



CHEMISTRY AND FUNCTIONAL DISTRIBUTION OF SULFOGLYCOLIPIDS

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ABBREVIATIONS

A ₂ Gro	diacylglycerol (acyl ₂ Gro), e.g. A _{16:0} A _{18:3} Gro designates 1-palmitoyl-2-linolenoylglycerol	SM4s-Glc	GlcCer I ³ -sulfate, HSO ₃ -3Glcβ-1Cer, glucosyl sulfatide
Cer	ceramide, <i>N</i> -acylsphingosine (d18:1/24h:0 designates a ceramide with sphingene and 2-hydroxy 24:0 fatty acid)	SM3	lactosyl sulfatide, LacCer II ³ -sulfate, HSO ₃ -3Galβ-4Glcβ-1Cer.
d18:1	sphingene (sphingosine)	SM2a	Gg ₃ Cer II ³ -sulfate, GalNAcβ-4(HSO ₃ -3)Galβ-4Glcβ-1Cer
d18:0	sphingane (dihydrosphingosine)	SM2b	Gg ₃ Cer III ³ -sulfate, HSO ₃ -3GalNAcβ-4Galβ-4Glcβ-1Cer
E ₂ Gro	<i>sn</i> -2,3-di- <i>O</i> -alkylGro, (alkyl) ₂ Gro, when the carbon chain length is known written as follows: E ₂₅ E ₂₀ Gro, <i>sn</i> -2-sesterterpenyl-3-phytanylglycerol	SB1a	Gg ₄ Cer II ³ ,IV ³ - <i>bis</i> -sulfate, HSO ₃ -3Galβ-3GalNAcβ-4(HSO ₃ -3)Galβ-4Glcβ-1Cer
EAGro	1-alkyl-2-acylGro, e.g. E _{14:0} A _{16:0} Gro designates 1-myristyl-2-palmitoylglycerol	SB2	Gg ₃ Cer II ³ ,III ³ - <i>bis</i> -sulfate, HSO ₃ -3GalNAcβ-4(HSO ₃ -3)Galβ-4Glcβ-1Ce r
GalSph	galactosyl sphingosine (psychosine)	SMGb ₄	Gb ₄ Cer IV ³ -sulfate, HSO ₃ -3GalNAcβ-3Galα-4Galβ-4Glcβ-1Cer
h	e.g. in 24 h:0,2-hydroxy fatty acid	SMGb ₅	Gb ₅ Cer V ³ -sulfate, HSO ₃ -3Galβ-3GalNAcβ-3Galα-4Galβ-4Glc β-1Cer
HSO ₃ -Chol	cholesterol 3-sulfate	SMGM1a	HSO ₃ -3Galβ-3GalNAcβ-4(NeuGcα2-3)Galβ-4Glcβ-1Cer
PAPS	3'-phosphoadenosine 5'-phosphosulfate	SMiGb ₄	iGb ₄ Cer IV ³ -sulfate, HSO ₃ -3GalNAcβ-3Galα-3Galβ-4Glcβ-1Cer
SM4g	HSO ₃ -3Galβ-1-Diradylglycerols (subscript S, M, and g stands for sulfate, mono- and glycerol respectively. Diradyl means both alkylacyl and diacyl types are included), seminolipid	SMiGb ₅	iGb ₅ Cer V ³ -sulfate, HSO ₃ -3Galβ-3GalNAcβ-3Galα-3Galβ-4Glcβ-1Cer
SM4s	GalCer II ³ -sulfate, HSO ₃ -3Galβ-1Cer, galactosyl sulfatide (s stands for sphingoid)	SMUnLc ₄ Cer	SGGL-1 (U stands for uronic acid), HSO ₃ -3GlcUβ-3Galβ-4GlcNAcβ-3Galβ-4Galβ-4Glcβ-1Cer
SM4s-6	GalCer I ⁶ -sulfate, HSO ₃ -6Galβ-1Cer	SMUnLc ₆ Cer	(SGGL-2), HSO ₃ -3GlcUβ-3-[Galβ-4GlcNAcβ-3] ₂ Galβ-4Glcβ-1Cer
		HSO ₃ -PtdGro	phosphatidylglycerosulfate
		SQ-A ₂ Gro	6-sulfo-α-D-quinovosyldiacylglycerol
		t18:0	trihydroxysphinganine (phytosphingosine)

The abbreviations and symbols for carbohydrates and lipids follow the recommendations of the IUPAC/IUBMB Commission of Biochemical Nomenclature.²¹³ In the present article the trivial name 'sulfatides', that has been used to designate Galβ-1Cer I³-sulfate, is used as a generic term for a class of amphiphiles containing sulfate esters and carbo-

hydrates,¹⁵⁹ in the same way as 'gangliosides' include all glycolipids containing sialic acids. For other trivial names for sulfatides see Section I. B.

I. INTRODUCTION

The purpose of this chapter is to summarize the recent progress on the purification, analysis, distribution and biological function of sulfatides with reference to the sulfoamphiphiles including cholesterol 3-sulfate (HSO₃-Chol), and sulfonoglycolipids. In particular, the quantitation data have been carefully assessed and structure determination processes evaluated as critically as possible. An attempt was made to cover the literature published later than 1975 and the subjects handled in the previous reviews in this series^{159,271} were described only briefly. To stay within space limitations, sometimes only reviews or the more recent works of the author were cited.

A. A Brief History

The first one of the mammalian sulfated glycolipids, galactosyl sulfatide (SM4s), was isolated by Thudichum in as early as 1884^{108,159,361,639} from the human brain. However, SM4s had long been neglected as dramatically underscored by the fact that the components were identified half a century later and the correct location of the sulfate ester was not determined until 1962.^{108,636}

In the next 20 years, numerous novel sulfatides were brought to light,³⁶¹ uncovering a distinct group of acidic amphiphiles comparable to gangliosides. The variety of sulfatides discovered at that time included 2,3,6,6'-tetraacyl-trehalose-2'-sulfate from *Mycobacteria*,¹⁴⁹ HSO₃-3Galpβ-6Manpα-2Glcα-1E₂₀E₂₀Gro from an extremely halophilic archaea,^{273,275} lactosyl sulfatide (SM3) from human kidney,³⁶⁶ seminolipid (SM4g) from mammalian testis,^{232,238} ganglioside sulfates from echinoderms,^{297,457} and the ganglio- and globo-series sulfatides from mammalian kidneys.^{400,561} Most of these compounds were discovered by metabolic labeling with [³⁵S]sulfate.^{159,238} The structure determination has greatly accelerated by the remarkable advances in MS as well as ¹H- and ¹³C-NMR. Figure 1 shows some representative sulfatides and a sulfonolipid with their major lipophilic species.

Since the 1970s, metabolism of SM4s in the myelin of the central nervous system of normal and metachromatic leukodystrophy (MLD) patients was extensively studied.¹⁰⁸ It had been assumed, however, that sulfatides might be functioning as one of the membrane matrices. The breakthrough was brought about by the recognition that SM4s was enriched in glandular epithelial tissues of mammals,^{267,361} and increased in the organs related to sodium excretion following NaCl loading to animals.²⁷⁰ This increase occurred at an individual cellular level⁴²⁶ by the upregulation of GalCer sulfotransferase.²³¹ Further, the metabolic activities of animals and the concentration of renal sulfatide sulfate could be stoichiometrically correlated by using the allometric equation.³⁹⁸ To elucidate the diversity of anionic glycolipids in the biosphere, two major principles of genetics and biology, the neutral theory of molecular evolution,²³⁸ and the allometric law³⁹⁸ respectively, were introduced. In fact, the anionic groups of amphiphiles including phosphates, phosphonates, sialic acids, and sulfates, were interchangeable in many tissues and cells.^{212,238} Recently, the discovery of several examples of horizontal descent of genes related to anionic glycolipids,⁵⁰¹ the upregulation of GlcCer in the myelin of galactosyltransferase-deficient mice⁷⁵ as well as the studies on sulfonoglycolipid-deficient strains of photosynthetic microorganisms¹⁵⁵ supported the above hypotheses. Between 1944 and 1956 the role of adenosine 3'-phosphate 5'-sulfophosphate (PAPS) was elucidated by Lipmann's group and others, although the purification of the GalCer sulfotransferase to homogeneity²¹¹ and cloning of the cDNA had to wait until 1997.²¹⁰ The cDNAs of cerebroside sulfatase and saposins were already cloned in the early 1990s.

The specific interaction of sulfatides with various mammalian proteins has stimulated studies on the possible ligand and receptor activities of sulfated amphiphiles.¹⁰⁷

Sulfatides interacted specifically with cell adhesive proteins including thrombospondins, and laminins,⁴⁶⁸ with vascular system proteins including selectins,⁶¹³ as well as with pathogenic viri and mycoplasmas.^{314,470} Sulfatides are increased in some human cancers originated from glandular epithelia including pulmonary cells, hepatocytes and renal tubular cells. Aberrant expression of some minor sulfatides in cancer tissues was tested for capability as tumor markers.^{136,205} The discovery of these biological characteristics of sulfatides increased publications on sulfatides sharply in the 1990s.

B. Classification and Nomenclature

Glycolipids can be classified into neutral and acidic groups.²¹³ Acidic glycolipids contain one or more anionic residues, including acidic carbohydrates (e.g. uronic acid, and sialic acid), esters of phosphate or sulfate (R-OSO₃H where R is the neutral glycolipid), C-P (phosphonate), and C-S (sulfonate, R-SO₃H) compounds.^{159,537} In addition to the ionic sulfate monoesters (half-esters of sulfuric acid), nonionic diesters (R-OSO₂O-sugar) and cyclic sulfates⁶⁰³ have been chemically synthesized but never isolated from biological sources.¹⁵⁹ Although sulfatides with two sulfate groups have sometimes been designated as disulfate,⁵³⁷ 'bis-sulfate' as in Gg₃Cer II³,III³-bis-sulfate,⁵⁵⁸ and Man α -2Glc α -3E₂Gro II²,6-bis-sulfate³⁶⁹ may be more appropriate. Sulfoamphiphiles can also be classified into sulfoglycoglycerolipids,^{238,273} sulfoglycosphingolipids,^{160,361,505,554} and steroid sulfates¹⁰⁵ according to their aglycons. Because the terms sulfatide, sulfatides, sulphatide(s), and sulfoglycolipid(s) are synonyms but recognized as the different words in the databases including Medline, the papers listed will be doubled by 'or-search' on the abstracts including all of these words in addition to the keyword 'sulfatides'. The Complex Carbohydrate Structure Database (CCSD) version 17 (1997), containing more than 48,000 records on glycoconjugate structures including sulfoglycoconjugates, is accompanied by the 'CarbBank' software operated in the Windows 95 system. The database and software can be downloaded through the WorldWide Web, www.ncbi.nlm.nih.gov/. LipidBank is a database which contains records on sulfoglycolipids [information available from Dr M. Oshima (E-mail: oshima@imcj.go.jp)] and LIPIDAT database contains informations concerning synthetic and biologically derived polar lipid phase behavior.³¹⁰

Some examples of trivial names and symbols for sulfatides currently under use include S-GalCer (sulfo-galactosylceramide);^{555,657} CSE (cerebroside sulfuric ester);¹⁵⁹ CBS (cerebroside sulfate),⁵³⁴ SGC (sulfogalactosylceramide), sulfoGalCer,³⁹⁵ and SUL²⁷ for SM4s,²³³ seminolipid,²³² SGD (sulfogalactosyldiglyceride), SGG (sulfogalactosyl glycerolipid),³⁴³ SGaAAG,^{135,238} and SulfoGalAAG³⁹⁵ for SM4g,²³³ S-LacCer for SM3;^{88,555} sulfoglucuronyl glycolipids (SGGL) for SMGlcUnLc₄Cer and SMGlcUnLc₆Cer; S_{tri1}, and S_{tri2} for SM2a and SB2.²⁴⁶

II. ISOLATION AND PURIFICATION

Isolation and purification of individual sulfatides are prerequisite for structural determination and quantitative analysis. It is desirable that the tissue concentration values are routinely corrected for recoveries.

A. Extraction

The media most frequently used for the extraction of total lipids containing sulfatides are chloroform/methanol/water mixtures.³²⁰ Folch-Suzuki procedure is the method of choice for relatively polar amphiphiles including gangliosides and sulfatides.^{50,116,160,480,505,547} In order to extract acidic glycolipids quantitatively, univalent cations must be present. These are usually supplied in adequate amounts by the tissue itself, but for the second and third extraction, an addition of Na⁺ or K⁺ is rec-

ommended,⁵³² while an excess of salt in the extraction medium will result in the precipitation of SM4s by the effect of salting out.¹⁸⁰

1. Folch–Suzuki Method

The standard extraction procedure for mammalian sulfoglycosphingolipids is as follows. Typically, rat kidneys (50 g) were extracted in three steps with: (1) 19 vol of chloroform/methanol (2:1, v/v); (2) 10 vol of chloroform/methanol/0.88% KCl (60:120:9, v/v); and (3) 10 vol of 40 mM sodium acetate in chloroform/methanol/water (30:60:8).⁵⁶⁰ The combined extracts were concentrated to dryness and treated with 20 ml of 0.2 M NaOH in methanol at 37°C for 1 hr to degrade acyl ester lipids. After adjustment of pH to 7.0 with 0.2 M HCl in methanol, the reaction mixture was partitioned in the Folch system. The clear upper phase was concentrated to approx. 1/10 vol, dialyzed, and lyophilized. The lyophilized extract was combined with the lower phase, made up to 500 ml of chloroform/methanol/water (5:10:1, v/v) by an addition of solvents, and applied to a column of DEAE–Sephadex.

Highly polar lipids (polysialosyl gangliosides, the ganglio-series sulfatides,⁵⁶³ SMUnLc₄Cer and SMUnLc₆Cer),⁶⁸ inorganic salts and nonlipids move into the upper aqueous phase (chloroform/methanol/KCl, 3:48:47) and less polar lipids (HSO₃-Chol, SM4g, SM4s, Gb₄Cer and GM4) can be recovered in the lower organic phase (chloroform/methanol/KCl, 84:14:1) of the Folch partition system. When a salt is not included in the aqueous phase substantial portions of HSO₃-Chol⁴²² and SM4s²²⁸ will be lost in the upper aqueous phase or in the ‘fluff’ which forms at the interface between the water and chloroform.³⁸⁸ In the presence of 0.1% KCl, for instance, 85% of dehydroepiandrosterone sulfate was found in the upper phase, while 90% of HSO₃-Chol was recovered in the lower phase.³⁸⁸ About 5% of the weight of the lower phase was KCl when this salt was used to wash extracts of brain tissues.⁴⁸⁰

It has been reported that several washings with the simulated (theoretical) upper phase (chloroform/methanol/water, 3:48:47, or more simply methanol/water, 1:1) were necessary to remove inorganic sulfate from the extract containing SM4s. In our experience, 3 washings were sufficient to remove inorganic [³⁵S]sulfate by Folch’s partition^{228,555} using the upper phases containing 0.88% KCl.⁴⁸⁰ The final lower phase, after 3 washings, contained 97.3 ± 0.5% of SM4s before washing, while when no salts were included in the upper phase, only 10.9 ± 0.1% was recovered in the lower phase after 3 washings.^{228,563} Lyso-SM4s,⁵⁶⁹ lyso-SM4g⁶⁰⁶ and lyso-SQ-A₂Gro¹¹⁸ were distributed almost quantitatively in the aqueous phase.

2. Bligh–Dyer Method

A method for extraction of lipids using a smaller amount of solvents was developed for economical and practical reasons. The Bligh–Dyer extraction method has often been applied to bacterial suspensions that contain a large amount of water^{272,273,498} or sometimes used for large-scale extraction of animal tissues. Typically, the suspensions were extracted with chloroform/methanol/water, 1:2:0.8 (v/v), then each vol of chloroform and water was added. The mixture was cleared by centrifugation, to separate upper aqueous and lower organic phases in a volume ratio of 0.9:1 with protein precipitates in the middle.³³² This extraction method may be unsatisfactory for plant lipid⁶³ and a larger amount of solvent may result in better recoveries.⁴¹⁹ In our experience, only 50% of human kidney SM3 was procured by Bligh–Dyer extraction.

3. 2-Propanol/Hexane/Water System

Halogenated hydrocarbon solvents, including chloroform, are ecologically hazardous and have strong absorption in the range below 245 nm used for U.V. detection of lipids after HPLC separation.¹⁹⁴ A hexane/2-propanol mixture has been used for a large-scale

extraction of phospholipids and glycolipids from brain tissue.¹⁸⁰ After treatment with NaOH, nonpolar lipids were effectively removed into the upper organic layer leaving a glycolipid fraction (containing 96% of SM4s) practically free of phospholipids. The periodic acid oxidation step of this procedure could be omitted.³⁸⁶ Lipids were also extracted from extraneural tissues such as erythrocyte membranes,¹⁶⁰ liver,^{204, 378} and hybridoma cells¹⁹⁴ with 2-propanol/hexane/water mixture or ethylacetate/methanol/water.¹⁹⁴

B. Desalting and Salt Forms

1. Removal of Salts and Hydrophilic Contaminants

Dialysis through nitrocellulose membranes (e.g. SpectraporTM dialysis tube with the pore size of Mw 3000) against water is the simplest and, in most cases satisfactory, method to remove not only salts but also other low molecular hydrophilic compounds such as oligosaccharides, nucleotide sugars or oligopeptides^{32, 50} from a suspension of sulfatides in water. Prolonged dialysis (> 48 hr) is usually avoided to minimize the loss of the monomers of gangliosides and SM4s^{47, 160} present in amounts below the critical micellar concentrations (CMC, approx. 150 $\mu\text{g}/\text{ml}$ for gangliosides) through nitrocellulose tubings. Because the CMC of SM4s/SM4g and lyso-SM4s/lyso-SM4g is reported to be > 3 μM ^{30, 455} or < 10–40 μM and < 100 μM respectively¹¹⁵ and the size of micelles is about 10⁶ Da,¹⁷⁹ being larger than those of gangliosides,⁴⁰⁴ the loss by dialysis may occur more slowly.

Removal of water-soluble contaminants from HSO₃-Chol, SM4s, and SM4g²²⁸ is easily achieved by Folch partition with high recoveries,⁵⁶³ although this procedure is not applicable to more hydrophilic amphiphiles such as lyso-SM4s (HSO₃-3GalSph),¹⁰⁹ SM3, SM2a and SB1a.⁵⁶³ Partition chromatographies in chloroform/methanol/water, 120:60:9,⁶²⁵ or chloroform⁶³ using Sephadex G-25 (superfine),^{109, 332, 480, 479, 555} or Sephadex LH-20 column²¹⁹ have been used to remove [³⁵S]PAPS from SM4s³⁵⁶ and SM3⁵⁵⁵ with satisfactory recoveries. Sephadex G-25 separated some higher acidic lipids including SMUnLc₄Cer probably due to their negative charges.²²¹ For a smaller scale, reversed phase chromatography using Sep-Pak C₁₈,^{244, 594} BondElut^{262, 505, 536} or silica gel RP-18²⁴⁴ was also able to remove the bulk of salts or nucleotides from SM4s,^{51, 137} lyso-SM4s,⁵⁷⁰ SM2a,⁴⁰² permethylated uronic methyl esters of SMUnLc₄Cer,³³⁶ SMUnLc₄Cer,¹⁶³ SMUnLc₆Cer,³⁸¹ gangliosides,⁴⁹ and the synthetic neoglycolipids coupled with phosphatidylethanolamine.¹¹² The recovery was approx. 79% for SM2a.⁴⁰² Salts were successfully removed from an HSO₃-Chol fraction by a silicic acid column.³⁹

2. Salt-Forms

As the pK_a of the sulfate half-ester, e.g. in sucrose polysulfate, is from 0.43 to 1.19,⁵⁹⁰ the salts of SM4s are fully ionized in solution above pH 3.5 (M⁺ SO₃⁻-O-3GalCer) and are more strongly acidic than phosphomonoesters, carboxyl groups of monosialosyl gangliosides or phosphatidylserine⁴⁵⁸ (cf. Figs 2 and 3). A concentrated solution of free sulfatide is strongly acidic and leads to cleavage of the ester bond. To avoid desulfation, free sulfatides should be neutralized by NH₄OH or NaHCO₃ and stored in the form of salts. The potassium or lithium salts have been recommended for the sake of stability.¹⁵⁹ When stored in water at 4°C,³² even the salts of SM4s gradually lost sulfate over several months. Dilute solutions of the K-salt of SM4s in the solvent mixture of Dabrowski (deuterated dimethylsulfoxide/deuterated water, 98:2) also released a few percent of sulfate ester at ambient temperature over several years.

The ammonium salts of sulfatides, eluted from a DEAE-Sephadex column with solvents containing ammonium acetate, were stable for years in a dessicator.⁴⁷⁸ Exceptionally, the ammonium salt of tetraacylated trehalose 2-sulfates obtained by elution with chloroform-methanol containing NH₄OH or ammonium acetate was unstable under most conditions of storage.¹⁴⁹ The spontaneous desulfation in this case was



Fig. 2. DEAE-Sephadex column chromatography of acidic lipids from boar testis. Acidic lipids were eluted with a gradient of ammonium acetate in chloroform/methanol/water, 5:10:1. Fractions were separated on a HPTLC plate in chloroform/methanol/acetone/acetic acid/water, 8:2:4:2:1, and visualized with orcinol/sulfuric acid. PI, phosphatidylinositol; PS, phosphatidylserine.

assumed to be autocatalytic owing to the sulfate anion produced,¹⁵⁹ and it was essentially inhibited by water. To avoid spontaneous solvolysis, Goren stored sulfatides as solutions in hexane, kept over a small amount of aqueous NaHCO_3 to suppress the hydrolysis and to continuously neutralize any traces of acid that were formed.

To convert the mixed cationic salt into K salt, less polar sulfolipids ($\text{HSO}_3\text{-Chol}$, SM4s, SM4g) are simply partitioned in a Folch system containing 0.88% KCl.²³² For more complete exchange, sulfolipids were once acidified by dissolution in chloroform/methanol/0.1–0.2 M HCl.^{275, 327, 602} The organic phase, containing the free acid form of sulfolipids, was immediately neutralized (to pH 8–9) by the addition of NH_4OH , pyridinium chloride,⁹⁴ NaOH, or KOH to final concentrations of 0.01–0.2 M to yield the desired salt form of the sulfolipid and was then freed from unbound cations by a desalting procedure. Alternatively, glycolipid sulfates were dissolved in chloroform/methanol/water, 1:1:0.05 and converted to their free acid forms by passing the solution through a small column of cation exchanger (H^+ form).³⁸³ The free acid in the combined eluates was converted to the ammonium salt by the addition of methanolic NH_4OH . The affinity of the sulfate group in SM4s for cations is similar to that of sialic acids in gangliosides⁴⁰⁴ and in the order of $\text{Ca}^{2+} > \text{Mg}^{2+} > \text{K}^+ > \text{Na}^+ > \text{Li}^+$.⁷

C. Chromatographic Procedures

Generally, column chromatographic separation of sulfatides is achieved by a combination of ion-exchangers and silica beads,^{228, 365, 480, 561} while for a relatively simple lipid mixture, such as the total lipid from halophilic bacteria, a silicic acid column only may be sufficient.^{332, 369} Elimination of gangliosides by treatment with sialidase may reduce the loading of lipids and improve resolution of SM2a,^{233, 559} and SMUnLc₄Cer.^{68, 221} Convenient synopses are available on the substrates, solvents for chromatographic separation and the retention volumes or R_f values of glyco-glycerolipids,¹¹⁶ and glycosphingolipids.⁵⁴⁷

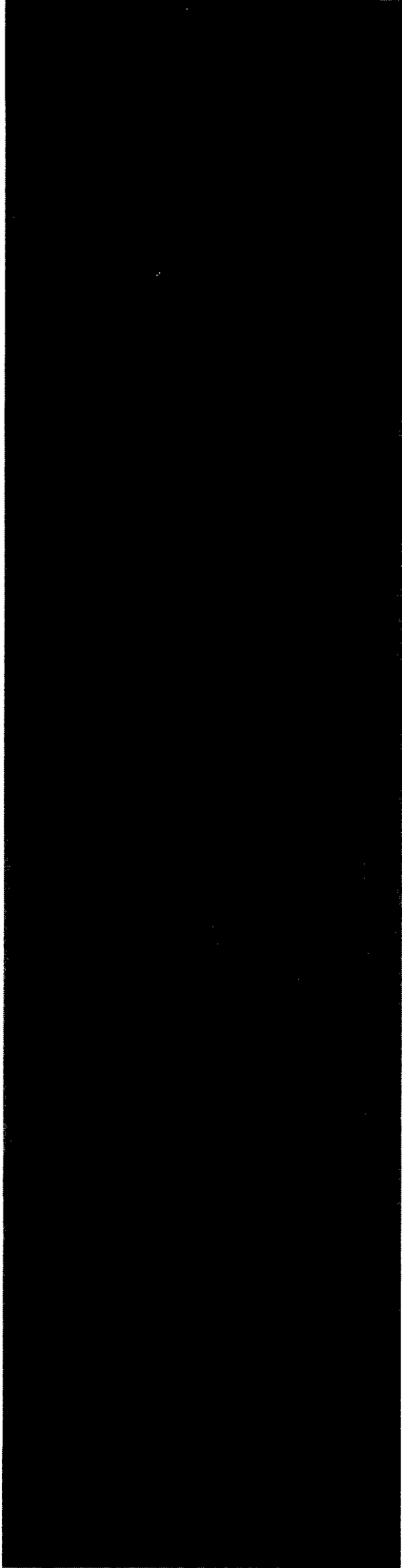


Fig. 3. Separation of acidic sphingoglycolipids from rat kidney by column chromatography using DEAE-Sephadex. Acidic lipids were eluted with a gradient of ammonium acetate in chloroform/methanol/water, 5:10:1. Each 1/2000 vol of the eluate of tube 1-120 and 1/200 vol of tube 121-288 were separated on HPTLC plates (5 x 10 cm, Merck) in chloroform/methanol/0.2% CaCl₂, 60:40:9 and stained with orcinol/sulfuric acid. BA, acidic glycolipid mixture from rat brain, containing a band under the double band of SM4s, is lyso-SM4g derived from SM4g by mild alkali treatment. The lowest four bands are gangliosides GM1, GD1a, GD1b, and GT1b.

1. Thin-layer Chromatography

Silica gel high-performance TLC (HPTLC) is indispensable for the studies of anionic amphiphiles³⁹¹ whereas cellulose TLC³³² is used less often for intact sulfatides.

(a) Adsorbents and solvent systems

Prewashing of the plate with methanol and activation of silica gel by heating at 100°C for 10 min or storing in a dessicator may eliminate soluble contaminants of silica gel and improve the resolution.⁶⁴⁴ With total or alkali-resistant lipid mixtures containing a substantial amount of cholesterol or detergents, double development of the plate, first with a nonpolar solvent system (e.g. chloroform/methanol/water, 90:12:1) to remove nonpolar contaminants to the solvent front^{400,494,555} and then with the second solvent system to separate the desired sulfatides, may improve the resolution greatly. Equilibration of tank vapor using a small motor-driven propeller accelerated saturation with the solvent system.²⁴⁴

In *neutral* solvent systems (Fig. 3) (e.g. chloroform/methanol/water, 65:25:4,^{130,232,342,479} SM4g can be separated from SM4s, although SM4s, lyso-SM4g, and SM2a migrate close to HSO₃-Chol,⁵⁸³ SM3, and ganglioside GM4 respectively. In chloroform/methanol/water (0.2% CaCl₂, 60:30:6,⁵⁶³ SM2a migrates a little slower than the neutral analogue, Gg₃Cer; while the *bis*-sulfo-derivative, SB2, migrates much slower than its monosulfo-counterpart SM2a.⁵⁶⁰ SM1a, the sulfate analogue of GM1a, is unique in migrating a little faster than Gg₄Cer.⁵⁶¹ Higher monosulfoglycolipids, SMiGb₄Cer and SMiGb₅Cer, comigrate with *bis*-sulfoglycolipids SB2, and SB1a respectively. SM4s, SM3, SM2a, and SB1a migrate much faster than the corresponding sialic acid derivatives, GM4, GM3, GM2 and GD1a. In chloroform/methanol/water solvent systems, lyso-SM4s migrates the distance corresponding to about 2/3 of SM4s,^{307,432} which is larger than those of GalSph (galactosylpsychosine) and GlcSph by the effect of the -NH₂ group. In contrast, lyso-SM4s comigrates with GalSph behaving like the sphingosylphosphocholine-HCl salt with the acidic and alkaline media.⁵⁶⁹ In *basic* solvent systems, e.g. chloroform/methanol/3.5 M NH₄OH, 60:40:9,⁵⁶⁸ SM4s comigrates with GalCer. Also HSO₃-Chol, and SM4s with nonhydroxy fatty acids are not separated from SM4s-Glc (118:0). In 1-propanol/15 N NH₄OH/H₂O, 80:5:15,^{323,425,433} bovine brain SM4s is separated into four bands on top of GalCer, because sulfatides migrate faster when ammonium ion concentration is higher.⁶³⁴

HSO₃-Chol,^{384,480} SM4g,^{119,228} SM4s with nonhydroxy fatty acids,^{17,51,234,314} SM4s-6,²³² SM4s-Glc with 118:0,²¹⁷ lyso-SM4g as well as mono- and *bis*-sulfo-Gg₃Cer⁵⁵⁸ can be neatly separated in *acidic* solvent systems (Fig. 2) including chloroform/methanol/acetone/acetic acid/water, 10:2:4:2:1. More polar acidic solvent systems may be suitable for more complex sulfo-glycolipids, including SM3 to SB1a,^{400,558} and mono- and *bis*-sulfated disaccharide E₂Gro.³⁶⁹ As in the neutral solvent systems, however, lyso-SM4g and SM4s with hydroxy fatty acids comigrate with SM3 and HSO₃-Chol respectively. Butanol/acetic acid/water, 6:2:2 has been successfully applied to SM4s and lyso-SM4s.⁵⁶⁹

Total lipid extracts from various sources have been separated by *two-dimensional*(2D) TLC^{116,237,480} using a couple of solvent systems selected from the basic, acidic or neutral solvent systems.^{237,466,547,574,606} Rat kidney acidic lipids were separated on a NH₂-silica gel HPTLC plate developed with chloroform/methanol/1% diethylamine, 50:47:15⁶²³ for the first-dimension. The bands were then transferred to an ordinary silica gel plate and developed to the second dimension in a neutral solvent system to separate mono- (SM4s, SM3), and *bis*-sulfoglycolipids (SB2, SB1a) from mono- (GM3, GM4) and disialosyl gangliosides (GD3) respectively. Alternatively, Analtech diphasic plate with an NH₂-modified silica gel lower hemisphere resulted in excellent separation of SM4g in sperm lipids.¹⁰

Extraction of sulfatides from silica gel is susceptible to low recovery (about 92% for HSO₃-Chol, 70% for SM4s²⁸² or SM4g, and much less for higher sulfatides). Usually the band of sulfatides is moistened with water before scraping to deactivate the adsor-

bent and individual glycolipid bands were eluted from the silica scrapings by using chloroform/methanol/water mixture (e.g. 10:10:1).^{383,412,539} To increase the recovery, the triglycosyl glycolipid sulfate was eluted from TLC silica gel with chloroform/methanol/0.1 N HCl, 1:2:0.8.²⁷⁵ The overall recoveries of SM4g from chicken retina,⁹⁵ rat brain,²²⁸ and human testis⁶⁰⁶ through purification procedures using Folch extraction, silicic acid or DEAE-cellulose columns, and preparative TLC were between 80 and 90% as assessed from the recoveries of [³⁵S]-labeled sulfatides or lipid galactose. Without an appropriate cleanup procedure, silicic acid and binder may contaminate in the extract from silica gel resulting in mass spectra of high noise levels.

Glycolipids and phospholipids can be efficiently transferred from an HPTLC plate to a polyvinylidene difluoride (PVDF) membrane by a simple blotting procedure.⁵⁷⁴ The developed HPTLC plate is dipped in a solvent mixture (2-propanol/0.2% CaCl₂/methanol, 40/20/7 v/v) for blotting, after which first a PVDF membrane, then a glass microfiber filter are placed on the plate and the assemblage pressed for 30 s by heating with a 180°C iron. The efficiency of transfer for each 5 μg of glycolipids was 68–93% (average 82%), and approx. 77% for SM4s.⁵⁷⁴ The glycolipids on the membrane can be detected chemically, or immunologically, and reextracted or analyzed by LSIMS.⁵⁷³

(b) Visualization

(i) *Chemical methods.* The cationic dye Azure A can stain sulfo-^{217,536} and sulfonconjugates³²¹ with satisfactory specificity. TLC plates are either dipped in or sprayed with a saturated solution of Azure A in 1 mM H₂SO₄ (2 g in 100 ml).^{217,504} The plates are washed by soaking in a mixture of methanol/40 mM H₂SO₄, 1:3, occasionally with gentle agitation and a few times of the change of the solution until the background staining is minimal. An excess of methanol in the washing solution may lead to the loss of nonpolar sulfatides such as SM4g, and an excess of water may destroy the layer of adsorbents. Aluminum-backed plates^{112,125,221,295,399} (e.g. Merck) are better than glass or plastic plates (e.g. the products of Marchery-Nagel, Brinkman²⁷ and Baker's²⁰⁴) and have the best strength with concomitant reduction in the resolution.⁵⁴⁹ As little as 250 pmol of SM4s is apparent to the naked eye and can be assayed by densitometry.^{369,387} The color may be retained for years at ambient temperature and without cover. Sulfatides with a sulfate ester on the internal galactose (SM2a, SM1a) are stained much less intensely than sulfatides with a sulfate ester on the terminal galactose (SM4s, SM3, and SM1b). Sialic acid-containing glycolipids and acidic phospholipids are stained weakly although the bands gradually change to brown in a few days unless the plate is covered with a glass plate or SaranTM (vinylidene polymer plastics) wraps. Also a dilute solution of toluidine blue stained SM4s on cellulose paper¹⁶² and methylene blue can detect sterol sulfates on TLC plates.⁴⁰

Exposure of the plate to iodine vapor,^{116,365,466,547} or immersing in a solution of Coomassie brilliant blue⁴⁰⁸ stain HSO₃-Chol and sulfatides nonspecifically. Observation under U.V. light after spraying rhodamine^{167,288} or primulin^{112,573} visualize all species of lipids. Primulin spray and observation under U.V. light did not interfere with mass spectrometry.⁵⁷⁴ When the saccharide is sulfated at position 2, or 3 of hexose or hexosamine, staining by periodate-Schiff reagent for detection of 1,2-glycol may be negative.¹¹⁶ Sulfated hexoses, 6-sulfoquinovose, and HSO₃-Chol, produce purple, greenish, and bluish purple color by anthrone sulfuric acid, Molisch⁶⁴⁷ or orcinol sulfuric acid stain²¹⁷ respectively.

(ii) *Densitometric and chemical analysis of sulfatides.* Azure A staining specific to sulfate ester has been successfully applied to quantitative assay of 0.5–4.0 nmol of sulfo-glycolipids (e.g. SM4s, bis-sulfo-ManGlc-E₂Gro)^{217,369} on TLC plate. It should be noted that every densitometric assay needs an appropriate standard compound because the molar absorption depends on the structure of sulfatides. Nonspecific staining with cupric acetate in aqueous phosphoric acid^{49,130,569,644} or

Coomassie blue and densitometry are able to quantitate 0.5–10 nmol of SM4s⁴⁰⁸ above 0.5–1.0 nmol of SM4s,⁵⁰⁴ and SM4g,¹¹ and HSO₃-Chol.³⁸⁴ Sulfatides were assayed densitometrically after staining with orcinol-H₂SO₄ reagent^{318, 412, 423, 453} when hexose composition is known. Alternatively, the band of glycolipid on silica gel powder scraped from the plate was hydrolyzed without extraction⁴²⁰ and component galactose analyzed enzymatically using galactose dehydrogenase and a fluorometer.^{97, 228, 606} HSO₃-Chol was able to be determined by the sophisticated gas chromatographic method after solvolysis using 5- α -cholestane,^{243, 427} [4-¹⁴C] HSO₃-Chol,³⁴⁸ or β -sitosteryl sulfate⁴⁰ as internal standards.

(iii) *Radioactivities.* [³⁵S]-labeled sulfo- and sulfono-glycolipids (cf. VI. A. 1) were detected by radioscan of TLC plates, autoradiography using X-ray films,^{111, 234, 632} or by using imaging plates. Cold SM4s can be ³H-labeled by tritiating with ³H₂ at the C4 and C5 positions of their Cer moiety.⁶¹⁶ High resolution is guaranteed for autoradiography while the sensitivity is not satisfactory.⁴⁶⁶ Detection of a doublet band of SM4s containing 200–300 dpm of [³⁵S]sulfate takes more than 1 month, although the β -emission energy of [³⁵S] is approx. 10-fold of ³H and comparable to ¹⁴C. Use of *En³Hance*TM and storage in a deep freezer at –80°C greatly improve the sensitivity of ³H-labeled compounds¹⁷³ but not substantially contribute to the intensity of [³⁵S]-bands. The densitometric quantitation of autoradiography film was linear between 170 and 300 dpm of [³⁵S]-SM4s and proportional to the counts obtained by the low-level liquid scintillation assay (Low-LevelTM, Packard) of the band of silica gel.⁴⁴³

Conventionally, the silica gel powder scraped from TLC plates was either directly counted with the scintillation solution (e.g. Aquasol),²³⁴ or sulfolipids extracted from the gel were assayed for radioactivities.^{632, 634} The limit of detection depended on specific radioactivities and detection methods. The detection/quantification imaging system BAS 2000 or 1500 (Fuji Film), with resolution higher than β -cameras and sensitivity higher than X-ray films, is rapidly replacing autoradiography and liquid scintillation counting.^{254, 425} The exposure time of radioluminography has been shortened to 1/10 of the time required for autoradiography with [³⁵S]-

Table 1. Antibodies interacting with sulfatides

Polyclonal
Sera from spontaneously diabetic BB rats, (+) SM4s (38%); sera from insulin-dependent diabetes patients, (+) SM4s (88%) ^{56, 122}
Rat sera (Heyman's nephritis) or the serum of rabbits immunized with rat proximal tubule, TLC-OL, ELISA, (+) SM2a, SB2 ²⁴⁶
Rabbit serum, antigen: SM4s, (+) SM4s; (–) GalCer, Gal-A ₂ Gro ⁵²¹ . Rabbit serum, SM4s, (+) SM4s, immunohistochemistry, (+) the initial part of the distal tubules or the thick ascending limb of rat kidney; (–) glomeruli, proximal tubules. ⁶⁴⁹ Rabbit sera (rSulf-1 and rSulf-2), SM4s, (+) SM4s. ⁴⁶² Rabbit sera (S antibodies), SM4s, (+ +) SM4s in sphingomyelin/cholesterol membrane; P antibodies, (+ +) SM4s in dipalmitoyl phosphatidylcholine/cholesterol membrane. ⁵³⁴
Sera (IgG ₁) of patients with HIV, TLC-OL, (+) SM4s, SM4g, SM3; (–), lyso-SM4s, SMUnLc ₄ Cer; immunohistochemistry, (+) oligodendroglial cells ⁸⁷
Sera of patients with cytomegalovirus (CMV) infection (IgG and IgM), TLC-OL, (+ + + high affinity) SMUnLc ₄ Cer, (+ + moderate affinity) SMUnLc ₄ Cer; (+ low affinity) 0.4 nmol SM4s, 0.5 nmol SM4g, 0.4 nmol SM3; (–) 20 nmol HSO ₃ -Chol ⁴³⁵
Sera from patients with autoimmune chronic active hepatitis (IgG), ELISA, (+) SM4s, SM4s-6 ⁵⁹³
Sera of patients infected with <i>Trypanosoma cruzi</i> (chronic chagasic disease) (IgA and IgM), ELISA, (+) HSO ₃ -Chol, 16% in healthy controls and 78% in carriers; (–) dehydroepiandrosterone sulfate ²⁰
Sera of patients with selective IgA deficiency, ELISA, (+) SM4s (32%), SMnLc ₄ Cer (11%) ²⁸
Sera of patients with leprosy (IgM), (+) SM4s, (–) HSO ₃ -Chol ⁶²⁷
Sera of patients with Guillain-Barré syndrome, (+), SM4s, ^{122, 123, 222, 435} SBA, TLC-OL, (+) SM4s (43%), diphosphatidylglycerol (48%), SMUnLc ₄ Cer and GM3 (11%) ⁵⁸⁷
Sera of patients with inflammatory demyelinating polyradiculoneuropathy, (+) SM4s, nLc ₄ Cer ³ IV-NeuAc. ¹²³
Sera from patients of inflammatory polyneuropathy after heart transplantation (IgM), (+) SM4s, GM1 ⁸
Sera of patients with predominantly sensory neuropathy, TLC-OL, ELISA, immunofluorescence microscopy, (+) SM4s, rat dorsal root ganglia neurons, human neuroblastoma ³³⁴
Sera from patients with autoimmune rheumatic diseases, (+) SM4s (absorbed by DNA, dextran sulfate, heparan sulfate and other anionic molecules) ¹⁴

—continued

Table 1—continued

Monoclonal
224-58 (IgM-κ), antigen: human brain myelin, ELISA, TLC-OL, (+) SM4s, SM4g, human Schwann cell membrane; (–) GalCer, gangliosides. ¹⁵⁰ (+) SM4g, SM4s, the apical ridge of boar spermatozoa; (–) SQ-A ₂ Gro ¹³⁵
2H12, JEG-3 human choriocarcinoma cell line, TLC-OL, ELISA, (+) SM4s, SM4g ⁶⁰⁰
412, ELISA, (++) SMUnLc ₄ Cer, (+) nLc ₄ Cer, (–) HSO ₃ -3GlcUCer. ⁵⁰³ (+) HNK-1 neoglycoprotein ¹⁶³
4A9E10 (IgG3), PLC/PRF/5 human hepatoma cells, TLC-OL, ELISA, cytofluorometry, (+) SB1a; (–) SM4s, SM3, SM2a, SB2, gangliosides. 2H6G5 (IgM), PLC/PRF/5, (+) SB1a; (±) SM3. 49-D6 (IgM), HepG2 hepatocellular carcinoma cells, TLC, ELISA, cytofluorometry, (+) SM3, SB1a. 7-E10 (IgM), HepG2, TLC, ELISA, (+) SM3, SB1a. 34-A4 (IgM), PLC/PRF/5, TLC, ELISA, (+) SM3, SB1a; ²⁰⁴ (+) SM4s ²⁰⁰
AGB43 (mouse IgG3κ), SM4s, TLC-OL, ELISA, (+) SM4s, SM4g, SM3; (–) SM2a, SB2, SB1a, HSO ₃ -Chol, gangliosides. ^{309,387,538}
AIC3IA2 (mouse IgG3), SM4s, (+) SM4s; (–) GalCer ²⁰⁶
AMR 20 (mouse IgM), GalCer, TLC-OL, ELISA, (+) GalCer, SM4s, SM4g, SM3, proteoglycans; (–) SM2a, SB2, SB1a, other neutral glycolipids, gangliosides ³⁰⁹
BMMK-33G (produced by lymphocytes of a breast cancer patient), TLC, ELISA, immunohistochemistry, (+) SM4s, human mammary glands, and other glandular epithelial cells ²²
CA 10 (IgM), SM4g, (+) SM4g, mouse spermatozoa, (–) SM4s, lyso-SM4s ⁹⁷
HF2-1/17 (human IgM from a lupus patient), TLC-OL, (+) SM4s, DNA ³⁹⁴
HNK-1 (mouse IgM), human T cell line HSB-2, ²⁴⁹ (++) SMUnLc ₄ Cer, (+) SMUnLc ₆ Cer. ^{249,381} IgM paraproteins from demyelinating polyneuropathy patients, ELISA, TLC-OL, (+) SMUnLc ₄ Cer, SMUnLc ₆ Cer, myelin-associated glycoprotein (MAG) ²²¹
L2 (334) (rat IgM), TLC-OL, (+) SMUnLc ₄ Cer. ²²¹ L9 (mouse IgM), acidic glycolipid mixture from the electric organ of <i>Torpedo marmorata</i> (Elasmobranchii), ELISA, TLC-OL, immunohistochemistry, (+) SMUnLc ₄ Cer, SMUnLc ₆ Cer, neuronal cell bodies ⁴⁴
M14-376 (human IgM), human lung cancer, TLC-OL, ELISA, immunohistochemistry, (+) SM4s, SM4g, kidney, testis, brain; (–) SM4s-Glc, lyso-SM4s, lyso-SM4g, SM3, SM2a, SM1a, SB2, SB1a, erythrocytes, granulocytes, lymphocytes ³⁷⁸
NGR50 (mouse IgG2a), SBA, (+) SMUnLc ₄ - and SMUnLc ₆ Cer (20 ng), human MAG, peripheral nerve; (–) rat MAG ⁶⁴²
O4 (mouse IgM), bovine corpus callosum, (+) SM4s, oligodendrocytes; ^{521,543} (–) GalCer, Gal-A ₂ Gro. ⁵²¹ ELISA, (+) SM4s, SM4g, oligodendrocytes; (±) cholesterol, HSO ₃ -Chol; (–) SMUnLc ₄ Cer. ²⁷ (+) SM4g, SM4s, lyso-SM4g; (–) SQ-A ₂ Gro. ¹³⁵ Immunohistochemistry, (+) Schwann cell lines; ⁵⁹⁴ (+) low grade and anaplastic astrocytomas; ³⁴⁶ (+) human oligodendrocytes, (–) human astrocytes from a 12-week fetus. ⁵⁰⁰
P3 (mouse IgM), GM3(NeuGc), SBA, TLC-OL, (+) NeuGc-containing gangliosides, SM4s, SM1b, human breast tumors ⁶¹⁴
OL-1 (rat mAb), (+) oligodendrocyte culture ¹²⁷
R (originally called 'mGalC', mouse IgG3), synaptic plasma membrane preparation, ELISA, (+) SM4s, SM4g, GalCer, Galβ-A ₂ Gro, GalSph, rat oligodendrocytes and Schwann cells; (–) SMUnLc ₄ Cer. ²⁷ TLC-OL, (+) HT-29, Caco-2 cells. ¹⁰³ (+) SM4s, SM4g, GalCer, Gal-EAGro; (–) SQ-A ₂ Gro ¹³⁵
SNH1 (mouse Ig), acidic glycolipid extract of melanoma (an upper phase of Folch partition), ELISA, TLC-OL, (+) SM4s, SM4g, SM3, SM2a, SB2, SB1a; (–) HSO ₃ -Chol, gangliosides, acidic phospholipids; I (+), SM4s; (+) tissues of neuroectodermal and hematopoietic origin, I (+), SM4s. ⁵⁰⁹
Sulf I (mouse IgG1), SM4s, TLC-OL, ELISA (+) SM4s, lyso-SM4s, SM4g, lyso-SM4g, SM3; (–) SM4s-Glc, ²¹⁷ SM2a, SB2, SB1a, glycosaminoglycans, sulfated glycoproteins; ^{84,125} (+) neutrophils. ¹⁷ (+) A and B cells of Langerhans islets, neutrophils and glandular epithelial tissues ⁵⁶
VESP 6.2, <i>Trypanosoma vespertilionis</i> , TLC-OL, ELISA, (+) SM4s ⁴⁵¹

Symbols and abbreviations: (+), positive interaction; (–), interaction not detected; I (+), inhibition of interaction; I (–), no inhibition of interaction; SBA, solid-phase binding assay; TLC-OL, TLC overlay; ELISA, enzyme-linked immunosorbent assay

Table 2. Molecules and cells interacting with sulfatides

Prokaryotes
Herpes simplex virus (HSV)-1, glycoprotein C, (+) SM4s, heparan sulfate ²⁰⁸
Human immunodeficiency virus (HIV), envelope protein gp120, TLC-OL and SBA, (+) SM4s, GalCer; (–) GlcCer, gangliosides. ⁶⁶³ TLC-OL using ¹²⁵¹ -gp120, (+) GalCer, GalSph, GM4, SM4s (5–10 nmol). ³⁶ ELISA, (+) SM4s, IV ³ -NeuAc-nLc ₄ Cer, GalCer, myelin-associated glycoprotein (MAG); (–) chondroitin sulfate, heparan sulfate. ⁶⁵⁶ TLC-OL (gp120), (+) GalCer, human colonic epithelial cell lines H-29 and Caco-2, ¹⁰³ (+) SQ-A ₂ Gro (0.2–1.0 μg/ml); ¹⁴⁸ (+) SM4g. ⁶¹⁸ Gp120, ELISA and by an immunosorbent assay on nitrocellulose paper, (+) SM4s, (–) GalCer, GM1; an immunosorbent assay on TLC plates, (+) GalCer, SM4s, GM1; (+) SM4s incorporated into the plasma membrane of lymphoid cells ⁶⁶²
Influenza virus, TLC-OL, (++) SM4s, GalCer, GD1a, nLc ₄ Cer IV ³ -NeuAc; (+) LacCer; (–) lyso-SM4s, GM3-NeuGc ⁵⁴⁹
<i>M. pneumoniae</i> cell, SBA, (+) SM4s, SM4g, SM3, laminins, glycoproteins; (–) gangliosides, neutral glycolipids; I (+) dextran sulfate, 3'-sialyllactose, human colon adenocarcinoma cell line (WiDr) ^{314,470}

—continued

Table 2—continued

M. pulmonis (rodents), TLC-OL, (+) SM4g, SM4s; (–) NeuAc-3LacNAc; I (+) dextran sulfate; I (–) SM4s, NeuAc-3LacNAc.³⁵⁵ *M. hominis*, *Ureaplasma urealyticum* (humans), *Ureaplasma diversum* (cattle), TLC-OL, (+) SM4g, SM4s.³⁵⁴ *M. hominis* (humans), TLC-OL, (+) SM4g (2 nmol), SM4s; SBA > 0.1 nmol/well, colon carcinoma cell line WiDr; I (+) dextran sulfate; I (–) 10⁻² M NaCl, HSO₃-Chol⁴⁴⁴

M. hyopneumoniae, SBA, TLC-OL, (+) SM4s, GM3, Gb₄Cer, cilia and ciliated cells of swine respiratory epithelia; (–) HSO₃-Chol; I (+), heparin⁶⁵²

Bordetella pertussis (virulent and avirulent strain), TLC-OL, (+) GalNAc β 4Gal-R (e.g. asialo GM2); (–) GM1, Gb₄Cer; *B. pertussis* (virulent strain), TLC-OL, (+) SM4s; I (+) dextran sulfate, fucoidan; (+) human colon adenocarcinoma cell (WiDr), I (+) heparin.⁴⁸ TLC-OL, (+) SM4s, (–) gangliosides, neutral glycolipids, a heparan sulfate proteoglycan¹⁷²

Escherichia coli, S-fimbriated, TLC-OL, (+) SM4s, SM4g, GalCer, LacCer; (–) gangliosides and other neutral glycolipids.⁴⁵⁴ *E. coli*, 987P fimbriae, TLC-OL, (+) SM4s (d18:1/hydroxy fatty acids, < 5 nmol), pig intestine.²⁸³

Helicobacter pylori, TLC-OL, (+) SM4s, SM3, GM3; flow cytometry, a gastric cancer cell line Kato III; (–) neutral glycolipids²⁵⁹

Helicobacter pylori, neutrophil-activating protein, TLC-OL, SBA, (+ + +) SM4s, SM1b, NeuAc(or NeuGc)nLc₄ or cCer; (–) neutral glycolipids, ganglio-series gangliosides⁵⁸⁹

Tetanus toxin, TLC-OL, (+) SM4s, SCLC (small cell lung cancer) cell lines; (–) lung tissue¹⁴⁴

Eukaryotes

Arthrobotrys oligospora lectin, TLC-OL, SBA, (+) SM4s, SM3, SM1b, HSO₃-Gal-ManGlcE₂Gro, acidic phospholipids⁴⁷⁶

Malaria circumsporozoite protein, the carboxy terminal region II, binding to sulfated glycoconjugate-Sepharose, (+) heparin, fucoidan, dextran sulfate; SBA, (+) SM4s, (±) HSO₃-Chol; binding of ¹²⁵I-protein, (+) hepatocytes,⁴⁴⁶ (+) SM4s, HSO₃-Chol.⁶² (+) SM4s³⁹⁰

Avian and human malaria plasmodium sporozoite, genes of thrombospondin-related anonymous protein (TRAP), contained sequences corresponding to thrombospondin-like sulfatide binding domain⁵⁸⁵

Antistatin (a salivary protein from the Mexican leech *Haementeria officinalis*), (+) SM4s, sulfated glycosaminoglycans; (–) HSO₃-Chol, gangliosides; I (+) dextran sulfate, heparin²⁰⁸

Melittin (a toxic bee peptide), (+) SM4s¹¹⁴

Laminins, TLC-OL, SBA, (+) SM4s, SM4s-6, SM3, SM2a, SB2, SB1a; I (+) fucoidan, dextran sulfate; I (–) chondroitin sulfate, hyaluronate, colominic acid, yeast phosphomannan; (–) neutral glycolipids, gangliosides.⁴⁷¹ The globular carboxyl domain of laminin α chain, SBA, (+) SM4s, heparin; after reduction and alkylation, (–) SM4s.⁵⁸³ SBA, TLC-OL, (+) SM4s, SM4g, SMUnLc₄Cer, SMUnLc₆Cer; I (+) heparin.⁵⁸¹ TLC-OL, (+) SM4s, SM3, SB2, heparin, heparan sulfate; erythrocytes (via SM4s at RBC surface membrane).⁴⁷² TLC-OL, (+) SM4s, SM4g, melanoma cell lines; (–) HSO₃-3Chol; SBA, I (+) for heparin, I (–) for SM4s, synthetic peptides from a thrombospondin type I repeats (e.g. KRFRKQDGGWSHWSPWSS) containing WSPW sequence.¹⁵⁷ (+), heparin, through the carboxyl terminal X-B-B-X-B-X, and B-X-B-X-B-X sequences.⁶¹ Laminin-1, SBA, electron microscopy, (+) a HNK-1 neoglycoprotein (through E8) (binding not abolished by reduction and alkylation or by urea treatment); heparin; I (+) heparin; I (–) SM4s.¹⁶³ E3 fragment of domain G4-G5, (+) SM4s, heparin; (–) a HNK-1 neoglycoprotein.¹⁶³ Laminin, (+) SMKT-R3 renal carcinoma cells, I (+) anti-laminin antibody, monoclonal antibody Sulf-I.²⁹⁴ TLC-OL, (+) SMUnLc₄Cer, mouse neurons and astrocytes.⁵⁰³ Laminin-1, laminin-2/laminin-4 and E8 fragment, mouse cerebellar neurons, cell adhesion (+); urea denatured E8, adhesion (–).¹⁶³ Laminin binding with α -dystroglycan or a schwannoma cell line RT4, I (+) SM4s, heparin, EDTA; I (–) chondroitin sulfate.³⁷²

Thrombospondins, TLC-OL, SBA, (+) SM4s, SM4s-6, SM3, SB2, SB1a, SM2a, heparin, fucoidan, (–) HSO₃-Chol; I (+) fucoidan, dextran sulfate, heparin;⁴⁷¹ (+) SMUnLc₄Cer, human melanoma cells.⁴⁶⁹ TLC-OL, (+) SMGb₄Cer,⁴⁰⁰ (+) SMGb₅Cer.³⁹⁹ (+) SM4s, heparin, melanoma cells; I (+) for heparin, I (–) for SM4s, synthetic peptides from a thrombospondin type I repeats (e.g. KRFRKQDGGWSHWSPWSS).¹⁵⁷

Thrombospondin, and laminin, TLC-OL, SBA, (+) SM4s; dynamic flow adhesion assay, (+) erythrocytes; I (+) high molecular weight dextran sulfate, chondroitin sulfate A²⁰⁰

Von Willebrand factor (high molecular coagulation factor VIII), (+) SM4s, SM4s-6, SM3, SB2, SB1a; I (+) dextran sulfate, I (–) other glycosaminoglycans.⁴⁷¹ (+) Human platelet;⁸³ rat and human platelets, 164–512 pmol/10⁹ platelets.⁴⁰ (+) SM4s, binds A1 domain.⁷³ Receptor binding, I (+) sulfobacins A and B, and SQ-A₂Gro from a gliding bacterium²⁶⁰

Properdin (human serum), SBA, (+) SM4s (> 62.5 nmol) and other sulfated glycosaminoglycans; (–), gangliosides, HSO₃-Chol, phospholipids; I (+) dextran sulfate, fucoidan.^{198,209} TLC-OL, ELISA, SM4g, SM4s (> 1 nmol), SM3, SB2, SM2a, SMUnLc₄Cer; (–) GM1, GD3; I (+) dextran sulfate.²⁴⁶

Vitronectin (mammalian blood plasma), the secondary structure prediction method, heparin, through X-B-B-X-B-X, and X-B-B-B-B-X-X-B-X sequences.⁶¹ SBA, (+) SM4s, HSO₃-Chol; (–) gangliosides; I for SM4s binding, (+) HSO₃⁻, fucoidan, dextran sulfate; (–) heparin, heparan sulfate; I for HSO₃-Chol binding, (–), HSO₃⁻, 1 M NaCl⁶⁴⁶

Fibronectin (an extracellular matrix protein), SBA, (+) heparin; (–) SM4s.⁵⁸³ SBA and cell lytic assay, (+) SMKT-R3 cells, I (–) anti-laminin antibody²⁹⁴

Amyloid P + Ca²⁺ (human serum), TLC-OL, (+) all sulfatides tested (0.1–5 nmol), GAG, mannose 6-phosphate; I (+) EDTA, glucose 6-phosphate, SM4s (5 μ M); I (–), lyso-SM4s³⁵⁷

Apolipoprotein E, (+) SM4s.¹⁵⁷ The secondary structure prediction method, (+) heparin, through X-B-B-B-X-B-X, and X-B-B-B-B-X-X-B-X sequences.⁶¹ (+) SM4g; I (+) synthetic peptides from a thrombospondin containing WSPW (e.g. KRFRKQDGGWSHWSPWSS)¹⁵⁷

—continued

Table 2—continued

Fibrinogen; (+) SM4s in 1:400 molar ratio¹⁷⁹

High molecular weight kinogen (participates in the intrinsic phase of blood coagulation as the complex with human factor XI), (+) SM4s, dextran sulfate, negatively charged surfaces (through domain 5)⁵⁸⁴

Human factor-H, TLC-OL, SBA, (+) SB2 (1 nmol), SM4s, SM4g, SM3²⁴⁶

Multicatalytic proteinase, human erythrocytes, TLC-OL using¹²⁵I-labeled enzyme, (+) SM4s, SM3; (–) neutral glycolipids, gangliosides⁴³⁸

NKR-P1 protein (natural killer cells), (+) heparin hexasulfotetrasaccharide > heparin tris-sulfodisaccharide > heparin bis-sulfodisaccharide > heparin monosulfodisaccharide > chondroitin 6-sulfate monosulfodisaccharide > chondroitin 4-sulfate monosulfodisaccharide > 3'-sulfated Le^a pentasaccharide > SB2 > SM2 > SB1a, SMUnLc₄Cer > SM1a > SM3 > SM4s in the order of larger amount of the protein bound (oligosaccharides as neoglycolipids).³⁵ (+) HSO₃-3Galβ-3(Fucα-4)-GlcNAcβ-3Gal-4Glc-neoglycolipid²⁹⁸

Human platelet, (+) SM4s; (–) HSO₃-Chol, gangliosides; I (+) von Willebrand factor, dextran sulfate, I (–), heparin⁸³

E-selectin + Ca²⁺, (–) SM4s.¹⁶⁹ E-selectin + Ca²⁺, TLC-OL, (+) HSO₃-3Galβ-4(Fucα-4)GlcNAcCer; HSO₃-3Galβ-4(Fucα-3)GlcNAc-1Cer.¹¹² TLC-OL with E-selectin transfected CHO cells, (+) HSO₃-3Galβ-3(Fucα-4)-GlcNAcβ-3Gal-4Glc-neoglycolipid.²⁹⁸ E-selectin-IgG + Ca²⁺, SBA, (+) 2,3-sialyl Le^x, SM4s; I (+) EDTA; (–) SMUnLc₄Cer.⁴¹⁶ Chimeras possessing the carbohydrate recognition domain of E-selectin, SBA, (–) SM4s.³⁰¹ E-selectin with substitution of EGF domain residues 124 and 128, SBA, (+) SM4s, HL60 cells⁴⁶⁷

L-selectin-IgG + Ca²⁺, TLC-OL, SBA, (+) SM4s, SM3, SB2, SB1a, sulfated fucans, SM4s-6, (±) HSO₃-Chol; (–) SM1a; I (+) SM4s, heparin.²²³ TLC-OL, SBA, (+) SM4s, SMUnLc₄Cer, heparin; I (–) EGTA; (+) sulfated Le^a/Le^x tetrasaccharides; I (+) EGTA.¹⁵³ L-selectin-IgG, TLC-OL, ELISA (30–100 pmol), (+) SM4s (50% of the interaction with sulfatides can be Ca²⁺-independent), SM4g, SM3, SM2a (weak), SB2, SB1a, synthetic sulfate analogues; Sia-Le^a; (–) gangliosides, neutral glycolipids.⁵⁵⁰ (+) HSO₃-3Galβ-3(Fucα-4)-GlcNAcβ-3Gal-4Glc-neoglycolipid.²⁹⁸ L-selectin-IgG + Ca²⁺, ELISA, (+) 2,3-sialyl Le^x; (–) octadecyl sulfate, sphingosine sulfate; L-selectin-IgG + Ca²⁺, (+) SM4s, SMUnLc₄Cer, desulfated SMUnLc₄Cer; I (–) EDTA.⁴¹⁷ L-selectin-IgG + Ca²⁺, SBA, (+) HEV-derived cell line, Ax.⁵⁸¹ The carbohydrate recognition domain of L-selectin, (+) SM4s³⁰¹

P-selectin-Ig + Ca²⁺, TLC-OL or ELISA, (+) SM4s, (–) Lyso-SM4s; I (+) SM4s, sulfated glycosaminoglycans; flow cytometry, granulocytes' lipids; I (+) SM4s.¹⁷ P-selectin-Ig + Ca²⁺, flow cytometry or SBA, (+) SM4s; I (+) heparin.²³ P-selectin-IgG + Ca²⁺, (+) sialyl Le^x, SM4s, SMUnLc₄Cer, desulfated SMUnLc₄Cer; I (–) EDTA; (–) octadecyl sulfate, sphingosine sulfate⁴¹⁷

Saposin B (sap-B), human, release of ³⁵S, (+) SM4s, lyso-SM4s, SM4g, lyso-SM4g.¹¹⁵ Tyrosine fluorescence measurement, (+) SM4s, Lyso-SM4s, lyso-SM4g, GM1, Gb₃Cer, sphingomyelin.³³⁰ TLC-OL, SBA, (+) SM4s, GM1.²⁰² (+) GalCer, GlcCer, SM4s; I (+) Mg²⁺, Ca²⁺, Zn²⁺.⁵³¹ TLC-OL, (+) SM4s, neutral glycolipids, gangliosides¹⁶⁴

Saposin C (sap-C), human, TLC-OL using each 15 nmol glycolipid, (+) SM4s, SM3, GM4, GM3, GM1, GM2, GalCer, LacCer, Gg₃Cer.¹⁶⁴

Tamm-Horsfall protein, (+) SM4s⁶⁴⁸

SLIP 1 (sulfolipid-immobilizing protein 1 isolated from male germ cells), (+) SM4g, SM4s³⁵⁰

Opioid peptides, (+) SM4s.³⁵⁶ Morphine, (+) phosphatidylserine.⁴ Levorphanol, (+) SM4s with hydroxy fatty acid (Kd 9.1×10⁻⁸), SM4s with nonhydroxy fatty acid (Kd 1×10⁻⁶)³⁵⁶

Myelin basic protein, (+) SM4s^{114, 493}

Amphoterin (a 30 kDa protein from rat brain), TLC-OL, SBA, (+) SM4s, SM4g, SMUnLc₄Cer, SMUnLc₆Cer, heparin, fucoidan; (–) bile acid sulfate, gangliosides, desulfated SMUnLc₄Cer^{382, 405}

Hepatic growth factor (HGF),¹²⁵I-HGF-OL, (+) SM4s, SM3, SB2, heparin, renal cell carcinoma (Grawitz) cell line SMKT-R3.²⁹¹ (–) SM2a, neutral glycolipids, gangliosides; I (+) dextran sulfate, heparin, fucoidan; I (–) keratan sulfate, hyaluronic acid²⁹⁵

Cytotactin (tenascin, a glial glycoprotein), SBA using radiolabeled cytotactin, SM4s (> 4 pmol); I (+) EDTA. (+) SM4s, HNK-1⁷⁹

Rat Schwann cells, TLC-OL, (+) SMUnLc₄Cer, SMUnLc₆Cer; (–) gangliosides.⁴¹⁶ Rat peripheral nerve myelin, TLC-OL, (+) SMUnLc₄ covalently attached to bovine serum albumin (BSA) via reductive amination; I (+) SMUnLc₄Cer⁵⁰⁷

Human melanoma cell line G361 (laminin-independent), SBA, (+) SM4s, SM4g > 100 pmol/cm²; I (+) fucoidan; (laminin-dependent) (+) SM4s, SM4g > 2.5 pmol/cm²; (±)HSO₃-Chol; I (+) fucoidan, dextran sulfate, anti-laminin antibody, anti-laminin-receptor polyclonal antibody, thrombospondins; (–) neutral glycolipids, gangliosides⁴⁷²

Human C32 melanoma cells, SBA, (+) SM4s, SM4g; I (+) thrombospondins, fucoidan, dextran sulfate; I (–) laminins, HSO₃-Chol⁴⁷²

Human melanoma cells, (+) thrombospondins⁴⁶⁹

Symbols and abbreviations: (+), positive interaction; (–), interaction not detected; I (+), inhibition of binding; I (–), no inhibition of binding; TLC-OL, TLC overlay; SBA, solid-phase binding assay; RIA, radioimmunoassay; ELISA, enzyme-linked immunosorbent assay

labeled sulfolipids, and the data can be transferred through a network to personal computers, analyzed for images and incorporation quantitated and calculated.

(iv) *TLC overlay*. Immunostaining using monoclonal antibodies^{121, 125} against sulfatides (Table 1) and overlay with adhesive proteins,^{468, 581, 613} lectins, bacteria and

vir^{263,391} (Table 2) are able to specifically locate individual sulfatide on TLC plates. Conversely, ligand specificities of the carbohydrate-binding proteins can be determined using synthetic neoglycolipids as immobilized probes on TLC plates.¹¹² The TLC overlay's potential disadvantage is that many monoclonal antibodies recognize antigens in unnaturally high densities but often are not capable of reacting with the same antigen expressed on the cell surface in low density.^{80,161,613} With Sulf I monoclonal antibody the lowest detectable amount of SM4s and SM3 on TLC was 5 pmol, whereas the lowest limit for SM4g was only 1 pmol.¹²⁵ The half-maximal binding was 1–3 pmol per well with SM4s on microtiter well while lyso-SM4g and lyso-SM4s showed very low sensitivities (> 1.5 nmol). After the washing and staining procedures used for the assay, only 36% of the standard [¹⁴C]-labeled SM4s remained on microtiter wells, and with lyso-SM4s, 50%,⁴⁷⁷ or even only 7%¹²⁵ of the ³H radioactivities remained indicating that the low sensitivities were due to the loss from the wells.

SMGb₅Cer and SMGb₄Cer were stained with the monoclonal anti-SSEA-3 antibody that interacted with the internal structure, R-3GalNAcβ-3Galx-4Galβ-R'. The structure 'R', HSO₃- of SMGb₄Cer and HSO₃-3Galβ- of SMGb₅Cer, did not interfere with the recognition by this antibody,^{399,400} while the monoclonal anti-SSEA-4 antibody, whose epitope was reported to be NeuAc2-3Galβ-3GalNAcβ-R, did not interact with SMGb₄Cer and SMGb₅Cer.⁴⁰⁰ This may indicate that a NeuAc residue on the terminal Gal of Gb₅Cer was essential for antibody binding and that a sulfate ester was not able to replace the sialic acid. Bands of SM4s (> 8–11 pmol) were detected with ¹²⁵I-labeled thrombospondins⁴⁶⁸ or hepatic growth factor (HGF)²⁹⁵ on TLC. The staining of SMGb₄Cer by thrombospondins was only 0.001 in comparison to the staining of SM4s.⁴⁰⁰ The bacterial-binding assay on TLC has the advantage that TLC plates present glycolipid receptors in a conformation similar to that of the eukaryotic cell membrane²⁶³ and discriminate the pathogenic prokaryotes using lipids containing SM4s from the tissue of 5 mg dry weight (Table 2).²⁵⁹

2. Ion-exchange Chromatography

Anion exchange column supports used at present include DEAE-cellulose,^{415,479,480} TEAE-cellulose,^{119,479} DEAE-Sephadex,^{217,385,559,567} DEAE-Sephacel,²³³ DEAE-Sephacel 6B,²⁰⁰ DEAE-Toyopearl,^{218,565,568} and QAE-Sephadex.⁶²⁴ Column chromatographies of anion exchangers have been routinely used for the initial phase of the purification of acidic lipids, including sulfatides (Figs 2 and 3), plant SQ-A₂Gro, sulfolipids of mycobacteria,¹⁴⁹ and halophilic archaea.²⁷³

Anionic lipids elute from DEAE-Sephadex columns with chloroform/methanol/ammonium acetate (5:10:1) system in the order of: (1) diphosphatidylglycerol,²³³ (2) monosialosyl gangliosides and free fatty acids; (3) monosulfoglycolipids (SM2a, SMiGb₄Cer/SMiGb₅Cer, SM2b, SM4s (d18:1/24:0), SM3 (d18:1/24:1), SM4s (d18:1/24h:0), SM4s-Glc (t18:0/24:0) in this order);^{217,233,564,565,568} (4) disialosyl gangliosides (GD3 and GD1a); (5) HSO₃-Chol and SM4s-Glc (t18:0/2-hydroxy fatty acids); (6) monosialosyl monosulfoglycolipid (SMGM1a);⁵⁶⁷ (7) bis-sulfated glycolipids (SB2 and SB1a),^{561,567} and (8) trisialosyl, and tetrasialosylgangliosides, and SMUnLc₄Cer.^{68,221,416} Sulfatides containing an *N*-acetylhexosamine eluted earlier than those composed of only hexoses.⁵⁵⁹ DEAE-Sephadex but not DEAE-Sephacel resolves SB2 and SB1a because the elution profiles are to some extent modified by the support (Sephadex, Sepharose, and Toyopearl) and the solvent (chloroform/methanol/water or methanol). SM4s-Glc (t18:0/hydroxy fatty acid) eluted later than HSO₃-Chol probably because glucose is adsorbed to Sephadex stronger than galactose. SM3²⁰⁴ and SMUnLc₄Cer²²¹ were effectively purified by elution with ammonium acetate in methanol.

Neutral lipids (e.g. cholesterol, and methyl esters of fatty acids released from acylglycerolipids) and zwitterionic phospholipids (e.g. sphingomyelin, and lyso-phosphatidyl-

choline) are collected in the pass-through fraction. Acidic phospholipids including diphosphatidylglycerol, and phosphatidylserine, phosphatidylinositol and their lyso-derivatives, eluted from the DEAE column along with monosulfoglycolipids (Fig. 2). The recoveries of SM4s from anion exchange columns ranged from 93²²⁸ to 95%.²⁷⁹ The assay mixture of GalCer sulfotransferase was first washed with Folch's partition and then applied to a small DEAE-Sephadex column to remove residual [³⁵S]sulfate, [³⁵S]PAPS and [³H]GalCer.^{137, 279}

Acidic oligosaccharides, derived from SM4s⁴⁸³ or SMUnLc₄Cer^{163, 416} by treatment with endoglycoceramidase, or ceramide glycanase respectively, were separated and quantitated using a Dionex⁴⁸³ or a Dowex column.¹⁶³ Three to five μg of SM4s was sufficient for the detection of the peak of the released HSO₃-Gal by pulsed amperometry.⁴⁸³

3. Adsorption Column Chromatography

Silicic acid is a potent adsorbent for acidic lipids of a wide polarity range.^{91, 479} The water in chloroform/methanol system attaches the effect of partition to the primarily adsorptive nature of silicic acids. SM4s and SM4g eluted usually with chloroform/methanol/water mixtures^{84, 318, 412, 558} closely after HSO₃-Chol.^{233, 559} A rat brain lipid mixture was able to be separated further into SM4g (E_{16:0}A_{16:0}), SM4s with nonhydroxy fatty acids and SM4s with hydroxy fatty acids (d18:1/24h:0) on a silica bead (Iatrobeds) (φ 60 μm) column²²⁸ (recovery, 88%). SM4s⁴⁴⁵ and more complex sulfatides, SM2a, SM2b, SB2, SB1a, SMiGb₄Cer, SMiGb₅Cer,^{565, 568} and SMGM1a⁵⁶⁷ were purified by HPLC using silica bead columns of 5–10 μm diameter. These sulfatides with the ceramide composed of d18:1 and nonhydroxy fatty acids usually elute significantly faster than those with an identical polar group but containing t18:0 and 2-hydroxy fatty acids.⁵⁶⁵

The salt form may change after passing through a silicic acid column because silicic acids contain cations including Na⁺, K⁺, Ca²⁺, and Mg²⁺.¹⁵⁹ Preequilibration with dilute salt solution may lead to a better separation of peaks because silicic acids have mild ion-exchange capacity²³⁶ and the addition of NaHCO₃, or KCl in the elution solvent may reduce the adsorptive strength resulting in better recoveries of sulfatides. Separation of highly polar sulfatides^{401, 407} including SMiGb₅Cer,⁵⁶⁴ SMGM1a,⁵⁶⁷ and HSO₃-8NeuGc2-6GlcCer³¹⁵ was also improved by using solvents containing up to 4 M of NH₄OH. For purification of radioactivity-labeled sulfatides in the total lipid extract from a few milligrams of tissues, nonradioactive SM4s^{32, 521} or SM2a²³⁰ was added as the carrier to guarantee high recoveries.

Chloroform/acetone systems have been used for the purification of less polar glycolipids from prokaryotes, plants, and animal brain because SM4g,^{95, 116, 232, 306, 381, 428, 511} SM4s,⁶⁰⁸ and HSO₃-2Man-4Glc-E₂Gro^{597, 598} show stronger affinity to acetone²³² than phospholipids. After washing out the acyl glycerides and cholesterol with chloroform, the glycolipids may be eluted from the column with solvent mixtures containing the increasing concentration of acetone in chloroform or dichloromethane, culminating with pure acetone^{32, 288} or acetone/methanol, 9:1.^{110, 610} Care must be taken in the formation of isopropylidene derivatives of galactolipids by acetone,³¹⁷ especially when the elution medium is acidic.³⁴⁹ HPLC using the combination of porous silica columns and solvents such as 2-propanol in hexane allows rapid separation of brain glycolipids with high resolution.⁸⁸

Florisil (aluminum silicate) has been a convenient column support to purify SM4s,^{180, 278, 480, 620, 621} SM3,⁴⁹¹ and HSO₃-Chol^{39, 647} with reasonable recoveries (e.g. SM4s, 85.5–87.0%²⁸²), while for more polar sulfatides, the adsorptive capacity of Florisil is too strong even after deactivation by the addition of 1–7% of water.^{189, 232} Use of paper or cellulose column chromatography for purification of SM4s was not successful.¹⁶²

Sulfatides were purified from the acidic glycolipids of platelets³⁹⁴ and the total lipid extracts from human spermatozoa¹¹ by the column of aminopropyl-bonded silica gel with recoveries between 89 and 98%. SMUnLc₄Cer and SMUnLc₆Cer were separated

from each other by HPLC with a Licrosorb-NH₂ column.⁶⁸ Florisil or silicic acids have been used as the adsorbent to separate peracetylated SM4s,⁴⁸⁶ SM4g,⁶⁰⁶ and SM1b.⁴⁷ Acetylation of hydroxyls drastically reduces the polarity of sulfatides and improves the recovery from the column. For acetylation, sulfatides should be incubated with acetic anhydride/pyridine at ambient temperature⁵⁶³ to prevent desulfation involving electrophilic attack by the acylium ion (CH₃C⁺=O) leading to scission of the S—O bond.⁶⁰³

A mixture of cholesterol, HSO₃-Chol, GalCer and SM4s was benzoylated to convert polar lipids except for HSO₃-Chol into nonpolar benzoate derivatives.²⁴³ HPLC determination of benzoylated and desulfated SM4s^{88,415,515,521} as well as perbenzoylated SM4s, SM4g, and lyso-SM4g⁵²⁹ have been reported. The recoveries of SM4s from rat spinal cord⁸⁸ and human urine⁴¹⁵ were 50–60%, and 30% respectively. The overall recovery of benzoylation/debenzoylation method of SM4s was 72%.⁵⁴⁴

The cerebellar amphoterin^{382,405} and anti-sulfatide antibodies⁴³⁵ were purified on affinity columns of SM4s-octyl-Sephadex. Purified SM4g was coupled to aminopropyl-glass with a photoactivated, heterobifunctional crosslinking agent, *N*-hydroxy-succinimidyl-azidobenzoate and used for the purification of antibodies.⁹⁷

III. ANALYTICAL METHODS

The selection of appropriate analytical methods, and reasonable recoveries in purification steps are essential to obtain reliable values for sulfatides as exemplified in the reports with widely different concentrations on the same tissue (Table 2). Lipid composition data should preferably be described in moles per unit fresh weight because direct comparison of lipid classes and sulfoamphiphiles become possible by using this mode of expression.⁴⁸⁰

A. Colorimetric Analysis

1. Chemical Analysis

Conventionally, inorganic sulfate released from sulfatides has been quantitated chemically⁴⁸¹ but recently the assay of either the chromogenic ion pair formed between sulfatides and Azure A,²⁸² or the chromogen or radioactivities obtained by the solid phase binding of specific antibodies or adhesive proteins are preferred.

(a) Sulfate

The assay of inorganic sulfate liberated from sulfoconjugates, as the precipitates of barium salts or by chelating barium ions with rhodizonate,⁴⁸² is reliable but laborious and not satisfactory in sensitivity.⁴⁸¹ Recently, an HPLC analysis of sulfate ion using 5–10 nmol of sulfatides was reported.²¹⁸

- (i) The *rhodizonate method*. The inorganic sulfate, released from sulfatide by HCl hydrolysis,^{232,588} Schöniger's oxygen combustion method⁶⁴⁷ or enzyme-catalyzed hydrolysis,⁴⁸² chelates with barium rhodizonate resulting in the destruction of the chromogen, which is assayed colorimetrically with the sensitivity of 1–10 nmol.⁵⁸⁸ Only oxidative mineralization in boiling HNO₃ released the sulfate from SQ-A₂Gro²⁷² because the negatively charged oxygen shields the carbon atom of the sulfonic acid from attacks by negative groups (e.g. Cl⁻).^{159,332}
- (ii) *Azure A*. The colored ion pair formation of the cationic dye, methylene blue, had been used primarily for the determination of relatively nonpolar sulfatides^{159,481} and later, and this assay method was modified for SM4s using Azure A as the dye.²⁸² The ion pair of the hydrophilic Azure A with sulfatides (absorption at 635 nm) is more lipophilic and partitioned into the organic phase of chloroform/methanol/0.1 M H₂SO₄ mixture. Azure A assay is the method of choice for SM4s, SM4g, and SDS for its sensitivity (above 0.5 nmol) and simplicity, although crude lipid preparations may yield optical densities higher than the sulfatide actually con-

tained^{519,615} because gangliosides, and acidic phospholipids interact weakly with this pigment.²⁸² In addition, the partition step involved in the procedure is susceptible to the lipophilicity of the aglycon. For instance, testicular SM4g yielded 1.22-fold of the molar extinction when human kidney SM4s was used as the standard,⁵⁶³ while more polar sulfatides such as SM3 and SB1a yielded only 0.82- and 0.19-fold respectively. Acetylation of sulfatides circumvented this difficulty.⁵⁶³ Peracetylated SMGb₄Cer,⁴⁰⁰ HSO₃-6ManGlc-E₂Gro,³⁶⁹ and SMGM1a,⁵⁶⁷ for instance, yielded optical densities of 0.097, 0.093, and 0.11/nmol respectively, comparable to that of SM4s, 0.101/nmol. However, caution is necessary to apply peracetylation/Azure A method to an unknown sulfatide because, for instance, (HSO₃)₂-2,6ManGlc-E₂Gro yielded only 1.5-fold absorption in comparison to the reference SM4s. Interestingly, TLC densitometry of the native form of this *bis*-sulfoglycolipid resulted in a value of 2.27 mol sulfate/mol *bis*-sulfoglycolipid.³⁶⁹

It should be noted that some lots of Azure A and rhodizonate^{481,482} yielded anomalously high background optical densities. The setback of the Azure A method, the possible contamination of the colored organic phase by the upper pigment phase, can be avoided by using a glass micropipette with a screw aspirator or by the use of a new mixture of solvents in which the desired layer floats above the excess dye.⁴⁵⁹

(b) Other constituents

Sulfoglycosphingolipids can be determined by fluorescamine method after hydrolysis of the band on silica gel (sensitivity 0.2 nmol, recovery 70%),⁶⁰⁶ or by the orcinol/sulfuric acid assay of silica gel powder containing glycolipids (50–200 nmol).⁴²⁰ A deep blue color of anthrone sulfuric acid reaction of 6-sulfoquinovose has an absorption maximum at 592 nm while glucose, galactose and glucose 6-sulfate peak at 625 nm. With free galactose liberated by acid hydrolysis of SM4g or SM4s on silica gel powder, the fluorescence intensity of NADH in the assay by β -galactose dehydrogenase was linear from 1 to 6 nmol of galactose.⁶⁰⁶ Fatty acids and trimethylsilylated methylglycosides obtained by methanolysis in HCl/methanol were quantitated gaschromatographically using heptadecanoic acid²³² and mannitol respectively, as internal standards,²²⁹ although mannitol should be used with caution for recovery especially when Ag₂CO₃ is used for neutralization.³⁹⁸ Also, cholesterol released from HSO₃-Chol,^{217,243,428,580} as well as Gal-1-alkylGro or GalGro²²⁸ from SM4g can be determined using appropriate internal standards.

2. Antibodies and Adhesive Proteins

Serum antibodies interacting with sulfatides have been identified in a number of autoimmune diseases and may be responsible for the pathology in some cases.⁵³⁴ On the other hand, sulfatides can be detected and determined by using artificial monoclonal antibodies^{121,204} and specific binding proteins^{112,468,550} as listed in Tables 1 and 2. The monoclonal antibodies Sulf I¹²⁵ and AGB43³⁰⁹ recognize SM4s, SM4g, their lyso-derivatives and SM3, while they did not interact with SM4s-Glc,²¹⁷ and other mono- and *bis*-sulfo glycolipids with sugar chains longer than disaccharides. M14-376 monoclonal antibody (human IgM) showed most stringent specificity interacting only with SM4s and SM4g.³⁷⁸ Obviously at least a part of the hydrophobic region of the sulfatides interacts with all the above antibodies.

(a) Solid phase binding assay

SM4s,^{122,506} SMUnLc₄Cer,^{381,417} and HSO₃-Chol²⁰ are adsorbed to plastic wells or silica gel layers in methanol,⁴⁰⁶ 50% aqueous solutions of ethanol, or chloroform/methanol and, after evaporation of the solvent, fixed with polyisobutylmethacrylate sometimes with the aid of admixed auxiliary lipids.^{27,200,468,468} TLC may, however, be less specific than the other assay systems, as binding on TLC plates can be altered by the polyisobu-

tylmethacrylate coating. Sulfatides are detected by using a ^{125}I -labeled ligand,⁴⁶⁸ or indirectly by using peroxidase conjugated secondary antibody¹²⁵ specific to the primary antibody, or streptococcal protein A.²⁰⁴ Microwell- and TLC-immunoassay are two orders of magnitude more sensitive than the conventional chemical methods. For instance, the densitometric response of TLC-ELISA of SM4s was linear between 15 and 250 pmol of SM4s.⁸⁴ A control TLC plate treated similarly, but stained by orcinol, showed that substantial amount of sulfatides remained on the plate after the immunostaining procedure.⁵⁵⁰ However, several reports cause concern regarding loss by background washing procedures.¹⁵³ The recovery of SM4s in microwells of ELISA ranged from 20%³³³ to 82%.³⁸¹ Even when ^{14}C -labeled sulfatides on microtiter well were incubated only once with Tris/BSA for 15 min, 73% of SM4s and only 13% of lyso-SM4s remained adsorbed to the wells.¹²⁵ Sulfatides with nonhydroxy-fatty acids were markedly better retained in microwells than sulfatides with hydroxy-fatty acids.

(b) *Histological and cytofluorometric analysis*

A considerable portion of polar lipids survives the routine dehydration procedure for paraffin embedding of tissues. Prior treatment of tissue sections with cold acetone, on the contrary, improved the staining.⁵³⁸ However, stronger extraction procedures using chloroform/methanol demolished the staining with the monoclonal antibodies^{44, 55, 210, 309, 316} or L-selectin,⁵⁸¹ suggesting that the cross-reactivity of the monoclonal antibodies with glycoproteins may be minimal. Arylsulfatase A activity was stained on frozen sections mounted on slides.⁶¹⁵ Human hepatocellular carcinoma cell lines,²⁰⁴ and SMKT-R3 renal carcinoma cells²⁹³⁻²⁹⁵ were analyzed by flow cytometry, after labeling with monoclonal antibodies or anti-laminin IgG then with fluorogenic second antibodies, producing distinct images of the cell populations expressing sulfatides on the cell surface.

B. *Spectroscopic Analysis*

About 30 years ago a few physical properties, including the melting point, 213°C ;²⁷⁸ optical rotation; and infrared absorption,⁶³⁷ were the only reliable parameters to identify SM4s. Today only $^1\text{H-NMR}$ can establish the full structure except for stereochemistry using a few to several hundred nmol of sulfatides^{369, 565} and when the amount is still meager, FAB, LSIMS⁵⁶⁶ or MALDI-TOF⁵⁶² can identify the partial structure in pmol or even fmol.

1. *Optical Rotatory Dispersion (ORD)*

Other optical rotation data reported for SM4s are, $[\alpha]_{\text{D}}^{18} = -0.14$ ($C = 3.56$ in pyridine),³⁶⁰ $[\alpha]_{\text{D}}^{24} = +2.84$,²⁷⁸ and $[\alpha]_{\text{D}}^{24} = -0.2$ ($c = 2.0$ in pyridine).⁵⁴⁸ ORD used today is much more sensitive and applied to the study of the configuration of component saccharides, e.g. SQ-A₂Gro⁵¹⁸ and HSO₃-2Man α -2Glc α -1E₂₀E₂₀Gro.⁵⁹⁸ The stereochemistry of the glycerol monoether from SM4g⁶⁰⁶ and glycerol tetra-ethers from lipids of a thermophilic archaea⁸⁹ was determined by optical rotation.

2. *Infrared Spectroscopy*

The infrared (IR) absorption was once frequently used for the characterization of sulfate esters in sulfatides,^{159, 232} lyso-sulfatides^{159, 307, 432, 569} and HSO₃-Chol.^{39, 388, 607} Recently, FT-IR apparatus replaced the conventional machines providing a better signal to noise ratio with several times smaller amounts of sulfatides (10–20 nmol).^{564, 565, 568}

The relatively broad absorption of the asymmetric O=S—O[−] stretching vibration at $1210\text{--}1265\text{ cm}^{-1}$ ^{68, 232, 273, 275, 557, 603, 637} is intense but overlaps with that of phosphate esters^{116, 370} (being not obvious, for instance, in the spectrum of HSO₃-PtdGro¹⁶⁷). Stretching mode of the Na⁺-SM4s decreased from 1219 to 1215 cm^{-1} on the addition of

Ca^{2+} probably due to bridge formation between two SM4s head groups.¹⁵⁹ After permethylation, the dimethyl ester of sulfate exhibited a stronger and more discrete sulfate ester ($-\text{O}-\text{SO}_2-\text{O}-$) doublet between 1405 and 1198 cm^{-1} .^{167,275} The $\text{R}-\text{O}-\text{SO}_2-\text{R}$ group of SQ-A₂Gro or deoxyceramide sulfonate¹² shows an asymmetric $\text{O}=\text{S}-\text{O}^-$ stretch at 1160–1170 cm^{-1} ,^{287,332,518} and symmetric $\text{O}=\text{S}-\text{O}^-$ stretch at 1030–1050 cm^{-1} .^{159,272,332} The $\text{O}=\text{S}-\text{O}^-$ of taurine amide was observed at 1096 cm^{-1} .^{29,665} The relative size of $\text{O}=\text{S}-\text{O}^-$ stretching vibration to OH absorption (3450 and 1060 cm^{-1}) depicted the sulfate/monosaccharide ratio and was used to discriminate *bis*-sulfoglycolipids from mono-sulfo compounds^{369,558,560} or SM4s from SM3.⁵³⁹ The peak height at 961 cm^{-1} due to $\text{SO}_3^{18}\text{O}^{2-}$ was used to quantitate SO_4^{2-} released by fission of $\text{O}-\text{S}$ bond.^{482,601}

For a primary sulfate group on C-6 of a hexopyranose in the C1(D) conformation, a sharp but relatively weak absorption of the C-O-S stretching vibration appears at 810–820 cm^{-1} , while the primary C-O-S of $\text{HSO}_3\text{-PtdGro}$ was observed at 840 cm^{-1} .¹⁶⁷ The sulfate at a secondary, equatorial position in the ring of a hexose,^{217,568} hexosamine^{149,560,565} and on C-8 of sialic acid³¹⁵ absorbs at 810–825 cm^{-1} , and the sulfate at a secondary, axial position at about 850 cm^{-1} .³⁶⁹ Although the above absorption has been frequently referred to as diagnostic of the sulfate location on sugar, there are often ambiguities because the presence of different aglycon groups and substituents on a glycoside may profoundly alter the position of the absorption near 850 cm^{-1} .¹⁸⁶

3. Mass Spectrometry

Because polar groups, especially sulfate, hinder the electron ionization (EI),⁵⁵⁹ the ionization of native sulfatides was carried out by field desorption (FD),¹⁷⁰ fast-atom bombardment (FAB),³⁶⁹ liquid secondary ion mass spectrometry (LSIMS), electrospray ionization (ESI),²¹⁵ or matrix-assisted laser desorption ionization (MALDI).^{562,570} FAB and LSIMS spectra of underivatized sulfatides, in both negative and positive ion detection modes, may be able to supply information on the number and the location of sulfate(s) on monosaccharides.^{215,324,437,566} Collision-induced dissociation (CID) and linked scan (or MS/MS) spectra contained product ions formed from a selected precursor (parent) ion or all precursors that give rise to a specific product (daughter) ion.⁵⁶⁶

(a) Electron ionization (EI)

Both molecule-related ions and carbohydrate sequence ions of sulfatides were obtained by EI-MS only after permethylation and desulfation.^{235,561} The peaks of the saccharide linked to C1 and C2 of the sphingoid with a fatty acid plus one proton were most intense. The sulfate(s) was localized by solvolytic desulfation of the permethylated sulfatides followed by C^2H_3 -remethylation of the hydroxyl that was originally occupied by the sulfate group.⁵⁶¹

(b) Fast atom bombardment (FAB), liquid secondary ion mass spectrometry (LSIMS) and electrospray ionization (ESI)

Soft ionization methods including LSIMS^{323,324,564–566} and FAB^{215,437} produce a series of sequence ions in addition to the molecule-related ions. These ionization patterns are essentially similar to those of gangliosides⁴⁵⁰ and unsaturated sulfated disaccharides obtained from glycosaminoglycans.⁹⁰ The amount of the sample for analysis can be reduced down to approx. 50–200 pmol and peracetylation or permethylation of sulfatides usually results in a better signal to noise ratio.^{218,339,369}

Positive ion LSIMS of the potassium salt of SM4g and lyso-SM4g showed the molecular ions in various cation forms including $[\text{M} - \text{H} + 2\text{Na}]^+$, $[\text{M} - \text{H} + \text{Na} + \text{K}]^+$, and $[\text{M} + \text{K}]^+$ ³²⁴ because Na^+ is abundantly present in water, test tubes, pipettes and the probe. Addition of 0.5% NaCl to the matrix intensified $[\text{M} - \text{H} + 2\text{Na}]^+$ and $[\text{M} - 2\text{H} + 3\text{Na}]^+$ ion from mono- and *bis*-sulfoglycolipids respectively, as well as positive ions containing ceramides ($\text{Yn} + \text{Na}$), which can be used to delineate the monosac-

charide sequence.⁵⁶² The advantage of positive ion detection is the relative preponderance of ions arising from lipophilic residues including EAGro,³²⁴ and sphingoids⁴⁵⁰ although the sensitivity is lower than the negative ion detection. The survey of *bis*- and *tris*-sulfoglycolipids led to the general formula for the molecule-related positive ions: $[M - nH + (n + 1)Na]^+$ where *n* represents the number of charges²¹⁵ in analogy with mono- and disialosylganglioside GM1 and GD1a which also yield $[M + Na]^+$ and $[M - H + 2Na]^+$ respectively.⁴⁵⁰

FAB or LSIMS spectra of HSO_3 -Chol,^{40, 384, 464} SM4g,^{10, 324} HSO_3 -3GalManGlcE₂Gro,^{126, 288} SMGM1a⁵⁶⁷ as well as other monosulfated glycolipids⁵⁶⁶ obtained in *negative* ion mode contained primarily the deprotonated molecule, $[M - H]^-$. The molecule-related ions of a potassium salt of SM4g,³²⁴ an ammonium salt of monosulfated glycolipid, e.g. HSO_3 -6ManGlc-E₂Gro,³⁸³ or di-ammonium salts of *bis*-sulfated glycolipids, e.g. $(HSO_3)_2$ -2,6ManGlc-E₂Gro,³⁶⁹ were similar to those of the corresponding sodium salts. Glycerol-type sulfatides, e.g. SM4g, and the E₂₀Gro-containing trihexosyl sulfatide, yielded a peak due to 'lyso-SM4g', and $[M - E_{20}]^-$ respectively, with minimal intensities of fragment ions from lipophilic moieties. Instead, the major negative ion from SM4g and $(HSO_3)_2$ -2,6ManGlc-E₂Gro was the ion consisting of the carbohydrate and glycerol.⁵⁶² When applied to *bis*-sulfoglycolipids (SB2, SB1a and $(HSO_3)_2$ -2,6Man α -2Glc α -1E₂₅E₂₀Gro), the molecule-related ions appeared in the form of sodium adduct ions, $[M + Na - 2H]^-$ accompanied by $[(M + Na - 2H) - NaSO_3 + H]^-$ ($[M - SO_3H]^-$) ions, which corresponds to the monodesulfated sulfolipid. SMUnLc₄Cer,⁶⁸ and SMGM1a,⁵⁶⁷ which have two species of the negatively charged group, also produced $[M + Na - 2H]^-$ accompanied by ions formed by the loss of a sulfate, $[M - SO_3H]^-$. SMGM1a contained, in addition, $[M + Cs - 2H]^-$. The mass number of the molecule-related negative ions in polysulfoglycolipids was generalized in the formula: $[M + (n - 1)Na - nH]^-$.²¹⁵ Sugar sequence ions, including $[HSO_3$ -Hex-*O*-HexNAc-*O*-Hex-*O*-Hex-*O*-Hex-*O*-Hex-*O*]- $^-$, and $[HSO_3$ -*O*-Hex-*O*-HexNAc-*O*-Hex(-*O*-SO₃H)-*O*-Hex-*O*]- $^-$ respectively, were abundantly obtained by the normal scan spectra of SMiGb₅Cer⁵⁶⁴ and SB1a.^{324, 566} The ions containing the sulfate and the parts of galactose $[SO_3OC(CH_2OH) = CHO]^-$ (*m/z* 169) and $[SO_3OCH = CHO]^-$ (*m/z* 139) were reported to arise from SM4s.⁴³⁷

The intense negative ion at *m/z* 97 (96.960) corresponded to the sulfate group plus hydrogen $[OSO_3H]^-$ (hydrogen sulfate anion) as confirmed by accurate mass measurement.³³⁹ Both *m/z* 80 ($[SO_3]^-$) and 97 were well visible when distinct from the matrix peaks³²³ and are the most convenient pair of ions to differentiate from a phosphate which yield the *m/z* 79/97 ($[PO_3]^-/[H_2PO_4]^-$) ion pair.

The single- and double-charged ions from synthetic mono-, *bis*-, and *tris*-sulfated Le^x-trisaccharide 1-propanols and from synthetic mono- and *bis*-sulfated sulfatides by ESI CID-MS/MS in positive²¹⁵ and negative modes²¹⁶ provided valuable information in identifying the sulfated sugar unit. Positive-mode ESI-mass spectrometry of a mixture of CaCl₂, GalCer and SM4s yielded $[GalCer-SM4s-Ca^{2+} - H]^+$ as the most stable noncovalent oligomer. These authors concluded that Ca²⁺ may mediate carbohydrate-carbohydrate interaction and might be involved in adhesion of the extracellular surfaces of the myelin sheath.³⁰⁸

(c) MALDI-TOF

Ions sublimated by matrix-assisted laser desorption ionization (MALDI) of lyso-SM4s⁵⁷⁰ and sulfatides⁵⁶² were analyzed by time-of-flight (TOF) mass spectrometry both in positive and negative modes. MALDI spectra can be obtained with a sulfatide mixture containing only a 1/100 of the sample required for LSIMS and the negative ion profiles of molecule-related ions are similar to those obtained with LSIMS of the same glycolipids. Mono- and *bis*-sulfoglycolipids produced intense $[M - H]^-$ and $[M + Na - 2H]^-$ accompanied by $[M - SO_3H]^-$ ions respectively. The spectra obtained in the positive ion mode showed $[M - nH + (n + 1)Na]^+$ (where *n* = 1 or 2) as well as the fragment ions

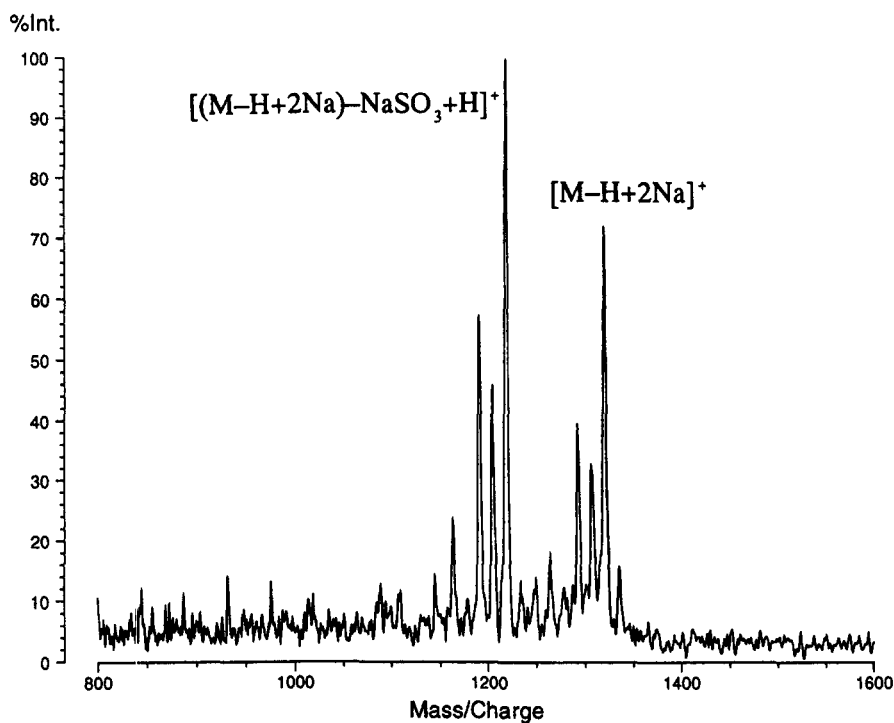
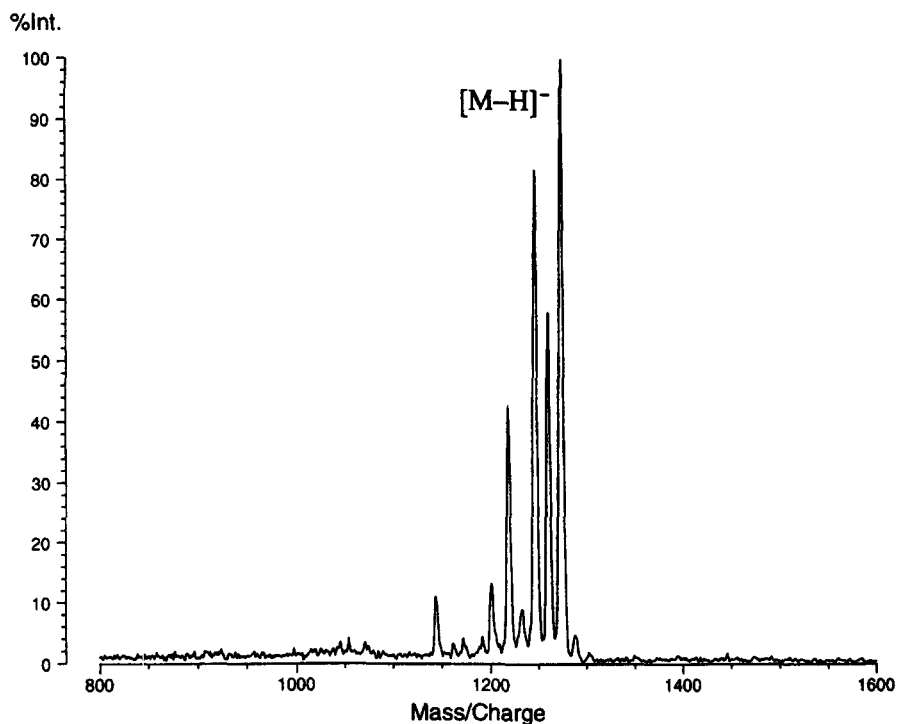


Fig. 4. MALDI-TOFMS spectra of SM2a from rat kidney in negative (upper panel) and positive ion modes (lower panel).

produced by the loss of one or two sulfate esters.⁵⁶² The initial setback of MALDI to produce only weak or no sequential ions is gradually improved by selection of novel matrices and by advanced hardware and ionization techniques including post source decay (PSD), (Fig. 4).

(d) Collision-Induced dissociation (CID)

The high and low-energy collision of the deprotonated molecule, i.e. $[M - H]^-$, produces sequence ions diagnostic for sulfatides.^{126, 215, 288, 437, 565, 567, 568} These include oligosaccharide ions produced by the cleavage of the glycosidic bonds on either side of the anomeric oxygen, sequentially from the nonreducing terminus (e.g. $[(H_3SO_3-O-Hex-(O)) - 2H]^-$, $[H_3SO_3-O-Hex-O-HexNAc-O-Hex(-O-SO_3H)-O-Hex-O-Hex-(O)-]^-$),^{565, 566} oligosaccharides linked to a lipophilic aglycon (e.g. $[(O)-HexNAc-O-Hex(-O-SO_3H)-O-Hex-OCer(t18:0/24:0)]^-$),⁵⁶⁸ $[H_3SO_3-O-Hex-O-Hex-O-CH=CHNH_2]^-$ ⁵⁶⁵ as well as the fragments produced by cleavage of the galactose (e.g. $[^-O_3SOC(CH_2OH)=CHOH]$ ⁴³⁷ or glucose^{215, 565} ring. $[M - \text{fatty acid}]^-$ and $[M - \text{alkyl}]^-$ were the most intense and characteristic product ions from SM4g and lyso-SM4g.⁵⁶² Low-energy CID-MS/MS of both SM4s and SM4s-6 produced ions of m/z 169 or 167 $[^-O_3SOC(CH_2OH)=CHOH]$ or its dehydrogenated form and 139 $[^-O_3SOCH=CHOH]$, while the ions at m/z 119 and 151 were specific to SM4s and SM4s-6 respectively.⁴³⁷ The CID-MS/MS of the $[M + Na]^+$ or $[M - H + 2Na]^+$ also supplied abundant information on the sulfated sugar unit as well as the sequence.^{215, 562}

(e) Aglycons and molecular species

The representative negative ions containing ceramides (Yn ions) include $[OCer]^-$,^{449, 564} $[SO_3-Hex-O-CH=CH-NH_2]^-$,⁴⁵⁰ $[O-Hex-OCer]^-$,^{450, 564} $[O-Hex-O-Hex-OCer]^-$,^{449, 564} $[O-Hex-O-Hex-O-Hex-OCer]^-$,⁵⁶⁴ $[O-HexNAc-O-Hex-O-Hex-OCer]^-$,⁴⁴⁹ $[(O)-Hex(-O-SO_3H)-O-Hex-OCer]^-$,^{324, 566} and $[(O)-HexNAc-O-Hex(-O-SO_3H)-O-Hex-OCer]^-$.⁵⁶⁶ Lyso-SM4s ($[M - \text{fatty acid}]^-$) was obtained only from SM4s with a 2-hydroxy fatty acid by FAB and low-energy CID in the negative mode, although the ion of the long-chain base was always detected by the positive ion mode.⁴³⁷ Peculiar ions ascribed to the mechanism called charge remote fragmentation (CRF) occur in some cases. For instance the high-energy CID spectra of the $[M - H]^-$ ion contained a series of ions at high mass region, that were evenly spaced by 14 Da ($[M - H - 16]^-$ and $[(M - H - 16) - (CH_2)_n]^-$), as a result of C—C bond cleavage in the ceramide moiety by CRF.⁵⁶⁵ The position of the double bond in 24:1 fatty acid was determined at C15 according to the intensities of CRF signals.⁴³⁷ FAB analysis of SM4s from bovine brain and erythrocytes showed an $[M - H]^-$ corresponding to a molecular species with d18:1/24:1 at m/z 888 and d18:1/16h:0 at 794³²³ respectively. Because other molecular species may yield incidentally $[M - H]^-$ at the same m/z , for instance with d18:1/22h:0 or t18:0/22:1 ceramide, comparison with $[M - H]^-$ of acetyl or methyl derivatives was necessary to delineate the number of hydroxyls.³⁹⁴

Using a specially designed, motorized TLC-FAB-MS probe with continuous desorption and scanning over a moving TLC plate, glycolipids with identical polar heads were well resolved into molecular species with differences in long-chain base and fatty acid.^{112, 264} The technique was applicable to SM4s and SM3 isolated from the human kidney (IV.B). The molecule-related ions were successfully recorded by TLC-LSIMS with ordinary probe and down to 1 pmol of neoglycolipids were identified.¹¹² Recently a simpler method of TLC blotting was developed, where glycolipids are transferred to polyvinylidene difluoride (PVDF) membrane [cf. II.C.1(a)].

4. ¹H-NMR

Thanks to the increase in sensitivity requiring only 10 nmol and 1 μ mol of sulfatides respectively, ¹H- and ¹³C-NMR spectroscopy developed into the method of choice for the structure determination of sulfatides. The monosaccharide composition, sequence, and linkage including anomeric configuration can be unequivocally determined by two dimensional ¹H homonuclear correlation spectroscopy (COSY),^{564, 567, 568} homonuclear Hartmann-Hahn (HOHAHA), nuclear Overhauser effect (NOE) analysis,⁵⁶⁸ and ¹H-¹³C heteronuclear COSY.³⁶⁹ The location of sulfates, and acyl esters^{10, 232} may be readily assigned by the chemical shift increment. The location of phosphate esters is determined,

in addition, by 3J and $^4J^{31P}$ - 1H and 13C - 31P couplings.³⁷⁰ Also, the structure of aglycon including 4-hydroxysphinganine (t18:0)^{565,568} and 1-alkyl-2-acylglycerol²³² is unequivocally determined. Moreover, some information on the conformation of molecule in solution was obtained (Iida-Tanaka, N. and Ishizuka, I. unpublished).

Most of the spectra since 1985 have been obtained in the solvent system of Dabrowski (deuterated dimethylsulfoxide/ 2H_2O , 98:2) with deuterium-exchanged glycolipids at 60°C facilitating the direct comparison of chemical shift data.⁸² The chemical shift and $^3J_{1,2}$ of anomeric protons can be measured with 10 nmol of sulfatides,⁵⁵⁹ and with the material above 100 nmol, 1H - 1H coupling constants of H1-H6 of sugar ring protons obtained by interpretation of 2D spectra, help to distinguish the monosaccharide species.^{9,131,369} Fortunately, the signals of the protons at α and β position to the sulfate ester (e.g. H6b, H6a, H5, and H2 of $(HSO_3)_2$ -2,6Man α -2Glc α -1E₂₀E₂₀Gro) emerge out of the bulk signal region due to the large downfield shift.³⁶⁹

The H1 (axial), H2 (ax.), H3 (ax.) and H4 (equatorial) protons of the 3-sulfated β -galactose in SM4s (Iida-Tanaka, N. and Ishizuka, I., in preparation), SM4g, SM3,³⁴⁷ SMiGb₅Cer,⁵⁶⁴ and SMGM1a⁵⁶⁵ resonated 0.09–0.12, 0.13–0.17, 0.64–0.71 and 0.29–0.37 ppm more downfield than those of the terminal galactose in GalCer, GalEAGro, LacCer, iGb₅Cer, and Gg₄Cer respectively. In contrast, the shift increments of the vicinal axial H2 of the internal 3-sulfated galactose of SM1a and SM2a were only 0.074 and 0.065 ppm respectively. The shift increments were largest with the α proton,¹⁸⁷ i.e. H3 of 3-sulfated glucose in SM4s-Glc (Δ 0.835 ppm), and H2 (Δ 0.70 ppm) of 2-sulfated mannose in $(HSO_3)_2$ -2,6Man α -2Glc α -1E₂Gro.³⁶⁹ The H1-H5, H6a and H6b of glucose 3-sulfate of SM4s-Glc (t18:0) resonated at more downfield by 0.122, 0.141, 0.835, 0.200, 0.080, 0, and -0.006 ppm respectively, in comparison to GlcCer (t18:0).²¹⁷ The smaller increment of the axial H4 of SM4s-Glc in comparison to the equatorial H4 of 3-sulfated galactose is in agreement with the results with synthetic oligosaccharides.¹⁸⁷ The chemical shift increment of H6a, H6b, and H5 of 6-sulfated GalCer (SM4s-6) was 0.300, 0.334, and 0.224 ppm respectively (Iida-Tanaka, N. and Ishizuka, I. unpublished), and H6a, H6b, and H5 of 6-sulfated mannose in HSO_3 -6Man α -2Glc α -1E₂Gro was 0.395, 0.351, and 0.158 ppm.³⁶⁹ In all the sulfated saccharides, the effect of sulfate esters on the coupling constant, $^3J_{H,H}$, and $^1J_{C,H}$ was minimal (Iida-Tanaka, N. and Ishizuka, I. In preparation).

The H1-H4 of the terminal 3-*O*-sulfated β -*N*-acetylgalactosamine in SM2b, SB2 and SMiGb₄Cer resonated at 0.01–0.05, 0.17–0.35, 0.39–0.59 and 0.45–0.47 ppm more downfield respectively, as compared with those of Gg₃Cer, SM2a, and iGb₄Cer respectively.⁵⁶⁵ The shift increment of H4 ring protons was significantly larger than those of vicinal H4s of the 3-*O*-sulfated galactose, probably due to the acetamide group. The law of additivity was applicable to the chemical shift increment of the protons of 3-sulfated GalNAc in SB2, and SM2b, vs. SM2a and Gg₃Cer. The shift increment of GalNAc-H3 in SM2b (0.579 ppm) added to that in SM2a (-0.146 ppm) made 0.433 ppm, which was roughly consistent with the increment of GalNAc-H3 in SB2 (0.472 ppm).⁵⁶⁵ Similar relationships were observed among GM1a(NeuGc) IV³ sulfate, GM1a(NeuGc), and Gg₄Cer.⁵⁶⁷

A hydroxyl or an electron-rich group in the aglycon which is spatially close to the ring proton but separated more than three bonds shifts the saccharide proton signals of sulfatides downfield.^{337,564} The shift increment of H1 of hexose by the hydroxyl at C4' of t18:0 and the 2-hydroxy group on the amide-linked fatty acid was approximately 0.02 ppm (Iida-Tanaka, N. and Ishizuka, I. In preparation). Inversely, when the amide on C2' of sphinganine was deacylated as in lyso-SM4s, chemical shift increments to H1' to H5' of d18:1, especially to the signals of H1', H3' and H5' were substantial.⁵⁶⁹ The dimethyl ester of HSO_3 -PtdGro¹⁶⁶ and permethylated 6-sulfo-Gal-ManGlc-E₂Gro from *H. salinarum* (*cutirubrum*)²⁷⁵ resonated at 3.98 ppm showed a sharp singlet at 4.0 ppm (three protons) attributable to the secondary S-*O*-CH₃ group (Fig. 5).

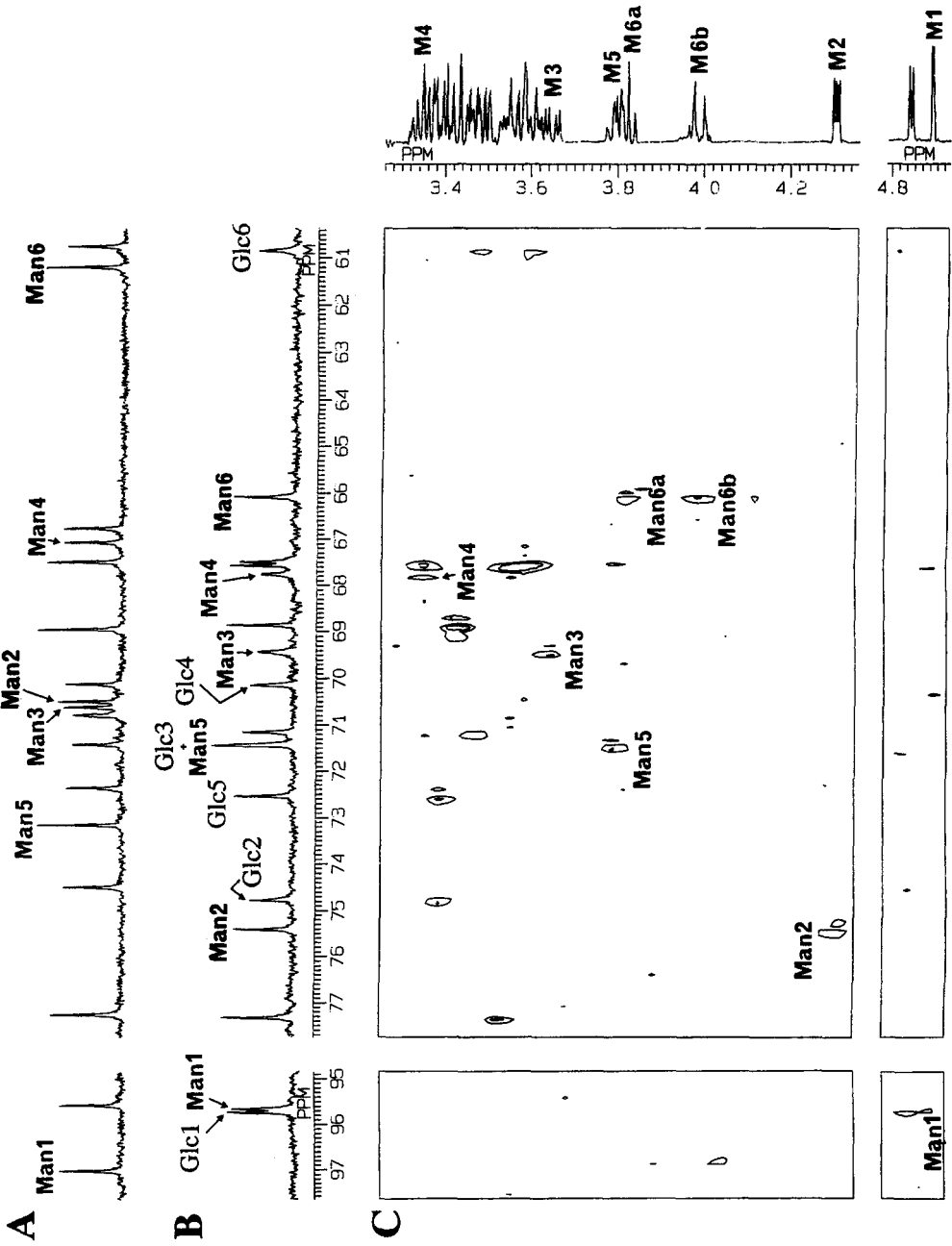


Fig. 5. Two dimensional H-C COSY spectrum (C) of $(\text{HSO}_3)_2\text{-}2,6\text{Man}_2\text{-}2\text{Glc}_2\text{-}1\text{E}_{20}\text{E}_{25}\text{Gro}$ from halophilic bacterium *Natrialba asiatica*. (A) and (B), the one dimensional carbon spectrum of this glycolipid and its desulfo-derivative respectively.

5. ^{13}C -NMR

One μmol of sulfatides is sufficient to obtain ^{13}C -NMR spectra either in one dimensional (1D) proton-decoupled or distortionless enhancement by polarization transfer (DEPT) mode, while several μmol of sulfatides are necessary to assign ^{13}C signals by relating to ^1H signals using two dimensional (2D) heteronuclear analyses such as ^1H - ^{13}C COSY,³⁶⁹ or ^{13}C - ^1H HMQC. The shift increment of the saccharide carbons by the sulfate ester was essentially similar to those with sulfated oligosaccharides.^{187,322} Downfield shifts of approx. 6 ppm for C-3 due to increased electronegativities accompanied by approx. 1.5 ppm of upfield shifts for the vicinal carbons at β -positions (C-2, C-4) were observed for the 3-sulfated galactose of SM4s, SM4g, SM3 (Iida-Tanaka, N. and Isizuka, I. In preparation), and lyso-SM4s⁵⁶⁹ as well as 2,6-bis-sulfated mannose of the sulfatide from a halotolerant archaea.³⁶⁹ ^{13}C chemical shifts are sensitive to the total environment of the particular carbon atom and may be affected by changes in molecular structure six or more atomic centers away¹⁵⁶ including 2-hydroxyl group of the fatty acid in ceramide. The downfield shift of protons at C-1, -3, and -5 of d18:1 in lyso-SM4s relative to GalSph indicated that the removal of the sulfate from SM4s might increase the degree of rotational freedom at C-2-C-5 of d18:1, thereby inducing the zigzag conformation of d18:1 in lyso-SM4s.⁵⁶⁹

IV. STRUCTURE

Variations in both hydrophilic and lipophilic residues contribute to the divergence of sulfatide structures in the biosphere.²³⁸ The variety of oligosaccharides may serve as the spacer at the cell surface.⁴⁹¹ Representative naturally occurring sulfatides including SM4g,¹⁴¹ and $\text{HSO}_3\text{-6E}_2\text{Gro}$ ²⁵⁷ were totally synthesized.

A. Sulfated Saccharide

The component sulfated monosaccharide in vertebrate glycolipids is exclusively the pyranosides (*p*) of β -Gal, β -Glc, β -GalNAc, and β -GlcU sulfated at the equatorial C3 hydroxyl. The saccharide structures of the major animal sulfatides, SM4s, SM3, SM2a, SB1a, and SMGb₅Cer are similar to the corresponding sialic acid analogues, gangliosides, GM4, GM3, GM2, GD1a, and V³-NeuAc-Gb₅Cer respectively.²³³ The sulfatides of echinoderms contain NeuGc sulfated at C8; archaeal sulfatides, α -Man*p* sulfated at C2 and/or C6 hydroxyl, β -3-sulfated Gal*p*, or *sn*-1-sulfated Gro; and mycobacterial sulfatides, 2-sulfated α -Glc*p*. Rat kidneys^{560,561} and an extremely halophilic archaea *Natrialba asiatica*³⁶⁹ contain bis-sulfated glycolipids that possess two sulfate groups attached to the separate monosaccharides and to the identical monosaccharide respectively. The 2-sulfate on mannose of a halophilic archaea is the only example of the axial glycolipid sulfate.

The sulfated saccharide interacts tightly with water molecules in the aqueous system. Water layers of thickness up to 44 Å, formed between the polar head groups in the gel state, were stable for several days.⁶ It has been known that proteoglycans are inflated with water to form gel with high internal osmotic pressure and thus contribute osmotically and, on the other hand, hamper the transport of ions. Hence the surface mucous exerts cellular osmoregulation by means of altering the ion gradients at the cell surface.⁷⁸ Similarly, sulfatides^{238,426} and $\text{HSO}_3\text{-Chol}$ ⁴¹⁸ may act as one of the ion barriers or ion traps on membranes and protect the cell against high extracellular osmolality in tight cooperation with the cell surface 'barrier mucus' (cf. IV. A), which provides the Donnan equilibrium.^{76,192} The sulfated glycosaminoglycan concentration and composition in 15 species of Crustacea, Pelecypoda and Gastropoda living in different degrees of salinity demonstrated a direct correlation between the logarithm of the sulfated glycosaminoglycans and the salinity for all the species analyzed.¹⁴⁶

B. Lipophilic Residue

The structure of lipophilic residues of glycolipids in eucaryotic cells varies considerably depending on the polar head groups⁵²⁵ and in turn the molecular species of sulfolipids with identical polar head groups vary with their specific tissue location.²³⁸ This diversity of lipophilic aglycons is essentially determined genetically but susceptible to the developmental stages and alteration of environments.⁴⁵ In contrast, the variation of the lipophilic residue concomitant with elongation or modification of carbohydrate chain is not prominent in archaea²⁷³ and eubacteria.^{117,238} For instance, in gram-positive cocci, fatty acid compositions of glycolipids are only slightly distinct from those of phospholipids.¹¹⁷ Although cellular accumulation of diacylglycerols, alkylacylglycerols, ceramides, long chain bases or sphingosine 1-phosphate might transduce hormonal signals, and regulate growth and apoptosis, the relation to sulfolipid has never been recognized.

1. Acyl-, Alkylglycerols and Ceramides

Diacylglycerols (A₂Gro): The aglycon of the major plant anionic glycolipid containing 6-sulfoquinovose is exclusively A₂Gro¹⁸⁸ (Fig. 1). The concentration of SM4g (1.69 μ mol/g) exceeds that of SM4s (1.35 μ mol/g) in the brain of Alaskan pollack indicating the dominance of glycerol-type sulfatides.⁵⁷⁷ SM4g in the brain of gadoid (cod) fishes contains mainly (>90%) A₂Gro with 18:1 and 16:0 fatty acids.

Alkylacyl- and dialkylglycerols (EAGro and E₂Gro): Alkylacyl- and diacylglycerols can be determined separately by TLC^{466,530} or gas chromatography.^{228,396} The mammalian testicular sulfatide (SM4g) contains almost exclusively a 1-alkyl-2-acyl-*sn*-glycerol²³² (E_{16:0}A_{16:0}Gro)^{606,631} (Fig. 1). This simple and saturated nature of the lipophilic residue is in striking contrast to testicular phospholipids. SM4g in aged humans contained alcohols and fatty acids of slightly longer chain than those in younger men.⁶⁰⁶ In HSO₃-3Gal-EAGro of the brain of rat³⁴³ and gadoid fishes,⁵⁷⁷ 16:1, 16:0 and 18:1 alcohols predominated. Vertebrate myelin SM4g contains varying proportions of EAGro.²³⁸ The relative amount of EAGro and A₂Gro forms of SM4g in the brain of rat at the peak of myelination (21 day) was about 1:1.^{227,452} The A₂Gro form of SM4g diminished more rapidly than the EAGro form⁴⁵² so that the ratio of EAGro to A₂Gro form at 175 days was about 13:1²²⁷ (V.C.2(a)). Diphytanylglycerylether (E₂₀E₂₀Gro) and its dimer, dibiphytanyl-diglyceroltetraether are the components of the membrane archaea.²⁷³ The genera *Halococcus* and *Natrialba* contain both E₂₀E₂₀Gro and E₂₅E₂₀Gro.²⁵⁶

Ceramides: Ceramides are distributed in plant, and fungi including yeast, and appeared in the animal kingdom from as early as sponge, the most primitive multicellular animal.¹⁹¹ The molecular species of sulfatides from rat kidneys were clearly depicted in LSIMS^{368,566,567} and MALDI-TOF spectra (Fig. 4). The presence of hydroxyl groups on C-2 of the fatty acid and C-4 of sphinganine enables sulfatides to form extensive hydrogen bonds on epithelial cell surface and thus strengthen the membrane^{264,359,657} and may also influence the transcellular influx of water.¹⁵⁸ It is notable that t18:0/24:0 predominated in SM4s-Glc from rat kidneys, whereas the major ceramide of the putative precursor GlcCer was t18:0/22 h:0.²¹⁷ Similarly, glycolipids in the intestinal cells of rat and man possess ceramides consisting of t18:0/2-hydroxy fatty acid.¹⁷³ Recently it was reported that the ceramides with 2-hydroxy fatty acids were selected as the substrate for galactosyltransferase at the endoplasmic reticulum,⁵¹⁰ and glycosphingolipids with hydroxy fatty acids including GalCer and SM4s were preferentially sorted to the basolateral membrane at the trans-Golgi network of MDCK cells⁶⁵⁷ shedding light on the regulatory mechanisms of the ceramide species.

2. Fatty Acids, Alcohols and Long-chain Bases

Saturated and monounsaturated fatty acids: The acyl component consisted predominantly of 16:0 (58%) in SQ-A₂Gro of the marine green alga, *Enteromorpha flexuosa*⁵¹⁸

and some green algae contained exclusively 16:0 acid in lyso-SQ-A₂Gro.¹³³ Whether the major fatty acid (14:0) of SQ-A₂Gro in sea urchin²⁸⁷ comes from algae or not is yet to be settled (V.B).

Mammalian testicular and sperm SM4g,²³² rat brain SM4g³⁴³ and porcine pancreatic SM4s⁴⁰⁷ contained 100, 80, and 60% respectively, of relatively shorter chain (16:0 and 18:0) saturated fatty acids. On the contrary, the predominant fatty acids of SM4s in the myelin of mammalian central nervous system were C24:1, and 2-hydroxy saturated.⁶¹⁸ SMUnLc₄Cer from mammalian central nervous system contained mainly relatively shorter chain fatty acids, 16:0, 18:0, and 18:1, amounting to 85% of the total fatty acids, whereas SMUnLc₄Cer from peripheral nervous system contained a large proportion (59%) of long-chain fatty acids (> 18:0).⁶⁷ Mammalian kidney sulfatides were rich in saturated 22:0, 23:0 and 24:0 acids^{401,561} (Fig. 4) and in particular, C22:0 which is found in quantities more than 10 times as much as those in the brain.³⁰² SM4s and SM3 of human liver,⁵³⁹ and SM4s of rabbit serum⁶⁵³ also contained 22:0 acid in proportions comparable to 24:0.

2-Hydroxy fatty acids: The ceramide of HSO₃-8NeuGc2-6Glc β -1Cer from sea urchin consisted of 4-hydroxysphinganine (t18:0), and 2-hydroxylated acids (22 h:1, 23 h:1, 24 h:1).³¹⁵ The typical ceramides of vertebrate brain SM4s were d18:1/24:1 and d18:1/24h:1.⁵⁷⁷ The content of 2-hydroxy acids in GalCer and SM4s was lowest in the brain of hatching chicken, shark and tuna.⁶³⁸ Substantially lower concentrations of hydroxy fatty acids were also found in the brains of cartilaginous deep-water fish compared with surface fish,³⁴⁴ and SM4s from Alaskan pollack and other gadoid fishes lacked in hydroxy fatty acids.⁵⁷⁷ Both GalCer²⁴⁷ and SM4s^{1,302} of mammalian brain contained high proportions of long chain fatty acids with a 2-hydroxy group in D-configuration.⁴⁴⁸ Nearly all of the 2-hydroxy fatty acids found in brain lipids were the constituents of these two glycolipids. In the GalCer-deficient animals, however, the ceramide with 2-hydroxy fatty acids was used for the synthesis of GlcCer and sphingomyelin.⁷⁵ During the first post-natal month of Wistar rats, the ratio of hydroxy- over non-hydroxy-species (HFA/NFA) of cerebral GalCer increased to 2.0, whereas the HFA/NFA ratio for cerebral SM4s declined to 0.6 in the same period.^{88,431} The developmental change of 2-hydroxylation in the nervous system may be regulated by thyroid hormones.⁵⁷⁸

SM4s and SM3 of the extraneural organs including human liver,⁵³⁹ mammalian kidneys,^{398,559} and rabbit serum⁶⁵³ were also characteristic for high contents of 2-hydroxy fatty acids. The contents of SM4s with 2-hydroxy fatty acids in the total SM4s in the kidney of human,³⁶² porcine intestine,⁵⁴⁸ and rabbit fundic mucosa⁴¹³ have been reported to be 89, 67 and 80% respectively, whereas SM3 of the kidney of human²⁶⁴ and house musk shrew contained only 0 and 25%³⁹⁸ respectively. A similar fatty acid pattern was found in human urine SM4s, which probably originated from the kidney.^{302,609}

Polyunsaturated and other fatty acids: The major fatty acids in the chloroplast SQ-A₂Gro were *trans*-hexadecenoic and 18:3.^{63,188,284} The major characteristic fatty acids of the sulfonolipid isolated from a marine diatom *Nitzschia alba* were 16:1- Δ^3 -*trans* for deoxyceramide sulfonate, and 20:5 for SQ-A₂Gro.¹² SQ-A₂Gro from the thermoacidophilic rod *Sulfolobus acidocaldarius* was unique in containing 17br, 17cyc, and 19cyc (cyc designates 11-cyclohexylundecanoic and 13-cyclohexyltridecanoic).³³² It has been reported that unsaturated fatty acids were abundantly present in the brain ganglioside of the stenothermic cold-water fish species,²¹ while no systematic studies have been performed on the lipophilic residues of sulfatides and the environmental temperature.

Long-chain alcohols: The synthetic sulfated oligosaccharides that have long alkyl groups (C12–C18) at the reducing end exhibited tens to hundreds times higher anti-HIV activities than those of the corresponding sulfated oligosaccharides without alkyl groups, probably due to the detergent effects.²⁷⁷

Long-chain bases: The predominant long-chain base of SM4s in the vertebrate nervous system is sphingenine (d18:1). For instance the long-chain bases of SM4s from the brain of Alaskan pollack and other gadoid fishes consisted exclusively of d18:1.⁵⁷⁷ The long-chain bases of SM4s from the salt gland of dogfish²⁶⁹ and the equine kidney^{181,570} were

unique in containing comparable amounts of dihydroxy and trihydroxy long-chain bases with 16–20 carbons. Bases with methyl branches at position 2, 3 or 4 from the methyl end were relatively abundant in shark brain SM4s.²⁶⁹ Sphinganine (d18:0) predominated in the long chain base fraction of SM4s from the brain of snake, tadpole, and frog.^{485,578}

The proportion of t18:0 was approx. 8% in SM4s from the whole human kidney, and SM4s and GM4 from rat kidney.⁵⁵⁹ Interestingly, the proportion of t18:0 was 8% in the cortex and 34% in the medulla of human kidneys, and 11% in the cortex and 27% in the medulla of bovine kidneys, forming a concentration gradient from the cortex to medulla.⁴⁹¹ On the other hand, more than 70% of the long chain base was t18:0 in SM2a, SM2b, and other rat kidney sulfatides with oligosaccharide chain longer than three monosaccharides.⁵⁶⁵

C. Chemical Modification

1. Sulfate

Partial (limited) acid or alkali catalyzed hydrolysis of the sulfate ester of sulfatides aims to remove the sulfate ester at a mild condition without destroying the remaining part of the molecule,^{456,603} while partial hydrolysis with stronger acid may provide sequence information. Solvolysis in dioxane or pyridine and acid catalyzed desulfation developed for steroid sulfates, are also effective especially for less polar sulfolipids. The discrimination of acid-catalyzed 'hydrolysis' and 'solvolysis' is marginal and both terms have often been used synonymously. Enzymatic release of inorganic sulfates was successfully applied to urinary metabolites of steroids and bile salts.⁴⁸² Invertebrate sulfatases⁶⁴⁵ or partially purified human placental arylsulfatase A¹⁶⁵ released approx. 70% of the sulfate esters from SM4s. However, there has been no broad specificity sulfatase available, which can quantitatively cleave the sulfate esters of sulfatides.⁴⁵⁶ 6-Sulfoquinovose was released from the lipophilic group by incubation with β -galactosidase from *E. coli*.²⁸¹

The sulfate ester of sulfoamphiphiles is easily solvolyzed in various organic, oxonium ion-forming solvents.¹⁵⁹ The HSO₃-Chol fraction was solvolyzed⁵⁸⁰ or 'hydrolyzed' in acidified organic solvents.²⁴³ The sulfate esters of more simple sulfatides including SM4g, SM4s or HSO₃-PtdGro can be quantitatively removed by solvolysis in dioxane at 100°C for 10 min^{159,228,232,318} when water is not present.¹⁶⁷ Even sulfatides with two or three monosaccharides were completely desulfated: SM2a (methanolic dimethylsulfoxide containing 4.5 mM H₂SO₄ at 80°C for 3 hr),⁵⁵⁹ HSO₃-ManGlcE₂Gro (dioxane/pyridine (1:1, v/v) at 100°C for 2 hr),^{597,598} HSO₃-GalManGlcE₂Gro (4 mM HCl in anhydrous tetrahydrofuran at ambient temperature for 90 min,²⁷³ and the ganglioside containing 8-O-sulfated NeuGc (dioxane or pyridine/dioxane at the same temperature.^{297,315} The cleavage of the sulfate ester was also complete in 4–5 hr when the Kantor and Schubert method of mild desulfation for chondroitin sulfates in 0.05 M methanolic HCl is applied to SM4s,^{362,415} SM3,⁵³⁵ SM4g,^{228,232} or HSO₃-3Gal β p-6(Galf α -3)Manp α -2Glc α -1E₂₀E₂₀Gro.⁵²⁷ Analogously, SMUnLc₆Cer was desulfated⁶⁸ in 16 hr with 0.1 N methanolic HCl.²²¹

On the other hand, solvolysis in dioxane or pyridine/HCl is often inadequate for sulfatides with more than two monosaccharides (e.g. SM1b, SMiGb₅Cer) or with the sulfate ester at the internal monosaccharide (e.g. SM2a).⁵⁶¹ Incubation with 5 mM HCl in dimethylsulfoxide containing 0.5% methanol at 80°C for 1–2 hr quantitatively converted SM2b,⁵⁶⁵ SMiGb₄Cer,⁵⁶⁴ and SMGM1a⁵⁶⁷ into Gg₃Cer, iGb₄Cer, and GM1a respectively, whereas SM2a with an internal sulfate ester was unchanged after 1 hr.⁵⁶⁵ After acid-catalyzed solvolysis of SB2 in 8 mM H₂SO₄ in 10% methanol in dimethylsulfoxide at 80°C, most of SB2 was converted into the monosulfo-analogue (SM2a) in 10 min, confirming that the sulfate ester on the terminal monosaccharide was much more labile to solvolysis,⁵⁶⁰ whereas the internal sulfate ester of SM2a was resistant.⁵⁶⁵ These obser-

vations suggested that the sulfate at C3 of the internal Gal is protected against acid by the GalNAc residue at C4 similarly to the internal sialic acid of GM1 ganglioside.²²⁹

Sulfate esters are stable under mild alkaline conditions used for deacylation.^{361,362,535,570} When 6-sulfates of Glc, Gal and Man were treated at higher temperatures (e.g. 0.1 M NaOH, 100°C) 3,6-anhydrohexosides were produced provided that C-3 has a free hydroxyl group.^{159,603} Conversely, alkali-treatment of SM4s, which contain a 3-*O*-sulfate and a free C6-hydroxyl, yielded a substantial amount of 3,6-anhydro-GalSph with the anhydrogalactose in the boat conformation.⁴³²

2. Limited Acid Hydrolysis and Periodate Oxidation

More than 60% of the sialic acid was released from GM3 in 0.1 M HCl at 37°C, whereas only 10% of the sulfate ester of SM4s was cleaved suggesting that the terminal sulfate ester was more resistant than the terminal sialic acid.⁴¹³ However, 5 mM HCl in dimethylsulfoxide containing 0.5% methanol cleaved preponderantly sulfate from SMGM1a to yield GM1a, whereas the treatment of the same sulfated ganglioside with 10 mM formic acid²²⁹ selectively cleaved sialic acid and yielded SM1b confirming that the effect of solvolysis is specific to sulfate esters.⁵⁶⁷ Also oligosaccharides with 8-*O*-sulfated *N*-acetylneuraminic acid were released from sea urchin sulfated gangliosides by mild acid treatment using 0.1 N trifluoroacetic acid.²¹⁸ The ester migration under acid conditions, akin to that observed with certain sugar phosphates, does not occur readily with sugar sulfates.⁶⁰³ Mild acid hydrolysis in the chloroform/methanol/water or 0.25 M methanolic HCl system was successfully applied to the identification of desulfated and partially hydrolyzed products from (HSO₃)₂-ManGlcE₂Gro,³⁶⁹ and HSO₃-GalManGlcE₂Gro²⁷⁵ respectively. 3-Sulfated galactose in SM4s and SM3³⁶⁶ was not attacked by periodate.^{228,362,637} The glucose at the nonreducing end was also resistant to periodate because of inaccessibility of the reagent to glucose due to micelle formation.

3. Lipophilic Residues

Stronger alkali and well-controlled conditions are necessary to chemically deacylate SM4s because the amide-linked fatty acid of ceramide is resistant to mild alkali and the yield is only 30–60%.^{125,307,432,569} Recently, a rapid method was developed to obtain lyso-sphingolipids in only 2 min using a microwave oven.⁵⁷⁰ Also a novel enzyme that hydrolyzes the *N*-acyl linkage between fatty acids and sphingosine bases in ceramides of various sphingolipids was purified from the culture filtrate of *Pseudomonas* sp. TK4, which released 59% of fatty acid from SM4s at the optimum condition.²⁴¹ The acyl ester on the glycerol of SM4g was released by a lipase activity of rat liver lysosome.⁴⁶⁵

The hydrazine-nitrous acid fragmentation procedure provided valuable information regarding the extent and position of sulfation on the various carbohydrate units. The ³H-labeled HSO₃-3AnTalH₂ [AnTalH₂ = 2,5-anhydro-D-talitol] was prepared from SB2 by hydrazine-nitrous acid treatment followed by NaB³H₄ reduction.⁹⁸ Similar treatment of bovine lutropin yielded HSO₃-4AnManH₂ and HSO₃-4AnTalH₂. The diacylglycerol residue can be released from SQ-A₂Gro by periodic acid oxidation in methanol and fission of the glycosidic linkage with 1,1-dimethylhydrazine.¹⁹³

4. Methylation

Acetolysis followed by reduction yields partially methylated alditols, are then separated by TLC²³⁰ or gaschromatography after acetylation,⁵⁶¹ providing clues to the position of the sulfate ester and glycoside attachment.⁵⁶⁵ Treatment with methylsulfinyl carbanion^{13,74} leads to deacylation of acyl esters, dephosphorylation, and *N*-methylation of fatty acid amide including ceramide,^{564,568} whereas desulfation during the process was minimal.^{559,564,568} By the use of NaOH instead of NaH,⁷⁴ the permethylated glycolipids recovered from the lower phase of Folch's partition showed homogeneous bands on

TLC without by-products or contaminants, indicating that further purification was not necessary.⁵⁶⁴ For methylation of the sulfate and phosphate esters of HSO₃-PtdGro, the free acid form of the sulfolipid was treated with ethereal diazomethane. The dimethyl ester was unstable in solution and decomposed within several hours to diphytanyl glycerol ether.¹⁶⁷

V. FUNCTIONAL DISTRIBUTION

Sulfatides are distributed in three widely separated phyla, halophilic archaea,²⁵⁶ Mycobacteria¹⁴⁹ and animals of the deuterostome lineage from echinoderms to vertebrates.²³⁸ On the other hand, the sulfonoglycolipid, SQ-A₂Gro, is distributed in cyanobacteria, green and brown algae and thylakoid membrane of chloroplasts (cyanobacterial symbiotes) of higher plants.^{63,498} Table 3 summarizes the structure and cellular concentration of sulfatides and related lipids classified according to the phylogenetic tree and the tissue of animals.

A. Prokaryotes and Plants

Extreme halophiles (Family *Halobacteriaceae*), which can grow in media containing up to 4 M of NaCl, belong to the distinct taxon Archaea (formerly Archaeobacteria) of the kingdom *Prokaryotae*.²⁵⁶ Sulfatides of halophilic archaea are useful taxonomic markers. Genera *Natrialba*, *Halorubrobacterium* and *Halobacterium* contain (HSO₃)₂-2,6Man α -2Glc α -1E₂₀E₂₀Gro (and E₂₅E₂₀Gro),³⁶⁹ HSO₃-2Man α -4Glc α -1E₂₀E₂₀Gro; and HSO₃-3Galp β -6Man α -2Glc α -1E₂₀E₂₀Gro + HSO₃-3Galp β -6(Galf α -3)Man α -2Glc α -1E₂₀E₂₀Gro respectively.²⁵⁶ *Halococcus* and *Haloferax* contain HSO₃-6Man α -2Glc α -1E₂₀E₂₀Gro, while HSO₃-PtdGro (E₂₀E₂₀ analogue) is present in *Haloarcula*, *Halobacterium*, and *Halorubrobacterium*.¹⁰⁰ *Haloferax mediterranei* maintains a value (1.3–1.6), for the number of negative charges per mol ionic lipid, which is comparable to that (1.7–2.4) for *H. salinarum* and for *Haloarcula marismortui* (1.7–2.3).³²⁶ With the growth of *Haloferax mediterranei* in media of increasing salt concentrations (3–7M), gradual increase in the relative proportion of the sulfated diglycosyldiether (S-DGD-1) (21–37 mol%) was observed. In addition, the proportion of the diphytanylether analogue of HSO₃-PtdGro was 1/3 as great in 7M salt as in 3M salt.³²⁵ A halotolerant strain of *Staphylococcus epidermidis* also responded to high NaCl concentrations (up to 4M) in the growth medium with an increase in the percentage of 6-glycerophosphoryl- β -gentiobiosyl-A₂Gro and cardiolipin.³⁰³ As a result, the average number of negative charge/mol phospholipid increased from 1.01 to 1.14. All data above supported that these anionic lipids play a role in controlling the ion permeability of the cytoplasmic membrane similarly to sulfatides of the plasma membrane of animal transporting tissues.²⁶⁷ Sulfoamphiphiles at the outer leaflet of the cell membrane appear to cooperate with the highly acidic surface layer glycoprotein as ion-barriers³⁷⁶ (cf. IV.A; V.C.1).

Sulfatides of *M. tuberculosis* were discovered in the search for a strain reactive to weakly basic phenazine dye, neutral red, located at or near the surface of virulent human and bovine strains of the pathogen.¹⁴⁹ Mycobacterial 'sulfolipids' contain the monosulfated α,α -trehalose core acylated with complex branched-chain fatty acids.¹⁹ Mycobacterial sulfatides elicit activities of neutrophils⁶³⁵ (V.C.3), although the possibility to be the major determinant of virulence in tuberculosis seems unlikely.¹⁴⁹

A nonphotosynthetic marine diatom *Nitzschia alba*, a eukaryote, contains sulfonoceramides (d18:1) and HSO₃-24-methylene-Chol.¹² Caprine (1-deoxy-15-methylhexadecaphinganine-1-sulfonic acid) and its acylated derivatives are the major components of the cell envelope of gliding bacteria.¹⁴⁵ The plant lacks sulfatides. Instead, photosynthetic microorganisms including cyanobacteria, algae, photosynthetic diatoms, and higher plants contain a sulfonoglycolipid (SQ-A₂Gro) accounting for as much as 40% of the total lipids.²⁷⁴ In eukaryotic photosynthetic organelles SQ-A₂Gro is located at the thylakoid membrane of chloroplasts,^{155,159} predominantly in the inner leaflet of the bilayer.

SQ-A₂Gro was localized in the photosystem II, suggesting that SQ-A₂Gro may be responsible for PSII activity by associating with the core and light-harvesting complexes of PSII.⁴⁹⁷

The SQ-A₂Gro deficient mutants of a photosynthetic purple bacterium *Rhodobacter sphaeroides*, and the cyanobacterium *Synechococcus* sp. did not have apparent phenotype including the photosynthetic electron transport system.¹⁵⁵ The growth of the wild type of these organisms under phosphate limitation resulted in increased amounts of SQ-A₂Gro with concomitant decrease of phosphoglycerolipids.³¹ By contrast, the SQ-A₂Gro-deficient mutant maintained a normal level of phosphatidylglycerol.¹⁵⁵ These authors concluded that SQ-A₂Gro plays no significant role in photoheterotrophic growth or photosynthetic electron transport in *R. sphaeroides* but may function as a surrogate for phospholipids, particularly phosphatidylglycerol, under phosphate-limiting conditions. Analogous examples of replacement of phosphate of lipids and polysaccharides with carboxylic acid residues have been reported in eubacteria.²³⁸ In contrast, another mutant of photosynthetic microorganism lacking in SQ-A₂Gro, with concomitant increase of phosphatidylglycerol showed alterations in photosynthetic activity.⁴⁹⁷

B. Invertebrates, Fishes, Amphibia, Reptiles and Birds

The protostome lineage, including Molluscs and Arthropods, lacks sulfatides²³⁸ and sialic acids,⁵⁰¹ instead contains lipids with other anionic or zwitterionic groups including uronic acid, aminoethylphosphonate,²¹² and phosphate esters of carbohydrates in insects.⁶²⁸ Sulfatides have been reported only from Deuterostomia (Echinodermata and Chordata).^{238,441} Echinoderms contain HSO₃-Chol, SM4s and/or gangliosides with sulfated *N*-glycolylneuraminic acid (NeuGc).²⁹⁶ Although the true metabolic origin of SQ-A₂Gro in the sea urchin is as yet unclear, it is interesting from a food-chain point of view that both the green alga and the sea urchin inhabit the same sea areas.²⁸⁷

Eptaretus burgeri (a cyclostome) is one of the most primitive vertebrates (the Chordate branch of Deuterostomia), and its myelin does not contain detectable amounts of glycolipids.⁵⁷⁵ The myelin structures are first apparent in cartilagenous fish species.^{441,575,578} Elasmobranchs, sharks and rays, may stand closer to higher vertebrates than to the Teleosts with their high concentrations of brain glycolipids including SM4s.^{311,441} The nerve membranes of Gadoid fishes (Alaskan pollack and Pacific cod) contained SM4g (seminolipid).^{228,576,577} In embryos of sweet water fish, medaka (teleost), SM4s was detected throughout the skin and alimentary canal by staining with a monoclonal antibody.¹¹³ SM3 was present in fish and avian testes³⁴² and was the major glycolipid of salmon eggs,³⁴⁷ whereas SM4s was detected in the testes of puffer (Pisces),⁶⁰⁶ turtle (Reptilia), and bullfrog (Amphibia).³⁴² Osmoregulatory organs of vertebrates including the gills of eel,⁶⁵⁴ the rectal gland of shark, and the salt gland of sea gull and duck²⁶⁷ contain SM4s (cf. V.C.1). Adaptation of eider duck to saline increased the concentration of SM4s in the salt gland dramatically suggesting a role for SM4s in NaCl excretion.²⁷⁰ On the other hand, the turnover of SM4s of the gills of eels adapted to seawater for six weeks was greatly enhanced, although the tissue concentration was not significantly different from that of eels adapted to fresh water.⁶⁵⁴ The skin and gills of tadpoles from a Chilean frog contained appreciable concentrations of SM4s that increased with differentiation.¹⁴⁷ SM4s also occupied about 87% of the total acidic glycolipids of *Xenopus laevis* oocytes.⁶⁶⁴ In oocytes and embryos of *X. laevis*, SM4s was distributed in the cytoplasm of vegetal hemisphere, and in neurula stage, in endoderm.³¹⁶ The concentration of SM4s in brain of *X. laevis*⁴⁴² and bullfrog tadpole⁵⁷⁸ changed quantitatively during their metamorphic stages.

The concentration of SM4s in chicken retina increased with age whereas SM4g decreased similarly to rat brain resulting in the contents of SM4g in sulfatides of 19.6%

in 18-day-embryo and 6.7% in adult (> 70 days after hatching).⁹⁵ The highest activity of GalCer sulfotransferase and arylsulfatase A in the brain and retina of chicken was observed just before hatching.⁹⁵ SM4s was the major glycosphingolipid of mature duck testes.³⁴²

C. Mammals

Mammalian epithelial tissues contain varieties of sulfatides, whose structure and concentration can be determined genetically and developmentally. Sulfatides are expressed at the outer leaflet of the plasma membrane of the glandular epithelial cells.³⁶¹

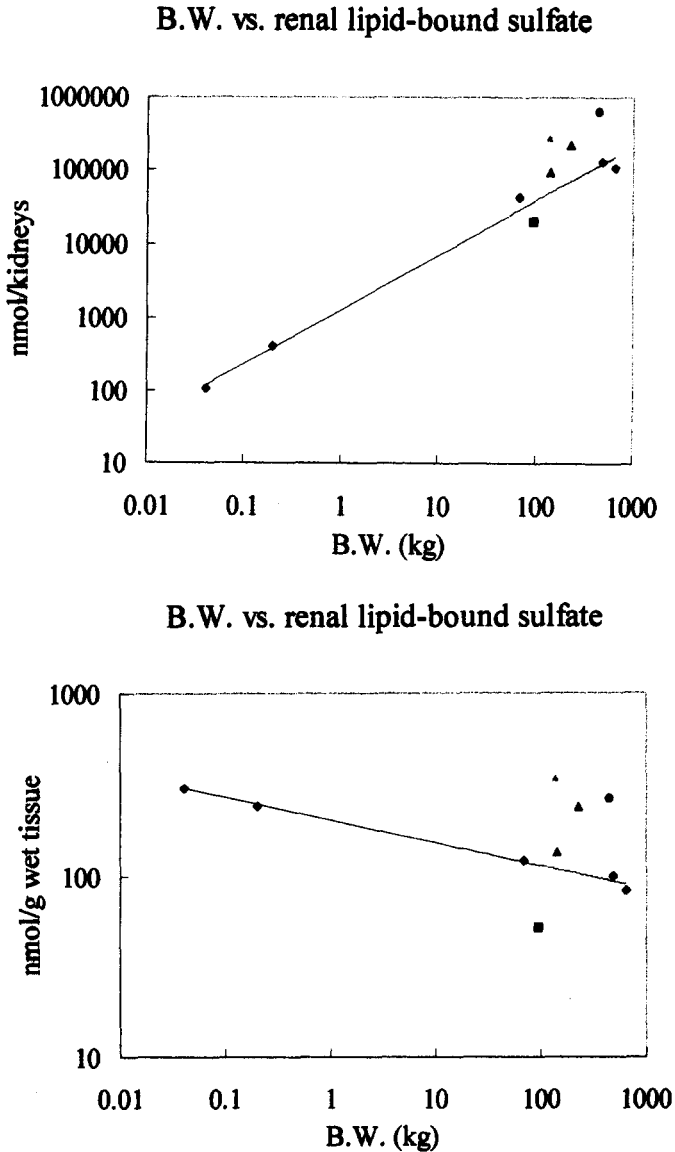


Fig. 6. Allometric relation between the renal concentration of the sulfatide sulfate and the body mass of mammals. ◆, From left to right, shrew, rat, human, horse and cow respectively; ▲, marine mammals; ■, porcine. The total sulfatide sulfate per animal (A) and the tissue concentration of the sulfatide sulfate (B) are plotted against body weight.

1. Kidney and Osmoregulatory Organs

(a) Kidney

The kidney of most of the mammals including rat, human, and rabbit contained significant amounts of HSO₃-Chol, and all epithelial cell lines derived from mammalian renal tubules incorporated [³⁵S]sulfate into HSO₃-Chol.²³³ The kidney contained the highest concentration of HSO₃-Chol among the visceral organs of adult rats^{217,243} and humans.³⁸⁸ The developmental pattern of HSO₃-Chol in the kidney was quite different from that in the brain and liver. The level of kidney HSO₃-Chol increased steadily from 6.5 nmol/g in 7-day-old rats, reached the adult level of approx. 86 nmol/g in 50-day-olds, and then stayed at the same level.²⁴³ Incorporation of [³⁵S]sulfate into SM4s of mouse kidney peaked between 35 and 42 days of age.^{171,233} In contrast, SM4s of mouse kidneys increased sharply after 2 weeks.⁹³ The major mammalian renal sulfatides are SM4s, whose concentrations are only high in the kidney of mammals next to the nervous tissue.³⁰² In 1982, the present author²³³ began to use symbols SM4s, SB1a, etc. to underline the similarity of the oligosaccharide portion of vertebrate SM4s and the ganglio-series 'sulfatides' enriched in rat kidney to that of GM4 and the ganglio-series gangliosides respectively. The human kidney contains SM3 and sulfatides of globo-series,⁴⁰⁰ while the ganglio-, globo- and isoglobo-series sulfatides are found in the rat kidney.⁵⁶⁵

The concentration (nmol/g) of acidic groups of glycolipids in the kidney of mesic mammals is regulated by two principles. The first rule dictates that the amount of glycolipid-bound sulfate (S in μ mol) of mesic terrestrial mammals is correlated with the body mass (M in kg) in the allometric equation $S = aM^b$ where a and b are constants (Fig. 6A). The total glycolipid sulfate (S , μ mol), total lipid-bound sialic acid (N , μ mol), and the total glycolipid anions (the sum of total glycolipid sulfate and ganglioside sialic acid, A , μ mol) per animal were $S = 1.29 M^{0.737}$ (or $\log S = 0.111 + 0.737 \log M$, correlation coefficient $r^2 = 0.992$), $N = 0.501 M^{0.846}$ ($r^2 = 0.988$) and $A = 1.85 M^{0.780}$ ($r^2 = 0.998$) respectively.³⁹⁸ The total amount of the renal lipid-bound sialic acid above (the exponent 0.846) was directly correlated to the kidney mass because the exponents for the kidney mass and total glomerular volume are about 0.85.⁵⁸ In contrast, the exponent for the total glycolipid sulfate was approx. 3/4 being close to those of the biochemical parameters,⁵⁹ e.g. the basal metabolic rate (0.76), nitrogen output (0.735); morphological parameters, e.g. the total proximal tubular volume (0.73); as well as the physiological parameters, e.g. the urine production rate (0.75), renal blood flow (RBF, 0.77), and the glomerular filtration rate (GFR, 0.72).

Biometrical handling of metabolic parameters in relation to body mass of animals has been called the allometric law and successfully applied to mammals of various body masses, ranging from a shrew of 2 g to a whale of 30 tons.⁵⁸ The above allometric equations can be better represented in μ M tissue concentrations of the sulfatide sulfate and the total anionic groups of mammals vs. body mass to cancel the contribution of kidney mass as follows: $S = 0.20 M^{-0.124}$ ($r^2 = 0.990$)⁴⁰³ (Fig. 6B). The GalCer sulfo-transferase activities in mice (182–550 pmol/hr/mg protein)^{93,541} were higher than human activities (42.0 pmol/hr/mg protein),⁴⁸⁹ substantiating the higher sulfatide expressions in smaller mammals. These equations were able to correlate the tissue concentration of glycolipids with other biological parameters of the animal for the first time.

The second rule predicts that the proportion of sulfatide sulfates in total anionic groups is progressively lower with increasing body mass. This rule can be derived from the equation of the first rule. The kidney of bovine (650 kg body mass) contained the glycolipid sulfate corresponding to approximately 1/3 of the total anionic groups³⁹⁸ with the remainder consisting of ganglioside sialic acids. The horse (500 kg), and human (70 kg) contained progressively higher proportions of renal sulfatides corresponding to 2/3, and 3/4 respectively, of total glycolipid acidic groups. Finally, among the smallest mammal examined, Japanese house musk shrew (body mass 0.04 kg), sulfatides occupied about 4/5 (82 mol%) of the renal glycolipid anion, setting the record ever observed in mammalian kidney. This may indicate that sulfate esters function better than sialic acids

as the ion barriers in animals with smaller body mass, which have higher metabolic activities and hence faster urine production per their body mass.⁵⁸ In a 'ouabain-resistant' mutant clone of MDCK, the sulfate/sialic acid ratio increased to 61% in comparison to 16% in the wild type MDCK⁴²³ probably compensating the lowered function of the sodium pump.

The concentration of SM4s was higher in the renal medulla of bovine,²⁶⁵ human,⁴⁹¹ and rabbit,⁶⁴⁹ forming a concentration gradient from the cortex to medulla. GalCer and Gal-A₂Gro-sulfotransferase activities were also higher in tubules and medulla.¹⁹⁵ In porcine kidney medulla of male, castrated male and female, the total sulfatide sulfate concentrations (ganglioside sialic acid in parentheses) were 194.0 (51.8), 184.3 (59.4), and 155.9 (49.4) nmol/g respectively, which were 6.1–6.7 times higher than the values for cortex, 32.0 (54.9), 27.5 (59.1), and 24.5 (59.1) nmol/g respectively.³⁷¹ The proportion of renal medulla in the whole kidney, represented as the relative medullary thickness (RMT), also increases with decreasing body mass. RMT is an expression of the proportion of the kidney that contains the loop of Henle and collecting ducts, the structure primarily responsible for concentrating urine, to overall kidney size.⁵⁹ RMT has a small inverse exponent ($M^{-0.108}$)³⁴ almost identical to that of the glycolipid sulfate, in whole kidney supporting the significant role of sulfatides in urine concentration.

The sulfatide sulfate occupied 59.7 and 65.3% respectively, of total glycolipid acidic groups in striped dolphin (body mass 140 kg) and sea lion (450 kg).³⁹⁷ These proportions are higher than those expected from their body masses. In addition, the calculated glycolipid sulfate concentrations in the kidney of terrestrial mesic mammals whose body masses were comparable to those of striped dolphin and sea lion respectively, were 110.3, and 94 nmol/g, whereas the values actually obtained were 133⁴⁰³ and 261 nmol/g,³⁹⁷ i.e. more than twice the level calculated for mesic terrestrial mammals. These results indicated that the amount of renal glycolipid sulfate of marine mammals, living solely on metabolic water, was higher than that of mesic mammals. Analogously, the incorporation of [³⁵S]sulfate into SM4s markedly increased in the compensatingly hypertrophied kidney of C₃H/He mice, after the unilateral resection of the kidney.⁶⁰⁸

A terrestrial mesic mammal of the size of porcine (95 kg) should contain 115.5 nmol/g of renal lipid-bound sulfate³⁷¹ (Toida, T., Matsumoto, H. and Ishizuka, I. unpublished) as calculated from the allometric equation. The concentrations (in parentheses % in total acidic lipids) actually obtained from male, castrated male and female porcine kidney were 52.1 (48.9%), 47.5 (46.4%) and 40.7 nmol/g (41.3%) respectively. Obviously, these lipid-bound sulfate concentrations were less than the half of the value predicted for mesic mammals, and the molar percentage in acidic lipids lower than that of a horse with the body mass several times larger than the porcine. These results suggested that the low concentration of renal acidic amphiphiles might be closely related to the behavior of pigs (and possibly in beavers) adapted to a water-rich environment where it is not necessary to concentrate urine to preserve water. The porcine has evolved from wild boar (*Artiodactyla*), and is closely related to hippopotamus of the same subclass suiform, the habitat for both being a place with a plentiful water supply.

The immunofluorescence observation using a polyclonal antibody against SM4s suggested that SM4s in the rat kidney distributed predominantly at luminal (apical) membrane of the thick ascending limb in contrast to Na⁺,K⁺-ATPase that was known to locate at basolateral membrane.⁶⁴⁹ MDCK cells, assumed to originate from collecting tubules based on hormone sensitivities,²³⁵ contained high concentrations of SM4s in contradiction to the negative staining of rabbit collecting tubules for SM4s.⁶⁴⁹ Histological studies using a specific monoclonal antibody,⁵⁶ and an L-selectin analogue⁵⁸¹ supported the tubular localization of sulfatides. Injection of a monoclonal antibody against sulfatides, Sulph I, into mice and rats showed homing to kidney tubules.⁵⁷ Kidney sections of rats, mice, pigs and monkeys were also intensely stained at the wall of the juxtaglomerular arterioles, and at the macula densa area, with Sulph I.⁵⁶ Rat or rabbit sera of Masugi's or Heymann's nephritis produced by immunization with proximal tubular

homogenate contained two populations of antibodies, specific to SB2 and SM2a respectively.²⁴⁶

In mesic species, the exponents for relative medullary thickness (RMT) ($M^{-0.108}$),^{34, 58, 59} the maximum urine concentrating ability ($M^{-0.097}$)³⁴ and the metabolic rate for kidney cortex slices ($M^{-0.13}$)⁷⁹ are close to the allometric slope, $M^{-0.124}$, of the renal sulfatide sulfate. Thus the kidney of smaller animals filters at a higher rate per gram, in other words, it produces urine at a higher rate per gram.⁵⁸ Electron microscopy has shown that smaller animals has more extensive basolateral infoldings.⁵ Basolateral membrane area per unit medullary thick ascending limb (mTAL) cellular volume ($M^{-0.075}$), as well as the volume of mitochondria as a percent of mTAL cellular volume ($M^{-0.056}$) increased with decreasing body mass. Thus, not only do mitochondria occupy more volume of mTAL cells, but those mitochondria are also more densely packed with cristae of smaller mammals.⁵ Inner mitochondrial membrane area per unit volume of mTAL cell cytoplasm scaled as $M^{-0.092}$, in reasonable agreement with the concentration of sulfatide sulfate (-0.124). The exponents for the filtration performed per gram of kidney, and the potassium uptake rate⁷⁹ were similar ($1.2 M^{-0.13}$, and $0.73 M^{-0.13}$), confirming the stoichiometry between the glycolipid sulfate and the activities of Na^+, K^+ -ATPase.^{270, 514}

(b) Cultured renal cells

Well-differentiated epithelial cell lines derived from various segments of the renal tubule, MDCK (canine), JTC-12 (cynomolgous monkey), LLC-PK₁ (porcine), and MDBK (bovine), contained sulfatides.^{235, 423, 555} SM3, and SM2a were often observed at high levels in the cultured cell lines, although not detectable or detected at quite low levels in the original tissues.²³³ The profile of glycosphingolipids of MDCK and Verots cells in culture with anisotonic media indicated that a hyposmotic medium reduced the concentrations of SM4s, SM3,⁴²⁴ and SB1a.⁴²⁵ On the contrary, the concentrations of sulfatides increased by maintaining the culture in hyperosmotic media, while the contents of most of the neutral glycolipids decreased. The hyperosmotic medium supplemented with nonelectrolytes, mannitol, sucrose or urea, also elevated the concentration of sulfatides.

The incorporation of radioactive sulfate into sulfatides⁴²⁴ and the activity of GalCer sulfotransferase²³¹ were elevated in cells of renal tubular origin, JTC-12, LLC-PK₁, MDBK,⁴²⁴ and Verots,⁴²⁵ adapted to high NaCl or mannitol. The *in vitro* stimulation of sulfotransferase at NaCl concentrations much higher than the intracellular environment⁵⁵⁵ may serve as an explanation for the stimulation of sulfatide synthesis in hyperosmolar media.²³¹ It has been established that the amount and turnover rate of sulfatides increases in adaptation to hyperosmolality by the intrinsic cellular mechanisms independent of the integral regulatory mechanisms including hormones, and the elevation of the synthesis is primarily responsible to the local barrier requirements of individual cell.⁴²⁶ Two clones (osmR-A and osmR-B), resistant to hyperosmotic media of 700 and 800 mosM/l respectively, were selected from MDCK cells. Even when cultured in isosmotic medium (300 mosM/l), the concentration of SM4s and SM3 in these hyperosmosis-resistant clones was 3.4–5.9 times higher than that in the wild-type MDCK. The rate of incorporation of [³⁵S]sulfate into sulfatides of osmR-A and osmR-B was also 1.9–6.7 times higher than that of MDCK.⁴²⁶

(c) Other osmoregulatory organs

Both HSO₃-Chol and SM4s were highly concentrated in the tracheal epithelia of rabbit.⁴⁶⁴ HSO₃-Chol,²⁴³ and SM4s⁶⁴⁵ were also contained in the stratum corneum of the skin, which functions primarily as a barrier against transepidermal water loss^{338, 661} and protects internal organs from desiccation. HSO₃-Chol, GlcCer and *N*-(*O*-linoleoyl)- ω -hydroxy fatty acyl GlcSph constituted comparable concentrations in the skin and epidermis of mammals and increased concurrently during the development.²⁵¹ Rabbit tracheal epithelial cells in primary culture undergo the terminal differentiation at confluence, with

concomitant accumulation of HSO₃-Chol by the cells, to yield cornified cells much in analogy to epidermal keratinocytes. Squamous differentiated tracheal cells also exhibited 20- to 30-fold higher levels of cholesterol sulfotransferase activity than those in undifferentiated cells did.⁴⁶⁴ Cholesterol sulfotransferase was concentrated in the basal and spinous layers of the skin, whereas arylsulfatase C, which hydrolyzes HSO₃-Chol, was rich in the stratum corneum and the granular layer.³⁸⁴ Guinea pig oral and nasal mucous membranes, which function as the first protective barrier against inhaled air, contained even higher concentrations of HSO₃-Chol than those in tracheal mucosa with higher cholesterol sulfotransferase and lower cholesterol sulfatase activities.¹⁹⁹

Aqueous humor is actively secreted from the basal surface of the epithelial cells of the ocular ciliary bodies, which participate in ion transport and osmoregulation. Electron microscopic autoradiography of the ciliary body epithelium (earlier these authors used 'ciliary processes') of rat eyes showed the synthesis of SM4s.³² In addition, an anti-SM4s monoclonal antibody stained the scattered spot-like structures in the choroid layer of murine and human eye,⁵⁶ especially pericytes, which play a role in the development of diabetic retinopathy.⁵⁵

2. Nervous System

HSO₃-Chol,²⁴³ SM4s,³⁰² SM4g,^{238,395} SMUnLc₄Cer, and SMUnLc₆Cer²⁴⁹ are the major sulfoamphiphiles of the mammalian nervous system. SM4s in the brain increased throughout the major portion of the life of man and some animals parallel to the total polar lipid of various regions.⁴⁸⁰

(a) Central nervous system

The age-dependent change of the HSO₃-Chol concentration in rat brain approximately paralleled the peak of myelination, although the subcellular distribution revealed that HSO₃-Chol was enriched in the nerve ending fraction (synaptic junctions).²⁴³ However, intracranial¹¹¹ or intraperitoneal²²⁸ administration of [³⁵S]O₄ did not label HSO₃-Chol of rat brain.¹¹¹ Fractionation of rat brain cells indicated that SM4s was concentrated in the myelin and oligodendrocytes, the cell producing myelin^{461,618} but not in neurons and astroglia.^{3,122} It has been histologically confirmed by using monoclonal antibodies, that SM4s is distributed in myelin⁵⁴³ and oligodendrocytes^{122,309} in rat brain. Both GalCer and SM4s were not expressed on astrocytes,^{122,309} whereas most of the astrocytoma tissues contained both GalCer and SM4s.²⁴⁵ Cell lines derived from astrocytoma and neuroblastoma contained only negligible amounts of SM4s⁸⁵ and did not incorporate [³⁵S] into SM4s. On the other hand, the frequency of staining for SM4s with the monoclonal antibody O4 indicated that low grade and anaplastic astrocytomas, classified on histological grounds as astrocytic, are often stained with antibodies that recognize oligodendrocytes and their progenitors.³⁴⁶ SM4s constitutes approx. 1/5 by weight of the glycolipids and 6 mol% of total lipids in the myelin membrane of the adult mammalian central nervous system and has a very slow turnover.^{54,248} The concentrations of SM4s were high in the order of brain stem > diencephalon > cerebellum > cerebrum, depending on the amount of myelin-rich nerve fibers.⁴³¹ GalCer and SM4s may be the indispensable components of myelin because mice lacking in GalCer and SM4s exhibit severe generalized tremoring after 16 days of age due to reduced action potentials.⁷⁵

The specific radioactivity of the lipid [³⁵S] in myelin-enriched fraction increased to about 15-fold²²⁶ of the activity in the homogenate of whole brain 24 hr after the intraperitoneal injection of inorganic [³⁵S]sulfate to 17-day old rats. The relative specific activity of sulfatides in myelin fraction was about 50, whereas those of other fractions, i.e. nuclei, synaptosomes, mitochondria and cytosol, were less than 1.0 indicating that SM4g was most enriched in the myelin fraction. Although SM4g has been thought to be a minor component of adult mammalian brain, it occupies approx. 42 mol% of the total cerebellar sulfatides in 14-day-old mouse⁵² and 16% of total sulfoglycolipids

of rat brain at the peak of myelination (18–21 days)^{228, 238, 395, 452, 525} but only 8.4% in adult (3–4 months) rats.⁴⁵² In the culture of embryonal mouse brain cells, the incorporation of [³⁵S]sulfate into SM4g was 17–21% of total sulfoglycolipids.³⁶ The *Jimpy*,⁴⁵² and *qk/qk* mice,³⁰⁶ deficient in myelin formation, showed reduced brain sulfotransferase activities, although the level of SM4g in testes was normal indicating that this mutation does not affect sulfatide levels in other organs.³⁰⁶ The SM4s concentration of cerebrospinal fluid (CSF)⁸⁴ in patients with vascular dementia (VAD)¹²⁴ and HIV¹⁴² was significantly higher than that in controls and Alzheimer disease, reflecting demyelination and damage of blood–brain barrier respectively.¹²² The values for gangliosides and SM4s of CSF in children and adults,⁸⁴ as well as in animals¹²² increased during development and aging.

The age for the most active synthesis of SM4s in mouse brain was 20 days^{54, 233} coinciding with the peak of myelination, whereas SM4g was synthesized with a peak around 14 days.⁵² The peak of these parameters in rat brain occurred a few days of age later²²⁸ and closely associated with the age of the most active myelination.^{228, 452, 542} After the age of active myelination, SM4s in rat²²⁷ and shrew⁶²⁴ continued to increase until 310 and 70 days respectively, in parallel with other typical myelin components including GalCer, sphingomyelin and phosphatidylinositol bisphosphate. The levels of the major galactolipids of rat spinal cord, GalCer and SM4s, increased linearly during the first two months after birth. At three months of age, constant levels were reached which were approx. 4-fold (GalCer) and 2.5-fold (SM4s) higher than in cerebral tissue of corresponding age.⁸⁸

SM4g and its synthetic activity were barely detectable in the brain of SD rat before 10 days of age,⁴⁵² and even in 14- to 16-day Wistar rats.²²⁸ Accumulation of SM4g, predominantly A₂Gro form, began with the onset of myelination (19 days), whereas after 22 days EAGro form was the major molecular species.²²⁸ SM4g then continued to increase in close parallel with SM4s until 31 days of age, reached a plateau at 48 days, then decreased dramatically after 68 days of age^{227, 228} while SM4s continues to increase. SM4s in myelin is one of the most metabolically stable components of mammals whereas SM4g turns over slightly more rapidly.^{52, 224, 248} The longer term experiment *in vivo* showed that A₂Gro form of SM4g in rat brain diminished more rapidly than the EAGro form.^{227, 452} Only about 1/10 (21.2 nmol/brain) of the level of SM4g at 48 days of age (total of both forms, 213 nmol/brain) remained at the age of 175 days and was not detected radiochemically and chemically at 310 days.²²⁷ SM4g (EAGro form) accumulated in rat spinal cord⁸⁸ in a similar fashion as Gal-A₂Gro but did not drop dramatically after 70 days as that in cerebrum. In primary cultures initiated from 19–21-day-old dissociated fetal SD rat brain,⁵²¹ GalCer, SM4s, and Galβ-A₂Gro were synthesized and accumulated by 8 days in culture. Thereafter the synthetic rates and levels of these glycolipids increased rapidly, reaching maximal values between 22–29 days in culture. The percentage of [³⁵S]sulfate incorporated into SM4g declined from 15% at day 2 *in vitro* to 5% at day 12.³⁰⁴

Sulfated glucuronyl neolacto-series sphingoglycolipids, SMUnLc₄Cer and SMUnLc₆Cer, were identified as the antigen of neuropathy IgM paraprotein.²²¹ SMUnLc₄Cer and SMUnLc₆Cer were localized predominantly in the postmigratory neuronal cells³⁸¹ as well as in BMEC (cultured bovine brain microvascular endothelial cells).²⁶¹ They may probably be related to external granulocytes of developing rat and the molecular layer containing dendrites of Purkinje cells in the adult rat cerebrum and cerebellum respectively.^{69, 71} Significant reduction in the content of the sulfoglucuronyl-neolacto series glycolipids⁶⁵ and *N*-acetylglucosaminyltransferase activity⁷¹ was observed in the cerebellum of the Purkinje cell abnormality mutant mice including Purkinje cell degeneration (*pcd/pcd*), *lurcher* (*Lc/+*), and *staggerer* (*sg/sg*). SMUnLc₄Cer and SMUnLc₆Cer were present during the embryonic stage of development of the forebrain³⁸¹ and in nerve growth cone membranes from fetal rat forebrain or brainstem²¹⁴ but disappeared soon after birth.

(b) *Peripheral nervous system*

Human peripheral nerve myelin from both sensory and motor nerves contained 1.1–2.5% of SM4s in the total lipid extract⁴³⁴ or 50 nmol/g.^{122, 552} Mouse monoclonal antibodies Sulf I^{55, 56} and 224–58¹⁵⁰ interacted specifically with SM4s and SM4g in the membrane of non-motor peripheral nerves including the vagal nerve and Schwann

Table 3. Distribution of sulfoamphiphiles in the biosphere

Prokaryotes and plants	
HSO ₃ -PtdGro (E ₂₀ E ₂₀ analogue), <i>Haloarcula marismortui</i> , 17 mol% of polar lipids. ¹⁰⁰ <i>Halobacterium salinarum</i> (<i>cutirubrum</i>), 4 mol% of polar lipids. ³²⁶ <i>Hb. sodomense</i> , 3% of the total lipid. ^{256, 597} <i>Hb. trapanicum</i> , 15 weight % of polar lipids. ^{256, 598}	
HSO ₃ -6Glc α -1E ₂₀ E ₂₀ Gro, chemical synthesis ²⁵⁷	
HSO ₃ -6Man α -2Glc α -1E ₂₀ E ₂₀ Gro (S-DGD-1), <i>Haloferax mediterranei</i> (strain R-4), 2.8 μ mol/g (25 mol% of polar lipids). ³²⁶ <i>Hb. saccharovororum</i> , 13% of the total lipid. ^{256, 326} <i>Halococcus saccharolyticus</i> strain, 19.5 mol% of polar lipids. ³⁸³ <i>Hb. salinarum</i> , (+). ³⁶⁸ Chemical synthesis ²⁵⁷	
HSO ₃ -2Man α -4Glc α -1E ₂₀ E ₂₀ Gro (S-DGD-3), <i>Hb. sodomense</i> , (+). ^{256, 597}	
HSO ₃ -2Man α -2Glc α -1E ₂₀ E ₂₀ Gro (S-DGD-5), <i>Hb. trapanicum</i> , 29% by weight of polar lipids. ⁵⁹⁸	
HSO ₃ -3Gal β -6Man α -2Glc α -1E ₂₀ E ₂₀ Gro (S-TGD-1), <i>Hb. salinarum</i> (<i>cutirubrum</i> or <i>halobium</i>), 24% of polar lipids. ^{256, 271, 275} 21%; ³²⁶ FAB, ¹ H-NMR, (+). ¹⁰² <i>Hb. saccharovororum</i> , (+). ^{256, 325}	
HSO ₃ -3Gal β -6(Galf α -3)Man α -2Glc α -1E ₂₀ E ₂₀ Gro (S-TeGD), <i>Hb. salinarum</i> , (+). ⁵²⁷ <i>Hb. saccharovororum</i> , (+). ^{256, 325}	
(HSO ₃) ₂ -2,6Man α -2Glc α -1E ₂₀ E ₂₀ Gro and E ₂₅ E ₂₀ Gro (S ₂ -DGD-1), halophilic bacterium <i>Natrialba asiatica</i> (former strain 172), (+). ³⁶⁹ Strain BIT, (+). ²³⁶	
2,3,6,6'-Tetraacyl-trehalose-2'-sulfate, Mycobacteria, (+). ^{149, 159}	
SQ-A ₂ Gro (17:0 branch), <i>Sulfolobus acidocaldarius</i> (glycolipid K), 9% of the total lipid or 43% of acidic lipids. ³³² SQ-A ₂ Gro [16:0, 18:0cyc, 18:1(ω 7)], freshwater bacterium <i>Caulobacter bacteroides</i> , 9% of the total lipid. ⁶⁶⁵ SQ-A ₂ Gro, anaerobic photosynthetic bacterium <i>Rhodospseudomonas viridis</i> , (-). ⁶⁶⁶ SQ-A ₂ Gro (16:0 and 18:1), photosynthetic (marine green) alga <i>Enteromorpha flexuosa</i> , 67 nmol/g (200 μ g/g dry cell). ⁵¹⁸ SQ-A ₂ Gro (18:3), green leaves of <i>Vicia faba</i> , 290 nmol/g. ²⁸⁴ The thylakoid membrane of chloroplasts, 70 μ mol/g. ²³⁸ <i>Rhodobacter spheroides</i> , cultured in 1.0 mM, Pi, 2.2 mol% of the total lipid; in 0.1 mM Pi, 16.6 mol% ^{31, 155}	
6-Sulfoquinovosyl-3',2'-O-acyl-1'-O-thioacyl-sn-Gro (thionsulfonolipid), a picoplankton cyanobacterium (<i>Synechococcus</i> sp.), 670 nmol/g (0.2% of dry cells). ²⁸¹	
Taurine-6GlcU α -3-sn-A ₂ Gro, <i>Hyphomonas jannaschiana</i> (a seawater bacterium), 34% of the total lipid or 18 μ mol/g (52 mg/g dry cell). ²⁹	
(HSO ₃) ₂ -3,4-(3-O-methyl-1-oxobutyl)-2Glc β -[15-hydroxy-2-oxy]-19-norkaur-16-en-18oic acid (atractyloside), <i>Atractylis gummifera</i> , (+). ⁵⁹⁰	
(HSO ₃) ₂ -1,12-docosanediol, a phytoflagellate <i>Ochromonas danica</i> , (+). ¹⁵⁹	
HSO ₃ -1Cer (d18:1/24:0), chemical synthesis ⁴³⁰	
HSO ₂ -1Cer (d18:1/16:1- Δ^3 -trans), a nonphotosynthetic diatom <i>Nitzschia alba</i> , (+). ¹²	
HSO ₃ -3(24-methylene)Chol, <i>Nitzschia alba</i> , (+). ¹²	
Invertebrates, fishes, amphibia, reptiles and birds	
HSO ₃ -Chol, sea star <i>Asterias rubens</i> , 560 nmol/g (1.3 mg/g dry tissue). ³⁹ Sea urchin <i>Anthocidaris crassispina</i> , 2.2 μ mol/g (1.04 mg/g, 15% of polar lipids). ⁶⁴⁷ Chemical synthesis. ^{607, 647}	
SQ-A ₂ Gro, sea urchin <i>Pseudocentrotus depressus</i> , eggs, 300 nmol/g; spermatozoa, 700 nmol/g. ²³⁹ SQ-A ₂ Gro (16:0), sea urchin <i>Anthocidaris crassispina</i> , shells, 30 nmol/g. ²⁸⁷	
SM4s, orange star fish, radial nerve, trace; SM4s (2-hydroxy acid), amphioxus, ventral nerve, 0.8 nmol/g. ⁴⁴¹	
HSO ₃ -8NeuGc α -2-6GlcCer (t18:0/15:0 and t18:0/22h:0), sea urchin gonads, (+). ²⁹⁷ Sea urchin eggs (t18:0/24h:0), 15 nmol/g (94 μ g/g dry eggs). ³¹⁵	
HSO ₃ -8NeuGc α -2-6Glc-8NeuGc α -2-6GlcCer, sea urchin eggs, (+). ⁴⁵⁷	
HSO ₃ -8NeuAcx2(-8NeuAcx2) _n -6GlcCer (t18:0/20:1,21:1,22:1) ($n = 0, 1, 2$, and 3), male sea urchin gonads, < 38 nmol/g. ²¹⁸	
SM4s (d18:1/24:1 and d18:1/24h:1), a coelacanth <i>Latimeria chalumnae</i> , brain, 3 μ mol/g. ⁵⁷⁹	
SM4s (d18:1, t18:0/22h:0, 24 h:1), spiny dogfish, salt (rectal) gland, 640 nmol/g (2.8 mg/g dry tissue), 470 nmol/g (determined by hexose analysis on the pure lipid fraction, 2.1 mg/g dry tissue). ²⁶⁹ SM4s (d18:1/24:1 and d18:1/24h:1), a ray <i>Torpedo marmorata</i> , electric organ, 1180 nmol/g (5.9 μ mol/g dry tissue). ¹⁷⁴ SM4s, a shark <i>C. longimanus</i> , brain, 3.1 μ mol/g (2.8 mg/g tissue). ³¹² Brown shark, brain, 3.6 μ mol/g. ⁴⁴¹ Skate fish testis, TLC (+). ³⁴²	
SM4s, eel (<i>Anguilla anguilla</i>), gills, [³⁵ S]labeling/TLC, adapted to fresh water, 45 nmol/g; seawater, 59 nmol/g. ⁶⁵⁴	
SM4s, teleost fishes, brain, 1.0–2.1 μ mol/g (0.9–1.9 mg/g). ³¹² SM4s (fatty acids are exclusively nonhydroxy), Alaskan pollack, brain, 1350 nmol/g (12 mol% of total glycolipids); spinal cord, 2660 nmol/g (13 mol%). ^{576, 577} SM4s (nonhydroxy fatty acids, 90%), common carp, brain, 370 nmol/g (20 mol%). ⁵⁷⁷ SM4s (nonhydroxy fatty acids predominate), killifish, brain, 1.6 μ mol/g; spinal cord, 1.4 μ mol/g. ⁴⁴¹ SM4s (d20:1/24:1), puffer (<i>Fugu rubripes rubripes</i>), testis, 17 nmol/g (15 μ g/g). ⁶⁰⁵ Medaka (teleost), skin and alimentary tract, TLC-OL, SM4s, (+). ¹¹³ Cod intestine, TLC (chemical staining), SM4s (-). ⁴⁷	

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Table 3—continued

SM4g, puffer, testis, (-).⁶⁰⁵ Alaskan pollack, brain, 1690 nmol/g (15 mol% of total glycolipids); spinal cord, 2580 nmol/g (13 mol%); common carp, brain, 0 nmol/g.⁵⁷⁷
 SM3, testes of rainbow trout and salmon, TLC, (+).³⁴² SM3 (d18:1/16:0), salmon milt, 426 nmol/g (2 mg/g dry weight)³⁴⁷
 SM4s (d18:1/24:1), bullfrog tadpole, metamorphosis XI–XVI, 250 nmol/g; bullfrog brain, 790 nmol/g; SM4s (d18:1/24:1), sciatic nerve, 1350 nmol/g.⁴⁸⁵ SM4s (d18:0/24h:1), adult frog brain, 800 nmol/g.⁵⁷⁸ SM4s (d18:0/24:1), axolotl brain, neonotous adult form, 560 nmol/g; metamorphosed adult form, 710 nmol/g.⁵⁷⁸ SM4s, Chilean frog (*Calyptocephalella caudiverbera*) skin, tadpole of 31 day, 210 nmol/g (2.0 mg/mg protein); 44-day tadpole, 640 nmol/g (6.2 mg/mg protein)⁶⁵¹
 SM4s, toad, convoluted oviduct, Azure A assay of the total organic phase lipid, 150 nmol/g (134 µg/g) in March, 260 nmol/g (232 µg/g) in May; uterine oviduct, 540 nmol/g (487 µg/g) in March, 750 nmol/g in May (671 µg/g).⁶¹⁵ *Xenopus*, ovaries, > 85% of the total acidic glycolipids³¹⁶
 SM4s (d20:1/22-24h:0), duck, salt gland, adapted to fresh water, 230 nmol/g; adapted to saline, 810 nmol/g.²⁶⁸ SM4s (d20:1/22-24h:0), eider duck, salt gland, 1.64 µ mol/g; herring gull, salt gland, 1.13 µ mol/g.²⁷⁰ Mature fowl testis, (+); immature duck testis, (±); mature duck testis, (++)³⁴² SM4s, adult chicken (> 70 days after hatching), retina, 24 nmol/retina or 295 nmol/g; brain, 2350 nmol/g⁹⁵
 SM4g, adult chicken, 1.8 nmol/retina⁹⁵

Mammalian

Kidney

H₂SO₄-Chol, rat (SD), hydrolysis and GC of the fraction purified on silicic acid columns, 107–132 nmol/g (249–307 µg/g dry tissue).²⁴³ Wistar rat, FeCl₃ assay of the fraction purified on DEAE-Sephadex and silicic acid columns, 89 nmol/g.²¹⁷ Rabbit, 275 nmol/g (2.75 nmol/mg protein)⁸¹
 Sheep, (+).²⁹⁹ Human, 430 nmol/g (200 µg/g); metachromatic leukodystrophy patient, 3.5 y, approx. 430 nmol/g.³⁸⁸ Human urine, 0.4 nmol/ml.⁴¹ Monkey renal cell line JTC-12 and canine kidney cell line MDCK,³⁵ S-label, (+)^{228, 233, 423}
 SM4s (d18:1/22h:0, 24 h:0), GC with mannitol as an internal standard, house musk shrew (*Suncus murinus*), 299 nmol/g (82 mol% of the total acidic glycolipid).³⁹⁸ SM4s, mouse (C57BL/6J-*cpk*, 3 weeks), densitometry after charring the TLC plate with copper sulfate/phosphoric acid, 457 nmol/g; cortex, 373 nmol/g; medulla, 935 nmol/g; cystic kidney, 34.4 nmol/g.⁹³ Prosaposin-deficient mouse, 5-fold increase¹³⁰
 SM4s (d18:1/22-24h:0), rat (Wistar), GC using mannitol as an internal standard, 182 nmol/g.⁵⁵⁹ SD rat (30–120 day), HPLC of benzoylated desulfated SM4s (d18:1/hydroxy fatty acids), 140–260 nmol/g (0.7–1.3 nmol/mg dry weight); SM4s (d18:1/nonhydroxy fatty acids), 60–160 nmol/g (0.3–0.8 nmol/mg dry tissue).²¹⁵ SM4s, 60-day-old SD rat (body weight 200 g), TLC densitometry of the purified SM4s after orcinol stain, 201 nmol/g.²¹⁷ Lewis rat, TLC-OL using the monoclonal antibody and densitometry, 278 nmol/g; rat and other mammals, distal tubules, (+).⁵⁶ Rat kidney Golgi complex fraction, 13 µ mol/g (117 µg/mg protein); plasma membrane fraction, 6.1 µ mol/g (55 µg/mg protein)⁶⁵⁰
 SM4s, rabbit glomeruli, preparative TLC, 16 nmol/g (8 nmol/100 mg dry weight); cortex, 26 nmol/g (13 nmol); cortical tubule, 54 nmol/g (27 nmol); medulla, 208 nmol/g (104 nmol);⁶⁴⁹ rabbit (6 month, New Zealand White) kidney, 96 nmol/g (0.96 nmol/mg protein)⁸¹
 SM4s, human (body weight 69 kg), DEAE-cellulose column, 81–112 nmol/g;³⁶⁶ 120 nmol/g (0.54 mg/g dry tissue).²⁶⁶ SM4s (d18:1/22:0), human, cortex, 59 nmol/g; SM4s (d18:1/24h:1), medulla, 169 nmol/g.⁴⁹¹ SM4s, human fetal (20 weeks), TLC densitometry, normal, 44 nmol/g (0.22 nmol/mg dry tissue); Krabbe disease, 78 nmol/g (0.39 nmol); Sandhoff disease, 8 nmol/g (0.04 nmol); metachromatic leukodystrophy, 52 nmol/g (0.26 nmol); prosaposin deficiency, 196 nmol/g (0.98 nmol).⁴⁶ SM4s, elevated in a patient of atypical Farber disease.¹³¹ Normal human urine, perbenzoylation and desulfation, 0.16 nmol/mg creatinine (67% in the sediment), urine of metachromatic leukodystrophy patients, 7.6 nmol/mg creatinine⁴¹⁵
 SM4s (d18:1/23h:0, 24 h:0), sheep, preparative TLC and gas chromatography, 22 nmol/g.²⁹⁹ SM4s (d18:1/23h:0, 24 h:0), horse (body weight 485 kg), 99.8 nmol/g.⁵⁹⁹ SM4s (d18:1, t18:0/23h:0, 24 h:0), (+).^{181, 570} SM4s (d18:1/24h:0), bovine (body weight 650 kg), 86 nmol/g (0.4 mg/g dry tissue); cortex, 22 nmol/g (0.1 mg/g); transition zone, 65 nmol/g (0.3 mg/g); medulla, 194 nmol/g (0.9 mg/g); large papilla, 86 nmol/g (0.4 mg/g).²⁶⁵
 SM4s, porcine (body weight 95–105 kg), male, renal cortex, 31.3 nmol/g, medulla, 193 nmol/g; castrated male, cortex, 26.0 nmol/g, medulla, 183 nmol/g; female, cortex, 23.6 nmol/g, medulla, 155 nmol/g.³⁷¹ (Toida T., Matsumoto H., and Ishizuka, I., unpublished results)
 SM4s (d18:1/22h:0), dolphin *Stenella coeruleoalba* (body weight 140 kg), 124 nmol/g.⁴⁰¹ SM4s (d18:1/22h:0), a sea lion *Eumetopias jubata* (suborder Pinnipedia, body weight 450 kg), 252 nmol/g.³⁹⁷
 SM4s, human renal cell carcinoma (Grawitz) tissue (adenocarcinoma), 1.0–72.0-fold the level in the uninvolved tissue.⁴⁹⁰ Renal cell carcinoma, 1.7-fold of the uninvolved tissue; Wilms' tumor, (-).⁴⁸⁹ SMKT-R3 human renal carcinoma cells, cytofluorometry, (+)²⁹³
 SM4s, JTC-12 cell line, 40 nmol/g (20 nmol/100 mg dry weight).^{225, 234} SM4s (d18:1/16:0), MDCK, 70 nmol/g (0.7 nmol/mg protein); MDCK, a 'ouabain-resistant' mutant, 190 nmol/g.⁴²³ SM4s + ³⁵S-label, MDCK strain I, (+); MDCK strain II, (+ + +)⁴²¹. MDCK strain I, (+); MDCK strain II, (-).¹⁷⁶ MDCK strain II, [³⁵S]sulfate and [³H]ceramide labeling, (+).⁶⁵⁷ MDCK cultured for 3–7 days in a hyposmotic medium (100 mosM/l), 46 nmol/g; cultured in a hypertonic medium (500 mosM/l), 93 nmol/g; in a medium made to 500 mosM/l by additions of mannitol, 120 nmol/g.⁴²⁴ MDCK, 23 nmol/10⁶ cells; fraction insoluble in Triton X-100, 13 nmol/10⁶ cells.⁴⁹ SM4s (d18:1/18:0), SGE1 rat renal glomerular epithelial cells, 8.0 ± 4.3 nmol/mg DNA; monolayer with domes 9.6 ± 1.2 nmol.⁵⁹⁵ EUE (human embryonic epithelium) line, Azure A assay of organic phase lipids, in the isotonic medium, 2.9 µ mol/g (28.0 µg/mg protein); 12.6 µg/g (120 mg/mg protein).⁴² Human renal cancer cell line SMKT-R3, Azure A staining, TLC-OL with monoclonal antibody Sulf I, [³⁵S]-labeling, (+)²⁹³

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Table 3—continued

SM4s-Glc (t18:0/24:0), Wistar rat kidney, TLC densitometry (orcino), 5.5 nmol/g.²¹⁷ SM4s-Glc, MDCK, DEAE-Sephadex column, (+)⁴²⁶
 SM4g, SD rat kidney, spleen, intestine, and liver, [³⁵S]-labeling, (-).³⁵⁴ Wistar rat kidney, (-)⁵⁵⁹
 SM3 (d18:1/18:0), house musk shrew, 0.5 nmol/g.³⁹⁸ SM3, prosaposin-deficient mouse, mildly elevated¹³⁰
 SM3 (t18:0/22-24:0), rat, 1.7 nmol/g;⁵⁵⁹ 1.6 nmol/g.²¹⁷ Human kidney, 30 nmol/g.^{366,535} SM3 (d18:1/24:0), human, cortex, 33 nmol/g; medulla, 11 nmol/g.⁴⁹¹ Human renal cell carcinoma (Grawitz) tissue (adenocarcinoma), 1.3–39.4-fold the level in the uninvolved tissue;⁴⁹⁰ 86.5-fold the level in the uninvolved tissue.⁴⁸⁹ Human Wilms' tumor, (-).⁴⁸⁹ SM3 (d18:1/22:0), dolphin *Stenella coeruleoalba*, 8.7 nmol/g.⁴⁰¹ SM3 (d18:1 and t18:0/22:0), a sea lion *Eumetopias jubata*, 9 nmol/g.³⁹⁷ Sheep, 3.3 nmol/g.²⁹⁹
 SM3, JTC-12, 53 nmol/g (0.53 nmol/mg protein).^{225,234} SM3 (d18:1/16:0), MDCK, 20 nmol/g; MDCK, a 'ouabain-resistant' mutant, 40 nmol/g.⁴²³ MDCK cultured for 3–7 days in a hypotonic medium (100 mosM/l), 18 nmol/g; cultured in a hypertonic medium (500 mosM/l), 54 nmol/g; in a medium made to 500 mosM/l by additions of mannitol, 50 nmol/g.⁴²⁴ MDCK strain II, [³⁵S]sulfate and [³H]ceramide labeling, (+).⁶⁵⁷ MDBK, ³⁵S-label, (+) as the major sulfate.^{233,234,424} Human renal cancer cell line SMKT-R3, Azure A staining, TLC-OL with monoclonal antibody Sul I, [³⁵S]-labeling, (+);²⁹³ Verots, [³⁵S]-labeling, (+)⁴²⁵
 SM2a (t18:0/24:0), Wistar rat, 24 nmol/g.⁵⁵⁹ SM2a, SD rat, (+).²⁴⁶ Human, (-).³⁹⁹ SM2a, JTC-12, ³⁵S-label, (+);^{230,424,555} porcine kidney cell line LLC-PK₁, ³⁵S-label, (+);⁴²⁴ Verots, ³⁵S-label, (+);⁴²⁵ human renal cancer cell line SMKT-R3, Azure A staining, [³⁵S]-labeling, (+).²⁹³
 SM2b (t18:0/24:0, 22:0), SD rat, 0.05 nmol/g.⁵⁶⁵ Human renal cancer cell line SMKT-R3, [³⁵S]PAPS-labeling, (+) (*in vitro* formation)²⁹²
 SB2 (t18:0/24:0), Wistar rat kidney, 11 nmol/g.⁵⁶⁰ SD rat, approx. 7.5 nmol/g.²⁴⁶ Human renal cancer cell line SMKT-R3, [³⁵S]PAPS-labeling, (+) (*in vitro* formation)²⁹²
 SB1a (t18:0/24:0), Wistar rat, 6 nmol/g.⁵⁶¹ SB1a, porcine, (+)³⁷¹
 SMGb₄Cer (d18:1/24:1, t18:0/24:1), human, 0.03 nmol/g.⁴⁰⁰ SD rat, 0.07 nmol/g.⁵⁶⁴
 SMiGb₄Cer (t18:0/24:0), SD rat, 0.27 nmol/g.⁵⁶⁴
 SMGb₅Cer (d18:1/24:0), human, 0.19 nmol/g.³⁹⁹
 SMiGb₅Cer (t18:0/24:0 and 22:0), SD rat, 0.11 nmol/g.⁵⁶⁸
 SMGM1a (II³-NeuGc, IV³-HSO₃-Gg₄Cer) (t18:0/24:0), SD rat, 120 pmol/g.⁵⁶⁷

Epidermis, keratinocytes, and eyes

HSO₃-Chol, guinea pig, dorsal epidermis, 790 nmol/g; dorsal dermis, 120 nmol/g.⁶⁰⁴ Murine skin, 15-day-fetus, 39 nmol/g (0.09 μg/mg dry weight); epidermis, 17-day-fetus, 211 nmol/g (0.49 μg/mg dry weight).²⁵¹ Human cohesive stratum corneum, upper arm, 1.2 ± 0.5%; palm, 1.3 ± 1.2% of the total lipid.⁵¹² X-linked ichthyosis patient, stratum corneum, elevated.⁶³⁰ Normal human, scale, 2.3% of the total lipid; X-linked ichthyosis 12.2%.⁶³⁰ Bovine hoof, (+).⁶⁰⁷ Horse hoof, 19.6% of horse hoof lipids, 10% of cow hoof lipids.⁶²⁶
 SM4s, guinea pig, dorsal epidermis, (-); dorsal dermis, 6 nmol/g.⁶⁰⁴ Human keratinocytes, Sulf-1 immunohistochemistry, (±).¹⁵⁴ Immunohistochemistry, (+) both acinar and ductal cells of the normal human mammary glands.²² H3630 breast carcinoma cell line,¹⁷ human breast ductal carcinoma cells, [³⁵S]incorporation, (+)⁵⁸³
 SM4s, rat, ciliary body epithelium, basal membrane.³⁵ S incorporation, (+).³² Choroidea of the eye, human, immunohistochemistry and TLC-OL, 30 nmol/g; Lewis rat and SD rat, 4.8 nmol/g, (+) pericytes.⁵⁵
 SM3, human breast ductal carcinoma cells, [³⁵S]incorporation, (+)⁵⁸³

Lung

HSO₃-Chol, rat (43-day-old), lung, 21.3–24.5 nmol/g (49.5–57.0 μg/g dry tissue).²⁴³ Rat lung, [³⁵S]-labeling, (+).³⁵⁴ Guinea pig, nasal mucosa, 136 nmol/g (0.68 μmol/g dry weight); oral mucosa, 194 nmol/g (0.97 μmol/g); tracheal mucosa, 26 nmol/g (0.13 μmol).¹⁹⁹ Rabbit, tracheal epithelial cells, (+)⁴⁶⁴
 SM4s, rat lung, undifferentiated small cell carcinoma, 2 nmol/g (11 μg/g dry tissue); adenocarcinoma, 8 nmol/g (36 μg/g dry tissue); human lung, 0.7 nmol/g (3 μg/g dry tissue); embryonal lung, 4 nmol/g (20 μg/g dry tissue); SM4s (d18:1/24:0) squamous cell carcinoma, 0.9 nmol/g (4 μg/g dry tissue).^{138,645}
 Immunohistochemistry, (+), human lung adenocarcinoma, 17 in 43 cases; large cell carcinoma, 5 in 25 cases.³⁷⁸ Small cell carcinoma, Azure A staining, (+)¹⁴⁴
 SM4s, porcine respiratory ciliated cells, (+)⁶⁵²

Central nervous system

HSO₃-Chol, brain, human, 4-day-old, (+);³⁸⁸ SD rat, 7.0–9.0 nmol/g (16–21 μg/g dry tissue).²⁴³
 SM4s, house musk shrew (*Suncus murinus*), 70 days, Azure A assay of the acidic glycolipids, cerebrum, 3.0 μmol/g; cerebellum, 4.2 μmol/g; bulbus olfactorius, 3.5 μmol/g.⁶²⁴
 SM4s, brain, mouse (16 days), 1017 nmol/g; *msd*, a myelin-deficient mutant, 365 nmol/g; reduced in other myelin-deficient mutants, *jp* and *qk*.³⁷⁵ Cerebellum, normal mouse, 1.9 μmol/g; *Lc/+* mutant mouse (Purkinje cell degeneration), 3.9 μmol/g; *wv/wv* mouse (Purkinje cell defect), 2.8 μmol/g; *sg/sg* mouse, 3.2 μmol/g; control of *rl/rl*, 2.9 μmol/g; *rl/rl* mutant, 3.5 μmol/g; *qk/qk* myelin-deficient mutant, 0.34 μmol/g.⁶⁶ Mice lacking in UDP-Gal: Cer galactosyltransferase, (-).⁷⁵ Mouse (32 days), cerebellum, 6 μmol/g.⁵⁴

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Table 3—continued

- SM4s, SD rat (22 day), brain, 2.0 $\mu\text{mol/g}$; 180 day, 4.8 $\mu\text{mol/g}$;²⁴ 120 day brain, SM4s (hydroxy fatty acid), 1.9 $\mu\text{mol/g}$; SM4s (nonhydroxy fatty acid), 1.9 $\mu\text{mol/g}$.⁵¹⁵ SM4s (d18:1/18:0, 24 h:0), SD rat, 15–20 days, silicic acid column and preparative TLC, myelin, 6.9 $\mu\text{mol/g}$ (4.4% of dry weight); astroglia, 0.2 $\mu\text{mol/g}$ (0.2%); neurons, 0.05 $\mu\text{mol/g}$ (0.1%).³ SM4s, Wistar rat, 84-Day-old, cerebrum, 4 $\mu\text{mol/g}$; 90-day-old, spinal cord, 11 $\mu\text{mol/g}$; 175 day, brain, 3.7 $\mu\text{mol/g}$; 310 day 4.9 $\mu\text{mol/g}$.²²⁷ Rat (14 day), cerebrum, SM4s (nonhydroxy fatty acids), 120 nmol/g; SM4s (hydroxy fatty acids) 63 nmol/g; cerebellum, SM4s (nonhydroxy fatty acids), 244 nmol/g; SM4s (hydroxy fatty acids); 112 nmol/g; brain stem, SM4s (nonhydroxy fatty acids), 724 nmol/g; SM4s (hydroxy fatty acids), 285 nmol/g.⁴³¹ SM4s + SM4g, Wistar rat, 6.8 $\mu\text{mol/g}$; chronically diazepam treated, 1.5 $\mu\text{mol/g}$.⁶²¹ SM4s + SM4g, 8-months-old SD rat, cerebellum, control, 1.08 $\mu\text{mol/cerebellum}$; alcohol-fed, 0.25 $\mu\text{mol/cerebellum}$.⁶²⁰ 8-months-old Wistar rat, brain, 6.8–6.9 $\mu\text{mol/g}$; fed ethanol and protein-deficient diet for 6 months, 5.3 $\mu\text{mol/g}$; 3 months after switching to a high protein diet, 8.4 $\mu\text{mol/g}$.⁶¹⁹
- SM4s, human brain, frontal lobe, white matter, 12.0 $\mu\text{mol/g}$; frontal lobe, gray matter, 2.2 $\mu\text{mol/g}$.⁴²⁰ Frontal lobe (1.7–11 y), 2.2–6.0 $\mu\text{mol/g}$; frontal lobe, metachromatic leukodystrophy, 8.2–13.3 $\mu\text{mol/g}$.⁴⁷⁷ Human brain, white matter, 8.2 $\mu\text{mol/g}$.⁴⁴¹ White matter, HPLC using a Nucleosil column, control, 5.1 weight%; alcoholics, 4.6%.⁴⁴⁵ SM4s (d18:1/24:1, 24 h:0), bovine brain, (+).⁴³⁰ SM4s, bear, cerebrum, 6.9 $\mu\text{mol/g}$; cerebellum, 5.3 $\mu\text{mol/g}$.⁴⁹⁹
- SM4s, mouse gliomas, 300 nmol/10⁹ cells.⁸⁵ Various neural tumors of rat brain, 0–8.4% of lipid weight.⁷² C6 glioma cell treated with disipramine, (+).⁵⁹² TLC-OL, (+), glioblastoma of grade IV; (–), glioblastoma multiforme, astrocytoma grade II, anaplastic astrocytoma.²⁴⁴ Oligodendrocytes, [³⁵S]-labeling, (+).^{27, 111} Human oligodendrogloma, immunohistochemistry, (+).⁵⁴³ SM4s [¹⁴C-stearic acid], chemical preparation⁹⁶
- SM4g (A₂Gro + EAGro forms), mouse cerebellum, 28-day-old, 900 nmol/g; 14 days, 42% of the total cerebellar sulfatides.⁵² SM4g (A₂Gro form), rat brain, [³⁵S]-labeling, (+).¹²⁰ SM4g (A₂Gro + EAGro forms), mouse (23-day-old), brain, 490 nmol/g; non-myelinating jimpy mouse, 69 nmol/g; Wistar rat (body weight 300 g), brain, 190 nmol/g.³⁴³ SD rat brain, before 10 days of age, (–); between 10–25 days, 0.3–0.4 $\mu\text{mol/brain}$; rat large myelin fraction 13.6 $\mu\text{mol/g}$ (136 nmol/mg protein).⁴⁵² SM4g, Wistar rat, brain, 48 days, A₂Gro form, 50 nmol/brain; EAGro form, 180 nmol/brain; A₂Gro + EAGro forms, 210 nmol/brain; 175 days (EAGro form), 20 nmol/brain; 175 days (A₂Gro form), 1.5 nmol/brain (total 21 nmol); 310 days, SM4g (A₂Gro + EAGro forms), not detected.^{227, 228} SM4g (EAGro form), 84-day-old Wistar rat, spinal cord, 600 nmol/g.⁸⁸ Wistar rat (17-day-old), central nervous system myelin, [³⁵S]sulfate incorporation, 16% of the total sulfatides.²²⁶
- SM4g, rabbit brain, approx. 50 nmol/g; human brain frontal lobe, (–).³⁴³ Adult human brain, (–); brain, metachromatic leukodystrophy patient, (–).²²⁸
- Lyso-SM4s, normal human frontal lobe, 50–180 nmol/g; frontal lobe, metachromatic leukodystrophy, 9–45 nmol/g.⁴⁷⁷ Lyso-SM4s (d18:1), chemical preparation^{125, 569, 616}
- SM4s-6(d18:1/24:0) (GalCer I⁶-sulfate), chemical synthesis^{364, 430, 572}
- SMUnLc₄Cer + SMUnLc₆Cer, mouse cerebellum, 400–600 nmol/g (4–6 ng/mg protein).⁶⁶ Cerebellum, murine mutants *lurcher* (*Lc/+*) and (*pcd/pcd*) (Purkinje cell degeneration), staggerer (*sg/sg*) (Purkinje cell defect), (–); *nr/nr* mutant, approx. 70% reduction⁶⁵
- SMUnLc₄Cer, rat (embryonic day 19), cerebral cortex, 1.4 nmol/g (11.2 $\mu\text{g/g}$ dry tissue).⁶⁷
- SMUnLc₄Cer + SMUnLc₆Cer, rat forebrain, embryonic stage, (+); soon after birth, (–).³⁸¹ Immunohistochemistry, growth cone of embryonal rat brain, neuronal cell bodies.⁴⁴ SMUnLc₄Cer (d18:1/16:0, 18:0 and 18:1), rat cerebellum, (+).⁶⁷ SMUnLc₄Cer, rat cerebral cortex, microvessel fraction, 4.8–6.6 nmol/g (76–106 ng/mg protein); cultured endothelial cells, 3.2 nmol/g (51 ng/mg protein).³⁷⁹ BMEC (cultured bovine brain microvascular endothelial cells), 7-day-culture, 4.1 nmol/g (65 ng/mg protein); 14-day-culture, 1.3 nmol/g (21 ng/mg protein).²⁶¹
- SMUnLc₄Cer, human: fetal (37 week) brain, 0.94 nmol/g (7.5 $\mu\text{g/g}$ dry tissue); cortex (adult), <0.1 $\mu\text{g/g}$ dry tissue; cerebellum, adult, 1.6 nmol/g (12.5 $\mu\text{g/g}$ dry tissue).⁶⁷
- Peripheral nervous tissue, Schwann cell and cerebrospinal fluid*
- H₃O₃-Chol, rat (43-day-old), adrenals, 22–30 nmol/g (51–69 $\mu\text{g/g}$ dry weight).²⁴³ Bovine adrenals, 3.2 nmol/g (1.5 $\mu\text{g/g}$); human adrenal tumor (Cushing's syndrome), 86–430 nmol/g (40–200 $\mu\text{g/g}$).⁹⁴ Schwann cell line D6P2T, (–).²⁶ G361 human melanoma cells, trace⁴⁶⁹
- SM4s, adult Wistar rat, superior cervical ganglia, 0.06 mol/mol phospholipid; nodose ganglia, 0.09; SM4s (d18:1/24h:0), vagus fibers, 0.12; SM4s (d18:1/24:0, 24 h:0), dorsal root ganglia, 0.14.¹⁷⁸ Dog, sciatic nerve, 70 nmol/g.⁴¹⁶ SM4s (d18:1/24:0, 24:1, 24 h:0), human femoral nerve, 3 $\mu\text{mol/g}$.⁵⁵¹
- SM4s, human peripheral nervous system (PNS) myelin-1, motor, 1.9% of the total lipid; sensory, 2.5%; PNS myelin-2, motor, 1.1%; sensory, 1.4%.⁴³⁴ Human cauda equina, 10 $\mu\text{mol/g}$.⁵⁵³ Human cauda equina, myelin, 62 mg/g dry myelin.⁵⁵³ Human spinal cord, 11 $\mu\text{mol/g}$ tissue; human spinal cord myelin, 39 mg/g dry myelin.⁵⁵³ Human motor nerve myelin, 53 nmol/g; motor nerve axons, 12 nmol/g; sensory nerve myelin, 49 nmol/g; sensory nerve axons, 12 nmol/g.⁵⁵² SM4s (d18:1/24:0, 24:1, 24 h:0), human skeletal muscle, 1 nmol/g.⁵⁵¹ Schwann cell line D6P2T, (+).²⁶ SM4s (hydroxy fatty acids), Schwann cell line S16 and S42, immunostaining, (+).⁵⁹⁴ Schwann cell line S16, [³⁵S]incorporation, (+).¹¹⁰ SM4s, mouse, G26-20 cells (one of the G26 series glioblastoma cell lines) 300 nmol/10⁹ cells.⁸⁵ Melanoma cell lines G361, and C32, [³⁵S]labeling, (+).⁴⁶⁹ The globular carboxyl domain of laminin A chain, a melanoma cell line A2058, (+).⁵⁸³ A melanoma cell line, TLC-OL, (+).⁵⁰⁹
- SM4s, human cerebrospinal fluid, 140 nmol/l.⁸⁴; control 145 ± 86 nmol/l; vascular dementia 307 + 118 nmol/l; Alzheimer, 178 ± 79 nmol/l.^{122, 124} Normal control, 99 nmol/l; AIDS patient, 395 nmol/l.¹⁴²
- SM4g, Schwann cell line D6P2T, (+).²⁶ Melanoma cell lines G361, and C32 (by [³⁵S]labeling), (–).⁴⁶⁹

Table 3—continued

SM3, melanoma cell lines G361, and C32, [³⁵S]labeling and TLC autoradiography, (+);⁴⁶⁹ a melanoma cell line A2058, (+)⁵⁸³
 SMUnLc₄Cer, dog sciatic nerve, 4.5 nmol/g; SMUnLc₆Cer, 3.3 nmol/g⁴¹⁶
 SMUnLc₄Cer, human spinal cord, 1.8 nmol/g; human, cauda equina, 64 nmol/g; femoral nerve, 39 nmol/g; cauda equina myelin, 112 nmol/g.⁵⁵² Human, sciatic nerve, adult, 8.2 nmol/g (65.0 μg/g dry tissue);⁶⁷ 53 nmol/g (0.85 μg/mg protein).¹⁶ Human cauda equina, 99.7 nmol/g (1.59 μg/mg protein).¹⁶ Adult human, motor (ventral root) nerve, 54 nmol/g; sensory (dorsal root) nerve, 72 nmol/g; motor nerve myelin, 193 nmol/g; motor nerve axons, 70 nmol/g; sensory nerve myelin, 199 nmol/g; sensory nerve axons, 80 nmol/g.⁵⁵² Schwann cells, (+)²⁴⁹
 SMUnLc₆Cer, rat, sciatic nerve, adult, 0.44 nmol/g (3.5 μg/g dry tissue)⁶⁷
 SMUnLc₆Cer, human spinal cord, trace; cauda equina, 15 nmol/g; femoral nerve, 10 nmol/g; motor nerve, 16 nmol/g; sensory nerve, 20 nmol/g; motor nerve myelin, 63 nmol/g; motor nerve axons, 25 nmol/g; sensory nerve myelin, 72 nmol/g; sensory nerve axons, 28 nmol/g.⁵⁵²
 SMUnLc₄Cer (SMUnLc₆Cer, human dorsal root ganglia, 64 nmol/g (1.02 μg/mg protein); human sympathetic ganglia, 2.5 nmol/g (0.04 μg/mg protein); dorsal root light myelin, 94 nmol/g (1.50 μg/mg protein); heavy myelin, 99 nmol/g (1.57 μg/mg protein); axolemma-enriched fraction, 107 nmol/g (1.71 μg/mg protein)^{15,16}
 SMUnLc₄Cer + SMUnLc₆Cer, melanoma G361 cells, [³⁵S]labeling, and TLC-OL with HNK-1 antibody, (+)⁴⁶⁹

Blood cells, serum, blood vessels and spleen

H₂O₂-Chol, human red blood cells, 5.0 nmol/g; human plasma, 3.1–3.9 nmol/ml.⁴¹ Serum, X-linked ichthyosis patient, elevated.⁶⁶¹ Human plasma, GC, normal control, 5.4 nmol/ml (253 μg/dl); cirrhosis patient, 9.6 nmol/ml (445 μg/dl); hypercholesterolemia patient, 8.9 nmol/ml (414 μg/dl).⁵⁸⁰ Human (>40-y), aorta (intima + media), 32 nmol/g (15 μg/g)⁹⁴
 H₂O₂-Chol, rat spleen, 33 nmol/g (77 μg/g dry tissue); thymus 1.8–1.9 nmol/g (4.2–4.5 μg/g dry weight).²⁴³ Rat spleen, [³⁵S]-labeling, (+)³⁵⁴
 SM4s (d18:1/16h:0), human erythrocytes, 0.62 nmol/g red blood cells (3.3 mg/6.7 kg wet cells).¹⁷⁵ Sheep, packed red blood cells, 5.4 nmol/g (4.3 mg/kg packed erythrocytes).⁴⁷¹ SM4s (d18:0, d18:1/16h:0), a bovine erythrocyte membrane preparation, 0.5 nmol/g (0.37 μg/g)³²³
 SM4s, human platelets, 1.5 nmol/g.⁴⁷¹ SM4s (d18:1/24:1 and 16 h:0), human, platelets, (+).³⁹⁴ SM4s, human granulocytes, flow cytometry and fluorescence microscopy, cell surface, (+); excreted into the medium, 5 μg/10⁸ cells/12 h; (+) breast carcinoma and myeloid cell lines; (–), leukemic T cell lines, HeLa and COS cells.¹⁸⁷ Leukocytes, immunodetection using the monoclonal antibody Sulf-I, (+).⁵⁶ SM4s (d18:1/24:0), bovine spleen, 1.1 nmol/g (1 μg/g tissue)⁴⁹⁶
 SM4s, porcine plasma, ¹H-NMR, (+).¹⁶⁵ SM4s (d18:1/22:0), mammalian sera: pig, 16 nmol/ml; goat, 13 nmol/ml; cow, 9 nmol/ml; horse, 8 nmol/ml; sheep 1.2 nmol/ml; human, 0.64 nmol/ml; dog 0.16 nmol/g; rat, mouse, chicken, not detected.⁶⁵³ SM4s, normal rabbit, serum, 3 nmol/ml; WHHL rabbit, serum, 121 nmol/ml; chylomicron, 1.2 mol% of phospholipids; VLDL, 1.9 mol% of phospholipids; LDL, 1.8 mol% of phospholipids; HDL, 1.4 mol% of phospholipids.¹⁸³ SM4s, normal rabbit, aorta (–); WHHL rabbit, aorta, atherosclerotic plaque, 280 nmol/g.¹⁷⁷
 SM3, porcine plasma, ¹H-NMR, (+)¹⁶⁵
 SMUnLc₄Cer, T-lymphocytes, (+)^{68,221}

Alimentary system

H₂O₂-Chol, human saliva, 4.3 nmol/ml.⁴¹ Rat stomach, [³⁵S]-labeling, (+).³⁵⁴ SD rat intestine, basolateral membrane, (+ +); H₂O₂-Chol, brush border (+).¹⁷³ Mammalian and cod fish intestine, (+).⁴⁷ Human feces, (+).³⁸⁸ H₂O₂-Chol + SM4s, human small intestinal mucosa, 5.2 mg/g dry weight¹⁰¹
 SM4s (20:0) + SM3 (18:0), rat sublingual gland, 940 nmol/g; SM4s (16:0) + SM3 (16:0, 18:0), rat submaxillary gland, 310 nmol/g.⁵²³ SM4s, rat (Wistar) glandular stomach, 28 nmol/g (136 nmol/g dry tissue).⁴¹⁴ SD rat small intestine, (–);¹⁷³ Wistar rat small intestine, (–) (Tadano-Aritomi, K. and Ishizuka, I. unpublished). Small intestine of hen, white rat, cat, rabbit, (+); mouse, black–white rat, guinea pig, (–).⁴⁷ SM4s, guinea pig gastric mucosa, 32 nmol/g (160 nmol/g dry tissue)³⁰⁰
 SM4s (d18:1/22–24h:0), rabbit, stomach, fundic mucosa, 79 nmol/g (394 nmol/g dry tissue); antral mucosa, 71 nmol/g; duodenum, 166 nmol/g; jejunum, 76 nmol/g.⁴¹⁴ SM4s (t18:0/22h:0, 23 h:0, 24 h:0), rabbit intestine, 31 nmol/g; WHHL rabbit intestine, 100 nmol/g.¹⁸³ SM4s (d18:1/24h:0), ileum, 12 nmol/g (58 nmol/g dry tissue); colon, 25 nmol/g (126 nmol/g)⁴¹³
 SM4s, dog fundic mucosa, 34 nmol/g (0.17 μmol/g dry mucosa); antral mucosa, 96 nmol/g (0.48 μmol/g dry mucosa).⁵²⁴ SM4s, dog intestine, 26 nmol/g.³⁷³
 SM4s, normal human gastric mucosa, 3.5–36 nmol/g; gastric cancer (tubular adenocarcinoma), 58–98 nmol/g.¹⁹⁰ Normal gastric mucosa, 56 nmol/g (0.25 μg/mg dry tissue); gastric mucosa (metaplasia), 89 nmol/g (0.40 μg/mg dry tissue); gastric carcinoma, 154 nmol/g (0.70 μg/mg dry tissue).³⁷⁷ SM4s, immunohistochemistry, (+) the chief cells of the stomach.²² Gastric cancer cell line, Kato III, (+) (a major acidic glycolipid).²⁵⁹ Esophagus mucosa, 3.3 nmol/g (16 nmol/g dry weight); SM4s (d18:1/24h:0, 26 h:0), fundic mucosa, 83 nmol/g (416 nmol/g); antral mucosa, 187 nmol/g (934 nmol/g); duodenum, 137 nmol/g (683 nmol/g); jejunum, 44 nmol/g (218 nmol/g); colon, 60 nmol/g (297 nmol/g); rectum, 31 nmol/g (153 nmol/g).^{412,414} Human colon (normal colonic mucosa), Azure A method using the Folch's lower phase, 20–170 nmol/g (0.20–1.70 nmol/mg protein); colonic cancer, 30–210 nmol/g (0.30–2.10 nmol/mg protein).⁵¹⁹

—continued

Table 3—continued

Human colon, the upper band, 27 nmol/g, the lower band, 56 nmol/g; colorectal cancer, the upper band 65 nmol/g, the lower band 11 nmol/g.³⁸⁷ Human, colon adenocarcinoma WiDr cell line, 90 nmol/g;³¹⁴ human colonic epithelial cell lines HT-29,¹⁰³ and Caco-2, (+);^{103,657} Flow cytometry, Colo 205 cell, (+)¹⁵⁴

SM4s (d18:1/24h:0), porcine gastric mucosa, 60 nmol/g.⁵²⁶ SM4s (22 h:0), pig intestine, 11 nmol/g (2 mg/22 kg)⁵⁴⁸

SM1b, C57B1/J mouse intestine, (+);⁴⁷ 7-week-old male, intestine, approx. 50 nmol/g (Tadano-Aritomi, K, and Ishizuka, I, unpublished).

Liver, gallbladder, and pancreas

HSO₃-Chol, rat (43-day-old) liver, 7–9 nmol/g (16.4–20.2 μg/g dry tissue).²⁴³ Human liver, 130 nmol/g (60 μg/g); human gallstones, 1.1 μmol/g (500 μg/g);⁹⁴ metachromatic leukodystrophy patient, 3.5 y, liver, 130 nmol/g.³⁸⁸ A patient with GM1-gangliosidosis, liver, 225 nmol/g.¹⁸²

SM4s, mouse liver, [³⁵S]label and silicic acid column, 416 nmol/g; rat liver, 22 days of age, 478 nmol/g; 117 day, 213 nmol/g;²⁴ CCL₄-induced liver inflammation (+).²⁵² SM4s, rabbit liver, 416 nmol/g.²³ SM4s (d18:1/22:0, 23:0, 24:0), rabbit, liver, 104 nmol/g; rabbit (WHHL), liver, 260 nmol/g.¹⁸³ SM4s (d18:1/24:0), rabbit hepatocyte plasma membrane, (+)⁵⁹³

SM4s, normal human liver, < 20 μg/g dry tissue.⁵³⁹ SM4s (d18:1/24h:0), metachromatic leucodystrophy (MLD) patient, liver, 62 nmol/g (280 μg/g dry tissue).⁵³⁹ Normal human liver, (+); liver, a patient of atypical Farber disease, (+ + +).¹³¹ SM4s (24:0), metachromatic leucodystrophy patient, liver, 3% of the total lipid; gallbladder, SM4s (d18:1/24:0), 35%.² SM4s (d18:1, d18:0/24h:0), a patient with GM1-gangliosidosis, liver, 69 nmol/g.¹⁸²

SM4s, [³⁵S]labeling/TLC, (+), a human hepatocellular carcinoma cell line PLC/PRF/5; a human cultured cholangiocarcinoma cell line, H-1; HepG2, KYN-1, KMCH-1 cell lines derived from hepatocellular carcinoma.²⁰⁴

SM3, liver, normal human, (–) (< 20 μg/g dry tissue); SM3 (d18:1/24h:0), metachromatic leukodystrophy, 39 nmol/g (208 μg/g dry tissue).⁵³⁹ SM3 (18:1), metachromatic leukodystrophy, 2% of the total lipid; gallbladder, 6%.²

SM3, [³⁵S]labeling/TLC, (+), human hepatocellular carcinoma cell lines PLC/PRF/5, HepG2, KYN-1, KMCH-1.²⁰⁴

SB1a, [³⁵S]labeling/TLC, (+), a human hepatocellular carcinoma cell line PLC/PRF/5.²⁰⁴

SM4s (d18:1/16:0 and 18:0), porcine pancreas, 10% by weight of the total acidic glycolipids.⁴⁰⁷ SM4s, human pancreas, immunological detection using monoclonal antibody Sulf I, 84 nmol/g; Langerhans islet, 150 pmol/100 islets (approx. 8.4 μmol/g islet tissue).⁵⁶ Lewis rat islet, 410 pmol/100 islets;⁵⁶ BB rat islet 87 pmol/100 islets; Wistar rat islet 181 pmol/100 islets; Lewis rat B cells 13 pmol/10⁵ cells.⁵⁶

Male reproductive organs

HSO₃-Chol, rat (43-day-old), testis, 15 nmol/g (34 μg/dry tissue).²⁴³ Rabbit treated with estradiol, testis, trace; prostate, 33 nmol/g (0.33 nmol/mg protein).⁸¹ Human semen, 14 nmol/ml,⁴¹ human spermatozoa (acrosome), (+).³⁵¹ Human seminal plasma 9.6 nmol/ml (445 μg/100 ml); sperm 32 nmol/10⁹ cells (15 μg/10⁹ cells).^{329,473} Rhesus monkey, sperm, 64 nmol/10⁹ cells (29.8 μg/10⁹ cells).³⁴⁸ Boar sperm plasma membrane, caput epididymidis, 0.8 nmol/10⁹ sperm; corpus, 0.6 nmol; cauda 1.4 nmol.⁴²⁸ Boar sperm acrosomal membrane, 140 nmol/g (1.4 nmol/mg protein).⁴²⁹ Boar sperm plasma membrane, before acrosome reaction, 410 nmol/g (4.1 nmol/mg protein); 1 hr after acrosome reaction, 40 nmol/g (0.4 nmol/mg).⁴²⁷

Desmosteryl sulfate, hamster epididymis, the major sterol sulfate.⁴⁷³ In rhesus monkey sperm not sulfated.³⁴⁸

SM4g (E_{16:0}A_{16:0}), C57 mouse, 800 nmol/g; W/W^t mutant, 120 nmol/g.³⁰⁶ Wistar rat, 7- to 10-day-old, testis, 100 nmol/g (1 nmol/mg protein); 30-day, 1400 nmol/g (14 nmol/mg protein); hypophysectomized rat, reduction of the SM4g concentration to 70% of the control.³⁰⁶ Mature rat, 2275 nmol/g testis (1.9 mg/g).¹¹⁹ SD rat testis, Azure A assay of the Folch lower phase lipids, 400 nmol/g, plasma membrane fraction, 13.5 μmol/g.⁵¹⁷ Rat, fed on vitamin A supplemented chow for 46 days, 556 nmol/g; fed on vitamin A deficient chow, 73 ± 46 nmol/g;⁵⁴⁶ Lewis rat, 4.9 μmol/g.⁵⁶ Rat (23 day) whole testis, 800 nmol/g (8 nmol/mg protein); rat spermatocyte, 4.4 μmol/g (44 nmol/mg protein).³⁴¹ Guinea pig, testis, 437 nmol/g.⁵⁴⁵ Rabbit treated with estradiol, testis, 400 nmol/g (0.40 nmol/mg protein).⁸¹

SM4g (E_{16:0}A_{16:0}), boar testis, about 300 nmol/g (0.8% by weight of the total lipid); boar spermatozoa, 1040 nmol/g cells.²⁵² Boar spermatozoa, perbenzoylation/HPLC, 980 nmol/g.⁵⁴⁴ Boar spermatozoa, 1.4 μmol/g cells.¹³⁴ SM4g plasma membrane of boar sperm obtained in epididymal caput, acetone fraction from silicic acid determined by Azure A method, 38.3 nmol/10⁹ sperm; corpus, 26.9 nmol; cauda, 19.6 nmol;⁴²⁸ boar sperm plasma membrane, 4.6–5.9 μmol/g;⁴²⁸ acrosomal membrane, 4.6 μmol/g (46 nmol/mg protein).^{427,429} Bull sperm, 8% of the total lipid.⁵¹¹

SM4g, human (2 m–11 y), (–).³⁴² SM4g (E_{16:0}A_{16:0}), human (40–50 y) testis, 170 nmol/g; seminoma, (–);²³⁸ 2–9 y, (–); 40 y, 159 nmol/g; 60–90 y, 25 nmol/g.⁶⁰⁵

SM4g (E_{16:0}A_{16:0}), chemical synthesis.¹⁴¹

Female reproductive organs

HSO₃-Chol, rat (43-day-old) uterus, 4.9 nmol/g (11.5 μg/g dry weight); ovaries 14.8 nmol/g (34.3 μg/g dry weight).²⁴³ Rabbit, endometrium, normal, 40 nmol/g; pseudopregnancy (day 4), 380 nmol/g.³⁸⁴

SM4s (d18:1/24h:0), human, uterine endometrium, proliferative (follicular) phase, 1–3 nmol/g (7–17 nmol/g dry tissue); secretory (luteal) phase, 23–49 nmol/g (115–245 nmol/g dry weight);³¹⁸ secretory phase, 12–46 nmol/g (60–230 nmol/g dry tissue).³¹⁹ Human, a uterine endometrial adenocarcinoma, 62 nmol/g (trace to 310 nmol/g dry tissue).³¹⁹ Human endometrial adenocarcinoma cell line Ishikawa, [³⁵S]sulfate incorporation, (+)²⁵⁵

—continued

Table 3—continued

SM4s, Lewis rat, ovary, 7 nmol/g. ⁵⁶ SM4s, normal human ovary, (-); SM4s (t18:0/22:0) mucinous cystadenocarcinoma (a clear cell adenocarcinoma), TLC-densitometry, 230 nmol/g (1.14 μ mol/g dry weight) (>90% of acidic glycolipids). ²⁸⁵ Choriocarcinoma cell line Rcho-1 (rat), TLC, (+). ⁵¹⁶ Human amnion, 338 pmol/g ¹⁸
SM3, human, uterine endometrium, secretory phase, 8 nmol/g (40 nmol/g dry tissue); SM3 (d18:1/24:0), human, a uterine endometrial adenocarcinoma, FAB, 46 nmol/g (trace to 230 nmol/g); SNG-II cell line derived from human endometrial adenocarcinoma, 158 nmol/g (790 nmol/g dry weight). ³¹⁹ Human endometrial adenocarcinoma cell line HEC 108, [³⁵ S]sulfate incorporation, (+). ²⁵⁵ Human ovarian clear cell carcinoma, (+). ²⁸⁵ Human amnion, 108 pmol/g ¹⁸
SM2a (d18:1/24:0), human, a uterine endometrial adenocarcinoma, FAB, (+); SNG-II cell line derived from human endometrial adenocarcinoma, 60 nmol/g (300 nmol/g dry weight). ³¹⁹ Human endometrial adenocarcinoma cell line HEC 108, SNG-M and SNG-II, [³⁵ S]sulfate incorporation, (+) ²⁵⁵
A sulfo-Gg ₄ Cer, human, a uterine endometrial adenocarcinoma, (+) ³¹⁹

Weights on dry weight or protein basis were recalculated into μ mol/g or nmol/g fresh tissue using the molecular weight of the major molecular species assuming the tissue water and protein content of 80 and 10% respectively unless otherwise specified.

cell respectively, supporting the results obtained by chemical analyses. The fatty acid composition of SM4s in human muscles and femoral nerve was very similar suggesting that SM4s of skeletal muscles was predominantly derived from the nerves in the muscle.⁵⁵¹

Guillain-Barré syndrome (GBS) has been characterized as an acute inflammatory demyelinating polyneuropathy, in which myelin is assumed to be the target of immune attack. The sera from patients with GBS displayed anti-SM4s antibodies in 67% of cases.^{122, 123} Serum antibodies from infants with congenital HCMV (human cytomegalovirus) infection interacted on TLC with the component sulfatides of the peripheral nerve, SMUnLc₆Cer and SMUnLc₄Cer, with high affinities, and with SM4g, SM4s, and SM3 with low affinities.⁴³⁵ Furthermore, the serum IgM from the HCMV infected patient interacted with a variety of sulfated glycolipids including SM4g, SM3, but not to HSO₃-Chol, suggesting that the sulfated sugar chains were the epitopes for the low-affinity interaction.⁴³⁵ Two glycolipids, SMUnLc₄Cer and SMUnLc₆Cer, from human peripheral nerve and cauda equina, react with monoclonal antibody HNK-1^{66, 70} and the IgM in patients with paraproteinemia and peripheral neuropathy.^{221, 656} They are present in human, monkey, bovine, dog, and cat peripheral nerves or in greatly reduced amounts in rat, mouse, rabbit, guinea pig and chicken.^{221, 249}

3. Blood, Blood Vessels, and Spleen

Red blood cells (RBC), white blood cells, blood plasma, thymus, and spleen contain low concentrations of HSO₃-Chol and/or sulfatides (Table 3).

SM4s is one of the major glycolipids in serum lipoproteins of various mammals except rodents^{183, 653} and enriched in the atherosclerotic plaque of Watanabe Hereditary Hyperlipidemia (WHHL) rabbit.¹⁷⁷ In serum lipoprotein of WHHL rabbit, an animal model for human familial hypercholesterolemia, SM4s content was elevated by about 40-fold (120 nmol/ml) (Table 3) over the normal level.¹⁸³ The intravenous administration of SM4s to WHHL rabbits lowered the level of triacylglycerols in serum, suggesting that SM4s might activate lipoprotein lipase or hepatic triacylglycerol lipase.⁵⁷¹

Negatively charged surfaces, phosphatidylinositol phosphate, SM4s, and dextran sulfate autoactivated the plasma contact activation system including human factor XII.⁴⁷⁵ Unlike heparin, SM4s failed to inhibit thrombin and coagulation factor Xa activities in the presence of antithrombin III (AT III). The SM4s micelle rather bound specifically to fibrinogen (400 mol SM4s per fibrinogen molecule) and thereby may interfere with both fibrin gel formation (anticoagulant activity) and platelet function¹⁷⁹ suggesting that SM4s may be an effective endogenous component for the prevention of thrombosis.¹⁸⁴ Actually, exogenous SM4s added to rabbit serum to 8 nmol/ml prolonged the fibrin-pre-

cipitation time.⁶⁵³ HSO₃-Chol showed no anticoagulant activity, while SM4s-6 was 20 times more potent anticoagulant, although the latter produced antibodies. Since AIDS drugs must be administered into the blood stream (cf. VII. D), the anticoagulant activity of sulfated polysaccharides may result in a significant side effect.^{276,277}

Staining with Sulf-I monoclonal antibody indicated that SM4s was expressed on the plasma membrane of granulocytes.^{17,56} P-selectin¹⁷ interacted with SM4s on the cell surface of granulocytes and several myeloid tumor cells. Exogenous SM4s may stimulate various cellular activities including the oxygen radical production of leukocytes (0.5×10^{-4} – 7×10^{-4} M)^{64,253} or neutrophils ($> 10^{-5}$ M),^{30,335} cytosolic Ca²⁺ level and oxidative burst of neutrophils via tyrosine kinase/MAP kinase system ($> 5 \times 10^{-5}$ M),^{30,622} monocyte phagocytosis of sulfatide-enriched human erythrocytes mediated possibly by thrombospondins;⁵¹³ accumulation of Ca²⁺, and secretion of interleukins with enhanced production of related mRNA in monocytes (4×10^{-4} M).⁷⁷ On the other hand, SM4s coated on *Staphylococcus aureus* cells stimulated phagocytosis of neutrophils, while had not effects on the superoxide anion release.⁶³⁵ Exogenous HSO₃-Chol was a better stimulant than SM4s, while SM4g, SM4s-6 and glucose 6-sulfate did not significantly stimulate the oxygen radical production of leukocytes.²⁵³ SM4s specifically triggered the increase of cytosolic free Ca²⁺ in neutrophils most probably through interaction with L-selectin.^{30,335} Secretion of TNF- α and IL-8 in monocytes (400 μ g/ml),⁷⁷ and neutrophils (> 10 μ g/ml)³³⁵ by SM4s, may correspond to the similar reaction elicited by 100 ng/ml of LPS. Currently drugs are under development to interfere with the interaction of selectins with their ligands. They include monoclonal antibodies to block the adhesive molecules, or competitive ligand saccharides, e.g. SM4s,³⁹³ 3'-sulfo-Lewis x, 3'-sulfo-Lewis a,⁶¹³ and lactose 6,6'-disulfate.³³ Not only vertebrate sulfatides, but also other acidic glycolipids including gangliosides, the monosulfated cord factor or TDM (trehalose dimycolate), and GlcU-containing sphingolipids from bacteria stimulated adhesion, phagocytosis, and phagosome-lysosome fusion of human neutrophils.³⁸⁰

4. Alimentary System and Liver

The mammalian saliva,⁴¹ gastrointestinal tract^{47,101,242} and feces³⁸⁸ contained HSO₃-Chol. The sublingual and submaxillary glands, and the gastrointestinal tract of mammals contained sulfatides^{81,361,537} (Table 3), whereas human saliva and gastric juice did not.⁴¹¹ The keratinizing squamous epithelium lining of the esophagus concomitantly showed high expression of prosaposin mRNA in the basal and spinous layers probably related to processing of sulfated lipids.⁵⁴⁰ The highest concentration of SM4s in the gastrointestinal tract was found with antral or duodenal mucosa of humans.⁴¹² In contrast, the concentrations of lipid-bound sialic acid in gastric mucosa, especially of fundic mucosa, were strongly reduced than those in the other parts of gastrointestinal tract⁵²⁴ resulting in the SM4s to lipid-bound sialic acid ratio of 1.4:2.3.⁴¹²

Sulfatides were isolated also from the intestine of mammals including porcine,⁵⁴⁸ dog,³⁷³ mouse,³³⁹ and WHHL rabbit,¹⁸³ and SM4s was even a major glycolipid for cat, rabbit, guinea pig, and hen intestine, reminiscent of the common developmental origin of intestinal epithelia and renal tubular cells.⁴⁷ Exceptionally mouse, rat and cod intestine did not contain SM4s, possibly due to the lack of GalCer similarly to some glioblastomas.²⁴⁴ Instead, HSO₃-Chol was enriched in the basolateral membrane of the rat small intestine, suggesting that HSO₃-Chol can replace SM4s.¹⁷³ Analogously, it appeared an opportunistic strategy for mice (C57B1/J strain) to sulfate Gg₄Cer in epithelial cells of small intestine, which lacks GalCer, to produce SM1b constituting at least 90% of the acidic glycolipids.³³⁹

Mucosal localization of SM4s has been demonstrated in the gastrointestinal tract of porcine,⁵²⁶ guinea pig,³⁰⁰ dog,⁵²⁴ rabbit,^{413,651} and human.^{101,190,377,412,519} The glycolipids were enriched in the epithelial but not in the nonepithelial compartment of the rat large intestine.¹⁷⁵ Both surface epithelial and glandular (parietal and chief) cells in the gastric mucosa of rabbit and human, but not the secretory granules, contained SM4s as

shown by immunofluorescence staining using a monoclonal antibody.⁵³⁸ It has been deduced that intestinal cell glycolipids make up even one third of the lipids in the apical membrane⁶⁵⁷ and essentially cover the whole surface of the luminal leaflet. The pre-epithelial mucosal defense activities⁷⁸ together with surface epithelial negative charges⁵³⁸ may protect mucosal cells against autodigestion by acid and pepsin, similarly to sucralfat (sucrose polysulfate aluminum complex), which has been widely used for treatment of gastritis and gastric ulcer.⁴¹⁴

[³⁵S]-labeling showed the presence of SM4s in rat liver,^{24, 151} although sulfatides were not detectable by chemical methods in the liver, except for the rabbit.¹⁸³ In the liver of patients of metachromatic leukodystrophy^{2, 539} and in a patient of atypical Farber disease,¹³¹ SM4s and SM3 were accumulated. Sera from patients with autoimmune chronic active hepatitis⁵⁹³ and systemic lupus erythematosus¹⁴ contained an IgG-class antibody to both the acidic glycosphingolipid fraction from rabbit hepatocyte plasma membrane, and SM4s. The reactivity of the hepatitis serum with SM4s was diminished by preincubation of the serum with SM4s-6 and SM4s itself, indicating that the antibody reacted with sulfated GalCer regardless of the position of the sulfate residue.

Sulf I monoclonal antibody immunogold staining showed the secretory granules of both A and B cells of the rat Langerhans islet contained SM4s. Furthermore, the SM4s synthesis was entirely abolished by destroying islets by streptozotocin treatment.^{56, 57} Assuming that 1% of the tissue weight of a pancreas consists of islets, SM4s concentration in human islets should be 8.4 μ mol/g islet tissue, which is about one third of the concentration in the white matter of the human brain.^{56, 57} SM4s antibody may be an IDDM (insulin dependent diabetes mellitus) marker because sera from 88% of newly diagnosed IDDM patients were anti-SM4s positive, and 76% were positive 6 months later while all healthy controls were negative.⁵⁵ IDDM patients' sera positive for SM4s stained secretory granules in A and B cells of rat islets of Langerhans.

5. Reproductive Organs

HSO₃-Chol was enriched at the plasma membrane of human,³³¹ and boar^{427, 428} spermatozoa and acrosomal membrane of boar.⁴²⁷ HSO₃-Chol or other steroid sulfates (HSO₃-Chol + desmosterol sulfate) of hamster⁴⁷³ and boar^{428, 429} spermatozoa increased during their epididymal transit. In contrast, the SM4g content of the sperm plasma membranes decreased by about 50% during epididymal transit from the caput to the cauda.⁴²⁸ Then, at capacitation in the female reproductive tract, drastic desulfation of HSO₃-Chol was observed at the plasma membrane of hamster.^{331, 473} Steroid sulfates, including HSO₃-Chol, have been known to stabilize certain membranes like the erythrocyte plasma membrane^{107, 329} and keratinocytes.^{107, 630}

In the testis of mammals including rat,^{238, 395} guinea pig,⁵⁴⁵ human,^{237, 342, 606} boar,²³² and bovine,³⁵⁴ SM4g (seminolipid) is the principal glycolipid of testis comparable in amount with HSO₃-Chol³⁴⁸ (w/w). In the lipid from ejaculated boar spermatozoa, SM4g comprised approx. 3% of the total lipid, that is more than 90% of the sum of glycolipids, and the concentration (1 μ mol/g wet cells) was approximately three times greater than that in the testis.²³² The *w/w^v* mouse testis, with a greatly reduced content of all germinal cells, contained a negligible amount of SM4g.³⁰⁶

Between 11 and 20 days in mice, intraperitoneally injected [³⁵S]sulfate was actively incorporated into SM4g of the testis.^{171, 253} There were only low amounts of lipid-bound galactose and lipid-bound sulfate in the testis of rats younger than 15 days,³⁰⁶ whereas between 15–22 days in rat,³⁰⁶ there was a dramatic increase in both lipid-bound sulfate and a sulfotransferase activity, although the total lipid concentration remained relatively constant.³⁹⁵ This age corresponds to the appearance of spermatocytes in rat testes and actually, isolated late spermatocytes contained SM4g at 5 times the level in the whole testis.³⁴¹ The activity of GalCer sulfotransferase in rat testes increased in maturing rats of 10–15 days, then reached a plateau from day 15–26 after birth, and finally decreased to one third of the maximal activity in rats of 50 days or older.^{305, 341, 517} [³⁵S]sulfate,⁵⁴⁵ [2-

^{14}C]Gro, and dihydroxy[U- ^{14}C]acetone⁵¹¹ were not incorporated into SM4g of spermatozoa, indicating that no synthesis of SM4g occurred after ejaculation. On day 28 after the pulse-labeling of rat with [^{35}S]sulfate, the majority of total radioactivities of the testicular lipid [^{35}S] declined and reached a minimum level on day 34.³⁴¹ Conversely, the radioactive level in the caudal epididymal sperm reached a maximum on day 32. Specific activities of the radiolabel in both the epididymides and the testes followed a similar trend. The highest specific activity on day 32 in the epididymis was up to 180 times greater than that in the testis.⁵⁸²

Neither SM4g nor Gal-EAGro was detected in the testes of infants or a child of prepubertal age.⁶⁰⁶ At 40 years, both the total lipid and SM4g contents were highest.⁶⁰⁶ Above 70 years of age, the concentration of SM4g decreased drastically, to approx. 1/6 of that in the 40 s, probably due to the reduction in the number of seminal tubules. In contrast, ganglioside content was highest in the testis of the aged, probably reflecting the fibrosis of the testis.

By using a monoclonal antibody specific to sulfatides, a dense staining inside each seminal tubule was noted in the rat testis, and in epididymis an intense labeling of the spermatozoa was seen.⁵⁶ Monoclonal antibodies specific to SM4g also showed that SM4g was enriched at the plasma membrane of spermatozoa.^{97, 135, 428, 447, 517} Actually, the isolated boar sperm acrosomal membrane contained an extremely high concentration of SM4g.⁴²⁸ In freshly ejaculated sperm cells, SM4g was present primarily at the apical ridge subdomain of the plasma membrane of the sperm head.¹³⁵ During capacitation of the sperm cells by Ca^{2+} ion,⁶⁴³ SM4g migrated rapidly from the apical ridge to the equatorial subdomain of the plasma membrane.¹³⁵ An addition of arylsulfatase A to the sperm cells desulfated SM4g, and the product Gal β -EAGro also appeared specifically at the equatorial ridge. The interaction and degradation of the sperm SM4g by *M. pulmonis* may play a role in the induction of infertility that follows infection with these organisms by interfering in sperm/egg receptor recognition.³⁵⁴ Prosaposins are contained in the seminal plasma²⁰² and the substrate glycolipids are located in, or at the exterior surface of the plasma membrane and acrosomal membrane of spermatozoa.^{134, 135, 618}

The presence of arylsulphatase A and SM4s in different tracts of *Rana esculenta* oviduct during different phases of the reproductive cycle was studied by histochemical and biochemical procedures.⁶¹⁵ The results indicate that enzyme activity and SM4s level show seasonal fluctuations connected with the phase of the sexual cycle. The SM4s concentration in human endometrium also changed associated with the menstrual cycle. The concentration at the proliferative phase, 1–3 nmol/g, was elevated to 23–49 nmol/g at the secretory (luteal) phase^{254, 318, 319} concomitant with the elevation of galactosyl-²⁵⁵ and sulfotransferase activities.²⁵⁴ In rabbit,³⁸⁴ and guinea-pig,⁴²² which are deficient in glandular organization, sulfatides were present in very small quantities in endometrium irrespective of the reproductive stage, whereas $\text{HSO}_3\text{-Chol}$ increased sharply at day 5 of pregnancy, which is the beginning of implantation.³⁸⁴ Pseudopregnancy of rabbit or administration of human chorionic gonadotropin after estrogen priming also stimulated $\text{HSO}_3\text{-Chol}$ synthesis and in the latter case, 15% of the total cholesterol was converted to the sulfated form concomitant with a 30-fold increase of cholesterol sulfotransferase activity.³⁸⁴ As in the rat intestine (V.C.4), $\text{HSO}_3\text{-Chol}$ in the endometria of these animals appears to replace SM4s. In the rat ovary, Sulph I (an anti-SM4s monoclonal antibody) distinctly labeled the ovum (oocytes), whereas the follicle epithelium, the theca and rest of the ovary were unstained.⁵⁶ In the rat oviduct (uterine tube), parts of the epithelium were labeled, whereas the uterus was negative.

6. Tumors

Sulfatides are expressed in mammalian malignancies.^{121, 137} For instance, sulfatides were increased in the adenocarcinomas including gastric^{190, 377} and renal cell carcinoma (Grawitz),⁴⁹⁰ but not in Wilms' tumor tissue.⁴⁸⁹ The accumulation of sulfatides was associated with the elevated activity of sulfotransferase in renal cell carcinoma,⁴⁹⁰ and gas-

tric adenocarcinoma.^{351, 377, 489} Human adenocarcinoma tissues, originated from the gastrointestinal tract,^{387, 519} and the related cell lines³¹⁴ also contained high concentrations of sulfatides (Table 3). It is interesting that the level of GalCer sulfotransferase was the highest in intestinal metaplasia.

In contrast to the differentiated, thus only weakly tumorigenic renal epithelial cell lines (MDCK, JTC-12),⁵⁵⁵ the specific activity of sulfotransferase toward GalCer in the renal cancer cell line SMKT-R3²⁹³ was 50-fold greater than that in normal human kidney tissue (42.0 pmol/mg protein/hr),⁴⁸⁹ and 8-fold greater than that in renal cell carcinoma tissues.²⁹³ SMKT-R3 contained SM4s, SM3, SM2a,²⁹³ and several other minor sulfatides²¹¹ concomitant with sulfotransferase activities synthesizing SM2b and SB2.²⁹² The level of SM4s in the lung of normal human adult and squamous cell carcinoma of lung was low,⁶⁴⁵ whereas pulmonary adenocarcinoma contained more than 10 times higher SM4s than the normal lung, accompanied by an elevated activity of both GalCer sulfotransferase⁶⁴⁵ and arylsulfatase A.^{139, 409}

In 29% of human hepatocellular carcinoma tissues, the accumulation of SB1a was observed.^{204, 205} In addition, approx. 88% of hepatoma tissues and 50% of cirrhotic liver contained a significant amount of SM3, although the level of the sulfotransferase activity in hepatoma tissues of human and rat was not distinguished from that of normal controls.²⁵⁸ Clonal tumor cell lines G26-20 and G26-24 of rat, supposed to be originated from neuroectoderm, incorporated [³⁵S]sulfate into SM4s.⁸⁵ Three types of glycosphingolipid (GSL) component profiles have been established for human intracranial gliomas.²⁴⁴ Only glycosphingolipid (GSL)-type III gliomas contained GalCer and SM4s.

Human testicular seminoma, an extremely undifferentiated tumor originated from the epithelium of seminiferous tubules, did not contain SM4g.²³⁸ On the contrary, uterine endometrial carcinoma tissues contained an elevated concentration of SM4s. A high concentration of SM3, SM2a and a sulfo-Gg₄Cer appeared after 70th doubling time of the endometrial adenocarcinoma cell lines SNG-M³¹⁹ and HEC 108²⁵⁵ in culture with concomitant defect of SM4s, arousing caution in the analysis of sulfatides in cultured cells. The differential expression of sulfatides between Ishikawa cells and the other three endometrial adenocarcinoma cell lines may be due to the absence of LacCer in the former and the lack of GalCer in the latter.²⁵⁵ Approximately 80% of the lipid [³⁵S]sulfate in human breast ductal carcinoma T47D cells was found in HSO₃-Chol.⁵⁸³

Sulfotransferase has been implicated as the potential tumor marker enzyme. Activities of glycolipid sulfotransferase in serum were elevated in 33% of patients with hepatocellular carcinoma.¹³⁶ In comparison, the sulfotransferase levels¹³⁷ in sera from patients with renal cell carcinoma were lower than those in hepatocellular carcinoma. It remains to be clarified why hepatocellular carcinoma, which is poor in SM4s, releases more sulfotransferase into serum than does renal, pulmonary and gastric¹⁹⁰ carcinomas,²⁵⁸ which are rich in the enzyme.

D. Molecular Evolution

The phylogenetic descendency of glycolipids has been discussed on the view point of the chemical structure of saccharides,⁶²⁹ glycosyl transferase genes,⁴⁸⁴ and Darwinian vs. neutral evolution of saccharide structure.²³⁸ The cloning of the cDNA of A, B, and O glycosyltransferases showed that the mutation of transferase to synthesize ABO blood group antigens resulted in no pressure in selection.⁶⁴⁰ Recently, inactivation of a specific glycosyltransferase gene is replacing the conventional 'experiments of the nature' to prepare animals with enzyme deficiencies. In many cases of deletion of a selected glycosyltransferase gene in knockout mice, the survival and the early development of the animal were apparently normal supporting the proposition that most of the oligosaccharide structures have been almost neutral in evolution.²³⁸ However, higher and more specific functions were usually affected. In other words, oligosaccharides are analogous to the 'axle grease' of an automobile.⁶¹¹ While its absence would markedly affect the ability of

the entire vehicle to function, the fine details of the composition of the grease should not be critical to the turning of the axle.

The concept of lipid class replacement by Rouser⁴⁸⁰ explained why one of the major phospholipid, sphingomyelin, should substitute phosphatidylcholine in biomembranes. The myelin structure of mice lacking the enzyme UDP-galactose:ceramide galactosyltransferase contained GlcCer and sphingomyelin with hydroxy fatty acids instead of GalCer and SM4s.⁷⁵ The apparently compensatory distribution of HSO₃-Chol and SM4s in the gastrointestinal tract, sperm, endometrium, and kidney of various species also supports the above concept. The reversible replacement of acidic phospholipids with GlcU-containing glycolipids or SQ-A₂Gro,¹⁵⁵ and the polysaccharide teichoic acid with teichuronic acids²³⁸ under phosphate limitation are representative examples in eubacteria and cyanobacteria. HSO₃-PtdGro in *H. mediterranei* (strain R-4) appears to be replaced by an equivalent increase in the amounts of HSO₃-6Man-2Glc-E₂₀E₂₀Gro (S-DGD).³²⁶ The major barrier amphiphiles including phosphatidylglycerol in bacteria,^{60,155} and sulfatides in halophilic archaea or vertebrates (for rev. ref. 238) increased in adaptation to high environmental osmolality. These anionic lipids arose most likely as the result of convergence of the structure in molecular evolution. Genes of the adaptive machinery can be supplied from phylogenetically widely separated species. A possible descent of sialic acid-related and polygalacturonate-related genes from a common ancestor gene has been suggested.⁵⁰¹ The genes related to the sialic acid metabolism such as sialidase found in prokaryotes may have been transferred from the host animal via phages horizontally to bacteria.⁴⁷⁴

Lipophilic residues also seem to be replaceable.²³⁸ The gram-negative rods of genus *Sphingomonas* contained GlcUCer³⁸⁰ and a 'lipid A-type', glucosamine-containing tetrasaccharide ceramide instead of lipopolysaccharide (LPS) usually found in the gram-negative bacterial membrane.²⁸⁰ The membrane fraction of *Sphingomonas paucimobilis*, the LPS-lacking gram-negative bacterium, contained glycosphingolipids that was assumed to have a function similar to that of the LPS of other gram-negative bacteria.²⁸⁰ The hybrid structure of this LPS is interesting in two ways: (1) ceramide with cyclopropanic sphinganine linked to a saccharide containing glucosamine amide, which in turn, is linked to 2-hydroxymyristic acid instead of 3-hydroxymyristic acid in normal lipid A, can replace LPS; (2) phylogenetically, the genes related to the metabolism of ceramide must have been transferred from the eukaryote host. Even the presumably advantageous molecular device such as LPS that has survived a billion years was able to be substituted by a glycosphingolipid!⁶⁵⁹ The neutral theory dictates that the great majority of evolutionary changes at the molecular level are not the sequel of Darwinian selection but the consequence of the random fixation of selectively neutral²⁸⁶ or very nearly neutral⁴⁴⁰ alleles under continued mutation pressure. Farther examples of the neutral evolution of sulfatide structures and replaceability with other anionic lipids will be accumulated in future.²³⁸

VI. BIOSYNTHESIS AND BIODEGRADATION

In as early as 1960, [³⁵S] labeling of rat *in vivo* demonstrated that brain, kidney, and liver incorporated sulfate into lipid fractions.^{24,151} Recently, a GalCer sulfotransferase was purified from human renal cell carcinoma,²¹¹ and the cDNA was cloned.²¹⁰ On the other hand, the sulfate is released from sulfatides by a lysosomal soluble enzyme and the activator protein on the lysosomal membrane.³⁰²

A. Biosynthesis

1. *In Vivo* Studies

Labeling of the whole animal or intact cell with [³⁵S]sulfate or [³⁵S]methionine^{24,382} and the chase of the incorporated glycolipid-[³⁵S]O₃⁻ reflected the turnover rate, the

difference between synthesis and degradation, of sulfatides. The turnover of sulfatides in the whole body of eels,⁶⁵⁴ rats^{24, 151, 233} and mice^{53, 171, 608} was studied by intraperitoneal injection of [³⁵S]sulfate to animals. The correction for the age-dependent blood level was necessary for the quantitative estimation of the turnover rate because the blood [³⁵S]sulfate level decreased with time after injection of the isotope.⁵⁴

The maximal incorporation of the isotope into rat brain lipids occurred 48 hr after the administration of radioactive sulfate, after which time the activity remained constant through the 16th day, but slowly fell thereafter; although on the 32nd day, the level was still 3/4 of that found on the second day¹⁵¹ and the specific radioactivity of [³⁵S]SM4s in myelin decreased only slightly even after 197 days.²⁴⁸ SM4g but not HSO₃-Chol was significantly labeled by [³⁵S]sulfate.^{120, 228} Incubation of rat brain slices with [³⁵S]sulfate labeled only SM4s⁵⁹⁶ whereas rat sciatic nerve slices incorporated [³⁵S]-radioactivities into both SM4s, and SMGlcUnLc₄Cer.⁶⁸ The half-life of kidney sulfatides was considerably shorter than that of testis and myelin.²³³ Intraperitoneal administration of [³⁵S]sulfate to rats labeled HSO₃-Chol,^{233, 559} SM4s, SM4s-Glc,²¹⁷ SM2a,⁵⁵⁷ SB2,⁵⁵⁸ and SB1a in kidneys.⁵⁶¹ In the testis of mice¹⁷¹ or rat,^{189, 305, 352, 517} intraperitoneally administered [³⁵S]sulfate labeled only SM4g. [³⁵S]Sulfate injected intratesticularly,^{119, 306, 341} or administered *per os*⁵⁸² into rat was also incorporated into SM4g in the testis whereas HSO₃-Chol was not significantly labeled. After incubation of guinea pig testicular slice in a medium containing [³⁵S]sulfate, autoradiography of the lipid fraction showed only SM4g.⁵⁴⁵

Primary cultures of animal cells and established cell lines are the convenient system for metabolic studies. Oligodendrocytes,²⁷ as well as Schwann cell lines^{26, 110} incorporated [³⁵S]sulfate into SM4s, SM4g and several sulfated gangliosides,¹¹¹ and melanoma cells into HSO₃-Chol, SM4s, SM3, SMUnLc₄Cer, and SMUnLc₆Cer.^{469, 583} Incubation with [³⁵S]sulfate of the renal tubular epithelial cell lines including MDCK,^{49, 234, 423} JTC-12,^{234, 555} MDBK,^{234, 424} LLC-PK₁,⁴²⁴ SMKT-R3 renal carcinoma cells^{291, 293} and Verots cell lines⁴²⁵ labeled HSO₃-Chol, SM4s, SM3, and some sulfatides of the ganglio-series. Hepatocellular carcinoma cell lines, a cholangiocarcinoma cell line,²⁰⁴ and endometrial adenocarcinoma cells^{255, 319} of humans incorporated [³⁵S]sulfate into SM4s and/or SM3. Photosynthetic bacteria¹⁵⁵ and myeloid cell lines¹⁷ were cultured in a commercial sulfate-free medium (CRCM-30, Sigma), or a medium with the sulfate salts substituted with chloride salts, then labeled with carrier-free [³⁵S]sulfate. However, the results of sulfation in sulfate-deprived (carrier-free) medium should be evaluated carefully⁶¹¹ since culture of cells using a sulfate deficient medium can result in undersulfation of glycosaminoglycans in some cell types⁶¹² and sulfation of each chain is an 'all or nothing' process.

Dissociated primary cultures of rat brain were incubated with varying quantities of galactose with a fixed amount of radioactivity per culture.⁵²¹ In a medium containing 1 mM galactose, more than 85% of the label was present in the carbohydrate moiety of glycolipids.⁵²¹ When cells were incubated with [³H]-hexose,²³⁰ [¹⁴C]hexose-, or [¹⁴C]serine,⁶² radioactivities were distributed in the fatty acid, sphingosine, and sugar moieties.

2. Sulfotransferase

Little has been reported on the properties of glycolipid sulfotransferase of halophilic archaea and mycobacteria.^{149, 273} The sulfotransferases of plant and animal cytosol sulfate flavonoids, steroids and other aryl compounds, but not glycolipids.⁶⁵⁸ In photosynthetic prokaryotes and plant chloroplasts, 6-sulfoquinovose is transferred from UDP-6-sulfoquinovose to the position *sn*-3 of A₂Gro by the catalysis of UDP-sulfoquinovose: diacylglycerol sulfoquinovosyltransferase.¹⁸⁸ When the gene of this enzyme, which catalyzes the last step of SQ-A₂Gro synthesis of photosynthetic bacterium *Rhodobacter sphaeroides*, was inactivated, the resulting sulfolipid-deficient mutant accumulated the precursor UDP-sulfoquinovose.¹⁵⁵

The *in vitro* formation of SM4s, via transfer of sulfate from 3'-phosphoadenosine 5'-phosphosulfate (PAPS) to GalCer catalyzed by a sulfotransferase, GalCer sulfotransferase (EC 2.8.2.11), has been demonstrated in brain,^{108,361} kidney,^{104,105,495} testis,^{171,289,353} rat gastric mucosa,³⁷⁷ lung,¹³⁸ human endometrium,^{255,319} and sera from patients with renal cell carcinoma.^{136,137}

The GalCer sulfotransferase activity was demonstrated in the Golgi-rich fraction from rat kidneys,¹⁰⁵ rat testis,²⁸⁹ and a Schwann cell line.¹¹⁰ The cytosolic sulfotransferases act on aryl compounds, hydroxysteroids, estrogens, and bile acids.^{137,210} The localization of GalCer sulfotransferase at the lumen of Golgi apparatus is similar to that of glycosaminoglycan sulfotransferases and tyrosylprotein sulfotransferase. The site of sulfation may be localized to the distal Golgi or trans-Golgi network¹¹⁰ because brefeldin, which destroys (or blocks vesicular transport) distal to medial Golgi, inhibited SM4s synthesis completely, while both hydroxy and nonhydroxy GalCer were synthesized. In contrast, 2-hydroxyceramide galactosyltransferase is thought to be located on the endoplasmic reticulum of myelinating rat brain,⁵¹⁰ and MDCK II cells.⁵¹

Some drugs affect the synthesis of sulfatides without modification of sulfotransferase. Incubation of the myeloid cell lines THP-1, and HL60,¹⁷ or a human renal carcinoma cell line SMKT-R3²⁹³ in the medium containing sodium selenate, an inhibitor of PAPS formation, resulted in a reduction of SM4s expression. Monensin, an inhibitor of vesicular transport from the medial- and trans-Golgi to the plasma membrane, also inhibited 75% of [³⁵S]O₄²⁻ incorporation into sulfolipids.⁶¹⁷ Brefeldin blocks, in a reversible manner, the anterograde movement of vesicular traffic from the endoplasmic reticulum (ER) to the Golgi, allowing retrograde movement back to the ER to continue.¹¹⁰

Honke and his group²¹¹ purified PAPS:GalCer sulfotransferase from a human renal cancer cell line SMKT-R3 through a combination of affinity chromatographies using GalSph, 3',5'-bisphosphoadenosine and heparin as ligands.²¹¹ The purified human renal sulfotransferase showed a highest specific activity of 18,000 μ mol/mg/hr²¹¹ and a single protein band with an apparent molecular mass of 54 kDa. pH optima were between 6.5,²¹¹ to 7.0⁵⁴² similar to other transverses of Golgi membrane. The putative hydrophobic transmembrane domain of GalCer sulfotransferase from human renal carcinoma cell contained 23 amino acid residues characteristic of type II transmembrane proteins. The cloned GalCer sulfotransferase showed homology neither to the cytosolic sulfotransferases nor to the Golgi glycosaminoglycan sulfotransferases.

3. Substrate Specificity

GalCer, that is abundant in mammalian oligodendrocytes, kidneys, and the gastrointestinal tract, was usually the best acceptor for GalCer sulfotransferase. LacCer,^{109,171,555} Gal-EAGro,^{171,487} and Gal-A₂Gro^{211,289,463,586} were also good acceptors. GalSph,⁵⁸⁶ GlcCer, Gg₄Cer, nLc₄Cer, Gg₃Cer, SM2a,²⁹² and Gb₄Cer did serve as acceptors although the relative activities were low.²¹¹ The glycolipids with the terminal α -Gal in Gal α -4GalCer^{171,463} or Gb₃Cer,^{171,211} and cholesterol⁴⁶³ did not serve as acceptors. GlcCer appeared not to be the substrate of sulfotransferase in the brain of mice deficient in GalCer.⁷⁵ Absence of sulfatides with an internal *N*-acetylgalactosamine 3-sulfate suggests that GalNAc 3-sulfate might be a termination signal against further sulfation or glycosylation of the glyco-amphiphiles. On the other hand, oligosaccharides including galactose and lactose did not serve as substrates or a competitive inhibitor for the purified enzyme from the renal carcinoma cell.²¹¹

Competition studies suggested that a single enzyme in the boar testis,¹⁷¹ as well as in cultured renal cells, MDCK, JTC-12,⁵⁵⁵ and SMKT-R3,²⁹³ sulfated GalCer and LacCer. Highly purified human renal sulfotransferase sulfated LacCer and Gal-EAGro rapidly establishing that a single enzyme synthesizes SM4s, SM4g, and SM3.²¹¹ A rat brain sulfotransferase was highly specific to GlcUnLc₄Cer and the rate-limiting step of the synthesis of SMUnLc₄Cer was the activity of a GlcNAc transferase.^{69,70} The UDP-*N*-acetylgalactosamine: SM3 *N*-acetylgalactosaminyltransferase activity of rat brain, which

synthesized SM2a, recognized both GM3 and SM3 as equally good acceptors.^{230, 402} This lack of stringency suggested that the GalNAc transferase step cannot be a rate-limiting step of glycolipid biosynthesis.

The optimal assay conditions for GalCer with nonhydroxy and hydroxy fatty acids, and LacCer were significantly different.^{171, 555} GalCer with hydroxy fatty acid, synthesized at the cytosolic leaflet of the endoplasmic reticulum of MDCK II cells, may be translocated to the luminal leaflet of the Golgi membrane whereas the activity to synthesize GlcCer and GalCer with nonhydroxy fatty acids may be located on the cytosolic leaflet of proximal Golgi.⁵¹ A ceramide containing a 2-hydroxy fatty acid may be preferentially converted to GalCer, Ga₂Cer, and SM4s by renal distal tubular cell line MDCK and colon carcinoma Caco-2 cells.⁶⁵⁷ In mouse brain, however, only one ceramide galactosyltransferase exists that catalyzes the synthesis of both nonhydroxy and hydroxy fatty acid GalCer.⁷⁵

4. Intracellular Traffic and Membrane Polarity

The apical membrane of polarized cells is enriched in glycosphingolipids.⁶⁵⁷ Luminal (apical), but not basolateral, membrane of transporting cells including MDCK was enriched in both phospholipids and glycolipids^{340, 655} including SM4s.⁴⁹ By transport experiments using Cer with shorter chain fatty acids, on the contrary, sphingomyelin, GalCer with 2-hydroxy fatty acids, LacCer and SM4s were transported preferentially to the basolateral cell surface when compared to GlcCer.⁶⁵⁷ When immature rat eyes were incubated in a medium containing [³⁵S]sulfate, there was a rapid *in vitro* uptake of the label into SM4s of the ciliary processes, the site of active transport.³² Electron microscopic autoradiography showed that initially (7 min-label) the label was localized in the apical cytoplasm where the Golgi apparatus is located but with longer periods (30–60 min-chase) of incubation in nonradioactive medium it moved to the infolded basal and lateral plasma membrane.

Exogenous glycosphingolipids are incorporated into the plasma membrane¹²⁸ or internalized by endocytosis,^{386, 655} packed in a vesicle and sent to lysosomes.¹²⁸ The fluorescent analogue of SM4s, *N*-lissamine rhodaminyl-(12-aminododecanoyl)-SM4s (LRh-SM4s)-albumin complexes (and not the fluorescent SM4s alone) was transferred into the plasma membrane by the cell. In order to incorporate glycolipids into SMKT-R3 cells, SM4s dissolved in a small amount of dimethylsulfoxide was exogenously added to the cells.³²⁸ Cytofluorometry showed that the incorporation of SM4s into the cells considerably increased the reactivity with Sulph-I and laminins in a dose-dependent manner.²⁹⁴ LRh-SM4s complexed to albumin were used to study the metabolic fate of SM4s in oligodendrocytes under microscope.^{386, 617} The degradation at 24 hr-pulse reached 18% in normal fibroblasts, whereas it was not detectable in metachromatic leukodystrophy cells. An exogenous [³H]labeled ceramide with hydroxy fatty acids adsorbed on BSA was preferentially incorporated into GalCer and SM4s in comparison to a ceramide with nonhydroxy fatty acids.⁶⁵⁷

B. Regulation

Lipophilic and hydrophilic hormones modify the expression and activities of sulfotransferases by the interplay with nuclear receptors and promoters of genes, and transmembrane cascades respectively.

1. Lipophilic Hormones

Incorporation of [³⁵S]sulfate into HSO₃-Chol fraction of rabbit tracheal epithelial cells increased 50–100-fold upon differentiation, suggesting that the accumulation of HSO₃-Chol can be a marker of differentiation of tracheal cells.⁴⁶⁴ This accumulation was the result of induction of cholesterol sulfotransferase and was completely blocked by the in-

clusion of retinoic acid analogues, the inhibitor of squamous differentiation, in the culture medium.⁴⁶⁴ Progesterone in association with estradiol, stimulated the synthesis of HSO₃-Chol in rabbit endometrium³⁸⁴ and HSO₃-Chol and hydrophobic sulfated conjugates in subcultured glandular epithelial cells of guinea-pig endometrium.⁴²²

Reduction of the level of SM4g to approx. 70% of the normal testis in hypophysectomized rats may reflect the suppression of spermatogenesis concomitant with the decrease in androgens of both adrenocortical and testicular origin.³⁰⁶ Retinoid (Vitamin A) deficiency also caused degeneration of seminiferous tubules and results in sterility of the rat. SM4g in the testes of rat fed a retinol-deficient diet for 46 days decreased to 13% of the control rats fed a retinol-deficient diet for 20 days and then supplemented with 140 µg/rat/day of retinol palmitate for 26 days.⁵⁴⁶ Vitamin K treatment for 3 days significantly enhanced brain sulfotransferase activities, whereas administration of the vitamin K antagonist, Warfarin, drastically reduced the enzyme in 2 weeks.⁵⁴¹

The thyroid hormones, thyroxine (T₄) and triiodothyronine (T₃), are essential for the maturation of oligodendrocytes and play a stimulatory role in myelination of the central nervous system. The rate of synthesis of sulfatides remained drastically diminished throughout a 70-day developmental period when brain cells from embryonic mice were grown in the presence of hypothyroid calf serum,³⁶ while the activity was restored to normal levels after 72 hr of exposure to a medium supplemented with exogenous T₃. In the presence of T₃, the incorporation of [³⁵S]sulfate into sulfatides (SM4s and SM4g) of oligodendroglial cultures obtained from the brains of 1-week-old rats exhibited a developmental profile which is comparable to that found in the developing brain *in vivo*³⁰⁴ whereas omission of T₃ resulted in lower rates of sulfatide synthesis. Incorporation of [³⁵S]sulfate into SM4s fraction of glioma cells, G26-19 and C26-20, was stimulated up to 2-6-fold in the presence of 5 × 10⁻⁶ M cortisol or dexamethasone in culture media with the concomitant 3-4-fold stimulation of sulfotransferase activities.^{86,250} Experiments with cycloheximide and actinomycin D showed that the effect of the hormone on glycolipid synthesis on these cells was mediated through *de novo* mRNA and protein synthesis.²⁵⁰ Testosterone, androsterone, and estradiol had no stimulatory effect.

2. Water-soluble Hormones and Growth Factors

[³⁵S]sulfate incorporation into HSO₃-Chol increased in JTC-12 and MDCK cell lines by additions of butyrate to culture medium.²³³ The incorporation into SM2a and GM2 in JTC-12 cells was also stimulated by [³⁵S]labeling *in vivo* depending on the dose of butyrate.⁵⁵⁶ Only a trace amount of radioactivity was incorporated normally into SM3 of MDBK cell line, whereas by additions of butyrate (2.5 mM), the incorporation into SM3 increased 20-fold. By incubation of the culture of C₆ glioma cell line with 50 mM disipramine, a tricyclic antidepressant, SM4s synthesis was substantially stimulated but HSO₃-Chol level was unchanged.⁵⁹² Prosapoin, saposin C, and peptides (prosaptides), encompassing the neurotrophic sequence located in the saposin C domain, also increased SM4s concentrations in primary and transformed Schwann cells and oligodendrocytes.²⁰¹

When growth factors of tyrosine kinase type, transforming growth factor α (TGF-α) (0-50 ng/ml²⁹⁰),²⁵ epidermal growth factor (EGF),²⁹⁰ and hepatic growth factor (HGF, 5-50 ng/ml),²⁹¹ were supplemented to the medium of the culture of SMKT-R3 cells, the sulfotransferase activity was increased markedly (approx. 300%). The incorporation of [³⁵S]sulfate into MDCK cells increased by renewal of the medium with 10% serum supplemented by hyperosmolar NaCl.⁴²⁶ On the other hand, supplementation of hypertonic NaCl after 2 days of serum depletion resulted in an increase of sulfotransferase corresponding to approx. 50% of that with the simultaneous addition of serum.^{231,443} Both of these increases were canceled by cycloheximide, an inhibitor of protein synthesis, suggesting that the synthesis of some proteins was responsible for the increased synthesis of sulfatides.^{291,426} On the contrary, removal of serum from the culture of mouse oligodendroglia cells resulted in approx. 2-fold enhancement of the sulfatide levels and

$\text{H}_2[^{35}\text{S}]\text{O}_4$ incorporation into SM4s within 24 hr with concomitant increase of sulfotransferase activity.²⁵⁰ Tyrosine kinase inhibitors, genistein and tyrphostin 51, reduced the enhancement of the sulfotransferase activity by EGF in a dose- and time-dependent manner.²⁵ Polycystic kidney disease is a disorder marked by aberrant renal tubular epithelial cell proliferation and transport abnormalities in *cpk/cpk* mice.⁹³ GlcCer, LacCer, and GM3 (V.C.1) displayed a large increase in the kidney of 3-week-old *cpk/cpk* mice, while SM4s and Cer concentrations decreased with concomitant decrease of GalCer sulfotransferase.

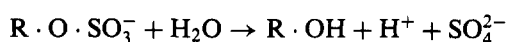
3. Sulfatide Modulation of Enzyme Activities

SM4s increased the activity of phospholipase C on the dilauroylphosphatidylcholine monolayer at various surface pressures³⁸ and activated protein kinase C to the same extent as phosphatidylserine did with phorbol esters.¹²⁹ $\text{HSO}_3\text{-Chol}$ activated η -, ϵ -, and ξ -isoforms of protein kinase C *in vitro* to an even greater extent.^{92,220} On the other hand, SM4s and $\text{HSO}_3\text{-Chol}$, as well as some detergents inhibited the activity of phosphatidylinositol-3-kinase.⁶³³ Exogenously added SM4s and anti-L-selectin antibodies enhanced tyrosine phosphorylation of MAP kinase in neutrophils.⁶²² The tyrosine kinase inhibitor, genistein, blocked the transient increase in intracellular Ca^{2+} and oxidative burst induced by SM4s, suggesting that this tyrosine phosphorylation serves to mediate signal transduction. Chymotrypsin-like activity of multicatalytic proteinase purified from human erythrocytes was selectively activated 2.5–3.5-fold by sulfated amphiphiles (SM4s, SM3, and SDS) but not by neutral glycolipids, gangliosides, and $\text{HSO}_3\text{-Chol}$. Heparin also activated the trypsin-like activity 2.5-fold, while other mucopolysaccharides did not.⁴³⁸

Sulfatides (SM4s, SM4g, 5 μM), and gangliosides (50 μM) inhibited the activity of DNA polymerase α , but acidic phospholipids did not.^{439,520} This inhibition of the enzyme activity by acidic glycolipids was suppressed by nonionic detergents suggesting that these acidic glycolipids, in the form of micelles, exerted influence on the enzyme activity by behaving as a polyanionic macromolecule.⁴³⁹ Human spleen lysosomal enzymes, glucocerebrosidases, were activated by acidic lipids (0.3–4.7 mM) including phosphatidylserine, a ganglioside, SM4s, synthetic monoacylglycerol sulfates and diacylglycerol sulfates.¹⁴³

C. Cerebroside Sulfatase and Saposins

Arylsulfatase A (EC 3.1.6.1) has been detected in most prokaryotes, plants, and the lysosome of animal tissues.¹⁰⁶ Only GalCer sulfatase (EC 3.1.6.8), which is the same enzyme as arylsulfatase A (EC 3.1.6.1), is responsible for cleavage of the sulfate esters of sulfatides. Sulfatides are not hydrolyzed by the other lysosomal enzymes arylsulfatase B (EC 3.1.6.12), and glucosamine-6-sulfatase which desulfate glycosaminoglycans. The third arylsulfatase, arylsulfatase C (EC 3.1.6.2), is in endoplasmic reticulum and desulfates steroid sulfates.⁵⁰² Arylsulfatase A catalyzes the cleavage of O—S bond similar to the acid hydrolysis of sulfoconjugates.⁴⁸²



Treatment of glucose 3-sulfate with arylsulfatase A in H_2^{18}O resulted in the formation of $\text{S}^{18}\text{O}^{16}\text{O}_3^{2-}$, supporting the cleavage of the O—S bond.⁶⁰¹

Arylsulfatases from alimentary tract of *Helix pomatia* were used for enzymatic desulfation of sulfatides immobilized on a 96-well plate.^{139,204} SM4s and SM4g were desulfated by an arylsulfatase fraction from the hepatopancreas of a mollusc *Charonia lampas*. The sulfate ester of these sulfatides was, however, scarcely hydrolyzed by 'glycosulfatase' I and II (EC 3.1.6.3), purified from the same source, which had been known to hydrolyze the sulfate ester linkage at C6 of glucose.¹⁸⁹ The acidic forms of the arylsulfatase from

various invertebrates possessed SM4s sulfohydrolase activities,³⁸⁹ whereas the arylsulfatase A fraction from ox liver showed 'glycosulfatase' activity.⁴⁸² The mRNA of arylsulfatase of sea urchin embryos reached a peak during development between the 32-cell stage and the prism stage.⁶⁶⁰

Mammalian arylsulfatase A hydrolyzes sulfate esters of sulfatides including SM4s,³⁶¹ SM4g,^{119, 134, 135, 641} lyso-SM4s (HSO₃-3GalSph),⁹⁹ ascorbic acid 2-sulfate, and tyrosine-*O*-sulfate¹⁰⁶ but not SM4s-6,³⁷⁴ SMUnLc₄Cer and SMUnLc₆Cer.⁶⁸ The human arylsulfatase A gene was about 3.2 kb long and contained eight exons.^{140, 313} Multiple transcription initiation sites were located between nucleotides -367 and -387 of this housekeeping gene, although its expression remained constant with EGF, TGF α or HGF treatment.²⁹¹ The genes coding for human arylsulfatase A, human steroid sulfatase, human glucosamine-6-sulfatase, and an arylsulfatase from sea urchin contained substantial amounts of homologous DNA sequences and probably originated from a common ancestral gene.⁵⁰²

Sphingolipid activator proteins (sap) or *saposins* are water-soluble, and heat-stable glycoproteins of about 1 kDa (80 amino acids) which interact with sphingolipids with oligosaccharide chains up to a length of three hexoses.^{128, 393, 492} The cDNA of prosaposin gene, the presumed 'housekeeping' gene located at chromosome 10q21 in humans, codes for a sequence of 524 amino acid residues in four domains (A, B, C and D).⁵⁴⁰ Probably alternative splicing of mRNA³³⁰ and post-translational proteolytic processing¹⁹⁶ generate four small, homologous proteins. *In situ* hybridization showed that prosaposin mRNA was expressed differentially in cell types and at developmental stages, and synthesized prosaposin was targeted via the mannose 6-phosphate receptor system to the lysosome or released into body fluids including serum, milk and semen.⁵⁴⁰ Saposins and saposin-like proteins contain three intradomain disulfide linkages, which create a common structural framework upon which amino acids in four amphipathic α helices can carry out diverse functions including transport of glycolipids.³⁹³

Saposin B (sap-B or SAP-1), a homodimeric glycoprotein, has a relatively wide specificity and interacts with GalCer, SM4s, sphingomyelin, phosphatidylserine, Gb₃Cer, Gg₄Cer, Ga₂Cer, as well as gangliosides GM1, GM2 and GM3⁵³¹ (Table 2). However, SM4s, but not Gb₃Cer and GM1 ganglioside, was accumulated in sap-B deficiency, suggesting that the physiological role of sap-B is solubilization of sulfatides.^{393, 508} Deglycosylated sap-B interacted with SM4s and GM1 ganglioside identical to native sap-B and stimulated the enzymatic hydrolysis of SM4s by arylsulfatase A to the same extent as native sap-B.²⁰³ Another saposin, glucosylceramidase-activator sap-C (SAP-2), and two additional potential activator proteins (sap-A and sap-D) are derived from a common precursor prosaposin by proteolytic processing. Sap-B, localized on lysosomal membrane, extracts a single sulfatide molecule from micelles or membranes probably behaving as a physiological detergent and binds it as a water-soluble 1:1 complex.¹²⁸ Deacylation of SM4s and SM4g (lyso-derivatives) or substitution of the fatty acid with acetic acid resulted in hydrolysis of the sulfate without a saposin probably because they form small micelles.^{115, 616}

Sulfatides are accumulated in metachromatic leukodystrophy (MLD) patients' myelin, liver, gallbladder, pancreatic islet cells, anterior pituitary, adrenal cortex (Table 3), and sweat glands.³⁰² MLD can be caused by mutations in two different genes, the arylsulfatase A⁴³⁶ and the prosaposin genes.¹⁹⁷ Patients with a complete defect of the prosaposin gene (due to a mutation in the initiation codon) exhibited complex biochemical abnormalities.¹⁹⁷ Mutant mice homozygous for an inactivated prosaposin gene exhibited two distinct clinical phenotypes, neonatally fatal and later-onset, with accumulation of glycolipids including SM4s.¹³⁰ X-linked ichthyosis is a genetic defect of the X chromosome with steroid sulfatase deficiency resulting in increased HSO₃-Chol in serum and stratum corneum giving rise to shedding of large scales containing a 5-fold increased HSO₃-Chol.^{630, 661} Multiple sulfatase deficiency (MSD) is an inborn error of metabolism with deficiency of arylsulfatase A, B, and C accompanied with accumulation of SM4s, SM3, HSO₃-Chol and heparan sulfate in the central nervous system, kidney and liver.^{502, 661}

The co- or posttranslational conversion of a cysteine (Cys69 and Cys91 in arylsulfatase A, and B respectively) to 2-amino-3-oxopropionic acid appeared to be required for generating catalytically active sulfatases.⁵⁰²

GM2 activator (SAP-3) stimulates the release of the terminal GalNAc from GM2, Gg₃Cer, SM2a,²³⁰ and Gb₄Cer by β -hexosaminidase A by binding with the substrate.¹⁶⁴ Microbial activator of endoglycoceramidase did not catalyze hydrolysis of SM4s or cerebroside.²⁴⁰

VII. INTERACTION WITH BIOMOLECULES

Sulfatides interact with various biomolecules specifically participating in cell adhesion, differentiation, and signal transduction.⁶¹¹

A. Electrostatic Interactions

As early as 1960, the high-affinity interaction of various neurotransmitters to SM4s was reported.^{152, 153} A particularly strong stereospecific and electrostatic bond can be formed between the protonated nitrogen of opioids and the sulfate group of SM4s³⁵⁶ or anionic phospholipids.⁴ Azure A (2×10^{-7} M) inhibited 85% of the specific electrostatic interaction of [³H]morphine and [³H]naloxone to synaptosomal membrane.³⁵⁶ Generally, proteins that interact with heparin also interact with fucoidan, dextran sulfate or sulfatides. The positively charged sequence (KKNKED) in L-selectin interacted with acidic phospholipids and fucoidan³⁶³ and sequences rich in basic amino acids are contained in the domains responsible to SM4s binding in thrombospondin and laminin. Tyrosine sulfate residues may also be able to cooperate with immediately adjacent sialylated oligosaccharides to generate P-selectin recognition of PSGL-1 (P-selectin glycoprotein ligand-1).³⁴⁵ In this instance the clustered patch may not be purely made up of saccharides, but would be a composite 'clustered anionic patch'.⁸⁰ From this point of view, most of the affinity of sulfated glycoconjugates with antibodies or 'specific' binding proteins⁶¹ may involve, primarily, ionic interactions.

In situ hybridization showed the colocalization of the mRNA of ceramide galactosyl transferase and myelin basic protein (MBP).⁵¹⁰ The developmental expression pattern was also similar to the myelination profile.⁵³³ It has been proposed that SM4s, together with other amphipathic compounds, interacts with hydrophobicity⁵²⁸ or electrostatic force⁴⁹³ with MBP, which contains predominantly hydrophobic and basic amino acids, stabilizing the compact structure of myelin. However, the physiological significance of this phenomenon has been questioned because MBP is an extrinsic membrane protein associated to the cytoplasmic leaflet of the bilayer and a direct interaction with SM4s at the exoplasmic leaflet is unlikely.⁶¹⁸ Due to its highly basic amino acid composition, Tamm-Horsfall glycoprotein (T-H) of the kidney, a glycosylphosphatidylinositol anchored protein, has been a potential candidate for ionic interaction with SM4s. Studies by using human T-H⁶⁴⁸ and polyclonal antibodies specific to SM4s⁶⁴⁹ showed that T-H and SM4s had a strictly superimposable localization on kidney tissue sections, that is at the luminal membrane of the thick ascending limb of the loop of Henle and the initial portion of the distal convoluted tubule beyond the macula densa. In view of the presence of 3-O-sulfated galactose in T-H, however, attention has to be paid to the possibility that antibodies recognizing 3-O-sulfated galactosyl residues may also cross-react with T-H.¹⁸⁵ Serum amyloid-P protein has been known to interact with a variety of anionic amphiphiles including hexose phosphates in the presence of Ca²⁺. Amyloid-P interacted preferentially with sulfatides with sulfate groups on the terminal Gal or GalNAc residues, whereas there was considerably weaker interaction with SM2a that has a sulfated penultimate Gal, and only trace binding to SM1a with an internal sulfated Gal.³⁵⁷

B. Proteins of the Extracellular Matrix and Blood

The extracellular matrix proteins including laminins, fibronectins and collagens, as well as plasma proteins including thrombospondins, have multidomain structures containing interaction sites for various types of anionic glycoconjugates including heparin and sulfatides⁶¹¹ (Table 2). For laminin interaction with sulfatides including SM4s, SM4g, SM3,^{294, 471} SMUnLc₄Cer and SMUnLc₆Cer,^{249, 381, 503} a sulfate group at the non-reducing terminus is necessary. Eight genetically distinct laminin chains (α 1, α 2, α 3, α 4, α 5, β 1, β 2, β 3, γ 1, γ 2) and ten different heterotrimeric assembly forms, laminins -1 to -10, are known so far. The major isoform, laminin-1, has a distinct affinity for heparin and SM4s mediated by fragment E3, which consists of the G4 and G5 domains of the carboxyl terminus of α 1 chain with two X-B-B-X-B-X, and three B-X-B-X-B-X sequences.^{583, 591} Urea treatment or reduction and alkylation of the fragments abolished SM4s binding⁵⁸³ but not the binding to heparin and HNK-1 neoglycoproteins¹⁶³ suggesting that sulfatide interaction specifically requires an intact three-dimensional structure. In contrast to SM4s and heparin, glycoconjugates with SMUnLc₄ appeared to bind to E-8 fragment of the domain G2 of laminin-1.¹⁶³

Thrombospondins (TSP) are large, homotrimeric glycoproteins.⁴³ TSP1 and TSP2 contain a globular NH₂-terminal domain, a procollagen homology domain, type I (TSP or properdin), and type II (EGF-like), and type III (Ca²⁺-binding) repeats, and a carboxyl terminal domain.⁴³ Thrombospondins interacted on solid phase with SM4s, SM3, SM4g,⁴³ SM2a, SMGb₄Cer,⁴⁰⁰ and SMGb₅Cer³⁹⁹ from human kidney. Human properdin, and H-Factor, the regulators of the alternative pathway of complement activation, interacted with SM4g, SM4s, SM3, SB2, and SM2a.²⁴⁶ A consensus sequence for the interaction with heparin and SM4s was proposed as Cys-Ser-Val-Thr-Cys-Gly-X-Gly-X-X-X-Arg-X-Arg (or -Lys) (= CSVTCGXGXXXR).²⁰⁷⁻²⁰⁹ TSP type I repeats, properdin (a serum protein),¹⁹⁸ antistasin (a leech salivary anticoagulant), and *Herpes simplex* I contain these motives. The coat protein,^{62, 390, 446} region II-plus of the³⁹⁰ circumsporozoite (CS) proteins, and thrombospondin-related anonymous protein (TRAP) of malaria parasites also contain an amino acid motif based around the sequence CSVTCG.⁵⁸⁵ This amino acid motif has been considered to confer on the CS protein the ability to bind specifically to host sulfated glycoconjugates and to the surface proteoglycans of hepatocytes or HepG2 cells.^{390, 522} CSVTCG has also been suggested to be an important determinant in interaction with other mammalian cells,⁴³ although it was shown recently that the downstream positively-charged residues without CSVTCG or WSPWS segment served as the potent ligand.⁶⁶⁷

Heparin and sulfated glycolipids had a stronger affinity to WSXW sequence in peptides obtained from thrombospondin, laminin, and apolipoprotein.¹⁵⁷ The heparin-binding domains of TSP1 and 2 were subsequently shown to contain BBXB sequences (e.g. KRFK), where B is the probability of a basic residue and X is a hydrophobic residue, and the type I repeat contains WSXW in addition to CSVTCG. However, TSP type I repeat 5 of properdin contains neither of the electroneutral sequences, the WSXW or the CSVTCG motif, while rich in basic amino acids.¹⁹⁸ XBBXB and XBBBXXBX were determined as the consensus sequences for glycosaminoglycan recognition in 49 regions in 21 proteins.⁶¹

It has been proposed that soluble laminins in the culture medium mediate adhesion of some melanoma cell lines to sulfatides (SM4s or SM4g, 25 fmol/mm²) on the plastic plate.⁴⁷² The interaction of ¹²⁵I-labeled laminin or thrombospondin to A2058 melanoma cells was inhibited by thrombospondin peptides containing WSXW sequence.¹⁵⁷ The peptide KRFKQDGGWSHWSPWSS inhibited binding by approx. 80–95%. Many other tumor cell lines have been found to attach on SM4s substrates using endogenous proteins interacting with SM4s (Table 2).

C. Selectins

All three (E-, P-, and L-) selectins possess the *N*-terminal carbohydrate-recognition domain homologous to C-lectins (Ca^{2+} -dependent vertebrate lectins) followed by an EGF-like domain, and recognize sialyl-Lewis x and its isomer sialyl-Lewis a.⁶¹³ L- and P-selectins also interact with diverse natural sulfoglycolipids and sulfated neoglycolipids in Ca^{2+} -dependent^{23, 112, 301, 467} or independent^{153, 363, 417, 550} modes (Table 2).

A soluble fusion protein of rat L-selectin-IgG, which contained the lectin and EGF domains,⁵⁵⁰ was assayed for the ligands by solid-phase binding assay. L-selectin-IgG interacted with sulfated glycolipids in the concentrations of 30–100 pmol/well. A synthetic SM4s analogue, $\text{HSO}_3\text{-3Gal-B30}$ (branched chain, C30 alcohol) interacted more preferentially with the chimera protein than $\text{HSO}_3\text{-2Gal-B30}$, or $\text{HSO}_3\text{-6Gal-B30}$. Also, $(\text{HSO}_3)_2\text{-3,6Gal-B30}$ and $(\text{HSO}_3)_3\text{-3,4,6Gal-B30}$ were more reactive than $\text{HSO}_3\text{-3Gal-B30}$, indicating that the interaction of L-selectin with its sulfated sugar ligands is position-specific but depends on the number of the sulfate group supporting the role of anion clusters for the interaction.^{393, 550} Soluble chimeric proteins with various domains switched between E- and L-selectins showed that the *N*-terminal lectin domain bound specifically to SM4s,³⁰¹ while EGF domain or the consensus repeats of E- and L-selectins had no influence on the specificity to carbohydrates.³⁰¹ In contrast, conservative substitution of Ser-124 and 128 in EGF domain residue was able to alter E-selectin binding such that it adhered to SM4s and heparin, and reduced P-selectin adherence to these ligands.⁴⁶⁷ The binding site for acidic phospholipids is located close to the lectin domain and contained a BBXBXX (i.e. KKNKED) sequence.³⁶³

By the use of rat L-selectin-IgG chimera, ligands for L-selectin were located in high endothelial cells in lymph nodes, the white matter, neurons, cerebellar Purkinje cells, and choroid plexus of the central nervous system, as well as in a straight portion of distal tubules and capillary blood vessels in medulla and pelvis of the kidney.⁵⁸¹ The staining on the white matter of the cerebellum and distal tubules of the kidney was completely abolished by treatment with organic solvents (C/M, 1:1), indicating that the major ligands for L-selectin-IgG chimera in these organs were glycolipids. However, the true, high-affinity ligand at the high-endothelial venules (HEV) may be diverse mucins with sialylated, sulfated, fucosylated lactosamine-type *O*-linked.⁸⁰

Using computer modeling techniques, site-specific mutagenesis, and ligand and cell binding assays, it was shown that P-selectin lectin domains bind to myeloid cells, SM4s and sialyl Lewis x oligosaccharide via an overlapping, but not identical set of residues located in a shallow cleft of the lectin domain that is similar to the mannose-binding protein saccharide-binding site proximal to a functional calcium binding site.^{23, 467} When Ala77 was substituted with lysine, P-selectin-carbohydrate binding specificity changed from sialyl Lewis x to oligomannose supporting the structural analogy with the lectin domain of rat mannose-binding protein.⁴⁶⁷

D. Microorganisms

Sulfatides interact with several viri including HIV (human immunodeficiency virus), cytomegalovirus⁴³⁵ and influenza virus.⁵⁴⁹ The ability of the surface envelope glycoproteins, gp120⁶⁵⁶ and GP41⁴⁶⁰ from different strains of HIV to interact with MAG, sulfatides or acidic phospholipids has been regarded as an important determinant in the development of neuropathy of AIDS, although CD4 molecule is the primary ligand for HIV.⁶⁶³ GalCer and SM4s were stained with ¹²⁵I-labeled gp120 but GlcCer, GM1, GD1a and neutral glycolipids from human erythrocytes were not.³⁷ In another experiment using nitrocellulose paper, however, gp120 bound only to SM4s and not to GalCer.⁶⁶² The interaction with a peptide of V3 domain of gp120 suggested that the GalCer and SM4s binding region may reside on the sequence of the V3 loop peptide.⁴⁰⁶ Fully sulfated Glc, Gal and lactose linked to lipophilic moieties,²¹⁹ the sulfated polysaccharides (e.g. curdlan sulfate),²⁷⁶ SQ-A₂Gro, as well as medium-molecular-weight sulfated alkyl

oligosaccharides¹⁴⁸ showed anti-HIV activities *in vitro*. By sulfated alkyl laminara- and malto-oligosaccharides (0.4 $\mu\text{g}/\text{ml}$) a 50% decrease of HIV-induced cytopathic effects was observed.^{276, 277} The infection of human T cells with the virus was markedly inhibited by treatment with the sulfated gangliosides at a concentration of 10 $\mu\text{g}/\text{ml}$, while the non-sulfated gangliosides had only weak antiviral activities.¹⁶⁸

The cell adhesion to sulfatides immobilized on TLC plates⁵⁰⁶ demonstrated that many prokaryotes use glycoconjugates as the receptors.²⁶³ S-fimbriated *E. coli* strains, the pathogens of neonatal meningitis, interacted with SM4s and SM4g, but the transformants which lack the *sfaA* gene showed no interaction with the glycolipids.⁴⁵⁴ *M. pneumoniae* bound avidly to pulmonary tissues as well as the WiDr human colon adenocarcinoma cell line, and also interacted on TLC plates with SM4s, SM4g and SM3,^{314, 470} while *M. hyopneumoniae* interacted with SM4s, GM3 and Gb₄Cer.⁶⁵² *Mycoplasma pulmonis* is associated with male infertility in humans, cattle and rodents probably by binding specifically to SM4g.³⁵⁴ The topology of mycoplasma interaction with rat sperm was consistent with the known topology of sperm SM4g.³⁵⁵ Moreover, dextran sulfate and heparin^{354, 444} inhibited the interaction between Mycoplasmas and cells or tissues that express SM4s or SM4g.

SM4s showed strong tetanus-toxin binding on HPTLC plates, with standard GD1b and GT1b as positive controls.¹⁴⁴ The absence of tetanus-toxin-binding gangliosides in SCLC (small-cell-lung-cancer) cell lines suggested that sulfatides might be responsible for the reaction of SCLC cells with the toxin. Sulfated glycolipids may serve as the mucosal receptor for colonization of *Helicobacter pylori*, which colonizes only in the gastric epithelium and is associated with gastritis and peptic ulcer.⁴¹² Viable *H. pylori* showed a strong interaction with SM4s, SM3 and GM3 by TLC-immunostaining.^{259, 404} Virulent strains of *B. pertussis*, the human respiratory pathogens, bound specifically to asialo GM1 and SM4s, suggesting that the *B. pertussis* adhesin FHA may utilize sulfated glycolipids and proteoglycans to initiate infection.^{48, 172}

VIII. PERSPECTIVES

The status of the present efforts to establish functions specific to sulfatides is more or less analogous to the trends in sulfated polysaccharides of the connective tissue described by Comper⁷⁶ and in sulfated glycoconjugates reviewed by Varki.⁶¹¹ The functional roles assigned to the anionic proteoglycans and polysaccharides of cell surface materials of prokaryotes or extracellular matrix of multicellular organisms contain a great insight applicable to sulfated glycolipids. The partial informational specificity of sulfated glycoconjugates has been discussed in terms of evolutionary flexibility of the extracellular matrix.⁷⁶ Matthews indicated, as early as 1975, that the response to environmental influences is an essential property of supporting tissues and that this characteristic is derived largely from the capacity for *fine modulation* of the structure of its constituent macromolecules.³⁶⁷ To name the major tools for the gross modulation against hyperosmolality, they are transporters such as Na⁺, K⁺-ATPase, and organic osmolytes including glucitol, glycerol, *myo*-inositol, betain, and glycerylphosphorylcholine.^{358, 410} As much evidence has suggested, the multi-step fine modulation may be accomplished by anionic polysaccharides and sulfated glycolipids forming multilayers on the cell surface.⁴²⁶

The recent results obtained by inactivation of the genes of sugar transferases in knockout mice were quite unexpected and some of them appeared to suggest the redundancy of anionic glycolipids in the biosphere. For instance, the mice with the inactivated gene of β -GalNAc transferase that transfers β -GalNAc at position 4 of the Gal in LacCer survived with unexpectedly slight growth retardation. In the above cases, GM3 and GD3¹³² replaced 'conventional' ganglio-series gangliosides in agreement with the neutral theory of glycolipid structure,²³⁸ and allometric principle.³⁹⁸ The major part of the function of GalCer and sulfatides in myelin appeared to be partly replaceable by other amphiphiles as shown by the knockout mice with inactivated ceramide galactosyltransferase.⁷⁵ However, the serious defect of these mice suggested that the replacement is not adequate

for the full expression of the myelin function. The biological molecules in the biosphere have either been evolutionally selected for their ability to adapt the environment or at least partly resulted from neutral diversification and random drift.²³⁸

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