Perspective on Mohammed et al., p. 1417

Chemoprevention of Pancreatic Cancer: Ready for the Clinic?
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Abstract
Advances in our molecular, clinical, and epidemiologic understanding of the risk and development of pancreatic cancer offer hope for preventing this disease, which is largely intractable once developed. This perspective on provocative, genetically engineered mouse model work reported by Mohammed et al. (beginning on page 1417 in this issue of the journal) examines the prospects for pancreatic cancer chemoprevention with epidermal growth factor receptor (EGFR) tyrosine kinase inhibitors (TKI). Despite having limited value in advanced pancreatic cancer, EGFR TKIs show promise in the setting of early pancreatic carcinogenesis.

Exocrine pancreatic cancer remains a largely intractable malignancy. Despite some advances in the radiological assessment of pancreatic cancer resectability and improvements in surgical technique, the overall 5-year survival of all patients diagnosed with pancreatic cancer is still only 2% to 3% (1). This poor survival persists despite extensive testing of chemotherapeutic agents and the integration of multiple modalities (primarily surgery, radiation therapy, and chemotherapy) into the management of patients with pancreatic cancer. The lack of progress against this malignancy is thought to be due to two elements inherent to its biology: (a) insidious presentation due to the lack of specific symptoms and signs, often leading to an advanced stage at diagnosis, and (b) striking therapeutic resistance.

The therapeutic resistance of pancreatic cancer is likely to be due to many factors but includes the high frequency of KRAS-activating mutations (KRAS*) and the extensive stromal reaction engendered as the malignancy develops. This extensive stroma is thought to lead to poor delivery of chemotherapeutic agents to the malignant cells (2).

Despite lack of progress in the treatment of established pancreatic cancer, steady advances are being made in our knowledge of patients who are at risk for developing this disease. Our current understanding of the risk for developing invasive pancreatic cancer allows patients at an increased risk to be divided into three general groups: (a) those individuals with known heritable risk factors, such as germ-line mutations in cyclin-dependent kinase inhibitor (CDKN2A), liver kinase B1 (LKB1), BRCA2, and PRSS1 (3–6), or individuals with two or more first-degree family members diagnosed with pancreatic cancer (7); (b) patients with mucinous cystic neoplasms of the pancreas (intraductal papillary mucinous neoplasm (IPMN) and mucinous cystic neoplasm (MCN); ref. 8); and (c) individuals with combinations of specific epidemiologic risk factors such as cigarette smoking, long-standing type II diabetes, and obesity (9, 10). So, although our ability to identify patients at risk of developing pancreatic cancer has improved, we have no interventions that can mitigate this risk other than partial or total pancreatectomy. Clearly, surgical resection is a radical intervention for patients whose lifetime risk of developing pancreatic cancer may be only elevated slightly over the baseline risk in the general population.

Like other epithelial cancers of the gastrointestinal tract, pancreatic cancer is thought to evolve through non-malignant precursor lesions termed pancreatic intra-epithelial neoplasia (PanIN), and these lesions progress through states of increasing cytologic atypia and dysplasia through the acquisition of increasing numbers of signature genetic alterations (11). The gatekeeper mutation for pancreatic cancer is KRAS*, with loss of tumor suppressor genes such as CDKN2A, p53, and Smad4/Dpc4 occurring very commonly as the PanIN lesions progress to carcinoma in situ and invasive pancreatic cancer. Recently, these pathologic and genetic observations derived from patients have been confirmed using transgenic mouse models in which the early development and progression of pancreatic cancer can be recapitulated through the expression of KRAS* and accelerated by engineered loss of CDKN2A or p53 specifically in pancreatic epithelium (12–14).

In this issue of the journal, Mohammed et al. report their study using the p48Cre/+; LSL-KRASG12D/+ transgenic mouse model of pancreatic cancer and show that the epidermal growth factor receptor (EGFR) tyrosine kinase inhibitor (TKI) gefitinib prevents progression of PanINs to invasive pancreatic cancer (15). They argue that "these results have important implications for human pancreatic cancer chemoprevention."
What is the evidence that examining such an intervention in patients at risk for pancreatic cancer is warranted? Qualitative protein expression data from human pancreatic cancer specimens have shown that EGFR is frequently overexpressed. However, genetic analyses have failed to identify mutations, amplification, or activating translocations affecting EGFR, suggesting that [at least in the advanced-disease setting] inhibition of EGFR would be anticipated to have only limited clinical effect. This fact has been borne out in prospective clinical trials that combined gemcitabine with the EGFR TKI erlotinib or the humanized monoclonal EGFR antibody cetuximab in patients with advanced pancreatic cancer (16, 17).

However, the study described by Mohammed et al. is provocative in that it suggests that targeting EGFR early in pancreatic carcinogenesis may be effective despite the limited value of this approach in advanced pancreatic cancer. So, are there data in addition to this study to suggest that gefitinib or other small-molecule EGFR TKIs represent a viable approach to pancreatic cancer chemoprevention? Right now, the picture looks mixed. As pointed out above, in the advanced pancreatic cancer setting, the effect of erlotinib is quite modest, and because we do not yet understand which pancreatic cancer patients are likely to benefit from erlotinib, the anticipated effect of small-molecule EGFR inhibitors in a chemopreventive setting may also be modest. Despite this, the Mohammed et al. study offers one hopeful experimental observation. That is, the effect of EGFR inhibition may be more significant early in pancreatic carcinogenesis. Reduction of PanIN-1 and PanIN-2 lesions was much more pronounced than was the effect on the more advanced PanIN-3 histology. This observation should be confirmed in other studies and with genetically engineered animal models of pancreatic carcinogenesis that incorporate additional genetic changes, but it suggests that the therapeutic resistance conferred by KRAS* may require the acquisition of additional genetic changes. Thus, although the mechanism of such a differential effect is unknown, this observation could suggest that EGFR inhibition may be effective early in pancreatic carcinogenesis, yet ineffective later—an observation that would be consistent with the results of the Mohammed et al. study and the limited effect of erlotinib in advanced-disease clinical trials. Last, early results from a study of erlotinib in patients diagnosed with IPMN have shown that at least one patient completely responded radiographically to this intervention.3

A general concern is whether the genetically engineered murine models that are completely dependent on a dominant oncogene (e.g., KRAS*) can inform us about a disease in humans that is much more genetically complex (18). Expression of KRAS* alone or expression of KRAS* coupled with loss of p53, CDKN2A, or other tumor suppressors leads to disease that evolves over months in mice and may not reproduce the full spectrum of genetic and biological heterogeneity present in human pancreatic cancer, a disease that may take decades to develop. Of course, the answer to the question of the applicability of results generated in genetically engineered mouse models to humans ultimately depends not so much on how well the transgenic models recapitulate aspects of the biology of human pancreatic cancer, but whether these models can prove predictive for the identification of clinically useful interventions. Because the predictive track records of subcutaneous and orthotopic models of pancreatic cancer have been poor, there is considerable hope that the transgenic models that incorporate our increasing understanding of the driving genetic events in pancreatic carcinogenesis will provide the needed insights.

Despite the challenges inherent in modeling human cancers in mice, there is a clear need to identify novel approaches/agents for prevention and therapy. Thus far, the animal models suggest that there is likely to be an important convergence between targets for therapy and prevention, and the positive effect of an EGFR inhibitor early in pancreatic carcinogenesis may be different (and more powerful) than the same drug administered to patients with advanced pancreatic cancer (19). The reasons for this difference remain unknown, but with further progression of cancer, there are a greater number of contributing molecular events, increasing biochemical complexity, cancer cell heterogeneity, and extensive tumor-stromal interactions. All of these alterations may conspire to make advanced pancreatic cancer highly resistant, yet early in pancreatic carcinogenesis, these changes are likely to be less fully developed, perhaps leading to drug vulnerabilities that are not encountered in the more advanced setting.

Our next hurdle will be to design clinical trials that can assess the ability of anticancer agents, administered early in pancreatic carcinogenesis, to have a beneficial clinical effect. Increasingly sophisticated transgenic animal models have also suggested that the sequence of mutational events can alter the morphology of pancreatic cancer precursor lesions. For example, concomitant expression of KRASG12D and haploinsufficiency of the Smad4/Dpc4 tumor suppressor gene (KRASG12D+/−;Dpc4floxFlo; p4GCre) gives rise to mucinous cystic neoplasms that can then progress to invasive ductal adenocarcinoma through loss of heterozygosity of Dpc4 and mutation of either p53 or p16 (20). As indicated above, patients with mucinous cystic neoplasms of the pancreas have very similar genetics and are known through clinical studies to be at risk for invasive pancreatic cancer. Furthermore, because of the (over)utilization of computerized abdominal imaging, increasing numbers of patients are being diagnosed with previously unsuspected pancreatic cysts, some of which will be confirmed to be mucinous cystic neoplasms.

Thus, gefitinib or other rationally targeted agents should be studied in the KRASG12D+/−;Dpc4floxFlo; p4GCre and KRASG12D+/−;Dpc4floxFlo; p4GCre transgenic models with the goal to then examine successful interventions in clinical trials of patients with newly identified mucinous cystic neoplasms (IPMN and MCN). As we learn more about how to effectively screen for pancreatic cancer precursor

3 S. Lipkin, personal communication, June 2010.
lesions, we will need to begin to examine the effects of medical interventions, reserving surgery for those patients with more advanced PanINs or progressing mucinous lesions or for those who do not respond to chemopreventive interventions. An enormous challenge will be to design these studies in a way that provides definitive evidence that the intervention inhibits pancreatic carcinogenesis without risking the development of invasive pancreatic cancer and to accomplish this with acceptable toxicity.

One approach to this problem is suggested in Fig. 1. As outlined in this figure, the primary goal would be to establish that the intervention increases the interval between entry into the trial and progression of the mucinous lesion size or development of radiographic findings that have been recommended by consensus to necessitate surgical intervention [time interval 1 (T1); ref. 21]. In addition, the time interval to progression for patients initially randomized to placebo (T2) can be compared with the open-label intervention (T3) such that these patients can serve as their own control. Unfortunately, due to the fact that we are only now gathering experience and natural history data on asymptomatic mucinous cystic lesions of the pancreas, there are many gaps in our knowledge. These questions include the following: (a) What is the rate of progression (enlargement) in MCNs and IPMNs over time? (b) Is the rate of progression the same for MCNs and IPMNs? (c) How much heterogeneity is there in the rate of progression for patients classified as MCN versus IPMN, and do these differences have a molecular basis? (d) Is the rate of change constant over time? (e) In the absence of development of worrisome signs indicating the need for surgery, what increase in cyst size defines progressive disease (0.5 cm is suggested in Fig. 1, but lesser or greater degrees of change could be considered)? (f) How reproducibly can we image small changes in cyst size, and what is the best imaging study to use [computed tomography versus magnetic resonance/magnetic resonance cholangiopancreatography versus endoscopic ultrasound versus some combination of these modalities]? Other issues to consider include how extensive the initial evaluation would need to be to diagnose IPMN and MCN: (a) Should all newly identified cysts be aspirated to confirm the presence of mucin? (b) Are there cytologic criteria that should be included in the diagnostic criteria (e.g., the presence of ovarian stroma in MCN)? (c) Should KRAS* testing be done on all patients? These and other eligibility and protocol design issues would need to be decided on as an intervention trial was developed.

In summary, whether or not gefitinib represents a reasonable intervention to inhibit pancreatic carcinogenesis, many important issues are raised by the provocative work of Mohammed et al. (15). It will be key to validate these observations with additional genetically engineered animal models, but as this study indicates, these models can be readily used to examine the effect of a variety of targeted interventions. It is hoped that we will discover that the models provide insights into interventions that can be rationally tested in well-designed clinical trials, leading to prevention of this dreaded malignancy.

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