MicroRNA-regulated transcriptome analysis identifies four major subtypes with prognostic and therapeutic implications in prostate cancer

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Abstract

MicroRNA (miRNA) deregulation plays a critical role in the heterogeneous development of prostate cancer (PCa) by tuning mRNA levels. Herein, we aimed to characterize the molecular features of PCa by clustering the miRNA-regulated transcriptome with non-negative matrix factorization. Using 478 PCa samples from The Cancer Genome Atlas, four molecular subtypes (S-I, S-II, S-III, and S-IV) were identified and validated in two merged microarray and RNAseq datasets with 656 and 252 samples, respectively. Interestingly, the four subtypes showed distinct clinical and biological features after comprehensive analyses of clinical features, multiomic profiles, immune infiltration, and drug sensitivity. S-I is basal/stem/mesenchymal-like and immune-excluded with marked transforming growth factor β, epithelial-mesenchymal transition and hypoxia signals, increased sensitivity to olaparib, and intermediate prognosis. S-II is luminal/metabolism-active and responsive to androgen deprivation therapy with frequent TMPRSS2-ERG fusion and a good prognosis. S-III is characterized by moderate proliferative and metabolic activity, sensitivity to taxane-based chemotherapy, and intermediate prognosis. S-IV is highly proliferative with moderate EMT and stemness, frequent deletions of TP53, PTEN, and RB, and the poorest prognosis; it is also immune-inflamed and sensitive to anti-PD-L1 therapy. Overall, based on miRNA-regulated gene profiles, this study identified four distinct PCa subtypes that could improve risk stratification at diagnosis and provide therapeutic guidance.

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Abbreviations: PCa, prostate cancer; mCRPC, metastatic castration-resistant prostate cancer; TCGA, The Cancer Genome Atlas; miRNAs, microRNAs; MIRcor, miRNA-correlated; SCNAs, somatic copy number alterations; k-NN, k-nearest neighbor; GEO, Gene Expression Omnibus; DFS, disease-free survival; OS, overall survival; BCR, biochemical recurrence; NMF, non-negative matrix factorization; FDR, false discovery rate; NTP, Nearest template prediction; SubMap, Subclass mapping; DEmiRs, differentially expressed miRNAs; GSEA, Gene Set Enrichment Analysis; GEP, gene expression profile; TNAs, tumor neoantigens; TMB, tumor mutation burden; KEGG, Kyoto Encyclopedia of Genes and Genomes; GO, Gene Ontology; ssGSEA, single-sample gene set enrichment analysis; CCLs, cancer cell lines; AUC, Area under the dose-response curve; CTLA-4, cytotoxic T-lymphocyte associated protein-4; PD-1, programmed cell death protein-1; PD-L1, programmed death-ligand 1; TGFb, transforming growth factor β; AR, androgen receptor; NEPC, neuroendocrine prostate cancer; EMT, epithelial-mesenchymal transition; IFN, interferon; MDSCs, myeloid-derived suppressor cells; Tregs, regulatory T cells; CAFs, cancer-associated fibroblasts; ADT, androgen deprivation therapy; ICB, immune checkpoint blockade.

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1. Introduction

Prostate cancer (PCa) is the second-most (14.1%) commonly diagnosed cancer and fifth (6.8%) leading cause of cancer death among men [1]. Despite the high incidence rate, a proportion of localized PCa is relatively indolent with good prognosis and therefore, these patients should be spared from invasive treatments and concomitant complications [2,3]. On the other hand, a subset of PCa is clinically significant and aggressive, and tends to progress into metastatic castration-resistant prostate cancer (mCRCPC) with limited therapeutic options [4,5]. Early identification of this subset of patients will allow for more active interventions and close follow-up, which could theoretically reduce relapse and prolong survival. Attributed to the inherent genetic heterogeneity of PCa [6], widely used risk stratification tools including serum prostate-specific antigen level, TNM and Gleason score often fail to pinpoint PCa patients with either aggressive or indolent forms of the disease [7–9]. As a result, these patients are at risk of undertreatment or overtreatment [2]. To address this challenge, genomic profiling has been exploited to characterize the molecular heterogeneity of PCa and has been proven to increase the accuracy of risk predictions in clinical practice [10,11]. Recently, The Cancer Genome Atlas (TCGA) used somatic mutations or transcript fusions to define PCa subtypes [12]. However, this classification fails to discriminate patients with distinct prognoses, and therefore, its use in routine clinical practice is severely compromised. Additionally, most published molecular classifications did not investigate whether different molecular subtypes respond differently to conventional therapies [12–15,10], which limited their application into clinical practice.

MicroRNAs (miRNAs) are a group of small non-coding regulatory RNA molecules that repress mRNA translation or promote degradation by binding to targeted mRNAs [16]. Mounting evidence suggests that miRNA aberrations occur frequently in PCa and play key roles in PCa initiation and progression [17,18]. Nevertheless, it remains unclear to what extent miRNA deregulation affects PCa heterogeneity, and thus molecular subtyping and therapeutic response. Herein, using miRNA-correlated (MIRcor) genes, we sought to develop and validate a novel molecular classification that could accurately identify PCa patients with differential prognoses and therapeutic responses. We intended to characterize the identified PCa subtypes at the epigenetic, genomic, transcriptional, immune, and clinical levels. We also sought to determine potential subtype-driven miRNAs in each subtype.

2. Materials and methods

2.1. Data collection and preprocessing

Mature miRNA expression (level 3, Illumina miRNA-Seq), processed mRNA expression (level 3, Illumina RNASeq), 450 K methylation data (level 3, Illumina HumanMethylation450 BeadChip), somatic copy number alterations (SCNAs, level 3, Affymetrix SNP 6.0 array), somatic mutation (level 4, MAF files), and clinical information of PCa patients up to November 2020 were obtained from TCGA database (http://cancergenome.nih.gov/). We filtered out miRNA, mRNA and CpG sites with >25% missing values across all samples and imputed the remaining missing values using the K-nearest neighbor (k-NN) approach in the ‘impute’ R package. We used log2 (normalized count + 1) as the expression levels of mRNA and miRNA. DNA methylation probes were filtered and normalized using the R ‘ChAMP’ package. We only selected CpG sites residing in the promoter region (TSS1500, TSS200, 5’UTR, and 1stExon). Somatic variants were detected using the TCGA MuTect2 pipeline and further analyzed by the R ‘maftools’ package. GISTIC2.0 (Gene Pattern) was used to infer recurrent SCNAs [19]. After removing ineligible samples, 478 treatment-naïve PCa patients with complete prognostic information were selected for subsequent analyses (Table S1).

For external validation, we retrieved 8 independent datasets with available GEPs and matched prognostic information, including GSE116918 (n = 248), GSE70769 (n = 92), GSE70768 (n = 111), GSE107299 (n = 65), and GSE54460 (n = 91) from the Gene Expression Omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/), and DKFZ-PRAD (n = 81), SU2C-PRAD (n = 80) and MSKCC-PRAD (n = 140) from the cBioPortal database (https://www.cbioportal.org/). Raw expression data from Affymetrix were background adjusted and normalized using the robust multiarray average algorithm (‘affy’ R package). The fragments per kilobase per million values in RNA sequencing cohorts were also transformed into transcripts per million values that were more comparable with the microarray data [20]. Next, the RNAseq- and microarray-based cohorts were merged into two meta-cohorts (Table S2 and Table S3), respectively. Batch effects were removed by ‘combat’ functions in the ‘sva’ R package (Fig. S1). Disease-free survival (DFS), overall survival (OS), and biochemical recurrence (BCR) rates were all considered as endpoint events but were analyzed separately.

2.2. Identification of miRNA-mRNA pairs

A two-step approach was adopted to identify miRNA-correlated genes as in other miRNA studies [21,22]. First, three miRNA databases were used to predict potential mRNA targets of miRNAs in the TCGA-PRAD dataset. Second, we computed the Pearson correlation coefficients of the intersecting miRNA-mRNA pairs, with Benjamini and Hochberg corrections (q-values) to control for multiple testing. We selected only those with q-values < 0.05 and correlation coefficients < -0.3 for downstream analysis because miRNAs have been widely shown and validated to reduce mRNA levels through complementary interactions between their seed sequences and the 3’ untranslated regions of targeted mRNAs [16].

2.3. Development and validation of molecular subtypes

Non-negative matrix factorization (NMF, R ‘NMF’ package) was applied to 570 MiRcor gene expression profiles with the setting of 30 iterations for k = 2 to k = 6. The optimal number of clusters was determined by taking into account the cophenetic coefficient and consensus maps from 2 to 6 clusters.

To validate the classification, we first identified differentially expressed genes among the four subtypes using the R ‘limma’ package. A false discovery rate (FDR) < 0.05 and log2 [fold change] >1 were used as the cutoff criteria when comparing each subtype with the rest. The gene signature for each subtype was defined as the unique and upregulated miRNAs in the corresponding subtype. Next, using subtype-specific signatures, the nearest template prediction (NTP) algorithm [23] (Gene Pattern) was employed to classify PCa patients in two merged datasets. To test the robustness of class assignment, we conducted subclass mapping (SubMap) analyses [24] (Gene Pattern) to evaluate the similarity of molecular subtypes between validation cohorts and the TCGA-PRAD cohort.

2.4. Associations of PCa subtypes with multiomics data in TCGA-PRAD

We first performed differential expression analysis of miRNAs by comparing each subtype to other tumor samples. FDR < 0.05 and log2 [fold change] >0.5 were used to define differentially expressed miRNAs (DEmiRs). We also computed and compared the mean expression of miRNAs of each subtype and adjacent
normal samples. Furthermore, to search for DEmiRs potentially driving PC subtypes, we defined miRNA gene sets using MIRcor genes (correlation coefficients < 0 and q values < 0.05) and conducted Gene Set Enrichment Analysis (GSEA) to determine contributions of candidate miRNAs to gene expression profiles (GEPs) of PCa subtypes. Normalized enrichment score > 1.5 and adjusted P values < 0.05 were used as the cutoff points.

For DNA methylation analysis, we calculated and compared the mean levels of promoter CpG methylation in each subtype. Differentially altered and mutated genes were detected by comparing their distributions among the subtypes. The focal and arm level changes of SCNAs were determined as the total number of altered mRNAs with absolute CNA values>0.2. The predicted aneuploidy score, tumor neoantigens (TNAs) and tumor mutation burden (TMB) data were extracted from a previous study by Thorsson et al. [25].

2.5. Biological characterization of PC subtypes

Kyoto Encyclopedia of Genes and Genomes (KEGG) and Gene Ontology (GO) were analyzed via the ‘clusterProfiler’ R package. Single-sample gene set enrichment analysis (ssGSEA, ‘GSVA’ R package) was performed on molecular pathways from the Molecular Signature Database and curated gene sets representing lineage plasticity, TMPRSS-ERG fusion, various biological processes and the immune microenvironment (Table S4). Meanwhile, we used the R ‘estimate’ package to infer tumor purity and estimate the degree of immune and stromal infiltration.

2.6. Exploration of therapeutic vulnerabilities across PC subtypes

Sequencing data of human cancer cell lines (CCLs) and drug sensitivity data against 482 CCLs were retrieved from the Cancer Cell Line Encyclopedia project (https://portals.broadinstitute.org/ccle/) and the secondary PRISM Repurposing dataset (https://depmap.org/portal/prism/), respectively. Area under the dose–response curve (AUC) values were used to measure drug sensitivity with lower AUC values indicating increased sensitivity. We applied ridge regression models (‘R pkRothic’ package) to estimate the AUC values of all PCa patients. Moreover, we used SubMap analysis to infer the efficacy of enzalutamide, anti-cytotoxic T-lymphocyte associated protein-4 (CTLA-4), anti-programmed cell death protein-1 (PD-1) and anti-programmed death-ligand 1 (PD-L1) therapies in PCa subtypes by evaluating the similarity of GEPs between the PCa subtypes and patients receiving corresponding therapies in previous studies [26–28].

2.7. Statistical analysis

All statistical and bioinformatics analyses were conducted with R software version 4.0.2 and Gene Pattern. Associations between PCa subtypes and categorical data were assessed by Fisher’s exact or chi-square tests. For continuous variables, the Student’s t-test or Wilcoxon rank-sum test was used to compare two groups while the Kruskal–Wallis test or ANOVA was used for more than two groups. Survival analyses were carried out by Kaplan–Meier curves and log-rank tests. A two-sided P value < 0.05 was recognized as statistically significant.

3. Results

3.1. Identifying the miRNA-regulated transcriptional subtypes in PCa

A flow chart was produced to summarize this study (Fig. 1). First, we used three miRNA databases to predict potential miRNA targets of 494 miRNAs among 16,766 mRNAs in 478 PCa samples of the TCGA-PRAD dataset. As shown in Fig. 2A, we obtained 281,308 miRNA-mRNA pairs from 487 miRNAs and 15,186 mRNAs in the miRDB database [29], 91,401 miRNA-mRNA pairs from 494 miRNAs and 12,791 mRNAs in the miRtar database [30], and 453,371 miRNA-mRNA pairs from 489 miRNAs and 16,178 mRNAs in the TargetScan v7.2 database [31], respectively. We then selected 14,046 pairs of miRNAs (n = 461) and mRNAs (n = 4565) that coexisted in the three databases. Eventually, using Pearson correlation coefficients (<-0.3; Fig. 2B) and q-values (<0.05; Table S5), we identified 570 MIRcor genes. KEGG and GO analyses revealed that these MIRcor genes were significantly enriched in epithelial cell proliferation and migration, cell cycle, reproductive system development, PI3K-Akt signaling pathway, p53 signaling pathway, T cell receptor signaling pathway, and transforming growth factor β (TGFβ) signaling pathway (Fig. 2C and Table S6). These results suggested that MIRcor genes might play important and diverse roles in regulating the biological functions of both prostate tumor cells and the tumor microenvironment.

Next, we applied NMF to 570 MIRcor genes and chose k = 4 as the optimal number of clusters according to the cophenetic correlation coefficients and consensus maps (Fig. 2D and Fig. S2). Four subclasses were therefore identified in the TCGA-PRAD cohort (S-I, n = 80; S-II, n = 213; S-III, n = 70; S-IV, n = 115). Kaplan–Meier analysis revealed that the four subclasses had distinct prognostic differences (DFS, p < 0.001; Fig. 2E). S-IV (24.1%) was associated with the poorest prognosis while the largest subtype, S-II (44.6%) showed the best prognosis. The prognoses of S-I (16.7%) and S-III (14.6%) were somewhere in between. These results indicated that the MIRcor gene expression profile could stratify PCa patients into distinct prognostic groups.

3.2. Validation of PCa subtypes across independent datasets

To verify the robustness of the clustering method, differential expression analysis was first conducted to reveal 443 significantly upregulated genes (Table S7) with 260 for S-I, 57 for S-II, 42 for S-III, and 84 for S-IV, as shown in Fig. 2F. Pathway analysis of subtype-specific genes (Fig. S3 and Table S8) showed that oxidative phosphorylation and mitochondrial protein complexes were significantly enriched in S-III and that mesenchymal-related pathways such as extracellular matrix modeling, focal adhesion and muscle contraction were overrepresented in S-I. S-IV-specific genes were more involved in cell division-relevant pathways while S-II-specific genes were significantly associated with secretory and transport processes. These results highlighted differentially active pathways among the four PCa subtypes and further supported the rationale of using MIRcor genes for class discovery.

Next, subtype-specific genes were used to assign class labels to each sample in two merged datasets by the NTP algorithm. SubMap analyses showed that the subtypes in both merged datasets were specifically and significantly correlated with the corresponding subtypes in TCGA-PRAD (all p < 0.01, Fig. 3A). Moreover, similar prognostic differences were verified in the two merged datasets (BCR, p < 0.001; Fig. 3B). All independent cohorts demonstrated similar results (GSE54460: BCR, p = 0.001; GSE70768: BCR, p < 0.001; GSE70796: BCR, p < 0.001; GSE107299: BCR, p = 0.047; GSE116918: BCR, p = 0.002; DKKFZ-PRAD: BCR, p < 0.001; MSKCC-PRAD: DFS, p < 0.001; SU2C-PRAD: OS, p = 0.002; Fig. 3C).

3.3. Associations of the PCa subtypes with miRNA, DNA methylation, somatic copy number alterations and mutations

We first analyzed the miRNA patterns among four PCa subtypes. Compared with normal samples, the four subtypes showed
higher levels of mean miRNA expression (Fig. 4A). No significant difference was observed among subtypes except that S-IV had a higher level of miRNA expression than S-II (p < 0.001). Differential analyses revealed that S-II exhibited fewer DEmiRs (n = 9) than other subtypes (S-I, n = 29; S-III, n = 17; S-IV, n = 30; Fig. 4B and Table S9). Next, GSEA was exploited to select DEmiRs that were overrepresented in the corresponding subtypes (Fig. 4C and Table S10). The results showed that four upregulated and two downregulated DEmiRs were significantly enriched in S-III and S-II, respectively. Hsa-miR-103a-3p enriched in S-III belongs to the miR-103 family, which is responsible for targeting genes involved in oxidative phosphorylation in PCa [18]. Of seven DEmiRs enriched in S-IV, upregulated hsa-miR-106-5p and downregulated hsa-miR-135a-5p had been found significantly associated with very high-risk aggressive prostate cancer at circulating levels [32]. S-IV was also overrepresented by downregulated hsa-miR-30d-5p, which inhibited PCa cell proliferation and invasion by targeting Ecto-5'-nucleotidase [33]. In contrast, overexpressed hsa-miR-30d-5p was significantly enriched in S-I and was correlated with low androgen receptor (AR) activity and inhibition of PCa cell growth [34,33]. Altogether, these results suggested that S-II was less affected by miRNA aberrations, whereas S-IV and S-I were heavily regulated. Furthermore, the identified subtype-driven miRNAs have been extensively linked to the distinct biological features of PCa subtypes (Fig. S4 and Table S11) with some experimentally validated to act as tumor-suppressive or oncogenic miRNAs.

To elucidate epigenetic and genomic alterations among PCa subtypes, we calculated methylation levels, somatic mutations, and SCNAS. Promoter methylation levels were comparable among S-II, S-III and S-IV except that S-I showed a lower level of methylation (p < 0.01; Fig. 4D). S-IV had the highest aneuploidy score and TMB, followed by S-III (p < 0.05; Fig. 4D). Both S-IV and S-III showed higher TNA loads than S-I and S-II (p < 0.05; Fig. 4D). Compared with other subtypes, S-IV had the highest burden in copy number gain and loss at both focal and arm levels (p < 0.001; Fig. 4E). Specifically, several tumor-suppressive genes (i.e. TP53, RB1, and PTEN) were more frequently deleted in S-IV (p < 0.001; Fig. 4F and Table S12). MYC and CCND1 amplifications were significantly observed in S-IV and least seen in S-II (p < 0.001). We also identified the top 20 frequently mutated genes in PCa (Fig. 4G and Table S13) and found that TP53 (31.30%, p < 0.001), FOXA1 (13.04%, p < 0.01), and LRPIB (8.70%, p < 0.05) were most frequently mutated in S-IV, followed by S-III.
3.4. Associations of PCa subtypes with clinical and biological features

We then compared the available clinicopathologic features among the four PCa subtypes (Tables S1–S3): elderly patients (>60 years, p = 0.004), surgical margins (p < 0.001) and lymph node invasion (p < 0.001) were significantly associated with S-IV in the TCGA-PRAD cohort. TMPRSS2-ERG fusion was significantly observed in S-II, S-III and S-IV when compared with S-I in both TCGA-PRAD (p = 0.035) and RNAseq cohorts (p = 0.031). Advanced ISUP (4 or 5, all p < 0.001) was significantly associated with S-I, S-III and S-IV in both TCGA-PRAD and microarray cohorts. Moreover, advanced pathological stage (T3 or T4, all p < 0.001) was remarkably correlated with S-IV in three cohorts. Notably, four subtypes were identified in each independent cohort, suggesting widespread intertumoral heterogeneity of PCa.

Using single-sample gene set enrichment analysis (ssGSEA), we next identified and validated distinct biological characteristics among the four subtypes in the TCGA-PRAD (Fig. 5A and Fig. S5A), microarray (Fig. 5B, Fig. S5B), and RNAseq cohorts (Fig. S6). S-I was significantly correlated with basal cell, stemness, neuroendocrine prostate cancer (NEPC), prostate gland development and mesenchymal-related (epithelial-mesenchymal transition [EMT], hypoxia, angiogenesis, Wnt, Notch, and TGFβ) signatures whereas S-II was enriched in luminal cells, ERG fusion, androgen response and metabolism-related (such as fatty acid metabolism, butanoate metabolism, steroid biosynthesis, and protein secretion) signatures. S-III exhibited elevated oxidative phosphorylation and moderate levels of proliferation as indicated by enriched DNA repair pathways (including base excision repair, nucleotide excision repair, homologous recombination, and mismatch repair). S-IV was characterized by moderate levels of NEPC, stemness, EMT and angiogenesis, as well as marked activities of DNA repair and cell cycle-related pathways such as E2F, G2M, and MYC targets.

3.5. Associations of PCa subtypes with immune infiltration

Next, we characterized the immune landscape of PCa subtypes in the TCGA-PRAD and microarray cohorts. The ESTIMATE algorithm revealed that S-I and S-IV had relatively higher immune and stromal scores and lower tumor purity than S-II and S-III (p < 0.05; Fig. 6A and 6C). Hence, we suspected that S-I and S-IV
were likely to be immune-related. To decode the immune microenvironments, ssGSEA was performed and revealed that S-IV was significantly enriched in immune-inflamed signatures (such as human leukocyte antigen signature, CD8 T effector cell, natural killer cell cytotoxicity, and T cell inflamed GEP; Fig. 6B and 6D, Fig. S7A and S7B). Although S-I displayed enrichment of immune-inflamed signatures similar to S-IV, it was found to have higher levels of myeloid-derived suppressor cells (MDSCs), regulatory T cells (Tregs), T-cell exhaustion, TGF\(\beta\) family members and receptors and cancer-associated fibroblasts (CAFs) than other subtypes. Thus, S-I showed an increased tendency toward an immune-excluded subtype. Overall, PCa subtypes in the RNAseq cohort demonstrated similar results (Fig. S8A-S8C).

3.6. Associations of PCa subtypes with therapeutic efficacy

To explore whether PCa subtypes responded differentially to conventional PCa therapies, we computed the AUC values of abiraterone, olaparib and taxane-based chemotherapy (Fig. 7A-7C) in PCa cohorts. As a result, S-II was significantly associated with increased sensitivity to abiraterone whereas S-I was more sensitive to olaparib than other subtypes. Although inconsistent results were demonstrated across PCa cohorts, S-III and S-IV patients were, in general, more susceptible to docetaxel and cabazitaxel.

We also used subMap to compare GEPs between PCa subtypes and 34 mCRPC patients treated with enzalutamide (160 mg/d). Data showed that S-I was significantly correlated with the enzalutamide-resistant group (all \(p < 0.01\); Fig. 7D). In contrast, the GEP of S-II was more similar to that of the enzalutamide-sensitive group (TCGA-PRAD, \(p = 0.172\); RNAseq cohort, \(p = 0.067\); Microarray cohort, \(p = 0.018\)). Since PCa subtypes correlated with different immune phenotypes, we asked whether they responded differently to immunotherapy. We first investigated the expression levels of PD-1, PD-L1 and CTLA-4, and found that they did not differ significantly between S-I and S-IV (except that S-I had slightly higher PD-L1 levels in the RNAseq cohort \([p < 0.05]\); Fig. S7C and S7D, Fig. S8D). SubMap analysis showed that S-I was significantly associated with anti-PD-L1 and anti-PD-1 nonresponders. In contrast, S-IV shared similar GEP with anti-PD-L1 responders (all \(p = 0.001\); Fig. 7D), indicating that S-IV may benefit from anti-PD-L1 therapy. Additionally, there were no significant associations between PCa subtypes and anti-CTLA-4 patients.

4. Discussion

The present study is the first to establish a robust PCa molecular classification based on miRNA-regulated GEPs, which offers new insights into the role of miRNAs in PCa heterogeneity and subty-
We identified and validated four PCa subtypes exhibiting distinct epigenetic, genomic, transcriptional, and clinicopathologic features. Of the four subtypes, S-I and S-IV exhibit immune-excluded and immune-inflamed phenotypes, respectively. Moreover, we uncovered subtype-driven miRNAs that could potentially serve as circulating diagnostic biomarkers and therapeutic targets for corresponding subtypes. Finally, possible treatment options for each subtype were suggested based on drug sensitivity analysis.

Fig. 4. The genomic and epigenetic landscape of the PCa subtypes in TCGA-PRAD. (A) Comparison of mean miRNA expression level among four PCa subtypes and normal samples. (B) A Venn diagram shows the number of unique and overlapping differentially expressed miRNAs (DEmiRs) among subtypes. (C) GSEA of subtype-specific DEmiRs identifies a subset of miRNAs that significantly contribute to the mRNA expression profiles of corresponding subtypes. (D) The levels of promoter methylation, somatic mutation, tumor neoantigen and aneuploidy were compared across subtypes. (E) The burden of copy number gain and loss at focal and arm levels in the four PCa subtypes. Oncoprints of top 20 somatic mutations (F) and somatic recurrent copy number alterations (G) among subtypes. The proportion of alteration of listed genes was shown on the right side. "ns" denotes no statistical significance, *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.
Whether prostate adenocarcinoma and NEPC originate from basal cells or luminal cells or both is constantly debated with conflicting results reported by previous studies [35–38]. Differences between S-I and S-II in our study support the hypothesis that PCa could be derived from either luminal or basal cell lineages, and that different cell origins give rise to distinct subtypes with divergent biological behaviors and drug responses. The identification of basal- and luminal-like subtypes has also been reported by previous studies [10,11]. Zhao et al. used PAM50 classifiers of breast cancer to assign PCa patients into luminal A, luminal B and basal-like subtypes and found that the luminal B subtype had worse prognosis than other subtypes [11]. Nevertheless, the direct application of this breast cancer classifier to PCa subtyping poses the risk of neglecting important genetic features of PCa.

Fig. 5. Associations of clinicopathologic and biological features with the PCa subtypes. Single-sample gene set enrichment analysis (ssGSEA) identified distinct biological characteristics among subtypes in the TCGA-PRAD (A) and Microarray (B) cohorts. Red indicates enrichment of signatures while blue suggests the opposite. NEPC, neuroendocrine prostate cancer. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Spratt et al. studied the heterogeneity of AR activity in primary treatment naïve PCa and identified a low AR-active and androgen deprivation therapy (ADT)-insensitive subclass that is in accordance with S-I characterized by overexpression of basal and NEPC signatures and decreased DNA repair [39]. These biological features also support the results of drug sensitivity in S-I. NEPC and low AR activity were extensively demonstrated to confer castration resistance [40]. Deficient DNA repair mechanisms, especially homologous recombination repair were associated with the antitumor activity of PARP inhibitors in patients with mCRPC [41]. In contrast to S-I, S-II was highly enriched in the androgen response and therefore, showed favorable response to ADTs. Although neither luminal nor basal signatures were overrepresented in S-III and S-IV, TMPRSS2-ERG fusion at both genomic and transcriptional levels was more enriched in S-II, S-III, and S-IV than in S-I, favoring the assumption that S-III and S-IV shared luminal cell origin with S-II. However, S-IV was also enriched in NEPC signatures, suggesting a possible role of lineage plasticity during S-IV evolution. This was supported by Dong et al. who used single-cell RNA sequencing and found that luminal-like adenocarcinoma cells could transdifferentiate into NE cells with loss of luminal markers [42].

Our data showed that basal-like S-I exhibited stem- and mesenchymal-like properties, which was consistent with a previous study uncovering intrinsic stem cell and EMT features in prostatic basal cells [43]. However, our study does not support the linkage between the basal cell signature and aggressive PCa [43]. Instead of S-I, S-IV was characterized as a more aggressive subtype of PCa with worse prognosis, significant enrichment in cell cycle-related pathways, and moderate levels of EMT and stemness. In support of this observation, lineage analysis by Wang et al.
revealed that PCa of luminal origin was more aggressive than basal origin prostate cancer and exhibited more active cell cycle-related pathways [38]. The most likely explanation for this is that although S-I is more migratory and mesenchymal-like, it is strikingly less proliferative than S-IV. Reasonably, proliferative capacity is essential to generate PCa macrometastasis once tumor cells reach distant sites with the help of EMT [44]. Increased CNV burden and aneuploidy also contributed to the worse prognosis of S-IV, as these factors were reported to promote aggressive PCa development and earlier biochemical relapse [45,46]. Specifically, we identified higher frequencies of \( \text{TP53} \) and \( \text{RB1} \) functional loss in S-IV. The biallelic loss of \( \text{TP53} \) and \( \text{RB1} \) has been associated with attenuated AR activity, increased cell proliferation and DNA repair activity, as well as exhibition of stemness and NE differentiation in
LNCaP cells [47]. Similar observations in S-IV support these associations, which highlight the important roles of TP53 and RB1 loss in the development of aggressive PCa.

Immune checkpoint blockade (ICB) treatments have been proven to be effective against several cancers including melanoma and bladder cancer. However, the use of ICB alone has thus far shown modest efficacy in advanced PCa as most PCa has insufficient immunogenicity and few tumor-infiltrating T cells [48–50]. It is, therefore, necessary to select a subset of PCa that may benefit from ICB. Our study showed that S-I was unlikely to respond to anti-PD-1 and anti-PD-L1 therapies because it was mostly immune-excluded, with enrichment in CAFs, MDSCs, and T cell exhaustion. Furthermore, increased TGFβ activity in S-I was reported to promote T cell exclusion and concomitant resistance to ICB [51]. Consequently, ICB resistance in S-I patients may be overcome by combining anti-TGFβ treatment. S-IV was mostly immune-inflamed and therefore, may benefit from anti-PD-L1 therapy. The higher TMB and TAN loads in S-IV could also increase the efficacy by driving infiltrating T cell responses, as evidenced by previous studies [27,52]. Taken together, these findings suggest that the current PCa subtypes offer the potential to provide ICB guidance.

Our study has several limitations that need to be optimized in the future. First, since we only used miRNA-regulated GEPs to classify PCa patients, our classification system is biased and may obscure important features determined by other omics data. Second, due to the lack of miRNA expression data, the classification pipeline used in the TCGA-PRAD dataset was not repeated in the validation datasets. Last, prospective cohorts and functional experiments are needed to validate the proposed classification and confirm our findings.

5. Conclusions

In summary, findings from the present study expand our understanding of PCa heterogeneity from the perspective of miRNAs regulation. While the proposed classification has the potential to predict clinical outcomes and inform personalized treatments for PCa patients, further experimental studies are required to verify this conclusion.

CRediT authorship contribution statement

Bing-Biao Lin: Conceptualization, Methodology, Software, Investigation, Visualization, Writing - original draft. Han-Qi Lei: Methodology, Investigation, Writing - original draft. Hai-Yun Xiong: Methodology, Investigation, Writing - original draft. Xing-Fu Fu: Methodology, Investigation, Visualization, Writing - review & editing. Fu Shi: Investigation, Writing - review & editing. Xiang-Wei Yang: Investigation, Writing - review & editing. Ya-Fei Yang: . Guo-Long Liao: Writing - review & editing. Yu-Peng Feng: Writing - review & editing. Dong-Gen Jiang: Conceptualization, Writing - review & editing. Funding acquisition, Supervision. Jun Pang: Conceptualization, Writing - review & editing. Funding acquisition, Supervision.

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Data availability

The data analyzed in this study are publicly available in TCGA, GEO, cBioPortal, CCLE, and PRISM Repurposing datasets.

Appendix A. Supplementary data

Figure S1. Identification and removal of batch effects during datasets merging. The principal component analysis demonstrated significant batch effects in the microarray-based (A) and RNA-sequencing (B) cohorts, respectively. Batch effects were removed in the microarray-based (C) and RNA-sequencing (D) cohorts.

Figure S2. Results of non-negative matrix factorization based on miRNA-correlated mRNAs. Consensus maps (A) and cophenetic coefficients (B) of PCa classifications with clustering numbers ranging from 2 to 6.

Figure S3. KEGG and GO analyses of subtype-specific upregulated miRNAs. Distinct pathway enrichments were shown for S-I (A), S-II (B), S-III (C) and S-IV (D).

Figure S4. KEGG and GO analyses of subtype-driven miRNAs. Active pathways determined by correlated miRNAs were only observed in S-I (A), S-II (B) and S-IV (C).

Figure S5. Biological features associated with the PCa subtypes. Boxplots showing ssGSEA scores of biological pathways across subtypes in the TCGA-PRAD (A) and Microarray (B) cohorts.

Figure S6. Clinicopathologic and biological features associated with the PCa subtypes in the RNAsq cohort. The heatmap (A) and boxplots (B) showing ssGSEA scores of biological pathways across subtypes. Red indicates enrichment of signatures while blue suggests the opposite. “ns” denotes no statistical significance, *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001. NEPC, neuroendocrine prostate cancer.

Figure S7. Immune characterization of the PCa subtypes. Boxplots showing ssGSEA scores of immune-related pathways for the PCa subtypes in the TCGA-PRAD (A) and Microarray (B) cohorts. The levels of immune checkpoint expression in the TCGA-PRAD (C) and Microarray (D) cohorts. “ns” denotes no statistical significance, *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001. HLA, human leukocyte antigen; TCR, T cell receptor; IFN, interferon; GEP, gene expression profile; MDSC, myeloid-derived suppressor cells; Treg cells, regulatory T cells; TGFb, transforming growth factor β; Pan F TBRs, pan tissue fibroblast TGF-β response signature. CTLA-4, cytotoxic T-lymphocyte associated protein-4; PD-1, programmed cell death protein-1; PD-L1, programmed death-ligand 1.

Figure S8. Immune characterization of the PCa subtypes in the RNAsq cohort. The immune score, stromal score and tumor purity were compared across subtypes (A). The heatmap (B) and boxplots (C) show ssGSEA scores of immune-related pathways in the four PCa subtypes. Red indicates enrichment of signatures while blue suggests the opposite. Boxplots (D) showing the levels of immune checkpoint expression. “ns” denotes no statistical significance, *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001.
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