Epigenetic modulation of AR gene expression in prostate cancer DU145 cells with the combination of sodium butyrate and 5'-Aza-2'-deoxycytidineline

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Abstract. The androgen receptor (AR) plays an essential role in the development and progression of prostate cancer. Castration-resistant prostate cancer (CRPC) is a consequence of androgen deprivation therapy. Unchecked CRPC followed by metastasis is lethal. Some CRPCs show decreased AR gene expression due to epigenetic mechanisms such as DNA methylation and histone deacetylation. The aim of this study was to epigenetically modulate the methylated state of the AR gene leading to targeted demethylation and AR gene expression in androgen-independent human prostate cancer DU145 cell line, representing the CRPC model with very low or undetectable AR levels. The cell treatment was based on single and combined applications of two epigenetic inhibitors, sodium butyrate (NaB) as histone deacetylases inhibitor and 5'-Aza-2'-deoxycytidine (Aza-dC) as DNA methyltransferases inhibitor. We found that the Aza-dC in combination with NaB may help reduce the toxicity of higher NaB concentrations in cancer cells. In normal RWPE-1 cells and even stronger in cancer DU145 cells, the combined treatment induced both AR gene expression on the mRNA level and increased histone H4 acetylation in AR gene promoter. Also activation and maintenance of G2/M cell cycle arrest and better survival in normal RWPE-1 cells compared to cancer DU145 cells were observed after the treatments. These results imply the selective toxicity effect of both inhibitors used and their potentially more effective combined use in the epigenetic therapy of prostate cancer patients.

Introduction

Castration-resistant prostate cancer (CRPC) develops over time as a consequence of androgen deprivation therapy (1,2) and this is enhanced by epithelial-mesenchymal transition and cancer stem cells (reviewed in ref. 3). Unchecked CRPC followed by metastasis is lethal (4) due to limited options for treatment. The mechanisms by which prostate cancer cells acquire castration resistance are numerous and include activation of alternative pathways with dominant PI3K/AKT signaling, androgen receptor (AR) gene mutations leading to promiscuous activations (5,6), and recently identified AR splice variants among other factors (7,8).

DNA methylation is one of the most intensely studied epigenetic modifications that appears to be a decisive event in the initiation and development of advanced CaP with the process of DNA hypermethylation preceding to global hypomethylation (9,10). Friedlander et al (4) compared CRPC and benign tissue methylation profiles with the finding of hypermethylated genomic DNA in CRPC patients. Moreover, in these patients, DNA methylations at individual CpG loci both within and outside CpG islands were found much more frequently than common copy number changes. Methylated DNA could thus be a treatment target for delaying the progression of the castration-resistant disease using PI3K/AKT inhibitors and hypomethylating agents (4).

AR plays an essential role in advanced CaP (11). Some CRPC expresses the AR gene in autocrine pathways and remains dependent on AR while other cancer cells show decreased AR
gene expression attributed to impaired AR protein stability, X chromosome loss or DNA methylation silencing and are independent of AR (7). Tian et al (12) showed a link between AR gene methylation and prostate cancer progression. These authors found that AR gene methylation in promoter regions was likely related to prostate stem/progenitor cell stemness and differentiation. Low expression of the AR in prostate cancer stem cells and LNCaP progenitor/stem cells were found to be due to high DNA methyltransferase 1/3 level and MBD2 promoter binding. Moreover, treatment of prostate cancer cells with 5 µM 5'-Aza-2'-deoxycytidine (Aza-dC) resulted in the inhibition of self-renewal/growth of prostate stem/progenitor cells in vitro, reduced prostate tumorigenicity in vivo followed by induction of the AR gene and functional protein expression in a time-dependent manner following 6 days incubation with 5 µM Aza-dC.

The hypothesis that epigenetically induced AR expression in CRPC with AR methylated pattern might revert some deleterious pathways and to some extent reduce the aggressiveness of the cancer cells, is supported by several studies. McCabe et al (13) for example found that Aza-dC inhibited aberrant de novo DNA methylation in the TRAMP mouse model and prevented CaP development during the drug administration. Zorn et al (14) reported the delayed emergence of androgen-independent CaP in castrated TRAMP mice after Aza-dC treatment. Moreover, combined treatment by castration and Aza-dC administration showed statistically significant longer survival than single treatment. In a preclinical study, Gravina et al (15) used 5-azacytidine for reactivation of AR gene expression, silenced by DNA methylation in the PC3 cell line. This led to re sensitisation to bicalutamide (BCLT) responsiveness and subsequent apoptosis. In detail, 5-azacytidine treatment increased the effect of BCLT therapy in AR-expressing and AR-deficient prostate cancer, both in vitro and in vivo. Co-treatment with both agents led to synergistic/additive effects in nude male mice xenografted with 22rv1 and PC3 cells (AR-expressing or AR-deficient cell lines, respectively) followed by significantly reduced tumor mass and delayed cancer progression. In the co-treated cell lines (22rv1 and PC3), increased cell cycle and apoptosis proteins expression were observed.

This study describes the epigenetic consequences of the combined treatment of two inhibitors, sodium butyrate (NaB) as a histone deacetylase (HDAC) inhibitor and 5'-Aza-2'- deoxycytidine (Aza-dC) as a DNA methyltransferase (DNMT) inhibitor, in both cancer and normal prostate cells. DNA methyltransferases (DNMTs) and histone deacetylases (HDACs) are co-regulators of the AR and could imply potential targets for affecting androgen receptor function and stability. These are now an option in epigenetic treatment (16) with the aim of overcoming the mechanism of hormonal resistance and in its consequences, of target therapy to regulation of the AR without therapy based on hormone treatment. Our previous results (17) showed demethylation of specific CpG sites in the AR gene in the DU145 prostate cancer cell line following co-treatment with Aza-dC and NaB. The focus of this study was the methylated AR gene with subsequent epigenetic modulation leading to targeted histone acetylation and AR gene re-expression. Our results imply that the epigenetic drugs used, depending on the concentration, affected the acetylation level of histones H3 and H4 in a vicinity of the AR gene promoter. In addition, the used epigenetic agents induced activation and maintaining of G2/M cell cycle arrest in RWPE-1 cells. Better survival in normal RWPE-1 cells compared to cancer DU145 cells implies the selective toxic effects of the used inhibitors.

Materials and methods

Cell culture, treatment conditions and viability assay. The androgen-independent human prostate cancer cell line DU145 was purchased from ATCC (Rockville, MD, USA), maintained in DMEM (Gibco) supplemented with 10% fetal bovine serum (FBS), 0.01% antibiotics, 2 mM L-glutamine. Non-tumorigenic, immortalized human prostate cell line RWPE-1 was kindly provided by the Department of Experimental Biology, Masaryk University (Brno, Czech Republic). The normal cell line RWPE-1 was cultivated in keratinocyte-SFM medium (kit) with L-glutamine, human recombinant epidermal growth factor (EGF) and bovine pituitary extract (BPE) (Gibco) supplemented with a final 0.01% concentration combination of penicillin and streptomycin, 0.01% concentration of amphotericin and 0.005% concentration of gentamycin. All cells were maintained at 37°C and 5% CO₂ atmosphere. Both cell lines were treated with NaB (Sigma-Aldrich), Aza-dC (Sigma-Aldrich) and their combinations for 2 and 6 days when medium and agents were changed after 2 days. Cell viability assays of cells were performed using 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide agent (MTT) as described (17). The cancer cell line DU145 was seeded on 96-well plates at 4,500 cells per well and RWPE-1 cells at 6,000 cells per well and increased to 40-50% confluent prior to the first treatment. The cell culture medium with inhibitors was changed after 2 days for 6-day treatments. All treatments were performed in triplicate. The percentage of viable cells was calculated as follows: average absorbance of treated cells/ average absorbance of control cells x 100.

mRNA analysis using RT-qPCR. The cells were seeded in 100-mm dishes (1x10⁶ cells for RWPE-1 and 7x10⁵ for DU145 cells) and treated with 5 µM Aza-dC, 5 mM NaB and their combinations (0.5 µM Aza-dC + 5 mM NaB), 0.5 µM Aza-dC + 5 mM NaB and 5 µM Aza-dC + 5 mM (NaB) for 2 and 6 days when medium and agents were exchanged after 2 days in prolonged 6-days culture. Total RNA from both cell lines was isolated with High Pure RNA Isolation kit (Roche) and 1,000-1,500 ng of total RNA was converted to cDNA using Transcriptor First Strand cDNA synthesis kit (Roche), in both cases according to the manufacturer's instructions. For following quantitative real-time PCR analyses, the total amount of 100 ng of cDNA from each sample was amplified using Taq-Man probes labelled with hexafluorescein and thermo-Start DNA Polymerase (AB gene) using the real-time PCR analyser Rotor-Gene RG-3000 (Corbett Research). Primers and probe sequences for AR, PSA (prostate specific antigen) and GAPDH genes are summarized in Table I. The experiments were performed in triplicate with a similar pattern of results.

Chromatin immunoprecipitation coupled with quantitative PCR (ChIP-qPCR). ChIP was performed on both cell lines as described (17) with the following modifications: briefly, the cells were seeded in 100-mm dishes (1x10⁶ cells for RWPE-1)
and in 150-mm dishes (1.9x10^6 cells for DU145). After a formaldehyde cross-linking terminated by glycine, RWPE-1 cells were washed with Dulbecco's phosphate-buffered saline (D-PBS) instead of standard PBS used for DU145 line. Cells were immunoprecipitated with 2 µg anti-acetyl histone H3 rabbit polyclonal antibody (cat. 06-599), and normal mouse IgG polyclonal antibody (cat. 06-866, both from Millipore), and for input, 100 µg of each chromatin lysates and 1 µl of proteinase K (100 µg/ml) were added to each tube, incubation at 55˚C for 3 h with gentle agitation was performed followed by DNA purification with QIAqwick® PCR Purification kit (Qiagen). Quantitative PCR analyses were performed on LightCycler 480 (Roche) with probe no 51 (Universal Probe Library cat. no. 04688481001) using ProbeFinder assay design software and primers amplifying AR gene promoter region (Table I). DNA from each input sample was diluted 10x and primer efficiency was tested. All experiments were repeated three times independently and all measurements were performed in triplicate.

Cell cycle analysis. Briefly, the cell lines were seeded in 6-well plates (2.2x10^5 cells for RWPE-1 and 1.5x10^5 cells for DU145) and treated with the same inhibitors and their combinations for 2 and 6 days as described above. Cells were harvested at indicated times after treatment, also detached cells were collected. Samples were fixed in cold 70% ethanol. After treatment with RnaseA, samples were stained with propidium iodide (PI). Cellular DNA content was analyzed using flow cytometry (BD FACSuite, BD, USA) and collected data were processed using BD FACSuite (BD). At least 10,000 cells per sample were analyzed.

Statistical analysis. Multifactoral analysis of variance (ANOVA) with post-hoc two-tailed Dunnett's t-test and Bonferroni multiple comparison test were used for cell viability assays and quantitative experiments. All statistical analyses were performed with the SPSS software version 15 (SPSS, Inc. Chicago, IL, USA) and the significance level was set at p<0.05 (two-tailed).

Results

Aza-dC decreases cytotoxicity of NaB in prostate cancer cells. Cell viability was analysed using MTT assay to compare cell cytotoxicity after treatment with one of the inhibitors used or inhibitor combinations. In DU145 cells treated with NaB or Aza-dC, or combinations of different concentrations of Aza-dC with NaB (Fig. 1), the cell viability was significantly lower following all treatments after 6 days than in 2-day experiments (ps<0.0004 for Aza-dC treatment, ps<0.0001 for co-treatment, Fig. 1B and C, respectively). The 6-day treatment with NaB (Fig. 1A) showed significantly lower cell viability, ranging from 2.5 mM concentration (p<0.002 and p<0.0001 for 4 and 5 mM NaB), and higher cell viability after treatment with 0.25 and 0.5 mM concentrations (p=0.0001 for both treatments) compared with the same conditions in the 2-day experiment. The images of prostate cancer cells DU145 (Fig. 2) demonstrate changes in cell viability after 2 and 6 days of cultivation with 5 mM NaB and with 0.5 µM Aza-dC + 5 mM NaB co-treatment compared with DMSO. Changes of cell viability in DU145 cells are comparable with the results of MTT assay (Fig. 1A and C) and suggest a lower cytotoxicity of 0.5 µM Aza-dC + 5 mM NaB combination used than 5 mM NaB treatment alone.

In normal RWPE-1 cells, the 6-day incubation led to significantly lower cell viability on treatment with NaB alone (p<0.0001; Fig. 1D) and NaB co-treatment with Aza-dC (p<0.0008; Fig. 1F) in all concentrations compared to 2 days. The highest 4 and 5 µM Aza-dC concentrations caused higher toxicity after 6 days than the 2-day experiment (Fig. 1E, p=0.028 and 0.007, respectively). In contrast, no differences in cell viability were observed on treatment with 0.5-2 µM Aza-dC concentrations between 2- and 6-day experiments.

A combination of the Aza-dC and NaB treatment induces AR gene re-expression in prostate cancer cells. When designing the RT-qPCR, we followed results from the MTT assay (Fig. 1) and from our previous bisulfite sequencing results in DU145 cells (17), where the most effective DNA demethylation effect was after 2-day treatment with 0.5 µM Aza-dC + 5 mM NaB in the AR gene. Here in DU145 cells (Fig. 3A and B), co-treatment with the combination of 0.5 µM Aza-dC + 5 mM NaB led to most significantly higher AR gene re-expression after both 2- and 6-day incubations. The 5 µM Aza-dC cell treatment had no effect on AR gene re-expression after 2 days (Ct undetectable; Fig. 3A) as was further confirmed by mRNA analysis, although a weak signal detection was observed after the 6-day exposure (Fig. 3B). As the 0.5 µM Aza-dC + 5 mM NaB co-treatment, the 5 mM NaB treatment and 5 µM Aza-dC

<table>
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<th>Reverse (5'-3')</th>
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<tr>
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<td>GGCTTTGGAGAAACAAGTGC</td>
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GOI-gene of interest.
+ 5 mM NaB combination were also effective in inducing significant AR mRNA re-expression.

In normal RWPE-1 cells (Fig. 3C and D), we found that AR gene expression was enhanced by all treatments with the highest effect after 5 mM NaB treatment, where only the 2-day experiment showed statistically significant differences between 5 mM NaB, 0.5 µM Aza-dC + 5 mM NaB and 5 µM Aza-dC + 5 mM NaB co-treatments.

Comparing the results from the mRNA analysis in both cell lines used (Fig. 3), a tendency to overall decrease in AR mRNA expression after 6-day treatment was observed. The treatments with 5 µM Aza-dC and 0.5 µM Aza-dC + 1 mM NaB in DU145 and RWPE-1 cells were weak or ineffective in re-expressing the AR. However, no PSA gene expression at either time course or in either cell line was detected (Ct undetectable).

The NaB increases histone H4 acetylation, but not histone H3 in prostate cancer cells affected with epigenetic modulators. Since histone deacetylation is a hallmark of silent condensed chromatin, to explore whether single NaB or its combinations with Aza-dC could be effective in initiating the AR gene re-expression through histone re-acetylation, we used the same treatment as described for mRNA analysis followed by chro-

Figure 1. DU145 (A-C) and RWPE-1 (D-F) cell viabilities after 2 and 6 days treated with NaB, Aza-dC and Aza-dC + NaB combinations. The data at each treatment point represent the means ± SE of triplicate wells. Asterisks show statistically significant differences between treatment and control DMSO. *p<0.05 value, **p<0.0001.

Figure 2. Visual appearance of DU145 cell line after treatment with 5 mM NaB and the combination 0.5 µM Aza-dC + 5 mM NaB after 2 and 6 days.
matin immunoprecipitation coupled with qPCR. Sonicated chromatin samples were immunoprecipitated with anti-acetyl histone H3-lysine 4, -lysine 9, -lysine 14 and -lysine 18 (H3K4, H3K9, H3K14 and H3K18, respectively) and anti-acetyl histone H4-lysine 5, -lysine 8, -lysine 12 and -lysine 16 (H4K5, H4K8, H4K12 and H4K16, respectively) antibodies. Normal IgG antibody served as a negative control, and sonicated chromatin samples were processed with no added antibody (NoAb sample) as a mock control (or noise control). The NoAb sample was processed as the standard sample used for IP without antibody and served as an internal control to minimize noise of the manual workflow. ChIP-qPCR results were analyzed by evaluating the signal of enrichment over noise normalized to input.

In DU145 cells treated for 2 days with 0.5 µM Aza-dC + 1 mM NaB and the 0.5 µM Aza-dC + 5 mM NaB (Fig. 4A), co-treatments were significantly more effective for histone H3 acetylation than the DMSO control. Monitored histone H4 sites were acetylated following all treatments except for the 5 µM Aza-dC treatment (Fig. 4A). In 6 days (Fig. 4B), the histone H3 acetylation increased after cell treatment with 5 mM NaB alone and with the 5 µM Aza-dC + 5 mM NaB combination, and decreased after treatments with 5 µM Aza-dC, 0.5 µM Aza-dC + 1 mM NaB and 0.5 µM Aza-dC + 5 mM NaB in comparison with untreated DMSO. We observed significant increase in histone H4 acetylation in the AR gene promoter region for all treatments, while the 5 µM Aza-dC treatment was ineffective (Fig. 4B).

In the RWPE-1, after 2-day treatment (Fig. 4C), we found significant decrease in the histone H3 acetylation targeted to the AR gene promoter for all used treatments except for 5 mM NaB. Significant upregulation of histone H4 acetylation was observed in cells treated with 5 mM NaB alone and with subsequent combinations of 0.5 µM Aza-dC + 5 mM NaB and 5 µM Aza-dC + 5 mM NaB. Downregulation of histone H4 acetylation was observed following treatment with 0.5 µM Aza-dC + 1 mM NaB combination but no significant change was detected after treatment with 5 µM Aza-dC alone. After 6 days (Fig. 4D), the RWPE-1 cells showed lower levels of histone H3 acetylation for all used treatments except 0.5 µM Aza-dC + 5 mM NaB application compare to DMSO.

Comparing histone H3 and H4 acetylations between 2- and 6-day experiments, we found a decrease in histone H3 acetylation in DU145 cells treated with 0.5 µM Aza-dC + 1 mM NaB (Fig. 4A) and 0.5 µM Aza-dC + 5 mM NaB (Fig. 4B). However, 6 days exposure (Fig. 4D) showed a decrease in histone H3 acetylation for all treatments except 0.5 µM Aza-dC + 5 mM NaB application compared to DMSO.
The DU145 cell line showed significantly decreased histone H4 acetylation after all treatments except the control DMSO and 5 µM Aza-dC applications that led to increased histone H4 acetylation. In normal RWPE-1 cells (Fig. 4C and D), the control DMSO, 5 µM Aza-dC and 0.5 µM Aza-dC + 5 mM NaB treatment showed higher levels of histone H3 acetylation in comparison with shorter 2-day incubation (p=0.036, p=0.025 and p=0.037, respectively). We found no significant changes in histone H3 acetylation for the other treatments. For increasing histone H4 acetylation in normal prostate cells effective was the 0.5 µM Aza-dC + 5 mM NaB combination. Other treatments were significantly less effective after 6-day exposure.

Cell cycle distribution explains the NaB induced cell death in prostate cancer cells. To determine the influence of Aza-dC and NaB on cell cycle regulation, exponentially growing RWPE-1 and DU145 cells were treated with the same treatment scheme as for RT-PCR and ChIP analysis. Cell cultures, including detached cells, were collected, stained with PI and DNA content was assessed by flow cytometry (Fig. 5). By comparing the profiles of control (DMSO) and Aza-dC treated cell cultures after 2 days, the accumulation of cells in S and G2/M stages was shown moderately for RWPE-1 cells and more prominent in DU145 cells. Prolonged Aza-dC treatment (6 days) revealed obvious difference in effectiveness of G2/M arrest between compared cell lines. While significant and specific accumulation of RWPE-1 cells in the G2/M compartment of the cell cycle was observed, the DU145 cells massively died, yielding predominant sub-G1 population reaching up to 80%. The treatment with 5 mM NaB resulted in a much more conclusive contrast between normal and cancer cells. The RWPE-1 cells survived 2-day NaB treatment effectively, employing the G2/M arrest, while 79% of DU145 cells was shifted to sub-G1 compartment, indicating cell death. After 6 days of the NaB treatment, more than one half of RWPE-1 cells were cycling, at the same time >90% of DU145 cells were dead, belonging to sub-G1 cell population.

Combination treatment at lowest concentration (0.5 µM Aza-dC +1 mM NaB) was more efficient than the Aza-dC treatment alone. G2/M accumulation was observed after 2-day treatment, with appearance of the sub-G1 cell population...
specifically in DU145 cells. Prolonged 6-day treatment led to divergent response of both cell lines, revealing effective G2/M arrest for RWPE-1 cell line and high degree of cell death for DU145 exhibiting <20% of cells in the cycle. Combination
therapy with higher concentrations (0.5 µM Aza-dC + 5 mM NaB and 5 µM Aza-dC + 5 mM NaB) resulted in generation of strong sub-G1 population in DU145, reaching >80% within 2 days of treatment. More than 95% of DU145 cells were shown to be sub-G1 for both concentrations after 6 days, suggesting that at 0.5 µM Aza-dC + 5 mM NaB combination treatment reached its plateau. Highest concentrations of combination treatment resulted in dose-dependent G2/M accumulation of RWPE-1 cells after 2 days, followed by appearance of sub-G1 cells after day 6.

Discussion

Histone modification and DNA methylation are associated with transcriptional repression and integrally linked with following synergistic/additive effects (19). In our study we used two types of epigenetic agents: sodium butyrate, a naturally occurring HDAC inhibitor known for its selective cell toxicity, and 5'-Aza-2'-deoxycytidine, an inhibitor DNMT, and one of the most promising and extensively used demethylating agents. The aim of this study was to select optimal concentrations or combination of concentrations of these inhibitors to achieve an epigenetic effect leading to AR gene restoration in androgen-independent prostate cancer cells.

The cancer DU145 cell line is androgen-independent and characterized by strong DNA methylation of the AR gene (17,20). This cell line is used as a model for simulating the conditions found in CRPC patients with changed genomic methylation patterns. We found that cancer line DU145 treated with the inhibitors used showed a substantial decrease in cell viability especially after 5 mM NaB administration and on 6-day exposure. NaB toxicity is a problem as to be effective as an inhibitor of histone deacetylases, NaB at the higher 5 mM concentration is needed. Comparing the required levels of cytotoxicity for NaB alone and NaB + Aza-dC co-treatments (0.5 and 5 µM Aza-dC with 5 mM NaB), it is clear that adding of the Aza-dC could have the same or better anti-proliferative effect with lower cell toxicity than 5 mM NaB alone in DU145 cells. The shorter 2-day treatments with used inhibitors had little effect, except for the NaB treatment that showed slight cytotoxicity in DU145 cells, and on the other hand growth stimulation in RWPE-1 cells. Similar results were reported by Paskova et al (21), where NaB (0.5, 1, 2.5 or 5 mM concentrations) had no toxic effect on normal RWPE-1 cells after 2-day treatment.

In accordance with the cell viability assay, the cell cycle distribution results showed similar effects of the used treatments, namely massive cell death upon 5 mM NaB treatment (administered alone and in co-treatment with Aza-dC) in DU145 cells compared to RWPE-1 cells. The finding that NaB inhibits cell viability and proliferation in DU145 cells and in certain other prostate cancer cell lines, in a time- and dose-dependent manner has been described in several studies (22-25). Pro-apoptotic activity induced by high doses of Aza-dC (~5 µM) has also been reported (26). In our study, Aza-dC alone also induced cell death (79% cells in sub-G1 cell population) in DU145 cells, however the co-treatments did not impair the toxic effects of NaB itself on the cell cycle. Although the cell death using a single inhibitor appears to be high, the combination of the two inhibitors might not have cumulative impact on prostate cancer cells. The appearance of the sub-G1 population of cells is apparently due to the DNA fragmentation of dead cells and could be a result of the pro-apoptotic NaB activity in cancer DU145 cells (24,25). A strong contrast of cell cycle distribution between both cell lines is given also by features of p-53 proficient RWPE-1 and p-53 deficient DU145 cells leading to p53-dependent activation and maintenance of G2/M cell cycle arrest and better survival in RWPE-1 cells compared to DU145 cells.

DU145 cells contain a methylated AR gene, thus the cell line has very low or undetectable AR gene expression (no Ct value in control DMSO treatment after 50 cycles was detected). The Ct value was established arbitrarily as Ct 50 at both time-points. Hence, the real re-expression of the AR gene could be considered higher than that calculated here for DU145 cells. In DU145, the most effective AR gene restoration was treatment with 0.5 µM Aza-dC + 5 mM NaB for both time-points, while the RWPE-1 cells did not show the same pattern and no significant difference between individual treatments was found after day 6. Although we noted an increased AR gene expression in RWPE-1, the level of mRNA was low compared to AR expression in DU145 cells. We observed a tendency to a decline in AR gene expression in a time-dependent manner apparent for both cancer and normal cell lines. The declining trend was also shown in histone H4 acetylation of the AR gene. As we changed the medium after 2 days, the half-life of inhibitors and recovery of remodulation enzymes, HDAC I and IIa that are targets of NaB (27), could result in lower AR gene re-expression and especially lower histone H4 acetylation.

We found significant re-acetylation of the histone H4 compared to control DMSO in cancer DU145 cells treated with the 0.5 µM Aza-dC + 1 mM NaB combination, while the same treatment induced significantly lower H4 acetylation levels in normal RWPE-1 cells. On the other hand, 0.5 µM Aza-dC + 5 mM NaB co-treatment had the same effect on both cell lines by increased enrichment of histone H4 acetylation while the 5 µM Aza-dC + 5 mM NaB had only moderate effect. This implies that low µM-concentration of Aza-dC together with mM-concentration of NaB could have either an additive/synergistic or antagonistic effect on chromatin remodeling.

The CRPC stadium harbourous heterogeneous features present with multiple alterations in the AR gene function. Besides mutations, copy number changes, deregulation of coregulators, and splice variants in AR gene, DNA methylation appears to be a minor modification contributing to AR gene dysfunction. However, based on a relatively recent study (12), the AR gene promoter methylation is likely related to prostate stem/progenitor traits and linked to enhanced castration resistance. Prostate tumor consists of a mixture of normal/benign, cancer and cancer stem cells with aberrant methylation profile. It is noteworthy that AR gene re-expression using epigenetic therapy leads to decreased proliferation and longer survival (13-15). AR signaling does not aggravate the development of the disease. It may act as a physiological regulator of AR downstream genes and sensitizes the cancer cells to other phases of therapy. For this reason, research efforts might be focused on keeping the active AR gene at low expression level to preserve the native AR signaling axis and to determine whether the restored AR gene could have regulation capability. Although
we showed a markedly increased AR mRNA expression and significant re-acetylation of histone H4 around AR gene promoter upon the co-treatment in DU145 cells, we assume that the corresponding AR gene regulation was not restored as the PSA level was not detected. Moreover, the high frequency of sub-G1 population of the dead cells was observed. To promote growth in androgen-independent CaP, the active AR should have selective and direct upregulation effect on M-phase cell cycle genes (28). Therefore, the cell cycle distribution results appear to be solely a consequence of NaB and Aza effects on cell death genes (24,25) and not by activation/restoration of AR. Modulation of the AR activity is mediated by the action of numerous coactivators and corepressors [histone modifiers, splice proteins, proteins of RNA metabolism and DNA repair, and cell cycle regulators (29)] and by phosphorylation of both AR and the mentioned co-regulators (30). It indicates further analysis is required aimed at mechanisms of action of epigenetic inhibitors and their potential role leading to completely active AR.

In conclusion, the impact of the combined treatment shows cancer cell reduction of their proliferative activity and changes in cell cycle distribution in comparison with normal cells with time-dependent effect. Further, the combined treatment both strongly increased histone H4 acetylation in the AR gene promoter and induced the AR gene re-expression in cancer cells in comparison with the normal cell line. Our results imply selective toxicity of used inhibitors with a suggestion that appropriately chosen inhibitor combination and concentration may have synergism/additive potential for therapeutic procedures in CRPC patients.

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References


