CASE REPORT

Malignant potential in a Brunner’s gland hamartoma

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Brunner’s gland hamartomas are rare tumours of the duodenum. These lesions have previously been described as being benign, with no malignant potential. A case report is presented of a Brunner’s gland hamartoma, whose histology revealed a focus of well marked epithelial dysplasia. This case suggests a dysplastic stage in the natural history of Brunner’s gland hamartoma, and questions the malignant potential of these lesions.

Brunner’s gland hamartomas are rare duodenal tumours and an unusual cause of gastrointestinal bleeding. Since the first description of a Brunner’s gland hamartoma in 1876 there has been controversy to the aetiology and pathogenesis of this lesion. Several case reports have described these lesions as non-dysplastic and entirely benign. However a case report is presented, representing a Brunner’s gland hamartoma with a dysplastic focus. This case report illustrates that these lesions do appear to become dysplastic and raises the question of malignant potential.

CASE REPORT

A 79 year old white man presented with a collapse and melaena. He had a history of stable angina for three years, mild chronic obstructive pulmonary disease, and a previous transient ischaemic attack. He was taking 75 mg of aspirin daily and took minimal alcohol. Examination revealed a soft non-tender abdomen and a blood pressure of 114/75 mm Hg, with no significant postural drop. Haematological investigations revealed anaemia, with a haemoglobin concentration of 92 g/l. On the second day the haemoglobin dropped to 72 g/l and an upper gastrointestinal endoscopy was performed.

At endoscopy a 2 cm pedunculated lesion, with an ulcerated crater, was noted in the second part of the duodenum (fig 1). There were no signs of active bleeding and the lesion was biopsied at this stage. The patient was transfused and subsequently maintained his haemoglobin.

Histology revealed a submucosal lesion with complex, crowded architecture and features consistent with a Brunner’s gland hamartoma. However, in the deeper layers of the specimen, some nuclear atypia with occasional mitotic figures was seen. Given these findings a repeat endoscopy was performed and the lesion was removed by snare polypectomy using coagulation current.

Histology of the resected specimen revealed it to predominantly composed of lobules of relatively bland appearing Brunner’s glands. However throughout the specimen there were foci of obvious dysplasia with some foci showing more pronounced cytological changes (fig 2).

DISCUSSION

Duodenal tumours, both malignant and benign, are rare. Brunner’s gland hamartomas and adenomas account for up to 1% of small bowel tumours. Normal Brunner’s glands are alkaline mucus-secreting and are found predominantly in the duodenum, extending to the proximal jejunum. Brunner’s gland hamartomas are most often found in the proximal duodenum (57%), representing the normal Brunner’s gland distribution. The majority are pedunculated (88%), and are commonly 1–2 cm in diameter. Clinical presentation is variable, and indeed many are asymptomatic and incidental findings. A recent series revealed the commonest presentations, in symptomatic patients, were gastrointestinal haemorrhage (37%) and obstructive symptoms (37%). Most symptomatic patients present in the fifth or sixth decade and there is equal sex distribution. Rarer presentations have included recurrent pancreatitis, obstructive jaundice, and biliary fistula. The diagnosis is usually made by radiographic and endoscopic techniques, and in the series by Levine et al, 92% were identified by small bowel studies. At endoscopy the classical findings are of a pedunculated polyp in the duodenal

Figure 1  Endoscopic view of pedunculated Brunner’s gland hamartoma in the second part of the duodenum.

Figure 2  Photomicrograph of resected specimen showing well marked epithelial dysplasia (haematoxylin and eosin, original magnification x 200).
bulb, though 11% can be sessile. Histology classically reveals normal overlying mucosa and a well circumscribed lesion within the submucosa. This tends to contain normal Brunner’s glands with glandular lobulation with intervening bands of paucicellular fibrous tissue. The Brunner’s glands have small, oval, basally orientated nuclei and apical mucinuos cytoplasm, staining positive with periodic acid Schiff. Series and case reports, in the literature, comment on the considerable variation in nomenclature from Brunner’s gland hyperplasia to adenoma to hamartoma. From reviewing the published literature it is clear that no previously examined Brunner’s case reports or series comment on the entirely benign nature of these lesions and many suggest conservative management in asymptomatic patients.

In conclusion, this case illustrates the potential for dysplasia in these lesions, previously felt to be entirely benign, and raises the question of potential malignant transformation. Indeed there appears to be a continuum between Brunner’s gland hyperplasia, hamartoma and adenoma, with a potential for dysplastic changes. Clearly, from the benign nature of Brunner’s gland hamartoma in all the previous case reports, these dysplastic changes appear to be a rare phenomenon. However given the findings in this case it would be dangerous to presume that all Brunner’s gland hamartomas are non-dysplastic and non-malignant.

References

Images in Medicine

A gangrenous foot

The incidence of Kaposi’s sarcoma in HIV infected individuals has declined since the advent of combination antiretroviral therapy in the mid-1990s. The patient, a 34 year old white homosexual man, presented with a swollen, ulcerated, and infected left foot. Amputation was considered but then an HIV antibody test was performed and this was reported positive with severe immunosuppression (CD4 cell count 20 mm⁻³) and plasma HIV viral load >750 000 copies/ml. A biopsy specimen from the foot lesions confirmed Kaposi’s sarcoma. There were also some small lesions suggestive of Kaposi’s sarcoma in his mouth and right conjunctiva along with widespread psoriasis. There was dramatic improvement after six weeks of antibiotics, a combination of three antiretroviral agents (stavudine, lamivudine, and efavirenz), localised radiotherapy, and treatment for psoriasis. This coincided with a rise in CD4 count and rapid drop in HIV viral load. Six months later, he was wearing a normal shoe and the foot was back to normal.

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