A deeper understanding of the tumor microenvironment in pancreatic cancer: the key to developing effective immunotherapies

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Dismal statistics in pancreatic ductal adenocarcinoma (PDAC)

The number of pancreatic cancer related deaths is projected to increase while many other cancers fall (1). Pancreatic cancer is the third leading cause of cancer related death and projected to overtake second by 2020 (1). Despite intense research efforts over many years, five-year survival after diagnosis with pancreatic cancer remains grim at around 8% (2). Only 20% of patients present with surgically resectable disease, and even for those, five-year survival is a mere 24.6% (3). For those with more advanced disease, median survival is just 4.2 months (3). Of the small minority of patients who present with resectable disease, standard of care includes surgical resection followed by a limited number of chemotherapeutic options which yield a 19-month median survival (3). Response to available chemotherapeutic regimens varies between patients and the reasons behind these varied responses are unclear. This leave clinicians with a strategy of administering a chemotherapy and abandoning it in lieu of another after disease progression. Regardless of therapeutic approach, and particularly in metastatic disease, the prognosis remains poor (4). There has been little improvement in these dismal statistics despite decades of research.

Immunotherapy in cancer treatment

Immunotherapy for cancer treatment has gained much interest in recent years. Outside the classic treatment with surgery, chemotherapy, and radiation therapy, immunotherapy offers another mechanism for fighting cancer. The National Cancer Institute (NCI) defines immunotherapy as “A type of therapy that uses substances to stimulate or suppress the immune system to help the body fight cancer, infection, and other diseases.” This can include monoclonal antibodies, vaccines, checkpoint inhibitors, and adoptive cell transfer. The past decade has seen fervent investigation into the relationship between the immune system and cancer. Targeting these pathways provides novel mechanisms for cancer therapies. The 2018 Nobel Prize in Physiology or Medicine was jointly awarded to James P. Allison and Tasuku Honjo for their efforts in checkpoint inhibition. The year of 2018 alone saw Food and Drug Administration (FDA) approval of immunotherapy in hepatocellular carcinoma, small cell lung cancer, colorectal cancer, lymphoma, and cervical cancer. Some of these efforts have yielded stunning outcomes with long-term survival in patients with previously poor prognoses. The results of a phase 1 trial on the use of recombinant poliovirus in the treatment of recurrent glioblastoma was...
the complex relationship between pancreatic cancer cells and the surrounding desmoplastic tumor microenvironment. The mechanisms behind the poor response of PDAC to current investigation are summarized in Table 1.

βig-h3 as a potential target for immunotherapy

In their recent paper in Gut, Goehrig et al. describe the impact of the stromal protein βig-h3 in pancreatic cancer (23). They make a compelling case for the importance of βig-h3 in immune tolerance to pancreatic cancer in a murine model and in culture. First, they demonstrate that βig-h3 is strongly expressed in Cre;KrasG12D;Ink4a/Arffl/fl (KIC) and pdx1-Cre;KrasG12D;Ip53R172H (KPC) mice compared to control, as well as human pancreatic cancer biopsies. βig-h3 is secreted primarily by the cancer-associated fibroblasts (CAF). The effects of βig-h3 within the tumor microenvironment highlight its potential as a target for future therapies. βig-h3 suppresses proliferation of antitumoral CD8+ T cells in vivo and in vitro, controls activation of macrophages and CD8+ T cells, and promotes M2 macrophage differentiation. Depletion of βig-h3 in vivo led to a decrease in neoplastic cells and reduced tumor volume. Further analysis of the tumors from Big-h3 depleted mice demonstrated more infiltrating CD8+ T cells with a less exhausted phenotype. The summation of these findings suggests that βig-h3 is a critical protein secreted by CAFs and involved in tumorigenesis as well as promoting an immunosuppressive tumor microenvironment through modulation of both the adaptive and innate immune systems.

βig-h3 represents an important target for future immunotherapies and the authors should be commended for their work. Blocking βig-h3 may help improve immune clearance of neoplastic cells in future therapies. A deeper understanding of the mechanisms through which this protein acts to promote an immunosuppressed tumor
microenvironment may identify other biologic targets of similar importance.

**Conclusions**

PDAC presents many unique challenges for development of effective immunotherapies when compared to other more immunogenic cancers. Investigators have described many mechanisms of immune evasion by PDAC and our understanding of this phenomenon continues to grow. Continued work to identify and attack these mechanisms of immune evasion and tumor progression may provide the breakthrough needed to help the immune system fight this deadly malignancy.

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None.

**Footnote**

*Conflicts of Interest:* The authors have no conflicts of interest to declare.

**References**


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**Table 1** Pathways under investigation in the tumor microenvironment of pancreatic ductal adenocarcinoma

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Mechanism</th>
<th>Citation</th>
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<tbody>
<tr>
<td>Angiotensin II</td>
<td>Protein kinase C dependent proliferation of cancer associated fibroblasts by SMAD7 induction</td>
<td>(14)</td>
</tr>
<tr>
<td>CCR2</td>
<td>Recruitment of immunosuppressive tumor associated macrophages to the tumor microenvironment</td>
<td>(15)</td>
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<tr>
<td>CD40</td>
<td>Promotes tumoricidal macrophage infiltration and stromal depletion</td>
<td>(16)</td>
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<tr>
<td>CDK4/6</td>
<td>Phosphorylation of retinoblastoma protein and entry into S phase</td>
<td>(17)</td>
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<tr>
<td>CXCR4</td>
<td>Binding of CXCL12 to CXCR4 promotes T cell suppression</td>
<td>(18)</td>
</tr>
<tr>
<td>FAK</td>
<td>Increased fibrosis and poor CD8+ T cell infiltration</td>
<td>(19)</td>
</tr>
<tr>
<td>Hh</td>
<td>Increased stromal desmoplasia via upregulation of the Gli family of transcription factors</td>
<td>(20)</td>
</tr>
<tr>
<td>JAK1/2</td>
<td>Upregulates inflammatory cytokines and STAT3 leading to disease progression</td>
<td>(21)</td>
</tr>
<tr>
<td>MyD88</td>
<td>CD4+ and CD8+ T cell suppression</td>
<td>(22)</td>
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CCR2, C-C chemokine receptor type 2; CD40, cluster of differentiation 40; CDK4/6, cyclin-dependent kinase 4/6; CXCR4, C-X-C chemokine receptor type 4; FAK, focal adhesion kinase; Hh, hedgehog; JAK1/2, Janus kinase 1/2; MyD88, myeloid differentiation primary response 88.

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