Original research article

Lectin-based analysis of human milk immunoglobulin G fucosylated variants in relation to milk maturation and perinatal risk factors

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A B S T R A C T

Background: Fucosylated glycotopes of milk immunoglobulin G (IgG) are ligands in reactions of biological recognition protecting newborns against infection and ensuring proper development.

Materials and methods: Relative amounts of IgG fucosyl-glycovariants in milk of mothers giving birth to term and premature newborns (term and preterm milk groups) were analysed by lectin-IgG-ELISA using α1,2-, α1,3-, and α1,6-fucose specific biotinylated Ulex europaeus (UEA), Tetragonolobus purpureus (LTA), and Lens culinaris (LCA) lectins, respectively.

Results: The term and preterm milk IgG glycovariants were highly reactive with UEA, LTA, and LCA, whereas maternal plasma IgG poorly or at all. During milk maturation the IgG of very preterm and preterm milk compared to term milk differed by lower relative amounts of UEA-, higher of LTA-, and nearly stable expression of LCA-reactive glycotopes. Moreover, lower α1,2- and higher α1,3- relative amounts of lectin-dependent milk IgG-fucosylated glycovariants were found to be associated with an infectious disease of lactating mothers.

Conclusion: The highly fucosylated glycovariants of human IgG given with mothers’ milk to immunologically immature newborn seem to be bifunctional molecules with potential therapeutic properties. The analysis of fucosylation status of milk IgG by simple lectin-IgG-ELISA may be helpful to control the immunological quality of milk for milk banking.

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Introduction

Among many human milk components supporting adequate growth and development of newborns during the first months of life (AAP, 2012; WHO, 2016a,b) the maternal immunoglobulins delivered to the babies are involved in naturally compensatory mechanisms of the unfavourable effects of neonatal immunodeficiency (Chirico et al., 2008). Immunoglobulin G (IgG) concentration in human milk is ranging from 20 to 117 mg/l (Arnold et al., 2007; Bondt et al., 2014) and is approximately 130–250 times lower than in human serum (5–15 g/l) (Arnold et al., 2005). Zhang et al. (2013) observed an increase of the relative abundance of IgG from week 1 to 12 months in ten milk samples from healthy mother. On the other hand, França et al. (2010, 2012) and Broadhurst et al. (2015) reported that the concentration of IgG did not differ significantly between the groups within any period of lactation, but pointed out the inherent physiological differences between mothers.

Several studies have demonstrated that human plasma IgG has three potential glycosylation sites situated in crystallisable fragment (Fc), antigen-binding fragment (Fab), and hinge region (Arnold et al., 2007; Bondt et al., 2014; Plomp et al., 2015). All IgG sub-isotypes contain a single N-linked glycan at Asn-297 of the Cy2 domain of Fc fragment. It is a complex-type biantennary glycan, whose pentasaccharide core is elongated by galactose and can be expanded by the addition of core fucose, and rarely by sialic acid (Arnold et al., 2007; Bondt et al., 2014; Shade and Anthony, 2013; Vidiarsson et al., 2014). The most abundant N-glycan of Fc region of IgG has a high level of core fucose (~94%) and low level of sialic acid (~10%) and bisecting GlcNAc (~10%) (Bondt et al., 2014).
Moreover, about 15–25% of plasma IgG contains a complex-type biantennary N-glycan at antigen-binding fragment (Fab) which is fucosylated (~69%), sialylated (~72%) and may contain bisecting GlcNAc (~45%) (Bondt et al., 2014; Vidarsson et al., 2014; Zauner et al., 2013). The third glycosylation place can be located in hinge region of human sera IgG3, and may contain three mono- or disialylated O-linked glycans (Plomp et al., 2015).

To date there are more than twenty IgG glycoforms (syn. glycovariants) described, whose mutual distribution is tissue-dependent and might be modified during growth and some diseases (Arnold et al., 2007; Gornik et al., 2012; Orczyk-Pawilowicz et al., 2012). The exposed residues of sialic acid and fucose on IgG are reported to take part in variety of biological functions, such as immunoregulation, inflammation progression, complement activation, opsonizing, mediation of antibody-dependent cellular cytotoxicity (ADCC) and can serve as ligand for bacterial adhesins (Bondt et al., 2014; Gornik et al., 2012; Thieker et al., 2016; Vidarsson et al., 2014; Zauner et al., 2013). Alterations of IgG glycosylation affect affinity binding of antigen and FcγR, antibody-mediated effector functions, release of pro-inflammatory factors (Arnold et al., 2007; Nimmerjahn and Ravetch, 2007), ADCC and antitumor activity as well as anti- and pro-inflammatory properties of IgG (Gornik et al., 2012; Kaneko et al., 2006; Shade and Anthony, 2013; Zauner et al., 2013). The core fucosylation α1,6 of IgG is known to have an influence on Fc region conformation. In consequence, a lack of a core fucose on IgG significantly decreases the ability of IgG to bind to FcγRIIa receptor on natural killer cells and macrophages. As suggested by Gornik et al. (2012) it leads to “safety switch” and in that way prevents elicitation of potentially destructive ADCC.

Fucosylated glycotopes are well known ligands in biological recognition. Alpha 1,2- and α1,3-linked fucoses are structural elements of the Lewis antigens, such as Lewis*, Lewis*, sialo-Lewis*, expressed on outer arms of N- and O-glycans. In contrast, α1,6-linked fucose is attached exclusively to the core part of N-glycans. The fucosylated glycotopes of IgG are essential for biological reactions, such as cell adhesion, signalling, cell growth and differentiation. All of the above events are particularly important during early human development and in protection of infants against infection (Becker and Lowe, 2003; Orczyk-Pawilowicz and Kątnik-Prastowska, 2011). Moreover, IgG may survive in the gastrointestinal tract of newborns and can serve as substitutes of S-IgA (Wilson and Ogra, 2011). This hypothesis is additionally supported by the fact that in bovine model, collostral IgG showed protective effect against enterotoxigenic E. coli (Freedman et al., 1998; Hurley and Theil, 2011).

In light of the significant role of fucosylated glycotopes in the biological recognition processes and since milk is not only food, but also specialized, high-quality medicine for premature newborns, we analysed expression of conformationally accessible fucose residues on human milk IgG, present in a milk sample in their native form, which in vivo might be recognized by endogenous intestinal receptors and/or bacterial lectins. In the study, the concentration of IgG and the expression of α1,6-, α1,3-, and α1,2-linked fucosyl residues on milk and plasma IgG of healthy and infected mothers giving birth to term and premature newborns were analysed in relation to lactation stages during human lactation from the 1st/2nd day to the 55th/40th day by lectin-based enzyme-linked immunoabsorbent assay (lectin-IgG-ELISA) using fucose-specific Lens culinaris (LCA), Tetragonolobus purpureus (LTA), and Ulex europaeus (UEA) agglutinins, respectively. Lectin-IgG-ELISA is a simple, quick and cost-effective method which allows for the simultaneous analysis of large number of samples without glycan separation steps as is required for high-performance liquid chromatography and mass spectrometry techniques.

**Materials and methods**

*Milk and plasma samples*

The milk samples called “term milk” (n = 186) were obtained from healthy lactating women (21–35 years old) who delivered healthy newborn at term (37 1/7–41 6/7 weeks of gestation) at the 1st Department and Clinic of Gynecology and Obstetrics at Wroclaw Medical University (Wroclaw, Poland). The milk samples called “preterm milk” (n = 106) were obtained from mothers (25–38 years old) who delivered preterm newborn (26 1/7–35 6/7 weeks of gestation) and who were treated at the Department and Clinic of Neonatology at Wroclaw Medical University (Wroclaw, Poland). Mothers who used tobacco products, illicit drugs or alcohol were not included in the study. Informed written consent was obtained from all mothers according the protocol approved by

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<th>Clinical characteristics of the preterm group.</th>
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<td><strong>Clinical parameter</strong></td>
<td><strong>Groups</strong></td>
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<tr>
<td>Gestation age</td>
<td>Extremely Preterm (EP) (26.5 ± 0.7 weeks)</td>
</tr>
<tr>
<td></td>
<td>Very Preterm (VP) (30.3 ± 1.7 weeks)</td>
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<td></td>
<td>Moderate Preterm (MP) (33.8 ±0.9 weeks)</td>
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<tr>
<td>Day of lactation</td>
<td>Early colostrum (1–3 days)</td>
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<td></td>
<td>Colostrum (4–7 days)</td>
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<td></td>
<td>Transitional milk (8–14 days)</td>
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<td></td>
<td>Mature milk (15–45 days)</td>
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<tr>
<td>Mode of delivery</td>
<td>Vaginal</td>
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<tr>
<td></td>
<td>Caesarean section</td>
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<tr>
<td>Mother’s status</td>
<td>Maternal infection</td>
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<tr>
<td></td>
<td>PROM</td>
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<tr>
<td></td>
<td>Hypertension</td>
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<tr>
<td>Newborn’s status</td>
<td>Birth weight (grams, mean ± SD)</td>
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<td></td>
<td>Antibiotic treatment (n)</td>
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<td>Hypertrophy (n)</td>
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According to WHO Preterm births (2016): EP, Extremely Preterm; VP, Very Preterm; MP, Moderately Preterm; PROM, Premature Rupture of Membranes.
Ethics Committee at Wroclaw Medical University (KB-30/2013 and KB-411/2015).

The term milk group consisted of mothers (n = 186), without signs of infection, giving birth to term single healthy newborns weight from 2650 to 3900 g (3360 ± 349 g), without genetic disorders. 83% of term milk samples were from mothers with secretor status. 52% of newborns in the “term” group were born by caesarean section.

The preterm milk group comprised of mothers (n = 40) with single pregnancy (Table 1) who delivered newborns assigned (WHO Preterm births, 2016) to the groups Extremely Preterm (EP: <28 0/7 weeks of gestation, n = 5), Very Preterm (VP: 28 1/7–32 0/7 weeks of gestation, n = 41), and Moderate Preterm (MP: 32 1/7–36 0/7 weeks of gestation, n = 60) and 73% of them had secretor status. Among mothers giving birth to premature newborns 21 mothers suffered from infections, 14 had a premature rupture of membranes before parturition, and 9 had hypertension during pregnancy.

To obtain plasma samples, blood samples from 40 lactating women on the 2nd postpartum day (n = 40) were taken with 3.2% sodium citrate.

Sample collection and preparation

The samples of human milk of term and preterm groups (from 1st to 55th and from 2nd to 40th days of lactation, respectively) were collected by a lactation consultant from the breast by manual expression and/or a breast pump at the end of nursing (hindmilk) by complete breast emptying, once per day, at the same time (8:00–10:00 a.m.). There was significant interindividual variation in volume of hindmilk, from 1 ml for early colostrum to 100 ml for mature milk. All milk samples were frozen and stored immediately at –20 °C until analysis. Skim milk (aqueous phase) was prepared by centrifugation at 3500g at 4 °C for 35 min, after which the fat layer and cells were removed. The aliquots of skim milk and plasma samples were stored at –20 °C until analysis.

The milk samples were divided into: (1) early colostrum (postpartum days 1–3; n = 40 for term and n = 6 for preterm milk), (2) colostrum (postpartum days 4–7; n = 75 for term and n = 15 for preterm milk), (3) transitional milk (postpartum days 8–14; n = 43 for term and n = 30 for preterm milk), (4) mature milk (postpartum days 15–55; n = 28 for term and n = 55 for preterm milk).

Additionally, the preterm milk samples were divided into: samples from mothers suffering an infectious disease (n = 50) and samples from mothers without infection (n = 56). Maternal urinary and upper respiratory tract infections were defined based on a positive microbiological culture and CRP higher than 100 mg/l.

IgG concentration

IgG concentration was determined by “sandwich type” ELISA using: 1) F(ab’)2 fragment of goat anti-human IgG (Jackson ImmunoResearch, USA) as a capture antibody, which extracted specifically IgG from a sample, 2) phosphatase-labelled rabbit anti-human IgG Fcγ fragment specific antibodies (Jackson ImmunoResearch, USA). For testing 100 μl of 1000-, 2500- and 5000-fold diluted milk, 400000- and 800000-fold diluted plasma, and IgG standard preparation from 0.2 to 12.5 ng/100 μl (Jackson ImmunoResearch, USA) were taken. The test was assayed with 4-nitrophenyl phosphate (SERVA, Heidelberg, Germany) as the enzyme substrate and absorbance was measured in a Stat Fax 2100 Microplate Reader (Awareness Technology Inc., Palm City, FL, USA) at 405 nm with 630 nm as the reference filter. All ELISA immunobinding and washing steps were carried out in TRIS-buffered saline (TBS, pH 7.5) containing 0.2% Tween 20.

Lectin-reactive fucosylvariants of IgG

Fucose expression on IgG was determined by a slightly modified lectin-IgG-ELISA (Orczyk-Pawilonicz et al., 2012) using specific biotinylated lectins (Vector Laboratories Inc., Burlingame, USA): Lens culinaris lectin (LCA), Tetragonolobus purpureus lectin (LTA), and Ulex europaeus lectin (UEA) showing binding preferences to fucose (Table 2) (Wu et al., 2009) linked by glycosidic linkages a1,6-, a1,3-, and a1,2-, respectively. However, the specificity of used lectins is not absolute and can be broader than just to fucose. F(ab’)2 fragment of goat anti-human IgG (Jackson ImmunoResearch, USA) diluted 1:1000 in 10 mM TBS, pH 8.5 showing a lack of reactivity with increasing concentrations of fucose-specific lectins in preliminary experiments, was used to coat a polystyrene microtitre ELISA plate, extract and specifically bind IgG from a sample. In preliminary experiments the amount of IgG as well as a concentration of lectins used in Lectin-IgG-ELISA fulfilling the conditions of proportionality have been selected. For the test, 100 μl of milk and plasma samples were taken which were pre diluted in 10 mM TBS pH 7.5, 10 mM CaCl2, 10 mM MgCl2, 0.05% Tween 20 and 0.5% glycerin, to a IgG concentration of 2 mg/l (200 ng per well). The presence of the respective fucosyl residues on IgG was detected by the reaction with the specific biotinylated lectins LCA (1:5000), LTA (1:5000), and UEA (1:500), respectively. The relative amount of formed IgG-lectin complex was determined with phosphatase-labelled ExtrAvidin (1:20000) (Sigma, St. Louis, MO, USA), and then detected by the reaction with 4-nitrophenyl phosphate. The absorbance (AU) was measured in a Stat Fax 2100 Microplate Reader. The absorbance values were found to be proportional to the increasing amount of fucose residue on IgG reactive with specific lectin.

All samples were analysed in duplicate. Controls were performed to demonstrate the specificity of the lectins as well as the absence of detectable endogenous reactive materials. The positive control for LCA, LTA, and UEA was a native haptoglobin and an asialo-haptoglobin and an asialo-haptoglobin preparation derived from ovarian cancer fluid (Kaźników et al., 1994). The negative control was an albumin preparation included in the test instead of milk and plasma samples. The background absorbances (TBS included instead of lectin, ExtrAvidin-AP, and samples, respectively) ranged from 0.04

<table>
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<th>Table 2</th>
<th>Major binding specificities of used lectins.</th>
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<tr>
<td>Origin and used abbreviation of lectin</td>
<td>Lens culinaris (LCA)</td>
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<tr>
<td>Binding preference</td>
<td>Fucα1,6-linked to proximal GlcNAc of the trimannosyl core of biantennary N-glycan</td>
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to 0.08 depending on the microtiter plate, the lectin used, and the day of the experiment.

**Statistical analysis**

The statistical analysis was performed with STATISTICA 12 (StatSoft, Inc., Tulsa, USA). The Kruskal–Wallis test was used for statistical significance. The results are shown as the mean ± SD and the median with 25th–75th percentiles. The correlations were estimated according to Spearman. A two-tailed p-value of less than 0.05 was considered significant.

**Results**

**IgG concentration**

The mean value of IgG concentration in milk samples of mothers (Fig. 1A) giving birth to a newborn at term (T group: 18.61 ± 15.6 mg/l), very preterm (VP group: 16.43 ± 6.1 mg/l), and moderate preterm (MP group: 20.06 ± 13.0 mg/l) was at similar level.

The mean value of IgG concentration was high in the early colostrum of the term (27.85 ± 23.2 mg/l) and preterm (41.74 ± 17.3 mg/l) groups, and then it significantly decreased in the colostrum of term (14.89 ± 12.5 mg/l, p < 0.003) and preterm milk (17.31 ± 10.1 mg/l, p < 0.02) groups. During further milk maturation IgG concentration was at stable similar level in the transitional as well as mature samples of the term (16.84 ± 11.9 and 18.06 ± 7.4 mg/l, respectively) and preterm milk groups (15.59 ± 8.4 and 17.49 ± 7.8 mg/l, respectively) (Table 3).

The concentration of term and preterm milk IgG showed no correlation with the day of lactation from the 1st/2nd day to the 55th/40th day, respectively (Fig. 2A). However, the concentration of IgG in milk groups showed individual differences and it ranged from the highest observed values of 99.6 mg/l at the 3rd day for term milk to the lowest value of about 4 mg/l at 8th day of lactation for term milk. The IgG concentration for the term and preterm milk group (mean value 18.61 ± 15.7 and 18.30 ± 10.9 mg/l, respectively) was about 280 times lower than in plasma of mothers at the 2nd postpartum day (5.17 ± 2.4 g/l).

**UEA-reactive α1,2-fucosylated variant of milk IgG**

The mean value of relative UEA reactivity with IgG of term milk samples (Fig. 1B) (0.53 ± 0.3 AU) was higher than the values for the very (0.38 ± 0.3 AU, p < 0.0006) and moderate (0.41 ± 0.2 AU) preterm milk groups.

The relative reactivity of UEA with term and preterm milk IgG showed the highest similar mean values in the early colostrum group (0.70 ± 0.3 and 0.78 ± 0.7 AU, respectively). During progression of lactation, the reactivity decreased in the colostrum group of the term milk (0.51 ± 0.3 AU; p < 0.004) and preterm milk (0.49 ± 0.2 AU). In subsequent stages of milk maturation, the reactivity remained almost unchanged in the term and preterm groups of transitional (0.46 ± 0.2 and 0.40 ± 0.2 AU, respectively) and mature (0.44 ± 0.2 and 0.37 ± 0.3 AU, respectively) milk groups. On the other hand, only the UEA reactivity with IgG of the term mature milk group (0.44 ± 0.2 AU) showed a significant higher (p < 0.02) mean value when compared with the reactivity for the mature preterm milk group (0.37 ± 0.3 AU) (Table 3).
The expression of UEA-reactive glycotopes on IgG showed a weak negative correlation with lactation from 1st/2nd to 55th/40th days in the term (r = -0.28; p < 0.05) and preterm (r = -0.31; p < 0.05) groups (Fig. 2B).

**LTA-reactive α1,3-fucosylated variant of milk IgG**

The mean value of LTA relative reactivity with IgG of milk (Fig. 1C) in the very (VP, 0.59 ± 0.3 AU, p < 0.003) and moderate (MP, 0.58 ± 0.4 AU, p < 0.03) preterm milk groups was higher than in the term milk group (0.45 ± 0.4 AU).

The relative reactivity of LTA with milk IgG was the lowest at the beginning of lactation in the early colostrum samples of term (0.28 ± 0.3 AU) and preterm (0.21 ± 0.3 AU) groups and it increased in the groups of colostrum of the term (0.50 ± 0.5 AU, p < 0.02) and preterm milk (0.54 ± 0.4 AU). During further milk maturation stages, the LTA reactivity with IgG of the term and preterm milk samples of transitional (0.42 ± 0.3 and 0.71 ± 0.4 AU, respectively) and mature milk (0.57 ± 0.3 and 0.54 ± 0.3 AU, respectively) did not change significantly compared with the values for the colostrum groups (Table 3). However, the high reactivity of LTA with milk IgG of transitional milk samples of the preterm group (0.71 ± 0.4 AU) was found to be significantly higher (p < 0.0002) than in the corresponding term milk group (0.42 ± 0.3 AU).

The expression of LTA-reactive glycotopes of IgG showed a weak positive correlation (r = 0.27; p < 0.05) with a lactation for the term milk group, whereas that of the preterm milk group did not (Fig. 2C).

**LCA-reactive α1,6-fucosylated variant of milk IgG**

The relative reactivity of LCA with IgG of milk (Fig. 1D) of mothers giving a newborn at term (T group: 1.16 ± 0.4 AU) and for those of the VP and MP preterm milk groups were found to be nearly at the same level (1.05 ± 0.3 and 1.10 ± 0.4 AU, respectively).

In the early colostrum group of term and preterm milk samples the relative reactivity of LCA with IgG was high (1.18 ± 0.4 and 1.07 ± 0.3 AU, respectively) and remained almost at the same level in the colostrum samples (1.24 ± 0.4 and 1.12 ± 0.5 AU, respectively). Following the period of 1st–7th days of lactation the reactivity of LCA with IgG decreased in transitional term milk group (1.07 ± 0.3 AU, p < 0.04), whereas for the preterm group was almost unchanged (1.07 ± 0.4 AU), and remained at the same level in the mature milk group of the term and preterm samples (1.05 ± 0.4 and 1.06 ± 0.3 AU, respectively) (Table 3, Fig. 2D). In contrast, 20 ng of lactating mothers’ plasma IgG was very weakly recognized by LCA (0.1 ± 0.01 AU) but not by UEA and LTA.

**Lectin-reactive fucosylated IgG variants in preterm milk in relation to infection of mothers**

The mean values of the relative reactivity of UEA and LTA with IgG of preterm milk of mothers having infections (Fig. 3A, B) were significantly lower (0.38 ± 0.3 AU, p < 0.009) and higher (0.69 ± 0.4 AU, p < 0.002), respectively than those values found for the preterm milk group without infection (0.48 ± 0.3 and 0.47 ± 0.4 AU, respectively).

**Discussion**

In spite of the fact that the binding specificity of lectins is not absolute and their using did not allow to determine the “true” glycan structures, they are especially helpful to observe the
Fig. 2. IgG concentration (A) and reactivity of fucose-specific lectins UEA (B), LTA(C) and LCA (D) with milk IgG over lactation of mothers giving birth to term and preterm newborns. A solid line indicates linear regression, and 95% confidence intervals are shown by dotted lines. Correlation coefficient (r) was calculated with lactation days according to Spearman and p-value lower than 0.05 was regarded as significant. R² value is the square of the correlation coefficient of the linear regression between the day of lactation and lectin reactivity [AU] or IgG concentration (mg/l). For details see under Fig. 1.
changes of terminal sugar expressions on glycoconjugates which in vivo might be ligand for endogenous selectins and/or bacterial lectins. Our results showed that during lactation human milk IgG was highly reactive, in contrast to poorly reactive maternal plasma IgG, with UEA, LTA, and LCA, lectins specific to fucose linked to oligosaccharide part by α1,2-, α1,3-, and α1,6-linkages, respectively (Table 3). The fucosylation pattern of IgG of the very preterm and preterm milk samples differed from those of the term milk samples by the lower α1,2-, higher α1,3-, and nearly stable α1,6- linked fucose expressions (Fig. 1). Moreover, the lower amounts of α1,2- and higher of α1,3-lectin-dependent milk IgG fucosylated glycovariants were found to be associated with infection of lactating mothers (Fig. 3).

The composition of human milk after preterm delivery is reported to be altered compared with full-term pregnancy (Mehta and Petrova, 2011; Molinari et al., 2012), and contains significantly higher concentrations of some immune proteins than term milk (Broadhurst et al., 2015; Trend et al., 2016). The highest concentration of IgG in the early colostrum (Table 3) seem to be a result of milk compaction at the beginning of lactation and in the case of preterm milk might be associated with abnormalities during the perinatal period (Table 1). In spite of above and in contrast to the total protein, lactoferrin, and S-IgA levels during lactation (Froehlich et al., 2010; Lönnerdal, 2013; Trend et al., 2016), the concentration of IgG was at relatively stable level of 18 mg/l in term and preterm milk. That value is consistent with that of about 20 mg/l given by Broadhurst et al. (2015) and lower than established by Koenig et al. (2005). Observed a large variation of IgG concentration among lactating mothers seems to be associated with the inherent physiological differences between individual mothers, ethnicity, diet, age, lifestyle choices, and the presence of subclinical disease (Broadhurst et al., 2015; Yuen et al., 2012).

Lectin-based analysis reveals the presence of α1,6-, α1,2, and α1,3- fucosylated IgG glycovariants in milk and their absence or negligible amount on lactating mothers’ plasma IgG (Table 3). Such significant differences may result from, at least partially, local milk IgG synthesis in mammary gland hormonally regulated (Hurley and Theil, 2011). However, the strong reaction of milk IgG with LCA, in contrast to very weak reaction of plasma IgG, may reflect some structural differences and altered accessibility of α1,6-linked fucose residue on milk and plasma IgG for the reaction with lectin. Shade and Anthony (2013) reported that oligosaccharides of Fc fragment of plasma IgG were concealed in the cavity between two Cy2 domains and thus, in our opinion, core α1,6-linked fucose of plasma IgG was not recognized by LCA in lectin-ELISA. Moreover, the milk IgG may have some fucosylated glycans attached to Fab fragment, which were not found on plasma IgG, but appeared for example at the Fab portion during pregnancy (Bondt et al., 2014) and myeloma IgG (Kinoshita et al., 1991).

In contrast to the relatively stable expression of α1,6- fucosylated glycopeptide on term and preterm milk IgG during milk maturation, the pattern of changes in the expression of α1,2- and α1,3-fucosylated glycopeptides on IgG was variable for UEA- and LTA-reactive glycovariants. The highest expression of glycopeptide α1,2-fucosylated, observed several hours after delivery in term and preterm early colostrum IgG (Table 3), although lower in the very and moderate preterm milk samples than in term milk group (Fig. 1B), decreased with milk maturation (Table 3, Fig. 2B). In contrast, the relative amount of LTA-reactive milk IgG glycovariant containing an α1,3-fucose, a part of inflammation-related LewisX antigen, was evidently higher in milk samples derived from very preterm and moderate preterm groups than in samples of the term group (Fig. 1C). However, for term and preterm group its expression was the lowest at the beginning of lactation, and it reached the highest level in the preterm transitional milk group (Table 3) formed in 50% of mothers who had an infection. In our opinion, the expression of LTA-reactive α1,3-fucosylated LewisX antigen was mainly associated with infection, in women who delivered preterm (Fig. 3B) and to a lesser degree with delivery-associated inflammatory phase, as well as postpartum hormonal imbalance (Groër et al., 2005). The high expression of α1,3-fucosylated glycopeptide on preterm milk IgG of the infection group correlated with the decrease of α1,2-fucosylated glycopeptide (Fig. 3A, B). That fact might be associated with the synthesis of sialyl-LewisX antigen, whose constrains the attachment of fucose through α1,2-linkage to outer antenna of N-glycans (Bieberich, 2014) and occurrence was related to inflammation (Van Dijk et al., 1998).

The described fucosylation pattern for milk IgG characteristic for term and preterm milk maturation overlapped in part with the trends for glycoconjugates (Nwosu et al., 2012), fucosylated glycovariants of fibronectin (Orczyk-Pawilonicz et al., 2015), α-1-acid glycoprotein (Orczyk-Pawilonicz et al., 2014), and that
for HMOs (De Leoz et al., 2012). The differences in fucosylation of term and preterm milk IgG during lactation might be related to immaturity of mammary gland of preterm delivered mothers and with interindividual variability. According to De Leoz et al. (2012) the lower and more variable fucosylation of HMOs of preterm milk than that of term milk is associated with dysregulation of breast cells (insufficient levels of hormones and not fully completed remodelling of mammary gland tissue) during preterm lactation. IgG acts not only as specific antibody, but due to the fucose exposed on surface is able to participate in reactions of biological recognition (Arnold et al., 2007; Bode, 2012; Newburg, 1996). The presence of the LCA-reactive α1,6-fucosylated IgG glycovariant in human milk is believed to have importance in immunoprotection of the incompletely developed immune system of newborns, namely differentiation of the intestinal epithelium of newborns and cell-cell signalling and adhesion (Becker and Lowe, 2003; Jakaitis and Denning, 2014). The high α1,2- and α1,6-fucosylation of some milk glycoproteins is reported to be associated with anti-pathogenic function of fucosylated glycotopes (Kątnik-Prastowska and Orczyk-Pawilowicz, 2011; Orczyk-Pawiłowicz et al., 2014; Royle et al., 2003) and might participate as a decoy for fucose-dependent pathogens via lectin-carbohydrate interactions and prevent adhesion to host epithelial cells. The maturation of immune and gastrointestinal systems of newborns starts during fetal development and is modulated by amniotic immunological components. Provision of maternal milk to newborns is a continuation of the immunoprotective and immunomodulatory effect of amniotic fluid (Goldblum and Hilton, 1999; Wagner, 2002).

Conclusion

Biologically active highly fucosylated milk IgG seems to constitute bifunctional molecules which are able to act as specific antibodies and moreover, thanks to their exposed monosaccharide determinants, to act as decoy for fucose-dependent pathogens via lectin-carbohydrate interactions and prevent adhesion to host epithelial cells. The maturation of immune and gastrointestinal systems of newborns starts during fetal development and is modulated by amniotic immunological components. Provision of maternal milk to newborns is a continuation of the immunoprotective and immunomodulatory effect of amniotic fluid (Goldblum and Hilton, 1999; Wagner, 2002).

Conflict of interests

The authors declare that they have no conflict of interests.

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