A microenvironment-inspired synthetic three-dimensional model for pancreatic ductal adenocarcinoma organoids

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Experimental in vitro models that capture pathophysiological characteristics of human tumours are essential for basic and translational cancer biology. Here, we describe a fully synthetic hydrogel extracellular matrix designed to elicit key phenotypic traits of the pancreatic environment in culture. To enable the growth of normal and cancerous pancreatic organoids from genetically engineered murine models and human patients, essential adhesive cues were empirically defined and replicated in the hydrogel scaffold, revealing a functional role of laminin–integrin αv/α6, signalling in establishment and survival of pancreatic organoids. Altered tissue stiffness—a hallmark of pancreatic cancer—was recapitulated in culture by adjusting the hydrogel properties to engage mechano-sensing pathways and alter organoid growth. Pancreatic stromal cells were readily incorporated into the hydrogels and replicated phenotypic traits characteristic of the tumour environment in vivo. This model therefore recapitulates a pathologically remodelled tumour microenvironment for studies of normal and pancreatic cancer cells in vitro.

Tumour cells are embedded in a complex environment encompassing conscripted host cells and a pathologically remodelled extracellular matrix (ECM). Coerced host cells such as immature myeloid cells and macrophages promote the development of an immunosuppressive environment, and cancer-associated fibroblasts (CAFs) deposit and remodel the ECM, resulting in a rigid and poorly perfused microenvironment. These features are particularly prominent in pancreatic ductal adenocarcinoma (PDA), where the stromal reaction constitutes, on average, 80% of the total tumour volume. Normalizing the dysfunctional microenvironment improves therapeutic efficiency, where targeting the stiff ECM or the stromal infiltrate improves efficacy of gemcitabine and immune-checkpoint inhibitors. Models that faithfully replicate the stromal environment are therefore essential to improve the development of therapeutic strategies.

A diverse range of preclinical models, including cell lines, genetically engineered murine models and patient-derived xenografts have been instrumental for target discovery and validation. Recently, organoids have gained traction as a complementary model system, enabling the establishment and three-dimensional (3D) culture of murine and patient-derived tumour cells at high efficiency. However, organoid models are typically established in animal-derived hydrogels, most prominently the basement membrane (BM) extract Matrigel, which suffers from batch-to-batch variability and ill-defined composition, disfavours stromal cell propagation and mostly fails to replicate the pathological ECM of human cancers. Consequently, interdependencies between tumour cells and the microenvironment are inadequately modelled in this system. Contemporary synthetic scaffolds, such as the hydrogel scaffolds based on polyethylene glycol (PEG), offer several advantages to cell- and tissue-derived matrices, including exquisite control over growth conditions.

Although current model systems have been successfully used for primary and induced pluripotent stem cell-derived intestinal organoids, matrices with tissue-matched adhesion characteristics capable of supporting both epithelial organoids and stromal cells are needed to better replicate the complexity of the tumour ecosystem.

Here, we set out to develop a rationally designed, synthetic 3D model for pancreatic organoids that simultaneously captures relevant physiological components of the tissue microenvironment. We envisaged that an ideal model would replicate essential cell–ECM interactions, mimic tissue stiffness ranges observed across normal...
Fig. 1 | Defining adhesive requirements of PCCs. a, Hierarchical clustering of matrisomal protein relative abundances (compared to normal pancreas). Histological grade (bottom) and selected matrisomal proteins (right side) are shown. b, Normalized relative abundance (NRA) of matrisomal proteins from three representative samples (intermediate represents a mixed sample containing PanIN and PDA). The solid line represents similar abundance between samples. c, Scatter plot displaying the variance of core-matrisome proteins over mean NRA in PanIN (left) or PDA (right). Vertical lines represent the 25% quantile and 75% quantile coefficient of variation. Horizontal line indicates median abundance. d, Outline of integrin adhesion complex (IAC) isolation (top) and identified proteins (bottom). Interactions of selected proteins identified in IACs isolated from KPC-1s after 3 h (left) or 12 h (right) are shown. Proteins are colour-coded according to their relative abundance. e, Time-course of green fluorescent protein (GFP)-labelled human SUIT-2 PCCs spreading on indicated ECM proteins. f, Schematic outline of blocking antibodies targeting integrins αvβ3, αvβ6 or αvβ1. g, Effect of blocking antibodies on GFP-labelled SUIT-2 cells spreading on L511- and L521-coated dishes. e,g, Bar graphs show mean values (n = 5), with each data point representing the median cellular area of >100 measured cells for indicated time points. In e and g, the number is the result of the statistical test (P-value) and the square brackets denote which samples are compared. Vertical dashed lines mark the time points selected for and visualized in the bar plots. Two-sided parametric Welch's t-test with Benjamini–Hochberg correction; error bars, s.e.m.
and tumour-bearing tissues, support co-culture of epithelial and stromal cells and facilitate growth and development of organoids directly from tissue samples.

Proteomic analysis of the tumour matrisome and functional validation of cellular adhesion requirements were used to develop a relevant PEG hydrogel scaffold. This system successfully...
replicates the entire stiffness range of normal and tumour-bearing tissue, supports growth of murine and human normal and cancerous organoids and essential stromal populations and is compatible with proteomic and single-cell analyses.

**Results**

**Defining adhesive requirements of pancreatic cancer cells (PCCs).** To reconstitute relevant ECM-receptor interactions within a custom-designed PEG environment, we first defined a matriome reference set for in vivo pancreatic tissues \(^2^0\). Normal pancreatic tissue (NP), pancreata with pancreatic intraepithelial neoplasia (PanIN), known precursor lesions of PDA, and overt PDA from \(\text{Kras}^{\text{G12D}}\) and \(\text{TP53}^{\text{R172H}}\)-driven murine models were collected for analysis \(^2^1–^2^3\) (Supplementary Fig. 1a). We confirmed the high reproducibility of the workflow by analysing ten normal pancreata by liquid chromatography with tandem mass spectrometry (LC–MS/MS) (Supplementary Fig. 1a–d) before analysing 12 tumour samples, which resulted in the identification and quantification of 147 matrisomal proteins, 83 of which predominantly exhibit structural functions (termed core matrisome) (Fig. 1a,b and Supplementary Figs. 2–5) \(^2^4\).

**Formation of hPDOs in defined PEG matrices.**

**a.** Schematic of hPDO establishment.

**b.** Representative brightfield images of hPDOs in 3D PEG CBF-0.5 hydrogels or Matrigel (d6). Images are representative of at least three independent experiments. Scale bar, 500 \(\mu\)m; inset scale bar, 100 \(\mu\)m.

**c.** H&E staining of organoids from b. Scale bar, 200 \(\mu\)m; inset scale bar, 50 \(\mu\)m.

**d.** Maximum intensity projection of hPDOs from four patients (hPDO1, hPDO2, hPDO3 and hPDO4) stained with Phalloidin and Hoechst to visualize organoid morphology and cellular structures. Images show representative morphology of organoids from each line. Scale bar, 200 \(\mu\)m.

**e.** IF analysis of indicated markers in hPDO2 organoids grown in PEG CBF-0.5 hydrogels or Matrigel (d6, \(n=2\)). Scale bar, 100 \(\mu\)m; inset scale bar, 25 \(\mu\)m. Images are representative of minimum five organoids.

**f.** Dual IF for pan-laminin and ITGA6 in hPDOs grown in PEG CBF-0.5 hydrogels (d6, \(n=2\)). Images are representative of minimum five organoids.

Dashed-outline rectangles in **b, c, e** and **f** correspond to insets.

We confirmed the high reproducibility of the workflow by analysing ten normal pancreata by liquid chromatography with tandem mass spectrometry (LC–MS/MS) (Supplementary Fig. 1a–d) before analysing 12 tumour samples, which resulted in the identification and quantification of 147 matrisomal proteins, 83 of which predominantly exhibit structural functions (termed core matrisome) (Fig. 1a,b and Supplementary Figs. 2–5) \(^2^4\).
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αL521-mediated tumour cell adhesion through β1 (Fig. 1d and Supplementary Fig. 6). As this indicates that PCCs
interacted adhesion to FN-coated or non-coated culture dishes (Fig. 1e,
in a cell-line-dependent manner, but consistently exhibited lim-
itsed adhesion to collagen-1
whereas pancreatic fibroblasts adhered to FN-coated dishes (Fig.
8 and Supplementary Videos 1 and 2). Murine and human
tumor and stromal cells were determined (Supplementary
met the relative in vivo abundance of collagen type I and
correlate with patient outcome (Supplementary Fig. 4). By contrast,
the BM proteins laminin 511 (L511, αβγγ) and laminin 521 (L521, αβγγ)
and the type 1 and 4 collagens (for example, Col1a1, Col1a2,
Col4a1 and Col4a2) were consistently highly abundant in both nor-
mal and disease-bearing pancreas (Supplementary Fig. 3b), with
laminin 511 and 521 exhibiting minimal variation across samples
(Fig. 1c and Supplementary Fig. 5c). Together, these findings high-
light important and potentially complementary roles of individual
matrisomal components, including FN, laminins (L322, L511 and
L521) and collagens, throughout PDA development.

Integrins constitute a family of cellular ECM receptors that
are fundamental to cellular adhesion and are commonly aber-
antly expressed in malignant disease. To identify which recep-
tors actively engage the ECM to form IACs, we seeded PCCs on
FN-coated plates for 3 or 12 h and then analysed mature IACs
by mass spectrometry (Fig. 1d and Supplementary Fig. 6).
Interestingly, although PCCs were seeded on FN, the isolated IACs
changed in composition from the canonical FN-binding integrins
αβββ after 3 h to laminin-binding integrins αβββ, αββ and αββ after 12 h, concurrent with the deposition of laminin 521 and
511 (Fig. 1d and Supplementary Fig. 6). This indicates that PCCs
autonomously engineer the ECM to match their specific adhesion
dependencies, we analysed the cell-derived matrix isolated from
PCCs and normal pancreatic ductal epithelial cells (HPDE) by mass
spectrometry. This analysis confirmed the deposition of a BM-rich
ECM that included both L511 and L521 (Supplementary Fig. 7).
Taken together, these data highlight a preference for L511- and
L521-mediated tumour cell adhesion through αβ and αβ-containing integrins.

To test these observations functionally, cell culture dishes were precoated with selected ECM ligands that had been iden-
tified in the proteomic analysis, and the adhesion kinetics of rel-
ant tumour and stromal cells were determined (Supplementary
Fig. 8 and Supplementary Videos 1 and 2). Murine and human
PCCs preferentially adhered to L511- and L521-precoated dishes,
whereas pancreatic fibroblasts adhered to FN-coated dishes (Fig.
1e and Supplementary Figs. 9 and 10). PCCs adhered to collagen-1
in a cell-line-dependent manner, but consistently exhibited lim-
ited adhesion to FN-coated or non-coated culture dishes (Fig. 1e,
Supplementary Figs. 9 and 10 and Supplementary Videos 3–9).
Function-blocking antibodies directed against integrins αβ and αβ
significantly reduced adhesion to L511- and 521-precoated dishes
(Fig. 1f,g and Supplementary Fig. 10), confirming the importance
of L511/L521–integrin αβββ-mediated PCC adhesion in vitro
(Fig. 1g and Supplementary Fig. 10).

Analysis of human PDA by immunohistochemistry (IHC) revealed that cyto keratin-positive PCCs express integrin subunits αβ
and β1, whereas β1 and αβ are broadly expressed across both tumour
and stromal cells. Integrins αβ and αβ appeared to be mostly inter-
cellular. Lama5 staining was observed basal to cyto keratin-positive
tumour cells (Supplementary Fig. 11), supporting relevant roles
of L511–integrin and L521–integrin αβββ, αβββ, and αββ-mediated
PCC adhesion as well as collagen- and FN-mediated adhesion
via integrins αβ and αβ, respectively, that may occur in vivo.
Thus, we reasoned that inclusion of these adhesion cues would be
needed to support epithelial and stromal cells within a synthetic
hydrogel scaffold.

Optimizing PEG hydrogel composition for pancreatic organoids.
To model relevant cell–ECM interactions for pancreatic epithelial
and stromal cells, we used an eight-arm PEG-based hydrogel system
in which adhesion-linker pre-functionalized vinyl sulfone-activated
PEG macromers (f-PEG-VS) are cross-linked via peptides sensi-
tive to matrix metalloproteinase (MMP) to form the hydrogel scaf-
fold (Fig. 2a–c). To support cell adhesion, we incorporated the
FN-mimetic peptide PHSRN-K-RGD (F) and the GFOGER pep-
tide (C), mimicking both fibrillar and network forming collagen,
as well as a BM-binding peptide to retain cell secreted matricellular
proteins more broadly (Fig. 2d and Supplementary Videos 1 and 2).
All adhesion-confering peptides were fully incorporated into the PEG scaffold with limited effect on the
differential elastic gel modulus (Supplementary Fig. 12).

Established mPCOs were disaggregated into single cells and
embedded into PEG-VS hydrogels prepared with different com-
binations of the adhesion-mimetic peptides (Fig. 2b,c). Whilst
FN-mimicking and BM-binding peptides in isolation did not sup-
port organoid growth, the collagen-mimicking peptide supported
expansion of mPCOs (Fig. 2b,c and Supplementary Fig. 13a).
This contrasts with observations made using human colorectal
and endometrial carcinoma organoid lines, which were sustained
by FN-mimetic peptides (Fig. 2d and Supplementary Videos 10 and 11),
demonstrating a complementary effect of the adhesion peptides
on organoid formation and growth. As the matricellular analysis
determined the relative in vivo abundance of collagen type I and

Fig. 4 | Recapitulating the stiffness range of PEG hydrogels. a, Box plot (top) and heat map (bottom) displaying stiffness frequency (Young’s modulus) for murine (left, n = 11 tumour, n = 3 NP) and human (right, n = 3) samples. Box plots represent individual measurements (>800 per sample). The bold line represents the median, boxes represent the 25th to 75th percentiles and whiskers indicate the 90% confidence interval. The genotype of murine samples is shown at the bottom of the panel. Cumulated relative stiffness frequency is shown on the right. Median stiffness for normal (grey) as well as median (purple), mean (orange) and upper 90% border (yellow) of murine cancerous samples. b, Illustration of how cross-linking density after PEG functionalization controls stiffness. c, AFM measurements of PEG CBF-0.5 hydrogels (n = 3; error bars, s.e.m). Linear regression (red) and 95% confidence interval shown (dashed line). d, Representative brightfield images (left) and maximum intensity projections of 4,6-diamidino-2-phenylindole (DAPI, right) from mPCOs grown at indicated stiffness (d4, n = 4). Dashed-outline boxes correspond to insets. Scale bar, 500 µm; inset scale bar, 100 µm (left). e, Violin plot of organoid volume (top) and frequency plots of numbers (bottom) from d. Dashed bold lines show median organoid volume. Number of analysed organoids shown above the plot. f, IF of yes-associated protein 1 (YAP1) and DAPI in mPCOs grown at indicated stiffness. IF images are representative of minimum five organoids (n = 2). Dashed-outline boxes correspond to insets. Scale bar, 100 µm; inset scale bar, 25 µm. g, Nuclear/cyttoplasmic ratio of YAP1 in individual organoids from f. Mean shown as bold line. Two-sided parametric Welch’s t-test used for 1.4, 3.1 or 8.2 kPa and two-sided non-parametric Wilcoxon test for 20.5 kPa with Benjamini–Hochberg correction. Bottom, middle and top lines of boxes show the 25th, 50th and 75th percentiles, respectively, and whiskers show 1.5 times the interquartile range from hinges. h, Normalized Ctgf expression in mPCOs grown at indicated stiffness (n = 3; parametric Welch’s t-test; error bars, s.e.m.). i, Mean organoid volumes (U) and numbers (J) of mPCOs grown at indicated stiffness (n = 4; two-sided parametric Welch’s t-test with Benjamini–Hochberg correction; error bars, s.e.m.). In g–j, the number is the result of the statistical test (P-value) and the square brackets denote which samples are compared.

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FN at 2:1 in all pancreatic matrisome samples (Supplementary Fig. 13c), we used the CBF-0.5 hydrogel formulation for all subsequent studies.

Using this formulation, organoids matured and developed macroscopically visible spheres throughout the entire gel within 3 to 5 days of seeding (Fig. 2d and Supplementary Fig. 14a). In addition, organoids exhibited similar growth kinetics (Supplementary Fig. 14c) and established cellular architecture and polarity comparable to Matrigel when assessed by electron microscopy (Fig. 2e) or IF (Fig. 2f and Supplementary Fig. 14b). Finally, organoids from both normal- and tumour-bearing pancreas were supported at similar efficacy, suggesting that cancer cells retain adhesion dependencies that mirror their non-malignant history (Supplementary Fig. 14d).

To determine whether CBF-0.5 hydrogels support the formation of a relevant ECM, we determined the deposition of ECM proteins in the PEG culture systems by LC–MS/MS (Fig. 2g and Supplementary Fig. 15). This resulted in the identification of 104 matrisomal proteins, including laminin-521, laminin-511, type I collagen and FN, as well as the FN-binding integrins α5β1 and αvβ3, the laminin-binding integrins α6β1, αvβ1, and αvβ3, and the collagen-binding integrin αvβ1 (Supplementary Fig. 15). Using a dual-IF analysis, we confirmed the basolateral localization of integrin αv in close proximity to organoid-derived laminin deposits in the PEG CBF-0.5 gels (Fig. 2h). Importantly, laminin deposits were also identified in PEG hydrogels containing only the GFOGER and PHSRN-K-RGD peptides, indicating that the interactions
between laminin and integrins αv and α6 are functionally important for organoid growth (Supplementary Fig. 16a,b). To functionally verify this interaction, we treated mPCOs grown in PEG CBF-0.5 hydrogels with lebein-2, a disintegrin that selectively blocks the laminin-binding integrins αvβ1, αvβ3 and α6β1, but does not target the collagen-binding integrins αvβ3 and αvβ1 (refs. 40–43). Lebein-2
interfere with adhesion of PCCs onto L511- and L521-coated dishes in vitro (Supplementary Fig. 16c,d) and significantly reduced mPCO formation and growth in PEG CBF-0.5 gels (Fig. 2i and Supplementary Fig. 16e–h). Together, these data demonstrate that relevant cell–ECM interactions are established in PEG hydrogels and that laminin–integrin interactions play a functionally important role for the growth of pancreatic cancer organoids. In conclusion, PEG CBF-0.5 hydrogels support the formation of complex cell–ECM interactions to enable the growth and polarization of pancreatic organoids with comparable efficacy to Matrigel.

**Human pancreatic ductal organoid (hPDO) formation in defined PEG matrices.** To determine whether the synthetic PEG CBF-0.5 hydrogel scaffold also supports expansion of human-patient-derived organoids, we seeded established hPDOS and tracked their growth and morphology (Fig. 3a). hPDOS were engrafted as fragments or single cells in PEG CBF-0.5 hydrogels, where they developed into fully mature spheres after 4–12 days of growth and exhibited similar morphology to hPDOS grown in Matrigel (Fig. 3b,c and Supplementary Fig. 17). Consistently with mPDOS, hPDOS exhibited a high degree of morphological heterogeneity in PEG and Matrigel culture conditions (Fig. 3c,d and Supplementary Fig. 17)⁴⁴. IF analysis confirmed that human organoids were polarized and retained expression of ZO-1, ITGA6, SOX9 and E-cadherin (Fig. 3e and Supplementary Fig. 17c–e). All hPDOS express integrin αi at the cell periphery, close to laminin deposits, indicating that laminin–integrin interactions are conserved in human pancreatic organoids (Fig. 3f). These data demonstrate that PEG CBF-0.5 hydrogels support murine and human pancreatic organoids growth in vitro.

**Recapitulating the stiffness range of PDA in PEG hydrogels.** Deregulated mechano-sensing is commonly observed in solid tumours, where a remodelled ECM promotes malignant traits⁴⁵–⁴⁶. As synthetic hydrogel scaffolds can be configured to recapitulate tissue stiffening⁴⁷–⁴⁸, we set out to determine whether the PEG hydrogel scaffold reliably replicates the stiffness ranges of normal and tumour-bearing pancreas.

To determine how tissue stiffness changes as normal healthy pancreas transitions into tumour-bearing tissue, we analysed 11 murine and 3 human fresh-frozen pancreatic cancerous samples by atomic force microscopy (AFM) and compared their stiffness profiles to those of three murine normal pancreata (Fig. 4a and Supplementary Figs. 18–20). Healthy murine pancreatic tissue exhibited a mean stiffness of 1.48±0.3 kPa, while murine tumour-bearing tissue displayed a median stiffness of 3.55 kPa and a mean stiffness of 8.48±2.2 kPa, with 90% of all measurements below ~20.1 kPa, confirming previous analyses⁴⁶–⁴⁸ (Fig. 4a and Supplementary Fig. 19). Similar stiffening was observed for the three human specimens (Fig. 4a and Supplementary Fig. 20). We confirmed these findings using second-harmonic generation (SHG) imaging for linearized collagen fibres, which are known to correlate with matrisomal stiffening⁴⁹ (Supplementary Fig. 21).

To model this environment in vitro, we tuned the stiffness of PEG CBF-0.5 hydrogels by increasing the molar ratio of the cross-linking peptide to the number of free vinyl sulfone (VS) arms (hereafter referred to as cross-linking density; Fig. 4b). We profiled the resulting hydrogels using AFM and confirmed the relative increase in stiffening by rheology (Supplementary Fig. 12e), demonstrating that the entire stiffness range from normal pancreas to PDA was recapitulated with high reproducibility (Fig. 4c).

To determine the effects of increased PEG hydrogel stiffening on organoid growth, we seeded murine organoids in hydrogels tuned to replicate the pathological stiffness of PDA (Fig. 4d,e and Supplementary Fig. 22). Organoids formed efficiently in hydrogels across the entire stiffness range, displayed elevated nuclear translocation of YAP1 and increased Ctgf levels in stiffened hydrogels, consistent with engagement of mechano-signalling (Fig. 4f–h and Supplementary Fig. 23). To further profile differences in the cellular response to increased hydrogel stiffening, we seeded one murine normal pancreatic organoid (mPNO) and three mPDOS in PEG hydrogels tuned at disease-relevant stiffnesses. While all organoids exhibited a growth advantage when seeded in stiffened gels (Fig. 4i,j and Supplementary Fig. 22)⁴⁴, differences existed in the optimal stiffness for growth across the mPDOS, possibly reflecting the stiffness of the tumour from which the organoids were established. These data demonstrate that the PEG CBF-0.5 hydrogels present a suitable and reproducible model to study effects of tissue stiffening.

**Modelling a heterocellular tumour microenvironment.** To determine whether diverse stromal cell populations from pancreatic tumours are retained in PEG hydrogels, we collected and disaggregated tumours from orthotopically implanted mPDOS and embedded these in PEG CBF-0.5 hydrogels (Supplementary Fig. 24). Strikingly, stromal cell populations were readily detected by IHC for pan-CCK, PDPN and CD45 after 8 days of culture, demonstrating the suitability of CBF-0.5 hydrogels for stromal co-cultures.

Next, we co-cultured mPDOS labelled with infrared fluorescent protein (iRFP), GFP-labelled murine pancreatic fibroblasts (PaFs) and cell-tracker-dye-labelled macrophages, derived from murine bone-marrow-derived myeloid cells (BMDM) treated with macrophage colony stimulating factor 1 (csf-1), in PEG CBF-0.5 hydrogels (Fig. 5a). Fluorescent image, IHC and flow analysis after 6 days of culture validated that viable PaFs and macrophages were retained in the PEG CBF-0.5 hydrogels (Fig. 5b and Supplementary Fig. 25). Next, the cellular dynamics within co-cultures were examined by live imaging for 3 days. Dynamic migratory behaviour of both tumour and stromal cells was observed within the PEG CBF-0.5 hydrogels (Supplementary Fig. 26). For example, individual tumour cells in close proximity to fibroblasts exhibited invasive behaviour and migrated away from the organoid structure (Supplementary Fig. 26 and Supplementary Videos 12–15). Moreover, consistently with previous observations, we also observed that fibroblasts exhibited an elongated, mesenchymal morphology; this was especially apparent with fibroblasts proximal to tumour cells⁵⁰ (Fig. 5c and Supplementary Fig. 26).

To further interrogate the interaction of tumour and stromal cells, we determined the production of functionally important soluble signals (Fig. 5d). Several cell-specific signals such as GM-CSF, CXCL12 and IL10 were identified across monocultures, whereas other signals (IL6 and TGFβ) increased robustly upon co-culture, demonstrating relevant tumour–stromal interactions (Fig. 5d). To further interrogate cellular phenotypes, we analysed co-cultures at single-cell resolution using cytometry by time of flight (CyTOF). Cells were liberated from PEG CBF-0.5 hydrogels after 6 days of co-culture and stained with antibodies directed against known markers of tumour cells, CAFs and macrophages, followed by analysis (Fig. 5e and Supplementary Fig. 27). Visualization using t-distributed stochastic neighbour embedding (t-SNE) distinguished cytokeratin- and EpiCAM-positive epithelial mPDOS from the CD45<sup>++</sup>/CD68<sup>++</sup> BMDM-derived macrophages and GFP<sup>++</sup> fibroblasts, revealing phenotypic diversity across tumour and stromal cell populations. For example, fibroblasts expressed markers of known subsets, such as αSMA<sup>++</sup> myCAs, MHCI<sup>++</sup> apCAs and MHCI<sup>++</sup>/αSMA<sup>++</sup> iCAs (refs. ⁵¹,⁵²) (Fig. 5f and Supplementary Fig. 28). This demonstrates that stromal cell populations are readily incorporated with organoids in PEG hydrogels and exhibit signalling, phenotypic and morphological behaviour consistent with in vivo models. Thus, PEG CBF-0.5 hydrogels can be readily utilized for the study of pancreatic cancer within a functionally relevant stromal context.
Discussion

Contemporary organoid models that replicate physiological characteristics and stromal abnormalities of the tumour microenvironment are needed to improve basic and translational cancer biology.

Here we combined biochemical and functional analysis of cellular adhesion dependencies to inform the design of a synthetic PEG scaffold. Interactions between cell-derived laminin 511 and laminin 521 and integrin αv or αv subunits are recapitulated within a complex cell–ECM niche to support organoid growth. Moreover, PEG hydrogels recapitulate the entire stiffness range of murine and human pancreatic cancers and engage signalling consistent with mechano-sensing. Lastly, stromal cell populations such as pancreatic fibroblasts and bone-marrow-derived macrophages were readily co-cultured and fibroblasts express markers and display morphology consistent with adaptation of myCAF, icAF and acpCAF subsets, demonstrating that stromal cells acquire relevant phenotypes within these defined growth settings.

This rationally designed and easy-to-use PEG system provides exquisite growth control in a reproducible and defined manner to support future basic and translational studies of human and murine pancreatic organoids within a relevant microenvironment.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41563-021-01085-1.

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Methods Mouse models. KPC (KrasLSL-G12D, Pdx-1 Cre); KPF (KrasLSL-G12D, Pdx-1 Flp), KC (KrasLSL-G12D, Pdx-1 Cre) and KF (KrasLSL-G12D, Pdx-1 Flp) animals have previously been described10,37. Animals were bred at CRUK Manchester (CRUK MI) or Beatson Institute (CRUK BI) under pathogen-free conditions. Mice were maintained in a purpose-built facility in a 12 h light–dark cycle at uniform temperature and humidity with continual access to food and water. The genotype of animals was confirmed by Transnetyx (Cordoba). For orthotopic tumour cell implantation studies, six-week-old female CD-1 nude animals were obtained from Charles River. All animal experiments were performed under a UK Home Office Licence and in accordance with the Animal (Scientific Procedures) Act of 1986 under project licence nos. 70/8745 and 70/8375 subject to review by the Animal Welfare and Ethical Review Body of Cancer Research UK Manchester Institute, University of Manchester (CRUK MI) and the University of Glasgow. Experiments are reported in accordance with Animal Research: Reporting of In Vivo Experiments (ARRIVE) 2.0 guidelines.

Patient samples. Research samples were obtained from the Manchester Cancer Research Centre (MCRC) Biobank with informed patient consent obtained prior to sample collection. The MCRC Biobank (ethics code 18/NW/0092) is licensed by the Human Tissue Authority (licence number 30004) and is ethically approved as a research tissue bank by the South Manchester Research Ethics Committee (Ref: 07/ H1003/161+5). The role of the MCRC Biobank is to distribute samples. For more information see www.mcrcc.man.ac.uk/Biobank/Ethics-and-Licensing.

Cell culture of murine and human PCC lines. Work with mammalian cancer cell lines and organoids was performed under sterile conditions. Cells were grown at 37 °C in 5% CO2 and humidified air. Murine KPC-1 cells were obtained from Jennifer Morrison at CRUK BI, UK; iKras cells were kindly shared by Ronald DePinho (MD Anderson Cancer Research Centre, TX, USA); KPC-43 cells were a kind gift from Kris Freese (CRUK MI); human SUIT-2 cells were obtained from ATCC and human H6c7 immortalized adult HPDE cells were obtained a kind gift from Kris Freese (CRUK MI); human SUIT-2 cells were obtained from ATCC and human H6c7 immortalized adult HPDE cells were obtained from CRUK MI. All cells were confirmed mycoplasma free and cultured in Dulbecco's MEM from ATCC and human H6c7 immortalized adult HPDE cells were obtained a kind gift from Kris Freese (CRUK MI); human SUIT-2 cells were obtained from ATCC and human H6c7 immortalized adult HPDE cells were obtained from CRUK MI. For passaging, following removal of the growth media, Matrigel domes were depolymerized in 0.01 µg ml-1 doxycycline (Sigma-Aldrich) at all times.

Establishment and culture of mPNOs or mPCOs. mPNOs were isolated from wild-type C57Black6c knockout pancreas or tumour-bearing KPC mice with histologically verified tumours. Tissues were minced (<0.5 mm in diameter) and enzymatically digested with 5 mg ml-1 Collagenase-II (Thermo Fisher Scientific) in advanced DMEM F12 (Gibco) supplemented with 1x foetal bovine serum (FBS, Life Technologies) and 1x HyClone solution (Invitrogen), hereafter referred to as DMEM 10% v/v FBS. HPDE cells were grown in keratinocyte serum-free medium (KSF) supplemented with 0.2 ng ml-1 EGF (Invitrogen), 30 µg ml-1 bovine pituitary extract (Invitrogen, catalogue no. 17005042) and 1x HyClone solution. PCC lines or HPDEs were allowed to reach a maximum confluence of 80% and split using 0.25% v/v Trypsin-Verse (Invitrogen). iKras cells were cultured in DMEM 10% v/v FBS supplemented with 1 µg ml-1 doxycycline (Sigma-Aldrich) at all times.

Culture and establishment of hPDOs. hPDOs were passaged at least once before usage for experimental procedures.

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In brief, hydrogels were crafted using Michael-type reaction chemistry inside a 1 ml syringe (BD) modified by removal of the tip. All hydrogels were prepared by reacting 1.5 mg of an eight-arm 20 kDa-PEG-VS macromer (2.4 mM PEG-VS (19.7 mM VS)) with 1 mM of GFOGER, 1 mM PHSRN-K-RGDS and 0.5 mM BM-binder. Final gelation, cells grown in hPCOM supplemented with 40 μM lebein-2 treatment. Media were half-replenished every day and new lebein-2 was added to the control. Comparing growth between Matrigel and PEG CBF-0.5 hydrogels and lebein-2 treatment assays, mPCOs were incorporated in PEG CBF-0.5 hydrogels.

For IHC, hydrogels were embedded in O.C.T., snap-frozen and sectioned at 7 μm. Sections were fixed for 15 min in 4% PFA at room temperature and washed in PBS. For podoplanin staining, cells were fixed in 1:1 acetone/methanol for 15 min at 4°C. Endogenous peroxidase was blocked using 3% H₂O₂ in PBST for 10 min at room temperature followed by PBST washes for 5× 2 minutes each. Non-specific antibody binding was blocked with 10% casein for 20 min at room temperature. Cells were then stained using primary rabbit anti-F-A (1:100, Cell Signaling Technology, catalogue no. 70076), rabbit anti-Pan-CK (1:100, Abcam, catalogue no. ab3977), rabbit anti-CDS (1:100, BD Biosciences, catalogue no. 550539, clone 30-411) or rabbit anti-Podoplanin (1:500, Thermo Fisher Scientific, catalogue no. MAS-29742) antibodies diluted in PBST for 1 h at room temperature. Following extensive washing with PBST, an interrupted goat anti-Rabbit secondary antibody (RTU, Dako, catalogue no. K4003) or HRP conjugated goat anti-Rat secondary antibody (RTU, Vector Laboratories, catalogue no. MP-7444) was diluted at 30 min at room temperature. Liquid DAB+ (Dako, catalogue no. K3468) chromogen was applied for 5 min at room temperature followed by counterstaining in Gill’s haematoxylin (Thermo Fisher Scientific, catalogue no. 6755006) for 30 s. After mounting in Pertex mounting medium (CellPath, catalogue no. SEA-0100-00A), slides were scanned on the Leica SCN400 (Leica Microsystems), using a ×20 objective lens under brightfield using the Leica LAS software to control the hardware and export the data to TIFF format. Raw Leica format files are initially stored onto a Synology RAID system and the exported to a read-only central storage solution to ensure data integrity.

Lebein-2 isolation and purification. Lebeins were isolated from the venom of Macrovipera lebetina obtusa (Latoxan, catalogue no. L1126) as described previously. In brief, upon gel filtration of the venom proteins on a Sepharose 6 column (GE Healthcare) in 20 mM sodium phosphate, pH 6.5, 50 mM sodium chloride and 1 mM EDTA, the relevant fractions were pooled, diluted with 10 mM MES (pH 5.7) and loaded onto a Mono S column (GE Healthcare). Lebein-2 was then eluted in a linear sodium chloride gradient from 80 to 140 mM and further purified on a C8 reversed-phase column (Nucleosil, Machery Nagel) in a linear gradient from 0.1% trifluoroacetic acid in water to 80% acetonitrile in 0.08% trifluoroacetic acid in water. After lyophilization, the lebeins were dissolved in a 50 mM HEPES, 150 mM NaCl solution (pH 7.4) and kept at −80 °C until use. Lebein-2 charges used in this study were validated for effectiveness in adhesion assays.

Comparison of growth between Matrigel and PEG CBF-0.5 hydrogels and lebein-2 treatment. To evaluate murine PDO (mPDO) growth rate, single-cell suspensions of IRFP-labelled mPDOs were generated and encapsulated in PEG CBF-0.5 or Matrigel at 1,000 cells per μl hydrogel as previously described. Growth was assessed by measuring IRFP signal using Odyssey imaging system (Lico). For lebein-2 treatment, cells were incorporated in PEG CBF-0.5 hydrogels as described previously at a concentration of 1,000 cells per μl hydrogel. Upon gelation, cells grown in hPCOM supplemented with 40 μg ml⁻¹ lebein-2 or a vehicle control. Media were half-replenished every day and new lebein-2 was added to the cells to maintain the inhibition. A 50 mM HEPES, 150 mM NaCl solution (pH 7.4) and kept at −80 °C until use. Lebein-2 charges used in this study were validated for effectiveness in adhesion assays.

Reverse transcription–quantitative PCR (RT–qPCR). RNA of murine organoids grown in synthetic gels was isolated using the TRIZol reagent (Invitrogen) and pooled for sequencing.
AFM. Tissue pieces isolated from the diseased portion of the pancreas and spanning the entire width of the tumour (for murine samples, d = 1–1.5 cm; d = 0.5–1 cm for human samples) were embedded in O.C.T. medium and snap-frozen on dry ice submerged in isopentane (Sigma-Aldrich). Sections 14 µm thick were cryosectioned from the frozen tissue blocks and stored at −20 °C on glass slides (Superfrost Plus, Thermo Fisher Scientific) until required for AFM. Prior to use, sections were brought up to room temperature and rested for approximately 30 min to allow the sections to fully adhere to the glass slide. Following this, sections were incubated in the dark. Siouxsie AFM 3000 AFM microscope was used to ensure complete removal of O.C.T. Sections were then air-dried overnight at room temperature. Each section was then rehydrated in ultrapure water for at least 10 min at room temperature immediately before data acquisition. AFM indentations were acquired using a Bruker Catalyst with a NanoDome V controller (Bruker UK Ltd, Coventry UK) operating under Nanoscope software (v.4.8). The system was mounted on a Nikon Eclipse 50i microscope (Nikon Instruments) equipped with a Hamamatsu ORCA-05G camera. For tissue sample indentations, 5 µm diameter gold colloids were mounted on cantilevers with a nominal spring constant of 0.1 N m⁻¹ (CP−aq−CONT-Au−B, qUbe). Cantilevers were calibrated before each experiment using the thermal oscillation method in air. For each tissue sample, force measurements were acquired from nine areas distributed over representative stromal and epithelial areas (as determined from an adjacent H&E stained section). In each area, data were obtained from 100 individual indentations, evenly spaced over a 25 µm² area, with a ramp size of 4–6 µm and a maximal indentation depth (trigger threshold) of 4% of the ramp size (1 µm sample depth). For biophysical profiling of hydrogels, gels were prepared 24 h before analysis. First, 20 µl of gel were pipetted onto pre-cleaned poly-l-lysine precoated glass slides (Thermo Fisher Scientific) and allowed to polymerize at 37 °C for 40 min. Gels were then immersed in PBS and stored overnight at 37 °C. Finally, the gels were washed twice in PBS before AFM measurements. For each PEG gel, ten force maps were obtained consisting of 25 individual indentations evenly acquired over 25 µm². Data analysis for both the tissue samples and the gels was performed using the Bruker NanoScope Analysis software suite v.4.8 after baseline correction. The Hertz model was used to determine the elastic properties of the sample (equation (2)), with F being the force, E being Young's modulus, µ being the Poisson's ratio, R being the radius of the indent, and d being the indentation. The maximum and minimal force fit boundaries of 80% and 20%, respectively, of the extension curve were used during all calculations. Values exceeding a Young's modulus of x µ = 2σ (where x is any one calculated modulus value, µ is the mean of the calculated moduli values and σ is standard deviation) within each force map were removed from the analysis. A Poisson's ratio of 0.5 was assumed during all calculations of the Young's modulus.

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F = \frac{3(1-\nu^2)}{2\pi} E R^2 d^4
\]

(IAC isolation. IACs of PCGs were isolated as previously described. In brief, for each condition and cell line, six dishes with a 10 cm diameter were coated with 10 µg ml⁻¹ FN (Sigma–Aldrich, catalogue no. F1141) in PBS containing calcium and magnesium (PBS±, Thermo Fisher Scientific) to stimulate FN-mediated adhesion. PCGs were cultured and passaged and single cells seeded onto each FN-coated 10 cm dish in 5 ml of DMEM 10% (v/v) FBS. Following incubation at 37 °C for either 3 h or overnight, the media were changed, and cells were washed twice with pre-warmed DMEM–HEPES (Sigma–Aldrich) to remove all non-adherent cells. IACs were then cross-linked by adding a 6 mM dimethyl 3,3'-dithiopropaniminodimethyl dithioldichloride (DTBP; Sigma–Aldrich, catalogue no. D2288). DMEM–HEPES solution for 5 min at room temperature. Cells were washed twice with ice-cold PBS and lysed in modified radioimmunoprecipitation array assay buffer (RIPA, 50 mM Tris-HCl (pH 7.6), 150 mM NaCl, 5 mM disodium EDTA (pH 8.0), 0.5% w/v SDS, 1% v/v Triton X-100 and 1% v/v sodium deoxycholate in ddH2O) for 20 min at 37 °C.

Cell bodies were removed using a high-pressure water wash. Protein complexes remaining attached to dishes were washed in cold PBS and recovered in 100 µl of adhesion complex recovery solution (125 mM Tris-HCl (pH 6.8), 1% w/v SDS and 1 mg ml⁻¹ glycerol) by scintillation (DTBP) trapping. Protein complexes were then precipitated overnight using four volumes of −20 °C acetone at −80 °C. Precipitated proteins were centrifuged at 16,000 g for 20 min at 4 °C and protein pellets washed with −20 °C acetone (16,000 g, 20 min, 4 °C). Proteins were then allowed to dry at room temperature for about 20 min. Precipitated complexes were resuspended in 2x reducing sample buffer (50 mM Tris-HCl (pH 6.8), 150 mM NaCl, 5% (v/v) glycerol, 4% (w/v) SDS, 0.004% (w/v) bromophenol blue and 15% (v/v) β-mercaptoethanol) and heated to 70 °C for 10 min and then heated for 3 min at 95°C.

In gel trypsic digest of adhesion complexes and LC–MS/MS. Protein samples were loaded on 4–15% polyacrylamide gels and electrophoretically separated for 3 h. Proteins were run off the gel. Gels were then stained with Coomassie Blue solution (Bio-Rad) for 10 min at room temperature, and protein bands were cut into 1 mm² pieces and transferred to a 96-well PVDF plate (FlilEx, Corning, catalogue no. 3504) with a pure size of 0.2 µm. Gel pieces were then washed with a freshly prepared 25 mM NH₄HCO₃ (Sigma–Aldrich) solution in H₂O followed by another wash in 100% (v/v) acetonitrile for 10 min. Gel pieces were then washed twice with 100% (v/v) acetonitrile for 5 min and gel pieces dried using a vacuum centrifuge for 30 min at room temperature. Proteins were reduced by incubation in 10 mM DTT diluted in 25 mM NH₄HCO₃, for 1 h at 56 °C and alkylated in 55 mM iodoacetamide diluted in 25 mM NH₄HCO₃, for 30 min at room temperature. Gels were washed with 1x acetonitrile amounts of 25 mM NH₄HCO₃ and acetonitrile for 5 min each and gels dried for 25 min at room temperature in the vacuum centrifuge. A 50 µl aliquot of 12 µg trypsin (Pierce) in 25 mM NH₄HCO₃ was added to each sample and incubated for 15 min at 4 °C followed by an overnight incubation at 37 °C. Upon digestion, peptides were extracted using 100% (v/v) acetonitrile in 0.2% (v/v) formic acid (FA, Thermo Fisher Scientific) for 20 min and from this time 1% (v/v) acetonitrile was added and 5% (v/v) FA in HPLC-grade water. Peptides were then washed using 0.1% (v/v) FA in HPLC-grade water and eluted in 30% (v/v) acetonitrile in 0.1% (v/v) FA, dried and resuspended in 5% (v/v) acetonitrile in 0.1% (v/v) FA for LC–MS/MS.

Peptides were analysed using an Ultimate 3000 RSLCnano system (Thermo Fisher Scientific) coupled to an Orbitrap Elite mass spectrometer (Thermo Fisher Scientific). Peptides were loaded onto a pre-column (300 µm × 180 mm internal diameter, Waters) and separated on a bridged ethyl hybrid (BEH) C18 column (250 mm × 75 mm internal diameter, 1.7 µm particle size, Waters) over a 2 h gradient from 8 to 33% (v/v) acetonitrile in 0.1% (v/v) FA at a flow rate of 200 nl min⁻¹. LC–MS/MS analysis was performed using a data-dependent mode to allow selection of peptide fragmentation in an automated manner. The resulting data were searched against the SwissProt database with species set to Mus musculus on an in-house Mascot server (Matrix Science; 2016) in Proteome Discoverer (Thermo Fisher Scientific, v.2.1). Search parameters included peptide modifications for carbamidomethylation (C) as static modification and oxidation (M, P and K) as well as deamination (N, Q) as dynamic modification. A decoy database search was performed to determine the peptide FDR. The peptide FDR was calculated using the FDR (false discovery rate) with the Percolator module. A 1% peptide FDR threshold was applied, peptides were filtered for high peptide confidence and minimum peptide length of six and finally, peptides without protein reference were removed. All proteins that exhibited a confidence of less than ‘high’ and with less than two uniquely identified peptides were excluded from further analysis.
ex extractions were performed in batches of five on two consecutive days to control for intra- and inter-extraction variability.

S-Trap-based tryptic digestion of ECM-enriched fractions and LC–MS/MS. ECM proteins were prepared for LC–MS/MS using S-Trap Midi Columns (Protifi). Samples were pulverized with frozen urea and resuspended in 8 M urea supplemented with 5% w/v SDS, 20 mM HEPES and 50 mM triethylammonium bicarbonate (TEAB) and harshly sonicated. Samples were then reduced in 5 mM tris(2-carboxyethyl)phosphine (Sigma–Aldrich) for 2 h at 37 °C and 1,400 r.p.m. (Thermomix) and alkylated using 15 mM S-methyl methanethiosulfonate (MMTS, Sigma–Aldrich) for 30 min at room temperature. Urea was then diluted to 2 M by adding 50 mM TEAB and proteins were deglycosylated using PNGaseF (New England Biolabs) for 2 h at 37 °C and 1,400 r.p.m. Following sonication, samples were centrifuged at 15,000 g for 30 min at 4 °C. The supernatant was passed through the columns. Trapped proteins were then washed with STBB in 4× and 1× washes and proteins were digested overnight at 37 °C in 50 mM TEAB using sequencing-grade trypsin at a ratio of 1:10 in a humidified incubator. Following digestion, peptides were eluted in 500 µl 50 mM TEAB followed by 500 µl 0.2% FA in LC–grade water and 500 µl 50% acetonitrile supplemented with 0.2% FA in LC–grade water at 4,000 g for 60 min each. Peptides were then dried down and desalted using a C18–Pak Vac Prep Module (WAT036820) according to the manufacturer’s instructions. In brief, following equilibration of the C18 cartridges with 100% acetonitrile supplemented with 0.1% FA and 0.05% TFA in LC-grade water, samples were loaded in 0.05% TFA in LC-grade water and washed with 0.05% TFA in LC-grade water followed by washing with 0.1% FA in LC-grade water. Peptides were then eluted in 30% acetonitrile in LC-grade water and followed by drying and elution with 2% FA and 98% water for 2 h. Peptides were resuspended in 0.05% v/v FA for LC–MS/MS. Next, 5.5 ng worth of peptide in 10 µl of 0.05% TFA were trapped on an PepMap 300 (C18, 300 µm × 5 mm, Thermo Fisher Scientific) column and separated on an Easy-Spray RSLC C18 column (75 µm × 500 mm, Thermo Fisher Scientific, catalogue no. ES803) at 200 nl min $^{-1}$ using the following gradient profile (minutes:%B): 0.0, 30-40, 40-60, 60:0, 100:0, 0.0, 100:0. Buffer A consisted of 0.1% FA in LC-grade water and buffer B of 100% acetonitrile. The eluent was directed into an Easy-Spray source (Thermo Fisher Scientific) with a temperature set at 60 °C and a source voltage of 1.9 kV. Data were acquired on QExactive HFX (Thermo Fisher Scientific) with precursor scan ranging from 375 to 1,200 m/z at 120,000 resolution and automatic gain control target of 3 × 10$^{6}$. The isolation window was set to 1.5 m/z. dd-MS$^2$ scans were conducted at 30,000 resolution and automatic gain control target of 1 × 10$^5$ and normalized collision energy set to 29%. At least four injections of each sample were conducted for each analysed tumour. For normal pancreas samples, at least five injections were conducted.

The resulting data were analysed in Progenesis (v.26.45.1656 Nonlinear Dynamics) using a H3 approach for each individual replicate and searched against S-Trap-based tryptic digestion of whole cell lysates from synthetic hydrogels. of six and finally, peptides without protein reference were removed.

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High-pH offline fractionation and LC–MS/MS. Peptides were fractionated as previously described$^{35}$. Briefly, peptides were separated on a Zorbax Extend-C18 column (4.6 mm × 150 mm, 3.5 μm, Agilent Technologies) at 250 µl min$^{-1}$ using the following gradient profile (minutes:%B): 5.0, 20.5, 30.0, 24.0, 26.75, 29.75, 30.0.5, 55.0.5. Buffer A consisted of LC–grade water supplemented with 0.1% v/v NH$_4$OH (pH 10.5, and buffer B of 100% acetonitrile. The eluent was directed onto 96–well microplate plates and four aqueous fractions in the elution window between 15:50 and 35:00 min were used, and all of the fractions were concatenated into 24 final fractions, with each containing 3.33 µg peptide on average. Following drying in a speedVac at 60 °C for 2 h, peptides were resuspended at 100 ng ml$^{-1}$ in 0.05% v/v TFA for LC–MS/MS. Next, 10 ng of peptides were trapped on a PepMap 300 (C18, 300 µm × 5 mm, Thermo Fisher Scientific) column and separated on an Easy-Spray RSLC C18 column (75 µm × 500 mm, Thermo Fisher Scientific, catalogue no. ES803) at 200 nl min$^{-1}$ using the following gradient profile (minutes:%B): 6.10, 40.24, 45.45, 46.80, 49.80, 50.1.0, 70.1.0. Buffer A consisted of 0.1% FA in LC-grade water and buffer B of 100% acetonitrile. The eluent was directed into an Easy-Spray source (Thermo Fisher Scientific) with a temperature set at 60 °C and a source voltage of 1.9 kV. Data were acquired on QExactive HFX (Thermo Fisher Scientific) with precursor scan ranging from 375 to 1,200 m/z at 120,000 resolution and automatic gain control target of 3 × 10$^6$. The isolation window was set to 1.5 m/z. dd-MS$^2$ scans were conducted at 30,000 resolution and automatic gain control target of 1 × 10$^5$ and normalized collision energy set to 29%. The resulting data were searched against the SwissProt database with species set to M. musculus on an in-house Mascot server (Matrix Science, 2016) in Proteome Discoverer (Thermo Fisher Scientific, v.2.1). Search parameters included peptide modifications for carboxymethylation (C) as static modification and oxidation (M, P and K) as well as deamidation (N, Q) as dynamic modification. A decoy database search was performed to determine the peptide FDR with the Percolator module. A 1% peptide FDR threshold was applied, peptides were filtered for high peptide confidence and a minimum peptide length of 6 and finally, peptides without protein reference were removed. Protein grouping was performed by applying strict parsimony principles. All proteins that exhibited a confidence lower than high and with fewer than two uniquely identified peptides were excluded from further analysis.

Electron microscopy. Samples were extensively washed with PBS and fixed with 4% formaldehyde and 2.5% glutaraldehyde in 0.1 M HEPES buffer (pH 7.2). Subsequently, samples were postfixed with 1% osmium tetroxide and 1.5% potassium ferricyanide in 0.1 M cacodylate buffer (pH 7.2) for 1 h, then 1% tannic acid in 0.1 M cacodylate buffer (pH 7.2) for 1 h and finally, in 1% uranyl acetate in water for 1 h. The samples were then dehydrated in ethanol, series-infiltrated with TAAB 812 resin and polymerized for 24 h at 60 °C. Sections were cut with a Leica EM UC6 ultramicrotome and observed with an FEI Tecnai 12 Biotwin microscope at 100 kV accelerating voltage. Images were taken with a Gatan Orius SC1000 CCD camera.

Live imaging of pancreatic cancer organoid co-cultures. CBF-0.5 hydrogels were dissociated using Sortase A pentamutant (SrtA), PBS was prepared as described previously and cultured into 10µl domes in 24-well Sensoplates (Greiner). Co-culture domes were imaged via the PerkinElmer Opera Phenix microscope. A Zeiss EC Plan Neofluar ×10 air objective (NA 0.3, WD 5.2 mm) or Zeiss W Plan-Apochromat ×20 water objective (NA 1.0, WD 1.17 mm) were used as indicated. Pre-scan directed z-stack acquisition of individual spheres was performed and used per well in the first experiment (n = 3) with a time window of 72 h starting from day 3 of culture. Videos and representative still images were generated from maximum intensity projections using Harmony 4.8 software. Representative images were exported from Harmony 4.8 in the portable network graphic format with scaling equally applied to all conditions.

ELISA. Concentrations of GM-CSF (catalogue no. DY415-05), IL-6 (catalogue no. DY406-05), IL-10 (catalogue no. DY400-05), IL-1β (catalogue no. DY401-05), TNFα (catalogue no. DY410-05), CXCL12 (catalogue no. DY453-05) and TGFβ (catalogue no. DY1679-05) (all from R&D Systems) in organoid cultures were measured using ELISA DuoSet kits as per the manufacturer’s instructions. In brief, CBF-0.5 hydrogels were dissociated using Sortase A pentamutant (SrtA), PBS was prepared as described previously and cultured into 10µl domes in 24-well Sensoplates (Greiner). Co-culture domes were imaged via the PerkinElmer Opera Phenix microscope. A Zeiss EC Plan Neofluar ×10 air objective (NA 0.3, WD 5.2 mm) or Zeiss W Plan-Apochromat ×20 water objective (NA 1.0, WD 1.17 mm) were used as indicated. Pre-scan directed z-stack acquisition of individual spheres was performed and used per well in the first experiment (n = 3) with a time window of 72 h starting from day 3 of culture. Videos and representative still images were generated from maximum intensity projections using Harmony 4.8 software. Representative images were exported from Harmony 4.8 in the portable network graphic format with scaling equally applied to all conditions.

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were prepared using 0.25% v/v Trypsin-Versene Solution. Following neutralization with DMEM 10% v/v FBS and centrifugation, cells were resuspended to a 106 cells ml−1 in PBS. Cells were incubated with LIVE/DEA Fixable Blue Dead Cell Stain (1:1,000, Thermo Fisher Scientific) at room temperature for 30 mins. Following washing in PBS, cells were resuspended in Cell Staining Buffer (CSM; Fluidigm) containing 5 mM EDTA, hereafter named CSM-1, supplemented with Fc block and subjected to surface staining with phycoerythrin-conjugated anti-CD45 (1:200, Le蟾ogen, clone 30-F11) for 45 min using the Fluidigm C10. Cells were washed in PBS, passed through a 70 μm filter and resuspended at a concentration of 1 x 106 cells ml−1 and analyzed on an LSRFortessa X-20 flow cytometer (BD) using 355 nm, 488 nm, 561 nm and 640 nm lasers, with 450/50, 515/20, 586/15 and 730/45 filter sets, respectively. Cell-line-specific events were selected based on GFP positivity (PaF), IRFP positivity (mPCO) and CD45 positivity (BMDM), and viable cells were selected based on negativity for LIVE/DEA blue stain.

Sample preparation for mass cytometry. To identify cells in synthesis phase (S phase), 5-iodo-2′-deoxyuridine (Sigma-Aldrich) was added to organoid culture medium at a final concentration of 25 μM and incubated for 30 min at 37°C. Cells were then either dissociated immediately or fixed in 4% PFA for 60 min at 37°C, washed twice in PBS (Fisher) and stored at 4°C until needed. Both fixed and unfixed PEG-VS hydrogels were dissolved using SrtA (Supplementary Methods). The resultant solution was transferred to a gentleMACS C-Tube (Miltenyi) containing 0.5 mg ml−1 Dispase II (Sigma), 0.2 mg ml−1 Collagenase IV (Gibco) and 0.20% v/v Collagenase (Gibco), made up to 5 ml in PBS and incubated using the gentleMACS Octo Dissociator (with heaters; Miltenyi) at 37°C for 50 min using a custom program adapted from Qin et al.57. C-tube was spun at 1,000 × g in the void volume before addition of 2 ml of ice-cold 20% methanol for 20 min at −20°C. Following incubation, cells were washed twice in 3 ml CSM-1 and the cell pellet was resuspended in the void volume. Next, 1 μl of Fc block was added for 5 min at room temperature, after which 50 μl of the heavy-metal tagged antibody master mix was added to 50 μl CSM and incubated for 45 min at room temperature (Supplementary Table 2). Following incubation, cells were washed twice with CSM-1. The resultant pellet was resuspended in 200 μl PBS, 4% PFA was added under vortex and it was stored overnight at 4°C.

FOX3 Fix/Permeabilization. Unfixed single-cell suspensions were washed in CSM-1. Cells were then pelleted by centrifugation; this and all subsequent centrifugations were performed at 1,000 × g at 4°C. Supernatant was removed and cells were vortexed in the void volume before the addition of 1 μl of 10% FBS (CSM-2) for 5 min at room temperature. Exposure to heavy-metal tagged antibody master mix was added to 50 μl CSM and incubated for 45 min at room temperature. Following incubation, cells were washed twice with CSM-1. FOX3 Fix/Perm Staining Buffer set (Invitrogen) was then used to fix and permeabilize the cells according to the manufacturer’s instructions. In brief, cells were resuspended in 1x Thermo FOX3 Fix Buffer and incubated for 30 min at room temperature. Next, 2 μl of 1x Thermo FOX3 perm buffer was added and cells pelleted. A further 2 μl of 1x Thermo FOX3 perm buffer was added and cells pelleted. The cell pellet was resuspended in the void volume and 1 μl of Fc block added for 5 min at room temperature. Next, 50 μl of the intracellular heavy-metal tagged antibody master mix was added to 50 μl CSM and incubated for 45 min at room temperature. Following antibody incubation, cells were washed twice with CSM-1. The resultant pellet was resuspended in 200 μl PBS, 4% PFA was added under vortex and it was stored overnight at 4°C.

All samples were processed similarly from this point. The following day, 1 μl of tridimensional fixation (Triton X-100) was added per 30 μl of cells followed by incubation for 60 min at room temperature. The cell suspension was diluted 1:3 in PBS before pelleting. Cells were washed in Maxpar wafer (Fluidigm) and resuspended in Maxpar wafer containing 15% (v/v) EQ beads (Fluidigm) to a concentration of 106 cells per ml and filtered twice (70 μm) before measuring on a Helios mass cytometer (Fluidigm) (300–300 events per second). Files were normalized against EQ beads (Fluidigm) and background subtracted (cytoBank processor http://www.cytobank.org/). Cell doublets and aggregates were removed based on event length. Live cell events were selected based on cleaved caspase-3 negativity. Cell-line-specific events were selected based on GFP positivity (PaF) and CD45 positivity (BMDM). Selected populations were exported as FCS files and uploaded to the CytoKit2 platform (v.2.0.1). Cells were clustered using FlowSOM and visualized using UMAP projections and expression overlays, and cell data with annotated clusters were exported for further downstream analysis of marker expression using R Studio v.4.0.0 (https://www.r-project.org).

Survival analysis. For comparison of matrisomal protein changes between studies, supplementary data files from Tian et al.63 were downloaded, and Log2FC between PDA and NP samples were calculated. To assess resuspending cells from either the murine or human cohort. For the human cohort, all samples were used regardless of differentiation grade. Using R v.4.0.0, we computed the Pearson or Spearman-based rank correlation of the Log2FC(PDA/WT) of matrisomal proteins identified and quantified in this study and Tian et al.63. For survival analysis, matrisomal proteins with a Log2FC(PDA/WT) ≥ 1 in this study and Tian et al.63 were used, and gene identifiers were transformed into entrezgene ids using the biomartR package64. Clinical information and gene expression data for human pancreatic cancer (TCGA-PAAD) were downloaded from cBioPortal (accessed 22 April 2020; refs. 65,66) and log-transformed. The Cox-proportional hazard regression P value and hazard ratio were determined in quintile-categorized patients (top 25% versus bottom 25%) by the matrisomal protein expression using the TCGA data. A survival analysis was done using the survival package (v.3.2-3, https://cran.r-project.org/web/packages/survival/index.html) and Kaplan–Meier survival plots were computed using the survminer package in R v.4.0.0.

Bioinformatics and statistics. Bioinformatics were conducted using the open-source R software (v.4.0.0). Proteins with matrisomal origin were annotated from proteomic datasets using the murine matrisome65. Proteins with BM origin were annotated from matrisome using the gene ontology term ‘basement membrane’ (GO:0005064; http://www.informatics.jax.org/go/terms/gor/GO:0005064, accessed 5 October 2020 (ref. 71)). Pathway annotations based on the Kyoto Encyclopedia of Genes and Genomes (KEGG) as defined in the supplementary pathways and biological process term annotations were obtained from the database for annotation, visualization and integrated discovery (DAVID) functional annotation clustering by assuming the whole genome as statistical background. For gene ontology and biological process term enrichment, only terms with at least three members, an FDR ≤ 0.05, Benjamini–Hochberg adjusted P value ≤ 0.05 and fold enrichment of ≥ 2.5 were included. For KEGG-pathway annotation, only pathways with at least three members, a P value ≤ 0.05 and fold enrichment ≥ 1.5 were included. MatrixCircos plots were generated using the open-source Circos R package72 by applying a custom-made Circos core-component library containing predefined and user-defined core-modules only. Circular visualizations of YAP1 nuclear/cyttoplasmic ratio were implemented using the circlize R package73. Integrin–ECM interactions were obtained from a publicly available manually curated cell–cell interaction database from the Bader lab (https://baderlab.org/CellCellInteractions), and only literature-based interactions with PubMed annotation were included in the analysis. For integrin–ECM interactions, only those interactions nucleating from an integrin α subunit were considered and integrin complexes containing an α and β subunit were subsequently manually assembled. For network analysis and protein–interaction network visualization, the cytoscape software package74 was used by incorporating the String database (STRING CONSORTIUM, v.10.5, 2018 (ref. 75)), MCODE and boundaryLayout apps. For String-based network visualization, networks were curated using a very stringent confidence of 0.95 by accepting text-mining, experimental, database, co-expression, neighbourhood, gene-fusion and co-occurrence interaction sources to obtain relevant integrin–ECM interactions. Statistics and visualizations were conducted using GraphPad Prism v.8 or R v.4.0.0. For statistical analysis of high-content imaging data, the non-parametric two-tailed Wilcoxon t-test was utilized with application of Benjamini–Hochberg correction and an FDR threshold of 0.05 in R. For statistical analysis of adhesion assays and PDO growth assessments in stiffened hydrogels, the parametric two-tailed Wilcoxon t-test was conducted with application of Benjamini–Hochberg correction and an FDR threshold of 0.05 in R. For statistical analysis of adhesion assays and PDO growth assessments in stiffened hydrogels, the parametric two-tailed Wilcoxon t-test was conducted with application of Benjamini–Hochberg correction and an FDR threshold of 0.05 in R. For statistical analysis of lebein-2 treated organoids, the two-tailed Student’s t-test was used, and results of YAP1 ratio between stiffened hydrogels, the parametric two-sided Student’s t-test was conducted to compare 1.4, 3.1 and 8.2 kPa conditions whilst the non-parametric Wilcoxon test was utilized for comparison of 1.4 and 20.5 kPa conditions upon removal of outliers and with application of Benjamini–Hochberg correction and an FDR threshold of 0.05 in R. Outliers were identified and removed from each population when exceeding 1.5 times the interquartile range and confirmed as outliers by the Grubb’s test function from the outliers package in R v.4.0.0. Normality was assessed in R using the Shapiro–Wilk test with a significance level of 0.05.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

All original source data are freely available. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE76 partner.
repository with the following dataset identifiers: NP matrisome atlas (PXD022555 and 10.6019/PXD022555); IAC datasets (PXD022487 and 10.6019/PXD022487); cell-derived matrix datasets (PXD022509 and 10.6019/PXD022509); 3D PEG-CBF-0.5 LG-Ms (PXD022530 and 10.6019/PXD022530); Tumour Matrisome LC–MS (PXD022767 and 10.6019/PXD022767). Raw CyTOF data, IF images and AFM force curves as well as source data for all figures (Figs. 1–5 and Supplementary Figs. 1–28) have been deposited to https://zenodo.org/record/4664132.

Code availability
All original R scripts have been deposited to https://zenodo.org/record/4664132 and are freely available.

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Author contributions
C.R.B., J.K., A. Brown, B.Y.L., J.D.H., D.L.S., L.G.G., M.J.H. and C.J. designed the research; C.R.B., J.K., A. Brown, A. Banyard, J.D.H., J.X., C.L., D.K., A.M., N.H., D.L.S., J.B., C.C. and B.Y.L. conducted experiments; C.R.B., J.K., A. Brown, A. Banyard and C.J. analysed data; A. Banyard, C.C., V.H.-G., L.S., J.A.E., B.S., X.Z., D.L.S., D.K., J.A., G.A. and C.H. provided technical support; J.P.M. maintained the genetically engineered mouse models and provided murine samples; J.P.M., L.G.G., J.A.E. and B.S. provided reagents and cell lines; M.A.G., J.G., L.F. and D.A.O. helped with clinical sample collection; L.F. provided pathological support; C.R.B. and C.J. wrote the paper and C.J. and L.G.G. oversaw the project. J.X. contributed to this work while an employee at CRUK MI.

Competing interests
L.G.G. has patent application pending related to the hydrogel system. The rest of the authors have no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41563-021-01085-1.
Correspondence and requests for materials should be addressed to Linda G. Griffith or Claus Jørgensen.
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- □ □ For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
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Software and code

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Data collection

Orbitrap Elite mass spectrometer (Thermo) operated under Thermo XCalibur (V4.1.31.9) was used to acquire IAC LC-MS/MS data.
Orbitrap Velos Pro (Thermo) operated under Thermo XCalibur was used to acquire COM LC-M-MS data.
QExactive HFX (Thermo) operated under Thermo XCalibur was used to acquire matrisome and PEG mPCO LC-MS/MS data.
PerkinElmer Opera Phenix microscope and Harmony 4.8 (Perkin Elmer) software was used to acquire confocal images.
Leica SP8 TCS inverted confocal microscope and LAS-X software suite (V3.5.5) (Leica) was used for second harmonic generation imaging.
Li-COR Odyssey and Li-COR ImageStudio (V 5.2) was used for western blot imaging.
Bruker Catalyst with a Nanoscope V controller (Bruker UK Ltd., Coventry UK) operating under nanoscope software (V9.1, Bruker) was used for AFM indentations.
FEI Tecnai 12 Biotwin microscope operated under Tecnai User Interface software (V1.9) was used to acquire EM images.
Helios mass cytometer (Fluidigm) operated under the CyTOF software (V7 Fludigm) was used to acquire CyTOF data.
Aria III SORP sorter (BD Biosciences) operated under BD Diva software (8.0) was used for FACS sorting.
Leica SCN-400 operated under the Leica SCN-400 software suite (V 2.2.0.3789) was used to scan histology slides.
Olympus VS-120 operated under the Olyvia software (V2.9, OLYMPUS) was used to scan IF slides.
QuantStudio 5 Real-Time-PCR instrument (Applied Biosystems, Thermo) operated under the QuantStudio Design & Analysis software (V1.5.1, 2016, Applied Biosystems, Thermo) was used for RT-qPCR analysis.

Data analysis

Proteome Discoverer (V2.1, ThermoFisher) was used to analyze qualitative LC-MS/MS data (IAC, COM, 3D-PEG LC-MS/MS).
Progenesis (V26.45.1656, Nonlinear Dynamics) was used to analyze qualitative LFQ matrisome LC-MS/MS data (matrisome datasets).
Harmony (V4.8, PerkinElmer) was used to analyze adhesion and high-content screening data and live imaging videos.
Imaris (V9.5, Bitplane) was used for 3D analysis of co-culture images.
ImageStudio Lite (V 5.2, Li-COR) was used to analyze western blot images.
ImageScope (V12.3.3, Leica) was used to analyze histology slides.
Olyvia (V2.9, OLYMPUS) has been used to acquire IF images.
QuPath (V0.2.3) was used to analyze IF images and export high-quality IF images from scanned slides.
QuantStudio Design & Analysis software (V1.5.1, 2016, Applied Biosystems, Thermo) was used to analyze and export RT-qPCR data.
CT-FIRE (V2.0, open source) was used to analyze SHG images. Fiji (V. 1.52e, NIH) was used to analyze microscopy images including IF and brightfield. NanoScope Analysis software (V4.8, Bruker) was used to analyze AFM data. Cytoscape (V3.8.2, National Resource of network biology) was used for network analysis using the plugins: MCODE (V2.0.0), String app (V1.6.0), boundarylayout (V1.1). Cytobank.org and Cytofkit2 package (version 2.0.1) were used to analyze Mass Cytometry data. Prisom GraphPad 8 (GraphPad Software Inc) was used for statistical analysis and data visualization. Cytobank.org and Cytofkit2 package (version 2.0.1) were used to analyze Mass Cytometry data.

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All original source data are freely available: The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifiers: NP matrisome atlas (PXD022555 and 10.6019/PXD022555); IAC datasets (PXD022487 and 10.6019/PXD022487); CDM datasets (PXD022509 and 10.6019/PXD022509); 3D PEG CBF-0.5 LC-MS (PXD022520 and 10.6019/PXD022520); Tumour Matrisome LC-MS (PXD022767 and 10.6019/PXD022767). Raw CyTOF data, IF images and AFM force curves as well as source data for all figures (Figures 1-5 and Supplemental Figures 1-28) have been deposited to https://www.zenodo.org/ (10.5281/zenodo.4664132). Clinical information and gene-expression data for human pancreatic cancer (TCGA-PAAD) were downloaded from cBioPortal (https://www.cbioportal.org, accessed: 22/04/2020).

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| Data exclusions | No data was excluded from the study. |
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| Randomization | Samples were prepared, treated, processed and analysed in random order. |
| Blinding | Blinding was conducted during the pathological scoring on murine pancreatic cancer samples by a trained pathologist who was blinded to genotype, stage and/or data generated from any animal in this study. The remaining experiments were not blinded as the investigators who set up the experiments analysed the data, which is incompatible with blinding. |

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- Animals and other organisms
- Human research participants
- Clinical data

**Methods**

- ChIP-seq
- Flow cytometry
- MRI-based neuroimaging

**Antibodies used**

**Antibodies used for Immunofluorescence/immunohistochemistry:** OCT embedded samples (fresh frozen) for Immunofluorescence:
- rabbit anti-ZO1 (1:100, Invitrogen, 61-7300)
- rat anti-CD49f (1:500, Merck Millipore, MAB1378, clone NKI-GoH3)
- rabbit anti-E-Cadherin (1:100, Cell Signalling, 3195, Clone 24E10)
- rabbit anti-Sox9 (1:300, Merck Millipore, AB5535)
- rabbit anti-Yap1 (1:50, Cell Signaling Technology, 4912s)
- rabbit anti-Laminin (1:50, abcam, ab11575)
- Alexa-488 conjugated donkey anti-Rabbit (1:400, Invitrogen, A21206)
- Alexa-488 conjugated donkey anti-Rat (1:400, Invitrogen, A21208)
- Alexa-647 conjugated goat anti-Rat (1:400, Invitrogen, A21247)

**OCT embedded murine samples (fresh frozen) for immunohistochemistry:**
- rabbit anti-F4/80 (1:100, Cell Signalling Technology, 70076)
- rabbit anti-PanCK (1:100, Abcam, ab9377)
- rat anti-CD45 (1:100, BioLegend, 550539, clone 3D-F11)
- rabbit anti-Podoplanin (1:500, Thermo Fisher, MA5-29742)
- HRP conjugated goat anti-Rabbit secondary antibody (RTU, Dako, K4003)

**OCT embedded human samples (fresh frozen) for immunohistochemistry:**
- rabbit anti-ITGA3 (1:1000, Merck Millipore, AB1920)
- rat anti-ITGAV (1:1000, Merck Millipore, MAB1378, Clone NKI-GoH3)
- rabbit anti-ITGA2 (1:2000, Merck Millipore, AB1936)
- mouse anti-ITGB1 (1:3000, Abcam, ab16821)
- rat anti-ITGB1 (1:3000, Biolegend, 550531, Clone 9E67)
- mouse anti-ITGAI (1:250, Merck Millipore, MAB2059, Clone ASC-8)
- mouse anti-DLAM (1:1000, Merck Millipore, MAB1924, Clone 4C7)
- mouse anti-PanCK (1:5000, Sigma Aldrich, C2931)

**Formalin fixed murine samples for immunohistochemistry (in addition to antibodies previously mentioned):**
- rabbit anti-F4/80 (1:100, Cell Signalling Technology, 70076)

**Function-blocking antibodies:**
- rabbit anti-ITGA6 (10 ug/mL, Merck Millipore, MAB1378, Clone NKI-GoH3)
- mouse anti-ITGA3 (10 ug/mL, ThermoFisher, 17-0494042, Clone PIBS)

**Antibodies for Immunoblotting:**
- mouse anti-LAMAS (1:1000, Mybiosource MBS20658)
- rabbit anti-GAPDH (1:5000, Abcam, ab181602)
- Dylight 800 (1:15,000, Cell Signalling Technologies) or 680 (1:15,000, Cell Signalling Technologies) 4xPEG pe-conjugated secondary antibodies

**Antibodies used for Flow cytometry:**
- brilliant violet 421 (BV-421) conjugated anti-CD45 (BioLegend Clone 30-F11, #103133)
- BV-421 conjugated CD-31 (BioLegend, clone MEC13.3, #102423)
- BV-421 conjugated Anti-EpCAM (BioLegend clone G8.8, #118225)
- Phycoerythrin (PE) conjugated CD-90 (Thermo, clone G7, #A14726)
- PE-Cyanine7 (PE-Cy7) conjugated anti-podoplanin (PDNP, BioLegend, clone 8.1.1, #127471)

**Antibodies used for Mass Cytometry:**
- CD68 (BioLegend #137002, Clone FA-11)
- CD3 (CST, #9579, Clone O3E9)
- CD11b (eBioScience, 14-0112-81, Clone MI/70, # 14-0112-82)
- ITGA2 (BioLegend, 103501, Clone HMa2)
- Podoplanin (BioLegend, 127402, Clone 8.1.1)
- Pan Cytokeratin (BioLegend, 628602, Clone C10)
- Vimentin (CST, #5741, Clone 023H3)
- Desmin (abcam, ab216616, Clone Y66)
Validation

All antibodies used in this study have been verified either by the direct manufacturer or a competing manufacturer for the exact same clone for the application for which this antibody has been used in this study. Immunoblotting: LamA5 (Mybiosource MBS20658) species reactivity: mouse. (Yun et al., Cell Death Dis 2014 5(2):e1049). Gapdh (ab181602) species reactivity: mouse (Lu et al. EMBO Mol Med 2021 13(1):e12798). IF and IHC: Zo1 (Invitrogen, 61-7300) species: mouse, human (Jorgens et al J Cell Sci 2017 1:177-189). CD49f (MAB1378, clone NKI-GoH3) species: human, mouse. E-Cadherin (Cell Signalling, 3195, Clone 24E10) species: mouse, human (Nikolopoulou et al Nat Commun 2019 10(1):2487). Sox9 (Merck Millipore, AB5535) species: mouse, human. Laminin (ab11575) species: human, mouse (Bisi et al., Nat. Commun. 2020 11(1):3516). F4/80 (Cell Signalling Technology, 70076) species: mouse (Lorschid et al JCI Insight 2019 4(22):e130835). PanCK (14-094042) species: mouse, human. Podoplanin (MAB2059, Clone ASC-8) species: human, mouse. Yapl (Cell Signaling Technology, #4912s) species: mouse, the specificity of the antibody was validated against isotype control (Rabbit IgG Control antibody (Vector Laboratories, I-1000-5). Itga3 (AB1920) species: human. Itga6 (MAB1378, Clone NKI-GoH3) species: human. Itga2 (AB1936) species human. Itgav (ab16821) species: human. Itgb1 (Clone 9EG7) species: human. Itgb4 (MAB2059, Clone ASC-8) species: human. LamA5 (MAB1924, Clone 4C7) species human. PanCK (C2931) species human, mouse. For IF/IHC applications, the antibody signal has been compared against a non-primary antibody control and only those antibodies with a significantly higher signal/noise and background ratio as well as correct localization have been used in this study.

Function blocking: Itga6 (MAB1378, Clone NKI-GoH3) species: human, Itga3 (17-049042, Clone PIBS) species: human. Samarelli, A. V. et al. Journal of Biological Chemistry, (2014) or Aumailley, M., Timpl, R. & Sonnenberg, Experimental Cell Research (1990).

For Mass Cytometry and flow cytometry applications, only antibodies reported by Fluidigm or manufacturers to be suitable for mass or flow cytometry respectively have been used in this study (Hutton et al Cancer Cell 2021 doi:10.1016.jccell.2021.060017 in press). CD68 (BioLegend #137002, Clone FA-11) species: mouse. cC3 (CST, #9579, Clone 03E9) species: mouse. CDllb (eBioScience, 14-0112-81, Clone M1/70, # 14-0112-82), species: mouse. ITGA2 (BioLegend, 103501, Clone HMa2), species: mouse. Podoplanin (BioLegend, 127402, Clone 8.1.1), species: mouse. Pan Cytokeratin (BioLegend, 628602, Clone CII), species: mouse. Vimentin (CST, #5741, Clone 01H5) species: mouse. Desmin (abcam, ab216616, Clone Y66) species: mouse. aSMA (ThermoFisher, 14-9760-82, Clone A4), species: mouse. Ki67 (ThermoFisher, 14-5698-82, Clone: SolA15), species: mouse. ITGA6 (Merc Millipore, MAB1378, Clone: NKI-GoH3), species: mouse. EpCam (BioLegend, 118202, Clone G8.8), species: mouse. CD206 (BioLegend, 141702, Clone Co68C2), species: mouse. Podoplanin (BioLegend, 127402, Clone 8.1.1), species: mouse. GFP (BioLegend, 338002, Clone FM264G). CD45 (BD Biosciences, 550994, Clone 30-FL), species: mouse. pRb (BD Biosciences, 558389, Clone J12-906), species: mouse. MHC Class 2 (eBioScience, 14-5321-81, Clone M5/114.15.2), species: mouse.
Animals and other organisms

Policy information about studies involving animals: ARRIVE guidelines recommended for reporting animal research

Laboratory animals
KPC (KrasLSL-G12D/+; Trp53LSL-R172H/+; Pdx-1-Cre), KPF (KrasFrt-S-Frt-G12D/+; Trp53LSL-Frt; Pdx-1-Flp), KC (KrasLSL-G12D/+; Pdx-1-Cre) and KF (KrasFrt-S-Frt-G12D/+; Pdx-1-Flp) animals.
Mice were maintained in purpose-built facility in a 12 hr light/dark cycle at uniform temperature and humidity with continual access to food and water. Age and gender of the laboratory animals are detailed in the source data (https://zenodo.org/record/4664132).

Wild animals
The study did not involve wild animals.

Field-collected samples
The study did not involve field collected samples.

Ethics oversight
All animal experiments were reported in accordance with the 'Animals (Scientific Procedures) Act of 1986' under project license No. 70/8745 (CRUK Manchester Institute) and 70/8375 (University of Glasgow). Conducted experiments with laboratory animals adhered to Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines and were subjected to constant review by the Animal Welfare and Ethical Review Boards of the Cancer Research UK Manchester Institute and the University of Glasgow (UoG).

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Human research participants

Policy information about studies involving human research participants

Population characteristics
For human patient-derived organoids and samples used for IHC/AFM, patient information and characteristics are disclosed in Supplementary Data 1.
For expression analysis in human PDAC patients, data was obtained from the cancer genome atlas (TCGA) using cBioPortal (https://www.cbioportal.org/, Cerami et al., 2012 & Gao et al., 2013). For population characteristics patient and clinical annotations for the PAAD:TCGA cohort were utilized.

Recruitment
Research samples were obtained from the Manchester Cancer Research Centre (MCRC) Biobank with informed patient consent obtained prior to sample collection.

Ethics oversight
The MCRC Biobank is licensed by the Human Tissue Authority (license number: 30004) and is ethically approved as a research tissue bank by the South Manchester Research Ethics Committee (Ref: 07/H1003/161+5). The role of the MCRC Biobank is to distribute samples. For more information see www.mcrc.manchester.ac.uk/Biobank/Ethics-and-Licensing.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Flow Cytometry

Plots

Confirm that:
- The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
- The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).
- All plots are contour plots with outliers or pseudocolor plots.
- A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation
For fluorescent activated cell sorting (FACS) of murine PaFs, a cell suspension was generated by gently lifting the cells using Accutase (Thermo Fisher Scientific) for 5 min at 37°C and following neutralization with DMEM 10% v/v FBS and centrifugation, PaFs were re-suspended in PBS supplemented with 1% v/v FBS, hereafter termed FACS buffer (FB). Cells were then washed and re-suspended in FB supplemented with Fc-block and subjected to surface staining with brilliant violet 421 (BV-421) conjugated anti-CD45 (1:20, BioLegend Clone 30-F11); BV-421 conjugated CD-31 (1:20, BioLegend, clone MEC13.3); BV-421 conjugated Anti-EpCAM (1:20, BioLegend clone G8.8); Phycoerythrin (PE) conjugated CD-90 (1:20 Thermo, clone G7); PE-Cyanine7 (PE-Cy7) conjugated anti-podoplanin (PDPN, 1:20, BioLegend, clone 8.1.1) for 30-45 min on ice, shaking. Following staining, excess antibody was removed using FB, cells were separated using a 70 μm filter (BD) to remove cell-clumps, re-suspended at a concentration of 5 x 10^6 cells/mL in FB and sorted.

For FACS of murine rFP-labelled mPDOs, cells were prepared as single cell suspension using TrypLE, filtered using a 70 μm cell strainer (BD) and retained in AdF Base medium till sorting. rFP-labelled mPDOs were sorted using an BD®Aria 3 SORP (BD) sorter using the 640 laser and 730/45 filter set.

Instrument
Aria III SORP sorter (BD Biosciences)
| Software          | BD Diva (BD Biosciences) |
|-------------------|--------------------------|
| Cell population abundance | Only highly positive cells for the selected markers isolated for FACS sorts. The sort was repeated after 7 days of growth to obtain a pure cell population. |
| Gating strategy   | For FACS of murine PaFs, a single-cell gate was first applied and cells were further sorted for BV421- and PE+/PE-Cyanine7+. For FACS purification of iRFP+ mPDOs, a single cell gate was applied and sorted for iRFP (640)+. For CyTOF experiments, the gating strategy has been uploaded with the raw data to https://www.zenodo.org/ under the identifier 10.5281/zenodo.4303962 and can be found in folder: U_2020_F5e-f,SF26,SF27_CyTOF_data.zip/Gating_strategy.pdf. |

☐ Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.