Enzymatic targeting of the stroma ablates physical barriers to treatment of pancreatic ductal adenocarcinoma

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SUMMARY

Pancreatic ductal adenocarcinomas (PDA) are characterized by a robust fibroinflammatory response. We show here that this desmoplastic reaction generates inordinately high interstitial fluid pressures (IFP), exceeding those previously measured or theorized for solid tumors, and induces vascular collapse, while presenting substantial barriers to perfusion, diffusion and convection of small molecule therapeutics. We identify hyaluronan, or hyaluronic acid (HA), as the primary matrix determinant of these barriers and show that systemic administration of an enzymatic agent can ablate stromal HA from autochthonous murine PDA, normalize IFP and re-expand the microvasculature. In combination with the standard chemotherapeutic, gemcitabine, the treatment permanently remodels the tumor microenvironment and consistently achieves objective tumor responses resulting in a near doubling of overall survival.

INTRODUCTION

Pancreatic ductal adenocarcinoma (PDA) is the fourth leading cause of cancer-related deaths in the United States (Jemal et al., 2010). The inherent biology of the disease makes it not only uniformly but also rapidly lethal: overall 5-year survival for PDA is less than 5% with a median survival of 4-6 months (Hidalgo, 2010). Gemcitabine, a deoxycytosine analog, represents the current standard-of-care for advanced disease and improves quality of life in a minority of patients while prolonging survival by several weeks (Burris et al., 1997). A recent advance involving targeted inhibition of epidermal growth factor receptor (EGFR) with erlotinib extends median survival by an additional fourteen days (Moore et al., 2007). Moreover, despite notable advances in surgical technique and post-operative care, and the
use of adjuvant chemical and radiotherapies, virtually all resected early stage patients also eventually succumb to recurrent and/or metastatic disease (Allison et al., 1998; Farnell et al., 2005; Oettle et al., 2007). Understanding the unusual resistance of pancreas cancer and finding ways to treat it at all stages of disease are clear and imperative needs.

Most preclinical drug evaluations to date have relied on in vitro assays and in vivo cell transplantation models. However, tumor cells arising in situ in the native organ are categorically distinct from the same cells grown in culture or engrafted into an immunocompromised mouse (Bissell and Radisky, 2001). Indeed, transplanted pancreas carcinoma cells respond readily to conventional chemotherapeutic agents (Hertel et al., 1990), in marked contrast to autochthonous tumors in mice (Olive et al., 2009) and humans (Tempero et al., 2003). This problem cuts both ways: in failing to fully appreciate the unique challenges imposed by the complex cancer “organ,” opportunities to target and exploit the most clinically relevant mechanisms may also be missed. Thus, there are non-cell autonomous factors, including unique cell-cell and cell-matrix interactions and evolving intratumoral physical dynamics, that contribute to the in vivo resistance of cancers.

Prior work in a number of experimental systems has suggested that altered intratumoral fluid dynamics can limit the effectiveness of systemic therapies (reviewed in Bouzin and Feron, 2007; Heldin et al., 2004; Jain, 1987; Tredan et al., 2007). Mammalian organs and tissues typically possess interstitial fluid pressures (IFP) at or below the intravascular pressures (IVP) in the terminal arterioles and capillaries that supply them. As first proposed by Starling in 1896, an IFP less than IVP permits ready perfusion and favors diffusive and convective forces necessary for solute and fluid flow into the interstitium (Starling, 1896, and see Supplemental Information). The solutes are consumed by cells while the venous vasculature and functional lymphatic channels return excess fluid to the systemic circulation (reviewed in Michel, 1984). While these processes governing the delivery and diffusion of cancer drugs have been studied extensively in xenograft models, very few experiments have been performed in autochthonous carcinomas and no rigorous analyses before and after perturbation of the tumor microenvironment have been described.

PDAs are characterized by an unusually intense desmoplastic reaction comprised of fibroblasts, immune cells and endothelial cells embedded within a dense and complex extracellular matrix (ECM). The ECM component hyaluronic acid (HA), or hyaluronan, is inordinately abundant, inviting speculation on its role in disease biology and resistance. HA is a large linear glycosaminoglycan (GAG), composed of repeating N-acetyl glucosamine and glucuronic acid units, and figures prominently in the architecture, integrity and malleability of tissues, particularly in dynamic processes such as embryogenesis and oncogenesis (Toole, 2004). Its viscoelastic properties underlie its unique role in tissue homeostasis and its appeal in cosmetic and clinical applications (Balazs and Denlinger, 1989). HA also communicates directly with cells in health and disease through interactions with several distinct types of surface receptors. However, the abundance of HA varies widely across tumors and it has been shown to both promote and suppress tumor growth (Toole, 2004). Even its potential contribution to IFP is controversial. Local injection of hyaluronidase into osteosarcoma xenografts has been shown to decrease IFP (Brekken et al., 2000; Eikenes et al., 2005), although enforced overexpression of HA in thyroid and colon carcinoma xenografts did not increase it (Jacobson et al., 2003). Thus, although HA has been implicated in a variety of cell autonomous and non-cell autonomous processes, its precise role(s) in therapeutic resistance of cancers has remained elusive and is likely context-specific (reviewed in Toole and Slomiany, 2008; Tredan et al., 2007).

As an alternative to transplanted cells and tumors, we have developed genetically engineered mouse (GEM) models of PDA that faithfully recapitulate the clinical syndrome,
histopathology, and molecular progression of human pancreas cancer from inception to invasion. Kras<sup>LSL-G12D/+;Cre (KC)</sup> (Hingorani et al., 2003) and Kras<sup>LSL-G12D/+;Trp53<sup>LSL-R172H/+;Cre (KPC)</sup> (Hingorani et al., 2005) mice conditionally express endogenous, physiologic levels of the respective oncogene and tumor suppressor gene mutations in pancreatic progenitor cells resulting in the stochastic development and spontaneous progression of preinvasive ductal lesions to invasive and metastatic carcinomas. In collaborative studies using the KPC model, we have recently suggested that the unusually hypovascular microenvironment of PDA compromises the delivery of even small molecule therapeutics such as gemcitabine. These studies showed further that targeting stromal fibroblasts by disrupting paracrine Hedgehog signaling between the epithelial and mesenchymal tumor compartments transiently increased the apparent microvascular density through neoangiogenesis and improved drug delivery, resulting in a meaningful if modest survival benefit (Olive et al., 2009). Interestingly, disease resistance rapidly re-emerged along with a return of the profound hypovascularity. Moreover, the mechanisms by which both the paucity and the collapse of vessels arose in the first place and the processes that sustained them remained unclear. In the present study, we pursue a mechanistic understanding of the biophysical barriers to perfusion, diffusion and convection imposed by the desmoplastic reaction in autochthonous PDA and investigate strategies to surmount them.

RESULTS

Cellular and molecular evolution of desmoplaxis in PDA

As part of a systematic effort to characterize the evolving stromal dynamics and potential therapeutic vulnerabilities during disease progression, we performed specific histochemical and immunohistochemical assays to identify components of the ECM and infiltrating cells in preinvasive, invasive and metastatic PDA in mice and humans (Figure 1 and Figures S1A-S1D). A robust and definable stromal reaction develops in association with early precursor lesions and includes a dense collagen content organized in a fibrillary structure in both primary tumors and metastases, as revealed by intravital second harmonic generation (SHG) imaging (Figures S1E-S1J). The tight association of this fibrillar collagen with epithelial carcinoma cells became evident in Kras<sup>LSL-G12D/+;Trp53<sup>LSL-R172H/+;R26<sup>LSL-GFP/+;Cre (KPGC) mice. Interestingly, the relative concentrations of collagen and glycosaminoglycans (GAG) appear to alter during disease progression, shifting toward higher GAG content at more advanced and metastatic stages of disease (Figure 1A-1J). One particular GAG, HA, was especially abundant (Figures 1K-1O), which was confirmed by treating tissue sections with hyaluronidase (Figures S1K-S1N). We note that significant HA deposition begins with early precursor lesions (Figures 1G and 1L). In addition, activated stromal pancreatic stellate cells (PSC), or myofibroblasts, also infiltrate early and are prevalent throughout disease progression (Figures 1P-1T). Conversely, a decrease in vessel number is seen with progression (Figures S1O-S1Q), as described previously (Olive et al., 2009), and is accompanied by a decrease in cross-sectional (luminal) area (Figure S1R).

This same overall molecular and cellular stromal composition is observed in human PDA (Figure 1E, 1J, 1O and 1T). We note, in particular, the expression of moderate to high levels of HA in all human PDAs tested (n = 30, Figures 1J and 1O and not shown), suggesting that this substrate may be an essential component of disease architecture, biology and resistance.

Dramatically elevated IFP in PDA

To directly measure IFP in normal pancreata and in autochthonous PDAs in live animals, we used a system coupling a miniature piezoelectric pressure transducer to a pressure control unit and data acquisition module (Figure 2A). Anesthetized animals underwent laparotomy
followed by needle-guided placement of the pressure transducer probe enabling real-time in vivo IFP measurements (Figures 2B and 2C). As expected, the normal pancreas possesses an IFP ranging between 8 – 13 mmHg (mean = 10.4 mmHg, n = 8) (Figure 2D). In contrast, we observed dramatically elevated IFPs in autochthonous PDA that far exceeded the normal range for the pancreas (Figure 2D). Tumor IFPs ranged from 75 – 130 mmHg (mean = 99 mmHg, n = 7, in KPC PDA; mean = 87 mmHg, n = 4, in KC tumors), which vastly exceeded typical arteriolar and capillary pressures of 40 – 80 mmHg and 15 – 40 mmHg, respectively (Sherwood, 1997) and rivaled mean arterial pressure (Janssen et al., 2000; Mattson, 1998). Pressures of this magnitude suggest not only major impediments to delivery and convection of small molecules, but also imply a profound reorganization and remodeling of the tumor architecture and the forces operant within it (see also Supplemental Information).

**Enzymatic targeting of stromal HA restores tumor IFP and functional perfusion**

We began our studies on the ability of HA to elevate IFP using reconstituted three-dimensional matrices. Purified primary murine PDA cells were embedded in matrices with differing HA concentrations, the tumor plugs implanted in immunodeficient mice and IFPs recorded (Figures S2A-S2C). HA concentrations of 3 mg/ml more than doubled the baseline IFP of carcinoma cells engrafted in collagen alone, suggesting that the presence of HA in the tumor can contribute directly to elevated IFP.

The abundance of HA in PDA and its demonstrated capacity to significantly elevate IFP prompted us to investigate whether a systemically delivered agent could ablate HA in the stroma of an autochthonous PDA and decrease pressures. We first tested the ability of intravenously administered PEGPH20 to deplete HA from normal tissues in wild-type (WT) mice and assessed for any untoward effects (Figures S2D-S2F). Several organs in the body including heart, lung, bowel and liver contain modest levels of detectable HA (Figure S2E), while joints possess very high levels (Figure S2F). HA was efficiently ablated from all of these sites except cartilage and the surrounding joint space. Cartilage represents an essentially avascular compartment completely excluding delivery of PEGPH20. Interestingly, depletion of HA from these sites had no discernible effects on organ function or animal health; animals remained active without apparent changes in energy level or mobility and also maintained their body weight (Figure S2D). We next demonstrated that intravenous administration of PEGPH20 to KC and KPC mice could deplete HA in autochthonous PDAs (Figure S2G). Moreover, after a single intravenous dose of PEGPH20, IFP was significantly decreased within 2 hours and approached the range for normal pancreata 24 hours post-treatment (Figure 2D). The restoration of near normal tumor IFP after degradation of stromal HA suggested that treatment with PEGPH20 should also significantly and rapidly improve blood flow if indeed mechanical compression of the vasculature was the primary mechanism limiting perfusion. To assess the effects on the tumor microvasculature of IFP normalization, we performed a systematic survey of vessel number and lumen diameter in WT pancreas as well as untreated and treated murine PDA (Figures 2E-2G). We found a significant shift in the distribution of CD31+ vessel diameters in murine and human PDA compared with the normal pancreas and, in particular, an almost complete absence of large diameter (> 10 μm) vessels (Figures 2E-2G and S2H and S2I). The normal pancreas contains a range of small to moderate sized vessels, more than 80% of which possess a readily discernible and patent lumen; conversely, the majority (~75%) of vessels in PDA lacked an apparent lumen, consistent with vascular collapse from elevated extrinsic fluid pressures. Following treatment with PEGPH20, vessel diameter was significantly increased and lumens clearly discernible in ~71% of CD31+ vessels (Figure 2G and S2H).
We next performed functional analyses of the vasculature with intravital multiphoton laser scanning microscopy (MPLSM). Normal pancreata revealed a rich, widely patent vasculature (Figure 2H) with excellent delivery of the small molecule therapeutic doxorubicin throughout the parenchyma (Figure 2K). In contrast, vessels in PDA were difficult to detect and appeared collapsed (Figure 2I) and penetration of doxorubicin into the tumor bed was extremely limited (Figure 2L). Even when rare perfusion was observed (Figure 2L), the distribution of drug into the tumor was low, suggesting limited diffusion and convection into the interstitium. A single intravenous dose of PEGPH20 resulted in the immediate appearance of significant numbers of patent vessels (Figures 2J and 2M) reflecting their increased diameters (Figure 2G). These changes were accompanied by a >6.5-fold increase in doxorubicin fluorescence intensity over baseline (p=0.013, n = 4 each for treated and untreated KPC cohorts; and p=0.04, n = 3 per group for KCPDAs). We note that overall vessel number did not significantly change (not shown), which together with the rapidity of the effect and lack of significant endothelial cell proliferation (see below) suggest that neoangiogenesis did not contribute significantly. Rather, it appears that the inordinately high IFP produces vascular collapse in PDA and reversal of these elevated pressures restores functional perfusion and unimpeded delivery of drug into the carcinoma.

**Combined enzymatic and cytotoxic chemotherapy alters tumor biology and disease response and increases overall survival in PDA**

We next sought to assess the effects of combination chemotherapy on the gross morphological, physicomechanical, cellular and molecular properties of autochthonous PDA. In previous studies of gemcitabine in KPC mice (Olive et al., 2009), we used a dose, route and schedule modeled on the original preclinical investigations of gemcitabine in subcutaneously engrafted tumors (Hertel et al., 1990; Schultz et al., 1993). For the present studies, we treated KPC cohorts according to the route and schedule used in the clinic. Specifically, we performed a randomized, placebo-controlled study of the combination regimen involving intravenous drug administration in cycles consisting of the 3-weeks ‘on’ and 1-week ‘off’ schedule used in patients (Figure 3A). Animals were first assessed with serial high-resolution ultrasound measurements until they achieved a requisite enrollment tumor diameter of 2-5 mm. Studies were performed on separate cohorts of animals to measure early and intermediate effects, and also to establish overall survival.

As expected, IFP normalized in Gem+PEGPH20 treated tumors, confirming that the addition of gemcitabine did not impede the ability of PEGPH20 to influence this physicomechanical property (Figure 3B). Interestingly, no change in baseline tumor IFP occurred in Gem+Placebo treated animals reflecting the inability of gemcitabine to effectively penetrate tumors and induce responses. The gross findings at necropsy underscored these differences as recovered tumors from Gem+Placebo treated animals (Figures 3C and S3A) revealed the hard, white, fibrous and overtly hypovascular composition typical of the untreated disease (Figure S3B). In contrast, tumors from animals receiving Gem+PEGPH20 were notably soft, highly vascularized and even hemorrhagic when transected (Figures 3D and S3C-S3F). A blood-rich microenvironment in combination treated tumors was readily apparent even by routine histology (Figure S3G-S3L). Quantitative analysis confirmed significant differences in vessel diameter between Gem+Placebo and Gem+PEGPH20 treated PDA (Figures 3E and 3F) without significant differences in vessel number (not shown). Importantly, these findings were observed in all mice immediately after initiating therapy (Figures S3G-S3L), during active treatment (Figures S3M and S3N) and persisted even after treatment cessation (Figures S3O-S3T). Even nearly 3 months after completing combination therapy, functional vessels were retained suggesting an irreversible change in tumor physiology (Figures S3S and S3T).
Objective measures of clinical response were frequently apparent after only one cycle of combination therapy. Notably, of the subset of Gem+Placebo treated animals that underwent quantitative volumetric imaging both before and after treatment, none experienced an objective tumor response affirming the collective clinical (Burris et al., 1997; Casper et al., 1994) and preclinical experience with this agent in autochthonous PDA (Olive et al., 2009) (Figures 4A and 4B). In contrast, 83% of Gem+PEGPH20 treated tumors decreased in size after one cycle of treatment (Figures 4A and 4B) with all tumors responding after 3 cycles (see example in Figure S4A). Consistent with its known mechanisms of inhibiting DNA and RNA synthesis (Hertel et al., 1990; Ueno et al., 2007), combination therapy both decreased proliferation and increased apoptosis of primary tumor cells compared with Gem+Placebo, contributing to the observed differences in objective responses (Figures 4C and 4D and S4B-S4E). Thus, when able to penetrate the tumor bed, gemcitabine can indeed be an effective agent against this disease.

The numbers of total stromal PSC (Figure S5A) and activated PSC (Figures 5A-5C and S5B) were also significantly decreased and the tumor stroma was extensively remodeled (Figures 5D-5G). Although unaffected by PEGPH20 alone, collagen content and distribution were both significantly decreased by the combined regimen, likely as a secondary consequence of depleting collagen-secreting activated PSC. Finally, no discernible effects on endothelial cell proliferation (Figure S5C) or apoptosis (Figure S5D) were observed, consistent with the expected restricted activity of a conventional cytotoxic agent to actively dividing cells. Thus, the depleted HA and diminished IFP seen with the combined enzymatic and cytotoxic regimen restores and preserves a functional vasculature while inducing tumor epithelial and stromal cells to die.

Similar restrictions and responses were observed in metastases. Metastatic lesions to the liver (Figures 1D, 1I, 1N, 1S) and lung (Figure 6A) possessed a robust and complex desmoplasia, with notable HA content and a paucity of vessels (Figures S6A-S6G). Perhaps surprisingly, given their typically smaller size compared with primary tumors, the metastatic deposits were also poorly perfused by small molecules (Figures S6H-S6K). As with primary tumors, PEGPH20 efficiently ablated HA from metastases (Figure 6B) and restored functional perfusion (Figure S6K). When given together with gemcitabine, the combination regimen decreased proliferation (Figure 6C) and increased apoptosis (Figure 6D) in metastases to both distant sites.

The aforementioned encouraging results notwithstanding, the most clinically relevant measure of efficacy is survival. We therefore performed a prospective, randomized, placebo-controlled trial in KPC mice with overall survival as the primary endpoint. Secondary endpoints included changes in intratumoral IFP and objective response rates, as well as metastatic disease burden.

Median overall survival increased from 55.5 days for Gem+Placebo to 91.5 days for Gem+PEGPH20, an 83% increase (Figure 6E). These results likely do not reflect the maximal achievable benefit from combination therapy as the treatment course was limited to 3 months because of sclerosing of the vein after repeated intravenous injections. In fact, only 29% of Gem+PEGPH20 treated animals died during the course of therapy and approximately 30% of animals lived an additional 4-11 weeks after completing 3 full months of treatment. In contrast, more than 80% of animals receiving Gem+Placebo died during treatment; of the remaining 4 animals, 3 died in the very next week after cessation of therapy (Figure 6E and Table S1).

The metastatic tumor burden was also significantly diminished with combination therapy (Figure 6F and Tables 1 and S1). The frequency of malignant ascites as well as metastases to
the liver, lungs, diaphragm and mesenteric lymph nodes were all decreased in the Gem +PEGPH20 cohort. Thus, the combination regimen effectively treated both the primary and metastatic disease burdens.

DISCUSSION

The defining features of PDA are a penchant for metastatic spread and a notorious resistance to chemical and radiotherapies. These features in turn establish the major hurdles to meaningful treatment of the disease. A perhaps prosaic, albeit unexpected and critically important, barrier to systemic therapy is a profoundly diminished functional vasculature in PDA. We show here that the primary mechanism limiting perfusion in PDA is significant collapse of resident vessels by inordinately high fluid pressures. Small molecules that are delivered to the tumor bed are subsequently impeded from penetrating the interstitium by constraints on diffusion and convection, completing a trifecta of stromal barriers to chemotherapy. Enzymatic degradation of hyaluronan results in a rapid reduction of IFP accompanied by the appearance of widely patent functioning vessels. Removing these barriers permits high concentrations of chemotherapy to reach the tumor, resulting in improved survival and revealing an unappreciated sensitivity of the disease to conventional cytotoxic agents.

The theory governing the movement of fluid and solute across a semi-permeable membrane dividing two compartments (e.g. vascular and extravascular) is readily derived from fundamental thermodynamic principles (Kedem and Katchalsky, 1958; Kedem and Katchalsky, 1961; Staverman, 1952) (reviewed in Ogston and Michel, 1978) and incorporates hydrostatic and osmotic pressure gradients as the primary determinants of fluid flow, and concentration gradients as the driving force for solute flux. The resultant equations, in turn, explain the inability to achieve effective chemotherapy concentrations in PDA given the extremely high IFPs observed (see Supplemental Information for a detailed treatment of these equations). They do not, however, adequately address how such dramatically elevated IFPs arise in the first place. The prevailing hypotheses envision the intravascular and interstitial compartments to be in contiguity or communication, suggesting that changes in IFP chronicle and directly reflect intratumoral vascular pressures largely as a consequence of “leaky” or damaged vessels (Boucher and Jain, 1992; Netti et al., 1999). However, the essentially complete collapse of the microvasculature in invasive PDA implies that this relationship no longer holds and forces other than intravascular pressure must be maintaining the dramatically elevated IFP. Indeed, we have observed a sustained elevation of tumor IFP even after the cessation of cardiac function (unpublished observations), underscoring the disconnect between intravascular pressure and IFP in this setting. Some models also postulate an elevated interstitial permeability in tumors (Jain, 1990) which appears not to be the case in PDA and, in any event, would favor not hinder delivery of (macro)molecules. Thus, despite an extensive history of studies in experimental systems and limited work in human cancers in situ (Boucher et al., 1991; Curti et al., 1993; Less et al., 1992), the magnitude of interstitial hypertension in solid tumors has clearly been underestimated, particularly with respect to PDA, and the mechanisms driving its genesis have remained elusive.

Plausible mechanisms to further elevate interstitial pressure include a sufficiently prolific ECM production to increase tumor density more rapidly than volume, and/or robust cell contractility that actively compacts the tumor (Reed et al., 1992). In this regard, the ability of a systemically delivered enzyme to dissipate the high IFPs in PDA not only holds great therapeutic promise but also provides insight into the underlying physicomechanical processes operant within this complex cancer environment (Figure 7). Intact HA functions as a hydrated gel generating an immobile fluid phase that secondarily also diminishes
compressibility (tethered collagen fibers under tension provide the primary framework for rigid structure). We propose that PEGPH20 first liberates water bound to HA through cleavage of the extended polymer into substituent units. Release of trapped water rapidly decreases IFP to a range of 20 – 30 mmHg, enabling collapsed arterioles and capillaries to open. Without additional treatment, HA is replenished, in part by α-SMA(+)

myofibroblasts, and intratumoral physical dynamics are restored. Together with a concomitant cytotoxic agent, however, the resultant death of activated PSC leads to the additional loss of collagen I content through decreased synthesis, while also unraveling its architecture, which itself has been implicated in promoting carcinoma invasion and metastasis (Provenzano et al., 2006; Wang et al., 2002), impeding drug delivery (Eikenes et al., 2004; Netti et al., 2000; Ramanujan et al., 2002), and conferring gemcitabine resistance (Dangi-Garimella et al., 2011). Thus, enzymatic degradation of HA removes the central barrier, allowing drugs to breach the previously impenetrable sanctuary of PDA. A subsequent feed-forward mechanism results in depletion of stromal fibroblasts and carcinoma cells, dissolution of the collagen network and irrevocable remodeling of the tumor microenvironment. Uniform objective responses ensue, at least until cells can evolve secondary mechanisms of resistance. This gap in time represents an unprecedented opportunity for therapeutic advantage. The persistence of patent vessels enables switching sequentially from one chemotherapeutic regimen to another in hopes of prolonging this window of vulnerability. In this context, re-visiting the extensive list of previously presumed “ineffective” agents against PDA (Nieto et al., 2008) represents an obvious starting point even as additional targets within the tumor stroma are identified and explored (Izeradjene and Hingorani, 2007).

In the context of our current findings, two very recent reports of combination regimens to treat advanced PDA are particularly notable for their shared pharmacokinetic features and their improved success over the current standard. One regimen combined gemcitabine with nab-paclitaxel, an albumin-coated formulation of taxol with an extended half-life of >10 hours in the circulation (Von Hoff et al., 2011). A second regimen, FOLFIRINOX, added bolus dosing of two conventional cytotoxic agents onto a base of prolonged (46 hours) continuous infusion of fluorouracil with each treatment cycle (Conroy et al., 2011). In each case, the sustained exposure to cytotoxic therapy may have helped surmount the relative barriers to perfusion imposed by the ECM of PDA.

Finally, the prospect that augmenting blood flow to a lethal carcinoma may increase its ability to seed metastatic deposits gives pause; however, the dramatically decreased metastatic tumor burden in animals treated with combination therapy is reassuring and no doubt contributes to the observed benefit in overall survival. Indeed, that Gemcitabine +PEGPH20 improves survival in animals with advanced and metastatic disease underscores its potential utility for the majority of patients who present with PDA. These findings also reflect the similar physicochemical principles operant in metastases as in the primary tumor; in fact, our molecular characterization of the ECM suggests that the metastatic microenvironment is further enriched in GAGs (including HA) relative to collagen as compared with primary tumors. Understanding the precise mechanism of achieving a decreased metastatic tumor burden has important implications for specific clinical uses of PEGPH20. The decreased metastatic burden may have been achieved by regression of established metastases (which we have observed) and/or decreased seeding of metastases either by killing primary tumor cells or altering their ability to spread. HA has been directly implicated in increasing the metastatic potential of cancer cells (Zhang et al., 1995). However, HA depletion may also inhibit the ability of already seeded cells to establish clinically relevant metastases. In other words, HA may represent a necessary component of the microenvironmental niche in order for a circulating tumor cell or nest of cells to blossom fully into a life-limiting lesion. Thus, there may be additional potential uses for and benefits
from a stromal targeting therapy including, for example, as part of an adjuvant regimen. Given that more than two thirds of patients with resected PDA eventually succumb to metastatic disease (Hidalgo, 2010), incorporating an enzymatic agent that degrades HA in the adjuvant setting may extend survival by preventing, or at least delaying, the establishment of a critical metastatic disease burden.

EXPERIMENTAL PROCEDURES

Mouse Strains

All animal studies were approved by the Institutional Animal Care and Use Committee of the Fred Hutchinson Cancer Research Center. Kras<sup>LSL-G12D/+;Trp53<sup>LSL-R172H/+;Cre</sup></sup> (KPC) mice have been previously described in detail (Hingorani et al., 2005). KPC mice conditionally express endogenous physiologic levels of activated Kras and point mutant Trp53 targeted to progenitor cells of the developing pancreas. The animals spontaneously develop the full spectrum of precursor ductal lesions (pancreatic intraepithelial neoplasias, PanINs) which progress to invasive and metastatic PDA, faithfully mimicking the clinical syndrome, histopathology, and molecular progression of the human disease from inception to invasion and metastasis. A subset of experiments were also performed on Kras<sup>LSL-G12D/+;Cre</sup> (KC) mice which similarly develop invasive and metastatic disease along the PanIN-to-PDA progression scheme, albeit with a slower time course than KPC mice (Hingorani et al., 2003). ROSA26<sup>LSL-EGFP</sup> mice (obtained from Jackson Laboratory) contain enhanced green fluorescent protein (EGFP) cDNA sequence flanked by LoxP sites knocked-in to the ROSA26 locus.

Measuring interstitial fluid pressure (IFP)

IFP measurements were performed using a Millar Mikro-Tip® pressure catheter transducer (SPR-1000, 0.33 mm diameter) with a dynamic pressure range from −50 to 300 mmHg, possessing a shielded side mounted sensor. The catheter was connected to PCU-2000 Pressure Control Unit and an ADInstruments PowerLab® data acquisition system (Millar Instruments, Inc.) (Ozerdem and Hargens, 2005; Thompson et al., 2010). Data were recorded and analyzed using LabChart software (Millar Instruments, Inc.). The system was calibrated to 0, 25 and 100 mmHg prior to each measurement following the manufacturer’s recommendations; calibration was re-confirmed after each study. To place the microtip pressure catheter, a 25 gauge needle (~2X the diameter of the probe) was first introduced into the tissue or tumor of interest and the probe inserted into the space after needle withdrawal.

Assessment of vascular function and molecular diffusion

To assess vascular function, 425 μl of biotin-conjugated <i>Lycopersicon esculentum</i> lectin (B1175-1mg, Vector Laboratories) was mixed with 75 μl of Streptavidin-AlexaFluor-488 (S32354, Molecular Probes) for whole tissue MPLSM imaging or with Streptavidin-AlexaFluor-647 (S21375) for analyses of tissue sections following co-administration of doxorubicin (Sigma). Mice were anesthetized by inhalation of isofluorane gas. Five minutes prior to euthanasia, 100 μl of the conjugated lectin was administered into the left ventricle. For co-administration experiments, 20 mg/kg doxorubicin solution was injected immediately after fluorescently-conjugated lectin. While under terminal anesthesia, mice were perfused with 4% paraformaldehyde. For histologic analyses, harvested tissues were additionally fixed overnight in 4% paraformaldehyde.

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Treatment schedule and survival study

Enrollment of KPC mice was based on primary tumor burden and informed by age. Specifically, animals between 12-14 weeks of age and with a palpable pancreas tumor were enrolled. Ultrasound imaging was performed in the initial 11 mice enrolled to confirm the physical exam findings of estimated tumor burden. KPC mice were genotyped and then randomized by one investigator (A.E.C.) to receive Gemcitabine+Placebo or Gemcitabine+PEGPH20, which were administered by another investigator (P.P.P.). Objective response was assessed by high-resolution ultrasound performed by a third investigator (C.C.) who was blinded to the treatments administered. The treatment schedule was modeled on the standard clinical management of patients with pancreas cancer (Figure 3A). Gemcitabine was administered intravenously (i.v.) once weekly at 50 mg/kg in a three weeks ‘on’ / one week ‘off’ schedule; PEGPH20 (or vehicle) was administered i.v. 24 hours prior to gemcitabine.

Statistical analyses

Data were tested to determine if they met the assumption of normality using the Kolmogorov-Smirnov test. Normally distributed two-group data were analyzed using a t-test. Multi-group data were analyzed using ANOVA followed with the Newman-Kuels multiple comparison post-test. In the instance that two-group data did not conform to normality, the nonparametric Mann-Whitney sum rank test was performed. Kaplan-Meier survival data were analyzed using a Log Rank test. Metastatic disease burden was analyzed using Fisher’s exact test.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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SIGNIFICANCE

More than 90% of cancer drugs brought to the clinic fail. The gap between preclinical promise and clinical reality arises, in part, from the inability of traditional model systems to sufficiently represent the cellular and structural complexity of the solid tumor “organ.” In failing to fully account for the tremendous obstacles to efficacy in autochthonous tumors, potential opportunities to identify and target the most clinically meaningful barriers are also lost. We have found that an unusually abundant matrix glycosaminoglycan fortifies the epithelium and creates a drug-free sanctuary in PDA. Ablating this constituent renders the cancer profoundly vulnerable to standard cytotoxics and holds promise as a distinct therapeutic strategy, while compelling reassessment of conventional chemotherapies previously thought to be ineffective.
HIGHLIGHTS

Extremely high interstitial fluid pressures (IFP) induce vascular collapse in PDA
Hyaluronan (HA) is the major determinant of elevated IFP in PDA
Systemic infusion of an enzyme, PEGPH20, ablates HA from PDA and normalizes IFP
PEGPH20 + gemcitabine therapy permanently remodels the stroma and increases survival
Figure 1. Evolution of the desmoplastic reaction in murine (mPDA) and human PDA (hPDA) (A-E) Masson’s trichrome histochemistry shows robust collagen deposition (blue) at all stages of disease. (F-J) Movat’s pentachrome histochemistry reveals collagen (yellow), GAGs and mucins (blue), and their co-localization (turquoise/green). (K-O) Histochemistry with HA-binding protein (HABP) reveals intense HA content beginning with preinvasive disease (PanIN). (P-T) Activated PSC express α-smooth muscle actin (α-SMA) and are abundant in preinvasive (Q), invasive (R) and metastatic mPDA (S) and hPDA (T), but not in normal pancreata (P). ac, acini; is, islet; d, duct; v, venule; h, hepatic parenchyma; *, metastatic lesions. Scale bars, 25 μm. See also Figure S1.
Figure 2. Elevated IFP compromises vascular function in PDA
(A) Experimental apparatus to measure IFP.
(B and C) IFP probe positioned in PDA at the head of the pancreas (hp). In (B) the probe ends in a tumor obscured by the overlying liver.
(D) IFP measurements in normal pancreata (N), untreated KPC and KC tumors and tumors 2 and 24 hours post-PEGPH20 treatment. *p<0.01 for difference between normal pancreata and PEGPH20 treatment groups; **p<0.05 for difference from all other groups with 1-way ANOVA and Newman-Keuls post-hoc multiple comparison test. Note that IFP levels in normal pancreata and PEGPH20-treated KC (mean = 16 mmHg, n = 4) and KPC (mean = 22 mmHg, n = 5) tumors 24 hours post-treatment were not statistically different. Box and whisker plots: boxes display the lower (25th) and upper (75th) quartiles with a line at the median; whiskers extend from the minimum to the maximum observation.
(E-G) Distribution of CD31+ vessel diameter in normal pancreata (E), untreated KPC tumors (F), and KPC tumors 24 hours post-PEGPH20 treatment (G). Insets show data for vessels of diameter >10 μm. Data represent evaluations from four independent sections from each of four separate animals for each condition and are significantly different (p<0.0001) in the pairwise comparisons with KPC.
(H-J) Multiphoton excitation of fluorescently-conjugated lectin (red) and second harmonic generation imaging of collagen (green) within intact normal pancreas (H), untreated KPC tumors (I), and KPC tumors 24 hours post-PEGPH20 treatment (J). Asterisks in (H) and (J) highlight examples of large functional vessels. Arrows in (I) and (J) indicate perfused (functional) vessels. Note when a rare functional vessel in untreated KPC tumors is identified, it is extremely small relative to those in normal pancreata and PEGPH20 treated tumors. Scale bar, 10 μm for (H-J).
(K-M) Direct fluorescence analysis of Alexa-647-conjugated lectin (red) and doxorubicin (green) in intact normal pancreas (K), untreated KPC tumors (L), and KPC tumors 24 hours after PEGPH20 treatment (M). Arrow in (K) shows example of widely patent, thin-walled
vessel and arrowheads indicate ductal epithelium. Large arrows in (L) and (M) highlight regions magnified in respective inset boxes. The left inset box in (L) shows signal in the far-red Alexa-647 channel (lectin) alone while the right inset box shows merged green (doxorubicin) and red (lectin) channels. Small arrows in inset box of panel (M) identify examples of specific nuclear staining from DNA-bound doxorubicin, as well as the presence of free doxorubicin in the tumor. Asterisks in (M) highlight examples of large, functional lectin-positive vessels loaded with doxorubicin. ac, acini; is, islet; d, duct. Scale bar, 50 μm for (K-M). See also Figure S2.
Figure 3. Gemcitabine+PEGPH20 combination therapy fundamentally changes the tumor vasculature

(A) Treatment schedule for administration of PEGPH20 (P) or placebo and gemcitabine (G). Mice undergoing gemcitabine monotherapy also received placebo (vehicle) injections. Each 4 week course represents one cycle of therapy.

(B) Tumor IFP at survival endpoint after treatment with gemcitabine (G; n=9) or Gem+PEGPH20 (GP; n=9), *p<0.0001. Box and whisker plots: boxes display the lower (25th) and upper (75th) quartiles with a line at the median; whiskers extend from the minimum to the maximum observation.

(C) Gross pathology of transected PDA at endpoint after gemcitabine monotherapy.

(D) Gross pathology of transected PDA at endpoint after combination therapy.

(E,F) Distribution of significantly different (p<0.0001) CD31+ vessel diameter in Gem (E) and Gem+PEGPH20 (F) treated KPC tumors at survival endpoint. Inset graphs show significantly different (p<0.0001) distribution also for subsets of vessels with diameters between 10-20 μm and >20 μm. Data are from four independent sections from five separate animals for each treatment arm. See also Figure S3.
Figure 4. Gemcitabine+PEGPH20 induces objective responses through reduced proliferation and increased apoptosis

(A) High-resolution ultrasound images of PDA before and after the indicated treatment. Scale bar, 1 mm. t, tumor.

(B) Quantitative analysis of tumor volume in Gem (G; n=5) and Gem+PEGPH20 (GP; n=6) treated mice after one cycle of therapy (*p=0.009). Box and whisker plots: boxes display the lower (25th) and upper (75th) quartiles with a line at the median; whiskers extend from the minimum to the maximum observation.

(C) Quantitative analysis of proliferation assessed as % of Ki-67+ cells per image field in untreated (U) and Gem (G) or Gem+PEGPH20 (GP) treated tumors after either two weekly doses of therapy (2 Dose; *p<0.035) or at endpoint (End; *p=0.0009).

(D) Quantitative analysis of apoptosis assessed as % cleaved caspase-3 (CC3)+ cells per image field in untreated (U) and Gem (G) or Gem+PEGPH20 (GP) treated tumors after two doses of therapy (*p=0.035) or at endpoint (*p=0.01). Data in (C and D) are plotted as mean ± SEM. See also Figure S4.
Figure 5. Combination therapy with Gemcitabine+PEGPH20 remolds the tumor stroma
(A and B) IHC for the PSC activation marker α-SMA in PDA at survival endpoint after the indicated treatment. Scale bar, 50 μm.
(C) Quantitative analysis of PSC activation in untreated tumors (U) or tumors treated with Gem (G), PEGPH20 (P) or Gem+PEGPH20 (GP) at survival endpoint (*p=0.005).
(D and E) Representative micrographs from intravital SHG imaging of collagen (green) combined with multiphoton excitation of endogenous fluorescence from NADH (red) in PDA at survival endpoint just prior to euthanasia. Scale bar, 50 μm.
(F) Mean SHG intensity in PDAs at endpoint as a measure of fibrillar collagen concentration (*p=0.028). (G) Percent area occupied by fibrillar collagen in tumors at endpoint (*p=0.028). Data in (C), (F) and (G) are plotted as mean ± SEM. See also Figure S5.
Figure 6. Gemcitabine+PEGPH20 combination therapy decreases metastatic tumor burden and improves survival

(A and B) HA expression in lung metastases from untreated (A) and PEGPH20-treated (B) animals.

(C) Quantitative analyses of proliferation assessed as % Ki-67+ cells in metastases to liver (filled circles) and lung (open squares) after gemcitabine monotherapy (G) or Gem +PEGPH20 (GP) combination therapy vs. (p=0.0012).

(D) Quantitative analysis of apoptosis assessed as % CC3+ cells in liver (filled circles) and lung (open squares) metastases (p=0.0001).

(E) Kaplan-Meier survival curves from time of enrollment in control (Con; n=16), Gem (n=16), PEGPH20 (n=15), and Gem+PEGPH20 treated KPC animals (n=14). (Black bar indicates maximum duration of therapy.) Median overall survival of Gem (55.5 days) and Gem+PEGPH20 (91.5) treated mice are significantly different (*p=0.004). Treatment with PEGPH20 alone showed a trend toward increased survival (median = 63 days) that did not reach statistical significance (p=0.1).

(F) Metastatic burden in Gem+PEGPH20 (GP) treated mice is significantly decreased compared with Gem treatment alone (G) (*p=0.014). See also Figure S6 and Table S1.
Figure 7. Altering physicomechanics and remodeling the stroma in PDA to therapeutic advantage

(A) Intratumoral mechanics in PDA impede diffusion and convection of small molecules.
(B) Enzymatic degradation of stromal HA decreases IFP and relieves physical constraints on small molecule perfusion which can reconstitute in the absence of additional therapy.
(C) Combined enzymatic and cytotoxic therapy permanently remodels the tumor microenvironment to favor the delivery and distribution of small molecules. Blue spheres represent chemotherapy molecules, vessels are shown in red, carcinoma cells in light blue, activated PSC in brown, collagen in green, and HA in yellow. $P_I$, interstitial fluid pressure; $P_V$, intravascular fluid pressure. See text for details.
## Table 1

Summary of metastatic disease burden in *KPC* mice

<table>
<thead>
<tr>
<th>Treatment</th>
<th>% mice with mets</th>
<th># macroscopic mets / mouse (mean +/- S.E.M.)</th>
<th># mets / mouse (mean +/- S.E.M.)</th>
<th>% mice with liver mets</th>
<th>% mice with lung mets</th>
<th>% mice with diaphragm mets</th>
<th>% lymph node involvement</th>
<th>% mice with malignant ascites</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gem+Placebo</td>
<td>93</td>
<td>1.7 +/- 0.35</td>
<td>2.4 +/- 0.35</td>
<td>73</td>
<td>40</td>
<td>40</td>
<td>73</td>
<td>47</td>
</tr>
<tr>
<td>Gem+PEGPH20</td>
<td>43</td>
<td>0.86 +/- 0.4</td>
<td>1.1 +/- 0.4</td>
<td>43</td>
<td>21</td>
<td>21</td>
<td>36</td>
<td>21</td>
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</tbody>
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