Dysregulation of spliceosome gene expression in advanced prostate cancer by RNA-binding protein PSF

Ken-ichi Takayama, Takashi Suzuki, Tetsuya Fujimura, Yuta Yamada, Satoru Takahashi, Yukio Homma, Yutaka Suzuki, and Satoshi Inoue

*Department of Functional Biogerontology, Tokyo Metropolitan Institute of Gerontology, Tokyo 173-0015, Japan; †Department of Pathology and Histotechnology, Tohoku University Graduate School of Medicine, Miyagi 980-8575, Japan; ‡Department of Urology, Graduate School of Medicine, The University of Tokyo, Tokyo 113-8655, Japan; §Department of Urology, Nihon University School of Medicine, Tokyo 173-0032, Japan; ‡Department of Medical Genome Sciences, Graduate School of Frontier Sciences, The University of Tokyo, Chiba 277-8562, Japan; and Division of Gene Regulation and Signal Transduction, Research Center for Genomic Medicine, Saitama Medical University, Saitama 350-1241, Japan

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Developing therapeutic approaches are necessary for treating hormone-refractory prostate cancer. Activation of androgen receptor (AR) and its variants’ expression along with the downstream signals are mostly important for disease progression. However, the mechanism for marked increases of AR signals and its expression is still unclear. Here, we revealed that various spliceosome genes are aberrantly induced by RNA-binding protein PSF, leading to enhancement of the splicing activities for AR expression. Our high-speed sequence analyses identified global PSF-binding transcripts. PSF was shown to stabilize and activate key long noncoding RNAs and AR-regulated gene expressions in prostate cancer cells. Interestingly, mRNAs of spliceosome-related genes are putative primary targets of PSF. Their gene expressions are up-regulated by PSF in hormone-refractory prostate cancer. Moreover, PSF coordinated these spliceosome proteins to form a complex to promote AR splicing and expression. Thus, targeting PSF and its related pathways implicates the therapeutic possibility for hormone-refractory prostate cancer.

androgen receptor | RNA-binding protein | PSF | NONO | prostate cancer

A ndrogen receptor (AR) regulates many genes central to the identity and behavior of prostate cancer cells (1). AR functions in a ligand-dependent manner in many prostate cancers, and androgen deprivation therapy is effective in inhibiting tumor growth. However, most of the patients acquire resistance to this therapy and eventually suffer from castration-resistant prostate cancer (CRPC) (2, 3). Previous studies have discovered the importance of enhanced AR downstream signaling (3, 4), expression of AR and its splice variants lacking the ligand binding domain (ARVs) in CRPC (5, 6). The variant AR-V7 was shown to regulate distinct androgen-independent activation of its downstream signals, which contributes to the development of CRPC (7). Thus, targeting AR, AR-V7, and its downstream signals could have efficacy in treating CRPC.

Long noncoding (Inc) RNAs function through interaction with epigenetic factors in cancer (8–11). Previous reports highlighted the lncRNA-mediated association of RNA-binding proteins or transcription factors with specific genomic regions for prostate cancer progression (10, 11). We have reported that androgen-induced lncRNA (named CTBP1-AS) in the antisense region of carboxyl terminal binding protein 1 (CTBP1) promotes castration-resistant tumor growth. CTBP1-AS interacts with an RNA-binding transcriptional and splicing factor, splicing factor proline and glutamine-rich (PSF/SFPQ), and represses cell cycle regulators by epigenetic mechanism (11). We also found that PSF and its interacting factor, non–POU-domain–containing octamer-binding protein (NONO), have an important role in prostate cancer cell proliferation, suggesting the possibility that these RNA-binding proteins are also implicated in the disease progression (11).

Results
PSF Promotes Castration-Resistant Prostate Tumor Growth. To investigate the role of PSF in prostate cancer, we analyzed the expression level of PSF. In prostate cancer cell lines, we observed increased expression of PSF in several prostate cancer cells such as DU145, LNCaP, and LNCaP-derived CRPC model cells, long-term androgen deprivation (LTAD) cells compared with

Significance
Elevated downstream signals of androgen receptor (AR) and its variants are important for prostate cancer progression. We show that an RNA-binding transcriptional and splicing factor, splicing factor proline and glutamine-rich (PSF/SFPQ), predicts worse prognosis of prostate cancer patients. Inhibition of PSF expression repressed treatment-resistant prostate tumor growth in our animal model. Our global analysis of PSF-binding RNAs revealed that PSF enhances AR-regulated genes and noncoding RNAs associated with prostate cancer progression. Interestingly, various splicing factors, which are primary targets of PSF, are upregulated in metastatic prostate tumors. These enhanced factors form complexes with PSF to promote AR expression and splicing. Our findings suggest a role of RNA-binding protein for AR activation for prostate cancer progression.

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1To whom correspondence should be addressed. Email: sinoue@tmig.or.jp.

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PNAS Early Edition | 1 of 6
normal prostate epithelial cells (PrEC) (SI Appendix, Fig. S1A). We evaluated the PSF protein expression in clinical tumor samples by immunohistochemical (IHC) analysis (Fig. L4) using anti-PSF specific antibody (SI Appendix, Fig. S1B). Low PSF staining was observed in benign prostate tissues, whereas high PSF staining was observed in a subset of tumor samples, and higher expression of PSF correlated with the patients’ cancer-specific survival after surgery and the PSA-free survival after hormone therapy (Fig. 1B and SI Appendix, Fig. S1C). According to public databases of expression profiles in prostate cancer clinical samples (such as Oncomine) (17–19), PSF mRNA expression is significantly elevated in metastatic or advanced prostate cancer samples, suggesting that PSF expression is associated with prostate cancer progression (Fig. 1C and SI Appendix, Fig. S1 D–F).

We previously reported that PSF-inhibited cell cycle-associated gene such as p53 and SMAD3 expression by transcriptional mechanism in AR-dependent prostate cancer cell lines (LNCaP, VCaP) and their CRPC models, LTAD cells (11). In this study, by using short interference (si) RNAs targeting PSF, we efficiently depleted the expression of PSF in other hormone-refractory prostate cancer cells such as DU145 and 22Rv1 cells (SI Appendix, Fig. S2A). We then observed the inhibited cell growth of these prostate cancer cells by siPSF treatment (SI Appendix, Fig. S2B). Next, we examined the roles of PSF using in vivo xenograft assays. Injection of PSF siRNA into the tumors significantly inhibited tumor growth derived from LNCaP cells (SI Appendix, Fig. S2 C and D). As for in vivo model of CRPC, we injected 22Rv1 cells into nude mice s.c., castrated the mice after the tumors formed, followed by injection of siPSF or siControl into the tumors. Interestingly, siPSF treatment reduced the growth of tumors derived from 22Rv1 cells (Fig. 1D). In tumors with reduced PSF expression, induced cell cycle regulators such as p53 and p21 proteins were also observed, suggesting that PSF could be a potential target for the treatment of CRPC (Fig. 1E). These data highlight a potential role of increased PSF in the CRPC tumor growth. (Additional results are shown in SI Appendix, Text S1 and Figs. S3–S5.)

**Global Mapping of PSF-Binding Transcripts Revealed That AR-Regulated Genes and Prostate Cancer-Associated IncRNAs Are Specific Targets of PSF.** To explore other functions of PSF in prostate cancer, we next analyzed RNAs bound with PSF using deep-sequence analysis (Fig. 2A). First, we performed RNA immunoprecipitation coupled with deep sequencing (RIP-seq) in LNCaP cells treated with vehicle or DHT. By mapping sequence tags to GenCODE, and NONCODE, we identified putative PSF target genes (Fold > 2, P < 10−7) including IncRNAs (Fig. 2B and SI Appendix, Fig. S6 A–C). We further performed RIP-seq in CRPC model cells (LTAD and 22Rv1) (Fig. 2C). In LTAD and 22Rv1 cells, we found more binding genes, which significantly overlapped with those of LNCaP cells. Interestingly, CTBP1-AS1 was identified as one of the most significant IncRNAs in this comprehensive study among NONCODE and RefSeq antisense transcripts (SI Appendix, Fig. S6B), suggesting the specific interaction between PSF and CTBP1-AS1 in LNCaP cells treated with DHT. In addition, we observed that DHT treatment changed the binding transcripts of PSF. We found that androgen-regulated genes such as C4MKK2 (20) are enriched among DHT-specific target genes of PSF (Fig. 2B). In contrast to the PSF targets identified by ChiP-seq (SI Appendix, Fig. S4A), pathway analysis showed that spliceosome genes are assumed to be the primary targets of PSF at RNA level in LNCaP cells (SI Appendix, Fig. S6D), suggesting this gene cluster is specifically regulated by PSF as an RNA-binding protein. Prostate cancer-associated signals were also significantly included in these pathways. We next performed crosslinking immunoprecipitation (CLIP)-seq (21, 22) in LNCaP cells to identify the peak positions of PSF bindings (Fig. 2D). PSF-binding sites were mainly distributed in the intron regions (Fig. 2E) of either protein-coding or noncoding genes (SI Appendix, Fig. S6E). Motif analysis (MEME) (23) revealed that GA-rich motifs were evident in the binding peaks of PSF (SI Appendix, Fig. S6F). We then identified putative PSF-binding genes including annotated RefSeq genes (Vehicle: 643 DHT: 495 genes) and non-coding genes among them (Vehicle: 173 DHT: 125) (Fig. 2F and SI Appendix, Fig. S6G). Different lists of PSF-binding genes in the absence and presence of DHT treatment implicated the involvement of AR in the PSF bindings to RNA. In fact, we found that transcripts bound with PSF in the presence of DHT were mainly derived from DHT-induced genes (Fig. 2G). Conversely, transcripts bound with PSF in the absence of DHT were derived from repressed genes by DHT. The list of PSF-binding genes obtained by CLIP-seq showed significant but partial overlap with those by RIP-seq, probably due to the different techniques (Fig. 2H). Those PSF-binding genes were validated by RIP assays (SI Appendix, Fig. S6H). Moreover, we found that PSF-binding genes in the presence of DHT were up-regulated in prostate cancer according to The Cancer Genome Atlas (TCGA) database (SI Appendix, Fig. S6I), suggesting the importance of PSF interaction with transcripts for prostate cancer progression.

To understand how PSF affects the binding transcripts, we performed directional RNA-seq to investigate the gene regulation by PSF. These results revealed that PSF mainly increases the expression of its binding transcripts (Fig. 2I). In addition, knockdown of PSF resulted in a marked decrease in DHT-induced expression of AR target genes (Fig. 2I). For IncRNAs, we observed both negative [NEAT1, MALAT1 (9, 24, 25)] and positive [CTBP1-AS1, SchlAP1 (26)] effects on expression in LNCaP cells (Fig. 2K). These effects were also confirmed by quantitative reverse transcription PCR (qRT-PCR) in both LNCaP
PSF Binding to AR Transcripts Is Enhanced in Hormone-Refractory Prostate Cancer Cells for Activating Their Expression and Splicing.

Next, we investigated the PSF function in CRPC model cells. Importantly, the pathway analysis of PSF-binding genes in 22Rv1 cells indicated the importance of PSF in induction of spliceosome genes, as observed in LNCaP cells (Fig. S7 A and B). These results suggest that prostate cancer-associated signals were also enhanced. For example, the binding of PSF to the introns of SchLAP1 and AR (Fig. S8 A and SI Appendix, Fig. S10 B–D) was enhanced in LTAD and 22Rv1 cells compared with LNCaP cells, suggesting the role of PSF in expressing these important signals in CRPC (SI Appendix, Fig. S10 E).

We then investigated the role of PSF in AR splicing and posttranscriptional regulation. We have detected CLIP signals in the intron and 3′-UTR regions of AR gene, suggesting the involvement of PSF in AR mRNA stability and splicing (SI Appendix, Fig. S10 F). First, we confirmed the binding of PSF with androgen (TTS, terminal transcription site; 5′UTR, 5′ untranslated region; 3′UTR, 3′ untranslated region). The number of significant PSF-binding genes annotated in RefSeq is shown. (D) Representative mapping of PSF-binding RNAs in the CTBP1-AS locus. We present the results of RIP-seq, CLIP-seq, and ChIP-seq (AR, AcH3) in LNCaP cells. Arrows indicate the direction of CTBP1 and CTBP1-AS. (E) Genomic location of the identified peaks in the presence and absence of androgen. TTS, spliceosome genes, and AR-regulated genes by binding to PSF. PSF-binding genes were annotated by CLIP-seq and RIP-seq. (F) The difference of PSF-binding genes identified by CLIP in the absence and presence of DHT. (G) PSF binds to androgen-induced genes significantly in the presence of DHT. The number of PSF-binding genes was counted, and χ² test was performed. (H) Overlap of identified genes by CLIP-seq with RIP-seq. (I) Regulation of PSF-binding genes in prostate cancer cells. LNCaP cells were treated with siControl or siPSF#1. After 48 h incubation, cells were treated with vehicle or DHT for 24 h. PSF-binding genes identified by RIP-seq were selected, and changes in their expression levels by PSF knockdown are summarized. (J) PSF regulates androgen-dependent gene activation. Heatmap results of RNA-seq in androgen-regulated genes are shown. (K) Regulation of lncRNAs by PSF binding. PSF-binding lncRNAs identified by CLIP-seq and RIP-seq are selected. (L) The ratio of miRNAs and lncRNAs in PSF-binding lncRNAs identified by CLIP. (M) Expressions of androgen-induced miRNAs (miR-125b2, miR-99a, and miR-21) were repressed by knockdown of PSF. LNCaP cells were treated with siControl, siPSF#1, or siPSF#2 for 48 h. We then measured mRNA expression levels by qRT-PCR in cells after treatment of vehicle or DHT for 24 h (n = 3). (N) miRNA stability of SchLAP1 was decreased by knockdown of PSF. LNCaP cells are treated with siControl, siPSF#1, or siPSF#2 for 48 h. To inhibit transcription, actinomycin-D (1 μM) was added. After incubation for indicated times, miRNA expression levels were measured by qRT-PCR (n = 3). Values represent the mean ± SD. *P < 0.05, **P < 0.01.

and VCaP cells (SI Appendix, Fig. S7 A and B). Interestingly, we found that lncRNAs overlapping with miRNAs (such as miR-99A/HG) were also included in the list of genes with PSF CLIP peaks (Fig. 2L). We then performed miRNA-qPCR, showing that the AR-regulated lncRNAs (27) were repressed by PSF knockdown. This suggests that production of androgen-mediated miRNAs could also be modulated by PSF binding to its primary transcripts (Fig. 2M and SI Appendix, Fig. S7C). To investigate the mechanism of PSF for up-regulation of its target genes, we analyzed transcript stability by incubating cells with actinomycin-D. Consistent with the result of RNA-seq, we observed reduced half-lives in these PSF-binding genes in VCaP and LNCaP cells by PSF knockdown (Fig. 2N and SI Appendix, Fig. S8). Thus, PSF was assumed to control the gene expression of coding and noncoding signals associated with prostate cancer, spliceosome genes, and AR-regulated genes by binding to these gene transcripts to increase stability. (Additional results are shown in SI Appendix, Text S2 and Fig. S9.)
Fig. 3. PSF bindings to the target transcripts are enhanced in CRPC. (A) Pathway analysis of PSF-binding genes in 22Rv1 cells is shown. (B) Representative mapping of PSF-binding NRAs in AR locus. (C) Expression levels of PSF, NONO, AR, and AR-V7 in prostate cancer cells. Lysates from LNCaP, LTAD, VCaP, and 22Rv1 cells were used for immunoblots with indicated antibodies. (D) Regulation of AR and AR-V7 expression by PSF and NONO in CRPC model cells. (Left) The 22Rv1 cells were treated with siPSF or siNONO for 72 h. AR and AR-V7 mRNA levels were measured by qRT-PCR (n = 3). (Right) Lysates from 22Rv1 cells transfected with siPSF or siNONO are used for immunoblots to detect AR and AR-V7 protein levels. (E) mRNA stability of AR and AR-V7 was decreased by knockdown of PSF. The 22Rv1 cells are treated with siControl, siPSF #1, or siPSF #2 for 48 h. To inhibit transcription, actinomycin-D (1 nM) was added. After incubation for indicated times, mRNA expression levels were measured by qRT-PCR (n = 3). (F) Immunohistochemistry of AR-V7 in prostate cancer tissues (n = 102) was performed. (G) AR-V7 expression (high, n = 18; low, n = 88) is a strong prognostic factor for prostate cancer patients. Cases with labeling index (LI) > 10% were determined to be AR-V7 high expression. (H) Positive correlation of PSF expression with AR and AR-V7 levels in prostate cancer tissues. LI in each group is shown. (I) NONO as well as PSF was up-regulated in metastatic prostate cancer tissues. Heatmap results using microarray database (GSE35988) is shown. (J) Immunohistochemistry of NONO in prostate cancer tissues (n = 102). (K) NONO expression (high, n = 51; low, n = 51) is a prognostic factor of poor outcome of prostate cancer patients. Values represent the mean ± SD. *P < 0.05, **P < 0.01.

AR and AR-V7, an AR-splicing variant associated with CRPC development, by RIP assays (SI Appendix, Fig. S10 B and G). In addition, Western blot analyses showed that AR-V7 protein expression was detected in VCaP and 22Rv1 cells (Fig. 3C). Interestingly, PSF expression was also up-regulated in these cell lines compared with LNCaP and LTAD, in which AR-V7 protein could not be detected. To determine whether PSF induces AR-V7 protein production or not, we performed qRT-PCR and Western blot analysis (Fig. 3D) by using siRNAs targeting PSF and its associated factor, NONO. Surprisingly, PSF knockdown in 22Rv1 cells inhibited full-length AR and AR-V7 mRNA and protein dramatically, whereas AR mRNA (SI Appendix, Fig. S11A) and protein (11) levels were not affected by PSF knockdown in LNCaP cells. Knockdown of NONO also effectively reduced the expression of AR-V7 as well as AR at mRNA and protein levels in 22Rv1 cells (Fig. 3D). Analysis of mRNA stability of AR and AR-V7 by inhibiting transcription indicated that PSF and NONO are involved in the posttranscriptional regulation of these mRNAs (Fig. 3E and SI Appendix, Fig. S11B). We also observed similar results in VCaP cells (SI Appendix, Fig. S11 C–E). As expected, our AR-V7 IHC analysis in clinical samples has shown that AR-V7 expression predicts poor prognosis of prostate cancer patients (Fig. 3 F and G). Importantly, AR-V7 immunoreactivity score is correlated with that of PSF expression level, suggesting the role of PSF in the production of AR-V7 protein (Fig. 3H). Microarray dataset (Fig. 3I) and our IHC analysis (Fig. 3J) revealed that expression of NONO was also elevated in the metastatic prostate cancer tissues and that high expression of NONO was associated with AR-V7 expression (SI Appendix, Fig. S11F) and poor prognosis of patients (Fig. 3K), suggesting the potential role of NONO in prostate cancer progression and AR-V7 expression by interaction with PSF.

Spliceosome Genes Are Primary Targets of PSF and Aberrantly Expressed in Hormone-Refractory Prostate Cancer. We next hypothesized that elevated expression of spliceosome genes by PSF binding might enhance splicing activity to induce oncogenic effects. Clinically, most of PSF-targeted spliceosome genes are up-regulated in metastatic prostate cancer tissues for further analysis. First, we observed up-regulation of spliceosome genes in addition to the up-regulation of AR and AR-V7 in LNCaP cells overexpressing PSF and NONO that we established previously (13) (Fig. 4B), indicating the role of PSF and NONO in these elevated expressions. We confirmed the up-regulation of these spliceosome genes in AR-V7 positive, VCaP, and 22Rv1 compared with LNCaP cells (Fig. 4C), while PSF and NONO knockdown repressed the expression of these spliceosome genes (Fig. 4D). Knockdown of these spliceosome genes inhibited AR and AR-V7 expression (Fig. 4E and SI Appendix, Fig. S12 A–C). Thus, these results revealed that PSF could increase the expression of spliceosome genes to regulate for AR/AR-V7 production.

We further investigated whether up-regulation of spliceosome genes could enhance splicing activity cooperating with PSF and NONO in CRPC. Our immunoprecipitation and Western-blotting analysis indicated that PSF interacts with these spliceosome components and was responsible for the complex formation of spliceosome (Fig. 4F). In addition, immunofluorescence analysis showed that this interaction occurred in the nucleus (Fig. 4G). RIP assay also indicated that these factors bind to AR transcripts and act as an integrator for PSF-mediated splicing of AR transcripts (Fig. 4H). Taken together, these results revealed that PSF also orchestrated its target splicing factors at protein levels to form a complex for splicing and protein expression in CRPC.

Finally, we evaluated the clinical significance of these PSF target genes in CRPC development and progression. By qRT-PCR analysis of PSF, AR-V7, and spliceosome genes in prostate cancer tissues and tissues obtained from CRPC patients including metastatic tissues (bone, lymph node, and liver), we found that all tested spliceosome genes were up-regulated in prostate cancer compared with the benign regions. Interestingly, several spliceosome genes are up-regulated in CRPC tissues (SI Appendix, Fig. S12D). Among those, we found expression of U2AF2, DDX23, CHERP, and HNRNPU was elevated in CRPC. AR-V7 expression was also correlated with PSF, consistent with the result of IHC (Fig. 3H). In addition, knockdown of these factors showed effective growth inhibition of 22Rv1 cells, particularly in cells where CHERP, HNRNPU, DDX23, SF3B3, and U2AF2 (28–32) were knocked down (Fig. 4I and SI Appendix, Fig. S12E). Overall, our findings suggest that aberrant overexpression of spliceosome genes in prostate cancer by PSF contributes to the progression from hormone-naïve prostate cancer to CRPC (Fig. 4I).

Discussion

In the present study, we demonstrate the importance of PSF in the progression of prostate cancer using deep-sequence–based approach, clinicopathological analysis, and public database. We explored a mechanistic insight by identifying global RNAs bound with PSF. We found that some lncRNAs and miRNAs are
important targets of PSF. Our analysis revealed that CTBP1-AS and SchLAP1 are positively regulated lncRNAs by PSF in prostate cancer cells. Interestingly, SchLAP1 has been known to be overexpressed in metastatic prostate tumors (26). The most unique finding of this study is that spliceosome genes are almost uniformly up-regulated in metastatic CRPC tissues. Importantly, our cell model-based analyses indicate that PSF is responsible for the expression regulation of these spliceosome genes. In addition, our experimental data indicate that PSF coordinates these factors to regulate PSF target genes.

It is notable that one of the most important targets of PSF in CRPC would be AR. Our RIP-seq analysis demonstrated that enhanced association of AR transcript with PSF in CRPC model, 22Rv1 cells, compared with LNCaP cells (Fig. 3B). This might be caused by upregulated PSF expression as shown by Western blot and qRT-PCR analyses (Fig. 3C). Thus, impact of PSF knockdown on AR expression level may be evident in 22Rv1 cells that express PSF abundantly. Since the AR and AR-V7 have been assumed as the driver for the hormone-refractory state (5–8), the up-regulation of PSF would be responsible for the up-regulation of AR and AR-V7 by aberrant splicing activity for CRPC development. We observed PSF forms a complex with other splicing factors and NONO in the intronic region of AR transcripts by RIP and Western blotting following immunoprecipitation analysis. In addition, AR mRNA stability was decreased by PSF knockdown (Fig. 3E), although PSF recruitment to the promoter region of AR was not detected in ChIP-seq analysis. Therefore, these findings suggest that PSF regulates AR-splicing process and promotes production of AR and its variants at mRNA level. Moreover, our comprehensive analysis unveiled the variety of PSF-binding transcripts associated with cancer development in addition to AR. By regulating spliceosome gene expressions and cooperating with those factors, PSF could activate such a broad range of oncogenic pathways as an RNA-binding protein. In the future study, it will be interesting to further analyze specific targets of each splicing factor for revealing the functions of PSF complexes in cancer progression.

Our clinicopathological analyses also demonstrated the importance of NONO, showing the higher expression of NONO correlated with poor prognosis of patients. Our cell model study showed that knockdown of NONO in 22Rv1 and VCaP cells inhibits the expression of AR-V7 more effectively than full-length AR, suggesting the cooperative function of RNA-binding protein complex including PSF and NONO in prostate cancer. These functions of RNA-binding proteins could be effective to enhance the expressions of prostate cancer-associated genes as well as AR.

We indicated the importance of PSF in the regulation of splicing machinery in the progression of aggressive prostate cancer. Our finding revealed a mechanism in which a broad range of splicing components were aberrantly regulated for cancer progression. The wide-ranging up-regulation of the splicing pathway in metastatic prostate cancer could affect the splicing complexes in cancer. RNA
splicing is mediated by an assembly, rearrangement, and disengagement of a set of small nuclear ribonucleoprotein (snRNPs) complexes (U1, U2, and either U4/5/6 or U11/U12) or other proteins onto the pre-mRNAs (33, 34). The broad coverage of the wide spectrum of these components in our study suggests the involvement of PSF in enhancing this machinery. Interestingly, frequent mutations of splicing pathway have been reported in several malignancies by whole-exome sequencing (35, 36). These clinical analyses provide an intriguing insight into the mechanism of cancer progression by splicing machinery, since RNA splicing system has essential cellular roles for the diversity of protein species using a limited number of genes. Loss of splicing activity by mutations would induce severe developmental abnormality and tumorigenesis (37). Conversely, such mutation has been rarely identified in prostate cancer (18, 38). Enhanced splicing machinery could be indispensable for the cancer progression in some types of malignancies (39). Moreover, the present study would illustrate the specificity of PSF function and splicing machinery in prostate cancer.

In summary, our global analysis of PSF functions revealed its target signals based on the RNA-binding ability in addition to cell cycle regulator control by epigenetic silencing of the transcription with histone modification. Importantly, we proposed a mechanistic link between RNA-binding proteins and AR as well as prostate cancer-associated signals, indicating the clinical and biological impact of PSF in the development of abnormal splicing machinery in hormone-refractory prostate cancer. Considering the potential function of AR and AR variants in the development of metastatic CRPC, PSF and its associated factors could have the potential for therapy targets.

Materials and Methods

All animal experimental protocols were performed in accordance with the guidelines of the Animal Ethics Committee of the University of Tokyo. Further details are provided in SI Appendix, SI Materials and Methods.

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