How grim is pancreatic cancer?

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Abstract

Pancreatic ductal carcinoma continues to be the most lethal malignancy with rising incidence. It is the fourth most common cause of cancer death in the western world due to its low treatment success rate. In addition, because of its rapid growth and silent course, diagnosis is often only established in the advanced stages. As one of the most aggressive malignancies, the treatment of this disease is a great challenge to clinicians. This paper reviewed the natural history of pancreatic cancer, the current clinical practice and the future in pancreatic cancer management.

Introduction

Pancreatic cancer is one of the most aggressive human malignancies, as 50% present with metastatic disease and 35% with locally advanced disease.1,2 It is the 13th commonest cancer with 200,000 cases per year worldwide, 6000 cases per year in the UK and the fourth leading cause of cancer death in the Western world.2 There is an increasing incidence of this disease affecting 8-12 per 100,000 of the population per year. Whether this increased incidence is real or whether it reflects advances in diagnostic imaging is unknown. Its poor prognosis is manifested in an overall median survival of 4.4 months, and a 5-year survival of 9.7%.3 Diagnostic problems arise because the symptoms are late and non-specific, there is no effective screening process and there is no specific high-risk group. Since conservative oncological therapies have failed to show any benefit in long-term survival, resection remains the only modality of treatment offering any possibility of cure.4,7 Unfortunately, only 10-20% with head and less than 3% of body/tail cancers are candidates for resection.1,5 In the past 20 years, there is also only a modest increase in long-term survival with a median survival of 12 months, and 5-year survival rate of 15-26% after potentially curative resection.7 This has led to the development, of wider resections in the hope of increasing long-term survival, or more conservative approaches to improve the quality of survival.1,4 The role of these newer procedures is controversial and the Whipple’s partial pancreaticoduodenectomy (PD) procedure is still favored by most surgeons.8-10 Traditional chemotherapy remains the standard treatment for advanced pancreatic cancer. Regimens like FOLFIRINOX (5-FU, leucovorin, irinotecan, and oxaliplatin) or gemcitabine and nab-paclitaxel have been used to palliate symptoms and prolong survival.11 This paper elucidates the grimness of pancreatic cancer and the potential for cure.

Risk factors and genetics

The etiology is largely unknown and the risk factors are listed in Table 1. Experimental studies have suggested that pancreatic carcinoma (PC) cell lines produce soluble factor(s) that can impair glucose metabolism and cause hyperglycemia.12-14 The 7-10% of pancreatic carcinoma are related to genetic factors with several well-defined genetic syndromes of familial pancreatic cancer (FPC) (Table 2).15,16 Although there are no specific genetic mutations identified for the majority of FPC (70%), relatives of FPC kindred have a high risk of pancreatic cancer.17 K-Ras gene mutations have been found in most pancreatic cancers. As a prediction of poor prognosis, the detection of K-ras mutations may be a useful prognostic factor for pancreatic cancer patients. K-Ras mutations are associated with a worse overall survival in pancreatic cancer patients, especially when mutations are detected inliquid biopsies or fresh frozen tumor tissue samples.18 Emerging studies show that cancer stem cells (CSCs) regulate several mechanisms underlying drug resistance, carcinogenesis, and metastases development in various types of cancer including pancreatic cancer.19,20 Recent studies dissected the role of microRNAs (miRNAs) and pancreatic CSCs on the modulation of pancreatic cancer etiology and progression, shedding light on their importance as potential therapeutic targets for pancreatic cancer.21,22 MicroRNAs are small noncoding RNAs involved in the regulation of gene expression at posttranscriptional level by binding to the 3 untranslated regions or the open reading frames of target genes. Oncogenic miR-21 lead to the repression of miRNA translation or to the degradation of target miRNAs. They modulate the proliferation and the chemoresistance of pancreatic cancer cells.23,24 In addition, there is a correlation between miR-21 expression and the clinical outcome of patients with pancreatic cancer through involvement of the PI3K/AKT pathway.25,26

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Pathophysiology and prognosis in ductal pancreatic cancer

The majority (95%) of pancreatic cancers develop as adenocarcinomas from the ductal cells of the exocrine pancreas. Of them 1% is of acinar origin, 1% is of non-epithelial origin and 3% are of uncertain cellular origin.25 It is associated with an accumulation of mutations with progressive morphological changes (Table 3).26 The current model proposes a progression from normal cuboidal to low columnar epithelium through a series of lesions termed pancreatic intraepithelial neoplasia to invasive carcinoma.26,27 Prognosis in patients with pancreatic cancer depends on the tumor stage at time of diagnosis and tumor biology. The pattern of growth is characterized by early spread into local tissues, lymphatic and perineural sites, venous invasion and peritoneal metastases. Free cancer cells in the peritoneal cavity have been observed in 20-40% of cases, even in patients undergoing resection.28 Regional lymph-node metastasis occurred in 30% of patients with very small primary cancers and 64% of T1 primary cancer had lymph node involvement.29 Careful histological studies in a large series of resected pancreatic cancers revealed cancer dissemination in the lymph nodes in 89%; lymph node metastases in 77%, intrapancreatic neural invasion in 92% and a neural and nerve plexus invasion outside the pancreas in 45%.30 Thus even though the surgeon may be able to offer resection to >20% of patients with pancreatic cancer, the possibility of cure is gravely limited by the extent of early or occult micrometastases.

Investigations

Because of the poor prognosis care is taken not to over investigate or embark on treatment strategies based on unrealistic expectations of patients or their families. It is wise to use the fewest, least invasive and least expensive tests for patients not suitable for major resection. The investigations begin with baseline blood tests including a full blood count, renal function (urea and electrolytes), liver function, blood glucose, amylase and the tumor marker carbohydrate antigen (CA) 19-9. Many patients are anemic as a result of nutritional deficiency and chronic blood loss. The occurrence of frank gastrointestinal bleeding suggests the diagnosis of ampullary or duodenal carcinoma but stool examination usually reveals occult blood in patients with carcinoma of the pancreatic head. A rapid elevation of serum bilirubin and alkaline phosphatase on serial measurements of liver function will suggest a diagnosis of periampullary malignancy. Transaminase levels become elevated to a lesser degree reflecting injury to the hepatocytes as a result of unrelied biliary obstruction. Blood glucose may be elevated.3 A transabdominal ultrasound should detect a pancreatic tumor and assess the liver, extrahepatic biliary tree and the pancreas. Its sensitivity of 70% falls to below 30% for tumors <2 cm in diameter.30 A pancreatic protocol helical computed tomography (CT) scan of the abdomen would confirm the diagnosis and provide more detailed information for accurate staging. By assessing the tumor size, local extension/infiltration, intra/extrahepatic dilatation and metastasis it

Table 1. Risk factors for pancreatic adenocarcinoma.

<table>
<thead>
<tr>
<th>Age</th>
<th>80% aged 70-80 years; 1% &lt;40 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td>Women &gt; men</td>
</tr>
<tr>
<td>Race</td>
<td>African Americans &gt; whites &gt; Asian Americans</td>
</tr>
<tr>
<td>Smoking, alcohol, coffee, high fat high protein low fiber diet</td>
<td>10× relative risk if &gt;2 packs/day</td>
</tr>
<tr>
<td>Chronic pancreatitis</td>
<td>5-15-fold ↑ risk</td>
</tr>
<tr>
<td>Hereditary pancreatitis</td>
<td>50-70-fold ↑ risk</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>2× relative risk diabetic onset &gt;3 years before diagnosis</td>
</tr>
<tr>
<td>Hereditary (4-16%)</td>
<td>3× relative risk diabetic onset &gt;2 years before diagnosis</td>
</tr>
<tr>
<td>Abdominal surgery</td>
<td>2.5× the relative risk:</td>
</tr>
<tr>
<td>Occupational</td>
<td>Prior partial gastrectomy, cholecystectomy</td>
</tr>
<tr>
<td>Hereditary (4-16%)</td>
<td>Naphthylamine, ethyl dichloride, benzidine, metal-gas workers, chemists</td>
</tr>
</tbody>
</table>

Table 2. Syndromic familial pancreatic cancer.

<table>
<thead>
<tr>
<th>Genetic syndrome</th>
<th>Germline mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hereditary pancreatitis</td>
<td>PRSS1</td>
</tr>
<tr>
<td>HNPCC (Lynch 11)</td>
<td>hMSH2, hMLH1</td>
</tr>
<tr>
<td>Familial breast cancer</td>
<td>BRAC-2</td>
</tr>
<tr>
<td>FAMMM syndrome</td>
<td>P16</td>
</tr>
<tr>
<td>FAP</td>
<td>APC</td>
</tr>
<tr>
<td>Peutz-Jeghers syndrome</td>
<td>STK11/LKB</td>
</tr>
</tbody>
</table>

HNPCC, hereditary non-polyposis colorectal cancer; FAMMM, familial atypical multiple mole melanoma; FAP, familial adenomatous polyposis.

Table 3. Frequency of mutations in pancreatic cancer.

<table>
<thead>
<tr>
<th>Mutation</th>
<th>Incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>K-ras</td>
<td>95</td>
</tr>
<tr>
<td>P16/CDKN2A</td>
<td>80</td>
</tr>
<tr>
<td>P21</td>
<td>75-85</td>
</tr>
<tr>
<td>TP53</td>
<td>50-75</td>
</tr>
<tr>
<td>Cyclin D1</td>
<td>95</td>
</tr>
<tr>
<td>DPC4/MADH4</td>
<td>55</td>
</tr>
<tr>
<td>Telomerase</td>
<td>95</td>
</tr>
<tr>
<td>BRCA-2</td>
<td>7-10</td>
</tr>
<tr>
<td>LKB1/STK11, M KK4, TGFβ 1/11, RB1</td>
<td>5</td>
</tr>
</tbody>
</table>

[Oncology Reviews 2016; 10:294]
Pre-operative staging

Preoperative staging is based on the AJCC TNM staging system (Table 4). The Japanese Pancreatic Society (JPS) classification is similar to TNM except lymph node groups are clearly defined. Staging laparoscopy, intraoperative liver US and peritoneal cytology may improve accuracy of staging and avoid unnecessary surgery in patients with occult peritoneal disease in up to 10%.38 The criteria for unresectability at staging laparoscopy are: i) histologically confirmed hepatic, serosal, peritoneal, or omental metastases; ii) extrapancreatic extension of tumor, e.g., mesocolic involvement; iii) histologically confirmed coeliac or high portal vein tumor involvement; iv) invasion or encasement of the coeliac axis, hepatic artery or SMA.4,39

Is there a role for biopsy?

Operative biopsy of pancreatic tumor is rarely necessary. Pre-operative ERCP biopsy, brushings, cytology or endoluminal US will differentiate chronic pancreatitis from carcinoma of pancreas and at least a fistula may occur but into the duodenum following an endoscopic biopsy. US or CT-guided fine-needle biopsy is avoided in patients with potentially resectable tumors as it can cause peritoneal seeding and spread of the tumor.40 In addition, there is a high rate of false negativity, especially if the tumor is very small or with a fibrotic reaction. However, whether benign or malignant, resection of an obstructive pancreatic lesion is beneficial to the patient. Intraoperative trucut biopsy of an inoperable pancreatic cancer at laparotomy is justifiable for documentation.36

Criteria for resectability and oncological standards of surgical resection

There is a growing consensus on the radiological definitions of resectable, borderline resectable and unresectable, and the National Comprehensive Cancer Network in the USA has endorsed a modified consensus from the Americas Hepato-Pancreato-Biliary Association,

Table 4. American Joint Committee on Cancer (AJCC) TNM staging of pancreatic cancer.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Tis Stage</th>
<th>N0</th>
<th>M0</th>
<th>Localized within pancreas</th>
</tr>
</thead>
<tbody>
<tr>
<td>T0</td>
<td>No evidence of primary tumor</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1</td>
<td>Tumor limited to the pancreas, &lt;2 cm in greatest diameter</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T2</td>
<td>Tumor limited to the pancreas, &gt;2 cm in greatest diameter</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T3</td>
<td>Tumor extends beyond pancreas but no involvement of celiac axis or SMA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T4</td>
<td>Tumor involves the celiac axis or SMA (unresectable)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N0</td>
<td>No regional lymph node metastases</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N1</td>
<td>Regional lymph node metastases</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M0</td>
<td>No distant metastasis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M1</td>
<td>Distant metastasis</td>
<td></td>
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</table>

SMA, superior mesenteric artery.
the Society of Surgical Oncology and the Society for Surgery of the Alimentary Tract. A tumor is potentially resectable if it can be technically removed with negative margins (R0 resection) without compromising the vascular supply to the liver (hepatic artery) or small bowel (superior mesenteric artery). Involvement of adjacent organs (e.g., duodenum or transverse colon), regional lymph nodes, portal vein (partial involvement), gastroduodenal artery, are not contraindications to resection, as these structures can be removed en bloc with the tumor to achieve an R0 (no tumor cells within 1 mm) resection. An R0 resection for ductal pancreatic cancer must include an N1 and N2 lymph node dissection, perivascular connective tissue dissection and a standardized retroperitoneal soft-tissue dissection. Favorable prognostic features include negative resection margin, negative lymph nodes, well/moderately differentiated carcinoma, primary <2 cm diameter and no perineural or lymphovascular invasion. Patients with negative lymph nodes have significantly higher 3- and 5-year survival rates than patients with positive lymph nodes. However, most R0 resections (70%) are actually R1 (one or more tumor cells within 1 mm) resections as a result of microscopically incomplete resection and the biological features of the tumor such as frequent neural invasion. This highlights the importance of specialized surgeons and pathologists in the treatment of this condition.

Extended vs standard lymphadenectomy

Several centers have reported a survival benefit with extended (radical) lymphadenectomy (standard plus resection of lymph nodes along arterial supply including an en-bloc lymphadenectomy of the hepatoduodenal ligament) compared with standard lymphadenectomy (resection of peripancreatic, periduodenal and perigastric lymph nodes). However, data from randomized control trials (RCTs) and a meta-analysis did not show any benefit, but a potentially increased morbidity with extended lymphadenectomy. Extended lymphadenectomy might thus not be recommended outside of adequately powered RCTs or specialist centers as it is also apparent that survival in patients with an R0 resection including N1 and N2 lymph node dissection is only marginally longer than in those with an R1 resection provided adjuvant chemotherapy is used. However, the surgical dexterity and precision of robotic surgery will facilitate extended lymphadenectomy with minimal morbidity.

Portal vein resection

Historically, portal vein involvement was a contraindication to resection. However, PV or SMV encasement is now considered to be related to tumor location, rather than biological behavior. Due to the unsatisfactory results of the standard Whipple’s (PD) resection and total pancreatectomy, Fortner in 1973 described regional pancreatectomy in order to achieve a negative resection margin and improve long-term survival. This consists of an en bloc resection of the tumor with an adequate peripancreatic soft tissue margin, regional lymph nodes with portal vein resection (Type I) or resection/reconstruction of a major artery (Type II). The reconstruction can very often be achieved by a direct end-to-end anastomosis of the venous remnants or the resected PV replaced if necessary with Dacron graft or saphenous interpositional vein graft. Early results were poor (morbidity 67%; mortality 23%) with 3% 3-year survival. Portal vein resection was associated with longer operating time, increased blood loss, increased perioperative morbidity and mortality as compared to standard PD (Whipple’s). Regional pancreatectomy is now associated with <4% mortality, 26% 5-year survival and reduced morbidity in high volume centers due to: i) advances in radiological and surgical techniques; ii) improved staging; iii) better patient selection; and iv) adjuvant chemotherapy. PD with PV resection should only be performed at centers with expertise in complex pancreatic surgery and only if an R0 resection can be achieved. However, PD with PV resection has similar survival compared to standard PD (in absence of SMA or coeliac axis involvement). Resection margins (R0 vs R1/R2) and lymph node status are more important than portal vein involvement per se in terms of long-term survival.

Management options

Role of multimodal treatment in pancreatic cancer

Surgery remains the only chance of cure for pancreatic cancer, but only 15-25% of patients present with resectable disease at the time of primary diagnosis. Important goals in clinical research must therefore be to allow early detection with suitable diagnostic procedures, to further broaden operation techniques and to determine the most effective perioperative treatment with either chemotherapy and/or radiation therapy. Multimodal treatment including adjuvant chemotherapy and radiotherapy for resectable pancreatic cancer was superior to conventional surgical treatment with a four-year survival rate of above 40% in the treatment group. Surgical resection without adjuvant treatment produces favorable five-year survival only in patients with early stage I pancreatic cancer. Tsuchiya et al. reported a 5-year survival rate of only 30% in patients with stage II cancer. The results of the Gastrointestinal Tumor Study Group using adjuvant radio-chemotherapy following resection of pancreatic cancer revealed a median survival of 21 months in the group randomized to treatment compared to 10.9 months in the control group. The two-year actuarial survival was 46%, and 18% in the control group and the five-year probability survival in the treatment group was above 20%. The European Study Group for Pancreatic Cancer (ESPAC-1-Trial) showed a 5-year survival benefit of adjuvant chemotherapy with gemcitabine for 6 months to patients with lymph node involvement (24% vs 9% 5-year survival). Locoregional chemotherapies administered via the hepatic artery of a regimen comprising 5-fluorouracil (3FU) folinic acid, mitoxantrone and cis-platinum for 6 months in Stage II and III cancers showed improvement of the median survival from 9.3 to 16 months compared to stage I cancers with resection only. While gemcitabine is the main drug in the chemotherapy, recent studies indicate the effectiveness of combination schemes, in particular, nab-paclitaxel plus gemcitabine over gemcitabine monotherapy or when FOLFIRINOX fails for metastatic pancreatic cancer. Additionally, therapies with a broader mechanism of action are emerging (stoma depletion, immunotherapy, anti-inflammation), raising hopes for more effective adjuvant and neoadjuvant treatment concepts, especially in the context of borderline resectability.

Surgical options

The surgical options entail: i) a realist approach of positive patient selection for radical resection or palliative stent or surgical bypass; ii) an activist approach of regional pancreatectomy plus transplantation; and iii) the nihilist approach of palliative stenting or bypass. The main factors in determining optimal treatment of pancreatic carcinoma is the tumor stage and the patient’s fitness for major surgical resection (Table 4). Surgical resection does not improve survival in patients with locally advanced or metastatic disease (Table 5 and Figure 1).

Resectable disease (Stage 1-11)

Resectable tumors are <4 cm in diameter and confined to the pancreas with no local invasion or metastases. Preoperatively, coagulopa-
thy should be corrected by the parenteral administration of vitamin K. To prevent the development of post-operative renal failure patients should be well hydrated and any electrolyte imbalance rectified. Prophylactic antibiotics are prescribed routinely, and some patients may benefit from preoperative parenteral nutrition. Patients with tumor in head of pancreas may benefit from: i) Whipple's PD which is the only hope for cure in <15%; ii) pylorus preserving pancreaticoduodenectomy; iii) PD with en bloc vascular resection and reconstruction (regional pancreatectomy). Although very few ductal carcinomas of the body and tail are resectable (3%) due to the advanced nature of these tumors at presentation these patients may benefit from distal pancreatectomy or total pancreatectomy for the rare slow-growing malignant tumors, which include cystadenocarcinoma and papillary-cystic neoplasm. Currently, minimally invasive techniques (laparoscopy and robotic techniques) are used for surgical treatment of early forms of pancreatic cancer with the advantages of a high degree of eligibility as well as significant reductions in length of stay, wound infections, and pancreatic fistula. The overall survival after laparoscopic PD is similar to open PD.6,7 Robotic-assisted PD is safely feasible in selected patients and the results in pancreatic cancer are encouraging but deserve further investigation.50

The poor survival rate associated with surgery alone for early-stage PC has led to adjuvant therapy becoming the standard of care after resection in an effort to prolong survival. Neoadjuvant and adjuvant chemotherapy or chemoradiotherapy improves the chance of cure for early PC. CT or magnetic resonance imaging (MRI) is used in the oncological follow-up of patients with resections for pancreatic carcinoma. It is started 6 months after the operation and in 6-month intervals for 2 years postoperatively. Then yearly intervals seem to be sufficient. In addition the serum levels of carcino-embryonic antigen (CEA) and CA 19-9 are also evaluated.3,4,7,8,67

Table 5. Management options.

<table>
<thead>
<tr>
<th>Resectable diseases (Stages I-II)</th>
<th>Locally advanced unresectable disease (Stage III)</th>
<th>Metastatic disease (Stage IV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surgical resection (with or without adjuvant chemotherapy or chemoradiotherapy)</td>
<td>Neoadjuvant chemoradiotherapy then restaging CT with or without resection if down-staged</td>
<td>Palliative chemotherapy (gemcitabine)</td>
</tr>
<tr>
<td></td>
<td>Palliative chemotherapy: 5-FU, folinic acid, and gemcitabine</td>
<td>Palliation of biliary and gastric outlet obstruction</td>
</tr>
<tr>
<td></td>
<td>Palliative chemoradiotherapy</td>
<td>Palliation of biliary and gastric outlet obstruction</td>
</tr>
<tr>
<td></td>
<td>Palliation of biliary and gastric outlet obstruction</td>
<td></td>
</tr>
</tbody>
</table>

CT, chemotherapy.

Figure 1. Summary of survival and resection percentages for patients with pancreatic cancer. Neoadj., neoadjuvant; Tx, treatment; Pall., palliative; Adj., adjuvant. Reproduced with permission from Gillen et al., 2010.63
Locally advanced (unresectable) disease (Stage III)

Of patients 35% with pancreatic cancer present with locally-advanced disease. The management options are: i) neoadjuvant chemoradiotherapy then re-staging CT and resection if down-staged as the best palliation is surgical resection; ii) palliative chemotherapy with 5-FU, folinic acid, and gemcitabine; iii) palliative chemoradiotherapy; and iv) palliation of biliary and gastric outlet obstruction.

Palliative surgery

It is recommended that biliary bypass is combined with gastric bypass (double bypass) in most patients with unresectable cancer of the pancreatic head. The mortality rate of biliary bypass is not increased by adding a gastroenterostomy. Following biliary bypass alone, 17% of patients will develop duodenal invasion and obstruction as the tumor enlarges requiring subsequent gastric bypass which carries an average mortality rate of 22%. However, while surgical bypass is an effective means of relieving biliary obstruction (hepaticojejunostomy) and avoiding duodenal obstruction (gastrojejunostomy), it is not without risk in this elderly ill patient population. The mean operative mortality of bypass in one series was no less than 19% (range 4-30%), and it was clear that much depended on patient selection. Given that the mean duration of survival following bypass is only 5.4 months, non-operative methods of stenting (endoscopic, percutaneous) have been embraced so enthusiastically, almost to the point at which surgical bypass is reserved for patients in whom stenting is technically not possible. Despite the success of stenting and its lower early mortality, stent blockage remains the major drawback with resulting cholangitis and return of jaundice and pruritus. However, while surgical bypass may enjoy a comeback in selected patients, particularly now that biliary-enteric and gastric bypass can be achieved by laparoscopic means with the additional benefit of unequivocal assessment of resectability.

Multivariate analysis of patients with unresectable pancreatic cancer revealed age, stage of disease serum albumin, serum C-reactive protein (CRP) levels as independent predictors of duration of survival. The presence of an acute phase response (C-reactive protein level) was the single most significant independent predictor with a median survival of 66 days with CRP <10 mg/L compared to 222 days in those with no acute phase response. Thus, the importance of cachexia as a cause of early death in pancreatic cancer and the elucidation of the relationship between cytokine profile, acute phase response and resting energy expenditure.

Metastatic disease (stage IV)

Of patients 50% with pancreatic cancer present with metastatic disease and the majority are beyond significant palliation. Management options of metastatic disease include palliative chemotherapy (gemcitabine), best supportive care, pain control with consideration of coeliac plexus block and palliation of biliary and gastric outlet obstruction. For patients with metastatic disease, the primary goals of treatment are palliation and improved survival, yet effective treatments for this population are limited, leading to extremely poor survival rates (5-9 months).

Gemcitabine has been the standard treatment for metastatic disease, primarily due to its effect on symptoms and favorable toxicity profile rather than a significant effect on survival. The combination of gemcitabine with oxaliplatin may confer an additional but small survival benefit in patients with good performance status and a younger age.

Arguments for surgical resection

The arguments for surgical resection are as follows: i) carcinoma of the pancreas is increasing in incidence and the use of modern diagnostic imaging techniques e.g., US, CT, MRI can pick up tumors at an early stage; ii) even though the chances of cure are <10%, the only hope for cure is by surgical resection; iii) if we exclude all patients from consideration for surgery we may exclude patients suffering from carcinoma of distal common bile duct, the duodenal mucosa and ampulla of Vater having a 5 year survival rate of 30%; iv) the operative mortality has fallen to 5% or less in experience hands; v) although it is considered a disease of the elderly more than 40% of men and 35% of women present under the age of 70 years (Figure 2).

Arguments against surgical resection

The arguments against surgical resection are: i) pancreatic carcinoma usually has an insidious presentation and physical signs of metastatic spread are commonly present at initial consultation; ii) it is a disease of elderly patients and 50% are >72 years. Many are unfit, weak, emaciated and suffer from other concomitant medical conditions. Endoscopic bypass is all that can be offered; iii) bypass procedures are all that can be achieved in the vast majority; iv) an unsuccessful resection for a carcinoma can result in a high mortality, a very high morbidity and an extremely costly period of treatment for the patient.

Why persistently poor long-term results after pancreatic resection?

The answer largely lies in the biological nature of pancreatic cancer, which demonstrates aggressive local invasion, and metastases during...
early development. Over 80% of patients have positive regional lymph nodes or distant metastases at the time of diagnosis. Studies have revealed that even in small pancreatic tumors, which have not spread through the pancreatic capsule and with a diameter of less than 2 cm, there are positive para-aortic lymph nodes in 40% and therefore classified as stage II disease. Thus small tumor size cannot automatically be equated with early tumor stage. The biological grade is most important. Pancreatic cancer extends preferentially along lymphatic channels and nerves and readily invades blood vessels. The close interaction of tumor cells and neural structures is mediated by molecular signals including the neural cell adhesion molecules (NCAM) which facilitates invasion and metastases. Nerves have been found to express these factors in abundance, thus providing a positive growth stimulus for pancreatic cancer cells. These may partially explain the early spreading nature and the fact that at least 50% of cases have a local recurrence within the tumor bed following resection. The implication is that by targeting NCAM, cytotoxic drugs may be delivered directly to the pancreatic cancer cells with better efficacy and less systemic side effects than systemic chemotherapy.

**Use of tumor associated antigens in diagnosis and follow-up**

The most widely used marker is CA 19-9 antigen as it is expressed in 86% of pancreatic cancers with a sensitivity of 89% compared to the sensitivity of 37% with CEA. 70% of patients with a tumor <4 cm already show elevated serum levels. Additionally CA 19-9 levels correlate with prognosis as it is more significantly lower in small resectable tumors than in larger ones. However, its sensitivity is not high enough for the primary diagnosis of pancreatic cancer. CA 19-9 is elevated in patients with non-malignant diseases, such as chronic pancreatitis or obstructive jaundice of various origins and in smoking. Its determination has a high clinical value if a CT scan indicate a pancreatic cancer and in the follow-up of patients following resection. If the CA 19-9 level returns to normal after tumor resection and increases during follow-up, then cancer relapse is extremely probable.

**The future**

Progress in identifying new therapies has been hampered by the genetic complexity of the disease with each tumor cell carrying an average of 63 mutations, and the lack of prognostic markers. Most alterations occur with very low frequency and so are challenging to exploit therapeutically. The future lies on the better understanding of the molecular oncology of pancreatic cancer, which entails the genetics and the pathophysiology of metastasis of pancreatic cancer. About 75% of human pancreatic adenocarcinomas have acquired a mutation in codon 12 of the K-ras gene and there could be a role for biological therapy countering the effects of specific mutant oncogenes. Inactivation of tumor-suppression genes occurs frequently during the development of cancers with loss of function of the p53 gene in 60% of primary ductal adenocarcinoma of the pancreas. Replacement of tumor suppressor functions may therefore be a useful therapeutic strategy. Several receptor tyrosine kinases, including epidermal growth factor receptor family (EGF-R, erbB-2, erbB-3) are frequently over-expressed in pancreatic cancer and involved in signal-transduction pathways. Inhibition of signal transduction may play a role in preventing metastases. Although the results of initial trials with EGFR inhibitor (erlotinib) in unselected patients have been disappointing, the agents may be useful in KRAS-mutant p53 wild type tumors which occurs in 15% of patients. New-onset diabetes may also be a potential clue to the early diagnosis of pancreatic cancer. Genetic data have been interpreted to suggest that development of invasive disease from precursor lesions occurs over a considerable length of time (17 years on average), with death following after 2-3 years, highlighting the importance of identifying early diagnostic markers of pre-invasive pancreatic cancer. The recent major break-through is in the identification of early protein markers (cancer exosomes) that may provide early diagnosis and represent a valid screening test. This would lead to early surgical intervention with a better chance of curing this essentially incurable disease. As familial aggregation and genetic susceptibility may play a role in as many as 10% of patients with pancreatic cancer, screening of first-degree relatives of this and other high risk groups for these protein markers would allow early intervention and improve long-term survival.

**Immunotherapy and vaccination**

Immune therapy is changing the current treatment paradigm for malignancy, especially with the recent development of antibodies that can modulate immune checkpoint pathways. Immunotherapy to treat pancreatic cancer is a promising approach due to its low toxicity and potential for creating lifelong immune response. Multiple large phase III trials using simple vaccination strategies have failed to modulate the immune response in pancreatic cancer. However novel strategies with whole cell vaccines using hyperacute rejections (algenpantucel-L) immunotherapy demonstrated 62% and 86% 12-month disease free survival and overall survival in resected pancreatic cancer patients. Combination of whole cell vaccine GVAX and mesothelin-secreting vaccine CRS-207 demonstrated an overall survival benefit in metastatic refractory pancreatic cancer patients. Anti-Gal is the most abundant natural antibody in humans, comprising about 1% of immunoglobulins. The anti-Gal ligand is a carbohydrate antigen called α-gal epitopes with the structure Galc1-3Galβ1-4GlcNAc-R. It is exploited in cancer vaccines to increase the immunogenicity of antigen-presenting cells (APCs). As cancer cells or ductal adenocarcinoma cells (PDAC) tumor lysates are processed to express α-gal epitopes, vaccination with these components results in in vivo opsonization by anti-Gal IgG in PDAC patients. The Fc portion of the vaccine-bound anti-Gal interacts with Fc receptors of APCs, inducing uptake of the vaccine components, transport of the vaccine tumor membranes to draining lymph nodes, and processing and presentation of tumor-associated antigens (TAAs). Cancer vaccines expressing α-gal epitopes elicit strong antibody production against multiple TAAs contained in PDAC cells and induce activation of multiple tumor-specific T cell.

A pancreatic tumor-specific biomarker characterized in humans and mice as an immunogenic onco-glycoprotein is efficient in dendritic cell vaccination. Thus murine DC, loaded with pancreatic tumor-specific glycoepitope C-ter-J28+, induces efficient antitumor adaptive immunity and represents a potential adjuvant therapy for patients afflicted with PDAC.

**Conclusions**

Pancreatic ductal adenocarcinoma is still a disease with a very poor prognosis. It is genetically very complex with a high diversity of mutations compared with other cancers. Early diagnosis with the new protein markers may lead to early intervention and better prognosis. The main surgical goal in performing an R0 resection facilitated by improved staging and patient selection would result in hospital mortality of <5% in specialist centers. As pancreatic carcinoma is largely resistant to standard chemotherapy, consideration of multimodal treatment including immunotherapy is necessary.
References


