Routine examination of appendices in Hong Kong

Drs Chan and Fu made a retrospective study of more than 11,000 resected appendices and described a number of relatively rare lesions, some of which had not been suspected clinically. They estimated that 0.74% of cases contained clinically important pathological findings which were likely to have been missed had the appendix not been examined histopathologically.

It was surprising that they made no mention of the not uncommon situation in which an appendix is removed, has equivocal macroscopic appearances, and is not microscopically to show gross acute serosal inflammation without evidence of intrinsic inflammatory change. This conclusion indicates a presence of a pronounced inflammatory focus within the abdomen but outside the appendix. Most of these cases occur in young women with a salpingitis that was not apparent at operation. Such information is surely of help to the clinician in the further management of these patients whose abdominal pain is unlikely to have been relieved by appendicectomy and who often benefit from postoperative antibiotics.

Reference


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Drs Chan and Fu comment:
The comment made by Dr Coghill is quite valid. We have indeed come across appendices in which clinically important inflammation is present only in the serosa, indicating the presence of an inflammatory focus within the abdomen other than the appendix. Without careful histopathological examination, some of these appendices could be taken to have acute appendicitis on external examination. Similarly, the finding of a perfectly normal appendix in a case of suspected acute appendicitis is also of clinical importance, as this should prompt the clinicians to investigate for a focus of inflammation elsewhere in the abdomen. Negative findings are therefore of definite clinical importance in some cases.

Using our method of study, the appendices showing serosal inflammation alone are mixed with a large group of acute appendicitis as both lesions are indexed by the same code. As a result, we have not been able to separate and study the effects of histopathological examination of this subgroup. The estimated 0.74% of clinically important pathological diagnoses made on routine histopathology must be taken as a baseline value. More accurate estimates will need further study, with the participation of surgeons.

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Embryonal carcinoma of testis simulating seminoma

I was interested to read the article by Alderdice and Johnston. I have looked carefully at the photomicrographs in the article and I can see no tumour which I would diagnose as seminoma, even in the so-called "semimoma like areas":

The authors use rather confusing terminology. They start by using the British terminology where "teratoma" implies a wide range of non-seminomatous germ cell tumours, but quickly change to American/WHO terminology. No reference is made to the British Testicular Tumour Panel and Registry, and the author's difficulties in differentiating between seminoma and malignant teratoma undifferentiated (MTU/embryonal carcinoma (EC)), may stem from their lack of acquaintance with the British publications. The British Testicular Tumour Panel and Registry emphasise the similarities between seminoma and MTU in some instances but "points in favour of a teratoma are...a greater degree of cellular pleomorphism and mitotic activity, and a tendency for the nuclei of the tumour cells to overlap one another..." The seminoma cell nucleus frequently contains a single prominent, rounded, often eosinophilic, nucleolus, which contrasts with the usually dense and hyperchromatic nucleoli, often multiple and of variable size, in the teratoma.

This description clearly excludes a diagnosis of seminoma in a "semimoma like area", especially as fig 2 closely resembles Pugh's figure illustrating a typical teratoma.

References


Dr Alderdice comments:
I feel that Dr Grigor concentrates too quickly on the fine nuclear detail and misses the main point behind the publication. On low power histological examination, each of these germ cell tumours bore a fibrous stroma containing abundant lymphocytes, and the first case had giant cell granuloma. The tumour cells were arranged for the most part in broad solid islands, and the initial impression of several consultant pathologists was that the overall pattern appeared seminomatous rather than teratomatous in differentiation.

Several of the nuclear points which Dr Grigor makes in favour of the diagnosis of teratoma—that is, cellular pleomorphism and mitotic activity—were mentioned in the discussion on diagnosis in the article, and both the letter writer and ourselves are in complete agreement as to the final diagnosis when the cellular detail is carefully examined.

The aim of this article was to illustrate the area of overlap between seminoma and embryonal carcinoma (malignant teratoma undifferentiated), and to point out that one does have to examine cellular and nuclear detail carefully after noting the overall pattern to reach the correct diagnosis. In this latter point we are in complete agreement.

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Letters to the Editor

It is regrettable that Alderdice and Johnston only referred to American publications and omitted to include British publications which would have given the correct diagnosis.

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