Testosterone modulates mitochondrial aconitase in the full-length human androgen receptor-transfected PC-3 prostatic carcinoma cells

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Abstract

*In vitro* studies indicated that dihydrotestosterone (DHT) stimulates the enzymatic activity of the mitochondrial aconitase (mACON) in androgen-sensitive prostatic carcinoma cells, LNCaP. Cell proliferation assay determined that DHT doubles the optimal proliferation response of LNCaP cells. The androgen-insensitive human prostatic carcinoma cells, PC-3, were overexpressed in the human androgen receptor to assess the involvement of the native androgen receptor in the regulation by DHT of mACON gene expression. A stable-transfected clone that expresses the full-length androgen receptor was selected and termed PCAR9. The results revealed that DHT-treated PCAR9 cells paradoxically not only reduced the enzymatic activity of mACON but also blocked the biosynthesis of intracellular ATP attenuating cell proliferation. Transient gene expression assay indicated that DHT divergently regulates the promoter activity of the mACON gene in LNCaP and PCAR9 cells. This study suggested that DHT regulates mACON gene expression and the proliferation of cells in a receptor-dependent model through modulation by unidentified non-receptor factors.


Introduction

The differentiation, development and maintenance of the prostate gland are under the direct control of androgen (Cunha *et al.* 1987). The epithelial compartment of the prostate gland in humans and other animals has the unique function of accumulating and secreting extremely high levels of citrate (Franklin *et al.* 1986). The net citrate secretion rate is determined by the rates of citrate synthesis and citrate oxidation; testosterone controls the balancing mechanism in the rat ventral prostate (Costello & Franklin 1991). Evidence is emerging that the enzymatic activity of many enzymes involved in metabolizing citrate in the prostate also depends on dihydrotestosterone (DHT) (Costello & Franklin 2002). Mutation of the androgen receptor is not a major mechanism in the process of prostate cells that become androgen growth independent (Taplin *et al.* 1995). It is well known that DHT has a positive effect on the proliferation of androgen-sensitive prostatic carcinoma (LNCaP) cells; however, early study has indicated that DHT treatment inhibits the proliferation of LNCaP cells which were long-term cultured in a medium without steroids (Liao *et al.* 1995). Other studies have also shown that DHT suppresses malignancy and triggers the cell cycle arrest of the androgen receptor-transfected bone metastatic prostate cells, PC-3 (Yuan *et al.* 1993, Heisler *et al.* 1997). Hence, the function of androgen and androgen receptor in the prostate has been suspected to vary with the state of malignancy.

Aconitase (aconitase hydratase, EC4·2·1·3) containing a \([4Fe–4S]\) cluster is regarded as the key enzyme responsible for the interconversion of citrate and isocitrate in the citric acid cycle (Emptage *et al.* 1983). Citrate oxidation via the Krebs cycle is the major pathway for energy production from carbohydrate and fat metabolism. From previous reports, the citrate that is synthesized by prostate cells is accumulated and secreted...
rather than oxidized. The limiting mitochondrial aconitase (mACON) activity prevents citrate from entering the Krebs cycle that induces accumulation and secretion of citrate from the prostate epithelial cells (Costello & Franklin, 1991). A bioenergetic hypothesis of prostate malignancy suggests that a normal citrate-producing prostatic epithelial cell becomes a citrate-oxidizing cell following a transformation in which mACON is not limiting (Costello & Franklin, 1994). Our previous in vitro study demonstrated that mACON-antisense stably-transfected PC-3 cells have lower citrate utility, intracellular ATP biosynthesis and cell proliferation than the mock-transfected cells (Juang, 2004a). Although early studies have demonstrated that androgen treatment promotes citrate oxidation and citrate secretion in LNCaP cells, in transient rat androgen receptor-transfected PC-3 cells, in the rat ventral prostate and in the pig prostate (Costello et al. 1995, 1996, Franklin et al. 1995, Juang et al. 1995), the regulation of mACON in the human prostate is not well understood.

Materials and methods

Cell culture and chemicals

PC-3 and LNCaP cell lines were obtained from the American Type Culture Collection. LNCaP cells are the unique tumor cell line derived from the supraclavicular lymph node metastasis from the human prostatic carcinoma which exhibits increased proliferation in response to androgens in vitro (Horoszewicz et al. 1983). The PC-3 cell line was isolated from the bone metastasis of a 62-year-old patient with stage IV prostatic cancer (Kaighn et al. 1979). 5α-Androstan-17β-ol-3 one (DHT) and charcoal (dextran-coated) were purchased from Sigma and the BCA protein concentration assay kit was purchased from Pierce (Rockford, IL, USA). Fetal calf serum (FCS) was purchased from HyClone (Logan, UT, USA) and RPMI 1640 medium and RPMI 1640 phenol red-free (RPMI-PRF) medium were purchased from Life Technologies. The FCS, which was treated with charcoal (1 g/500 ml FCS) for 24 h to remove the steroids, was considered to be the charcoal–dextran-treated FCS (CD-FCS) in this study. PC-3 and LNCaP cells were maintained in an RPMI 1640 medium that contained 10% FCS and the medium was changed twice a week.

Construction of the androgen receptor overexpression vector

The full-length human androgen receptor cDNA clone was kindly provided by Dr S Liao (Chang et al. 1988). Two fragments of human androgen receptor cDNA were digested: one was digested with EcoRI and HindIII, and the other was digested with HindIII and XbaI resulting in 2.3 and 1.3 kb fragments respectively. The overexpression vector, pcDNA3 (Invitrogen), was digested with EcoRI and XbaI, and the linearized plasmid DNA was ligated with the 2.3 kb EcoRI–HindIII and the 1.3 kb HindIII–XbaI human androgen cDNA fragments, resulting in the insertion of the full-length androgen receptor cDNA in the polyadenylate region that was controlled by the human cytomegalovirus immediate-early (CMV) promoter (pcDNA3AR). Proper ligation was confirmed by extensive restriction mapping and sequencing.

Stable transfection of androgen receptor into PC-3 cells

PC-3 cells were transfected with pcDNA3AR overexpression vector using lipofectin as described by the manufacturer (Life Technologies). Briefly, the transfected cells were incubated for 48 h in RPMI 1640 supplemented with 10% fetal bovine serum (FBS). Then, the cells were incubated for 2 weeks in the same medium supplemented with Geniticin (G418; PAA laboratories, Linz, Austria) to a final concentration of 800 µg/ml. The resulting Geniticin-resistant cells were plated in a 96-well plate with a limiting dilution. One selected single, resistant colony was then examined to evaluate the expression of the androgen receptor using immunoblotting assay and cell immunocytochemical staining. One clone that expressed the highest concentration of the full-length androgen receptor (110 kDa) in the nuclei was selected and termed the PCAR9. The mock-transfected PC-3 cells (PCDNA) were transfected with controlled pcDNA3 expression vector that lacked the androgen receptor cDNA insert. The PCDNA cells were clonally selected in the same way as the PCAR9 cells.

Immunocytochemical staining of cells

LNCaP, PCAR9 and PCDNA cells were grown on the surface of the cover slides for 48 h. Cells were
washed with PBS and then fixed with cold acetone for 15 min. Cells were immunocytochemically stained using a VECTASTAIN ABC kit according to the manufacturer’s instructions (Vector; Burlingame, CA, USA). Cells were detected using 1:500 of anti-bovine mitochondrial aconitase antiserum (kindly donated by Dr R B Franklin) and rabbit anti-human androgen receptor N-terminal (N-20) and C-terminal (C-19) polyclonal antibodies (Santa Cruz Biotechnology, Santa Cruz, CA, USA).

Reverse transcription (RT)-PCR

Total RNA was isolated using Trizol reagent and the cDNA was synthesized using the superscript III preamplification kit according to the manufacturer’s instructions (Life Technologies). Template-primer annealing was conducted at 50 °C for 2 h. Excess RNA was degraded by RNase H treatment. Table 1 shows the primers used for synthesis of the human androgen receptor cDNA in RT-PCR. The PCR reaction was carried out in a thermal cycler (Thermolyne, IA, USA), for 35 cycles at 94 °C for 1 min, 55 °C for 1 min and 72 °C for 1 min. The PCR products were separated by 2% agarose gel electrophoresis and visualized using ethidium bromide staining.

Chloramphenicol acetyltransferase (CAT) assay

PCAR9 and PCDNA cells were transiently transfected with pMSG-CAT (Amersham) using lipofectin as described above. pMSG-CAT contains the mouse mammary tumor virus (MMTV) promoter as a mammalian positive control vector for the glucocorticoid inducible expression of CAT. After the transfected cells were grown for 12 h in RPMI-PRF medium without FCS, the cells were cultured in the same medium with 2% CD-FCS, with or without 10 nM DHT, for an additional 48 h. Cells were harvested and resuspended in 100 µl 0·25 M Tris buffer (pH 7·5) for CAT assay as described elsewhere (Juang et al. 1995).

Mitochondrial aconitase activity assay

Cells were suspended in 100 µl 0·25 M sucrose buffered with 50 mM HEPES and 0·007% digitonin on ice for 5 min. After centrifugation the pelleted mitochondrial material was resuspended in 100 µl HDGC (20 mM HEPES (pH 7·5), 1 mM dithiothreitol (DTT), 10% glycerol, 2 mM trisodium citrate, 0·5 mg/l leupeptin, 0·7 mg/l pepstatin and 0·2 mM phenylmethylsulfonyl fluoride (PMSF)). The membrane of the mitochondrial particle was broken apart in the presence of 1% Triton X-100. Reaction was carried out in 100 mM Tris--HCl, 1 mM MgCl₂, 1 mM NADP⁺, 1 U isocitrate dehydrogenase (ICDH) and 1 mM citrate (Drapier & Hibbs 1996). The mACON enzymatic activities of the cells were adjusted according to the concentration of protein in the mitochondrial extract, which was measured using the BCA protein assay kit.

Cell proliferation assay

The cellular proliferation of LNCaP, PCAR9 and PCDNA cells in response to DHT was measured by the conversion of an MTS tetrazolium salt

| Table 1 Primers required by RT-PCR to synthesize the human androgen receptor (AR) and β-actin cDNAs |
|---------------------------------|---------------------------------|
| Primer                         | Sequence                        | Location     |
| AR1sense                       | 5′-CCAAGACCTACCGAGGAGCT-3′       | +47 to +66   |
| AR1antisense                   | 5′-GCTGTGAAGTTGCTTCC-3′         | +355 to +336 |
| AR2sense                       | 5′-CGGACGAGGATGACTCAG-3′        | +458 to +475 |
| AR2antisense                   | 5′-TCTTCAGTGCTTTGCCCTGC-3′      | +914 to +895 |
| AR3sense                       | 5′-CTGAGCTTCGCAACTTACAC-3′      | +2172 to +2191|
| AR3antisense                   | 5′-TGTTAGAAGCGCTTGTAGCA-3′      | +2576 to +2577|
| β-Actin-sense                  | 5′-GAAGATCAAGATCATGCTTCTCC-3′   | +975 to +998 |
| β-Actin-antisense              | 5′-CTGGCTCTCAAGTAGCTACAGG-3     | +1698 to +1676|

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(3-(4,5-dimethylthiazol-2-yl)-5 (3-carboxymethoxyphenyl)-2-(4-sulphopheny)-2H-tetrazolium, inner salt) into a formazan product by viable cells and absorbance at 490 nm using the cell Titer96AQueous cell proliferation assay (Promega). In this assay 5000 cells were cultured to each well of a 96-well plate in RPMI 1640 medium with 10% FCS for 48 h, and in RPMI-PRF medium with 2% CD-FCS for 24 h thereafter. The cells were then incubated with 100 µl of 0, 0·1, 1, 10 and 100 nM DHT in the same medium for an additional 72 h. MTS dye solution was added and the plates were read 3 h later. The number of cells in each well was counted using an enzyme-linked immunosorbent assay (ELISA) microplate reader (Dynex Technologies, Chantilly, VA, USA). The medium and DHT treatments were changed daily.

**Immunoblot assay of androgen receptor, mitochondrial aconitases and β-actin**

Cells (1 × 10⁶ cells/flask) at 80–90% confluence were incubated with RPMI-PRF medium with 2% CD-FCS and various concentrations of DHT. Cells were lysed with lysing buffer (62·5 mM Tris (pH 6·8), 2% SDS, 10% glycerol, 5% β-mercaptoethanol, 7 M urea, 5 µg/ml leupeptin, 1 mM PMSF). The concentrations of protein in the aliquot samples were measured using the BCA protein assay kit. Equal amounts of protein (60 µg/lane) were analyzed by the ECL detection system, as described by the manufacturer (Amersham). The proteins were probed with 1:500 anti-bovine mitochondrial aconitase antiserum, 1:500 diluted rabbit anti-human androgen receptor N-terminal (N-20) and C-terminal (C-19) polyclonal antibodies or 1:1000 diluted anti β-actin antiserum (C11; Santa Cruz Biotechnology).

**Citrate secretion and intracellular ATP assay**

Cells (1 × 10⁵ cells/well) were seeded into each well of a six-well plate. After being treated for 48 h with DHT at various concentrations, the media were collected, vacuum-dried, resuspended in 300 µl water, and deproteinized by adding 300 µl 70% (w/w) perchloric acid. The mixture was centrifuged at 1000 g for 15 min to remove the protein. The supernatant was mixed with 200 µl 0·75 M K₂CO₃ on ice for an additional 15 min. The precipitate and supernatant were separated by further centrifugation at 1000 g for 15 min. The citrate concentrations in the supernatants were determined by tracking NADH oxidation from the coupled reactions (Möllering & Gruber 1966). The reaction system contained 1 ml glycylglycine buffer (0·5 M)/ZnCl₂ (0·6 mM) (pH 7–8), 20 µl malate dehydrogenase (MDH; 600 U/ml)/l-lactate dehydrogenase (LDH; 1375 U/ml), 100 µl NADH (10 mM) and 500 µl sample. The reaction was initiated by adding 20 µl citrate lyase (40 U/ml). Cells were collected by centrifugation at 300 g for 10 min. The concentrations of intracellular ATP were determined using an ATP assay kit, as described by the manufacturer (CalBiochem, San Diego, CA, USA). The 100 µl of cell extract in the HEPES buffer (25 mM; pH 7·75) was mixed with the ATP-releasing reagent and the reaction was initiated by injecting firefly luciferin-luciferase solution. The peak height of light emission was measured using a LumiCount (Packard Bioscience, Meriden, CT, USA). The ATP concentrations were determined directly from the standard curve. The concentration of citrate in the medium and the intracellular ATP were adjusted according to the concentrations of proteins of the whole-cell extract, as determined using the BCA protein assay kit.

**Luciferase and β-galactosidase assay**

LNCaP or PCAR9 cells were plated onto 24-well plates at 1 × 10⁴ cells/well 1 day prior to transfection. The cells were transiently transfected using TransFast transfection reagent (0·6 µg/well; Promega), 1 µg/well reporter vector and 0·5 µg/well β-galactosidase (β-GAL) expression vector (pCMVSPORTβgal) in the serum reduction medium (OPTI-medium; Life Technologies) for 4 h. The cells continued to be incubated in RPMI 1640 medium with 10% FCS overnight. Before they were treated with DHT, the cells were washed and incubated for an additional 48 h in the same medium, the reactants were washed twice in PBS and the reaction was terminated by adding 200 µl of luciferase cell culture lysis reagent (Promega). Twenty microliters of cell lysate were used for luciferase assay using luciferase assay kit, and 100 µl cell lysate was used for the β-GAL enzyme assay as described by the manufacturer (Promega). The luciferase activity was determined in relative light units (RLUs) using...
LumiCount (Packard BioScience) and adjusted according to the β-GAL enzymatic activity.

Statistical analysis

Results are expressed as the means ± s.e. obtained from at least three independent replications of each experiment. Statistical significance was determined by paired t-test analysis using the SigmaStat program for Windows version 2.03 (SPSS Inc, Chicago, IL, USA).

Results

In vitro studies indicated that DHT (10 nM) markedly stimulates not only the enzymatic activity but also the protein level of mACON in the androgen-sensitive prostatic carcinoma cell, LNCaP (Fig. 1A and C). The MTS assays revealed that DHT doubles the optimal proliferation response of LNCaP cells (Fig. 1B). The increase in mACON activity was associated with an increase in the intracellular biosynthesis of ATP in the LNCaP cells after DHT treatments (Fig. 2A). However, the citrate concentration in the medium also increased after LNCaP cells were treated with the DHT (Fig. 2B).

The human androgen cDNA was stably transfected to an androgen-insensitive human prostate carcinoma cell line, PC-3, to evaluate the function of the native androgen receptor in the effect of the DHT on mACON gene expression and cell proliferation. The clonally selected PC-3 cells transfected with androgen receptor overexpression vector (pcDNA3AR) or mock-overexpression vector (pcDNA3) were maintained in the RPMI 1640 medium with 10% FCS and 800 µg/ml G418. The concentrations of androgen receptor in the selected clones were determined by immunoblotting assay. The clone, PCAR9, that expressed the highest concentration of full-length androgen receptor, was chosen. Cell immunocytochemical staining revealed that PCAR9 cells expressed androgen receptors that were fixed in the nuclei in the same way as LNCaP cells. However, when LNCaP, PCDNA and PCAR9 cells were immunostained with mACON antibody, the results showed that the mACON of those cells was specifically expressed in the cytoplasm (Fig. 3). Immunoblotting assay, using N-terminal (Fig. 4A) or C-terminal rabbit anti-human androgen receptor polyclonal antibodies, revealed an immunoreactive protein with an apparent molecular weight of 110 kDa in PCAR9 cells, as in LNCaP cells; however, this immunoreactive androgen receptor is absent from mock-transfected PC-3 cells, PCDNA (Fig. 4B). Results from RT-PCR also verified that both PCAR9 cells and LNCaP cells expressed the full-size androgen receptor mRNA (Fig. 4C).

PCAR9 and PCDNA cells were transiently transfected with the positive control vector that contained the androgen-sensitive MMTV promoter fused to a CAT reporter gene. The results of treating cells with 10 nM DHT for 48 h showed that DHT treatment doubled CAT activity in PCAR9 cells; however, DHT treatment did not affect the CAT activity in PCDNA cells (Fig. 5). The PCAR9 cells seemed to express functional AR protein that was transcriptionally active in an amount that sufficed to induce the transcription of the CAT gene. This stable transfectant, which continues to respond to androgen in vitro, provides an accurate tool for studying wild-type human androgen receptor activation and its regulation by androgen.

The regulation by DHT treatments of the proliferation of PCAR9 and PCDNA cells was examined. The results from MTS assays indicated that DHT treatments do not considerably affect the relative proliferation of PCDNA cells. On the contrary, the considerable inhibition by DHT treatment of the growth of PCAR9 cells, unlike that of LNCaP cells, was dose dependent (Fig. 6A).

Considerably more endogenous mitochondrial aconitase is present in PCDNA cells (0.81 ± 0.01 U/µg mitochondria) than in PCAR9 cells (0.62 ± 0.01 U/µg mitochondria) [q;6 units in Fig. 6B are ‘U/µg mitochondria’ (i.e. not µg as in text); which is correct; please check units on all Figures]. The results showed that DHT treatments did not affect the enzymatic activity of mACON in PCDNA cells; however, DHT treatment (10 nM) inhibited enzymatic activity of mACON by 50% in PCAR9 cells (Fig. 6B). Moreover, the intracellular ATP level of PCAR9 cells dropped 60% after 10 nM DHT treatment (Fig. 6C). Although the DHT treatment inhibits the enzymatic activity of mACON in PCAR9 cells, the rate of citrate secretion from PCAR9 cells was also markedly reduced by 40% after 10 nM DHT treatment (Fig. 6D). Immunoblot assay indicated that DHT significantly
down-regulates the protein levels of mACON in PCAR9 cells (Fig. 7A). The results suggested that DHT regulates the enzymatic activity of mACON in PCAR9 cells at the transcriptional level.

The human mACON gene, containing 18 exons (GenBank accession no. HSACO2 G01-HSACO2 G17), was cloned and sequenced in our laboratory (Juang 2004b). An EcoRI-digested 9 kb DNA fragment was cloned and identified as the 5'-flanking region of the mACON gene. Two DNA fragments were cloned into the pbGL3 reporter vector (Progema) designed as pbGL743 (−1013 to +38) and pbGL188 (−158 to +38). Transient gene expression assays indicated that the putative promoter was located at the DNA fragment (−158 to +38) of the 5'-flanking region of the human mACON gene (Fig. 7B). Although DHT treatment (10 nM) doubled the activity of the reporter activity when the pbGL188 vectors were transiently transfected into the LNCaP and PCAR9 cells, DHT treatment (10 nM) was found to enhance the promoter activity of the mACON gene when pbGL743 vectors were transiently transfected into LNCaP cells. In contrast, the same treatment inhibits the promoter activity in pbGL743-transfected PCAR9 cells (Fig. 7C and D). DHT was found to operate by a divergent regulative mechanism on the mACON gene in LNCaP and PCAR9 cells. These results implied that the regulation by DHT of the transcription of the mACON gene in LNCaP or PCAR9 cells may depend on the receptor and the modulation of unknown non-receptor factors.

**Discussion**

The growth of the normal prostate is well known to depend largely on androgens and DHT, which are very important in the prostatic pathophysiology (Lopez-Otin & Diamandis 1998). Early in vivo studies found that testosterone tissue specifically induces the biosynthesis of mACON and its

**Figure 1** Modulation by DHT of enzymatic activity and gene expression of mACON and the proliferation of LNCaP cells. (A) LNCaP cells were treated with different concentrations of DHT, as indicated, in an RPMI-PRF medium with 2% CD-FCS for 24 h. The enzymatic activities of mACON were determined as described in Materials and methods. Experimental data are presented as means±S.E. (n=5) of the enzymatic activity induced by DHT treatments (***P<0.001). (B) LNCaP cells (5000 cells/well) were treated with DHT (0.1–100 nM) in 100 µl RPMI-PRF medium with 2% CD-FCS for 72 h. The number of cells were counted using an MTS assay kit. Each point of the curve represents the mean-percentage stimulated of 490 nm absorbance induced by DHT treatments relative to that of a sample that had been mock-treated (**P<0.05; ***P<0.001). (C) LNCaP cells were treated with different concentrations of DHT, as indicated, in the RPMI-PRF medium with 2% CD-FCS for 24 h. Cells were harvested and lysed to extract protein for use in the immunoblot assay.
enzymatic activity in the rat ventral prostate but not in the liver or kidneys (Costello et al. 1995). The results obtained herein indicated that DHT induced the enzymatic activity of mACON in LNCaP cells in a manner consistent with an earlier study which suggested that treatment with androgen increases the oxidation of citrate in LNCaP cells (Franklin et al. 1995).

The LNCaP cells express the androgen receptor with a single-base mutation that produces a change from threonine to alanine in the androgen-binding domain. The growth of LNCaP cells is induced by both the agonist and the antagonist of androgen (Kokontis et al. 1991). Most studies agree that PC-3 cells do not express androgen receptor proteins by immunoblotting assay or mRNA, as detected by Northern blotting analysis and S1 nuclease protection assay (Webber et al. 1997). Androgen treatment does not affect the growth of PC-3 cells, even though some studies have stated that PC-3 cells express very low levels of functional androgen receptor (Tilley et al. 1995, Chlenski et al. 2001). However, this study revealed that treatment of androgen receptor-transfected PC-3 cells with DHT reduces the enzymatic activity of mACON. The androgen suppresses the growth of the androgen receptor-transfected PC-3 cells herein in a manner consistent with other in vitro or in vivo studies of the androgen receptor-transfected PC-3 cells conducted in different independent laboratories (Yuan et al. 1993, Umekita et al. 1996, Heisler et al. 1997, Shen et al. 2000, Evangelou et al. 2002, Pizzi et al. 2003).

Immunoblot assay and RT-PCR assay in this study showed that the PCAR9 cells express almost the same concentration of full-length androgen receptor as do the LNCaP cells. The activity of recombinant human androgen in PCAR9 cells appears to be transcriptionally functional by activating the MMTV–CAT reporter gene. Heisler et al. (1997) found that the growth inhibition of androgen receptor-transfected PC-3 cells was inhibited in both the higher and the lower androgen receptor-expressing cell lines and suggested that the observed inhibition of growth is not caused by the overexpression of the androgen receptor. Yuan et al. (1993) indicated that the morphology of androgen receptor-transfected PC-3 cells was distinct from that of cells of the parental cell line. Their studies suggested that DHT treatments block the progression of androgen receptor-transfected PC-3 cells through the cell cycle, resulting in growth inhibition and apoptosis. However, the PCAR9 and PCDNA cells herein did not exhibit significantly different morphologies. DHT treatments were not found to induce apoptosis with DNA fragmentation in the PCAR9 cells (data not shown). All the cells in this study were incubated in the RPMI-PRF medium with 2% CD-FSC for 48 h, cells were harvested to determine intracellular ATP levels (A) and the media were collected for citrate assay (B), as described in Materials and methods. Experimental data are presented as mean±S.E. (n=6) of the citrate concentrations and ATP levels induced by DHT treatments, relative to those of the sample that had undergone the control treatment (** P<0.05; ***P<0.001).

Figure 2 DHT upregulates citrate utility and the bioenergy of LNCaP cells. Cells were treated with different concentrations of DHT (0.1–100 nM) in 1 ml RPMI-PRF medium with 2% CD-FSC for 48 h, cells were harvested to determine intracellular ATP levels (A) and the media were collected for citrate assay (B), as described in Materials and methods. Experimental data are presented as mean±S.E. (n=6) of the citrate concentrations and ATP levels induced by DHT treatments, relative to those of the sample that had undergone the control treatment (** P<0.05; ***P<0.001).
2% CD-FCS for 48 h before and after DHT treatment. The low percentage of FCS may have abolished the effect of the cell cycle in this study.

The results obtained herein indicated that DHT inhibits the biosynthesis and the enzymatic activity of mACON, halting the proliferation of androgen receptor-transfected PC-3 cells. Our previous study, using antisense strategy to block the endogenous mACON expression of PC-3 cells, demonstrated the positive correlation among mACON, bioenergy and cell proliferation (Juang 2004a). The androgen receptor-transfected PC-3 cells blocked the mACON enzymatic activity after DHT treatments which was suspected to reduce the cellular bioenergetic synthesis and attenuate cell proliferation.

Results from this study, however, indicated that the medium citrate concentration was decreased after treating PCAR9 cells with DHT. The capability of citrate synthesis and oxidation in prostatic epithelial cells is important in determining the net amount of citrate secretion, and enzymes involved in the metabolism of citrate in the prostate are controlled by each hormone in unison (Costello & Franklin 2002). Previous study has shown that DHT modulates not only the oxidation of citrate but also the synthesis of citrate in LNCaP cells (Franklin et al. 1995). It seems that DHT may also modulate some enzymes that are important in metabolizing citrate in PCAR9 cells.

Immunoblot assays and transient gene expression assays indicated that DHT regulates mACON at the transcriptional level in both LNCaP and PCAR9 cells although the regulation is in the opposite manner in each type of cell. Results showed that DHT upregulates the reporter activity in both LNCaP and PCAR9 cells when the cells were transfected with the pbGL188 reporter vector. It seems that DHT upregulates the gene expression of mACON via an alternative signaling pathway since simple sequence analysis did not reveal a consensus androgen response element (AGAAGANNNTGTTCT; Nelson et al. 1999) within the DNA fragments. Several studies have revealed that, when the exogenous androgen receptor is stably overexpressed into PC-3 cells, some factors determine whether the transcriptional effect is the ligand-dependent or ligand-independent situation (Gkonons et al. 2000, Evangelou et al. 2002, Pizzi et al. 2003). One study
on the α-subunit of the pituitary glycoprotein hormones, luteinizing hormone, have similarly stated that the modulation of transcriptional activity by androgen receptor is independent of direct DNA binding but involves interactions between the cAMP response element (Heckert et al. 1997). A putative cAMP response element (ACGTCA) has been found within the promoter region of the human mACON gene (Juang 2004b). Another report also indicated that androgen receptor can interact with the RNA polymerase IIH and the positive elongation factor P-TEFb, which is independent of the androgen response element in androgen receptor-transfected PC-3 cells (Lee et al. 2000).

However, when cells were transiently transfected with the pbGL743 that contained the full-length (−1013 to +38) of the 5′-flanking region of the human mACON gene, the promoter activity was enhanced in LNCaP cells but suppressed in PCAR9 cells after treatment with DHT. The divergent regulation of DHT appears to be mediated by the region of 1013–159 bp upstream of the translational starting site of the human mACON gene. Several reports from independent laboratories have indicated that additional coregulators and cis-regulatory elements are important in the regulation of genes by androgen (Lu et al. 2000, Robyr et al. 2000, Haendler et al. 2001). The functions of numerous coregulators of nuclear receptors may be selectively modulated by secondary signal transduction pathways that determined...
whether the modulators act as coactivators or corepressors, although the mechanisms by which the recruitment of corepressors or coactivators regulate the function of the nuclear receptor are still not well understood (Fernandes & White 2003). Previous study has indicated that the consensus ratio between the coactivator and the androgen receptor determines the regulative function of the androgen receptor in the prostate (McKenna & O’Mallery 2002). Therefore, DHT may activate the androgen receptor with unidentified coregulatory proteins and then this transcriptional unit binds to the cis-regulatory element, which is presented in the 5′-flanking region of mACON gene, thus determining whether the DHT-bound androgen receptor acts as a coactivator or a corepressor in LNCaP and PCAR9 cells. The precise mechanism of the modulation of the transcription of human mACON gene by DHT must be further investigated.

This study successfully cloned cells that express the functional full-length androgen receptor to human androgen-insensitive prostate carcinoma cells, PC-3. The pathways by which the androgen receptor stimulated the growth of LNCaP cells, but arrested that of PCAR9 cells, are yet to be explained. The paradoxical results concerning mACON gene expression following the treatment

Figure 6 Effect of DHT on cell proliferation, enzymatic activity of mACON, intracellular ATP and citrate secretion of the stable androgen receptor-transfected PC-3 cells. (A) PCAR9 cells (▲) and PCDNA cells (●) were incubated with DHT (0–1000 µM) in 100 µl RPMI-PRF medium with 2% CD-FCS for 3 days and the cell number was determined by MTS assay. Each point of the curve represents the mean percentage±S.E. (n=8) stimulations of 490 nm absorbance induced by DHT treatments relative to that of a sample that had been mock-treated. (B) PCAR9 cells (▲) and PCDNA cells (●) were treated with different concentrations of DHT (0·1–100 nM) in RPMI-PRF medium with 2% CD-FCS for 48 h. The mACON enzymatic activities in the mitochondrial fraction were assayed as described in Materials and methods. PCAR9 cells were treated with different concentrations of DHT (0·1–100 nM) in 1 ml RPMI-PRF medium with 2% CD-FSC for 48 h. Cells were harvested to determine intracellular ATP levels (C) and the media were collected for citrate assay (D). Data from experiments are presented as mean±S.E. (n=6) of the intracellular ATP levels and citrate concentration induced by DHT treatments relative to no-treatment controls (**P<0.05).
of LNCaP and PCAR9 cells with DHT, suggested that DHT regulates the expression of the mACON gene expression in a receptor-dependent manner, by modulating unknown non-receptor factors.

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Androgen regulates mitochondrial aconitase


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