



Proceedings

# An In Silico Approach to Enzymatic Synthesis of Fucooligosaccharides Using $\alpha$ -L-Fucosidase from Thermotoga maritima $^{\dagger}$

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Abstract: Fucooligosaccharides comprise the primary group of human milk oligosaccharides. Due to their beneficial properties, a series of synthetic methods have been proposed to obtain them. Enzymatic methods show great promise, and  $\alpha$ -L-fucosidase from *Thermotoga maritima* has emerged as a powerful catalyst for their production. Nonetheless, the enzyme's limited substrate scope has delayed its wider application. The present work aims to compare the relative reactivity of fucose, pNP-fucose, and ethyl-fucose, while also exploring the molecular interactions of these fucosyl-donors with the enzyme through a combination DFT and docking analysis. The HOMO-LUMO band gaps range from -7.14571 to -4.24429 eV, with  $\alpha/\beta$ -pNP-fucose and  $\alpha$ -fucose being the three most reactive compounds. Moderate association energies between -6.4 to -5.5 kcal·mol<sup>-1</sup> were found in the docking analysis, with  $\alpha$ -pNP-fucose and both anomers of ethyl-fucose demonstrating the poorest affinity. In the case of  $\alpha/\beta$ -lactose affinity to the  $\beta$ -fucose/enzyme complex, no significant differences were shown. We conclude that the best fucosyl-donors for transfucosylation are those that maintain an enzyme affinity and reactivity similar to pNP-fucose.

**Keywords:** Fucooligosaccharides;  $\alpha$ -L-fucosidase; DFT study; molecular docking

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#### 1. Introduction

Fucooligosaccharides (FucOS) are the main oligosaccharides in human milk comprising 65–77% of the total oligosaccharide content [1]. Due to their antimicrobial, immunomodulatory, and prebiotic activities, as well as their promise to function as developmental cognitive enhancers, their incorporation into commercial formulations has become highly desirable [1,2]. As their isolation is complex due to their low abundance in animal milk, attention has turned to synthesis [1,2]. Fermentation is the most efficient, with generally recognize as safe (GRAS) certification from the U.S. Food and Drug Administration (FDA) permitting the addition of 2'-fucosyllactose (2'FL) to infant formula [2]. Another synthetic alternative with recent promising results is the use of isolated en-

Chem. Proc. 2021, 3, 10 2 of 6

zyme, which requires the yield optimization of FucOS by fucosyl-transferases. Unfortunately, this approach presents the inconvenience of requiring nucleotide sugars as fucosyl-donors, which are more expensive than those used for fucosidases [1,3]. Consequently, FucOS synthesis by fucosyl-hydrolases like the  $\alpha$ -L-fucosidase from *Thermotoga maritima* has gained importance, as this pathway allows the use either of less expensive fucosyl-donors or even agro-industrial waste [3–8]. However, this enzymatic route provides lower yields than the transferase, or involves the release of toxic compounds such as p-nitrophenol [5–8]. Alternatives are highly desirable. Thus, the present work aimed to determine the relative reactivity of three non-classical fucosyl-donors through an in silico study to propose substrate alternatives for the enzymatic synthesis of FucOS.

#### 2. Methods

#### 2.1. Geometry Optimization and HOMO-LUMO Parameters

All compounds were totally geometry optimized through the Density Functional Theory (DFT) with the B3LYP/6-311++G(2d,2p) basis set using water as solvent. The highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) density surfaces were visualized with Gabedit 2.5.0. [9].

#### 2.2. Molecular Docking for Hydrolysis and Transfucosylation Process

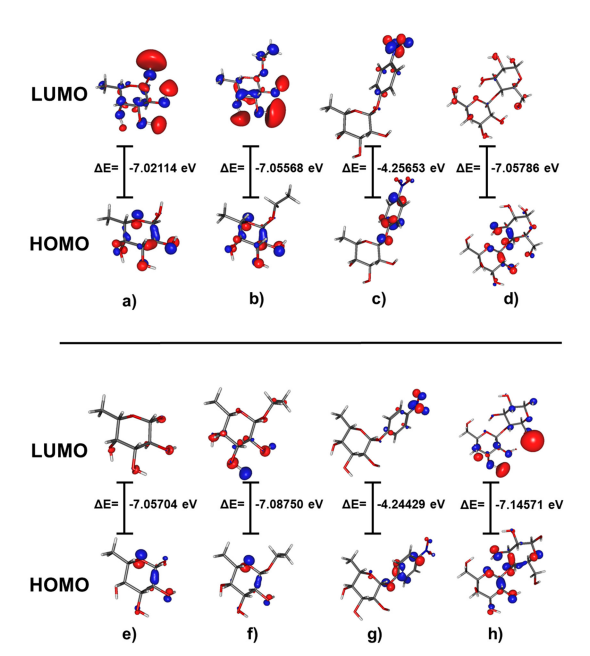
The A chain of the  $\alpha$ -L-fucosidase crystal from *Thermotoga maritima* (PDB: *10DU*) was prepared with the DockPrep tool implemented in Chimera 1.13.1 [10], prior to its use as the receptor for molecular docking. All molecular dockings were performed by Autodock VINA [11] through the PyRx software [12], taking as reference the amino acids from the active site cited by Sulzenbacher et al. [13], with coordinates for the search space centered on x: –20.63, y: 19.03, and z: 63.32, with a grid cube with dimensions of 25.00 Å. In the case of the hydrolysis, a single docking step was performed for each fucosyl-donor and the receptor. Meanwhile for the transfucosylation process, a sequential docking sequence was employed. First,  $\beta$ -fucose was docked to the enzyme in order to form the pre-complex, then docking was performed again with lactose. The best binding mode for each interaction was obtained and its interactions were processed with the BIOVIA Discovery Studio Visualizer© v19.1.0.18287 [14].

# 3. Results and Discussion

#### 3.1. HOMO-LUMO Parameters

The HOMO-LUMO frontier orbitals and the bandgaps were calculated for both anomers of fucose, ethyl-fucose, p-nitrophenyl (pNP)-fucose, and lactose (Figure 1). In general,  $\beta$ -anomers showed a lower bandgap compared with the  $\alpha$ -anomers, except for the pNP-fucose anomers, which showed similar bandgaps. As the magnitude of the HOMO-LUMO gap directly relates to chemical reactivity, where larger bandgaps predict lower reactivity [15,16], we predict that the  $\beta$ -anomers of fucose, ethyl-fucose, and lactose should prove less reactive than their  $\alpha$ -anomers; in contrast, both anomers of pNP-fucose should demonstrate similar reactivity. Ranking the series,  $\alpha/\beta$ -pNP-fucose with its electron withdrawing nature, unsurprisingly, is predicted to be the most reactive followed by  $\alpha$ -fucose. In general, the  $\alpha$ -anomers promise greater reactivity than the  $\beta$  anomers (again, with the exception of the hot pNP electrophile) based on the distribution of density in the frontier molecular orbitals.

Chem. Proc. **2021**, 3, 10 3 of 6



**Figure 1.** HOMO and LUMO surfaces (blue: positive and red: negative) of (a)  $\alpha$ -fucose, (b)  $\alpha$ -ethyl-fucose, (c)  $\alpha$ -pNP-fucose, (d)  $\alpha$ -lactose, (e)  $\beta$ -fucose, (f)  $\beta$ -ethyl-fucose, (g)  $\beta$ -pNP-fucose, and (h)  $\beta$ -lactose, as well as their HO-MO-LUMO band gap.

## 3.2. Molecular Docking for Hydrolysis Processes

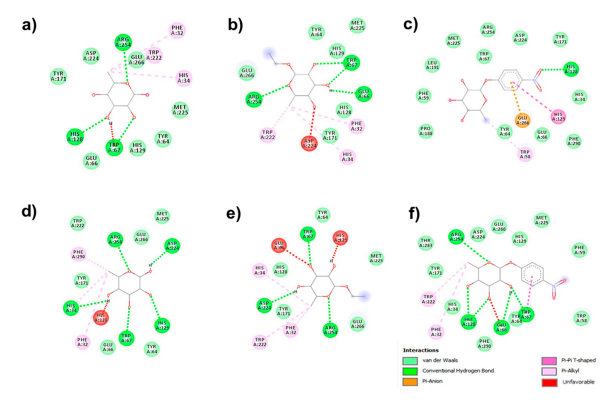
Single molecular docking simulated the hydrolysis catalyzed by *T. maritima's*  $\alpha$ -L-fucosidase. Docking scores varied between –6.4 and –5.5 kcal·mol<sup>-1</sup>, with  $\beta$ -pNP-fucose showing the best affinity to the receptor, while  $\alpha$ -ethyl-fucose presented the worst (Table 1). A lower substrate—enzyme affinity could be correlated with lower complex stability and a tendency to destroy it [17]. Thus, among the six different tested molecules,  $\alpha$ -ethyl-fucose,  $\beta$ -ethyl-fucose, and  $\alpha$ -pNP-fucose appear to be the fucosyl donors most readily hydrolyzed once bound to the enzyme.

Chem. Proc. 2021, 3, 10 4 of 6

<b>Table 1.</b> Coupling energies obtained for each fucosyl-donor and the $\alpha$ -L-fucosidase from
Thermotoga maritima.

Fucosyl-Donor	Coupling Energy (kcal·mol-1)
$\alpha$ -fucose	-6.0
β-fucose	-6.3
lpha-ethyl-fucose	-5.5
β-ethyl-fucose	-5.8
$\alpha$ -pNP-fucose	-5.9
β-pNP-fucose	-6.4

The key non-covalent interactions were identified (Figure 2). Those identified for  $\beta$ -fucose/enzyme docking (Figure 2d), agree with those reported by Sulzenbacher et al. [13]. All showed essential  $\pi$ -interactions between the C-5 methyl group and aromatic residues on the enzyme. The last interactions propitiate that the sugar ring takes a perpendicular position to this hydrophobic region, favoring van der Waals interactions observed with the rest of the molecule sites [13]. An important change was found for the interaction with the Asp224, the amino acid responsible for the nucleophilic attack that forms the covalent glycosyl-intermediate [13]. This pre-reaction interaction is a hydrogen-bond for  $\beta$ -fucose; but is van der Waals type for  $\alpha$ -fucose and  $\alpha/\beta$ -pNP-fucose; is an unfavorable repulsive interaction for  $\alpha$ -ethyl-fucose; and a hydrogen bond with the C-4 OH for  $\beta$ -ethyl-fucose, indicating the different binding modes of these two substrates. These insights could explain the differential docking energies and suggests that hydrolysis would prove more challenging.



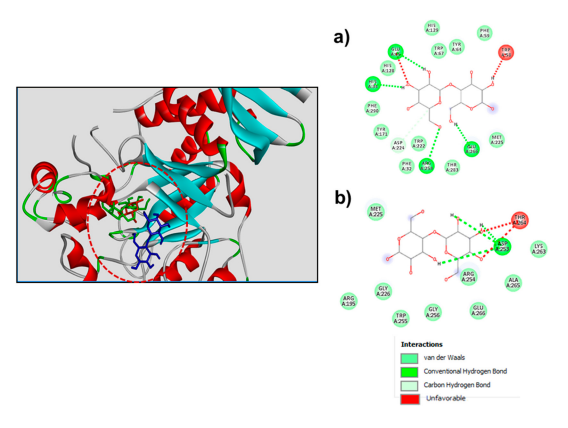
**Figure 2.** Mappings in 2D of the binding interactions among fucosyl-donors and the  $\alpha$ -L-fucosidase from *Thermotoga maritima*: (a)  $\alpha$ -fucose, (b)  $\alpha$ -ethyl-fucose, (c)  $\alpha$ -pNP-fucose, (d)  $\beta$ -fucose, (e)  $\beta$ -ethyl-fucose, and (f)  $\beta$ -pNP-fucose.

## 3.3. Molecular Docking for Studying Transfucosylation

In order to simulate a transfucosylation process, a sequential docking was performed. First, the  $\beta$ -fucose/enzyme complex was established by a single docking. The

Chem. Proc. 2021, 3, 10 5 of 6

results were consistent with the reports of Sulzenbacher et al. [13], with  $\alpha$ - or  $\beta$ -lactose, providing docking scores of –5.7 and –5.8 kcal·mol<sup>-1</sup>, respectively. However, the conformation was different for each (Figure 3). While  $\alpha$ -lactose adopts a position near the  $\beta$ -fucose binding site,  $\beta$ -lactose adheres preferentially to a more distal site.  $\beta$ -lactose has only a series of weak van der Waals interactions holding it in place, including with the key Glu266 residue, while  $\alpha$ -lactose forms strong hydrogen bonds with both Glu66 and Glu266, the amino acids responsible for the activation of acceptor groups for transfucosylation [13]. This suggests that the alpha anomer should be far more reactive, and indicates that mutarotation likely forms this anomer prior to transformation.



**Figure 3.** Binding position of  $\alpha$ -lactose (green colored) and  $\beta$ -lactose (blue colored) in the complex  $\beta$ -fucose (orange colored)/enzyme, as well as the molecular interactions of (**a**)  $\alpha$ -lactose and (**b**)  $\beta$ -lactose to the receptor.

On the other hand, the effect of the fucosyl-donors on the transfucosylation process can be related to the reactivity showed in the HOMO-LUMO gap, because previous in vitro studies have shown the effective transference of pNP-fucose to lactose to synthesize FucOS [5,6], while other studies have found low yields or long process when fucose itself is used as the donor [7,8]. Thus, ethyl-fucose could show similar results to those obtained with fucose mainly due to both molecules showing similar reactivity. Finally, according to *in vitro* results obtained earlier and the in silico insights found here, it is possible to hypothesize that fucosyl-donors with similar structure and/or reactivity to that of pNP-fucose could act as good substrates for transfucosylation with the  $\alpha$ -L-fucosidase from *Thermotoga maritima*.

#### 4. Conclusions

In silico insights of the reactivity and molecular interactions with the  $\alpha$ -L-fucosidase from *Thermotoga maritima* obtained for each fucosyl-donor and *in vitro* results from literature allowed us to conclude that the best fucosyl-donors for transfucosylation reaction will be those with similar reactivity to pNP-fucose, so that the synthesis of compounds

Chem. Proc. 2021, 3, 10 6 of 6

with similar structures, but lower toxicity, should be prioritized to find next generation fucsosyl transfer agents.

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