Evaluation of the cytotoxic potential of Annona muricata Linn leaf extract against N2a-neuroblastoma cells

Brindha Durairaj, B. Bala Preedeba

Department of Biochemistry, PSG College of Arts and Science, Coimbatore, Tamil Nadu, India

Abstract

Background: Plants are rich in bioactive compounds with various pharmacological properties, and they can be used effectively to treat various diseases. The ability of Annona muricata, preventing progression of cancer cells by initiating programmed cell death, was focused in the present study. Objective: The objectives of this study were to determine the antioxidant and anticancer potential of A. muricata ethanolic extract (AME) of leaves. Materials and Methods: The antioxidant potential of AME was determined by 2,2-diphenylpicrylhydrazyl (DPPH) radical scavenging activity, 2,2’-azinobis(3-ethyl-benzothiozoline)-6-sulfonic acid ammonium salt (ABTS) radical cation scavenging activity, ferric reducing antioxidant power (FRAP) assay, phosphomolybdenum assay, and the cytotoxic potential of AME over N2a cell lines (neuroblastoma cells) was determined using 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide, lactate dehydrogenase (LDH), intracellular reactive oxygen species (ROS), and measurement of mitochondrial membrane potential (MMP) assay. Results: The 50% scavenging ability of the extract in DPPH assay was found to be 14 µg/mL. The ABTS radical scavenging potential was measured to be 10507.3 mg/g of extract. The FRAP was found to be 10.38 mM Fe(II)/mg extract. The effective phosphomolybdenum reduction was measured at a concentration of 53.33 mg ascorbic acid equivalents/g extract. The in vitro cytotoxicity of AME with escalating concentrations (5–85 µg/ml) was first screened, and it was shown to inhibit N2a cells with IC50 value of 55 µg/ml. A significant increase in cytosolic enzyme LDH confirms the cell membrane damage. The extent of cell damage was confirmed by a significant increase in the level of intracellular ROS and a decrease in MMP. Furthermore, to characterize and identify the major active constituents present in AME, Fourier-transform infrared and gas chromatography-mass spectrometry analysis were carried out. Conclusion: The present results demonstrated that AME could be a promising candidate for developing anticancer agents which helps further to combat neuroblastoma with an effective treatment strategy.

Key words: Annona muricata, antioxidant, cytotoxicity, lactate dehydrogenase, neuroblastoma, reactive oxygen species

INTRODUCTION

Neuroblastoma is the second most common malignant pediatric cancer which affects children under the age of 5 years.[1] In 2013, the American Cancer Society reported that neuroblastoma accounts for ~7% of childhood cancer and is responsible for ~15% of deaths in children younger than the age of 15 years. It affects the postganglionic sympathetic nervous system derived from the neural crest cells during embryonic development. Initially, it develops in the adrenal gland and metastasizes to the liver, bone, bone marrow, lymph nodes, neck, and chest. This tumor comprises of heterogeneous population of cells that differ morphologically, biochemically, and genetically from one another.[2] The available treatment strategy includes the drugs vincristine, cisplatin, Neostar, and nitrogenous mustard compounds, but their extensive use confers for alteration in fundamental cellular processes which leads to multidrug resistance.[3] Even the multimodal therapy such as surgery, radiotherapy in conjugation with chemotherapy, and monoclonal antibody-based immunotherapy for high-risk neuroblastoma remains...
unsuccessful.\cite{4,5} Hence, there is a need for alternative medicine to combat neuroblastoma. Phytomedicine, as they possess antioxidants and other phytoconstituents with greater pharmacological property, offers promising effects for cancer prevention with fewer or no side effects. In recent decades, in recent decades, the extensive research over plant compounds for their therapeutic property has been revealed, especially over the leaves of Annonaceae family (annonaceous acetogenins) for their anticancer properties\cite{6,7} and their promising effects against several diseases on various cell lines.\cite{8-11} Based on these evidence, we sought to investigate the cytototoxic potential of *A. muricata* ethanolic extract (AME) against N2a cells.

**MATERIALS AND METHODS**

**Collection and Identification of AME**

The leaves of *Annona muricata* were collected from Sulur region, Coimbatore, Tamil Nadu. The taxonomic identity of the plant was confirmed and a voucher specimen (No: BSI/SRC/5/23/2015/ Tech.153) has been deposited at the Herbarium of the Botany Department, Tamil Nadu Agricultural College, Coimbatore, Tamil Nadu. The leaves of the plants were made dust free, and the surface of leaves was washed well with tap water and was air dried in the shade for 10–15 days. The dried leaves were weighed, powdered, and used further for extraction process.

**Extraction of AME**

The powdered leaves of *A. muricata* were weighed and immersed in 50% ethanol solvent and placed in 4°C for 7 days by occasional shaking (cold maceration). The extract was filtered using Whatman No. 1 filter paper and was concentrated by drying in hot air oven. The powder of 50% ethanolic extract collected was used for further assays.

**Chemicals**

2,2-diphenylpicrylhydrazyl (DPPH), 2,2’-azinobis(3-ethyl-benzothiazoline)-6-sulfonic acid ammonium salt (ABTS), trolox (6-hydroxy-2,5,7,8-tetra methylchroman-2-carboxylic acid), 2,4,6-Tris (2-pyridyl)-s-triazine (TPTZ), 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT), dimethyl sulfoxide (DMSO), Eagle’s minimal essential medium (MEM), and fetal bovine serum (FBS) were obtained from HiMedia Laboratories (Mumbai), Sigma-Aldrich (USA), and Merck (Bengaluru). All the chemicals and solvents used were of analytical reagent grade.

**Cell Line**

The neuroblastoma (N2a) cell line was obtained from the National Centre for Cell Science, India. The cells were seeded into plates and flasks in 1:1, MEM supplemented with 10% FBS, antibiotic, and antimycotic solution (Sigma, St. Louis, MO, USA) in a conditions of 5% CO\textsubscript{2} and 95% air at 37°C. The media were changed on alternate days and treated with different concentrations (5–85 µg/mL) of plant extract dissolved in DMSO and water (1:9 v/v).

**In Vitro Antioxidant Studies**

**DPPH radical scavenging activity**

The antioxidant activity of the extract was determined in terms of hydrogen-donating or radical scavenging ability using the stable radical DPPH.\cite{12} Sample with varying concentration was used and the volume was adjusted to 100 µL with methanol. About 3 mL of a 0.1 mM methanolic solution of DPPH was added to the aliquots of samples and standards (BHT and rutin) were vortexed well. Negative control was prepared by adding 100 µL of methanol in 3 mL of 0.1 mM methanolic solution DPPH. The tubes were allowed to stand in the dark for 15 min at room temperature. The absorbance of the sample was measured at 517 nm against the blank. Radical scavenging activity of the samples was expressed as IC\textsubscript{50} which is the concentration of the sample required to inhibit 50% of DPPH concentration.

**ABTS radical cation scavenging activity**

The total antioxidant activity (TAA) of the samples was measured by ABTS radical cation decolorization assay.\cite{13} ABTS\textsuperscript{+} was produced on the reaction of 7 mM ABTS aqueous solution with 2.4 mM potassium persulfate in the dark for 12–16 h at the room temperature. Before assay, this solution was diluted with ethanol (about 1:89 v/v) and equilibrated at 30°C to produce an absorbance of 0.700 ± 0.02 at 734 nm. After the addition of 1 mL of diluted ABTS\textsuperscript{+} solution to 10 µL of plant extract, all the test tubes were vortexed well and incubated in the dark for 30 min at room temperature. Triplicate determinations were made at each dilution of the standards (BHT and rutin) plant extracts, and the absorbance was read against the blank at 734 nm. The unit of TAA is defined as the concentration of Trolox having equivalent antioxidant activity expressed as µM TE/g of plant extract.

**Ferric reducing antioxidant power (FRAP) assay**

The antioxidant capacities of different extracts of samples were estimated.\cite{14} FRAP reagent (900 µL), prepared freshly and incubated at 37°C for 30 min, was mixed with 90 µL of distilled water and 30 µL of test sample or blank. The test samples and reagent blank were incubated at 37°C for 30 min in a water bath. The final dilution of the test sample in the reaction mixture was 1/34. The FRAP reagent was prepared by mixing 2.5 mL of 20 mM TPTZ in 40 mM HCl, 2.5 mL of 20 mM FeCl\textsubscript{3}, 6H\textsubscript{2}O, and 25 mL of 0.3 M acetate buffer (pH 3.6). At the end of incubation, the absorbance readings were taken immediately at 593 nm against the reagent blank, using a spectrophotometer. Methanolic solutions of known Fe (II)
concentration, ranging from 100 to 2000 μM (FeSO$_4$·7H$_2$O), were used for the preparation of the calibration curve. The parameter equivalent concentration was defined as the concentration of antioxidant having a ferric-TPTZ reducing ability equivalent to that of 1 mM FeSO$_4$·7H$_2$O. Equivalent concentration was calculated as the concentration of antioxidant giving an absorbance increase in the FRAP assay equivalent to the theoretical absorbance value of a 1 mM concentration of Fe (II) solution. BHT, rutin, and quercetin were used as standards.

**Phosphomolybdenum assay**

The antioxidant activities of extracts were evaluated by the formation of green phosphomolybdenum complex.[15] About 100 μL aliquots of extract was added with 3 mL of reagent solution (0.6 M sulfuric acid, 28 mM sodium phosphate, and 4 mM ammonium molybdate). The test tubes were covered with foil and incubated in a water bath at 95°C for 90 min. After the samples had cooled to room temperature, the absorbance of the mixture was measured at 695 nm against the reagent blank. The results were reported by means of mean value expressed as mg ascorbic acid equivalent (AAE)/g extract. BHT, rutin, and quercetin were used as reference standards.

**Characterization of AME Extract**

**Fourier-transform infrared (FT-IR) analysis**

FT-IR spectra were obtained from KBr (potassium bromide) pellets prepared using 1 mg of powdered whole plant. The pellets were analyzed in the absorption mode of FT-IR, and all spectra were recorded from 4000 to 400 cm$^{-1}$ at a data acquisition rate of 2 cm$^{-1}$ using a FT-IR spectrophotometer (Shimadzu).

**Gas chromatography–mass spectrometry (GC-MS) analysis**

GC-MS analysis was carried out using Clarus 600 GC system equipped with Clarus 600 C mass spectrometer (PerkinElmer precisely, USA). An Elite-5MS fused silica capillary column coated with a 5% diphenyl/95% dimethyl polysiloxane stationary phase (60 m × 0.25 mm, film thickness 0.10 m, PerkinElmer precisely, USA) was used for GC-MS. The injector temperature was kept at 200°C, whereas the oven temperature was programmed from 70°C to 300°C for a total run time of 30 min. Helium was used as carrier gas at a flow rate of 1 mL/min. An appropriate blank was run, from which the solvent delay was fixed to 4 min. The electron ionization mode with ionization energy of 70 eV, ion source temperature of 200°C, GC interface temperature of 240°C, scan interval of 0.2 s, and fragments ranging from 50 to 600 m/z were set for the MS analysis. About 1 μL of the extract was injected manually in a splitless mode. The mass spectra of the respective peaks obtained in the GC-MS were compared with the mass fragmentation patterns of standards in the Wiley library.

**MTT assay**

Cell viability was defined as the ability of cells to metabolically reduce the MTT to a purple formazan dye. Cells were seeded into individual 96-well plates and incubated under the above conditions. After a day of incubation, cells were treated with various concentration of plant extract ranging from 5 to 85 μg/mL. To obtain IC$_{50}$ values, absorbance was measured at 570 nm in an ELISA multiplate reader and images were taken in Olympus CKX41 microscope equipped with digital camera (E-330). The percentage inhibition of growth was calculated using the formula, % Cell viability = 100−[100×(Ac−At)/Ac], where At = Absorbance value of test compound and Ac = Absorbance value of control.

**Lactate dehydrogenase (LDH) assay**

The cytotoxicity was quantified in terms of plasma membrane damage by measuring the amount of LDH released using LDH kit (Agappe-11407002) according to the manufacturer’s instructions. LDH activity was measured through the oxidation of lactate to pyruvate with simultaneous reduction of nicotinamide adenine dinucleotide (NAD$^+$) at a wavelength of 340 nm. The rate of increase in enzyme activity due to the formation of reduced NADH is directly proportional to the LDH activity in the sample. The N2a cells were plated at a density of 5 × 10$^4$ cells/well in 24-well plate for 24 h, and cells were treated with various concentrations of plant extracts (5–85 μg/mL) for different time intervals (3–24 h). The cells were precipitated by centrifugation at 2500 rpm for 5 min at 4°C, and the supernatant (100 μL) was mixed with 1000 μL reaction mixture and the percentage activity was calculated. The total LDH activity was measured by lysis (2% Triton X-100) of untreated cells. The cells were seeded in Petri dishes (1 × 10$^6$ cells) and then treated with different concentrations (5–85 μg/mL) of extract for 24 h.

**Measurement of intracellular reactive oxygen species (ROS)**

The intracellular ROS was estimated to measure oxidative stress induced by the toxins using oxidation-sensitive dye 2’,7’-dichlorodihydrofluorescein diacetate (DCFH-DA).[16] The assay is based on the principle that the non-fluorescent fluoroscein DCFH-DA derivatives will emit fluorescence after being oxidized by the radicals generated by the toxins. The non-ionic, non-polar DCFH-DA crosses cell membranes and is hydrolyzed enzymatically by intracellular esterases to non-fluorescent DCFH-DA which is oxidized to highly fluorescent dichlorofluorescein (DCF) in the presence of ROS.[17] Therefore, the intracellular DCF fluorescence can be used as an index to quantify the overall oxidative stress generated in the cells. The cells were plated at a density of 5 × 10$^4$ cells/well in 24-well plate for 24 h and the cells were treated with various concentrations of plant extracts (5–85 μg/mL). Cells were then treated with 20 μM DCFH-DA for 30 min, and intracellular ROS was measured by fluorimetric detection of DCF oxidation at an excitation
wavelength of 485 nm and an emission wavelength of 522 nm. The DCF fluorescence intensity is proportional to the amount of ROS formed intracellularly. The fluorescent intensity percentage was measured using fluorescent microscope (ZEES).

**Mitochondrial membrane potential (MMP) assay**

The electrical potential across the inner mitochondrial membrane was measured using the fluorescent dye rhodamine 123 to estimate the mitochondrial membrane integrity.[18] The cells were cultured in 24 well plates and treated as mentioned earlier. After treatment, rhodamine 123 (10 µg/ml) was added to the cells and incubated for 60 min at 37°C. Then, the cells were collected after washing twice with PBS, and the fluorescence was detected at an excitation wavelength of 485 nm and an emission wavelength of 535 nm using Hidex plate chameleon™ V (Finland).

**Statistical Analysis**

The data were analyzed using one-way ANOVA and the data are expressed as the standard mean value. The graph was generated using GraphPad Prism software.

**RESULTS**

**In Vitro Antioxidant Analysis of A. muricata Leaf Extract**

The analysis of antioxidant potential would help in determining the total reducing capacity of the plant extract.

**DPPH Scavenging Activity**

To check the ability of the extracts to stabilize the DPPH radical in-vitro DPPH measurement assay was selected. DPPH was used as a substrate to evaluate the antioxidant activity of natural products from plant and microbial sources. The results are depicted in Figure 1.

Values are mean of triplicate determination (n = 3) ± standard deviation. From Figure 1, it was evident that DPPH scavenging activity was increased with an increase in the concentration of the extract. The 50% scavenging ability of the extract was found to be at 14 µg/mL, and it was found to be higher when compared to the inhibitory concentrations exerted by rutin and BHT standards.

**ABTS Assay**

The ABTS assay is based on the inhibition of absorbance of the radical cation ABTS+ that has a long characteristic wavelength absorption spectrum. The efficiency of ABTS radical scavenging potential of A. muricata leaf extract was estimated to identify the total reducing potential of extract, was found to be 10,507.3 mg/g of extract, and was found to be similar to the reducing potential of the natural antioxidant rutin and greater than that of BHT. The ability of reducing capacity by standard as well as leaf extract is described in Table 1.

**Phosphomolybdenum Assay**

The phosphomolybdenum reduction potential of the leaf extract refers to the ability of the extract to reduce TPTZ-Fe (III) complex to TPTZ-Fe (II). The reducing ability of hydroalcoholic extracts by quenching the ions was compared with rutin and quercetin standards, and the potential was found to be 10.38 mM Fe(II)E/mg extract. Moreover, the concentration of antioxidant in the sample is given in Table 1.

**FT-IR Analysis of AME**

The functional group in the compound can be identified based on the IR peaks that are obtained based on the percentage transmission. The percentage transmission at particular wavelength determines the functional group of the
compound. The peaks obtained for the hydroethanolic extract of *A. muricata* are shown in Figure 2.

**GC-MS Analysis of AME**

GC-MS was performed to identify the active constituents of AME. The GC-MS chromatogram and spectrum of leaf extract are shown in Figure 3.

**MTT Assay**

The cytotoxic effect of *A. muricata* leaf extract with various concentrations ranging from 5–85 µg/mL was observed and it was found to be increase in the concentration of extracts was indirectly proportional to the cell viability, and the result is depicted in Figure 4. After 24 h of extract treatment, all cells exhibited a significant decrease in cell viability even at lower concentration. Moreover, 55 µg of the extract decreased the cell viability to 50%. Moreover, this concentration was fixed to be the IC$_{50}$ concentration and was further confirmed by determining the LDH assay.

**LDH Assay**

Cell membrane damage can be confirmed by measuring the leakage of extracellular LDH enzyme, and it was found that the amount of enzyme measured was directly proportional to the treated concentration of extract. The percentage LDH release with varying concentration of extract is shown in Figure 5.

**Effect of AME on ROS**

Oxidative stress occurs due to the increase in intracellular ROS level and serves as a major cause for the death of the cell. The rise in free radical creates oxidative stress that leads to the activation of mitochondria-mediated induction of intrinsic apoptotic pathway. The ability of extract in generating ROS is represented in Figure 6.

**Determination of MMP**

Loss of MMP is another indicator that indicates the initiation of apoptosis. Hence, the membrane potential was checked by

<p>| Table 1: The reducing potential of hydroalcoholic extract of <em>A. muricata</em> by ABTS, FRAP, and phosphomolybdenum assays |</p>
<table>
<thead>
<tr>
<th>Extracts</th>
<th>ABTS$^*$ (µM TE/g extract)</th>
<th>FRAP (mM Fe (II) E/mg extract)</th>
<th>Phosphomolybdenum reduction (mg AAE/g extract)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydroalcoholic extract of <em>A. muricata</em></td>
<td>10507.3±139.83</td>
<td>10.387±1.131</td>
<td>60.66±7.68</td>
</tr>
<tr>
<td>7BHT</td>
<td>8423.95±96.32</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rutin</td>
<td>9942.69±109.84</td>
<td>99.19±1.90</td>
<td>437.19±6.85</td>
</tr>
<tr>
<td>Quercetin</td>
<td>-</td>
<td>307.47±82.70</td>
<td>494.74±7.95</td>
</tr>
</tbody>
</table>

Values are mean of triplicate determination ($n=3$)±standard deviation. TE: Trolox equivalents, AAE: Ascorbic acid equivalents, Fe (II) E: Fe (II) equivalents, ABTS: 2,2'-azinobis (3-ethyl-benzothiozoline)-6-sulfonic acid ammonium salt, FRAP: Ferric reducing antioxidant power, *A. muricata*: *Annona muricata*

Figure 2: Fourier-transform infrared analysis of *Annona muricata* leaf extract
Durairaj and Preedeba: Cytotoxic potential of *Annona muricata* extract against N2a cells

DISCUSSION

The results of the *in vitro* antioxidant analysis revealed that the plant possesses compounds that could act as a potent antioxidant, and this property of AME extract makes the platform for the use of this plant for treating any disease. FT-IR analysis reveals the functional groups present in the extract by means of peak ranges, and based on the absorption spectrum, the presence of functional groups in the extract was observed to be the compound containing amino group, compound containing aromatic ring, and other compounds with functional groups such as mono substitution group, carbonyl, and methyl groups. GC-MS using the Wiley library, and based on the retention time of the compounds, the active constituents of the AME were identified which includes dodecane, 4H-Pyran-4-one, 1,5-anhydro-6-deoxyhexo-2,3-diolose, methyl triethylsilyl ether, trimethoxyvinylsilane, benzenepropanol, o-chloromethoxybenzene, 7-tetradecene, coumaran, 2,3-dihydro-benzofuran, cyclotetradecane, 13-octadecenal, oxirane, and 3-phenylpropanamide. The compounds observed in the study was similar with the results obtained from ethanol extract of *A. muricata*. The cytotoxic assay reveals that the *A. muricata* leaf extract is highly effective to treat neuroblastoma in dose-dependent manner. The previous study reported the cytotoxic potential of various solvent extracts, and among the studied extracts, ethyl acetate extract of *A. muricata* leaf was found to have very low inhibitory concentration over the lung cancer cells. Results of the present study were in correlation with the earlier works, suggesting that the *A. muricata* leaf ethanol extract is effective in treating neuroblastoma.
The assay revealed that the increase in the leakage of LDH with respect to damage of membrane in a dose-dependent manner, the stable cytosolic enzyme, LDH catalyzes the oxidation of L-lactate to pyruvate. On membrane damage in cells, it is released into the culture medium, suggesting the loss of membrane integrity. Thus, the fixation of inhibitory concentration requires the LDH test and the result suggests a similar concentration as well, and the IC$_{50}$ value was found to be 55 µg/ml. The present study revealed the potential of extract to induce apoptosis by generating ROS. Hence, the efficiency of extract at the concentration of 55 µg/mL could induce apoptosis in tumor cells. During the process of apoptosis, there will be a great loss in MMP due to the formation of pores, and this determination of MMP would help in serving as an indication for the initiation of apoptosis. When the cells were treated with escalating concentration of AME, the MMP decreased significantly in a dose-dependent manner. The 50% of loss in membrane potential is known to be the best concentration to act as a cytotoxic substance, and it was found to be as 55 µg/mL for AME which was best noted for the generation of ROS and for decreasing the MMP of the cells.

The chemical profiling of the AME identified the major constituents of the extract which plays a major in anticancer activities such as 4H-Pyran-4-one, benzofuran, 2-amino-3-hydroxypyridine, and n-hexadecanoic acid. Further, to determine the membrane damage and to understand the importance of oxidative stress, the ROS measurement was done, which revealed that AME works by inducing intrinsic pathway of apoptosis by generating intracellular ROS in mitochondria, generating pores, and decreasing the MMP of tumor cells. Further, the apoptosis initiation can be confirmed by extending the study for pathway analysis.

CONCLUSION

The chemical profiling of the AME identified the major constituents of the extract which plays a major in anticancer activities such as 4H-Pyran-4-one, benzofuran, 2-amino-3-hydroxypyridine, and n-hexadecanoic acid. Further, to determine the membrane damage and to understand the importance of oxidative stress, the ROS measurement was done, which revealed that AME works by inducing intrinsic pathway of apoptosis by generating intracellular ROS in mitochondria, generating pores, and decreasing the MMP of tumor cells. Further, the apoptosis initiation can be confirmed by extending the study for pathway analysis.

ACKNOWLEDGMENT

The authors are thankful to the director and the scientists of DRDO-BU, Coimbatore, for providing all the facilities required to perform the present study.

REFERENCES

15. Prieto P, Pineda M, Aguilar M. Spectrophotometric quantitation of antioxidant capacity through the formation of a phosphomolybdenum complex: Specific application to the determination of Vitamin E. Anal Biochem 1999;269:337-41.

Source of Support: The work was carried out with the support of DRDO-BU, Coimbatore and PSG College of Arts and Science, Coimbatore. Conflict of Interest: None declared.