Expression of a hyperactive androgen receptor leads to androgen-independent growth of prostate cancer cells

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Abstract

Cellular changes that affect the androgen receptor (AR) can cause prostate cancer to transition from androgen dependent to androgen independent, which is usually lethal. One common change in prostate tumors is overexpression of the AR, which has been shown to lead to androgen-independent growth of prostate cancer cells. This led us to hypothesize that expression of a hyperactive AR would be sufficient for androgen-independent growth of prostate cancer cells. To test this hypothesis, stable lune cancer prostate (LNCaP) cell lines were generated, which express a virion phosphoprotein (VP)16-AR hybrid protein that contains full-length AR fused to the strong viral transcriptional activation domain VP16. This fusion protein elicited as much as a 20-fold stronger transcriptional activity than the natural AR. Stable expression of VP16-AR in LNCaP cells yielded androgen-independent cell proliferation, while under the same growth conditions the parental LNCaP cells exhibited only androgen-dependent growth. These results show that expression of a hyperactive AR is sufficient for androgen-independent growth of prostate cancer cells. To study the molecular basis of this enhanced growth, we measured the expression of soluble guanylyl cyclase-α1 (sGCα1), a subunit of the sGC, an androgen-regulated gene that has been shown to be involved in prostate cancer cell growth. Interestingly, the expression of sGCα1 is androgen independent in VP16-AR expressing cells, in contrast to its androgen-induced expression in control LNCaP cells. RNAi-dependent inhibition of sGCα1 expression resulted in significantly reduced proliferation of VP16-AR cells, implicating an important role for sGCα1 in the androgen-independent growth of these cells.

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Introduction

Prostate cancer is an epithelial-derived cancer (Cussenot et al. 1994) that involves the action of androgens and androgen receptor (AR), a ligand-dependent transcription factor (Chang et al. 1988). This liganded AR is essential for the growth and survival of primary tumors (androgen-dependent prostate cancer cells; reviewed by Jenster 1999). The primary treatment for prostate cancer is aimed at disrupting the androgen–AR interaction (Eder et al. 2000, Jiang et al. 2004). In the prostate, two main forms of androgens are secreted: testosterone and dihydrotestosterone (DHT). Although DHT is the more potent form, both forms are involved in the development of both the prostate and male secondary sex characteristics (Singh et al. 2000). The liganded AR forms a homodimer and binds to the androgen-responsive elements (AREs) found on the promoters of target genes to regulate gene expression (reviewed in Kastner et al. 1995, Mangelsdorf & Evans 1995, Mangelsdorf et al. 1995, Thummel 1995, Beato & Klug 2000).

The AR is important not only in the development of prostate cancer but also in its progression to the usually lethal androgen-independent form (reviewed in Jenster 1999). Several mechanisms for the transition of prostate cancer from a hormone-sensitive to hormone-refractory form have been proposed. First, the AR pathway is bypassed by activation of oncogenes and/or inactivation of tumor suppressor genes (Jenster 1999). In some cases, this can lead to the loss of AR expression by either gene deletion/mutation or DNA methylation, as exemplified by several prostate cancer cell lines (Russell & Kingsley 2003). Another mechanism is ligand-independent activation of AR, which can be mediated by several growth factors, including keratinocyte growth factor, epidermal growth factor, and interleukin-6 (Smith et al. 2001, Roznovanu et al. 2005). AR mutations that broaden ligand specificity are common in late-stage prostate cancer (Brinkmann et al. 1995, Marcelli et al. 2000), and recent evidence has shown that such mutations cannot only initiate prostate cancer but also cause it to progress to a hormone-refractory stage in transgenic mice (Han et al. 2005).
Lastly, AR gene amplifications are observed in 28–30% of recurring tumors following anti-androgen therapy, leading to increased levels of AR and presumably to a more sensitized and active AR pathway (Jenster 1999). In support of this, it was recently demonstrated that AR overexpression is sufficient to convert androgen-dependent prostate cancer to an androgen-independent form (Chen et al. 2004).

Since these earlier studies demonstrate that hormone-independent prostate cancer cells exhibit increased AR expression and, thus, transcriptional activity (Kokontis et al. 2005), and AR function has been linked to the development and progression of androgen-independent prostate cancer form, we sought to study the direct effect of AR transcriptional activity on the growth of prostate cancer cells. Hence, we hypothesized that expression of a hyperactive AR can convert the growth of prostate cancer cells from androgen dependent to androgen independent. Our data show that fusion of the activation function of virion phosphoprotein (VP)16 (Tiley et al. 1992) to the AR renders a receptor that has significantly higher transcriptional activity than the native AR. Stable expression of VP16-AR in prostate cancer cells yielded androgen-independent cell proliferation. Importantly, this androgen-independent growth correlated with androgen-independent expression of soluble guanylyl cyclase-α1 (sGCα1), an AR-regulated gene that has been previously implicated in hormone-refractory prostate cancer cell growth (Cai et al. 2007a). These data show that mechanisms that elevate AR transcriptional activity can lead to hormone-refractory growth of prostate cancer cells and suggest that androgen-independent expression of sGCα1, and perhaps other AR-regulated genes, may be responsible for the cell growth.

Materials and methods

Plasmids

To make VP16-AR/pCI-Neo, hAR/pSG5 (Bubulya et al. 1996) was digested with BamH1/DraI and inserted into pcDNA3.1/Neo(+), generating hAR/pcDNA3.1/Neo(+). The VP16 transactivation domain (amino acids 414–490; Berger et al. 1990) was synthesized by PCR and inserted into the BamH1 site of AR/pcDNA3.1/Neo(+), making VP16-hAR/pcDNA3.1/Neo(+). VP16-hAR was digested out of VP16-hAR/pcDNA3.1/Neo(+) with NheI and NotI and inserted using these same restriction sites into pCI-Neo, yielding VP16-AR/pCI-Neo.

The reporter plasmids MMTV-CAT, PSA-CAT, and hKLK2CAT have been described previously (Bubulya et al. 1996). Transfection efficiency was standardized according to β-galactosidase activity, which comes from the transfected pCH110 plasmid (Shenk et al. 2001). The PSA-Luc (Shenk et al. 2001) and ARE4-Luc (Chen et al. 2006a) luciferase reporters have been described previously.

Transient transfection and reporter gene assays

Transient transfection with reporter gene plasmids of LNCaP and stable cell lines was carried out using the CaPO4 method as described previously (Chen et al. 2006b). For all transfection experiments, cells were transfected with 2 μg pCH110 and enough pTL1 to bring the final plasmid amount to 10 μg per dish (Chen et al. 2006b). Whole cell extracts were prepared and subjected to CAT (Bubulya et al. 2001) or luciferase assays (Cai et al. 2007a). CAT assay results were quantified using the Bio-Rad Molecular Imager FX and Quantify One software (Philadelphia, PA, USA).

Generation and growth of stable cell lines

Generation of stable LNCaP cell lines using a pCI-Neo-based plasmid has been already described (Chen et al. 2006b). Several cell lines were generated, which express VP16-AR, from which three lines (V94, V134, and V149) were selected for study in this work. C14 is a control LNCaP cell line stably transfected with an empty pCI-Neo vector (Chen et al. 2006b). For the experiments described here, these LNCaP stable cell lines were grown in RPMI 1640 complete medium (Sigma) containing 1–5% fetal bovine serum (FBS; Hyclone Logan, UT, USA), depending on the experiment, and 0.1 mg/ml neomycin.

Cellular proliferation assay

C14 or V94 cells were seeded at 3×10⁴ cells per well in 24-well plates, with 1 ml RPMI 1640 phenol-free medium containing either 1 or 5% dextran coated charcoal (DCC)-FBS (Hyclone). After 2-day incubation, the cells were treated with ethanol (vehicle control) or 1 nM R1881. After 0-, 3-, and 6-day incubation periods, the cells were treated with ethanol (vehicle control) or M Casodex were added together with ethanol or R1881. Note that the cells with siRNA transfection were treated with R1881, 24 h after transfection, as described previously (Cai et al. 2007a). Fifty nanomolar of control or sGCα1 siRNA were used (Cai et al. 2007a).

Semi-quantitative RT-PCR and real-time quantitative PCR analyses

The TRlzol reagent (Invitrogen) was used to isolate total RNA from C14, V94, V134, and V149 cells and subjected to either semi-quantitative reverse transcription
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PCR (RT-PCR) or real-time quantitative PCR (Q-RT-PCR) analysis using SYBR Green (iScript; Bio-Rad). The PCRs were carried out utilizing the following upstream and downstream primers respectively, for each gene: PSA, 5'-GGACGATGGAACCAGAGGAG-3’ and 5'-CCCATGAACGTCTTCTCTGA-3’; sGcz1, 5'-AGCAGTGAGAGCTGGAT-3’ and 5'-CTCATCCAGGTGGAGTCGA-3’; TMPRSS2, 5’-CAGTGCCCCAGCAGCACCT and 5’-ACACCATTCCTCGTCTCTC-3’; and GAPDH, 5'-GAAGGAGATTGATGCTG-3’ and 5'-AGGAGGATTGATGCTG-3’. Note that GAPDH was used as a control for RNA amount.

Western blotting
Cell extracts were prepared by boiling cells for 5–10 min in SDS sample buffer (Chen et al. 2006b). Nitrocellulose blots were probed with antibodies against sGcz1 (Cayman Chemical, Ann Arbor, MI, USA), AR (Upstate, Billerica, MA, USA), VP16 (Clonetech), cyclin-dependent kinase 1 (CDK1) (Cell Signaling, Boston, MA, USA), CDK2 (Santa Cruz Biotechnology, Santa Cruz, CA, USA), CDK4 (Santa Cruz Biotechnology), cyclin D (Santa Cruz Biotechnology), retinoblastoma (Rb) (Cell Signaling), PSA (Biodesign, Carmel, NY, USA), p27 (Santa Cruz Biotechnology), sGcz1 (Cayman Chemical), β-actin (Abcam, Cambridge, MA, USA), and β-tubulin (Chemicon, Billerica, MA, USA), and then developed using the chemiluminescence detection system from Amersham. The cells were treated with 1 nM R1881, with or without 10 μM Casodex or 10 μg/ml cycloheximide (CHX). Note that β-actin or β-tubulin was used as a control for protein amount.

Immunocytochemistry
LNCaP cells were grown in 2% DCC-FBS on glass coverslips and treated with either ethanol (carrier) or 1 nM R1881 for 2 days. The cells were then fixed with formaldehyde and incubated with Image-IT FX signal enhancer (Molecular Probes, Carlsbad, CA, USA). After rinsing with PBS, the cells were incubated with an anti-AR antibody (Santa Cruz Biotechnology). This was followed by staining with secondary antibody (Molecular Probes) and development with ProLong Gold antifade reagent (Molecular Probes). Images were viewed using a confocal microscope (Olympus Fluoview IX70, Center Valley, PA, USA).

Results
Expression of VP16-AR results in markedly higher AR transcriptional activity
Since studies have demonstrated that hormone-independent prostate cancer cells exhibit increased AR expression and transcriptional activity (Kokontis et al. 2005) and AR function has been linked to the development and progression of androgen-independent prostate cancer form (Chen et al. 2004), we hypothesized that expression of a hyperactive AR can convert the growth of prostate cancer cells from androgen dependent to androgen independent. In order to test this hypothesis, a hyperactive AR was generated through the fusion of the activation domain of VP16, a strong transcriptional activator (Tiley et al. 1992), with a full-length AR protein to produce the fusion protein VP16-AR (Fig. 1A). In transient transfection experiments, this fusion protein exhibited strongly androgen-dependent transcriptional activity on different promoters, three- to sixfold stronger than endogenous AR in LNCaP cells (Fig. 1B). As shown in Fig. 1C, VP16-AR protein is expressed in this experiment, but this expression is significantly weaker than that of endogenous AR. In addition, and interestingly, androgen has no effect on the stability of this transiently expressed VP16-AR protein, while it is significant on the endogenous AR (Fig. 1C). To better compare the relative activities of VP16-AR and AR, we transfected COS cells that do not express endogenous receptor. This experiment showed that VP16-AR has a fourfold higher activity than AR on PSA-Luc (Shenk et al. 2001) and nearly 20-fold higher on ARE4-Luc (Chen et al. 2006a) (Fig. 1D). The expression levels of the two proteins are comparable (Fig. 1E), arguing that increased activity of VP16-AR is not due to increased protein expression.

To study the biological effects of this hyperactive AR, stable LNCaP cell lines were generated expressing VP16-AR. All VP16-AR-expressing cell lines have elevated R1881-induced AR activity, ranging from 3- to 35-fold better than the endogenous activity found in control C14 cells (Fig. 1F). Interestingly, and surprisingly, V272 cells that exhibited the highest AR activity grew very poorly (Chen & Shemshedini unpublished results), suggesting that high AR activity that is supra-physiological can be detrimental to the growth of prostate cancer cells. Therefore, we elected to use for the studies the below cell lines with moderately elevated, and thus more physiological, AR activity.

Expression of VP16-AR leads to androgen-independent growth of prostate cancer cells
The stable LNCaP cell lines were first monitored for the expression of VP16-AR. RT-PCR was used to show that VP16 was expressed in VP16-AR cells (V94, V134, and V149) but not in control cells (C14) (Fig. 2A). As expected, the three VP16-AR stable cell lines exhibited a markedly higher AR mRNA expression than C14 cells in either the presence or absence of R1881 (Fig. 2A). Western blotting demonstrated expression of VP16-AR protein in all three V cell lines but not in C14 cells.
Additionally, R1881 increased the levels of VP16-AR and the anti-androgen Casodex (bicalutamide) inhibited this positive androgen effect (Fig. 2B), as previously observed with endogenous AR (Bai et al. 2005). This result shows that androgen affects stably expressed VP16-AR but not transiently expressed VP16-AR (Fig. 1C). Further analysis using different concentrations of R1881 revealed that the VP16-AR protein was equally sensitive to the androgen stabilizing effect as native AR (Fig. 2C). To directly measure the stability of these two proteins, the protein synthesis inhibitor CHX was used. CHX had a similar effect on AR and VP16-AR protein levels, and the kinetics of recovery of protein levels induced by R1881 was also similar for AR and VP16-AR (Fig. 2D), indicating that these two proteins do not differ significantly in stability. To examine the expression of these two proteins in cells, immunocytochemistry was performed using an AR antibody that can detect both endogenous AR and VP16-AR. As shown in Fig. 2E, C14 cells express small levels of AR scattered throughout the cell without androgen. Treatment with R1881 caused increased AR levels and nuclear localization (Fig. 2E), as observed previously (Mora et al. 1996). Interestingly, the V cell lines express higher protein levels than the C14 cells, representing signal from AR and VP16-AR, and positive signal is found in the cytoplasm and nucleus (Fig. 2E). Addition of R1881 enhanced these protein levels and induced nuclear localization, just as in C14 cells. Collectively, these data suggest that in the absence of R1881 the V cells have some nuclear VP16-AR protein, which may be responsible for the androgen-independent growth of these cells (Fig. 3). To test the hypothesis that expression of a hyperactive AR can convert the growth of prostate cancer cells from androgen dependent to androgen independent, the VP16-AR-expressing cells were monitored for cell growth. We performed an MTT growth assay to measure the proliferation of C14, V94, and V149 cells grown in...
steroid-reduced culture medium (5% DCC-extracted serum). Under these growth conditions, C14 cells did not grow in the absence of R1881 (Fig. 3A and B). However, and importantly, both V94 (Fig. 3A) and V149 (Fig. 3B) cells exhibited a time-dependent increase in cell number in the absence of androgens, and this growth is statistically higher than that of C14 cells. In fact, the androgen-independent growth of V149 cells is comparable with the androgen-induced growth of C14 cells (Fig. 3B). Collectively, these data show that expression of VP16-AR in LNCaP cells leads to androgen-independent cell growth. Interestingly, despite the androgen-independent growth of V94 and V149 cells, this growth is enhanced by R1881 treatment, and V94 cells grew statistically better than C14 cells in the presence of R1881 (Fig. 3A and B).

To demonstrate that the growth of V94 and V149 cells in the absence of exogenous androgen was indeed androgen independent, cell growth was measured in the presence of Casodex. As shown in Fig. 3C, Casodex suppressed the R1881-induced growth of C14, V94, and V149 cells, but had no effect on the R1881-independent growth of V94 and V149 cells. Interestingly, Casodex almost completely inhibited the growth of C14 cells, while it only reduced the growth of V94 and V149 cells to the level observed in the absence of R1881 (Fig. 3C). These data show that the VP16-AR-expressing cells exhibit a growth phase that is insensitive to Casodex and thus independent of androgens.

Our data above (Fig. 3A–C) demonstrate that V94 and V149 cells grow better in the presence of R1881 than in its absence, suggesting that their growth is androgen inducible. To get a relative measure of the androgen sensitivity of these cells, their growth was compared with that of C14 cells under very low serum (1% DCC-extracted serum) conditions and different concentrations of R1881. As shown in Fig. 3B, the growth profiles of V94 and V149 cells were similar to C14 cells. All three cell lines exhibited a growth response at an R1881 concentration of 0.01–0.1 nM and optimal growth at 1 nM (Fig. 3D). Interestingly, the V149 cells elicited the highest growth response to R1881 treatment and C14 cells the lowest (Fig. 3D). Collectively, these data show that V94 and V149 cells exhibit androgen-independent proliferation that is androgen inducible.

**VP16-AR-expressing cells display altered expression of cell cycle regulatory proteins**

To begin to understand the molecular mechanism responsible for the androgen-independent growth of
VP16-AR-expressing cells, we used western blotting to measure the expression of several cell cycle regulatory proteins (Fig. 4). Rb protein expression was repressed by androgen in C14 cells, as shown previously (Taneja et al. 2002). Interestingly, Rb expression was significantly reduced in V94 cells, but only in the absence of androgen, when compared with C14 cells. In contrast to C14 cells, however, R1881 induced Rb expression in V94 cells. V94 cells expressed reduced levels of p27 when compared with C14 control cells, in both the presence and absence of R1881. As shown in Fig. 4, CDK1 and CDK2 are expressed each as one protein band in C14 cells, while in V94 cells there is a second, more slowly migrating, band that appears for both proteins. On the other hand, CDK4 expression is the same in C14 and V94 cells.

Expression of sGC\(\alpha_1\) is androgen independent in VP16-AR-expressing prostate cancer cells

To understand the molecular basis of the androgen-independent growth of the VP16-AR cells, we first measured sGC\(\alpha_1\) expression in our VP16-AR-expressing cells. Interestingly, the expression of sGC\(\alpha_1\), as measured by semi-quantitative RT-PCR (Fig. 5A and B), QRT-PCR (Fig. 5C), and western blotting (Fig. 5D), is androgen independent in V94, V134, and V149 cells, mimicking what has been observed in the other androgen-independent prostate cancer cells (Cai et al. 2007a; Cai & Shemshedini unpublished results). This androgen-independent expression is not a common feature of all androgen-regulated genes, since TMPRSS2 (Lin et al. 1999) is androgen induced in both the C14 and VP16-AR cell lines (Fig. 5A–C). We have also confirmed androgen regulation in V94 cells of the PSA and hKLK2 genes (Fig. 5A; Cai & Shemshedini unpublished observations). These findings, together with our data from other androgen-independent prostate cancer cells (Cai et al. 2007a; Cai & Shemshedini unpublished observations), strongly argue that androgen-unresponsive expression of sGC\(\alpha_1\) may be a common property of hormone-refractory prostate cancer cells.

Previous studies have shown that Casodex can induce expression of PSA in prostate cancer cells that over-express AR (Chen et al. 2004). To determine whether this is also true for our VP16-AR-expressing cells, PSA expression was measured by semi-quantitative RT-PCR in cells treated with Casodex. As shown in Fig. 5E, Casodex had a weak negative effect on PSA expression in V94 and V149 cells, the same as C14 control cells.
Interestingly, however, the basal expression of PSA is higher in the VP16-AR cells than in the control cells. By contrast, R1881 was able to induce PSA expression in all three cell lines (Fig. 5E). Similar results were obtained for PSA protein expression with R1881 and Casodex, although the protein levels varied in different cell lines (Fig. 5F). These results demonstrate that Casodex does not have agonistic activity in VP16-AR-expressing cells.

**Discussion**

Transcriptionally active AR is present in all forms of prostate cancer (Trapman & Cleutjens 1997, Meehan & Sadar 2003). Among the multiple mechanisms implicated in the conversion of prostate cancer growth from androgen sensitive to androgen insensitive (reviewed in Jenster 1999), overexpression of AR has been recently shown to be sufficient for this transition (Chen et al. 2004). In support of this, several earlier studies have demonstrated that hormone-independent prostate cancer cell lines exhibit increased AR expression and transcriptional activity (Kokontis et al. 1998, Culig et al. 1999, Gao et al. 1999, Lu et al. 1999, Thalmann et al. 2000). In addition, clinical studies have shown that about 30% of advanced prostate tumors exhibit AR overexpression (Visakorpi et al. 1995, Koivisto et al. 1997, Palmberg et al. 1997). These studies collectively led us to hypothesize that increased AR transcriptional activity, like increased AR expression, can promote androgen-independent growth of prostate cancer cells. This hypothesis was directly tested by expression in LNCaP cells of VP16-AR, a fusion protein that harbors androgen-inducible transcriptional activity that is tenfold higher than the endogenous AR. We demonstrate in this study that cells expressing VP16-AR do indeed exhibit androgen-independent growth under the same culture conditions that yield only androgen-dependent growth for parental LNCaP cells (Fig. 3).

Interestingly, however, the basal expression of PSA is higher in the VP16-AR cells than in the control cells. By contrast, R1881 was able to induce PSA expression in all three cell lines (Fig. 5E). Similar results were obtained for PSA protein expression with R1881 and Casodex, although the protein levels varied in different cell lines (Fig. 5F). These results demonstrate that Casodex does not have agonistic activity in VP16-AR-expressing cells.

sGCζ1 is involved in the androgen-independent growth of VP16-AR-expressing prostate cancer cells

To directly test whether sGCζ1 is involved in V94 cell proliferation, a growth assay was performed, in which sGCζ1 expression was diminished by siRNA transfection. Transfection of sGCζ1 siRNA markedly down-regulates the endogenous expression of sGCζ1 mRNA in V94 cells, when compared with control siRNA (Fig. 6A), as has been observed in C14 cells (Cai et al. 2007a). The same effect was observed on sGCζ1 protein levels in V94 and C14 cells (Fig. 6B). This reduction in sGCζ1 expression results in significantly decreased growth of C14 cells (Fig. 6C), as shown previously (Cai et al. 2007a). Most significantly, sGCζ1 siRNA also inhibited the growth of V94 cells both in the presence and absence of androgens (Fig. 6D). These findings strongly suggest that sGCζ1 is required for the androgen-independent growth of V94 cells.

**Figure 4** VP16-AR-expressing prostate cancer cells exhibit differential expression of cell cycle proteins. C14 and V94 cells were grown for 2 days in the presence or absence of 1 nM R1881, as indicated, and subjected to western blotting to measure the expression of VP16-AR, AR, Rb, p27, CDK1, CDK2, CDK4, cyclin D1, and β-tubulin. Note that the V94 cells express two proteins detected by the anti-AR antibody, endogenous AR, and VP16-AR (upper band).
In view of the recent study reporting that increased AR expression is sufficient to convert prostate cancer cell growth from hormone sensitive to hormone refractory (Chen et al. 2004), it is possible that our VP16-AR cells simply represent another example of androgen-independent growth in response to AR overexpression. However, the data provided argue against this and suggest that our cells may represent another cellular state of hormone-refractory prostate cancer. First, and most importantly, we have no evidence that the antagonist Casodex has any agonistic activity in VP16-AR cells, as observed in AR overexpressing cells (Chen et al. 2004). Indeed, the VP16-AR cells exhibited no detectable Casodex-induced expression of PSA (Fig. 5E and F) or of several other androgen-regulated genes (Hsieh & Shemshedini unpublished observations). Secondly, AR overexpression in LNCaP cells increased their sensitivity to R1881-induced growth (Chen et al. 2004), while expression of VP16-AR had no effect (Fig. 3D).

Thus, our VP16-AR cells are likely to represent a mechanism of androgen independence, which depends more on elevated AR transcriptional activity than expression level. Since no more than 30% of advanced prostate tumors exhibit increased AR expression (Visakorpi et al. 1995, Koivisto et al. 1997, Palmberg et al. 1997), it is possible that our VP16-AR cells may mimic one cancer cellular state that is found among the remaining 70% of tumors lacking increased AR expression. The question remains of what other mechanisms in addition to AR gene amplification can lead to AR overexpression. One can speculate that mutations in the AR gene promoter disrupting the activities of silencer elements can lead to higher AR expression. It is also possible that mutations that disrupt the activity of transcriptional repressors that act on the AR promoter can have the same effect. Perhaps these mechanisms are operating in some prostate tumors, but empirical evidence needs to be obtained in the future.
The molecular changes responsible for the hormone-refractory cancer state in tumors expressing normal AR protein levels are poorly understood. Several mechanisms have been suggested (reviewed in Jenster 1999), including AR mutations that broaden ligand specificity, generation of signaling pathways that lead to ligand-independent activation of AR, and overexpression of AR coactivators. All these provide means by which to activate AR, but they do not address the issue of AR-regulated gene expression that may be responsible for the hormone-refractory cancer state. Our VP16-AR cells provide some insight into this, with our discovery that these cells express the sGCα1 gene, an AR-regulated gene (Cai et al. 2007a), in a hormone-independent manner (Fig. 5A–C). sGCα1 is a component of sGC, the enzyme that catalyzes cGMP synthesis in response to nitric oxide (reviewed in Hanafy et al. 2001). Our earlier study showed that sGCα1 expression is androgen independent in C81 cells (Igawa et al. 2002). Furthermore, we demonstrated that sGCα1 protein expression levels are directly related to the growth capacity of LNCaP cells (Cai et al. 2007a). While the molecular mechanism responsible for the sGCα1 role in cell proliferation is not yet known, it is clear from our data thus far that the sGCα1 protein levels are directly proportional to the growth capacity of prostate cancer cells (Cai et al. 2007a). As expected, we find in this study that endogenous sGCα1 expression is also required for the androgen-independent growth of VP16-AR cells (Fig. 6D), suggesting that androgen-independent expression of sGCα1 may be responsible for the hormone-refractory growth of LNCaP cells. Importantly, the transition to androgen-independent expression of sGCα1 is not unique to LNCaP cells, but may be a common property of prostate cancer cells since we have observed a similar expression pattern in hormone-refractory CWR22-Rv1 (Wainstein et al. 1994) and Mda-P109 (Navone et al. 1997) cells (Cai & Shemshedini unpublished observations).

It is interesting to note that androgen-independent expression is not observed globally on AR-regulated genes in hormone-refractory cells, but in fact is target gene specific. Our VP16-AR cells here (Fig. 5A and B) and C81 cells (Igawa et al. 2002) exhibit androgen-induced expression of two hallmark AR-regulated genes, PSA and TMPRSS2. The contrasting expression patterns of PSA and TMPRSS2 when compared with AR induces androgen-independent cell growth  ·  C-L HSIEH, C CAI and others 21

Figure 6  sGCα1 is involved in the androgen-independent growth of V94 cells. (A) V94 cells were transfected with sGCα1 siRNA or an unrelated control siRNA, and expression of sGCα1 and sGCβ1 was measured by semi-quantitative RT-PCR. (B) Western blotting shows reduced sGCα1 protein expression in C14 or V94 cells transfected with sGCα1 siRNA. (C) C14 or (D) V94 cells were transfected with sGCα1 siRNA or control siRNA and grown for 0–6 days in the presence or absence of 1 nM R1881, as indicated, and measured for cell number. Bar graphs represent the average of three independent experiments plus the s.d. Asterisks indicate statistical significance (P<0.05) on the effect of sGCα1 siRNA on cell growth.
sGCα1 suggest that genes important in prostate cancer biology are selected for hormone-independent expression. This is supported by another novel AR-regulated gene, ETV1, and also elicits androgen-independent expression in VP16-AR (Cai & Shemshedini unpublished observations) and C81 cells (Cai et al. 2007b). ETV1 is involved in prostate cancer cell invasiveness (Cai et al. 2007b) and sGCα1 in cell proliferation (Cai et al. 2007a), both processes essential to the progression of prostate cancer. For now, we do not know how many more AR-regulated genes transition to androgen-independent expression or the molecular basis of the gene selection process for such expression. These will be the two important objectives of future work.

Previous work has suggested that reduced Rb protein expression may be responsible for androgen-induced proliferation of LNCaP cells (Taneja et al. 2002). In support of this, we observed that R1881 repressed Rb expression in LNCaP cells (C14 cells; Fig. 4). Interestingly, Rb expression is even lower in the V94 cells without androgen than C14 cells with androgen (Fig. 4), suggesting that reduced levels of Rb in V cells may be involved in their androgen-independent growth. Surprisingly, however, Rb levels are higher in V94 cells treated with androgen than without, indicating a complex function of Rb in prostate cancer cell growth. On the other hand, protein levels of p27 (Kip1) are consistently and significantly lower in V94 than C14 cells (Fig. 4). In view of previous findings showing that increased levels of p27 are associated with decreased prostate cancer cell growth and increased cell cycle arrest (Deep et al. 2007), our finding suggests that decreased p27 expression in V cells may be in part responsible for the increased growth capacity of these cells. Supporting this conclusion is the previous finding that decreased expression of p27 is associated with androgen-independent proliferation of LNCaP cells (Murillo et al. 2001). Previous data also suggest that increased levels of CDKs and cyclins are found in proliferating LNCaP cells (Deep et al. 2007). While we observed no significant change in CDK expression between C14 and V94 cells, a second larger band was detected for both CDK1 and CDK2 in V94 cells (Fig. 4). These larger proteins may represent phosphorylated forms of CDK1 and CDK2, which may be involved in androgen-independent growth of our LNCaP cell lines. Collectively, these data, together with the surprising finding that cyclin D1 expression is reduced in V94 when compared with C14 cells (Fig. 4), show the complex nature of cell cycle regulatory proteins in LNCaP cells. Future work will be directed to a biochemical analysis of the larger CDK proteins and determining what roles these proteins, as well as Rb and p27, play in cell cycle progression of VP16-AR-expressing cells.

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