Proliferating Brenner tumour

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Summary
A new case of proliferating Brenner tumour is added to the 18 previously reported cases, and the literature relating to this particular kind of tumour is reviewed.

Introduction
The following case of a proliferating Brenner tumour represents the nineteenth report in the literature (Miles and Norris, 1972; Roth and Sternberg, 1971; Chang, Roberts and Homesley, 1977; Hallgrimsson and Scully, 1972). It is important to recognize this intermediate proliferative variety, as it may easily be confused with a malignant Brenner tumour.

Case report
A 62-year-old white female, nulligravida, 10 years post-menopause, was admitted to hospital because of a low abdominal mass discovered upon routine gynaecological examination. Physical examination on admission was unremarkable except for the presence of a movable mass located in the left lower abdomen. At exploratory laparotomy, a left ovarian tumour was discovered and total abdominal hysterectomy with bilateral salpingo-oophorectomy was performed. The postoperative course was satisfactory and the patient was discharged on the 7th day.

Pathological findings
The left ovary was replaced by an ovoid mass that measured 11 × 7 × 4 cm. The external surface was bosselated, grey-white and glistening. The fallopian tube was attached to the mass in one area. The bisected specimen was partially cystic and partially solid (Fig. 1). The cystic area was multilocular with large and small cystic spaces containing watery to slightly mucoid fluid. The largest cyst measured 4 cm in diameter, thickness of wall 0·1 cm. The solid portion showed 2 distinct areas: at the periphery a hard, whitish-yellow, gritty homogeneous tissue and a large soft papillary mass growing into the cystic spaces (Fig. 2). The right ovary and both fallopian tubes were normal. Microscopically the peripheral solid area was typical of the Brenner tumour and consisted of islands of squamous transitional epithelial cells with varying density of intervening stroma and many calcifications disposed in the epithelial nests and in the stroma (Fig. 3). In the soft papillary intracytic growth the Brenner epithelium assumed a papillary appearance and consisted of 10–20 layers of epithelial cells, which generally maintained their uniform appearance (Fig. 4). There was focal atypism consisting of hyperchromatic cells, but none of the epithelial areas had cytological characteristics of malignancy and no stromal invasion was seen. The lesion resembled a low-grade papillary transitional cell carcinoma of the urinary bladder. Throughout the tumour was a mucinous transformation of epithelial cells, resulting in small to larger cavities of varying sizes, containing a mucicarmine and periodic acid Schiff-positive material. Elsewhere the cyst lining was smooth and composed of mucinous epithelium and partially of transitional epithelium.

Discussion
Meyer (1932) introduced the term Brenner tumour for the ovarian neoplasm, which was first described by Brenner (1907). Since its first description this tumour has been the subject of numerous papers attempting to establish its histogenesis and endocrine activity. The structure of these tumours is characterized by the presence of epithelial cell foci or columns distributed within a prominent fibrous stroma. These cells show remarkable uniformity and tendency for mucinous transformation. Although
characteristically benign, about 42 malignant Brenner tumours have been reported (Miles and Norris, 1972).

Roth and Sternberg (1971) described an unusual form of Brenner tumour – an intermediate neoplasm between the usual benign Brenner tumour and its frankly malignant counterpart, and called it 'proliferating Brenner tumour'. The proliferating Brenner tumour is characterized by an unusual degree of proliferation of Brenner epithelium resembling low-grade papillary transitional cell carcinoma of the urinary bladder.

Lack of cellular atypism, mitosis and stromal invasion are the criteria which differentiate this entity from the malignant counterpart (Roth and Sternberg, 1971). Generalization concerning this variety of Brenner tumour is difficult, because of the small number of reported cases. They are generally
large, mixed cystic and solid tumours. The characteristic macroscopic feature is the presence of soft papillary masses growing into cystic spaces, as is well illustrated in the present case. The presence of Sudanophile (or lipid) granules in the stromal and epithelial cells has been considered as histological evidence of the hormonal activity of Brenner tumours (Ming and Goldmann, 1962).

Proliferating Brenner tumour may cause postmenopausal bleeding owing to endometrial hyperplasia, but the presenting symptom may be an abdominal mass without other clinical complaints. In the present case, lipid stains were negative and there was no clinical or pathological support for endocrine activity. There is still no complete agreement as to the histogenesis of Brenner tumour. The appearance of the proliferating Brenner tumour resembling low-grade papillary transitional cell carcinoma of the urinary bladder, gives additional support to the concept that it originates from the coelomic epithelium, which is capable of transitional (urothelial) and mucinous (uterino-cervical) differentiation, both of which are also characteristics of Brenner tumours. Proliferating Brenner tumour is biologically benign and the resection of the ovary containing the tumour is the treatment of choice.

References
CASE REPORTS


