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Siddabasave Gowda B. Gowda, Shu-Ping Hui *et al.* A facile method for monitoring sphingomyelin synthase activity in HeLa cells using liquid chromatography/mass spectrometry

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1. Introduction

Sphingolipids are a major class of lipids that, comprises various metabolites with diverse functions. Changes in the biological processes of sphingolipids are associated with metabolic syndromes, including obesity, dyslipidemia, insulin resistance, cardiovascular problems, and type 2 diabetes mellitus.¹ Sphingomyelin (SM) is a sphingolipid metabolite produced by the transfer of phosphorylcholine from phosphatidylcholine to ceramide (Cer) in a reaction catalyzed by sphingomyelin synthase (SMS).² Mammalian SMSs exist in three isoforms: SMS1, SMS2, and SMSr.³ SMS1 and SMS2 share 57% sequence



Punith M. Sundaraswamy,‡^a Yusuke Minami,‡^b Jayashankar Jayaprakash,‡^a Siddabasave Gowda B. Gowda, ^b *^{a,c} Hiroyuki Takatsu,^d Divyavani Gowda,^c Hye-Won Shin^d and Shu-Ping Hui*^c

Sphingomyelin synthase (SMS) is a sphingolipid-metabolizing enzyme involved in the *de novo* synthesis of sphingomyelin (SM) from ceramide (Cer). Recent studies have indicated that SMS is a key therapeutic target for metabolic diseases such as fatty liver, type 2 diabetes, atherosclerosis, and colorectal cancer. However, very few SMS inhibitors have been identified because of the limited sensitivity and selectivity of the current fluorescence-based screening assay. In this study, we developed a simple cell-based assay coupled with liquid chromatography/tandem mass spectrometry (LC-MS/MS) to screen for SMS inhibitors. HeLa cells stably expressing SMS1 or SMS2 were used for the screening. A non-fluorescent unnatural C6-Cer was used as a substrate for SMS to produce C6-SM. C6-Cer and C6-SM levels in the cells were monitored and quantified using LC-MS/MS. The activity of ginkgolic acid C15:1 (GA), a known SMS inhibitor, was measured. GA had half-maximal inhibitory concentrations of 5.5 µM and 3.6 µM for SMS1 and SMS2, respectively. To validate these findings, hSMS1 and hSMS2 proteins were optimized for molecular docking studies. *In silico* analyses were conducted to assess the interaction of GA with SMS1 and SMS2, and its binding affinity. This study offers an analytical approach for screening novel SMS inhibitors and provides *in silico* support for the experimental findings.

similarity and have identical enzyme activities⁴ but different subcellular locations.⁵ SM production is catalyzed by SMS1 in the Golgi apparatus and by SMS2 on the plasma membrane.^{2,5,6}

Recent, studies in mice have shown that SMS deficiency prevents the advancement of atherosclerosis,⁷ insulin resistance,^{8,9} and obesity,⁹ while *in vitro* inhibition of SMS attenuated endotoxin-mediated macrophage inflammation and the development of atherosclerosis.¹⁰ SM accumulation is commonly observed in metabolic diseases such as fatty liver disease,¹¹ colorectal cancer,¹² and coronary artery disease.¹³ Hence, SMS may serve as a therapeutic target for the development of novel treatments for metabolic diseases, and the discovery of SMS inhibitors is a promising strategy for achieving this goal.

A cell-based method coupled with high-performance liquid chromatography (HPLC) to detect fluorescent substrates and products is a widely used assay method to screen for SMS inhibitors.¹⁴ Natural inhibitors such as malabaricone C¹⁵ and ginkgolic acid (GA)¹⁶ have been discovered using this assay method. Despite the assay being rapid, its sensitivity is poor, interference peaks often occur in the HPLC chromatogram, and the results are not quantitative. LC-MS/MS has a higher

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^aGraduate School of Global Food Resources, Hokkaido University, Kita-9, Nishi-9, Kita-Ku, Sapporo 060-0809, Japan. E-mail: gowda@gfr.hokudai.ac.jp

^bGraduate School of Health Sciences, Hokkaido University, Kita-12, Nishi-5, Kita-ku, Sapporo 060-0812, Japan

^cFaculty of Health Sciences, Hokkaido University, Kita-12, Nishi-5, Kita-ku, Sapporo 060-0812, Japan. E-mail: keino@hs.hokudai.ac.jp; Tel: +81-11-706-3693

^dGraduate School of Pharmaceutical Sciences, Kyoto University, Kyoto 606-8501, Japan

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[‡]These authors contributed equally to this work.

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sensitivity and selectivity than HPLC. Efforts have been made to identify SMS inhibitors through virtual screening combined with an LC-MS/MS approach.^{4,17} Although this previous LC-MS/MS-based method allows the quantitation of SMS activity, it is time-consuming, involves complex steps, and does not provide *in silico* insights.⁴ Reports on improved methods to screen for SMS inhibitors using *in vitro* and *in silico* approaches are limited. In this study, we aimed to establish a cell-based assay coupled with LC-MS/MS to screen for novel SMS inhibitors and to provide an *in silico* method to validate the inhibition potential through molecular docking and dynamic simulation studies.

2. Materials and methods

2.1. Materials

Liquid chromatography/mass spectrometry (LC/MS)-grade solvents including isopropanol, chloroform, and methanol were purchased from Wako Pure Chemical Industries, Ltd (Osaka, Japan). Ammonium acetate (1 mol L^{-1} , eluent additive for LC/MS) was purchased from Sigma-Aldrich (St Louis, MO, USA). C6-Cer (d18:1/6:0) and C6-SM (d18:1/6:0) were purchased from Avanti Polar Lipids, Inc. (Alabaster, USA). Protease inhibitor cocktail (EDTA-free) was purchased from Nacalai Tesque, Inc. (Kyoto, Japan). GA C15:1 was purchased from Nagara Science Co. Ltd (Gifu. Japan). Bovine serum albumin (BSA) was purchased from Thermo Fisher Scientific (Tokyo, Japan).

2.2. Targeted analysis of C6-Cer and C6-SM by LC-MS/MS

Selected reaction monitoring channels for C6-Cer and C6-SM were established using a TSQ quantum access triple quadrupole mass spectrometer in positive ionization mode (Thermo Fisher Scientific, Inc., Waltham, MA, USA). The optimized ion source parameters were as follows: spray voltage: 3500 V, vaporizer temperature: 270 °C, capillary temperature: 300 °C, sheath gas (nitrogen): 30 psi, auxiliary gas (nitrogen): 40 psi, and collision gas (argon): 1.5 mTorr. Separation was achieved using an ultra-fast LC system (Shimadzu Corp., Kyoto, Japan) equipped with a Hypersil GOLD C8 column (50 mm × 2.1 mm, 6 µm; Thermo Fisher Scientific Inc.). The oven and sample tray temperatures were maintained at 40 °C and 4 °C, respectively. The mobile phases were A: 10 mM CH₃COONH₄ with 0.1% acetic acid, B: isopropanol, and C: methanol. The gradient elution was set as follows: 0-1 min (30% B, 30% C), 1-3 min (40% B, 50% C), 3-7 min (20% B, 70% C), 7-8 min (100% C), 8-10 min (30% B, 30% C), and re-equilibration for 2 min. The flow rate was 0.4 mL min⁻¹.

2.3 Plasmids

cDNAs encoding full-length SMS1 and SMS2 were obtained by RT-PCR from total RNA of HeLa cells. To obtain cDNA encoding SMS2, a sense primer (5'-GCCGGATCCACCatggatatcatagagacagcaaaac-3') and an antisense primer (5'-CGGGTCGACggtcgatttctcattgtcttcacc-3') were used. To obtain cDNA encoding SMS1, a sense primer (5'-gactgcctgctgtctgccagtac-3') and an antisense primer (5'gttcttagcacttcggacaatttgtc-3') were used, and the PCR product was re-amplified using a sense primer (5'-GCCGGATCCACCatgaaggaagtggtttattggtcac-3') and an antisense primer (5'-CGGCTCGAGtgtgtcattcaccagccggctg-3') to generate a restriction enzyme site. Both PCR products were introduced into a pMXs-puro expression vector with a C-terminal FLAG-His6 tag. The pMXs-puro vector and pEF-gagpol plasmid were kind gifts from Toshio Kitamura (The University of Tokyo). The pCMV-VSVG-RSV-Rev plasmid was a kind gift from Hiroyuki Miyoshi (RIKEN BioResource Center).

2.4 Establishment of cells stably expressing SMS

HeLa cells stably expressing C-terminally FLAG-tagged hSMS1 and hSMS2 were established according to a previously reported method.¹⁸ In brief, the pMXs-puro-derived vector for the expression of FLAG-tagged hSMS was co-transfected with pEF-gag-pol and pCMV-VSVG-RSV-Rev into HEK293T cells for retroviral production. The retroviruses were concentrated and used to infect HeLa cells to establish stable cell lines. Infected cells were selected in a medium containing puromycin (1 μ g mL⁻¹). A mixed population of drug-resistant cells was used for further analyses.

2.5. Immunoblot and immunofluorescence analyses

Immunoblot analysis was conducted as previously reported.¹⁸ Cells were lysed in lysis buffer (20 mM HEPES (pH 7.4), 150 mM NaCl, 1 mM EDTA, and 1% Nonidet P-40) containing a protease inhibitor mixture (Nacalai Tesque) at 4 °C for 30 min. The lysates were centrifuged at maximum speed in a microcentrifuge at 4 °C for 20 min to remove cellular debris. The lysates were incubated in SDS sample buffer containing β-mercaptoethanol at 37 °C for 2 h and subjected to SDS-PAGE and immunoblot analysis using mouse anti-DYKDDDDK (1E6) (Wako, Osaka, Japan) and anti-β-actin (C4) (Santa Cruz, Dallas, TX, USA) antibodies. Horseradish peroxidase-conjugated secondary antibodies were purchased from Jackson ImmunoResearch Laboratories (West Grove, PA, USA). The immunoblots were developed using ImmunoStar reagents (Wako) and recorded on an ImageQuant 800 (GE Healthcare, Chicago, IL, USA). Immunofluorescence was analyzed as previously described¹⁹ and visualized using an Axiovert 200MAT microscope (Carl Zeiss, Thornwood, NY, USA). Monoclonal rabbit anti-ATP1A1 (EP1845Y) was purchased from Abcam (Cambridge, UK) and monoclonal mouse anti-GM130 was purchased from BD Biosciences (Franklin Lakes, NJ, USA). Alexa Fluor-conjugated secondary antibodies were from purchased Invitrogen (Waltham, MA, USA).

2.6. Cell culture and protein assay

HeLa/hSMS1 and HeLa/hSMS2 cells were maintained in an incubator at 37 $^{\circ}$ C under a humidified atmosphere of 5% CO₂ in minimal essential medium (MEM) supplemented with 10% (v/v) fetal bovine serum and 1% penicillin–streptomycin–neo-mycin mixture (all reagents were sourced from Thermo Fisher Scientific, Tokyo, Japan). To determine the cellular protein

content, the cells were pelleted by centrifugation (1000 rpm, 10 min) and stored at -80 °C until use. In a 15 mL tube, 1 mL of 20 mM Tris buffer (pH 7.5) was added to the cell pellet, and the tube was vortexed. Then, 25 µL of the cell lysate was transferred into a 96-well plate in triplicate. To each well, 200 µL of a reaction reagent, consisting of a 50:1 mixture of reagent A and reagent B from the Bicinchoninic protein assay kit (Thermo Fisher Scientific), was added. The plate was incubated at 37 °C for 30 min, and the absorbance at 562 nm was measured using a Wallac 1420 ARVO Mx plate reader. To construct a calibration curve, a BSA solution with a concentration of 2000 µg mL⁻¹ was serially diluted to concentrations of 2000, 1000, 100, 10, 1, and 0.1 μ g mL⁻¹ using PBS. The absorbance of the BSA standards was measured as described above. The protein contents in the cell pellets were determined based on the calibration curve.

2.7. In vitro SMS assay

In vitro SMS activity was assayed as previously reported,¹⁴ with a modification. Lysis buffer (20 mM Tris-HCl, pH 7.5, containing protease inhibitor mixture) was added to HeLa/hSMS1 and HeLa/hSMS2 cell pellets and the samples were sonicated for 5 s. Cell lysates with protein concentrations of 0.1 μ g μ L⁻¹, 0.5 μ g μ L⁻¹, 0.75 μ g μ L⁻¹, and 1.0 μ g μ L⁻¹ respectively, were prepared. One hundred microliters of each cell lysate with known protein concentration was transferred to a 1.5 mL Eppendorf tube in quadruplicate and 1 μ L of C6-Cer (5 μ M in ethanol) was added. The quadruplicate mixtures were incubated at 37 °C for 5, 30, 60, and 120 min, respectively. Then, the enzymatic reaction was stopped by adding 350 μ L of a chloroform/methanol solution (2:1, v/v). The mixtures were vortexed (1300 rpm, 10 min) and centrifuged (15 000 rpm, 5 min), and the lower chloroform layer was transferred to a new Eppendorf tube. The chloroform extracts were dried under a vacuum using a centrifuge evaporator at 4 °C. The residue was redissolved in 100 µL of methanol, vortexed, and centrifuged (15000 rpm, 5 min), and the centrifugate was transferred into an LC vial for subsequent analysis. After optimization of the protein concentration and incubation, the experiments were conducted in the presence of a GA C15:1 at various concentrations. The detailed workflow of the SMS activity assay with or without inhibitor is shown in Fig. 1. The limit of detection, limit of quantification, extraction recovery, matrix effect (in cell lysates), and accuracy of the method were determined, and the results were provided in the ESI Table S1.† The LC-MS/MS raw data were processed using Xcalibur 2.2 software, and the peak intensities of C6-Cer and C6-SM were exported. The percentage of inhibition was calculated using the following equations:

$$SM(\%) = \frac{\text{intensity of SM}}{\text{intensity of Cer} + \text{intensity of SM}} \times 100$$
$$Inhibition(\%) = \left(1 - \frac{SM(\%)}{SM(\%) \text{ in neg. ctrl}}\right) \times 100$$

2.8. Methods for in silico analysis of SMS

2.8.1 SMS protein structure optimization. Three-dimensional structures of the target hSMS1 and hSMS2 are not available; hence, the target protein structures were optimized using AlphaFold 2.2.0. version.²⁰ Computational methods have been used for decades to predict three-dimensional protein models in the absence of experimental structures. Before AlphaFold was developed, the two main approaches were homology modeling and *ab initio* methods.²¹ The deep neural network of the AlphaFold algorithm has demonstrated outstanding accuracy



Fig. 1 Workflow of the SMS inhibition assay using HeLa cells expressing SMS1 and SMS2.

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in predicting three-dimensional structures of proteins with previously unknown folds by combining features derived from homologous templates and multiple sequence alignment. The canonical human phosphatidylcholine: ceramide choline phosphotransferase 1 [SGMS1] amino acid sequence (UniProt ID: Q86VZ5) was used to generate a structural model of SMS1 and the canonical human phosphatidylcholine: ceramide choline phosphotransferase 2 [SGMS2] amino acid sequence (UniProt ID: Q8NHU3) was used to generate a structural model of SMS2 using a locally installed AlphaFold 2.2.0 version, with default settings. In addition, a recent implementation of AlphaFold2 that allows different protein conformations to be sampled was used to obtain the structural models.²² The following parameters were applied: max msa clusters = 32, which sets the maximum number of sequence clusters selected at random and sent to the AlphaFold2 neural network; and max_extra_msa = 64, which sets the maximum number of additional sequences needed to calculate additional summary statistics. Note that the ideal values of these parameters vary depending on the target protein. Therefore, the minimal values of the AlphaFold2 advanced notebook were utilized, and these worked well for modeling the carrier's various conformations. To increase the number of models obtained to 40, the number of random seeds was set to 8. Subsequently, the minimization option was disabled and the number of recycles was set to one.²³

2.8.2 Molecular dynamics. To obtain a stable conformation of the modeled SMS protein, 10 ns molecular dynamics was performed using the Desmond membrane simulation protocol, which is widely used for simulating lipid bilayers and transmembrane proteins using the molecular dynamics software package Desmond.²⁴ This protocol is a powerful tool for studying the behavior of biomolecules in the complex environment of lipid bilayers.²⁵ The protocol involves several steps, including system preparation, equilibration, production, and analysis. During the equilibration phase, the system is allowed to relax into its minimum energy state, whereas in the production phase, the system is simulated for a set period, allowing for the observation of molecular dynamics and the calculation of various properties of interest. The analysis phase involves post-processing of the simulation data to obtain insights into the system's behavior.

2.8.3 Molecular docking studies. In molecular docking, ligand structure and orientation within a specified binding site are predicted. Molecular docking analysis was employed to study the interaction between GA C15:1 and the optimized SMS1 and SMS2, using the AutoDock Vina 1.5.6 software. The two-dimensional structure of GA C15:1 was downloaded from PubChem and converted to PDB format using the Marvin JS tool.²⁶ The ligands were prepared and saved in PDBQT format using AutoDock Vina 1.5.6. The optimized structures of the receptors SMS1 and SMS2 were acquired in the PDB format. The bounded atoms were removed during protein structure optimization. Polar hydrogen was added to the receptors using the AutoDock Vina 1.5.6. tool.²⁷ A grid box was generated for both receptors, targeting the active site of the receptor with a

size of *X*, *Y*, *Z*. The active site of SMS1 fell within the coordinates with values x = 3.16, y = 1.03 and z = 11.48, and for SMS2, the values were x = 12.25, y = 4.58 and z = -6.05. A molecular docking analysis was then carried out using AutoDock Vina 1.5.6. The binding energies between the ligand and target were expressed in kcal mol⁻¹. Docking was analyzed and visualized using Biovia Discovery Studio 2021 Visualizer. Further molecular dynamic simulation studies were carried out for 100 ns to assess the stability of the protein–ligand complex.²⁸

3. Results and discussion

3.1. Detection of the SMS substrate (C6-Cer) and product (C6-SM) by LC-MS/MS and its stable expression in HeLa cells

Authentic standards C6-Cer (i.e., Cer (d18:1/6:0)) and C6-SM (i.e., SM (d18:1/6:0)) were selected as a substrate and product of SMS, respectively, to monitor their activity in cell lysates. Ceramides with shorter acyl chains have faster enzymatic reaction rates; hence, the unnatural C6-Cer was selected as a substrate for both SMS1 and SMS2. Using a triple-quadrupole mass spectrometer, the selected reaction monitoring (SRM) channels were optimized for C6-Cer and C6-SM. Elution of C6-Cer and C6-SM was achieved within 4 min using a reverse-phase C₈ column coupled with an LC system. The SRM optimization curves, extracted ion chromatograms, and MS/MS spectra of C6-Cer and C6-SM are shown in Fig. 2. C6-Cer and C6-SM ionized to yield $[M + H]^+$ product ions of m/z 398.6 and m/z 563.7, respectively. The optimization curves (Fig. 2A) show that m/z 264.3 and m/z184.1 were the most intense product ions of C6-Cer and C6-SM, which eluted at 3.72 and 3.48 min respectively (Fig. 2B). The MS/MS spectra of C6-Cer and C6-SM and their fragmentation patterns are shown in Fig. 2C. These MS/MS spectra are similar to those of Cer and SM with different acyl chains, as reported in our previous study.²⁹

A previous study utilized (6-((N-(7-nitrobenz-2-oxa-1,3diazol-4-yl)amino)hexanoyl)sphingosine) C6-NBD-Cer and C6-NBD-SM as a fluorescence substrate and product, respectively, to monitor the in vitro activity of SMS with an HPLC fluorescence detector.14 However, fluorescent-based detection methods are less sensitive than LC/MS.³⁰ Chen et al.⁴ previously used LC-MS/MS to monitor SMS1 and SMS2 activities based on C6-Cer and C6-SM levels. The authors overexpressed SMS1 and SMS2 in HEK293 cells and established the first quantitative method for SMS activity. However, their method is time-consuming and requires large amounts of protein. To establish a functional biochemical system, we opted to express recombinant SMS1 and SMS2 proteins in HeLa cells. These recombinant proteins had a FLAG tag at their C-terminus, allowing us to monitor their expression levels. Immunoblot and immunofluorescence analyses were conducted to confirm the stable expression of hSMS1 and hSMS2 in HeLa cells (Fig. 3). Both SMS1 (413 amino acids) and SMS2 (365 amino acids) were detected as proteins with an approximate mole-



Fig. 2 Optimization of the single-reaction monitoring channels for C6-cer and C6-SM. (A) Optimization curves for C6-Cer and C6-SM. (B) Extracted ion chromatograms of C6-Cer and C6-SM. (C) MS/MS spectra of C6-Cer and C6-SM acquired using a triple quadrupole mass spectrometer.



Fig. 3 Stable expression of hSMS1 and hSMS2 in HeLa cells. (A) HeLa cells stably expressing C-terminally FLAG-tagged hSMS1 or hSMS2 were lysed, and the expression levels of hSMS1-FLAG and hSMS2-FLAG were analyzed by immunoblotting with anti-DYKDDDDK (for detecting the FLAG-tag) and anti-β-actin (as an internal control) antibodies. (B) HeLa cells stably expressing hSMS1-FLAG or hSMS2-FLAG were fixed, permeabilized, and incubated with anti-DYKDDDDK, anti-GM130 (a Golgi complex marker), and anti-ATP1A1 (a plasma membrane marker), and subsequently with Alexa Fluor 555-conjugated anti-mouse IgG2b, Alexa Fluor 488-conjugated anti-mouse IgG1, and Alexa Fluor 647-conjugated anti-rabbit antibodies. Bars, 20 μm.

cular weight of 40 kDa (Fig. 3A). SMS1-FLAG was exclusively detected in the Golgi complex, whereas SMS2-FLAG was observed in the plasma membrane, as described previously,^{2,5,6} as well as in the Golgi complex (Fig. 3B). Findings in a previous study that also stably expressed SMS1/SMS2 in human HeLa cells³¹ support our results.

3.2. *In vitro* evaluation of SMS activity using the natural inhibitor GA

To evaluate *in vitro* SMS activity, the amount of protein and reaction incubation time needed to be optimized. The results of the optimization using SMS1 and SMS2 are shown



Fig. 4 Optimization of the incubation time and amount of protein for the SMS activity assay in the presence of GA. (A) Optimization of the incubation time and protein concentration for the SMS inhibition assay using HeLa/SMS1 and HeLa/SMS2 cells. (B) Extracted ion chromatograms of C6-SM (*i.e.*, SM(d18:1/6:0)) and C6-Cer (*i.e.*, Cer(d18:1/6:0)) peaks with and without inhibitor. (C) Results of *in vitro* assays using GA, and its IC_{50} values for hSMS1 and hSMS2.

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in Fig. 4A. The peak intensity for the enzymatic product C6-SM slightly increased with increasing protein concentration; however, the change was not substantial. Therefore, 0.1 μ g μ L⁻¹ was selected as an optimal concentration for both SMS1 and SMS2 inhibition assays. The production of C6-SM in HeLa, HeLa/SMS1, and HeLa/SMS2 cells was also examined over time, and the results are provided in ESI Fig. S1.† C6-SM was produced at higher levels in SMS-expressing cells than in HeLa control cells. C6-SM production reached a maximum after a 60 min incubation at 37 °C. Hence, 0.1 μ g μ L⁻¹ of protein and a 60 min incubation were considered optimal conditions for the SMS1 and SMS2 enzymatic reactions. Zama *et al.*¹⁴ previously developed a cell-based assay using ZS/SMS1 and ZS/SMS2 cells and a fluorescent substrate and used a similar incubation time in their experi-

ments. However, in a previous method using C6-NBD-Cer as a substrate for ZS/SMS1 or ZS/SMS2, the incubation time was 3 h.¹⁶ In contrast, our method using C6-Cer and HeLa/ SMS2 cells requires an incubation time of 60 min, which suggests the robustness of our assay method. To perform a precise quantitative assessment of SMS activity, we employed an LC-MS/MS technique to measure C6-Cer and C6-SM levels. The ion chromatograms of C6-Cer and C6-SM with and without inhibitor (*i.e.*, GA) are shown in Fig. 4B. The results demonstrate that C6-SM production was decreased in the presence of the inhibitor GA.

GA is a potent inhibitor of both SMS1 and SMS2 isolated from *Ginkgo biloba* stem. Given its commercial availability and SMS inhibition potency, we tested our assay method using GA. The half-maximal inhibitory concentrations (IC_{50}) of GA (at



Fig. 5 SMS protein-structure optimization and *in silico* insights. (A) Optimized structures of hSMS1 and hSMS2 were estimated using AlphaFold2. (B) Interaction of GA with hSMS1 and hSMS2 (orange: ribbon structure of amino acid residues in SMS1 and SMS2 protein; green: ball-stick representation of the carbon atoms in GA; red: ball-stick representation of the oxygen atoms in GA). (C) Two-dimensional interaction of GA with hSMS1 and hSMS2 (orange: ribbon structure of structure) atoms in GA). (C) Two-dimensional interaction of GA with hSMS1 and hSMS2 (conventional hydrogen bond between C=O and SER A:269 of SMS1 protein, conventional hydrogen bond between C=O and ASN A:307, C-OH and SER A:310, and O-H and ARG A:311 of SMS2 protein).

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various concentrations) for SMS1 and SMS2 are shown in Fig. 4C. The IC_{50} values of GA for SMS1 and SMS2 were 5.5 μ M and 3.5 μ M, respectively. A previous study using the fluorescence method reported an IC_{50} value of 1.5 μ M for both SMS1 and SMS2.¹⁶ Many factors can explain these differences, including the cell type used, SMS expression level, and method of substrate and product detection used. Despite this, the results are close to the values reported in the previous study, with 2–3-fold variations. Our assay system provides a specific and sensitive method to evaluate the structural and functional relationships of both enzymes.

3.3. *In silico* insights into GA as a potent inhibitor of SMS1 and SMS2

To validate the in vitro results, in silico analysis was performed to optimize the hSMS1 and hSMS2 protein structures. To our knowledge, crystal structures of hSMS1 and hSMS2 have not been reported to date. To optimize the protein structures, AlphaFold 2.2.0 was used. First, sequence analysis was performed using the gene sequences of the proteins from the National Center for Biotechnology Information (NCBI) database (https://www.ncbi. nlm.nih.gov/accessed on 2023/04/08). The gene sequence of hSMS1, with 400 residues, was compared with more than 4000 sequences, whereas that of hSMS2, with 350 residues, was compared with 3500 sequences. The detailed comparison is shown in ESI Fig. S2A.[†] Based on this analysis, five models were generated for each protein. Among the five models, model 5 ranked first for both proteins, with a predicted local distance difference test (pLDDT) score of 79 for SMS1 and of 73.9 for SMS2. The other models had lower pLDDT scores than model 5, as shown in ESI Fig. S2B.† The predicted template modeling score of model 5 was 0.638 for SMS1 and 0.709 for SMS2. Hence, this modeled structure was used for molecular dynamics simulation studies. Molecular dynamics were simulated for both proteins to obtain a stable conformation for up to 10 ns. The best conformational structures of SMS1 and SMS2 from these simulations (Fig. 5A) were used for molecular docking analysis. Previously, an effort has been made to optimize the hSSM1 structure using Ambalone and Yasara software.32

The locations of the binding sites of GA in hSMS1 and hSMS2 are unknown. We predicted the structures of the complexes formed by GA with hSMS1 and hSMS2 by molecular docking studies using AutoDock Vina 1.5.6. GA interacted with hSMS1 and hSMS2 with binding energies of -6.6 kcal mol⁻¹ and -8.2 kcal mol⁻¹, respectively. The docking interaction modes were obtained using Discovery Studio Visualizer for evaluation, and the results are shown in Fig. 5B. GA interacts with the amino acid residue SER269 of hSMS1, and with ARG311, SER310, and ASN307 of SMS2 through conventional hydrogen bonding (Fig. 5C). To assess the stability of the GAhSMS1/hSMS2 complexes, molecular dynamics simulations were carried out. The simulation revealed a good root mean square deviation for both GA-hSMS1/hSMS2 complexes, with values of 3.5 Å and 6.1 Å, respectively (as shown in ESI Fig. S3[†]), indicating good stability. A previous study revealed a similar unique interaction between hSMS1 protein and tricyclodecan-9-yl-xanthogentate (D609), a well-known SMS inhibitor.³³ A similar strategy was used to predict small-molecule inhibitors of hSMS1 by virtual screening followed by *in vitro* enzymatic assays.

4. Conclusion

SMS plays a pivotal role in maintaining sphingolipid homeostasis by facilitating the conversion of Cer into SM within the framework of sphingolipid metabolism. As SMS is a potential key therapeutic target in various metabolic diseases, it is important to develop a method to monitor its levels. We developed an LC-MS/MS-based method to monitor SMS1 and SMS2 activity using C6-Cer and C6-SM as a substrate and product, respectively. *In vitro* and *in silico* studies were conducted using a known SMS inhibitor, GA. Both *in vitro* and *in silico* results supported the inhibition of hSMS1 and hSMS2 by GA. This method has the potential to be used in screens for novel SMS inhibitors.

Author contributions

P. M. S.: formal analysis; validation; investigation; methodology; visualization; writing – review and editing. Y. M.: formal analysis; validation; investigation; methodology; visualization; data curation. J. J.: formal analysis; validation; investigation; methodology; visualization. H. T.: formal analysis; validation; data curation, writing – review and editing. H. W. S: formal analysis; validation; data curation, writing – review, and editing. D. G.: data curation; methodology; writing – review and editing. S. G. B. G.: conceptualization; formal analysis; validation; investigation; supervision; project administration; writing – original draft; funding acquisition. S. P. H.: Supervision; resources; writing – review and editing.

Conflicts of interest

The authors have no conflict of interest to declare.

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