A stratification model of hepatocellular carcinoma based on expression profiles of cells in the tumor microenvironment

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Abstract

Background: A malignancy of the liver, hepatocellular carcinoma (HCC) is among the most common and second-leading causes of cancer-related deaths worldwide. A reliable prognosis model for guidance in choosing HCC therapies has yet to be established.

Methods: A consensus clustering approach was used to determine the number of immune clusters in the Cancer Genome Atlas and liver cancer-RIKEN, JP (LIRI_JP) datasets. The differentially expressed genes (DEGs) among these groups were identified based on RNA sequencing data. Then, to identify hub genes among signature genes, a co-expression network was constructed. The prognostic value and clinical characteristics of the immune clusters were also explored. Finally, the potential key genes for the immune clusters were determined.

Results: After conducting survival and correlation analyses of the DEGs, three immune clusters (C1, C2, and C3) were identified. Patients in C2 showed the longest survival time with the greatest abundance of tumor microenvironment (TME) cell populations. MGene mutations in Fibroblast growth factor-19 (FGF19) and catenin (cadherin-associated protein),β1 (CTNNB1) were mostly observed in C2 and C3, respectively. The signature genes of C1, C2, and C3 were primarily enriched in 5, 23, and 26 pathways, respectively.

Conclusions: This study sought to construct an immune-stratification model for the prognosis of HCC by dividing the expression profiles of patients from public datasets into three clusters and discovering the unique molecular characteristics of each. This stratification model provides insights into the immune and clinical characteristics of HCC subtypes, which is beneficial for the prognosis of HCC.

Keywords: Hepatocellular carcinoma, Immunotherapy, Tumor microenvironment, Tumor treatment, Stratification model

Background

Despite the treatments and diagnostic methods that have emerged over time for hepatocellular carcinoma (HCC), the long-term survival of HCC patients remains poor [1]. Currently, Since the development of molecular biology and molecular immunology, immunotherapy has gained considerable attention in cancer treatment. Safety and efficacy of nivolumab, an anti-PD-1 immune checkpoint inhibitor was previously evaluated in a clinical trial of advanced HCC patients and the outcome was promising [2]. Providing
novel insight into the prognosis of and treatment options for HCC was the purpose of the current study by establishing an immune-stratification model by classifying the HCC tumor microenvironment (TME) into three subtypes.

**Methods**

**Data collection and preprocessing**

We obtained transcriptional profile and mutation spectrum of TCGA (The Cancer Genome Atlas) cohort from the TCGA website (https://portal.gdc.cancer.gov/repository). Survival information of the 377 HCC samples in TCGA cohort were downloaded from the TCGA Pan-cancer Clinical Data Resource. 21 samples without survival data were excluded. We downloaded raw RNA-Seq read count data and clinical data of the LIRI_JP cohort from Liver Cancer-RIKEN,JP (LIRI_JP) dataset of the International Cancer Genome Consortium (https://dcc.icgc.org/projects/LIRI-JP). In addition, we normalized all raw counts data into transcript per million (TPM) for downstream analyses.

**Immune score, stromal score, and microenvironment cell population (MCP)-counter**

The Estimation of Stromal and Immune Cells in Malignant Tumour Tissues Using Expression Data (ESTIMATE) algorithm was used to calculate immune and stromal scores. Based on transcriptomic data, MCP-counter evaluated the absolute abundance of eight immune and two non-immune stromal cell populations by using the MCPcounter package of the R software program (R Foundation for Statistical Computing, Vienna, Austria).

**Consensus clustering for TME cell populations**

Based on the NbClust and ConsensusClusterPlus R packages, the number of clusters was determined for the TCGA and LIRI_JP cohorts using a consensus clustering algorithm. A 1000-times repetition of the process ensured the stability of classification.

**Signature genes associated with different TME subtypes**

According to the TME cell population–infiltrating patterns, patients were divided into three groups—namely, C1, C2, and C3. The expression differences between groups were characterized by log2 fold-change. Subsequently, pathway enrichment analysis was performed using the Enrichr (http://amp.pharm.mssm.edu/Enrichr/) database and the top five pathways with values of \( p < 0.05 \) were added into bubble plots.

**Identification of hub genes among signature genes of different subtypes**

The WGCNA package in R was applied to calculate the correlation coefficient based on the RNA sequencing data (expression values were log-transformed). Then, correlation data were imported into the Cytoscape version 3.0 software program (Cytoscape Consortium, San Diego, CA, USA; https://cytoscape.org/) to construct the gene co-expression network. Co-expression network edges were specified to have correlation coefficients > 0.7 for C2 and C3.

**Statistical analysis**

Normality was verified using Kolmogorov–Smirnov and Shapiro–Wilk tests (\( p > 0.05 \) signifies the variables that were normally distributed). The cutoff values of absolute abundance of TME cell populations and gene expression were evaluated using the survminer R package. A univariate Cox proportional hazards regression model was established to calculate the hazard ratios (HRs) for univariate analyses. All statistical and computational analyses were conducted by R programming (https://www.r-project.org/) and GraphPad Prism version 8.0 (GraphPad, San Diego, CA, USA; https://www.graphpad.com/), and a two-tailed value of \( p < 0.05 \) was considered to be statistically significant.

**Results**

**Landscape of TME cell populations in HCC**

Our study was systematically described with a flowchart (Fig. 1a). Totals of 345 and 202 patients from the TCGA and LIRI_JP datasets, respectively, were enrolled in this study. Based on the gene-expression matrix of several distinct experimental groups, the MCP-counter was used to determine the abundance of different TME cell populations. Subsequently, a correlation heatmap of TME cell populations in the TCGA cohort (Fig. 1b) was constructed to investigate the relationship between the eight immune and two non-immune stromal cell populations. It has been shown that there is a significant correlation between the eight immune cell types and two stromal cell types in the heatmap, and the same result has also been confirmed in the LIRI_JP cohort (Fig. 1c). Based on the abundance of TME cells in the patients, two categories were created:
Fig. 1 (See legend on previous page.)
high or low abundance based on the optimal cutoff value determined by the cutpoint function in the survminer package. The results of survival analysis indicated that high infiltration by T-cells (HR=0.65, 95% confidence interval [CI]=0.44–0.94, $p=0.036$), cytotoxic lymphocytes (HR=0.62, 95% CI=0.43–0.90, $p=0.020$), a B lineage (HR=0.65, 95% CI=0.45–0.94, $p=0.034$), CD8$^+$T-cells (HR=0.60, 95% CI=0.42–0.86, $p=0.005$), and natural killer (NK) cells (HR=0.70, 95% CI=0.49–0.99, $p=0.046$) correlated with a significantly favorable prognosis, whereas a monocytic lineage (HR=1.47, 95% CI=1.02–2.12, $p=0.043$) was associated with a poorer outcome. In the LIRI_JP group, similar results were also observed in that the prognosis was improved in conjunction with the high infiltration of T-cells, cytotoxic lymphocytes, B lineage, CD8$^+$T-cells, and NK cells. Different from the TCGA cohort, however, LIRI_JP patients with a high infiltration of myeloid dendritic cells, neutrophils, fibroblasts, and endothelial cells also had favorable outcomes. The Kaplan–Meier curve for the above-identified cell populations and that for the rest of the TCGA and LIRI_JP cohorts are shown in Fig. 1d and e, respectively.

Identification of TME subtypes associated with prognostic subtypes

According to the optimal cluster number (k = 3, Fig. 2a) determined by the NbClust and ConsensusCluster-Plus packages, patients in the TCGA cohort were classified into three groups (C1, C2, and C3). The details of K-means consensus clustering are shown in Fig. S1. A significant prognostic difference was observed ($p=0.004$, Fig. 2b) among these three groups, for C2, the median survival time (MST) is longer ($n=58$, MST=3125 days) than C1 ($n=129$, MST=1508 days) or C3 ($n=167$, MST=1560 days). Another HCC dataset from the LIRI_JP cohort was adopted to validate its prognostic value, though there was no statistical difference of the median survival time for the clusters in the LIRI-JP cohort, the similar trend that C2 has a longer median survival time for C2 than C1 or C3 can be observed ($p=0.2$, Fig. 2b). To confirm the difference between the three subtype classifications, the absolute abundance of TME cell populations for the TCGA cohort (Fig. 2c) was estimated, and results revealed that seven types of TME cells were involved (T-cells, CD8$^+$ T-cells, NK cells, cytotoxic lymphocytes, myeloid cells, and monocytic cells; all $p<0.001$) had the highest abundance in C2, while C3 had the lowest abundance of these seven TME cell types. Additionally, As for neutrophils ($p=0.114$), endothelial cells ($p=0.191$), and fibroblasts ($p=0.268$) there were no significant differences between C1 and C2. A similar result was also validated in the LIRI_JP cohort (Fig. 2d) in that the amounts of T-cells, CD8$^+$ T-cells, cytotoxic lymphocytes, NK cells, B-cell lineage cells, and monocytic lineage cells (all $p<0.001$) were highest in C2 and lowest in C3, respectively. Additionally, ESTIMATION is used to determine immune and stromal scores, and significant differences between the three clusters were found (Fig. 2e). This provides powerful evidence for the validity of the subtype classification.

TME subtypes in the TCGA cohort: clinical characteristics

Clinical characteristics of different subtypes in the TCGA cohort, including age at diagnosis, gender, TNM stage, histologic grade, vascular invasion, and genetic changes (tumor protein p53/TP53, CTNNB1, and FGF19) are shown in Fig. 3a and Table 1. According to the results of the analysis, CTNNB1 and FGF19 had different genetic alterations among the three TME types. For example, the frequency of CTNNB1 gene mutations was higher in C3 (37.7%) than C1 (13.2%) or C2 (24.1%), and the frequency of FGF19 genetic alterations was higher in C2 (20.7%) than C1 (3.9%) or C3 (4.2%). In addition, C3 had a higher number of somatic mutations per megabase (TMB) than C1 or C2 ($p<0.001$), while between C1 and C2, TMB did not differ significantly ($p=0.990$, Fig. 3b). Subsequently, a low-TMB and high-TMB group of patients was formed based on the median value of TMB, and there were significant differences in the amounts of T-cells ($p=0.022$), monocytic lineage cells ($p<0.001$), myeloid dendritic cells ($p<0.001$), neutrophils ($p=0.037$), endothelial cells ($p<0.001$), and fibroblasts ($p<0.001$) between the two groups (Fig. 3c).

Clinical characteristics of TME subtypes in the LIRI_JP cohort

The clinical characteristics of different subtypes in the LIRI cohort, as well as age, gender, and stage of the tumor at diagnosis, are shown in Table 2. The results suggested that age, gender, and tumor stage showed no significant differences among the three TME subtypes.

Transcriptome feature of the TME subtypes

Differential expression analysis was conducted according to the cutoff criteria of an adjusted $p$ value of <0.01 and $\log_2$ (fold-change) of $\geq 1$. Differentially expressed genes (DEGs) in C1 and C2, C2 and C3, and C1 and C3, respectively, were 400, 781, and 358 (Fig. 4a). Then, after combining the high-expression and low-expression genes and deleting the duplication genes, signature genes were identified, and the numbers of signature genes in C1, C2, and C3 were 64, 213, and 142, respectively (Figs. 4b and 4c). A heatmap was then constructed to visualize the expression level of signature genes of C1, C2, and C3 (Fig. 4d).
Fig. 2 Unsupervised clustering of tumor microenvironment (TME) cell populations in the Cancer Genome Atlas (TCGA) cohort.  

- **a** Consensus matrix heatmap from K-means consensus clustering identified 3 different clusters—namely, C1, C2, and C3.  
- **b** Kaplan-Meier curves showed the overall survival rates for patients with hepatocellular carcinoma (HCC) in TCGA and Liver Cancer-RIKEN,JP (LIRI_JP) cohorts. Patients were divided by different TME subtypes. Violin plot of the absolute abundance of TME cell populations distinguished by different TME subtypes in the TCGA cohort and LIRI_JP cohort.  
- **c** and **d** Immune and stromal scores were determined using ESTIMATION for patients of 3 TME subtypes in the TCGA cohort. Medians, interquartile ranges, and minimum/maximum are shown in boxplots, and the differences between each group were compared by using the Mann–Whitney U test.

Abbreviation: ns, no significance. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. A log-rank test was used to determine the statistical significance of differences.
To explore the functional implications of these signature genes of different subtypes, KEGG pathway enrichment analysis was performed. The results of KEGG enrichment analysis revealed that the signature genes of C1, C2, and C3 were primarily enriched in five KEGG pathways (for C1, extracellular matrix–receptor interaction, the Wnt/PI3K/Akt signaling pathway, focal adhesion, and amoebiasis), 23 pathways (the top five for C2 were primary immunodeficiency, cytokine–cytokine receptor interaction, intestinal immune network for immunoglobulin A production, hematopoietic cell lineage, and cell-adhesion molecules), and 26 pathways (the top five for C3 are neuroactive ligand-receptor interaction, protein digestion and absorption, GABAergic synapse, and breast cancer), respectively (Table S1). The results are visualized in bubble plots (Fig. 4e). Moreover, we investigated the correlation of immune cells with mutations of TP53 and CTNNB1 and found that the distribution of immune cells differed significantly between the low-TMB and high-TMB groups (Fig. 3b, c).

Fig. 3 Clinical characteristics of different TME subtypes. **a** Heatmap describing the absolute abundance of tumor microenvironment (TME) cell populations in TME subtypes C1, C2, and C3. Group information; mutation status of TP53, CTNNB1, and FGF19; vascular invasion; American Joint Committee on Cancer stage; histological grade; and gender are shown as patient annotations. **b** Boxplot of the tumor mutation burden in three TME subtypes. The differences were compared by using the Mann–Whitney U test. **c** Violin plot of the absolute abundance of TME cell populations between the low-TMB and high-TMB groups distinguished by median value. The differences between each subtype were compared using Student’s t-test.
CTNNB1, which are two representative genes in liver cancer. It was demonstrated that two types of immune cells (fibroblasts and cells of monocytic lineage) have different relative amounts of TME cell populations when comparing between TP53-mutant and TP53 wild-type HCC cells. Analogously, eight immune cells also showed different relative amounts of TME cell populations between CTNBB1-mutant and CTNBB1 wild-type HCC cells. Furthermore, CTNNB1 wild-type HCC cells, compared to CTNBB1-mutant HCC cells, had greater relative amounts of TME cell populations across all eight immune cells (Fig. S2).

Identification of hub genes of 3 TME subtypes
The signature genes in the three TME subtypes were analyzed and identified by gene co-expression network analysis to determine the core genes for C1, C2, and C3. After excluding the gene pairs with correlation coefficients of < 0.7 in C2 and C3 and those with coefficients of < 0.3 in C1, 45 signature genes for C1, 57 signature genes for C2, and 40 signature genes for C3 were included to construct the co-expression network. Co-expression networks were constructed using Cytoscape (Fig. 5a). Genes were represented by nodes in the network, and interactions between genes were represented by edges.

Network degrees, which describe the number of genes that regulate each other, represented the size and the color shades of the cycle node. Each node in the network represented the number of enriched genes in the gene set, and different colors and sizes represented different degrees of closeness, with the most central genes in the network having the largest degree values. Subsequently, the cytoHubba plugin in Cytoscape was applied to identify hub genes among signature genes of C1 (the top three genes were PNCK, SOHLH1, and CT45A1), C2 (the top three genes were SLAMF7, CD27, and A T cell immunoreceptor with immunoglobulin and ITIM domains [TIGIT]), and C3 (the top three genes were MOXD1, LUM, and CCDC80), respectively. Also, lollipop plots were used to visualize the correlation between hub gene expression and total number of TME cells. The results show that there was a weak correlation between hub genes of C1 and all TME cell populations (correlation coefficient < 0.3), hub genes correlate strongly with C2 and the eight immune cell populations (correlation coefficient > 0.5), and a strong correlation exists between hub genes for C3 and the two stromal cell populations (correlation coefficient > 0.5) (Fig. 5b). This result was reproduced in the LIRI_JP cohort (Fig. 5c).

Prognostic value of hub genes
The prognosis of the four hub genes was further analyzed and is shown in Fig. S2. The results of survival

Table 1  Clinical characteristics of patients of different subtypes in the Cancer Genome Atlas cohort

| Variable     | C1   | C2   | C3   | p value |
|--------------|------|------|------|---------|
| Gender       |      |      |      |         |
| Female       | 47   | 19   | 51   | 0.561   |
| Male         | 82   | 39   | 116  |         |
| Age (years)  |      |      |      | 0.371   |
| < 60         | 56   | 31   | 77   |         |
| ≥ 60         | 73   | 26   | 90   |         |
| Grade        |      |      |      | 0.330   |
| I            | 17   | 9    | 25   |         |
| II           | 69   | 25   | 75   |         |
| III          | 39   | 20   | 58   |         |
| IV           | 1    | 3    | 8    |         |
| AJCC stage   |      |      |      | 0.896   |
| I            | 55   | 29   | 82   |         |
| II           | 31   | 14   | 36   |         |
| III          | 27   | 13   | 42   |         |
| IV           | 3    | 0    | 2    |         |
| Vascular invasion |    |      |      | 0.752   |
| None         | 71   | 32   | 93   |         |
| Micro        | 33   | 11   | 45   |         |
| Macro        | 6    | 3    | 5    |         |
| TP53         |      |      |      | 0.447   |
| Mutant       | 35   | 21   | 52   |         |
| Wild-type    | 94   | 37   | 115  |         |
| CTNNB1       |      |      | <0.001|
| Mutant       | 17   | 14   | 63   |         |
| Wild-type    | 112  | 44   | 104  |         |
| FGF19        |      |      | <0.001|
| Amplification| 5    | 12   | 7    |         |
| Wild-type    | 124  | 46   | 160  |         |

Table 2  Clinical characteristics of patients of different subtypes in the LIRI_JP cohort

| Variable     | C1   | C2   | C3   | p value |
|--------------|------|------|------|---------|
| Gender       |      |      |      |         |
| Female       | 24   | 11   | 14   | 0.208   |
| Male         | 63   | 25   | 65   |         |
| Age (years)  |      |      |      | 0.163   |
| < 60         | 22   | 6    | 11   |         |
| ≥ 60         | 65   | 30   | 68   |         |
| Grade        |      |      |      | 0.279   |
| I            | 16   | 5    | 9    |         |
| II           | 41   | 19   | 37   |         |
| III          | 27   | 7    | 26   |         |
| IV           | 3    | 5    | 7    |         |
Fig. 4  Identification of signature genes of three tumor microenvironment (TME) subtypes and functional annotation. a Volcano plots of the differentially expressed genes between C1 and C2, between C2 and C3, and between C1 and C3, respectively. b Diagram depicting the process of identification of the signature genes of three TME subtypes. c Venn diagram showing the signature genes of three TME subtypes with no duplication. d Three different subtypes of signature genes shown as a heatmap. Abbreviation: FPKM, fragments per kilobase of exon per million reads mapped. e Bubble plots of Kyoto Encyclopedia of Genes and Genomes pathway analysis of TME subtypes’ signature genes

(See figure on next page.)

Fig. 5  Identification of hub genes from signature genes of different subtypes. a Co-expression network of signature genes of different subtypes. The correlation between genes (expression values were log-transformed and verified to be normally distributed) were estimated by Pearson’s correlation analysis. Genes are represented by cycle nodes, whose size represents how many genes are connected to the node, and in gene networks, edges represent interactions between genes. Lollipop plots show the correlation between the expression levels of hub genes and the absolute abundance of TME cell populations in the Cancer Genome Atlas cohort b and Liver Cancer-RIKEN, JP cohort c. The size of nodes represents the power of the interrelation between genes and TME cell populations, and the shade of nodes represents the significance of the correlation
Fig. 5 (See legend on previous page.)
analysis indicated that all three hub genes of C1 (PNCK: HR = 1.84, 95% CI = 1.29–2.64, \( p = 0.001 \); SOHLH1: HR = 1.71, 95% CI = 1.18–2.48, \( p = 0.002 \); and CT45A1: HR = 2.36, 95% CI = 1.27–4.40, \( p < 0.001 \)), two of the three hub genes for C2 (SLAMF7: HR = 0.63, 95% CI = 0.42–0.94, \( p = 0.041 \) and CD27: HR = 0.62, 95% CI = 0.42–0.90, \( p = 0.019 \)), and one of the three hub genes for C3 (MOXD1: HR = 1.61, 95% CI = 1.09–2.39, \( p = 0.032 \)) had significant prognostic value.

Discussion

Based on this analysis, three subtypes of immune microenvironment were identified for HCC (C1, C2, and C3). With the highest absolute abundance of TME cell populations and the longest survival time, the potential prognostic value of the PD-1 programmed death ligand 1 (PD-L1) blockade for the C2 patients is worth further investigation. The present analysis showed that patients in C2 had the best prognosis with the largest number of TME cells. This result is in accordance with that of another study on HCC classification, which showed that the patients with the immune-high subtype of TME had a significantly better prognosis [3]. The efficacy of immune checkpoint inhibitors (ICIs) is associated with the level of immune infiltration. Greater infiltration of cytotoxic T-cells is directly related to a better clinical response [4]. It has been demonstrated in previous studies [5–7] that the immune-high subtype correlates with PD-L1 expression in tumor and immune cells (mainly macrophages) and PD1 positivity in CD8+ T cells. In clinical trials for HCC, only CTLA-4 and PD-1/PD-L1 inhibitors have been tested to date. In the body, by interacting with PD-1 and PD-L1 on antigen-presenting cells, PD-L1 maintains self-tolerance and inhibits T cell activation [8, 9]. ICIs boost the immune response by depleting the tumor’s cytotoxic T-cells [10]. Hence, it is valuable to study the potential efficacy of ICIs in HCC patients in C2 further.

CTNNB1 and TP53 are two major mutation genes in HCC [11]. Previous studies [12] have shown that HCC with a CTNNB1 mutation is generally well differentiated. As a therapeutic biomarker, therapies targeting CTNNB1 in HCC have reached promising outcomes [13]. In this analysis, the lowest infiltration of immune cells together with the highest mutations of CTNNB1 may indicate a lack of response to ICIs. Simultaneously, in this current study, though the TMB in C3 was highest among the three subtypes, this result not may indicate a better response to ICIs [14]. Bagaev et al. [15] previously showed that TMB predicts the immunotherapy effect unsatisfactorily in a pan-cancer analysis (area under the receiver operating characteristic curve [AUC] for TME = 0.82, AUC of TMB = 0.56). An analysis based on TCGA data also found that a higher TMB in HCC is associated with a poorer survival prognosis and negative immune checkpoint activity; the infiltration level of immune cells was higher in the low-TMB group than the high-TMB group, and the heavy mutation load may inhibit immune cell infiltration in HCC. Consistent results were also observed in the C3 group of this study, i.e., a high TMB (Fig. 3b) and a low level of immune cell infiltration (Figs. 2c and 2d). For further verification, we divided patients into two groups according to their level of TMB, and the results showed that the high-TMB group had lower immune cell infiltration (Fig. 3c). This result may contradict the universal notion that high TMB might yield numerous neo-antigens that will activate the anti-tumor immune response, therefore more experiments are necessary.

FGF19 amplification was found in C2 patients in this analysis, which is reported to be overexpressed in a subtype of HCC patients [16]. FGF19 signals through its receptor, fibroblast growth factor receptor 4 (FGFR4), can induce hepatocyte proliferation as well as glycogen synthesis. FGFR4 inhibitors, including BLU-554 and FGR401, have been studied as potential treatments for HCC patients that act by interfering with FGF19–FGFR4 signaling and have achieved promising results in phase I and II clinical trials [17]. Hence, further studies in this regard are desirable, as are clinical trials designed to ascertain the therapeutic value of combining immunotherapy and molecular-targeted therapy for C2 patients.

A more thorough understanding of the complex mechanism of TME may stimulate further investigation into combination therapies [18]. The top three hub genes (PNCK, SOHLH1, and CT45A1) identified in C1 correlated with patient prognosis. Human breast cancers overexpress PNCK compared to benign breast tissues as PNCK plays an important role in mammary development [19]. Furthermore, CT45A1 overexpression has been confirmed in various cancers with a weak tumorigenic effect [20]. Studies have shown that [21, 22] overexpression of CT45A1 in breast cancer cells significantly upregulates several oncogenic and metastatic genes, indicating that CT45A1 may be a promising biomarker for targeted tumor therapy [21].

The top three hub genes (SLAMF7, CD27, and TIGIT) identified in C2 have great potential to be biomarkers of immunotherapy. SLAMF7 has been regarded previously as a target for immunotherapy in multiple myeloma [23]. CD27 belongs to the TNF (tumor necrosis factor) receptor superfamily and it is expressed exclusively on lymphocytes, and CD27 signaling involving its ligand, CD70, promotes T-cell expansion, survival, and differentiation as well as B-cell and NK cell activation [24]. Varilumab, a CD27 agonist, has already been tested in multiple early-phase clinical trials, either alone or in combination with
antiPD-1 therapy. Mouse models treated with CD27 ago-
nists have also revealed these drugs’ preventive role in
tumor formation or progression [25]. For cancer immu-
notherapy, T-cell immunoglobin and TIGIT, as well as
CTLA-4, PD-1, T-cell immunoglobin 3 and lympho-
cyte activation gene 3, are the most commonly targeted
checkpoints [26]. In melanoma and colon cancer mouse
models, anti-TIGIT therapy results in tumor regression
and improved survival. Although the development of
TIGIT inhibitors is still in its early phase, several clin-
ical trials of TIGIT inhibitors are now ongoing [27]. The
synergistic effect of combination therapy involving PD-1
inhibitors and TIGIT inhibitors in mouse models sug-
gests that it may be possible to treat lung cancer patients
with upregulation of both PD-1 and TIGIT [28]. In sum-
mary, molecular-targeted therapy alone, immunotherapy
alone, or a combination of both may be a potential ther-
apeutic strategy for C2 HCC patients.

Among the top three hub genes detected in C3, the
expression of LUM was found to be regulated during liver
fibrogenesis. Though LUM is considered to be essential
for hepatic fibrosis, its function in hepatocarcinogenesis
has not yet been determined [29, 30]. The signature genes
of C1 were primarily enriched in five KEGG pathways
(extracellular matrix–receptor interaction, Wnt signaling
pathway, Focal adhesion, PI3K–Akt signaling pathway,
and amoebiasis). Thus, the therapies targeting molecules
in these signaling pathways may be interesting items to
explore further in the future.

In general, this study classified the TME of HCC
according to varied immune compositions, gene expres-
sion levels, and prognoses. On the one hand, it is hoped
that this study can develop a deeper understanding of
HCC tumor biology and therapeutic response. On the
other hand, the new patient-stratification method pro-
posed in this study is expected to provide some guidance
for the management of HCC patients in the future and
may contribute to precision medicine;however, we still
recognize the limitations of our study. To reduce the bias,
we analyzed and constructed an immun-stratification
model based on the TCGA dataset and validated it in the
LIRI_JP dataset. However, all data in this study originate
from online datasets, so further experimental validation
and patient tumor sample validation are necessary.

Conclusion
Immunotherapies and/or FGFR4 inhibitors for HCC
patients in C2, CTNNB1-targeted therapies for those in
C3, and therapies targeting the molecules Wnt–β-catenin
pathway and/or the PI3K pathway for those in C1, may
be potential therapeutic strategies that warrant further
investigation.

Abbreviations
CTLA-4: Cytotoxic T-lymphocyte-associated protein-4; CTNNB1: Catenin
(cadherin-associated protein), β1; DEGs: Differentially expressed genes; FGFR19:
Fibroblast growth factor-19; FGFR4: Fibroblast growth factor receptor 4; ESTI-
MATE: Estimation of Stromal and Immune cells in Malignant Tumour Tissues
Using Expression Data; AJCC: American Joint Committee on Cancer; HCC:
hepatocellular carcinoma; HR: hazard ratio; IC: Immune checkpoint inhibitor;
KEGG: Kyoto Encyclopedia of Genes and Genomes; MST: Median survival time;
PD-1: Programmed cell death protein 1; PD-L1: Programmed death ligand 1;
TCGA: The Cancer Genome Atlas; TMB: Tumor mutation burden; TME: Tumor
microenvironment.

Supplementary Information
The online version contains supplementary material available at https://doi.
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Additional file 1: Table S1. Results of the Kyoto Encyclopedia of Genes
and Genomes enrichment analysis

Additional file 2: Fig. S1. Identification of optimal k of the Cancer
Genome Atlas. Fig. S2. Correlation of the immune cells with TP53 and
CTNNB1. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001

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Authors’ contributions
Luo Z.L. and Chen F.S. designed the study. Zhang L.M. and Yang C. conducted
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revised the manuscript. Other authors collected and processed the data. All
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Availability of data and materials
The data that support the findings of this study are openly available in the
Cancer Genome Atlas database (https://cancergenome.nih.gov) and the Inter-
national Cancer Genome Consortium portal database (https://dcc.icgc.org/
projects/LIRI_JP). The data that support the findings of this study are openly
available in the cBioPortal website (http://www.cbioportal.org/index.do). The
data that support the findings of this study are openly available in the Enrichr
database (http://amp.pharm.mssm.edu/Enrichr/).

Declarations
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Competing interests
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