Glutathione-s-transferase pi expression in leukaemia: a comparative analysis with mdr-1 data

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Summary Drug resistance in haemopoietic cells may be partly related to the expression of the glutathione-s-transferase (GST) pi and mdr-1 genes. We have used RNA slot blotting techniques to investigate the expression of GST pi in peripheral blood and bone marrow of nineteen patients with myelodysplastic syndrome (MDS), eighteen patients with acute myeloblastic leukaemia (AML), and thirty-two patients with chronic lymphocytic leukaemia (CLL). We found increased expression of GST pi in 8 of 9 MDS, (7 peripheral blood, 1 bone marrow) 12 of 18 AML (3 peripheral blood, 7 bone marrow; 4 of 5 untreated, 1 of 5 secondary, 7 of 11 relapse or refractory) and in the peripheral blood of 24 of 32 CLL (3 of 7 untreated, 21 of 25 treated) relative to normal controls. Increased expression of GST pi can occur at any stage of disease and shows no clear relation to mdr-1 expression except, possibly, in CLL. In 3 AML patients GST pi transcript levels were the same or lower on relapse compared to presentation. Upregulation of the GST pi gene could not be demonstrated in 2 CLL patients in response to treatment with intermittent chlorambucil.

Resistance to cytotoxic drugs, either at presentation, or, more frequently, at relapse, is commonly encountered during the treatment of leukaemia. It is well established in vitro that cellular acquisition of the multidrug resistant (MDR) phenotype confers loss of sensitivity to a wide range of structurally unrelated toxins (Biedler & Riehm, 1970; Kartner et al., 1985). Several distinct drug resistance processes are now recognised which may be involved in clinically resistant leukaemia (Holmes, 1990b). The mdr-1 gene encodes the transmembranous P-glycoprotein (PGP, Mr = 170,000) which acts as an energy-dependent efflux pump and is associated with decreased intracellular accumulation of drugs (Chen et al., 1986; Juliano & Ling, 1976). The function of the protein encoded by the mdr 2 gene in man is unknown. Mdr-1 gene amplification and increased expression have been seen in the drug-resistant CEM/VLB/100 leukaemia cell line when compared to the parental sensitive CCRF-CEM line. However, mdr 1 gene amplification has not been observed in AML or MDS (Holmes et al., 1989). Elevated mdr 1 mRNA levels have been reported in many forms of leukaemia (Goldstein et al., 1989; Holmes et al., 1989, 1990a).

The glutathione-s-transferases (GST) have also been implicated in drug resistance (Hayes & Wolf, 1988). They comprise four gene families which map to distinct chromosomal locations. Three of these gene families encode cytosolic proteins which have been classified as alpha, mu, and pi (Mannervik et al., 1985). GST isozymes catalyse the conjugation of electrophilic drugs, toxins and carcinogens to reduced glutathione which leads to detoxification. They also detoxify toxic hydroperoxides and bind to sequester other foreign compounds (Ketterer et al., 1986). Increased expression of a variety of GST subunits has been observed in cell lines made resistant to cytotoxic drugs. (Hayes & Wolf 1988). In certain cases concomitant over expression of both GST pi and mdr-1 mRNA have been observed (Cowan et al., 1986). Over expression of both mdr-1 and GST has also been seen in carcinogen-induced preneoplastic lesions in rat liver indicating that there may be co-ordinate expression of these proteins (Ktahara et al., 1984). Elevated levels of GST pi, mRNA have been found in a variety of haematological malignancies but no clear relationship with chemotherapy can be determined. (McQuaid et al., 1989; Moscow et al., 1989).

The myelodysplastic syndrome (MDS) represents a group of conditions characterised by peripheral cytopenias and ineffective haemopoiesis. Many patients go on to develop acute myeloblastic leukaemia (AML). In addition, AML can arise de novo, or secondary to chemotherapy for malignancy, or other toxic bone marrow damage. Chemotherapy may produce a remission in AML but the disease is characterised by relapse which is often resistant to further cytotoxic treatment. CLL is a disease of slow progression with accumulation of malignant lymphocytes in blood, bone marrow and lymphoid tissue. Drug treatment can result in suppression of lymphocyte count but with time re-emergence of the malignant clone occurs and cure is not usually possible.

To identify the incidence and possible association of GST pi and mdr-1 expression in leukaemia patients we have studied any treatment related changes in the expression of these genes; we have investigated their expression in haemopoietic cells of normal control subjects and untreated and treated patients with MDS, AML and CLL.

Materials and methods

Methods for mdr-1 expression have previously been published (Holmes et al., 1989, 1990a). GST pi expression was investigated in total peripheral leucocytes of ten normal control subjects (4 men and 6 women) aged 23 to 83 years, and total bone marrow of one normal male aged 75 years. None of the normal control subjects had any significant exposure to drugs or environmental toxins. Nine patients with MDS of varying sub-type, 18 with AML of varying type and stage of disease and 32 patients with CLL, both untreated and treated of varying stage have also been investigated. For those patients with MDS, none of whom had received cytotoxic therapy, one bone marrow and eight peripheral blood samples were studied. For all cases of AML blood or bone marrow contained greater than 90% blasts. Only peripheral blood was studied in CLL and all samples contained greater than 70% lymphocytes. All cases of AML were treated with adriamycin, cytosine and thioguanine at presentation and if refractory, or on relapse, with either bisantrene or mitoxantrone and cytosine. CLL patients were all treated with intermittent or continuous low dose chlorambucil.

DNA and total RNA were extracted by lysis in guanidine thiocyanate and centrifugation in a caesium chloride gradient (Maniatis et al., 1982). Concentrations of mRNA were determined spectrophotometrically and duplicate slot blots were made with five doubling dilutions of total mRNA (10 µg to

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0.625 μg) on to Hybond N (Amersham) membranes. Each slot blot included the CEM line and its drug-resistant derivative CEM/VLB/100 as negative and positive controls respectively for mdr expression (Beck et al., 1979). Hybridisation was carried out using a 414 bp fragment of the mdr cDNA P5L-18 (Scotto et al., 1986) as described by Holmes et al. (1989) and then to a full length GST pi cDNA (Kano et al., 1987). Probes were labelled with 32PdCTP by primer extension (Boehringer). Quantitation of RNA was carried out by hybridisation with a human B-actin (PHA4.1) probe (Khalili et al., 1983). The blots were finally washed with 2 × SSC, (sodium chloride, sodium citrate), 1% SDS (sodium dodecyl sulphate), at 65°C for 1 hour, and 0.1 × SSC 1% SDS at 60°C for 30 minutes for GST pi and B actin respectively. Kodak XAR-5 film exposed at −70°C for 1–5 days was used for autoradiography. Relative levels of GST pi expression were determined by densitometry.

**Results**

Results are summarised in Table I. The data relating to mdr-1 expression in the same samples examined here have been published previously (Holmes et al., 1989, 1990a).

Representative data for relative levels of GST pi expression in a normal control, CEM and CEM/VLB/100 cell line and patients with increased expression in MDS, AML and CLL are shown in Figure 1. Total peripheral leucocytes from the same 10 normal controls and total bone marrow from one individual demonstrated a range of GST pi mRNA expression of 1 to 3 arbitrary units. It was hoped that analysis of GST pi expression in normal myeloid and lymphoid fractions in peripheral blood could be obtained as had been completed with mdr analysis. However, no signal could be detected from the relevant membranes and no RNA was available to construct further blots. All peripheral blood samples contained 30–40% lymphocytes whilst the one bone marrow examined held less than 5% lymphocytes. It is not feasible to obtain data on GST pi expression on normal blast cells since approximately 10⁶ cells are required for RNA extraction. Therefore, no truly adequate control exists for the analysis of GST pi expression in AML. We have compared GST pi RNA level in CLL to those in total peripheral blood. Ideally data on normal lymphoid fractions would have provided better controls.

All pre-leukaemic and leukaemic samples contained GST pi mRNA within a 28-fold range of variations in expression. Eight of nine (one bone marrow, seven total peripheral blood) patients with MDS of all sub-types demonstrated GST pi mRNA (4-10 units) above the normal range. Elevated RNA levels were seen in 12 of 18 cases of AML (5 peripheral blood, 7 bone marrow) including 4 of 5 untreated AML (6–8 units), one of 5 secondary AML (secondary to previous chemotherapy and radiotherapy for carcinoma of

| Table I | GST pi expression in normal control subjects, cell lines and leukaemic samples |
|---------|---------------------------------|
| Results | Relative levels of RNA GST pi |
| Normal controls | Mean +/− | Standard Error | Range |
| Total peripheral leucocytes | 10 | 2.3 ± 0.2 | 1–3 |
| Total bone marrow | 1 | 2 |
| Cell lines | |
| CEM (parental drug sensitive lymphoblastic line) | 9 |
| CEM/VLB/100 (drug resistant mutant line) | 9 |
| Patients with increased GST pi expression | |
| | Mean +/− | Standard Error | Range |
| MDS | 8/9 | 5.6 ± 0.8 | 4–10 |
| AML Untreated | 4/5 | 7 ± 0.6 | 6–8 |
| Secondary | 1/5 | 6 | 6 |
| Refractory | 7/11 | 10.1 ± 2.9 | 4–24 |
| CLL Untreated | 3/7 | 13.7 ± 6.8 | 5–27 |
| Treated | 21/25 | 10.7 ± 1.58 | 4–28 |

Figure 1 A = GST pi; B = B actin. Numbers 1–5 represent doubling dilutions of RNA (10 μg to 0.625 μg). Representative RNA slot blots of parental sensitive (CEM) and drug resistant (CEM/VLB/100) cell lines, normal leucocytes and patients with MDS, AML and CLL showing increased GST pi expression. One arbitrary unit represents the level of expression demonstrated in normal leucocytes. Relative levels of GST expression were determined by densitometry. Slot 1, lane A does not give a representative signal of GST pi expression due to either RNA overloading or the presence of an excess of RNA sample buffer. This signal is not used in any subsequent densitometric analysis. Levels of GST pi expressions are determined from the remaining 4 slots.
the breast, 6 units), and 7 of 11 cases of relapsed or refractory AML (4–24 units). Repeated measurements on three patients with GST pi levels (6, 8, 13 units) at presentation had the same or decreased expression after chemotherapy (2, 3, 6 units). Of the CLL patients 3 of 7 untreated (5–27 units) and 21 of 25 treated (4–28 units) demonstrated increased mRNA in peripheral blood. Sequential measurements on two patients could not demonstrate a change in gene expression with exposure to intermittent chlorambucil.

Gene amplification of GST pi could not be demonstrated in any of the patient material screened. Of all patients, those with normal GST pi and mdr-1 expression include none with MDS, one case of relapsed secondary AML, two untreated and four treated patients with CLL. Out of a total of 59 patients, only 7 patients had neither gene upregulated. In view of the possible association between GST pi and mdr-1 expression, a comparison was made between the relative levels in the same samples. There was no obvious association except for a possible weak correlation between GST pi and mdr-1 mRNA levels in CLL (Figure 2).

Discussion

As it is likely that malignant cells can use several mechanisms to achieve a drug-resistant state and maintain a proliferative advantage we have investigated the possible associations and relevance of two mechanisms that are currently considered to be important. Moscow et al. (1989) reported over expression of PGP and GST pi in the Adr

\[ ^{\text{r}} \text{MCF-7 breast carcinoma} \]

line and rat hyperplastic nodules. They further investigated 23 cases of lymphoma and observed generally higher levels of mdr-1 mRNA in low grade compared to high grade tumours whilst GST pi levels were uniformly low. Analysis of 7 cases of pre-B ALL suggested that increased GST pi expression is present before chemotherapy and may be higher at relapse.

To date, there have been no studies comparing GST pi and mdr-1 expression in human pre-leukaemia, AML and CLL. McQuaid et al. (1989) have investigated GST pi expression alone in 2 cases of MDS, 9 of AML and 1 case of ALL and the modest increase in expression was higher compared to four normal controls. Their data further suggested that, in two cases of AML and two of lymphoma, that transcription levels of GST pi fell after the introduction of chemotherapy indicating that perhaps gene upregulation is not an important mechanism in the detoxification of cytotoxics.

We have observed lower levels of GST pi mRNA in normal total peripheral leucocytes and total bone marrow. Data on GST pi expression in normal lymphoid and mdr-1 fractions are not available for technical reasons, as mentioned previously. In contrast, mdr-1 mRNA levels whilst low in normal lymphocyte populations can be high in normal total peripheral leucocytes and bone marrow suggesting that myeloid cells and possibly monocytes express high levels of mdr-1 mRNA. (Holmes et al., 1990a).

We have previously demonstrated mdr-1 gene amplification and increased RNA expression in the drug resistant CEM/VLB/100 leukaemic line when compared to the parental sensitive CEM line (Holmes et al., 1989). However, both the sensitive and resistant lines express similar levels of GST pi mRNA compared to normal peripheral blood and bone marrow. This suggests, in agreement with Moscow et al. (1989), that GST pi upregulation may be an inherent feature of lymphoblastic leukaemia. However, ALL, particularly in children, is chemosensitive raising a question concerning the relevance of increased GST pi expression. These data also suggest that the mdr-1 and GST pi gene may act independently to confer drug resistance.

Elevated levels of mdr-1 mRNA as compared to the basal expression of the drug sensitive CEM leukaemic cell line are more common in secondary and relapsed/refractory AML than in untreated AML (Holmes et al., 1989). Repeated measurements in three patients with AML with basal levels of mdr-1 mRNA at presentation, that is comparable to the CEM line, have shown that levels can be significantly increased at relapse. In contrast, GST pi levels are more commonly elevated above the normal range for blood and bone marrow in primary than secondary AML and repeated measurements on the same three patients suggest that RNA levels remain the same or fall after chemotherapy. This suggests that, unlike mdr-1, GST pi is not inductive by the therapeutic agents used. Furthermore, although GST pi may be upregulated initially, other drug resistance mechanisms may be relevant in subsequent clinical drug resistance.

For those patients with the pre-leukaemic syndrome, MDS, elevations of GST pi above the normal range for peripheral blood and bone marrow and mdr-1 mRNA relative to the drug sensitive CEM line are commonly observed. All patients demonstrated over expression of either GST pi, or mdr-1.

We have previously demonstrated increased levels of mdr-1 mRNA in approximately 50% of both untreated and treated cases of CLL, with upregulation of the mdr-1 gene occurring in response to intermittent chemotherapy with alkylating agents (Holmes et al., 1990a). GST pi upregulation appears more common in treated than untreated CLL although no clear relationship could be demonstrated in two patients treated with chlorambucil. There are now various reports which suggest an association between GST pi and mdr-1 expression (Kitahara et al., 1984; Cowan et al., 1986; Burt et al., 1988).

In AML and CLL patients all patterns of GST pi and mdr-1 expression were observed suggesting that these mechanisms act independently. Only in CLL does there appear to be a possible association between GST pi and mdr expression (\( r = 0.335, \ p = 0.05 \)). No pattern could be found with either GST pi or mdr-1 expression and age, sex of patient, FAB type of AML and MDS or stage of CLL.

![Figure 2 GST pi vs mdr RNA expression. Using Spearman's Rank Order Correlation test gives non significant negative correlation in MDS (\( r = 0.412 \)) and AML (\( r = 0.314 \)). Positive association of GST pi and mdr mRNA expression in CLL group (\( r = 0.335, \ p = 0.05 \), two tailed test).](image-url)
Within the AML and CLL groups there were patients, albeit a minority, (7 out of 50) with clinically resistant disease and no apparent involvement of either of these drug resistance mechanisms, which suggests that other processes may be clinically relevant.

Potmesil et al. (1988) have demonstrated low levels of topoisomerase II in CLL, suggesting this as a clinically relevant route to a drug-resistant state for those tumours with a large population of non-proliferating cells.

Although the clinical relevance of these data is not clear, the relationship of mdr-1 and GST pi expression to chemotherapy may provide important considerations for the future design of therapeutic regimes and suggest a complex relationship between drug resistance mechanisms in leukaemia.

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