Intraductal Papillary Mucinous Neoplasia (IPMN)

Highlights from the "2010 ASCO Gastrointestinal Cancers Symposium". Orlando, FL, USA. January 22-24, 2010

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Summary
The diagnosis and treatment of intraductal papillary mucinous tumors (IPMN) of the pancreas has evolved over the last decade. IPMN is a disease of the ductal epithelium and represents a spectrum of disease, ranging from benign to malignant lesions, making the early detection and characterization of these lesions important. As with villous adenomas of the colon, not all IPMNs will develop into adenocarcinoma. Definitive management is surgical resection for appropriate candidates, as benign lesions harbor malignant potential. Growing controversy revolves around issues of natural history, management of small-branch-duct lesions, ability to predict malignancy and/or progression, and surveillance strategies. Given these controversies, novel methods are needed to help in detecting and classifying IPMNs’ malignant potential so that appropriate treatment can be administered. The authors review abstracts from the 2010 ASCO Gastrointestinal Cancers Symposium held in January 2010, including biomarkers helping to classify IPMNs: IL-8 and IL-1β from IPMN cyst aspirates (Abstract #133), and Foxp3/CD4/CD25 cells (Abstract #148) in peripheral blood. Future studies will hopefully provide insight into the many unanswered questions.

Introduction
Intraductal papillary mucinous neoplasms (IPMN) have been reported to account for approximately 7% of clinically diagnosed pancreatic neoplasms and up to 50% of incidentally detected pancreatic cysts [1]. Since the initial description of IPMN by Ohashi et al. [2], the incidence of these mucin-producing epithelial tumors of the exocrine pancreas has been increasing [3]. This is probably attributed to improvements in technology and diagnostic imaging as well as more distinct nomenclature [1]. The World Health Organization (WHO) introduced the term “intraductal papillary mucinous tumor” in 1996, which was later renamed as IPMN in 2000 [4, 5].

IPMN may manifest as recurrent pancreatitis, with or without hyperamylasemia, steatorrhea, diabetes, and weight loss. On the other hand, patients may be entirely asymptomatic, with a tumor found on imaging performed for a different indication. On imaging, IPMN appears as a dilated pancreatic duct, full of mucin, which extrudes through a bulging papilla. Tumors may arise from the main duct, side-branches, or they may display a mixed pattern of involvement.

Updates from the 2010 ASCO Gastrointestinal Cancers Symposium

Abstract #133: Cyst fluid cytokines to distinguish low- and high-risk intraductal papillary mucinous neoplasms (IPMN) [10]

Cyst formation in the pancreas may induce or produce an immunological response and could result in release of cytokines or other soluble factors into cyst fluid that...
could correlate with the risk of having a high risk IPMN. Before surgical resection, pancreatic cyst fluid was aspirated from 40 patients. Patients were then grouped based on the grade of dysplasia in resected tissue. Cyst fluid cytokine levels (IL-2, 4, 5, 8, 10, 12, 13, TNF-alpha, IFN-gamma) were determined using a multiplex ELISA methodology.

Of the cytokines measured, both IL-1beta and IL-8 levels were found elevated in patients’ cyst fluid with high-grade dysplasia. IL-1beta levels in low-grade dysplasia were 0.2±0.1 pg/mL and 539±255 pg/mL in high grade group. Cyst IL-8 levels were 2,893±836 pg/mL and 8,089±2,288 pg/mL in low vs. high grade lesions respectively.

IL-1beta is a cytokine produced principally by mononuclear phagocytes but also by various other cell types including keratinocytes, epithelium and cells of the central nervous system. Elevated levels of IL-1beta have been implicated in sepsis, cachexia, rheumatoid arthritis, chronic myelogenous leukemia, asthma, psoriasis, inflammatory bowel disease, anorexia, AIDS, and graft-versus-host disease associated with bone marrow transplants. In addition, medical literature indicates that IL-1beta is one of the key mediators of immunobiological responses to physical stress. IL-8 is produced by stimulated monocytes, macrophages, fibroblasts, endothelial cells, keratinocytes, melanocytes, hepatocytes, chondrocytes, and a number of tumor cell lines. Elevated concentrations of IL-8 have been observed in patients with psoriasis, rheumatoid arthritis, chronic polyarthritis, tumor development and hepatitis C.

The source of these cytokines could be interesting and future studies may want to investigate the immunophenotype of cells found in cyst in addition to immunohistological localization of these cytokines and associated cells in the tumor microenvironment to see how they are interrelated and if there are any other biomarkers that may be predictive of malignant IPMN.

**Table 1. Review of select abstracts from 2010 ASCO Gastrointestinal Cancers Symposium.**

<table>
<thead>
<tr>
<th>Abstract</th>
<th>Tool/Marker</th>
<th>Highlight</th>
<th>Comment</th>
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<tbody>
<tr>
<td>#133</td>
<td>Cyst cytokines from aspirates</td>
<td>Cyst IL-1beta and IL-8 are elevated in patients with high grade IPMN compared to low grade IPMN</td>
<td>Immunophenotype of immune cells in aspirated fluid is not reported in abstract</td>
</tr>
<tr>
<td>#148</td>
<td>Foxp3/CD4/CD25 T-regulatory cells from peripheral blood and resected tumor</td>
<td>Patients with no recurrence had low peripheral T-regulatory cells compared to plasma or transcriptional profile of these cells in abstract</td>
<td>No information given regarding cytokine profile of cells in abstract</td>
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**Abstract #148: Evaluation of Foxp3/CD4/CD25/T cell (Treg) in peripheral blood as a biomarker for the aggressiveness of intraductal papillary mucinous neoplasm [11]**

Differentiating and predicting the benign and potentially malignant forms of IPMN located in ductal adenomas is sometimes a challenge. More methods are needed to help in predicting benign from malignant forms of IPMN. Recent studies have found increased Foxp3/CD25/CD4 cells in IPMN tumors and their presence heralds a worse prognosis [12]. Foxp3/CD25/CD4 cells can also be found in peripheral blood, and Ikemoto et al. previously reported that the percentage of peripheral Foxp3+, CD4+, CD25 T-cells were increased in patients with advanced pancreatic cancer [13].

The work presented at the 2010 ASCO Gastrointestinal Cancers Symposium is a continuation of this line of research in the setting of benign and cancerous IPMN types focusing on the percentage of peripheral Foxp3+ T-regulatory cells in tumor and blood in relation to histological aggressiveness of resected IPMNs and recurrence. They were able to show that patients with peripheral T-regulatory cells levels below 2.5% had no recurrence after surgery. Conventional tumor markers, CEA, CA 19-9, Span-1, and DUPAN-2 showed no correlation to the aggressive nature of the IPMNs and preoperative imaging only had a weak correlation. Histological aggressiveness was also positively correlated with the number of Foxp3+ T-regulatory cells in resected tumors. These results are promising because the peripheral Foxp3 levels seemed to correlate better than many conventional tumor markers with tumor aggressiveness. Since cyst fluid contains cytokines and other factors that may help recruit T-regulatory cells, it would be interesting to isolate these T-regulatory cells and determine what type of immunokines they are.

**Table 2. Biomarkers helpful in differentiating benign and malignant IPMN or invasive pancreatic cancer.**

<table>
<thead>
<tr>
<th>Method</th>
<th>Sensitivity</th>
<th>Specificity</th>
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<tbody>
<tr>
<td>Carcinoembryonic antigen (CEA; cystic) over 200 ng/mL [20]</td>
<td>47% (malignant vs. benign IPMN)</td>
<td>40%</td>
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<tr>
<td>Carbonic anhydrase 19-9 (CA19-9; cystic) over 10,000 U/mL [21]</td>
<td>80% (malignant vs. benign IPMN)</td>
<td>50%</td>
</tr>
<tr>
<td>Span-1 (serum) over 400 U/mL [22, 23]</td>
<td>81-94% (pancreatic cancer detection)</td>
<td>75%</td>
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<tr>
<td>DUPAN-2 [24]</td>
<td>48-80% (pancreatic cancer detection)</td>
<td>65%</td>
</tr>
<tr>
<td>PAM4 [25]</td>
<td>77% (pancreatic cancer detection)</td>
<td>95%</td>
</tr>
<tr>
<td>EUS-FNA-cytology [21]</td>
<td>75% (malignant vs. benign IPMN)</td>
<td>91%</td>
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</table>
producing to see if any of these are potential biomarkers for malignant potential of IPMN (Table 2).

**Discussion**

IPMN is a disease of the ductal epithelium. There is a progression in the degree of dysplasia from IPMN adenoma, to borderline IPMN with dysplasia, to IPMN with carcinoma in situ, to invasive carcinoma. This progression is similar to the progression from adenoma to carcinoma that is seen in colorectal carcinoma and the progression of pancreatic intraepithelial neoplasm seen in solid epithelial tumors of the pancreas (Figure 1) [14]. An accumulation of molecular abnormalities is associated with progression from IPMN adenoma to invasive carcinoma, but the time course for progression and the malignant potential for each lesion are unknown. According to early reports [15, 16] invasive carcinoma was present in 35% to 50% of patients diagnosed with IPMN.

The data from these two abstracts raise some interesting possibilities for determining the nature of IPMNs found incidentally or even from symptomatic patients. Cystic cytokines obtained from EUS-FNA, along with peripheral and tissue CD4-regulatory cells, represent two additional markers to help classify the malignant potential of IPMNs and also giving us a window into what is evolving in the tumor microenvironment and host to allow the tumor to progress. Recent work has helped us to appreciate that the tumor microenvironment can be an active participant in the formation or maintenance of a malignancy [17]. Local and systemic cytokines and T-regulatory cells are known to be players in development and maintenance of a malignant phenotype [18, 19]. The ability to monitor these changes locally in the case of cystic cytokines and systemically, in the case of peripheral Foxp3/CD4/CD25 cells is promising. Because of genomic instability, it is likely that IPMNs will evolve different ways to manipulate the local and systemic environment to favor their survival, and in all likelihood, there are other biomarkers which may be discovered that are common in the development of malignant IPMN. As more data is gathered, algorithms combining multiple parameters will help give physicians greater confidence regarding the nature of newly discovered IPMNs. These two abstracts are promising and clinicians should initiate more studies to determine sensitivity and specificity compared to other methodologies to see if incorporating these two biomarkers offers any improvements in classifying IPMNs.

**Conflict of interest**

The authors have no potential conflicts of interest.

**References**


