CASE REPORT

Pure testicular carcinoid associated with intratubular germ cell neoplasia

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This case report describes for the first time a case of pure testicular carcinoid preaortic lymph node metastases in a 25 year old patient with carcinoid syndrome. The simultaneous occurrence of intratubular germ cell neoplasia in the surrounding testicular tissue was identified by OCT4 and placental-like alkaline phosphatase positivity. This confirmed that the tumour had a germ cell origin in the testis, rather than being a metastasis from an extragenital carcinoid.

About 90 cases of carcinoid tumour of the testis have been reported to date. Histologically, most correspond to pure patterns and a few occur as a component of testicular teratoma. From a diagnostic viewpoint, it is important to differentiate primary testicular carcinoids from either metastases from extragenital carcinoids or testicular neoplasms of non-germ cell origin resembling carcinoids, such as granulosa cell tumours.

We report, for the first time, a case of pure testicular carcinoid with lymph node metastases and carcinoid syndrome. In this tumour, the concomitant presence of intratubular germ cell neoplasia (ITGCN) in the surrounding testicular tissue helped to identify its germ cell origin in the testis and to eliminate the possibility of a metastasis from an extragenital carcinoid.

CASE REPORT

A 25 year old patient presented with dull scrotal pain. He had had a mass in the left testis since childhood, and since adolescence had experienced bouts of watery diarrhoea and flushing episodes, considered allergic in nature by his family practitioner.

Physical examination revealed a hard left testicular mass, which ultrasound showed to be a solid, nodular, heterogeneous lesion with isolated irregular calcifications. Both α fetoprotein and β human chorionic gonadotrophin concentrations were within normal limits. A left orchidectomy was performed. After a histopathological diagnosis of pure testicular carcinoid, we measured 5-hydroxyindole acetic acid concentrations and found them to be raised at 330 ng/ml. Both diarrhoea and flushing were reduced after surgery. The presence of a carcinoid tumour elsewhere was initially discarded by full body scintigraphy with 111In-octreotide. However, 11 months later an abdominal computerised tomography scan revealed preaortic lymph node involvement, confirmed surgically by the finding of masses at lymphadenectomy. However, 5-hydroxyindole acetic acid concentrations remained within normal limits. Surgical inspection of intestine and liver failed to demonstrate a mass. The caecal appendix was unremarkable. The patient is presently alive and symptom free.

PATHOLOGY

The left testis contained a 6 × 5.5 × 4.5 cm, white/yellow, hard, nodular tumour with only a thin rim of testicular tissue. No gross involvement of the tunica albuginea or adnexa was evident. Sampling comprised a total of 15 tissue blocks. Microscopically, the tumour was an insular-type carcinoid arranged in nests and acini separated by broad collagenous tracts (fig 1A). Coagulative necrosis was present in the central area. Cells were regular and homogeneous, lacking atypia or mitoses, but there were microscopic foci of capsular and adnexal invasion. Immunohistochemistry revealed pronounced cytoplasmic positivity to synaptophysin and chromogranin (fig 1B); placental-like alkaline phosphatase (PLAP) was negative in the carcinoid areas. Intratubular extension of carcinoid was not detected. The surrounding testicular parenchyma was reduced to an atrophic band revealing widespread foci of ITGCN, both at a distance and in the vicinity of the carcinoid (fig 2A), showing cells with irregular nuclei and prominent nucleoli, (fig 2B) some displaying mitoses. Immunohistochemically, they revealed

Figure 1 (A) Characteristic nests of insular type of carcinoid are (B) positive for chromogranin.

Abbreviations: ITGCN, intratubular germ cell neoplasia; PLAP, placental-like alkaline phosphatase
membranous PLAP positivity (fig 3A), but they were synaptophysin and chromogranin negative. OCT4 (Santa Cruz Biotechnology, Santa Cruz, California, USA), a novel marker for ITGCN,4 was positive in some, but not all, of the atypical germ cell nuclei of the ITGCN of neighbouring and distant tubules (fig 3B).

Metastatic tumour, measuring 0.3 to 1 cm, in the four preaortic lymph nodes, was morphologically identical to the testicular primary.

DISCUSSION

Our case provides an insight into the origin of testicular carcinoid. As with other heterotopic single tissue testicular tumours, such as epidermoid cysts, chondroma, etc., the origin of testicular carcinoid remains debatable, and consequently the World Health Organisation classification of testicular tumours5 includes carcinoid in the miscellaneous section.

Several histogenetic possibilities have been considered for these neoplasms. An origin from argentaffin cells was proposed by some,6 but was subsequently discarded because of the absence of such cells in the testis. A germ cell origin seems a more likely possibility and this is evident when the carcinoid is a component of testicular teratoma. Nevertheless, pure carcinoids are more frequent and are considered to be either monodermal germ cell tumours or the remaining component of burnt out teratoma.7

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The finding of a concomitant precursor lesion of germ cell tumours, such as ITGCN,8 would support such an origin. However, two recent series of testicular carcinoids1 2 were unable to find ITGCN; an absence difficult to explain once adequate sampling is discarded. Our present case reports for the first time ITGCN in the testicular tissue surrounding a pure carcinoid. The tubules adjacent to the carcinoid showed characteristically atypical intratubular cells with membranous PLAP positivity and nuclear positivity for OCT4, a highly specific marker for ITGCN.3 Our case shows similarities to a previous one,9 where a testicular mature cartilaginous nodule was thought to be a monodermal teratoma because it was associated with ITGCN. Similarly, a pure Wilms tumour of the testis was shown to be a type of monodermal teratoma10 by the identification of i(12p)—a characteristic chromosomal marker of testicular germ cell tumours—in the tumour karyotype. This finding is a more accurate demonstration of a germ cell origin than the more broadly non-specific study by fluorescence in situ hybridisation analysis of X chromosome gain in a series of testicular carcinoids.1

From both clinicopathological and diagnostic viewpoints, it is important to distinguish between carcinoid as a primary testicular monodermal teratoma as opposed to a metastasis from an extragenital carcinoid of non-germ cell origin.11 The last situation is extremely rare, with only 10 cases published.
in the literature, most of which originated in the gastro-intestinal tract.1

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The patient gave his informed consent for this case report to be published

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