The MitoNEET Ligand NL-1 Mediates Antileukemic Activity in Drug-Resistant B-Cell Acute Lymphoblastic Leukemia

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ABSTRACT

Disease relapse in B-cell acute lymphoblastic leukemia (ALL), either due to development of acquired resistance after therapy or because of de novo resistance, remains a therapeutic challenge. In the present study, we have developed a cytarabine (Ara-C)-resistant REH cell line (REH/Ara-C) as a chemoresistance model. REH/Ara-C-1 was not crossresistant to vincristine or methotrexate; 2) showed a similar proliferation rate and cell surface marker expression as parental REH; 3) demonstrated decreased chemotaxis toward bone marrow stromal cells; and 4) expressed higher transcript levels of cytidine deaminase (CDA) and mito-NEET (CISD1) than the parental REH cell line. Based on these findings, we tested NL-1, a mitoNEET inhibitor, which induced a concentration-dependent decrease in cell viability with a comparable IC50 value in REH and REH/Ara-C. Furthermore, NL-1 decreased cell viability in six different ALL cell lines and showed inhibitory activity in a hemosphere assay. NL-1 also impaired the migratory ability of leukemic cells, irrespective of the chemotherapeutic agent used, in a chemotaxis assay. More importantly, NL-1 showed specific activity in inducing death in a drug-resistant population of leukemic cells within a coculture model that mimicked the acquired resistance and de novo resistance observed in the bone marrow of relapsed patients. Subsequent studies indicated that NL-1 mediates autophagy, and inhibition of autophagy partially decreased NL-1-induced tumor cell death. Finally, NL-1 showed antileukemic activity in an in vivo mouse ALL model. Taken together, our study demonstrates that mitoNEET has potential as a novel antileukemic drug target in treatment refractory or relapsed ALL.

Introduction

In B-cell acute lymphoblastic leukemia (ALL) patients, there is a positive correlation between drug resistance and poor prognostic clinical outcome, independent of age, gender, and initial leukemic burden (Pieters et al., 1991; Kaspers et al., 1997). Emergence of drug resistance after chemotherapy is a common phenomenon in newly relapsed ALL patients, and unfortunately the drug-resistance profile worsens with subsequent relapse (Klumper et al., 1995; Pogorzala et al., 2015). Additionally, throughout disease progression, leukemic cells are constantly modifying their bone marrow (BM) microenvironment to form a leukemic niche, which affords de novo chemoresistance (Colmone et al., 2008). This de novo resistance leads to survival of leukemic cells after therapy, and the residual surviving cell number is a measure of minimal residual disease. Importantly, minimal residual disease is correlated with a poor prognosis in patients (Stow et al., 2010).

To study drug resistance in leukemic cells, we previously developed an in vitro coculture model consisting of human-derived ALL cells cultured in the presence of either primary human bone marrow stromal cells (BMSCs) or primary human osteoblasts (HOBs) (Moses et al., 2016b). Using this model, we developed a protocol to isolate and characterize a subset of phase dim (PD) leukemic cells buried under the BM-derived adherent layer (Slone et al., 2016b). PD cells share the characteristics typical of relapsed/refractory leukemic cells, in that they are quiescent and resistant to chemotherapy, have impaired microRNA biogenesis, and are relevant to the current study on manifested altered mitochondrial metabolism (Moses et al., 2016a,b; Slone et al., 2016a). This drug-resistant phenotype was context dependent and transient, since removing the PD cells from the coculture restored their sensitivity to chemotherapy treatment. In the present study, we have developed a cytarabine (Ara-C)-resistant REH cell line (REH/Ara-C) and cocultured it with BMSCs or HOBs to mimic both the acquired and de novo drug-resistant phenotype generally found in the BM of relapsed ALL patients.

Central to the rationale of this investigation is work by others in which mitochondrial dysfunction has been shown to
be associated with chemoresistance and has been targeted as a strategy for anticancer therapy (Neuzil et al., 2013). Evasion of programmed cell death is one of the major mechanisms of drug resistance and cancer progression (Hanahan and Weinberg, 2011), and targeting mitochondrial function may be a therapeutic option in targeting this population of cells (Fu et al., 2017; Kuntz et al., 2017).

MitoNEET (CISD1) is a mitochondrial outer membrane protein that plays a critical role in maintaining mitochondrial homeostasis and is frequently overexpressed in cancer (Bai et al., 2015). This 2Fe-2S cluster-containing protein was first discovered as a secondary target for the anti-diabetic drug pioglitazone (Colca et al., 2004). In the present study, we have used a small molecule ligand of mitoNEET called NL-1, which was derived from pioglitazone (Geldenhuys et al., 2010) and demonstrated its antileukemic activity in both acquired and de novo drug-resistant cells (Fig. 1). NL-1, pioglitazone, and rosiglitazone belong to the glitazone or thiazolidinedione class of compounds. Additionally, we show that NL-1-mediated death in leukemic cells requires the activation of the autophagic pathway. These results led us to propose the use of NL-1 for the treatment of relapsed ALL, leveraging the vulnerability of the mitochondria as a therapeutic strategy.

Materials and Methods

**Cell Cultures.** Nine human B-cell ALL cell lines were purchased from ATCC or DSMZ, SUP-B15 (CRL-1299), TOM1 (ACC 578), and JM1 (CRL-10423) cells were maintained in RPMI 1640 containing 10% FBS, 0.05 mM β-mercaptoethanol, and 1x streptomycin/penicillin. REH (CRL-8286), NALM-1 (CRL-1567), NALM-6 (ACC 128), BV-173 (ACC 20), RS4;11 (CRL-1873), and SD-1 (ACC 366) were maintained in RPMI 1640 containing 10% FBS and 1x streptomycin/penicillin. Primary CD3+ T cells, CD19+ B cells, peripheral blood mononuclear cells, and bone marrow mononuclear cells were purchased from AllCells (Allcells.com) and maintained in Lymphocyte Growth Medium-3 (Lonza) containing 10% FBS and 1x streptomycin/penicillin. HOBs were purchased from PromoCell (Heidelberg, Germany) and cultured according to the vendor’s recommendations.

A chemotherapy-resistant REH ALL cell line (REH/Ara-C) was developed by exposing REH cells to increasing concentrations of Ara-C starting at 0.2 μM. Following a 1-week period, if cell viability of the Ara-C-treated culture was determined to be greater than 70%, the concentration of Ara-C was increased by 0.1 μM. The resulting REH/Ara-C cell line was maintained in growth media containing 1 μM Ara-C, and all experiments utilizing REH/Ara-C cells were carried out following 48 hours of exposure to 1 μM Ara-C.

Deidentified primary BMSCs were provided by the WVU Cancer Institute Biospecimen Processing Core and the WVU Pathology Laboratory Tissue Bank. The ALL cell lines used in the current study were authenticated by short tandem repeat analysis (University of Arizona Genetics Core, Tucson, AZ) and maintained in 6% CO2 in normoxia at 37°C.

**Cell Proliferation Assays.** ALL cell lines were plated at 5 × 10⁴ cells/well in a 96-well plate. Following treatment with chemotherapy (pharmacologic inhibitors or NL-1), viable cells were quantitated using a Cell Counting Kit-8 (Dojindo Molecular Technology Inc.) according to the manufacturer’s instructions.

**Cell Surface Expression Analysis.** Cell surface proteins were stained with fluorochrome-conjugated antibodies to CD19 (PerCP), CD34 (PE), CD38 (APC), CD44 (FITC), and CD45 (PE-Cy5) in staining buffer (PBS + 10% FBS) for 1 hour at room temperature. Cells were washed with PBS, fixed with 1% paraformaldehyde, and resuspended in PBS for analysis of mean fluorescence intensity by flow cytometry (BD LSR Fortessa). Fluorochrome-conjugated isotype controls were used for measuring nonspecific binding.

**Chemotaxis Assay.** BMSCs or HOBs were plated at 90% confluence in 24-well Transwell plates (Costar). After overnight incubation, untreated ALL cells or pretreated NL-1 (60 μM) cells (1 hour pretreatment) were plated at 1.5 × 10⁵ cells in the top of the Transwell insert. The cells were allowed to migrate toward adherent layers for 4 hours. Migrated leukemic cells were collected, fixed by 2% paraformaldehyde, and counted using flow cytometry. SDF-1 (100 ng/ml) served as the positive control for the chemoattractant and media alone served as the negative control.

**Real-Time Quantitative Reverse Transcriptase Polymerase Chain Reaction.** REH and REH/Ara-C cells were plated at 1 × 10⁶ cells/ml and cultured overnight, and then the cells were pelleted and processed for RNA extraction using an RNaseasy kit (Qiagen) according to the manufacturer’s instruction. RNA (50 ng) was then amplified by real-time polymerase chain reaction using the Power SYBR Green RNA-to-cDNA 1-step kit (ThermoFisher Scientific) with primers specific for cytidine deaminase (CDA), deoxyctydin kinase, ribonuclease reductase M1, and ACTB (VHPS-110) purchased from RealTimePrimers.com, and primers for SCL19A1 (FWD 5’-GAA CCC GTT TGA GCT CGG TA-3’) and REV 5’-CCC GAG AGT CAC TGG TTC ACA-3’) purchased from Integrated DNA Technologies. The gene expression in REH and REH/Ara-C was normalized to ACTB and the fold difference in gene expression between the two cell lines was subsequently analyzed.

**Hemosphere Assay.** SD1 leukemic cells were plated at 1 × 10⁵ cells/well in a 96-well plate and incubated for 4 days to allow sphere formation. Following confirmation of spheroids in culture by light microscopy, the cells were treated with 60 μM NL-1 for 72 hours. At the end of the treatment, the resultant spheroids were counted via microscopy and the images were captured (DM IL LED microscope/camera; Leica). Cells were collected at the end of the study and counted with Trypan Blue (0.4% v/v) to determine the number of live cells.

**Coculture Study.** Coculture conditions have been previously described (Slone et al., 2016b). Briefly, 1 × 10⁶ REH or REH/Ara-C cells were seeded on either BMSC or HOB adherent layers and

![Fig. 1. Structure of NL-1. The thiazolidinedione (TZD) warhead scaffold is shown in blue, alongside pioglitazone and rosiglitazone, which are classic TZDs.](image-url)
maintained in 5% O2. The coculture was fed every 4 days, and on the ninth day the coculture was treated with 60 μM NL-1 for 72 hours. Following exposure, ALL cells that were in suspension were collected [suspended (S) cells]. The ALL cells that were buried under the adherent layer were trypsinized and separated from BMSCs or HOBs by size exclusion using Sephadex G-10 columns (GE Healthcare). These buried ALL cells were designated as PD cells and have been previously described to share relevant characteristics with cells that contribute to treatment-resistant minimal residual disease (Moses et al., 2016a; Slone et al., 2016a).

**Autophagy Detection Assay.** REH cells were plated at 1 × 10^6 cells/ml and then treated with 5 μM chloroquine for 1 hour. Following chloroquine treatment, the cells were exposed to 60 μM NL-1 for 6 hours. At the end of the treatment, the autophagic flux was measured using CYTO-ID Autophagy Detection Kit 2.0 (Enzo Life Sciences) according to the manufacturer's instructions. Briefly, treated cells were washed with 1x assay buffer and resuspended in CYTO-ID green stain solution for 30 minutes at 37°C. Following incubation, the cells were thoroughly washed and cell density was adjusted to 5 × 10^4 cells/ml. Then, 100 μl of this suspension was plated in triplicate in a 96-well plate and the fluorescence intensity was measured with Biotek Plate Reader Synergy HT using a FITC filter (excitation 480 nm, emission 530 nm).

**Cell Death Assay.** Primary immune cells were plated at 1 × 10^6 cells/ml in 24-well plates. The cells were treated with 60 μM NL-1 for 24 hours and then stained using a LIVE/DEAD Fixable Dead Cell Stain Kit (ThermoFisher Scientific) following the manufacturer's instructions. The number of dead cells was then analyzed using flow cytometry.

**In Vivo Study.** All animal procedures were reviewed and approved by the WVU Institutional Animal Care and Use Committee. Ten 6–8 month old female NOD.Cg-Prkdcs-cild129Scid1Scid2Prkdcscid Il2rgtm1Wjl/C2 mice were injected intravenously with 1 × 10^7 luciferase-expressing TOM1 ALL cells. Two days postinjection, the mice were injected with luciferin and the tumor burden was analyzed using an IVIS SpectrumCT (Perkin-Elmer). After confirmation of tumor engraftment, the mice were injected intravenously with 1 × 10^6 cells/ml in 24-well plates. The cells were treated with 60 μM NL-1 for 24 hours and then stained using a LIVE/DEAD Fixable Dead Cell Stain Kit (ThermoFisher Scientific) following the manufacturer's instructions. The number of dead cells was then analyzed using flow cytometry.

**Statistical Analysis.** In vitro experiments were performed in triplicate at least three independent experiments, unless otherwise stated. For studies consisting of more than two treatment groups, statistical significance between groups was determined using one-way ANOVA followed by a post hoc Tukey's test. For the chemotaxis assay, two-way ANOVA followed by a post hoc Sidak's multiple comparisons test was used to determine statistical significance. For the in vivo study, an unpaired one-tailed t test was performed to determine the statistical significance between the NL-1–treated and control vehicle–treated mice. Data are shown as mean ± S.E.M., and P < 0.05 was considered statistically significant.

## Results

**Development of a Chemotherapy-Resistant ALL Cell Line.** The emergence of chemotherapy-resistant tumors presents a major clinical challenge in the long-term treatment of ALL patients. To generate a model to address chemotherapy-resistant disease, the REH ALL cell line was treated with increasing doses of Ara-C over the course of several months to generate a cell line that is resistant to this drug (REH/Ara-C). A dose-response curve using Ara-C was performed to compare the REH/Ara-C cells to the parental REH cells. As expected, there was a dose-dependent decrease in the number of viable REH cells, with an IC_{50} value of 19.5 nM; however, Ara-C had no effect on the number of viable REH/Ara-C cells, even at doses 10 times higher than the IC_{50} value for the parental line (Fig. 2A). The chemotherapy resistance in the REH/Ara-C cell line was specific to that treatment, since those cells remained sensitive to other chemotherapy reagents including methotrexate (Fig. 2B) and vincristine (Fig. 2C), showing IC_{50} values comparable to the parental cell line, REH (Fig. 2D).

**REH/Ara-C Cell Line Shows Increased CISD1 Levels.** Further experiments were performed using the REH and REH/Ara-C cells to assess the functional or expression differences that may contribute to the differential sensitivity to chemotherapy. REH and REH/Ara-C had similar cell proliferation rates; the modest difference noted at day 4 was not statistically significant (Fig. 3A). Analysis of cell surface receptors demonstrated that CD19, CD34, CD38, CD44, and CD45 were present at the same levels in both REH and REH/Ara-C cell lines (Fig. 3B). Interestingly, the chemotactic ability of the REH/Ara-C cells to migrate toward BMSCs, HOBs, and SDF-1 was diminished when compared with the REH parental cell line. However, no difference was observed between the two cell lines when media were used as the chemotaxtractant (Fig. 3C). Gene expression analysis showed a 2.6-fold increase in the number of CDA transcripts (CDA) and a 1.7-fold increase in mitoNEET (CISD1) transcripts (Fig. 3D); however, deoxycytidine kinase (DCK), ribonuclease reductase M1 polypeptide (RRM1), and solute carrier family 19 (folate transporter) member 1 (SCL19A1) transcripts were comparable to the parental cell line.

**NL-1 Shows Anticancer Activity in ALL Cell Lines.** Based on the increased expression of mitoNEET in the drug-resistant cell line, we tested the activity of a mitoNEET ligand, NL-1. Treatment with NL-1 reduced the number of viable cells in both REH and REH/Ara-C cell lines in a concentration-dependent manner (Fig. 4A). The IC_{50} value of NL-1 was comparable in REH (47.35 ± 7.7 μM) and REH/Ara-C (56.26 ± 8.8 μM) cells (Fig. 4B). Furthermore, NL-1 successfully decreased the number of viable cells in a concentration-dependent manner in six additional ALL cell lines (Fig. 4C). SUPB15 was the most sensitive to NL-1 with an IC_{50} value of 29.48 ± 10.66 μM, and NALM6 demonstrated the most resistance to NL-1 with an IC_{50} value of 94.26 ± 2.60 μM (Fig. 4D). In contrast, TOM1, BV173, NALM1, and JM1 all had similar IC_{50} values of around 60 μM for NL-1 (Fig. 4D). Furthermore, we used SD1 ALL cells to form hemospheres for evaluation of NL-1 in a model of tumor stem cell–like behavior. In contrast to the REH cells, SD1 cells will form hemospheres suited for this experimental design to identify compounds, which will target this stem cell population. SD1 cells formed 10–15 spheroids/well. After the spheroids formed, they were treated with NL-1. The NL-1 treatment consistently reduced the size and number of preformed SD1 spheroids (Fig. 5, A and B). Additionally, we found that the number of live cells (Fig. 5C) and the percentage of viability (Fig. 5D) of the cells was reduced when treated with NL-1.

**NL-1 Impairs Chemotaxis in ALL Cells.** Soluble factor gradients driving chemotaxis play an important role in leukemic niche development (Möhle et al., 2000; Colmone et al., 2008). Our initial studies showed that parental REH and drug-resistant REH/Ara-C cells both had comparable chemotaxis profiles, with REH/Ara-C showing only a modest decrease in its migratory capacity. NL-1 pretreatment inhibited the chemotactic ability of both REH (Fig. 5E) and...
REH/Ara-C (Fig. 5F) cells to migrate toward multiple chemo-attractants. This inhibitory NL-1 activity was most pronounced in REH and REH/Ara-C cells migrating toward the BMSCs, although there were also significant differences in all of the conditions. The cells treated with NL-1 showed a dose-dependent decrease in chemotaxis in both the REH (Fig. 5G) and REH/Ara-C (Fig. 5H) cells.

**NL-1 Induces Death in the Drug-Resistant Phase Dim ALL Cell Population.** To model the bone marrow microenvironment, which can provide a site of sanctuary for leukemia, our laboratory has previously developed a coculture model where ALL cells are grown with either primary human BMSCs or HOBs. The leukemia cells form three populations in the coculture: those floating in suspension (the S cells), those loosely adhered to the top of the stromal cell layer, and the cells that bury underneath the stromal cell layer (phase dim cells; PD). The PD leukemic cells consistently show increased resistance to a variety of chemotherapies. To determine whether NL-1 could affect cells that are protected by both BM microenvironment cues and acquired drug resistance, NL-1 was tested in cocultures including either REH or REH/Ara-C cells. Treatment of BMSC and HOB cocultures with NL-1 decreased the number of live cells recovered from the S and PD cell populations (Fig. 6, A and C) and decreased viability of the PD population (Fig. 6, B and D), where the number of live cells were determined with Trypan Blue live staining. In these conditions, the S cells showed a small but significant decrease in the number of live cells.
in the REH/Ara-C suspension cells, but the viability was not decreased.

**NL-1–Mediated Induction of Autophagy.** MitoNEET has been shown to play a role in mitophagy during autophagy (Lazarou et al., 2013). To determine if NL-1 modulates autophagic flux in ALL cells, we used a fluorescent dye that specifically labels autophagosomes. Treatment with NL-1 increased the fluorescence intensity compared with untreated cells, indicating an increase in autophagy (Fig. 6E). The specificity of the assay was confirmed by pretreating the cells with chloroquine, an inhibitor of autophagy. Treatment with ALL cells with chloroquine before the addition of NL-1 slightly reduced the increase in fluorescence (Fig. 6E). To investigate if autophagy is critical in NL-1–mediated induction of death in leukemic cells, we pretreated cells with increasing doses of chloroquine followed by NL-1 and then analyzed cell viability. Treatment with NL-1 by itself caused death in REH cells (Fig. 6F). Pretreatment with 5 μM chloroquine to inhibit autophagy reduced the amount of NL-1–mediated cell death significantly in both the REH and REH/Ara-C cells (Fig. 6F). We repeated this pharmacological inhibitory study in TOM1 and observed a similar reduction in NL-1–mediated death (Supplemental Material).

**NL-1 Has Antileukemic Activity In Vivo.** To determine whether NL-1 would negatively affect the viability of other hematopoietic cells, we treated primary immune cells with NL-1 and found that NL-1 did not substantially decrease cellular viability in those cells (Fig. 7A). NL-1 caused the highest increase in death in CD3 T cells (12.00% ± 0.66%), followed by peripheral blood mononuclear cells (11.00% ± 5.00%), then CD19 B cells (7.57% ± 5.44%), and finally bone marrow mononuclear cells (2.00% ± 1.00%). To determine whether NL-1 would affect leukemic cell growth in vivo, we evaluated NL-1 in a pilot study with a single dose of NL-1, where we injected TOM1 ALL cells expressing luciferase into NOD.Cg−Prkdc<sup>scid</sup> Il2rg<sup>tm1Wjl/SjJ</sup> mice. After confirming leukemic cell engraftment by IVIS luminescent imaging, the mice were treated with 1 mg/kg NL-1 every day for 5 days. The dose was based on our previous pharmacokinetic study (Pedada et al., 2014). Out of the 10 mice, one control mouse did not show tumor progression after engraftment and was not included in the study. On day 14 postinjection, the NL-1–treated group showed significantly lower tumor burden than the control vehicle–treated group (Fig. 7B).

**Discussion**

In the present study, we have developed an Ara-C–resistant leukemic cell line (REH/Ara-C) to model the development of chemotherapy resistance observed in patients with ALL. REH/Ara-C cells overexpress CDA and CISD1 (mitoNEET protein) transcripts. This overexpression did not result in crossresistance to methotrexate or vincristine but was associated with altered migratory capacity toward BMSCs, HOBs, and SDF-1. Targeting the mitochondrial protein mitoNEET (CISD1) with NL-1 decreased the viability in several ALL cell lines with comparable IC<sub>50</sub> values. Moreover, NL-1 efficiently disrupted preformed spheroids in a hemosphere assay and was able to inhibit the chemotaxis of both parental REH and REH/Ara-C cell lines. More importantly, NL-1 was able to specifically decrease the viability of REH/Ara-C cells in a coculture model that models the combination of acquired and de novo drug resistance found in relapsed patients. The NL-1–mediated cell death was attributed to its ability, in part, to induce autophagy since pretreatment with the autophagy inhibitor chloroquine was able to reduce NL-1–mediated cell death. Finally, NL-1 was nontoxic to normal immune cells and showed significant antileukemic activity in a pilot study using a mouse model of ALL.

B-cell ALL relapsed patients are treated with a combination of chemotherapy and hematopoietic stem cell transplantation (Eckert et al., 2013). The antitumor effect of Ara-C varies widely in the clinic and is related to genetic variation in the key genes involved in the uptake, activation, and metabolism of Ara-C within the tumor cells (Lamba, 2009). SCL19A1 is a folate transporter that is responsible for the uptake of Ara-C and has been shown to be downregulated in Ara-C–resistant cells (Gati et al., 1997). Once inside the cells,
Ara-C is activated by deoxycytidylate deaminase (DCK) and its expression has been reported to be decreased in relapsed patients (Kakihara et al., 1998). Ara-C is inactivated by cytidine deaminase (CDA), which has been shown to be upregulated in drug-resistant cells (Schröder et al., 1998). Finally, ribonucleotide reductase M1 (RRM1) catalyzes and maintains...
the intracellular pools of dCTP, which directly competes with activated Ara-C for incorporation within the DNA (Liliemark and Plunkett, 1986). In our study, CDA was upregulated in the REH/Ara-C cell line and may be the cause of resistance to Ara-C. Since the upregulated CDA has little role to play in modulation of vincristine and methotrexate activity, not surprisingly, these drugs showed similar sensitivity in REH/Ara-C compared with REH cells. This lack of crossresistance has been previously demonstrated in T-lymphoblastic leukemia cells that were resistant to Ara-C (Cai et al., 2008). Since there are no clinical therapies available to mitigate the overexpression of CDA, we sought to identify a druggable target for treatment of relapsed ALL patients. Our previous studies have shown that the drug resistance phenotype is accompanied by mitochondrial dysfunction (Moses et al., 2016b). In light of this, we looked at the mitochondrial protein mitoNEET expression in REH/Ara-C cells and found it to be upregulated in the drug-resistant cells compared with parental REH cells.

MitoNEET has been previously described as being overexpressed in cancer cells of epithelial origins (Sohn et al., 2013). Specifically, constitutive overexpression of mitoNEET increases tumor size in breast cancer, and conversely knocking down the expression of mitoNEET decreased proliferation and tumor development in human breast cancer cells (Sohn et al., 2013; Tamir et al., 2015). Furthermore, a bioinformatics study of gene expression analysis has revealed that mitoNEET expression is elevated in several other solid tumors.
Interestingly, autophagy plays a key role in ALL survival and Conversely, overexpression of mitoNEET confers autophagic resistance and promotes tumor growth (Salem et al., 2012). Interestingly, autophagy plays a key role in ALL survival and leukemia transformation in the bone tumor microenvironment (Altman et al., 2011). Several drugs used in ALL therapy use activation of autophagy as a mechanism for inducing death in the leukemic cells. For example, glucocorticoids activate the autophagic machinery to induce death in primary ALL cells (Laane et al., 2009). Furthermore, idarubicin mediated cytotoxic autophagy in ALL cells that could be partially reversed by pretreating cells with pharmacologic inhibitors of autophagy (Ristic et al., 2014). Similarly, everolimus induced autophagy-mediated death in ALL primary cell lines and in NOD/SCID mice xenografted with ALL cell lines, which could be partially reversed by genetic ablation of beclin-1, a key regulator of autophagy (Crazzolara et al., 2009; Neri et al., 2014). In the present study, NL-1 treatment in ALL cells led to increased autophagy, which could be reduced by pretreating cells with an autophagy inhibitor, chloroquine (Kimura et al., 2013). More importantly, inhibition of autophagy by the use of chloroquine also significantly reduced NL-1-mediated cell death in ALL. These studies suggest that the decrease in proliferation and increased cell death of the PD cells may in part be due to involvement of the autophagic pathways. Future studies will focus on elucidating the key mechanisms important for cell death seen with NL-1 treatment.

In addition to survival, autophagy also plays a key role in chemotaxis. Specifically, repression of autophagy facilitates membrane trafficking and compartmentalization of critical proteins to aid in the directional chemotactic migration of cells (Coly et al., 2017). Chemotaxis plays a very important role in the niche development within the BM of leukemia patients (Gómez et al., 2015). Previous studies using chemokine inhibitors like AMD3100 reduced tumor burden and increased leukemic death when used in combination with chemotherapy (Fei et al., 2010). Our group has demonstrated the utility of inhibiting chemotaxis as a mechanism for inducing death in drug-resistant ALL cells (Nair et al., 2018). In the present study, NL-1 was sufficient to inhibit ALL chemotaxis, irrespective of the chemoattractant used in the study. The combination of the autophagy-induced death and chemotaxis might partially explain the in vivo antileukemic activity observed in our study. However, future studies to include evaluation of a stable cell line with impaired autophagy will be required to definitively validate this correlative observation made during our study.

(Stelzer et al., 2011). MitoNEET is an iron-sulfur 2Fe-2S cluster protein located on the outer mitochondrial membrane, which acts as a redox sensor and regulates mitochondrial bioenergetics (Mittler et al., 2019). However, the role of mitoNEET as a potential drug target in ALL had not yet been investigated. In our studies, we have found that mitoNEET, along with CDA, is overexpressed in our acquired drug-resistant cell line, REH/Ara-C. Overexpression in REH/Ara-C cells was not accompanied by changes in cell proliferation or cell surface receptor expression when compared with the parental cell line, REH.

Impaired apoptosis is one of the most crucial phenotypes associated with resistance to different classes of drugs in ALL (Holleman et al., 2003). Since mitochondria modulate the molecular players involved in apoptosis, targeting the same is an attractive strategy for therapy in cancer (Indran et al., 2011). Toward this goal, we have used a selective mitoNEET ligand, called NL-1, a derivative of the antidiabetic drug pioglitazone (Geldenhuys et al., 2010, 2011). The use of NL-1 as an anticancer agent has not been reported, and our studies indicated that NL-1 has significant antileukemic activity. Our data show that NL-1 was effective in decreasing the cell viability of different ALL cell lines and had similar IC₅₀ values in the REH and resistant REH/Ara-C cell lines. We had previously demonstrated the enrichment of ALL cells with overexpression of stem cell markers like SOX2 in hemospheres (Nair et al., 2018). We evaluated the effect of NL-1 in the SD1 ALL cells since these have a propensity to form hemospheres: NL-1 effectively disrupted preformed spheroids in a hemosphere assay, and the cells had significantly decreased viability (Zhang et al., 2015). We found that NL-1 was able to target and induce death in the PD cells of the REH/Ara-C cells in coculture with either BMSCs or HOBs. This observation is very notable since the coculture study mimics the acquired drug-resistant phenotype associated with relapse in ALL patients (Mudry et al., 2000; Fortney et al., 2001; Moses et al., 2016b).

Previous studies have reported that suppression of mitoNEET is accompanied by mitochondrial dysfunction and accumulation of autophagosomes (Sohn et al., 2013). Conversely, overexpression of mitoNEET confers autophagic resistance and promotes tumor growth (Salem et al., 2012). Interestingly, autophagy plays a key role in ALL survival and leukemia transformation in the bone tumor microenvironment (Altman et al., 2011). Several drugs used in ALL therapy use activation of autophagy as a mechanism for inducing death in the leukemic cells. For example, glucocorticoids activate the autophagic machinery to induce death in primary ALL cells (Laane et al., 2009). Furthermore, idarubicin mediated cytotoxic autophagy in ALL cells that could be partially reversed by pretreating cells with pharmacologic inhibitors of autophagy (Ristic et al., 2014). Similarly, everolimus induced autophagy-mediated death in ALL primary cell lines and in NOD/SCID mice xenografted with ALL cell lines, which could be partially reversed by genetic ablation of beclin-1, a key regulator of autophagy (Crazzolara et al., 2009; Neri et al., 2014). In the present study, NL-1 treatment in ALL cells led to increased autophagy, which could be reduced by pretreating cells with an autophagy inhibitor, chloroquine (Kimura et al., 2013). More importantly, inhibition of autophagy by the use of chloroquine also significantly reduced NL-1-mediated cell death in ALL. These studies suggest that the decrease in proliferation and increased cell death of the PD cells may in part be due to involvement of the autophagic pathways. Future studies will focus on elucidating the key mechanisms important for cell death seen with NL-1 treatment.
In summary, we have demonstrated that targeting of mitoNEET by using the first-in-class compound NL-1 is a clinically viable strategy to consider in order to overcome the drug resistance observed in relapsed/refractory ALL patients. This corroborates findings published by other groups, which indicate that targeting mitochondrial respiration in ALL is a tractable therapeutic strategy (Fu et al., 2017; Kuntz et al., 2017). In addition to its antileukemic activity, NL-1 also showed a nontoxic profile when primary immune cells were exposed to it. NL-1 efficiently reduced viability in several ALL cell lines harboring different molecular drivers of leukemogenesis. Induction of death of leukemic cells in a coculture model of acquired and de novo drug resistance indicates the promise of targeting mitoNEET in combination with the standard of care in the treatment of ALL.

Authorship Contributions

Participated in research design: Goldenhues, Nair, Ptikel, Martin, Gibson.
Conducted experiments: Nair, Ptikel.
Performed data analysis: Goldenhues, Nair, Ptikel.
Wrote or contributed to the writing of the manuscript: Nair, Ptikel, Martin, Gibson.

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