New era of research on cancer-associated glycosphingolipids

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Cancer-associated glycosphingolipids have been used as markers for diagnosis and targets for immunotherapy of malignant tumors. Recent progress in the analysis of their implications in the malignant properties of cancer cells revealed that cancer-associated glycosphingolipids are not only tumor markers, but also functional molecules regulating various signals introduced by membrane microdomains, lipid rafts. In particular, a novel approach, enzyme-mediated activation of radical sources combined with mass spectrometry, has enabled us to clarify the mechanisms by which cancer-associated glycosphingolipids regulate cell signals based on the interaction with membrane molecules and formation of molecular complexes on the cell surface. Novel findings obtained from these approaches are now providing us with insights into the development of new anticancer therapies targeting membrane molecular complexes consisting of cancer-associated glycolipids and their associated membrane molecules. Thus, a new era of cancer-associated glycosphingolipids has now begun.

KEYWORDS
cancer marker, cancer-associated antigen, cluster, enzyme-mediated activation of radical sources, ganglioside, glycosphingolipid, lipid raft

1 | GLYCOSPHINGOLIPIDS AS FUNCTIONAL MOLECULES ON THE CELL SURFACE

Cancer-associated glycosphingolipids have been considered as tumor markers,1 and used as diagnostic markers and targets of cancer treatment.2 There have been a number of reports on the specific expression of various glycosphingolipids in individual cancers. These include ganglioside GD3 in melanomas3 and T-cell lymphoblastic leukemia,4,5 GD2 in neuroblastomas,6 osteosarcomas,7,8 small cell lung cancers9,10 and breast cancers,11 and globotriaosylceramide in Burkitt's lymphomas.12 Sialyl Lewis A has also been used as a tumor marker of pancreatic, gastrointestinal,13 and other epithelial cancers;14 this structure is carried on both glycosphingolipids and mucins.2

Remodeling of carbohydrate structures on proteins and lipids by artificial manipulation of glycosyltransferase genes has enabled us to further analyze the roles of these cancer-associated glycosphingolipids in cancer cells in addition to mere cancer markers.15 Generally speaking, disialyl glycosphingolipids confer malignant properties in various cancer systems. In particular, tandem-repeated sialic acid-structures such as those in GD3 and GD2 enhanced the malignant
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Properties of cancer cells, such as cell proliferation, cell invasion and migration, cell adhesion, and metastasis.\textsuperscript{15-17} In contrast, monosialyl gangliosides, such as GM1, GM3, and GM2 often suppress malignant properties of various cancer cells.\textsuperscript{18-20} Therefore, GD3 synthase is a key enzyme determining cell phenotypes based on the expression patterns of gangliosides, as shown in Figure 1.

\section*{2 | ENIGMA IN THE ACTION MECHANISMS FOR GLYCOSPHINGOLIPIDS HAS BEEN GETTING DISCLOSED}

How these glycosphingolipids affect the phenotypes and behaviors of cells in various manners has also been rigorously analyzed, and molecular mechanisms by which glycosphingolipids modify cell signaling based on the interaction with other molecules existing on the same membrane (cis-binding) or on the different cell surface (trans-binding) have been reported.\textsuperscript{21} Although soluble and/or cell surface ligand molecules that gain access from outside cells are important as trans-interacting molecules with glycosphingolipids,\textsuperscript{22} knowledge on cis-interaction on the cell surface has recently advanced. In this group, growth factor receptors and adhesion receptors such as the integrin family are mainly defined as cis-interacting molecules with glycosphingolipids on the cell surface, leading to the modification of cell signals mediated by those receptors.\textsuperscript{16,17} Figure 2 shows a representative example of signal convergence in melanomas. Activation signals mediated by growth factor receptors are merged with adhesion signals by integrins, resulting in the convergence of signals, leading to the markedly enhanced promotion of downstream molecules such as AKT, p130Cas, and paxillin, and increased malignant properties of melanoma cells.\textsuperscript{16,17}

Results on increased adhesion signals by the integrin-FAK axis are presented in Figure 2B.

The recent development of a new approach to define interacting molecules with cancer-associated glycosphingolipids on the surface of living cells has markedly promoted understanding of these interactions, namely EMARS/MS, developed by Honke and Kotani.\textsuperscript{23} To analyze molecular clustering on the cell surface, co-immunoprecipitation has been used. Cross-linking analysis, including photoaffinity labeling, with the aid of chemical cross-linkers has also been used, leading to partly successful results with a restricted range of clustered molecules. Morphological visualization with fluorescence microscopy or electron microscopy has been used to obtain useful findings. However, all target molecules of these approaches need to be known at the starting points. Compared with these current methods, EMARS is a very powerful approach. The advantages of this approach can be summarized as follows: (a) using living cells to analyze events on the cell surface, corresponding to the size of microdomains; (b) no need for special equipment, and easy to carry out; and (c) applicable for comprehensive analysis of clustering molecules with particular targets, enabling the identification of molecular profiles including those of unknown molecules.\textsuperscript{24}

Using "cancer-specific" mAbs, we have identified candidate molecules that might physically and functionally interact with cancer-associated glycosphingolipids on the cell surface within ca. 300 nm, as shown in Figure 3.\textsuperscript{23,24} Thus, how glycosphingolipids are...
expressed on the outer layer of the cell membrane and exert their roles in the regulation of cell phenotypes is now being clarified.

Consequently, EMARS/MS could be a very powerful approach to identify cooperating molecules with cancer-associated glycosphingolipids. The features of molecular assembly depending on the changes in cell states can also be defined. Furthermore, we can investigate the roles of molecular complexes consisting of cancer-associated glycosphingolipids and EMARS-defined molecules. Downstream molecules activated by these molecular complexes have also been reported, as described in the following section.

3 | CANCER-ASSOCIATED GLYCOSPHINGOLIPIDS PLAY ROLES THROUGH COMPLEX FORMATION WITH MEMBRANE MOLECULES

As described above, ganglioside GD3 has been considered as a melanoma-specific glycolipid antigen,25 and been used as a target of Ab therapy and/or immune cell-mediated therapy for malignant melanomas.26,27 Roles of GD3 in melanomas have been reported based on experiments using cells transfected with GD3 synthase cDNA, showing its effects on the activation of signaling molecules located downstream of growth factor receptors28 or adhesion receptors, integrins.27 An EMARS/MS analysis (Figure 3) with melanoma cells using anti-GD3 mAb revealed various membrane molecules as candidate molecules that associate with GD3.29 Among them, Neogenin-1 was defined as a GD3-associating molecule and also as a GEM/raft-residing molecule exclusively in GD3+ cells.30 Neogenin-1 was shown to be involved in enhancement of the malignant properties of melanomas such as increased cell growth, invasion, and migration, indicating that Neogenin-1 might be an effector molecule exerting the effects of melanoma-specific ganglioside, GD3 (Figure 4A). Consequently, GD3 expression resulted in the shift of Neogenin-1 (Figure 4B) but also of γ-secretase to lipid rafts from the nonraft compartment, leading to increased cleavage of the intracytoplasmic domain of Neogenin-1. Neogenin intracytoplasmic domain promoted expression of multiple genes such as GPR126, STXB5, MMP16, SPATA31A1, and S6K, as identified by ChIP-sequencing, possibly playing central roles in enhancing the malignant properties of melanomas30 (Figure 4A).

Using a replication-competent avian leukemia virus splice acceptor system,31 we analyzed murine gliomas with a focus on the expression of gangliosides on the cell surface.32 Astrocytes transfected with
PDGFB showed some levels of GD3 expression, and GD3+ and GD3– astrocytes were separated, enabling us to analyze the effects of GD3/GD2 expression on astrocytes. Thus, it was shown that GD3/GD2 expression increased cell growth, invasion, and migration. To clarify the mechanisms by which GD3/GD2 enhance the malignant properties of gliomas, EMARS/MS was carried out, and PDGFRα was defined as a GD3-associating molecule in murine gliomas. Only GD3 formed a molecular complex with PDGFRα, and the Src family Yes was also included in the complex, leading to the formation of a ternary molecular complex adjacent to the cell membrane. This complex activated paxillin, resulting in the increased invasiveness that is the most notable feature of gliomas (Figure 5A).

Application of the EMARS/MS approach resulted in the detection of GD3-associating molecules in human melanoma and murine glioma cells, both of which are membrane receptors that regulate cell growth/differentiation. In contrast, a similar approach led to the detection of an amino acid transporter in SCLC. Among human lung cancers, SCLC show high frequencies of metastasis and recurrence. Therefore, novel Ab therapy targeting SCLC-specific antigens has been long expected. As a result of EMARS/MS using anti-GD2 mAb and GD2 synthase-transfectants of SK-LC-17, approximately 20 molecules were identified as candidates of GD2-associating molecules. These molecules consisted of CD109, uPAR, EphA2, and CD44. Among them, ASCT2, a glutamine transporter, was most clearly localized in lipid rafts of GD2+ cells. Then we examined roles of ASCT2 in SCLC cells. The expression of ASCT2 in SCLC cells resulted in increased proliferation and invasion. ASCT2 colocalized with GD2 in GEM/rafts, and they were associated with each other, as shown by multiple approaches such as co-immunoprecipitation and the proximity ligation assay. Consequently, ASCT2 co-operated with GD2 and increased the uptake of glutamine, and increased activation levels of downstream molecules of mTOR1, such as p-p70 S6K1 and p-S6 (Figure 5B).
Neogenin was defined as a ganglioside GD3-associating molecule in melanoma cells by enzyme-mediated activation of radical sources/mass spectrometry. A, Under GD3 expression, Neogenin colocalized with GD3 and γ-secretase in lipid rafts, resulting in increased generation of Neogenin intracytoplasmic domain (NeICD). Eventually, NeICD promotes the expression of various genes and enhances function of GD3 (modified from Kaneko et al30 with permission). B, Floating pattern (intracellular localization) of Neogenin was analyzed by sucrose density gradient ultracentrifugation of Triton X-100 extracts from GD3+ (G5) and GD3− (V9) melanoma sublines. A shift of Neogenin to glycolipid-enriched microdomain (GEM)/rafts was clearly indicated by immunoblotting (ref. 30).

Action of ganglioside-associating molecules defined by enzyme-mediated activation of radical sources/mass spectrometry (EMARS/MS) analysis in murine gliomas and human small-cell lung cancer (SCLC) cells. A, Platelet-derived growth factor receptor-α (PDGFRα) was defined as a GD3-associating molecule in murine gliomas by the EMARS/MS approach. Specific association of GD3 with PDGFRα was identified by co-immunoprecipitation, and activated Yes was also found in the immune complex. Subsequently, it was shown that membrane molecular complexes consisting of GD3, PDGFRα and Yes promote glioma invasiveness by activating paxillin.32 B, By EMARS/MS, a glutamine transporter, ASCT2, was defined as a GD2-associating molecule in SCLC. ASCT2 cooperates with GD2, resulting in the increased activation levels of mTOR1 downstream molecules such as p70S6K1 and S6 in SCLC cells.35 These signals finally induce increased cell growth and migration.
GLYCOSPHINGOLIPIDS REGULATE ARCHITECTURE AND FUNCTION OF GEM/RAFTS

The majority of the cancer-associated glycosphingolipids mentioned above are considered to localize in GEM/rafts. Indeed, many of the EMARS/MS-defined molecules targeting glycosphingolipids were detected in GEM/rafts. Among GEM/raft-residing molecules, such as cholesterol, sphingomyelin, GPI-anchored proteins, and glycosphingolipids, the latter 2 are unique as they make up relatively uniform lipids and polymorphic nonlipid portions. The polymorphic portions, ie, proteins or carbohydrates, are exposed to the outside of the cell surface, interacting with outward ligands. Glycosphingolipids are particularly unique because they interact with outward ligands, but also with cis-residing molecules on the same cell membrane, playing roles by forming molecular complexes consisting of assembled glycolipids and membrane molecules.

From our experience, alteration of the glycosphingolipid composition in cells and tissues resulted in marked changes in the architecture and function of GEM/rafts in cultured cells, and also in experimental animals. Neo-expression of GD3 induced marked changes in the intracellular localization of integrins, Src-family kinases, and Neogenin in melanoma cells. A lack of gangliosides except GM3 resulted in the dispersion of GPI-anchored proteins, as well as raft marker proteins from GEM/rafts in mouse brain tissues. Thus, expression patterns of glycosphingolipids determine the features of GEM/rafts, ie, profiles of GEM/rafts and properties of GEM/rafts as a platform for signal regulation. Lipid rafts have been considered to take part in the complex protein-protein interactions for the generation of signal transduction. Lipid rafts can change size (from nanoscale to raft “phase”) and composition responding to intra- or extracellular stimuli, as shown in Figure 6. Therefore, how altered carbohydrates in cancer cells regulate the composition and function of GEM/rafts and cell signals, and whether individual cancer-associated glycosphingolipids form distinct molecular clusters and exert unique signaling, remain to be clarified in the future.

DEVELOPMENT OF NOVEL TECHNOLOGY ENABLES US TO FURTHER UNDERSTAND GLYCOSPHINGOLIPIDS

In addition to the EMARS/MS approach, development of novel technologies has brought about a new era of cancer-associated glycosphingolipids. Single molecule imaging with high resolution (millisecond scale) has made it possible to observe the spatiotemporal dynamics of molecular clustering on cell surfaces. It has become possible to observe processes of actual formation of molecular complexes on the cell surface. The results of these imaging experiments have been markedly changing the concept of GEM/rafts. Minimal units of nanoscale GEM/rafts might be dimer formation of glycosphingolipids with the same structure, whereas GPI-anchored proteins should play central roles in the solid and microscale membrane domains. Analysis of interactions between cancer-associated glycosphingolipids and interacting molecules with high-resolution single molecule imaging would reveal new mechanisms for the generation of “cancer-associated glyco-signals”. One more novel technique that has supported single molecule imaging is that of chemical synthesis technology of fluorescence-labeled glycosphingolipids. A new method to label glycosphingolipids developed by Komura et al is based on the incorporation of fluorescent dye in the carbohydrate portion of glycosphingolipids by maintaining their primary natures. This point is markedly different from the fluorescent ceramide analogue that was widely used in the trafficking, transfer, intracellular localization, and metabolism of glycosphingolipids. Modification of the ceramide moiety might generate confusing data, as lipid portions are essential for the intracellular trafficking and localization of glycosphingolipids. The new method has
enabled us to obtain reasonable results concerning the dynamics of glycosphingolipids.

For a long time, glycosphingolipids have been considered to be regulatory molecules, particularly in the nervous system, mainly because of their abundant expression in neuronal cells. However, it was reported that fine structures of glycosphingolipids are not essential for the existence of bio-organisms, but they are pivotal regulators of various cells and organs based on the regulation of cell signals in GEM/rafts. What should be emphasized here is that the cluster formation of glycosphingolipids with homopolymer mode could be critical for recognition by ligand molecules. Recently, heteropolymers have also been reported to sometimes be essential for interactions with cis- and trans-acting ligands such as lectins.

These molecular clusters in GEM/rafts should affect the assembly/dispersion of other membrane molecules in/from GEM/rafts, leading to the functional regulation of those membrane molecules. Furthermore, molecular complex formation in a vertical direction through the cell membrane is also crucial for the determination of cell fates.

Now, glycosphingolipids are emerging as regulators of membrane environments and cancer microenvironments, not just markers of cell differentiation and malignant transformation.

Based on novel findings on the functions and modes of action of cancer-associated glycosphingolipids, novel therapeutic trials have been carried out, or considered. For example, when a novel signaling pathway activated by cancer-associated glycosphingolipids is found, key molecules in the pathway could be targets of therapy. In fact, p130Cas and paxillin, which are activated in GD3+ melanomas, were examined as possible target molecules for siRNA therapy. Membrane proteins that are identified as partners of cancer-associated glycolipids might be better targets for treatment. Downstream signal molecules of these partner membrane molecules could be more specific for cancer cells. Furthermore, identification of new epitopes formed by molecular complexes consisting of cancer-associated glycolipids and their partners might be expected as new targets. If particular molecules in glycosylation machineries are considered to be cancer-specific, and to play key roles, they could also be new targets of cancer therapy.

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CONFLICT OF INTEREST

Authors declare no conflict of interest for this article.

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