Angiomyolipomas are rare lesions, often arising in the kidney, and are part of a group of tumours with a diverse appearance and evidence of dual melanocytic and smooth muscle differentiation known as PEComas (tumours of perivascular epithelioid cell origin). This report describes an unusual case of a colonic PEComa in a 40 year old woman. Unlike most of the previous colonic angiomyolipomas/PEComas reported in the literature, this case formed a large, mainly extrinsic mass and was monotypic, and composed entirely of the myomatous component with no adipose tissue or typical vasculature.

Angiomyolipoma is an uncommon, usually benign, renal tumour showing a characteristic histological triad of adipose tissue, vessels, and modified smooth muscle with immunohistochemical and ultrastructural evidence of dual melanocytic and smooth muscle differentiation. The tumour may be found sporadically or in association with tuberous sclerosis, an autosomal dominantly inherited disorder, characterised by mental retardation, epilepsy, and adenoma sebaceum. Recently, similar tumours at other sites have been shown to have a similar origin.

We present the case of a 40 year old woman with a PEComa (tumour of perivascular epithelioid cell (PEC) origin) of the colon without evidence of tuberous sclerosis. Unlike the previous colonic PEComas/angiomyolipomas reported, this case formed a large, mainly extrinsic mass and was monotypic, composed entirely of the myomatous component with no adipose tissue or typical vasculature.

CASE REPORT

A 40 year old woman was admitted to University Hospital, Nottingham, UK with a history of abdominal bloating and constipation for one year. Her family history and past medical history were unremarkable. Colonoscopy revealed a large extraluminal mass in the medial wall of the caecum. There were no mucosal abnormalities. At laparotomy a large mobile mass arising from the medial caecal wall was removed.

The right hemicolecctomy specimen contained a mass arising in the caecum measuring 80 × 60 × 90 mm. This was predominantly extrinsic but formed a polypoid protrusion into the caecum. The tumour had a solid, haemorrhagic cut surface with areas of necrosis. It had a lobulated outline mainly of classic histological type. Only one previous case,9 reported cases were small (1–3 cm) polypoid tumours, to make when it arises outside its typical renal location because of its rarity and the histological variation.

Angiomyolipoma/PEComa may be a challenging diagnosis to make when it arises outside its typical renal location because of its rarity and the histological variation.

Angiomyolipoma/PEComa may be a challenging diagnosis to make when it arises outside its typical renal location because of its rarity and the histological variation. Typically, angiomyolipoma shows a mixture of adipose tissue, fat, and abnormal vessels, but it may show a wide range of histological appearances. These include tumours where the muscle component is dominant, including epithelioid variants and those showing considerable nuclear atypia. Gastrointestinal involvement is rare, with only seven previous cases reported in the colon and one each in the duodenum, stomach, and appendix. The previously reported cases were small (1–3 cm) polypoid tumours, mainly of classic histological type. Only one previous case, a 3 cm lesion in the caecum, was entirely epithelioid.

Abbreviations: PEC, perivascular epithelioid cell
myomatous but without necrosis. Another 1 cm lesion in the appendix showed an epithelioid and typical vascular component without fat. Thus, our present case is the second reported case in the gastrointestinal tract to consist entirely of the myomatous component, and is also the largest example reported in the gastrointestinal tract.

The combination of a rare tumour at an unusual site with an atypical histological appearance presents diagnostic challenges. Our differential diagnosis included gastrointestinal stromal tumour, leiomyosarcoma, melanoma, and extrarenal rhabdoid tumour. Although gastrointestinal stromal tumour may contain epithelioid areas they are rarely as pleomorphic as in this case. The weak staining with CD117 and the strong positivity for HMB45 precluded this diagnosis. Epithelioid leiomyosarcoma was probably the most important differential diagnosis to consider. Like this tumour, it can present a combination of spindled and epithelioid morphology and may be very pleomorphic. HMB45 positivity has been described in the clear cell component of uterine epithelioid leiomyosarcomas, although these rare tumours may also represent part of the PEComa spectrum. In our case, HMB45 was strongly positive throughout the tumour in spindled and epithelioid areas; this has not been described in leiomyosarcoma. Melanoma was excluded because of lack of staining for S100, whereas rhabdoid tumour was unlikely given the lack of positivity for keratins and minimal positivity for desmin in...
the epithelioid area, HMB45 positivity, and absence of typical ultrastructural features.

Although classic renal angiomyolipomas were regarded as universally benign lesions it is becoming increasingly clear that PEComas should be regarded as tumours of uncertain malignant potential.11–13 Tumours with a poor outcome are usually monotypic myomatous variants, especially those with dominant pleomorphic epithelioid components. Other factors likely to portend poor outcome include presence of haemorrhage and necrosis, local invasiveness, and high mitotic activity. The case presented here had a monotypic appearance with a prominence of epithelioid features, areas of haemorrhage and necrosis, and mucosal invasion, all features that are regarded as potentially malignant. However, to date, the patient remains well with no evidence of recurrence or metastasis.

In summary, we have presented a case of colonic PEComa, which we believe to be the largest so far reported in the gastrointestinal tract and the first to show morphological evidence to suggest potential malignant behaviour.

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REFERENCES