Overexpression of Anti-apoptotic Molecules and Bax Translocation to Mitochondria by Pharbitis Nil Extracts in AGS

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Conventional medicines have usually sorted to a number of treatments such aseoperation, radiotherapy, and chemotherapy. The existing anti-cancer agents, designed to eradicate cancer cells, have strong toxicities, also with leading to harmful side effects. Recently, a number of researches on natural products have been actively carried out in efforts to develop new treatments that can decrease side effects or increase anti-cancer effects. We performed this study to understand the molecular basis underlying the antitumor effects of Pharbitis nil, and Plantago asiatica, which have been used for herbal medicinal treatments against cancers in East Asia. We analyzed the effects of these medicinal herbs on proliferation and on expression of cell growth/apoptosis related molecules, with using an AGS gastric cancer cell line. The treatment of Pharbitis nil dramatically reduced cell viabilities in a dose and time-dependent manner, but Plantago asiatica didn’t. FACS analysis and Annexin V staining assay also showed that Pharbitis nil induce apoptotic cell death of AGS. Expression analyses via RT-PCR and Western blots revealed that Pharbitis nil didn’t increase expression of the p53 and its downstream effector p21waf1, and that the both increased expression of apoptosis related Bax and cleavage of active caspase-3 protein. We also confirmed the translocation of Bax to mitochondria. Collectively, our data demonstrate that Pharbitis nil induce growth inhibition and apoptosis of human gastric cancer cells, and these effects are correlated with down- and up-regulation of growth-regulating apoptotic and tumor suppressor genes, respectively.

Key words: Pharbitis nil, apoptosis, anti-tumor effect, Bax, medicinal herbs

Introduction

Pharbitis nil, and Plantago asiatica are representative anticancer medicinal herbs that have been traditionally used for cancer treatments in China, Japan and Korea. Previous studies also demonstrated that Pharbitis nil exhibits anti-tumor and antifungal effects by antifungal peptides7. Plantago asiatica has been widely used to treat for cancers and liver diseases and were reported to carry immune suppressive effects.2,3

It is well known that tumor development is accelerated by disruption of the balance between cell proliferation and cell death, which is maintained through regulations of various signal transduction pathways.5,6 Active cell death, known as apoptosis or programmed cell death, is caused by various physiologic and non-physiologic cell injuries including DNA damage. It has been demonstrated that various cell proliferation-and apoptosis-signal transduction pathways are built on complicated networks between oncogenes and tumor suppressor genes such as p53 and its downstream factor p21.6,10 For example, transcription and translation of anti-apoptotic Bcl-2 and pro-apoptotic Bax are down-and up-regulated by p53, respectively.7,11

p53 controls various genetic expressions and plays an important role in cell proliferation and in modulation of signal transduction pathways. Accumulation of p53 in cells after DNA damage leads to cell cycle arrest and apoptosis induction. In addition, p53 is involved in repair of damaged DNA and thus prevents accumulation of mutations and suppresses tumor development.7,12

In our study, we analyzed effects of Pharbitis nil, and Plantago asiatica on proliferation and apoptosis of a human gastric cancer cell line AGS. Here we demonstrated first that Pharbitis nil induced growth inhibition and apoptosis of AGS cells and these effects are correlated with down-and up-regulation of growth-regulating pro-apoptotic and tumor...
suppressor genes, respectively, suggesting that the anticancer effects of *Pharbitis nil* might be associated with their regulatory capabilities of tumor-related genes expressions.

**Materials and Methods**

1. Cell culture

   The AGS human gastric carcinoma cell line was obtained from American Type Culture Collection (ATCC, Rockville, MD) and grown in RPMI 1640 (Life Technologies, Inc., Rockville, MD) containing 10% FBS (HyClone Laboratories, Inc., Logan, UT) and 1% gentamicin in a 5% CO₂ humidified atmosphere. Subconfluent monolayers of cells were employed in all experiments.

2. Preparation of medicinal herbs

   The raw herbs were purchased from Omni-herb Inc. (Seoul, Korea) and extracted [*Pharbitis nil* (570g), and *Plantago asiatica* (350g)] by sonication of dried roots of these plants in 80% ethanol, following by a freeze-drying process of the ethanol extract. Freeze dried powder forms of the extracts [*Pharbitis nil* (14.8g), and *Plantago asiatica* (82.5g)] were dissolved in RPMI 1640 medium (Life Technologies, Inc.) to 10 mg/ml, vortexed at room temperature for 1 min, and incubated at 37°C for 1 hr while rotating before use. These solutions were centrifuged at 12,000 rpm for 5 min to remove any insoluble ingredients. The supernatant was passed through a 0.22-μm filter for sterilization and diluted with RPMI 1640 culture medium to final concentrations of 6.25 - 1,000 μg/ml.

3. Growth inhibition assay

   To determine the inhibition effect of these four herbal extracts on proliferation of AGS cells, the percentage of growth inhibition was determined by measuring MTT dye absorbance of viable cells in the absence or presence of medicinal herb extracts. Ten thousand cells per well were seeded onto a well of 96-well plates (Nunc, Roskilde, Denmark) for 24 h, treated with various concentrations of these extracts, and incubated for 3 days at 37°C. Subsequently, 50 μl of MTT (Sigma) at a concentration of 2 mg/ml was added to each well, and cells were incubated for an additional 4 h at 37°C. The supernatant was aspirated, and 150 μl of DMSO were then added to the wells to dissolve any precipitate present. The absorbance was then measured at a wavelength of 570 nm using an ELX800 microplate reader (Bio-Tek Instruments, Inc., Winooski, VT). The IC₅₀'s were calculated assuming the survival rate of untreated cells to be 100%. To examine the time-dependent inhibition effects, cells were treated with *Pharbitis nil*, which shown cytotoxic effects at the various times, cell viability was measured by a Trypan blue exclusion assay. To evaluate how much these herbal medicines have cytotoxic effects to cancer cells as compared with normal epithelial cell line, AG5 and RIE1 cells, from rat intestine, were seeded to wells of 96 well plates in the presence of normal culture media at 1.0 x 10⁶ cells/well. Twenty four hours later, cells were treated with indicated concentrations of *Pharbitis* nilextracts for additional 72 hrs. Then MTT assay was performed as explained in Material and methods. Data shown is representative from three independent experiments, in which each condition was in triplicate. Data were shown in mean standard deviation (SD).

4. Flow cytometric cell cycle or DNA content analysis

   A total of 5 x 10⁵ cells were seeded in 60mm dishes and incubated for 24 h at 37°C. *Pharbitis nilextracts at indicated various concentrations was directly added to the dishes and incubated for an additional 12, 24, 48 hrs. During harvests, both cells detached (probably apoptotic) and adherent were combined, fixed by addition of 4ml 70% ethanol, and stored at -20°C at least 30 min. Cells were then pelleted, washed twice with ice-cold PBS, incubated in PBS containing 10 μg/ml of RNase A (Sigma) for 15 min at 37°C, and stained with 10 μg/ml of propidium iodide (PI). The relative DNA content per cell of samples was obtained by measuring the fluorescence of PI that bound stoichiometrically to DNA. The cell cycle was analyzed using a FACSCalibur flow cytometer (Becton Dickinson, San Jose, CA) and a ModFit LT V2.0 computer program.

5. Apoptosis assay

   To evaluate the apoptotic cell death, annexin V staining was carried out using an annexin V assay kit (PharMingen) according to the manufacturer’s recommendations. In brief, both floating and adherent cells were collected. Prepared cells were washed twice with ice-cold PBS and then resuspended in binding buffer (10 mM HEPES/NaOH, pH 7.4, 140 mM NaCl, 2.5 mM CaCl₂). Both 5 μl of annexin V-FITC and 10 μl of 20 μg/ml PI were then added to these cells, which were later analyzed with a FACSCalibur flow cytometry (Becton Dickinson, San Jose, CA).

6. DNA fragmentation assay

   Cells were collected, washed with PBS twice and then lysed in 100 ul of lysis buffer (50 mM Tris (pH 8.0), 10 mM EDTA, 0.5% sodium lauryl sarcosinate and 1 mg/ml protease K) for 3hr at 56°C and treated with 0.5 μg/ml RNase A for an
additional 1 hr at 56°C. DNA was extracted by the phenol/chloroform/isoamyl alcohol (25/24/1) method before loading. Loading buffer (50 mM Tris, 10 mM EDTA, 1% (w/v) low melting point agarose, 0.25% (w/v) bromphenol blue) and samples were loaded onto a prepsolidified, 2% (w/v) agarose gel containing 0.1 ug/ml ethidium bromide. Agarose gels were electrophoresed at 50 V for 90 minutes in TBE buffer. Gels were observed and photographed under ultraviolet light.

7. Quantitative RT-PCR analysis of gene expression

Total cellular RNA was extracted from cultured cells by a single-step method. One ug of extracted RNA was reverse-transcribed to cDNA in a 20 ul reaction using MoMuLV (Gibco) and random hexamer primers. Two separate cDNAs were prepared from each RNA and diluted 1:4 or 1:8 with distilled, sterile H2O prior to PCR. For quantitative evaluation of gene expression, we initially performed PCR with exon-specific primer sets for all targets including a housekeeping standard GAPDH. The sequences of oligonucleotide primers will be provided upon request. PCR was performed with increasing cycle numbers (21, 24, 27, 30, 33, 36, 39, and 42 cycles) and diluted cDNAs (1.0, 1.2, 1.4, and 1.8). Each cycle comprised denaturation at 95°C for 1 min, annealing at 58-62°C for 1 min, and polymerization at 72°C for 1 min. Ten ul of the PCR products were resolved on 2% agarose gels (FMC, Rockland, ME). Quantitative analysis of gene expression was confirmed through scanning of ethidium bromide-stained gels, using a laser densitometry. Measurement of signal intensity was performed using the Molecular Analyst program (version 2.0) on an IBM compatible computer.

8. Western blot analysis

AGS cells in 100 mm dishes were treated with or without Pharbitis nil extracts for indicated periods. After incubation, cells were washed with ice-cold PBS and lysates were prepared using a lysis buffer containing 20 mM Tris-Cl (pH 7.4), 100 mM NaCl, 1% NP40, 0.5% sodium deoxycholate, 5 mM MgCl2, 0.1 mM phenylmethylsulfonyl fluoride, 0.1 mM pepstatin A, 0.1 mM antipain, 0.1 mM chymostatin, 0.2 mM leupeptin, 10 µg/ml aprotinin, 0.5 mg/ml soybean trypsin inhibitor, and 1 mM benzamidine. After incubation of the lysates on ice for 30 min, whole cell extracts were cleared by a centrifugation at 13,000 rpm for 20 min. Twenty ug of protein were fractionated by SDS-PAGE denaturing gels and transferred onto a nitrocellulose membrane. The membrane was blocked for 1 h in the 20 mM Tris-buffered saline (TBS) buffer containing 5% skim milk and 0.1% Tween 20 and then probed with specific antibodies for indicated molecules. The protein was detected using chemiluminescence method (Amersham Pharmacia Biotech) followed by autoradiography.

9. Subcellular fractionation

Cells were lysed in isotonic mitochondrial buffer in mitochondria lysis buffer (210 sucrose, 70mmanit, 10mM Hepes, pH 7.4,EDTA) containing 1 mM phenylmethylsulfonyl fluoride, 5 ug/ml leupeptin, 5 ug/ml aprotinin, and 0.7 ug/ml pepstatin. After homogenization with a Dounce homogenizer, cell lysates were centrifuged at 1,000 x g for 10 to discard nuclei and unbroken cells. The postnuclear supernatant was centrifuged at 10,000 x g for 15 to pellet mitochondria-enriched heavy membrane fraction, and the resulting supernatant was further centrifuged at 100,000 x g for 30 to obtain cytosolic fraction. The membrane fractions were resuspended in Triton X-100 lysis buffer containing protease inhibitors. Protein concentration was determined by BCA assay (Pierce Chemical, Rockford, IL) and total proteins (50 ug) from each fraction were subjected to immunoblot analysis.

10. Data analysis

Results shown are representative of at least three independent experiments performed in triplicate and are presented as the means ± standard deviation (SD).

Results

1. Growth inhibition of AGS gastric cancer cells by Pharbitis nil, and Plantago asiatica

To determine treatment concentrations, we initially examined effects of two medicinal herbs on cell viability using MIT assay. The growth inhibition of AGS by the Pharbitis nil, and Plantago asiatica extracts were determined by quantifying viable cells in the absence or presence of treatment of each herbal extract at various concentrations for 3 days. As shown in figure 1A, Pharbitis nil extracts significantly decreased cell viability in a dose-dependent manner. Based on the growth inhibition curves, the concentration of Pharbitis nil extracts was required for 50% inhibition of growth (IC50) for AGS cells was about 12.5 ug/ml. The maximal inhibition of cell growth (>80%) were achieved at 60 ug/ml in AGS cells. A time-dependent growth inhibition of AGS cells by Pharbitis nil extracts were also observed at the various times, and maximal growth inhibitions were shown within 2 days in Pharbitis nil extracts after the treatments (Fig. 1B). Meanwhile, the parallel treatment of the extracts to a normal epithelial cell line from rat intestine (RIEI) showed much less strong effects on inhibition of
viability (with an IC at about 80 μg/ml, respectively, Fig. 1C). Therefore, Pharbitis nil extracts could induce growth inhibition of gastric cancer cells such as AGS cells.

Fig. 1. Medicinal herb-mediated inhibition of cell viability in gastric AGS cells. (A) inhibition of cellular growth by four medicinal herbs in a dose-dependent manner. AGS cells seeded to wells of 96 well plates in the presence of normal serum containing media at 1.0 x 10^5 cells/well. Twenty-four hours later, cells were treated with indicated concentrations of four medicinal herbal extracts for further 72 hrs. Then MTT assay was performed as explained in Material and methods. (B) inhibition of cellular growth by four medicinal herbs in a time-dependent manner. AGS cells seeded to 60 mm culture dishes in the presence of normal serum containing media at 5 x 10^5 cells/dish. Twenty-four hours later, cells were treated with 12.5 μg/ml of Pharbitis nil extracts for the indicated time. Then Trypan blue exclusion assay was performed and viable cell counted. (C) Inhibition of AGS cell growth by the Pharbitis nil extracts. AGS or RPE1 cells were seeded to wells of 96 well plates in the presence of normal culture media at 1.0 x 10^5 cells/well. Twenty-four hours later, cells were treated with indicated concentrations of Pharbitis nil extracts for additional 72 hrs. Then MTT assay was performed as explained in Material and methods. Data shown is representative from three independent experiments. PN, Pharbitis nil PA, Panax ginseng.

2. Cell cycle analysis of Pharbitis nil

Based on the results from the preliminary study, two medicinal cytotoxic herbs, Pharbitis nil extracts were chosen to examine their cytotoxic effects. To further characterize whether these herbs affect apoptotic population after the treatments, cells were treated with 12.5 μg/ml concentrations, respectively, for 12 to 48 hours, before flow cytometry analyses. While no detectable changes in cell cycle distribution were observed until 48 hours after treatment, significant increases of cell population with subG1 DNA content, as time passed after treatment of Pharbitis nil extracts were found at 24, and 48 hours (Fig. 2A and 2B). However, cells in control condition showed no significant apoptotic population.

Fig. 2. Treatment of Pharbitis nil extracts induced apoptosis. (A) Pharbitis nil extracts induced apoptosis. Cells in 60 mm culture dishes were treated with Pharbitis nil extracts at 12.5 μg/ml for indicated periods. The treatments were done by a direct addition of Pharbitis nil extracts solution into culture media. The solution was made of RPMI1640 culture media as explained in the Materials and methods. After incubation, cells floating and adherent were harvested and combined before PI staining and flow cytometric analysis for subG1 population and cell cycle as explained in the materials and methods. (B) Apoptotic population by the treatments of Pharbitis nil were shown with histogram.

3. Annexin V staining assay and DNA fragmentation assay of Pharbitis nil

Next we tried to determine if Pharbitis nil extracts induced apoptotic cell death of AGS cells. As shown in fig. 2A and 2B, untreated cells did not show any significant apoptosis, whereas cells were becoming rapidly apoptotic with time after treatment with the extract. In addition to flow cytometric cell cycle analysis, we performed another approach to detect apoptotic cells using annexin V staining. In this assay, Pharbitis nil extracts showed apoptotic populations of about 13.75% and
17.15%, respectively, when cells were treated with 12.5 μg/ml and *Pharbitis nil* extracts for 24 h. To demonstrate apoptotic cell death by *Pharbitis nil* extracts, we also examined DNA fragmentation assay, and in this study, we got the evidence of DNA fragmentation of morphological changes of nuclei in AGS gastric cells.

![Fig. 3. The externalization of phosphatidylserine during Pharbitis nil-induced apoptosis.](image)

Fig. 3. The externalization of phosphatidylserine during Pharbitis nil-induced apoptosis. (A) Annexin V staining combined with PI staining was performed in control cells (left histogram) and cells treated with 12.5 μg/ml of *Pharbitis nil* (right histogram) extracts for the 24 hr and then analyzed by flow cytometry. Early apoptotic cells were localized in the lower right quadrant of a dotplot graph using Annexin V FITC versus PI/EID DNA fragmentation assay was examined to confirm the apoptotic change of nuclei after treatment of *Pharbitis nil* extracts, 12.5 μg/ml respectively, for 36 hr.

4. Transcriptional up-regulation of the tumor suppressor genes p53 and p21Waf1 expression

We next examined, using semi-quantitative RT-PCR assay, the possibility that growth inhibition and apoptosis induction by *Pharbitis nil* were associated with their regulatory effects on the cell cycle- and apoptosis-controlling genes expressions. As shown in Fig. 4, treatment of resulted in increase of transcription of the p53 tumor suppressor gene and its downstream effector p21Waf1 at 48 hours after the treatment. In contrast to *Pharbitis nil* also showed a strong growth arrest and apoptosis induction activity, but did not increase p53 and p21Waf1 expression. We also analyzed mRNA expression of p16INK4A, another tumor suppressor gene which also plays a critical role in regulation of cell cycle progression and apoptosis, but its expression was not modulated by any of the two herbal medicines tested. Next, we examined expression of representative anti-apoptotic genes, such as Bcl-2 and Bcl-XL, and a pro-apoptotic gene Bax. Whereas no effects on Bcl-2 and Bcl-XL expression were observed, *Pharbitis nil* showed a strong stimulating effect on Bax gene expression. Induction of the Bax gene transcription by *Pharbitis nil* was dependent on the treatment time. A representative example of gene expression analysis was shown in Fig. 4.

![Fig. 4. A semi-quantitative RT-PCR analysis for cell cycle- and apoptosis-related gene expression.](image)

5. Western Blot analysis

Based on the apoptotic analysis by determination of cells with subG1 DNA contents and RT-PCR analysis on cell cycle arrest- and apoptosis-related genes transcription, we have then examined expression levels of cell apoptotic molecules. In this analysis, we have confirmed intracellular apoptotic events biochemically, by examining the expression levels of pro-apoptotic molecules such as active caspase 3, and Bax, and an anti-apoptotic molecule of Bcl2. Cell lysates were prepared at various time points after treatment of *Pharbitis nil* extracts, and used for Western blots. As shown in the figure 5, the level of anti-apoptotic Bcl2 was not altered significantly by the both herbal extract treatments, but the expression of pro-apoptotic Bax increased gradually with the treatment. Inductions of p53 and its downstream targets including Bax and p21Waf1 were not obvious in case of *Pharbitis nil* treatment but showed obvious activation of caspase 3 (Fig. 5A). It is well known that apoptotic stimuli trigger Bax translocation to mitochondria where active Bax causes subsequent activation of certain downstream effector caspases 3 and its substrate PAPR cleavage. To examine Bax activation in response to *Pharbitis nil* extract treatment, we performed subcellular fragmentation assay that recognizes the translocation of Bax protein. In this subcellular fractionation/immunoblot analysis confirmed Bax
translocation to mitochondria in AGS cells treated with Pharbitis nil extracts (Fig. 5B). Taken together, apoptosis by the treatments of Pharbitis nil might be caused by their abilities in regulations of expression levels of growth arrest and apoptosis-related genes.

![Diagram of cell cycle and apoptosis](image)

**Discussion**

Pharbitis nil, and Plantago asiatica are representative herbal medicines that have traditionally been used for cancer treatments in East Asia, such as China, Japan and Korea. In the present study, we analyzed effects of Pharbitis nil, and Plantago asiatica on proliferation and apoptosis of a human gastric cancer cell line AGS. Our cell proliferation assay showed that Pharbitis nil strongly inhibited cell proliferation of AGS, whereas Plantago asiatica exhibited no detectable effects on cellular growth (Fig. 1A). Pharbitis nil also showed the inhibition effects of cellular proliferation in time-dependent manner and cancer cell specific cytotoxic effects (Fig. 1B, IC). Based on the results of cytotoxicities of the two medicinal herbs (i.e., Pharbitis nil), we have done further studies. In the cell cycle analysis of our study, Pharbitis nil extracts showed no detectable changes in cell cycle distribution after treatment, but significant increases of cell population with subG1 DNA content were observed at 24, and 48 hours (Fig. 2A and 2B). In the annexin V staining, Pharbitis nil extracts also showed apoptotic changes in AGS cell lines. We also performed DNA fragmentation assay, DNA fragmentation which means late marker of apoptosis, in this assay, after treatment of Pharbitis nil extracts for 36 hr, significant DNA fragmentations were detected in both medicinal herbs (Fig. 3A and 3B).

Being consistently, in our semi-quantitative RT-PCR analysis, Pharbitis nil did not alter the levels of p53 and p21 expression but Pharbitis nil showed a strong stimulating effect on transcription and induction of expression of Bax. Apoptosis is a genetically programmed event that can take place by a variety of internal or external stimuli and these signals are regulated by two distinct pathways, involving either death receptor (extrinsic) or mitochondria. In the mitochondrial pathway, a variety of death signals triggers the release of several pro-apoptotic proteins, in these proteins Bax is a crucial mediator of the mitochondrial pathway for apoptosis, and loss of this proapoptotic Bcl-2 family protein contributes to drug resistance in human cancers. Previous studies have shown that tumor cells with mutational inactivation of apoptosis-inducing genes, such as a tumor-suppressor gene p53, failed to respond to chemotherapeutic agent-induced DNA damage, and that a number of apoptosis-related genes such as Bcl-2 family members display both pro-survival and pro-apoptotic functions by forming homodimers or heterodimers. Bax, a tumor suppressor, mediates the p53-induced apoptosis and it increases sensitivity to chemotherapy-induced apoptosis. Meanwhile when Bcl-2 is activated or prevalent, apoptosis is prohibited. Abnormal overexpression of Bcl-2 has frequently been observed in many types of human cancers, and relative expression levels of Bcl-2 to Bax were reported to determine the sensitivity to apoptosis. In addition, p53’s DNA binding property and its ability of controlling gene transcription and transduction are usually lost by mutation in human cancers and p21 is highly activated by p53, and p53-induced p21 leads to arrest of the cell cycle and/or apoptosis.

According to our apoptotic analysis by determination of cells with subG1 DNA contents, annexin V staining and RT-PCR analysis, it is likely that the growth inhibition by Pharbitis nil extracts might involve apoptosis. When we also analyzed cyclinD1 levels after the treatments, growth inhibition by Pharbitis nil was not observed. We have confirmed the idea by analyzing intracellular events biochemically, such as the
expression of a pro-apoptotic molecule, Bax, and anti-apoptotic molecules including Bcl-2 and cleavage of active caspase 3 and also got evidence of Bax protein translocation to mitochondria by subcellular fragmentation. Collectively, in this subcellular fractionation/immunoblot analysis confirmed that cytotoxic effects of *Pharbitis nil* extracts in AGS cells were mediated by mitochondrial apoptotic pathway. And our observations thus suggests that anticancer effect of *Pharbitis nil* might be associated with their ability to regulate tumor-related genes expressions.

**References**