A Cancer Cell-Intrinsic GOT2-PPARδ Axis Suppresses Antitumor Immunity

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ABSTRACT

Despite significant recent advances in precision medicine, pancreatic ductal adenocarcinoma (PDAC) remains near uniformly lethal. Although immune-modulatory therapies

hold promise to meaningfully improve outcomes for patients with PDAC, the development of such therapies requires an improved understanding of the immune evasion mechanisms that characterize the PDAC microenvironment. Here, we show that cancer cell-intrinsic glutamic-oxaloacetic transaminase 2 (GOT2) shapes the immune microenvironment to suppress antitumor immunity. Mechanistically, we find that GOT2 functions beyond its established role in the malate-aspartate shuttle and promotes the transcriptional activity of nuclear receptor peroxisome proliferator-activated receptor delta (PPAR\delta), facilitated by direct fatty acid binding. Although GOT2 is dispensable for cancer cell proliferation in vivo, the GOT2-PPARδ axis promotes spatial restriction of both CD4+ and CD8+ T cells from the tumor microenvironment. Our results demonstrate a noncanonical function for an established mitochondrial enzyme in transcriptional regulation of immune evasion, which may be exploitable to promote a productive antitumor immune response.

SIGNIFICANCE: Prior studies demonstrate the important moonlighting functions of metabolic enzymes in cancer. We find that the mitochondrial transaminase GOT2 binds directly to fatty acid ligands that regulate the nuclear receptor PPAR δ , and this functional interaction critically regulates the immune microenvironment of pancreatic cancer to promote tumor progression.

See related commentary by Nwosu and di Magliano, p. 2237.

INTRODUCTION

Dual functions for glutamic-oxaloacetic transaminase 2 (GOT2) are described in the literature. The far better studied function is as a mitochondrial transaminase, implicated in the maintenance of the malate-aspartate shuttle and redox homeostasis (1-4). However, a limited body of evidence indicates a role for GOT2 in fatty acid binding and trafficking (5-10), though this role remains poorly understood and has not been investigated in cancer. In these studies, GOT2 is often referred to as plasma membrane fatty acid binding protein (FABPpm) due to its membrane-proximal localization in hepatocytes and the ability of GOT2/FABPpm antiserum to disrupt fatty acid trafficking in metabolic cell types, including hepatocytes and cardiomyocytes (5, 11-13). In light of recent work from our group and others documenting the importance of fatty acid

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(PDAC) growth, at least in part through its fatty acid trafficking function. GOT2 is overexpressed in human PDAC (19), and although transmembrane fatty acid transporters were variably expressed, GOT2 was consistently expressed in human PDAC per two independent RNA sequencing (RNA-seq) data sets (Supplementary Fig. S1A and S1B), as was the cytosolic isoform GOT1. We set out to determine whether GOT2 plays a role in PDAC progression in vivo and, if so, to understand the relevance of its established mitochondrial role versus its less characterized role in spatial regulation of fatty acids. RESULTS

trafficking for solid tumor progression (14-18), we considered that GOT2 may promote pancreatic ductal adenocarcinoma

To assess the significance of GOT2 for PDAC progression, we generated several loss-of-function systems using short hairpin RNA (shRNA) or CRISPR/Cas9 and using human and murine PDAC cells (Supplementary Fig. S2A). Cas9 and single-guide RNAs (sgRNA) were introduced by transient transfection, and Cas9 was no longer expressed by the time cells were used for in vivo studies; GOT1 levels were unchanged (Supplementary Fig. S2B). Across all cell lines tested, only two showed proliferation defects (Fig. 1A; Supplementary Fig. S2C). These defects were modest, and, in one of the two lines, a reduction in proliferation was seen only upon inducible GOT2 knockdown, suggesting that PDAC cells have sufficient metabolic plasticity to adapt to GOT2 loss and maintain proliferative capacity. However, when sgGot2 PDAC cells were transplanted into pancreata of immune-competent syngeneic hosts, tumor growth was severely compromised (Fig. 1B). Consistent with in vitro results, proliferation among tumor cells was not impaired in vivo (Fig. 1C). An independent model also revealed a critical role for GOT2 in PDAC growth, whether GOT2 was knocked down with shRNA (Fig. 1D) or knocked out with CRISPR/Cas9 (Fig. 1E; Supplementary Fig. S2D). Though shRNA-mediated knockdown had a less dramatic effect on tumor growth, we



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Figure 1. GOT2 promotes pancreatic tumor progression without impacting proliferation. **A**, Viable cell measurements in the indicated PDAC cell lines. Data are presented as mean ± SEM from biological triplicates. *****, *P* < 0.0001 by one-way ANOVA. Dox, doxycycline; RLU, relative light unit. Numbers in parentheses are the short hairpins (shRNAs) used for the GOT2 knockdowns. **B**, PDAC tumor weight at the experimental endpoint, 34 days after orthotopic transplantation of 688M cells into immune-competent hosts. Ctrl: *n* = 8, sgGot2 a: *n* = 9, sgGot2 b: *n* = 10. Data are presented as mean ± SEM. ns = not significant. *****, *P* < 0.0001 by one-way ANOVA. **C**, IHC staining of tumors in **B** for Ki-67 (proliferation) and pan-cytokeratin (panCK; tumor cells), with a DAPI counterstain (nuclei). Representative images are shown on the left (scale bars = 50 µm), with quantification on the right (ctrl: *n* = 6, sgGot2: *n* = 5). Data are presented as mean ± SEM. ns = not significant. ****, *P* < 0.001 by one-way ANOVA. **C**, IHC staining of tumors in **B** for Ki-67 (proliferation) and pan-cytokeratin (panCK; tumor cells), with a DAPI counterstain (nuclei). Representative images are shown on the left (scale bars = 50 µm), with quantification on the right (ctrl: *n* = 6, sgGot2: *n* = 5). Data are presented as mean ± SEM. ns = not significant test. **D**, PDAC tumor weight at the experimental endpoint, 22 days after orthotopic transplantation of FC1245 cells into immune-competent hosts. Ctrl: *n* = 5, shGot2 b: *n* = 3. Data are presented as mean ± SEM. ns = not significant. ***, *P* < 0.001 by one-way ANOVA. **E**, PDAC tumor weight at the experimental endpoint, 18 days after orthotopic transplantation of FC1245 cells into immune-competent hosts. Ctrl: *n* = 5, shGot2 b: *n* = 3. Data are presented as mean ± SEM. ns = not significant. ***, *P* < 0.001 by one-way ANOVA. **E**, PDAC tumor weight at the experimental endpoint, 18 days after orthotopic transplantation of FC1245 cells into immune-competent hosts. Ctrl: *n* = 5, sgG

noted a partial recovery of GOT2 expression in these tumors by the experimental endpoint (Supplementary Fig. S2E). These results indicated that GOT2 is dispensable for PDAC cell proliferation but required for tumor growth *in vivo* and raised the possibility that cancer cell-intrinsic GOT2 promotes growthpermissive regulation of the tumor microenvironment. To gain insight into GOT2 function in an intact tumor microenvironment, we identified transcriptional programs with expression inversely correlated with *GOT2* transcript abundance in The Cancer Genome Atlas (TCGA) RNAseq data (Fig. 2A). Pathway analysis of this group of genes revealed enrichment for genes associated with lymphocyte

Figure 2. PDAC cell-intrinsic GOT2 suppresses T cell-dependent immunologic control of tumor growth. **A**, Metascape pathway analysis depicting the top transcriptional programs inversely correlated with *GOT2* expression in human PDAC. **B** and **C**, IHC staining of control and sgGot2 688M tumors for T-cell marker CD3 (**B**) and subtype markers CD4 and CD8 (**C**). Representative images are shown on the left (scale bars = 50 µm), with quantification on the right (ctrl: *n* = 5, sgGot2 a: *n* = 4, sgGot2 b: *n* = 4). Data are presented as mean ± SEM. *, *P* < 0.05; **, *P* < 0.01; ***, *P* < 0.001 by one-way ANOVA. **D**, IHC staining of control and shGot2 FC1245 tumors for T-cell markers CD3 and CD8. Representative images are shown on the left (scale bars = 50 µm), with quantification on the right (ctrl: *n* = 5, shGot2 a: *n* = 5, shGot2 b: *n* = 3). Data are presented as mean ± SEM. **, *P* < 0.01; ***, *P* < 0.001 by one-way ANOVA. **D**, IHC staining of control and sgGot2 or shGot2 PDAC for macrophage marker F4/80 and immunosuppressive factor arginase-1 (Arg1). Representative images are from 688M tumors (scale bar on 20× images = 10 µm, scale bar on 63× images = 5 µm). Quantification of double-positive cells out of total F4/80⁺ cells in the 688M and FC1245 models is on the right; data are presented as mean ± SEM. **, *P* < 0.01; ****, *P* < 0.0001 by an unpaired t test. **F**, Multiplex IHC staining of control and sgGot2 FC1245 tumors for the indicated markers (large images, scale bar = 50 µm). G, Quantification of CD3 IHC on 688M PDAC at the indicated time points after transplantation (ctrl 11: *n* = 7, sgGot2 a d11: *n* = 6, ctrl d19: *n* = 3, sgGot2 a d19: *n* = 3, sgGot2 a d27: *n* = 4). *, *P* < 0.01; ***, *P* < 0.0001 by an unpaired t test. **F**, Multiplex IHC staining of control and sgGot2 FC1245 tumors for the indicated markers (large images, scale bar = 50 µm). G, Quantification of CD3 IHC on 688M PDAC at the indicated time points after transplantation (ctrl 11: *n* = 7, sgGot2 a d11: *n* = 6, ctrl d19: *n*







differentiation, activation, and adhesion and led us to question whether cancer cell-intrinsic GOT2 regulates the abundance and/or activity of intratumoral T cells. Transcripts positively correlated with GOT2 were involved in metabolic processes (Supplementary Fig. S2F). We quantified T cells in two independent GOT2 loss-of-function models and found that T-cell frequencies were increased in sgGot2 or shGot2 tumors compared with controls, including CD4⁺ and CD8⁺ T cells (Fig. 2B-D). As PDAC contains high numbers of immunesuppressive myeloid cells, including abundant macrophages, which contribute to T-cell exclusion (20-22), we assessed macrophage abundance and phenotype in these tumor tissues. We found that loss of GOT2 in cancer cells increased total macrophage abundance while decreasing the frequency of Arg1⁺ macrophages out of total macrophages (Fig. 2E), consistent with macrophage polarization to a less immunesuppressive phenotype permissive to T-cell recruitment. To further characterize the immune infiltrates of these tumors, we performed multiplex IHC (see Methods for details), which revealed relatively abundant Ki-67+CD8+ T cells (Fig. 2F) and CD11b⁺ myeloid cells (Supplementary Fig. S2G) in proximity to pan-cytokeratin⁺ tumor cells lacking GOT2 compared with controls. Quantification revealed that GOT2-null PDAC cells were associated with the increased presence of CD11b⁺PD-L1⁺ and CD11b+Ki-67+ cells (Supplementary Fig. S3A and S3B); increased presence of total TCF1/7+ and PD-1+ T cells as well as CD8+Ki-67+ and CD8+PD-1+Ki-67+ T cells; and increased trends for CD8+GRZB+ and CD8+PD-1+GRZB+ T cells, though these trends did not reach statistical significance (Supplementary Fig. S3C-S3H). To address whether the differences in T-cell abundance were secondary to differences in tumor size, we performed a time course and harvested tumors soon after transplantation to quantify intratumoral T cells. At 11 days after transplantation, a time point when tumors are small in control and sgGot2 tumors but not yet different in size (Supplementary Fig. S3I), T-cell frequencies were already increased in the GOT2-null setting (Fig. 2G); T-cell frequency was also increased at days 19 and 27 after transplantation, though tumors are different in size by these early time points. To further understand the impact of cancer cell-intrinsic GOT2 on the immune microenvironment, we analyzed the myeloid compartment of these tumors and identified pronounced changes to macrophage and dendritic cell populations including increases in proliferating macrophages and cDC1 (Supplementary Fig. S4A-S4M). We next asked whether these T cells were in fact functional in suppressing tumor progression. To address this, we treated control and sgGot2 tumors with depleting antibodies against CD4⁺ and CD8⁺ T cells. This intervention had no impact on the growth of control tumors, consistent with previous studies documenting a lack of T cellmediated antitumor immunity in mouse models of PDAC (23). However, the growth of sgGot2 tumors was restored upon T-cell neutralization (Fig. 2H), indicating that GOT2 promotes PDAC progression at least in part by suppressing T cell-dependent antitumor immunity.

We next addressed the mechanism by which cancer cellintrinsic GOT2 influences the immune microenvironment, taking potential enzymatic and fatty acid binding functions into account. To begin to address this, we examined GOT2 localization in PDAC cells and found that a pool of this canonically mitochondrial protein localizes to the nucleus in murine premalignant lesions and PDAC as well as human PDAC in vivo (Fig. 3A and B; Supplementary Fig. S5A). We note that, although all human PDAC specimens examined showed evidence of nuclear GOT2 in pan-cytokeratin⁺ tumor cells, tumor cells with GOT2 restricted to mitochondrial and membrane-proximal regions and without nuclear GOT2 were also observed across these samples. This nuclear GOT2 pool was also evident in vitro, whether we analyzed endogenous or exogenous, His-tagged GOT2 (Supplementary Fig. S5B-S5E). We reasoned that the intact proliferation of GOT2null tumors indicated the presence of metabolic adaptation mechanisms to retain redox balance, and this motivated us to consider noncanonical functions of GOT2 related to its putative fatty acid binding capacity. The previously unappreciated nuclear pool of GOT2 led us to hypothesize that GOT2 regulates nuclear trafficking of fatty acids either into or within the nucleus. Nuclear fatty acid trafficking has been reported to be regulated by fatty acid binding proteins (24, 25), and nuclear fatty acids have established functional significance as ligands for the peroxisome proliferator-activated receptor (PPAR) members of the nuclear receptor superfamily of transcription factors (26). This three-member family is activated by fatty acid ligands, and although PPARα and PPARγ display tissuerestricted expression and PPARy is downregulated during PDAC progression (27, 28), PPARδ is ubiquitously expressed and was expressed in all PDAC lines examined whether or not GOT2 was inhibited (Supplementary Fig. S6A). Analysis of human PDAC RNA-seq data revealed a correlation between the expression of GOT2 and the PPARS regulome (Fig. 3C; Supplementary Fig. S6B). Importantly, PPAR& promotes tumorigenesis via tissue-specific metabolic and immunemodulatory mechanisms (29-32), prompting us to test a functional relationship between GOT2 and PPAR8 that may underlie the phenotypes of GOT2-null PDAC.

Transcriptional activity from a PPAR response element (PPRE) was reduced in GOT2-null PDAC cells (Fig. 3D), indicating that GOT2 positively regulates PPARS activity. Unlike steroid-activated nuclear receptors, which are sequestered in the cytoplasm in the absence of ligand and translocate to the nucleus upon ligand engagement, PPARo is constitutively nuclear and bound to DNA but undergoes a conformational change upon binding of nuclear fatty acids to enable interaction with coactivator complexes, altered DNA binding, and induction of target gene expression (33). Further supporting positive regulation of PPARδ transcriptional activity by GOT2, nuclear extracts from control and sgGot2 cells were applied to wells containing immobilized, PPRE-containing DNA, followed by incubation with a PPAR δ antibody and a peroxidase-conjugated secondary antibody. Results of this assay indicated reduced PPARS transcriptional activity in GOT2-null PDAC cells (Fig. 3E). To identify immune-modulatory factors putatively regulated by the GOT2-PPARδ axis, we performed a cytokine array on a conditioned medium from PDAC cells and found reductions in myeloid cellmodulating factors REG3G and M-CSF with GOT2 loss (Supplementary Fig. S6C and S6D) and thus included the associated genes in our analyses. Chromatin immunoprecipitation (ChIP) for PPAR8 and acetylated histone H3K9, a marker of active promoters, followed by qPCR also supported



Figure 3. GOT2 positively regulates PPAR& activity. **A**, IHC staining for GOT2 or GOT2 and panCK in pancreas tissues from *Kras*^{LSL-G12D/+};*Pdx*1-*Cre* (KC) mice at 6 or 12 months of age (representative of *n* = 4 per time point). Scale bars = 20 µm. **B**, IHC staining for GOT2 or GOT2 and panCK in human PDAC (representative of *n* = 5). Fluorescent images: scale bar = 5 µm, brightfield image: scale bar = 20 µm. Arrowheads indicate examples of tumor cells with nuclear GOT2. **C**, Scatter plot depicting the correlation of *GOT2* expression with expression of PPAR& target genes in human PDAC per TCGA RNA-seq data (*n* = 177). FPKM, fragments per kilobase of transcript per million mapped reads. **D**, Luciferase assay for PPRE activity in the indicated cell lines, normalized to Renilla, presented as mean ± SEM. ****, *P* < 0.001 by one-way ANOVA (688M) or an unpaired t test (FC1245). **E**, PPAR& transcriptional activity assay, reading out binding to immobilized DNA containing PPREs, in the indicated cell lines. Data are presented as mean ± SEM from four (FC1245) or three (8988T) independent experiments. **, *P* < 0.01 by an unpaired t test. **F** and **G**, ChIP for H3K9Ac (**F**) and PPAR& (**G**) in control or sgGot2 688M PDAC cells, followed by qPCR for proximal promoter regions of the indicated genes. Data were normalized to an intergenic region (Int. B) and are presented as mean ± SEM from biological triplicates. ns = not significant. *, *P* < 0.001; ****, *P* < 0.001 by one-way ANOVA. I, Luciferase assay for PPRE activity in sgGOT2 PDAC cells reconstituted with whicle (DMSO) or the PPAR& synthetic agonist GWSO1516 (GW; 100 nmol/L). Data are presented as mean ± SEM from four independent experiments. ****, *P* < 0.001 by one-way ANOVA. (continued on next page)

a reduction of PPAR δ transcriptional activity in the absence of GOT2 (Fig. 3F and G). Some of these genes previously linked to PPAR δ activity appear potentially to be indirect targets. Expression of PPAR δ target genes was also reduced in GOT2-null PDAC cells, and the synthetic PPAR δ agonist GW501516 restored target gene expression, indicating that these genes are indeed regulated by PPAR δ (Fig. 3H). Among the genes with clear relevance to our *in vivo* phenotype was *PTGS2*, which encodes COX2. Recently reported gain- and loss-of-function experiments revealed that COX2 promotes



Figure 3. (Continued) J, PDAC tumor weight at the experimental endpoint, 22 days after orthotopic transplantation of FC1245 cells into immunecompetent hosts (n = 4-5 per arm). Data are presented as mean ± SEM. ns = not significant. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P <

T-cell exclusion from the PDAC microenvironment, consistent with our results, and that PTGS2 expression correlated with poor patient survival (34). COX2 has also been implicated in the suppression of antitumor immunity in other cancer types (35). We further investigated the regulation of COX2 downstream of GOT2 and found that COX2 protein levels were reduced in GOT2-null PDAC cells in vitro and in vivo (Supplementary Fig. S7A and S7B). Although GOT2 lacks a nuclear localization sequence (NLS), we added an NLS to GOT2 and found that increasing GOT2 nuclear localization increased PPARS activity (Fig. 3I; Supplementary Fig. S7C). Although NLS-GOT2 did not increase tumor growth compared with wild-type GOT2 in vivo, increased GOT2 nuclear targeting reduced T-cell abundance in these tumors, and Arg1+ macrophage abundance was not significantly altered (Fig. 3J-N; Supplementary Fig. S7D and S7E).

Reconstitution with wild-type GOT2 restored immune suppression as expected, though CD8 T-cell abundance was not reduced all the way down to control levels (Fig. 3K-N). These results together support the hypothesis that GOT2 promotes the transcriptional activity of PPAR δ in PDAC cells.

As we were prompted to investigate a GOT2-PPAR δ functional interaction based on the putative fatty acid binding function of GOT2, we investigated this role further. For this, we analyzed the crystal structure of human GOT2 (36) and identified five putative fatty acid binding sites based on hydrophobicity (Fig. 4A and B). We then performed *in silico* docking studies for known fatty acid ligands for PPAR δ and identified a potential interaction between arachidonic acid and GOT2 hydrophobic site 2 (Fig. 4C). This modeled interaction yielded a docking score of -7.6 kcal/mol, which is very similar to the docking score calculated for arachidonic acid in



Figure 4. GOT2 binds to the PPARô ligand directly. A, Hydrophobic site maps on the GOT2 protein, indicating putative fatty acid binding domains. Red: hydrogen bond acceptor, blue: hydrogen bond donor, green: hydrophobic. B, Plot of the hydrophobic area of the putative fatty acid binding sites depicted in A. C, Docking model of arachidonic acid in site 2 on the GOT2 protein, with bioenergetic docking score (–7.6 kcal/mol) indicated below. D, Competitive fatty acid binding assay measuring radioactivity upon incubating purified human GOT2 with ³H-arachidonic acid (1 µmol/L) and the indicated concentrations of cold lipid species. Ara, cold arachidonic acid. E, Left, model of arachidonic acid bound to GOT2, indicating amino acid residues that potentially facilitate binding. Based on this model, K234, K296, and R303 were selected for mutation to alanine. Right, conservation of GOT2 amino acid sequence, including the three residues predicted to support arachidonic acid binding, among higher vertebrates. (continued on next page)

the ligand binding domain of PPAR_γ (-7.0 kcal/mol; ref. 37), an interaction that is known to be direct and functionally significant. To probe this interaction further, we performed competitive fatty acid binding assays using purified GOT2 protein and radiolabelled arachidonic acid. In addition to cold arachidonic acid, we used cold oleic acid, as this was previously reported to bind to GOT2 (ref. 10; though our analysis revealed a distinct fatty acid binding domain from this previous study) as well as prostaglandin D2 (PGD₂), a downstream metabolite of arachidonic acid that we predicted to serve as a negative control and not to bind directly to GOT2 based on computational modeling. The competitive binding assay showed that arachidonic acid indeed bound to GOT2 directly, and although cold arachidonic acid readily displaced radiolabeled ligand, our negative control lipid (PGD₂) was unable to compete away the arachidonic acid signal even when PGD₂ concentration exceeded that of arachidonic acid by three orders of magnitude (Fig. 4D), supporting a specific interaction. Oleic acid had a modest effect on binding, suggesting that oleic acid may bind to the arachidonic acid-bound site but at a lower affinity or may bind to a separate site on the protein. To assess a relationship between GOT2 and arachidonic acid trafficking in cells, we performed mass spectrometry to measure arachidonic acid in whole cells and nuclei; levels were unchanged at the whole-cell level between control and sgGot2 cells (Supplementary Fig. S7F), but nuclear levels were below a reliable level of detection. We developed an assay to measure nuclear arachidonic acid accumulation by spiking fluorescent arachidonic acid into our culture medium and measuring fluorescent signal in isolated nuclei, which revealed a reduction in nuclear arachidonic acid accumulation in two GOT2 loss-of-function cell lines (Supplementary Fig. S8A). Though significant, these differences were modest, suggesting that GOT2 may regulate arachidonic acid within the nucleus as opposed to predominantly regulating its nuclear import. To address the functional significance of GOT2 fatty acid binding, we looked closely at the putative fatty acid binding pocket we identified and selected three key amino acid residues that we predicted to be critical for arachidonic acid binding at that site (Fig. 4E). We note conservation of this amino acid sequence across mammalian species and between GOT1 and GOT2. We mutated these three residues on His-tagged GOT2 and used this triplemutant GOT2 (tmGOT2) or wild-type GOT2 (wtGOT2) to reconstitute sgGot2 PDAC cells. Although wtGOT2 localized to mitochondria and nuclei, tmGOT2 exhibited a reduction in nuclear localization compared with the wild-type protein (Fig. 4F), raising the possibility that fatty acid binding at this site promotes GOT2 nuclear trafficking, perhaps via interaction with a chaperone. To further probe the relationship between nuclear GOT2 and PPARo, we immunoprecipitated GOT2 from whole cells and found that wtGOT2 interacts with PPARo, whereas this interaction is reduced upon disruption of fatty acid binding (Fig. 4G). After confirming that tmGOT2 retains enzymatic activity (Supplementary Fig. S8B and S8C), we assessed PPARo activity and found that target gene expression and transcriptional activity were reduced in cells expressing tmGOT2 compared with wtGOT2 (Fig. 4H and I). We next transplanted immune-competent mice with control, sgGot2, sgGot2 + wtGot2, or sgGot2 + tmGot2



Figure 4. (Continued) **F**, Western blots indicating nuclear and whole-cell abundance of wtGOT2 and tmGOT2 (both His-tagged) in reconstituted sgGOT2 FC1245 PDAC cells. Nuclear GOT2 quantification appears to the right. **, *P* < 0.01 by an unpaired t test. **G**, Western blots from whole-cell lysates or His pulldowns from the cells depicted in **F**. ****, *P* < 0.001 by an unpaired t test. **H**, qPCR for the indicated PPAR&-regulated genes in FC1245 stable cell lines, normalized to 36b4. Data are presented as mean ± SEM from biological triplicates. ns = not significant. **, *P* < 0.001; ****, *P* < 0.0001 by one-way ANOVA. **I**, PPAR& transcriptional activity assay in the indicated FC1245 stable cell lines. Data are presented as mean ± SEM from three to six biological replicates. ns = not significant. **, *P* < 0.01; ****, *P* < 0.0001 by one-way ANOVA. **J**, PDAC tumor weight at the experimental endpoint, 18 days after orthotopic transplantation of the indicated FC1245 cells. Ctrl: *n* = 5, sgGot2 a + wtGOT2: *n* = 4, sgGot2 a + tmGOT2: *n* = 5. Ctrl and sgGot2 arms here are also depicted in Fig. 1E. Data are presented as mean ± SEM. ns = not significant. **, *P* < 0.001 by one-way ANOVA. **L**, PAR& transcriptional activity assay in the indicated FC1245 stable cell lines. Data are presented as mean ± SEM from three to six biological replicates. ns = not significant. **, *P* < 0.001 by one-way ANOVA. **J**, PDAC tumor weight at the experimental endpoint, 18 days after orthotopic transplantation of the indicated FC1245 cells. Ctrl: *n* = 5, sgGot2 a + wtGOT2: *n* = 4, sgGot2 a + tmGOT2: *n* = 5. Ctrl and sgGot2 arms here are also depicted in Fig. 1E. Data are presented as mean ± SEM. ns = not significant. **, *P* < 0.001 by one-way ANOVA. **K** and **L**, IHC staining (scale bars = 50 µm) and quantification for T cells (**K**; CD3) and macrophages (**L**; F4/80 and Arg1) in PDAC harboring wtGOT2 or tmGOT2 (*n* = 5 per arm). Data are presented as mean ± SEM. ns = not significant. *, *P* < 0.001 **ky** or an unpaired *t* test (**L**).

PDAC cells. We noted that wtGot2 had a strong nuclear localization in vivo, whereas tmGot2 was heterogeneous, with strong nuclear signal in some cells and a diffuse staining pattern in others (Supplementary Fig. S8D). Although wtGot2 completely rescued tumor growth as expected, tmGot2 only partially rescued tumor growth (Fig. 4J) and immune suppression (Fig. 4K and L). Disruption of GOT2 transaminase activity (1) did not affect tumor growth (Supplementary Fig. S9A-S9C). To further test the importance of enzymatic activity for the nuclear function of GOT2, we treated inducible shGOT2 8988T PDAC cells, sensitive to acute GOT2 inhibition with respect to proliferation, with the pantransaminase inhibitor aminooxyacetate (AOA; ref. 38). We found that although AOA reduced the proliferation of these cells, transaminase inhibition had no impact on PPARS activity (Supplementary Fig. S9D-S9G). Together, these results indicate a significant role for the fatty acid binding region in GOT2-mediated PDAC progression.

Based on these results, we hypothesized that PPARS activation would restore PDAC growth in the GOT2-null setting. We treated PDAC cells with GW501516, as we found this to override the limitation on PPARS activity in sgGot2 cells in vitro, and observed no increase (in fact, a decrease) in proliferation (Fig. 5A). However, GW501516 treatment in vivo rescued the growth of GOT2-null PDAC without impacting control tumor growth (Fig. 5B) and restored immune suppression with respect to intratumoral T-cell abundance (Fig. 5C) and induction of COX2 expression (Fig. 5D). As GW501516 acts systemically, we next specifically activated PPARo in PDAC cells by introducing a fusion of PPARo with the VP16 transactivation domain from herpes simplex virus (39), to enable ligand-independent activation, into control and sgGot2 PDAC cells (Supplementary Fig. S10A) at sufficiently low copy number to avoid detectable PPARS overexpression. Although VP16-PPAR8 increased neither proliferation in vitro (Fig. 5E) nor PDAC growth in the control



Figure 5. PPAR& activation restores tumor growth and T-cell exclusion in the absence of GOT2. A, Viable cell measurements in control or sgGot2 PDAC cells treated with vehicle or 100 nmol/L GW501516. RLU, relative light unit. B, PDAC tumor weight at the experimental endpoint, 30 days after orthotopic transplantation of the control or sgGot2 cells, with daily i.p. injection of vehicle or 4 mg/kg GWS01516. Ctrl: n = 5 per cohort, sgGot2: n = 4 per cohort. Data are presented as mean ± SEM. ns = not significant.*, P < 0.05; **, P < 0.01 by one-way ANOVA. C, IHC staining of control and sgGot2 688M tumors treated with vehicle or GW501516 as in B for the T-cell marker CD3. Representative images are shown above (scale bars = 50 µm), with quantification below (ctrl: n = 5, ctrl + GW501516: n = 5, sgGot2: n = 3, sgGot2 + GW501516: n = 4). Data are presented as mean ± SEM. ns = not significant. **, P < 0.01 by one-way ANOVA. D, IHC staining for PTGS2/COX2 in control or sgGot2 PDAC treated with vehicle or GW501516 (representative of n = 3-5 per cohort). Scale bars = 50 µm. E, Viable cell measurements in control or sgGot2 PDAC cells stably transduced with empty vector or VP16-PPAR6. Data are presented as mean ± SEM. F and G, PDAC tumor weight at the experimental endpoint in the indicated 688M (F) and FC1245 (G) lines. 688M: Ctrl: n = 5, sgGot2 a: n = 4, ctrl VP16-PPAR8: n = 4, sgGot2 a VP16-PPAR8: n = 4, endpoint = day 27. FC1245: Ctrl: n = 5, sgGot2 a: n = 5, ctrl VP16-PPAR8: n = 5, sgGot2 a VP16-PPAR8: n = 4, endpoint = day 18. Ctrl and sgGot2 FC1245 arms here are also depicted in Fig. 1E. ns = not significant. *, P < 0.05; **, P < 0.01; ***, P < 0.001; *** P < 0.0001 by one-way ANOVA. H, GPCR for PPARô-regulated genes in the indicated FC1245 stable cell lines, normalized to 36b4. Data are presented as mean \pm SEM from biological triplicates. *, P < 0.05; **, P < 0.01; ****, P < 0.001 by one-way ANOVA. **I**, PDAC tumor weight at experimental endpoint (day 18) in ctrl (n = 4) and shPpard (n = 5 per hairpin) FC1245 tumors. **, P < 0.01 by one-way ANOVA. **J**, Quantification of CD3 IHC on the tumors from **I** (scale bars = 50 µm). ns = not significant. *, P < 0.05 by one-way ANOVA. K, Heat map depicting differentially expressed (DE) genes in control and sgGot2 FC1245 PDAC cells, untreated or treated with 500 nmol/L GW501516 for 24 hours (n = 3 per group), identified by RNA-seq using cutoff criteria $P_{adi} < 0.01$ and logFC < -1 or > 1 in at least one comparison. (continued on next page)





group, genetic PPARδ activation significantly albeit partially rescued tumor growth in sgGot2 tumors in two independent models (Fig. 5F and G). Consistent with these findings, VP16-PPAR δ increased expression of target genes such as Ptgs2 in sgGot2 cells (Fig. 5H). PPAR8 knockdown in PDAC cells similarly reduced tumor growth and immune suppression (Fig. 5I and J; Supplementary Fig. S10B). Motivated by these results, we performed RNA-seq on control and sgGot2 PDAC cells treated with GW501516 to understand the transcriptional programs regulated by GOT2 and PPARo. Genes downregulated with loss of GOT2 were enriched for inflammatory mediators and for genes upregulated by PPARS agonist in macrophages in a previous study (40), whereas genes upregulated by PPARS agonist and downregulated with GOT2 loss within PDAC cells overlapped significantly and genes in the overlap were enriched for immune-modulatory genes (Fig. 5K-M; Supplementary Fig. S11A-S11C). Genes in the overlap included classic PPARδ target genes, such as *Hmgcs2*, as well as targets identified in this study, such as Reg3g. Together, these results indicate that GOT2 promotes PDAC progression and immune suppression by activating PPARδ.

DISCUSSION

Our study indicates that GOT2 plays a critical role in promoting a tumor-permissive immune microenvironment in the pancreas. This function is attributable at least in part to direct fatty acid binding and activation of nuclear receptor PPAR\delta. GOT2 loss did not affect the whole-cell abundance of any of the fatty acids measured, supporting a mechanism functionally downstream of fatty acid uptake. That said, analysis of our RNA-seq results for expression of fatty acid transporters (FATP1-6 encoded by SLC27a1-6, CD36) showed that Slc27a1 (encoding FATP1) was significantly downregulated with loss of GOT2, raising the possibility that uptake of other fatty acids may be indirectly affected by GOT2. The same transporter, Slc27a1, was significantly upregulated by PPARS agonism, consistent with transcriptional regulation by a GOT2-PPARS axis. Although diverse impacts on the tumor immune contexture resulted from perturbation of the GOT2-PPARS axis, the increased infiltration of cDC1 into tumors lacking cancer cell-intrinsic GOT2 was particularly promising in light of recent studies demonstrating the paucity of dendritic cells in PDAC as a cause of immune suppression (41, 42). Conversely, their increase in the GOT2-null setting is consistent with and may indeed enable antitumor immunity, which should be tested directly in future studies. Our results

agree with a recent study showing that dietary or pharmacologic activation of PPARS cooperates with oncogenic KRAS to drive pancreatic tumorigenesis by promoting immune-suppressive changes to the myeloid compartment (43). We note that our studies made use of orthotopic, syngeneic models of PDAC. Although these models have meaningful strengths relevant to human PDAC including an immune-suppressive phenotype and reflect the genetically engineered, spontaneous models from which they were derived with respect to the lack of T cell-mediated antitumor immune response in control mice, these models also present limitations. For example, these models do not progress through relevant stages of premalignant lesions, and our study leaves in question the significance of GOT2 in regulation of the immune microenvironment and disease progression in early neoplasia, which will best be addressed in spontaneous models of PDAC.

Further studies are needed to understand the mechanisms regulating GOT2 subcellular localization, the precise molecular mechanism by which GOT2 promotes PPAR8 transcriptional activity, and whether this functional interaction extends beyond pancreatic cancer cells. This last point is poised for investigation thanks to the recent development of a Got2-flox mouse strain (44). Motivated by such studies, analysis of GOT2 expression across tissues shows the highest expression in tissues known to be functionally regulated by PPARô, including skeletal muscle, heart, and liver (https://gtexportal.org/ home/). We speculate that the normal function of this GOT2-PPARo axis is to provide high levels of localized fatty acid ligand directly to constitutively nuclear PPARo via GOT2 fatty acid binding and transient nuclear shuttling into proximity of PPARδ, which we expect occurs at specific but not all PPARδ binding sites in light of our transcriptional profiling results. Investigating the therapeutic potential of targeting this GOT2-PPARδ axis is an appealing next step. With this goal in mind, we note that GOT2 has been subject to little characterization or functional study in immune cells directly. That said, analysis of publicly available RNA-seq data from human immune cell subsets showed that whereas GOT2 expression is low in most cell types examined, expression is appreciable in naïve CD4 and CD8 T cells (https://dice-database.org/). This gains potential significance in light of a recent study showing that exogenous GOT2 expression increased antitumor activity of CAR T cells in preclinical models, which was attributed to GOT2 metabolic function (45). Together with our findings, these results raise the possibility that therapeutic strategies parsing the enzymatic and fatty acid binding functions may be important. Although diverse mechanisms contribute to

immune evasion in PDAC (46), targeting GOT2 may be part of a potential treatment approach to foster an immune response against this deadly cancer.

METHODS

Animals

All experiments were reviewed, approved, and overseen by the institutional animal use and care committee at Oregon Health & Science University (OHSU) in accordance with NIH guidelines for the humane treatment of animals. C57BL/6J (000664, for models with FC1245; ref. 47) or B6129SF1/J (101043, for models with 688M; ref. 48) mice from The Jackson Laboratory were used for orthotopic transplant experiments at 8 to 10 weeks of age. Tissues from 6- or 12-month-old Kras^{LSL-G12D/+};Pdx1-Cre (KC) mice were kindly provided by Dr. Ellen Langer (OHSU).

Human Tissue Samples

Human patient PDAC tissue samples donated to the Oregon Pancreas Tissue Registry program (OPTR) in accordance with full ethical approval were kindly shared by Dr. Jason Link and Dr. Rosalie Sears (OHSU), and written informed consent from patients was received for all human tissue samples used in this study.

Plasmids

The pCMX-VP16-PPARD plasmid was kindly provided by Dr. Vihang Narkar (University of Texas Health Science Center at Houston; ref. 39). The VP16-PPARD element was cloned into the lentiviral vector. To construct pLenti VP16 PPARD, the VP16-PPARD element was amplified by PCR using sense primer 5'-GGGGACAAGTTTG TACA AAAAAGCAGGCTTAATGGCCCCCCGAC-3' and antisense primer 5'-GGGGACCACTTTGTACAAGAAAGCTGGGTTTTAGTAC ATGTCCTTGTAGATTTCCTGGAGCAGG-3'. PCR product was inserted into pDONR 221 entry clone using Gateway BP Clonase II enzyme (Thermo Fisher 12535029). Entry clone VP16 PPARD element was swapped into the expression region of pLenti CMV Puro DEST (Addgene #17452) using LR Clonase II enzyme (Thermo Fisher 11-791-020) to generate the pLenti VP16 PPARD construct. The pCMV3 plasmid containing C-terminal His-tagged human GOT2 cDNA was purchased from Sino Biological (HG14463-CH) and cloned into the lentiviral vector pLenti CMV Puro DEST (Addgene #17452) using the same approach as pLenti VP16 PPARD. pLenti wtGOT2 PCR product was generated using sense primer 5'-GGGGACAAGTTTGTACAAAA AAGCAGGCTTAATGGCCCTGCTGCACT-3' and antisense primer 5'-GGGGACCACTTTGTACAAGAAAGCTGGGTTTTTAGTGATGGT GGTGATGATGGTGG-3'. Triple-mutant GOT2 was constructed using the Q5 Site-Directed Mutagenesis Kit (New England Biolabs E0552S) in two subsequent steps. Two sets of primers were used to generate three site mutations: primer set 1 for K234A mutation (F: 5'-AACAGTGGTG GCGAAAAGGAATCTC-3'; R: 5'-GCTATTTCCTTCCACTGTTC-3') and primer set 2 for K296A and R303A mutations (F: 5'-GTCTG CGCAGATGCGGATGAAGCCAAAGCGGTAGA GTC-3'; R: 5'-CATA GTGAAGG CTCCTACACGC-3'). pLenti tmGOT2 was then generated using the same approach and primers as pLenti wtGOT2. pLenti aspartate transaminase mutant GOT2 (atamGOT2) was constructed in two consecutive steps using pLenti wtGOT2 as a template, with point mutations selected based on a human genetics study (1). The first step generated the R262A mutant using primers F: 5'-CTGTGGGCCACTTCATCGAA-3' and R: 5'-CCCAGGCATCCTTA TCACCATC-3'. The second step added the R337G mutant to the existing R262A mutant using primers F: 5'-CCAGATTTGG GAAAACAATGGC-3' and R: 5'-GGTGTTCAGAATGGCAGCA-3'. The pLenti NLS wtGOT2 construct was generated by inserting one cMYC NLS sequence into the C-terminus region using sense primer 5'-GTGAAACTGGATCTCGAGGGAGGCTCTCACCATC-3' and antisense primer 5'-TCTCTTAGCAGCAGGACCCCCCTTGGTGG-3'.

Cell Lines

Human pancreatic cancer cell lines MIA PaCa-2, PA-TU-8988T, Panc1, HPAF-II, and Capan-2 were obtained from ATCC and grown in Dulbecco's modified Eagle medium (DMEM) containing 10% FBS. Nontransformed, TERT-immortalized human pancreatic ductal epithelial cells were kindly provided by Dr. Rosalie Sears (OHSU; ref. 49). PA-TU-8988T cells harboring doxycycline-inducible shGOT2 were kindly provided by Dr. Costas Lyssiotis (University of Michigan). FC1245 PDAC cells were generated from a primary tumor in Kras^{LSL-G12D/+};Trp53^{LSL-R172H/+};Pdx1-Cre mice and were kindly provided by Dr. David Tuveson (Cold Spring Harbor Laboratory; ref. 47). 688M PDAC cells were generated from a liver metastasis in Kras^{LSL-G12D/+}; Trp53^{LSL-R172H/+};Pdx1-Cre;Rosa26^{LSL-tdTomato/+} mice and were kindly provided by Dr. Monte Winslow (Stanford University School of Medicine; ref. 48). Cell lines were routinely tested for Mycoplasma at least monthly (MycoAlert Detection Kit, Lonza).

The pSpCas9(BB)-2A-Puro(PX459) v2.0 plasmid (Addgene #62988) was used to clone guide sequences targeting Got2 per the supplier's protocol: sgRNA A: GACGCGGGTCCACGCCGGT, sgRNA B: ACG CGGGTCCACGCCGGTG. The 688M or FC1245 cell line was transfected with a control plasmid or plasmid containing either of the sgGot2 sequences and subject to selection with 2 µg/mL puromycin for 4 days. Single-cell clones were expanded and screened for GOT2 protein expression by Western blot.

GOT2 shRNA vectors were purchased in bacterial glycerol stocks from Sigma-Aldrich Mission shRNA (mouse shRNA A: TRCN0000325948, shRNA B: TRCN00000325946; human sh24: TRCN0000034824, sh27: TRCN0000034827), as were PPARD shRNA vectors (shRNA 6001: TRCN0000026001, shRNA 6007: TRCN0000026007). Briefly, bacterial cultures were amplified in an ampicillin growth medium from glycerol stocks for use in the purification of plasmid DNA. Subsequently, the purified plasmid was transfected to packaging cells HEK293T for the production of lentiviral particles. FC1245 cells were then infected and puromycin selected to generate stable GOT2 or PPARD knockdowns, with validation by Western blot. Lentivirus preparation for stable cell line generation was done with pMD2.G envelope plasmid (Addgene #12259) and psPAX2 packaging plasmid (Addgene #12260) in 293T-LentiX cells. Briefly, 5 µg of pMD2.G, 5 µg of psPAX2, and 10 µg of plasmid DNA (shGOT2 KD, shPPARD KD, VP16-PPARdelta, wtGOT2, tmGOT2, NLS-wtGOT2, atamGOT2, or scramble ctrl) were combined with 600 µL Opti-MEM and 20 µL lipofectamine 2000 for 20 minutes at room temperature. Dishes (10 cm) of 293T-LentiX were kept in 0% FBS DMEM, and the mixture was added in a dropwise manner. Twelve hours later, media were changed to 10% FBS DMEM. At 24 and 48 hours after transduction, media were collected and filtered through a 0.25-µm filter, aliquoted, and frozen at -80°C. For lentiviral transduction of human and mouse cell lines, cells were plated to 6-well plates. Polybrene (10 µg/mL; EMD Millipore TR-1003-G) was added to 1 mL 10% FBS DMEM and 300 µL of filtered lentivirus media. Twenty-four hours later, media were changed to a fresh 10% FBS DMEM. Fortyeight hours after initial transduction, cells were treated with 2 μ g/mL puromycin (Thermo Fisher A1113803) or 4 μ g/mL puromycin depending on the cell line. A control well of nontransduced cells was used as an indicator for proper selection. Protein knockdown was validated by Western blot. For transaminase inhibition in vitro, AOA (Millipore Sigma; C13408) was used at 1 mmol/L as previously described (50).

Western Blotting

PDAC cells were treated as described in the text, and whole-cell lysates were prepared in RIPA buffer containing protease inhibitor cocktail (Sigma-Aldrich 11836170001). Alternatively, subcellular fractions were prepared using the Cell Fractionation Kit #9038 purchased from Cell Signaling Technology following the manufacturer's instructions. Briefly, cells were collected with scraping, washed in PBS, and



pelleted (350 \times g 5 minutes). Cells were resuspended in 500 µL PBS and 100 µL reserved for whole-cell lysis in RIPA buffer + cOmplete mini EDTA-free protease inhibitor cocktail. The remaining cell pellet was centrifuged (500 $\times\,g$ 5 minutes), PBS was decanted, and 500 μL CIB + 5 μ L protease inhibitor and 2.5 μ L PMSF was added. Solutions were vortexed and stored on ice for 5 minutes. Lysates were centrifuged $(500 \times g 5 \text{ minutes})$; the supernatant was collected as the cytosolic fraction. The remaining insoluble pellet was washed with CIB, and the supernatant was decanted. MIB (500 μ L) + 5 μ L protease inhibitor and 2.5 µL PMSF was then added to the cell pellet. After vortexing for 15 seconds, solutions were incubated on ice for 5 minutes and centrifuged (8,000 \times g 5 minutes). The supernatant was collected as the membrane and organelle fraction. Pellet was then washed in MIB, and the supernatant was decanted. CyNIB (250 μ L) + 2.5 μ L protease inhibitor + 1.25 µL PMSF was then added to the pellet containing nuclei. The solution was sonicated for 5 seconds at 20% power $3 \times$ to prepare nuclear lysate. For Western blot, 60 µL 4× LDS loading buffer with 10× reducing agent was added for every 100 µL of supernatant per fraction. Samples were boiled for 5 minutes at 95°C and centrifuged for 3 minutes at 15,000 \times g; 15 µL of each fraction along with 15 µL of the whole-cell lysate was loaded for Western blotting. Alternatively, to generate total nuclear and cytosolic fractions, NE-PER Nuclear and Cytoplasmic Extraction Reagent Kit (Thermo Fisher) was used according to the manufacturer's protocol. Where indicated, His-tagged GOT2 protein was immunoprecipitated using the His-tag isolation and pulldown Dynabeads system (Thermo Fisher) using the manufacturer's protocol. Protein concentration was quantitated using the BCA Protein Assay Kit (Pierce). Equal amounts of protein were loaded in each lane and separated on a 4% to 12% Bis-Tris NuPAGE gel (Invitrogen), and then transferred onto a PVDF membrane. Membranes were probed with primary antibodies and infrared secondary antibodies: GOT2 (Thermo Fisher PA5-77990), β-Actin (Santa Cruz sc-47778), PPAR& (Abcam ab178866), His (R&D Systems MAB050-100), COX2 (Abcam ab15191), COX IV (Cell Signaling Technology 11967S), AIF (Abcam ab1998), PMCA1 (Novus Biologicals 5F10), Lamin A/C (Cell Signaling Technology 4777S), Tom20 D8T4N (Cell Signaling Technology 42606S), HSC 70 (Santa Cruz sc-7298), Cas9 (Novus Biologicals NBP2-36440SS), GOT1 (Sigma-Aldrich AV48205), anti-rabbit Alexa Fluor Plus 680 (Thermo Fisher A32734), and anti-mouse Alexa Flour Plus 800 (Invitrogen A32730). Protein bands were detected using the Odyssey CLx infrared imaging system (LI-COR Biosciences).

Immunofluorescence

Cells plated on coverslips were fixed in 10% neutral buffered formalin for 10 minutes at room temperature, washed three times with PBS, and permeabilized with 0.1% Triton X-100 for 10 minutes at room temperature. When MitoTracker staining was performed, cells plated on coverslips were stained with 100 nmol/L MitoTracker (Thermo Fisher M22462) at 37°C for 15 minutes prior to fixation. Following permeabilization, coverslips were blocked for 1 hour at room temperature in blocking solution (Aqua block buffer, Abcam ab166952) and then transferred to a carrier solution (Aqua block) containing diluted primary antibodies: GOT2 (Sigma-Aldrich HPA018139), COX IV (Cell Signaling Technology 11967S), COX2 (Abcam ab15191), and His (R&D Systems MAB050-100). Coverslips were incubated with the primary antibody at 4°C overnight and then washed five times for 5 minutes each in PBS, following which secondary Alexa Flour-conjugated antibodies diluted in the same carrier solution (1:400) were added to the coverslips for 1 hour at room temperature. After the secondary antibody incubation, coverslips were washed five times for 5 minutes each in PBS and mounted with Vectashield mounting media containing DAPI (Vector Laboratories; H-1500). Images were captured on a Zeiss LSM 880 laser-scanning inverted confocal microscope at the OHSU Advanced Light Microscopy Shared Resource, and a 40×/1.1 NA water objective or 63×/1.4 NA oil objective was used to image the samples.

IHC

Mice were anesthetized and euthanized according to institutional guidelines. Pancreatic tumors were excised carefully and fixed overnight in 10% phosphate-buffered formalin. Tissue samples were paraffin-embedded and sectioned by the OHSU Histopathology Shared Resource. Human PDAC tissue sections from formalin-fixed, paraffinembedded blocks were obtained from the OPTR. In brief, tissue sections were deparaffinized and rehydrated through an ethanol series and ultimately in PBS. Following antigen retrieval, tissue samples were blocked for 1 hour at room temperature in a blocking solution (8% BSA solution) and then transferred to a carrier solution (8% BSA solution) containing diluted antibodies: GOT2 (Sigma-Aldrich HPA018139), COX IV (Cell Signaling Technology 11967S), COX2 (Abcam ab15191), CD3 (Abcam ab5690), CD4 D7D2Z (Cell Signaling Technology 25229S), CD8 (Abcam ab203035), F4/80 (Cell Signaling Technology 70076T), and Arginase-1 (ARG1; Sigma-Aldrich ABS535). Sections were incubated overnight at 4°C and then washed five times for 5 minutes each in PBS. For fluorescence imaging, secondary Alexa Flour-conjugated antibodies diluted in the same carrier solution (1:400) were added to the sections for 1 hour at room temperature. Sections were then washed five times for 5 minutes each in PBS and were mounted with Vectashield mounting media containing DAPI. For DAB chromogen imaging, sections were stained with primary antibody as described above, then the samples were incubated in polymeric horseradish peroxidase (HRP)-conjugated secondary antibody (Leica PV6121) for 1 hour, followed by five 5-minute 1× TBST washes. HRP was detected using DAB chromogen (3,3'-diaminobenzidine) solution (BioCare Medical BDB2004) prepared per the manufacturer's instructions. Tissues were exposed to chromogen solution until a brown precipitate was detected produced from oxidized DAB where the secondary poly-HRP antibody is located. As soon as DAB chromogen was detected, tissue slides were washed in diH₂O, counterstained in hematoxylin, dehydrated, and cleared for mounting. Stained tissue sections were scanned on a Leica Biosystems Ariol digital fluorescence scanner or Leica Biosystems Aperio brightfield digital scanner. Quantification was performed for single stains using QuPath quantitative pathology and bioimage analysis software v0.2.3. For costains (CD8/ GRZB and F4/80/ARG1), manual counting was performed on at least 10 high-powered fields per tumor sample.

Multiplex IHC

Sequential IHC was performed on 5-µm formalin-fixed, paraffinembedded sections as previously described (51, 52). Briefly, slides were deparaffinized, which was followed by heat-mediated antigen retrieval in pH 6.0 Citra solution (BioGenex HK086) followed by blocking in Dako Dual Endogenous Block (Dako S2003) and then 10 minutes of protein blocking with 5% normal goat serum and 2.5% BSA in TBST. Primary antibodies were applied in sequential order as listed in Box 1. After washing off primary antibody in TBST, either anti-rat (414311F), anti-mouse (414131F), or anti-rabbit (414141F) Histofine Simple Stain MAX PO HRP-conjugated polymer (Nichirei Biosciences) was applied for 30 minutes at room temperature, followed by AEC chromogen (Vector Laboratories SK-4200). Whole-slide digital imaging was performed following each chromogen development. Heat and chemical stripping between cycles and rounds was performed as previously described (51, 52). DNA was stained with hematoxylin (Dako S330130-2) for the purposes of image computation.

Regions of interest were selected, and then images were coregistered in MATLAB version R2018b using the SURF algorithm in the Computer Vision Toolbox (The MathWorks, Inc.). Image processing and cell quantification were performed using FIJI (53) and CellProfiler Version 3.5.1 (54). AEC signal was extracted for quantification and visualization in FIJI using a custom macro for color deconvolution. Briefly, the FIJI plugin Color_Deconvolution [H AEC] was used to separate hematoxylin, followed by postprocessing steps for signal cleaning and

BOX 1: ANTIBO	DIES USED FOR	RMULTIPLEX	10			
	Ly	ymphoid panel:				
	Cycle 1	Cycle 2	Cycle 3	Cycle 4		
- .	Round 1	Round 1	Round 1	Round 1		
larget				<u>CD3</u>		
Company/Product	CSI (96615)	CSI (84651)	CST (2203S)	(RM-9107-S)		
Clone	Polyclonal	D7D5W	C63D9	SP7		
Dilution	1/100 DT 1 br	1/100	1/200.	1/300 DT 1 br		
Secondary species	Rabbit	Rabbit	Rabbit	Rabbit		
AEC	1 h 5 min	1 h 20 min	15 min	53 min		
Target	Round 2 Hematoxylin	Round 2 CD45	Round 2 CD8			
Company/Product	Dako (S330130-2)	BD Bio (550539)	eBioscience			
			(14-0808-82)			
Clone	NA Ready to use	30-F11	4SM15			
Duration	1 min	1/50. RT, 1 hr	0N, 4°C			
Secondary species	NA	Rat	Rat			
AEC	NA	52 min	30 min			
	Cycle 5	Cycle 6	Cycle 7	Cycle 8		
- .	Round 1	Round 1	Round 1	Round 1		
Company/Droduct	PD-L1	(1222E7)	Granzyme B	Ki-67		
Company/Product	(15004)	ADCall (155557)	ADCall (4059)	(15580)		
Clone	E1L3N	EPR1334	Polyclonal	Poylclonal		
Dilution	1/50 ON 4°C	1/3,000 RT 1 br	1/200 ON	1/1,000 RT 1 br		
Secondary species	Rabbit	Rabbit	Rabbit	Rabbit		
AEC	23 min	17 min	10 min	24 min		
				Round 2		
Target				panCK		
Company/Product				Abcam		
Clone				AE1/1E3		
Dilution				1/100		
Duration				RT, 1 hr Mouse		
AEC				22 min		
				Round 3		
Target				Hematoxylin		
Company/Product				рако (5330130-2)		
Clone				NA		
Dilution				Ready to use		
Duration Secondary species				1 min		
AEC				NA		
Myeloid panel:						
	Cycle 1	Cycle 2	Cycle 3	Cycle 4		
_	Round 1	Round 1	Round 1	Round 1		
larget	F4/80	CSFIR	CD11c	CD206		
Company/Product	Serotec CI-A3-1	Santa Cruz F2412	CST (97585) D1V9Y	Abcam (64693) Polyclonal		
Dilution	1/200	1/1,000	1/100	1/2,000		
Duration	RT, 1 hr	RT, 1 hr	RT, 1 hr	RT, 1 hr		
Secondary species	Rat 22 min	Raddit 16 min	Raddit 15 min	Kabbit 43 min		

(continued)



Target			Round 2 CD45	Round 2 MHCII
Company/Product			BD Bio (550539)	eBioscience
Clone Dilution Duration Secondary species AEC			30-F11 1/100 ON, 4°C Rat 30 min	(eBi14-532) M5/144.15.2 1/200 ON, 4°C Rat 63 min
	Cycle 5	Cycle 6	Cycle 7	
	Round 1	Round 1	Round 1	
Target	PD-L1	Ki-67	CD11b	
Company/Product	CST (13684)	Abcam (15580)	Abcam (133357)	
Clone	E1L3N	Polyclonal	EPR1334	
Dilution	1/50	1/1,000	1/6,000	
Duration	, RT, 1 hr	RT, 1 hr	RT, 1 hr	
Secondary species	Rabbit	Rabbit	Rabbit	
AEC	23 min	24 min	8 min	
	Round 2	Round 2	Round 2	
Target	Ly6G	panCK	Hematoxylin	
Company/Product	eBioscience (551459)	Abcam (ab27988)	Dako (S330130-2)	
Clone	1A8	AE1/1E3	NA	
Dilution	1/200	1/100	Ready to use	
Duration	ON, 4°C	RT, 1 hr	1 min	
Secondary species	Rat	Mouse	NA	
AEC	6 min	22 min	NA	

BOX 1: ANTIBODIES USED FOR MULTIPLEX IHC (*CONTINUED***)**

Abbreviations: BD Bio, BD Biosciences; CST, Cell Signaling Technology; NA, not applicable; ON, overnight; RT, room temperature.

background elimination. AEC signal was extracted in FIJI using the NIH plugin RGB_to_CMYK. Color-deconvoluted images were processed in CellProfiler to quantify single-cell mean intensity signal measurements for every stained marker. CellProfiler outputs were loaded into FCS Express 7 Image Cytometry RUO (De Novo Software) software, and hierarchical gating was carried out to classify immune cell populations.

Proliferation Assays

PDAC cells were seeded into 96-well plates at 2×10^3 cells per well in DMEM containing 10% FBS. Cells were treated as indicated in the text with 100 nmol/L GW501516 (Cayman Chemical 10004272) at the time of cell seeding or 5 mg/mL doxycycline (Sigma-Aldrich D9891) 48 hours prior to cell seeding. GW501516 and doxycycline treatments were both replenished every 48 hours for extended time points. After 72 hours or at the time points indicated in the text, cells were lysed with the CellTiter-Glo Luminescent Cell Viability Assay reagent (Promega), and luminescence was read using a GloMax plate reader.

ChIP

ChIP was performed as described previously (55). Briefly, PDAC cells were fixed in 1% formaldehyde, and nuclei were isolated and lysed in buffer containing 1% SDS, 10 mmol/L EDTA, 50 mmol/L Tris-HCl pH 8.0, and protease inhibitors and sheared with a Diagenode Bioruptor to chromatin fragment sizes of 200 to 1,000 base pairs. Chromatin was immunoprecipitated with antibodies to PPARδ (Abcam ab178866) or acetylated histone H3K9 (Cell Signaling Technology 9649). PPARδ binding or histone acetylation at known PPARδ target gene promoter regions was assessed by ChIP-qPCR, and enrichment values were normalized to a control intergenic region of the genome.

The following primer sequences were used: Intergenic F: tggtgcttcttggtcaatca, R: aggacaaaacagcaaccaaca, *Angptl4* F: tcagcctaccagggaggagaa, R: ggaggaaagggcgtacaaat, *Reg3 g* F: actgcacccatacctgacaa, R: ccctaggatggtgtgtcagg, *Ptgs2* F: gttcttgcgcaactcactga, R: agtgctgagttccttcgtga, *Csf1* F: gaaggctgctcatcccattg, R: ggggcctaaaacatgtgcaa, and *Cxcl2* F: gcttgaacacacacacacct, R: ctatgggaccctgggatgtc.

Gene Expression Analysis by qPCR

The isolated total RNA (1 µg) was reverse-transcribed to produce cDNA using the iScript Reverse Transcription Supermix kit (Bio-Rad). Real-time PCR was performed using SYBR Green supermix (Bio-Rad). The cDNA sequences for specific gene targets were obtained from the human genome assembly (http://genome.ucsc.edu), and gene-specific primer pairs were designed using the Primer3 program (https://primer3.ut.ee/). Relative gene expression was normalized using the 36B4 housekeeping gene. The following primer sequences were used: human and mouse 36B4 (RPLP0): F: 5'-GTGCT GATGGGCAAGAAC-3'; R: 5"-AGGTCCTCCTTGGTGAAC-3"; mouse Ptgs2: F: 5'-TGAGTGGGGTGATGAGCAAC-3'; R: 5'-TTCAGAGGCAAT GCGGTTCT-3'; mouse Pdk4: F: 5'-TGAACACTCCTTCGGTGCAG-3'; R: 5'-GTCCACTGTGCAGGTGTCTT-3'; mouse Csf1: F: 5'-ATGAG CAGGAGTATTGCCAAGG-3', R:5'-TCCATTCCCAATCATGTGGCTA-3'; and mouse Reg3g: F: 5'-ATGCTTCCCCGTATAACCATCA-3', R: 5'-GGCCATATCTGCATCATACCAG-3'.

Analysis of GOT2 Expression and PPAR δ Activity in TCGA-PAAD

To assess the correlation between GOT2 and PPAR δ activity in a larger compendium of human PDAC samples, we downloaded the

pancreatic cancer RNA-seq fragments per kilobase million gene expression data from TCGA-PAAD using the Bioconductor (56) R package (version 2.15.3). We built the PPAR δ target gene signature using the PPARô-regulated genes defined in a previous study (40). Then, we used the gene set variation analysis (GSVA) algorithm (57) with the default settings, as implemented in the GSVA R package (version 1.34.0), to calculate the enrichment of PPARS target genes as PPARS activity score for each sample and plotted this against GOT2 expression.

Metascape Analysis

The PDAC TCGA Firehose Legacy database provides mRNA expression data for coexpression analysis accessible through cBioPortal. The data set includes Spearman correlation analysis and P values for each gene comparison. The data set was used to identify genes negatively or positively correlated with GOT2 expression in patients with PDAC. A list of genes with a Spearman correlation value of equal to or less than -0.25 (negative correlation) or equal to or greater than 0.25 (positive correlation) and a P value of less than 0.01 was generated. The list of genes was submitted to the online bioinformatics tool Metascape for the identification of enriched gene ontology clusters in the data set. The output from the Metascape analysis was graphed using GraphPad Prism.

Cytokine Array

Control and sgGOT2 FC1245 cells were respectively seeded into a 6-well plate at 5×10^5 cells per well in growth medium (DMEM with 10% FBS and 1% penicillin and streptomycin). The next day, the growth medium was removed and replaced with 1 mL of serum-free DMEM. The supernatant was collected 48 hours after media change and spun down at $1,500 \times g$ for 10 minutes before proceeding to cytokine array using the Proteome Profiler Mouse XL Cytokine Array Kit (R&D ARY028). The assay was completed as per the manufacturer's instructions. Membranes were blocked for 1 hour at room temperature using Array Buffer 6, followed by overnight incubation of supernatant samples at 4°C. After overnight incubation, blots were washed three times for 10 minutes per wash using 1× wash buffer. Diluted detection antibody cocktail (1.5 mL) was then added to the membranes and incubated for 1 hour on a shaker at room temperature. Wash steps were repeated similarly to prior wash steps. Membranes were then incubated with 2 mL of diluted IRDye 800CW Streptavidin (Li-COR Biosciences 926-32230) on a shaking platform for 30 minutes at room temperature. Wash steps were repeated after streptavidin incubation, and membranes were read using the Odyssey CLx infrared imaging system (Li-COR Biosciences).

Orthotopic PDAC Modeling

The orthotopic transplant method used here was described previously (58). In brief, 8- to 10-week-old, wild-type male C57BL/6J (for FC1245) or B6129SF1/J (for 688M) mice were orthotopically transplanted as described previously with 5×10^3 FC1245 cells or 8×10^4 688M cells in 50% Matrigel (Corning 356231) and 50% DMEM. For experiments with 688M cells harboring VP16-PPARD, 6×10^4 688M cells were used. For pharmacologic activation of PPARo, mice were treated with vehicle (5% PEG-400, 5% Tween-80 in diH2O) or with 4 mg/kg GW501516 in vehicle by i.p. injection once daily. Vehicle was created and autoclaved before use. GW501516 was created in 10 mmol/L stock in DMSO and stored in 250 µL aliquots at -20°C (one for each day of treatment). On the day of treatment, a vial was thawed and diluted 1:10 in vehicle, and mice were dosed at 4 mg/kg. For T-cell depletion experiments, mice received an i.p. injection of 0.2 mg of \alphaCD8 (2.43), aCD4 (GK1.5), or an IgG2b isotype control (LTF-2) diluted in 100 µL sterile PBS. Antibodies were purchased from Bio X Cell and were administered beginning 2 days preimplantation with 6×10^4 688M cells and every 4 days thereafter until euthanasia, as previously described (23). Mice were euthanized when control animals were moribund, and tumors were excised, weighed, and immediately fixed in formalin.

Long-Chain Fatty Acid Binding Site Prediction

The arachidonic acid binding site on the human GOT2 surface is predicted using the molecular modeling technique. Druggable hotspot identification has long been used to predict and explore the allosteric pockets that accommodate substrate and drug-like molecules (59, 60). A similar approach was taken to identify a plausible arachidonic acid binding site by probing the GOT2 3D protein structure (ref. 36; PDBID:5AX8). The protein structure was prepared using the protein prep tool of Maestro-2014-3 (Schrödinger, LLC). Arachidonic acid is a 20-carbon long-chain fatty acid (LCFA) with greasy carbons and a carboxylate group. The available structural information suggests that the binding pocket must be hydrophobic with the positively charged residues to accommodate LCFA (61, 62).

The SiteMap (63) calculation accounts for the prediction of pockets, characterized by cavity volume, chemical, and physical properties as that of known druggable sites. Five sites were predicted on the GOT2 structure, and these sites had a site score of >0.8, composed of hydrophobic, hydrogen bond acceptor, and donor volumes. The top-ranked site 1 is a catalytic site, and sites 2 to 5 are allosteric. Arachidonic acid docked against all the predicted sites. The Induced-Fit docking protocol (64) adopted here allows both the ligand and the surrounding residues of protein to be flexible. A total of five docking runs were performed on the predicted site. The docking grid boxes were defined based on the residues suggested by the SiteMap analysis (site 1: N215, H210; site 2: N270, F239; site 3: A260, W226, H373, G385, Q390; site 4: R337, G254; site 5: N332, D93). The site 2 ~25 Å away from the catalytic site resulted in a binding pose with favorable energy and interaction complementarity between the protein and ligand. Compared with other sites, site 2 has increased hydrophobic volume, which may recognize LCFA-like arachidonic acid. Triple mutants K234A/K296/R303 were proposed to validate the predicted binding pose. K234 interacts with the carboxylate group of LCFA. K296, which is in proximity to making ionic interaction (in dynamics) and perturbation of the positive charge to neutral alanine residues, prevents the charged interaction. From the docking pose, R303 is making the hydrophobic interaction with the lipid tail of arachidonic acid. R303A mutation reduces the hydrophobic interaction by the side chain of arginine. The proposed triple mutations have the potential to abolish the arachidonic acid binding.

Fatty Acid Binding Assay

Reactions were carried out in binding buffer (0.003% digitonin in 1× PBS) containing 1 µmol/L of purified human GOT2 protein (AA30-430) and 0.5 µci/mL [3H]-arachidonic acid. After incubation for 1 hour at 4°C, the mixture was incubated with preequilibrated TALON Metal Affinity Resin (Takara, 635502) at 4°C for 1 hour and loaded onto a column and washed with binding buffer, then binding buffer with 0.01% BSA, and then binding buffer again. The protein-bound [3H]-arachidonic was eluted with elution buffer (50 mmol/L sodium phosphate, 300 mmol/L sodium chloride, 150 mmol/L imidazole; pH 7.4.) and quantified by scintillation counting. For competition experiments with unlabeled lipids, the assays were carried out in the presence of ethanol containing the indicated unlabeled sterol (0-1 mmol/L).

Luciferase Assay

The PPRE x3-TK-Luc (PPAR response element driving luciferase) plasmid #1015 was purchased from Addgene, and the Renilla plasmid (pRL-SV40) was generously provided by Dr. Ellen Langer (OHSU). Cells were transfected with 2.5 µg PPRE x3-TK-Luc, 15 ng pRL-SV40, and 4 µL Lipofectamine 2000 in 6-well plates. Briefly, cells were plated at 1×10^6 per well of a 6-well plate and allowed to adhere overnight. Plasmids were combined in 150 µL Opti-MEM, whereas Lipofectamine 2000 was combined in a separate tube with 150 μ L Opti-MEM. After 5 minutes, the tubes were combined. The mixture (300 µL) was added, in a dropwise manner, to 700 µL of Opti-MEM on each well for transfection. The cells were incubated overnight at 37°C, collected, counted, and replated to white-walled, 96-well plates in triplicates. Twenty-four hours later, a dual-luciferase assay was completed following the manufacturer's instructions: Dual-Luciferase Reporter Assay System (Promega E1910). Briefly, cells were lysed in white-walled, 96-well plates with 20 μ L 1× Passive Lysis Buffer and shaken on a room temperature shaker. LARII (100 μ L) was added to each well, and luminescence was measured over 5 seconds. Stop and Glo (100 μ L) was then added, and Renilla activity was measured with luminescence over 5 seconds. Activity was calculated by normalizing the luciferase signal to Renilla for each well.

PPAR δ Transcription Factor Activity Assay

Nuclear lysates were prepared using a detergent-free fractionation protocol. Cells were scraped and collected from 10-cm dishes, washed with PBS, pelleted ($450 \times g5$ minutes), resuspended in PBS, and 1/5 of the volume was reserved for whole-cell lysis in RIPA (Amresco N653-100 mL) + cOmplete EDTA-free protease inhibitor cocktail (Sigma-Aldrich 11836170001). The remaining 4/5 of cell suspension was centrifuged ($450 \times g 5$ minutes), PBS was removed, and cells were lysed on ice for 15 minutes in lysis buffer (5× of cell pellet volume). Lysis buffer consisted of 10 mmol/L HEPES pH 7.9, 1.5 mmol/L MgCl2, and 10 mmol/L KCl with 1 mmol/L DTT and EDTA-Free cOmplete mini protease inhibitor cocktail. Lysates were centrifuged ($450 \times g 5$ minutes), supernatant was decanted, lysis buffer was added (2× cell volume), and suspensions were ground on ice with a plastic homogenizer 10× in 1.5 mL Eppendorf tubes. Lysates were centrifuged (10,000 × g 20 minutes), and supernatant was collected as a cytosolic fraction. The remaining pellet was washed with 200 µL lysis buffer (10,000 \times g 5 minutes), the supernatant was decanted, and extraction buffer was added (2/3× cell pellet volume). Extraction buffer consisted of 20 mmol/L HEPES pH 7.9, 1.5 mmol/L MgCl2, 0.42M NaCl, 0.2 µmol/L EDTA, 25% glycerol (V/V), and 1 mmol/L DTT and cOmplete mini EDTA-free protease inhibitor cocktail. Nuclei were ground with a plastic homogenizer in 1.5 mL Eppendorf tubes 20× and incubated at 4°C with gentle shaking for 10 minutes. Samples were centrifuged (20,000 \times g 5 minutes), and the supernatant was transferred to cold Eppendorf tubes as a nuclear fraction. Lysates were measured with BCA, and an equal protein amount was added per sample for each well. The manufacturer's instructions were followed for the PPARo transcription factor kit (Abcam ab133106). Briefly, CTFB was prepared and added to blank and nonspecific binding wells, nuclear lysates were added to each sample well containing immobilized PPRE-containing DNA, and the plate was incubated overnight at 4°C without agitation. The next day, wells were washed 5× in 1× wash buffer and incubated in PPAR8 primary antibody (1:100) for 1 hour at room temperature in the dark, without agitation. Wells were washed 5× in 1× wash buffer and incubated in goat anti-rabbit HRP conjugate (1:100) for 1 hour at room temperature in the dark without agitation. Wells were washed 5× in 1× wash buffer, and 100 µL developing solution was added to each well. The plate was incubated for 15 to 45 minutes on a room temperature shaker, in the dark, until color developed. Stop solution (100 μ L) was added to the wells, and the absorbance at 450 nm was taken.

RNA-seq

Control or sgGot2 FC1245 PDAC cells were untreated or treated with 500 nmol/L GW501516 for 24 hours, and RNA was isolated by TRIzol extraction per the manufacturer's instructions and cleaned up using an RNeasy kit (Qiagen). Treatments were performed in triplicate. RNA-seq libraries were prepared using the Illumina TruSeq Stranded mRNA kit using 300-ng total RNA as input and 15 rounds of amplification. Libraries were sequenced on an Illumina NextSeq 500 (~30 million reads per sample). The quality of the raw sequencing files was evaluated using FastQC (https://www.bioinformatics.babraham.ac.uk/projects/

fastqc/) combined with MultiQC (ref. 65; http://multiqc.info/). Trimmomatic (66) was used to remove any remaining Illumina adapters. Reads were aligned to Ensembl's GRCm38 along with its corresponding annotation, release 104. The program STAR (ref. 67; v2.7.3a) was used to align the reads to the genome. STAR has been shown to perform well compared with other RNA-seq aligners (68). Since STAR utilizes the gene annotation file, it also calculated the number of reads aligned to each gene. RNA-SeQC (69) and another round of MultiQC were utilized to ensure alignments were of sufficient quality. Gene-level differential expression analysis was performed in open source software R (R Core Team). Gene-level raw counts were filtered to remove genes with extremely low counts in many samples following the published guidelines (70), normalized using the trimmed mean of M-values (71), and transformed to log-counts per million with associated observational precision weights using the voom (72) method. Gene-wise linear models for control versus knocked out for Got2 (sgGot2) were used for differential expression analyses using limma with empirical Bayes moderation (73) and false discovery rate adjustment. Gene set enrichment analysis using fgsea was used to compare differentially expressed genes from control and sgGot2 PDAC cells to PPARo target genes defined in a previous study (40). Gene ontology and pathway analyses were performed using Ingenuity Pathway Analysis or Enrichr. Two-way comparison (control versus sgGot2) and four-way comparison (control versus sgGot2 \pm GW501516) were performed with independent sgGot2 clones.

Nuclear Fatty Acid Uptake Assay

MIA PaCa-2 ctrl and sh27 cells were plated at 5×10^5 in a 6-well plate and allowed to adhere overnight. The media were changed to 0% FBS DMEM, and the cells were incubated for 24 hours. The media was changed to 0.5% fatty acid-free BSA DMEM with either chloroform (ctrl) or 2.5 µmol/L NBD-arachidonic acid (Avanti Polar Lipids 810106C). Media were made before being added to cells, heated to 37°C, and vortexed until fatty acid was completely in solution. Cells were incubated at 37°C for durations indicated in the manuscript and collected and fractionated using the detergent-free method described above (PPARo transcription factor activity assay). Nuclear lysates were placed in a white-walled, 96-well plate, and fluorescence was measured at 480 nm excitation and 540 nm emission. Lysate concentration was measured using a BCA kit. FC1245 cells were plated at 5×10^5 per well and treated as described above, but treatment was reduced to 2 µmol/L NBD-arachidonic acid for 15 minutes due to lipid toxicity in this cell line.

Aspartate Aminotransferase Assay

The AST Activity Assay Kit (Sigma-Aldrich MAK055) was used to determine aspartate aminotransferase activity per the manufacturer's instructions. Briefly, this assay determines the transfer of an amino group from aspartate to alpha-ketoglutarate in the generation of glutamate, which produces a colorimetric product (450 nm) that is proportional to aspartate aminotransferase activity in the sample. For this assay, PA-TU-8988T cells with stable expression of doxycyclineinducible GOT2 shRNA were transiently transfected with wtGOT2 and tmGOT2. After 48 hours, these cells were exposed to doxycycline for 48 hours to knock down endogenous GOT2 in cells with GOT2 shRNA. Cells were seeded at 5×10^6 and collected via trypsin disassociation after cells were adhered. The cells were then resuspended in 1 mL of ice-cold 1× PBS and 200 μ L (1 × 10⁶ cells) and were collected for the AST assay, and 800 μ L (4 × 10⁶ cells) were collected for protein concentration estimation and Western blot protein expression analysis. Using AST assay kit buffers, cells were lysed to obtain a supernatant that was combined with the kit reagent master mix to detect glutamate in a colorimetric reaction. The samples were read every 5 minutes for 30 minutes. AST activity and concentration in the samples were determined using instructions from the manufacturer.

Free Fatty Acid Measurements

Samples were subjected to an LC-MS analysis to detect and quantify levels of free fatty acids in sample extracts. A fatty acid extraction was carried out on each sample using 100% methanol as the homogenization solvent. Whole-cell pellets (1 \times 10⁶ cells/sample) were lysed with 1,000 µL of methanol and ~100 µL of zircon beads (0.5 mm). Manual disruption with a p1000 pipette tip was performed to assist with initial pellet suspension in the extraction buffer. The methanol extracts were centrifuged (21,000 $g \times 3$ minutes) and transferred to glass LC-MS inserts for analysis. The LC column was a WatersTM BEH-C18 $(2.1 \times 100 \text{ mm}, 1.7 \,\mu\text{m})$ coupled to a Dionex Ultimate 3000 system, and the column oven temperature was set to 25°C for the gradient elution. The flow rate was 0.1 mL/minute and used the following buffers: (A) water with 0.1% formic acid and (B) acetonitrile with 0.1% formic acid. The gradient profile was as follows: 60% to 99% B from 0 to 6 minutes, hold at 99% B from 6 to 10 minutes, 99% to 60% B from 10 to 11 minutes, hold at 60% B from 11 to 15 minutes. Injection volume was set to 1 µL for all analyses (15 minutes of total run time per injection).

MS analyses were carried out by coupling the LC system to a Thermo Q Exactive HFTM mass spectrometer operating in heated electrospray ionization mode. Data acquisition was 10 minutes with a negative mode full MS scan (profile mode) and one microscan, with an AGC target of 3e6 and a maximum IT of 100 ms at 120,000 resolution, with a scan range from 160 to 400 m/z. Spray voltage was 3.5 kV, and the capillary temperature was set to 320°C with a sheath gas rate of 35, aux gas of 10, and max spray current of 100 µA. The acquisition order of samples and standard curve points was randomized, with blank matrix controls before and after each standard curve point to assess carryover (none detected). The resulting free fatty acid peaks were quantified by measuring the relative intensities (peak heights) of the high resolution extracted ion chromatogram (XIC) for each fatty acid across the samples and external standard curve samples ranging from 10 µg/mL to 100 ng/mL. All fatty acids were detected as the negative mode [M-H] ion, and retention times of the fatty acids were defined using a cocktail of authentic standards. For each XIC, the theoretical m/z of each fatty acid (±5 ppm) was used to extract the peak height (24 seconds retention time window, 12 seconds retention time tolerance) as follows: lauric acid (199.1704 m/z, 2.3 minutes), myristic acid (227.2017 m/z, 3.1 minutes), palmitoleic acid (253.2173 m/z, 3.4 minutes), palmitic acid (255.2330 m/z, 4.1 minutes), oleic acid (281.2486 m/z, 4.4 minutes), stearic acid (283.2643 m/z, 5.1 minutes), arachidic acid (311.2956 m/z, 6.0 minutes), nervonic acid (365.3425 m/z, 6.9 minutes), and lignoceric acid (367.3582 m/z, 7.5 min). The resulting standard curve points (in duplicate) were fit to a linear regression (GraphPad Prism 8), and this equation was used to interpolate the concentration of fatty acids in the sample extracts, as prepared.

Statistical Analysis

All statistical analyses were performed using GraphPad Prism 9.0 Software (GraphPad Software Inc.).

Data Availability

All sequencing data from this study have been deposited in the Sequence Read Archive under BioProject ID PRJNA782676.

Authors' Disclosures

L.M. Coussens reports personal fees from (P30) Koch Institute for Integrated Cancer Research, Massachusetts Institute of Technology, Bloomberg-Kimmel Institute for Cancer Immunotherapy, Sidney Kimmel Comprehensive Cancer Center at Johns Hopkins, (P50) Dana-Farber Cancer Center Breast SPORE, (P30) Dana-Farber/Harvard Cancer Center, (P30) University of California, San Diego Moores Cancer Center, (P30) The Jackson Laboratory Cancer Center, (P01) Columbia University Medical Center, Prostate P01, (P50) MD Anderson Cancer Center Gastrointestinal SPORE, the Cancer Research Institute, the

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Authors' Contributions

J. Abrego: Conceptualization, resources, data curation, formal analysis, validation, investigation, visualization, methodology, writing-original draft, writing-review and editing. H. Sanford-Crane: Conceptualization, resources, data curation, formal analysis, funding acquisition, investigation, methodology, writing-review and editing. C. Oon: Conceptualization, data curation, formal analysis, validation, investigation, visualization, methodology, writing-review and editing. X. Xiao: Data curation, investigation, writing-review and editing. C.B. Betts: Conceptualization, data curation, formal analysis, investigation, visualization, methodology, writing-review and editing. D. Sun: Data curation, formal analysis, visualization, writing-review and editing. S. Nagarajan: Resources, formal analysis, investigation, visualization, methodology, writing-review and editing. L. Diaz: Data curation, formal analysis, investigation, writing-review and editing. H. Sandborg: Data curation, formal analysis, validation, investigation, writing-review and editing. S. Bhattacharyya: Data curation, formal analysis, investigation, visualization, writingreview and editing. Z. Xia: Resources, data curation, formal analysis, supervision, investigation, visualization, writing-review and editing. L.M. Coussens: Resources, data curation, formal analysis, supervision, investigation, visualization, methodology, writing-review and editing. P. Tontonoz: Data curation, formal analysis, supervision, investigation, writing-review and editing. M.H. Sherman: Conceptualization, resources, data curation, formal analysis, supervision, funding acquisition, validation, investigation, visualization, methodology, writing-original draft, writing-review and editing.

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