Early multidrug resistance, defined by changes in intracellular doxorubicin distribution, independent of P-glycoprotein

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Summary Resistance to multiple antitumour drugs, mostly antibiotics or alkaloids, has been associated with a cellular plasma membrane P-glycoprotein (Pgp), causing energy-dependent transport of drugs out of cells. However, in many common chemotherapy resistant human cancers there is no overexpression of Pgp, which could explain drug resistance. In order to characterise early steps in multidrug resistance we have derived a series of P-glycoprotein-positive (Pgp+/+) and P-glycoprotein-negative (Pgp−−/) multidrug resistant cell lines, from a human non-small cell lung cancer cell line, SW-1573, by stepwise selection with increasing concentrations of doxorubicin. These cells were exposed to doxorubicin and its fluorescence in nucleus (N) and cytoplasm (C) was quantified with laserscan microscopy and image analysis. The fluorescence N/C ratio in parent cells was 3.8 and decreased both in Pgp+/+ and Pgp−−/ cells with increasing selection pressure to 1.2–2.6 for cells with a resistance factor of 7–17. N/C ratios could be restored partly with verapamil only in Pgp+/+ cells. N/C ratio measurements may define a general Pgp-independent type of defense of mammalian cells against certain anticancer agents which may precede Pgp expression in early doxorubicin resistance.

The development of resistance of human cancers to potent anticancer agents has been classically ascribed to the selection and outgrowth of pre-existing or newly occurring subpopulations of resistant tumour cells (Carl, 1989; Coldman & Goldie, 1985; Skipper, 1986). Great progress in the understanding of the mechanism of one type of in vitro derived resistance, the so-called multidrug resistance (MDR), was recently obtained from the successful cloning and transfection of the mdr1 gene, which codes for a 170 kDa plasmamembrane protein, called P-glycoprotein (Pgp) (Gottesman & Pastan, 1988; Hammond et al., 1989; Juranka et al., 1989; Lincke et al., 1990). The role of this glycoprotein was elucidated by studying the phenotype of cell lines, transfected with mdr1 (Gottesman & Pastan, 1989; Hammond et al., 1989; Juranka et al., 1989; Lincke et al., 1990) or selected in vitro for resistance to anticancer agents, such as anthracyclines and vinca alkaloids, colchicine and actinomycin D (Biedler et al., 1988; Bradley et al., 1988). Thus Pgp was proven to confer multidrug resistance. Further it was proposed that Pgp functions as a plasma membrane pump for several classes of lipophilic drugs. Evidence for this was based on the predicted amino acid sequence of Pgp, which contained two consensus ATP binding sites, and on the demonstration of ATP-dependent drug binding to Pgp and energy-dependent drug efflux from Pgp+/+ cells (Broxterman et al., 1988; Choi et al., 1988; Gros et al., 1986). However, it has been shown that in some Pgp+/+ cells with high levels of doxorubicin resistance the net change in cellular accumulation of drug was relatively modest, which was taken as argument against an important role of such a drug export protein in the resistance mechanism in these cells (Siegfried et al., 1983). Recently we showed that in the Pgp+/+ cells 278040 and CH4CS, the resistance to doxorubicin could be quantitatively accounted for, when, in addition to doxorubicin accumulation, the distribution of doxorubicin between nuclear sites and cytoplasmic localisations was taken into consideration (Schuurhuis et al., 1989). The data suggested that these Pgp+/+ cells had developed a resistance mechanism characterised by a decrease in net cellular doxorubicin accumulation, including the nuclear target sites, while accumulation in cytoplasmic structures was relatively unaffected.

We have now investigated whether such a defence mechanism of mammalian cells against certain types of anticancer agents is necessarily dependent on Pgp overexpression and we have therefore especially focussed on low level resistant cells. Hereto we used two series of variants selected by doxorubicin exposure from the human non-small cell lung cancer cell line, SW-1573, which both displayed multidrug resistance.

Materials and methods

Cells and cell culture

The human non-small cell lung cancer cell line was obtained from Dr H. Joene (Department of Anthropogenetics, Free University, Amsterdam; Keizer et al., 1989) and was cultured in Dulbecco’s modified Eagle’s medium + 7.5–10% foetal calf serum (GIBCO Europe, Paisley, Scotland). The resistant sublines were derived by continuous exposure to increasing doxorubicin concentrations: e.g. 2R50 (2R80 etc) means that these cells were continuously cultured in 50 (80 etc) mM doxorubicin, until they were harvested 7–14 days before the experiments by short trypsinisation. There was no significant difference in cell size between all the SW-1573 sublines (diameter SW-1573: 16.2 ± 1.2 µm in nine experiments). Cell cycle distribution as measured with flow cytometry (G1, S and G2 + M phase, resp., expressed as % of total number of cells and determined in two independent experiments) was as follows. SW-1573: 54, 37 and 9, resp.; SW-1573/2R50: 59,33 and 9, resp.; SW-1573/2R120: 62, 20 and 19, resp.; SW-1573/4R50: 41, 40 and 19, resp.; SW-1573/4R120: 54, 28 and 18, resp. Cell doubling times were 22 ± 1 (SW-1573), 32 ± 3 (SW-1573/2R50), 38 ± 3 (SW-1573/2R120), 45 ± 5 (SW-1573/2R160), 26 ± 1 (SW-1573/4R50) and 28 ± 2 (SW-1573/4R120), determined in 3–5 independent experiments (M ± s.d.).

Drug cytotoxicity

Doxorubicin resistance factor (RF) and dose-modifying factor (DMF) were calculated from 50% cell-growth inhibitory
concentrations of doxorubicin (IC₅₀) determined in a cell proliferation assay; IC₅₀ of SW-1573 cells was 0.12 ± 0.02 μM (mean ± s.d. of 5 exp.), 2 h drug exposure in a waterbath at 37°C in culture medium lacking NaHCO₃ but containing 20 mM HEPES). After 2 h exposure to doxorubicin and 32 μM verapamil (for DMF), the cell culture medium was refreshed and after another 4 h with or without verapamil in the same medium the cells were allowed to grow in NaHCO₃ containing medium for three cell doubling-times, again in the presence of 32 μM verapamil for DMF determinations (Schuurhuis et al., 1987). The 2 h exposure was carried out in medium lacking NaHCO₃ in order to be able to compare the cytotoxicity data directly with drug accumulation and N/C ratio measurements (see below). The IC₅₀ value for SW-1573 cells was lower (factor 2) in medium lacking NaHCO₃ than in growth medium but no major differences in resistance factors were found when the two media were compared. Resistance factors for the other drugs were determined in a cell proliferation assay by a continuous incubation with drugs as described (Broxterman et al., 1989; Mosman, 1983). A concentration of 32 μM verapamil was used in order to obtain maximal effects on doxorubicin accumulation, cytotoxicity and intracellular drug distribution, since in case low level MDR cells are used these parameters do not differ much from those in sensitive cells. Previously dose-dependent effects of verapamil on these parameters have been shown (Schuurhuis et al., 1987, 1989, 1990).

Drug accumulation
Doxorubicin accumulation and accumulation-enhancement factor by co-incubation with 32 μM verapamil (AEF) were determined by exposure of adhered cells to 0.5 μM [¹⁴C]doxorubicin (2 h, 37°C) in culture medium lacking NaHCO₃, three rapid cold washes and subsequent trypsinisation of the cells; further procedures were essentially as described (Schuurhuis et al., 1987).

Flow cytometry
For quantification of Pgp, 10⁶ unfixed cells were incubated with the monoclonal antibody MRK-16 (4 μg ml⁻¹) or an irrelevant mouse IgG (5 μg ml⁻¹) for 1 h in a volume of 200 μl at 20°C. Samples were washed three times with phosphate buffered saline + 1% bovine serum albumin (PBS-BSA) and incubated with 100 μl rabbit-antimouse IgG-fluorescein isothiocyanate (100 μg ml⁻¹), DAKO immunoglobulins, Copenhagen, Denmark) for 45 min at 20°C. After washing the cells three times with PBS-BSA, the cells were resuspended in 500 μl PBS-BSA and fluorescence was measured with a FACSTAR Plus, Becton Dickinson Medical Systems (Sharon, Ma).

| Table 1 Doxorubicin resistance and accumulation in SW-1573 variants |
|----------------|----------------|----------------|
| Cell line | RF (dose) | DMF* | AEF* |
| SW-1573 | 1 | 1.62 ± 0.34 | 100 |
| 4R50 | 7.7±3.8 | 7.3±1.4 | 12.2±0.5 | 5.9±0.1 |
| 4R80 | 11.1±7.2 | 12.8±1.4 | 10.2±0.2 | 7.4±0.4 |
| 4R120 | 13.2±2.9 | 10.0±1.0 | 9.2±0.7 | 5.5±0.4 |
| 4R160 | 16.7±2.0 | 16.1±2.0 | 7.1±0.2 | 10.0±1.1 |
| 2R50 | 6.9±0.3 | 2.5±0.1 | 34.0±2.0 | 1.3±2.0 |
| 2R80 | 9.5±2.0 | 5.5±1.4 | 35.4±2.5 | 1.4±2.0 |
| 2R120 | 11.1±2.8 | 6.3±0.2 | 41.9±2.0 | 1.3±2.0 |
| 2R160 | 63 ±12 | 17.2±1.3 | 4.7±0.3 | 9.3 ± 2.7 |

*DMF, dose modifying factor = IC₅₀ minus Vp/IC₅₀ plus Vp, AEF, accumulation enhancement factor = drug accumulation plus Vp/drug accumulation minus Vp. Data are means ± s.d. of 2–3 experiments each performed at least in duplicate (cytotoxicity) or triplicate (accumulation). **Significantly different from 1 (DMF), from SW-1573 (Vp) levels (fourth column) or from accumulation -Vp (fifth column): *P<0.02; **P<0.01 (Student’s-test). Non-Pgp cell lines, as measured with RNAase protection assay (Baas et al., 1990; Zinn et al., 1983). Vp, verapamil.

Determination of N/C doxorubicin fluorescence ratios
Trypsinised cells were allowed to adhere on tissue culture petri dishes (Costar, Cambridge, Ma) for 24 h. Cells were incubated with doxorubicin for 1 h under the same conditions as described for drug accumulation and cytotoxicity experiments in Table I and quickly washed with PBS to reduce background fluorescence. PBS was lacking glucose to prevent drug efflux and redistribution. Dox concentrations were chosen to obtain equal net cellular drug amounts in all cell lines (4 μM in SW-1573 cells). Thirty to fifty cells were recorded for each treatment using laser scanning microscope and fluorescence ratios were quantified by delineating nuclei and cytoplasm interactively using digital image analysis as described (Schuurhuis et al., 1989).

Determination of the ratio intercalated doxorubicin
The ratios of intercalated doxorubicin vs non-intercalated fluorescent doxorubicin were determined as described (Lankelma et al., 1990). SW-1573 and SW-1573/2R160 cells were loaded for 1 h at 37°C with 4 and 20 μM doxorubicin, respectively, in order to obtain about equal intracellular drug amounts.

RNAase protection assay
Ten μg RNA samples were hybridised with a ³²P-labelled 301 nucleotide human mdrl cDNA specific probe, obtained from F. Baas, Neth. Cancer Inst., and analysed by RNAase protection assay as essentially as described (Baas et al., 1990; Zinn et al., 1983). A 3-actin probe was used to control for amounts of analysed RNA.

Results and discussion
Two separate series of resistant cells were selected by continuous exposure to 50, 80, 120 and 160 nm doxorubicin. The cross-resistance pattern of the 2R series has recently been described (Baas et al., 1990; Kuiper et al., 1990); it was shown that all sublines from this series had a decreased doxorubicin and vincristine accumulation, while Pgp expression, which was detectable with a sensitive RNAase protection assay in the parent cell line, was lost during an early selection step, but reappeared, strongly overexpressed, in a later selection step (Baas et al., 1990; Kuiper et al., 1990). In

| Table 2 Cross-resistance in SW-1573 variants |
|----------------|----------------|----------------|
| Cell line | RF (diano) | RF (vincer) | RF (etopos) |
| SW-1573 | 1 | 1 | 1 |
| 4R50 | 3.0±1.3 | 213±80 | 161±53 |
| 4R80 | 3.4±1.6 | 222±76 | 185±70 |
| 4R120 | 4.9±1.5 | 299±55 | 149±45 |
| 4R160 | 6.6±3.7 | 413±108 | 202±55 |
| 2R50 | 3.3±0.4 | 58±1.1 | 32±1.1 |
| 2R80 | 5.9±1.6 | 16±3.1 | 31±4.1 |
| 2R120 | 3.7±0.1 | 17±3.2 | 23±3.7 |
| 2R160 | 35±9 | 480±170 | 146±33 |

Resistance factors (RF) were determined in a continuous incubation assay with drug as described (Broxterman et al., 1989; Mosman, 1983). Data are means ± s.d. from 3–4 separate experiments. The RF data for daunorubicin, vincristine, gramicidin D and etoposide for the 2R series are based on Kuiper et al., 1990 and are shown for comparison. Data are given as mean ± s.d. of 3 experiments performed in triplicate. Fluorescence ratio for 2780AD cells (Schuurhuis et al., 1987) was 37 (mean of seven experiments). Pgp negative cell lines (see Table 1). ND, not determined.
a separate selection with doxorubicin, the 4R series was derived (Tables I and II) which appeared to have overexpression of Pgp from the first selection step on, as determined at protein level with the monoclonal antibody MRK-16 (Table II) and with a RNAase protection assay using a mdr1 specific probe (Figure 1). Remarkably the 4R50, 4R120, 4R160 as well as the 2R160 cells have similar amounts of Pgp, while the 4R80 cells for unknown reasons have higher amounts (Table II). In separate experiments it was shown that cells with a higher degree of resistance (2780RD, Schuurhuis et al., 1987) have a higher amount of Pgp (see legends of Table II), indicating non-saturability of the method used. Both the 2R and 4R series display a MDR-like phenotype with, relative to doxorubicin and daunorubicin resistance (Tables I and II), high vincristine and gramicidin D resistance in Pgp/+ cells and high etoposide resistance in Pgp/- cells (Table II). Doxorubicin resistance could be modulated by coincubation with verapamil in 4R.Pgp/+ cells to a large extent (Table I). In 2R.Pgp/- cells, however, the resistance modulation with verapamil was less effective (Table I). Both the 2R and 4R series show an impairment of doxorubicin accumulation, which was more prominent, however, in the 4R series (Table I). Especially in the 2R series the accumulation defect would not be sufficient to fully account for the observed resistance factors. Doxorubicin accumulation, like doxorubicin resistance, could be modulated more effectively in the Pgp/+ cells (Table I). Thus, while interaction of verapamil with Pgp seems to allow an effective modulation of doxorubicin resistance and accumulation, it does not prove that doxorubicin resistance and impairment of accumulation in these Pgp/+ cells is directly caused by Pgp.

We have shown before by a quantitative approach using laser scan microscopy and image analysis that a decrease in doxorubicin fluorescence nucleus/cytoplasm (N/C) ratio could be measured in the low level mdr Pgp/+ cell line 8226/dox 4 (Broxterman et al., 1990). Those data suggested a correlation of doxorubicin fluorescence N/C ratio with Pgp expression. The present SW-1573 experimental system allowed us to study the doxorubicin fluorescence N/C ratio’s in cells with increasing levels of multidrug resistance in Pgp/+ as well as Pgp/- cells, derived from the same parent cells.

Figure 2 shows a decrease in N/C ratio with increasing selection pressure in Pgp/+ as well as in Pgp/- cells. Interestingly, N/C ratios were slightly lower in Pgp/- cells compared to Pgp/+ cells at the same levels of resistance, while on the other hand doxorubicin accumulation was less in the Pgp/+ cells (Table I). This suggests that the relative contribution of decreased drug accumulation and altered drug

**Table III** Effect of verapamil on doxorubicin fluorescence ratios in SW-1573 variants

| Cell line | N/C | N/C (+ verapamil) |
|-----------|-----|------------------|
| SW-1573   | 3.77±0.18 | 3.87±0.44       |
| 4R50      | 2.60±0.18  | 3.27±0.25       |
| 4R80      | 1.89±0.37  | 4.02±0.18*      |
| 4R120     | 2.02±0.17  | 3.04±0.10d      |
| 4R160     | 1.16±0.24  | 2.61±0.11*      |
| 2R50      | 2.18±0.16  | 1.70±0.03*      |
| 2R80      | 1.72±0.24  | 1.81±0.23       |
| 2R120     | 1.75±0.13  | 1.77±0.37       |
| 2R160     | 0.67±0.12  | 2.39±0.69*      |

N/C ratios (doxorubicin fluorescence in nucleus/doxorubicin fluorescence in cytoplasm) were measured as described for Figure 2. * P<0.01, Student’s t-test. Significantly different from N/C ratios minus verapamil: P<0.05; **P<0.02; ***P<0.01. Values in brackets represent factor increase of N/C ratios by verapamil. Data are from 2–6 independent experiments each performed in duplicate.
distribution to the resistance phenotype may differ depending on the presence of Pgp. The fact that 4R160 cells have a lower N/C ratio than e.g. 4R50 cells, despite similar amounts of Pgp, indicates that the non-Pgp resistance mechanism might also be present in the 4R cells. However, it cannot be excluded that the high resistance to VP-16 in Pgp− cells would be caused in part by additional changes such as an altered topoisomerase II activity (Baas et al., 1990).

Again, like for doxorubicin accumulation, for an effective measurement of doxorubicin fluorescence N/C ratio by verapamil the presence of Pgp seems to be a prerequisite (Table III, compare e.g. 2R120 with 4R120 cells). Remarkably, verapamil-induced changes in accumulation (Table I) together with changes in N/C ratios (Table III) can account for resistance modulation (DMF, Table I) in Pgp+/− cells but not in Pgp−/− cells, which leaves the possibility of additional actions of verapamil which affect doxorubicin cytotoxicity. In the 4R series the 4R80 cells show a somewhat exceptional behaviour: the amount of Pgp as estimated with flow cytometry is higher than in the other 4R cells (Table II). In line with this stimulation of doxorubicin accumulation as well as reversal of resistance by verapamil was more prominent in the 4R80 cells (Table I). Also the N/C ratio in 4R80 cells was relatively low, while its modification with verapamil was relatively high (Table III).

In this study doxorubicin fluorescence N/C ratios are used operationally to probe mechanisms of drug resistance, related to changes in intracellular drug distribution. These ratios as such do not reflect the actual concentrations of doxorubicin in each compartment, since they do not take into account the quenching of doxorubicin fluorescence at different localisations of the drug: quenching of doxorubicin fluorescence due to DNA intercalation has been estimated at 95% (Chaires et al., 1982), while cytoplasmic fluorescence may be largely unaffected (Budge & Triton, 1985; Tarasiuk et al., 1989). We have used an independent technique to show that N/C ratio changes reflect changes in the actual compartmental amounts of doxorubicin. With this technique the ratio intercalated drug/non-intercalated fluorescent drug can be measured (Lankelma et al., 1990). It was found that this ratio was 4.2 ± 0.6 in SW-1573 cells and 1.2 ± 0.2 in SW-1573/2R160 cells (M ± s.e.m. of two independent experiments). N/C ratio changes might also be found if certain drug binding factors are saturated at the relatively high drug concentrations which were used in the fluorescence assays. This is unlikely, however, since N/C ratios were shown to be independent of doxorubicin concentrations in the medium (J.H.M. de Lange, N.W. Schipper, G.J. Schuurhuis et al., manuscript submitted).

The molecular mechanism(s) responsible for the induction of decreases in doxorubicin N/C ratio are not yet known; in the SW-1573 cell lines the presence of Pgp is not required for this decrease to occur. One explanation would be the presence of a pump protein present at vesicular membranes oriented to pump drug inside such vesicles. However, in the case of Pgp, the antigenic determinant of MRK-16 could be detected only on the Golgi stack but not in other vesicles (Willingham et al., 1987). Moreover, similar amounts of Pgp, as estimated with MRK-16 binding, do not seem to correlate with resistance factor or N/C ratios (compare e.g. 2R160 and 4R160 cells in the Tables I−III). Thus, even after the induction of Pgp overexpression, the early, non-Pgp mediated mechanism, may still be operative. Since cytoplasmic pH was elevated in some resistant SW-1573 isolates, compared to SW-1573 parent cells as measured in medium without bicarbonate (Keizer & Joenje, 1989), one possibility to be considered is that anthracyclines are forced to accumulate to a higher extent in an acidic vesicular compartment in the cell (Beck, 1987; Keizer et al., 1989; Sehested et al., 1987). Since even in steady-state the 2R/Pgp− cells accumulated less doxorubicin (Table I) or vincristine (Kuiper et al., 1990), an active extrusion process could be present. Active drug extrusion processes for different types of drugs, not related to Pgp overexpression, have in fact been reported recently (Henderson & Tsuji, 1990; Hindingburg et al., 1989; McGrath et al., 1989a,b). In such highly resistant cells anthracycline resistance could also be related to differences in drug binding to nuclear and cytoplasmic binding sites (Hindingburg et al., 1989). Since different independently selected, low resistant isolates from SW-1573 cells showed the Pgp−/− multidrug resistance (Baas et al., 1990), which now has been shown to be related to decreased doxorubicin N/C ratio’s, a mutation in one gene product could be responsible for the observed resistance pattern in Pgp−/− cells. The remarkably low resistance factors of 4R160 compared to 2R160 for both anthracyclines and etoposide, despite a similar Pgp expression further suggests that such a mechanism still prevails in the resistance of 2R160 Pgp+/− cells for these drugs. The combined measurement of doxorubicin accumulation and fluorescence N/C ratio’s in tumour cells, isolated directly from patient’s tumours would allow to investigate the role of drug transport resistance in failure of chemotherapy. Further, applying immunohistochemical staining techniques after N/C ratio measurements would allow a direct comparison of functional changes (reflected by N/C ratio changes) and the presence of resistance-related antigenic determinants (e.g Pgp) in individual cells.

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