Evaluating Sphingosine and its Analogues as Potential Alternatives for Aggressive Lymphoma Treatment

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Key Words
Sphingosine • Ceramide • Autophagy • Apoptosis • Protein kinase C • Diffuse large B cell lymphoma

Abstract
Background: Ceramide (Cer) and sphingosine (Sph) interfere with critical cellular functions relevant for cancer progression and cell survival. While Cer has already been investigated as a potential drug target for lymphoma treatment, information about the potency of sphingosine is scarce. The aim of this study therefore was to evaluate Sph and its synthetic stereoisomer L-threo-sphingosine (Lt-Sph) as potential treatment options for aggressive lymphomas.

Methods: Diffuse large B cell lymphoma (DLBCL) cell lines were incubated with Sph and Lt-Sph and consequently analysed by flow cytometry (FACS), enzyme-linked immunosorbent assay (ELISA), liquid chromatography coupled to triple-quadrupole mass spectrometry (LC/MS/MS), electron microscopy, and Western blot. Results: Sph induced cell death and blocked cell growth independently of S1P receptors in different DLBCL cell lines. Three different modes of Sph-mediated cell death were observed: Apoptosis, autophagy, and protein kinase C (PKC) inhibition. Generation of pro-apoptotic Cer accounted only for a minor portion of the apoptotic rate. Conclusion: Sph and its analogues could evolve as alternative treatment options for aggressive lymphomas via PKC inhibition, apoptosis, and autophagy. These physiological responses induced by different intracellular signalling cascades (phosphorylation of JNK, PARP cleavage, LC3-II accumulation) identify Sph and analogues as potent cell death inducing agents.

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Introduction

DLBCL is the most common of the aggressive lymphomas with estimated 70,000 new cases and 19,000 deaths in the United States in 2013 [1]. It is characterized by highly heterogenous morphology, biology, and clinical presentation [2]. Different patterns of gene expression give rise to 2 distinct subtypes of DLBCL: Germinal center-like (GCB) and activated B cell-like (ABC) [3]. A major characteristic of ABC DLBCL is the constitutive activation of the nuclear factor-kappa B (NF-κB) pathway [4]. A majority of GCB DLBCL was found to be dependent on the phosphatidylinositide 3-kinase (PI3K) and protein kinase B (Akt) pathway [5]. In contrast to slow-growing indolent non-Hodgkin's lymphomas (NHL), rapidly growing DLBCL is often curable with a success rate greater than 50% [1]. ABC DLBCL has the worst prognosis with 3-year overall survival rates of around 40% [6].

Sphingolipids like Sph, sphingosine 1-phosphate (S1P), and Cer are important determinants for cell fate [7, 8]. While S1P signalling through G protein-coupled cell surface S1P receptors is considered as a pro-survival factor [9], Cer and Sph both induce cell death, albeit using different signalling pathways [7, 8]. The exact mechanisms of both lipids are not completely understood (reviewed in [10, 11]). Cer for example interferes with mitochondrial functions [12, 13], but also upregulates apoptosis-inducing proteins like Bcl-xS and caspase-9 [14]. It directly activates PKC-ζ and binds to cathepsin D to support its proteolytic maturation and activation [15, 16]. Cer also induces autophagy. Dependent on the cell system used, the mechanism involves the induction of ER stress [17], suppression of Akt [18], activation of JNK [19], up-regulation of Beclin 1 [20], and expression of the mitochondrial BH3-only protein BNIP3 [21]. Little is known about the signalling pathways that transmit Sph-induced apoptosis. It inhibits PKC and mitogen-activated protein kinase (MAPK) [22, 23], but may also act through ceramide synthase-dependent conversion to Cer [24]. It can be rapidly phosphorylated by sphingosine kinases (SphK) type 1 and 2 to S1P [25, 26] which has antiapoptotic functions predominantly by activating S1P receptors [9, 27]. These two different metabolic conversions may explain antipodal observations demonstrating down- or upregulation of antiapoptotic Bcl-2 proteins in different cell systems [28, 29].

The generation of pro-survival S1P by SphKs supports tumor growth (reviewed in [30]). Current strategies for cancer treatment therefore include inhibition of SphKs, particularly SphK1, as a potentially new therapeutic avenue [31]. While prevention of S1P generation is generally regarded as the main anti-cancer effect due to abrogation of pro-survival S1P signalling, the effect of concomitant Sph accumulation has typically not been considered as being effective. Different interventions in sphingolipid metabolism like deficiency of SphK2 [32] or the S1P degrading enzyme S1P-lyase [33] however result in increased Sph concentrations. Moreover sphingosine analogues were tested for their ability to inhibit SphKs [34]. SphK inhibitors that are structurally related to Sph may not only prevent Sph phosphorylation, but could also share functional properties of Sph. We therefore investigated the effect of Sph on different DLBCL in order to better understand the idiosyncratic functions of Sph in the context of aggressive lymphomas, and to explore Sph accumulation and Sph analogues as alternative treatment options for aggressive lymphomas.

Materials and Methods

Chemicals

The following chemicals were used throughout the study: S1P (Sigma), Sph (Sigma), Lt-Sph (Avanti Polar Lipids), C17-Sph (Avanti Polar Lipids) sphinganine (Avanti Polar Lipids), C15-Cer (Matreya), C16-Cer (Matreya), phosphatidylinerseine (PS, Sigma), camptothecin (Sigma), fumonisin B1 (Cayman), 4-deoxypyridoxine (DOP, Sigma).

Cell culture

HT, HBL-1, and U2932 cells were grown in RPMI 1640 (Life Technologies) and OciLy19 in Iscove’s modified Dulbecco’s medium (IMDM) with 10% fetal bovine serum (FBS, Biochrom), 1 mM sodium...
pyruvate (PAA Laboratories), 100 units/ml penicillin G (PAA Laboratories), 100 µg/ml streptomycin (PAA Laboratories), and 2 mM L-glutamine (PAA Laboratories) [35].

Analysis of cell growth

Cells were plated in 96-well plates at 100,000 cells/well and treated with the indicated concentrations of the respective compound. Immediately and 1-4 days later, cells were lysed with CellTiter-Glo luminescent cell viability assay reagent according to the manufacturer’s protocol (Promega), and the resulting luminescence was determined with the Victor3 plate reader (PerkinElmer).

Determination of apoptosis by FACS

Cells were plated in 24-well plates at 100,000 cells/ml and treated with 3 µM (OciLy19) or 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph, and with 20 µM camptothecin for 4 h. Subsequently cells were washed twice with ice-cold binding buffer (0.01 M Hepes/NaOH (pH 7.4), 0.14 M NaCl, 2.5 mM CaCl2). Five µl fluorescein conjugated annexin V (annexin V-FITC, Immunotools) was added to 100 µl cell suspension in binding buffer and incubated at room temperature for 15 min. Five µl of 50 µM propidium iodide was added immediately before FACS analysis using the FACSCalibur (Becton Dickinson).

Lipid quantification

Lipid quantification was done as described [36, 37]. Biological samples (1 ml of medium or 10E6 cells) were adjusted to 1 ml sample volume with 1 M NaCl in H2O and transferred into a glass centrifuge tube. After addition of 1 ml of methanol and 200 ml of 6 M HCl, the samples were vortexed. Chloroform (2 ml) was added, and the samples were again vigorously vortexed for 2 min. After the samples were centrifuged for 3 min at 1,900 x g, the lower chloroform phase was transferred to another glass centrifuge tube. After the lipid extraction was repeated with 2 ml of chloroform, the chloroform phases were combined and vacuum dried in a speed-vac for 45 min at 50 °C. The QTrap triple-quadrupole mass spectrometer (ABSciex) interfaced with a Merck-Hitachi LaChrom chromatograph and autosampler was used for electrospray ion (ESI) LC/MS/MS analysis. Positive ion ESI LC/MS/MS analysis was employed for detection of all analytes. The multiple reaction monitoring transitions for the detection were as follows: C17-Sph m/z 286/268, C15-Cer m/z 524/264, Sph m/z 300/282, spinganine m/z 302/284, S1P m/z 380/264, C16-Cer m/z 538/264, C24-Cer m/z 650/264. Liquid chromatographic resolution of all analytes was achieved using a MultoHigh RP 18-3 µm column (2 mm x 60 mm, CS Chromatographie Service). The elution protocol was composed of a 9 min column equilibration with 10% solvent A (methanol) and 90% solvent B (1% formic acid) followed by sample injection and a 20 min period with 100% solvent A. Samples were infused into the ESI source through an electrode tube at a rate of 300 µl/min. Standard curves were generated by adding increasing concentrations of the analytes to 300 pmol of C17-Sph and C15-Cer (internal standards). Linearity of the standard curves and correlation coefficients were obtained by linear regression analyses. All mass spectrometry analyses were performed with Analyst 1.4 (ABSciex).

Cell cycle analysis

Cells were plated in 24-well plates at 100,000 cells/ml and treated with 3 µM (OciLy19) or 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph in the presence or absence of 15 µM PS for 3 days. Cells were harvested and resuspended in 400 µl hypotonic lysis buffer (50 µg/ml propidium iodide in 0.5 phosphate buffered saline (PBS) and 0.1% Triton X-100). After incubation at room temperature for 1 h, the liberated nuclei were analyzed by FACS with the FACSCalibur (Becton Dickinson).

Staining of acidic vesicular organelles

Cells were plated in 24-well plates at 100,000 cells/ml and treated with 3 µM (OciLy19) or 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph, and with 10 µM camptothecin for 1 day. Subsequently acridine orange was added to the cell suspension at a final concentration of 1 µg/ml for 15 min. Cells were washed twice with ice-cold PBS and analyzed by FACS with the FACSCalibur (Becton Dickinson).

Electron microscopy

Samples were fixed in 2.5% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.3) overnight. Samples were washed, postfixfixed with 2% osmium tetroxide in 0.1 M cacodylate buffer for two hours, dehydrated
with graded ethanol solutions, and embedded in Epon (SERVA). Semithin sections were stained with Richardson’s stain [38]. Ultrathin sections were stained with uranyl acetate and lead citrate [39]. The samples were analyzed on a transmission electron microscope EM 906 (Zeiss, Oberkochen). Sample preparation was done by Petra Schrade (Electron Microscopy Facility, Charité - University Medical School Berlin, Germany).

**PKC activity assay**

PKC activity was tested using the PepTag fluorescent protein kinase assay (Promega) according to the manufacturer’s protocol. All PepTag PKC assay reaction components were combined on ice, and PKC activity was assayed in a final volume of 25 µl of the following mixture: 5 µl of 5x PKC reaction buffer (100 mM HEPES, pH 7.4, 6.5 mM CaCl₂, 5 mM dithiothreitol, 50 mM MgCl₂, 5 mM ATP), 5 µl of PepTag C1 peptide (PLSRTLSVAAK, 0.4 µg/µl in water), 5 µl of freshly sonicated PKC activator solution (1 mg/ml PS in water), 1 µl of peptide protection solution, 3.75 µl of water, 1.25 µl of either vehicle (methanol), Sph, or Lt-Sph, and 4 µl of supplied PKC (2.5 µg/ml in PKC dilution buffer containing 100 µg/ml FBS and 0.05% Triton X-100). Before adding PKC, the mixture was preincubated at 30 °C for 2 min. After the addition of PKC, the entire reaction mixture was incubated at 30 °C for 30 min. The reaction was stopped by incubation at 95 °C for 10 min. Before loading samples on an agarose gel (0.8% agarose in 50 mM Tris-HCl buffer, pH 8.0), 2 µl of 80% glycerol was added to the sample. Electrophoresis was run at 100 V for 30 min in 50 mM Tris-HCl, pH 8, and was imaged immediately under UV light. Signals were quantified using ImageJ (NIH).

**Quantification of IL-10**

Enzyme-linked immunosorbent assay (ELISA) was used to quantify human interleukin-10 (IL-10) in the supernatant of DLBCL cell lines. 300,000 HBL-1 cells/300 µl were grown for 24 h and 60,000 U2932 cells/300 µl were grown for 6 h in 96-well plates (TPP) in the presence and absence of 3-5 µM Sph and Lt-Sph. Subsequently cells were centrifuged at 300 x g and the supernatant was harvested. Maxisorp 96-well plates (NUNC) were coated with the coating antibody provided by the IL-10 ELISA set and processed according to the manufacturer’s protocol (Immunotools). Plates were developed with 3,3',5,5'-tetramethylbenzidine (TMB) substrate solution (eBioscience). The reaction was stopped with 1N HCl. Absorbance at 450 nm was detected with the Victor plate reader (PerkinElmer). Standard curves were generated with 3-300 pg/ml IL-10 and used for quantification.

**Determination of AKT phosphorylation and PARP cleavage**

Cells were plated in 6-well plates at 100,000 cells/ml and treated with 3 µM (OciLy19) or 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph. Subsequently cells were transferred into 96-well plates at a density of 100,000 cells per well and tested for the presence of AKT (protein kinase B), phospho-AKT (pAKT), cleaved poly (ADP ribose) polymerase (PARP), and tubulin with colorimetric in-cell ELISA kits according to the manufacturer’s protocol (Pierce Biotechnology). Normalization was performed by whole cell staining with Janus green.

**Determination of mTOR and JNK phosphorylation**

Cells were plated in 6-well plates at 100,000 cells/ml and treated with 3 µM (OciLy19) or 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph. After harvesting they were tested for the presence of the c-Jun N-terminal kinase (JNK), phospho-JNK (pJNK), and Ser2448 phosphorylated mammalian target of rapamycin (mTOR) with ELISA kits according to the manufacturer’s protocol (Abcam). Protein concentrations were 130 µg/ml (OciLy19) and 190 µg/ml (HT, HBL-1, U2932) for JNK measurements, 160 µg/ml (OciLy19) and 190 µg/ml (HT, HBL-1, U2932) for pJNK measurements, and 160 µg/ml (OciLy19) and 190 µg/ml (HT, HBL-1, U2932) for mTOR measurements, respectively.

**Western-blot analysis**

Western blots were performed according to standard protocols. Cells were plated in 6-well plates at 100,000 cells/ml and treated with 3 µM (OciLy19) or 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph. After harvesting, 200,000 cells were lysed in 20 mM Tris–HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA, pH 8.0, 1% Triton-X-100, 20 mM NaF, 0.1 mM Na₃VO₄, and complete protease inhibitor cocktail (Roche Applied Science). Lysates (12 - 23 µg) were subjected to 8-16% Bis-Tris gels (GE Healthcare) according to the manufacturer’s
protocol, and proteins were transferred to Hybond-P polyvinylidene difluoride (PVDF) membranes (GE Healthcare) by wet blotting. Membranes were subsequently blocked with 5% SlimFast chocolate powder (Allpharm-Vertriebs-GmbH) in Tris-buffered saline and probed with 1:1000 dilutions of the following primary antibodies (Cell Signaling Technology) overnight at 4 °C: Rabbit anti-LC3A (clone D50G8), rabbit anti-calnexin (clone CSC59), rabbit anti-Ero1-Lα, or rabbit anti-IRE1α (clone 14C10). After incubation with a specific horseradish peroxidase (HRP)-labeled secondary antibody against rabbit (Cell Signaling Technology #7074, 1:2000 dilution), signals were visualized with the enhanced chemiluminescent detection system (ECL) according to the manufacturer’s instructions (GE Healthcare).

Statistical analysis
A two-tailed unpaired Student t test was used to determine the significance of differences (*p<0.05, **p<0.01, and ***p<0.001).

Results
Sph inhibited cell growth and induced apoptosis in DLBCL cell lines
To investigate the impact of cellular Sph accumulation on lymphoma cell growth, different DLBCL cell lines (OciLy19, HT, HBL-1, U2932) were incubated with increasing concentrations of Sph. Determination of cell growth each day over 4 days revealed concentration-dependent growth inhibition after treatment with Sph in all DLBCL cell lines tested, with HBL-1 cells being most resistant to higher Sph concentrations, which were thus treated with 2-fold higher sphingolipid concentrations (Fig. 1A). All DLBCL cell lines also responded to Lt-Sph with decreased growth, with HT cells showing a lower impact of Lt-Sph than Sph. Lt-Sph is a stereoisomer of naturally occurring D-erythro-sphingosine (Sph). It can be phosphorylated by cells, but the resulting Lt-S1P does not bind and activate S1P receptors, excluding their role in Lt-Sph-induced growth inhibition [40-42].

To test the onset of apoptosis as one possible reason for the observed growth retardation in examined cells, DLBCL cell lines were treated for 4 h with Sph and Lt-Sph, and with 20 µM camptothecin as positive control. Camptothecin is a cytotoxic quinoline alkaloid which induces apoptosis by inhibiting the topoisomerase type 1 (topo1) [43]. All cell lines were treated with 5 µM Sph and Lt-Sph except OciLy19, which were more sensitive to Sph and Lt-Sph incubation and were therefore treated with 3 µM Sph and Lt-Sph. Apoptosis was detected by FACS after staining apoptotic cells with annexin V and dead cells with propidium iodide. All DLBCL cell lines responded with 28-59% apoptosis to camptothecin after 4 h. Sph also induced a high apoptotic rate of 23-32%, while Lt-Sph was less efficient with apoptotic rates between 12-22% (Fig. 1B). U2932 cells did not respond to Lt-Sph treatment with apoptosis at all (Fig. 1B). Most of the propidium iodide and annexin V double-positive cells after Lt-Sph treatment did not pass the annexin V single-positive stage and were therefore not considered as apoptotic cells.

Apoptosis is mainly driven by activation of death proteases known as caspases [44], although caspase-independent apoptosis has also been described [45]. Caspases cleave proteins that are essentially required for cellular function and cell survival. One of these many target proteins of caspases is PARP, and cleaved PARP is commonly used as an indicator for caspase-induced apoptosis [46]. Incubation of DLBCL cell lines with Sph and Lt-Sph as mentioned before induced significant PARP cleavage only in OciLy19 DLBCL predominantly after Sph treatment (Fig. 1C). A significant but less pronounced increase in PARP cleavage was also found after Lt-Sph treatment. No important differences in PARP cleavage were observed in similarly treated HT, HBL-1, and U2932 DLBCL cell lines (Fig. 1C). A related pattern was observed for JNK phosphorylation (Fig. 1D). pJNK was significantly increased only in OciLy19 DLBCL treated with Sph, but not Lt-Sph. Both stimuli did not induce pJNK in HT, HBL-1, or U2932 DLBCL cell lines, which obviously depended on different signalling pathways.
The role of ceramide in Sph-induced apoptosis

Sph can either be phosphorylated to S1P, which binds and activates S1P receptors, or it can be acylated to ceramide, which also exerts pro-apoptotic activities using caspase-dependent and –independent pathways. To test the relevance of ceramide generation for apoptosis induction after Sph treatment, DLBCL cell lines were treated with the ceramide synthase inhibitor fumonisin B1. The apoptotic rate of OciLy19, HT, and U2932 cell lines after treatment with camptothecin, Sph, and Lt-Sph, and 20 µM camptothecin (CPT) was not significantly altered by addition of 25 µM fumonisin B1, while HBL-1 cells showed a significant decrease of the apoptotic rate in Sph treated cells after fumonisin B1 treatment (Fig. 2A). The activity of fumonisin B1 was confirmed by LC/MS/MS quantification of the ceramide synthase substrate sphinganine in control and fumonisin B1 treated cells. Fumonisin B1 was shown to increase the amount of sphinganine via inhibition of ceramide synthases [47]. All DLBCL cell lines demonstrated a significant increase in sphinganine levels after treatment with fumonisin B1 (Fig. 2B). Increases were 2.5-fold and 10-fold for HT and OciLy19, and 32-fold and 39-fold for HBL-1 and U2932 cells, respectively. Triple-quadrupole mass spectrometry revealed similar Sph and S1P accumulation with and without fumonisin B1 in all DLBCL cell lines (Fig. 2C). Intracellular Cer accumulation was only detectable in Sph treated HBL-1 and U2932, but not...
in HT and OciLy19 DLBCL cell lines (Fig. 2D). It was efficiently blocked by 25 µM fumonisn B1 (Fig. 2D). Thus, intracellular Cer accumulation was only evident in Sph-treated HBL-1 and U2932 DLBCL and did not account for most of the observed apoptosis except for HBL-1 cells which showed significantly reduced apoptosis. OciLy19 and HT DLBCL did not show any alterations in ceramide levels nor in the apoptotic rate after ceramide synthase inhibition.

**Autophagy contributed to Sph-induced cell death**

In order to find out more about the underlying mechanisms involved in Sph-induced cell death, we looked for the development of acidic vesicular organelles 1 day after addition of Sph, Lt-Sph, and camptothecin (CPT). Acidic vesicular organelles are characteristic for autophagy and can be detected by FACS with acridin orange, which accumulates in acidic compartments and produces bright red fluorescence. All 4 tested DLBCL cell lines demonstrated an increase
Fig. 3. Analysis of autophagy in DLBCL treated with 3 µM (OciLy19) and 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph, and 10 µM camptothecin. (A) FACS analysis of acidic vesicular organelles stained with acridine orange 1 day after treatment. Shown are histograms of 1 representative experiment out of 3, and linear means ± SD of 3 experiments, normalized to control (=100). *p<0.05, **p<0.01, ***p<0.001. (B) Electron microscopy pictures of untreated U2932 cells (control) and U2932 cells treated with 5 µM Sph for 1 and 3 days. Potential autophagic vesicles are marked with an arrow. Potential autophagosomal structures in colored squares are shown at higher magnification (m) in equivalently colored frames to demonstrate the presence of double membranes characteristic for autophagosomes. (C) Western-blot analysis of the autophagic marker LC3-II and the ER stress markers PERK, calnexin, Ero1-Lα, and IRE1α in combination with ELISA results of the phosphorylated signaling molecules pAKT and mTOR (pSer2448) 4 h after treatment with 3 µM (OciLy19) and 5 µM (HT, HBL-1, U2932) Sph and Lt-Sph with and without prior addition of 25 µM fumonisin B1 (FB1). Shown are representative Western-blot results from 2-3 independent experiments and means ± SD of ELISA results for AKT and mTOR phosphorylation, untreated control = 1.0, n=3. pAKT values were normalized to AKT values, the relative density of LC3-II Western blot signals was normalized to the respective beta-actin expression.
of red fluorescence as determined by FACS after treatment with Sph and Lt-Sph, but not with camptothecin (Fig. 3A). While HBL-1 and U2932 DLBCL cell lines responded equally well to both stimuli, OciLy19 and HT revealed a greater shift in the red fluorescence after Lt-Sph stimulation than after Sph-stimulation. U2932 cells did not respond to Lt-Sph treatment with apoptosis (Fig. 1B, 2A), but they showed a similar increase in acidic compartments compared to Sph treatment (Fig. 3A). Electron microscopy of U2932 cells 1 and 3 days after treatment with Sph uncovered the presence of potential autophagosomes or secondary lysosomes, respectively. In comparison to control cells, Sph-treated cells developed increased vacuole formation one and three days after Sph addition (Fig. 3B). Examination of these vacuoles at higher magnification revealed the presence of double-membranes which are characteristic for autophagosomes (Fig. 3B). The presence of these double-membrane vesicles together with the observed increase in acidic compartments are in support for the additional induction of autophagy after treatment with Sph and Lt-Sph (Fig. 3A-B). Further evidence derived from the analysis of the autophagy marker light chain 3 (LC3), which is cleaved at the carboxy terminus to the LC3-I form immediately after its synthesis. LC3-I is converted to LC3-II via lipidation and becomes associated with autophagic vesicles. The conversion of LC3 to the lower migrating form LC3-II is indicative for ongoing autophagy in Western blot analyses [48]. Treatment of DLBCL cell lines with Sph and Lt-Sph led to an increase of LC3-II in U2932 DLBCL, while no specific signal was observed in OciLy19, HT, and HBL-1 DLBCL cell lines (Fig. 3C). LC3-II formation was not inhibited by fumonisin B1 (Fig. 3C). Neither the protein kinase AKT nor its downstream target molecule and suppressor of autophagy mTOR were influenced by Sph or Lt-Sph treatment, indicating the dispensability of the AKT/mTor signalling pathway in this context (Fig. 3C). Further analysis of markers for endoplasmic reticulum (ER) stress signalling including PKR-like ER kinase (PERK), calnexin, endoplasmic oxidoreducin-1 (Ero1), and the inositol-requiring enzyme 1 (IRE1) also eliminated a

Fig. 4. Influence of Sph and Lt-Sph on PKC activity and IL-10 production in DLBCL cell lines. (A) Measurement of PKC activity in the presence and absence of indicated molar ratios of PS/Sph and PS/Lt-Sph. Shown are means ± SEM, n=2, *p<0.05. (B) Measurement of IL-10 in the supernatant of HBL-1 and U2932 DLBCL cell lines after treatment with the indicated µM concentrations of Sph and Lt-Sph. Shown are means ± SD, n=3, ***p<0.001.
pertinent role of ER stress-related pathways for induction of the observed cell death (Fig. 3C). Thus, U2932 DLBCL responded to Sph and Lt-Sph treatment with increased autophagy-related LC3-II generation, while no detectable changes of this protein were observed with OciLy19, HT, and HBL-1 DLBCL cell lines.

PKC inhibition by Sph and Lt-Sph

The observed differences of DLBCL cell lines to respond to Sph and Lt-Sph persuaded us of investigating PKC signalling as a third potential contributor to Sph and Lt-Sph induced cell death. Recent findings indicated that PKC signalling is an important survival factor of ABC, but not GCB DLBCL [4]. In vitro PKC activity assays demonstrated that both Sph and Lt-Sph inhibited PKC in a concentration-dependent manner, with Sph being more effective than Lt-Sph in competing with the PKC activator PS for binding to the C1 domain (Fig. 4A). Since PKC signalling was shown to induce the production of the downstream effector cytokine IL-10, which promotes proliferation and survival of B cells, quantification of IL-10 was performed by ELISA. While the DLBCL cell lines OciLy19 and HT hardly produced any IL-10 (data not shown), the DLBCL cell lines HBL-1 and U2932 produced IL-10 at physiologically relevant levels. Addition of Sph and Lt-Sph reduced the amount of secreted IL-10 within 4 hours (Fig. 4B). The effect was similar with Sph and Lt-Sph, and HBL-1 cells were more sensitive than U2932 cells in this assay. PKC activity was therefore compromised by Sph and Lt-Sph treatment, and addition of Sph and Lt-Sph reduced the amount of IL-10 release in the IL-10 expressing DLBCL cell lines HBL-1 and U2932.

The PKC activator PS partially rescued HBL-1 and U2932 DLBCL from cell death

To test the relevance of PKC inhibition for Sph and Lt-Sph induced cell death, OciLy19, HT, HBL-1, and U2932 DLBCL cell lines were incubated with 15 µM of the PKC activator PS in order to compete with the PKC inhibitors Sph and Lt-Sph for binding to the C1 domain [22]. As part of the regulatory domain or the amino-terminus of the PKCs, the C1 domain
is present in all PKC isoforms. PS was ineffective on OciLy19 and HT DLBCL cell lines, but HBL-1 and U2932 were partially rescued from Lt-Sph induced cell death in the presence of 15 µM PS. A significant amount of HBL-1 and U2932 cells remained in the G1 phase of the cell cycle 3 days after co-treatment with 5 µM Lt-Sph and 15 µM PS, while the same cells treated with Lt-Sph alone were mostly shifted to the sub-G1 phase (Fig. 5A). PS similarly rescued HBL-1, but not U2932 cells from cell death during co-treatment with 5 µM Sph, again indicating the activity of additional pathways leading to cell death in U2932 cells after Sph treatment. Quantitative analysis of several different experiments demonstrated 20-30% decrease of HBL-1 and U2932 cells in the sub-G1 phase and a similar increase of both DLBCL cell lines in the G1 phase after co-treatment with 5 µM Lt-Sph and 15 µM PS compared to single treatment with 5 µM Lt-Sph alone (Fig. 5B). No significant changes were observed with OciLy19 and HT DLBCL cell lines. Sph-induced cell death was prevented by PS only in HBL-1 cells, with 40-50% less cells in the sub-G1 phase and equivalently more cells in the G1 phase.

**Discussion**

Sphingolipids like S1P, Sph, and ceramide have long been implicated as regulators of cell fate [7, 8]. Ceramide was suggested as a target molecule to combat drug resistance of tumors [49], and an anti-S1P antibody is currently a promising candidate for treatment of solid tumors [50]. Cell culture experiments with DLBCL cell lines that were incubated with Sph and Lt-Sph manifested significant inhibition of cell growth (Fig. 1A), while incubation of cells with similar concentrations of S1P was ineffective (data not shown). These results suggest that Sph accumulation could be an alternative treatment option for DLBCL. To this end we investigated several different signaling pathways that are potentially involved in Sph- and Lt-Sph-induced cell death (Table 1).

Apoptosis induced by Sph has been described before [23, 24, 29], but the physiological relevance remained unclear. The Sph analog FTY720 also induces apoptosis, but only at concentrations similar to Sph, which are never reached during treatment [51]. But how could relevant Sph concentrations been attained in vivo? One possible way would be the pharmacological inhibition of the S1P-lyase. Sph concentrations reach levels after S1P-lyase inhibition up to 100 µmol/kg tissue weight in mice, which would be sufficient for the described cell death inducing effects in aggressive lymphomas [33]. LX2931 is an S1P-lyase inhibitor that passed phase 2a clinical studies for treatment of the autoimmune disease rheumatoid arthritis [52]. Our results suggest that Sph accumulation could be an alternative treatment option for DLBCL. To this end we investigated several different signaling pathways that are potentially involved in Sph- and Lt-Sph-induced cell death (Table 1).

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**Table 1.** Summary of effects of Sph and Lt-Sph on investigated DLBCL cell lines. * Acridine orange staining only; # IL-10 production only; + strong response; o weak response; - no response

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<th>Lt-Sph</th>
<th>HBL-1 Sph</th>
<th>Lt-Sph</th>
<th>U2932 Sph</th>
<th>Lt-Sph</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell growth block</td>
<td>+</td>
<td>+</td>
<td>0</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Apoptosis</td>
<td>-</td>
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<td>-</td>
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<td>-</td>
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<tr>
<td>Autophagy</td>
<td>-</td>
<td>-</td>
<td>0</td>
<td>-</td>
<td>0</td>
<td>+</td>
<td>0</td>
<td>+</td>
</tr>
<tr>
<td>PKC inhibition</td>
<td>-</td>
<td>-</td>
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</tr>
</tbody>
</table>
S1P result in prolonged internalization and functional inhibition of S1PR1 [33], which may also disrupt pro-survival signaling of S1P, at least partially.

Genetic profiling of the cell lines used indicated that OciLy19 and HT belong to the group of GCB-DLBCL, while HBL-1 and U2932 share main characteristics with ABC-DLBCL [35, 54]. This likely explains the different PKC-mediated responses in OciLy19 and HT versus HBL-1 and U2932 DLBCL cell lines since survival of ABC-DLBCL is much more reliant on constitutive PKC signaling than GCB-DLBCL. Intriguingly HBL-1 and U2932 cell lines also produced more Cer than OciLy19 and HT upon incubation with Sph, and Cer production contributed to Sph-induced apoptosis in HBL-1 cells (Fig. 2A). Several reports indicate that Cer activates PKCζ [16, 55], which may contribute to constitutive PKC signaling in ABC-DLBCL. Sph itself was able to interfere with PKC signaling (Fig. 4A) and counteracted pro-survival signals of PKC as a competitive inhibitor of the PKC activator PS. Lt-Sph also inhibited PKC in vitro, but with lower efficacy (Fig. 4A). Lt-Sph was more stable than Sph in cell culture (data not shown), which may explain its similar efficacy as competitive PKC inhibitor in HBL-1 cells and even better efficacy in U2932 cells compared to Sph.

Besides accumulation of endogenous Sph by S1P-lyase inhibition, SphK inhibitors may also be used for DLBCL treatment due to their structural similarities to Sph [34]. Potential effects of those inhibitors certainly need to be tested individually. This study provides the basis to specifically look at apoptosis, PKC-inhibition, and autophagy in DLBCL. Since S1P is typically considered as a pro-survival factor [7], increased cell death observed after treatment with inhibitors of the S1P-producing SphKs are usually assigned to the presence of lower S1P levels. This study however demonstrates that accumulation of the SphK substrate Sph can significantly impair DLBCL growth and survival, suggesting that SphK inhibiting analogues of Sph may be similarly effective. Agents enhancing endogenous cellular Sph levels like LX2931 [52] and Sph analogues like Safingol [56] may achieve their activity not only by altering S1P levels and signaling, but also by inducing apoptosis, PKC-inhibition, and autophagy as shown in this study. Future drug candidates in the sphingolipid field for cancer treatment may therefore shift from inhibitors of S1P signaling to non-phosphorylatable Sph analogues. An autonomous phosphorylation independent inhibitory activity of Sph and FTY720 was also demonstrated for the TRPM7 channel [57], which further supports specific cellular activity of Sph.

Sph induced cell death and blocked proliferation by various means, including apoptosis, autophagy, and PKC inhibition. This variability of cellular functions seems to be important for the overall efficacy of Sph to prevent lymphoma cell growth. Despite striking differences in the efficacy of Sph and Lt-Sph on different DLBCL, the final impact on cell growth and mortality was similar due to the combined activity of different cellular pathways. The broadband impact of Sph across different types of DLBCL introduces sphingosine as a cytostatic drug for lymphoma treatment. Current targeting strategies include the disruption of cancer-specific cellular functions like PKC signalling in ABC DLBCL [35]. Sphingosine accumulation e.g. by S1P-lyase inhibition or by application of Sph analogues provides a different strategy which could be an interesting alternative for combination therapy.

Disclosure Statement

The authors declare no conflict of interests.

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