Glutathione S-transferase expression in the human testis and testicular germ cell neoplasia

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Summary Glutathione S-transferase (GST) isoenzyme expression is altered in a variety of neoplasms and the enzymes are implicated in metabolism of carcinogens and resistance to drugs, including cisplatin. We have studied GST Alpha, Pi, Mu and microsomal isoenzyme expression by immunohistochemistry in normal and cryptorchid testes, intratubal germ cell neoplasia (ITGCN), seminoma and non-seminomatous germ cell tumours. In 16 stage II–IV malignant teratoma intermediate (MTI) both orchidectomy and post-treatment residual surgical masses were studied.

All four isoenzymes were strongly expressed in Leydig and Sertoli cells. GST Pi was absent from normal spermatogonia but strongly expressed by the neoplastic germ cells of ITGCN and seminoma. GST Pi was strongly expressed in all elements of teratoma, irrespective of differentiation. There were no qualitative differences in expression between primary and post-chemotherapy metastases. GST Alpha expression in teratoma correlated with epithelial differentiation. GSTs may be important in normal spermatogenesis and protection of germ cells from teratogens and carcinogens. They may have a role in testicular tumour drug resistance but this role is not well defined. GST Pi is a new marker for ITGCN.

Materials and methods

Cases

A total of 62 cases were studied. These were formalin-fixed, paraffin embedded biopsies from archives (1977–1990) where clinical follow-up data were available. These included 5 normal testes, seven cryptorchid testes and 11 cases of ITGCN. Thirty-nine cases of invasive germ cell tumours were studied composed of 22 malignant teratoma intermediate (MTI), eight malignant teratoma undifferentiated (MTU), six seminoma and three yolk sac tumours (British classification) (Pugh & Cameron, 1976, pp. 202–204). The age range of the tumour cases at time of diagnosis was 15–51 years. The normal testes were of histologically normal cases in males of 50 years and over, who had not received any hormonal therapy. In 16 stage II–IV MTI (Royal Marsden Hospital stage) (Heydig et al., 1980) both orchidectomy specimens and post-treatment residual surgical masses were available for study (Table I).

Immunostaining

Polyclonal antisera raised in the rabbit to GST Pi, Mu, Alpha and microsomal classes were the kind gift of Dr J.D. Hayes, University Department of Clinical Biochemistry, Edinburgh. Isoenzyme preparation, rabbit immunisation and characterisation of antibodies have been previously described (Morgenstern et al., 1990; Hayes et al., 1983; Hayes et al., 1986).

Sections of formalin-fixed, paraffin-embedded tissues were cut at 3 μm, dewaxed in xylene and incubated with primary antibody at a dilution of 1:200 in phosphate buffered saline for a period of one hour at 20°C. Detection was by biotinylated swine anti-rabbit immunoglobulin (Dako) and streptavidin-biotin-peroxidase complex (Dako). Visualisation

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Table I  Treatment regimes and survival data for 16 stage II–IV malignant teratoma intermediate.

| Case | Stage at presentation | Treatment | Status | Duration of follow-up (months) |
|------|-----------------------|-----------|--------|-------------------------------|
| 1    | IVL1                  | BEP × 4   | DIED   | 8                             |
| 2    | II B                  | BPV/E × 4 | A + W  | 72                            |
| 3    | II B                  | BEP × 4   | A + W  | 48                            |
| 4    | II C                  | BPV/E × 4 | A + W  | 66                            |
| 5    | II C                  | BPV/E × 4 | A + W  | 62                            |
| 6    | II A                  | BEP × 4   | A + W  | 62                            |
| 7    | IV H +                | PVB × 5   | DIED   | 25                            |
| 8    | II A                  | PVB × 6   | DIED   | 17                            |
| 9    | II A                  | PVB × 5   | DIED   | 10                            |
| 10   | II B                  | BPV/E × 6 | A + W  | 36                            |
| 11   | IV CL2                | PVBEM × 6 | A + W  | 60                            |
| 12   | II C                  | BEP × 6   | A + W  | 13                            |
| 13   | III A                 | BPV/E × 4 | A + W  | 59                            |
| 14   | IV CL2                | BEP × 5   | A + W  | 55                            |
| 15   | III C                 | PBVEM × 6 | A + W  | 55                            |
| 16   | II C                  | BEP × 4   | A + W  | 94                            |

B = Bleomycin, E = Etoposide, V = Vinblastine, P = Cisplatin, M = Methotrexate, A + W = Alive and well.

using 3,3′-diaminobenzidine as substrate produced a brown insoluble precipitate. Sections were lightly counterstained with haematoxylin. Liver tissue was used as a control, GST Pi staining bile duct epithelium and GST Alpha, Mu and microsomal classes staining hepatocytes.

Results

Normal and cryptorchid testes

Sertoli cells and Leydig cells showed strong positive cytoplasmic staining for GST Pi, Alpha, Mu and microsomal classes in all 5 normal testes. GST Pi was also present in nuclei (Figure 1). Spermatagonia and primary spermatocytes were negative in all normal testes. Faint reactivity of secondary spermatocytes and spermatids for GST Pi and Mu classes was seen in each case. In seven testes where histologically normal epididymal tissues were present, the epididymal lining epithelium was strongly positive for both GST Pi and Alpha. Six cases were positive for GST Mu and four for GST microsomal class.

In the seven cryptorchid testes there was a similar pattern of Leydig and Sertoli cell GST expression to the normal testis but the intensity appeared stronger (Figure 2).

**ITGCN**

The neoplastic germ cells in 11 cases of ITGCN showed strong nuclear and peripheral cytoplasmic staining for GST Pi (Figure 3). Faint staining for GST Pi and Mu positivity was present in five cases and GST microsomal in four cases.

**Seminoma**

GST Pi was strongly expressed in all eight cases (Figure 4) with occasional negative cells and showed a similar cellular distribution to ITGCN. GST Alpha was positively expressed in seven cases, GST Mu in five cases, and GST microsomal in four cases.

**Teratoma**

All well differentiated epithelial elements of the 6 stage I MTI and 16 stage II–IV MTI testes showed strong expression of GST Pi (Figure 5a) and Alpha classes (Figure 5b). Mesenchymal elements (cartilage and smooth muscle), where present, were uniformly negative for GST Alpha class but strongly positive for GST Pi class (Figure 5). Undifferentiated elements (embryonal carcinoma) showed strong diffuse predominantly nuclear positivity for GST Pi in all cases. GST Alpha was negative for undifferentiated elements in three of 22 cases of MTI. In 19 positive cases of MTI the undifferentiated elements showed a weak background positivity with strong focal staining for GST Alpha related to areas of tubular and papillary differentiation (Figure 6). GST Mu and microsomal were positive in all cases of MTI but did not show any consistent relation to epithelial or mesenchymal differentiation.

There was no significant difference between staining pat-
The eight cases of MTU showed strong diffuse mainly nuclear staining for GST Pi. GST Alpha expression was similar to the undifferentiated elements of MTI; with focal strong positivity associated with tubular formations. All eight cases expressed GST Mu and four cases were positive for GST microsomal class. Three cases of yolk sac tumour showed positive expression of GST Pi, Alpha, Mu and microsomal classes.

**Discussion**

We have demonstrated that GST Pi, Alpha, Mu and microsomal expression is present in and largely confined to the Sertoli and Leydig cell compartments of the normal human testis. Normal spermatogenesis is critically dependent on the close association of active Sertoli cells and developing germ cells (Ritzen et al., 1981). Sertoli cells are largely responsible for germ cell o xo-reductive enzyme systems (Voganathan et al., 1989a) and glutathione production (Li et al., 1989). Testes of mature rodents contain high concentrations of GSH (Mushawar & Koepppe, 1973) and increase in testicular GSH parallels spermatogenic cell development (Grosshans & Calvin, 1985). GSH protects against germ cell mutagenesis by ethyl methanesulphonate in the F-344 rat and this is dependent on enzymatic conjugation by GST (Teaf et al., 1985). GSTs may be an important enzyme system in detoxification of xenobiotics in the human testis but the role of individual carcinogens and teratogens is largely unknown. The demonstration of GST expression in the epididymis parallels previous studies in the rat testis (Hales et al., 1980). At this site they may be important in final stages of spermatozoal maturation and provide protection against teratogens. The function of GSTs in the Leydig cell has not been defined. They have a role in steroid transport and isomerisation (Listowsky et al., 1988) and this may also be of importance in the Sertoli cell. GST Alpha is present in the corresponding cell in human ovary, the enzymatically active stromal cell (EASC) (Rahilly et al., 1991). The nuclear distribution of GST Pi has been previously described but its significance remains uncertain (Rahilly et al., 1991).

We demonstrated GST Pi expression in ITGCN but not in normal spermatagonia. This is consistent with its expression in preneoplasia in other organs (Sato, 1989) and it shows a striking parallel with expression of placental alkaline phosphatase (PLAP). This latter enzyme is expressed in fetal germ cells, ITGCN (Hustin et al., 1987) and seminoma and is used as a marker in routine diagnosis. GST Pi appears to reliably distinguish ITGCN from normal spermatagonia but is not such a specific marker. It has been shown that cisplatin-based chemotherapy may not eradicate ITGCN in some cases (Fossa & Aass, 1989; Chong et al., 1986) implying a drug-

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**Figure 4** Expression of the GST Pi isoenzyme in seminoma. (bar = 50 μm)

**Figure 5** a Expression of GST Pi isoenzyme in MTI chondrocytes and well differentiated epithelium. (bar = 125 μm). b Expression of GST Alpha isoenzyme in MTI well differentiated epithelium. The chondrocytes are negative. (bar = 125 μm)

**Figure 6** Focal expression of GST Alpha isoenzyme in embryonal carcinoma (MTU) in relation to tubulo-papillary formations. (bar = 125 μm)

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terms for GSTs Pi and Alpha in MTI Stage I and MTI Stage II–IV.

The pattern of expression in post-treatment metastatic deposits of MTI Stage II–IV was similar to the primary tumour and reflected the component elements (differentiated epithelium, mesenchyme, embryonal carcinoma). There were no significant differences in isoenzyme expression between survivors and those who died of their disease.
resistant phenotype and/or protection by the blood-testis barrier; and this may be important in development of a second primary tumour in the contralateral tests.

In seminoma and teratoma GST P1 was expressed in all tumour elements, irrespective of degree or line of differentiation. We were unable to demonstrate any qualitative difference between untreated primary tumours and post-chemotherapy metastatic deposits nor correlate expression with survival. However, the immunohistochemical method is relatively insensitive to changes in levels of expression and the study was necessarily limited by the availability of tissue from patients for whom follow-up data was available. GST P1 expression has been correlated with cisplatin resistance in human tumour cell lines, (Nakagawa et al., 1988; Teicher et al., 1987) but the implications in the clinical situation are uncertain. Raised cellular GSH levels protect against Bleomycin cytotoxicity (Russo et al., 1984) but there is no direct evidence of GSH/GST involvement in resistance to vinca alkaloids or etoposide (VP-16).

We have shown that GST Alpha class expression is related to epithelial differentiation and that in embryonal carcinoma (MTU) expression is focal and related to tubal and papillary formations. In post-chemotherapy surgically resected masses differentiated elements are often the only viable component. It is believed that this is due to selective destruction of more primitive elements rather than chemotherapy-induced differentiation (McCartney et al., 1984); and implies that differentiated elements are drug resistant, even though they behave in a less malignant fashion.

In summary GSTs are widely expressed in normal and neoplastic testicular tissues. They may be important in spermatogenesis and resistance to teratomas and carcinogens in the normal tests. They show characteristic patterns of expression in testicular tumours but their role in drug resistance is not well defined. GST P1 is a new marker for ITGCN.

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