Pseudomyxoma peritonei (PMP) is classified into pathologically and prognostically distinct categories, such as disseminated peritoneal adenomucinosis (DPAM) and peritoneal mucinous carcinomatosis. There is overwhelming evidence that DPAM arises from a mucinous adenoma of the appendix. The one exception to this is the presentation of a mature ovarian cystic teratoma as PMP where the appendix is normal. We report such a case and discuss the presentation, histopathology, and treatment options.

CASE REPORT
A 39 year old woman presented with a three month history of abdominal distension and an episode of abnormal per vaginal bleeding. Clinical examination revealed a mass arising out of the pelvis and ultrasound examination demonstrated a large multicystic complex mass containing fluid and solid components, arising from the left ovary. The tumour marker CA125 was normal at 11 KU/litre. At laparotomy, copious amounts of free mucin and tenacious mucinous deposits were seen on the surface of the omentum and bowel. There was a 10 x 12 cm left ovarian mass and a small right ovarian cyst. Peritoneal lavage, hysterectomy, bilateral salpingo-oophorectomy, omentectomy, and appendicectomy were performed. After an uneventful recovery she was referred to our PMP centre for further management. Nine months after surgery, an abdominal computed tomography scan shows no evidence of PMP and tumour markers such as carcinoembryonic antigen, CA125, and CA19-9 are within normal limits.

HISTOPATHOLOGY
Macroscopic examination revealed a multiloculated, cystic, left ovarian mass measuring 14 x 12.5 x 7.5 cm. The cut surfaces revealed multiple cysts with smooth walls containing mucinous material. One of the cysts contained sebaceous and hair. The appendix measured 4 cm with no focal lesions identified on the cut surface.

Paraffin wax embedded sections of the left ovary confirmed the presence of a mature cystic teratoma (dermoid cyst) and a mucinous tumour. The teratomatous component consisted of a cyst lined by keratinising squamous epithelium containing skin appendages, such as sebaceous and apocrine glands (fig 1A). The mucinous component was multiloculated with the cysts lined by a single layer of mucin secreting epithelium (fig 1B). There were intestinal changes characterised by the presence of goblet cells and occasional Paneth cells. Sections of the appendix demonstrated mucinous material on the serosal surface. No abnormal epithelial proliferation was seen along the entire length of its lumen. The surfaces of the uterus and omentum also demonstrated mucinous material but without epithelial elements.

On immunoperoxidase staining, the mucinous epithelium in the ovary was cytokeratin 20 (CK20), carcinoembryonic antigen, and BerEP4 positive and CK7 negative.

Abbreviations: CK, cytokeratin; DPAM, disseminated peritoneal adenomucinosis; PMCA, peritoneal mucinous carcinomatosis; PMP, pseudomyxoma peritonei
DISCUSSION
Although the association of appendiceal mucinous tumours/mucocoele with PMP has long been recognised,2 the true origin of PMP in women who have simultaneous appendiceal and ovarian tumours has been a dilemma. Advances in immunohistochemical and molecular genetic techniques have provided compelling evidence that the source of PMP is an appendiceal mucinous tumour.3,4 In the past decade, detailed classifications of PMP have emerged with descriptions of distinct pathological characteristics of prognostic relevance. PMP has been classified into DPAM, characterised histologically by bland to low grade adenomatous mucinous epithelium and an associated appendiceal mucinous neoplasm; peritoneal mucinous carcinomatosis (PMCA), often with an associated mucinous adenocarcinoma of the gastrointestinal tract; and PMCA with intermediate or discordant features, which has combined features of DPAM and PMCA.7 It has been suggested that the term PMP should be used to describe a clinical syndrome of mucinous ascites along with the pathological diagnosis of DPAM.6

There are four important points to consider in the above case. (1) The appendix was macroscopically and microscopically normal, suggesting a primary ovarian mucinous tumour. (2) The morphology and immunophenotype of the epithelium lining the mucinous cysts in the ovary suggested a gastrointestinal origin (CK20+/CK7-). (3) The presence of squamous epithelium and skin appendages in the ovary indicated the presence of a mature cystic teratoma. (4) Peritoneal mucinous deposits were devoid of lining mucinous epithelium.

The association of ovarian teratomas, mucinous ovarian tumours, and PMP has been reported in two recent studies. The first was a study of mucinous tumours of the ovary (of intestinal type) in which three cases of PMP with an associated ovarian dermoid cyst were reported.8 Of these, two had a normal appendix, did not demonstrate peritoneal DPAM lesions, and had no evidence of disease recurrence after five years and 16 years. The third patient in whom neither the appendix was removed nor the peritoneal lesions sampled was alive without clinical evidence of recurrence after a follow up period of four years and nine months. The CK immunophenotype of these tumours was not reported.

The second study reported three cases of mucinous tumours in ovarian cystic teratomas,9 where the appendix was microscopically normal and the mucinous tumours had a CK20+/CK7- immunophenotype. The peritoneal lesions had histological features suggestive of PMP/DPAM in two cases and PMCA with intermediate or discordant features in the third. No follow up data were reported for this group of patients.

"The lack of lining epithelium in the peritoneal mucinous deposits, combined with normal tumour markers and normal imaging, make it difficult to outline the treatment that should be offered to our patient"

An appendiceal mucinous neoplasm can be found in almost all cases of DPAM/PMP with the notable exception of those arising in the gastrointestinal component of a mature cystic teratoma. Primary ovarian mucinous tumours are usually CK7 positive and show variable expression of CK20,10 unlike those associated with mature teratomas, which have the CK20+/CK7- phenotype.10

The above factors, particularly the lack of lining epithelium in the peritoneal mucinous deposits, combined with normal tumour markers and normal imaging, make it difficult to outline the treatment that should be offered to our patient. If her diagnosis is PMP/DPAM, then she is at risk of recurrent PMP and should undergo cytoreduction and hyperthermic intraperitoneal chemotherapy, the currently accepted treatment.11 However, the similarity of her pathology to that of the three patients in the first of the above studies suggests that she could be treated conservatively, because up to the time of reporting PMP had not recurred in these patients.

The above considerations have led to a "wait and watch" approach to the management of our patient, with plans to carry out six monthly tumour marker measurements and annual computed tomography scans.

Take home messages

- We report a patient with a mature ovarian cystic teratoma in whom the appendix was normal and who presented with pseudomyxoma peritonei
- The pathological, histological, and tumour marker characteristics of our patient make it difficult to classify this tumour into a distinct category (such as disseminated peritoneal adenomucinosis or peritoneal mucinous carcinomatosis) and advise the patient on the appropriate treatment
- Therefore, we have decided to take a "wait and watch" approach, and plan to carry out six monthly tumour marker measurements and annual computed tomography scans

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Accepted for publication 22 December 2004

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