Inhibition of Anaplastic Lymphoma Kinase (ALK) Activity Provides a Therapeutic Approach for CLTC-ALK-Positive Human Diffuse Large B Cell Lymphomas

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Abstract

ALK positive diffuse large B-cell lymphomas (DLBCL) are a distinct lymphoma subtype associated with a poor outcome. Most of them feature a t(2;17) encoding a clathrin (CLTC)-ALK fusion protein. The contribution of deregulated ALK-activity in the pathogenesis and maintenance of these DLBCLs is not yet known. We established and characterized the first CLTC-ALK positive DLBCL cell line (LM1). LM1 formed tumors in NOD-SCID mice. The selective ALK inhibitor NVP-TAE684 inhibited growth of LM1 cells in vitro at nanomolar concentrations. NVP-TAE684 repressed ALK-activated signalling pathways and induced apoptosis of LM1 DLBCL cells. Inhibition of ALK-activity resulted in sustained tumor regression in the xenotransplant tumor model. These data indicate a role of CLTC-ALK in the maintenance of the malignant phenotype thereby providing a rationale therapeutic target for these otherwise refractory tumors.

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Introduction

Expression of anaplastic lymphoma kinase (ALK) fusion proteins resulting from chromosomal translocations involving chromosome band 2p23 is a hallmark of anaplastic large cell lymphomas (ALCL) [1]. The most frequently detected ALCL translocation t(2;5) encodes a nucleophosmin (NPM)-ALK fusion protein with constitutive tyrosine kinase activity which is clearly implicated in the pathogenesis of ALCL [2,3].

Diffuse large B cell lymphomas (DLBCL) harbouring ALK fusion proteins were first described in 1997 [4]. With few exceptions these ALK-translocated DLBCLs display a fine granular cytoplasmic ALK-staining characteristic for the fusion of clathrin (CLTC) with ALK caused by the reciprocal translocation t(2;17)(p23;q23) [5,6,7]. These DLBCLs are further characterized by the expression of immunoglobulin light chain kappa or lambda, plasma cell associated antigens CD38 and CD138, and epithelial membrane antigen (EMA), but lack expression of CD30 antigen and many other B- and T-cell markers [4,6,7]. From the published case reports based on approximately 50 patients, these lymphomas seem to be associated with a poor outcome in children and adults compared to both ALK-positive ALCL and ALK-negative DLBCL when treated with current chemotherapy regimens [8,9].

Small molecule inhibitors of the ALK kinase have recently been developed [9,10]. However, their therapeutic potential in ALK positive DLBCL has not been studied so far in part due to the lack of representative preclinical models. We report the characterization of the first CTLC-ALK positive DLBCL cell line (LM1), the establishment of a pre-clinical model to study the role of CLTC-ALK activity in DLBCL lymphomagenesis, and demonstrate that these lymphomas display activation of ALK signalling pathways and are potently suppressed in vitro and in vivo by a selective ALK inhibitor.

Materials and Methods

Ethics Statement

The tissue donor was included in a protocol approved by the Institutional Review Board of the Justus-Liebig University in 1999 that included the use of biopsy material for further biological studies. In accordance, the parents of the patient gave a written informed consent that included the use of tumor material and normal bone marrow for cell banking as well as for the establishment of the tumor cell line and use of the cells for further studies. Procedures involving animals followed National Institutes of Health guidelines and were approved by and done according to guidelines of the Animal Institute Committee of the...
Cell lines and chemicals

The DLBCL cell lines Karpas422 and LM1, the ALCL cell lines SUDHL1 and Karpas299 and the BL cell line DGH75 were grown in medium containing 90% RPMI and 10% FCS supplemented with antibiotics, L-glutamine and HEPES. The cell lines Karpas422, Karpas299, SUDHL1 and DGH75 were obtained from the Deutsche Sammlung von Mikroorganismen und Zellkulturen (DSMZ) repository that performs authentication and genotyping. Cells were maintained in these conditions during the experiments and NVP-TAE684 (hereon TAE-684) was added from a concentrated DMSO stock solution to the 10% serum-containing culture medium. The ALK-inhibitor TAE-684 (CID16036120) was synthesized in N. Gray’s laboratory [10].

Reverse transcriptase polymerase chain reaction (RT-PCR) and sequencing

Total RNA was extracted from cell lines or frozen tumor material with Trizol reagent according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA). cDNA synthesis was performed with 1 µg of total RNA, random hexamers or oligo-dT and Superscript-II/III reverse transcriptase (Invitrogen). Reverse Transcriptase (RT)-PCR conditions and primers were previously described [11]. Additional primers are shown in Table S1. In experiments involving TAE-684, LM1 cells were treated with DMSO or TAE-684 10 nM for 12 h and the RNA isolated using RNAeasy Plus kit (Qiagen) following the manufacturer instructions. cDNA was synthesized using High Capacity RNA-to-cDNA kit (Applied Biosystems). We amplified specific genes using the Fast SYBR Green conditions (initial step of 20 sec at 95°C followed by 40 cycles of 1 sec at 95°C and 20 sec at 60°C). The CT value of the housekeeping gene (RPL13A) was subtracted from the correspondent genes of interest (ΔCT). The standard deviation of the difference was calculated from the standard deviation of the CT values (duplicates). Then, the ΔΔCT values of the TAE-684-treated cells were expressed relative to their respective DMSO-treated cells using the ΔΔCT method. The folds of expression for each gene in cells treated with the drug relative to control treated cells is determined by the expression: 2-ΔΔCT. Results were represented as fold of expression with the standard error of the mean (SEM) for 2 series of duplicates. The CLTC-ALK specific RT-PCR fragment from frozen tumor tissue at the time point of relapse was cloned in the PCR 2.1 TOPO vector (Invitrogen). Sequencing analysis of the CLTC-ALK plasmid was performed on an ABI PRISM 3100 (Applied Biosystems, Foster City, CA) automated sequencing analyzer using standard sequencing methods.

Immunoblotting, phospho-array and flow cytometry

Cell lysates were prepared using 50 mM Tris pH 7.4, 150 mM NaCl and 1% NP-40 lysis buffer. Lysates for nuclear and cytoplasmatic fractions were obtained using a fractionation kit (Biovision) following the manufacturer’s instructions. Protein concentrations were determined using the BCA kit (Pierce). Fifty micrograms of protein lysates were resolved by SDS-PAGE, transferred to nitrocellulose membrane, and probed with the indicated specific primary antibodies: rabbit to Akt (Cell Signaling), rabbit to STAT3 (Cell Signaling), rabbit to p44/p42 MAPK (Cell Signaling), mouse anti RPS6 (Cell Signaling), rabbit anti phosphorylated Akt (Ser473) (Cell Signaling), rabbit anti phosphorylated p44/p42 MAPK (Thr202/Tyr204) (Cell Signaling), rabbit anti phosphorylated RPS6 (Ser235/236) (Cell Signaling), rabbit anti phosphorylated STAT3 (Tyr705) (Cell Signaling) and mouse to ALK (BD Bioscience, San Jose, CA). Membranes were then incubated with a peroxidase-conjugated correspondent secondary antibody. Detection was performed using an ECL detection system. Relative levels of protein phosphorylation in LM1 cells treated with DMSO or TAE-684 10 nM for 24 h were determined using a phospho-array (Proteome Profiler Array ARY003, R&D Systems, Inc., Minneapolis, MN) following the manufacturer instructions. The scanned film image was analyzed using the ImageJ freeware (NIH, Bethesda, MD). The spot density of the proteins of interest was normalized using the spot density of the positive controls. A detailed protocol and localization of the proteins in the array can be accessed in http://www.rndsystems.com/pdf/ary003.pdf. Flow cytometry was performed with a BD FACScalibur (BD Bioscience, San Jose, CA) using CD30-FITC and CD45-APC antibodies for surface staining and ALK-PE for intracellular staining. All antibodies were from BD Bioscience (San Jose, CA).

IGHV mutation analysis

IGHV mutation analysis was performed by multiplex PCR using the BIOMED2 protocol. Sequences were compared with published germ line VH, D, and JH genes using the International ImMunoGeneTics (IMGT) database (http://www.imgt.org). Mutational status was calculated as percent deviation from the closest matching germ line V_H segment.

Single nucleotide polymorphism (SNP) array analysis

The Genome-Wide Human SNP Array 6.0 (Affymetrix, Santa Clara, CA) has been used according to the protocol provided by the manufacturer (Affymetrix). Microarrays were washed and stained with the Fluidics Station 450 (Affymetrix) and scanned with the GeneChip Scanner 3000 (Affymetrix) using the Command Console software (Affymetrix). The Birdseed v2 algorithm was used to genotype tumor samples. Copy number analysis, loss of heterozygosity analysis and segmentation was calculated using Genotyping Console software version 3.0.2 (Affymetrix).

Growth inhibition determination

Cell lines were grown at their respective concentration that were sufficient to keep the untreated cells in exponential growth over the 48 h drug exposure time. We determined cell viability by using a fluorometric resazurin reduction method (CellTiter-Blue, Promega, Madison, WI) following the manufacturer’s instructions. The fluorescence (560 Ex/590 Em) was determined using the 48 h drug exposure time. We determined cell viability by using a fluorometric resazurin reduction method (CellTiter-Blue, Promega, Madison, WI) following the manufacturer’s instructions. The fluorescence (560 Ex/590 Em) was determined using the Synergy4 microplate reader (Biotek Inc, Winooski, VT). Fluorescence was determined for six replicates per treatment condition or controls. A detailed protocol and localization of the proteins in the array can be accessed in http://www.rndsystems.com/pdf/ary003.pdf. Flow cytometry was performed with a BD FACScalibur (BD Bioscience, San Jose, CA) using CD30-FITC and CD45-APC antibodies for surface staining and ALK-PE for intracellular staining. All antibodies were from BD Bioscience (San Jose, CA).

Activated STAT DNA binding assay.

The DNA-binding capacity of STAT3 and STAT3a was assayed by plate-based assay (TransAM, Active Motif, Carlsbad, CA) following the manufacturer instructions. Briefly, 5×10^5 LM1 and Karpas422 cells were treated with TAE-684 10 nM or DMSO control for 4 h.
Five micrograms of cell lysates were added to wells containing pre-
adsorbed STAT consensus oligonucleotides (5'-TTCCCGGAA-
3'). For control treated cells the assay was performed in the
absence or presence of 20 pmol of competitor oligonucleotides
that contains either a wild-type or mutated STAT consensus
binding site. Interferon-treated HeLa cells (5 µg per well) were
used as positive controls for the assay. After incubation and
washing, rabbit polyclonal anti-STAT3a or anti-STAT3
antibodies (1:1000, Active Motif) were added to each well,
followed by HRP-anti-rabbit secondary antibody (1:1000, Active
Motif). After HRP substrate addition, absorbance was read at
450 nm with a reference wavelength of 655 nm (Synergy4, Biotek,
Winooski, VT). In this assay the absorbance is directly
in a multiplex assay. Results are expressed in relative fluorescent
units (RFU) normalized to cell number.

Cell-Cycle-Analysis

Experimentally growing LM1 and Karpas299 cells were
incubated with 10 nM TAE-684 or DMSO for 4, 12 and 24 h.
Cells were fixed with 70% ethanol and incubated for 2 h at 4°C.
After washing with ice cold PBS the cells were incubated with
50 µg/ml RNAse A and 50 µg/ml propidium iodide at 37°C for
30 m. Cell cycle distribution was analyzed with a FACS-Calibur
flow cytometer (BD Bioscience, San Jose, CA).

Detection of apoptotic cells with Annexin-V

Distribution of apoptotic, death and viable cells were deter-
mined by using Annexin-V-PE Apoptosis detection Kit I
according to the manufacturers instructions (BD Bioscience,
San Jose, CA). Briefly, 4×10^5 proliferating LM1 and Karpas299 cells
were treated with DMSO or 10 nM TAE684 for 24 h After
washing with PBS, cells were stained with Annexin-V-PE and 7-
AAD at RT for 15 m. Cells were analysed on a FACS-Calibur
with Cell Quest Pro software (BD Biosciences, San Jose, CA).

Caspase-7 and -3 activity

The activity of caspase-7 and caspase-3 was determined using
the Ape-ONE caspase 3/7 assay (Promega). Cell lines were treated
with TAE-684 10 nM or control for 4 h followed by 1 h exposure
to the pro-fluorescent Z-DEVD-R110 substrate. Activation of Z-
DEVD-R110 by the activity of caspases 3 and 7 allows the R110
group to become intensely fluorescent (Exitation wavelength = 488
nm/ Emission wavelength = 521 nm), which was measured using the Synergy4 micro-
plate reader in four replicates. Caspase-7 and -3 activity was
related to the cell number determined by CellTiter-Blue (Promega)
in a multiplex assay. Results are expressed in relative fluorescent
units (RFU) normalized to cell number.

Cell proliferation

LM1 cell proliferation was determined by measuring incorpora-
tion of the nucleoside analog 5-ethyl-2'-deoxyuridine (EdU) into
newly synthesized DNA following the manufacturer instructions
(Click-it EdU, Invitrogen) with modification for suspension cells.
LM1 cells were treated with DMSO or TAE-684 5, 10 and 20 nM
for 1 h following incubation with EdU reagent for additional 23 h.
Experiment was carried out in 4 replicates. EdU incorporation was
measured by the abundance of a fluorescent product (Amplex
UltraRed reagent, Invitrogen) and normalized to the viable cellular
number determined by dye exclusion (ethidium bromide and
orange G staining, easycount, Immunicom).

Mice xenotransplant models

Six to eight-week old male SCID and NOD-SCID mice were
purchased from the National Cancer Institute (Bethesda, MA) or
from Charles River Laboratories International Inc, (Wilmington,
MA). Mice were subcutaneously injected in the left flank with low-
passage human LM1 and Karpas422 DLBCL cells. Tumor
volume was monitored every other day using electronic digital
calipers (Fisher Scientific, Hampton, NH) in two dimensions.
Tumor volume was calculated using the formula: Tumor Volume
\( = \frac{a \times b^2}{2} \) where \( a \) is the smallest diameter and \( b \) is the largest diameter.
When tumors reached a palpable size (approximately 75 to 100 mm^3), the mice
were randomly assigned to different treatment arms; in conse-
quence these experiments were all performed once tumors had
fully formed in the animals. TAE-684 was dissolved in vehicle
(10% pyrrolidinone, 90% PEG300, both reagents from Sigma,
Saint Louis, MO) and administered by oral gavage. Mice were
weighed twice a week. All mice were euthanized by cervical
dislocation under anesthesia when at least 2/10 tumors reached
15 mm in any dimension that for the cell lines used corresponded
approximately to 5 weeks. Directly after euthanasia, all organs and
tissues underwent careful macroscopic and microscopic (hema-
toyxlin-eosin staining) examination for signs of toxicity.

Immunohistochemistry

Slides were stained using standard procedures using Envision
reagents (Dako, Glostrup Denmark) following the manufacturer
instructions. Microscopic pictures were acquired using a final 400-
X magnification with an Axioscope 40 microscope (Zeiss, Jena,
Germany) corresponding to a 0.5 mm picture diameter at room
temperature with a Color Vision 3 camera (Soft Imaging Systems,
Münster, Germany). Pictures were adjusted in respect of sharpness
and brightness using Adobe Photoshop 5.0 software (Adobe
Systems Inc, San Jose, CA).

Results

Establishment of the CLTC-ALK positive DLBCL cell line
LM1

The cell line LM1 was established from the bone marrow of a
13 year-old girl suffering from a systemic relapse of a CLTC-ALK-
positive DLBCL. The patient initially presented with a rapidly
growing cervical and supraclavicular mass. Histopathological
evaluation demonstrated large ALK-positive lymphoma cells
suggestive of anaplastic large cell lymphoma of T- or O-lineage
(ALKL) and treatment was initiated accordingly [12]. The patient
progressed locally after the first course of chemotherapy and an
additional biopsy was taken. Revision of the histology of the initial
biopsy as well as analysis of the second biopsy revealed the
presence of ALK-positive DLBCL with expression of CD38,
VS38c, CD38 and EMA, fine granular cytoplasmic ALK-staining
and expression of the immunoglobulin kappa light chain as well as
gamma heavy chain (Figure 1). Negativity for CD30, T cell
markers (CD5, CD5, CD43, CD56, CD2, granzyme B) as well as
CD20 and CD79a further confirmed the diagnosis. Molecular
cytogenetics as well as RT-PCR for CLTC-ALK transcripts
revealed t(2;17) (p23;q23) with expression of CLTC-ALK in
the cells of the relapsed tumor (Figure 2A).

Despite subsequent intensive chemotherapy, the lymphoma
progressed again locally. Highly intensive chemotherapy with
autologous stem cell rescue and concomitant local radiotherapy
was then administered, resulting in complete remission. This
was followed by allogeneic blood stem cell transplantation. However,
the patient relapsed 53 days later both locally and in the bone
marrow. The infiltrating lymphoma cells were positive for CLTC-
ALK, and were isolated for cell line derivation. These cells were kept under in vitro culture conditions using RPMI supplemented with penicillin/streptomycin, 4 mM L-glutamine and 20% fetal calf serum in a humidified incubator at 37°C with 5% CO₂.

We determined the ability of these cells to propagate in vitro and whether they maintained the phenotype of the parental tumor. The immunophenotype of the cells in culture (from hereon named “LM1 cells”) was confirmed to be the same as the primary tumor: The cells expressed CD138, VS38c, CD38 and EMA, showed fine granular cytoplasmic ALK-staining and expression of the immunoglobulin kappa light chain as well as gamma heavy chain (Figure 1). Like the primary tumors, LM1 cells were negative for CD30, T cell markers, CD20 and CD79a [Figure 1 and not shown]. The expression of the CLCTC-ALK fusion could be demonstrated by RT-PCR in both the primary tumor and in the LM1 cell line [Figure 2A]. Sequencing analysis indicated the presence of the CLCTC-ALK fusion transcript [Figure 2B].

Immunoblot analysis with an ALK antibody showed exclusive 17ptel-13p13.1, 17q22, 19q, and Xp21.1-q21.31, Xq21.33-q22.1, and a germline identity of only 86.61%.

Bone marrow and the established LM1 cell line detected a number of (p11)x2,inc[cp 15]. SNP analysis of mononuclear cells from the patient showed der(4)t(4;15)(p14;q15),add(7)(q34), der(4)(4;15)(p14q15),add(7)(q34~35)x2,der(9)(9;13)(q24q12)x2,add(17)(p11)x2,inc[cp 15]. SNP analysis of mononuclear cells from the patient bone marrow and the established LM1 cell line detected a number of changes associated to the cell line including chromosomal gain in 1q, 3q13.31qter, 8q, 11p13 and 19p as well as chromosomal loss in 1p, 2q22.1qter, 4q12qter, 7q36.3, 10, 13q11-q21.32, 13q21.33-q22.2, 17p13.1p13.11, 17q22, 19q, and Xp21.1-q21.31, Xq21.33-q22.1, Xq22.3qter. No regions of partial uniparental disomy were identified [Figure 2D and Table S2]. Moreover, 94.7% of the SNPs were determined by flow cytometry in the LM1 cell line before and after engraftment in the NOD-SCID mouse model. The ALCL cell line SU-DHL1 was used as positive control for CD30 and ALK expression while the BL cell line DG75 served as negative control.

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Figure 1. Immunophenotyping analysis of the original lymphoma and the derived cell line LM1. A: Determination (from top to bottom) of ALK, CD138, CD30 and Ig kappa chain expression and distribution by immunohistochemistry in the original tumor (left column), LM1 cell line grown in vitro (middle column) and LM1 xenograft tumor in NOD-SCID mice (right column). B: The expression of CD30 (left column) and ALK (right column) were determined by flow cytometry in the LM1 cell line before and after engraftment in the NOD-SCID mouse model. The ALCL cell line SU-DHL1 was used as positive control for CD30 and ALK expression while the BL cell line DG75 served as negative control.

ALK kinase inhibition induces cell death in LM1 cells in vitro

The selective ALK inhibitor TAE-684 was shown to have activity against NPM-ALK positive ALCL cell lines in vitro and in vivo [10]. In order to determine whether an ALK inhibitor also had activity in CLCTC-ALK positive DLBCL, we exposed LM1 cells to increasing concentrations of TAE-684. The NPM-ALK positive ALCL cell lines KarPas299 and SUDHL1 were used as positive controls [10] while the ALK negative DLBCL cell line KarPas422 served as negative control. In agreement with previous publications [10], SUDHL1 and KarPas299 were susceptible to TAE-684 (GI50 of 21.5 and 30.8 nM respectively) while KarPas22 was resistant (Figure 3A). TAE-684 inhibited the growth of LM1 at low nanomolar concentrations (GI50; 8.5 nM) (Figure 3A).

To further characterize the biological effects of ALK inhibition on the growth and survival of the LM1 cell line, we performed proliferation, cell cycle and apoptosis analysis on cells treated with either TAE-684 or DMSO control. LM1 cells were treated with increasing concentrations of TAE-684 (DMSO, 5, 10 and 20 nM) for 24 h and assessed for proliferation by a nucleoside analog (EdU) DNA incorporation assay. Treatment with TAE-684 decreased the EdU incorporation in LM1 cells (Figure 3B) indicating that exposure to TAE-684 inhibited proliferation. Since different NPM-ALK positive ALCL cell lines have been reported to respond differentially with either apoptosis or G1-cell cycle arrest [10], we wished to determined whether the effect on proliferation was due to preferential cell cycle arrest, cell death or a combination of both. We analyzed cell cycle distribution by flow cytometry DNA deconvolution at 4, 12 and 24 h after treatment. TAE-684 10 nM caused G1 cell cycle arrest at 24 h in KarPas299 cells but not in LM1 (Figure S1). There was no cell cycle arrest in LM1 at any of time points analyzed, suggesting that cell death is the main mechanism for growth inhibition in this cell line. Accordingly, TAE-684 exposure for 24 h induced apoptosis in a dose-dependent manner in LM1 cells as detected by Annexin-V staining (Figure 3C) and caspase-7 and -3 activation (Figure 3D). Apoptosis induction was morphologically confirmed with ethidium bromide and orange G staining under fluorescence microscopy. Collectively, these data suggest that inhibition of ALK kinase activity by TAE-684 reduces the growth of LM1 cells by preferentially inducing apoptosis.

TAE-684 inhibits signalling downstream of CLCTC-ALK

Fusions of ALK have oncoenic potential as their aberrant kinase activity enhances cell proliferation and survival [3]. Similarly to most normal and oncogenic tyrosine kinases, ALK fusions activate many interconnected and redundant pathways. The most relevant and characterized pathways are the ERK, JAK-STAT3 and P3K-AKT pathways [3,13]. To determine what pathways are preferentially affected with TAE-684 in LM1 cells, we performed a phosphoprotein array in these cells treated with DMSO and TAE-684 at 10 nM for 24 h. The most affected protein in the array was STAT3 (Figure 4A and Figure S2). STAT3 phosphorylation in tyrosine 705 decreases 5 fold after TAE-684 (Figure 4A). Additional proteins with significant decreases were: p70S6K (3.2 fold), STAT1 (7.0 fold), STAT13 (2.6 fold), FAK1 (2.4 fold), LCK (2 fold).
fold and STAT5a/b Y699 (2 fold) (Figure 4A and Figure S2).

There were more modest reductions in the phosphorylation of other proteins such as p90RSK, ERK1/2, AKT, c-JUN, STAT1, STAT2 and several members of the SRC family (LYN, HCK, FYN, FRG) among others. We validated some of these changes in an independent experiment using immunoblots (Figure 4B).
addition to changes in AKT, ERK1 and STAT3 phosphorylation following TAE-684 treatment, we found a decrease in phospho-RPS6S235/S236, a protein not included in the array (Figure 4B). In contrast to STAT3 [3], the role of STAT5 in ALK fusion-mediated lymphomagenesis is more controversial. [3,14]. To determine whether STAT3 or STAT5 signalling are functional in CLTC-ALK in DLBCL, we performed DNA binding assays on lysates of LM1 and Karpas422 cells treated with DMSO or TAE-684 10 nM for 24 h. In concordance with the protein levels, the baseline activity of STAT3 was higher in LM1 compared to Karpas422. The concentration of TAE-684 that inhibits 50% the growth of cell lines (GI50) is shown in parentheses. B: Cell proliferation determined by EdU incorporation in DMSO- and TAE-684-treated cells (X-axis, concentration in nM). Results are expressed as percentage to control after normalization to viable cells. C: Apoptosis in LM1 cells measured by Annexin-V (and 7-AAD for nuclear staining) after incubation with DMSO or TAE-684 10 nM for 24 h (one representative of triplicate experiments is shown). D: Caspase-3 and -7 activity in LM1 cells determined after 24 h of exposure to DMSO- or TAE-684- at several concentration (X-axis). Results are expressed in relative fluorescent units (Y-axis) normalized to cell number.

Figure 3. In vitro effect of the ALK inhibitor TAE-684. A: The cell lines SU-DHL1 (ALCL), Karpas299 (ALCL) and LM1 (DLBCL) harboring ALK translocations and the cell line Karpas422 (as negative control) were exposed in six replicates to 8 concentrations (from 1 to 400 nM) of the ALK inhibitor TAE-684 or vehicle control (DMSO) for 48 h and analyzed for viability. Dose-response curves were plotted. The X-axis shows the dose of TAE-684 in nM. The Y-axis shows the effect of TAE-684 as compared to vehicle control on cell viability. The experiment was done in triplicates. The goodness of fit for the experimental data to the median-effect equation (linear correlation coefficient) obtained from the logarithmic form of this equation was equal or higher than 0.90 for each curve. The concentration of TAE-684 that inhibits 50% the growth of cell lines compared to control treated cells (GI50) is shown in parentheses. B: Cell proliferation determined by EdU incorporation in DMSO- and TAE-684-treated cells (X-axis, concentration in nM). Results are expressed as percentage to control after normalization to viable cells. C: Apoptosis in LM1 cells measured by Annexin-V (and 7-AAD for nuclear staining) after incubation with DMSO or TAE-684 10 nM for 24 h (one representative of triplicate experiments is shown). D: Caspase-3 and -7 activity in LM1 cells determined after 24 h of exposure to DMSO- or TAE-684- at several concentration (X-axis). Results are expressed in relative fluorescent units (Y-axis) normalized to cell number.

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Establishment of a CLTC-ALK Lymphoma Model

A

B

DMSO TAE-684

p-AktS473

Akt

p-Erk1T202/Y204

Erk1

p-RPS6S235/S246

RPS6

p-STAT3Y705

STAT3

C

D

E

TAE-684

CLTC-ALK

PI3K

PLCG

Diacylglycerol

AKT

PKC

mTOR

p70S6K

p90RSK

FOXO3a

RPS6

Cdc25A

MYC

BCL2

BCL2L1

Src / Fyn / Yes / Frg / Lck / Hck

Lyn

FAK

RAS

RAF

MEK1/2

ERK1/2

JNK

Fos

c-Jun

Cyclin D

Decreased proliferation / increased apoptosis
DLBCL as NPM-ALK in ALCL.

Induces similar survival and proliferative signalling cascades in that constitutive ALK activity of CLTC-ALK fusion proteins including those in phosphoprotein levels and mRNA abundance, in the CLTC-ALK related pathways with TAE-684 treatment, FOSB, FOSL1, JUN and JUND did not change significantly and MYC (FOSL2, JUNB, CDC25A, CCND1, CCND2, CCND3, BCL2 treated with TAE-684 10 nM for 12 h, we found a decrease in (did not change significantly after TAE-684 in either cell line (Figure 4C). The impact of CLTC-ALK inhibition on the cellular transcriptional activity was determined by the mRNA abundance of several target genes related to these pathways. In LM1 cells treated with TAE-684 10 nM, STAT3 activity levels decreased significantly (37.1 ± 1.8 AU to 20.9 ± 1 AU, p < 0.001) in LM1 cells, but not in Karpas422 cells (8.2 ± 1.6 AU to 7.3 ± 1.2 AU) (Figure 4C). In contrast, the activity of STAT5 did not change significantly after TAE-684 in either cell line (Figure 4C). The impact of CLTC-ALK inhibition on the cellular transcriptional activity was determined by the mRNA abundance of several target genes related to these pathways. In LM1 cells treated with TAE-684 10 nM, STAT3 activity levels decreased significantly (37.1 ± 1.8 AU to 20.9 ± 1 AU, p < 0.001) in LM1 cells, but not in Karpas422 cells (8.2 ± 1.6 AU to 7.3 ± 1.2 AU) (Figure 4C). In contrast, the activity of STAT5 did not change significantly after TAE-684 in either cell line (Figure 4C).

Figure 4. TAE-684 inhibits signalling downstream of CLTC-ALK in LM1 cells. A: Phosphoprotein abundance in LM1 cells treated with DMSO (D) or TAE-684 10 nM for 4 h (T). The heat map represents the relative abundance normalized by the positive control in the proteome array (see also Supp. Fig. 2). The particular protein residue phosphorylated is shown as superscript. B: Western blot for phospho-ALK, ALK, phospho-Erk1, Erk1, phospho-Rps6, Rps6, phospho-STAT3 and STAT3 in cytosol lysates of LM1 cells treated with DMSO or TAE-684 10 nM for 4 h. C: STAT 3 and STAT5 DNA binding activity (Y-axis, arbitrary units, AU) in Karpas422 and LM1 cells treated with DMSO (representing the baseline activity) and after 4 h of exposure to 10 nM of TAE-684. Experiments represent duplicates with SEM. D: Transcript abundance for Fosl2, JUNB, CDC25A, CCND1, CCND2, CCND3, BCL2 and MYC in LM1 cells treated with DMSO (control) or TAE-684 10 nM for 12 h. Data was normalized to RPL13A levels. Data is presented as fold decrease compared to DMSO treated cells. E: Signalling pathways modified in LM1 cells by treatment with TAE-684. Phosphoproteins (transcripts are represented below the double dotted line) with significant decreases by TAE-684 are shown in green boxes, with no changes in yellow boxes and those not determined in white boxes.

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Karpas422 cells (Figure 4C), as determined by the respective DNA binding capacity, whereas the DNA-binding of STAT5 was only slightly higher in LM1 compared to Karpas422 (Figure 4C).

After 4 h of treatment with TAE-684 10 nM, STAT3 activity levels decreased significantly (37.1 ± 1.8 AU to 20.9 ± 1 AU, p < 0.001) in LM1 cells, but not in Karpas422 cells (8.2 ± 1.6 AU to 7.3 ± 1.2 AU). (Figure 4C). In contrast, the activity of STAT5 did not change significantly after TAE-684 in either cell line (Figure 4C). The impact of CLTC-ALK inhibition on the cellular transcriptional activity was determined by the mRNA abundance of several target genes related to these pathways. In LM1 cells treated with TAE-684 10 nM, we found a decrease in Fosl2, JUNB, CDC25A, CCND1, CCND2, CCND3, BCL2 and MYC (Figure 4D) transcript abundance. Other target genes related to these pathways (Bcl2L1, Mcl1, Cdh1, Fos, Fosb, Fosl1, Jun and Jund) did not change significantly under the experimental conditions (data not shown). The changes in the CLTC-ALK related pathways with TAE-684 treatment, including those in phosphoprotein levels and mRNA abundance, are summarized in Figure 4E. Taken together, our data suggest that constitutive ALK activity of CLTC-ALK fusion proteins induces similar survival and proliferative signalling cascades in DLBCL as NPM-ALK in ALCL.

TAE-684 induces complete regression of LM1 xenografts

In order to evaluate the anti-lymphoma activity of TAE-684 in vivo, the LM1 cell line was injected into the right flank of 10 NOD-SCID mice and allowed to form tumors. Once palpable tumors were detected, pairs of mice were randomized to receive either TAE-684 10 mg/kg/day 5 days per week for 2 weeks (n = 5) or vehicle (n = 5). The drug and vehicle were administered by oral gavage. The ALK fusion negative DLBCL cell line Karpas422 was also implanted in NOD-SCID mice (n = 10) and treated in the same way. TAE-684 induced regression of the LM1 tumors by the second week and complete remission by the third week (Figure 5A). Remission was sustained without recurrence of tumors in any of the animals for 13 additional weeks after which the experiment was terminated and the animals sacrificed (Figure 5A). In contrast, Karpas422 xenografted tumors were unaffected by the drug and grew at the same rate as vehicle controls (Figure 5B). In both models, macroscopic and microscopic examination of the animals showed no signs of disease or organ toxicity. The adjusted body weight between treated and controls was similar for Karpas422 and LM1 animals (Figure 5C).

Discussion

Here we describe the establishment of the first CLTC-ALK positive DLBCL cell line from the bone marrow of a patient with chemotherapy-resistant lymphoma. This cell line, LM1, carries the same phenotypic and genotypic characteristics as the malignant cells from the subject. LM1 forms tumors in mice with a similar growth pattern compared to other established mouse xenograft models of human DLBCL. LM1 can thus serve as pre-clinical testing platform for the role of CLTC-ALK in lymphomagenesis and developing molecular targeted therapy approaches for CLTC-ALK positive DLBCL.

Our data gained from pharmacological inhibition of ALK-activity in vitro and in vivo suggest that CLTC-ALK mediates DLBCL lymphomagenesis and maintenance by constitutive ALK kinase activity. This observation is in line with data indicating that CLTC-ALK transforms fibroblasts as efficiently as other ALK-fusion proteins [15]. Additionally, our data lend further support to the notion that ALK-fusion proteins confer high oncogenic potential to transformed cells of different origin independently of the fusion partner and induce both B- and T-cell lymphomas in transgenic mice [3,16].

Several small molecule kinase inhibitors have been developed blocking ALK-kinase activity and signal transduction in a concentration-dependent manner [10,18,19]. This development opens the possibility of targeted therapy for ALK-positive malignancies. Patients with ALK-positive ALCL have a good overall survival due, in part, to effective relapse strategies including immunotherapeutic approaches [17,18]. In contrast to ALCL, the available reports suggest that ALK-positive DLBCL is often a chemorefractory disease associated with a poorer outcome [8,9]. These patients might, therefore, be candidates for clinical trials with ALK-inhibitors. The high in vitro and in vivo sensitivity of LM1 cells to ALK inhibition supports the rationale testing these compounds for ALK-positive DLBCLs.

NPM-ALK-positive cells show activation of signaling pathways, such as Src kinases [19], PI3K-AKT [20], ERK [21] and STAT3 and 5 [13,14,22]. Functional studies suggest a pivotal role of STAT3 and the PI3K-AKT pathway in NPM-ALK mediated lymphomagenesis whereas a role for STAT5 is more controversial [13,14,22]. While ectopic expression of CLTC-ALK in fibroblasts induced less STAT3 phosphorylation than other ALK-fusion proteins [15], a recent immunohistological study detected ubiquitous STAT3 hyperphosphorylation in two CLTC-ALK positive DLBCL cases compared to ALK negative DLBCL [23]. In our study CLTC-ALK positive DLBCL cells exhibited constitutive STAT3 activity as well as activation of Akt and ERK. Inhibition of ALK-activity decreased the activity of these three signaling pathways in LM1 cells suggesting that CLTC-ALK employs similar signaling cascades than NPM-ALK.

Taken together, our data demonstrate that LM1 is a bona fide model of the DLBCL subtype featuring the CLTC-ALK translocation and indicate that growth of CLTC-ALK positive DLBCL is dependent on ALK kinase. Patients diagnosed with ALK positive DLBCL may, therefore, be candidates for therapeutic trials of ALK inhibitors. The incorporation of ALK status determination into the histopathological characterization of DLBCL could help identifying these patients more readily.
Figure S1 Cell cycle analysis. LM1 and Karpas299 (positive control) cells were assessed for cell cycle distribution by propidium iodide staining and flow cytometry after treatment with TAE-684 10 nM or DMSO for 24 h. One representative experiment from triplicates is shown. (TIF)

Figure S2 Phosphoprotein array. Scanned image of the phosphoprotein array in LM1 cells treated with DMSO (left) or TAE-684 10 nM for 4 h (right). Certain proteins of interest with the correspondent phosphorilated residue are identified. A complete map of the array can be found at http://www.rndsystems.com/pdf/ary003.pdf (TIF)

Supporting Information

Figure S1 Cell cycle analysis. LM1 and Karpas299 (positive control) cells were assessed for cell cycle distribution by propidium iodide staining and flow cytometry after treatment with TAE-684 10 nM or DMSO for 24 h. One representative experiment from triplicates is shown. (TIF)

Figure 5. Effect of the ALK inhibitor TAE-684 on the growth of LM1 and Karpas422 xenografts. A: Tumor growth plot in LM1 xenografted mice treated with vehicle (blue circles) or TAE-684 at 10 mg/kg/day 5 days per week for 2 weeks (red circles). The mice were followed without treatment until the tumors reached 1500 mm³ when the mice were killed (beginning of week 5). The treated mice were followed without further treatment for a total of 16 weeks when they were also killed. The treated mice had no macro or microscopic evidence of tumor relapse. The Y-axis indicates tumor volume (in mm³) and X-axis days of treatment. B: The same experiment as in panel A but using Karpas422 mice. In this case all the mice were killed when the tumors reached 1500 mm³ (at the end of week 5). C: Adjusted body weight at week 5 in the Karpas422 and LM1 mice for the control and TAE-684-treated animals. The error bars represent the SEM.
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Table S1 Additional RT-PCR primers.

Table S2 Chromosomal gains and losses in the LM1 cell line. Chromosomal gain and losses were determined by SNP array. The cytoband and linear position (bp) are in accordance to the NCBI Build 36.1.

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