Stabilized epithelial phenotype of cancer cells in primary tumors leads to increased colonization of liver metastasis in pancreatic cancer

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AUTHOR CONTRIBUTIONS
J.L.C. helped design the experimental strategy, performed experiments to generate the genetically engineered mouse models, analyzed the data related to the genetically engineered mouse models, performed culture experiments, collected data, helped with scRNA-seq sample collection and discussion, generated the figures, and helped write the manuscript. S.Y. helped generate the genetically engineered mouse models and helped conceptually design and perform the scRNA-seq experiments and analysis. P.C.d.S. helped with assay optimization, multiplex data analysis, sample collection, and manuscript editing. X.Z. helped generate the KPC;ST. S.B. generated the spatial analysis algorithms and data, developed the computer code for analyses, analyzed the data, and participated in discussions related to the spatial and survival analyses. K.M.M. helped with immunohistochemical staining and manuscript editing. A.R. conceived and designed the analytic strategy for spatial infiltration analysis, supervised S.B. toward the implementation of the analysis algorithms, provided intellectual input, and participated in discussions related to the spatial and survival analysis. J.K.B. helped conceive and design the analytic strategy for the Visiopharm analysis of the mouse immune multiplex, provided intellectual input, and participated in discussions related to the spatial analysis. A.D.R. provided the KPPC-Zeb1-flox mouse line. R.K. conceptually designed the strategy for this study, participated in discussions, provided intellectual input, supervised experimental discussion, and helped write the manuscript.

SUPPLEMENTAL INFORMATION
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DECLARATION OF INTERESTS
The authors declare no competing interests.
SUMMARY

Pancreatic ductal adenocarcinoma (PDAC) is therapeutically recalcitrant and metastatic. Partial epithelial to mesenchymal transition (EMT) is associated with metastasis; however, a causal connection needs further unraveling. Here, we use single-cell RNA sequencing and genetic mouse models to identify the functional roles of partial EMT and epithelial stabilization in PDAC growth and metastasis. A global EMT expression signature identifies ~50 cancer cell clusters spanning the epithelial-mesenchymal continuum in both human and murine PDACs. The combined genetic suppression of Snail and Twist results in PDAC epithelial stabilization and increased liver metastasis. Genetic deletion of Zeb1 in PDAC cells also leads to liver metastasis associated with cancer cell epithelial stabilization. We demonstrate that epithelial stabilization leads to the enhanced collective migration of cancer cells and modulation of the immune microenvironment, which likely contribute to efficient liver colonization. Our study provides insights into the diverse mechanisms of metastasis in pancreatic cancer and potential therapeutic targets.

Graphical Abstract

In brief

The necessity of partial epithelial to mesenchymal transition for cancer metastasis is unclear. Applying single-cell RNA sequencing and genetic models to stabilize the epithelial phenotype, Carstens et al. show a continuum of epithelial and mesenchymal pancreatic cancer cells that use different mechanisms to migrate, escape the immune system, and metastasize.
INTRODUCTION

Pancreatic ductal adenocarcinoma (PDAC) is a treatment-resistant, deadly disease with a 5-year survival rate of 9% and metastatic disease emerging in nearly 98% of patients (Siegel et al., 2019; Hidalgo, 2010). Such sobering statistics strongly urge an in-depth mechanistic analysis of PDAC metastatic disease. The acquisition of mesenchymal features by epithelial cells, traditionally called an epithelial-to-mesenchymal transition (EMT), has been studied in multiple cancers and correlates with metastasis and therapy resistance (Gao et al., 2019; Williams et al., 2019). EMT is not a simple binary process, either epithelial or mesenchymal, but instead a plastic continuum of partial EMT (pEMT) states between the epithelial and mesenchymal poles (Jolly et al., 2018; Kalluri and Weinberg, 2009; Lovisa et al., 2015, 2016; Brabletz et al., 2018; Aiello et al., 2018; Williams et al., 2019; da Silva-Diz et al., 2018; Yang et al., 2020). Such pEMT cells have been suggested as being important in cancer metastasis and drug resistance (Yang et al., 2020). The role of EMT in therapy resistance has been validated by multiple studies, but evidence for its requirement for metastasis is still evolving (Brabletz et al., 2018; Dongre and Weinberg, 2019; Williams et al., 2019; Liu et al., 2014).

It is well documented that EMT occurs in PDAC and is associated with cancer cell migration. The evidence includes gene expression analysis, live imaging, and cytology using *in vitro* systems, transplant models, genetically engineered mouse models (GEMMs), and analysis of human samples (Arumugam et al., 2009; Rhim et al., 2012; Von Burstin et al., 2009; Özdemir et al., 2014; Zheng et al., 2015; Chen et al., 2018; Krebs et al., 2017; Beerling et al., 2016; Reichert et al., 2018; Wang et al., 2019). However, histopathological and gene expression analyses of human PDAC samples show that the majority of metastatic PDACs are epithelial, with minimal evidence of intermediate morphologies (McDonald et al., 2012; Liu et al., 2014; Hayashi et al., 2020). Such observations with human PDAC samples were also recapitulated in studies with PDAC model systems (Liu et al., 2014; Zheng et al., 2015; Chen et al., 2018). Such studies suggest pEMT may not be critical for all forms of PDAC metastasis, which requires careful unraveling.

Previous studies demonstrated that the depletion of an EMT driver, either Snail or Twist, had no effect on PDAC metastasis in mice (Zheng et al., 2015). In mice lacking Zeb1, metastases were reduced in number, and the primary tumor and remaining metastatic nodules displayed a well-differentiated epithelial phenotype (Krebs et al., 2017). Furthermore, transplantation studies have shown that epithelial PDAC cells can migrate and metastasize, and the re-expression of the epithelial adherens junction protein, E-cadherin, in mesenchymal PDAC cells enhances their migratory ability (Liu et al., 2014). An alternative to EMT-mediated motility is collective epithelial cell migration (Berton et al., 2009; Friedl et al., 2012; Friedl and Wolf, 2010). This form of migration has been a hypothesized mechanism for the circulation of cancer cell clusters, which is associated with poor clinical outcomes (Aceto et al., 2014; Fabisiewicz and Grzybowska, 2017; Giuliano et al., 2018). These data support the notion that migration, invasion, and metastasis of cancer cells may adapt different mechanisms outside the classical pEMT paradigm. However, the quantitation of cancer cells along the pEMT spectrum and the impact on the metastasis of stabilized EMT states remains to be elucidated.
Here, we use single-cell RNA sequencing (scRNA-seq) analysis of epithelial, pEMT, and mesenchymal cancer cells in human and mouse PDACs to identify the entire spectrum of the EMT continuum. We further explored the impact of stabilized pEMT states on PDAC metastasis via the use of multiple GEMMs that exhibit a stabilized epithelial phenotype through the deletion of EMT-driving transcription factors (deletion of Snail and Twist together or deletion of Zeb1). We demonstrate that the stabilization of an epithelial phenotype in PDAC cells enhances liver metastasis. In addition, we provide evidence for collective migration and immune microenvironment modifications as possible processes for epithelial PDAC metastasis. Collectively, we demonstrate that metastasis in PDACs may involve diverse mechanisms.

RESULTS

Single-cell RNA-seq of human PDAC reveals 51 different cancer cell phenotypes across the EMT continuum

To examine the prevalence of different EMT states in human PDAC, we evaluated the single-cell RNA-seq (scRNA-seq) dataset of >57,000 individual cells from 27 PDAC patient primary tumors (Peng et al., 2019). The cancer cell clusters were identified from the total population based on an epithelial gene expression signature (Figures 1A and S1A). RNA expression values of all cancer cells were subjected to Markov affinity-based graph imputation of cells (MAGIC) smoothing and clustering (van Dijk et al., 2018) based on the 315 genes identified by Thiery and colleagues as representing a universal EMT gene signature (referred to here as the Thiery EMT signature) (Figure 1B) (Tan et al., 2014). The clustering clearly demonstrated 51 different clusters of EMT in this dataset (Figure 1B). The RNA expression values for epithelial genes, divided by mesenchymal genes, were used to generate an E/M score for each cluster to organize them across the EMT spectrum from epithelial to mesenchymal (Figures 1C, 1D, S1B, and S1C). The 51 clusters were further sub-grouped into epithelial, pEMT, or mesenchymal phenotypes based on mathematical cutoffs (see Method details; Figure 1D). Eighteen clusters (34.09%) were predominantly of an epithelial phenotype (clusters 1–18, referred to here as E1–E18), and 10 clusters (24.48%) were predominantly of a mesenchymal phenotype (clusters 42–51, referred to here as M1–M10) (Figure 1D). The majority of the cancer cells (41.44%) clustered into 23 different pEMT states (clusters 19–41, referred to here as P1–P23) (Figure 1D). Next, the scRNA-seq analysis was computed for each individual PDAC patient using the above-described E/M score. This analysis revealed that 18.52% of the patients exhibited a predominately epithelial phenotype, 66.67% of the patients showed mainly pEMT phenotypes, and 14.81% revealed a mesenchymal phenotype (Figures S1D–S1G; Table S1).

To evaluate the correlation between the E/M score and the survival of PDAC patients, we next queried The Cancer Genome Atlas (TCGA) pancreatic cancer database with the Thiery EMT signature. PDAC tumors were classified as either epithelial, pEMT, or mesenchymal, and compared to previously reported classifiers by Moffitt et al. (2015) (basal versus classical), Collisson et al. (2011) (classical versus quasimesenchymal), and Chan-Seng-Yue et al. (2020) (basal A/B versus classical A/B), using the University of California, Santa Cruz (UCSC) Xena browser. In the TCGA cohort, which comprises early-stage/resectable tumors,
none of the classifiers correlated with a survival advantage (Figure S1H). Despite this outcome in the TCGA cohort, we did observe a significant correlation between the Thiery classifier and the Collisson et al. (2011) (classical versus quasimesenchymal) and Chan-Seng-Yue et al. (2020) (basal A/B versus classical A/B) classifiers, which have been previously shown to predict survival (Table S1). These data show that the EMT program, demonstrated by the Thiery EMT signature, correlates with previously reported classifiers that suggest poorer patient outcome in the advanced tumor setting.

scRNA-seq of murine PDAC identifies 56 cancer cell phenotypes across the EMT spectrum

To further evaluate the EMT spectrum in murine PDAC, we conducted scRNA-seq of Pdx1-Cre; KrasLSL-G12D; p53R172H/+ (KPC) mice that develop clinically relevant primary and spontaneously metastasizing PDAC. Previous studies demonstrated a significant reduction in EMT upon the complete deletion of EMT-driving transcription factors Snail or Twist (Zheng et al., 2015). However, it remained unclear whether such manipulation affected the full spectrum of EMT states, or led to an incomplete EMT inhibition, or led to a compensation via the upregulation of other pEMT states not governed by either Snail or Twist (Brabletz et al., 2018).

To mechanistically address this issue, we generated a cancer cell-specific, dual deletion of both Snail and Twist by crossing SnailcKO; YFPLSL mice with KPC; TwistcKO; YFPLSL mice (KPC;ST) (Zheng et al., 2015). PDAC tumors from KPC and KPC;ST mice were subjected to scRNA-seq. Similar to the scRNA-seq analysis of human PDAC, murine cancer cells were clustered based on the expression of epithelial gene transcripts, coupled with the enrichment of captured cancer cells via the engineered YFP (yellow fluorescent protein) signal (Figures 2A and S2A). Using MAGIC smoothing and clustering algorithms (van Dijk et al., 2018) of cancer cells based on the mouse correlates of the 315 genes of the Thiery EMT signature (294 genes) (Tan et al., 2014) followed by the generation of E/M scores, we identified 56 clusters encompassing the EMT spectrum (Figures 2B–2D and S2B–S2E). The mesenchymal clusters were largely positive for the canonical EMT transcription factors with overlapping and unique populations (Figure S2F). We also confirmed our Thiery EMT signature analyses with a principal-component analysis (PCA) of the MAGIC algorithms (Figures S2G–S2J). Overall, the spectrum of phenotypes exhibited in the aggregated mouse tumors are similar to those observed in the human PDAC analysis (Figures 2D and 2E). As in the human analyses, the 56 clusters were further sub-grouped into epithelial, pEMT, or mesenchymal phenotypes based on mathematical cutoffs (see Method details; Figure 2D). Eighteen clusters (34.45%) were predominantly of an epithelial phenotype (E1–E18), 18 clusters (23.92%) were predominantly pEMT states (clusters 19–36, referred to here as P1–P18), and 20 clusters (41.62%) were predominantly of a mesenchymal phenotype (clusters 37–56, referred to here as M1–M20) (Figure 2D). The KPC PDAC cells predominately exhibited a mesenchymal phenotype (Figures 2E and S2B–S2J). The loss of both Snail and Twist in the KPC;ST mice significantly stabilized the epithelial phenotype of cancer cells with more representation of clusters on the epithelial end of the EMT spectrum (Figures 2E and S2B–S2J), proving that Snail and Twist can affect the entire spectrum of EMT in the cancer cells.
To further appreciate the emergence of cancer cells within the EMT spectrum in relation to disease progression in the KPC and KPC;ST mice, we next performed scRNA-seq of PDAC tumors from early- and late-stage PDAC. Early-stage KPC PDAC cells presented with predominantly epithelial cluster phenotypes (clusters 1–26) and progressed to overtly mesenchymal cluster phenotypes in the late-stage KPC PDAC tumors (clusters 38–56) (Figure 2E). While the early KPC;ST PDAC cells were comparable to the early KPC PDAC cells in showing predominantly epithelial cluster phenotypes (clusters 1–26), the late-stage KPC;ST PDAC cells significantly maintained/stabilized epithelial cluster phenotypes (clusters 1–26) due to the loss of EMT-promoting Snail and Twist in the cancer cells (Figures 2E and S2B–S2F). Trajectory and pseudotime analyses indicate that the early-stage cancer cells are similar between the models and then progress through late-stage epithelial phenotypes with branches that later evolve into pEMT and mesenchymal types (Figures S2K–S2M). This pattern of cancer cell evolution is concordant with recent studies of whole exon sequencing in patients (Hayashi et al., 2020). The scRNA-seq data demonstrate that the KPC PDAC tumors exhibit multiple states within the EMT spectrum as observed in human PDAC. A significant impact on the EMT spectrum and the suppression of pEMT and mesenchymal phenotypes can be observed, with a combined deletion of Snail and Twist, using the Thiery EMT signature of 294 murine gene transcripts. This clearly suggests that Snail and Twist can regulate multiple signaling pathways to influence epithelial plasticity and the EMT program, and compensatory mechanisms are not realized when they are concurrently deleted, highlighting their rate-limiting role in this regard.

Stabilized epithelial phenotype of PDAC cells enhances liver metastasis

We next evaluated the impact of a stabilized epithelial phenotype in the KPC;ST mice on PDAC progression and metastasis. The combined loss of Snail and Twist had no impact on overall survival (Figure 3A; Table S2), disease-free survival (Figure 3B), and primary tumor burden (Figure 3C). Primary tumor histology was unchanged between the groups (Figures 3D and 3E). The gene expression analysis of EMT markers Zeb1, Vimentin, and Col1a1 were significantly decreased in YFP sorted primary pancreatic cancer cell lines from KPC;ST mice, along with complete loss of Snail1 and Twist1 (Figure S3A). There was no significant difference in the gene expression of E-cadherin (Cdh1) but immunolabeling for YFP, E-cadherin (E-cad), and Cytokeratin 8 (CK8) protein expression showed a significant increase in the percentage of YFP+E-cad+ cells (Figures S3A and S3B). These data confirm previous reports that the protein expression of E-cadherin may not correlate with gene expression (Aiello et al., 2018). Furthermore, a significant decrease in the percentage of cells co-expressing YFP and Snail/Slug, Twist, Zeb1, vimentin, α smooth muscle actin (αSMA), or fibroblast-specific protein 1 (FSP1) was observed (Figures 3F and S3B). The Snail antibody used in this study binds both Snail and Slug (Snail2). Staining with an antibody specific for Slug revealed that Slug is non-nuclear, limited to the PanIN lesions, and did not decrease in the expression in KPC;ST mice (Figures 3F and S3B). These data show that a stabilized epithelial phenotype does not affect primary PDAC progression.

We next assessed the impact of the dual loss of Snail and Twist on PDAC metastasis. There was no change in the vascular dissemination of cancer cells, as measured by quantitating circulating YFP+ tumor cells (Figure S3C). Gross pathological assessment showed no
difference in ascites or macroscopic metastatic nodules on the lung, liver, or other peritoneal organs (diaphragm, kidney/adrenal gland, gut, peritoneum, spleen, or reproductive organs) (Table S3). Microscopic analyses revealed insignificant differences in the incidence of metastatic disease (the number of mice per cohort with at least one lesion per specified organ; Table S3) or the number of metastatic lesions per organ (Figure 3G). However, the cumulative metastatic area in the liver was significantly larger in the KPC;ST mice (Figure 3H). Similarly, there were significantly more disseminated single YFP+ in the livers of KPC mice (Figure S3D). This was not due to an increased proliferation within the metastases, suggesting mechanisms outside of growth rate for the increase in metastatic liver colonization (Figure S3E).

Next, we performed scRNA-seq on cancer cells isolated from the livers of late-stage KPC and KPC;ST mice and compared it to the scRNA-seq of the corresponding primary tumors. The KPC PDAC primary and their corresponding liver metastatic cancer cells were indistinguishable in regard to the spectrum of E/M phenotypes that were identified, suggesting a potential capacity of each cluster from the primary tumor to metastasize or exhibit potential plasticity of a given cluster to expand into all primary tumor-associated clusters at the metastatic site (Figures 3I, S2B, and S2C). This is consistent with previous studies reporting similar mutation and expression profiles of primary and metastatic cancer cells in PDAC patients (Hayashi et al., 2020). However, the metastatic KPC;ST cancer cells isolated from the liver displayed an enrichment for unique E/M phenotypes not represented in KPC primary or metastatic tumors or the KPC;ST primary tumor (clusters 30, 32, and 34). Gene set enrichment analysis (GSEA) (Subramanian et al., 2005; Mootha et al., 2003) of the top expressing genes in these clusters showed an increase in membrane dynamics, lipid metabolism, and immune responses, notably interactions with platelets, coagulation, and complement (Figure S3F). This suggests that the stabilization of an epithelial phenotype in KPC;ST primary tumors may lead to unique metastatic adaptations, including modulation of the immune response. These data demonstrate that pancreatic cancer cells with a stabilized epithelial phenotype can efficiently metastasize, with an increased growth advantage in the liver.

Stabilized epithelial PDAC cells migrate by collective cell migration

We further examined the cancer cell-intrinsic pathways associated with stabilized epithelial PDAC cells. YFP+ cancer cells were freshly sorted from the primary pancreatic tumors and the most common metastatic sites, liver and lung, of the KPC and KPC;ST mice for global expression profiling. GSEA of the KPC;ST primary tumors compared to the KPC control tumors showed 3,569 and 647 pathways up- and downregulated, respectively. The most highly upregulated pathways were associated with membrane dynamics, immune regulation, metabolism, and, notably, ameboid and epithelial migration (Figure 4A). Using Ingenuity Pathway Analysis (Krämer et al., 2014), we confirmed that the loss of Snail and Twist are critical nodes for regulating these pathways (Figures S4A and S4B). Similar pathway expression patterns were observed when examining metastatic cancer cells (3,616 and 600 pathways, respectively) (Figure 4B). Also, a common metastatic signature (3,152 and 1,064 pathways, respectively) between KPC;ST and KPC PDAC cells was noted (Figure S4C). String analysis of the core enriched genes in the ameboid migration gene set show RhoA as
a central signaling hub (Szklarczyk et al., 2019). Using qPCR, we further confirmed the elevated expression of several genes in the epithelial/ameboid gene set (Figure S4D). These data suggest that the stabilization of epithelial PDAC in the primary tumor results in adaptive mechanisms that promote liver metastatic outgrowth.

To functionally validate the potential role of collective epithelial migration as a mechanism for metastasis in KPC;ST mice, we performed a scratch assay of YFP-sorted primary cancer cell lines. We used several primary cell lines isolated from KPC and KPC;ST tumors as well as a primary cell line isolated from an adenoviral-Cre-induced KPC; E-cadherin-KO (knockout) tumor to represent a stabilized mesenchymal phenotype. The proliferation rates, as determined by the frequency of passaging of the cell lines, were comparable between the groups. In general, both the KPC and KPC;ST lines maintained cell-cell contact in a uniform sheet with the absence of mesenchymal single cells (Figures 4C and 4D). However, unlike the ubiquitous epithelial morphology of the KPC;ST lines, the KPC lines were heterogeneous with some lines displaying leading cells with mesenchymal morphology (Figure S4E). The KPC; E-cadherin-KO cell line demonstrated pronounced single-cell mesenchymal morphology. Over the course of the scratch closure, both the KPC and the KPC;ST lines exhibited a steady decrease in single cells (particle counts) reflective of collective sheet migration (Figure 4E) (Friedl et al., 2012; Friedl and Wolf, 2010; Friedl et al., 2014). Alternatively, the KPC; E-cadherin-KO cell line maintained as single cells with clearly distinguishable boundaries (Figures 4C and 4D). The epithelial migration of the KPC and KPC;ST cells resulted in a more complete scratch closure than the single-cell mesenchymal migration of the KPC; E-cadherin-KO cell line (Figure 4F). Of note, the heterogeneity of the KPC lines also followed this pattern, with the line displaying the most collective phenotype (978U) clustering with the KPC;ST lines (Figures S4F and S4G). This suggests that the epithelial mode of migration is not due solely to the absence of Snail and Twist. These data confirm that epithelial PDAC cells migrate in vitro by collective sheet migration and that the loss of Snail and Twist does not prevent migration.

**Epithelial PDAC cells associate with more T cells**

To investigate the possibility of a distinct immune tumor microenvironment, we evaluated the number of infiltrating immune cells determined by scRNA-seq within KPC or KPC;ST tumors. Both the early KPC and KPC;ST tumors had comparable levels of immune cells (Figure 5A). However, the late-stage KPC tumors displayed a dramatic reduction in the percentage of immune cells, while the KPC;ST tumors maintained a higher level of immune populations, notably T cells (Figure 5A). These data were confirmed using multiplex immunohistochemistry (Figure 5B), which showed a significant inverse correlation between CD8+ T cells and EMT+ cells (CK8+ αSMA+ and YFP+ and any mesenchymal marker+; see Figure 3) (Figures 5C and 5D). PDAC patient tumors’ scRNA-seq showed no significant correlations in the percentages of infiltrating immune cells across the E/M spectrum tumors (Figures 5E and S5A). As tumor tissues are heterogeneous, we next investigated whether differences in immune infiltration could be observed in areas that contain more epithelial or mesenchymal cancer cells as opposed to bulk tumor counts. We have previously published the importance of the spatial localization of T cells in predicting clinical outcomes (Carstens et al., 2017). We re-examined our published tissue array for EMT (CK8+ αSMA+). As
suggested by examining patient survival using the Thiery EMT signature, the presence of EMT alone did not predict survival (Figure 5F and 5G). However, the presence of EMT inversely correlated with T cells, which by themselves were a predictor of improved survival (Figures 5G, 5H, and S5B) (Carstens et al., 2017). We further determined that EMT+ cancer cells had significantly fewer T cells within cell-cell contact distances (Figures 5I and S5C). These data demonstrate that epithelial and mesenchymal PDAC cells elicit differential effects on the infiltrating immune microenvironment.

Epithelial stablization via Zeb1 ablation also enhances liver metastasis

We show that the deletion of Snail and Twist is sufficient to suppress Zeb1 (Figure 3F). To specifically address the role of Zeb1-induced EMT in PDAC progression, we deleted Zeb1 in cancer cells of a second version of the KPC model with homozygous loss rather than a dominant negative mutation of p53 (KrasG12D; p53f/f; p48-Cre; YFP_LSL, hereafter referred to as KPPC). In comparison to the p53mut KPC model, the KPPC model displays a rapidly progressing primary tumor but with low distant metastases (Timpson et al., 2011). There was no delay in tumor onset in KPPC mice compared to KPPC with heterozygous Zeb1 deletion (KPPC;Zf+o) or homozygous Zeb1 deletion (KPPC;ZcKO) as measured by the emergence of a first palpable pancreatic nodule (Figure 6A). The tumor onset kinetics agree with previously published KPC; Zeb1cKO data (Krebs et al., 2017). However, there was a significant delay in tumor progression as reflected in improved survival and a slower increase in tumor burden and histology over time, but with equivalent tumor burdens and histologies at endpoint (Figures 6B–6G). The reduction of cancer cell-specific Zeb1 protein expression was confirmed through immunolabeling of the primary tumor in both the KPPC;Zf+ and the KPPC; Zeb1cKO (Figures S6A and S6B) as well as quantified with the co-expression of mesenchymal markers and the lineage tracer YFP (Figure 6H). Similar to the dual loss of Snail and Twist, which reduce the cancer cell expression of Zeb1, we also observed the stabilization of an epithelial phenotype (Figure 6H). Next, we assessed the rate of metastasis. The tumors in the KPPC model are locally invasive to the surrounding gut, resulting in bowel obstruction and/or a blocked bile duct and jaundice, and were associated with a lower incidence of liver or lung metastases, when compared to the KPC mice (Table S2). We observed a significant increase in the rate of macroscopic liver metastasis in the KPPC; Zeb1cKO compared to KPPC (Tables S2 and S3). Microscopically, we observed a significant increase in the number and size of liver metastasis in both the KPPC; Zeb1f+ and KPPC; Zeb1cKO (Figures 6G, 6I, and S6C; Tables S2 and S3). The increased liver metastasis trended with an increase proliferation, although not striking, again suggesting other mechanisms beyond proliferation as seen in the KPC;ST model (Figure S6D). These data confirm that the stabilization of the PDAC epithelial phenotype enhances liver metastasis even in a low metastatic KPPC PDAC model.

DISCUSSION

Recently, the importance of the EMT program in various biological processes, including cancer, and challenges in studying EMT were summarized with guidance for more rigorous in vivo genetic modeling studies (Yang et al., 2020). In this regard, advances in next-generation and single-cell sequencing technology provide an opportunity for a more nuanced
analysis and appreciation of the full spectrum of the EMT continuum and addressing its role in metastasis.

Here, we provide single-cell quantitation of the EMT spectrum in human and KPC murine PDAC primary and metastatic tumors. Human and KPC PDAC displayed over 50 different phenotypic clusters of cancer cells across the EMT continuum based on the Thiery EMT signature using 315 human and 294 mouse gene transcripts. Our study clearly identifies that the EMT program of cancer cells is highly heterogeneous, with many cells existing in pEMT and mesenchymal states, along with different epithelial states. We discover that the EMT program is highly plastic and exists in an interchangeable continuum influenced substantially by transcription factors such as Snail, Twist, and Zeb1, which both overlap and represent unique subpopulations. We also identify that the genetic loss of Snail and Twist (associated with suppression of Zeb1) substantially affects the plastic continuum of the EMT program and stabilizes the epithelial phenotype with a shift toward epithelial cancer cells, away from the pEMT and mesenchymal states. These results suggest that a compensation for the loss of Snail and Twist was not encountered in this model system and that the EMT program, as represented by hundreds of genes, was affected at a broad level, favoring an epithelial phenotype at the expense of pEMT and mesenchymal states. We further demonstrate that a large-scale inhibition of the EMT program does not impair metastasis, but rather is associated with increased liver metastasis, likely due to the fitness of the cancer cells with an epithelial phenotype. Importantly, the number of metastatic lesions does not change, suggesting that the extravasation and intravasation steps of cancer cell dissemination from the primary tumor are not impared due to epithelial stabilization via inhibition of the EMT program, which is supported by previous studies (Zheng et al., 2015). These data suggest that inhibition of partial or hybrid EMT states does not impact metastasis; it appears that the system can compensate for the absence of mesenchymal features and accomplish metastasis with similar efficiency.

A previous study showed that the loss of Zeb1 in the cancer cells of KPC mice leads to decreased metastasis associated with a trend toward decreased primary tumor growth, and an insignificant difference in disease-free survival of the mice (Krebs et al., 2017). These results suggested that decreased metastasis could be a direct reflection of the rate of primary tumor progression (Krebs et al., 2017). To further validate that epithelial stability can lead to increased liver metastasis, we used a genetic mouse model of pancreatic cancer that is recalcitrant to metastasis (Timpson et al., 2011). This genetic mouse model substitutes the mutant p53 allele in the KPC mice with an allele leading to the loss of p53 (p53F/F), and the resulting KPPC mice develop primary PDAC, but rarely metastasize. Deletion of Zeb1 in KPPC mice led to a decreased rate of primary tumor growth but an insignificant difference in disease-free survival of these mice when compared to the KPPC mice. The loss of Zeb1 enabled the emergence of significant liver metastasis associated with the epithelial stabilization of the EMT program. These results further support the idea (Reichert et al., 2018) that suppression of an EMT program leads to epithelial stabilization, resulting in the fitness of cancer cells to form efficient liver metastatic colonies.

The concept of epithelial versus mesenchymal cancer cells and its implications on clinical prognosis is well established in the PDAC field. Basal-like or quasi-mesenchymal gene
expression subtypes have been associated with poor patient outcomes (Porter et al., 2019). In addition, model studies have provided a causative link between mesenchymal traits and therapeutic resistance (Shah et al., 2007; Arumugam et al., 2009; Wellner et al., 2009; Krebs et al., 2017; Gaianigo et al., 2017). However, patients with epithelial/classical phenotypes represent approximately two-thirds of the clinical population and still have poor prognosis with metastatic disease (Porter et al., 2019). Furthermore, 76%–80% of PDAC metastasizes to the liver, regardless of expression subtypes (Houg and Bijlsma, 2018; Hayashi et al., 2020). The data presented here are in agreement with these clinical observations and suggest a link between the presentation of classical/epithelial subtypes and the preponderance of liver metastasis.

In summary, our data suggest that the EMT program on a single-cell level within the primary tumor is heterogeneous, and cancer cells can exist within multiple states within the full spectrum of the EMT continuum. Suppression of this plastic EMT spectrum leads to epithelial stabilization and contributes to the fitness of the cancer cells for liver metastatic colonization, without any impact on the dissemination of the cancer cells. Our data suggest that the dissemination of epithelial cancer cells is facilitated by collective migration and collaboration with the immune system. Successful colonization and growth are considered to be the most challenging steps of metastatic disease, and the present study highlights the importance of epithelial stabilization in this process and offers insights into potential therapeutic interventions.

STAR METHODS

RESOURCE AVAILABILITY

Lead contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Raghu Kalluri, MD, PhD (rkalluri@mdanderson.org).

Materials availability—Zeb1-floxed allele reported in this study was generated from the Knockout Mouse Project (KOMP) plasmid (tm1a(KOMP)Wtsi).

There are restrictions to the availability of the mouse lines and generated cell lines from these mouse models due to institutional property rights and require an MTA for transfer. Please contact the Lead author for more information.

Data and code availability—The accession number for the murine single-cell RNA-sequencing data reported in this paper is GEO: GSE165534. The accession number for the murine YFP sorted cancer cell microarray is GEO: GSE164612.

Original data have been deposited into Mendeley Data: https://dx.doi.org/10.17632/sxc5zk4m9w.1.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mouse studies—Characterization of disease progression and genotyping for the Pdx1-Cre; LSL-KrasG12D; P53R172H+/−; R26-LSL-EYFP (herein referred to as KPC), Pdx1-Cre;
KPC; Snai1\textsuperscript{cKO} and KPC; Twist\textsuperscript{cKO} were bred to generate the KPC; Snail\textsuperscript{cKO}; Twist\textsuperscript{cKO} dual conditional knockout line (herein referred to as KPC;ST). The KPC;ST mice were maintained on a mixed background, bred according to Mendelian ratios, and developed normally (data not shown). KPC;Snail\textsuperscript{F+}; Twist\textsuperscript{F+} mice showed no differences in all assays investigated from the KPC;Snail\textsuperscript{++}; Twist\textsuperscript{++} mice and were included in the KPC control group.

All mice were housed under standard housing conditions at MD Anderson Cancer Center (MDACC) animal facilities, and all animal procedures were reviewed and approved by the MDACC Institutional Animal Care and Use Committee. Investigators were not blinded for group allocation, but were blinded for the assessment of the phenotypic outcome assessed by histological analyses. Males and females were utilized equally. No randomization method was used and all mice of the desired genotype were enrolled in the studies. Age, disease progression and gender are reported for each mouse in Table S2.

**Primary tissue sorting and cell line generation**—Primary cancer cell lines were generated similar to a previous publication (Zheng et al., 2015). In brief, tumor tissue was minced with scissors in a 1.5 mL tube, and incubated at 37°C in 4 mg/mL Collagenase IV (GIBCO, 17104019) and 4 mg/mL Dispase II (GIBCO, 17105041) in serum-free RPMI in a 6-well dish for 1 hour. Cells were resuspended in 10 mL of complete media (RPMI with 20% FBS and 1% PSA) and strained through a 70 um strainer and pelleted to removed digestion media. For cell line generation, the cells were then resuspended in 2 mL of complete media and plated on 6-well Collagen IV coated plates (Corning BioCoat, 62405–636), after reaching approximately 80% confluency the cells were then passaged onto standard T25 tissue culturing flasks. For single cell sorting, the pelleted cells were washed twice in PBS, resuspended in FACS buffer (PBS with 2% FBS) and sorted for YFP positivity by flow cytometry.

The stabilized mesenchymal KPC; E-cadherin-KO control line was isolated from the primary tumor generated by the adenoviral-Cre mediated transformation of the pancreas in a KP;E-cadherin\textsuperscript{FF} mouse. In brief, the previously published Cdh1\textsuperscript{L/+} allele (Derksen et al., 2006) was crossed into the KPC background. Adenoviral vector injections were roughly based upon a previous publication (Gierut et al., 2014). Cre negative KP;E-cadherin\textsuperscript{FF} mice were injected in the tail of the pancreas with 5x10\textsuperscript{6} pfu in 25 μL. (5 μL of 1x10\textsuperscript{9} pfu virus + 23 μL of MEM + 2 M CaCl\textsubscript{2}) of adenoviral Cre (Vector BioLabs 1045) per mouse. Tumor tissue was collected at necropsy and a cell line was isolated as described, above.
All cell lines used in the study were validated for genotype, recombination purity (to exclude stromal cell contamination) and EMT expression status through qPCR. Details for genotype and sex of each cell line are provided in the Key resources table.

**METHOD DETAILS**

**Single-cell RNA-sequencing**—Patient scRNA-seq data were mined from Peng et al. (2019). Whole pancreatic tumor tissue from six KPC (3 early and 3 late) and six (1 early and 5 late) KPC:ST mice and liver tissue from 3 KPC late and 4 KPC:ST late mice were processed to obtain a single-cell suspension for each tissue. Tumor tissue was minced and digested with Collagenase IV (4 mg/mL) and Dispase (4 mg/mL) in serum-free DMEM, shaking at 37°C for 30 minutes. The suspension was then passed through a 70 um and 40 um filter and washed twice with DMEM containing 20% FBS. Cells were stained for 30 minutes on ice in 1:100 of eFluor 780 fixable viability dye (eBioscience 65-0865-14). Cells were sorted on a BD FACSAria for all live or live/endogenous YFP+ as appropriate. scRNA-seq on these samples was conducted at the MDACC Advanced Technology Genomics Core. Single-cell Gel Bead-In-Emulsions (GEMs) generation and barcoding, post GEM-RT cleanup and cDNA amplification, library construction and Illumina-ready sequencing library generation were prepared following the manufacturer’s guidelines. High Sensitivity dsDNA Qubit kit was used to estimate the cDNA and library concentration. HS DNA Bioanalyzer was used for the quantification of cDNA. DNA 1000 Bioanalyzer was used for the quantification of libraries. The “c-loupe” files were generated by using Cell Ranger software pipelines following the manufacturer’s guidelines. Cells from the unfractionated tumor were encapsulated using 10X Genomics’ Chromium controller and Single-Cell 3′ Reagent Kits v2. Following capture and lysis, cDNA was synthesized and amplified to construct Illumina sequencing libraries. The libraries from approximately 1,000–4,000 cells per sample were sequenced with Illumina Nextseq 500 or HiSeq 3000. The run format was 26 cycles for read 1, 8 cycles index 1, and 124 cycles for read 2. scRNA-seq data were processed by the Advanced Technology Genomics Core at MDACC.

Library Seurat (version 3.6.1)(Butler et al., 2018; Stuart et al., 2019), dplyr and cowplot were loaded into R (version 3.6.1) to explore QC metrics, filter cells, normalize data, cluster cells, and identify cluster biomarkers. To filter out low-quality cells, a threshold with a minimum of 200 and a maximum of 4000–7000 genes per cell was used. Cells with more than 10% of the mitochondrial genome were also removed from further analysis. To remove the influence of technical characteristics from downstream analyses, “sctransform” package was used for normalization. “RunUMAP” function was used for clustering the cells. “FindAllMarkers” function was used to identify the specific markers for each cell cluster as well as downstream analysis on gene expression. “DoHeatmap” function was used to show the top genes in each cluster. “VlnPlot” function was used to show expression probability distributions across cell clusters of the genes selected to assign the cell type identity and the genes of interest.

The MAGIC algorithm utilizes nearest neighbor graphing and a diffusion operator to restore or “smooth” missing transcripts from the single-cell expression data based on the expression of similar cells (van Dijk et al., 2018). MAGIC smoothing of the cancer cell cluster was run.
using the library Matrix and library Rmagic (van Dijk et al., 2018) based off of a 315 EMT gene signature (Tan et al., 2014). The exact resolution/cluster numbers were selected based on the consistency of the expression heatmaps within each cluster. To validate our results obtained by MAGIC processing using the 315 Thiery gene signature (305 genes for human and 294 genes for the mouse were discoverable in our dataset), we utilized a second MAGIC based method utilizing principle component analysis (PCA): magic(data, genes = “pca_only”). 3D scatterplots of the results were plotted and color-coded for CDH1, VIM, FN1, and ZEB1.

The post-MAGIC Thiery EMT Signature expression values were used to define cancer cell EMT stage. An “E/M score” was calculated by taking the Log2 of the percentage of positive epithelial genes (from the scaled.data) divided by the percentage of positive mesenchymal genes: \( \log_2(\text{EPI/MES}) \). Log2 values to set general boundaries for clinical correlate analyses and comparison of mouse signatures to the human were determined as follows: Epithelial \( \geq 1.5 \), Partial-EMT \( < 1.5 \) and \( > -1.5 \), and Mesenchymal \( \leq -1.5 \).

Cancer cells, defined by Seurat, were further analyzed by Monocle 3 (Haghverdi et al., 2018) to construct single-cell trajectories and order the cells in pseudotime. The figures related to pseudotime were generated by “learn_graph,” “plot_cells,” and “order_cells.” The area enriched for both KPC early and KPC:ST early cells was defined as the root nodes for pseudotime. Modules of co-regulated genes were found by “find_gene_modules.”

**Histology and histopathology**—Histology and histopathological scoring were previously described (Zheng et al., 2015). Formalin-fixed tissues were embedded in paraffin and sectioned at 5 μm thickness. Sections were stained with hematoxylin and eosin (H&E). Histopathological measurements were made by scoring H&E stained tumors for relative percentages of each histopathological phenotype: normal (non-neoplastic), PanIN, well-differentiated PDAC, moderately-differentiated PDAC, poorly-differentiated PDAC, or necrosis. The weighted pathology scores were calculated by taking the relative percentages for each phenotype and assigning a weighted score (Normal > 5% = 1; PanIN > 30% = 2; PDAC > 30% = 4; Poorly-differentiation PDAC > 5% = 5; and Necrosis > 5% = 6). These scores were then summed to produce the weighted pathology scores presented in Figure 4. When tumor histology was missing or of poor quality, the mice were excluded from all histological analyses and this was determined blinded from genotype information.

Microscopic metastases were observed in H&E stained tissue sections of at least four and three depths, 150 μm apart for the liver and lung, respectively. These data have been presented as a contingency table (Table S3) and represented as the number of positive tissues (one or more lesions in a tissue) out of the number of tissues scored. The “Any” metastasis score is the number of mice positive for a secondary lesion found anywhere throughout the body out of the total number of mice scored. Metastatic lesion numbers and areas were processed by circling every metastatic lesion in each tissue section using scanned slides and the Aperio ImageScope software. Data represented as indicated numbers per tissue depth.

**Immunohistochemical staining and analyses**—Antigen retrieval conditions, antibody concentration and company details are outlined in Table S4. In brief, tissues were fixed in 10% formalin overnight, dehydrated, embedded in paraffin and 5-μm-thick sections...
were deparaffinized for subsequent staining. Immunofluorescence was performed by antigen retrieval in heated citric acid buffer (pH 6.0) or Tris-EDTA with 0.1% Tween 20 (pH 9.0) by microwave (EZ Retriever microwave, BioGenex) or pressure cooker as indicated. Unless otherwise indicated, each combination of staining was put through sequential rounds of staining, each including a protein block, followed by a primary antibody and a corresponding secondary antibody directly conjugated to the indicated fluorophore or HRP-conjugated polymer followed by covalent binding of an Opal fluorophore using tyramide signal amplification (Perkin Elmer Opal7 Kit). Sections were either stained with DAPI (1:20,000) and mounted with VectaShield mounting media or mounted with a DAPI containing mounting media (Sigma). For immunohistochemical staining, sections underwent antigen retrieval, incubation in hydrogen peroxide (3%), protein block, primary and secondary antibody incubations with biotinylated conjugated antibody, amplified with streptavidin-HRP conjugation (ABC kit, VectorLabs) and developed by DAB according to the manufacturer’s recommendations.

Representative images were taken using a Leica DM1000 light microscope, a Vectra3 Multispectral Imaging System (Akoya Biosciences), or a Zeiss AxioScan.Z1 at 20x magnification and analyzed using cell segmentation and scoring on inForm (v2.4.6; Akoya) or QuPath (Bankhead et al., 2017) (for Ki-67). For EMT quantification, YFP+ cancer cells were identified by setting a threshold with the inForm scoring algorithms and then hand counted as being EMT+ (co-expression of any mesenchymal marker and YFP) and presented as the average percentage double positive out of YFP positive cells per image per mouse. Immune population data on a patient TMA was mined from previously published data (Carstens et al., 2017). Spectrally unmixed images were scored for CK8+aSMA+ cells using inForm phenotyping. Mouse immune population data were unmixed using inForm and analyzed for tissue segmentation, cell segmentation and phenotyping (aka phenomapping) using Vis analysis software from Visiopharm. XY coordinates for each cell extracted from multiplexed images were used to compute the extent of T cell infiltration around cancer cells using the L-function as described (Carstens et al., 2017).

**CTC assays**—Blood (1,000 μL) was collected at necropsy and incubated with 13 mL of ACK lysis buffer (A1049201, GIBCO) for five minutes at room temperature to lyse red blood cells. Cell pellets were washed twice with PBS and resuspended in 300 μL of PBS containing 2% FBS and analyzed for the number of YFP+ cells by flow cytometry (BD FACS Aria IIIu Cell Sorter).

**Quantitative reverse-transcriptase PCR analyses (qPCR)**—RNA was extracted from cultured primary PDAC cells using TRIzol (15596026, Life Technologies). cDNA synthesis was performed using the Applied Biosystems cDNA synthesis kit according to the manufacturer’s directions. Quantitative PCR was performed to analyze the gene expression profiles of the listed genes using SYBR Green PCR Master Mix in a 7300 Sequence Detector System (Applied Biosystems), and measurements were standardized to the expression of the 18s housekeeping gene. The expression data are presented as fold change ($2^{\Delta \Delta Ct}$) with the averaged control group normalized to a fold value of 1. dCt were used to
measure statistical significance in observed changes. Genes and primer sequences are in Key resources table.

**Microarray analyses**—YFP⁺ cells were single-cell sorted at the South Campus Flow Cytometry core (BD FACSria IIIu Cell Sorter). RNA was isolated using either Trizol or PicoPure RNA isolation kit (Applied Biosystems). Total RNA was quantified using an RNA6000 Pico or Nano assays (Aligent), whole transcriptome analysis was performed using the WT Plus Assay (Applied Biosystems) and run using the Mouse Clarion D array (Applied Biosystems) by the MDACC Sequencing and Genome Core Program. Gene expression analysis was performed using the TAC4.0 version 2 software. Gene set enrichment analysis (GSEA) was performed for KEGG, biocarta, c5bp, and hallmarks gene sets using GSEA software version 4.0.0 and MSigDB 7.0 (Broad Institute). Gene expression microarray data have been deposited in GEO: GSE164612. Ingenuity pathway analysis (QIAGEN) was run on the expression values of the primary cancer cells using 2.1 fold and p < 0.05 as cutoffs.

**Migration assays**—Primary PDAC cell lines that had been confirmed by qPCR for expression of EMT genes, were plated densely in 3 wells of a 24-well plate for each line. When the cells reached confluency, the media was removed and a vertical scratch was made in the center of each well with a P100 pipette tip. Wells were washed twice in PBS and the media was replaced (RPMI, 20% FBS, 1% PSA). Each well was imaged using a 4x phase objective every 30 minutes for 40 hours using the Olympus BX51 Live Imaging Microscope and Scope software. The experiment was repeated three times for lines 28B and 407B and four times for all others. Exported tif images were processed using FIJI (ImageJ, NIH) and the PHANTAST cell segmentation plugin (Jaccard et al., 2014). The sigma value was kept constant at 1.2 and the epsilon value was adjusted between 0.05 and 0.15 to compensate for lighting changes of each well, but kept constant over the time course for each well. Images were then inverted and particle count and area (confluency) were analyzed.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical analyses were performed using unpaired two-tailed t tests, one-way and two-way ANOVA with Tukey’s multiple comparisons test using GraphPad Prism version 8 and 9, as stipulated in the figure legends. χ² analyses, using SPSS statistical software, were performed comparing control to conditional knock-out groups across multiple histological parameters. Results are outlined in Table S3. χ² analyses were performed on scRNA-seq patients comparing each of the defined E/M groups across all provided clinical and histopathological parameters. Results are outlined in Table S1. Analysis of TCGA Pancreatic Cancer (PAAD) was performed using the UCSC Xena browser (Goldman et al., 2020). Kaplan-Meier plots were drawn for survival analysis and the log rank Mantel-Cox test was used to evaluate statistical differences, using GraphPad Prism 8. Data met the assumptions of each statistical test, where variance was not equal (determined by an F-test) Welch’s correction for unequal variances was applied. Error bars represent s.e.m. Multiple visual fields were averaged to produce a single value for each animal which was then averaged again to represent the mean bar for the group in each graph. p < 0.05 was considered statistically significant.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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### Highlights

- Single-cell RNA-seq reveals a continuum of cancer epithelial-mesenchymal transition
- Genetic deletion of both *Snai1* and *Twist1* or *Zeb1* stabilizes epithelial phenotypes
- Epithelial cancer cells have increased liver metastasis and collectively migrate
- Mesenchymal-like cancer cells recruit fewer T cells
Figure 1. Single-cell RNA sequencing (RNA-seq) of human PDAC reveals 51 different cancer cell phenotypes across the EMT continuum

(A) Seurat uniform manifold approximation and projection (UMAP) clustering of whole-tissue single-cell populations. The red dashed line highlights the cancer cells isolated for further analysis.

(B) UMAP clustering of patient cancer cells post-MAGIC using the Thiery EMT signature.

(C) Same as (B), recolored to reflect the E/M score.

(D) E/M score for each cluster, with the percentage of cancer cells in each E/M phenotype and corresponding expression heatmap of the Thiery EMT signature. Epithelial genes are above the red line. The blue line indicates the chosen cutoff between epithelial, partial, and mesenchymal groups.

(E) Percentage of cancer cells in each E/M phenotype.

See also Figure S1.
Figure 2. Single-cell RNA seq of murine PDAC identifies 56 cancer cell phenotypes across the EMT spectrum

(A) Seurat UMAP clustering of whole-tissue single-cell populations. The red dashed line highlights the cancer cells isolated for further analysis.

(B) UMAP clustering of murine pancreatic cancer cells post-MAGIC using the Thiery EMT signature. (C) Same as (B), recolored to reflect the E/M score.

(D) E/M score for each cluster and corresponding expression heatmap of the Thiery EMT signature. Epithelial genes are above the red line. The blue line indicates the chosen cutoff between epithelial, partial, and mesenchymal groups, with the aggregated percentage of cancer cells indicated.

(E) Pie charts for the percentage of cancer cells in each E/M phenotype for aggregated human and mouse and each experimental mouse cohort. See also Figure S2.
Figure 3. Stabilized epithelial phenotype of PDAC cells enhances liver metastasis
(A) Overall survival curve of KPC control (n = 17) and KPC;ST (n = 16) mice.
(B) Disease-free survival curve of KPC control mice (n = 18) and KPC;ST (n = 30) mice as determined by first palpable pancreatic nodule.
(C) Percentage of tumor burden (tumor-bearing pancreas weight/body weight) in grams of endpoint KPC control (n = 14) and KPC;ST (n = 15) mice.
(D) Representative histological micrographs (2003) stained with H&E of primary pancreatic tumors and liver and lung metastatic lesions; 100 μm scale bar. The yellow line outlines the border of metastatic lesions. T marks tumor area and L marks liver or lung, respectively.
(E) Relative percentage area of indicated epithelial phenotypes in primary pancreatic tumors of endpoint KPC control (n = 12) and KPC;ST (n = 14) mice. Two-way ANOVA.
(F) Quantification of the percentage double positive of the indicated mesenchymal marker: Snail/Slug, Slug, Twist, Zeb1, Vimentin, αSMA, and FSP1 with YFP lineage tracing out the...
total number of YFP+ cells per 200 × image (5–10 images) per mouse: KPC (n = 9 or 10) and KPC;ST (n = 8) mice. Means ± SEMs.

(G) Quantification of the number of metastatic lesions per tissue depth in the liver (KPC control, n = 25 depths, 8 mice, and KPC;ST, n = 32 depths, 12 mice), lung (KPC control, n = 11 depths, 5 mice, and KPC;ST, n = 13 depths, 8 mice), and other tissues (KPC control, n = 13 depths, 11 mice, and KPC;ST, n = 10 depths, 9 mice).

(H) Quantification of the metastatic area per tissue depth for the liver (KPC control, n = 25 depths, 8 mice, and KPC;ST, n = 32 depths, 12 mice), lung (KPC control, n = 11 depths, 5 mice, and KPC;ST, n = 13 depths, 6 mice), and other tissues (KPC control, n = 13 depths, 11 mice, and KPC;ST mice, n = 10 depths, 9 mice).

(I) Pie charts for the percentage of cancer cells in each E/M phenotype across indicated experimental cohort. Unless otherwise specified, data are presented as means ± SDs and significance determined by an unpaired two-tailed t test. ns, not significant, *p < 0.05, **p < 0.01.

See also Figure S3.
Figure 4. Stabilized epithelial PDAC cells migrate by collective cell migration

(A and B) Select enriched gene sets comparing primary tumors (A) and metastatic tumors (B). All pathways displayed are significant with a nominal p value, false discovery rate (FDR) q value, and family-wise error rate (FWER) p value < 0.05.

(C–F) Representative micrographs (40×) (C), matching inverted PHANTAST cell segmentation masks (D) and quantifications (E and F) of 3–4 replicate experiments with increasing passage numbers of cancer cell lines isolated from the primary tumors of KPC control, KPC;ST, and KP-EFF-AdCre (n = 3, 3, and 1 line isolated from individual mice, respectively) plated to confluency. The red boxes indicate digital zoom of areas magnified in far-right panel. Quantifications of the particle counts (E) and percentage of cellular confluency (F) with linear regressions (****p < 0.001) of the grouped values post-scratch for the indicated time points.

See also Figure S4.
Figure 5. Epithelial PDAC cells associate with more T cells
(A) Immune cell percentages from scRNA-seq for each GEMM cohort.
(B) Cell population percentages from multiplexed immunohistochemistry for indicated genotype.
(C) Correlation between CD8+ T cells and CK8+ αSMA+ cancer cells in KPC and KPC;ST tumors.
(D) Correlation between CD8+ T cells and YFP+CK8+AnyMesenchymal marker+ (Vimentin, αSMA, Zeb1, Snail, Twist, Slug, and FSP1) cancer cells in KPC and KPC;ST tumors.
(E) Immune cell percentages from scRNA-seq for each patient grouped by E/M classifications.
(F) Survival of patients high or low for EMT determined by multiplexed immunohistochemistry (split on the median).
(G) Cell population percentages from multiplexed immunohistochemistry for patients high or low for EMT (CK8+ αSMA+).

(H) Correlation between CD8+ T cells and CK8+ αSMA+ cancer cells.

(I) L-function area under the curve values (reflecting the number of cells) for CD8+ T cell within 20 μm of CK8+ or CK8+ αSMA+ cancer cells; t test. Unless otherwise specified, data are presented as means ± SDs and significance determined by one-way ANOVA. ns, not significant, *p < 0.05, **p < 0.01, ****p < 0.0001. See also Figure S5.
Figure 6. Epithelial stabilization via Zeb1 ablation also enhances liver metastasis
(A) Disease-free survival curve of KPPC (n = 3), KPPC;Z^F/+ (n = 12), and KPPC; Zeb1^K0(n = 5) mice as determined by first palpable pancreatic nodule.
(B) Overall survival of KPPC (n = 4), KPPC;Z^F/+ (n = 7), and KPPC; Z^K0(n = 5) mice. Log-rank tests.
(C) Percentage of tumor burden (tumor-bearing pancreas weight/body weight) in grams plotted by age of necropsy (n = 4, 11, 5 mice, respectively); lines represent linear regression.
(D) Weighted sum of pathology scores for each mouse plotted by age of necropsy (n = 4, 7, and 5 mice); linear regression line.
(E) Percentage tumor burden of endpoint mice (n = 4, 6, 5 mice, respectively).
(F) Relative percentages of epithelial tumor histology of endpoint mice (n = 4, 7, 5 mice, respectively), two-way ANOVA.
(G) Representative histological micrographs (200×) stained with H&E of primary pancreatic tumors and liver and lung tissues. The yellow line outlines the border of metastatic lesions. T marks tumor area, and L marks liver or lung, respectively.
(H) Quantifications of the percentage of double positive out of YFP^+ cells per field of view (200× magnification) of primary pancreatic tumors (n = 4, 5, and 5 mice, respectively) and indicated mesenchymal markers: Snail/Slug, Slug, Twist, Zeb1, Vimentin, αSMA, or FSP1. SEM, ANOVA.
(I) Quantification of the metastatic area per tissue depth for liver (KPPC, n = 16 depths, 4 mice, KPPC;Z^{F/+}, n = 48 depths, 12 mice, and KPPC; Z^{cKO}, n = 20 depths, 5 mice), lung (KPPC, n = 12 depths, 4 mice, KPPC;Z^{F/+}, n = 36 depths, 12 mice, and KPPC; Z^{cKO}, n = 15 depths, 5 mice), and other tissues (KPPC, n = 50 depths, 4 mice, KPPC;Z^{F/+}, n = 72 depths, 12 mice, and KPPC; Z^{cKO}, n = 30 depths, 5 mice). Unless otherwise specified, data are presented as means ± SDs, scale bar: 100 μm, and significance determined by a one-way ANOVA. ns, not significant, *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001. 
See also Figure S6.
### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Rabbit monoclonal anti-Slug | Cell Signaling Technology | Cat# 9585; RRID:AB_2239535 |
| Rabbit polyclonal anti-FSP1 | Agilent (originally DAKO) | Cat#A5114; RRID:AB_2335679 |
| Rabbit polyclonal anti-Zeb1 | Novus | Cat# NBP1-05987; RRID:AB_2273178 |
| Rat monoclonal anti-cytokeratin 8 | DSHB | Cat# TROMA-I; RRID:AB_531826 |
| Chicken polyclonal anti-GFP | Aves | Cat# GFP-1020; RRID:AB_10000240 |
| Mouse monoclonal anti-Twist1 | Novus | Cat# NP2-37364; RRID:AB_2801339 |
| Rabbit monoclonal anti-Vimentin | Cell Signaling Technology | Cat# 5741; RRID:AB_10695459 |
| Rabbit polyclonal anti-Snail + Slug | Abcam | Cat# ab180714; RRID:AB_2728773 |
| Mouse monoclonal anti-aSMA | Agilent (originally DAKO) | Cat# M0851; RRID:AB_2223500 |
| Mouse monoclonal anti-Ecadherin | Cell Signaling Technology | Cat# 3195; RRID:AB_2291471 |
| Rabbit monoclonal anti-Ki-67 | Thermo Scientific | Cat# RM-9106-S; RRID:AB_149707 |
| Rabbit on Rodent HRP Polymer | Biocare Medical | Cat# RMR622 |
| Mouse-on-Mouse HRP-Polymer | Biocare Medical | Cat# MM620 |
| Polink-1 HRP Rat-NM for DAB Bulk Kit | GBI labs | Cat# D35–110 |
| Polink-2 Plus HRP Chicken IgY for DAB Kit | GBI labs | Cat# D84–60 |
| Goat polyclonal anti-Rabbit Alexa Fluor 488 | Invitrogen | Cat# A-11008; RRID:AB_143165 |
| Goat polyclonal anti-Chicken Alexa Fluor 594 | Invitrogen | Cat# A-11042; RRID:AB_2534099 |
| Goat polyclonal anti-Rat Alexa Fluor 647 | Invitrogen | Cat# A-21247; RRID:AB_141778 |
| Goat polyclonal anti-Rabbit, Biotinylated | Vector Laboratories | Cat# BA-1000; RRID:AB_2313606 |
| **Bacterial and virus strains** | | |
| Ad-CMV-iCre | Vector Biolabs | Cat#1045 |
| **Biological samples** | | |
| mouse model pancreatic tumor and mixed organ FFPE blocks | This paper | N/A |
| **Critical commercial assays** | | |
| Opal 7 color IHC Kit | Akoya Biosciences | NEL811001KT |
| TSA Coumarin System | Akoya Biosciences | NEL703001KT |
| VECTASTAIN Elite® ABC System | Vector Labs | PK-6100 |
| Power SYBR Green PCR Master Mix | Thermo Fisher | 4367659 |
| Fluoroshield mounting media | Sigma Aldrich | F618 |
| Vectashield Mounting Media | Vector Laboratories | H1000 |
| **Deposited data** | | |
| Raw and analyzed microarray data | This paper | GEO:GSE164612 |
| Raw and analyzed scRNA-seq data | This paper | GEO:GSE165534 |
| GDC TCGA Pancreatic Cancer (PAAD) | TCGA | [https://xenabrowser.net/datapages/](https://xenabrowser.net/datapages/) |
### REAGENT or RESOURCE | SOURCE | IDENTIFIER
--- | --- | ---
Patient scRNA-seq data | Peng et al., 2019 | N/A
Source Data | N/A | Mendeley Data: https://dx.doi.org/10.17632/xxc5zk4m9w.1

**Experimental models: Cell lines**

| Control Line 978U; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Rosa26-LSL-EYFP \textsuperscript{F/F}; Female | This paper | N/A |
| Control Line H444; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Rosa26-LSL-EYFP \textsuperscript{F/F}; Female | This paper | N/A |
| Control Line 0739; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Rosa26-LSL-EYFP \textsuperscript{F/F}; Male | This paper | N/A |
| KPC-ST Line 309P; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Snai1 \textsuperscript{F/F}, Twist1 \textsuperscript{F/F}; Rosa26-LSL-EYFP \textsuperscript{F/F}; aSMA-RFP; Female | This paper | N/A |
| KPC-ST Line 404P; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Snai1 \textsuperscript{F/F}, Twist1 \textsuperscript{F/F}; Rosa26-LSL-EYFP \textsuperscript{F/F}; aSMA-RFP; Female | This paper | N/A |
| KPC-ST Line 407P; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Snai1 \textsuperscript{F/F}, Twist1 \textsuperscript{F/F}; Rosa26-LSL-EYFP \textsuperscript{F/F}; aSMA-RFP; Female | This paper | N/A |
| KPC Ecad KO Line 28B; Adenoviral-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+ (may contain Rosa26-LSL-EYFP or aSMA-RFP) | This paper; Dr. Rhim | N/A |
| KPPC; P48-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{F/F}, Zeb1 \textsuperscript{F/F}; (may contain Rosa26-LSL-EYFP) | This paper; Dr. Rhim | N/A |
| KPPC;Z \textsuperscript{F+}; P48-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{F/F}, Zeb1 \textsuperscript{F/F}; (may contain Rosa26-LSL-EYFP) | This paper; Dr. Rhim | N/A |
| KP;E \textsuperscript{FF}-AdCre \textsuperscript{+}; Adenoviral-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Cdh1 \textsuperscript{F/F}, Rosa26-LSL-EYFP \textsuperscript{F/F}; Female | This paper | N/A |

**Experimental models: Organisms/strains**

| KPC; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+ (may contain Rosa26-LSL-EYFP or aSMA-RFP) | This paper | N/A |
| KPC;ST; Pdx1-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Snai1 \textsuperscript{F/F}, Twist1 \textsuperscript{F/F}; Rosa26-LSL-EYFP \textsuperscript{F/F}; aSMA-RFP; Female | This paper | N/A |
| KPPC; P48-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{F/F}; (may contain Rosa26-LSL-EYFP) | This paper; Dr. Rhim | N/A |
| KPPC;Z \textsuperscript{F+}; P48-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{F/F}, Zeb1 \textsuperscript{F/F}; (may contain Rosa26-LSL-EYFP) | This paper; Dr. Rhim | N/A |
| KP;E \textsuperscript{FF};AdCre \textsuperscript{+}; Adenoviral-Cre; Kra5 \textsuperscript{G12D}, p53 \textsuperscript{R172H}/+; Cdh1 \textsuperscript{F/F}; (may contain Rosa26-LSL-EYFP or aSMA-RFP) | This paper | N/A |

**Oligonucleotides**

| mouse Snai1 Forward; CACACGCTGCCTTGTTCT | Lovisa et al., 2015 | N/A |
| mouse Snai1 Reverse; GGTCAAGCAAAGACACAGTTT | Lovisa et al., 2015 | N/A |
| mouse Twist1 Forward; CTGCCCTCGGACAAGCTGAG | Lovisa et al., 2015 | N/A |
| mouse Twist Reverse; CTAGTGGGACGCGGACATGG | Lovisa et al., 2015 | N/A |
| mouse Zeb1 Forward; GCCTCAGGATAAATGACGGC | Harvard Primer Bank | N/A |
| mouse Zeb1 Reverse; GCCTCAGGATAAATGACGGC | Harvard Primer Bank | N/A |
| mouse Cdh1 Forward; GTCAACACCTCAACAGCCTGAG | Lovisa et al., 2015 | N/A |
| mouse Cdh1 Reverse; GTTTTGCTCAAGCCCTTCAC | Lovisa et al., 2015 | N/A |
| mouse Vim Forward; CTGTAGAAGGATGAAATCTCCT | Lovisa et al., 2015 | N/A |
| mouse Vim Reverse; GTCAGCCGTTGGAACGTCC | Lovisa et al., 2015 | N/A |
| mouse Col1a1 Forward; CTCTCTTTAGGGGCCACT | Lovisa et al., 2015 | N/A |
| mouse Col1a1 Reverse; CCACGTCTCACCCATTGGGG | Lovisa et al., 2015 | N/A |
| mouse 18s Forward; GTAACACCTCAACAGCCTGAG | Lovisa et al., 2015 | N/A |
| mouse 18s Reverse; CCATCCAATCGGTAGTAGGGC | Lovisa et al., 2015 | N/A |
| mouse Rhoa Forward; AGCTTGTGTAAGACATGCTTG | Harvard Primer Bank | N/A |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| mouse Rhoa Reverse; GTGTCCCATAAAGCCAACACTCTAC | Harvard Primer Bank | N/A |
| mouse Rock1 Forward; GACTGGGGACAGTTTGGAGAC | Harvard Primer Bank | N/A |
| mouse Rock1 Reverse; GGGCATCCCAATCCATCCACG | Harvard Primer Bank | N/A |
| mouse Rac1 Forward; GAGACGGAGCTGTGGTAAAA | Harvard Primer Bank | N/A |
| mouse Rac1 Reverse; ATAGGCCCAGATTCTACTGTT | Harvard Primer Bank | N/A |
| mouse Sdc1 Forward; GAGACGGAGCTGTGGTAAAA | Harvard Primer Bank | N/A |
| mouse Sdc1 Reverse; ACGGGCCCTCAACAGTCAG | Harvard Primer Bank | N/A |
| mouse Ctsb Forward; TTCCTGATCTCTTCTCTTGCC | Harvard Primer Bank | N/A |
| mouse Ctsb Reverse; ACAGTGCCACACAGCTCTCTTC | Harvard Primer Bank | N/A |
| mouse Lamp1 Forward; CAGCACTCTTTTGAGGTGAAAAAC | Harvard Primer Bank | N/A |
| mouse Lamp1 Reverse; ACGATCTGAGAACCATTCGCA | Harvard Primer Bank | N/A |
| mouse Max Forward; ACCATAATGCACAGACTGGAACGAAA | Harvard Primer Bank | N/A |
| mouse Max Reverse; GTCCCCGAAACTGTGGAAGGC | Harvard Primer Bank | N/A |
| mouse Jun Forward; CCTTCTACAGCAGATGCCCTC | Harvard Primer Bank | N/A |
| mouse Jun Reverse; GGTTCAAGGTCTGCTCTGTTT | Harvard Primer Bank | N/A |
| mouse Sec24b Forward; GGTTCCAGCAGACAGTCCCAATG | Harvard Primer Bank | N/A |
| mouse Sec24b Reverse; GGAGTCCCCGAAATTGTTGTT | Harvard Primer Bank | N/A |
| mouse Ndufb11 Forward; CCTCCAGGCGCTTAATCGC | Harvard Primer Bank | N/A |
| mouse Ndufb11 Reverse; GGTTCTCCGCCTAGCCTTTTC | Harvard Primer Bank | N/A |
| mouse Apoe Forward; CCGACAGATGCTCTAGCCG | Harvard Primer Bank | N/A |
| mouse Apoe Reverse; CCGAGTATCCAGAAAGC | Harvard Primer Bank | N/A |
| Software and algorithms | | |
| Prism 8 | GraphPad | N/A |
| InForm v2.4.6 | Akoya Biosciences | N/A |
| Vis 6.2019 | Visiopharm | N/A |
| UCSC Xena browser | Goldman et al., 2020 | N/A |
| Seurat (version 3.6.1) | Butler et al., 2018; Stuart et al., 2019 | N/A |
| library Matrix and library Rmagic | van Dijk et al., 2018 | N/A |
| Monocle 3 | Haghverdi et al., 2018 | N/A |
| L-Function | Carstens et al., 2017 | N/A |
| FIJI with Phantast plugin | NIH; Jaccard et al., 2014 | N/A |
| QuPath | Bankhead et al., 2017 | N/A |
| SPSS Statistics | IBM | N/A |