Recent research has demonstrated that aberrant sphingolipid signaling is an important mechanism of chemoresistance in solid tumors. Sphingosine kinase (Sphk), the primary enzyme metabolizing the sphingolipid ceramide into sphingosine-1-phosphate (S1P), is a primary mediator of breast cancer promotion, survival and chemoresistance. However, to date the mechanism of Sphk-mediated drug resistance is poorly understood. Using the dual sphingosine kinase isozyme inhibitor, SKI-II (4-[4-(4-chloro-phenyl)-thiazol-2-ylamino]-phenol), we explored the effects of sphingosine kinase inhibition on multi-drug-resistant breast cancer cells. We demonstrate that SKI-II alters endogenous sphingolipid signaling and decreases cancer proliferation, survival and viability. Furthermore, pharmacological inhibition of Sphk1/2 induced intrinsic apoptosis in these cells through modulation of the NF-κB pathway. SKI-II decreases NF-κB transcriptional activity through altered phosphorylation of the p65 subunit. Taken together, these results suggest that Sphk may be a promising therapeutic target in chemoresistant cancers.

Introduction

NF-κB is a ubiquitously expressed transcription factor known to regulate a variety of cellular functions, including apoptosis, survival, and differentiation. It consists of five subunits: RelA (p65), c-Rel, RelB, NF-κB1 (p50/p105), NF-κB2 (p52/p100), that are sequestered in the cytoplasm under resting conditions (1). The NF-κB signaling cascade is activated through binding of death receptors on the cell surface, which are members of the tumor necrosis factor (TNF) superfamily that contain a death domain involved in protein-protein interactions (2). Upon ligand binding, the intracellular domain of the death receptor recruits a number of scaffolding proteins forming a ‘death inducing signaling complex’ or DISC. The DISC subsequently binds the inactive form of caspase-8 via the death domain (1). These pro-caspase-8 molecules are then cleaved to form active caspase-8, thus triggering a cascade that activates downstream caspasas, including caspases 3, 6 and 7, which leads to apoptosis (3,4).

Upon TNF binding to the TNF receptor, inhibitory κ kinase (IκK) is activated and phosphorylates IκB, which in turn degrades and releases the transcription factor allowing the NF-κB to translocate to the nucleus and alter gene transcription (2). NF-κB is constitutively activated in cancer and mediates cancer progression and metastasis through regulation of a variety of specific genes, including the inhibitor of apoptosis (IAP) and matrix metalloproteinase (MMP) gene families (5). NF-κB gene transcription is tightly regulated through phosphorylation of the various subunits prior to nuclear binding (6). Each subunit has multiple phosphorylation binding sites, with p50 and p65 believed to the biologically relevant subunits for breast cancer. The p65 subunit is phosphorylated by several intracellular enzymes including protein kinase A (PKA), Akt, GSK1, and mitogen and stress activated protein kinase-1 (MSK1) (7,8).

Altered sphingolipid signaling is known to be a primary mechanism of solid tumor promotion, progression and survival (9,10). The enzyme sphingosine kinase (Sphk) converts the pro-apoptotic sphingolipid ceramide into its pro-survival metabolite sphingosine-1-phosphate (SIP) (11). Sphk induces cancer mitogenesis through both increased production of SIP as well as SIP-independent cellular mechanisms (9). The two isoforms of sphingosine kinase, Sphk1 and Sphk2, are differently regulated and were initially believed to have opposing roles in the cells (12). Recent evidence suggests that the isoforms have similar roles in certain cancers, including breast, and promote cancer survival and chemoresistance (13,14). However, to date the mechanism of Sphk induced drug
resistance is not well understood, in part as a result of the scarcity of small molecular inhibitors targeting sphingosine kinase. Recently, crosstalk between the NF-κB and sphingolipid signaling pathways has been under investigation. It is now known that Sphk1 interacts with TRAF2 on the intracellular portion of the TNF receptor and that TNF can increase Sphk activity (15,16). Sphk also regulates TNF-induced expression of certain adhesion molecules (17). More recently, siRNA knockdown of Sphk1 in A549 lung cancer cells was shown to decrease phosphorylation and translocation of the p65/p50 complex to the nucleus (18). Our laboratory has recently demonstrated that isozyme selective inhibition of Sphk2 with ABC294640 can alter NF-κB transcriptional activity in metastatic cancer cells (19).

As mentioned above, only recently have small molecular inhibitors targeting Sphk been developed (20). In this study, we use the SKI-II, the first commercially available Sphk inhibitor, to pharmacologically target Sphk (21). While initially believed to selectively block Sphk1, it is now known to inhibit both Sphk1 and Sphk2 through competitive inhibition of the intermediate sphingosine (20). SKI-II was utilized to study the effects of sphingosine kinase inhibition on TNF-induced multidrug resistance in metastatic breast cancer cells.

Materials and methods

Reagents. SKI-II (4-[4-(4-chloro-phenyl)-thiazol-2-ylamino]-phenol), etoposide and doxorubicin were purchased from Sigma-Aldrich. Dimethyl sulfoxide (DMSO) was purchased from Fisher Scientific.

Cell culture. ER-negative MCF-7TN-R and MDA-MB-231 cells were cultured as previously described (22,23). Briefly, the MDA-MB-231 cell line was obtained from the American Type Culture Collection (ATCC, Manassas, VA). The MCF-7TN-R cell line is a MCF-7 derived, ER/PR/HER2 negative, chemoresistant breast cancer cell line generated in the laboratory (22,24,25). These cells were derived by growing MCF-7 cells in increasing concentrations of TNFα until resistance was established. The culture flasks were maintained in a tissue culture incubator in a humidified atmosphere of 5% CO2 and 95% air at 37°C.

Lipidomics analysis. Endogenous lipid levels were quantified by mass spectrometry (Lipidomics Core, Medical University of South Carolina) according to published methods (19,22,26). Briefly, cells were collected, fortified with internal standards and extracted with ethyl acetate/isopropanol alcohol. Electrospray ionization followed by tandem mass spectrometry (ESI/MS/MS) analyses of sphingoid bases, sphingoid base 1-phosphates, ceramides, and sphingomyelins were performed on a Thermo Finnigan TSQ 7000 triple quadrupole mass spectrometer.

Clonogenic survival assay. Colony assays were performed as described in previously published methods (22,24). Cells were plated in 6-well plates at a density of 1000 cells per well in full DMEM media. Twenty-four hours later, cells were treated with SKI-II (0.1-10 μM) and then monitored for colony growth. Ten days later the cells were fixed with 3% glutaraldehyde. Following fixation for 15 min, the plates were washed and stained with a 0.4% solution of crystal violet in 20% methanol for 30 min, washed with PBS, and dried. Colonies of ≥30 cells were counted as positive. Results were normalized to DMSO vehicle treated control cells.

Cell proliferation immunoﬂuorescence assay. Proliferation assays were performed as previously described (19,26). Cells were plated at a density of 10,000 cells per well in a 96-well plate in 10% DMEM media and allowed to attach overnight. The following day, cells were treated with DMSO or SKI-II for 24 h. At endpoint, cells were fixed using 100 μl of 3.7% formaldehyde in PBS for 10 min. Formaldehyde was removed and cells were permeabilized using cold methanol for 5 min at room temperature and washed twice with PBS. One hundred μl of 3% FBS in PBS blocking buffer was then added. After 30 min, blocking buffer was removed and cells were incubated for 1 h with Ki-67 (BD Pharmingen, San Diego, CA) antibody. Cells were then washed with PBS and stained with DAPI nuclear stain for 5 min before imaging. For staining quantification, numbers of positively stained cells were expressed as a percentage of the total number of cells per field of view/image. The vehicle control was then set to 1 for comparison with SKI-II treatment.

Cell viability assay. Viability assays were performed as previously described (23,24). Briefly, cells were plated at a density of 7500 cells per well in a 96-well plate in phenol-free DMEM supplemented with 5% FBS and allowed to attach overnight. Cells were then treated with SKI-II (ranging from 10 nM to 100 μM) alone or in combination with etoposide or doxorubicin for 24 h. Following treatment, 20 μl of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT, 5 mg/ml) reagent was incubated in each well for 4 h. Cells were lysed with 20% SDS in 50% dimethylformamide. The pH and absorbances were read on an ELx808 Microplate reader (Bio-Tek Instruments, Winooski, VT) at 550 nm, with a reference wavelength of 630 nm.

Cell death detection ELISA. The induction of apoptosis was determined as previously described using the nucleosome ELISA kit (Roche, Boulder, CO) (19). Cells were plated at 10,000 cells per well in 96-well plates and treated for 24 h with varying concentration of drug. The induction of apoptosis was determined by the amount of nucleosomes in the cytoplasm per the manufacturer’s protocol. Absorbances were read on an ELx808 Microtect plate reader (Bio-Tek Instruments) at 405 nm.

Caspase activation assay. Caspase activation was determined as previously described using the Caspase-Glo 9 Assay (Promega Corp., Madison, WI) per the manufacturer’s protocol (19). Cells were plated at a density of 10,000 cells per well in a 96-well plate in phenol-free DMEM supplemented with 5% FBS and allowed to attach overnight. Cells were then treated with indicated drug for 24 h. Following treatment, Caspase-Glo-9 Buffer, Caspase-Glo-9, substrate and MG-132 inhibitor reagent mix was added to cells in a 1:1 ratio with media for 35 min. Following incubation, luminescence was read in an Autoluminat Plus luminometer (Berthold Technologies, Bad Wildbad, Germany).
Western blot analysis. Protein analysis was performed as described (19,26,29). Cells were plated at 50-60% confluency in 10-cm² culture flasks in 10% DMEM for 48 h. They were then treated with DMSO or SKI-II for 1 h. Following treatment, cells were detached with PBS-EDTA and centrifuged. After removing supernatant, cells were lysed in 60-100 µl lysis buffer (Mammalian Protein Extraction Reagent, MPER, and Halt protease inhibitor (Pierce, Rockford, IL); and PhoSTOP phosphatase inhibitor (Roche). Lysed cells were centrifuged for 10 min at 12,000 G at 4°C to separate protein from cell debris. The supernatants were combined with loading buffer (5% 2-mercaptoethanol in 4X LDS Loading buffer, Invitrogen), boiled for 5 min, and loaded onto a 4-12% Bis polyacrylamide gel (Invitrogen) followed by polyacrylamide gel electrophoresis at 150 V for 1.25 h. Protein was transferred to nitrocellulose membranes using the iBlot (Invitrogen) transfer unit. Nitrocellulose membranes were blocked in 5% milk (BioRad Laboratories) Tris Buffered Saline with Tween-20 (TBS-T) for 1 h at room temperature. Cells were washed briefly with 1X TBS-T (USB, Cleveland, OH) and primary antibodies were diluted in 5% BSA (Bovine Serum Albumin, Sigma-Aldrich, St. Louis, MO) TBS-T according to manufacturer's recommended dilutions. Antibodies for tubulin, p65 (C22B4), phospho-p65 (93H1), IkB (polyclonal), phospho-IκB (14D4), IκB (L570) and phospho-IκB (16A6) (1:1000) were purchased from Cell Signaling Technology, Inc. (Beverly, MA). Membranes were incubated in primary antibody overnight at 4°C with gentle agitation. Secondary infrared conjugated antibodies (LI-COR Biosciences, Lincoln, NE) were diluted in 5% BSA-TBST solution at 1:10,000 and membranes were incubated for 1 h under gentle agitation at room temperature. Membranes were scanned using the LI-COR Odyssey imager and software (LI-COR Biosciences) to detect total and phosphorylated protein levels in cell lysates. Western blots and protein quantification were performed following three independent experiments, with representative blots shown. Protein levels were quantified using densitometry analyses.

Statistical analysis. Statistical analyses were performed as previously described (19,30). Briefly, the IC₅₀ values were calculated from concentration-response curves using GraphPad Prism 5.0 (GraphPad software), using the equation:

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Y = \text{Bottom} + \frac{\text{Top-bottom}}{1 + 10^{\log EC_{50} - X}}
\]

assuming a standard slope, where the response goes from 10 to 90% of maximal as X increases over two log units. Differences in IC₅₀ were compared using Student's unpaired t-test with p<0.05 as the limit of statistical significance. Experiments comparing multiple concentrations to the control were tested with one-way ANOVA with Bonferroni post-test to compare individual concentrations. Combination effects were calculated using the formula: \(\frac{[X1a+X2b]}{[X1a+X2b]},\) wherein X1 is the effect of drug 1 at concentration a, X2 is the effect of drug 2 at concentration b and \([X1a+X2b]\) is the effect of the combination of both drugs at concentrations a and b, respectively (31). All statistical analyses were done using GraphPad Prism 5.0 (GraphPad software).
to the clinical relevance of short-term viability assays, with some studies demonstrating a poor predictive value between these and clinical models (34). Therefore, we determined the effects of sphingosine kinase inhibition on long-term metastatic cancer clonogenic survival to better determine the therapeutic potential of this target. Long-term treatment of SKI-II results in MDA-MB-231 and MCF-7TN-R IC$_{50}$ values of 2.51±1.08 µM (p<0.001) and 2.70±1.05 µM (p<0.001), respectively (Fig. 2B). These results in the low micro-molar range are similar to those of current clinical therapeutics.

Inhibition of cancer proliferation is a necessary characteristic of any clinical chemotherapeutic. The effect of SKI-II on cancer proliferation was determined using Ki-67 immunofluorescence assays. Ki-67 is a nuclear protein expressed only during mitogenic phases of the cell cycle (35,36). As seen in Fig. 3, pharmacological inhibition of SKI-II has potent anti-proliferative properties in MDA-MB-231 cells, decreasing Ki-67 staining by 80.23±4.87% (p<0.001). Of note, SKI-II was less effective in the MCF-7TN-R cell line, decreasing staining by 20.97±5.55% (p<0.001). This suggests that the primary viability effects of SKI-II may not be related to its anti-proliferative effects.

Pharmacological inhibition of sphingosine kinase-1/2 enhances chemotherapeutic-induced intrinsic apoptosis in chemoresistant cancer. Given the decreased anti-proliferative effect of SKI-II in MCF-7TN-R cells compared to MDA-MB-231 cells, we examined whether SKI-II induced apoptosis to exert its anti-viability properties. MCF-7TN-R cells were treated with SKI-II and analyzed for cytoplasmic histone-associated oligonucleosome DNA fragments. As seen in Fig. 4A, treatment with SKI-II resulted in increased apoptosis by 11.90±2.96-fold (p<0.05) compared to vehicle control (Fig. 4A). This finding is interesting given that the MCF-7TN-R cell line is resistant to TNF-induced apoptosis. Given that sphingolipids are known to regulate both intrinsic and extrinsic apoptotic pathways, we investigated the effect of SKI-II on the downstream apoptotic protein caspase-9, a commonly used marker for intrinsic cell death. As seen in Fig. 4B, pharmacological inhibition of Sphk1/2 resulted in a corresponding increase in caspase-9 activity, suggesting that SKI-II-induced apoptosis is at least partially through the intrinsic cell death pathway.

The ability of therapeutic agents to act synergistically is beneficial in the treatment of any cancer. Therefore, we determined whether pharmacological inhibition of Sphk1/2 enhanced the effects of the clinical chemotherapeutics doxorubicin and etoposide. MCF-7TN-R breast cancer cells were pre-treated for 1 h with 10 µM of SKI-II followed by 24 h of treatment with a chemotherapeutic agent and compared to isolated treatment.
with each drug. Following treatment, cells were measured for apoptosis, caspase-9 activity and cell viability.

As expected, etoposide and doxorubicin also induced apoptosis through the intrinsic pathway. Pre-treatment with SKI-II enhanced the apoptotic effect of both etoposide and doxorubicin in MCF-7TN-R cells (Fig. 5). Treatment with SKI-II alone showed modest induction of apoptosis and caspase-9 compared to etoposide and doxorubicin alone. However, treatment with SKI-II and etoposide increased apoptosis by 20.99±4.83 fold (p<0.05) and caspase-9 activity by 4.97±0.74 (p<0.05) fold. Similarly, following SKI-II and doxorubicin treatment, apoptosis increased by 33.33±9.94-fold (p<0.05) and caspase-9 activity by 5.26±1.22-fold (p<0.05). These results suggest that combination treatment with sphingosine kinase inhibition can synergistically enhance the apoptotic effects of clinical chemotherapeutics.

Modulation of the NF-κB signaling through inhibition of Sphk1/2. The ability of Sphk1/2 inhibition to induce apoptosis in these TNF resistant cells led us investigate the mechanism of cell death. Our laboratory previously discovered increased basal p65 transcriptional activity in MCF-7TN-R cells compared to the parental MCF-7 cell line. Therefore, we hypothesized that Sphk1/2 inhibition may be targeting this pathway to restore chemosensitivity to these cells. Both the parental and acquired resistant cell lines were utilized to determine affect of Sphk1/2 signaling on death receptor cascades. Pharmacological inhibition of Sphk1/2 dose-dependently decreased p65 promoter

Figure 3. Varying anti-proliferative effects of Sphk inhibition in de novo acquired drug resistance. (A) MDA-MB-231 cells and (B) MCF-7TN-R cells were treated with vehicle or SKI-II (10 μM) for 48 h. Following treatment, cells were fixed and stained with anti-Ki-67 (red) and nuclei counter stained with DAPI (blue). (A) Representative images of cells at x250. (B) Quantification of cells positive for Ki-67 staining from 10 fields of view per treatment. Data is represented as percent positive cells as compared to total cells normalized to vehicle control. Bars represent mean values ± SEM of three independent experiments.
transcriptional activity in both MCF-7 and MCF-7TN-R cells, suggesting Sphk1/2 is involved in endogenous NF-κB gene regulation (Fig. 6A).

The NF-κB subunit p65 has several phosphorylation sites, which are differentially activated by specific kinase cascades. The PI3K/Akt signaling pathway can activate p65 at the Ser365 phosphorylation site. SKI-II has previously been shown to decrease Akt activation in breast cancer cells. Therefore, we hypothesized that SKI-II decreased phosphorylation of p65 to exert its transcriptional effect. As seen in Fig. 6C, treatment with SKI-II markedly blocked TNF-induced phosphorylation of the p65 subunit at the Ser365 site, without significant altering upstream total or phosphorylation protein levels of IκB. Taken together, these results show that pharmacological inhibition of Sphk1/2 blocks p65 transcription activity through decreased phosphorylation at the Ser365 activation site.

**Discussion**

Acquired chemoresistance is an important mechanism of clinical treatment failure in cancer. Recent studies have demonstrated the importance of sphingolipid signaling in cancer drug resistance (14). In this study, we show that inhibition of both Sphk1 and Sphk2 with the small molecule inhibitor SKI-II decreases viability, proliferation, and survival in both de novo and acquired drug resistance models of metastatic breast cancer. Furthermore, we demonstrate that inhibition of Sphk1/2 can induce apoptosis through the intrinsic cell death pathway. Of note, combination treatments of SKI-II with clinical chemotherapy enhances the apoptotic potential of doxorubicin and etoposide in a synergistic fashion.

While the NF-κB signaling pathway has long been known to promote cancer progression, drug resistance and metastasis, targeting this pathway in an effective and non-toxic fashion has proven difficult. Herein, we demonstrate that pharmacological inhibition of Sphk1/2 blocks NF-κB transcriptional activation through decreased phosphorylation of the p65 subunit. This mechanism likely accounts for the increased efficacy SKI-II and in combination with chemotherapeutics in acquired-drug-resistance breast cancer cells. SKI-II is well-tolerated in vivo, and these results suggest that combination therapy may allow
Figure 6. SKI-II inhibits NF-κB transcriptional activity. (A) MCF-7 and (B) MCF-7TN-R cells were transiently transfected with pGL3-NF-κB-Luciferase plasmid. Following transfection, cells were treated with vehicle, TNFα, SKI-II or SKI-II + TNFα. Cells treated with TNFα were set to 1. Data points and error bars represent the mean ± SEM of three independent experiments. (C) MCF-7 cells were treated with vehicle or SKI-11 (10 µM) for 6 h and analyzed by western blotting for phosphorylated and total IςB. Data shown are representative of three independent experiments.

for a decrease in systemically toxic chemotherapeutics to reach the same anti-tumor effects.

While we have demonstrated one mechanism of SKI-II-induced NF-κB signaling blockade, further investigation is necessary to further understand the crosstalk between the sphingolipid and NF-κB pathways. Sphk/SIP signaling is known to activate a number of intracellular kinase signaling cascades, including Ras, Akt, c-Jun N-terminal Kinase (JNK), Rho and ERK. Many of these kinase cascades are also known to phosphorylate the various Rel family subunits. It is possible that further primary effects of Sphk inhibition on these kinases have downstream secondary effects on the NF-κB phosphorylation. Furthermore, pharmacokinetic studies are needed to elucidate the mechanism of synergy between Sphk inhibition and traditional chemotherapies. The studies presented suggest that pharmacological targeting of Sphk1/2 may be a promising therapeutic avenue in the treatment of drug resistant cancer.

References


