**CD8+ effector memory T cells Marker HLA-B Is a Risk Factor for Atherosclerosis and Prostate Cancer: An Insight Integrating Single-Cell Expression Quantitative Trait Locus and Mendelian Randomization Analyses**

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**Abstract**

**Background:** atherosclerosis (AS) and Prostate cancer (PCa), though traditionally classified as distinct diseases, share inflammatory and immune dysregulation features, supporting the emerging concept of pan-vascular diseases. This study investigates common immune mechanisms and potential targets across these conditions.

**Methods:** We integrated single-cell RNA sequencing and Mendelian randomization (MR) to profile T cell subsets in PCa, AS, and healthy controls. eQTL mapping and bidirectional MR were conducted to identify causal genes, followed by cell-cell communication and trajectory analyses.

**Results:** CD8+ effector memory T cells (CD8\_EM) were enriched in both PCa and AS, displaying shared immune activation patterns. HLA-B, a key CD8\_EM marker, was causally linked to PCa and highly expressed in AS. HLA-B+ CD8\_EM cells showed stronger signaling and communication activities in diseased tissues.

**Conclusion:** HLA-B may serve as a shared immunological driver in PCa and AS, highlighting its role in the broader context of pan-vascular pathology. These findings offer new insights into shared disease mechanisms and immunotherapeutic strategies for vascular-related malignancies.

**Keywords:** CD8⁺ Effector Memory T Cells, HLA-B, Atherosclerosis, Prostate Cancer, Single-Cell RNA Sequencing

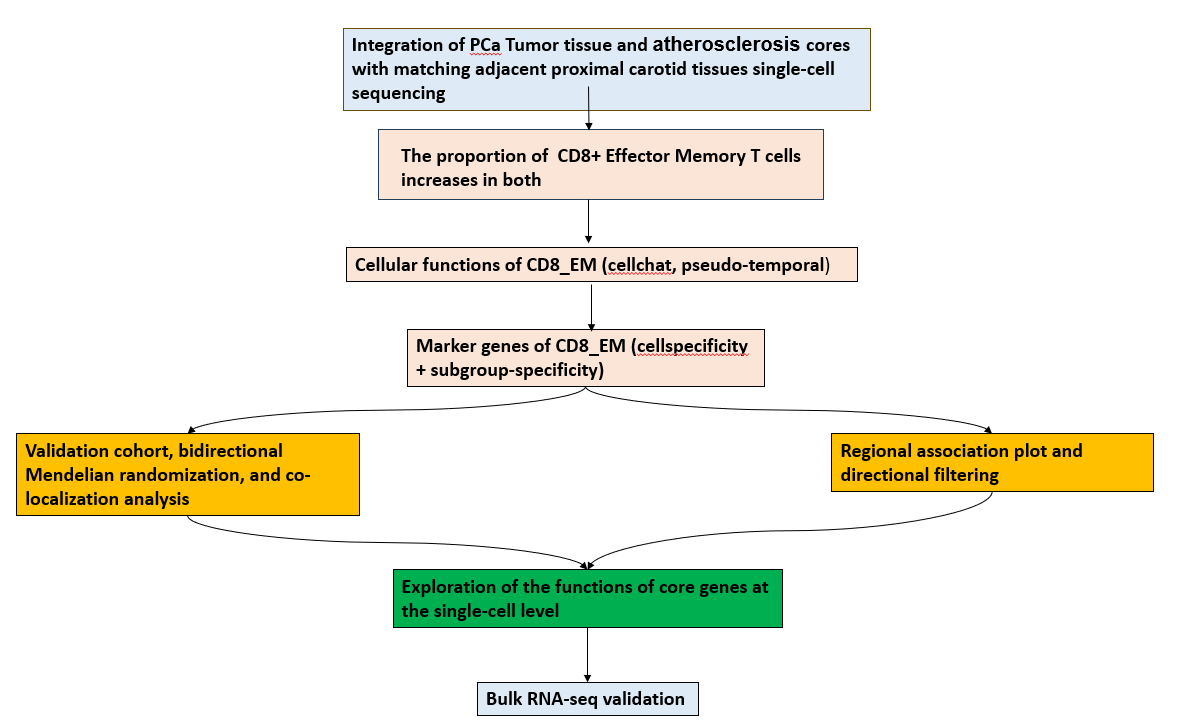
**Introduction**

Prostate cancer (PCa) is the most prevalent urogenital tumor in men globally, affecting over 1.4 million individuals and causing more than 375,000 deaths annually[1]. As one of the most common male malignancies, it primarily targets the prostate gland. The cancer often progresses slowly and initially relies on androgens, but it can evolve into castration-resistant prostate cancer (CRPC), resistant to standard androgen deprivation therapy [2, 3]. Early detection includes PSA testing, rectal exams, and imaging [4]. Treatment options encompass surgery, radiotherapy, hormone therapy, chemotherapy, and increasingly, immunotherapy and targeted therapies for advanced cases [5].

Atherosclerosis (AS) is a systemic vascular disorder characterized by arterial wall inflammation and plaque formation, representing a major component of panvascular diseases[6, 7]. It is a leading cause of heart disease, stroke, and peripheral artery disease, underscoring its role in global morbidity and mortality. The condition begins with endothelial cell activation, which triggers vessel narrowing and inflammation, leading to plaque formation. Notably, AS shares pathophysiological similarities with cancer, including chronic inflammation, metabolic reprogramming, and immune dysregulation[8, 9]. For instance, recent studies highlight the oncogenic-like behavior of vascular smooth muscle cells (SMCs), such as their ability to undergo proliferation, resist apoptosis, and remodel the extracellular matrix—phenomena also observed in tumor cells[10, 11]. These parallels have led to the emergence of "pan-cardio-oncology", a paradigm that seeks to unravel shared molecular mechanisms between vascular diseases and cancer[12, 13].

Inflammation may bridge lipid abnormalities and AS progression [8, 9]. Studies suggest that high-sensitivity C-reactive protein (hsCRP) independently predicts cardiovascular events [10, 11]. In rodent models, mutations in genes linked to clonal hematopoiesis, such as Dnmt3a, Tet2, or Jak2 V617F, accelerate AS and activate pro-inflammatory pathways [12, 13]. These insights advance understanding and treatment of AS, though further research is needed to fully uncover its mechanisms and risk mitigation strategies. Importantly, clonal hematopoiesis-associated mutations have also been implicated in the development of certain cancers, including PCa, underscoring the genetic and immunological overlap between AS and malignancies [13].

This study aims to identify potential therapeutic targets shared between AS and (PCa through Mendelian randomization (MR) analysis, integrating eQTL data from blood samples with two independent PCa Genome-Wide Association Study (GWAS) datasets. We specifically focus on the association between the HLA-B gene and PCa. The findings from this study could provide new insights into the shared molecular mechanisms underlying these diseases and highlight potential therapeutic targets for both AS and PCa. The workflow for this study is outlined in Figure 1.



**Methods**

**Data collection**

We downloaded STAR count data and clinical information for 52 normal samples and 499 PCa samples from the UCSC Xena website (<https://xena.ucsc.edu/>). TPM format data was extracted and normalized using log2 (TPM + 1). We retained samples with RNAseq data and clinical information for further analysis.

Additionally, the datasets (GSE100927, GSE159677, GSE216860, and GSE141445) were retrieved from the GEO database (<https://www.ncbi.nlm.nih.gov/>). We conducted transcriptomic analysis of human peripheral arteries in the carotid, femoral, and popliteal regions from atherosclerotic and control tissues. Furthermore tumor cells from 13 PCa patients ([GSM4203181](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4203181)),normal tissues from 6 healthy individuals (GSM6696842,GSM6696843,GSM6696844,GSM6696845,[GSM6696846](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM6696846),[GSM6696847](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM6696847)), and atherosclerotic cores with matching adjacent proximal carotid tissues from 3 AS patients ([GSM4837523](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4837523), [GSM4837524](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4837524), [GSM4837525](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4837525), [GSM4837526](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4837526), [GSM4837527](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4837527),[GSM4837528](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM4837528)) were selected for 10X single-cell sequencing. The SNPs used as genetic instruments were sourced from a comprehensive European GWAS (<https://gwas.mrcieu.ac.uk/>)[14-16].

**Single-cell data processing and analysis**

Single-cell RNA-seq data from normal, atherosclerotic, and tumor tissues were processed using the Seurat pipeline. Individual Seurat objects were created for each sample and merged into a unified object for integrated analysis. Cells with high mitochondrial or red blood cell gene expression were excluded during quality control. Following normalization, principal component analysis (PCA) was conducted on the top 2,000 highly variable genes. Batch effects were corrected using the Harmony algorithm, and clustering was performed using UMAP for dimensionality reduction. Cell-type annotation was based on canonical markers from the CellMarker database, visualized via FeaturePlot and DimPlot. After manual verification, differential expression analysis was performed using a fold change threshold of 1.5 and an adjusted p-value < 0.05, with results visualized through volcano plots and heatmaps. For developmental trajectory reconstruction, Monocle was employed, while CellChat was used to model intercellular communication.

**CD8\_EM key marker gene eQTL and PCa's mendelian randomization analysis**

CD8\_EM marker genes were identified through comparison with other T cell subsets. Gene symbols were converted to ENSEMBL IDs, and SNPs associated with these markers were extracted from the "ieu-b-4809" GWAS dataset. SNPs passing an eQTL p-value threshold of 5 × 10⁻⁸ were retained. R² and F-statistics were calculated to assess instrument strength. TwoSampleMR was used to estimate causal associations between candidate genes and PCa.

**Validation set MR analysis, bidirectional Mendelian MR and colocalization analysis**

Validation analysis was conducted by harmonizing exposure and outcome datasets, followed by causal inference using the mr\_modified function, which also calculated the phenotypic variance explained (PVE). Forest plots were generated for visualization. For bidirectional MR, SNPs from the reverse exposure dataset (“ebi-a-GCST90018905”) were merged with outcome data to generate a harmonized PCa gene set. eQTL data were processed using vcfR, and coloc.abf was applied to assess colocalization using Bayes factor estimation. The results were interpreted according to standard colocalization hypotheses and evidence strength.

**Regional association plot**

Construct a regional association plot. By reading the genotype data from eqtl-a-ENSG00000234745.vcf (in VCF format) and the related association data, extract the eQTL information associated with the target gene. Based on this, select the eQTLs located within the specified region and organize them into a format suitable for plotting a regional association plot. The plotting process uses the locuscomparer package to visualize the association information from both eQTL and GWAS, providing an intuitive graphical presentation for subsequent analysis.

**Single-cell Functional Analysis of Exposure-Associated Genes**

To investigate the functional role of target genes at the single-cell level, we visualized their expression patterns using DotPlot and FeaturePlot in Seurat. Developmental trajectories were reconstructed with the Slingshot package, with dimensionality reduction performed via UMAP. Key trajectory-driving genes were identified using the find\_switch\_logistic\_fastglm function, and gene expression dynamics were illustrated along pseudotime using plot\_timeline\_ggplot.

Intercellular communication analysis was conducted with the CellChat tool. This included identifying ligand–receptor pairs, mapping interactions to protein–protein interaction networks, and calculating communication probabilities. The network structure was visualized using netVisual\_circle and netVisual\_bubble. Additionally, differential gene expression analysis was performed by integrating bulk transcriptomic datasets to validate single-cell findings and assess consistency across platforms.

**Trajectory inference**

We used the monocle3 package in R [17, 18] to infer T cell differentiation trajectories. The process began with data preprocessing, including quality control, normalization, and dimensionality reduction. Parameters were then configured following the guidelines in the official documentation, leading to cell state inference and transcriptome trajectory analysis.

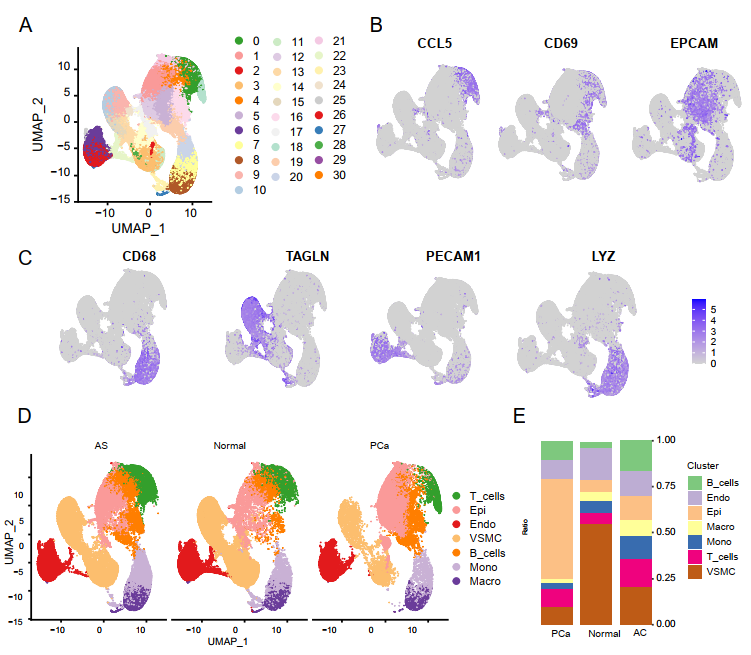
**Data analysis**

All data analyses were performed based on R 4.2.1, with *p* < 0.05 considered statistically significant.

**Results**

**Single-cell transcriptomic analysis of PCa, AS and normal groups**

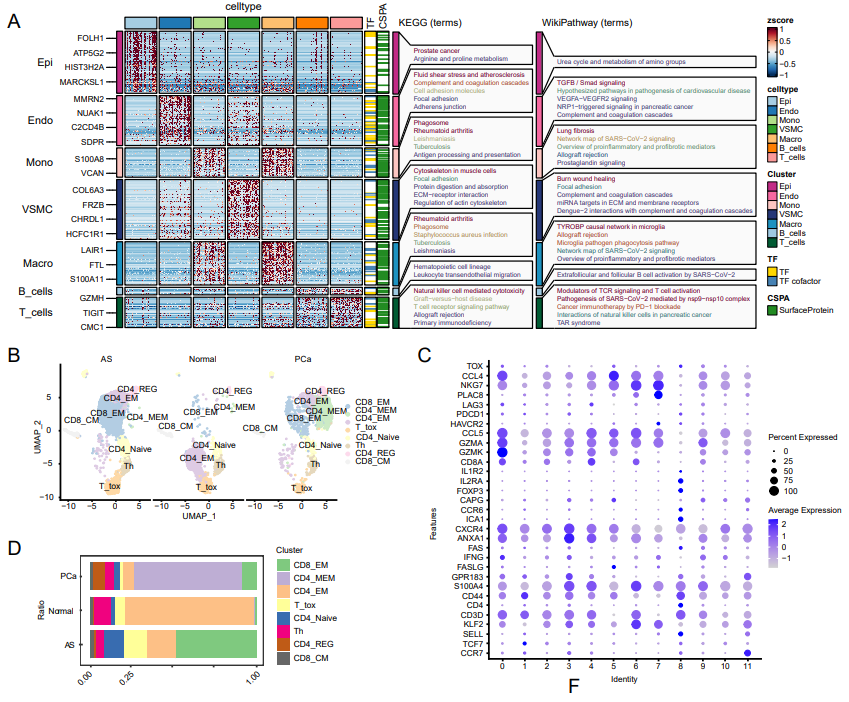
In this study, we selected 13 PCa samples, 6 healthy controls, and 3 groups of AS samples from the datasets GSE159677, GSE216860, and GSE141445 for 10X single-cell RNA sequencing analysis. Low-quality data was initially filtered out (Fig.S1 A-B), leaving 64,568 cells for further analysis. To reduce batch effects across samples, we used the Harmony method for integration and standardization, followed by normalization, PCA for dimensionality reduction, and clustering (Fig.S1 C). Visualization of each cluster was achieved using UMAP based on the top 15 principal components (Fig.S1 D). Specific marker genes were then used to identify and annotate different cell subpopulations in the single-cell RNA sequencing data. We annotated the clusters using the RenameIdents function, aligning each with its corresponding cell type. Cell distribution was visualized with UMAP and DimPlot (Fig.2 A). FeaturePlot in Seurat was used to display the spatial distribution of marker genes for various cell types, including T cells, epithelial cells, endothelial cells, vascular smooth muscle cells, B cells, macrophages, and monocytes. (Fig.2 B, C). Manual annotation renamed 30 clusters, visualized by UMAP and DimPlot, segmented by tissue type (Fig.2 D). Figure 2E shows significant differences in T and epithelial cell proportions across samples, highlighting the importance of understanding T cell functions in cancer and AS for effective treatments.



**KEGG and WikiPathway enrichment analysis and single-cell transcriptomics of T cells**

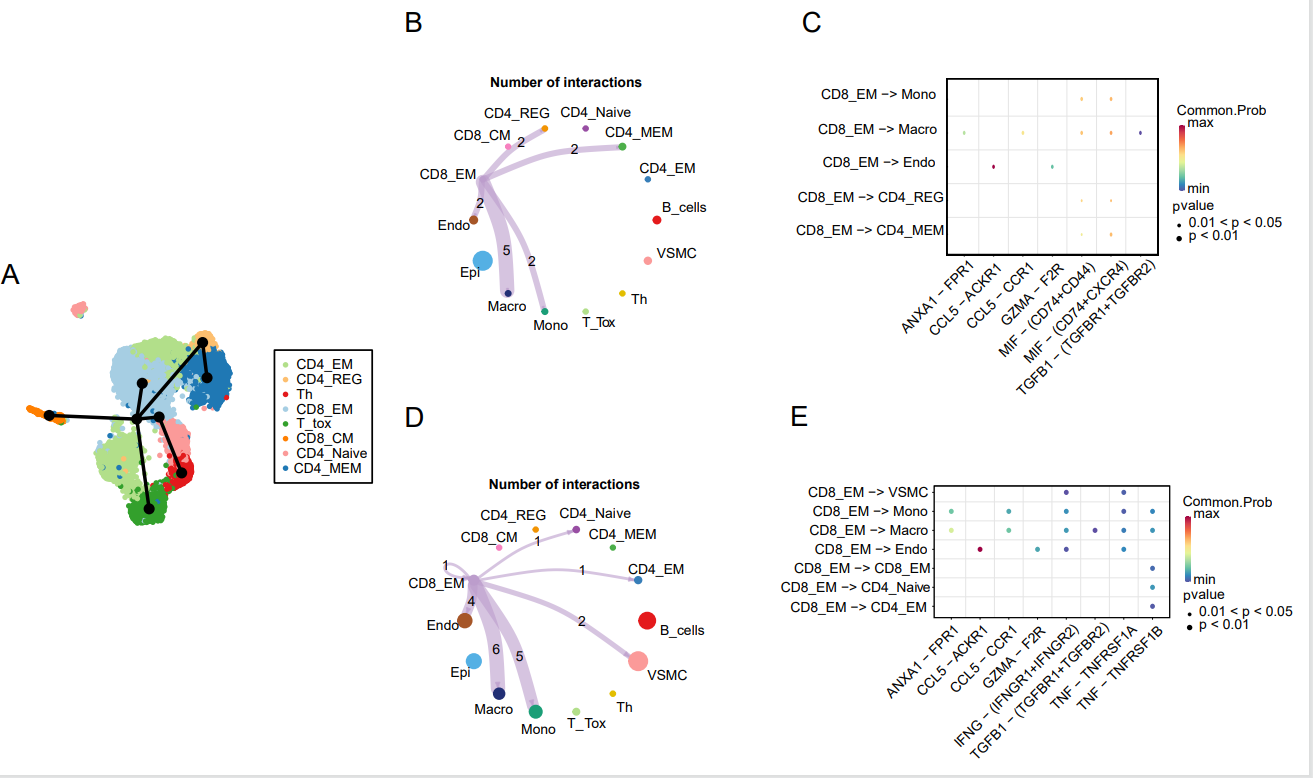
We performed differential gene expression analysis across seven cell types—T cells, epithelial cells, endothelial cells, vascular smooth muscle cells, B cells, monocytes, and macrophages—defining genes with fold change ≥1.5 or ≤−1.5 as differentially expressed. To explore functional roles of differentially expressed T cell genes, we conducted enrichment analyses using TF, CSPA, GO\_BP, KEGG, and WikiPathway databases (Fig. 2A). KEGG analysis revealed enrichment in NK cell-mediated cytotoxicity, while WikiPathway indicated involvement in NK cell interactions in PCa, TCR signaling, and T cell activation. These findings highlight the distinct biological roles of T cells in immune responses and disease.

We then isolated T cell subpopulations from the single-cell RNA-seq data, followed by standard preprocessing, dimensionality reduction, clustering, and manual annotation. Eight T cell clusters were identified and visualized by tissue type (Fig. 3B). A dot plot illustrated marker gene expression across subpopulations (Fig. 3C), and a cell proportion plot showed a significant increase in CD8\_EM cells in cancer and AS groups versus normal (Fig. 3D). These results offer valuable insights for downstream analysis.

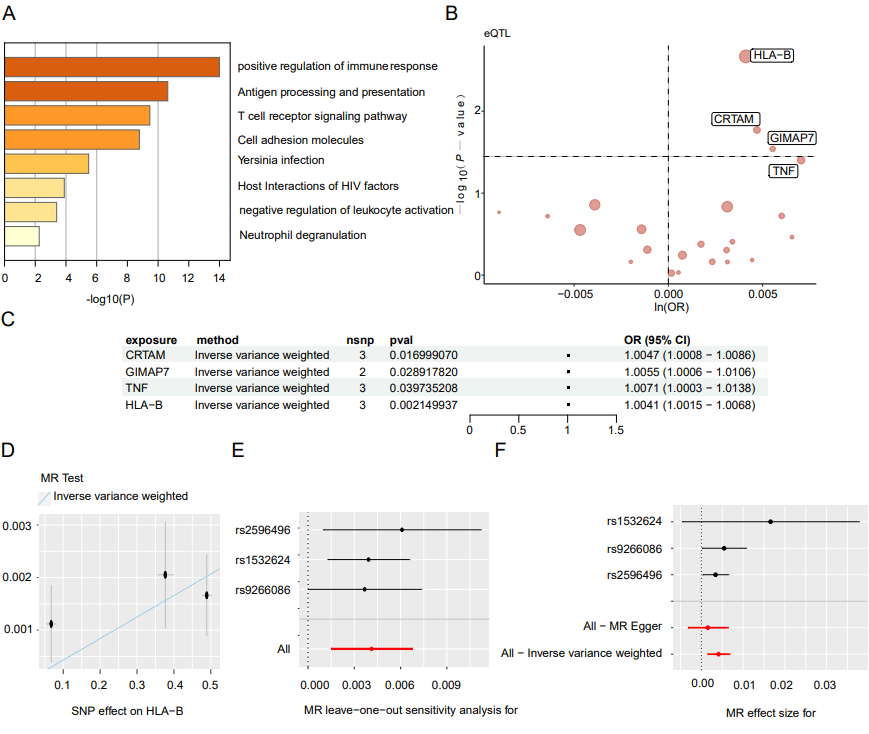


**Trajectory analysis of T cell types and cell communication**

Using single-cell RNA sequencing and the Slingshot R package, we reconstructed T cell differentiation trajectories. UMAP visualization effectively captured the continuous nature of T cell development, identifying CD4\_Naïve as the root and CD8\_EM as a differentiated branch, suggesting that CD8\_EM cells arise from CD8⁺ T cells after antigen stimulation (Fig. 4A). To further explore the functional role of CD8\_EM cells in PCa and AS, we performed cell-cell communication analysis. In both diseases, CD8\_EM cells exhibited frequent interactions with endothelial cells, primarily via the CCL5–ACKR1 signaling pathway (Fig. 4B–E). These findings underscore the significance of CD8\_EM cells as central mediators of immune communication in the disease microenvironment.

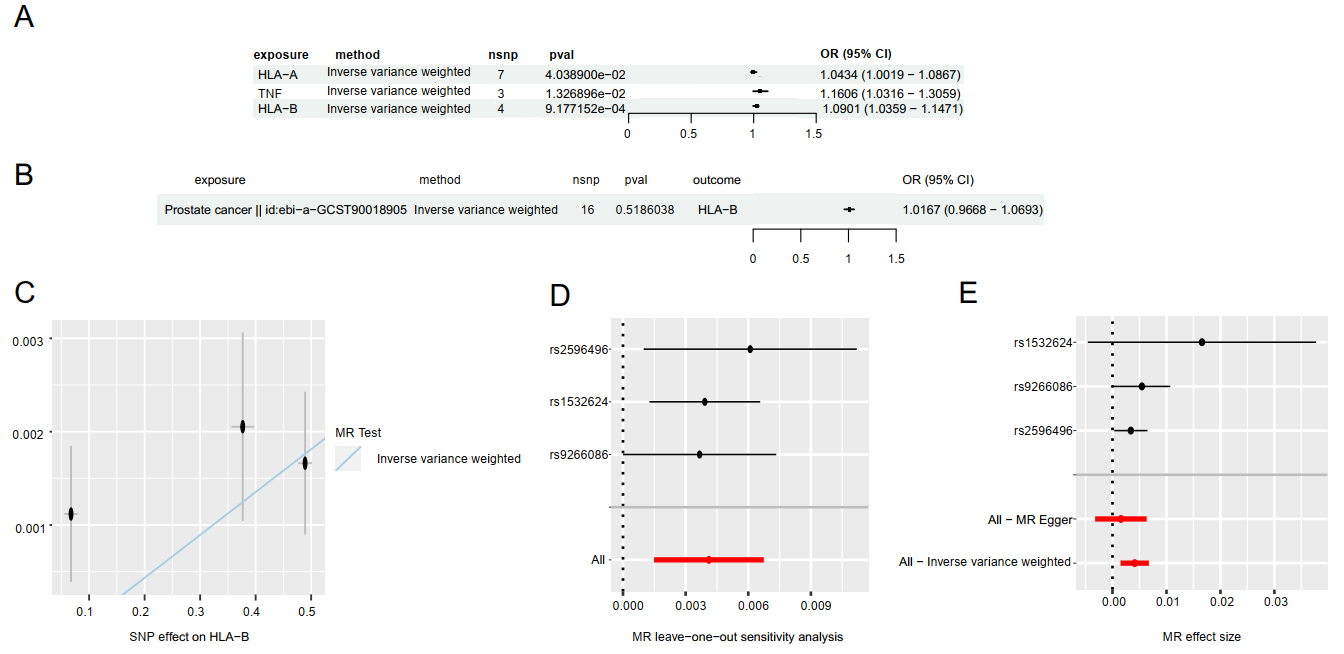


**MR analysis using key marker genes identifies one novel causal gene for PCa**

We identified 30 key genes specifically expressed in CD8\_EM cells using Seurat (Table S1). These genes distinguish CD8\_EM T cells from other subtypes and serve as crucial markers for downstream analyses. Enrichment analysis via Metscape linked them to pathways such as "positive regulation of programmed cell death," "antigen processing and presentation," and "T cell receptor signaling" (Fig. 5A). To assess potential associations with prostate cancer (PCa), we performed two-sample Mendelian randomization (MR) using scRNA-seq data. Gene symbols were converted to ENSEMBL IDs via org.Hs.eg.db. SNPs related to target genes were extracted as exposures, and PCa outcome data were obtained from the EBI database. Integration of these datasets revealed several genes significantly associated with PCa risk: CRTAM (OR=1.0047, p=0.0169), GIMAP7 (OR=1.0055, p=0.0289), TNF (OR=1.0071, p=0.039), and HLA-B (OR=1.0041, p=0.0021). A volcano plot (-log10 p vs ln(OR), Fig. 5B) and forest plots (Fig. 5C–F) visualize these associations. No significant heterogeneity or horizontal pleiotropy was observed for CRTAM, TNF, and HLA-B (Table S2–S3), supporting the robustness of our findings. These results point to potential genetic markers for PCa risk and offer directions for future biological validation.

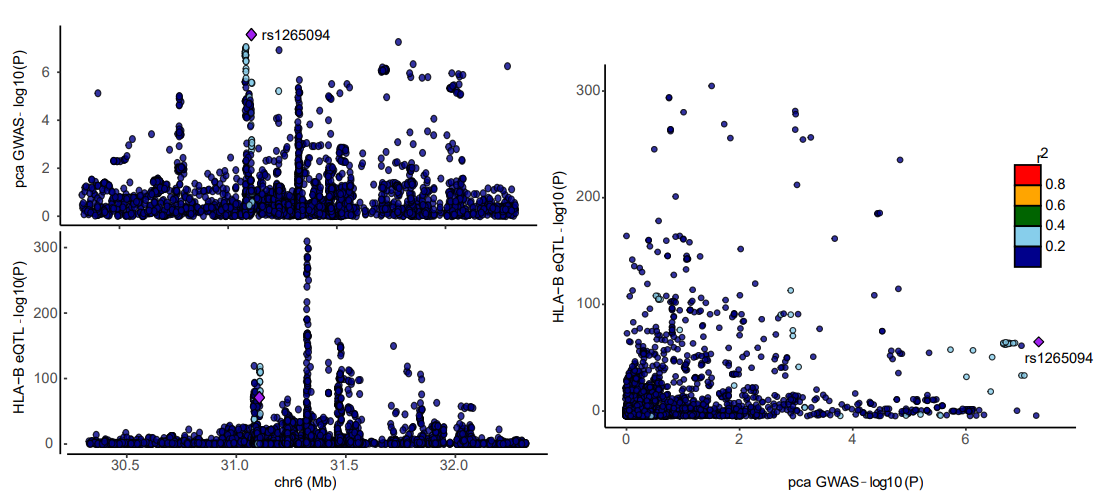
**MR analysis of gene HLA-B with PCa risk**

To validate our findings, we conducted a MR validation analysis. By intersecting results with the IEU dataset, we identified the TNF and HLA-B genes, with HLA-B showing the smallest p-value, indicating a stronger association with PCa. Further investigation revealed that HLA-B mutations are significantly associated with PCa risk (OR = 1.0901, 95% CI: 1.0359−1.1471) (Fig. 6A). Reverse MR analysis showed no significant association when treating PCa as the exposure and HLA-B as the outcome (Fig. 6B). We also tested horizontal pleiotropy, heterogeneity, validity, and robustness (Fig. 6C-E, Table S4). To sum up, we investigated the association between the HLA-B gene and PCa using MR.



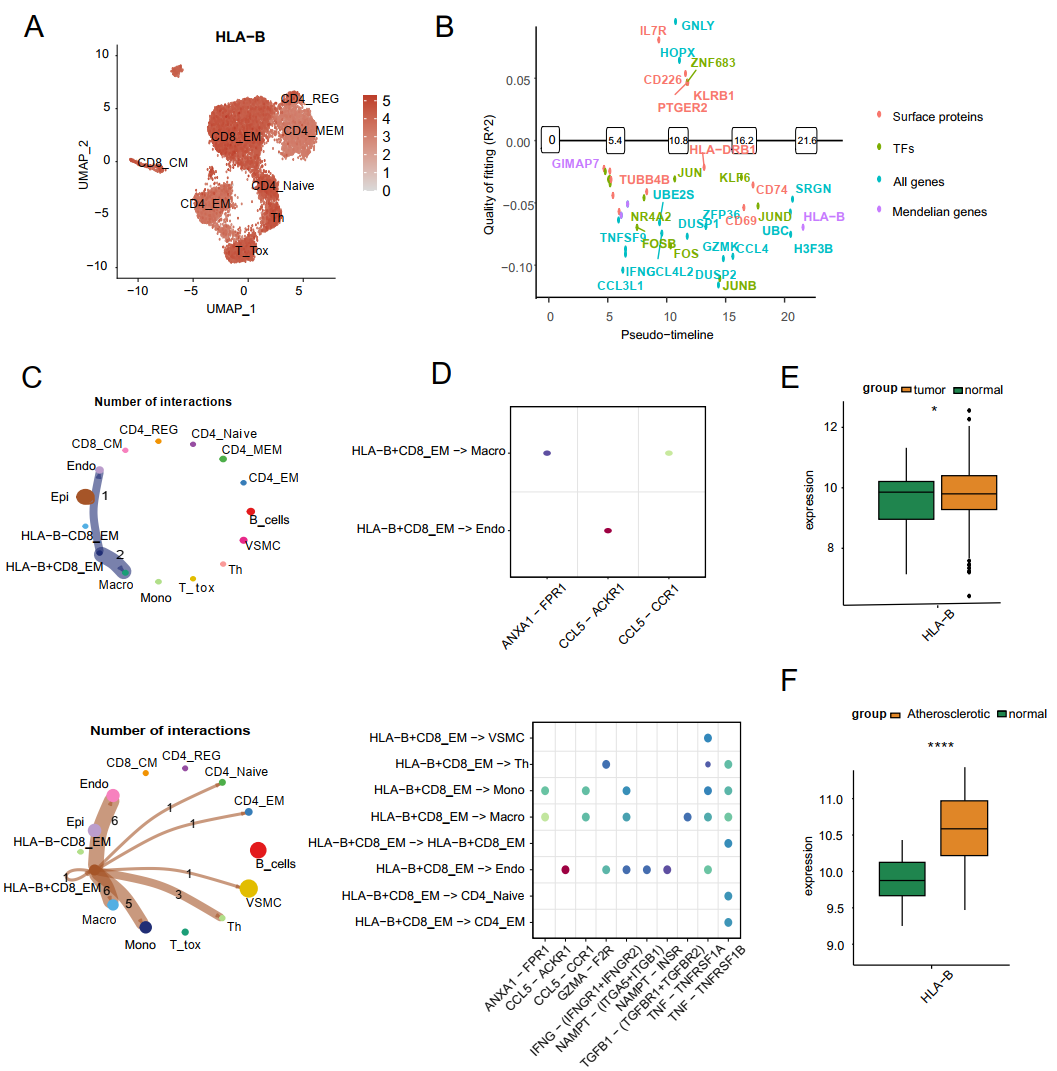
**Investigating the association between the HLA-B gene and PCa using Mendelian Randomization**

In this study, we presented the eQTL regional association plot for the HLA-B gene and the PCa GWAS results. By comparing the association strengths, we identified several SNPs that showed strong associations with both phenotypes. Notably, certain SNPs related to HLA-B eQTLs (eg: rs1265094) also exhibited significant associations in the PCa GWAS. This provides preliminary evidence for a potential link between the HLA-B gene and PCa development (Figure 6).



**Single-cell transcriptome analysis reveals the critical role of HLA-B in T cell function within PCa and AS**

We analyzed the expression dynamics of HLA-B across cell populations and pseudotime. FeaturePlot revealed that HLA-B is highly expressed in specific cell clusters (Fig. 8A), with increased expression during early-to-mid stages of T cell development and a decline in later stages (Fig. 8B). In both PCa and AS groups, CellChat analysis showed that HLA-B⁺ CD8\_EM cells exhibited stronger intercellular communication and more active ligand-receptor interactions than their HLA-B⁻ counterparts (Fig. 8C). Notably, the CCL5–ACKR1 signaling pathway was prominently enriched in HLA-B⁺ CD8\_EM-mediated interactions (Fig. 8D). Differential expression analysis confirmed that HLA-B⁺ CD8\_EM cells are linked to PCa-associated genes. Furthermore, HLA-B expression levels were significantly higher in PCa and AS samples compared to healthy controls (Fig. 8E–F), reinforcing its potential role as a functional immune modulator in disease states.



**Discussion**

PCa, one of the most common male malignancies, initially relies on androgens but can progress to CRPC. It is often described as an "immune-cold" tumor, characterized by fewer gene mutations in tumor cells, reduced antigen presentation, and lower CD8+ T cell activation, alongside higher levels of immunosuppressive molecules like cytokines and chemokines [19, 20]. CD8+ T cells recognize antigen peptides presented by MHC class I molecules and eliminate target cells through perforin and granzyme release, making their function crucial in anti-tumor immunity and increasingly vital in cancer immunotherapy [21, 22]. CD8+ T cells include two circulating memory subpopulations: CD8\_EM and CD8\_CM. CD8\_EM, or effector memory CD8+ T cells, are a subset within the CD8+ T cell lineage that can rapidly respond to tumor recurrence, playing a crucial role in the adaptive immune system[23, 24]. They are primarily found in peripheral blood, peripheral tissues, and secondary lymphoid organs[25]. Although CD8\_CM cells have stronger antitumor effects, CD8\_EM cells are more widely distributed throughout the body, particularly in non-lymphoid tissues, enabling a faster response to infections or tumors [26].

The CD8+ T cells induced by this vaccine are crucial for protective immune responses and play a key role in PCa treatment, particularly through their involvement in immune surveillance and antitumor activity. However, in PCa patients, CD8\_EM cells may become dysfunctional and inactive due to prolonged antigen exposure in chronic infections or the tumor microenvironment (TME) [27-30]. Within the TME, immune suppression occurs through (1) differential mutations that regulate immune responses [31-33] and (2) tumor-infiltrating T cells that induce checkpoint molecules and recruit immunosuppressive cells [34-36]. In PCa tissue, the number of CD8\_EM cells is lower compared to peripheral blood, with reduced cytokine secretion capacity. Many PCa patients exhibit immune ignorance, partly due to the tumor microenvironment (TME) suppressing CD8\_EM cell activity through mechanisms like the CTLA4, PD-1, PD-L1/2, and TGF-β pathways. CTLA-4 and PD-1 are immune checkpoint molecules that inhibit T cell activation[37], while PD-L1/2 binds to PD-1, further suppressing T cell function[38]. TGF-β enhances this suppression by promoting an immunosuppressive environment[39]. Enhancing CD8\_EM cell infiltration and activity through immunotherapy is a key approach to boosting antitumor immunity in PCa patients. Studies have shown that using anti-CTLA-4, anti-PD-1, or anti-PD-L1 antibodies, as well as TGF-β inhibitors, can partially restore the activity and function of CD8\_EM cells within tumors [40-43]. The subpopulation and functional state of CD8\_EM cells can serve as important biomarkers for predicting and monitoring the effectiveness of PCa immunotherapy. In summary, the subpopulation and functional state of CD8\_EM cells are crucial for immune responses in tumor immunity. Further exploration of their biology and function, coupled with immunotherapeutic strategies, could advance antitumor treatments.

This study systematically examined the composition and function of immune cells in AS and PCa patients as well as healthy individuals using scRNA-seq analysis. Our preliminary data confirmed the presence of T cells, B cells, epithelial cells, and others in the tumor tissue of PCa patients and the atherosclerotic core and matching carotid tissue in AS patients, consistent with existing literature[44-46]. Notably, both AS and PCa patient groups showed an increased proportion of effector memory CD8+ T cells (CD8\_EM) compared to healthy controls, potentially linked to disease characteristics.

Additionally, single-cell trajectory analysis has revealed the dynamic differentiation process of T cells, offering new insights into T cell development and activation. The analysis of cell communication networks also showed enhanced interactions between CD8\_EM T cells and endothelial cells, suggesting their role in regulating other immune cells during disease progression. This study not only identifies disease-related immune cell changes but also proposes potential molecular mechanisms through multi-omics analysis. For example, we found that increased HLA-B expression might contribute to the reduced function of CD8+ T cells. HLA-B is a T cell receptor gene involved in antigen recognition and is part of the major histocompatibility complex I (MHC-I) [47]. While high HLA-B expression can enhance the immune system's ability to recognize and kill tumor cells, its high polymorphism means different HLA-B alleles are linked to PCa susceptibility [48-51]. In the tumor microenvironment, excessive HLA-B expression may cause immune cells, including T and B lymphocytes, and NK cells, to lose their function [52]. Therefore, regulating HLA-B expression levels could be a new strategy to enhance T cell antitumor activity.

It is important to acknowledge certain limitations in this study. First, due to the limited sample size, the results require validation in larger cohorts. Additionally, the predicted functional roles of key genes need further experimental validation, and molecular mechanisms should be explored using animal models. Furthermore, the inherent batch effects of scRNA-seq technology may affect the interpretation of results, so future research should increase sample size and use standardized procedures for verification. Lastly, MR analysis may produce false positives due to genetic pleiotropy, necessitating functional validation of new candidate genes.

**Conclusion**

This study provides a comprehensive integrative analysis combining single-cell transcriptomics and Mendelian randomization to uncover shared immune mechanisms underlying AS and PCa. We identified (CD8\_EM as a common immunological feature in both diseases, characterized by increased proportions and enhanced cell-cell communication. Among CD8\_EM-associated genes, HLA-B emerged as a robust genetic risk factor for PCa and was also highly expressed in AS tissues. Functional analyses revealed that HLA-B⁺ CD8\_EM cells exhibit stronger intercellular signaling and play a central role in immune modulation within diseased microenvironments. Bidirectional and validation MR analyses supported the causal role of HLA-B in PCa risk, while single-cell trajectory and pseudotime analyses highlighted its involvement in T cell development and dysfunction. These findings suggest that HLA-B may serve as a potential immunological driver and therapeutic target across pan-vascular conditions. Future studies should validate these observations in larger cohorts and explore the therapeutic implications of modulating HLA-B–mediated immune responses.

**Figure legend**

**Figure 1.** The workflow of the present study

**Figure 2**. Single-cell transcriptomics overview across various samples. (**A)**.UMAP plot colored by various cell clusters. (**B**)and(**C**). UMAP plots showing representative marker genes for each cell type. (**D**). After manual annotation, seven distinct cell groups were identified: T cells, epithelial cells, endothelial cells, vascular smooth muscle cells, B cells, monocytes, and macrophages. (**E**). The proportion of each cell type across different samples is displayed as shown in the figure.

**Figure 3:** In-depth analysis and visualization of T cell subpopulations. (**A**). KEGG and WikiPathway enrichment analysis. (B). Manually annotated T cell subpopulations.(**C**). Dot plot illustrating the expression of characteristic genes across various cell subpopulations.(**D**). Distribution of T cell subpopulations across different samples.

**Figure 4:** Cellular Differentiation and Communication Analysis in AS and PCa. (**A**). UMAP plot showing the cell differentiation trajectory. (**B, D**). Cell communication networks constructed for CD8\_CM and other cells in PCa and AS samples.(**C, E**). Differentially enriched signaling pathway analysis between CD8\_EM and other cells in PCa and AS samples.

**Figure 5** Shows the Mendelian Randomization analysis: (**A**). Bar chart of gene set enrichment analysis. (**B**). A volcano plot illustrating the association between key genes and PCa risk. (**C**). A forest plot describing the association between key genes and PCa risk.(**D**). A scatter plot showing the genetic association between "HLA-B" and PCa. (**E**). A forest plot of single nucleotide polymorphisms related to "HLA-B" and their causal relationship with PCa. (**F**). The validity and robustness of the Mendelian Randomization results, further validating the identified associations with minimal bias and confounding factors.

**Figure 6**. Validation cohort and bidirectional Mendelian Randomization (**A**). Mendelian Randomization analysis shows the association between HLA-B gene variants and PCa risk. (**B**). Reverse Mendelian Randomization illustrates the relationship between PCa and the HLA-B gene. (**C**). Scatter plot in the validation cohort displays the genetic association between "HLA-B" and PCa. (**D**). Forest plot in the validation cohort examines the causal relationship of single nucleotide polymorphisms related to "HLA-B" and PCa. (**E**). Demonstrates the validity and robustness of Mendelian Randomization in the validation cohort.

**Figure 7**. Regional association map

**Figure 8**. Single-cell transcriptome analysis reveals the critical role of HLA-B in T cell metabolism and function in PCa. (**A**). UMAP plot showing the distribution of HLA-B gene expression in cells. (**B**). Gene on/off states in pseudo-time, reflecting gene expression dynamics during cell development or transcriptional processes. (**C**). Intercellular communication between HLA-B+CD8\_EM and HLA-B-CD8\_EM cell subpopulations in PCa and AS analyzed using the CellChat tool, with the top showing results from the PCa group and the bottom showing results from the AS group. (**D**). Differential signaling pathway enrichment involving HLA-B+CD8\_EM and HLA-B-CD8\_EM cells compared to other cells, shown in order from top to bottom for the PCa group and the AS group. (**E**). This boxplot illustrates the expression levels of the HLA-B gene in PCa group and Normal Group (**F**). This boxplot illustrates the expression levels of the HLA-B gene in AS group and Normal Group.

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**Declarations**

**Ethics approval and consent to participate**

Prior to participation in this study, all patients provided informed consent for the use of their information and specimens for research purposes.

**Consent for publication**

Not applicable.

**Data availability statement**

Availability of data and materials: The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Conflict of interest**

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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**Author contributions**

Quan Zhang Study conceptualization and design. Data collection. Study analysis and interpretation of data. Draft and revise manuscript. Prepare tables and figures. Approved submitted version. Bochen Pan (Corresponding author) Data collection. Draft and revise manuscript. Study analysis and interpretation of data. Approved submitted version.

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**Supplementary material**

The Supplementary Material for this article can be found online at: