C18:3-GM1a induces apoptosis in Neuro2a cells: enzymatic remodeling of fatty acyl chains of glycosphingolipids

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Abstract GM1a [Galβ1-3GalNAcβ1-4(NeuAcα2-3)Galβ1-4Glcβ1-1Cer] is known to support and protect neuronal functions. However, we report that α-linolenic acid-containing GM1a (C18:3-GM1a), which was prepared using the reverse hydrolysis reaction of sphingolipid ceramide N-deacylase, induced apoptosis in neuronal cells. Intraneuronal DNA fragmentation, chromatin condensation, and caspase activation, all typical features of apoptosis, were observed when mouse neuroblastoma Neuro2a cells were cultured with C18:3-GM1a but not GM1a containing stearic acid (C18:0) or oleic acid (C18:1). The phenotype of Neuro2a cells induced by C18:3-GM1a was similar to that evoked by lyso-GM1a. However, lyso-GM1a caused a complete disruption of lipid microdomains of Neuro2a cells and hemolysis of sheep erythrocytes, whereas C18:3-GM1a did neither. C18:3-GM1a, but not lyso-GM1a, was found to be abundant in lipid microdomains after the removal of loosely bound GM1a by BSA. The activation of stress-activated protein kinase/c-Jun N-terminal kinase in Neuro2a cells was observed with lyso-GM1a but not C18:3-GM1a. These results indicate that the mechanism of apoptosis induced by C18:3-GM1a is distinct from that caused by lyso-GM1a. This study also clearly shows that fatty acid composition of gangliosides significantly affected their pharmacological activities when added to the cell cultures and suggests why naturally occurring gangliosides do not possess polyunsaturated fatty acids as a major constituent.

Supplementary key words ganglioside • polyunsaturated fatty acids • neuroblastoma • lipid microdomain

Glycosphingolipids (GSLs), found in the outer leaflet of the plasma membranes of vertebrates, are thought to play functional roles in various cellular events, such as cell proliferation and differentiation (1–6) and cellular interaction and recognition, including microbial infection (7–9).

There is much evidence that sphingolipids dynamically cluster with cholesterol to form lipid microdomains, so-called detergent-insoluble membranes (DIMs), detergent-insoluble GSL-enriched domains, GSL-enriched membranes, or rafts (10–12). Lipid microdomains have specific proteins that are anchored to membranes by saturated acyl chains (e.g., Src family protein tyrosine kinases and glycosylphosphatidylinositol-anchored proteins) (13). Most sphingolipids have saturated fatty acyl chains, whereas glycosphingolipids are rich in unsaturated fatty acyl chains. This allows sphingolipids to pack tightly with cholesterol to form domains in the glycosphingolipid bilayer and provides sphingolipids with their high melting temperature (14). However, the question of why cells do not synthesize GSLs with polyunsaturated fatty acids as a major constituent has not been answered.

Exogenously added GSLs, especially gangliosides, sialic acid-containing GSLs, show various biological activities in neuronal cells. For example, neurite formation, a typical phenotype for neuronal differentiation, can be induced by gangliosides, notably by GM1a [Galβ1-3GalNAcβ1-4(NeuAcα2-3)Galβ1-4Glcβ1-1Cer] (5) and GQ1b (1). Antiapoptotic effects of GM1a have also been observed (15). For the past decade, clinical applications of GM1a for neurological disorders such as Alzheimer’s disease (16), Parkinson’s disease (17), and spinal cord injury (18) have been reported. However, these studies and clinical applications did not focus on the role of the ceramide (Cer) moiety of gangliosides in the activity. On the other hand, GSLs are known to show heterogeneity not only in their carbohydrate moiety but also in their lipid moiety. For ex-
ample, gangliosides from bovine brain contain more than seven species of fatty acids in the Cer moiety, although stearic acid (C18:0) is a major constituent (19). Recently, it was suggested that the structure of Cer, especially the composition of fatty acids, could influence the localization and functions of GSLs on plasma membranes, possibly by direct interaction with membrane proteins, cholesterol, and phospholipids (20, 21). Thus, the function of exogenously added GSLs could be changed with the conversion of their fatty acid moiety. The present study indicates that remodeling of fatty acyl chains of GSLs is easily performed by sphingolipid ceramide N-deacylase (SCDase), an enzyme capable of catalyzing reversible reactions by which the N-acyl linkage of Cer in GSLs is hydrolyzed or synthesized (22, 23).

Lysosphingolipids, which lack fatty acyl chains of sphingolipids, are known to be accumulated in some cases of sphingolipid storage diseases (24–27). We have reported that the accumulation of lysosphingolipids, but not parental sphingolipids, are known to be accumulated in some cases of sphingolipid storage diseases (24–27). We have reported that the accumulation of lysosphingolipids, but not parental sphingolipids, triggers the apoptotic cascade in neurononal cells of patients with sphingolipidosis (28). We found in this study that GM1a containing α-linolenic acid (C18:3-GM1a) also induces apoptosis in Neuro2a cells. The fact that lyso- and PUFA-containing gangliosides showed strong cell toxicity may partly explain why cells do not synthesize these aberrant gangliosides as a major constituent. The physiological relevance of lyso-GSLs has been studied by many researchers (28–30), whereas that of PUFA-containing gangliosides has not yet been clarified, although the latter gangliosides have actually been detected as a minor constituent of various tissues (31–33). We report here that exogenously added C18:3-GM1a was abundant in the lipid microdomains, whereas lyso-GM1a disrupted them, both leading to apoptosis but in a different manner.

MATERIALS AND METHODS

**Materials**

GM1a was prepared from bovine brain gangliosides using the sialidase-producing marine bacterium *Pseudomonas* sp. YF-2 as described previously (34). Serric acid (C18:0), oleic acid (C18:1-n-9), linoleic acid (C18:2-n-6), α-linolenic acid (C18:3-n-3), eicosa- and docosa-hexaenoic acid (C20:5-n-3), and docosa-hexaenoic acid (C22:6-n-3) were obtained from Sigma (St. Louis, MO). SCDase was prepared from the culture supernatant of *Pseudomonas* sp. TK4 as described (22, 35) or purchased from Takara Bio (Otsu, Japan). Lyso-GM1a was prepared from GM1a using SCDase (36) or the SCDase-producing *Pseudomonas* sp. TK4 as a microbial catalyst, as reported previously (37). N-2 supplement was purchased from Invitrogen (Carlsbad, CA). [3H]thymidine and ECL Plus reagents were purchased from Amersham Biosciences (Little Chalfont, UK). [14C]serric acid and [14C]linolenic acid were purchased from American Radiolabeled Chemicals (St. Louis, MO). Ac-DEVD-MCA and zVAD-fmk were obtained from Peptide Institute, Inc. (Minoh, Japan). Specific polyclonal antibodies directed at extracellular signal-regulated kinase (ERK) 1/2, p44/42 mitogen-activated protein kinase (MAPK), phospho-ERK, and phospho-SAPK/JNK (stress-activated protein kinase/ c-Jun N-terminal kinase) were obtained from Cell Signaling Technology (Beverly, MA), and monoclonal antibody (clone N-CAM 13) directed toward neural cell adhesion molecule (NCAM) was from BD Biosciences PharMingen (San Diego, CA). OptiPrep was purchased from Nycomed Pharma (Oslo, Norway).

**Synthesis and purification of the reconstructed GM1a with a single fatty acid molecule**

Ten micromoles of lyso-GM1a and 25 μmol of fatty acid were incubated at 37°C for 18 h with 100 mU of SCDase in 40 ml of 25 mM phosphate buffer, pH 7.0, containing 0.1% Triton X-100. The reaction mixture was applied to a Sep-Pack Plus C18 cartridge to remove salts that had been previously equilibrated with distilled water. After a wash with 20 ml of distilled water, the GM1a was eluted from the cartridge with 1 ml of methanol and 10 ml of chloroform-methanol (2:1, v/v), and purified by normal-phase HPLC using a silica column (AQUASIL SS-1251, 4.6 × 250 mm, Senshu Pak; Senshu Scientific Co., Tokyo, Japan) with a linear gradient of chloroform-methanol-distilled water from 65:25:4 (v/v) to 5:4:1 (v/v). In a typical experiment, 5 μmol of C18:3-GM1a was obtained when 25 μmol of α-linolenic acid and 10 μmol of lyso-GM1a were used as the substrates.

**Electrospray ionization-liquid chromatography-mass spectrometry**

Fifty nanomoles of the reconstructed GM1a was analyzed by electrospray ionization-liquid chromatography-mass spectrometry (ESI-LC-MS) with an API 365 (Applied Biosystems, Foster City, CA). The GM1a was introduced into the equipment with 50% methanol containing 5 mM ammonium acetate as a solvent at a flow rate of 5 μl/min. The mass spectrometer was operated in the positive-ion mode (the ion-spray voltage was set at 5,000 V, and the orifice voltage was 60 V).

**Cell culture**

Neuro2a cells were cultured in DMEM supplemented with 60 mg/l kanamycin and 10% FBS at 37°C in an atmosphere of 95% air and 5% CO₂.

**DNA fragmentation, chromatin condensation, and nuclear fragmentation analyses**

DNA fragmentation was analyzed by agarose gel electrophoresis. In brief, fragmented genomic DNA was extracted by incubating cells in a 10 mM Tris-HCl buffer, pH 7.5, containing 10 mM EDTA and 0.5% Triton X-100 at 4°C for 10 min. Under these conditions, intact genomic DNA was not extracted. The cell suspension was centrifuged at 15,000 rpm for 20 min, and the resulting supernatants were incubated at 37°C for 1 h in the presence of RNase A (400 μg/ml) and then for an additional 1 h in the presence of proteinase K (400 μg/ml). The fragmented DNA was precipitated with isopropanol and then analyzed by 2% agarose gel electrophoresis. For analyses of chromatin condensation and nuclear fragmentation, the cells were washed with PBS and fixed in a 1% glutaraldehyde solution at 4°C for 75 min. After fixation, cells were collected by centrifugation at 800 × g for 5 min, washed with PBS two times, and resuspended in PBS. The cell nuclei were stained with Hoechst 33258 (Wako, Osaka, Japan) and examined with a fluorescence microscope with an ultraviolet combination filter. The DNA fragmentation was also determined by flow cytometry (EPICS XL; Beckman Coulter, Fullerton, CA).

**Cholera toxin staining and fluorescence microscopy**

Neuro2a cells, cultured on a cover glass, were treated with 40 μM GM1a for given periods in serum-free DMEM containing 20
µg/ml BSA and the N-2 supplement. To remove the GM1a loosely attached to cell surfaces, the cells were washed with DMEM containing 0.2% fatty acid-free BSA. After BSA treatment, cells were fixed with 3% paraformaldehyde in PBS for 15 min. After being rinsed with PBS and 0.1 M glycine in PBS, cells were permeabilized with 0.1% digitonin in PBS. After treatment with blocking buffer (5% skim milk in PBS) for 15 min, the samples were incubated with biotin-labeled cholera toxin B-subunit (CTB; diluted 1:1,000 with blocking buffer) at 4°C overnight followed by Cy3-labeled streptavidin at room temperature for 2 h. Samples were examined with a confocal laser-scanning microscope (Digital Eclipse C1; Nikon, Tokyo, Japan).

**SDS-PAGE and Western blotting**

SDS-PAGE was carried out according to the method of Laemmli (38). Protein transfer onto a polyvinyl difluoride membrane was performed using a Trans-Blot SD Semi-Dry Cell (Bio-Rad, Hercules, CA). After treatment with 3% skim milk in TBS containing 0.1% Tween 20 (T-TBS) for 1 h, the membrane was incubated with primary antibody at 4°C overnight. After three washes with T-TBS, the membrane was incubated with horseradish peroxidase-conjugated secondary antibody at room temperature for 2 h. After another three washes with T-TBS, the ECL reaction was performed for 5 min as recommended by the manufacturer, and chemiluminescent signals were visualized on an ECL™ Mini-camera (Amersham Biosciences).

**TLC blotting**

After sucrose density gradient centrifugation, fractions were dialyzed with distilled water, dried with a SpeedVac concentrator, and applied to a plastic TLC plate (Polygram SIL G; Macherey-Nagel, Duren, Germany), which was developed with chloroform/methanol/0.02% CaCl₂ (5:4:1, v/v). After treatment with 1% methanol/0.02% CaCl₂ for 5 min and washed with PBS and 0.1 M glycine in PBS, cells were permeabilized with 0.1% digitonin in PBS. After treatment with blocking buffer (5% skim milk in PBS) for 15 min, the samples were incubated with biotin-labeled cholera toxin B-subunit (CTB; diluted 1:1,000 with blocking buffer) at 4°C overnight followed by Cy3-labeled streptavidin at room temperature for 2 h. Samples were examined with a confocal laser-scanning microscope (Digital Eclipse C1; Nikon, Tokyo, Japan).

**Sucrose and OptiPrep density gradient analyses**

Sucrose gradient analysis was performed by the method described previously (39). Briefly, Neuro2a cells cultured in a 35 mm dish (1 × 10⁶) were suspended in 250 µl of ice-cold homogenizing buffer (10 mM Tris-HCl, pH 7.5, containing 150 mM NaCl, 5 mM EDTA, 5 µg/ml leupeptin, 5 µg/ml pepstatin, and 5 µg/ml chymostatin) containing 1% Triton X-100, allowed to stand for 20 min on ice, and homogenized using a Teflon glass homogenizer. The cell lysate was centrifuged for 5 min at 1,500 g and 4°C to remove nuclei and large cellular debris. The supernatant fraction was mixed with an equal volume of the homogenizing buffer containing 85% sucrose (w/v). A discontinuous sucrose concentration gradient, 1.1 ml of 30% and 0.6 ml of 5% sucrose in the same buffer, was layered over the lysate. Samples were centrifuged at 200,000 g for 18–20 h at 4°C. After centrifugation, 220 µl fractions were collected from the top of the gradient (10 fractions) and subjected to trichloroacetic acid precipitation. OptiPrep gradient analysis was performed as described previously with some modifications (40). The cell lysate was centrifuged at 1,300 g for 5 min at 4°C to remove nuclei and large cellular debris. The supernatant fraction was mixed with an equal volume of 50% (v/v) OptiPrep in the homogenizing buffer. A discontinuous OptiPrep concentration gradient, 1.1 ml of 20% and 0.6 ml of 10% OptiPrep in the same buffer, was layered over the lysate. Samples were centrifuged at 200,000 g for 18–20 h at 4°C. After centrifugation, 220 µl fractions were collected from the top of the gradient (10 fractions) and subjected to trichloroacetic acid precipitation.

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Fig. 1. Enzymatic synthesis of GM1a [Galβ1-3GalNAcβ1-4(NeuAcα2-3)Galβ1-4Glcβ1-1Cer] having different fatty acyl chains by sphingolipid ceramide N-deacylase (SCDase). A: Four nanomoles of lyso-GM1a was incubated with 10 nmol of various fatty acids at 37°C for 18 h with 40 µU of SCDase in 20 µl of 25 mM phosphate buffer, pH 7.0, containing 0.1% Triton X-100. When eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) were used as a substrate, 120 µU of SCDase was used for the reaction. After incubation, reaction efficiencies were determined as described previously (23). Values are means for duplicate determinations. B: Electrospray ionization-liquid chromatography-mass spectrometry analyses were conducted with an API 365 mass spectrometer using the positive-ion mode. C: Fatty acid methyl esters of reconstructed GM1a were analyzed by gas chromatography (GC-14A; Shimadzu Co., Kyoto, Japan) with a flame-ionization detector and a capillary column (HR-SS-10, 0.25 mm × 30 m; Shinwa Chemical Industries, Kyoto, Japan). C18:0-GM1a, GM1a with stearic acid; C18:1-GM1a, GM1a with oleic acid; C18:2-GM1a, GM1a with linoleic acid; C18:3-GM1a, GM1a with α-linolenic acid; C20:5-GM1a, GM1a with eicosapentaenoic acid; C22:6-GM1a, GM1a with docosahexaenoic acid.
RESULTS

Condensation of various fatty acids to lyso-GM1a by SCDase and identification of the reconstructed GM1a

The extent of GM1a synthesis with unsaturated fatty acid was examined using the condensation reaction of SCDase. The efficiencies of all reactions reached ~80% when stearic acid (C18:0), oleic acid (C18:1), linoleic acid (C18:2), or α-linolenic acid (C18:3) was used as a substrate, whereas the value was 20–30% when eicosapentaenoic acid (C20:5) or docosahexaenoic acid (C22:6) was used (Fig. 1A). The saturated fatty acyl chains of C12-C18 were well condensed, but those of C2-C6 and C22-C26 were less condensed to lyso-GM1a by SCDase (23). These results suggest that the length of fatty acyl chains is more important in determining the efficiency of the condensation reaction by the SCDase than is the degree of unsaturation of the fatty acid, if the reaction period is long enough with an excess amount of enzyme. After the purification of the reconstructed GM1a, as described in Materials and Methods, the structure of GM1a was determined by ESI-LC-MS using the positive-ion mode. As shown in Fig. 1B, the characteristic pseudomolecular ions (M+H)+ and (M+Na)+ were found at values of the expected molecular masses corresponding to the reconstructed GM1a with a different fatty acid. Furthermore, the fatty acid molecule of each reconstructed GM1a was determined by gas chromatography after methanolysis of the sample. It was confirmed that each reconstructed GM1a contains the expected single fatty acid (Fig. 1C). These results clearly indicated that unsaturated fatty acids were successfully condensed by the SCDase to lyso-GM1a to generate a GM1a having each unsaturated fatty acid.

Growth inhibition of Neuro2a cells by the GM1a reconstructed with unsaturated fatty acids

The addition of GM1a composed of d18:1 sphingosine and a stearic acid (C18:0-GM1a) or an oleic acid (C18:1-GM1a), both of which are naturally occurring gangliosides, did not affect the incorporation of [3H]thymidine by Neuro2a cells (Fig. 2A). On the other hand, this index...
was markedly decreased in a dose-dependent manner by the addition of C18:2 and C18:3-GM1a, the Cer of which is composed of a d18:1 sphingosine and a linoleic acid (C18:2) and o-linolenic acid (C18:3), respectively. As shown previously (28), lyso-GM1a, which lacks the fatty acyl chain of GM1a, was very toxic to the cells (Fig. 2A).

Features of apoptosis induced by C18:3-GM1a

Sueyoshi, Maehara, and Ito (28) demonstrated that lyso-GM1a induced apoptosis in Neuro2a cells. Thus, the possibility that the suppression of DNA synthesis by C18:3-GM1a is derived from apoptosis was examined. After treatment of cells with C18:3-GM1a, the genomic DNA showed a ladder fragmentation pattern on agarose gel electrophoresis, which is one of the features typical of apoptosis (Fig. 2B). In addition, C18:3-GM1a-treated cells showed chromatin condensation and nuclear fragmentation after staining with Hoechst 33258 (Fig. 2C). These results indicated that not only lyso-GM1a but also C18:3-GM1a induced the apoptosis of Neuro2a cells. Because DNA fragmentation was observed not only in Neuro2a cells but also in human leukemia cell line HL60 cells (Fig. 2D), the C18:3-GM1a-induced apoptosis is not specific for neuronal cells. Apoptotic cells were also quantified with a flow cytometer. The ratio of apoptotic cells reached 30% of total cells after treatment with C18:3-GM1a for 24 h (Fig. 3E), whereas it was less than 10% with C18:0-GM1a (Fig. 3C). Interestingly, after the removal of loosely bound GM1a by BSA treatment, apoptosis was still observed at almost the same rate (Fig. 3D). These results indicated that treatment with C18:3-GM1a, but not C18:0-GM1, significantly induced apoptosis in Neuro2a cells.

Next, caspase activity was measured using the fluorescence substrate Ac-DEVD-MCA, which is specifically hydrolyzed by caspase-3. In Neuro2a cells, an increase of caspase-3-like activity was observed after the addition of C18:3-GM1a followed by incubation at 37°C for 18–24 h (Fig. 4A). Simultaneously, a DNA ladder was detected by agarose gel electrophoresis (Fig. 4B). The addition of an inhibitor for caspases, zVAD-fmk, to the Neuro2a cell culture inhibited the increase in protease activity (Fig. 4A) but not the DNA fragmentation (Fig. 4B). This result suggested that a caspase-3-independent signaling pathway is involved in the C18:3-GM1a-induced apoptosis.

Incorporation and distribution of exogenously added gangliosides in Neuro2a cells

To quantify the amount of gangliosides that enter the Neuro2a cells, cell-bound gangliosides were determined after removing the loosely bound gangliosides by treatment of the cells with fatty acid-free BSA. The amounts of C18:0-GM1a, C18:3-GM1a, and lyso-GM1a bound to cells decreased to half after BSA treatment, suggesting that half of the gangliosides/lyso-gangliosides were attached loosely to cells (Fig. 5A). The amount of C18:3-GM1a incorporated in the cells was almost the same as that of C18:0-GM1a and was approximately twice that of lyso-GM1a after BSA treatment. It is noted that the incorporation of gangliosides increased slowly during the course of incubation regardless of the difference in the fatty acyl chain (Fig. 5A). The distribution of exogenously added gangliosides in cells was compared with and without treatment with BSA by confocal laser-scanning microscopy after staining with biotin-labeled CTB (Fig. 5B). It is noteworthy that both C18:0- and C18:3-GM1a were detected mainly on the cell surface (arrows) and partly in intracellular compartments (arrowheads) during incubation, regardless of the treatment of cells with BSA. At the cell surface but not at the intracellular compartment, gangliosides were found to decrease in quantity on treatment of the cells with BSA, consistent with the result shown in Fig. 5A. This result confirmed that loosely bound, but not incorporated, gangliosides were removed by BSA treatment. Apoptosis could be caused by the C18:3-GM1a incorporated into plasma membranes, because apoptosis was induced after treatment of the cells with BSA (Fig. 3), although the possibility that C18:3-GM1a incorporated into intracellular compartments could cause the apoptosis cannot be ruled out at present.

Metabolism of exogenously added C18:0-GM1a and C18:3-GM1a in Neuro2a cells

To investigate how GM1a is metabolized in Neuro2a cells after its incorporation, 14C-labeled C18:0- and C18:3-
GM1α (labeled at fatty acyl chains) were added to the cell cultures. Both 14C-labeled GM1α forms were incorporated into Neuro2a cells and partially metabolized to GM2, GM3, and more complex gangliosides, mainly GD1α (Fig. 6). The metabolic patterns of the two 14C-labeled GM1α forms were almost the same, although the metabolic rate of C18:3-GM1α was slower than that of C18:0-GM1α. It is noted that neither GM1α was metabolized to Cer under the conditions used, which are known to induce apoptosis, indicating that apoptosis was not induced by Cer generated from the metabolism of C18:3-GM1α.

**Different effects of C18:3-GM1α and lyso-GM1α on erythrocytes and MAPKs**

The apparent features of the apoptosis induced by C18:3-GM1α seem to be almost the same as, if not identical to, those caused by lyso-GM1α (28). Therefore, it was investigated whether or not the mechanisms underlying the apoptosis induced by C18:3-GM1α and lyso-GM1α are the same. As shown in Fig. 7A, lyso-GM1α caused the hemolysis of sheep erythrocytes in a dose-dependent manner, whereas C18:0-GM1α and C18:3-GM1α did not. This result may indicate that the lyso-GM1α-induced apoptosis of Neuro2a cells is caused by a nonspecific detergent-like effect, perhaps by physical damage to the plasma membranes, and is different from the apoptosis induced by C18:3-GM1α.

Neuritogenesis of Neuro2a cells was induced by adding gangliosides in which the activation of MAPK was observed (41). Considering this report, the involvement of MAPK activation in C18:3-GM1α-induced apoptosis was examined. The treatment of Neuro2a cells with lyso-GM1α decreased the phosphorylation level of ERK1/2 and simultaneously increased that of SAPK/JNK (Fig. 7B). On the

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**Fig. 4.** Activation of caspase-3-like enzymes by C18:3-GM1α. A: Time course of caspase-3-like activity. Neuro2a cells were incubated in DMEM containing the N-2 supplement and 20 μg/ml BSA at 37°C with 40 μM C18:3-GM1α in the presence or absence of zVAD-fmk, a specific inhibitor of caspases, for the times indicated. Cytosolic extracts were prepared from cells, and the caspase-3-like activity in the lysates was measured using the fluorescent substrate Ac-DEVD-MCA. The fluorescence of the cleaved substrates was determined using a microplate reader (CS-9500PC; Shimadzu Co., Kyoto, Japan) set at an excitation wavelength of 360 nm and an emission wavelength of 460 nm. B: Time course for genomic DNA fragmentation. Genomic DNA was extracted from the cells and then analyzed by electrophoresis. Details are described in Materials and Methods.

**Fig. 5.** Incorporation of GM1α in Neuro2a cells. A: Cells (1 × 10⁵) were incubated with 40 μM C18:0-GM1α, C18:3-GM1α, and lyso-GM1α at 37°C for the times indicated. After treatment of the cells with 0.2% fatty acid-free BSA solution, lipids were extracted from cells and then analyzed by TLC using chloroform/methanol/0.02% CaCl₂ (5:4:1, v/v). For the control experiment, the BSA treatment was omitted. B: Cells were treated with 40 μM C18:0-GM1α and C18:3-GM1α for the indicated times. After incubation with gangliosides, cells were treated with (BSA+) or without (BSA−) 0.2% fatty acid-free BSA solution and then stained with biotin-labeled cholera toxin B-subunit (CTB). Signals were observed by confocal laser-scanning microscopy as described in Materials and Methods. Values are expressed as the mean with SD (n = 3).
other hand, C18:3-GM1a did not affect the phosphorylation of SAPK/JNK, although it decreased that of ERK1/2 similar to lyso-GM1a (Fig. 7B). C18:0-GM1a did not cause any change in the activity of MAPKs under the conditions used. Another MAPK, p38 MAPK, was not affected by the addition of either C18:3-GM1a or lyso-GM1a (data not shown).

**Distribution of exogenously added gangliosides in lipid microdomains**

In the sucrose density gradient analysis, endogenous GSLs, including GM2 and GM1a, were concentrated mainly in DIM fractions (Fig. 8A, fractions 2–4). The exogenously added C18:0-GM1a and C18:3-GM1a were detected mainly in bulk membrane-lipid fractions (Fig. 8B, fractions 6–8) and partly in DIM fractions (Fig. 8B, fractions 2–4) without treatment of the cells with BSA. However, after the removal of loosely bound GM1a by BSA, half of the GM1a was detected in DIM fractions, indicating that the incorporated GM1a, but not loosely bound GM1a, was concentrated in lipid microdomains similar to endogenous GM1a (Fig. 8C, fractions 2–4). Interestingly, the behavior of C18:3-GM1a in the sucrose density gradient was almost the same of that of C18:0-GM1a but completely different from that of lyso-GM1a, which was not detected in DIM fractions and was distributed only in bulk membrane-lipid fractions (Fig. 8A, fractions 6–8). In summary, C18:3-GM1a was incorporated and distributed in cell membranes possibly in the same manner as C18:0-GM1a and concentrated in DIM fractions after its incorporation, similar to endogenous GSLs, whereas lyso-GM1a was detected solely in bulk membrane lipid fractions.

**Effects of C18:3-GM1a and lyso-GM1a on lipid microdomains**

NCAM-120, a member of the immunoglobulin superfamily that is expected to be localized to lipid microdomains of neuronal cells (42), was analyzed by Western blotting using a specific monoclonal antibody after the treatment of cells with gangliosides. It was found that NCAM-120 was completely excluded from DIM fractions of lyso-GM1a-treated cells but was included in the fractions of C18:3-GM1a-treated cells when DIM was prepared by the sucrose density gradient method using Triton X-100 (Fig. 9A). However, when the membrane was fractionated using an OptiPrep density gradient without detergents, NCAM-120 disappeared from the low-density fractions and could only be detected in high-density fractions in both lyso-GM1a- and C18:3-GM1a-treated cells (Fig. 9B). C18:0-GM1a did not affect the distribution of NCAM-120 in sucrose or OptiPrep density gradients. These results suggested that the addition of lyso-GM1a to the Neuro2a culture almost completely disrupts the structure of the lipid microdomains (DIM fractions), whereas C18:3-GM1a does not disrupt but somehow affects the structure, judging from the behavior of a lipid microdomain-associated protein, NCAM-120, in sucrose and OptiPrep density gradients. In contrast to these aberrant gangliosides, naturally occurring C18:0-GM1a has no apparent effect on the structure of...
lipid microdomains when added to the cell culture at a concentration of 40 μM (Fig. 9A, B).

**DISCUSSION**

It is generally accepted that gangliosides contain saturated fatty acids but not PUFAs in the Cer moiety. Unexpectedly, we found in the literature that PUFAs are widely distributed in gangliosides as a minor constituent [e.g., mouse fibroblast 1 cells contain 1.3% and 7.2% of PUFAs in the fatty acid portion of monosialoganglioside and disialoganglioside, respectively (32); bovine serum contains 1.7% and 1.1% of C18:3 fatty acids in monosialoganglioside and disialoganglioside, respectively (32); human milk contains 0.39% and 0.17% of α-linolenic acid in GM3 and GD3, respectively (33); and GD3 of bovine kidney contains ~2% of C18:3 fatty acid (31)]. Nevertheless, little attention has been paid to the physiological significance of PUFA-containing gangliosides. We found that C18:3-GM1a at relatively high concentration (<40 μM) induced apoptosis in Neuro2a cells, whereas at lower concentration (5 μM) it induced neurite formation of PC12 cells in the absence of nerve growth factor (data not shown). Considering these results and the ubiquitous distribution of PUFA-containing gangliosides, the physiological and pharmacological functions of PUFA-containing gangliosides should be further clarified not only in vitro but also in vivo. In this context, a series of reports demonstrating that remodeling of the fatty acyl chain of GM1a enhanced the neuroprotective effects is particularly interesting (43, 44).

In the apoptosis induced by C18:3-GM1a, a caspase-3-independent signaling pathway could be involved, because the addition of an inhibitor for caspases, zVAD-fmk, to the Neuro2a cell culture inhibited the increase in the protease activity but not the DNA fragmentation. A pathway that is not inhibited by zVAD-fmk has also been observed in lyso-GM1a-induced apoptosis in Neuro2a cells (28) and...
adenovirus death factor-induced apoptosis (45). Recently, it was reported that endonuclease G released from mitochondria was involved in caspase-independent DNA degradation in the apoptosis of mouse embryonic fibroblasts, which lack the caspase-activated DNase (46). Thus, it is likely that endonuclease G-like enzymes along with caspase-like enzymes are involved in the C18:3-GM1a and lyso-GM1a-induced DNA fragmentation in Neuro2a cells. It should be noted that ZVAD-fmk does not affect the activity of endonuclease G.

The activation of the SAPK/JNK cascade seems to be essential for apoptosis induced by stress, including ionizing radiation and chemotherapeutic drugs (47, 48). On the other hand, the activation of ERK promotes cell proliferation. The balance of the activation between the SAPK/JNK and ERK pathways is thought to determine the fate of cells (i.e., death or survival) (49, 50). In this context, it is interesting that the inhibition of ERK (inhibition of the ERK pathway) was caused by both C18:3-GM1a and lyso-GM1a, whereas the activation of SAPK/JNK was attributable only to lyso-GM1a. SAPK seems to be activated through the nonspecific stress applied to the plasma membranes by lyso-GM1a but not C18:3-GM1a, although the mechanism underlying the apoptosis induced by both aberrant gangliosides remains to be clarified.

Recently, GSLs were found to be enriched with cholesterol to form lipid microdomains on plasma membranes (12). It is believed that the difference in the hydrophobic region of gangliosophospholipids and sphingolipids (i.e., the former contains mainly unsaturated fatty acids and the latter mainly saturated fatty acids) causes the phase separation to form lipid microdomains (21, 51). C18:3-GM1a, which contains polyunsaturated fatty acid, was expected to be excluded from lipid microdomains. Unexpectedly, however, C18:3-GM1a was found in DM fractions at almost the same level as C18:0-GM1a. This result can be explained by either the "hydrogen bonding network hypothesis," in which sphingolipids are considered to bind each other with hydrogen bonds between hydroxy and amido groups, or the "polar head group interaction hypothesis," in which GSLs are considered to aggregate via lateral carbohydrate-carbohydrate interactions (52). It is noteworthy that lyso-GM1a was not detected in DM fractions, possibly because the lipid destroyed the structure of lipid microdomains.

To address the significance of the fatty acid moiety of GSLs, a specific and convenient method is still required to reconstruct GSLs with unsaturated fatty acids. Until now, the only choice was a chemical procedure (53) that could generate by-products and is time-consuming. SC&Dase, an enzyme capable of hydrolyzing the Nacyl linkage between fatty acid and Cer in various GSLs and sphingomyelin (22), was found to catalyze the reverse reaction, in which a fatty acid is condensed to a lyso-GSL to yield a GSL (23). Cloning of SC&Dase from Shewanella alga G8 revealed that the reversible reactions were catalyzed by a single protein (54). SC&Dase is a superior reagent for remodeling GSLs with not only saturated fatty acids (37, 55, 56) but also unsaturated fatty acids (this study). Thus, the method using SC&Dase to remodel fatty acyl chains in gangliosides will facilitate further development in GSL research.

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