Molecular Medicine Reports 22: 2076-2084, 2020

Abstract. Abeliophyllum distichum Nakai is a Korean endemic plant of the Oleaceae family that contains acteoside, a glycosylated caffeic acid, with neuroprotective, anti-inflammatory and antibacterial properties. Previous studies, involving Accelerated Chromatographic Isolation, a high-performance liquid chromatography-photodiode array detector and a liquid chromatograph-mass selective detector, isolated and identified acteoside in A. distichum (aad) and documented its antioxidant and anti-inflammatory activities. The aim of the present study was to determine whether AAD could protect from DNA damage by reducing oxidative stress. AAD treatment protected plasmid DNA against damage to DNA double-strands induced by reactive oxygen species (ROS) and decreased the levels of phosphorylated p53 and γ-H2aX in ROS-treated NIH 3T3 cells. These findings suggested that AAD could reduce ROS-mediated cellular damage and may represent an effective, natural antioxidant with the ability to protect genetic material.

Introduction

Molecules obtained from medicinal plants have traditionally been used by people to maintain good health. According to the World Health Organization, >80% of the world's population use traditional medicinal plants to improve their health (1). A previous study has suggested that medicinal plants contain chemical components and exhibit biological activities that are useful for physiological processes including antioxidant, anti-inflammatory and anti-cancer activity in the body (2). The most widely used of these plant bioactive components are alkaloids, flavonoids and phenolic compounds (3,4). Due to their aromatic structure, phenolic compounds can act as electron donors, and this antioxidant potential can relatively reduce reactive oxygen species (ROS) levels (5). DNA is under constant stress resulting from environmental factors or cellular metabolism (6). ROS cause DNA damage through oxidative stress, thereby playing an important role in tumor initiation (7). In response to DNA damage, cells activate complex signaling networks to either repair the damage or promote cell death (8). Oxidative DNA damage has been implicated in the pathogenesis of several diseases including cancer, inflammation, and heart disease (9). Thus, it is important for organisms to suppress the excessive generation of ROS.

ROS-mediated DNA damage influences cell survival through the modification of histone H2AX (10-14). The tumor suppressor p53 regulates baseline and DNA damage-inducible levels of ROS (15-17). p53 prevents DNA damage through an activated phosphatidylinositol 3-kinase-related kinase (PI3K) that plays a pivotal role in activating DNA repair proteins (18). Phosphorylated H2AX (γ-H2AX) serves an important role in the DNA damage response to oxidative stress by recruiting genes involved in DNA repair (19). Biological processes that eliminate ROS are important in the DNA damage response and in preventing the progression of numerous diseases (20).

Abeliophyllum distichum Nakai is a deciduous flowering plant belonging to the Oleaceae family. A. distichum is a monotypic genus and has been regarded as one of the important plant resources for research on its genetic property and bioactivity (21). Additionally, it is known to contain some glycosides including acteoside, isoacteoside, rutin and hirsutrin (22). Acteoside exhibits several pharmacological properties such as anti-hepatotoxic (23), anti-inflammatory (24), antinociceptive (25) and antioxidant activities (26-28). It also improves sexual function (29). However, to the best of our knowledge, whether acteoside from A. distichum (AAD) can prevent oxidative DNA damage remains unknown. Therefore, the aim of the present study was to determine whether AAD could exert a protective effect over oxidative DNA damage.

Materials and methods

Plant material and extraction. A. distichum Nakai [voucher no. Park1001(ANH)] was collected from the Misun-Hyang Theme Park. An extract from air-dried A. distichum was
obtained by immersion in 80% methanol for 3 days and a using a sonicator (40 KHz; 700 W; Hwashin Technology Co., Ltd.). The methanol-soluble extracts were filtered and concentrated using an N-1110S vacuum evaporator (Eyela). The extracts were sequentially fractionated three times with 1:1 ratio of petroleum ether, 1:1 ratio of chloroform and 1:1 ratio of ethyl acetate. The ethyl acetate fraction from A. distichum was harvested and refrigerated at 4°C until use.

Isolation and purification. Acteoside was isolated from the ethyl acetate fraction (500 g) of A. distichum using an Isolera™ Spectra Accelerated Chromatographic Isolation system at 25°C with SNAP KP-SIL (100 g; 170×40 mm) and SNAP Ultra (25 g; 80 x 35 mm) cartridges (both from Biotage AB). The mobile phases consisted of chloroform (solvent A) and methanol (solvent B). The flow rate was kept at 50 ml/min for a total run volume of 1,600 ml.

The system was run with a gradient program: 0-32 min, 6-50% B and the peaks of interest were monitored at 335 nm by a photodiode array (PDA) detector with SNAP KP-SIL cartridges. Subsequently, using the same mobile phases, the flow rate was kept constant at 75 ml/min for a total run volume of 600 ml. Then the system was run with a gradient program: 0-32 min, 6-50% B and the peaks of interest were monitored at 335 nm by a PDA detector using SNAP Ultra cartridges. Lastly, 2.5 g AAD was further analyzed by high-performance liquid chromatography (HPLC)-PDA and liquid chromatography-mass spectrometry (LC-MS) for further analysis and identification.

Identification of acteoside by HPLC analysis. AAD (1 mg/ml) was analyzed in a Waters 2695 HPLC system equipped with Waters 2996 PDA using an Xbridge-C18 column (250x4.6 mm; 5 µm; all from Waters Corporation). The binary mobile phase consisted of acetonitrile (solvent A) and water containing 1% acetic acid (solvent B). Both solvents were filtered through a 0.45-µm filter prior to use. The flow rate was kept constant at 1.0 ml/min for a total run time of 40 min at 25°C. The system was run with a gradient program: i) 0-5 min, 10-15% A: 90-85% B; ii) 5-15 min, 15% A: 85% B; iii) 15-40 min, 15-40% A: 85-60% B. The sample injection volume was 10 µl. The peaks of interest were monitored at 335 nm by a PDA and compared with an acteoside standard (V4015-10MG; Sigma-Aldrich; Merck KGaA).

Identification of acteoside by LC-MS. LC-MS analysis was performed using a Nexera X2 Ultra high-performance liquid chromatography/mass selective detector at 25°C, equipped with an LCMS-8050 quadrupole mass spectrometer (all from Shimadzu Corporation). The liquid chromatographic system consisted of a binary pump, on-line vacuum degasser and thermostatic column compartment, connected in-line to the mass spectrometer. Data acquisition and analysis were carried out using Shimadzu LabSolutions 5.0 (Shimadzu Corporation). Samples (3 µl) were injected into the HPLC system using a binary gradient of deionized water (solvent A) and acetonitrile (solvent B) and isocratically eluted in 50% solvent B for 1 min at a flow rate of 0.3 ml/min. The mass spectrometer was fitted to an atmospheric pressure electrospray ionization source operated in negative ion mode. The electrospray capillary voltage was set to 4.0 kV, with a nebulizing gas (10.7 bar) flow rate of 3 l/min and a drying gas temperature of 300°C. MS data were acquired in the Scan mode (mass range m/z 300-800) and the Product ion scan mode.

1,1-Diphenyl-2-picryl hydrazyl (DPPH) radical scavenging activity. DPPH radical scavenging activity was measured as previously described (30), with minor modifications. A 300 µM DPPH solution (Sigma-Aldrich; Merck KGaA) in 95% alcohol was prepared. The solution was adjusted to an absorbance value of 1.00 at 515 nm. A 40-µl volume of sample was mixed with 760 µl DDPH solution, then incubated for 20 min in the dark at 25°C. L-ascorbic acid was used as a positive control. Absorbance was measured at 515 nm using an ultraviolet (UV)/visible spectrophotometer (Xma-3000PC; Human Corporation). The radical scavenging activity was measured as a decrease in the absorbance of DPPH and calculated using the following formula:

\[ \text{DPPH radical scavenging activity} \% = \frac{1-(A_{\text{Sample}} - A_{\text{Blank}})/A_{\text{control}}}{100} \times 100 \]

where 'A_{\text{Sample}}' = Absorbance values of DPPH radicals following treatment with sample, 'A_{\text{Blank}}' = Absorbance values of ethanol and 'A_{\text{control}}' = Absorbance values of DPPH radicals.

2,2’-Azino-bis (3-ethylbenzothiazoline-6-sulfonic acid) diaminonitrobenzene (ABTS) radical scavenging activity. ABTS radical scavenging activity was measured as previously described (31), with some modifications. A 7.4 mM ABTS and 2.6 mM potassium persulfate solution (both from Sigma-Aldrich; Merck KGaA) in distilled water was prepared 24 h prior to use. The solution was adjusted to an absorbance value of 0.70 at 732 nm. A 40 µl volume of sample was mixed with 760 µl ABTS solution, then incubated for 20 min in the dark at 25°C. L-ascorbic acid was used as a positive control. Absorbance was measured using a UV/visible spectrophotometer at 732 nm. The radical scavenging activity was measured as a decrease in the absorbance of ABTS and calculated using the following formula: ABTS radical scavenging activity (%) = \[1-(A_{\text{Sample}} - A_{\text{Blank}})/A_{\text{Control}}\] x100 where 'A_{\text{Sample}}' = Absorbance values of ABTS radicals after treatment with sample, 'A_{\text{Blank}}' = Absorbance values of H₂O and 'A_{\text{Control}}' = Absorbance values of ABTS radicals.

Inhibitory effect of AAD on oxidative DNA damage using ΦX-174 RF1 plasmid DNA. The inhibitory effect on oxidative DNA damage was measured as previously described (32). Conversion of supercoiled plasmid DNA to open circular forms has been used as an index of DNA damage (32). In the absence of OH radicals and Fe²⁺ ions, plasmid DNA mainly exists in a supercoiled form. However, in the presence of OH radicals and Fe²⁺, plasmid DNA is cleaved, converting its supercoiled form into open-circular forms. AAD (0-200 µM) was pre-incubated with 5 mM FeCl₂ or 0.4 mM FeSO₄ + 0.4 mM H₂O₂ for 15 min at 25°C. The ΦX-174 RF1 plasmid DNA (50 µg/ml; Promega Corporation) was then added to the mixtures. A solution containing 50% glycerol (v/v), 40 mM EDTA, and 0.05% bromphenol blue (Sigma-Aldrich; Merck KGaA) was added to stop the reaction. Reaction mixtures were analyzed by agarose gel electrophoresis on 1% gels with DNA SafeStain (Lamda Biotech). DNA in the gel was...
visualized using the Chemi-Doc imaging system (Bio-Rad Laboratories, Inc.) controlled by Image Lab software 6.0 (Bio-Rad Laboratories, Inc.).

**Cell culture.** The murine skin fibroblast cell line NIH 3T3 was purchased from American Type Culture Collection. Cells were cultured under sterile conditions at 37°C in a humid incubator with 5% of CO2, in Dulbecco's modified Eagle's medium supplemented with 10% bovine calf serum (Biowest), 1% antibiotics (penicillin/streptomycin; Hyclone; Cytiva) and 0.2% anti-mycoplasma agents (Plasmocin prophylactic; InvivoGen). The oxidative damage was induced by 150 µM FeSO4 + 600 µM H2O2 for the cell viability, western blotting, and PCR assays.

**Cell viability.** Cells (1.0x10⁴ cells/ml) were treated with aad (0‑200 µg/ml) and incubated at 37°C for 24 h. After 24 h, cells were treated with 10 µl alamarBlue ® cell viability reagent (Thermo Fisher Scientific, Inc.). After 2 h, cell viability was measured using a UV/Visible spectrophotometer at 570 nm.

**Immunoblotting.** Cells were lysed using radioimmunoprecipitation assay buffer (Thermo Fisher Scientific, Inc.) supplemented with a protease inhibitor cocktail (Sigma-Aldrich; Merck KGaA). The protein concentration in the sample was determined using the Bradford protein assay (Bio-rad laboratories, inc.). Proteins (20 µg) were separated by SDS-PAGE on 10% gels, then transferred to a PVDF (Bio-Rad Laboratories, Inc.). The membrane was blocked for 1 h at 20°C with 5% bovine serum albumin (BSA; Biosesang) in TBS containing 1% Tween-20 (TBS-T), then incubated with primary antibodies in 3% BSA overnight at 4°C. The following primary antibodies were used at 1:1,000 (all from Abcam): i) Anti-γH2AX (Ser139) polyclonal antibody (cat. no. ab11174); ii) anti-H2AX polyclonal antibody (cat. no. ab11175); iii) anti-phosphorylated-p53 (p-p53; Ser15) polyclonal antibody (cat. no. ab1431); iv) anti-p53 polyclonal antibody (cat. no. ab131442); and v) anti-GAPDH monoclonal antibody (cat. no. ab8245). After washing with TBS-T, membranes were incubated with anti-rabbit IgG monoclonal antibody (cat. no. 18-8816-33; Rockland Immunochemicals Inc.; 1:5,000) or anti-mouse IgG monoclonal antibody (cat. no. 18-8817-33; Rockland Immunochemicals Inc.; 1:5,000) horseradish peroxidase-conjugated secondary antibodies for 1 h. Immunoreactive bands were visualized using an ECL western blotting substrate and the Chemi-Doc imaging system (both from Bio-Rad Laboratories, Inc.). Bands were analyzed using the ImageJ software v1.51K (National Institutes of Health).

**Reverse transcription (RT)-semi-quantitative PCR (qPCR).** Total RNA was extracted from NIH 3T3 cells using the NucleoSpin™ RNA Plus kit (Macherey-Nagel; Thermo Fisher Scientific, Inc.). cDNA was synthesized using ReverTra Ace-α™ (Toyobo Life Science), according to the manufacturer's protocol. qPCR was carried out using Quick Taq® HS Dye Mix (Toyobo Life Science). Amplification was initiated at 94°C for 2 min, followed by 30 amplification cycles at 94°C for 30 sec (denaturation), then 54.2°C for 30 sec (annealing/extension) using Thermal cycler (T100™; Bio-rad laboratories, inc.). Primer sequences using Primer 3 program 0.4.0 (http://bioinfo.ut.ee/primer3-0.4.0/) are presented in Table I. DNA density was visualized by 2% agarose gel (A1200; Biosesang) with 1% DNA SafeStain (C138; Lambda Biotech Corporation). Transcription levels were normalized to GAPDH. Density was analyzed using the ImageJ software 1.51K (National Institutes of Health).

**RT-qPCR analysis.** cDNA samples were generated using the aforementioned method. qPCR was carried out using the QuantiTect® SYBR-Green PCR kit (Qiagen GmbH) on a 7500 Real-Time PCR system (Applied Biosystems; Thermo Fisher Scientific, Inc.). Primers were designed using the Primer 3 program 0.4.0 (http://bioinfo.ut.ee/primer3-0.4.0/; Table I). Thermocycling conditions consisted of a 15-min initial denaturation at 95°C, followed by 40 amplification cycles at 94°C for 15 sec (denaturation), 60°C for 30 sec (annealing) and 72°C for 30 sec (extension).
Transcription levels were normalized to GAPDH. The $2^{\Delta\Delta Cq}$ formula was used to analyze mRNA expression levels, where $\Delta\Delta Cq = (Cq_{\text{target}} - Cq_{\text{GAPDH}})_{\text{sample}} - (Cq_{\text{target}} - Cq_{\text{GAPDH}})_{\text{control}}$ (33). Data were analyzed using the ABI 7500 Software 2.3 (Applied Biosystems; Thermo Fisher Scientific, Inc.).

Immunofluorescence staining. γ-H2AX and p-p53 are markers for DNA damage and repair (34). NIH 3T3 cells were treated with AAD (0, 100 µM) and oxidative damage (150 µM FeSO₄ + 600 µM H₂O₂) at 37°C for 24 h. Following treatment, samples were prepared on sterilized glass coverslips (BD Biosciences). Cells were then fixed with 4% paraformaldehyde for 15 min at 20°C, then blocked for 1 h with 3% BSA (Biosesang) and 0.1% Triton X-100 for permeabilization at 20°C. Cells were incubated with anti-γ-H2AX (cat. no. ab11174; 1:500) and anti-p-p53 (cat. no. ab1431; 1:500) overnight at 4°C, then with Alexa Fluor 488 (cat. no. ab150077; 1:500) and Alexa Fluor 568-conjugated secondary antibodies (cat. no. ab175471; 1:500; all antibodies from Abcam) for an additional 1 h at 20°C in the dark. Nuclei were stained using DAPI (Invitrogen; Thermo Fisher Scientific, Inc.) at 37°C. Slides were mounted and images captured using a CKX53 fluorescence microscope (magnification, x400; Olympus Corporation).

Statistical analysis. All data were analyzed using GraphPad Prism 5.0 (GraphPad Software, Inc.) and are presented as the mean ± standard deviation. All experiments were performed ≥3 times. Data of DPPH and ABTS radical scavenging activity were analyzed using paired Student's t-test. Other data were analyzed by One-way ANOVA followed by Tukey's post hoc test were used to compare the means across multiple groups. P<0.05 was considered to indicate a statistically significant difference.

Figure 1. HPLC chromatograms and structure of AAD. (A) HPLC chromatogram and UV-spectra of AAD. (B) HPLC chromatogram and UV-spectra of standard acteoside. (C) LC-MS spectra of AAD with m/z 623. (D) Structure of acteoside. AAD, acteoside from Abeliophyllum distichum; AU, arbitrary units; HPLC, high-performance liquid chromatography; UV, ultraviolet.

Results

Isolation, purification and identification of AAD. Acteoside standard and AAD extract were analyzed using HPLC-PDA. The standard (Fig. 1A) and the sample (Fig. 1B) displayed similar retention times (20.3 min) and UV spectra. The LC-MS data (Fig. 1C) also exhibited 623 m/z (M-H⁻). Based on these observations, the AAD sample was identified as an acteoside (Fig. 1D). The purity of AAD was >95%, based on a calibration curve ($y = 16089x - 93989, R^2 = 0.9971$).

Antioxidant activity of AAD. AAD eliminated DPPH radicals in a dose-dependent manner (Fig. 2A). DPPH radicals were scavenged at all AAD concentrations, except 0.32 µM, although not to the same extent as with L-ascorbic acid. The half-maximal inhibitory concentration (IC₅₀) values of AAD and L-ascorbic acid were 8.81 and 5.08 µg/ml, respectively. Similarly, AAD also eliminated ABTS radicals in a dose-dependent manner (Fig. 2B). The IC₅₀ values of AAD and L-ascorbic acid were 6.47 and 10.49 µg/ml, respectively. Furthermore, at concentrations of 8 and 40 µM, the scavenging activity of AAD was significantly higher compared with L-ascorbic acid. These results indicated that AAD effectively scavenged DPPH and ABTS radicals in vitro.

Protective effect against oxidative stress-induced DNA damage of AAD. The protective effects of AAD on oxidative DNA damage were evaluated in a DNA cleavage assay using ΦX-174 RF I DNA. The untreated control group (lane 1) was used as a positive control and assigned an open circular (OC) DNA band density value of 100%. In lane 2, the groups receiving Fe²⁺ or OH⁻ without AAD treatment were used as a negative control and given a value of 0%.
aad inhibited Fe\textsuperscript{2+}-induced oxidative DNA damage in a dose-dependent manner, as suggested by the significant increase in band density at all concentrations, compared with the negative control (47.14±1.64% at 0.32 µM aad vs. 97.66±3.41% at 200 µM; Fig. 3a). Additionally, aad also significantly protected plasmid DNA from OH\textsuperscript{·} radicals-induced oxidative damage in a dose-dependent manner (7.09±0.25% at 0.32 µM aad vs. 88.05±3.09% at 200 µM; Fig. 3B). These results suggested that aad could protect plasmid DNA from oxidative damage.

AAD protects NIH 3T3 cells against oxidative DNA damage. To determine whether AAD had any cytotoxic effect, NIH 3T3 cells were cultured in AAD concentrations ranging between 25 and 200 µM, and cell viability was then assessed following 24-h incubation. AAD had no effect on cell viability. In oxidative damage conditions, AAD increased cell viability compared with untreated cells (Fig. 4).

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Cytotoxicity of AAD in NIH 3T3 cells. To determine whether AAD had any cytotoxic effect, NIH 3T3 cells were cultured in AAD concentrations ranging between 25 and 200 µM, and cell viability was then assessed following 24-h incubation. AAD had no effect on cell viability. In oxidative damage conditions, AAD increased cell viability compared with untreated cells (Fig. 4).
of AAD, oxidative damage significantly increased phosphorylation of p-53 and H2AX in NIH 3T3 cells. However, under oxidative damage conditions, the levels of p-p53 and γ-H2AX were significantly reduced by treatment with AAD. This reduction in p-p53 and γ-H2AX was dose-dependent (Fig. 5A and B). Additionally, gene expression levels of p53 and H2AX followed the same pattern, as indicated by RT-semi-qPCR (Fig. 5C and D) and RT-qPCR (Fig. 5E).

Immunofluorescence staining of p-p53 and γ-H2AX was also performed (Fig. 6). The levels of p-p53 and γ-H2AX increased following oxidative damage. However, p-p53 and γ-H2AX levels were reduced in the presence of AAD compared with untreated oxidative damaged cells. These results suggested that AAD could regulate the expression of p-p53 and γ-H2AX.

**Discussion**

Due to the rarity of *A. distichum* Nakai, previous studies have focused on its ecological characterization, genomic sequencing and biological properties (35-40). Bremer et al (41) demonstrated that a number of plants in the Oleaceae family, such as the *Abeliophyllum*, *Forsythia* and *Jasminum* genera, contain acteoside. Although the potential bioactivities of acteoside may have important uses in industry and research, AAD remains poorly characterized.
Several plant-sourced compounds, such as vitamin C, phenolics and flavonoids, can act as natural antioxidants by reacting with free radicals, chelating catalytic metals and scavenging oxygen (42,43). Antioxidants suppress oxygen, electrons and hydrogen atoms that are generated during intracellular metabolism. They also protect cells against harmful effects of ROS and associated oxidative stress generated in aerobic organisms (44). For these reasons, numerous studies on the regulation of ROS and the role of natural antioxidants derived from plants have been conducted (45-47).

In the present study, HPLC and LC-MS analyses were carried out to characterize an AAD extract. The AAD sample had the same retention time and a similar UV spectrum compared with a standard acteoside. Scale bar, 40 µm. (A) Fluorescence microscopy shows expression of anti-p-p53 antibody in NIH 3T3 cells treated for 24 h with 150 µM FeSO4 and 600 µM H2O2 (oxidative damage) in the presence or absence of 100 µM AAD. Green, p-p53; Blue, DAPI. (B) Fluorescence microscopy shows expression of anti-γ-H2AX antibody in NIH 3T3 cells treated for 24 h with 150 µM FeSO4 and 600 µM H2O2 (oxidative damage) in the presence or absence of 100 µM AAD. Green, γ-H2AX, Blue, DAPI. Scale bar, 40 µm. Control (0 µM AAD; no oxidative damage); AAD, acteoside from Abeliophyllum distichum; H2AX, H2A histone family member X; p/γ, phosphorylated; DAPI, 4,6-diamidino-2-phenylindole.

Figure 6. AAD regulates p-p53 and γ-H2AX in NIH 3T3 cells. (A) Fluorescence microscopy shows expression of anti-p-p53 antibody in NIH 3T3 cells treated for 24 h with 150 µM FeSO4 and 600 µM H2O2 (oxidative damage) in the presence or absence of 100 µM AAD. Green, p-p53; Blue, DAPI. Scale bar, 40 µm. (B) Fluorescence microscopy shows expression of anti-γ-H2AX antibody in NIH 3T3 cells treated for 24 h with 150 µM FeSO4 and 600 µM H2O2 (oxidative damage) in the presence or absence of 100 µM AAD. Green, γ-H2AX, Blue, DAPI. Scale bar, 40 µm. Control (0 µM AAD; no oxidative damage); AAD, acteoside from Abeliophyllum distichum; H2AX, H2A histone family member X; p/γ, phosphorylated; DAPI, 4,6-diamidino-2-phenylindole.

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In the present study, HPLC and LC-MS analyses were carried out to characterize an AAD extract. The AAD sample had the same retention time and a similar UV spectrum compared with a standard acteoside. Additionally, based on LC-MS analysis, the AAD peak was observed at 623 m/z (M-H), which coincided with the molecular weight of acteoside (48). Consequently, based on HPLC-PDA and LC-MS data, the AAD extract used in this study was identified as acteoside. Additionally, the effects of the antioxidant properties of AAD on oxidative DNA damage were also examined. Our previous studies suggested that A. distichum had potential antioxidant activities (49,50). DPPH and ABTS radicals are the most common and most stable chromogens used to estimate antioxidant activity of biological materials. Furthermore, DPPH radical scavenging and ABTS radical cation decolorization assays can both be used to evaluate antioxidant activities in a relatively short time (51,52). Oxidation of ABTS with potassium persulfate generates ABTS+ radical cations with the transfer of one electron. Under prolonged oxidative conditions, these radicals can generate di-cation ABTS+ radicals (53,54). In the present study, AAD scavenged both DPPH and ABTS radicals.

DNA damage leads to an irreversible covalent modification of the DNA molecule. This can occur through single- or double-strand breaks (DSBs) in the DNA sugar-phosphate backbone, disruption of the or N-glycosyl bonds linking nucleobases to the sugar, as well as chemical modification of nucleobase residues (55,56). Oxidative DNA damage represents one of the most frequent consequences of exposure to exogenous environmental stimuli or endogenous genotoxic agents. These reactions are associated with the activity of ROS, such as hydroxyl radicals, peroxides, peroxides or single oxygen (57,58). For instance, the oxidative DNA damage produced by the Fenton reaction plays a major role in the aging process, as well as several diseases, including Alzheimer’s disease, cancer and multiple sclerosis (59-62). In the present study, the protective effects of AAD against oxidative DNA damage were evaluated using non-cellular and cellular models. A ΦX-174 RF I plasmid DNA cleavage assay was conducted using OH radicals and Fe2+ ions. Hydrogen peroxide itself cannot directly oxidize DNA molecules, but can react with transition metals, such as Fe2+ to form OH radicals. These OH radicals attack DNA structures, leading to sugar fragmentation, strand scission and base adducts (63). The Fe2+ ion is an essential transition metal element in humans. In the present study, AAD decreased oxidative DNA damage in a dose-dependent manner. Therefore, due to its scavenging capacity, AAD may prevent cell damage caused by OH- radicals and Fe2+ ions. This type of DNA damage is one of the most genotoxic types occurring in cells, either resulting in cell cycle arrest and DNA repair, or elimination of the injured cells. Appropriate cellular responses to DNA DSBs are important for tumor suppression and maintaining genetic stability (64,65).

One of the first cellular responses to the introduction of DSBs is the phosphorylation of H2AX, a sensitive marker for DNA DSBs. DSBs induce phosphorylation of H2AX at its C-terminal serine residues (Ser136 and Ser139) (66). The phosphorylated formed of H2AX, called γ-H2AX, is detectable at the sites of the damage following the introduction of DSBs (67,68). Therefore, γ-H2AX plays a distinct role in the DNA damage response. Additionally, the p53 pathway is also a key effector of the DNA damage response and is activated by several stimuli that induce DNA lesions (69). The expression of phosphorylated p53 (p-p53) may represent a negative regulator of the cellular response to DNA damage (70). In the present study, AAD treatment reduced the phosphorylation of γ-H2AX and p-p53 in NIH 3T3 cells under oxidative damage conditions. Moreover, the expression levels of H2AX and p53 were also reduced by AAD treatment. Although several other possible mechanisms may also explain these observations, the present study identified a relationship between the antioxidant properties of AAD and inhibition of DNA damage. In conclusion, the present study demonstrated that AAD displayed antioxidant properties that could protect DNA against oxidative damage.
Acknowledgements

Not applicable.

Funding

This research was supported by The Basic Science Research Program through The National Research Foundation of Korea funded by The Ministry of Education (grant no. NRF-2016R1D1A1B03934869).

Availability of data and materials

All data generated or analyzed during this study are included in this published article.

Authors' contributions

TWJ and JHP drafted the manuscript and performed the experiments. TWJ and JSC performed the experiments and data interpretation. TWJ contributed materials and analytical tools and helped in data analysis. TWJ, JSC and JHP designed the experiments, provided critical suggestions for the manuscript, and reviewed and revised the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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