Neurofibromatosis and small bowel adenocarcinoma: an unrecognised association

T J Jones and T L Marshall

From the Department of Morbid Anatomy, The London Hospital Medical College, London, and the Department of Histopathology, Central Pathology Laboratory, Stoke-on-Trent, Staffordshire

SUMMARY We present a review of the reported cases and describe a fifth of neurofibromatosis with small bowel adenocarcinoma and concurrent leiomyomas, indicating a previously unrecognised association.

Neurofibromatosis is known to be associated with an increased incidence of neoplasms including phaeochromocytoma glomata and neurofibrosarcoma.1 Gastrointestinal involvement is well documented, predominantly consisting of neurofibromas, ganglioneuromas and schwannomas.1,2 Epithelial tumours including colonic,3 pancreatic4 and small intestinal adenocarcinomas have all been reported.5,6

Case report

The patient, a 59 year old white man initially presented in July 1980 with a soft, well circumscribed mass 1-5 cm in diameter on the right thigh. This was excised and histology showed it to consist of spindle cells in a fibrillary stroma, appearances typical of a neurofibroma. On further clinical examination he was found to have a café-au-lait area on the left thigh; his father had been diagnosed as having neurofibromatosis in the past (no biopsy available). He was therefore diagnosed as a case of neurofibromatosis and remained asymptomatic for the next five years.

He next presented in February 1985 with intermittent colicky abdominal pain associated with vomiting, diarrhoea and 10 kg weight loss. On examination his abdomen was distended, with increased bowel sounds and tenderness over the right iliac fossa. Initial laboratory investigations showed a neutrophil leucocytosis (WCC 19-6×10 with 80% neutrophils). A diagnosis of small bowel obstruction was made and laparotomy revealed an area of stenosis in the middle third of the ileum with multiple serosal nodules. Thirty centimetres of ileum were resected and submitted for histological examination. On opening the bowel an ulcerated stenosing lesion was noted 15 cm from the distal plane of excision along with multiple nodules on the serosal surface situated principally on the mesenteric border.

Microscopy of the stenosing lesion showed it to be a moderately differentiated adenocarcinoma (Fig. 1) which had penetrated through to the serosal surface with metastases in the mesenteric lymph nodes (Fig. 2). The serosal nodules were shown to consist of spindle cells with cigar shaped nuclei arranged in interlacing bundles (Fig. 3); they arose from the muscularis propria and extended into the submucosa and onto the serosal surface. No neural elements were seen. Tissue from the nodules was placed in glutaraldehyde, postfixed in osmium tetroxide and examined using an AEI Corinith 500 Electron microscope. This showed intracellular filaments, adherent basement membrane like material and pinocytotic vacuoles. The overall appearances were of tumours of smooth muscle origin.

Discussion

Neurofibromatosis with involvement of the gastrointestinal tract is well documented1,2 and usually consists of neurofibromas, ganglioneuromas, and leiomyomas; much rarer is the development of
Fig. 1  Section showing infiltrating adenocarcinoma. H&E. ×50

Fig. 2  Metastatic adenocarcinoma in subcapsular region of mesenteric lymph node. H&E. ×50
epithelial malignancy. Cases of adenocarcinoma of the colon and pancreas with neurofibromatosis have been described. In addition there have been four case reports of small bowel adenocarcinoma. In three of these the neurofibromatosis was of the peripheral type; in the remaining case the neural tumours were limited to the ileum and so were of the visceral type.

The first case was reported in 1974 in a 48 year old man who presented with small bowel obstruction and was found to have adenocarcinoma of the ileum with multiple nodules protruding from the mucosal surface. Microscopy showed these to consist of plexiform proliferation of neurites and Schwann cells interpreted as being plexiform neurofibromas. Multiple lymph node metastases were present. Cutaneous evidence of neurofibromatosis was not seen. The patient died in the immediate postoperative period.

The second case was reported in 1980 in a 75 year old man who presented with anorexia, weight loss, and cutaneous neurofibromatosis. A large ulcerated, fungating tumour was present in the second part of the duodenum in the region of the ampulla. Biopsy of the tumour showed it to be a typical villous peripanillary adenocarcinoma interpreted as arising from the duodenal mucosa. A laparotomy was carried out which showed multiple nodules on the serosa of the jejunum and ileum which were interpreted as being metastatic deposits; a biopsy procedure was done and the serosal nodules biopsied. Microscopy showed these to consist of spindle cell tumours interpreted as being leiomyomas. The patient died on the third day postoperatively.

The third case was reported in 1982 in a 72 year old woman with cutaneous neurofibromatosis and weight loss. Small bowel radiography showed a polypoid mass in the ileum which was resected and shown to be a moderately differentiated adenocarcinoma admixed with a ganglioneuroma. Lymph node metastases were present. The patient was discharged one month after surgery.

The fourth case was also reported in 1982 in a 29 year old man with jaundice, hepatomegaly and cutaneous neurofibromatosis. Laparotomy revealed a mass in the ampulla of Vater and a Whipples pancreaticoduodenectomy was carried out. Microscopy showed the mass to be a well differentiated adenocarcinoma with dysplastic changes in the surrounding duodenal mucosa and in the terminal common bile duct. Four lymph nodes contained metastatic carcinoma. The patient died four months postoperatively.

The above cases together with the one which is reported here indicate a previously unrecognised association between neurofibromatosis and adenocarcinoma of the small bowel including the peripanillary region. Three of the reported cases also had intestinal stromal tumours: one with multiple leiomyomas, one ganglioneuroma, and one plexiform neurofibroma. In the latter this was the only manifestation of neurofibromatosis. The present case had multiple ileal leiomyomas. All the resected cases including the present had local lymph node metastases at resection, indicating the usual advanced stage of small bowel adenocarcinoma at presentation. We believe that the association is unlikely to be caused by chance because of the very low incidence of small intestinal adenocarcinoma (0.5/100000 population) and forming less than 1% of all intestinal neoplasms: the possibility of the two conditions occurring together spontaneously is therefore very slight.

References


