Recent Advances in Androgen-Receptor Splicing Variants Related to the Mechanism and Treatment of Castration-Resistant Prostate Cancer

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Introduction
Prostate cancer is one of the most common malignant solid tumors in men. In the United States in 2020, prostate cancer is predicted to cause 191930 new cases, accounting for 21% of the total number of new solid tumors in all men. [1] China and Asia faster-growing incidence of prostate cancer (PCa), [2] The death rate from prostate cancer in urban areas of mainland China increased by nearly 50% in 2009, compared with 2004. At present, there are significant regional differences in the detection of prostate cancer in China, and the patients with advanced metastatic prostate cancer account for a large proportion in the whole patient population. [3]-[6] Although endocrine therapy is the standard regimen for the treatment of advanced metastatic prostate cancer, almost all patients will eventually develop drug resistance and enter Castration-resistant prostate cancer (CRPC) after receiving endocrine therapy within 18 to 36 months, [7] resulting in significantly increased mortality and poor quality of life. Therefore, more in-depth research is needed in the mechanism and treatment of CRPC. Androgen receptor (AR) is the focus of research, and androgen receptor splicing variant (AR-Vs) is a new and important breakthrough point. This review provides a theoretical basis for further research on androgen receptor splicing variants related to the mechanism and treatment of CRPC.

2.1 Diagnostic Criteria of CRPC
In the definition of CRPC, "castration" includes surgical castration and drug castration. Regardless of the castration mode, 95% of the androgens will be removed in the future from the hypothalamic-pituitary-gonadal axis. Diagnosis of CRPC in the 2017 edition of the European Association of Urology (EAU) guidelines includes three criteria: (1) Serum testosterone reaching castration level (< 50 ng/dl, or < 1.7 nmol/L); (2) Prostate-specific antigen (PSA) progression: 3 times in a row during the interval of 1 week, the PSA increased by more than 50% from the minimum value, and the absolute value of PSA increase was >2 ng/ml; (3) Imaging progress: bone scan found two or more bone metastatic lesions or large soft tissue lesions meeting Response Evaluation Criteria in Solid Tumors. [8] The CRPC consists of the following 2 stages: (1) Androgen independent and AR dependent stages; (2) Androgen independent and AR independent stages.[9]

2.2 Androgen Receptor (AR) and Androgen Receptor splicing Variant (AR-Vs)
Androgen receptor (AR) is a member of the steroid hormone receptor family. [11] It contains n-terminal trans-activation domain (encoded by exon 1), DNA binding domain (encoded by exon 2 and 3), short hinge region (encoded by exon 4), and c-terminal ligand-binding domain (LBD; Exon 4-8 encoding). [12] AR is activated by binding to androgen, and plays a key role in male physiology and pathology, involved in all stages of prostatic tumor initiation, development, and therapeutic resistance. [10] Prostate cancer by a variety of mechanisms such as AR gene amplification, mutation and steroid metabolism, cell signal transduction and regulation of the change of the core protein produces resistance to androgen deprivation therapy (ADT). [13] The expression of androgen receptor splicing variants (AR - Vs) lacking ligand-binding domain is a kind of new mechanism. [7] The discovery of the constitutive active AR splicing variant has become a new clinical trial and novel method for targeting AR.[13]

As a truncated subtype of AR, the androgen-receptor splicer variant (AR-Vs) lacks ligand-binding domain and still has constitutive activity in the absence of circulating androgens. [14] It activates AR reporter genes in the absence of ligand, thereby promoting the progression of prostate cancer. [17][18] Compared with primary prostate cancer, the expression of androgen-receptor splicing variant with truncated C terminal in CRPC was significantly increased. [15] The expression of AR-Vs AR-V1, AR-V7 and AR-V567es in CRPC bone metastasis was increased and indicated poor prognosis. [19] AR-V7 in androgen receptor splicing variant was particularly important, and more and more evidences emphasized that the expression of AR splicer variant could be used as a potential
3. Several androgen-receptor splicing variants that play an important role in CRRP

3.1 Androgen receptor splicing variant 567es (ARv567es)
ARv567es is produced by splicing the 5th, 6th and 7th exons of AR. [22] The expression of ARv567es can often be detected in human metastatic prostate cancer. [19][20]ARv567es can induce castration resistance in prostate tumors of human xenograft, and the ratio of androgen receptor full length(ARFl) expression is positively correlated with castration resistance. [22] In the standard taxane treatment of castration-resistant prostate cancer (CRPC), the effect of docetaxel treatment was significantly enhanced because ARv567, after interacting with the microtubules of prostate cells, increased the sensitivity of taxane-induced microtubules stabilization. [21]ARv567es can be used as a biomarker for early recurrence of CRRP and is associated with a particularly poor prognosis of CRRP. [19][22]ARv567es promotes the development of castration-resistant prostate cancer by inducing tumorigenesis de novo, [20] acting as constitutively active receptors, increasing the expression of androgen receptor full length(ARFl) and enhancing the transcriptional activity of AR. [22][20]

3.2 Androgen receptor Splicing Variant 7(AR-V7)
Ar-v7 /AR3 is one of the major splicing variants expressed in castration-resistant prostate cancer tissues in humans. It has constitutive activity and its transcriptional activity is not regulated by androgens or antiandrogens. [23] Under the condition of androgen deprivation and the absence of ligand, AR-v7 in the nucleus of PCa can drive the expression of classical androgen reactive gene and endure castration resistance to prostate cancer. [27][18][23]

During the progression of PCa, especially in hormone-refractory prostate cancer (HRPC), AR-V7 was significantly up-regulated and predicted biochemical recurrence after surgical treatment (P = 0.012). [31][23][27] The increased expression of AR-Vs (AR-V1, AR-V7 and AR-V567es) in bone metastasis in CRPC not only indicates the correlation with advanced bone metastasis of prostate cancer, but also indicates a very poor prognosis. [19]

3.2.1 AR-V7 is related to endocrine therapy for castration-resistant prostate cancer
Compared with hormone-sensitive prostate cancer, AR-V1 and AR-V7 in HRPC were significantly increased in both protein expression and mRNA expression, the mRNA expression of more than 20 times (n = 25), [24][27] especially after treatment with abiraterone or enzalutamide, the expression of ar-v7 was more significant, ar-v7 was associated with decreased response to castration-resistant prostate cancer (CRPC) endocrine therapy and decreased overall survival. [31]

In 168 cases treated with androgen deprivation (ADT) prostate cancer patients showed that the positive expression of AR-V7 significantly reduced the response rate of prostate-specific antigen (PSA) (P < 0.001)to androgen deprivation therapy and significantly reduced progression-free survival (P < 0.0001)and overall survival of castration-resistant prostate cancer(HR: 4.826; 95% CI: 2.960-7.869; P < 0.001). [7]

Among the patients with metastatic castration resistant prostate cancer treated with enzalurudine or abirutone, the PSA response rate, prostate specific antigen progression-free survival(PSA PFS) and overall survival (OS) of AR-V7-positive patients were significantly lower than those of AR-V7-negative control group. [31][30][34] In the enzalutamide group, AR-V7-positive patients had a PSA response rate (0% vs. 53%, P = 0.004), PSA PFS (median 1.4 months vs. 6.0 months,P <0.001), no clinical or radiological progression (median 2.1 months vs. 6.1 months,P <0.001), and OS (median 5.5 months vs. unmet,P = 0.002), relative to AR-V7-negative patients. Also in treatment of male abiraterone, AR-V7-positive patients had a PSA response rate (0% vs. 68%, P = 0.004), PSA PFS (median 1.3 months vs. unmet,P = 0.001), no clinical or radiological progression (median 2.3 months vs. unmet,P <0.001) and OS (median 10.6 months vs. unmet,P = 0.006), relative to AR-V7-negative patients. In patients with metastatic castration-resistant prostate cancer treated with enzalutamide or abirutone, AR-V7- is not only an independent predictor of PSA response rate and PFS, [30] but is also associated with resistance to enzalutamide or abirutone in castration-resistant prostate cancer. [30][28][34]

3.2.2 AR-V7 is related to taxane chemotherapy in castration-resistant prostate cancer
Although AR-V7 expression of Circulating Tumor Cells (CTC) in metastatic castration-resistant prostate cancer(mCRPC) was not associated with major resistance to taxane chemotherapy, [35] it was shown in 37 metastatic CRPC patients treated with taxane (docetaxel or cabataxel) and 62 CRRP patients treated with enzalutamide or abirutone: In the AR-V7 negative patients, there was no difference in the outcomes of the three treatments, while in the AR-V7-positive patients, the PSA response rate, PSA PFS and PFS of the taxanes treatment group were all higher than that of the enzalutidine treatment group or the abiraterone treatment group,with Prostate-specific antigen responses (41% vs 0%, P <0.001), PSA PFS hazard ratio [HR] 0.19[95%ci, 0.07-0.52] P = 0.001, PFS HR 0.21[95%ci, 0.07-0.59], P = 0.003), [35] when positive AR-V7 was detected in circulating tumor cells (CTCs) before treatment, the risk of death of choosing taxanes was lower than that of using androgen receptor signaling (ARS) inhibitors (HR, 0.24; 95% CI, 0.10-0.57; P = 0.035)[29] In summary, the clinical outcome of taxanes was better than that of...
enzalutamide or abiridine, and AR-V7 detection could be used as a treatment biomarker in CRPC. [35] AR-v7 is also a dynamic marker in abiridine, enzalutamide endocrine therapy, or taxane chemotherapy. "AR-V7 can be transformed into a positive or negative state", and the continuous AR-V7 test in blood provides a method to observe the evolution of tumors in real time. [33]

3.2.3 Mechanism of AR-V7 induced castration resistance in CRRP
Under the condition of ADT enhancement, increased AR binding to the pre-mRNA and splicing factor, increase RNA splicing enhancer and its binding protein (U2AF65 and ASF/SF2) to AR - V7 3'splice sites raise, lead to the increase of AR - V7 level in prostate cancer cells. [24] Although the expression of AR - V7 protein relative to AR - FL is low, the low level of AR - V7 are not sufficient to restore the AR activity, but after androgen deprivation induced by fast, [26] the increased ar-v7 was mediated by AR-fl to endure ligand-independent trans-activation of AR, [32] enabling the tumor to retain the basic AR activity needed for survival until a more effective mechanism for AR activation was arise.[26] AR3 may directly raise AKT1 expression, [23] regulate a variety of tumor autocrine/paracrine factor (Tgf beta2 and Itg1), raised some epithelium - interstitial conversion related genes such as single or superposition approach to driving of castration resistant prostate cancer. [25] In CRRP, the expression of ar-v7 is positively correlated with UBE2C. [14]

3.3 AR4, AR5 and AR45
AR4 and AR5 lacking ligand-binding domains are two novel AR slection variants in hormone-insensitive prostate cancer cells. [23]AR45 is a natural variant of the human androgen receptor (AR). AR45 is mainly expressed in the heart and skeletal muscle. AR45 can inhibit the function of AR and reduce the proliferation rate of androgen-dependent LNCap cells. [36]

3.4 ARV1, ARV9, ARV12, ARV13 and ARV14
ARV1 and ARV9 are similar in structure, both of which do not possess the basic amino acids of the two-part nuclear localization sequence, and can be conditionally activated according to the level of androgens. ARV13 and ARV14 had no functional effect, and ARV12 had constitutive activity. The expression levels of ARV12 and ARV9 in CRPC specimens were significantly increased, and ARV12 was positively correlated with patients with Gleason score of 8 or above (P = 0.017). [37]

3.5 AR8
AR8 is a new AR splicing variant, mainly located on the plasma membrane, upregulation in castration-resistant prostate cancer cells. AR8 promotes the binding of Src and AR to the EGF receptor in response to EGF treatment and enhances the tyrosine phosphorylation of AR to promote the progression of prostate cancer in the context of androgen deprivation.

[38]

Conclusion
At present, significant breakthroughs have been made in the research on the mechanism and treatment of prostate cancer, especially in the aspects of endocrinology and chemotherapy, which have improved the overall survival time and quality of life for patients. However, when prostate cancer inevitably progresses to the castration resistance stage, the prognosis is still extremely poor. [7] In CRRP, AR-Vs (AR-V7,AR-V567es) was significantly correlated with overall survival, progression-free survival, postoperative biochemical recurrence, and advanced bone metastasis. This paper preliminarily demonstrated the mechanism of AR-Vs in the treatment of abiraterone and enzalutamide resistance in castration-resistant prostate cancer, and also provided directional selection for the treatment of abiraterone, enzalutamide and taxo. [29][35] However, the specific mechanism of ar-vs in CRRP still needs to be further studied. It is expected that ar-vs can be used as a therapeutic target to develop effective target inhibitors with little side effects, so as to improve the outcome of poor prognosis of CRRP and improve the overall survival and quality of life of patients. AR-Vs is the key research direction in the future.[16][17]

The authors declare no conflict of interest.

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