Different mechanisms of decreased drug accumulation in doxorubicin and mitoxantrone resistant variants of the MCF7 human breast cancer cell line

C.W. Taylor, W.S. Dalton, P.R. Parrish, M.C. Gleason, W.T. Bellamy, F.H. Thompson, D.J. Roe & J.M. Trent

Section of Hematology/Oncology, Department of Internal Medicine, Arizona Cancer Center, University of Arizona, Tucson, Arizona 85724, USA.

Summary We selected two drug resistant variants of the MCF7 human breast cancer cell line by chronic in vitro exposure to doxorubicin (MCF7/D40 cell line) and mitoxantrone (MCF7/Mitox cell line), respectively. The cell lines are similar in growth characteristics including doubling time, DNA synthetic phase and cell size. Resistance to mitoxantrone conferred only partial resistance to doxorubicin; whereas resistance selected for doxorubicin appeared to confer complete resistance to mitoxantrone. Both agents selected for cross resistance to the Vinca alkaloids. MCF7/D40 cells display a classic-multi-drug resistance phenotype with expression of P-glycoprotein, decreased drug accumulation relative to the parental line and reversal of drug accumulation and drug resistance by verapamil. MCF7/Mitox cells likewise display resistance to multiple drugs, but in contrast to MCF7/D40 cells do not express P-glycoprotein by immunoblot or RNA blot analysis. Net drug accumulation in MCF7/Mitox cells was decreased relative to the parental cells but there was no selective modulation of drug accumulation or in vitro drug resistance by the addition of verapamil. Efflux of mitoxantrone was enhanced in both the MCF7/D40 and MCF7/Mitox cell lines relative to the MCF7/S cell line. We conclude that the two drug resistant cell lines have different mechanisms of decreased drug accumulation.

Breast cancer is responsive to a wide variety of single and combination chemotherapy regimens. Unfortunately, an initial response to chemotherapy most patients with metastatic breast cancer ultimately develop recurrences (Dalton, 1990). In such patients, clinical drug resistance (failure to respond to drugs which were initially effective) is a common phenomenon. Clinical drug resistance is likely due to a number of factors such as tumour growth kinetics, development of pharmacologic sanctuaries due to loss of vascular supply, development of hypoxia in large tumours, and development of structural and metabolic changes in individual tumour cells.

Doxorubicin (DOX) is the most active single agent currently available for the treatment of breast cancer (Tormey, 1975). For previously treated and untreated patients the response to DOX as a single agent varies between 28 and 43% respectively. In addition, DOX is an integral component of many combination chemotherapy regimens for breast cancer. Mitoxantrone (Mitox) belongs to a new class of synthetic chemotherapeutic agents, the anthracyclines (Alberts et al., 1980). Numerous comparisons have been made between DOX and Mitox in that both compounds possess dihydroxyquinones and are believed to intercalate DNA (Shenkenberg & Von Hoff, 1986). Mitox is active as a single agent in patients with breast cancer (Smyth et al., 1984; Yap et al., 1981).

Individual tumour cells may develop resistance to a broad range of structurally unrelated drugs and thus display the multidrug resistance (MDR) phenotype (Gerlach et al., 1986). A 170,000 dalton cell membrane protein, termed P-glycoprotein is over-expressed in some MDR cells and functions as a drug efflux-pump (Gerlach et al., 1986; Pastan & Gottesman, 1987). A DOX-resistant MCF7 breast cancer cell line (DOXR MCF7) was previously reported and shown to have a decreased intracellular drug accumulation, over-expression of P-glycoprotein and over-expression of an anionic form of the enzyme glutathione transferase (Batist et al., 1986; Cowan et al., 1986). Another DOX-resistant breast cancer cell line (MDA-AD5) was found to express P-glycoprotein gene sequences (Fuqua et al., 1987). In comparison to DOX resistance, little has been reported about the development and mechanisms of Mitox resistance in human breast cancer cells. Recent reports, however, indicate that resistance to Mitox in colon cancer (WiDr) and leukaemia (HL-60) cell lines conveys a unique MDR phenotype unrelated to the over-expression of P-glycoprotein (Dalton et al., 1986; Harker et al., 1989).

We have selected DOX and Mitox resistant variants of the MCF7 human breast cancer cell line and provide evidence that these two drug resistant cell lines share some similarities in MDR phenotype and both have decreased drug accumulation and enhanced drug efflux. The decrease in drug accumulation appears to be partially responsible for the MDR; however, the mechanisms of decreased drug accumulation in the two cell lines are different.

Material and methods

Cell culture

The MCF7 parent cell line was obtained from the American Type Culture Collection (ATCC, Rockville, MD). Authenticity was confirmed by cytogenetic analysis. The MCF7 cells were adherent to plastic and grown in RPMI 1640 medium supplemented with 10% (v/v) foetal bovine serum, 1% (v/v) penicillin (100 units ml\(^{-1}\)), 1% (v/v) streptomycin (100 µg ml\(^{-1}\)), and 1% (v/v) L-glutamine (GIBCO, Grand Island, NY). Cells were maintained at 37°C in a 5% CO\(_2\)-95% air atmosphere.

Drugs

DOX was obtained from Adria Laboratories (Columbus, OH); Vincristine (VCR) from Eli Lilly Laboratories (Indianapolis, IN); Verapamil (VER) from Knoll Pharmaceuticals (Whippany, NJ); Mitox from Lederle/Cyanamid (Pearl River, NY); amascine (m-AMSA) from Ben Venue Laboratories, Inc. (Bedford, OH). Vinblastine (VBL) and 5-fluoracil (5-FU) were obtained from LyphoMed, Inc. (Rosemont, IL). Etopo-
side (VP-16) and mitomycin-C (MitoC) were obtained from Bristol Laboratories (Evansville, IN). Cisplatin (CDDP), melphalan (L-PAM) and gramamicidin-D (Gram-D) were obtained from Sigma Chemicals (St. Louis, MO). 

Selection of drug resistant cells

MCF7 parental cells were exposed to DOX or Mitox, each at an initial concentration of 1 x 10^{-6} M. Fresh drug was added weekly to keep the concentration approximately the same (approximately 3 x 10^{-7} M weekly). As allowed by cell growth, the concentration of each drug was slowly increased in a multiple step procedure. Over a period of 19 months the concentration of DOX was increased from 1 x 10^{-6} M to 7 x 10^{-5} M. An additional 12 months were required to reach the final DOX concentration (4 x 10^{-5} M, representing a 40-fold increase) and full development of the DOX resistant variant (MCF7/D40) for a total selection time of 31 months. The Mitox resistant cell line (MCF7/Mitox) was selected in a similar fashion by increasing the Mitox concentration from 1 x 10^{-4} M to 8 x 10^{-4} M over a 6 month period and maintaining this concentration over the next 18 months. Prior to any experiments, cells were maintained in drug-free medium for 1 week.

In vitro drug assays

Cytotoxicity was determined using a modified MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium) dye assay (Carmichael et al., 1987; Denizot & Lang, 1986). This is a colorimetric assay based on the ability of viable cells to reduce MTT to a blue formazan product. In dose response curves the data are presented as the per cent of an optical density value obtained for untreated cells. Cells were plated into 96-well microtitre plates (Falcon, Becton Dickinson and Company, Oxnard, CA) at 1 x 10^4 cells/well in 0.2 ml of media containing appropriate concentration of drug with replicates of six (standard deviations were within ±10% of the mean values) VER (6 μg ml^{-1}, 13.2 μM) was added 15 min prior to drug and was present continuously during the period of incubation with drug. After incubation for 4 days at 37°C, 50 μl of MTT dye (2 mg ml^{-1}) was added to each well and incubated for 4 h. Plates were then centrifuged at 500 g for 5 min, media aspirated, dimethyl sulfoxide (DMSO) added to each well (100 μl), plates mechanically agitated for 5 min and optical density at 570 nm determined on a microplate reader (Dynatech Labs, Alexandria, VA). Each experiment was repeated a minimum of three times and data from a representative experiment are shown. A clonogenic assay (Malinm & Perry, 1967) was used to confirm results obtained with the MTT assay for Mitox with and without VER. The concentration of drug which produced a 50% inhibition of cloning efficiency (IC_{50}) was calculated by linear regression analysis of the linear portion of the dose response curves.

Cell growth characteristics

For the determination of doubling time, cell growth curves were established for each cell line by plotting cell number versus growth time. Doubling time was determined directly from the linear portion of the plot. The fraction of cells in S-phase and relative cell size were determined using flow cytometry (FACStar Flow Cytometer, Becton Dickinson) after propidium iodide staining (Krishan, 1975; Deitch et al., 1982). Actual tumour cell diameters were determined by direct measurements of 100 tumour cells per cell line using an inverted microscope equipped with a reticle and an internal standard. The results are expressed as the mean cell diameter and the standard deviation.

Chromosome analysis

Cultures were harvested for karyotypic analysis, slides prepared and G-banding performed as previously described (Trent & Thompson, 1987). A minimum of 30 cells per cell line were analysed, with results expressed according to ISCN recommendations (Harden & Klinger, 1985). Karyotypic information on the parental MCF7 cell line has been presented previously (Osborn et al., 1987).

Drug accumulation

Cellular accumulation of drug was determined during a 1 h exposure of an aliquot of cells (1 x 10^6 cells plated 2 days previously into a 35 mm² petri dish) to 12^4 DOX-5.0 μM, 12^4 Mitox-10.0 μM, and 3^4 VCR-1.6 μM at 37°C. These drug concentrations were chosen based on the relative specific activities of the three radiolabelled agents in an attempt to achieve similar counts per minute (c.p.m.) in MCF7/S cells. One hour after the addition of radiolabelled drug, the cells were washed twice with ice cold phosphate buffered saline (PBS) trypsinised, incubated with 0.2 N NaOH for 2 h and neutralised with 0.2 N HCl (equivalent volumes). The amount of radiolabel (c.p.m.) was determined using liquid scintillation counting. Each experiment was repeated a minimum of three times and data from a representative experiment are shown.

For all experiments involving VER, cells were incubated in media containing 6 μg ml^{-1} (13.2 μM) VER for 15 min prior to drug exposure and during the 1 h drug exposure.

Mitoxantrone efflux studies

Cells were incubated with 12^4 Mitox for 1 h as described in the Drug Accumulation section. At the end of 1 h, cells were placed on ice and washed once with ice cold PBS, drug-free RPMI complete media (2% foetal bovine serum) was added, and cells were incubated at 37°C for up to 60 min. At the end of each time period, cells were washed, trypsinised, digested, and counted as described previously. Efflux at each time point was determined by:

\[
\frac{\text{c.p.m.}}{\text{c.p.m.}} \times 100
\]

Each experiment was repeated a minimum of three times and data from a representative experiment are shown.

Non-protein sulphydryl (NPSH) measurements

The NPSH content of the MCF7 cell lines was measured using the method of Sedlak and Lindsay (1968). A total of 5 x 10^5 cells was washed twice with ice cold PBS (pH 7.4) and transferred to a microcentrifuge tube where they were lysed by sonication (model 250 Branson sonifier, Danbury, CT). Cellular protein was precipitated by addition of 5% sulfosalicylic acid. The cell lysate was then centrifuged at 12,000 g for 5 min at room temperature and 1 ml aliquot of this supernatant was transferred to a tube containing 0.2 M Tris buffer (pH 8.9). To each tube, 100 μl of 0.01 N 5,5'-dithiobis(2-nitrobenzoic acid) in absolute methanol was added. The contents were mixed and the absorbance of each sample was measured at 412 nm. The concentration of NPSH in the sample was determined by comparing the optical density reading of the sample to a standard curve constructed using reduced glutathione. NPSH levels were analysed 4 days post cell passage in all lines. Protein measurements were determined according to the method of Lowry et al. (1951).

Immunoblot analysis for P-glycoprotein

Plasma membranes were purified from 2 x 10^6 cells (Riordan & Ling, 1979). Polyacrylamide gel electrophoresis (50 μg protein per lane) was performed according to the method of Laemmli (1970). Protein was transferred from sodium dode-
cyl sulphate-polyacrylamide gels onto nitrocellulose filter paper according to the method of Towbin et al. (1979). The nitrocellulose filters were then incubated overnight at room temperature in tris-PBS (TPBS) containing the C219 monoclonal antibody (IgG) (kindly provided by Dr Victor Ling, Ontario Cancer Institute, Toronto, Ontario, Canada) at a concentration of 10 ng ml$^{-1}$ (Kartner et al., 1985). Following wash steps to remove unbound antibody, the filters were incubated overnight at room temperature in 50 ml of TPBS containing 25 µg of $^{32}$P-labelled rabbit anti-mouse IgG (specific activity – 600 µCi ml$^{-1}$, New England Nuclear, Boston, MA). The filters were again washed to remove the unbound secondary antibody, dried, and exposed to X-Omat AR film (Kodak). A membrane preparation from 8226/DOX40 multiple myeloma cells was added as a positive control for P-glycoprotein (Dalton et al., 1986).

RNA analysis

RNA was isolated from MCF7 cells using guanidium isothiocyanate and cesium chloride centrifugation (Maniatis et al., 1982). Slot blot analysis of total cellular RNA was carried out as described (Davis et al., 1986) using a 640 bp cDNA fragment of mdr-1, p-chip-1, isolated from a colchicine-resistant CHO cDNA library (Riordan et al., 1985). The probe was oligolabelled according to the method of Feinberg and Vogelstein (1983). Human multiple myeloma cells displaying the multidrug resistant phenotype (8226/DOX40) were included as positive controls and the drug sensitive parent cells (8226/S) as negative controls. The blots were also probed with $^{32}$P-labelled human tubulin cDNA (American Type Culture Collection, Rockville, MD) to confirm the amount of RNA contained in each sample.

Statistical methods

The P-values for the comparison of drug accumulation with and without VER were calculated using a two-sample independent t-test. The P-values for the difference in drug accumulation between cell lines were calculated using Tukey’s Studentised Range Test and the Bonferroni multiple comparisons procedure (Miller, 1981). The 95% confidence limits of the doubling times were determined from the slope of the linear portion of the growth curves (Mood et al., 1974). The slopes of the Mitox efflux curves were evaluated by Analysis of Covariance (Snedecor & Cochran, 1980).

Results

Growth characteristics

The growth characteristics of the sensitive, parental line (MCF7/S), DOX resistant variant (MCF7/D40) and Mitox resistant variant (MCF7/Mitox) are listed in Table I. The doubling times for MCF7/S, MCF7/D40 and MCF7/Mitox cells were similar. No differences between the fraction of cells in S-phase were demonstrated between the three cell lines.

The mean cell diameter varied slightly among the three cell lines, but the confidence intervals (standard deviation) were overlapping.

Dose response curves

Dose response curves for MCF7/S and MCF7/D40 cells after exposure to DOX (with and without VER) as determined by the MTT assay are depicted in Figure 1a. Similar curves for MCF7/Mitox cells after exposure to Mitox (with and without VER) are depicted in Figure 1b. Significant resistance to DOX and Mitox was observed for both MCF7/D40 and MCF7/Mitox cells, respectively. However, the addition of

---

**Table 1 Growth characteristics**

| Characteristic | MCF7/S | MCF7/D40 | MCF7/Mitox |
|---------------|--------|----------|------------|
| Doubling time, hours | 41.6 | 35.2 | 40.3 |
| (95% confidence limits) | (32.1–59.0) | (25.5–56.8) | (35.0–47.6) |
| *Fraction of cells in S-phase | 38% | 38% | 31% |
| *Relative cell size | 1.00 | 1.09 | 1.00 |
| Mean cell diameter in microns (SD) | 18.5 | 21.0 | 17.5 |

* Determined using propidium iodide and flow cytometry. * Determined using flow cytometry, MCF7/S normalised to 1.00.
VER affected resistance only for the MCF7/D40 cell line. VER alone was minimally toxic to the cells (data not shown). The 50% inhibitory concentration (IC₅₀) of DOX (with and without VER) against MCF7/D40 cells and the IC₅₀ of Mitox (with and without VER) against MCF7/Mitox cells are shown in Table II. With the addition of VER the IC₅₀ of DOX against MCF7/D40 cells decreased 10-fold (30 μM vs 3.1 μM). In contrast, the IC₅₀ of Mitox against MCF7/Mitox cells was essentially unchanged with VER exposure (64 μM vs 91 μM). The experiments performed with the MTT assay were repeated using a clonogenic assay (Malinin & Perry, 1967) for Mitox with and without VER in all three cell lines (Table II). The IC₅₀ from the clonogenic assays were generally lower than those obtained from the MTT assays. However, the relative degree of resistance between the cell lines was similar. In addition, the IC₅₀ from the clonogenic assay for Mitox against MCF7/Mitox cells remains unchanged with VER exposure (7.7 μM vs 7.4 μM).

Cross resistance patterns

Table II demonstrates the sensitivities of the 3 MCF7 cell lines to a number of chemotherapeutic agents. The data are expressed as the IC₅₀ and as the relative amount of resistance of MCF7/D40 and MCF7/Mitox cells to MCF7/S cells (IC₅₀ ratio). A high degree (75 fold) of resistance to DOX was observed in MCF7/D40 cells. In addition, MCF7/D40 cells were cross resistant to a number of other drugs: VCR (190 fold), Mitox (153 fold), VLB (93 fold), m-AMSA (41 fold). The addition of VER partially reversed the resistance of MCF7/D40 cells to DOX, VCR and Mitox. MCF7/Mitox cells displayed a very high degree of resistance to Mitox (1208 fold) which was not reversed by VER. A lesser degree of cross resistance to DOX (8.3 fold) and VCR (22 fold) was observed. VER was ineffective in reversing resistance to both DOX and VCR in MCF7/Mitox cells. The MCF7/Mitox cell line was cross resistant to VLB (43 fold), m-AMSA (42 fold) VP-16 (26 fold) and CDDP (21.5 fold). Only low levels of resistance to L-PAM, Mitox, Gram D or SUF were seen in either MCF7/D40 or MCF7/Mitox cells.

**Drug accumulation studies**

The 1 h net intracellular accumulations of ¹⁴C DOX, ³H VCR and ¹³C Mitox are displayed in Figure 2. In both the MCF7/D40 and MCF7/Mitox cell line, accumulation of ¹³C DOX (Figure 2a), ³H VCR (Figure 2b) and ¹³C Mitox (Figure 2c) was decreased relative to the MCF7/S cell line. The MCF7/Mitox cell line had less net intracellular drug accumulation for DOX and Mitox compared to the MCF7/D40 cell line (P < 0.05 for DOX and Mitox) and the MCF7/S parent cell line (P < 0.05 for DOX and Mitox). Both resistant cell lines had decreased VCR net intracellular accumulation relative to MCF7/S (P < 0.05 for MCF7/D40 and MCF7/Mitox) but were not different from each other (P > 0.05).

The effects of VER in increasing DOX and VCR intracellular drug accumulation were limited to the MCF7/D40 resistant cell line (41% and 298% increased, respectively) with no effect observed in the MCF7/S or MCF7/Mitox cell lines. A slight but significant increase in Mitox accumulation was seen in both the MCF7/S (25%) and MCF7/Mitox (24%) cell lines with a greater increase seen in the MCF7/D40 cell line (63%).

Curves of Mitox efflux versus time in minutes for MCF7/S, MCF7/D40 and MCF7/Mitox cells were depicted in Figure 3. At each time point tested, relatively less drug persisted in MCF7/Mitox and MCF7/D40 cells as compared with MCF7/S cells. At the 5 min time point MCF7/D40 and MCF7/Mitox cells contained 34% (2149 c.p.m.) and 24%

**Table II Cross resistance patterns**

| Drug | MCF7/S | MCF7/D40 | MCF7/Mitox |
|------|--------|----------|------------|
| DOX⁴ | 0.4    | 30.0     | 3.3        |
| DOX + VER | 0.37  | (75)  | (8.3)      |
| VCR  | 0.0039 | 0.74     | 0.086      |
| VCR + VER | 0.0031 | 0.065   | 0.16       |
| Mitox| 0.053  | 8.1      | 64         |
| Mitox + VER | [0.011]ⱉ | [0.14] | [7.7]     |
| VLB  | 0.0070 | 0.65     | 0.30       |
| CDDP | 1.3    | (4.6)    | (21.5)     |
| VP-16| 2.9    | 33       | 76         |
| m-AMSA| 0.86  | 35       | 36         |
| L-PAM| 16     | 40       | 91.4       |
| MitoC| 0.72   | 5.3      | 6.3        |
| Gram D| 0.15 μg ml⁻¹ | 1.2 μg ml⁻¹ | 0.98 μg ml⁻¹ |
| 5FU  | 30     | 103      | 117        |

¹⁰⁰% Inhibitory Concentration, as determined by MTT assay; ⁴Ratio of IC₅₀ for the drug resistant cell line to MCF7/S; ⁵Abbreviations: DOX = Doxorubicin; VP-16 = Etoposide; VCR = Vinristine; m-AMSA = Amsacrine; VER = Verapamil; L-PAM = Melphalan; Mitox = Mitoxantrone; MitoC = Mitomycin C; VLB = Vinblastine; CDDP = Cisplatinum; SFU = 5-Fluorouracil; ⁶IC₅₀ as determined by clonogenic assay.
(1526 c.p.m.) respectively, of the amount of drug contained in MCF7/S cells (6299 c.p.m.). These values were statistically significant at \( P < 0.05 \) for MCF7/S cells compared to both MCF7/D40 and MCF7/Mitox cells. Furthermore, in MCF7/Mitox and MCF7/D40 cells drug efflux was more rapid during the initial 5 min as compared to MCF7/S cells. Statistical comparison of the slopes of the efflux curves during this time period revealed \( P = 0.055 \) and 0.0063 for MCF7/S cells compared to MCF7/D40 and MCF7/Mitox cells, respectively. At time points beyond 10 min the slopes of the efflux curves were not significantly different.

**P-glycoprotein detection**

An immunoblot probed for the presence of P-glycoprotein using the C219 antibody is seen in Figure 4. The 8226/DOX40 cell line was included as a positive control for P-glycoprotein (Dalton et al., 1986). P-glycoprotein was detected in the MCF7/D40 cell line but was not seen in the MCF7/S and MCF7/Mitox cell lines.

**RNA analysis**

Slot blot analysis revealed no evidence of mdr-1 message in the MCF7/S or MCF7/Mitox cell lines but was found to be expressed in the MCF7/D40 line thus confirming our findings using western blotting (Figure 5). Thus, selection with the anthracycline DOX resulted in an expression of the P-glycoprotein message while selection with the antracenedione mitoxantrone did not.

**Cytogenetic studies**

Chromosomal analysis of the MCF7 parent cell line and its DOX and Mitox resistant sublines were performed using G-banding analysis. Detailed cytogenetic analysis of the parental line has been published previously (Osborne et al., 1987). There were obvious similarities between all three cell lines clearly indicating their common origin. All three cell lines demonstrated a near-triploid chromosome number (72–80) with numerous structural and numeric alterations (Table III). The MCF7/D40 cell line deviated the most significantly from the parental cell line (primarily by the addition of new marker chromosomes). The MCF7/Mitox cell line more closely resembled the parental line with the only differences being a loss of the marker chromosomes 1p−, 6q−, 7pHSR, 13q+, 16q+ and the addition of one marker unique to this cell line (3p+). The parental cell line (which is known to be amplified for N-ras) (Graham et al., 1985) displayed cytologic evidence of gene amplification in the form of a homogeneous staining region. This marker was lost in both resistant sublines and neither displayed double minutes or other HSRs.
Table III  Cyto genetic analysis of MCF7 parental and resistant sublines

| Marker | MCF7/S | MCF7/D40 | MCF7/Mitox |
|--------|--------|----------|------------|
| 1p-    | +      | +        | +          |
| 2q+    | +      | +        | +          |
| der(2) |        |          |            |
| 2q-    | +      | +        | +          |
| 3p-    | +      | +        | +          |
| 3p+    | +      | +        | +          |
| 5p-    | +      | +        | +          |
| 6p+    | +      | +        | +          |
| 6q+    | +      | +        | +          |
| 7p+    | +      | +        | +          |
| 7pHSR  | +      | +        | +          |
| inv(7) |        |          |            |
| 9p+    | +      | +        | +          |
| 9q+    |        |          |            |
| 12q-   | +      | +        | +          |
| 13q+   | +      | +        | +          |
| 16q+   | +      | +        | +          |
| 17q+   | +      | +        | +          |
| t(12;19)| +   | +        | +          |
| t(22;22) |    |          |            |
| Xq-    | +      | +        | +          |
| Xq+    | +      | +        | +          |
| t(X:? )|        |          |            |

Non-protein sulfhydryl (NPSH) measurements

The MCF7/S cell line was found to have significantly elevated amounts of NPSH compared to the other two cell lines (25.43 ± 0.34 nmol 10^{-6} cells; P < 0.001; Student’s t-test). NPSH levels in MCF7/D40 and MCF7/Mitox cells were decreased at 16.23 ± 0.28 and 15.83 ± 0.21 nmol 10^{-6} cells respectively (not significant, Student’s t-test). Total protein measurements revealed no differences among the three cell lines (data not shown).

Discussion

In this report we describe the development of two multidrug-resistant cell lines established from a common parental cell line using DOX and Mitox as selecting agents. Of note is the fact that Mitox resistance developed much quicker (6 months) than DOX resistance (greater than 2 years). In establishing the drug resistant variants the concentrations of the selecting agents were increased as rapidly as allowed by cell growth. Indeed, the concentration of DOX could be increased only very slowly in the selection of the MCF7/DOX cell line. However, in spite of a different degree of selection pressure for the MCF7/Mitox cell line (lower final Mitox concentration, shorter selection time) relative to the MCF7/DOX cell line, a greater degree of Mitox resistance (1208 fold vs 75 fold) developed. Formal fluctuation analysis tests were not performed and the rate of mutation to resistance for the individual drugs cannot be stated.

The DOX resistant (MCF7/D40) and Mitox resistant (MCF7/Mitox) cell lines have both similarities and differences in cross resistance to other agents. A broad range of drugs with varied mechanisms of action were studied: DNA binding/intercalating agents (DOX, Mitox), tubulin binding agents (VCR, VLB), alkylating agents (L-PAM, CDDP, Mit-C) topoisomerase II inhibitors (DOX, Mitox, VP-16, mel-Amsa), an antimetabolite (5 FU) and a cell membrane ionophore (Gram-D). In addition to being resistant to DOX, MCF7/D40 cells were highly resistant to the tubulin binding agents, moderately resistant to the topoisomerase agents and displayed lower levels of resistance to the alkyllating agents and the antimetabolite 5 FU. Cross-resistance to Mitox was also high in the MCF7/D40 cell line. The MCF7/Mitox cell line displayed a number of unique characteristics. It developed a very high degree of resistance (1208 fold) to the primary selecting agent (Mitox) with relatively minor cross resistance to DOX (8.3 fold) and VCR (22 fold). In addition, MCF7/Mitox cells were partially cross resistant to CDDP, an agent not commonly associated with the multidrug-resistance phenotype (21.5 fold resistant).

Other Mitox resistant cell lines have been reported and are similar to the Mitox resistant MCF7 cell line in that P-glycoprotein is not over-expressed. Wallace et al. first reported a human colon carcinoma cell line selected for resistance to Mitox (WiDr) (Wallace et al., 1987). This line was further characterised by Dalton et al. demonstrating decreased intracellular accumulation of both Mitox and DOX with no increased expression of P-glycoprotein (Dalton et al., 1988). Harker et al. recently reported an HL-60 leukaemic cell line selected for resistance to Mitox which also lacked P-glycoprotein overexpression (Harker et al., 1989).

Marsh et al. reported a multi-drug resistant HL-60 cell line which over-expressed a 150,000 dalton membrane protein distinct from P-glycoprotein (Marsh & Center, 1988). These investigators felt that the P150 protein was involved in the resistance mechanism and contributed to the decreased intracellular drug accumulation. Whether the Mitox resistant cell lines over-express a novel drug-resistance related membrane protein remains to be determined. Studies are currently in progress to further describe the mechanism of enhanced drug efflux in the MCF7/Mitox cell line.

Other drug resistance mechanisms unrelated to drug transport may be important in conferring drug resistance. A number of 'atypical' MDR cell lines have recently been reported. Danks et al. reported a leukaemic cell line (CEM/VIM-1) selected for resistance to the epipodophyllotoxin, VM-26 (Danks et al., 1987). This cell line was felt to display 'atypical' MDR because it was not cross resistant to Vinca alkaloids and did not display a decreased cellular accumulation of drug. Slovak et al. reported two DOX resistant cell lines which had different mechanisms of resistance; one associated with P-glycoprotein, the other more atypical in its mechanism of resistance (Slovak et al., 1988). Mirski et al. reported a DOX resistant human small cell lung cancer cell line (Mirski et al., 1987) which displayed a typical cross resistance pattern to VP-16, Vinca alkaloids and colchicine but did not over-express P-glycoprotein.

In summary, this study demonstrates that two compounds (DOX and Mitox) with structural similarities result in different mechanisms of resistance when used as selecting agents in the same human breast cancer cell line. Reduced drug accumulation secondary to enhanced drug efflux accounts, at least in part, for the multidrug resistant phenotype in both resistant cell lines. However, only the DOX resistant cell line over-expresses P-glycoprotein compared to the drug sensitive parent cell line. It remains to be determined why DOX, a natural product, induces P-glycoprotein mediated MDR; whereas, Mitox, a synthetic compound induces a different mechanism of MDR.

The authors would like to thank Judith Gooley for her technical assistance and B. Kathryn Monroe for typing the manuscript. This work was supported in part by National Cancer Institute Grants CA17094, CA4304) and CA41183. C.W.T. was a recipient of a Clinical Oncology Career Development Award from the American Cancer Society.

References

ALBERTS, D.S., GRIFFITH, K.S., GOODMAN, G.E., HERMAN, T.S. & MURRAY, E. (1980). Phase I clinical trial of mitoxantrone: a new anthracyclinedione anticancer drug. Cancer Chemother. Pharmacol., 5, 11.

BATIST, G., TULPULE, A., SINHA, B.K., KATKI, A.G., MYERS, C.E. & COWAN, K.H. (1986). Overexpression of a novel anionic glutathione transferase in multidrug-resistant human breast cancer cells. J. Biol. Chem., 261, 15544.
CARMICHAEL, J., DEGRAFF, W.G., GAZDAR, A.F., MINNA, J.D. & MITCHELL, J.B. (1987). Evaluation of a tetrazolium-based semi-automated colorimetric assay: Assessment of chemosensitivity testing. Cancer Res., 47, 936.

COWAN, K.H., BATIST, G., TULPULUE, A., SINHA, B.K. & MYERS, C.E. (1996). Similar biochemical changes associated with multidrug resistance in human breast cancer cells and carcinogen-induced resistance to zoanobiotics in rats. Proc. Natl Acad. Sci. USA, 83, 9128.

DALTON, W.S., CRESS, A.E., ALBERTS, D.S. & TRENT, J.M. (1988). Cytogenetic and phenotypic analysis of a human colon carcinoma cell line resistant to mitoxantrone. Cancer Res., 48, 1882.

DALON, W.S., DURIE, B.G.M., ALBERTS, D.S., GERLACH, J.H. & CRESS, A.E. (1986). Characterization of a new drug-resistant human myeloma cell line that expresses P-glycoprotein. Cancer Res., 46, 1321.

DALTON, W.S. (1991). Management of systemic metastases and the sequential therapy of advanced disease. In: Bland, K.I. & Cope-land, E.M. (eds). The Breast: A Comprehensive Textbook of the Management of Benign and Malignant Diseases, pp. 877–899. W.B. Saunders Company.

DANS, M.K., YALOWICH, J.C. & BECK, W.T. (1987). Atypical multiple drug resistance in a human leukemic cell line selected for resistance to teniposide (VM-26). Cancer Res., 47, 1297.

DAVIS, L.G., DIBNER, M.O. & BATTEY, J.F. (1986). Preparation and analysis of RNA from eukaryotic cells. In Basic Methods in Molecular Biology. p. 129. Elsevier: New York.

DEITCH, A.D., LAW, H. & WHITE, R.D. (1982). A stable propidium iodide staining procedure for flow cytometry. J. Histochem. Cytochem., 30, 307.

DENIZOT, F. & LANG, R. (1986). Rapid colorimetric assay for cell growth and survival: Modifications to the tetrazolium dye procedure giving improved sensitivity and reliability. J. Immunol. Methods, 89, 271.

FEINBERG, A.P. & VOGELSTEIN, B. (1983). A technique for radiolabeling DNA restriction endonuclease fragments to high specific activity. Anal. Biochem., 132, 6.

FUQUA, S.A.W., MORETTI-ROJAS, J.M., SHINEIDER, S.L. & MCGUIRE, W.L. (1987). P-glycoprotein expression in human breast cancer cells. Cancer Res., 47, 2103.

GERLACH, J.H., KARTNER, N., BELL, D.R. & LING, V. (1986). Multidrug resistance. Cancer Surveys, 5, 25.

GRAHAM, K., RICHARDSON, C., MINDEN, M., TRENT, J. & BUICK, R. (1985). Varying degrees of amplification of the N-ras oncogene in the human breast cancer cell line MCF-7. Cancer Res., 45, 2201.

HARDEN, D.G. & KLINGER, H.P. (1985). International System for Human Cytogenetic Nomenclature (ISCN). In Cytogenet and Cell Genes., Jessen, J.T. & Kaehling, M. (eds). 21, 1.

HARKER, W.G., SLADE, D.L., DALTON, W.S., MELTZER, P.S. & TRENT, J.M. (1989). Multidrug resistance in mitoxantrone-selected HL-60 leukemia cells in the absence of P-glycoprotein overexpression. Cancer Res., 49, 4542.

KARTNER, N., EYBERNDEN-PORELLE, D., BRADLEY, G. & LING, V. (1985). Detection of P-glycoprotein in multidrug-resistant cell lines by monoclonal antibodies. Nature, 316, 820.

KRISHAN, A. (1975). Rapid flow cytofluorometric analysis of mammalian cell cycle by propidium iodide staining. J. Cell. Biol., 66, 188.

LAEMMLI, U.K. (1970). Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature, 227, 680.

LOWRY, O.H., ROSENNBROUGH, N.J., FARR, A.L. & RANDALL, R.J. (1951). Protein measurements with the folin phenol reagent. J. Biol. Chem., 193, 265.