Subpopulations of cancer-associated fibroblasts link the prognosis and metabolic features of pancreatic ductal adenocarcinoma

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**Background:** Cancer-associated fibroblasts (CAFs) are a vital constituent of the tumor microenvironment (TME) and have several activities, but the effect of CAF heterogeneity on the molecular features and clinical outcomes of pancreatic ductal adenocarcinoma (PDAC) remains unknown.

**Methods:** An algorithm “scFrac” based on single-cell sequencing data from the Gene Expression Omnibus was introduced to emulate the enrichment of CAF subtypes in a TCGA-PDAC cohort and their prognostic influence, and confirmed by an external validation group (66 patients with PDAC) with multiplex immunohistochemistry staining. A comprehensive analysis including metabolic profile and transcription factor regulon activity was carried out among CAF subtypes.

**Results:** Three distinct CAF populations were confirmed: myofibroblast (myCAF), inflammatory CAF (iCAF), and antigen-presenting CAF (apCAF). These subtypes expressed distinct metabolic profiles and transcriptional regulon activity. KEGG pathway annotation demonstrated that complement and coagulation cascades, as well as cytokine-cytokine receptor interaction were dominant in iCAFs, and pathways related to focal adhesion, and ECM-receptor interaction showed dominance in myCAFs, while antigen processing and presentation were the top enriched pathways in apCAFs. iCAFs trended to glycolysis with CREB3L1, EGR2 and SOX4 activation, whereas myCAFs depend on the tricarboxylic acid cycle and its derivatives with NRF2, CEBPD and YBX1 activation. iCAF is a protective factor associated with an inflammatory phenotype, but myCAF is an important factor in the poor prognosis of PDAC.

**Conclusions:** We identified distinct molecular characteristics of 3 CAF subtypes in PDAC and plotted their metabolism profile. We introduced a novel algorithm, scFrac, for exploring how CAF subgroups dysregulate cancer biology, and also shed a new therapeutic light on targeting the CAF subtype in TME.

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Introduction

Pancreatic ductal adenocarcinoma (PDAC) is one of the most malignant tumors of digestive tract and the seventh leading cause of cancer death in all sexes, with a median survival of 6 months and a 5-year survival rate of <8% (1). Despite improvements in surgical techniques and therapy regimens, existing systemic medications for advanced PDAC patients give only transient relief, underscoring the need for novel therapeutic solutions.

PDAC is distinguished by extensive desmoplasia, which accounts for 90% of the total tumor volume and includes extracellular matrix (ECM), immune cells, vasculature, and cancer-associated fibroblasts (CAFs) (2,3). Of these, CAFs are the majority. CAFs may originate from bone marrow-derived mesenchymal stem cells attracted to the tumor, or may be transformed from adipocytes, pericytes, or endothelial cells, but are predominantly derived from tissue resident fibroblasts in the pancreas. CAFs have a wide range of functions due to their adaptability and plasticity. The alterations in extracellular matrix production and tumor mechanics that result in a large part from the action of CAFs have complex consequences for tumors. Increased tissue stiffness triggers pro-survival and pro-proliferation signaling in cancer cells (4). Increased mechanical stress could collapse blood vessels, leading to hypoxia, promoting more aggressive cancer phenotypes, and reducing drug delivery (5-8). Contrary to these tumor-facilitated principles, 2 studies indicated that removing stroma completely resulted in more aggressive tumors with worse overall survival (OS), highlighting the complexities of the stromal-tumor interaction (9,10).

This paradoxical phenomenon has provoked wide debate accompanied by the formulation of a hypothesis that subtypes of CAFs with distinct phenotypes exist in PDAC. To address this possibility, several groups have performed gene-expression profiling on genetically engineered PDAC mouse models or patient samples. Öhlund et al. identified 2 distinct CAF subpopulations, one with elevated α-SMA expression and instead secreted interleukin (IL)-6 and additional inflammatory mediators were termed inflammatory CAFs (iCAFs). Furthermore, CAFs were shown to be dynamic, with variable phenotypes depending on their spatial and biochemical niche within the PDAC microenvironment (11). A recent study based on single-cell RNA-seq (scRNA-seq) analysis also highlighted the existence of different CAF populations in PDAC. Beside the classical myCAFs and iCAFs, a third population named antigen-presenting CAFs (apCAFs) was found, expressing the MHC II complex and CD74 that were able to present antigen to T cells in vitro (12). The findings all suggest the existence of multiple subpopulations of CAFs with distinct transcriptional programs. However, the functions of these subgroups remain unclear, particularly the relationship between these subgroups and patient prognosis. In this study, we take advantage of all published human PDAC scRNA-seq samples and identified ~7,000 CAFs for robust cluster discovery, and introduced a novel algorithm, scFrac, allowing calculation of enrichment score of all subgroups of CAFs in bulk RNA-seq of PDAC, to determine the effect of the different CAF subtypes on patient survival based on the patient cohort of our center.

We present the following article in accordance with the REMARK reporting checklist (available at https://atm.amegroups.com/article/view/10.21037/atm-22-407/rc).

Methods

Clinical samples

An external validation group of 66 patients from the Shanghai General Hospital and Minhang Hospital between 2016 and 2021 were enrolled in this study. All procedures performed in this study involving human participants were in accordance with the Declaration of Helsinki (as revised in 2013). All clinical samples were collected with informed consent from the patients, and the research was approved by the ethics committees of Shanghai General Hospital, Shanghai Jiao Tong University School of Medicine (China).
Data collection

The FPKM-normalized mRNA expression profiles and clinical features of ~170 patients with PDAC were downloaded from The Cancer Genome Atlas (https://portal.gdc.cancer.gov/). For single-cell RNA-seq data, 24 human PDAC tumor samples were collected from the Genome Sequence Archive (GSA) with accession number CRA001160 [NGDC-GSA (cncb.ac.cn)]. WE also collected 4 mouse KPC tumor samples from the Gene Expression Omnibus with accession number GSE129455.

scRNA-seq data procession

We applied Seurat 4.0 for scRNA-seq data procession. We routinely excluded single cells with <1,000 or >5,000 UMIs. To remove batch effects among samples, we applied the “IntergrateData” method. The “FindAllMarkers” method was used to identify major cell types. Each cell type was recognized by marker genes using “CellMarker” database (http://biocc.hrbmu.edu.cn/CellMarker/).

Gene-set enrichment analysis

Single-sample gene-set enrichment analysis (ssGSEA) was performed to evaluate the enrichment of the gene set of interest in TCGA-PDAC samples using R package GSVA. Gene signature collections were downloaded from the Molecular Signatures Database (http://www.gsea-msigdb.org/gsea/msigdb).

Evaluation of metabolic activity at single-cell resolution

A newly introduced algorithm, “scMetabolism”, generally supports the quantification and visualization of metabolism at single-cell resolution. We applied the algorithm to decipher the metabolic activity of 78 metabolic-associated pathways in human PDAC CAFs.

Transcription factor (TF) analysis

In an attempt to understand the regulatory activity differences of TFs in the CAF subtypes, single-cell regulatory network inference and clustering (SCENIC) was used to score the regulon. The adj.P value < 0.05 was considered as significant. The activity of the top regulons in each subgroup was displayed by heatmap.

Relative enrichment fraction of CAF subtypes in TCGA-PDAC

To allow for the convenient estimation of cell type proportions discovered by scRNA-seq in bulk-sample RNA-seq data, we introduced an SVR-based machine learning algorithm, “scFrac” (https://github.com/shenxiaotianCNS/scFrac). Similar deconvolution method has been introduced to analyze human and mouse sequence data (13,14). To our knowledge, this is the first time the full scRNA data set has been used to generate the X matrix. In this algorithm, we valued and made full usage of each single cell in the scRNA-seq, by deciphering the mixed matrix to the true, real, equal single-cell level, and using the summary of cell fractions belonging to the same subtype as the final enrichment fraction. Generally, biological mixed samples are modeled as a system of linear equations: a single mixture matrix X with n genes can be viewed as the product of Y and F, where Y is the n*m single-cell matrix containing n genes in X and m cells identified by scRNA-Seq. F represent a vector of size m, referring to the relative cell proportions for each cell. The sum of m is referred to as M, which is considered the relative enrichment fraction of the cell type.

Survival analysis

R package “Survminer” was used to performed survival analysis. Gene set used in this manuscript was downloaded from http://www.gsea-msigdb.org/gsea/msigdb. These calculation steps are taken from publicly available resources on the web.

Statistical analysis

Using R, Pearson correlation coefficients were calculated; survival analysis and Kaplan-Meier curves were constructed using the “survival” R package, and log-rank tests were used to determine significance. All statistical P values were two-tailed, with a significance level of P<0.05 being considered statistically significant. R 3.6.3 software was used to process all data.

Multiplex immunofluorescence (IF)

The IF was accomplished by staining 4 μm-thick formalin-
fixed, paraffin-embedded whole-tissue sections sequentially with standard primary antibodies and then pairing with the TSA 7-color kit (D110071-50T, WiSee Bio), followed by DAPI staining. For instance, deparaffinized slides were incubated with antibody for 30 minutes and then treated for 10 minutes with an anti-rabbit/mouse horseradish peroxidase (HRP) secondary antibody (#A10011-60, Yuanxibio). The IF labeling was created for a rigorously enforced 10 minutes using TSA 620 in accordance with the manufacturer’s recommendations. Slides were washed in TBST buffer and then transferred to a warmed citrate solution (90 °C) for 15 minutes. Slides were chilled to room temperature in the same solution. The slides were rinsed with Tris buffer in between each stage. The procedure was repeated with each antibody/fluorescent dye. Following that, each slide was treated with two drops of DAPI, rinsed in distilled water, and cover-slipped manually. The slides were dried in the air, mounted with anti-fade mounting media, and photographed using the Aperio Versa 8 tissue imaging equipment (Leica). Indica Halo software was used to analyze the images. The antibodies used in this study were CXCL12 (ab9797), α-SMA (ab7817), S100A4 (ab155326), pan-CK (ab7753), and PDGFR-alpha (ab203491).

Results

Identification of 3 subtypes of CAFs in PDAC

We first explored the heterogeneity of CAFs in human and mice pancreatic cancer using 2 single-cell sequencing data collections (12,15). A total of 6,933 CAFs in human tumor tissue and 4,000 CAFs in a mouse KPC model were included. Similar to previous findings (16), in the murine KPC model, 3 subtypes of CAF were identified: iCAF, myCAF and apCAF (Figure 1A). Notably, Elyada et al. did not discover apCAFs in human PDAC using single-cell analysis, probably due to the restricted cell number (12). In a larger PDAC data collection with ~60,000 cells, we identified a subpopulation of CAFs with the feature of high CD74 and MHCII molecule expressions (Figure 1B), indicating the presence of apCAFs in human PDAC, and they were transcriptome distinct against other subtype of CAFs. The markers of the CAF subtypes between humans and mice showed high consistency (Figure 1C,1D). Specifically, IL6, CXCL12, and DPT were overexpressed in iCAFs; HPOX (NDUF4L2), ACTA2, and MYL9 were highly expressed in myCAFs; and CD74, HLA-DRB1 (H2-Ab1), and HLA-DRA were upregulated in apCAF. Taken together, these results suggested 3 subpopulations of CAFs existed in human PDAC and the mouse KPC model.

Molecular characterization of CAFs in PDAC

We established the gene signatures of each CAF subtype using their marker genes, which were chosen using a “1 vs. others” strategy. Overall, 204, 159 and 193 genes were used to construct the iCAF, myCAF and apCAF signatures, respectively. The most different expressed markers are shown in Figure 2A. KEGG pathway annotation demonstrated that complement and coagulation cascades, and cytokine-cytokine receptor interaction were dominant in iCAFs, and pathways related to focal adhesion, and ECM-receptor interaction showed dominance in myCAFs, while antigen processing and presentation were the top enriched pathways in apCAFs (Figure 2B-2D), which was consistent with the functional annotation of markers of CAF subtypes in the murine KPC model (12). We calculated the “ssGSEA enrichment score” of the signatures in the TCGA-PDAC cohort to score the enrichment of the 3 subtypes of CAFs in human pancreatic cancer, and calculated the Pearson correlation value between the enrichment score and “HALLMARK” gene sets to reveal their connection (Figure 2E). R value >0.7 or <−0.3 was considered as significant. Results indicated that all types of CAFs were inflammatory regulators. Some malignant-related “HALLMARKs”, such as epithelial-mesenchymal transition (EMT) or angiogenesis, were all positively correlated with CAFs. Oxidative phosphorylation (OXPHOS), however, negatively correlated with the iCAF and apCAF signatures. Hence, CAF subtypes exhibit specific genetic signatures and functional roles in PDAC.

Metabolic heterogeneity of CAFs

The results showed that metabolic-related hallmarks such as hypoxia, OXPHS and adipogenesis correlated with different subtypes of CAF, indicating that other than cytokine secreting, matrix regulating or antigen-presenting, some more significant biological differences, such as metabolism alterations, may be present in the CAF subtypes. We then applied “scMetabolism” to evaluate the CAF metabolic activity at a single-cell resolution and found the 3 subtypes of CAFs were highly different metabolically. iCAF showed the highest activity in carbohydrates, fatty
acid, and amino acid metabolism, with the highest score in 47 of 78 metabolic pathways. myCAF had lowest metabolic activity, with the highest score in 13 pathways. apCAF was moderate, with 18 pathways having the highest score (Figure 3A). Notably, iCAF had the highest score for glycolysis, while myCAF had the highest score for the tricarboxylic acid (TCA) cycle and TCA derivatives such as glycine and serine metabolism. myCAF also had the highest score for OXPHOS (Figure 3B). These results suggest metabolic reprogramming from OXPHOS to glycolysis during CAF reshaping, especially towards iCAF. As shown in Figure 3C, the iCAF signature showed the highest correlation with glycolysis, while myCAF and apCAF had much lower R values (R=0.43, 0.084 and 0.2, respectively). One of the main reasons for increased glycolysis is hypoxia in the TME. Further exploration proved the iCAF signature
also had the highest correlation with the hypoxia signature (0.81 vs. 0.56 and 0.63). Taken together, local hypoxia in PDAC may reshape fibroblasts towards iCAF subtype, with increased glycolysis.

Distinct transcriptional activity and gene regulatory networks of CAF subtypes

TFs and their downstream-regulated genes constitute a
complex and intermingled network of gene regulation that determines the cell-to-cell variability in cancer. We applied “SCENIC” analysis to infer the activity of each TF and its regulated genes (which is the regulon) in the CAF subtypes. SCENIC (Figure 4A) showed a similar clustering of CAFs as with “Seurat” (Figure 4B). We can assess the difference in regulator activity between CAF subtypes using the regulator activity scores of TFs (Figure 4C). By comparing the regulon specificity score (RSS), we examined the crucial regulon of each cell type and visualized the top 5 regulons (Figure 4D). For myCAF, NRF2, CEBPD and YBX1 were the top regulon. Notably, a study has revealed a connection between fibroblast CEBPD and post-chemotherapy angiogenesis (17), and YBX1 has been identified as a target to suppress the myofibroblast phenotype in oral submucous fibrosis (18). For iCAF, we identified CREB3L1, EGR2 and SOX4 as the top regulon, among which CREB3L1 is a well-known hypoxia-induced TF (19). Moreover, SOX4 has been found to be activated by hypoxia signaling in non-small-cell lung cancer cells (20). For apCAF, MAFB, SPL1 and REL were the top regulon. Collectively, these results indicated different TF activity and gene regulatory networks among the CAF subtypes in line with their specific functions.

Association of CAF subtypes and different survival prognosis

In order to assess the composition of CAF subtypes in the TCGA-PDAC cohort, we built an SVM-based deconvolution algorithm to analyze the cell component of a mixture using the Seurat object, Seurat-derived cell markers and FPKM-normalized RNA-seq data as input. We tested the algorithm using a pseudo pooled RNA-seq data comprising 1,000 cells withdrawn from 24 tumor tissue samples, which showed a high consistency with scRNA data (Figure 5A). For fibroblasts, the Pearson correlation R value...
Figure 4 Distinct transcriptional activity and gene regulatory networks of CAF subtypes. (A,B) SCENIC (A) and Seurat (B) dimension reduction algorithms reveal similar CAF subgroups; (C) heatmap showing the regulon activity of CAF subtypes; (D) regulon specificity score of the regulon activity of CAF subtypes. CAF, cancer-associated fibroblast. RSS, regulon specificity score. iCAF, inflammatory cancer-associated fibroblast; myCAF, myofibroblast cancer-associated fibroblast; apCAF, antigen-presenting CAF.

was 0.93 respectively (Figure 5B).

We assessed the prognostic difference between high or low abundance groups. Interestingly, the higher iCAF was related to a better prognosis, whereas higher myCAF was related to a worse one (log-rank P=0.003 and 0.021). apCAF seemed to have no prognostic impact (P=0.18) (Figure 5C). We then validated the prognostic impact of iCAF and myCAF in an independent cohort using multiplex immunohistochemistry (miHIC). The cohort included 66 patients with PDAC, as an external validation group, from
Figure 5  Different survival prognosis of CAF subtypes. (A) Actual (left) and predicted (right) cellular composition of 24 pancreatic cancer samples deconvoluted by scRNA-seq and an SVR-based algorithm. (B) Pearson correlation of predicted and actual fibroblast composition in 24 pancreatic cancers. (C) Kaplan-Meier curves showing the survival difference between inflammatory CAF (upper) and myofibroblastic CAF (middle) and antigen-presenting CAF (lower) high or low subgroups in a TCGA-PDAC cohort. (D) Images of multiplex immunohistochemistry of PDGFRα+/CXCL12+ inflammatory CAF and S100A4+/α-SMA+ myofibroblastic CAF (left), high iCAF enrichment (upper) and high myCAF enrichment (lower) in PDAC co-hort. Scale bars indicating 50 μm and 200 μm in lefthand/righthand panels (×20) and middle panel (×5). (E) Kaplan-Meier curves showing the survival difference between inflammatory CAF (left), and myofibroblastic CAF (right) high or low subgroup in the external validation cohort. CAF, cancer-associated fibroblast; PDAC, pancreatic ductal adenocarcinoma; iCAF, inflammatory cancer-associated fibroblast; myCAF, myofibroblast cancer-associated fibroblast.
the Shanghai General Hospital and Minhang Hospital between 2016 and 2021. Four widely recognized molecules were selected as biomarkers to distinguish iCAF from myCAF: PDGFRα and CXCL12 for iCAF, and S100A4 and a-SMA for myCAF. In addition, we used pan-CK as a marker for PDAC to identify cancerous and non-cancerous tissues (Figure 5D) (21). The mIHC results demonstrated that patients with a higher number of iCAF infiltrations had a longer OS with statistical difference (P<0.05), which was consistent with the results of our preceding analysis (Figure 5E). We also compared the clinical features, such as tumor size, TNM stage, CA125 and CA199 levels, between iCAF-high and iCAF-low, or myCAF-high and myCAF-low subgroups in the external validation group. We found that iCAF was related to a lower TNM stage (P=0.0367) (Tables 1,2). Collectively, we discovered and validated the diverse prognostic impact of iCAF and myCAF in PDAC patients.

**Association of higher abundance of iCAF with an inflammatory phenotype**

Preceding findings suggested that the iCAF signature in PDAC is associated with more active metabolism. We further investigated how iCAFs influence cancer biology in PDAC. PDAC samples were divided into 2 groups based on the abundance of iCAFs (low or high). PCA plot showed a difference between the 2 groups (Figure 6A). Gene analysis of the 2 groups was performed, showing the ~6,000 genes were differently expressed between groups. KEGG annotation of the upregulated genes showed that for the iCAF-high group, inflammatory pathways, such as the JAK-STAT and PI3K-AKT signaling pathways, or metabolic pathways, such as the Wnt and PPAR signal pathways, were enriched (Figure 6B). For the iCAF-low group, bile secretion and ECM-receptor interaction were enriched (Figure 6C). We evaluated the metabolic activity in the iCAF-high or low groups using a ssGSEA algorithm, which, in line with the previous finding (16), showed higher metabolic activity, including lipid metabolism and glucose metabolism, was enriched in the iCAF-high group (Figure 6D, left). Moreover, we compared the immune cell specific chemotaxis genes between groups, and annotated them by the immune cell they attracted. Interestingly, the iCAF-high group was more attractive to immune cells, such as leukocytes and monocytes (Figure 6D, right). The tumor inflammation signature (TIS) is a widely applied gene signature to evaluate inflammatory state based on antigen-presented cell abundance, T/natural killer cell abundance, interferon activity and T cell exhaustion (22). Higher TIS score is related to benefit from PD1 treatment. We found a significantly higher expression of TIS signature genes.

### Table 1 Baseline characteristic between iCAF high patient and iCAF low patient in PDAC from Shanghai General Hospital and Minhang Hospital

<table>
<thead>
<tr>
<th>Clinical features</th>
<th>iCAF-high, mean ± SD</th>
<th>iCAF-low, mean ± SD</th>
<th>P/χ²</th>
</tr>
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<tr>
<td>Age, years</td>
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<td>Male</td>
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<tr>
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<tr>
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<td>1b</td>
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</tr>
<tr>
<td>2a</td>
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<tr>
<td>2b</td>
<td>9</td>
<td>13</td>
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<td>CA125, U/mL</td>
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iCAFs, inflammatory cancer-associated fibroblasts; PDAC, pancreatic ductal adenocarcinoma; TNM, tumor node metastasis.
in the iCAF-high group compared with the iCAF-low group (Figure 6E). Taken together, these findings suggested that higher iCAF abundance in PDAC is related to more reprogramming of metabolism and higher inflammatory state, and may predict a better response to immune therapy.

**Discussion**

PDAC patients are often diagnosed at a late stage, when curative surgical approaches are not feasible. Rarely do patients benefit from chemotherapy, due to the mesenchymal-rich character of PDAC. The existence of heterogeneous and distinct subgroups of CAFs has been widely recognized and traditionally divided into myCAFs and iCAFs, but recently a new subgroup, apCAF with antigen-presenting function, has been reported, enriching our understanding of CAFs (12). Moreover, a study based on single-cell sequencing showed that CAFs were categorized as CD105⁺ or CD105⁻ CAFs; CD105⁺ pancreatic CAFs were permissive for tumor development in vivo, whereas CD105⁻ fibroblasts were strongly tumor suppressive, providing a novel perspective on CAF types (23).

Previously, it has been considered that abundant CAFs and deposited collagen affected the perfusion of blood vessels through compression, preventing drugs and immune cells from entering the tumor and limiting the therapeutic effect (16). However, elimination of CAFs and the ECM does not prolong patient survival, which reflects the complexity and plasticity of CAFs. In this study, we found 3 spatially and functionally distinct CAF populations in human and murine PDAC and also plot the metabolism profile of CAF, indicated a potential association between metabolic rewiring and CAF subgroups. We introduced a novel algorism, scFrac, allowing calculation of enrichment score of all subgroups of CAFs in bulk RNAseq of PDAC, providing convenience for exploring how CAF subgroups dysregulate cancer biology. Interestingly, higher iCAF related to a better prognosis, but higher myCAF related to a worse one. A recent study showed that myCAF composition was a poor prognostic factor in 9 cancer types, including PDAC (24). Another single-cell sequencing study based on melanoma, head and neck squamous cell carcinoma and lung cancer revealed iCAFs were potentially associated with a molecular microenvironment of enhanced immune cell activation, whereas myCAFs were associated with a molecular microenvironment that favored ECM remodeling (25). When we cannot obliterate CAFs, perhaps reshaping and altering their phenotype, that is, turning foes into pals, would be a promising endeavor. Thus, it is crucial to clarify their features and functions as well as the

<table>
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<tr>
<th>Clinical features</th>
<th>myCAF-high, mean ± SD</th>
<th>myCAF-low, mean ± SD</th>
<th>P/χ²</th>
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myCAF, myofibroblast cancer-associated fibroblast; PDAC, pancreatic ductal adenocarcinoma; TNM, tumor node metastasis.
principles of their mutual transformation.

Although there is mounting evidence of interconversion among distinct CAF subtypes, the mechanisms by which the CAF composition varies during cancer development and which CAF subtype is associated with prognosis and therapeutic sensitivity remain unknown. In comparison to normal pancreatic fibroblasts, human PDAC CAFs exhibited low levels of lipid storage genes such as fatty acid-binding proteins, but human and mouse CAFs showed high levels of the vitamin D receptor protein (26). In vitro treatment of human CAFs with the strong vitamin D analog calcipotriol resulted in the production of lipid droplets and a reduction in α-SMA expression in the majority of samples. CAFs isolated from murine PDAC may be shifted between the α-SMA-high and IL-6-producing states through TGF-β and IL-1 signaling, implying substantial fibroblast plasticity (27). Additionally, the fact that fibroblasts and the α-SMA promoter are sensitive to a variety of extracellular stimuli, including substrate stiffness, supports the notion that the α-SMA-high, ECM-producing-high state is reversible (28-32). Another research discovered that inhibiting the hedgehog pathway decreased myCAF and raised iCAF numbers, which coincides with a drop in cytotoxic T cells and an increase in regulatory T cells, associated with greater immunological suppression (33). Overall, these results suggest the potential for interconversion of the subtypes of CAFs. Our results revealed that CAFs were metabolically heterogenic, notably iCAF trending in glycolysis, while myCAF trended in TCA cycle and TCA derivatives such as glycine and serine metabolism. Targeting and reprogramming these metabolic pathways may provide novel therapeutic approaches for better clinical outcomes.

![Figure 6](image-url) Association of higher abundance of iCAFs with an inflammatory phenotype. (A) PCA dimension reduction plot showing the different clustering of iCAF-high and iCAF-low PDAC subgroups. (B,C) KEGG annotation of the differently expressed genes highly expressed in iCAF-high PDAC (B) or iCAF-low PDAC (C). (D) Volcano plots comparing the enrichment difference of metabolic pathways (left) or chemotaxis gene-expression differences (right) between iCAF-high or -low PDAC subgroups. (E) Tumor inflammatory gene signature expression differences between iCAF-high or -low subgroups. ns, not significant; *, P<0.05; **, P<0.01; ***, P<0.001; ****, P<0.0001. iCAFs, inflammatory cancer-associated fibroblasts; PDAC, pancreatic ductal adenocarcinoma. FPKM, Fragments Per Kilobase of exon model per Million mapped fragments.
Patient outcomes will be improved through multidisciplinary advancements in imaging, surgery, radiation, and systemic therapies. Although clinical progress has been modest, our understanding of the molecular biology of PDAC and the TME continues to expand and will eventually inform rational therapeutic approaches that will result in clinical benefit.

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Footnote

Reporting Checklist: The authors have completed the REMARK reporting checklist. Available at https://atm.amegroups.com/article/view/10.21037-atm-22-407/rc


Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://atm.amegroups.com/article/view/10.21037-atm-22-407/coif). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All procedures performed in this study involving human participants were in accordance with the Declaration of Helsinki (as revised in 2013). All clinical samples were collected with informed consent from the patients, and the research was approved by the ethics committees of Shanghai General Hospital, Shanghai Jiao Tong University School of Medicine (China) (No. 2021-130) and Minhang Hospital, Fudan University (China) (No. 2021-066-01K).

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