Anticancer drug clustering based on proteomic profiles and a sensitivity database in a lung cancer cell line panel

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Abstract. Previously, we performed a molecular pharmacological study that applied a combination of DNA microarray-based gene expression profiling and drug sensitivity tests in vitro with a view to designing an improved chemotherapeutic strategy for advanced lung cancer. Utilizing recent key technological advances in proteomics, particularly antibody array-based methodologies, the current study aimed to examine the benefit of protein expression profiling in an analogous molecular pharmacological context. We performed protein expression analysis in a panel of lung cancer cell lines via an antibody array approach. Using a modified NCI program, we related cell line-specific proteomic profiles to the previously determined cytotoxic activity of a selection of commonly used anticancer agents, namely docetaxel, paclitaxel, gemcitabine, vinorelbine, 5-fluorouracil (5-FU), SN38, cisplatin (CDDP) and carboplatin (CBDCA). In addition, we compared these results with those obtained from our prior DNA microarray-based transcriptomic study. In our expression-drug correlation analysis using antibody array, gemcitabine consistently belonged to an isolated cluster in a reproducible manner (5). In this prior mRNA expression-drug correlation analysis, previous transcript expression-drug sensitivity correlation analysis. These results suggest that chemotherapy regimens that include gemcitabine should be evaluated in second-line chemotherapy in cases where the first-line chemotherapy did not include this drug. Protein expression-drug sensitivity correlations in lung cancer cells in vitro may provide useful information in determining the most appropriate therapeutic options for lung cancer patients.

Introduction

There are many anticancer agents being used in the treatment of lung cancer, such as cisplatin (CDDP), carboplatin (CBDCA), docetaxel, paclitaxel, vinorelbine, gemcitabine, 5-fluorouracil (5-FU) and CPT-11. A number of combination therapy regimens employing platinum compounds have proven to be effective (1) and are widely applied to initial treatment for unresected non-small cell lung cancer (NSCLC) (2). In addition, docetaxel and pemetrexed have been reported to be effective in the context of second-line chemotherapy for NSCLC (3,4). However, at present, the effect of these therapies on improving patient survival remains far from satisfactory (1-3). Therefore, it is desirable to find more appropriate therapeutic opportunities based on informed insight. Previously, we performed a gene expression profile analysis of a series of lung cancer cell lines using both cDNA filter and high-density oligonucleotide arrays (5). In parallel, we also examined the sensitivity of these cell lines to commonly used anticancer agents (docetaxel, paclitaxel, gemcitabine, vinorelbine, 5-FU, SN38, CDDP and CBDCA) in vitro via an MTT assay (5). We related the cytotoxic activity of each of these agents to the corresponding transcriptomic expression profile in each of the cell lines using a modified NCI program to determine an improved chemotherapeutic strategy for advanced lung cancer (5). In this prior mRNA expression-drug correlation analysis, gemcitabine consistently belonged to an isolated cluster in a reproducible manner (5).

Recent technological advances in protein analysis, such as antibody array approaches, allow for large-scale analysis...
of protein expression (6-8). We related the known cytotoxic activity of the above-mentioned anticancer agents to corresponding proteomic profiling patterns, as determined using an antibody array approach, across our lung cancer cell line panel. In addition, we compared transcriptomic and proteomic data from the same set of lung cancer cell lines. The essential aim of this study was to aid in the selection of appropriate drug combinations for the treatment of lung cancer.

Materials and methods

Cell lines. The following panel of cell lines was used: PC9, PC7, PC14, A549, PK-2, RERF-LC-KJ, RERF-LC-MS, RERF-LC-AI, PC1, PC3, PC10, ABC-1, EBC-1 and QG56. The PC1, PC3, PC7, PC9, PC10, PC14 and QG56 cell lines were obtained from IBL (Gumma, Japan). The A549 cell line was obtained from the American Type Culture Collection (Rockville, MD, USA). The PK-2 cell line was obtained from the Health Science Research Resources Bank (Osaka, Japan), from which RERF-LC-MS, EBC-1 and ABC-1 cell lines were also purchased. PC1, PC3 and PC10 cell lines were provided by S. Hirohashi (National Cancer Center Research Institute, Tokyo, Japan). RERF-LC-KJ and RERF-LC-AI cell lines were obtained from the RIKEN Cell Bank (Ibaraki, Japan). PC7, PC9, PC14, A549, RERF-LC-KJ, RERF-LC-MS, PC3 and ABC-1 are adenocarcinoma cell lines. PK-2, RERF-LC-AI, PC1, PC10, EBC-1 and QG56 are squamous cell cancer cell lines.

Antibody microarray analysis. Antibody microarray assays were performed using an Antibody Microarray system (cat #AA-001; Lab Vision Co., CA, USA) according to the manufacturer’s instructions. In brief, whole cell extracts were prepared and protein samples biotinylated using appropriate buffers from the kit. Antibody microarray slides (containing 722 antibodies in triplicate) were initially blocked by soaking in blocking buffer. Hybridization buffer was added to biotin-labeled protein samples. Antibody microarray slides were set in a humidity chamber, coverslips were slowly placed onto the slides and the samples were applied to the microarray surface. After incubation for 2 h, the slides were washed three times with washing buffer from the kit. Following this, the slides were applied to a second detection chamber and incubated with Cy3-tagged detection antibody for 45 min at room temperature. The slides were washed three times with washing buffer. The slides were imaged with a GenePix 4000B microarray scanner (Axon Instruments/Molecular Devices, ON, Canada), with protein expression levels quantified by measuring the intensity of the signals using Array-Pro Analyzer (Media Cybernetics, MD, USA). Signal intensities were normalized by comparison with the average values of all probes. All the experiments were repeated at least twice.

Data analysis. Data analysis for the correlation coefficients that related the drug activity patterns to protein expression profiles was principally performed by a modified NCI program (10). The symbol (A) (G1) refers to the drug activity matrix in which the rows represent the anticancer drugs and the columns represent the human lung cancer cell lines. The symbol (T) (protein expression) refers to the protein expression matrix in which the rows represent individual protein and the columns represent the cell lines. In order to analyze the relationship between protein expression and drug activity, we generated a protein-drug correlation matrix (AT) (correlation coefficient) in which the rows represent the proteins and the columns represent the drugs. Firstly, we subtracted its mean value from the matrix (A) in the direction of the row and columns for a pre-treatment. Secondly, we normalized each element in the matrix (A) by subtracting its row-wise mean and dividing by its row-wise standard deviation; normalized (T) was generated in a similar way. Finally, we took the inner product of the matrix (A) and the transpose of the matrix (T). The resulting matrix (AT) implied the Pearson correlation coefficients that reflected the relationship between drug activity and protein expression.

The Pearson correlation coefficient r is given by the following formula:

\[ r = \frac{1}{n-1} \sum_{i=1}^{n} \left( \frac{A_i - \bar{A}}{\Phi_x} \right) \left( \frac{T_i - \bar{T}}{\Phi_y} \right) \]

\[ = \frac{1}{n-1} \sum_{i=1}^{n} \left( A_i - \overline{A} \right) \left( T_i - \overline{T} \right) \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} \left( A_i - \overline{A} \right)^2 \frac{1}{n-1} \sum_{i=1}^{n} \left( T_i - \overline{T} \right)^2} \]

where \( \Phi_x \) is the standard deviation of X; and \( \bar{X} \) is the mean of X.

Hierarchical clustering helps to comprehend huge volumes of data. With cluster analysis, the elements are divided into groups that show similar patterns by calculating the distances between their respective rows and columns. The AT-clustered image map (CIM), indicating the correlation coefficients between protein expression and drug sensitivity in the human lung cancer cell lines, was obtained by the linkage-average clustering method, also known as UPGMA (un-weighted pair-group method using arithmetic average). The statistical algorithms and the graphical outputs described here were implemented in MATLAB 6.5 Release 13 (MathWorks Inc., USA).

Results

Clustering on the basis of drug activity and proteomic expression patterns. We used an antibody array, representing 722 antibodies, to perform proteomic profile analysis of 14 human lung cancer cell lines. To avoid the influence of cell culture artifacts, we separately cultured each cell line in six bottles (11). Signal intensities were normalized by comparison with the average values of all probes. Drug sensitivity tests, namely via MTT analysis, had been previously performed on the same panel of cell lines (5). Eight anticancer drugs currently used for lung cancer chemotherapy, namely docetaxel, paclitaxel, gemcitabine, vinorelbine, 5-FU, SN38, CDDP and CBDDA, were selected for these analyses. We then analyzed the protein profiling data in relation to the
activity profiles of the eight drugs examined. The drugs were clustered on the basis of Pearson correlation coefficients that related their activity patterns across the 14 cell lines to the proteomic profile information obtained from the same cell lines (10). The AT-matrix CIM summarized the relationship between drug sensitivity and protein profiling data, as it allows the visualization of patterns of similarity in large sets of high-dimensional data (Fig. 1A) (10). In this analysis, gemcitabine was located in a separate cluster (Fig. 1A). Various proteins were commonly listed that were associated with gemcitabine sensitivity (Table I). Most of these factors were different from our previous transcriptome analysis. Catenin γ was commonly listed as a factor associated with gemcitabine sensitivity using proteomic and transcription analyses.
Clustering on the basis of drug activity and gene expression patterns. We previously performed an analogous gene expression profiling screen using Affymetrix GeneChip arrays (5). We re-analyzed the transcript expression profiling data in relation to the activity profiles of the eight drugs within the same panel of lung cancer cell lines, receiving similar results with respect to drug clusters (Fig. 1B). In this analysis, gemcitabine was again located in separate clusters (Fig. 1B).

Discussion

The NCI-60 set of cell lines, which are representative of tumors from nine different tissue origins, has been profiled by various laboratories at the DNA, RNA and protein levels, as well as correlated with pharmacologic sensitivity (8,10,12,13). In order to aid in the selection of appropriate drug combinations for the treatment of lung cancer, we previously established a human lung cancer cell panel (5). In our previous study, we used a DNA microarray-based gene expression profiling approach, together with assessment of the cytotoxic activity of several widely applied anticancer agents (5). In this report, we related gene expression and drug sensitivity patterns in our cell line panel (5). Our present analysis using proteomic profiling confirmed this prior transcript expression-drug sensitivity correlation analysis. Overall, our data might provide a rational basis, at the molecular pharmacological level, for the combination of chemotherapeutic regimens involving drugs associating with discrete gene-protein expression clusters. Indeed, gemcitabine was deemed, in particular, from our studies, to be a good candidate for the treatment of recurrent or refractory NSCLC.

Using transcriptomic profiling techniques and clinical response data, it is sometimes difficult to consistently reproduce gene-drug sensitivity correlation data. Such data are often influenced by sampling methods, sample preservation status, tumor size and tumor environment status including tumor vessels and inflammation. In our study, the influence of these confounding factors would be expected to be minimal as cancer cell lines were used. However, cell lines differ from tumor cells and should therefore be considered as surrogates.

A question might be raised as to which technologies should be used for determining gene-drug correlations using the NCI approach. Current methods for proteomic profiling include two-dimensional PAGE, antibody microarrays and other mass spectrometry-based techniques (7,8,14,15). Transcript expression profiling via use of DNA microarrays is technologically more advanced and easier than protein profiling. As most biological functions are executed by proteins, it is important to evaluate the utility of transcript profiling for prediction of protein expression. There have been several reports in which gene expression and protein profiling technologies have been used in order to compare the same samples (16,17). These prior reports have shown modest correlation between transcript and protein profiling data. In our study, drug clustering by transcript profiling coincided with that of protein profiling. Transcript profiling may continue to be useful for these analyses. The identified factors associated with drug sensitivities were different between both analyses. There are several possible explanations for this difference including rates of translation, post-translational modification and protein degradation.

In the treatment of lung cancer, a number of combination therapy regimens employing platinum compounds have proven to be effective (1) and are widely applied as first-line treatment for unresected NSCLC; for example, CDDP + docetaxel, CBDCA + paclitaxel, CDDP + gemcitabine, CDDP + CPT-11, CDDP + paclitaxel and CDDP + vinorelbine (2). In addition, docetaxel and pemetrexed have been reported to be effective in the context of second-line chemotherapy for NSCLC (3,4). It is consequently desirable to find more appropriate therapeutic opportunities based on informed insight. The gene-drug correlations using the NCI program in these studies may be a valuable tool for the identification of determinants of anticancer drug activity in tumors and for the design of cancer chemotherapy and may be useful for improved therapeutic options in the treatment of specific tumor types.

Our present analysis using proteomic profiling confirmed the results of the transcript expression-drug sensitivity correlation analysis. The molecular pharmacological approach taken here suggests that chemotherapy regimens involving gemcitabine should be evaluated in second-line chemotherapy when the initial chemotherapy does not include this drug. A total design approach to cancer chemotherapy through gene-drug correlations using the NCI program may yield improved therapeutic options.

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References


