Differential effects of PXD101 (belinostat) on androgen-dependent and androgen-independent prostate cancer models

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Abstract. Histone deacetylase inhibitors (HDACi) are promising epigenetic cancer chemotherapeutics rapidly approaching clinical use. In this study, we tested using in vitro and in vivo models the differential biological effects of a novel HDAC inhibitor [belinostat (PXD101)], in a wide panel of androgen-sensitive and androgen-independent tumor cells. Belinostat significantly increased acetylation of histones H3 and H4. Belinostat potently inhibited the growth of prostate cancer cell lines (IC50 range from 0.5 to 2.5 µM) with cytotoxic activity preferentially against tumor cells. This agent induced G2/M arrest and increased significantly the percentage of apoptosis mainly in androgen-sensitive tumor cells confirming its growth-inhibitory effects. The cell death mechanisms were studied in three different prostate cancer cell lines with different androgen dependence and expression of androgen receptor; LAPC-4 and 22rv1 (androgen-dependent and expressing androgen receptor) and PC3 (androgen-independent not expressing androgen receptor). Belinostat induced the expression of p21 and p27, acetylation of p53 and G2/M arrest associated with Bcl2 and Bcl-Xl downmodulation and significant reduction of survivin, IAPs and Akt/pAkt and increased caspase-8 and -9 expression/activity. Belinostat effectiveness was dependent on the androgen receptor (AR), since the stable transfection of AR greatly increased the efficacy of this HDAC inhibitor. These observations were correlated using in vivo models. We demonstrated that belinostat preferentially resulted in antitumor effect in androgen-dependent tumor cells expressing AR. Our findings provide evidence that belinostat may be a promising anticancer drug for prostate cancer expressing AR, supporting its clinical role in prostate cancer.

Introduction

Prostate cancer is a major health problem and is the second highest cause of deaths from cancer each year in males in the European countries. The hormone therapy for prostate cancer targets the ligand-binding domain of the androgen receptor using drugs that reduce the serum level of testosterone. This is often administered in combination with competitive androgen receptor (AR) antagonists. The outcome of such a treatment follows a predictable course comprised of an initial response, a period of quiescence during which the tumor does not proliferate, followed by progression of the disease. This indicates that androgen directed anti-androgen therapy alone is not sufficient for achieving long-term remission, thus identification of other targets that can be used in conjunction with the current hormonal treatment is warranted. Histone deacetylases (HDACs) are a group of co-repressors of transcriptional activators, including AR (1). These set of proteins regulate gene expression by altering nucleosome conformation at the chromatin level and the stability of several large complexes of transcription factors. Prostate cancer begins as a hormone-sensitive tumor and its growth is stimulated by the activation of AR in conjunction with multiple nuclear co-regulatory proteins such as histone deacetylases. The histone deacetylases modulate transcriptional activity of hormonal receptors including AR by altering the stability of the transcriptional pre-initiation complex and/or modifying the chromatin structure. HDACs associate with other co-repressors and form large protein complexes including Sin3 (Sin3A/Sin3B/HDAC1/HDAC2), HDAC3/GPS/TBL/NCoR/Sin3 complex (2). Wang and co-workers (3) demonstrated that HDAC1 is overexpressed in 35% of prostate cancer, in metastatic tumors and in androgen-independent cell lines. These findings were consistent with the previously reported increase in HDAC1 activity, mRNA expression, and protein expression in prostate

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cancer cell lines (1) and cancers that are refractory to hormone treatment (5). HDAC2 had been found to be increased in human colon cancer and overexpressed in about 30% of patients with prostate cancer (3). HDAC4 was also overexpressed in 23% of prostate cancer specimens (1,3) whereas an altered intracellular localization of HDAC4 in prostate cancers was described in prostate cancer that had become refractory to hormone treatment (6), suggesting that HDAC4 may be involved in the late events of prostate cancer progression. HDAC5 expression was increased in 57% of prostate cancer specimens (3). In addition, it has been demonstrated that HDAC5 is able to associate with known AR co-repressors such as SMRT and N-CoR (7), it may also play a role in tumorigenesis of prostate cancer.

Several HDAC inhibitors have been developed for treatment of ematological and solid tumors. These agents represent a relatively new and interesting class of anticancer agents. Histones play an important role in maintaining chromatin structure, and posttranslational modifications to these molecules serve to regulate chromatin density. As a result of HDAC-inhibition, histones become hyper-acetylated and DNA is maintained in a relatively open conformation that is conducive to interaction with transcription factors. Increased levels of histone acetylation lead to relaxation of the chromatin structure, allowing access of transcription factors and increased transcription, while decreased levels of acetylation are associated with repressed transcription. Consistent with this scenario, HDAC inhibitors have been shown to alter the transcription of a number of genes. In addition, these compounds have been shown to mediate tumor cell differentiation, growth-inhibition and death, and a number of such HDAC inhibitors are in clinical trials. In contrast to the relatively detailed understanding that has been attained regarding the role that histones and their modifications play in maintaining chromatin structure, the mechanism(s) by which HDAC inhibitors mediate anticancer effects are poorly understood.

While histones clearly represent a predominant class of proteins which exhibit HDAC inhibitors-mediated hyperacetylation, a number of non-histone proteins have also been shown to be hyper-acetylated and functionally influenced as a consequence of HDAC inhibitors. Thus, HDAC inhibitors mediate a variety of transcriptional and non-transcriptional alterations in cancer cells. The ability of HDAC inhibitors to affect the activity of a variety of disparate proteins and pathways in tumor cells makes them attractive anticancer agents and also makes the understanding of their mechanism of action a daunting task.

We have been studying a novel hydroxamate, belinostat (previously called PXD101), which inhibits HDAC activity in cancer cell lysates with nM IC₅₀ potency and induces the hyperacetylation of proteins such as histones and tubulin in tumor cells (8). This compound, which is considered a pan-HDAC inhibitor on account of its inhibitory activity on a number of purified recombinant HDACs, was previously shown to inhibit tumor cell growth in vitro and in animal models both as mono-therapy (9,10) as well as in combination with other anticancer agents and (11,12) it is currently being evaluated as an anticancer agent in clinical trials (reviewed in ref. 13). To support the clinical development of belinostat in prostate cancer, we tested, in vitro and in vivo models, the differential biological effects of this agent in a wide panel of androgen-sensitive and androgen-independent tumor cells.

Materials and methods

Reagents. All the materials for tissue culture were purchased from Hyclone (Cramlington, NE, USA). Plastic ware was obtained from Nunc (Roskilde, Denmark). Antibodies were purchased from Santa Cruz (Santa Cruz, CA, USA) unless otherwise indicated. PXD101 (Belinostat) was kindly provided from TopoTarget A/S (Copenhagen, Denmark).

Cell lines. One non-tumor prostate epithelial cell line (BPH-1), 14 PCA cell lines [CWR22 (14), CWR22R-2152 (14), CWR22R-2272 (14), CWR22R-2274 (14), CWR22R-2524 (14), 22r1 (14), LAPC-4 (15), LnCaP, LnCaP-C81 (16), C4-2B (17), PC3, PTEN-transfected PC3 (18), AR-transfected PC3 (19), DU145, AR-transfected DU145, named DU-AR, (20), VCaP (21) and DuCaP (21) cells were used (a complete characterization of the above mentioned cells is reviewed in Pienta et al (22)). CWR22, LAPC-4, LnCaP, VCaP and DuCaP cell lines are androgen-dependent whereas CWR22R derivatives, 22r1, LnCaP-C-81, C4-2B, PC3 and DU145 cell lines are androgen-independent (22).

Growth assays. Briefly, PCa cells (3x10⁴ cells/well) were seeded in 24-well tissue culture plates (Costar, Corning, NY) and left to attach and grow for 24 h. Cells were incubated with belinostat at 0.05-5 µM for 2-72 h dependent on the read out. After this time, cells were maintained in the appropriate culture conditions. Morphological controls were performed every day with an inverted phase-contrast photomicroscope (Nikon Diaphot, Tokyo, Japan), before cell trypsinization and counting. Cells trypsinised and resuspended in 1.0 ml of saline were counted using the NucleoCounter™ NC-100 (automated cell counter systems, Chemotec, Cydevang, Denmark). All experiments were conducted in triplicate. NucleoCounter NC-100 allows to determine also the number of dead cells present in a cell sample. Therefore we considered viable and dead cells as separated entities. IC₅₀ values were calculated by the GraFit method (Erithacus Software Limited, Staines, UK) considering the slopes of inhibition curves obtained for each group of tests. The effect on cell proliferation was measured by taking the mean cell number respect to controls in the time for the different treatment groups.

Apoptosis and cell cycle analysis. Cells (1x10⁴) were fixed for 30 min in 70% ethanol, pelleted by centrifugation (720 g; 5 min). After removal of ethanol by centrifugation, cells were incubated and resuspended in 1 ml of DNA staining solution (PBS containing 200 mg/ml RNase A, 20 mg/ml propidium iodide plus 0.1% Triton X-100) and left at room temperature for 60 min. All cells were then measured on a FACSScan flow cytometer (Becton-Dickinson, UK) and analysed using Cell Quest software (Becton-Dickinson). Resulting DNA distributions were analyzed for the proportion of cells in G₁, S and G₂/M phases of the cell cycle. Apoptotic cells were detected by both the quantifiable peak in sub-G₁ phase, corresponding to the red fluorescence light emitted by sub-diploid nuclei of cells. For statistical comparisons apoptosis was quantified by using a quantitative ELISA tests for DNA fragmentation (Trevigen, Gaithersburg, MD).

Preparation of cell lysates and Western blot analysis. Following treatments, cells grown in 90-mm diameter Petri dishes were
washed with cold PBS and immediately lysed with 1 ml lysis buffer containing a protease and phosphatase inhibitor cocktail. Lysates were electrophoresed in 7% SDS-PAGE, and separated proteins transferred to nitrocellulose and probed with the appropriate antibodies using the conditions recommended by the suppliers.

**In vivo experiments.** Male CD1 nude mice (Charles River, Milan, Italy) were kept in line with University guidelines (University of L’Aquila, Medical School and Science and Technology School Board Regulations, in compliance with Italian government regulation no. 116 of January 27, 1992 on the use of laboratory animals). Before any invasive manipulation, mice were anesthetized with a ketamine (25 mg/ml)/xylazine (5 mg/ml) mixture. All mice received s.c. flank injections of 1x10⁶ in 250 µl of 12 mg/ml Matrigel (Becton-Dickinson, Franklin Lakes, NJ, USA) of 22Rv1 and PC3 cells. Treatments were started when a mean tumor volume of 100 mm³ was reached. Before tumor inoculation mice were randomly assigned to 4 experimental groups of 12 mice each. For each cell line one control group received an intraperitoneal (i.p.) injection of 100 µl PBS for 18 consecutive days and one group received 20 and 40 mg/kg/dose bid for 18 consecutive days. The effects on tumor growth of different treatments were evaluated by: i) tumor volume measured during and at the end of experiment by biweekly measurement of tumor diameters with a Vernier caliper (length x width). Tumor volume was calculated according to the formula: TV (mm³) = π x d² x D/2, where d and D are the shortest and longest diameters, respectively (23). ii) Tumor weight, measured at the end of experiment. Experiments were stopped 30 sec/day after the first day of treatment, mice were sacrificed by carbon dioxide inhalation and tumors were removed surgically. Tumor growth delay (TGD) was determined as % TGD = ((T-C)/C)₁₀₀, where T and C are the mean times in days required to reach the same fixed tumor volume in the treated group and control group, respectively (24,25). In vivo drug combination studies were evaluated by CalcuSyn (Biosoft, Cambridge, UK). For the calculation of CI, the values of cell kill for a fixed tumor volume were considered (determined by the log cell kill (LCK)). Log cell kill was determined as LCK = (T-C)/ (3.3-Td), where Td represents the mean control group doubling time required to reach a fixed tumor volume, expressed in days, whereas T and C are the same values as described above (24,25).

Half of the tumor was directly frozen in liquid nitrogen for protein analysis and the other half fixed in paraformaldehyde overnight for immunohistochemical analyses. Masson's trichrome staining, CD31, Ki67, K18, VEGF, pAkt (Ser⁴²⁸) and Her2 expression was evaluated in 4 µm tissue sections cut from blocks selected for the presence of representative tumor tissue. Negative controls were incubated only with universal negative control antibodies under identical conditions, processed, and mounted. Expression of K18, VEGF, pAkt (Ser⁴²⁸) and Her2 was subjectively graded and quantified as 0, 1, 2, 3, 4, and 5 representing not detectable, weak, moderate, strong, and very strong staining, respectively. Tissue disorganization and vessel distribution were also analyzed using a variation of Masson trichrome staining (MBS-HS443, HD supplies, Aylesbury, UK).

**Statistical analysis.** Continuous variables were summarized as mean and standard deviation (SD) and were compared by the unpaired Student's t-test. Dichotomous variables were summarized by absolute and/or relative frequencies and were compared by the exact Fisher's test or χ² test when appropriate. For multiple comparisons the level of significance was corrected by multiplying the P-value by the number of comparisons performed (n) according to Bonferroni correction. All tests were two-sided and were determined by Monte Carlo significance. P-values <0.05 were considered statistically significant. All statistical analyses were performed using the SPSS® statistical analysis software package, version 10.0.

**Results**

**Belinostat results in acetylation of histonic and non-histonic proteins in vitro.** It has been demonstrated that HDAC inhibitors increase histone H3 and H4 acetylation. In Fig. 1A we demonstrate that belinostat resulted in acetylation of H3 and H4 histones in LnCaP cell line in a dose-dependent manner. At the same time, we also demonstrate that belinostat was able to induce acetylation of the non-histonic protein p53 involved in cell proliferation and apoptosis (Fig. 1). As shown in Fig. 1B acetylation of histones occurred in other cell models (PC3, 22rv1, LAPC-4 and LnCaP cell lines) with kinetics and EC₅₀ and EC₃₀ (concentration of belinostat able to result in acetylation of 20 and 50% of H3 and H4 histones) with characteristics typical of the different cellular models. Fig. 1C demonstrates that 0.1 µM Belinostat, responsible for a sub-total acetylation of histones, induced acetylation of H3 and H4 histones as well as of p53 within 8 h of treatment, raising maximal values after 16-24 h of culture. Histone acetylation was maintained at high levels until at least 72 h.

**Belinostat results in differential antiproliferative and cytotoxic effects in androgen-sensitive and androgen-independent tumor cells in vitro.** Dose-dependent experiments with belinostat ranging from 0.05 to 5.00 µM were used to calculate the IC₅₀ and IC₃₀ values for proliferation, viability and apoptosis in the different prostate cancer cell models. In Fig. 2A we show the growth inhibition measured after 24-, 48- and 72-h incubation at 0.05-1 µM belinostat in a panel of cancer cell lines with high hormone sensitivity (LAPC-4 and LnCaP) and low hormone sensitivity (22rv1 and PC3) and in a non-tumor cell line (BPH-1). In these cellular models, belinostat inhibited cell proliferation in a dose-dependent manner in cancer cells. Overall the antiproliferative effects of belinostat was greater in androgen-dependent than androgen-independent tumor cells with IC₂₀ and IC₅₀ values significantly lower for androgen-dependent models. In non-tumor cell line BPH-1 belinostat resulted in antiproliferative effect only at the higher drug concentrations (Fig. 2A). Using the NucleoCounter NC-100 we determined also the number of dead cells as marker of cytotoxic effect of Belinostat. This drug induced cell toxicity in a time-dependent manner (Fig. 2B). The increment of mortality was strongly dose-dependent and clearly documentable in cancer cells but not in BPH-1 cells. As in the case of the antiproliferative effects, belinostat were significantly more effective in AD than AI cell lines with EC₂₀ and EC₅₀ values significantly lower for androgen-dependent models (Fig. 2B).
Figure 1. (A) Dose-dependent effects of belinostat measured after 24 h of treatment in LnCaP cell cultures. (B) Summarized effects of histone H3 and H4 acetylation on PC3, 22rv1, LnCaP, DU145, C4-2B and LAPC-4 cell lines. (C) Time-dependent effects of 0.1 µM belinostat administered to LnCaP cell culture. Each data are representative of three individual experiments. Each Western blot lane was loaded with 40 µg of proteins and normalized vs actin.

Figure 2. (A) Variation in cell viable number in the time and at different drug concentration in the PC3, 22rv1, LnCaP and LAPC-4 cell lines: the number of cells are normalized as percent of inhibition vs control to calculate IC_{20} and IC_{50} values, representation of growth inhibition curves in non-tumor BPH-1 (IC_{50} > 5 µM) cell line and neoplastic PC3 (IC_{50} = 1.35±0.32 µM), 22rv1 (IC_{50} = 1.23±0.45 µM), LnCaP (IC_{50} = 0.84±0.16) and LAPC-4 (IC_{50} = 0.68±0.15 µM) cell line. (B) Variation in cell death number in the time and at different drug concentration in PC3, 22rv1, LnCaP and LAPC-4 cell lines: the number of cells/hour are normalized as percent vs control to calculate efficacy concentration (EC_{20} and IC_{50}) values, comparison of the effects in prostate cancer cell lines vs non-tumor/BPH-1 cell line.
Figure 3. Effects of belinostat at different time of culture on proliferation (A-D), apoptosis (E-H) and caspase activation (I-N) in PC3 and DU145 after AR transfection.
if AR modified belinostat sensitivity, two androgen-independent and AR-negative tumor cells were stably transfected with the wild-type AR and then subjected to increasing concentrations of 10-12 M belinostat (Fig. 3A-D). The re-expression of a functional wild-type AR in androgen defective PC3 and DU145 cells resulted in a significant slow-down of basal growth rate of these cells compared to parental ones in standard culture conditions. Belinostat significantly decreased the number of viable PC3 and DU145 parental cells starting from the higher concentrations of 1 and 5 µM (Fig. 3A). The same tumor cells transfected and forced to express AR acquired more sensitivity to belinostat, which significantly decreased the number of viable cells starting from 0.1 µM belinostat. The same trend was clearly visible if apoptosis was used as efficacy indicator. An increment in the DNA fragmentation (Fig. 3E-H) as well as in the activity of caspase-3, -8 and -9 (Fig. 3I-N) was observed upon treatment with belinostat in PC3 and DU145 cells transfected with AR when compared with parental cells.

Sub-toxic concentrations of belinostat modulate cell cycle transition and results in apoptosis in vitro. Sub-toxic concentrations of belinostat corresponding to IC<sub>20</sub> values were associated with significant modifications of cell cycle phase distribution. At these concentrations, belinostat induced a preferential accumulation of tumor cells in the G<sub>2</sub>/M (Fig. 4A) phase of cell cycle the increase of cyclin D1, cdk4, p-Rb, p16, p21 and p27 and decrease of cyclin A and B1. The levels of p53 were also increased in p53 expressing LnCaP, 22rv1 and LAPC-4 prostate cancer cells. In Fig. 4B and C a representative Western blot analysis is shown performed in PC3 (Fig. 4B, p53 and AR defective) and 22rv1 (Fig. 4C, p53 and AR expressing) PCa cell lines treated with 0.5 µM belinostat. The accumulation of tumor cells in G<sub>2</sub>/M was associated with a dose-dependent increase of apoptosis. Fig. 4D shows the FACS appearance of 22rv1 tumor cell line treated with 0.2, 0.5 and 1.0 µM belinostat. For statistical purpose the quantification of apoptosis among the different cell lines was performed by a quantitative ELISA tests for DNA fragmentation.
Fig. 4E shows a dose-dependent effect in terms of apoptosis in PC3, 22rv1, LnCaP and LAPC-4 cell lines. Apoptotic effects of belinostat were significantly greater in androgen-sensitive (LAPC-4 and LnCaP) than androgen-independent cell lines (22rv1 and PC3). The pathways involved in the Belinostat-induced apoptosis were also studied at the concentration of 0.1 µM. We observed a modulation of mediators involved in the apoptotic mechanisms. Specifically a down-modulation of Bcl2, Bcl-Xl, survivin, XIAP, c-IAP-1, TRAF-1 with a simultaneous up-modulation of Bax, Bcl-Xs, cytochrome c and smac/Diablo. The intrinsic moiety was dependent on FasL up-modulation and caspase-8 activation. Fig. 4F and G show the modulation of pro- and anti-apoptotic proteins in LnCaP (F) and 22rv1 (G) representing two AR expressing PCa cell lines having or not androgen sensitivity. These changes were time-dependent and achieved maximal effect after 36-48 h from beginning of treatment.

Anticancer activity of PXD101 in human prostate xenografts. For in vivo studies, two models of tumor cells with graded androgen sensitivity and AR expression were chosen. The PC-3 xenograft model, androgen-insensitive, no AR expression (26) and 22rv1, androgen-sensitive, AR expression (27). As indicated in Table I, the treatment with 20 and 40 mg/kg twice daily i.p. of belinostat was well tolerated by nude mice xenografted with PC3 and 22rv1 cells. As surrogate end-point of toxicity the changes in the body weights of xenografted mice were assessed after treatments. No significant difference was observed in the body weights of xenografted mice treated with belinostat at the dosage of 20 and 40 mg/kg suggesting a low toxicity profile. In Fig. 5 we show the...
trend over time of tumor volumes generated by PC3 and 22rv1 cells upon belinostat treatment. At the treatment dose of 20 mg/kg belinostat, tumor growth of PC-3 and 22rv1 xenografts was slowed down by 33.7 and 44.0% compared to controls. The tumor growth of PC-3 and 22rv1 xenografts was further slowed down at the treatment dose of 40 mg/kg belinostat with a decrease of 46.3 and 66.5%, respectively. Upon treatment with 20 and 40 mg/kg belinostat, the tumor growth delay (TGD) was 7.5 (±2.1) and 14.4 (±3.6) days in PC3 xenografts (p<0.001) and 11.6 (±3.4) and 16 (±2.2) days in 22rv1 xenografts (p<0.001). However, in terms of TGD, belinostat was more effective in 22Rv1 (androgen-sensitive but not -dependent) than in PC3 model (androgen-insensitive) at both dosages as clearly documented in Table I. A complete or partial response, defined as the disappearance or a reduction of >50% of a volume, was documented only in 22rv1 xenografts treated with 20 mg/kg (1 out of 10) and 40 mg/kg (3 out of 10) with a dose-dependent effect (Table I). Belinostat resulted in a significant modulation of Ki67 labeling index levels, apoptosis and angiogenesis compared to controls. In particular, the angiogenesis and the number of tumor cells labeled with Ki67 were significantly reduced compared to controls whereas apoptosis levels increased in both xenograft models in a dose-dependent manner. Immunohistochemical studies in PC3 xenografts (Fig. 5C) revealed a reduction of angiogenesis as documented

Figure 5. (A) Tumor proliferation in PC3 and 22rv1 xenografts in presence of 20 and 40 mg/kg/bid i.p. (B) Microscopic appearance of PC3 and 22rv1 tumors stained with Martius yellow-brilliant crystal scarlet blue technique. The presence of large blood vessels (stained in yellow/orange) near cancer cell nests of untreated PC3 tumors is shown. Untreated PC3 tumors show the presence of numerous and small blood vessels scattered in the tumor mass and delineating single tumor cell nests. Belinostat reduces blood vessel number. Vessels appear dilated with unstructured capillary bed dispersed in collagen I deposits (azure staining) with deposition of fibrin clots (pale pink staining) and formation of dense collagen I deposits (fibrosis) which envelop tumor cell nests. Evaluation of neoangiogenesis through CD31 stain. VEGF and p-Akt staining were reduced after belinostat treatment. Single pictures are at magnification x400.
Table I. Antitumor activity of PXD101 in 22rv1 and PC3 xenograft.

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<th>Treatment</th>
<th>Drug</th>
<th>Dose mg/kg</th>
<th>Weight of mice Gr ± SD</th>
<th>Tumor weight (mg ± SD)</th>
<th>TGD (days)</th>
<th>PI Ki67%</th>
<th>Apoptosis (%)</th>
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by CD31 Masson's trichrome staining. The immunostaining for VEGF was also sensitively reduced after belinostat treatment as well as K18 and p-Akt.

Discussion

The increase in HDAC and DNMT expression could represent an active role of HDACs and DNMTs in prostate cancer or a cancer epi-phenomenon. It has been reported previously that hydroxamic acid-based inhibitors such as belinostat, are potent inducers of differentiation and/or apoptosis in transformed cells in culture including prostate cancer cells (10,28,29). Here we confirm that Belinostat, a member of hydroxamic acid-based inhibitors, suppresses, in vitro and in vivo, the growth of a wide panel of prostate tumor cells with graded androgen dependence at micromolar concentrations. Here we demonstrate that belinostat can suppress the growth of both androgen-dependent and -insensitive prostate cancer cells in culture at micromolar concentrations, and that belinostat suppress tumor growth to a higher extent in nude mice bearing human androgen-dependent 22rv1 prostate tumors than androgen-independent PC-3 prostate tumors. This strongly indicates a functional relevance of increased HDACs in prostate cancer rather than cancer epi-phenomenon. In addition, HDACs regulate expression of many genes related to this event. In the belinostat-treated tumors, expression of Bax and caspase-3 increased with a contextual decrease of the anti-apoptotic gene Bcl-2. However, belinostat results in a cytotoxic and pro-apoptotic activity of P21 and P27 and with the preferential accumulation of tumor cells in the G/M phase of the cell cycle. The cell cycle block was followed by apoptosis, as described for other HDAC inhibitors, and was associated to both intrinsic and extrinsic apoptotic program (37). Apoptosis induction by belinostat was manifested at higher concentrations and was associated with the expression of many genes related to this event. In the belinostat-treated tumor cells, expression of Bax and caspase-3 increased with a contextual decrease of the anti-apoptotic gene Bel-2. However, belinostat was well tolerated by nude mice and tumor growth was significantly suppressed by 33.7 and 44.0%, and by 46.3 and 66.5%, after treatment with 20 and 40 mg/kg belinostat bid, respectively, in PC-3 and 22rv1 xenografts. A complete or partial remission of the tumor was observed in 22rv1 xenografts treated with 20 mg/kg/bid (1 out 10) and 40 mg/kg/bid (3 out 10).

In conclusion the evidence here reported indicates that belinostat results in a cytotoxic and pro-apoptotic activity preferentially in androgen-sensitive preclinical models and this activity is influenced by AR and this supports further studies...
to evaluate the effects of belinostat with hormone manipulation therapies such as castration or treatments with the anti-androgen bicalutamide.

References


