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$Bcl-x_L$ inhibition enhances Dinaciclib-induced cell death in soft-tissue sarcomas.

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Abstract:

Soft-tissue sarcomas (STS) are an uncommon and heterogeneous group of malignancies that result in high mortality. Metastatic STS have very bad prognosis due to the lack of effective treatments. Dinaciclib is a model drug for the family of CDK inhibitors. Its main targets are cell cycle regulator CDK1 and protein synthesis controller CDK9. We present data supporting Dinaciclib ability to inactivate in vitro different STS models at nanomolar concentrations. Moreover, the different rhythms of cell death induction allow us to further study into the mechanism of action of the drug. Cell death was found to respond to the mitochondrial pathway of apoptosis. Anti-apoptotic Bcl-x_L was identified as the key regulator of this process. Bcl-x_L showed a slower decay curve after protein synthesis disruption that in tolerant cell lines was enough to delay apoptosis, as its action cannot be countered by the relative low levels of pro-apoptotic BH3 proteins BIM and PUMA. Combination of Dinaciclib with BH3-mimetics led to quick and massive apoptosis induction in vitro, but in vivo assessment was prevented due to liver toxicity. Additionally, Bcl-x₁ inhibitor A-1331852 also synergized with conventional chemotherapy drugs as Gemcitabine. Thus, Bcl-x_L targeted therapy arises as a major opportunity to the treatment of STS.

Keywords: CDK inhibitors, soft-tissue sarcoma, apoptosis, Bcl-x_L, BH3-mimetics

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Introduction

Soft-tissue sarcomas (STS) are a group of tumors derived from mesenchymal precursors with scarce incidence and rich variability (1). Tumors arising from non-epithelial extraskeletal tissue are generally accounted as STS (2). There has been much improvement in the understanding of the drivers of STS entities: (i) STSs driven by specific chromosome fusions leading to generation of anomalous transcription factors (like *FUS-CHOP* in myxoid liposarcoma) or chromatin remodelers (*SS18-SSX* in synovial sarcoma); (ii) STS that rely on specific mutations (*KIT* in gastrointestinal stromal tumors) and (iii) other STS driven by more complex genomic rearrangements (like leiomyosarcomas or some fibrosarcomas) (3,4).

STS incidence is difficult to estimate due to their variability, and some reports claim that the usual figures could be underestimations (1,5). Clinical prognosis and therapeutic outcome is also highly variable in STS (2). When it is possible, the complete clinical resection make full recovery achievable. However, almost half of the patients will develop metastatic disease. Five-year survival rates are still below 50 %. So, the weight of STS in total cancer death toll is clearly disproportionate to its incidence (4,6).

Thus, STS can benefit for new therapeutic approaches (6). Among the molecular targeted drugs in development, the group of Cyclin-Dependent Kinases (CDKs) inhibitors is one of those concealing major interest (7). CDKs constitute a wide family of Ser/Thr protein kinases that require binding with cyclins to act. This coupling enables a complex panorama of interactions that keep track on the activation/suppression of specific pathways during cell cycle (8). Several CDK inhibitors have been identified and tested *in vitro* as anti-cancer agents (7,9). The initial aim of CDK inhibitor strategy was the disruption of cell cycle sequence-of-events in order to induce cell death (9,10). But it was soon understood that CDKs exert more powerful effects over other processes of cell physiology like transcription regulation, RNA splicing or protein folding (9).

Dinaciclib is a promising CDK inhibitor, extensively proved pre-clinically (11). Its known affinities encompass CDK1 (IC₅₀=3 nM), CDK2 (IC₅₀=1 nM), CDK5 (IC₅₀=1 nM) and CDK9 (IC₅₀=4 nM) (12). CDK2 is a key cell cycle regulator in meiosis but its significance in somatic cells is relative (8). CDK5 role is less clear. It has been associated with neuronal metabolism via multiple interactions, including proteins from

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the cytoskeleton (13). Most studies concerning Dinaciclib activity have been focused on the CDK1 control of mitotic entry and CDK9 regulation of gene transcription (14–16). CDK9-dependent down regulation of anti-apoptotic Bcl-2 family member Mcl-1 is commonly regarded as the main mechanism of action of this drug (17,18). Some Phase I clinical trials (mostly in pediatric leukemia) have also been performed with Dinaciclib. Anti-cancer activity was found to be encouraging, but not sufficient for planning monotherapy treatments. Further use of Dinaciclib is thought to rely on combination therapies (14,19,20).

Combination therapies constitute a hot spot in oncology research. It has become clear its benefits avoiding tumor evolution in favor of drug resistant phenotypes (21). Moreover, combination therapies work better than monotherapy even in the absence of synergistic behavior (22). BH3-mimetics are a new class of anti-cancer drugs particularly interesting for these combinations. They are aimed to disturb the balance of the different proteins of the Bcl-2 family, thus favoring apoptosis triggering (23,24). Alone, BH3-mimetics have been successfully used in chronic lymphocytic leukemia since the FDA approval of Venetoclax (25). BH3-mimetics work better when the cells are already undergoing an apoptotic signaling process that has been compensated by expression or activity changes in the Bcl-2 family of proteins. Cells became addicted to these compensative mechanisms creating then an Achilles' heel for cancer cells (24). Recent screenings are showing that BH3-mimetics boost the cytotoxic potential of a panoply of chemicals, including CDK9 inhibitors (26).

Our aim in the present study is to seek the suitability of Dinaciclib in a series of STS *in vitro* models as cell death inductor, fully characterizing the cellular response to treatment. We have found that Dinaciclib is capable of inducing cell death as single agent. The cellular context, particularly the Bcl-2 family balance, at every model is decisive for the precise behavior after Dinaciclib incubation. Our data support that Bcl- x_L inhibition status is central for treatment tolerance. Moreover, Bcl- x_L specific inhibitors synergize with Dinaciclib to avoid such tolerance.

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Materials & Methods

Cell culture.

Cell lines (**Table 1**) were cultured in RPMI-1640 supplemented with 10 % (v/v) heatinactivated FBS, 1 % (v/v) Penicillin-Streptomycin and 1 % (v/v) HEPES (Gibco). All cell lines were incubated in an AutoFlow NU-4750 Incubator (NuAire) at 37 °C in a humidified atmosphere of 5 % CO₂ in air. Otherwise specified, plasticware were purchased from Jet BioFil.

Cell line	Origin	Provider	
SW982	Synovial Sarcoma	ATCC	
1273-99	Synovial Sarcoma	Dr. Carmen de Torres, <i>Hospital Sant Joan</i> <i>de Deu</i> (Esplugues de Llobregat, Spain)	
SK-LMS-1	Leiomyosarcoma	ATCC	
SK-UT-1	Leiomyosarcoma	CLS	
402-91	Myxoid Liposarcoma	Prof. Roberto Mantovani (Università degli Studi di Milano, Milan, Italy)	
1765-92	Myxoid Liposarcoma	Prof. Roberto Mantovani	
SW872	Dedifferentiated Liposarcoma	ATCC	
HT-1080	Fibrosarcoma	ATCC	

Table 1: List of cell lines employed in the study.

Authenticity of the cell lines was routinely confirmed by STR profiling analysis done at qGenomics SL (Esplugues de Llobregat, Barcelona, Spain).

Exponential growing cells were used for all experiments. Depending on cell size and proliferation rates, 80,000-180,000 cells were seeded in 12-well plates (or 1-2 million cells in P100 dishes) and allowed to grow for 24 h prior to treatment. Cells were routinely observed and photographed by means of an IX70 inverted epifluorescence microscope (Olympus).

For 3D cultures, we seeded 80,000 cells mixed in 1:1 dilution of medium and Matrigel[®] (Corning Inc.) in 24-well plates already coated with a solidified layer of pure Matrigel[®]. Cells were kept in these conditions for 72 h to allow microsphere formation prior to

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treatment. Cell-permeable DNA-binding dye Hoechst-33342 (Molecular Probes) was added ($10 \mu g/mL$ final concentration in medium) for nuclei visualization.

Chemicals and treatments.

Dinaciclib (CAS 779353-01-4) was purchased from Selleckchem and diluted in DMSO (Sigma-Aldrich) to a 100 μ M concentration. Used dilutions were made in complete medium. Pan-caspase inhibitor N-benzyloxycarbonyl-Val-Ala-Asp(O-Me) fluoromethyl ketone (z-VAD-fmk, CAS 187389-52-2) was purchased from ApexBio and diluted in DMSO to form a 50 mM stock solution. Used dilutions (40 μ M) were made in complete medium. Generic BH3-mimetic ABT737 (CAS 852808-04-9) was obtained from ApexBio and diluted in DMSO to an initial 10 mM stock. The Bcl-x_L specific inhibitor A-1331852 (CAS 1430844-80-6) was purchased from ChemieTek and a 3 mM initial stock solution in DMSO was employed. The nucleoside analog Gemcitabine (CAS 122111-03-9) was obtained as hydrochloride salt from LC Laboratories and diluted in DMSO to form a 1 mM stock. In every case, further dilutions were made taking into account not to surpass 0.1% (v/v) content of DMSO in media.

Proliferation and long-term clonogenic assay.

For proliferation assessment, 3,000-5,000 cells were seeded in 96-well plates and treated with increasing concentrations of Dinaciclib, ABT737, A-1331852 or combinations of drugs for variable times up to 72 h. At desired time points, cells were incubated with WST-1 reagent (Roche) diluted 1:10 in complete medium up to 2 h in the incubator. Soluble formazan salt production from WST-1 was measured by absorbance (λ =440 nm) in a PowerWaveXS plate reader spectrophotometer (BioTek Instruments). IC₅₀ values were calculated using Prism software (GraphPad).

Long-term analyses of post-treatment survival were performed by seeding 180,000-300,000cells in 6-well plates. Cells were cultivated with 25 nM Dinaciclib for 72 h prior drug wash-out and kept in culture up to a week. At that point cells were transferred to new plates to recover Dinaciclib Resistant strains or were stained with 0.5 % (w/v) Crystal Violet (Sigma-Aldrich) solution in 1:5 Methanol:PBS for 20 min and washed with water. Images reflect representative results of at least three independent experiments.

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siRNA experiments.

Cells were transfected using DharmaFECT (Dharmacom, GE HealthCare) following manufacturer's instructions. ON-TARGETplus Non-Targeting Control Pool (Dharmacom) was used as reference and customized siRNA sequences were purchased from Sigma-Aldrich: 5'-AAUAACACCAGUACGGACGGG-3' for Mcl-1 and 5'-ACAAGGAGAUGCAGGUAUUUU-3' for Bcl-x_L.

Flow cytometry.

For the simultaneous quantification of plasma membrane integrity and mitochondrial transmembrane potential ($\Delta \psi_m$), living cells were collected and stained with 1 µg/mL propidium iodide (PI, Molecular Probes) and 40 nM of DiOC₆(3) (DiOC, Sigma-Aldrich) for 30 min at 37 °C. DiOC is a cyanine derivative that accumulates in functional mitochondria and dissipates when $\Delta \psi_m$ is lost. PI is a cell impermeable cationic stain (DNA intercalating) that accumulates in cell nuclei once the plasma membrane is no longer a barrier. So, we can identify cells committed to apoptosis (low levels of DiOC but still excluding PI), cells that suffered sudden necrosis (positive to PI but still retaining DiOC labeling) and cells that progress from apoptosis to secondary necrosis (negative to DiOC and positive to PI). Live cells (positive to DiOC and negative to PI) were not represented.

For cell cycle analysis, cells were fixed in 70 % ice-cold ethanol and labeled with 50 μ g/mL PI in PBS containing 500 μ g/mL RNase (Sigma-Aldrich). This approach also allows cell death quantification as subG₁ events. Cytofluorometric determinations were performed employing a Gallios flow cytometer and data were statistically evaluated using Kaluza software (Beckman Coulter). Only the events characterized by normal forward scatter (FSC) and side scatter (SSC) parameters were included in sub-sequent analyses.

Western blot analysis.

Cells were lysed with RIPA Buffer (ThermoFisher Scientific) containing Protease Inhibitor Cocktail Tablets and Phospatase Inhibitor Cocktail Tablets (Roche). Lysates were sonicated and centrifuged at 13,000 rpm for 20 min (at 4 °C). Protein content was determined with BCA assay (Pierce Biotech.). Lysate aliquots (50 µg) were resolved by

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8-12 % SDS-PAGE and transferred onto nitrocellulose membranes (Bio-Rad Laboratories). After blocking with 5 % skimmed milk in PBS containing 0.2 % (v/v) Tween-20 (Sigma-Aldrich) for 1 h (RT), membranes were incubated overnight at 4 °C with the appropriate primary antibody dilution (**Table 2**).

Target	Provider	Catalog number	Dilution
Noxa	Calbiochem	OP180	1:500
p16 ^{INK4a} (CDKN2A)	Abcam	ab108349	1:1000
Bax	Cell Signaling	5023	1:2000
Дах	Technology	5025	
p21 ^{Waf1/Cip1}	Cell Signaling	2947	1:1000
P21	Technology	2)41	
PUMA	Cell Signaling	12450	1:1000
IUMA	Technology		
Bmf	Abcam	ab181148	1:2000
Bak	Cell Signaling	12105	1:2000
Dak	Technology	12105	
Bcl-2	Cell Signaling	4223	1:1000
DCI-2	Technology	7223	
Bcl-x _L	Cell Signaling	2764	1:2000
Del AL	Technology		
CDK1 (cdc2)	Cell Signaling	9116	1:1000
	Technology		
Mcl-1	Cell Signaling	4572	1:1000
	Technology		
Cyclin A2	Cell Signaling	4656	1:1000
	Technology		
Chk1	Cell Signaling	2360	1:1000
U	Technology		
XIAP	BD Transduction	610762	1:5000
	Laboratories		
Cyclin B1	Cell Signaling	4138	1:1000
- 5 -	Technology		
BIM	Cell Signaling	2933	1:1000
	Technology		
CDK9	Cell Signaling	2316	1:1000
	Technology		
PARP	Cell Signaling	9542	1:1000
	Technology		
phospho(Ser10)-	Cell Signaling	9706	1:1000
Histone H3	Technology		

 Table 2: List of primary antibodies employed for western blotting and/or immunofluorescence.

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Blots were then incubated at room temperature (RT) for 1 h with Goat anti-Rabbit/anti-Mouse IgG (H+L) horseradish peroxidase–conjugated secondary antibodies (#31460 and #31430, Thermo-Fisher Scientific) and bands were revealed in Hyperfilm ECL (Amersham) by enhanced chemiluminescence employing ECL Western Blotting Substrate (Pierce Biotech.). Immunodetection of β -actin (#ab49900) or α -tubulin (#ab28439, both from Abcam) were used as loading reference.

Immunofluorescence.

Cells (30,000-80,000) were seeded over glass coverslips placed into 24-well plates prior to treatment. At desired end-points, cells were fixed by 30 min incubation in 4 % (w/v) paraformaldehyde (Sigma-Aldrich) in PBS. Cells were sequentially permeabilized with 0.1 % (w/v) SDS in PBS (USB Corp.) and subjected to antigen blocking with 10 % (v/v) FBS in PBS (both for 20 min). Primary antibodies employed were Bax, $p21^{Waf1/Cip1}$, $p16^{INK4a}$ and phospho(Ser10)-Histone H3 (**Table 2**). Secondary antibodies were Goat anti-Mouse or anti-Rabbit IgG (H+L) Secondary Antibodies, conjugated either with Alexa Fluor[®] 488 or 594 (ThermoFisher Scientific: #A-11001, #A-11005, #A-11008 and #A-11012). DNA counterstaining was performed upon 10 min incubation with 5 µg/mL solution of DAPI (Molecular Probes). Photographs were taken with either an Axio Observer.Z1 (Carl Zeiss AG) or a DM6000B (Leica Microsystems) epifluorescence microscope. Images were analyzed with Image J software (freely available from the National Institutes of Health at the address https://imagej.nih.gov/ij/).

In vivo experimentation.

For animal experimentation, Dinaciclib was purchased to ApexBio as powder and formulated in 20 % hydroxypropyl beta-cyclodextrin (HP β CD, Sigma-Aldrich) in deionized water. The formulation was stored at 5 °C, used within 7 days and warmed to RT and vortexed for 3 sec before intraperitoneal (i.p.) administration. ABT737 was also obtained from ApexBio and formulated in 30 % propylene glycol, 5 % Tween 80, 10 % DMSO, 3.3 % dextrose (all from Sigma-Aldrich) in water. Aliquots were kept at RT and vortexed before i.p. administration. A-1331852 was provided by Abbvie under Material Transfer Agreement and formulated in 60 % Phosal 50 PG (standardized phosphatidylcholine, PC, concentrate with at least 50 % PC and propylene glycol; gently provided by AbbVie), 27.5 % poly-ethylene glycol (PEG) 400, 10 % ethanol, and

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2.5 % DMSO (all from Sigma-Aldrich). First, A-1331852 powder was suspended in DMSO and ethanol until a uniform cloudy suspension was obtained. PEG 400 and Phosal were then added and the solution was mixed by vortexing. The solution was allowed to sit for approximately 30 min after adding all the excipients helped to achieve a clear solution. A sonicator was also used for less than 10 min. Formulated compound was stored in an amber bottle at room temperature to protect it from light. A-1331852 formulation was administered *per os* (p.o.) by using an oral gavage. Dosages and administration regimes tested are summarized in **Supplementary Figure 5A & C**.

Female Athymic (Hsd:Athymic Nude $Foxn1^{nu}$) mice (n=20) were purchased at Envigo. Female SCID[®] Beige (CB17.Cg-*Prkdc^{scid}Lyst^{bg-J}*/Crl) mice (n=40) were acquired at Charles River. Animal care procedures were followed according to the Institutional Guidelines for the Care and Use of Laboratory Animals. Ethics approval was provided by the locally appointed ethics committee. IDIBELL animal facility abides to the Association for Assessment and Accreditation of Laboratory Animal Care (AAALAC) regulations.

Five million (5×10^6) SK-LMS-1 cells in 1:1 RPMI:Matrigel[®] were subcutaneously (s.c.) injected in the right dorsal flank of the mice. Tumor growth was followed by a caliper and volume was extrapolated using the formula vol= $(L \times l^2)/2$ were "L" refers to the major and "I" to the minor diameter measured. Drug treatments started when tumors reached a volume of approximately 100 mm³. Mice were blindly randomized into the different groups by using a 6-faced dice.

After the end of the experiment, tumors and organs of interest were extracted and fixed in 3.7 % Formol solution in PBS (Sigma-Aldrich) for 24 h. Afterwards, samples were paraffin embedded and prepared for Hematoxylin-Eosin (H&E) staining following standard protocols. Microscopical imaging was performed by means of a Nikon Eclipse 80i workstation. Photographs were analyzed using GNU Image Manipulation Program (GIMP) software (freely available at <u>http://www.gimp.org</u>).

Statistical Analysis.

Unless otherwise stated, experiments were performed thrice. Data were analyzed for statistical significance using Student's t test, using either Calc (The Document

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Foundation) or Excel (Microsoft Corporation) software; $p \le 0.05$ was regarded as significant. Synergistic behavior of drugs in combination was assessed by means of the comparison of the surviving fractions according to the method of Valeriote & Lin (27). Mice survival curves were analyzed using Prism (GraphPad Software Inc.).

Results

Dinaciclib induces cell death in STSs independently of CDK1 and CDK9 expression levels.

To have a base for inferring the response of STS to Dinaciclib, we analyzed the expression of CDK9, CDK1 and its partners Cyclin B1 and A2 in a series of cell lines representative of major groups of STS: synovial sarcoma SW982 and 1273-99, leiomyosarcoma SK-LMS-1 and SK-UT-1, myxoid liposarcoma 402-91 and 1765-92, dedifferentiated liposarcoma SW872 and myxofibrosarcoma HT-1080 (**Figure 1A**). Protein expression was variable among the cell lines, from the rather homogenous 43 kDa (low band) of CDK9 to the more variable CDK1. SW872 expressed comparatively undetectable levels of CDK1 and its partners. However, the levels of the 55 kDa (high band) of CDK9 were among the highest.

The same set of cell lines was used to test the ability of Dinaciclib to impair cellular viability (read as WST-1 assay, **Figure 1B**). SW982, SW872 and HT-1080 showed fewer decreases in viability at low concentrations, but at 27 nM differences among the cell lines were reduced. IC₅₀ calculations failed for SW872 and HT-1080 due to high disparity of the data (**Supplementary Figure 1A**), but actual values should be close to the 12.760 nM obtained for SW982.

However, when observed under the microscope, the response to Dinaciclib was more diverse (**Figure 1C**). Rather than any form of cell death, most cell lines (like SK-LMS-1) showed a stop in proliferation with almost the same number of cells as seeded and few detached. Only 402-91 and 1273-99 showed an extensive accumulation of detached, dead cells. To clarify this dual effect, we performed an analysis of apoptotic induction by cytofluorometric quantification of the levels of the vital stains $DiOC_6(3)$ and propidium iodide (DiOC-PI). Comparison among the 8 cell lines treated with 25 nM Dinaciclib for 72 h evidenced the two major phenotypes from the microscopical

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observations (**Figure 1D**). Only 1273-99 and 402-91 presented values of apoptotic cell death exceeding the 50 %. Dinaciclib concentration of 25 nM was chosen under experimental conditions as the minimal value above which cell death reached a plateau as soon as 24 h of treatment either in responsive or in tolerant cell lines (**Supplementary Figure 1B**).

In order to confirm that Dinaciclib induced mitochondrial apoptosis we studied the translocation of pro-apoptotic protein Bax to the mitochondria (**Figure 1E**) and the caspase-3 activation reporter PARP (**Supplementary Figure 1C**). Both features were confirmed in the sensitive cell lines 1273-99 and 402-91. Furthermore, the addition of 40 μ M pan-caspase inhibitor z-VAD-fmk to the system for 36 h was sufficient to abrogate the induction of cell death in both cell lines (**Supplementary Figures 1D and E**).

Dinaciclib potential as cell death inducer was conserved also in 3D tissue simulation. 402-91 cells embedded in a Matrigel[®] matrix grew as microspheres. By treating the microspheres with Dinaciclib for 24 h prior staining with vital dye H-33342, we clearly observed a reduction of the size of the spheres but also the accumulation of bright pyknotic nuclei generated by apoptosis (**Figure 1F**).

Taken together, our results show the potential of Dinaciclib as inducer of apoptotic cell death in STS at nanomolar concentrations.

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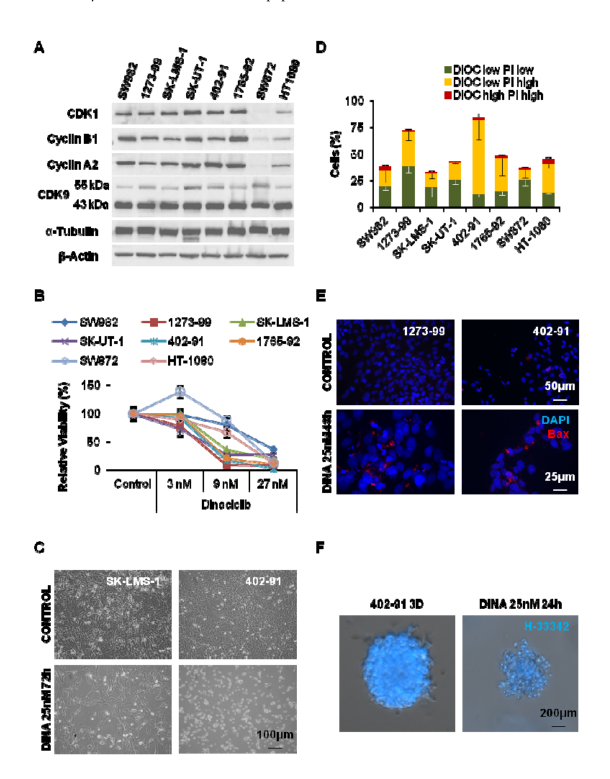


Figure 1: CDK inhibitor Dinaciclib is able to successfully inactivate sarcoma cell lines in a nanomolar range. (A) Representative western blot showing expression levels of Dinaciclib targets CDK1 and CDK9, and CDK1 partners Cyclin B1 and Cyclin A2 among different soft-tissue sarcoma derived cell lines. (B) Viability,

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measured by means of WST-1 reduction test, in a set of different STS cell lines incubated with increasing concentrations of Dinaciclib for 72 h; comparing with cells treated only with vehicle. (C) Microscopic imaging showing different cell behavior among cell lines when treated with Dinaciclib. (D) Comparison of apoptotic induction, cytofluorometric measurement of mitochondrial vital dye DiOC and death marker PI, in different STS derived cell lines treated with 25 nM Dinaciclib up to 72 h reveals different sensitivity among the cell lines. (E) Immunofluorescent detection of pro-apoptotic protein Bax showing its accumulation in *puncta* (mitochondria) in sensitive cells treated with 25 nM Dinaciclib. (F) 402-91 cells cultured as 3D spheroids by embedding into a Matrigel[®] matrix and treated by Dinaciclib show loss of size and accumulation of pyknotic nuclei as revealed by staining with Hoechst-33342 (H-33342).

Dinaciclib effects on cells last after wash-out.

To complete our understanding of cellular response to Dinaciclib, we performed longterm experiments analyzing cell response after drug wash-out. During the first days after Dinaciclib removal, cell lines from the tolerant group (like SW982) underwent cell death albeit the identification of a number of viable surviving colonies (**Figure 2A**). Cells from the sensitive lines (like 1273-99) presented rare, if any, surviving colonies. Additionally, we allowed those colonies to grow into sub-cultures that we dubbed Dinaciclib Resistant (DR). Those DR lines were newly challenged with 72 h incubation of Dinaciclib 25 nM. DRs from the tolerant group did not show any evident difference in behavior from the parental culture. But those derived from the sensitive cell lines appeared to have acquired a better fitness against the drug (**Figure 2B**). Nevertheless, when different experiments were summed up there was no statistical significance in this trend to a reduced cell death induction (**Figure 2C**).

Therefore, our data suggests that Dinaciclib exerts a long-term effect in cell viability with the additional value that surviving colonies do not show a strong selection towards drug resistance.

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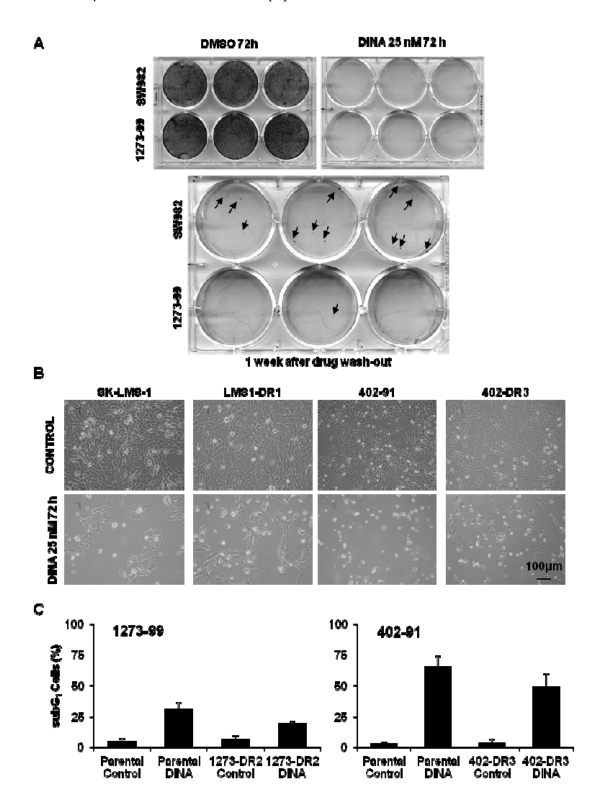


Figure 2: Dinaciclib did not drive strong selection of resistant clones after drug wash-out. (A) Crystal violet staining images showing resistant colonies (arrows) arising from highly sensitive 1273-99 cell line (low row) and more tolerant SW982

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(upper row) after Dinaciclib wash-out. (B) Microscopic imaging showing polyclonal Dinaciclib Resistant (DR) strains response after being newly subjected to 72 h of treatment. (C) Cytofluorometric measurement of death induction (subG₁ DNA content in PI profiles) for parental and derived DR strains from sensitive 1273-99 and 402-91 cell lines after a new cycle of 72 h incubation with Dinaciclib 25 nM (DINA).

Sensitive and tolerant cell lines differ in cell death protein expression patterns.

As the known targets of Dinaciclib yielded no clear correlation with drug response (**Figure 1**), we performed an extensive analysis of proteins involved in cell cycle control and mitochondrial apoptosis among examples of responsive (402-91 and 1273-99) and tolerant (SW982 and SK-LMS-1) cell lines (**Figure 3A**).

For most proteins little variation among the cell lines was observed. However, other proteins did shown remarkable differences in expression between the two groups. Tumor suppressor p53 was almost undetectable in Dinaciclib tolerant cells when compared with the responsive ones. In this regard, it is important to note that SK-LMS-1 cell line has been described as TP53 mutant (28). SK-LMS-1 expressed also relatively low levels of p21^{Waf1/Cip1} (p21) protein, a p53 target. Finally, CDK4 inhibitor p16^{INK4a} (p16) presented a pattern very similar to p53. If we compare this data with the study of cell cycle changes, we can see that only cell line SW982 presented a statistically significant arrest in G_2/M phase (Supplementary Figure 2A). In any case, most of the Dinaciclib tolerant cell lines did share the same trend. To analyze the precise nature of this G₂/M blockade we visualized treated and untreated cells immunolabelled for Ser10phosphorylated Histone H3 (p-H3) and p21. Our results confirmed that the arrest was not due to accumulation of mitotic events but rather to the arrest of cells in the G₂ phase (or in a tetraploid G_1). In concordance with the protein expression data p21 was not induced in SK-LMS-1 cells (Supplementary Figure 2B). To complete the analysis, we found that senescence-related p16 translocated towards the nuclei in tolerant cell lines treated with Dinaciclib, in a way similar to the sensitive ones (whose signal was clearly more intense) (Supplementary Figure 2C). Thus, tolerance to Dinaciclib seems to be linked to the generation of a partial G_2 arrest without a clear or sustained induction of senescence, as cells finally die in the following days (Figure 2A).

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Among the mitochondrial apoptotic control machinery, major differences were found in the small pro-apoptotic BH3-only proteins PUMA and BIM. Both were found comparatively over-expressed in Dinaciclib sensitive cell lines. Minor differences were also found in Noxa, BMF or Mcl-1 but with no correlation with the Dinaciclib response.

The major effect of Dinaciclib via inhibition of CDK9 is the interruption of mRNA transcription (17,29). On this basis, we analyzed how our main targets evolved upon Dinaciclib incubation (**Figure 3B**). As expected, most of the proteins showed a radical decrease in expression after Dinaciclib incubation. In concordance with the data from **Supplementary Figure 2C**, p21 was clearly induced in SW982 cell line. Cell cycle modulators p16 and p53 levels were kept in Dinaciclib sensitive cells. More importantly, anti-apoptotic regulator Bcl-x_L expression seemed invariable or slightly reduced by Dinaciclib. Only in 402-91 cells repression of Bcl-x_L was observed after 48 h of treatment. In contrast, Mcl-1 was early and completely suppressed in all cell lines. Finally, Dinaciclib responsive cell lines retained for longer time the, already higher, levels of pro-apoptotic BIM and PUMA.

Summing up, differences in key proteins among STS cell lines can explain particular features of Dinaciclib response.

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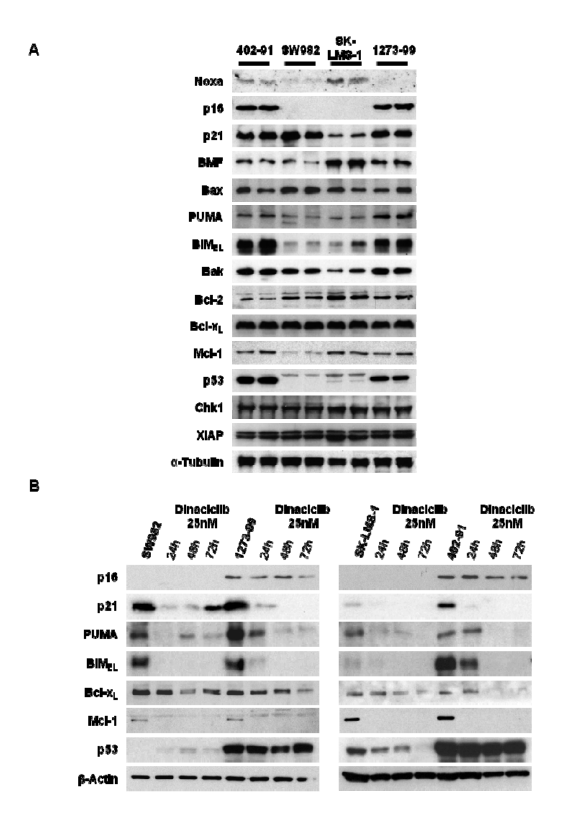


Figure 3: Dinaciclib highly sensitive cell lines present characteristic levels of key regulators of the cell cycle and apoptosis triggering. (A) Representative western blots showing protein expression patterns in control duplicates from myxoid

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liposarcoma 402-91, synovial sarcoma 1273-99 and SW982 and leiomyosarcoma SK-LMS-1. A representative sample of equal charges marker α -tubulin is also showed. (B) Representative western blots showing the decay rhythm of key components of cell cycle and apoptosis regulation at selected end-points of Dinaciclib incubation. A representative sample of equal charges marker β -actin is also showed.

Bcl-x_L silencing enhances sensitivity to Dinaciclib in SW982 and SK-LMS-1 cell lines.

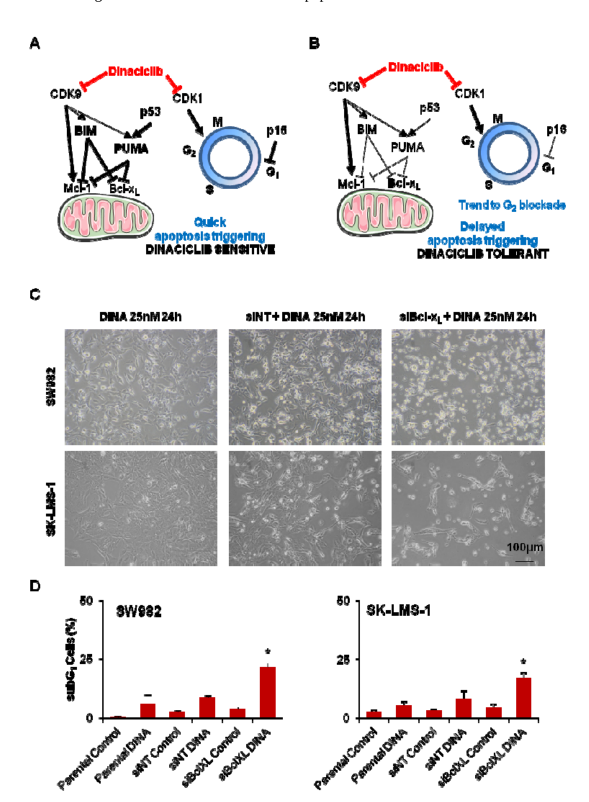
Based on our protein expression data, we developed a hypothetical framework to study the mechanistic of Dinaciclib action. Dinaciclib sensitive cells undergo quick apoptosis triggering due to their relative retention of higher levels of pro-apoptotic proteins (BIM, PUMA) after quick disappearing of anti-apoptotic Mcl-1. The high levels of p16 prevent surviving cells to get arrested in G_2 phase (**Figure 4A**). On the other hand, Dinaciclib tolerant cells had relatively lower levels of BIM and PUMA, and they are lost faster after CDK9 inhibition than in the responsive ones. Remaining Bcl-x_L prevents apoptosis to be triggered and cells proceed to G_2 when other mechanisms (like p21 induction in SW982) are activated (**Figure 4B**). In time, the induced damage will force the cells to die (**Figure 2A**).

To check the coherence of our hypothesis, we have reviewed the data presented by Booher and co-workers in 2014 showing correlation between *MCL1:BCL2L1* mRNA ratio and viability after 24 h 100 nM Dinaciclib treatment (18). By reducing the 250 samples to the ones appertaining to STS, we still found a statistically significant correlation, meaning that those cell lines more depending on Mcl-1 were also prone to die after Dinaciclib (**Supplementary Figure 3A**).

For experimental validation, we addressed the role of Mcl-1 in Dinaciclib-induced cell death by siRNA silencing (**Supplementary Figure 3B**). Although Mcl-1 was successfully depleted, no major increase in cell death was observed in SW982 cells (**Supplementary Figure 3C**). However, depleting Bcl- x_L in Dinaciclib tolerant cell lines before Dinaciclib addition (**Supplementary Figure 3D**) clearly resulted in an increase of the detachment of cells (**Figure 4C**). When cell death induction was quantified, the increase of cell death at 24 h of treatment was deemed statistically significant (**Figure 4D**).

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Therefore, our results supported our hypothesis that $Bcl-x_L$ is the more relevant counteracting mediator of Dinaciclib-induced apoptosis in STS.



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Figure 4: Bcl- x_L is the key regulator of apoptosis triggering after Dinaciclib treatment. (A & B) Schematics of the proposed mechanisms of action for the triggering of apoptosis in STS cell lines sensitive (A) and tolerant (B) to Dinaciclib. (C) Representative microscopic imaging of SW982 (above) and SK-LMS-1 (below) cell lines treated with 25 nM Dinaciclib for 24 h without prior challenge or after 24 h treatment with alternatively a non targeting (siNT) or a Bcl- x_L directed (siBcl- x_L) siRNA. (D) Cytofluorometric quantification of cell death induction (measured as subG₁ DNA content in PI profiles of fixed cell populations) after sequential treatment with siRNA and Dinaciclib. Data are presented as means \pm SD. Statistical significance was achieved by the Student's *t* test from at least three different experiments: * $p \leq 0.05$.

BH3-mimetics are able to synergistically enhance and accelerate cell death in STS.

To assess the possibility of a combination therapy to enhance Dinaciclib action we chose two different BH3-mimetics: one with a broad action spectrum, but excluding Mcl-1 (ABT-737) and one specific only for Bcl- x_L (A-1331852). BH3-mimetics activity over Dinaciclib tolerant cells as monotherapy was negligible (**Supplementary Figure 4A & C**) but its effect was notable when combined with 25 nM Dinaciclib (**Supplementary Figure 4B & D**). Cytofluorometric measurement showed that both combinations synergized (27) in terms of apoptosis induction after 24 h of treatment at very low concentrations (**Figure 5A-B**). Moreover, A-1331852 was more efficient than ABT-737. These results confirmed our operational hypothesis regarding Dinaciclib mechanism of action.

The next logical step was trying the drug combination *in vivo*. Our first approaches were aimed to assess treatment toxicity and cell growth using Hsd:Athymic Nude*Foxn1*^{*nu*} mice. We tried different combination regimes of Dinaciclib and ABT-737 based on literature (18,30) (**Supplementary Figure 5A**), but we always observed liver toxicity linked to massive hepatocyte death (**Figure 5C**). Moreover, human SK-LMS-1 showed a poor efficiency of nesting and growing as sub-cutaneous tumors in those mice (**Supplementary Figure 5B**). In consequence, we moved to the more immunodepressed CB17.Cg-*Prkdc*^{*scid*}*Lyst*^{*bg-J*}/Crl mice trying a lower dosage of Dinaciclib (as it was able to kill the mice even as monotherapy) and A-1331852 instead of ABT-737 (**Supplementary Figure 5C**). SK-LMS-1 cells grew better in these mice

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(Supplementary Figure 5D), but an important percentage of tumors appeared as intraperitoneal masses despite being injected sub-cutaneously (Figure 5D). Intra-peritoneal tumors grew by surrounding, engulfing and finally invading the kidney (Supplementary Figure 5E). Those mice were excluded from the final experiments. Final *in vivo* testing of anti-tumoral combined activity of both drugs was unsuccessful in terms of mice survival (Figure 5E). The failure was allegedly due to a lack of overall effect of the drugs, as can be inferred for the unchanged mice weight (Supplementary Figure 5F). Summing up, we could not reach a therapeutic regime in which the drugs showed anti-tumor activity without induction of deadly toxicity.

Notwithstanding this set back, we sought to understand whether the key role of $Bcl-x_L$ in Dinaciclib-induced cell death was a particular feature or part of a more general behavior. Other groups have already shown a positive cooperation of BH3-mimetics with CDK inhibitors and anthracyclines (26,31). So, we tested another commonly used anti-tumor drug in STS therapy: nucleoside analogue Gemcitabine. Our results showed that A-1331852 synergized with Gemcitabine in terms of cell death induction after 48 h of combined treatment (**Figure 5F**).

Taking together, our results unveil the specificities of cellular response to CDK inhibitor Dinaciclib and positively identify $Bcl-x_L$ as the key player of apoptosis induction in these systems. More pharmaceutical research will be required to achieve a safe way for clinical practice, probably through the development of new delivery methods or new molecules with reduced side-effects. Still, this question falls beyond the scope of our study.

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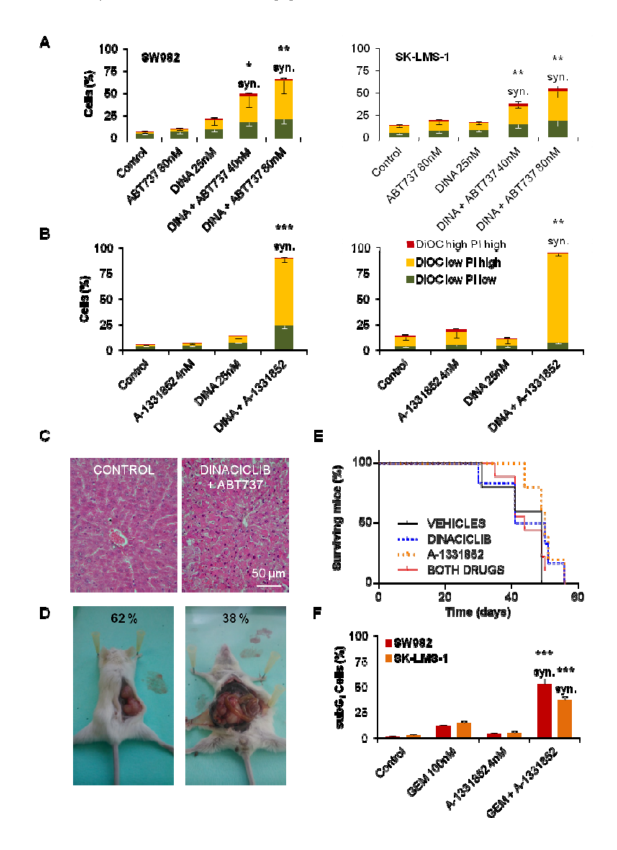


Figure 5: BH3-mimetics targeting Bcl-xL synergize with Dinaciclib to achieve quick apoptosis. (A) Cytofluorometric quantification of cell death via DiOC-PI

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staining of SW982 (left) and SK-LMS-1 (right) cell lines treated for 24 h with Dinaciclib 25 nM, ABT-737 80 nM or a combination of both drugs. (B) Cytofluorometric quantification of cell death via DiOC-PI staining of SW982 (left) and SK-LMS-1 (right) cell lines treated for 24 h with Dinaciclib 25 nM, A-1331852 4 nM or a combination of both drugs. (C) Microscopical imaging of H&E stained liver sections from Hsd:Athymic NudeFoxn1^{nu} mice control (left) and treated (right) after 3 i.p. injections with 40 mg/Kg Dinaciclib and 7 i.p. injections of ABT-737 who suffered sudden death at day 8. (D) Photographs showing typical subcutaneous tumor (left) and atypical intra-peritoneal tumor (right) caused by SK-LMS-1 cells in CB17.Cg-Prkdc^{scid}Lyst^{bg-J}/Crl mice. Percentages of occurrence are presented on the images. (E) Kaplan-Meier graph showing the survival of CB17.Cg-Prkdc^{scid}Lvst^{bg-J}/Crl mice inoculated (s.c.) with 5×10⁶ SK-LMS-1 cells and afterwards treated with i.p. 20 mg/Kg Dinaciclib, p.o. 25 mg/Kg A-1331852 or both drugs. Control mice were treated with both drugs vehicles. Time 0 represents the beginning of the treatment. (F) Cytofluorometric quantification of cell death (as subG₁ in PI profiles in fixed cells) of SW982 and SK-LMS-1 cells treated with 100 nM Gemcitabine (GEM), 4 nM A-1331852 or a combination of both drugs for 48 h. Data are presented as means \pm SD. Statistical significance was achieved by the Student's *t* test from at least three different experiments: $p \le 0.05$; $p \le 0.001$; *** $p \leq 0.0001$. syn. denotes synergistic behavior as determined according Valeriote & Lin.

Discussion

Dinaciclib is one of the most promising compounds in the group of CDK inhibitors because is able to act by meddling in the progression of late steps of the cell cycle, the entry in mitosis, transcriptional regulation and even the unfolded protein response (15,32,33). It has been studied in clinical trials on cancer entities such as acute leukemia and other pediatric malignancies (11,14,20). Dinaciclib was well tolerated, but its therapeutic effect as single agent did not achieve any total remission. However, in several neoplasias it is considered a good candidate for combination therapies (7,33,34), including BH3-mimetics (26).

Here we present new data on Dinaciclib pre-clinical results in STS cell lines. Our results show that Dinaciclib is able, as single agent, to impede cell proliferation in a nanomolar

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range (**Figure 1B**) and to induce cell death at least in myxoid liposarcoma 402-91 and synovial sarcoma 1273-99 cells (**Figure 1D**). Our model of 3D culture also proves that Dinaciclib is able to infiltrate and kill cells in solid tumors (**Figure 1F**). These results align with previous pre-clinical reports that have also proved the utility of Dinaciclib in different tumor entities (16,18,33). Moreover, our long-term studies after Dinaciclib removal showed that the deregulation induced by the drug was enough to promote a delayed death (five days after drug withdrawal) and that the remnant colonies did not develop an enhanced resistance to re-challenge (**Figure 2C**). The lack of reinforced tolerance to Dinaciclib in DR lines opens the gate to the establishment of repeated administrations or combination procedures (35,36).

To do so, a proper cell death mechanism characterization is paramount (37). In this sense, our results proved that the main mechanism involved in Dinaciclib killing is the mitochondrial apoptotic pathway (Figure 1D-E and Supplementary Figure 1B-E). Other in vitro studies published with Dinaciclib also pointed to apoptotic mechanisms but they provided a less detailed characterization of the cell death (16,17,19). Dinaciclib action involves two clearly different processes: cell cycle inhibition and alteration in protein production and homeostasis (9,32). The study of those cell lines more tolerant to Dinaciclib is helpful as their delayed death allow us to observe other variables. Cell cycle profiling showed a trend to an arrest in G₂-to-M transition that was only significant in SW982 (Supplementary Figure 2A). SW982 cells are also the only cell line able to induce an increase in p21 protein (Figure 3B and Supplementary Figure **2B**). Our data agreed to previous reports in other cancer entities, in which the cell cycle arrest was revealed when apoptosis was inhibited by Bax/Bak DKO (18). But this mechanism may not be universal as most of our Dinaciclib tolerant cell lines did not reach statistically significant mitotic arrest. However, the sensitive cell lines indeed did increase their G₂/M and postM (tetraploidy) phases when apoptosis was abrogated by z-VAD-fmk (data not shown). So, it appears that the effect on the cell cycle deregulation in our models may be an important contributor in the apoptotic triggering. Further research is needed to understand if other proteins (like p21) are required for G_2 arrest after CDK1 inhibition.

The role of cell cycle regulator p16 is intriguing. It is one of the multiple transcripts of the *CDKN2A* locus, all of them with key functions in the homeostasis of the cell cycle, tumor suppression, senescence and aging (38–42). The expression of p16 is not directly

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regulated by p53 but rather by Jun-B and Ras via Ets-1/2 (38). However, there are references cross-linking Ets-1/2 transcription factors with Bcl-2 family expression (43–46) and even p53 activity (47). In our settings, the sensitive cell lines 1273-99 and 402-91 exhibit higher levels of p16 than the tolerant ones (**Figure 3A and Supplementary Figure 2C**). Hence, there is supporting data to consider that expression levels of p16 could be linked to an apoptotic-permissive unbalance among the Bcl-2 family that enhances Dinaciclib sensitivity. As has been proved in other cancer entities, we consider interesting to explore the use of p16 as a predictive marker for cell cycle targeted therapies (39,42).

Dinaciclib causes a general depletion of protein content in the cell via RNA Polymerase II dephosphorylation with different impact depending on protein distinctive half-life (7,9,17,18). From our list of proteins of interest (Figure 3B), we focused in Bcl-x_L as it maintained its levels essentially constant after Dinaciclib incubation. Delayed depletion of $Bcl-x_L$ was seen only in the most sensitive cell line (402-91). In contrast, PUMA and BIM levels decreased later in Dinaciclib responsive cell lines. Mitochondrial apoptosis is basically regulated by the inhibition status of large anti-apoptotic members of Bcl-2 family of proteins. This inhibition status relies on the balance with the pro-apoptotic members of the group, particularly the small BH3-only proteins (48). Published literature has focused in anti-apoptotic regulator Mcl-1 levels as the main mediator of Dinaciclib cytotoxicity because its expression drops drastically after treatment (17–19). Recently, a more comprehensive study from Inoue-Yamauchi and co-workers lead to a more complex panorama in which every member of the Bcl-2 family has a role in the apoptotic triggering (26). For STS, the limited cohort from Booher and co-workers provides correlation between Dinaciclib sensitivity and Mcl-1/Bcl-x_I ratio (Supplementary Figure 3A). However, our data showed that Mcl-1 levels are already low in SW982 cells (Figure 3A) and quickly become almost undetectable upon Dinaciclib treatment in all cell lines (Figure 3B). So, we envisaged a mechanistic hypothesis that put remnant Bcl-x_L interaction with BIM and/or PUMA in the decision point of apoptosis triggering in STS cell lines (Figure 4A-B). When Dinaciclib is added, its effect over CDK9 causes a quick depletion in anti-apoptotic protein Mcl-1 but the effect over BIM or PUMA is weaker and negligible on p53. PUMA and BIM are thus capable to initiate mitochondrial apoptosis by inhibition of Bcl-x_L. Yet SW982 and SK-LMS-1 have low levels of BIM and PUMA. In the presence of Dinaciclib, the

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remaining levels of Bcl- x_L are enough to sustain cell viability for longer time. Then, the effects over the cell cycle (via CDK1) become detectable as a trend (significant in SW982) to a G₂ blockade. Silencing experiments with siRNA confirmed our suspicion, as Bcl- x_L induced a clear increase in detached cells whereas Mcl-1 inhibition exerted no major difference (**Figure 4C-D and Supplementary Figure 3B-C**).

 $Bcl-x_L$ involvement as key mediator of apoptosis triggering has been previously described in several contexts including cell cycle alterations and other sarcoma entities (49–51). Among the different Bcl-2 family interactions, $Bcl-x_L$ has strong affinity for pro-apoptotic BH3-only proteins PUMA and BIM (49,50,52,53). BH3-mimetics are a new class of drugs precisely designed to benefit of this interaction to promote cell death (24). Since the recent FDA approval of Venetoclax, these molecules are in the spotlight of cancer research, albeit some groups are calling to improve the accurateness of data analysis (54). A more detailed research in their precise mechanism of action is needed in order to design meaningful combinations for treatment. Nevertheless, we decided to go ahead and test our working hypothesis by designing drug combination testing in vitro. Two BH3-mimetics were chosen: ABT-737, a rather generic compound that, inhibits Bcl-2, Bcl-w and Bcl- x_L proteins (55) and A-1331852, a newer and Bcl- x_L specific molecule (56). Neither of them induced a relevant cell death accumulation as monotherapy, but when combined with Dinaciclib they showed a fast synergistic induction of cell death in both SW982 and SK-LMS-1 cells (Figure 5A-B and Supplementary Figure 4). The effects were clearly better with A-1331852 than with ABT-737, providing more support to our mechanistic hypothesis. Regretfully, we encountered too many problems at escalating our settings to in vivo experimentation (Figure 5C-E and Supplementary Figure 5). The main issue was the development of deadly liver toxicity, that was already glimpsed in literature (57). Our data supports the need of more pharmaceutical research in order to generate a safer treatment protocol.

In the past few years, other groups have recognized the potential of BH3-mimetics in combination approaches to kill cancer cells. As previously commented, Inoue-Yamauchi and co-workers, using screening technologies in a small-cell lung cancer model, found that anti-tumor properties of CDK9 inhibitors and anthracyclines were boosted when combined with those molecules (26). Research in STS has also pointed to the suitability of such combinations. Possible combinations for BH3-mimetics include anthracyclines, histone deacetylase inhibitors, tubulin poisons and etoposide (31,58–

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60). Here, we add to the list Dinaciclib and Gemcitabine (Figure 5F) and we point that the major focus should be put in $Bcl-x_L$ inhibitors. Both, our data and the previous literature support the conclusion that $Bcl-x_L$ targeted molecules behave better than generic BH3-mimetics. The existence of a single molecular route to explain this multiple activity seems unlikely. Bcl- x_L has been identified as a key player of cell death triggering after mitotic arrest (50). This can explain the synergy reported with tubulin poisons or other compounds that induced strong mitotic arrest. But, it is not the case with Dinaciclib, as we obtained no relevant mitotic arrest linked to the cell death triggering. The G_2 blockage reported for SW982 occurs at 72 h and combined cell death is verified at 24 h. Thus, in our setting it is more relevant the quick loss of protein players once Dinaciclib has disturbed the process of protein synthesis. The higher life time of Bcl-x_I positions it as the cornerstone of apoptotic induction. For Gemcitabine we cannot discard that the mechanism proceeds more similarly to the observed for mitotic disruptors. In this case, combined action takes more time to be verified (48 h vs. 24 h with Dinaciclib) showing that BH3-mimetics did not exert a pro-apoptotic activity by themselves. Indeed, their function is simply to push towards cell death *after* another injury has been detected in the cell.

Taking together, we have extensively showed that Dinaciclib (or an improved CDK9 inhibitor) is a suitable candidate for clinical trials in STS. It is capable to effectively kill STS cell lines in a relevant fashion, without inducing a strong selection towards drug resistance. Moreover, when combined with BH3-mimetics its action gets boosted in a remarkable way. Pharmacological research is, however, required in order to address the elevated toxicity linked to this drug.

Acknowledgments

The mitochondria draw in Figure 4, was taken from Servier Medical Art (https://smart.servier.com/) under the terms of a Creative Commons Attribution 3.0 Unported License. The mice draw in Supplementary Figure 5 was taken and adapted from Wikimedia Commons image archive (https://commons.wikimedia.org/wiki/File:Mouse_line_drawing.jpg), licensed under the Creative Commons Attribution 2.5 Generic license. We thank David Herrero-Martín for helpful discussion.

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Author's Contributions

Conception and design: S. R-V., O. M. T. and X. GdM. **Development of methodology:** S. R-V., M. F-G. and O. M. T. **Acquisition of data:** S. R-V., M. F-G., R. L-A., A. C-P., N. M-M. and S. G-M. **Analysis and interpretation of data:** S. R-V., M. F-G. and O. M. T. **Writing and/or revision of manuscript:** S. R-V., O. M. T. and X. GdM. **Funding acquisition:** S. R-V., O. M. T. and X. GdM.

Bibliography

- Oda Y, Yamamoto H, Kohashi K, Yamada Y, Iura K, Ishii T, et al. Soft tissue sarcomas: From a morphological to a molecular biological approach. Pathol Int. 2017 Sep 1;67(9):435–46.
- Clark MA, Fisher C, Judson I, Thomas JM. Soft-tissue sarcomas in adults. N Engl J Med. 2005 Aug 18;353(7):701–11.
- 3. Jain S, Xu R, Prieto VG, Lee P. Molecular classification of soft tissue sarcomas and its clinical applications. Int J Clin Exp Pathol. 2010 Jan;3(4):416–28.
- Abeshouse A, Adebamowo C, Adebamowo SN, Akbani R, Akeredolu T, Ally A, et al. Comprehensive and Integrated Genomic Characterization of Adult Soft Tissue Sarcomas. Cell. 2017 Nov 2;171(4):950–965.e28.
- Ducimetière F, Lurkin A, Ranchère-Vince D, Decouvelaere A-V, Péoc'h M, Istier L, et al. Incidence of sarcoma histotypes and molecular subtypes in a prospective epidemiological study with central pathology review and molecular testing. Najbauer J, editor. PLoS One. 2011 Jan;6(8):e20294.
- 6. Demicco EG, Maki RG, Lev DC, Lazar AJ. New therapeutic targets in soft tissue sarcoma. Adv Anat Pathol. 2012 May;19(3):170–80.
- Cicenas J, Kalyan K, Sorokinas A, Jatulyte A, Valiunas D, Kaupinis A, et al. Highlights of the Latest Advances in Research on CDK Inhibitors. Cancers (Basel). 2014 Jan 27;6(4):2224–42.
- Malumbres M, Barbacid M. Cell cycle, CDKs and cancer: a changing paradigm. Nat Rev Cancer. 2009 Mar 1;9(3):153–66.
- 9. Sánchez-Martínez C, Gelbert LM, Lallena MJ, de Dios A. Cyclin dependent kinase (CDK) inhibitors as anticancer drugs. Bioorg Med Chem Lett. 2015 Sep

Rello-Varona et al. 2018

1;25(17):3420-35.

- Castedo M, Perfettini J, Roumier T, Kroemer G. Cyclin-dependent kinase-1: linking apoptosis to cell cycle and mitotic catastrophe. Cell Death Differ. 2002 Dec;9(12):1287–93.
- Gorlick R, Kolb EA, Houghton PJ, Morton CL, Neale G, Keir ST, et al. Initial testing (stage 1) of the cyclin dependent kinase inhibitor SCH 727965 (dinaciclib) by the pediatric preclinical testing program. Pediatr Blood Cancer. 2012 Dec 15;59(7):1266–74.
- Parry D, Guzi T, Shanahan F, Davis N, Prabhavalkar D, Wiswell D, et al. Dinaciclib (SCH 727965), a novel and potent cyclin-dependent kinase inhibitor. Mol Cancer Ther. 2010 Aug 1;9(8):2344–53.
- Shah K, Rossie S. Tale of the Good and the Bad Cdk5: Remodeling of the Actin Cytoskeleton in the Brain. Mol Neurobiol. 2018 Apr 13;55(4):3426–38.
- Gojo I, Sadowska M, Walker A, Feldman EJ, Iyer SP, Baer MR, et al. Clinical and laboratory studies of the novel cyclin-dependent kinase inhibitor dinaciclib (SCH 727965) in acute leukemias. Cancer Chemother Pharmacol. 2013 Oct;72(4):897–908.
- Paruch K, Dwyer MP, Alvarez C, Brown C, Chan T-Y, Doll RJ, et al. Discovery of Dinaciclib (SCH 727965): A Potent and Selective Inhibitor of Cyclin-Dependent Kinases. ACS Med Chem Lett. 2010 Aug 12;1(5):204–8.
- Desai BM, Villanueva J, Nguyen T-TK, Lioni M, Xiao M, Kong J, et al. The anti-melanoma activity of dinaciclib, a cyclin-dependent kinase inhibitor, is dependent on p53 signaling. PLoS One. 2013 Jan 18;8(3):e59588.
- Gregory GP, Hogg SJ, Kats LM, Vidacs E, Baker AJ, Gilan O, et al. CDK9 inhibition by dinaciclib potently suppresses Mcl-1 to induce durable apoptotic responses in aggressive MYC-driven B-cell lymphoma in vivo. Leukemia. 2015 Jun;29(6):1437–41.
- Booher RN, Hatch H, Dolinski BM, Nguyen T, Harmonay L, Al-Assaad A-S, et al. MCL1 and BCL-xL levels in solid tumors are predictive of dinaciclib-induced apoptosis. PLoS One. 2014 Jan 7;9(10):e108371.
- 19. Johnson AJ, Yeh Y-Y, Smith LL, Wagner AJ, Hessler J, Gupta S, et al. The

Bcl-x_L interferes in *Dinaciclib-induced* apoptosis in sarcomas.

novel cyclin-dependent kinase inhibitor dinaciclib (SCH727965) promotes apoptosis and abrogates microenvironmental cytokine protection in chronic lymphocytic leukemia cells. Leukemia. 2012 Dec 30;26(12):2554–7.

- Nemunaitis JJ, Small KA, Kirschmeier P, Zhang D, Zhu Y, Jou Y, et al. A firstin-human, phase 1, dose-escalation study of dinaciclib, a novel cyclin-dependent kinase inhibitor, administered weekly in subjects with advanced malignancies. J Transl Med. 2013 Jan;11(1):259.
- McGranahan N, Swanton C. Clonal Heterogeneity and Tumor Evolution: Past, Present, and the Future. Cell. 2017 Feb;168(4):613–28.
- Palmer AC, Sorger PK. Combination Cancer Therapy Can Confer Benefit via Patient-to-Patient Variability without Drug Additivity or Synergy. Cell. 2017 Dec 14;171(7):1678–1691.e13.
- Ni Chonghaile T, Letai A. Mimicking the BH3 domain to kill cancer cells. Oncogene. 2008 Dec;27 Suppl 1:S149-57.
- 24. Montero J, Letai A. Why do BCL-2 inhibitors work and where should we use them in the clinic? Cell Death Differ. 2018 Jan 27;25(1):56–64.
- Adams JM, Cory S. The BCL-2 arbiters of apoptosis and their growing role as cancer targets. Cell Death Differ. 2018 Jan 3;25(1):27–36.
- Inoue-Yamauchi A, Jeng PS, Kim K, Chen H-C, Han S, Ganesan YT, et al. Targeting the differential addiction to anti-apoptotic BCL-2 family for cancer therapy. Nat Commun. 2017 Jul 17;8(May):16078.
- Valeriote F, Lin H s. Synergistic interaction of anticancer agents: a cellular perspective. Cancer Chemother Rep. 1975;59(5):895–900.
- Stratton MR, Moss S, Warren W, Patterson H, Clark J, Fisher C, et al. Mutation of the p53 gene in human soft tissue sarcomas: association with abnormalities of the RB1 gene. Oncogene. 1990 Sep;5(9):1297–301.
- Mariaule G, Belmont P. Cyclin-dependent kinase inhibitors as marketed anticancer drugs: where are we now? A short survey. Molecules. 2014 Jan 11;19(9):14366–82.
- 30. Vandenberg CJ, Cory S. ABT-199, a new Bcl-2-specific BH3 mimetic, has in vivo efficacy against aggressive Myc-driven mouse lymphomas without

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Rello-Varona et al. 2018

provoking thrombocytopenia. Blood. 2013 Mar 21;121(12):2285-8.

- de Graaff MA, de Rooij MAJ, van den Akker BEWM, Gelderblom H, Chibon F, Coindre J-M, et al. Inhibition of Bcl-2 family members sensitises soft tissue leiomyosarcomas to chemotherapy. Br J Cancer. 2016 May 24;114(11):1219–26.
- Blachly JS, Byrd JC. Emerging drug profile: cyclin-dependent kinase inhibitors. Leuk Lymphoma. 2013 Oct;54(10):2133–43.
- Vymětalová L, Kryštof V. Potential Clinical Uses of CDK Inhibitors: Lessons from Synthetic Lethality Screens. Med Res Rev. 2015 Nov 25;35(6):1156–74.
- 34. Fabre C, Gobbi M, Ezzili C, Zoubir M, Sablin M-P, Small K, et al. Clinical study of the novel cyclin-dependent kinase inhibitor dinaciclib in combination with rituximab in relapsed/refractory chronic lymphocytic leukemia patients. Cancer Chemother Pharmacol. 2014 Nov;74(5):1057–64.
- Dean M, Fojo T, Bates S. Tumour stem cells and drug resistance. Nat Rev Cancer. 2005 Apr;5(4):275–84.
- Minchinton AI, Tannock IF. Drug penetration in solid tumours. Nat Rev Cancer. 2006 Aug;6(8):583–92.
- Rello-Varona S, Herrero-Martin D, Lopez-Alemany R, Munoz-Pinedo C, Tirado OM, Herrero-Martín D, et al. "(Not) all (dead) things share the same breath": identification of cell death mechanisms in anticancer therapy. Cancer Res. 2015 Mar 15;75(6):913–7.
- Kim WY, Sharpless NE. The regulation of INK4/ARF in cancer and aging. Cell. 2006 Oct 20;127(2):265–75.
- LaPak KM, Burd CE. The molecular balancing act of p16(INK4a) in cancer and aging. Mol Cancer Res. 2014 Mar 1;12(2):167–83.
- Collado M, Blasco MA, Serrano M. Cellular senescence in cancer and aging. Cell. 2007 Jul 27;130(2):223–33.
- 41. Puig-Butille JA, Escámez MJ, Garcia-Garcia F, Tell-Marti G, Fabra Á, Martínez-Santamaría L, et al. Capturing the biological impact of CDKN2A and MC1R genes as an early predisposing event in melanoma and non melanoma skin cancer. Oncotarget. 2014 Mar 30;5(6):1439–51.
- 42. Sheppard KE, McArthur GA. The cell-cycle regulator CDK4: an emerging

 $Bcl-x_L$ interferes in Dinaciclib-induced apoptosis in sarcomas.

therapeutic target in melanoma. Clin Cancer Res. 2013 Oct 1;19(19):5320-8.

- Dong L, Jiang CC, Thorne RF, Croft A, Yang F, Liu H, et al. Ets-1 mediates upregulation of Mcl-1 downstream of XBP-1 in human melanoma cells upon ER stress. Oncogene. 2011 Aug 25;30(34):3716–26.
- 44. Yu Z, Shah DM. Curcumin down-regulates Ets-1 and Bcl-2 expression in human endometrial carcinoma HEC-1-A cells. Gynecol Oncol. 2007 Sep;106(3):541–8.
- 45. Sevilla L, Zaldumbide A, Carlotti F, Dayem MA, Pognonec P, Boulukos KE. Bcl-XL expression correlates with primary macrophage differentiation, activation of functional competence, and survival and results from synergistic transcriptional activation by Ets2 and PU.1. J Biol Chem. 2001 May 25;276(21):17800–7.
- 46. Nakazawa Y, Suzuki M, Manabe N, Yamada T, Kihara-Negishi F, Sakurai T, et al. Cooperative interaction between ETS1 and GFI1 transcription factors in the repression of Bax gene expression. Oncogene. 2007 May 24;26(24):3541–50.
- Xu D, Wilson TJ, Chan D, De Luca E, Zhou J, Hertzog PJ, et al. Ets1 is required for p53 transcriptional activity in UV-induced apoptosis in embryonic stem cells. EMBO J. 2002 Aug 1;21(15):4081–93.
- 48. Youle RJ, Strasser A. The BCL-2 protein family: opposing activities that mediate cell death. Nat Rev Mol Cell Biol. 2008 Jan;9(1):47–59.
- Michels J, Kepp O, Senovilla L, Lissa D, Castedo M, Kroemer G, et al. Functions of BCL-X L at the Interface between Cell Death and Metabolism. Int J Cell Biol. 2013;2013:705294.
- Bah N, Maillet L, Ryan J, Dubreil S, Gautier F, Letai A, et al. Bcl-xL controls a switch between cell death modes during mitotic arrest. Cell Death Dis. 2014 Jan;5:e1291.
- Baranski Z, de Jong Y, Ilkova T, Peterse EFP, Cleton-Jansen A-M, van de Water B, et al. Pharmacological inhibition of Bcl-xL sensitizes osteosarcoma to doxorubicin. Oncotarget. 2015;1–12.
- Ming L, Wang P, Bank A, Yu J, Zhang L. PUMA Dissociates Bax and Bcl-X(L) to induce apoptosis in colon cancer cells. J Biol Chem. 2006 Jun 9;281(23):16034–42.

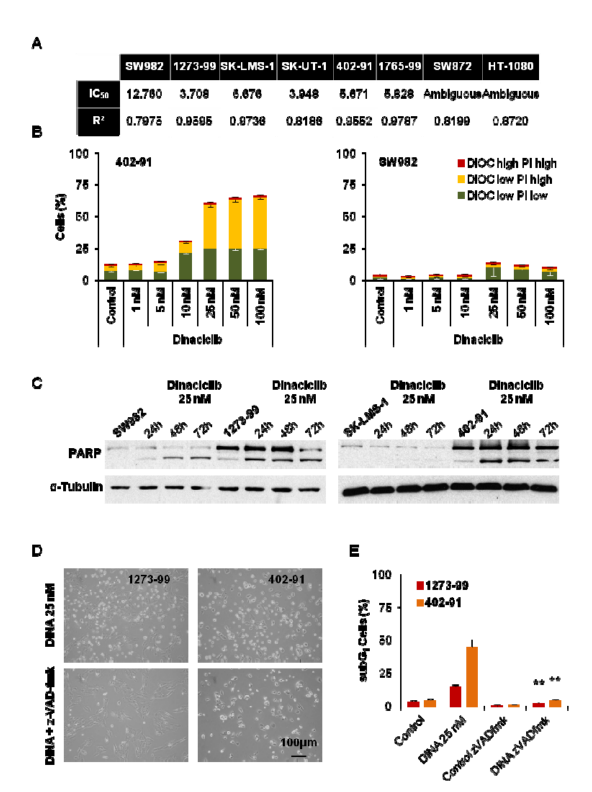
Rello-Varona et al. 2018

- 53. Alfredsson J, Puthalakath H, Martin H, Strasser A, Nilsson G. Proapoptotic Bcl-2 family member Bim is involved in the control of mast cell survival and is induced together with Bcl-XL upon IgE-receptor activation. Cell Death Differ. 2005 Feb;12(2):136–44.
- S Soderquist R, Eastman A. BCL2 Inhibitors as Anticancer Drugs: A Plethora of Misleading BH3 Mimetics. Mol Cancer Ther. 2016 Sep;15(9):2011–7.
- 55. Ren D, Tu H-C, Kim H, Wang GX, Bean GR, Takeuchi O, et al. BID, BIM, and PUMA are essential for activation of the BAX- and BAK-dependent cell death program. Science. 2010 Dec 3;330(6009):1390–3.
- 56. Leverson JD, Phillips DC, Mitten MJ, Boghaert ER, Diaz D, Tahir SK, et al. Exploiting selective BCL-2 family inhibitors to dissect cell survival dependencies and define improved strategies for cancer therapy. Sci Transl Med. 2015 Mar 18;7(279):279ra40.
- 57. Hikita H, Takehara T, Shimizu S, Kodama T, Shigekawa M, Iwase K, et al. The Bcl-xL inhibitor, ABT-737, efficiently induces apoptosis and suppresses growth of hepatoma cells in combination with sorafenib. Hepatology. 2010 Oct;52(4):1310–21.
- Faqar-Uz-Zaman SF, Heinicke U, Meister M, Vogler M, Fulda S. BCL-x L selective BH3 mimetic sensitizes rhabdomyosarcoma cells to chemotherapeutics by activation of the mitochondrial pathway of apoptosis. Cancer Lett. 2017 Sep 23;
- Heinicke U, Haydn T, Kehr S, Vogler M, Fulda S. BCL-2 selective inhibitor ABT-199 primes rhabdomyosarcoma cells to histone deacetylase inhibitorinduced apoptosis. Oncogene. 2018 Jun 1;1.
- Jones KB, Su L, Jin H, Lenz C, Randall RL, Underhill TM, et al. SS18-SSX2 and the mitochondrial apoptosis pathway in mouse and human synovial sarcomas. Oncogene. 2013 May 2;32(18):2365–71, 2375.e1-5.

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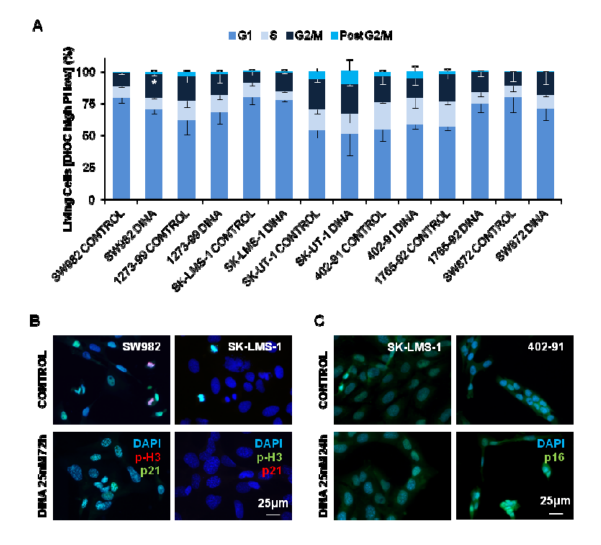
Supplementary Figure 1 (related to Figure 1): Dinaciclib successfully induce apoptosis in sensitive cell lines. (A) IC₅₀ values obtained from WST-1 reduction test represented in Figure 1B. (B) Cytofluorometric measurement of mitochondrial vital dye DiOC and death marker PI showing apoptotic induction in 402-91 cell line (left), but not in SW982 cells (right), when treated with increasing concentrations of Dinaciclib for 24 h. (C) Representative western blots showing PARP protein cleavage after treatment with 25 nM Dinaciclib at different end times. (D) Representative microscopic imaging showing that Dinaciclib-induced apoptosis can be abrogated with co-incubation with 40 μ M z-VAD-fmk for 36 h. (E) Cytofluorometric quantification of cell death (as subG₁ in PI profiles in fixed cells) of 1273-99 and 402-91 cells treated with 25 nM Dinaciclib and 40 μ M z-VAD-fmk for 36 h. Data are presented as means \pm SD. Statistical significance was achieved by the Student's *t* test from at least three different experiments: ** $p \le 0.001$.

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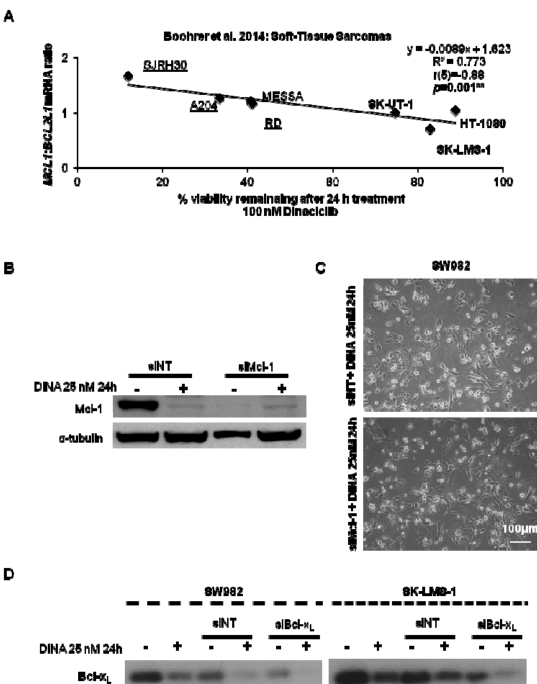
Supplementary Figure 2 (related to Figure 3): Differences in protein expression levels explain particular responses to Dinaciclib treatment. (A) Cytofluorometric measurement of cell cycle phases by means of H-333258 profiling of DiOC high PI low (living) cells in control conditions or after 72 h treatment with 25 nM Dinaciclib (DINA). SubG₁ cells were excluded. (B) Microscopic imaging showing mitotic marker phosphorylated histone H3 (p-H3) and checkpoint protein p21^{Waf1/Cip1} (p21) in control conditions or after 72 h treatment with 25 nM DINA. (C) Microscopic imaging showing senescence marker p16^{INK4a} (p16) in control conditions or after 24 h treatment with 25 nM DINA. Data are presented as means \pm SD. Statistical significance was achieved by the Student's *t* test from at least three different experiments: * $p \le 0.05$.



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Supplementary Figure 3 (related to Figure 4): Mcl-1 is not involved in Dinaciclib tolerance in STS cell lines. (A) Reanalysis of Boohrer *et al.* data on Dinaciclib tolerance and Mcl-1 and Bcl- x_L expression (mRNA) constrained to the "Soft-Tissue Sarcoma" subset. Cell lines names are noted. Rhabdomyosarcoma cell lines are underlined. Cell lines also included in our study are highlighted in bold. (B) Representative western blot showing Mcl-1 expression in SW982 cell line in the combined presence of siRNA constructs and Dinaciclib. (C) Representative microscopic imaging showing SW982 cell response to sequential silencing of Mcl-1 prior treatment with Dinaciclib 25 nM. (D) Representative western blot showing Bcl- x_L expression in SW982 and SK-LMS-1 in sequential silencing of the target prior of Dinaciclib incubation. Data are presented as means \pm SD. Statistical significance was achieved by the Student's *t* test from at least three different experiments: ** $p \le 0.001$.

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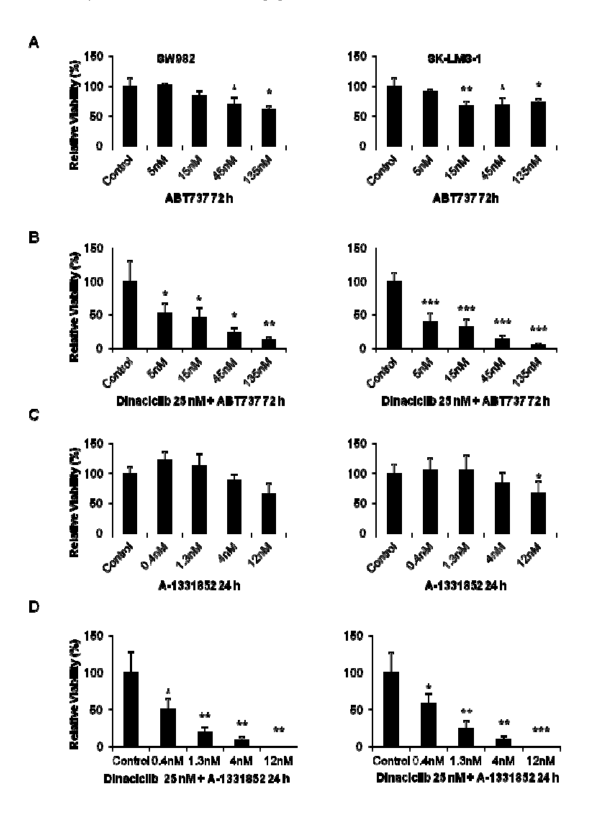




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Supplementary Figure 4 (related to Figure 5): BH3-mimetics are unable to induce relevant harm to STS cell lines as monotherapy. (A to D) Viability measured by means of WST-1 reduction test in SW982 (left) and SK-LMS-1 (right) cell lines after: (A) 72 h treatment with increasing concentrations of ABT-737; (B) 72 h treatment with increasing concentrations of ABT-737 combined with 25 nM Dinaciclib; (C) 24 h treatment with increasing concentrations of A-1331852; (D) 24 h treatment with increasing concentrations of A-1331852 combined with 25 nM Dinaciclib. Data are presented as means \pm SD. Statistical significance was achieved by the Student's *t* test from at least three different experiments: $*p \le 0.05$; $**p \le 0.001$; $***p \le 0.0001$.

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Supplementary Figure 5 (related to Figure 5): **Combined treatment with Dinaciclib and BH3-mimetics shows deathly liver toxicity.** (A) Treatment regimes using Dinaciclib and ABT-737 in Hsd:Athymic Nude*Foxn1^{nu}* mice. Mortality rates are indicated. (B) Tumor success rate and growth curve for SK-LMS-1 cell line in Hsd:Athymic Nude*Foxn1^{nu}* mice. (C) Treatment regime with Dinaciclib and A-1331852 tested in CB17.Cg-*Prkdc^{scid}Lyst^{bg-J}*/Crl mice. (D) Tumor success rate and growth curve for SK-LMS-1 cell line in CB17.Cg-*Prkdc^{scid}Lyst^{bg-J}*/Crl mice. (E) Microscopical imaging of H&E staining of one intra-peritoneal sample tumor showing tissue transition from renal cortex to tumor mass. * indicate renal corpuscles. (F) Mean CB17.Cg-*Prkdc^{scid}Lyst^{bg-J}*/Crl mice weight variation during combination treatment with Dinaciclib and A-1331852. Time 0 represents the beginning of the treatment.

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A Dinaciclib 40 mg/Kg (i.p.) ABT737 75 mg/Kg (i.p.) meming aftemoon 50 % + moming aftemoon 80 % + В a a 2.0749x San) Řª = -5.225 Tumorvolu (mm³) Tumor SUCC868 80% Ç Time (days) Dinaciciib 20 mg/Kg (i.p.) A-1331852 25 mg/Kg (p.o.) moming afternoon 17%† D y = e^{0.09}x Sent) = -0.761 R Tumorvolu (mm²) Tumor 100% Time (days) Ε F ♦VEHICLES DNACICLIB elghrvariation 1 day 8 (%) A-1331852 BOTH DRUG8 Ĩ Ô ō 500 µm Treatment time (days)