A Small Molecule Polyamine Oxidase Inhibitor Blocks Androgen-Induced Oxidative Stress and Delays Prostate Cancer Progression in the Transgenic Adenocarcinoma of the Mouse Prostate Model

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Abstract

High levels of reactive oxygen species (ROS) present in human prostate epithelia are an important etiologic factor in prostate cancer (CaP) occurrence, recurrence, and progression. Androgen induces ROS production in the prostate by a yet unknown mechanism. Here, to the best of our knowledge, we report for the first time that androgen induces an overexpression of spermidine/spermine N1-acetyltransferase, the rate-limiting enzyme in the polyamine oxidation pathway. As prostatic epithelia produce a large excess of polyamines, the androgen-induced polyamine oxidation that produces H2O2 could be a major reason for the high ROS levels in the prostate epithelia. A small molecule polyamine oxidase inhibitor N,N′-butanedienyl butanediamine (MDL 72,527 or CPC-200) effectively blocks androgen-induced ROS production in human CaP cells, as well as significantly delays CaP progression and death in animals developing spontaneous CaP. These data show that polyamine oxidation is not only a major pathway for ROS production in prostate, but inhibiting this pathway also successfully delays CaP progression. [Cancer Res 2009;69(19):7689–95]

Introduction

Advanced hormone refractory metastatic prostate cancer (CaP) is a major cause of cancer deaths among U.S. men. Most CaP patients at the time of initial diagnosis have androgen-dependent tumors that regress quickly after radical prostatectomy or radiation therapy. Unfortunately, in ~15% of the patients, the cancer recurs within a few years as an advanced hormone refractory and often-metastatic disease. Most commonly used cancer chemotherapeutic agents have little effect against advanced, metastatic CaP. Therefore, development of agents to prevent CaP occurrence, recurrence, and progression to the advanced stage is warranted. Reactive oxygen species (ROS) such as hydrogen peroxide, superoxide, hydroxyl-free radical, and nitric oxide levels are relatively higher in prostate epithelial cells than are in most other tissues (1, 2). Direct evidence linking ROS with an increase in tumor development in the prostate has been established (3–5). The ability of ROS to alter growth or apoptosis-related genes either by direct mutagenic effects on DNA or by alterations in gene expression and cellular signaling suggest a potential role for ROS in both initiation and progression of CaP (1–16). Oberley and colleagues (16) performed immunohistochemistry to analyze human malignant and normal prostate tissues in archival paraffin blocks. They reported that oxidative stress-induced enzymes and oxidative damage to DNA bases are relatively more abundant in malignant CaP compared with that in normal human prostate tissues. Ho and her colleagues (17, 18) used similar methods to confirm that the ROS-induced damage in spontaneously formed prostate neoplasm of transgenic adenocarcinoma of mouse prostate (TRAMP) animals are relatively more than that in normal prostatic lumen of the same animal. It has also been shown that ROS play a key role in the androgen-independent growth of androgen-dependent CaP cells (19). Therefore, understanding the biochemical pathway that modulates cellular ROS levels could yield a novel and effective therapeutic strategy to delay or even prevent CaP occurrence, recurrence, and progression. Androgen induces oxidative stress by producing ROS in normal and malignant prostatic epithelial cells (5, 10–15). The results initially published from our laboratory (10–14) have been independently confirmed by other laboratories (15, 18). Ho and colleagues (18) conclusively showed that androgen induces ROS production in rat prostatic tissues. The biochemical mechanism of androgen-induced ROS production in the prostate, however, has not yet been reported.

Our DNA microarray data from androgen-treated and untreated LNCaP human CaP cells suggest that androgen induces over-expression of spermidine/spermine N1-acetyltransferase (SSAT) mRNA. SSAT is the first enzyme in polyamine catabolic pathway. Polyamines are essential components of the seminal fluid. Large excess of polyamines are produced and secreted by prostatic epithelial cells and polyamine catabolism produces the ROS H2O2 (20–22). Thus, an induction of a rate-limiting enzyme of the polyamine catabolic pathway may be a key factor in producing a high level of ROS in the prostatic tissue. Here, we report data from quantitative reverse transcription–PCR (qRT-PCR): lack of androgen-induced ROS production in cells

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transfected with small interfering RNA (siRNA) against SSAT, SSAT enzyme activity, and cellular polyamine levels in LNCaP cells. These data confirm that androgen induces both SSAT expression and enzyme activity in androgen-treated LNCaP human CaP cells. A small molecule inhibitor of N1-acetyl polyamine oxidase (APAO) N,N’-butanedienyl butanediamine (MDL 72,527 or CPC-200; refs. 23, 24) completely blocks androgen-induced ROS production in LNCaP cells, as well as in the prostatic lumen of the TRAMP animals. CPC-200 treatment also inhibited tumor growth and significantly increased the life expectancy of TRAMP animals. These data clearly show that polyamine oxidation is the major biochemical pathway for generating oxidative stress in the prostatic epithelial cells. Blocking polyamine oxidation is a valid strategy for lowering oxidative stress in the prostate and prevents CaP progression, showing a direct link between prostatic ROS and CaP progression.

Materials and Methods

Materials

CPC-200 has been synthesized by Prof. Patrick Woster following a previously published procedure (23). The LNCaP human prostate carcinoma cell line was purchased from the American Type Culture Collection. All enzymes and assay kits were purchased from manufacturers described in the Methods (see below).

Methods

**Tissue culture.** Cells were maintained in DMEM supplemented with 5% fetal bovine serum, nonessential amino acids, and 1% streptomycin-penicillin in a humidified 95% air/5% CO2 atmosphere. Cells were harvested by treatment for 3 to 5 min with STV (sulfate A, 0.05% trypsin, 0.02% EDTA) at 37°C following a previously published procedure (12).

**Androgen deprivation.** Cells collected for experiments were counted and cultured in medium containing 4% charcoal-stripped serum plus 1% nonstripped serum (F1/C4) for 48 h. This combination of stripped and nonstripped serum was previously shown to sufficiently deplete androgen content while limiting adverse growth effects not related to hormone depletion that occur with the use of 5% stripped serum (12). Concurrently in each experiment, cells were seeded in 96-well tissue culture plates at a density of 2,500 cells per well in 100 μL medium for the measurement of ROS as an indicator of redox status. DNA levels were measured as an indicator of growth (see below). LNCaP cells were seeded in F1/C4 at a density of 1 × 106 cells per 10-cm tissue culture plate for protein estimation for Western blot analysis (see below). Protein estimation was carried out following standard procedure previously published from our laboratory (10).

**qRT-PCR.** Total RNA from LNCaP cultures was isolated using the RNeasy kit (Qiagen) according to the manufacturer-supplied protocol. For qRT-PCR analysis, cDNA was produced from total RNA using Superscript (Invitro-gene) according to the manufacturer’s instructions, with thermal cycling parameters of j 95°C for 10 min followed by 40 cycles of a 96°C denaturation for 15 s and a 60°C annealing/extension for 1 min using an iCycler (Bio-Rad), following a published procedure (10). To control for variability in efficiency of cDNA synthesis between samples, CDNA levels of the genes under investigation were normalized to the cDNA levels of glyceraldehyde-3-phosphate dehydrogenase (GAPDH). PCR primer sequences were designed using Oligo 5.0 software (National Biosciences) and synthesized at the University of Wisconsin Biotechnology Center. The PCR primer set sequences were as follows: GAPDH forward primer, AAA TTC CAT GGC ACC GTC AA; and reverse primer, TCT CCG TCC TGG AAG ATG GT; SSAT forward primer, CGA GCT CGA GAG GGG CCT GGT CCG CAA A and reverse primer, GTT CGA ATT CTA AAG CTT TGG AAT GGG TGC TCA. siRNA transfection and isolation of stable LNCaP clone sSSAT.

**Construction of psF1-H1-siSSAT vector.** Stable expression of siRNA for SSAT was used to suppress SSAT expression in LNCaP cells. Oligonucleo-

tides for silencing SSAT were designed based on the published sequence (25). The oligonucleotides were synthesized by Invitrogen. The annealed oligonucleotides were inserted into psF1 vector (SBI, System Biosciences).

**Transduction of LNCaP cells by pPACKaged psF1-H1-siSSAT or psF1-H1-siLuc.** LNCaP cells stably expressing psF1-H1-siSSAT vector were established using lentiviral system from SBI, following the manufacturer’s instructions. Briefly, the day before transfection, 5 × 105 293NT cells were seeded in a 10-cm plate in complete medium without antibiotics. The following day, 2 μg expression vector psF1-H1 carrying siSSAT or vector control (psF1-H1-siLuc) were separately mixed with 10 μg pPACK (Packaging Plasmid Mix; SBI) and 30 μL of Lipofectamine (Invitrogen) in medium without serum and antibiotics. The mixture was added to 293NT cells in 2% serum without antibiotics. After overnight incubation at 37°C in a 5% CO2 incubator, the medium was replaced with fresh 2% serum containing antibiotics, and the incubation was continued for another 48 h at 37°C in a 5% CO2 incubator. The supernatant was harvested by centrifugation at 5,000 rpm for 5 min and used for transducing 1 × 104 LNCaP cells, which were plated in polylysine plate (BD Bioscience) the day before transduction. Puromycin (Sigma; 1 μg/mL) in complete medium was used for selection of stably transduced cells for psF1-H1-si SSAT and vector control. The silencing of SSAT in these cells was verified by qRT-PCR (see below).

**Reduced 2’-7’-dichlorofluorescein diacetate oxidation assay.** The 96-well culture plates were assayed for estimation of ROS levels in intact cells using the fluorescent dye 2’,7’-dichlorofluorescein diacetate (DCF; Molecular Probes, Inc.) following a published procedure (12). In brief, cell cultures were washed with 200 μL Kreb's Ringer buffer warmed to 37°C, incubated as usual at 37°C in 100 μL Kreb's Ringer buffer containing 10 μg/mL (final concentration) DCF dye for 45 min. Each 96-well culture plate was scanned on a CytoFluor 2350 plate scanner (Applied Biosystems) using the 485/530 nm filter excitation and emission set and then frozen at −70°C for the subsequent analysis of DNA content.

**DNA assay.** For DNA analysis, each culture plate froze to −70°C was thawed/equilibrated to room temperature in the dark. Hoechst dye was then added to each well in 200 μL of high salt TNE buffer [10 mmol/L Tris, 1 mmol/L EDTA, 2 mol/L NaCl (pH 7.4)] following a published procedure (12). After further incubation at room temperature for over 2 h under protection from light, culture plates were scanned on a CytoFluor 2350 scanner using the 360/460 nm filter excitation and emission set. The DCF fluorescence units were normalized to the Hoechst-DNA fluorescence units for each well and used as a measure of the level of ROS being generated. The DNA fluorescence units were also used as a measure of cell growth.

**SSAT assay.** SSAT assay was performed following essentially the same procedure published elsewhere (26). Cells in monolayer were washed twice with PBS and resuspended in 5 mmol/L HEPES buffer (pH 7.2) containing 1 mmol/L DTT. Cells were lysed by three 30-s pulses of sonication. Cell lysates were centrifuged, and cytoplasts were collected and stored at −80°C. On the day of the assay, the cytoplasts were incubated with 150 pmol of spermidine and 500 pmol of 15C-acetyl CoA (GE Healthcare/Amersham) in 25 μL HEPES buffer for 30 min. The reaction was stopped by cooling in ice and by an addition of chilled 10 μL of hydroxyamine hydrochloride and then heating in a boiling water bath. After centrifugation, the supernatant was spotted on a phosphocellulose filter, washed, and counted. Cytosolic protein contents were determined by Bradford method, and the results are expressed as picomoles of acetyl spermidine synthesized per minute per milligram protein.

**Polyamine analysis.** A known number of cells (>1 × 105) were taken from harvested samples and centrifuged at 800 × g at 4°C for 5 min. The cells were washed twice with chilled Dulbecco’s isotonic phosphate buffer (pH 7.4) by centrifugation at 1,000 rpm at 4°C and resuspended in the same buffer. After the final centrifugation, the supernatant was decanted and 250 μL of 8% sulfosalicylic acid were added to the cell pellet. The cells were sonicated, and the mixture was kept at 4°C for at least 1 h. After further centrifugation at 8,000 × g for 5 min, the supernatant was removed for analysis following a published high-performance liquid chromatography (HPLC) procedure (27). Because polyamine levels vary with environmental conditions, control cultures were sampled for each experiment.
Animals. TRAMP mice were a kind gift from Dr. Norman Greenberg, and a new colony has been established and maintained at the University of Wisconsin. FVB mice were obtained from Harlan Sprague-Dawley and bred with TRAMP females to produce TRAMP × FVB[F1] mice for these studies. Male TRAMP × FVB mice were confirmed positive for the TRAMP transgene by PCR following published protocols (28). Animal care and use were in accordance with protocols approved by the University of Wisconsin-Madison School of Medicine and Public Health Animal Care and Use Committee and the NIH Guide for the Care and Use of Laboratory Animals.

Results

Androgen induces SSAT mRNA expression. LNCaP cells cultured in androgen-depleted medium (see Materials and Methods) were treated for 96 hours with 0.05 and 1 nmol/L of the androgen analogue R1881. Results published from our laboratory have shown that 1 nmol/L R1881, which closely resembles the androgen levels in normal male human serum, produces high ROS levels in LNCaP cells after a 96-hour treatment, whereas 0.05 nmol/L R1881 causes a minor decrease in cellular ROS (10–14). We have performed qRT-PCR experiment to detect the expression of SSAT mRNA in cells treated with 0, 0.05, and 1.0 nmol/L R1881. The expression of GAPDH mRNA has been used as a control to normalize the qRT-PCR data. The results for the 96-hour treatment are shown in Fig. 1A. qRT-PCR data show that a 25-fold to 30-fold increase in the SSAT mRNA level only in 1.0 nmol/L R1881–treated cells and not in untreated or 0.05 nmol/L R1881–treated cells. The time course of SSAT mRNA production after treatment with 1 nmol/L R1881 is shown in Fig. 1B. The induction of SSAT mRNA production has not been observed up to 48 hours and has been observed only between 48 and 72 hours of androgen exposure. The ROS levels of LNCaP cells treated with varying concentrations of R1881 for different times are shown in Fig. 1C. The ROS levels do not increase for 48 hours after treatment and start increasing only between 48- and 72-hour treatment. Thus, these data show that the androgen-induced SSAT gene expression that causes polyamine oxidation and ROS production runs parallel to the time course of androgen-induced ROS production in LNCaP cells. It is also evident that SSAT gene expression is induced only at androgen concentration that increases ROS but not at androgen concentration that does not increase cellular ROS.

Androgen-induced changes in cellular polyamine level confirm an increase in SSAT enzyme activity. To test if the increase in SSAT mRNA level also translates into higher SSAT enzyme activity, we have tested the effect of androgen on cellular polyamine and acetyl polyamine (Ac-polyamine) levels in untreated LNCaP cells and cells treated with 1 nmol/L R1881 for 96 hours. We have also pretreated the cells with 25 μmol/L APAO inhibitor CPC-200 to test the efficacy in inhibiting APAO as observed by the effect on androgen-induced changes in cellular polyamine levels. Under our culture conditions, it has been reported that 25 μmol/L CPC-200 completely inhibits APAO in most cell lines, including CaP cells (23, 24). The polyamine and Ac-polyamine levels in cells treated with 0 and 1 nmol/L R1881 for 96 hours were increased changes in polyamine and spermidine levels by 6-fold to 10-fold, decreases spermine levels by one-half, and markedly increases N-acetyl spermidine and N-acetyl spermine levels that are undetectable in untreated cells. These results confirm that androgen treatment not only increases the SSAT mRNA level (Fig. 1) but also enhances SSAT enzyme activity, which causes the increase in spermidine and spermine catabolites—Ac-polyamines, putrescine, and spermidine. CPC-200 treatment alone has little effect on most cellular polyamine levels (except for a small increase in acetyl spermine level). In 1 nmol/L R1881–treated cells, however, CPC-200 pretreatment almost completely blocks the R1881–induced increase in putrescine and spermidine levels and causes 3–4-fold increase in N-acetyl-spermidine and N-acetyl-spermine levels without appreciably changing spermine level. These data, in addition to a small but significant increase in N-acetyl-spermine level in cells treated only...
with CPC-200, show that 25 μmol/L CPC-200 efficiently blocks 
APAO enzyme activity, thus inhibiting polyamine oxidation and 
increasing cellular N-acetyl-polyamine levels. In addition, the 
increase in Ac-polyamines also shows that CPC-200 alone has 
little effect on the SSAT gene expression and/or SSAT enzymatic 
activity in the androgen-treated cells.

**Androgen-induced ROS production is relatively lower in** 
**SSAT siRNA transfected LNCaP cells.** To further confirm that 
SSAT is the major player in androgen-induced ROS production in 
LNCaP cells, we have constructed one LNCaP cell clone stably 
transfected with siRNA against SSAT (siSSAT). The ability of the 
siRNA to reduce SSAT mRNA level was first confirmed using qRT- 
PCR. The results are shown in Fig. 3A. These data have been 
normalized to the 18S rRNA level. The results show that 1 nmol/L 
R1881-induced increase in SSAT mRNA level in the siSSAT clone is 
nearly 80% less than that observed for 1 nmol/L R881–treated 
LNCaP cells transfected with a control vector. The acetylated 
polyamine levels in the vector control and siSSAT-transfected cells 
are shown in Table 1. Cells expressing siSSAT show over 4.5-fold 
decrease in acetylated spermidine and over 30-fold decrease in 
acetylated spermine level, which confirms that the decrease in 
mRNA level parallels a decrease in SSAT enzyme activity. The SSAT 
enzyme activity in R1881-treated and untreated LNCaP cells, cells 
transfected with a control vector, and siSSAT-transfected cells are 
shown in Fig. 3B. In both wild-type LNCaP cells and cells 
transfected with a control vector, the SSAT enzyme activity has 
been increased by over 2-fold. In siSSAT transfected cells, however, 
the enzyme activity is less than half of the other two cell lines and 
the activity does not change after R1881 treatment. The effects of 
1 nmol/L R1881 treatment on the ROS levels in LNCaP cells and 
siSSAT clone as determined by a DCF dye oxidation assay are 
shown in Fig. 3C. R1881 treatment has no significant effect on ROS 
production in siSSAT clone compared with a nearly 1.5-fold 
increment induced by R1881 in LNCaP cells transfected with the 
control vector. The difference in percentage induction of ROS levels 
in wild-type versus siSSAT clone is statistically significant with a P 
value of <0.001 as determined using a two-tailed Student’s t test.

**CPC-200 treatment completely blocks androgen-induced** 
**ROS production in LNCaP cells.** To test if CPC-200 treatment can 
also block the androgen-induced ROS production in the wild-type 
LNCaP cells, we have determined the relative changes in ROS levels in 
LNCaP cells that are either untreated or pretreated for 24 h with 
25 μmol/L CPC-200 and exposed to graded concentrations of R1881 
for 96 h. The results of ROS measurement in CPC-200 pretreated 
and untreated cells exposed to increasing androgen concentrations 
are shown in Fig. 4. Data points, SDs, and P values are calculated 
from the reading of six wells of a 96-well plate, wherein each plate 
was run in triplicate and the experiment was repeated twice. CPC- 
200 (25 μmol/L) pretreatment not only completely blocks the 
R1881-induced ROS production, the ROS levels of CPC-200 
pretreated cells are actually even lower than that of control 
androgen-untreated cells. These data confirm that polyamine 
catabolism is one of the major causes of the ROS production of 
prostate cells in general and androgen-induced enhancement of 
ROS in androgen-dependent CaP cells in particular.

We have not observed any effect of CPC-200 pretreatment on the 
androgen receptor level in both androgen-treated and untreated 
cells (data not shown), indicating that the decrease in cellular ROS 
is not due to changes in cellular androgen receptor level in CPC-
200–treated cells.

**CPC-200 treatment markedly reduces tumor formation** 
in TRAMP animals and significantly increases life expectancy. 
To determine if CPC-200 treatment induced reduction in the 
ROS levels and delays prostate tumor progression in vivo, we tested

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![Putrescine (Pu), spermidine (Sd), spermine (Sm), N-acetyl spermidine (N-Ac-Sd), and N-acetyl spermine (N-Ac-Sm) levels of LNCaP cells treated as follows: control (Con), 25 μmol/L CPC-200 for 120 h (CPC-200), 1 nmol/L R1881 for 96 h (R1881), 25 μmol/L CPC-200 for 24 h followed by 1 nmol/L R1881 for 96 h (CPC-200 + R1881). All data points and SDs are calculated from the results of two separate experiments run in triplicates.](image-url)
its effect on tumor formation in TRAMP × FVB hybrid animals (29) that spontaneously develop palpable prostate tumors by ~12 to 16 weeks of age, and all die due to CaP. We tested a dose of 25 mg/kg of CPC-200 given i.p. biweekly over a period of 10 weeks for a total of six treatments. This dose is slightly above the 20 mg/kg dose required to completely inhibit mouse APAO enzyme activity in vivo and is well below the maximum tolerated dose of CPC-200 in mice (100 mg/kg daily for 14 days; ref. 24). This dose has shown no overt toxicity and no observable side effects in mice. In our study, no overt sign of toxicity, abnormal behavior, body weight loss, or any other observable symptom was detectable, either short term after each of the successive CPC-200 injections or long term until the end of the study.

The difference in median survival time was assessed using a Wilcoxon rank sum test. Across two independent studies that include a total of 26 animals per arm, we have observed median survival of 12.0 weeks versus 17.5 weeks after first treatment for vehicle control versus CPC-200–treated mice, yielding a statistically significant (P = 0.03) improvement in median survival by 5.5 weeks for CPC-200–treated animals (Fig. 5). As treatment has begun at an average age of 8 weeks in these studies, this equates to median survival ages of ≥20 weeks for control and ≥25 weeks for CPC-200–treated animals. Therefore, six injections of CPC-200 have resulted in >25% increase in overall life expectancy.

### Table 1. Acetylated polyamine levels (nmol/10^6 cells) in vector control and siSSAT clones of LNCaP cells

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<tr>
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<th>Vector control</th>
<th>siSSAT</th>
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<tr>
<td></td>
<td>N-Ac-Sd</td>
<td>N-Ac-Sm</td>
</tr>
<tr>
<td>Control untreated</td>
<td>NQ</td>
<td>ND</td>
</tr>
<tr>
<td>1 nmol/L R1881</td>
<td>33.3 ± 6.9</td>
<td>6.3 ± 2.6</td>
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<tr>
<td>1 nmol/L CPC-200</td>
<td>7.2 ± 2.5</td>
<td>0.21 ± 0.15</td>
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NOTE: Acetylated spermidine and spermine levels in untreated and R1881-treated LNCaP cells transfected with either control vector or vector expressing siSSAT. Mean ± SDs of two separate experiments run in triplicates. All data are in nmol/10^6 cells. Abbreviations: N-Ac-Sd, N-acetyl spermidine; N-Ac-Sm, N-acetyl spermine; NQ, not quantifiable; ND, not detectable.

Figure 3. A, qRT-PCR amplification of SSAT mRNA from LNCaP cells stably transfected with siSSAT vector and scrambled vector control (see Materials and Methods for detail). Cells are either grown in the presence of 0 nmol/L or 1 nmol/L R1881. All data are normalized to corresponding 18S rRNA expression. The data are average of readings from six identically treated wells repeated twice in triplicates, and error bars represent the exponent of the variance of data distribution. B, SSAT enzyme activity in picomoles of acetyl spermidine formed per milligram protein per minute in wild-type LNCaP cells, LNCaP cells transfected with control vector, and LNCaP cells transfected with siSSAT either vehicle treated or treated with 1 nmol/L R1881 for 96 h. Each data point and SDs are calculated from cell extracts from three independent experiments, and each extract is run in two separate assays that are run in triplicates. C, cellular ROS levels as determined by the DCF assay (see text) for LNCaP cells stably transfected with vector control and siSSAT either untreated or treated with 1 nmol/L R1881. All data were normalized to the ROS level of untreated control LNCaP cells grown in the absence of androgen (F1/C4 medium). Data points and SDs were calculated from the results of two separate experiments, each run in triplicate 96-well plates.

Figure 4. ROS levels in cells treated with graded R1881 concentration that are either vehicle treated or treated with 25 μmol/L CPC-200. All data are normalized to the ROS level of untreated control LNCaP cells grown in the absence of androgen (F1/C4 medium). Data points and SDs are calculated from the results of two separate experiments, each run in triplicate 96-well plates.
polyamine levels in R1881 (Fig. 2) and the reduction of ROS levels in cells treated with 25 μmol/L CPC-200 (Fig. 4) suggest that most of the ROS production in R1881-treated cells is due to SSAT induction followed by oxidation of acetylated polyamines by constitutively expressed APAO rather than a direct oxidation of spermine by PAOh1.

It is also to be noted that CPC-200 pretreated LNCaP cells growing in the presence of 1 mmol/L R1881 have even less ROS than do untreated cells (Fig. 4). This suggests that the basal level of ROS produced in LNCaP cells growing in the absence of androgen may also be due to low grade oxidation of cellular polyamines that is now blocked by CPC-200 treatment.

As spermine acts as a scavenger for ROS (30), androgen-induced decrease in polyamine levels may also cause the observed increase in ROS levels. CPC-200 (25 μmol/L) pretreatment, however, does not reverse androgen-induced reduction of cellular spermine levels (Fig. 2), although it completely blocks the ROS production (Fig. 4). Therefore, R1881 induced depletion of the cellular spermine level can be ruled out as a major contributor to the increase in the ROS levels.

Our Western analysis for androgen receptor protein expression showed no effect of CPC-200 treatment on androgen receptor levels (data not shown). Thus, we rule out the possibility of CPC-200–induced changes in androgen receptor expression as a cause for the changes in growth and/or ROS.

We have established preclinical efficacy of CPC-200 using the TRAMP × FVB mouse model of prostate carcinogenesis. CPC-200 at a well-tolerated dose of 25 mg/kg given i.p. once every 2 weeks for a total of six treatments significantly inhibited the growth of tumors in this model as evidenced by improved survival (Fig. 5). In our studies, the majority of mice (>90%) were sacrificed due to reaching a predefined tumor size per our animal protocol, thus survival is a surrogate for tumor burden and improvement in survival thus equates with an inhibition of tumor growth by CPC-200 in our TRAMP × FVB model. Therefore, the significant 5.5-week improvement (P = 0.03) in median survival for CPC-200 using this model (Fig. 5) shows the ability of CPC-200 to slow prostate tumor growth. The efficacy of CPC-200 against prostate tumor progression in this animal model supports the hypothesis that polyamine oxidation and resultant increase in ROS play an important role in prostate carcinogenesis and strongly implicates the potential of CPC-200 as a new therapeutic agent for CaP.

Several mechanisms such as expression (or nuclear translocation) of specific transcription factors, such as hypoxia-induced transcription factor-1α, NF-κB, activator protein-1, etc. (9, 11, 28, 29), have been suggested as a probable mode of regulation of specific genes that may control cellular redox status. Enhanced mitochondrial activity (13), suppression of glutathione S-transferase-π expression, and a reduction in the level of total glutathione specifically in CaP cells (8) have also been suggested as probable pathways for an increase in ROS production in CaP. A direct effect of androgen in regulating any of these pathways has yet to be shown. To the best of our knowledge, this is the first report of androgen-induced regulation of a rate-limiting enzyme of a specific biochemical pathway (polyamine catabolism) that is directly related to cellular oxidative stress induction. Because of the high levels of polyamines present in human prostate and CaP cells, this pathway seems all the more important in regulating oxidative stress in the prostate gland and specifically in CaP cells. The spermine level in LNCaP cells reported here is by far the highest among most of the cell lines reported thus far (ref. 31 and related references therein). A higher basal metabolism of spermine may be
one reason for the relatively higher ROS level in LNCaP cells compared with other cell lines (12). A close inspection of the SSAT gene sequence reveals that there are four glucocorticoid response elements (GRE) but no androgen response element (ARE) upstream of the SSAT transcription start site. It has been reported that androgen receptor binds and activates GRE containing promoters (32). Its efficiency of activating promoters containing GRE, however, is much lower than that of activating promoters containing ARE. This may be one reason why the SSAT activation and the consequent ROS production was observed only when the cells were treated with high concentration (≥0.5 nmol/L) of R1881 but not with low concentration (≤0.05 nmol/L) of R1881 (ref. 12; Fig. 1).

Lastly, a decrease in cellular polyamine levels has been related to a delay in tumor growth and progression both in cell culture as well as in animals and humans (33–38). In this report, we observed a significant delay in tumor growth by CPC-200 (Fig. 5) although there is an increase, and not a decrease, in cellular spermidine and putrescine levels and no observable change in cellular spermine level. Therefore, change in cellular polyamine levels is probably not a cause for the delay in tumor progression in the TRAMP × FVB animals developing spontaneous prostate tumor. The detail study of the polyamine and Ac-polyamine levels in the TRAMP and animal tumor tissue has now been undertaken to confirm this point.

To the best of our knowledge, this is the first report of SSAT induction by a hormone (hormone analogue) R1881. Identification of androgen-induced polyamine catabolism leading to enhanced oxidative stress in the prostate cells and significant inhibition of CaP progression by blocking this pathway should open up a new avenue for CaP chemoprevention.

Disclosure of Potential Conflicts of Interest

H.S. Basu and G. Wilding: employment, ownership interest, and/or consultant/advisory board, Colby Pharmaceutical Company. The other authors disclosed no potential conflicts of interest.

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