Suppressor role of androgen receptor in proliferation of prostate basal epithelial and progenitor cells

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Abstract

Early studies have reported the differential roles of androgen receptor (AR) in different types (luminal, basal intermediate, and stromal) of prostate cancer cells. In vivo mouse model tumor studies using the total prostate epithelial knockout mice (pes-ARKO) also revealed that AR played a suppressive role in proliferation of the CK5+/CK8+ progenitor/intermediate cells but a positive role in the CK5-/CK8+ luminal epithelial cells. Using three different resources (one human basal epithelial cell line, one mouse basal epithelial originated progenitor cell line, and a basal epithelium-specific ARKO mouse model), we here demonstrated that the AR in basal epithelial cells of normal prostate plays a suppressive role in their proliferation but a positive role in differentiation into luminal epithelial cells. These results led us to conclude that ARs may play a negative role to suppress CK5+ basal epithelial and progenitor cell proliferation, yet play an essential role to drive basal epithelial cells into more differentiated states. These results may explain why differential AR expression in different cell types within normal prostate is needed and suggest that ARs in prostate basal epithelial cells, although expressed at a very low level, are necessary to maintain the balance between progenitor cells and differentiated luminal epithelial cells. 


Introduction

Normal adult prostatic epithelium includes three types of cells: neuroendocrine cells, luminal epithelial cells, and basal epithelial cells that are aligned along the basement membrane. Cells in the luminal layer express androgen receptor (AR), prostate-specific antigen (PSA), cytokeratin-8 (CK8), and CK18. In contrast, cells of the basal layer express CK5 and CK14 (Verhagen et al. 1988, Hayward et al. 1996, Xue et al. 1998a, Johnson et al. 2000), and AR expression is either undetectable (Prins et al. 1991, Uzgare et al. 2004) or at a very low level (Bonkhoff et al. 1998). More than 80% of proliferating cells are observed in the basal layer, which is thought to contain CK5+/CK8− stem cells that give rise to CK5+/CK8low progenitor/intermediate cells and can then be further differentiated into the secretory luminal CK5−/CK8high cells (Bonkhoff et al. 1994, Xue et al. 1998b).

Under normal conditions, adult prostate epithelial cells remain in homeostasis with infrequent turnover. However, the homeostasis is broken upon androgen depletion. When an adult male is castrated, the serum testosterone level decreases rapidly and the prostate regresses with the majority of luminal epithelial cells dying through apoptosis, whereas the majority of basal epithelial cells remain alive (Bonkhoff et al. 1994). These contrasting results suggest that response to androgen deprivation treatment is different in normal luminal epithelial cells vs basal epithelial cells.

Using established mouse and human basal epithelial/progenitor cells as well as an in vivo basal-ARKO (Cyp19a1−/−) mouse model, we report that ARs may play a negative role to suppress proliferation of basal epithelial/progenitor cells, yet AR expression also plays essential roles to drive these cells into a more differentiated state.

Materials and Methods

Development of basal-ARKO mice

All research was approved and conducted following the rules/regulations of the Institutional Review Board (IRB) and University Committee of Animal Research (UCAR),
which was fully credited by AAALAC (ID: A-3292-01), at the University of Rochester Medical Center. The generation of basal-ARKO mice was carried out by mating female mice harboring floxed AR alleles with male transgenic mice expressing Cre recombinase under the control of the CK5 promoter. The detailed strategies of mating and genotyping were described (Yeh et al. 2002). Basal-ARKO mice carrying floxed AR and Cre were selected after genotyping.

### Histological examination of mice

Briefly, mice were killed using CO₂. Ventral prostate (VP), anterior prostate (AP), and dorsal lateral prostate were removed, fixed in 10% neutral buffered formalin, and subjected to paraffin sectioning. Tissue sections from VP at different ages were stained with hematoxylin and eosin and their morphology examined. The images were acquired using an E800 microscope (Nikon) and a SPOT camera (Diagnostic Instruments Sterling Heights, MI, USA) and were analyzed using SigmaScan Pro Sysat software (version 5.0; SPSS Inc., Chicago). For staining of tissues, antibodies of AR (1:200, N20; Santa Cruz, Bio-technologies), anti-CD44 (1:250; N20, Santa Cruz), anti-PSCA (1:200; N20, Santa Cruz), CD49f, and CD44 (eBioscience), washed again with 1% FBS in PBS; and then analyzed by flow cytometry according to the previous publication (Lai et al. 2009).

### Cell culture

Cultures of mouse prostate epithelial cells (mPrE) were maintained in RPMI medium containing 10% fetal bovine serum (FBS). Normal human basal cells from Lifeline Cell Technology (Lifeline-basal) were maintained in ProstaLife prostate epithelial cell culture medium (Lifeline, Frederick, MD, USA) supplemented with LifeFactors (transforming growth factor-β, epinephrine, insulin, transferrin, and hydrocortisone) provided with medium.

#### MTT cell growth assay

Cells were plated onto 24-well plates. At various time points indicated, MTT solution was added onto cells for 30 min in 5% CO₂, isopropanol was used to dissolve the MTT salt, and ODs were measured at 570 nm.

### Immunohistochemical staining/immunofluorescence staining of cells/tissues

Cells were seeded on four-well chamber slides and fixed with methanol. After fixation, cells were washed with PBS three times for 5 min, then cells were blocked with 1% FBS for 1 h, washed with PBS three times, and then incubated with primary antibodies in 3% BSA in PBS overnight at 4°C. Antibodies used were as follows: anti-Ki67 (1:1000; National Chemical Lab, Pune, India), anti-p63 (1:250; Santa Cruz), anti-AR (1:250; N20, Santa Cruz), anti-PSCA (1:200; N20, Santa Cruz), anti-CD44 (1:250; N20, Santa Cruz), anti-CK5 (1:250; Covance), anti-CK8 (1:250; Abcam), and anti-sca-1 (1:250; eBioscience, San Diego, CA, USA). Cells were then incubated with 1:200 diluted biotinylated secondary antibody (Vector Laboratories, Burlingame, CA, USA) and ABC solution (Vector Laboratories) for immunohistochemistry (IHC), or with fluorescent secondary antibodies for immunofluorescence (IF) (either Alexa 594 or Alexa488 tagged). For IHC, cells were stained by AEC (Dako, Carpenteria, CA, USA), followed by Mayor’s hematoxylin counterstaining.

Formalin-fixed paraffin wax-embedded tissues were cut into 5 μm sections, placed on poly-L-lysine-coated slides, deparaffinized, rehydrated, subjected to antigen retrieval for 10 min, blocked with normal horse serum, and incubated with appropriate primary monoclonal antibodies, appropriate fluorescein-tagged secondary antibody (for IF), or visualized by VECTASTAIN ABC peroxidase system and diaminobenzidine (DAB) kit (Vector Laboratories) (for IHC).

### Self-renewal analysis

Single cell suspension (1×10³, in 50 μl medium) was mixed with 50 μl cold Matrigel and the mixture was placed along the rim of the 24 wells with minimum of triplicate experiments. The culture plates were placed in a 37°C incubator for 10 min to let the mixture solidify and 500 μl medium was then added into the well. Cells were grown in culture media and sphere numbers were counted after 7–14 days.

### Western blot analysis

Cell lysates were resolved with 10% SDS gel electrophoresis (SDS–PAGE), blotted with AR antibody (N20; Santa Cruz), and incubated with secondary antibodies conjugated with horseradish peroxidase. Proteins were visualized using the Pierce ECL Western Blotting Substrate (Thermo-Fisher Scientific, Rochester, NY, USA) according to the manual’s instructions. Glyceraldehyde 3-phosphate dehydrogenase (GAPDH) served as loading controls.

### RNA extraction and real-time quantitative PCR (qRT-PCR)

Total RNAs were extracted with Trizol (Invitrogen). mRNAs (2 μg) were used for reverse transcription to cDNAs using Superscript III (Invitrogen). Quantitative qRT-PCR was performed using cDNA, specific gene primers, and SYBR green master mix (Bio-Rad) on an iCycler iQ Multi-color real-time PCR machine (Bio-Rad).

### Flow cytometry analysis of cell markers

Cells were detached with 5 mM EDTA and washed with 1% FBS in PBS (flow washing buffer); stained with sca-1, CD49f, and CD44 (eBioscience), washed again with 1% FBS in PBS, resuspended in PBS; and then analyzed by flow cytometry according to the previous publication (Lai et al. 2009).
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Lentivirus infection
293T cells were transfected with a mixture of DNAs (Lentiviral vectors pWPI) (Addgene, Cambridge, MA, USA) containing AR (vectors used as controls), pMD2.G packaging plasmid, and psPAX2 envelope plasmid (4:1:1 ratio) using Lipofectamine (Invitrogen) according to the transfection protocol. Culture media containing virus was collected 32 h after transfection, filtered, and added onto cultured cells with 2 μg/ml polybrene. Cells were refreshed with culture media 24 h later and cultured for 3 days. As the pWPI vector contains green fluorescent protein (GFP), transfection efficiency was monitored by fluorescence microscopy.

Apoptosis assay
The in situ cell death detection kit (Roche Pharmaceuticals) was used according to the manufacturer’s instruction for detection of apoptotic cells.

β-Galactosidase assay
Six weeks after mating of CK5-Cre/floxed AR and ROSA26, prostate lobes were taken, processed, and embedded with OCT media. The 5 μm sections were used to perform the β-galactosidase assay according to the manufacturer’s instruction.

Luciferase assay
Cells were plated in 24-well plates and transfected with MMTV-luc containing androgen response element (ARE) sequence using Lipofectamine (Invitrogen) according to the manufacturer’s instruction. After transfection, RPMI medium containing charcoal-stripped medium (for mPrE) and basal cell medium (without serum, for Lifeline-basal cells) was added into the culture with addition of various concentrations of dihydrotestosterone (DHT), 0 (ethanol as vehicle control), 1, and 10 nM, and incubated for 48 h. pRL-TK was used as internal control. Luciferase activity was measured by Dual-Luciferase Assay (Promega) according to the manufacturer’s manual.

Statistical analysis
Values were expressed as mean ± s.d. The Student’s t-test was used to calculate P values. P < 0.05 was considered significant.

Results
AR suppresses growth of prostate basal epithelial and progenitor cells
To investigate the potential role of AR in proliferation of prostate basal epithelial and progenitor cells, we selected two established cell lines, human basal epithelial cell line (named Lifeline-basal) and a mouse progenitor cell line (mPrE). Figure 1A includes results of various studies using the mPrE cell line whereas Fig. 1B shows results using the Lifeline-basal cell line.

The mPrE cell line was originally derived from normal mouse prostate tissues (Wang et al. 2000). The morphology of mPrE cells is shown in Supplementary Figure 1A, see section on supplementary data given at the end of this article. The initial examination of characteristics of these cells revealed that they are of basal epithelial origin (Wang et al. 2000) and the IF staining result showed positive staining with basal cell markers CK5 and p63 (Supplementary Figure 1B). The flow cytometric analysis result (Supplementary Figure 1C) further demonstrated that 98% of these cells stained positive for stem/progenitor markers, sca-1 and CD49f, whereas the immunoglobulin G control showed almost no signals for sca-1 and CD49f. These cells were not positively stained for the intermediate cell marker, prostate stem cell antigen (data not shown). Together, these results indicate that the mPrE cells are basal epithelial originated progenitor cells. These mPrE cells express little AR as shown in western blot (Fig. 1A-a), qRT-PCR (Fig. 1A-b), and IF staining (Fig. 1A-d) analyses. To investigate whether the low level of AR is critical in self-renewal/proliferation of these cells, we forcedly expressed AR into the cells via viral infection and MTT assay, sphere formation assay, and Ki67 labeling experiments were performed. Interestingly, the proliferation of the mPrE cells was shown to be significantly suppressed upon AR expression (MTT assay result at day 6, Fig. 1A-c, and Ki67 IF staining result, Fig. 1A-d), indicating that AR plays a negative role to suppress their proliferation. The MTT assay was performed at 0 (vehicle control), 1 (the prostate intra tissue DHT level after castration) (Titus et al. 2005a,b), and 10 nM DHT (the prostate intra-tissue DHT level before castration), and the suppressive effect of AR on proliferation was observed at all androgenic concentrations. Self-renewal tests were also performed and the results showed that the AR-expressing mPrE cells have lesser self-renewal ability compared with control cells (Fig. 1A-e). All these results indicate the suppressive role of AR in self-renewal/proliferation of mPrE progenitor cells.

The Lifeline-basal cells (also named as HPrE), which show characteristics of basal epithelial cells, were originally derived from a normal human donor. These cells were also used in our study to investigate the AR effects on proliferation of basal cells. The morphology of these cells is shown in Supplementary Figure 2A, see section on supplementary data given at the end of this article. These human basal cells showed positive staining with CK5 and p63 but failed to express CK8 (Supplementary Figure 2B). Similar to the mPrE cell line data, a low expression of AR was observed (protein level in Fig. 1B-a and mRNA level in Fig. 1B-b). We also tested whether the proliferation of these cells was also suppressed upon addition of AR. As shown in Fig. 1B-c, the addition of AR suppressed cell growth at all androgenic concentrations tested starting on day 3. IF staining analysis result showing lower numbers of Ki67 positively stained cells upon AR addition (Fig. 1B-d) also indicated that AR plays a suppressive role in proliferation of these basal epithelial cells.
Figure 1 AR expression suppresses self-renewal/proliferation of mPrE (A) and Lifeline-basal epithelial cells (B). (a) Western blot analysis of AR expression. Cell extracts were obtained from mPrE/Lifeline-basal cells and LNCaP cells (used as positive control) and analyzed by western blot analysis. (b) qRT-PCR analysis results of AR mRNA expression. The mRNAs of mPrE/Lifeline-basal cells were analyzed by qPCR analysis. The human prostate cell lines LNCaP and PC3 were used as positive and negative controls respectively. (c) MTT assays at three different DHT concentrations, 0 (vehicle control), 1, and 10 nM, are shown. Western blot results are shown to indicate AR overexpression. (d) Ki67 IF staining results after infection of cells with lentivirus carrying either vector or AR. After infection, cells were plated and immunofluorescence (IF) staining was performed using Ki67 antibody. AR staining presents different levels of AR expression after the infection. (A–e) Self-renewal test on Matrigel after infection of mPrE cells with lentivirus carrying either vector or AR. Quantitation shown at right. *P < 0.03.
Mice lacking AR in basal epithelial cells led to increased proliferation of CK5⁺ cells

In our earlier studies, the increase in CK5⁺ cells was also detected when AR was knocked out in prostate epithelium in TRAMP derivatives of prostate epithelium-specific ARKO (pes-ARKO-TRAMP) mice (Niu et al. 2008). As an extension of this finding and our in vitro results shown in Fig. 1B, we tried to elucidate the significance of the low level of AR in basal epithelial cells, although the endogenous AR level is very low. We developed a mouse model in which AR is specifically knocked out in basal epithelial cells (basal-ARKO) by mating male CK5-Cre mice with female floxAR mice (Yeh et al. 2002). The Supplementary Figure 3A, see section on supplementary data given at the end of this article, shows the strategy for developing basal-ARKO mice and Supplementary Figure 3B genotyping shows Cre and floxAR expressions in transgenic mice. We bred ROSA26R mice with the CK5-Cre, floxed AR/AR mice to monitor the gene deletion efficiency. In the ROSA26R mice, generalized transcription of the β-gal (Glb1) gene is prevented by a preceding transcriptional stop sequence, which is flanked by two loxP sites. In crosses of ROSA26R transgenic mice with promoter-derived Cre transgenic mice, the transcriptional stop sequence was removed by Cre-mediated excision and GLB1 protein will be expressed (Soriano 1999). As shown in Fig. 2A, we observed blue staining in almost all basal epithelium areas, but not in luminal epithelium, suggesting basal epithelium-specific AR knockout. The blue staining was more obvious in VP and dorsolateral prostate lobes than AP lobe (data not shown). To further verify that AR was knocked out in basal epithelium but not in luminal epithelium, we performed IF double staining of VPs of CK5-Cre/WTAR:ROSA26 mice and the GLB1 activity is from the AR-containing basal epithelial cells. In the CK5-Cre/WTAR:ROSA26, indicating that the GLB1 activity is from the AR-containing basal epithelial cells. However, the GLB1 activity was detected only in basal epithelium in CK5-Cre/floxAR:ROSA26 mice and the luminal epithelium did not show blue color (Fig. 3A). It may be that the AR knocked out basal epithelial cells in the CK5-Cre/floxAR:ROSA26 mice were not able to differentiate into luminal epithelial cells, so blue staining was not observed. The luminal epithelium was probably due to the incomplete AR knockout in 20% of basal epithelial cells. These results indicate that basal AR, although the level is very low, is essential to drive cells to differentiate.

The AR role in basal epithelial cells in their differentiation process

We found that AR knockout in basal epithelial cells promoted their proliferation (Figs 1 and 2). We next investigated whether AR in basal epithelial cells plays a role in the differentiation process into terminally differentiated luminal epithelial cells by exploiting the basal-ARKO mice model. We generated CK5-Cre/WTAR:ROSA26 and CK5-Cre/floxAR:ROSA26 mice by crossing the CK5-Cre/WT AR and CK5-Cre/floxAR mice with the ROSA26 mice respectively. We found that GLB1 activity (blue color) was expressed in all prostate tissues including luminal epithelium in the CK5-Cre/WTAR:ROSA26, indicating that the GLB1 activity is from the AR-containing basal epithelial cells. However, the GLB1 activity was detected only in basal epithelium in CK5-Cre/floxAR:ROSA26 mice and the luminal epithelium did not show blue color (Fig. 3A). Similar phenotypes of membrane disorganization were reported when p63 was depleted from the basal cell layer (Liu et al. 2009). They suggested that this phenotype could promote tumor invasion and progression, but whether this phenotype is correlated with any function and any other phenotype remains unclear.

We analyzed the distribution of CK5⁺ and CK8⁺ cells in VP tissues of basal-ARKO and control WT littermate mice (4 weeks old) and found that the CK5⁺ cells (basal epithelial including stem/progenitor/intermediate cells) (Fig. 2C, stained green, arrow) were significantly increased in VPs of 4-week-old basal-ARKO mice compared with WT control mice. The increased CK5⁺ cell numbers could be due to higher proliferation and/or lower apoptotic death. We could detect significantly higher numbers of Ki67 positively stained cells in 4- and 6-week-old basal-ARKO mice compared with WT mice (Fig. 2D). In contrast, we failed to observe significant differences in apoptotic death rate between these two types of mice (data not shown). These results indicate that AR, plays a negative role to suppress proliferation of basal cells, which is in agreement with the in vitro studies previously shown (Fig. 1).

Unlike the luminal cells, the prostate basal cells are long-lived and basal cell differentiation is associated with the irreversible maintenance of self-renewal capacity. These observations are consistent with a stem cell-like behavior of basal cells and indicate that basal cells maintain self-renewal proliferation capacity even in the absence of AR. AR expression in basal cells is associated with slow proliferation and increased proliferation of AR KO mice. This study demonstrates that AR in basal cells is highly correlated with increased proliferation of basal cells in vivo. We showed that the increased proliferation of basal cells in AR KO mice is associated with increased proliferation of AR KO mice. Unlike the luminal cells, the prostate basal cells are long-lived and basal cell differentiation is associated with the irreversible maintenance of self-renewal capacity. These observations are consistent with a stem cell-like behavior of basal cells and indicate that basal cells maintain self-renewal proliferation capacity even in the absence of AR. AR expression in basal cells is associated with slow proliferation and increased proliferation of AR KO mice. This study demonstrates that AR in basal cells is highly correlated with increased proliferation of basal cells in vivo. We showed that the increased proliferation of basal cells in AR KO mice is associated with increased proliferation of AR KO mice.
the transactivation of AR was confirmed by luciferase assay using MMTV-luciferase including ARE. As shown in Fig. 4A and B, the luciferase activity was increased upon AR transfection, confirming that the transfected AR was functional. Then the expression/activation of several signaling pathways and anti-apoptotic protein in mPrE and Lifeline-basal cells was analyzed by western blot analysis. We found decreases in AKT (AKT1) signal (shown by p-AKT level) and BCL2 expression in both cell lines, whereas no difference was detected in other signals including ERK (MAPK1), WNT (WNT2), and c-MYC (MYC) (Fig. 4C and D). These results suggest that AKT signal and BCL2 are critical in promoting self-renewal/proliferation of these cells.

**Discussion**

Earlier *in vitro* studies using different prostate epithelial cell lines led to different conclusions that androgen might be able to either stimulate or suppress prostate epithelial cell proliferation.
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Using orthotopic injection of CK5+ basal intermediate type of human prostate cancer (PCa) PC3 cells, it was demonstrated that PC3 tumor growth was inhibited when AR was introduced (Niu et al. 2008). The increase in CK5+ cells was also detected when AR was knocked out in prostate epithelium in TRAMP derivatives of prostate epithelium-specific ARKO (pes-ARKO–TRAMP) mice (Niu et al. 2008). These results indicated that AR might play a suppressive role in proliferation of PCa CK5+ cells. Recently, we have also found that the AR role in self-renewal/proliferation of the CK5+ PCa stem/progenitor cells is opposite (suppressor) to the stimulator AR role in proliferation of the CK8+ nonstem/progenitor cells (Lee et al., manuscript submitted). Here, we found similar results showing the suppressor role of AR in self-renewal/proliferation of the normal prostate CK5+ progenitor and basal epithelial cells (Figs 1 and 2).

The androgen/AR roles in differentiation of prostate epithelium have been extensively studied (Ling et al. 2001, Whitacre et al. 2002, Garraway et al. 2003, Berger et al. 2004). When the AR gene was selectively knocked out in epithelial cells, the AR–deficient cells did not terminally differentiate (Simanainen et al. 2007, 2009), which impedes prostate lobes’ growth and function (Wu et al. 2007). However, these results were due to the AR knockout in total epithelium, and up-to-now no basal epithelium-specific AR knockout mouse models have been developed and thus the AR role in basal epithelium has not been studied extensively.

The indication of importance of very low AR expression levels in basal epithelial cells is of significance. The very low levels of AR in the basal compartment may be important in providing a good environment for stem/progenitor cells’ self-renewal/proliferation. However, the AR in basal epithelial cells enables these cells to differentiate into luminal epithelial cells (Fig. 4A), and we have observed higher expression of differentiated cell markers upon AR expression (Fig. 4B and C). So it can be speculated that maintaining the low level of AR in basal epithelial cells might be essential to maintain the balance between progenitor and differentiated luminal epithelial cells and total lack of AR in basal epithelial cells might impede their ability to differentiate.

Taken together, we conclude that the low level of AR expression in basal epithelial cells is essential to determine which axis these cells are destined to: when AR level is low, their proliferation is promoted and may facilitate self-renewal/proliferation of stem/progenitor cells, but when the AR levels increase, their destination is toward differentiation. These circumstances and the driving forces of AR expression toward proliferation or differentiation remain unclear.

After androgen deprivation via castration/androgen deprivation therapy (ADT), in normal prostate tissues, an extensive and rapid cell loss was observed, which appeared to affect mainly the luminal cells. In the process of prostate regeneration, induced by testosterone administration, it was observed that the luminal compartment rapidly regained its normal thickness with keratin expression pattern

Figure 3 AR expression induces differentiation of basal epithelial cells. (A) CK5-Cre/WTAR:ROSA26 and CK5-Cre/floxAR:ROSA26 mice were generated by crossing the CK5-Cre/WT AR and CK5-Cre/floxAR mice with the ROSA26 reporter mice respectively. The mice were killed at 6 weeks of age and GLB1 staining in VP was monitored. GLB1 activity was shown in blue staining and insets represent magnified staining. Arrows indicate luminal cells and arrowheads point out basal cell layer. (B) The mPrE and (C) Lifeline–basal cells were infected with lentivirus carrying AR/vector and grown in three different androgenic concentrations (0, 1, and 10 nM), and the expressions of differentiation markers, PSA, CK8, and Nkx3.1, were examined together with basal marker, CK5, by qRT-PCR analysis.

(Nishi et al. 1988, Altuwaijri et al. 2007, Simanainen et al. 2009, Liao et al. 2010). However, which type of prostate epithelial cells could be suppressed by androgen remained unclear, especially when using in vivo animal models to prove which epithelial cell type of prostate could be stimulated or suppressed by androgen.

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intermediate between basal (CK5+/CK8−/ARlow) and luminal (CK5−/CK8low/ARhigh) cells (Verhagen et al. 1988). In prostate tumors, the intermediate-type cells (CK5low/CK8high/ARlow) were also found increased after ADT (Verhagen et al. 1992, Schalken 2004, Niu et al. 2008). These results also support our hypothesis that the AR expression drives cells into differentiation to generate intermediate cells and into terminal differentiated luminal cells. However, the mechanism by which the AR expression in basal cells and progenitor cells is increased upon castration/ADT is unclear. It was suggested that AR can become transactivated in the low-androgen environment of castration/ADT through a number of different mechanisms, including amplification and mutation of the receptor, cross talk with other signaling pathways, and altered regulation by coregulatory proteins (Lamont & Tindall 2011). Recently, it has been shown that an enhancer in the AR second intron contributes to increased AR expression at low androgen levels (Cai et al. 2011). Therefore, we can speculate that these factors might contribute to the increased level of AR in those cell types and the subsequent differentiation process.

The suppressive role of AR in progenitor cells and basal cells can be applied to PCa therapeutic approach. Much recent evidence indicates that PCa contains stem/progenitor cells that might play essential roles in prostate tumorigenicity and metastasis; therefore, the therapeutic strategies targeting these cells are emerging (Maitland & Collins 2008, Klarmann et al. 2009, Pfeiffer & Schalken 2009, Li & Tang 2011).

Figure 4 AR expression decreases signals related to cell growth/proliferation/survival in mPrE and Lifeline-basal cells. (A) mPrE and (B) Lifeline-basal cells were infected with lentivirus carrying AR or vector and transfected with MMTV-luciferase containing ARE. After transfection, cells were grown in three different androgenic concentrations (0, 1, and 10 nM) and luciferase activity was measured by dual-luciferase assay (Promega) according to the manufacturer’s manual. pRL-TK was used as internal control. (C) mPrE and (D) Lifeline-basal cells were prepared similar to A and B, and the expression levels of the indicated signal molecules were analyzed by western blot analysis.
We speculate that the suppressor role of AR, can be applied to the
development of new therapies to target self-renewal/
proliferation of stem/progenitor cells and the basal cells as we
observed in this study that introducing AR could block self-
renewal/proliferation of these cells. As an introduction of AR
into the cells is not practically possible, we might be able to
develop a therapy targeting the downstream signaling
molecules we revealed in these studies.

**Supplementary data**

This is linked to the online version of the paper at http://dx.doi.org/10.1530/JOE-11-0474.

**Declaration of interest**

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

**Funding**

This work was supported by NIH grants (CA122840 and CA127300) and Taiwan Department of Health Clinical Trial and Research Center of Excellence (DOH99-TD-B-111-004) to China Medical University, Taiwan.

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Received in final form 24 February 2012
Accepted 5 March 2012
Made available online as an Accepted Preprint 5 March 2012