Altered stability of etoposide-induced topoisomerase II–DNA complexes in resistant human leukaemia K562 cells

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Summary K562 leukaemia cells were selected for resistance using 0.5 μM etoposide (VP-16). Cloned K/VP-5 cells were 30-fold resistant to growth inhibition by VP-16 and 5-13-fold resistant to m-AMSA, adriamycin and mitoxantrone. K/VP-5 cells did not overexpress P-glycoprotein; VP-16 accumulation was similar to that in K562 cells. VP-16-induced DNA damage was reduced in cells and nuclei from K/VP-5 cells compared with K562 cells. Topoisomerase II protein was reduced 3-7-fold and topoisomerase IIIs and topoisomerase IIb mRNAs were each reduced 3-fold in resistant cells. After drug removal, VP-16-induced DNA damage disappeared 1.7 times more rapidly and VP-16-induced DNA–topoisomerase II adducts dissociated 1.5 times more rapidly in K/VP-5 cells than in K562 cells. ATP (1 mm) was more effective in enhancing VP-16-induced DNA damage in nuclei isolated from sensitive cells than in nuclei from resistant cells. In addition, ATP (0.3–5 mm) stimulated VP-16-induced DNA–topoisomerase II adducts to a greater extent in K562 nuclei than in K/VP-5 nuclei. Taken together, these results indicate that resistance to VP-16 in a K562 subline is associated with a quantitative reduction in topoisomerase II protein and, in addition, a distinct qualitative alteration in topoisomerase II affecting the stability of drug-induced DNA–topoisomerase II complexes.

DNA topoisomerase II (topoisomerase II) is a nuclear matrix-associated DNA-binding protein responsible for transient cleavage of DNA, allowing the passage of DNA double strands through formed DNA breaks to relieve torsional stress during replication and transcription (Wang, 1985; Liu, 1989; Osheroff, 1989). Topoisomerase II also allows for separation of daughter DNA strands during mitosis and is thought to play a role in recombinational events (Wang, 1985). Topoisomerase II is a target for a number of clinically effective antineoplastic agents including m-AMSA, doxorubicin, mitoxantrone, VM-26 and VP-16 (Chen et al., 1984; Tewey et al., 1984a,b; Zwelling, 1985; Minford et al., 1986; Zhang, 1990). These drugs interfere with topoisomerase II activity by stabilising topoisomerase II/DNA binding and strand breakage, a result of blockade of the religation/rescaling reaction which follows topoisomerase II-mediated strand breakage (Chen et al., 1984; Nelson et al., 1984). Drug resistance associated with alterations in the level, activity and/or phosphorylation state of topoisomerase II has been reported in both murine and human malignant cell lines selected for resistance in the presence of topoisomerase II inhibitors (Glisson et al., 1986; Odamiri et al., 1986; Pommier et al., 1986a,b; Danks et al., 1987; Drake et al., 1987; Per et al., 1987; Davies et al., 1988; Ferguson et al., 1988; Defle et al., 1989; Harker et al., 1989; 1991; Roberts et al., 1989; Zwelling et al., 1989; Fernandes et al., 1990; Matsuo et al., 1990; Charcosset et al., 1991; Cole et al., 1991; Friche et al., 1991; Long et al., 1991; Sugawara et al., 1991; Takano et al., 1991; Webb et al., 1991; Patel & Fisher, 1993; Sullivan et al. 1993). In several resistant cell lines, quantitative reduction of topoisomerase II expression has been correlated with the level of drug resistance in the absence of alterations of drug transport and may represent the major or sole determinant for resistance (Per et al., 1987; Cole et al., 1991; Long et al., 1991; Takano et al., 1991; Webb et al., 1991). In other cell lines selected for resistance in the presence of VP-16, VM-26, anthracyclines or mitoxantrone, there is not only reduced topoisomerase II expression but also reduced drug accumulation, often (but not always) correlated with amplification of the mdrl gene and overexpression of the 150–180 kDa P-glycoprotein drug efflux pump (Ferguson et al., 1988; Defle et al., 1989; Matsuo et al., 1990; Politi et al., 1990; Friche et al., 1991; Long et al., 1991; Kamath et al., 1992; de Jong et al., 1993). Qualitative alterations in topoisomerase II activity have also been reported (Danks et al., 1988; Zwelling et al., 1989; Sullivan et al., 1989, 1993) in cell lines selected for resistance in the presence of topoisomerase II inhibitors (Gupta et al., 1983; Beran & Anderson, 1987; Danks et al., 1987; Sullivan et al., 1993). In several of these resistant human leukaemia cell lines point mutations have been identified within or near nucleotide-binding consensus sequences of the topoisomerase II gene (Bugg et al., 1991; Hinds et al., 1991; Lee et al., 1992; Danks et al., 1993; Chan et al., 1993). Point mutations have also been identified near the active-site tyrosine 804 in resistant cell topoisomerase IIIs obtained from CCRF CEM cells selected for resistance in the presence of VM-26 or VP-16 (Danks et al., 1993; Patel & Fisher, 1993). In resistant cell lines exhibiting qualitative changes in topoisomerase II activity there are neither quantitative alterations in topoisomerase II expression nor changes in drug accumulation or overexpression of P-glycoprotein. Thus, stable resistance to topoisomerase II inhibitors can be manifest as (1) altered topoisomerase II expression alone or (2) together with the P-glycoprotein multiple drug resistance phenotype or (3) as a result of a mutation(s) resulting in a qualitative change in topoisomerase II activity.

In this paper, we describe a subline of human leukaemia K562 cells selected for resistance in the presence of VP-16 which has no alterations in drug accumulation compared with sensitive K562 cells. These stably resistant K/VP-5 cells have decreased expression of topoisomerase II mRNA and protein compared with parental K562 cells. In addition to this quantitative decrease in topoisomerase II, K/VP-5 cells exhibit qualitative changes in topoisomerase II activity related to stability of drug-induced topoisomerase II–DNA complexes and alterations in nucleotide dependency for formation of topoisomerase II–DNA complexes. Thus, resistance to topoisomerase II inhibitors in this novel resistant cell line is due to both qualitative and quantitative changes at the level of topoisomerase II.

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Materials and methods

Drugs and chemicals

[U-3H]VP-16 was obtained from Moravek Biochemicals (Brea, CA, USA) and was more than 95% radiochemically pure as measured by high-performance liquid chromatography (Sinkule & Evans, 1984). [2-14C]Thymidine (53 mCi mmol\(^{-1}\)), [3-3H]thymidine, [methyl-3H]thymidine (20 Ci mmol\(^{-1}\)) and [α-32P]dCTP were obtained from New England Nuclear (Boston, MA, USA). VP-16 and VM-26 were provided by Bristol-Myers Squibb (Wallington, CT, USA). Amrascine, adriamycin and mitoxantrone were obtained from the Drug Investigational Branch of the National Cancer Institute. Vinblastine, vincristine, podophyllotoxin, camptothecin and protease K were obtained from Sigma (St Louis, MO, USA). Water-insoluble drugs were prepared in dimethylsulphoxide (DMSO) such that solvent concentrations did not exceed 1% in the culture medium or buffer after drug treatment. DMSO was also included in control flasks at equivalent levels. Protein dye reagent and agarose were purchased from BioRad Laboratories (Richmond, CA, USA). L-F. Liu, Johns Hopkins University, Baltimore, MD, USA, generously provided antitoma (ID3) against human DNA topoisomerase II as well as the DNA topoisomerase II cDNA probe (ZilN69) contained in the plasmid pC15 and the DNA topoisomerase I cDNA probe contained in the plasmid pSH15-3. F.H. Drake, SmithKline Beecham Laboratories (King of Prussia, PA, USA), kindly provided a sample of purified DNA topoisomerase II.

Cells, media and incubation conditions

Human leukaemia K562 cells were grown in suspension culture in Dulbecco’s modified Eagle medium (DMEM) plus 10% fetal calf serum and L-glutamine. L1210 cells were grown in DMEM, 10% horse serum and L-glutamine. Resistant K/VP.5 cells were grown in DMEM supplemented with 20% fetal calf serum. K/VP.5 cells were stably resistant to VP-16 and continuous exposure to K562 cells to 0.5 μM VP-16 for 1 year, after which clones were isolated by limiting dilution (Norman & Thompson, 1977). The K/VP.5 clone had a doubling time of 22 h compared with the parental K562 doubling time of 18 h. The resistant clone has been stably resistant in the absence of drug for 2 years. Cytogenetic analysis indicated that both parental and resistant cells were hyperdiploid. No homogeneously staining regions or double minute chromosomes were present in K/VP.5 cells.

Drug-induced growth inhibition and drug accumulation

Log-phase sensitive and resistant cells were adjusted to 1 × 10⁵ cells ml\(^{-1}\) and incubated with various concentrations of a number of drugs for a period of 48 h, after which cells were counted on a model ZFB Coulter counter (Coulter Electronics, Hialeah, FL, USA). The extent of growth (beyond the starting concentration of 1 × 10⁵ cells ml\(^{-1}\)) in drug-treated vs control cells was ultimately expressed as per cent inhibition of control growth. The 50% growth-inhibitory concentration for each drug in each cell line was calculated from replicate dose–response curves generated from separate experiments.

For drug accumulation studies, K562 and K/VP.5 cells were suspended in a pH 7.4 buffer (buffer L) of 110 mM sodium chloride, 5 mM potassium chloride, 1 mM magnesium chloride, 5 mM sodium dihydrogen phosphate, 25 mM 4-(2-hydroxy-ethyl)-1-piperazineneethanesulfonic acid and 10 mM glucose, at a final concentration of 0.5 × 10⁵ cells ml\(^{-1}\). Cells were stirred in specially designed flasks by revolving Teflon paddles in a 37°C water bath, as described previously (Yalowich et al., 1987). One millilitre portions of cell suspension containing [3H]VP-16 were periodically injected into ten volumes of buffer L solution at 1°C. Cell fractions were then separated by centrifugation and washed twice with 0.85% sodium chloride at 0°C. The washed pellet was drawn up into a plastic pipette tip, extruded onto a polyethylene tare and dried overnight at 70°C. The dried pellets were weighed, placed in a glass scintillation vial and dissolved in 0.25 ml of 1 M hydroxide solution for 90 min at 70°C. The digest was neutralised with 0.25 ml of 1 M hydrochloric acid; 4 ml of aqueous counting scintillant (Amersham Corp., Arlington Heights, IL, USA) was added, and radioactivity was determined by liquid scintillation counting. Results yield cellular drug content expressed as nmol per g dry weight. Intracellular water content was determined from the difference between the wet and dry weights of cell pellets minus the [4C]inulin space, as described elsewhere (Yalowich & Goldman, 1984). Molar intracellular drug concentration was then determined from the molar content of cell VP-16 and the intracellular water volume.

DNA damage assays

For DNA damage experiments, K562 and K/VP.5 cells were labelled for 48 h with [2-14C]thymidine (0.02 μCi ml\(^{-1}\)). L1210 cells were labelled for 16 h with [methyl-3H]thymidine (0.1 μCi ml\(^{-1}\)). Unlabelled thymidine was added to allow for a final thymidine concentration of 1 μM in the culture medium.

Isolated nuclei were prepared by washing 3C-labelled whole cells in an ice-cold buffer A (pH 6.4) containing 1 mM potassium dihydrogen phosphate, 5 mM magnesium chloride, 150 mM sodium chloride and 1 mM EGTA (Filipski & Kohn, 1982). The cells were resuspended in 1 ml of this buffer, and an additional 9 ml of buffer A containing 0.3% Triton X-100 was added to lyse the cells. After incubation on ice for 30 min, 40 ml of buffer A was added, and nuclei were pelleted by centrifugation at 1,000 r.p.m. for 10 min in an IEC model HN-SII tabletop centrifuge. Nuclei density was adjusted to 1 × 10⁵ ml⁻¹ in cold buffer. After warming at 37°C for 15 min, the nuclei were treated with VP-16 and other agents in the presence or absence of 1 mM ATP and processed for measurement of DNA damage as described below.

Drug-mediated DNA damage was assessed using the alkaline elution technique for high-frequency single-strand breaks (Kohn et al., 1976). Intact K562 and K/VP.5 cells previously labelled with [2-14C]thymidine were suspended at 5 × 10⁵ cells ml⁻¹ in buffer A. These cells were incubated with various concentrations of VP-16 and other agents for 30 min at 37°C. L1210 cells (5 × 10⁵) containing [3H]DNA which received 1,500 rad irradiation were added as internal standards to 7.5 × 10⁵ drug-treated K562 cells containing [4C]DNA. After two washings in cold buffer A, cells were layered onto a polyvinyl chloride filter (pore size 0.8 μm; Gelman Sciences, Ann Arbor, MI, USA) and lysed with a solution of 2% sodium dodecyl sulphate, 10 mM disodium EDTA and 0.5 mg ml⁻¹ proteinase K. The DNA was eluted from the filter with tetrapropylammonium hydroxide, pH 12.1. The elution flow rate was 0.16 ml min⁻¹, with a fractional interval of 5 min. Cells containing [3H]DNA were irradiated on ice with a 100 Ci source (Mark Irradiator; J.L. Sheppard and Associates, Glendale, CA, USA). The yield of VP-16 and other drug-induced DNA single-strand breaks (SSBs) was quantitated as the fraction of [4C]DNA remaining on the filter when either 60% or 75% of the 3H-labelled internal standard DNA remains. A calibration curve for relating the frequency of VP-16-induced DNA SSBs to a corresponding effect of radiation (radiation equivalent damage) using 4C-labelled cells was obtained by plotting radii vs [4C]DNA retention at 60% or 75% retention of the [3H]DNA internal standard.

Determination of DNA–protein cross-link (DPC) frequency was accomplished as previously described (Ross et al., 1979). Aliquots of [3H]thymidine-labelled K562 and K/VP.5 cells (5 × 10⁵ ml⁻¹) or nuclei (1 × 10⁵ ml⁻¹), drug-treated or controls, were irradiated on ice with 3,000 rad prior to elution, as were internal control [3H]-labelled L1210 cells. Cells or nuclei were layered onto polyvinylchloride–acrylic co-polymer filters (Metrical DM-800; Gelman Sciences) along with 7.5 × 10⁵ internal control cells, and
lysed with 5 ml of a solution of 0.2% Sarkosyl, 2 M sodium chloride, 0.04 M disodium EDTA, pH 10, which was allowed to flow through the filters by gravity, as was a 3 ml addition of 0.04 M disodium EDTA, pH 10. DNA was eluted with tetrapropylammonium hydroxide, pH 12.1, at a flow rate of 0.035 ml min⁻¹. Fractions were collected at 90 min intervals for 12 h. DPC frequencies were calculated according to the bound-to-one terminus model (Ross et al., 1979).

\[
\text{DPC (rad equivalents)} = \frac{(1 - r)^{-1} - (1 - r)_{PD}^{-1}}{r_{PD}}
\]

where \( r \) and \( r_P \) are the fractional filter retentions of DNA extrapolated to zero time from \( [\text{H}] \) thymidine-labelled control and drug-treated cells (or nuclei) respectively and \( r_P \) is the radiation dose administered (3,000 rad). The greater the DNA retention of drug-treated cells relative to the controls, the greater the DPC frequency.

**DNA topoisomerase II–DNA covalent complex formation assay—intact cells and nuclei**

Mid-log cells (1.5–2.0 x 10⁶ cells ml⁻¹) were labelled overnight with 0.5 pCi ml⁻¹ [methyl-³H] thymidine (0.5 Ci mmol⁻¹) and 0.1 μCi ml⁻¹ [U-³H]jeuneine (318 mCi mmol⁻¹) in DMEM containing 5% FBS. Cells were then pelleted and resuspended in fresh DMEM/5% FBS and incubated for 1 h at 37°C. For experiments measuring the stability of topoisomerase II–DNA covalent complexes after VP-16 removal, cells were washed, resuspended in buffer L and equilibrated to 37°C. VP-16 was added to 20–200 μM and incubation continued for an additional 15 min at 37°C. Cells were pelleted for 90 s at 2,400 × g, washed with ice-cold buffer L and resuspended in 1 x 10⁶ cells ml⁻¹ in L buffer at 37°C. At selected intervals, 1 x 10⁶ cells were removed and added to 10 ml of ice-cold PBS and held on ice until all time points were collected. Cells were pelleted, lysed, cellular DNA sheared and protein–DNA complexes precipitated with SDS and potassium chloride as described by Zwelling et al. (1989). For experiments examining the effect of ATP on VP-16-induced stabilisation of topoisomerase II–DNA cleavable complexes, nuclei were isolated from cells as described above and adjusted to 2 x 10⁶ nuclei ml⁻¹ in buffer A (pH 7.4) containing 0.5 mM ATP. After prewarming at 37°C for 5 min, 200 μM VP-16 or 0.4% DMSO (control) was added and incubation continued at 37°C for an additional 15 min. Nuclei were pelleted, lysed, cellular DNA sheared and complexes precipitated with SDS–potassium chloride exactly as for whole cells.

**Topoisomerase II catalytic activity**

Topoisomerase II-containing extracts of nuclei were prepared from 1–2 x 10⁶ K562 and K/VP.5 cells as previously described (Danks et al., 1988). When the aqueous volume contributed by the nuclei is taken into consideration, the final sodium chloride concentration of the nuclear extracts varied between preparations from 0.7 to 0.85 M. 

**Western blots of cell lysates**

Lysates from 2–5 x 10⁶ K562 or K/VP.5 cells were prepared by the addition of an equal volume of 2 x gel loading buffer (0.1 M Tris–HCl, pH 6.8, 20% glycerol, 2% SDS, 0.5 M β-mercaptoethanol). Lysates were boiled for 5 min, sonicated to reduce viscosity and protein content determined using the BioRad protein assay. Samples containing 10 μg were electrophoresed through 6% SDS–polyacrylamide gel (Laemmli, 1970) and electroblotted at 400 mA overnight to nitrocellulose (Towbin et al., 1979) using a Hoefer (San Francisco, CA, USA) Transfot apparatus. Blots were incubated with human topoisomerase II-specific rabbit antiserum, IIG (from L.F. Liu), affinity-purified topoisomerase IIG-specific antibody FHD22 (Webb et al., 1993) or affinity-purified topoisomerase IIG-specific antibody FHD21 (Chung et al., 1989). The affinity-purified, isofrom-specific antibodies were provided by F.H. Drake (SmithKline Beecham). Bound antibodies were detected using alkaline phosphatase-conjugated goat-anti-rabbit IgG and the ProteinA Blot AP System of Promega Biotech (Madison, WI, USA). Levels of topoisomerase II were quantitated by scanning positive films of photographed blots using a Visage 110 image analyser (Ann Arbor, MI, USA).

**Southern blot analysis of cellular RNAs**

Total cellular RNA was isolated from 1–2 x 10⁶ K562 and K/VP.5 cells (Chomczynski & Sacchi, 1987) RNA (10 μg) was electrophoresed through 1.2% agarose gels containing 0.3 M formaldehyde, 40 mM 3(N-morpholino) propane sulfonic acid (MOPS) pH 7.0, 10 mM sodium acetate trihydrate, 1 mM EDTA, transferred to Nytran (Schleicher & Schuell, Keene, NH, USA) by capillary blotting overnight, baked for 2 h at 80°C and hybridised using standard techniques (Maniatis et al., 1982). Specific RNAs were detected by autoradiography at ~70°C using intensifying screens and Kodak XAR-5 X-ray film.

**Probe synthesis and labelling**

The human topoisomerase IIG cDNA probe (ZII69) contained in the plasmid pC15 was provided by L.F. Liu. A human topoisomerase IIG cDNA probe, synthesised by polymerase chain reaction (PCR) amplification using topoisomerase IIG-specific primers, was provided by D.P. Suttle (St Jude’s Children’s Research Hospital, Memphis, TN, USA). The β-globin cDNA probe in plasmid pHb2 was provided by K.B. Tan. Cloned inserts were removed from plasmids pC15 and pHb2 by digestion with restriction enzymes and isolation on agarose gels. cDNA fragments were used as templates for probe synthesis using Klenow fragment, [γ-³²P]dCTP, (3,000 Ci mmol⁻¹) and random primer initiation (Feinberg & Vogelstein, 1983).

**Results**

**Drug accumulation and growth-inhibitory effects in sensitive and resistant cells**

The K/VP.5 subline is 30-fold resistant to the 48 h growth-inhibitory effects of the selecting agent VP-16, and equally cross-resistant to VM-26 (Table I). In addition, there was lower (5- to 13-fold) cross-resistance to m-AMSA, adriamycin and mitoxantrone, three other agents that are known to inhibit topoisomerase II. K/VP.5 cells cultured in the absence of VP-16 have maintained this level of resistance for 2 years. In contrast, K/VP.5 cells were not cross-resistant to the microtubule inhibitors vincristine, vinblastine and podophyllotoxin, the antimetabolite arabinosyl cytosine or to the DNA topoisomerase I inhibitor camptothecin (Table I). Figure 1 indicates that the steady-state concentrations of VP-16 in K/VP.5 cells were slightly higher than in parental K562 cells when cells were exposed to 2.5–20 μM [HIJVP-16.
Table 1  Cross-resistance of K/VP.5 cells to anti-cancer agents

| Anti-cancer agent | IC_{50} values (nm) | Relative resistance |
|-------------------|---------------------|---------------------|
|                   | K562                | K/VP.5              |                     |
| VP-16             | 121 ± 48 (21)^f     | 3640 ± 1617 (21)    | 30                  |
| VM-26             | 10.3 ± 0.5 (4)      | 369 ± 63 (4)        | 36                  |
| Amsacrine         | 41.4 ± 22.3 (8)     | 524 ± 240 (8)       | 12.7                |
| Adriamycin        | 38.6 ± 10.8 (7)     | 178 ± 71 (8)        | 4.8                 |
| Mitoxantrone      | 14.7 ± 4.8 (5)      | 66.0 ± 21.4 (5)     | 4.5                 |
| Camptothecin      | 5.2 ± 0.9 (4)       | 5.9 ± 1.5 (4)       | 1.1                 |
| Vinblastine       | 8.2 ± 5.2 (9)       | 5.7 ± 2.8 (13)      | 0.7                 |
| Vincristine       | 5.8 ± 2.7 (7)       | 4.1 ± 1.1 (5)       | 0.7                 |
| Podophyllotoxin   | 7.6 ± 7.9 (7)       | 9.1 ± 4.4 (4)       | 1.2                 |
| ARA-C             | 4.7 ± 0.2 (3)       | 6.8 ± 1.5 (3)       | 1.4                 |

*50% inhibitory concentration in a 48 h growth inhibition assay. *IC_{50} of K/VP.5 cells divided by that of the parental K562 cell line. *Mean ± s.d.; numbers in parentheses, numbers of experiments run on different days.

These results are consistent with those previously reported (Beran & Anderson, 1987), in which accumulation of mAMSA was slightly increased in HL-60 cells selected for resistance in the presence of mAMSA. Accounting for intracellular water (Yalowich & Goldman, 1984), which was 5.19 ± 0.74 and 5.37 ± 0.13 ml g^{-1} dry weight for K/VP.5 and K562 cells respectively, the intracellular VP-16 concentration was 22.2 μM and 30.2 μM for K562 and K/VP.5 cells respectively, when the intracellular VP-16 concentration was 20 μM. These results demonstrate that VP-16 does not concentrate extensively within cells and suggests that decreased drug accumulation does not play a role in resistance to VP-16 in K/VP.5 cells. In addition, averaging results from 4–7 separate experiments, there were no significant differences in VP-16 unidirectional efflux in sensitive and resistant cells loaded to steady-state drug concentrations in the presence of 5 μM [3H]VP-16 (t_{1/2} = 103 ± 12 s and 108 ± 2 s for K562 and K/VP.5 cells respectively) using methods previously reported (Yalowich, 1987). Finally, using the C219 monoclonal antibody, there was no overexpression of P-glycoprotein (180 kDa) in cell membranes from K/VP.5 compared with K562 cells (M. Meyers & J.C. Yalowich, unpublished data).

**Drug-Induced DNA single-strand breaks and DNA–protein cross-linking**

DNA single-strand break frequency was reduced in K/VP.5 cells compared with parental K562 cells when these two lines were incubated with varying concentrations of VP-16 (Figure 2). A similar reduction in single-strand breaks was observed in resistant cells in the presence of mAMSA and adriamycin (not shown). VP-16 (10–100 μM)-induced single-strand breaks were also reduced in isolated K/VP.5 nuclei compared with K562 nuclei (results not shown). In three separate experiments in which nuclei were incubated with 10 μM mAMSA, the single-strand break frequency was 1857 ± 238 and 671 ± 46 radiation equivalents (mean ± s.e.) for K562 and K/VP.5 nuclei respectively. When the DNA topoisomerase I inhibitor camptothecin (0.1–1.0 μM) was added to isolated nuclei from K562 and K/VP.5 cells, there was no difference in drug-induced DNA strand break frequency (not shown). Finally, DNA–protein cross-linking (DPC) induced by VP-16, VM-26 and mAMSA was decreased in K/VP.5 compared with K562 cells (Figure 3).

**Topoisomerase II catalytic activity**

Decatentation activity of nuclear extract topoisomerase II (0–2 μg) isolated from K562 and K/VP.5 cells indicated that topoisomerase II catalytic activity was reduced 5- to 7-fold in resistant compared with sensitive cells as measured by the amount of protein required for 50% decatentation (Figure 4). Since decreased topoisomerase II levels could account for reduced topoisomerase II catalytic activity, the next set of experiments examined topoisomerase II expression in K562 and K/VP.5 cells.

**Topoisomerase II levels: Western blot analysis**

The amount of topoisomerase II in whole-cell lysates and in 0.8 M sodium chloride nuclear extracts from K562 and K/VP.5 cells was determined by Western blotting utilising rabbit IIB3 antisera to human DNA topoisomerase II (Figure 5). Two bands were observed in the lane containing purified topoisomerase II from P388 leukaemia cells (Drake et al., 1987) as well as in lanes corresponding to K/VP.5 nuclear extracts. The molecular weights of these bands were 170 ± 3 kDa and 179 ± 3 kDa (mean ± s.e.; n = 5), corresponding to previously characterised topoisomerase IIα and topoisomerase IIβ isoforms of this enzyme respectively (Chung et al., 1989; Drake et al., 1989). A pronounced reduction in the level of topoisomerase II in the lower molecular weight band (topoisomerase IIα) in both whole-cell lysates and nuclear extracts was observed in resistant K/VP.5 cells. Levels of topoisomerase IIα were quantitated by densitometric scanning of immunolabelled 170 kDa bands from

![Figure 1](image_url)  
**Figure 1**  Steady-state cellular concentrations of VP-16 in K562 (○) and K/VP.5 (■) cells exposed to 2.5–20 μM [3H]VP-16. Steady-state concentrations of VP-16 were achieved within 10–15 min, regardless of extracellular VP-16 concentration. For each experiment, 3–5 measurements at steady state were obtained at each drug concentration. Points, mean of three separate experiments; bars, s.d.
whole-cell lysates or nuclear extracts. The topoisomerase II signals (area under scanned peak) obtained with various amounts of K562 protein served as a standard curve from which signals for K/VP.5 were quantitated. Using whole-cell lysates, we observed a 7.8 ± 2.3-fold reduction in the level of topoisomerase II (M, 170,000) in K/VP.5 compared with K562 cells (mean ± s.e., from three separate lysoate preparations). In nuclear extracts there was a 4.5-fold decrease in the level of topoisomerase II in K/VP.5 compared with K562 cells (not shown). The higher molecular weight band (topoisomerase IIβ) showed much less staining in both sensitive and resistant cell lysates and nuclear extracts and could not be accurately quantitated. However, we independently quantitated levels of each topoisomerase II isoform by probing Western blots (not shown) of whole-cell lysates with polyclonal antibodies that specifically recognize either topoisomerase IIα (antibody FHD22) or topoisomerase IIβ (antibody FHD21). Results from three independent blots showed that topoisomerase IIα and topoisomerase IIβ were reduced 5.6 ± 1.6-fold and 2.7 ± 0.1-fold respectively (mean ± s.e.) in K/VP.5 compared with K562 cells.

Figure 2 DNA single-strand breaks in K562 (O) and K/VP.5 (●) cells treated for 30 min in the presence of various concentrations of VP-16. A calibration curve for relating the frequency of VP-16-induced single-strand breaks to a corresponding effect of radiation (radiation equivalent DNA damage) was obtained by plotting: rads vs. [3H]DNA retention at 75% retention of the [3H]DNA internal standard. Points, mean of 5–10 separate experiments; bars, s.e.

Figure 3 DNA–protein cross-linking produced by VP-16, VM-26, and mAMSA in K562 (open bars) and K/VP.5 (shaded bars) cells. Cells were treated with drugs at 37°C for 30 min at the indicated concentrations. DNA–protein cross-linking was quantitated by alkaline elution from polyvinyl chloride filters (0.8 μm) and expressed as radiation equivalents (see Materials and methods). Columns, mean of six separate experiments; bars, s.e.

Figure 4 Decatenation of kinetoplast DNA as a function of nuclear extract topoisomerase II. Salt (0.8 M)-extracted topoisomerase II from the nuclei of K562 and K/VP.5 cells was incubated with 1 μg of [14C]labelled kinetoplast DNA in the presence of 1 mM ATP for 30 min at 30°C. Decatenation of DNA was measured subsequent to centrifugal separation of catenated from decatenated DNA as described in Materials and methods.
t-test). Since the efflux rate of VP-16 from K562 and K/VP.5 cells does not differ ($t_{1/2} \approx 2$ min for both lines, see above), and since VP-16 has been considered a non-intercalating drug that does not bind directly to DNA, the more rapid reversal of VP-16-induced DNA damage in K/VP.5 cells compared with K562 cells suggests that resistant cells contain an altered binding site(s) for VP-16: either an altered form of topoisomerase II and/or an altered modulator of topoisomerase II activity. In addition, VP-16 efflux from both cell lines ($t_{1/2} \approx 2$ min) is more rapid than the rate of DNA damage reversal from either cell line ($t_{1/2} = 9.7-16.1$ min), indicating that VP-16 transport is not rate limiting to the process of reversal of VP-16-induced DNA damage. Hence, these data suggest that altered stability of VP-16-induced topoisomerase II–DNA covalent complexes in resistant cells leads to more rapid reversal of DNA damage.

The specificity of VP-16-induced DNA damage and its more rapid reversal in K/VP.5 compared with K562 cells was revealed by experiments in which equivalent single-strand breaks were introduced into the DNA of K562 and K/VP.5 cells by exposure of cells at 4°C to 1,500 rad of gamma-irradiation. There was no difference in the rate of reversal of DNA damage when cells were warmed to 37°C. In three experiments the $t_{1/2}$ for reversal of radiation-induced DNA damage was 11.1 ± 1.7 and 10.1 ± 2.2 min (mean ± s.e.) in K562 and K/VP.5 cells respectively. These results again are consistent with the idea that VP-16-induced DNA damage and its more rapid reversal in K/VP.5 cells is determined by interaction with an intracellular target that is altered in these resistant cells.

**Stability of VP-16-induced topoisomerase II–DNA complexes**

The stability of VP-16-induced topoisomerase II–DNA complexes was measured subsequent to a 15 min incubation of K562 cells with 20 μM VP-16 and of K/VP.5 cells with 200 μM VP-16. At these VP-16 concentrations, the steady-state level of topoisomerase II–DNA complex was similar (within 15%) in sensitive vs resistant cells. After suspension of cells in drug-free buffer, there was a more rapid dissociation of the topoisomerase II–DNA complex in resistant compared with sensitive cells (Figure 8). In eight separate paired experiments, the half-life for reversal of covalent complexes of topoisomerase II–DNA in K/VP.5 cells averaged 6.2 ± 0.3 min as compared with 9.0 ± 0.6 min for K562 cells (mean ± s.e.; $P = 0.003$, Student’s paired t-test). Thus, a significant reduction in stability of VP-16-induced topoisomerase II binding to DNA may be a factor in decreased DNA damage and/or a more rapid rate of recovery from DNA damage in resistant K/VP.5 cells.

![Figure 5](image-url)  
**Figure 5** Topoisomerase II protein levels from K562 and K/VP.5 cells. PBS-washed cells were lysed, sonicated, electrophoresed through SDS–polyacrylamide gels, and electroblotted to nitrocellulose. The blot was labelled first with topoisomerase II-specific polyclonal antisera then alkaline phosphatase-conjugated goat anti-rabbit antibody as described in Materials and methods. Shown are duplicate loadings of 10 μg of protein from lysates of K562 and K/VP.5 cells (corresponding to lysates from c. 1.5 × 10⁶ cells). The last lane contains purified topoisomerase II from P388 cells. Numbers to the right of the figures are molecular size markers (in kDa).

![Figure 6](image-url)  
**Figure 6** Topoisomerase IIa mRNA levels in K562 and K/VP.5 cells. RNA was purified from mid-log phase cells and 10 μg was electrophoresed through formaldehyde-containing agarose gels. After transfer to a nylon membrane, RNAs were hybridised to 3²P-labelled topoisomerase IIa and β₂-microglobulin cDNA probes, and autoradiographed as described in Materials and methods. Topoisomerase IIa mRNA signals were corrected for β₂-microglobulin levels. Averaging results from three separate blots, there was a 2.91- ± 0.95-fold decrease in topoisomerase IIa mRNA in K/VP.5 compared with K562 cells (mean ± s.d.; $P < 0.02$, Wilcoxon’s signed-ranks test).

![Figure 7](image-url)  
**Figure 7** Topoisomerase IIβ mRNA levels in K562 and K/VP.5 cells. RNA was purified, electrophoresed, immobilised and hybridised to 3²P-labelled topoisomerase IIβ and β₂-microglobulin cDNA probes, and autoradiographed exactly as for Figure 7. Topoisomerase IIβ mRNA signals were corrected for β₂-microglobulin levels. Averaging results from three separate blots, there was a 2.87- ± 0.3-fold decrease in topoisomerase IIβ mRNA in K/VP.5 compared with K562 cells (mean ± s.d.; $P < 0.01$, Wilcoxon’s signed-ranks test).
Table II Effects of ATP (1 mM) on VP-16-induced single-strand break frequency and DNA-protein cross-linking in sensitive K562 and resistant K/VP.5 nuclei

| Type of DNA lesion       | Experimental condition | K562 (25 μM VP-16) | K/VP.5 (100 μM VP-16) |
|--------------------------|------------------------|--------------------|-----------------------|
| Single-strand breaks (SSBs) | Control               | 476 ± 46 (9)       | 442 ± 30 (9)           |
|                          | + ATP                  | 1322 ± 86 (11)     | [277] ± 20 (4)         |
| DNA–protein cross-links (DPCs) | Control               | 497 ± 28 (5)       | 301 ± 47 (5)           |
|                          | + ATP                  | 1136 ± 96 (5)      | [229] ± 83 (5)         |

*Nuclei were incubated with the indicated concentrations of VP-16 for 30 min in the presence or absence of 1 mM ATP, following which alkaline elution methodology was used for quantitation of radiation equivalent SSBs and DPCs as described in Materials and methods. *Data are expressed as mean ± S.E. Numbers in parenthesis indicate the number of individual experiments run on different days. **Numbers in brackets represent the percentage increase in SSBs in the presence of ATP. The percentage increase in K562 nuclei is significantly greater than in K/VP.5 nuclei (*P = 0.0001, Student’s t-test adjusted for non-equal sample variances). *Numbers in brackets represent the percentage increase in DPCs in the presence of ATP. The percentage increase in K562 nuclei is significantly greater than in K/VP.5 nuclei (*P = 0.039, Student’s paired t-test).

Figure 8 Stability of topoisomerase II–DNA covalent complexes in VP-16-treated K562 and K/VP.5 cells. Cells were prelabelled with [methyl-3H]thymidine and [U-14C]leucine, and exposed to 20 μM VP-16 (K562) or 200 μM VP-16 (K/VP.5) for 15 min. Cells were washed free of drug, and at the indicated time points (10 s to 30 min) aliquots were removed and potassium chloride–SDS-precipitable complexes isolated as described in Materials and methods. Results were normalised to 14C counts as an internal standard for cell number, and are expressed relative to potassium chloride–SDS-precipitable 3H counts recovered at the end of VP-16 treatment (0 min).

Nucleotide-dependent VP-16-induced formation of single-strand breaks, DNA–protein cross-links and topoisomerase II–DNA complexes

ATP (1 mM) stimulates VP-16 (25 μM)-induced SSBs almost 3-fold in nuclei isolated from K562 cells (Table II). In contrast, when the VP-16 concentration was increased to 100 μM to achieve a similar SSB frequency in K/VP.5 nuclei, ATP enhanced drug-induced DNA damage less than 2-fold. Similarly, ATP-mediated enhancement of VP-16-induced DNA–protein cross-linking was significantly less in resistant K/VP.5 than in sensitive K562 nuclei (Table II). Since topoisomerase II catalytic activity is dependent on binding (but not hydrolysis) of ATP (Osheroff, 1989), and since ATP is known to stimulate VP-16-, m-AMSA-, ellipticine- and 5-iminoaunorobicin-induced DNA damage in isolated nuclei from L1210 cells (Glisson et al., 1984; Pommier et al., 1984), our results are consistent with the hypothesis that there is an altered functional interaction of ATP with topoisomerase II in resistant K/VP.5 cells. ATP stimulation of VP-16 (200 μM)-induced topoisomerase II–DNA covalent complexes was less in K/VP.5 than in K562 nuclei (Figure 9). At 1 mM ATP, covalent complex formation was increased 8-fold for K562 cells but only 2-fold for K/VP.5 cells. In addition, using a non-hydrolysable form of ATP, 5′-adenylyl-imidodiphosphate (1 mM), there was less stimulation of VP-16-induced topoisomerase II–DNA complex formation in K/VP.5 nuclei (1.7–2.0-fold) than in K562 (3.2±0.4-fold) nuclei (mean ± S.E., *P < 0.05; Student’s paired t-test; data not shown from four separate experiments). These data further support a qualitative alteration in resistant cell topoisomerase II affecting nucleotide-dependent VP-16-induced stabilisation of topoisomerase II–DNA complexes.

Topoisomerase II catalytic activity

After normalising for the difference in topoisomerase II protein in nuclear extracts obtained from sensitive and resistant cells, 2-fold greater ATP concentration was required for 50% deacatenation of 3H-labelled kinetoplast DNA using nuclear extract topoisomerase II from K/VP.5 compared with K562 cells (Figure 10). When ATP concentration was fixed at 1 mM, there was no significant difference in VP-16-induced
inhibition of catalytic decatenation using nuclear extract preparations from sensitive and resistant cells (Figure 11).

Discussion

The results presented in this study indicate that resistance to VP-16 and cross-resistance to other topoisomerase II inhibitors in K/VP.5 cells is associated with alterations in both the levels and the drug-induced DNA-binding activity of topoisomerase II. Drug-induced DNA strand breaks, DNA–protein cross-links and topoisomerase II catalytic activity were reduced in K/VP.5 compared with K562 cells and nuclei (Figures 2–4); these results correlate with the reduction in levels of topoisomerase II protein (Figure 5) and parallel the reduced levels of topoisomerase II mRNA (Figure 6) in resistant compared with sensitive cells. Together these results indicate a quantitative reduction in topoisomerase II expression in K/VP.5 cells. Previously, we performed topoisomerase II immunoblot depletion experiments which demonstrated that topoisomerase II from resistant cells was not bound to DNA at VP-16 concentrations which did induce topoisomerase II–DNA binding in sensitive cells (Ritke & Yalowich, 1993). Similar topoisomerase II immunoblot depletion experiments demonstrated qualitative changes in topoisomerase II in a VP-16-resistant small-cell lung cancer cell line which also exhibited quantitative topoisomerase II alterations (Mirski et al., 1993). Together, these results suggested qualitative alterations in topoisomerase II in resistant K/VP.5 cells.

In order to better characterise qualitative alterations in resistant cell topoisomerase II function in the present study, we have used different VP-16 concentrations to achieve similar levels of DNA damage or topoisomerase II–DNA binding in sensitive vs resistant cells. In this manner we have ‘normalised’ for differences in topoisomerase II protein in K562 and K/VP.5 cells. Under these experimental conditions, the more rapid reversal of drug-induced DNA damage in resistant K/VP.5 cells compared with K562 cells and the attenuated effect of ATP on the stimulation of VP-16-induced DNA strand breaks and DPC in nuclei from resistant cells (Table II) strongly suggests that, in addition to the quantitative changes reflected by reduced topoisomerase II levels, there are qualitative alterations in topoisomerase II function in K/VP.5 cells. This conclusion is further supported by measurement of the stability of VP-16-induced topoisomerase II–DNA covalent complexes which was reduced in K/VP.5 cells (Figure 8) and by the demonstration of a 2-fold increase in ATP requirement for drug-induced K/VP.5 cell topoisomerase II–DNA binding and subsequent catalytic activity (Figures 9 and 10).

The selection technique used to obtain K/VP.5 cells, intermittent then continuous low-concentration (0.5 μM) exposure to VP-16, may relate to the dual phenotypic changes in this cell line, i.e. quantitative and qualitative topoisomerase II alterations. Multifactorial resistance characteristics have been reported using intermittent topoisomerase II inhibitor exposure (Long et al., 1991; Sugawara et al., 1991) or stepwise increases in drug exposure (Ferguson et al., 1988; Matsuo et al., 1990; de Jong et al., 1993) to select for resistant cell lines. However, unlike those studies, the multifactorial resistance reported here for K/VP.5 cells does not include reduced intracellular drug accumulation. More discreet mutational or regulatory changes in topoisomerase II have been observed in cells treated with relatively high concentrations of topoisomerase II inhibitory drug or with mutagens (Bugg et al., 1991; Hinds et al., 1991; Lee et al., 1992; Chan et al., 1993; Danks 1993). Even though our selection procedure used relatively low concentrations of VP-16 (0.5 μM), the fact that K/VP.5 cells have retained 30-fold resistance to VP-16 in the absence of drug for more than 2 years suggests that a stable mutational alteration has occurred during the acquisition of resistance.

At least two targets for mutation which are not mutually exclusive may be considered as sources for the quantitative and qualitative changes of topoisomerase II observed in the K/VP.5 cell line. First, chronic VP-16 exposure may have selected for an alteration in the topoisomerase II gene itself. A mutation of the primary sequence could produce a less stable mRNA, although the published nucleotide sequence for topoisomerase II (Tai & Pflugfelder et al., 1988) reveals no known consensus mRNA stability determinants (Cleveland & Yen, 1989). Alternatively, a mutation in the topoisomerase II gene may also result in a RNA that is less efficiently transcribed. The consequence of such a mutation would be a reduction in topoisomerase II mRNA and decreased translation. Previously, we reported a reduction in the stability of topoisomerase II mRNA in K/VP.5 cells that parallels the reduction of topoisomerase II mRNA levels (Ritke &

Figure 10 Effects of ATP on nuclear extract topoisomerase II decatenation of kinetoplast DNA. Sodium chloride (0.8 M)-extracted topoisomerase II from the nuclei of K562 and K/VP.5 cells was incubated with 1 μg of [3H]-labelled kinetoplast DNA in the presence of 0–1 mM ATP for 30 min at 30°C. Decatenation of kinetoplast DNA was measured subsequent to centrifugal separation of catenated from decatenated DNA as described in Materials and methods. In order to normalise for the 5-fold differences in topoisomerase II levels in nuclear extracts from these two cell lines, 0.36 μg and 1.8 μg of nuclear extract protein was used for K562 and K/VP.5 cells respectively. Points represent the mean for 3–6 separate experiments; bars, s.e. 

Figure 11 Inhibition of topoisomerase II catalytic decatenation activity by VP-16. Nuclear extract topoisomerase II from the nuclei of K562 and K/VP.5 cells was incubated with 1 μg of [3H]-labelled kinetoplast DNA in the presence of 1 mM ATP and 0–100 μM VP-16 for 30 min at 30°C. Decatenation of kinetoplast DNA was measured as described in Figure 10 and Materials and methods. Nuclear extract protein content was normalised for differences in topoisomerase II content in K562 and K/VP.5 cells. Inhibition of decatenation is expressed relative to decatenation activity observed in the absence of VP-16. Points represent the mean of four separate experiments; bars, s.e. The 50% inhibitory concentrations were 22.0 ± 1.7 and 29.3 ± 4.1 μM VP-16 for K562 and K/VP.5 cells respectively (P = 0.21).
Yalowich, 1993). In addition, no change in topoisomerase II transcription rate was observed in K/VP.5 compared with K562 cells (Ritke & Yalowich, 1993). Mutations in the coding sequence of the topoisomerase II gene could also result in an alteration in the protein conformation or post-translational modification of this enzyme, thus decreasing the stability of its binding to DNA. Our data demonstrating an attenuation of ATP stimulation of VP-16-induced topoisomerase II–DNA binding in resistant cells correspond to previously published data in VM-26-selected resistant CCRF-CEM cells (Danks et al., 1989) and suggest a mutation in or near ATP-binding domains of topoisomerase II consistent with identified topoisomerase II mutations in several other resistant cell lines (Bugg et al., 1991; Hinds et al., 1991; Lee et al., 1992; Chan et al., 1993; Danks et al., 1993) However, topoisomerase II cDNA sequence analysis has revealed no changes in the region which includes consensus nucleotide-binding domains (nucleotides 1134–1597) comparing K562 and K/VP.5 cells (Ritke et al., submitted).

In addition, single-strand conformational polymorphism analysis of topoisomerase II cDNA from K/VP.5 cells has uncovered no evidence of mutations (Ritke et al., submitted). Single-strand conformation polymorphism analysis has been used previously to identify point mutations in topoisomerase II cDNA from drug-resistant cell lines (Danks et al., 1993). These results suggest that, despite the stable biochemical changes in K/VP.5 topoisomerase II which implicate mutations in the gene coding for this enzyme, there may be other genetic changes in resistant cells that affect topoisomerase II function. Sequence analysis of the entire coding sequence of K/VP.5 and K562 topoisomerase II cDNA is under way and will definitively reveal whether a point mutation(s) in K/VP.5 cells is related to qualitative and/or quantitative topoisomerase II alteration not revealed by the present work.

A second set of results for genetic alteration of VP-16-selected K/VP.5 cells may be a regulator, effector or co-factor of topoisomerase II. Topoisomerase II has been shown to be phosphorylated in intact cells at serine and threonine residues (Saijo et al., 1990; Kroll & Rowe, 1991; Cardenas et al., 1992) and in vitro serves as a substrate for casein kinase II, protein kinase C and p34<sup>ck<sup>2</sup> kinase (Ackerman et al., 1985; 1988; Sahyoun et al., 1986; Cardenas et al., 1992; Devore et al., 1992; Cao et al., 1993). In addition, the activity and degree of phosphorylation of topoisomerase II has been found to increase during cell cycle progression from G<sub>1</sub> to G<sub>2</sub>–M phase (Heck et al., 1989; Wessner et al., 1991; Saijo & Enomoto, 1992). These studies suggest a protein kinase as a candidate topoisomerase II co-factor and a mutational target in resistant K/VP.5 cells. Mutation and subsequent altered activity of a protein kinase that phosphorylates topoisomerase II might reduce the stability of this enzyme, resulting in a decreased level of protein in resistant cells. A change in topoisomerase II phosphorylation in resistant cells might also influence ATP binding/hydrolysis and affect catalytic activity, as has recently been demonstrated using Dro sophila topoisomerase II (Corbett et al., 1993). Altered phosphorylation of topoisomerase II in resistant cells may change the stability of topoisomerase II binding to DNA such that VP-16 stabilisation of the protein–DNA complex is compromised. Recent studies in this laboratory indicate that topoisomerase II phosphorylation is reduced at least 2-fold in K/VP.5 compared with parental K562 cells (Ritke et al., submitted). Therefore, altered topoisomerase II phosphorylation in K/VP.5 cells correlates with decreased VP-16-induced topoisomerase II–DNA binding stability in these resistant cells (Figure 8).

Based on studies presented here, we conclude that selection for a low level of resistance to VP-16 resulted in a quantitative reduction of topoisomerase II expression as well as distinct qualitative changes affecting VP-16-induced stability of topoisomerase II–DNA binding. The multiple stable phenotypic changes exhibited by the novel K/VP.5 cell line provide an opportunity to increase our understanding of the post-transcriptional and/or post-translational modifications in DNA topoisomerase II which regulate its activity in acquired drug resistance.

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**Abbreviations:**
Topoisomerase II DNA topoisomerase II; VP-16 (etoposide), 4'-demethylepipodophyllotoxin 9-(4,6-D-ethylenedioxy-β-D-glucopyranoside); M-AMSA (amsacrine), 4',9-acridinylamino)methane-sulphon-m-anisidide; SSB, single-strand break; DPC, DNA–protein cross-links; SDS, sodium dodecyl sulphate; DMSC, dimethylsulphoxide; EDTA, ethylenediamine tetraacetic acid; EGTA, ethyleneglycol-bis-(β-aminoethyl ether) N,N',N',N',-tetraacetic acid.

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