RESEARCH ARTICLE

Anti-Genotoxicity Evaluation of *Cratoxylum Formosum* Dyer Leaves by Comet Assay and Micronucleus Test

Ratsada Prphasawat*, Narongsak Munkong

Abstract

*Cratoxylum formosum* Dyer is the Thai vegetable which commonly consumed a fresh leaves. In this study, we extracted *Cratoxylum formosum* with water and tested the extract for genotoxicity and anti-genotoxicity effects. We carried out the experiment using micronucleus test and comet assay in TK6 cells. In micronucleus experiment, we used cytokinesis-block proliferation technique to stop cell division which produced a cell at binucleated (BNC) stage. The comet assay was carried out after pre-treatment the cell with *C. formosum* for 18 h. The results revealed not increased the micronucleus frequency of *C. formosum* at concentration ranging from 50-150 µg/ml. In contract, it showed that the combination between *C. formosum* at various concentrations (25, 50, 75, 100, 150 and 200 µg/ml) and mitomycin C could decrease significantly in frequency of micronuclei. The mean of micronucleus frequency in the sample were 23.17 ±3.33, 23.33 ±4.72, 21.00 ±3.61, 11.33 ±3.21, 16.67 ±2.08, and 23.33±1.53 MN/ 1,000 BNC, respectively whereas the MMC-treated group was 33.67 ± 8.96 MN/ 1,000 BNC. The comet assay result showed that pre-treatment with *Cratoxylum formosum* (25, 50, 100, 200 µg/ml) could inhibit the hydrogen peroxide induced DNA damage by 6.95, 12.99, 17.61, and 26.39 respectively.

Keywords: *Cratoxylum formosum* Dyer- comet assay- micronucleus test

Introduction

*Cratoxylum formosum* Dyer (CF) was a plant mostly found in many Southeast Asia countries including Thailand. Many people used this plant as food. They are commonly consumed the fresh young leaf or sometimes used a leaf as an ingredient in soup. There are studies found that its leaf have many biological activities such as anti-oxidant and anti-mutagenic properties. Moreover, the aqueous extracts from leaf possess strong free radical scavenging and protective effect to vascular also (Yingngam et al., 2014; Nakahara et al., 2002; Maisuthisakul et al., 2007). However, there are little data on DNA effect of its leaf.

According to OECD guideline, a well know genotoxicity assays which is the comet assay and micronucleus (MN) test have been approved for detect DNA damage (Pant et al., 2015; Araldi et al., 2015). The alkaline comet assay or single-cell gel electrophoresis (SCGE) is a sensitive and specific technique for measuring DNA damage. The damage expressed as comet after the electric current pulled the charged DNA from the nucleus. The comet cell composed of the head of normal DNA and the tail of DNA break. Thereby, the intensity of the DNA in tail is correlated to DNA damage (Collins, 2004; Tice et al., 2000). In the MN test is assay for detect the DNA damage result from clastogenic and aneugenic activities which lead to micronuclei formation in the cytoplasm during mitosis. Thus, an increase in the micronuclei frequency is an indication of chromosomal damage (OECD, 2010; Fenech, 2000). Therefore, the combination of two different assays has been mostly used to assess of DNA damage in many cells and tissue.

The aim of the present study is to evaluate the potential in vitro anti-genotoxic effect of *C. formosum* leaves using the comet assay and MN test. These tests using TK6 lymphoblastoid cell line were performed.

Materials and Methods

Extraction of *C. formosum* leaves

Fresh leaves of CF were harvested and collected from Phayao Province, North of Thailand. The leaves were cleaned by tap water and air-dried. Two hundred-fifty grams of fresh leaves were blended with 2.5 L distilled water and heated for 1 h at 80°C in a water bath. Following, filtration through Whatman no.1 filter paper using a suction apparatus, the extract was lyophilized giving a reddish brown color. The dry extracted was weight and kept at -20°C. The yield of freeze-dried powder from fresh leaves was about 8%W/V.

Cell culture and maintaining

The TK6 human lymphoblastoid cell line (CRL-8015) were purchased from American Type Culture Collection Department of Pathology, School of Medicine, University of Phayao, Phayao, Thailand. *For correspondence: ratsada.pr@gmail.com*
Cytotoxicity test

The cytotoxicity test was done in our laboratory. CF extract was investigated in TK6 cells by WST-1 assay (Ngamwongsatit et al., 2008). In the experiment, TK6 cells were cultured in the presence of CF leaf extracts at various concentrations for 24 h and the percentage of the cell viability was evaluated by WST-1 method. Then, cell viability values were calculated as inhibitory concentration (IC50). The leaf extract of CF showed the low cytotoxicity expressed as IC50 values of 381.48±36.13 µg/ml as reported previous.

Micronucleus test

Prior to the anti-genotoxicity assay, the genotoxicity of CF leaves were evaluated at selected doses based on viability greater than 70%. In this study the extract at doses of 50, 100 and 150 µg/ml were incubated in TK6 cells for 24 h at 37°C. The anti-genotoxicity experiment was performed at doses (>70% cell viability) of 25, 50, 75, 100, 150 and 200 µg/ml RPMI in combination with a known mutagen, mitomycin C (MMC) at 0.8 µg/ml) for 24 h (Elhajouji, 2010; Aardema et al., 2011; Fenech, 2007). After treatment, a cytochalasin B solution (Cyt B 3 µg/ml) was added during 6 h to collect the cells at a binucleated stage 13. Following washing and harvesting steps, treated cells were prepared as monolayer on glass slides using cytoospin equipment (Shandon, UK). Slides were left to dry at room temperature and then fixed in cold methanol for 30 mins. Cells on slides were stained using 10% Giemsa solution. Micronuclei (MN) formations were scored in 1000 binucleated (BNC) cells under light microscope (40x)(Figure 1).

Comet assay

In the present study, alkaline condition (pH>13) was used in the lysing and electrophoresis steps which produce DNA denature. After seeding the TK6 cells (2×10⁴ cells/ml) into a 6-well plate, cells were treated for 18 h with CF leaf extract at doses of 25, 50, 100, 200 µg/ml RPMI. By the end of the treatment time, cells were harvested by to remove extract-containing medium. Cells were then treated with 50 µM H2O2 for 5 mins at 4°C for induced DNA damage. Following treatment, cells were washed twice with cold HBSS and re-suspended in RPMI medium to be ready for comet assay.

The comet assay was performed following the method described by Tice (2003). In brief, 20 µl of cell suspension was mixed with 75 µl of 0.5% low melting point (LMP) agarose at 37°C, layered onto a pre-coated slide with 0.75% normal melting point (NMP) agarose and covered with a coverslip allowing gel-solidification on a flat surface ice box. The coverslip was gently removed and 95 µl of 0.75% normal melting point (NMP) agarose and covered with a coverslip allowing gel-solidification on a flat surface ice box. The coverslip was gently removed and 95 µl of 0.75% normal melting point (NMP) agarose and covered with a coverslip allowing gel-solidification on a flat surface ice box. After coverslip removal, the slides immersed into a lysis ice box. The coverslip was gently removed and 95 µl of 0.75% normal melting point (NMP) agarose and covered with a coverslip allowing gel-solidification on a flat surface ice box. After coverslip removal, the slides immersed into a lysis

solution (2.5M NaCl, 10 mM Na2EDTA-2H2O, 10 mM Trisma base, pH 10 with 1% triton X-100, 10% DMSO) for 2 h at 4°C. After lysis, slides were exposed to freshly make alkaline electrophoresis buffer (200 mM EDTA, 10 N NaOH, pH 13) for 20 min to allow DNA unwinding. The slides were then placed on an electrophoresis tank filled with sufficient electrophoresis solution and kept in an ice bath (4°C). Electrophoresis was carried out for 20 mins at constant 25V and a current of 300mA using a power pac supply. Then, the slides were neutralized in 0.4 M Trizma base buffer (pH 7.5) and stained with 100 µl of 20 µg/ml Ethidium bromide. At least 50 cells per slide and per treatment were randomly analyzed for comet images using the fluorescence microscope (at 40x magnification) connected to a computer equipped with an automated image analysis system (Comet assay III, Perceptive Instrument, UK).

Results of comet assay was used two parameters as indicator of DNA damage like tail length (TL= a distance of damage-DNA migration) and tail moment (TM; a DNA damage intensity) values.

Statistical analysis

The experiment was repeated in triplicate for MN test and duplicate for the comet assay. Data was expressed as mean ±SD of the medians were calculated for each group of treatments and then analyzed by SPSS software program. The statistical approach included the one-way ANOVA followed by LSD test which was to evaluate the significance of the differences in micronucleus frequency and comet assay data.

Results

Micronucleus test

In this study, we performed the micronucleus test to investigate the mutagenic and anti-mutagenic properties of the CF leaf extracts at various concentrations. Maximum concentrations were defined as those inducing around 40-50% toxicity which was 200 µg/ml of CF extract. Result of MN test showed that negative control (untreated cells) exhibited a baseline frequency of micronuclei of 1.33 MN/1,000 BNC. After exposure, CF leaf extracts did not induce the MN frequency in TK6 cells. The mean micronucleus frequencies in the CF treatment were 3.00, 2.50, and 2.67 MN/ 1,000 BNC for doses of 50, 100, and 150 µg/ml, whereas the induction of micronucleus found in MMC-treated cells were 5 MN/ 1,000 BNC.

For the anti-genotoxicity assay, the combination of CF leaf extract and MMC was performed in the MN test using TK6 cells. The mean frequency of micronucleus cells in negative control group was 2.33 ±3.21 MN/1,000 BNC. In contrast, MMC-treated group induced a significant increase in MN frequency from negative control group by 15 folds (66.67 ±2.08, 23.33±1.53 MN/ 1000 BNC, respectively.

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Both assays are difference in endpoints of DNA damage. The MN test indicates DNA damage as clastogenic or aneugenic activities, whereas the comet assay for detecting DNA strand, cross-links and alkaline-labile sites which utilized a wide variety of cells and tissues (Pilar et al., 2016; Zapata et al., 2016). However, in comet assay was relatively sensitivity than the MN test because the comet assays can detection of carcinogens which gave negative results in the MN test. Therefore, we suggested the used of both techniques for assessing whether the chemical can induced DNA damage (Pant et al., 2015; Araldi et al., 2015).

In consideration of the result of this study, the protective effect might be the phenolic compound found in C. formosum leaf due to anti-oxidant activity. These would lead to being oxidized the H2O2 to attack cellular component (Kaur and Kapoor, 2001). Similar results were reported that the extraction for phenolic compounds from C. formosum has protective ability against H2O2 –induced HEK293 cells death (Yingngam et al., 2014).

Our results reveal the absence of genotoxicity of C. formosum leaf extract in TK6 cells. For the anti-genotoxicity test, C. formosum leaf extract possess a moderate inhibition of DNA damage both in vitro comet assay and micronucleus test. Moreover, we suggest the in vivo data are required before its approval.

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