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Fruit extract of the medicinal plant *Crataegus oxyacantha* exerts genotoxic and mutagenic effects in cultured cells

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**ABSTRACT**

*Crataegus oxyacantha*, a plant of the Rosaceae family also known “English hawthorn, haw, maybush, or whitethorn,” has long been used for medicinal purposes such as digestive disorders, hyperlipidemia, dyspnea, inducing diuresis, and preventing kidney stones. However, the predominant use of this plant has been to treat cardiovascular disorders. Due to a lack of studies on the genotoxicity of *C. oxyacantha*, this investigation was undertaken to determine whether its fruit extract exerts cytotoxic, genotoxic, or clastogenic/aneugenic effects in leukocytes and HepG2 (liver hepatocellular carcinoma) cultured human cells, or mutagenic effects in TA100 and TA98 strains of *Salmonella typhimurium* bacterium. Genotoxicity analysis showed that the extract produced no marked genotoxic effects at concentrations of 2.5 or 5 µg/ml in either cell type; however, at concentrations of 10 µg/ml or higher significant DNA damage was detected. The micronucleus test also demonstrated that concentrations of 10 µg/ml or higher produced clastogenic/aneugenic responses. In the Ames test, the extract induced mutagenic effects in TA98 strain of *S. typhimurium* with metabolic activation at all tested concentrations (2.5 to 500 µg/ml). Data indicate that, under certain experimental conditions, the fruit extract of *C. oxyacantha* exerts genotoxic and clastogenic/aneugenic effects in cultured human cells, and with metabolism mutagenicity occurs in bacteria cells.

**ARTICLE HISTORY**

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**Introduction**

*Crataegus oxyacantha* L., a plant of the Rosaceae family that reaches a length of 25 to 30 m, was originally found in Europe, North America, and West Asia, but due to medicinal potential, this plant has since been cultivated in other locations, such as South America. In springtime, *C. oxyacantha* L. forms large white or pink inflorescence with a distinct aroma; while in the fall, these floral clusters become small shiny red fruits (Weihmayr & Ernst, 1996). Extracts of this shrub’s leaves and flowers and more recently its fruit have been used for several health conditions, especially heart problems (Rigelsky & Sweet, 2002; Zhang et al., 2001; Nascimento, 2009; Belščakcvitanović et al., 2014; Alp et al., 2015). This fruit has a long history of use as a diuretic, to treat dyspnea and renal calculus as well as sedative and anxiolytic effects (Rigelsky & Sweet, 2002). However, most pharmacological studies on this species focused on cardiotonic properties of a mixture of its aerial parts. The protective effect of *C. oxyacantha* against adverse cardiovascular activity is attributed to flavonoid constituents, particularly oligomeric protoanthocyanidins (OP), which exert significant antioxidant activity (Wang et al., 2013; Chang et al., 2005; Jayalakshmi & Niranjali-Devaraj, 2004).

The oligomeric protoanthocyanidins (OP) are largely concentrated in the leaves, fruits, flowers, and fruits. Oligomeric protoanthocyanidins are responsible for pigmentation of the fruits. Jayalakshmi and Niranjali-Devaraj (2004)
demonstrated that the extract of this plant reduced oxidative stress in the myocardium after reperfusion and appeared to inhibit apoptosis, resulting in a cardioprotective effect. The standard dose of *Crataegus* aerial parts extract is 300 to 600 mg three times daily of an extract standardized to contain 2 to 3% flavonoids or 18 to 20% procyanidins (Ammon & Hänel, 1981).

Studies on fruit extracts of other *Crataegus* species (*Crataegus microphylla*) showed protection of cultured human lymphocytes against the genotoxic effects induced by radiation (Hosseinimehr et al., 2007; 2009; 2011). The *Crataegus* genus is considered one of the oldest pharmacological sources and widely used in folk medicine (Bahorun et al., 2003).

At present, few studies examined the genetic toxicity of *C. oxyacantha* extract. Tabach et al. (2009) developed a preclinical toxicological assessment of a phytotherapeutic product—CPV, which consist of dry aerial parts of *C. oxyacantha* (26.7%), *Passiflora incarnata* (33.3%), and *Valeriana officinalis* (40%). The parameters assessed included weight, behavior, estrous cycle, teratogenicity in dogs, rats, or mice, and mutagenicity in bacteria (Ames test). Tabach et al. (2009) reported only negative results, indicating that the CPV mixture was apparently devoid of risk for humans.

Considering the widespread popular use of extract from aerial parts of *C. oxyacantha*, and lack of studies specifically assessing the genetic toxic potential of fruit extract, this investigation was conducted to determine the cytotoxic, genotoxic, and clastogenic/aneugenic potential in vitro, using human leukocytes and HepG2 cells, as well as mutagenic potential in bacteria cells. Among the available safety tests, the micronucleus (MN) test, comet assay, and Ames test are recommended by international regulatory agencies for toxicological evaluation of natural products (Choy, 2001; Claxton et al., 2010; OECD 489, 2014; Franco et al., 2015; Trindade et al., 2016). Thus, these tests were utilized to determine the safety of *C. oxyacantha*.

**Materials and methods**

**Botanical material**

*C. oxyacantha* fruit was collected in Turkey and purchased from certified distributor in Brazil. A sample voucher (LFF00297) was deposited in the Phytochemical Laboratory of José do Rosário Vellano University (Alfenas, MG). Dried and powdered *C. oxyacantha* fruit, in a ratio of herbal drug to drug preparation of 4:7:1, was extracted at room temperature with methanol 70% v/v. The macerated fruit was filtered and concentrated under reduced pressure in a rotary evaporator, resulting in crude methanolic extract (MeOH).

**Chemicals**

*C. oxyacantha* extract was diluted in Dulbecco’s modified Eagle’s medium (DMEM) (Gibco) to be used with HepG2 cells and RPMI-1640 medium (Gibco) to be employed with human lymphocytes. Methyl methane sulfonate (MMS) (Aldrich, CAS number 66-27-3) and benzo(a)pyrene (Sigma-Aldrich) were utilized as the positive control due to known potential for DNA damage, recognizable in the comet and MN assays. The other main chemicals were obtained from the following suppliers: trypsin (Sigma-Aldrich), phytohemagglutinin (Sigma-Aldrich), cytocalasin-B (Sigma-Aldrich), normal melting point (NMP) agarose (Invitrogen), low-melting point (LMP) agarose (Invitrogen), fetal calf serum (FCS) (Gibco), ethylenediaminetetraacetic acid (EDTA) (Merck), Triton X-100 (J.T.Baker), trypan blue (Sigma-Aldrich), thiazolyl blue tetrazolium bromide ethidium bromide (Sigma), dimethyl sulfoxide (DMSO) (Sigma-Aldrich), Giemsa (Synth), and Histopaque-1077 (Sigma-Aldrich).

**Phytochemical analysis of extract**

The 70% methanolic extract was analyzed using liquid chromatography attached to a mass detector. The chromatographic conditions were as follows: C18 column (100 x 2.1 mm, 1.7 mc), temperature 30 °C, injection volume 2 µl, using gradient mode for the mobile phase with mixture I (water: THF: formic acid 890:90:20) and mixture II (methanol:acetonitrile:formic acid 400:400:200) over a 0.2-ml/min flow. A triple quadrupole detector operating with an electrospray ionization source in the negative mode (ESI -), with desolvation and source temperatures of 300 °C and 120 °C, respectively, over a m/z 100 to m/z 1000 range.
The solutions were directly infused into the mass spectrometer electrospray ionization source (ESI). The chromatographic separation was performed using a 2.1 x 100 mm C18 column packed with 1.7 μm particles (Zorbax XDB C18, Agilent) at 30°C. Mobile phase A consisted of water:tetrahydrofuran:formic acid 0.1%, 890/90/20 v/v/v, while mobile phase B consisted of methanol:acetonitrile:formic acid 0.1%, 400/400/200 v/v/v. The following gradient was used as follows: (0-20 min) 50% B, (20-40 min) 100% B, and (40-45 min) 100% A. The flow rate was 0.2 ml/min at room temperature, and the injection volume was 2 μl.

Equipment: A Waters Acquity UPLC was used. A Waters triple–triple TDQMS/MS mass spectrometer with an electrospray ionization source (ESI) was utilized as a detector. Argonium was employed as a collision gas in the sequential mass analysis. Analyst software (Masslynx) was used for equipment control and data, acquisition and analysis. The analyses were monitored in the full-scan mode, and the mass lines intended for analysis were selected for collision-induced dissociation.

Cultured human cells

Peripheral blood mononuclear cells (PBMC) used in this study were obtained by venipuncture from two healthy volunteers (a man and a woman under 25 years of age), according to OECD guidelines (OECD 487, 2014). The peripheral blood leukocyte (PBL) donors provided written informed consent at the time of donation, as determined by the Ethics Committee. HepG2 cells (human hepatoma cell line) were obtained from the Cell Bank of Rio de Janeiro, Brazil). The PBMC were maintained in cell culture flasks containing RPMI medium, whereas the HepG2 cells were grown in DMEM medium, supplemented with both antibiotics and 10% FCS. Cultures were incubated at 37°C, 5% CO₂, and 95% relative humidity, with complete asepsis. This study was approved by the Human Ethics Committee of the Universidade Estadual Paulista (Marília, SP) on December 4, 2013 (protocol 0839/2013).

Determination of cell viability and cytotoxicity

Cell viability was tested with trypan blue staining, which indicates cell membrane integrity. A total of 2 x 10⁵ cells were placed in each well of a 24-well plate. The concentrations tested were 2.5, 5, 10, 25, 50, 100, 250, 500, 1000, 1250, 2500, and 5000 ug/ml. The culture plates were maintained in an incubator at 37°C, 5% CO₂. The cellular viability test was performed for both cell types according to Strober (2001) method. The culture medium itself was used as a negative control, and Triton X-100 diluted in medium without fetal bovine serum (FBS) culture was used as positive control. Each concentration was tested twice, and each test was performed in triplicate. After 24-hr incubation in the presence of the test substance, cells were harvested and analyzed under an optical microscope, using a Neubauer chamber as base. Non-viable cells were stained blue, and 100 cells per well were counted.

To assess cytotoxicity, a MTT test (3- [4,5-dimethylthiazol-2-yl] -2,5-diphenyltetrazolium) was conducted, with the HepG2 cells Mosmann (1983) with some modifications. The test estimates percent cells capable of transforming MTT into formazan, that is, percent metabolically viable cells. This transformation is indicated by a shift to purple coloration in the solution, which is due to mitochondrial metabolism. The metabolism involves a reduction that utilizes NADH and similar molecules for electron transfer, which leads to the formation of a product which precipitates and accumulates at the bottom of each well. For precipitate dissolution, DMSO was added to the medium, and the plate was read by custom software applied to a spectrophotometer (Riss, 2013).

To perform the MTT assay, 1x10⁴ cells were added to each 96-well plate, which were then filled with culture medium until 2ml was reached. After 24-hr incubation period, 1.4-ml medium was removed. Twenty µl of each test extract concentration was prepared, and volume was made up to 2 ml. Controls were the same as those employed in the trypan blue staining test. After another 24-hr incubation period, 1.4-ml medium was removed and 200 µl MTT was added to each well. The plate was incubated again for a further 4 hr, MTT was removed, and then, 200 µl DMSO was added. After 5 min, the plate was read with a spectrophotometer, at 540 nm filter.

Comet assay

The alkaline comet assay was performed according to Tice et al. (2000). An aliquot of 2 x 10⁵ cells was
placed into 24-well plates in 2-ml respective culture medium per well at 37°C and incubated for 4 hr with the extract, at concentrations of 2.5, 5, 10, 50, and 100 ug/ml; these concentrations were selected on the basis of cell viability and cytotoxicity tests. The positive control was 75 μM MMS, and the negative control was culture medium. Analysis was performed immediately at 400x magnification under fluorescence microscopy with a 515–560 nm excitation filter and a 590 nm barrier filter. All experiments were performed three times and in duplicate.

To determine the extent and distribution of DNA damage, single cell gel electrophoresis (SCGE) assay was conducted by examining at least 100 randomly selected and non-overlapping cells (50 cells per coded slide) per culture well was conducted in a blind fashion. These cells were scored visually (Azqueta et al., 2011), according to tail size into the following four classes: class 0- no tail; class 1- tail shorter than the diameter of the head (nucleus); class 2- tail length 1- to twofold greater than the diameter of the head; and class 3- tail length more than twofold the diameter of the head. Headless comets (class 4) with almost all the DNA in the tail or those with a wide tail were excluded from the evaluation given the likelihood that they were dead cells (Fairbairn et al., 1995; Hartmann & Speit, 1997). The score for each treatment was obtained by multiplying the number of nucleoids observed in each damage class by the value of the class (0, 1, 2, or 3).

**Cytokinesis-block micronucleus (CBMN) test**

The CBMN test was performed according to the Fenech (2000) protocol, having been recently revised and standardized by the OECD (OECD 487, 2014). Cells were cultured and treated with the same five concentrations evaluated in the comet assay. The positive control was 150 μM MMS for PBL and 2 μM benzo[a]pyrene for HepG2 cells, and the negative control was the culture medium itself.

For the test with PBL, whole blood (0.4ml) was added to 5-ml culture medium (RPMI) that had been supplemented with 10% FBS plus 10 µl/ml phytohemagglutinin (PHA) to stimulate mitogenesis. The cultures were incubated at 37°C, under 95% air and 5% CO₂ in a humidified incubator for 72 hr. After 44-hr incubation, the human leukocytes were treated with five different concentrations of the test extract. After 4-hr treatment, cytochalasin B (6 μg/ml) was added to each culture to block cytokinesis. The cells were harvested by centrifugation (5 min at 850 x g), and pellets were resuspended in a chilled hypotonic solution of 0.075 M KCl for 5 min. The cells were then washed once with 5 ml cold methanol: acetic acid solution (3:1, v/v). The fixation procedure was repeated three times. Formaldehyde (1%) was added after the final repetition to preserve the cytoplasm. The cell suspension was placed on slides and stained with 5% Giemsa dye diluted in phosphate buffer (pH 6.8) for 5 min.

In the MN test using HepG2 cells, culture flasks in triplicate were incubated for 24 hr to enable cell growth. Cells were washed again and incubated with five concentrations of extract for 24 hr. Cells were then washed and incubated with cytochalasin B for another 28 hr. After this point, the procedure was equivalent to the leukocytes. A light microscope (Zeiss, Primo Star) was used to quantify micronucleated cells on the slides, with 1000 binucleate cells scored per culture flask at 100x magnification (Fenech, 2006; OECD 487, 2014). As a measure of cytotoxicity, the nuclear division index (NDI) was calculated according the formula NDI = [M1 + 2(M2) + 3(M3) + 4(M4)] /N, where M1–M4 indicate the number of cells with 1–4 nuclei per 500 cells counted (N) (for each culture flask).

**Ames test**

*C. oxyacantha* fruit extract was evaluated in a bacterial mutation assay system, with *Salmonella typhimurium* tester strains TA98 (frameshift) and TA100 (base-pair substitution), using a pre-incubation methodology both with (+S9) and without (-S9) metabolism (Maron & Ames, 1983). The bacteria strains were kindly provided by Dr. B.N. Ames (Berkeley, CA, USA). The strains were grown overnight from frozen cultures for 12–14 hr in Oxoid Nutrient Broth No. 2. The metabolic activation mixture (S9 fraction), prepared from Sprague–Dawley mice livers treated with the polychlorinated biphenyl mixture Aroclor 1254 (500
mg/kg), was freshly prepared before each test. The metabolic activation system consisted of 4% S9 fraction, 1% 0.4 M MgCl2, 1% 1.65 M KCl, 0.5% 1 M D-glucose-6-phosphate disodium and 4% 0.1M NADP, 50% 0.2 M phosphate buffer, and 39.5% sterile distilled water. For the mutagenic assessment, six different concentrations of the extract (2.5, 5, 10, 100, 250, or 500 µg/ml /plate) were tested. The sample concentrations were selected based upon a preliminary toxicity test that determined the highest non-toxic and lowest toxic concentrations. Samples were considered toxic when there was a thinning of the auxotrophic background (i.e., background lawn) accompanied by a decrease in histidine revertants (His+). The concentrations of the test substances were added to 0.5 ml 0.2 M phosphate buffer or to 0.5 ml 4% S9 mixture with 0.1 ml bacterial culture and then incubated at 37°C for 20–30 min. Two-ml surface agar was added, tubes were mixed, and then, the mixture was poured into a Petri dish containing 20-ml minimal agar. The Petri dishes were incubated at 37°C for 66 hr and His+ revertant colonies counted manually. The test was performed in triplicate. Distilled water was used as negative control. The mutagen used as positive control in the tests without S9 mix was 4-nitroquinoline N-oxide (4NQO). For the tests carried out in the presence of S9 mix, the positive control was 2-anthramine (2-AA).

**Statistical analysis**

The results obtained in the CBMN and comet assays were subjected to analysis of variance (ANOVA) followed by Student’s t-test (CBMN) and Tukey’s test (comet assay). GraphPad Prism® software (version 5.02) was used to perform the statistical analyses. In both tests, the results were considered statistically significant at p< 0.05.

For the AMES test, data were analyzed with package SALANAL 1.0 (U.S. Environmental Protection Agency, Monitorin Systems Laboratory, Las Vegas, NV, from Research Triangle Institute, RTP, NC, USA). T Data (revertants/plate) were assessed with ANOVA using the Bernstein et al. (1982) followed by linear regression.

**Results**

According to chromatographic analysis, total flavonoid content of *C. oxyacantha* fruit extract was 2.7%. Four major compounds were identified and quantified: vitexin (m/z 431.2) 0.1%, isovitexin (m/z 431.2) 0.14%, hyperoside (m/z 463.1) 0.63%, and vitexin-2-O-rhamnoside (m/z 577.31) 1.79%. Vitexin-2-O-rhamnoside was the major compound; transition was monitored by 577.31 > 292.80 m/z, which refers to the loss of the sugar moiety, rhamnoside, followed by the loss of C4H8O4, [M - H] - = 577.31; [M - H - C6H12O5] - = 413.25 and transition [M - H - C6H12O5 - C4H8O4] - = 292.83 (Figure 1(a)–1(c)).

In the PBL assessment, extract concentrations above 100 µg/ml resulted in less than 80% cell viability, while for HepG2 cells, decreased viability was observed starting at 250 µg/ml concentration (data not shown). Cytotoxicity results for the MTT test were the same as trypan blue exclusion test. These assay results led to the selection of extract concentrations of 2.5, 5, 10, 50, and 100 µg/ml for assessing genotoxic and clastogenic/aneugenic potential of the extract. The cytotoxicity test performed on TA98 and TA100 *Salmonella* strains led to the selection of the following concentrations: 2.5, 5, 10, 50, and 100 µg/ml.

The results of the comet assay (genotoxicity) are shown in Tables 1 and 2. When the positive control was compared with negative control, a significant difference was noted confirming the test efficacy for detecting DNA damaging agents. Significant increases in total number of cells with DNA damage were found in cells treated with concentrations ≥ 5 µg/ml. In the cells (nucleoids) where DNA damage was observed, it was predominantly minor (class 1). However, in cells treated with extract concentrations of 10 or 50 µg/ml, class 2 DNA damage was higher. Data generated in treated PBL were equivalent to those in HepG2 cells (Table 2).

The results of the CBMN test are presented in Tables 3 and 4. Extract concentrations ≥ 10 µg/ml induced a significant rise in micronucleated binucleate cells compared to negative controls. This effect was detected in both PBL and HepG2 cells, although in HepG2 cells, the frequency of micronucleated cells was higher (Table 4).

Table 5 shows the number of revertant *Salmonella typhimurium* colonies, and TA98 and
TA100 strains per plate, after treatment with the test extract, in both the presence (+S9) and absence (-S9) of metabolic activation. The mutagenicity test showed that all tested concentrations of the extract, after metabolism by S9 fraction, produced mutagenic effects in the TA98 strain.
Table 2. Mean frequency and standard deviation of damaged cells, average distribution between the classes of damage, and average scoring for the assessment of genotoxicity of *C. oxyacantha* fruits extract in HepG2 cells.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Total 1</th>
<th>Comet class</th>
<th>Scores</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Control</td>
<td>10.67 ± 0.57</td>
<td>89.33 ± 0.57</td>
<td>10.00 ± 1.00</td>
</tr>
<tr>
<td>2.5 µg/mL</td>
<td>12.67 ± 0.57</td>
<td>87.33 ± 0.57</td>
<td>11.33 ± 0.57</td>
</tr>
<tr>
<td>5.0 µg/mL</td>
<td>18.33 ± 0.57*</td>
<td>81.67 ± 0.57</td>
<td>17.67 ± 0.57</td>
</tr>
<tr>
<td>10 µg/mL</td>
<td>39.33 ± 1.15*</td>
<td>60.67 ± 1.15*</td>
<td>13.33 ± 1.52</td>
</tr>
<tr>
<td>50 µg/mL</td>
<td>54.67 ± 0.57*</td>
<td>45.33 ± 0.57*</td>
<td>23.00 ± 2.00*</td>
</tr>
<tr>
<td>100 µg/mL</td>
<td>68.33 ± 0.57*</td>
<td>31.67 ± 0.57*</td>
<td>48.33 ± 1.15*</td>
</tr>
<tr>
<td>MMS 75 µM (Positive control)</td>
<td>65.00 ± 1.00*</td>
<td>35.00 ± 1.00*</td>
<td>52.33 ± 0.57*</td>
</tr>
</tbody>
</table>

*Significantly different from the negative control (p < 0.05).

1 Total number of damaged cells (class 1 + 2 + 3).

Table 3. The micronucleus frequency and Nuclear Division Index (NDI) in human lymphocytes treated with *C. oxyacantha* fruits extract, and respective controls.

<table>
<thead>
<tr>
<th>Test substance</th>
<th>Treatment</th>
<th>Binucleated cells With MN (2000 cells scored)</th>
<th>NDI/1000 cells</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Period (h)</td>
<td>Concentration (µg/mL)</td>
<td>Nº</td>
</tr>
<tr>
<td>Control</td>
<td>28</td>
<td>0</td>
<td>6</td>
</tr>
<tr>
<td>MMS (Positive control)</td>
<td>28</td>
<td>150*</td>
<td>97 a</td>
</tr>
<tr>
<td><em>Crataegus oxyacantha</em></td>
<td>28</td>
<td>2.5</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>28</td>
<td>5.0</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>28</td>
<td>10</td>
<td>14 a</td>
</tr>
<tr>
<td></td>
<td>28</td>
<td>50</td>
<td>16 a</td>
</tr>
<tr>
<td></td>
<td>28</td>
<td>100</td>
<td>16 a</td>
</tr>
</tbody>
</table>

Note. MN, micronucleus; SD, standard deviation; MMS, Methyl methanesulfonate; *Concentration, µM,* a Significantly different from the negative control (p < 0.05).

Table 4. The micronucleus frequency and Nuclear Division Index (NDI) in HepG2 cells treated with *C. oxyacantha* fruits extract, and respective controls.

<table>
<thead>
<tr>
<th>Test substance</th>
<th>Treatment</th>
<th>Binucleated cells With MN (3000 cells scored)</th>
<th>NDI/1500cell</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Period (h)</td>
<td>Concentration (µg/mL)</td>
<td>Nº</td>
</tr>
<tr>
<td>Control</td>
<td>24</td>
<td>0</td>
<td>13</td>
</tr>
<tr>
<td>B[a]P (Positive control)</td>
<td>24</td>
<td>150*</td>
<td>146 a</td>
</tr>
<tr>
<td><em>Crataegus oxyacantha</em></td>
<td>24</td>
<td>2.5</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>5.0</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>10</td>
<td>21 a</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>50</td>
<td>23 a</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>100</td>
<td>26 a</td>
</tr>
</tbody>
</table>

Note. MN, micronucleus; SD, standard deviation; B[a]P, Benzo[a]Pyrene; *Concentration, µM,* a Significantly different from the negative control (p < 0.05).

Table 5. Mutagenic Assessment of *C. oxyacantha* fruits Extract in *Salmonella typhimurium* TA98 and TA100 Strains with (+S9) and Without (-S9) Metabolic Activation.

<table>
<thead>
<tr>
<th>Treatments (µg/mL)</th>
<th>Number of revertants (Mean ± SD)/plate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>TA 98 -59</td>
</tr>
<tr>
<td>0</td>
<td>23.00 ± 3.94</td>
</tr>
<tr>
<td>2.5</td>
<td>28.00 ± 1.41</td>
</tr>
<tr>
<td>5.0</td>
<td>26.67 ± 6.43</td>
</tr>
<tr>
<td>100.0</td>
<td>28.33 ± 2.08</td>
</tr>
<tr>
<td>250.0</td>
<td>27.00 ± 2.65</td>
</tr>
<tr>
<td>500.0</td>
<td>25.00 ± 5.20</td>
</tr>
</tbody>
</table>

Note. Data are expressed as mean and standard deviation of the number of revertants.

1 Negative control: dimethyl sulfoxide (DMSO, 100 µL/plate); a Significantly different from the negative control by ANOVA (p < 0.05).
strain. In contrast, the extract without S9 fraction induced no marked mutagenic effects in both strains.

Discussion

Cell viability and cytotoxicity tests carried out with *C. oxyacantha* fruit extract demonstrated that concentrations above 100 µg/ml resulted in decreased cell viability and/or population of human cells in culture. Further, concentrations above 500 µg/ml were cytotoxic to TA98 and TA100 strains of *Salmonella*. The cytotoxicity of the fruit extract observed in our *in vitro* study was higher than that reported for animals with the extract of aerial parts. The German Commission E monography states that mice and rats have been safely given a standardized extract at doses up to 3g/kg body weight (Blumenthal et al., 1998). In humans, the acute oral toxicity of hawthorn was 6 g/kg.

The results of the present study showed that extract concentrations of 5 µg/ml or higher produced genotoxic effects in two types of mammalian cells. The CBMN results indicate that *C. oxyacantha* fruit extract mediated clastogenic and/or aneugenic effects in both tested types of cultured human cells at concentrations of 10 µg/ml or higher. Our study also demonstrated that the TA98 strain, after extract metabolism by S9 fraction, induced a significant increase in number of revertant colonies, indicating that the chemical components of the extract, after undergoing metabolism, might form mutagenic products capable of inducing insertion and/or deletion of DNA base pairs (frameshift mutations).

Tabach et al. (2009) developed a preclinical toxicological evaluation of a phytotherapeutic product termed CPV, which consists of *C. oxyacantha* aerial part extract (26.7 %) in association with *Passiflora incarnata* (33.3 %) and *Valeriana officinalis* (40 %) extracts. Several parameters were investigated in their study including, teratogenicity, mutagenicity, and genotoxicity. Tabach et al. (2009) reported that all of the results were negative, indicating that CPV presented no apparent marked toxicity and appeared devoid of risk for humans. It is possible that the differences between positive genotoxicity results noted in our *in vitro* study compared to Tabach et al. (2009) may be attributed to the fact that a mixture of three plant extracts was used in which the proportion of *C. oxyacantha* aerial part extract (leaves, flowers, and fruits) was smaller and of a different chemical composition than the *C. oxyacantha* fruit extract tested in our study.

The chemical characterization of *C. oxyacantha* fruit extract illustrated significant flavonoid content, including vitexin, isovitexin, hyperoside, and vitexin-2-O-rhamnoside as major compounds. Choo et al. (2012) conducted an *in vivo* acute toxicity test of vitexin and isovitexin by administering them to diabetic and normoglycemic rats and found no apparent signs of toxicity or significant body weight change during the study period. Wenjuan-Wei et al. (2014) examined the effects of vitexin-2-O-rhamnosideo and vitexin-4-O-glucoside on cell growth and apoptosis noting that these compounds exerted no marked cytotoxic effects on human cells derived from adipose tissue. However, few available studies exist, evaluating genetic toxicity of the major components or similar in *C. oxyacantha* fruit extract. Several investigators demonstrated the cytotoxic/mutagenic potential assessment of some flavonoids present in *C. oxyacantha* or other plant genus. Jhoo et al. (2007) identified C-glycoside flavonoids 2-O-rhamnosylvitexin and schaftoside as potential mutagenic compounds in Kava (*Piper methysticum*) by the umu test, a sensitive test for point mutations. Snijman et al. (2007) evaluating the antimutagenic activity of the major flavonoids of rooibos (*Aspalathus linearis*), by using the Ames test, observed that flavonoid–mutagen interactions ranged from antimutagenic, comutagenic, and promutagenic to mutagenic, while vitexin, isovitexin, and hyperoside exhibited only antimutagenic effects. Mohammed et al. (2014) reported potent cytotoxic activity of flavonoids constituents from *Gleditsia triacanthos* L. leaves, against liver, breast, cervix, larynx, and colon cancer cell lines, with vitexin displaying intermediate actions.

In general, toxicity assessments of different formulations of *Crataegus* extract in animals and humans demonstrated low toxicity and few side effects (Ammon & Hänel, 1981). Nascimento (2009) tested the potential toxicity of a mix of
extracts from *Passiflora incarnata*, *Crataegus oxyacantha*, and *Salix alba* on humans with clinical, cardiology, and laboratory exams and found that the administration of this phytotherapeutic compound twice a day for 28 d produced no apparent toxicity. Transient side effects including dizziness, gastrointestinal complaints, headaches, and heart palpitations were reported only occasionally (Daniele et al., 2006).

Under the experimental conditions employed in the present study, the fruit extract of *C. oxyacantha* produced genotoxic and clastogenic/aneugenic effects in human PBL and HepG2 cells, as well as induced frameshift mutation in the TA98 strain of *S. typhimurium* after metabolism by S9 liver enzymes. Despite the therapeutic potential of *C. oxyacantha* fruit extract for humans, these effects raise concern and suggest the need for further *in vitro* and *in vivo* studies investigating genetic toxicity.

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**References**


