Human milk 3’-Sialyllactose is positively associated with language development during infancy

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ABSTRACT

Background: Genetic polymorphisms leading to variations in human milk oligosaccharides (HMO) composition have been reported. Alpha-Tetrasaccharide (A-tetra), an HMO, has been shown to only exist between 3'-Sialyllactose (3'-SL) and age, provided the original work is properly cited

Methods: We enrolled typically developing children (2–25 months old; mean, 10 months old) who were at least partially breastfed at the study visit. The Mullen Scales of Early Learning (MSEL) were used as the primary outcome measure to assess early cognitive development. Linear mixed effects models were employed by stratifying children based on A-tetra levels (A-tetra+ or A-tetra-) to assess associations between age-removed HMO concentrations and both MSEL composite scores and the 5 subdomain scores.

Results: A total of 99 mother-child dyads and 183 HM samples were included (A-tetra+: 57 samples, 33 dyads; A-tetra−: 126 samples, 66 dyads). No significant association was observed between HMOs and MSEL when all samples were analyzed together. The composite score and 3’-sialyllactose (3’-SL) levels were positively associated (t < 0.002; effect size (EF), 13.12; 95% CI, 5.36–20.80) in the A-tetra + group. This association was driven by the receptive (adjusted P = 0.015; EF, 9.95; 95% CI, 3.91–15.99) and expressive (adjusted P = 0.048; EF, 7.53; 95% CI, 2.51–13.79) language subdomain scores. Furthermore, there was an interaction between 3’-SL and age for receptive language (adjusted P = 0.03; EF, -14.93; 95% CI, -25.29 to -4.24).

Conclusions: Our study reports the association of 3’-SL and cognition, particularly language functions, in typically developing children who received HM containing detectable A-tetra during infancy.

Keywords: human milk, early cognitive development, Mullen Scales of Early Learning, human milk oligosaccharides, sialyllactose, language, random mixed effects model, breastfed infants

Introduction

A wealth of evidence has documented the health benefits of breastfeeding during infancy (1). Human milk (HM) contains a wide array of nutrients and bioactives; among the latter are a vast amount and diversity of human milk oligosaccharides (HMOs) (2, 3). HMOs, the third most abundant solid component of HM, are formed in the mammary glands starting from lactose by the action of a series of glycosyltransferases that elongate lactose with 1 or several moieties of N-acetyl-lactosamine, lacto-N-biose, fucose (Fuc), and sialic acid (4). Several lines of evidence have revealed the potential health benefits of HMOs (3, 5–14). In particular, both sialyllactose (SL) (15–20) and 2’-fucosyllactose (2’-FL) (21–24) have been implicated to benefit early cognitive development in preclinical studies, suggesting that early life dietary supplementation with SL and 2’-FL improved learning ability and spatial cognition and reduced stress-induced anxiety (15–18, 21–23). Furthermore, Berger et al. (24) reported that greater exposure to 2’-FL relative to other HMOs at 1 month but not at 6 months of age was associated with better cognitive development at 24 months. More recently, Oliveros et al. (25) reported a positive association between 1 month postpartum HM 2’-FL levels and 6 month motor scores, whereas 6’-sialylactose (6’-SL) levels were positively associated with cognitive and motor scores at 18 months.

While the aforementioned studies have reported positive health benefits of HMOs during infancy, it is important to note that variations of HMO composition resulting from genetic polymorphisms, particularly fucosyltransferase genes (5), have been reported. Austin et al. (26) assessed 10 HMOs of HM samples obtained from Chinese women. They reported that with
the exception of alpha-Tetrasaccharide (A-tetra), the majority of HMOs were above the limit of detection (LoD). Specifically, of the 446 samples evaluated, only 65 HM samples (~14.6%) exhibited an A-tetra level greater than the LoD (A-tetra+). While the underlying biological mechanisms determining the levels of A-tetra in HM samples are largely unknown, Kobata (27) reported that A-tetra may only be present in mothers with a blood type A, which is consistent with the percentage of the Chinese population with an A blood type reported by Austin et al. (26). These findings raised 2 major considerations regarding the levels of A-tetra in HM. First, the presence (A-tetra+) or absence (A-tetra-) of A-tetra is associated with blood type, suggesting potential genetic origins. Second, given a large portion of HM samples are expected to be A-tetra-, statistical approaches accounting for this major factor should be considered. Taken together, in this study we aimed to examine the potential associations between HMOs and cognitive development, assessed by the Mullen Scales of Early Learning (MSEL; primary outcome) during infancy.

Methods

Informed consent for both their own participation and the participation of their infant was obtained from parents prior to enrolling in the study. All study activities were approved by the University of North Carolina at Chapel Hill (UNC) and University of Minnesota (UMN) Institutional Review Boards. A subset of the Baby Connectome Project (BCP) who were younger than 3 years old were also enrolled in the BCP-Enriched study. Of subjects enrolled in BCP-Enriched study who were breastfed (including exclusive/predominant and mixed feeding) at the scheduled visits were included in the data analyses reported herein. Additional details of the BCP study can be found in Howell et al. (28) and Supplemental Information 1. The exclusively/predominantly breastfed infants were the infants who were fed less than 4 teaspoons or 20 g per day of nonformula and complementary foods/liquids (water, apple juice, etc.). In contrast, the mixed breastfed infants were the infants who were fed more than 50% human milk and some formula and other foods/liquids. The mixed formula–fed infants were those who were fed more than 50% formula, but also some human milk and other foods/liquids. Subjects were enrolled at both UNC (Site A) and UMN (Site B) using site-based research registries for identifying research participants between birth and 3 years of age. Recruitment strategies were further supplemented by local newborn nurseries, institutional centers with research activities focusing on early brain development, university listservs, and local flyers. The inclusion criteria were birth at gestational age 37–42 weeks, birth weight appropriate for gestational age, and absence of major pregnancy and delivery complications. The exclusion criteria included cases in which the child was adopted; the presence of a first-degree relative with autism, an intellectual disability, schizophrenia, or bipolar disorder; birth weight < 2000 grams; neonatal hypoxia (10-minute APGAR [Appearance, Pulse, Grimace, Activity, and Respiration] <5); illness requiring a newborn intensive care unit stay > 2 days; chromosomal or major congenital abnormality; abnormal magnetic resonance on a previous MRI; significant medical illness or developmental delay or significant medical and/or genetic conditions affecting growth, development, or cognition (including visual/hearing impairment); contraindication for MRI; and the mother having preeclampsia, having a placental abruption, living with HIV, or using alcohol or an illicit drug during pregnancy. Finally, both hearing and vision functions were also assessed.

HM collection and macronutrient analyses

HM samples were obtained using a hospital-grade, electric Medela Symphony breast pump from the right breast at each visit. The milk samples were collected until no more HM could be extracted, so that the collective sample was representative of the nutrient concentration of the right breast for 1 feeding. In addition, collections were standardized to the second feed of the day whenever possible to avoid diurnal variation of milk nutrients. The weight and volume of milk collected were recorded, and the milk was then vortexed for 2 minutes at the highest speed. The volume of milk was measured using a graduated cylinder with special attention to avoid bubbles. Subsequently, 2.5 mL of milk was used for midinfrared spectroscopic analyses (MIRIS Human Milk Analyzer; Miris AB, Uppsala, Sweden). This step was to ensure that the total fat content (as an indicator of the quality of milk sampling) fell within the expected range. Finally, an aliquot of the minimum 30 mL of volume was transferred from the collection bottle to a 50 mL polypropylene Falcon tube. Using...
repeat pipette and an appropriate tip, 11 aliquots of 1 mL were made in 1-mL Eppendorf tubes and 9 aliquots of 2 mL were made in 2-mL Eppendorf tubes for storage in a −80 °C freezer within 30 minutes from the end of the time of collection.

**HMO analyses**

A representative 1-mL aliquot of collected HM was shipped on dry ice to Neutron Spa (Italy) for HMO quantification by LC with fluorescence detection after labeling with 2-amino-azobenzamide using the method of Austin and Bénet (29). The following HMOs were quantified using standard curves with authentic HMO standards: 2'-FL, 3-fucosyllactose (3-FL), 3'-sialyllactose (3'-SL), 6'-SL, lacto-N-tetraose (LNT), lacto-N-neotetraose (LNNT), lacto-N-fucopentaose I (LNFP-I), and A-tetra.

The milk samples collected by Site A were all analyzed in 1 batch, whereas the milk samples collected by Site B were analyzed in 2 batches (1 along with the samples from Site A and the other with only samples from Site B).

**Mullen Scales of Early Learning**

The MSEL, a widely used tool to assess infant cognitive development, is comprised of 5 subdomains, including fine motor, gross motor, visual reception, receptive language, and expressive language. An early learning composite (ELC) score corresponding to the Developmental Quotient score for infants can be derived using the scores of all subdomains, excluding the gross motor subdomain. The MSEL was administered at every visit by trained staff who were not blinded to subjects’ milk exposures.

**Statistical modeling and analyses**

R version 3.6.3 (R Studio) was used for all statistical analyses. It is well known that HMO concentrations change across lactation (5). Thus, in order to remove the potential confounding effects from age, age effect was removed from the HMOs using smooth spline regression prior to conducting association analyses (30). Nevertheless, because the quantities of some of the HMOs are known to be below the LoD of the applied assays (undetectable), the age effect removal procedures were only applied to those with HMO concentrations greater than the LoD. In addition, log transformation was applied whenever needed in order to minimize the heteroskedasticity and satisfy the linear assumption of the linear models applied (31).

The associations between the 8 age-removed HMO concentrations and concurrently collected MSEL scores (as the response variable), including the ELC score and each of the subdomain MSEL t-scores, were tested. The data acquisition process and construction of the final data set for the association analysis are depicted in Supplemental Figure 1. To account for the longitudinal aspect, a random linear mixed effects model was used, with 2 random intercepts for the infant and MSEL examiner (32). As implicated by Verbyla (33), the restricted maximum likelihood (REML) approach can be used for model comparisons in addition to parameter estimation when the original likelihood is used with the REML estimates. Thus, the REML approach was used for fitting the models and for model comparison. The Akaike Information Criterion (AIC) was calculated using the original likelihood and the REML estimates (34). Potential HMO analysis batch and site effects were also controlled and were considered as binary variables.

Three statistical models were employed in our study, and detailed information is provided below. Model 1 was used to analyze all 8 of the HMOs for all samples without any stratifications.

The MSEL data represent the ELC or each of the subdomain scores. We applied Model 1 to assess potential associations between MSEL scores and all 8 measured HMOs for the entire population, without any stratifications.

As indicated above, our study aimed to determine whether associations between HMOs and cognitive development differ between A-tetra+ and A-tetra- groups. We stratified children as being either A-tetra+, meaning receiving HM containing more A-tetra than the LoD (4.4 mg/L), or otherwise as being A-tetra-(29). In addition, if A-tetra values were greater than 0 but lower than the LoD, these values were assigned as 0. For the A-tetra-subjects, the analysis batch and site were used as independent variables with 2 random intercepts, whereas in addition to the variables used for the A-tetra-subjects, 8 age-removed HMOs were also included for the A-tetra+ subjects.

Finally, to evaluate whether the HMOs exhibiting significant associations with MSEL have different associations at different ages, an indicator variable for age cutoff point was included as both main and interaction terms in Model 3. To choose the appropriate age cutoff point, the AIC was compared between models with different cutoff ages.

**Results**

A total of 99 mother-child dyads were enrolled in this study, and 183 HM samples were collected. The mean duration of breastfeeding was 14.4 ± 4.95 months. Among the 99 child participants, 80 participants were exclusively/predominantly breastfed, 15 were mixed breastfed, and 1 was mixed formula fed. Although feeding practice information was not available from the remaining 3 participants, they were all breastfed at the time of visits. With the exception of 1 participant who exhibited nystagmus, all children had normal vision and hearing tests. The age distributions for the study visits are shown in Figure 1. The numbers of participants with 1, 2, 3, and 4 study visits were 49, 24, 18, and 8, respectively. Detailed subject information; the weights, volumes, measured fat, carbohydrates, and energy of the milk samples; and 8 HMO concentrations for all, A-tetra-, and A-tetra- subjects are summarized in Table 1. HM samples from 2 subjects showed A-tetra levels greater than the LoD at some visits but 0s during the other visits (40, 28, 0, and 43 mg/L in 1 subject and 35, 0, 45, and 0 mg/L in the other subject). These 2 subjects were assigned to the A-tetra+ group. No significant differences in demographics were observed between the A-tetra+ and A-tetra- groups (t-test for all parameters, with the exception of household incomes, where a chi-square test of independence was used). The sex variable was not significant, and thus was not included in the model. While 3-FL concentrations were significantly different between the 2 groups, the inclusion or exclusion of 3-FL in the linear mixed effect models did not yield significant differences in the fitted results. Thus, results using all of the 8 HMOs are reported below. Relations of patterns of human milk exposure and
HMO concentrations between A-tetra+ and A-tetra- groups were also studied (Supplemental Tables 1 and 2), and we concluded that no distinct difference existed between these groups (Supplemental Information 2). Table 2 shows the MSEL ELC and individual subdomain scores, which are all within the anticipated normal ranges. While variables such as socioeconomic status and education levels of parents have been shown to impact early cognitive development of offspring, Tables 1 and 2 show that the demographic information and MSEL scores did not differ significantly between the A-tetra+ and A-tetra- subgroups, justifying the exclusion of these variables from the models.

The HMO concentrations are highly age dependent (Figure 2), with 2’-FL, LNFP-I, LNNT, LNT, and 6’-SL decreasing with age, whereas A-tetra, 3-FL, and 3’-SL increase with age. For 2’-FL, 6’-SL, and LNFP-I, age has a significant quartic effect (adjusted P values < 0.001 for all 3 effect sizes (EFs): 1837.19, 338.66, and 996.60 for quartic age, respectively; 95% CIs, 974.16–2701.50, 217.20–459.51, and 578.77–1412.53 for quartic age, respectively), whereas 3-FL, A-tetra, 3’-SL, LNT, and LNNT exhibited significant quadratic effects (adjusted P values < 0.01 for all 5 EFs, -1655.18, 56.27, 170.88, 1232.48, and 200.51, respectively; 95% CIs, -2729.39 to -582.43, 21.81–90.82, 51.72–288.45, 697.81–1773.43, and 103.11–299.45, respectively). Notably, about 70% (129 samples), 21% (39 samples), and 12% (22 samples) of the HM contained A-tetra, LNFP-I, and 2’-FL below the LoDs (4.4, 2.0, and 3.9 mg/L, respectively) (29). Among these 3 HMOs, HM samples exhibiting 2’-FL levels below the LoD also had undetectable LNFP-I, and all the undetectable LNFP-I samples also had undetectable A-tetra.

For age-removed 3’-SL, log transformation was necessary in order to minimize the heteroskedasticity and satisfy the linearity assumption. Comparisons between distributions with and without the log-transformation of 3’-SL are provided in Supplemental Information 3. Furthermore, evaluations of intra-subject variability of 3’-SL from before and after age effects were removed are provided in Supplemental Information 3.

No significant associations among cognition and HMO concentrations were observed when all samples without stratifications were analyzed using Model 1. In contrast, a significantly positive association existed between the ELC scores and age-removed 3’-SL (P = 0.002; EF, 13.12; 95% CI, 5.36–20.80), with a power of 0.89 when children were stratified into A-tetra+ and A-tetra- groups (Model 2), suggesting that a higher 3’-SL concentration is associated with a higher ELC score in children receiving HM with measurable levels of A-tetra (Table 3). More specifically, controlling for other HMOs and site and batch effects, the ELC score increased by 13.12 for every unit increase of the age-removed 3’-SL. We further determined whether the observed positive association between 3’-SL and the MSEL ELC score was specific to particular cognitive subdomains by evaluating each MSEL subdomain score using Model 2. The Holm-Bonferroni method was used to correct for multiple comparisons (35). Significant and positive associations between 3’-SL and the receptive (adjusted P = 0.015; EF, 9.95; 95% CI, 3.91–15.99) and expressive (adjusted P = 0.048; EF, 7.53;
### TABLE 1  Demographic and human milk information of the participants

|                                      | Total          | A-tetra+        | A-tetra-         | P value$	extsuperscript{2}$ |
|--------------------------------------|----------------|-----------------|------------------|-------------------------------|
| **Subjects**                         | 99 subjects    | 33 subjects     | 66 subjects      |                               |
| Sex, male                            | 41             | 19              | 22               |                               |
| Age, months                          | 10.13 (3.98)   | 10.85 (4.21)    | 9.8 (3.84)       | 0.11                          |
| Birth weights, kg                    | 3.5 (0.42)     | 3.47 (0.48)     | 3.51 (0.39)      | 0.59                          |
| Birth lengths, cm                    | 51.56 (2.39)   | 51.39 (2.18)    | 51.63 (2.47)     | 0.57                          |
| Gestational ages, months             | 9.27 (0.25)    | 9.2 (0.19)      | 9.32 (0.28)      | 0.21                          |
| % of vaginal births                  | 0.88           | 0.8             | 0.93             |                               |
| Human milk volume, mL                | 55.75 (31.94)  | 53.63 (30.05)   | 56.78 (32.89)    | 0.53                          |
| Human milk weight, g                 | 56.29 (32.00)  | 52.97 (28.74)   | 57.87 (33.44)    | 0.33                          |
| Fat, g/100mL                         | 5.02 (1.73)    | 5.1 (1.73)      | 5.02 (1.74)      | 0.97                          |
| Carbohydrate, g/100mL                | 6.87 (0.66)    | 6.77 (0.89)     | 6.92 (0.47)      | 0.27                          |
| Energy, kcal/100mL                   | 78.21 (16.5)   | 77.57 (17.28)   | 78.59 (16.13)    | 0.74                          |
| Household income, USD, n             |                |                 |                  |                               |
| <25k                                 | 1              | 1               | 0                | 0.26                          |
| 25k–35k                              | 4              | 0               | 4                |                               |
| 35k–50k                              | 2              | 2               | 0                |                               |
| 50k–75k                              | 17             | 5               | 12               |                               |
| 75k–100k                             | 14             | 3               | 11               |                               |
| 100k–150k                            | 19             | 7               | 12               |                               |
| 150k–200k                            | 12             | 4               | 8                |                               |
| >200k                                | 9              | 5               | 4                |                               |
| Mother education (grad level proportion) | 0.61 (0.49) | 0.64 (0.48) | 0.59 (0.49) | 0.59                          |
| HMO concentrations$	extsuperscript{3}$ |                |                 |                  |                               |
| 2'-FL                                | 1276.72 (982.39) | 1444.67 (717.31) | 1200.75 (1075.16) | 0.07                          |
| 3-FL                                 | 1580.37 (1031.32) | 1340.37 (595.38) | 1688.93 (1162.62) | 0.01                          |
| A-tetra                              | 30.23 (75)     | 30.37 (39.78)   | 31.1 (50.59)     | 0.69                          |
| 3'-SL                                | 190.42 (99.76) | 202.22 (142.23) | 185.08 (73.04)   | 0.39                          |
| 6'-SL                                | 32.25 (47.4)   | 30.37 (39.78)   | 31.1 (50.59)     | 0.69                          |
| LNT                                  | 401.08 (264.17) | 397.6 (326.14)  | 402.65 (232.26)  | 0.92                          |
| LNNT                                 | 64.94 (65.97)  | 70.35 (65.76)   | 62.5 (66.18)     | 0.46                          |
| LNFP-I                               | 248.42 (286.92) | 256.08 (237.48) | 244.96 (307.5)   | 0.79                          |

1Values show means (SDs). Abbreviations: A-tetra, alpha-Tetrasaccharide; A-tetra-, undetectable alpha-Tetrasaccharide; A-tetra+, alpha-Tetrasaccharide positive; LNFP-I, lacto-N-fucopentaose I; LNT, lacto-N-tetraose; LNNT, lacto-N-neotetraose; USD, United States dollar; 2'-FL, 2'-fucosyllactose; 3-FL, 3-fucosyllactose; 3'-SL, 3'-sialyllactose; 6'-SL, 6'-sialyllactose.

2The P values are for the differences among subjects between A-tetra+ and A-tetra- groups using t-tests for all parameters, with the exception of household incomes, where the chi-square test of independence was used.

3Although our subjects had multiple visits, HMO concentration results shown in this table treated each sample independently.

95% CI, 2.51–13.79) language subscores were observed, with powers of 0.86 and 0.73, respectively, (Table 3). In other words, with other variables controlled, a unit increase in the age-removed 3'-SL was associated with 9.95 and 7.53 higher receptive and expressive language t-scores, respectively. In addition to applying Model 2, we also separately analyzed the A-tetra+ and A-tetra- subjects using Model 1. Similar results as those observed using Model 2 were found. Nevertheless, a single model (Model 2) comprising all data had more statistical power and was more robust than Model 1 when separately analyzing the A-tetra+ and A-tetra- groups. More detailed discussion is provided in Supplemental Information 4.

For the batch and site effects, subjects from batch 1 of Site B had a 9.38 significantly lower (P < 0.001; 95% CI, -13.90 to -4.96) ELC score when compared to those from batch 2 of Site B (Table 3). The gross motor and visual reception t-scores were 6.80 and 9.06 lower, respectively, in Site B subjects from batch 1 than in those from batch 2 (both were significant differences at adjusted P values < 0.001; 95% CI, -9.87 to -3.61 and -12.51 to -3.61).

### TABLE 2  Mullen Scales of Early Learning scores of the participants

|                                | Total (n = 183) | A-tetra+ (n = 57) | A-tetra- (n = 126) | P value$	extsuperscript{2}$ |
|--------------------------------|----------------|------------------|--------------------|-------------------------------|
| Composite score                | 106.18 (12.89) | 104.39 (12.71)   | 106.99 (12.94)     | 0.2                           |
| Gross motor                    | 51.58 (9.13)   | 53.53 (9.89)     | 50.7 (8.66)        | 0.07                          |
| Visual reception               | 53.6 (10)      | 52.37 (9.81)     | 54.16 (10.08)      | 0.26                          |
| Fine motor                     | 54.66 (11.08)  | 53.65 (9.42)     | 55.11 (11.76)      | 0.37                          |
| Receptive language             | 50.52 (9.75)   | 50.49 (10.14)    | 50.54 (9.61)       | 0.98                          |
| Expressive language            | 53.42 (8.89)   | 52.05 (8.95)     | 54.04 (8.82)       | 0.17                          |

1Values are shown as means (SDs). Although our subjects had multiple Mullen Scales of Early Learning assessments over time, the results shown in this table treated each visit independently. Abbreviations: A-tetra-, undetectable alpha-Tetrasaccharide; A-tetra+, alpha-Tetrasaccharide positive.

2The P values are for the differences among subjects between A-tetra+ and A-tetra- groups using t-tests.
3’-SL is associated with language development

FIGURE 2 The distributions of the 8 HMOs in mg/L with age. Each HMO has 256 samples from 114 subjects from the final HMO data set. The light and dark blue filled circles in the 2’-FL, A-tetra, and LNFP-I scatter plots indicate human milk below and above the LoD, respectively. About 70% (129 samples), 21% (39 samples), and 12% (22 samples) of the human milk contained A-tetra, LNFP-I, and 2’-FL, respectively, below the LoDs of these HMOs (4.4, 2.0, and 3.9 mg/L for A-tetra, LNFP-I, and 2’-FL, respectively). The y-axis represents the HMO concentration in mg/L. The x-axis represents age in months. The red lines represent the mean values at different ages for each HMO, excluding the undetectable values for 2’-FL, A-tetra, and LNFP-I. The HMOs are highly age dependent, with 2’-FL, LNFP-I, LNNT, LNT, and 6’-SL decreasing and A-tetra, 3-FL, and 3’-SL increasing with age. For 2’-FL, 6’-SL, and LNFP-I, age has a significant quartic (adjusted \( P \) values < 0.001 for all 3) effect, whereas 3-FL, A-tetra, 3’-SL, LNT, and LNNT exhibit significant quadratic (adjusted \( P \) values < 0.01 for all 5) effects. Abbreviations: A-tetra, alpha-Tetrasaccharide; HMO, human milk oligosaccharide; LNFP-I, lacto-N-fucopentaose I; LoD, limit of detection; LNNT, lacto-N-neotetraose; LNT, lacto-N-tetraose; 2’-FL, 2’-Fucosyllactose; 3-FL, 3-fucosyllactose; 3’-SL, 3’-sialyllactose; 6’-SL, 6’-sialyllactose.

The visual reception \( t \)-score was 5.85 lower in Site A subjects compared to the Site B subjects from batch 2 (significant at an adjusted \( P \) value of 0.02; 95% CI, -9.55 to -2.23). The results of the gross motor and visual reception \( t \)-scores are not included in Table 3.

We further studied whether the observed positive associations of 3’-SL and the language subdomain scores had interactions with age. Using the AIC together with attempts to balance the sample sizes of the 2 age subgroups, 12 months was chosen as the cutoff point. Figure 3 shows that among the A-tetra+ subjects, the associations of age-removed 3’-SL levels on the ELC scores were consistent across ages, but younger subjects had higher ELC scores for the same age-removed 3’-SL level compared to the older subjects. For the receptive language \( t \)-scores, the increments of the \( t \)-scores per age-removed 3’-SL level were significantly greater for the older subjects. Specifically, for 1 increase in the age-removed 3’-SL level, the receptive language \( t \)-score increased by about 1.82 and 16.74 for infants younger and older than 1 year old, respectively, resulting in a difference of 14.93 between the younger and the older groups (adjusted \( P = 0.03 \) for the interaction between age and age-removed 3’-SL; 95% CI, -25.29 to -4.24).

Discussion

The present study evaluated the potential associations between 8 HMOs and cognitive development during infancy. Based on the A-tetra level in HM that they received, children were stratified into A-tetra+ and A-tetra- subgroups. Levels of 3’-SL were significantly and positively associated with the ELC scores in A-tetra+ subjects, and the associations were specific to expressive and receptive language scores. Furthermore, larger effects of 3’-SL on receptive language were observed for children older than 12 months as compared to for those younger than 12 months. To the best of our knowledge, these are the first reported human results demonstrating positive associations between 3’-SL and cognitive development, particularly language development, during infancy.

Development of language ability during infancy

Because language is a trait only humans possess, it is not surprising that all previously reported preclinical studies were unable to uncover the potential benefits of HMOs on language function. The MSEL assesses 2 specific language functions: receptive and expressive language. Developmentally, it has been widely recognized that cognitive maturation follows different temporal orders depending on the functional domains (36–38). Therefore, we investigated a plausible age-related association between 3’-SL and language function. We found that the associations between expressive language and 3’-SL were statistically indistinguishable between infants younger and older than 12 months of age. In contrast, larger effects of 3’-SL on receptive language were observed for children older than
TABLE 3

Statistical results of associations between ELC, expressive language, and receptive language scores and HMOs.

| Batch 1 | Intercept | Site A 2'-FL | 3-FL | 3'-SL | 6'-SL | LNT | LNNT | LNFP-I | A-tetra | A-tetra+ |
|---------|-----------|--------------|------|-------|-------|-----|------|--------|---------|---------|
|         | Estimate  | -1.012       | 0.008 | 0.048 | 0.017 | 0.019| 0.0008| 0.0001 | 0.0003 | 0.0005 |
|         | SE        | 0.151        | 0.005 | 0.045 | 0.013 | 0.003| 0.003 | 0.0006 | 0.0004 | 0.0005 |
|         | t         | -6.674       | 0.003 | 0.038 | 0.028 | 0.019| 0.004 | 0.001  | 0.005  | 0.009  |
|         | P         | 0.011       | 0.000 | 0.000 | 0.000 | 0.000| 0.000 | 0.000  | 0.000  | 0.000  |

| Batch 1 | Intercept | Site A 2'-FL | 3-FL | 3'-SL | 6'-SL | LNT | LNNT | LNFP-I | A-tetra | A-tetra+ |
|---------|-----------|--------------|------|-------|-------|-----|------|--------|---------|---------|
|         | Estimate  | 13.119       | 0.008 | 0.048 | 0.017 | 0.019| 0.0008| 0.0001 | 0.0003 | 0.0005 |
|         | SE        | 0.151        | 0.005 | 0.045 | 0.013 | 0.003| 0.003 | 0.0006 | 0.0004 | 0.0005 |
|         | t         | 86.674       | 0.003 | 0.038 | 0.028 | 0.019| 0.004 | 0.001  | 0.005  | 0.009  |
|         | P         | 0.000       | 0.000 | 0.000 | 0.000 | 0.000| 0.000 | 0.000  | 0.000  | 0.000  |

| Batch 1 | Intercept | Site A 2'-FL | 3-FL | 3'-SL | 6'-SL | LNT | LNNT | LNFP-I | A-tetra | A-tetra+ |
|---------|-----------|--------------|------|-------|-------|-----|------|--------|---------|---------|
|         | Estimate  | -1.012       | 0.008 | 0.048 | 0.017 | 0.019| 0.0008| 0.0001 | 0.0003 | 0.0005 |
|         | SE        | 0.151        | 0.005 | 0.045 | 0.013 | 0.003| 0.003 | 0.0006 | 0.0004 | 0.0005 |
|         | t         | -6.674       | 0.003 | 0.038 | 0.028 | 0.019| 0.004 | 0.001  | 0.005  | 0.009  |
|         | P         | 0.011       | 0.000 | 0.000 | 0.000 | 0.000| 0.000 | 0.000  | 0.000  | 0.000  |

**Note:**
- Estimations are adjusted for batch number (Se, Le) and group (+, -).
- Bolded values indicate statistical significance at P < 0.05.
- SE values are calculated using the Holm-Bonferroni method.
- P-values are calculated using the Holm-Bonferroni method.

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**Secretor and Lewis statuses**

Differences in HMO secretion in mothers based on their Lewis blood group have been reported (42, 43). Among the blood groups, secretor-positive (Se+) and Lewis positive (Le+), mothers secreted all of the oligosaccharides examined, but secretor-negative (Se-), Lewis-negative (Le-), and Se+/Le- mothers did not secrete α1,2-fucosylated compounds and α1,4-fucosyloligosaccharides. Newburg et al. (44) further demonstrated that the differences in blood groups resulted in different compositions of HM and affected the immune system. To further determine whether the associations between HMOs and early cognitive development depend on the secretor and Lewis statuses, exploratory analyses were conducted. Results and discussion are provided in Supplemental Information 4. In short, given the limited sample sizes of the Se+/Le- and Se-/Le-groups (Supplemental Table 3), analyses focused on the Se+/Le+ and Se-/Le+ groups. No significant association was observed for Se-/Le+ group, whereas significant associations between 3'-SL and the ELC score (P = 0.02) and receptive language subscore (adjusted P = 0.045) were observed for the Se+/Le+ group using Model 1. We further evaluated whether these findings were independent of the A-tetra status using Model 2 for the Se+/Le+ group. Significant associations of 3'-SL with ELC (P = 0.001), receptive language (adjusted P = 0.015), and expressive language (adjusted P = 0.016) scores were observed in the A-tetra+ group. Therefore, the positive associations observed using Model 1 are most likely driven by the A-tetra+ subjects. Together, it appears that secretor and Lewis statuses do not account for A-tetra levels in our study population. Nevertheless, studies with larger and balanced sample sizes among the 4 Lewis blood groups are warranted to better understand the implications of genetic polymorphisms and early cognitive development.

**A-tetra+ and A-tetra-**

Austin et al. (26) reported that ∼16.7% of HM samples obtained from Chinese women were A-tetra+, and these women were all blood type A. In the context of our study, the percentage of the US population with blood type A ranges between ∼34% and 36% (45), which is consistent with our results, where 33.3% of mothers had A-tetra- HM, suggesting that the A-tetra level may be a proxy for blood type, and consequently HM grouping. However, it is worth noting that 2 women in our
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FIGURE 3  Relationship between 3'-SL and composite score, E.L., and R.L. by age groups ≤12 months (red) and >12 months (blue), respectively. Among the A-tetra+ group. Each point represents the samples from A-tetra+ subjects. There are 40 samples in the younger age group and 17 in the older group. The red and blue lines show the fitted linear regression lines using linear mixed effects models by age groups. The associations of age-removed 3'-SL on the composite score were consistent throughout ages, but younger subjects had higher composite scores for the same age-removed 3'-SL compared to the older subjects. For the R.L. t-scores, the increments of the t-scores per age-removed 3'-SL were significantly greater for the older subjects. Specifically, for 1 increase in the age-removed 3'-SL, the R.L. t-score increased by about 1.82 and 16.74 for infants younger and older than 1 year old, respectively, resulting in the difference of 14.93 between the younger and the older groups. The adjusted P value for the interaction term between age and age-removed 3'-SL was 0.03 (95% CI, -25.29 to -4.24). The P value for the interaction term for composite score was 0.538, and the adjusted P value for the interaction term for E.L. was 1. Thus, for the composite score and E.L., the effect of age-removed 3'-SL was consistent between age groups. The y-axis represents the MSEL t-score levels and the x-axis represents the residuals of 3'-SL. Abbreviations: A-tetra+, detectable alpha-Tetrasaccharide; E.L., expressive language; MSEL, Mullen Scales of Early Learning; R.L., receptive language; 3'-SL, 3'-sialyllactose.

study showed temporally varied A-tetra levels in the HM samples between visits. Therefore, other biological underpinnings leading to women secreting detectable compared with undetectable A-tetra should be considered. We offer hypothetical explanations. Glycosylation disorder, a defec tion of glucosidase I, was shown to be present in patients with accumulated A-tetra in their urine (46). Furthermore, soluble Klotho (sKlotho), which is an ectodomain shedding of antiaging membrane, was shown to affect the α2–3-sialyllactose in glycolipid gangliosides and possess cellular effects interfered by lipid rafts, where the α2–3-sialyllactose are supplemented (47). Thus, it is conceivable that both glucosidases and sKlotho play a role between women secreting detectable compared with undetectable A-tetra. Nevertheless, future studies will be needed to gain additional insights into how A-tetra concentrations may vary with time.

Beyond 3'-SL

Berger et al. (24) reported that higher exposures of 2'-FL relative to other HMOs at 1 month of age were associated with better cognitive ability, assessed at 24 months of age. Oliveros et al. (25) showed a positive association between 2'-FL levels at 1 month of age and 6-month motor scores, whereas 6'-SL levels were positively associated with cognitive and motor scores at 18 months. However, no association was found between 2'-FL nor 6'-SL and cognitive development in our study. Although many factors may explain the discrepancies between our results and those reported by these 2 studies, the differences in the study designs may account for the observed discrepancies. Specifically, both the Berger et al. (24) and Oliveros et al. (25) studies utilized a classic longitudinal design, whereas an accelerated longitudinal design was employed in our study. As a result, data from our cohort cannot be used to evaluate how differences in exposure to HMOs at early ages may lead to diverse cognitive outcomes at later ages. Instead, association analyses of 8 commonly reported HMOs and cognitive development were conducted in our study.

While we reported the results of 8 quantitative HMOs, HMOs with the 24 Maltotriose quantitative HMOs were also analyzed. Given the limited sample size in our study, exploratory analyses were conducted using variable selection via backward elimination. The main findings were similar to those reported above (Supplemental Information 5).

Limitations

Our study has several potential limitations. We did not collect data on either the time taken for the acquisition of HM samples after the last feeding nor the frequency of breastfeeding. Factors other than socioeconomic status and maternal factors, such as the first language spoken at home, multiple birth status, and maternal parity, might modulate children’s cognitive ability. Lack of diversity in the study population hampers the generalizability of our results. The use of an accelerated longitudinal design makes it difficult for prediction analyses. Insufficient age coverage hinders the discovery of any age interaction between 3'-SL and expressive language. The biological links between 3'-SL and language development remain elusive. Finally, since the study focuses on the role of oligosaccharides, our data cannot determine whether other milk components, such as fatty acids,
also influence cognitive development. Detailed discussions of these limitations are provided in Supplemental Information 6.

Conclusion

When children were stratified based on A-tetra levels in the HM that they received, a significant positive association between 3'-SL and the ELC score was observed in A-tetra+ subjects. This association was driven by the receptive and expressive language subdomain scores. Our results support the potential cognitive benefits of HMOs in the A-tetra+ subgroups of infants.

The authors’ responsibilities were as follows – BRH, JTE, WL: designed the research; KB, BRH, HCH, JTE: conducted the research; SC, ZZ, TL, DW, WL: analyzed data or performed the statistical analysis; SC, JH, NS, WL: wrote the paper; and all authors: had primary responsibility for the final content and read and approved the final manuscript. JH and NS are employees of Société des Produits Nestlé SA. WL is a consultant of and received travel support from Nestlé SA. SC, ZZ, TL, KB, BRH, HCH, JTE, and DW, no conflicts of interest.

Data Availability

Details described in the manuscript, code book, and analytic code will be made available upon request pending application and approval by the authors.

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