Epithelial markers in pancreatic carcinoma: immunoperoxidase localisation of DD9, CEA, EMA and CAM 5-2


Abstract
Paraffin wax embedded, formalin fixed sections of 22 adenocarcinomas of the exocrine pancreas were stained with four mouse monoclonal antibodies: DD9-E7, an antibody raised against a human pancreatic tumour xenograft; carcinoembryonic antigen (CEA); epithelial membrane antigen (EMA); and cyto-keratin (CAM 5-2). An indirect immunoperoxidase technique without enzyme pre-digestion and an affinity-purified sheep anti-mouse peroxidase conjugate were used. All of the tumours were positive for DD9-E7, EMA, and CAM 5-2. Twenty out of 22 were focally positive for CEA and the staining was often weak. As all of these adenocarcinomas were DD9-E7 positive, absence of staining for DD9-E7 in a tumour makes the diagnosis of adenocarcinoma of the exocrine pancreas very unlikely, and this is of value in distinction from endocrine carcinomas with a marked acinar pattern. The weak CEA staining distinguished pancreatic carcinomas from colorectal tumours. Because the distribution of staining for EMA and CAM 5-2 was no different from that previously seen in adenocarcinomas from other sites, these markers are likely to be of limited value in the differential diagnosis of abdominal adenocarcinomas of uncertain origin.

In England and Wales there has been a substantial increase in adenocarcinomas of the exocrine pancreas over the past 75 years. The mortality trebled from an age standardised rate for both sexes of 2.9 per 100 000 in the quinquennium 1911–1915 to 10.6 in men and 8.3 in women in 1971–1975. Since 1974 the mortality has remained fairly stable, with a mean of 2971 a year for men and 2857 for women. There were slightly more deaths from this cause in men than in women until 1986. Since then the mortality in men has fallen somewhat from 2950 in 1986 to 2905 in 1988, and risen in women from 3093 to 3103. Carcinoma of the pancreas is uncommon under the age of 40, and the maximum mortality from this cause is at age 65–79. The prognosis is dismal, with a three year survival of the order of 3% and very few long term survivors. Most patients present late, often with metastatic deposits demonstrable at surgery or laparotomy and a resectability rate of only 12%. The prognosis of cystadenocarcinomas of the pancreas seems to be better than that of solid ductal carcinomas. In a study of 1001 patients, which included all types of exocrine and endocrine pancreatic carcinomas and ampullary carcinomas, the resectability rate of pancreatic cystadenocarcinomas, which occurred in patients with a mean age of 45 compared with a mean age of 67 for solid tumours, was 67% (six of nine). The three year survival was 33% (three of nine), though all three patients finally died of recurrence three, nine, and 16 years later.

The diagnosis of the primary tumour remains difficult for clinicians, while for the histopathologist there are various areas of difficulty. These include (i) determining whether metastatic tumour in nodes, liver, bone, or in other sites could have arisen in the pancreas; (ii) deciding whether an adenocarcinoma in the lung could represent a metastasis from an occult pancreatic carcinoma and therefore be unsuitable for resection; and (iii) distinguishing chronic pancreatitis from well differentiated adenocarcinoma, especially in needle biopsy specimens.

The pronounced desmoplastic response evoked by many pancreatic adenocarcinomas may be of some help in the distinction from other tumours, and many are well differentiated with a rather angular glandular pattern, rather unlike the smoother more ring-like glands seen in gastric or colorectal carcinomas. They are not, however, sufficiently different morphologically from those arising elsewhere in the gastrointestinal tract for a confident diagnosis of a metastatic deposit. In biopsy material origin from other sites cannot be excluded unless there is both a mass in the pancreas felt by the surgeon and pancreatic tissue in the biopsy specimen.

In an initial study DD9-E7 supernatant, raised by immunisation of nude mouse hairy lintermates (nu+) with a homogenised xenograft of GEF pancreatic adenocarcinoma cells, was used. It had been shown to react strongly with 12 of 14 adenocarcinomas of the exocrine pancreas and with a variety of other normal and neoplastic tissues. Staining of polymorphs and macrophages suggested that the antigen against which DD9-E7 was directed was similar to normal crossreacting antigen, NCA. Western blot analysis has shown that DD9 recognises a protein epitope on a family of glycoproteins (80–115 kilodaltons) which are distinct from the sialoglycoside and mucin-like pancreatic tumour antigens recognised by other monoclonal
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antibodies CA 19–9, Du Pan-2, or Ca-50. The epitope to which the antibody binds is not found in CEA or NCA-2, but is present in NCA-1, suggesting that NCA-1 and the DD9 antigen share a common determinant (AG Grant, unpublished data).

Methods
Twenty two surgically resected or biopsied unequivocal primary pancreatic ductal adenocarcinomas were selected from the surgical files of St Thomas’s or St George’s Hospitals, including the adenocarcinoma from which the GER cell line had been derived. Blocks from needle biopsy specimens taken at laparotomy and showing tumour infiltrating fat or fibrous tissue without evidence of pancreatic invasion were excluded.

DD9-E7 used in this study was an ascites preparation produced in beige mice. The anti-CEA was also an ascites preparation (Amersham International plc, Buckinghamshire). It does not stain polymorphs or macrophages and does not seem to recognise an NCA determinant. Anti-EMA was a hybridoma supernatant raised using a preparaton of human milk fat globule membranes (Dakopatts Ltd, Buckinghamshire), and CAM 5-2 was also a supernatant, raised by immunisation with the human colon carcinoma cell line HT29.

An indirect immunoperoxidase technique, without enzyme predigestion, and an affinity purified sheep anti-mouse peroxidase conjugate prepared by periodate oxidation were used. The method was similar to that already published, except that incubation with the first antibody was carried out overnight instead of for one hour.

The positive controls used in this study were a moderately differentiated colorectal carcinoma for CEA and CAM 5-2, a ductal carcinoma of the breast for EMA, and a metastatic deposit in the liver of a pancreatic adenocarcinoma for DD9-E7. A monoclonal antibody directed against prostatic acid phosphatase, PASE/4LJ, was used as an irrelevant negative control. This antibody stains some pancreatic islet cells in formalin fixed tissue, but not benign or malignant exocrine pancreatic tissue. Prostatic chippings showing benign hyperplasia were used as a positive control for PASE/4LJ.

Results
All of the tumours were positive for DD9-E7, EMA, and cytokeratin, and 20 of 22 were positive for CEA (figs 1–6). Most of the residual pancreatic islets were negative with all four antibodies. A few cells in some islets, however, were positive for cytokeratin. Some positive staining for DD9-E7, EMA, and cytokeratin was found in normal pancreas, as well as in the areas of chronic pancreatitis so frequently found surrounding pancreatic exocrine carcinomas. Although normal pancreatic tissue was negative for CEA, there was positive staining in foci of chronic pancreatitis.

The distribution of staining for DD9 was on the luminal membrane of malignant acini and in the luminal contents (fig 1). Positivity was also often seen as a cytoplasmic globule above the nucleus, separated from the luminal membrane, and corresponding to the position of mucigen granules (fig 2). The two tumours negative with the supernatant used in the previous study were positive in this study when ascitic fluid was used. This was likely to be due either to the increase in immunoglobulin content or to prolongation of the first antibody incubation time.

Though some of the tumours were strongly positive for CEA (fig 3), several were weakly or focally positive. Staining was on the luminal membrane or in the cytoplasm, or both, but the globular staining seen with DD9-E7 (fig 2) was not seen with CEA. The normal pancreas surrounding the tumour was
negative for CEA, but there were areas of positivity in foci of chronic pancreatitis.

In some tumours CAM 5-2 was localised just under the plasma membrane of the tumour cells, in a “pericellular pattern” (fig 4). In others it was linear and just under the luminal membrane; in the rest it was cytoplasmic. In some tumours the staining was weak and patchy, but in most it was strong. Some residual pancreatic islets showed focal areas of positivity.

The staining for EMA was mainly on the luminal membrane (fig 5), but there was also some cytoplasmic staining (fig 6). In the residual exocrine pancreas intercellular canaliculi similar to those seen in eccrine sweat glands were positive.

**Discussion**

The presence of DD9 in all 22 primary pancreatic adenocarcinomas, and its absence in up to 50% of other adenocarcinomas, suggests that its presence in a metastatic tumour, especially with a weak or only focally positive CEA, makes the tumour more likely to be of pancreatic than of gastric or of colorectal origin. Conversely, a negative stain for DD9 makes the exocrine pancreas a highly unlikely primary site. This has already proved useful clinically in distinguishing endocrine pancreatic carcinomas with pronounced acinar differentiation from exocrine carcinomas with their very much worse prognosis. Bronchial tumours may also be negative for CEA and some contain DD9 (E Heyderman, unpublished data), so DD9 staining does not distinguish primary adenocarcinomas of the lung from metastatic pancreatic tumours. As far as other sites are concerned, prostatic deposits can virtually be excluded by the absence of stainable prostatic acid phosphatase, and most thyroid carcinomas are positive for thyroglobulin.

The presence of endogenous peroxidase activity in leucocytes, especially eosinophils, makes it essential to inhibit this activity adequately when studying the distribution of staining with antibodies to NCA related epitopes. Positive staining for polymorphs can make interpretation difficult in foci heavily infiltrated by acute inflammatory cells. This was not a major problem in the pancreatic tumours studied here, but we have found staining for DD9 in squamous carcinomas of the lungs difficult to assess for this reason (E Heyderman, unpublished data). DD9 immunostaining, however, could be of value in determining whether cells in the bone marrow in proliferative states or in leukaemia are of myeloid origin, and this antibody, like others recognising an NCA epitope, could be of value in immunoscintigraphy of occult abscesses.

Tsutsumi et al used a rabbit antibody specifically raised against NCA. They showed the presence of NCA in cryostat sections of pancreatic adenocarcinomas, but not in formalin-fixed, paraffin wax embedded blocks from the same tissues. All of their tumours, however, were positive with the Dako anti-CEA before absorption with a spleen extract. The Dako antibody is well recognised to have NCA activity, and pre-absorption with an NCA preparation is required before it can be used to show CEA and not NCA. Presumably, the rabbit NCA antibody used by Tsutsumi et al recognises a different, more labile NCA epitope. In another study monoclonal and polyclonal NCA antibodies stained most pancreatic carcinomas and foci of chronic pancreatitis. The monoclonal antibody 374 which recognises a shared CEA, NCA 95, and NCA 55 determinant, stained normal ductal cells and 36 of 40 pancreatic carcinomas. The monoclonal antibody 47D10, which was positive in 95% of pancreatic adenocarcinomas but not in foci of chronic pancreatitis, also stains granulocytes, and may recognise an NCA-related antigen. This antibody, however, is different from DD9 in that it recognises a carbohydrate epitope.
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There have been several studies on the localisation of CEA in pancreatic carcinomas. In the two studies undertaken in 1972 and 1978, only three out of six and two out of seven of the pancreatic carcinomas were positive, but neither of the antisera used showed positivity in normal colonic mucosa, so that sensitivity was low. More recent studies have shown results very similar to ours, with all or nearly all pancreatic adenocarcinomas showing some CEA.

It has been noted in some studies that staining for CEA was stronger in better differentiated pancreatic tumours but this was not notable in our series, nor in those of others. The weak staining was not always in close association with the tumour epithelial elements. However, in some cases, strong positivity was observed only in a subpopulation of epithelial cells. It is interesting to note that CEA staining was not observed in normal pancreatic acinar cells, and that the staining was restricted to the cytoplasm of the tumour cells. The absence of CEA staining in normal pancreas led us to hope that its demonstration could be of value in the differentiation of tumour from chronic pancreatitis, but, as found by others, both adenocarcinomas and foci of chronic pancreatitis were positive, though such foci were found to be negative with the F6 CEA antibody.

EMA and cytokeratin, while of possible value for tumour immunolocalisation in patients where specificity may not be required, do not distinguish pancreatic adenocarcinomas from those arising elsewhere.

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The photomicrographs were taken using Ektor Gold 25.

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Figure 5  The EMA stain is mainly luminal with the contents of the malignant acini also positive.

Figure 6  In this tumour there is a patchy cytoplasmic positivity for EMA.


