Bacterial immunogenic $\alpha$-galactosylceramide identified in the murine large intestine: dependency on diet and inflammation

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Abstract The glycosphingolipid, $\alpha$-galactosylceramide ($\alpha$GalCer), when presented by CD1d on antigen-presenting cells, efficiently activates invariant natural killer T (iNKT) cells. Thereby, it modulates immune responses against tumors, microbial and viral infections, and autoimmune diseases. Recently, the production of $\alpha$GalCer by Bacteroidetes from the human gut microbiome was elucidated. Using hydrophilic interaction chromatography coupled to MS, we screened murine intestinal tracts to identify and quantify $\alpha$GalCers, and we investigated the $\alpha$GalCer response to different dietary and physiologic conditions. In both the ecum and the colon of mice, we found 1–15 pmol of $\alpha$GalCer per milligram of protein; in contrast, mice lacking microbiota (germ-free mice) and fed identical diet did not harbor $\alpha$GalCer. The identified $\alpha$GalCer contained a $\beta$-($\beta$)-hydroxylated hexadecanoyl chain $\n$-linked to C18-sphinganine, which differed from what has been reported with Bacteroides fragilis. Unlike $\beta$-anomeric structures, but similar to $\alpha$GalCers from B. fragilis, the synthetic form of the murine $\alpha$GalCer induced iNKT cell activation in vitro. Last, we observed a decrease in $\alpha$GalCer production in mice exposed to conditions that alter the composition of the gut microbiota, including Western type diet, colitis, and influenza A virus infection. Collectively, this study suggests that $\alpha$GalCer is produced by commensals in the mouse intestine and reveals that stressful conditions causing dysbiosis alter its synthesis. The consequences of this altered production on iNKT cell-mediated local and systemic immune responses are worthy of future studies.—von Gerichten, J., D. Lamprecht, L. Opála, D. Soulard, C. Marsching, R. Pilz, V. Sencio, S. Herzer, B. Galy, V. Nordström, C. Hopf, H-J. Gröne, F. Trottein, and R. Sandhoff. Bacterial immunogenic $\alpha$-galactosylceramide identified in the murine large intestine: dependency on diet and inflammation. J. Lipid Res. 2019. 60: 1892–1904.

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Besides recognition of classical major histocompatibility complex class I or II molecules presenting oligopeptides by T lymphocytes, a wide variety of nonclassical nearly monomorphic major histocompatibility complex molecules, including CD1, may be recognized by $\alpha$β and $\gamma$δ T cells as

Abbreviations: APC, antigen-presenting cell; CID, collision-induced dissociation; DC, dendritic cells; DSS, dextran sodium sulfate; FT-ICR, Fourier transformation-ion cyclotron resonance; GalCer, galactosylceramide; $\alpha$GalCer, $\alpha$-galactosylceramide; $\alpha$GalCerA, $\alpha$-galactosylceramide (d(17/18)/19:0/βH17:0) from Bacteroides fragilis, which incorporates three isobranched sphingoid bases with a total of 17, 18, and 19 carbon atoms; $\alpha$GalCerA10, $\alpha$-galactosylceramide(d18/0/βH16:0) from murine large intestine; GlcCer, glucosylceramide; HexCer, hexosylceramide; HILIC, hydrophilic interaction chromatography; IAV, influenza A virus; IL-2, interleukin-2; iNKT, invariant natural killer T; KRN7000, $\alpha$-galactosylceramide(d18/0/26:0); LCB, long chain base (of ceramide); McL, McLaFerty (fragment); MLL, mouse large intestine; MRM, multiple reaction monitoring; NKT, natural killer T; RP18, reversed phase 18-Catoms; TCR, T cell receptor; WTD, Western type diet.

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well as natural killer T (NKT) cells. These “unconventional” T cells recognize lipids, metabolites, and modified peptides (1). The glycosphingolipid, α-galactosylceramide (αGalCer), presented by CD1d of antigen-presenting cells (APCs) is one of the most potent stimulators of invariant NKT (iNKT) cells (2, 3). The interaction of the presented lipid-CD1d complex with the T cell receptor (TCR) of iNKT cells can activate these cells and trigger the release of a variety of cytokines and chemokines modulating immune responses (4, 5).

Originally, αGalCer was found in the marine sponge, Agelas mauritianus, in 1993 (6), and still, to date, very little is known about natural contact of mammalian organisms with this sphingolipid either through microbial sources or by endogenous production. Later, Bacteroides fragilis, a commensal of the human gut, was demonstrated to produce αGalCer (αGalCer_H) (7). Based on this work, two more strains of the human gut microbiome, Bacteroides vulgatus and Prevotella copri, were demonstrated to produce αGalCer (8). Recent work has shown that early exposure to αGalCer_H reduces iNKT cell number in the mouse colon, thereby protecting against colitis (9). Hence, there is a need to identify structure-specific sphingolipids in the gut microbiome to elaborate the role of bacterial sphingolipids in the host-immune response.

As mentioned, the immunogenic αGalCers belong to the class of sphingolipids that all contain a long chain base (LCB) 2-amino alcohol. In most cases, the LCB is amide linked with a fatty acyl chain to form ceramides, which often are modified at C1-position to carry a head group, such as various glycans, phosphorylcholine, phosphate, or acyl chains. The LCB as well as the fatty acyl residue can vary in their chain lengths, methylation, and degree of saturation and hydroxylation, and thereby create a huge variety of ceramide backbones, which contribute additionally to the variation and complexity of the sphingolipid class (10, 11). Sphingolipids are known to play important roles in proliferation, apoptosis, differentiation, and migration of eukaryotic cells (12–16). In contrast, only a few bacteria are known to produce sphingolipids (17–19). Among them is B. fragilis, which has recently been described to produce αGalCer with a C17, C18, or C19 iso-branched sphinganine LCB and a β-hydroxylated C17 iso-branched fatty acyl residue (7, 9). These structures were also present in Bacteroides vulgatus and P. copri (8). Human pathology with diseases such as diarrhea, inflammatory bowel disease, and cancer were connected to the abundance of B. fragilis in the human gut (7, 20–22). In the mouse system, there is evidence that the gut microbiota can produce iNKT cell agonists (23, 24). However, the nature of these agonists has not yet been clearly defined.

MS has become the most accurate tool to measure lipids. Although, structural stereoisomers often cannot be separated by MS alone, the combination of MS with LC can be suited to distinguish them. Our group recently published a method to successfully separate α- and β-anomeric isomers of glucosylceramide (GlcCer) and GalCer (8). This method has now enabled us to screen mouse intestinal tissue and content for hexosylceramide (HexCer) structures with an α-glycosidic linkage produced by bacteria. In the current report, we describe the identification of an αGalCer structure originating from the gut microbiome. This αGalCer was absent in microbiota-deficient mice. Because the low concentration of this compound did not allow us to obtain sufficient purified material for structural NMR analysis, we chemically synthesized the assumed structure in an α- and β-anomeric form. This allowed us to reveal the differential behavior of these compounds in hydrophilic interaction chromatography (HILIC) and MS collision-induced dissociation (CID) similar to previously described differential CID-fragmentation patterns (7, 25–28). By that, we subsequently confirmed the mouse intestinal tract-derived compound αGalCer(d18:0/βH16:0) containing C18-sphinganine and a β-hydroxylated palmitic acid. Furthermore, this α-anomeric compound turned out to stimulate iNKT cells in a similar manner as the αGalCers from B. fragilis. We further show that the production of αGalCer(d18:0/βH16:0) largely fluctuates according to the diet, local inflammatory conditions, and infection. These data imply that alteration of the microbiome composition impacts strongly on αGalCer production.

MATERIALS AND METHODS

Sphingolipids and reagents

αGalCer(d18:1/16:0), βGalCer(d18:1/16:0), αGalCer(d18:0/16:0), βGalCer(d18:0/16:0), βGalCer(d18:1/βH18:0), αβGalCer(d18:1/αS,h18:0), αβGalCer(d18:1/24:1), ββGalCer(d18:1/16:0), C18-α-galactosylsphingosine, and C18α-glucosylsphingosine were purchased from Avanti Polar Lipids. Kerasin (nonhydroxylated acyl chain containing βGalCers from bovine brain) and phrenosin (α-hydroxylated acyl chain containing βGalCers from bovine brain) were from Matreya. C18-β-galactosylsphingosine (psychosine) was purchased from Abcam. Racemic β-hydroxypalmitic acid, 1-hydroxybenzotriazole hydrate, 4-dimethylaminopyridine, anthrone, and N-(3-dimethylaminopropyl)-N′-ethylcarbodiimide were provided by Sigma-Aldrich. A mix of βGlCer(d18:1/14:0, 19:0, 25:0) and βGalCer(d18:1/31:0) was used as internal standard. These internal standard compounds had previously been chemically synthesized from the corresponding commercially available βGlCer and βGalCer and were used for quantification without further purification. Hexamethyldisilazane and 1-(3-dimethylaminopropyl)-N′-ethylcarbodiimide were purchased from VWR International; ammonium acetate and ammonium formate (99–100%) was purchased from Sigma-Aldrich.

Solvents and additives

Anhydrous (dry) tetrahydrofuran was purchased from Sigma-Aldrich. Chloroform, methanol, isopropanol, water, all LC-MS Chromasolv®-grade, and 2-butanol (>99.5%) for GC were from Honeywell Specialty Chemicals Seelze GmbH; propionitrile for synthesis was from Merck Darmstadt; formic acid (99–100%) was from VWR International; ammonium acetate and ammonium formate (≥99%) for MS-analytics were from Sigma-Aldrich.

2 Ceramide anchor description of hexosylceramides: In GalCer(d18:0/βH16:0), the first code (d18:0) refers to the sphingoid base (LCB), in this case with two hydroxyl groups (d), 18 carbon atoms, and no double bonds (0), and the second code (βH16:0) refers to the amide linked acyl chain (fatty acyl) containing, in this case, one hydroxyl group (h) in β-position, 16 carbon atoms and no double bond (0).
Mice
Tissue was isolated from euthanized C57BL/6N, C57BL/6J, and NMRI mice. Mice were obtained from the animal facilities of the German Cancer Research Center, the EMBL in Heidelberg, and the Pasteur Institute in Lille, as well as Charles River, Sulzfeld. Except for germ-free mice (NMRI-held in the animal facility of the German Cancer Research Center), all mice were kept under specific pathogen-free conditions. All animal procedures were approved by the respective governmental institutions and performed in accordance with federal laws. Duodenum was cut out 2 cm following stomach, and jejunum was taken from the middle part (1 cm) of caeca. With a distance of 2 cm from cecum, colon tissue samples with the length of 1 cm were cut and immediately stored at −80°C. Feces were obtained from the rectum. C57BL/6 mice, backcrossed at least 10 times in C57BL/6, were a gift from Dr. L. Van Kaer (Vanderbilt University, Nashville, TN).

Bacteria
According to previous publication (31), B. fragilis (DSM-2151) was grown anaerobically on Columbia agar plates with 5% sheep blood for 48 h. One culture plate was harvested and transferred in a tube containing 200 μl of methanol.

Chemical synthesis of hexosylceramides
For the synthesis and quantification of αGlcCer, αGalCer, and βGalCer standards carrying a β-hydroxylated fatty acid, see the supplemental Methods.

Lipid extraction
For the enrichment of HexCers containing GalCer(d18:0;h16:0) from a larger amount of mouse caeca and lipid extraction from small amounts of mouse tissue as well as lipid extraction from bacteria, see the supplemental Methods.

HILIC- and reversed-phase LC-MS² analysis of GalCer
Aliquots corresponding to 10 μg of mammalian tissue dry weight or rather 2 μg of dried bacteria were mixed with internal lipid standards for analysis by LC-MS² using an Acquity I-class UPLC and a Xevo TQ-S “triple quadrupole” instrument, both from Waters. Using a CSH C18 column (2.1 × 100 mm, 1.7 μm; Waters), lipids were measured in reversed-phase-LC mode with a gradient between 57% solvent A (50% methanol) and 99% solvent B (100% methanol) at a flow rate of 0.1 mL/min. For the enrichment of HexCers containing GalCer(d18:0;h16:0) from a larger amount of mouse caeca and lipid extraction from small amounts of mouse tissue as well as lipid extraction from bacteria, see the supplemental Methods.

Fourier transformation-ion cyclotron resonance (FT-ICR)-MS analysis of αGalCerMLI
Fourier transformation-ion cyclotron resonance (FT-ICR)-MS analyses were performed on a solariX 7T XR magnetic resonance mass spectrometer (Bruker Daltonics) equipped with a dual ESI/ MALDI ion source. Spectra were acquired in ESI positive ion mode with a size of 4M data points and active absorption mode; transfer time-of-flight was 0.700 ms, and the Q1 mass was set on the mass of interest to achieve highest sensitivity. Calibration was performed on clusters of sodium formate followed by internal lock mass calibration for MS² on the precursor ion of interest. For each spectrum in MS mode, 20 scans with an accumulation time of 0.01 s were added, and in MS² mode, 200–800 scans with an accumulation time of 1.1 s were added. CID fragmentation was performed with a collision energy of 25 V after isolating the respective precursor with a width of ±0.1 Da.

Activation of iNKT cells with lipids
To generate dendritic cells (DCs), bone marrow precursors from wild-type mice were cultured in complete IMDM medium supplemented with 10% FCS and 1% of a supernatant from a granulocyte-macrophage colony-stimulating factor-expressing cell line for 14 days. To investigate iNKT cell reactivity, 1 × 10⁵ DCs (>90% purity) were cultured with 1 × 10⁵ mouse iNKT hybridoma DN32.D3 (a gift from A. Bendelac, Chicago, IL) in the presence of purified glycolipids or vehicle in complete RPMI medium supplemented with 5% FCS for 24 h. Glycolipids were added from a stock solution of 0.1 mg/ml in PBS containing 0.05% Tween 20. Coculture supernatants were collected, and cytokine (interleukin-2 (IL-2)) production was measured by ELISA (R&D Systems, Minneapolis, MN). To assess activation of primary iNKT cells, splenocytes (10⁶ cells/well, 96-well plate) were exposed to purified glycolipids or vehicle alone for 48 h. Supernatants were collected, and IFN-γ concentration was measured by ELISA (R&D Systems).

Treatment of mice with Western type diet
Male C57BL/6 mice (age 23–36 weeks) were housed in a controlled environment with free access to food and water. The mice were fed a Western type diet (WTD) (EF TDB8317, smnff special diets GmbH, Germany, containing 21.1% fat, 14.4% polysaccharides, 34.3% sugar, and 0.21% cholesterol) or a standard rodent diet (control; Altromin, Lage, Germany; #1310, containing 5.1% fat, 35% polysaccharides, and 5% sugar) for 7 days and euthanized afterwards by cervical dislocation. Tissue was removed quickly, weighed, and stored at −80°C until sample preparation.

Induction of colitis in mice
Nine- to 11-week-old C57BL/6 male mice were given 2% dextran sodium sulfate (DSS) (colitis grade, molecular weight 36,000–50,000; MP Biomedicals GmbH, Eschwege, Germany) in drinking water and euthanized, respectively, after 2 or 4 days of treatment. Induction of colitis was verified by performing an occult blood test (Hemoccult; Beckman Coulter GmbH, Krefeld, Germany) after 4 days of exposure to DSS.

Lung infection of mice with influenza A virus
Eight-week-old male C57BL/6 mice were purchased from Janvier (Le Genest-St-Isle, France). For infection with influenza A virus (IAV), mice were maintained in a biosafety level 2 facility in the Animal Resource Center at the Pasteur Institute, Lille. Mice
were anesthetized and administered intranasally with 50 µl of PBS containing [or not (mock)] 30 plaque-forming units of the high-pathogenicity murine-adapted H3N2 IAV strain, Scotland/20/74, as reported previously (32). Mock-treated and IAV-infected mice were euthanized at day 7 postinfection.

RESULTS

Screening of lipid extracts from mouse intestine by HILIC-MS² revealed the presence of a potential microbial αGalCer isomer

We previously reported a HILIC-MS² method to successfully separate GlcCers and GalCers as well as α- and β-HexCer isomers, showing the detection of α-glycosidically linked GalCers (αGalCers) from bacteria (8). Based on published structures for bacterial αGalCer, we screened mouse intestinal tissue for these and similar αGalCer structures by MRM in positive ESI mode. The mass spectrometric transitions used a sphingoid base-based fragment, indicating by that the size or chain length of the incorporated sphingoid base. In total cecum of C57BL/6N mice, we initially found a HexCer peak that migrated at the same retention time as αGalCer(d17:0;βh17:0), one of the main αGalCers from B. fragilis (αGalCerBf). The caecal HexCer had the same chemical formula, but appeared with the MRM transition for a C18-sphinganine (d18:0)-containing compound. Therefore, the attached hydroxylated fatty acid could only contain 16 carbons, HexCer(d18:0;h16:0). This peak, however, was absent in the cecum of germ-free NMRI mice (Fig. 1A–C). When separating cecum tissue from its luminal content, this peak disappeared from the washed caecum and increased in the content of caecum (Fig. 1D). Further investigations showed that this peak was basically absent in the duodenum and jejunum, but present in the content of cecum and colon as well as in feces (Fig. 1D). As it is present basically in the mouse large intestine (MLI), we name the structure here αGalCerMLI. When comparing corresponding tissue of germ-free NMRI mice with control NMRI mice, this compound was only found in control mice but not in germ-free mice (Fig. 1E). These data implicate a bacterial origin of the αGalCerMLI peak, which is supported by its enrichment in the corresponding tissue contents and depletion in intestinal tissue lacking content (Fig. 1D).

The HILIC-MS² results lead us to the hypothesis that, as attributed to αGalCerBf, this peak from the mouse large intestine would contain an α-glycosidically linked galactosyl residue and a β-hydroxylated acyl chain, the latter N-linked to a dihydrosphingosine base, but, in contrast to αGalCerBf, with an even number of C-atoms in both the LCB and the fatty acyl chain.

Fig. 1. HILIC-MS² reveals a potential microbial αGalCer(t34:0) obtained from murine large intestine samples. A–D: HILIC-MS². A: Extracted ion chromatogram (EIC): Detection of αGalCer(t34:0) from purified sphingolipids of B. fragilis (termed αGalCerBf) with SRM transition m/z 740.5 ([M + H]+) to m/z 270.3 ([Sa(d17:0) + H – H2O]+), i.e., containing d17:0-sphingoid base and h17:0-acyl chain. B, C: EIC: Detection of GalCer(d18:0/h16:0) with SRM transition m/z 740.5 ([M + H]+) to m/z 284.3 ([Sa(d18:0) + H – H2O]+) from total cecum (including luminal content) lipid extracts of mice with commensals (control) (B) and mice not harboring commensals (germ-free) (C). This GalCer(d18:0/h16:0) with identical retention time to αGalCerBf is described further on as αGalCerMLI. D: Quantities of αGalCerMLI at retention time = 3.57 min in extracts of tissues from 14-week-old male C57BL/6N mice including luminal content (total), tissues without luminal content (washed), and corresponding released luminal contents (n = 3). E: RP18-LC-MS². Quantification of αGalCerMLI from 14- to 18-week-old male and female NMRI mice using specific transitions for αHexCer as outlined in the Materials and Methods. Extracts of tissues included luminal content, both from commensal containing (control) and germ-free mice. Note that αGalCerMLI is enriched in intestinal luminal contents, but disappears in tissues of germ-free mice (control, n = 6; germ free, n = 3). *P < 0.05; **P < 0.01; ***P < 0.001.
Analyzing sodium adducts of GalCers by LC-MS² displays structural information on anomeric linkage of the sugar moiety and β-hydroxylation of the acyl chain

We next turned to prove the anomeric linkage of the sugar moiety and the position of the hydroxyl group in the acyl chain. Due to the low concentration of αGalCerMLI in cecum tissue with no more than 80 ng per single mouse cecum, we excluded NMR studies and focused on LC-MS² analysis relying on characteristic CID-generated product ion patterns for structure elucidation: First, Brennan et al. (25) previously demonstrated differences in the fragmentation spectra between α- and βGalCer-containing nonhydroxylated acyl chains when selecting the sodiated ions [GalCer + Na]⁺ (fragment c in Fig. 2A) over [GalCer + Na – (hexose – H₂O)]⁺ (fragment b in Fig. 2A) decreased significantly in the α-anomer, which we also saw analyzing commercially available GalCer(d18:1/16:0) and GalCer(d18:0/16:0) standards (Fig. 2B–E). Second, Brown et al. (7) reported a specific product ion to appear from the deprotonated αGalCer ion (–ESI) containing a β-hydroxylated fatty acyl chain. This ion corresponds to a similar product ion reported in the CID-spectra of a deprotonated ceramide ion from mouse skin, which also contains a β-hydroxylated fatty acyl chain and likely is derived by McLafferty (McL) fragmentation (33). This type of β-cleavage apparently depends on β-hydroxylation of the acyl chain.

To evaluate the fragmentation behavior of GalCer with β-hydroxylated acyl chains, we synthesized and quantified GalCer standards with a β-hydroxylated acyl chain [α-, βGalCer(d18:1/βh16:0); α-, βGalCer(d18:0/βh16:0)] as described in Materials and Methods. Due to the incorporation of racemic β/(R/ S)-hydroxylated palmitic acid into the corresponding chiral galactosylsphingosines, each synthesis yielded a mixture of two diastereomeric compounds, which migrated as double bands on thin-layer chromatography as well as double peaks on reversed phase 18 C-atoms (RP18-LC) (supplemental Fig. S1). On RP18-LC, β-hydroxylated GalCers eluted earlier than α-hydroxylated GalCers, which again eluted in front of the corresponding nonhydroxylated compounds (supplemental Fig. S1). CID-product ion spectra derived from the first peak of the double peak did not differ from the corresponding second peak for all four compound mixtures (supplemental Fig. S2). Comparison of CID-product ion spectra of α- and βGalCers containing a C18-LCB and a C16-acyl chain revealed a specific McL fragment [m/z 526 for α/βGalCer(d18:1/βh16:0) and m/z 528 for α/βGalCer(d18:0/βh16:0)] to appear in the spectra of compounds with a β-hydroxylated acyl chain (Fig. 2F–I), but not in spectra of those compounds containing nonhydroxylated or α-hydroxylated acyl chains [m/z 526 and 528 not observed for α/βGalCer(d18:1/16:0) and α/βGalCer(d18:0/16:0), Fig. 2B–E; m/z 542 or 524 not observed for βGalCer(d18:1/α/(R/ S)18:0), Fig. 2J, K]. The McL fragment in GalCers with a β-hydroxylated acyl chain was accompanied by another peak due to the subsequent loss of the sugar moiety from the McL fragment (supplemental Fig. S2). Spectra of compounds with an α-hydroxylated acyl chain contained a characteristic low intense fragment e (m/z 512) not present in spectra of GalCer with nonhydroxylated or β-hydroxylated acyl chains (Fig. 2J, K). The other fragments (a, b, c, and d), which are due to water, sugar, or acyl chain loss, appeared in all compounds as well as m/z 203 representing the release of sodiated galactose (supplemental Fig. S2). To address differences in the product ion spectra arising from the anomeric linkage of the galactosyl residue, we compared the CID spectra of corresponding pairs of αGalCer and βGalCer standards with identical ceramide anchor. The fragmentation pattern detected was qualitatively the same for corresponding pairs of α- and β-glycosidically linked GalCers. Differences were observed in the relative abundance of these fragments. As reported previously (25), the α- and β-anomers of GalCer with nonhydroxy acyl chains differed in the ratio of fragment b over fragment c in favor of b in spectra of the αGalCers. In spectra of GalCers containing β-hydroxylated acyl chains, fragment c almost disappeared. However, we found a quantitative difference in the abundance of the McL fragment over fragment b in favor of the McL fragment in β-anomeric compounds (Fig. 2F–I). These results encouraged us to adopt the b/McL fragment ratio as a marker for the anomeric structure of GalCers with β-hydroxylated acyl chains. Therefore, we determined the optimal collision energies for the corresponding MS² transitions (supplemental Fig. S3). With optimized parameters, all standards were then subjected to RP18-LC-MS² and monitored by MRM transitions of the molecular sodium adducts to fragments a, b, c, d, e, and McL. Standards with β-hydroxylated acyl chains again revealed double peaks, as they represented diastereomeric mixtures as stated above (supplemental Fig. S4). Relative quantification of the transitions revealed a highly significant (more than 15-fold) difference in the b/McL ratio between corresponding α- and β-anomeric compounds, independent of the R-/orientation of the β-hydroxy group (supplemental Fig. S5).

αGalCerMLI enriched from mouse cecum is indistinguishable from the synthetic αGalCer(d18:0;βh16:0) by RP18-LC-MS²

Next, we enriched GalCers from a pool of mouse caeca to increase signal intensities for further analysis, as described in the Materials and Methods. By RP18-LC-MS², the enriched sample contained two peaks of HexCer(d18:0:h16:0). The first and main peak (αGalCerMLI) eluted identically to the second peak of the synthetic α- and βGalCer(d18:0;βh16:0) standards but 0.15 min later than αGalCer(d17:0;βh17:0) from B. fragilis (Fig. 3A). Monitoring the above identified fragments in relative quantity by MRM, the αGalCerMLI peak was again not distinguishable from αGalCer(d18:0;βh16:0), but the b/McL ratio clearly differed from the corresponding β-anomer (Fig. 3B). Additionally, the fragmentation patterns of all three αGalCers detected in B. fragilis (αGalCerαβ) were identical to the major peak from the enriched cecum sample, except for differences in the chain length of the sphingoid base and acyl chain. The fragmentation pattern of the second minor HexCer(d18:0:h16:0)
Fig. 2. Product ion spectra of sodium adducts of synthetic GalCers. A: Structure of GalCers with nonhydroxylated and α-hydroxylated acyl chains (upper structure), and with β-hydroxylated acyl chains (lower structure) including fragmentation pattern to fragments a, b, c, McL, d, and e. B–K: Production ion spectra induced by CID (45 V) of αGalCer(d18:1/16:0) (B), βGalCer(d18:1/16:0) (C), αGalCer(d18:0/16:0) (D), βGalCer(d18:0/16:0) (E), αGalCer(d18:1/βh16:0) (F), αGalCer(d18:0/βh16:0) (G), βGalCer(d18:1/βh16:0) (H), βGalCer(d18:0/βh16:0) (I), βGalCer(d18:1/α(R)h18:0) (J), and βGalCer(d18:1/α(S)h18:0) (K). Note the difference in the b/c-ratio between α- and βGalCers with nonhydroxylated acyl chains (B–E), the unique McL fragment of GalCers with β-hydroxylated acyl chains and the difference in the b/McL-ratio between corresponding α- and βGalCers (F–I) as well as the unique fragment e from GalCers with α-hydroxylated acyl chains (J, K).
only GalCer can be separated from GlcCer with HILIC-MS2. 

Fig. 3. Comparison of GalCer(d18:0/h16:0) from mouse cecum with synthetic GalCer standards and αGalCer from B. fragilis by MRM on RP18-LC. A: Total ion chromatograms resulting from MRM transition of the sodiated compounds to fragments a, b, c, McL, d, and e (see Fig. 2A, supplemental Table S5). The RP18-LC gradient contained 10 mM of ammonium acetate as additive. B: Relative abundance of the monitored transitions for synthetic GalCer with a C18-base and a nonhydroxylated or β-hydroxylated C18:0-acyl chain, GalCer with a C18-base, and an α-hydroxylated C18:0 acyl chain, αGalCerMLI enriched from mouse cecum as well as αGalCerBf with a C17- (17), C18- (18), or C19-base (19) from B. fragilis (Bf), each containing a βR- or βS-hydroxylated C17:0-acyl chain. Synthetic GalCers with a β-hydroxylated acyl chain represent each a diastereomeric mixture containing βR- and βS-hydroxylated C16:0 acyl chains. These diastereomeric mixtures elute as double peaks (see A), which are annotated in B with (1.) and (2.) for the first and the second peak. Note that an αGalCerBf from mouse cecum [main peak (A)] migrates with the second peak of synthetic αGalCer(d18:0/βh16:0) at 3.48 min. Furthermore, this αGalCerMLI peak does not differ by fragmentation pattern from synthetic αGalCer(d18:0/βh16:0) nor from any αGalCer of B. fragilis. However, it significantly differs from that of βGalCer(d18:0/βh16:0). The αGalCerMLI from cecum eluted 0.15 min later than αGalCer(d17:0/βh17:0) from B. fragilis, likely due to the iso-branching of the sphingoid base and acyl chain in the latter. The minor HexCer(d18:0/h16:0) peak of the cecum sample (here at 3.76 min) migrates with βGlcCer(d18:0/ah16:0) generated by hydration from mouse small intestine GlcCer(d18:1/ah16:0) (supplemental Fig. S6) (n = 3).

peak (retention time = 3.76 min) from the enriched cecum sample did not reveal a McL fragment, but revealed at a low intensity the fragment e, which had been observed in the standards containing α-hydroxylated acyl chains (Fig. 3B). These results confirm that the major HexCer(d18:0/h16:0) of mouse caeca, i.e., αGalCerMLI, is a αGalCer(d18:0/βh16:0) containing a β-hydroxylated fatty acyl and an α-glycosidic sugar linkage.

HILIC-MS2 adds further evidence for the presence of αGalCer(d18:0/βh16:0) in mouse caeca

In our previous publication (8), we could show that not only GalCer can be separated from GlcCer with HILIC-MS2, but also αGalCer from βGalCer. The method had been set up with HexCer standards containing nonhydroxylated acyl chains. Therefore, we compared the retention behavior of a β-hydroxylated acyl chain containing an α/βGalCer standards with αGalCerMLI from mouse cecum and with αGalCerBf by HILIC-MS2. As by RP18-LC, every GalCer standard with a β-hydroxylated acyl chain revealed a double peak by HILIC-MS2 due to the racemic presence of either the βR- or the βS-hydroxylated acyl chain. However, in contrast to GalCers with nonhydroxylated acyl chains, the double peaks of α- and βGalCer with β-hydroxylated acyl chains eluted very similarly, with a minimal but significant difference of 0.03 min. Therefore, an equal mixture of the α- and βGalCer standards, both containing a β-hydroxylated acyl chain, revealed no baseline separation, but yielded two broad peaks, each with a shoulder (Fig. 4A–C). When αGalCerMLI from cecum was spiked with synthetic βGalCer(d18:0/βh16:0), the second of the double peaks broadened roughly by 50% (FWHH: first, 0.036 ± 0.003 min; second, 0.053 ± 0.009 min) and revealed a shoulder (Fig. 4F). Clearly αGalCerMLI differed in structure from the corresponding βGalCer. Next, we spiked synthetic αGalCer(d18:0/βh16:0) into the enriched αGalCerMLI from cecum. This time, the second of the standard double peaks remained a sharp peak (FWHH: first, 0.035 ± 0.002 min; second, 0.034 ± 0.001 min) without a shoulder, but, as expected, increased in intensity compared with the first peak (Fig. 4G, supplemental Fig. S7). These data further confirm the α-anomeric linkage of galactose in αGalCerMLI from mouse cecum. In addition, αGalCerBf(d17:0/βh17:0) migrated identically to this αGalCerMLI (Fig. 4D, E), which implies a βR-configuration for the hydroxyl group in the acyl chain.

Because α- and βGalCers with a β-hydroxylated acyl chain do not separate as much as corresponding GalCers with nonhydroxylated acyl chains, we also synthesized αGlcCers with a β-hydroxylated acyl chain. αGlcCer(d18:0/βh16:0) gives rise to a b/ McL ratio of 1.78 ± 0.08 in MS2, more related to αGalCer(d18:0/βh16:0) (b/McL = 2.93)
which are annotated as acid, the latter being used as a racemic compound for synthesis. As (D), h16:0) (F), GalCer(d18:0/h16:0) (C), synthetic from mouse cecum. Mixture of synthetic GalCer from B. fragilis in comparison with GalCer(d18:0/h16:0) in mouse cecum. Mixture of synthetic GalCer(d18:0/h16:0) (b/McL = 0.19). On the RP18-LC-MS2 system, both GalCer(d18:0/h16:0) peaks, which are eluting almost identically, elute in front of the second αGalCer(d18:0/h16:0). Because the latter corresponds to the compound found in the large intestine, this is an αGalCer and not an αGlcCer (supplemental Fig. S8).

High-resolution FT-ICR-MS2 confirms the presence of HexCer(d18:0/βh16:0) in mouse cecum and the structural identity of CID-fragments

We subjected the αGalCer_{MLI} enriched sample from mouse caeca to high-resolution FT-ICR-MS2 and detected a peak at m/z 740.56468 differing by 0.167 ppm from the theoretical mass of the chemical formula of GalCer(d18:0/h16:0). CID of this peak revealed a product ion at m/z 528.35097 corresponding with a mass error of 0.534 ppm to the theoretical McL fragment and a product ion at m/z 486.34035 corresponding with an error of 0.473 ppm to fragment d, as annotated in the GalCer structure with a β-hydroxylated acyl chain (Fig. 2A), as well as a product ion at m/z 348.28759 corresponding to the McL ion with further loss of hexose (C6H12O6) and an error of 0.833 ppm (supplemental Table S5). The latter is also observed in the product ion spectra of GalCer(d18:0/βh16:0) recorded by LC-ESI-(triple quadrupole)MS2 (supplemental Fig. S2). The presence of the McL fragment confirmed the β-hydroxylated acyl chain and, again, the b/McL ratio of 3.58 resembled the ratio of 2.87 found in synthetic αGalCer and the ratio of 2.82 found in αGalCer_Bf but clearly distinguished from the b/McL ratio of 0.13 found in synthetic βGalCer on the FT-ICR instrument.

Synthetic αGalCer_{MLI} activates iNKT cells

Presented on CD1d by APCs, αGalCer binds the TCR of iNKT cells and induces the formation of an immunological synapse. This formation leads to iNKT cell activation and cytokine secretion. To investigate the effects of synthetic αGalCer, a DC/iNKT cell coculture system was used (34). When tested in vitro, all synthetic αGalCers were capable of inducing, in a dose-dependent fashion, IL-2 production by the NKT cell hybridoma; the activation of which depending solely on the TCR (Fig. 5A). In contrast, all corresponding βGalCer structures failed to induce iNKT cell activation. The IL-2 production induced by the synthetic analog of αGalCer_{MLI}, i.e., αGalCer(d18:0/βh16:0), was comparable to that of αGalCer_{Bf}. The difference (lower IL-2 production) relative to the positive control, αGalCer(d18:0/βh16:0) (KRN7000), which contains the very long chain cerotic acid (26:0) [αGalCer(t18:0;26:0)], is in line with a previous report (7). Similar to KRN7000, αGalCer(d18:1/24:1) with the very long chain nervonic acid (24:1) activated iNKT cells at lower concentrations than the other αGalCers, which comprised shorter C16-fatty acids. We then assessed the effect of αGalCer structures on primary iNKT cells. To this end, splenocytes from wild-type and CD1d−/− mice were exposed to purified glycolipids, and IFN-γ concentration was measured by ELISA. As depicted in Fig. 5B, synthetic αGalCers activated, in a CD1d-dependent manner, the release of IFN-γ by splenocytes. Collectively, synthetic αGalCer_{MLI}s activate iNKT cells in a CD1d-dependent manner.
Stressful conditions alter the content of αGalCerMLI in the large intestine

Alterations of the intestinal microbiota composition are associated with a variety of diseases, including obesity, diabetes, and inflammatory bowel disease (35). Furthermore, diet is a major driving factor for the establishment of the gut microbiome, and a fat-rich diet is associated with the suppression of several bacteria, including a number of Bacteroides (36). Likewise, studies in mice revealed that obesity is associated with a division-wide increase in the relative abundance of the Firmicutes and a corresponding division-wide decrease in the relative abundance of the Bacteroidetes (37), which may produce αGalCerMLI. To study the impact of diet on αGalCerMLI levels in the gut, C57BL/6 mice were fed for 7 days with WTD, which is rich in fat and sugar. After 7 days, total cecum was analyzed for the presence of αGalCerMLI. Relative to the control, mice fed WTD had an almost 3-fold decrease of αGalCerMLI in the intestine (Fig. 6A).

We next turned to investigate the effect of a local inflammation (colitis) on the concentration of αGalCerMLI. To this end, mice were treated with the heparin-like polysaccharide, DSS, a compound commonly used to induce inflammatory disorders of the colon such as inflammatory bowel diseases, principally, ulcerative colitis and Crohn’s disease. This experimental model induces colitis in mice. Of note, Bacteroides were reported to significantly decrease in the colon after DSS treatment (38–41). Mice (C57BL/6) were treated with DSS for 4 days and samples were collected after 2 and 4 days. αGalCerMLI levels decreased progressively over time down to less than 3% on day 4 (Fig. 6B). Because the intestinal epithelial barrier can turn leaky during colitis, antigens and bacteria may get carried to the liver, which in mice is rich in NKT cells. Therefore, we also analyzed liver for the presence of αGalCerMLI in the colitis model. Levels were below the detection limit in all samples (Fig. 6B, supplemental Fig. S9).
Finally, we evaluated the influence of influenza infection on αGalCerMIL. Recently, gut dysbiosis has been reported upon experimental (H1N1 and H5N1) IAV infection with a peak at day 7 postinfection ([42–46] and unpublished observations, F. Trottein). Interestingly, the levels of caecal and fecal αGalCerMIL were reduced by more than 2-fold at day 7 post-influenza infection (Fig. 6C).

DISCUSSION

NKT cells are innate lymphocytes at the interface between innate and adaptive immunity, which impact initiation and regulation of immune responses (47). This may impact, for example, anti-tumor immunity in the liver, which appeared to be controlled by liver NKT cells via the gut microbiome (48, 49). This process appears to depend on Hexb-subunit (54). This may impact, for example, anti-tumor immunity in the liver, which appeared to be controlled by liver NKT cells via the gut microbiome (48, 49).

Via the cell surface receptor, CD1d, APCs present lipids to the TCR of invariant (or type I) NKT (αNKT) cells (47). Whereas a series of lipids that bind to CD1d have been published, only a few allow the formation of a stable CD1d-lipid-TCR trimeric complex, which leads to activation of αNKT cells, either in the thymus or in the periphery in the context of inflammation and infection. Lipids of endogenous or microbial origin are processed and loaded on CD1d inside APCs and are thereafter presented on the cell surface (47).

Pathogen-derived α-anomeric glycolipids include α-galacturonolyceramide [αGalACer(t18:0;14:0)] from Sphingomonas sp., 6-acetyl-α-glucosylcholesterol from Helicobacter pylori, α-galactosyldiacylglycerol [αGalDAG(18:1/16:0)] from Borrelia burgdorferi, and α-glucosyldiacylglycerol [αGlcDAG(18:1/16:0)] from Streptococcus pneumoniae (50–53). Most of these pathogen-derived lipids do not activate αNKT cells as strongly as the synthetic αGalCer agonist, KRN7000 (2), which contains a very long acyl chain and α-phosphorylosinosine. It was based on the naturally strong agonist, Agelasphin 9b [αGalCer(t18:0/α(R)h24:0)] (6).

Endogenous self-lipids have been proposed to replace pathogenic lipids on CD1d to activate αNKT cells in cases of infection. This process appears to depend on Hexb-subunit (54). Hexb is required for the formation of hexosaminidases (HexA and HexB) involved in glycosphingolipid degradation/processing in lysosomes. In this lysosomal compartment also CD1d loading takes place (55).

Recently host-derived α-glycosylceramides (56, 57) and simple gangliosides GM3 and GD3 (34) containing mono-unsaturated nervonic acid have been discussed to serve as endogenous lipid ligands. The latter were enriched after Toll-like receptor stimulation of APCs (34, 58). The former are αGalCers containing sphingosine and nonhydroxylated acyl chains, apparently produced by immune cells (56, 57). Similar αGalCer structures, but with sphinganine and saturated nonhydroxylated acyl chains, were also reported to be present in cow milk and likely occur in the thymus, supporting the potential of mammalian cells to produce endogenous αGalCers (25, 59). In our hands, αGalCer containing nervonic acid, especially, appeared to stimulate αNKT cells more efficiently than αGalCers with shorter (C16-C17) acyl chains (Fig. 5).

Interestingly, not only the positive selection of αNKT cells in the thymus, but also the neonatal maturation of αNKT cells in the gut epithelium is CD1d restricted (24, 50, 60). For the latter, gut microbiota-derived αGalCer or related compounds might be critical (24, 60). In this context, αGalCers synthesized by the commensal B. fragilis exert either an inhibitory effect preventing proliferation or a stimulatory effect favoring the development of αNKT cells. In studies with neonatal mice lacking access to bacterial sphingolipids, treatment with B. fragilis glycosphingolipids (GSL-B717/αGalCerα) reduced colonic αNKT cell numbers and conferred protection against oxazolone-induced colitis (9); however, these αGalCers from B. fragilis bind to CD1d and activate both mouse and human αNKT cells, in vitro and in vivo (7).

In the current study, we screened mouse intestinal tissue for αGalCers with our physico-chemical HILIC-MS² method. We did not detect significant amounts of αGalCer displaying the same structures as those of B. fragilis. However, we detected a signal for a very closely related structure and showed that it is indeed an αGalCer containing a β-hydroxylated acyl chain [αGalCerα, i.e., αGalCer(d18:0/βh16:0)]. This αGalCerα was, on the contrary, basically absent in corresponding lipid extracts from B. fragilis (supplemental Fig. S6I–L). Identification was achieved in comparison to corresponding synthetic α- and β-anomeric GalCers and related compounds. Although characteristically different retention times for α- and β-anomeric compounds diminished in GalCers containing a β-hydroxylated acyl chain compared to those with a nonhydroxylated acyl chain, the intensity ratios of specific fragments obtained from the sodiated molecular ions could be used to clearly annotate the anomeric linkage of the glycosidic bond. In contrast to GalCers with nonhydroxylated acyl chains, we did not use the ratio of fragments b and c (see Fig. 2), which had been reported lately (25), but the ratio of fragment b to the fragment specific for the presence of β-hydroxylated fatty acids, the McL fragment. Interestingly, this fragment is nicely observed from the sodiated but not the protonated molecular ions. As the McL fragment appears much more intense than fragment c, the sensitivity to screen biological samples for the presence of αGalCers with a β-hydroxylated acyl chain increases manifold using corresponding MRM transitions, while adding structural information. Both features, α-glycosidic linkage and β-hydroxylation of the acyl chain, are given by the αGalCer structures found in B. fragilis. Interestingly, αGalCer(d17:0;βh17:0) from B. fragilis (αGalCerα) migrated with HILIC identically to αGalCerMLII as well as the second of the double peaks of synthetic αGalCer(d18:0/βh16:0) (Fig. 4). This αGalCerα has previously been shown to contain a β-hydroxylated acyl chain (7). Because the hydrophobic part of the ceramide anchor rather does not interact with the stationary phase in HILIC, it is therefore likely that the second peak of the synthetic standard corresponds to the β-hydroxylated isomer. This implies that αGalCerMLII, like αGalCerβ, contains a β-hydroxylated

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acetyl chain. Differences arise in the length of the sphingoid base and N-linked acyl chain, but the total amount of C-atoms in the ceramide anchor of αGalCerMLI was identical to the main αGalCer (GSL-B717) of B. fragilis. Therefore, it is not surprising that the synthetic αGalCerMLI activates iNKT cells in a coculture system with similar results as αGalCer. αGalCerMLI was present in cecum (an extremely rich source of commensals) but not in duodenum and jejunum, arguing for a microbial instead of a host-derived origin. Moreover, germ-free mice did not reveal this compound in their intestinal tract samples, here too arguing for a microbial origin of the αGalCerMLI. The structure of the iNKT-stimulatory αGalCerMLI described in the current study might well be the enterogenous bacterial αGalCer-like glycolipid reported by Wei et al. (23). In the later study, the authors used the monoclonal antibody, L363, described to recognize a CD1d-complex with αGalCer or with a structurally related lipid.

As we found the identical structure in C57BL/6 and NMRI mice (Fig. 1) and in mice from four different facilities [three academic (Fig. 6) and one commercial (supplemental Fig. S10)], we exclude a strain-specific or facility-specific event. Nevertheless, absolute concentrations may depend on strain (supplemental Fig. S10). Because germ-free mice, lacking αGalCerMLI, obtained the same food as control mice, we exclude food as a source of this compound. Therefore, we conclude that the common microbiota of the mouse gut produce the αGalCerMLI identified here. Considering that microbial αGalCer components were found in Bacteroides and Prevotella species of the human gut microbiome (7, 8), we hypothesized that the identified αGalCerMLI may be part of the Bacteroidetes taxa.

Previous publications showed the correlation between specific diets and the composition of the microbiome, especially the change in the ratio of Firmicutes versus Bacteroidetes [see reviews (61, 62)]. As this αGalCerMLI corresponds in functional activity to αGalCer, it may be speculated that, together with corresponding symbionts, it contributes to the microbiome-dependent neonatal maturation of iNKT cells in the gut epithelium, as has been reported lately (24, 50, 60). Both, αGalCer and αGalCerMLI are not such strong inducers of iNKT cell activation as is the synthetic agonist, KR7000, with its very long acyl chain, as demonstrated for αGalCer in supplemental Fig. 3 of Ref. 7. This may be attributed to the fact that the microbial compounds integrate a shorter acyl chain (C16 but not C26) and do not present a 4-hydroxy group in their sphingoid base (63–65). The incorporation of a shorter (C16-C17) acyl chain into αGalCers of commensals (going along with lower iNKT activation) might have developed evolutionarily to prevent inflammation against commensals. The β-hydroxylation of the N-linked acyl chain, however, appears to have little influence, if at all. The IL-2 and IFN-γ responses were similar upon stimulation with nonhydroxylated or β-hydroxylated acyl chains containing αGalCer (Fig. 5). To date, there is no established enzymatic pathway integrating β-hydroxylated acyl chains into mammalian sphingolipids, and it may turn out that β-hydroxylation could preferentially hint to a microbial origin of corresponding sphingolipids, although a ceramide with a β-hydroxylated acyl chain, i.e., Cer(d18:1;βh26:0), had been described for mouse epidermis (66, 67), another organ colonized with commensals. In contrast to the odd and iso-branched sphingoid base found in αGalCerbg, the even and presumably unbranched sphingoid base of αGalCerMLI is also found in mammalian tissue, which therefore cannot be excluded as a corresponding source for the base. Likewise, uptake and potential processing of αGalCerMLI in mammalian lysosomes eventually could lead to de-acetylation, e.g., by acidic ceramidase, leading to “mammalian-like” αGalCer. To reveal such processes, lysosomal degradation of exogenous αGalCer should be investigated in more detail.

Assuming that the αGalCerMLI was of microbial origin, we thought its intestinal levels should depend on the composition of the microbiota and should reflect changes in the microbiome composition described in the literature.

Because diet is a major driving factor for the establishment of the gut microbiome, we evaluated the influence of WTD and observed an almost 3-fold decrease of αGalCerMLI with this diet after only 1 week. Other studies in mice have previously revealed that obesity is associated with an increased relative abundance of Firmicutes and a decreased relative abundance of Bacteroidetes (37, 68). Bacteria described to produce αGalCer with β-hydroxylated acyl chains indeed belong to the latter phylum (7, 8). Low levels of Bacteroides were also reported in obesity/metabolic disorder (36, 69) in line with a relative depletion of Bacteroides in neonates exposed to maternal high-fat gestational diet (70).

Bacteroidetes were also reported to decrease in a commonly used model of experimental colitis induced by DSS (41). Decreased Bacteroides levels are associated with inflammatory bowel disease and ulcerative colitis (36, 71) and a decreased abundance of Erysipelotrichales, Bacteroidales, and Clostridiales going along with an increase of Enterobacteriaceae, Pasteurellaceae, Veillonellaceae, and Fusobacteriaceae were observed in new-onset cases of pediatric Crohn’s disease (72). The loss of Bacteroidetes would correlate the strong decrease of αGalCerMLI which we observed in a model of DSS-induced colitis. The over 30-fold decrease indicates special sensitivity of αGalCer-producing bacteria to inflammatory disorders of the colon.

Infections primarily not targeting the intestinal tract nevertheless impact the gut microbiome. IAV infection, for example, leads to enteric dysbiosis in the mouse system, and possibly in humans [(42–46) and unpublished observations, F. Trottein]. Our data show that mice infected 7 days earlier with IAV (the peak of lung inflammation) displayed a reduced amount of αGalCerMLI relative to noninfected animals. Several changes in the gut microbiome have been observed after infection with IAV. Especially, the Bacteroidetes S24-7 family to which the potential αGalCer-producing Bacteroides belong decreased on day 7 postinfection [(42) and unpublished observations, F. Trottein].

Collectively, our data reveal for the first time the presence of an αGalCer [αGalCerMLI, which is αGalCer(d18:0;βh16:0)] in the digestive tract of mice, which is very likely of microbial
origin, and demonstrate that stressful conditions altering the microbiome composition impact strongly on its production. These findings raise interesting questions concerning the potential impact of such αGalCer alteration on functions and activation status of iNKT cells and a corresponding impact on global immunity.

Note added in proof
There was an error in Supplemental Table 3 in the version of this article that was published as a Paper in Press on September 4, 2019. In column 2, the precursor ion labels [M+H]+ and [M+H3O]+ were inadvertently switched. This error has now been corrected in the online version.

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