Acute GI bleeding by multiple jejunal gastrointestinal autonomic nerve tumour associated with neurofibromatosis type I

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INTRODUCTION

Neurofibromatosis type I is an autosomal dominant disorder with variable penetrance. The major diagnostic criteria include skin lesions such as café-au-lait spots, axillary and inguinal frecklings, multiple neurofibromas, Lisch nodules (pigmented iris hamartomas), optical gliomas and bone lesions and a first degree family history of NF (1). The responsible gene for NF1 codes for a GTPase-activating protein (neurofibromin), which regulates the activity of the protooncogene p21 ras. Therefore neurofibromatosis patients are at an increased risk to develop malignant tumours (2).

Gastrointestinal (GI) involvement in von Recklinghausen’s disease occurs in three principal forms: hyperplasia of the submucosal and myenteric nerve plexuses and mucosal ganglioneuromatosis lead to disordered gut motility, gastrointestinal stromal tumours (GISTs) showing varying degrees of neuronal or smooth muscle differentiation and carcinoids of the periampullary region of
the duodenum (3). The identification of gastrointestinal stromal tumours which arise from autonomic nerves in the bowel wall (gastrointestinal autonomic nerve tumours, GANT’s), originally described as plexosarcomas, is based on ultrastructural similarities with the myenteric plexus of the gastrointestinal tract (3,4). These tumours represent a distinct subcategory of gastrointestinal stromal tumours. There is also increasing evidence in the literature for the association of von Recklinhausen’s disease and GANT’s of the small bowel (5,6). Here we describe a surgical emergency due to GI-bleeding caused by GANT’s in a patient with von Recklinghausen’s disease.

**CASE REPORT**

A 72-year old female patient presented with a history of melena. Von Recklingshausen’s disease was diagnosed 30 years ago. Physical examination showed a clearly reduced nutritional status and the typical café au lait spots as well as multiple skin neurofibromas. During physical assessment blood was found upon digital examination of the rectum. To exclude a bleeding source in the stomach or duodenum a gastroscopy was performed which showed no abnormalities. At colonoscopy blood was found in the terminal ileum, however, the active bleeding source could not be identified. Because GI bleeding persisted, two units of whole banked blood were transfused over the first two days. A dual-phase angio-CT-scan (Somatom Volume Zoom, Siemens, Erlangen, Germany) after i.v. application of 120 ml iodinized contrast media was performed, which localized a bleeding focus in extraluminal tumours of the proximal jejunum (Fig. 1).

After laparotomy, multiple extraluminal tumours were found on the serosal side of the proximal jejunum (Fig. 2). The proximal resection margin was chosen directly after Treitz ligament to allow a 2 cm distance to the most proximal tumour and a 20 cm long segment was resected. On the opened specimen, one larger tumour had eroded into the mucosal surface. Bowel continuity was restored by means of a retrocolical latero-terminal duodeno-jejunos- tomie.

Pathological examination of the resected jejunum showed multiple small tumour nodules on the serosa in close proximity to several larger maximally 2.5 cm tumours. These larger tumours infiltrated all layers of the bowel wall and some eroded the mucosa. Tumours consisted of uniform spindle-formed cells with an eosinophilic cytoplasm (Fig. 3A). Locally a hyperplasia of the plexus myentericus could be found. Smaller tumours in the serosa and subserosa showed a clearly positive staining for CD 34, NSE and S100 (Fig. 3B). No staining could be found for neurofilaments and SMA. Larger tumours with infiltration of the mucosa also stained strongly positive for C-Kit.
These are typical features of gastrointestinal stromal tumors of autonomous nerve origin. The postoperative period was uneventful. The patient was advised to attend a regular follow up. In the three year follow up, no recurrence could be detected.

DISCUSSION

Neurofibromatosis is one of the most frequent hereditary diseases. While also several rare mosaic forms are described neurofibromatosis 1 is the commonest form with up to 20 times as frequent as NF2 (7). Neurofibromatosis 1 and 2 are autosomal dominant diseases with half of the cases being attributed to new mutations. There is an increased risk to develop malignant tumours in NF1. The responsible gene which has been localized to chromosome 17 (gene locus 17q11.2) codes for a GTP-ase which is expressed in all cell types at low amounts, in high amounts it can be found in neuroectodermal cells (8,9). Physiologically, neurofibromin 1 binds with the gap120-like to p21 ras which leads to inactivation and consequentially to cellular differentiation and to an inhibition of cell growth. This inhibitory function is lost with a mutated and defective neurofibromin gene product (9).

The identification of GANT’s, originally described under the name plexosarcoma has been based on ultrastructural similarities with the myenteric plexus of the gastrointestinal tract (10). Ultrastructurally, the presence of dense core granules, cell processes, neurotubules, dystrophic axons, synapse-like structures and skenoid fibers are described to be characteristic for these tumours (11,12). Gain of function c-KIT-mutations were found in extracellular, juxamembrane and kinase domains presumably contributing to the molecular initiation of these neoplasms (5).

GANT’s normally tend to be semimalignant. Nevertheless complete en bloc surgical resection, when possible, is the cornerstone of their therapy (13). A more radical resection in the present case would have been a pylorus preserving pancreaticoduodenectomy since also metastatic disease and tumour recurrence have been described in patients with GANT’s. In our patient, however, the tumours had manifested themselves by severe GI-bleeding and therefore a less radical approach was pursued, especially since liver metastases or intraperitoneal spread are a rare phenomenon and tend to occur mainly in patients with tumours that have ruptured spontaneously or which have been violated by the surgeon.

Gastrointestinal tumour manifestation in neurofibromatosis has been described and typically varies from gut neuronal tissue hyperplasia to gastrointestinal stromal and endocrine tumours. Also other neoplasms can be found (14) but this is generally considered a rare event. GANT’s are a rare but distinct subcategory of gastrointestinal stromal tumours. In the majority of the cases which have been described so far, the primary site of the tumour was the stomach, duodenum, jejunum or ileum, but also oesophagus, rectum, bladder or colon have been reported (6). Stromal tumours remain mainly asymptomatic until the tumour reach a sufficient size to produce abdominal pain or mucosal ulceration with GI-bleeding as reported in our case. At this point, the clinical symptoms depend on the site of the manifestation. Symptoms could also be related to the luminal obstruction but this is a late event due to the serosal origin of the tumour (15). GI-bleeding as a consequence of erosion of the mucosa by GIST has so far only been reported in one patient with neurofibromatosis (16). This report should alert surgeons and gastroenterologists to the rare GI-tract manifestations of tumours in von Recklingshausen’s disease. In as much as GANT’s have to be treated by surgery and especially in the case of GI-bleeding as reported here, we propose a dual phase angio-CT as first cornerstone of the diagnostic of neurofibromatosis patients with any gastrointestinal symptoms.

REFERENCES


