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Loop engineering of an α-1,3/4-L-fucosidase for improved synthesis of human milk oligosaccharides

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Graphical abstract

Highlights

- Targeted loop engineering of a GH29 fucosidase for improved transfucosylation
- 3-fold yield increase in the enzyme’s conversion efficacy on the fucosyl donor
- Enzymatic synthesis of fucosylated human milk oligosaccharides LNFP II and LNFP III

Abstract

The α-1,3/4-L-fucosidases (EC 3.2.1.111; GH29) BbAfCB from Bifidobacterium bifidum and CpAfC2 from Clostridium perfringens can catalyse formation of the human milk oligosaccharide (HMO) lacto-
N-fucopentaose II (LNFP II) through regioselective transfucosylation of lacto-N-tetraose (LNT) with 3-fucosyllactose (3FL) as donor substrate. The current work exploits structural differences between the two enzymes with the aim of engineering BbAfcB into a more efficient transfucosidase and approaches an understanding of structure-function relations of hydrolytic activity vs. transfucosylation activity in GH29. Replacement of a 23 amino acids long α-helical loop close to the active site of BbAfcB with the corresponding 17-amino acid α-helical loop of CpAfc2 resulted in almost complete abolishment of the hydrolytic activity on 3FL (6000 times lower hydrolytic activity than WT BbAfcB), while the transfucosylation activity was lowered only one order of magnitude. In turn, the loop engineering resulted in an α-1,3/4-L-fucosidase with transfucosylation activity reaching molar yields of LNFP II of 39±2% on 3FL and negligible product hydrolysis. This was almost 3 times higher than the yield obtained with WT BbAfcB (14±0.3%) and comparable to that obtained with CpAfc2 (50±8%). The obtained transfucosylation activity may expand the options for HMO production: mixtures of 3FL and LNT could be enriched with LNFP II, while mixtures of 3FL and lacto-N-neotetraose (LNnT) could be enriched with LNFP III.

**Abbreviations**

2'FL: 2’-fucosyllactose; 3FL: 3-fucosyllactose; A:D: acceptor-to-donor ratio; BbAfcB: α-1,3/4-L-fucosidase from *Bifidobacterium bifidum*; BiAfcB: α-1,3/4-L-fucosidase from *Bifidobacterium longum* subsp. *infantis*; CpAfc2: α-1,3/4-L-fucosidase from *Clostridium perfringens*; Fuc: fucose; GH: glycoside hydrolase; Gal: galactose; Glc: glucose; GlcNAc: N-acetylglucosamine; HMO: human milk oligosaccharide; LNT: lacto-N-tetraose; LNnT: lacto-N-neotetraose; LNFP: lacto-N-fucopentaose; Sia: sialic acid; WT: wild type.

**Keywords:** fucosidase, GH29, human milk oligosaccharides, protein engineering, transglycosylation
Introduction

Human milk oligosaccharides (HMOs) are soluble complex glycans which are present in human milk at concentrations of 5-15 g/L, often exceeding the concentration of protein thus being the third most abundant component in human milk. In the human colostrum, concentrations are even higher [1]. In contrast, HMOs are virtually absent from bovine milk, which is the basis of infant formula. Approx. 1% of the HMOs are adsorbed, while the majority is either metabolized by the gut microbiota of the infant or excreted in the faeces [2, 3]. HMOs function as prebiotics and antimicrobial agents in the gut of breastfed infants, and they further protect the infant against pathogens by functioning as soluble decoy receptors for pathogen adhesion as well as through a number of immunomodulating effects. HMOs may also be involved in infant brain development [1]. No single HMO has all these effects at once, suggesting why as many as 200 different HMO structures have been identified in human milk [4, 5].

Five monosaccharides, i.e. galactose (Gal), glucose (Glc), N-acetylglucosamine (GlcNAc), fucose (Fuc), and sialic acid (Sia) derivative N-acetyl-neuraminic acid, form the HMO building blocks. All HMO molecules have lactose (Gal-β1,4-Glc) at the reducing end can be elongated in β1,3- or β1,6-linkages by two different disaccharide moieties, namely Gal-β1,3-GlcNAc (type 1) or Gal-β1,4-GlcNAc (type 2). The HMO backbone can be further modified with Sia and/or Fuc substitutions [1, 5]. Among them, the fucosylated species are the most abundant – at least in breastmilk of approx. 80% of the population since the degree and type of HMO fucosylation are linked to the secretor and Lewis blood status group of the mother [6].

Several routes to production of HMOs in vitro exist, one of them being the use of glycosidases catalysing transglycosylation [7, 8, 9]. Advantages of glycosidases include easy expression, robustness, and the option of using inexpensive naturally occurring donor substrates. Indeed, the α-1,3/4-L-fucosidase BiAfCB from Bifidobacterium longum subsp. infantis (EC 3.2.1.111) was recently shown to catalyse transfucosylation of the HMO core structure lacto-N-tetraose (LNT)
with another HMO, namely 3-fucosyllactose (3FL), as fucosyl donor leading to formation of lacto-N-fucopentaose II (LNFP II) [10] (Figure 1). For this WT enzyme, transfucosylation yields on 3FL ranged from 6% to 12% depending on the substrate concentration, but through elaborate engineering of BiAfcB mutants transfucosylation yields on 3FL of up to 60% were achieved [10, 11]. We recently studied two WT α-L-1,3/4-fucosidases (EC 3.2.1.111) – BbAfcB from Bifidobacterium bifidum and CpAfc2 from Clostridium perfringens: In terms of their transfucosylation potential CpAfc2 was superior to the BbAfcB producing molar transfucosylation yields of up to 39% (on the donor 3FL) in production of LNFP II [12]. These α-L-fucosidases belong to the glycoside hydrolase family 29 subfamily B (GH29B) which comprises retaining α-L-fucosidases with high specificity for branched α-1,3/4-fucosylations [13]. In turn, GH29B α-1,3/4-L-fucosidases have also been shown to possess high regioselectivity in transfucosylation [10, 12].

When comparing the transfucosylation and hydrolytic activities of BbAfcB and CpAfc2 it was evident that CpAfc2 had markedly higher transfucosylation activity than BbAfcB using 3FL and LNT for formation of LNFP II, and that BbAfcB in turn had significantly higher hydrolytic activity on 3FL than CpAfc2 [12]. Unfortunately, C. perfringens is a potential pathogen, so the use of an enzyme derived from this organisms for production of HMOs for infant formulae may be controversial. In contrast, B. bifidum is considered a beneficial probiotic microbe. Consequently, the current work set out to exploit any structural differences between the two GH29B α-1,3/4-L-fucosidases with the aim of engineering of BbAfcB for improving its transfucosylation activity and/or reducing its hydrolytic activity. Comparison of homology models of BbAfcB and CpAfc2 with the crystal structure of the well-studied BiAfcB, reveal that BbAfcB and CpAfc2 appear similar in terms of active site structure and substrate-interacting residues. However, the models of the active site region suggest that an α-helical loop on the side of the active site constitutes a major difference between BbAfcB and CpAfc2. Since the transsialylation yield of a sialidase from Trypanosoma rangeli could be significantly improved by replacing a loop near the active site with the corresponding loop of a native transsialidase from T. cruzi [14], a similar approach was pursued here: The identified α-helical loop in
BbAfcB was replaced with that of CpAfc2, hypothesizing that the shape of the loop in CpAfc2 provide better shielding of the active site from the aqueous environment and that this feature could be transferred to BbAfcB by loop engineering.

**Materials & Methods**

*Chemicals*

The fucosylated oligosaccharides 3-fucosyllactose (3FL), lacto-N-tetraose (LNT), lacto-N-neotetraose (LNnT), and lacto-N-fucopentaose V (LNFP V) were purchased from Elicityl Oligotech (Crolles, France). All other chemicals were purchased from Sigma-Aldrich (Steinheim, Germany).

*Homology modelling*

Structure homology models were prepared using the *Homology Modeling* function of YASARA Structure, version 16.9.23 (YASARA Biosciences GmbH, Vienna, Austria) [15]. The program identified automatically the following template structures in structural databases: 4ZRX, 3UES, 3MO4, 2BER, 2X0Y for BbAfcB and 4ZRX, 5K9H, 3EYP, 3RB5, 2OZN for CpAfc2. For each of the templates up to 5 homology models were prepared by the program, which afterwards were automatically ranked. Based on the ranking a hybrid model was prepared by the program, which finally was equilibrated by molecular dynamics simulation using the YASARA macro md_refine without changes [16]. The refined models with the lowest energy were quality checked using QMEAN4 [17] and subsequently used for comparison with the BiAfcB structure 3UET. All figures showing 3D structures were prepared using PyMol (The PyMOL Molecular Graphics System, Version 1.1 Schrödinger, LLC). Sequence alignments were made with MUSCLE 3.8 [18].

*Cloning, expression and purification of α-L-fucosidases*
The sequence of α-L-1,3/4-fucosidase BbAfCB from *Bifidobacterium bifidum* JCM 1254 (GenBank BAH80310.1; EC 3.2.1.111) was slightly truncated and codon-optimized for expression in *Escherichia coli* previously [12]. The resulting pET22b+/bbafcb plasmid was used as a template for deletion of the encoded loop 759-AAYNDGVSDKPKPQMAPDGKL-781 and introduction of the new loop DIEKMKERENPTYLNNG originating from α-L-1,3/4-fucosidase CpAfC2 from *Clostridium perfringens* ATCC 13124 (GenBank ABG83106.1; EC 3.2.1.111) [12, 19] by PCR using the forward primer 5’-AACCCGACCTATCTGAACAACGGCAGCATGAGCAGCGTGCTGAGCGAG-3’ and the reverse primer 5’-TTCACGTTCTTCTTTTTCGATATCTTGCGGGCTCCATTCGGTCTGACGCGCC-3’ where underlined nucleotides correspond to the new loop sequence (Table A.1; Table A.2). The plasmid template was removed by DpnI digestion. After purification using Illustria GFX PCR DNA and Gel Band Purification kit (GE Healthcare, Chicago, IL, USA), the PCR product was phosphorylated using T4 polynucleotide kinase (Thermo Fisher Scientific, Waltham, MA, USA) at 37°C during 1 hour and ligated overnight at 16°C using T4 DNA ligase (Thermo Fisher Scientific, Waltham, MA, USA). The ligated PCR product was then transformed into competent *E. coli* DH5α cells, prepared with the Mix & Go *E. coli* Transformation kit (Zymo Research, Irvine, CA, USA). Plasmid was extracted using QIAprep Spin Miniprep kit (Qiagen, Hilden, Germany) and checked by sequencing (Macrogen Europe, Amsterdam, the Netherlands).

Expression and purification of *BbAfCB*, *BbAfCBmut*, and *CpAfC2* were carried out in *E. coli* BL21 (DE3) as described previously [12]. Protein purity was confirmed by SDS-PAGE and protein concentration was determined with the BCA Protein Assay Kit (Thermo Fisher Scientific, Waltham, MA, USA) using bovine serum albumin (BSA) standards.

*Hydrolytic activity and HPAEC-PAD analysis*

Hydrolytic activity was determined for *BbAfCB*, *BbAfCBmut*, and *CpAfC2* using 0.1 mM 3FL in 50 mM buffer (acetate buffer pH 5.5 for *BbAfCB* and *BbAfCBmut* according to the pH optimum of *BbAfCB* established by Ashida et al. [20], and phosphate buffer pH 7.0 for *CpAfC2* according to the pH
optimum for CpAfc2 [19]) and enzyme concentrations between 0.005 µM and 5 µM. Negative controls without enzyme were included. All reactions took place at 40°C, were monitored for up to 20 min (180 min for \textit{BbAfcB}_{mut}), and stopped by mixing a sample 1:1 with 1 M Na$_2$CO$_3$. The amounts of substrate and hydrolysis products were quantified by high-performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD). Specific hydrolytic activity was defined as µmol of product released per minute (U) per µmol of enzyme.

Enzyme stability at 40°C was determined by incubation of the fucosidases in the reaction buffer at 40°C followed by assessment of their remaining hydrolytic activity after 0, 1, 3, 5, 12, and 24 hours of incubation as described above.

HPAEC-PAD analysis was carried out on a Dionex ICS3000 system (Dionex Corp., Sunnyvale, CA, USA) using a CarboPac™ PA1 (4 mm x 250 mm) analytical column equipped with a CarboPac™ PA1 (4 mm x 50 mm) guard column (Dionex Corp., Sunnyvale, CA, USA) and a flow rate of 1 mL/min at 25°C. The eluent system comprised MilliQ water (A), 500 mM NaOH (B), and 500 mM NaOAc with 0.02% (w/v) NaN$_3$ (C). After enzymatic hydrolysis, 3FL was quantified by isocratic elution at 90:10:0 (% A:B:C) for 20 min using external standards of L-fucose, lactose, and 3FL. Strongly retained anions were washed from the column by elution with 10:10:80 (% A:B:C) for 5 min followed by re-equilibration of the column at 90:10:0 (% A:B:C) for 10 min.

\textit{Transfucosylation activity and LC-ESI-MS analysis}

For transfucosylation of LNT and LNnT, 3FL was used as donor substrate for reactions catalysed by \textit{BbAfcB}, \textit{BbAfcB}_{mut} and \textit{CpAfc2}. Enzyme concentration was 0.5 µM and reaction took place at 40°C for up to 24 hours in 100 mM buffer (acetate buffer pH 5.5 for \textit{BbAfcB} and \textit{BbAfcB}_{mut}, phosphate buffer pH 7.0 for \textit{CpAfc2}). Reactions were terminated by heating at 99°C for 15 min. For reactions with an acceptor-to-donor ratios (A:D) of 10, donor substrate (3FL) concentration was 10 mM and acceptor substrate (LNT or LNnT) concentration was 100 mM. For reactions with an A:D of 1, concentrations of donor and acceptor substrates were 20 mM. Additional experiments were conducted with \textit{BbAfcB}
and \( \text{BbAfcB}_{\text{mut}} \) at the same conditions (reaction time up to 60 min) using A:Ds of 2 (10 mM 3FL and 20 mM LNT) and 5 (20 mM 3FL and 100 mM LNT). Transfucosylation products were identified and quantified by liquid chromatography electrospray ionization mass spectrometry (LC-ESI-MS) using an external LNFP V standard. Transfucosylation yields were calculated as molar yields based on the donor substrate 3FL, i.e. the percentage of LNFP formed compared to the initial 3FL concentration. Initial transfucosylation rates were calculated as mM LNFP formed per min of reaction over the time where the product formation curve was linear (2 min for \( \text{BbAfcB} \), 5 min for \( \text{CpAfc2} \), and 20-30 min for \( \text{BbAfcB}_{\text{mut}} \)).

Identification and quantification of LNFP transfucosylation products were performed by LC-ESI-MS on an Amazon SL iontrap (Bruker Daltonics, Bremen, Germany) coupled to an UltiMate 3000 UHPLC from Dionex (Sunnyvale, CA, USA) equipped with a porous graphitized carbon column (Hypercarb PGC, 150 mm × 2.1 mm, 3 µm; Thermo Fisher Scientific, Waltham, MA, USA) as described previously [12]. LNFP isomers LNFP II, LNFP III, and LNFP V were identified from MS\(^2\) fragmentation patterns [21].

**Statistics**

One-way ANOVA for determination of statistical significance was performed with JMP®, version 13.0.0 (SAS Institute Inc., Cary, NC, USA). Statistical significance was established at \( p < 0.05 \).

**Results**

**Homology modelling**

The homology models of \( \text{BbAfcB} \) and \( \text{CpAfc2} \) were inspected visually and their quality was checked using the QMEAN4 Z-score [17]. The model for \( \text{BbAfcB} \) showed a dimeric structure, but since there was no significant difference in between the two monomers it was decided to continue with only one of them. The catalytic domains for both models showed the expected \((\beta/\alpha)_8\)-barrel, though one of
the β-strands was not modelled as such (Figure A.1). However, aligning the models with the crystal structures of \textit{BiAfcB} (3UET) [22] and another GH29B member, namely α-L-1,3/4-fucosidase BT2192 from \textit{Bacteroides thetaiotaomicron} (4OZO) [23], revealed that even in the crystal structure this part of the barrel was not a perfect β-strand; thus, this could be a natural feature of these enzymes. The QMEAN4 Z-scores for the full-length models of \textit{BbAfcB} and \textit{CpAfc2} were -2.65 and -2.61, respectively (Figure A.2). Deleting the C-terminal non-catalytic domain for the quality check improved the Z-score for \textit{BbAfcB} only marginally (-2.51), but for \textit{CpAfc2} the score significantly improved to -0.62 (Figure A.2). The low quality score for the \textit{BbAfcB} model might be caused by the long single α-helices present, which especially influence the packing check negatively. However, since the model of the catalytic domain for \textit{CpAfc2} was of good quality, the models were used to identify potential targets to engineer \textit{BbAfcB} for improved transglycosylation performance.

Based on inspection of the homology models and comparison to the well-studied crystal structure of \textit{BiAfcB}, it was evident that they share all substrate-interacting residues in terms of position in the three-dimensional structure (Figure A.1; Figure A.3; Table A.3). However, \textit{CpAfc2} featured an α-helical loop region on the side of the active site, approx. 13 Å from the substrate. In \textit{BbAfcB}, the major part of this α-helical loop was further away and more disordered (Figure 2) and in \textit{BiAfcB} it was even found to be a large disordered structure, because the structure of this loop could not be determined (Figure A.3). The most significant structural difference in the compared structures was this loop region, which was moreover found to be highly variable in the three sequences. Hence, the hypothesis behind the current work was that replacement of the loop sequence in \textit{BbAfcB} with that of \textit{CpAfcB}, would improve the transfucosylation ability of \textit{BbAfcB} by inducing a better shielding of the active site from the aqueous environment. This hypothesis was further supported by inspection of the surface representation of the aligned models, in which it became obvious that especially R266 in the \textit{CpAfc2}-loop might indeed shield the active site of \textit{BbAfcB}, even though there is still room for water molecules in the active site (Figure A.4).
To define the exact parts of the loop in \( Bb \text{AfcB} \) to be exchanged with the one from \( Cp \text{Afc2} \), the homology models were aligned in PyMol and inspected visually. To minimize the risk of losing the correct foldability and also a possible dimerization of \( Bb \text{AfcB} \), the starting and end points of the loops to be exchanged were carefully selected based on backbone alignment. The backbones of the amino acid residues A759 in \( Bb \text{AfcB} \) and D259 in \( Cp \text{Afc2} \) aligned well, so these residues were defined as starting points of the loops to be exchanged (Figure A.1). Although true backbone alignment was not achieved until L799 in \( Bb \text{AfcB} \) and S280 in \( Cp \text{Afc2} \), detailed inspection of the modelled loop structures revealed that G782 of \( Bb \text{AfcB} \) and E276 of \( Cp \text{Afc2} \) aligned well enough, so that these two positions were defined to terminate the loops to be exchanged (Figure A.1). This led to the final definition of the loop 759-AAYNDGVDKVSLKPGQMAPDGKL-781 in \( Bb \text{AfcB} \) to be exchanged with the loop 259-DIEKMKERENPTYLNNG-275 from \( Cp \text{Afc2} \).

**Expression of recombinant α-1,3/4-L-fucosidases**

\( Bb \text{AfcB} \), \( Bb \text{AfcB}_{\text{mut}} \) loop mutant, and \( Cp \text{Afc2} \) were successfully expressed in \( E. \text{coli} \) BL21 (DE3) cells using LB medium. After \( \text{Ni}^{2+} \) affinity chromatography purification and desalting by gel filtration expression levels reached 24, 23, and 38 mg of purified enzyme per litre of culture, respectively. SDS-PAGE analysis showed that all purified extracts were satisfactorily pure (Figure A.5).

**Hydrolytic activity**

The hydrolytic activities of \( Bb \text{AfcB} \) and \( Cp \text{Afc2} \) on 3FL were similar to the previously reported values [12]: 18 U/μmol enzyme for \( Cp \text{Afc2} \) and 59 U/μmol enzyme for \( Bb \text{AfcB} \). However, for \( Bb \text{AfcB}_{\text{mut}} \) the hydrolytic activity on 3FL was practically non-existent: 0.01 U/μmol enzyme, i.e. 6000 times lower than the WT enzyme (Table 1).

**Transfucosylation activity**
Transfucosylation activity of BbAfcBmut, BbAfcB, and CpAfc2 was monitored for 3 hours using 3FL as fucosyl donor and LNT or lacto-N-neotetraose (LNnT) as acceptors. The acceptor-to-donor ratios (A:D) were either 10 (acceptors 100 mM LNT or LNnT and donor 10 mM 3FL) or 1 (20 mM of each substrate). For WT BbAfcB, maximum levels of LNFP II and LNFP III were obtained after 2 minutes of reaction, which was the shortest reaction time measured; after this, product levels decreased to 0 within 2 hours of reaction (Figure 3). At A:D 10, the maximum yield of LNFP II was 14%, whereas it was 3.3% for LNFP III (Figure 3; Table 2). For CpAfc2, the transient maxima were reached after 30 minutes and yields were 50% for LNFP II and 17% for LNFP III at A:D 10 (Figure 3; Table 2). For BbAfcBmut, no transient maxima were observed within 3 hours of reaction (Figure 3). Instead, maximum yields were obtained after 3 hours reaching 39% for LNFP II and 11% for LNFP III on the 3FL donor at A:D 10 (Figure 3, Table 2).

Interestingly, the highest apparent initial transfucosylation rates were obtained with BbAfcB, being approx. 1.5 times higher than those of CpAfc2 at A:D 10 and 2-3 times higher at A:D 1 (Table 2). In comparison, initial transfucosylation rates were approx. one order of magnitude lower for BbAfcBmut compared to the two WT enzymes (Table 2).

Additional samples were taken after 24 hours of reaction in order to investigate the degree of product hydrolysis. For both WT enzymes BbAfcB and CpAfc2, product degradation was complete after 24 hours (Table 2). However, for BbAfcBmut no significant decrease in LNFP II levels were observed, whereas the level of LNFP III did decrease at A:D 10 though not at A:D 1 (Table 2). After incubation at 40°C for 24 hours in reaction buffer, the remaining activities were 77%, 100%, and 81% for BbAfcBmut, BbAfcB, and CpAfc2, respectively. For BbAfcBmut, a half-life of 66 hours was determined, whereas it was 82 hours for CpAfc2.

In addition to LNFP II, small amounts of LNFP V (i.e. the LNFP isomer where Fuc is α1,3-linked to the reducing end Glc moiety of LNT) were detected in the reaction with 3FL and LNT catalysed by BbAfcBmut (Figure A.6). After 3 hours of reaction, the yield of LNFP V was 5% at A:D 10 and 0.6% at
A:D 1 corresponding to 11% and 5% of the total transfucosylation yield, respectively. No LNFP V was detected in the reaction catalysed by CpAfc2, and for WT BbAfcB yields never exceeded 0.5%.

In most cases, significantly higher yields were obtained at A:D 10 compared to A:D 1 (Table 2). To further explore the influence of A:D and substrate concentration, BbAfcB and BbAfcB\textsubscript{mut} were employed at A:D 1, 2, 5, and 10 combining two different donor concentrations (10 mM and 20 mM) with two different acceptor concentrations (20 mM and 100 mM) (Table 3; Figure 4). In this way, the effect of donor concentration, acceptor concentration, and their molar ratio (A:D) on product formation and yield were investigated simultaneously. In all cases, the same enzyme showed similar reaction progress over time independent of the A:D and substrate concentration employed (Figure 4). To clearly investigate the effect of both parameters, initial transfucosylation rates were divided by the initial 3FL concentration (10 or 20 mM). Doing so, almost identical values were obtained for reactions with identical LNT acceptor concentrations (Table 3). In turn, the initial transfucosylation rates increased 4 times for BbAfcB\textsubscript{mut} and CpAfc2 when the acceptor concentration was increased 5 times (Table 3). For BbAfcB this was not the case, as a 5-fold increase in acceptor concentration led to a 1.5-fold increase in initial transfucosylation rate only (Table 3). Similar trends were observed for maximum LNFP II yields obtained within 60 minutes of reaction (Table 3).

**Discussion**

Examples of loop deletions exist as a strategy for improving transglycosylation activity in microbial enzymes with polymeric substrates, e.g. xyloglucan endo-hydrolases (using xyloglucan endo-transglucosylases as template) [24] and α-amylases (using cyclodextrin glucanotransferases as template) [25]. However, point mutations – single or multiple – in and around the active site comprise by far the most pursued strategy for exoglycosidases [26-29], and is also the only protein engineering strategy previously used on GH29 α-L-fucosidases [10, 11, 22, 30]. However, for a sialidase from *Trypanosoma rangeli* it was shown that exchanging a loop of seven amino acids
approx. 10 Å from the active site with the seven amino acids found in the corresponding loop in the native transsialidase from *Trypanosoma cruzi* significantly improved transsialylation yields [14]. The loop mutation mainly reduced the hydrolytic activity, whereas it had little effect on the transsialylation rate [29]. On a more general level, the interest in the flexible loop regions and their impact on enzyme function is increasing and as a consequence engineering of loop regions is becoming a more commonly pursued strategy [31].

Enzymes of the GH29 family have a (β/α)₈ or TIM-barrel-like fold (although one α-helix may be missing), where all the conserved active site residues are on the catalytic βα-loops, i.e. the loops that follow after the β-strands [32, 33]. The active sites of GH29 members exhibit a pocket topology which is common for exoglycosidases [34]. Conservation is strict among the ligand interacting residues inside the active site pocket, but there is large diversity in the overall shape due to variation in the loops surrounding the active site [32]. Crystal structures provided evidence that some of the loops undergo conformational changes upon substrate binding and that at least two of the loops take part in the catalytic process in BiAfcB (Table A.3) [22]. Thus, the loops near the active site could indeed be an interesting target for α-L-fucosidase engineering. The loop targeted in the current work appears highly flexible as it was disordered and not properly crystallized in the crystal structure of BiAfcB (Table A.3) [22]. In the (β/α)₈-barrel structure, mutations in the catalytic βα front loops often do not cause stability issues since they are separated in space by the central barrel structure from the αβ back loops, which confer stability [33, 35]. This separation could give functional versatility in enzymes with (β/α)₈-barrel-like folds, and the (β/α)₈-barrel is therefore considered an ideal scaffold for rational design of novel enzyme activities [33, 35, 36]. Indeed, several reports exist on loop grafting, i.e. replacing βα-loops from one enzyme of the (β/α)₈-barrel fold class with those from another, for transferring substrate specificity and function although not yet for transglycosylation purposes [36, 37].

The loop mutation in the *T. rangeli* sialidase introduced a change in net charge of +3 and it was hypothesized that it altered the water network in the active site [14]. In the current work, a 23-
amino acid α-helical loop with a net charge of 0 was replaced by a 17-amino acid α-helical loop with a net charge of -1. Rather than altering the water network it was hypothesized that this would lead to better shielding of the active site pocket from the surrounding aqueous environment, thus reducing the competing hydrolytic activity. A closed active site topology, a hydrophobic entrance to the active site, or obstruction of catalytic water channels to exclude water from the active site have all been shown to have positive effects on transglycosylation for GH33 sialidases, GH1 β-glucosidases, and GH13 neopullulanase [38-42]. Indeed, the study of the homology models of CpAfc2 and BbAfcB suggested that the α-helical loop in CpAfc2 could have a more shielding effect than the more open and disordered loop found in BbAfcB, although the models indicate that there is still ample room for water molecules in the active site (Figure 2; Figure A.4). While these were indeed models – and the corresponding loop in BiAfcB so flexible that it could not be crystallized [22] – the results emphasize that the exchange of the loop in BbAfcB with that of CpAfc2 did indeed have a major effect on the transfucosylation performance of BbAfcB, mainly by almost completely abolishing its hydrolytic activity while maintaining an initial transfucosylation rate which was approx. one order of magnitude lower than those observed in WT BbAfcB and CpAfc2 (Table 1; Table 2; Table 3). A drop in activity rate is often observed in glycosidases engineered for improved transglycosylation, especially when mutations are in the negative subsite (i.e. the donor binding site) and/or when more than one amino acid is substituted [27, 28]. Exchange of a loop in the T. rangeli sialidase with the corresponding one of T. cruzi transsialidase also reduced the hydrolytic activity significantly, indicating that this strategy – which is similar to the one employed here – particularly targets the undesirable hydrolytic activity [14, 29]. The loop mutation did not influence the expression levels of recombinant BbAfcB and BbAfcBmut in E. coli, which were similar. Enzyme stability at extended reaction times was slightly influenced by the loop mutation: whereas BbAfcB retained full activity after 24 hours of incubation at 40°C, the remaining activity of BbAfcBmut was 77% and thus more similar to that of CpAfc2, which was 81%. However, 77% is an appreciable remaining activity and it should be feasible to run the reaction...
past 24 hours in order to obtain higher transfucosylation yields with \textit{Bb}Afc\textsubscript{Bmut} at A:D 1. At A:D 10, maximum yields were obtained within 24 hours of reaction (Table 2).

For glycosidase-catalyzed transglycosylation, product maxima are often transient due to hydrolysis [8]. Interestingly, despite having the highest initial transfucosylation rate, \textit{Bb}Afc\textsubscript{B} gave the lowest transfucosylation product yields. From this, it is evident that transfucosylation rate is not the only important factor. Indeed, when employing glycosidases for transglycosylation, hydrolytic rates on substrate as well as on product are just as important as the transglycosylation rate in determining the maximum transglycosylation yield (Table 2; Figure 3). The fact that maximum product levels were obtained at the shortest measured reaction time with \textit{Bb}Afc\textsubscript{B} (Figure 3) is indeed linked to the fact that this enzyme had significantly higher hydrolytic activity than \textit{Bb}Afc\textsubscript{Bmut} and \textit{Cp}Afc\textsubscript{2} (Table 1). For \textit{Cp}Afc\textsubscript{2} the transient maximum occurred at a later stage (30 min; Figure 3), whereas the product hydrolysis was limited for \textit{Bb}Afc\textsubscript{Bmut}, which had very low hydrolytic activity on 3FL compared to the two WT enzymes (Figure 3; Table 1). As discussed above, the transfucosylation activity was also approx. one order of magnitude lower, but due to the dramatic decrease in product hydrolysis, appreciable levels of LNFP II and LNFP III were reached with \textit{Bb}Afc\textsubscript{Bmut} (Table 2). While \textit{Cp}Afc\textsubscript{2} produced even higher yields at A:D 10, the yields were similar with \textit{Cp}Afc\textsubscript{2} and \textit{Bb}Afc\textsubscript{Bmut} at A:D 1 (Table 2).

It is well established that both substrate concentration and A:D affect transglycosylation rates and yields [8]. The transfucosylation data on LNT with 3FL as donor catalyzed by \textit{Bb}Afc\textsubscript{B} or \textit{Bb}Afc\textsubscript{Bmut} suggested that the progress and mechanism of the reactions were independent of A:D and substrate concentration (Figure 4). Dividing initial transfucosylation rates and transfucosylation yields by the initial donor substrate concentration revealed similar values for reactions with identical acceptor concentrations, indicating that the effect of donor substrate concentration was linear in this range (Table 3). Analogously, there was a clear effect of the acceptor concentration: For \textit{Bb}Afc\textsubscript{Bmut} a 4-fold higher initial rate and a 5.5-fold higher yield was obtained when the acceptor concentration was increased 5 times (Table 3). A similar trend was observed for \textit{Cp}Afc\textsubscript{2}, although the increase in
yield was only 3-fold (Table 3). For WT BbAfcB, increases in rate and yield were only 1.5-fold (Table 3). Most likely, initial transfucosylation rates were heavily influenced by the large hydrolytic activity of this enzyme, and the effect of A:D was therefore not as evident (Table 1; Table 3). Indeed, the transient maximum was passed within the first two minutes of reaction, and the initial transfucosylation rate and maximum yield may therefore be higher than established here (Figure 4).

The transfucosylation activity was tested on two different HMO acceptor substrates of equal size, namely LNT and LNnT which differ only in the linkage between the non-reducing Gal moiety and GlcNAc (β1,3 in LNT and β1,4 in LNnT; Figure 1). Both were used as substrates for transfucosylation with 3FL, resulting in formation of HMO pentaose structures LNFP II and LNFP III, respectively, as identified by LC-ESI-MS/MS (Figure 1). However, for WT α-1,3/4-L-fucosidases BbAfcB and CpAfc2 initial transfucosylation rates were 4-5 times higher on LNT compared to LNnT (Table 2). For BbAfcBmut the difference was more moderate, the initial rate being only 2 times higher for LNT (Table 2). Similarly, maximum yields were higher for LNFP II than for LNFP III, the difference between the enzymes being less distinct (Table 2). The slightly larger preference for LNnT observed in BbAfcBmut compared to the WT α-1,3/4-L-fucosidases could be linked to the less strict regioselectivity observed for the engineered enzyme in the transfucosylation of LNT, where formation of minor amounts of LNFP V was observed. Although the crystal structure of orthologue BiAfcB indicates that the reducing end of LNFP II protrudes from the enzyme, while the non-reducing end Gal moiety is buried in the active site pocket (Figure 2; Figure A.1) [22], the formation of LNFP V indicates that BbAfcBmut must be able to accommodate LNT in a reverse manner. This may be a result of the loop engineering since only negligible amounts of LNFP V were observed in reactions catalyzed by WT BbAfcB. However, no LNFP V was observed in reactions catalysed by CpAfc2, indicating that it was not the loop as such, but rather its effect on BbAfcBmut. LNFP V is present in human milk, albeit only in very small amounts [43].

The only previously reported engineering of BbAfcB is its transformation into an α-L-fucosynthase, giving an LNFP II yield of 41% when using 34 times more enzyme than in the current
work [22]. While this strategy is appealing due to its simple, generic nature, especially α-glycosynthases struggle with low donor substrate stability [22, 44]. GH29B α-1,3/4-L-fucosidases are particularly appealing in terms of engineering for improved transfucosylation activity – and for use in transfucosylation in general – since they appear to maintain their high regioselectivity in the transfucosylation reaction, while the GH29A α-L-fucosidases often do not [10, 12, 22]. So far, the only other example of engineering of a GH29B α-1,3/4-L-fucosidase is the extensive work on BjAfcB [10, 11]. Together with these results, the current work emphasizes the potential of protein engineering for improved transfucosylation performance among GH29B enzymes, as well as for targeted loop engineering in general. In addition, the work highlights how the competition between transglycosylation and hydrolysis rates governs glycosidase-catalysed transglycosylation and why reduction of hydrolytic activity is an important strategy in transglycosidase engineering.

The use of glycosidase-catalysed transglycosylation represents one strategy for in vitro production of HMOs [7, 8]. An alternative strategy for in vitro HMO production is the use of prokaryotic fucosyltransferases; however, their expression is challenging and their use requires a multi-enzyme setup for regeneration of the required expensive substrate GDP-L-fucose [9]. For fairly simple HMO structures, fermentation of engineered Escherichia coli appears to be a viable strategy. For instance, HMOs including 2′-fucosyllactose (2′FL), 3FL, LNT, and LNnT have been produced by engineered E. coli with yields generally around 10 g/L [7, 45-47]. However, for more complex structures such as LNFPs yields are generally much lower and only LNFP I and LNFP III have been reported as fermentation products with yields below 1 g/L [48, 49]. A process combining fermentation products with regioselective enzymatic transfucosylation catalysed by α-L-fucosidases could lead to a mixture of 3FL and LNT enriched with LNFP II, which is the seventh most abundant HMO in general and the second most abundant HMO in non-secretor milk [6].

Being of lactic acid bacterial origin, the use of engineered BbAfcB for production of human milk oligosaccharides may be a more suitable strategy than employing CpAfc2 from pathogenic C. perfringens. The newly obtained transfucosylation activity could be exploited to expand the
complexity of HMO production: mixtures of 3FL and LNT – e.g. of fermentation origin – could be enriched with LNFP II, while mixtures of 3FL and LNnT could be enriched with LNFP III.

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References


Figure legends

**Figure 1.** Transfucosylation reactions catalysed by α-L-1,3/4-fucosidases. Top: Formation of lacto-N-fucopentaose II (LNFP II) from 3-fucosyllactose (3FL) and lacto-N-tetraose (LNT). Bottom: Formation of LNFP III from 3FL and lacto-N-neotetraose (LNNT). The fucosyl moieties are shown in blue.

**Figure 2.** Homology model of BbAfcB (white) indicating the catalytic residues D703 and E746 (yellow). The ligand comprising Fuc (orange), GlcNAc, and non-reducing end Gal (purple-blue) moieties from LNFP II is from the crystal structure of BiAfcB (PDB: 3UET). The BbAfcB α-helical loop (759-AAYNDGVKVLKPGQMAPDGLK-781) subjected to mutation in the current work is shown in black. The position and shape of corresponding α-helical loop (259-DIEKMKERENPTYLNNG-275) in the CpAfc2 homology model to which it was changed is indicated in pink.

**Figure 3.** Transfucosylation catalysed by the α-L-1,3/4-fucosidases. Molar transfucosylation yields ([LNFP II/III]/[3FL]) obtained in transfucosylation catalysed by BbAfcBmut (squares), BbAfcB (triangles), and CpAfc2 (circles) using 3FL as fucosyl donor substrate and either LNT (left) or LNNT (right) as acceptor substrate (n = 2). Acceptor-to-donor ratios (A:D) were either 10 (10 mM 3FL and 100 mM LNT/LNNT; filled symbols) or 1 (20 mM of both substrates; open symbols).

**Figure 4.** Effect of acceptor-to-donor ratio (A:D) and substrate concentration on transfucosylation. The reaction was catalysed by BbAfcBmut (squares) and BbAfcB (triangles) for up to 60 min using 3FL and LNT (n = 2). For A:D 1 and 5 (dashed lines) the 3FL donor concentration was 20 mM, whereas it was 10 mM for A:D 2 and 10 (solid lines). For A:D 1 and 2 (open symbols) the LNT acceptor concentration was 20 mM, whereas it was 100 mM for A:D 5 and 10 (filled symbols). Transfucosylation activity is given as molar LNFP II yield on donor (%; left) and as LNFP II concentration (mM; right).
Figure 1.
Figure 2.
Figure 3.
Figure 4.
### Tables

**Table 1.** Hydrolytic activity of α-1,3/4-L-fucosidases on 3FL ($n = 2$).

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Hydrolytic activity on 3FL ($\mu\text{mol min}^{-1} (\mu\text{mol enzyme})^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CpAfc2</td>
<td>$18 \pm 3^b$</td>
</tr>
<tr>
<td>BbAfcB</td>
<td>$59 \pm 1.5^a$</td>
</tr>
<tr>
<td>BbAfcB$_{\text{mut}}$</td>
<td>$0.01 \pm 0.001^c$</td>
</tr>
</tbody>
</table>

$^a-c$Significant difference ($p < 0.05$) between enzymes.
Table 2. Initial transfucosylation rate, maximum conversion yield ([LNFP II/III]/[3FL]₀), and yield after 24 hours in transfucosylation catalysed by BbAfcB, BbAfcBₘᵤₜ, and CpAfc2 using 3FL as donor substrate and either LNT or LNnT as acceptor substrate (n = 2) at acceptor-to-donor ratios (A:D) of either 10 or 1.

<table>
<thead>
<tr>
<th>Acceptor</th>
<th>Enzyme</th>
<th>A:D</th>
<th>3FL (mM)</th>
<th>Initial transfucosylation rate (μM min⁻¹)</th>
<th>Maximum yield of LNFP II/III (%)</th>
<th>LNFP II/III yield after 24 hours (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LNT</td>
<td>BbAfcBₘᵤₜ</td>
<td>10</td>
<td>10</td>
<td>46 ± 6d\text{x}</td>
<td>39 ± 2b\text{x} (3.9)</td>
<td>34 ± 11</td>
</tr>
<tr>
<td></td>
<td>BbAfcB</td>
<td>10</td>
<td>10</td>
<td>709 ± 17\text{ab,}\text{x}</td>
<td>14 ± 0.3\text{c} (1.4)</td>
<td>0.7 ± 0.04*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>20</td>
<td>24 ± 4d\text{v}</td>
<td>13 ± 4\text{c,}\text{v} (2.7)</td>
<td>13 ± 4</td>
</tr>
<tr>
<td></td>
<td>CpAfc2</td>
<td>10</td>
<td>10</td>
<td>517 ± 56\text{bc,}\text{x}</td>
<td>39 ± 2b\text{x} (3.9)</td>
<td>0.4 ± 0.1*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>20</td>
<td>273 ± 0.3\text{c,}\text{d,}\text{v}</td>
<td>16 ± 3\text{c,}\text{v} (3.2)</td>
<td>0.4 ± 0.6*</td>
</tr>
<tr>
<td>LNnT</td>
<td>BbAfcBₘᵤₜ</td>
<td>10</td>
<td>10</td>
<td>23 ± 2\text{c,}\text{v}</td>
<td>11 ± 0.3\text{c,}\text{v} (1.1)</td>
<td>5.4 ± 2*</td>
</tr>
<tr>
<td></td>
<td>BbAfcB</td>
<td>10</td>
<td>10</td>
<td>163 ± 16\text{a,}\text{v}</td>
<td>8.1 ± 2\text{a,}\text{v} (1.6)</td>
<td>0.0 ± 0.0*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>20</td>
<td>12 ± 2\text{c,}\text{d,}\text{v}</td>
<td>8.1 ± 2\text{a,}\text{v} (1.6)</td>
<td>0.0 ± 0.0*</td>
</tr>
<tr>
<td></td>
<td>CpAfc2</td>
<td>10</td>
<td>10</td>
<td>101 ± 5\text{bc,}\text{z}</td>
<td>17 ± 3\text{c,}\text{y} (1.7)</td>
<td>0.1 ± 0.03*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>20</td>
<td>77 ± 6\text{b,}\text{z}</td>
<td>6.2 ± 2\text{c,}\text{y} (1.2)</td>
<td>0.04 ± 0.002*</td>
</tr>
</tbody>
</table>

\text{a-d}Significant difference (p < 0.05) between values for each parameter comparing enzymes on the same acceptor substrate (LNT or LNnT) at both A:Ds. \text{^x-y}Significant difference (p < 0.05) between values for each parameter comparing acceptor substrates for the same enzyme at both A:Ds. *For yields after 24 hours, an asterisk indicates whether this value is significantly different from the maximum yield, i.e. if significant product hydrolysis took place within 24 hours.

Table 3. Effect of acceptor-to-donor ratio (A:D) and substrate concentrations on initial transfucosylation rate (divided by [3FL]₀ to clearly observe the effect of varying donor and acceptor concentrations) and on maximum LNFP II yield (given as molar yield on donor, i.e. [LNFP II]/[3FL]₀) obtained within 60 min in the transfucosylation catalysed by BbAfcB, BbAfcBₘᵤₜ, and CpAfc2 using 3FL and LNT (n = 2).

<table>
<thead>
<tr>
<th>A:D</th>
<th>3FL (mM)</th>
<th>LNT (mM)</th>
<th>Initial transfucosylation rate (μM min⁻¹ [3FL]₀⁻¹)</th>
<th>Maximum LNFP II yield in 60 min (mM [3FL]₀⁻¹)</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td>BbAfcBₘᵤₜ</td>
<td>BbAfcB</td>
</tr>
<tr>
<td>1</td>
<td>20</td>
<td>20</td>
<td>1.2 ± 0.2\text{d}</td>
<td>14 ± 0.0\text{c}</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>20</td>
<td>1.2 ± 0.1\text{d}</td>
<td>51 ± 0.4\text{b}</td>
</tr>
<tr>
<td>5</td>
<td>20</td>
<td>100</td>
<td>4.8 ± 0.2\text{d}</td>
<td>70 ± 5.3\text{a}</td>
</tr>
<tr>
<td>10</td>
<td>10</td>
<td>100</td>
<td>4.6 ± 0.6\text{d}</td>
<td>71 ± 1.7\text{a}</td>
</tr>
</tbody>
</table>
*Significant difference ($p < 0.05$) between values for each parameter.