Potential of lacto-\textit{N}-biose I as prebiotic for infant health: A review

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Abstract

Prebiotic is one alternative in the prevention of disease in infants. Generally, it is available as oligosaccharide which may occur naturally, but can also be added as a dietary supplement for food, beverage or formula. Lacto-\textit{N}-biose I (LNB), a kind of prebiotic has not been widely examined in regard to its activities as a bifidogenic factor. Naturally, it is available in a compound form in human milk oligosaccharide (HMO) as the main constituent of human milk rather than fat and protein. HMOs also have prebiotic activity in the body and play an important role in providing nutrition for the infant health. LNB is potential to be used in food ingredient, especially infant food formula regarding the prebiotic effect and it could be enzymatically synthesized using enzymes involved in the LNB biosynthesis pathway by microorganisms.

Keywords: Human milk oligosaccharide; infants health; lacto-\textit{N}-biose I; prebiotic; synthesis

1. Introduction

Infant disease is a problem related to the nutritional status of an infant in Indonesia. Gastroenteritis is a common disease occurred in infant [1]. It can be prevented by breastfeeding that can provide functional prebiotics [2]. Prebiotics are food ingredients that can not be digested in the intestine to give benefits to the body through the development of intestinal microorganisms providing a health effect for the body. They generally improve the population of beneficial microorganisms and reduce any harmful microorganisms in the body. In a proper dosage, the consumption of prebiotics can treat and support the control of diseases such as cancer, colon cancer, constipation, diabetes mellitus, and liver. In addition, prebiotics can regulate the growth of intestinal microorganisms, prevent gastrointestinal infections, modulates the immune system, increase the bioavailability of minerals, regulate metabolic disorders associated with obesity and diabetes, and reduce the risk of cancer [3].

Human milk oligosaccharide (HMO) is a major component contained in milk. It is known to have a prebiotic activity for the growth of infant intestinal microorganisms. It is composed of oligosaccharide type 1 containing lacto-\textit{N}-biose I (LNB) as the core structure and available in large numbers. LNB is the prebiotic candidate that has not been widely examined in regard to its activities as a bifidogenic factor. Several studies have reported that LNB significantly enhances the activity of microflora that lives in an infant’s intestine [4].

Naturally, LNB exists in HMO, but the recent study revealed that it can be synthesized by enzymatic reaction [5] and has the potential to be used as an ingredient [6]. LNB promotes the growth and activity of intestinal microorganisms, especially in the infant’s intestine.

2. Human Milk Oligosaccharide (HMO) and Its Prebiotic Activity

HMO is a building block of human milk in addition to lactose and fat. Its structure is very complex. HMO biosynthesis in the mammary gland starts with both the lactose core production of galactose and \textit{\beta}-galactotransferase in the presence of \textit{\alpha}-lactalbumin, which catalyzes glucose [7].

HMO has a varied structure and diverse bioactive properties. It consists of a combination of 5 monosaccharides constituent groups, i.e. D-glucose, \textit{\textit{N}}-acytetylglucosamine, L-fucose, and sialic acid. Certain monosaccharides have also been detected in other mammalian milk [8]. HMO cannot be hydrolyzed by the enzyme in the stomach and small intestine. Due to the complexity of its structure, the composition of HMO in human milk cannot be replicated in infant formula. HMO is more than just a substrate that stimulates the growth of microorganisms that are desirable in the infant’s intestine [9].

2.1. HMO affecting intestinal functions and immune protection

Some undigested parts of HMO will pass the lower digestive tract [10], which are selectively used for the metabolism of beneficial intestinal microorganisms in an infant’s body [11]. The bifidogenic activity of HMO can promote the growth of microorganisms on infant’s intestine better than the formula milk [12]. HMO has some beneficial effects on the development of the neonatal intestine, such as a) promoting the growth of beneficial microorganisms; b) protection against

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HMO is an inborn immune system component of human milk which protects children against infections. The protective effect of breastfeeding is caused by fucosylated $\alpha(1\rightarrow2)$-oligosaccharide bond. Fucosylated oligosaccharides can prevent diarrheal diseases through various mechanisms such as inhibiting activity of Escherichia coli toxin by binding and restricting the exposure to receptors, Campylobacter inhibition of intestinal cell adhesions and Norovirus suppression competing with intestinal epithelial tissues HMO can reduce pathogen infection with antimicrobial activity as an anti-adhesive [14].

2.2. HMO as a nutrient for brain development

Premature infants who are given by breastfeeding have a good development at the age of 1.8 years old and a higher level of intelligence at 7 years old [15]. Brain development and cognitive are partly dependent on gangliosides containing sialic acid and polysialic acid-glycoprotein [16]. The concentration of sialic acid in the brain twice as many among the few months before birth and a few years after birth [17]. This substance is a vital ingredient for acid formation in the brain before and after the births. Human milk is rich in sialic acid and a post-mortem analysis at this stage of neonatal shows that the ganglioside and the concentration of sialic acid bound protein are significantly higher in the brain of breastfed infants than the one by formula milk fed containing low sialic acid [16]. HMO is the main carrier of sialic acid that can trigger the brain development, which plays an important role at the intelligence level in infants fed by human milk [8].

2.3. HMO and systemic effect

HMO is known to have systemic impacts on endothelial cell leucocyte adhesion or interaction between platelets and neutrophils. T-cells in the blood are exposed by HMO. It makes the number of INFγ that produces CD3+CD4+, CD3+CD8+ T cells increases along with Interleukin 13 coding genes that produces CD3+CD8+ T cells. HMO affects the maturation of lymphocytes and promotes a shift in the response of T-cells to achieve the more balanced production of Th1/Th2-cytokines [7].

At the neutral fraction of HMO, lacto-$N$-fucopenatose (LNFP) III, and lacto-$N$-neotetraose (LNnT) have been shown to affect the ability of peritoneal macrophages to suppress the T-cell response of naive CD4+ [18]. LNFP III prompts the task of macrophages in vitro and increases the production of prostaglandin E2, IL-10 and TNFα [18]. Fucosylated HMO influences leucocyte infiltration and model flow activation using TNFα-activated endothelial cells and leucocytes in the isolated human umbilical vein. In the ex vivo models along the fresh human blood, the formation of the neutrophil complex and the activation of neutrophils in the presence of HMO are reduced [8].

2.4. HMO and intestinal microorganisms

The equilibrium of microorganisms in human’s intestines is one of the important factors in increasing the body’s immune system [19]. Intestines are the organs that are sensitive to environmental influences. Intestinal microorganisms and the intestinal epithelium have a very close relationship, especially in the digestive system [20]. Its mechanism occurs through the walls of the intestinal mucosa and microorganisms respond both beneficial and pathogenic [7]. Therefore, it is important to note the role of HMO and its influence in the selective microorganisms growth in the intestine.

Bifidobacterium longum subsp. infantis uses HMO is the only source of carbon, whereas HMO cannot be taken up by other gut microbes. Genome Bifidobacterium longum subsp. Infantis strain ATCC 15697 comprises a substantial gene cluster comprising of numerous glycosides and allegedly HMO metabolism-specific transporters [21]. The prevalence of clusters in HMO, as shown in the comparative study of genomics, correlates with their survival. Thus, not all enzymes were used in the bifidobacterial HMO metabolism at the time [22].

There are enzymes that degrade HMO in each strain bifidobacteria. The existence of such external fucosidase enzymes and lacto-$N$-biosidase produced by bacteria cells and specific transporter for LNB [Gal-$β$-(1→3)-GlcNAc] and GNB [Gal-$β$-(1→3)-GalNAc] proves that Bifidobacteria with several methods can particularly utilise HMOs [23]. B. bifidum is a microorganism that can degrade the HMO with a specific mechanism. It uses LNB to hydrolyze LNT, LNB I, and lactose. LNB was subsequently released, put in a cell by ABC transporters specifically for LNB and converted into C$_{6}$H$_{16}$NO$_{5}$P in the presence of lacto-$N$-biose I phosphorylase (LNBP) [24].

3. Lacto-$N$-Biose I

Lacto-$N$-biose I [LNB; Gal($β$1-3)GlcNAc] is an important role of disaccharide in the carbohydrate structure on several glycoconjugates (figure 1). This compound presents in HMO but it is not available in the form of a free disaccharide and active as building blocks of oligosaccharides type I containing lacto-$N$-tetraose as its core structure. Oligosaccharide type I is only contained in milk, and animals such as monkeys, while oligosaccharides Type II is detected in mammalian milk. In addition, LNB glycan is found in the domain of glycolipid and glycoprotein on several cell types, including epithelial cells in

![Fig. 1. Lacto-$N$-biose I](image-url)
intestine and blood group antigen [25].

3.1. Biosynthesis of LNB in bifidobacteria

The biosynthesis of LNB specifically is done by bifidobacteria such as B. bifidum which hydrolyzes HMO type 2 outside the cell by using sialidase, fucosidase, and lacto-N-biosidase. Then, LNB as the outcome enters the cell by ABC-type transporter and are transformed into Gal1P (Galactose-1-phosphate) and N-acetyhexosamine using LNBp that is first isolated from the cell-free extract of B. bifidum DSM20082.38 [26]. In the same hydrolysis activity of B. bifidum on HMO type II, it starts from the non-reducing end through the activity of sialidase, fucosidase, β-galactosidase and β-N-acetyhexosaminidase [27].

Biosynthesis of LNB in B. longum subsp. infantis directly incorporates LNT into the cytoplasm and is subsequently hydrolyzed to LNB and lactose. However, the metabolic pathway of LNB is encoded by lnpABCD operon that encodes LNBp, NahK, GalT and GalE correspondingly [28]. On that path, LNBp phosphorylates and hydrolyzes LNB into Gal1P, then transformed into Glu1P through GalT and GalE activity and some N-acetylgalcosamine (GlcNAc), phosphorylated via NahK produces GalNAc1P. B. bifidum has an identical operon containing two previously identified sugar kinases between the LNBp and NahK coding regions [29].

In majority of organisms galactose is metabolized to Gal1P without utilizing ATP that the two pathways are different. In the Leloir pathway, for example, α-Gal, phosphorylated by GalK utilizes one ATP molecule, but in the LNB/GNB system, LNBp creates Gal1P without utilizing ATP through galactoside phosphorylation. It is concluded that the other sections of the LNB/GNB in its pathway differ from those of Leloir in the use of N-acetylgalcosamine. By utilizing ATP, GalNAc has been converted into GlcNAc1P. In Leloir it appears like GalK. No clusters genes in the Bifidobacterium longum NCC2705 encoding any kinds of enzyme connected with the pathway of Leloir. GalK (BL1210) and GalT (BL1211), which are discovered in the genome, except the lnpABCD cluster, are the only gene cluster containing 2 Enzymes from Leloir’s pathway. Nonetheless, Gal M, GalK, GalT and GalE are generally present in operons of other species in the genes coding enzymes of the Leloir pathway for galactose metabolism. A metabolic pathway for galactose as a carbonyl source for Bifidobacterium longum is used in the simplified LNB/GNB pathway. Therefore, LNB is a bifidogenic factor for intestinal tract through HMO and carbohydrate metabolism mucin, because intestinal bifidobacteria is useful for health. In addition, LNB/GNB pathway is associated with symbiosis between humans and bifidobacteria [31].

3.2. The prebiotic activity of LNB

As a Bifidus factor, the prebiotic activity of LNB is found by bifidobacteria, especially B. bifidum, B. breve and B. longum. However, LNB shows no effect of prebiotics on lactic acid and other bacteria (Bacteroides, Clostridium, Enterococcus, Eubacterium, Lactobacillus, Lactococcus, and Ruminococcus) [4]. In addition, LNB can be utilized by many strains of bifidobacteria comprising of 208 strains of 10 species and 4 subspecies [32]. The whole strains that naturally live in infant’s gastrointestinal tract can grow by using a single LNB as a carbohydrate source, while adults, human teeth, and courageous domestic animals cannot breakdown the LNB by a strain present in the digestive tract.

As shown in table 1, the recent study evaluated the bifidogenic activity of lacto-N-biose I on the growth of microorganisms isolated from infant feces. In addition, comparative testing with other prebiotic compounds such as lactulose, raffinose, galacto-oligosaccharides, and mannanooligosaccharides was performed in vitro. The results of formula milk inoculated into the medium containing 1% LNB showed a significant improvement of the population of bifidobacteria as similar with other prebiotic compounds. But the populations of B. bifidum treated LNB 1% higher compared to the others. It is also proven from the lactic acid concentration of lower and higher acetic acid showing the larger prebiotic activity of LNB [6].

3.3. The effect of LNB on the immune system

Based on the previous research, the LNB immunomodulatory behavior employing mice's naïve immune cell expressing Receptor T-cells (TCR) especially chicken ovalbumin (OVA) demonstrated that the LNB exposure greatly decreased the Ag-specific production of IL-4 [33]. IL-4 is the leading development of the ag-specific Th2 cell in association with cytokine. LNB has the same lactose affinity according to another study [34]. Gal-3 that binds lactose can inhibit the relationship between TCR and Gal-3. It changes the effect of Gal-3, which binds LNB to damage the function partially and only reduce the secretion of Ag-specific IL-4. In general, by the modification of APCs signal, LNB affects Ag-specific immune response [33].

4. Synthesis of Lacto-N-Biose I

LNB synthesis was first introduced by Flowers and Jeanloz [35] that produced the derivatives of muramic acid and N-acetylmuramic acid. LNB synthesis was carried out to produce the products with high yields and high purity. This was due to the result of LNB synthesis used as a food ingredient that may provide a prebiotic effect on the formula. In addition, it took a shorter time to produce LNB for increasing productivity. Figure 2 represents a comparison between biosynthetic pathway in human and enzymatic synthesis of lacto-N-biose I [36-37].

4.1. Enzymatic synthesis of LNB

Research related to synthesis of lacto-N-biose I was in developed [5,26,28,36-37]. As briefly seen in table 2. LNB can
be synthesized using glycosynthase isolated from *Thermus thermophilus* through a trans-glycolytic mechanism [5]. The synthesis of LNB in scale-up has been conducted. It produced LNB up to 1.4 kg of sucrose and GlcNAc with the addition of UDP-Glc and phosphates using 4 enzymes such as sucrose phosphorylase (SP), GalT, GalE and LNB as the catalyst for 700 hours [28]. Such methods could be expected to be used on an industrial scale using a bioreactor system with an immobilized enzyme. In addition, the present invention is applicable to strengthen the hypothesis that in infant formulas, LNB may be employed as food additives similar to the LNB content in human milk.

LNB synthesis according to Nishimoto and Kitaoka started with the phosphorylation of sucrose into α-glucose-1-phosphate (Glc1P) with the addition of SP as a catalyst. Glc1P was transformed into UDP-Glc at the same time that UDP-Gal was transformed into Gal1P by GalT. As a result of the GalE behavior, the UDP-Glc was changed into UDP-Gal, where GalT was utilized by the UDP-Gal. LNB and then synthesized from Gal1P and GlcNAc by LNB [28].

4.2. The enzyme involved in LNB synthesis

The enzyme acts as a catalyst in a reaction. In certain conditions, the chemical reactions that run in the presence of a catalyst can be done faster. In LNB synthesis, enzyme plays an important role in catalyzing the chemical reactions. Enzymes that contribute to LNB synthesis so far quite a lot, dependent on the synthesis method. Some enzymes are generally present in the LNB pathway occurred in the infant's intestinal microorganisms to degrade LNB as a carbon source to stimulate activity and growth. Lacto-N-Biose I Phosphorylase (LNBp; 1,3-β-galactosyl-N-acetylgalactosamine phosphorylase) is active to phosphorylate LNB into Gal1P in the presence of GlcNAc. In CAZy database, LNBp is classified into glycoside hydrolase family 112 based on similarity of structure with the β-galactosidase GH42. Phosphorylase generally can hydrolyze sugar as the substrate which plays an important role as a source of energy, particularly for anaerobic bacteria. This is because phosphorylation directly produces sugar phosphorylated without requiring ATP. This energy can be obtained from the substrate which has greater energy than other sugars, especially in an anaerobic condition that only has 3 ATP molecules available through a glycolytic pathway from glucose 6-phosphate [24].

*LnpB* has shown an interaction in the production of using GlcNAc and ATP, α-GlcNAc 1-phosphate (GlcNAc1P) and it indicated that *LnpB* was a novel kinase anomeromic (NahK; N-acetylhexosamine 1-kinase) [31]. This protein showed a similar action on GlcNAc which produces GlcNAc1P with low activity in several monosaccharides. NahK had a number EC2.7.1.162 and was active in hexose, which phosphorylated its substrate in position 6.

*LnpC* and *lnpD* have been demonstrated as GalT (UDP-glucose-hexose 1-phosphate-uridylyltransferase; EC 2.7.7.12) and GalE (UDP-glucose 4-epimerase; EC 5.1.3.2) amino acid

**Table 1. The prebiotic effect of LNB in several experiments**

<table>
<thead>
<tr>
<th>Microorganism</th>
<th>Information</th>
<th>Results</th>
</tr>
</thead>
</table>
| *Bifidobacterium longum* subsp. *infantis* ATCC15697 [40] | Comparison of the substrate specificity of LNB with pNP-β-Gal, pNP-β-Fuc, lactose, LacNAc, LNSII, and LNT | kₐ/K₀ value of LNB was higher, but it was lower than LNT, lactose, and LacNAc<br><br>Strains of Bifidobacteria can grow. *B. pseudocatenulatum, B. animalis* subsp. *animalis, B. pseudolongum* can utilize LNB<br><br>*B. bifidum* JCM1254 and JCM7004, *B. longum* subsp. *infantis* JCM1210 and JCM1222<sup>7</sup>, *B. longum* subsp. *longum* JCM1217<sup>7</sup> and JCM7054, *B. breve* JCM1192<sup>7</sup> and *B. scardovi* JCM12489<sup>3</sup> growth 2x > control |<br>-
| 208 strains consisted of 10 species and 4 subspecies of Bifidobacteria [32] | LNB used as a source of carbon addition of lopA gene (LNBp) |<br>-
| 18 strains of Bifidobacteria, *Clostridium, 13 Enterobacter* and 8 lactic acid bacteria [4] | Bacteria were inoculated into media containing LNB and the absorbance was evaluated in OD₅₀<br><br>Comparing the prebiotic effect of LNB with other prebiotics: lactulose, raffinose, galacto-oligosaccharide, and mannan-oligosaccharide |<br>The population of Bifidobacteria given LNB significantly grow as large as other prebiotics. The activity of *B. bifidum* was higher than other prebiotics. The residual compound of LNB treatment such as lactic acid was lower while acetic acid was higher than other prebiotics |<br>-
| Bifidobacteria [6] |<br>The activity of gen cluster Gnb REFGBCD was evaluated on several substrates, including LNB | LNB was hydrolyzed on Gnb G (phospho-β-galactosidase from *L. casei*) |<br>-
| *Lactobacillus casei* [41] |<br>The activity of gen cluster Gnb REFGBCD was evaluated on several substrates, including LNB | - |
sequence based accordingly. GalT and GalE are galactose metabolic enzymes associated in the Leloir pathway. Both of them become active in the change of Gal1P into Glc1P and allow the substrate to enter into the glycolytic pathway. Gal1P can be formed from LNB through phosphorylation by LNBP with GalT and GalE, indicating the transformation into Glc1P of LNB galactose component provided by an enzyme expressed in the gene cluster that produced energy [24].

Sucrose phosphorylase (SP) plays an important role in LNB synthesis on Nishimoto and Kitaoka’s method, which is active as a catalyst which phosphorylates sucrose into UDP-Glc. In this method, SP with the pET30 vector can process LNB synthesis from sucrose with the addition of UDP-Glc GlcNAc and phosphates to produce a yield of 99.6%. This enzyme has a number EC 2.4.1.7 [31].

Transglycosidase (1,4-α-D-glucan-(D-glucose)6-α-D-glucosyltransferase) can be used for LNB synthesis. This enzyme obtained from Thermus thermophilus that catalyzes transglycosylation of phenyl-2-amino-1-thio-β-D-glucopyranoside as the substrate into phenyl β-D-galactopyranosyl-(1-3)-2-deoxy -2-amino-1-thio-β-D-glucopyranoside which produces 2-acetamido analog and tipped LNB formation as the final product [5]. This enzyme has a number EC 2.4.1.24.

Table 2. Enzymatic synthesis of LNB in several experiments

<table>
<thead>
<tr>
<th>Microorganism Source</th>
<th>Enzyme</th>
<th>Substrate</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xanthomonas manihotis [38]</td>
<td>β-D-galactosidase</td>
<td>p-nitophenyl β-D-galactopyranoside dan 2-acetoamido-2-deoxy-D-glucopyranose</td>
<td>22</td>
</tr>
<tr>
<td>Bifidobacterium bifidum [26]</td>
<td>Lacto-N-biose I (LNBP)</td>
<td>Gal1P and GlcNAc</td>
<td>76</td>
</tr>
<tr>
<td>Bifidobacterium bifidum [31]</td>
<td>Sucrose phosphorylase, GalT, GalE, and LNBP</td>
<td>GlcNAc and Gal1P</td>
<td>96.9</td>
</tr>
<tr>
<td>Thermus thermophilus [5]</td>
<td>Transglycosidase</td>
<td>GlcNAc</td>
<td>80</td>
</tr>
<tr>
<td>Bifidobacterium longum and Ruminoccus albus NE1 [42]</td>
<td>LNBP and lactose phosphorylase</td>
<td>GlcNAC and Lactose</td>
<td>75.3</td>
</tr>
</tbody>
</table>

Fig. 2. Comparison of biosynthesis pathway (a) and enzymatic synthesis of lacto-N-biose I (b). LNB is connected to the galactosyl moiety of terminal lactose through a β-1,3-binding with an extra β-1,6-binding in branched HMO.
5. Conclusion

HMO is one of the major components of milk playing an important role as a source of nutrients for the infant’s health as well as prebiotics for infant intestinal microorganisms to improve health. LNB is a disaccharide that is bound to the core structure of HMO contained in large numbers. In addition, LNB has a prebiotic activity by stimulating growth and activity of microorganisms such as bifidobacteria which are useful for health. LNB has a great potential to be used as a food ingredient, especially for infant formula. In addition, LNB can be synthesized enzymatically, chemically, and chemoenzymatically using enzymes involved in the biosynthesis pathway LNB by microorganisms. The synthesis of good lacto-N-biose I has been widely used to obtain high productivity and efficiency. Although some methods require long processing times with complex stages, a continuous system or a one-pot system can be used to solve this problem. For the future prospect, LNB production on a large scale also needs to be carried out for commercialization purposes. In addition, as a prebiotic candidate, lacto-N-rose I has the prospect of being used as a food ingredient because of its functionality. This is related to its application as a food additive in infant formula. Therefore, regulations related to their use need to be issued to increase the potential for their use in food.

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