Synergistic interaction of gemcitabine and paclitaxel by modulating acetylation and polymerization of tubulin in non-small cell lung cancer cell lines

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Background: The combination of gemcitabine (GEM) and paclitaxel (PTX) was appealing for clinical exploration due to different mechanisms of action and partially non-overlapping toxicities.

Purpose: The aim of this study was to elucidate a potential effect of this combination on the proliferation of two non-small cell lung cancer (NSCLC) cell lines, A549 and H520.

Materials and methods: Cell lines were treated with GEM and PTX for 48 hours to evaluate the half maximal inhibitory concentration (IC50). To determine the combination index (CI), cell lines were exposed to GEM and PTX, in a constant ratio of IC50, by various combination treatments. GEM’s effect on tubulin was assessed by western blotting and immunofluorescent staining. GEM was combined with nanoparticle albumin-bound-paclitaxel (NP) in evaluating tumor growth inhibition.

Results: The IC50 of GEM and PTX in A549 and H520 were 6.6 nM and 46.1 nM, and 1.35 nM and 7.59 nM, respectively. Among the sequences explored (GEM → PTX, PTX → GEM, and GEM plus PTX simultaneously [GEM+PTX]), GEM → PTX produced a mean CI <1 in both cell lines. Western blotting and immunofluorescent staining revealed the intention expressions of acetylated tubulin protein and enhancement of tubulin polymerization within GEM → PTX group. A combination order GEM → NP also worked synergistically to suppress tumor growth.

Conclusion: The GEM → PTX sequence may represent a promising candidate regimen for the treatment of NSCLC.

Keywords: IC50, combination index, tubulin

Introduction

Globally, lung cancer is a highly lethal form of cancer; in 2018, approximately 19% of the 9.6 million cancer deaths recorded were associated with lung malignancy.¹ Approximately 85–90% of lung cancer cases were diagnosed specifically as non-small cell lung cancer (NSCLC).² An improved understanding of the mechanisms underlying lung cancer would lead to the discovery of novel molecular-targeted therapies and immunotherapies. However, anti-cancer cytotoxic drugs are still important, particularly because of recurrent disease and acquired drug resistance.³

Gemcitabine (GEM), a deoxycytidine analog, is used as single or combination chemotherapy for solid tumors including NSCLC and pancreatic cancer.⁴ Insertion of
its active phosphorylated metabolite, difluorodeoxyuridine triphosphate (dFdTTP), into DNA synthesis will induce cell apoptosis.\(^5\) The combination of GEM and other anti-cancer drugs are recommended for systemic therapy in advanced NSCLC.\(^6\)

Paclitaxel (PTX) is a microtubule-interfering drug that promotes the polymerization of tubulin.\(^7\) Suppression of the microtubules leads to dynamic changes in a cell including mitotic block and cell apoptosis.\(^8\) Targeting drugs to specific cellular pathways that drive cancer cells is a highly promising treatment modality; therefore, a PTX-based combination was essential in the treatment of advanced NSCLC.\(^9\) Nanoparticle albumin-bound-paclitaxel (NP) is a novel drug that delivers PTX in a manner which makes the most of its advantageous pharmacokinetic profile.\(^10\)

The combination of GEM and PTX was appealing for clinical exploration because these drugs exhibit different mechanisms of action and partially non-overlapping toxicities.\(^11\) Previous studies showed that this combination had similar activity in comparison with a carboplatin-based agent but lower efficacy against cisplatin-containing doublets.\(^12\) Subsequent phase II studies, involving first-line and second-line chemotherapy, confirmed a high disease control rate but raised awareness of the safety profile.\(^11,13\)

Interestingly, most previous studies which have investigated the interaction of GEM and PTX have been performed by focusing on the function of PTX as an agent which reinforces the action of GEM. Indeed, PTX was shown to enhance the anti-tumor activity of GEM by increasing levels of the GEM-metabolizing enzyme, deoxycytidine kinase (dCK), which eventually helps to concentrate GEM in cancer cells.\(^14\) In addition, a recent study showed that the administration of NP elevated the concentration of GEM by reducing the levels of cytidine deaminase (CDA).\(^15\) However, few studies have focused on the function of GEM as a reinforcing agent of PTX.

In this study, we evaluated the potential effect of GEM in combination with PTX (in vitro) and NP (in vivo). Our hypothesis was that GEM would enhance the anticancer activities of PTX and influence the synergism between the two drugs.

### Materials and methods

#### Cell culture and reagents
The human NSCLC cell lines, A549 (adenocarcinoma) and H520 (squamous cell carcinoma), were obtained from the American Type Culture Collection (Manassas, VA, USA). The culture medium was RPMI 1640 (Sigma, St. Louis, MO, USA) supplemented with 10% fetal bovine serum (FBS) and 1% penicillin-streptomycin (Wako, Osaka, Japan) under 5% CO\(_2\) at 37 °C. GEM and PTX were purchased from Mochida Pharmaceuticals Co., Ltd. (Tokyo, Japan) while NP was obtained from Taiho Pharmaceuticals Co., Ltd. (Tokyo, Japan).

For immunoblotting, mouse monoclonal anti-β-tubulin antibody was also used for immunofluorescent staining (Atlas Antibodies, Bromma, Sweden).

#### Cell viability assay and drug combination studies

To determine the half maximal inhibitory concentration (IC\(_{50}\)), we carried out a cell proliferation assay using the Cell Counting Kit-8 (Dojindo, Kumamoto, Japan) based on the manufacturer’s instructions. Cell suspensions of A549 and H520, at a concentration of 1×10\(^4\) cell/100 μl, were seeded in a 96-well plate. The next day, cells were treated with GEM or PTX in various concentrations for a total of 48 h. Absorbances were analyzed at a wavelength of 450 nm using a microplate reader (iMark, Biorad). The IC\(_{50}\) was then determined using the four-parameter logistic function \(y = D + (A-D)/(1 + 10^{(x-K_{logC})})\), with parameter C representing the estimation of IC\(_{50}\).

Combination studies were completed by incubating 1×10\(^4\) cell/100 μl in the 96-well plates for 24 h. Next, the cell lines were simultaneously exposed to the drugs in a different order: GEM and PTX sequentially (GEM→PTX), vice versa PTX and GEM sequentially (PTX→GEM) for 24 h per drug, or GEM and PTX simultaneously (GEM+PTX) for 48 h at a constant ratio. Medium containing the first drug was removed and the cells were washed with phosphate-buffered saline (PBS) before adding the second drug. Based on the principles of the median-effect equation by Chou,\(^16\) drug interactions were analyzed using specific software (Calceusyn, Biosoft, Oxford, UK) and results presented as a combination index (CI).

The mean CI for every fraction affected (Fa, % inhibition of cellular proliferation) at 0.5, 0.75, and 0.9 was described as the final CI. The range of CI <0.1, 0.1–0.3, 0.3–0.7, 0.7–0.85, 0.85–0.9, 0.9–1.1, 1.1–1.2, 1.2–1.45, 1.45–3.3, and >10 describes very strong synergism, strong synergism, synergism, moderate synergism, slight synergism, nearly additive, slight antagonism, moderate antagonism, antagonism, strong antagonism, and very strong antagonism, respectively.\(^16\)
Western blot analysis
Measurement of acetylated tubulin protein was commenced by incubating cells in a 10 cm dish before treating with single drugs, or a combination of drugs, in a different sequence for a total of 48 h. On ice, we then applied cell lysis buffer (Cell Signaling Danvers, MA, USA) containing proteinase inhibitors (Roche Applied Science, Indianapolis, IN, USA) to detach cells, aided by manual scraping. An equal amount of extracted protein was then separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE; Wako, Osaka, Japan) and transferred onto nitrocellulose membranes. After blocking, membranes were incubated with a 1:100,000 dilution of primary antibody (anti-acetylated tubulin) overnight at 4 °C. Blotted protein samples were then detected with an enhanced chemiluminescence kit from Thermo Fisher Scientific (Waltham, MA, USA).

Immunofluorescence
Cell suspensions at a concentration 1 x 10⁶/ml were seeded into 2 well chambers on Lab-Tek chamber slides (Thermo Fisher Scientific, Waltham, MA, USA): GEM (6.6 nM in A549 or 46.1 nM in H520) and PTX (1.35 nM in A549 or 7.59 nM in H520) sequentially (GEM→PTX); vice versa PTX and GEM sequentially (PTX→GEM) for 24 h per drug or GEM and PTX simultaneously (GEM+PTX) for 48 h at a constant ratio. After the administration of drugs for a total of 48 h, the cells were fixed with 20 °C methanol with EGTA (a calcium chelator) and permeabilized with 1% Triton. One hour after blocking, incubation overnight with primary antibody at a dilution of 1:250 and 1:500 for α-tubulin and β-tubulin, respectively. Nuclei were counterstained with 4’, 6-diamidino-2-phenylindole (DAPI) and the expression of tubulin was analyzed using a fluorescence microscope.

Apoptosis assay
The cells were incubated in 6-well plates and treated with their IC₅₀ values of GEM, PTX, and concurrent GEM+PTX for 48 h. In sequence group, GEM or PTX was administrated for 24 h followed by combination GEM+PTX in next 24 h. Following drug treatment, cells were then collected and washed twice with cold PBS. After centrifuge, cells were resuspended within 100 μl solution binding buffer at a concentration of 1.0×10⁶ cells/ml and added with 5 μl Fluorescein isothiocyanate (FITC) Annexin V and 5 μl Propidium Iodide (PI). Cells were incubated for 15 min at room temperature under dark conditions. 400 μl binding buffer was added and apoptotic cells were identified using the BD Pharmingen FITC Annexin V Apoptosis Detection kit (San Diego, CA, USA), according to the manufacturer’s instructions.

In vitro scratch assay
To assess the effect of combination drugs on cell migration, a wound-healing assay was performed. The cells were seeded on 6 well plates and incubated at 37 °C until cells reach 80–100% confluence. A scratch was made through confluent monolayer and each group of cells was exposed to single and combination for 48 h, GEM or PTX for 24 h followed combination for 24 h.

Scratch zones representative for each group were photographed at 0, 24, and 48 h by BZ-X700 microscope (Japan). Relative wound closure (RW) is the comparison of the distance of final wound (Wf) and initial wound (Wi); RW =([Wf−Wi]/Wi).

In vivo growth inhibition assay
Twenty-four female BALB/c nude mice were purchased from SLC (Shizuoka, Japan). The research was approved by the Institutional Animal Care and Use Committee of Kobe University ( Permit Numbers: P171201) and carried out according to the Kobe University Animal Experimentation Regulations. Mice were inoculated subcutaneously with 5.0×10⁶ cells/100 μl suspension of cell lines. When the tumor volume (TV) reached 100 mm³, mice were divided into six groups (n=4 per group). Each mouse was treated with sterile PBS intraperitoneally (i. p.) (day 1, 8, 15), GEM 50 mg/kg i. p. (day 1, 8, 15), NP 75 mg/kg intravenously (i. v.) (day 1, 8, 15), GEM i. p. (day 0, 7, 14) followed by NP i. v. (day 1,8,15) (GEM→NP), GEM combined NP (day 1, 8, 15) (GEM+NP) and an NP i. v. (day 0, 7, 14) followed by GEM i. p. (day 1, 8, 15) (NP→GEM).

We then determined a range of parameters: longest diameter (a), widest diameter (b), TV =a x b²/2, TVₐ= the tumor volume on n day, TV₀= the tumor volume on day 0, relative tumor volume (RTV)= TVₐ/TV₀, body weight (BW), and relative body weight (RBW)= BWₐ/BW₀. TV and BW were measured twice a week after injection of an anti-cancer drug.

Statistical analysis
Data are presented as mean ± standard error (SE). Independent two-sample t-test was used to determine the statistical difference between all combination drugs and control groups. A P-value <0.05 was considered to be statistically significant.
Results
The sensitivity of NSCLC cells to GEM and PTX
The sensitivities of A549 and H520 to anticancer drugs are summarized in Table 1. A549 cells were more sensitive than H520. As predicted, PTX inhibited cell proliferation more conclusively in both A549 and H520 than GEM. The IC$_{50}$ value of PTX in A549 and H520 cells were 1.35 nM and 7.59 nM, respectively (Figure 1). The IC$_{50}$ value of GEM in A549 cells (6.6 nM) was approximately 7-fold stronger compared to the IC$_{50}$ value of GEM in H520 cells (46.1 nM) (Figure 1).

The synergistic effects of GEM and PTX
Calculations were carried out at a constant molar ratio of PTX: GEM of 1:4.8 and 1:6.1 for A549 and H520 cells, respectively. Table 2 shows the mean CI for cell lines while Figure 2 illustrates the median-effect and the combination index plots.

The drug interaction for the GEM→PTX sequence produced synergism with a mean CI of 0.91252 and 0.26651 for A549 and H520 cells, respectively. Various synergistic interactions were observed at a Fa of 0.5, 0.75, and 0.9 with CI values of 0.96330, 0.90915, 0.86511 for A459 cells, and 0.10741, 0.22417, 0.46795 for H520 cells, respectively.

The strongest synergism in H520 was obtained at low Fa. There were also synergisms in reverse sequence with a CI of 0.77576 in H520 cells, and concurrent sequence with aCI of 0.91081 and 0.92726 for H520 and A549 cells. Overall, the mean CI of GEM+PTX and PTX→GEM sequences did not meet the criteria of synergism.

GEM-mediated activation of tubulin acetylation
To clarify the synergistic role of GEM in the GEM→PTX sequence, we performed Western blot analysis of acetylated tubulin protein, the main target of PTX. PTX alone, and the combination of drugs, increased the level of acetylated tubulin protein. Among the variant sequences, the highest expression of acetylated tubulin protein was observed in cells treated with GEM→PTX (Figure 3).

The effect of GEM on microtubule polymerization
PTX-treated cell lines with increased levels of polymerized $\alpha$- and $\beta$-tubulin protein were characterized by thicker and denser microtubule bundles (green and red) forming ring-like structures around the nucleus (blue) (Figure 4). As evidenced by immunofluorescence, the sequence of GEM→PTX in both of the cell lines appeared to enhance polymerization. Cells exposed to GEM, and controls, showed an extensive fine microtubule structure within the cytoplasm.

Table 1 The sensitivity of cell lines to gemcitabine and paclitaxel

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Drug</th>
<th>IC$_{50}$ (nM)</th>
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<tbody>
<tr>
<td>A549</td>
<td>GEM</td>
<td>6.6</td>
</tr>
<tr>
<td>A549</td>
<td>PTX</td>
<td>1.35</td>
</tr>
<tr>
<td>H520</td>
<td>GEM</td>
<td>46.1</td>
</tr>
<tr>
<td>H520</td>
<td>PTX</td>
<td>7.59</td>
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Abbreviations: GEM, gemcitabine; PTX, paclitaxel.

Figure 1 Inhibitory effect of gemcitabine (GEM) or paclitaxel (PTX) on human non-small cell lung cancer (NSCLC) cell lines. After incubating cell lines with various concentrations of drugs for a total of 48 h, we measured the absorbance at a wavelength of 450 nm. The IC$_{50}$ of GEM in A549 and H520 cells was 6.6 nM and 46.1 nM, respectively. The IC$_{50}$ of PTX in A549 and H520 cells was 1.35 nM and 7.59 nM, respectively.
To confirm treatment-induced apoptosis, Annexin-V and Propidium Iodide was performed to allow specific differentiation of the apoptotic and necrotic cell. The combination using GEM as a first drug followed by GEM + PTX resulted in a significance number of apoptotic cells, 8% and 9.4%, in A549 and H520 cell line respectively. Administration PTX→GEM+PTX and concurrent GEM + PTX produced a different result in each cell line (Figure 5A and B).

**Apoptosis assay**

**Scratch assay**

The migration of cells in the scratch assay was followed over 48 hrs. Both cell lines displayed a different migratory profile during the wound healing process. A549 cells migrated to the scratched area and approached each other.
to close the wound faster than H520 cells. Although relative wound closure in A549 was almost similar in all treatment group (Figure 6A), administration of GEM for 24 hrs as the first drug continued by the next 24 hrs GEM +PTX in H520 cell line leads to a significant decrease of migration rate (0.82) compared with control (Figure 6B).

### Tumor growth inhibition assay

We evaluated the efficacy of drug combinations to inhibit tumor growth by RTV. The GEM→NP sequence grouped showed a significant tumor suppression in both of cell lines (Figure 7A and B). It was important for us to consider the progression of tumor size. Tumor growth observed in control and GEM treatments of A549 cells was terminated on day 7 and 18, respectively, and mice in control and GEM of H520 were only followed to day 14. In both types of tumor-bearing mice, there were tendencies for tumor regrowth after stopping the treatment injections. The smallest body weight might be influenced by the lowest growth of tumor size in group GEM→NP.

### Discussion

This study utilized two major types of lung cancer cell line, the adenocarcinoma cell line A549 and the squamous cell carcinoma cell line H520. A549 was most sensitive to anti-cancer drugs (Table 1). The result was consistent with the results of previously published reports. On the other hand, our experiments on the H520 cell line showed that it was more sensitive to anti-cancer drugs than described in a previous report.
Because of GEM and PTX have different action modes, the combination of these drugs could potentiate the action of these as single agents. Intriguingly, synergism was observed in the GEM → PTX sequence; this observation was apparent for both cell lines (Table 2). The strongest synergism, with a CI of 0.10741 at 0.5 Fa, was apparent in H520 cells. This finding is of considerable clinical importance even though synergism at a high-level Fa for cancer treatment is more critical.

This leaves us with the question of whether synergism and antagonism could be predicted? An earlier pharmacodynamic study showed that a low dose of GEM (1 ng/ml) induced cell cycle arrest in phase S of the cell cycle, and assisted the efficacy of the subsequent drug. A low dose of PTX will stabilize microtubule during mitosis at phase G₂-M and lead to apoptosis rather than necrosis. Meanwhile, other studies described that low PTX may arrest of the cell only at phase G₁ and also both of G₂-M and/or G₁. Furthermore, low PTX may induce aberrant mitosis resulting in aneuploidy and lead to apoptosis. We assume that administration of low GEM as the first drug in combination produce a transient accumulation cell in phase S that will facilitate efficacy of the next drug, low PTX. In reverse sequence, exposure to low PTX probably caused cells to arrest in phase G₁, thereby the number of cells in phase S was reduced leading to insufficiency of cell cycle-specific cytotoxicity of GEM.

Synergism in combination using low GEM as a first drug were also represented by apoptosis and migration assay. Apoptosis was more pronounced in the sequence combination in which GEM preceded GEM+PTX. The migration rate of cells was also decreased after a scratch made and followed for 48 hrs. The increases in apoptotic cell and deceleration of migration assay may be related in the scheduling of drug combination.

Growth inhibition in GEM→NP seemed more favorable. The maximum tolerated dose (MTD) of NP was between 120–240 mg/kg and the lethal dose was 240 mg/kg. These might be explained that the more cell arrested by GEM in phase S will facilitate the low dose of NP (50 mg/kg) lead to apoptosis of the cell and suppress the tumor growth. GEM works effectively in phase S so that the use of NP as the first drug or concurrent combination will decrease the activities of GEM.

**Figure 4** The effect of gemcitabine (GEM) in polymerization microtubule. Cell lines were prepared for immunostaining β-tubulin (green) and α-tubulin (red) after stimulation with the indicated drugs for a total of 48 h. Nuclei were counter-stained with 4', 6-diamidino-2-phenylindole (DAPI) (blue). Scale bar, 100 μm. Immunofluorescence staining revealed alterations of cellular microtubule structures characterized by the formation of a microtubule bundle. The GEM→PTX sequence appeared to enhance α- and β-tubulin polymerization.
Figure 5 Apoptosis assay. Flow cytometry analysis of both cell line that treated with sequence drug GEM→GEM+PTX resulted in significance meaning. (A) A percentage apoptotic of A549 cell within group GEM→GEM+PTX and vice versa were greater than concurrent GEM+PTX. (B) In the case of H520 cell line, GEM→GEM+PTX and concurrent GEM+PTX produced a similar percentage of apoptotic. *P<0.05 compared with control; † not significant.

Figure 6 Wound healing assay. (A) Relative wound closure of A549 cell line in all group combination drugs were similar and showed no significant results. (B) However, scratch assay showed the most significant deceleration of cell migration was in drug combination GEM→GEM+PTX-treated H520 cell. *P<0.05 compared with control; † not significant.
Hence, the single use of NP might be inhibited the tumor growth similar to group NP→GEM and concurrent.

Microtubule stabilization, resulting in cell apoptosis, is a cornerstone that underlies drug-drug interaction. Microtubules, which consist of α and β tubulin, are highly dynamic structures and represent the target of tubulin-binding anti-cancer drugs. PTX interacts with an amino-terminal region of tubulins against depolymerization. Even at low concentrations, PTX can still exert the ability to suppress the dynamics of a microtubule. Tubulin acetylation is an indicator of microtubule stability. In contrast, histone deacetylase 6 (HDAC6), a class Ib deacetylase, is known to deacetylate substrates such as tubulin. Indeed, in a previous study, a combination of trichostatin A, a histone deacetylase inhibitor, and PTX resulted in a significant increase in tubulin acetylation and microtubule stabilization. On the other hand, GEM can influence microtubules indirectly. For example, GEM can play a vital role in apoptosis driven by caspase-3. The cleavage of the C-terminal ubiquitin-binding zinc finger of HDAC6 will promote apoptosis. Therefore, GEM can enhance the acetylation of tubulin and polymerization of microtubules, thus reinforcing the anti-cancer effect of PTX.

In conclusion, the present study provides evidence that the synergistic drug combination of low GEM and low PTX/NP can inhibit the growth of NSCLC cell lines. Furthermore, the administration of low GEM as a first drug in the combination was proven to enhance the anti-tumor activity of low PTX by modulating tubulin acetylation and microtubule polymerization.

**Abbreviation list**

NSCLC, non-small cell lung cancer; GEM, gemcitabine; dFdCTP, difluorodeoxycytidine triphosphate; PTX, paclitaxel; NP, nanoparticle albumin-bound-paclitaxel; dCK, deoxycytidine kinase; CDA, cytidine deaminase; FBS, fetal bovine serum; IC₅₀, half maximal inhibitory concentration; PBS, phosphate-buffered saline; CI, combination index; Fa, fraction affected; TV, tumor volume; i.p, intraperitoneally; i.v, intravenously; RTV, relative tumor volume; BW, body weight; RBW, relative body weight; HDAC6, histone deacetylase 6; RW, relative wound closure; Wf, final wound; Wi, initial wound.

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Disclosure
The authors report no conflicts of interest in this work.

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