ORIGINAL RESEARCH Infiltration of a Unique CD8⁺CD274⁺ Cell Subgroup in Hepatocellular Carcinoma is Associated with Poor Clinical Outcomes

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Introduction: Immune checkpoint (IC) inhibitor-related immunotherapies have attracted considerable attention in hepatocellular carcinoma (HCC). High IC expression and high tumor infiltrating lymphocyte levels are the current indicators of sensitivity to IC inhibitors. Thus, it is imperative to apply precision medicine strategies for patient selection.

Methods: Six independent HCC cohorts were used for analysis at the single-cell and tissue levels. Multiplex immunofluorescence and immunochemistry staining assays were used to validate our results. A series of methodologies were used for immune-related evaluations.

Results: Herein, we uncovered a unique CD8⁺CD274⁺ cell subpopulation that is associated with tumor progression and poor survival in HCC at the single-cell level. We assessed this subset at the tissue level and found that the prognostic significance of CD274 is dependent on CD8A expression in HCC. Subsequently, we identified a unique high-risk subpopulation that showed high CD8A expression coupled with intense CD274 expression in multiple HCC cohorts. CD8A^{High}CD274^{High*} subgroup was correlated with malignant indexes and remained an independent prognostic factor when considering the influence of these indexes. Molecular characteristic analyses showed that the CD8A^{High}CD274^{High*} subgroup harbored more mutations, had higher immune response activity and presented enrichment of cancer-related biological processes. Moreover, this high-risk subpopulation in HCC was characterized by high immune cell infiltration, low tumor purity, and enrichment of cancer-related signatures. Finally, cases with this phenotype demonstrated higher immunomodulator and IC levels and greater sensitivity to IC inhibitors.

Conclusion: Our findings illustrate that some HCC patients may have a poor prognosis despite high CD8⁺ T-cell infiltration. These patients would probably benefit from IC inhibitor-based combination treatment.

Keywords: hepatocellular carcinoma, CD8A, CD274, molecular subtype, tumor microenvironment, immune overdrive

Introduction

Liver cancer is the seventh most common cancer type in the world and remains a global health challenge in the 21st century. Hepatocellular carcinoma (HCC) has a high degree of malignancy and accounts for greater than 90% of primary liver cancers.¹ In addition to surgical treatment, new drug therapies for HCC have attracted much attention. With the launch of precision medicine, especially the advent of the immunotherapy era, increasing attention has been given to studies of immune checkpoint (IC) inhibitors in the precision treatment of HCC.² Programmed death receptor-1 (PD-1) is a critical IC that inhibits T-cell activation and cytokine production through its two ligands, CD274 (commonly known as PD-L1) and PD-L2. CD274 inhibition is one of the most popular clinical research and treatment strategies. CD274

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inhibitors can reactivate the immune response function of T cells on tumors by inhibiting IC activity, resulting in antitumor effects. However, the benefits of CD274 inhibitors cannot be accurately captured in clinical practice. Currently, patients who are sensitive to CD274 or PD-L1 antibodies mainly have the following characteristics: 1) high CD274 expression, ie, patients with an expression rate of 1% can be treated with this therapy, but those with an expression rate over 50% exhibit better efficacy; 2) high tumor mutational burden, ie, treatment is effective in patients with a tumor mutational burden of > 20/Mb; 3) high levels of tumor infiltrating lymphocytes.^{3,4} However, the efficacy of these inhibitors in the treatment of HCC remains limited. Thus, it is now imperative to apply precision medicine strategies for patient selection.

The immune response is complex and may be highly heterogeneous among tumors, even tumors from the same cancer type, probably due to the complex and heterogeneous tumor microenvironment (TME).^{5,6} Thus, TME-based molecular subtyping systems have been developed for precision therapy in several cancer types. Previous studies on the Immunoscore showed that high immune cell infiltration predicts prolonged survival in multiple cancer types, including HCC.^{7,8} However, recent studies have identified unique subgroups with high risk that feature high levels of immune infiltration and IC expression, and only two genes need to be examined to stratify patients into these groups.^{5,9} Hence, the identification of a novel TME-based immune molecular subtyping system is urgently needed in HCC to personalize cancer immunotherapies.

In this study, we analyzed tumors at the single-cell level and identified a unique CD8⁺CD274⁺ cell subpopulation in HCC. We assessed this subset at the tissue level and found a risk subpopulation among HCC patients showing high CD8A expression coupled with intense CD274 expression. Furthermore, molecular and immune characteristics of this subgroup were explored in HCC datasets.

Materials and Methods

Data and Resources

We downloaded The Cancer Genome Atlas (TCGA) HCC primary tumor RNA-Seq data, mutation and related clinical information from the Genomic Data Commons (GDC) Data Portal (<u>https://portal.gdc.cancer.gov/</u>). Overall survival (OS) data of the TCGA cohort were obtained from the integrated TCGA Pan-cancer clinical data resource.¹⁰ HCC microarray expression data GSE76427 were available on GEO (<u>http://www.ncbi.nlm.nih.gov/geo</u>). HCC RNA-Seq data LINC-JP were downloaded from ICGC (<u>https://dcc.icgc.org/releases/current/Projects/</u>). HCC protein data were obtained from CPTAC (<u>https://proteomics.cancer.gov/programs/cptac</u>). HCC single-cell RNA sequencing (scRNA-Seq) data were obtained from a previous study (<u>https://db.cngb.org/search/project/CNP0000650</u>). A total of 80 pathological paraffin sections and medical records were obtained from patients who underwent radical resection at The Second Affiliated Hospital of South China University of Technology (Guangzhou, China) between December 2016 and December 2020 (<u>Table S1</u>). These patients did not receive immunotherapy, chemotherapy, or radiofrequency ablation. All experimental procedures were approved by the Ethics Committee of The Second Affiliated Hospital of South China University of Technology. Written informed consent was provided by all patients before the study.

Single-Cell Level Analysis

The HCC scRNA-Seq dataset CNP0000650 contains information on 19 samples, including 6 recurrent tumor samples and 12 orthotopic tumor samples. We selected tumor in situ samples for follow-up analysis. The specific data quality control process was strictly based on the literature published in a previous study, and 12,185 cells were selected for further analysis. The Seurat R package was used to perform unsupervised clustering of the single cells using the read count matrix as input.¹¹ For the clustering of cells, the top 20 principal components (PCs) were selected for principal component analysis (PCA) with a resolution parameter equal to 0.8. For the clustering of T lymphocytes, the top 12 PCs were selected with a resolution parameter equal to 0.6. The Scissor R package used bulk phenotypes to identify cell subsets that are highly correlated with phenotypes from scRNA-seq data.¹² To obtain the appropriate sparsity, we finally determined the penalty value as alpha = 0.02 after many iterations. The CellChat R package was used to infer, visualize

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Mutation Analysis

by using Monocle2.14

The Maftools package from R software was used to analyze Mutation Annotation Format files in the HCC CD8A^{High}CD274^{High*} and other subgroups of the TCGA HCC cohort as previously described.^{15,16}

Enrichment Analysis

GSEA 4.3.0 software was used to estimate the recurrence risk and assess the enrichment of biological processes in the HCC CD8A^{High}CD274^{High*} subgroup. Predefined gene sets are available on the Molecular Signatures Database (MSigDB) website (https://www.gsea-msigdb.org/gsea/msigdb/collections.jsp). HOSHIDA_LIVER_CANCER_LATE_RECURRENCE_UP and WOO_LIVER_CANCER_RECURRENCE_UP were defined as gene sets whose expression correlated with a higher risk of late recurrence of HCC. Gene ontology biological process and KEGG pathway database and hallmark gene sets, respectively. Single-sample gene set enrichment analysis (ssGSEA) was performed via the GSVA package in R software to calculate the enrichment degree of KEGG_PATHWAYS_IN_CANCER, HALLMARK_ANGIOGENESIS, and KEGG_JAK_STAT_SIGNALING_PATHWAY signatures in the TCGA HCC cohort.

Immune Evaluation

Four independent approaches were used to evaluate immune infiltrates in the TCGA HCC cohort: ESTIMATE, analysis of a 22 immune cell type signature (LM22) via CIBERSORTx, Pan-cancer immunogenomics and TIMER2.^{17–21} Multiplatform integrative molecular subtyping of the TCGA HCC cohort and revealed the following groups: iClust1, iClust2 and iClust3.²² Six immune subtypes were identified in 33 TCGA cancer types: C1 (wound healing), C2 (IFN-g dominant), C3 (inflammatory), C4 (lymphocyte depleted), C5 (immunologically quiet) and C6 (TGF- β dominant).²³ Four immune/fibrotic TME subtypes were detected and conserved in TCGA human cancers: immune-enriched/fibrotic (IE/F), immune-enriched/non-fibrotic (IE), fibrotic (F) and immune-depleted (D).²⁴ Cancer-associated fibroblast (CAF) level estimation was performed with the MCPCOUNTER and EPIC algorithms via the TIMER 2.0 (http://timer.cistrome.org/).^{25,26} Immunomodulator genes were collected from a previous TCGA immune subtype study.²³ Tumor Immune Dysfunction and Exclusion (TIDE) analysis was performed at http://tide.dfci.harvard.edu/. The TIDE score computed for each tumor can serve as a surrogate biomarker to predict the response to IC blockade. A more negative TIDE score probably predicts a better IC blockade response.²⁷

Immunohistochemistry (IHC) Assay

Formalin-fixed and paraffin-embedded pathological wax blocks were sequentially sectioned. Next, the slices were dewaxed, dehydrated and subjected to EDTA (pH = 8.0) at high temperature and pressure for 3 min. Then, the endogenous peroxidase activity was blocked by incubation with $0.3\%H_2O_2$ at $37^{\circ}C$ for 10 min, sealed with 5% goat serum at $37^{\circ}C$ for 20 min. Primary antibodies, including anti-CD8A (China, LBP, IHC-R024) and anti-CD274 (UK, Abcam, ab210931), were incubated overnight at 4°C and finally stained with DAB. All slides were counterstained with hematoxylin and sealed with neutral resin. A Lecai SOPTOP HS6 scanner, ImageJ and scoring systems were used to analyze the results according to a previous study.²⁸

Multiplex Immunofluorescence (mIF) Assay

Multiplex staining of FFPE tissue was performed using a multiple fluorescent immunohistochemical staining kit (absin, abs50012) according to the manufacturer's instructions. CD274 (UK, Abcam, ab210931) and CD8 (China, LBP, IHC-R024) antibodies were sequentially applied, followed by horseradish peroxidase-conjugated secondary antibody incubation and tyramide signal amplification. After the sections were washed 3 times with PBS, multispectral images were obtained by using the Mantra System (Zeiss LSM880), which captures fluorescent spectra at 20-nm wavelength intervals from 420–720 nm with identical exposure times. For each slide, five fields of immune cell-enriched tumoral area were selected for image capture.

Statistical Analysis, Code Availability and Visualization

OS was evaluated by Kaplan–Meier survival analysis and Log rank test as previously described:^{9,16} HCC samples of each cohort were divided into high and low expression groups according to CD8A or CD274 level, and *P* values based on two groups were examined based on Kaplan–Meier survival analysis. The lowest log-rank *P* value was selected from the 10th to 90th percentiles of these samples. Patient survival was analyzed by R software 4.1.0 and GraphPad Prism 9. The Mann–Whitney test was used in the two-group comparisons, and the Kruskal–Wallis test was used in the three-group comparisons in GraphPad Prism 9. All reported *P* values were two-sided. Chi-squared test analyses were performed using GraphPad Prism 9. Analyses were performed using R software 4.1.0 with the GSVA (1.40.1) package (ssGSEA) for Pan-cancer immunogenomics and enrichment score calculation; the survival (3.2–11) package for two-group OS analysis; the ESTIMATE (1.0.13) package for ESTIMATE immune score and tumor purity estimation; the Seurat (4.0.3) package for clustering and annotation of HCC single-cell samples; and the Scissor (2.0.0) package for identifying high-and low-risk subpopulations at the single-cell level by integrating bulk sequencing data. The Monocle (2.18.0) package can be used to model the changes in cells by comparing cell features over time. A *P* value less than 0.05 was regarded as statistically significant. Figure generation and data analysis were performed with GraphPad Prism 9 and R software 4.1.0.

Results

A Unique CD8⁺ T-Cell Subpopulation is Related to a Poor Prognosis in HCC

To construct a global tumor niche atlas, we identified and visualized 9 major clusters using the T-distributed stochastic neighbor embedding (t-SNE) method (Figure 1A). Nonimmune cells primarily consisted of endothelial cells, hepatic stellate cells, apparently normal epithelial cells and HCC malignant cells (Figure 1B). Furthermore, the most abundant immune cells in the TME of HCC patients were CD4⁺/CD8⁺ lymphocytes, followed by myeloid cells, NK cells and B cells (Figure 1B). To determine the molecular characteristics of cell subsets of the HCC TME that are related to the prognosis, we performed an analysis using the scissors algorithm. Notably, we found that some malignant tumor cells and myeloid cells in the HCC TME are related to a poor prognosis in patients. Interestingly, among prognosis-related cells, most CD8⁺ T cells were associated with poor survival (Figure 1C and D). Recently, researchers found that tumor occurrence and progression are the results of the joint action of tumors and other nonmalignant stromal cells in the TME.²⁹ Thus, the CellChat algorithm was used to analyze and construct the possible cell communication network between various cell populations in the HCC TME (Figure 1E). Consistent with the abovementioned findings, cell communication levels among three subsets mainly related to a poor prognosis (CD8⁺ T cells, malignant tumor cells and myeloid cells) suggest that the three are closely related in terms of the number and intensity of receptor-ligand relationships (Figure 1F). The presence of leukocytes called tumor infiltrating lymphocytes (TILs) in tumors is associated with better treatment outcomes for most types of cancer. CD8⁺ TILs can find tumor antigens on the surface of cancer cells and destroy them.³⁰ To explore this phenomenon, we extracted the most active receptor-ligand relationship pairs between the three cell clusters (Figure 1G). In addition to activating MIF and CXCL signaling pathways related to macrophage and T-cell recruitment, malignant tumor cells and myeloid cells mainly induce signal activation of IL10 and TGF- β to regulate CD8⁺ T cells. These pathways are related to the functional depletion of CD8⁺ T cells.³¹ Thus, the HCC TME contains a series of CD8⁺ T cells related to a poor prognosis and pathways that repress antitumor immune responses.

$\text{CD8}^{+}\text{CD274}^{+}$ T Cells are Related to a Poor Prognosis and Tumor Development in HCC

To further locate the previously identified CD8⁺ TIL cell subsets related to a poor prognosis, we further performed unsupervised clustering of T cells (Figure 2A). The reclustering of T cells revealed 14 populations, including two NK subtypes (CD16 and CD160), four subtypes of CD4⁺ T cells (CD4-CCR7, CD4-IL2, Treg-FOXP3, and Treg-LAYN), and seven subtypes of CD8⁺ T cells (CD8-CCR6, CD8-LAG3, CD8-GZMK, CD8-GZMH, CD8-CD274, CD8-XCL1, CD8-NR4A1) (Figure 2A). These results showed that the CD8⁺ T subsets associated with a poor prognosis are mainly CD8⁺CD274⁺ cells, and these cells highly express a variety of IC inhibitory molecules (Figure 2B and C). Further pseudotime analysis suggested that CD8⁺ cells tended to change to the CD8⁺ CD274⁺ phenotype with tumor development (Figure 2D). Correspondingly, the cytotoxic function of CD8⁺



Figure I scRNA-seq profiling of the tumor ecosystem in primary HCC. (A) T-distributed stochastic neighbor embedding (t-SNE) plot showing the annotation and color codes for cell types in the HCC ecosystem. (B) The expression of marker genes in the indicated cell types. (C) The relationship between cells and the prognosis of HCC patients. Red represents poor survival, blue represents favorable survival, and gray is meaningless. (D) The proportions of prognosis-related cells in myeloid cell, malignant cell and CD8⁺ T-cell types. (E) The regulatory network between various groups of cells in the HCC tumor microenvironment based on the receptor ligand exchange mode. The connecting line between cell groups represents the regulatory relationship between the two cells, and the thickness of the line represents the number of receptor ligand pairs (left) and interaction intensity (right). (F) The number and intensity of regulation between receptor–ligand pairs among cell groups. (G) The regulatory relationship of a specific signaling pathway between cell groups.

cells was gradually lost, and the cells were gradually depleted (Figure 2E). We next investigated the transcriptional changes associated with transitional states and observed that the $CD8^+$ T-cell clusters could be categorized into five phase groups (Figure 2F). $CD8^+CD274^+$ cells were predominantly Phase 5 cells. Phase 5 was characterized by high levels of T-cell exhaustion-related TFs, including TOX, TOX2, HOPX and IRF8, and genes involved in the PD-1 and mitotic pathways, further confirming the exhausted state of these cells.³² Hence, we found an exhausted $CD8^+$ T-cell phenotype that may be related to the poor prognosis of HCC at the single-cell transcriptome level. We further confirmed its authenticity in HCC tumor tissue by using mIF staining assays (Figure 2G). We found that $CD8^+CD274^+$ cells appeared in the tumor tissues of patients with HCC, featured by "circles" because they are mainly expressed on the cell surface. These results suggest that this unique $CD8^+$ T-cell type may play important roles in HCC; in particular, the exact correlation of CD8A and CD274 should be further explored.

The Prognostic Significance of CD274 is Associated with CD8A Expression in HCC

Analyses of single-cell data suggest the importance of $CD8^+CD274^+$ T cells, particularly the unique relationship between CD8A and CD274 in the TME of HCC. Previous studies have explored correlations between CD8A and macrophage-expressed CD274 in colorectal cancer (CRC).⁵ Thus, TCGA HCC RNA-Seq and clinical data were used to assess the OS of different CRC subpopulations based on CD8A and CD274 expression. First, we divided patients into two groups based on CD8A and CD274 expression levels based on our previous TME-related studies.^{9,33} High CD8A expression was associated with favorable survival in HCC according to several peaks with optimal cutoffs of *Pscore* > 1.3 (equivalent to



Figure 2 The subtypes of T cells and cell transition states in primary HCC. (A) t-SNE projections of subclustered T cells, labeled in different colors. Cell type annotations are provided in the figure. (B) The expression of selected gene sets, including markers of naïve and resident cells, inhibitory molecules, cytokines, costimulatory markers, transcription factors, and cell type markers, in T-cell subtypes. (C) The t-SNE plot showing cell origins by color, showing the relationship between cells and the prognosis of HCC patients. (D) Pseudotime-ordered analysis of CD8⁺ T cells from primary HCC samples. (E) 2D pseudotime plot showing the dynamics of cytotoxic (upper panel) or exhausted signals (lower panel) in CD8⁺ T cells from primary HCC samples. (F) The dynamic changes in gene expression along with pseudotime (lower panel). The distribution of CD8 subtypes during the transition (divided into 5 phases), along with pseudotime. Subtypes are labeled by colors (upper panel). (G) Immunofluorescence images showing the colocalization of CD274 and CD8 in HCC tissue. Scale bar for all images = 50 μ m.

Log rank test P < 0.05) (Figure 3A). High expression of CD274 was also associated with favorable OS in HCC according to only one peak with an optimal cutoff of *Pscore* > 1.3 (Figure 3B and C). Then, we applied the commonly used median value to define CD8A high and low expression groups and performed survival analysis based on the optimal cutoff of CD274 expression. CD274^{High} status was still associated with favorable survival in CD8A^{Low} HCC, even though P >



Figure 3 The prognostic significance of CD274 is dependent on CD8A expression in HCC. (**A**) Dotplots showing HR (up) and Pscore (bottom) for different CD8A expression cutoffs in TCGA HCC patients. The dotted line represents HR = 1 (up) and Pscore= 1.3 (equivalent to P<0.05, bottom). (**B**) Similar to (**A**), but for CD274. The Orange arrow represents the optimal cutoff. (**C**) Kaplan-Meier plot showing the OS for CD274 in TCGA HCC patients according to the optimal cutoff. (**D**) Pearson correlation between CD8A and CD274 expression in TCGA HCC patients. Vertical blue lines represent the median value of CD8A. The horizontal dark blue line represents the optimal cutoff of CD274 in CD8A^{Low} patients. (**E**) Similar to (**A**), but for CD274 in CD8A^{Low} patients. (**F**) Kaplan-Meier plot showing the OS for CD274 in CD8A^{Low} patients according to the optimal cutoff. (**G**-1) Similar to (**D**-**F**), but for CD274 in CD8A^{High} cases. (**J** and **K**) Kaplan-Meier plot showing the OS for CD274 in CD8A^{High} patients (**J**) or CD8A^{Low} patients (**K**) in the LIRI-JP HCC cohort according to the optimal cutoff.

0.05 (Figure 3D–F). However, among CD8A^{High} HCC patients, the CD274^{High*} subgroup had poorer OS than the CD274^{Low*} subgroup (Figure 3G–I). Additional LIRI-JP HCC expression data were analyzed, and similar results were obtained (Figure 3J and K). Collectively, these data indicate that CD274 expression predicts the opposite OS trend in the CD8A^{Low} and CD8A^{High} HCC groups.

CD8A^{High}CD274^{High*} Patients Represent a Novel High-Risk Subgroup in HCC

To validate the subtyping effect of CD8A^{High}CD274^{High*} patients based on our approach in the aforementioned analyses, we identified this subgroup in the TCGA HCC, GSE76427 and LIRI-JP cohorts, and this subpopulation showed the worst survival outcomes (Figure 4A–C). Moreover, this correlation was also confirmed at the protein level by analyzing the CPTAC HCC dataset (Figure 4D). Hence, a unique high-risk subpopulation, CD8A^{High}CD274^{High*} was identified in HCC. Moreover, GSEA was performed to determine whether the signatures associated with HCC recurrence were enriched in CD8A^{High}CD274^{High*} patients. We found that the CD8A^{High}CD274^{High*} subgroup was significantly correlated with increased risks of early and late recurrence in the TCGA HCC dataset (Figure 4E). These results demonstrated that the CD8A^{High}CD274^{High*} subgroup is associated with poor survival and recurrence.

To evaluate the prognostic value of CD8A/CD274, we explored the clinical features among these subtypes in the TCGA HCC dataset. First, CD8A^{High}CD274^{High*}T3+T4, tumor node metastasis (TNM) stage III+IV or vascular invasion were found to be associated with poor survival according to the univariable Cox analysis (Figure 4F). Then, multivariable Cox analysis revealed that CD8A^{High}CD274^{High*}T3+T4 or N1 stage III+IV were associated with poor survival (Figure 4G). Thus, further multivariable Cox analysis was performed based on these three indexes, and CD8A^{High}CD274^{High*} remained an independent prognostic factor when considering the influence of the other two indexes (Figure 4H). Finally, the CD8A^{High}CD274^{High*} patients were observed to have a higher tumor stage, higher tumor grade and vascular invasion (such as stage III disease, grade 3 disease and macrovascular invasion) (Figure 4I).

To further validate these genomic findings, primary tumor tissues were collected from 80 HCC patients at The Second Affiliated Hospital of South China University of Technology. As shown in Figure 5A–D, the CD8A^{High}CD274^{High*} cases showed increased immune cell infiltration, and this phenotype indicated the worst survival outcomes and highest relapse risk, which was consistent with our observations of public HCC cohorts. Importantly, cases with CD8A^{High}CD274^{High*} remained an independent prognostic factor after multivariable Cox analysis (Figure 5E and F). Taken together, our analyses based on mRNA and protein expression data show that the CD8A^{High}CD274^{High*} subgroup has a higher risk of poor clinical outcomes than other patients with HCC.

Molecular Characteristics of the CD8A^{High}CD274^{High*} Subgroup

We first estimated the gene mutation rates. Missense variations were the most common mutation type in both subgroups. We found significantly higher mutation rates in the CD8A^{High}CD274^{High*} subgroup than in the other patients (Figure 6A and B). In particular, TP53 mutation was more common in the CD8A^{High}CD274^{High*} subgroup than in the other subgroups. Next, GSEA was performed based on GO BP, KEGG and Hallmark signatures and demonstrated that immune-related pathways and cancer-related pathways were enriched in the CD8A^{High}CD274^{High*} subgroup (Figure 6C and E). For example, the T-cell receptor signaling pathway, E2F target and cell cycle signatures were enriched in the CD8A^{High}CD274^{High*} subgroup (Figure 6F). These results suggest that the CD8A^{High}CD274^{High*} subgroup an active immune response and cancer-related biological processes.

The CD8A^{High}CD274^{High*} High-Risk Subpopulation is Characterized by an Immune Overdrive TME Phenotype

To further investigate the immune activity level in the CD8A^{High}CD274^{High*} high-risk subgroup in HCC, we first estimated the total immune infiltration level in the TCGA HCC dataset based on mRNA expression and observed the highest overall immune infiltration level in the CD8A^{High}CD274^{High*} subgroup compared with other subgroups (Figure 7A). Then, the total immune infiltration level was estimated in the CPTAC HCC dataset based on protein expression, and similar results were observed (Figure 7B). Moreover, the CD8A^{High}CD274^{High*} subgroup showed the



Figure 4 CD8A^{High}CD274^{High*} patients represent a novel high-risk subtype in HCC. (**A**) Pearson correlation between CD8A and CD274 expression in TCGA HCC patients (left). Vertical blue lines represent the median value of CD8A. The horizontal dark blue line represents the optimal cutoff of CD274 in CD8A^{High} patients. Kaplan–Meier plots showing the OS for each CD8A/CD274 subtype in TCGA HCC patients. (**B**–**D**) Similar to (**A**), but in GSE76427 (**B**), LIRI-JP (**C**) and CPTAC protein (**D**) HCC cohorts. (**E**) GSEA plots showing that the CD8A^{High}CD274^{High*} subgroup was significantly correlated with a higher risk of recurrence in TCGA HCC patients. (**F** and **G**) Univariable and multivariate COX regression models showing the effect of the CD8A/CD274 stratification system, TNM stage, grade and vascular invasion indexes on OS in TCGA HCC patients. (**H**) Multivariate Cox regression models showing the effect of CD8A/CD274, T stage and N stage on OS (**G**) in TCGA HCC patients. (**I**) Pie charts showing the chi-squared test of clinicopathologic factors for CD8A/CD274 stratification system TNM stage, grade and vascular invasion indexes from the TCGA HCC cohort.



Figure 5 Wet-laboratory experimental validation in our independent HCC cohort. (A) IHC staining assay of each CD8A/CD274 subtype in the validation HCC cohort. Scale bar for all images = $100 \mu m$. (B) Kaplan–Meier plots showing the OS for each CD8A/CD274 subtype in the validation HCC cohort. (C and D) Stacked bar plot showing the chi-squared test of survival (C) and recurrence (D) status for each CD8A/CD274 subgroup in the validation HCC cohort. (E and F) Univariable and multivariate COX regression models showing the effect of the CD8A/CD274 stratification system, TNM stage and grade on OS in our validation cohort.

highest levels of five major immune cell types: B cells, CD4⁺ T cells, CD8⁺ T cells, Tregs and macrophages (Figure S1). We next estimated tumor purity using the ESTIMATE algorithm, and the CD8A^{High}CD274^{High*} subgroup had the lowest levels of tumor purity among these subgroups (Figure 7C and D). These data indicate that CD8A^{High}CD274^{High*} HCC is characterized by a TME with high immune cell infiltration and low tumor purity.

We next wondered why the CD8A^{High}CD274^{High*} subgroup shows a high risk of poor clinical outcome despite high immune cell infiltration and low tumor purity. We investigated the proportions of the previously described HCC and immune subtypes in each subgroup of the CD8A/CD274 subtyping system.^{22–24} We first analyzed the TCGA HCC subtyping system and observed that the CD8A^{High}CD274^{High*} high-risk subpopulation largely overlapped with the iCluster:1 subtype (Figure 7E). Then, TCGA immune and TME subtypes were used, and the CD8A^{High}CD274^{High*} high-risk subpopulation showed a high proportion of samples with the C2, IE/F and IE subtypes (Figure 7F–G). Interestingly,



Figure 6 Molecular characteristics of the CD8A^{High}CD274^{High*} subgroup. (**A**) Mutation profiles of the 20 most mutated genes in the CD8A^{High}CD274^{High*} subgroup or non-CD8A^{High}CD274^{High*} samples among TCGA HCC tumor samples. (**B**) Boxplot showing the mutation rate of total genes grouped by CD8A^{High}CD274^{High*} and non-CD8A^{High}CD274^{High*} patients from the TCGA HCC cohort. (**C**) Barplot showing the top 20 significantly enriched GO BP signatures in the CD8A^{High}CD274^{High*} subgroup from the TCGA HCC cohort. (**E**) Barplot showing the top significantly enriched HALLMARK signatures in the CD8A^{High}CD274^{High*} subgroup from the TCGA HCC cohort. (**E**) Barplot showing the top significantly enriched HALLMARK signatures in the CD8A^{High}CD274^{High*} subgroup from the TCGA HCC cohort. (**E**) Barplot showing the top significantly enriched KEGG signatures in the CD8A^{High}CD274^{High*} subgroup from the TCGA HCC cohort. (**E**) Barplot showing that the CD8A^{High}CD274^{High*} subgroup was significantly correlated with higher T-cell receptor signaling pathway, E2F target and cell cycle signatures in TCGA HCC patients.

Figure 7 TME characteristics of the CD8A^{High}CD274^{High*} subgroup. (**A** and **B**) Boxplots showing the total immune infiltrates based on ESTIMATE, CIBERSORT×, Pancancer immunogenomic analysis signature and TIMER for each CD8A/CD274 subtype in TCGA (a) and CPTAC protein (b) HCC cohorts. (**C** and **D**) Boxplots showing the total tumor purity based on the ESTIMATE algorithm for each CD8A/CD274 subtype in the TCGA (c) and CPTAC protein (d) HCC cohorts. (**E**–**G**) Stacked bar-plot showing the Chi-squared test of TCGA HCC subtype (e), TCGA immune subtype (f) and TCGA TME subtype (g) for each CD8A/CD274 subgroup in TCGA HCC cases. (**H**) Boxplots showing the levels of KEGG pathways in cancer and hallmark angiogenesis signatures for each CD8A/CD274 subtype in the TCGA HCC cohort. (**J**) Boxplots showing the expression of MYBL2, PLK1, and MKI67 for each CD8A/CD274 subtype in the TCGA HCC cohort. (**J** and **K**) Similar to (**H**), but for the KEGG JAK/STAT signaling pathway and hallmark TGF-β signaling signatures. (**L**) Boxplot showing the expression of ICs for each CD8A/CD274 subtype in the TCGA HCC cohort. (**M**) Boxplot showing the TIDE score for each CD8A/CD274 subtype in the TCGA HCC cohort (left). Stacked bar-plot showing the Chi-squared test of IC inhibitor-response status for each CD8A/CD274 subgroup in the TCGA HCC cohort (right).

iCluster:1 cases exhibit specific features, such as high tumor grade and macrovascular invasion. We found that the CD8A^{High}CD274^{High*} high-risk subpopulation demonstrated higher levels of KEGG pathways in cancer and hallmark angiogenesis signatures than the CD8A^{High}CD274^{Low*} and CD8A^{Low} subpopulations (Figure 7H). In addition, the iCluster:1 phenotype was associated with the upregulation of proliferation marker genes, such as MYBL2, PLK1, and MKI67. Our analyses validated that these genes were overexpressed in the CD8A^{High}CD274^{High*} high-risk subpopulation (Figure 7I). Moreover, C2 has a strong CD8 signal and a high proliferation rate. Immunomodulator genes are most highly expressed in the C2 subtype. IE/F cases are distinguished by angiogenesis and CAF activation. IE has high levels of immune infiltration, and assessment of activated signaling pathways revealed increased JAK/STAT pathway activation. which may be associated with increased T-cell activity. Further analyses showed that the CD8A^{High}CD274^{High*} high-risk subpopulation had the highest levels of the KEGG JAK/STAT signaling pathway and hallmark TGF-β signaling signature enrichment in the TCGA HCC dataset, and this population also had high expression of ICs (Figure 7J-L). The CD8A^{High}CD274^{High*} high-risk subpopulation also exhibited the highest level of CAFs in the TCGA HCC cohort (Figure S2). Immunomodulators play a critical role in cancer immunotherapy, and many immunomodulator agonists and antagonists are currently being used in clinical oncology.²³ Notably, most of the immunomodulator genes were overexpressed in the CD8A^{High}CD274^{High*} high-risk subpopulation compared with the other subgroups (Figure S3). Remarkably, the CD8A^{High}CD274^{High*} high-risk subpopulation had the lowest TIDE score and a higher proportion of IC inhibitor-sensitive cases in the TCGA HCC cohort (Figure 7M).

Together, these results demonstrate that both immune cell infiltration and IC genes are overrepresented in the CD8A^{High}CD274^{High*} high-risk subpopulation, reflecting an immune overdrive TME phenotype. Furthermore, both the JAK/STAT pathway and TGF- β signaling are active in the CD8A^{High}CD274^{High*} high-risk subpopulation, reflecting a counterbalancing phenotype between immunostimulatory and immunosuppressive mechanisms.

Discussion

In this scenario, we uncovered a unique CD8A^{High}CD274^{High*} subpopulation at the HCC single-cell level. This subpopulation was next identified at the tissue level and associated with poor clinical outcomes. Moreover, the CD8A^{High}CD274^{High*} high-risk subpopulation featured an immune overdrive phenotype in HCC.

CD8⁺ T cells are characterized by CD8A gene expression, which is a cytotoxic T-cell lineage marker. The CD274 gene is located on chromosome 9p24.1 and encodes the PD-L1 protein. The critical IC PD-L1 suppresses antitumor immunity through an interaction with its receptor PD-1 on the cell membrane. The binding of PD-L1 and PD-1 results in the transmission of an inhibitory signal, which reduces the proliferation of antigen-specific T cells in lymph nodes. Blockade of the PD-L1/PD-1 interaction promotes the antitumor immune response and provides survival benefits for patients. This is a major breakthrough in cancer therapy. Interestingly, PD-L1 also has nonimmune checkpoint functions; it harbor nuclear transcriptional activity and is involved in the pyroptosis pathway.³⁴ Several studies have noted that a high CD8⁺ T-cell level is associated with a longer OS duration.^{7,8} Interestingly, recent studies have revealed that a series of special CD8 T-cell subpopulations were associated with poor prognosis despite high immune cell infiltration in digestive system cancers.^{5,35} However, these analyses were only performed at the tissue level. The availability of singlecell transcriptomic data provides an unprecedented opportunity to understand the TME in great depth. Through comprehensive analysis of scRNA-seq data from 12 HCC cases, we realized the comprehensive molecular characteristics of the TME of liver cancer, including malignant tumor cells and nonmalignant stromal cells. On this basis, the precise, key cell subpopulation causing the poor prognosis of HCC patients includes some malignant tumor cells and most monocytes. This finding is consistent with previous research results and demonstrates the accuracy of our analysis and methods to a certain extent.^{36,37} However, we accidentally found that some $CD8^+$ T cells are strongly associated with a poor prognosis. Moreover, we found that $CD8^+$ T cells gradually tended to exhibit a change in the $CD8^+CD274^+$ phenotype in the process of tumor progression based on cell communication and pseudotime analyses. The important feature of this phenotype is overexpressed multiple ICs, and this characteristic is probably related to the imbalance of IL10, TGF-β and arginine metabolic pathways mediated by tumor cells and macrophages.³⁸ Based on the mIF staining assay, we identified a T-cell phenotype with high expression of both CD8 and CD274. Previous studies have shown that CD8⁺CD274⁺ T cells have predictive value in NSCLC patients who received IC inhibitors.³⁹ Reportedly, PD-L1 molecules are expressed on a higher percentage of peripheral blood CD8⁺ T cells in advanced gastric cancer patients than in those of healthy controls.⁴⁰ In addition, chemoradiation and immunotherapy decreased the proportion of CD8⁺CD274⁺ T cells in cervical biopsy specimens.⁴¹ Thus, CD8⁺CD274⁺ T cells probably have an immunosuppressive role in the TME, although CD8+ T cells are generally considered antitumor cells.³⁴ PD-L1 inhibits antitumor immunity via interaction with its receptor PD-1 membrane. Therefore, CD8⁺ CD274⁺ T cells may be adjacent to PD-1⁺ cells to play an immunosuppressive role.⁴² Moreover, in our analysis at the tissue level, we found that the CD274^{high*} group was associated with favorable survival according to the best cutoff value in HCC. Only patients with CD274^{high*} expression in the CD8A^{high} subgroup showed poor survival and high recurrence risk, indicating that the prognostic significance of CD274 is dependent on CD8A expression in HCC. These findings are consistent with the results of our single-cell level analysis and similar to those of previous TCGA-related studies of melanoma and CRC.⁵

Based on our analyses using TCGA mRNA and CPTAC protein HCC datasets, this CD8A^{High}CD274^{High*} high-risk subpopulation features a TME with high immune cell infiltration and low tumor purity, respectively. In addition, this subpopulation has higher JAK/STAT pathway activation, which may be associated with increased T-cell activity.²⁴ We wondered why this subpopulation has high immune infiltration and low tumor purity but exhibits worse clinical outcomes. The CD8A^{High}CD274^{High*} high-risk subpopulation showed high expression levels of ICs, such as CD274, CTLA4, HAVCR2, IDO1, LAG3, PDCD1, PDCD1LG2, TIGIT and TNFRSF9. These ICs provide negative signals to decrease T-cell responses and limit antitumor immune responses, resulting in tumor development.⁴³ Some cell types, such as CAFs, are associated with immunosuppression and have protumorigenic and immunosuppressive properties in human cancers.⁴³ In addition, this high-risk subtype also exhibited obviously increased cancer-related biological processes, such as tumor proliferation and angiogenesis, particularly increased TGF- β signaling, which is one of the primary mechanisms mediating immune evasion.⁴⁴ The characteristics of this TME type in the CD8A^{High}CD274^{High*} subgroup are similar to the immune overdrive phenotype described in previous studies in CRC, which involves the infiltration of CD8 T cells, macrophages expressing CD724, and Tregs overexpressing CTLA4.^{5,9} In addition, a recent study suggests that CD8⁺ T cells help to induce nonalcoholic steatohepatitis (HCC) rather than invigorate or execute immune surveillance.⁴⁵ Hence, this study suggests that relying on evaluation of a single immune marker may lead to the misclassification of risk in HCC and underscores the importance of extending the Immunoscore study.

The CD8A^{High}CD274^{High*} subpopulation may provide novel guidelines for HCC patient selection in IC-related therapies. The liver is not only the largest immune organ of the body but also a well-known immune tolerance organ. The liver is exposed to foreign antigens from the gastrointestinal tract for long periods, thus forming its unique immune tolerance microenvironment.⁴⁶ Immunosuppressive cytokines play a key role in HCC tumor formation, progression, metastasis, immunosuppression and drug resistance. On the other hand, HCC has significant TME heterogeneity, and there is no difference in the outcomes of patients treated with PD-L1 inhibitors. Numerous PD-L1 inhibitors with low response rates are available; thus, the efficacy of PD-L1 inhibitors is not ideal. Major unmet challenges in HCC checkpoint immunotherapy include the discovery and validation of predictive biomarkers, advancing treatment to earlier stages of the disease, the application of treatment to patients with liver dysfunction and the discovery of more effective combinatorial or sequential approaches. To overcome this challenge, the body's immune response to the tumor is stimulated, immune tolerance is weakened, and immune cells are reinvigorated to play their antitumor role. Thus, it is necessary to identify individuals likely to benefit from immunotherapy.⁴⁷ The CD8A^{High}CD274^{High*} subgroup is characterized by high immune infiltration (such as T cells) and low tumor purity, so this subgroup represents an ideal population to target for immunotherapy. Remarkably, this subgroup showed a high expression of ICs, some of which are related to promoting tumor immune escape, such as the TGF- β pathway.⁴⁸ TIDE analysis also showed that the CD8A^{High}CD274^{High*} subgroup had a lower TIDE score and a higher proportion of IC inhibitor-response cases. In addition, the CD8A^{High}CD274^{High*} subgroup showed high levels of tumor mutations, including a high number of somatic mutations in the tumor genome of this subgroup, likely indicating the better efficacy of PD-1/PD-L1 therapy.⁴⁹ Interestingly, the CD8A^{High}CD274^{High*} status was correlated with enhancement of angiogenesis, which plays an essential role in tumor progression. Similar to IC inhibitors, antiangiogenic agents target TME components other than tumor cells and synergize with IC inhibitors by promoting CD8⁺ T lymphocyte infiltration and activation. A recent study suggested that IC inhibitor- and antiangiogenesis-based neoadjuvant/conversion therapy has good

efficacy and feasibility in late-stage gastric cancer.⁵⁰ Hence, targeting the TGF- β pathway and angiogenesis probably enhances the efficacy of IC inhibitors in CD8A^{High}CD274^{High*} HCC cases.

A limitation of our study is the lack of biopsy samples from advanced HCC patients treated with IC inhibitors. We acknowledge that it is difficult to collect samples from patients treated with IC inhibitors, including HCC patients. However, the interesting aspect and strengths of this study are clear. First, the existence of the CD8A^{High}CD274^{High*} subpopulation was validated in multiple HCC cohorts, including Western and Eastern HCC patient cohorts, and at both the mRNA and protein levels. This unique subpopulation exists at both the single-cell level and bulk tissue level. Importantly, this high CD8 T cell infiltration subpopulation is significantly associated with poor OS, which is considered the gold standard in terms of outcomes in clinical cancer trials. CD8A/CD274 stratification also remained an independent prognostic variable for survival after adjusting for other indexes. Moreover, the use of multiple methodologies of immune evaluation is likely to overcome potential biases and add confidence to our findings. Finally, this subtyping system only requires the examination of the expression levels of CD8A and CD274, which can be readily performed via IHC staining assay.

Conclusions

In summary, our comprehensive analyses uncovered a new CD8A^{High}CD274^{High*} subpopulation that could classify patients into an immune-overdrive high-risk subgroup in HCC. Notably, the combined evaluation of CD8A and CD274 expression is a strong predictor of HCC prognosis than evaluation of either marker alone. CD8 T cells overexpressing CD274 might be used as biomarkers and therapeutic targets in HCC. HCC patients in the CD8A^{High}CD274^{High*} immune-overdrive high-risk subpopulation may benefit from more active IC inhibitor-based combination treatment. This study will help in the development of related therapeutic strategies.

Abbreviations

HCC, hepatocellular carcinoma; PD-1, programmed death receptor-1; TME, tumor microenvironment; PCA, principal component analysis; PCs, principal components; CRC, colorectal cancer; MSigDB, Molecular Signatures Database; ssGSEA, single-sample gene set enrichment analysis; LM22, 22 immune cell type signature; CAF, cancer-associated fibroblasts; IHC, immunohistochemistry; mIF, multiplex immunofluorescence; OS, overall survival; t-SNE, T-distributed stochastic neighbor embedding; TILs, tumor infiltrating lymphocytes; TNM stage, tumor node metastasis stage.

Data Sharing Statement

Available public datasets are described in Materials and Methods. The data that support this study are available from the corresponding author upon reasonable request.

Ethics Approval and Consent to Participate

HCC tissues of pathological paraffin section and medical records were obtained from those who underwent radical resection at The Second Affiliated Hospital of South China University of Technology (Guangzhou, China) between December 2016 and December 2020. All experimental procedures were approved by the Ethics Committee of The Second Affiliated Hospital of South China University of Technology (Approval number: k-2022-031-02). Written informed consent was provided by all patients before the study. We confirmed that this study was conducted in accordance with the Declaration of Helsinki.

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Author Contributions

All authors made a significant contribution to the conception, study design, execution, acquisition of data, analysis and interpretation and took part in drafting and revising the article. All authors have agreed on the journal to which the article has been submitted and agree to be accountable for all aspects of the work.

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Disclosure

The authors declare that they have no competing interests in this work.

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