Cytotoxic Evaluation of Plant Essential Oils in Human Skin and Lung Cells

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ABSTRACT

Plant essential oils are defined as fragrant volatile oils extracted from leaves, stems, fruits, flowers, and roots of a plant. Such oils are composed of multiple components and multiple functions. By accumulation of inductive information, various plant essential oils have been studied for using in therapeutic medicine for various diseases. Despite of the apparent advantages of essential oils as a source of therapeutic medicines, plant essential oils have many limitations, including cytotoxic side effects. Therefore, it is necessary to evaluate the toxicity and the mechanisms of cytotoxicity of such oils. In this study, we evaluated the cytotoxicity to human-derived cell lines of 10 plant essential oils provided by National Institute of Forest Science (i.e., Larix kaempferi; Abies holophylla; Zanthoxylum ailanthoides; Pinus parviflora; Tsuga sieboldii; Chamaecyparis pisifera; Cryptomeria japonica; Pinus densiflora; Illicium anisatum; Pinus thunbergii). Cytotoxicity evaluations were accomplished by using CCK-assays and PCR-based cytotoxicity-related marker gene analyses with A549 cell line, and the Detroit551 cell line which are lung and skin cell line. The genes were analyzed included caspase-3 has a role in cell apoptosis, and the other cyclinA, cyclinB, cyclinD, and cyclinE regulated cell cycling for the cell proliferation. By examining the five cytotoxicity-related marker genes by performing real-time PCR and examined the cytostatic gene regulation associated with the various essential oils. The results of this study showed that the degree of cytotoxicity and the cytostatic gene regulation which could give precious information for using the plant essential oil for the clinical usages.

Keywords: essential oil, cytostatic effect, cytotoxicity

1. INTRODUCTION

A plant essential oil is a volatile natural compound that is obtained as a secondary product of the plant and is characterized by a strong smell (Galindo et al., 2010). The components of a plant essential oil depend on the species of plant, and of the plant (Kahkonen et al., 1999; Lee et al., 2014). Oils may be extracted from different parts of the plant, such as leaves, fruit peels, seeds, bark, wood wicks, and flowers, and are usually obtained via steam or hydro-distillation (Waseem and Low, 2015). Over the centuries, plant
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essential oils have been interested as sources of natural products (Prabuseenivasan et al., 2006). Plant essential oils have been widely used in the food and perfume industries until now due to their fragrance and flavor (Lee et al., 2017). Extracts having characteristics that can kill insects have been used as insecticides, while extracts having antibiotic and antifungal activity have been used as alternative treatments for diseases (Zhang et al. 2016). However, despite the great interest in the long-term usage of plant essential oils, science-based research into essential oils has been not revealed (Lambert et al., 2001; Sienkiewicz et al., 2014).

Cytotoxicity was determined by using CCK-assays and PCR-based cytotoxicity-related marker gene analyses. The genes analyzed included caspase-3, which has a role in cell apoptosis, and cyclinA, cyclinB, cyclinD, and cyclinE, which have roles in each stage of the cell cycle. Caspase-3 is a mammalian homolog of the ced-3 gene and has an essential role in the execution of cell apoptosis (Namura et al., 1998). In mammals, caspasas (mainly caspase-3) are reported to be activated in the proteolytic cascade, leading to inadequate activation or rapid inactivation of key structural proteins, significant signal transduction, homeostasis, and repair enzymes (Porter and Janicke, 1999).

The cell cycle is divided into four stages; (Bravo and Celis, 1980) An S phase in which DNA is synthesized, an M phase in which mitosis occurs, and the G1 and G2 phase occur between S and G phases (Elledge, 1996). The signaling pathway regulating the progression of the cell cycle is known to be mainly related to the G1 phase (Grana and Reddy, 1995; Elledge, 1996). Cyclins are thought to be involved in interactions with upstream regulatory proteins, subcellular localization, timing of activation of Cdks, and substrate specificity and its catalytic subunit (Ohtsubo et al., 1995).

In this study, we suggest guidelines for the degree of toxic concentrations in Korean plant essential oils when we will apply in various fields. In this study, The essential oils evaluated for IC50 from larch (Larix kaempferi), fir (Abies holophylla), paulownia (Zanthoxylum ailanthoides), white pine (Pinus parviflora), tsuga (Tsuga sieboldii), cypress (Chamaecyparis pisifera), cedar (Cryptomeria japonica), red pine (Pinus densiflora), anise (Illicium anisatum), and black pine (Pinus thunbergii). We also determined the essential oil-induced cytostatic effect with four cell cycle markers (cyclinA, cyclinB, cyclinD, and cyclinE) and one cell death marker (caspase-3).

2. MATERIALS and METHODS

2.1. Extraction of Plant Essential Oils

Plant essential oils are extracted by hydrodistillation method. Add 1 kg of plant leaves and 10L of distilled water on heating mantle (MS-DM608, MTOPS®, Republic of Korea) at 102°C. Volatile steam is condensed in dean stark trap. Acquired essential oils are dehydration with sodium sulfite anhydrous, stored at 4°C.
2.2. CCK Assay

A549 and Detroit551 cells (Korean cell line bank, Republic of Korea) were seeded at a density of 0.5×10^3 cells per well and the medium used for cell culture was DMEM high glucose media (Biowest, France) supplemented with penicillin-streptomycin solution (Biowest, France), Plasmocin™ prophylactic (Biowest, France) and fetal bovine serum. Cells were incubated in a CO₂ atmosphere at 37°C for 24 h prior to treating with essential oil. The plant essential oils were prepared in four concentrations 10⁻⁸, 10⁻⁶, 10⁻⁴, and 10⁻². After 24 h, the cells are treated with diluted plant essential oil and incubated for 24 h. Subsequently, after removing the medium with the plant essential oil from cells, the cells are washed with PBS solution (WELGENE, Republic of Korea). EZ-Cytox reagent (DoGenBio, Republic of Korea) was used for evaluating cell viability. The absorbance was measured at 450 nm with Epoch microplate spectrophotometer (BioTek, USA). Cell viability (%) was determined by calculating the optical density (OD) by using the formula OD sample / OD control (vehicle)×100 for each concentration range. A cell survival curve was calculated from these values, and the IC₅₀ value was established. After conducting the CCK assay, the same procedure was repeated with plant essential oils that were diluted to concentrations of 10⁻⁶, 10⁻⁵, 10⁻⁴, 10⁻³, and 10⁻².

2.3. RNA Extraction

Cells were seeded in 6-well plates at a density of 0.3×10⁻⁵ cells per well and incubated at 37°C in 5% CO₂ atmosphere. Plant essential oils in DMEM high glucose media (Biowest, France) were prepared in three concentrations: IC₅₀ from the CCK assay, 10 times more than the IC₅₀, and 10 times less than the IC₅₀. When cell confluence was 70%, the cells were treated with the appropriate concentration from preliminary experiment (10⁻⁵–10⁻³ or 10⁻³–10⁻²) of plant essential oil. Harvesting of cells and RNA extraction were accomplished by using TRI reagent (Invitrogen, USA) at 24 h after plant essential oil treatment. The purity and concentration of the extracted RNA were measured by using an Epoch microplate spectrophotometer (BioTek, USA).

2.4. Synthesis of cDNA

The extracted RNA was synthesized by reverse transcription. The concentration of RNA was titrated to 1 µg/µL and DEPC-treated water was added to adjust the total volume to 10.5 µL. The mixture contained Moloney murine leukemia virus (MMLV) reverse transcriptase (Invitrogen, USA), Ribonuclease inhibitor (Invitrogen, USA), random primers (6-mers; TaKaRa, Japan), 100 mM DTT (Invitrogen, USA), dNTP, and 5×first strand buffer. The reaction mixture was incubated at 37°C for 60 min. After synthesizing cDNA, the enzyme mixture were incubated at 95°C for 5 min for inactivation. After cooling for 5 min on ice, the cDNA was stored at -20°C prior to use.

2.5. Real-time Polymerase Chain Reaction

For real-time polymerase chain reaction (RT-PCR), a mastermix of the following reaction components was prepared: cDNA sample (2 µL), 0.25 µL 50× ROX dye, 6.25 µL 2× prime Q-mastermix SYBR (GeNetBio, Korea), 0.5 µL forward primer, and 0.5 µL reverse primer. RT-PCR was performed using 7300 RT-PCR system (Applied Biosystems, Foster City, CA, USA) and the procedure sequence was as follows: cDNA denatured at 95°C for 30 sec and annealed at 58°C for 30 sec, elongation is performed at 72°C for 30 sec, and the whole cycle is repeated 40 times. The degree of gene expression was determined by using RQ software (Applied Biosystems).

The primer sequences used are as follows: 5’-CATA CTCCACAGCACCCTGGTA-3’ (forward) and 5’-CTGT
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TGCCACCTTTCGGTGA-3’ (reverse) for caspase-3; 5’-TACTTTCTGACATCAGCAGCC-3’ (forward) and 5’-TGATTCAAGGCAAGCTTTGTC-3’ (reverse) for cyclinA; 5’-CTTTCGACTTCTCAGAGA-3’ (forward) and GTAGAGTTGGTGTCCATTCACC-3’ (reverse) for cyclinB; 5’-CTCGGTGCCTACTTTCA-3’ (forward) and 5’-CTCCTCGCATTCTGGTC-3’ (reverse) for cyclinD; and 5’-GGTTTCAGGGRATCAGTGGT-3’ (forward) and 5’-TTTCTTTTGCTGGGGCTTTG-3’ (reverse) for cyclinE. The primer sequence for Rn18s was used as the primer for the control group: 5’-CTCAA CACGGGAAACCTCAC-3’ (forward) and 5’-CGCTCACCAACTAAGAACG-3’ (reverse). All primers were synthesized from Macrogen (Republic of Korea).

2.6. Data Analysis

All data were analyzed by applying nonparametric one-way analysis of variance followed by Tukey’s test for multiple comparisons. All experiments consisted of three separate trials. Statistical analysis was performed by using Prism Graph Pad (v.4.0; GraphPad Software, USA).

3. RESULTS and DISCUSSION

3.1. Two-step Identification of ICso Value of Each Essential Oils

ICso values were obtained by performing CCK-8 assays. To determine the test concentration, the cells were treated with four concentrations of CCK-8, followed by further subdivision within the CCK-8 assay. The ICso values for the 10 plant essential oils tested are presented in Figs. 1 and 2. In each cell line, cytotoxic degree of each essential oil was classified as low toxicity (~1×10^-4), intermediate toxicity (~1×10^-5), or high toxicity (1×10^-6~). The A549 cell line showed
low cytotoxicity to *L. kaempferi*, *Z. ailanthoides*, and *C. pisifera* treatment, while *A. holophylla*, *P. parviflora*, *T. sieboldii*, *C. japonica*, *P. densiflora*, *I. anisatum*, and *P. thunbergii* treatments were intermediately toxic to A549 cells. In the Detroit551 cell line, *A. holophylla*, *Z. ailanthoides*, *P. parviflora*, *T. sieboldii*, and *C. pisifera* produced low cytotoxicity, and *L. kaempferi*, *C. japonica*, *P. densiflora*, *I. anisatum*, and *P. thunbergii* were moderately toxic.

### 3.2. Expression of Caspase-3 Gene Affected by Plant Essential Oils

Expression levels of caspase-3 gene after essential oil treatment of A549 and Detroit551 cells, a marker gene related to cell apoptosis, were determined by performing RT-PCR. The expression level of caspase-3 in the A549 and Detroit551 cell line after treatment with plant essential oil is shown in Fig. 3. The expression level of caspase-3 gene decreased in high concentrations of *Z. ailanthoides*, *C. japonica*, and *I. anisatum*, while it increased in low concentration *T. sieboldii*. The expression of caspase-3, a cytotoxicity marker, increased at concentrations near the IC₅₀ values of *L. kaempferi*, and *Z. ailanthoides* in Detroit551 cell line.

### 3.3. Expression of CyclinA, CyclinB, CyclinD, and CyclinE Affected by Plant Essential Oils

The expression levels of cyclinA, cyclinB, cyclinD, and cyclinE in the A549 cell as plant essential oil treatment at different concentrations are shown in Fig. 4. The concentrations of essential oils are determined...
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Fig. 3. The expression level of Caspase-3 in A549 and Detroit551 cells after treatment with essential oil. Relative expression of caspase-3 in A549 cell (A), and Detroit-551 cell (B). Based on the IC_{50} values obtained above, three concentrations were set so that the difference was 10 times back and forth. The expression levels of Caspase-3 in each group were determined by real time polymerase chain reaction after treatment with three concentrations of essential oil. *P < 0.05 vs. vehicle. Data are presented as mean±SD values.

with IC_{50} value from CCK assay. The cell-cycle control gene, cyclinA, showed a concentration-dependent decrease in all samples except C. pisifera. CyclinB showed a concentration-dependent decrease in cell-cycle marker protein in all 10 oils tested. In the A549 cell line, cyclinD expression was significantly decreased in Z. ailanthoides, C. japonica, and I. anisatum oils, while cyclinE expression showed a decrease in all species except P. thunbergii. Expression of cyclinE was observed at a concentration approximately 10 times the IC_{50} value when treated with P. thunbergii oil. The expression levels of cyclinA, cyclinB, cyclinD, and cyclinE in the Detroit551 cell line according to plant essential oil concentrations are shown in Fig. 5. CyclinA, a cell-cycle regulator, showed a concentration-dependent decrease in Detroit551 cells for all species except C. pisifera. This tendency was the same as the A549 cells, a lung cell line. CyclinB analysis showed a concentration-dependent decrease in all species except C. pisifera. The cyclinD results showed that expression of the cell-cycle marker protein was significantly different among several oils. Extracts of L. kaempferi, Z. ailanthoides, C. japonica, and I. anisatum showed a concentration-dependent decrease, while the other species did not show a decreasing tendency. The expression level of cyclinE showed that there was a significant decline in L. kaempferi, T. sieboldti, C. japonica, and I. anisatum.
Fig. 4. The expression level of CyclinA, CyclinB, CyclinD, CyclinE in A549 cells after treatment with essential oil. Based on the IC₅₀ values obtained above, three concentrations were set so that the difference was 10 times back and forth. The expression levels of CyclinA (A), CyclinB (B), CyclinD (C), CyclinE (D) in each group were determined by real time polymerase chain reaction after treatment with three concentrations of essential oil. *P < 0.05 vs. vehicle. Data are presented as mean±SD values.
Fig. 5. The expression level of CyclinA, CyclinB, CyclinD, CyclinE in Detroit551 cells after treatment with essential oil. Based on the IC$_{50}$ values obtained above, three concentrations were set so that the difference was 10 times back and forth. The expression levels CyclinA (A), CyclinB (B), CyclinD (C), CyclinE (D) in each group were determined by real time polymerase chain reaction after treatment with three concentrations of essential oil. *P < 0.05 vs. vehicle. Data are presented as mean ± SD values.
3.4. Discussion

Essential oils have been used in traditional medicine. Researches in this area is growing by systematic and scientific approaches for the biological activities of phytotherapeutics targets in last few decades. The availabilities of essential oils seem to have a great potential as anti-inflammatory, anti-bacterial, anti-cancer therapeutic agents and aromatherapies (Bravo and Celis, 1980; Pagano et al., 1992; Elledge, 1996; Min et al., 2017). Their complex chemical composition makes it difficult to examine the biological effect, which is likely the show biological activity of each component. However, information regarding their mechanism of action is still lacking and far from being deciphered. Indeed, it was difficult that the mechanism and biological effects of complex chemical compositions in cell toxicity or cytostatic effects (Russo et al., 2015). As though, we tried to understand the effects of plant essential oils by CCK-8 assay and cytotoxicity-related marker gene analyses with two cell line A549 and Detroit551 for evaluation of their cytotoxic and antiproliferative actions.

The IC50 values determined by the CCK-8 assay showed low toxicities in the A549 cell line from *L. kaempferi*, *Z. ailanthoides*, and *C. pisifera* oils, whereas extracts of *A. holophylla*, *P. parviflora*, *T. sieboldii*, *C. japonica*, *P. densiflora*, *I. anisatum*, and *P. thunbergii* produced moderate toxicity. In the Detroit551 cell line, *A. holophylla*, *Z. ailanthoides*, *P. parviflora*, *T. sieboldii*, and *C. pisifera* produced low toxicity, while *L. kaempferi*, *C. japonica*, *P. densiflora*, *I. anisatum*, and *P. thunbergii* produced moderate toxicity. In addition to determining toxicity levels, we examined the expression levels of five marker genes to assess the cell death pathway. Expression of caspase-3 in the A549 cell line decreased under high concentrations of *Z. ailanthoides*, *C. japonica*, and *I. anisatum*, whereas *T. sieboldii* oil produced an increase in caspase-3 expression at low concentration. There were no concentration-dependent increases or decreases in the expression of caspase-3 gene in all tested essential oil samples; thus, confirming that the decrease in cell numbers of the lung cell line was not a result of apoptosis (Citron et al., 2000). Subsequently, we evaluate cell cycle regulator genes cyclin, which regulate cell proliferation. The expression level of cyclinA, a cell-cycle marker, in the A549 cell line is shown in Fig. 4. The concentration-dependent expression of cyclinA was decreased in all oil samples except that extracted from *C. pisifera*. This suggests that cytotoxicity through inhibition of cell proliferation occurred in all essential oils except that from *C. pisifera*. The expression level of cyclinB is shown in Fig. 4, which shows that the expression of cyclinB decreases in a concentration-dependent manner in all species. This suggests that the cytostatic effect of these oils in lung cell lines is due to inhibition of cell proliferation rather than to apoptosis (Jeong et al., 2009). Unlike cyclins A, B, and E, the expression of cyclinD was markedly decreased when treated with *Z. ailanthoides*, *C. japonica*, and *I. anisatum*. Therefore, it is thought that cytotoxicity results at high concentration of those essential oils is due to inhibition of specific phases in the cell cycle (Schmidt et al., 2002). The expression of cyclinE showed a concentration-dependent decrease in all oils except *P. thunbergii*. Decrease of cyclinE expressions were observed at a concentration approximately 10 times the IC50 values of *Larix kaempferi*, *Tsuga sieboldii*, *Cryptomeria japonica*, *Pinus densiflora* and *Illicium anisatum*. These results suggest that different types of essential oils affect different phases of the cell proliferation.

The expression level of caspase-3 in the Detroit551 cell line increased at concentrations near the IC50 values for *L. kaempferi*, and *Z. ailanthoides*. Therefore, when essential oil extracts of *L. kaempferi* and *Z. ailanthoides*
at a high concentration are treated to Detroit551 cells, the decrease in cell numbers would be the result of apoptosis (Namura et al., 1998). These results are opposite to the result with A549 cell line. In previous researches, cytotoxic effect due to apoptosis could be shown depends on cell type (Russo et al., 2015). Expression of cyclinA in the Detroit551 cell line decreased following treatment by all oils except that of C. pisifera; a similar trend was observed in the A549 cell line. CyclinB expression in the Detroit551 cell line showed a concentration-dependent decrease after treatment by all oils except C. pisifera oil. Cytotoxicity of the essential oils of these species in skin cell lines is due to inhibition of cell proliferation rather than due to apoptosis (Goldwasser et al., 1996). The results of the cyclinD analysis showed the greatest differences among the tested species; L. kaempferi, Z. ailanthoides, C. japonica, and I. anisatum showed a concentration-dependent decrease, while the other species did not show a decreasing tendency. Finally, analysis of cyclinE showed that there were significant decreasing trends following treatment with L. kaempferi, T. sieboldii, C. japonica, and I. anisatum oils.

In conclusion, cell growth is induced by cyclinD which is required for all cell cycle stage, cyclinE for G1/S transition, cyclinA for G2 phase, cyclin B for Mitosis. Phosphorylation of pRB proteins by the cyclin-dependent kinases (CDKs) releases E2F factors, promoting the transition to next cycle (Duronio and Xiong 2013). Regulation of cyclin could be used for cancer treatment with carefully chosen dosage. But in ahead of usage, cytotoxic evaluation (identifying IC50). Expressions of all marker genes in the A549 cell line after treatment with the plant essential oils of Z. ailanthoides, C. japonica, and I. anisatum were down-regulated. After L. kaempferi, A. holophylla, P. parviflora, and T. sieboldii treatment, only cyclinA, cyclinB, and cyclinE were down-regulated. Down-regulation of cyclinB and cyclinE was observed in C. pisifera treatment, and down-regulation of cylinA and cyclinB was observed in P. thunbergii treatment. In the Detroit551 cell line, changes in expression level of caspase-3 were observed in two species' essential oils: L. kaempferi, and Z. ailanthoides. CyclinA, cyclinB, cylinD, and cylinE gene markers were all down-regulated by L. kaempferi essential oil treatment, whereas caspase-3 genes were down-regulated in three genes, cyclinA, cyclinB, and cyclinD. Essential oils of A. holophylla, P. parviflora, P. densiflora, P. thunbergii affected only two genes, cyclinA and cyclinB, which were down-regulated. CyclinA, cyclinB, and cyclinE were down-regulated by T. sieboldii, and all five marker genes were down-regulated by C. japonica and I. anisatum essential oils.

4. CONCLUSION

With those results, in turn, essential oils can regulate depends on its own chemical contents and concentration, the right selection of kind and concentration of essential oils are important for the source of medicine and cosmetic usage.

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