VentX expression in tumor-associated macrophages promotes phagocytosis and immunity against pancreatic cancers

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Introduction
Pancreatic ductal adenocarcinoma (PDA) is a lethal malignancy that has no effective treatment. The tumor microenvironment (TME) of PDA employs a multitude of immune derangement strategies to protect PDA from immune elimination. Tumor-associated macrophages (TAMs) have been implicated in the pathogenesis of immune suppression of the PDA TME; however, its underlying mechanisms remained largely unknown. Using primary patient samples, our studies showed that, in comparison with macrophages isolated from normal pancreatic tissues, the phagocytosis activity of the PDA TAMs was significantly reduced. We found that the expression of homeobox protein VentX, a master regulator of macrophage plasticity, was significantly decreased in the PDA TAMs. We demonstrated that VentX was required for phagocytosis and that restoration of VentX expression in PDA TAMs promoted phagocytosis through the regulation of the signaling cascades involved in the process. Using an ex vivo culture model of primary human PDA, we showed that VentX-modulated TAMs transformed the PDA TME from a protumor milieu to an antitumor microenvironment by rectifying differentiation, proliferation, and activation of PDA-infiltrating immune cells. Using NSG-PDX models of primary human PDAs, we showed that VentX-modulated TAMs exerted strong inhibition on PDA tumorigenesis in vivo. Taken together, our data revealed a central mechanism underlying immune evasion of PDA and a potential novel venue to improve PDA prognosis.
and immune surveillance, is significantly impaired in PDA TAMs. To define the mechanism underling the TAM inertia, we found that the expression of homeobox protein VentX, a master regulator of macrophage plasticity (14), is drastically reduced in TAMs of all cases of PDA tested in this study. We show that restoration of VentX expression in PDA TAMs recovered its phagocytosis function. Mechanistically, we show that VentX promoted the expression of cell surface receptors, such as TLRs, and controls intracellular signaling cascades involved in phagocytosis. Consistent with its role in polarizing TAMs from the protumor M2-like phenotype to the antitumor M1-like phenotype, we demonstrated that VentX-modulated TAMs transform the PDA TME from immune suppressive to immune activating by alternating the molecular and cellular composition at the PDA TME. Using a preclinical model of primary human PDA, we show that VentX-modulated TAMs exerted strong inhibition of PDA tumorigenesis in vivo. Taken together, our studies suggests a key regulatory role of VentX in governing the fundamental function of macrophages and a central function of VentX-TAMs in modulating immunity at the PDA TME. Our study suggests a potentially novel venue for PDA immunotherapy, functioning in part by promoting phagocytosis and rectifying immune derangement at the PDA TME.

Results

Characterization of the PDA TME. To better understand the mechanism underlying immune suppression at the PDA TME, we characterized the composition and function of immune cells at the PDA TME. TAMs and T cells were isolated from fresh PDA and adjacent normal tissues, which were verified by a board-certified pathologist. Consistent with prior immunohistochemistry findings (15), in comparison with adjacent normal tissues, the PDA TME contained significantly increased numbers of macrophages, which predominately displayed a characteristic M2-like phenotype (Figure 1 and Supplemental Figure 1). Consistent with their immune-suppressive function in the PDA TME, the PDA TAMs expressed increased levels of immune-suppressive cytokines, such as IL-4, IL-10, IL-13, and TGF-β, but decreased levels of proinflammatory cytokines, IL-1β, IL-8, IL-12B, and TNF-α (Figure 2A). In addition, the PDA TAMs expressed increased levels of immune checkpoint inhibitors, such as PD-L1 and Vista (Figure 2B). Similar to the TME of colorectal cancer (CRC), the prominent presence of M2-like TAMs in the PDA TME was accompanied by a significantly increased number of immune-suppressive Tregs. In contrast, there were very limited numbers of cytotoxic effector CD8 T cells in the PDA TME (Supplemental Figure 2; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.137088DS1).

VentX expression is decreased in PDA TAMs. As the key executor of both innate and adaptive immunity, TAMs have been implicated in immune suppression at the TME (16). Based on our recent findings on homeobox protein VentX, a master regulator of macrophage plasticity, and its implications in the pathogenesis of immune suppression at the TME (14, 17, 18), we further explored the potential involvement of VentX in the immunopathogenesis of PDA. We found that, similar to its involvement in CRC (18), the expression of VentX is significantly decreased in TAMs of all cases of PDA tested in this study (Figure 2, C and D).

Restoration of VentX expression in PDA TAMs promotes phagocytosis. Phagocytosis constitutes the essential effector function of macrophages (19). While implicated in antitumor immunity, the role and mechanisms of phagocytosis in the pathogenesis of PDA has remained poorly defined. Using primary clinical samples, we sought to determine whether the phagocytosis function of TAMs was compromised in the PDA TME. To achieve our goal, PDA TAMs and control macrophages were isolated from PDA and adjacent normal pancreatic tissues. Using an ex vivo phagocytosis assay of CFSE-labeled pancreatic cancer cells, we found that the phagocytic activities of PDA TAMs were significantly impaired in comparison with that of the macrophages isolated from adjacent normal pancreatic tissues (Figure 3A). Consistent with its functional plasticity, previous studies suggested that phagocytosis is regulated by M1 and M2 signals (20). As such, our findings that VentX was significantly downregulated in PDA TAMs prompted us to explore whether VentX regulates phagocytic function of PDA TAMs. Therefore, we transfected the primary PDA TAMs with plasmid encoding GFP-labeled VentX (GFP-VentX, a chimeric protein of GFP and VentX) or control GFP and found that ectopic expression of VentX led to 2.5- to 3-fold increases of PDA TAM phagocytosis (Figure 3B). The effects of VentX on PDA TAM phagocytosis can be readily visualized with phase-contrast and fluorescent microscopy examination of the coculture of PDA TAMs and CellTrance Yellow–labeled tumor cells (Figure 3, B and C). To rule out the possibility that VentX promotion of phagocytosis is cell-type specific, we tested the effects of VentX expression on PDA TAM phagocytosis of leukemia cells and found that VentX promoted phagocytosis indiscriminately (Supplemental Figure 3). In addition to its effects in promoting phagocytosis, when
A time-course observation of phagocytosis was performed, we found that, after the CFSE-labeled cells were engulfed during the first 24 hours, the CFSE dyes became diffusely distributed throughout the cell in 24–48 hours (Figure 3C). There was no formation of foamy cells or accumulation of vacuole of uncleared phagocytic material, as observed in M2 macrophages (21). Indeed, consistent with our prior findings in CRC-TAMs (18), we found that ectopic expression of VentX in PDA TAMs promotes the polarization of PDA TAMs from the M2-like phenotype to the M1-like phenotype (Supplemental Figure 4). We showed that VentX inhibited the cell surface expression of M2 marker CD163 but promoted the expression of M1 marker CD80 (Supplemental Figure 4A); VentX also promoted the expression and secretion of proinflammatory cytokines TNF-α, IL-1β, and IL-12 but inhibited the expression of immune-suppressive cytokines TGF-β, IL-4, and IL-10 (Supplemental Figure 4, C and D). Moreover, we found that VentX expression in PDA TAMs inhibited the expression of immune checkpoint blockade proteins PD-L1 and Vista and promoted the expression and secretion of effector molecules, such as nitrate (Supplemental Figure 4, B and E).

To explore the mechanisms underlying VentX promotion of phagocytosis, we performed a targeted screen of cell surface receptors and signaling pathways implicated in phagocytosis. We found that VentX promoted the expression of TLR2 and TLR9 and downstream signaling molecules MyD88 and MAP kinase p38 in PDA TAMs (Figure 4A). In comparison, ectopic expression of VentX in PDA TAMs exerted little effect on the expression of SIRPα, the cellular receptor for CD47, the “do-not-eat-me” signal implicated in cancer immune evasion (22) (Figure 4B). Instead, we found that VentX expression inhibits the phosphorylation of SHP-1 and SHP-2, the SIRPα downstream signaling molecules implicated in inhibition of phagocytosis. Moreover, we found that VentX expression promoted the phosphorylation of FAK kinase implicated in promoting phagocytosis (Figure 4C). Consistent with the essential requirement of VentX in the phagocytic function of PDA TAMs, we found that when the PDA TAMs were treated with morpholino-oligo against VentX, the stimulation effects of LPS were abolished (Figure 4D). Our findings suggested an essential role of VentX in regulating the balance of signaling cascades to promote phagocytosis.

**Figure 1. Characterization of pancreatic ductal adenocarcinoma.**
(A) Representative images of H&E and vimentin immunostaining of pancreatic ductal adenocarcinoma (PDA) and control adjacent normal pancreatic tissue. Scale bar: 200 μm. (B) FACS analysis of CD68+ macrophages in pancreatic cancers. CD68+ macrophages in adjacent normal pancreatic tissue and PDA were analyzed by a flow cytometer. Blue shading represents isotype control, and red shading represents CD68+ cells. (C) Accounting of CD68+ macrophages in pancreatic cancer and adjacent normal pancreatic tissues. n = 6. **P < 0.01, paired Student’s t test. (D) The percentage of surface expression of M2 markers (CD163 and CD206) and M1 surface markers (CD40 and CD80) on CD68+ macrophages of adjacent normal pancreatic tissues and PDA TAMs from the same patients. Data are shown as the mean ± SD of 4 independent experiments. *P < 0.05, **P < 0.01, 1-way ANOVA with multiple comparisons was performed.
Figure 2. Characterization of tumor-associated macrophages in pancreatic ductal adenocarcinoma. (A) The mRNA expression levels of indicated M1 and M2 cytokines in tumor-associated macrophages (TAMs) versus control macrophages, as determined by qRT-PCR. Results represent mean ± SD of 4 independent experiments. *P < 0.05, paired Student’s t test. (B) Cell surface expression of checkpoint inhibitor PD-L1 and VISTA mRNA in TAMs and control macrophages isolated from adjacent normal pancreatic tissues. The expression level of PD-L1 was determined by flow cytometer analysis, and representative graphs are shown. n = 4. The mRNA expression levels of VISTA were determined by qRT-PCR. Results represent mean ± SD of 8 independent experiments. *P < 0.01, paired Student’s t test. (C) Paired comparison of qRT-PCR measurements of VentX mRNA expression in macrophages isolated from normal control tissues and TAMs of 17 patients. The relative VentX mRNA expression levels in normal macrophages were arbitrarily designated as 1. **P < 0.01, TAMs vs. adjacent normal macrophages, paired Student’s t test. n = 17. (D) Western blot analysis of endogenous VentX protein levels in macrophages isolated from normal control tissues and TAMs of 3 patients. The numbers indicate the relative fold difference of VentX proteins, as determined by ImageJ (NIH) software scanning. N, normal control tissue; Tu, tumor.
VentX-TAMs transforms the PDA TME and promotes immunity against PDA. A multitude of immune derangements of the PDA TME has been implicated in PDA evasion of current immunotherapeutic modalities. Macrophages are key executors of both innate and adaptive immunity. Our findings that VentX promoted PDA TAM phagocytosis and polarized PDA TAMs from the M2-like phenotypes to the M1-like phenotypes prompted us to examine the effects of VentX-modulated PDA TAMs on the immune state of the PDA TME. Fresh isolated PDA TAMs were transfected with plasmid encoding GFP or GFP-VentX. The transfected TAMs were then cocultured with 1 μM CellTrance Yellow-labeled Panc-1 cells for 24 hours. The effects of the treatment were determined by phase-contrast and fluorescent microscopy (left) and flow cytometry analysis (right). Arrowheads indicated phagocytosed cancer cells. Scale Bar: 50 μM. The representative figure was shown. n = 3. (C) Phagocytosis and intracellular digestion of CellTrance Yellow-labeled Panc-1 cells. TAMs transfected with control GFP or GFP-VentX were incubated with CellTrance Yellow-labeled Panc-1 cells. Representative images of the phagocytosis were revealed by fluorescent microscopy after 24 hours (top) and 48 hours (bottom). Scale Bar: 20 μm. Quantification of phagocytic cells was performed by measuring green fluorescence intensity after phagocytosis. The graph represents the pixel intensity of fluorescence. Data presented are mean ± SD. n = 3. **P < 0.01. At least 10 cells were scanned for each time point.
of the M2-like TAMs and marked increases in the number of cytotoxic CD8 T cells (Figure 5D). To explore the mechanisms underlying the alternation of cellular composition at the PDA TME, we cocultured CD8 T cells with autologous PDA TAMs transfected with GFP-VentX or control GFP and found that VentX abolished the inhibitory effects of PDA TAMs on CD8 proliferation and activation (Supplemental Figure 5). Together with the effects of VentX on PDA TAM phenotypes and the expression of immune checkpoint inhibitors (Supplemental Figure 4), our data suggested that VentX-TAMs transform the immune status of the PDA TME by rectifying a multitude of molecular and cellular derangements at the PDA TME. As the M1/M2 ratio and the CD8/Treg ratio have been implicated in PDA prognosis (11), we sought to explore the therapeutic potential of VentX-TAMs on PDA. As VentX does not have a murine homolog (23), we transplanted a small piece of primary human PDA tissue subcutaneously at the dorsal lateral side of NSG mice to generate the NSG patient-derived xenograft (NSG-PDX) model of primary human PDA. Seven days after transplantation, the NSG-PDX model of primary human PDA (NSG-PDA) mice were tail vein injected with VentX-TAMs or control-TAMs. Consistent with the tumor chemokine-traction of macrophages (24, 25), we found that the injected TAMs accumulated in lymph nodes on the side of transplanted tumors (Figure 6A). In addition, we also observed the homing of the injected VentX-TAMs to the implanted PDA (Supplemental Figure 6B). The effects of VentX-TAMs on the growth of PDA in vivo was observed for up to 6 weeks. As shown in Figure 6B, in comparison with the mice treated with control-TAMs, VentX-TAMs exerted strong inhibition on the growth of PDA in the NSG-PDX model of primary human PDA.
Discussion
Phagocytosis underlies the fundamental function of macrophages for antigen presentation and immunosurveillance. However, the role of phagocytosis in tumorigenesis remains an area of active debate (26). Previous studies suggested that inhibition of phagocytosis may play an important role in tumor evasion of immune surveillance. It was shown that the expression of CD47, the do-not-eat-me signal, is elevated in a variety of human cancers. On the other hand, the expression of CD163, a scavenger receptor, is increased in the M2-like TAMs, and studies suggested that M2 macrophages phagocytose antibody-opsonized target cancer cells more efficiently than M1 macrophages (27, 28). Despite speculation, systemic examination of macrophage function in the pathogenesis of PDA has been scarce. In our current studies, using primary human pancreatic cancer as a model, we found that the phagocytosis function of PDA TAMs is significantly impaired in comparison with macrophages isolated from control normal pancreatic tissue. We found that VentX expression is drastically impaired in PDA TAMs compared to control TAMs.
reduced in TAMs of all cases of PDA tested in this study. The importance of these studies is suggested by our findings that restoration of VentX expression in PDA TAMs revokes the phagocytosis function of TAMs and that VentX-TAMs transform immunity by altering the molecular and cellular composition at the PDA TME. Interestingly, mechanistic exploration showed that VentX expression did not alter the expression of SIRP, the cellular receptor of CD47. On the other hand, we did find that ectopic expression of VentX inhibits phosphorylation of SHP-1 and SHP-2, SIRP downstream signaling molecules implicated in the inhibition of phagocytosis. Moreover, our data showed that VentX promotes the expression of TLRs and downstream signaling molecules as well as FAK signaling, which has been implicated in promoting phagocytosis. Our data are consistent with the prior notion of M1 and M2 signaling in the regulation of phagocytosis. We show here that this process is controlled by VentX, a human homolog of the Xenopus ventral homeobox protein Xom (29–31). Since the initial discovery of macrophages as the key regulator of immunity more than one hundred years ago, the cell-intrinsic factor that controls macrophage plasticity has remained elusive (32, 33). By leveraging the power of developmental study, which led to the discovery of VentX as the key regulator of macrophage plasticity (14, 17, 18, 30, 31, 34), our current findings suggested a fundamental mechanism of immune regulation, which underlies the central mechanism of immune suppression at the PDA TME. Currently, the mechanism underlying decreased VentX expression in PDA TAMs remains unknown. Our data suggest that further exploration along these lines may provide better mechanistic insight and potentially novel therapeutic opportunities to improve PDA prognosis.

Methods

Collection of pancreatic tissue samples. A total of 19 patients with pancreatic adenocarcinoma, who were scheduled for surgical resection at Brigham and Women’s Hospital, consented to have a portion of resected tissues and blood collected for research purposes. The characteristics of the patients with pancreatic cancer, whose specimens were used for this study, are listed in Table 1. Approximately 5–10 g tissues were collected from the tumor mass or adjacent normal tissues. Tumor samples and control tissues were verified by board-certified pathologists at Brigham and Women’s Hospital.

Preparation of lymphocytes and macrophages from pancreatic and tumor tissues. Lymphocytes were isolated essentially as described previously (14, 18, 35). Briefly, dissected fresh normal pancreatic tissues and pancreatic tumors were rinsed in a 10-cm Petri dish with Ca^{2+}-free and Mg^{2+}-free HBSS (Life Technologies) containing 2% FBS and 2 mM Dithiothreitol (MilliporeSigma). The pancreatic and tumor tissues were then cut into approximately 0.1-cm pieces with a razor blade and incubated in 5 mL HBSS containing 5 mM EDTA (MilliporeSigma) at 37°C for 1 hour. The tissues were then passed through a gray mesh (100 microns). The flow-throughs containing lymphocytes and epithelial cells were then analysis by flow cytometry.
To isolate the macrophages, normal pancreatic tissues and pancreatic tumors were rinsed with HBSS, cut into approximately 0.1-cm pieces with a razor blade, and then incubated in HBSS (with Ca²⁺ and Mg²⁺) containing 2% FBS, 1.5 mg/mL Collagenase D (Roche), and 0.1 mg/mL DNase I at 37°C for 1 hour. The digested tissues were then passed through a gray mesh (70-micron) filter. The flow-throughs were collected, washed, and resuspended in RPMI 1640 medium. Normal tissue macrophages and TAMs were further purified using the EasySep Human Monocyte/Macrophage Enrichment kit without CD16 depletion (StemCell Technologies, catalog 19085) according to the manufacturer’s instructions. The isolation process did not lead to activation of macrophages, and the purity of isolated macrophages was above 95% (14, 35, 36). Greater than 98% of cells isolated were viable by propidium iodide staining tests.

**FACS analysis.** Phenotypic analysis of macrophages and lymphocytes was performed using flow cytometry after immunolabeling of cells with fluorescence dye–conjugated antibodies. The following antibodies were used: PE-conjugated anti-CD3 (OKT3), -CD25 (BC96), -CD14 (61D3), -CD40 (2D10.4), -CD163 (GH161), and -PD-IL (J105); FITC-conjugated anti-CD4 (RPA-T4) and -CD68 (Y182A); and APC-conjugated anti-CD8 (OKT8), -CD4 (OKT4), and -SIRPα (clone 15-414) (eBioscience). Intracellular staining of anti-Foxp3 (236A/E7) APC and -IFN-γ (4S.B3) PE antibodies was performed following the protocol provided by the manufacturer (eBioscience). Isotope control labeling was performed in parallel. Antibodies were diluted as recommended by the manufacturer (eBioscience). Labeled cells were acquired using the BD LSRRFortessa at the Flow Cytometry Core of the Dana-Farber Cancer Institute with FACS Diva software (BD Biosciences) and analyzed using FlowJo 10.1 software (Tree Star). Typically, 20,000 events were analyzed per sample according to the standard FACS analysis procedure. Results are expressed as the percentage of positive cells.

**Quantitative RT-PCR.** Total RNA was isolated by the TRIzol reagent (Life Technologies), and RNA amounts were measured by NanoDrop 2000 (Thermo Scientific). Equal amounts of RNA were used for first-strand cDNA synthesis with the SuperScript III First-Strand Synthesis System (Life Technologies) according to the manufacturer’s protocol. The AccuPrime Taq DNA polymerase system (Life Technologies) was used to amplify VentX cDNA with conventional PCR. Quantitative measurements of VentX and other cDNA were carried out with SYBR Green (LightCycler 480 SYBR Green Master kit, Roche Life Science), using a Mastercycler ep Gradient S (Eppendorf). GAPDH was used as a housekeeping gene to normalize mRNA expression. The primers used are listed in Supplemental Table 1. Relative expression profiles of mRNAs was calculated using the comparative Ct method (DDCT method).

**Cytokine measurement.** Levels of IL-1β and TNF-α were quantified using ELISA kits obtained from eBioscience. Analyses were conducted according to the manufacturer’s instructions. Triplicate wells were plated for each condition. Nitrite concentrations in tissue culture supernatants were determined using the Griess reagent kit (Molecular Probes), as described previously (14).

**Transfection assays.** Plasmids encoding GFP-VentX and control GFP were described previously (31). Transfection of GFP-VentX and GFP into macrophages was carried out using the Human Macrophage Nucleofector Kit (VVPA-1008, Lonza). Briefly, 2 × 10⁶ cells were resuspended in 100 μl nucleofector solution with 5 μg plasmid DNA for 20 minutes on ice. Transfections were performed on a Nucleofector 2b Device (Lonza). After transfection, cells were placed on ice immediately for 1 minute and then cultured in prewarmed RPMI 1640 complete medium, containing 10% FBS and 1% antibiotic-antimycotic solution (Gibco, 15240062) for 48 hours before transfected cells were used for experiments.

For VentX knockdown experiments, pancreatic TAMs were transfected with morpholino oligonucleotides (MOs) (Gene Tools) using the Human Macrophage Nucleofector Kit (Lonza) as previously described (14, 18). Briefly, 2 × 10⁶ cells were resuspended in 100 μl nucleofector solution, with 2.5 nmol of either VentX MO (5′-TACTCAACCCCTGACATAGAGGGTA-3′) or a control MO oligonucleotide (5′-CCTCTTACCT-CAGTTACAATTATA-3′) and electroporated with the Nucleofector II Device (Lonza). Cells were then immediately removed from the device and placed on ice for 1 minute. The cells were then incubated overnight with 1 ml prewarmed Human Monocyte Nucleofector Medium (Lonza Bioscience) containing 2 mM glutamine and 10% FBS. Cells were then resuspended in complete RPMI medium and stimulated with 10 ng/mL LPS (LiporeSigma) or control PBS overnight for phagocytosis assays.

**Phagocytosis assays.** For the phagocytosis assay, PANC-1 cells (human pancreatic cancer cell line, MilliporeSigma) or primary human leukemia cells (Brigham and Women’s Hospital) were labeled with 1 μM CellTrance Yellow reagent using the CellTrace CFSE proliferation kit (Thermo Fisher Scientific). 5 × 10⁶ TAMs transfected with GFP or GFP-VentX were plated in each well of 12-well tissue culture plates (Cor-
ing). The TAMs were then incubated in RPMI complete medium for 2 hours before mixing with $1 \times 10^6$ of CSFE Yellow–labeled cancer cells. After the indicated incubation time, the TAMs were washed repeatedly and detached with 0.25% Trypsin/EDTA; phagocytosis was analyzed by a flow cytometer. Microscopic analysis of phagocytosis was performed with a Nikon Eclipse Ti fluorescence microscopy. Images were captured at magnifications of ×20 or ×40 using a color camera and the NIS Elements imaging software (Nikon). Brightness and contrast for representative images were adjusted equally among groups.

**Western blot analysis.** Western blot analysis was performed as described previously (14, 18). Briefly, cells were lysed in 1× RIPA buffer (Abcam) mixed with phosphatase and protease inhibitor cocktails (Cell Signaling Technology [CST]). Proteins were resolved by 4%–15% Tris-Glycine Gel (Bio-Rad) electrophoresis. Primary antibodies used included VentX (Abcam, ab105352, 1:500), β-actin (CST, 4967, 1:2000), anti–phospho-SHP-1 (Tyr564) (CST, catalog 8849, 1:500), SHP-2 (Tyr542) (CST, catalog 3751, 1:500), and phosphor-FAK (Tyr397) (CST, catalog 3283, 1:500) antibodies. Their protein control antibodies included SHP-1 (CST, catalog 3759, 1:1000), SHP-1 (CST, catalog 3397, 1:1000), and FAK (CST, catalog 3285, 1:1000). The relative fold difference of the band density was determined by ImageJ software (NIH).

**T cell proliferation assays.** T cell proliferation was performed essentially as described previously (18, 37). Briefly, blood CD8 cells of pancreatic patients were enriched by the Easysep human CD8 enrichment kit (StemCell Technologies, catalog 19053) following the manufacturer's instructions. $1 \times 10^6$ CD8 cells were labeled with 1 μM of CellTrance Yellow and then activated with human T-Activator CD3/CD28 Dynabeads (Gibco, catalog 11131D) at a bead-to–T cell ratio of 1:4 in the presence of 10 ng/mL IL-2 (PeproTech). Activated CD8 was then mixed with 0.25 × $10^6$ GFP-VentX– or GFP-transfected TAMs of the same patients. The mixtures were incubated in completed RPMI 1640 at 37°C, 5% CO₂ for 5 days. Cells were then stained with an anti–CD8-APC–conjugated antibody and analyzed by a flow cytometer.

**Cocultures of tumor tissues and macrophages.** Tumor tissues were washed with 1× PBS buffer plus antibiotics and then cut into 0.5-cm pieces. Tissues were mixed and cultured with 0.5 × $10^6$ GFP-VentX– or GFP-transfected macrophages of the same patient in 2 mL RPMI 1640 complete medium, supplemented with 1% antibiotic-antimycotic solution (Gibco). The cultures were incubated at 37°C, 5% CO₂, for 5 days. The tissues were then subjected to cell isolation and analyzed by a flow cytometry or immunohistochemistry studies.

**NSG-PDX model of human pancreatic cancers.** Animal models of primary human pancreatic cancers were developed as described previously (38). Briefly, 8-week-old NOD.Cg-Prkdc<sup>scid</sup> Ikr2g<sup>−/−</sup>/SzJ mice (commonly known as NOD/SCID/γ [NSG] mice) were purchased from The Jackson Laboratory and maintained under specific pathogen–free conditions. Pancreatic tumors were cut into about 0.5-cm pieces and then surgically implanted into subcutaneous space on the dorsal side of NSG mice. After 1 week of implantation, 0.5 × $10^6$ TAMs transfected with GFP-VentX or control GFP were injected into the mice through tail vein. The tumor growth was monitored twice weekly and measured by a caliper for 6 weeks.

In vivo tracing of injected TAMs was carried out using CFSE-stained TAMs or macrophages, according to the manufacturer’s instructions (Molecular Probes). Briefly, $1 \times 10^6$ TAMs were incubated in 5 mL of

| Characteristics | Number (n = 19) | (%) |
|-----------------|----------------|-----|
| **Age (y)**     |                |     |
| ≤60             | 2              | 10.5|
| 60–79           | 15             | 79.0|
| ≥80             | 2              | 10.5|
| **Sex**         |                |     |
| Male            | 12             | 63.2|
| Female          | 7              | 36.8|
| **Pathology type** |              |     |
| Adenocarcinoma  | 19             | 100 |
| **Stage**       |                |     |
| I               | 4              | 21.1|
| II              | 12             | 63.2|
| III             | 3              | 15.7|
| IV              | 0              | 0   |
5 μM CFSE staining solution in a 37°C water bath for 20 minutes. Cells were then incubated with 20 mL RPMI 1640 complete medium for 5 minutes to removed unbounded dye. After centrifugation, cells were dissolved in PBS and then tail-vein injected into NSG mice. Tumor tissues and lymph nodes were dissected out 3 days after injection, and TAMs were isolated as CFSE-positive cells.

**Immunohistochemistry.** Immunohistochemistry was performed following established protocols (39). Briefly, pancreatic tumors or adjacent normal tissues were fixated in formalin (Thermo Fisher Scientific) for at least 48 hours. The tissues were then embedded in paraffin and sectioned. CD8 (Dako, clone C8/144B, 1:100), CD163 (Vector, clone 10D6, 1:250), Vimentin (Dako, clone M0725, 1:50), and H&E staining was performed on the Specialized Histopathology Core at Dana-Farber/Harvard Cancer Center. All immunohistochemistry was performed on the Leica Bond automated staining platform, using the Leica Biosystems Refine Detection Kit with EDTA antigen retrieval. The images of whole slides were scanned by a Pannoramic MIDI II digital slide scanner and analyzed with Caseviewer and Quant center software (3DHistech).

**Statistics.** Two-tailed Student’s t test or 1-way ANOVA were used for statistical analysis in Prism (version 8, GraphPad). Data are presented as mean ± SD. Tumor growth curves were analyzed by repeated measurement 2-way ANOVA using Holm-Šidák multiple comparison test. The level of significance was indicated by the P value. P values of 0.05 or less were considered statistically significant.

**Study approval.** All patients provided written informed consent, and the Institutional Review Board of the Brigham and Women’s Hospital approved these studies (2006P1354). Tumor samples and control tissues were verified by board-certified pathologists at Brigham and Women’s Hospital. All animal experiments were approved by the Institutional Animal Care and Use Committee at the Brigham and Women’s Hospital (2016N000353).

**Author contributions**
YL, HG, and ZZ designed the research, analyzed the data, and prepared figures. YL and HG performed the experiments. RB, WR, LZ, and TC provided tumor samples, clinical information, and analysis. ZZ wrote the manuscript.

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