Introduction

The human prostate is unique in that it possesses the ability to accumulate high levels of intracellular zinc. Multiple studies have demonstrated that decreasing levels of intracellular zinc appear to be an important factor in the development and progression of prostate cancer (1,2). In fact, the inability to accumulate intracellular zinc by prostate cells often precedes the initial histopathological changes associated with prostate cancer. Cellular zinc handling becomes increasingly dysfunctional as prostate cancer progresses to castration-independent growth (3,4).

The zinc content of normal prostatic epithelium, benign prostatic hyperplastic tissue and cancerous prostate glands has been measured at 1018, 1142 and 146 μg/g of dry tissue, respectively (4). Recent mechanistic studies have revealed a strong association between the development of prostate cancer and downregulation of the zinc uptake transporters hZIP1 and hZIP3. The expression of hZIP1 and hZIP3 genes was markedly downregulated in adenocarcinomatous glands and in prostatic intraepithelial neoplastic foci when compared with adjacent normal peripheral zone glandular epithelium and benign hyperplastic glands (5–7). Moreover, we recently reported that overexpression of hZIP1 transporter has strong functional effect on the malignant potential of prostate cancer cells via inhibition of natural factor-kappaB-dependent pathways (8).

Although genetic alterations in prostate cancer have long been studied, the role of epigenetic changes during prostatic malignant transformation has recently garnered more attention. Epigenetic changes alter target gene expression without changing the cells DNA sequence. Inactivation of tumor suppressor genes by epigenetic changes is frequently observed in human cancers, particularly as a result of histone modification and/or DNA methylation.

Promoter methylation is one of the most common epigenetic mechanisms associated with altering gene expression. In a variety of tumors, CpG-rich regions, i.e. CpG islands, exhibit aberrant DNA hypomethylation resulting in abnormal transcriptional repression and gene inactivation (9). Specific to prostate cancer tumorigenesis, many of the inactivated genes in these CpG islands encode proteins that act as tumor suppressors, resulting in prostate cancer initiation, progression and, perhaps, an association with a more aggressive prostate cancer phenotype (10,11).

Recent studies have shown that the inhibition of DNA methyltransferase activity by 5-aza-2′-deoxycytidine (5-aza-CdR) prevented prostate cancer tumorigenesis in a mouse model (12). In the present report, we examine the effects of the demethylating agent 5-aza-CdR on the accumulation of intracellular zinc as well as the expression of zinc uptake transporters hZIP1 and hZIP3 in DU-145 and LNCaP prostate cancer cell lines. Recently, we reported that specificity protein 1 (SP1) and CAMP responsive element binding protein 1 are important transcription factors in the regulation of the hZIP1 zinc transporter gene (13). In the current study, we also demonstrate the importance of SP1 and activator protein (AP)-2 alpha proteins as transcription factors in the regulation of the hZIP3 zinc transporter in RWPE-1 cells. Furthermore, we were able to document the critical role of AP-2 alpha in regulating hZIP1 gene transcription in the RWPE-1 normal prostatic epithelial cell line.

In addition, we show that the epigenetic mechanisms of gene silencing caused by promoter hypomethylation in prostate cancer cells are indirectly involved in transcriptional downregulation of the zinc transporters hZIP1 and hZIP3. Since the SP1 and AP-2 alpha proteins play an important role in the transcriptional regulation of hZIP1 and hZIP3 genes, the loss of expression of any of these transcription factors would have a dramatic impact on the expression of the zinc uptake transporters. Specifically, we appear to demonstrate that hypomethylation of AP-2 alpha’s promoter region results in decreased expression of AP-2 alpha in DU-145 and LNCaP prostate cancer cells. This decreased expression results in a notable transcriptional downregulation of the hZIP1 and hZIP3 zinc transporters and, thus, reduced ability to accumulate intracellular zinc.

Materials and methods

Cell lines and culture conditions

The hormone-dependent LNCaP, hormone-independent DU-145 prostate cancer cell lines and the RWPE-1 normal prostate epithelial cell line were obtained from ATCC (Rockville, MD). Cells were cultured in RPMI 1640 (Bio-Whittaker, Walkersville, MD) medium supplemented with 10% fetal bovine serum (HyClone, Logan, UT), gentamycin (50 mg/l), sodium pyruvate (1 mM) and non-essential amino acids (0.1 mM). RWPE-1 cells were maintained in Keratinoocyte-Serum Free medium (Invitrogen, Carlsbad, CA) supplemented with 5 mg/ml of human recombinant epidermal growth factor and 0.05 mg/ml of bovine pituitary extract.

Antibodies and reagents

Antibodies to SP1 (sc17824X) and AP-2 alpha (sc-184X) were obtained from Santa Cruz Biotechnology (Santa Cruz, CA). Antibodies to GAPDH (#2118) were obtained from Cell Signaling Technology (Danvers, MA). 5-aza-CdR was purchased from Sigma–Aldrich Corporation (St Louis, MO).

Computer analysis of promoter regions

The analysis of 5′-flanking region of hZIP3 gene was performed using Gene2Promoter online programs (www.genomatix.de). Prediction of transcriptional factor-binding sites was carried out using the AliBaba 2.0,
Plasmid construct

The 5’-flanking region of the hZip1 gene has been recently cloned into the luciferase reporter vector (13). Promoter region (~154/+100 relative to transcription start) of hZip3 gene site was amplified by polymerase chain reaction (PCR) with 5’-ACTTGGGTACCGTTCGCGCCCAGGGCGTTGTTG-3’ forward and 5’-ATCTTGGTACCGTTCGCGCCCAGGGCGTTGTTG-3’ reverse primers containing restriction sites (underlined) using T7-Primer–genomic DNA as a template and cloned into XhoI/KpnI sites of the pcDNA3.1 basic reporter vector (Promega, Madison, WI). Site-directed mutagenesis of transcription factor–binding sites was performed by PCR with overlapped complementary primers containing the targeted binding site (Supplementary Table 1 is available at Carcinogenesis Online). All mutations in reporter constructs were confirmed by sequencing using GL2 primer (Promega) in the Fox Chase Cancer Center Automated DNA Sequencing Facility.

Electrophoretic mobility shift assay and Gel-Supershift assays

Electrophoretic mobility shift assay and Gel-Supershift assays were performed as described previously (13) using a set of double-stranded DNA oligos (Supplementary Table 2 is available at Carcinogenesis Online) and specific antibodies.

Assessment of cell growth

Cells were plated in 96-well plates (5 × 103 to 10 × 103 cells per well) and incubated for 48 h in complete cell growth medium. The metabolic capacity of cells was then evaluated using the CellTiter-Blue Cell Viability Assay (Promega) according to the manufacturer instructions.

Reverse transcription and quantitative PCR

Total RNA was isolated from cells using the Mini RNA isolation-II Kit (Zymo Research Corporation, Orange, CA). Total RNA was treated with RNase-free DNase set (Invitrogen, Gaithersburg, MD) and 50 ng of random hexamer primers. Then, complementary DNA samples were diluted and 5 μl of diluted complementary DNA were amplified by TaqMan quantitative polymerase chain reaction (qPCR) using hZip1- and hZip3-specific primers and probes (IDT DNA Technologies, Coralville, IA). GAPDH Mini qPCR assay (IDT DNA Technologies) was used as an internal amplification control. Specific primers and probes are indicated in Supplementary Table 2 is available at Carcinogenesis Online.

Chromatin immunoprecipitation

Chromatin immunoprecipitation (ChIP) experiments were performed using the EZ-Chip assay kit (Upstate, Charlotteville, VA). Briefly, 1 × 106 cells were cross-linked with 1% formaldehyde and washed twice with ice-cold phosphate-buffered saline containing a cocktail of protease inhibitors (Roche, Germany). Cells were resuspended in 500–1000 μl of sodium dodecyl sulfate lysis buffer on ice and then sonicated with four sets of 10 s pulses by a Kontes Ultrasonic Cell Disrupter to an average DNA size of 200–400 bp. Protein concentrations of chromatin samples were measured using BCA Protein Assay (ThermoScientific, Rockford, IL) and 300 μg of chromatin protein were used in the immunoprecipitation step. The chromatin was precleared with protein A/G-agarose beads (Santa Cruz Biotechnology, Santa Cruz, CA) for 1 h followed by an overnight incubation at –4°C with 2 μg of specific antibodies. Normalized anti-mouse IgG (Santa Cruz Biotechnology) antibodies were used to control immunoprecipitations. Chromatin–antibody complexes were collected by reincubation with protein A/G PLUS-Agarose beads for 1 h. Chromatin was eluted from the beads and cross-links were reversed by incubation at 65°C for 4 h. ChIP DNA was recovered and purified using ChIP DNA Clean and Concentrator Kit (Zymo Research Corporation).

About 2% of input DNA and 2–8% of eluted ChIP DNA were subjected to TaqMan qPCR with specific primers and probes (Supplementary Table 4 is available at Carcinogenesis Online). Samples were run in triplicates for both input DNA and ChIP-DNA in a 10 μl reaction using TaqMan Universal Master Mix according to the manufacturer’s instructions (Applied Biosystems). Reactions were carried out in an Applied Biosystems 7500 Real-Time PCR System.

Relative quantification of the ChIP-DNA was performed with the comparative Ct method. The ΔCt values were computed by subtracting the average of normalized against input Ct values of GAPDH ChIP-DNA samples from the average of normalized against input Ct values of hZip1 or hZip3 ChIP-DNA samples recovered with anti-AP-2alpha antibodies. In order to determine the fold change of target ChIP-DNA treated with 5-aza-CdR cells over target ChIP-DNA in untreated cells, the ΔΔCt was calculated by subtracting the ΔCt values of treated cells from the ΔCt values of untreated cells. The relative amount of target ChIP-DNA was finally calculated by the formula 2−ΔΔCt.

Tissue samples

Tissue specimens were obtained from the Department of Urology, Cancer Research Center RAMS, Russia and analyzed at the Institute of Carcinogenesis, Cancer Research Center, Moscow, Russia. All of the prostate tumors had been histopathologically confirmed as stage pT2cN0MO Gleason score 6 prostate cancer. All patients had not received any hormonal or radiation therapy prior to radical prostatectomy. Representative samples of confirmed prostate cancer as well as adjacent normal prostate tissue were taken from each prostate specimen and were freshly frozen at the time of surgery. Genomic DNA samples were isolated from the frozen tissues using STR Genomic DNA3M-Tissue MiniPrep (Zymo Research Corporation) according to manufacturer’s instructions.

Results

Treatment with 5-aza-CdR increases zinc uptake in prostate cancer cells

Since prostate cancer development and progression is characterized by a sequentially increasing dysregulation of zinc metabolism, we chose to use in our experiments the hormone-dependent LNCaP and the hormone refractory DU-145 prostate cancer cell lines as well as the normal prostate epithelial cell line RWPE-1. Firstly, we compared the ability of the normal prostate epithelial cell line RWPE-1 and the prostate cancer cell lines DU-145 and LNCaP to accumulate zinc by staining the cells with the fluorescent zinc indicator FluoZin-3. As demonstrated in Figure 1A, intracellular zinc levels were significantly decreased in the prostate cancer cell lines DU-145 and LNCaP when compared with RWPE-1 cells.

To assess whether epigenetic factors participate in the regulation of intracellular zinc uptake in prostate cancer cells, DU-145 and LNCaP cells were treated with the inhibitor of DNA methyltransferases 5-aza-CdR. Treatment with 5-aza-CdR resulted in significantly increased intracellular zinc accumulation in DU-145 and LNCaP cells as shown in Figure 1B. The possible toxic effect of 5-aza-CdR on cell line survival was evaluated by propidium iodide staining following 48 h of incubation of a 5 μM concentration of 5-aza-CdR with LNCaP, DU-145 and RWPE-1 cells; respectively. PI staining did not reveal a significant increase in the induction of apoptosis between the cells treated with the demethylating and those not treated with the demethylating agent (data not shown).

Upregulation of hZip1 and hZip3 expression by 5-aza-CdR

Next, we sought to examine effects of 5-aza-CdR treatment on the transcriptional regulation of the hZip1 and hZip3 genes. Using RT–
LNCaP cells that were treated with 5-aza-CdR. Cells were treated with 5-aza-CdR for 48 h and then cell media was changed and incubation was continued for another 24 h. Cells were harvested and subjected to qRT–PCR analysis. The qPCR, we found that hZip1 and hZip3 messenger RNA levels were increased in both DU-145 and LNCaP cells treated with 5-aza-CdR (Figure 1C). These results imply that treatment with 5-aza-CdR engenders increased intracellular zinc uptake by upregulation of the main zinc transporters.

Given that treatment with 5-aza-CdR increases zinc uptake in prostate cancer cells and that this increase coincides with upregulated expression zinc transporters, we proposed that DNA methylation may be partially responsible for the decreased zinc accumulation in prostate cancer cells. Thus, we analyzed the methylation status of promoter regions of hZip1 and hZip3 genes. Using bisulfite sequencing, we found hZip1 and hZip3 promoter regions to be hypomethylated in both prostate cancer DU-145 and LNCaP cell lines (data not shown). Based on these findings, we hypothesized that 5-aza-CdR treatment could have a secondary effect on hZip1 and hZip3 genes by activation of silenced transcription factors involved in regulation of hZip1 and hZip3 expression.

Transcriptional regulation of hZip1 and hZip3 promoters: role of SP1 and AP-2alpha

Since some of 5-aza-CdR’s effects on hZip1 and hZip3 expression appeared to involve the potential derepression of epigenetically silenced transcription factors, we investigated mechanisms of transcriptional regulation of hZip1 and hZip3 genes. Although the role of SP-1 and CAMP responsive element binding protein-1 proteins in the regulation of the hZip1 promoter was recently shown (13), the impact of the AP-2-binding site at –58/–67 position (relative to transcription start site) of the hZip1 promoter region still remains unclear. Since the AP-2 consensus sequence has been found, the pGL3-hZip1-luc (–366/+315) reporter vector was subjected to site directed mutagenesis. The Luciferase Reporter Assay revealed the high significance of the AP-2-binding site for transcriptional regulation of the hZip1 promoter. As demonstrated in Figure 2A, mutation of the AP-2alpha-binding site dramatically impaired transcriptional activity of the hZip1 (–366/+315) region.

In order to ascertain which transcriptional factors play a significant role in the regulation of hZip3 promoter activity, we performed a computer analysis of this promoter region using the AliBaba 2.0, TRANSFAC, Match-1.0, and MathInspector™ computer programs. Since multiple SP-1 and AP-2-like consensus sequences were predicted in the hZip3 promoter region, a pGL3 luciferase reporter vector containing a hZip3 (–154/+100) promoter region was constructed. Transcriptional activity of the hZip3 promoter region was then measured after single and combined mutations of the transcription factor binding sites were created. The results of native and mutated hZip3 promoter activity demonstrated the important role of SP-1 and AP-2alpha in transcriptional regulation of the hZip3 promoter region as illustrated on Figure 2C.

Analysis of SP-1 and AP-2alpha binding to hZip1 and hZip3 promoters

Next, we performed Gel-Supershift assays to evaluate the ability of SP-1 and AP-2alpha proteins to bind hZip1 and hZip3 promoters. For the assays, several double-stranded DNA oligos and their mutated analogs spanning the SP-1 or AP-2alpha-binding sites within hZip1 and hZip3 promoters were generated. The SP-1 protein was able to bind the hZip3 promoter region in the –120/–114 (relative to transcription start site) position (Figure 2D). As demonstrated on Figure 2E, using the following set of probes spanning the appropriate AP-2-binding sites within hZip1 and hZip3 promoters, we were able to demonstrate the ability of the AP-2alpha transcription factor to bind hZip1 (–67/–58), hZip3 (–147/–138), hZip3 (–25/–15) and hZip3 (+83/+92) promoter regions.

AP-2alpha expression is downregulated in DU-145 and LNCaP cells due to promoter hypermethylation

Since SP-1 and AP-2alpha have key roles in hZip1 and hZip3 promoter activities, we compared the levels of expression of these proteins in DU-145 and LNCaP prostate cancer cell lines as well as in the normal prostate epithelial cell line RWPE-1. SP1 is a ubiquitously expressed protein and western blot assay revealed no difference in the level of SP1 protein expression in prostate cancer and normal cells. In
contrast, the level of the AP-2alpha protein was found to be decreased in both the DU-145 and LNCaP cell lines as compared with the RWPE-1 cell line (Figure 3A).

It has previously been shown that downregulation of the AP-2alpha transcription factor is caused by hypermethylation of its promoter region in breast cancer cells (15). Additionally, it was reported that the AP-2alpha protein expression is lost in prostate cancer cells (16). However, the methylation status of the AP-2alpha promoter region in the prostate cancer cell lines DU-145 and LNCaP to date had not been characterized. Thus, we analyzed the methylation of the AP-2alpha promoter region relative to the +1 position of AP-2alpha transcript (accession # NM_003220) by bisulfite sequencing. As demonstrated in Figure 3D, the AP-2alpha promoter region was found to be unmethylated in the normal prostate epithelial cell line RWPE-1. In contrast, we observed high methylation levels of this region in both the DU-145 and LNCaP prostate cancer cell lines.

Treatment with 5-aza-CdR results in AP-2alpha promoter demethylation, upregulation of its expression and binding to hZip1 and hZip3 promoters

Based on the data from our preceding experiments, we examined the effect of 5-aza-CdR treatment on the methylation level of the AP-2alpha promoter region in DU-145 and LNCaP cells. As demonstrated in Figure 3E, 5-aza-CdR treatment resulted in reduced methylation of the AP-2alpha promoter regions in both DU-145 and LNCaP cells. Furthermore, demethylation of the AP-2alpha promoter coincided with an increased expression of AP-2alpha protein levels in those cell lines (Figure 3B), whereas SP-1 protein levels remained without any change.
changes. Next, ChIP–qPCR analysis was performed to examine the effect of 5-aza-CdR treatment on binding of AP-2alpha with hZip1 and hZip3 promoters in vivo in DU-145 and LNCaP cells. As we theorized, increased binding of the AP-2alpha transcription factor with hZip1 and hZip3 promoter regions occurred in both DU-145 and LNCaP cells after 5-aza-CdR treatment (Figure 4).

The impact of AP-2alpha transcription factor in regulation of hZip1 and hZip3 genes expression, cell growth and zinc accumulation in prostate cancer cells

To investigate if restored expression of AP-2alpha would result in upregulated zinc transporters expression and increased zinc accumulation, we overexpressed this transcription factor in DU-145 and LNCaP prostate cancer cell lines. We performed the transient transfection of DU-145 and LNCaP cells with a pcDNA3.1 vector containing AP-2alpha open-reading frame. The relative messenger RNA levels of hZip1 and hZip3 zinc transporters were measured by qRT–PCR in DU-145 and LNCaP cells transfected with pcDNA3.1-AP-2alpha vector and control pcDNA3.1 vector. Twenty-four hours after transfection, the population of transfected cells was separated from non-transfected cells using fluorescence-activated cell sorting and incubated for additional 24 h. Cells were then harvested and subjected to RT–qPCR analysis and western blotting. As illustrated in Figure 5A, we observed increased hZip1 and hZip3 messenger RNA levels in DU-145 and LNCaP cells transfected with AP-2alpha expression construct. We then sought to examine the functional significance of increased zinc transporters expression. Consistent with the ability of
ChIP-DNA samples recovered with anti-AP-2alpha antibodies. DU-145 and hZip3 promoters ChIP-DNA were normalized to the GAPDH promoter triplicates. The representative data of one of the three ChIP–qPCR harvested and ChIP–qPCR was performed. The qPCR reactions were ran in additional 24 h incubation in cell medium alone. After incubation, cells were harvested and ChIP-qPCR was performed. The qPCR reactions were run in triplicates. The representative data of one of the three ChIP-qPCR experiments are presented. Error bars represent ±SD of triplicates.

**Fig. 4.** ChIP-qPCR assay representing association of AP-2alpha at hZip1 and hZip3 promoter regions in vivo. Target DU-145 (A) or LNCaP (B) ChIP-DNA increased after cells were treated with 5-aza-CdR. Bars represent fold change of $2^{-\Delta\Delta Ct}$ of target ChIP-DNA in treated cells over $2^{-\Delta\Delta Ct}$ of target ChIP-DNA in untreated cells. Normalized against input Ct values of hZip1 and hZip3 promoters ChIP-DNA were normalized to the GAPDH promoter ChIP-DNA samples recovered with anti-AP-2alpha antibodies. DU-145 and LNCaP cells were preincubated with 5 μM of 5′-aza-CdR for 48 h with an additional 24 h incubation in cell medium alone. After incubation, cells were harvested and ChIP-qPCR was performed. The qPCR reactions were run in triplicates. The representative data of one of the three ChIP-qPCR experiments are presented. Error bars represent ±SD of triplicates.

hZip1 and hZip3 to enhance intracellular zinc uptake (1,17), the accumulation of labile zinc was notably increased in cells transfected with the AP-2alpha expression vector (Figure 5C) cultured in the presence of physiologically relevant concentrations of zinc. Also, we evaluated the effect of AP-2alpha expression in DU-145 and LNCaP prostate cancer cells on cellular growth using the CellTiter-Blue Assay. As displayed in Figure 5D, we observed a 25 and 33.1% inhibition in prostate cancer cell growth for DU-145 cells and LNCaP cells, respectively.

**Analysis of AP-2alpha promoter methylation in clinical specimens**

Results of several previous studies have shown that the loss of AP-2alpha expression occurs early in the development of prostate adenocarcinoma (16) as well as during breast cancer progression (15). Our results demonstrate that AP-2alpha promoter region hypermethylation in DU-145 and LNCaP prostate cancer cell lines coincides with decreased AP-2alpha expression. Based on these results, we wanted to examine whether AP-2alpha promoter hypermethylation occurs in tissue samples from prostate glands surgically removed for localized prostate cancer. Samples of genomic DNA were isolated from frozen prostate tumors and adjacent normal tissues of five patients. After bisulfite conversion, DNA samples were amplified with specific primers and then cloned into pGEM-T Easy vector. Eight individual clones of each PCR product representing individual alleles of the AP-2alpha 5′-flanking region were sequenced. Figure 6 demonstrates that methylation of the AP-2alpha 5′-flanking region in all prostate cancer tissues was found to be markedly increased in comparison with the adjacent normal prostatic tissue except for tumor #2 where the methylation level of the AP-2alpha 5′-flanking region remained hypermethylated compared with the adjacent normal tissue.

**Discussion**

The process whereby a normal prostatic epithelial cell transforms into a cancerous cell involves the loss of the ability to accumulate intracellular zinc. This transformation remains one of the sine qua non cellular changes during prostatic carcinogenesis. Recent studies reveal a strong association between the development of prostate cancer and the downregulation of zinc uptake transporters hZip1 and hZip3 (6,18). Therefore, in the present study, we examined the epigenetic mechanisms that might be responsible for the downregulation of hZip1 and hZip3 genes in DU-145 and LNCaP prostate cancer cell lines.

DNA hypermethylation is one of the best-characterized epigenetic changes noted in neoplastic development and can be associated with gene silencing (10,11,19). Recent studies have shown that 5-aza-CdR appears to interrupt tumorigenesis by inhibition of DNA methyltransferase. In the present study, the expression of hZip1 and hZip3 zinc uptake transporters as well as the ability to accumulate intracellular zinc was notably increased after 5-aza-CdR treatment. Interestingly, an analysis of the promoter regions of the hZip1 and hZip3 genes did not exhibit hypermethylation in either cell line. Thus, we do not regard hypermethylation of the hZip1 and hZip3 promoters as the primary mechanism responsible for downregulation of these genes in DU-145 and LNCaP prostate cancer cells. It is probably that the epigenetic regulation of hZip1 and hZip3 gene expression is a secondary event. Indeed, it is known mechanism while downregulation of transcriptional regulators caused by their promoters hypermethylation has affected expression of their target genes (20,21).

Recently, we were able to characterize the transcriptional regulation of the hZip1 promoter (13) in contradistinction to the hZip3 promoter, whose regulatory control remained unclear. In our present work, we determined the essential role of SP1 and AP-2alpha proteins in transcriptional regulation of hZip3 expression. Additionally, we characterized the AP-2-like binding site in the hZip1 promoter, which has been shown to have an important regulatory role on hZip1 expression. Since SP1 and AP-2alpha proteins were found to be critical for transcriptional regulation of both hZip1 and hZip3 genes, we compared the expression of these transcriptional factors in DU-145 and LNCaP prostate cancer cell lines as well as in the normal prostatic epithelial cell line RWPE-1. The level of SP1 expression remained unchanged; therefore, we focused our investigations on the AP-2alpha transcription factor, which has been shown to be downregulated in both DU-145 and LNCaP cells in comparison with RWPE-1 cells.

AP-2alpha is a well-characterized transcription factor that is required for growth, normal morphogenesis and apoptosis in mammalian cells (22–24). It binds to the GC-rich promoter regions of various genes (25) where it has been shown to be involved in both transcriptional activation of some genes (26–28) and repression of other genes (29–31). Furthermore, the AP-2alpha has been associated with cellular transformation and anchorage-independent growth (32) as well as being under the stimulatory control of the ERBB2 oncogene (33).

Conversely, different investigators have demonstrated an antitumor activity of the AP-2alpha protein (29,34). For example, the loss of AP-2alpha has been strongly associated with the aggressiveness and progression of numerous types of cancer, such as gliomas, melanoma, breast, prostate and colorectal cancers (15,35–37). In particular, the AP-2alpha protein has been associated with luminal differentiation of prostatic epithelium, and its expression is lost in high-grade prostatic intraepithelial neoplastic and prostate cancer (16,38). It was also
reported that the loss of AP-2alpha affects the normal expression profiles of many genes involved in carcinogenic pathways (29,39).

The downregulation of AP-2alpha protein expression caused by hypermethylation of its promoter region has been reported in hematological as well as in breast cancer and other solid cancers (15,40). Little is presently known regarding the methylation patterns of AP-2alpha promoter in prostate cancer cell lines. Thus, based on our finding of AP-2alpha downregulation in prostate cancer cell lines, we hypothesized that the mechanism of AP-2alpha downregulation in prostate cancer cells involves hypermethylation of its promoter region. Using bisulfite sequencing, we were able to confirm this hypothesis by finding high methylation levels of the AP-2alpha promoter region in DU-145 and LNCaP prostate cancer cells. In contrast, the same promoter region in RWPE-1 normal prostate epithelial cells was unmethylated. Next, we examined the effects of the demethylating agent 5-aza-CdR on methylation of AP-2alpha promoter regions in the DU-145 and LNCaP cell lines. Bisulfite sequencing revealed a reduction in methylation levels in both DU-145 and LNCaP cells, which coincided with an elevation of AP-2alpha protein levels in those cells.

Our data coupled with previous published reports suggest that the loss of the AP-2alpha protein is caused by its promoter hypermethylation during breast cancer progression (15) as well as being an early event in prostate cancer development (16). To test this supposition, we examined methylation patterns of the AP-2alpha promoter region from patients’ tissues who had Gleason score 6 prostate cancer. Gleason score 6 prostate cancer is considered to be typical for early well-differentiated disease. Patients with Gleason score 6 disease generally have a very favorable prognosis and a low risk of recurrence after therapy (41). Bisulfite sequencing of the AP-2alpha promoter region in five human tissue samples with prostate cancer revealed that 80% of these patients had a high methylation level at the AP-2alpha promoter region. Despite the fact that the level of AP-2alpha promoter methylation was lower in human prostate cancer tissue than compared with the prostate cancer cell lines DU-145 and LNCaP, the degree of methylation in human prostate cancer tissues was still significantly higher than compared with adjacent normal prostate tissue as well as in normal prostate epithelial RWPE-1 cells. These data suggest that the methylation of AP-2alpha promoter region in prostate cancer cells is an early event in neoplastic development.

In order to test our theory that the AP-2alpha protein plays a significant role in the transcriptional regulation of both hZip1 and hZip3 genes, we re-established an altered expression of AP-2alpha in the DU-145 and LNCaP prostate cancer cell lines. We transfected DU-145 and LNCaP prostate cancer cells with an AP-2alpha expression vector that resulted in the upregulation of the hZip1 and hZip3 genes. As a result of this upregulation, these transfected cells demonstrated a significantly higher zinc uptake. The re-establishment of normal intracellular zinc levels is a particularly attractive target for therapeutic manipulation given the importance of dysregulation of zinc metabolism in the development and progression of prostate cancer. Recent studies have demonstrated that the activation of Ras responsive element binding protein-1 resulted in the activation of dysregulation of zinc metabolism in the development and progression of prostate cancer. Recent studies have demonstrated that the activation of Ras responsive element binding protein-1 resulted in the upregulation of the hZip1 zinc transporter downregulation in prostate cancer cells (42). In addition to this work, the present study, to our knowledge, represents

regulation of intracellular zinc uptake, transfected DU-145 and LNCaP cells were incubated with 2 μM of zinc for 1 h, washed twice in phosphate-buffered saline and incubated with the cell-permeable fluorescent zinc indicator FluoZin-3 (5 μM) for additional 15 min. Fluorescence was examined by flow cytometry. Data is representative of three independent experiments. *P < 0.01; **P < 0.05 compared with pcDNA3.1-AP-2alpha transfected cells. (D) The effect of AP-2alpha expression on cell growth was examined using the CellTiter-Blue Assay. DU-145 and LNCaP prostate cancer cells were transfected as described above, plated in a 96-well plate (5 × 10³ to 10 × 10³ cells per well) in triplicates and incubated for 48 h in complete cell growth medium. After the 48 h incubation period ended, the CellTiter-Blue Assay was performed. Error bars represent ±SD of triplicate fluorescence means.

![Fig. 5](image-url)

Fig. 5. The DU-145 and LNCaP cell lines were transiently transfected with the pcDNA3.1-AP-2alpha expression vector and pcDNA3.1 empty vector. Twenty-four hours after transfection, cells were subjected to fluorescence-activated cell sorting. After another 24 h of incubation in cell medium, cells were harvested and RT-qPCR assay (A), western blot assay (B) were performed. (C) In order to assess whether transfection of DU-145 and LNCaP cells with the pcDNA-AP-2alpha expression vector has an effect on the
Supplementary Tables 1–4 can be found at http://carcin.oxfordjournals.org/

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