Organoid Models of Human and Mouse Ductal Pancreatic Cancer


1Hubrecht Institute, Royal Netherlands Academy of Arts and Sciences (KNAW), University Medical Centre Utrecht and CancerGenomics.nl, 3584 CT Utrecht, the Netherlands
2Foundation Hubrecht Organoid Technology (HUB), 3584 CT Utrecht, the Netherlands
3Cold Spring Harbor Laboratory, Cold Spring Harbor, NY 11724, USA
4Lustgarten Foundation Pancreatic Cancer Research Laboratory, Cold Spring Harbor, NY 11724, USA
5Graduate Program in Molecular and Cellular Biology, Stony Brook University, Stony Brook, NY 11794, USA
6Rubenstein Center for Pancreatic Cancer Research, Memorial Sloan Kettering Cancer Center, New York, NY 10065, USA
7Weill Medical College at Cornell University, New York, NY 10065, USA
8Watson School of Biological Sciences, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY 11724, USA
9Graduate Program in Genetics, Stony Brook University, Stony Brook, NY 11794, USA
10Department of Pathology, University Medical Centre Utrecht, 3584 CX Utrecht, the Netherlands
11Department of Surgery, University Medical Center Utrecht, 3584 CX Utrecht, the Netherlands
12Department of Pathology, Memorial Sloan Kettering Cancer Center, New York, NY 10065, USA
13The Sol Goldman Pancreatic Cancer Research Center, Johns Hopkins University School of Medicine, Baltimore, MD 21231, USA
14Co-first author
15Present address: Gurdon Institute-University of Cambridge, Tennis Court Road, Cambridge CB2 1QN, UK
*Correspondence: h.clevers@hubrecht.eu (H.C.), dtuveson@cshl.edu (D.A.T.)
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SUMMARY

Pancreatic cancer is one of the most lethal malignancies due to its late diagnosis and limited response to treatment. Tractable methods to identify and interrogate pathways involved in pancreatic tumorigenesis are urgently needed. We established organoid models from normal and neoplastic murine and human pancreas tissues. Pancreatic organoids can be rapidly generated from resected tumors and biopsies, survive cryopreservation, and exhibit ductal- and disease-stage-specific characteristics. Orthotopically transplanted neoplastic organoids recapitulate the full spectrum of tumor development by forming early-stage neoplasms that progress to locally invasive and metastatic carcinomas. Due to their ability to be genetically manipulated, organoids are a platform to probe genetic cooperation. Comprehensive transcriptional and proteomic analyses of murine pancreatic organoids revealed genes and pathways altered during disease progression. The confirmation of many of these protein changes in human tissues demonstrates that organoids are a facile model system to discover characteristics of this deadly malignancy.

INTRODUCTION

Mortality due to pancreatic cancer is projected to surpass that of breast and colorectal cancer by 2030 in the United States (Rahib et al., 2014; Siegel et al., 2013). This dire scenario reflects an aging population, the improvement of outcomes for breast and colorectal cancer patients, the advanced stage at which most patients with pancreatic cancer are diagnosed, and the lack of durable treatment responses in pancreatic cancer patients. Indeed, effective therapeutic strategies for patients with pancreatic ductal adenocarcinoma (PDAC) have been difficult to identify (Abbruzzese and Hess, 2014).

The therapeutic resistance of PDAC has been explored in a variety of cell culture and animal model systems, with clinically actionable findings encountered only occasionally (Villarroy et al., 2011). Patient-derived xenografts (PDXs) have yielded insights into PDAC, but their generation requires a large amount of tissue, and they take multiple months to establish (Kim et al., 2009; Rubio-Viqueira et al., 2006). Genetically engineered mouse models (GEMMs) of PDAC have also been generated as a parallel system for fundamental biological investigation and preclinical studies (Perez-Mancera et al., 2012). These GEMMs...
accurately mimic the pathophysiological features of human PDA, including disease initiation from preinvasive pancreatic intraepithelial neoplasms (PanINs) (Hingorani et al., 2003; Pérez-Mancera et al., 2012) and were used to discover that PDA possesses a deficient vasculature that impairs drug delivery (Erkan et al., 2009; Jacobetz et al., 2013; Koong et al., 2000; Olive et al., 2009; Provenzano et al., 2012). Although GEMMs have informed PDA therapeutic development (Beatty et al., 2011; Frese et al., 2010; Provenzano et al., 2012), they are expensive and time consuming (Pérez-Mancera et al., 2012). In addition, both human PDA and GEMMs exhibit an extensive stromal component that decreases the neoplastic cellularity, making it difficult to isolate and characterize the epithelium-derived malignant cells in pancreatic neoplastic tissues.

To study neoplastic cells, dissociated human tumors are often grown in two-dimensional (2D) culture conditions (Sharma et al., 2010), which do not support growth of untransformed, nonneoplastic pancreatic cells. Three-dimensional (3D) culture strategies have been developed to study normal, untransformed cells but so far have only allowed minimal propagation (Agbunag and Bar-Sagi, 2004; Lee et al., 2013; Means et al., 2005; Rovira et al., 2010; Seaberg et al., 2004). A comprehensive 3D cell culture model of murine and human PDA progression would facilitate investigation of genetic drivers, therapeutic targets, and diagnostics for PDA.

To address this deficiency, we sought to generate normal and neoplastic pancreatic organoids by modifying approaches we previously pioneered to culture intestinal (Sato et al., 2009), gastric (Barker et al., 2010), colon carcinoma (Sato et al., 2011), hepatic (Huch et al., 2013b), pancreatic (Huch et al., 2013a), and prostatic organoids (Gao et al., 2014; Karthaus et al., 2014). We developed 3D organoids from normal and malignant murine pancreatic tissues and used this model system to investigate PDA pathogenesis. Pancreatic organoids derived from wild-type mice and PDA GEMMs accurately recapitulate physiologically relevant aspects of disease progression in vitro. Following orthotopic transplantation, organoids from wild-type mouse normal pancreata are capable of regenerating normal ductal architecture, unlike other 3D model systems. We further developed methods to generate pancreatic organoids from normal and diseased human tissues, as well as from endoscopic needle biopsies. Following transplantation, organoids derived from murine and human PDA generate lesions reminiscent of PanIN and progress to invasive PDA. Finally, we demonstrate the utility of organoids to identify molecular pathways that correlate with disease progression and that represent therapeutic and diagnostic opportunities.

RESULTS

Murine Pancreatic Ductal Organoids Expressing Oncogenic Kras Recapitulate Features of PanINs

Recently, we derived continuously proliferating, normal pancreatic organoids from adult murine ductal cells (Huch et al., 2013a). We optimized this approach to generate models of PDA progression. We manually isolated small intralobular ducts and established organoid cultures from C57Bl/6 mouse normal pancreata and pancreatic tissues that contained low-grade murine PanIN (mPanIN-1a/b) from Kras^+/LSL-G12D, Pdx1-Cre ("KC") mice (Figure 1A). KC mice develop a spectrum of preinvasive ductal lesions that mirror human PanINs and, upon aging, stochastically develop primary and metastatic PDA (Hingorani et al., 2003). Ducts from KC pancreata were often larger and exhibited higher grades of dysplasia compared to those from wild-type mice (Figure 1A). After 1–3 days in culture, organoid growth was observed from isolated ducts (Figure 1A). We created a collection of 10 murine normal (mN) and 9 PanIN (mP) organoid cultures that we have continuously propagated for over 20 passages and successfully cryopreserved (Table S1A available online). mP organoids exhibited recombination of the conditional Kras^+/LSL-G12D allele and higher levels of Kras-GTP when compared to mN organoids (Figure 1B).

To determine the contribution of different pancreatic lineages to the organoids, we evaluated the expression of pancreatic lineage markers in these cultures. Genes associated with the ductal lineage (Ck19 and Sox9) (Cleveland et al., 2012) were enriched in the mN and mP organoids compared to total pancreatic tissues, which contain relatively few ductal cells (Figure 1C). In addition, the mP organoids upregulated genes indicative of a PanIN disease state (Muc5ac, Muc6, and Tff1) relative to mN, with no difference in Klf4 (Figure 1D) (Prasad et al., 2005). GFP-transduced mN and mP organoids were orthotopically transplanted into syngeneic C57Bl/6 or Nu/Nu mice. mN organoids quickly formed ductal structures comprised of simple cuboidal cells that persisted for up to 1 month (n = 9/27 transplants) but were not observed after 2 months (n = 0/13 transplants) (Figure 1E and Table S1B). In comparison, mP organoids formed small cysts lined with a single layer of simple cuboidal ductal cells interspersed with mucin-containing columnar epithelial cells. Although we could not demonstrate that the mP transplants were contiguous with the native ductal system, they resembled preinvasive mPanIN (Figure S1C). These dysplastic epithelial cells persisted for 2 months or longer (n = 16/18 transplants), were GFP and Ck19 positive, expressed the mPanIN-associated mucin Muc5ac, and stained prominently with Alcian blue (Figure 1E and Table S1C). In addition, when compared to mN transplants, mP transplants had increased proliferation and a robust stromal response, which are characteristics of autochthonous mPanIN tissue (Figures S1A–S1C). The ability of transplanted mP organoids to form lesions with many of the features of mPanINs demonstrates the utility of this system as a model for early pancreatic neoplasia.

Multiple cellular origins have been proposed for the development of PDA, with the pancreatic acinar cell hypothesized to be a major contributor to PDA initiation (De La O et al., 2008; Gidekel Friedlander et al., 2009; Guerra et al., 2003; Habbe et al., 2008; Kopp et al., 2012; Morris et al., 2010; Sawey et al., 2007). However, recent studies have suggested that transformation of pancreatic ductal cells can also give rise to PDA (Pylayeva-Gupta et al., 2012; Ray et al., 2011; von Figura et al., 2014). Acinar cells isolated from wild-type pancreata are unable to form organoids in our conditions (Huch et al., 2013a). Therefore, our pancreatic ductal organoid system offers a unique opportunity to determine whether ductal cells can give rise to mPanIN. To assess whether expression of oncogenic Kras in pancreatic ductal organoids is sufficient to induce mPanIN formation in vivo, we derived...
Figure 1. Oncogenic Kras<sub>G12D</sub> Expression in Pancreatic Ductal Organoids Is Sufficient to Induce Preinvasive Neoplasms

(A) Hematoxylin and eosin (H&E) staining of murine pancreatic tissue used to prepare organoids (top). Arrows indicate mouse normal or PanIN ductal structures. Ducts embedded in Matrigel immediately following isolation (middle) and organoids 3 days postisolation (bottom). Arrowheads mark isolated ducts and growing organoids. Scale bars, 50 μm.

(B) Immunoblots for Kras, pan Ras, Kras-GTP by RBD-GST pull-down, and Tubulin in mN and mPanIN (mP) organoids. PCR confirmation of Cre-mediated recombination of the Kras<sub>LSL-G12D</sub> allele (bottom).

(C) qRT-PCR of ductal (Pdx1, Ck19, Sox9, and Hnf6), acinar (Ptf1a, Cpa1, and Amy), and endocrine (Ngn3, Chga, and Ins2) lineage markers in mN and mP organoids. Means of three biological replicates are shown. Error bars indicate SEMs. Values were normalized to mouse normal pancreas.

(D) qRT-PCR of genes indicative of PanIN lesions (Muc5ac, Muc6, Tff1, and Klf4) in mN and mP organoids. Values were normalized to mN organoids. Means of three biological replicates are shown. Error bars indicate SEMs. **p < 0.01 by two-tailed Student’s t test.

(E) H&E, Alcian blue staining, and immunohistochemistry (IHC) of orthotopic, syngeneic transplants of GFP-transduced mN and mP organoids. Scale bars, 200 μm.

(F) Immunoblots for Kras, pan Ras, Kras-GTP by RBD-GST pull-down, and Tubulin in Kras<sup>+/LSL-G12D</sup> organoids transduced with adenoviral-Cre (Ad-Cre) or adenoviral-blank (Ad-Bl). PCR confirmation of Cre-mediated recombination of the Kras<sub>LSL-G12D</sub> allele (bottom).

(G) H&E, Alcian blue staining, and IHC of orthotopic syngeneic transplants of organoids transduced with Ad-Bl (Kras<sup>+/LSL-G12D</sup>, R26<sub>LSL-YFP</sub>) and Ad-Cre (Kras<sup>G12D</sup>, R26<sub>YFP</sub>) 2 weeks posttransplant. Scale bars, 200 μm.

See also Figure S1 and Table S1.
organoids from ducts harboring the conditional \textit{Kras}^{LSL-G12D} allele (Hingorani \textit{et al}., 2003). Following activation of \textit{Kras} by adenoviral-Cre (Ad-Cre) infection, \textit{Kras}^{G12D} organoids maintained expression of genes specific to ductal cells and not acinar or endocrine lineages (Figures S1D and S1E). Recombination of the \textit{Kras}^{LSL-G12D} allele was confirmed by PCR, and levels of GTP-bound \textit{Kras} were increased relative to control-infected organoids (Figure 1F). In addition, expression of \textit{Kras}^{G12D} resulted in the upregulation of genes associated with human PanIN (Figure S1F). The \textit{Kras}^{G12D}-expressing organoids demonstrated increased proliferation relative to control organoids (Figure S1G).

Finally, \textit{Kras}^{G12D} organoids formed mPanIN-like structures with columnar cell morphology when implanted orthotopically into syngeneic mice (Figure 1G). This morphology contrasted with the normal-appearing ductal architecture formed by transplanting \textit{Kras}^{+/LSL-G12D} organoids or wild-type m\textit{N} (Figures 1E and 1G). The ability of mPanIN-like structures to develop from \textit{Kras}^{G12D} expressing ductal organoids following transplantation demonstrates that ductal cells are also competent to form mPanINs.

**Tumor-Derived Organoids Provide a Model for Murine PDA Progression**

We prepared pancreatic ductal organoids from multiple murine primary tumors (mT) and metastases (mM) from KC and \textit{Kras}^{+/LSL-G12D}, \textit{Trp53}^{+/LSL-R172H}, Pdx1-Cre ("KPC") mice, which develop mPDA more rapidly than KC mice (Figures 2A and Table S2A) (Hingorani \textit{et al}., 2005). mT and mM organoids exhibited recombination of the \textit{Kras}^{LSL-G12D} allele, as well as increased levels of \textit{Kras}-GTP and \textit{Kras} protein (Figure 2B). mT and mM organoids had increased levels of S6 phosphorylation, but not of Erk or Akt phosphorylation (Figure 2B).

Orthotopic transplantation of mT organoids initially generated low- and high-grade lesions that resembled mPanIN (Figure 2C and Table S2B). Over longer periods of time (1–6 months), transplants developed into invasive primary and metastatic mPDA (Figure 2C and Table S2B). mT organoids grafted with a similar efficiency upon orthotopic transplantation in Nu/Nu mice (91.7%) compared to C57Bl/6 mice (85%), but disease progression was accelerated in Nu/Nu hosts (Table S2B). Although most mM organoid transplants required several months to progress from early mPanIN-like lesions to invasive and metastatic cancer (Figure 2C and Table S2B), mM organoids rapidly formed invasive mPDA within 1 month (Table S2C). The ability of organoid transplants to reproduce the discrete stages of disease progression contrasts with the rapid formation of advanced mPDA following transplantation of 2D cell lines (Figures S2A–S2C) (Olive \textit{et al}., 2009).

Tumors derived from transplanted mT and mM organoids exhibited prominent stromal responses and resembled autochthonous tumors from KPC mice (Figure S2A) (Olive \textit{et al}., 2009). This stromal response is often absent in tumors formed from 2D cell lines (Figure S2A) (Olive \textit{et al}., 2009). Low vascular density and high vessel-to-tumor distance were also observed, demonstrating the close resemblance of the organoid transplantation models to autochthonous mPDA, in contrast to transplanted 2D cell lines (Figures S2A–S2C) (Olive \textit{et al}., 2009).

Loss of heterozygosity (LOH) for Trp53 has been reported as a common feature of mPDA based on studies of 2D cell lines (Hingorani \textit{et al}., 2005). Therefore, we assayed for Trp53 LOH in our murine 3D organoids. All mT organoids prepared from KPC tumors maintained expression of p16, did not exhibit Trp53 LOH, and maintained a stable karyotype, whereas most mM organoids lost the wild-type Trp53 allele and were aneuploid (Figures 2D, 2E, and S2D). We generated 2D cell lines from mT and mM organoids but found that mM organoids were unable to propagate in 2D. mM-derived 2D cell lines exhibited Trp53 LOH and were aneuploid (Figures 2E and S2D).

To determine whether organoids are suitable for genetic cooperation experiments, shRNAs targeting p53 and p16/p19 were introduced into mM organoids (Figure S2E). Although the proliferation of mM organoids increased upon knockdown of either p53 or p16/p19 (Figure S2G), only p53 knockdown enabled 2D growth and colony formation (Figure S2F; data not shown). Also, only p53 knockdown promoted progression of mM organoid transplants to invasive carcinoma within 3 months (Figure S2H). This contrasts with a previous report that Kr\textit{a}s mutation and biallelic loss of p16/p19 promoted mPDA (Aguirre \textit{et al}., 2003; Bardeesy \textit{et al}., 2006) and may reflect differences in the genetic system or the initiating cellular compartment. Nevertheless, the cooperation between p53 depletion and oncogenic Kr\textit{a}s demonstrates that organoids are a facile system to evaluate genetic mediators of PDA progression.

**Human Pancreatic Organoids Model PanIN to PDA Progression**

We modified our culture conditions to support the propagation of human normal and malignant pancreatic tissues. Isolation of ductal fragments was not always feasible because some normal pancreatic tissue samples were predigestion in preparation for islet transplantation. Therefore, we directly embedded digested material into Matrigel. This approach achieved an isolation efficiency of 75%–80% for human normal (hN) organoids (Figures 3A and S3 and Table S3). hN organoids were aneuploid (Figures 2D, 2E, and S2D). We generated 2D cell lines from hN organoids but found that hN organoids were unable to propagate in 2D. All hN-derived 2D cell lines exhibited Trp53 LOH and were aneuploid (Figures 2E and S2D).

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Figure 2. Modeling Murine PDA Progression with Tumor- and Metastasis-Derived Organoids

(A) H&E staining of murine tissue from which tumor and metastasis organoids were derived (top). Arrowhead indicates metastasis. Scale bars, 50 μm. Digested murine tissues embedded in Matrigel immediately following isolation (middle) and organoids 3 days postisolation (bottom). Scale bars, 200 μm.

(B) Immunoblots of selected signaling effectors, Kras-GTP and Ras-GTP by RBD-GST pull-down, and tubulin. PCR confirmation of Kras<sup>LSL-G12D</sup> recombination in mP, mT, and mM organoids (bottom).

(C) H&E staining of tumors and metastases (Met) derived from mT organoid orthotopic transplants. Scale bars, 200 μm (top) and 50 μm (bottom).

(D) Loss of heterozygosity of the wild-type Trp53 allele determined by PCR (top) and immunoblot analysis of Trp53, Smad4, p16, and Tubulin. mM3L, derived from a liver metastasis.

(E) Karyotypes of organoids and monolayer (2D) cell lines.

See also Figure S2 and Table S2.
Figure 3. Human Pancreatic Ductal Organoids Recapitulate Features of Normal and Neoplastic Ducts

(A) Representative images (top) and H&E staining (middle) of human organoid cultures established from normal tissues (hN1-2), resected primary tumors (hT1-2), a resected metastatic lung lesion (hM1), and a fine-needle aspiration biopsy of a metastatic lesion (hFNA2). H&E staining of the resected tissues from which the organoids were derived (bottom). Scale bars, 500 μm (top), 250 μm (middle), and 500 μm (bottom).

(B) Representative images of hN and hT organoids cultured for 2 weeks (1 passage) in human complete media or in human complete media lacking the indicated factors. Scale bars, 500 μm.

(C) Number of passages hN and hT organoids could be propagated in the absence of the indicated factors. Scale bars, 500 μm.

(D) Targeted sequencing analysis of human organoids. Genes altered in more than one sample and/or known to be mutated in PDA are shown. If multiple mutations were found in a gene, only one mutation per gene is shown. Color key for the type of genetic alterations is shown. Met indicates organoids derived from metastatic samples.

See also Figure S3 and Tables S3 and S4.
organoids were predominantly and stably diploid (Figure 4B). The PDA-associated biomarker CA19-9 (Makovitzky, 1986) within 1 month (n = 9/12) (Figures 4D and S4A and Table S4D). (n = 2/23), whereas hT organoids efficiently generated a specific background of human cancer. The hN and hT organoids are therefore reflective of normal and neoplastic human pancreatic ductal cells and offer a model system that was not dissociated prior to suspension in Matrigel (Figures 3A and S3 and Table S3). This approach is broadly applicable to PDA patients and enables serial sampling.

Targeted sequencing of 2,000 cancer-associated genes was performed on hN and hT organoids. As expected, no mutations were detected in the hN organoid cultures. These analyses identified oncogenic KRAS mutations in the majority of tumor-derived samples (n = 8), as well as mutations in TP53 (n = 7), SMAD4 (n = 5), and CDKN2A (n = 4) (Figure 3D and Table S4). We also noted amplification of known oncogenes, such as MYC (n = 4), and loss of tumor suppressors, including TGFBR2 (n = 3) and DCC (n = 5). Importantly, the same KRAS mutations observed in several hT organoids were confirmed in the primary PDA from which they were derived (Table S4). The allele frequency of oncogenic KRAS variants in hT1–hT5 and hFNA2 ranged from ~50–100%. In contrast, the KRAS<sup>GT20D</sup> allele frequency in hFNA1 was only 1% (Table S4), which may result from coexistence of wild-type ductal cells. Although KRAS mutations were not detected in hT8 (Figure 3D and Table S4), the presence of mutations in known PDA genes (ARID1A and MLL3) suggests that hT8 contains malignant cells (Table S4).

To further characterize the cell types present in primary PDA organoids, we evaluated the expression of pancreatic lineage markers. hN and hT organoids expressed markers of ductal cells, but not other pancreatic lineages (Figure 4A). The karyotypes of hT organoids were highly aneuploid, whereas the hN organoids were predominantly and stably diploid (Figure 4B). The PDA-associated biomarker CA19-9 (Makovitzky, 1986) was also elevated in hT relative to hN organoids (Figure 4C). The hN and hT organoids are therefore reflective of normal and neoplastic human pancreatic ductal cells and offer a model system to explore pancreatic cancer biology in the more genetically complex background of human cancer.

Following orthotopic transplantation into Nu/Nu mice, hN organoids produced normal ductal structures at low efficiency (n = 2/23), whereas hT organoids efficiently generated a spectrum of low- and high-grade, extrudal PanIN-like lesions within 1 month (n = 9/12) (Figures 4D and S4A and Table S4D). The hT-derived transplants initially formed well-defined hollow lesions lined by a single layer of columnar epithelial cells with apical mucin and basally located, relatively uniform nuclei. The nuclei were small and lacked the pleomorphism and hyperchromasia often seen in invasive PDA. These lesions progressed over several months to infiltrative carcinoma comprised of poorly defined and invasive glands (Figures 4D and S4A and Table S4). A prominent desmoplastic reaction was present in hT-derived PanIN-like structures and PDA, including the deposition of a collagen-rich stroma and the recruitment of αSMA-positive cells (Figure 4B). The mutation or loss of TP53 or SMAD4 in hT1 and hT2 was also detected by IHC in these tumors (Figure S4C and Table S4). Overall, hT organoids represent a transplantable model of human pancreatic cancer progression.

**Gene Expression Analysis of Murine Pancreatic Ductal Organoids Implicates Candidate Genes in PDA Progression**

The mouse organoids were prepared from syngeneic mice, offering the ability to discern gene expression changes in organoids and determine whether these changes correlate with PDA progression. We harvested RNA from mN (n = 7), mP (n = 6), and mT (n = 6) organoids and generated strand-specific RNA-seq (RNA-seq) libraries. Sequences were mapped to the mm9 version of the mouse genome, and relative transcript abundances (transcripts per million) of 29,777 mouse genes were determined (Table S5). Principal component analysis revealed that mN organoids were distinct from mP and mT organoids (Figure 5A and Table S5).

Genes whose levels differed significantly among mN, mP, and mT organoids were identified. 772 genes were found downregulated and 863 genes upregulated in mP relative to mN organoids (Figure 5B and Table S5). When mT organoids were compared to mN organoids, 2,721 genes were downregulated and 2,695 were upregulated. In addition, 823 genes were downregulated and 640 genes were upregulated in mT relative to mP organoids. Distinct patterns of gene expression were found in the data set (Figure 5C). The majority of genes differentially expressed in mP relative to mN organoids changed in a similar manner in mT relative to mN organoids (Figure 5D). However, a much larger cohort of genes changed in expression in mT relative to mN than in mP relative to mN organoids (Figure 5D), suggesting that mP organoids represent an intermediate state between mN and mT organoids.

The glycosyltransferase Gcnt3 and putative protein disulfide isomerase Agr2 were among the most upregulated genes in both mP and mT organoids and have been demonstrated to be elevated in human PDA (Figure 5E) (Dumartin et al., 2011; Zhao et al., 2014). The most upregulated gene in both mP and mT relative to mN organoids was the acyl-CoA synthetase Acsm3 (Figure 5E). RNA-seq results were confirmed by qRT-PCR for 35 out of 40 genes (Table S5), including the upregulation of Agr2, Acsm3, Gcnt1, Gcnt3, and Ugdh and the downregulation of Ptprd in mP and mT organoids (Figure 5F and Table S5). Among the genes upregulated in mP and mT relative to mN organoids, Gcnt1, Gcnt3, Acsm3, Agr2, Syt16, Nt5e, and Ugdh were upregulated following the Ad-Cre-induced expression of oncogenic Kras<sup>GT20D</sup>, suggesting that these genes are activated downstream of mutant Kras<sup>GT20D</sup> (Figure S5A). To determine whether organoid RNA-seq profiles resembled gene expression patterns in vivo, we compared our organoid RNA-seq data to a published transcription profile of murine pancreatic tumors upon Kras<sup>GT20D</sup> inactivation (Ying et al., 2012). Genes differentially expressed upon inactivation of oncogenic Kras overlapped significantly with those up or downregulated in mP or mT relative to mN organoids (Figure S5B). These analyses demonstrate the ability of the organoid system to identify molecular alterations associated with PDA progression.
Proteomic Alterations in Murine Pancreatic Ductal Organoids Predict Pathways Associated with PDA Progression

As an orthogonal method to investigate molecular alterations in murine pancreatic organoids, we characterized the global proteomes of mN (n = 5), mP (n = 4), and mT (n = 5) organoids. Protein lysates were processed using amine-reactive isobaric tags for relative and absolute quantification (iTRAQ) mass spectrometry (Wiese et al., 2007). Samples were run in four 8-plex experiments and merged using an approach that normalizes the data to common samples included across all experiments (Extended Experimental Procedures). Upon merging, 6,051 unique protein isoforms were quantified in all samples. We applied linear regression modeling on the normalized intensity peak values and identified 710 protein isoform expression changes between mN and mP organoids (Figure 6A). 1,047 protein isoforms changed expression between mN and mT organoids, and 63 differentially expressed proteins were identified between mP and mT (Figure 6A). The relatively small number of protein expression changes identified between mP and mT organoids reflects their biological similarity (Figure S6A).

mN organoids showed unique proteomic profiles from their mP and mT counterparts (Figures 6B and 6C). To compare the proteomic and RNA-seq data, we collapsed the unique protein isoforms were quantified in all samples. We applied linear regression modeling on the normalized intensity peak values and identified 710 protein isoform expression changes between mN and mP organoids (Figure 6A). 1,047 protein isoforms changed expression between mN and mT organoids, and 63 differentially expressed proteins were identified between mP and mT (Figure 6A). The relatively small number of protein expression changes identified between mP and mT organoids reflects their biological similarity (Figure S6A).

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isoforms into their corresponding 4,155 genes. Some protein expression changes (e.g., 123/150 for downregulated and 96/151 for upregulated mP proteins) did not reflect corresponding transcriptional changes, indicating that protein stability may play a role in cancer progression, particularly in mP organoids (Figure 6D). Nonetheless, the proteomic data validated many of the expression changes identified by RNA-seq (Figure 6D), including upregulation of Gcnt3, Agr2, and Ugdh (Table S6).

Additionally, of the 1,599 genes whose expression levels changed in mT relative to mN organoids that were measured by mass spectrometry, 301 (19%) showed corresponding protein changes (Figure 6D).

Gene Set Enrichment Analysis (GSEA) on the RNA-seq and proteomic data (Subramanian et al., 2005) revealed elevated expression of genes and proteins involved in glutathione metabolism and biological oxidations in mP relative to mN organoids (Figures 6E, S6B, and S6C and Table S7), which is consistent with elevations in reactive oxygen species metabolism previously reported in KrasG12D cells (DeNicola et al., 2011; Ying et al., 2012). Enrichment of proteins involved in glutathione metabolism was also found in mT relative to mN organoids (Table S7). Additionally, we identified a significant positive enrichment of proteins involved in the steroid biosynthesis, cholesterol biosynthesis, one carbon pool by folate, and pyrimidine metabolism pathways (Figures 6E, S6B, and S6C and Table S7), which is consistent with an earlier report (Ying et al., 2012). Similar pathways were enriched in mP relative to mN organoids (cholesterol biosynthesis, one carbon pool by folate, and pyrimidine
metabolism) (Figures S6B and S6C and Table S7), whereas fatty acid metabolism and TCA cycle/respiratory electron transport pathways were downregulated (Figure S6C and Table S7). The increase in anabolic and decrease in catabolic pathways suggest that complex alterations in fatty acid and nucleotide metabolism occur during PDA progression.

Interestingly, we also found broad upregulation of the nucleoporin family at both the RNA and protein levels in the mT relative to mN organoids (Figures 6E and 6F and Table S6). The individual nucleoporins NUP214, NUP153, and NUPL1 were previously identified in shRNA dropout screens in PDA cell lines (Cheung et al., 2011; Shain et al., 2013). Furthermore, amplification of NUP153 was detected in one human PDA cancer cell line, and elevation of NUP88 was detected in human primary PDA (Cheung et al., 2011; Gould et al., 2000; Shain et al., 2013). This systematic analysis of molecular alterations in pancreatic organoids implicates nuclear transport as a pathway correlated with pancreatic cancer progression.

In Vivo Mouse and Human Validation of Candidates Associated with PDA Progression in Organoids

To demonstrate that the mouse organoid culture system represents a biological resource for the accurate discovery of genes associated with PDA progression, we selected 16 genes...
upregulated in mT organoids for validation in primary tissue specimens by IHC and immunofluorescence (IF) (Figure 7A). These 16 genes included enzymes, membrane proteins, structural proteins, and secreted ligands, which could represent candidate biomarkers and therapeutic targets. Of the 14 antibodies that generated a detectable signal on murine pancreatic tissue sections, 13 antibodies confirmed the increased expression of the candidate protein in mPanIN and mPDA lesions in concordance with the RNA-seq and proteomic data (Figures 7A, 7B, and S7A). 11 of the 13 candidate antibodies were compatible for evaluation in human tissues, and 7 of these candidates were upregulated in human PDA when compared to normal pancreatic ductal tissues (Figures 7C, 7D, and S7B).

**Figure 7. Increased Levels of ACSM3, NT5E, and GCNT3 Correlate with Mouse and Human PDA Progression**

(A) IHC analysis of 14 candidate genes in mouse adjacent normal ducts, mPanIN and mPDA. Differential expression is indicated as – (negative), + (weak), ++ (moderate), or +++ (strong). Only the ductal component of the normal pancreas was scored.

(B) IHC analysis of Acsm3, Nt5e, and Gcnt3 in mouse normal ducts, mPanIN and mPDA tissues. Arrow indicates adjacent normal ducts in mPanIN tissues. Arrowhead indicates mPanIN or mPDA. Scale bars, 50 μm.

(C) IHC analysis of seven candidate genes in human normal pancreas, hT orthotopic transplants, and PDA tissues. Differential expression is indicated as – (negative), + (weak), ++ (moderate), or +++ (strong). Only the ductal component of the normal pancreas was scored.

(D) IHC analysis of ACSM3, NT5E, and GCNT3 in human normal pancreas and PDA tissues. Arrow indicates normal ducts, and arrowhead indicates PDA. Scale bars, 50 μm.

See also Figure S7.
in orthotopic transplants of HT organoids into Nu/Nu mice (Figure 7C). These results indicate that the organoid culture system accurately models PDA progression and can serve as a resource for the discovery and genetic dissection of pathways driving human pancreatic tumorigenesis.

**DISCUSSION**

We have established pancreatic organoids as a tractable and transplantable system to probe the molecular and cellular properties of neoplastic progression in mice and humans. In contrast to prior reports (Agbunag and Bar-Sagi, 2004; Kovács et al., 2010; Seaberg et al., 2004), our culture conditions prevent the rapid exhaustion of normal ductal cells in vitro and generate a normal ductal architecture following orthotopic transplantation. Importantly, the ability to passage and transplant both normal and neoplastic ductal cells enables a detailed analysis of molecular pathways and cellular biology that is not possible when neonatal pancreatic fragments are propagated in air-liquid interfaces or when induced pluripotent cells are employed (Agbunag and Bar-Sagi, 2004; Kim et al., 2013; Li et al., 2014). Our finding that nucleoporins are broadly upregulated in the neoplastic murine organoids, coupled with the known associations of nucleoporins to cell proliferation and cell transformation, presents a class of proteins to investigate in pancreatic cancer progression (Gould et al., 2000; Köhler and Hurt, 2010). Furthermore, the ability to systematically characterize human pancreatic cancer organoids that lack KRAS mutations, such as hT8, will reveal driver genes for PDA. Finally, because organoids can be readily established from small patient biopsies, they should hasten the development of personalized approaches for pancreatic cancer patients.

**EXPERIMENTAL PROCEDURES**

**Animals**

\( \text{Tp53}^{+/LSL-G12D}, \text{Kras}^{+/LSL-G12D}, \text{and} \text{Pdx1-Cre} \) strains in C57Bl/6 background were interbred to obtain \( \text{Pdx1-Cre; Kras}^{+/LSL-G12D} \) (KC) and \( \text{Pdx1-Cre; Kras}^{+/LSL-G12D}, \text{Tp53}^{+/LSL-R172H} \) (KPC) mice (Hingorani et al., 2005).

The \( \text{R26}^{SLE-YFP} \) strain was interbred to get the desired genotype. C57Bl/6 and athymic Nu/Nu mice were purchased from Charles River Laboratory and Jackson Laboratory. All animal experiments were conducted in accordance with procedures approved by the IACUC at Cold Spring Harbor Laboratory (CSHL).

**Murine Pancreatic Ductal Organoid Culture**

Detailed procedures to isolate normal pancreatic ducts have been described previously (Huch et al., 2013a). In brief, normal and preneoplastic pancreatic ducts were manually picked after enzymatic digestion of pancreas with 0.012% (w/v) collagenase XI (Sigma) and 0.012% (w/v) dispase (GIBCO) in DMEM media containing 1% FBS (GIBCO) and were seeded in growth-factor-reduced (GFR) Matrigel (BD). For tumors and metastases, bulk tissues were minced and digested overnight with collagenase XI and dispase and trypsin inhibitor (1 mg/ml, Sigma). Following digestion, cells were embedded in GFR Matrigel and cultured in human complete medium with the addition of PGE2 (1 \( \mu \)M, Tocris). Normal samples were processed as above, except that the collagenase digestion was done for a maximum of 2 hr in the presence of soybean trypsin inhibitor (1 mg/ml, Sigma). Following digestion, cells were embedded in GFR Matrigel and cultured in human complete medium with the addition of PGE2 (1 \( \mu \)M, Tocris).

Additional experimental details and methods can be found in the Extended Experimental Procedures.

**HUMAN PANCREATIC TUMOR AND NORMAL ORGANOID CULTURE**

Tumor tissue was minced and digested with collagenase II (5 mg/ml, GIBCO) in human complete medium (see below) at 37°C for a maximum of 16 hr. The material was further digested with TrypLE (GIBCO) for 15 min at 37°C, embedded in GFR Matrigel, and cultured in human complete medium (AdDMEM/F12 medium supplemented with HEPEs [1x, Invi-rogen], Glutamax [1x, Invitrogen], penicillin/streptomycin [1x, Invitrogen], B27 [1x, Invitrogen], Primocin [1 mg/ml, InvivoGen], N-acetyl-L-cysteine [1 mM, Sigma], Wnt3a-conditioned medium [50% v/v], Rspo1-conditioned medium [10% v/v, Bovine Kuro], Noggin-conditioned medium [10% v/v or recombinant protein [0.1 \( \mu \)g/ml, Peprotech], epidermal growth factor [EGF, 50 ng/ml, Peprotech], Gastrin [10 nM, Sigma], fibroblast growth factor 10 [FGF10, 100 ng/ml, Preprotech], Nicotinamide [10 mM, Sigma], and A83-01 [0.5 \( \mu \)M, Tocris]).

**ACCESSORY NUMBERS**

All RNA-seq data are available at Gene Expression Omnibus (GEO) under accession number GSE63348. The proteomic raw data are available at PeptideAtlas under accession number PASS00625. The targeted DNA-seq data are available at EMBL European Nucleotide Archive under the accession number ERP006373.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, seven figures, and seven tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2014.12.021.

**AUTHOR CONTRIBUTIONS**

S.F.B. initiated the project, developed the methods for isolating mouse and human organoids, and characterized human organoids (Figures 1A, 3, 4B, 4D, and S4C and Tables S3 and S4). C.-I.H. developed transplantation models for organoids and performed shRNA knockdown and histological and karyotypic analyses (Figures 1A, 1E, 1G, 2A, 2C–2E, 4D, 7, S1A–S1C, S2A–S2F, S2H, S4A, S4B, S7B, and Tables S1, S2, and S4). L.A.B. performed RNA-seq on mouse organoids and analyzed RNA-seq and proteomic data (Figures 5, 6, S5B, and S6 and Tables S5, S6, and S7). I.C.C. conducted proteomic evaluation of mouse organoids and analyzed proteomic data (Figures 6 and S6C). D.D.E. developed mouse organoid methods and evaluated CA19-9 levels in human organoids (Figures 1A, 2A, 4C, S3, and Table S6). V.C. developed human organoid methods, performed molecular analyses of organoids, and prepared material for DNA sequencing and sequencing of Kras (Figures 1C, 1D, S3, 3D, 4A, S1D–S1F, and S6A and Tables S3, S4, and S5).

M.J. performed and analyzed the DNA sequencing of human organoids (Figures 3D and Table S4). Mouse and human organoid preparation and characterization were performed by M.P.-S., H.T., M.S.S., T.O., D.O., A.H.-S., C.M.A.-A., M.L., E.E., B.A., M.E.F., G.N.Y., B.D., B.C., K.W., K.H.Y., Y.P., M. Huch, A.G., F.H.M.M., and S.D.L. Sequencing analyses were performed by Y.H., Y.J., M. Hammel, J.N., E.C., and R.V.B. Pathological analyses were performed by the University of Illinois at Chicago and University of Miami Miller School of Medicine. All human experiments were approved by the ethical committees of the University Medical Centre Utrecht or the IRBs of MSKCC, MDACC, WCMC, and CSHL. Written informed consent from the donors for research use of tissue in this study was obtained prior to acquisition of the specimen. Samples were confirmed to be tumor or normal based on pathological assessment.

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