Pancreatic ductal adenocarcinomas (PDACs) are classically immunologically cold tumors that have failed to demonstrate a significant response to immunotherapeutic strategies. This feature is attributed to both the immunosuppressive tumor microenvironment (TME) and limited immune cell access due to the surrounding stromal barrier, a histological hallmark of PDACs. In this issue of the JCI, Sharma et al. employ a broad glutamine antagonist, 6-diazo-5-oxo-l-norleucine (DON), to target a metabolic program that underlies both PDAC growth and hyaluronan production. Their findings describe an approach to converting the PDAC TME into a hot TME, thereby empowering immunotherapeutic strategies such as anti-PD1 therapy.
Disrupting a converging metabolic target turns up the immunologic-heat in pancreatic tumors

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The tumor microenvironment of pancreatic ductal adenocarcinomas

Immunotherapy has changed the treatment paradigm for a number of previously deadly cancers, including metastatic non–small cell lung cancers and melanomas. However, current immunotherapies have had little success against other cancers, including pancreatic ductal adenocarcinomas (PDACs). PDACs are stereotypically known to be “immunologically cold” tumors, in which the presence of immune cells and their activity are limited by the immune-suppressive tumor microenvironment (TME). The relative lack of effector T cells in the pancreatic TME is largely attributed to the presence of a barrier, termed stromal desmoplasia, surrounding the PDAC cells. Importantly, this histopathological hallmark of PDACs can also serve as a potential therapeutic target toward enhancing both drug delivery and immune responses.

Hyaluronan, a nonsulphated glycosaminoglycan in the extracellular matrix, is secreted by PDAC cells (1), and its high deposition within the pancreatic TME is associated with poor prognosis (2). Efforts to target hyaluronan directly, however, have been met with mixed responses. Mouse model studies have shown that enzymatic degradation of hyaluronan by administering a pegylated human recombinant PH20 hyaluronidase (PEGPH20) improves intratumoral vascularity and, subsequently, drug delivery and efficacy (3, 4). While one early phase clinical trial showed that adding PEGPH20 to gemcitabine and nab-paclitaxel (one standard of care regimen for PDAC) improves responses to therapy (5), another showed that combining PEGPH20 with FOLFIRI-NOX chemotherapy (the other standard of care for PDAC) is less effective than chemotherapy alone (6). These observations indicate the complexity of targeting the stroma, consistent with its known role in restraining PDAC growth (7, 8).

In this issue of the JCI, Sharma and colleagues used mouse models of chronic pancreatitis (caerulein induced) and PDAC (Kras-Tp53 driven, KPC mice), and human pancreatic cancer samples from a deidentified tissue microarray and The Cancer Genome Atlas, to confirm that markers relevant to the HBP were highly represented in PDAC models. The authors then inhibited the HBP rate-limiting enzyme glutamine fructose-6 phosphate (F6P) amidotransferase 1 (GFAT1) with siRNA in human PDAC cell lines. Notably, self-renewal gene expression and clonogenicity decreased. Furthermore, treatment of KPCs with DON led to reduced viability, invasiveness, and migration potential. Consistent with these in vitro findings, in vivo treatment of xenograft and KPC-fibroblast coimplanted orthotopic syngeneic mouse models with DON led to significantly decreased tumor growth, Ki67 positivity, and metastatic spread. The authors then observed that the antitumor effects of DON were indeed associated with features of active extracellular matrix remodeling, including decreased hyaluronan and collagen I, changes in metalloproteases, lower IL-27 in KPC mice, lower IL-6, and higher IFN-γ production by fibroblasts. Thus, the authors rigorously

Therapeutic effects of glutamine antagonism in PDAC models

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demonstrated the therapeutic utility of DON through multiple models (9).

Importantly, Sharma et al. investigated the effects of DON on the immune TME. They employed both single-stain immunohistochemistry and flow cytometry to demonstrate DON-associated tumor infiltrating CD68+ monocytes and CD8+ T cells. They also evaluated survival and tumor volume in the orthotopic mouse model using both wild-type and CD8-knockout mice and showed that the antitumor effects conferred by DON are dependent on CD8+ T cells. However, caution should be taken in interpreting DON’s effects on the immune TME, especially within the CD68+ population, due to the lack of subtyping of these cells and analysis of their functional states in this study. The authors then further interrogated the immune-activating effects of DON by testing the in vivo response of KPC tumors to anti-PD1 therapy. It is well known that both human and KPC PDACs fail to respond to anti-PD1 therapy alone. Surprisingly, this study showed that DON induces susceptibility to anti-PD1 therapy, generating superior efficacy in combination over monotherapy (9).

Translational implications and future considerations
This study by Sharma et al. provides a number of new insights into the PDAC TME. First, it highlights a metabolic program that can be pharmacologically inhibited to disrupt tumor proliferation, metastasis, and stromal composition. Second, it implies that glutamine antagonism can in fact induce a marked change in the antitumor immunologic response, effectively converting the “cold” into a “hot” TME, and can elicit significant responses to anti-PD1 therapy (9). The immunologic phenomenon is particularly intriguing, since leveraging the immune system to treat PDACs remains a difficult task. Similar to what Sharma et al. observed, another recent study has shown that breaking down the stromal barrier with PEGPH20 is associated with increased memory T cell infiltration and improved survival when combined with a GM-CSF–secreting pancreatic tumor vaccine (10). Thus, these observations provide important proof of concept and excite possible translational efforts.

To realize the translatability of the authors’ findings, there are at least two major caveats to address: targeting a physiologically relevant pathway and developing a viable drug. As studies recognize, the tumor-restraining function of the stroma is an important therapeutic target. Thus, targeting a signaling pathway responsible for both carcinogenesis and stromal barrier functions is theoretically attractive. In fact, this approach has previously been investigated in the form of Hedgehog inhibitors. However, despite the well-established role of aberrantly activated Hedgehog signaling in pancreatic carcinogenesis (11) and stromal development (12) and despite the successful preclinical assessment of Hedgehog inhibition approaches (13), clinical trials have failed to demonstrate benefit (14). This example suggests that predicting the clinical outcome is difficult when perturbing a pathway involved in carcinogenesis and stromal barrier functions as well as in stromal development for normal physiologic benefit. Another critical question is whether a clinically applicable glutamine antagonist could be developed. While the authors discuss DON as the candidate drug for future trials, prior trials with DON have been challenged by significant toxicities (15); the rate of mucositis, which is a highly morbid adverse effect, was greater than 80% in multiple trials when used at low daily dosing with which efficacy was observed. Thus, alternative formulations or candidate drugs must be explored. For example, the use of DON prodrugs with more preferable tissue distribution and therapeutic index has recently been proposed (15).

In summary, this study by Sharma et al. demonstrates the therapeutic effect of DON on suppressing PDAC growth that occurs through the attraction of CD8+ T cells into the TME (9). Concurrent anti-PD1 therapy may further activate these T cells. Thus, these results suggest the unique opportunity to convert PDACs from a cold to a hot immunological state. Finally, future drug development and translational efforts to target this “integrated metabolic node” will be worthwhile.

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