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Glutathione S-transferase P1 (GSTP1) directly influences platinum drug chemosensitivity in ovarian tumour cell lines

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Background: Chemotherapy response in ovarian cancer patients is frequently compromised by drug resistance, possibly due to altered drug metabolism. Platinum drugs are metabolised by glutathione *S*-transferase P1 (*GSTP1*), which is abundantly, but variably expressed in ovarian tumours. We have created novel ovarian tumour cell line models to investigate the extent to which differential *GSTP1* expression influences chemosensitivity.

Methods: Glutathione S-transferase P1 was stably deleted in A2780 and expression significantly reduced in cisplatin-resistant A2780DPP cells using Mission shRNA constructs, and MTT assays used to compare chemosensitivity to chemotherapy drugs used to treat ovarian cancer. Differentially expressed genes in *GSTP1* knockdown cells were identified by Illumina HT-12 expression arrays and qRT–PCR analysis, and altered pathways predicted by MetaCore (GeneGo) analysis. Cell cycle changes were assessed by FACS analysis of PI-labelled cells and invasion and migration compared in quantitative Boyden chamber-based assays.

Results: Glutathione S-transferase P1 knockdown selectively influenced cisplatin and carboplatin chemosensitivity (2.3- and 4.83-fold change in IC₅₀, respectively). Cell cycle progression was unaffected, but cell invasion and migration was significantly reduced. We identified several novel *GSTP1* target genes and candidate platinum chemotherapy response biomarkers.

Conclusions: Glutathione S-transferase P1 has an important role in cisplatin and carboplatin metabolism in ovarian cancer cells. Inter-tumour differences in *GSTP1* expression may therefore influence response to platinum-based chemotherapy in ovarian cancer patients.

Ovarian cancer often presents at an advanced stage where treatment is rarely curative (Kristensen and Trope, 1997). Chemotherapy with platinum-based drug regimens (combining cisplatin or carboplatin with paclitaxel) can be initially effective, but longer-term treatment is frequently compromised by the development of drug-resistant disease (Vaughan *et al*, 2011). Platinum drugs are thought to act by promoting the formation of intra-strand DNA crosslinks, thus inhibiting DNA translation and replication (Eastman, 1987). Although several candidate drug resistance mechanisms, including impaired DNA repair, decreased drug uptake, increased drug efflux and detoxification, have been proposed (reviewed by Galluzzi *et al*, 2012), we are still some way from understanding the molecular basis of drug resistance, and from developing biomarkers to predict the onset and monitor the development of drug-resistant disease in cancer patients.

In recent work, we used qRT–PCR analysis to quantify intertumour differences in the expression of multiple candidate genes associated with disease progression and chemotherapy response in ovarian tumours (Smith *et al*, 2012). One of the most abundantly expressed genes in ovarian tumours and tumour cell lines was

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glutathione S-transferase P1 (GSTP1, EC 2.5.1.18), a polymorphic phase II drug-metabolising enzyme, which conjugates the antioxidant tri-peptide glutathione with many toxic hydrophobic and electrophilic xenobiotics to facilitate elimination (reviewed by Hayes and Pulford, 1995). Increased GSTP1 expression has been reported in pre-neoplastic lesions in chemically induced animal cancer models (Satoh et al, 1985) and in many human tumours, relative to surrounding normal tissues (Shea et al, 1988). Although a useful neoplasia biomarker, it is not known whether increased GSTP1 expression directly influences carcinogenesis, or is simply a bystander effect, where expression of this highly inducible gene is increased as part of the adaptive response mounted by the neoplastic cell. In support of a direct role in carcinogenesis, however, GSTP1 inhibits the stress-inducible c-Jun N-terminal kinase (JNK) in vitro, and JNK activity is reduced in GSTP1 null mice (Adler et al, 1999; Yin et al, 2000; Elsby et al, 2003). Additional roles for GSTP1 in the regulation of genes including TRAF2 (Wu et al, 2006), CDK5 (Sun et al, 2011) and the FAS death receptor (Anathy et al, 2012) have also been described.

Purified GSTP1 conjugates cisplatin in vitro (Hagrman et al, 2004), and GSTP1 expression is increased in human tumour cell lines either inherently or made resistant to chemotherapy drugs including cisplatin and various alkylating agents (Black and Wolf, 1991; McLellan and Wolf, 1999), although a direct association linking differential glutathione conjugation with platinum drug resistance in human ovarian tumours has not been convincingly demonstrated. In support of a functional association, heterologous GSTP1 expression in Saccharomyces cerevisiae (Black et al, 1990) or of various GSTP1 alleles in Escherichia Coli influenced sensitivity to platinum and additional chemotherapy drugs (Ishimoto and Ali-Osman, 2002). In contrast, similar experiments in breast cancer MCF7 cells revealed only modest effects on platinum sensitivity (Peklak-Scott et al, 2008), while partial GSTP1 knockdown in the adriamycin-resistant human colorectal cancer cell line M7609 increased sensitivity not only to the selection agent, but to cisplatin, melphalan and etoposide (Ban et al, 1996).

We therefore created novel ovarian tumour cell line models in which *GSTP1* expression is stably deleted, to investigate whether differential *GSTP1* expression directly influenced chemosensitivity to platinum drugs, and to other drugs routinely used in the treatment of ovarian cancer patients. We have used whole-genome transcriptional profiling analysis and various quantitative phenotypic assays to identify gene expression and associated phenotypic differences in parental and *GSTP1* knockdown cell lines.

MATERIALS AND METHODS

Ovarian tumour cell lines. A2780 and cisplatin-resistant derivative A2780DPP cells were obtained from ATCC (LGC Standards, Teddington, UK), via Cancer Research UK Cell Services. A2780 cells were maintained in RPMI-1640 media supplemented with 10% FBS, and A2780DPP cells (derived *in vitro* following longterm cisplatin selection; Behrens *et al*, 1987) in RPMI-1640 media supplemented with 15% FBS and $1 \,\mu$ M cisplatin. Both cell lines were maintained in 37 °C incubators, supplemented with 5% CO₂.

Creation and characterisation of stable *GSTP1* **knockdown cell lines.** Glutathione *S*-transferase P1 expression was stably knockeddown in A2780 and expression significantly reduced in A2780DPP cells by RNA interference using Mission shRNA plasmids (Sigma-Aldrich, Gillingham, UK). Five unique *GSTP1*-specific shRNA plasmids (clones TRCN0000083773, TRCN0000083774, TRCN000 0083775, TRCN0000083776 and TRCN0000083777) and a negative control plasmid (empty vector control, SHC001) were purchased as glycerol stocks and plasmid DNA extracted using plasmid DNA maxi prep kits (Qiagen, Manchester, UK) according to the manufacturer's instructions. A2780 and A2780DPP cells (2.5×10^5) cells per well in six-well plates) were transfected with each plasmid using lipofectamine (Life Technologies, Paisley, UK), and shRNA-containing cells selected with puromycin. Individual cell colonies were picked using cloning cylinders, harvested for mRNA analysis (A2780DPP cells) or expanded to 75 cm² tissue culture flasks and harvested for mRNA and protein analysis (A2780 cells). *GSTP1* knockdown in A2780 cells was confirmed by qRT–PCR analysis and western blotting, and by qRT–PCR analysis in A2780DPP cells. Cell growth rates were compared by plating 1×10^5 cells from each cell line in individual wells of six-well dishes (day 0). Cells were harvested daily by trypsinisation (days 2–10) and counted using a haemocytometer.

RNA extraction and qRT-PCR analysis. Cells were grown to 80% confluency in 75 cm² flasks, harvested by trypsinisation, counted using a haemocytometer, and 1×10^7 cells used for RNA extraction using a Qiagen RNeasy mini kit (Qiagen), following the manufacturer's protocol for mammalian cells, with additional on column DNAse digestion (RNAse free DNAse kit, Qiagen). RNA yield and integrity were initially assessed from absorbance readings at 260 and 280 nm using a Nanodrop 1000 spectrophotometer (Thermo Fisher Scientific, Loughborough, UK), and confirmed using an Agilent Bioanalyzer 2100 and RNA 6000 Nano LabChip Kit (Agilent Technologies, Wokingham, UK) according to the manufacturer's guidelines.

RNA was reverse transcribed into cDNA (50 ng RNA per 20 μ l RT reaction) using High Capacity RNA to cDNA kits (Life Technologies) according to the manufacturer's instructions, and the expression of GSTP1 (Taqman probe ID Hs00168310_m1) and the loading control 18S ribosomal RNA (Hs03003631 g1) assessed by qRT-PCR analysis, as previously described (Smith et al, 2012), where 20 μ l individual reaction mixes (per well) contained 10 μ l Taqman Universal PCR Master Mix (Life Technologies), $1 \mu l$ genespecific Taqman probe, $1 \mu l$ cDNA and $8 \mu l$ sterile water. Each reaction was performed in triplicate and run on the Standard Real-Time PCR program on a 7900 Taqman real-time PCR system (Applied Biosystems, Warrington, UK) using pre-defined thermal cycling conditions (50 °C for 2 min, 94.5 °C for 10 min, 40 cycles of (97 °C for 30 s and 59 °C for 1 min)). Similarly, the expression of ALX1 (Hs00232518_m1), CDH2 (Hs00983056_m1), FOXC1 (Hs00559473_s1), LAYN (Hs00379511_m1), TM4SF (Hs00371661_m1) and VCAN (Hs00171642_m1) was additionally investigated by qRT-PCR analysis in A2780 and A2780/GSTP1 cells. Analysis was performed using SDS 2.3 software (Applied Biosystems); optimal experimental baselines and thresholds were chosen for each gene, and gene expression quantitated by cycle threshold (Ct) values. Relative expression values were determined by comparing the expression of each target gene with the invariant 'loading control' 18S ribosomal RNA, as previously described (Smith et al, 2012). All samples were analysed in triplicate and gene expression calculated relative to 18S ribosomal RNA ± compound error ((s.d. target gene)² + (s.d. 18S ribosomal RNA)²)^{1/2}, where s.d. = standard deviation of the mean of triplicate replicates.

Whole-genome microarray mRNA analysis in A2780 and A2780/GSTP1 cells. RNA was prepared from A2780 and A2780/GSTP1 cells and integrity assessed as described above. Each RNA sample was converted to biotinylated amplified cRNA using an Illumina TotalPrep RNA Amplification kit (Life Technologies) according to the manufacturer's guidelines, and cRNA quality and concentration confirmed on the Agilent Bioanalyzer 2100, as described above. cRNA samples were hybridised in triplicate on Illumina Human HT-12 BeadChip Arrays (Illumina, Little Chesterford, UK) using standard protocols optimised by the Genetics Core, Wellcome Trust Clinical Research Facility, University of Edinburgh.

Bioinformatics analysis. Gene expression data were analysed using Bioconductor 2.7 (http://bioconductor.org), running on R 2.12.1. Normalised probeset expression measures were calculated using log₂ transformation and quantile normalisation using the 'Lumi' package. To identify significant differences in gene expression in A2780 and A2780/GSTP1 cells, moderated Student's *t*-tests were performed using empirical Bayes statistics in the 'Limma' package, and resulting *P*-values adjusted for multiple testing using the false discovery rate (FDR) Benjamini and Hochberg method (Smyth, 2004); probe sets with adjusted *P*-value FDR q < 0.05 were called differentially expressed. Differentially expressed probes were also subjected to Metacore Pathway analysis (Thomson Reuters, London, UK) to identify enrichment of pathways and processes, using hyper-geometric distributions to determine the most enriched gene sets (FDR q < 0.05).

Protein expression and western blotting. Cells for protein extraction were plated in six-well dishes, cultured until confluent, growth media removed and washed with ice-cold PBS before lysis in 0.5 ml RIPA buffer (50 mM Tris-HCl, 150 mM NaCl, 5 mM EDTA, 0.1% SDS, 0.5% sodium deoxycholate, 1% Nonidet P-40) supplemented with 2% protease inhibitor cocktail (Sigma-Aldrich). Lysed cells were centrifuged (13 000 r.p.m; 10 min) to pellet cell debris, and protein concentrations of the resulting cell supernatants determined by Bradford Assay (Bio-Rad, Hemel Hempstead, UK), relative to a standard curve prepared from serial dilutions of bovine serum albumin (0–1 mg ml⁻¹), with absorbance readings at 595 nm.

Glutathione S-transferase P1 expression was analysed in protein extracts from each cell line by western blotting, following SDS-PAGE. Each protein sample (40 μ g) was diluted in equal volumes of $5 \times$ sample buffer (10% SDS, 250 mM Tris-HCl (pH 6.8), 1 mg ml⁻¹ bromophenol blue, 0.5 м DTT, 50% glycerol), denatured and separated by SDS-PAGE using 12% Mini-PROTEAN 3 polyacrylamide gels (Bio-Rad) in Tris-glycine buffer (25 mM Tris pH 8.3, 250 mM glycine, 0.1% SDS). Following electrophoresis, proteins were transferred to nitrocellulose membranes in Trisglycine-methanol buffer (48 mM Tris pH 8.3, 39 mM glycine, 0.037% SDS, 10% methanol) and nonspecific antibody binding blocked by incubation for 2 h in TBST (25 mM Tris-HCl pH 7.6, 150 mM NaCl, 0.05% Tween-20) containing 5% milk powder. Membranes were then incubated overnight with a rabbit polyclonal GSTP1 primary antibody ((Henderson et al, 1998), diluted 1:1000) or a mouse monoclonal β -actin antibody (sc-47778, Santa Cruz Biotechnology Inc., Heidelberg, Germany, diluted 1:1000), washed in PBST (PBS supplemented with 0.05% Tween-20) and incubated for 1 h with a swine anti-rabbit polyclonal secondary antibody (Dako PO399, diluted 1:1000, GSTP1, Dako, Ely, UK) or goat anti-mouse polyclonal secondary antibody (Dako PO447, diluted 1:1000, β -actin). Immunoblots were developed using an ECLchemiluminescence kit (Merck Millipore, Watford, UK) according to the manufacturer's instructions.

Analysis of cellular glutathione levels. Total cellular glutathione (GSSG + GSH) levels were compared in A2780 and A2780/GSTP1 cells using a Glutathione Assay kit (Sigma-Aldrich) according to the manufacturer's guidelines. Cells were harvested, washed in PBS, counted using a haemocytometer and re-suspended to a final concentration of 1×10^8 cells ml⁻¹ in PBS. Cells were then pelleted, de-proteinised with 5% 5-sulfosalicyclic acid and glutathione levels assessed using a kinetic assay in which catalytically active glutathione reduces 5,5-dithiobis(2-nitrobenzoic acid) (DTNB) to 5-dithiobis(2-nitrobenzoic acid) (TNB). TNB production was measured spectrophotometrically at 412 nm, with kinetic reads at 1-min intervals for 5 min, and glutathione levels extrapolated from a standard curve of serial dilutions of reduced glutathione (positive values for both GSH and GSSG are obtained in the reaction).

Ovarian cell line chemosensitivity assays. MTT assays (Mosmann, 1983) were used to compare chemosensitivity of A2780 and A2780/GSTP1 cells to the GSTP1 model substrate ethacrynic acid and chemotherapy drugs cisplatin, carboplatin, paclitaxel, doxorubicin, topotecan and gemcitabine. Each cell line was plated in a 96-well plate (5000 cells per well) and treated in triplicate with serial dilutions of each drug. Ethacrynic acid was used at concentrations from 2.5 to 80 μ M, and chemotherapy drugs at concentrations selected to mimic typical peak plasma levels in ovarian cancer patients (range 0-200% (peak plasma); cisplatin $0-25 \,\mu\text{M}$, carboplatin $0-85 \,\mu\text{M}$, paclitaxel $0-32 \,\mu\text{M}$, doxorubicin $0-6\,\mu\text{M}$, gemcitabine $0-190\,\mu\text{M}$ and topotecan $0-56\,\mu\text{M}$; Konecny et al, 2000). Cells were drug treated for 72 h, media removed and $100 \,\mu\text{l}$ of a 0.5 mg ml⁻¹ MTT solution (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide in phenol red-free DMEM) added and cells incubated at 37 °C for 3 h. The resulting formazan crystals were solubilised in DMSO, quantitated spectrophotometrically at 570 nm and the percentage of live cells remaining following each drug treatment calculated (assigning a value of 100% to vehicletreated cells). IC50 values were calculated from log dose-response curves using Prism 6 software (GraphPad Software Inc., La Jolla, CA, USA).

Cell invasion and migration assays. Cell invasion and migration was assessed using 24-well InnoCyte Cell Invasion and Migration Assays (Merck Millipore) according to the manufacturer's guide-lines. Cell invasion was assessed by plating cells in serum-free medium in invasion chambers (8 μ m membranes) coated with basement membrane matrix, which prevents non-invasive cells from passing through the membrane pores. Similarly, cell migration was compared in A2780 and A2780/GSTP1 cells by plating each cell line in serum-free medium in migration chambers. In both assays, serum-containing medium was added to each well to induce cell migration, assessed by staining cells attached to the lower side of the membrane with the fluorescent cell permeable dye Calcein-AM, and measuring fluorescence at 485 nm (excitation) and 520 nm (emission). Negative control A2780 cells were additionally treated with the anti-migratory drug Latrunculin A.

FACS analysis. Cell cycle parameters were compared in A2780 and A2780/GSTP1 cells by flow cytometry following propidium iodide labelling of cell line DNA. Cells were untreated, or treated with 12.66 μ M cisplatin (to represent typical 100% patient peak plasma levels) for 1, 4 or 24 h then fixed in ice-cold 70% ethanol overnight at -20 °C, re-suspended in PBS and stained by incubation with propidium iodide (40 μ g ml⁻¹, Sigma-Aldrich) and RNAse A (100 μ g ml⁻¹, Sigma-Aldrich) for 30 min at 37 °C in the dark. Cell cycle parameters (10 000 cells per sample) were analysed using a FACScan flow cytometer (Becton Dickinson, Oxford, UK) and Cellquest Pro software to determine cell cycle phases and cells in sub-G₀/G₁.

Statistical analysis. All statistical tests were performed using the SPSS statistics package version 20.0 (IBM, New York, NY, USA). Independent sample *t*-tests were used to assess differences in gene expression identified by qRT–PCR analysis and invasion and migration changes in A2780 and A2780/GSTP1 cells.

RESULTS

We previously showed that *GSTP1* is abundantly expressed in human ovarian tumours, and highlighted marked inter-tumour differences in *GSTP1* expression (Smith *et al*, 2012). To investigate whether individuality in *GSTP1* expression influences chemotherapy response, we used shRNA-mediated gene silencing to stably knockdown *GSTP1* in the ovarian tumour cell line A2780, created from a chemotherapy-naive ovarian cancer patient (Behrens *et al*, 1987). A2780 cells were transfected with each of five *GSTP1*specific Mission shRNA pLKO.1-puro plasmids (773, 774, 775, 776 and 777) and a negative control plasmid as described in Materials and Methods section. Following lipofectamine selection, multiple antibiotic-resistant colonies were screened for *GSTP1* knockdown

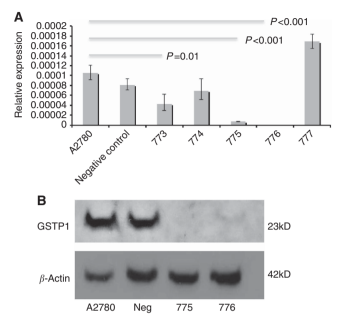


Figure 1. Generation of stable GSTP1 knockdown cell lines. A2780 ovarian tumour cells were transfected with various GSTP1 Mission shRNA constructs (clones 773, 774, 775, 776 and 777) and an empty vector negative control plasmid as described in Materials and Methods section. Following puromycin selection, GSTP1 expression was compared in A2780 cells and in each novel daughter cell line (**A**) by qRT–PCR analysis, relative to 18S ribosomal RNA (significant differences in gene expression (P<0.05) are highlighted) and (**B**) by western blot analysis of clones 775 and 776 to confirm loss of GSTP1 protein expression.

by qRT–PCR analysis, and gene knockdown confirmed in cells transfected with plasmids 773, 775 and 776 (Figure 1A), with maximum reduction in shRNA construct 775 and 776-transfected cells. *GSTP1* protein expression was evaluated by western blotting, with complete loss of *GSTP1* expression confirmed in shRNA construct 775-transfected cells and >95% reduction in protein expression in construct 776-transfected cells (Figure 1B). GSTP1-775 knockdown cells (A2780/GSTP1) were therefore selected for use in additional experiments, following additional confirmation of increased total cellular glutathione levels (A2780 83.16 ± 1.67 μ M and A2780/GSTP1 116.85 ± 4.67 μ M), consistent with reduced *GSTP1* catalytic activity.

MTT assays were then used to compare chemosensitivity of A2780 and A2780/GSTP1 cells to the diuretic and wellcharacterised GSTP1 substrate ethacrynic acid. GSTP1 conjugates and detoxifies ethacrynic acid (Ahokas et al, 1984), which was significantly more toxic to GSTP1 null cells (1.3-fold increase in IC₅₀, P = 0.004, Figure 2A). Sensitivity to platinum-based chemotherapy drugs was then compared in A2780 and A2780/GSTP1 cells and significant differences in chemosensitivity observed following treatment with cisplatin (2.28-fold decrease in IC₅₀, P = 0.03, Figure 2B) and carboplatin (4.83-fold decrease in IC₅₀, P = 0.007, Figure 2C), and a less pronounced decrease in sensitivity to paclitaxel (1.58-fold increase in IC_{50} , P = 0.02, Figure 2D). Chemosensitivity to additional drugs used to treat ovarian cancer was also compared but no significant differences in IC₅₀ values identified (doxorubicin 0.13 vs 0.11 µM, gemcitabine 3.32 vs 3.34 µM and topotecan 1.00 vs 1.12 µM in A2780 and A2780/GSTP1 cells, respectively (all *P*-values > 0.05)). To confirm our findings, MTT assays were repeated in GSTP1-776 cells and very similar changes in cisplatin, carboplatin and paclitaxel chemosensitivity identified (data not shown).

As *GSTP1* knockdown significantly increased cisplatin and carboplatin chemosensitivity in A2780 cells, we further investigated whether *GSTP1* knockdown in the related platinum-resistant A2780DPP cell line (Behrens *et al*, 1987) could re-sensitise these drug-resistant cells to platinum-based chemotherapy. A2780DPP cells were therefore transfected with shRNA constructs 775 and 776, optimised for *GSTP1* knockdown in the experiments described

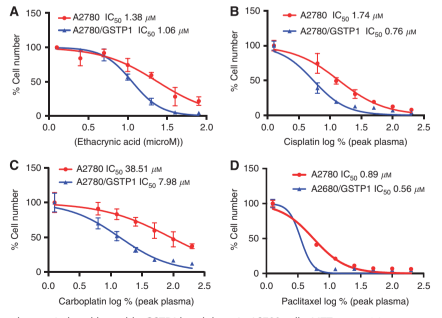


Figure 2. Chemosensitivity changes induced by stable GSTP1 knockdown in A2780 cells. MTT cytotoxicity assays were used to compare chemosensitivity to (A) ethacrynic acid, (B) cisplatin, (C) carboplatin and (D) paclitaxel in A2780 and A2780/GSTP1 cells as described in Materials and Methods section. Cisplatin, carboplatin and paclitaxel were used at doses representative of typical peak plasma levels in ovarian cancer patients. Each assay was performed in triplicate—mean data from three independent experimental replicates is illustrated.

above. In contrast to drug-sensitive A2780 cells, however, although puromycin-resistant colonies were initially formed in A2780DPP cells in multiple replicate experiments, the colonies formed were relatively small (10-20 cells per colony) and were viable for only 48 h (Figure 3A). As our control transfections resulted in viable colony formation in both A2780 and A2780DPP cells, we hypothesised that GSTP1 knockdown may be lethal in A2780DPP cells, which are grown in medium containing 1 µM cisplatin to maintain the drugresistant phenotype. To confirm GSTP1 knockdown in A2780DPP cells, we isolated individual puromycin-resistant colonies 24 h after colony formation using cloning cylinders, and extracted sufficient RNA to confirm GSTP1 knockdown by qRT-PCR analysis (Figure 3B). Unfortunately, we were unable to harvest sufficient A2780DPP/GSTP1 knockdown cells to perform western blotting, cytotoxicity assays or more detailed phenotypic characterisation, but our confirmation of GSTP1 knockdown, together with selective platinum toxicity in puromycin-resistant clones is consistent with an essential role for GSTP1 in glutathione conjugation and cisplatin detoxification. We further attempted to confirm this hypothesis by knocking down GSTP1 in A2780DPP cells grown in the absence of cisplatin selection, but found that control untransfected A2780DPP cells did not retain a stable drug-resistant phenotype over the 6-week experimental period in the absence of cisplatin selection (data not shown).

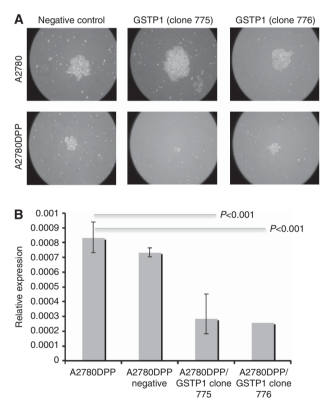


Figure 3. GSTP1 knockdown is toxic to cisplatin-resistant A2780DPP cells. A2780 and cisplatin-resistant A2780DPP cells were transfected with GSTP1 Mission shRNA plasmids 775 and 776 or an empty vector negative control plasmid, as described in Materials and Methods section, and GSTP1 knockdown cells identified by puromycin selection. (A) Viable puromycin-resistant colonies were obtained from all transfections in A2780 cells, but only from A2780DPP cells transfected with an empty vector negative control plasmid. (B) qRT–PCR analysis was used to confirm GSTP1 knockdown in A2780DPP cells (illustrated relative to the expression of 18S ribosomal RNA) following transfection of GSTP1 shRNA clones. Cells from small colonies with limited viability were collected using cloning cylinders, and RNA extracted as described in Materials and Methods section.

To investigate the cellular phenotypes resulting from *GSTP1* knockdown in A2780 cells, Illumina HT-12 Expression BeadChip Arrays were used to compare gene expression in A2780 and A2780/GSTP1 cells. Each HT-12 array contains >47 000 unique probe sets, corresponding to >28 000 coding transcripts; 2671 probes were significantly more highly expressed, and 2717 probes less highly expressed in A2780/GSTP1 cells. Of these, \geq 2-fold differences in gene expression were identified for 336 gene transcripts—the most significantly differentially expressed genes are summarised in Supplementary Table 1 (Supplementary Information), and changes in gene expression predicted by BeadChip Array analysis confirmed by qRT–PCR analysis for selected up- and downregulated genes (Figure 4). It was of particular interest to note that no compensatory changes in additional GST isoforms were identified in A2780/GSTP1 cells.

Pathway and process enrichment analysis was then used to identify common differentially regulated pathways or processes in A2780 and A2780/GSTP1 cells (Supplementary Table 2). Consistent with these predictions, the most significant differences in gene expression were seen for genes (e.g., N-cadherin (CDH2), versican (VCAN), L6 cell surface antigen (TM4SF1) and layilin (LAYN)) associated with cell invasion, migration, metastasis and the epithelial-mesenchymal transition. Consistent with known functions of the GSTs, several differentially regulated pathways and processes were associated with lipid metabolism and with the oxidative stress response, while additional interesting associations suggested altered cell cycle regulation and differences in apoptosis, cell invasion and migration. We therefore used quantitative cell growth, invasion and migration assays and FACS analysis to compare A2780 and A2780/ GSTP1 cell phenotypes. We did not observe significant differences in cell growth rates, where A2780 and A2780/GSTP1 cells had doubling times of 0.91 and 1.03 days, respectively (P = 0.462), or in cell cycle parameters, assessed by FACS analysis of propidium iodide-labelled untreated cells and stressed cells acutely treated with cisplatin (Figure 5A). In contrast, consistent with the gene expression and pathway/process differences described above, we found that both cell invasion (Figure 5B) and migration (Figure 5C) was significantly reduced in A2780/GSTP1 cells.

DISCUSSION

Conjugation of glutathione with platinum-based chemotherapy drugs is an important detoxification mechanism, which promotes drug clearance, limits the formation of DNA crosslinks and reduces toxicity (Peklak-Scott *et al*, 2008). It is therefore logical that the expression of glutathione-conjugating enzymes including *GSTP1* is increased as an adaptive response in drug-resistant tumour cells (Black and Wolf, 1991; McLellan and Wolf, 1999) and that reduced *GSTP1* activity may influence chemosensitivity. Several studies, however, have described increased *GSTP1* expression in drug-resistant cell lines following exposure to drugs which are not detoxified by glutathione conjugation (Wang *et al*, 1989), or which are not *GSTP1* substrates (Tew, 1994) a direct role for *GSTP1* in platinum chemosensitivity therefore remains to be unequivocally established.

We have shown for the first time that stable deletion of *GSTP1* in A2780 ovarian tumour cells significantly and selectively increases sensitivity to cisplatin and carboplatin, drugs routinely used to treat ovarian cancer, a notoriously drug-resistant and clinically intractable disease. Importantly, loss of *GSTP1* expression in the cisplatin-resistant A2780 subline A2780DPP was toxic to the cells in the presence of cisplatin, consistent with an important catalytic role for *GSTP1* in glutathione conjugation in the detoxification pathway of platinum drugs. Consistent with this hypothesis, we found significant differences in intracellular glutathione levels and in sensitivity to the GST substrate ethacrynic acid in *GSTP1* null cells. Although the role of *GSTP1* in ovarian

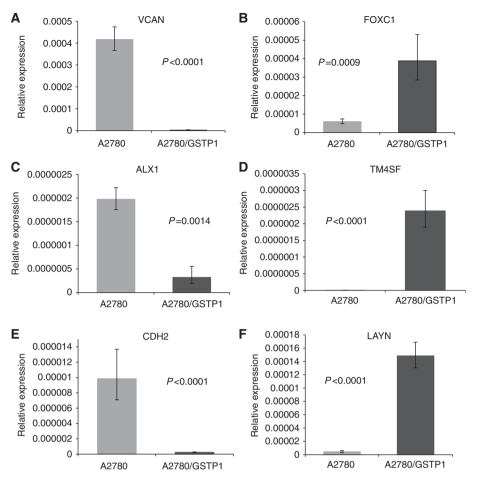


Figure 4. qRT–PCR analysis confirms gene expression changes predicted by Illumina Beadchip microarray analysis of A2780 and A2780/GSTP1 cells. qRT–PCR was performed, as described in Materials and Methods section, to confirm differential expression of selected genes predicted by comparative Illumina HT-12 Beadchip Array analysis to be differentially expressed in A2780 and A2780/GSTP1 cells—(A) VCAN, (B) FOXC1, (C) ALX1, (D) TM4SF, (E) CDH2 and (F) LAYN. All samples were analysed in triplicate; gene expression is illustrated relative to the expression of 18S ribosomal RNA.

tumour chemosensitivity has not been studied, transient siRNAmediated *GSTP1* knockdown in a diverse panel of leukaemia and lymphoma cell lines has recently also been reported to influence cisplatin sensitivity (Chen *et al*, 2013).

Glutathione S-transferase P1 has also been proposed to have a non-catalytic role in promoting cell proliferation by binding to and inhibiting JNK (Adler et al, 1999)-JNK activity is increased when GSTP1 activity is reduced, either by small molecule GSTP1 inhibitors or in GSTP1 null mice (Henderson et al, 1998). It is logical to propose, therefore, that stable GSTP1 deletion in A2780 ovarian tumour cells may result in INK activation. Consistent with this hypothesis, interrogation of our gene expression microarray data sets revealed increased expression of several JNK regulatory genes including the toxicity and JNK inducer TAJ (TNFRSF19, 8.33-fold, adjusted *P*-value 1.25×10^{-18} , Supplementary Table 1), and JNK target genes including the p53-inducible proteins TP53I3 (PIG3, 5.02-fold, adjusted P-value 1.73×10^{-16}) and CDKN1A (p21, 4.87-fold, adjusted P-value 9.62×10^{-15}) in GSTP1 knockdown cells. Although many of the genes up- and downregulated by GSTP1 deletion are associated with the RAS/MAPK pathway (Supplementary Table 1), we did not detect a significant difference in RAS/MAPK pathway activation, using a quantitative RAF-1binding/Ras GTPase ELISA to assess pathway activation in GSTP1 knockdown cells (data not shown). In contrast, several differentially expressed genes (e.g., CDH2) and associated pathways and processes suggested that GSTP1 knockdown may significantly inhibit cell invasion and migration-these predictions were

tion and the development of drug resistance. We were therefore surprised to find that GSTP1 knockdown did not significantly influence cell growth rate or alter cell cycle progression in A2780 cells, although similar findings have been reported in a series of ovarian cancer cell lines made resistant to cisplatin, carboplatin and paclitaxel (Li et al, 2004), suggesting limitations of the use of immortalised cancer cell line models. Similarly, and consistent with a previous report of comparable GSTP1 expression in A2780 cells and the cisplatin-resistant derivatives C70 and C200 (Townsend et al, 2002), we did not detect a significant increase in GSTP1 expression in platinum-resistant A2780DPP cells. In contrast, increased GSTP1 expression has been described in many drugresistant cell lines (Kuroda et al, 1991; Kotoh et al, 1997; Tozawa et al, 2008; Yang et al, 2009), including ovarian tumour lines resistant to platinum drugs (Lewis et al, 1988; Parekh and Simpkins, 1996; Yanagie et al, 2009). Recent data from our own laboratory, where we see increased GSTP1 expression in a novel drug-resistant A2780 subline immediately following de novo platinum selection, further support the hypothesis that increased GSTP1 expression may not be maintained in long-term culture of immortalised tumour cells. In light of these concerns, we are currently extending our analysis to additional immortalised cell lines, with different histologies and genetic backgrounds and to primary cell lines derived from ascites from drug-sensitive and drug-resistant ovarian cancer patients.

confirmed experimentally, and are consistent with the hypothesis

that increased GSTP1 expression promotes neoplastic transforma-

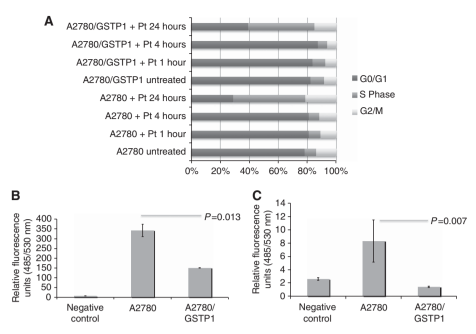


Figure 5. Phenotypic comparison of A2780 and A2780/GSTP1 knockdown cells. (A) FACS analysis of PI-labelled cells was used, as described in Materials and Methods section, to compare cell cycle progression of A2780 and A2780/GSTP1 cells, before and following cisplatin challenge. Each sample was analysed in triplicate—the relative proportions of cells in each phase of the cell cycle at each time point is illustrated. Boyden chamber-based (B) invasion (using cell culture media supplemented with 10% FBS as chemoattractant) and (C) migration (ECMatrix) assays were then used to compare predicted phenotypes in negative (Latrunculin A-treated) controls, A2780 and A2780/GSTP1 cells as described in Materials and Methods section.

Glutathione S-transferase P1 expression in cancer patients may also be influenced by the presence of allelic variants-GSTP1 Ile₁₀₅Val (rs1695) and GSTP1 Ala₁₁₄Val (rs1138272), with homozygote rare allele frequencies of approximately 12% and 2% in the Caucasian population, respectively (Zimniak et al, 1994; Ali-Osman et al, 1997; Harries et al, 1997; Sachse et al, 2002). Unlike GSTM1 and GSTT1, however, where common polymorphisms result in complete gene deletions (Board, 1981), variant GSTP1 alleles differ from the consensus reference sequence by single aminoacid substitutions, resulting in less pronounced, less frequent and less well-characterised phenotypes (Peklak-Scott et al, 2008). In contrast, however, our recent gene expression profiling experiments in human ovarian tumours describe marked (>70-fold) inter-tumour differences in GSTP1 expression, suggesting that individuality in glutathione-conjugating activity, although not genetically determined, could significantly influence response to platinum-based chemotherapy in ovarian cancer patients.

Studying the influence of inter-individual differences in GSTP1 expression on disease progression or chemotherapy response in ovarian cancer patients is challenging-the disease is frequently diagnosed at an advanced stage, limiting the availability of matched normal and tumour samples, and serial matched drugsensitive and drug-resistant tumour biopsies are rarely available. Increased *GSTP1* expression in ovarian tumours (n = 41), relative to unmatched healthy ovarian samples (n = 12) and benign tumours (n = 25) has been described in a small Chinese study (Cheng et al, 2000) and, consistent with our findings, GSTP1 expression in ovarian cyst fluid has been correlated with higher relapse rates following platinum-based chemotherapy (Boss et al, 2001). In similar studies, GSTP1 expression in ovarian cyst fluid correlated both with serum CA125 levels and reduced patient survival (Kolwijck et al, 2009), while in a small series (n = 30)of matched first and second-look laparotomies, increased GSTP1 expression was associated with disease progression, assessed by both more frequent relapses and reduced survival (Surowiak et al, 2005).

Confirmation of a direct role for GSTP1 in chemotherapy response is important, not only in the prediction of response to platinum-based chemotherapy, but as a candidate response biomarker for new generation chemotherapy drugs, designed to exploit increased GSTP1 expression in tumours relative to surrounding normal tissues. For example, TLK-286 (Telcyta, canfosfamide) was identified as the lead candidate in a rationally designed series of selectively toxic glutathione analogues (Lyttle et al, 1994), which continues to be evaluated both as a single agent and in combination chemotherapy in phase II and III clinical trials in ovarian cancer and other solid tumours. TLK-286 is metabolically activated by GSTP1 and cytotoxicity has been correlated with GSTP1 expression (Rosario et al, 2000; Dourado et al, 2013). Increased GSTP1 expression in tumours and in drug-resistant cells is also the target of a new class of GST suicide inhibitors, including 7-nitro-2,1,3-benzoxadiazole (NBDHEX) (Ricci et al, 2005), which acts to induce apoptosis by promoting dissociation of the GSTP1/ JNK1 complex in leukaemia cell lines (Turella et al, 2005). Similar effects were recently observed in mesothelioma cell lines, where NBDHEX synergised with cisplatin (De Luca et al, 2013).

We have described marked inter-tumour differences in GSTP1 expression but, unlike fibroblast growth factor (FGF) family genes, pre-treatment tumour GSTP1 expression was not influenced by tumour histology or associated with altered survival (Smith et al, 2012). Our study did not have sufficient power to perform a meaningful assessment of the potential role of GSTP1 in chemotherapy response, although we believe that lack of routine access to comparable clinical biopsy or tumour samples from ovarian cancer patients pre- and post-treatment may significantly limit tumour biomarker utility. It is interesting to note, however, that postchemotherapy GSTP1 expression was associated with progression-free survival in a small study (n = 41 patients) where pre- and postchemotherapy biopsies were available (Saip et al, 2005). We are therefore currently recruiting ovarian cancer patients to clinical studies in which we are collecting serial serum and ascites samples from matched drug-sensitive and drug-resistant patients for

quantitative biomarker profiling. GSTP1 expression has previously been investigated by immunostaining in ovarian ascites samples, and shown to correlate with both primary tumour expression and cisplatin chemosensitivity (Kase et al, 1998). Our microarray data sets provide numerous examples of GSTP1-dependent gene expression changes, and may therefore allow us to identify additional biomarkers, which correlate with GSTP1 activity. Additional candidate GSTP1 biomarkers have recently identified in studies describing an inverse association between GSTP1 expression, chemosensitivity and expression of the regulatory microRNA miR-513a-3p in cisplatin-resistant A549 lung cancer cells (Zhang et al, 2012). Similarly, interleukin-6 (IL-6) production has been shown to be elevated in both serum and ascites samples from ovarian cancer patients and to be inversely associated with platinum chemosensitivity and survival (Scambia et al, 1995). Of obvious relevance to our findings, GSTP1 has recently been shown to be an IL-6 target gene (Wang et al, 2010), while inhibition of GSTP1 activity by an IL-6 or IL-6 receptor mAb correlates with increased platinum (and paclitaxel) sensitivity in renal cancer cells (Mizutani et al, 1995). It will therefore be of particular interest to compare GSTP1 and IL-6 levels in serum and ascites samples from drug-sensitive and drug-resistant ovarian cancer patients.

In summary, and consistent with the findings of a recent similar study in mesothelioma (Chen *et al*, 2014), we have shown that *GSTP1* selectively influences sensitivity to cisplatin and carboplatin in ovarian tumour cells. Additional studies to evaluate the role of *GSTP1* and co-regulated genes as clinical response biomarkers of disease progression and platinum chemosensitivity in ovarian cancer patients are underway in our laboratory.

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