Loss of epithelial AR increase castration resistant stem-like prostate cancer cells and promotes cancer metastasis via TGF-β1/EMT pathway

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Background: Previous study has reported that loss of epithelial androgen receptor (AR) may promote tumor progression and cause TRAMP mouse model die earlier. The detail mechanisms, however, remain unclear.

Methods: Immunohistochemistry assay, Western blot and real-time PCR were used to detect the expression of epithelial and mesenchymal markers. RNA extraction, RT-PCR, quantitative RT-PCR, BrdU incorporation assays, flow cytometry and other experimental techmics were also used in present work.

Results: Decreased expression of epithelial markers (Cytokeratin 8, NKX3.1 and E-cadherin) and increased expression of mesenchymal markers (α-SMA, Vimentin, and N-cadherin) in were found in AR knockout TRAMP tumors. Further investigation indicated that AR signal deprivation is associated with cell morphology transition, high cell mobility, high cell invasion rate and resistance to anoikis in TRAMP prostate tumor cells. Together, these findings implied knockout AR in TRAMP prostate tumor may lead to EMT, which may result in earlier metastasis, and then cause TRAMP mice die earlier. TGF-β1 is responsible for EMT in AR knockout TRAMP tumor cells.

Conclusions: In conclusion, ADT therapy induced hormone refractory prostate cancer may gain the ability of metastasis through cell’s EMT which is a phase of poor differentiation. Anti-EMT drugs should be developed to battle the tumor metastasis induced by ADT therapy.

Keywords: Prostate cancer; androgen receptor; TGF-β; EMT; metastasis

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Introduction

Prostate cancer (PCa) is the second leading cause for cancer mortality of men in USA and the most commonly confirmed non-skin cancer, worldwide (1). Failure to cure patients with tumor metastasis is the most primary reason for death from prostate cancer. In PCa early period, androgens were responsible for tumor development. Therefore, androgen deprivation therapy (ADT) can be implemented to suppress tumor growth (2). However, the final recurrence of PCa is on account of the androgen independence tumor, one of the reasons for tumor high metastasis, replacing the initial most composition of androgen dependence tumor. Currently, there is no effective treatment for cancer metastasis.

For the development and growth of prostate, interactions between androgens and epithelial-mesenchymal is indispensable. Within the prostate glands epithelium and stroma, androgen receptor (AR)-associated androgen signaling pathway is observed (3,4). An absence of functional AR/androgen signaling mice lead to abnormal prostate (5,6). Groundbreaking research on prostate development exhibited that apoptosis, bud formation, epithelial cell identify, proliferation, and ductal branching are identified by specification of stroma AR, not epithelium (7,8). Epithelium AR, however, promotes cell proliferation stimulated by androgen, which has been observed and evidenced by many anaplastic PCa cells research (9-11). ADT has become the key therapy of tumor metastasis or advanced PCa based on this notion and premise.

Our previous study has showed that the AR functions were different in different prostate cells, such as: as tumor inhibitor to depress tumor metastasis in epithelial basal intermediate cells, as survival factor and expansion factor to advance the tumor survival and development. Meantime, it also can cause loss of epithelial AR mice die earlier (12,13). The detail mechanisms, however, remain unclear.

Mesenchymal cells characteristics replaced lots of epithelial cells initial properties in epithelial cells for the reason of Epithelial-mesenchymal transition (EMT) (14). Therefore, it is a key role in order to ascertain tumor cells spread (15). According to current study, it has been observed that EMT-related signaling pathways were negative correlation for AR density (16). Therefore, EMT may be easily occur when AR axis are disrupted during tumor progression.

Using the pes-ARKO-TRAMP Mice, it was observed that knockdown-AR could induce EMT of the pes-ARKO-TRAMP Mice tumor cells. This outcome beyond expectation may explain the reason of failure of androgen deprivation therapy in almost long-term treatment patients. Moreover, it also can provide an urgent need to develop new approaches that gives castration therapy at earlier stages and prevents EMT to treat prostate cancer.

Methods

Cell culture and reagents

The LNCaP human early androgen-dependent prostate cancer cell line and CWR22rv1 (CRL-2505) castration resistant prostate cancer cell line were purchased from the China Type Culture Collection, cultured in recommended medium with supplement of 10% fetal bovine serum (FBS; Gibco, California, USA) and incubated at 37 °C within humidified atmosphere including 5% CO2. Meantime, 25 U/mL penicillin and 25 μg/mL streptomycin were added to them.

Generation of pes-ARKO-TRAMP mice

To product TRAMP-floxed female mice, mice with the conditional AR allele (floxed AR, C57BL/6) were used to mate with TRAMP mice. Then to generate pes-ARKO-TRAMP male and littermates Wt-AR-TRAMP mice, ARR2PB2-Cre male mice (C57BL/6N) were used to mate with TRAMP-floxed AR female mice. TRAMP (FVB) were purchased from Jackson Laboratory and Probasin Cre (pb-Cre)(C57BL/6) mice from NCI. Using R26R/β-Gal transgenic female mice were mated with pes-ARKO-TRAMP male mice to generate pes-ARKO-TRAMP β-Gal transgenic mice. After 12 weeks feeding, mice were killed and then fresh tissue samples were exteriorized to conduct β-Gal staining. The microscopy was used to identify the β-Gal-positive cells.

Tumor orthotopic transplantation in nude mice

8-wk nude mice without thymus were selected to orthotopically implant CWR22rv1-AR+/+ cells and CWR22rv1-AR+/- cells in their prostate anterior lobe by opening abdomen under anesthesia and sterile condition. We choose 25-gauge needle to inject the CWR22rv1-AR+/+ cells and CWR22rv1-AR+/- cells into one anterior lobe. The cells were calculated and suspended in 50 μL Matrigel with 10⁶ cells/μL and then sutured the abdomens. 12 weeks later, we killed the mice, incised the xenograft and...
metastatic tumors, followed by fixing and embedding the samples in paraffin for future research.

**RNA extraction, RT-PCR, and quantitative RT-PCR**

TRIzol reagent (Invitrogen) was used to extract total RNA from tissues and cells, under manufacturer’s instructions. Each isolated RNA sample (5 μg) was reverse-transcribed into cDNA (20 μL) using a miScript II RT Kit (Invitrogen) according to the manufacturer’s instruction. After diluting the cDNA 10 times with water, we took out 2 μL cDNA for PCR amplification template ABI 7500 System (Applied Biosystems, USA) was applied to carry out the RT-PCR reactions. Beacon Designer 2 software were used to design primers. The β-actin was as control in each PCR reaction. According to the β-actin calculated threshold (CT) values levels, we deduced the samples CT values. Agarose-gel electrophoresis informed the absence of nonspecific amplification products. All the assays’ procedures repeat in triple.

**Immunohistochemistry staining**

We sliced the paraffin-embedded samples and stained the slice. The anti-CK8 (Abcam), anti-CK5 (Covance), the rabbit anti-AR (C19) (Santa Cruz Biotechnology), rabbit anti-Tag (Santa Cruz), rabbit anti-Tag (Santa Cruz) was employed as primary antibodies. Afterwards, the biotinylated secondary antibody was used to recognize primary antibody followed by using peroxidase substrate DAB kit (Vector) to visual. Image J software approximately quantified positive signals strength.

**BrdU incorporation assays**

We fabricated Bromodeoxyuridine (BrdU) (sigma) solution (10 mg/mL). Beginning with 24 h before sacrifice, mice were injected BrdU solution (10 μg/g) by intraperitoneal injection every 6 h. After incising, fixing and embedding the tumors in paraffin, we marked samples by using BrdU Staining Kit (Zymed) according to recommend protocol.

**Obtain selected prostate cells by Laser-Capture microdissection**

Samples were stored in O.C.T. placing in −80 °C condition as standby. For separating the epithelium and stroma cells, we first cut 5-m specimens sections, placed them on slides. Subsequently, HistoGene LCM Frozen Section Staining Kit (Arcturus) was immediately used to stain the specimens’ sections according to manufacturer’s recommend. Prostatic epithelium and stroma cells were separated into different capes by laser transferring under the Pixcell II LCM system. PicoPure RNA Isolation Kit (Arcturus), a kind of RNA extraction buffer, was used to merge capes and extract RNA according to instruction after microdissecting. Then, we amplified and reverse transcribed the RNA, and analyzed genes’ changes by qRT-PCR.

**Flow cytometry**

Trypsin-EDTA were used to digest the cells, separate cells (1×10⁶) for each other, and then cells were fixed by 70% ethanol at 4 °C condition. 12 h later, we centrifuged the cells treated before during 7 min by 1,000×g at 4 °C condition, suspended cells in PBS with 0.05 mg/mL Rnase A(Sigma), afterwards incubated cells during 30 min at room temperature. 30 min later, propidium iodide (10 mg/mL) were performed to stain the cells. Then we used flow cytometry (FACSCalibur, BD Company) to analyzed 10,000 cells and the result were analyzed by ModFit software (Verity Software House, Inc.).

**Statistics**

We showed the data in the mean + standard deviation (SD). Two-sided Student’s t test was used to compare two groups and Kaplan-Meier analysis and log-rank tests were performed to analyze Survival curves.

**Results**

**AR loss led to the expression changes of EMT markers**

We found ARKO prostate tumors lost more cytokeratin markers than the AR Wt prostate tumors. From 16 to 24 wks, all prostate tumors gradually lost CK8 expression. Although pes-ARKO-TRAMP prostate tumors contains more basal intermediate cell population in 16 wks than Wt TRAMP tumor in the same age, ARKO tumors finally decreased both CK5 and CK8 staining in the later tumor stage (Figure 1A). Immunohistochemical staining showed tumors from pes-ARKO-TRAMP prostate expressed less E-cadherin than tumors from Wt-TRAMP in 16 wks, 20 and 24 wks (Figure 1B). But in the 24 wks prostate tumors, pes-ARKO-TRAMP tumors expressed higher
Figure 1 Loss of epithelial AR in pes-ARKO-TRAMP prostate led to higher expression of EMT markers in prostate tumor. (A) ARKO prostate tumors lost more cytokeratin markers than the AR Wt prostate tumors. Prostate samples from pes-ARKO-TRAMP and Wt-TRAMP were double stained wnt1, CK5 (green) and CK8 (red). From 16 to 24 wks, all prostate tumors gradually lost CK8 expression. Although pes-ARKO-TRAMP prostate tumors contains more basal intermediate cell population (CK5/CK8 double stained yellowish cells) in 24 wks (right upper panel) than Wt TRAMP tumor in the same age, ARKO tumors finally decreased both CK5 and CK8 staining in the later tumor stage (right lower panel). Figure 1A (Wt TRAMP 16 ws) was quoted from: Niu Y, Altuwaijri S, Lai KP, et al. Androgen receptor is a tumor suppressor and proliferator in prostate cancer. Proc Natl Acad Sci U S A 2008;105:12182-7—the first picture of Figure 3C. (B) E-cadherin, as an important marker of EMT, was Immunohistochemical staining in 16, 20 and 24 wks Wt and ARKO TRAMP tumors. Tumors from pes-ARKO-TRAMP prostate expressed less E-cadherin than tumors from Wt-TRAMP. (C) N-cadherin (green), which was mainly expressed in the stroma of 16 wks Wt-TRAMP prostate (left, upper panel), was also expressed in the epithelial of 16 wks ARKO-TRAMP prostate (right, upper panel). And in the 24 wks prostate tumors, pes-ARKO-TRAMP tumors expressed higher level of N-cadherin than Wt-TRAMP tumors. (D) Integrin β1 and MMP9 staining were also stronger in the pes-ARKO-TRAMP tumors than that in Wt-TRAMP tumors. (E) Using Western-blot assay, epithelial markers as NKX3.1, cytokeratin (Pan), and E-cadherin were found declined, and mesenchymal markers containing Vimentin and N-cadherin were found increased in pes-ARKO-TRAMP tumors compared with same aged Wt-TRAMP tumors. (F) The relative RNA expression levels of integrinβ1, Wnt1, Wnt11, FZD, snail1, snail2, MMP2, MMP9, N-cadherin and E-cadherin were detected by Real-time PCR.
levels of N-cadherin than Wt TRAMP tumors (Figure 1C). Integrin β1 and MMP9 staining were also stronger in the pes-ARKO-TRAMP tumors than that in Wt TRAMP tumors (Figure 1D); Using Western-blot assay, epithelial markers as NKKX3.1, cytokeratin (Pan), and E-cadherin were found decline, and mesenchymal markers including Vimentin and N-cadherin were found increase in pes-ARKO-TRAMP tumors compared with same aged Wt TRAMP tumors (Figure 1E). The relative RNA expression levels of integrinβ1, Wnt1, Wnt11, FZD, snail1, snail2, MMP2, MMP9, N-cadherin and E-cadherin were identical with Western-blot assay results (Figure 1F). These results indicate that high expression of mesenchymal markers appeared in pes-ARKO-TRAMP prostate when epithelial AR lost.

**AR loss increased EMT phenotypes in term of cell morphology, detachment, motility and invasion**

We subsequently found that after 48hrs of cell culture, the primary cells from ARKO-TRAMP prostate tumor were clustered with elongated, spindle-like cells, while the primary cells from Wt-TRAMP prostate tumor were scattered with round and polygonal-like cells (Figure 2A). Meanwhile, the cell invasion rate was higher in the primary cultured ARKO cells than that in Wt cells (Figure 2B). We confirmed that cell invasion was significantly increased in AR knockdown CWR22rv1 (marked as CWR22rv1-AR+/-) cells comparing with their parental cells (marked as CWR22rv1-AR+/+) (Figure 2C). We also detected Cell motility in a wound healing assay, both CWR22rv1-AR+/- and CWR22rv1-AR+/+ cells were grown to nearly confluency, and monolayers were scratched with a pipette tip. Cell motility was photographed 24hr after wound scratch. We found that the migration of the AR knockdown CWR22rv1-AR+/- cells was higher than CWR22rv1-AR+/+ cells, and 10 nM DHT suppressed the migration of both cells (Figure 2D). And CWR22rv1-AR+/- cells were easier to be detached by EDTA than CWR22rv1-AR+/+ cells, and 10 nM DHT suppressed the detachment of both cells (Figure 2E). 10 nM DHT protected LnCaP cells from detachment treated by EDTA (Figure 2F). Our data confirmed that loss of AR increase EMT phenotypes in term of cell morphology, detachment, motility and invasion.

**AR loss resulted in fewer anoikis cells**

A Poly-Hema coated plate or a tissue culture control plate was performed to seed both CWR22rv1-AR+/- and CWR22rv1-AR+/+ cells (50,000 cells/well). Culturing cells for 24 hours, then we determined cell activity by Calcein AM (green), while stained death anoikis-like cell by EthD-1 (red) (Figure 3A,B). The anoikis cells were counted by flowcytometry and were subsequently quantitated shown in (Figure 3C,D) LNCaP cells with or without 10 nM DHT treatment were also submitted to detect the anoikis by flowcytometry. The quantitative data was shown in (Figure 3E). These data were shown that decreased expression and function of AR in CWR22rv1 cells and LnCaP cells resulted in fewer anoikis cells.

**AR loss developed larger xenograft tumors and early lymph nodes metastasis**

Both CWR22rv1-AR+/- cells and CWR22rv1-AR+/+ were orthotopically transplanted into the prostate anterior lobe of nude mice. 12 wks following implantation, mice were sacrificed, and tumors and lymph nodes were harvested (Figure 4A). The xenograft primary tumors in the prostate were weighted (Figure 4B) and the weight of the metastatic lymph nodes was also scaled (Figure 4C). We found that AR knockdown tumors from CWR22rv1-AR+/- xenografts and lymph nodes expressed less E-cadherin than that of CWR22rv1-AR+/+ xenografts and lymph nodes (Figure 4D). Snail1 expression was higher in CWR22rv1-AR+/- xenograft tumors (Figure 4E). Mesenchymal markers of α-SMA, N-cadherin, vimentin, MMP9, and integrin-α5 were significantly increased in CWR22rv1-AR+/- xenograft tumors comparing with those in CWR22rv1-AR+/+ xenograft tumors (Figure 4F). Epithelial markers of CK8, CK5, E-cadherin, NKKX3.1 and PSA, except PSP94, were significantly decreased in CWR22rv1-AR+/- xenograft tumors comparing with those in CWR22rv1-AR+/+ xenograft tumors (Figure 4G). Signaling difference, which may involve in EMT, was also investigated and compared (Figure 4H). These observations implicated that Snail1, α-SMA, N-cadherin, vimentin, MMP9, and integrin-α5 expression increased in response to AR loss, possibly facilitated CWR22rv1-AR+/- cells tumor metastasis.

**AR loss induced EMT may be responsible for early cancer metastasis**

Subsequently, we found that at 20 wks, pes-ARKO-TRAMP mice developed larger tumors than the Wt-TRAMP mice. The large tumors of pes-ARKO-TRAMP mice began to
Figure 2 Loss of AR increases EMT phenotypes in terms of cell morphology, detachment, motility and invasion. (A) After 48 hrs of primary culture, the primary cells from ARKO-TRAMP prostate tumor are clustered with elongated, spindle-like cells, while the primary cells from Wt-TRAMP prostate tumor are scattered with round and polygonal-like cells. (B) Meanwhile, the cell invasion rate is higher in the primary cultured ARKO cells than that in Wt cells (C) We confirmed that cell invasion was significantly increased in AR knockdown CWR22rv1 (marked as CWR22rv1-AR+-) cells comparing with their parental cells (marked as CWR22rv1-AR+/+). (D) Cell motility was detected in a wound healing assay. Both CWR22rv1-AR- and CWR22rv1-AR+/+ cells were grown to nearly confluency, and monolayers were scratched with a pipette tip. Cell motility was photographed 24 hrs after wound scratch. The migration of the AR knockdown CWR22rv1-AR- cells was higher than CWR22rv1-AR+/+ cells, and 10 nM DHT suppressed the migration of both cells. (E) And CWR22rv1-AR- cells were easier to be detached by EDTA than CWR22rv1-AR+/+ cells, and 10 nM DHT suppressed the detachment of both cells. (F) 10 nM DHT protected LNCaP cells from detachment treated by EDTA.
Figure 3 Decreasing the expression and function of AR in CWR 22rv1 cell and LNCaP cells resulted in fewer anoikis cells. (A) A Poly-Hema coated plate or a tissue culture control plate was performed to seed both CWR22rv1-AR+/+ and CWR22rv1-AR+/- cells (50,000 cells/well). Culturing cells for 24 hours, then we determined cell activity by Calcein AM (green), while stained death anoikis-like cell by EthD-1(red). (B) The anoik1s cells were counted by flowcytometry and were subsequently quantitated shown in (C). (D) LNCaP cells with or without 10 nM DHT treatment were also submitted to detect the anoikis by flowcytometry. The quantitative data was shown in (E).
Figure 4 Androgen receptor signaling blocking may induce EMT and metastasis in vivo and in vitro. (A) Both CWR22rv1-AR+/+ cells and CWR22rv1-AR+-/ cells were orthotopically implanted into the anterior lobe of nude mice. 12 wks following implantation, mice were sacrificed, and tumors and lymph nodes were harvested. (B) The xenograft primary tumors in the prostates were weighted. (C) The weight of the metastatic lymph nodes was also scaled. (D) Samples from the xenograft tumors and metastatic lymph nodes of each group were double stained by androgen receptor (red) and E-cadherin (green). AR knockdown tumors from CWR22rv1-AR+-/ xenografts and lymph nodes expressed less E-cadherin than that of CWR22rv1-AR+/+ xenografts and lymph nodes. (E) Snail1 expression was higher in CWR22rv1-AR+-/ xenograft tumors (F) Mesenchymal markers of α-SMA, N-cadherin, vimentin, MMP9, and integrin-α5 were significantly increased in CWR22rv1-AR+-/ xenograft tumors comparing with those in CWR22rv1-AR+/+ xenograft tumors. (G) Epithelial markers of CK5, CK8, E-cadherin, NNX3.1 and PSA, except PSP94, were significantly decreased in CWR22rv1-AR+/+ xenograft tumors comparing with those in CWR22rv1-AR+-/ xenograft tumors. (H) Signaling difference, which may involve in EMT, was also investigated and compared.
develop metastatic tumors in lymph node from 20 wks, which were larger and more lesion foci than those of Wt-TRAMP mice (Figure 5A). The tumor samples from 20 wks pes-ARKO-TRAMP and Wt-TRAMP prostate were double stained by T-antigen (Red) and N-cadherin (green). N-cadherin expression was increased in ARKO tumors (Figure 5B). The increased α-SMA, decreased AR and CK8 expressions were detected by Real-time PCR in ARKO tumors and lymph nodes (Figure 5C). Illustrating the evolitional events in which the tumor cells keep transiting (Figure 5D). Our data showed that AR loss induced EMT transition may be responsible for the early metastasis of the pes-ARKO-TRAMP tumors again.

**The mechanism may involve in the AR loss triggered EMT**

In view of the widely acknowledged role of TGF-β as a potent EMT inducer, we examined TGF-β1 signaling. We found that of TGF-β1, phospho-Smad2/3, phospho-AKT, phospho-JNK, phospho-GSK3β, β-catenin and snail1 proteins expression increased by IHC staining and Western-blot assay in ARKO mice than that in WT mice (Figure 6A,B). Furthermore, we found that TGF-β1 signaling through phosphor-AKT to activate snail1 and to result in EMT, which is characterized by decreased E-cadherin and increased α-SMA expression, was blocked by PI-3K inhibitor LY294002 (Ly) (Figure 6C). Subsequently, we used Real-time PCR to test the concept that the TGF-β1 signaling is important for the EMT transition of CWR22rv1 cells. In the left panel, without any treatment, AR knockdown CWR22rv1-AR+/− cells expressed more mesenchymal markers, such as α-SMA, N-cadherin, MMP9 and integrin β1, than those of CWR22rv1-AR+/+ cells. 10ng/ml TGF-β1 treatment further induced the mesenchymal markers expression in CWR22rv1-AR+/− cells (middle panel). By using TGF-β1 receptor II siRNA (marked as TGF-bRIIIsi), we confirmed that the expression of these mesenchymal markers was significantly reduced in CWR22rv1-AR+/− cells (right panel) (Figure 6D). In addition, the Real-time PCR results showed the TGF-β1 signaling and blocking TGF-β1 signaling significantly changed the expression of epithelial markers, such as CK8, E-cadherin, PSA, and Nkx3.1, in CWR22rv1-AR+/− cells (Figure 6E). The Real-time PCR results showed the TGF-β1 effects on the Wnt, snail, and ZEB signaling (Figure 6F). TGF-β1 induced and TGF-bRIIIsi suppressed CWR22rv1-AR+/− cells invasion in CWR22rv1-AR+/− cells (Figure 6G). A clear consequence of the results here presented is that TGF-β1 induced EMT sensitizes prostate tumor cells to respond to AR loss.

The relevant mechanisms are illustrated in Figure 7. Androgen-binding AR inhibits TGF-β-related Pathway. However, through binding signal molecules to TGF-β receptors (TGF-β R1 and TGF-β R2), the receptors are phosphorylated activating the SMAD signaling pathway and the non-SMAD signaling pathway. The SMAD signaling pathway begins with the co-SMAD gaining phosphorylation from TGF-β receptor, and then binding to SMAD4 to form SMAD complex. The complexes enter the nucleus and bind the transcription factors of Snail genes promoting the expression of Snail1. Because Snail1 is the key protein of EMT, it can induce the production of EMT. Non-SMAD pathways include TAK1 signaling and PI-3k signaling pathway, etc. The TAK1 signaling pathway starts with the phosphorylation of TAK1 factors, which lead to JNK phosphorylation acting on c-JUN to regulate the transcriptional activity of cells and promote cell proliferation and inhibit cells apoptosis. And PI-3K signaling pathway is caused by PI-3K factors binding phosphoric acid groups which lead to phosphorylation of AKT. Inhibition of GSk3β and β-catenin Phosphorylation by phospho-AKt reduces the degradation of β-catenin, and then β-catenin converging in nucleus promotes cell proliferation and inhibits apoptosis by binding transcription factors. Furthermore, phospho-AKt factors can also lead to EMT by directly promoting the expression of EMT-related key proteins.

**Discussion**

EMT is one mark of epithelial cancer progression. Tumor cells lose cell-cell junctions and polarity and acquire many mesenchymal characteristics, the matrix invasion making cell metastasis (17,18). In this research, we have provided a lot of evidence that AR signal deprivation in TRAMP prostate tumor lead to epithelial-mesenchymal transition. We generated pes-ARKO-TRAMP mice that spontaneously developed prostate cancers without AR in cancer epithelial cells. We found that loss of epithelial AR in pes-ARKO-TRAMP mice prostate led to the epithelial markers decrease (E-cadherin, Cytokeratin 8 and Nkx3.1) and mesenchymal markers increase (Vimentin, α-SMA, and N-cadherin). Compared with the loss of E-cadherin, N-cadherin up-regulating expression has been previously described during the progression of various cancer types and is thought to be necessary for tumor cells to gain invasive
Figure 5 AR loss induced EMT transition may be responsible for the early metastasis. (A) At 20 wks, pes-ARKO-TRAMP mice developed larger tumors than the Wt-TRAMP mice. The large tumors of pes-AR KO-TRAMP mice began to develop metastatic tumors in lymph node from 20 wks, which were larger and more lesion foci than those of Wt-TRAMP mice. The last picture of Figure 5A was quoted from: Niu Y, Altuwaijri S, Yeh S, et al. Targeting the stromal androgen receptor in primary prostate tumors at earlier stages. Proc Natl Acad Sci U S A 2008;105:12188-93—the second picture of Figure 5A. (B) The tumor samples from 20 wks pes-ARKO-TRAMP and Wt-TRAMP prostate were double stained by T-antigen (Red) and N-cadherin (green). N-cadherin expression was increased in ARKO tumors (C) The increased α-SMA, decreased AR and CK8 expressions were detected by Real-time PCR in ARKO tumors and lymph nodes (D) Illustrating the evolutionary events in which the tumor cells keep transiting.
Figure 6 The mechanisms which may involve in the AR loss triggered EMT. (A) IHC staining of TGF-β, phospho-Smad2/3, phospho-AKT, phospho-JNK, phospho-GSK3β, β-catenin and snail1 were carried out. (B) The expression of TGF-β, phospho-Smad2/3, phospho-ERK 1/2, phospho-AKT, phospho-GSK3β, snail1 and α-SMA were confirmed in Western blot assay. (C) TGF-β1 signaling through phosphor-AKT to activate snail1 and to result in EMT, which is characterized by decreased E-cadherin and increased α-SMA expression, was blocked by PI3K inhibitor LY294002 (Ly). (D) Using Real-time PCR to test the concept that the TGF-β1 signaling is important for the EMT transition of CWR22rv1 cells. In the left panel, without any treatment, AR knockdown CWR22rv1-AR+/- cells expressed more mesenchymal markers, such as α-SMA, N-cadherin, MMP9 and integrin β1, than those of CWR22rv1-AR+/+ cells. 10 ng/mL TGF-β1 treatment further induced the mesenchymal markers express in CWR22rv1-AR+/- cells (middle panel). Using TGF-β1 receptor II siRNA (marked as TGF-bRllsi), the expression of these mesenchymal markers was significantly reduced in CWR22rv1-AR+/- cells (right panel) (E) The Real-time PCR results to show the TGF-β1 signaling and blocking TGF-β1 signaling significantly changed the expression of epithelial markers, such as CK8, E-cadherin, PSA, and Nkx3.1, in CWR22rv1-AR+/- cells (F) The Real-time PCR results to show the TGF-β1 effects on the Wnt, snail, and ZEB signaling. (G) TGF-β1 induced and TGF-bRllsi suppressed CWR22rv1-AR+/- cells invasion in CWR22rv1-AR+/- cells.
Figure 7: Androgen-binding AR inhibits TGF-β-related Pathway. Through binding signal molecules to TGF-β receptors (TGF-βR1 and TGF-βR2), the receptors are phosphorylated to activate the SMAD signaling pathway and the non-SMAD signaling pathway. The SMAD signaling pathway begins with the co-SMAD gaining phosphorylation from TGF-β receptor, and then binding to SMAD4 to form SMAD complex. The complex enters the nucleus and binds the transcription factor of Snail gene promoting the expression of Snail1. Because Snail1 is the key protein of EMT, it can induce the production of EMT. Non-SMAD pathways include TAK1 signaling and PI-3K signaling pathway, etc. The TAK1 signaling pathway starts with the phosphorylation of TAK1 factor, which leads to JNK phosphorylation acting on c-JUN to regulate the transcriptional activity of cells and promote cell proliferation and inhibit cells apoptosis. And PI-3K signaling pathway is caused by PI-3K factor binding phosphoric acid which leads to phosphorylation of AKT. Inhibition of GSk3β and β-catenin Phosphorylation by phospho-AKt reduces the degradation of β-catenin, and then β-catenin converging in nucleus promotes cell proliferation and inhibits apoptosis by binding transcription factors. Furthermore, phospho-AKt factors can also lead to EMT by directly promoting the expression of EMT-related key proteins.
properties (19). Significantly, the functional importance of decreased E-cadherin levels has also been demonstrated in PCA cells with its inverse correlation with cellular motility and protease expression (20, 21). Our results unexpectedly imply that AR signal deprivation may promote escape of TRAMP PCA cells in primary site and metastasis. This is according with a recent report revealing that EMT was easily induced by the condition of low intracellular AR levels (22).

Subsequently, we also found the diameter of pes-ARKO-TRAMP mice tumor or lymph node metastatic tumors were significantly larger than their WT TRAMP littermates. And orthotopic implantation of AR knockdown CWR22rv1-AR+/- cells developed larger xenograft tumors and bigger metastatic lymph nodes than those from parental CWR22rv1-AR+/- cells. Mesenchymal-cell markers gain and epithelial-cell markers loss, particularly at the leading edge or invasive front of solid tumors, has been observed in human tumor specimens and is associated with tumor progression to metastasis (23). Our findings indicated that AR signal deprivation had effect on promoting the progression and metastasis of tumor, and support for our opinion.

Epithelial tumor cells lose many characters and meanwhile acquire protein mesenchymal-cell markers and signaling activities associated with mesenchymal cells facilitating migration and survival in an anchorage-independent environment and ultimately metastasis (23, 24). Interestingly, concomitant to the molecular alterations occurred on absence of E-cadherin and gain of Vimentin, N-cadherin, and ^\alpha$-SMA, we also found that AR signal deprivation is associated with cell morphology transition, high cell mobility, high cell invasion rate and resistance to anoikis in TRAMP prostate tumor cells. These data suggest that AR signal deprivation plays an important role on the changes of prostate epithelial cell morphology and promotes metastasis of tumor.

We present evidence for molecular mechanism that contributes to explain our finding, TGF-$\beta$1 signaling and Wnt signaling, as well as AKT and snails, are responsible for EMT change in AR knockout TRAMP tumor cells. TGF-$\beta$, a kind of multifunctional factor, regulates cellular responses in broad range (25). TGF-$\beta$ binds to type II and type I serine/threonine kinase receptors and transmits intracellular signals. Smads are the major transducer of TGF-$\beta$-signaling; Smad2 and Smad3 are phosphorylated by the TGF-$\beta$ type I receptor and form complexes with Smad4. These complexes accumulate in the nucleus and regulate transcription of target genes (26). The mitogen-activated protein kinases' [MAPK; p38 MAPK, c-jun NH2-terminal kinase (JNK), and extracellular signal-regulated kinase (ERK)-1/2] activation modulated Smad2/3 or other downstream transcription factors activity to increase TGF-$\beta$ signaling complexity (27-31) (including Snail1, Snail2, Notch, and others). The interference between TGF-$\beta$ signaling and androgen has been elaborated (32).

A direct interaction between AR and TGF-$\beta$1 has been causally implicated in other human tumors (33). Androgens can inhibit TGF-$\beta$1-induced transcriptional activity in prostate cancer cells (34), an interaction that is regulated by AR-associated protein 55 (ARA55/Hic-5; LIM protein superfamily). Overexpression of ARA55 inhibits TGF-$\beta$-mediated up-regulation of SMAD transcriptional activity in rat prostate epithelial cells, as well as human prostate cancer cells, via an interaction between ARA55 and SMAD3 mediated through the MH2 domain of SMAD3 and the C terminus of ARA55 (35). Our results found the AR loss result in TGF-$\beta$, p-Smad2/3, p-AKT, p-JNK, p-GSK3$\beta$, Snail1 proteins expression increased. In several cancer cells, E-cadherin expression can be repressed by Snail regulated by TGF-$\beta$/Smad pathway (36-38). Our outcomes show that after knockdown AR, TGF-$\beta$1 signaling through phosphor-AKT to activate snail1 and to result in EMT, which is characterized by increased ^\alpha$-SMA expression and decreased E-cadherin, was blocked by PI-3 K inhibitor LY294002 (Ly). We also showed using TGF-$\beta$1 receptor II siRNA (marked as TGF-$\beta$-bRIIsi), the expression of these mesenchymal markers (^\alpha$-SMA, N-cadherin, MMP9 and integrin $\beta$1) was significantly reduced in CWR22rv1-AR+/- cells. The TGF-$\beta$1 signaling and blocking TGF-$\beta$1 signaling significantly changed the expression of epithelial markers, such as CK8, E-cadherin, PSA, and Nkx3.1, in CWR22rv1-AR+/- cells. Our data also showed the TGF-$\beta$1 effect on the Wnt and ZEB signaling. These results suggest that knockdown AR can independently induce EMT, potentially bypassing the effect elicited by TGF-$\beta$.

Conclusions

Taken together, our data reveal that PCA cell metastatic ability can be promoted by absence of AR by modulating EMT. This research put forward innovation view for the androgen-mediated EMT effect. ADT therapy induced hormone refractory prostate cancer may gain the ability of metastasis through cell's EMT which is a phase of poor differentiation. Anti-EMT drugs should be developed to
battle the tumor metastasis induced by ADT therapy. The signaling pathways which involved in EMT progress need further elucidation.

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Footnote

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/tau.2020.03.02). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was approved by research medical ethics committee of the Second Hospital of Tianjin Medical University (Grant No.KY2016K004).

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