The structure of the three galactosyllactose trisaccharides, and (B) work flow of their *in vitro* fermentation by infant faecal microbiota and impact on microbiota composition and metabolite production.

oligosaccharides (HMOs).²¹ GLs are found in higher concentrations in colostrum compared to mature milk and have been linked to anti-inflammatory properties in the intestine *in vitro/ex vivo*.²¹ To further dissect the role of specific GLs in shaping the microbiome, chemically pure samples are needed. Purification of GLs from GOS mixtures is highly challenging due to the micro-heterogeneous nature of these samples. Chemical GL synthesis can generate pure material, but needs to be scaled to produce the gram scale quantities needed for research into their potential health benefits.^{22,23} Chemical procedures to prepare GLs at this scale are currently unavailable to the best of our knowledge.

Herein, we report the development of a gram-scale chemical synthesis of 3'-GL, 4'-GL, and 6'-GL starting from readily available lactose. The general synthetic strategy for the GL trisaccharides 13–15 employs the synthesis of three different lactose-derived acceptors (4, 7 and 8), which are extended on the non-reducing end with either a β 1,3-, β 1,4-, or β 1,6-galactopyranosyl residue using galactosyl donor 9. Next, we investigated the impact of 3'-GL, 4'-GL and 6'-GL on infant gut microbiota *in vitro* by fermentation experiments using infant faecal samples. We found that GLs are important components in GOS as they clearly contribute to its effects on infant microbiota.

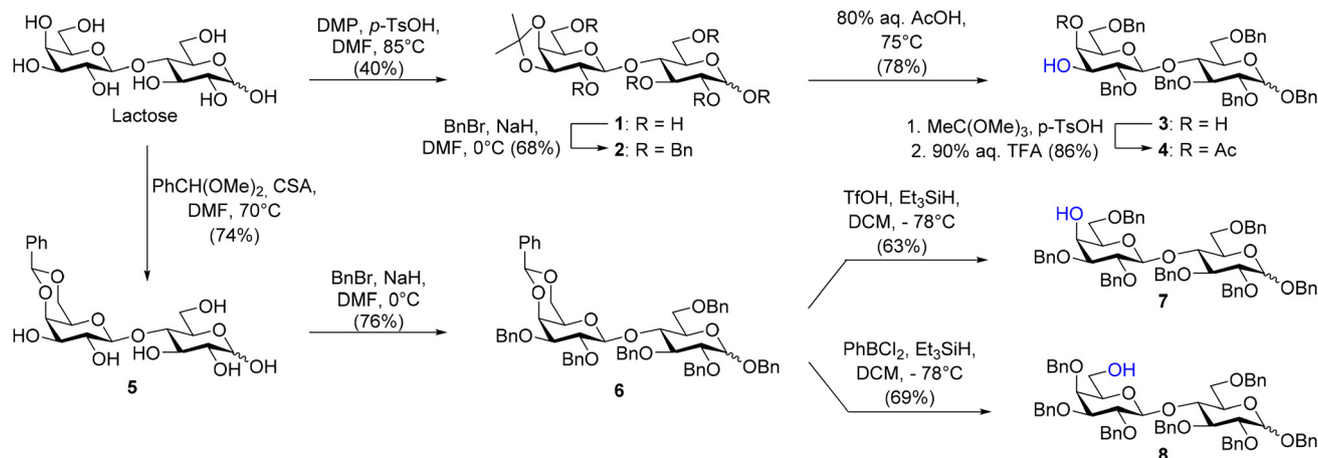
Results and discussion

Synthesis of 3'-, 4'- and 6'-GL

The synthesis of 3'-GL, 4'-GL, and 6'-GL started with the regioselective protection of lactose to afford the disaccharide acceptors (please see ESI S3–13† for experimental details). C-3' hydroxyl lactose-derived acceptor 4 needed to construct 3'-GL was achieved in three steps from lactose (Scheme 1). First, lactose was protected with a 3',4'-O-isopropylidene followed by benzylation of the remaining hydroxy groups with benzyl bromide and sodium hydride to obtain 2 in 27% overall yield.^{24,25} The low yield was attributed to the formation of multiple unidentified byproducts during the 3',4'-O-isopropylidene protection. Potential byproducts include the kinetically favored 4',6'-O-isopropylidene.²⁵ Moreover, ring opening of the glucose residue can also result in the formation of additional byproducts.²⁶ The 3',4'-O-isopropylidene was cleaved by heating 2 in 80% aqueous acetic acid to afford the corresponding *cis*-diol 3 in 78% yield. A small sample of 3 was acetylated using Ac₂O in pyridine to confirm the correct regioselectivity of this sequence by NMR. A downfield shift of the C-3' and C-4' protons was observed in the ¹H-NMR compared to 3 confirming the correct regiochemistry (Fig. S11–13†).

The synthesis of the disaccharide acceptors needed to prepare 4'-GL and 6'-GL was achieved from central benzylidene





Scheme 1 Gram scale synthesis of lactose acceptor building blocks (4, 7 and 8).

intermediate **5** (Scheme 1). A 4',6'-*O*-benzylidene acetal was installed on lactose, followed by benzylation of the remaining hydroxy group using benzyl bromide and sodium hydride to afford **6** in a moderate yield of 56% over two steps.²⁷ The 4',6'-*O*-benzylidene was reductively opened to afford either the C-4' hydroxyl or C-6' hydroxyl derivative **7** and **8**, respectively. The regioselectivity of the reductive ring opening depends on the (Lewis) acid used during the reaction. Triflic acid (TfOH) in combination with triethyl silane as the reducing agent provided the C-4' hydroxyl **7** in 63% yield. In contrast, the use of dichlorophenylborane in combination with the same reducing agent provided the C-6' hydroxyl **8** in 69% yield (Scheme 1).²⁸ A small sample of the compound **7** and **8** was acetylated to determine the regioselectivity. As expected, a downfield shift of the C-4' and C-6' protons was observed in the ¹H-NMR compared to **7** and **8**, respectively (Fig. S31–33 and Fig. S45–47†).

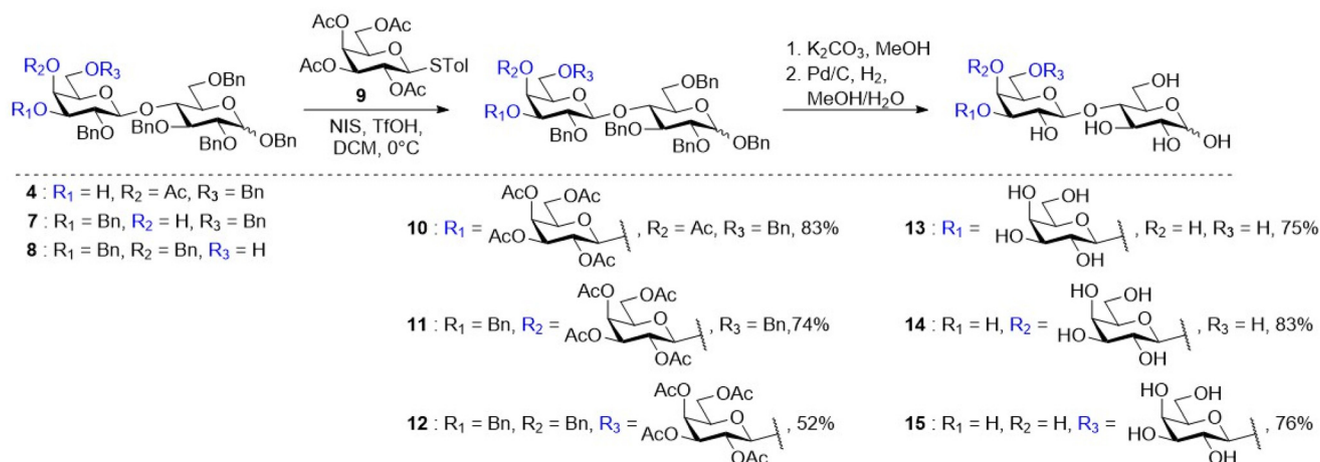
With gram quantities of the disaccharides available, the synthesis of galactose-derived thioglycoside donor **9** was achieved in one step from the commercially available β-D-galactose pentaacetate using *p*-thiocresol and boron trifluoride etherate (BF₃·OEt₂).²⁹ Regioselective glycosylation of acceptor **3** was attempted by treating *cis*-diol **3** with thiogalactoside donor **9** using the NIS/TfOH promotor system similar to the report of Craft and Townsend.^{30,31} Unfortunately, thin layer chromatography (TLC) analysis indicated the formation of numerous side products and it was therefore not possible to isolate the protected 3'-GL (**16**) (Scheme S1†) in good yield and purity. Since glycosylation of diol **3** was unsuccessful, another strategy was pursued. The *cis*-diol was subjected to a simple two-step, one-pot procedure to selectively protect the axial C-4' hydroxyl with an acetyl ester.³⁰ In the first step, **3** was protected with trimethylorthoacetate, followed by an orthoester rearrangement using 90% aqueous TFA to afford **4** in a yield of 86% (Scheme 2). With this acceptor (**4**), TfOH/NIS promoted glycosylation with glycosyl donor **9** resulted in the successful formation of **10** in a yield of 83% with full β-selectivity (Fig. S22†). Glycosylation of the C-4' acceptor **7** was also achieved using donor **9** via a TfOH/NIS promoted glycosylation resulting in

the formation of **11** in a good yield of 74% with full β-selectivity (Fig. S38†). Finally, glycosylation of acceptor **8** with donor **9** under the same conditions afforded **12** in a moderate yield of 52% with full β-selectivity (Fig. S53†). This moderate yield is the result of byproduct formation during the glycosylation reaction. This byproduct was identified as the acetylated acceptor which is presumably formed by orthoester formation with the activated donor followed by rearrangement. Since acetylated acceptor (**19**) was already synthesized to prove the correct regioselectivity of **8** (*vide supra*), the ¹H-NMR spectra of **19** and the byproduct obtained in the glycosylation of 6'-GL were compared (Fig. S48†). This indeed indicated that the obtained byproduct was the acetylated acceptor **19**. All three molecules were deprotected in a two-step process. Deacetylation was achieved by treatment with potassium carbonate in methanol, followed by hydrogenolysis using Pd/C and hydrogen gas to afford 3'-GL (**13**), 4'-GL (**14**) and 6'-GL (**15**) in 75%, 83% and 76% over two steps, respectively (Scheme 2). Overall, 1.42 g of 3'-GL (11% yield, seven steps, 95.4% purity), 2.18 g of 4'-GL (22% yield, six steps, 95.0% purity) and 2.25 g of 6'-GL (15% yield, six steps, 98.1% purity) were obtained. The GLs were subsequently used to evaluate their effect on infant microbiota composition by performing *in vitro* fermentations in infant faecal microbiota.

The effects of 3'-GL, 4'-GL and 6'-GL on infant faecal microbiota *in vitro*

To determine the effect of the galactosylactoses on healthy human infant intestinal microbiota, the TNO i-screen model was used.³² To this end, the faecal microbiota from formula- or breastfed infants were incubated with pure samples of 3'-GL, 4'-GL and 6'-GL (18 h) and compared to an untreated control. At the end of the experiment, the effects of the GLs on the composition of the microbiota was established (see ESI, S15 and 16† for details). An overview of the impact of the different GLs on the top 25 most abundant microbiota taxa can be found in Fig. S1.† Overall, the GLs appear to enhance the abundance of *Bifidobacterium* species and reduce





Scheme 2 The glycosylation and deprotection scheme used for the synthesis of 3'-, 4'-, and 6'-GL.

Bacteroides. DESeq analysis of the data shows that each GL stimulates a certain pattern of bifidobacterial species, with 4'-GL stimulating the most broad pattern of *Bifidobacterium* (Fig. 2 and 3). In the formula-fed infant microbiota, 4'-GL was most bifidogenic; enhancing levels of *B. longum*, *B. breve*, *B. pseudocatenulatum* and *B. kashiwanohense*. 6'-GL increased the abundance of *B. longum* and *B. breve*. 3'-GL was the only GL significantly stimulating the growth of *B. bifidum*, in addition to increasing *B. longum* and *B. breve*. 4'-GL showed the highest overall bifidogenic potential, followed by 6'-GL and 3'-GL. All GLs significantly reduced *Clostridium butyricum* levels and selectively decreased certain members of the genus *Bacteroides* (Fig. 2).

In the breastfed infant microbiota, 6'-GL and 4'-GL were slightly more bifidogenic compared to 3'-GL. All GLs clearly stimulated the growth of *B. longum*. 3'-GL and 6'-GL enhanced

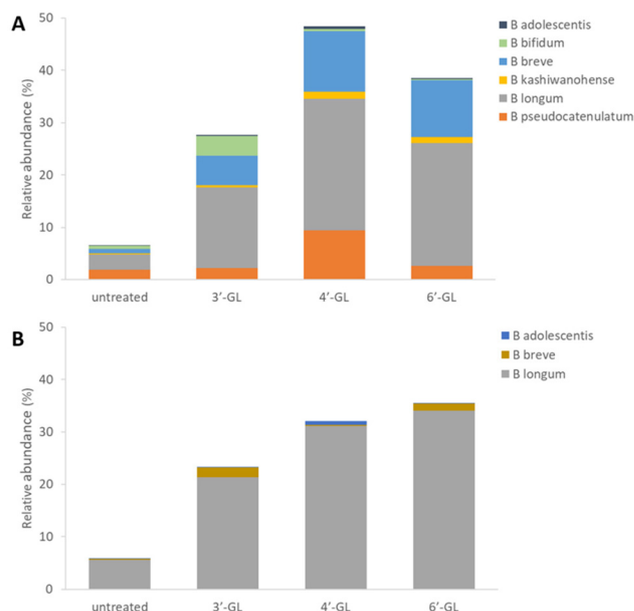


Fig. 3 Bar graphs showing the relative abundances of the different *Bifidobacterium* species in infant microbiota after exposure to 3'-GL, 4'-GL or 6'-GL (panel A: formula fed infants, panel B: breastfed infants).

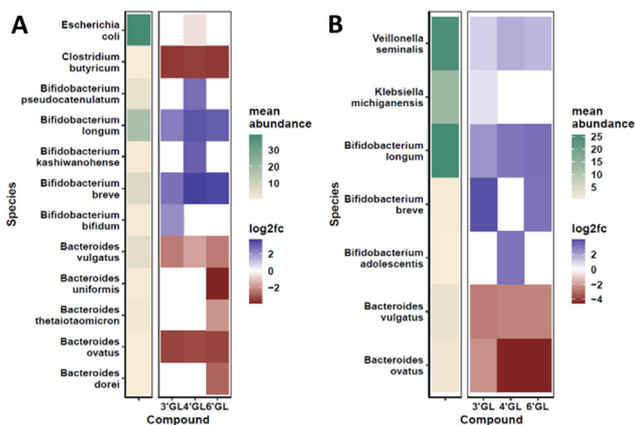


Fig. 2 Visual representation of DESeq2 analysis on the effects of 3'-GL, 4'-GL and 6'-GL on the composition of the infant microbiota (panel A: formula fed infants, panel B: breastfed infants). The microbial taxa represented here are the ones that significantly changed in abundance after exposure to the selected compounds compared to the untreated control.

B. breve, whereas 4'-GL showed a minor increase in *B. adolescentis*. Similar to the results for the formula-fed infant microbiota, the GLs decreased the abundance of certain members of the genus *Bacteroides* (Fig. 2 and 3).

A bifidogenic effect for some of the GLs has been described before. Li *et al.* found that 3'-GL and 6'-GL purified from GOS could enhance *Bifidobacterium* growth *in vitro* in adult faecal microbiota samples.³³ In that study, microbiota changes were identified at genus level, using fluorescent *in situ* hybridization (FISH). With that technique, effects at bacterial strain level could not be detected. To our knowledge, our study is the first to investigate the impact of all individual galactosyllactoses on infant microbiota at bacterial strain level.



To characterize the *B. longum* species into even more detail, qPCR analyses were performed quantifying the levels of its subspecies *B. longum* subsp. *longum* and *B. longum* subsp. *infantis* in the microbiota samples. The different GLs all stimulated *B. longum* subsp. *longum* and *B. longum* subsp. *infantis* to a similar extent in the formula-fed infant microbiota (the relative abundances of both subspecies had roughly a 1 : 1 ratio). In the breastfed infant microbiota, *B. longum* subsp. *longum* was enhanced by the GLs slightly more than *B. longum* subsp. *infantis* (data not shown).

The slight differences we found in bifidogenicity of the GLs between microbiota from infants that were previously formula-fed or breastfed, could in part also be driven by the nutrition received by the infants. In human milk, the levels of 6'-GL are higher early in lactation as compared to 3'-GL and 4'-GL.^{34,35} On the other hand, infant formula in The Netherlands is often supplemented with galacto-oligosaccharides (GOS). GOS are a complex mixture of oligosaccharides also containing GLs,¹⁹ with 4'-GL being the most abundant.³⁶ Thus, the breastfed infant microbiota has seen more 6'-GL as compared to the other GLs and the formula-fed microbiota has seen relatively more 4'-GL. In response, the microbiota may have adapted its composition and/or metabolism to efficiently ferment these respective GLs.

Upon fermentation of substrates, the microbiota can produce different metabolites, such as short-chain fatty acids (SCFA). Fig. 4 shows the levels of SCFA produced by the infant microbiota during fermentation of the GLs. All GLs increased the total concentration of SCFA in the supernatant of the microbiota fermentations, with 4'-GL and 6'-GL showing a significant increase as compared to the control (Fig. 4). This increase was mainly driven by higher levels of acetic acid produced upon fermentation of the GLs (Fig. 4). This is consistent with other studies reporting that galacto-oligosaccharides can enhance acetic acid production by microbiota *in vitro*,³⁷ as well

as in clinical trials in infants³⁸ and adults.³⁹ The increased levels of acetic acid found upon fermentation of the GLs probably reflects their enhancement of *Bifidobacterium* species growth, as these are known producers of acetic acid.⁴⁰ SCFA have been found to play a role in intestinal development (serving as energy source for intestinal cells), but also to influence immune function, energy metabolism and gut-brain communication.⁴¹ Acetic acid production in particular has been shown to be important for cross-feeding between *Bifidobacterium* and butyrate-producing bacteria; and potentially also for reducing overproduction of gas in the infant intestine.⁴⁰ Thus, SCFA are an important factor in infant gut- and overall health.

Overall, our results show a clear bifidogenic effect of all the GLs, with 4'-GL and 6'-GL having a slightly stronger effect as compared to 3'-GL. This indicates that the GLs play an important role in the microbiota-shaping effect of GOS. Future development of GOS ingredients which are further enriched in GL-content may hold potential for enhanced bifidogenicity.

Conclusion

A chemical synthetic approach was developed for the gram-scale chemical synthesis of 3'-, 4'- and 6'-GL. These GLs were obtained in moderate to good overall yields and with very high purity. The effects of these compounds on the composition of infant microbiota was investigated. These GLs were found to enhance *Bifidobacterium* abundance in infant faecal microbiota as tested by *in vitro* fermentation and supported SCFA production. We expect that the gram-scale availability of GLs enables further investigation of their functionality in other, potentially more physiologically relevant, *in vitro* model systems and/or in *in vivo* studies.

Conflicts of interest

There are no conflicts to declare.

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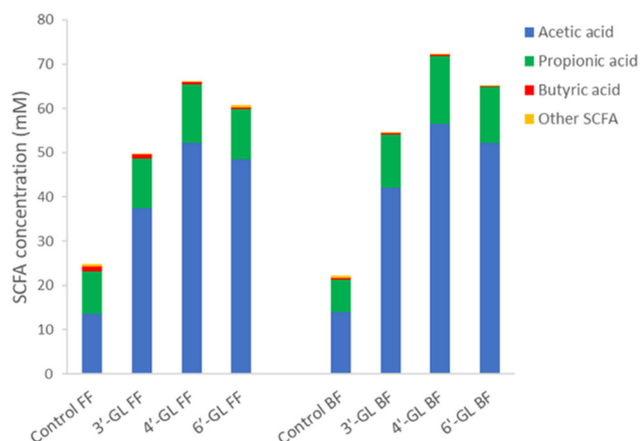


Fig. 4 SCFA concentrations in the supernatant of infant microbiota fermentations. "Other SCFA" include lactic acid, succinic acid, isobutyric acid, 2-methylbutanoic acid, isovaleric acid and valeric acid. FF = formula fed infant microbiota, BF = breastfed infant microbiota.



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