Potentiation of in vitro apoptotic effects of δ-tocotrienol and jerantinine A on human lung adenocarcinoma cells

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ABSTRACT

Introduction: The apoptotic effects of single-compound and combined sub-effective concentrations of δ-tocotrienol and jerantinine A on human lung adenocarcinoma (A549) cells were investigated.

Methods: Assays including cell viability, histochemical and immunofluorescence staining techniques, flow cytometry and enzyme activity were used.

Results: The combination of δ-tocotrienol with jerantinine A at sub-effective concentrations induced a synergistic effect and improved selective toxicity towards cancerous A549 cells over normal lung fibroblast (MRC5) cells compared to the single-compound jerantinine. Morphological features of apoptosis were evident on treated A549 cells. Combined sub-effective concentrations of δ-tocotrienol with jerantinine A induced a predominantly G2/M cell cycle arrest and characterised by a disruption of microtubular networks mediated via caspase 8, 9 and 3 enzymatic activities.

Conclusion: These findings demonstrated improved potency in vitro and reduced dose-related toxicity of jerantinine A to normal cells through prospective combined treatment between low-concentration δ-tocotrienol and jerantinine A for lung cancer.

Implication for health policy/practice/research/medical education: The combination of δ-tocotrienol and jerantinine A at low dosage demonstrated improved apoptotic effect and reduced toxicity towards non-cancerous cells and such could potentially serve as future chemotherapeutic regimen against lung cancer.

non-selective toxicity towards the normal healthy cells.

**Materials and Methods**

**Human cell lines and culture conditions**

Human lung adenocarcinoma (A549) and normal lung fibroblast (MRC5) cells, purchased from ATCC, USA were used in this study. All cell lines were cultured and maintained in optimal conditions as previously described (7).

**Neutral red uptake assay for evaluation of cell viability**

The viability of both A549 and MRC5 cells was determined based on the neutral red uptake assay described previously (8) in a 96-well plate format where A549 and MRC5 cells were pre-seeded at a density of 5 × 10⁴ per well for 24 hours. These cells were independently treated with either δ-tocotrienol (at 0-300 μg/mL concentrations) or jerantinine A (at 0-40 μg/mL concentrations) for 72 hours. Dimethyl sulfoxide (DMSO) was used as a vehicle control (negative) and vinblastine served as an anticancer drug control (positive). For combined treatment, δ-tocotrienol (at 0-24 μg/mL concentrations) was combined with jerantinine A at its fixed IC₅₀ concentration (0.35 μg/mL). The IC₅₀ values of single-compound and combined treatments were determined using the GraphPad Prism 5 software and presented as mean ± SEM (standard error of mean) of triplicates completed in three independent experiments. Based on the combined treatment experiment, the combination index (CI) and dose reduction index (DRI) values were determined as previously described (9). The CI envisions the type of pharmacological interaction between the combined chemotherapeutic agents. CI values of < 1, > 1 or = 1 indicate synergistic, antagonistic or additive pharmacological interactions, respectively (10). The DRI represents the fold reduction on potent concentrations of δ-tocotrienol and jerantinine A used in the combined treatment over their individual IC₅₀ values generated from the single-compound treatments.

**Histochemical staining for examination of cellular morphologies**

Each well of the 2-well chamber slides (SPL Life Sciences, Korea) was seeded with 5 × 10⁴ A549 cells for 24 hours. The A549 cells were then individually incubated with respective IC₅₀ concentration of δ-tocotrienol, jerantinine A or vinblastine (positive control) for another 24 hours. The cells treated with plain media containing DMSO were served as the negative control. For combined treatment, A549 cells were incubated with a combination of δ-tocotrienol (2.77 μg/mL) and IC₅₀ concentration of jerantinine A (0.35 μg/mL). A protocol previously described for haematoxylin and eosin (H&E) histochemical staining (7) was adopted and H&E-stained cells were captured at 40X magnification under Nikon 80i microscope (Nikon, Japan).

**Flow cytometry for cell cycle analysis**

A549 cells (1 × 10⁴) pre-seeded per well in 6-well plates (SPL Life Sciences, Korea) were incubated with similar concentrations of single-compound and combined treatments as stated in histochemical staining section for 24 and 48 hours. Cell cycle analysis was conducted using BD Accuri F4 Flow cytometer (BD Biosciences, USA) and based on the previously published procedures (8). Data were analysed using FCS express 5 research edition (De Novo Software, USA) and presented as mean ± standard deviation (SD) of three repeats.

**Immunofluorescence staining for morphological detection of microtubules**

A total of 5 × 10⁴ A549 cells were seeded in 2-well chamber slides. Following 72-hour incubation with separate treatments consisting of single-compound IC₅₀ and combined sub-effective concentrations (2.77 μg/mL of δ-tocotrienol and 0.35 μg/mL of jerantinine A) as well as positive and negative controls, A549 cells were stained using an immunofluorescence method as previously described (11). The slides were visualised at 40X magnification under Nikon eclipse Ti confocal microscope (Nikon, Japan).

**Colorimetry for determination of caspase enzymatic activity**

A549 cells pre-seeded at a density of 1 × 10⁴ per dish in 90 mm petri dishes (SPL Life Sciences, Korea) for 24 hours, were separately treated with single-compound (IC₅₀) and combined sub-effective concentrations (2.77 μg/mL of δ-tocotrienol and 0.35 μg/mL of jerantinine A) at different time points (1-48 hours). Caspase 8, 9 and 3 enzymatic activities were determined using Gene Tex colorimetric caspase 8 (GTX 85543), caspase 9 (GTX 85538) and caspase 3 (GTX 8558) kits according to manufacturer’s instruction (Gene Tex, USA). The absorbance values for treated and control samples were measured at 405 nm and presented as mean ± SD generated from triplicate experiments.

**Statistical analysis**

Statistical analysis of data was conducted using one-way ANOVA. Dunnett’s test was used to assess for a significant difference between treated and untreated groups, as indicated at P < 0.001 (***)**, P < 0.01 (**), P < 0.05 (*) levels.

**Results**

Both δ-tocotrienol and jerantinine A had growth inhibitory effects on A549 cells with IC₅₀ values of 12.40 ± 1.09 μg/mL and 1.42 ± 1.31 μg/mL, respectively. Similar potent inhibitory effect was evident for the positive control vinblastine (IC₅₀ = 0.03 ± 1.45 μg/mL) on A549 cells. However, jerantinine A induced similar growth inhibitory effect on normal MRC5 cells (0.97 ± 0.11 μg/
mL). The combination of δ-tocotrienol and jerantinine A at sub-effective concentration induced synergistic effect (CI = 0.47) and caused a 4-fold dose reduction on the concentrations of δ-tocotrienol and jerantinine A compared to respective individual IC$_{50}$ values (Table 1). Haematoxylin and eosin staining of individual and combined treatments showed morphological hallmarks of apoptotic cell death including cellular shrinkage and nuclear chromatin condensation on A549 cells incubated with IC$_{50}$ concentrations of vinblastine, δ-tocotrienol, jerantinine A and combined sub-effective concentrations of δ-tocotrienol and jerantinine A. Formation of membrane blebs was particularly evident on cells incubated with IC$_{50}$ of jerantinine A (Figure 1a).

As shown in Figure 1b, c, d, cell cycle analysis revealed the induction of G0/G1 and G2/M arrests on A549 cells by single-compound treatments of δ-tocotrienol and jerantinine A, respectively after 24- and 48-hour incubation periods. On the other hand, the combined low concentrations of δ-tocotrienol and jerantinine A induced G2/M arrest on A549 cells. Further investigation on the effects of single-compound and combined treatments on microtubules, was conducted. Indeed, immunofluorescence study revealed that jerantinine A inhibited or caused a disruption of microtubular network in A549 cells (Figure 2a). This was characterised by cell shrinkage and decrease in immunofluorescence intensity. Similar anti-microtubular effect was observed in A549 cells incubated with the microtubule-destabilising agent, vinblastine (positive control).

An evaluation of caspase enzymatic activity revealed that both single-compound IC$_{50}$ and combined sub-effective concentrations of δ-tocotrienol with jerantinine A induced caspase 8, 9 and 3 enzymatic activities (Figure 2b, c, d). Interestingly, the combined treatment had boosted caspase 9 and 3 enzymatic activities, respectively up to 1.90 and 1.95 folds, which were considerably higher than those of single-compound IC$_{50}$ of δ-tocotrienol and jerantinine A.

**Discussion**

Combined treatment of phytochemicals has been suggested to improve potency and minimize non-selective toxicity. Herein, jerantinine A induced similar toxicity to A549 cells and normal MRC5 cells conforming to a previous report on its non-selective toxicity on normal cells (5).

In contrast, unlike the single-compound treatments of δ-tocotrienol and jerantinine A, the combined treatment demonstrated more than 2-fold selective toxicity on A549 cells thereby minimizing toxicity on normal MRC5 cells. Indeed, the study herein conformed to previous suggestion that combined therapy could potentially minimize non-selective toxicity and improve potency of phytochemicals (6).

Further investigation using H&E staining technique revealed the manifestation of the morphological features of apoptosis on treated A549 cells. The cells undergoing apoptosis are single or cluster of cells that appear as circular or oval shaped mass with dark cytoplasm and dense purple nuclear chromatin upon staining with H&E dyes (12). Similar morphological observation of these apoptotic features herein strongly indicates that single-compound and combined low concentrations of δ-tocotrienol and jerantinine A induced apoptosis on A549 cells. An evaluation of cell cycle revealed that induction of G0/G1 and G2/M arrests A549 cells by single doses of δ-tocotrienol and jerantinine A, respectively, whereas, the combined treatment induced G2/M arrest. Microtubule-targeting agents have been shown to induce G2/M cell cycle arrest (13). Therefore, further investigation revealed that jerantinine A herein caused disruption of microtubules. This corroborated the previous mechanistic study that had demonstrated the potential anti-microtubular effect of jerantinine A (5). In contrast, untreated cells (vehicle control) and cells treated with IC$_{50}$ of δ-tocotrienol showed an intact α-tubulin structure. Interestingly, combined low concentrations of δ-tocotrienol and jerantinine A caused a considerable disruption of microtubules in A549 cells. This may be attributed solely to the IC$_{50}$ of jerantinine A or could probably be potentiated by the presence of low-concentration δ-tocotrienol. To date, there is no study confirming the potentiation effect of a tocotrienol isomer on microtubules but this possibility cannot conclusively be ruled out. Nevertheless, the study has demonstrated an improved potency of δ-tocotrienol thereby corroborating the previous combination studies on tocotrienols (3). Microtubule-targeting agents have been shown to induce apoptosis predominantly via the mitochondria pathway and caspase-independent pathway mediated via the activation of lysosomal cathepsin B (14).

**Table 1.** Combination index (CI) and dose reduction index (DRI) values of combined sub-effective concentrations of δ-tocotrienol (0-24 μg/mL) with jerantinine A at its fixed IC$_{50}$ concentration (0.35 μg/mL) on A549 cells

<table>
<thead>
<tr>
<th>Cells</th>
<th>Jerantinine A (JA, μg/mL) [used at IC$_{50}$]</th>
<th>New concentration of δ-tocotrienol (δ-T3, μg/mL) that induced 50% growth inhibition in combined treatment</th>
<th>CI</th>
<th>DRI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(Mean ± SEM)</td>
<td>(Mean ± SEM)</td>
<td>δ-T3 + JA</td>
<td>δ-T3</td>
</tr>
<tr>
<td>A549</td>
<td>0.35 ± 0.32</td>
<td>2.77 ± 1.09</td>
<td>0.47</td>
<td>4.48</td>
</tr>
<tr>
<td>MRC5</td>
<td>0.35 ± 0.32</td>
<td>5.57 ± 1.22</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

DRI represents the fold decrease in required potent concentration to induce 50% cell growth inhibition following a combined treatment. Pharmacological interaction between the combined treatment is reflected by either synergistic (CI < 1), antagonistic (CI > 1) or additive (CI = 1) effects.
On the other hand, tocotrienols have been shown to induce apoptosis via death receptor and mitochondria pathways, although the roles of caspase enzymes and caspase-independent apoptosis vary depending on the cellular microenvironment (8,15). Herein, the individual doses of δ-tocotrienol and jerantinine A induced caspase 8, 9 and 3 activities. In fact, an improved caspase 9 and 3 enzymatic activity was evident for the combined sub-effective treatment compared to individual treatments. This suggests a potentially improved potency following the combined treatment. Death receptor and mitochondria pathways of apoptosis are mediated via caspase 8 and 9 enzymatic activities, respectively and both pathways can be linked at caspase 3 following the truncation of BID protein (12). The different patterns of cell cycle arrest and anti-microtubular effects induced by single-compound treatments of δ-tocotrienol and jerantinine A as well as combined treatment strongly suggest that the combined treatment may be mediated via multi-targeted pathways. This conforms to a recent study that demonstrated the cytotoxic potency of combined low-concentration treatment of γ-tocotrienol and jerantinine A, which was also able to activate caspase 8, 9 and 3 enzymatic activities and disrupt microtubules in a different cell type, i.e. brain cancer (6). This suggests that the combination of different isomers of tocotrienol such as γ and δ with jerantinine derivatives might induce synergistic apoptotic effects against a broad spectrum of cancer cell lines. However, it is worthwhile to emphasize that the current combined δ-tocotrienol and jerantinine A treatment induced

Figure 1. Evaluation of cellular morphologies and cycle arrests in A549 cells. (a) Microscopic observations on H&E-stained A549 cells receiving the single-compound treatments with IC\textsubscript{50} of δ-tocotrienol, jerantinine A, and vinblastine as well as combined treatment involving 2.77 μg/mL of δ-tocotrienol and 0.35 μg/mL of jerantinine A for 24 hours. Cell cycle profiles show (b) representative flow cytometric images and (c, d) percentages of A549 cells in each phase. A549 cells were incubated with plain media containing DMSO (untreated control), IC\textsubscript{50} of δ-tocotrienol, jerantinine A and combined sub-effective concentrations of δ-tocotrienol with IC\textsubscript{20} of jerantinine A for (c) 24 hours and (d) 48 hours. Single-compound IC\textsubscript{50} of δ-tocotrienol and jerantinine A induced G0/G1 and G2/M cell cycle arrests, respectively. The combined sub-effective concentrations of δ-tocotrienol with IC\textsubscript{20} of jerantinine A induced G2/M cell cycle arrest. P $<$ 0.001 (***) and P $<$ 0.05 (*) indicate a statistical significant difference between treated and untreated groups. Abbreviation: h = hour(s).
higher caspase 8 and 9 enzymatic activities in lung cancer cells than those observed in brain cancer cells treated by γ-tocotrienol and jerantinine A combination in the previous study (6). This even strengthens the speculation that the induction of apoptosis on A549 cells by current combined treatment is mediated via both death receptor and mitochondria pathways (12).

**Conclusion**

This study showed that the combined sub-effective concentrations of δ-tocotrienol and jerantinine A caused inhibition of microtubular networks and activation of caspase enzymatic activities in addition to the established mechanism of action of δ-tocotrienol. This demonstrates an improved potency of δ-tocotrienol. Besides, the sub-effective concentrations used in the combined treatment have minimized toxicity to MRC5 cells. Yet, this still necessitates more mechanistic and in vivo investigations in order to confirm the putative mechanisms of action and further develop the combined treatment against lung cancer.

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Authors’ contributions
IBA was involved in the experiment design, conducted the experiments and drafting of manuscript. HSL was involved in experimental design, manuscript corrections, proof reading and data analysis. All read and confirmed the final version of the manuscript.

Conflict of interest
The authors declare that there are no conflicts of interest.

Ethical considerations
All ethical issues have been checked by the authors and the study complies with ethical considerations.

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References