Abstract. Drug resistance is one of the major obstacles to chemotherapy of ovarian cancer. Studies with cell lines can serve as an initial screen for agents that might modulate drug resistance. To establish more appropriate models of drug resistance and explore whether the differences exist in the different drug resistant sublines selected by different treatments, we induced SKOV3 cell line using cisplatin (CDDP) and Taxol over a period of 16 months by the pulse (SKOV3/CDDP-P and SKOV3/Taxol-P) and intermittent incremental (SKOV3/CDDP-80 and SKOV3/Taxol-25) method, respectively. The resistant phenotype of the four resistant sublines, SKOV3/CDDP-P, SKOV3/CDDP-80, SKOV3/Taxol-P and SKOV3/Taxol-25, was very stable and the resistance index was 4.12, 11.50, 261.98 and 622.76, respectively. In cell morphology, the cells from pulse treatment had remarkable changes compared with the cells from intermittent incremental treatment. SKOV3/CDDP-80 and SKOV3/Taxol-P grew more slowly than SKOV3/CDDP-P and SKOV3/Taxol-25. Multidrug resistance gene 1, multidrug resistance protein 1, lung resistance protein and glutathione S-transferase pi mRNA expression of SKOV3/CDDP-P and SKOV3/Taxol-25 had greater changes than that of SKOV3/CDDP-80 and SKOV3/Taxol-P. The results suggest there are great differences between the resistant cell lines resulting from pulse and intermittent incremental method. The resistant cells selected by the intermittent method were more resistant than the cells selected by the pulse method. The two resistant sublines selected by the pulse method may serve as appropriate models for the study of mechanisms of drug resistance in ovarian cancer.

Introduction

Ovarian cancer is the leading cause of gynecological cancer mortality. Despite the fact that first-line chemotherapy is effective in reducing tumor burden following optimal cytoreductive surgery, the 5-year survival rate for stage III and IV disease is ~20-30% (1). One of the major reasons for the low 5-year survival rate is the appearance of drug resistance. The combination chemotherapy of CDDP and Taxol has been confirmed as the first-line therapeutic protocol via long-term prospective studies of clinical trials and resistance to anti-tumor drugs has been well defined in ovarian cancer (2). The elucidation of drug resistant mechanisms is insufficient to overcome clinical resistance.

Many resistant tumor cells in humans are gradually acquired during chemotherapy. The resistant cell lines, selected by exposure to anti-tumor agents, have been valuable tools for the illumination of the factors underlying drug resistance since the first induced resistant cell lines in vitro were used by stepwise augmentation treatment 40 years ago. Until now the most common method of establishing resistant cell lines is to use increasing continuous administration (3) and low-dosage intermittent incremental inducement (4) with various and inconsistent dosages. The two methods are significantly different from clinical chemotherapy, in which a pulse protocol with certain interval is commonly adopted. The pulse treatment in previous studies was seldom used compared with the two above methods (5,6). Various dosages were selected for the inducement of the resistant cells in different studies. The usage of the resistant cell lines has greatly promoted the understanding of mechanisms of resistance and drug resistance-associated genes, such as multidrug resistance gene 1 (MDR1) and glutathione S-transferase pi (GST-pi) (7,8). However, the crucial problem existing in the present studies is that studies with cells in culture might not always mirror the situation in clinical tumors and contradictory evidence concerning the mechanisms of drug resistance have been reported (9-12). The situation may be due, at least in part, to the difference of the resistant cells selected by dissimilar approach and a failure in combination of the laboratory and the clinic. In view to the previously inconsistent evidence, it is very possible that different mechanisms are involved in drug resistance.
Materials and methods

Drugs. CDDP was purchased from F.H. Faulding & Co. Ltd. (Melbourne, Victoria, Australia) and stored at a concentration of 3.33 mM at room temperature. Taxol was obtained from Bristol-Myers Squibb Co. (Princeton, NJ, USA) and stored at a concentration of 10 nM diluted in DMEM at room temperature.

Cell lines and culture conditions. SKOV3 is a human epithelial ovarian cancer cell line obtained from Cell Culture Center, Institute of Basic Medical Sciences, Chinese Academy of Medical Sciences. The SKOV3/CDDP-P (SD-P) was generated by selecting SKOV3 cells for growth with CDDP at pulse treatment of 100 μM for 2 h, and then the treated cells grew in drug-free medium. Further treatment was not administered until the cells were in exponential phase. The cells were exposed to CDDP in 10, 20, 40 and 80 μM and 10 times each dosage. The cells were incubated in CDDP-containing medium for 48 h and were subjected to another drug treatment when the cells became confluent. Similarly, SKOV3/Taxol-P (ST-P) was induced by 20 pulse treatments with 2.5 μM Taxol for 1 h each time. SKOV3/Taxol-25 (ST-25) was selected by intermittently growing in small dosage of Taxol at 10 nM, 15 nM and 25 nM for 24 h and 10 times each dosage. The four resistant subclones were established over a period of 16 months. All the cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal calf serum, 100 U/ml penicillin and streptomycin. Cells were kept at 37°C in a humidified atmosphere of 5% CO2 and 95% air. These cell lines grew in monolayers and were passaged when cultures were 70-80% confluent. No experiments were performed until all the cells had been maintained in medium-free drugs for 2 months.

Drug sensitivity assay. Cells were harvested from exponential phase and digested using 0.25% trypsin-EDTA. Single-cell suspensions were prepared. Cells were counted using a hemocytometer and then dispersed within 96-well microtiter plates. Six duplicate wells were used for each determination. Plates were maintained at 37°C in a humidified atmosphere of 5% CO2 and 95% air. A 24-h preincubation time was allowed prior to addition of drugs. CDDP or Taxol were added to each well in six to eight grades. After incubation of 72 h with drugs, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (Sigma, St. Louis, MO, USA) solution (100 μg) was added to each well and the plates were incubated at 37°C for 4 h. Following incubation, 10 μl of extraction buffer containing 20% sodium dodecyl sulfate, 50% N,N-dimethylformamide, pH 7.4 was added to each well to dissolve the formazan crystals overnight. Absorbance at 540 nm on each well was measured using Immunoskan 340 (Labsystems, Vantaa, Finland). Control wells for absorbance readings contained cell-free medium. All experiments were performed at least three times. Resistance index (RI) equals the ratio of the inhibitory concentration 50% (IC50) values of resistant to sensitive cells.

Morphological observations. For light microscopy, exponentially growing cells were transferred to 30-mm dishes containing sterile glass slides and allowed to adhere in 5% CO2 at 37°C for 24 h. Then the cells were exposed to CDDP or Taxol for different time periods as described above. The treated cells were immediately plated in drug-free growth medium after drug treatment. Following maintenance in drug-free growth medium for 24 h or several days when the cells were 70-80% confluent, the slides were washed, fixed in methanol for 10 min and stained by the Wright-Giemsa method. For electron microscopy, the harvested cells were fixed with 3% glutaraldehyde in 0.1 M PBS (pH 7.4) at 4°C for 2 h, postfixed in 1% osmium tetroxide overnight at 4°C, dehydrated with a graded series of acetone and embedded in Araldite, which were polymerized at 60°C for 48 h. Ultrathin sections (50-80-nm) were stained with uranyl acetate and lead citrate and then observed with EM400T transmission electron microscope (Philips, Eindhoven, The Netherlands).

Growth curves. Single-cell suspensions were prepared. Aliquots containing 2,000 cells were seeded into 30-mm dishes preloaded with 2 ml medium. Three duplicate wells were used for each determination. Four cell counts for each replicate from each cell line were made every 24 h for 7 days. The data were subjected to linear regression analysis, in which the doubling time (T50) was calculated from the formula: T50 = ln2/slope.

Cell cycle analysis by flow cytometry. Monodispersed cells ~1x10^6 were harvested during the exponential growth phase. The cells were washed with PBS, fixed in 70% ethanol, and stored at -20°C overnight. The fixed cells were washed twice in PBS, resuspended in PBS containing 200 μg RNase A (Sigma) and incubated at 37°C for 30 min. The samples were stained with 20 μg propidium iodide protected from light for 30 min and then analyzed on a FACS calibur (BD Company, Franklin Lakes, NJ, USA).

Semiquantitative RT-PCR. Total RNA from cells of ~5x10^6 was isolated with TRIzol Reagent (Sigma) according to the supplier’s instructions. The RNA concentration was determined.
spectrophotometrically at 260 nm. The quality of the isolated RNA was checked by agarose gel electrophoresis by means of the presence of 28s and 18s rRNA. The isolated RNA was used for the preparation of first-strand cDNA by reverse transcription. The RNA samples were incubated in 20 μl of a reaction buffer containing 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 5 mM MgCl2, 2.5 μM oligo dT-adaptor primer, 1 mM dNTP each, 20 U RNase inhibitor, and 2.5 U AMV reverse transcriptase (Takara, Japan) at 42˚C for 60 min. The PCR mixture of 20 μl contained 10 mM Tris-HCl (pH 8.3), 50 mM KCl, 1.5 mM MgCl2, 0.25 mM dNTP each, 1 U Taq plus polymerase, 2.5 pmol each of the primers and 1 μg cDNA samples. The PCR conditions included an initial denaturation of 2 min at 95˚C, followed by 30 cycles consisting of 20 sec denaturation at 95˚C, 1 min primer annealing at 58˚C, 50 sec elongation at 72˚C in PTC-150 minicycler (MJ Research, USA). A negative control using water instead of DNA was used for each PCR. The sequence of each primer and product length is shown in Table I. The resulting DNA fragments were separated by electrophoresis on a 1.5% agarose gel containing 0.5 μg/ml of ethidium bromide. Actin was used as internal standards. The band intensities were obtained by QCapturePro software and the ratio of each gene to actin was calculated.

**Statistical analysis.** Results were expressed as the mean ± SD of 3-5 repeated experiments. Data analysis was carried out utilizing the SPSS12.0 statistical software package. Continuous variables were analyzed using Student's t-test. Differences between groups were considered significant at P<0.05. The reported P-values were two-sided.

**Results**

**Resistant phenotype of the four sublines.** We finished the establishment of CDDP-resistant and Taxol-resistant cell sublines of SKOV3 over a period of 16 months. The resultant CDDP- or Taxol-resistant sublines were then maintained and passaged in drug-free medium for at least two months. The stability of drug resistance was examined at monthly intervals. The drug sensitivity data of the five cell lines are shown in Table II and Fig. 1. The resistant phenotype was very stable because the values of IC_{50} and RI showed no significant

**Table I. DNA sequence of the forward and reverse primers, length of product and annealing temperature for RT-PCR.**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Length (bp)</th>
<th>Annealing temperature (˚C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-actin</td>
<td>ACACGGCGCCCATCTACGAGGAGGGGCCGGACTCGTCATAC</td>
<td>621</td>
<td>58</td>
</tr>
<tr>
<td>MDR1</td>
<td>CCCATGGATCAATCAGG</td>
<td>157</td>
<td>58</td>
</tr>
<tr>
<td>MRP1</td>
<td>ACCAAGCAGTATCAGGGTGCCTGC</td>
<td>428</td>
<td>58</td>
</tr>
<tr>
<td>LRP</td>
<td>ACAACTACGTGTGGATCTCGGCTTCAGTCCGACATA</td>
<td>350</td>
<td>58</td>
</tr>
<tr>
<td>GST-pi</td>
<td>GGTTGTGACCGTGAGGACTCATGGATCAGCAGCAAG</td>
<td>397</td>
<td>58</td>
</tr>
</tbody>
</table>

**Table II. IC_{50} and RI values of the resistant cell lines by pulse or intermittent method using CDDP or Taxol.**

<table>
<thead>
<tr>
<th>Cell line</th>
<th>IC_{50} (μM)</th>
<th>CDDP</th>
<th>Taxol</th>
<th>RI</th>
</tr>
</thead>
<tbody>
<tr>
<td>SKOV3</td>
<td>6.67±2.58</td>
<td>80.79±10.82</td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td>SD-P</td>
<td>27.24±8.60</td>
<td>ND</td>
<td>4.12</td>
<td></td>
</tr>
<tr>
<td>SD-80</td>
<td>76.07±3.85</td>
<td>ND</td>
<td>11.50</td>
<td></td>
</tr>
<tr>
<td>ST-P</td>
<td>ND</td>
<td>328.83±58.60</td>
<td>261.98</td>
<td></td>
</tr>
<tr>
<td>ST-25</td>
<td>ND</td>
<td>757.46±80.85</td>
<td>622.76</td>
<td></td>
</tr>
</tbody>
</table>

ND, the value of IC_{50} that was not detected.
change during 4 months in drug-free medium (data not shown).

Morphological changes between the pulse and intermittent methods. SKOV3 showed epithelial-like shape and adhered to the disk. It had a distinct cell border and uniform cell size and shape. After exposure to CDDP or Taxol as described above, most of the cells died and cell morphology changed becoming longer and irregular in shape, varied in size, with unclear cell borders. Especially, SD-80 and ST-25 had significant changes. Most of the cells were in neuron-like shape, with some pseudopodiums, and aberrant nucleus and cytosolic granules. Cell morphology gradually recovered in the logarithmic phase. There were no apparent differences between SKOV3 and its sublines from the pulse method by CDDP or Taxol. However, SD-80 and ST-25 were still in multiform shape, in various sizes and unclear borders (Fig. 2A). Electron microscopy of SKOV3 cells showed regular nuclei and nonuniform chromatin distribution within the nuclei. In the resistant cells, there was no significant difference between the pulse method and low-dosage intermittent inducement. The chromatin pattern of resistant cells was finely dispersed compared with SKOV3 cells, which were coarse and aggregated. The resistant cells showed expansion of mitochondria and formation of small vacuoles in cytoplasm, great amount of double membrane and pinocytic vesicles in the cell surface, and appearance of nuclear bags. Especially, SD-P and SD-80 exhibited unusually abundant vacuoles, of which some were empty, and some contained granules (Fig. 2B).

Cell growth and $T_d$ of five cell lines. All the five cell types were in detention phase and cell number had no significant change on the second day after seeding. SKOV3 was in logarithmic phase and cell number was much more than SD-P, SD-80 and ST-P since the third day (Fig. 3). $T_d$ of SKOV3,
that grew slowly had significant difference from SD-P in Td compared with SKOV3 (P<0.01, P<0.001, P<0.01). SD-80 in G0/G1 phase compared with SD-P. The cells of ST-P and every phase of cell cycle, but SD-80 had a trend of increase. There was no statistical difference between SD-P and SD-80 in G2/M phase decreased significantly (P<0.05, P<0.01). ST-25, which grew faster than ST-P (P<0.01), had no significant difference in Td compared with SKOV3.

Cell cycle distribution and DNA content. Cell cycle distribution of each cell line is shown in Table III. The cells of SD-P, SD-80, ST-P and ST-25 in G0/G1 phase increased markedly (P<0.01) and the cells of SD-P and SD-80 in S phase and G2/M phase decreased significantly (P<0.05, P<0.01). There was no statistical difference between SD-P and SD-80 in every phase of cell cycle, but SD-80 had a trend of increase in G0/G1 phase compared with SD-P. The cells of ST-P and ST-25 in S phase decreased compared with SKOV3 (P<0.01). ST-P cells in G2/M phase decreased compared with SKOV3 and ST-25 (P<0.05).

Expression of drug resistance-associated genes. The mRNA expression of MDR1, multidrug resistance protein 1 (MRP1), lung resistance protein (LRP) and GST-pi is shown in Fig. 4. MDR1 was not detected and MRP1, LRP and GST-pi could be detected in the parental SKOV3 cell line. In SD-P, MDR1 was up-regulated, MRP1 and LRP were down-regulated and GST-pi had no change compared with SKOV3 and SD-80. In SD-80, the other three genes had no change except for MDR1 that was up-regulated compared with SKOV3. There was no difference between ST-P and SKOV3 in expression of four drug resistance-associated genes. Compared with SKOV3, expression of MDR1 and LRP increased, whereas expression of MRP1 and GST-pi decreased in ST-25.

Discussion

We induced two resistant sublines by CDDP and Taxol with different approaches, respectively. SD-P and ST-P was generated from pulse treatment similar to that used in the clinic with pulse dosage according to plasma peak concentration in patients. SD-80 and ST-25 was produced by conventional methods. To our knowledge, this is the first study comparing the difference of drug resistance between different methods of inducement and demonstrating the importance of the resistant cell model similar to the clinical situation in ovarian cancer. In RI values, SD-80 had a 2.8-fold increase compared with SD-P and ST-25 had a 2.4-fold increase compared with ST-P, suggesting the resistant cells selected by the intermittent method was more resistant than the cells selected by the pulse method, which was consistent with a previous study (13). From the RI data generated in the identical period, we assumed that the extent to drug resistance of pulse treatment may be more similar to the clinical chemoresistance because the observation that cells selected for resistance in vitro acquire higher levels of resistance than those observed in tumors of patients that are refractory to CDDP-based chemotherapy have been underscored (14). Some studies showed that the resistant cells needed to grow in drug-containing medium to maintain the stability of drug resistance, otherwise the characteristic of drug resistance would be lost (15). However, in our study, the resistant phenotype was very stable and the values of IC50 and RI had no significant change in 4 months in drug-free medium, suggesting the resistant sublines selected by the pulse method may serve as appropriate models for the study of mechanisms of drug resistance in ovarian cancer.

Although the changes of subcellular structure by electron microscopy were similar, the morphology by light microscopy was different between the cells selected by the pulse method and the intermittent method. SD-80 that grew more slowly had more significant changes than SD-P suggesting biological characteristics such as cell cycle and cytoskeleton regulation would appear essential changes following the development of drug resistance. However, the Taxol-resistant cells showed inconsistent changes between morphology and cell growth and proliferation compared with SKOV3. ST-P from the pulse method had no significant change in cell morphology, but proliferation index decreased significantly. On the contrary, ST-25 from intermittent treatment that had significant changes in cell morphology altered less prominently in proliferation index and cell cycle than ST-P. These results showed dissimilar dosages of inducement that affected the property of drug resistance to different degrees.

Mechanisms of resistance to CDDP and Taxol include the following aspects: 1) decreased drug accumulation; 2) increased repair of drug induced damage; 3) altered gene expression and drug target; 4) increased resistance to apoptosis (16,17). The abundant vacuoles in the resistant cells by electron microscopy may contribute to drug uptake and metabolism and dysfunction to drug target, and the changes of subcellular structure such as mitochondria and membrane surface may be involved in increased protection against oxidative stress and efflux of drug, which is supported by previous observations.
epigenetic changes influence upon the occurrence of clinical resistant cases. The anti-tumor agents in clinical patients may have certain resistance. Studies with cell lines can serve as an initial screen exploring mechanisms of drug resistance and reversing clinical intermittent incremental exposure are.

drug resistance to CDDP or Taxol using the pulse or it remains to be answered what the respective mechanisms of that drug resistance is multi-factorial and underscoring the different for dissimilar methods of drug treatment, indicating associated genes and mechanisms of drug resistance was inducement may lead to distinct expression of drug resistance—note that the same drugs but different approaches of

Hospital (Grant 200203 *). We thank Dr Bing Liu for critical

Acknowledgements

this study was supported by the Key foundation of PUMC Hospital (Grant 200203*). We thank Dr Bing Liu for critical comments on the study and colleagues for excellent technical assistance (Department of Cell Biology, Institute of Basic Medical Sciences).

References


