Abstract. With increases in the mortality rate and number of patients with prostate cancer (PCa), PCa, particularly the advanced and metastatic disease, has been the focus of a number of studies globally. Over the past seven decades, androgen deprivation therapy has been the primary therapeutic option for patients with advanced PCa; however, the majority of patients developed a poor prognosis stage of castration resistant prostate cancer (CRPC), which eventually led to mortality. Due to CRPC being incurable, laboratory investigations and clinical studies focusing on CRPC have been conducted worldwide. Clarification of the molecular pathways that may lead to CRPC is important for discovering novel therapeutic strategies to delay or reverse the progression of disease. A sustained androgen receptor (AR) signal is still regarded as the main cause of CRPC. Increasing number of studies have proposed different potential mechanisms that cause CRPC, and this has led to the development of novel agents targeting the AR-dependent pathway or AR-independent signaling. In the present review, the major underlying mechanisms causing CRPC, including several major categories of AR-dependent mechanisms, AR bypass signaling, AR-independent mechanisms and other important hypotheses (including the functions of autophagy, PCa stem cell and microRNAs in CRPC progression), are summarized with retrospective pre-clinical or clinical trials to guide future research and therapy.

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1. Introduction
Prostate cancer (PCa) is the most frequently diagnosed cancer in men from the Western world (1). The incidence of PCa has exceeded lung cancer, and is the second leading cause of mortality in malignant tumors in males (1). The number of novel diagnosed cases in 2016 was 180,890 in the USA alone (2). In China, the incidence of PCa is increasing year by year with the change in living habits and the aging population (3). In 2015, there were 60,300 novel cases of PCa and 26,600 patients with PCa succumbed (3). Androgen deprivation therapy (ADT) has been the primary treatment option for male patients with advanced symptomatic PCa for almost seven decades and generally results in decreased prostate specific antigen (PSA) levels. By contrast, following a median of 18-24 months of endocrine therapy, almost all patients progressed to a poor prognosis stage -castration resistant prostate cancer (CRPC), which was previously called hormone-refractory prostate cancer. However, the stage of CRPC is not hormone-refractory based on the understanding that the androgen axis continues to be activated and promotes the growth of CRPC (4). Almost all patients succumbed due to no existing treatment to control or cure the mortality-causing CRPC; therefore, the mechanisms underlying the initiation and development of CRPC remains challenging to decipher, which requires focus from international academic research, and will help to improve the survival and the quality of life of patients with PCa. In the present review, the current studies regarding the molecular and cellular mechanisms leading to CRPC are summarized for the development of future molecularly targeted therapies for PCa.

2. Definition of CRPC
CRPC refers to the continuous progression of PCa following ADT. In recent years, the definition of castration resistance
has changed substantially. With the emergence of novel CRPC therapy, it is particularly important to determine the exact definition of castration resistance. In March 2015, Gallen gathered 41 experts in the field from 17 countries and regions around the world, and held the first session of the advanced PCa Consensus Conference (5). Almost all experts in the session agreed that the diagnosis of CRPC should meet the following 2 conditions: i) The serum testosterone level of the castrated is <1.7 nmol/l; and ii) indicating biochemical progression. Biochemical progression is characterized that the PSA expression levels have increased twice in a row from an interval of 1 week or ≥3 consecutive measurements with the lowest value increased >50% and ≥2 g/l, and ≥2 increases in novel lesions based on bone scanning or soft tissue lesions with the corresponding evaluation criteria of the solid tumor. Currently, only symptom progression is not sufficient to diagnose CRPC. The understanding of PCa biology and underlying disease resistance mechanisms has grown over the last few decades, and has been translated into improved and clinically meaningful treatment strategies for males with advanced PCa (6). Understanding the mechanisms underlying the progression of PCa from hormone sensitive to castration resistant is the key to develop future therapy. In the present review, a number of the major underlying mechanisms leading to castration resistance are discussed.

3. Androgen receptor (AR)-dependent mechanisms underlying the resistance leading to CRPC

AR. The growth and survival of PCa tumors depend on androgens, the male sex steroid hormones, of which testosterone and dihydrotestosterone (DHT) are the principal members that activate and bind to the AR (7). High expression of AR occurs in the majority of patients with CRPC, and the expression of AR-regulated genes recovered in tumors after patients received ADT implies that AR transcription activity is activated again (7).

AR, as a member of the steroid hormone super family, has four functional motifs, including the ligand-binding domain (LBD), DNA-binding domain (DBD), hinge region and the amino-terminal domain (N-terminal domain, NTD) that contains the phosphorylation sites essential for transcriptional activity (8) (Fig. 1). When AR binds with a steroid ligand, including DHT, it can prevent ubiquitination and proteasome degradation by interacting with heat-shock proteins (HSP) (9). The AR is phosphorylated prior to dimerization, which leads to a conformational change, which displaces the HSP upon ligand binding (10). Dimerized ligand-binding complex will then be transported into the nucleus (10). Ligand binding is vital to dimerization and translocation of AR to the nucleus. Following AR being trafficked into the nucleus, it may mediate transcription and various growth signaling pathways, including cellular proliferation or anti-apoptosis and androgen-regulated genes, including PSA, through binding to androgen response elements in the promoter or enhancer regions of DNA (11). A number of studies revealed that the AR may be transported into the nucleus through the microtubule complex (12). Evacuation of the ligand in androgen-responsive prostate cells results in an exportation of the AR from the nucleus (9).

In the process of hormone-responsive PCa to CRPC, the AR and its cross-signaling pathways exist a number of changes with various forms, which provides an important clue for the exploration of the mechanisms underlying CRPC.

Amplification of the AR gene and overexpression of the AR protein. AR gene amplification and overexpression of the AR protein have been frequently observed in clinical studies. The AR gene amplification leading to an overexpression of the AR protein is the most common genetic change among patients with CRPC (13). Of the patients with CRPC, >80% of them had significant augmentation of AR mRNA and protein levels (14). By contrast, AR gene amplification was rarely observed in untreated PCa (14). Research determined that the AR gene had a high frequency of amplification in circulating tumor cells (CTCs) of patients with CRPC (15). Fluorescence in situ hybridization studies utilizing tissue microarrays discovered that AR gene amplification is present in only 2% of the primary PCa tumor samples and none of the samples of benign prostate hyperplasia, compared with 23.4% of CRPC tumors (16). Using reverse transcription-polymerase chain reaction (RT-PCR) analysis, it was confirmed that the AR gene amplification is indeed reflected at the mRNA level, where the expression of AR mRNA in CRPC tumors with AR amplification was increased two fold, compared with those CRPC tumors without AR amplification (17). On the other hand, increased AR was an unique gene expression alteration that was sustained in different CRPC xenograft tumor samples (18).

Elevated AR protein levels are also linked to CRPC. Using multiple isogenic tumor xenograft models, it was demonstrated that the expression of the AR protein increased in recurrent tumor samples, compared with paired androgen-sensitive samples (18,19). Notably, in a CWR22 xenograft model that imitates the transition from androgen-sensitive to recurrent growth, the expression of the AR protein was gradually reduced during the 120-day castration period, and then regrowth occurred in recurrent tumors (19). In addition, not only was the gene amplification and elevated mRNA expression directly augmenting the AR protein levels, increased protein half-life can also contribute to the elevated levels of AR protein in CRPC (20). Through gene amplification and enhanced transcriptional induced overexpression of AR causes AR hypersensitive to low level of androgen (21). Furthermore, it can be concluded that under the lowest concentration of androgens, tumor cells proliferate continuously (22), which indicated that overexpression of AR may contribute to the development of CRPC formation.

AR mutations. AR mutations in the early stage of PCa are rare, but it is commonly occurs in CRPC, particularly advanced PCa following systematic hormone therapy. There are >100 point mutations that appear in AR, and the majority of them present in the NTD or LBD region (4,23). In recent years, there are a number of studies regarding AR mutations including T878A, W742C, L702H and F877L (24-26). The mutations in the hinge and LBD regions of AR result in increased AR transactivation activity and decreased ligand specificity. A large number of AR mutations can be activated by adrenal androgens and other steroid hormones; additionally, a number...
of mutations can turn AR antagonists (flutamide and bicalutamide) into potential agonists. Compared with endocrine therapy, AR antagonists in the treatment of metastatic PCa will cause the rate of AR mutations in the LBD region to be increased.

The T878A mutation is one of the most common forms of AR mutations. It was found that the AR T878A mutation occurs in patients with CRPC following prolonged ADT; however, not hormone-sensitive patients (27-29). Furthermore, it has been reported that this mutant was expressed in the tumors of patients with CRPC whose PSA levels were significantly decreased by flutamide withdrawal (29). This discovery has led to the hypothesis that expression of the AR T878A mutation is responsible to the beneficial effect of withdrawing anti-androgen therapy in patients with CRPC. This mutation also leads to an expansion of binding specificity of the ligand of AR, improving the sensitivity of steroid hormones such as progesterone and estrogen (30). A number of the mutations can turn the anti-androgen substances into agonists. For example, the F877L mutation activated by enzalutamide or ARN-509 (31). Recent studies indicated that the mutations T878A, H875T/Y, W742C and L702H exist in 15-20% of CRPC tumor samples, which emphasized that the LBD region is a mutational hotspot (32). Utilizing circulating free DNA from patients with CRPC, which has been determined to contain genomic DNA with the AR mutations, demonstrated that using sequencing to detect those point mutations aforementioned could possibly act as biomarkers for patients at risk of developing CRPC (33). Additionally, combination therapy may overcome part of the secondary resistance with AR mutations, an ongoing clinical trial (34), using combination therapy of abiraterone and enzalutamide in patients with CRPC may demonstrate this hypothesis. If those studies using blood-based detection to distinguish AR mutations in patients with CRPC are clinically validated, then clinicians may have accessible biomarkers for individual treatment selection, which may improve the clinical outcome for patients.

Expression of AR splice variants (AR-Vs). AR-Vs have been demonstrated to be associated with resistance to ADT of PCa as well as the resistance to abiraterone and enzalutamide (35). AR-Vs are truncated forms of AR, which lack the LBD region, where just the NTD and DBD regions were transcriptionally activated, independent of the presence or absence of ligand-binding or antagonist effect (36). Hu et al (37) explored the AR intronic sequences in the human expressed sequence database, and determined that CE 1-4 and identified AR variants (AR-V 1-7) lacked the LBD region. This trait of ligand-independence makes truncated AR variants potentially important to disease progression and patient treatment response, as they can activate AR target genes independent of the androgen (36). The current studies discovered splice variants include AR3, AR4, AR5 and AR-V1V7 (38). The AR transcripts AR-V3 and AR-V4 have the same sequences to AR1/2/2b and AR1/2/3/2b; therefore, novel variants were identified as AR-V1, AR-V2, AR-V5, AR-V6 and AR-V7 (39). A number of the AR-Vs (including AR-V7) primarily localize in the nucleus, while others (including AR-V1, AR-V4 and AR-V6) localize mainly in the cytoplasm. Notably, cytoplasm-predominant AR-Vs frequently co-expressed with the nucleus-predominant AR-Vs as well as the full-length AR (AR-FL) (40). Not only are AR-Vs predominantly products of alternative splicing, but also can they be products of nonsense mutations (ARQ640X) or proteolytic cleavage (41) (Fig. 2).

Detected at the mRNA and protein levels, AR-V7 is the most frequently studied and abundant AR variants in clinical samples from patients with CRPC (42). A retrospective study have indicated that the presence of AR-V7 was associated with more advanced malignance and reduced survival in patients with metastatic CRPC (mCRPC) (43). Previously, the study of Antonarakis et al (44) have carried out a prospective study, which utilized CTC samples of patients receiving abiraterone or enzalutamide therapy to evaluate the prognostic role of AR-V7 detected at the mRNA levels by RT-PCR. They discovered that compared with those patients without detectable circulating AR-V7, the appearance of AR-V7 in CTCs was associated with lower PSA response rate, reduced progression-free-survival (PFS) and reduced overall survival (OS) (44). Notably, the level of AR-V7 was higher in males who had been treated with enzalutamide and abiraterone, whilst the level of AR-V7 was lowest in males who had not received either agent therapy. In addition, when analyzing
serial CTCs over time, the study demonstrated that all males with a baseline of AR-V7 remained AR-V7 positive during the process of abiraterone and enzalutamide therapy, whereas 14% of males with negative AR-V7 level at baseline changed to AR-V7 positive during treatment, and these patients had an intermediate clinical outcome (44).

It was confirmed that AR-V1, AR-V4 and AR-V6 can dimerize with AR-V7 and AR-FL; therefore, AR-V7 and androgen-bound AR-FL induced nuclear localization of AR-V1, AR-V4 and AR-V6. Additionally, these variants weakened the efficiency of enzalutamide to inhibit androgen-bound AR-FL nuclear localization. It is notable that the impact of nuclear localization of AR-V4 and AR-V6 on AR transactivation differs from that of AR-V1 (45). Nuclear localization results in an enhanced ability of AR-V4 and AR-V6 to transactivate canonical AR targets and AR variants specific targets, and promote castration resistant cell growth (45); however, when AR-V1, which lacks inherent transcriptional activity, activates AR-FL in an androgen-independent way, it significantly inhibits AR-V7 transactivation (45). These data illustrated the important role of complicated interactions among different AR-Vs and AR-FL in castration resistant disease, and dissecting these complex interactions may help to develop effective strategies to target AR variants signaling (45). In recent research, AR-NTD targeting drugs have benefits over drugs targeting the AR-LBD due to the NTD being essential for the transcriptional activities of AR-FL and AR-Vs; therefore, AR-NTD antagonists, including EPI-002, patients who are resistant to abiraterone or anti-androgens with constitutive activated expression of AR-Vs could achieve therapeutic responses from EPI-002 (46,47). As a result, with gradual clarification of the AR-Vs formation, it has a possibility to become a novel targeted therapy for the treatment of patients with CRPC in the future.

**Altered expression and function of AR co-regulators.** AR co-regulators are protein factors, which were associated with AR transcription activation, and serve an auxiliary role in activation or inhibition of AR-mediated transcription (48). AR co-regulators have the function of activating transcription activity of AR under an extremely low concentration of androgens (48). It has also been demonstrated that co-regulator factors serve an important role in the development of CRPC formation. Normally, AR recruits a series of co-regulator complexes, which can either enhance or repress transcriptional activity. Many of these co-regulators are enzymes that serve to modulate other proteins through phosphorylation, methylation, acetylation or ubiquitylation, in a complex form (49,50). They have also been identified as molecular chaperones, recruiters of transcriptional machinery and RNA splicing regulators (49,50). According to their different effects on AR transcription, AR co-regulators can be divided into co-activators [including P160/SRC and p300/CREB binding protein (CBP)], co-repressors (including NcoR 1 and NcoR 2), pioneer factors (including forhead box A1, GATA2 and cardiotrophin 1) and cooperators (including ETS-1, adaptor associated protein complex 4 and NKX3-1) (51-55). There are >150 different molecules that have already been identified as co-activators or co-repressors of AR (56). When AR interacts with a co-activator, it can lead to a significant abnormal activation of AR, which results in increased AR transcriptional activity (50); therefore, changing the ability of AR transcriptional activation is essential for AR to achieve the maximum transcriptional effect. For example, FKBP51, as an AR target gene and a co-activator of AR, was determined to be upregulated in relapsed LAPC-4 tumors that were grown in castrated mice (57). The formation of a super chaperone complex was improved by FKBP51 via recruiting p23, to adenosine 5'-triphosphate (ATP)-bound Hsp90, which in turn keeps AR in a conformation with high affinity for ligand-binding, thus promoting androgen-induced transcriptional activity and growth (58). Other important pathways, including p300/CBP, which promotes androgen-independent interleukin (IL)-6 mediated AR activation in the presence of the signal transducer and activator of transcription 3 (STAT3), lysine demethylase 1A (LSD1A), jumonji domain-containing 2c (JMJD2c) and lysine demethylase that demethylates the histone H3 proteins and then promotes AR induced transcription (59). On the other hand, when AR is combined with co-repressor, the opposite effect occurs in reducing the AR transcriptional activity, although they have been observed at decreased levels in CRPC (60). Additionally, the majority of cooperators serve...
the role of increasing the AR transcription, whilst pioneer factors can interact with condensed chromatin and directly regulate the accessibility of chromatin (61). It is important that the interaction between pioneer factors and chromatin is prior to the androgen induction, which means that these factors can direct AR to chromatin aggregation at low or no levels of androgens (54,55).

**Aberrant post-translational modifications of AR.** Post-translational modifications of AR mainly include acetylation, methylation, ubiquitination, sumoylation and phosphorylation (62). The molecular effects of these transformations induce decreased apoptosis and elevated transcriptional activity to the androgen-responsive genes (62). Research indicated that hypomethylating drugs can be used as a treatment in delaying the appearance of CRPC (63). Inhibiting DNA methylation can reverse the mechanism underlying castration resistance, the potential underlying mechanism may be associated with the downregulation of DNA methyltransferase 1-dependent STAT3 activity. Many serine/threonine and tyrosine kinases are involved in AR phosphorylation, including Aurora-A, PIM1 kinase, cyclin-dependent kinase 1 (CDK1), Src and Ack1 (64-67). Aurora kinase (68) is the most extensively studied clinically. In a phase II trial (69), 60 randomly selected patients with mCRPC were randomized into two different treatment schedules based on using danusertib or not, an inhibitor of Aurora kinase. All of the patients had been previously treated with docetaxel. Only two patients had a PSA response, and the best radiographic response was an indicated stable disease in 27% of patients. The median PFS was 12 weeks in both groups. Due to of the disappointing results, no further research on this compound is being pursued. The study indicated that aurora kinase was upregulated in anaplastic or neuroendocrine PCa (70). More recent clinical trials (NCT01799278 and NCT01848067) are using these agents to target the small cell and anaplastic PCa. Research for these targets is actively continuing, although no late-phase clinical trials are currently planned or ongoing. These phosphorylation modifications, which enable AR to respond to low levels of androgen, and being the targets of small molecule inhibitors could be a promising treatment for patients with CRPC.

**Synthesis of adrenal androgens and intratumoral androgens.** Normally, testosterone mainly originates from the testes, with 5-10% coming from adrenal glands. In general, medical or surgical castration can reduce circulating levels of serum testosterone by >90% to maintain its circulating concentration below normal levels of castration (1.25 pmol g⁻¹) (70); however, androgen concentration in prostate tissues remains sufficient for activation of AR (70). The androgen concentration in prostate tissues is significantly lower than that of serum androgen in patients following ADT (71). PCa may act as synthetic androgen within tissues through a number of unknown mechanisms; therefore, PCa tissues can synthesize androgen using prostate tissues and not rely on androgen cycle following ADT, and this condition may result in failure of the procedure (71). Recently, utilizing autopsy and tissue biopsy studies to compare androgen levels in hormone-sensitive and hormone-resistant tumors from patients has revealed high levels of continuous androgen production in tumor tissues of patients with CRPC (72). Studies have considered that sustained production of androgen within prostate tissues possibly results from the conversion of weak-bioactivity androgen-like androstenedione (AD) and dehydroepiandrosterone (DHEA) (73,74), which are produced by adrenal glands or de novo steroidogenesis of androgen from the cholesterol within tumor tissues (74). Although circulations of AD and DHEA significantly decrease in patients with CRPC following treatment with abiraterone (CYP17 inhibition), a sustained pool of DHEA-S, which is a sulfated form of DHEA, and the predominant adrenal androgen in circulation can serve as precursors for transformation into testosterone and DHT in prostate tissues (75). The role of intratumoral androgen in promoting CRPC cells growth was proven by large phase III clinical trials, which aimed to test the testosterone concentration of tumors in patients receiving novel anti-androgen therapies of enzalutamide and abiraterone (76,77). All patients with mCRPC were tested and indicated improved OS, compared with those who received placebo treatment.

**4. AR by-pass signaling: Glucocorticoid receptor (GR) and progesterone receptor (PGR) expression**

Steroid receptors, including AR, estrogen receptor-α, estrogen receptor-β, PGR, GR and mineralocorticoid receptor, are derived from the same material and feature high homology, particularly in the DBD binding region (79). GR and PR are more relevant to AR (80). Almost all PCa were discovered to express AR, whereas GR is only present in 30% of PCa cases; however, GR expression increases in patients following ADT (81). Recent studies have indicated that AR and GR possess the same chromatin binding sites and regulate the expression of a large number of AR-specific genes (82). Resistance of enzalutamide to novel anti-androgen therapies of enzalutamide and abiraterone (76,77). All patients with mCRPC were tested and indicated improved OS, compared with those who received placebo treatment.

Montgomery et al have summarized a series of clinical trials to assess the contributions of glucocorticoids and GR in patients with CRPC (84,85). In these trials, analysis of patients in clinical cohorts who received glucocorticoids indicated poor prognostic features, compared with patients who did not receive such treatment; therefore, stimulation of GR signaling may pose negative effects to patients who have received androgen-targeted therapies (84-86). In such conditions, the presence of increased glucocorticoids in serum and absence of androgen possibly enables the selection of ‘promiscuous’ AR variants with mutations in the LBD, allowing their activation by glucocorticoids (87); therefore, GR or other nuclear steroid receptors can bypass AR pathways and promote the development of CRPC.

On the other hand, PGR is also a member of the steroid hormone nuclear receptor family, and it is structurally associated with AR. As with GR, PR may have the ability to transcriptionally regulate a subset of AR target genes in PCa and thereby
bypass AR (88). A large retrospective analysis carried out recently demonstrated the association of high PR staining in primary PCa with its clinical recurrence (89). A recent research on a cohort of >500 patients also revealed the association of high PR expression in cancer cells with reduced clinical failure-free survival (89). This evidence indicated that PR antagonists may possess therapeutic effects, indicating that PR signaling can be an important carcinogenic target in the treatment of PCa.

5. AR-independent pathways

Phosphoinositide 3-kinase-protein kinase B-mechanistic target of rapamycin (PI3K-Akt-mTOR) pathway and CRPC. The PI3K-Akt-mTOR pathway has been known as a potential driver of CRPC. PI3K is activated by G protein-coupled receptors or tyrosine kinase receptors. Activation of PI3K leads to phosphorylation of AKT and activation of mTOR and subsequent downstream effects, including cellular proliferation, survival and angiogenesis (90,91). Previous studies have demonstrated that alterations of components in the PI3K-Akt-mTOR pathway occur in 100% of metastatic tumors and 42% of primary prostate tumors (4,91,92); therefore, targeting the PI3K-Akt-mTOR pathway is considered a promising therapeutic approach for patients with CRPC. Loss of the phosphatase and tensin homolog (PTEN) gene has been frequently proven to result in constituting activation of the PI3K-Akt-mTOR pathway in the majority of advanced PCa cases (93). A recent study used either targeted drugs or gene knockout in mouse models, whereas cell lines have indicated alterations in PI3K and PTEN activities, demonstrating changes in AR expression and AR transcriptional activity (94). Kato et al (92) reported the combined use of the AR-V inhibitor EPI-002 and PI3K/mTOR inhibitor BEZ235 to evaluate their therapeutic efficacy on LNCaP95 cells in CRPC models in vitro and in vivo. Results demonstrated that compared with single drug cohorts, drug combination can significantly inhibit LNCaP95 cell growth without increasing toxicity. These data indicated that co-targeting the PI3K-Akt-mTOR pathway and AR-NTD to block AR-FL and AR-Vs can be a novel approach for achieving improved antitumor efficacy for patients with CRPC who have indicated resistance to abiraterone or other anti-androgen treatments by a mechanism involving expression of AR-Vs.

Src signaling pathway and its inhibitors. The Src kinase family (SFK) comprises of proteins with protein tyrosine kinase. Src kinase activation has been indicated to be relevant to
androgen-independent cell growth, inhibition of anti-apoptotic pathways, cell migration and adhesion and tumor invasion. In PCa, Src signaling upregulates vascular endothelial growth factor (VEGF) and IL-8 to promote angiogenesis (95) and activates the nuclear factor-KB pathway of osteoclasts and various tumor necrosis factor receptors involved in resistance to apoptosis signal transmission to participate in bone metastasis of PCa (95). Dasatinib, an inhibitor of Src kinases, has been confirmed to slow down the growth of PCa tumors and to be as effective as CRPC treatment based on a phase III clinical trial (96). Saracatinib (AZD-0530) is a dual inhibitor of Src and AbI, and the research of Yang et al (97) demonstrated that this drug can inhibit proliferation and invasion in bone metastasis models of PC3 cell lines and alleviate bone destruction; a phase II clinical trial with Saracatinib is currently ongoing.

**Growth factor pathways.** Activation of the growth factor signaling pathway can also enhance AR signaling pathway and promote CRPC. Growth factor receptors, including insulin-like growth factor I receptor, IL-6 receptor and epidermal growth factor receptor, can mediate the growth of critical downstream pathways, including mitogen-activated protein kinase (MAPK), PI3K/AKT and signal transducer and activator of transcription signaling (98). Activation of PI3K-Akt and Ras-Raf-MAPK pathway and expression of receptor tyrosine kinases, including Her-2/neu, have already been observed to enhance AR stability and activity in CRPC samples (99). A previous study demonstrated that Her-2/neu promotes the growth of xenograft cells and androgen independence through Ack1-kinase, which phosphorylates AR at tyrosine-267 and activates AR activity (100). Targeting growth factor signaling pathways has provided a novel therapeutic strategy. In a phase II clinical trial (101), application of cabozantinib (XL-184), an inhibitor of tyrosine kinase of c-Met and VEGF, resulted in PFS of 23.9 weeks in patients with CRPC, whereas the placebo group yielded a PFS of 5.9 weeks. Following treatment with cabozantinib for 12 weeks, bone scan of 68% of patients indicated improvement; whereas, results for 12% of patients demonstrated that bone metastatic lesions completely subsided. XL-184 can also relieve pain of patients with CRPC to improve their quality of life (101).

**6. Expression of microRNAs (miRNAs) in CRPC**

miRNAs are small, non-encoding, and conserved genetic single-stranded RNAs. miRNAs can target gene mRNA 3’ untranslated region through complete or incomplete complementary binding, resulting in degradation of target gene mRNA or inhibition of translation; which functions as a gene expression regulation factor in cell metabolism, proliferation, differentiation and cell death and cell survival (102,103).

Extensive research indicated that expression profiles of miRNAs in PCa become increasingly important due to their application in diagnosis, staging, predicting prognosis and response to the treatment (104). An earlier study confirmed upregulation of miRNA-96, -182, -183 and -375 and downregulation of miRNA-16, -31, -125b, -145, -149, -181b, -184, -205, -221 and -222 in PCa tumor tissues (105).

In a recent study, using miRNA microarray to detect miRNA expression between primary PCa tumors and CRPC tumors has indicated divergent results, where 75 miRNAs were differentially expressed in primary PCa, 88 in CRPC tumors, and 22 changed expressions of miRNAs overlapping between primary and CRPC samples (106). Altogether, these data implicate that changes in miRNA expression can contribute to resistance to ADT.

The most frequently studied miRNAs associated with PC progression include miRNA-16, miRNA-34a/34b, miRNA-143, miRNA-101 and miRNA-200c. Cell experiments indicated that exogenous expression of miRNA-16 can inhibit the recurrent growth of 22Rv1 cells, androgen-independent DU145 cells and PC3 cells, but not androgen-dependent LNCaP cells (107). miRNA-34a has been demonstrated to specifically downregulate AR expression (108), indicating that loss of miRNA-34a can increase AR expression, as observed in PCa cell lines, xenograft models and human CRPC tumors (17). Observation of patient with PCa samples has demonstrated that high miRNA-34b expression is correlated with longer OS, whereas the opposite is correlated with higher Gleason score (15-17,109). One assumption indicated that induced miRNA-34b expression can increase the aggressiveness of PCa. Expression levels of miRNA-200c were significantly reduced in primary PCa patients of tumors who relapsed following prostatectomy, compared with those without relapse (110). miRNA-101 expressions notably decreased in metastatic tumors, compared with those of primary samples (111). All these data implicated that miRNAs cannot only potentially promote tumor progression and sustain resistance to ADT but also act as biomarkers for patient outcomes. Analysis of radical prostatectomy specimens indicated that miRNA-301a, miRNA-449b and miRNA-182 can act as biomarkers to predict biochemical recurrence following prostatectomy (112-114). Recently, comparing Gleason scores and lymph node status with expression levels of miRNA-4516 and miRNA-601 has demonstrated predictive roles of these miRNAs in biochemical recurrence following post-prostatectomy salvage radiation therapy (115), supporting the utilization of miRNAs in clinically used predictive models.

In conclusion, miRNAs serve important roles in initial PCa pathogenesis, progression and development of CRPC. miRNAs can also function as attractive biomarkers due to they are relatively stable in biological fluids, easy to measure and are resistant to storage handling (104); therefore, abnormal levels of expressed miRNA in tumor tissues, serum or plasma and urine can be a promising biomarker for diagnosis, prognostic prediction, or treatment efficacy of patients with PCa.

**7. Defects in DNA damage repair (DDR) in CRPC**

Defects and mutations in DDR genes have been reported in CRPC and in localized aggressive cancers (116). For example, mutations and loss of BRCA2 were observed in 12% of PCa cases (117). Germline mutations in BRCA increase the probability of lymph node involvement and distant metastases, which confer PCa with more aggressive phenotypes (118). CRPC frequently features homologous DNA recombination function deletion, expression of TMPRSS2-ERG gene fusion and the DDR protein poly (ADP-ribose) polymerase (PARP)-1,
which is recruited to sites of AR targets and serves an important role in AR transcription, which is essential for the activity of TMPRSS2-ERG in PCs (119). In a phase II trial (120), patients who had recurred following >2 rounds of CRPC therapies were treated with olaparib, a PARP inhibitor and a base excision repair protein. Results demonstrated that patients who exhibited genomic defects in DNA repair genes (including BRCA1, BRCA2 and ATM) manifested a 6.3-month increase in OS, compared with those without these defects. These results indicated that blockade of DNA damage response may promote CRPC, whereas genomic instability improves responsiveness of patients with CRPC to treatment for target DNA repair (120).

AR indicated potential links with DDR. By using a combination of anti-androgen therapy and radiotherapy to treat aggressive PCa, large clinical trials demonstrated significant augmentation in efficacy, indicating the potential role for AR inhibition in weakening DDR (121,122). AR enhances the expression and activity of key DDR factors, including DNA-dependent protein kinase (DNAPK); in turn, DNAPK sustains AR transcription (122). ADT induces reduction in transcription of key DDR genes, resulting in higher levels of DNA damage following radiotherapy, specifically non-homologous end-joining (121). Such results indicated that ADT affects genomic instability prior to castration or promotes development of CRPC through repressing DDR. MYB protein was observed to replace AR function in regulating DDR by modulating an overlapping set of genes. Silencing of MYB gene or a number of its targets (TOPB1, ATR and checkpoint kinase 1) synergizes with PARP inhibitor olaparib, and such condition can significantly suppress PCa growth in in vitro and in vivo experiments and increase cytotoxicity of PCs (123); therefore, combination of MYB pathway inhibitor and PARP activity suppressor can be a viable clinical strategy.

8. Autophagy and CRPC progression

Autophagy is a genetically programmed cellular stress regulation mechanism occurring in all eukaryotic cells. This process is one of the major protein degradation processes that prevail through the lysosomal pathway (124). Autophagy mediates the clearance of damaged organelles and long-lived proteins, leading to the formation of double-membrane structures called autophagosomes (Fig. 4). Cells can recycle amino acids and other macromolecular materials for biosynthesis via autophagy. Normal cells clear chemical carcinogens and impaired organelles caused by radiation or oxidative stress mainly through the action of mitochondria, thereby protecting cell DNA against damage from reactive oxygen species, ensuring genomic stability and reducing incidence of cell malignant transformation (125); therefore, autophagy is crucial in maintaining hereditary stability.

Recent studies indicated that autophagy may be associated with resistance to chemotherapy in malignant tumors. For example, the studies of Jiang et al (126,127) have reported that overexpression of mitochondrial-associated leucine-rich pentatricopeptide repeat motif-containing protein, an inhibitor of autophagy, and low levels of autophagy activator microtubule-associated protein 1S (MAP1S), a member of MAP1 family, can be both used as independent biomarkers for patients with late-stage PCs and poor prognosis. Patients receiving multiple chemotherapy, particularly for oxygen-deprived tumor cells, demonstrated significant increase in the degree of nutrient limitation. Ischemia, hypoxia and lack of nutrition can activate autophagy. Autophagy modifies metabolism by degradation and recycling of intracellular substances. Through this mechanism, tumor cells can induce adaptive responses to stress and survive. In the late stage of tumor progression, autophagy becomes an important survival mechanism for tumor cells that cannot directly obtain nutrition from circulation in the central body of solid tumors (128,129). Facial tumors and those determined that have poor living environments can eliminate large molecules or mitochondria damaged by ionizing radiation and cell toxicity through enhancing autophagic activity and blocking transmission of the mitochondrial apoptosis signal cascade (129). Autophagy can also prevent pro-apoptotic factors, including cytochrome and apoptosis-inducing factor diffusion by separating mitochondria. These mechanisms may form a resistance of tumor cells to radiotherapy and chemotherapy and prevent their apoptosis. Patients with CRPC frequently use chemotherapy based on docetaxel and mitoxantrone as palliative therapy. However, with repeated administration, resistance to chemotherapy becomes prevalent in patients, but overall effect is poor (130,131). Existing studies imply that autophagy serves an important role in development of CRPC (130-133).

Using enzalutamide to block AR as a long-term treatment can activate AMP-dependent protein kinase [AMP-activated protein kinase (AMPK)], whereas mTOR is inhibited by mTOR modulin (132,133). Studies have indicated that AMPK can directly regulate mTOR activities to match energy imbalance, thus increasing the ratio of AMP/ATP (132,133). Hypoxia is one of the indirect causes by which anti-androgen therapy can improve autophagic activities (130), whereas blockage of angiogenesis by androgen withdrawal and nutrient synthesis under hypoxia further stimulates autophagy. The information aforementioned can confirm the effects of AR on autophagy surges but fail to fully elucidate autophagy mechanisms. For example, ADT can still be used to develop PC3 cells that lack AR for upregulating autophagy (132). Possible reasons include changes in proto-oncogenes, including PTEN, altered tumor suppressor genes, including cell tumor antigen P53 and abnormal metabolism of AR downstream (134). Bennett et al (131) indicated that the use of synergistic autophagy inhibitor 3-methyladenine with bicalutamide results in 1.5-fold higher cell mortality rate than using bicalutamide alone. Colquhoun et al (135) also demonstrated that using bicalutamide with metformin in treating LNCaP cells will significantly decrease the colony formation rate. This phenomenon also occurs in PC3 cell lines although the latter requires higher doses. Notte et al (136) discovered that two key pathways of taxol inducing autophagy; the first is paclitaxel-inhibited mTOR, whereas the second participates in activation of c-Jun N-terminal kinase, Bcl-2 and Bcl-XL phosphorylation and activation of autophagy. Notably, although these pathways perform similar functions, treatment time with paclitaxel and environmental pressures required for activation of autophagy remarkably differ. LNCaP-AI is an androgen-dependent PCs cell line. When cultured in an androgen-deficient environment, tolerance of the LNCaP-AI cell line to docetaxel is 2.5 times greater than that of common LNCaP cell lines (131). Recovery of androgen of the LNCaP-AI cell line in culture medium also indicated sensitivity to
docetaxel (131). When re-adding 3-methyladenine to inhibit autophagy, cytotoxicity of docetaxel increases; thus, autophagy is a factor influencing docetaxel resistance (131).

Kinases serve important roles in the metabolism and proliferation of PCa cells and thus affect the progression of the disease. A previous study established PCa models in xenogenic mice, with one group receiving saracatinib treatment and another receiving saracatinib combined with chloroquine treatment. Following 14 days of observation, the rate of tumor growth inhibition reached 64%, whereas that of single-saracatinib group totaled 26% (137). Current limitation of inhibitors of SFK may be due to simultaneous emergence of therapeutic effects and therapeutic resistance caused by overlapping molecular pathways (137). An autophagy inhibitor may overcome this limitation.

At present, many clinical trials focus on regulation of autophagy in PCa. However, regulation of autophagy still lacks specificity, and influences on non-tumor tissues are also uncertain. In the future, the aim is to gain further insights into dynamic changes in autophagy during PCa development and treatment. Through specific autophagy regulation, inducing cell apoptosis and malignization may be a viable strategy for treatment of PCa.

9. Prostate stem cells

Stem cells are a type of cell that indicate multiple differentiation potential, self-renewal, absence of differentiation or low differentiation and exist in multicellular organisms (138). In recent years, constant research demonstrated that prostate stem cells are closely associated with the occurrence and development of PCa. Prostate epithelial stem cells exist in the basal layer, which generates intermediate amplification cells, resulting in highly differentiated luminal epithelial cells of the prostate (139). Mesenchymal stem cells exist in the prostatic stroma, and they are induced by the microenvironment to generate various prostate stromal cells, including fibroblasts and muscle cells. Prostate epithelial stem cells and mesenchymal stem cells do not depend on the presence of androgen but can respond to this hormone (140). Richardson et al. (141) confirmed that ~1% of human prostate basal cells express cell surface markers, CD133, cells with phenotype of α2β1 and hi/CD133+ possess high proliferative potential in vitro and can reconstruct prostatic acini in immunodeficient male nude mice. A previous study also isolated a type of PCa cell similar to prostatic basal cells from androgen-dependent PCa; these cells can also survive without androgen (140). The study of Collins et al. (142) successfully isolated PCa stem cells with the phenotype of CD44+/α2β1 hi/CD133+ from PCa tissues utilizing a transplantation assay based on isolated cells xenografted into immunodeficient mice; they first proved the existence of PCa stem cells, and this result may explain why patients with PCa will indicate potential castration resistance and metastasis following ADT.

PCa stem cells are generally believed to be caused by mutations of prostate stem cells, and development of PCa is possibly associated with mutations in stem cells of the prostate (143). Stem cells from PCa or the normal prostate possess unlimited self-renewal ability and the capacity for multipotential differentiation, they improve self-protection through a variety of ways, including anti apoptosis. Stem cells activate membrane transport proteins and strengthen DNA repair capacity and they differentiate into cancer tissues similar to primary prostate carcinoma following transplantation in vitro (138). However, PCa stem cells and prostate stem cells also exhibit differences. Normal prostate stem cell proliferation is frequently associated with tissue damage or other conditions, whereas malignant proliferation of PCa stem cell is usually associated with loss of body control (140,144). By contrast, normal prostate stem cells can differentiate into mature tissues and perform corresponding functions unlike PCa stem cells, which can only differentiate and mature. Compared with normal prostate stem cells, PCa stem cells are more prone to accumulate mutations (140,144). Although prostate stem cells have thus far not been purified in vitro, based on their previous experience, Wang et al. (145) retrieved stem cell-like cells from PCa tissues in 2014. At present, PCa stem cells and prostate basal stem cells express a number of similar markers, including CD44, integrin α2β1 and CD133, which can be used to identify and isolate stem cell populations. Clinically, prostate stem cell surface markers, including prostate stem cell antigen, can be used to assist in the diagnosis of PCa or metastatic tumors of the prostate (141,142,146). Recent evidence has demonstrated that prostate cancer stem-like cells (PCSCs) serve a critical role in stimulating CRPC evolution and resistance to abiraterone and enzalutamide (147). For CRPC, the drug control of PCa
stem cells may be a novel targeted therapy direction for future PCa treatments. A precautionary PCa vaccine can also be developed based on characteristics of PCa stem cells to prevent the progression of human PCa.

10. Conclusion
Currently, PCa, particularly advanced and metastatic, has been a continuous burden on the healthcare system; therefore, the clinical research on CRPC has become a hot spot. Multiple mechanisms underlying the resistance to AR targeted therapies have been identified, including AR overexpression with or without amplification, AR mutations, AR-Vs, intratumoral DHT synthesis, GR overexpression, defects of DDR genes and loss of AR, with other therapies to be discovered. In addition to the aforementioned underlying mechanism, including the studies of PSMA, which contains high expression in androgen resistance prostate cells and has been demonstrated to promote the proliferation, invasion and apoptosis of PCa cells (148); additionally, the p100/p52 pathway, one of the NF-κB pathways, the process of p100 to p52 in PCa via molecules, including B-cell activating factor, CD40 ligand and STAT3, leads to significant hyperplasia and induces the growth of castration-resistant cells (149). The majority of those advancements in the mechanism understanding of CRPC have provided numerous potential biological targets for the CRPC treatment, including AR antagonist abiraterone and enzalutamide, AR-NTD antagonist EPI-002, PARP inhibitor olaparib, PI3K/mTOR inhibitor BEZ235, chemotherapy drugs cabazitaxel and immunotherapy sipuleucel-T, which can improve the OS and disease-free survival of patients with CRPC (47,92,150-152). In addition, the majority of these pharmaceutical agents are undergoing clinical trials and are being approved for the treatment of CRPC by FDA (6,153); however, all of these agents merely slow the progression of disease, and all patients will inevitably deteriorate into an incurable stage. The changes in the AR signaling pathway, multiple genes and signaling pathways are involved in the process of PCa from early androgen dependence to late CRPC (154), and the underlying mechanism remains at a limited understanding. The interconnection of these signaling pathways form a molecular signaling network, inhibiting a target alone or blocking the occurrence and progression of a signaling pathway is not sufficient to prevent CRPC. To conclude, the focus of the study is to identify and control the key genes targeting these pathways, and effectively prevent PCa from CRPC transformation and evolution, which has an important role in adjustment of CRPC therapeutic strategies, discovery of novel drug targets and improvement of the therapeutic effect.

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JXH and HYQ drafted the manuscript; LX and JGG were responsible for the collection of the relevant literature. All authors have read and approved the final manuscript.

Ethics approval and consent to participate
This article does not contain any studies with human participants performed by any of the authors.

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Competing interests
The authors declare that they have no competing interests.

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